Herbal Medicines

Second Edition by Joanne Barnes
The use of complementary and alternative medicines (CAM) has increased rapidly over recent years and the global market for herbal products has risen dramatically.

Designed for healthcare professionals, *Herbal Medicines* addresses the issues of quality, safety and efficacy by providing scientifically rigorous, impartial, evidence-based, information on medicinal herbs.

- 152 structured monographs detailing phytochemical, pharmacological and clinical aspects of herbal medicines
- Enables the healthcare professional to advise on the rational and safe use of herbal medicines
- Presents various appendices that group herbs with specific actions, and highlights potential interactions with conventional medicine
- Written by experts, *Herbal Medicines* reliably informs on quality, safety, interactions and efficacy

‘...This is one of the best examples so far of a useful guide for physicians and clinical herbalists...'  

The American Herb Association
Use search to find specific words or phrases anywhere in the text.
**Enter a search**

<table>
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<th>ENTER AS</th>
<th>EFFECT</th>
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<tr>
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<td>documents containing either (or both) of the words</td>
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<td>cat -dog</td>
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- Enter the term you wish to search for in the search box at the top of the page as
  - a single word (e.g. *aspirin*)
  - a phrase or multi-word term enclosed in double quotes (e.g. "renal failure")
  - Several terms separated by spaces (e.g. *warfarin aspirin*)
- Select the sections in which you would like to search
- Click on the search button.
Help

All the basic functions for searching and viewing Herbal Medicines are displayed on a single screen which is divided into 2 main parts:

- **Function buttons and search box**
- **Document display**
Function buttons and search box

There are 5 function buttons on the top bar:

- **Home**
  To go to the home page, click on this button.

- **Contents**
  To show the table of contents, click on this button.

- **Advanced search**
  To perform an advanced text search click on this button.

- **Print**
  To print the current page, click on this button.

- **Help**
  To show help information, click on this button.

The **search box** allows you to search *Herbal Medicines* directly.
Use it to find words or phrases anywhere in the text.

**See related:**

**Entering a search**
Document display

This is the complete text of the document you have selected.

The top line gives the position of the document in the hierarchy, and links to the previous and next documents.

If the document has been displayed in response to a search, the search terms ('hits') are highlighted. If two or more search terms have been entered, a different colour is used to highlight different hits. If the search included synonyms (e.g. adrenaline/epinephrine), those terms will be highlighted in the same colour.

Notes:

See document area for instructions about moving up and down the document and changing the size of the display.

See related:

Printing a document

Moving around Herbal Medicines

Moving around a document
Three new monographs have been added to *Herbal Medicines* online: Butterbur, Mistletoe and Rhodiola. There are 152 monographs now available online.
149 monographs now available

One new monograph has been added to *Herbal Medicines* so that 149 herbs are now included. Furthermore, one of the previous monographs has been extensively rewritten and all the others have been updated. A new section on the chemical constituents of plants used as herbal medicines has been added and 73 general references are now included, reflecting the growth in the literature on medicinal herbs. This increase in the literature, together with additional knowledge on the quality, safety, efficacy and legal requirements for herbal medicinal products, has led to a clear need to update on this subject for healthcare professionals.
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Access keys

Most browsers support jumping to specific links by typing keys defined within the software application. The access keys defined for *Herbal Medicines* are listed below.

access key s
  skip navigation
access key 1
  home page
access key 3
  contents page
access key 4
  search box
access key 6
  help
access key 8
  copyright page
access key 0
  accessibility statement
access key [
  previous page
access key ]
  next page
Using access keys

The way in which access keys are used depends on your operating system and web browser. Listed below are the methods of access for common browsers.

- **Microsoft Windows Operating System**
  - Microsoft Internet Explorer
    - Alt [key] Enter
  - Mozilla/Netscape
    - Alt [key]
  - Opera
    - Shift + Esc [key]

- **Apple Macintosh Operating System**
  - Microsoft Internet Explorer
    - Ctrl [key] Enter
  - Mozilla
    - Ctrl [key] Enter
  - Opera
    - Shift + Esc [key] Enter

- **Linux Operating System**
  - Mozilla
    - Alt [key]
Visual design

*Herbal Medicines* uses cascading style sheets for visual layout. The default stylesheet uses only relative font sizes, compatible with the user-specified "text size" option in visual browsers. For example, if you're using Internet Explorer, you can make your default text size larger under the "View" menu, "Text Size", "Larger" (or "Largest"). A print stylesheet is used to eliminate the need for an additional print page, or a "text-only" version. If your browser or browsing device does not support stylesheets at all, the content of each page is still readable.
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'...This is one of the best examples so far of a useful guide for physicians and clinical herbalists...'  

The American Herb Association
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<td>Herbal Ingredients containing Amines or Alkaloids, or with Sympathomimetic Action</td>
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Help

All the basic functions for searching and viewing *Herbal Medicines* are displayed on a single screen which is divided into 2 main parts:

- **Function buttons and search box**
- **Advanced search**
Function buttons and search box

There are 5 function buttons on the top bar:

- **Home**
  To go to the home page, click on this button.

- **Contents**
  To show the table of contents, click on this button.

- **Advanced search**
  To perform an advanced text search click on this button.

- **Print**
  To print the current page, click on this button.

- **Help**
  To show help information, click on this button.

The **search box** allows you to search *Herbal Medicines* directly.

Use it to find words or phrases anywhere in the text.

See related:

**Entering a search**
Advanced search

Advanced search allows you to perform a text search in specific sections of the text.

When you perform an advanced search, the results display in the main document area, showing how your search has been handled.

Use the tick boxes to choose which sections you want to search.

See related:

Examples of searches

Entering a search
Welcome to Help for *Herbal Medicines*

Click on the links below for information about these topics

- **About Herbal Medicines**
- **How to cite electronic Herbal Medicines**
- **Structure of Herbal Medicines**
- **The Herbal Medicines interface**
  - **Top bar**
    - Function buttons and search box
      - Home Button
      - Contents Button
      - Advanced search Button
      - Print Button
      - Help Button
      - Search box
      - Search Button
  - **Document area**
    - Contents list
    - Document display
    - Text search results display
    - Advanced search
- **How to find information**
  - How to browse using the contents list
  - How to use the Search function
    - Advanced searching features
  - How to view the documents retrieved by a search
- **Navigation**
  - How to move around Herbal Medicines
  - How to move around a document
  - How to use links
  - Links between documents within Herbal Medicines
  - Links to external sources
  - Reference citation lists
- **How to print a document**
- **Examples of searches**
Document display

This is the complete text of the document you have selected.

The top line gives the position of the document in the hierarchy, and links to the previous and next documents.

If the document has been displayed in response to a search, the search terms ("hits") are highlighted. If two or more search terms have been entered, a different colour is used to highlight different hits. If the search included synonyms (e.g. adrenaline/epinephrine), those terms will be highlighted in the same colour.

Notes:

See document area for instructions about moving up and down the document and changing the size of the display.

See related:

Printing a document

Moving around Herbal Medicines

Moving around a document
Home Button

Click on the Home button to go to the site home page.
Contents Button
Contents Button

Click on the Contents button to see the contents list.

See related:

Table of contents
Advanced search Button

Click on the "Advanced search" button to perform a text search in specific sections of the text, or if you want help with using logical operators.

See related:

Search box

Entering a search
Help contents > Top bar > Function buttons and search box > Print Button
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Click on the Print button to print the current document.

See related:

Printing
Help Button

Click on the Help button for context-sensitive help.
Search box

The search box allows you to search *Herbal Medicines* directly.

Use it to find words or phrases anywhere in the text.

**See related:**

[How to use the text search function](#)
How to use the Search function

Use the Search function to find a specific word or phrase.

1. Enter the term you wish to search for in the box at the top of the screen as:
   - a single word (e.g. *arrow*)
   - a phrase enclosed in double quotes (e.g. "*aplastic anaemia*")
   - several terms combined with logical operators - see [advanced searching](#) for more information
2. Click on the **Search** button or hit **Enter**.
   - To search within specified sections click on the [Advanced search](#) button

**Notes:**

Searching is 'free text' so you can search for any word, but allow for spelling variations.

*Herbal Medicines* uses International Non-proprietary Names for titles and British English spelling for its editorial content.

You may not always get the result you expect when searching within sections because sections are sometimes combined (for example Adverse Effects and Precautions).

**See related:**

[Advanced searching](#)

[Using the contents list](#)

[To view the documents retrieved by a search](#)
Document area

The main document area is used to display:

- the text of a **document**
- the **table of contents**
- the **advanced search** page
- **text search results**

To move up or down

- use the scroll bar on the right of the screen
- use the 'Page Up' or 'Page Down' keys
- use the arrow keys
How to print a document

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Print documents from Herbal Medicines using the print functions on your browser.

For example, to print a complete document with Microsoft Internet Explorer

- select the document you wish to print
- on the toolbar, click File
- select Print from the dropdown menu
- in the Print Range box, select All for the complete document or a page range
- in the Copies box, specify the number of copies
- click OK

Notes:

Some documents are very long. Print Preview may not tell you how many pages you will print.

We recommend that you print a complete document. If you select part of a document by highlighting it, the printout may lose its formatting and be difficult to read.

Remember to write the source on the copy using the recommended style for citing Herbal Medicines
Links between documents within *Herbal Medicines*

Related records are linked in the same way whether they are within the same document or in a different document.

1. To display the linked document, click on the link text. The linked document will be displayed in the document area.
2. To return to the original document, click on the browser **Back** button.

Some examples of links include those between

- two monographs
- a reference citation and the full reference details
- a monograph and the General references

**See related:**

**Using links**

**Links to external sources**
How to move around a document

- Click anywhere in the document display to activate that part of the screen
- Use the arrow keys or Page Up and Page Down keys on your keyboard
- Use the scroll bar (or wheel on your mouse, if available)

See related:

Document display
We are grateful to Paul Weller, Linda Horrell and Louise McIndoe of the Pharmaceutical Press for their active help and encouragement with this work. Our thanks also go to Pepi Hurtado-Lopez and Suzannah Harris for their assistance in preparation of the manuscript.
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Overview

*Herbal Medicines* is an electronic database developed from the book of the same name, first published by Pharmaceutical Press in 1986.

Interest in herbal medicines has continued to grow since publication of the first edition of *Herbal Medicines* in 1996. This is shown in several ways, for example, by increased retail sales of herbal medicinal products in Europe and the USA, as well as the greater awareness among the public and healthcare professionals about natural health products and complementary therapies. Industrially produced new herbal products, mainly based on single-herb extracts standardised for a specific active ingredient, continue to be developed. Good-quality clinical research to support the reputed effects of many herbs is lacking. Nevertheless, several herbal medicines have been investigated in well-designed clinical trials and, for a small number of herbs, systematic reviews and meta-analyses of randomised controlled trials have been published.

The safety of herbal products continues to be a matter of concern, even though toxic herbs have been eliminated from products manufactured in developed countries. There is, for example, a growing appreciation that herbal medicines may interact with concurrently used pharmaceutical medicines. The potential for herb–drug interactions has been highlighted by the recognition that St. John’s wort (*Hypericum perforatum*) may interact with certain prescription medicines, including HIV protease inhibitors, oral contraceptives, selective serotonin reuptake inhibitors (SSRIs), theophylline, ciclosporin and warfarin. The havoc that may be caused by the use of a toxic herb, either for perceived medicinal purposes, or through inadvertent or deliberate substitution of a medicinal herb, is well illustrated by species of *Aristolochia* used in traditional Chinese medicine. Nephrotoxic and carcinogenic effects of *Aristolochia* in humans have been reported in Europe, China, Japan and the USA. Disquiet over the safety of herbs used in herbal products, whether licensed as medicines or sold as unlicensed products, is reflected by medicines regulatory authorities world-wide, highlighting the need for health-care professionals to be aware of such problems.

Since publication of the first edition of *Herbal Medicines*, it has become apparent that the text is a valuable source of information for all groups of healthcare professionals, and that it is used throughout Europe, North America and Australasia. The section ‘Introduction to the Monographs’ was completely revised for the second edition of *Herbal Medicines* in 2002 to include a summary of the current position of herbal medicines, mainly in Europe, as well as commenting on the possible future regulation of herbal
medicinal products in the European Union. Seven new herbal monographs were added, 10 of the previous monographs were extensively revised and rewritten, and a further 33 were updated.

Four new monographs have been added to *Herbal Medicines* since the publication of the second print edition, so that 152 herbs are now included. Furthermore, one of the previous monographs has been extensively rewritten and all the others have been updated. A new section on the chemical constituents of plants used as herbal medicines has been added and 73 general references are now included, reflecting the growth in the literature on medicinal herbs. This increase in the literature, together with additional knowledge on the quality, safety, efficacy and legal requirements for herbal medicinal products, has led to a clear need to update on this subject for healthcare professionals.

This electronic reference source will therefore be regularly updated in the future, with additional monographs added and revisions made to existing monographs. New print editions of *Herbal Medicines* will also be published. However, with the advantages that electronic reference sources offer in terms of searchability and currency, healthcare professionals are advised to consult the most up-to-date electronic version of *Herbal Medicines* (for further information contact the Pharmaceutical Press).

The authors welcome comment on the content and functionality of *Herbal Medicines*; please contact them via the publishers.

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Notes on *Herbal Medicines*

**Sub-sections**

- Purpose and Scope
- Introduction to the Monographs
- The Herbal Monographs

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A general disillusionment with conventional medicines, coupled with the desire for a ‘natural’ lifestyle has resulted in an increasing utilisation of complementary and alternative medicine (CAM) across the developed world.

A study of long–term trends in the use of CAM therapies in the United States of America reported that the use of CAM therapies has increased steadily since the 1950s.(1) Use of CAM has increased independent of gender, ethnicity and level of education, but is more common in younger people. The use of herbal medicine increased particularly in the 1970s and then again in the 1990s. The report concluded that the continuing demand for CAM will affect delivery of healthcare for the foreseeable future. Several other studies have documented the growing use of CAM in the United Kingdom, with the most common complementary therapies reported as acupuncture, homeopathy, herbal medicine and manipulative therapies, chiropractic and osteopathy.(2)

In their report on complementary and alternative medicine, the House of Lords Select Committee on Science and Technology’s Subcommittee on Complementary and Alternative Medicine highlighted the lack of comprehensive information on the use of herbal medicines in the UK.(3) Estimates of herbal medicine use are available, but it is difficult to gauge usage accurately as many products are considered to be food supplements. Nevertheless, a national telephone survey of a nationally representative sample of 1204 British adults found that around 7% of those contacted had used herbal medicines in the previous year.(4) In another survey, over 5000 randomly selected adults in England (not the UK) were sent a postal questionnaire on their use of CAM.(5) Around 20% of the respondents had bought an over–the–counter herbal remedy in the previous 12 months.

Estimates of expenditure on herbal medicines vary, but data generally show that the global market for herbal products has grown rapidly in the past decade. In the USA, annual retail sales of herbal medicines were estimated to be US$ 1.6 billion in 1994,(6) and almost US$ 4 billion in 1998.(7) Retail sales
of herbal products in the European Union (EU) were estimated to be US$7000 million in 1996.\(^8\) A detailed analysis of the European herbal medicines market reported that Germany and France make up more than 70% of the market share.\(^9\) In 1997, total sales of herbal products (using wholesale prices) were US$1.8 billion in Germany and US$1.1 billion in France. In the UK, retail sales of herbal products are reported to have increased by 43% in the period from 1994 to 1998, with retail sales of licensed herbal medicinal products reported to be £50 million in 1998.\(^3\)

These figures demonstrate that herbal medicinal products are being used increasingly by the general public on a self–selection basis to either replace or complement conventional medicines. Against this background of increasing usage of herbal medicines by the public, several major public health issues have raised concerns about these products. The substitution of toxic *Aristolochia* species in traditional Chinese medicines (TCM) has resulted in cases of serious renal toxicity and renal cancer in Europe, China and America.\(^10\) The emergence of interactions between St. John’s wort (*Hypericum perforatum*) and certain prescription medicines has necessitated regulatory action world–wide and has highlighted the need for healthcare professionals to have up–to–date scientific information on the quality, safety and efficacy of these products.\(^11\)

Pharmacists need to be able to advise the consumer on the rational and safe use of all medicines. To fulfil this role with respect to herbal medicines, a pharmacist should be reliably informed of their quality, safety and efficacy. Also, many other healthcare professionals are becoming increasingly aware of their patients’ use of herbal medicines and need to be informed of the suitability of these products for use as medicines.

This handbook brings together in one text a series of monographs on 148 herbs commonly present in herbal medicinal products sold through pharmacies in the UK. Various appendices are also presented, grouping together herbs with specific actions, and highlighting potential interactions with conventional medicines.

As a preface to the monographs, an overview of UK and European legislation concerning herbal products is provided, together with issues pertaining to their quality, safety and efficacy. In addition to retail purchase, herbs can be obtained by picking the wild plant or from a herbal practitioner. This handbook does not discuss the self–collection of plant material for use as a herbal remedy or the prescribing of herbal medicines by herbal practitioners.
All living organisms produce numerous chemical substances that are termed natural products. Those natural products that are common to all life forms are known collectively as primary metabolites and are exemplified by carbohydrates, proteins and fats. Thus, many of the chemical building blocks of primary metabolism are found in all medicinal plants (e.g. amino acids, common sugars, such as glucose, and fatty acids). In addition to primary metabolites, plants also produce other compounds with a more restricted distribution and these are referred to collectively as secondary metabolites. Plants are a rich source of secondary metabolites and some of these are of such limited distribution that they are found only in a particular genus (e.g. *Papaver*), or even in only a single species (e.g. *Papaver somniferum*). On the other hand, some secondary metabolites are widely distributed throughout many of the plant families. It is not always understood why particular plants produce specific secondary metabolites, but some of them are known to have definite functions; for example, some are toxic and form a defence against predators, while some are attractive to insects and aid pollination. Whatever their roles are within plants, many of them have pharmacological actions and this has been exploited to provide medicinal drugs such as codeine, morphine, digoxin and quinine. Some secondary metabolites have proved to be too toxic for human use (e.g. aconitine from aconite), but investigations of their mode of action have stimulated research into synthetic analogues as potential therapeutic agents.

For more details of plant secondary metabolites, the reader is recommended to consult specialist texts, for example on the biosynthesis of medicinal natural products (e.g. Dewick, 2002\(^1\)), on scientific background to herbal medicines (e.g. Evans, 2002;\(^2\) Heinrich *et al.*, 2003\(^3\)) and on specific chemical structures (e.g. Harborne and Baxter, 1993;\(^4\) *Dictionary of Natural Products* CD-ROM\(^5\)).

Numerous secondary metabolites are derived from a common biosynthetic precursor; for example, shikimic acid is involved in the formation of coumarins, lignans, phenylpropanes and tannins. Although each of these groups of secondary metabolites is different chemically they all contain a common structural feature, namely a C6–C3 moiety. Among the most...
prevalent of secondary metabolites are alkaloids, glycosides and phenols. Examples of these types of secondary metabolite can be found in many of the medicinal plants.
General References

General references are cited with the prefix 'G' in the monographs (eg 'G1').


Dr Joanne Barnes
BPharm, PhD, MRPharmS, FLS

Jo Barnes is a pharmacist and Lecturer in Phytopharmacy at the Centre for Pharmacognosy & Phytotherapy at the School of Pharmacy, University of London. Her main research interests are the clinical efficacy and safety of herbal products (phytomedicines), the role of the pharmacist in ensuring the safe and effective use of complementary medicines, and the implications of complementary medicine use for pharmaceutical care.

Dr Barnes holds a PhD from the University of London and a postgraduate certificate in pharmaco vigilance and pharmacoepidemiology from the London School of Hygiene and Tropical Medicine.

Dr Barnes was previously a research fellow in the Department of Complementary Medicine at the University of Exeter, where she was also Senior Editor of the review journal Focus on Alternative and Complementary Therapies (FACT). She is a member of the editorial board or editorial advisory committee of several journals, including Complementary Therapies in Medicine, the International Journal of Pharmacy Practice, Phytotherapy Research, FACT and Drug Safety. She is a member of the Royal Pharmaceutical Society’s Science Committee’s working group on complementary/alternative medicine, and the Independent Review Panel on Classification of Borderline Products.

Her career has included employment as a clinical pharmacist at Queen’s Medical Centre in Nottingham, a medical information pharmacist at Fisons Pharmaceuticals, and a period as a pharmacist at the Royal London Homoeopathic Hospital. She has also worked in pharmaceutical publishing, as a features editor for Adis International Ltd in Auckland, New Zealand, before returning to the UK to begin an academic career. Dr Barnes is a regular contributor to the Pharmaceutical Journal.
Dr Linda A Anderson
BPharm, PhD, FRPharmS

Linda Anderson obtained her first degree in Pharmacy and her PhD in Pharmacognosy at the Welsh School of Pharmacy in Cardiff. She was a postdoctoral research and teaching fellow at the School of Pharmacy, University of London from 1981 to 1987.

Dr Anderson joined the Medicines Control Agency (Department of Health, UK) in 1987.

Within the MCA, she was initially involved in the assessment of new chemical entities and is now mainly involved with abridged applications and has specific responsibility for herbal products. She is Principal Assessor to the Committee on Safety of Medicine’s (CSM) Sub-Committee on Chemistry, Pharmacy and Standards (CPS) and is UK delegate to the European Committee on Proprietary Medicinal Products (CPMP) Quality Working Party. Dr Anderson is also UK Delegate to the European Medicines Evaluation Agency (EMEA) Working Group on Herbal Medicinal Products and is Vice-Chair of the British Pharmacopoeia Committee on Crude Drugs and Galenicals. She is a member of the Royal Pharmaceutical Society’s Science Committee’s working group on complementary/alternative medicine.

Linda Anderson has been awarded a Fellowship of the Royal Pharmaceutical Society of Great Britain (2001).
David Phillipson graduated BSc in Pharmacy (1956), MSc (1959), and DSc (1979) from the University of Manchester, and PhD (1965) from the University of London. He was Lecturer in Pharmacognosy (1961–1972) at the Department of Pharmacy, Chelsea College, London, and Senior Lecturer (1973–1979), Reader in Phytochemistry (1979–1981), and Professor and Head of the Department of Pharmacognosy (1981–1994) at the School of Pharmacy, University of London. On retirement he became Emeritus Professor of Pharmacognosy. In 1995 he was appointed as Wilson T S Wang Distinguished International Visiting Professor at the Chinese University of Hong Kong from January to June. His research included investigations of the chemistry and biological activities of natural products from higher plants with special interests in indole and isoquinoline alkaloids, and plants used in traditional medicines for the treatment of malaria and other protozoal diseases. Collaboration with the pharmaceutical industry included the application of radioligand–receptor binding assays in the search for natural products with activity in the central nervous system.

David Phillipson has received awards from the Phytochemical Society of Europe including the Tate and Lyle Award (1982), Medal (1994), and the Pergamon Prize for Creativity in Plant Biochemistry (1996). He was awarded the Korber Foundation Prize for Achievement in European Science (1989) in collaboration with Professor M.H. Zenk of the University of Munich and four other European colleagues, was presented with the Harrison Memorial Medal of the Royal Pharmaceutical Society of Great Britain (1999), and with the Sir Hans Sloane Medal of the Faculty of the History and Philosophy of Medicine and Pharmacy, Society of Apothecaries (2001). In 1985 he was Science Chairman of the British Pharmaceutical Conference and has been Secretary (1977–1982), Vice-Chairman (1982–1984, 1986–1988) and Chairman (1984–1986) of the Phytochemical Society of Europe. He has been awarded Fellowships of the Royal Pharmaceutical Society of Great Britain (1980) and of the School of Pharmacy, University of London (1998).

He has supervised 33 PhD students and 11 postdoctoral researchers, publishing some 210 full research papers, 150 short communications, 42 review articles, and has edited six books on natural products. Collaborative research was established with scientists in many countries world–wide and in 1989 he was appointed Honorary Professor of the Chinese Academy of Medical Sciences at the Institute of Medicinal Plant Research and Development, Beijing, China. For 19 years he was a member of the Natural
Products Group of the International Foundation for Science, Sweden, helping to award research grants to individual young scientists in developing countries. He has been a member of a number of national and international committees.
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## Monographs

- Bayberry
- Bilberry
- Bloodroot
- Blue Flag
- Bogbean
- Boldo
- Boneset
- Borage
- Broom
- Buchu
- Burdock
- Burnet
- Butterbur

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- Calamus
- Calendula
- Capsicum
- Cascara
- Cassia
- Cat’s Claw
- Celery
- Centaury
- Cereus
- Chamomile, German
- Chamomile, Roman
- Chaparral
- Cinnamon
- Clivers
- Clove
- Cohosh, Black
- Cohosh, Blue
- Cola
- Coltsfoot
- Comfrey
- Corn Silk
- Couchgrass
- Cowslip
- Cranberry

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<td>Drosera</td>
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- Echinacea
- Elder
- Elecampane
- Ephedra
- Eucalyptus
- Euphorbia
- Evening Primrose
- Eyebright

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<td>Ginseng, Eleutherooccus</td>
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<td>Ginseng, Panax</td>
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<td>Horehound, White</td>
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Monographs

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### Monographs

- Marshmallow
- Maté
- Meadowsweet
- Melissa
- Milk Thistle
- Mistletoe
- Motherwort
- Myrrh

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<td>Pleurisy Root</td>
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<td>Poplar</td>
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<td>Prickly Ash, Northern</td>
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<td>Prickly Ash, Southern</td>
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<td>Pulsatilla</td>
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Monographs: Q

Monographs

- Quassia
- Queen’s Delight

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Monographs: R

Monographs

- Raspberry
- Red Clover
- Rhodiola
- Rhubarb
- Rosemary

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Monographs: S

- Sage
- Sarsaparilla
- Sassafras
- Saw Palmetto
- Scullcap
- Senega
- Senna
- Shepherd’s Purse
- Skunk Cabbage
- Slippery Elm
- Squill
- St. John’s Wort
- Stone Root

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- Tansy
- Thyme

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Monographs: V

Monographs

- Valerian
- Vervain

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Monographs

- Wild Carrot
- Wild Lettuce
- Willow
- Witch Hazel

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## Monographs: Y

### Monographs

- Yarrow
- Yellow Dock
- Yucca

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Species (Family)

Various *Yucca* species (Liliaceae/Agavaceae) including

i. *Yucca schidigera* Roezl ex Ortgies

ii. *Yucca brevifolia* Engelm.

iii. *Yucca glauca*
Synonym(s)

i. Mohave Yucca, *Yucca mohavensis* Sarg.

ii. Joshua Tree, *Yucca arborescens* Trel.
Part(s) Used

Whole plant
Pharmacopoeial and Other Monographs

Martindale 33rd edition\textsuperscript{(G67)}

PDR for Herbal Medicines 2nd edition\textsuperscript{(G36)}
Legal Category (Licensed Products)

Yucca is not listed in the GSL.
Terpenoids
Various saponins have been isolated from different *Yucca* species, including tigogenin and chlerogenin,\(^1\) yuccagenin and kammogenin,\(^2\) sarsaspogenin, markogenin, higogenin, neo–tigogenin, neo–gitogenin, hecogenin, gloriogenin, and diosgenin (trace)\(^3\) and smilagenin.
Yucca filamentosa L. (bear grass) is listed by the Council of Europe as a natural source of food flavouring (category N3). This category indicates that there is insufficient information available for an adequate assessment of potential toxicity.\(^{G16}\) The yucca plant has been used traditionally as a major foodstuff by Indian tribes. In the USA, both *Y. schidigera* and *Y. brevifolia* are approved for food use.\(^{G41}\)
Herbal Use

Yucca has been used for the treatment of arthritis, diabetes and stomach disorders. Concentrated plant juice has been used topically to soothe painful joints.
Dosage

None documented.
Pharmacological Actions

In vitro and animal studies

In the rat, anti-inflammatory activity against carrageenan-induced inflammation has been documented for a saponin-containing leaf extract from *Yucca schottii*.\(^{(2)}\) Yucca saponin extract, from *Y. schidigera*, is reported to exhibit approximately half the haemolytic activity of commercial soap bark saponin.

Antitumour activity against B16 melanoma has been documented for a polysaccharide-containing extract of *Y. glauca*.\(^{(4)}\) The extract was found to be inactive towards L1210 or P388 leukaemias.

Clinical studies

A saponin-containing yucca extract has been reported to reduce symptoms of swelling, pain and stiffness in approximately 75 of 150 arthritic patients given the extract in a double-blind study.\(^{(5)}\) The onset of a positive response was found to vary from days to weeks or months. A saponin-containing yucca extract has also been documented to reduce blood pressure, abnormal triglyceride, and high cholesterol concentrations in a double-blind study involving 212 arthritic and hypertensive patients.\(^{(6)}\) Optimum results were obtained in conjunction with diet and exercise. Yucca extracts have also been reported to provide relief from headaches and to improve circulation and gastrointestinal function.\(^{(5,6)}\)
Side-effects, Toxicity

Limited toxicity data are available for yucca. A 12-week study in rats concluded that yucca was non-toxic. A saponin-containing yucca extract was given to more than 700 arthritic patients with no signs of toxicity documented. The yucca saponins are regarded to be a safe food supplement since they are not thought to be absorbed from the gastrointestinal tract, thereby reducing the dangers of systemic haemolytic activity.⁵
Contra–indications, Warnings

**Pregnancy and lactation**
There are no known problems with the use of yucca during pregnancy and lactation. However, it is advisable not to exceed amounts normally ingested as a food.
Pharmaceutical Comment

Limited phytochemical information is available for yucca, steroidal saponins being the only documented constituents. Human studies have reported a yucca saponin extract to have a beneficial effect on certain symptoms of arthritis such as pain and stiffness, and to reduce blood pressure and serum triglyceride and cholesterol concentrations. The traditional use of yucca as a foodstuff would indicate it to be of low toxicity.
References

See also General References G16 G32 G36 G41.


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Appendix 1 Potential Drug–Herb Interactions

Until the emergence of reports of important interactions between St. John’s wort (*Hypericum perforatum* L.) and certain conventional drugs (see St. John’s Wort), very few interactions involving herbal products had been reported in the medical literature. Furthermore, there has been very little experimental and clinical research in this area.

The following list of potential drug–herb interactions has been compiled on the basis of known herbal constituents and their reported pharmacological actions. It should be emphasised that many drug interactions are harmless and many of those that are potentially harmful occur only in a small proportion of patients and may then vary in severity from patient to patient. Healthcare professionals should be alert to undeclared use of herbal medicines as a possible cause of unexplained toxicity or lack of effect of conventional medicines.

Suspected drug–herb interactions involving licensed or unlicensed herbal products should be reported to the regulatory authorities, as for any other suspected adverse reaction to drugs or herbs.

### Drug/therapeutic category affected interacting Possible effects

#### Gastrointestinal system

<table>
<thead>
<tr>
<th>Herbal ingredients interacting</th>
<th>Possible effects</th>
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<tbody>
<tr>
<td>Herbal ingredients irritant to gastrointestinal tract. See Appendix 13</td>
<td>Exacerbation of symptoms Risk of systemic side–effects</td>
</tr>
<tr>
<td>Herbal ingredients with laxative activity. See Appendix 2</td>
<td>Antagonism</td>
</tr>
<tr>
<td>Herbal ingredients with laxative activity. See Appendix 2</td>
<td>Potentiation; increased risk of side–effects</td>
</tr>
<tr>
<td>Cardiovascular system</td>
<td>Cardioactive herbal ingredients. See Appendix 3</td>
</tr>
<tr>
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<tr>
<td>Herbal ingredients containing hydroxyanthracene laxatives. See Appendix 2</td>
<td>Potentiation; increased risk of side–effects</td>
</tr>
<tr>
<td>St. John’s wort</td>
<td>Risk of reduced therapeutic effect of digoxin</td>
</tr>
<tr>
<td>Herbal ingredients containing hydroxyanthracene laxatives. See Appendix 2</td>
<td>Potentiation; increased risk of hypokalaemia</td>
</tr>
<tr>
<td>Herbal ingredients with diuretic activity. See Appendix 4</td>
<td>Potentiation; increased risk of hypokalaemia</td>
</tr>
<tr>
<td>Herbal ingredients with hypotensive activity. See Appendix 5</td>
<td>Difficulty in controlling diuresis; hypotension</td>
</tr>
<tr>
<td>Anti–arrhythmic activity</td>
<td>Herbal ingredients containing hydroxyanthracene laxatives. See Appendix 2</td>
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<tr>
<td>Cardioactive herbal ingredients. See Appendix 3</td>
<td>Interference/antagonism with existing therapy</td>
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<tr>
<td>Herbal ingredients with</td>
<td>Antagonism if hypokalaemia</td>
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<tr>
<td>Category</td>
<td>Interaction</td>
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<tr>
<td>Beta–adrenoceptor blocking drugs</td>
<td>Diuretic activity. See Appendix 4</td>
</tr>
<tr>
<td></td>
<td>Cardioactive herbal ingredients. See Appendix 3</td>
</tr>
<tr>
<td></td>
<td>Herbal ingredients with significant amine content or sympathomimetic activity. See Appendix 14</td>
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<td>Antihypertensive therapy</td>
<td>Potential antagonism</td>
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<td>Potential risk of severe hypertension</td>
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<td>Potentiation</td>
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<td></td>
<td>Risk of potentiation/interference with existing therapy</td>
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<tr>
<td>Lipid-lowering drugs</td>
<td>Herbal ingredients with hypolipidaemic activity. See Appendix 7</td>
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<td>Additive effect</td>
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<td><strong>Nitrates and calcium–channel blockers</strong></td>
<td><strong>Cardioactive ingredients. See Appendix 3</strong></td>
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<td>Cohosh, blue</td>
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<td>Herbal ingredients with hypertensive activity. See Appendix 5</td>
<td>Interference with therapy</td>
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<td>Herbal ingredients with anticholinergic activity</td>
<td>Reduced sublingual absorption of glycercyl trinitrate</td>
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<tr>
<td>Herbal ingredients with significant sympathomimetic amine content. See Appendix 14</td>
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<tr>
<td>Herbal ingredients with hypertensive activity. See Appendix 5</td>
<td>Increased risk of hypertension</td>
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<td>Increased risk of hypertension</td>
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<th><strong>Risk of antagonism or potentiation</strong></th>
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<p>| <strong>Herbal ingredients with</strong> | | |
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<td>Horse–chestnut</td>
<td>Plasma protein binding</td>
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<td>St. John’s wort</td>
<td>Risk of reduced therapeutic effect of warfarin</td>
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**Respiratory System**

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<td>May increase arrhythmogenic potential of terfenadine</td>
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<td>Electrolyte imbalance may increase arrhythmogenic potential of terfenadine</td>
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<td>Allergic Disorders</td>
<td>Potentiation of drowsiness associated with antihistamines</td>
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**Central Nervous System**
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<td>Herbal ingredients claimed to have sedative activity. See Appendix 8</td>
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<td>Ginseng</td>
<td>Increased risk of ginseng side–effects</td>
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<td><strong>Antipsychotics</strong></td>
<td>Herbal ingredients with diuretic activity. See Appendix 4</td>
<td>Potentiation of lithium therapy; increased risk of toxicity; diuretics reported to reduce lithium clearance</td>
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<td>Herbal ingredients with anticholinergic activity</td>
<td>Risk of interference with therapy; anticholinergic drug reported to reduce plasma phenothiazine concentrations</td>
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<td>Evening primrose</td>
<td>Potential risk of seizures</td>
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<td><strong>Antidepressants</strong></td>
<td>Herbal ingredients containing sympathomimetic amines. See Appendix 14</td>
<td>Risk of hypertensive crisis with monoamine–oxidase inhibitors (MAOIs)</td>
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<td>Ginseng</td>
<td>Suspected phenelzine interaction</td>
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<td>White horehound</td>
<td>Hydroxytryptamine antagonism, <em>in vivo</em></td>
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<td>Antagonism; contra–indicated in patients with depressive illness</td>
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<td>Risk of increased serotonergic effects in patients taking selective serotonin reuptake inhibitors (SSRIs)</td>
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<thead>
<tr>
<th>Drugs used in nausea and vertigo</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Herbal ingredients with sedative activity. See Appendix 8</strong></td>
</tr>
<tr>
<td>May potentiate sedative side–effects</td>
</tr>
<tr>
<td><strong>Herbal ingredients with anticholinergic activity</strong></td>
</tr>
<tr>
<td>Antagonism</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Analgesics</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Herbal ingredients with diuretic activity. See Appendix 4</strong></td>
</tr>
<tr>
<td>Increased risk of toxicity with anti–inflammatory analgesics</td>
</tr>
<tr>
<td><strong>Herbal ingredients with corticosteroid activity, e.g. bayberry, liquorice. See Appendix 10</strong></td>
</tr>
<tr>
<td>Possible reduction in plasma–aspirin concentrations</td>
</tr>
<tr>
<td><strong>Herbal ingredients with sedative activity. See Appendix 8</strong></td>
</tr>
<tr>
<td>May potentiate sedative side–effects</td>
</tr>
<tr>
<td><strong>St. John’s wort</strong></td>
</tr>
<tr>
<td>Risk of increased serotonergic effects, with possibility of increased risk of side–effects</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Antiepileptics</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Herbal ingredients with sedative activity. See</strong></td>
</tr>
<tr>
<td>May potentiate sedative side–effects</td>
</tr>
</tbody>
</table>
Appendix 8

Borage  May increase risk of seizure

Evening primrose oil  May increase risk of seizure

Ground ivy  May increase risk of seizure

Sage  May increase risk of seizure

Herbal ingredients with significant salicylate content (meadowsweet, poplar, willow)  Transient potentiation of phenytoin therapy may occur. See Appendix 6

Herbal ingredients with significant folic acid content  Plasma phenytoin concentration may be reduced

St. John’s wort  Risk of reduced therapeutic effect of anticonvulsants (carbamazepine, phenobarbitone, phenytoin)

Drugs for parkinsonism  Herbal ingredients with anticholinergic activity  Potentiation; increased risk of side–effects

Herbal ingredients with cholinergic activity  Antagonism

Infections

Antifungal drugs  Herbal ingredients with anticholinergic activity  Risk of reduced absorption of ketoconazole
HIV protease inhibitors

St. John’s wort

Risk of reduced blood concentrations of anti-HIV drugs, with possible loss of HIV suppression

Endocrine system

Antidiabetics

Herbal ingredients with hypo- or hyperglycaemic activity. See Appendix 9

Herbal ingredients with diuretic activity. See Appendix 4

Potentiation/antagonism of activity

Antagonism

Drugs for hypo- and hyperthyroidism

Herbal ingredients with significant iodine content e.g. fucus

Horseradish, myrrh

Interference with therapy

Interference with therapy

Corticosteroids

Herbal ingredients with diuretic activity. See Appendix 4

Herbal ingredients with corticosteroid activity e.g. bayberry, liquorice. See Appendix 10

Risk of increased potassium loss

Increased risk of side-effects, e.g. water and sodium retention

Sex hormones

Herbal ingredients with hormonal activity. See Appendix 10

Possible interaction with existing therapy
### Obstetrics and gynaecology

<table>
<thead>
<tr>
<th>Oral contraceptives</th>
<th>Herbal ingredients with hormonal activity. See Appendix 10</th>
<th>Possible interaction with existing therapy; may reduce effectiveness of oral contraceptive</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>St. John’s wort</td>
<td>Risk of reduced blood concentrations of oral contraceptives, breakthrough bleeding and unintended pregnancy</td>
</tr>
</tbody>
</table>

### Malignant disease and immunosuppression

<table>
<thead>
<tr>
<th>Methotrexate</th>
<th>Herbal ingredients with significant salicylate content. See Appendix 6</th>
<th>Increased risk of toxicity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Drugs affecting immune response</td>
<td>Herbal ingredients with immunostimulant activity. See Appendix 11</td>
<td>Potentiation or antagonism</td>
</tr>
<tr>
<td></td>
<td>St. John’s wort</td>
<td>Risk of reduced therapeutic effect of ciclosporin</td>
</tr>
</tbody>
</table>

### Musculoskeletal and joint diseases

<table>
<thead>
<tr>
<th>Systemic lupus erythematous</th>
<th>Alfalfa</th>
<th>Antagonism; contra–indicated</th>
</tr>
</thead>
<tbody>
<tr>
<td>Probenecid</td>
<td>Herbal ingredients with significant salicylate content. See Appendix 6</td>
<td>Risk of inhibition of probenecid</td>
</tr>
<tr>
<td>Eye</td>
<td>Herbal ingredients with significant salicylate content. See Appendix 6</td>
<td>Increased risk of toxicity</td>
</tr>
<tr>
<td>-----------------------------</td>
<td>------------------------------------------------------------------------</td>
<td>-----------------------------</td>
</tr>
<tr>
<td>Skin</td>
<td>Herbal ingredients with potential allergenic activity. See Appendix 12</td>
<td>Allergic reaction; exacerbation of existing symptoms</td>
</tr>
<tr>
<td></td>
<td>Herbal ingredients with phototoxic activity. See Appendix 12</td>
<td>Phototoxic reaction; exacerbation of existing symptoms</td>
</tr>
<tr>
<td>Anaesthetics</td>
<td></td>
<td></td>
</tr>
<tr>
<td>General anaesthetics</td>
<td>Herbal ingredients with hypotensive activity. See Appendix 5</td>
<td>Potentiation of hypotensive effect</td>
</tr>
<tr>
<td>Competitive muscle relaxants</td>
<td>Herbal ingredients with diuretic activity. See Appendix 4</td>
<td>Risk of potentiation if hypokalaemia occurs</td>
</tr>
<tr>
<td>Depolarising muscle relaxants</td>
<td>Cardioactive herbal ingredients. See Appendix 3</td>
<td>Risk of arrhythmias</td>
</tr>
</tbody>
</table>
### Appendix 2 Laxative Herbal Ingredients

<table>
<thead>
<tr>
<th>Drug</th>
<th>Effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aloes</td>
<td>Hydroxyanthracene constituents</td>
</tr>
<tr>
<td>Cascara</td>
<td>Hydroxyanthracene constituents</td>
</tr>
<tr>
<td>Eyebright</td>
<td>Iridoids, <em>in vivo</em></td>
</tr>
<tr>
<td>Frangula</td>
<td>Hydroxyanthracene constituents</td>
</tr>
<tr>
<td>Horehound, White</td>
<td>Large doses</td>
</tr>
<tr>
<td>Ispaghula</td>
<td>Bulk laxative</td>
</tr>
<tr>
<td>Plantain</td>
<td>Iridoids, <em>in vivo</em> (much less than senna)</td>
</tr>
<tr>
<td>Rhubarb</td>
<td>Hydroxyanthracene constituents</td>
</tr>
<tr>
<td>Senna</td>
<td>Hydroxyanthracene constituents</td>
</tr>
<tr>
<td>Yellow Dock</td>
<td>Hydroxyanthracene constituents</td>
</tr>
</tbody>
</table>
## Appendix 3 Cardioactive Herbal Ingredients

<table>
<thead>
<tr>
<th>Drug</th>
<th>Effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>Broom</td>
<td>Alkaloid constituents: cardiac depressant activity</td>
</tr>
<tr>
<td>Calamus</td>
<td>Anti–arrhythmic activity</td>
</tr>
<tr>
<td>Cereus</td>
<td>Tyramine: cardiotonic amine</td>
</tr>
<tr>
<td>Cola</td>
<td>Caffeine</td>
</tr>
<tr>
<td>Coltsfoot</td>
<td>Cardiac calcium–channel blocking activity</td>
</tr>
<tr>
<td>Devil’s Claw</td>
<td>Activity <em>in vivo</em></td>
</tr>
<tr>
<td>Fenugreek</td>
<td>Activity <em>in vitro</em></td>
</tr>
<tr>
<td>Figwort</td>
<td>Cardioactive glycoside constituents, activity <em>in vitro</em></td>
</tr>
<tr>
<td>Fumitory</td>
<td>Alkaloid constituent: cardioactive</td>
</tr>
<tr>
<td>Ginger</td>
<td>Activity <em>in vivo</em></td>
</tr>
<tr>
<td>Ginseng, Panax</td>
<td>Activity <em>in vivo</em></td>
</tr>
<tr>
<td>Golden Seal</td>
<td>Berberine: cardioactive alkaloid</td>
</tr>
<tr>
<td>Hawthorn</td>
<td>Tyramine: cardiotonic amine; activity <em>in vivo</em></td>
</tr>
<tr>
<td>Horehound, White</td>
<td>Activity <em>in vivo</em></td>
</tr>
</tbody>
</table>
Lime Flower  Activity reputed with excessive ingestion

Maté  Caffeine

Mistletoe  Viscotoxin, negative inotropic effect

Motherwort  Cardiac glycoside constituents; activity *in vitro*

Parsley  Apiole poisoning, high doses

Pleurisy Root  Cardenolides, active *in vitro* and *in vivo*

Prickly Ash, Northern  Interaction with Na\(^+\)K\(^+\) ATPase

Prickly Ash, Southern  Interaction with Na\(^+\)K\(^+\) ATPase

Quassia  Activity *in vitro*

Shepherd’s Purse  Activity *in vitro*

Squill  Cardiac glycoside constituents

Wild Carrot  Depressant activity *in vivo*
# Appendix 4 Diuretic Herbal Ingredients

<table>
<thead>
<tr>
<th>Drug</th>
<th>Effect</th>
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</thead>
<tbody>
<tr>
<td>Agrimony</td>
<td>Activity <em>in vivo</em></td>
</tr>
<tr>
<td>Artichoke</td>
<td>Reputed action</td>
</tr>
<tr>
<td>Boldo</td>
<td>Irritant oil</td>
</tr>
<tr>
<td>Broom</td>
<td>Reputed action</td>
</tr>
<tr>
<td>Buchu</td>
<td>Reputed action</td>
</tr>
<tr>
<td>Burdock</td>
<td>Reputed action</td>
</tr>
<tr>
<td>Celery</td>
<td>Reputed action</td>
</tr>
<tr>
<td>Corn Silk</td>
<td>Activity <em>in humans</em></td>
</tr>
<tr>
<td>Couchgrass</td>
<td>Activity <em>in vivo</em></td>
</tr>
<tr>
<td>Dandelion</td>
<td>Activity <em>in vivo</em></td>
</tr>
<tr>
<td>Devil’s Claw</td>
<td>Reputed action</td>
</tr>
<tr>
<td>Elder</td>
<td>Activity <em>in vivo</em></td>
</tr>
<tr>
<td>Guaiacum</td>
<td>Reputed action</td>
</tr>
<tr>
<td>Java Tea</td>
<td>Activity <em>in vivo</em></td>
</tr>
<tr>
<td>Plant</td>
<td>Description</td>
</tr>
<tr>
<td>-------------</td>
<td>----------------------------------</td>
</tr>
<tr>
<td>Juniper</td>
<td>Reputed action; terpinen-4-ol</td>
</tr>
<tr>
<td>Nettle</td>
<td>Activity in humans</td>
</tr>
<tr>
<td>Pokeroott</td>
<td>Activity <em>in vivo</em></td>
</tr>
<tr>
<td>Shepherd’s Purse</td>
<td>Activity <em>in vivo</em></td>
</tr>
<tr>
<td>Squill</td>
<td>Activity <em>in vivo</em></td>
</tr>
<tr>
<td>Uva-Ursi</td>
<td>Reputed action</td>
</tr>
<tr>
<td>Yarrow</td>
<td>Activity <em>in vivo</em></td>
</tr>
<tr>
<td>Drug</td>
<td>Effect</td>
</tr>
<tr>
<td>---------------------</td>
<td>---------------------------------------------</td>
</tr>
<tr>
<td><strong>Hypotensive</strong></td>
<td></td>
</tr>
<tr>
<td>Agrimony</td>
<td>Hypotensive, <em>in vivo</em></td>
</tr>
<tr>
<td>Asafoetida</td>
<td>Hypotensive, <em>in vivo</em></td>
</tr>
<tr>
<td>Avens</td>
<td>Hypotensive, <em>in vivo</em></td>
</tr>
<tr>
<td>Calamus</td>
<td>Hypotensive, <em>in vivo</em></td>
</tr>
<tr>
<td>Cat’s Claw</td>
<td>Hypotensive, <em>in vivo</em></td>
</tr>
<tr>
<td>Celery</td>
<td>Hypotensive, human and <em>in vivo</em></td>
</tr>
<tr>
<td>Cohosh, Black</td>
<td>Hypotensive, <em>in vivo</em></td>
</tr>
<tr>
<td>Corn Silk</td>
<td>Hypotensive, <em>in vivo</em></td>
</tr>
<tr>
<td>Cowslip</td>
<td>Hypotensive, then hypertensive <em>in vivo</em></td>
</tr>
<tr>
<td>Devil’s Claw</td>
<td>Hypotensive, <em>in vivo</em></td>
</tr>
<tr>
<td>Elecampane</td>
<td>Hypotensive, <em>in vivo</em></td>
</tr>
<tr>
<td>Fenugreek</td>
<td>Hypotensive</td>
</tr>
<tr>
<td>Fucus</td>
<td>Hypotensive</td>
</tr>
<tr>
<td>Plant</td>
<td>Effect</td>
</tr>
<tr>
<td>-------------</td>
<td>-------------------------------</td>
</tr>
<tr>
<td>Fumitory</td>
<td>Hypotensive, <em>in vivo</em></td>
</tr>
<tr>
<td>Garlic</td>
<td>Hypotensive, human and <em>in vivo</em></td>
</tr>
<tr>
<td>Ginger</td>
<td>Hypotensive</td>
</tr>
<tr>
<td>Ginseng, Panax</td>
<td>Hypotensive, human and <em>in vivo</em></td>
</tr>
<tr>
<td>Golden Seal</td>
<td>Hypotensive, alkaloid effect</td>
</tr>
<tr>
<td>Hawthorn</td>
<td>Hypotensive, <em>in vivo</em></td>
</tr>
<tr>
<td>Horehound, White</td>
<td>Vasodilator (volatile oil)</td>
</tr>
<tr>
<td>Horse-chestnut</td>
<td>Hypotensive, <em>in vivo</em></td>
</tr>
<tr>
<td>Horseradish</td>
<td>Hypotensive, <em>in vivo</em></td>
</tr>
<tr>
<td>Java Tea</td>
<td>Hypotensive, <em>in vivo</em></td>
</tr>
<tr>
<td>Juniper</td>
<td>Hypotensive, <em>in vivo</em></td>
</tr>
<tr>
<td>Mistletoe</td>
<td>Hypotensive, <em>in vivo</em></td>
</tr>
<tr>
<td>Nettle</td>
<td>Hypotensive, <em>in vivo</em></td>
</tr>
<tr>
<td>Parsley</td>
<td>Hypotensive, <em>in vivo</em></td>
</tr>
<tr>
<td>Plantain</td>
<td>Hypotensive, <em>in vivo</em></td>
</tr>
<tr>
<td>Herb</td>
<td>Effect Description</td>
</tr>
<tr>
<td>---------------------</td>
<td>------------------------------------------</td>
</tr>
<tr>
<td>Pokeroot</td>
<td>Hypotensive, <em>in vivo</em></td>
</tr>
<tr>
<td>Prickly Ash, Northern</td>
<td>Hypotensive, <em>in vivo</em></td>
</tr>
<tr>
<td>Prickly Ash, Southern</td>
<td>Hypotensive, <em>in vivo</em></td>
</tr>
<tr>
<td>Sage</td>
<td>Hypotensive</td>
</tr>
<tr>
<td>Shepherd’s Purse</td>
<td>Hypotensive</td>
</tr>
<tr>
<td>Squill</td>
<td>Vasodilator, <em>in vivo</em></td>
</tr>
<tr>
<td>St. John’s Wort</td>
<td>Hypotensive, <em>in vivo</em></td>
</tr>
<tr>
<td>Thyme</td>
<td>Hypotensive, <em>in vivo</em></td>
</tr>
<tr>
<td>Vervain</td>
<td>Hypotensive</td>
</tr>
<tr>
<td>Wild Carrot</td>
<td>Hypotensive, <em>in vivo</em></td>
</tr>
<tr>
<td>Yarrow</td>
<td>Hypotensive, <em>in vivo</em></td>
</tr>
</tbody>
</table>

**Hypertensive**

<table>
<thead>
<tr>
<th>Herb</th>
<th>Effect Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bayberry</td>
<td>Hypertensive, myricitrin mineralocorticoid side–effect</td>
</tr>
<tr>
<td>Broom</td>
<td>Hypertensive, alkaloid effect, stated to be contra–indicated in hypertensive individuals</td>
</tr>
<tr>
<td>Capsicum</td>
<td>Hypertensive, increased catecholamine secretion</td>
</tr>
</tbody>
</table>
Cohosh, Blue  Hypertensive, methylcytisine has nicotinic action, alkaloid effect

Cola  Hypertensive, caffeine

Coltsfoot  Hypertensive, pressor activity

Ephedra  Hypertensive, human and *in vivo*

Gentian  Stated to be contra–indicated in hypertensive individuals

Ginger  Hypertensive

Ginseng, Panax  Hypertensive, human and *in vivo*

Liquorice  Hypertensive, mineralocorticoid side–effect

Maté  Hypertensive, caffeine

Vervain  Hypertensive
<table>
<thead>
<tr>
<th>Drug</th>
<th>Effect</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Anticoagulants</strong></td>
<td></td>
</tr>
<tr>
<td>Alfalfa</td>
<td>Coumarin constituents</td>
</tr>
<tr>
<td>Angelica</td>
<td>Coumarin constituents</td>
</tr>
<tr>
<td>Aniseed</td>
<td>Coumarin constituents</td>
</tr>
<tr>
<td>Arnica</td>
<td>Coumarin constituents</td>
</tr>
<tr>
<td>Asafoetida</td>
<td>Coumarin constituents, anticoagulant <em>in vivo</em></td>
</tr>
<tr>
<td>Bilberry</td>
<td>Inhibits platelet aggregation, human, <em>in vivo, in vitro</em></td>
</tr>
<tr>
<td>Boldo</td>
<td>Coumarin constituents</td>
</tr>
<tr>
<td>Capsicum</td>
<td>Coumarin constituents</td>
</tr>
<tr>
<td>Cat’s Claw</td>
<td>Inhibits platelet aggregation, <em>in vitro</em></td>
</tr>
<tr>
<td>Celery</td>
<td>Coumarin constituents</td>
</tr>
<tr>
<td>Chamomile, German</td>
<td>Coumarin constituents</td>
</tr>
<tr>
<td>Chamomile, Roman</td>
<td>Coumarin constituents</td>
</tr>
<tr>
<td>Clove</td>
<td>Eugenol inhibitor of platelet activity</td>
</tr>
</tbody>
</table>
Dandelion  Coumarin constituents
Fenugreek  Coumarin constituents
Feverfew  Inhibits platelet aggregation
Fucus  Anticoagulant action
Garlic  Interaction with warfarin reported
Ginger  Inhibition of platelet activity
Ginkgo  Inhibition of platelet activity
Ginseng, Panax  Reduction of blood coagulation
Horse-chestnut  Coumarin constituents
Horseradish  Peroxidase stimulates synthesis of arachidonic acid metabolites
Liquorice  Inhibition of platelet activity
Meadowsweet  Salicylate constituents
Nettle  Coumarin constituents
Passionflower  Coumarin constituents
Poplar  Salicylate constituents
Prickly Ash, Northern
Coumarin constituents

Prickly Ash, Southern
Coumarin constituents

Quassia
Coumarin constituents

Red Clover
Coumarin constituents

Willow
Salicylate constituents

Coagulants

Agrimony
Coagulant, human

Golden Seal
Heparin antagonist

Mistletoe
Lectins, agglutinating activity

Yarrow
Coagulant, *in vivo*
# Appendix 7 Hypolipidaemic and Hyperlipidaemic Herbal Ingredients

<table>
<thead>
<tr>
<th>Drug</th>
<th>Effect</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Hypocholesterolaemic</strong></td>
<td></td>
</tr>
<tr>
<td>Alfalfa</td>
<td>Hypocholesterolaemic, <em>in vivo</em></td>
</tr>
<tr>
<td>Artichoke</td>
<td>Hypocholesterolaemic, <em>in vivo</em>, human</td>
</tr>
<tr>
<td>Bilberry</td>
<td>Hypocholesterolaemic, <em>in vivo</em></td>
</tr>
<tr>
<td>Capsicum</td>
<td>Hypocholesterolaemic, <em>in vivo</em></td>
</tr>
<tr>
<td>Cohosh, Black</td>
<td>Hypocholesterolaemic, <em>in vivo</em></td>
</tr>
<tr>
<td>Fenugreek</td>
<td>Hypocholesterolaemic, <em>in vivo</em>, human</td>
</tr>
<tr>
<td>Garlic</td>
<td>Hypocholesterolaemic, <em>in vivo</em>, human</td>
</tr>
<tr>
<td>Ginger</td>
<td>Hypocholesterolaemic, <em>in vivo</em></td>
</tr>
<tr>
<td>Ispaghula</td>
<td>Hypocholesterolaemic, human</td>
</tr>
<tr>
<td>Milk thistle</td>
<td>Hypocholesterolaemic, <em>in vivo</em></td>
</tr>
<tr>
<td>Myrrh</td>
<td>Hypolipidaemic, <em>in vivo</em>, human</td>
</tr>
<tr>
<td>Plantain</td>
<td>Hypocholesterolaemic, <em>in vivo</em></td>
</tr>
</tbody>
</table>
Scullcap  Hypocholesterolaemic, *in vivo*

Senega  Hypolipidaemic, *in vivo*

Tansy  Hypocholesterolaemic, *in vivo*

**Hypercholesterolaemic**

Hydrocotyle  Hypercholesterolaemic, *in vivo*
## Appendix 8 Sedative Herbal Ingredients

<table>
<thead>
<tr>
<th>Drug</th>
<th>Effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calamus</td>
<td>Potentiates barbiturate sleeping time</td>
</tr>
<tr>
<td>Celery</td>
<td>In vivo</td>
</tr>
<tr>
<td>Centaury</td>
<td>Reputed action</td>
</tr>
<tr>
<td>Chamomile, German</td>
<td>Human</td>
</tr>
<tr>
<td>Couchgrass</td>
<td>In vivo</td>
</tr>
<tr>
<td>Elecampane</td>
<td>In vivo</td>
</tr>
<tr>
<td>Ginseng, Eleutherococcus; Ginseng, Panax</td>
<td>CNS depressant and stimulant</td>
</tr>
<tr>
<td>Golden Seal</td>
<td>In vivo</td>
</tr>
<tr>
<td>Hawthorn</td>
<td>CNS depressant; potentiates barbiturate sleeping time</td>
</tr>
<tr>
<td>Hops</td>
<td>In vivo</td>
</tr>
<tr>
<td>Hydrocotyle</td>
<td>In vivo</td>
</tr>
<tr>
<td>Jamaica Dogwood</td>
<td>In vivo</td>
</tr>
<tr>
<td>Nettle</td>
<td>CNS depression, <em>in vivo</em></td>
</tr>
<tr>
<td>Plant</td>
<td>Action</td>
</tr>
<tr>
<td>---------------------</td>
<td>--------------------------------------------------</td>
</tr>
<tr>
<td>Passionflower</td>
<td><em>In vivo</em></td>
</tr>
<tr>
<td>Sage</td>
<td><em>In vivo</em></td>
</tr>
<tr>
<td>Scullcap</td>
<td>Reputed action</td>
</tr>
<tr>
<td>Senega</td>
<td>CNS depressant, <em>in vivo</em></td>
</tr>
<tr>
<td>Shepherd’s Purse</td>
<td>Potentiates barbiturate sleeping time</td>
</tr>
<tr>
<td>St. John’s Wort</td>
<td>Traditional use, biflavonoids</td>
</tr>
<tr>
<td>Valerian</td>
<td>Human, <em>in vivo</em></td>
</tr>
<tr>
<td>Wild Carrot</td>
<td><em>In vivo</em></td>
</tr>
<tr>
<td>Wild Lettuce</td>
<td><em>In vivo</em>, related species</td>
</tr>
</tbody>
</table>
### Hypoglycaemic and Hyperglycaemic Herbal Ingredients

<table>
<thead>
<tr>
<th>Drug</th>
<th>Effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>Agrimony</td>
<td>Hypoglycaemic, <em>in vivo</em></td>
</tr>
<tr>
<td>Alfalfa</td>
<td>Hypoglycaemic, manganese, human</td>
</tr>
<tr>
<td>Aloes/Aloe vera</td>
<td>Hypoglycaemic, <em>in vivo</em></td>
</tr>
<tr>
<td>Burdock</td>
<td>Hypoglycaemic, <em>in vivo</em></td>
</tr>
<tr>
<td>Celery</td>
<td>Hypoglycaemic, <em>in vivo</em></td>
</tr>
<tr>
<td>Corn Silk</td>
<td>Hypoglycaemic, <em>in vivo</em></td>
</tr>
<tr>
<td>Damiana</td>
<td>Hypoglycaemic</td>
</tr>
<tr>
<td>Dandelion</td>
<td>Hypoglycaemic, <em>in vivo</em></td>
</tr>
<tr>
<td>Elecampane</td>
<td>Hypoglycaemic</td>
</tr>
<tr>
<td>Eucalyptus</td>
<td>Hypoglycaemic, <em>in vivo</em></td>
</tr>
<tr>
<td>Fenugreek</td>
<td>Hypoglycaemic, human</td>
</tr>
<tr>
<td>Garlic</td>
<td>Hypoglycaemic, <em>in vivo</em>, human</td>
</tr>
</tbody>
</table>
Ginger  Hypoglycaemic, *in vivo*

Ginseng, Panax  Hypoglycaemic

Ispaghula  Hypoglycaemic, humans

Java Tea  Hypoglycaemic *in vivo*

Juniper  Hypoglycaemic *in vivo*

Marshmallow  Hypoglycaemic, *in vivo*, human

Myrrh  Hypoglycaemic, *in vivo*

Nettle  Hypoglycaemic, *in vivo*

Sage  Hypoglycaemic, *in vivo*

Senega  Hypoglycaemic, *in vivo*

Tansy  Hypoglycaemic, *in vivo*

**Hyperglycaemic**

Elecampane  Hyperglycaemic

Ginseng, Panax  Hyperglycaemic

Hydrocotyle  Hyperglycaemic, human

Liquorice  Reduced $K^+$ aggravates glucose tolerance
Rosemary Hyperglycaemic, *in vivo*
# Appendix 10 Hormonally Active Herbal Ingredients

<table>
<thead>
<tr>
<th>Drug</th>
<th>Effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>Agnus Castus</td>
<td>Many uses in hormonal imbalance disorders</td>
</tr>
<tr>
<td>Agrimony</td>
<td>Oestrogenic</td>
</tr>
<tr>
<td>Alfalfa</td>
<td>Oestrogenic, <em>in vivo</em></td>
</tr>
<tr>
<td>Aniseed</td>
<td>Oestrogenic</td>
</tr>
<tr>
<td>Bayberry</td>
<td>Mineralocorticoid</td>
</tr>
<tr>
<td>Cohosh, Black</td>
<td>Oestrogenic</td>
</tr>
<tr>
<td>Fucus</td>
<td>Hyper-/hypothyroidism reported</td>
</tr>
<tr>
<td>Ginseng, Eleutherococcus; Ginseng, Panax</td>
<td>Oestrogenic, human</td>
</tr>
<tr>
<td>Horseradish</td>
<td>May depress thyroid activity</td>
</tr>
<tr>
<td>Liquorice</td>
<td>Mineralocorticoid activity, human; oestrogenic <em>in vivo, in vitro</em></td>
</tr>
<tr>
<td>Motherwort</td>
<td>Oxytocic</td>
</tr>
<tr>
<td>Pleurisy Root</td>
<td>Oestrogenic</td>
</tr>
<tr>
<td>Red Clover</td>
<td>Oestrogenic <em>in vivo</em></td>
</tr>
<tr>
<td>Herb</td>
<td>Characteristics</td>
</tr>
<tr>
<td>--------------</td>
<td>------------------------------------------------------</td>
</tr>
<tr>
<td>Saw Palmetto</td>
<td>Oestrogenic and anti–androgenic <em>in vivo</em>; human use in prostate cancer</td>
</tr>
<tr>
<td>Vervain</td>
<td>Inhibition of gonadotrophin activity</td>
</tr>
<tr>
<td>Wild Carrot</td>
<td>Oestrogenic</td>
</tr>
</tbody>
</table>
## Appendix 11 Immunomodulating Herbal Ingredients

<table>
<thead>
<tr>
<th>Drug</th>
<th>Effects</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alfalfa</td>
<td>Stimulant, <em>in vitro</em></td>
</tr>
<tr>
<td>Boneset</td>
<td>Stimulant, <em>in vitro</em></td>
</tr>
<tr>
<td>Calendula</td>
<td>Stimulant, <em>in vitro</em></td>
</tr>
<tr>
<td>Cat’s Claw</td>
<td>Stimulant, <em>in vitro</em></td>
</tr>
<tr>
<td>Chamomile, German</td>
<td>Stimulant, <em>in vitro</em></td>
</tr>
<tr>
<td>Drosera</td>
<td>Stimulant and depressant (<em>in vitro</em>)</td>
</tr>
<tr>
<td>Echinacea</td>
<td>Stimulant, <em>in vitro</em>, <em>in vivo</em></td>
</tr>
<tr>
<td>Ephedra</td>
<td>Inhibits complement pathway, <em>in vitro</em></td>
</tr>
<tr>
<td>Ginseng, Eleutherococcus</td>
<td>Stimulant, <em>in vivo</em>, human</td>
</tr>
<tr>
<td>Mistletoe</td>
<td>Stimulant, <em>in vivo</em>, human; suppressant (high doses), human</td>
</tr>
<tr>
<td>Saw Palmetto</td>
<td>Stimulant, <em>in vivo</em></td>
</tr>
<tr>
<td>Drug</td>
<td>Effect</td>
</tr>
<tr>
<td>--------------------</td>
<td>------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Agnus Castus</td>
<td>Allergic effects reported</td>
</tr>
<tr>
<td>Angelica</td>
<td>Furanocoumarins, photosensitivity, contact allergy</td>
</tr>
<tr>
<td>Aniseed</td>
<td>Furanocoumarins, photosensitivity, contact allergy</td>
</tr>
<tr>
<td>Apricot</td>
<td>Contact allergy, kernels</td>
</tr>
<tr>
<td>Arnica</td>
<td>Contact allergy</td>
</tr>
<tr>
<td>Artichoke</td>
<td>Sesquiterpene lactone constituents</td>
</tr>
<tr>
<td>Asafoetida</td>
<td>Irritant gum, contact allergy</td>
</tr>
<tr>
<td>Boneset</td>
<td>Sesquiterpene lactone constituents</td>
</tr>
<tr>
<td>Calendula</td>
<td>Individuals sensitive to plants from the Compositae/Asteraceae families</td>
</tr>
<tr>
<td>Cassia</td>
<td>Allergic reactions, mainly contact</td>
</tr>
<tr>
<td>Celery</td>
<td>Furanocoumarins, photosensitivity</td>
</tr>
<tr>
<td>Chamomile, German</td>
<td>Sesquiterpene lactone constituents</td>
</tr>
<tr>
<td>Chamomile, Roman</td>
<td>Sesquiterpene lactone constituents</td>
</tr>
<tr>
<td>Plant</td>
<td>constituent/property</td>
</tr>
<tr>
<td>------------------</td>
<td>----------------------------------------------------------</td>
</tr>
<tr>
<td>Cinnamon</td>
<td>Contact allergy</td>
</tr>
<tr>
<td>Corn Silk</td>
<td>Allergic reactions</td>
</tr>
<tr>
<td>Cowslip</td>
<td>Allergic reactions</td>
</tr>
<tr>
<td>Dandelion</td>
<td>Sesquiterpene lactone constituents</td>
</tr>
<tr>
<td>Elecampane</td>
<td>Sesquiterpene lactone constituents</td>
</tr>
<tr>
<td>Euphorbia</td>
<td>Histamine potentiating properties</td>
</tr>
<tr>
<td>Feverfew</td>
<td>Sesquiterpene lactone constituents</td>
</tr>
<tr>
<td>Fucus</td>
<td>Iodine may aggravate/trigger acne</td>
</tr>
<tr>
<td>Garlic</td>
<td>Sulfur–containing compounds, allergic reaction</td>
</tr>
<tr>
<td>Ginger</td>
<td>Dermatitis in sensitive individuals</td>
</tr>
<tr>
<td>Ginkgo</td>
<td>Fruit pulp and seeds: severe allergic reactions</td>
</tr>
<tr>
<td>Gravel Root</td>
<td>Sesquiterpene lactone constituents</td>
</tr>
<tr>
<td>Guaiacum</td>
<td>Irritant resin</td>
</tr>
<tr>
<td>Holy Thistle</td>
<td>Sesquiterpene lactone constituents</td>
</tr>
<tr>
<td>Hops</td>
<td>Contact allergy</td>
</tr>
<tr>
<td>Hydrangea</td>
<td>Contact allergy</td>
</tr>
<tr>
<td>Plant Name</td>
<td>Effect</td>
</tr>
<tr>
<td>-------------------------</td>
<td>---------------------------------</td>
</tr>
<tr>
<td>Hydrocotyle</td>
<td>Photosensitivity</td>
</tr>
<tr>
<td>Ispaghula</td>
<td>Rare cases of allergy</td>
</tr>
<tr>
<td>Juniper</td>
<td>Contact allergy</td>
</tr>
<tr>
<td>Lady’s Slipper</td>
<td>Contact allergy</td>
</tr>
<tr>
<td>Meadowsweet</td>
<td>Potentiation of histamine bronchospastic properties</td>
</tr>
<tr>
<td>Motherwort</td>
<td>Dermatitis, photosensitisation</td>
</tr>
<tr>
<td>Parsley</td>
<td>Furanocoumarins, photosensitivity</td>
</tr>
<tr>
<td>Pilewort</td>
<td>Contact allergy, protanemonin</td>
</tr>
<tr>
<td>Plantain</td>
<td>Contact allergy</td>
</tr>
<tr>
<td>Pleurisy Root</td>
<td>Contact allergy</td>
</tr>
<tr>
<td>Pulsatilla</td>
<td>Contact allergy, protoanemonin</td>
</tr>
<tr>
<td>Rosemary</td>
<td>Dermatitis, photosensitisation</td>
</tr>
<tr>
<td>St. John’s Wort</td>
<td>Photodermatitis, hypericin</td>
</tr>
<tr>
<td>Tansy</td>
<td>Sesquiterpene lactone constituents</td>
</tr>
<tr>
<td>Wild Carrot</td>
<td>Furanocoumarins, photosensitivity</td>
</tr>
<tr>
<td>Drug</td>
<td>Effects</td>
</tr>
<tr>
<td>-----------------</td>
<td>--------------------------------------------------</td>
</tr>
<tr>
<td>Alfalfa</td>
<td>Irritant, canavanine in seeds</td>
</tr>
<tr>
<td>Arnica</td>
<td>Irritant to mucous membranes</td>
</tr>
<tr>
<td>Asafoetida</td>
<td>Irritant gum</td>
</tr>
<tr>
<td>Blue Flag</td>
<td>Irritant gum and oil</td>
</tr>
<tr>
<td>Bogbean</td>
<td>Irritant to GI tract</td>
</tr>
<tr>
<td>Boldo</td>
<td>Irritant oil</td>
</tr>
<tr>
<td>Buchu</td>
<td>Irritant oil</td>
</tr>
<tr>
<td>Capsicum</td>
<td>Capsaicinoids, mucosal irritants</td>
</tr>
<tr>
<td>Cassia</td>
<td>Irritant to mucous membranes, oil</td>
</tr>
<tr>
<td>Cinnamon</td>
<td>Irritant to mucous membranes, oil</td>
</tr>
<tr>
<td>Cohosh, Blue</td>
<td>Irritant to mucous membranes; spasmogenic in vitro</td>
</tr>
<tr>
<td>Cowslip</td>
<td>Irritant saponins</td>
</tr>
<tr>
<td>Drosera</td>
<td>Plumbagin, irritant</td>
</tr>
<tr>
<td>Eucalyptus</td>
<td>Irritant oil</td>
</tr>
</tbody>
</table>
False Unicorn  
Large doses may cause vomiting

Figwort  
Purgative effect

Garlic  
Raw clove

Ground Ivy  
Irritant oil

Guaiacum  
Avoid if inflammatory condition

Horse-chestnut  
Saponin constituents, contra–indicated in existing renal disease

Horseradish  
Irritant oil

Hydrangea  
May cause gastro–enteritis, hydrangin

Jamaica Dogwood  
Irritant to humans

Juniper  
Irritant oil

Lemon Verbena  
Irritant oil

Lime Flower  
Irritant to kidney, oil

Nettle  
Tea irritant to stomach

Parsley  
Irritant oil

Pennyroyal  
Toxic and irritant oil
<table>
<thead>
<tr>
<th>Plant</th>
<th>Effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pilewort</td>
<td>Irritant sap</td>
</tr>
<tr>
<td>Pleurisy Root</td>
<td>Gastrointestinal irritant</td>
</tr>
<tr>
<td>Pokeroot</td>
<td>Irritant saponins</td>
</tr>
<tr>
<td>Pulsatilla</td>
<td>Irritant to mucous membranes</td>
</tr>
<tr>
<td>Queen’s Delight</td>
<td>Diterpene constituents</td>
</tr>
<tr>
<td>Sage</td>
<td>Irritant oil</td>
</tr>
<tr>
<td>Sarsparilla</td>
<td>Saponins</td>
</tr>
<tr>
<td>Senega</td>
<td>Saponins</td>
</tr>
<tr>
<td>Skunk Cabbage</td>
<td>Inflammatory and blistering to skin</td>
</tr>
<tr>
<td>Squill</td>
<td>Saponins</td>
</tr>
<tr>
<td>Thyme</td>
<td>Irritant oil</td>
</tr>
<tr>
<td>Willow</td>
<td>Salicylates</td>
</tr>
<tr>
<td>Witch Hazel</td>
<td>Irritant to stomach</td>
</tr>
<tr>
<td>Drug</td>
<td>Effects</td>
</tr>
<tr>
<td>-----------------</td>
<td>------------------------------</td>
</tr>
<tr>
<td>Agnus Castus</td>
<td>Alkaloids</td>
</tr>
<tr>
<td>Alfalfa</td>
<td>Alkaloids</td>
</tr>
<tr>
<td>Aniseed</td>
<td>Anethole, sympathomimetic</td>
</tr>
<tr>
<td>Arnica</td>
<td>Betaines, choline</td>
</tr>
<tr>
<td>Bloodroot</td>
<td>Alkaloids</td>
</tr>
<tr>
<td>Bogbean</td>
<td>Alkaloids</td>
</tr>
<tr>
<td>Boldo</td>
<td>Alkaloids</td>
</tr>
<tr>
<td>Borage</td>
<td>Alkaloids</td>
</tr>
<tr>
<td>Broom</td>
<td>Alkaloids, amines</td>
</tr>
<tr>
<td>Calamus</td>
<td>Amines</td>
</tr>
<tr>
<td>Capsicum</td>
<td>Sympathomimetic</td>
</tr>
<tr>
<td>Cat’s Claw</td>
<td>Alkaloids</td>
</tr>
<tr>
<td>Centaury</td>
<td>Alkaloids</td>
</tr>
</tbody>
</table>
Cereus Tyramine
Cohosh, Black Alkaloids
Cohosh, Blue Alkaloids
Cola Alkaloids
Coltsfoot Alkaloids
Comfrey Alkaloids
Corn Silk Amines
Echinacea Alkaloids
Ephedra Alkaloids
Eyebright Alkaloids
Fenugreek Choline, trigonelline
Fumitory Alkaloids
Gentian Alkaloids
Ginkgo Seed: alkaloids
Leaf: MAOI activity
Ginseng, Panax MAOI potentiation, suspected phenelzine interaction
Golden Seal  Alkaloids
Gravel Root  Alkaloids
Hawthorn  Tyramine
Horehound, White  Alkaloids
Hydrocotyle  Alkaloids
Ispaghula  Alkaloids
Jamaica Dogwood  Alkaloids
Liferoot  Alkaloids
Lobelia  Alkaloids
Maté  Alkaloids, amines
Mistletoe  Histamine release
Motherwort  Alkaloids
Nettle  Choline
Parsley  Myristicin, sympathomimetic
Passionflower  Alkaloids (traces or absent)
Plantain  Alkaloids
Pleurisy Root       Sympathomimetic

Pokeroot          Betalains

Prickly Ash, Northern Alkaloids

Prickly Ash, Southern Alkaloids

Quassia           Alkaloids

Sassafras         Alkaloids

Shepherd’s Purse  Choline, tyramine

Skunk Cabbage     Alkaloids

St. John’s Wort   MAOI activity, in vitro

Stone Root        Alkaloids

Valerian          Alkaloids

Vervain           Sympathomimetic

Yarrow            Betonicine, stachydrine, betaine
Aloe Vera, Angelica, Arnica, Bilberry, Bloodroot, Blue Flag, Boldo, Boneset, Borage, Buchu, Calendula, Cassia, Cat’s Claw, Centaury, Chamomile (German), Chamomile (Roman), Cohosh (Black), Coltsfoot, Comfrey, Cowslip, Dandelion, Devil’s Claw, Echinacea, Elder, Ephedra, Evening Primrose, Feverfew, Figwort, Gentian, Ginger, Ground Ivy, Horse-chestnut, Hydrocotyle, Juniper, Liquorice, Milk Thistle, Myrrh, Nettle, Plantain, Pokeroot, Prickly Ash (Southern), Rosemary, Sarsaparilla, Shepherd’s Purse, Uva-Ursi, Willow, Yarrow, Yucca
Appendix 16 Antispasmodic Herbal Ingredients

Angelica, Aniseed, Asafoetida, Calendula, Capsicum, Cassia, Celery, Cohosh (Blue), Cinnamon, Clove, Cowslip, Echinacea, Elecampane, Euphorbia, Chamomile (German), Hops, Jamaica Dogwood, Lime Flower, Raspberry, Chamomile (Roman), Rosemary, Sage, Scullcap, Tansy, Thyme, Valerian

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Appendix 17 Herbal Ingredients containing Coumarins

Alfalfa, Angelica, Aniseed, Arnica, Asafoetida, Bogbean, Boldo, Buchu, Capsicum, Cassia, Celery, Chamomile (German), Chamomile (Roman), Dandelion, Fenugreek, Horse-chestnut, Horseradish, Liquorice, Meadowsweet, Nettle, Parsley, Passionflower, Prickly Ash (Northern), Quassia, Wild Carrot, Wild Lettuce

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Appendix 18 Herbal Ingredients containing Flavonoids

Agnus Castus, Agrimony, Angelica, Aniseed, Apricot, Arnica, Artichoke, Bayberry, Bilberry, Bogbean, Boldo, Boneset, Broom, Buchu, Burdock, Burnet, Calendula, Celery, Cereus, Chamomile (German), Chamomile (Roman), Chaparral, Clivers, Coltsfoot, Corn Silk, Couchgrass, Cowslip, Damiana, Dandelion, Devil’s Claw, Drosera, Elder, Ephedra, Eucalyptus, Euphorbia, Eyebright, Fenugreek, Feverfew, Figwort, Frangula, Fumitory, Ginkgo, Gravel Root, Ground Ivy, Hawthorn, Hops, Horehound (Black), Horehound (White), Horse-chestnut, Hydrangea, Hydrocotyle, Java Tea, Juniper, Lemon Verbena, Lime Flower, Liquorice, Marshmallow, Maté, Meadowsweet, Melissa, Milk thistle, Mistletoe, Motherwort, Nettle, Parsley, Passionflower, Plantain, Pulsatilla, Raspberry, Red Clover, Rhubarb, Rosemary, Sage, Sarsaparilla, Saw Palmetto, Scullcap, Senna, Shepherd’s Purse, Squill, St. John’s Wort, Thyme, Uva-Ursi, Wild Carrot, Wild Lettuce, Willow, Witch Hazel, Yarrow
Appendix 19 Herbal Ingredients containing Iridoids

Agnus Castus, Bilberry, Bogbean, Centaury, Clivers, Devil’s Claw, Eyebright, Figwort, Gentian, Ispaghula, Motherwort, Plantain, Scullcap, Uva-Ursi, Valerian, Vervain

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Appendix 20 Herbal Ingredients containing Saponins

Alfalfa, Aloe Vera, Bogbean, Burnet, Calendula, Chaparral, Cohosh (Blue), Corn Silk, Cowslip, False Unicorn, Fenugreek, Ginseng (Eleutherococcus), Ginseng (Panax), Hawthorn, Horehound (White), Horse-chestnut, Hydrangea, Hydrocotyle, Jamaica Dogwood, Lime Flower, Milk Thistle, Pokerooot, Pulsatilla, Red Clover, Sarsaparilla, Senega, Senna, Stone Root, Thyme, Witch Hazel, Yucca
Appendix 21 Herbal Ingredients containing Tannins

Agrimony, Apricot, Arnica, Artichoke, Avens, Bayberry, Bilberry, Blue Flag, Boldo, Borage, Burnet, Calamus, Cascara, Cassia, Chamomile (German), Cinnamon, Clivers, Cohosh (Black), Cola, Coltsfoot, Comfrey, Corn Silk, Cowslip, Damiana, Drosera, Elder, Ephedra, Eucalyptus, Eyebright, Feverfew, Frangula, Gentian, Ground Ivy, Hawthorn, Holy Thistle, Hops, Horsechestnut, Ispaghula, Juniper, Lady’s Slipper, Lime Flower, Marshmallow, Meadowsweet, Mistletoe, Motherwort, Nettle, Pilewort, Plantain, Poplar, Prickly Ash (Northern), Prickly Ash (Southern), Queen’s Delight, Raspberry, Rhubarb, Sage, Sassafras, Saw Palmetto, Scullcap, Slippery Elm, Squill, St. John’s Wort, Stone Root, Tansy, Thyme, Uva-Ursi, Valerian, Vervain, Willow, Witch Hazel, Yarrow, Yellow Dock

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Appendix 22 Herbal Ingredients containing Volatile Oils

Agnus Castus, Agrimony, Angelica, Aniseed, Arnica, Artichoke, Asafoetida, Avens, Blue Flag, Boldo, Boneset, Buchu, Burdock, Burnet, Calamus, Calendula, Capsicum, Cassia, Celery, Chamomile (German), Chamomile (Roman), Chaparral, Cinnamon, Clove, Cohosh (Black), Coltsfoot, Couchgrass, Damiana, Ephedra, Elder, Elecampane, Eucalyptus, Eyebright, Feverfew, Garlic, Gentian, Ginger, Ginseng (Eleutherococcus), Ginseng (Panax), Golden Seal, Ground Ivy, Holy Thistle, Hops, Horehound (Black), Horseradish, Hydrocotyle, Java Tea, Juniper, Lemon Verbena, Lime Flower, Liquorice, Lobelia, Meadowsweet, Melissa, Motherwort, Myrrh, Parsley, Pennyroyal, Prickly Ash (Northern), Queen’s Delight, Red Clover, Rosemary, Sage, Sassafras, Saw Palmetto, Senna (trace), Skunk Cabbage, Squill, St. John’s Wort, Stone Root, Tansy, Thyme, Uva-Ursi, Valerian, Wild Carrot, Witch Hazel, Yarrow

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Category Natural source of flavouring

Plants, animals and other organisms, and parts of these or products thereof, normally consumed as food items, herbs or spices in Europe for which it is considered that there should be no restrictions on use.

1 Flavouring preparations, which are not themselves consumed as food but which are derived from plants, animals and other organisms, and parts of these or products thereof, normally consumed as food items, herbs or spices in Europe. These preparations, on the basis of the information available, are not considered a risk to health in the quantities used.

Plants, animals and other organisms, and parts of these or products thereof, and preparations derived therefrom, not normally consumed as food items, herbs or spices in Europe.

2 These source materials and preparations, on the basis of the information available, are not considered to constitute a risk to health in the quantities used.

Plants, animals and other organisms, and parts of these or products thereof, normally consumed as food items, herbs or spices in Europe which contain defined ‘active principles’ or ‘other chemical components’ requiring limits on use levels.

Flavouring preparations, which are not themselves consumed as food but which are derived from plants, animals and other organisms, and parts of these or products thereof, normally consumed as food items, herbs or spices in Europe which contain defined ‘active principles’ or ‘other chemical components’ requiring limits on use levels.

These source materials and preparations are not considered to constitute a risk to health in the quantities used provided that the
Plants, animals and other organisms, and parts of these or products thereof, and preparations derived therefrom, not normally consumed as food items, herbs or spices in Europe which contain defined ‘active principles’ or ‘other chemical components’ requiring limits on use levels.

These source materials and preparations are not considered to constitute a risk to health in the quantities used provided that the limits set for the ‘active principles’ or ‘other chemical components’ are not exceeded.

Plants, animals and other organisms, and parts of these or products thereof, and preparations derived therefrom, for which additional toxicological and/or chemical information is required.

These could temporarily be acceptable provided that any limits set for the ‘active principles’ or the ‘other chemical components’ are not exceeded.

Plants, animals and other organisms, and parts of these or products thereof, and preparations derived therefrom, which are considered to be unfit for human consumption in any amount.

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Advanced search

Advanced search allows you to perform a text search in specific sections of the text.

When you perform an advanced search, the results display in the main document area, showing how your search has been handled.

Use the tick boxes to choose which sections you want to search.

See related:

Examples of searches

Entering a search
Help contents > Examples of searches
Examples of searches

Here are some examples of how to find answers to common questions

- How do I do a full text search?
About Herbal Medicines

Herbal Medicines is an electronic database developed from the book of the same name, first published by Pharmaceutical Press in 1986. It contains monographs of over 100 herbal medicines, with information on the quality, safety, efficacy and legal requirements. In addition, there is a section on the chemical constituents of plants used as herbal medicines, and a list of general references.
Help contents > How to cite electronic *Herbal Medicines*
How to cite electronic *Herbal Medicines*

When you cite *Herbal Medicines*, we recommend the following style:


This citation should always be added to copies of documents printed from *Herbal Medicines*. 
Structure of *Herbal Medicines*

There are two main types of information in *Herbal Medicines*:

- monographs
- appendices

You can read more about these in the Overview, Notes and Introduction.
The **Herbal Medicines** interface

All the basic functions for searching and viewing *Herbal Medicines* are displayed on a single screen.

The screen is divided into 2 main parts:

- **the top bar**
- **the document area**
The top bar

The **top bar** is used to display:

- Function buttons:
  - [Home](#)
  - [Contents](#)
  - [Advanced search](#)
  - [Print](#)
  - [Help](#)
- The [search box](#)
The document area

Underneath the top bar is the document area which is used to display:

- documents - pages from Herbal Medicines
- the table of contents
- text search results
Top bar

The top bar is used to display the **Function buttons and search box**.
Function buttons and search box

There are 5 function buttons on the top bar:

- **Home**
  To go to the home page, click on this button.

- **Contents**
  To show the table of contents, click on this button.

- **Advanced search**
  To perform an advanced text search click on this button.

- **Print**
  To print the current page, click on this button.

- **Help**
  To show help information, click on this button.

The **search box** allows you to search *Herbal Medicines* directly. Use it to find words or phrases anywhere in the text.

See related:

**Entering a search**
Help contents > Top bar > Function buttons and search box > Search Button
Search Button

Click on the "Search" button to perform a text search when you have entered your search terms in the search box.

If you click on the "Search" button when the search box is empty, information about how to use text search is displayed.

See related:

Search box

Entering a search
Contents list

The contents list shows the hierarchical arrangement of documents in *Herbal Medicines*. You can use the contents list to view the contents of any monograph or appendix and select documents to view.

To view the contents list, click on the **Contents button** on the top bar

*Herbal Medicines* is organised into 3 sections. To locate a document:

- expand the contents list by clicking on any ⬇️ symbol.
- continue to expand the list in the same way until you find a document you wish to view
- click on the title to display the document

or

- Click on the section title to display a detailed contents list
- follow links under the "Sub-sections" heading to display individual monographs

Click on a ⬇️ symbol to collapse the table to that level.

Notes:

Some browsers are unable to expand the table of contents levels. In this case, click on the section you require in the table of contents, and use the list of sub-sections in the main document area.
Text search results display

When you perform a text search, the results display in the main document area, showing how your search has been handled.

The display includes:

- An explanation of what has been searched
- the number of matching documents
- the list of documents retrieved
- a link to Related articles

The list of documents includes the titles of the documents and their locations. This will help you select the most relevant records to view.

See related:

Examples of searches

Entering a search
Help contents > How to find information
How to find information

There are two alternative ways of finding information in *Herbal Medicines*:

- **searching for a term**
- **browsing using the contents list**

See related:

**Examples of searches**
How to browse using the contents list

The contents list (sometimes referred to as the menu) gives a hierarchical view of the structure (see Structure of Herbal Medicines).

- Expand the contents list by clicking on any symbol
- Continue to expand the list in the same way until you find a document you wish to view
- Click on the link or the symbol to display the document.
Advanced searching features

You can refine your search by combining search terms using logical operators.
Advanced text searching

ENTER AS  EFFECT

heart failure   documents containing either (or both) of the words
+heart +failure documents containing both the words
liver heart +failure documents containing liver and/or heart which must also contain failure
"heart failure" documents containing both words as a phrase
cat -dog documents containing the first word but not the second

Notes:
Move the cursor over any section titles in Search in... to display the full title of the section or record type.
You can select several boxes in Search in... and the search will retrieve any of those sections containing the search term.
If you do not select any sections, the system defaults to searching all document types.
How to view the documents retrieved by a search

The results of a search will be displayed when you click the **Search** button after entering search terms.

Select the documents you want to see from the list and click on the title to display.

The search terms ('hits'), including any synonyms matched, are highlighted. If two or more search terms have been entered, a different colour is used to highlight different hits or families of hits.

To return to the list of search results and select a different document use the Return to search link in the **breadcrumb trail** or the **Back** button on the browser.

See related:

**Search results display**

**Document display**
Navigation

- To move around *Herbal Medicines*
  - from the **breadcrumb trail**
  - from the **contents list**

- To move around a document
  - from the **document area**
  - from the **contents list**
  - using **links**

- To move from one document to another
  - **within Herbal Medicines**
  - **to documents on the internet**
How to move around *Herbal Medicines*

A set of links is provided at the top of each screen to help you navigate around *Herbal Medicines*.

The top line links to the next and previous document in the hierarchy

« Next | Previous »

Leave the cursor over the link for a moment to see the title of the document.

If the screen contains a document displayed as the result of a search, the top line will also show the links

Return to search and Hide highlighting

- Use Return to search to return to your list of search results
- Use Hide highlighting to see the document without highlighting when you have finished with the search

The next line shows the location of the document within the hierarchy. This is known as a 'breadcrumb trail'.

Home > Contents > Monographs > C

Each entry in the breadcrumb trail acts as a link to a more general document, sub-section, or contents list.

**Notes:**

Next | Previous links are also available at the bottom of the screen.

Once you choose Hide highlighting, both that link and Return to search link will disappear. If you then wish to check the results of your last search, use the Back button on your browser to return to the search results list. Selecting a document will highlight the search terms as usual.

The first link in the breadcrumb trail is always Home and will take you to the
MedicinesComplete home page.

See related:

Document display
How to use links

Active links (sometimes referred to as 'hyperlinks') are indicated by underlined text.

- position the mouse over the link text (the cursor will usually change to a pointing hand)
- click on the link
- the linked document will be displayed.

Notes:
To return to a previous screen, use the 'Back' button on your browser toolbar

See related:
Links between documents within Herbal Medicines
Links to external sources
Links to external sources

Links are included to external documents available via the Internet.

If you have web access, external documents can be retrieved by clicking on the link in the same way as internal links.

External documents will appear in a separate browser window. To return to Herbal Medicines close or minimise the new window.

The main links to external sources currently available are

- links from reference citations to Medline records (from PubMed)
- links to relevant websites (on-line information)
- in the Contact us section, links to email addresses (in this case, your e-mail authoring software will be launched and the address inserted, ready for your message).

See related:

Using links

Links between documents within Herbal Medicines
Reference citation lists

References cited in monographs are linked by the first author's name and publication date to the full citations listed at the end of each monograph.

Notes:

Citations listed at the end of individual records include a link to a Medline abstract in PubMed where possible.
St. John’s Wort
Species (Family)

*Hypericum perforatum* L. (Hypericaceae)
**Synonym(s)**

Hypericum, Millepertuis
Part(s) Used

Herb
Pharmacopoeial and Other Monographs

American Herbal Pharmacopoeia\(^{(G1)}\)

BHP 1996\(^{(G9)}\)

BP 2002\(^{(G71)}\)

Complete German Commission E\(^{(G3)}\)

ESCOP 1996\(^{(G52)}\)

Martindale 33rd edition\(^{(G67)}\)

Mills and Bone\(^{(G50)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)

Ph Eur 2004\(^{(G72)}\)

USP26/NF21\(^{(G73)}\)
Legal Category (Licensed Products)

GSL (for external use only)\(^{(G37)}\)
Constituents
See Reference 1 and General References G1 G2 G22 G40 G48 G52 G62 G64.

**Anthraquinone derivatives (naphthodianthrones)**
Hypericin, pseudohypericin and isohypericin; protohypericin and protopseudohypericin (biosynthetic precursors of hypericin and pseudohypericin, respectively) are present in fresh material. Cyclopseudohypericin is also stated to be present. The hypericin content (approximately 0.1–0.15%) includes both hypericin and pseudohypericin and is sometimes referred to as ‘total hypericins’.

**Flavonoids**
Flavonols (e.g. kaempferol, quercetin), flavones (e.g. luteolin) and glycosides (e.g. hyperoside, isoquercitrin, quercitrin, rutin), biflavonoids including biapigenin (a flavone) and amentoflavone (a biapigenin derivative) and catechins (flavonoids often associated with condensed tannins). The concentrations of rutin, hyperoside and isoquercitrin have been reported as 1.6, 0.9 and 0.3%, respectively.

**Prenylated phloroglucinols**
Hyperforin (2.0–4.5%) and adhyperforin (0.2–1.9%).

**Tannins**
8–9%. Type not specified. Proanthocyanidins (condensed type) have been reported.

**Other phenols**
Caffeic, chlorogenic, p-coumaric, ferulic, p-hydroxybenzoic and vanillic acids.

**Volatile oils**
0.05–0.9%. Major component (not less than 30%) is methyl-2-octane (saturated hydrocarbon); others include n-nonane and traces of methyl-2-decane and n-undecane (saturated hydrocarbons), α- and β-pinene, α-terpineol, geraniol, and traces of myrcene and limonene (monoterpenes), caryophyllene and humulene (sesquiterpenes).

**Other constituents**
Acids (isovalerianic, nicotinic, myristic, palmitic, stearic), carotenoids, choline, nicotinamide, pectin, β-sitosterol, straight-chain saturated hydrocarbons ($C_{16}$, $C_{30}$) and alcohols ($C_{24}$, $C_{26}$, $C_{28}$).
Food Use

St. John’s wort is listed by the Council of Europe as a natural source of food flavouring (herb: category 5) (see Appendix 23).\(^{(G17)}\)
Herbal Use

St. John’s wort is stated to possess sedative and astringent properties. It has been used for excitability, neuralgia, fibrositis, sciatica, wounds, menopausal neurosis, anxiety and depression and as a nerve tonic. St. John’s wort is used extensively in homeopathic preparations as well as in herbal products. Modern interest is focused on its use as an antidepressant.
Dosage

**Dried herb**
2–4 g or by infusion three times daily.\(^{(G7)}\)

**Liquid extract**
2–4 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)

**Tincture**
2–4 mL (1 : 10 in 45% alcohol) three times daily.\(^{(G7)}\)

The doses of St. John’s wort extract used in clinical trials involving patients with mild to moderate depression generally range from 350 to 1800 mg daily (equivalent to 0.4 to 2.7 mg hypericin daily, depending on the extract).\(^{(14)}\)
Pharmacological Actions

The major active constituents are considered to be hyperforin (a prenylated phloroglucinol) and hypericin (a naphthodianthrone), although other biologically active constituents, e.g. flavonoids and tannins, are also present.\(^{(15)}\) Several pharmacological activities, including antidepressant, antiviral and antibacterial effects, have been documented for extracts of St. John’s wort and/or its constituents. The pharmacology and pharmacodynamics of St. John’s wort have been reviewed.\(^{(1,16,G1G5G55)}\)

**In vitro and animal studies**

**Antidepressant activity**

The precise mechanism of action for the antidepressant effect of St. John’s wort is unclear. Initially, attention was focused on hypericin as the constituent of St. John’s wort believed to be responsible for the herb’s antidepressant effects. Inhibition of monoamine oxidase (MAO) type A and B in rat brain mitochondria *in vitro* was described for hypericin.\(^{(17)}\) However, other studies have demonstrated only weak or no MAO inhibition.\(^{(18–20)}\)

*In vitro* receptor binding and enzyme inhibition assays carried out using hypericum extract demonstrated significant receptor affinity for adenosine, GABA\(_A\), GABA\(_B\), benzodiazepine and MAO types A and B, although, with the exception of GABA\(_A\) and GABA\(_B\), the concentrations of hypericum required were unlikely to be attained after oral administration in humans.\(^{(21)}\) Other biochemical studies have reported that the hypericum extract LI 160 is only a weak inhibitor of MAO-A and MAO-B activity, but that it inhibits the synaptosomal uptake of serotonin (5-hydroxytryptamine or 5-HT), dopamine and noradrenaline (norepinephrine) with approximately equal affinity and also leads to a downregulation of β-receptors and an upregulation of 5-HT\(_2\) receptors in the rat frontal cortex.\(^{(22)}\) The effects of fluoxetine and hypericin- and flavonoid–standardised hypericum extracts (LI 160, 0.3% hypericin and 6% flavonoids and Ph–50, 0.3% hypericin and 50% flavonoids) on the concentrations of neurotransmitters in brain regions were studied in rats.\(^{(23)}\) All three preparations induced a significant increase in 5-HT concentrations in the rat cortex, both LI 160 and Ph–50 caused increases in noradrenaline (norepinephrine) and dopamine in the rat diencephalon and Ph–50 also induced an increase in the noradrenaline (norepinephrine) content in the brainstem, areas that are implicated in depression.\(^{(23)}\) In studies using the rat forced swimming test, an experimental model of depression, hypericum extracts induced a significant reduction in immobility.\(^{(24)}\)
Hyperforin has now emerged as being one of the major active constituents of importance in antidepressant activity. Hyperforin has been shown to be an uptake inhibitor of 5-HT, dopamine, noradrenaline (norepinephrine), GABA and L-glutamate in synaptosomal preparations\(^\text{25}\) and to inhibit 5-HT uptake in rat peritoneal cells in a dose-dependent manner.\(^\text{26}\) Studies have also described discrepancies between observed and theoretical IC\(_{50}\) values, indicating that hyperforin is not the only component of hypericum extract that is responsible for the observed effects.\(^\text{26,27}\) It has been reported that the mode of action of hyperforin in serotonin uptake inhibition seems to be associated with the elevation of free intracellular sodium ion concentrations\(^\text{28}\) and that this may be secondary to activation of the Na\(^+\)/H\(^+\) exchange as a result of a decrease in intracellular pH.\(^\text{29}\) Hyperforin was shown to inhibit 5-HT reuptake in washed platelets but not in fresh platelet-rich plasma, suggesting that plasma–protein binding could be a limiting factor for 5-HT uptake inhibition in vivo.\(^\text{30}\)

A commercial extract of St. John’s wort has exhibited psychotropic and antidepressant activities in mice.\(^\text{31}\) Pure hyperforin and hypericum extracts also demonstrated antidepressant activity in a despair behaviour test in rats.\(^\text{26}\)

In other experimental models of depression, including acute and chronic forms of escape deficit induced by stressors, hypericum extract was shown to protect rats from the consequences of unavoidable stress.\(^\text{32}\) Flavonoid fractions and flavonoids isolated from these fractions have been reported to have antidepressant activity in experimental studies (forced swimming test) in rats.\(^\text{33}\)

**Antimicrobial activity**
A leaf extract has been documented as enhancing the immunity of mice towards *Staphylococcus aureus* and *Bordetella pertussis*;\(^\text{34}\) hyperforin is reported to be antibacterial with activity against *S. aureus*.\(^\text{8}\) Antibacterial activity of hyperforin against multiresistant *S. aureus* and Gram-positive bacteria, including *Streptococcus pyogenes* and *Corynebacterium diphtheriae*, has been reported.\(^\text{35}\) However, it has been emphasised that the antibacterial effects of hyperforin are only observed at high concentrations.\(^\text{36,37}\) Hyperforin did not exhibit any growth inhibitory effect against Gram-negative bacteria, such as *Enterococcus faecalis*, *Escherichia coli* and *Pseudomonas aeruginosa* or against *Candida albicans*.\(^\text{35}\) Further antibiotic constituents have been isolated from St. John’s wort: imanine and novoimanine.\(^\text{38,39}\) Novoimanine was reported to be the most effective topical agent against *S. aureus*.\(^\text{38}\) Herb extracts are reported to exhibit more pronounced activity
against staphylococci, shigellae and *E. coli* than are decoctions.\(^{(39,40)}\)

**Antiviral activity**

Flavonoid and catechin–containing fractions have exhibited antiviral activity, inhibiting the influenza virus by 83–100\%.\(^{(41)}\) Hypericin and pseudohypericin have been reported to inhibit several encapsulated viruses *in vitro*, including herpes simplex types 1 and 2\(^{(42,43)}\) and human immunodeficiency virus type 1 (HIV-1).\(^{(44–47)}\) Hypericin has also been reported to inactivate murine cytomegalovirus (MCMV) and Sindbis virus.\(^{(47)}\) The antiviral activity of hypericin appears to involve a photoactivation process.\(^{(47,G1)}\)

**Other effects**

*In vitro* studies using a hamster vas deferens smooth muscle cell line demonstrated that hyperforin induces the release of calcium ions from mitochondrial or other sources followed by activation of cellular metabolism.\(^{(48)}\) It is not known whether this activity contributes to the antidepressant effects of hyperforin.

Oral administration of a single dose of St. John’s wort (100, 200, 400, 600 or 800 mg/kg) to two strains of alcohol–preferring rats significantly reduced alcohol intake in both strains.\(^{(49)}\) In another study in experimental alcoholism, acute intraperitoneal administration of St. John’s wort (10–40 mg/kg), fluoxetine (1–10 mg/kg) and imipramine (3–30 mg/kg) reduced alcohol intake in a dose–dependent manner in a 12–hour, limited access, two–bottle choice (ethanol/water) procedure.\(^{(24)}\) Depression and alcoholism are thought to have some neurochemical similarities, such as low brain serotonin concentrations.\(^{(50)}\)

It has been suggested that biflavonoids may be the sedative principles in St. John’s wort since CNS activity has been documented for biflavonoid constituents in another plant, *Taxus baccata*.\(^{(3)}\)

An extract of St. John’s wort was found to suppress inflammation and leukocyte infiltration induced by carrageenan and prostaglandin E\(_1\) (PGE\(_1\)) in mice.\(^{(51)}\) *In vitro*, hypericin has been shown to inhibit tumour necrosis factor–induced activation of the transcription factor NF-κB,\(^{(52)}\) specific growth factor–regulated protein kinases\(^{(53–55)}\) and the release of arachidonic acid and leukotriene B\(_4\).\(^{(56)}\) In a rabbit model of proliferative vitreoretinopathy (PVR), intravitreal injection of hypericin 0.1 mL (10 or 100 μmol/L, but not 1 μmol/L) inhibited the progression of PVR when compared with severity in control eyes five days after hypericin administration.\(^{(57)}\) It was suggested that, as protein kinase C is important in the cellular reactions occurring in
PVR, modulation of protein kinase C by hypericin may be a factor in this system. Hypericin and pseudohypericin have been reported to inhibit 12–lipoxygenase activity; the products of lipoxygenase–catalysed reactions, such as leukotrienes, may be involved in inflammatory reactions.\(^{(58)}\)

Other compounds may contribute to the anti–inflammatory properties of St. John’s wort.\(^{(37)}\) Anti–inflammatory and anti–ulcerogenic properties have been documented for amentoflavone, a biapigenin derivative.\(^{(4)}\) Analgesic activity in mice has been reported for the total flavonoid fraction;\(^{(59)}\) the active principle was stated to be of the quercetin type.

Both water–soluble imanine and imanine were reported to reduce blood pressure and increase the frequency and depth of breathing following intravenous administration (50 mg/kg) to rabbits.\(^{(38)}\) A study of the vasoconstrictor action of water–soluble imanine and imanine on the isolated rabbit ear indicated that their hypotensive action was not due to a direct effect on the vasculature.\(^{(38)}\) When perfused through the isolated frog heart, both water–soluble imanine and imanine were found to cause cardiac systolic arrest at a dilution of $1 \times 10^{-5}$.\(^{(38)}\) Proanthocyanidin–containing fractions isolated from St. John’s wort have been reported to inhibit contractions of the isolated guinea–pig heart induced by histamine, PGF\(_{2\alpha}\) and potassium chloride.\(^{(60)}\)

A tonus–raising effect on isolated guinea–pig and rabbit uteri has been documented for a crude aqueous extract.\(^{(61)}\) Of the group of plants investigated, St. John’s wort was reported to exhibit the weakest uterotonic activity.

Tannins isolated from St. John’s wort are stated to have mild astringent activity.\(^{(62)}\) The anthraquinone derivatives documented for St. John’s wort do not possess any purgative action.\(^{(62)}\)

*In vitro* cytotoxicity against human colon carcinoma cells (CO 115) has been described for hyperforin–related constituents isolated from *Hypericum calycinum* and *Hypericum revolutum*.\(^{(63)}\)

**Clinical studies**

Clinical trials with extracts of St. John’s wort have focused mainly on its effects in patients with depression, although there have been several studies exploring its use in other conditions, including seasonal affective disorder, chronic fatigue and premenstrual syndrome.

**Depression**
Initially, hypericin was thought to be responsible for the antidepressant activity of St. John’s wort, although, more recently, experimental\(^{25,26}\) and clinical evidence\(^6^{4}\) has emerged to indicate that hyperforin is one of the major constituents required for antidepressant activity.

The precise mechanism of action of St. John’s wort’s antidepressant effect remains unclear (see Pharmacological Actions, \textit{In vitro} and animal studies). A double-blind, placebo-controlled, crossover study in 12 healthy male volunteers investigated the effects of a single dose of St. John’s wort extract (LI 160) (2700 mg, 9 × 300-mg tablets standardised to 0.3% hypericin) on plasma concentrations of growth hormone, prolactin and cortisol.\(^{65}\) A significant increase in plasma growth hormone concentration and a significant decrease in plasma prolactin concentration were observed following St. John’s wort administration relative to placebo administration. Plasma cortisol concentrations were unchanged. These findings suggest that this dose of St. John’s wort extract may increase aspects of brain dopamine function in humans, although further studies are required to confirm this, assess dose-response relationships and determine whether there is evidence for effects on dopaminergic systems in patients with depression treated with St. John’s wort.\(^{65}\) Another study, which used a randomised, three-way, crossover design, investigated the effects of a single dose of St. John’s wort extract (LI 160S) (600 or 300 mg) or placebo on hormone concentrations in 12 healthy male volunteers.\(^{66}\) Compared with placebo, St. John’s wort extract (600 mg) increased cortisol secretion between 30 and 90 minutes after dosing, indicating an influence of St. John’s wort on certain CNS neurotransmitters. There was no difference between the three groups with regard to adrenocorticotropic hormone (ACTH), growth hormone and prolactin secretion.\(^{66}\)

A systematic review and meta-analysis of randomised controlled trials of preparations of St. John’s wort extract included 23 trials involving a total of 1757 patients with depressive disorders.\(^{67}\) This has been updated to include new studies and published as a Cochrane review of 27 randomised controlled trials of St. John’s wort extract in patients with ‘neurotic depression’ and mild to moderately severe depressive disorders.\(^{14}\) Seventeen of these trials (involving 1168 patients) compared St. John’s wort preparations with placebo (16 studies used preparations containing St. John’s wort extract as the sole herbal ingredient and one involved a combination product of St. John’s wort extract with four other herbal ingredients); the ten other trials (involving 1123 patients) compared St. John’s wort extracts with conventional antidepressant or sedative drugs, including amitriptyline, imipramine, desipramine and maprotiline (eight trials used single-ingredient preparations and two used combinations of St. John’s wort and valerian). St. John’s wort
extracts were administered at doses ranging from 350 to 1800 mg; the hyperforin content of the preparations tested was not known. Most trials lasted for 4–6 weeks, although some studies were conducted for three months.

The results of the meta–analysis showed that St. John’s wort preparations were significantly superior to placebo in the short–term treatment of mild to moderately severe depressive disorders (rate ratio 2.47 and 95% confidence interval (95% CI) 1.69–3.61). St. John’s wort preparations were found to be as effective as conventional antidepressant agents (single preparations, rate ratio 1.01 and 95% CI 0.87–1.16), although for several reasons – for example, the use of low doses of conventional antidepressants and the trials involving small numbers of patients – this evidence was considered inadequate for establishing whether St. John’s wort was as effective as conventional antidepressant drugs. Further studies comparing St. John’s wort preparations with standard antidepressant agents in well–defined patient groups and over longer periods were considered necessary.

Another meta–analysis employed tighter inclusion criteria for trials in an effort to increase the validity of the analysis. It included only randomised, blinded, controlled trials of St. John’s wort as a single preparation, which involved patients with depressive disorders as defined by the standard criteria ICD-10 (International Statistical Classification of Diseases and Related Health Problems), DSM-IIIR (Diagnostic and Statistical Manual) or DSM-IV and which used the Hamilton Depression (HAMD) Scale for measuring clinical outcomes. Six such trials involving 651 patients with mainly mild to moderately severe depressive disorders were included; two trials were placebo controlled and four compared St. John’s wort with standard anti depressants. The studies lasted for 4–6 weeks and the doses of St. John’s wort extract ranged from 200 to 900 mg daily; the range for total hypericin administered was 0.75–2.7 mg daily.

This meta–analysis showed that the response rate for St. John’s wort was significantly greater than that for placebo (73.2 versus 37.9%, respectively, relative risk 1.48 and 95% CI 1.03–1.92) and similar to that observed with tricyclic antidepressants (64 versus 6.4% for St. John’s wort and tricyclic antidepressants, respectively, relative risk 1.11 and 95% CI 0.92–1.29). Despite the stringent inclusion criteria for trials in this meta–analysis, it was concluded that further studies are required in order to address methodological problems before it can be concluded that St. John’s wort is an effective antidepressant.

At least four randomised, controlled trials of monopreparations of St. John’s
worts involving patients with depressive disorders\(^{(64,69–71)}\) have been published since the Cochrane review.\(^{(14)}\) Two trials compared St. John’s wort against placebo only,\(^{(64,69)}\) one compared St. John’s wort with fluoxetine\(^{(70)}\) and one was a three-arm study comparing St. John’s wort with imipramine and placebo.\(^{(71)}\)

In a randomised, double-blind, multicentre study, 162 patients with mild to moderate depression received St. John’s wort extract (ZE117) (250 mg) twice daily (equivalent to 1 mg hypericin daily) or placebo for six weeks.\(^{(69)}\) At the end of the study, 56% of St. John’s wort-treated patients compared with 15% of placebo recipients were classified as responders according to recognised criteria. The proportions of patients reporting adverse events were similar between groups (7.4 and 6.2% for St. John’s wort and placebo, respectively).

Another randomised, double-blind, multicentre trial compared two different extracts of St. John’s wort with placebo in 147 patients with mild or moderate depression according to DSM-IV criteria.\(^{(64)}\) Patients received St. John’s wort extract (300 mg, WS 5573, containing 0.5% hyperforin or 300 mg, WS 5572, containing 5% hyperforin) or placebo three times daily for six weeks. Patients who received the extract containing 5% hyperforin showed the largest reduction in Hamilton Rating Scale for Depression scores from baseline values. Furthermore, 49% of these patients were classified as treatment responders (according to recognised criteria), whereas 38.8 and 32.7% of patients who received 0.5% hyperforin and placebo recipients, respectively, were classified as responders. The proportions of patients reporting adverse events were similar (28.6 versus 28.6 versus 30.6% for 5% hyperforin, 0.5% hyperforin and placebo, respectively). These findings were the first to show that the therapeutic effect of St. John’s wort in mild to moderate depression depends on its hyperforin content.\(^{(64)}\)

In a study comparing St. John’s wort with a selective serotonin reuptake inhibitor, 161 patients aged 60–80 years with mild or moderate depression according to ICD-10 criteria were randomised to receive St. John’s wort extract (LoHyp–57) (400 mg) twice daily or fluoxetine (10 mg) twice daily for six weeks.\(^{(70)}\) Neither the hypericin nor the hyperforin content of the St. John’s wort extract were stated in a published report of the study. At the end of the treatment period, 71.4% of St. John’s wort recipients and 72.2% of fluoxetine recipients were classified as responders according to recognised, pre-defined criteria. Similar efficacy for both St. John’s wort and fluoxetine was demonstrated when data from subgroups of patients with mild and moderate depression were analysed. The numbers of patients developing adverse reactions with a possible or probable relationship to treatment were 12 and 17 for St. John’s wort and fluoxetine, respectively, leading to
cessation of treatment in six and eight cases, respectively.\(^{(70)}\)

In a randomised, double-blind, multicentre trial in a primary care setting, 263 patients with moderate depression received St. John’s wort extract (350 mg) three times daily (STEI 300, containing 0.2–0.3% hypericin and 2–3% hyperforin, \(n = 106\)), imipramine (100 mg) daily (in three divided doses of 50, 25 and 25 mg, titrated from 50 mg on day 1 and 75 mg on days 2–4, \(n = 110\)) or placebo (\(n = 47\)) for eight weeks.\(^{(71)}\) Hypericum was found to be more effective than placebo after six weeks of treatment and to be as efficacious as imipramine after 8 weeks of treatment. In addition, both St. John’s wort and imipramine were shown to improve quality of life, as measured by the SF-36, to a greater extent than placebo. Adverse events were reported by 22% of St. John’s wort recipients, 46% of imipramine recipients and 19% of placebo recipients.

This study was criticised for its use of a relatively low dose of imipramine, such that the trial shows only that a comparatively high dose of St. John’s wort seems to be as effective as a comparatively low dose of imipramine.\(^{(72)}\) Nevertheless, this\(^{(71)}\) and other new trials\(^{(64,69)}\) have confirmed that St. John’s wort extracts are more effective than placebo in mild to moderately severe depression.\(^{(72)}\) However, further trials comparing St. John’s wort with standard antidepressants, particularly newer classes of agents such as the selective serotonin reuptake inhibitors, are still required. A large placebo-controlled trial comparing St. John’s wort extract (900–1800 mg daily) with the selective serotonin reuptake inhibitor sertraline (50–150 mg daily) in patients with major depression according to DSM-IV criteria is ongoing in the United States.\(^{(73)}\) Published abstracts of randomised, double-blind, controlled trials have reported superiority of St. John’s wort extract over placebo\(^{(74)}\) and equivalent efficacy between St. John’s wort and fluoxetine (20 mg) daily in mild to moderate depression\(^{(75,76)}\) and between St. John’s wort and imipramine (150 mg) daily.\(^{(76)}\)

In a dose-ranging trial involving 348 patients with mild to moderate depression according to ICD-10 criteria, patients were randomised to receive St. John’s wort extract three times daily equivalent to either 1 mg (\(n = 119\)), 0.33 mg (\(n = 115\)) or 0.17 mg (\(n = 114\)) hypericin for six weeks.\(^{(77)}\) At the end of the treatment period, there was a significant reduction in HAMD scores compared with baseline values. The response rates (according to recognised criteria) were 68, 65 and 62% for 1, 0.33 and 0.17 mg hypericin, respectively; the differences between groups were not statistically significant. Thus, the study showed that there was no dose-dependent effect of hypericin in St. John’s wort extracts.
Seasonal affective disorder
The effects of St. John’s wort extracts have been investigated in studies involving subjects with seasonal affective disorder (SAD),\(^ {78,79}\) although as yet there have not been any trials that have included a placebo control group. Twenty individuals with SAD were randomised to receive St. John’s wort (LI 160) (300 mg) three times daily (equivalent to 0.9 mg hypericin) with or without bright light therapy.\(^ {78}\) After four weeks, there were significant reductions in HAMD scores in both groups compared with baseline values and there were no statistically significant differences between groups. Another study evaluated data from individuals with mild to moderate SAD who had used St. John’s wort (300 mg) three times daily (equivalent to 0.9 mg hypericin) with \((n = 133)\) or without light therapy \((n = 168)\) for eight weeks.\(^ {79}\) The study was not randomised and involved data collection by postal questionnaires. Data from 301 returned questionnaires were suitable for analysis. Significant reductions in the mean SAD scores were observed in both groups compared with baseline values; the differences in the SAD scores between groups were statistically non–significant.

Antiviral activity
Antiviral activity has been reported for hypericin against human immuno deficiency virus (HIV) and hepatitis C.\(^ {80–82}\) Several uncontrolled studies in HIV-positive patients who received St. John’s wort extract have reported immunologic and clinical benefits, including increases in CD4 cell counts in some patients.\(^ {83,84}\) In a phase I, dose–escalating study, 30 HIV-positive patients with CD4 cell counts <350 cells/mm\(^3\) received intravenous synthetic hypericin twice weekly (0.25 or 0.5 mg/kg body–weight), three times weekly (0.25 mg/kg) or oral hypericin daily (0.5 mg/kg).\(^ {85}\) Sixteen patients discontinued treatment early because of toxic effects, and phototoxicity in several other patients prevented completion of dose escalation. Antiretroviral activity as assessed by significant changes in HIV p24 antigen level, HIV titre, HIV RNA copies and CD4 cell counts was not observed.

Other studies
The potential for the use of St. John’s wort in 20 individuals presenting with fatigue\(^ {86}\) and in 19 women with self–reported premenstrual syndrome\(^ {87}\) has also been explored in uncontrolled pilot studies. Significant improvements in perceived fatigue and in symptoms of depression and anxiety were seen after six weeks’ treatment with St. John’s wort (equivalent to 0.9 mg hypericin daily) compared with baseline values\(^ {86}\) and in overall premenstrual syndrome scores after treatment with St. John’s wort (equivalent to 0.9 mg hypericin daily) for two menstrual cycles.\(^ {87}\) Thus, there is scope for conducting randomised controlled trials of St. John’s wort in
In a randomised, double-blind, placebo-controlled trial, 179 women with menopause-related psycho vegetative symptoms received a combination preparation of St. John’s wort and black cohosh (Cimicifuga racemosa) or placebo for six weeks. The results indicated that the combination product had a significantly greater effect on the symptoms than did placebo. Postmarketing surveillance studies have been carried out with extracts of St. John’s wort in patients with psychovegetative disorders and in women with menopausal symptoms of psychological origin (see Side–effects, Toxicity). Improvements in symptom scores compared with baseline values following treatment with St. John’s wort extracts were reported in all studies; these studies did not involve a control group.

A randomised, double-blind, phase I study involving 55 healthy volunteers who received St. John’s wort (900 mg) daily (containing 0.5% hyperforin), St. John’s wort (900 mg) daily (containing 5.0% hyperforin) or placebo for eight days investigated the effects on quantitative electroencephalogram as an indicator of drug–induced pharmacological action. Reproducible central pharmacodynamic effects were apparent in both groups of St. John’s wort recipients compared with placebo recipients. The effects were greater in subjects who received extract containing 5.0% hyperforin than in those who received extract containing 0.5% hyperforin.

Placebo–controlled, crossover studies investigating the effects of St. John’s wort (0.9 and 1.8 mg) on the sleep polysomnogram of healthy subjects reported that both doses of St. John’s wort significantly increased rapid eye movement (REM) sleep latency compared with placebo, but had no effect on REM sleep duration or other parameters of sleep architecture.

In a randomised, double-blind, placebo-controlled trial involving 23 overweight but otherwise healthy adults, subjects who received treatment with St. John’s wort (900 mg) daily, Citrus aurantium extract (975 mg) daily and caffeine (528 mg) daily lost significantly more body weight than did subjects in the placebo and no–treatment control groups.

A placebo–controlled, crossover study in 19 healthy volunteers who received St. John’s wort for 15 days either alone or in combination with ethanol (to achieve a blood alcohol concentration of 0.05%) reported that there were no differences between the two groups in sense of well–being or adverse events.

A randomised, double-blind, placebo-controlled, six–week trial involving 72 long–distance runners and triathletes reported significant improvements in
endurance capacity in subjects who received vitamin E with St. John’s wort compared with subjects who received vitamin E alone or placebo.\(^{95}\)

**Pharmacokinetics**

Detailed pharmacokinetic studies have been carried out with the hypericin–standardised St. John’s wort extract LI 160.\(^{96}\) Administration of single oral doses of LI 160 (300, 900 and 1800 mg) to healthy male volunteers resulted in peak plasma hypericin concentrations of 1.5, 7.5 and 14.2 ng/mL for the three doses, respectively. Peak plasma concentrations were seen with hypericin after 2.0–2.6 hours and with pseudohypericin after 0.4–0.6 hours. The elimination half–life of hypericin was between 24.8 and 26.5 hours. Repeated doses of LI 160 (300 mg) three times daily resulted in steady–state concentrations after four days.\(^{96}\) Oral administration of the St. John’s wort extract WS 5572 (300 mg, equivalent to 14.8 mg hyperforin) resulted in peak plasma concentrations of 150 ng/mL being reached 3.5 hours after administration.\(^{97}\) The elimination half–life was 9 hours. Following repeated doses of 300 mg three times daily, the estimated steady–state plasma hyperforin concentrations were 100 ng/mL. Other studies investigating the pharmacokinetics of hypericum and hypericin have been summarised.\(^{1,G1}\)
A review of safety data for St. John’s wort obtained from reports of randomised controlled trials, drug monitoring and postmarketing surveillance studies (98–101) and national and international drug safety monitoring bodies has been published. (102) Collectively, the data indicate that St. John’s wort is well-tolerated. Adverse effects are generally mild; the most common adverse effects reported are gastrointestinal symptoms, dizziness, confusion and tiredness/sedation. In placebo–controlled trials, the frequency of adverse effects with St. John’s wort is similar to that for placebo. (102) Photosensitivity appears to be an extremely rare event with recommended doses of St. John’s wort (see below). (102)

Several postmarketing surveillance studies of the St. John’s wort extracts HYP811, (89,103) LI 160 (90,104) and Neuroplant (105) have since been published. These studies provide further confirmation of the tolerability of St. John’s wort extracts taken at recommended doses for short–term treatment (usually 4–6 weeks, although one study monitored 111 women for 12 weeks (90)). The frequency of adverse reactions in 6382 patients with mild depression who took St. John’s wort for six weeks was reported to be 0.125% (mainly skin reactions). (105)

A systematic review and meta–analysis of randomised controlled trials of St. John’s wort in patients with mild to moderately severe depressive disorders reported that, in the trials comparing St. John’s wort with standard antidepressants, the proportions of patients reporting side–effects were 26.3 and 44.7%, respectively (rate ratio 0.57 and 95% CI 0.4–0.69). (14) However, further studies investigating the long–term safety of St. John’s wort were advised. Another meta–analysis which employed tighter inclusion criteria reported that tricyclic antidepressants were associated with a higher proportion of side–effects than were St. John’s wort preparations (47 versus 26.4%, respectively, relative risk 1.72 and 95% CI 1.30–2.14). (68) Randomised controlled trials (64,69–71) published since the Cochrane meta–analysis (14) and published abstracts (74–76) have also reported that St. John’s wort has a more favourable short–term safety profile than standard antidepressants (70,71,75,76) and that the frequency of adverse events seen with St. John’s wort is similar to that for placebo (64,69,71,74) (see Clinical studies). In a comparative trial of St. John’s wort and fluoxetine, the frequency of adverse reactions associated with St. John’s wort was higher than expected, although it was stated that the effects reported were similar to those known to occur with fluoxetine. (70) The observation that the frequency of adverse effects is lower in placebo–controlled trials of St. John’s wort than in comparative trials with standard antidepressants has been made.
A review has attempted to compare the safety profile of St. John’s wort systematically with that of several conventional antidepressants. A review has attempted to compare the safety profile of St. John’s wort systematically with that of several conventional antidepressants. (102)

Photosensitivity
Sensitivity to sunlight following the ingestion of hypericum or hypericin is known as hypericism.

Delayed hypersensitivity or photodermatitis has been documented for St. John’s wort following the ingestion of a herbal tea made from the leaves. (107) Hypericin is stated to be the photosensitising agent present in St. John’s wort. (82, G33 G47) A review of the photodynamic actions of hypericin has been published. (108) In a double-blind, crossover, single-dose study in 13 healthy volunteers who received placebo or St. John’s wort extract (LI 160) (900, 1800 and 3600 mg containing 0, 2.81, 5.62 and 11.25 mg total hypericin, respectively), no evidence of photosensitivity was observed with or without St. John’s wort following skin irradiation with both UV-A and UV-B light 4 hours after dosing. (109) In a multiple-dose study in which 50 volunteers received St. John’s wort (LI 160) (600 mg) three times daily (equivalent to 5.6 mg total hypericin daily) for 15 days, a moderate increase in UV-A sensitivity was observed. (109) However, the doses used were higher than those recommended therapeutically. In another single-dose study, administration of St. John’s wort (LI 160) (1800 mg, equivalent to 5.4 mg total hypericin) to 12 healthy volunteers resulted in a mean serum total hypericin concentration of 43 ng/mL and a mean skin blister fluid concentration of 5.3 ng/mL. (110) After administration of St. John’s wort (300 mg) three times daily for seven days in order to achieve steady-state concentrations, the mean serum total hypericin concentration was 12.5 ng/mL and the mean skin blister fluid concentration was 2.8 ng/mL; these concentrations are below those estimated to be phototoxic (>100 ng/mL). (110)

The consumption of large quantities of St. John’s wort by grazing animals has been associated with the development of photosensitivity. (111, G22 G51) Mice given 0.2–0.5 mg of the herb were found to develop severe photodynamic effects. (G22) Studies using cell cultures of human keratinocytes incubated with hypericin or St. John’s wort extract and exposed to UV-A resulted in a reduction in the LC50 (lethal concentration) with hypericin, but only a mild reduction with hypericum. (112) From these findings it has been estimated that at least 30 times the therapeutic dose would be necessary to produce phototoxic effects in humans. (112) Experimental evidence has suggested that a solution of hypericin can react with visible and UV light to produce free
radical species and that this may lead to damage of proteins in the lens of the eye.\textsuperscript{(113)} There are no reports of cataract formation in individuals who have taken St. John’s wort.

A study reported that HIV-positive patients treated with oral hypericin (0.05 mg/kg) for 28 days developed mild symptoms of photosensitivity on exposure to sunlight and that two patients developed intolerable symptoms of photosensitivity when the dose was increased to 0.16 mg/kg.\textsuperscript{(114)} In a dose-escalating study involving 30 HIV-infected patients treated with oral (0.5 mg/kg daily) or intravenous hypericin (starting dosage 0.25 mg/kg twice or three times weekly), 16 patients discontinued treatment before completing eight weeks of therapy because of moderate or severe phototoxicity; severe cutaneous phototoxicity was observed in 11 out of 23 evaluable patients.\textsuperscript{(85)} Other serious clinical or laboratory adverse events were infrequent: elevation of alkaline phosphatase and hepatic aminotransferase concentrations to more than five times normal values was noted in two and three patients, respectively.

**Other effects**

\textbf{In humans} A case of subacute toxic neuropathy possibly related to the use of St. John’s wort and subsequent exposure to sunlight has been reported.\textsuperscript{(115)} A woman developed stinging pains in areas exposed to the sun (face and hands) four weeks after starting treatment with St. John’s wort (500 mg/day, extract and hypericin content not stated); the report did not state whether the woman was using any other products. Her symptoms improved three weeks after stopping St. John’s wort and disappeared over the next two months.

There have been reports of sensory nerve hypersensitivity occurring in individuals who have taken St. John’s wort preparations (tablets or tinctures).\textsuperscript{(116)}

Cases of mania\textsuperscript{(117,118)} and hypomania\textsuperscript{(119,120)} have been reported in individuals taking St. John’s wort preparations. Two cases of mania were reported in patients with bipolar depression who began self-treatment with standardised St. John’s wort extract (900 mg) daily\textsuperscript{(118)} and one in a patient experiencing a moderate depressive episode who was taking both sertraline and St. John’s wort (dosage not known).\textsuperscript{(117)} A case of hypomania was reported in a woman with panic disorder and unipolar major depression who had discontinued sertraline treatment one week before starting St. John’s wort tincture.\textsuperscript{(119)} Two cases of hypomania were reported in individuals with no history of bipolar disorder.\textsuperscript{(120)} A man who had received electroconvulsive therapy and who had previously taken various...
antidepressant drugs, including venlafaxine, fluvoxamine, moclobemide and nortriptyline, experienced a hypomanic episode six weeks after starting St. John’s wort (dosage not stated). A man with symptoms of post–traumatic stress disorder was diagnosed with an acute manic episode after three months of self–treatment with St. John’s wort (dosage not stated).\(^\text{120}\)

Several of these reports stated that the symptoms had resolved after stopping treatment with St. John’s wort, although in one case the patient improved but remained agitated despite cessation of St. John’s wort.\(^\text{120}\)

None of the cases involved rechallenge with St. John’s wort and, in all cases, there were other pharmacological factors and/or underlying illnesses that could have been responsible for or contributed to the precipitation of mania.

**IN ANIMALS AND Experimental studies investigating the genotoxic potential and mutagenic activity of St. John’s wort extracts in vitro and in vivo have been summarised.**\(^\text{G1 G52}\) In vivo studies and most in vitro studies provided negative results, indicating a lack of mutagenic potential with defined St. John’s wort extracts.\(^\text{G52}\) Mutagenic activity observed in an in vitro Ames test was attributed to the presence of quercetin, although other studies have found no mutagenic potential with a St. John’s wort extract and it has been stated that there is no valid evidence for the carcinogenicity of quercetin in humans.\(^\text{G1 G52}\)

Dietary administration of St. John’s wort to rats was found to have no affect on various hepatic drug–metabolising enzymes (e.g. aminopyrine, \textit{N}-demethylase, glutathione \textit{S}-transferase and epoxide hydrolase) or on copper concentrations in the liver (see Contra–indications, Warnings, Drug interactions). No major effects were observed on hepatic iron or zinc concentrations and no significant tissue lesions were found in four rats fed St. John’s wort in their daily diet for 119 days (10% for first 12 days and 5% thereafter because of unpalatability).\(^\text{121}\)

Cytotoxic constituents related to hyperforin have been isolated from two related \textit{Hypericum} species (see In vitro and animal studies).
Contra-indications, Warnings

Individuals with sensitivity towards St. John’s wort may experience allergic reactions. The use of St. John’s wort is not advised in known cases of photosensitivity and, in view of the potential of hypericin as a photosensitising agent, therapeutic UV treatment should be avoided whilst using St. John’s wort.\(^{(G1)}\)

It has previously been suggested that excessive doses of St. John’s wort may potentiate monoamine oxidase inhibitor therapy.\(^{(122)}\) However, as monoamine oxidase inhibitory activity has not been reported \textit{in vivo} with St. John’s wort, this warning is no longer considered necessary. In addition, avoidance of foodstuffs, such as those containing tyramine (e.g. cheese, wine, meat and yeast extracts) and medicines containing sympathomimetic agents (e.g. cough/cold remedies), which interact with MAOIs, is not considered necessary.

\textbf{Drug interactions}

Recent evidence has emerged from spontaneous reports\(^{(123)}\) and published case reports\(^{(124–127)}\) of interactions between St. John’s wort and certain prescribed medicines, leading to a loss of or reduction in the therapeutic effect of these prescribed medicines. Drugs that may be affected include indinavir, warfarin, ciclosporin, digoxin, theophylline and oral contraceptives. Drug interaction studies in healthy volunteers have provided supporting evidence of interactions between St. John’s wort and phenprocoumon\(^{(128)}\) and digoxin\(^{(129)}\) and have provided evidence that St. John’s wort may induce some cytochrome P450 (CYP) drug–metabolising enzymes in the liver,\(^{(128,130,131)}\) namely CYP3A4, CYP1A2 and CYP2C9, as well as affecting P-glycoprotein (a transport protein). Other studies have failed to find significant effects on CYP isoenzymes,\(^{(132–134)}\) although the numbers of volunteers may have been too small and the duration of St. John’s wort administration too short to exclude an inductive effect truly.\(^{(133,134)}\)

There have been other reports of increased serotonergic effects in patients taking St. John’s wort concurrently with selective serotonin reuptake inhibitors (e.g. sertraline, paroxetine).\(^{(135,136)}\)

Also of concern is that the content of active constituents can vary between different preparations of St. John’s wort; thus, the degree of enzyme induction may vary.

Collectively, these data led the UK Committee on Safety of Medicines (CSM) to issue advice to pharmacists, doctors and patients on the use of St. John’s
The CSM’s advice for healthcare professionals for patients taking St. John’s wort and certain drugs can be summarised as follows.

**Warfarin, Ciclosporin, Digoxin, Theophylline and Anticonvulsants (Carbamazepine, Phenobarbital and Phenytoin)** There is a risk of reduced therapeutic effect, e.g. risk of transplant rejection, seizures and loss of asthma control. Advice is to check plasma drug concentrations (with warfarin, the patient’s International Normalised Ratio should be checked) and to stop St. John’s wort therapy. In addition, dose adjustment may be necessary.

HIV protease inhibitors (indinavir, nelfinavir, ritonavir and saquinavir) and HIV non-nucleoside reverse transcriptase inhibitors (efavirenz and nevirapine) There is a risk of reduced blood concentrations with possible loss of HIV suppression. Advice is to measure HIV RNA viral load and to stop St. John’s wort.

**Oral Contraceptives** There is a risk of reduced blood concentrations, breakthrough bleeding and unintended pregnancy. Advice is to stop St. John’s wort.

**Triptans (Sumatriptan, Naratriptan, Rizatriptan and Zolmitriptan) and Selective Serotonin Reuptake Inhibitors (Citalopram, Fluoxetine, Fluvoxamine, Paroxetine and Sertraline)** There is a risk of increased serotonergic effects with the possibility of an increased risk of adverse reactions. Advice is to stop St. John’s wort.

Patients already taking any of the above drugs should be advised not to start taking St. John’s wort and users of other medicines should be advised to seek professional advice before using St. John’s wort. Topical medicines and non-psychotropic medicines that are excreted renally are not likely to interact with St. John’s wort. In addition, topical or homeopathic preparations of St. John’s wort are not likely to interact with prescribed medicines.

**Pregnancy and lactation**

Slight *in vitro* uterotonic activity has been reported for St. John’s wort (see *In vitro* and animal studies).

There is a report of a 38-year-old woman who started taking St. John’s wort (900 mg/day) at her 24th week of pregnancy, taking the last dose 24 hours before delivery. The pregnancy was unremarkable except for late onset of thrombocytopenia. Another report described a 43-year-old woman who discontinued fluoxetine and methylphenidate upon becoming pregnant and started taking St. John’s wort (900 mg/day). The report does not state the
outcome of the pregnancy,\(^{(139)}\) although it is assumed that had adverse events occurred, they would have been stated. In view of the lack of toxicity data, St. John’s wort should not be used during pregnancy and lactation.
Pharmaceutical Comment

The chemical composition of St. John’s wort has been well studied. Documented pharmacological activities provide supporting evidence for several of the traditional uses stated for St. John’s wort. Many pharmacological activities appear to be attributable to hypericin and to the flavonoid constituents; hypericin is also reported to be responsible for the photosensitive reactions that have been documented for St. John’s wort. With regard to the antidepressant effects of St. John’s wort, hyperforin rather than hypericin, as originally thought, has emerged as one of the major constituents responsible for antidepressant activity. However, further research is required in order to determine which other constituents contribute to the antidepressant effect.

Evidence from randomised, controlled trials has confirmed the efficacy of St. John’s wort extracts over placebo in the treatment of mild to moderately severe depression.\(^{(14)}\) Other randomised controlled studies have provided some evidence that St. John’s wort extracts are as effective as some standard antidepressants in mild to moderate depression. However, there is still a need for further trials in order to assess the efficacy of St. John’s wort extracts compared with that of standard antidepressants, particularly newer antidepressant agents such as the selective serotonin reuptake inhibitors. In addition, there is generally a need for further studies in well-defined groups of patients, in different types of depression and conducted over longer periods in order to determine long-term safety.\(^{(14)}\) St. John’s wort does appear to have a more favourable short-term safety profile than standard antidepressants, a factor that is likely to be important in patients continuing to take medication. Concerns have been raised over interactions between St. John’s wort and certain prescribed medicines (including warfarin, ciclosporin, theophylline, digoxin, HIV protease inhibitors, anticonvulsants, selective serotonin reuptake inhibitors, triptans and oral contraceptives); advice is that patients taking these medicines should stop taking St. John’s wort, generally after seeking professional advice as dose adjustment may be necessary. With the exception of oral contraceptives, patients taking these prescribed medicines should not be self-treating with over-the-counter medicines, including herbal medicines, without first seeking professional advice.

In view of the lack of long-term safety data for St. John’s wort and its reported photosensitising ability, excessive use of St. John’s wort should be avoided.
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Purpose and Scope

*Herbal Medicines* is intended to serve as a reference work for pharmacists, doctors and other healthcare professionals, assisting in their provision of advice on the use of herbal medicines to members of the public. *Herbal Medicines* is not intended to represent a guide to self–diagnosis and self–treatment with herbal medicines, and should not be used as such.

The term ‘herbal medicine’ (or ‘herbal remedy’) is used to describe a marketed product, whereas ‘herbal ingredient’ refers to an individual herb that is present in a herbal medicine. ‘Herbal constituent’ is used to describe a specific chemical constituent of a herbal ingredient. Thus, as examples, Valerian Tablets are a herbal product, valerian is a herbal ingredient, and valtrate is a herbal constituent of valerian.

The main criterion for inclusion of a herbal ingredient in the text is its presence in herbal medicines that are used in the UK, particularly those which are sold through pharmacies. In addition, herbs which have recently been the subject of media or scientific interest have also been included. The aim of *Herbal Medicines 2004* is to draw the attention of the reader to the reputed actions and uses of herbal ingredients, and to whether or not these have been substantiated by *in vitro* animal or human studies. In addition, any known or potential toxicities of herbal ingredients, and how these may influence the suitability for inclusion in herbal medicines or for use with conventional medicines, are also discussed.

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The introductory section to the 152 monographs on the individual herbal ingredients contained in *Herbal Medicines* discusses the legal aspects of herbal medicines including licensed medicines and non-licensed products in the UK and within the European Union (EU). All medicines are assessed for their quality, safety and efficacy and, in the context of herbal medicines, there are often specific criteria which are not encountered in the assessment of other medicines. As a first line in ensuring the safety and efficacy of herbal medicines there is a series of guidelines for quality assessment and this is briefly discussed. In terms of safety, it is a popular conception that just because herbs are ‘natural’ then they must also be safe. This is a misconception, and it is emphasised that some herbal ingredients have the capability to cause adverse effects, whilst some are decidedly toxic. Within the context of the 152 monographs on herbal ingredients, most have documented adverse effects, or the potential to interact with other medication, and few can be recommended for use during pregnancy.

Three tables in the Introduction and 22 appendices after the monographs summarise the safety aspects of these herbal ingredients and give information on biologically active herbal ingredients and their active principles. Clinical efficacy has not been established for the majority of the herbal ingredients described in this handbook and, in some instances, there is a lack of documentation for chemical constituents and for pharmacological actions.

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The Herbal Monographs

Some 152 monographs on individual herbal ingredients found in herbal products are included, the title used for the monograph being their preferred common name. A data sheet–type format was chosen for the monographs because it was felt important to arrange the relevant information in a format familiar to pharmacists and doctors. Although conventional data sheets are written for products, it was decided to draw up the data sheets for herbal ingredients and not for products.

The headings used in the herbal monographs are listed on below with a brief explanation of the information held under them.

<table>
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<tr>
<th>Monograph title</th>
<th>Common name for the herbal ingredient; if more than one common name exists, this is the chosen preferred name.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species (Family)</td>
<td>Preferred botanical name with authority, together with the plant family.</td>
</tr>
<tr>
<td>Synonym(s)</td>
<td>Other common or botanical names.</td>
</tr>
<tr>
<td>Part(s) Used</td>
<td>Plant part(s) traditionally used in herbal medicine.</td>
</tr>
<tr>
<td>Pharmacopoeial and Other Monographs</td>
<td>Key pharmacopoeial monographs and texts on herbal medicines.</td>
</tr>
<tr>
<td>Legal Category</td>
<td>Legal category of the herb with respect to licensed products. For the majority of herbal ingredients this will be the General Sales List (GSL).</td>
</tr>
<tr>
<td>Constituents</td>
<td>Main documented chemical constituents grouped into categories such as alkaloids (type specified), flavonoids, iridoid glycosides, saponins, tannins, triterpenes, volatile oil and other constituents for miscellaneous and minor chemical components.</td>
</tr>
</tbody>
</table>
Food Use

Provides an indication as to whether the herbal ingredient is used in foods. The Council of Europe (COE) category, which reflects the opinion of the COE on the suitability of the herbal ingredient for use as a food flavouring, is quoted where applicable.

Also where applicable, the Food and Drugs Administration (FDA) listing is stated, e.g. Generally Regarded as Safe (GRAS), and Herb of Undefined Safety.

Herbal Use

States the reputed actions and uses of the herbal ingredients, based on information from several sources. In some instances, current investigations of particular interest are included.

Dosage

States the traditional dose of the herbal ingredient, mainly from the British Herbal Pharmacopoeia (BHP) and German Commission E monographs, giving doses for plant part used (herb, rhizome, leaf), liquid extract and infusion. Where possible, dosages used in clinical trials are included.

Pharmacological Actions

Describes any documented pharmacological actions for the herbal ingredient. This is further divided into a section on *In vitro and animal studies* and a *Clinical studies* section, which describes studies involving humans.

Side–effects, Toxicity

Details documented side–effects to the herbal ingredient and toxicological studies. If side–effects or toxicity are generally associated with any of the constituents in the herbal ingredient, or with its plant family, then these are mentioned here. See also Table 1 and Table 2 in the Introduction to the Monographs.

Contra–indications, Warnings

Describes potential contra–indications and potential side–effects, and individuals who may be more susceptible to a particular side–effect. This section should be used in conjunction with Appendices 1–23. Comments on Pregnancy and lactation are included; a summary is provided in Table 3 of the Introduction to the Monographs.
This section is designed to give the reader an overall summary of the monograph contents, indicating the extent of phytochemical, pharmacological and clinical data available for the herbal ingredient, whether or not proposed herbal uses are justified, concerns over safety and, based on this information, whether or not the herbal ingredient is considered suitable for use as a herbal medicine.

References are included at the end of the text on each monograph. There is considerable literature on herbal plants and general references have been selected for use with the handbook. These General References, referred to as G1 to G65, are listed after the Introduction. For some well-known herbal ingredients only general references are cited. The majority of the monographs also contain specific references which are cited at the end of each monograph.
Herbal medicines are also referred to as herbal remedies, herbal products, herbal medicinal products, phytomedicines, phytotherapeutic agents and phytopharmaceuticals. The use of herbal medicines in an evidence- or science–based approach for the treatment and prevention of disease is known as (rational) phytotherapy. This approach to the use of herbal medicines contrasts with traditional medical herbalism in the UK, which uses herbal medicines in a holistic manner and mainly on the basis of their empirical and traditional uses. Plants have been used medicinally for thousands of years by cultures all over the world. According to the World Health Organization, 80% of the world’s population uses plant-based remedies as their primary form of healthcare;\(^{(12)}\) in some countries, herbal medicines are still a central part of the medical system, such as Ayurvedic medicine in India and traditional Chinese medicine. Herbal medicine has a long history and tradition in Europe.

Although these two approaches – traditional/holistic and rational/evidence-based – are entirely different, in some instances they use the same terminology. For example, traditional herbalism is also described as ‘phytotherapy’ and refers to preparations of plant material as ‘herbal medicines’.

Furthermore, herbal medicines and homeopathic remedies are often mistaken by the layperson to be similar. However, homeopathy is based on the principle of ‘like should be treated by like’, and involves the administration of minute doses of remedies that, in larger doses, produce symptoms in a healthy person mimicking those expressed by people who are ill. Many, but not all, homeopathic remedies originate from plants. By contrast, herbal medicine (phyto therapy) involves the use of dried plant material or extracts of plant parts in therapeutic doses to treat the symptoms exhibited. In this respect, it is similar to conventional medicine.

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Regulatory Controls on Herbal Medicines

Sub-sections

- Herbal medicinal products in Europe
- Future regulation of herbal medicinal products in the EU
- Current regulatory position of herbal products in the UK
- Licensed herbal medicinal products
- Herbal remedies exempt from licensing
- Control of herbal ingredients in the UK
- Review of herbal medicines in the UK
- Regulatory control of herbal medicines world-wide

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In order to ensure public health, medicinal products must be safe, efficacious and of suitable quality for use. To obtain a marketing authorisation (product licence) within the EU, manufacturers of herbal medicinal products are required to demonstrate that their products meet acceptable standards of quality, safety and efficacy.
Echinacea

Echinacea is widely used throughout Europe for the prevention and treatment of colds and other upper respiratory tract infections. A recent Cochrane review of 16 clinical trials has reported that the overall results suggest that some products may have an effect greater than placebo, but that overall the results were inconclusive (see Echinacea).\(^{(118)}\)

Garlic

Numerous studies and systematic reviews have investigated the effects of garlic preparations in lowering raised serum cholesterol concentrations (see Garlic). Generally, the studies report beneficial results for garlic. However, the evidence at present is insufficient to recommend garlic as routine treatment for hypercholesterolaemia. One of the major problems in assessing the evidence available on garlic is the wide variation in the chemical composition of the products available, compared with fresh garlic. Further controlled studies are needed using standardised preparations to investigate efficacy in reducing serum lipids, blood pressure, platelet aggregation and antimicrobial activity (see Garlic).

Ginger

Some clinical studies have reported ginger to be an effective prophylactic against motion sickness, although subsequent studies have found ginger to be ineffective (see Ginger).

Ginkgo

Ginkgo is widely used in France and Germany in authorised herbal medicinal products for the treatment of circulatory insufficiencies (peripheral and cerebral). Currently, no licensed herbal medicinal products containing ginkgo are available in the UK.

Several systematic reviews have been carried out analysing the available evidence on the effects of ginkgo in cerebral insufficiency, dementia, tinnitus and intermittent claudication (see Ginkgo). Overall the results suggest some beneficial effects, but further studies are needed.
Ginseng is widely renowned for its adaptogenic properties in Eastern countries, where it is used to help the body cope with stress and fatigue, and to promote recovery from illness or imbalance such as hypertension or hypoglycaemia. Generally, it is only recommended to be used for certain individuals with specific illnesses. By comparison, in the UK, ginseng is mainly self-administered and taken in the form of tablets or capsules containing dried extracts of the root. Ginseng products available in the UK are sold as food supplements, often in combination with vitamins and minerals. A wealth of research describing a wide range of pharmacological activities, particularly on the hypothalamic and pituitary regions of the brain, has been documented for ginseng (see Ginseng, Eleutherococcus).

**Saw palmetto**

Saw palmetto is widely used in Europe, particularly in Germany, for symptoms associated with benign prostatic hypertrophy (BPH). In the UK, saw palmetto is licensed in a number of products for the symptomatic relief of short-term, male urinary discomfort. Results of clinical trials indicate that saw palmetto is a potential agent for the symptomatic treatment of BPH (see Saw Palmetto).

**Valerian**

Valerian is widely used in Europe for nervous tension and for promoting sleep. The therapeutic indications proposed by the EMEA HMPWP include relief of temporary, mild nervous tension and temporary difficulty in falling asleep. A systematic review of randomised, double-blind, placebo-controlled trials of valerian reported inconsistencies in methodology between studies, and that the evidence for efficacy was inconclusive (see Valerian). It is unclear whether the active principles in valerian are associated with the volatile oil, the iridoid components termed valepotriates or with some other, as yet unidentified, group of constituents.
Conclusion

The use of herbal medicinal products, including use in addition to or instead of conventional medicines, is continuing to increase. Healthcare professionals need to be aware that patients may be taking herbal medicinal products, and need to understand their effects and be aware of the potential problems associated with their use. This handbook provides the reader with factual information on almost 150 herbal ingredients present in herbal medicinal products in European and other developed countries. Herbal medicinal products can offer an alternative to conventional medicines in non–life–threatening conditions, providing they are of adequate quality and safety, and are used in an appropriate manner by suitable individuals.


17. *Monographs on the Medicinal Uses of Plant Drugs*, Fascicules 1 and 2


than Veterinary Drugs) (Prescription Only) Order 1983, as amended.


71. Schaumburg HH et al. Alopecia and sensory polyneuropathy from thallium


74. Cuncha J *et al.* Arsenic and acute lethal intoxication. *Hong Kong Pharm J* 1998; **7**: 50–53.


90. Ljunggren B. Severe phototoxic burn following celery ingestion. *Arch Dermatol* 1990; **126**: 1334–1336. (PubMed)


14. The Cochrane Library.
16. Wurglics M *et al*. Comparison of German St John’s Wort Products according to hyperforin and total hypericin content. *J Am Pharm Assoc* 2001; **41**: 560–566.
A typical alkaloid is chemically basic (alkaline) and contains a secondary or tertiary amine function within a heterocyclic ring (e.g. codeine). Alkaloids may be classified by their chemical skeleta (Figure 1) into the following major types: pyrrolidine (e.g. betonicine from white horehound); pyridine (e.g. gentianine from gentian); piperidine (e.g. lobeline from lobelia); pyrrolizidine (e.g. symphytine from comfrey); quinolizidine (e.g. sparteine from broom); quinoline (e.g. quinine from cinchona); isoquinoline (e.g. boldine from boldo); indole (e.g. harman from passionflower); tropane (e.g. hyoscine from belladonna); imidazole (e.g. pilocarpine from jaborandi); and xanthine (e.g. caffeine from maté). Biosynthetically related compounds that do not follow the above definition of an alkaloid may also be referred to as alkaloids, for example phenylalkylamines that do not contain an \(N\)-heterocyclic ring (e.g. ephedrine from ephedra), or that are not basic (e.g. colchicine from colchicum). Examples of medicinal plants that contain alkaloids are given in Appendix 14. For further information on alkaloids the reader is referred to other texts (e.g. references 1, 3 and 6).
A glycoside consists of two components, an aglycone (non-sugar) part and a sugar part. The aglycone portion may be of several different types of secondary metabolite (Figure 2), including coumarin (e.g. scopolin from horse-chestnut, see also Appendix 17), flavonoid (e.g. rutin from buchu, see also Appendix 18), or hydroxyanthracene (e.g. cascaroside A from cascara, see also Appendix 2). The sugar moiety is linked to the aglycone by a direct carbon-to-carbon bond (C-glycoside), or through an oxygen-to-carbon bond (O-glycoside). Cyanide glycosides, (e.g. amygdalin from apricot) release toxic hydrogen cyanide when cells are damaged and act as a defence mechanism. Glucosinolates (e.g. sinigrin from horseradish) contain nitrogen and sulphur and are pungent. Hydroxyanthracene glycosides are the active principles of the laxative herbs cascara and senna (see also Appendix 2).
Many of the aromatic constituents of plants contain hydroxy substituents and are phenolic. There is a wide variety of phenolics in medicinal plants and they range in chemical structure from simple phenolic acids (Figure 3), e.g. caffeic acid from artichoke, to complex tannins (see Appendix 21).

Where chemical structures are included in a monograph they may be the active principles or they may be compounds that can be used as chemical markers for that plant, i.e. they are present in significant quantities or are otherwise characteristic for a particular plant. As some compounds are of common occurrence in medicinal plants, their chemical structures are not necessarily included within a monograph. Some of the commonly encountered natural products, including mono-, sesqui-, di- and tri-terpenes, flavonoids and tannins, are briefly summarised below.
Terpenes are derived from two C5 units (isopentane), dimethylallylpyrophosphate and isopentenylpyrophosphate. The monoterpenes contain two isopentane units (C10) and are constituents of many volatile oils (see Appendix 22). Some of the more common monoterpenes are shown in Figure 4. Sesquiterpenes contain three isopentane units (C15) and occur as different skeletal types, e.g. eudesmane, germacrene, guaiane (Figure 5). A large number of sesquiterpenes contain a γ-lactone ring and these are known as sesquiterpene lactones. Some sesquiterpene lactones are allergenic (see Appendix 12). Examples of different skeletal types of molecule, including eudesmolide, germacranolide and guaianolide (e.g. constituents of comfrey) and pseudoguaianolide (e.g. matricine, chamomile), are illustrated in Figure 5. Diterpenes are derived from four isopentane units (C20) and examples of abietane (e.g. carnosic acid from sage), daphnane, kavane, labdane (e.g. rotundifaraine from agnus castus), taxane and tigliane are given in Figure 6. The ginkgolides from ginkgo are examples of complex diterpenes. Triterpenes are derived from six isopentane units (C30), and some commonly occurring compounds are illustrated in Figure 7, e.g. campesterol, β-sitosterol, stigmasterol, α- and β-amyrin, oleanoic and ursolic acids. Cardiac glycosides (cardenolides, see Appendix 3) and saponins (see Appendix 20) are examples of triterpenes that are less widely distributed in plants than the triterpenes illustrated in Figure 7.
Flavonoids

Flavonoids are biosynthesised from a phenylpropane unit (C6–C3), derived via shikimic acid and phenylalanine, and a C6 unit from three molecules of malonyl-CoA. They are widely distributed in the plant kingdom and occur in many medicinal plants (see Appendix 18). There are five major types: chalcones, flavanones, flavones, flavonols and anthocyanins (Figure 8). The flavones and their 3-hydroxy analogues (flavonols) are the most widespread. The five aglycones kaempferol, quercetin, myricetin, apigenin and luteolin, as well as the quercetin glycosides quercitrin and rutin are among the most commonly present in medicinal plants (Figure 9).
Tannins

Tannins are also common constituents of many medicinal plants (see Appendix 21) and they occur as two major types – the hydrolysable tannins and the non-hydrolysable (condensed) tannins. The hydrolysable tannins are esters of sugars with phenolic acids and they are either gallotannins (galloyl esters of glucose), e.g. pentagalloyl glucose, or ellagitannins (hexahydrodiphenic acid, derived from two units of gallic acid, esters with glucose), e.g. agrimoniin from agrimony. Non-hydrolysable tannins, also known as condensed tannins or proanthocyanidins, are polymers of catechin or gallocatechin linked by C-C bonds (e.g. cola tannins). Examples of some chemical structures of hydrolysable and non-hydrolysable tannins are given in Figure 10.


Agnus Castus
Species (Family)

*Vitex agnus-castus* L. (Verbenaceae)
Synonym(s)
Chasteberry, Chaste Tree, Monk’s Pepper
Part(s) Used

Fruit
Pharmacopoeial and Other Monographs

American Herbal Pharmacopoeia\(^{(G1)}\)

BHP 1996\(^{(G9)}\)

Complete German Commission E\(^{(G3)}\)

ESCOP 1997\(^{(G52)}\)

Martindale 33rd edition\(^{(G67)}\)

Mills and Bone \(^{(G50)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents

See General Reference G40.

**Alkaloids**
Vitcin

**Diterpenes**
Rotundifuran (labdane–type), vitexilactone and 6β,7β-diacetoxy–13–hydroxy–labda–8,14–diene.\(^{(1)}\)

**Flavonoids**
Flavonol (kaempferol, quercetagetin) derivatives, the major constituent being casticin. Other identified flavonoids include penduletin and chrysoplenol D.\(^{(2–4)}\)

**Iridoids**
In the leaf: 0.3% aucubin, 0.6% its \(p\)-hydroxybenzoyl derivative agnuside and 0.07% unidentified glycosides.\(^{(3,5)}\)

**Other constituents**
Fatty acids, including stearic and palmitic, volatile oil 0.5% with cineol and pinene as main components; castine (a bitter principle).\(^{(6,7)}\)
Food Use

Agnus castus is not used in foods.
Herbal Use

Traditionally, agnus castus has been used for menstrual problems resulting from corpus luteum deficiency, including premenstrual symptoms and spasmodic dysmenorrhea, for certain menopausal conditions, and for insufficient lactation.\(^{(G4 G49)}\) The German Commission E approved it for internal use for irregularities of the menstrual cycle, premenstrual complaints and mastodynia.\(^{(G3)}\)
Dosage

**Fruit**
0.5–1.0 g three times daily;\((G4)^9\) by contrast, 30–40 mg daily of crushed fruit.\((G3)\)

**Tincture**
1 : 5 (g/mL), 50–70% ethanol (v/v) 0.15–0.2 mL.\((G4)\)
Pharmacological Actions

In vitro and animal studies

Agnus castus does not contain any oestrogenic constituents but has been reported to diminish release of follicle-stimulating hormone from the anterior pituitary whilst increasing the release of luteinising hormone and prolactin.\(^{(8,9)}\)

Extracts of agnus castus act at dopamine receptors and affect prolactin release. Dopamine D\(_2\)-receptor binding of extracts has been demonstrated for three different dopamine receptors (rat striatum, calf striatum and human recombinant receptors) and for two separate ligands (sulpiridine and spiroperidol).\(^{(1,10,11)}\) The active compounds acted as dopamine agonists and were characterised as labdane diterpenes. The two most active diterpenes, rotundifuran and 6β,7β-diacetoxy-13-hydroxy-labda-8,14-diene, had IC\(_{50}\) values (calf striatum preparation, \(^3\)H-spiroperidol ligand) of 45 μg/mL (124 nmol/mL) and 79 μg/mL (194 nmol/mL), respectively.\(^{(11)}\) A lyophilised extract of agnus castus (5 mg/mL) was similar in activity to \(10^{-4}\) mol/L dopamine in receptor–ligand binding assays, displacing \(^3\)H-spiroperidol from calf brain striatal preparations.\(^{(10)}\)

An extract of agnus castus inhibited release of acetylcholine from \(^3\)H-choline loaded rat brain striatal cells on electrical stimulation, and had an IC\(_{50}\) value of 30 μg/mL.\(^{(11)}\)

Hexane fractions of agnus castus bind to human opiate receptors with IC\(_{50}\) values of 20 μg/mL (μ-receptors) and 10 μg/mL (κ-receptors).\(^{(12)}\)

Agnus castus extracts and fractions dose–dependently stimulated galactosidase activity in yeast cells; this may be indicative of oestrogen–receptor binding. Radioligand–receptor binding assays have demonstrated that agnus castus only binds weakly to oestrogen receptors in comparison to dopamine receptor binding.\(^{(12)}\)

Clinical studies

A proprietary preparation containing an alcoholic extract of agnus castus (0.2% w/w) has been available in Germany since the 1950s. It is used in the treatment of breast disease and pain, ovarian insufficiency, and dysfunctional uterine bleeding.\(^{(8,13-17)}\) The clinical effects of agnus castus have been reviewed.\(^{(18,19)}\) Details of clinical studies described in these reviews are summarised below.
Effects on prolactin secretion
Several open, uncontrolled studies involving small numbers of women with fertility disorders, hyperprolactinaemia and menstrual disorders have explored the effects of treatment with extracts of agnus castus (e.g. Mastodynon and Agnucaston; Bionorica), generally at doses equivalent to 30–40 mg drug for several months. These studies report decreased prolactin concentrations at the end of treatment, compared with baseline values.\(^{18,19}\)

A double-blind, placebo-controlled study involving women with cyclic mastalgia compared the effects of Mastodynon tablets (\(n = 32\)) and Mastodynon solution (\(n = 31\)) with those of placebo (\(n = 38\)) over three menstrual cycles. Prolactin concentrations in the two treated groups were significantly reduced, compared with those in the placebo group.

Other studies have reported that women with normal basal prolactin concentrations do not experience significant reduction in the concentration following treatment with agnus castus.

In a double-blind, placebo-controlled trial, 37 women with deficiencies in corpus luteal phase and latent hyperprolactinaemia received an agnus castus preparation (\(n = 17\)) or placebo (\(n = 20\)) for three menstrual cycles. In the treated group, the luteal phase was extended to 10.5 days from an initial 3.4–5.5 days.

The effects of agnus castus have also been investigated in men. In an open, uncontrolled study, 20 healthy men were given a commercial preparation of agnus castus extract (BP 1095E1) at doses ranging from 120 to 480 mg extract (3–12 times higher than doses used in women) for 14 days.\(^{20}\) The lowest dose was reported to increase serum prolactin concentrations, whereas the higher dose decreased prolactin concentrations.

Effects on mastodynia
In an open, uncontrolled trial, 825 women with mastodynia received Mastodynon for three months. At the end of the study, 465 patients (56\%) said they were symptom-free, and a further 198 patients (24\%) reported that their symptoms had improved. A subsequent trial involving 121 women reported that 75\% of participants experienced relief of symptoms.

In a randomised, double-blind, placebo-controlled study, 104 patients with breast pain were treated for three menstrual cycles with either Mastodynon solution (\(n = 34\)), tablets (\(n = 32\)) or placebo (\(n = 38\)).\(^{19}\) Patients in both treated groups claimed to have reduced breast pain; the findings were reported to be statistically significant for treatment, compared with placebo.
Another randomised, double-blind, placebo-controlled trial involving 97 patients with cyclic mastalgia compared Mastodynon solution \((n=48)\) with placebo \((n=49)\).\(^{21}\) After two menstrual cycles, pain intensity, as assessed by visual analogue scale (VAS) scores was reduced in both groups; the reduction was significantly greater in the agnus castus group \((p=0.006)\). However, at the end of the study (after three cycles), the reduction in pain intensity between the two groups was no longer statistically significant \((p=0.064)\). It was reported that Mastodynon was well-tolerated.

**Effects on premenstrual syndrome**

Five postmarketing surveillance studies involving more than 5000 women have monitored the effects of agnus castus preparations in premenstrual syndrome (PMS).\(^{19}\)

A randomised, double-blind, controlled trial involving 175 women with PMS compared Agnolyt (Madaus) one capsule daily (equivalent to 3.5–4.2 mg dry extract) \((n=90)\) and pyridoxine two capsules daily \((n=85)\) over three menstrual cycles.

Therapeutic response was measured using a premenstrual tension scale (self-assessment) and a clinical global impression (CGI) scale (physician assessment). Reductions in PMS scale scores (of around 48%) were reported for both groups.\(^{22}\) It was concluded that the two treatments were equally effective in the treatment of symptoms of PMS.

In a randomised, double-blind, placebo-controlled, parallel-group trial, 170 women with PMS received agnus castus extract ZE 440 (60% ethanol, extract ratio 6 to 12 : 1; standardised for casticin) 20 mg daily \((n=86)\) or placebo \((n=84)\) for three menstrual cycles.\(^{23}\) The main outcome measure was the participant’s self-assessment of PMS symptoms (irritability, mood alteration, anger, headache, breast fullness and bloating). At the end of the third cycle, improvements in the PMS symptoms were significantly greater in the agnus castus group, compared with the placebo group \((p<0.001)\). Clinical global impression scores for severity of condition, global improvement and overall risk/benefit were also significantly better for agnus castus, compared with placebo \((p<0.001)\). Mild adverse events were reported by four agnus castus recipients and three placebo recipients. All resolved without treatment. It was concluded that agnus castus dry extract is an effective and well-tolerated treatment for symptoms of PMS.\(^{23}\)

Agnus castus has also been reported to be effective in the treatment of endocrine disorders such as menstrual neuroses and dermatoses \(^{24}\) and has been used for the treatment of acne.\(^{25,26}\)
A lactogenic action has been documented for agnus castus;\textsuperscript{(27)} chemical analysis of the breast milk revealed no changes in composition.

The precise mode of action of agnus castus and the active constituents has not been established. However, it is thought to act on the pituitary-hypothalamic axis rather than directly on the ovaries.
Side–effects, Toxicity

Agnus castus is generally well–tolerated, although allergic reactions (which resolved following discontinuation of agnus castus therapy), headaches and an increase in menstrual flow have been reported.\(^8,24\)
Contra-indications, Warnings

Agnus castus has dopaminergic activity and should not be used with dopamine receptor antagonists or agonists.

**Pregnancy and lactation**

In view of the documented pharmacological actions and lack of toxicity data, the use of agnus castus during pregnancy should be avoided. Agnus castus has been reported to stimulate milk secretion without altering the composition of the breast milk.\(^{(24,27)}\) Nevertheless, agnus castus should be avoided during lactation until further information is available.
Pharmaceutical Comment

The chemistry and pharmacology of agnus castus are well–documented. There is clinical evidence indicating that raised prolactin concentrations are reduced following agnus castus treatment, and that there are beneficial effects in mastodynia and in symptoms of PMS.
References

See also General References G1 G3 G5 G9 G31 G36 G40 G43 G49 G50.


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Species (Family)

*Agrimonia eupatoria* L. (Rosaceae)
Synonym(s)

Agrimonia
Part(s) Used
Herb
Pharmacopoeial and Other Monographs

BHP 1996 \( ^{G9} \)

BP 2002 \( ^{G71} \)

Complete German Commission E \( ^{G3} \)

Martindale 33rd edition \( ^{G67} \)

PDR for Herbal Medicines 2nd edition \( ^{G36} \)

Ph Eur 2004 \( ^{G72} \)
Legal Category (Licensed Products)

GSL$^{(G37)}$
Constituents
See References 1 and 2, and General References G2 G22 G31 G40 G64.

Acids
Palmitic acid, salicylic acid, silicic acid and stearic acid.

Flavonoids
Apigenin, luteolin, luteolin–7–glucoside, quercetin, quercitrin, kaempferol and glycosides.\(^{(3)}\)

Tannins
3–21%. Condensed tannins in herb; hydrolysable tannins (e.g. ellagitannin).

Vitamins
Ascorbic acid (vitamin C), nicotinamide complex (about 100–300 μg/g leaf), thiamine (about 2 μg/g leaf) and vitamin K.

Other constituents
Bitter principle, triterpenes (e.g. α-amyrin, ursolic acid, euscapic acid), phytosterols and volatile oil 0.2%.
Agrimony is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that agrimony can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\(^G16\)
Herbal Use

Agrimony is stated to possess mild astringent and diuretic properties.\(^1\) It has been used for diarrhoea in children, mucous colitis, grumbling appendicitis, urinary incontinence, cystitis, and as a gargle for acute sore throat and chronic nasopharyngeal catarrh.\(^{G2\ G7}\)
Dosage

*Dried herb*
2–4 g by infusion three times daily.\(^{[G7]}\)

*Liquid extract*
1–3 mL (1 : 1 in 25% alcohol) three times daily.\(^{[G7]}\)

*Tincture*
1–4 mL (1 : 5 in 45% alcohol) three times daily.\(^{[G7]}\)
Pharmacological Actions

**In vitro and animal studies**

Significant uricolytic activity has been documented for agrimony infusions and decoctions (15% w/v), following their oral administration to male rats at a dose of 20 mL/kg body weight (equivalent to 3 g dry drug).\(^4\) Diuretic activity was stated to be minimal and elimination of urea unchanged. A hypotensive effect in anaesthetised cats has been documented for an agrimony extract given by intravenous injection; blood pressure was lowered by more than 40%.\(^5\)

Marked antibacterial activity against *Staphylococcus aureus* and α-haemolytic streptococci has been reported for agrimony.\(^6\)

An aqueous ethanol extract of the herb was tested for immunomodulative activity in the peritoneal cavities of mice.\(^7\) Immunostimulant activity resulted in an increase in phagocytic activity and increases in the activities of lysozyme and peroxidase. *Agrimonia eupatoria* given in the diet of mice for 12 days prior to intraperitoneal administration of streptozotocin resulted in a reduction in hyperglycaemia.\(^8\) Further investigation revealed stimulation of 2-deoxyglucose transport, glucose oxidation and incorporation of glucose into glycogen in mouse abdominal muscle. An aqueous extract (0.25–1 mg/mL) stimulated insulin secretion from a BRIN-BD11 pancreatic B cell line.\(^9\) These findings demonstrate that *A. eupatoria* aqueous extract given orally to mice has antihyperglycaemic, insulin–releasing and insulin–like activity.\(^9\)

A related species, *A. pilosa*, has also been investigated. *In vivo* antitumour activity in mice has been attributed to the tannin agrimoniin\(^10\) which has not been reported as a constituent of *A. eupatoria*. Agrimoniin was administered intraperitoneally into ascites–type and solid tumours in rodents.\(^11\) At doses of greater than 10 mg/kg, given before or after intraperitoneal inoculation with MM2 cells, it completely rejected tumour growth in mice.\(^11\) Solid tumours of MH134 and Meth-A were inhibited by agrimoniin, and the number of peripheral blood cells was increased, indicating that agrimoniin has antitumour activity and that it exerts its effect by enhancing the immune response. *In vitro* studies have reported that agrimoniin induces the cytotoxicity of murine peritoneal exudate cells,\(^12\) and that it induces interleukin 1 in human peripheral blood mononuclear cells and in mouse adherent peritoneal exudate cells *in vivo*.\(^13\) Several phloroglucinols isolated from *A. pilosa* have demonstrated activity against *Staphylococcus aureus*,\(^14\) and a methanol extract of the herb inhibited HIV-1 protease activity.\(^15\) An aqueous suspension of *A. pilosa* herb (1 mg/kg and
5 mg/kg) given orally or intraperitoneally significantly reduced blood glucose concentrations in streptozotocin–induced diabetic rats.\(^{(16)}\)

**Clinical studies**

The successful treatment of cutaneous porphyria in a group of 20 patients receiving agrimony infusions has been described.\(^{(17)}\) An improvement in skin eruptions together with a decrease in serum iron concentrations and in urinary porphyrins was noted.

A compound herb preparation containing agrimony has been used to treat 35 patients suffering from chronic gastroduodenitis.\(^{(18)}\) After 25 days of therapy, 75% of patients claimed to be free from pain, 95% from dyspeptic symptoms and 76% from palpitation pains. Gastroscopy was said to indicate that previous erosion and haemorrhagic mucous changes had healed. No side-effects or signs of toxicity were documented.
Side-effects, Toxicity

None documented for *A. eupatoria*. A polar fraction containing flavonoids and triterpenes, but not tannins, produced a negative result in the Ames test.\(^{(1)}\)

In mice, agrimonin has been documented to cause stretching and writhing reactions when administered by intraperitoneal injection, and cyanosis and necrosis at the site of intravenous injection.\(^{(11)}\) These reactions were considered to be inflammatory reactions. The LD\(_{50}\) of agrimonin in mice has been estimated as 33 mg/kg (by intravenous injection), 101 mg/kg (by intraperitoneal injection), and greater than 1 g/kg (by mouth).\(^{(11)}\) Cytotoxic activity has been reported for *A. pilosa*\(^{(10)}\) (see *In vitro* and animal studies).
Contra-indications, Warnings

Excessive doses may interfere with existing drug treatment for high or low blood pressure, and anti coagulant therapy. In view of the tannin constituents, excessive use should be avoided.

Pregnancy and lactation
Agrimony is reputed to affect the menstrual cycle.\textsuperscript{G22} In view of the lack of toxicity data, excessive use of agrimony should be avoided during pregnancy and lactation.
Relatively limited information is available on the chemistry of agrimony, although more is known about the tannin constituents of the related species *A. pilosa*. Human studies have indicated that agrimony may be useful in the treatment of certain cutaneous and gastrointestinal disorders, although further studies are needed to confirm these reports. The tannin constituents may justify the astringent activity attributed to the herb. In view of the lack of toxicity data, excessive use of agrimony should be avoided.
References

See also General References G2 G3 G9 G16 G22 G31 G36 G37 G40 G42 G64.


Species (Family)

Medicago sativa L. (Fabaceae/Leguminosae)
Synonym(s)

Lucerne, Medicago, Purple Medick
Part(s) Used

Herb
Pharmacopoeial and Other Monographs

BHP 1996\(^{G9}\)

Martindale 33rd edition\(^{G67}\)

PDR for Herbal Medicines 2nd edition\(^{G36}\)
Constituents
See General References G19 G22 G41 G64.

**Acids**
Lauric acid, maleic acid, malic acid, malonic acid, myristic acid, oxalic acid, palmitic acid and quinic acid.

**Alkaloids**
Pyrrolidine-type (e.g. stachydrine, homostachydrine); pyridine-type (e.g. trigonelline) in the seeds only.

**Amino acids**
Arginine, asparagine (high concentration in seeds), cystine, histidine, isoleucine, leucine, lysine, methionine, phenylalanine, threonine, tryptophan and valine. The non–protein toxic amino acid canavanine is present in leaves (0.9–1.2 mg/g), stems (0.6–0.9 mg/g) and seeds (5–14 mg/g).^{G19}

**Coumarins**
Medicagol.

**Isoflavonoids**
Coumestrol, biochanin A, daidzein, formononetin and genistein.

**Saponins**
2–3%. Hydrolysis yields aglycones, medicagenic acid, soyasapogenols A–F and hederagenin.\(^1\) Sugar chain components include arabinose, galactose, glucuronic acid, glucose, rhamnose and xylose.

**Steroids**
Campesterol, cycloartenol, β-sitosterol (major component), α-spinasterol and stigmasterol.

**Other constituents**
Carbohydrates (e.g. arabinose, fructose, sucrose, xylose), vitamins (A, B\(_1\), B\(_6\), B\(_{12}\), C, E, K), pectin methylesterase, pigments (e.g. chlorophyll, xanthophyll, β-carotene, anthocyanins), proteins, minerals and trace elements.

See reference G22 for more detailed chemical information.
Food Use

Alfalfa is widely used in foods and is listed by the Council of Europe as a source of natural food flavouring (categories N2 and N3). These categories indicate that alfalfa can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\(^{(G16)}\) In the USA, alfalfa is listed as GRAS (Generally Recognised As Safe).\(^{(G41)}\)
Herbal Use

The herb was not valued by ancient civilisations and is not detailed in classical herbals. Herbal use probably developed in the USA where claims have been made for it in the treatment of arthritis, high cholesterol, diabetes and peptic ulcers.\(^{(2, G19 G32)}\) Reputedly, the herb has bactericidal, cardiotonic, diuretic, emetic, emmenagogue and oestrogenic properties.\(^{(2)}\) Commercial preparations including teas, tablets and capsules are available.\(^{(G19)}\) Alfalfa is stated to be a source of vitamins A, C, E and K, and of the minerals calcium, potassium, phosphorus and iron. It has been used for avitaminosis A, C, E or K, hypoprothrombinaemic purpura, and debility of convalescence.\(^{(G7 G64)}\)
Dosage

**Dried herb**
5–10 g as an infusion three times daily.\(^{(G7)}\)

**Liquid extract**
5–10 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

*In vitro* and animal studies

Alfalfa top (stem and leaves) saponins have been reported to decrease plasma cholesterol concentrations without changing high-density lipoprotein (HDL) cholesterol concentrations, decrease intestinal absorption of cholesterol, increase excretion of neutral steroids and bile acids, prevent atherosclerosis and induce the regression of atherosclerosis.\(^{(3)}\)

Hypocholesterolaemic activity has been reported for root saponins, when given to monkeys receiving a high-cholesterol diet.\(^{(4)}\) Alfalfa herb fed to monkeys reduced hypercholesterolaemia and atherosclerosis; the effect may be partially due to the saponin constituents.\(^{(G19)}\) In mice fed with alfalfa (6.25% of diet) for 12 days before administration of streptozotocin, hyperglycaemia was reduced compared with values for control animals.\(^{(5)}\)

Oestrogenic activity in ruminants has been documented for coumestrol and the isoflavone constituents.\(^{(G22 G41)}\)

An investigation into the effect of various herbs on hepatic drug metabolising enzymes in the rat, showed that alfalfa potentiated the activity of aminopyrine \(N\)-demethylase but had no effect on glutathione \(S\)-transferase or epoxide hydrolase activities.\(^{(6)}\)

The seeds are reported to contain trypsin inhibitors.\(^{(G41)}\) Saponins isolated from the aerial parts have been reported to stimulate the lipolytic activity of neopancreatinum (a mixture of porcine pancreatic enzymes including lipase, amylase and proteases).\(^{(7)}\)

Alfalfa root saponins have been documented to exhibit selective toxicity towards fungi.\(^{(1,8,9)}\) A medicagenic acid glycoside with low haemolytic activity, isolated from alfalfa root, was found to exhibit both strong inhibitory and fungitoxic activities towards several medically important yeasts including *Candida* species, *Torulopsis* species, *Geotrichum canadidum* and *Rhodotorula glutinis*.\(^{(8)}\) It has been proposed that the antimycotic activity of alfalfa saponins is related to their ability to complex steroids and that fungi sensitive to the saponins may contain relatively more steroids in their membranes.\(^{(8)}\) Antifungal properties have also been documented for medicago.\(^{(G41)}\)

The saponin constituents are documented to be haemolytic and to interfere with vitamin E utilisation, and are believed to be one of the causes of ruminant bloat.\(^{(G41)}\) Haemolytic activity is associated with the medicagenic acid glycosides and not the hederagenin and soyasapogenol glycosides.
The effects of polysaccharides from medicago on mice lymphocytes in vitro indicated immunopotentiating activity.\(^{(10)}\)

**Clinical studies**

In a short–term study involving three normolipidaemic individuals given alfalfa seeds (80–60 g daily), serum cholesterol concentrations were reported to be reduced.\(^{(G19)}\) In another small study in which heat–treated alfalfa seeds (40 g three times daily for eight weeks) were taken by eight type-IIA hyperlipoproteinaemic patients and three type IIB patients, a significant decrease was noted in total serum cholesterol concentrations, low–density lipoprotein (LDL) cholesterol and apolipoprotein B. The LDL cholesterol concentration fell by less than 5% in two of the 11 patients.\(^{(11)}\)

The manganese content of alfalfa (45.5 mg/kg) is reported to be the active principle responsible for a hypoglycaemic effect documented for the herb.\(^{(12)}\) A diabetic patient, treated with soluble insulin but poorly controlled, found that an alfalfa extract adequately controlled his diabetes. When administered separately, only small doses of manganese chloride (5–10 mg) were required to have a hypoglycaemic effect. However, no effect was seen on the blood sugar concentrations of non–diabetic controls or of other diabetic patients, who were also administered manganese. It was concluded that manganese lowered the blood sugar concentration in this particular diabetic patient because he was unable to utilise manganese stored in his body.\(^{(12)}\)
Side-effects, Toxicity

Both alfalfa seed and herb have been reported to induce a systemic lupus erythematosus (SLE)-like syndrome in female monkeys.\(^{3,13,G19,G32}\) This activity has been attributed to canavanine, a non–protein amino acid constituent which has been found to have effects on human immunoregulatory cells in vitro.\(^{14}\) Reactivation of quiescent SLE in humans has been associated with the ingestion of alfalfa tablets which, following analysis, were found to contain canavanine.\(^{15}\) It was not stated whether the tablets contained seed or herb material. Canavanine is known to be toxic to all animal species because it is a structural analogue of arginine and may interfere with the binding of this amino acid to enzymes and its incorporation into proteins.\(^{16,G19}\) Alfalfa seeds are reported to contain substantial quantities of canavanine (8.33–13.6 mg/kg), whereas the herb is stated to contain amounts that are considerably less.\(^{16,17}\)

Pancytopenia has been associated with human ingestion of ground alfalfa seeds (80–160 g/day), which were taken to lower plasma cholesterol concentrations.\(^{18}\)

Dietary studies using alfalfa top saponins (ATS) in the diet of rats and monkeys showed no evidence of toxicity and serum lipid concentrations were lowered.\(^{3,19,20}\) In addition, when ATS were given to cholesterol–fed animals, a reduction in serum lipid concentrations was observed.\(^{3,19,20}\) ATS are reported to be free of the SLE-inducing substance that is present in the seeds.\(^3\)

Negative results were documented for alfalfa when tested for mutagenicity using Salmonella strains TA98 and TA100.\(^{21}\)
Contra–indications, Warnings

Individuals with a history of SLE should avoid ingesting alfalfa. Ingestion of large amounts of alfalfa (exceeding amounts normally consumed in the diet) should be avoided in view of the documented oestrogenic activity and potential anticoagulant activity. Excessive doses may interfere with anticoagulant therapy and with hormonal therapy, including the oral contraceptive pill and hormone replacement therapy. Alfalfa may affect blood sugar concentrations in diabetic patients because of the manganese content.

Pregnancy and lactation

Alfalfa seeds are reputed to affect the menstrual cycle and to be lactogenic.\(^{(G30)}\) Although the safety of alfalfa herb has not been established, it is probably acceptable for use during pregnancy and lactation provided that doses do not exceed the amounts normally ingested as a food. Alfalfa seeds should not be ingested during pregnancy or lactation.
Pharmaceutical Comment

The chemistry of alfalfa is well documented and it does appear to be a good source of vitamins and minerals, thereby supporting the herbal uses. However, normal human dietary intake of alfalfa is low and excessive ingestion should be avoided in view of the many pharmacologically active constituents (e.g. canavanine, coumarins, isoflavones and saponins), which may give rise to unwanted effects if taken to excess. Oestrogenic effects are generally associated with the ingestion of large amounts of the herb, such as in fodder for poultry and cattle. Reports of a possible SLE-inducing capacity for alfalfa, particularly the seeds, also suggests that excessive ingestion is not advisable. In view of the reports of arthralgia, alfalfa should not be recommended for the treatment of arthritis.
References

See also General References G5 G9 G16 G19 G22 G24 G30 G31 G32 G36 G37 G41 G43 G64.

15. Roberts JL, Hayashi JA. Exacerbation of SLE associated with alfalfa


Aloe Vera
Aloe barbadensis Mill., Aloe ferox Mill. and hybrids with Aloe africana Mill. and Aloe spicata Baker (Liliaceae)
Synonyms
Aloe Gel, Aloe vera Tourn. ex L., Aloe vera (L.) Webb
Part Used
Leaf gel
Pharmacopoeial and Other Monographs

Martindale 33rd edition($G67$)

PDR for Herbal Medicines 2nd edition($G36$)

WHO volume 1 1999($G63$)
Legal Category (Licensed Products)

Aloe vera is not included in the GSL.
Constituents


Aloe vera is reported to contain mono- and poly saccharides, tannins, sterols, organic acids, enzymes (including cyclooxygenase),\(^1\) saponins, vitamins and minerals.\(^2\)

**Carbohydrates**

Glucomannan and other polysaccharides containing arabinose, galactose and xylose.

**Lipids**

Includes cholesterol, gamolenic acid and arachidonic acid.\(^1\) The polar, non–polar and fatty acid composition has been investigated.\(^1\)
Food Use

Aloe vera is not used in foods.
Herbal Use

Traditionally, aloe vera has been used in ointments and creams to assist the healing of wounds, burns, eczema and psoriasis. (G2 G6 G41 G64)
Dosage

None documented.
Pharmacological Actions

Aloe vera refers to the mucilaginous tissue located in the leaf parenchyma of Aloe vera or related Aloe species. However, many documented studies for Aloe vera have utilised homogenised leaf extracts which therefore combine aloe vera with aloes, the laxative preparation obtained from the bitter, yellow juice also found in the leaf (see Aloes). Unless otherwise specified, the following studies will refer to a total leaf extract.

In vitro and animal studies

Gel preparations have been reported to be effective against radiation burns, skin ulcers and peptic ulcers.\(^{(2)}\) However, the gel was also found to be ineffective against drug- and stress–induced gastric and peptic ulcers in rats.\(^{(2)}\)

Anti–inflammatory activity has been observed in various rat and mouse models that received subcutaneous injections of Aloe vera leaf extract.\(^{(3)}\) A positive response was noted in wound–healing (10 mg/kg, rat; 100 mg/kg, mouse), mustard oedema (10 mg/kg, rat) and polymorphonuclear leukocyte infiltration (2 mg/kg, mouse) tests, although no activity was demonstrated in the antifibrosis test (cotton pellet granuloma) (400 mg/kg, rat).

Anti–arthritic and anti–inflammatory activity has been documented for a cream containing homogenised Aloe africana leaves, ribonucleic acid, and ascorbic acid, following topical application to rats which had been injected (day 0) with Mycobacterium butyricum to cause adjuvant arthritis.\(^{(4)}\) This model is considered a good experimental tool for studying rheumatoid arthritis.\(^{(4)}\) The cream was found to be active when applied both as a prevention (days 1–13) and as a regression (days 21–35) treatment.\(^{(4)}\) Subsequent work suggested that anthraquinone compounds (anthraquinone, anthracene and anthranilic acid) may be the active components in the aloe leaf mixture.\(^{(5)}\) These compounds are, however, constituents of aloes rather than aloe vera (see Aloes). Aloe vera juice (presumably containing the anthraquinones contained in aloe preparation) has been applied directly to open pressure sores to assist in their healing.\(^{(6)}\) The aloe vera extract exhibited an anaesthetic reaction, antibacterial action and increased local microcirculation.\(^{(6)}\)

Endogenous cyclooxygenase in Aloe vera has been found to convert endogenous arachidonate to various prostanoids, namely PGE\(_2\) (major), TXB\(_2\), PGD\(_2\), PGF\(_{2\alpha}\), and 6–keto-PGF\(_{1\beta}\).\(^{(1)}\) The production of these compounds, especially PGE\(_2\), has been associated with the beneficial effect of an aloe
Hypoglycaemic actions have been documented for aloes extracts (see Aloes).

Clinical studies

Enhancement of phagocytosis in adult bronchial asthma has been attributed to a non-dialysable fraction of the extract, consisting of active components that are a mixture of polysaccharide and protein or glycoprotein. Despite the nature of these proposed active components, it has been proposed that activity of the fraction may be related to the previous observation that aloe vera synthesises prostaglandins from endogenous arachidonic acid using endogenous cyclooxygenase. In this current study, activity of the aloe vera extract required dark storage at 4–30°C for a period of 3–10 days. These conditions are reported to be favourable for the hydrolysis of phospholipids, thus releasing arachidonic acid for synthesis of prostanoids. In addition, activity was dependent on patients not having received prior treatment with a corticosteroid. The gel has been reported to be effective in the treatment of mouth ulcers.
Side-effects, Toxicity

None documented.
Contra–indications, Warnings

Hypoglycaemic activity has been documented for an aloe vera extract, although it is unclear whether this is associated with the true aloe vera gel or aloe extract.\(^9\)

Pregnancy and lactation

The external application of aloe vera gel during pregnancy is not thought to be any cause for concern. However, products stated to contain aloe extracts or aloe vera may well contain gastrointestinal stimulant anthraquinone components that are well recognised as the active constituents in aloe (laxative). As such, ingestion of such preparations during pregnancy and lactation should be avoided.
Aloe vera is obtained from the mucilaginous tissue in the centre of the *Aloe vera* leaf and consists mainly of polysaccharides and lipids. It should not be confused with aloes, which is obtained by evaporation of water from the bitter yellow juice that is drained from the leaf. Unlike aloes, aloe vera does not contain any anthraquinone compounds and does not, therefore, exert any laxative action. Studies have reported an anti-inflammatory and anti-arthritic action for total leaf extracts but the activity seems to be associated with anthraquinone compounds. Hypoglycaemic activity has been reported for aloe vera extract. Aloe vera is a source of gamolenic acid. The literature on burn management with aloe vera gel preparations is confused and further studies are required.\(^{10}\)
References

See also General References G5 G6 G18 G19 G22 G29 G31 G32 G36 G41 G43 G63 G64.


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Species (Family)

i. *Aloe barbadensis* Mill. (Liliaceae)

ii. *Aloe ferox* Mill. and its hybrids with *Aloe africana* Mill. and *Aloe spicata* Baker
Synonym(s)

Part(s) Used

Dried leaf juice
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)
BHP 1996\(^{(G9)}\)
BP 2002\(^{(G71)}\)
Complete German Commission E\(^{(G3)}\)
ESCOP 1997 (Cape Aloes)\(^{(G52)}\)
Martindale 33rd edition\(^{(G67)}\)
PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Ph Eur 2004\(^{(G72)}\)
USP26/NF21\(^{(G73)}\)
WHO volume 1 1999\(^{(G63)}\)
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents

See General References G2 G20 G22 G41 G52 G64.

**Anthranoids**(1–6,G20 G52)
Cape aloes anthranoids are qualitatively identical to leaf exudate of *A. ferox*. Anthrones (up to 30%), mainly the C-glycosides aloins A and B (= barbaloin, isobarbaloin, stereo isomers of 10–glucosyl–aloe–emodin anthrone); other glycosides include 8-O-methyl–7–hydroxy aloins A and B, aloinosides A and B (aloin–11-O-α-L-rhamnosides). Small quantities of 1,8–dihydroxyanthra quinoid aglycones, including aloe–emodin and chrysophanol are also present.

**Chromones**(1,2,7,G20 G52)
Major constituents are aloesin (2–acetonyl–5–methyl–8–glucosyl chromone) and aloeresin E. Lesser quantities of isoaloeresin D, 8-C-glucosyl–7-O-methyl–aloesol and related glycosides which may be esterified at the glucose moiety by either cinnamic, *p*-coumaric or ferulic acids, are also present. Non-glycosylated chromones include 7–hydroxy–2,5–dimethylchromone, furoaloesone, 2–acetonyl–7–hydroxy–8-(3–hydroxyacetonyl)-5–methyl chromone and 2–acetonyl–8-(2–furoylmethyl)-7–hydroxy–5–methylchromone.(8,9)

**Phenyl pyrones**(1,2)
Glycosides include aloenin and aloenin B.

**Other constituents**(G52)
Cinnamic acid and 1–methyltetralin.
Food Use

Aloes is listed by the Council of Europe as a natural source of food flavouring (category N3).\(^{(G16)}\) This category indicates that aloes can be added to foodstuffs in the traditionally accepted manner, although there is insufficient information available for an adequate assessment of potential toxicity. The concentration of aloin present in the final product is limited to 0.1 mg/kg; 50 mg/kg in alcoholic beverages.\(^{(G16)}\) In the USA, aloes is listed as GRAS (Generally Recognised As Safe).\(^{(G41)}\)
Herbal Use

Aloes is recommended for the treatment of atonic constipation and suppressed menstruation. (G2 G49 G64)
Dosage

Dried juice

50–200 mg or equivalent three times daily.\(^{(G6,G49)}\)

In view of potential adverse effects, the dose recommended for adults and children aged over 10 years is 10–30 mg of hydroxyanthracene derivatives (calculated as barbaloin) once daily at night.\(^{(G52)}\) Use of aloes as a laxative in self-treatment of constipation for more than two weeks is not recommended.\(^{(G52)}\)
Pharmacological Actions

The activity of aloes can be attributed to the anthranoid glycoside content. The glycosides are metabolised by glycosidases in the intestinal flora to form active anthrones. The laxative action is due to an increase in motility of the large intestine by inhibition of the Na\(^+\)/K\(^+\) pump and chloride ion channels; enhanced fluid secretion occurs due to stimulation of mucus and chloride ion secretion.\(^{G52}\)

In vitro and animal studies

Nine hours after oral administration, aloes produced diarrhoea at doses of 5 g/kg (in 20% of rats) and 20 g/kg (in 100% of rats).\(^{10}\) Pretreatment of rats with the nitric oxide (NO) synthase inhibitor N-nitro-L-arginine methyl ester given intraperitoneally reduced diarrhoea induced by aloes (20 g/kg) 9 hours after oral administration. The results suggest that endogenous NO modulates the diarrhoeal effects of Cape aloes.\(^{10}\)

Inhibitory effects of aqueous extracts of five genera of Aloe, including A. ferox and A. barbadensis, and aloe powder (Japanese Pharmacopoeia) on histamine release from rat peritoneal mast cells induced by antigen were investigated in vitro.\(^{11}\) All extracts tested inhibited histamine release in a concentration-dependent manner under the test conditions. Aloe ferox extract, Japanese Pharmacopoeia aloes and barbaloin strongly inhibited histamine release (IC\(_{50}\) 0.16, 0.07 and 0.02 μg/mL, respectively).\(^{11}\)

Aqueous extracts of aloes are said to elevate the rate of ethanol oxidation in vivo.\(^{12}\) Oral administration of aloin (300 mg/kg) to rats 12 hours prior to administration of alcohol (3 g/kg) resulted in a significant decrease (40%) in blood alcohol concentration.\(^{12}\) Pretreatment with intraperitoneal aloe–emodin 2 hours prior to alcohol administration significantly reduced blood alcohol concentrations; it was hypothesised that aloin is metabolised to aloe–emodin which exerts its effect on alcohol metabolism.\(^{12}\) Activity–guided fractionation of the leaves of A. aborescens resulted in the isolation and characterisation of elgonica–dimers A and B (dimeric C-glycosides of anthrone emodin–10′-C-β-D-glucopyranoside and aloe–emodin) as potent inhibitors of cytosolic alcohol dehydrogenase and aldehyde dehydrogenase activities in vitro.\(^{13}\)

Aloe–emodin and an alcoholic extract of aloes have been reported to possess antitumour activity.\(^{G41}\)

Hypoglycaemic activity has been shown in alloxan–diabetic mice for aloes\(^{14}\)
Barbaloin is active *in vitro* against *Mycobacterium tuberculosis* and *Bacillus subtilis* (minimum inhibitory concentration 0.125 mg/mL and 0.25 mg/mL, respectively).\(^{(G18)}\)

**Clinical studies**

The purgative action of the anthraquinone glycosides is well recognised (*see* Senna), although aloes is reported to be more potent than both senna and cascara.\(^{(G41,G45)}\) Orally ingested anthranoid glycosides are not metabolised until they reach the colon. In humans, the intestinal flora break down *O*-glycosides readily and *C*-glycosides to some extent. The main active metabolite is aloe–emodin–9–anthrone.\(^{(G52)}\)

An aloes extract in doses too small to cause abdominal cramps or diarrhoea had a significant hypoglycaemic effect in five patients with non–insulin–dependent diabetes.\(^{(14)}\)
Side-effects, Toxicity

Aloes is a potent purgative that may cause abdominal pains, gastrointestinal irritation leading to pelvic congestion and, in large doses, may result in nephritis, bloody diarrhoea and haemorrhagic gastritis.\(^{(G41 G44)}\) Like all stimulant purgatives, prolonged use of aloes may produce watery diarrhoea with excessive loss of water and electrolytes (particularly potassium), muscular weakness and weight loss.\(^{(G44)}\)

Tests of the possible carcinogenicity of hydroxyanthraquinones and their glycosides showed that exposure to certain aglycones and glycosides may represent a human cancer risk.\(^{(17)}\) Most of the aglycones tested were found to be mutagenic and some, such as emodin and aloe–emodin, were genotoxic in mammalian cells.

Administration of dried aloes extract 50 mg/kg per day for 12 weeks to mice did not result in the development of severe pathological symptoms, although a raised sorbitol dehydrogenase concentration was suggested to be indicative of liver damage.\(^{(G20)}\) No mutagenic effects in *Salmonella typhimurium* and Va7 cells, or DNA repair induction in rat hepatocytes, were observed. These negative results are due to the inability of the test systems to release mutagenic anthranoids from the C-glycosides.\(^{(G20)}\) Retrospective and prospective studies have shown no causal relationship between anthranoid laxative use and colorectal cancer.\(^{(G20)}\)
Contra–indications, Warnings

Aloes has been superseded by less toxic laxatives.\(^{G45}\) The drastic purgative action of aloes contra–indicates its use in individuals with haemorrhoids and existing kidney disease. Hypokalaemia resulting from laxative abuse potentiates the action of cardiac glycosides, interacts with anti–arrhythmic drugs, and with drugs which induce reversion to sinus rhythm, e.g. quinidine. Concomitant use with thiazide diuretics, adrenocorticosteroids and liquorice may aggregate electrolyte imbalance. In common with all purgatives, aloes should not be given to patients with inflammatory disease of the colon (e.g. Crohn’s disease, ulcerative colitis), appendicitis, intestinal obstruction, abdominal pain, nausea or vomiting. Aloes colours alkaline urine red. Long–term use should be avoided and a doctor should be consulted within two weeks of treatment initiation if symptoms persist.

_Pregnancy and lactation_

In view of the irritant and cathartic properties documented for aloes, its use is contra–indicated during pregnancy.\(^{G44}\) Anthraquinones may be secreted into breast milk and, therefore, aloes should be avoided during lactation (see Senna).

Aloes is reputed to be an abortifacient and to affect the menstrual cycle.\(^{G22}\)
Aloes and aloe gel are often confused with each other. Aloes is obtained by evaporation of water from the bitter yellow juice drained from the leaves of A. vera. Commercial ‘aloin’ is a concentrated form of aloes.\(^{(G41)}\) Aloe gel is prepared by many methods, but is obtained from the mucilaginous tissue in the centre of the leaf and does not contain anthraquinones (see Aloe Vera). Aloes is a potent purgative which has been superseded by less toxic drugs such as senna and cascara. Generally, the use of unstandardised preparations containing anthraquinone glycosides should be avoided, since their pharmacological effect is unpredictable and they may cause abdominal cramp and diarrhoea. In particular, the use of products containing combinations of anthraquinone laxatives should be avoided.
References


Angelica
Species (Family)

*Angelica archangelica* L. (Apiaceae/Umbelliferae)
Synonym(s)

Archangelica officinalis Moench and Hoffm.
Part(s) Used
Fruit, leaf, rhizome, root
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)

BP 2002\(^{(G71)}\)

Complete German Commission E\(^{(G3)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)

Ph Eur 2004 (root)\(^{(G72)}\)
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents

See General References G2 G22 G32 G41 G48 G57 G58 G64.

The literature mainly refers to constituents of the root.

**Coumarins**
Over 20 furanocoumarins, including angelicin, archangelicin, bergapten, isoimperatorin and xanthotoxin.\(^{(1,G2)}\) Also the coumarins osthol (major constituent in rhizome/root, 0.2%) and umbelliferone.\(^{(G2)}\) The root also contains the furanocoumarins 2′-angeloyl-3′-isovaleryl vaginate,\(^{(2)}\) heraclenol-2′-O-senecioate and heraclenol-2′-O-isovalerate.\(^{(3)}\)

**Volatile oils**
0.35–1.3% in root and fruit. 80–90% monoterpenes, including α- and β-phellandrene, α- and β-pinene, sabinene, α-thujene, limonene, linalool, borneol\(^{(1,4)}\) and four macrocyclic lactones.

**Other constituents**
Archangelone (a flavonoid), palmitic acid, caffeic and chlorogenic acids, sugars (fructose, glucose, sucrose, umbelliferose).
Angelica is widely used in foods. Angelica is listed by the Council of Europe as a natural source of food flavouring (stem: category 1; other parts and preparations: category 4, with limits on coumarin and furanocoumarin) (see Appendix 23). In the USA, angelica is listed as GRAS (Generally Recognised As Safe).
Herbal Use\textsuperscript{(1,G32)}

Angelica is stated to possess antispasmodic, diaphoretic, expectorant, bitter aromatic, carminative, diuretic and local anti-inflammatory properties. It has been used for respiratory catarrh, psychogenic asthma, flatulent dyspepsia, anorexia nervosa, rheumatic diseases, peripheral vascular disease, and specifically for pleurisy and bronchitis, applied as a compress, and for bronchitis associated with vascular deficiency.\textsuperscript{(G2 G7 G49 G64)} The German Commission E monograph states that angelica can be used for lack of appetite and dyspeptic complaints such as mild stomach cramps and flatulence.\textsuperscript{(G4)} Many related species, including \textit{A. sinensis} (dong quai) are traditionally used in Chinese medicine.\textsuperscript{(G57)}
**Dosage**

**Dried leaf**
2–5 g by infusion three times daily.\(^{(G7)}\)

**Leaf liquid extract**
2–5 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)

**Leaf tincture**
2–5 mL (1 : 5 in 45% alcohol) three times daily.\(^{(G7)}\)

**Dried rhizome/root**
Daily dose 4.5 g\(^{(G2)}\) or 1–2 g by infusion three times daily.\(^{(G4, G7)}\)

**Rhizome/root liquid extract**
0.5–2.0 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)

**Rhizome/root tincture**
0.5–2 mL (1 : 5 in 50% alcohol) three times daily.\(^{(G7)}\)

**Fruit**
1–2 g.\(^{(G49)}\)
Pharmacological Actions

In vitro and animal studies

Minimal anti-inflammatory activity (1% inhibition of carrageenan-induced rat paw oedema) has been documented for fruit extracts (100 mg/kg body weight by mouth) given 45 minutes before eliciting oedema. This was compared with 45% inhibition by indometacin (5 mg/kg by mouth). Angelica is reported to possess antibacterial and antifungal properties. Antibacterial activity against Mycobacterium avium has been documented, with no activity exhibited against Escherichia coli, Bacillus subtilis, Streptococcus faecalis or Salmonella typhi. Antifungal activity was reported in 14 of 15 fungi tested.

A methanolic extract of A. archangelica root showed antispasmodic activity against spontaneous contractions of circular smooth muscle (IC$_{50}$ 265 μg/mL) and inhibited acetylcholine- and barium chloride–induced contractions of longitudinal smooth muscle (IC$_{50}$ values 242 and 146 μg/mL, respectively).

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Extracts of A. archangelica root exhibit calcium channel–blocking activity. A series of isolated coumarins showed activity, the most active being archangelicin with an IC$_{50}$ 1.2 μg/mL (verapamil 2.0μg/mL) as a calcium channel antagonist as assessed by inhibition of depolarisation in GH4C1 rat pituitary cells.

Sixteen coumarin compounds isolated from A. archangelica were tested for activity in cyclooxygenase 1 (COX-1) and 5–lipoxygenase (5-LO) inhibition assays in vitro. None of the compounds demonstrated activity against COX-1, but osthol and oxypeucedanin isovalerate were active in the 5-LO assay.

In rabbits, a uterotonic action has been documented for Japanese angelica root following intraduodenal administration of a methanolic extract (3 g/kg). A. sinensis is reported to have induced uterine contraction and relaxation.

Clinical studies

None documented for angelica (A. archangelica). The furanocoumarin constituent bergapten (5–methoxypsoralen) has been used in the PUVA (psoralen (P) and high-intensity long-wavelength ultraviolet irradiation) treatment of psoriasis.
Side-effects, Toxicity

Both angelica and the root oil have been reported to cause photodermatitis and phototoxicity, respectively, following external contact.\(^{(G32)}\) Angelica contains furanocoumarin constituents which are known to cause photosensitisation. Concern has been expressed at the possible carcinogenic risk of the furanocoumarin bergapten. Seven species of plants known to cause dermatitis were analysed for psoralen, 8–methoxypsoralen (xanthotoxin) and 5–methoxypsoralen (bergapten). The highest total yield was obtained from \(A.\ archangelica\).\(^{(12)}\)

The root oil has been reported to be non–irritant and non–sensitising on animal and human skin.\(^{(6)}\)

Root and fruit oils obtained by steam distillation are claimed to be devoid of furanocoumarins, although extracts may contain them.\(^{(G41)}\)

Toxicity studies have been documented for the root oil.\(^{(6)}\) Acute LD\(_{50}\) values have been reported as 2.2 g/kg body weight (mouse, by mouth) and 11.16 g/kg (rat, by mouth). Death was attributed to liver and kidney damage, although animals surviving for three days completely recovered with a reversal of organ damage. An acute LD\(_{50}\) (rabbit, dermal) value was reported to be greater than 5 g/kg. Subacute toxicity studies, lasting eight weeks, suggested that the tolerated dose in the rat was 1.5 g/kg, although at lower doses the animals weighed less than the controls.\(^{(6)}\)

Furanocoumarins isolated from a related Chinese species, \(Angelica koreana\), have been reported to affect the hepatic metabolism of hexobarbitone. The compounds were found to cause a marked inhibition of drug metabolism in the first phase and an acceleration in the second phase, and were thought to be drug–metabolising enzyme inhibitors rather than enzyme inducers. Furanocoumarins investigated included imperatorin and oxypeucedanin, which are also documented as constituents of \(A.\ archangelica\). It has been reported that a related Chinese species, \(Angelica sinensis\), may be hepatoprotective and prevent the reduction of hepatic glycogen.
Contra–indications, Warnings

Angelica may provoke a photosensitive allergic reaction because of the furanocoumarin constituents. In addition, excessive doses may interfere with anticoagulant therapy because of the coumarin constituents.

The use of bergapten in cosmetic and suntan preparations is stated to be ill–advised by some regulatory authorities,\(^{(G45)}\) in view of the concerns regarding the risk of skin cancer. The International Fragrance Association recommends that angelica root oil be limited to a maximum of 0.78% in products applied to skin which is then exposed to sunshine.\(^{(G58)}\)

**Pregnancy and lactation**

Angelica root is reputed to be an abortifacient and to affect the menstrual cycle. In view of this and the photosensitising constituents, the use of angelica during pregnancy and lactation in amounts exceeding those used in foods should be avoided.
The chemistry of angelica is well documented. Although the traditional use of Chinese angelica species, such as *A. sinensis* and *A. acutiloba*, is well established in oriental medicine, there is limited documented pharmacological information available for *A. archangelica*, the species most commonly used in Europe, to justify its herbal uses. In view of the presence of known pharmacologically active constituents, especially bergapten, consumption of amounts exceeding normal human dietary intake should be avoided. Angelica contains furanocoumarins which are known to possess photosensitising properties.

The related species *A. sinensis* (dong quai) is popular in traditional Chinese medicine (TCM) and occurs in about 70% of all TCM prescriptions to treat dysmenorrhoea, postnatal disturbances, anaemia, constipation and chronic pelvic infections.\(^{(13)}\) Western naturopaths recommend the use of dong quai in hypertension, for modification of high blood sugar concentrations, regulation of the immune system, liver detoxification, anaemia and to relieve allergic conditions. Several unlicensed over–the–counter (OTC) products containing dong quai are readily available.

The chemistry of dong quai is similar to that of *A. archangelica*, with coumarins and volatile oil being major components.\(^{(13)}\) In addition, a series of phthalides (e.g. ligustilide, butylphthalide, butylidenephthalide) have been isolated. Pharmacological investigations have shown that phthalides and coumarins have antispasmodic activity. The volatile constituents generally exert a hypotensive effect. A polysaccharide component is active against Ehrlich ascites tumours in mice and has immunostimulating activity,\(^{(14)}\) and protects the gastric mucosa against ethanol- and indometacin–induced damage.\(^{(15)}\) Clinical investigation has failed to support the claims for relieving menopausal symptoms.\(^{(16)}\) *A. sinensis* has been reported to be effective in improving abnormal protein metabolism in patients with chronic hepatitis or hepatic cirrhosis.\(^{(17)}\)

The furanocoumarins are phototoxic and have photocarcinogenic potential, but need ultraviolet (UV) light for activation. An extract of dong quai administered subcutaneously to rabbits did not affect prothrombin time given alone, but did after concurrent administration of a single dose of warfarin.\(^{(18)}\) Elevation of prothrombin time was noted in a patient stabilised on warfarin who began taking dong quai.\(^{(19)}\) Coagulation values returned to normal one month after discontinuing use of dong quai.
References

See also General References G2 G3 G9 G10 G16 G22 G31 G33 G36 G37 G41 G44 G48 G49 G57 G58 G64.


Aniseed
Species (Family)

*Pimpinella anisum* L. (Apiaceae/Umbelliferae)
Synonym(s)
Part(s) Used

Fruit
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)
BP 2002\(^{(G71)}\)
Complete German Commission E\(^{(G3)}\)
ESCOP 1997\(^{(G52)}\)
Martindale 33rd edition\(^{(G67)}\)
PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Ph Eur 2004\(^{(G72)}\)
Legal Category (Licensed Products)

GSL$^{G37}$
Constituents
See General References G2 G22 G41 G52 G58 G64.

Coumarins
Scopoletin, umbelliferone, umbelliprenine; bergapten (furanocoumarin).

Flavonoids
Flavonol (quercetin) and flavone (apigenin, luteolin) glycosides, e.g. quercetin–3–glucuronide, rutin, luteolin–7–glucoside, apigenin–7–glucoside; isoorientin and isovitexin (C-glucosides).

Volatile oils
2–6%. Major components are trans-anethole (80–95%), with smaller amounts of estragole (methyl chavicol), anise ketone (p-methoxyphenylacetone) and β-caryophyllene. Minor components include anisaldehyde and anisic acid (oxidation products of anethole), linalool, limonene, α-pinene, pseudoisoegenol–2–methyl butyrate, acetaldehyde, p-cresol, cresol, hydroquinone, β-farnesene, α-, β- and γ-himachalene, bisabolene, d-elemene, ar-curcumene and myristicin.(1)

Other constituents
Carbohydrate (50%), lipids 16% (saturated and unsaturated), β-amyrin (triterpene), stigmasterol (phytosterol) and its palmitate and stearate salts.
Food Use

Aniseed is used extensively as a spice and is listed by the Council of Europe as a natural source of food flavouring (category N2). This category allows small quantities of aniseed to be added to foodstuffs, with a possible limitation of an active principle (as yet unspecified) in the final product.\(^{(G16)}\) In the USA, aniseed is listed as GRAS (Generally Recognised As Safe).\(^{(2,G41)}\)
Herbal Use

Aniseed is stated to possess expectorant, antispasmodic, carminative and parasiticide properties. Traditionally, it has been used for bronchial catarrh, pertussis, spasmodic cough, flatulent colic; topically for pediculosis and scabies; its most specific use is for bronchitis, tracheitis with persistent cough, and as an aromatic adjuvant to prevent colic following the use of cathartics.\(^{(G2 \ G7 \ G64)}\)

Aniseed has been used as an oestrogenic agent.\(^{(3)}\) It has been reputed to increase milk secretion, promote menstruation, facilitate birth, alleviate symptoms of the male climacteric and increase libido.\(^{(3)}\)
**Dosage**

**Dried fruit**
Adults: 1.0–5.0 g crushed fruits in 150 mL water as an infusion several times daily.\(^{(G52)}\) Children: 0–1 year old, 1.0 g of crushed fruits as an infusion; 1–4 years of age, 2.0 g; over 4 years, use adult dose.\(^{(G52)}\)

**Oil**
0.05–0.2 mL three times daily.\(^{(G7)}\)

**Spirit of anise**
(BPC 1949) 0.3–1.0 mL three times daily.

**Distilled anise water**
(BPC 1934) 15–30 mL three times daily.
Pharmacological Actions

The pharmacological effects of aniseed are largely due to the presence of anethole, which is structurally related to the catecholamines adrenaline, noradrenaline and dopamine. Anethole dimers closely resemble the oestrogenic agents stilbene and diethylstilbestrol.\(^3\)

**In vitro and animal studies**

**Antimicrobial, antifungal and insecticidal activities**

The volatile oil has antibacterial, antifungal and insecticidal activities.\(^{G41}G52\)

Anethole, anisaldehyde and myristicin have exhibited mild insecticidal properties.\(^{G41}\)

**Antispasmodic activity**

Anise oil (200 mg/L) was shown to antagonise carbachol–induced spasms in a guinea–pig tracheal muscle preparation.\(^{G52}\)

**Secretolytic and expectorant effect**

Application of aniseed (6.4 g/140 mL) to isolated ciliated epithelium of frog trachea induces small increases in transport velocity.\(^{G52}\)

Dilutions of anise oil increased respiratory tract fluid in anaesthetised guinea–pigs, rats and cats. A similar action was observed in anaesthetised rabbits inhaling anise oil.\(^{G52}\)

The reputed lactogogic action of anise has been attributed to anethole, which exerts a competitive antagonism at dopamine receptor sites (dopamine inhibits prolactin secretion), and to the action of polymerised anethole, which is structurally related to the oestrogenic compounds stilbene and stilboestrol.\(^3\)

**CNS activities**

Whole plant aqueous infusions have been reported to delay (but not prevent) the onset of picrotoxin–induced seizures and to reduce the mortality rate in mice following intraperitoneal injection.\(^4\)

Aniseed has also been found to slightly elevate γ-aminobutyric acid (GABA) concentrations in brain tissue.\(^4\)

The anticonvulsant effect is much weaker with aniseed than with conventional drug treatment and therefore its use as an anticonvulsant in Arabic folklore is not supported.\(^4\)

Anise oil diluted in sesame oil (0.25–1.0 mL/kg) given intraperitoneally to mice increased in a dose–dependent manner the dose of pentylenetetrazole needed to induce clonic and tonic seizures.\(^5\)

Activity was also observed against tonic seizures induced by maximal electric shock. Motor impairment was observed at higher doses of anise oil. Pentobarbital–induced sleeping time was prolonged by intraperitoneal administration of anise oil.
(50 mg/kg) to mice.\(^{(52)}\)

**Other activities**

Oral administration of anethole (250–1000 mg/kg) to Swiss albino mice with Ehrlich ascites tumour in the paws indicated antitumour activity.\(^{(6)}\) The conclusions were based on biochemical changes (nucleic acids, proteins, malondialdehyde, glutathione), survival rate and tumour weight. Anise oil given to rats (100 mg/kg given subcutaneously) stimulated liver regeneration after partial hepatectomy.\(^{(52)}\)

**Clinical studies**

Aniseed is mainly used for the treatment of dyspeptic complaints and catarrh of the upper respiratory tract.\(^{(2} G^{41} G^{52})\) There is a lack of documented clinical studies with aniseed.
Contact dermatitis reactions to aniseed and aniseed oil have been attributed to anethole.\(^7\) Reactions have been reported with products, such as creams and toothpastes, flavoured with aniseed oil.\(^5\) The volatile oil and anethole have been stated to be both irritant and sensitising.\(^3\) Two female workers in a cake factory developed severe dermatitis, and patch tests indicated sensitivity to anise oil and to anethole.\(^8\) Soreness, dryness and cracking of lips and perioral skin occurred in an individual using a herbal (fennel) toothpaste; anethole was reported to be the sensitising agent.\(^9\)

Bergapten is known to cause photosensitivity reactions and concern has been expressed over the possible carcinogenic risk of bergapten.\(^{10}\)

Ingestion of as little as 1–5 mL of anise oil can result in nausea, vomiting, seizures, and pulmonary oedema.\(^7\)

The LD\(_{50}\) values per kg body weight for anise oil and trans-anethole are 2.7 g and 2–3 g, respectively.\(^5\) Mild liver lesions were observed in rats fed repeated anethole doses (695 mg/kg) for an unspecified duration.\(^3\) Hepatic changes have been described in rats fed anethole in their daily diet (1%) for 15 weeks,\(^2\) although at a level of 0.25% there were no changes after one year. Rats fed with 0.1% trans-anethole in their diet for 90 days showed no toxic effects, but higher concentrations (0.3%, 1.0% and 3.0%) resulted in liver oedema.\(^5\) In therapeutic doses, anethole is reported to cause minimal hepatotoxicity.\(^2\) Trans-anethole given orally to rats (50–80 mg/kg) resulted in dose-dependent anti-implantation activity.\(^{10}\) Significant oestrogenic activity was observed, but no anti-oestrogenic, progestational, anti-progestational, androgenic or antiandrogenic activity.\(^{10}\)

Oral administration (1% of diet) of trans-anethole to rats resulted in induction of parathion-degrading drug enzymes.\(^5\) Male Wistar rats were treated with trans-anethole (125 or 250 mg/kg) by gavage for 10 days and the activities of liver microsome and cytosol phase I and II biotransformation enzymes were determined.\(^{11}\) There was no effect on cytochrome P450, but UDP-glucuronyltransferase activity in the cytosol towards the substrates 4-chlorophenol and 4-hydroxyphenol was significantly increased for both doses. It was concluded that trans-anethole preferentially induces phase II biotransformation in rat liver \textit{in vivo}.\(^{11}\)

The safety of trans-anethole (4-methoxy propenylbenzene) has been reviewed by the Expert Panel of the Flavour and Extract Manufacturer Association (FEMA).\(^2\) The evaluation was based on whether the hepatotoxic metabolite anethole epoxide is produced. At low levels of exposure, trans-
anethole is efficiently detoxicated in rodents and humans, primarily by O-demethylation and ω-oxidation, respectively, while epoxidation is only a minor pathway. At higher doses in rats, a metabolic shift occurs resulting in epoxidation and formation of anethole epoxide. The continuous intake of high doses of trans-anethole induces a continuum of cytotoxicity, cell necrosis and cell proliferation. In chronic dietary studies in rats, hepatotoxicity resulted when the daily production of anethole epoxide exceeded 30 mg anethole epoxide per kg body weight. Neither trans-anethole nor anethole epoxide showed any evidence of genotoxicity. The Expert Panel concluded that the hepatocarcinogenic effects in female rats occur via a non-genotoxic mechanism and are secondary to hepatotoxicity caused by continuous exposures to high hepatocellular concentrations of anethole epoxide. Trans-anethole was reaffirmed as GRAS, based on a thorough study of the scientific literature. Because trans-anethole undergoes efficient metabolic detoxication in humans at low levels of exposure, the neoplastic effects in rats associated with dose-dependent hepatotoxicity are not indicators of any significant risk to human health from the use of trans-anethole as a flavouring substance.
Contra-indications, Warnings

Aniseed may cause an allergic reaction. It is recommended that the use of aniseed oil should be avoided in dermatitis, and inflammatory or allergic skin conditions.\(^{(G31 \ G58)}\) Aniseed should be avoided by persons with known sensitivity to anethole.\(^{(G52)}\) Bergapten may cause photosensitivity in sensitive individuals. The documented oestrogenic activity of anethole and its dimers may affect existing hormone therapy, including the oral contraceptive pill and hormone replacement therapy, if excessive doses are ingested. In view of the structural similarity reported between anethole and myristicin, consumption of large amounts of aniseed may cause neurological effects similar to those documented for nutmeg.

**Pregnancy and lactation**
Traditionally, aniseed is reputed to be an abortifacient\(^{(G22)}\) and also to promote lactation. The safety of aniseed taken during pregnancy and lactation has not been established; however, there are no known problems provided that doses taken do not greatly exceed the amounts used in foods. It has been proposed that aniseed and preparations used at recommended dosages may be used during pregnancy and lactation.\(^{(G52)}\)
Pharmaceutical Comment

The chemistry of aniseed is well studied and documented pharmacological activities support some of the herbal uses. Aniseed is used extensively as a spice and is widely used in conventional pharmaceuticals for its carminative, expectorant and flavouring properties. Aniseed contains anethole and estragole which are structurally related to safrole, a known hepatotoxin and carcinogen. Although both anethole and estragole have been shown to cause hepatotoxicity in rodents, aniseed is not thought to represent a risk to human health when it is consumed in amounts normally encountered in foods. Anethole was reaffirmed as GRAS in 1997 on the basis of the recognised metabolic detoxication of \textit{trans}-anethole in humans at low levels of exposure (1 mg/kg body weight).\textsuperscript{(2)} For medicinal use, it is recommended that treatment should not be continued for extended periods.
References


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Apricot
Species (Family)

Prunus armeniaca L. (Rosaceae)
Synonym(s)
None
Part(s) Used

Kernel (seed), expressed oil
Pharmacopoeial and Other Monographs

Martindale 33rd edition (G67)
Apricot is not included in the GSL. Amygdalin (a cyanogenetic glycoside) is classified as a POM.\(^{(1)}\)
**Constituents**

**Acids**
Phenolic. Various quinic acid esters of caffeic, \( p \)-coumaric and ferulic acids.\(^{(2)}\)
Neochlorogenic acid major in kernel, chlorogenic in fruit.\(^{(3)}\)

**Glycosides**
Cyanogenetic. Amygdalin (mandelonitrile diglucoside). Cyanide content of kernel varies from 2 to 200 mg/100 g.\(^{(3)}\)

**Tannins**
Catechins, proanthocyanidins (condensed).\(^{(4)}\)

**Other constituents**
Cholesterol, an oestrogenic fraction (0.09%) containing estrone (both free and conjugated) and \( \alpha \)-estradiol.\(^{(5)}\)

**Other plant parts**
Leaves and fruit contain various flavonol (kaempferol, quercetin) glycosides including rutin (major).\(^{(5)}\)
Food Use

Apricot fruit is commonly eaten. Apricot is listed by the Council of Europe as a natural source of food flavouring (category N1 and N2). These categories limit the total amount of hydrocyanic acid permitted in the final product to 1 mg/kg. Exceptions to this are 25 mg/kg for confectionery, 50 mg/kg for marzipan and 5 mg/kg for fruit juices. In the USA, apricot kernel extract is listed as GRAS (Generally Recognised As Safe).
Herbal Use

Traditionally, the oil has been incorporated into cosmetic and perfumery products such as soaps and creams.\(^{(G^{34})}\)
Dosage

None documented. Traditionally, apricot kernels have not been utilised as a herbal remedy.
Pharmacological Actions

During the late 1970s and early 1980s considerable interest was generated in apricot from claims that laetrile (a semi–synthetic derivative of amygdalin) was an effective treatment for cancer. Two review papers(6,7) discuss these claims for laetrile together with its chemistry, metabolism and potential toxicity.

The claims for laetrile were based on three different theories. The first claimed that cancerous cells contained abundant quantities of β-glucosidases, enzymes which release hydrogen cyanide from the laetrile molecule as a result of hydrolysis. Normal cells were said to be protected because they contained low concentrations of β-glucosidases and high concentrations of rhodanese, an enzyme which converts cyanide to the less toxic thiocyanate. However, this theory was disproved when it was shown that both cancerous and normal cells contain only trace amounts of β-glucosidases, and similar amounts of rhodanese. In addition, it was thought that amygdalin was not absorbed intact from the gastrointestinal tract.(6,7)

The second theory proposed that following ingestion, amygdalin was hydrolysed to mandelonitrile, transported intact to the liver and converted to a β-glucuronide complex. This complex was then carried to the cancerous cells, hydrolysed by β-glucuronidases to release mandelonitrile and subsequently hydrogen cyanide. This theory was considered to be untenable.(7)

A third theory proposed that laetrile is vitamin B$_{17}$, that cancer is a result of a deficiency of this vitamin, and that chronic administration of laetrile would prevent cancer. Again this was not substantiated by any scientific evidence.(7)

A retrospective analysis of the use of laetrile by cancer patients reported that it may have slight activity.(6,7) However, a subsequent clinical trial concluded that laetrile was ineffective in cancer treatment. Furthermore, it was claimed that patients taking laetrile reduced their life expectancy as a result of lack of proper medical care and chronic cyanide poisoning.(6,7)

In order to reduce potential risks to the general public, amygdalin was made a prescription–only medicine in 1984.(1)
Side-effects, Toxicity

Laetrile and apricot kernel ingestion are the most common sources of cyanide poisoning, with more than 20 deaths reported.\(^6,7\) Apricot kernels are toxic because of their amygdalin content. Hydrolysis of the amygdalin molecule by \(\beta\)-glucosidases, heat, mineral acids or high doses of ascorbic acid (vitamin C) yields hydrogen cyanide (HCN), benzaldehyde, and glucose. \(\beta\)-Glucosidases are not generally abundant in the gastrointestinal tract, but they are present in the kernels themselves as well as certain foods including beansprouts, carrots, celery, green peppers, lettuce, mushrooms and sweet almonds. Hydrolysis of the amygdalin molecule is slow in an acid environment but much more rapid in an alkaline pH. There may therefore be a delay in the onset of symptoms of HCN poisoning as a result of the transit time from the acid pH of the stomach to the alkaline environment of the small intestine.

**Acute poisoning**

Cyanide is rapidly absorbed from the upper gastrointestinal tract, diffuses readily throughout the body and promptly causes respiratory failure if untreated. Symptoms of cyanide toxicity progress rapidly from dizziness, headache, nausea, vomiting and drowsiness to dyspnoea, palpitations, marked hypotension, convulsions, paralysis, coma and death, which may occur from 1 to 15 minutes after ingestion. Antidotes for cyanide poisoning include nitrite, thiosulfate, hydroxocobalamin, cobalt edetate and aminophenol.\(^6,7\)

**Chronic poisoning**

Principal symptoms include increased blood thiocyanate, goitre, thyroid cancer, lesions of the optic nerve, blindness, ataxia, hypertonia, cretinism and mental retardation.\(^6\) These symptoms may develop as a result of ingesting significant amounts of cyanide, cyanogenetic precursors in the diet, or cyanogenetic drugs such as laetrile. Demyelinating lesions and other neuromyopathies reportedly occur secondary to chronic cyanide exposure, including long-term therapy with laetrile. Agranulocytosis has also been attributed to long-term laetrile therapy.\(^6,7\)

Individual reports of adverse reactions and cyanide poisoning in patients using laetrile have been documented.\(^G45\)

Normally, low concentrations of ingested cyanide are controlled naturally by exhalation or by rapid conversion to the less toxic thiocyanate by the enzyme rhodanese. Oral doses of 50 mg of hydrogen cyanide (HCN) can be fatal. This is equivalent to approximately 30 g kernels which represents about 50–60
kernels, and approximately 2 mg HCN/g kernel. Apricot seed has also been reported to contain 2.92 mg HCN/g.\(^8\) A 500–mg laetrile tablet was found to contain between 5 and 51 mg HCN/g.

There may be considerable variation in the number of kernels required to be toxic, depending on the concentration of amygdalin and β-glucosidases present in the kernels, the timespan of ingestion, the degree of maceration of the kernels, individual variation in hydrolysing, and detoxifying abilities.

Systemic concentrations of β-glucosidases are low and therefore toxicity following parenteral absorption of amygdalin is low. However, cyanide poisoning has been reported in rats following intraperitoneal administration of laetrile, suggesting another mechanism of hydrolysis had occurred.\(^6,7\)

It is thought that cyanogenetic glycosides may possess carcinogenic properties. Mandelonitrile (amygdalin = mandelonitrile diglucoside) is mutagenic and stimulates guanylate cyclase.\(^6,7\)
Contra-indications, Warnings

Apricot kernels are toxic due to their amygdalin content. Following ingestion hydrogen cyanide is released and may result in cyanide poisoning. Fatalities have been reported following the ingestion of apricot kernels. Contact dermatitis has been reported for apricot kernels.(9)

Pregnancy and lactation

Apricot kernels are toxic and should not be ingested. The ingestion of cyanogenetic substances may result in teratogenic effects.(6) However, one case has been reported where no acute toxicity was noted in the infant when laetrile was used during the third term of pregnancy. It was unknown whether chronic effects would be manifested at a later date.(6) Breeding rats fed ground apricot kernels had pups with normal birth weights, but with lower survival rates and lower weaning weights.(3)
Pharmaceutical Comment

Interest in apricot kernels was generated as a result of claims in the late 1970s that laetrile, a semi-synthetic derivative of the naturally occurring constituent amygdalin, was a natural, non-toxic cure for cancer. Apricot kernels were seen as an alternative source for this miracle cure. These claims have since been disproved and it has been established that laetrile (amygdalin) is far from non-toxic, particularly if administered orally. Fatal cases of cyanide poisoning have been reported following the ingestion of apricot kernels.
References

See also General References G10 G16 G32 G34 G43 G57.

Species (Family)

_Arnica montana_ L. (Asteraceae/Compositae)

_Arnica chamissonis_ Less. _ssp. foliosa_ (Nutt.) Maguire also allowed in German Pharmacopoeia.\(^{G52}\)
Synonym(s)

Leopard’s Bane, Mountain Tobacco, Wolf’s Bane
Part(s) Used

Flower
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)

BP 2002\(^{(G71)}\)

ESCOP 1997\(^{(G52)}\)

Martindale 33rd edition\(^{(G67)}\)

Mills and Bone\(^{(G50)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)

Ph Eur 2004\(^{(G72)}\)
Legal Category (Licensed Products)

GSL, for external use only. (G37)
Constituents
See References 1–3, and General References G2 G22 G41 G52 G64.

**Alkaloids**
Traces of non–toxic alkaloids tussilagine and isotussilagine\(^4\) but these are reportedly artefacts produced during extraction.\(^5\)

**Amines**
Betaine, choline and trimethylamine.

**Carbohydrates**
Mucilage, polysaccharides including inulin.

**Coumarins**
Scopoletin and umbelliferone.

**Flavonoids**
Betuletol, eupafolin, flavonol glucuronides\(^{1–3}\) hispidulin, isorhamnetin, kaempferol, laciniatin, luteolin, patuletin, quercetin, spinacetin, tricin and 3,5,7–tri hydroxy–6,3′,4′-trimethoxyflavone.

**Terpenoids**
Sesquiterpene lactones of the pseudoguaianolide–type, 0.2–0.8%.\(^{G52}\) Pharmacopoeial standard not less than 0.4%.\(^{G15 G28}\) Helenalin,\(^6\) 11α,13–dihydrohelenalin and their esters with acetic, isobutyric, methacrylic, tiglic and other carboxylic acids.\(^{G52}\) Diterpenes including \(\alpha\text–\text{labd}–13\text–\text{ene}–8α,15\text–\text{diol}.\(^7\)

**Volatile oils**
Up to 1%, normally about 0.3%. Thymol and thymol derivatives.

**Other constituents**
Amino acid (2–pyrrolidine acetic), bitter principle (arnicin), caffeic acid, carotenoids, fatty acids, phytosterols, polyacetyl enes, resin, tannin (unspecified).
Food Use

Arnica is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that arnica can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\(^{G16}\) In the USA, arnica is listed by the Food and Drugs Administration (FDA) as an ‘unsafe herb’,\(^{G22}\) and is only approved for food use in alcoholic beverages.\(^{G41}\)
Herbal Use

Arnica is stated to possess topical counter-irritant properties. It has been used for unbroken chilblains, alopecia neurotica, insect bites, gingivitis, aphthous ulcers, rheumatoid complaints and specifically for sprains and bruises.

German Commission E approved external use for injuries and consequences of accidents, e.g. haematoma, dislocation, contusions, oedema due to fracture, rheumatoid muscle and joint pains, inflammation of oral and throat region, furunculosis, inflammation caused by insect bites and superficial phlebitis.

Arnica is mainly used in homeopathic preparations; it is used to a lesser extent in herbal products.
Dosage

*Tincture of arnica flower*
(BPC 1949) 2–4 mL for external application only.

**Preparations**
Ointments, creams, gels, compresses made with 5–25% v/v tinctures, 5–25% v/v fluid extracts, diluted tinctures or fluid extract (1 : 3–1 : 10), decoctions 2.0 g drug/100 mL water. (G3 G4)
Pharmacological Actions

*In vitro* and animal studies

**Antimicrobial activity**
Arnica has been reported to exhibit bactericidal properties against *Listeria monocytogenes* and *Salmonella typhimurium*.\(^{(G41)}\) Helenalin and related sesquiterpenes from arnica have antimicrobial activity against *Bacillus subtilis* and *Staphylococcus aureus*,\(^{(8)}\) *Corynebacterium insidiosum*, *Micrococcus roseus*, *Mycobacterium phlei*, *Sarcinia lutea* and *Proteus vulgaris*.\(^{(G52)}\) Antifungal activity against *Trichophyton mentagrophytes*, *Epidermaphyton* spp. and *Botrytis cinerea* is reported for helenalin.\(^{(8,G52)}\)

**Antitumour activity**
The cytotoxicity of 21 flavonoids and five sesquiterpene lactones from *Arnica* spp. has been investigated *in vitro* in studies using GLC\textsubscript{4} (a human small cell lung carcinoma) and COLO 320 (a human colorectal cancer) cell lines.\(^{(9)}\) The most potent compound, helenalin, had an IC\textsubscript{50} value of 0.44 μmol/L against GLC\textsubscript{4} and 1.0 μmol/L against COLO 320 after 2 hours exposure.\(^{(9)}\) Some of the individual flavonols and flavones of arnica at non–toxic concentrations significantly reduced helenalin–induced cytotoxicity *in vitro*.\(^{(10)}\)

**Anti–inflammatory activity**
Moderate (29%) anti–inflammatory effect in the carageenan rat paw model has been reported for arnica.\(^{(11)}\) Helenalin is a potent inhibitor in this test and in chronic adjuvant arthritis tests in rats.\(^{(12)}\) The α-methylene-γ-lactone moiety of sesquiterpenes is required for activity, and the potency of helenalin is enhanced by the presence of the 6–hydroxy group.\(^{(13)}\) The mode of action of sesquiterpene lactones as anti–inflammatory agents is at multiple sites. At a concentration of 5 × 10\textsuperscript{−4} mol/L, the compounds uncoupled oxidative phosphorylation of human polymorphonuclear leukocytes, elevated cyclic adenosine monophosphate (cAMP) levels of rat neutrophils, and rat and mouse liver cells, and inhibited free and total lysosomal enzyme activity.\(^{(12)}\) Human polymorphonuclear neutrophil chemotaxis was inhibited at 5 × 10\textsuperscript{−4} mol/L, whereas prostaglandin synthetase activity was inhibited at concentrations of 10\textsuperscript{−3} mol/L. Helenalin and 11α-13–dihydrohelenalin inhibited collagen–induced platelet aggregation, thromboxane formation and 5–hydroxytryptamine secretion in a concentration–dependent manner.\(^{(14)}\)

**Other activities**
Helenalin has potent activity in the hotplate tail flick analgesic test in
Helenalin has also been reported to possess immunostimulant activity \textit{in vitro}, \(^{(15)}\) while high molecular weight polysaccharides have been found to exhibit immunostimulant activity \textit{in vivo} in the carbon clearance test in mice. \(^{(15,16)}\)

Arnica contains an adrenaline–like pressor substance and a cardiotonic substance. \(^{(G24)}\)

**Clinical studies**

A gel preparation of arnica flowers applied externally to the limbs of 12 male volunteers was more effective than placebo in the treatment of muscle ache. \(^{(G50\ G52)}\) In a randomised, double–blind, placebo–controlled study, 89 patients with venous insufficiency received arnica gel (20\% tincture) or placebo. \(^{(G50)}\) It was reported that arnica treatment produced improvements in venous tone, oedema and in feeling of heaviness in the legs.
Arnica is poisonous if taken internally. It is irritant to mucous membranes and ingestion may result in fatal gastroenteritis, muscle paralysis (voluntary and cardiac), increase or decrease in pulse rate, palpitation of the heart, shortness of breath, and may even lead to death. Helenalin is stated to be the toxic principle responsible for these effects. Thirty millilitres of a 20% arnica tincture, taken by mouth, was reported to produce serious, but not fatal, symptoms. The topical application of arnica has been documented to cause dermatitis. Arnica is a strong sensitisers, with the sesquiterpene lactone constituents implicated as the contact allergens: they possess an α-methylene group exocyclic to a γ-lactone ring, which is recognised as an immunological prerequisite for contact allergy. Helenalin is also reported to possess cytotoxic activity and this has been attributed to its ability to alkylate with sulfhydryl groups. Helenalin was not mutagenic in the Salmonella typhimurium assay.
Contra–indications, Warnings

Arnica should not be taken internally except in suitable homeopathic dilutions.\(^{(G42)}\)

Externally, arnica is poorly tolerated by some people, precipitating allergic reactions in sensitive individuals.\(^{(G42)}\) It should only be applied to unbroken skin and withdrawn at the first sign of reaction.\(^{(G7)}\) Toxic allergic skin reactions have occurred following application of the tincture.\(^{(G33)}\)
The chemistry and pharmacology of arnica are well documented, but there is a paucity of clinical data. Anti-inflammatory properties associated with sesquiterpene lactones justify the herbal uses, although allergenic and cytotoxic properties are also associated with this class of constituents. Arnica is not suitable for internal use, although it is present in some homeopathic products. External use of arnica tincture, which is included as an ingredient in some cosmetics, hair shampoos and bath preparations, may cause an allergic reaction. The pyrrolizidine alkaloids tussilagine and isotussilagine, reportedly present in arnica, are non-toxic. Moreover, they are artefacts produced during the extraction process with methanol.
References


Artichoke
Species (Family)

*Cynara scolymus* L. (Asteraceae/Compositae)
Synonym(s)

Globe Artichoke.

Globe artichoke should not be confused with Jerusalem artichoke, which is the tuber of *Helianthus tuberosa* L.
Part(s) Used

Leaf
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)

Complete German Commission E\(^{(G3)}\)

Martindale 33rd edition\(^{(G67)}\)

Mills and Bone\(^{(G50)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL$^{(G37)}$
Constituents

See References 1 and 2, and General Reference G41.

**Acids**
Phenolic, up to 2%. Caffeic acid, mono- and dicaffeoylquinic acid derivatives, e.g. cynarin (1,5-di-O-caffeoylquinic acids) and chlorogenic acid (mono derivative).

**Flavonoids**
0.1–1%. Flavone glycosides e.g. luteolin–7β-rutinoside (scolymoside), luteolin–7β-D-glucoside and luteolin–4β-D-glucoside.

**Volatile oils**
Sesquiterpenes β-selinene and caryophyllene (major); also eugenol, phenylacetaldehyde, decanal, oct–1–en–3–one, hex–1–en–3–one, and non-trans-2–enal.

**Other constituents**
Phytosterols (taraxasterol and β-taraxasterol), tannins, glycolic and glyceric acids, sugars, inulin, enzymes including peroxidases,\(^3\) cynaropicrin and other sesquiterpene lactones, e.g. grosheimin, cynarotriol.\(^4,5\) The root and fully developed fruits and flowers are devoid of cynaropicrin; highest content reported in young leaves.\(^6\)
Food Use

Artichoke is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that artichoke can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\(^{(G16)}\) In the USA, artichoke leaves are approved for use in alcoholic beverages only, with an average maximum concentration of 0.0016% (16 ppm).\(^{(G41)}\)
Herbal Use

Artichoke is stated to possess diuretic, choleretic, hypocholesterolaemic, hypolipidaemic, and hepatostimulating properties. (7-9) Modern use of artichoke is focused on its use in the treatment of hyperlipidaemia, hyperlipoproteinaemia, non-ulcer dyspepsia and conditions requiring an increase in choleresis. There is also interest in the potential hepatoprotective properties of globe artichoke, although this has not yet been tested in controlled clinical trials. (10,11)
Dosage

The German Commission E recommends an average daily dose of 6 g drug, or an equivalent dose of extract (based on the herb-to-extract ratio) or other preparations, for dyspeptic problems.\(^{\text{G3 G56}}\) A recommended dosage regimen for liquid extract (1 : 2) is 3–8 mL daily.\(^{\text{G50}}\)

Dosages used in clinical trials of globe artichoke leaf extract have assessed the effects of dosages of up to 1.92 g daily in divided doses for up to six months.\(^{\text{12}}\)
Pharmacological Actions

Several pharmacological properties have been documented for artichoke leaf, including inhibition of cholesterol biosynthesis, hypolipidaemic, anti-oxidant and hepatoprotective activity. It remains unclear which of the constituents of artichoke are responsible for its pharmacological activities. The dicafeoylquinic acids, which include cynarin, are likely to be an important group of constituents in this respect.\(^{11, G50}\) The sesquiterpene lactones, such as cynaropicrin, and flavonoids, such as luteolin glycoside, may also exert biological effects.\(^{11}\)

**In vitro and animal studies**

**Hypolipidaemic, hypocholesterolaemic and choleretic activity**

Hypolipidaemic, hypocholesterolaemic and choleretic activities are well documented for globe artichoke leaf extract and particularly for the constituent cynarin; this literature has been reviewed.\(^{10, 11}\) Globe artichoke leaf extract not only increases choleresis and, therefore, cholesterol elimination, but also has been shown to inhibit cholesterol biosynthesis.\(^{10}\) Preparations of globe artichoke leaf extract inhibit cholesterol biosynthesis in a concentration–dependent manner in studies in cultured rat hepatocytes.\(^{13, 14}\) Low concentrations (<0.1 mg/mL) of globe artichoke extract achieved around 20% inhibition, whereas 65% inhibition was noted with concentrations of 1 mg/mL. Luteolin was considered to be one of the most important constituents for this effect, and it was suggested that a possible mechanism of action might be indirect inhibition of hydroxymethylglutaryl-CoA reductase (HMG-CoA).\(^{14}\) Other in vitro studies have documented a concentration–dependent inhibition of de novo cholesterol biosynthesis in cultured rat and human hepatocytes for globe artichoke leaf extract 0.03–0.1 mg/mL.\(^{15}\)

Several other experimental studies have documented lipid–lowering effects for globe artichoke leaf extract and cynarin in vivo.\(^{10, 11}\) A study in rats explored the hypocholesterolaemic, hypolipidaemic and choleretic effects of purified (containing 46% caffeoylquinic acids, calculated as chlorogenic acid) and total extracts of globe artichoke leaf (containing 19% caffeoylquinic acids, calculated as chlorogenic acid).\(^{7}\) The purified extract was found to be more potent than the total artichoke extract: purified extract 25 mg/kg intraperitoneally reduced plasma triglyceride and cholesterol concentrations by 33% and 45%, respectively, whereas reductions of only 18% and 14%, respectively, were observed with the total extract (100 mg/kg intraperitoneally).\(^{7}\) Both purified (25 mg/kg intraperitoneally) and total extract (200 mg/kg intraperitoneally) significantly enhanced bile secretion.
following treatment, compared with baseline values; the increase in bile secretion seen with the purified extract was still statistically significant 3 hours after treatment. The more potent pharmacological activities observed with the purified extract were attributed to the higher concentration of monoaetoxyquinic acids (e.g. chlorogenic, neochlorogenic) compared with dicaetoxyquinic acids (e.g. cynarin) present.

Another study investigated the effects of cynarin on total cholesterol concentrations in serum and liver of rats given ethanol (6 g/kg/day by gavage over three days). In rats given ethanol alone, serum cholesterol concentrations rose significantly by 44%, compared with controls (p < 0.01). Rats given ethanol plus cynarin (30 mg/kg intraperitoneally 30 minutes before gavage) showed a significant reduction in serum cholesterol concentrations, compared with controls (p < 0.05).

**Antioxidant and hepatoprotective activity**

*In vitro*, a luteolin-rich globe artichoke leaf aqueous extract (flavonoid content around 0.4% w/w) retarded low-density lipoprotein (LDL) oxidation in a concentration-dependent manner (determined by a prolongation of the lag phase to conjugated diene formation). The same tests carried out with the pure aglycone luteolin at concentrations of 0.1–1 μmol/L showed that this constituent had a similar concentration-dependent effect on LDL oxidation in this model. Luteolin-7-O-glucoside also demonstrated a concentration-dependent reduction in LDL oxidation, but was less potent than luteolin.

Several *in vitro* and *in vivo* studies have investigated the antioxidative and hepatoprotective properties of globe artichoke leaf extracts, and their constituents, against liver cell damage induced by different hepatotoxins.

The hepatoprotective effect of polyphenolic compounds isolated from artichoke has been investigated *in vitro* using rat hepatocytes. Cynarin was the only compound reported to exhibit significant cytoprotective activity, with a lesser action demonstrated by caffeic acid. A standardised extract of globe artichoke (Hepar-SL forte) significantly inhibited the formation of malondialdehyde induced by tert-butylhydroperoxide (t-BHP) in a concentration-dependent manner within 40 minutes of incubation, compared with control. The protective antioxidant effect of globe artichoke was reported to be significant, compared with control, even at a concentration of 1 μg/mL. A reduction in t-BHP-induced cell death with globe artichoke extract was also observed. Further studies assessed the antioxidative and protective potential of the same extract (Hepar-SL forte) in cultures of primary rat hepatocytes exposed to t-BHP. Incubation of cultured hepatocytes with globe artichoke extract and t-BHP inhibited t-BHP-induced malondialdehyde
formation in a concentration–dependent manner. Globe artichoke extract was significantly effective, compared with control, at concentrations down to 0.001 mg/mL. Furthermore, concentrations of globe artichoke extract down to 0.005 mg/mL significantly enhanced hepatocyte survival. The antioxidative effect of the extract was not affected by various pretreatments (including tryptic digestion, boiling and acidification), although it was sensitive to alkalinisation. Incubation with the globe artichoke constituents chlorogenic acid and cynarin resulted in significant inhibition, and incubation with both compounds was reported to have a synergistic effect, although an additive effect may be a more accurate description of the findings. Chlorogenic acid and cynarin were not solely responsible for the antioxidant effect, as reduction of malondialdehyde formation by the extract was at least twofold that seen with the chlorogenic acid and cynarin.\(^{(19)}\) The antioxidative and hepatoprotective potential of globe artichoke extract was confirmed in other studies which also indicated that several constituents of the extract may contribute to the effects.\(^{(20)}\)

The effects of globe artichoke leaf extract and its constituents have also been investigated for activity against oxidative stress in studies using human leukocytes.\(^{(21)}\) Globe artichoke leaf extract demonstrated a concentration–dependent inhibition of oxidative stress induced by several agents, such as hydrogen peroxide and phorbol–12–myristate–13–acetate, that generate reactive oxygen species. The constituents cynarin, caffeic acid, chlorogenic acid and luteolin also showed concentration–dependent inhibitory activity in these models.

*In vivo* hepatoprotectivity against tetrachloromethane–induced hepatitis has been documented for globe artichoke leaf extract (500 mg/kg) administered orally to rats 48 hours, 24 hours and 1 hour before intoxication.\(^{(9)}\) Concentrations of liver transaminases were significantly lower in rats given globe artichoke leaf extract, compared with those in controls. A hepatoregenerating effect has also been described for an aqueous extract of globe artichoke leaf administered orally to rats for three weeks following partial hepatectomy.\(^{(22)}\) Regeneration was determined by stimulation of mitosis and increased weight in the residual liver when animals were sacrificed in globe artichoke-treated rats, compared with controls. In a similar study, aqueous extract of globe artichoke leaf (0.5 mL daily for five days preceding hepatectomy) was found to be more potent than a root extract.\(^{(23)}\)

**Clinical studies**

Several clinical trials have explored the choleretic and hypolipidaemic properties of globe artichoke leaf extract, and its effects in patients with
A randomised, double-blind, placebo-controlled, crossover trial involving 20 male volunteers assessed the choleretic effects of a single intraduodenal dose (1.92 g in 300 mL water) of the globe artichoke leaf extract Hepar-SL forte.\(^{(24)}\) Intraduodenal bile secretion, the primary outcome variable, was measured using multichannel probes starting 30 minutes after drug administration and continuing for up to 4 hours afterwards. An increase in bile secretion was observed in both groups; maximal increases for globe artichoke leaf extract and placebo were 152% at 60 minutes after drug administration, and 40% at 30 minutes, respectively. Differences between globe artichoke leaf extract and placebo were statistically significant at 30, 60 and 90 minutes after drug administration \((p < 0.01)\) and at 120 and 150 minutes after drug administration \((p < 0.05)\). In another randomised controlled trial, 60 patients with dyspepsia received a combination preparation containing extracts of globe artichoke 50 mg, boldo \((\textit{Peumus boldus})\) 30 mg and chelidonium \((\textit{Chelidonium majus})\) 20 mg per tablet, or placebo, three times daily for 14 days.\(^{(25)}\) The volume of bile secreted, measured using a duodenal probe, increased significantly in the treatment group, compared with the placebo group \((p < 0.01)\). Also, an improvement in symptoms was reported for 50% of the treatment group, compared with 38% of the placebo group. There are also clinical studies in the older literature (some of which were placebo-controlled trials, whereas others were open, uncontrolled studies) which report choleretic effects with globe artichoke leaf extract. These trials have been summarised in several reviews.\(^{10,24,G50}\)

The effects of globe artichoke leaf extract have also been monitored in several postmarketing surveillance (phase IV) studies in patients with non-specific gastrointestinal complaints, including dyspepsia,\(^{(12)}\) functional biliary tract complaints, constipation and gastric irritation.\(^{(11)}\) The studies monitored the effects of globe artichoke leaf extract (Hepar-SL forte; one capsule contains 320 mg standardised aqueous extract; drug-extract ratio: 3.5 to 5.5 : 1) up to six capsules daily for six weeks\(^{(11)}\) or six months.\(^{(12)}\) Both studies reported improvements in clinical symptoms and reductions in serum total cholesterol and triglyceride concentrations, compared with baseline values. A subgroup analysis of 279 patients with at least three of five symptoms of irritable bowel syndrome reported significant reductions in the severity of symptoms and favourable evaluations of overall effectiveness by both physicians and participants.\(^{(26)}\) The findings from postmarketing surveillance studies provide supporting data for the effects of globe artichoke leaf extract, but these are open studies and do not include a control group and, therefore, are not designed to assess efficacy.
The efficacy of globe artichoke leaf extract in patients with hyperlipoproteinaemia has been assessed in a randomised, double-blind, placebo-controlled, multicentre trial involving 143 patients with initial total cholesterol concentrations of >7.3 mmol/L (>280 mg/dL). Participants received globe artichoke leaf extract (CY450; drug-extract ratio: 25 to 35 : 1) 1800 mg daily in two divided doses, or placebo, for six weeks. At the end of the study, mean total cholesterol concentrations had decreased by 18.5% to 6.31 mmol/L and by 8.6% to 7.03 mmol/L in the CY450 and placebo groups, respectively (p < 0.0001). CY450 treatment also led to a significant reduction in LDL cholesterol, compared with placebo (p = 0.0001). There was no difference between CY450 recipients and placebo recipients in blood concentrations of the liver enzyme gamma-glutamyltransferase (GGT).

A published abstract reports the findings of a previous randomised, double-blind, placebo-controlled trial of a globe artichoke leaf extract (Hepar-SL forte; 640 mg three times daily for 12 weeks) involving 44 healthy volunteers. Mean baseline total cholesterol concentrations for participants in this study were low (placebo group: 203.0 mg/dL; globe artichoke extract group: 204.2 mg/dL). Subgroup analysis suggested lipid-lowering effects with globe artichoke extract for participants with baseline total cholesterol concentrations of >200 mg/dL. However, numbers of participants included in this analysis were small. The study indicates only that trials in patients with hyperlipoproteinaemia are required.

A series of three open, uncontrolled studies involved the administration of pressed globe artichoke juice (obtained from fresh leaves and flower buds) 10 mL three times daily for up to 12 weeks to a total of 84 patients with secondary hyperlipidaemia (total cholesterol ≥260 mg/dL). After six weeks’ treatment, total cholesterol, LDL cholesterol and triglyceride concentrations decreased, whereas high-density lipoprotein cholesterol tended to increase. Another uncontrolled study involved the administration of cynarin to 17 patients with familial type IIa or type IIb hyperlipoproteinaemia for whom blood lipid concentrations were maintained with dietary treatment alone. Cynarin was taken 15 minutes before meals at either 250 mg or 750 mg daily dose. Over a period of three months, cynarin was reported to have no effect on mean serum cholesterol and triglyceride concentrations. The results were in agreement with the findings of some previous workers, but also in contrast to other studies that have reported cynarin to be effective in lowering serum concentrations of cholesterol and triglycerides when taken in daily doses ranging from 60 mg to 1500 mg.
Side-effects, Toxicity

A randomised, double-blind, placebo-controlled trial involving 143 patients with hyperlipoproteinaemia reported a similar frequency of adverse events for globe artichoke leaf extract (CY450) and placebo groups.\((27)\) A total of 28 adverse events were reported during the study, 26 of which related to mild changes in laboratory values. The relationship to the globe artichoke was considered to be ‘unlikely’ in all cases.

Postmarketing surveillance (phase IV) studies have monitored patients with non-specific gastrointestinal complaints receiving treatment with globe artichoke leaf extract (Hepar-SL forte; up to 1.92 g daily for six weeks\((11)\) or six months\((12)\)). In one study involving 533 patients with non-specific gastrointestinal complaints, including dyspepsia, functional biliary tract complaints, constipation and gastric irritation, seven adverse events (weakness, hunger, flatulence) were reported (1.3% of participants).\((11)\) No serious adverse events were reported. A second postmarketing surveillance study involved 203 patients with symptoms of dyspepsia who received globe artichoke leaf extract up to 1.92 g daily for up to six months.\((12)\) It was reported that no adverse events were recorded during the study, and that the physician’s overall judgement of tolerability was given as ‘good’ or ‘excellent’ in 98.5% of cases.

Allergic contact dermatitis, with cross-sensitivity to other Compositae plants, has been documented for globe artichoke.\((30,G51)\) A case of occupational contact urticaria syndrome in a 20-year-old woman has been reported in association with globe artichoke. The woman developed acute generalised urticaria, angioedema of the hands, forearms and face, and respiratory symptoms after handling globe artichokes. The clinical history and results of skin-prick tests indicated that the woman had developed type I allergy to globe artichoke antigen(s).\((31)\) An isolated case of allergy to ingested globe artichoke has also been described.\((32)\)

Cynaropicrin and other sesquiterpene lactones with allergenic potential have been isolated from globe artichoke.\((30,G53)\) Purified globe artichoke extract is more toxic than a total extract. LD\(_{50}\) values (rat, by intraperitoneal injection) have been documented as greater than 1000 mg/kg (total extract) and 265 mg/kg (purified extract).\((7)\)
Contra-indications, Warnings

Globe artichoke yields cynaropicrin, a potentially allergenic sesquiterpene lactone.\textsuperscript{(G51)} Individuals with an existing hypersensitivity to any member of the Compositae family may develop an allergic reaction to globe artichoke.

**Pregnancy and lactation**

In view of the lack of toxicity data, excessive use of globe artichoke should be avoided during pregnancy and lactation.
Pharmaceutical Comment

Globe artichoke is characterised by the phenolic acid constituents, in particular cynarin. Experimental studies (in vitro and in vivo) support some of the reputed uses of artichoke. Traditionally, the choleretic and cholesterol-lowering activities of globe artichoke have been attributed to cynarin. However, studies in animals and humans have suggested that these effects may in fact be due to the monocaffeoylquinic acids present in globe artichoke (e.g. chlorogenic and neochlorogenic acids). Clinical trials investigating the use of globe artichoke and cynarin in the treatment of hyperlipidaemia generally report positive results. However, further rigorous clinical trials are required to establish the benefit of globe artichoke leaf extract as a lipid- and cholesterol-lowering agent. Hepatoprotective and hepatoregenerating activities have been documented for cynarin in vitro and in animals (rats). However, these effects have not yet been documented in clinical studies.
References

See also General References G3 G9 G16 G19 G36 G41 G43 G49 G50 G51.

16. Wójcicki J. Effect of 1,5–dicaffeylquinic acid (Cynarine) on cholesterol levels in serum and liver of acute ethanol–treated rats. Drug Alcohol...
Depend 1978; 3: 143–145. ([PubMed])


Asafoetida
Species (Family)

*Ferula* species including:

i. *Ferula assafoetida L. (Ferula rubricaulis*) Boiss

ii. *Ferula foetida* (Bunge) Regel (Apiaceae/Umbelliferae)
Synonym(s)
Asafetida, Asant, Devil’s Dung, Gum Asafetida
Part(s) Used

Oleo gum resin obtained by incising the living rhizomes and roots.
Pharmacopoeial and Other Monographs

BHC 1992\(^{G6}\)

BHP 1996\(^{G9}\)

Martindale 33rd edition\(^{G67}\)

PDR for Herbal Medicines 2nd edition\(^{G36}\)
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents


**Gum fraction**
25%. Glucose, galactose, L-arabinose, rhamnose and glucuronic acid.

**Resins**
40–64%. Ferulic acid esters (60%), free ferulic acid (1.3%), asaresinotannols and farnesiferols A, B and C, coumarin derivatives (e.g. umbelliferone), coumarin–sesquiterpene complexes (e.g. asacoumarin A and asacoumarin B).\(^{(1)}\) Free ferulic acid is converted to coumarin during dry distillation.

**Volatile oils**
3–17%. Sulfur–containing compounds with disulfides as major components, various monoterpenes.\(^{(1)}\)

The oleo gum resins of different *Ferula* species are not identical and many papers have documented their phytochemistry,\(^{(2–11)}\) reporting polysulfanes,\(^{(2–11)}\) complex acetylenes,\(^{(3)}\) phenylpropanoids\(^{(7)}\) and many sesquiterpene derivatives.\(^{(2,4,5,6,8,9)}\)

C-3 prenylated 4–hydroxycoumarin derivatives (e.g. ferulenol) are thought to represent the toxic principles in the species *Ferula communis*.\(^{(12)}\)
Food Use

Asafoetida is used widely in foods. Asafoetida (essential oil, fluid extract and gommo-oleoresin) is listed by the Council of Europe as a source of natural food flavouring (category 5) (see Appendix 23).\(^{(G17)}\) Asafoetida is approved for food use in the USA.\(^{(G41)}\)
Herbal Use

Asafoetida is stated to possess carminative, antispasmodic and expectorant properties. It has been used for chronic bronchitis, pertussis, laryngismus stridulus, hysteria and specifically for intestinal flatulent colic.\(^{(G6\ G7)}\)
Dosage

*Powdered resin*
0.3–1 g three times daily\(^{G6 \ G7}\)

*Tincture of asafoetida*
(BPC 1949) 2–4 mL
Pharmacological Actions

**In vitro and animal studies**

Asafoetida has been reported to possess anticoagulant and hypotensive properties.\(^{(G41)}\) Asafoetida is an ingredient of a plant mixture reported to have antidiabetic properties in rats.\(^{(13)}\) However, when the individual components of the mixture were studied asafoetida was devoid of antidiabetic effect with myrrh and aloe gum extracts representing the active hypoglycaemic principles.\(^{(14)}\)

Oestrogenic activity in rats has been documented for carotane sesquiterpenes and ferujol (a coumarin) isolated from *Ferula jaeschkeana*.\(^{(15,16)}\)

**Clinical studies**

A protective action against fat–induced hyperlipidaemia has been documented for asafoetida and attributed to the sulfur compounds in the volatile oil fraction of the resin.\(^{(17)}\) Two double–blind studies have reported the efficacy of asafoetida in the treatment of irritable bowel syndrome to be just below the 5% significance level in one study\(^{(18)}\) and at 1% in the other.\(^{(19)}\)
Side–effects, Toxicity

Asafoetida is documented to be relatively non–toxic; ingestion of 15 g produced no untoward effects.\(^{(G^{45})}\) A report of methaemoglobinaemia has been associated with the administration of asafoetida (in milk) to a five–week–old infant for the treatment of colic.\(^{(20)}\) Asafoetida was found to exert an oxidising effect on fetal haemoglobin but not on adult haemoglobin.

Toxic coumarin constituents of a related species, *Ferula communis*, have been documented to reduce prothrombin concentrations and to cause haemorrhaging in livestock.\(^{(21,G^{51})}\)

Two other species, *Ferula galbaniflua* and *Ferula rubicaulis*, are stated to contain a gum that is rubefacient and irritant, causing contact dermatitis in sensitive individuals.\(^{(G^{51} G^{58})}\)

A weak sister chromatid exchange–inducing effect in mouse spermatogonia\(^{(22)}\) and clastogenicity in mouse spermatocytes\(^{(23)}\) has been documented for asafoetida. Chromosomal damage by asafoetida has been associated with the coumarin constituents.
Contra-indications, Warnings

Asafoetida should not be given to infants because of the oxidising effect on fetal haemoglobin resulting in methaemoglobinaemia.\(^{(20)}\) The gum of some \textit{Ferula} species is reported to be irritant and therefore may cause gastrointestinal irritation or induce contact dermatitis in some individuals. Excessive doses may interfere with anticoagulant therapy and with hypertensive and hypotensive therapy.

\textbf{Pregnancy and lactation}

Asafoetida has a folkloric reputation as an abortifacient and an emmenagogue.\(^{(G30)}\) However the use of asafoetida during pregnancy is probably acceptable, provided doses do not exceed amounts normally ingested in foods. In view of the toxic effect to infants (e.g. methaemoglobinaemia), asafoetida should be avoided during breast feeding.
Asafoetida is a complex oleo gum resin consisting of many constituents that vary according to the different species used. Asafoetida is commonly used in foods but little scientific evidence is available to justify the herbal uses. In view of the known pharmacologically active constituents, asafoetida should not be taken in amounts exceeding those used in foods.
References


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Species (Family)

*Geum urbanum* L. (Rosaceae)
Synonym(s)
Benedict’s Herb, Colewort, Geum, Herb Bennet
Part(s) Used

Herb
Pharmacopoeial and Other Monographs

BHP 1983\(^{(G7)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

Avens is not included in the GSL. (G37)
Constituents

See General References G40 G49 G64.

Limited information is available on the herb. Constituents reported include bitter principles, resin, tannins and volatile oil.

Other plant parts

The root has been more extensively studied and is reported to contain a phenolic glycoside (gein), yielding eugenol as the aglycone and vicianose (disaccharide) as the sugar component;\(^1\) 30% tannin, including gallic, caffeic and chlorogenic acids (pseudotannins generally associated with condensed tannins);\(^1\) a bitter substance, a flavonoid, and volatile oil.
Food Use

Avens is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that avens can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\textsuperscript{(G16)}
**Herbal Use**

Avens is stated to possess antidiarrhoeal, antihaemorrhagic, and febrifugal properties. It has been used for diarrhoea, catarrhal colitis, passive uterine haemorrhage, intermittent fevers, and specifically for ulcerative colitis.\(^{(G7 G64)}\)
Dosage

*Dried herb*
1–4 g or by infusion three times daily.\(^{(G7)}\)

*Liquid extract*
1–4 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

*In vitro* and animal studies

A 20% aqueous decoction of avens, administered by intravenous injection, has been reported to produce a reduction in blood pressure in cats.\(^2\) Tannins are generally known to possess astringent properties.
Side-effects, Toxicity

None documented.
Contra-indications, Warnings

In view of the reported tannin constituents and the lack of toxicity data, it is advisable to avoid excessive use of avens.

**Pregnancy and lactation**
Avens is reputed to affect the menstrual cycle. In view of the lack of phytochemical, pharmacological and toxicological data, the use of avens during pregnancy should be avoided.
Pharmaceutical Comment

Limited phytochemical or pharmacological data are available for avens, although reported tannin constituents would indicate an astringent action thus supporting the traditional use in diarrhoea and haemorrhage. In view of the lack of toxicity data, excessive use should be avoided.
References

See also General References G2 G7 G16 G30 G31 G36 G37 G40 G49 G64.


Species (Family)

*Myrica cerifera* L. (Myricaceae)
Synonym(s)
Candleberry Bark, Myrica, Wax Myrtle Bark
Part(s) Used
Root bark
Pharmacopoeial and Other Monographs

BHP 1996\textsuperscript{(G9)}

Martindale 33rd edition\textsuperscript{(G67)}

PDR for Herbal Medicines 2nd edition\textsuperscript{(G36)}
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents

See General References G22 G41 G48 G64.

Flavonoids
Myricitrin.

Tannins
3.9% (bark), 34.82% (total aqueous extract).

Terpenoids
Myricadiol, taraxerol and taraxerone.\(^{(1)}\)

Other constituents
Albumen, red dye, gum, resin, starch, wax containing palmitic, myristic and lauric acid esters.
Food Use

Bayberry is not used in foods.
Herbal Use

Bayberry is stated to possess antipyretic, circulatory stimulant, emetic and mild diaphoretic properties. It has been used for diarrhoea, colds and specifically for mucous colitis. An infusion has been used as a gargle for a sore throat, and as a douche for leucorrhoea. Powdered root bark has been applied topically for the management of indolent ulcers.\(^{G7 \ G41 \ G64}\)
Dosage

*Powdered bark*
0.6–2.0 g by infusion or decoction three times daily.\(^{(G7)}\)

*Liquid extract*
0.6–2.0 mL (1 : 1 in 45% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

*In vitro and animal studies*

Myricitrin has been reported to exhibit choleretic, bactericidal, paramecicidal, and spermatocidal activity; myricadiol has mineralocorticoid activity.\(^{(G41)}\) Tannins are known to possess astringent properties.
Side–effects, Toxicity

A total aqueous extract, tannin fraction, and tannin–free fraction from bayberry were all reported to produce tumours in NIH black rats, following weekly subcutaneous injections for up to 75 weeks.(2,3) The number of tumours that developed were stated to be statistically significant for the tannin fraction and tannin–free fraction. Analysis of the tannin–free fraction revealed the presence of four phenolic compounds, one of which was identified as myricitrin. No tumours were reported in a later study, in which rats were given subcutaneous injections of total aqueous extract for 78 weeks.

Large doses may cause typical mineralocorticoid side–effects (e.g. sodium and water retention, hypertension).
Contra–indications, Warnings

Large doses may interfere with existing hypertensive, hypotensive or steroid therapy. Excessive use of tannin–containing herbs is not recommended.

**Pregnancy and lactation**
The safety of bayberry has not been established. In view of the possible mineralocorticoid activity and the reported carcinogenic activity, the use of bayberry during pregnancy and lactation should be avoided.
Pharmaceutical Comment

Limited chemical information is available for bayberry. Documented tannin constituents justify some of the herbal uses. In addition, mineralocorticoid activity has been reported for one of the triterpene constituents. In view of this and the tannin constituents, excessive use of bayberry should be avoided.
References

See also General References G9 G11 G22 G31 G32 G36 G37 G41 G44 G48 G64.


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Bilberry
Species (Family)

Vaccinium myrtillus L. (Ericaceae)
Synonym(s)

Blueberry, bogberry, huckleberry, *Myrtilus niger* Gilib., *Vaccinium angulosum* Dulac, *Vaccinium montanum* Salisb., whortleberry
Part(s) Used

Fruit (berries), leaves
Pharmacopoeial and Other Monographs

Complete German Commission E\(^{(G3)}\)

Martindale 33rd edition (Myrtillus)\(^{(G67)}\)

Mills and Bone\(^{(G50)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)

Ph Eur 2004\(^{(G72)}\)
Legal Category (Licensed Products)

Bilberry is not included in the GSL.
Constituents
See General References G2 G55.

Berries

*Flavonoid glycosides*
Anthocyanins (particularly glycosides of delphinidin, cyanidin, petunidin, peonidin, malvidin),\(^1,2\) quercetin–3–glucuronide and hyperoside.\(^3\)

*Polyphenols*
Catechin, epicatechin and tannins.

*Other constituents*
Pectins\(^1\) and vitamin C.

Leaves

*Flavonoids*
Quercetin and its glycosides (hyperoside, quercitrin).\(^1\)

*Phenolic acids*
Caffeic, \(p\)-coumaric, \(p\)-hydroxybenzoic, protocatechuic and melilotic.\(^4\)

*Other constituents*
Tannins and iridoids.\(^1\)
Food Use

Bilberries are used in foods.\(^{(1)}\) Bilberry is listed by the Council of Europe as a natural source of food flavouring (category N1). This category indicates that there are no restrictions on the use of bilberry in foods.\(^{(G16)}\)
Herbal Use

Bilberry is stated to possess astringent, tonic and antiseptic properties and has traditionally been used in the treatment of diarrhoea, dysentery, haemorrhoids, gastrointestinal inflammations, mouth infections, scurvy and urinary complaints.\(^1\) It has also been used in diabetes, gout and rheumatism and applied locally in eye inflammation, burns and skin infections.\(^1\)
Dosage

*Dried fruit*
20–60 g daily as a decoction for the treatment of diarrhoea.\(^{(G2)}\)
Pharmacological Actions

Several pharmacological activities have been documented for bilberry, including ophthalmic activity and anti-inflammatory, wound-healing, anti-ulcer, anti-atherosclerotic and vasoprotective properties. The biochemical, biological, pharmacological and clinical effects of bilberry have been reviewed.\(^{(1)}\)

**In vitro and animal studies**

An anthocyanidin extract of *V. myrtillus* has been reported to act as a superoxide anion scavenger\(^{(1,5)}\) and as an inhibitor of lipid peroxidation in rat liver microsomes\(^{(1,5,6)}\) and in mouse liver tissue *in vivo*\(^{(5)}\) and to inhibit potassium ion loss induced by free radicals in human erythrocytes.\(^{(1)}\) *V. myrtillus* extract is stated to have a potent protective antioxidant action on human low-density lipoproteins (LDLs) *in vitro* during copper-mediated oxidation.\(^{(7)}\) Oxidative activity is recognised as a major process in tissue damage in a variety of pathological conditions, such as atherosclerosis and carcinogenesis. In addition, oxidative stress is thought to be involved in brain ageing and age-related neurodegenerative disease. A study in rats reported that, compared with rats fed a control diet, dietary supplementation of blueberry (bilberry) extract for eight weeks reversed age-related deficits in several neuronal and behavioural parameters, such as enhancement of dopamine release from striatal slices and a water maze performance test.\(^{(8)}\)

*V. myrtillus* anthocyanins have been reported to inhibit aggregation of human platelets *in vitro* in a dose-dependent manner\(^{(9)}\) and, in rats, *V. myrtillus* anthocyanins administered orally at doses ranging from 5 to 400 mg/kg have been shown to prolong bleeding time markedly.\(^{(10)}\) Inhibition of platelet aggregation has also been reported in humans treated with *V. myrtillus* anthocyanins (see Clinical studies).\(^{(11)}\)

*In vitro* inhibition of elastase, a proteolytic enzyme involved with elastic fibre and connective tissue degeneration and with some pathological vascular conditions, has been demonstrated in studies using anthocyanins extracted from *V. myrtillus*.\(^{(12)}\)

The hypolipidaemic activity of oral administration of extracts of *V. myrtillus* leaves has been demonstrated in rats.\(^{(13,14)}\) In genetically hyperlipidaemic rats, plasma triglyceride and cholesterol concentrations, but not free fatty acids, decreased significantly.\(^{(13)}\) In streptozotocin-induced diabetic rats, plasma glucose concentrations as well as plasma triglyceride concentrations decreased significantly compared with values in control rats.\(^{(14)}\) In further
experiments using blueberry and clofibrate, both preparations reduced plasma triglyceride concentrations in a dose–dependent manner in rats fed a hyperlipidaemic diet and in ethanol–treated normolipidaemic rats.\(^{(14)}\) Blueberry, however, did not prevent fructose–elicited increases in plasma triglyceride concentrations. Other studies in glucose–loaded mice failed to demonstrate hypoglycaemic activity following oral administration of blueberry leaf extract.\(^{(15)}\)

Several \textit{in vitro} studies have demonstrated the relaxing effects of \textit{V. myrtillus} anthocyanins on isolated vascular smooth muscle preparations, including the thoracic vein and splenic and coronary arteries.\(^{(16–18)}\) There is evidence that the mechanism for this smooth muscle relaxant effect is via stimulation of prostaglandin release within vessel walls.\(^{(19)}\)

Effects of \textit{V. myrtillus} anthocyanins on enhancing arterial vasomotion (rhythmic variation of arteriole diameter in the microvasular network which influences microvascular blood flow and the formation of interstitial fluid) have been shown in experimental models, including the cheek pouch microcirculation of hamsters.\(^{(20)}\) This model has also been used to investigate the effects of \textit{V. myrtillus} anthocyanins on ischaemia–reperfusion injury.\(^{(21)}\) Oral administration for two and four weeks of Myrtocyan, a commercially available product comprising bilberry anthocyanin complex, reduced the increase in capillary permeability, decreased leukocyte adhesion and improved capillary perfusion compared with controls. In rats, oral administration of \textit{V. myrtillus} anthocyanins for 12 days before the induction of hypertension (by ligation of the abdominal aorta) limited the increase in vascular permeability and maintained a normal blood–brain barrier permeability.\(^{(22)}\)

Components of bilberry have been reported to exhibit potential anticarcinogenic activity \textit{in vitro} as demonstrated by inhibition of the induction of ornithine decarboxylase (ODC) by the tumour promoter phorbol 12–myristate 13–acetate (TPA).\(^{(23)}\)

Myrtocyan and one of its anthocyanin constituents have been shown to have anti–ulcer activity in various experimental models of acute gastric ulcer and in chronic ulcer induced by acetic acid.\(^{(24)}\) The mechanism for this may be by potentiation of the defensive barriers of the gastrointestinal mucosa, such as the secretion of gastric mucus or stimulation of cellular regeneration.\(^{(24)}\)

Extracts of \textit{V. myrtillus} leaves have demonstrated antibacterial activity against several species, including \textit{Staphylococcus aureus} and \textit{Escherichia coli}, as determined by the hole–plate diffusion method and the microdilution broth
method.\textsuperscript{(25)} \textit{V. myrtillus} fruit extracts were less active.

The pharmacokinetics of \textit{V. myrtillus} anthocyanins have been studied in rats.\textsuperscript{(26)} Following a single oral administration, plasma anthocyanin concentrations peaked after 15 minutes and declined rapidly within 2 hours. No hepatic first-pass effect was observed; elimination occurred mostly through the urine and bile.

**Clinical studies**

Clinical studies with extracts of \textit{V. myrtillus} fruits (berries) have focused mainly on its therapeutic applications in certain ophthalmological conditions and in altered microcirculation and peripheral venous insufficiency. The clinical efficacy of \textit{V. myrtillus} fruits has been reviewed.\textsuperscript{(1)}

A study involving 30 healthy subjects with normal platelet aggregation investigated the effects of administration of \textit{V. myrtillus} anthocyanins (Myrtocyan) (480 mg) daily, ascorbic acid 3 g daily and \textit{V. myrtillus} anthocyanins plus ascorbic acid on collagen- and ADP-induced platelet aggregation.\textsuperscript{(11)} Platelet aggregation in blood samples taken from participants after 30 and 60 days’ treatment was clearly reduced in all subjects compared with baseline values. The reduction in platelet aggregation was greater in subjects who received \textit{V. myrtillus} anthocyanins alone than in those who received ascorbic acid alone and was most marked in subjects who received both preparations. Platelet aggregation returned to baseline values when tested 120 days after discontinuation of treatment.\textsuperscript{(11)}

Early studies involving healthy subjects and patients with visual disorders who received \textit{V. myrtillus} extracts alone or in combination with β-carotene and vitamin E reported improvements in night vision and faster adjustment to darkness and restoration of visual acuity following exposure to a bright flash of light.\textsuperscript{(1)} Other studies reported improvements in retinal sensitivity and the visual field in patients with myopia or glaucoma following short- or long-term (six months) treatment with \textit{V. myrtillus} anthocyanins.\textsuperscript{(1)} However, all these studies appear to have been uncontrolled. Other uncontrolled studies in small numbers of patients with retinal pathologies have reported improvements in retinal function, compared with pretreatment values (e.g. ref. 27).

In a randomised, double-blind, placebo-controlled trial, 40 patients with diabetic and/or hypertensive retinopathy received Myrtocyan (160 mg) twice daily or placebo for one month.\textsuperscript{(28)} At the end of the study, the placebo group received Myrtocyan for one month. It was reported that 77–90\% of treated patients experienced improvement compared with the pretreatment period, as determined by ophthalmoscopy and fluorescein fundus angiography.\textsuperscript{(28)}
However, there does not appear to have been a statistical comparison between the treatment and placebo groups. A similar placebo–controlled trial involving 40 patients with early–phase diabetic retinopathy who received Myrtocyan for 12 months also reported improvements in Myrtocyan–treated patients.\(^{(29)}\)

In a randomised, double–blind trial involving 51 patients with mild senile cortical cataract who received *V. myrtillus* anthocyanins plus vitamin E twice daily for four months, treated patients showed significant improvements in lens opacity compared with placebo recipients.\(^{(30)}\)

Studies involving patients with peripheral vascular disorders of various origins are stated to have demonstrated clinical benefits with *V. myrtillus* extracts.\(^{(1)}\) Other studies in patients with ulcerative dermatitis secondary to post–thrombotic or venous varicose stasis, capillary fragility secondary to liver disorders and other conditions, or chronic venous insufficiency have been reported to have shown improvements in clinical signs and symptoms.\(^{(1)}\) However, several of these studies appear to have been uncontrolled (e.g. refs 31–33) and/or included only small numbers of patients (e.g. refs 31 and 32). A double–blind, placebo–controlled study involving 47 patients with peripheral vascular disorders reported reductions in subjective symptoms, such as paraesthesia, pain and heaviness and improved oedema in patients treated with Myrtocyan (480 mg/day) for 30 days.\(^{(1)}\) A single–blind study involving 60 patients with venous insufficiency who received Myrtocyan (480 mg/day) or placebo for 30 days reported significant improvements in oedema, paraesthesia, cramp–like pain and pressure sensation in Myrtocyan–treated patients compared with pretreatment values in these patients.\(^{(1)}\)

*V. myrtillus* anthocyanins have been investigated in a variety of other disorders.

A randomised, double–blind, placebo–controlled trial of *V. myrtillus* anthocyanins (320 mg/day) taken for three days before menstruation was conducted involving 30 patients with chronic primary dysmenorrhoea.\(^{(34)}\) Significant differences between the active treatment and placebo groups were reported for several symptoms investigated, including nausea and vomiting and breast tenderness; there was no effect on headache.

A trial involving 60 patients who had undergone haemorrhoidectomy who were randomised to receive *V. myrtillus* anthocyanins (320–480 mg/day) postoperatively in addition to usual medical care or to no additional treatment reported reductions in itch and oedema, but no effect on other symptoms, in bilberry recipients.\(^{(35)}\)
Other studies, all of which were uncontrolled, have reported beneficial effects following administration of *V. myrtillus* extracts in patients with fibrocystic mastopathy\(^{(36)}\) and type II diabetes mellitus,\(^{(37)}\) in infantile dyspepsia\(^{(38)}\) and in pregnant women with lower limb venous insufficiency and acute–phase haemorrhoids.\(^{(39)}\)
Side-effects, Toxicity

A review of clinical trials of *V. myrtillus* extracts stated that no adverse effects had been observed, even following prolonged treatment.\(^1\) However, most trials involved relatively small numbers of patients and, therefore, would only be able to detect very common acute adverse effects.

The same review summarised the results of an unpublished postmarketing surveillance study which had involved 2295 subjects who had taken Myrto cyan, usually 160 mg twice daily for 1–2 months, for lower limb venous insufficiency, capillary fragility, functional changes in retinal microcirculation or haemorrhoids. Ninety–four subjects reported side–effects, mainly relating to the skin and gastro intestinal and nervous systems.\(^1\)

Long–term consumption of bilberry leaves may lead to toxicity. Chronic administration of doses of 1.5 g/kg per day or more to animals has been reported to be fatal.\(^{G2}\)

Unpublished animal toxicity data for Myrtocyan have also been summarised.\(^1\) In mice and rats, the LD\(_{50}\) for Myrtocyan is over 2000 mg/kg and, in dogs, single doses of 3000 mg/kg produced no adverse effects other than marked darkening of urine and faeces (demonstrating absorption). Oral daily doses to rats and dogs of 125–500 and 80–320 mg/kg, respectively, for six months did not induce mortality or toxic effects.\(^1\) Pharmacokinetic studies of *V. myrtillus* anthocyanins in rats demonstrated that anthocyanins are removed rapidly from the systemic circulation within 2 hours of oral administration.\(^{26}\)
Contra–indications, Warnings

In view of the inhibitory effects of *V. myrtillus* anthocyanins on platelet aggregation, the use of bilberry concurrently with other antiplatelet agents and anti coagulants may enhance the risk of bleeding.

**Pregnancy and lactation**

In an uncontrolled study, *V. myrtillus* anthocyanin extract (Tegens) (80 or 160 mg) twice or three times daily for three months was administered to pregnant women with lower limb venous insufficiency and acute–phase haemorrhoids with no apparent adverse effects.\(^{(39)}\) However, the safety of bilberry has not been established and, in view of the lack of toxicity data, the use of bilberry during pregnancy and lactation should be avoided.
The chemistry of bilberry is well documented and there is good evidence that the anthocyanin constituents are responsible for the pharmacological effects of bilberry.

Documented scientific evidence from *in vitro* and animal studies provides supportive evidence for some of the uses of bilberry. There have been several clinical studies investigating the effects of bilberry in a range of conditions. However, many studies have been uncontrolled, involved only small numbers of patients and had other methodological flaws. Further, well–designed clinical trials are required to establish the efficacy of bilberry.

There are some limited toxicity and safety data for bilberry which together with data on adverse effects reported in clinical trials provide some support for the safety of bilberry when used at recommended doses in the short term. However, further data on the long–term safety of bilberry use are required and, therefore, excessive use of bilberry should be avoided.

Patients wishing to use bilberry for medicinal purposes should be advised to consult a pharmacist, doctor or other suitably trained health care professional for advice.
References

See also General References G2 G3 G31 G36 G43 G50 G55.


31. Coget J, Merlen JF. Étude clinique d’un nouvel agent de protection vasculaire le difrarel 20, composé d’anthocyanosides extraits du *Vaccinium*


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Bloodroot
Species (Family)

*Sanguinaria canadensis* L. (Papaveraceae)
Synonym(s)
Red Indian Paint, Red Root, Sanguinaria, Tetterwort
Part(s) Used

Rhizome
Pharmacopoeial and Other Monographs

BHP 1983\(^{(G7)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

Bloodroot is not included in the GSL."^{G37}"
Constituents

See General References G22 G41.

Alkaloids
Isoquinoline type. 3.0–7.0%.\(^{(1)}\) Sanguinarine (approx. 1%), sanguidimerine, chelerythrine, protopine; others include oxysanguinarine, \(\alpha\)- and \(\beta\)-allocryptopine, sanguilutine, dihydrosanguilutine, berberine, coptisine and homochelidonine.

Other constituents
Resin, starch, organic acids (citric, malic).

Alkaloid content of other plant parts recorded as 0.08% (leaf), 1.8% (root).
Food Use

Bloodroot is listed by the Council of Europe as a natural source of food flavouring (category N3). This category indicates that bloodroot can be added to foodstuffs in the traditionally accepted manner, although there is insufficient information available for an adequate assessment of potential toxicity.\(^{(G16)}\)
Herbal Use

Bloodroot is stated to act as an expectorant, spasmolytic, emetic, cathartic, antiseptic, cardioactive, topical irritant and escharotic (scab-producing). Traditionally it is indicated for bronchitis (subacute or chronic), asthma, croup, laryngitis, pharyngitis, deficient capillary circulation, nasal polyps (as a snuff), and specifically for asthma and bronchitis with feeble peripheral circulation.\(^{(G7)}\)
Dosage

**Rhizome**
0.06–0.5 g (1–2 g for emetic dose) three times daily.\(^{(G7)}\)

**Liquid extract**
0.06–0.3 mL (1 : 1 in 60% alcohol) (1–2 mL for emetic dose) three times daily.\(^{(G7)}\)

**Tincture**
0.3–2 mL (1 : 5 in 60% alcohol) (2–8 mL for emetic dose) three times daily.\(^{(G7)}\)
Pharmacological Actions

Activities documented for bloodroot are principally attributable to the isoquinoline alkaloid constituents, in particular sanguinarine. In the last 10 years, interest has focused on the use of sanguinarine in dental hygiene products. Unless otherwise stated, the following actions refer to sanguinarine.

*In vitro* and animal studies

Considerable antimicrobial activity has been documented against both Gram–positive and Gram–negative bacteria, *Candida* and dermatophytes (fungi), and *Trichomonas* (protozoa).\(^{(2)}\) In addition, anti–inflammatory activity has been described against carrageenan–induced rat paw oedema.\(^{(3)}\)

Prolongation of the ventricular refractory period has been attributed to an inhibition of Na\(^+\)K\(^+\) ATPase.\(^{(4,5)}\) However, a single intravenous injection of sanguinarine to anaesthetised dogs reportedly exerted no effect on cardiovascular parameters monitored.\(^{(4)}\)

*In vitro* inhibition of bone resorption and collagenase has been documented.\(^{(2)}\)

Clinical studies

Many studies have investigated the efficacy of bloodroot extracts in oral hygiene.\(^{(2)}\) Preparations containing bloodroot extracts, such as oral rinses and toothpastes, have been reported to significantly lower plaque, gingival and bleeding indices.\(^{(2)}\) Alteration of the oral microbial flora, or development of resistant microbial strains has not been observed with the use of bloodroot extracts.\(^{(2)}\)
Side-Effects, Toxicity

None documented for bloodroot. Much has been documented concerning the potential toxicity of the alkaloid constituents in bloodroot, in particular of sanguinarine.

In the 1920s contamination of cooking oil with Argemone mexicana seed oil was proposed as the causative factor for epidemic dropsy and associated glaucoma, with sanguinarine considered the toxic component of the seed oil.\(^1,6,7\) However, subsequent workers disputed this theory and the toxicity of A. mexicana oil has been attributed to a fatty acid constituent.\(^1,7\)

Conclusions reached in the 1960s over the carcinogenic potential of sanguinarine have more recently been disproved.\(^8\) In addition, negative mutagenic activity has been observed in the Ames test (microbial, with and without activation).\(^8\)

Sanguinarine is poorly absorbed from the gastrointestinal tract. This is reflected in stated acute oral LD\(_{50}\) values (rat) of 1.7 g/kg (sanguinarine) and 1.4 g/kg (sanguinaria extract), compared with an acute intravenous LD\(_{50}\) (rat) value of 28.7 mg/kg (sanguinarine).\(^1\) Symptoms of diarrhoea, ataxia and reduced activity were observed in animals receiving high oral doses of sanguinarine.\(^5\) The acute dermal toxicity (LD\(_{50}\)) of sanguinarine is stated to be greater than 200 mg/kg in rabbits.\(^1\) The first experimental study of sanguinarine toxicity (1876) reported prostration and severe respiratory distress as the most marked signs of oral toxicity.\(^1\) However, in more recent short-term toxicity studies no toxic signs were observed in the fetuses of rats following maternal administration of 5–30 mg/kg/day of sanguinarine.\(^1\)

The reproductive and developmental toxicity potential of an S. canadensis extract has been evaluated in rats and rabbits.\(^8\) Developmental toxicity (increase in postimplantation loss, slight decrease in fetal and pup body weights) was only evident at maternally toxic doses. No effect was reported on reproductive capabilities, on parturition or on lactation. It was concluded that oral ingestion of sanguinaria extract has no selective effect on fertility, reproduction, or on fetal or neonatal development.\(^8\)

Hepatotoxicity has been documented in rats following a single intraperitoneal administration (10 mg/kg) of sanguinarine.\(^5\) Toxicity was indicated by an increase in serum alanine aminotransferase and serum asparate aminotransferase activity, and by a significant reduction in microsomal cytochrome P450 and benzfetamine N-demethylase activities.\(^5\) Macroscopic lesions were also observed but the authors stated that the two events could
not be conclusively directly related.\textsuperscript{(5)} No hepatotoxicity has been observed in short-term toxicity studies involving oral administration of sanguinarine.\textsuperscript{(1)} Animal studies have indicated sanguinarine to be non-irritant and to exhibit no allergenic or anaphylactic potential.\textsuperscript{(4)} Human patch tests have shown sanguinarine to be non-irritant and non-sensitising.\textsuperscript{(4)}
Contra–indications, Warnings

None documented.

**Pregnancy and lactation**
Animal studies have indicated bloodroot to be non–toxic during pregnancy (see above). However, in view of its pharmacologically active constituents, use of bloodroot during pregnancy and lactation is best avoided.
Pharmaceutical Comment

Bloodroot is characterised by isoquinoline alkaloid constituents (benzophenanthridine–type), predominantly sanguinarine. A wide range of pharmacological activities has been documented for this class of compounds including antimicrobial, anti-inflammatory, antihistaminic, cardiotonic and antiplaque.\(^{(1,8)}\) Other benzophenanthridine alkaloids have been associated with cytotoxic activities. However, recent interest over the potential use of bloodroot in oral hygiene has stimulated considerable research into both sanguinarine and bloodroot extracts. Results have indicated that products such as oral rinses and toothpastes containing either sanguinaria extracts or sanguinarine may be of value in dental hygiene, and are of low toxicity.
References

See also General References G7 G16 G22 G31 G36 G37 G41 G43.


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Species (Family)

*Iris versicolor* L. or *Iris caroliniana* Watson (Iridaceae)
Part(s) Used

Rhizome
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL$^{G37}$
Constituents
See General References G22 G40 G48 G64.

**Acids**
Isophthalic acid 0.002%, salicylic acid, lauric acid, stearic acid, palmitic acid and 1-triacontanol.

**Volatile oils**
0.025%. Furfural.

**Other constituents**
Iridin, β-sitosterol, iriversical\(^1\) and tannin.
Food Use

Blue flag is not used in foods.
Herbal Use

Blue flag is stated to possess cholagogue, laxative, diuretic, dermatological, anti-inflammatory and anti-emetic properties. It has been used for skin diseases, biliousness with constipation and liver dysfunction, and specifically for cutaneous eruptions.\(^{(G7 \, G64)}\)
**Dosage**

**Dried rhizome**
0.6–2.0 g or by decoction three times daily.\(^{(G6 G7 G10)}\)

**Liquid extract**
1–2 mL (1 : 1 in 45% alcohol) three times daily.\(^{(G6 G7 G10)}\)
Pharmacological Actions

None documented.
Side-effects, Toxicity

It has been stated that the fresh root of blue flag can cause nausea and vomiting.\(^\text{G42}\)

Furfural, a volatile oil constituent, is known to be irritant to mucous membranes causing lachrymation, inflammation of the eyes, irritation of the throat, and headache.\(^\text{G48}\) Whether these irritant properties are attributable to the volatile oil of blue flag has not been established. Acute oral toxicity (rat, LD\(_{50}\)) for furfural has been documented as 127 mg/kg body weight.\(^\text{G48}\)

Iridin has been reported to be poisonous in both humans and livestock.\(^\text{G22}\) However, it is unclear whether this substance is the same iridin documented as a constituent of blue flag.
Contra-indications, Warnings

Only small doses of the dried root are advisable, because of the risk of nausea and vomiting.\(^\text{G}42\) In view of the possible irritant nature of the volatile oil, blue flag may not be suitable for internal use, especially in sensitive individuals.

**Pregnancy and lactation**

The safety of blue flag has not been established. In view of this, together with the documented irritant properties of some of the constituents, blue flag should not be taken during pregnancy.
Pharmaceutical Comment

Little is known about the phytochemical, pharmacological or toxicological properties of blue flag and its constituents, although related species are known to be toxic. In view of these factors, the use of blue flag is best avoided.
References

See also General References G6 G9 G10 G22 G31 G36 G37 G40 G42 G43 G48 G64.


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Bogbean
Species (Family)

*Menyanthes trifoliata* L. (Menyanthaceae)
Synonym(s)

Buckbean, Marsh Trefoil, Menyanthes
Part(s) Used

Leaf
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)

BP 2002\(^{(G71)}\)

Complete German Commission E\(^{(G3)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)

Ph Eur 2004\(^{(G72)}\)
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents
See General References G2 G62 G64.

Acid
Caffeic acid, chlorogenic acid, ferulic acid, \( p \)-hydroxybenzoic acid, protocatechuic acid, salicylic acid, vanillic acid;\(^{(1,2)} \) folic acid and palmitic acid.\(^{(2)} \)

Alkaloids
Gentianin and gentianidine (pyridine-type); choline.\(^{(2)} \)

Coumarins
Scopoletin.\(^{(2)} \)

Flavonoids
Hyperin, kaempferol, quercetin, rutin and trifolioside.\(^{(1,2)} \)

Iridoids
7′,8′-Dihydrofoliamethin, foliamethin, loganin, menthiacolin and sweroside.\(^{(2–4)} \)

Other constituents
Carotene, ceryl alcohol, enzymes (e.g. emulsin, invertin), \( \alpha \)-spinasterol, an unidentified substance with haemolytic properties.\(^{(2)} \) \( \alpha \)-Spinasterol has been reported to be a mixture of five sterols with \( \alpha \)-spinasterol and stigmast-7-enol as major components.\(^{(5)} \)
Food Use

Bogbean is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that bogbean can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\textsuperscript{(G16)}
Herbal Use

Bogbean is stated to possess bitter and diuretic properties. It has been used for rheumatism, rheumatoid arthritis, and specifically for muscular rheumatism associated with general asthenia. (G2 G7 G8 G64)
Dosage

*Dried leaf*  
1–2 g or by infusion three times daily.\(^{(G7)}\)

*Liquid extract*  
1–2 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)

*Tincture*  
1–3 mL (1 : 5 in 45% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

**In vitro and animal studies**

A choleretic action has been described for caffeic acid and ferulic acid; a stomachic secretive action has been reported for protocatechuic acid and \( p \)-hydroxybenzoic acid. The iridoids possess bitter properties.\(^{(1)}\) The bitter index (BI) of bogbean is stated to be 4000–10 000 (compared to gentian BI 10 000–30 000).\(^{(G62)}\) Bogbean extracts have antibacterial activities.\(^{(6,7)}\)
Side-effects, Toxicity

Large doses of bogbean are stated to be purgative and may cause vomiting.\(^8\) An unidentified substance with haemolytic activity has been isolated from bogbean.\(^2\)
Contra-indications, Warnings

Excessive doses may be irritant to the gastrointestinal tract, causing diarrhoea, griping pains, nausea and vomiting.\(^8\)

**Pregnancy and lactation**

The safety of bogbean has not been established. In view of the lack of toxicity data and possible purgative action, the use of bogbean during pregnancy and lactation should be avoided.
Pharmaceutical Comment

The chemistry of bogbean is well studied, but no pharmacological information is available to justify the herbal uses. In view of the lack of toxicity data, excessive doses should be avoided.
References

See also General References G2 G3 G8 G9 G16 G31 G36 G37 G43 G56 G62 G64.

Species (Family)

*Peumus boldus* Molina (Monimiaceae)
Synonym(s)

Boldus, *Boldus boldus* (Mol.) Lyons
Part(s) Used

Leaf
Pharmacopoeial and Other Monographs

BHP 1996$^{(G9)}$

BP 2002$^{(G71)}$

Complete German Commission E$^{(G3)}$

ESCOP 1996$^{(G52)}$

Martindale 33rd edition$^{(G67)}$

PDR for Herbal Medicines 2nd edition$^{(G36)}$

Ph Eur 2004$^{(G72)}$

WHO volume 1 1999$^{(G63)}$
Legal Category (Licensed Products)

GSL\textsuperscript{(G37)}
Constituents

See General References G2 G22 G41 G52 G58 G62 G64.

Alkaloids
Isoquinoline–type. 0.25–0.7%. Pharmacopoeial standard not less than 0.1% alkaloid calculated as boldine. Boldine 0.06% (major, disputed), isoboldine, 6a,7-dehydroboldine, isocorydine, isocorydine-N-oxide, norisocorydine, laurolitsine, laurotetanine, N-methyllaurotetanine, reticuline (aporphines); (−)-pronuciferine (proaporphine) and sinoacutine (morphinandienone).\(^1\text{-}^4\)

Flavonoids
Flavonols (e.g. isorhamnetin) and their glycosides.\(^5\text{-}^6\)

Volatile oils
2.5%. Some 38 components have been identified, including \(p\)-cymene 28.6%, ascaridole 16.1%, 1,8-cineole 16.0%, linalool 9.1%, terpinen-4-ol 2.6%, α-terpineol 0.9%, fenchone 0.8% and terpinolene 0.4%.

Other constituents
Coumarin 0.5%, resin and tannin.
Food Use

Boldo is listed by the Council of Europe as a natural source of food flavouring (category N3). This category indicates that boldo can be added to foodstuffs in the traditionally accepted manner, although insufficient information is available for an adequate assessment of potential toxicity.\(^{(G16)}\) In the USA, boldo is approved for food use in alcoholic beverages only.\(^{(G64)}\)
Herbal Use

Boldo is stated to possess cholagogue, liver stimulant, sedative, diuretic, mild urinary demulcent, and antiseptic properties. It has been used for mild digestive disturbances, constipation, gallstones, pain in the liver or gall bladder, cystitis, rheumatism, and specifically for cholelithiasis with pain. The German Commission E approved use for treatment of dyspepsia and mild spastic gastrointestinal complaints.
**Dosage**

*Dried leaf*
60–200 mg or by infusion three times daily;\(^{(G7)}\) 2–5 g as a tea.\(^{(G52)}\)

*Liquid extract*
0.1–0.3 mL (1 : 1 in 45% alcohol) three times daily.\(^{(G7)}\)

*Tincture*
0.5–2.0 mL (1 : 10 in 60% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

**In vitro and animal studies**

Boldo has exhibited choleretic (highest activity in rats), diuretic, stomachic and cholagogic properties.\(^{(G41 G52)}\) The choleretic activity may be due to synergy between flavonoids and alkaloids.\(^{(G52)}\) Experiments in rats have failed to demonstrate choleretic activity after oral administration of 400 or 800 mg/kg aqueous ethanolic extract, intraduodenal administration of 200 mg or 800 mg/kg, and intravenous administration of 32.5–130 mg/kg of a dry ethanolic extract.\(^{(7)}\)

An aqueous ethanolic extract (equivalent to 0.5–1.0 mg/mL dried ethanolic extract) and also boldine (33 μg/mL) gave significant hepatoprotection against t-butyl hydroperoxide–induced hepatotoxicity in rat hepatocytes *in vitro*.\(^{(7)}\) Boldine at a concentration of 0.015 mol/L inhibited microsomal lipid peroxidation in a rat liver preparation by 50%.\(^{(8)}\) A dried aqueous ethanolic extract (0.06–0.115%) of boldine at a dose of 500 mg/kg gave 70% protection against carbon tetrachloride–induced hepatotoxicity in mice, and boldine alone (10 mg/kg) gave 49% protection.\(^{(7)}\) An aqueous ethanolic extract of boldo at doses of 50 and 100 mg/kg administered intraperitoneally showed anti–inflammatory activity in the rat paw carrageenan–induced oedema test, whereas boldine alone appeared to be inactive.\(^{(7)}\)

Boldine showed concentration–dependent relaxant activity on isolated rat ileum (EC\textsubscript{50} 1.7 × 10\textsuperscript{-4} mol/L), and acted as a competitive antagonist of acetylcholine and as a non–competitive antagonist of barium.\(^{(9)}\) Boldine at low micromolar concentrations prevented oxidation in rat brain homogenate and lipid peroxidation of red cell plasma membranes, led to inactivation of lysozymes, indicating high reactivity of free radicals.\(^{(10)}\)

Boldo essential oil contains terpinen–4–ol, the irritant and diuretic principle in juniper oil.

**Clinical studies**

Boldo, in combination with cascara, rhubarb and gentian, has been reported to exhibit a beneficial effect on a variety of symptoms such as loss of appetite, digestion difficulties, constipation, flatulence and itching.\(^{(11,12)}\) Rhubarb and gentian were found to be more effective with respect to appetite–loss related symptoms, and boldo and cascara more effective in constipation–related symptoms.
Two preparations containing extracts of boldo and cascara have been documented to increase biliary flow without altering the lithogenic index or bile composition.\(^{(13)}\) Boldine may provide relief to patients with gallstones for whom surgery is not an option or drugs have not been effective.\(^{(G56)}\) The choloretic action of boldine releases bile and its diuretic action increases fluid secretion, possibly cleansing sediment or bacteria from the biliary tract.\(^{(G56)}\) Treatment of 12 human volunteers with boldo dry extract resulted in prolongation of intestinal transit time.\(^{(G4)}\)

Ascaridole, a component of the volatile oil, previously found a clinical use as an anthelmintic agent.\(^{(14)}\) However, this use has declined with the development of synthetic compounds with lower toxicity and a wider range of activity.
Side-effects, Toxicity

Boldo volatile oil is stated to be one of the most toxic oils. Application of the undiluted oil to the hairless backs of mice has an irritant effect. The oil contains irritant terpenes, including terpinen-4-ol, the irritant principle in juniper oil.

An acute oral LD$_{50}$ value for boldo oil has been given as 0.13 g/kg body weight in rats, with doses of 0.07 g/kg causing convulsions. The acute dermal LD$_{50}$ in rabbits has been reported as 0.625–1.25 g/kg. No acute toxicity was observed in rats given oral doses of 3 g/kg of dry aqueous ethanolic extract. In mice, an aqueous ethanolic extract (1 : 1) had an LD$_{50}$ of 6 g/kg (intraperitoneal administration). The LD$_{50}$ values of total alkaloids and of boldine in mice were 420 and 250 mg/kg (intraperitoneal administration), respectively. Total alkaloids (intraperitoneal administration) given to dogs produced vomiting, diarrhoea and epileptic symptoms with a recovery after 50 minutes.

Boldine was not genotoxic as indicated by the SOS chromotest with Escherichia coli, or in the Ames test, and did not induce mutations in Saccharomyces cerevisiae. Boldine did not induce an increase in the frequency of chromosome aberrations in human lymphocytes in vitro, or in mouse bone marrow cells in vivo. There were no signs of genotoxicity in mouse bone marrow, as assessed by the micro nucleus test.
Contra–indications, Warnings

Excessive doses of boldo may cause renal irritation, because of the volatile oil, and should be avoided by individuals with an existing kidney disorder. Boldo is contraindicated in individuals with obstruction of bile duct or severe liver disease. For gallstone patients, it should only be used after consultation with a physician.\textsuperscript{(G3)} Ascaridole is toxic and use of the oil is not recommended.\textsuperscript{(G58)}

Pregnancy and lactation
The safety of boldo taken during pregnancy has not been established. In view of the potential irritant nature of the volatile oil, the use of boldo during pregnancy should be avoided.
Pharmaceutical Comment

The chemistry of boldo is well documented, and some pharmacological data are available. Clinical studies have described choleretic activity, although further well-designed studies are required to establish this. The reputed diuretic and mild urinary antiseptic properties of boldo are probably attributable to the irritant volatile oil. In view of the toxicity data and the irritant nature of the volatile oil, excessive use of boldo should be avoided. Boldo is not recommended for long-term use. (G56)
References


Boneset
Species (Family)

*Eupatorium perfoliatum* L. (*Asteraceae/Compositae*)
**Synonym(s)**

Feverwort, Thoroughwort. Snakeroot has been used to describe poisonous *Eupatorium* species.
Part(s) Used
Herb
Pharmacopoeial and Other Monographs

BHP 1983\(^{(G7)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR\(^{®}\) for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL – as Boneset and Eupatorium.\(^{(G37)}\)
Constituents
See General References G22 G41 G48 G64.

Flavonoids
Flavonol (kaempferol, quercetin) glycosides including astragalin, hyperoside and rutin; eupatorin (flavone) and dihydroflavonols.\(^1\)

Terpenoids
Sesquiterpene lactones including euperfolin and euperfolitin (germacranolides), eufoliatin (guianolide), eufoliatorin (dilactone guaiane) and euperfolide.\(^2\) Sesquiterpenes, diterpenes (dendroidinic acid, hebeclinolide), triterpenes (α-amyrin, dotriacontane) and sterols (sitosterol, stigmasterol).

Other constituents
Volatile oil, resin, wax, tannic and gallic acids, bitter glucoside, inulin, polysaccharides and sugars.
Food Use

Boneset is not used in foods.
Herbal Use

Boneset is stated to possess diaphoretic and aperient properties. Traditionally, it has been used for influenza, acute bronchitis, nasopharyngeal catarrh, and specifically for influenza with deep aching, and congestion of the respiratory mucosa. (G7 G64)
Dosage

**Herb**
1–2 g or by infusion three times daily.\(^{(G7)}\)

**Liquid extract**
1–2 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)

**Tincture**
1–4 mL (1 : 5 in 45% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

*In vitro* and animal studies

Immunostimulant activity (*in vitro* stimulation of granulocyte phagocytic activity) has been demonstrated by high dilutions (10\(^{-5}\)–10\(^{-7}\) g/100 mL) of various sesquiterpene lactones isolated from *E. perfoliatum*\(^{(3)}\). In addition, immuno stimulating actions (granulocyte, macrophage and carbon clearance tests) have been documented for polysaccharide fractions from *E. perfoliatum*\(^{(3,4)}\).

An ethanol extract of the whole plant has exhibited weak anti–inflammatory activity in rats\(^{(G41)}\). Many activities have been documented for flavonoid compounds including anti–inflammatory activity.
Side-effects, Toxicity

Contact dermatitis has been reported for *Eupatorium* species but not specifically for boneset (*E. perfoliatum*).\(^{(G51)}\) Cytotoxic properties have been documented for a related species, *E. cannabinum*, and are attributed to the sesquiterpene lactone eupatoriopicrin. This compound has not been documented as a constituent of boneset. Hepatotoxic pyrrolizidine alkaloids (PAs) have been isolated from various *Eupatorium* species although none have been documented as constituents of boneset (*E. perfoliatum*).\(^{(5)}\)

Instances of allergic and anaphylactic reactions have been associated with the sesquiterpene lactone constituents in German chamomile, although no reactions specifically involving boneset have been documented.

The US Food and Drugs Administration (FDA) has classified boneset as a herb of undefined safety.\(^{(G22)}\)
Contra–indications, Warnings

The allergenic potential of sesquiterpene lactones is well recognised. Individuals with a known hypersensitivity to other members of the Asteraceae family (e.g. chamomile, feverfew, ragwort, tansy) should avoid using boneset. Individuals with existing hypersensitivities/allergies should use boneset with caution.

Pregnancy and lactation
The safety of boneset taken during pregnancy has not been established. In view of the lack of toxicity data and the possibility of constituents with allergenic activity, the use of boneset during pregnancy and lactation should be avoided.
Pharmaceutical Comment

The constituents of boneset are fairly well documented and include many pharmacologically active classes such as flavonoids, sesquiterpene lactones (typical for the Asteraceae family) and triterpenes. Immunostimulant activity (in vitro) has been reported for sesquiterpene lactone and polysaccharide components, possibly supporting the traditional use of boneset in influenza. Many pharmacological studies have focused on the cytotoxic/antitumour actions of sesquiterpene lactone components of various Eupatorium species, although these actions have not been reported for sesquiterpene lactones isolated from boneset. Little is known regarding the toxicity of boneset. Hepatotoxic pyrrolizidine alkaloids, which have been documented for other Eupatorium species, have not been reported for boneset.
References

See also General References G7 G20 G22 G31 G32 G36 G37 G41 G48 G51 G64.


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Borage
Species (Family)

*Borago officinalis* L. (Boraginaceae)
**Synonym(s)**

Beebread, Bee Plant, Burrage, Starflower (oil)
Part(s) Used

Herb
Pharmacopoeial and Other Monographs

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Product)

Borage is not included in the GSL.\(^{(G37)}\)
Constituents

See General References G22 G64.

Alkaloids
Pyrrolizidine–type. Lycopsamine, intermedine, acetyllycopsamine, acetylintermedine, amabiline, supinine and thesinine (unsaturated).\(^{(1,2)}\) Concentrations reported as 0.01% and 2–10 ppm for commercial dried samples. Alkaloid concentrations reportedly the same for fresh and dried samples; fresh samples revealed alkaloids as the free base in the roots and mainly as \(N\)-oxides in the leaves.

Mucilages
11.1%. Yielding glucose, galactose and arabinose.

Oil
Rich in fatty acids, in particular gamolenic acid.

Other constituents
Acids (acetic, lactic, malic, silicic), cyanogenetic compounds and tannins (up to 3%).
**Food Use**

Borage is occasionally used in salads and soups.
Borage is stated to possess diaphoretic, expectorant, tonic, anti-inflammatory and galactogogue properties.\(^3\) Traditionally, borage has been used to treat many ailments including fevers, coughs and depression.\(^{3,G42}\) Borage is also reputed to act as a restorative agent on the adrenal cortex.\(^3\) Borage oil (starflower oil) is used as an alternative source to evening primrose oil for gamolenic acid.
Dosage

Infusion
Two 5–mL spoonfuls of dried herb to one cup boiling water three times daily.(3)

Tincture
1–4 mL three times daily.(3)
Pharmacological Actions

In vitro and animal studies

Borage oil has been reported to attenuate cardiovascular reactivity to stress in rats.\(^4\)

Clinical studies

The effect of borage seed oil on the cardiovascular reactivity of humans to acute stress has been studied in 10 individuals, who each received a total daily dose of 1.3 g for 28 days.\(^4\) The individuals were required to undertake an acute psychological task requiring sensory intake and vigilance (Stroop colour test). Borage oil was found to attenuate cardiovascular reactivity to stress, indicated by a reduction in systolic blood pressure and heart rate and by increased task performance. The specific mechanisms by which borage exerts this effect were unknown but a central mechanism of action of the fatty acids was suggested in view of the simultaneous reduction in heart rate and blood pressure.\(^4\)
Side–effects, Toxicity

No side–effects of borage have been identified. Borage contains low concentrations of unsaturated pyrrolizidine alkaloids, which are known to be hepatotoxic in both animals and humans (see Comfrey).\(^{(5)}\)
Contra–indications, Warnings

Evening primrose oil is recommended to be used with caution in epileptic patients, especially in those with schizophrenia and/or those taking phenothiazines (see Evening Primrose); as borage oil is used similarly it should also be used with caution. In view of the known toxic pyrrolizidine alkaloid constituents, excessive or prolonged ingestion of borage should be avoided. In particular, infusions (e.g. herbal teas) containing borage should be avoided.

Pregnancy and lactation

In view of the documented pyrrolizidine constituents and lack of toxicity data, borage should not be used during pregnancy or lactation.
Limited information is available on the constituents of borage. No documented pharmacological data were located to support the traditional uses, although the mucilage content supports the use of borage as a demulcent. Interest has focused on the volatile oil as a source of gamolenic acid. Borage contains known toxic pyrrolizidine alkaloids, although at concentrations considerably lower than comfrey for which human toxicity has been documented. However, it would seem wise to avoid excessive or prolonged ingestion of borage. It is unclear whether borage oil, currently available in food supplements, contains any pyrrolizidine alkaloids.
References

See also General References G18 G20 G22 G31 G32 G36 G42 G43.


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Broom
Species (Family)

Sarothamnus scoparius (L.) Koch. (Leguminosae/Papilionaceae)
Synonym(s)

*Cytisus scoparius* (L.) Link, Hogweed, Scoparius, *Spartium scoparium* L.
Part(s) Used

Flowerhead
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

Broom is not included in the GSL.\textsuperscript{(G37)}
Constituents
See General References G2 G40 G41 G48 G62 G64.

Alkaloids
Quinolizidine–type. 0.8–1.5%. Sparteine 0.3–0.8% (major component); minor alkaloids include cytisine (presence disputed), genisteine (d-α-isosparteine), lupanine, oxyisparteine and sarothamine.

Amines
Epinine, hydroxytyramine and tyramine.

Flavonoids
Scoparin and vitexin.

Other constituents
Amino acids, bitter principles, carotenoids, fat, resin, sugars, tannin, wax and volatile oil.
Food Use

Broom is listed by the Council of Europe as a natural source of food flavouring (category N3). This category indicates that broom can be added to foodstuffs in the traditionally accepted manner, although there is insufficient information available for an adequate assessment of potential toxicity.\textsuperscript{(G16)}
Herbal Use

Broom is stated to possess cardioactive, diuretic, peripheral vasoconstrictor and antihaemorrhagic properties. It has been used for cardiac dropsy, myocardial weakness, tachycardia, profuse menstruation and specifically for functional palpitation with lowered blood pressure.\(^{(G2 \ G7 \ G64)}\) Broom is also reported to possess emetic and cathartic properties.\(^{(G41)}\)
Dosage

*Dried tops*
1–2 g as a decoction.\(^{(G7)}\)

*Liquid extract*
1–2 mL (1 : 1 in 25% alcohol).\(^{(G7)}\)

*Tincture*
0.5–2.0 mL (1 : 5 in 45% alcohol).\(^{(G7)}\)
Pharmacological Actions

The pharmacological actions of broom are primarily due to the alkaloid constituents.

In vitro and animal studies

Sparteine is reported to exhibit pharmacological actions similar to those of quinidine. Low doses administered to animals result in tachycardia, whereas high doses cause bradycardia and may lead to ventricular arrest. Sparteine has little effect on the central nervous system (CNS), but peripherally, paralyses motor nerve terminals and sympathetic ganglia as a result of a curare-like action.\(^{(G44)}\)

The flowers, seeds, root and whole herb have been used to treat tumours.\(^{(G41)}\)

Clinical studies

None documented for broom. However, sparteine is known to decrease the irritability and conductivity of cardiac muscle and has been used to treat cardiac arrhythmias,\(^{(G44)}\) restoring normal rhythm in previously arrhythmic patients.\(^{(G2)}\) Sparteine is reported to have a quinidine-like action rather than a digitalis-like action.\(^{(G2)}\) Sparteine is also stated to be a powerful oxytocic drug, which was once used to stimulate uterine contractions.
Side-effects, Toxicity

The alkaloid constituents in broom are toxic. Sparteine sulfate has been reported to be a cardiac depressant and can also produce respiratory arrest.\(^{(G44)}\) Symptoms of poisoning are characterised by tachycardia with circulatory collapse, nausea, diarrhoea, vertigo and stupor.
Contra–indications, Warnings

Broom is stated to be inappropriate for non–professional use.\(^{(G49)}\) Its use is contra–indicated in individuals with high blood pressure\(^{(G49)}\) or a cardiac disorder, because of the alkaloid constituents.

Pregnancy and lactation

The use of sparteine is contra–indicated during pregnancy.\(^{(G42)}\) Sparteine is stated to be a powerful oxytocic drug and is cardio toxic. Broom should not be taken during lactation.
Pharmaceutical Comment

The chemistry of broom is well documented. The pharmacological actions are primarily due to the alkaloid constituents. Sparteine, the major alkaloid component, is a cardiac depressant with actions similar to those of quinidine. Although these actions support the documented traditional herbal uses, broom is not suitable for self-medication.
References
Buchu
Species (Family)

*Agathosma betulina* (Berg.) Pillans (Rutaceae)
Synonym(s)

*Barosma betulina* Bart. & Wendl., Round Buchu, Short Buchu

Note that Oval Buchu refers to *Agathosma crenulata* (L.) Pillans (synonym *Barosma crenulata* (L.) Hook.) and Long Buchu refers to *Agathosma serratifolia* (Curt.) Spreeth (synonym *Barosma serratifolia* (Curt.) Willd.).

\(G^{12}\)
Part(s) Used

Leaf
Pharmacopoeial and Other Monographs

BHC 1992\(^6\)

BHP 1996\(^9\)

Martindale 33rd edition\(^{67}\)

Mills and Bone\(^{50}\)

PDR for Herbal Medicines 2nd edition\(^{36}\)
Legal Category (Licensed Products)

GSL$^{(G37)}$
Constituents

See General References G2 G22 G41 G48 G64.

**Flavonoids**
Diosmetin, quercetin, diosmin, quercetin–3,7-diglucoside, rutin.

**Volatile oils**
1.0–3.5%. Over 100 identified compounds, including diosphenol, limonene, menthone and pulegone as the major components.

**Other constituents**
Mucilage, resin. Coumarins have been reported for many other *Agathosma* species.\(^1\)
Food Use

Buchu is listed by the Council of Europe as a natural source of food flavouring (category N3). This category allows buchu to be added to foodstuffs in the traditionally accepted manner, although there is insufficient information available for an adequate assessment of potential toxicity.\(^{(G16)}\) In the USA, buchu volatile oil is approved for food use with concentrations usually up to about 0.002% (15.4 ppm).\(^{(G16 \ G41)}\)
Buchu is stated to possess urinary antiseptic and diuretic properties. It has been used for cystitis, urethritis, prostatitis, and specifically for acute catarrhal cystitis. (G2 G7 G8 G64)
Dosage

Dried leaf
1–2 g by infusion three times daily. (G6 G7)

Liquid extract
0.3–1.2 mL (1:1 in 90% alcohol). (G6 G7)

Tincture
2–4 mL (1:5 in 60% alcohol). (G6 G7)
Pharmacological Actions

*In vitro and animal studies*

None documented for buchu. Diosmin has documented anti-inflammatory activity against carrageenan–induced rat paw oedema, at a dose of 600 mg/kg body weight.\(^{(2)}\)
Side-effects, Toxicity

None documented for buchu. The volatile oil contains pulegone, a known hepatotoxin (see Pennyroyal).\(^{(G20)}\) The oil may cause gastrointestinal and renal irritation.
Contra-indications, Warnings

Excessive doses of buchu should not be taken in view of the potential toxicity of the volatile oil. Buchu should be avoided in kidney infections.\(^{(G42)}\)

**Pregnancy and lactation**

The safety of buchu has not been established. In view of this, together with the potential toxicity and irritant action of the volatile oil, the use of buchu during pregnancy and lactation should be avoided.
Pharmaceutical Comment

Limited chemical data are available for buchu. No scientific evidence was found to justify the herbal uses, although reputed diuretic and anti-inflammatory activities are probably attributable to the irritant nature of the volatile oil and the flavonoid components, respectively. In view of the lack of documented toxicity data, together with the presence of pulegone in the volatile oil, excessive use of buchu should be avoided.
References


Burdock
Species (Family)

*Arctium majus* Bernh. (*Asteraceae/Compositae*)
Synonym(s)

*Arctium lappa* L. and other *Arctium* species, Lappa
Part(s) Used

Root
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

Martindale 33rd edition\(^{(G67)}\)

Mills and Bone\(^{(G50)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL (G^37)
Constituents
See General References G2 G6 G22 G41 G48 G64.

Acids
Acetic acid, butyric acid, caffeic acid, chlorogenic acid, gamma–guanidino-n-butyric acid, η-guanidino-n-isovaleric acid, trans-2-hexenoic acid, isovaleric acid, lauric acid, linoleic acid, linolenic acid, myristic acid, oleic acid, palmitic acid, propionic acid, stearic acid and tiglic acid.\(^{1-3}\)

Aldehydes
Acetaldehyde, benzaldehyde, butyraldehyde, caproicaldehyde, isovaleraldehyde, propion aldehyde and valeraldehyde.\(^1\)

Carbohydrates
Inulin (up to 45–50%), mucilage, pectin and sugars.

Polyacetylenes
0.001–0.002% dry weight. Fourteen identified compounds include 1,11-tridecadiene–3,5,7,9-tetrayne (50%), 1,3,11-tridecatriene–5,7,9-triyne (30%) and 1-tridecen–3,5,7,9,11-pentayne as the major components;\(^4\) arctinone–a, arctinone–b, arctinol–a, arctinol–b, arctinal, arctic acid–b, arctic acid–c, methyl arctate–b and arctinone–a acetate (sulfur–containing acetylenic compounds).\(^5,6\)

Other constituents
Fats (0.4–0.8%), fixed and volatile oils (0.07–0.18%), sesquiterpene lactones (arctiopicrin),\(^7\) bitters (lappatin), resin, phytosterols (sitosterol and stigmasterol), tannin\(^8\) and lignan–type compound.\(^9-11\)

Other species
Flavonol (kaempferol, quercetin) glycosides are present in Arctium minus (Hill) Bernh.\(^3\)
Food Use

Burdock is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that burdock can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\(^{(G16)}\)
Herbal Use

Burdock is stated to possess diuretic and orexigenic properties. It has been used for cutaneous eruptions, rheumatism, cystitis, gout, anorexia nervosa, and specifically for eczema and psoriasis. (G2 G6 G7 G8 G60)
Dosage

*Dried root*
2–6 g or by infusion three times daily.\(^{(G7)}\)

*liquid extract*
2–8 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)

*Tincture*
8–12 mL (1 : 10 in 45% alcohol) three times daily.\(^{(G7)}\)

*Decoction*
500 mL (1 : 20) per day.\(^{(G7)}\)
Pharmacological Actions

**In vitro and animal studies**

The roots and leaves of burdock plants not yet flowering are stated to possess diuretic, hypoglycaemic and antifurunculous properties.\(^7\) A burdock extract (plant part not stated) was reported to cause a sharp, long–lasting reduction in the blood sugar concentration in rats, together with an increase in carbohydrate tolerance and a reduction in toxicity.\(^{12}\) The antimicrobial activity documented for burdock has been attributed to the polyacetylene constituents,\(^4\) although only traces of these compounds are found in the dried commercial herb.\(^{G62}\)

Furthermore, arctiopicrin is stated to be a bitter with antibiotic activity against Gram–positive bacteria.\(^{7,13}\) Antibacterial activity against Gram–positive (e.g. *Staphylococcus aureus, Bacillus subtilis, Mycobacterium smegmatis*) and Gram–negative (*Escherichia coli, Shigella flexneri, Shigella sonnei*) bacteria has been documented for burdock leaf and flower, whereas the root was only found to be active towards Gram–negative strains.\(^{14}\)

**In vivo** uterine stimulant activity has been reported.\(^{G30}\)

Protection against mutagenic activity has also been documented for burdock.\(^{9,15,16}\)

Burdock reduced the mutagenicity to *Salmonella typhimurium* (TA98, TA100) of mutagens both requiring and not requiring S9 metabolic activation.\(^{10}\) A lignan–like structure was proposed for the desmutagenic factor.\(^9\) **In vivo** studies have shown that fresh or boiled plant juice from burdock may cause a significant reduction in DMBA–induced chromosome aberrations.\(^{16}\)

Burdock has been reported to exhibit antitumour activity.\(^{17}\)

The addition of dietary fibre (5%) from burdock roots to the diet of rats has been documented to provide protection against the toxicity of various artificial food colours.\(^{18}\)
**Side–effects, Toxicity**

A single report of human poisoning with burdock has been documented.\(^{(19)}\) The patient exhibited symptoms of atropine–like poisoning following the ingestion of a commercially packaged burdock root tea. Atropine is not a constituent of burdock, and subsequent analysis indicated that the tea was contaminated with a herbal source of solanaceous alkaloids, possibly belladonna leaf. This report served to highlight the problems which may arise with inadequate quality control of herbal preparations.

The carcinogenicity of burdock was investigated in 12 rats fed dried roots (33% of diet) for 120 days, followed by a normal diet until 480 days.\(^{(20)}\) Ten of the 12 rats survived 480 days and no tumours were detected. A urinary bladder papilloma and an oligodendroglioma were observed in one rat but these were considered to have been induced spontaneously.

Burdock has been reported to exhibit antitumour properties (see *In vitro* and animal studies).
Contra–indications, Warnings

Excessive doses may interfere with existing hypo glycaemic therapy (see \textit{In vitro} and animal studies).

\textbf{Pregnancy and lactation}

\textit{In vivo} uterine stimulant action has been reported.\textsuperscript{(G30)} In view of this, and the lack of toxicity data, the use of burdock during pregnancy and lactation should be avoided.
Pharmaceutical Comment
The chemistry of burdock and related *Arctium* species has been well studied. Various pharmacological activities have been reported in animals although none support the reputed herbal uses. Documented bitter constituents, however, may explain the traditional use of burdock as an orexigenic. In view of the lack of toxicity data, excessive use of burdock should be avoided.
References


Species (Family)

*Sanguisorba officinalis* L. (Rosaceae)
Synonym(s)
Garden Burnet, Greater Burnet, Sanguisorba
Part(s) Used

Herb
Pharmacopoeial and Other Monographs

BHP 1983\(^{(G7)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL (Sanguisorba)\textsuperscript{(G37)}
Constituents

See General References G40 G64.

All phytochemical data located refer to the underground plant parts and not to the herb.

**Flavonoids**
Flavones, unstable flavonol derivatives.

**Saponins**
Ziyu glycosides I and II (major glycosides),\(^1\) pomolic acid as aglycone (not tomentosolic acid as documented in earlier work), sanguisorbin 2.5–4.0%.

**Tannins**
Numerous compounds (condensed and hydrolysable) have been isolated, including 3,3,4–tri-O-methylellagic acid.\(^2\)--\(^6\)

**Other constituents**
Volatile oil, ascorbic acid (vitamin C) in the fresh plant.
Food Use

Burnet is not used in foods.
Herbal Use

Burnet is stated to possess astringent, antihaemorrhagic, styptic and antihaemorrhoidal properties. It has been used for ulcerative colitis, metrorrhagia, and specifically for acute diarrhoea.\(^{(G7 \ G64)}\)
Dosage

**Dried herb**
2–6 g or by infusion three times daily.\(^{(G7)}\)

**Liquid extract**
2–6 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)

**Tincture**
2–8 mL (1 : 5 in 45% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

*In vitro* and animal studies

None documented for burnet. The roots have been reported to contain an antihaemorrhagic principle, 3,3,4-tri-O-methylellagic acid. (2)
Side-effects, Toxicity

None documented.
Contra-indications, Warnings

None documented.

**Pregnancy and lactation**
In view of the lack of phytochemical, pharmacological and toxicity data, the use of burnet during pregnancy and lactation should be avoided.
Pharmaceutical Comment

The chemistry of burnet herb does not appear to have been studied, although data are available for the underground plant parts. If present in the herb as well as the root, the tannin constituents would support the reputed astringent and antihaemorrhagic actions of burnet. In view of the lack of toxicity data and the possible high tannin content of the herb, excessive use of burnet should be avoided.
References

See also General References G7 G36 G37 G40 G64.


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Species (Family)

Synonym(s)
Part(s) Used

Aerial parts, leaf, rhizome, root
Pharmacopoeial and Other Monographs

German Commission E (root and leaf).\(^{(G3)}\)
Legal Category (Licensed Products)

Butterbur is not on the GSL. There are no licensed products containing butterbur available in the UK.
Constituents

See General reference G2

Two chemovars of *P. hybridus* have been described, which differ markedly in the profile of sesquiterpenes present in the root/rhizome and leaf. The furanopetasin chemovar contains furanoeremophilane sesquiterpenes, whereas the petasin chemovar contains esters of the eremophilane sesquiterpenes petasol, isopetasol and neopetasol.\(^{(1-3)}\) The furanopetasin chemovar contains exclusively furanoeremophilane sesquiterpenes: petasin sesquiterpenes are not found in this chemovar.\(^{(2,4)}\) Eremophilanlactones have been reported as constituents of the furanopetasin chemovar,\(^{(1)}\) but it has been shown that they are produced by rearrangement of 9-hydroxy-furanoeremophilanes during storage of raw plant material.\(^{(5)}\)

Both chemotypes contain unsaturated pyrrolizidine alkaloids,\(^{(6)}\) and these compounds occur throughout all parts of the plant.\(^{(G19)}\)

Leaf

**Alkaloids**

Pyrrolizidine type, mainly senecionine and integerrimine.\(^{(7)}\) Content may vary from 0.02 to 1.50 parts per million (ppm),\(^{(8)}\) or <0.1 to 3.9 or 27.7 ppm (for leaves and leafstalks, respectively).\(^{(7)}\)

**Flavonoids**

Astragalin, isoquercitrin, quercetin.\(^{(G2)}\)

**Sesquiterpenes**

Furanopetasin chemovar: furanoeremophilane type, including 9-hydroxy-furanoeremophilanes and 9-oxo-furanoeremophilanes.\(^{(2)}\) Petasin chemovar: eremophilane type, mainly petasin and neopetasin (2.8 and 1.9 mg/g of dried plant material, respectively), also isopetasin, and the sulfur-containing compounds neo-S-petasin, S-petasin and iso-S-petasin (= S-isopetasin; trace).\(^{(3)}\)

**Other constituents**

Tannins, mucilage, volatile oil (about 0.1%), triterpenoid saponins (traces).\(^{(G2)}\)

Rhizome, root

**Alkaloids**
Pyrrolizidine type, mainly senecionine, its isomer integerrimine and senkirkine.\(^9\) Concentration can vary from 2 to 500 ppm.\(^{10}\) Values reported following specific investigations include 5–90 ppm\(^8\) and 6–105 (mean 49.2) ppm (calculated as senecionine);\(^7\) concentrations of senecionine and senkirkine have been reported as 11.3 and 25.8 ppm, respectively.\(^{11}\)

**Sesquiterpenes**

Furanoeremophilane type,\(^1\) mainly furanopetasin and 9-hydroxy-furanoeremophilane (approximately 16–21\% and 30–37\%, respectively),\(^2\) as well as furanoeremophilane, 2-senecioyl-furanopetasol, 2-tigloyl-furanopetasol and 2-methylthioacryloyl-furanopetasol (6–10\%).\(^2\) Petasin chemovar: eremophilane type, mainly petasin (9.4, 7.0 and 7.0 mg/g of dried plant material for rhizomes, roots and runners, respectively), also isopetasin, neopetasin, iso-S-petasin, neo-S-petasin and S-petasin.\(^3\) Minor compounds include 3-methylcrotonyl- and methacryloyl-esters of petasol, neopetasol, isopetasol, 3-desoxyneopetasol and 3-desoxyisopetasol.\(^{12}\)

**Volatile oil**

A mixture of at least 26 compounds of which 1-nonene, 1-undecene, albene, furanoeremophilane and fukinanolide are major components; minor compounds include the novel sesquiterpene hydrocarbons petasitene and pethybrene.\(^{13}\)

**Quality of crude material and marketed products**

As with other medicinal plants, the profile of constituents in different parts of *P. hybridus* can vary both qualitatively and quantitatively depending on various factors, and this has implications for safety and efficacy. The petasin content of the underground parts of plants (petasin chemovar) grown in four different locations in Switzerland ranged from 6.6 to 13.8 mg/g across regions.\(^{14}\) Furthermore, although plants showed an approximately similar relative distribution of the six main petasin sesquerpenes (petasin, isopetasin, neopetasin, iso-S-petasin, neo-S-petasin and S-petasin), there were exceptions: for example, in plants grown in the Krauchthal region, the content of S-petasin was approximately half that of neo-S-petasin, whereas in plants grown in the Chessiloch region, S-petasin content was greater than that of neo-S-petasin. Raised temperatures and storage conditions also affect the profile of constituents in *P. hybridus* (petasin chemovar) roots. Under moderate drying conditions (40°C in the dark), isomerisation occurs: sesquiterpene esters with petasol and neopetasol skeletons form the corresponding esters with isopetasin skeletons.\(^{15}\) The isomerisation of neopetasin to petasin occurs more readily than that of petasin to isopetasin (half-life for neopetasin and petasin: 49 and 205 days, respectively). In the
furanopetasin chemovar, 9-hydroxy-furanoeremophilanes rearrange to the corresponding eremophilanlactones during drying and storage of raw plant material.\(^5\)

Some manufacturers of herbal medicinal products containing butterbur state that the unsaturated pyrrolizidine alkaloids are removed from the product before it is released for sale.\(^{16}\) One method used to achieve acceptably low concentrations (the German Commission E advised that the maximum daily intake is 1 μg)\(^{11}\) of these alkaloids is by multiple applications of a method involving a cation-exchange resin;\(^{17}\) other methods of separating pyrrolizidine alkaloids from several plants, including \textit{P. hybridus}, containing these compounds have been described.\(^{11}\) A different approach to achieving extremely low concentrations of pyrrolizidine alkaloids is to cultivate clones of wild-type plants previously selected for almost absence of these constituents. Achievement of an initial step – \textit{in vitro} propagation of wild-type plants – in this approach has been documented.\(^{18}\)
Food Use

Butterbur is not used in foods.
Herbal Use

Butterbur root has been used traditionally for the treatment of fevers, wheezing and colds and as a diuretic and vermifuge.\(^{19,G34}\) It is also reputed to be a heart stimulant, cardiac tonic and antispasmodic, and to promote menstruation. It has been used externally for treating persistent sores and other skin problems.

Butterbur root was approved by the German Commission E for use as supportive therapy for acute spastic pain in the urinary tract, particularly if urinary stones were present.\(^{G3}\) Modern-day use of butterbur and clinical trials of products containing butterbur have focused on its use in seasonal allergic rhinitis and migraine prophylaxis.
The German Commission E advised that the dose of butterbur should be such that the daily intake of unsaturated pyrrolizidine alkaloids, including their $N$-oxides, is not greater than 1 μg.\(^{(G3)}\) The duration of use of butterbur should not exceed a total of 4–6 weeks in a year.\(^{(G3)}\)

**As an antispasmodic and for other traditional uses**

Dosages described in the herbal literature are typically preparations equivalent to 4.5–7 g dried butterbur root daily in the form of an ethanolic or lipophilic extract.\(^{(G3, G69)}\)

Dosages of butterbur extract used in clinical trials have varied. In trials involving adults with seasonal allergic rhinitis, one study used a butterbur root extract (containing not less than 15% petasins calculated as isopetasin; Petaforce) (D Martin, Bioforce (UK) Ltd, personal communication, 10 June 2004) 50 mg twice daily for two weeks,\(^{(20)}\) whereas another study tested a butterbur leaf extract (ZE-339) one tablet four times daily (equivalent to 32 mg petasins daily) for two weeks.\(^{(21)}\) Trials of butterbur in migraine prophylaxis have tested a carbon dioxide extract of butterbur rhizome (Petadolex) at doses of 50 or 75 mg twice daily (equivalent to at least 15 or 22.5 mg petasins daily, respectively) for 12 or 16 weeks.\(^{(22, 23)}\)
Pharmacological Actions

Several pharmacological activities, including anti-inflammatory, anti-allergic and antispasmodic effects, have been described for butterbur root/rhizome and leaf following preclinical studies. Extracts of butterbur have been tested clinically in patients with seasonal and perennial allergic rhinitis (hayfever), bronchial asthma and chronic obstructive bronchitis, and in prophylaxis of migraine headaches. The precise chemical constituents responsible for the documented effects of butterbur are not clearly understood, although several preclinical studies have shown that the petasin constituents, such as petasin and isopetasin, are important for certain pharmacological activities. These compounds may have several intracellular targets, and there is some evidence that the stereoisomers may have different affinities at these, even bringing about different effects.$^{(24)}$

However, other in vitro studies have reported that petasin is inactive,$^{(25)}$ and activities for the furanoeremophilane and eremophilanlactone constituents have been documented.$^{(26)}$

Published papers and texts often use the words root and rhizome interchangeably, although this is not strictly correct. This monograph uses terms exactly as published in the reference material, since it is not possible to say otherwise with certainty precisely which plant part was meant. Published reports also often fail to state which chemovar of P. hybridus was used.

In vitro and animal studies

Pharmacokinetics

Petasins have been shown to be bioavailable following studies in rabbits fed butterbur leaf extract (containing 50 mg petasin) twice daily for seven days; petasins were detected by gas chromatography-mass spectrometry (GC-MS) within the low nanogram range.$^{(27)}$

Anti-allergic and anti-inflammatory properties

Inhibition of peptido-leukotriene (LT) biosynthesis in murine peritoneal macrophages has been documented following in vitro experiments using several different extracts of butterbur root and leaf.$^{(25,28)}$ A petroleum ether extract of P. hybridus ground root showed the greatest inhibitory activity of LT synthesis from mouse peritoneal macrophages stimulated with calcium ionophore A23187 ($10^{-6}$ mol/L): 12.6, 95.1 and 100% inhibition at concentrations of P. hybridus extract of 6.3, 31.0 and 94.0 μg/mL, respectively ($p < 0.01$ versus control).$^{(25)}$ By contrast, a pyrrolizidine-alkaloid-free extract showed no activity in this assay, and the addition of the
Pyrrolizidine alkaloids senkirkine and senecionine to this extract and to the macrophage-containing tissue culture system also failed to result in inhibition of LT synthesis. A report of this work does not describe the constituents of the pyrrolizidine alkaloid-free extract, so the latter results for lack of activity are difficult to interpret.

Further work involving fractions of the active P. hybridus root extract indicated that isopetasin and the oxopetasan esters (= eremophilalanlactones, the presence of these is due to rearrangement of 9-hydroxy-furanoeremophilanes during drying and storage as stated above) were important for activity: the content of isopetasin and oxopetasan esters in the fractions correlated with inhibition of LT synthesis in a concentration-dependent manner. By contrast, petasin was not important for activity, but appeared to reduce the activity of isopetasin in this assay.

Other in vitro studies, however, have reported that petasin is one of the major active compounds of P. hybridus extract. Isolated petasin and a standardised high-pressure carbon dioxide extract of P. hybridus (containing 14.1% petasin; ZE-339, Zeller) inhibited the synthesis of cysteinyl-LTs in eosinophils and LTB₄ in neutrophils which had been primed with granulocyte-macrophage colony-stimulating factor and subsequently stimulated with platelet-activating factor (PAF) or the anaphylatoxin C5a. Petasin, ZE-339 and the positive control zileuton (a 5-lipoxygenase inhibitor) achieved similar inhibition of cysteinyl-LTs and LTB₄ in eosinophils and neutrophils, respectively, in these models (IC₅₀ (concentration for 50% inhibition) ≤ 24 μg/mL in each case).

In other in vitro studies using a similar model of eosinophil stimulation, ZE-339 showed greater inhibition of cysteinyl-LT synthesis than did a fraction containing petasin, isopetasin and neopetasin. When each of these latter compounds was tested separately in this model, all inhibited C5a- and PAF-induced cysteinyl-LT synthesis in a concentration-dependent manner and to the same extent as zileuton (positive control). However, the release of eosinophil cationic protein (ECP) from eosinophils (a measure of eosinophil degranulation) was inhibited by petasin, but not by isopetasin, neopetasin or zileuton. Similarly, petasin, but not isopetasin, neopetasin or zileuton, suppressed cytosolic phospholipase A₂ activity, 5-lipoxygenase translocation and PAF- and C5a-induced increases in intracellular calcium ion concentrations. These findings suggest that the different petasin constituents of butterbur may act upon different intracellular signalling molecules.

Gastroprotective effects
A petroleum ether extract of *P. hybridus* ground root protects against gastrointestinal damage induced by ethanol and indometacin (indomethacin) *in vivo*.\(^{28}\) *P. hybridus* root extract 0.83, 2.5 and 7.5 mg/kg administered intragastrically to fasted rats 30 minutes before intragastral administration of 1.5 mL ethanol significantly reduced gastric mucosal damage in a dose-dependent manner, compared with control (*p* < 0.05). In an experiment involving normally fed rats, indometacin (8 mg/kg orally) was administered with *P. hybridus* root extract (6.3, 12.5 or 25 mg/kg orally), cimetidine (50 mg/kg orally) or control, followed by a second dose 6 hours later, and sacrifice after a further 16 hours. Compared with control, all doses of *P. hybridus* root extract significantly inhibited indometacin-induced intestinal damage (*p* < 0.05), and *P. hybridus* root extract 2 × 25 mg/kg inhibited intestinal damage to a greater extent than did cimetidine (*p* < 0.05).\(^{28}\)

**Other activities**

Extracts of butterbur rhizome from both petasin and furanopetasin chemovars of *P. hybridus* have been reported to inhibit the binding of radioactive ligands at dopamine D\(_2\) and histamine H\(_1\) receptors to a similar extent. The greatest inhibition was observed for extracts with the highest content (>60%) of eremophilane constituents. For example, for a rhizome extract from the petasin chemovar, mean (standard error of mean; SEM) IC\(_{50}\) values for inhibition of dopamine D\(_2\) and histamine H\(_1\) radioligand binding were 38 (3) and 155 (52), respectively.\(^{26}\) Fractions of the petasin chemovar extract inhibited radioligand binding in a concentration-dependent manner, with the highest affinities displayed by the petasin and desoxyneopetasol fractions. This was confirmed in assays with individual constituents in which the lowest IC\(_{50}\) values were determined for petasin and isopetasin. The furanopetasin chemovar constituents hydroxy-furanoeremophilane and furanopetasin also inhibited radioligand binding, as did the eremophilanlactones eremophilanolide and hydroxy-eremophilanolide, albeit with markedly lower affinities than the eremophilanes.\(^{26}\)

Petasin, isopetasin, S-petasin and iso-S-petasin, isolated from *Petasites formosanus*, a species related to *P. hybridus*, relaxed histamine-, carbachol-, potassium chloride- and LTD\(_4\)-induced precontractions of isolated guinea-pig trachea in a concentration-dependent manner.\(^{31}\) S-Petasin was more potent than petasin and isopetasin for precontractions induced by all four reagents. S-Petasin relaxed the precontractions induced by all four reagents in a non-selective manner, with IC\(_{50}\) values of <10 μmol/L, whereas iso-S-petasin selectively relaxed precontractions induced by carbachol and potassium chloride, with IC\(_{50}\) values of around 10 μmol/L for each. The reason for these
differences in the respective profiles of activity, and the influence of isomerisation (S-petasin to iso-S-petasin) is not clear. The documented relaxant effects of S-petasin and iso-S-petasin from *P. formosanus* on precontracted guinea-pig trachea may be due to antispasmodic and antimuscarinic properties, respectively.\(^{32}\) Following a study using isolated guinea-pig atria, it has been reported that iso-S-petasin may act preferentially on tracheal muscarinic M\(_3\) receptors, rather than cardiac M\(_2\) receptors.\(^{33}\)

Hypotensive activity has been documented for S-petasin and iso-S-petasin isolated from *P. formosanus*. In rats, a dose-dependent hypotensive effect occurred following intravenous administration of S-petasin or iso-S-petasin (0.1–1.5 mg/kg body weight for each); *in vitro* experiments indicated that these compounds have a relaxant effect on precontracted rat aortic ring segments and that this may be due in part to blockade of calcium ion (Ca\(^{2+}\)) channels in vascular smooth muscle cells by S-petasin and iso-S-petasin.\(^{34,35}\)

S-Petasin from *P. formosanus* has been shown to have negative chronotropic activity *in vivo*. In anaesthetised rats, S-petasin (1.0–1.5 mg/kg body weight, intravenously) induced bradycardia in a dose-dependent manner within a few seconds of administration and the effect persisted for up to 11 minutes after administration.\(^{36}\) The highest administered dose of S-petasin (1.5 mg/kg body weight intravenously) evoked a maximal reduction in heart rate of approximately 25%. S-Petasin had a negative inotropic effect in isolated rat atria and depressed the amplitude of contraction of rat cardiac myocytes. Iso-S-petasin also depresses cardiac contraction as demonstrated by *in vitro* studies involving ventricular myocytes.\(^{37}\) The mechanism for the observed negative cardiac chronotropic and inotropic effects of S-petasin and iso-S-petasin may be through inhibition of cardiac L-type voltage-dependent Ca\(^{2+}\) channels.\(^{36,37}\) Further work has shown that S-petasin decreases the amplitude of L-type Ca\(^{2+}\) currents in NG108-15 cells (a mouse neuroblastoma and rat glioma hybrid cell line) in a concentration-dependent manner (IC\(_{50}\) = 11 μmol/L).\(^{38}\)

**Clinical studies**

Several clinical trials of butterbur extracts have been conducted and have involved individuals with, for example, seasonal and perennial allergic rhinitis and migraine. While many of these studies have reported beneficial effects with butterbur, rigorous investigation of the efficacy of butterbur extracts is limited and further large, well-designed clinical trials are required.
Pharmacokinetics

There are limited pharmacokinetic data for the constituents of butterbur extracts. The bioavailability of petasin has been reported following a study involving healthy volunteers who received a single dose of two ($n = 24$) or four tablets ($n = 24$) of butterbur leaf extract (ZE-339) orally; each tablet contained 8 mg petasins. The time to maximal petasin concentrations ($t_{\text{max}}$) was around 1.6 hours in each group (mean (standard deviation, SD): 1.62 (0.50) and 1.61 (0.93) hours for the lower and higher dose groups, respectively), whereas the mean (SD) half-life was 7.16 (4.61) and 7.62 (3.34) hours for the lower and higher dose group, respectively. Maximal mean (SD) petasin concentrations were 25.5 (14.8) and 58.1 (26.7) ng/mL for the lower and higher dose groups, respectively.

Another study described monitoring compliance by measuring serum petasin concentrations using an enzyme-linked immunosorbent assay, but these results were not reported.

Therapeutic effects

**Seasonal and perennial allergic rhinitis** In a randomised, double-blind, double-dummy, parallel-group study involving 125 individuals with seasonal allergic rhinitis, the effects of a carbon dioxide extract of butterbur leaf (ZE-339) one tablet four times daily (equivalent to 32 mg petasins daily) were compared with those of the non-sedating antihistamine cetirizine 10 mg each evening. At the end of the two-week study, butterbur recipients ($n = 61$) achieved similar scores to cetirizine recipients on the SF-36 (Short Form 36; a self-assessment scale for medical and health outcomes), the primary outcome measure. There were also no differences between groups with respect to secondary outcome measures, including the clinical global impression score. The overall frequency of adverse events was similar in both groups (see Side-effects, Toxicity).

The statistical power of the study only allowed the conclusion to be drawn that butterbur was not inferior to cetirizine (i.e. the study does not demonstrate equivalence of the two preparations), and the study has been criticised for its choice of subjective outcome measures and interpretation.

A subsequent study used an objective outcome measure – the adenosine
AMP monophosphate (AMP) nasal provocation test (AMP is important in the pathway leading to the release of allergic inflammatory mediators, such as histamine, cysteinyl leukotrienes and prostaglandins) – to assess the effects of butterbur. In a randomised, double-blind, crossover trial, 20 individuals with seasonal allergic rhinitis ceased any existing treatment for the condition (antihistamines and/or intranasal corticosteroids) one week before starting treatment with butterbur 50 mg (Petaforce; Bioforce), or placebo, twice daily for two weeks.\(^{(20)}\) No further details of the preparation were provided in the report of the study, although Petaforce marketed in the UK contains butterbur root extract 25 mg per capsule (containing not less than 15% petasins calculated as isopetasin).

At the end of each treatment period, participants underwent AMP challenge (described as two applications to each nostril of a 400 mg/mL solution delivered via a pump actuator spray device) and spontaneous recovery was monitored by measuring peak nasal inspiratory flow (PNIF) at regular intervals over 1 hour. Time to recovery was significantly attenuated and the maximum fall in PNIF was significantly less in butterbur recipients compared with placebo recipients \( (p = 0.028 \text{ and } 0.036, \text{ respectively}) \).\(^{(20)}\) The study design did not incorporate a wash-out period between the two phases of the study, and it is not clear whether there could have been any carry-over effect in patients who received butterbur during the first phase of the study and what influence, if any, this may have had on the results.

The effects of butterbur extract (Petaforce) 50 mg twice daily in perennial allergic rhinitis were compared with those of fexofenadine 180 mg daily in a randomised, double-blind, placebo-controlled, crossover study involving 16 individuals. Participants stopped their existing treatment for allergic rhinitis one week before starting their randomised treatment; treatments were taken for one week, with a one-week wash-out period between each randomised treatment. At the end of the study, the maximum percentage fall from baseline values in PNIF after nasal AMP challenge was significantly attenuated in the butterbur and fexofenadine groups, compared with the placebo group (mean (SEM) for butterbur, fexofenadine and placebo: 34 (3), 39 (3) and 46 (3), respectively; \( p < 0.05 \)).\(^{(43)}\) The total nasal symptom score, a secondary outcome measure, was also significantly improved for the butterbur and fexofenadine groups, compared with the placebo group \( (p < 0.05) \). This study is limited in that the duration of treatment was short, and the sample size calculation was not based on detecting differences in nasal symptom scores, so a larger study is needed to confirm the latter result.

In a small, uncontrolled, open study, six patients with seasonal \( (n = 4) \) or
perennial (n = 2) allergic rhinitis received butterbur extract (ZE-339) two tablets three times daily (equivalent to 48 mg petasins daily) for five days and underwent a series of tests each day 90 minutes after taking their first daily dose. At the end of the study, nasal symptom scores and self-rated quality-of-life scores had improved significantly, compared with baseline values. Significant reductions in the concentrations of some, but not all, inflammatory mediators measured were also noted (e.g. a decrease in \( \text{LTB}_4 \) concentration in nasal lavage fluids) after five days, compared with baseline values. However, because of the methodological limitations of the study (e.g. small sample of patients, no control group, no masking of treatment) these changes cannot be attributed to treatment with butterbur extract.

**Asthma and Bronchitis**

At present, there is insufficient evidence from well-designed randomised controlled trials to support the efficacy of butterbur extracts in asthma and bronchitis.

In a randomised, double-blind, crossover study, 16 patients with atopic asthma who had been stabilised on inhaled corticosteroid therapy for at least three months before the study received butterbur (Petaforce) 25 mg twice daily (no further details stated in the report although Petaforce marketed in the UK contains not less than 15% petasins calculated as isopetasin), or placebo, for one week. Participants stopped any treatment with long-acting \( \beta_2 \)-agonists one week before and for the duration of the study. At the end of the study, bronchial hyper-responsiveness in response to AMP bronchial challenge was significantly improved in butterbur recipients compared with placebo recipients (\( p < 0.05 \)), and concentrations of inflammatory markers (exhaled nitric oxide, serum eosinophil cationic protein and peripheral blood eosinophil count) were significantly suppressed in the butterbur group, compared with the placebo group (\( p < 0.05 \) for each).

In a preliminary trial, 80 individuals (aged 6–85 years) with mild or moderate asthma received a lipophilic carbon dioxide extract of butterbur rhizome (Petadolex; standardised to contain at least 15% petasins) 50 mg three times daily (equivalent to 22.5 mg petasins daily; dose reduced for children depending on age) for a minimum of two months following a two-week run-in phase. A report of this study attributes several improvements to administration of butterbur, including reductions in the number, duration and severity of asthma attacks, increases in peak flow and forced expiratory volume in one second (\( \text{FEV}_1 \)), and reduced use of existing asthma medications. However, such conclusions may be flawed because of the design and methodological limitations of the study and because no statistical analysis was undertaken.
**Allergic Skin Reactions** A randomised, double-blind, double-dummy, controlled, crossover skin trial assessed the effects of butterbur on the histamine and allergen cutaneous response in 20 atopic patients with asthma or allergic rhinitis and sensitisation to at least one common household allergen, such as house dust mite, on skin prick testing.\(^{(46)}\) Participants received butterbur (Petaforce) 50 mg twice daily (no further details given), fexofenadine 180 mg daily, montelukast 10 mg daily, or placebo, for one week; existing treatment with antihistamines and leukotriene receptor antagonists was stopped one week before and for the duration of the study, although existing treatment for asthma or allergic rhinitis was continued. Each day, approximately 2 hours after taking the first daily dose of study medication, each participant underwent skin prick testing with the allergen that had previously been shown to provoke the greatest response in that individual, as well as with histamine (1.7 mg/mL; no further details reported) and 0.9% sodium chloride as control. Mean histamine and allergen wheal and flare responses were significantly attenuated by fexofenadine, compared with placebo, but not by butterbur or montelukast.\(^{(46)}\)

**Migraine Prophylaxis** The rationale for the use of butterbur in migraine prophylaxis is centred around the understanding that vasoconstrictive and neurogenic inflammatory processes are involved in the generation of migraine headaches, and that butterbur and certain of its isolated constituents have been shown in preclinical studies to have anti-inflammatory properties.\(^{(22)}\) However, at present, rigorous clinical investigation of the effects of butterbur extracts in preventing migraine is limited.

In a randomised, double-blind, parallel-group trial, 60 hospital outpatients with migraine (minimum of three attacks per month for the three months prior to the study and a minimum of two attacks in the four-week run-in phase) received a carbon dioxide extract of *P. hybridus* rhizome (Petadolex) two capsules twice daily (equivalent to 100 mg extract or 15 mg petasins daily), or placebo, for 12 weeks. None of the participants were previous users of the butterbur extract.\(^{(22)}\) Inclusion and exclusion criteria for the study were in accordance with the International Headache Society guidelines.\(^{(47)}\) The frequency of migraine attacks in the last four weeks of the study was significantly lower in the butterbur group, compared with the placebo group: the mean (SD) numbers of attacks during the month were 1.7 (0.9) and 2.6 (1.1), for the butterbur and placebo groups, respectively; \(p < 0.05\). The number of days with migraine during the last four weeks of the study also decreased significantly in the butterbur group, compared with the placebo group (mean (SD) number of days at baseline and weeks 8–12: 3.4 (1.6) and 1.7 (0.9) versus 3.0 (1.3) and 2.6 (1.2), for butterbur
and placebo, respectively; $p < 0.05$). There were no statistically significant differences between the two groups in the duration and intensity of migraine headaches at the end of the study.\(^{(22)}\)

The study has several limitations which should be considered when interpreting the results. A sample size calculation was not carried out, baseline characteristics (apart from age and variables relating to migraine attacks) of participants were not provided in a report of the study, so it is not clear if the randomisation was successful (i.e. whether or not the two groups were similar at baseline) and differences at baseline in the frequency of migraine attacks are not considered in the analysis. There are further flaws in the statistical analysis, for example, the analysis was carried out only with those participants who adhered to the protocol (i.e. an intention-to-treat analysis was not carried out), efficacy parameters were not defined a priori,\(^{(48)}\) the results are reported without 95% confidence intervals and precise $p$-values are not provided.

Against this background, an independent re-analysis of the data from this study was undertaken. Data entry from original case report forms was completely repeated under the principles of good clinical practice (which includes double data entry and consistency checks), and all four primary efficacy criteria and data from all three time points (4, 8 and 12 weeks) were evaluated equally weighted in an attempt to avoid bias by post-hoc selections of efficacy criteria.\(^{(48)}\) All analyses were undertaken with data from the intention-to-treat population (i.e. all patients who were randomised and took the study medication at least once).

The new analysis confirmed the findings of the original analysis and stated that all 12 primary efficacy criteria (number of attacks, number of days with attacks, mean duration and mean intensity of attacks at all three time points) were significantly reduced in the butterbur group, compared with the placebo group, and the most conservative analysis showed that 7 of these 12 variables were still statistically significant.\(^{(48)}\) However, the possibility of bias in this retrospective re-analysis cannot be excluded entirely, and other methodological issues remain, such as the lack of a sample size calculation.

A larger randomised, double-blind, placebo-controlled trial of a butterbur extract for migraine prophylaxis has been conducted but at present has been reported only as a conference abstract so full details are not available and it is not possible to judge the methodological quality of the study. In the trial, 245 patients with migraine (minimum of three attacks per month for the three months prior to the study and a minimum of two attacks in
the four-week run-in phase) received an extract of butterbur rhizome (Petadolex) 50 mg twice daily, 75 mg twice daily, or placebo, for 16 weeks.\(^{(23)}\) In total, 202 participants completed the study and were included in the analysis; it is not clear whether this was the per-protocol or the intention-to-treat population. At the end of the study, the mean number of migraine attacks was reduced by 48%, 34% and 26%, compared with baseline values, for the butterbur extract 150 mg daily, 100 mg daily and placebo groups, respectively (\(p < 0.001\) for butterbur extract 150 mg daily versus placebo).
Side-effects, Toxicity

As with many herbal medicines, the safety of butterbur preparations has not yet been adequately assessed: only limited preclinical and clinical safety and toxicity data are available (see below). One of the main issues regarding the clinical use of butterbur extracts concerns the unsaturated pyrrolizidine alkaloid constituents which are known to be hepatotoxic in humans, and have been shown to be carcinogenic and mutagenic in preclinical studies (see Comfrey). These alkaloids may be present in low concentrations in all parts of the plant (see Constituents).

Several manufacturers of products containing butterbur include in their manufacturing processes steps to remove the unsaturated pyrrolizidine alkaloids to concentrations below the detectable limit (e.g. <35 parts per billion\(^{49}\)). However, there remains the possibility that toxic quantities of unsaturated pyrrolizidine alkaloids may be present in poorly processed products, and any products which contain *Petasites* species as a result of botanical misidentification or adulteration.

Butterbur is a member of the Asteraceae family, and members of this family may cause allergic reactions in sensitive individuals, especially those with an existing hypersensitivity to other members of the Asteraceae/Compositae. No reports of allergic reactions to butterbur were identified.

Clinical safety data

Swissmedic, the competent authority for licensing medicines in Switzerland, has revoked the licences for certain products containing butterbur root extract following spontaneous reports received in Germany of liver damage associated with their use.\(^{50}\) At the time of writing, the German authority had not taken any regulatory action, but was monitoring the situation. Further details of the reports are not yet available.

There is only limited information available for butterbur from long-term postmarketing surveillance studies, and clinical trials reported to date generally have involved only small numbers of participants, mostly with allergic rhinitis or migraine who are generally otherwise healthy, and have involved ingestion of butterbur preparations for relatively short periods of time (up to four months). Some clinical trials have not adequately reported data on safety, and two randomised, double-blind, placebo-controlled trials involving individuals with seasonal allergic rhinitis or asthma who received butterbur extracts and in which participants continued to take their existing medication (inhaled or intranasal corticosteroids, long-or short-acting \(\beta_2\)-
agonists\(^{(44,46)}\) have either not assessed or not reported data on safety parameters at all.

In a randomised, double-blind, trial involving 125 individuals with seasonal allergic rhinitis who received butterbur extract (ZE-339) equivalent to 32 mg petasins daily, or cetirizine 10 mg daily, for two weeks, the frequency of adverse events was similar in both groups (proportions of butterbur and cetirizine recipients who reported adverse events were 10% and 11%, respectively).\(^{(21)}\) Eight of the 12 adverse events reported by cetirizine recipients and 2 of the 10 adverse events reported by butterbur recipients were classified as fatigue or drowsiness. Raised liver enzyme activity (no further details provided) and pruritus were reported for one butterbur recipient each, but not for cetirizine recipients.

In a randomised, double-blind, placebo-controlled, crossover study, 20 patients with seasonal allergic rhinitis stopped taking any existing treatment one week before receiving butterbur (Petaforce) 50 mg twice daily for two weeks (no further details of the preparation were provided in a report of the study, although Petaforce marketed in the UK contains not less than 15% petasins calculated as isopetasin). At the end of the study, liver function test values (blood concentrations of alanine aminotransferase (ALT), bilirubin, alkaline phosphatase (ALP) and albumin) were reported to be similar for butterbur and placebo groups,\(^{(20)}\) although no statistical analysis was carried out. However, this study involved only small numbers of participants and was conducted over a short time period and, therefore, cannot provide adequate data on safety.

Open, uncontrolled studies of butterbur extracts (ZE-339 and Petadolex) in patients with seasonal and/or perennial allergic rhinitis or asthma have reported that butterbur was well tolerated,\(^{(39,45)}\) although the design of these studies renders them unsuitable for an adequate assessment of safety. In one of these studies, in which participants (aged 6–85 years) with mild or moderate asthma received a lipophilic carbon dioxide extract of butterbur rhizome (Petadolex; standardised to contain at least 15% petasins) 50 mg three times daily (reduced for children depending on age) for a minimum of two months, seven participants reported 11 adverse events.\(^{(45)}\) These included abdominal pain, flatulence, sneezing, allergic conjunctivitis, allergic rhinitis and halitosis (reported by children) and hair loss, coughing, dyspnoea, difficulty exhaling and depression (reported by adults). None of these was judged by the study physician to be causally related to butterbur ingestion and none led to withdrawal of participants from the study.

In a randomised, double-blind, placebo-controlled trial involving 60
individuals with migraine who received a carbon dioxide extract of *P. hybridus* rhizome (Petadolex) two capsules twice daily (equivalent to 100 mg extract daily), or placebo, for 12 weeks, no adverse events were reported for the butterbur group and no statistically significant changes from baseline values were reported for vital signs and physical examination results.\(^{(22)}\) However, two butterbur recipients withdrew from the study, one because of a suspected pregnancy; the other participant did not provide a reason. An independent re-analysis of the data from this trial added that, at the end of the study, three butterbur recipients had liver function test (ALT, aspartate transaminase (AST)) values which were higher than normal ranges and significantly higher than baseline values, and that bilirubin concentration and erythrocyte count were significantly higher than baseline values for the butterbur group, compared with the placebo group.\(^{(48)}\) These changes were not regarded as being clinically relevant,\(^{(48)}\) although no numerical data were provided to support this judgement.

Only limited data relating to safety are available from a larger randomised, double-blind, placebo-controlled trial involving 245 patients with migraine who received an extract of butterbur rhizome (Petadolex) 50 mg twice daily, 75 mg twice daily, or placebo, for 16 weeks.\(^{(23)}\) Data from 230 participants were included in a safety analysis. The most frequently reported adverse events that were considered possibly related to treatment were gastrointestinal symptoms (not specified) which occurred in 24, 30 and 8% of participants in the butterbur extract 75 mg, butterbur extract 50 mg and placebo groups, respectively.\(^{(23)}\)

A review of safety data for a specific preparation of butterbur rhizome – an extract (Petadolex) standardised to contain a minimum of 15% petasins and processed to achieve a concentration of unsaturated pyrrolizidine alkaloids of less than 0.08 ppm – includes data from four postmarketing surveillance studies involving a total of 188 patients (145 with migraine, including 50 children and adolescents aged 6–17 years). Adverse events deemed to be possibly or probably causally related to ingestion of the butterbur product included eructations (belching; \(n = 7\)), bad taste/smell of the product (2) and skin rash (1).\(^{(51)}\) However, it is unclear why these four studies included such a small number of participants: a single postmarketing surveillance study would normally be expected to include many more participants.

The review also refers to 93 spontaneous suspected adverse drug reaction (ADR) reports (75 of which originated from Germany, where the manufacturer is located) received by the product manufacturer from 1976 to the end of June 2002. In total, 27 of these reports were determined to be possibly \((n = 19)\) or probably \((8)\) causally related to butterbur administration;
the latter included one case of reversible cholestatic hepatitis.\(^{(51)}\) The review states an overall frequency of suspected ADRs, calculated on the basis of sales figures and the total number of spontaneous ADR reports received by the manufacturer. However, for several reasons, these types of data should not be used to calculate frequencies of suspected ADRs.

**Preclinical safety data**

There is limited data from animal toxicity studies for butterbur preparations, although a review\(^{(51)}\) summarises data from unpublished toxicity studies. Acute toxicity studies in rats yielded LD\(_{50}\) values for a single-dose oral administration and single-dose intraperitoneal administration of ≥ 2.5 and approximately 1 g/kg body weight, respectively. A chronic toxicity study in rats (\(n = 200\)) carried out over 26 weeks established a no adverse effect level for the lower dose range tested (oral administration; no further details provided).\(^{(51)}\)

Information on mutagenicity and genotoxicity of butterbur extracts is limited to summaries of unpublished data. An extract of butterbur rhizome (Petadolex) produced a negative result in the Ames test for mutagenicity using several strains of *Salmonella typhimurium*.\(^{(52)}\) In an *in vitro* test for clastogenic activity in which the butterbur extract was incubated with cultured human peripheral lymphocytes, the mean number of chromosomal aberrations was within the reference range of the negative control.\(^{(53)}\) This result was obtained irrespective of whether or not the test included metabolic activation using a rat liver postmitochondrial fraction. In contrast, mitomycin C and cyclophosphamide, as positive controls, induced chromosomal damage.

There is some evidence from preclinical studies that S-petasin, a constituent of butterbur root, rhizome and leaf, has effects on certain endocrine systems. In rats, S-petasin (10 \(\mu\)g/kg body weight, intravenously) reduced basal plasma corticosterone concentrations to a significantly greater extent than did control at 30 minutes after administration (\(p < 0.05\)), although there were no statistically significant differences between groups when corticosterone concentrations were measured at 1, 2 and 3 hours after administration.\(^{(54)}\) The same dose of S-petasin also significantly reduced adrenocorticotropic hormone (ACTH)-induced increases in plasma corticosterone concentrations at 30 minutes, but not longer intervals, after administration, compared with control. Similar effects were observed following *in vitro* experiments: S-petasin significantly reduced basal, ACTH- and forskolin (an adenylyl cyclase activator)-stimulated corticosterone release from rat adrenal gland cells (zona fasciculata reticularis) in a concentration-dependent manner. Results of further *in vitro* experiments suggested that the mechanism for the observed
Effects is in part through inhibition by S-petasin of the enzymes adenylyl cyclase (which catalyses the formation of cyclic AMP), and cytochrome P450 side-chain cleavage and 11β-hydroxylase, which are important in the biosynthesis of corticosterone.\(^{54,55}\)

S-Petasin (1 μg/kg body weight, intravenously) administered as a single dose to small numbers of adult male rats reduced basal plasma testosterone concentrations, compared with control (mean (SEM): 0.81 (0.06) and 1.31 (0.21) ng/mL for S-petasin and control, respectively; \(p < 0.05\)).\(^{56}\) Incubation of S-petasin at concentrations of 0.14–14.4 μg/mL with rat testicular interstitial cells led to a concentration-dependent inhibition of testosterone release. S-Petasin also inhibited forskolin-, human chorionic gonadotrophin- and androstenidione-induced stimulation of testosterone secretion from rat testicular interstitial cells \textit{in vitro}. 
Contra-indications, Warnings

In view of the known toxicity, the German Commission E recommended that the maximum daily intake of unsaturated pyrrolizidine alkaloids is 1 μg, and that the duration of use should not exceed 4–6 weeks in a year. In Switzerland, there are concerns regarding reports in Germany of hepatotoxicity associated with the use of certain products containing butterbur root extract (see Side-effects, Toxicity).

Drug interactions

No reports of drug interactions with butterbur extracts were identified. Certain constituents of butterbur have been documented in preclinical studies to have hypotensive activity and negative chronotropic and negative inotropic effects. Against this background, and on a theoretical basis, the possibility of interactions with hypertensive and antihypertensive medicines should be considered. Likewise, several constituents of butterbur have been documented to displace the binding of ligands at dopamine D₂ and histamine H₁ receptors, so there is a theoretical possibility of interactions with medicinal agents acting at these receptors.

Pregnancy and lactation

In view of the lack of safety data and the possible hepatotoxic effects of poorly processed butterbur extracts, the use of products containing butterbur is contraindicated in pregnancy and by breastfeeding women.
The chemistry of butterbur is well documented. There is evidence from preclinical studies to indicate that the eremophilane sesquiterpene constituents (e.g. petasin and isopetasin), found in the petasin chemovar, are important for activity, although there are some conflicting reports and it is not clear precisely which constituents are most active. Furthermore, certain activities have been documented for the furanoeremophilane sesquiterpenes found in the furanopetasin chemovar of \textit{P. hybridus}.

There is evidence from preclinical studies of the anti-inflammatory, anti-allergic and antispasmodic properties of butterbur extracts and/or their constituents, hence supporting some of the traditional uses. Precise mechanisms of action have not yet been elucidated, although inhibition of leukotriene synthesis, phospholipase A$_2$ activity and 5-lipoxygenase translocation, and effects on intracellular calcium ion concentrations have been documented in \textit{in vitro} studies. The implications of binding of certain butterbur constituents at dopamine D$_2$, histamine H$_1$ and muscarinic receptors, described following \textit{in vitro} studies, require further investigation.

There is a paucity of well-designed, randomised clinical trials of butterbur extracts. Existing trials generally have involved individuals with seasonal allergic rhinitis and migraine and have tested the effects of only a small number of herbal medicinal products containing butterbur extracts. While most of these studies have reported beneficial effects for butterbur extracts, methodologically rigorous investigation of efficacy in allergic rhinitis and migraine prophylaxis is limited and large, well-designed clinical trials are required. Similarly, at present, there is insufficient evidence from randomised clinical trials to support the use of butterbur extracts in patients with asthma and bronchitis.

There is also only limited information on the safety of butterbur extracts and further investigation of the safety of butterbur extracts is required. On the basis of small, randomised clinical trials, certain butterbur extracts appear to be well-tolerated when taken at recommended doses for relatively short periods (clinical studies have involved ingestion for up to four months); adverse events reported most frequently are gastrointestinal symptoms. Chronic toxicity studies are lacking, and there is a lack of clinical information on the safety of long-term use. Against this background, butterbur should not be used for long periods, nor at higher than recommended doses (see below). Spontaneous reports of liver damage associated with the use of a butterbur root extract have been received in Germany; no further details were available at the time of writing. On the basis of this information, the Swiss
regulatory authority for medicines revoked the licences for certain butterbur products marketed in Switzerland; regulatory action has not been taken in other countries.

No drug interactions have been reported for butterbur extracts. Based on data from preclinical studies, there is a theoretical possibility that butterbur extracts potentially could interact with hyper- and antihypertensive medicines, leukotriene receptor antagonists and agonists/antagonists at dopamine D₂, histamine H₁ and muscarinic M₃ receptors.

Both chemotypes (petasin and furanopetasin) of *P. hybridus* contain unsaturated pyrrolizidine alkaloids,⁶ and these compounds occur in low concentrations throughout all parts of the plant.⁴¹ These constituents are known to be hepatotoxic in humans, and have been shown to be carcinogenic and mutagenic in preclinical studies (see Comfrey). Several manufacturers of products containing butterbur include in their manufacturing processes steps to remove the unsaturated pyrrolizidine alkaloids to concentrations below the detectable limit (e.g. <35 parts per billion⁴⁹). However, there remains the possibility that toxic quantities of unsaturated pyrrolizidine alkaloids may be present in poorly processed products, and any products which contain *Petasites* species as a result of botanical misidentification or adulteration.

There is a view that butterbur is unsuitable for use as a tea or infusion.⁴⁶ This is because the *N*-oxides of unsaturated pyrrolizidine alkaloids, which can be formed from unsaturated pyrrolizidine alkaloids during storage of plant material but which can also be present in raw plant material, are more water soluble than the parent compounds and are, therefore, extracted more easily during preparations of herbal teas and infusions.⁴¹

The dose of butterbur should be such that the daily intake of unsaturated pyrrolizidine alkaloids, including their *N*-oxides, is not greater than 1 μg.⁴³ The duration of use of butterbur should not exceed a total of 4–6 weeks in a year.⁴³
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See also General References G2, G3, G19.

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Calamus
Species (Family)

*Acorus calamus* L. (Araceae)

Synonym(s)
Sweet Flag
Part(s) Used

Rhizome
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL^{G37}
Constituents

See General References G19 G22 G41 G58.

**Amines**
Dimethylamine, methylamine, trimethyl amine and choline.

**Volatile oil**
1.5–3.5%. β-Asarone content varies between genetic species: 96% in tetraploid (Indian), 5% in triploid (European) and 0% in the diploid (North American) species.\(^{(1-4)}\) Other identified components include calamenol (5%), calamene (4%), calamone (1%), methyl eugenol (1%), eugenol (0.3%) and the sesquiterpenes acolamone, acoragermacrone and isoacolamone. Considerable qualitative and quantitative differences have been reported between the volatile oil from different genetic species, and between the volatile fraction of an alcoholic extract and the essential oil from the same variety (European).\(^{(3,4)}\)

**Tannin**
1.5%.

**Other constituents**
Bitter principles (e.g. acorin), acoric and palmitic acids, resin (2.5%), mucilage, starch (25–40%), sugars.
Food Use

The level of β-asarone permitted in foods is restricted to 0.1 mg/kg in foods and beverages, 1 mg/kg in alcoholic beverages and in foods containing Acorus calamus or Asarum europaeum. Calamus is listed by the Council of Europe as a source of natural food flavouring (category N3). This category indicates that calamus can be added to foodstuffs in the traditionally accepted manner, although there is insufficient information available for an adequate assessment of potential toxicity. Calamus is classified as an ‘unsafe herb’ by the US Food and Drugs Administration (FDA), and the use of the rhizome and its derivatives (oil, extracts) are prohibited from use in human food.
Herbal Use

Calamus is stated to act as a carminative, spasmolytic and diaphoretic. Traditionally it has been indicated for acute and chronic dyspepsia, gastritis and gastric ulcer, intestinal colic and anorexia.\(^{(G7)}\)
Dosage

**Rhizome**
1–3 g or by infusion three times daily.\(^{(G7)}\)

**Liquid extract**
1–3 mL (1 : 1 in 60% alcohol) three times daily.\(^{(G7)}\)

**Tincture**
2–4 mL (1 : 5 in 60% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

**In vitro and animal studies**

Numerous documented studies have concentrated on activities associated with the oil. The pharmacology and toxicology of calamus oil have been reviewed.\(^{(5)}\) Unless specified, all of the following actions refer to those exhibited by the oil.

Spasmolytic action *in vitro* versus various spasmogens in different smooth muscle preparations including tracheal, intestinal, uterine, bronchial and vascular has been reported for European and Indian varieties.\(^{(5-8)}\) In one study activity was associated with a lack of β-asarone,\(^{(6)}\) whereas oils with either low or high levels of β-asarone have also exhibited activity.\(^{(5,7)}\) The pattern of spasmolytic activity has been compared to that of papaverine, and a direct musculotropic action has been proposed.\(^{(8)}\) Unlike papaverine an acetylcholine–like action has also been observed with low dilutions of the oil and asarone.\(^{(8)}\)

Inhibition of monoamine oxidase activity and a stimulation of D- and L-amino oxidase has been reported.\(^{(5)}\) The mechanism for this activity, involving serotonin and adrenaline, has been disputed, and an alternative mechanism involving depression of hypothalamic function has been proposed.\(^{(9)}\)

*In vitro* oil rich in β-asarone has been reported to reduce phenylbutazone–induced ulcers in the rat by 5–60%, although no effect was observed on stress- or ethanol-induced ulcers.\(^{(7)}\) No spasmolytic activity was reported for oil free from or with low levels of β-asarone.

A sedative action and a potentiation of barbiturate effect (increased sleeping time, reduction in body temperature) have been described in a number of small animals (mice, rats, rabbits and cats) following intravenous or intraperitoneal administration of European (alcoholic and aqueous extracts) and Indian varieties.\(^{(5)}\) Dexamfetamine has been found to block the potentiating action of the Indian variety on barbiturate sleeping time.\(^{(5)}\) Potentiation of morphine activity has been reported for the European variety.

The Indian oil has been reported to deplete levels of serotonin and noradrenaline in the rat brain following intraperitoneal administration.\(^{(5)}\) The mechanism of action was suggested as similar to that of reserpine, and a potentiation of the amfetamine–detoxifying effect of reserpine has also been described.\(^{(5)}\) In contrast, the central action of the European variety has been stated to not resemble that of reserpine.\(^{(5)}\) Anti-adrenergic activity
demonstrated by antagonism of dexamfetamine–induced agitational symptoms has been reported for the Indian variety in various small animals.\(^{(5)}\)

Anticonvulsant, anti–arrhythmic (like quinidine) and hypotensive (apparently not due to a nervous mechanism) activities in small animals have also been reported for the Indian variety.\(^{(5)}\)

α-Asarone, isolated from *Asarum europaeum* (Aristolochiaceae), has a local anaesthetic activity similar to that of benzocaine.\(^{(10)}\)

Antifungal activity has been documented for β-asarone\(^{(11)}\) and for the oil (weak).\(^{(5)}\) Insecticidal and leech repellant properties have been reported for the oil and may be synergised by synthetic pine oil.\(^{(5)}\) Antibacterial activity primarily versus organisms responsible for gut and throat infections has been documented,\(^{(12)}\) although a lack of antibacterial activity has also been reported.\(^{(5)}\)
Side-Effects, Toxicity

Concerns over the toxicity of calamus centre around the volatile oil and in particular on the β-asarone content. The level of β-asarone in the oil varies considerably between the different genetic species of calamus (see Constituents).

Feeding studies (rat) using the Indian oil (high β-asarone) have shown death, growth depression, hepatic and heart abnormalities, and serous effusion in abdominal and/or peritoneal cavities. A two–year study involving diet supplemented with calamus oil at 0, 500, 1000, 2500 and 5000 ppm, reported growth depression, and malignant duodenal tumours after 59 weeks at all levels of dietary supplementation. Tumours of the same type were not noted in the controls.

Genotoxic activity (strong induction of chromosomal aberrations, slight increase in the rate of sister chromatid exchanges) has been exhibited by β-asarone in human lymphocyte cultures in the presence of microsomal activation. Mutagenic activity (Ames) has been documented for root extracts, a tincture and β-asarone in one (TA100) of the various Salmonella typhimurium strains (TA98, 100, 1535, 1537, 1538) tested, but only in the presence of a microsomal activation mix. Lack of mutagenicity has also been reported for an organic extract, when tested in the above Salmonella typhimurium strains (except TA1538) with and without activation.

Acute toxicities (LD₅₀) quoted for the volatile oil from the Indian variety (high β-asarone content) include 777 mg/kg (rat, oral), >5 g/kg (guinea–pig, dermal), 221 mg/kg (rat, intraperitoneal). The oleoresin is stated to be toxic at 400 and 800 mg/kg (mouse, intraperitoneal). The LD₅₀ of asarone in mice is stated to be 417 mg/kg (oral) and 310 mg/kg (intraperitoneal).

Generally the oil is considered to be non–irritant, non–sensitising and non–phototoxic. However, bath preparations containing the oil have reportedly caused erythema, and dermatitis has been reported in hypersensitive individuals.
Contra-indications, Warnings

The toxicity of calamus oil has been associated with the β-asarone content.\(^{(16)}\) It has therefore been advised that only roots free from, or with a low content of β-asarone should be used in human phytotherapy.\(^{(16)}\) In foods and beverages, the level of β-asarone permitted in the final product is restricted (see Food use).

Use of the isolated oil is not recommended.\(^{(G49 \text{ G58})}\) External contact with the oil may cause an irritant reaction in sensitive individuals.

Calamus may potentiate monoamine oxidase inhibitor (MAOI) therapy (\textit{in vitro} MAOI activity, amine constituents), although the clinical significance of the \textit{in vitro} action has not been established.

\textbf{Pregnancy and lactation}

In view of the toxic properties associated with calamus, it should not be used during pregnancy or lactation. It is not known whether β-asarone is excreted into the breast milk. In general, the topical application of any undiluted oil is not recommended. Application of preparations containing calamus oil may provoke an irritant reaction and is therefore best avoided.
The phytochemistry of calamus, especially the oil, has been extensively investigated. Three genotypes (diploid, triploid and tetraploid) have been identified which are chemically distinct with respect to the β-asarone content. Spasmolytic and anti-ulcer effects documented for the oil support the traditional herbal uses of calamus. In addition, bitter principles documented as constituents may account for the use of the root in anorexia. However, in view of the toxic properties documented for the oil and associated with β-asarone, it has been recommended that only β-asarone-free calamus root should be used in phytotherapy. Use of the oil is not recommended due to its carcinogenic activity and its ability to cause kidney damage, tremors and convulsions.\textsuperscript{(G58)} Studies carried out to investigate the mutagenic potential of calamus have produced conflicting results.
References


Calendula
Species (Family)

*Calendula officinalis* L. (Compositae)
Synonym(s)
Gold-bloom, Marigold, Marybud, Pot Marigold
Part(s) Used

Flower
Pharmacopoeial and Other Monographs

BHP 1996\textsuperscript{(G9)}

BP 2002\textsuperscript{(G71)}

Complete German Commission E\textsuperscript{(G3)}

ESCOP 1996\textsuperscript{(G52)}

Martindale 33rd edition\textsuperscript{(G67)}

Mills and Bone\textsuperscript{(G50)}

PDR for Herbal Medicines 2nd edition\textsuperscript{(G36)}

Ph Eur 2004\textsuperscript{(G72)}

WHO volume 1 1999\textsuperscript{(G63)}
Legal Category (Licensed Products)

GSL (external use only)\(^{(G37)}\)
Constituents

See General References G2 G48 G52 G53 G62 G64.

**Flavonoids**
Pharmacopoeial standard not less than 0.4% flavonoids.\(^{G15 \ G28}\) Flavonol (isorhamnetin, quercetin) glycosides including isoquercitrin, narcissin, neohesperidoside, and rutin.\(^{1}\)

**Polysaccharides**
Three polysaccharides PS-I, -II and –III have a \((1\rightarrow3)\)-β-D-galactan backbone with short side chains at C-6, comprising \(α\)-araban-(1\rightarrow3)-araban, \(α\)-L-rhamnan-(1\rightarrow3)-araban or simple \(α\)-L-rhamnan moieties.\(^{2}\)

**Terpenoids**
Many components, including \(α\)- and \(β\)-amyrin, lupeol, longispinogenin, oleanolic acid, arnidiol, brein, calenduladiol, erythrodiol, faradiol, faradiol–3–myrctic acid ester, faradiol–3–palmitic acid ester,\(^{3}\) helantriols A1, B0, B1 and B2, lupeol, maniladiol, urs–12–en–3,16,21–triol, ursadiol; oleanolic acid saponins including calendulosides C–H;\(^{4}\) campesterol, cholesterol, sitosterol, stigmasterol and taraxasterol (sterols).\(^{5}\)

**Volatile oils**
Terpenoid components include menthone, isomenthone, caryophyllene and an epoxide and ketone derivative, pedunculatine, \(α\)- and \(β\)-ionone, a \(β\)-ionone epoxide derivative, dihydro actinidiolide.\(^{6}\)

**Other constituents**
Bitter (loliolide),\(^{7}\) arvoside A (sesquiterpene glycoside),\(^{8}\) carotenoid pigments\(^{9}\) and calendulin (gum).\(^{9}\)
Food Use

Calendula is not used in foods. In the USA, calendula is listed as GRAS (Generally Recognised As Safe).\(^{(G65)}\)
Calendula is stated to possess antispasmodic, mild diaphoretic, anti-inflammatory, anti-haemorrhagic, emmenagogue, vulnerary, styptic and antiseptic properties. Traditionally, it has been used to treat gastric and duodenal ulcers, amenorrhea, dysmenorrhea and epistaxis; crural ulcers, varicose veins, haemorrhoids, anal eczema, proctitis, lymphadenoma, inflamed cutaneous lesions (topically) and conjunctivitis (as an eye lotion). The German Commission E approved internal and external use for inflammation of oral and pharyngeal mucosa and external use in treatment of poorly healing sores.
Dosage

*Dried florets*
1–4 g or by infusion three times daily.\(^{(G7)}\)

*Liquid extract*
0.5–1.0 mL (1 : 1 in 40% alcohol) three times daily.\(^{(G7)}\)

*Calendula Tincture*
(BPC 1934) 0.3–1.2 mL (1 : 5 in 90% alcohol) three times daily.\(^{(G7)}\)

*External use*
Tincture–liquid extract (1 : 1) in 40% alcohol or tincture 1 : 5 in 90% alcohol. Apply to wounds as such and dilute 1 : 3 with water for compresses. Ointment 2.5%.\(^{(G52)}\)
Pharmacological Actions

In vitro and animal studies

Anti-inflammatory, antibacterial and antiviral activities have been reported for calendula.\(^{(10)}\) Weak anti-inflammatory activity in rats (carrageenan-induced oedema) has been reported.\(^{(11,12)}\) An aqueous ethanolic extract had mild dose-dependent action in the mouse croton oil test with 20% inhibition being reached at a dose of 1200 μg/ear, whereas a carbon dioxide extract exhibited 70% inhibition at the same concentration.\(^{(5,13)}\) The activity was shown to be due to the triterpenoids, the most active being a monoester of faradiol. Further separation of the triterpenoids has shown that the three most active compounds in the croton oil mouse test are faradiol-3-myristic acid ester, faradiol-3-palmitic acid ester and 4-taraxosterol.\(^{(3)}\)

A polysaccharide enriched extract showed strong concentration-dependent adhesive properties on porcine buccal membranes \textit{ex vivo}.\(^{(14)}\) Fluorescent labelled rhamnogalacturonan indicated the presence of polysaccharide layers on buccal membranes, leading to the suggestion that irritated buccal membranes may be smoothed by mucilage.

The formation of new blood vessels is an essential part of the wound-healing process. Angiogenic activity has been shown for a freeze-dried aqueous extract of calendula utilising the chick chorioallantoic membrane (CAM) assay.\(^{(15)}\) The number of microvessels in calendula-treated CAMs was significantly higher than in the control \((p < 0.0001)\). Furthermore, calendula-treated CAMs were positive for the glycosaminoglycan hyaluronan (HA) associated with neovascularisation. The presence of HA was not demonstrated in control CAMs.

A combination of allantoin and calendula extract applied to surgically induced skin wounds in rats has been reported to stimulate physiological regeneration and epithelisation.\(^{(16)}\) This effect was attributed to a more intensive metabolism of glycoproteins, nucleoproteins and collagen proteins during the regenerative period in the tissues.\(^{(16)}\) Allantoin applied on its own was found to exert a much weaker action.\(^{(16)}\)

A proprietary cream containing a combination of plant extracts, including calendula, has been reported to be effective in dextran and burn oedemas and in acute lymphoedema in rats. Activity against lymph oedema was primarily attributed to an enhancement of macrophage proteolytic activity.\(^{(17)}\) Slight increases in foot oedema were attributed to a vasodilatory action.

The trichomonacidal activity of calendula has been associated with the
An *in vitro* uterotonic effect has been described for calendula extract on rabbit and guinea–pig preparations.\(^{(18)}\)

Immunostimulant activity, assayed using granulocyte and carbon clearance tests, of calendula extracts has been attributed to polysaccharide fractions of high molecular weight.\(^{(19)}\) Polysaccharides PS-I, -II and -III have immunostimulant activity at concentrations of $10^{-5}$ to $10^{-6}$ mg/mL, stimulating phagocytosis of human granulocytes *in vitro*.\(^{(2)}\) A dry 70% ethanolic extract was not directly mitogenic, and was inhibitory in the mitogen–induced lymphocyte assay, causing stimulation at concentrations of 0.1–10 μg/mL, and inhibition at higher concentrations.\(^{(20)}\)

A 70% methanolic extract of calendula was successively extracted with ether, chloroform, ethyl acetate and *n*-butanol, leaving a residual aqueous extract. Each of the five extracts were concentrated and dissolved in 50% ethanol to produce 6% (w/v) solutions which were assessed for activity on liposomal lipid peroxidation induced by Fe$^{2+}$ and ascorbic acid. The ether, butanol and water extracts showed antioxidant activity.\(^{(21)}\)

The triterpenoid constituents of calendula are reported to be effective as spermicides and as antiblastocyst and abortion agents.\(^{(G53)}\)

*In vitro* cytotoxic activity and *in vivo* antitumour activity (against mouse Ehrlich carcinoma) have been documented for calendula extracts.\(^{(7)}\) The most active fraction *in vivo* (saponin–rich) was not the most active *in vitro*.\(^{(10)}\)

A 70% aqueous ethanolic extract had marked antiviral activity against influenza virus and herpes simplex virus.\(^{(G52)}\) A dichloromethane–methanol (1 : 1) extract exhibited potent anti-HIV activity in an *in vitro* MTT/tetrazolium–based assay.\(^{(22)}\) Uninfected Molt–4 cells were completely protected for up to 24 hours from fusion and subsequent death caused by co–cultivation with persistently infected U-937/HIV-1 cells. The organic extract caused a significant concentration- and time–dependent reduction of HIV-1 reverse transcriptase.\(^{(22)}\)

In a study in mice fed for three weeks with a diet containing either 0.1% or 0.4% of a calendula extract (containing 37% of esters of the carotenoid lutein), mammary tumour cells were infused into the mammary glands. Tumour latency increased, and tumour growth was inhibited in a dose–dependent manner by dietary lutein. In addition, dietary lutein was reported to enhance lymphocyte proliferation.\(^{(23)}\)
Clinical studies

A proprietary cream preparation containing several plant extracts, including calendula, has been reported to reduce pain associated with postmastectomy lymphoedema, although there was no significant clinical difference in the reduction of oedema between controls and experimental groups.\(^{(17)}\) Calendula tincture 20% has been reported to be useful in the treatment of chronic suppurative otitis.\(^{(24)}\) Calendula extracts are used to accelerate healing and to reduce inflammation.\(^{(9)}\) Thirty patients with burns or scalds were treated three times daily with a hydrogel containing 10% aqueous ethanolic extract of calendula for 14 days in an open, uncontrolled, pilot study.\(^{(25)}\) Improvement was noted for reddening, swelling, blistering, pain, soreness and heat sensitivity.
Side-effects, Toxicity

An aqueous extract of calendula had an LD$_{50}$ of 375 mg/kg (intravenous administration) and an LD$_{100}$ of 580 mg/kg (intraperitoneal administration) in mice.$^{(G52)}$ Aqueous ethanolic extracts (drug/extract ratio 1 : 1 and 0.5 : 1, 30% ethanol) had LD$_{50}$ values of 45 mg/mouse (subcutaneous administration) and 526 mg/100 g in rat (intravenous administration). An aqueous extract was not toxic following chronic administration to mice. Six saponins at doses of 400 μg were non-mutagenic in the Ames test using *Salmonella typhimurium* TA98 with and without S9 activation mixture.$^{(G52)}$ In *vitro* cytotoxicity has been reported for calendula extracts.$^{(10)}$ Extracts have been reported to be non-carcinogenic in rats and hamsters.$^{(G52)}$
Contra-indications, Warnings

Calendula may cause an allergic reaction in sensitive individuals, especially those with an existing hypersensitivity to other members of the Asteraceae/Compositae.

**Pregnancy and lactation**

Calendula is traditionally reputed to affect the menstrual cycle. An uterotonic effect (*in vitro*) has been reported, and the triterpenoid constituents are reported to be effective as spermaticides and as antiblastocyst and abortion agents. In view of the lack of toxicity data, the use of calendula is best avoided during pregnancy and lactation.
Pharmaceutical Comment

Phytochemical studies have reported four main groups of constituents, for calendula, namely flavonoids, polysaccharides, volatile oil and triterpenes. The latter seems to represent the principal group, with many compounds isolated including pentacyclic alcohols, glycosides (saponins) and sterols. Animal studies have reported wound-healing and anti-inflammatory effects, supporting the traditional uses of calendula in various dermatological conditions. The anti-inflammatory effect is due to the triterpenoid constituents although flavonoids may contribute to the activity. The reputed antispasmodic effect may be attributable to the volatile oil fraction. In addition, immunostimulant activity has been reported for high molecular weight polysaccharide components. Despite the popularity of calendula in herbal preparations there is little substantial clinical evidence to support its use.
References


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Capsicum
Species (Family)

Synonym(s)
Cayenne, Chilli Pepper, Hot Pepper, Paprika, Red Pepper, Tabasco Pepper
Part(s) Used

Fruit
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)

Complete German Commission E (Paprika)\(^{(G3)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)

Ph Eur 2004\(^{(G72)}\)

USP26/NF21\(^{(G73)}\)
Legal Category (Licensed Products)

GSL(\textsuperscript{G37})
Constituents
See General References G22 G41 G64.

Capsaicinoids
Up to 1.5%, usually 0.11%. Major components capsaicin (48.6%), 6,7-dihydrocapsaicin (36%), nordihydrocapsaicin (7.4%), homodihydrocapsaicin (2%) and homocapsaicin (2%).

Volatile oils
Trace. Over 125 components have been isolated with at least 24 characterised.

Other constituents
Carotenoid pigments (capsanthin, capsorubin, carotene, lutein), proteins (12–15%), fats (9–17%), vitamins including A and C.

Other plant parts
The plant material contains solanidine, solanine and solasodine (steroidal alkaloidal glycosides) and scopoletin (coumarin).
Food Use

Capsicum (chilli) peppers are widely used as a spice. Capsicum is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that capsicum can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\(^{(G16)}\) In the USA, capsicum is stated to be GRAS (Generally Recognised As Safe).\(^{(G41)}\)
Capsicum is stated to possess stimulant, antispasmodic, carminative, diaphoretic, counterirritant, antiseptic and rubefacient properties. Traditionally, it has been used for colic, flatulent dyspepsia without inflammation, chronic laryngitis (as a gargle), insufficiency of peripheral circulation and externally for neuralgia including rheumatic pains and unbroken chilblains (as a lotion/ointment). The German Commission E approved external use for treatment of painful muscle spasms in shoulder, arm and spine; arthritis, rheumatism, lumbago and chilblains. (G3)
Dosage

**Fruit**
30–120 mg three times daily.\(^{(G7)}\)

**Capsicum Tincture**
(BPC 1968) 0.3–1.0 mL; capsaicin content 0.005–0.01%.\(^{(G4)}\)

**Stronger Tincture of Capsicum**
(BPC 1934) 0.06–2.0 mL.

**Oleoresin**
0.6–2.0 mg.\(^{(G44)}\)

**Oleoresin, internal**
1.2 mg (maximum dose), 1.8 mg (maximum daily dose).\(^{(G37)}\)

**Oleoresin, external**
2.5% maximum strength.\(^{(G37)}\)

**Creams, ointments**
0.02–0.05%.\(^{(G4)}\)
Pharmacological Actions

The action of capsaicin on nervous, cardiovascular, respiratory, thermoregulatory and gastrointestinal systems has been reviewed.\(^{(1)}\) Capsaicin has been used as a neurochemical tool for studying sensory neurotransmission.\(^{(1)}\)

**In vitro and animal studies**

Infusion of capsaicin (200 μg/kg, by intravenous injection) has been reported to evoke dose–dependent catecholamine secretion (adrenaline, noradrenaline) from the adrenal medulla of pentobarbitone–anaesthetised rats.\(^{(2)}\)

The addition of capsaicin (0.014%) to a high–fat (30%) diet fed to rats was found to reduce serum–triglyceride concentrations but to have no effect on serum cholesterol or pre-β-lipoprotein concentrations.\(^{(3)}\) Capsaicin was thought to stimulate lipid mobilisation from adipose tissue. Lipid absorption was unaffected by capsaicin supplementation.\(^{(3)}\)

Activities of two hepatic enzymes, glucose–6–phosphate dehydrogenase and adipose lipoprotein lipase, were elevated in rats when capsaicin was added to the diet.\(^{(3)}\) Capsicum extracts fed orally to hamsters have been reported to significantly decrease hepatic vitamin A concentrations.\(^{(4)}\) Serum vitamin A concentrations were not affected.\(^{(4)}\)

Both the gastric and duodenal mucosae are thought to contain ‘capsaicin–sensitive’ areas which afford protection against acid- and drug–induced ulcers when stimulated by hydrochloric acid or by capsaicin itself. Stimulation causes an increase in mucosal blood flow and/or vascular permeability, inhibits gastric motility, and activates duodenal motility.\(^{(5)}\) Desensitisation of these areas, using a regimen involving subcutaneous or oral administration of capsaicin, is thought to remove the protection.\(^{(5)}\) However, capsaicin desensitisation was found to have little effect on peripheral responses to stress (i.e. ulcer formation) but did enhance central responses (increase in plasma corticosterone concentration) in rats.\(^{(6)}\) The increase in plasma corticosterone concentration observed in capsaicin–desensitised rats was similar in stressed and non–stressed animals.\(^{(6)}\)

Capsaicin was found to influence adrenal cortical activity independently of the presence of a stress factor and may represent a stressor in itself.\(^{(6)}\) Capsaicin desensitisation was not found to influence basal gastric acid secretion in non–stressed rats, but did lower pentagastrin–stimulated gastric output.\(^{(6)}\)
However, other results have reported that capsaicin desensitisation does increase acid secretion.\(^6\)

Capsicum (leaf and stem) has been reported to exhibit uterine stimulant activity in animal studies.\(^{30}\)

Pharmacokinetic studies in rats have reported that capsaicin is readily transported via the gastrointestinal tract and absorbed through non–active transport into the portal vein.\(^2\) Capsaicin is partly hydrolysed during absorption and the majority is excreted in the urine within 48 hours.\(^{2,7}\) Dihydrocapsaicin–hydrolysing enzyme is present in various organs of the rat but principally in the gastrointestinal tract and the liver. The biotransformation pathway of dihydrocapsaicin in the rat has been studied.\(^7\) Metabolites are mainly excreted as glucuronide conjugates in the urine.\(^7\)

**Clinical studies**

Ingestion of red chillies (10 g in wheatmeal) by controls and duodenal ulcer sufferers has been reported to have no significant effect on acid or pepsin secretion, or on sodium, potassium and chloride concentrations in the gastric aspirate.\(^8\) There was reported to be no apparent change (qualitative or quantitative) in mucous and no gastric mucosal erosion was evident.\(^8\) However, in contrast, capsicum has been shown to increase acid concentration and DNA content (indicating exfoliation of epithelial cells) of gastric aspirates in both control subjects and patients with duodenal ulcers.\(^1\) A study involving 18 healthy volunteers suggested that chilli (20 g in 200 mL water) protected against aspirin–induced gastroduodenal mucosal injury, compared with control (water).\(^9\)

Capsicum is applied externally as a counter–irritant in many preparations used for rheumatism, arthritis, neuralgia and lumbago. Clinical studies of topical preparations containing capsaicin have investigated its effectiveness in the treatment of chronic post–herpetic neuralgia, shingles, diabetic neuropathy, rhinopathy and neuropathic pain in cancer patients.\(^{4}\)

A systematic review of randomised, double–blind, placebo–controlled trials of topical capsaicin included 13 trials involving patients with diabetic neuropathy, osteoarthritis, post–herpetic neuralgia, postmastectomy pain and psoriasis.\(^{10}\) All the included trials reported that capsaicin was superior to placebo. However, the review drew cautious conclusions because blinding may have been compromised by the irritant effects of capsaicin.
Side-effects, Toxicity

Capsicum contains pungent principles (capsaicinoids) that are strongly irritant to mucosal membranes. Inhalation of paprika can produce a form of allergic alveolitis.\(^{(G51)}\)

Chronic administration of capsicum extract (0.5 μg capsaicin/kg body weight) to hamsters has been reported to be toxic.\(^{(4)}\) Treated animals did not survive beyond 17 months whereas all untreated controls survived beyond this period. In addition, eye abnormalities were observed in the treated animals. This effect was attributed to the depletion of substance P in primary afferent neurons by capsaicin, causing a loss of corneal pain sensation and subsequently the loss of protective corneal reflexes.\(^{(4)}\)

It is thought that metabolism of capsaicin and related analogues may reduce their acute toxicity.\(^{(7)}\) LD\(_{50}\) values stated for capsaicin in mice include 0.56 mg/kg (intravenous), 7.56 mg/kg (intraperitoneal), 9.00 mg/kg (subcutaneous) and 190 mg/kg (oral). In rats, an intraperitoneal LD\(_{50}\) of 10 mg/kg has been reported for capsaicin.\(^{(7)}\) The toxicity of capsaicinoids has reportedly not been ascribed to any one specific action but may be due to their causing respiratory failure, bradycardia and hypotension.\(^{(7)}\)
Contra-indications, Warnings

Capsicum may cause gastrointestinal irritation, although it has been stated that capsicum does not influence the healing of duodenal ulcers and does not need to be avoided by patients with this condition.\(^{(1)}\) Excessive ingestion may cause gastroenteritis, hepatic or renal damage.\(^{(G42)}\) Capsicum may interfere with monoamine oxidase inhibitors (MAOIs) and antihypertensive therapy (increased catecholamine secretion), and may increase the hepatic metabolism of drugs (glucose–6–phosphate dehydrogenase and adipose lipoprotein lipase activity elevated).

Pregnancy and lactation

There are no known problems with the use of capsicum during pregnancy, although it may cause gastrointestinal irritation and should therefore be used with caution. Doses should not greatly exceed amounts normally ingested in foods. It is not known whether the pungent components in capsicum are secreted into the breast milk.
Pharmaceutical Comment

Capsicum is commonly used in both foods and in medicinal products. The capsaicinoids are principally responsible for the biological activity of capsicum. These pungent principles are thought to stimulate and aid digestion and to act as a counter-irritant when applied externally. Capsaicin has also been used as a neurochemical tool for studying sensory neurotransmission. Topical creams containing capsaicin 0.025% and 0.075% are licensed in the UK for the treatment of pain in osteoarthritis, and painful diabetic neuropathy and post-herpetic neuralgia, respectively.\(^{11}\) Capsicum oleoresin and capsaicin are ingredients of a number of over-the-counter (OTC) topical preparations for relief of pain in muscle, tendon and joints.\(^{11}\)

Conflicting reports have been documented concerning the effect of capsicum on acid secretion and on ulcer healing. Capsaicin-sensitive areas of the gastric and duodenal mucosa are thought to provide protection against mucosal damage. It has been suggested that this protection is lost if the sensory fibres are desensitised. Whether oral consumption of capsicum by humans can cause desensitisation is unclear. The toxicity of capsicum extracts observed in animals is considered to be due to the capsaicinoid components. However, ingestion of capsicum in the diet is not thought to represent a health risk. Capsicium should not be ingested in doses greatly exceeding amounts normally used in foods.
See also General References G3 G5 G9 G12 G16 G22 G30 G31 G32 G33 G36 G37 G41 G43 G48 G51 G56 G61 G64.

Cascara
*Species (Family)*

*Rhamnus purshiana* DC. (*Frangula purshiana* (DC). A. Gray ex J.C. Cooper) (Rhamnaceae)
Synonym(s)
Cascara Sagrada, Rhamni Purshianae Cortex, Rhamnus
Part(s) Used

Bark
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)
BP 2002\(^{(G71)}\)
Complete German Commission E\(^{(G3)}\)
ESCOP 1997\(^{(G52)}\)
Martindale 33rd edition\(^{(G67)}\)
PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Ph Eur 2004\(^{(G72)}\)
USP26/NF21\(^{(G73)}\)
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents


**Anthracene glycosides**

Pharmacopoeial standard, not less than 8% hydroxyanthracene glycosides.\(^{(G15 \ G28)}\) Cascarosides A and B are anthrone C- and O-glycosides being 8-O-β-D-glucosides of 10-S-deoxyglucosyl aloe–emodin anthrone (aloin A) and of 10-R-deoxyglucosyl aloe–emodin anthrone (aloin B), respectively. Cascarosides C and D are the 8-O-β-D-glucosides of 10-(R)-(S)-deoxyglucosyl chrysophanol anthrone (chrysaloin A and B, respectively). Cascarosides E and F are the 8-O-β-D-glucosides of 10-deoxyglucosyl emodin–9–anthrone. The cascarosides comprise 60–70% of the total hydroxy anthracene complex. Aloins A and B, chrysaloins A and B account for 10–30% of the total hydroxy anthracene complex. The remaining 10–20% is a mixture of hydroxyanthracene O-glycosides including mono glucosides of aloe–emodin, chrysophanol, emodin and physcion.

**Other constituents**

Linoleic acid, myristic acid, syringic acid, lipids, resin and tannin.
Food Use

Cascara is listed by the Council of Europe as a natural source of food flavouring (category N4). This category indicates that while the use of cascara for flavouring purposes is recognised, it cannot be classified into the categories N1, N2 or N3 because of insufficient information.\(^{(G16)}\) In the USA, cascara has been approved for food use.\(^{(G41)}\)
Herbal Use

Cascara is stated to possess mild purgative properties and has been used for constipation. The German Commission E approved for use for treatment of constipation.\(^\text{G3}\)
Dosage

*Dried bark*
0.3–1 g single daily dose.\(^{(G3)}\)

*Infusion*
1.5–2 g in 150 mL water.\(^{(G3)}\)

*Cascara Liquid Extract*
(BP 1980) 2–5 mL.

*Preparations*
Equivalent to 20–30 mg hydroxy anthracene derivatives calculated as cascaroside A, daily.\(^{(G3)}\)
Pharmacological Actions

The laxative action of anthraquinone glycosides is well recognised (see Senna). Cascara has a laxative action.\(^{(G45)}\)

Clinical studies

Studies involving elderly patients suggest that cascara treatment, compared with placebo, leads to relief of constipation and increased bowel movements.\(^{(1)}\)
Side–effects, Toxicity

The side–effects and toxicity documented for anthraquinone glycosides are applicable (see Senna).\(^{(G22)}\)
Contra–indications, Warnings

Cascara is contra–indicated for patients with intestinal obstruction, acute intestinal inflammation, e.g. Crohn’s disease, colitis, appendicitis, abdominal pain of unknown origin, and in children under 12.\(^{(G3)}\) Cascara should not be used over an extended period of time.\(^{(G3)}\)

**Pregnancy and lactation**

Cascara should not be used during pregnancy and lactation.
Pharmaceutical Comment

The chemistry of cascara is characterised by the anthraquinone derivatives, especially the cascarosides. The laxative action of these compounds is well recognised. Cascara has been used extensively in conventional pharmaceutical preparations. Stimulant laxatives have largely been superseded by bulk-forming laxatives. However, the use of non-standardised anthraquinone-containing preparations should be avoided since their pharmacological effects will be variable and unpredictable. In particular, the use of products containing combinations of anthraquinone laxatives is not advisable.
References


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Species (Family)

*Cinnamomum cassia* Bl. (Lauraceae)
Synonym(s)
Cassia Bark, Cassia Lignea, Chinese Cinnamon, *Cinnamomum aromaticum* Nees, False Cinnamon
Part(s) Used

Bark
Pharmacopoeial and Other Monographs

BHP 1996\textsuperscript{(G9)}

Martindale 33rd edition\textsuperscript{(G67)}

PDR for Herbal Medicines 2nd edition\textsuperscript{(G36)}
Legal Category (Licensed Products)

GSL (oil)\textsuperscript{G37}
Constituents

See General References G41 G58 G59 G62 G64.

**Volatile oils**

1–2%. Mainly composed of cinnamaldehyde (75–90%). Other major components include salicylaldehyde, methylsalicylaldehyde, and methyl eugenol. Eugenol is reported to be absent. Cassia oil contains no monoterpenoids or sesquiterpenoids.\(^1\)

**Other constituents**

Calcium oxalate, coumarin, mucilage (higher content compared to cinnamon), resins, sugars and tannins (condensed). Complex diterpenoids have been isolated from cinnamomum cortex, for which *C. cassia* is used as a source.\(^1\)
Food Use

Cassia bark and oil are extensively used as food flavourings. A temporary estimated acceptable daily intake of cinnamaldehyde is 700 μg/kg body weight. In the USA, cassia is listed as GRAS (Generally Recognised As Safe).\(^{(G41)}\)
Herbal Use

Cassia is stated to possess carminative, antispasmodic, anti-emetic, antidiarrhoeal and antimicrobial properties. It has been used for flatulent dyspepsia, flatulent colic, diarrhoea, the common cold, and specifically for colic or dyspepsia with flatulent distension and nausea.\(^{(G7)}\) Cassia bark is also documented to possess astringent properties.\(^{(G41\ G64)}\) Carminative and antiseptic properties are documented for the oil.\(^{(G41)}\)
Dosage

*Dried bark*
0.5–1 g or by infusion three times daily.\(^{(G7)}\)

*Oil of cassia*
(BPC 1949) 0.05–0.2 mL three times daily.\(^{(G7)}\)
Pharmacological Actions

In vitro and animal studies

Anti-ulcerogenic properties have been described for two propionic derivatives isolated from cassia.\(^{(2)}\) An in vivo study using rats reported activity against a variety of ulcerogens including serotonin, phenylbutazone, ethanol, water immersion and stress. The compounds were thought to act by improving gastric blood flow rather than by inhibiting gastric secretion.

Many pharmacological investigations have been carried out on cinnamomi cortex, for which sources include *C. cassia* (cassia) and *Cinnamomum zeylanicum* (cinnamon). These studies have either looked at the volatile oil, in particular the major constituent cinnamaldehyde, or at parts excluding the oil.\(^{(1)}\)

Activities documented for cinnamaldehyde include CNS stimulation (low dose), sedation (high dose), hypothermic and antipyretic actions;\(^{(1,G41)}\) antibacterial and antifungal activity, acceleration of catechol amine (mainly adrenaline) release from the adrenal glands, weak papaverine-like action, increase in peripheral blood flow, hypotension, bradycardia and hyperglycaemia have also been reported.\(^{(1)}\) However, these actions are of low potency and, in addition, much of the cinnamaldehyde content of cassia is thought to be lost by evaporation and auto-oxidation during decoction of the crude drug. The contribution of cinnamaldehyde to the overall therapeutic efficacy of cassia has therefore been doubted.\(^{(1)}\)

Actions observed for essential oil-free aqueous extracts have been reported to be weak, and the only appreciable effects are prolongation of barbiturate-induced sedation and a slight reduction of acetic acid-induced writhing.\(^{(1)}\)

In vivo inhibitory activity against complement formation has been documented and attributed to the diterpenoid and condensed tannin constituents.\(^{(1)}\) Anti-inflammatory activity exhibited by the Japanese plant *Cinnamomum sieboldii* Meisn (also used as a source for cassia bark), has been attributed to a series of condensed tannin constituents.\(^{(1)}\) Antiplatelet aggregation and antithrombotic actions have also been reported. These actions, together with the documented anti-inflammatory activity, are thought to contribute to the suppression of thrombus formation in certain diseases.\(^{(1)}\)

Antitumour activity has been described and the activity depends on the plant source used.\(^{(1)}\)
Side–effects, Toxicity

Allergic reactions, mainly contact sensitivity, to cassia oil and bark have been reported.\(^{(G51 \ G58)}\) Cinnamaldehyde in toothpastes and perfumes has also been reported to cause contact sensitivity.\(^{(G51)}\) Cassia oil is stated to cause dermal and mucous membrane irritation.\(^{(G58)}\) The irritant and sensitising properties of cassia oil have been attributed to cinnamaldehyde.\(^{(G58)}\) The dermal LD\(_{50}\) value for cassia oil is stated as 320 mg/kg body weight.\(^{(G58)}\)
Contra-indications, Warnings

Contact with cassia bark or oil may cause an allergic reaction. Cassia oil is stated to be one of the most hazardous oils and should not be used on the skin in concentrations of more than 0.2%. \((G58)\)

Pregnancy and lactation

There are no known problems with the use of cassia during pregnancy, provided that amounts taken do not exceed those generally used in foods.
Pharmaceutical Comment

Cassia is similar in composition to cinnamon and both are widely used as flavouring agents in foods, and in pharmaceutical and cosmetic preparations. Cassia oil is stated to be inferior in flavour to cinnamon oil. The reputed herbal uses of cassia have been attributed to the oil. Cassia contains an irritant and sensitising principle in the oil, cinnamaldehyde, and should not be used in amounts generally exceeding those used in foods. It has been recommended that the oil should never be applied topically.
References

See also General References G9 G11 G19 G20 G31 G36 G37 G41 G43 G51 G58 G59 G62 G64.


Cat’s Claw
Species (Family)

*Uncaria tomentosa* (Willd.) DC., *Uncaria guianensis* (Aubl.) Gmel. (Rubiaceae)
Synonym(s)

Life-giving Vine of Peru, Savéntaro, Uña de gato
Part(s) Used
Roots, root bark, stem bark and leaves
Pharmacopoeial and Other Monographs

None
Legal Category (Licensed Products)

Cat’s claw is not included in the GSL.(G37)
Constituents

Alkaloids
Both *U. tomentosa* and *U. guianensis* yield oxindole alkaloids, includingisorhynchophylline and rhynchophylline and their N-oxides, mitraphylline and the indole alkaloids dihydrocorynantheine, hirsutine and hirsuteine.\(^{(1)}\) *U. tomentosa* also contains isomitraphylline, its N-oxide, dihydrocorynantheine N-oxide and hirsutine N-oxide.\(^{(1)}\)

There are two chemotypes of *U. tomentosa* which differ markedly in their patterns of alkaloids present in the root bark; in addition, the alkaloidal pattern of individual plants changes with time.\(^{(2,3)}\) One chemotype primarily contains the pentacyclic oxindole alkaloids pteropodine, isopteropodine, mitraphylline, isomitraphylline, uncarine F and speciophylline,\(^{(4)}\) whereas the other chemotype primarily contains the tetracyclic oxindole alkaloids rhynchophylline and isorhynchophylline.\(^{(1)}\) Although a particular plant may contain either tetracyclic or pentacyclic oxindole alkaloids predominantly, both types of alkaloids can co–occur in the same plant.\(^{(3,5)}\)

Other constituents
Quinovic acid glycosides have been isolated from both species.\(^{(6–9)}\) Polyhydroxylated triterpenes\(^{(10)}\) and steroids (β-sitosterol, stigmasterol, campesterol)\(^{(11)}\) occur in *U. tomentosa*. An unidentified South American species of *Uncaria* (presumably either *U. guianensis* or *U. tomentosa*) contains polyphenols ((−)-epicatechin and procyanidins).\(^{(4)}\)
Food Use

Cat’s claw is not used in foods.
Herbal Use

Cat’s claw is stated to possess anti-inflammatory, antiviral, antioxidant, immunostimulating, antirheumatic and anticancer properties. It is native to the Amazon and has been used traditionally to treat gonorrhoea, dysentry, arthritis, rheumatism, gastric ulcers and various tumours.\(^{(12)}\) It is also reputed to be a contraceptive.
Dosage
Commercial products (tablets, capsules) contain varying amounts of material, ranging from 25 to 300 mg standardised extract and from 400 mg to 5 g of plant material. (G31)
Pharmacological Actions

Several pharmacological activities have been documented for cat’s claw, including anti-inflammatory, antimutagenic, antitumour, antioxidant and immuno stimulating properties. The pharmacological activities of cat’s claw have been reviewed.\(^{(12,13)}\)

**In vitro and animal studies**

Certain oxindole alkaloids isolated from *U. tomentosa* (isopteropodine, pteropodine, isomitraphylline, isorhynchophylline) have been shown to enhance phagocytosis markedly in vitro.\(^{(14)}\) Pentacyclic oxindole alkaloids from *U. tomentosa* have been reported to induce the release of a lymphocyte proliferation–regulating factor from human endothelial cells; tetracyclic oxindole alkaloids were found to reduce the activity of pentacyclic oxindole alkaloids on these cells in a dose–dependent manner.\(^{(12,15)}\) Stem bark extracts of *U. tomentosa* have also been shown to stimulate interleukin 1 (IL-1) and interleukin 6 (IL-6) production in vitro in rat alveolar macrophages in a dose–dependent manner (range 0.025–0.1 mg/mL) and to potentiate the production of IL-1 and IL-6 in lipopolysaccharide–stimulated macrophages.\(^{(16)}\)

Extracts and fractions of *U. tomentosa* bark have shown no mutagenic effect but demonstrated a protective antimutagenic effect in vitro against 8–methoxypsoralen- and UVA-induced photomutagenesis in *Salmonella typhimurium* TA102.\(^{(17)}\) It was suggested that this antimutagenic activity may be due to an antioxidant effect of *U. tomentosa*.\(^{(17)}\)

**In vitro** antioxidant activity of stem bark and root extracts of *U. tomentosa* has been demonstrated in an assay using *tert*-butylhydroperoxide–initiated chemoluminescence in rat liver homogenates.\(^{(18)}\) Extracts also prevented free radical–mediated DNA sugar damage.\(^{(18)}\)

**In vitro** antitumour activity of water extracts of *U. tomentosa* (C-Med–100) has been shown in a human leukaemic cell line (HL-60) and a human Epstein–Barr virus (EBV)-transformed B lymphoma cell line (Raji).\(^{(19)}\) The suppressive effect of *U. tomentosa* extracts on tumour cell growth appear to be mediated through induction of apoptosis.\(^{(19)}\) The pentacyclic oxindole alkaloids uncarine C and uncarine E from *U. guianensis* have been identified as cytotoxic and DNA-damaging agents in a yeast–based assay.\(^{(20)}\) These alkaloids also showed moderate cytotoxicity to several mammalian cell lines, including human lung carcinoma.\(^{(20)}\) **In vitro**, aqueous extracts of *U. tomentosa* bark appear to interact with oestrogen receptor–binding sites.\(^{(21)}\)
Rhynchophylline has been reported to inhibit rat and rabbit platelet aggregation *ex vivo*. Studies in cats and dogs have reported that rhynchophylline has a negative inotropic effect which can contribute to a hypotensive effect. Rhynchophylline and isorhynchophylline have been reported to have negative chronotropic and inotropic effects in isolated guinea-pig atria. Isorhynchophylline has been reported to have hypotensive effects in rats and dogs.

Trichloromethane/methanol and aqueous extracts of cat’s claw (*U. tomentosa*) bark have demonstrated anti-inflammatory activity in the rat paw carrageenan-induced oedema test; a quinovic acid glycoside was identified as one of the active principles. An aqueous extract of cat’s claw (*U. tomentosa*) bark was reported to protect against oxidant-induced stress *in vitro* and to attenuate indometacin-induced chronic intestinal inflammation in rats. Cat’s claw extract was found to prevent the activation of the transcription factor NF-κB, which suggests a mechanism for the anti-inflammatory activity of cat’s claw.

Quinovic acid glycosides have demonstrated antiviral activity in *in vitro* tests against the RNA virus vesicular stomatitis virus. Two quinovic acid glycosides also demonstrated *in vitro* activity against rhinovirus type 1B.

Receptor-binding assays using dihydrocorynantheine isolated from the branchlet and hook of *Uncaria sinensis* (and also found in *U. tomentosa*) have shown that this alkaloid is a partial agonist for serotonin receptors.

**Clinical studies**

There is a lack of clinical evidence to support the activities of cat’s claw. A decoction of *U. tomentosa* bark ingested daily for 15 days by a smoker decreased the mutagenicity induced in *S. typhimurium* TA98 and TA100 by the subject’s urine; urine from a non-smoker who ingested the same regimen of *U. tomentosa* did not show any mutagenic activity before, during or after treatment.

In an uncontrolled study, 13 HIV-positive individuals who refused to receive other therapies ingested 20 mg daily of an extract of *U. tomentosa* root (containing 12 mg total pentacyclic oxindole alkaloids per gram) for 2.2–5 months. The total leukocyte number in the group was unchanged, compared with pretreatment values, whereas the relative and absolute lymphocyte count increased significantly. No significant changes in T4/T8 cell ratios were observed.
Side-effects, Toxicity

There has been a report of acute renal failure in a Peruvian woman with systemic lupus erythematosus who had added a product containing cat’s claw (one capsule four times daily, obtained from a local herbal shop) to her regimen of prednisone, atenolol, metolazone, furosemide and nifedipine.\(^{(29)}\) The patient had a serum creatinine concentration of 3.6 mg/dL and was diagnosed with acute allergic interstitial nephritis. She was advised to discontinue cat’s claw and, one month later, her renal function had improved (serum creatinine 2.7 mg/dL).

Data on the acute oral toxicity of *U. tomentosa* aqueous root extract (containing 35 mg total pentacyclic oxindole alkaloids per gram) in mice and four-week oral toxicity of an aqueous extract of *U. tomentosa* root (containing 7.5 mg total oxindole alkaloids per gram) in rats administered 1000 mg/kg/day have been summarised.\(^{(12)}\) The acute median LD\(_{50}\) to mice was found to be greater than 16 g/kg body weight. In the study in rats, a slight but statistically significant increase in the percentage of lymphocytes and a decrease in the percentage of neutrophil granulocytes were seen. In addition, an increase in the relative weight of the kidneys in rats of both sexes was noted, although kidney histology was normal.

*In vitro*, extracts of *U. tomentosa* have been shown to possess antitumour activity and to stimulate production of the cytokines IL-1 and IL-6, both of which are known to initiate a cascade of defence activities of the immune system. Oxindole alkaloids from *U. tomentosa* have been reported to enhance phagocytosis *in vitro* (see *In vitro* and animal studies).

The *in vitro* toxicity of aqueous extracts of *U. tomentosa* has been evaluated in bioassays using Chinese hamster ovary (CHO) cells and bacterial cells (*Photobacterium phosphoreum*).\(^{(30)}\) At the concentrations used (10−100 mg/mL), the extracts did not show a significant cytotoxic effect in CHO cells and demonstrated a non–toxic effect in the bacterial cells used.
Contra–indications, Warnings

In view of its immunostimulant properties, cat’s claw may interfere with immunosuppressive therapy. Extracts of the pentacyclic chemotype of U. tomentosa should be avoided where there is a risk of organ rejection in patients undergoing transplants; this includes bone marrow transplants.\(^{13}\)

In view of the inhibitory effects of rhynchophylline on platelet aggregation,\(^{22,23}\) it has been stated that cat’s claw is contra–indicated in patients receiving anticoagulants and in those with coagulation disorders.\(^{G31}\)

Animal studies have reported hypotensive effects with rhynchophylline\(^{24}\) and isorhynchophylline;\(^{26}\) thus, cat’s claw should be used with caution in patients receiving antihypertensive agents.\(^{G31}\)

**Pregnancy and lactation**
The safety of cat’s claw has not been established. In view of the lack of toxicity data, use of cat’s claw during pregnancy and lactation should be avoided. In addition, use in children (<3 years) is not advised.
Pharmaceutical Comment

The chemistry of cat’s claw is well documented. Reported pharmacological activities are mainly associated with the oxindole alkaloids and the quinovic acid glycosides.

The two species *U. tomentosa* and *U. guianensis* may be confused. In addition, there are two chemotypes of *U. tomentosa*, one predominantly producing pentacyclic oxindole alkaloids, and the other tetracyclic oxindole alkaloids. Since the tetracyclic oxindole alkaloids have been reported to antagonise the immunostimulant effect of the pentacyclic oxindole alkaloids on human cells *in vitro*, it has been stated that mixtures of the two chemotypes of cat’s claw are unsuitable for therapeutic use unless certified to contain less than 0.02% tetracyclic oxindole alkaloids.

Documented scientific evidence from *in vitro* and, to a lesser extent, animal studies provides some supportive evidence for some of the uses of cat’s claw. However, there is a lack of clinical data, and well–designed clinical trials involving adequate numbers of patients and using standardised preparations manufactured from the appropriate chemotype are necessary.

In view of the lack of toxicity and safety data, excessive use of cat’s claw should be avoided. Individuals wishing to use cat’s claw concurrently with conventional medicines should first seek advice from an appropriate healthcare professional.
References


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Species (Family)

*Apium graveolens* L. (Apiaceae/Umbelliferae)
Synonym(s)

Apii Fructus, Celery Fruit, Celery Seed, Smallage
Part(s) Used
Fruit
Pharmacopoeial and Other Monographs

BHC 1992\textsuperscript{(G6)}

BHP 1996\textsuperscript{(G9)}

Martindale 33rd edition\textsuperscript{(G67)}

PDR for Herbal Medicines 2nd edition\textsuperscript{(G36)}
Legal Category (Licensed Products)

GSL(G37)
Constituents
See General References G2 G6 G22 G38 G41 G48 G57 G58 G64.

**Flavonoids**
Apigenin, apiin, isoquercitrin and others.\(^{(1)}\)

**Furanocoumarins**
Apigravin, apiumentin, apiumoside, bergapten, celerin, celereoside, isoimperatorin, isopimpinellin, ostheno, rutaretin, seselin, umbelliferone and 8-hydroxy-5-methoxypsoralen.\(^{(1–9)}\)

Low concentrations (not exceeding 1.3 ppm) of furanocoumarins have been identified in commercial celery,\(^{(10)}\) although concentrations are reported to rise considerably in diseased stems.\(^{(11)}\)

**Volatile oils**
2–3%. Many components including limonene (60%) and selenine (10–15%), and various sesquiterpene alcohols (1–3%), e.g. α-eudesmol and β-eudesmol, santalol.\(^{(12,13)}\) Phthalide compounds, 3-\(\text{-n}\)butyl phthalide and sedanenolide, provide the characteristic odour of the oil (presence of sedanolide and sedanonic anhydride disputed).\(^{(14,15)}\)

**Other constituents**
Choline ascorbate,\(^{(16)}\) fatty acids (e.g. linoleic, myristic, myristicic, myristoleic, oleic, palmitic, palmitoleic, petroselinic and stearic acids).
Food Use

Celery is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that celery can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product. Celery stem (not the fruit) is commonly used in foods. In the USA, celery seed is listed as GRAS (Generally Recognised As Safe).
Herbal Use

Celery is stated to possess antirheumatic, sedative, mild diuretic and urinary antiseptic properties. It has been used for arthritis, rheumatism, gout, urinary tract inflammation, and specifically for rheumatoid arthritis with mental depression. (G2 G6 G7 G8 G64)
Dosage

*Dried fruits*
0.5–2.0 g or by decoction 1 : 5 three times daily.\(^{(G7)}\)

*Liquid extract*
0.3–1.2 mL (1 : 1 in 60% alcohol) three times daily.\(^{(G7)}\)

*Liquid Extract of Celery*  
(BPC 1934) 0.3–1.2 mL.
Pharmacological Actions

**In vitro and animal studies**

In mice, sedative and antispasmodic activities have been documented for the phthalide constituents.\(^{17,G22}\) Celery seed oil has been reported to exhibit bacteriostatic activity against *Bacillus subtilis*, *Vibrio cholerae*, *Staphylococcus aureus*, *Staphylococcus albus*, *Shigella dysenteriae*, *Corynebacterium diphtheriae*, *Salmonella typhi*, *Streptococcus faecalis*, *Bacillus pumilus*, *Streptococcus pyogenes* and *Pseudomonas solanacearum*.\(^9\) No activity was observed against *Escherichia coli*, *Sarcina lutea* or *Pseudomonas aeruginosa*.

Apigenin has exhibited potent antiplatelet activity *in vitro*, inhibiting the aggregation of rabbit platelets induced by collagen, ADP, arachidonic acid and platelet–activating factor (PAF), but not that induced by thrombin or ionophore A23187.\(^{18}\)

Studies with celery plant extracts have demonstrated anti–inflammatory activity in the mouse ear test and against carrageenan–induced rat paw oedema,\(^{19}\) and a hypotensive effect in rabbits and dogs after intravenous administration.\(^{G41}\) In addition, hypoglycaemic activity has been documented.\(^{G22}\)

Celery juice has been reported to exhibit choleretic activity and the phthalide constituents are stated to possess diuretic activity.\(^{13}\)

**Clinical studies**

None documented for celery fruit. Hypotensive activity was reported in 14 of the 16 hypertensive patients given a celery plant extract.\(^{G41}\)
None documented for celery fruit. Photosensitivity reactions have been reported as a result of external contact with celery stems.\textsuperscript{(20,21,G51)} These reactions have been attributed to the furanocoumarin constituents which are known to possess photosensitising properties.\textsuperscript{(11,22)} The concentrations of these compounds are reported to increase considerably in diseased celery stems.\textsuperscript{(11,22)} It is thought that psoralen, the most potent phototoxic furanocoumarin, acts as a transient precursor for other furanocoumarins and does not accumulate in celery.\textsuperscript{(5,11)}

Instances of allergic and anaphylactic reactions to celery have also been documented\textsuperscript{(23)} following oral ingestion of the stems.\textsuperscript{(24)} Celery allergy is reported to be mediated by IgE antibodies and an association between pollen and celery allergy has been postulated, although the common antigen had not been determined.\textsuperscript{(25)}

Cross-sensitivities to celery have been documented in patients with existing allergies to dandelion and wild carrot.\textsuperscript{(G51)}

Acute LD\textsubscript{50} values (rats, by mouth; rabbits, dermal) have been reported as greater than 5 g/kg body weight.\textsuperscript{(26)} Celery seed oil is stated to be non-irritant, non-phototoxic and non-sensitising in humans.\textsuperscript{(26,G58)}
Contra–indications, Warnings

Celery fruit contains phototoxic compounds, furanocoumarins, which may cause photosensitive reactions. Celery fruit may precipitate allergic reactions, particularly in individuals with existing plant, pollen or food allergies. Diseased celery stems (indicated by a browning of the stem) should not be ingested.

Pregnancy and lactation

Celery fruit is reputed to affect the menstrual cycle and to be abortifacient.\(^{(G30)}\) Uterine stimulant activity has been documented for the oil,\(^{(G22 G30)}\) and the use of celery fruits is contra–indicated during pregnancy.\(^{(G49)}\) This does not refer to celery stems that are commonly ingested as a food, although excessive consumption should be avoided.
Pharmaceutical Comment

Celery fruit should not be confused with the commercial celery stem, which is commonly eaten as a food. The chemistry of celery fruit is well studied and the phototoxic furanocoumarin constituents are well documented. Phototoxicity appears to be associated with the handling of the celery stems, especially diseased plant material. Limited scientific evidence is available to justify the herbal uses of celery, although bacteriostatic activity has been documented for the oil. Celery fruit should be used cautiously in view of the documented allergic reactions.
References


18. Teng CM et al. Inhibition of platelet aggregation by apigenin from *Apium


Species (Family)

*Centaurium erythraea* Rafin. (Gentianaceae)
Synonym(s)

Part(s) Used

Herb
Pharmacopoeial and Other Monographs

BHP 1996\textsuperscript{G9}

BP 2002\textsuperscript{G71}

Complete German Commission E\textsuperscript{G3}

ESCOP 1999\textsuperscript{G52}

Martindale 33rd edition\textsuperscript{G67}

PDR for Herbal Medicines 2nd edition\textsuperscript{G36}

Ph Eur 2004\textsuperscript{G72}
Legal Category (Licensed Products)

GSL\textsuperscript{(G37)}
Constituents

See General References G2 G41 G64.

**Acids**
Phenolic. Protocatechuic, \( m \)- and \( p \)-hydroxybenzoic, vanillic, syringic, \( p \)-coumaric, ferulic, sinapic and caffeic, hydroxyterephthalic and 2,5-dihydroxyterephthalic acids among others.

**Alkaloids**
Pyridine–type. Traces of gentianine, gentianidine, gentioflavine and others.

**Monoterpenoids**
Iridoids (bitters). \(^{1,2}\) Gentiopicroside (about 2%) as major, others include centapicrin, gentioflavoside, sweroside and swertiamarin; intensely bitter \( m \)-hydroxybenzoylesters of sweroside and catapicrin.

**Triterpenoids**
Includes \( \alpha \)- and \( \beta \)-amyrin, erythrodiol, crataegolic acid, oleanolic acid and sitosterol.

**Xanthones**
Highly methylated xanthones, including eustomin and 8-demethyleustomin.

**Other constituents**
Flavonoids, fatty acids, in, alkanes and waxes.
Food Use

Centaury is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that centaury can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\(^{(G16)}\) In the USA, the bitter properties of centaury are utilised in alcoholic and non–alcoholic beverages with maximum permitted doses between 0.0002\% and 0.0008\%.\(^{(G41)}\)
Centaury is reputed to act as a bitter, aromatic and stomachic. Traditionally, it has been used for anorexia and dyspepsia.
Dosage

*Herb*
2–4 g or by infusion three times daily.\(^{G7}\)

*Liquid extract*
2–4 mL (1 : 1 in 25% alcohol) three times daily.\(^{G7}\)
Pharmacological Actions

Centaury is stated to have bitter tonic, sedative and antipyretic properties.\(^{(G41)}\) The antipyretic activity is stated to be due to the phenolic acids.\(^{(G45)}\) Gentiopicrin is stated to have antimalarial properties.\(^{(G48)}\)

**In vitro and animal studies**

Anti-inflammatory activity has been documented in two rat models; sub chronic inflammation (air pouch granuloma and polyarthritis) test,\(^{(3)}\) and the carrageenan rat paw oedema test (19% compared to 45% with indometacin).\(^{(4)}\) Antipyretic activity has also been exhibited by a centaury extract against experimentally induced hyperthermia in rats, although pretreatment with the extract did not prevent hyperthermia.\(^{(3)}\) In the same study, no analgesic activity could be demonstrated in mice (writhing syndrome and hotplate models).\(^{(3)}\) Gentiopicroside (30 mg/kg/day intra peritoneally) inhibited tumour necrosis factor (TNF) production in carbon tetrachloride–induced and bacillus Calmette–Guérin/lipopolysaccharide–induced models of hepatic injury in mice.\(^{(G52)}\)

In rats, anticholinesterase activity has been demonstrated for swertiamarin in a dose–dependent manner following oral administration, demonstrated by inhibition of carbachol–induced contraction of proximal colon.\(^{(G52)}\) In mice, gentianine has central nervous system (CNS)-depressant activity at oral doses of 30 mg/kg, demonstrated by inhibition of spontaneous movement and prolonged hexobarbital–induced sleeping time.\(^{(G52)}\) Anti-ulcerogenic and inhibitory gastric secretion in rats (100 mg/kg) have been shown for gentianine.\(^{(G52)}\)
Side–effects, Toxicity

An alcoholic extract of centaury (200 mL/plate) was antimutagenic in *Salmonella typhimurium* strains TA8 and TA100.\(^{(G52)}\)
Contra–indications, Warnings

Centaury is contra–indicated for individuals with peptic ulcers.\(^{(G52)}\)

**Pregnancy and lactation**

The safety of centaury taken during pregnancy has not been established. In view of the lack of toxicity data, use of centaury during pregnancy and lactation is best avoided.
Pharmaceutical Comment

There is little published information specifically concerning *C. erythraea*. Bitter components support the traditional use of centaury as an appetite stimulant, although it is said to be less active than comparable bitter herbs, such as gentian. In view of the lack of pharmacological and toxicological data, excessive use should be avoided.
References

See also General References G2 G3 G9 G15 G16 G28 G31 G36 G37 G43 G48 G52 G56 G64.


Species (Family)

*Selenicereus grandiflorus* (L.) Britt. & Rose (Cactaceae)
Synonym(s)

*Cactus grandiflorus*, *Cereus grandiflorus* Mill., Night Blooming Cereus
Part(s) Used

Stem
Pharmacopoeial and Other Monographs

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

Cereus is not included in the GSL.\(^{(G37)}\)
Constituents
See General References G22 G40 G64.

**Alkaloids**  
Isoquinoline-type. Unidentified alkaloids.\(^1\)

**Amines**  
Tyramine\(^2\), hordenine,\(^3\) previously referred to as cactine.

**Flavonoids**  
Rutin, kaempferitrin, hyperoside, isorhamnetin-3-β-(galactosyl)-rutinoside.

**Other constituents**  
Resin
Food Use

Cereus is not used in foods.
Herbal Use

Cereus is reputed to act as a cardiac stimulant and as a partial substitute for digitalis, although there is no proof of its therapeutic value. Cereus has been used in cases of dropsy and various cardiac affections. (G10 G64)
Dosage

Liquid extract of cereus
(BPC 1934) 0.06–0.6 mL.

Tincture of cereus
(BPC 1934) 0.12–2.0 mL.
Pharmacological Actions

*In vitro and animal studies*

None documented for cereus. Cereus is reported to contain a cardiotonic amine, tyramine, which has positive inotropic activity.
Side-effects, Toxicity

The fresh juice of cereus is irritant to the oral mucosa, causing a burning sensation, nausea and vomiting. Diarrhoea has also been reported following cereus consumption. (G22)
Contra–indications, Warnings

In view of the documented tyramine content, excessive doses of cereus may interact with concurrent monoamine oxidase inhibitor (MAOI) treatment and may affect patients with an existing cardiac disorder.

**Pregnancy and lactation**
The safety of cereus has not been established. In view of the limited information available on cereus, its use during pregnancy and lactation should be avoided.
Pharmaceutical Comment

Little phytochemical or pharmacological information has been documented for cereus, although the presence of tyramine, a cardiotonic amine, may support the traditional use of cereus as a cardiac stimulant. Cardiac complaints are not considered to be suitable for self-medication.
References

See also General References G10 G22 G31 G36 G37 G40 G48 G64.

Chamomile, German
Species (Family)

*Matricaria recutita* L. (Asteraceae/Compositae)
Synonym(s)

*Chamomilla recutita* (L.) Rauschert, Hungarian Chamomile, *Matricaria chamomilla* L., Matricaria Flowers, Sweet False Chamomile, Wild Chamomile
Part(s) Used

Flowerhead
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)
BHP 1996\(^{(G9)}\)
BP 2002\(^{(G71)}\)
Complete German Commission E\(^{(G3)}\)
ESCOP 1999\(^{(G52)}\)
Martindale 33rd edition\(^{(G67)}\)
Mills and Bone\(^{(G50)}\)
PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
USP26/NF21\(^{(G73)}\)
WHO volume 1 1999\(^{(G63)}\)
Constituents
See General References G2 G6 G22 G38 G41 G48 G52 G64.

Coumarins
Umbelliferone and its methyl ether, heniarin.

Flavonoids
Apigenin, apigetrin, apiin, luteolin, quercetin, quercimeritrin and rutin.

Volatile oils
0.24–1.9%. Pharmacopoeial standard not less than 4 mg/kg blue oil. (G15 G28) Main components are (−)-α-bisabolol (up to 50%) (1) and chamazulene (1–15%). (2) Others include (−)-α-bisabolol oxide A and B, (−)-α-bisabolone oxide A, spiroethers (e.g. cis- and trans-en–yn–dicycloether), sesquiterpenes (e.g. anthecotulid), cadinene, farnesene, furfural, sphanthulenol and proazulenes (e.g. matricarin and matricin).

Chamazulene is formed from matricin during steam distillation of the oil. It varies in yield depending on the origin and age of the flowers. (2)

Other constituents
Amino acids, anthemic acid (bitter), choline, polysaccharide, plant and fatty acids, tannin and triterpene hydrocarbons (e.g. triacontane).
Food Use

German chamomile is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that chamomile can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\textsuperscript{(G16)} German chamomile is commonly used in herbal teas. In the USA, German chamomile is listed as GRAS (Generally Recognised As Safe).\textsuperscript{(G41)}
German chamomile is stated to possess carminative, antispasmodic, mild sedative, anti-inflammatory, antiseptic and anticatarrhal properties. It has been used for flatulent nervous dyspepsia, travel sickness, nasal catarrh, nervous diarrhoea, restlessness and specifically for gastrointestinal disturbance with associated nervous irritability in children. It has been used topically for haemorrhoids, mastitis and leg ulcers. German Commission E approved use for gastrointestinal spasms and inflammatory diseases of the gastrointestinal tract and externally for skin and mucous membrane inflammation and bacterial skin diseases including oral cavity and gums. It is also approved for inflammations and irritations of the respiratory tract (by inhalation) and ano-genital inflammation (baths and irrigation).
Dosage

*Dried flowerheads*
2–8 g or by infusion three times daily.\(^{(G7)}\)

*Liquid extract*
1–4 mL (1 : 1 in 45% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

In vitro and animal studies

A wide range of pharmacological activities have been documented for German chamomile, including antibacterial, anti-inflammatory, antispasmodic, anti-ulcer, antiviral and hypouraemic activities.

Anti-inflammatory and anti-allergic activity

Anti-allergic and anti-inflammatory activities\(^{2,3}\) are well documented for German chamomile. The azulene components of the volatile oil are thought to contribute by inhibiting histamine release and they have been reported to prevent allergic seizures in guinea-pigs.\(^{2}\) Aqueous alcoholic extracts inhibited 5-lipoxygenase and cyclooxygenase activity, and oxidation of arachidonic acid, and a supercritical carbon dioxide extract had IC\(_{50}\) values of 6–25 µg/mL for these three activities.\(^{G52}\) The active compounds identified included apigenin, chamazulene, \(\text{cis}-\text{en}-\text{yn spiroether}\) and (−)-α-bisabolol.\(^{G52}\) Matricin, the precursor to chamazulene, is reported to be a more effective anti-inflammatory agent than chamazulene.\(^{2,4}\) Anti-inflammatory activity has also been documented for the sesquiterpene bisabolol compounds, with greatest activity reported for (−)-α-bisabolol,\(^{2,5}\) and for \(\text{cis}\)-spiroether.\(^{2}\) Anti-inflammatory activity (rat paw carrageenan test) has also been documented for a \(\text{cis}\)-spiroether against dextran induced oedema; no activity was observed against oedema induced by serotonin, histamine or bradykinin.\(^{6}\) In addition, flavonoids are known to possess anti-inflammatory activity.

Sedative activity

Apigenin competitively inhibited binding of flunitrazepam to the central benzodiazepine receptor, but lacked activity at other receptors, including muscarinic, \(\alpha_1\)-adrenoreceptor and GABA\(_A\).\(^{G52}\) High-performance liquid chromatography (HPLC) fractions of a methanol extract displaced flunitrazepam from its receptors in rat cerebellum membranes and muscimol from GABA receptors in rat cortical membranes, due to the presence of GABA in the fractions. Prolongation of hexobarbital-induced sleeping time and reduction in activity of mice have been documented.\(^{G52}\)

Anti-ulcerogenic activity

Anti-ulcerogenic activity in rats has been reported for (−)-α-bisabolol; the development of ulcers induced by indometacin, stress or ethanol was inhibited.\(^{2,7}\)
Antimicrobial and antiviral activities
German chamomile oil has been reported to have antifungal activity and antibacterial activity against Gram–positive bacteria. The coumarin herniarin has antibacterial and antifungal activities in the presence of UV light. Antibacterial activity has been documented for the coumarin constituents. An ethanolic extract of the entire plant has been reported to inhibit the growth of poliovirus and herpesvirus.

Antispasmodic activity
Antispasmodic activity on the isolated guinea–pig ileum has been documented for the flavonoid and bisabolol constituents. Greatest activity was exhibited by the flavonoids, especially apigenin which was found to be more than three times as potent as papaverine. (−)-α-Bisabolol activity was found to be comparable to that of papaverine, while the total volatile oil was considerably less active. Smooth muscle relaxant properties have also been documented for a cis-spiroether.

Enhancement of uterine tone in the guinea–pig and rabbit has been reported for an aqueous extract at a concentration of 1–2 mg extract/cm$^3$.

Other activities
High molecular weight polysaccharides with immunostimulating activity have been isolated from German chamomile. The oil has been reported to increase bile secretion and concentration of cholesterol in the bile, following the administration of 0.1 mL/kg by mouth to cats and dogs. A dose of 0.2 mL/kg was stated to exhibit hypotensive, and cardiac and respiratory depressant properties.

The ability of the volatile oil to regenerate liver tissue in partially hepatectomised rats has been attributed to the azulene constituents.

The volatile oil has been documented to reduce the serum concentration of urea in rabbits with experimentally induced uraemic conditions.

Clinical studies
German chamomile extracts have been reported to exhibit anti–inflammatory, anti peptic and antispasmodic activities on the human stomach and duo denum.

Anti–inflammatory and wound–healing effects
Clinical studies investigating the anti–inflammatory effects and wound–healing properties of German chamomile preparations have been
A summary of this information is given below.

A cream containing German chamomile extract was reported to have effects equivalent to 0.25% hydrocortisone, and superior effects to 0.1% diflucortolone and 5% bufexamax in inflammatory dermatoses, as assessed in 161 patients. Studies involving healthy volunteers who received German chamomile preparations have reported that German chamomile ointment was superior to 0.1% hydrocortisone acetate in dermatitis, and German chamomile cream (20 mg/g) reduced visual sores and redness of skin in an adhesive–tape stripping test. A randomised, double-blind trial involving 25 participants indicated that a cream containing an aqueous alcoholic extract of German chamomile was more effective than hydrocortisone against UVB-induced erythema.

In an open study involving 98 patients with cancer, an extract preparation (containing 50 mg α-bisabolol and 150–300 mg apigenin–7–glucoside/100 g) used three times daily was reported to reduce oral mucositis caused either by irradiation or chemotherapy. However, a double-blind, placebo-controlled trial involving 164 patients showed that a mouthwash containing German chamomile did not decrease 5-fluorouracil-induced stomatitis.

German chamomile has been reported to be an effective treatment for mucosal infections. Diluted extracts administered as a mouthwash 5 or 6 times daily provided cooling and astringent effects.\(^2\)

A cream containing German chamomile has produced additional anti-inflammatory, slight anaesthetic, cooling and deodorant effects in patients with cutaneous leg infections, when used in conjunction with existing treatment.\(^2\)

The healing effects of German chamomile ointment and dexapanthenol 5% cream administered for six days were reported to have comparable effects in a study involving 147 female patients who underwent episiotomy during childbirth. A standardised extract (50 mg α-bisabolol and 3 mg chamazulene/100 g) significantly decreased weeping wound area and drying of wound in 14 patients following removal of tattoos. An ointment preparation improved haemorrhage, itching, burning and oozing due to haemorrhoids in a study involving 120 patients.

**Sedative effects**

Sedative effects have been documented for German chamomile. Oral administration of a German chamomile extract was reported to induce a deep sleep in 10 of 12 patients undergoing cardiac catheterisation.\(^2\)
Side-effects, Toxicity

Reports of allergic reactions to chamomile are common, although in the majority of cases the plant species is not specified.\(^{(15)}\) Two reports of anaphylactic reactions to chamomile (species unspecified) have been documented\(^{(16,17)}\) and in both cases the individuals concerned had an existing hypersensitivity to ragweed (member of Asteraceae/Compositae). The symptoms they experienced included abdominal cramps, thickness of the tongue and a tight sensation in the throat,\(^{(17)}\) angioedema of the lips and eyes, diffuse pruritus, a full sensation of the ears, generalised urticaria, upper airway obstruction, and pharyngeal oedema.\(^{(16)}\) Both patients made a full recovery following medical treatment. Patients with an existing hypersensitivity to German chamomile have demonstrated cross-sensitivities to other members of the family Asteraceae/Compositae\(^{(18,G51)}\), and also to celery (family Umbelliferae).\(^{(G51)}\)

Allergic skin reactions have been documented following external contact with German chamomile.\(^{(2,19,G51)}\) Consumption of chamomile tea may exacerbate existing allergic conditions and the use of a chamomile enema has been documented to cause asthma and urticaria.\(^{(G51)}\)

The allergenic properties documented for chamomile have been attributed to anthecotulid, a sesquiterpene lactone present in low concentrations,\(^{(15)}\) and to matricarin, a proazulene which has produced positive patch tests in patients with an existing sesquiterpene lactone hypersensitivity.\(^{(G51)}\)

Sesquiterpene lactones have been implicated in the allergenic activity of many plants, especially those belonging to the Asteraceae/Compositae family (see Feverfew). The prerequisite for allergenic activity is thought to be an exocyclic α-methylene group.\(^{(20)}\)

The flowerheads contain anthemic acid, which is reported to act as an emetic in large doses.\(^{(G22)}\)

The acute toxicity of chamomile oil (German and Roman) is reported to be low.\(^{(21)}\) Oral and dermal LD\(_{50}\) values in rabbits have been documented as greater than 5 g/kg,\(^{(21)}\) and the application of undiluted oil to the hairless backs of mice, to rabbit skin, and to human skin was not found to produce any observable irritation.\(^{(21)}\) An LD\(_{50}\) value (mouse, by mouth) for German chamomile oil has been documented as 2.5 mL/kg.\(^{(13)}\) The acute oral toxicity of (−)-α-bisabolol in mice and rats is reported to be low at approximately 15 mL/kg.\(^{(22)}\) The subacute oral toxicity of (−)-α-bisabolol has been
estimated to be between 1.0 and 2.0 mL/kg in rats and dogs.\(^{22}\) An LD\(_{50}\) value (mouse, intraperitoneal injection) for \(cis\)-spiroether has been stated as 670 mg/kg.\(^{6}\)
Contra-indications, Warnings

In view of the documented allergic reactions and cross-sensitivities, German chamomile should be avoided by individuals with a known hypersensitivity to any members of the Asteraceae/Compositae family. In addition, German chamomile may precipitate an allergic reaction or exacerbate existing symptoms in susceptible individuals (e.g. asthmatics). Excessive doses may interfere with existing anti coagulant therapy, because of the coumarin constituents.

The use of chamomile preparations for teething babies is not recommended.

Pregnancy and lactation

German chamomile is reputed to affect the menstrual cycle\(^{(G30)}\) and extracts are reported to be uterotonic.\(^{(2,11)}\) Teratogenicity studies in rats, rabbits and dogs have been documented for \((-\)-\(\alpha\)-bisabolol, with the oral toxic dose stated as 1–3 mL/kg.\(^{(22)}\) A dose of 3 mL/kg was found to increase the number of fetuses reabsorbed and reduce the body weight of live offspring.\(^{(22)}\) \((-\)-\(\alpha\)-Bisabolol administered orally (250 and 500 mg/kg) to pregnant rats has been reported to have no effect on the fetus.\(^{(1)}\) In view of the documented information, the excessive use of chamomile during pregnancy and lactation should be avoided.
Pharmaceutical Comment

The chemistry of German chamomile, especially of the volatile oil component, is well documented and is similar to that of Roman chamomile.\(^{(24)}\)

Pharmacological activity is associated with the flavonoid and volatile oil fractions. A wide range of pharmacological actions have been documented (e.g. anti–inflammatory and antispasmodic activities) and many of these support the reputed herbal uses\(^{(23,24)}\). A small number of studies in patients and healthy volunteers have reported anti–inflammatory, wound healing and sedative effects. Toxicity studies to date have indicated chamomile to be of low toxicity, although allergic reactions are documented.
References


Chamomile, Roman
Species (Family)

*Chamaemelum nobile* (L.) All. (Asteraceae/Compositae)
Synonym(s)

_Anthemis nobilis_ L.
Part(s) Used

Flowerhead
Pharmacopoeial and Other Monographs

BHC 1992(G6)
BHP 1996(G9)
BP 2002(G71)
Martindale 33rd edition(G67)
PDR for Herbal Medicines 2nd edition (English Chamomile)(G36)
Ph Eur 2004(G72)
USP26/NF21(G73)
Legal Category (Licensed Products)

GSL$^{(G^{37})}$
Constituents
See General References G2 G6 G22 G41 G48 G64.

**Coumarins**
Scopoletin–7–glucoside.

**Flavonoids**
Apigenin, luteolin, constituents in, quercetin and their glycosides (e.g. apiin, luteolin–7–glucoside, and rutin).

**Volatile oils**
0.4–1.75%. Angelic and tiglic acid esters (85%);\(^{(1)}\) others include 1,8–cineole, \textit{l-}trans–pinocarveol, \textit{l-}trans–pinocarvone, chamazulene, farnesol, nerolidol; germacranolide–type sesquiterpene lactones (0.6%);\(^{(2)}\) including nobilin, 3–epinobilin, 1,10–epoxynobilin, 3–dehydrobenzoin; various alcohols including amyl and isobutyl alcohols, anthemol.\(^{(1–4)}\) Chamazulene is formed from a natural precursor during steam distillation of the oil, and varies in yield depending on the origin and the age of flowers.\(^{(1)}\)

**Other constituents**
Anthemic acid (bitter), phenolic and fatty acids, phytosterol, choline and inositol.
Food Use

Roman chamomile is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that Roman chamomile can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\textsuperscript{(G16)} Chamomile is commonly used as an ingredient of herbal teas. In the USA, Roman chamomile is listed as GRAS (Generally Recognised As Safe).\textsuperscript{(G41)}
Herbal Use

Roman chamomile is stated to possess carminative, anti-emetic, antispasmodic, and sedative properties. It has been used for dyspepsia, nausea and vomiting, anorexia, vomiting of pregnancy, dysmenorrhoea, and specifically for flatulent dyspepsia associated with mental stress. (G2 G6 G7 G8 G64)
Dosage

*Dried flowerheads*
1–4 g or by infusion three times daily.\(^{(G7)}\)

*Liquid extract*
1–4 mL (1 : 1 in 70% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

German and Roman chamomile possess similar pharmacological activities (see Chamomile, German for a fuller description of documented pharmacological actions).

**In vitro and animal studies**

Few studies have been documented specifically for Roman chamomile. The azulene compounds are reported to possess anti–allergic and anti–inflammatory properties; their mechanism of action is thought to involve inhibition of histamine release (see Chamomile, German). The volatile oil has been documented as having anti–inflammatory activity (carrageenan rat paw oedema test), and antidiuretic and sedative effects following intraperitoneal administration of doses up to 350 mg/kg body weight to rats.(5)

The azulenes have been reported to stimulate liver regeneration following oral, but not subcutaneous, administration.

The sesquiterpenoids nobilin, 1,10–epoxynobilin and 3–dehydronobilin have demonstrated *in vitro* antitumour activity against human cells.(1) The concentration of hydroxyisonobilin required for cytotoxic activity is reported to be low enough to warrant further investigations (ED$_{50}$ 0.56 μg/mL versus HeLa; ED$_{50}$ 1.23 μg/mL versus KB; arbitrary acceptable test level 4 μg/mL).
Side-effects, Toxicity

Instances of allergic and anaphylactic reactions to chamomile have been documented (see Chamomile, German) The allergenic principles in chamomile are thought to be the sesquiterpene lactones.\(^1\) Roman chamomile yields nobilin, a sesquiterpene lactone that is reported to be potentially allergenic.\(^1\) However, Roman chamomile oil has also been reported to be non-irritant and non-sensitising to human skin.\(^2\) Animal studies have indicated the oil to be either mildly or non-irritant, and to lack any phototoxic effects.\(^2\)

Large doses of Roman chamomile are stated to act as an emetic\(^{G44}\) and this has been attributed to the anthemic acid content.\(^6\)

The acute toxicity of Roman chamomile in animals is reported to be relatively low.\(^1\) Acute LD\(_{50}\) values in rabbits (dermal) and rats (by mouth) have been stated to exceed 5 g/kg.\(^2\)
Contra-indications, Warnings

In view of the documented allergic reactions and cross-sensitivities (see Chamomile, German), Roman chamomile should be avoided by individuals with a known hypersensitivity to any members of the Asteraceae/Compositae family. In addition, Roman chamomile may precipitate an allergic reaction or exacerbate existing symptoms in susceptible individuals (e.g. asthmatics). Excessive doses may interfere with anticoagulant therapy because of the coumarin constituents.

The use of chamomile preparations in teething babies is not recommended.

Pregnancy and lactation

Roman chamomile is reputed to be an abortifacient and to affect the menstrual cycle. In view of this and the potential for allergic reactions, the excessive use of chamomile during pregnancy and lactation should be avoided.
Pharmaceutical Comment

The chemistry of Roman chamomile, particularly of the volatile oil, is well documented and is similar to that of German chamomile. Limited pharmacological data are available for Roman chamomile, although many actions have been reported for German chamomile. In view of the similar chemical compositions, many of the activities described for German chamomile are thought to be applicable to Roman chamomile and thus support the traditional herbal uses. Roman chamomile is stated to be of low toxicity, although allergic reactions (mainly contact dermatitis) have been reported.
References


Species (Family)

*Larrea tridentata* (DC.) Coville (Zygophyllaceae)
Creosote Bush. *L. tridentata* (south-western USA and northern Mexico) is now regarded as a separate species to *Larrea divaricata* Gav. (north-western Argentina). \(^1\)
Part(s) Used
Herb
Pharmacopoeial and Other Monographs

Martindale 33rd edition (G67)
Legal Category (Licensed Products)

Chaparral is not included in the GSL.(G37)
Constituents
See General Reference G22.

Amino acids
Arginine, aspartine, cystine, glutamic acid, glycine, isoleucine, leucine, phenylalanine, tryptophan, tyrosine and valine.

Flavonoids
More than 20 different compounds reported, including isorhamnetin, kaempferol and quercetin and their glycosidic and ether derivatives; gossypetin, herbacetin, and their acetate derivatives;\(^{(1-7)}\) two C-glucosyl flavones.

Lignans
Major constituent nordihydroguaiaretic acid (NDGA) (up to 1.84%), norisoguaiacin, dihydro guaiaretic acid, partially demethylated dihydroguaiaretic acid, 3′- demethoxyisoguaiacin.\(^{(8-10)}\)

Resins
20%. Phenolic constituents on external leaf surfaces of *L. divaricata* and *L. tridentata* are reported to be identical, containing a number of flavone and flavonol glycosides, and two lignans (including NDGA).\(^{(5)}\)

Volatile oils
Many identified terpene components include calamene, eudesmol, limonene, α- and β-pinene, and 2-rossalene.\(^{(11)}\)

Other constituents
Two pentacyclic triterpenes,\(^{(12)}\) saponins.

Other plant parts
A cytotoxic naphthoquinone derivative, larreantin, has been isolated from the roots.\(^{(13)}\)
Food Use

Chaparral is not used in foods, although a related species, *Larrea mexicana* Moric., also termed creosote bush, is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that creosote bush can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\(^{G16}\)

In the USA, NDGA is no longer permitted to be used as an antioxidant in foods following the results of toxicity studies in animals (see Side-effects, Toxicity).
Herbal Use

Chaparral has been used for the treatment of arthritis, cancer, venereal disease, tuberculosis, bowel cramps, rheumatism and colds.\(^{(G60)}\)
Dosage

None documented.
Pharmacological Actions

In vitro and animal studies

Amoebicidal action against *Entamoeba histolytica* has been reported for a chaparral extract (0.01%).\(^\text{(14)}\) This action may be attributable to the lignan constituents, which are documented as both amoebicidal and fungicidal.\(^\text{(9)}\) NDGA has been reported to have antimicrobial activity against a number of organisms including *Penicillium* spp., *Salmonella* spp., *Streptococcus* spp., *Staphylococcus aureus*, *Bacillus subtilis*, *Pseudomonas aeruginosa* and various other pathogens and moulds.\(^\text{(8,15)}\)

NDGA is an antioxidant, and has been documented to cause inhibition of hepatic microsomal enzyme function.\(^\text{(15–17)}\)

Clinical studies

Medical interest in chaparral increased following claims that an aqueous infusion of the herb had caused the regression of a malignant melanoma in the cheek of an 85-year-old man.\(^\text{(18)}\) However, results of a subsequent study that investigated the antitumour action of chaparral, as a tea, were inconclusive.\(^\text{(G60)}\)
Side-effects, Toxicity

Acute hepatitis has been associated with chaparral ingestion.\(^{19–21}\) Contact dermatitis to chaparral has been reported.\(^{22,23}\) Chaparral-induced toxic hepatitis has been reported for two patients in different parts of the USA. The adverse effects were attributed to ingestion of a herbal nutritional supplement derived from the leaves of chaparral. Five cases of serious poisoning in the USA and another three in Canada have been linked to chaparral-containing products.\(^{20,24}\) Some patients have developed irreversible reno-hepatic failure. Early investigations into the toxicity of NDGA concluded it to be low.\(^{15}\) NDGA has been administered to humans, by intramuscular injection, in doses of up to 400 mg/kg body weight for 5–6 months, with little or no toxicity reported.\(^{15}\) Documented oral LD\(_{50}\) values for NDGA include 4 g/kg (mouse), 5.5 g/kg (rat) and 830 mg/kg (guinea-pig).\(^{15}\) Results of chronic feeding studies (two years, 0.25–1.0% of diet) in rats and mice reported no abnormalities in histological tests of the liver, spleen and kidney. Inflammatory caecal lesions and slight cystic enlargement of lymph nodes near the caecum were observed in rats at the 0.5% feeding level. At this point NDGA was considered to be safe for food use. However, two later studies in rats (using NDGA at up to 3% of the diet) reported the development of cortical and medullary cysts in the kidney.\(^{15}\) On the basis of these findings, NDGA was removed from GRAS (Generally Recognised As Safe) status in the USA and is no longer permitted to be used as an antioxidant in foods.\(^{15}\)
Contra–indications, Warnings

In view of the reports of acute hepatitis associated with chaparral ingestion, and the uncertainty regarding NDGA toxicity, consumption should be avoided. Excessive doses may interfere with monoamine oxidase inhibitor (MAOI) therapy, because of the documented amino acid constituents.

Pregnancy and lactation

In vitro utero activity has been documented for chaparral.\(^{(G30)}\) In view of the concerns regarding toxicity, chaparral should not be ingested during pregnancy or lactation.
Pharmaceutical Comment

The chemistry of chaparral is well studied and extensive literature has been published on the principal lignan component, NDGA. However, little documented evidence is available to justify the herbal uses of chaparral. In view of the concerns over the hepatic toxicity, the use of chaparral as a herbal remedy cannot be recommended.
References

See also General References G16 G18 G20 G22 G30 G31 G32 G36 G37 G43 G60.


Cinnamon
Species (Family)

i. *Cinnamomum zeylanicum* Bl. (Lauraceae)

ii. *Cinnamomum loureirii* Nees

iii. *Cinnamomum burmanii* (Nees) Bl.
Synonym(s)

i. Ceylon Cinnamon, *Cinnamomum verum* J.S. Presl., True Cinnamon

ii. *Cinnamomum obtusifolium* Nees var. *loureirii* Perr. & Eb., Saigon Cassia, Saigon Cinnamon

iii. Batavia Cassia, Batavia Cinnamon, Padang-Cassia, Panang Cinnamon
Part(s) Used

Inner bark
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)

BP 2002\(^{(G71)}\)

Complete German Commission E\(^{(G3)}\)

Martindale 33rd edition\(^{(G67)}\)

Ph Eur 2004\(^{(G72)}\)

WHO volume 1 1999\(^{(G63)}\)
Legal Category (Licensed Products)

GSL\textsuperscript{G37}
Constituents

See General References G2 G41 G48 G58 G59 G62 G64.

Tannins

Condensed.

Volatile oils

Up to 4%. Cinnamaldehyde (60–75%), benzaldehyde and cuminaldehyde; phenols (4–10%) including eugenol, and methyl eugenol, pinene, phellandrene, cymeme and caryophyllene (hydrocarbons), eugenol acetate, cinnamyl acetate and benzyl benzoate (esters), linalool (an alcohol). Of the various types of cinnamon bark the oil of C. zeylanicum is stated to contain the highest amount of eugenol. Cinnamon oil differs from the closely related cassia oil in that the latter is reported to be devoid of eugenol, monoterpenoids and sesquiterpenoids (see Cassia).

Other constituents

Calcium oxalate, cinnzeylanin, cinnzeylanol, coumarin, gum, mucilage, resins and sugars.

Other plant parts

Cinnamon leaf oil contains much higher concentrations of eugenol, from 80 to 96% depending on the species. A cinnamon leaf oil of Chinese origin, Cinnamomum japonicum Sieb., contains a high concentration of safrole (60%) and only about 3% eugenol.
Cinnamon is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that cinnamon can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\(^\text{G16}\) It is commonly used as a spice in cooking, although at levels much less than the stated therapeutic doses. The acceptable daily intake of cinnamaldehyde has been temporarily estimated as 700 μg/kg body weight.\(^\text{G45}\) In the USA, cinnamon is listed as GRAS (Generally Recognised As Safe).\(^\text{G41}\)
Herbal Use

Cinnamon is stated to possess antispasmodic, carminative, orexigenic, antidiarrhoeal, antimicrobial, refrigerant and anthelmintic properties. It has been used for anorexia, intestinal colic, infantile diarrhoea, common cold, influenza, and specifically for flatulent colic, and dyspepsia with nausea.\(^{(G7)}\) Cinnamon bark is also stated to be astringent, and cinnamon oil is reported to possess carminative and antiseptic properties.\(^{(G2 G41 G64)}\)
**Dosage**

*Dried bark*
0.5–1.0 g as infusion three times daily.\(^{(G7)}\)

*Liquid extract*
0.5–1.0 mL (1 : 1 in 70% alcohol) three times daily.\(^{(G7)}\)

*Tincture of Cinnamon*
(BPC 1949) 2–4 mL.
Pharmacological Actions

In vitro and animal studies

Cinnamon oil has antifungal, antiviral, bactericidal and larvicidal properties.\(^{(G4)}\) A carbon dioxide extract of cinnamon bark (0.1%) has been documented to suppress completely the growth of numerous microorganisms including *Escherichia coli*, *Staphylococcus aureus* and *Candida albicans*.\(^{(G4)}\) (See Cassia for details of the many pharmacological actions documented for cinnamaldehyde and cinnamomi cortex (cinnamon bark).)

Antiseptic and anaesthetic properties have been documented for eugenol\(^{(1)}\) and two insecticidal compounds, cinnzeylanin and cinnzeylanol, have been isolated.\(^{(G4)}\) Tannins are known to possess astringent properties.

Weak tumour-promoting activity on the mouse skin and weak cytotoxic activity against HeLa cells has been documented for eugenol.\(^{(G4)}\)
Side-effects, Toxicity

None documented for cinnamon bark. Cinnamon oil contains cinnamaldehyde, an irritant and sensitising principle.\(^{(G58)}\) The dermal LD\(_{50}\) of the oil is reported to be 690 mg/kg body weight (see Cassia). The accepted daily intake of eugenol is up to 2.5 mg/kg.\(^{(G45)}\)
Contra-indications, Warnings

Contact with cinnamon bark or oil may cause an allergic reaction. (G51) Cinnamon oil is stated to be a dermal and mucous membrane irritant, and a dermal sensitiser. (G58) It is a hazardous oil and should not be used on the skin. (G58) The oil should not be taken internally.

**Pregnancy and lactation**

There are no known problems with the use of cinnamon during pregnancy and lactation, provided that doses do not greatly exceed the amounts used in foods.
Pharmaceutical Comment

The reputed antimicrobial, antiseptic, anthelmintic, carminative and antispasmodic properties of cinnamon are probably attributable to the volatile oil. The astringent properties of tannins may account for the claimed antidiarrhoeal action. Cinnamon should not be used in amounts greatly exceeding those used in foods.

Species (Family)

*Galium aparine* L. (Rubiaceae)
Synonym(s)
Cleavers, Galium, Goosegrass
Part(s) Used
Herb
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents
See General References G6 G34 G40 G44 G48 G64.

**Acids**
Caffeic acid, \( p \)-coumaric acid, gallic acid, \( p \)-hydroxybenzoic acid, salicylic acid and citric acid.\(^{(1)}\)

**Coumarins**
Unspecified. Scopoletin and umbelliferone reported for related species *Galium cruciata* and *Galium tauricum*.\(^{(2)}\)

**Iridoids**
Asperuloside (rubichloric acid), monotropein.\(^{(3,4)}\)

**Tannins**
Unspecified;\(^{(5)}\) gallic acid is usually associated with hydrolysable tannins.

**Other constituents**
Alkanes (\( C_{19} \)–\( C_{31} \)),\(^{(4)}\) flavonoids.

**Other plant parts**
Anthraquinones have been documented for the roots, but not for the aerial parts.\(^{(1)}\)
Food Use

Clivers is not used in foods.
Herbal Use

Clivers is stated to possess diuretic and mild astringent properties. It has been used for dysuria, lymphadenitis, psoriasis, and specifically for enlarged lymph nodes. (G6 G7 G8 G64)
Dosage

*Dried herb*
2–4 g or by infusion three times daily.\(^{(G6 \ G7)}\)

*Liquid extract*
2–4 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G6 \ G7)}\)

*Expressed juice*
3–15 mL three times daily.\(^{(G6 \ G7)}\)
Pharmacological Actions

*In vitro and animal studies*

None documented for clivers. Asperuloside and monotropein have been reported to elicit a mild laxative action in mice.\(^6\) The action was stated to be approximately 15 times less potent than that of senna, and of shorter duration.

*Clinical studies*

None documented. Tannins are known to possess astringent activities.
Side-effects, Toxicity

None documented.
Contra–indications, Warnings

It has been stated that diabetics should only use the expressed juice with caution\(^{(G34)}\) although no pharmacological data were located to support this statement.

**Pregnancy and lactation**
In view of the lack of pharmacological and toxicological information, the use of clivers during pregnancy should be avoided.
Pharmaceutical Comment

Limited chemical information is available for clivers. No scientific evidence was found to support the herbal uses, although documented tannin constituents may account for the reputed mild astringent action. In view of the paucity of toxicity data, excessive use of clivers should be avoided.
See also General References G6 G9 G34 G36 G37 G40 G43 G48 G49 G64.

Clove
Species (Family)

*Syzygium aromaticum* (L.) Merr. & Perry (Myrtaceae)
Synonym(s)

Part(s) Used
Clove (dried flowerbud), leaf, stem
Pharmacopoeial and Other Monographs

- BHP 1996$^{(G9)}$
- BP 2002$^{(G71)}$
- Complete German Commission E$^{(G3)}$
- Martindale 33rd edition$^{(G67)}$
- PDR for Herbal Medicines 2nd edition$^{(G36)}$
- Ph Eur 2004$^{(G72)}$
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents

See General References G2 G22 G41 G48 G58 G64.

**Volatile oils**
Clove bud oil (15–18%) containing eugenol (80–90%), eugenyl acetate (2–27%), β-caryophyllene (5–12%). Others include methyl salicylate, methyl eugenol, benzaldehyde, methyl amyl ketone and α-ylangene.

Leaf oil (2%) containing eugenol 82–88%.

Stem oil (4–6%) with eugenol 90–95%. A more comprehensive listing is provided elsewhere. (G22)

**Other constituents**
Campesterol, carbohydrates, kaempferol, lipids, oleanolic acid, rhamnetin, sitosterol, stigmasterol and vitamins.
Food Use

Clove is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that clove can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\(^{(G16)}\) Clove is commonly used in cooking, and as a flavouring agent in food products. In the USA, clove is listed as GRAS (Generally Recognised As Safe).\(^{(G41)}\)
Herbal Use

Clove has been traditionally used as a carminative, anti-emetic, toothache remedy and counter-irritant.\(^{(G2 \ G41 \ G64)}\)

Clove oil is stated to be a carminative, occasionally used in the treatment of flatulent colic\(^{(G54)}\) and is commonly used topically for symptomatic relief of toothache.\(^{(G45)}\)
Dosage

*Clove*
120–300 mg.\(^{(G44)}\)

*Clove oil*
0.05–0.2 mL.\(^{(G44)}\)
Pharmacological Actions

*In vitro* and animal studies

The anodyne and mild antiseptic properties documented for clove oil have been attributed to eugenol.\(^{[G41]}\) Clove oil is stated to possess antihistaminic and antispasmodic properties.\(^{[G41]}\) Eugenol, eugenol acetate and methyl acetate are reported to exhibit trypsin–potentiating activity.\(^{[G41]}\)

Antibacterial, hypoglycaemic and potent CNS-depressant activities have been documented for *Syzygium cuminii* L., a related species cultivated in India.\(^{[1]}\)

**Clinical studies**

A tincture of cloves (15% in 70% alcohol) was effective in treating athlete’s foot.\(^{[G41]}\)
Side-effects, Toxicity

None documented for the bud, leaf or stem of cloves. Clove oil is stated to be a dermal and mucous membrane irritant; (G58) contact dermatitis, cheilitis, and stomatitis have been reported for clove oil. (G51) The irritant nature of the oil can be attributed to the eugenol content. Eugenol is also stated to have sensitising properties. (G51) An LD$_{50}$ (rat, by mouth) value for clove oil is stated as 2.65 g/kg body weight. (G22)

In humans, the accepted daily intake of eugenol is up to 2.5 mg/kg body weight. (G45)
Contra-indications, Warnings

None documented for the bud, leaf or stem. It is recommended that clove oil should be used with caution orally and should not be used on the skin.\(^{(G58)}\) Repeated application of clove oil as a toothache remedy may result in damage to the gingival tissue.\(^{(G45)}\) In view of the irritant nature of the volatile oil, concentrated clove oil is not suitable for internal use in large doses. Eugenol is a powerful inhibitor of platelet activity and it is recommended that caution be taken for patients on anticoagulant therapy.\(^{(G58)}\)

**Pregnancy and lactation**

There are no known problems with the use of clove during pregnancy or lactation, provided that doses taken do not greatly exceed the amounts used in foods.
Pharmaceutical Comment

The pharmacological properties documented for cloves are associated with the volatile oil, in particular with eugenol which has local anaesthetic action. Cloves should not be taken in doses greatly exceeding those used in foods and caution should be exerted in patients taking anticoagulant or anti platelet therapy.
References

See also General References G2 G3 G9 G15 G16 G22 G25 G31 G36 G37 G41 G43 G48 G51 G64.


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Species (Family)

*Cimicifuga racemosa* Nutt. (Ranunculaceae)
Synonym(s)
Actaea Racemosa Radix, Black Snakeroot, Cimicifuga, Macrotys Actaea
Part(s) Used

Rhizome, root
Pharmacopoeial and Other Monographs

BHC 1992\textsuperscript{G6}
BHP 1996\textsuperscript{G9}
BPC 1934\textsuperscript{G10}
Complete German Commission E\textsuperscript{G3}
Martindale 33rd edition\textsuperscript{G67}
Mills and Bone\textsuperscript{G50}
PDR for Herbal Medicines 2nd edition\textsuperscript{G36}
Legal Category (Licensed Products)

GSL$\text{G37}$
Constituents
See References 1 and General References G6 G22 G41 G64.

Alkaloids
Quinolizidine–type. N-Methylcytisine and other unidentified compounds.

Tannins
Type unspecified. Tannic and gallic acids are usually associated with hydrolysable tannins.

Terpenoids
Triterpene glycosides, principally the xylosides actein (aglycone: acetylacteol) and cimicifugoside (also known as cimigoside; aglycone: cimigenol),\(^{(2–8,G6)}\)
also 26–deoxycimicifugoside, cimi aceroside A, 27–deoxyactein,\(^{(9,10)}\)
cimiracemosides A–H,\(^{(9–11)}\) and cimicifugosides H-3, H-4 and H-6.\(^{(12)}\)

Other constituents
Acetic acid, butyric acid, formic acid, hydroxycinnamic acid esters of fukiic and piscidic acids (fukinolic acid, cimicifugic acids A, B, E, F), caffeic acid, ferulic acid,\(^{(13)}\) isoferulic acid, oleic acid, palmitic acid, salicylic acid, racemosin, formononetin,\(^{(3)}\) phytosterols, cimicifugin 15–20%, acteina (resinous mixture) and volatile oil.
In the USA, black cohosh is listed by the Food and Drugs Administration (FDA) as a ‘Herb of Undefined Safety’. Black cohosh is not used in foods.
Herbal Use

Black cohosh is stated to possess antirheumatic, antitussive, sedative and emmenagogue properties. It has been used for intercostal myalgia, sciatica, whooping cough, chorea, tinnitus, dysmenorrhoea, uterine colic, and specifically for muscular rheumatism and rheumatoid arthritis. (G6 G7 G8 G32 G64) Modern use of black cohosh is focused on its use in treating peri- and postmenopausal symptoms. (1,14,G50)
Dosage

**Dried rhizome/root**
40–200 mg daily.\(^{(G6)}\)

**Liquid extract**
Ethanolic extracts equivalent to 40 mg dried rhizome/root daily.\(^{(G3 G50)}\)

**Tincture**
0.4–2 mL (1 : 10 in 60% ethanol) daily.\(^{(G6)}\)

Several clinical trials of black cohosh have used a standardised black cohosh extract (Remifemin; each 20 mg tablet contained 1 mg triterpene glycosides, calculated as 27-deoxyactein) 40 mg twice daily for up to 24 weeks.\(^{(15–17)}\)
Pharmacological Actions

Several pharmacological activities, including hormonal, cardiovascular, circulatory and anti-inflammatory activities, have been documented for black cohosh and/or its constituents. The triterpene glycosides and flavonoids are considered to be the active components of black cohosh.\(^{18, G56}\)

**In vitro and animal studies**

**Hormonal activity**

A methanolic extract of the rhizome of black cohosh reduced the serum concentration of luteinising hormone (LH) in ovariectomised rats, and exhibited a binding affinity to oestrogen receptors in isolated rat uterus.\(^{2}\) In vivo, the activity of the methanolic extract was significantly reduced following enzymatic hydrolysis of glucosides present. Subsequent in vitro studies isolated three compounds with endocrine activity, including an isoflavone, formononetin. Formononetin was found to exhibit competitive oestrogen receptor activity, but did not cause a reduction in serum concentrations of LH.\(^{2}\) Recent research found that formonenetin could not be detected in commercial preparations of black cohosh, although other flavonoids were present.\(^{19}\)

In ovariectomised rats, administration of a lipophilic extract of black cohosh (140 mg by intraperitoneal injection for three days) led to a significant reduction in serum LH concentrations, compared with control \((p < 0.01)\), whereas no effect was observed with a hydrophilic extract (216 mg intraperitoneally for three days).\(^{16}\) Subsequent studies using fractions of the lipophilic extract demonstrated that constituents inhibited LH secretion and/or exhibited activity in an oestrogen receptor-binding assay.

In an in vitro study in oestrogen receptor–positive breast cancer cells, black cohosh extract did not stimulate cancer cell growth, i.e. it did not exhibit oestrogen–like effects, but at a concentration of 2.5 \(\mu g/mL\) led to a marked inhibition of breast cancer cell proliferation.\(^{20}\)

Oestrogenic activity has been documented in vitro for fukinolic acid, a hydroxycinnamic acid ester of fukiic acid, in oestrogen–dependent MCF-7 cells (a breast cancer cell line).\(^{13}\) Fukinolic acid at concentrations of \(5 \times 10^{-7}\) mol/L and \(5 \times 10^{-8}\) mol/L led to significantly increased cell proliferation (mean (standard deviation): +120 (6%) and +126 (5%), respectively), compared with control. These effects were reported to be equivalent to those of estradiol at \(10^{-10}\) mol/L. By contrast, in other in vitro studies, a methanol extract of black cohosh rhizomes and roots did not
demonstrate oestrogenic activity in several assays, including binding affinity for oestrogen receptors α and β, stimulation of pS2 mRNA expression in S30 cells (S30 is a subclone of an oestrogen receptor–negative breast cancer cell line), and induction of alkaline phosphatase in an oestrogen receptor–positive endometrial adenocarcinoma cell line.\(^{(21)}\)

**Other activities**

Anti–inflammatory and analgesic activity for constituents of black cohosh has been documented following *in vitro* and *in vivo* studies (mice and rats).

*In vitro*, caffeic acid, fukinolic acid and cimicifugic acids A, B, E and F inhibited the activity of neutrophil elastase in a concentration–dependent manner.\(^{(22)}\) (Raised plasma concentrations of neutrophil elastase are a typical feature of active inflammation. Neutrophil elastase contributes to the destruction of basement membranes during inflammation.) Caffeic acid inhibited the enzyme with an IC\(_{50}\) of 16 μg/mL (93 μmol/L), whereas fukinolic acid had an IC\(_{50}\) of 0.1 μg/mL (0.23 μmol/L), relative to controls. Of the cimicifugic acids, A and B were the strongest inhibitors of the enzyme, with IC\(_{50}\) values of 2.2 μmol/L and 11.4 μmol/L, respectively.

Compared with controls, a methanol extract, a butanol–soluble fraction and a water–soluble fraction obtained from *Cimicifuga* rhizome inhibited carrageenan–induced rat paw oedema by 73–76%, 80–84% and 46–54%, respectively, compared with controls, 30–120 minutes after injection of 0.1 mL carrageenan 1%.\(^{(23)}\) The same fractions (100 mg/kg by intraperitoneal injection), compared with controls, demonstrated analgesic activity determined by significant reductions in acetic acid–induced writhing in mice, and the methanol extract and butanol–soluble fraction also displayed analgesic activity in a tail–flick test (demonstrated by increased latency time upon infrared light exposure). All three fractions (40 μg/mL) inhibited bradykinin and histamine receptor–mediated contractions of guinea–pig ileum, and all inhibited lipopolysaccharide–induced 6–keto prostaglandin F\(_{1α}\) (PGF\(_{1α}\)) formation in macrophages. Incubation of macrophages with lipopolysaccharide and the water–soluble fraction 10 μg/mL almost completely blocked (99% inhibition) lipopolysaccharide–induced 6–keto-PGF\(_{1α}\) formation. Lipopolysaccharide–induced 6–keto-PGF\(_{1α}\) formation in macrophages is related to selective expression of cyclooxygenase 2 (COX-2). The inhibitory effects of fractions of *Cimicifuga* rhizome in this model, and their inhibitory effects on bradykinin and histamine receptor–mediated reactions are possible mechanisms for the observed anti–inflammatory and analgesic activities.
In vitro studies using rat aortic strips have investigated the vasoactive effects of constituents of *Cimicifuga* species.\(^{(24)}\) Cimicifugic acid D and fukinolic acid \((3 \times 10^{-4} \text{ mol/L})\) caused a sustained relaxation of aortic strips precontracted with noradrenaline (norepinephrine) in preparations with or without endothelium. By contrast, cimicifugic acid C inversely caused a weak contraction, and fukiic acid and cimicifugic acids A, B and E showed no vasoactivity at the concentration tested.

A resinous component, termed acteina, has exhibited a hypotensive action in both unanaesthetised rabbits and anaesthetised cats. The effect in unanaesthetised dogs was found to be inconsistent.\(^{(25)}\) An effective dose of acteina \(1 \text{ mg/kg body weight}\) was recorded, with maximum hypotension attained using \(10 \text{ mg/kg}\). It was stated that acteina may act by an effect on the vasomotor centres.

Triterpene compounds in black cohosh have been shown to possess hypocholesterolaemic activity *in vivo*, and an inhibitory effect on phytohaemagglutin–induced proliferative response *in vitro*. These activities were thought to be linked to molecular characteristics between the identified triterpenes and intermediates in cholesterol biosynthesis.\(^{(26)}\)

Triterpenoid constituents from several *Cimicifuga* species, including actein from *C. racemosa*, have been investigated for antimalarial activity against *Plasmodium falciparum in vitro*.\(^{(27)}\) Cimicifugoside (isolated from *C. simplex*) and actein were among the compounds with potent antimalarial activity \((\text{EC}_{50} 5.0 \text{ μmol/L} \text{ and } 10.0 \text{ μmol/L}, \text{ respectively})\), although activity was 2- to 3–fold less than that of positive controls (quinine, chloroquine and pyrimethamine).

The root of a related species, *Cimicifuga dahurica*, has been reported to exhibit antibacterial activity towards Gram–positive (*Bacillus subtilis*, *Mycobacterium smegmatis*, *Staphylococcus aureus*), and Gram–negative (*Escherichia coli*, *Shigella flexneri*, *Shigella sonnei*) organisms.\(^{(28)}\)

In ovariectomised rats, ethyl acetate–soluble fractions from the rhizome of the related species *Cimicifuga heracleifolia* and *C. foetida* administered orally at doses of \(100 \text{ mg/kg/day}\) for 42 days led to a significant increase in bone mineral density of the lumbar spine, compared with that in untreated ovariectomised control rats.\(^{(29)}\)

**Clinical studies**

Clinical trials of extracts of black cohosh have investigated mainly its effects in women with peri- and/or postmenopausal symptoms.
Menopausal symptoms

In a randomised, double-blind, placebo-controlled trial, 80 women (mean (standard deviation) age: 51.2 (3.1) years) with menopausal symptoms received standardised black cohosh extract (Remifemin; each 20-mg tablet contained 1 mg triterpene glycosides, calculated as 27-deoxyactein) 40 mg twice daily, conjugated oestrogens 0.625mg daily, or placebo, for 12 weeks. At the end of the study, somatic and psychological symptoms, measured by the Kupperman Menopausal Index and the Hamilton Anxiety Scale, had improved significantly only in women who received black cohosh, compared with those who received oestrogen or placebo. Similarly, a significant increase in proliferation of vaginal epithelium was noted only in the black cohosh group. However, there is a view that the dose of oestrogen used in the study was too low to provide a useful comparison.

A randomised, double-blind, placebo-controlled trial involving 85 women with a history of breast cancer assessed the effects of black cohosh (no details of extract provided) one tablet twice daily for 60 days on the frequency and intensity of hot flushes. Participants were stratified according to whether or not they were using tamoxifen. Both treatment and placebo groups reported decreases in the number and intensity of hot flushes, compared with baseline values. There were no statistically significant differences between the two groups, and subgroup analysis of tamoxifen users and non-users did not reveal any statistically significant differences. Changes in other parameters measured during the study (other menopausal symptoms, serum concentrations of follicle-stimulating hormone (FSH) and luteinising hormone (LH)) were also not statistically significant between groups.

In a placebo-controlled study involving 110 women with menopausal symptoms, black cohosh extract (Remifemin) two tablets daily for two months significantly reduced serum LH concentrations, compared with placebo ($p < 0.05$). There was no significant difference between the two groups with respect to serum FSH concentrations.

Most other studies of black cohosh involving women with menopausal symptoms have an open and/or uncontrolled design and, therefore, do not provide an unbiased assessment of efficacy. Generally, these studies report significant improvements in menopausal symptoms, compared with baseline values, after at least four weeks’ treatment. Some studies involved administration of black cohosh extract for up to 12 weeks. These studies have been summarised elsewhere. Details of two of these studies are provided below.

In an open study, 60 women with at least one intact ovary who had
undergone hysterectomy and who were experiencing menopausal symptoms were randomised to receive estriol (1 mg daily), conjugated oestrogens (1.25 mg daily), oestrogen–progestagen sequential therapy (dose not specified), or black cohosh extract (Remifemin) 40 mg twice daily, for 24 weeks.\(^{(17)}\) At the end of the study, improvements in Kupperman Index scores were significantly lower, compared with baseline values, in all groups. There were no statistically significant differences between groups.

Another open, controlled study involving women with menopausal symptoms (\(n = 60\)) assessed the effects of black cohosh extract administered as a tincture (80 drops daily), compared with oestrogen (0.625 mg daily) or diazepam (2 mg daily), over a 12–week period.\(^{(31)}\) Cytological responses (proliferation and maturation of vaginal epithelial cells) were observed for participants in the black cohosh and oestrogen groups, but not in the diazepam group. For all three groups, improvements in neurovegetative and psychological symptoms (e.g. self–assessed depression) were reported.

**Other conditions**

Black cohosh has been reported to cause peripheral vasodilatation and an increase in peripheral blood flow, following the administration of a resinous constituent, acteina (500 \(\mu\)g/kg body weight), to patients suffering from peripheral arterial disease.\(^{(25)}\) The blood pressure of conscious individuals, both normotensive and hypertensive, was stated to be unaffected. The chemical composition of acteina is undefined.

In a randomised, double–blind trial, 82 patients with osteoarthritis or rheumatoid arthritis received a proprietary combination herbal preparation containing black cohosh 35 mg (other ingredients: white willow bark, guaiacum resin, sarsaparilla and poplar bark; Reumalex), or placebo, two tablets daily for two months.\(^{(32)}\) At the end of the study, there was a small, but statistically significant improvement in pain symptoms (as assessed by the Arthritis Pain Scale) in the treatment group, compared with the placebo group (\(p < 0.05\)).
Side-effects, Toxicity

A review of the literature on black cohosh\(^{18}\) describes a postmarketing surveillance study involving 629 women with menopausal symptoms who received standardised black cohosh extract as a tincture (80 drops daily) for 6–8 weeks.\(^{33}\) Tolerability was rated as ‘good’ in 93% of patients; mild, transient gastrointestinal symptoms were noted in 7% of patients.

A randomised, double-blind, placebo-controlled trial involving 80 women with menopausal symptoms who received standardised black cohosh extract (Remifemin) 40 mg twice daily, conjugated oestrogens, or placebo, for 12 weeks reported that non-specific adverse events, such as headaches, not considered treatment-related occurred in all three groups.\(^{15}\) Another controlled trial (\(n = 60\)) of the same extract reported that tolerability of black cohosh extract was ‘good’.\(^{17}\) In a randomised, double-blind trial involving 85 women with a history of breast cancer who received black cohosh (no details of extract provided) one tablet twice daily, or placebo, for 60 days, 10 adverse events occurred in the treatment group, compared with three in the placebo group.\(^{30}\) Three events were considered serious (treatment group: hysterectomy, recurrence of breast cancer; placebo group: appendectomy) all of which occurred in women who were also receiving tamoxifen. Minor adverse events (including constipation, weight gain, cramping, indigestion, vaginal bleeding) were not thought to be treatment related.

Older reference texts state that overdose may produce symptoms of nausea, vomiting, dizziness, visual and nervous disturbances, together with reduced pulse rate and increased perspiration.\(^{G22 \ G42 \ G49}\)
Contra-indications, Warnings

*In vitro* studies investigating the oestrogenic activity of extracts of black cohosh and their constituents report conflicting results (*see In vitro and animal studies, Hormonal activity*). Some studies have documented that certain constituents of black cohosh bind to oestrogen receptors,\(^{(2,16)}\) and others have reported oestrogenic activity *in vitro* for fukinolic acid, a hydroxycinnamic acid ester of fukiic acid, in an oestrogen receptor–positive breast cancer cell line (MCF-7 cells).\(^{(13)}\) These findings contrast with those of a previous *in vitro* study in oestrogen receptor–positive breast cancer cells, which reported that black cohosh extract did not stimulate cancer cell growth, i.e. it did not exhibit oestrogen–like effects, but at a concentration of 2.5 μg/mL led to a marked inhibition of breast cancer cell proliferation.\(^{(20)}\)

There is a view that the therapeutic effects of black cohosh extract are not attributable to oestrogenic effects, and that there is clinical evidence, such as lack of vaginal cell proliferation, as well as *in vitro* evidence to support this view.\(^{(18)}\) The German Commission E monograph states that there are no known contraindications to the use of black cohosh. Concern has been expressed that herbs with oestrogenic activity might stimulate breast cancer cell growth and oppose the effects of competitive oestrogen receptor antagonists, such as tamoxifen.\(^{(34)}\) A randomised, double–blind, placebo–controlled trial involving 85 women with a history of breast cancer assessed the effects of black cohosh (no details of extract provided) on hot flushes (*see Clinical studies, Menopausal symptoms*).\(^{(30)}\) Participants were stratified according to whether or not they were using tamoxifen. One woman receiving both black cohosh and tamoxifen experienced a recurrence of breast cancer, although it was reported that the woman had an increase in carcinoembryonic antigen when she entered the trial (this had not been reported to the referring physician).

Further study is required to establish whether black cohosh has oestrogenic activity. Herbal medicines with oestrogenic activity should be avoided in women with oestrogen–dependent tumours, such as breast cancer.\(^{(G50)}\)

**Pregnancy and lactation**

*In vitro* studies using rat uterus have indicated that black cohosh binds to uterine oestrogen receptors. Black cohosh has been used traditionally to assist labour. However, as there are insufficient data on the use of black cohosh during pregnancy and also during lactation, it is contra–indicated during these periods.\(^{(G49 \ G50 \ G56)}\)

There is an isolated report of a child born with no spontaneous breathing and
who subsequently experienced brain hypoxia and seizures following the oral administration of black cohosh and blue cohosh by a midwife in an attempt to induce labour in a woman who had had an uneventful pregnancy.\(^{35}\) The report has been criticised as it did not provide any further details of the dose or formulation of the herbs, and as the authors of the report make several assumptions about the clinical activity of the herbs on the basis of studies in animals.\(^{36}\)
Pharmaceutical Comment

The chemistry of black cohosh is well studied, although most of the documented information concerns the triterpene constituents. Most of the reputed traditional uses of black cohosh are not supported by data from experimental or clinical studies. One exception is the use of black cohosh in rheumatism and rheumatoid arthritis – there are data from in vitro and in vivo studies in rodents which indicate that black cohosh extracts have anti-inflammatory activity.\(^{(22,23)}\) Other pharmacological actions have been observed in both animals and humans which provide supporting data for the hormonal activity of the herb. However, there are conflicting reports on the oestrogenic activity of black cohosh. As it is known that there are at least two types of oestrogen receptor, there is a view that research relating to the oestrogenic effects of black cohosh needs to be considered against this background.\(^{(1)}\)

Little is known about the toxicity of black cohosh and excessive use should be avoided. It has been stated that the duration of use should not exceed three months.\(^{(G56)}\) Further study is required to determine whether black cohosh has oestrogenic activity before definitive statements about its use in women with oestrogen-dependent tumours can be made.

There is evidence from one published, randomised clinical trial to indicate that black cohosh extract is more effective than placebo in the treatment of menopausal symptoms. However, another randomised clinical trial involving women with a history of breast cancer who were experiencing menopausal symptoms found no effect for an unspecified black cohosh extract on the frequency and intensity of hot flushes. There are supporting data for the effects of black cohosh extracts on menopausal symptoms from open, uncontrolled studies and from postmarketing surveillance studies that assessed effectiveness as well as safety. However, further randomised clinical trials are required to establish the effects of black cohosh in women with menopausal symptoms. There is also a lack of information on the toxicity of black cohosh.
References

See also General References G3 G5 G6 G9 G22 G31 G32 G36 G41 G42 G43 G49 G50 G56 G64.

15. Stoll W. (Phytopharmacon influences atrophic vaginal epithelium: double–


20. Jacobson JS et al. Randomized trial of black cohosh for the treatment of...


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Species (Family)

*Caulophyllum thalictroides* (L.) Mich. (Berberidaceae)
Synonym(s)
Caulophyllum, Papoose Root, Squaw Root
Part(s) Used
Rhizome, root
Pharmacopoeial and Other Monographs

BHP 1983\(^{(G7)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents

See General References G22 G41 G48 G64.

**Alkaloids**
Quinolizidine and isoquinoline-types. Anagyrine, baptifoline, magnoflorine, methylcytisine (caulophyline). Other unidentified minor tertiary alkaloids.\(^1\)

**Saponins**
Caulosaponin and cauloside D yielding hederagenin on hydrolysis.\(^2\)

**Other constituents**
Citrullol, gum, resins, phosphoric acid, phytosterol and starch.

**Other Caulophyllum species**
A related species, *C. robustum* Maxim., is rich in triterpene glycosides (caulosides A–G), most of which possess hederagenin as their aglycone.
Food Use

Blue cohosh is not used in foods.
Blue cohosh is stated to possess antispasmodic, emmenagogue, uterine tonic and antirheumatic properties. Traditionally, it has been used for amenorrhoea, threatened miscarriage, false labour pains, dysmenorrhoea, rheumatic pains, and specifically for conditions associated with uterine atony.\textsuperscript{(G7 G64)}
Dosage

*Dried rhizome/root*
0.3–1.0 g or by decoction three times daily.\(^{(G7)}\)

*Liquid extract*
0.5–1.0 mL (1 : 1 in 70% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

In vitro and animal studies

A blue cohosh extract exhibited stimulant properties on the isolated guinea-pig uterus, although subsequent in vivo studies in cats, dogs and rabbits demonstrated no uterine activity.\(^3\) Antifertility actions documented in rats were reported to be caused by inhibition of ovulation\(^4\) and by interruption of implantation.\(^5\)

Smooth muscle stimulation has been documented for a crystalline glycoside constituent on the uterus (in vitro), the small intestine (in vitro), and the coronary blood vessels (in vivo) of various small mammals.\(^6\) The glycoside was also reported to cause erythrolysis and to be of an irritant nature. An earlier study that used a crystalline glycoside identified as caulosaponin, reported a variety of actions including an oxytocic effect on the isolated rat uterus, constriction of coronary and carotid blood vessels, a toxic action on cardiac muscle, and a spasmogenic action on the isolated intestine.\(^6\)

Methylcytisine is stated to have a nicotinic–like action, causing an elevation in blood pressure and stimulating both respiration and intestinal motility.\(^\text{G60}\)

An alcoholic extract of the aerial parts of blue cohosh produced up to 55% inhibition of inflammation in the carrageenan rat paw test\(^7\).
Side–effects, Toxicity

Powdered blue cohosh is stated to be irritant, especially to mucous membranes.\(^{(G51)}\) The leaves and seeds are reported to contain methylcytisine and some glycosides that can cause severe stomach pains. Children have been poisoned by eating the bright blue bitter–tasting seeds.\(^{(G22)}\) Caulosaponin is reported to be cardiototoxic, causing constriction of coronary blood vessels, to produce intestinal spasms, and to possess oxytocic properties.\(^{(G60)}\)
Contra-indications, Warnings

Blue cohosh may interfere with existing therapy for angina, and may irritate gastrointestinal conditions. Excessive doses may cause a rise in blood pressure, because of the methylcytisine constituent and give rise to other symptoms of nicotine poisoning.

Pregnancy and lactation

Blue cohosh should not be taken in pregnancy; it is reputed to be an abortifacient and to affect the menstrual cycle.\(^{(G30)}\) Some texts give conflicting advice. It has been documented that blue cohosh should be avoided by pregnant women,\(^{(G22)}\) only be taken once labour has commenced,\(^{(G49)}\) only taken in small doses during the first trimester of pregnancy,\(^{(G7)}\) or only be used under expert supervision.\(^{(G42)}\)
Pharmaceutical Comment

Limited data are available on the chemistry of blue cohosh. Documented pharmacological actions support some of the reputed traditional uses, although many of these are not suitable indications for self-medication. No evidence regarding antirheumatic properties was located, although anti-inflammatory action has been documented for the aerial plant parts. In view of the potential toxicity associated with blue cohosh, it should be used with caution.
References

See also General References G7 G10 G20 G22 G30 G31 G32 G36 G37 G41 G42 G48 G49 G51 G60 G64.


Species (Family)

i. *Cola nitida* A. Chev. (Sterculiaceae)

ii. *Cola acuminata* Schott & Endl. and related species
Synonym(s)
Cola Seed, Guru Nut, Kola Nut
  i. *Sterculia acuminata* Beauv.
Part(s) Used

Cotyledon
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

BP 2002\(^{(G71)}\)

Complete German Commission E\(^{(G3)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)

Ph Eur 2004\(^{(G72)}\)
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents


**Alkaloids**
Xanthine-types. Caffeine (0.6–3.0%), theobromine (up to 0.1%).

**Tannins**
Condensed type, catechins.

**Other constituents**
Betaine, cellulose, enzyme, fats, a glucoside, protein, red pigment and sugars.
Food Use

Cola is listed by the Council of Europe as a natural source of food flavouring (cola and cola nut extract: category 4, with limits on caffeine) (see Appendix 23). Cola is commonly used in foods. In the USA, it is listed as GRAS (Generally Recognised As Safe).
Herbal Use

Cola is stated to possess CNS stimulant, thymoleptic, antidepressant, diuretic, cardioactive and antidiarrhoeal properties. It has been used for depressive states, melancholy, atony, exhaustion, dysentery, atonic diarrhoea, anorexia, migraine and specifically for depressive states associated with general muscular weakness. (G6 G7 G8 G64)
Dosage

*Powdered cotyledons*
1–3 g or by decoction three times daily.\(^{(G6\ G7)}\)

*Liquid Extract of Kola*
(BPC 1949) 0.6–1.2 mL (1 : 1 in 60% alcohol).

*Tincture of Kola*
(BPC 1934) 1–4 mL (1 : 5 in 60% alcohol).
Pharmacological Actions

The xanthine constituents, caffeine and theobromine, are the active principles in cola. The pharmacological properties of caffeine are well documented and include stimulation of the CNS, respiratory system and skeletal muscle, cardiac stimulation, coronary dilatation, smooth muscle relaxation and diuresis. (G41) Cola-containing beverages are stated to provide active doses of caffeine. (G45)
Side–effects, Toxicity

Side–effects commonly associated with xanthine– containing beverages include sleeplessness, anxiety, tremor, palpitations and withdrawal headache.\(^{(G54)}\)
Contra–indications, Warnings
Consumption of cola should be restricted in individuals with hypertension or cardiac disorders, because of the caffeine content.

Pregnancy and lactation
It is generally recommended that caffeine consumption should be restricted during pregnancy, although conflicting reports have been documented regarding the association between birth defects and caffeine consumption. In view of this, excessive consumption of cola during pregnancy should be avoided. Caffeine is excreted in breast milk, but at concentrations too low to represent a hazard to breastfed infants.\(^{(G45)}\) As with all xanthine–containing beverages, excessive consumption of cola by lactating mothers should be avoided.
Pharmaceutical Comment

The principal active constituent in cola is caffeine. The reputed herbal uses of cola can be attributed to the actions of caffeine, and precautions associated with other xanthine-containing beverages are applicable to cola.
References


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Coltsfoot
Species (Family)

*Tussilago farfara* L. (*Asteraceae/Compositae*)
Synonym(s)

Farfara
Part(s) Used

Flower, leaf
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1983\(^{(G7)}\)

Complete German Commission E\(^{(G3)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL(G37)
Constituents
See General References G2 G22 G48 G64.

**Acids**
Caffeic acid, caffeoyltartaric acid, ferulic acid, gallic acid, \( p \)-hydroxybenzoic acid, and tannic acid (phenolic); malic acid and tartaric acid (aliphatic).\(^{(1)}\)

**Alkaloids**
Pyrrolizidine–type. Senkirkine 0.015% and senecionine (minor) (unsaturated)\(^{(2,3)}\) and tussilagine (saturated).\(^{(4)}\)

**Carbohydrates**
Mucilage (water–soluble polysaccharides) 7–8% yielding various sugars following hydrolysis (e.g. arabinose, fructose, galactose, glucose, uronic acid and xylose); inulin (polysaccharide).\(^{(5)}\)

**Flavonoids**
Flavonols (e.g. kaempferol, quercetin) and their glycosides.\(^{(1)}\)

**Tannins**
Up to 17% (type unspecified).

**Other constituents**
Bitter (glycoside), choline, paraffin (fatty acid), phytosterols (sitosterol, stigmasterol, taraxasterol), triterpene (amyrin), tussilagone (sesquiterpene)\(^{(6)}\) and volatile oil.
Food Use

Coltsfoot is not commonly used as a food but it is listed by the Council of Europe as a source of natural food flavouring (category N4). This category indicates that although coltsfoot is permitted for use as a food flavouring, there are insufficient data available for an assessment of toxicity to be made. (G16)
Herbal Use

Coltsfoot is stated to possess expectorant, antitussive, demulcent and antcatarrhal properties. It has been used for asthma, bronchitis, laryngitis and pertussis. (G2 G7 G49 G64)
Dosage

**Dried herb**
0.6–2.0 g by decoction three times daily.\(^{(G6)}\)

**Liquid extract**
0.6–2.0 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)

**Tincture**
2–8 mL (1 : 5 in 45% alcohol) three times daily.\(^{(G7)}\)

**Syrup**
2–8 mL (liquid extract 1 : 4 in syrup) three times daily.\(^{(G7)}\)
Pharmacological Actions

In vitro and animal studies

Antibacterial activity has been documented for coltsfoot against various Gram-negative bacteria including *Staphylococcus aureus*, *Proteus hauseri*, *Bordetella pertussis*, *Pseudomonas aeruginosa* and *Proteus vulgaris*.(7–9)

Anti-inflammatory activity comparable to that of indometacin, determined in Selye’s experimental chronic inflammation test, has been attributed to water-soluble polysaccharides in coltsfoot.(10) Weak acute anti-inflammatory activity has been reported for coltsfoot when tested against carrageenan-induced rat paw oedema.(11,12)

Platelet-activating factor (PAF) is known to be involved in various inflammatory, respiratory and cardiovascular disorders. The aggregating action of PAF is known to be weaker if intracellular concentrations of calcium are low. A sesquiterpene, L-652,469, isolated from coltsfoot buds has been reported to be a weak inhibitor of both PAF receptor binding and calcium channel blocker binding to membrane vesicles.(13) This combination of actions was found to effectively block PAF-induced platelet aggregation. L-652,469 was also found to be active orally, inhibiting PAF-induced rat paw oedema.(13) Interestingly, L-652,469 was reported to interact with the cardiac calcium channel blocker receptor complex (dihydropyridine receptor), but was also found to be a calcium channel blocker.(13)

Tussilagone has been reported to be a potent cardiovascular and respiratory stimulant.(6,14) Dose-dependent pressor activity following intravenous injection has been observed in the cat, rat and dog.(14) The pressor effect is stated to be similar to that of dopamine, but without tachyphylaxis. A significant stimulation of respiration was also observed.(6) Cardiovascular and respiratory effects are thought to be mediated by peripheral and central mechanisms, respectively.(6)
Side-effects, Toxicity

Coltsfoot has been reported to be phototoxic in guinea–pig skin.\(^{(15)}\)

Pyrrolizidine alkaloids with an unsaturated pyrrolizidine nucleus are known to be hepatotoxic in both animals and humans (see Comfrey). Of the pyrrolizidine alkaloids documented for coltsfoot, senecionine and senkirkine are unsaturated. Chronic hepatotoxicity has been described in rats following the incorporation of coltsfoot into their diet at concentrations ranging from 4 to 33\%.\(^{(16)}\) After 600 days, it was found that rats fed more than 4\% coltsfoot had developed hepatic tumours (haemangioendothelial sarcoma) while none were observed in the control group. Furthermore, histological changes associated with pyrrolizidine alkaloid toxicity, such as centrilobular necrosis of the liver and cirrhosis, were observed in many of the rats who had ingested coltsfoot but who had not developed tumours.\(^{(16)}\) The hepatotoxicity of coltsfoot was attributed to senkirkine, which is present at a concentration of only 0.015\%, thus highlighting the dangers associated with chronic exposure to low concentrations of pyrrolizidine alkaloids.

Newborn rats have been found to be more susceptible than weanlings to the hepatotoxic effects of senkirkine despite lacking the hepatic microsomal enzymes required for the formation of the toxic pyrrolic metabolites.\(^{(17)}\) Fatal hepatic veno–occlusive disease has been documented in a newborn infant whose mother had regularly consumed a herbal tea during pregnancy.\(^{(18)}\) Analysis of the herbal tea revealed the presence of 10 different plants including coltsfoot and a *Senecio* species (known source of pyrrolizidine alkaloids, see Liferoot). The mother exhibited no signs of hepatic damage, suggesting an increased sensitivity of the fetal liver to pyrrolizidine alkaloid toxicity.

Pre–blooming coltsfoot flowers are reported to contain the highest concentration of alkaloids.\(^{(3)}\) Considerable loss of both senkirkine and senecionine has been observed upon prolonged storage of the dried plant material.\(^{(3)}\) Senkirkine and senecionine are both easily extracted into hot water and, therefore, would presumably be ingested in a herbal tea prepared from the fresh plant.\(^{(3)}\) A cup of tea prepared from 10 g pre–blooming flowers has been estimated to contain a maximum of 70 μg senecionine and 1.4 mg senkirkine. Tea from the young leaves or mature plant would presumably contain considerably less alkaloids.\(^{(3)}\) These concentrations are not considered to represent a health hazard compared to the known hepatotoxicity of senecionine (intravenous LD\(_{50}\) 64 mg/kg body weight, mice).\(^{(3)}\) However, prolonged exposure to low concentrations of pyrrolizidine alkaloids has resulted in hepatotoxicity (see Comfrey).
Tussilagine LD$_{50}$ (mice, intravenous injection) has been determined as 28.9 mg/kg.$^{(14)}$
Contra-indications, Warnings

Excessive doses of coltsfoot may interfere with existing antihypertensive or cardiovascular therapy. In view of the known pyrrolizidine alkaloid content, excessive or prolonged ingestion should be avoided. In particular, herbal teas containing coltsfoot should be avoided.

Pregnancy and lactation
Coltsfoot should not be taken during pregnancy or lactation in view of the toxicity associated with the pyrrolizidine alkaloid constituents. Coltsfoot is reputed to be an abortifacient. (G30)
The majority of the traditional uses associated with coltsfoot can be attributed to the mucilage content. However, coltsfoot also contains toxic pyrrolizidine alkaloids albeit at a low concentration. The risk of exposure to low concentrations of pyrrolizidine alkaloids is unclear although hepatotoxicity following prolonged exposure has been documented (see Comfrey). The regular or excessive consumption of coltsfoot, especially in the form of herbal teas, should therefore be avoided.
References


Comfrey
Species (Family)

Symphytum officinale L. (Boraginaceae)
Synonym(s)

Consolidae Radix, Symphytum peregrinum Ledeb., Symphytum Radix

Related species include Prickly Comfrey (Symphytum asperum), Quaker and Russian Comfrey (Symphytum uplandicum, hybrid of S. officinale × S. asperum)
Part(s) Used

Leaf, rhizome, root
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

Complete German Commission E\(^{(G3)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR® for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL (external use only)\(^{(G37)}\)
Constituents
See General References G2 G6 G22 G41 G48 G64.

Alkaloids
Pyrrolizidine-type. 0.3%. Symphytine, symlandine, echimidine, intermidine, lycopsamine, myoscorpine, acetyllycopsamine, acetylintermidine, lasiocarpine, heliosupine, viridiflorine and echiumine.\(^{(1-5)}\)

Carbohydrates
Gum (arabinose, glucuronic acid, mannose, rhamnose, xylose); mucilage (glucose, fructose).

Tannins
Pyrocatechol-type. 2.4%.

Triterpenes
Sitosterol and stigmasterol (phytosterols), steroidal saponins and isobauerenol.

Other constituents
Allantoin 0.75–2.55%, caffeic acid, carotene 0.63%, chlorogenic acid, choline, lithospermic acid, rosmarinic acid and silicic acid.
Food Use

Comfrey is occasionally used as an ingredient of soups and salads. It is listed by the Council of Europe as natural source of food flavouring (category N4). This category indicates that although comfrey is permitted for use as a food flavouring, insufficient data are available to assess toxicity.\textsuperscript{G16}
Herbal Use

Comfrey is stated to possess vulnerary, cell-proliferant, astringent, antihaemorrhagic and demulcent properties. It has been used for colitis, gastric and duodenal ulcers, haematemesis, and has been applied topically for ulcers, wounds and fractures.\(^{(G2\ G6\ G7\ G8\ G49\ G64)}\)
Dosage

*Dried root/rhizome*
2–4 g in a decoction three times daily.\(^{(G7)}\)

*Root, liquid extract*
2–4 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)

*Ointment symphytum root*
10–15% root extractive in usual type ointment basis three times daily.\(^{(G7)}\)

*Dried leaf*
2–8 g or by infusion three times daily.\(^{(G7)}\)

*Leaf, liquid extract*
2–8 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

The classical pharmacology of pyrrolizidine alkaloids is overshadowed by the well-recognised toxicity of this class of compounds. Consequently, the majority of data documented for comfrey involve toxicity. Many useful reviews have been published on the toxicity of pyrrolizidine alkaloids in humans (see below).\(^{(5-11)}\)

**In vitro and animal studies**

Wound-healing and analgesic activities have been documented in rats administered comfrey extract orally.\(^{(12)}\) Percutaneous absorption of pyrrolizidine alkaloids obtained from comfrey is reported to be low in rats, with minimal conversion of the pyrrolizidine alkaloid N-oxides to the free pyrrolizidine alkaloids in the urine (reduction of the N-oxides is required before they can be metabolised into the reactive pyrrolic esters).\(^{(13,14)}\)

Rosmarinic acid has been isolated from comfrey (\textit{S. officinale}) as the main constituent with \textit{in vitro} anti-inflammatory activity.\(^{(15)}\) Biological activity was determined by inhibition of malonic dialdehyde formation in human platelets. Minor components, chlorogenic and caffeic acids, were not found to exhibit any significant activity. The pyrrolic esters have been reported to possess mild antimuscarinic activity, which is more pronounced in the non-hepatotoxic esters of saturated amino alcohols.\(^{(16)}\) Conversely, the free amino alcohols are reported to exert indirect cholinomimetic action involving the release of acetylcholine from postganglionic sites in the guinea-pig ileum.\(^{(16)}\)

Comfrey has been reported to stimulate the activity of the hepatic drug-metabolising enzyme aminopyrine N-demethylase in rats.\(^{(17)}\)

A comfrey extract has been reported to enhance uterine tone \textit{in vitro}.\(^{(18)}\) The action of comfrey was reported to be weaker than that exhibited by German chamomile, calendula and plantain, but stronger than that shown by shepherd’s purspe, St. John’s wort and uva-ursi.

**Clinical studies**

The antimuscarinic properties of certain pyrrolic esters have been utilised. Two non-hepatotoxic pyrrolizidine alkaloids, sarracine and platyphylline, have been used for the treatment of gastrointestinal hypermotility and peptic ulceration.\(^{(16)}\)
Side-effects, Toxicity

Two reports of human hepatotoxicity associated with the ingestion of comfrey have been documented.\(^{(19,20)}\) One case involved a 13-year-old boy who had been given a comfrey root preparation in conjunction with acupuncture to treat Crohn’s disease.\(^{(19)}\) The boy was diagnosed with veno-occlusive disease of the liver and the authors concluded comfrey to be the only possible causal factor of the liver disease. The second case involved a 49-year-old woman diagnosed with veno-occlusive disease.\(^{(20)}\) She had been taking various food supplements including a herbal tea and comfrey-pepsin pills. Pyrrolizidine alkaloids were identified in both the tea (stated to contain ginseng) and the comfrey-pepsin pills. The authors estimated that over a period of six months the woman had ingested 85 mg of pyrrolizidine alkaloids, equivalent to 15 μg/kg body weight per day. This report highlighted the potential toxicity associated with chronic ingestion of relatively small amounts of pyrrolizidine alkaloids.

The toxicity of pyrrolizidine alkaloids is well recognised. Pyrrolizidine alkaloids with an unsaturated pyrrolizidine nucleus are metabolised in the liver to toxic pyrrole metabolites.\(^{(8)}\) Acute toxicity results in hepatic necrosis, whereas chronic toxicity typically results in veno-occlusive disease characterised by the presence of greatly enlarged liver cells.\(^{(8,10)}\)

Reports of human hepatotoxicity associated with pyrrolizidine alkaloid ingestion have been documented.\(^{(5,8-10,21-30)}\) Many of these reports have resulted from crop (and subsequently flour and bread) contamination with Crotalaria, Heliotropium and Senecio species and from the use of pyrrolizidine-containing plants in medicinal ‘bush’ teas. In addition, pyrrolizidine alkaloid poisoning has been associated with the use of herbal teas in Europe and the United States.\(^{(20,25-27)}\) The diagnosis of veno-occlusive disease in a newborn infant who subsequently died highlights the susceptibility of the fetus to pyrrolizidine alkaloid toxicity.\(^{(30)}\) In this case, the mother had consumed a herbal tea as an expectorant during pregnancy. The tea, which was purchased from a pharmacy in Switzerland, was analysed and found to contain pyrrolizidine alkaloids. The mother did not exhibit any signs of hepatotoxicity.

Interestingly, liver function tests in 29 chronic comfrey users have been reported to show no abnormalities.\(^{(31)}\)

The hepatotoxicity of pyrrolizidine alkaloids is well documented in animals.\(^{(5)}\) In addition, carcinogenicity has been described in rats fed a diet supplemented with comfrey.\(^{(32)}\) The mutagenicity of comfrey has been
attributed to lasiocarpine,\textsuperscript{(23)} which is known to be mutagenic and carcinogenic. However, other workers have reported a lack of mutagenic activity for comfrey following assessment using direct bacterial test systems (Ames), host mediated assay (Legator), liver microsomal assay and the micronucleus technique.\textsuperscript{(33,34)}
Contra-indications, Warnings

In view of the hepatotoxic properties documented for the pyrrolizidine alkaloid constituents, comfrey should not be taken internally. The topical application of comfrey-containing preparations to broken skin should be avoided.

Pregnancy and lactation

The safety of comfrey has not been established. In view of the toxicity associated with the alkaloid constituents, comfrey should not be taken during pregnancy or lactation.
Comfrey is characterised by its pyrrolizidine alkaloid constituents. The hepatotoxicity of these compounds is well known, and cases of human poisoning involving comfrey have been documented. Human hepato toxicity with pyrrolizidine–containing plants is well documented, particularly following the ingestion of *Crotalaria*, *Heliotropium* and *Senecio* species. Comfrey has traditionally been used topically for treating wounds. Percutaneous absorption of pyrrolizidine alkaloids present in comfrey is reported to be low, although application of comfrey preparations to the broken skin should be avoided.

Licensed herbal products intended for internal use are not permitted to contain comfrey.

The inclusion of comfrey in products intended for topical application is permitted, provided the preparation is only applied to the unbroken skin and that its use is restricted to ten days or less at any one time.

As a result of a report by the Committee on Toxicity of Chemicals in Food to the Food Advisory Committee and the Ministry of Agriculture, Fisheries and Food, the health food trade voluntarily withdrew all products, such as tablets and capsules, and advice was issued that the root and leaves should be labelled with warnings against ingestion. It was considered that comfrey teas contained relatively low levels of pyrrolizidine alkaloids and did not need any warning labels.\(^{35}\)
References


17. Garrett BJ et al. Consumption of poisonous plants (Senecio jacobaea, Symphytum officinale, Pteridium aquilinum, Hypericum perforatum) by rats: Chronic toxicity, mineral metabolism, and hepatic drug–metabolizing


Corn Silk
Species (Family)

Zea mays L. (Gramineae)
Synonym(s)
Stigma Maydis, Zea
Part(s) Used

Stigma, style
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

Corn silk is not included in the GSL.(G37)
Constituents
See General References G2 G6 G40 G41 G44 G49 G64.

**Amines**
0.05%. Type not specified, although hordenine is listed for the genus *Zea*.

**Fixed oils**
1.85–2.25%. Contain glycerides of linoleic, oleic, palmitic and stearic acids.

**Saponins**
3% (unspecified).

**Tannins**
Up to 11.5–13% (unspecified).

**Other constituents**
Allantoin, bitter glycosides (1%), cryptoxanthin, cyanogenic compound (unidentified),\(^1\) flavone, gum, phytosterols (e.g. sitosterol, stigmasterol), pigments, resin, vitamins (C and K).
Corn silk is listed as a natural source of food flavouring (category N2). This category indicates that corn silk can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product. In the USA, corn silk is listed as GRAS (Generally Recognised As Safe).\(^{(G41)}\) The fruits are classified as category N1 with no restriction on their use.\(^{(G16)}\) Corn (maize) oil and flour are commonly used in cooking.
Herbal Use

Corn silk is stated to possess diuretic and stone-reducing properties. It has been used for cystitis, urethritis, nocturnal enuresis, prostatitis, and specifically for acute or chronic inflammation of the urinary system.\(^{G2 \ G6 \ G7 \ G8 \ G64}\)
Dosage

*Dried style/stigma*

4–8 g or by infusion three times daily.\(^{G6\ G7}\)

*Liquid Extract of Maize Stigmas*

(BPC 1923) 4–8 mL.

*Tincture*

5–15 mL (1 : 5 in 25% alcohol) three times daily.\(^{G6\ G7}\)

*Syrup of Maize Stigmas*

(BPC 1923) 8–15 mL.
Pharmacological Actions

In vitro and animal studies

Corn silk is stated to possess cholagogue, diuretic, hypoglycaemic, and hypotensive activities in laboratory animals.\(^2,\text{G}41\) Utilising aqueous extracts, a methanol-insoluble fraction has been reported to exhibit diuretic activity in rabbits,\(^\text{G}41\) and an isolated crystalline component has been documented to have a hypotensive action and to stimulate uterine contraction in rabbits.\(^3\) The latter two actions were thought to involve a cholinergic mechanism. The action of corn silk extract on experimental periodontolysis in hamsters has been documented.\(^4\)

Cryptoxanthin is stated to possess vitamin A activity,\(^\text{G}48\) and tannins are known to possess astringent properties.

Clinical studies

It has been stated that an aqueous extract is strongly diuretic in humans,\(^\text{G}41\) and that clinical studies have indicated corn silk to be effective in kidney and other diseases.\(^\text{G}41\) No further information on human studies was located to support these statements.
Side-effects, Toxicity

Allergic reactions including contact dermatitis and urticaria have been documented for corn silk, its pollen and for starch derived from corn silk.\(^{G51}\) Cornstarch is considered to be a known allergen.\(^{G51}\) The toxicity of a methanol-insoluble fraction of an aqueous corn silk extract has been reported to be low in rabbits. The effective intravenous dose for a diuretic action was documented as 1.5 mg/kg body weight compared to the lethal intravenous dose of 250 mg/kg.\(^{G41}\) Corn silk contains an unidentified toxic principle,\(^{1,2}\) and is listed as being capable of producing a cyanogenetic compound.\(^1\)
Contra-indications, Warnings

Corn silk may cause an allergic reaction in susceptible individuals. Excessive doses may interfere with hypoglycaemic drug therapy (*in vivo* hypoglycaemic activity has been documented) or with hypertensive or hypotensive therapy (*in vivo* hypotensive activity reported), and prolonged use may result in hypokalaemia because of the diuretic action.

**Pregnancy and lactation**
Corn silk has been documented to stimulate uterine contractions in rabbits. In view of this, doses of corn silk greatly exceeding amounts used in foods should not be taken during pregnancy or lactation.
Pharmaceutical Comment

Limited information is available on the constituents of corn silk. Extracts have been reported to exhibit diuretic actions in both humans and animals, thus justifying the reputed herbal uses. However, no additional data were located to support these reported actions. In view of the lack of toxicity data, excessive use of corn silk should be avoided.
References

See also General References G2 G6 G9 G10 G16 G36 G37 G40 G41 G48 G49 G51 G64.


Species (Family)

Agropyron repens (L.) Beauv. (Gramineae)
Synonym(s)
Agropyron, Dogs Grass, Quackgrass, Triticum, *Triticum repens* L., Twitchgrass
Part(s) Used

Rhizome
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)

BP 2002\(^{(G71)}\)

Complete German Commission E\(^{(G3)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)

Ph Eur 2004\(^{(G72)}\)

WHO volume 1 1999\(^{(G63)}\)
Legal Category (Licensed Products)

GSL (Agropyron)\(^{G37}\)
Constituents
See General References G2 G7 G40 G41 G53 G64.

Carbohydrates
Fructose, glucose, inositol, mannitol, mucilaginous substances (10%), pectin, triticin.

Cyanogenic glycosides
Unspecified.

Flavonoids
Tricin and other unidentified flavonoids.

Saponins
No details documented.

Volatile oils
0.05%. Agropyrene (95%). Presence of agropyrene has been disputed,\(^{(1)}\) with the oil reported to consist mainly of the monoterpenes carvacrol, trans-anethole, carvone, thymol, menthol, menthone and \(p\)-cymene and three sesquiterpenes.

Other constituents
Fixed oil, vanillin glucoside.
Food Use

Couchgrass is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that couchgrass can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\(^{(G16)}\) In the USA, couchgrass is listed as GRAS (Generally Recognised As Safe).\(^{(G41)}\)
Herbal Use

Couchgrass is stated to possess diuretic properties. It has been used for cystitis, urethritis, prostatitis, benign prostatic hypertrophy, renal calculus, lithuria, and specifically for cystitis with irritation or inflammation of the urinary tract. (G2 G7 G64)
Dosage

*Dried rhizome*
4–8 g or in decoction three times daily.\(^{(G7)}\)

*Liquid extract*
4–8 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)

*Tincture*
5–15 mL (1 : 5 in 40% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

In vitro and animal studies

Couchgrass is stated to exhibit diuretic and sedative activities in rats and mice, respectively.\(^{(G41)}\) Broad antibiotic activity has been documented for agropyrene and its oxidation product.\(^{(G41)}\) An ethanolic extract was found to exhibit only weak inhibition (14%) of carrageenan–induced inflammation in the rat paw.\(^{(2)}\)

Couchgrass has been reported to be phytotoxic with flavonoid components implicated as the active constituents.\(^{(3)}\)
Side-effects, Toxicity

None documented for couchgrass. An unspecified cyanogenetic glycoside has been reported as a constituent of couchgrass, although no further details were located.\textsuperscript{G7}
Contra-indications, Warnings

In view of its reputed diuretic action, excessive or prolonged use of couchgrass should be avoided since this may result in hypokalaemia.

Pregnancy and lactation

In view of the limited pharmacological and toxicological data, the use of couchgrass during pregnancy and lactation should be avoided.
Pharmaceutical Comment

Limited chemical data are available for couchgrass and little scientific evidence was located to justify the traditional herbal uses. Agropyrene is regarded as the main active principle in couchgrass on account of its antibiotic effect, although the presence of agropyrene in the volatile oil has been disputed.\(^{(1)}\) In view of the lack of toxicity data, excessive ingestion should be avoided.
Species (Family)

Primula veris L. (Primulaceae)
Synonym(s)
Paigle, Peagle, Primula, *Primula officinalis* (L.) Hill.
Part(s) Used

Flower
Pharmacopoeial and Other Monographs

BHP 1983\(^{(G7)}\)

Complete German Commission E (Primrose flower)\(^{(G3)}\)

ESCOP 1997\(^{(G52)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL(G37)
Constituents

See General References G2 G40 G49 G59 G62 G64.

Carbohydrates
Arabinose, galactose, galacturonic acid, glucose, rhamnose, xylose and water-soluble polysaccharide (6.2–6.6%).

Flavonoids
Apigenin, isorhamnetin, kaempferol, luteolin and quercetin.\(^1\)

Phenols
Glycosides primulaveroside (primulaverin) and primveroside.

Quinones
Primin and other quinone compounds.

Saponins
Primula acid in sepals but saponins absent from other parts of the flower.

Tannins
Condensed (e.g. proanthocyanidin B2), pseudotannins (e.g. epicatechin, epigallocatechin).\(^1\)

Other constituents
Silicic acid and volatile oil (0.1–0.25%).

Other plant parts
Saponins have been documented for the underground parts.\(^1\) ‘Primulic acid’ is a collective term for the saponin mixture.\(^2\) Primulic acid A glycoside (5–10%) yields primulagenin A as aglycone together with arabinose, galactose, glucose, glucuronic acid, rhamnose and xylose.\(^3,4\) The saponin content of the roots is stated to peak at two years.\(^5\) After five years of storage the saponin content was reported to have decreased by 45%.
Food Use

Cowslip is not commonly used in foods. A related species, *Primula eliator*, is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that *Primula eliator* can be added to foodstuffs, provided that the concentration of coumarin does not exceed 2 mg/kg.\(^{(16)}\) Coumarins, however, are not documented as constituents of *Primula veris*, the subject of this monograph.
Herbal Use

Cowslip is stated to possess sedative, antispasmodic, hypnotic, mild diuretic, expectorant and mild aperient properties. It has been used for insomnia, nervous excitability, hysteria and specifically for anxiety states associated with restlessness and irritability. (G2 G7 G64)
Dosage

**Dried flowers**
1–2 g as an infusion three times daily.$^G7$

**Liquid extract**
1–2 mL (1 : 1 in 25% alcohol) three times daily.$^G7$
Pharmacological Actions

In vitro and animal studies

The saponin fraction has been reported to cause an initial hypotension followed by a long-lasting hypertension in anaesthetised animals.\(^6\)

*In vitro*, the saponins have been documented to inhibit prostaglandin (PG) synthetase, but to a lesser extent than aspirin because of insignificant protein binding; to exhibit a slight anti-inflammatory effect against carrageenan rat paw oedema; to contract isolated rabbit ileum; and to possess analgesic and antigranulation activity.\(^6\)

Flavonoid and tannin constituents have been documented for cowslip. A variety of activities has been reported for flavonoids including anti-inflammatory and antispasmodic effects. The tannins are known to be astringent.
Side-effects, Toxicity

Allergic contact reactions to related *Primula* species have been documented; quinone compounds are stated to be the allergenic principles with primin described as a strong contact allergen.\(^7\) Two positive patch test reactions to cowslip have been recorded, although allergenicity was not proven.\(^{G51}\) An \(\text{LD}_{50}\) value (mice, intraperitoneal injection) for the saponin fraction is documented as 24.5 mg/kg body weight compared to a value of 9.5 mg/kg for reparil (aescin). Haemolytic activity has been reported for the saponins, and an aqueous extract of cowslip is stated to contain saponins that are toxic to fish. Saponins are stated to be irritant to the gastrointestinal tract.

The toxicity of cowslip seems to be associated with the saponin constituents. However, these compounds have only been documented for the underground plant parts, and not for the flowers which are the main plant parts used in the UK.
Contra-indications, Warnings

Cowslip may cause an allergic reaction in sensitive individuals. Excessive doses may interfere with hypo- or hypertensive therapy or cause gastrointestinal irritation.

Pregnancy and lactation
The safety of cowslip has not been established. In view of the lack of toxicity data, use of cowslip during pregnancy and lactation should be avoided.
Pharmaceutical Comment

The chemistry of cowslip is not well documented and it is unclear whether saponins reported as constituents of the underground plant parts are also present in the flowers. Little pharmacological information has been documented to justify the herbal uses of cowslip. In view of the lack of toxicity data, excessive use of cowslip should be avoided.
References

See also General References G2 G3 G7 G15 G16 G36 G37 G40 G44 G49 G51 G52 G59 G62 G64.


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Species (Family)

*Vaccinium macrocarpon* Ait, *Vaccinium oxycoccus* (Ericaceae)


Synonym(s)

Large Cranberry (*V. macrocarpon*) is the species grown for commercial purposes.\(^1\) *V. oxycccus* is European Cranberry, Mossberry and Small Cranberry.
Part(s) Used

Fruit (whole berries)
Pharmacopoeial and Other Monographs

Martindale 33rd edition\(^{(G67)}\)
Legal Category (Licensed Products)

Cranberry is not included in the GSL.$^{(G37)}$
Constituents

**Acids**
Citric, malic, quinic and benzoic acids are present.\(^{(2)}\)

**Carbohydrates**
Fructose and oligosaccharides.

**Phenolics**
Anthocyanins and proanthocyanidins.

**Other constituents**
Trace glycoside has been isolated from *V. oxyccoccus*.\(^{(3)}\) Cranberries are also a good source of fibre. Cranberry juice cocktail contains more carbohydrate than do products (i.e. soft or hard gelatin capsules) based on cranberry powder (prepared from rapidly dried fruits), whereas the latter contain more fibre.\(^{(2)}\) Alkaloids (*N*-methylazatricyclo type) have been isolated from the leaves.\(^{(4)}\)
Cranberries are commonly used in foods;\(^{(5)}\) cranberry juice cocktail (containing approximately 25% cranberry juice) is widely available.\(^{(2,5)}\) Cranberry is listed by the Council of Europe as a natural source of food flavouring (fruit: category 1) (see Appendix 23).\(^{(G17)}\)
Cranberry juice and crushed cranberries have a long history of use in the treatment and prevention of urinary tract infections.\(^{(1)}\) Traditionally, cranberries have also been used for blood disorders, stomach ailments, liver problems, vomiting, loss of appetite, scurvy and in the preparation of wound dressings.\(^{(5)}\)
Dosage

The doses used in clinical trials of cranberry for prevention of urinary tract infections have been variable. One study used 300 mL cranberry juice cocktail (containing 30% cranberry concentrate) daily for six months.\(^{(6)}\)
Pharmacological Actions

Documented activity for cranberry is mainly of its use in the prevention and treatment of urinary tract infections; its role in urinary tract infection has been reviewed.\(^1\)

Initially it was thought that the antibacterial effect of cranberry juice was due to its ability to acidify urine and, therefore, to inhibit bacterial growth. However, recent work has focused on the effects of cranberry in inhibiting bacterial adherence and on determining anti-adhesion agents in cranberry juice. Bacterial adherence to mucosal surfaces is considered to be an important step in the development of urinary tract infections;\(^7\) it is facilitated by fimbriae (protein aceous fibres on the bacterial cell wall) which produce adhesins that attach to specific receptors on uroepithelial cells.\(^8\)

\textbf{In vitro and animal studies}

In \textit{in vitro} studies using human urinary tract isolates of \textit{Escherichia coli}, cranberry cocktail (which contains fructose and vitamin C in addition to cranberry juice) inhibited bacterial adherence to uroepithelial cells by 75% or more in over 60% of the clinical isolates.\(^9\) In addition, urine from mice fed cranberry juice significantly inhibited \textit{E. coli} adherence to uroepithelial cells when compared with urine from control mice.\(^9\) However, these studies did not define the bacteria tested in terms of the type of fimbriae they might have expressed (specific fimbriae mediate bacterial adherence to cells).

Irreversible inhibition of adherence of urinary isolates of \textit{E. coli} expressing type 1 and type P fimbriae has been demonstrated with cranberry juice cocktail.\(^10\) It was thought that fructose might be responsible for the inhibition of type 1 fimbriae\(^10\) and an unidentified high molecular weight substance responsible for type P fimbriae inhibition.\(^11\) Further \textit{in vitro} studies in which cranberry juice was added to the growth medium of P-fimbriated \textit{E. coli} duplicated immediate inhibition of adherence, but also showed the loss of fimbriae with cellular elongation after long-term exposure; such changed bacteria are unable to adhere to urothelium.\(^12\)

Proanthocyanidins extracted from cranberries have been shown to inhibit the adherence of P-fimbriated \textit{E. coli} to uroepithelial cell surfaces at concentrations of 10–50 μg/mL, suggesting that proanthocyanidins may be important for the stated effects of cranberry in urinary tract infections.\(^13\)

The effects of a high molecular weight constituent of cranberry juice on adhesion of bacterial strains found in the human gingival crevice have also
been investigated.\(^{14}\) A non-dialysable material derived from cranberry juice concentrate used at concentrations of 0.6–2.5 mg/mL reversed the interspecies adhesion of 58% of 84 bacterial pairs. Gram-negative dental plaque bacteria appeared to be more sensitive to the inhibitory effects of the cranberry constituent on adhesion.\(^{14}\)

Crude extracts of cranberry have been reported to exhibit potential anticarcinogenic activity in vitro as demonstrated by inhibition of the induction of ornithine decarboxylase (ODC) by the tumour promoter phorbol 12–myristate 13–acetate (TPA).\(^{15}\) The greatest activity appeared to be in the polymeric proanthocyanidin fraction which had an IC\(_{50}\) for ODC activity of 6.0 μg. The anthocyanidin fraction and the ethyl acetate extract were either inactive or relatively weak inhibitors of ODC activity.

A cranberry extract with a polyphenolic content of 1548 mg gallic acid equivalents per litre inhibited low-density lipoprotein (LDL) oxidation in vitro.\(^{16}\)

Cranberry juice has demonstrated marked in vitro antifungal activity against Epidermophyton floccosum and against several Microsporum and Tricho phyton species, but had no effect against Candida albicans.\(^{17}\) Benzoic acid and/or other low molecular weight constituents of cranberry juice were reported to be responsible for the fungistatic action.

Clinical studies

Clinical trials investigating the use of cranberries for the treatment and prevention of urinary tract infections have been subject to Cochrane systematic reviews; both of these systematic reviews sought to include all randomised or quasi-randomised controlled trials.\(^{18,19}\)

Prevention of urinary tract infections

Four trials\(^{6,20–22}\) were included in a systematic review of cranberries for prevention of urinary tract infections; three trials compared the effectiveness of cranberry juice versus placebo or water and one trial compared cranberry capsules with placebo.\(^{19}\) Three of the four trials reported beneficial effects for cranberry compared with placebo on at least one of the outcomes (number of symptomatic or asymptomatic urinary tract infections, side-effects, adherence to therapy). However, the methodological quality of the trials was found to be poor and the reliability of the results questionable. It was stated that 'on the basis of the available evidence, cranberry juice cannot be recommended for the prevention of urinary tract infections in susceptible populations'.\(^{19}\)
The largest study of cranberry juice for the prevention of urinary tract infections was a double-blind, placebo-controlled trial involving 153 women (mean age 78.5 years) randomised to receive 300 mL cranberry juice cocktail \((n = 72)\) or an indistinguishable placebo \((n = 81)\) daily for six months.\(^6\) The odds of experiencing bacteriuria with pyuria were significantly lower in cranberry-treated subjects than in those who received a placebo beverage \((p = 0.004)\). A randomised, controlled, crossover study was conducted involving 38 persons (mean age 81 years) who had had hospital treatment and were waiting to be transferred to a nursing home.\(^{21}\) Subjects received cranberry juice (15 mL) mixed with water or water alone twice daily for four weeks before crossing over to the alternative regimen. Seventeen participants completed the study and, of the seven from whom data were suitable for comparison, there were fewer occurrences of bacteriuria during the period of treatment with cranberry juice.\(^{21}\)

The role of cranberry in the prevention of urinary tract infections in younger women has been explored in a randomised, double-blind, placebo-controlled, crossover trial involving 19 non-pregnant, sexually active women aged 18–45 years.\(^{22}\) Participants received capsules containing 400 mg cranberry solids daily (exact dose not stated) or placebo for three months before crossing over to the alternative regimen. Ten subjects completed the six-month study period. Of the 21 incidents of urinary tract infection recorded among these participants, significantly fewer occurred during periods of treatment with cranberry than with placebo \((p < 0.005)\).\(^{22}\)

A randomised, physician-blind, crossover study investigated the efficacy of cranberry cocktail (30% cranberry concentrate) (15 mL/kg/day) for six months in 40 children (age range 1.4–18 years, mean age 9.35 years) with neuropathic bladder and managed by clean intermittent catheterisation; water was used as a control.\(^{20}\) No benefit was reported for cranberry compared with control.

A randomised, double-blind, placebo-controlled, crossover trial of the effects of consumption of cranberry concentrate on the prevention of bacteriuria and symptomatic urinary tract infection has been carried out in children \((n = 15)\) with neurogenic bladder receiving clean intermittent catheterisation.\(^{23}\) Children drank 2 oz of cranberry concentrate or placebo daily for three months before changing to the alternative regimen. At the end of the study, the number of urinary tract infections occurring under each regimen was identical \((n = 3)\). There was no significant difference between cranberry treatment and placebo with regard to the number of collected urine samples testing positively for a pathogen (75% of samples for both cranberry and placebo) \((p = 0.97)\). It was concluded that cranberry concentrate had no
Treatment of urinary tract infections

Although several trials investigating the effectiveness of cranberry juice and cranberry products for treating urinary tract infections were found, none of these trials met all the inclusion criteria for systematic review. Two of the studies found did report a beneficial effect with cranberry products, although both contained methodological flaws and no firm conclusions can be drawn from these studies. Thus, it was stated that ‘at the present time, there is no evidence to suggest that cranberry juice or other cranberry products are effective in treating urinary tract infections’ (see Pharmaceutical Comment).

Other studies

Early studies involving the administration of large amounts of cranberry juice to human subjects reported reductions in mean urinary pH values. A crossover study involving eight subjects with multiple sclerosis reported that administration of cranberry juice and ascorbic acid was more effective than orange juice and ascorbic acid in acidifying the urine. However, neither treatment consistently maintained a urinary pH lower than 5.5, the pH previously determined as necessary for maintaining bacteriostatic urine. Inhibition of bacterial adherence (see In vitro and animal studies) has been observed with urine from 22 human subjects who had ingested cranberry cocktail 1–3 hours previously. Protection against bacterial adhesion has also been reported in a study involving urine collected from ten healthy male volunteers who had ingested water, ascorbic acid (500 mg twice daily for 2.5 days) or cranberry (400 mg three times daily for 2.5 days) supplements. Urine samples were used to determine uropathogen adherence to silicone rubber in a parallel plate flow chamber; urine obtained after ascorbic acid or cranberry supplementation reduced the initial deposition rates and numbers of adherent E. coli and Enterococcus faecalis, but not Pseudomonas aeruginosa, Staphylococcus epidermidis or C. albicans.

Other preliminary studies have explored the use of cranberry juice in reducing urine odours, in improving peristomal skin conditions in urostomy patients and in reducing mucus production in patients who have undergone entero–uroplasty.

The ingestion of cranberry juice by subjects with hypochlorhydria due to omeprazole treatment or atrophic gastritis has been shown to result in increased protein–bound vitamin B₁₂ absorption, although the clinical benefit of ingesting cranberry juice along with a meal (i.e. with the buffering action
(33) Possible mechanisms by which the ingestion of an acidic drink such as cranberry juice could result in improved protein–bound vitamin B\textsubscript{12} absorption include increased release of vitamin B\textsubscript{12} from protein by direct action of acid on the vitamin B\textsubscript{12}–protein bond and a pH-sensitive bacterial binding activity of vitamin B\textsubscript{12} that is altered in an acidic environment.\textsuperscript{(33)}
Side–effects, Toxicity

None documented for cranberry, although diarrhoea is possible if large quantities are consumed.\(^{(G31)}\) One study has reported that no subjects withdrew because of undesirable side–effects,\(^{(22)}\) although this study involved only a small number \((n = 19)\) of patients. A systematic review of cranberry products for the prevention of urinary tract infections reported that the drop–out rates in the four studies included \((6,20–22)\) were high \((20–55\%)\).\(^{(19)}\) In one of these studies, of 17 withdrawals during cranberry treatment (a further two occurred during the control period), nine participants gave the taste of cranberry as the reason for withdrawal.\(^{(20)}\)

It has been claimed that ingesting large amounts of cranberry juice may result in the formation of uric acid or oxalate stones secondary to a constantly acidic urine and because of the high oxalate content of cranberry juice.\(^{(1)}\) However, it has also been stated that the role of cranberry juice as a urinary acidifier has not been well established.\(^{(34)}\) The use of cranberry juice in preventing the formation of stones which develop in alkaline urine, such as those comprising magnesium ammonium phosphate and calcium carbonate, has been described.\(^{(26)}\)
Contra–indications, Warnings

The calorific content of cranberry juice should be borne in mind. Patients with diabetes who wish to use cranberry juice should be advised to use sugar–free preparations. Patients using cranberry juice should be advised to drink sufficient fluids in order to ensure adequate urine flow.\(^{(G31)}\) Although a constituent of cranberry juice has been reported to have potential for altering the subgingival microbiota, some commercially available cranberry juice cocktails may not be suitable for oral hygiene purposes because of their high dextrose and fructose content.\(^{(14)}\)

It has been stated that cranberry should be used with caution in patients with benign prostatic hypertrophy or urinary obstruction, because there is the theoretical possibility that cranberry may enhance the elimination of drugs excreted in urine.\(^{(G31)}\) Interference with dipstick tests for glucose and haemoglobin in urine has been reported in a study involving 28 patients who had drunk 100 or 150 mL of low–sugar or regular cranberry juice daily for seven weeks;\(^{(35)}\) ascorbic acid in cranberry juice was reported to be the component responsible for interference resulting in negative test results.

**Pregnancy and lactation**

There are no known problems with the use of cranberry during pregnancy. Doses of cranberry greatly exceeding the amounts used in foods should not be taken during pregnancy and lactation.
Pharmaceutical Comment

Limited chemical information is available for cranberry. Documented *in vitro* and animal studies provide supporting evidence for a mechanism of action for cranberry in preventing urinary tract infections. However, little is known about the specific active constituent(s); proanthocyanidins have been reported to be important.\(^{13}\)

Preliminary clinical trials of cranberry for the prevention of urinary tract infections have generally been uncontrolled and/or involved only small numbers of patients. The validity of the results of a controlled trial involving relatively large numbers of (female only) patients\(^6\) was questioned because of methodological shortcomings in the study design, particularly the method of randomisation.\(^{36,37}\) Other controlled studies claiming to involve random assignment to treatment\(^{20–22}\) either did not employ true randomisation\(^{21}\) or the method of randomisation was not stated.\(^{20,22}\) In addition, these four controlled studies\(^{6,20–22}\) differed in the formulations of cranberry, doses and treatment periods used. Therefore, clinical studies do not provide compelling evidence for the efficacy of cranberry in the prevention of urinary tract infections, nor do they provide evidence that it is not efficacious. However, the findings do indicate that the area warrants further investigation. Cochrane systematic reviews of cranberry for the treatment and prevention of urinary tract infections have stated that, at present, there is no reliable evidence to suggest that cranberry juice or other cranberry products are effective.\(^{18,19}\) It has also been stated that properly randomised, double-blind, placebo-controlled, parallel group trials using appropriate outcome measures are needed in order to determine the efficacy of cranberry products in the prevention and treatment of urinary tract infections.\(^{18,19}\) Prevention trials should be of at least 6–months’ duration in order to take into account the natural course of the illness.\(^{19}\)

Patients wishing to use cranberry for urinary tract infections should be advised to consult a pharmacist, doctor or other suitably trained health care professional for advice.
See also General References G31 G43.


18. Kilbourn JP. Interference with dipstick tests for glucose and hemoglobin in
urine by ascorbic acid in cranberry juice. *Clin Chem* 1987; **33**: 1297. (PubMed)


Damiana
Species (Family)

*Turnera diffusa* Willd. var. *aphrodisiaca* Urb. (Bignoniaceae/Turneraceae) and related species indigenous to Texas and Mexico.
Synonym(s)

Part(s) Used

Leaf, stem
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents
See General References G6 G22 G40 G41 G64.

Carbohydrates
Gum 13.5%, starch 6%, sugars.

Cyanogenic glycosides
Tetraphyllin B. \(^{(1)}\)

Phenolic glycoside
Arbutin (up to 0.7%). \(^{(2)}\)

Tannins
3.5%. Type unspecified.

Volatile oils
0.5–1.0%. At least 20 components including 1,8-cineole (11%), \(p\)-cymene (2%), \(\alpha\)- and \(\beta\)-pinene (2%), thymol, \(\alpha\)-copaene, \(\delta\)-cadinene and calamene. The presence of 1,8-cineole and \(p\)-cymene has been disputed. \(^{(2)}\)

Other constituents
Acids (fatty, plant), alkanes (e.g. hexacosanol–1 and triacontane), damianin (7%) (a bitter principle), flavone, \(\beta\)-sitosterol, resin (6.5%). \(^{(3)}\)
Food Use

Damiana is used in foods and is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that damiana can be added to foodstuffs in small quantities with a possible limitation of an active principle (as yet unspecified) in the final product.\(^{(G16)}\) In the USA, damiana is approved for food use.\(^{(G41)}\)
Herbal Use

Damiana is stated to possess antidepressant, thymoleptic, mild purgative, stomachic and reputedly aphrodisiac properties.\(^{(4)}\) It has been used for depression, nervous dyspepsia, atonic constipation, coital inadequacy, and specifically for anxiety neurosis with a predominant sexual factor.\(^{(G6 \ G7 \ G8 \ G64)}\)
Dosage

*Dried leaf*
2–4 g or by infusion three times daily.\(^{(G6 \ G7)}\)

*Liquid Extract of Damiana*
(BPC 1934) 2–4 mL.
Pharmacological Actions

In vitro and animal studies

Hypoglycaemic activity has been reported in mice following both oral and intraperitoneal administration of damiana.\(^5\) An ethanolic extract was stated to exhibit CNS-depressant activity although no other experimental details were available.\(^6\)

Antibacterial activity against *Escherichia coli, Proteus mirabilis, Pseudomonas aeruginosa* and *Staphylococcus aureus* has been documented for a mixed herbal preparation, with some of the activity attributed to damiana.\(^7\) The same herbal preparation was also reported to inhibit acetylcholine-induced spasm of the isolated guinea–pig ileum, although none of the antispasmodic activity was attributed to damiana.\(^7\)

Arbutin is stated to be responsible for the urinary antiseptic properties (see Uva-Ursi). However, the arbutin content of damiana is much less than that quoted for uva–ursi (0.7% and 5 to 18%, respectively).

The roots of various *Turnera* species have exhibited utero–activity.\(^G30\)

Clinical studies

A herbal preparation containing damiana as one of the ingredients was reported to have a favourable effect on the symptoms of irritable bladder associated with functional and neurohormonal disorders, and on bacterial bladder infections.\(^7\)
Side–effects, Toxicity

Tetanus–like convulsions and paroxysms resulting in symptoms similar to those of rabies or strychnine poisoning have been described in one individual following the ingestion of approximately 200 g damiana extract; cyanide poisoning was considered to be a possible cause. No other reported side–effects for damiana were located.

High doses of arbutin (e.g. 1 g) are considered to be toxic, although the concentration of arbutin documented for damiana (1 g arbutin is equivalent to more than 100 g plant material) is probably too low to warrant concerns over safety.
Contra–indications, Warnings

Excessive use should be avoided because of the presence of cyanogenetic glycosides and arbutin; damiana may interfere with existing hypoglycaemic therapy.

Pregnancy and lactation
The safety of damiana has not been established. In view of the lack of toxicity data and possible cyanogenetic constituents, doses greatly exceeding amounts used in foods should not be taken during pregnancy or lactation.
Pharmaceutical Comment

There is limited chemical information available on damiana. There has been little documented evidence to justify the herbal uses, and the reputation of damiana as an aphrodisiac is unproven.\(^{(7,8)}\) In view of the lack of toxicity data and reported cyanogenetic and arbutin constituents, excessive use of damiana should be avoided.
See also General References G6 G9 G10 G16 G22 G30 G31 G32 G36 G37 G40 G41 G43 G64.

Dandelion
Species (Family)

*Taraxacum officinale* Weber (Asteraceae/Compositae)
Synonym(s)

Part(s) Used

Leaf, root
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

Complete German Commission E\(^{(G3)}\)

ESCOP 1996\(^{(G52)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL(G37)
Constituents

See General References G2 G6 G8 G22 G41 G48 G52 G57 G64.

**Acids**
Caffeic acid, \( p \)-hydroxyphenylacetic acid, chlorogenic acid,\(^{(1)}\) cichoric acid, monocaffeyl tartaric acids\(^{(2)}\) linoleic acid, linolenic acid, oleic acid and palmitic acid.

**Coumarins**
Cichoriin and aesculin.\(^{(2)}\)

**Flavonoids**
Luteolin–7–glucoside and luteolin–7–diglucosides.\(^{(2)}\)

**Minerals**
Potassium 4.5% in leaf, 2.45% in root.\(^{(3)}\)

**Resin**
Undefined bitter complex (taraxacin).

**Terpenoids**
Sesquiterpene lactones taraxinic acid (germacranolide) esterified with glucose,\(^{(4)}\) and eudesmanolides.\(^{(5)}\)

**Vitamins**
Vitamin A 14 000 iu/100 g leaf (compared with 11 000 iu/100 g carrots).

**Other constituents**
Carotenoids, choline, inulin, pectin, phytosterols (e.g. sitosterol, stigmasterol, taraxasterol, homotaraxasterol), sugars (e.g. fructose, glucose, sucrose), triterpenes (e.g. \( \beta \)-amyrin, taraxol, taraxerol).
Food Use

Dandelion is used as a food, mainly in salads and soups. The roasted root and its extract have been used as a coffee substitute.\(^{(G41)}\) Dandelion is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that dandelion can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\(^{(G16)}\) In the USA, dandelion is listed as GRAS (Generally Recognised As Safe).\(^{(G41)}\)
Dandelion is stated to possess diuretic, laxative, cholagogue and antirheumatic properties. It has been used for cholecystitis, gallstones, jaundice, atonic dyspepsia with constipation, muscular rheumatism, oliguria, and specifically for cholecystitis and dyspepsia. The German Commission E approved use of root and herb for disturbance of bile flow, stimulation of diuresis, loss of appetite and dyspepsia. Root is used in combination with celandine herb and artichoke for epigastric discomfort due to functional disorders of the biliary system.
Dosage

Dried leaf
4–10 g or by infusion three times daily.\(^{(G6 \ G7)}\)

Leaf, liquid extract
4–10 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G6 \ G7)}\)

Leaf tincture
2–5 mL.\(^{(G3)}\)

Leaf, fresh juice
5–10 mL.\(^{(G52)}\)

Dried root
2–8 g or by infusion or decoction three times daily.\(^{(G6 \ G7)}\)

Root, tincture
5–10 mL (1 : 5 in 45% alcohol) three times daily.\(^{(G6 \ G7)}\)

Liquid Extract of Taraxacum
(BPC 1949) 2–8 mL.

Juice of Taraxacum
(BPC 1949) 4–8 mL.
Pharmacological Actions

In vitro and animal studies

A diuretic effect in rats and mice has been documented for dandelion extracts, following oral administration.\(^{(6)}\) Herb extracts were found to produce greater diuresis than root extracts; a dose of 50 mL (equivalent to 2 g dried herb/kg body weight) produced an effect comparable to that of furosemide 80 mg/kg. By contrast, no significant increases in urine volume or sodium excretion were observed in mice following oral administration of either leaf or root extracts, or of purified fractions.\(^{(3)}\) Similarly, oral and intravenous administration of an ethanolic extract of dandelion root failed to produce a diuretic effect in laboratory animals.\(^{(7)}\)

Moderate anti–inflammatory activity against carrageenan–induced rat paw oedema has been documented for a dandelion root extract.\(^{(8)}\) An 80% ethanol extract of root (100 mg/kg orally) inhibited oedema by 43% in the carrageenan–induced rat paw oedema test at 3 hours.\(^{(7)}\)

Bile secretion was doubled in dogs by a decoction of fresh root (equivalent to 5 g dried plant); similar activity has been observed for rats.\(^{(G52)}\)

Hypoglycaemic activity has been described in normal, but not in diabetic rabbits, following oesophageal administration of dandelion.\(^{(9)}\) Doses greater than 500 mg/kg produced a significant blood glucose concentration which had returned to normal after 24 hours. The maximum decrease produced by a dose of 2 g/kg was reported to be 65% of the effect produced by tolbutamide 500 mg/kg. Sulphonylureas (e.g. tolbutamide) act by stimulating pancreatic beta–cells and a similar mechanism was proposed for dandelion.

In vitro antitumour activity has been documented for an aqueous extract of dandelion, given by intraperitoneal injection, in the tumour systems ddY-Ehrlich and C3H/He-MM46.\(^{(10)}\) The mechanism of action was thought to be similar to that of tumour polysaccharides such as lentinan.

Clinical studies

There is a lack of well–designed clinical studies investigating the effects of dandelion.
Side-effects, Toxicity

Contact allergic reactions to dandelion have been documented\textsuperscript{(11,G51)} and animal studies have reported dandelion to have a weak sensitising capacity.\textsuperscript{(12)} Sesquiterpene lactones are thought to be the allergenic principles in dandelion.\textsuperscript{(4)} These compounds contain an exocyclic $\alpha$-methylene $\beta$-lactone moiety, which is thought to be a prerequisite for allergenic activity of sesquiterpene lactones.

The acute toxicity of dandelion appears to be low, with LD$_{50}$ values (mice, intraperitoneal injection) estimated at 36.8 g/kg and 28.8 g/kg for the root and herb, respectively.\textsuperscript{(6)} No visible signs of toxicity were observed in rabbits administered dandelion 3, 4, 5 and 6 g/kg body weight by mouth for up to seven days.\textsuperscript{(9)} In addition, no behavioural changes were recorded.
Contra-indications, Warnings

Treatment with dandelion is contraindicated for patients with occlusion of bile duct, gall bladder empyema and obstructive ileus.\(^{(G3 \ G52)}\) Dandelion may precipitate an allergic reaction in susceptible individuals, although no reports following the ingestion of dandelion have been documented. Dandelion may potentiate the action of other diuretics and may interfere with existing hypoglycaemic activity.

Pregnancy and lactation

There are no known problems with the use of dandelion during pregnancy, provided that doses do not greatly exceed the amounts used in foods.
Pharmaceutical Comment

Dandelion is a well–known traditional herbal remedy, although limited scientific information, particularly clinical research, is available to justify the reputed uses. Several investigations have failed to demonstrate significant diuretic effects in laboratory animals and have proposed that any diuretic activity is due to the high potassium content of the leaf and root. Dandelion has also been used in foods for many years. Animal studies indicate dandelion to be of low toxicity. However, excessive ingestion of dandelion, particularly in amounts exceeding those normally consumed in foods, should be avoided.
References


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Devil’s Claw
Species (Family)

Harpagophytum procumbens DC. (Pedaliaceae)
Synonym(s)
Harpagophytum, Grapple Plant, Wood Spider
Part(s) Used

Secondary root tuber
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)
BHP 1996\(^{(G9)}\)
BP 2002\(^{(G71)}\)
Complete German Commission E\(^{(G3)}\)
ESCOP 1996\(^{(G52)}\)
Martindale 33rd edition\(^{(G67)}\)
PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Ph Eur 2004\(^{(G72)}\)
Legal Category (Licensed Products)

Devil’s claw is not included in the GSL.\(^{(G37)}\)
Constituents


Carbohydrates
Fructose, galactose, glucose and myo-inositol (monosaccharides), raffinose, stachyose (46%) and sucrose (oligosaccharides).\(^1\)

Iridoids
Harpagide, 8-\(O\)-(\(p\)-coumaroyl)-harpagide, harpagoside, procumide, 6′-\(O\)-(\(p\)-coumaroyl)-procumbide, and procumboside (glucosides).\(^2\) Pharmacopoeial standard: not less than 1.2% harpagoside, calculated with reference to the dried drug.\(^{G15\ G25}\)

Phenols
Acetoside and isoacetoside (glycosides), and a bioside.\(^3\)

Other constituents
Amino acids and flavonoids (kaempferol, luteolin).

Other plant parts
The flower, stem and ripe fruit are reported to be devoid of harpagoside; the leaf contains traces of iridoids.\(^4\)
Food Use

Devil’s claw is not used in foods.
Herbal Use

Devil’s claw is stated to possess anti-inflammatory, antirheumatic, analgesic, sedative and diuretic properties. Traditionally, it has been used as a stomachic and a bitter tonic, and for arthritis, gout, myalgia, fibrositis, lumbago, pleurodynia and rheumatic disease. Modern use of devil’s claw is focused on its use in the treatment of rheumatic and arthritic conditions, and low back pain.
Dosage

Painful arthrosis and tendonitis

1.5–3 g dried tuber as a decoction, three times daily; 1–3 g drug or equivalent aqueous or hydroalcoholic extracts;\(^{(G52)}\) liquid extract 1–3 mL (1 : 1, 25% ethanol) three times daily.\(^{(G6)}\)

Loss of appetite or dyspepsia

**Dried tuber**

0.5 g as a decoction, three times daily.\(^{(G6)}\)

**Tincture**

1 mL (1 : 5, 25% ethanol) three times daily.\(^{(G6)}\)

Clinical trials of devil’s claw root extracts for the treatment of low back pain have tested doses ranging from 600 to 2400 mg daily, orally, in two or three divided doses (equivalent to up to 100 mg harpagoside (depending on the concentration of the extract)).\(^{(5–7)}\) In a clinical trial in osteoarthritis, participants received capsules containing powdered cryoground devil’s claw root 2610 mg daily.\(^{(8)}\) Other clinical trials in arthrosic conditions have used daily doses of devil’s claw of 2.4 g dried tuber and 2.46 g hydroalcoholic extract.\(^{(9)}\) Clinical trials in various rheumatic conditions have used daily doses of devil’s claw of 0.75–2 g dried tuber and 1.23 g aqueous extract in two or three divided doses.\(^{(9)}\)
Pharmacological Actions

The active constituents of devil’s claw are widely held to be the iridoid glucosides although, of these, it has not been definitively established whether harpagoside is the most important pharmacologically active constituent of the whole extract. Other compounds present in the root may contribute to the pharmacological activities of devil’s claw.\(^{(9,10)}\) It has also been suggested that harpagogenin, formed by \textit{in vivo} acid hydrolysis of harpagoside, may have biological activity.\(^{(11)}\)

\textbf{In vitro and animal studies}

Animal studies using aqueous extracts of devil’s claw have suggested that the extract may be inactivated by passage through the acid environment of the stomach.\(^{(10,12)}\) One study compared the anti-inflammatory activities of aqueous devil’s claw extract administered by different routes. Intraperitoneal and intraduodenal administration led to a significant reduction in the carrageenan-induced rat paw oedema test, but there was no effect following oral administration.\(^{(12)}\) In another study, aqueous devil’s claw extract pretreated with hydrochloric acid to mimic acid conditions in the stomach showed no activity in pharmacological models of pain and inflammation.\(^{(10)}\)

Transformation of the iridoids harpagide, harpagoside and 8-\(O-(p\)-coumaroyl)-harpagide into the pyridine monoterpene alkaloid aucubinine B, chemically or by human intestinal bacteria \textit{in vitro}, has been documented.\(^{(13,14)}\) However, it is not known if aucubinine B is formed \textit{in vivo} by intestinal bacteria and, therefore, whether it contributes to the pharmacological activity of devil’s claw.\(^{(14)}\)

Animal studies of the anti-inflammatory activity of devil’s claw have reported conflicting results. Activity differs depending on the route of administration of devil’s claw, and the model of inflammation, whether acute or subacute.

Weak anti-inflammatory activity has been reported in rats following intravenous administration of devil’s claw extract.\(^{(15)}\) Anti-inflammatory activity of harpagoside has been demonstrated in experimental models, including the croton oil-induced granuloma pouch test, and for harpagogenin, the aglucone of harpagoside, in the croton oil-induced granuloma pouch test and in formalin-induced arthritis in rats.\(^{(16)}\) Dried aqueous extract of devil’s claw administered by intraperitoneal injection demonstrated significant activity in the carrageenan-induced oedema test in rats, an acute model of inflammation.\(^{(10)}\) The effect on oedema was dose dependent for doses of devil’s claw extract 100–400 mg/kg, and reached a maximum 3 hours after
carrageenan injection. Other studies in rats have reported significant reductions in oedema using the same model following pretreatment with intraperitoneal\(^{(12,17)}\) and intraduodenal, but not oral, dried aqueous extract of devil’s claw.\(^{(12)}\) Other studies have reported that dried aqueous extract of devil’s claw administered orally had no effect on carrageenan- or *Mycobacterium butyricum*-induced oedema in rat paw.\(^{(18,19)}\) In addition, oral dried aqueous extract of devil’s claw had no significant effect in adjuvant-induced arthritis in rats.\(^{(18)}\) By contrast, in these studies, both indometacin and aspirin displayed significant anti-inflammatory activity.\(^{(18,19)}\)

Analgesic activity has also been documented for devil’s claw in animal studies. Pretreatment with dried aqueous devil’s claw extract at doses of 100 mg/kg and above, administered intraperitoneally, resulted in peripheral analgesic activity demonstrated by a significant reduction in the number of writhings induced by acetic acid in mice.\(^{(10)}\) However, no effect was observed in the hotplate test, indicating a lack of central analgesic activity with devil’s claw extract. The peripheral analgesic properties of intraperitoneal dried aqueous extract of devil’s claw have been confirmed in other studies for doses of 400 mg/kg and above.\(^{(17)}\) These studies also reported peripheral analgesic and anti-inflammatory properties for the related species *Harpogophytum zeyheri*.\(^{(17)}\)

A clear mechanism of action for the purported anti-inflammatory effects of devil’s claw has yet to be established. *In vitro*, devil’s claw (100 mg/mL) had no significant effect on prostaglandin (PG) synthetase activity, whereas indometacin (316 μg/mL) and aspirin (437 μg/mL) caused 50% inhibition of this enzyme.\(^{(19)}\) In other *in vitro* studies in human whole blood samples, devil’s claw extracts and fractions of extracts were tested for their effects on thromboxane B\(_2\) (TXB\(_2\)) and leukotriene (LT) biosynthesis.\(^{(20)}\) TXB\(_2\) is an end-product of arachidonic acid metabolism by the cyclooxygenase 1 (COX-1) pathway. Inhibition appeared to be dependent on the harpagoside content of the extracts or fractions.\(^{(20)}\) Harpagoside (100 μmol/L), but not harpagide (100 μmol/L), inhibited calcium ionophore A23187–stimulated release of TXB\(_2\) from human platelets.\(^{(21)}\) However, harpagoside and harpagide had no significant inhibitory effect on calcium ionophore A23187–stimulated release of PGE\(_2\) and LTC\(_4\) from mouse peritoneal macrophages.\(^{(21)}\) In *in vitro* inhibition of tumour-necrosis-factor-α (TNF-α) synthesis in lipopolysaccharide–stimulated human monocytes by a hydroalcoholic extract of devil’s claw (SteHap 69) has also been documented.\(^{(22)}\)

Crude methanolic extracts of devil’s claw have been shown to be cardioactive *in vitro* and *in vivo* in animals. A protective action against ventricular
Arrhythmias induced by aconitine, calcium chloride, epinephrine (adrenaline)/chloroform and reperfusion has been reported for devil’s claw given intraperitoneally or added to the reperfusion medium.\(^{23,24}\) The crude extract was found to exhibit greater activity than pure harpagoside.\(^{24}\) In isolated rabbit heart, low concentrations of a crude methanolic extract had mild negative chronotropic and positive inotropic effects,\(^{23}\) whereas high concentrations caused a marked negative inotropic effect with reduction in coronary blood flow.\(^{23}\) In anaesthetised dogs, harpagoside administered orally by gavage caused a decrease in mean aortic pressure and arterial and pulmonary capillary pressure.\(^{25}\)

*In vitro*, harpagoside has been shown to decrease the contractile response of smooth muscle to acetylcholine and barium chloride on guinea–pig ileum and rabbit jejunum.\(^{26}\) Harpagide was found to increase this response at lower concentrations, but antagonised it at higher concentrations.\(^{26}\) On the basis of these studies in isolated smooth muscle, it was suggested that the constituents of devil’s claw may influence mechanisms regulating calcium influx.\(^{26}\)

Methanolic extracts have also exhibited hypotensive properties in normotensive rats, causing a decrease in arterial blood pressure following oral doses of 300 mg/kg and 400 mg/kg body weight.\(^{23}\)

Devil’s claw extracts possess weak antifungal activity against *Penicillium digitatum* and *Botrytis cinerea*.\(^{27}\)

**Clinical studies**

**Pharmacokinetics**

There is little published information on the pharmacokinetics of devil’s claw extract in humans. A pharmacokinetic study involving a small number of healthy male volunteers \((n = 3)\) measured plasma harpagoside concentrations after oral administration of devil’s claw extract (WS1531 containing 9% harpagoside) 600, 1200 and 1800 mg as film–coated tablets.\(^{20}\) Maximal plasma concentrations of harpagoside were reached after 1.3–1.8 hours, and were 8.2 ng/mL and 27.8 ng/mL for doses of harpagoside of 108 and 162 mg, respectively (corresponding to 1200 and 1800 mg devil’s claw extract, respectively). Other studies involving small numbers of healthy male volunteers indicated that the half–life ranged between 3.7 and 6.4 hours. Other results suggested that there may be low oral absorption or a considerable first–pass effect with devil’s claw extract, although this needs further investigation.\(^{20}\)
Pharmacodynamics

A study involving healthy volunteers investigated the effects on eicosanoid production of orally administered devil’s claw (four 500-mg capsules of powder, containing 3% glucoiridoids, daily for 21 days).\(^{28}\) No statistically significant differences on PGE\(_2\), TXB\(_2\), 6-keto-PGF\(_{1\alpha}\) and LTB\(_4\) were observed following the period of devil’s claw administration, compared with baseline values. By contrast, in a subsequent study involving whole blood samples taken from healthy male volunteers, a biphasic decrease in basal cysteinyl-leukotriene (Cys-LT) biosynthesis, compared with baseline values, was observed following oral administration of devil’s claw extract (WS1531 containing 9% harpagoside) 600, 1200 and 1800 mg as film-coated tablets.\(^{20}\)

Therapeutic activity

The efficacy and effectiveness of devil’s claw has been investigated in more than 10 clinical studies involving patients with rheumatic and arthritic conditions, and low back pain.\(^{9,29}\) These studies have involved different methodological designs, including several uncontrolled studies, and different preparations of devil’s claw, including crude drug and aqueous extracts. These studies have been summarised elsewhere,\(^{9,29,56}\) and several are discussed in detail below.

A randomised, double-blind, placebo-controlled study involving 118 patients with acute exacerbations of chronic low back pain investigated the effects of devil’s claw extract 800 mg three times daily (equivalent to 50 mg harpagoside daily) for four weeks.\(^{7}\) There was no statistically significant difference between the devil’s claw and placebo groups in the primary outcome measure – consumption of the opioid analgesic tramadol over weeks 2–4 of the study – among the 109 patients who completed the study. This was an unusual choice of primary outcome measure as it gives no direct indication of the degree of pain experienced by participants. There was a trend towards improvement in a modified version of the Arhus Low Back Pain Index (a measure of pain, disability and physical impairment) for devil’s claw recipients compared with placebo recipients, although this did not reach statistical significance. A greater proportion of patients in the devil’s claw group were pain-free at the end of the study, although this was only a secondary outcome measure.

On the basis of these findings, a subsequent randomised, double-blind, placebo-controlled trial involving 197 patients with exacerbations of low back pain tested the effects of two doses of devil’s claw (WS1531) extract against placebo.\(^{6}\) Participants received devil’s claw extract 600 mg or 1200 mg daily (equivalent to 50 mg and 100 mg harpagoside daily,
respectively), or placebo, for four weeks. There was a statistically significant difference \( (p = 0.027) \) between devil’s claw and placebo with respect to the primary outcome measure – the number of patients who were pain-free without tramadol for at least five days during the last week of the study. However, numbers of patients who were pain-free were low (3, 6 and 10 for placebo, devil’s claw 600 mg daily and devil’s claw 1200 mg daily, respectively). Furthermore, this is a non-standard outcome measure. Arhus Low Back Pain Index scores improved significantly in all three groups, compared with baseline values, although there was no statistically significant difference between groups.

In a randomised, double-blind, placebo-controlled study involving patients with non-specific low back pain, 65 participants received devil’s claw extract (LI-174, Rivoltan), or placebo, 480 mg twice daily (equivalent to 24 mg harpagoside daily) for four weeks.\(^5\) There was a significant improvement \( (p < 0.001) \) in visual analogue scale (VAS) scores for muscle pain in the devil’s claw group, but not the placebo group, compared with baseline values, after two and four weeks’ treatment. Differences in VAS scores between the two groups were statistically significant after four weeks’ treatment \( (p < 0.001) \). Significant differences between the two groups in favour of devil’s claw after four weeks’ treatment were also observed with several other parameters, including muscle stiffness and muscular ischaemic pain.

A randomised, double-blind trial has compared the efficacy of devil’s claw extract with that of diacerein in 122 patients with osteoarthritis of the knee and hip.\(^8\) Participants received powdered cryoground devil’s claw (Harpadol) 2.61 g daily, or diacerein 100 mg daily, for four months. VAS scores for spontaneous pains improved significantly in both groups, compared with baseline values, and there were no differences between devil’s claw and diacerein with respect to VAS scores.

In a placebo-controlled study involving 89 patients with rheumatic complaints, devil’s claw recipients (who received powdered crude drug 2 g daily for two months) showed significant improvements in sensitivity to pain and in motility (as measured by the finger-to-floor distance), compared with placebo recipients.\(^30\)

Open, uncontrolled studies involving patients with rheumatic and arthritic disorders report conflicting results for the effectiveness of devil’s claw. One study involved 13 patients with arthritis, rheumatoid arthritis or psoriatic arthropathy who received tablets of devil’s claw aqueous extract 1.23 g daily for six weeks in addition to their conventional drug treatment. There were no significant changes after 6 and 12 weeks in pain, early morning stiffness, and
the Ritchie Articular Index (a method of assessing joint tenderness), compared with baseline values.\(^{31}\) By contrast, other open uncontrolled studies of devil’s claw involving patients with rheumatic disorders (who received devil’s claw powder 1.5 g daily for 60 days)\(^{32}\) or arthrosis (who received devil’s claw aqueous extract, containing 2.5% iridoid glycosides, 3–9 g daily for 6 months)\(^{33}\) reported improvements in pain and ‘complaints’ at the end of the treatment period compared with baseline values.

Another study involved 45 patients with osteo- or rheumatoid arthritis who received devil’s claw root extract 2.46 g daily for two weeks in addition to non-steroidal anti-inflammatory drug (NSAID) treatment, followed by devil’s claw extract alone, for four weeks.\(^{34}\) It was reported that there were no statistically significant changes in pain intensity and duration of morning stiffness during the period of treatment with devil’s claw extract alone. In subgroups of patients with rheumatoid arthritis and those with osteoarthritis, small decreases were observed in concentrations of C-reactive protein and creatinine, respectively. The design of this study in terms of the treatment regimen (NSAID followed by devil’s claw extract without a washout period), however, renders the results difficult to interpret.
Side-effects, Toxicity

Randomised, placebo-controlled trials involving patients with rheumatic and arthritic conditions who have received devil’s claw extracts or powdered drug at approximately recommended doses for four weeks have reported mild, transient gastrointestinal symptoms (such as diarrhoea, flatulence) in a small proportion (less than 10%) of devil’s claw recipients.\(^{(5-7)}\) No serious adverse events were reported, although one patient withdrew from one study because of tachycardia.\(^{(7)}\) In an open, uncontrolled study, one patient withdrew after four days’ treatment with devil’s claw aqueous extract 1.23 g daily because of several symptoms, including frontal headache, tinnitus, anorexia and loss of taste.\(^{(31)}\)

In a randomised, controlled trial comparing devil’s claw extract with diacerein in patients with osteoarthritis, numbers of patients ending the study prematurely because of suspected adverse drug reactions were 8 and 14 for devil’s claw and diacerein recipients, respectively.\(^{(8)}\) In total, 26 diacerein recipients and 16 devil’s claw recipients reported one or more adverse events \((p = 0.042)\). The numbers of adverse events attributed to the treatment was significantly lower for devil’s claw than for diacerein \((10 \text{ versus } 21; p = 0.017)\). The most frequently reported adverse event, diarrhoea, occurred in 8.1% and 26.7% of devil’s claw and diacerein recipients, respectively.

There is an isolated report of conjunctivitis, rhinitis and respiratory symptoms in a 50-year-old woman who had experienced chronic occupational exposure to devil’s claw.\(^{(35)}\)

The mechanism of action of devil’s claw remains unclear, in particular, whether it has significant effects on the mediators of acute inflammation. Data from \textit{in vitro} and clinical studies in this regard do not yet give a clear picture (\textit{see In vitro and animal studies and Clinical studies, Pharmacodynamics}). It has been stated that adverse effects associated with the use of NSAIDs are unlikely to occur with devil’s claw, even during long-term treatment.\(^{(G50 \ G52)}\) While there are no documented reports of gastrointestinal bleeding or peptic ulcer associated with the use of devil’s claw, the latter statement requires confirmation. Use of devil’s claw in gastric and duodenal ulcer is contraindicated, although this appears to be because of the drug’s bitter properties.\(^{(G50)}\)

Acute and subacute toxicity tests in rodents have demonstrated low toxicity of devil’s claw extracts. In a study in mice, the acute oral lethal dose \((LD) LD_0\) and \(LD_{50}\) were greater than 13.5 g/kg body weight.\(^{(19)}\) In rats, clinical, haematological and gross pathological findings were unremarkable following
administration of devil’s claw extract 7.5 g/kg by mouth for seven days. Hepatic effects (liver weight, and concentrations of microsomal protein and several liver enzymes) were not observed following oral treatment with devil’s claw extract 2 g/kg for seven days.\(^{(19)}\) Other studies in mice have reported acute oral acute intravenous LD\(_0\) values of greater than 4.64 g/kg and greater than 1 g/kg, respectively.\(^{(15)}\) For an extract containing harpagoside 85%, acute oral LD\(_0\), acute intravenous LD\(_0\) and acute intravenous LD\(_{50}\) values were greater than 4.64 g/kg, 395 mg/kg and 511 mg/kg, respectively.\(^{(15)}\)
Contra–indications, Warnings

Devil’s claw is stated to be contra–indicated in gastric and duodenal ulcers,\(^{(G3 \ G52)}\) and in gallstones should be used only after consultation with a physician.\(^{(G3)}\) On the basis of pharmacological evidence of devil’s claw’s cardioactivity, the possibility of excessive doses interfering with existing treatment for cardiac disorders or with hypo/hypertensive therapy should be considered.

**Pregnancy and lactation**

It has been stated that devil’s claw has oxytocic properties,\(^{(36)}\) although the reference gives no further details and the basis for this statement is not known. In addition, there is no further evidence to substantiate the statement. However, given the lack of data on the effects of devil’s claw taken during pregnancy and lactation, its use should be avoided during these periods.
Pharmaceutical Comment

The chemistry of devil’s claw has been well documented. The iridoid constituents are thought to be responsible for the reputed anti-inflammatory activity of devil’s claw, although it is not known precisely which of these are the most important for pharmacological activity, and the importance of other compounds. There is conflicting evidence from in vitro, animal and human studies regarding the anti-inflammatory activity of devil’s claw and possible mechanisms of action. Several randomised trials using devil’s claw extracts standardised on harpagoside content have reported superiority over placebo for some aspects of low back pain and rheumatic complaints. However, some studies used non-standard outcome measures and carried out several post-hoc analyses. Further studies have used recognised, predefined outcome measures to establish the therapeutic value of standardised devil’s claw extracts in patients with arthritic and rheumatic conditions.

On the basis of randomised controlled trials involving patients with arthritic and rheumatic disorders, devil’s claw extracts appear to have a favourable short-term adverse effect profile when taken in recommended doses. Mild, transient gastrointestinal effects, such as diarrhoea and flatulence, may occur. Chronic toxicity studies and clinical experience with prolonged use are lacking, so the effects of long-term use are not known. On this basis, and in view of the possible cardioactivity of devil’s claw, devil’s claw should not be used for long periods of time at doses higher than recommended. Further studies involving large numbers of patients are required.

Some commercial extracts of devil’s claw root may have been prepared not only from the roots of *H. procumbens*, but also from the roots of *H. zeyheri*, which are similar macroscopically. However, the two species differ in the concentration of the constituents harpagoside and 8-0-p-coumaroyl-harpagide. On this basis it has been stated that the species can be distinguished chemically by determining the ratio harpagoside:8-0-p-coumaroyl-harpagide. The ratio is stated to be near one for *H. zeyheri* and between 20 and 38 for *H. procumbens* which has a low 8-0-p-coumaroyl-harpagide content. While this ratio may be sufficient for chemotaxonomic differentiation, it may not be adequate for quality control. Other studies have demonstrated that the harpagoside content of several powdered dry extracts of devil’s claw from different manufacturers varies, and that each extract has a unique profile of other constituents.
References


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Drosera
Species (Family)

Drosera rotundifolia L. (Droserceae)
Synonym(s)

Sundew
Part(s) Used

Herb
Pharmacopoeial and Other Monographs

BHP 1983$^{(G7)}$

Complete German Commission E (Sundew)$^{(G3)}$

Martindale 33rd edition$^{(G67)}$

PDR for Herbal Medicines 2nd edition (Sundew)$^{(G36)}$
Legal Category (Licensed Products)

Drosera is not included in the GSL.\(^{(G37)}\)
Constituents

See General References G2 G40 G48 G53 G64.

Flavonoids
Kaempferol, myricetin, quercetin and hyperoside.\(^{(1)}\)

Quinones
Plumbagin,\(^{(2)}\) hydroplumbagin glucoside\(^{(3)}\) and rossoliside (7–methyl–hydrojuglone–4–glucoside).\(^{(4)}\)

Other constituents
Carotenoids, plant acids (e.g. butyric acid, citric acid, formic acid, gallic acid, malic acid, propionic acid), resin, tannins (unspecified) and ascorbic acid (vitamin C).
Food Use

Drosera is not used in foods.
Herbal Use

Drosera is stated to possess antispasmodic, demulcent and expectorant properties. It has been used for bronchitis, asthma, pertussis, tracheitis, gastric ulceration and specifically for asthma and chronic bronchitis with peptic ulceration or gastritis.\(^{(G2 \text{ G7 G64})}\)
Dosage

**Dried plant**
1–2 g or by infusion three times daily.\(^{(G7)}\)

**Liquid extract**
0.5–2.0 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)

**Tincture**
0.5–1.0 mL (1 : 5 in 60% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

**In vitro and animal studies**

Drosera is reported to prevent acetylcholine- or histamine–induced broncho spasm, and to relax acetyl choline- or barium chloride–induced spasm of the isolated intestine.\(^{(5)}\) Drosera is stated to possess antitussive properties and has been reported to prevent coughing induced by excitation of the larynx nerve in the rabbit.\(^{(5)}\) These antispasmodic actions have been attributed to the naphthoquinone constituents.\(^{(G53)}\)

Antimicrobial properties have also been documented for the naphthoquinones.\(^{(6)}\) *In vivo*, plumbagin is reported to exert a broad spectrum of activity against Gram–positive and Gram–negative bacteria, influenza viruses, pathogenic fungi, and parasitic protozoa. *In vitro*, a plumbagin solution (1 : 50 000) was reported to exhibit activity against staphylococci, streptococci and pneumococci (Gram–positive bacteria), but to lack activity against *Haemophilus pertussis* (Gram–negative bacteria).\(^{(5)}\) Plumbagin administered orally to mice for five days, was found to be ineffective against *Lamblia muris* and tuberculosis infection. *Microsporum* infections in guinea–pigs were treated successfully by local applications of 0.25–0.5% solutions (in 40% alcohol) or of 1% emulsions.\(^{(6)}\)

An aqueous drosera extract was reported to possess pepsin–like activity.\(^{(G53)}\)

*In vitro*, drosera extracts and plumbagin, in concentrations of 0.01–1.0 mg/mL, have been documented to exert a cytotoxic or immunosuppressive effect in human granulocytes and lymphocytes.\(^{(2)}\) Lower concentrations were reported to exhibit immunostimulating properties. Plumbagin possesses chemotherapeutic properties, but is irritant when administered at therapeutic doses.\(^{(6)}\)
Side-effects, Toxicity

None documented for drosera. Plumbagin is stated to be an irritant principle\(^{G51}\) and an LD\(_{50}\) (mice, intraperitoneal injection) has been reported to be 15 mg/kg body weight.\(^{G48}\)

Cytotoxic properties have been documented for drosera and plumbagin (see *In vitro* and animal studies).
Contra-indications, Warnings

None documented.

**Pregnancy and lactation**  
The safety of drosera has not been established. In view of the lack of toxicity data, the use of drosera during pregnancy and lactation should be avoided.
Pharmaceutical Comment

Limited chemical information is available for drosera. Documented animal studies support some of the herbal uses. Reported immunostimulant and immunosuppressant activities may warrant further research into the pharmacological activities of drosera. In view of the lack of chemical and toxicity data, excessive use of drosera should be avoided.
References

See also General References G2 G3 G7 G31 G36 G37 G40 G43 G45 G48 G51 G53 G64.

Echinacea
Species (Family)

i. *Echinacea angustifolia* (DC.) Hell. (Asteraceae/Compositae)

ii. *Echinacea pallida* (Nutt.) Britt.

iii. *Echinacea purpurea* (L.) Moench
Synonym(s)

Black Sampson, Coneflower

i. *Brauneria angustifolia*

ii. *Brauneria pallida* (Nutt.) Britt.
Part(s) Used

Rhizome, root. *E. purpurea* herb (aerial parts) is also used
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHMA 2003\(^{(G66)}\)

BHP 1996\(^{(G9)}\)

Complete German Commission E 1998\(^{(G3)}\)

ESCOP 1999\(^{(G52)}\)

Expanded German Commission E 2000\(^{(G4)}\)

Martindale 33rd edition\(^{(G67)}\)

Mills and Bone\(^{(G50)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)

WHO volume 1 1999\(^{(G63)}\)
Legal Category (Licensed Products)

GSL$^{(\text{G37})}$
Constituents

See References 1–5, and General References G2, G6, G52, G63 and G69.

Alkylamides

At least 20, mainly isobutylamides of straight-chain fatty acids with olefinic and/or acetylenic bonds,\(^{(1,6-9)}\) e.g. isomeric dodeca-\(2E,4E,8Z,10E/Z\)-tetraenoic isobutylamide,\(^{(10)}\) present in the roots and aerial parts of *Echinacea angustifolia* and *Echinacea purpurea*, but mainly absent from *Echinacea pallida*. Isobutylamides from the roots of *E. purpurea* contain mainly 2,4-dienoic units whilst those of *E. angustifolia* contain mainly 2-monoene units.\(^{(8)}\) The synthesis of the acetylenic amide \(N-(2\text{-methylpropyl})-2E\text{-undecene-8,10-diynamide, a constituent of } E. \text{ angustifolia root, has been reported.}^{(11)}\) *E. purpurea* root reportedly contains 0.01–0.04% alkylamides.\(^{(G52)}\)

Phenylpropanoids

Caffeic acid glycosides (e.g. echinacoside, verbascoside, caffeoylechinacoside), caffeic acid esters of quinic acid (e.g. chlorogenic acid = 5-caffeoylquinic acid, isochlorogenic acid = 3,4- and 3,5-dicaffeoylquinic acid, cynarin = 1,3-dicaffeoylquinic acid) and of tartaric acid (e.g. caftaric acid = 2-caffeoyltartaric acid, cichoric acid = 2,3-dicaffeoyltartaric acid).\(^{(12)}\) Varying mixtures of caffeic acid derivatives are present in the three species, with echinacoside being the major component of the roots of *E. angustifolia* and *E. pallida*\(^{(12)}\) (0.5–1.0%),\(^{(G52)}\) and cichoric acid being a major component of *E. purpurea* roots (0.14–2.05%),\(^{(13)}\) and aerial parts (1.2–3.1%).\(^{(14,G52)}\) Cynarin is reportedly present in *E. angustifolia* root,\(^{(10,12)}\) but not in the roots of the other two species.

Polysaccharides

Polysaccharides PS1 (a methylglucuron-arabinoxylan, mol. wt 35 kDa), PS2 (an acidic rhamnoarabinogalactan, mol. wt 450 kDa) and a xyloglucan (mol. wt 79 kDa) have been isolated from *E. purpurea* herb.\(^{(5,G52)}\) Polysaccharides and glycoproteins are present in *E. purpurea* herb and *E. pallida* root.\(^{(G52)}\) The pressed juice from the aerial parts of *E. purpurea* (and the herbal medicinal product Echinacin prepared from the juice) contain heterogeneous polysaccharides (mol. wt <10 kDa), inulin-type fractions (mol. wt 6 kDa) and an acidic highly branched arabinogalactan polysaccharide (mol. wt 70 kDa).\(^{(15)}\) The pressed juice of *E. purpurea* aerial parts has yielded an arabinogalactan-protein (AGP) comprising 83% polysaccharide (galactose-arabinose 1.8 : 1), uronic acids (4–5%) and protein (7%) with high concentrations of serine, alanine and hydroxyproline.\(^{(16)}\) The AGP (mol. wt
1.2 \times 10^6 \text{ Da}) has a highly branched polysaccharide core of 3-, 6-, and 3,6-linked galactose residues with terminal arabinose and glucuronic acid units.\(^{(16)}\)

**Volatile oils**

*E. pallida* root (0.2–2.0\%)\(^{(G52)}\) mainly contains polyenes and polyacetylenes including pentadeca-1,8Z-diene and a range of ketoalkenes and ketoalkenynes (ketopolyacetylenes), principally pentadeca-8Z-ene-2-one, pentadeca-8Z,11Z-diene-2-one, pentadeca-8Z,13Z-diene-11yne-2-one, tetradeca-8Z-ene-11,13-diyne-2-one and others.\(^{(17,G52)}\) These alkenes are unstable and readily oxidise to 8-hydroxy derivatives.\(^{(G52)}\) The alkenes of *E. pallida* and *E. purpurea* root are distinctly different from those of *E. angustifolia* which are mainly alkylketones.\(^{(9)}\) The volatile oil from the aerial parts of the three species contains borneol, bornyl acetate, germacrene D, caryophyllene and other components.\(^{(G2,G52)}\)

**Other constituents**

A series of other constituents has been reported including the saturated pyrrolizidine-type alkaloids tussilagine and isotussilagine (0.006\%) from *E. angustifolia* and *E. purpurea*.\(^{(18)}\) Flavonoids, including quercetin, kaempferol, isorhamnetin and their glycosides\(^{(G52)}\) and also anthocyanins, are present in the aerial parts of *E. purpurea* (0.48\%).\(^{(G2)}\) The major flavonoid of the aerial parts of *E. angustifolia* has been identified as patuletin-3-rutinoside,\(^{(19)}\) and not rutin as previously reported.\(^{(20)}\) Free phenolic acids, including \(p\)-coumaric, \(p\)-hydroxybenzoic and protocatechuic acids, have been isolated from the aerial parts of *E. angustifolia* and *E. purpurea*.\(^{(21)}\) Other miscellaneous compounds reported include betaine, fatty acids, simple sugars, sterols and vanillin.

**Quality of plant material and commercial products**

Alkylamide concentrations vary between species and between different parts of the plant.\(^{(22)}\) Commercial root samples of *E. purpurea* have been shown to vary in their alkylamide content (0.12–1.2\%).\(^{(14)}\) In Germany, 25 commercial echinacea preparations were assayed for their alkylamide (dodeca-2E,4E,8Z,10E/Z-tetraenoic acid isobutylamide) and cichoric acid contents.\(^{(23)}\) Some products were highly concentrated, whereas others had no detectable concentrations of alkylamide or cichoric acid. Large differences were observed between comparable products from different manufacturers.

Of 25 commercial echinacea products purchased in the USA only 14 (56\%) passed assessments for their quality.\(^{(24)}\) Six were inadequately labelled, three of them not stating the species used, one not stating the plant part and two
liquid preparations had no concentrations given for their echinacea content. The remaining 19 products were assessed for their stated content of particular species and for claimed concentrations of phenols. Twelve of these products were labelled as containing only *E. purpurea* and two of them failed, as one contained only 54% of the expected concentration of phenols and the other had three times the accepted concentration of microbes as set out in World Health Organization (WHO) guidelines. Two products were allegedly prepared from *E. angustifolia* and both failed, one having only one-third of the stated phenolic content and the other having no detectable echinacoside. Five further products allegedly containing a mixture of species were also assessed and one failed because echinacoside could not be detected.

Analysis of 59 commercial products available in the US revealed that 10% had no measurable echinacea content, 48% were not consistent with their labels in respect of the species present, and of 21 standardised preparations, 57% did not meet the standards stated on their labels; often products did not contain the species stated.\(^{(25)}\)

A fresh plant product of echinacea herb has been shown to possess three times the amount of alkylamide than a product prepared from dried plants and this has been attributed to loss on drying.\(^{(26)}\) The alkylamide and cichoric acid content of six commercial preparations of *E. purpurea* expressed juice have been shown to be variable (0.1–1.8 mg/mL and 0.0–0.4%, respectively).\(^{(27)}\) Ten commercial preparations of echinacea were analysed for their betaine content and concentrations ranged from 0.04 to 0.64%.\(^{(28)}\)

The concentrations of some constituents may be affected during growing, drying or storage of the plant material. The yields of some constituents are affected when plants are grown under conditions of drought stress.\(^{(29)}\) Analysis of roots of *E. angustifolia* dried at a range of temperatures between 23°C and 60°C indicated that there were no significant changes in the alkylamide content, whereas 25% and 45% of the echinacoside content was lost at 30°C and 60°C, respectively.\(^{(30)}\) By contrast, roots of *E. purpurea* at −18°C in deep-freeze for 64 weeks were found to have lost 40% of their alkylamide content.\(^{(31)}\) An aqueous–alcoholic extract of *E. purpurea* and its dried extract were stored at different temperatures for seven months and then assayed for their alkylamide and phenylpropanoid content.\(^{(32)}\) The amount of the major alkylamide (dodeca-2E,4E,8Z,10E/Z-tetraenoic acid isobutylamide) in the liquid preparation was not significantly affected by storage at 25°C and 40°C, whereas the cichoric acid content declined. However, the reverse occurred for the dried extract when there was a significant loss of alkylamide at storage temperatures of 25°C and 40°C but no significant loss of cichoric acid content.
The effects of drying temperatures on the constituents of all three echinacea species have been investigated.\textsuperscript{(33)} The results showed that there was an increase in cichoric acid for \textit{E. purpurea} and \textit{E. pallida}. Furthermore, increased moisture content resulted in higher concentrations of echinacoside for \textit{E. angustifolia} and \textit{E. pallida} and of chlorogenic acid in \textit{E. angustifolia}. The polysaccharide contents were significantly decreased by raised moisture levels in the roots of \textit{E. angustifolia} and \textit{E. pallida}.

The presence of colchicine in commercial echinacea products in the USA has been reported,\textsuperscript{(34)} although subsequent analysis of 17 commercial echinacea products purchased in pharmacies in Chicago, USA, did not detect colchicine in any of the samples.\textsuperscript{(35)}
Herbal Use

Echinacea has a long history of medicinal use for a wide variety of conditions, mainly infections, such as syphilis and septic wounds, but also as an ‘anti-toxin’ for snakebites and blood poisoning.\textsuperscript{(36,G50)} Traditionally, echinacea was known as an ‘anti-infective’ agent, and was indicated in bacterial and viral infections, mild septicaemia, furunculosis (persistent recurring episodes of painful nodules in the skin) and other skin conditions, including boils, carbuncles and abscesses.\textsuperscript{(G6, G7, G60, G69)} Other traditional uses listed include naso-pharangeal catarrh, pyorrhoea (periodontitis) and tonsillitis, and as supportive treatment for influenza-like infections and recurrent infections of the respiratory tract and lower urinary tract and, externally, for poorly healing superficial wounds.\textsuperscript{(G66)}

Current interest in the medicinal use of echinacea is focused on its immunomodulatory or immunostimulant effects, particularly in the treatment and prevention of the common cold, influenza and other upper respiratory tract infections (see Clinical studies).
Dosage

In standard herbal reference texts

Dosages for oral administration (adults) recommended in older standard herbal reference texts are the same for several indications; examples are given below.

E. angustifolia root\(^{(G6)}\) and/or E. pallida root\(^{(G7)}\)
For various indications, including chronic viral and bacterial infections, skin complaints, prophylaxis of colds and influenza\(^{(G6)}\), mild septicaemia, furunculosis,\(^{(G6,G7)}\) naso-pharyngeal catarrh, pyorrhoea and tonsillitis.\(^{(G7)}\)

Dried root/rhizome
1 g by infusion or decoction three times daily.\(^{(G6,G7)}\)

Liquid extract
0.5–1.0 mL (1 : 5 in 45% alcohol) three times daily,\(^{(G6)}\) or 0.25–1.0 mL (1 : 1 in 45% alcohol) three times daily.\(^{(G7)}\)

Tincture
2–5 mL (1 : 5 in 45% alcohol) three times daily,\(^{(G6)}\) or 1–2 mL (1 : 5 in 45% alcohol) three times daily.\(^{(G7)}\)

In more recent texts

Dosages for oral administration (adults) described in more recent texts are provided for more specific indications.

As adjuvant therapy and for prophylaxis of recurrent infections of the upper respiratory tract (common colds); treatment should not exceed eight weeks’ duration.\(^{(G3,G52)}\)

E. pallida root
Hydroethanolic extract corresponding to 900 mg crude drug daily,\(^{(G52)}\) e.g. tincture (1 : 5 in 50% ethanol by volume) from dry extract (7–11 : 1 in 50% ethanol).\(^{(G3)}\)

E. purpurea herb
6–9 mL expressed juice daily.\(^{(G3,G52)}\)

E. purpurea root
3 × 60 drops of tincture (1 : 5 in 55% ethanol), equivalent to 3 × 300 mg
crude drug daily.\textsuperscript{G52}

\textbf{E. angustifolia root}

1–3 g daily

\textbf{In clinical trials}

Echinacea preparations (i.e. containing different echinacea species and plant parts) and, therefore, dosage regimens tested in clinical trials have varied widely (see Clinical studies). Trials of echinacea preparations for the prevention of upper respiratory tract infections have typically involved an 8- or 12-week duration of treatment; trials of echinacea preparations for the treatment of upper respiratory tract infections typically involve administration of the study medication for 6–10 days.
Pharmacological Actions

There is a vast scientific literature on the pharmacological activities of echinacea. Research has focused on investigating the immunomodulatory activity of echinacea preparations, although other activities, such as antiviral, antifungal, anti-inflammatory and antioxidant properties have also been explored. Effects on the immune system may play a role in some of these other activities.

The pharmacological activities of echinacea preparations cannot be attributed to a single constituent or group of constituents. Rather, several groups of constituents – the alkylamides, caffeic acid derivatives, polysaccharides and polyenes – appear to be important for activity.

In vitro and animal studies

Immunomodulatory activity

Currently, there is a view that immunomodulatory, rather than immunostimulatory, is the most appropriate term to describe the immunological effects of echinacea,\(^ {37}\) although ‘immunostimulatory’ is still used and is ubiquitous in the earlier scientific literature on echinacea. It has been suggested that broad stimulation of the various highly complex components of the immune system is unlikely to be beneficial, since some immune responses are harmful.\(^ {37}\)

The immunological effects of a wide range of echinacea preparations comprising different species, plant parts and types of extract, have been investigated extensively in vitro and in vivo. Collectively, the data indicate that echinacea preparations do have effects on certain indices of immune function, although at present there is no clear picture as to which specific preparations have the greatest activity. The immunological activity of echinacea has been reviewed,\(^ {37,38}\) and a summary of some of the scientific literature on the immunological effects of echinacea is given below.

Enhancement of macrophage function has been documented for various preparations of echinacea in vitro and in vivo in studies using a range of methods, such as the carbon clearance test and measurement of cytokine production, as indicators of macrophage activity.\(^ {39-41}\) In vitro experiments with human macrophages found that fresh pressed juice and dried juice from the aerial parts of *E. purpurea* stimulated production of cytokines, including interleukin 1 (IL-1), IL-10, and tumour necrosis factor α (TNFα).\(^ {42}\)

Other studies have reported that purified polysaccharides from *E. purpurea*
induced macrophage production of IL-1,\(^{(43)}\) and that a polysaccharide arabinogalactan isolated from plant cell cultures of *E. purpurea* induced TNFα and interferon β2 production by murine macrophages.\(^{(44)}\) Polysaccharides obtained from plant cell cultures of *E. purpurea* have also been shown previously to have immunological activity *in vitro*.\(^{(45)}\) In another series of *in vitro* experiments, *E. purpurea* induced macrophage activation (as assessed by TNFα production) following simulated digestion (incubation of echinacea with gastric fluid) in an attempt to mimic effects following oral administration.\(^{(46)}\) Other work has demonstrated that *E. purpurea* dry root powder (containing 1.5% total polyphenols, calculated as chlorogenic acid) increased the resistance of splenic lymphocytes to apoptosis; splenic lymphocytes were obtained from mice administered the echinacea preparation orally at dosages of 30 or 100 mg/kg daily for 14 days.\(^{(47)}\)

In an *in vitro* study, peripheral blood mononuclear cells (PBMCs) from healthy individuals and from patients with chronic fatigue syndrome and acquired immune deficiency syndrome (AIDS) incubated with increasing concentrations of extracts of *E. purpurea* led to enhanced natural killer function of PBMCs.\(^{(48)}\) *In vivo*, oral administration of *E. purpurea* root extract has been reported to increase numbers of natural killer cells in normal,\(^{(49)}\) leukaemic,\(^{(50)}\) and ageing mice.\(^{(51)}\)

*In vivo* studies in rats have shown that administration of water–ethanol extracts (100 μL twice daily by oral gavage for four days) of *E. purpurea* roots and aerial parts containing defined concentrations of cichoric acid, polysaccharides and alkylamides stimulated phagocytic activity of macrophages; activity was increased with increasing concentrations of the three components.\(^{(52)}\) Subsequently, an increase in lipopolysaccharide-stimulated nitric oxide release was observed by macrophages obtained from the spleens of rats previously treated with the echinacea extracts. A similar set of experiments demonstrated stimulation of alveolar macrophage function by alkylamides administered to healthy rats.\(^{(53)}\)

A proprietary preparation containing *E. purpurea* root extract and liquorice (*Glycyrrhiza glabra*) root extract stimulated phagocytosis *in vitro* and *in vivo*, as demonstrated by the carbon clearance test, following oral administration to mice.\(^{(54)}\) The combination product produced a greater immunostimulatory effect in this test than did either extract tested alone. Another combination preparation, comprising aqueous–ethanol extracts of *E. purpurea* and *E. pallida* root, *Baptisia tinctoria* root and *Thuja occidentalis* herb, administered orally via the diet or drinking water to mice for seven days enhanced the antibody response to sheep red blood cells.\(^{(55)}\)
In contrast with the extensive body of research supporting the immunostimulatory effects of echinacea preparations, some recent work has reported a lack of effect. No evidence of natural killer cell activity or antibody formation was found in studies involving rats fed various preparations of echinacea, including an alcoholic extract of *E. purpurea* root and an alcoholic extract of the roots of *E. angustifolia*, *E. purpurea* and *E. pallida*, in their diet.\(^{[56]}\)

**Antiviral activity**

Antiviral activity has been described for various different preparations of echinacea following *in vitro* studies. An ‘indirect’ antiviral effect was documented in experiments involving addition of glycoprotein-containing fractions obtained from *E. purpurea* root to mouse spleen cell cultures.\(^{[57]}\) Interferon-α and -β produced by the cells were then tested for activity against vesicular stomatitis virus. These glycoprotein-containing fractions were also tested directly against herpes simplex virus (HSV) and were reported to reduce the number of plaques by up to 80%, although raw data were lacking and statistical tests do not appear to have been carried out.

In other *in vitro* studies, the antiviral activity of an aqueous solution of *E. purpurea* herb was tested against aciclovir-susceptible and aciclovir-resistant strains of HSV-1 and HSV-2.\(^{[58]}\) In aciclovir-susceptible strains of HSV-1 and HSV-2, median ED\(_{50}\) (effective dose) values for the echinacea preparation were 1 : 100 (range 1 : 25 to 1 : 400) and 1 : 200 (range 1 : 50 to 1 : 1600), respectively. Similarly, for aciclovir-resistant HSV-1 and HSV-2, median ED\(_{50}\) values (range) were 1 : 100 (1 : 50 to 1 : 400) and 1 : 200 (1 : 50 to 1 : 3200), respectively.

An *n*-hexane extract of *E. purpurea* root, an ethanolic extract of *E. pallida* var. *sanguinea* herb and the isolated constituent cichoric acid were the most potent inhibitors of HSV-1 in *in vitro* studies designed to assess light-activated antiviral activity.\(^{[59]}\) The minimum inhibitory concentrations (MIC) for these preparations were 0.12, 0.026 and 0.045 mg/mL, respectively.

Other *in vitro* studies using mouse fibroblasts found that preincubation with *E. purpurea* herb juice and methanolic and aqueous extracts of *E. purpurea* root resulted in resistance to influenza A2, herpes, and vesicular stomatitis virus infection for 24 hours.\(^{[60]}\)

**Antifungal and antibacterial activities**

Activity against several yeast strains, including *Saccharomyces cerevisiae* and *Candida albicans*, has been described for *n*-hexane extracts of *E. purpurea* roots.\(^{[61]}\) Antifungal activity was observed under near ultraviolet light.
irradiation and, in some cases, was also light independent. The pure polyacetylenic compound trideca-1-ene-3,5,7,9,10-pentayne, isolated from *E. purpurea* root extracts, demonstrated marked light-mediated inhibition of growth of *S. cerevisiae*.\(^{(61)}\) Anti-*Candida* activity for *E. purpurea* extracts has also been described previously.\(^{(37)}\) Other studies in mice have described a dose-dependent protective effect for polysaccharide fractions from *E. purpurea* plant cell cultures against lethal-dose infection with *C. albicans* and *Listeria monocytogenes* when administered intravenously within less than 18 hours of the infection dose.\(^{(62)}\) A similar finding was reported when such polysaccharide fractions were administered to immunosuppressed mice both before and after lethal dose infection with *C. albicans* and *L. monocytogenes*.\(^{(63)}\)

Antibacterial activity against *Escherichia coli*, *Proteus mirabilis*, *Pseudomonas aeruginosa* and *Staphylococcus aureus* has been demonstrated for a multi-herbal preparation containing *E. purpurea* root extract, although it was stated that the observed antibacterial effects were most likely attributable to one of the ingredients, extract of onion.\(^{(64)}\)

**Anti-inflammatory activity**

*In vivo* anti-inflammatory activity has been reported for a polysaccharide fraction (PSF) obtained from *E. angustifolia* roots in the carrageenan-induced rat paw oedema test and in the croton oil mouse ear test, with the PSF administered intravenously and topically, respectively.\(^{(65)}\) The isolated PSF was twice as active as the total aqueous extract in the carrageenan-induced oedema test, and about half as active as indomethacin in the croton oil test. An aqueous extract of *E. angustifolia* roots was also reported to be more effective than benzydamine (a topical non-steroid anti-inflammatory drug (NSAID)) in the croton oil test.\(^{(66)}\) Further work using fractions of an aqueous extract of *E. angustifolia* roots administered topically to mice in the croton oil test attributed the observed anti-inflammatory activity mainly to intermediate and high molecular weight fractions.\(^{(67)}\)

Oral administration of higher (100 mg/kg) but not lower (30 mg/kg) doses of *E. purpurea* dry root powder (containing 1.5% total polyphenols, calculated as chlorogenic acid) inhibited carrageenan-induced paw oedema in mice; the effect was stated to be similar to that of indomethacin 0.25 mg/kg, although this was not tested statistically.\(^{(68)}\) Further exploration suggested that the observed effect may be due to downregulation of cyclooxygenase 2 (COX-2) expression by the echinacea preparation. *In vitro* inhibition of cyclooxygenase 1 (COX-1) and, to a lesser extent, COX-2 has been described for alkamides isolated from *E. purpurea* roots,\(^{(69)}\) and *in vitro* inhibition of 5-lipoxygenase (5-LO) and cyclooxygenase (from sheep seminal microsomes) has been
reported for polyunsaturated alkamides from *E. angustifolia* roots.\(^{(70)}\)

Anti-inflammatory and cicatrising activities have been reported for gel preparations containing echinacoside 0.4 mg and *E. pallida* root extract 100 mg following studies in rats with experimental skin abrasions and excision wounds.\(^{(71)}\) These effects were observed 48 and 72 hours after topical administration, and were stated to be greater than those observed for *E. purpurea* root extract and control. However, no statistical analysis was reported.

The wound-healing properties documented for echinacea have been attributed in part to a polysaccharide fraction, which is thought to inhibit the action of hyaluronidase.\(^{(72)}\) Ethanol extracts of *E. purpurea* roots and aerial parts have been reported to inhibit fibroblast-induced collagen contraction, although the significance of this activity for wound healing needs to be investigated.\(^{(73)}\) Other studies have documented a protective effect for echinacoside, isolated from *E. angustifolia* root, and other caffeoyl esters against free radical-induced degradation of collagen, an experimental model for skin damage caused by exposure to ultraviolet light.\(^{(74)}\)

**Other activities**

A long-chain alkene from *E. angustifolia* is stated to possess antitumour activity *in vivo*, inhibiting the growth of Walker tumours in rats and lymphocytic leukaemia (P388) in mice.\(^{(75)}\)

In an assay of the mosquitocidal activity of alkamides isolated from dried *E. purpurea* roots, a mixture of dodeca-2\(E\),4\(E\),8\(Z\),10\(E\)-tetraenoic acid isobutylamide and dodeca-2\(E\),4\(Z\),8\(Z\),10\(Z\)-tetraenoic acid isobutylamide at a concentration of 100 μg/mL achieved 87.5% mortality of *Aedes aegyptii* L. mosquito larvae within 15 minutes. Several other alkamides assayed also demonstrated mosquitocidal activity, but required longer incubation periods and were less effective.\(^{(69)}\)

Free radical-scavenging activity has been documented for alcoholic extracts of the roots and leaves of *E. purpurea*, *E. angustifolia* and *E. pallida in vitro*.\(^{(10)}\)

Dodeca-2\(E\),4\(E\),8\(Z\),10\(E/Z\)-tetraenoic isobutylamides found in *Echinacea* species (but isolated in this experiment from *Echinacea atrorubens* root) were transported across Caco-2 monolayers, an *in vitro* model for intestinal absorption, over a 6-hour period.\(^{(76)}\) Transport kinetics did not differ significantly following modification of the model (by preincubation of Caco-2 cells with lipopolysaccharide and phorbol 12-myristate-13-acetate) to mimic inflammation.
Clinical studies

Pharmacokinetics
There is a lack of data on the clinical pharmacokinetics of echinacea preparations (see *In vitro* and animal studies, Other activities). One study reported that dodeca-$2E,4E,8Z,10E/Z$-tetraenoic acid isobutylamides (alkamides) were detectable in blood one hour after oral administration of 65 mL of a concentrated ethanolic extract of *E. purpurea* herb (containing 4.3 mg isobutylamides) on an empty stomach to a single healthy volunteer.\(^{(77)}\)

Therapeutic effects
Clinical trials of preparations containing echinacea have focused on testing effects in preventing and treating the common cold and other upper respiratory tract infections (URTIs); some preliminary studies have explored the effects of echinacea in other infections, such as genital herpes, and as an adjunctive treatment in cancer chemotherapy. The rationale for the use of echinacea in these conditions is for its immunomodulatory activity. Collectively, the findings of studies of echinacea are difficult to interpret as studies have assessed preparations containing different species of echinacea and/or different plant parts of echinacea, administered as monopreparations or in combination with other herbal ingredients, and products manufactured by different processes and with different formulations. Hence, the different preparations tested will vary quantitatively and qualitatively in their chemical composition (i.e. will contain different profiles and concentrations of chemical constituents).

**IMMUNOMODULATORY ACTIVITY** One of the first systematic reviews of studies of echinacea-containing preparations assessed evidence of their immunomodulatory effects.\(^{(78)}\) The review included 26 controlled clinical trials, of which six investigated the treatment of URTIs and influenza-like syndromes, seven explored the treatment of other infections, such as sinusitis, bronchitis and candida, six studied the prophylaxis of URTIs and influenza-like syndromes, four tested the reduction by echinacea of adverse effects of antineoplastic treatment and three explored the effects on immunological parameters in patients with infections or malignancies.\(^{(78)}\)

Most studies reported that echinacea-containing preparations were superior to placebo in the indications tested. However, trials included in the review tested different species, parts and preparations (e.g. pressed juice, extract) of echinacea administered via different routes (including oral and parenteral) and with different dosage regimens. In addition, many studies were of poor methodological quality (only eight achieved more than 50% of
the maximum score in an assessment of quality), several preparations tested included other herbs in addition to echinacea, and the review included trials involving patients with a range of conditions, so evidence for the immunomodulatory activity of echinacea from this review can only be considered tentative at best.

The same research group carried out another systematic review of five of their randomised, placebo-controlled studies (four were also conducted double-blind) which investigated the immunomodulatory activity of preparations of echinacea in healthy volunteers.\(^\text{79}\) Again, there were marked differences between the preparations tested in the studies included in the review: combination homeopathic preparations containing \(E.\) \textit{angustifolia} at potencies of D1 and D4 (which can be considered to contain reasonable quantities of starting material) for intravenous administration; ethanolic extracts of \(E.\) \textit{purpurea} root and \(E.\) \textit{pallida} root for oral administration; ethanolic extract of 95\% \(E.\) \textit{purpurea} herb and 5\% \(E.\) \textit{purpurea} root. In two of the five studies, phagocytic activity of polymorphonuclear neutrophil granulocytes (the primary outcome measure) was significantly increased in the echinacea groups, compared with the placebo groups, although no such effects were noted in the other studies.

Recent studies investigating the immunomodulatory activity of echinacea species administered to healthy volunteers have reported different findings. In a randomised, double-blind, placebo-controlled trial, compared with a placebo group, volunteers who received extracts of \(E.\) \textit{purpurea} and \(E.\) \textit{angustifolia} with or without the addition of an arabinogalactan extracted from \(Larix\) \textit{occidentalis} (larch) for four weeks were found to have increased concentrations of complement properdin (thought to be an indication of immune system stimulation).\(^\text{80}\) Other small placebo-controlled studies have reported stimulatory effects following 28 days’ oral pretreatment with pressed juice of \(E.\) \textit{purpurea} on the exercise-induced immune response in athletes,\(^\text{81}\) and of administration of purified polysaccharides from cell cultures of \(E.\) \textit{purpurea} to healthy volunteers.\(^\text{82}\) By contrast, a double-blind, placebo-controlled, crossover study involving 40 healthy volunteers found that oral administration of freshly expressed juice of \(E.\) \textit{purpurea} herb, or placebo, for two weeks did not enhance phagocytic activity of polymorphonuclear leukocytes or monocytes, or affect TNF\(\alpha\) and IL-1 production.\(^\text{83}\)

Preliminary studies have assessed the effects of a combination preparation containing extracts of \(E.\) \textit{angustifolia}, \textit{Eupatorium perfoliatum} (boneset) and \textit{Thuja occidentalis} (thuja) on cytokine production in patients who have undergone curative surgery for various solid malignant tumours,\(^\text{84}\) and
the immunostimulatory effects of a regimen comprising intramuscular *E. purpurea* extract, low-dose intramuscular cyclophosphamide and intravenous thymostimulin in patients with advanced colorectal cancer.\(^{(85)}\) In another study, the effects of a polysaccharide fraction of *E. purpurea* herb obtained from cell cultures in reducing the adverse effects of cancer chemotherapy were explored in patients with advanced gastric cancer receiving palliative therapy with etoposide, leucovorin and 5-fluorouracil.\(^{(86)}\) Although these studies reported some positive findings with echinacea, no firm conclusions can be drawn because of the nature of the study designs, therefore further research in this area is required.

**Upper Respiratory Tract Infections** (Numerous studies have explored the effects of echinacea preparations in preventing or treating the common cold and other URTIs. Overall, several, but not all, studies have reported beneficial effects for certain echinacea preparations, compared with placebo, for the prevention and treatment of URTIs. However, for the reasons given (see Therapeutic effects), current consensus is that there is insufficient evidence to recommend any specific echinacea preparations, or to advise on optimal dose and treatment duration.

**Prophylaxis** A Cochrane systematic review included 16 randomised and quasi-randomised controlled trials – involving a total of almost 3400 participants – of extracts of echinacea for preventing \((n = 8)\) or treating \((n = 8)\) URTIs.\(^{(87)}\) The eight ‘prevention’ trials comprised five which were placebo-controlled \((n = 1272\) participants), and which largely were considered to be of adequate methodological quality, and three \((n = 1139\) participants) in which the control group received no treatment. The five placebo-controlled trials tested combination echinacea preparations \((n = 2)\) or monopreparations of *E. purpurea* herb or root, or *E. angustifolia* root \((n = 3)\), administered orally typically for 8–12 weeks. Two of these studies reported a statistically significant reduction in the incidence of URTIs in echinacea recipients, compared with placebo recipients (odds ratios, 95% confidence interval \((CI)\): 0.45, 0.22–0.92 and 0.27, 0.11–0.66). One of these studies also found that in participants who did acquire infections, the duration was significantly shorter in those who had received echinacea compared with placebo recipients, although two other studies reported no difference in this outcome.

The three other ‘prevention’ trials all involved children and compared a combination preparation containing extracts of *E. angustifolia* and *E. pallida* root, *Baptisia tinctoria* root and *Thuja occidentalis* herb, as well as several homeopathic dilutions, with no treatment. All three studies reported that the frequency of infection was significantly lower in the treatment
compared with no treatment group (pooled odds ratio 0.36; 95% CI 0.28–0.46), although the methodological quality of all three studies was considered inadequate.\(^{(87)}\)

Several new trials of echinacea preparations in the prevention of the common cold have been completed since the Cochrane review, but have not shown beneficial effects for echinacea preparations, compared with placebo, on main outcome measures.\(^{(88,89)}\)

A randomised, double-blind, placebo-controlled trial involved 302 healthy volunteers recruited from military institutions and an industrial plant who received an ethanolic extract of \(E.\ purpurea\) root or \(E.\ angustifolia\) root (drug : extract ratio, 1 : 11 in 30% alcohol), or placebo, 50 drops twice daily on five days per week (Monday to Friday) for 12 weeks.\(^{(88)}\) In an intention-to-treat analysis \((n = 289)\), the proportion of participants who experienced at least one URTI was 32% (95% CI: 23–41%) for \(E.\ angustifolia\) recipients, 29% (95% CI: 20–38%) for \(E.\ purpurea\) recipients, and 37% (95% CI: 27–47%) for placebo recipients; these differences were not statistically significant \((p = 0.55)\). Similarly, there were no statistically significant differences between groups in time to occurrence of the first URTI \((p = 0.49)\), or in the duration of infections \((p = 0.29)\), although it is possible that the study was not large enough to detect differences. However, a greater proportion of echinacea recipients believed they had benefited from the study medication than did placebo recipients (78, 70 and 56% for \(E.\ angustifolia\), \(E.\ purpurea\) and placebo, respectively; \(p = 0.04)\).\(^{(88)}\)

In another randomised, double-blind, placebo-controlled trial, involving 109 individuals who had experienced more than three colds or respiratory infections in the previous year, a fluid extract of \(E.\ purpurea\) prepared from the aerial parts of fresh flowering plants, administered at a dose of 4 mL twice daily for eight weeks, had no statistically significant effect compared with placebo on the incidence of colds and URTIs (rate ratio for number of participants in each group with at least one cold or URTI = 0.88, 95% CI: 0.60–1.22).\(^{(89)}\) Likewise, there was no statistically significant difference between groups in the duration and severity of occurring colds or URTIs.

A further study tested the effects of echinacea for the prevention of colds due to experimental rhinovirus infection.\(^{(90)}\) Adult volunteers \((n = 117\) enrolled) with a serum titre of neutralising antibody to rhinovirus of \(\leq 1 : 4\) received echinacea (300 mg) or placebo three times daily for 14 days prior to and for five days after challenge with rhinovirus \((n = 92\) challenged due to study withdrawals). It is not stated in a report of the study\(^{(90)}\) whether
random allocation to study group was undertaken, or whether participants were masked (blind) to treatment allocation, although a blinding check before virus challenge found that 30 (60%) of the 50 echinacea recipients and 19 (45%) of the 42 placebo recipients thought they were receiving the ‘active’ treatment \( (p = 0.21) \).

The study did not provide evidence to suggest that echinacea had effects over those of placebo: rhinovirus infection occurred in 22 (44%) of echinacea recipients and in 24 (57%) of placebo recipients (rate ratio = 0.77; \( p = 0.3 \)), ‘clinical’ colds developed in 50 and 59% of echinacea and placebo recipients, respectively \( (p = 0.77) \), and there was no difference in mean total symptom scores (11.4, 95% CI 3.9–18.9 and 13.6, 95% CI 7.5–19.7 for echinacea and placebo, respectively). However, the study involved small numbers of participants and a sample size calculation was not reported, hence it is possible that the study was not large enough to be able to detect a difference if one existed. Additionally, information on the species of echinacea, plant part used, type of preparation (e.g. extract) and route of administration used was not provided in a report of this study.\(^{(90)}\) It was stated that the preparation contained cichoric acid 0.16% and almost no echinacosides or alkamides, but with this limited information, it is not possible to say with certainty which species is likely to have been used, although it may have been \textit{E. purpurea}.  

\textit{Treatment} The Cochrane systematic review described above (see Prophylaxis) included eight randomised, placebo-controlled trials of echinacea preparations for the treatment of URTIs.\(^{(87)}\) These trials tested three different combinations of echinacea extracts and two monopreparations, taken orally typically for 6–10 days. Six studies reported statistically significant beneficial effects for echinacea recipients, compared with placebo recipients, on outcome measures such as duration of illness or symptoms (e.g. ‘running nose’). However, heterogeneity of the studies precluded any further summary of the results. In addition, several of the studies had methodological flaws or their methodological quality could not be determined due to a lack of detail about the study designs in published reports. For these reasons, although the majority of the studies described reported positive results for echinacea preparations, it was not possible to recommend any specific product for the treatment of the common cold and further research is necessary.\(^{(87)}\)

Several new trials of echinacea preparations in the treatment of URTIs have been completed since the Cochrane review (Table 1) and have reported conflicting results.\(^{(91-94)}\)
The most recent study, a randomised, double-blind, placebo-controlled, community-based trial involving 148 students with common colds of recent onset, assessed the effects of capsules containing unrefined *Echinacea purpurea* herb (62 mg), root (62 mg) and *E. angustifolia* root (123 mg). Analysis of samples of the preparation by independent laboratories found that they contained cichoric acid and alkamides (0.5 to <1.0%), and echinoside, chlorogenic acid and caffeoyltartaric acid (all >0.1% to <0.5%). Participants took four capsules six times during the first 24 hours of the onset of a cold, followed by four capsules three times daily until symptoms resolved, or for up to 10 days. Among the 142 participants who completed the study, there was no difference in the mean duration of cold symptoms (6.27 and 5.75 days for the echinacea and placebo groups, respectively; difference: −0.52 days, 95% CI −1.09 to 0.22 days), even though the study had an adequate sample size: with a sample size of 150 participants, the study would have had 80% power to detect a benefit of two days’ duration.

Three different preparations and doses of *E. purpurea* were tested in a randomised, double-blind, placebo-controlled trial in healthy adults. The four arms of the study were: 6.78 mg *E. purpurea* crude extract, based on 95% herb and 5% root (Echinaforce); 48.27 mg *E. purpurea* crude extract, based on 95% herb and 5% root; 29.60 mg *E. purpurea* crude extract, based on root only; and placebo. In total, 246 participants experienced symptoms typical of the onset of a common cold and took their allocated study medication two tablets three times daily until they felt better, or for up to seven days. According to an intention-to-treat analysis, the two echinacea extracts prepared from both *E. purpurea* herb and root were significantly more effective than *E. purpurea* root extract and placebo in reducing symptoms as assessed by the investigator (the primary outcome measure): the relative reductions in the mean complaint index for these preparations were 58.7% (95% CI 48.7–68.7; \( p = 0.045 \) versus placebo) and 58.1% (95% CI 47.7–69.7; \( p = 0.027 \) versus placebo).

Statistically significant effects for an extract of *E. purpurea* herb (Echinacin) on median duration of illness were reported in another randomised, double-blind, placebo-controlled trial involving 80 adults who experienced onset of a cold (median duration six and nine days for echinacea and placebo, respectively; \( p = 0.0112 \)). Participants started taking their allocated
medication on first experiencing symptoms, and continued treatment (5 mL twice daily) until symptoms resolved.

A further placebo-controlled study involving adults with early symptoms of a cold (n = 95) explored the effects of a combination preparation containing *E. purpurea* and *E. angustifolia* herb and extract of *E. purpurea* root, as well as lemongrass leaf and spearmint leaf as flavourings, formulated as a tea.\(^{94}\) When prepared as directed, tea prepared from one bag was stated to provide 31.5 mg of phenolic compounds, calculated as caftaric acid, cichoric acid, chlorogenic acid and echinacoside. The results suggested a statistically significant difference between the treatment and placebo groups in self-rated effectiveness – mean (standard deviation, SD) effectiveness score 4.13 (0.96) and 2.78 (0.95) for echinacea tea and placebo, respectively (\(p < 0.001\)) – although the mean (SD) duration of symptoms was significantly longer in the echinacea group, compared with the placebo group (4.33 (0.93) and 2.34 (1.09) for echinacea tea and placebo, respectively; \(p < 0.001\)). In addition, the study had several methodological limitations. For example, although stated to be randomised, the study did not involve true randomisation (participants were allocated to groups alternately), the ‘placebo’ tea contained low doses of several herbs (peppermint leaf, sweet fennel seed, ginger rhizome, papaya leaf, alfalfa leaf and cinnamon bark), and outcomes were self-assessed only.

Several trials of echinacea preparations in the prevention of URTIs provide data on duration and severity of infections occurring in participants (see Prophylaxis). While these data have some relevance to treatment, they should not be grouped together, since the dosage regimens are entirely different – in ‘prevention’ trials, participants may have received study medication for several weeks or more before experiencing an infection, whereas in ‘treatment’ trials, participants usually start study medication immediately after the onset of symptoms.

**OTHER INFECTIONS** A randomised, double-blind, placebo-controlled, crossover trial assessed the effects of an extract of *E. purpurea* herb (95%) and root (5%) (Echinaforce) on the incidence and severity of recurrent genital herpes in 50 patients who had not been exposed to aciclovir or similar medicines within 14 days of enrollment into the study and who had had at least four recurrences of genital herpes within the previous 12 months.\(^{95}\) Study medication, or placebo, was taken orally 800 mg twice daily for six months. The study did not show any significant difference between the two groups on the outcomes measured (frequency and duration of recurrences, pain score, CD4 cell count, neutrophil count), although there was a high dropout rate during the study.
A systematic review of studies exploring the immunomodulatory effects of echinacea-containing preparations included seven controlled clinical trials in infections such as sinusitis, bronchitis and candida (see Immunomodulatory activity). (78)
Side-effects, Toxicity

Data on numbers of participants experiencing adverse events were provided by several studies included in a Cochrane systematic review of 16 randomised and quasi-randomised controlled trials of extracts of echinacea for preventing or treating URTIs (see Clinical studies, Prophylaxis and Treatment). Four placebo-controlled ‘prevention’ trials of echinacea reported these data: in three trials, involving a total of around 1000 participants, the frequency of adverse events in the echinacea group was similar to that in the corresponding placebo group; in one trial, adverse events did not occur in either the echinacea or placebo groups. Three ‘treatment’ trials provided adverse event data: in two studies, adverse events were not observed in either the echinacea or the placebo groups, and in one, numbers of patients experiencing adverse events in the echinacea and placebo groups were similar (four and five, respectively).

New clinical trials published since the Cochrane review also report that there was no statistically significant difference in the frequency of adverse events noted for echinacea and placebo. Where adverse events were reported, most commonly these were mild gastrointestinal symptoms. Another review of clinical data, mostly from clinical trials, concluded that oral administration of the expressed juice of *E. purpurea* herb is well-tolerated. The review included data from an unpublished post-marketing surveillance study involving over 1200 individuals aged 2 to 20 years who used oral *E. purpurea* lozenges for 4–6 weeks for URTIs, and which indicated that unpleasant taste was the most frequently reported adverse event.

On the basis of these limited data, it seems that the risk of acute adverse effects with echinacea is very small. However, it is not possible to draw firm conclusions from these data for several reasons – different echinacea preparations and regimens were tested, different patient populations (adults, children) were involved, and echinacea preparations were administered for only a short time period, particularly in the ‘treatment’ trials. In addition, since clinical trials usually have the statistical power only to detect common, acute adverse effects and, as there is a lack of data on the safety of the longer term use of echinacea preparations, there is a need for further evaluation of the safety of different echinacea preparations.

The low number of reports of suspected adverse reactions associated with echinacea preparations set against estimates of the high frequency of use of echinacea has been used as an argument for the safety of echinacea. However, this argument is flawed since it fails to consider that under-reporting of suspected adverse reactions associated with herbal medicines is
likely at several levels,\(^{(98, G21)}\) and that in general reporting systems for herbal medicines are not well-established. The use of sales data to estimate the frequency of an adverse reaction can be misleading at best and, in addition, the argument takes no account of the differences in preparations of echinacea. The UK Committee on Safety of Medicines and Medicines and Healthcare Products Regulatory Agency yellow card scheme for adverse drug reaction reporting received 27 reports of suspected adverse drug reactions (ADRs) associated with echinacea preparations for the period 1996 to end of June 2003.\(^{(99)}\)

**Allergic reactions**

*Echinacea* species belong to the Asteraceae (Compositae, daisy) family, members of which are known to cause allergic reactions. Individuals with allergic tendencies, particularly those with known allergy to other members of the Asteraceae family (e.g. chamomile) should be advised to avoid echinacea preparations containing aerial parts.\(^{(G50)}\)

Isolated spontaneous reports of suspected ADRs associated with the use of echinacea preparations include allergic skin reactions.\(^{(96)}\) In Australia, detailed assessment of five cases of allergic reactions temporally associated with echinacea (anaphylaxis, 2; acute asthma attack in an echinacea-naive individual, 1; recurrent mild asthma, 1; macropapular rash, 1), three of which reported positive rechallenge, revealed that three patients had positive skin-prick test results for echinacea.\(^{(100)}\) One case report described a 37-year-old woman with atopy who experienced anaphylaxis 30 minutes after ingesting several dietary supplements (vitamins B\(_12\) and E, an iron preparation, ‘folate’, vitamin B complex, a multivitamin preparation, zinc, antioxidants, a garlic and onion preparation, evening primrose oil) and 15 minutes after taking 5 mL of an echinacea preparation, stated to be equivalent to *E. angustifolia* whole plant extract 3825 mg and *E. purpurea* dried root 150 mg.\(^{(101)}\) The woman took promethazine and was observed in an emergency department for 2 hours, and her symptoms resolved without further treatment. Two weeks later she gave a positive skin-prick test to the echinacea product, but not to ‘crude’ extracts of the other supplements she had taken. She had been taking echinacea for 2–3 years and had previously taken the same product without experiencing any adverse effects. A causal association in this case has been questioned.\(^{(102)}\)

Positive skin-prick test results for echinacea were also reported for 20% of 100 echinacea-naive atopic individuals, and over 50% of 26 Australian suspected ADR reports of hypersensitivity associated with echinacea involved individuals with atopy.\(^{(100)}\) Echinacea has previously been reported to have
produced positive patch test reactions in four individuals with a previous history of plant dermatitis.\textsuperscript{(G51)} These reports raise hypotheses that require testing in formal studies.

An isolated report describes a 41-year-old man who experienced four episodes of erythema nodosum after using an echinacea preparation at each onset of an influenza-like illness.\textsuperscript{(103)} The man had been using echinacea intermittently for 18 months, as well as loratadine on an as required basis and St. John’s wort for the previous six months. Each episode of erythema nodosum responded to conventional treatment including prednisone. The man was advised to discontinue treatment with echinacea and, after one year, had not experienced any further recurrences. However, the report does not provide any details (species, plant part, formulation, dosage regimen) of the echinacea preparation involved and therefore is difficult to interpret. Causality has not been established.

Other reactions

A case of hypokalaemic renal tubular acidosis due to Sjögren’s syndrome (a symptom complex of unknown aetiology, marked by keratoconjunctivitis sicca, xerostamia, with or without lachrymal and salivary gland enlargement, respectively, and presence of connective tissue disease, usually rheumatoid arthritis, but sometime systemic lupus erythematosus, scleroderma or polymyositis) has been reported in a 36-year-old woman.\textsuperscript{(104)} She was stated to have begun taking echinacea, St. John’s wort and kava two weeks before becoming ill, but the report does not provide any further details of the echinacea species contained in the product(s), nor of the types of preparations, formulations, dosages and routes of administration of any of the herbal medicines listed. The woman was hospitalised with severe generalised muscle weakness and tests revealed she had a serum potassium ion concentration of 1.3 mEq/L. She was given electrolyte replacement for four days, after which the muscle weakness resolved, and was started on hydroxychloroquine 200 mg daily for ‘probable’ Sjögren’s syndrome. The authors suggested that ingestion of echinacea may have aggravated an autoimmune disorder, although rechallenge with echinacea was not undertaken, and causality has not been established.\textsuperscript{(104)}

Another report describes a 49-year-old woman who presented with a five-day history of numbness and weakness in her right arm.\textsuperscript{(105)} For the previous seven weeks she had received echinacea comp 2 mL mixed with 5 mL of her venous blood intramuscularly twice weekly to prevent infections and ‘boost’ her immune system. The injection was stated to contain \textit{E. angustifolia} D2 1.1 mL, Aconitum D4 0.3 mL and Lachesis (bushmaster snake venom) D8
0.3 mL. (D nomenclature relates to a homeopathic dilution step; D2 is equivalent to a 1 in 100 dilution, whereas D8 is equivalent to a 1 in 100 000 000 dilution).\(^{(106)}\) The woman was admitted to hospital with mild spastic paresis and fluctuating numbness of the right arm and was described as having acute disseminated encephalomyelitis; symptoms resolved after treatment with methylprednisolone 500 mg daily by intravenous infusion. Causality has not been established.

**Toxicology**

In general, animal studies with different preparations and fractions of echinacea species have indicated low toxicity.\(^{(37)}\) In acute toxicity studies involving polysaccharide fractions from *E. purpurea* administered by intraperitoneal injection to small numbers of mice, the LD\(_{50}\) (lethal dose) for female mice was 2500 mg/kg.\(^{(107)}\) Other acute toxicity studies using a preparation comprising pressed juice from *E. purpurea* herb have provided LD\(_{50}\) values in mice of >30 000 mg/kg and >10 000 mg/kg for oral and intravenous administration, respectively, and in rats of >15 000 mg/kg and >5000 mg/kg for oral and intravenous administration, respectively.\(^{(108,109)}\) Further experiments showed no evidence of mutagenic activity in bacteria and mammalian cells *in vitro* and *in vivo* in mice.\(^{(109)}\)

High concentrations of *E. purpurea* (8 mg/mL) have been reported to reduce sperm motility, sperm penetration of hamster oocytes and to be associated with sperm DNA denaturation *in vitro*; no such effects were observed with low concentrations.\(^{(110,111)}\) These findings are difficult to interpret since there is a lack of detail regarding the preparation of *E. purpurea*, and their clinical relevance is questionable.

The pyrrolizidine alkaloids isotussilagine and tussilagine have been documented for echinacea, although they possess a saturated pyrrolizidine nucleus and are not thought to be toxic.

*In vivo* antitumour activity and *in vitro* stimulation of TNF\(\alpha\) secretion have been reported for echinacea. In addition to its antitumour effects, TNF is stated to be a mediator of cachexia and the manifestations of endotoxic shock. Concern has been expressed over the possible toxicity of TNF.\(^{(G67)}\)
Contraindications, Warnings

It has been stated that echinacea is contraindicated in patients with progressive systemic diseases, such as tuberculosis, leukaemia and leukaemia-like diseases, collagen disorders, multiple sclerosis and other autoimmune diseases.\(^{(G56)}\)

In the UK, some products also advise against use in AIDS and HIV infections. The basis for these statements appears to be a theoretical one, based on evidence that echinacea preparations have immunomodulatory activity; there is an opposing view that echinacea is not harmful in autoimmune diseases.\(^{(G50)}\)

At present, there is a lack of reliable clinical evidence to support these views, although in view of the seriousness of the conditions listed, it is appropriate to avoid use in these disorders until further information is available.

Interactions

There are no reported drug interactions for echinacea, although on the basis of its documented immunomodulatory activity, as a general precaution, echinacea should only be used with caution in patients taking immunosuppressant drugs.

The effects of echinacea products available in Canada on inhibition of the human cytochrome P450 drug metabolising enzyme CYP3A4 have been tested in vitro using a fluorometric mitrotitre plate assay.\(^{(112)}\)

In the study, 10 mL samples of preparations of *E. angustifolia* roots, *E. purpurea* roots and herb, and a 1 : 1 blend of *E. angustifolia* and *E. purpurea* (plant parts not specified) were standardised to contain ethanol 55% and used as stock solutions. Samples of serial dilutions of these preparations, as well as different concentrations of the pure compounds echinacoside and cichoric acid, were assayed. The blend of *E. angustifolia* and *E. purpurea*, and *E. purpurea* herb showed ‘moderate’ inhibition of CYP3A4: median (95% CI) inhibitory concentration (IC\(_{50}\)) values (% of full strength preparation) were 6.73 (4.75, 10.09) and 8.56 (5.95, 13.05), respectively. Echinacoside also showed moderate inhibitory activity (median IC\(_{50}\) values (95% CI) 6.29 (2.07, 71.56)), whereas cichoric acid showed low inhibitory activity.\(^{(112)}\)

A study in mice fed both melatonin and an extract of *E. purpurea* root in their diet reported reduced numbers of proliferating myeloid cells in the spleen and bone marrow.\(^{(113)}\)

Further research is needed to determine whether these findings have clinical importance.

Pregnancy and lactation
There is a lack of data on the safety of echinacea preparations taken during pregnancy and lactation and, given that the benefits of specific echinacea preparations have not been established definitively, excessive use during these periods should be avoided as a general precaution.

A cohort study compared numbers of live births, and spontaneous and therapeutic abortions occurring among women who had taken echinacea preparations during pregnancy ($n = 206$, 112 of whom took echinacea during the first trimester) with those occurring among a control group of 206 women matched for disease (URTI), maternal age and alcohol and cigarette use.$^{114}$ The exposed group of women had telephoned a hospital teratogen information service regarding the use of echinacea during pregnancy; the unexposed group had also telephoned the service for this reason, but subsequently did not use echinacea or used a non-teratogenic antibiotic instead.

There were no statistically significant differences between the two groups in assessed outcomes including number of live births, spontaneous and therapeutic abortions, gestational age, birth weight and rates of malformations. In the exposed group there were six major and six minor malformations, compared with seven major and seven minor malformations in the control group.$^{114}$ The study has several limitations, particularly the small sample size, meaning that the study would have the statistical power only to detect common malformations, and self-report of exposure, since it is possible that misclassification could have occurred (e.g. exposed women reported as unexposed). In addition, participants used a range of different preparations of echinacea at different dosage regimens, so the study does not provide adequate evidence for any specific preparation. Further study is required to establish the safety profile of echinacea during pregnancy.
The chemistry of echinacea is well documented (see Constituents). The three species are chemically dissimilar. *E. purpurea* and *E. angustifolia* both contain amides as their major lipophilic constituents, but of differing structural types. By contrast, the lipophilic fraction of *E. pallida* is characterised by polyacetylenes and contains only very low concentrations, if any, of amides. The polyene constituents are stated to be susceptible to auto-oxidation, resulting in the formation of artefacts during storage.\(^{(G2)}\)

Commercial echinacea samples and marketed echinacea products may contain one or more of the three echinacea species mentioned above. Analysis of commercial samples of raw echinacea material and marketed echinacea products has shown that in some cases the echinacea species assigned to the sample or product was incorrect, and that the pharmaceutical quality and labelling of some finished products was inadequate (see Quality of plant material and commercial products). Users and potential users of echinacea products should be made aware of the possible differences between products and the implications of this for efficacy and safety.

Documented scientific evidence from *in vitro* and animal studies supports some of the uses for echinacea, particularly the reputed immunostimulant properties,\(^{(37)}\) although immunostimulant activity has been disputed following one series of studies (see *In vitro* and animal studies, Immunomodulatory activity).\(^{(56)}\) Reported pharmacological activities have been documented for the polyene and high molecular weight polysaccharide constituents, as well as the alkylamides and caffeic acid derivatives.

Several, but not all, clinical trials of echinacea preparations have reported effects superior to those of placebo in the prevention and treatment of URTIs. However, evidence of efficacy is not definitive as studies have included different patient groups and tested various different preparations and dosage regimens of echinacea.\(^{(87,97)}\) As such, there is insufficient evidence to recommend any specific echinacea products, or to advise on optimal dose and treatment duration (see Clinical Studies). Further well-designed clinical trials using well-defined, standardised preparations are necessary in order to establish efficacy.

There is a lack of clinical research on the anti-inflammatory and wound-healing properties of echinacea preparations documented *in vitro* and in animal studies. Several other areas of interest, related to the immunostimulant effects of echinacea, such as prevention of recurrence of genital herpes and other infections, and reduction of adverse effects...
associated with antineoplastic treatment, also require further clinical investigation.

Another area that requires further study is whether certain groups of constituents, such as the polysaccharides, are active after oral administration and, if so, what is the mechanism of action since polysaccharides usually would be broken down into simple inactive sugars.\(^{(37)}\) There is a lack of data on the pharmacokinetics of echinacea preparations, although very preliminary studies have reported transportation of isobutylamides across Caco-2 cells, an \textit{in vitro} model of intestinal absorption,\(^{(76)}\) and detection of these compounds in blood obtained from a single healthy volunteer.\(^{(77)}\)

On the basis of the available (limited) safety data, which come mostly from short-term clinical trials of echinacea preparations for the prevention and treatment of URTIs in otherwise generally healthy individuals, echinacea appears to be well-tolerated. However, firm conclusions cannot be drawn from these limited data, and further investigation is required to establish the safety profile of different echinacea preparations. At present, the main safety issues are the possibility of allergic reactions, and concern about the use of echinacea by patients with progressive systemic diseases, such as tuberculosis, leukaemia, collagen disorders, multiple sclerosis and other autoimmune diseases (see Side-effects, Toxicity and Contra-indications, Warnings). In view of the lack of toxicity data, excessive use of echinacea should be avoided. In placebo-controlled trials of echinacea preparations for the prophylaxis of URTIs, treatment was taken typically for 8–12 weeks.
References

See also General References G2, G3, G4, G5, G6, G7, G9, G18, G32, G34, G36, G37, G50, G52, G54, G56, G63, G66, G67 and G69.

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Species (Family)

*Sambucus nigra* L. (Caprifoliaceae)
Synonym(s)

Black Elder, European Elder, Sambucus

*Sambucus canadensis* L. refers to American Elder
Part(s) Used

Flower
Pharmacopoeial and Other Monographs

BHC 1992\textsuperscript{(G6)}
BHP 1996\textsuperscript{(G9)}
BP 2002\textsuperscript{(G71)}

Complete German Commission E\textsuperscript{(G3)}

Martindale 33rd edition\textsuperscript{(G67)}

PDR for Herbal Medicines 2nd edition\textsuperscript{(G36)}

Ph Eur 2004\textsuperscript{(G72)}
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents

See General References G2 G6 G41 G62 G64.

Flavonoids
Flavonols (kaempferol, quercetin), quercetin glycosides (1.5–3.0%) including hyperoside, isoquercitrin and rutin.

Triterpenes
α- and β-amyrin, oleanolic and ursolic acids.

Volatile oils
0.3%. 66% fatty acids (primarily linoleic, linolenic and palmitic) and 7% alkanes (C$_{19}$, C$_{21}$, C$_{23}$ and C$_{25}$). Numerous other constituent types have been identified including ethers and oxides, ketones, aldehydes, alcohols and esters.$^{(1)}$

Other constituents
Chlorogenic acid, tannin, mucilage, plastocynin (protein)$^{(2)}$, pectin and sugar.

Other plant parts
Leaf Sambunigrin (0.042%), prunasin, zierin and holocalin (cyanogenetic glycosides)$^{(3)}$, choline, flavonoids (rutin, quercetin), sterols (sitosterol, stigmasterol, campesterol), triterpenes (α- and β-amyrin palmitates, oleanolic and ursolic acids), alkanes, fatty acids, tannins and others.$^{(G41)}$

Bark Lectin (mol. wt 140 000) rich in asparagine/aspartic acid, glutamine/glutamic acid, valine and leucine$^{(4)}$, phytohaemagglutinin$^{(5)}$, triterpenoids (α-amyrenone, α-amyrin, betulin, oleanolic acid, β-sitosterol).$^{(6)}$
Food Use

Elder is listed by the Council of Europe as a source of natural food flavouring (categories N1 and N2). Category N1 refers to the fruit and indicates that there are no restrictions on quantities used. Category N2 refers to the restrictions on the concentrations of hydrocyanic acid that are permitted, namely 1 mg/kg in beverages and foods, 1 mg/kg for every per cent proof of alcoholic beverages, 5 mg/kg in stone fruit juices, 25 mg/kg in confectionery and 50 mg/kg in marzipan.\(^{(G16)}\) In the USA, the flowers have a regulatory status of GRAS (Generally Recognised As Safe).\(^{(G41)}\)
Elder is stated to possess diaphoretic and anticatarrhal properties. Traditionally, it has been used for influenza colds, chronic nasal catarrh with deafness and sinusitis.\(^{(G8)}\) Elder is also stated to act as a diuretic, laxative and local anti-inflammatory agent.\(^{(G2\ G6\ G7\ G8\ G41\ G49\ G64)}\)
Dosage

Dried flower
2–4 g by infusion three times daily.\(^{(G6 G7)}\)

Liquid extract
2–4 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G6 G7)}\)
Pharmacological Actions

In vitro and animal studies

Elder is stated to possess diuretic and laxative properties. (G41)

Moderate (27%) anti-inflammatory action in carrageenan-induced rat paw oedema has been documented for an elder preparation given one hour before carrageenan (100 mg/kg, by mouth). (7) Indomethacin as a control exhibited 45% inhibition at a dose of 5 mg/kg. (7)

An infusion made from the flowers of elder, St. John’s wort herb and root of soapwort (Saponaria officinalis) has exhibited antiviral activity against influenza types A and B (in vivo and in vitro) and herpes simplex virus type 1 (in vitro). (8)

A diuretic effect in rats exceeding that exerted by theophylline has been reported for elder. (9) An infusion and extracts rich in potassium and in flavonoids all caused diuresis. Greatest activity was exerted by the combined potassium- and flavonoid–rich extracts.

In vitro antispasmodic activity (rat ileum, rabbit/guinea–pig intestine) and spasmogenic activity (rat uterus) have been reported for lectins isolated from elder. (10)

A lectin isolated from elder bark was found to be a lactose–specific haemagglutinin with a slightly higher affinity for erythrocytes from blood group A. (4) Unlike many other plant lectins, the lectin did not inhibit protein synthesis. (4) The carbohydrate–binding properties of a lectin isolated from elder bark have been studied. (11)

Phytohaemagglutinins are biologically active extracts isolated from various plants and represent a class of lectin. They are associated with haemagglutination and mitogenic, antigenic and immunosuppressant properties. (5) In vitro, phytohaemagglutinin has been found to stimulate production of an interferon–like substance in human leukocytes. (G45)

Hepatoprotective activity against carbon tetrachloride–induced toxicity has been reported for triterpenes isolated from Sambucus formosana Nakai. (12)

Clinical studies

None documented for elder. Phytohaemagglutinin extracts have been used clinically to treat drug–induced leucopenia and some types of anaemia. (5)
blastogenic response of lymphocytes to phytohaemagglutinin has been used extensively as a measure of immunocompetence.\(^{(G45)}\)
Side–effects, Toxicity

No reported side–effects specifically for elder were located. Human poisoning has occurred with *Sambucus* species.\(^{(13)}\) The roots, stems and leaves and, much less so, the flowers and unripe berries, are stated to contain a poisonous alkaloid and cyanogenic glycoside causing nausea, vomiting and diarrhoea.\(^{(13)}\) The flowers and ripe fruit are stated to be edible without harm.\(^{(13)}\)

The effects of a lectin isolated from elder bark on mammalian embryonic and fetal development has been studied.\(^{(5)}\) The lectin exerted mainly a toxic effect and, to a lesser degree, a teratogenic effect when administered subcutaneously to pregnant mice. In view of the high doses administered, the authors stated that the results did not indicate a potential hazard to human fetuses exposed to lectins.\(^{(5)}\)
Contra–indications, Warnings

Excessive or prolonged use may result in hypokal aemia in view of the documented diuretic effect. Plant parts other than the flowers are reported to be poisonous and should not be ingested.

Pregnancy and lactation
The safety of elder taken during pregnancy has not been established. In view of the lack of toxicity data, the use of elder during pregnancy and lactation should be avoided.
Pharmaceutical Comment

Phytochemical details have been documented for elder, with flavonoids and triterpenes representing the main biologically active constituents. Anti-inflammatory, antiviral and diuretic effects have been observed in \textit{in vivo} studies, thus supporting the herbal uses of elder. No documented studies in humans were found. Potentially toxic compounds have been reported for the bark (lectins) and the leaves (cyanogenetic glycosides); the flowers are suitable for use as a herbal remedy.
References


5. Paulo E. Effect of phytohaemagglutinin (PHA) from the bark of Sambucus nigra on embryonic and foetal development in mice. Folia Biol (Kraków) 1976; 24: 213–222.

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Species (Family)

*Inula helenium* L. (*Asteraceae/Compositae*)
An elecampane extract has been referred to as helenin. Alantolactone is also known as elecampane camphor, alant camphor, helenin and inula camphor.
Part(s) Used

Rhizome, root
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL$^{(G37)}$
Constituents

See General References G2 G6 G41 G64.

**Carbohydrates**
Inulin (up to 44%), mucilage.

**Terpenoids**
β- and γ-sitosterols, stigmasterol and damaradienol (sterols), friedelin.

**Volatile oils**
1–4%. Mainly contains sesquiterpene lactones including alantolactone, isoalantolactone and dihydroalantolactone (eudesmanolides), alantic acid and azulene.

**Other constituents**
Resin.
Food Use

Elecampane is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that elecampane can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\textsuperscript{(G16)}

In the USA, elecampane is only approved for use in alcoholic beverages.\textsuperscript{(G41)}
Herbal Use

Elecampane is stated to possess expectorant, antitussive, diaphoretic and bactericidal properties. Traditionally, it has been used for bronchial/tracheal catarrh, cough associated with pulmonary tuberculosis and dry irritating cough in children.\(^{(G2\ G6\ G7\ G8\ G64)}\)

Alantolactone has been used as an anthelmintic in the treatment of roundworm, threadworm, hookworm and whipworm infection.\(^{(G44\ G45)}\)
Dosage

**Rhizome/root**
1.5–4.0 g or by decoction three times daily.\(^{(G6 \ G7)}\)

**Liquid extract**
1.5–4.0 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G6 \ G7)}\)

**Alantolactone**
300 mg daily for two courses of 5 days, with an interval of 10 days. Children, 50–200 mg daily.\(^{(G44)}\)
**Pharmacological Actions**

**In vitro and animal studies**

Elecampane infusion has exhibited a pronounced sedative effect in mice.\(^{\text{G41}}\) Alantolactone has been reported to exhibit hypotensive, hyperglycaemic (large doses) and hypoglycaemic (smaller doses) actions in animals.\(^{\text{G41}}\) Antibacterial properties have also been documented. Alantolactone and isoalantolactone have been reported to exhibit high bactericidal and fungicidal properties *in vitro*.\(^{\text{G41}}\)

The volatile oil has been reported to exert a potent smooth muscle relaxant effect *in vitro* on guinea–pig ileal and tracheal muscle.\(^{\text{1}}\)

Various activities have been documented for *Inula racemosa*: an extract lowered plasma insulin and glucose concentrations in rats 75 minutes after oral administration,\(^{\text{2}}\) counteracted adrenaline–induced hyperglycaemia in rats,\(^{\text{2}}\) exhibited negative inotropic and chronotropic effects on the frog heart,\(^{\text{2}}\) and provided a preventative and curative action against experimentally induced myocardial infarction in rats.\(^{\text{3}}\) Pretreatment was found to be most effective.\(^{\text{3}}\)

Sesquiterpene lactones with antitumour activity have been isolated from *Helenium microcephalum*.\(^{\text{4,5}}\)

**Clinical studies**

Alantolactone has been used as an anthelmintic in the treatment of roundworm, threadworm, hookworm and whipworm infection.\(^{\text{G44 G45}}\)

*Inula racemosa* has been reported to prevent ST-segment depression and T-wave inversion in patients with proven ischaemic heart disease,\(^{\text{2}}\) and to have a beneficial effect on angina pectoris.\(^{\text{6}}\)
Side-effects, Toxicity

Elecampane has been reported to cause allergic contact dermatitis.\(^{(G51)}\) Sensitising properties have been documented for the volatile oil\(^{(G51 G58)}\) and for alantolactone and isoalantolactone.\(^{(7)}\) *In vitro* cytotoxicity has been reported for alantolactone and isoalantolactone.\(^{(8)}\)
Contraindications, Warnings

Elecampane may cause an allergic reaction, particularly in individuals with an existing allergy or sensitivity to other plants in the Asteraceae family. Elecampane may interfere with existing hypoglycaemic and antihypertensive treatment.

Pregnancy and lactation
The safety of elecampane taken during pregnancy has not been established. In view of the lack of toxicity data, the use of elecampane during pregnancy and lactation should be avoided.
The pharmacological actions documented for elecampane seem to be attributable to the sesquiterpene lactone constituents, in particular alantolactone and isoalantolactone. The demulcent action of mucilage and reported \textit{in vivo} antispasmodic activity of the volatile oil support the traditional uses of this remedy in coughs. In addition, alantolactone has been utilised as an anthelmintic. A number of interesting cardiovascular activities have been documented for a related species, \textit{I. racemosa}. Whether the constituents responsible for these actions are also present in elecampane is unclear. In view of the paucity of toxicity data for elecampane, excessive or prolonged use should be avoided.


7. Stampf JL et al. The sensitising capacity of helenin and two of its main constituents the sesquiterpene lactones alantolactone and isoalantolactone: a comparison of epicutaneous and intradermal sensitising methods in different strains of guinea pig. *Contact Dermatitis* 1982; **8**: 16–24. (PubMed)

Species (Family)

*Ephedra sinica* Stapf., *E. equisetina*, *E. intermedia*, *E. geriardiana*, *E. major* and other *Ephedra* species that contain ephedrine (Ephedraceae)
Synonym(s)
Cao Ma Huang (Chinese Ephedra), Herba Ephedrae, Ma Huang. Ephedra (and some other herbs) has also been referred to as ‘herbal ecstasy’. 
Part(s) Used

Aerial parts
Pharmacopoeial and Other Monographs

Complete German Commission E\(^{(G^3)}\)

Martindale 33rd edition\(^{(G^67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G^36)}\)

WHO volume 1 1999\(^{(G^63)}\)
Ephedra is not included in the GSL.\textsuperscript{(G37)}

Ephedra is included in Parts II and III of SI 2130.\textsuperscript{(1)} This allows supply of ephedra (maximum dose of 600 mg and a maximum daily dose of 1800 mg) following a one–to–one consultation with a practitioner.

Ephedrine and pseudoephedrine are not included on the GSL. Both are prescription–only medicines (POM), but can be supplied through pharmacies at certain permitted doses, as follows. Ephedrine for internal preparations: maximum dose 30 mg, maximum daily dose 60 mg; nasal preparations, ephedrine 2%. Pseudoephedrine hydrochloride for internal preparations: maximum dose 60 mg, maximum daily dose 240 mg; prolonged–release preparations: maximum dose 120 mg, maximum daily dose 240 mg. Pseudoephedrine sulfate for internal preparations: maximum dose 60 mg, maximum daily dose 180 mg.
Constituents

Alkaloids
0.5–2.0%. Mainly (−)-ephedrine (30–90% in most species, except E. intermedia) and (+)-pseudoephedrine, also (−)-norephedrine, (+)-nor pseudoephedrine, (−)-methylephedrine and (+)-methylpseudoephedrine. (2,G63)

Volatile oil
Mainly terpenoids (e.g. α-terpineol, limonene, tetramethylpyrazine, terpinen-4-ol, linalol). (3)

Other constituents
Tannins (catechin, gallic acid), ephedrains (glycans) and acids (citric, malic, oxalic).

Roots

Alkaloids
Ephedroxane, ephedradines A to D, feruloylhistamine and maokonine. (4)

Flavonoids
A flavonoflavonol (ephedrannin A), bisflavonols (mahuannins A to D). (4)
Food Use

Ephedra is not used in foods.
Herbal Use

Ephedra has traditionally been used for the treatment of bronchial asthma, hayfever, coughs and colds, fever, urticaria, enuresis, narcolepsy, myasthenia gravis, chronic postural hypotension and rheumatism. (G32 G34 G36 G49 G54 G63 G64)

It is stated to have vasoconstricting, bronchodilating and central stimulating properties. (G56) Modern interest in ephedra is focused on its use in cough and bronchitis, (G56) and in nasal congestion due to hay fever, allergic rhinitis, common cold and sinusitis. (G63) There is also interest in the potential of ephedra as an appetite suppressant.
Dosage

**Herb**
1.2–2.3 g cut herbs containing approximately 1.3% (13 mg/g) total alkaloids.\(^{(G4)}\)

**Extract**
Adults: 15–30 mg alkaloids (maximum daily dose 300 mg), calculated as ephedrine.\(^{(G56)}\)

**Tincture**
6–8 mL (1 : 4) three times daily.\(^{(G36)}\)

In 1997, the US Food and Drugs Administration (FDA) proposed restrictions on the use of ephedra, although these restrictions have not, to date, been implemented. The FDA proposals included a restriction on the maximum dose of ephedrine: 8 mg taken every 6 hours to a maximum daily dose of 24 mg for no more than seven days.\(^{(5,G56)}\)
The pharmacological properties of ephedra are due to the presence of ephedrine, pseudoephedrine and other ephedra alkaloids (see Constituents). Ephedrine and pseudoephedrine are sympathomimetic agents that have direct and indirect effects on both α- and β-adrenoceptors, as well as stimulating the central nervous system (CNS). Pseudoephedrine is stated to have less pressor activity and fewer CNS effects than ephedrine.

**In vitro and animal studies**

Pharmacological activities documented for ephedrine and/or pseudoephedrine in vitro or in vivo (animals) include smooth muscle relaxant, cardiovascular, anti-inflammatory, immuno modulatory, CNS stimulatory and antimicrobial effects. The pharmacology of ephedra and its constituent alkaloids has been reviewed.

Ephedrine and pseudoephedrine have been stated to have a relaxant effect on bronchial smooth muscle in isolated rabbit lung and bronchi. Relaxant effects on gastrointestinal smooth muscle have also been noted.

Ephedrine has been shown to cause vasoconstriction and to have hypertensive effects in several animal models. Maokonine, a constituent of ephedra root, has been reported to be have hypertensive effects in anaesthetised rats. By contrast, other constituents of ephedra roots, such as ephedrannin A and feruloylhistamine, have been reported to have hypotensive activity. An aqueous extract of ephedra and its alkaloid fraction increased blood pressure, heart rate and blood glucose concentration in anaesthetised dogs following intravenous administration.

Anti-inflammatory activity has been documented for ephedrine and pseudoephedrine in carrageenan–induced hind–paw oedema in mice. Oral administration of ephedrine and pseudoephedrine also inhibited hind–paw oedema induced by histamine, serotonin, bradykinin and prostaglandin E₁. Crude extracts of ephedra have been reported to inhibit complement in vitro. Further investigation, using an aqueous extract of E. sinica leaves, showed that the classical complement–inhibiting component of ephedra inhibited the classical complement pathway in sera from several species, including human, pig, guinea–pig, rat and rabbit.

In vitro antibacterial activity against several species, including *Staphylococcus aureus*, has been reported.
In vitro studies have assessed the cytotoxicity of extracts of ephedra prepared under various conditions (e.g. using ground or unground material boiled for 0.5 or 2 hours) against a range of cell lines, including a human hepatoblastoma cell line (HepG2), a mouse neuroblastoma cell line (Neuro–2a) and a mouse fibroblastoma cell line.\(^{(15)}\) Ephedrine and ephedra extracts prepared from ground material appeared to be significantly more cytotoxic in these cell lines than did preparations from unground material. Also, Neuro–2a cells were more sensitive to ephedra extracts than were the other cell lines tested. Findings of this in vitro work also indicated that ephedra contains toxins other than ephedrine, as IC\(_{50}\) values were lower (i.e. indicating greater cytotoxicity) for ephedra extracts than for ephedrine alone.

Clinical studies

Pharmacokinetics
Ephedrine and pseudoephedrine are readily absorbed from the gastrointestinal tract and are excreted, largely unchanged, in the urine.\(^{(G43)}\) Small amounts of metabolites following hepatic metabolism may be produced. The half–lives of ephedrine and pseudoephedrine range from 3 to 6 hours and from 5 to 8 hours, respectively, depending on urinary pH.\(^{(G43)}\)

In a study involving 12 healthy volunteers aged 23–40 years, four capsules of an ephedra product were administered twice, 9 hours apart. Each capsule contained ephedra 375 mg (\(E.\sinica\)), with a mean (standard deviation (SD)) ephedrine content of 4.84 (0.45) mg.\(^{(16)}\) The half–life was reported to be 5.2 hours, maximum plasma concentration (\(C_{\text{max}}\)) was 81.0 ng/mL, the time to reach \(C_{\text{max}}\) (\(t_{\text{max}}\)) was 3.9 hours, and clearance was 24.3 L/hour.

In a randomised, crossover study, 10 healthy volunteers received ephedrine 25 mg or one of three ephedrine–containing nutritional supplements on one day during different phases of the study, each with a one–week washout period.\(^{(17)}\) Following single–dose administration of ephedrine 25 mg, mean (SD) half–life, \(C_{\text{max}}\), \(t_{\text{max}}\) and clearance were found to be 5.37 (1.67) hours, 86.5 (15.4) ng/mL, 2.81 (1.35) hours and 28.5 (5.92) L/hour, respectively.

Therapeutic effects
The pharmacological properties of ephedrine and pseudoephedrine in humans have been documented and include cardiovascular, bronchodilator and CNS stimulant effects.\(^{(G43 \ G63)}\)

Ephedrine is stated to raise blood pressure by increasing cardiac output and also by peripheral vasoconstriction. Ephedrine relaxes bronchial smooth
muscle, reduces intestinal tone and motility, relaxes the bladder wall and reduces the activity of the uterus. Ephedrine is a CNS stimulant; this has led to its investigation for use in assisting weight loss.

A randomised, double-blind, placebo-controlled trial assessed the effects of a herbal combination preparation which included ephedra and other herbal ingredients.\(^{18}\) In the study, 67 overweight to obese individuals (body mass index 29–35 kg/m\(^2\)) received the ephedra-containing preparation, or placebo, for eight weeks. Among the 48 participants who completed the study (24 in each group), a greater mean (SD) weight loss was noted for the treatment group, compared with the placebo group (4.0 (3.4) kg versus 0.8 (2.4) kg, respectively; \(p < 0.006\)).
Side-effects, Toxicity

The most common adverse effects of ephedrine and pseudoephedrine are tachycardia, anxiety, restlessness and insomnia. Tremor, dry mouth, impaired circulation to the extremities, hypertension and cardiac arrhythmias may also occur with ephedrine, and skin rashes and urinary retention have been reported for pseudoephedrine. There are isolated reports of hallucinations in children following use of pseudoephedrine.

In the US, adverse effects have been reported following self-treatment with products containing ephedra alkaloids marketed for several uses, including as an aid to weight loss, to increase athletic performance, and as an alternative to illegal drugs of abuse.

A review assessed 140 reports of adverse events related to the use of products containing ephedra alkaloids, usually combined with caffeine, submitted to the US FDA between June 1997 and March 1999. The main reasons for use of these products were weight loss (59%) and to increase athletic performance (16%); the reason for use was unknown in 17% of cases. Thirty-one per cent of cases (n = 43) were considered to be ‘definitely’ or ‘probably’ related to the use of products containing ephedra alkaloids, and a further 31% (n = 44) were judged to be ‘possibly’ related; for 29 cases, insufficient information was available to assess causation, and 24 cases were deemed to be ‘unrelated’ to use of these products. In several cases, individuals were thought to be ingesting up to 60 mg ephedra alkaloids daily. Of the 87 cases where causality was assessed, cardiovascular symptoms (mainly hypertension, palpitations, tachycardia) were the most common adverse events (47%). The most common CNS events were stroke (n = 10) and seizures (n = 7). Where events were ‘definitely’ or ‘probably’ related (n = 43), clinical outcomes were death (three cases), permanent impairment (seven) and ongoing treatment (four); a full recovery occurred in 29 cases.

In a randomised, double-blind, placebo-controlled trial of a herbal supplement containing ephedra (72 mg/day) and guarana (240 mg/day), as well as other herbal ingredients, 23% (n = 8) of participants in the treatment group withdrew from the study because of adverse events (e.g. dry mouth, insomnia, headache) that may have been treatment-related; there were no withdrawals among placebo recipients.

There are isolated reports of myocarditis, exacerbation of autoimmune hepatitis, acute hepatitis, nephrolithiasis and psychiatric complications associated with the use of ephedra-containing products.
There is a report of sudden death associated with ephedrine toxicity in a 23–
year-old man.\(^{(25)}\) Several other reports also document psychosis and renal
calculi following chronic use or misuse of ephedrine.\(^{(G18)}\)

In a study involving 12 normotensive adults who ingested four capsules each
containing 375 mg powdered ephedra, followed by four more capsules nine
hours later, a statistically significant increase in heart rate, compared with
baseline values, was noted in six participants, although effects on blood
pressure were variable.\(^{(16)}\)

A study involving 47 dogs who were considered to have accidentally ingested
herbal products containing ephedra and guarana reported that most dogs
(80\%) developed clinical signs of toxicosis, within eight hours of ingestion,
which persisted for up to 48 hours.\(^{(26)}\) Hyperactivity, tremors, seizures and
behaviour changes were reported in 83\% of dogs; other signs and symptoms
included vomiting, tachycardia and hyperthermia.
Contra–indications, Warnings

Ephedra is stated to be contra–indicated in coronary thrombosis, diabetes, glaucoma, heart disease, hypertension, thyroid disease, phaeochromocytoma and enlarged prostate.\(^{(G63)}\) Another source states that ephedrine (and, therefore, ephedrine–containing products) should be used with caution in patients with diabetes, ischaemic heart disease, hypertension, hyperthyroidism, renal impairment and angle–closure glaucoma, and that in patients with prostate enlargement, ephedrine may increase difficulty with micturition.\(^{(G43)}\) It has been recommended to reduce the dose or discontinue treatment if nervousness, tremor, sleeplessness, loss of appetite or nausea occur with use of ephedra preparations.\(^{(G63)}\)

Ephedrine–containing products should be avoided in patients receiving monoamine oxidase inhibitors as concomitant treatment may lead to a hypertensive crisis.\(^{(G43)}\) Ephedrine should also be avoided or used with caution in patients undergoing anaesthesia with cyclopropane, halothane or other volatile anaesthetics. There may be an increased risk of arrhythmias in patients receiving ephedrine together with cardiac glycosides, quinidine or tricyclic antidepressants, and there is an increased risk of vasoconstrictor or pressor effects in patients receiving ergot alkaloids or oxytocin.\(^{(G43)}\)

There is a report of a professional sportsman who tested positive for norpseudoephedrine after having consumed a liquid herbal product listing ephedra as one of the 15 ingredients.\(^{(27)}\)

**Pregnancy and lactation**

There are no reliable data on the use of ephedra during pregnancy and lactation. The safety of ephedra during pregnancy and lactation has not been established and its use should be avoided.
Pharmaceutical Comment

The activities of ephedra are due to the presence of the ephedra alkaloids; of these, the pharmacological effects of ephedrine and pseudoephedrine are most well-documented and support their modern uses. There is less information on the pharmacological effects of ephedra extracts and clinical trials, in particular, are generally lacking.

In view of the safety concerns regarding the use of ephedra products, individuals wishing to use these products should be advised to consult an appropriately trained health care professional. Pharmacists and other health care professionals should be aware that ephedra may be included in unlicensed herbal products and food supplements under the name Ma Huang. Such products will not include reference to ephedra in the labelling.
References

See also General References G5 G18 G29 G31 G32 G34 G36 G43 G49 G54 G56 G63 G64.

5. Blumenthal M. Ephedra update: industry coalition asks FDA to adopt national labeling guidelines on ephedra; offers co-operative research with NIH. Herbal Gram 2000; 50: 64–65.


Eucalyptus
Species (Family)

*Eucalyptus globulus* Labill. (Myrtaceae)
Synonym(s)
Fevertree, Gum Tree, Tasmanian Bluegum
Part(s) Used

Leaf
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)

BP 2002\(^{(G71)}\)

Complete German Commission E\(^{(G3)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)

Ph Eur 2004\(^{(G72)}\)
Legal Category (Licensed Products)

GSL$^{(G37)}$
Constituents
See General References G2 G22 G41 G48 G64.

**Flavonoids**
Eucalyptrin, hyperoside, quercetin, quercitrin and rutin.

**Volatile oils**
0.5–3.5%. Eucalyptol (cineole) 70–85%. Others include monoterpenes (e.g. α-pinene, β-pinene, d-limonene, p-cymene, α-phellandrene, camphene, γ-terpinene) and sesquiterpenes (e.g. aromadendrene, alloaromadendrene, globulol, epiglobulol, ledol, viridiflorol), aldehydes (e.g. myrtenal) and ketones (e.g. carvone, pinocarvone).

**Other constituents**
Tannins and associated acids (e.g. gallic acid, protocatechuic acid), caffeic acid, ferulic acids, gentisic acid, resins and waxes
Eucalyptus is listed by the Council of Europe as a natural source of food flavouring (leaves, flowers and preparations: category N4, with limits on eucalyptol) (see Appendix 23).\(^{(G17)}\) Both eucalyptus and eucalyptol (cineole) are used as flavouring agents in many food products.\(^{(G41)}\) In the USA, eucalyptus is approved for food use and eucalyptol is listed as a synthetic flavouring agent.\(^{(G41)}\)
Herbal Use

Eucalyptus leaves and oil have been used as an antiseptic, febrifuge and expectorant. (G2 G41 G64)
Dosage

**Eucalyptol**
(cineole BPC 1973) 0.05–0.2 mL.

**Eucalyptus Oil**
(BPC 1973) 0.05–0.2 mL.

**Fluid extract**
2–4 g.

**Oil for local application**
30 mL oil to 500 mL lukewarm water.
Pharmacological Actions

*In vitro* and animal studies

Hypoglycaemic activity in rabbits has been documented for a crude leaf extract rich in phenolic glycosides. Purification of the extract resulted in a loss of activity.\(^{(G41)}\) Expectorant and antibacterial activities have been reported for eucalyptus oil and for eucalyptol.\(^{(G41)}\) Various *Eucalyptus* species have been shown to possess antibacterial activity against both Gram–positive and Gram–negative organisms. Gram–positive organisms were found to be the most sensitive, particularly *Bacillus subtilis* and *Micrococcus glutamious*.\(^{(1)}\)

*In vitro* antiviral activity against influenza type A has been documented for quercitrin and hyperoside.\(^{(G41)}\)

Clinical studies

Eucalyptus oil oil has been taken orally for catarrh, used as an inhalation and applied as a rubefacient.\(^{(G45)}\) A plant preparation containing tinctures of various herbs including eucalyptus has been used successfully in the treatment of chronic suppurative otitis.\(^{(2)}\) The efficacy of the preparation was attributed to the antibacterial and anti–inflammatory actions of the herbs included.
Side-effects, Toxicity

Externally, eucalyptus oil is stated to be generally non-toxic, non-sensitising and non-phototoxic.\(^{(G58)}\) Undiluted eucalyptus oil is toxic and should not be taken internally. A dose of 3.5 mL has proved fatal.\(^{(G45)}\) Symptoms of poisoning with eucalyptus oil include epigastric burning, nausea and vomiting, dizziness, muscular weakness, miosis, a feeling of suffocation, cyanosis, delirium and convulsions.
Contra–indications, Warnings

Eucalyptus may interfere with existing hypoglycaemic therapy. Eucalyptus oil should be diluted before internal or external use.

_Pregnancy and lactation_

Eucalyptus oil should not be taken internally during pregnancy.
Pharmaceutical Comment

Eucalyptus is characterised by its volatile oil components. Antiseptic and expectorant properties have been attributed to the oil, in particular to the principal component eucalyptol. The undiluted oil is toxic if taken internally. Essential oils should not be applied to the skin unless they are diluted with a carrier vegetable oil.
See also General References G2 G3 G9 G15 G16 G19 G22 G28 G31 G36 G37 G41 G43 G48 G58 G64.


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Euphorbia
Species (Family)

*Euphorbia hirta* L. (Euphorbiaceae)
Synonym(s)

_Euphorbia capitata_ Lam., _Euphorbia pilulifera_ L., Pillbearing Spurge, Snakeweeds
Part(s) Used

Herb
Pharmacopoeial and Other Monographs

BHP 1983\(^{G7}\)

Martindale 33rd edition\(^{G67}\)
Legal Category (Licensed Products)

GSL$^{G37}$
Constituents

See General References G41 G48 G64.

**Flavonoids**
Leucocyanidin, quercetin, quercitrin and xanthorhamnin.

**Terpenoids**
α- and β-Amyrin, taraxerol and esters, friedelin; campesterol, sitosterol and stigmasterol (sterols).

**Other constituents**
Choline, alkanes, inositol, phenolic acids (e.g. ellagic, gallic, shikimic), sugars and resins.
Food Use

Euphorbia is not used in foods.
Herbal Use

Euphorbia is stated to be used for respiratory disorders, such as asthma, bronchitis, catarrh and laryngeal spasm. It has also been used for intestinal amoebiasis. (G7 G64)
Dosage

*Herb*
120–300 mg or as an infusion. (G7)

*Liquid Extract of Euphorbia*
(BPC 1949) 0.12–0.3 mL.

*Euphorbia Tincture*
(BPC 1923) 0.6–2.0 mL.
Pharmacological Actions

In vitro and animal studies

Euphorbia has been reported to have antispasmodic and histamine-potentiating properties.\(^{(G41)}\) Smooth muscle relaxing and contracting activities have been exhibited by euphorbia in vitro (guinea-pig ileum) and have been attributed to shikimic acid and to choline, respectively.\(^{(1)}\)

In vivo antitumour activities have been documented for euphorbia.\(^{(G41)}\)

Antibacterial activity in vitro versus both Gram-positive and Gram-negative bacteria has been documented for euphorbia.\(^{(2)}\) Stem extracts were slightly more active than leaf extracts. In vitro amoebicidal activity versus Entamoeba histolytica has been reported for a euphorbia decoction.\(^{(3)}\)
Side–effects, Toxicity

None documented for euphorbia. Carcinogenic properties in mice have been reported for shikimic acid, although no mutagenic activity was observed in the Ames assay. (G41)
Contra-indications, Warnings

None documented.

**Pregnancy and lactation**
The safety of euphorbia has not been established. Euphorbia has been reported to cause both contraction and relaxation of smooth muscle. In view of the lack of pharmacological and toxicity data, the use of euphorbia during pregnancy and lactation should be avoided.
Pharmaceutical Comment

There is little published information concerning euphorbia, although documented actions observed in animals do support the traditional herbal uses. There is a lack of information concerning toxicity, although the documented constituents of euphorbia do not indicate any obvious toxic component. Nevertheless, excessive or prolonged ingestion should be avoided.
References

See also General References G7 G10 G31 G37 G41 G44 G48 G64.


Evening Primrose
Species (Family)

*Oenothera* species including *Oenothera biennis* L. (Onagraceae)
Synonym(s)

King’s Cureall
Part(s) Used

Seed oil
Pharmacopoeial and Other Monographs

Martindale 33rd edition\textsuperscript{(G67)}

Mills and Bone\textsuperscript{(G50)}

PDR for Herbal Medicines 2nd edition\textsuperscript{(G36)}
Legal Category (Licensed Products)

Evening primrose is not included in the GSL.\textsuperscript{(G37)} Gamolenic acid is a prescription–only medicine.
Constituents

**Fixed oils**
14%. cis-Linoleic acid (LA) 72% (65–80%), cis–gammalinolenic acid (gamolenic acid, GLA) 2–16%, oleic acid 9%, palmitic acid 7% and stearic acid (3%).\(^{(1-5)}\)
Food Use

Evening primrose root has been used as a vegetable with a peppery flavour.\(^5\) The seed oil has been used as a food supplement for many years. LA and gamolenic acid are both essential fatty acids (EFAs), with LA representing the main EFA in the diet, whilst gamolenic acid is found in human milk, in oats and barley, and in small amounts in a wide variety of common foods.\(^{4,5}\)
Herbal Use

An infusion of the whole plant is reputed to have sedative and astringent properties, and has traditionally been used for asthmatic coughs, gastrointestinal disorders, whooping cough and as a sedative pain killer.\(^{(5)}\) Externally, poultices were reputed to ease bruises and to speed wound-healing.\(^{(5)}\)

Evening primrose oil (EPO) is licensed for the treatment of atopic eczema, and cyclical and non-cyclical mastalgia. Other conditions in which evening primrose oil is used include premenstrual syndrome, psoriasis, multiple sclerosis, hypercholesterolaemia, rheumatoid arthritis, Raynaud’s phenomenon, Sjögren’s syndrome, postviral fatigue syndrome, asthma and diabetic neuropathy.\(^{(1-3,5)}\)
Dosage

Recommended doses for evening primrose oil are specific to the condition being treated.

Daily doses for a licensed evening primrose oil product are 6–8 g (adults) and 2–4 g (children) in atopic eczema. In cyclical and non-cyclical mastalgia, a daily dose of 3 to 4 g is recommended. These doses are based on a standardised gamolenic acid content of 8%. No special precautions are noted for the elderly. The oil may be swallowed directly, mixed with milk or another liquid, or taken with food.

A patient may need to receive evening primrose oil for a period of three months before a clinical response is observed.
Pharmacological Actions

The pharmacological actions of evening primrose oil have been reviewed.\(^{(1–3,5)}\)

The actions of evening primrose oil are attributable to the essential fatty acid content of the oil and to the involvement of these compounds in prostaglandin biosynthetic pathways.

Gamolenic acid and its metabolite dihomogamma-linolenic acid (DGLA) are precursors of both the inflammatory prostaglandin \(E_2\) (PGE\(_2\)) series via arachidonic acid (AA), and of the less inflammatory prostaglandin \(E_1\) (PGE\(_1\)) series. Actions attributed to PGE\(_1\) include anti-inflammatory, immunoregulatory and vasodilatory properties, inhibition of platelet aggregation and cholesterol biosynthesis, hypotension and elevation of cyclic AMP (inhibits phospho lipase A\(_2\), see below).\(^{(1–3)}\)

Dietary supplementation with gamolenic acid has been noted to have a favourable effect on the DGLA : AA ratio. Although an increase in arachidonic acid concentrations is also seen, this is much smaller and less consistent compared to the increase seen for DGLA.\(^{(3)}\) Contributory factors to this negative effect on arachidonic acid are PGE\(_1\) and 15-hydroxy-DGLA. The latter inhibits conversion of arachidonic acid to inflammatory lipoxygenase metabolites including leukotrienes, whilst PGE\(_1\) inhibits the enzyme phospholipase A\(_2\) which is required for the mobilisation of arachidonic acid from phospholipid membrane stores.\(^{(3)}\) In addition, DGLA desaturation to arachidonic acid is a rate-limiting step in humans and proceeds very slowly.\(^{(3)}\)

Gamolenic acid is not normally obtained directly from dietary sources and the body relies on metabolic conversion from dietary LA. This conversion is readily saturable and is considered to be the rate-limiting step in the production of gamolenic acid. A reduced rate of LA conversion to gamolenic acid has been observed in a number of clinical situations including ageing, diabetes, cardiovascular disorders and high cholesterol concentrations, high alcohol intake, viral infections, cancer, nutritional deficits, atopic eczema and premenstrual syndrome.\(^{(1–3)}\) Direct dietary supplementation with gamolenic acid effectively bypasses this rate-limiting conversion step and has a beneficial effect on the ratio of inflammatory:non-inflammatory prostaglandin compounds.

Evening primrose oil represents a good source of both LA and, more
importantly, of gamolenic acid. Numerous papers have been published on the biochemical rationale for the therapeutic uses of evening primrose oil and on its efficacy in various disease states associated with low concentrations of gamolenic acid. The use of evening primrose oil in various disease states which include atopic eczema, premenstrual syndrome including mastalgia, diabetic neuropathy, rheumatoid arthritis, Sjögren’s syndrome, cardiovascular, renal, hepatic and gastrointestinal disorders, viral infections, endometriosis, schizophrenia, alcoholism, Alzheimer’s disease and cancers has been reviewed.  

Atopic eczema
An inherited slow rate of 6–desaturation (LA to gamolenic acid conversion) has been documented in this condition. Normal or elevated concentrations of LA are associated with reduced concentrations of their metabolites. Randomised, double-blind, placebo-controlled trials have shown gamolenic acid to produce a highly significant improvement in all features of atopic eczema, especially in itch.  

The requirement for topical and oral steroids, histamines and antibiotics was also reduced. However, attention has been drawn to the conflicting evidence of clinical trials on evening primrose oil. Two large trials have not shown evidence of benefit whereas other trials have resulted in benefits, particularly for patients with moderate or severe eczema. Adequate doses of evening primrose oil for treatment of atopic eczema are 160–320 mg of gamolenic acid daily in children aged 1–12 years and 320–480 mg in adults for three months.  

Cyclical/non-cyclical mastalgia
PGE\textsubscript{1} is thought to modulate the action of prolactin. Abnormal concentrations may result in an excessive peripheral action of prolactin. Several placebo-controlled studies have demonstrated that gamolenic acid is better than placebo in the treatment of both premenstrual syndrome and breast pain. Overall, cyclical mastalgia responds better than non-cyclical to all treatments (danazol, bromocriptine, evening primrose oil).

Premenstrual syndrome
The use of evening primrose oil for the treatment of premenstrual syndrome has been rationalised on the grounds that hypersensitivity to prolactin is due to low levels of PGE\textsubscript{1}. High levels of linoleic acid and low levels of gamma-linolenic acid have been observed for patients with premenstrual syndrome. Several clinical studies have been reported and the conclusions vary from no beneficial effects being observed to marked improvements.
Diabetic neuropathy
Diabetes has been associated with reduced ability to desaturate essential fatty acids, with deficits resulting in abnormal neuronal membrane structure. Animal studies have shown that diabetic neuropathy can be either prevented or reversed by the provision of gamolenic acid as evening primrose oil. In humans, a double-blind, placebo-controlled trial has demonstrated reversal of diabetic neuropathy by gamolenic acid.\(^{(17)}\)

Multiple sclerosis
The results of clinical trials on the use of evening primrose oil for the treatment of multiple sclerosis are contradictory.\(^{(1,2)}\) Patients with recent onset or less severe forms of the disease are more likely to respond. Linoleic acid may have a beneficial effect on the severity and duration of relapses and on the progression of the disease.\(^{(1)}\) It is suggested that linoleic acid is involved in the immunosuppressive effect at the cellular level and may be of use when combined with a low animal fat/high polyunsaturated fat diet. \(^{(2)}\)

Rheumatoid arthritis
A randomised, double-blind trial has demonstrated a significant improvement in subjective symptoms of rheumatoid arthritis (RA) (indicated by a reduction in required non-steroidal anti-inflammatory drug treatment) in the active group receiving evening primrose oil compared with the placebo group. However, no objective changes were observed in any of the biochemical indicators of RA.\(^{(1-3)}\)

Sjögren’s syndrome
This disease is associated with the loss of secretions from exocrine glands throughout the body, but especially from the salivary and lacrimal glands. One of the features of EFA deficiency is exocrine gland atrophy. Placebo-controlled trials have shown a modest improvement in tear flow together with relief of lethargy, a prominent feature of the syndrome.\(^{(1,3)}\)

Coronary heart disease
Abnormal intake and metabolism of EFAs (both \(n\)-3 and \(n\)-6) are thought to be important risk factors for coronary heart disease (CHD), resulting in enhanced cholesterol and triglyceride biosynthesis, enhanced platelet aggregation and elevated blood pressure. Dietary supplementation with foods or oils rich in LA (\(n\)-6) or in marine (\(n\)-3) EFAs have been found to decrease significantly the risk of CHD, although it is considered that an optimum balance between \(n\)-3 and \(n\)-6 EFAs may well be important.\(^{(1-3,18)}\) Gamolenic acid has been reported to decrease blood pressure and platelet aggregation in both animal and human studies.\(^{(3)}\)
Renal disease
Renal tissue is especially rich in EFAs, and prostaglandins of the E series are believed to be important in maintaining adequate renal blood flow. Administration of gamolenic acid to animals has been reported to prevent or attenuate renal damage. A single placebo–controlled trial involving postrenal transplant patients demonstrated better graft survival rate for the group receiving evening primrose oil (45 patients) compared with the placebo group (44 patients).\(^{(3)}\)

Liver disease
PGE\(_1\) has been administered to patients with liver failure, and has been observed to exert some cytoprotective effect and to maintain the normal function of the liver. There is little experience of gamolenic acid supplementation in liver disease.\(^{(17)}\)

Gastrointestinal disorders
A double–blind placebo–controlled crossover trial has indicated a beneficial effect of evening primrose oil on irritable bowel syndrome exacerbated by pre menstrual syndrome. A beneficial effect superior to that of fish oil or placebo has been reported for evening primrose oil in ulcerative colitis. A protective effect of gamolenic acid against gastric ulceration has yet to be shown in humans.\(^{(3)}\)

Viral infections/postviral fatigue
A single placebo–controlled study has demonstrated significant beneficial effects in patients with well–defined postviral fatigue (PVF) receiving evening primrose oil compared with those receiving placebo. Symptoms arrested were muscle weakness, aches and pains, lack of concentration, exhaustion, memory loss, depression, dizziness and vertigo.\(^{(1,3)}\)

Endometriosis
A placebo–controlled trial has shown that gamolenic acid in combination with eicosapentaenoic acid (\(n\)-3 EFA metabolite) reduced symptoms of endometriosis in 90% women, whereas 90% of the placebo group reported no relief from symptoms.\(^{(3)}\)

Schizophrenia
It is believed that EFAs, in particular PGE\(_1\), antagonise the excessive central dopaminergic activity that is thought to be a possible cause of schizophrenia. Low concentrations of LA in plasma phospholipids have been observed in populations of schizophrenics from Ireland, England, Scotland, Japan and the USA. It is thought that a poor recovery rate from the disease is associated
with the presence of saturated fats in the diet, but not with unsaturated fats. Various open and placebo–controlled trials of gamolenic acid and DGLA supplementation have reportedly produced mixed results. Administration of evening primrose oil with co–factors known to be important in EFA metabolism (zinc, pyridoxine, niacin and vitamin C) enhanced the improvements in memory loss, schizophrenic symptoms and tardive dyskinesia that were observed in evening primrose oil–treated compared with placebo–treated patients.\(^{(1–3)}\)

**Alcoholism**
Evening primrose oil has been documented to reduce symptoms in the first three weeks of withdrawal, indicated by a reduced requirement for tranquillisers, and to significantly improve the rate of return to normal liver function. However, in the longer term, evening primrose did not affect the relapse rate.\(^{(3)}\)

**Dementia**
Alzheimer’s disease and other forms of dementia are associated with low serum concentrations of EFAs. A single placebo–controlled trial in patients with Alzheimer’s disease reported improvements in cerebral function in the evening primrose oil group compared with the placebo group.

**Hyperactivity in children**
Hyperactive children tend to have abnormal levels of essential fatty acids. No improvements in behavioural patterns and no changes in blood fatty acids were observed in one trial with evening primrose oil.\(^{(2)}\)

**Cancer**
*In vitro* studies have observed that malignant cells die following exposure to gamolenic acid and related fatty acids at concentrations that are non–lethal to normal cells. *In vitro* studies have shown gamolenic acid to inhibit the growth of various human cancer cell lines, and *in vivo* studies have described an inhibitory effect of gamolenic acid on tumour growth. Human studies are currently ongoing to assess the impact of gamolenic acid supplementation in various human cancers.\(^{(3)}\)
Side–effects, Toxicity

Evening primrose oil appears to be well tolerated with very few side–effects reported, despite it being available for many years in a number of countries as a food supplement.\(^{(3)}\) Mild gastrointestinal effects, indigestion, nausea and softening of stools and headache have occasionally occurred.\(^{(3,5)}\) It has been noted that there may be an increased risk of temporal lobe epilepsy in schizophrenic patients being treated with epileptogenic drugs such as phenothiazines.\(^{(6)}\) In cases of overdosage, symptoms of loose stools and abdominal pain have been noted. No special treatment is required.\(^{(6)}\)

Toxicity studies have indicated evening primrose oil to be non–toxic.\(^{(3)}\) The two principal components in evening primrose oil are LA and gamolenic acid. LA is commonly ingested as part of the diet. It has been estimated that the concentration of gamolenic acid provided by evening primrose oil is comparable to that metabolised in the body from normal dietary LA.\(^{(4)}\) In addition, it has been calculated that a breastfed infant receives a higher proportion (mg/kg) of LA and gamolenic acid from human milk compared to that received from evening primrose oil.\(^{(4)}\)
Contra–indications, Warnings

Evening primrose oil may have the potential to make manifest undiagnosed temporal lobe epilepsy, especially in schizophrenic patients and/or those who are already receiving known epileptogenic drugs such as phenothiazines.\(^{(6)}\) No epileptic events have been reported in patients not being treated with phenothiazines.\(^{(6)}\)

Pregnancy and lactation

Animal studies have indicated evening primrose oil to be non–teratogenic.\(^{(6)}\) However, data on the safety of evening primrose oil during human pregnancy are not available and therefore the risk of taking evening primrose oil during pregnancy should be carefully considered against the perceived benefit to the patient. Both LA and gamolenic acid are normally present in breast milk (see Side–effects, Toxicity) and therefore it is reasonable to assume that evening primrose oil may be taken while breast feeding.
Interest in the seed oil of the evening primrose plant lies in its essential fatty acid content, in particular in the linoleic acid (LA) and gamolenic acid (GLA) content. Both of these compounds are prostaglandin precursors and dietary gamolenic acid supplementation has been shown to increase the ratio of non-inflammatory to inflammatory prostaglandin compounds.

The use of evening primrose oil in various disease states associated with low gamolenic acid concentrations has been extensively investigated and a vast body of published literature is available. The beneficial effects of evening primrose oil in treating atopic eczema and mastalgia (cyclical/non-cyclical) have been recognised with product licences granted to evening primrose oil-containing preparations for these indications. However, doubt has also been expressed over the effectiveness of evening primrose oil in eczema. Alternative natural oil sources such as blackcurrant or borage (see Borage) that offer a higher gamolenic acid yield compared to evening primrose oil have been identified, although these oils have not been found to exhibit the same biological effects as those observed for evening primrose oil.

Evening primrose oil is reported to be virtually non-toxic with only minor adverse effects such as headache and nausea occasionally associated with its use. The range of potential uses for evening primrose oil is extensive and results of further human studies are awaited to establish its efficacy in various therapeutic conditions.
References

See also General References G5 G29 G32 G36 G43 G50 G56 G64.

Eyebright
Species (Family)

*Euphrasia* species including

i. *Euphrasia brevipila* Burnat & Greml

ii. *Euphrasia officinalis* L.

iii. *Euphrasia rostkoviana* Hayne (Scrophulariaceae)
Synonym(s)

Euphrasia
Part(s) Used
Herb
Pharmacopoeial and Other Monographs

BHP 1983\(^{(G7)}\)

Martindale 33rd edition\(^{(G67)}\)

Mills and Bone\(^{(G50)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

Eyebright is not included in the GSL.\(^{(G37)}\)
Constituents
See General References G2 G22 G40 G64.

Unless otherwise stated, constituents listed are for *E. officinalis*.

**Acids**
Caffeic acid, ferulic acid.\(^1\)

**Alkaloids**
Unidentified tertiary alkaloids, choline, steam volatile bases\(^1\)

**Amino acids**
Glycine, leucine and valine.

**Flavonoids**
Four compounds (unidentified). Quercetin and rutin stated to be absent.\(^1\)
Quercetin, quercitrin and rutin have been documented for *E. rostkoviana*.

**Iridoids**
Aucubin 0.05%. Additional glycosides have been reported for related *Euphrasia* species including catalpol, euphroside, eurostoside, geniposide, ixoroside and mussaenoside for *E. rostkoviana*.\(^2–5\)

**Phenethyl glycosides**
Dehydroconiferyl alcohol–4-β-D-glucoside\(^3\) and eukovoside (3,4–dihydroxy–4–phenethyl-O-α-L-rhamnoside(13)-4-O-isofeuoyl-β-D- glucoside)\(^4\) from *E. rostkoviana*.

**Tannins**
About 12%. Condensed and hydrolysable; gallic acid is among the hydrolysis products.\(^1\)

**Volatile oils**
About 0.2%. Seven major and numerous minor components, mainly unidentified; four of the major compounds are thought to be aldehydes or ketones.\(^1\)

**Other constituents**
Bitter principle, β-carotene, phytosterols (e.g. β-sitosterol, stigmasterol),\(^1\)
resin, carbohydrates (e.g. arabinose, glucose, galactose) and vitamin C.
Food Use

Eyebright is listed by the Council of Europe as a natural source of food flavouring (category N3). This category indicates that eyebright can be added to foodstuffs in the traditionally accepted manner, although there is insufficient information available for an adequate assessment of potential toxicity. (G16)
Herbal Use

Eyebright is stated to possess anticitarrhal, astringent and anti-inflammatory properties. Traditionally it has been used for nasal catarrh, sinusitis and specifically for conjunctivitis when applied locally as an eye lotion.\(^{(G2 G7 G64)}\)
Dosage

**Dried herb**
2–4 g or by infusion three times daily.\(^{(G7)}\)

**Liquid extract**
2–4 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)

**Tincture**
2–6 mL (1 : 5 in 45% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

*In vitro* and animal studies

None documented for eyebright. Caffeic acid is bacteriostatic,\(^{(1)}\) and a purgative action in mice has been documented for iridoid glycosides.\(^{(6)}\) The purgative action of aucubin is approximately 0.05 times the potency of sennosides, with onset of diarrhoea stated to occur more than 6 hours after aucubin administration.\(^{(6)}\) Tannins are known to possess astringent properties.
Side-effects, Toxicity

It has been stated that 10–60 drops of eyebright tincture could induce toxic symptoms including mental confusion and cephalalgia, raised pressure in the eyes with lachrymation, pruritus, redness, swelling of the eyelid margins, dim vision, photophobia, weakness, sneezing, nausea, toothache, constipation, cough, dyspnoea, insomnia, polyuria and diaphoresis.\(^{G22}\)
Contra–indications, Warnings

The use of eyebright for ophthalmic application has been discouraged.\(^{(G60)}\)

Pregnancy and lactation

The safety of eyebright has not been established. In view of the lack of pharmacological and toxicity data, the use of eye bright during pregnancy and lactation should be avoided.
Pharmaceutical Comment

Limited information is available regarding the constituents of eyebright and it is unclear which *Euphrasia* species is most commonly utilised. In addition, eyebright is also used as a common name for plants other than *Euphrasia* species. Little scientific information was found to justify the reputed herbal uses, although tannin constituents would provide an astringent effect. The use of home-made preparations for ophthalmic purposes should be avoided. Little is known regarding the toxicity of eyebright and, in view of the reported presence of unidentified alkaloids, it should be used with caution avoiding excessive doses.
References

See also General References G2 G7 G16 G22 G31 G32 G36 G37 G40 G44 G50 G60 G64.


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Species (Family)

*Chamaelirium luteum* (L.) A. Gray (Liliaceae)
Synonym(s)
Part(s) Used
Rhizome, root
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL[^G37]
Constituents

See General References G40 G48 G64.

Limited chemical information is available on false unicorn. It is stated to contain a steroidal saponin glycoside, chamaelirin, and another glycoside helonin.
Food Use

False unicorn is not used in foods.
Herbal Use

False unicorn is stated to possess an action on the uterus. Traditionally it has been used for ovarian dysmenorrhea, leucorrhoea and specifically for amenorrhoea. It is reported to be useful for vomiting of pregnancy and threatened miscarriage.\(^{(G7 \ G8 \ G64)}\)
**Dosage**

*Dried rhizome/root*
1–2 g or by infusion three times daily.\(^{(G7)}\)

*Liquid extract*
1–2 mL (1 : 1 in 45% alcohol) three times daily.\(^{(G7)}\)

*Tincture*
2–5 mL (1 : 5 in 45% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

None documented.
Side–effects, Toxicity

No reported side–effects or documented toxicity studies were located. It is stated that large doses of false unicorn may cause nausea and vomiting.\footnote{G7}
Contra–indications, Warnings

None documented.

**Pregnancy and lactation**
The safety of false unicorn has not been established. In view of the lack of phytochemical, pharmacological and toxicity data, and its reputed action as a uterine tonic, the use of false unicorn during pregnancy and lactation should be avoided.
The chemistry of false unicorn is poorly documented and no scientific evidence was located to justify the herbal uses. In view of this and the lack of toxicity data, the use of false unicorn should be avoided.
References

See General References G9 G31 G36 G37 G40 G43 G48 G64.
Fenugreek
Species (Family)

Trigonella foenum-graecum L. (Leguminosae)
**Synonym(s)**

Bockshornsame
Part(s) Used

Seed
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)

BP 2002\(^{(G71)}\)

Complete German Commission E\(^{(G3)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)

Ph Eur 2004\(^{(G72)}\)
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents

See General References G2 G41 G48 G64.

**Alkaloids**
Pyridine–type. Gentianine, trigonelline (up to 0.13%), choline (0.05%).

**Proteins and amino acids**
Protein (23–25%) containing high quantities of lysine and tryptophan. Free amino acids include 4–hydroxyisoleucine (0.09%), histidine, lysine and arginine.

**Flavonoids**
Flavone (apigenin, luteolin) glycosides including orientin and vitexin, quercetin (flavonol).

**Saponins**
0.6–1.7%. Glycosides yielding steroidal sapogenins diosgenin and yamogenin (major), with tigogenin, neotigogenin, gitogenin, neogitogenin, smilagenin, sarsasapogenin, yuccagenin;\(^1\) fenugreekine, a sapogenin–peptide ester involving diosgenin and yamogenin;\(^2\) trigofoenosides A–G (furostanol glycosides).\(^3–6\)

**Other constituents**
Coumarin,\(^7\) lipids (5–8%),\(^8\) mucilaginous fibre (50%),\(^8\) vitamins (including nicotinic acid) and minerals.
Food Use

Fenugreek is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that fenugreek can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\(^{(G16)}\) In the USA, fenugreek extracts are permitted in foods at concentrations usually below 0.05%. In addition, fenugreek is listed as GRAS (Generally Recognised As Safe) in the USA.
Herbal Use

Fenugreek is stated to possess mucilaginous demulcent, laxative, nutritive, expectorant and orexigenic properties, and has been used topically as an emollient and vulnerary. Traditionally, it has been used in the treatment of anorexia, dyspepsia, gastritis and convalescence, and topically for furunculosis, myalgia, lymphadenitis, gout, wounds and leg ulcers. (G2 G7 G22 G64)
Dosage

Seed
1–6 g or equivalent three times daily. (G49)
Pharmacological Actions

In vitro and animal studies

Hypocholesterolaemic activity has been reported for fenugreek in rats\(^9,\text{G41}\) and alloxan–diabetic dogs.\(^10\) Activity has been attributed to the fibre and saponin fractions, and not to lipid or amino acid fractions.\(^9,\text{10}\) Studies have reported a reduction in cholesterol but not triglyceride concentrations,\(^9\) or in both cholesterol and triglyceride concentrations, but without significant alterations in high–density lipoprotein (HDL) and low–density lipoprotein (LDL) concentrations.\(^\text{10}\)

Hypoglycaemic activity has been observed in rabbits, rats and dogs, and attributed to the defatted seed fraction (DSF),\(^8\) trigonelline, nicotinic acid and coumarin.\(^7,\text{11}\) Oral administration of DSF reduced hyperglycaemia in four alloxan–diabetic dogs, and reduced the response to an oral glucose tolerance test in eight normal dogs, whereas the lipid fraction had no effect on serum glucose and insulin concentrations.\(^8\) The high fibre content (50%) of DSF was thought to contribute to its antidiabetic effect although the initial rate of glucose absorption was not affected.\(^8\) Nicotinic acid and coumarin were reported to be the major hypoglycaemic components of fenugreek seeds, following administration to normal and alloxan–diabetic rats.\(^7\) The hypoglycaemic action exhibited by coumarin was still significant 24 hours post administration.\(^7\) In addition, a slight antidiuretic action was noted for coumarin.\(^7\) Trigonelline inhibited cortisone–induced hyperglycaemia in rabbits if administered (250 mg/kg) concomitantly or two hours before, but not two hours after, cortisone.\(^\text{11}\) In addition, trigonelline exhibited significant hypoglycaemic activity in alloxan–diabetic rats (50 mg/kg), lasting 24 hours.\(^\text{11}\)

A stimulant action on the isolated uterus (guinea–pig), especially during late pregnancy, has been noted for both aqueous and alcoholic extracts.\(^\text{G41}\) An aqueous extract is stated to increase the number of heart beats in the isolated mammalian heart.\(^\text{G41}\)

In vitro antiviral activity against vaccinia virus has been reported for fenugreekine, which also possesses cardiotonic, hypoglycaemic, diuretic, antiphlogistic and antihypertensive properties.\(^2\)

Clinical studies

A transient hypoglycaemic effect was observed in 5 of 10 diabetic patients who received 500 mg oral trigonelline whilst fasting.\(^\text{11}\) Increasing the dose
did not increase this effect, and 500 mg ingested three times a day for five days did not alter the diurnal blood glucose concentration.\textsuperscript{(11)} Hypoglycaemic activity in healthy individuals has been reported for whole seed extracts, with slightly lesser activity exhibited by gum isolate, extracted seeds and cooked seeds.\textsuperscript{(12)} The addition of fenugreek to an oral glucose tolerance test reduced serum glucose and insulin concentrations. Chronic ingestion (21 days) of extracted seeds (25 g seeds daily incorporated into two meals) by non–insulin–dependent diabetics improved plasma glucose and insulin responses (no control group), and reduced 24–hour urinary glucose concentrations.\textsuperscript{(12)} Furthermore, in two diabetic insulin–dependent subjects, daily administration of 25 g fenugreek seed powder reduced fasting plasma–glucose profile, glycosuria and daily insulin requirements (56–20 units) after eight weeks. A significant reduction in serum cholesterol concentrations in diabetic patients was also noted.\textsuperscript{(12)}
No reported side-effects were located for fenugreek. Acute toxicity values (LD$_{50}$) documented for fenugreek alcoholic seed extract are 5 g/kg (rat, oral) and 2 g/kg (rabbit, dermal).\(^{(13)}\) The alcoholic seed extract is reported to be non-irritating and non-sensitising to human skin and non-phototoxic (mice, pigs).\(^{(13)}\) Coumarin is a toxic seed component.\(^{(7)}\) Acute LD$_{50}$ (rat, oral) values per kilogram documented for various seed constituents are 5 g (trigonelline), 8.8 g (nicotinic acid), 7.4 g (nicotinamide) and 0.72 g (coumarin).\(^{(7)}\)
Contra–indications, Warnings

Hypoglycaemic activity has been reported for fenugreek, which may therefore interfere with existing hypoglycaemic therapy. Caution is advisable in patients receiving monoamine oxidase inhibitor (MAOI), hormonal or anticoagulant therapies in view of amine, steroidal saponin and coumarin constituents, respectively, although their clinical significance is unclear. Cardioactivity has been documented in vitro. The absorption of drugs taken concomitantly with fenugreek may be affected (high mucilaginous fibre content).

Pregnancy and lactation

Fenugreek is reputed to be oxytocic(G22) and in vitro uterine stimulant activity has been documented. In view of this, and the documented pharmacologically active components, the use of fenugreek during pregnancy and lactation in doses greatly exceeding those normally encountered in foods is not advisable.
Fenugreek seeds contain a high proportion of mucilaginous fibre, together with various other pharmacologically active compounds including steroidal and amine components. The majority of the traditional uses of fenugreek are probably attributable to the mucilage content. In addition, hypocholesterolaemic and hypoglycaemic actions have been documented for fenugreek in both laboratory animals and humans. The mechanism by which fenugreek exerts these actions is unclear. Proposed theories include a reduction in carbohydrate absorption by the mucilaginous fibre,\(^{12}\) and an effect on cholesterol metabolism, cholesterol absorption and bile acid excretion by the saponin components.\(^{8}\) Toxicity studies indicate fenugreek seeds to be relatively non–toxic, although the presence of pharmacologically active constituents would suggest that excessive ingestion is inadvisable.

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Species (Family)

*Tanacetum parthenium* (L.) Schultz Bip. (Asteraceae/Compositae)
Synonym(s)

Part(s) Used

Leaf, aerial parts
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)
BHP 1996\(^{(G9)}\)
BP 2002\(^{(G71)}\)
ESCOP 1996\(^{(G52)}\)
Martindale 33rd edition\(^{(G67)}\)
Mills and Bone\(^{(G50)}\)
PDR for Herbal Medicines\(^{(G36)}\)
Ph Eur 2004\(^{(G72)}\)
USP26/NF21\(^{(G73)}\)
Legal Category (Licensed Products)

Feverfew is not included in the GSL.\textsuperscript{(G37)}
Constituents

See General References G6 G22 G49 G64.

**Terpenoids**
Sesquiterpene lactones: germacranolides (GE), guaianolides (GU) and eudesmanolides (EU). The structural feature common to all three types is an α-unsaturated γ-lactone moiety, and examples of each type include parthenolide, 3-β-hydroxy-parthenolide, costunolide, 3-β-hydroxycostunolide, artemorin, 8-α-hydroxyestafiatin and chrysanthemomonin (novel dimeric nucleus) (GE); artecanin, chrysanthemolin A (canin) and B (stereoisomers), chrysanthemolide, partholide, two chlorine-containing sesquiterpene lactones (GU); magnolialide, reynosin, santamarine, 1-β-hydroxyarbusculin and 5-β-hydroxyreynosin (EU).'(1–5)

**Volatile oils**
(0.02–0.07%). Various monoterpene and sesquiterpene components (e.g. camphor, borneol, α-pinene derivatives, germacrene, farnesene and their esters).

**Other constituents**
Pyrethrin, flavonoids, tannins (type unspecified) and melatonin.'(6)
Food Use

Feverfew is not generally used in foods.
Feverfew has traditionally been used in the treatment of migraine, tinnitus, vertigo, arthritis, fever, menstrual disorders, difficulty during labour, stomach-ache, toothache and insect bites. Modern use of feverfew is focused on its effects in the prevention and treatment of migraine.
Dosage

Limited information is available regarding the traditional dose of feverfew. The doses that have been recommended for migraine prophylaxis are as follows.

*Leaf (fresh)*

2.5 leaves daily with or after food.

*Leaf (freeze-dried)*

50 mg daily with or after food.

*Aerial parts (dried)*

50–200 mg daily; equivalent to 0.2–0.6 mg parthenolide daily.\(^{G6 \ G52}\)

Clinical trials of feverfew for the prevention of migraine have assessed the effects of, for example, 143 mg of a dried alcoholic extract of feverfew daily (equivalent to 0.5 mg parthenolide),\(^{7}\) and capsules containing powdered feverfew leaf 50 mg daily,\(^{8,9}\) for one to six months.
Pharmacological Actions

In vitro and animal studies

Feverfew extracts have been documented to inhibit platelet aggregation and prostaglandin, thromboxane and leukotriene production, although feverfew has also been reported to have no effect on cyclooxygenase (the mechanism by which non-steroidal anti-inflammatory drugs inhibit prostaglandin production).\(^{(10-12)}\) Instead, feverfew is thought to act by inhibiting the enzyme phospholipase A\(_2\), which facilitates the release of arachidonic acid from the phospholipid cellular membrane.\(^{(11-13)}\) The clinical significance of this action has been questioned.\(^{(14)}\) In addition, *in vitro* experiments have shown that feverfew extracts inhibit the interaction of human platelets with collagen substrates.\(^{(15,16)}\) Feverfew has been shown to inhibit granule secretion in blood platelets and neutrophils, which has been associated with the aetiology of migraine and rheumatoid arthritis, respectively.\(^{(17)}\) Feverfew was also found to inhibit the release of vitamin B\(_{12}\)-binding protein from polymorphonuclear leukocytes, but to be ineffective against platelet and polymorphonucleocyte secretion induced by calcium ionophore A2318.\(^{(17)}\) Sesquiterpene lactone constituents of feverfew containing an α-methylene butyrolactone unit are thought to be responsible for the antisecretory activity.\(^{(18)}\) Their inhibitory effect on platelet aggregation is thought to involve neutralisation of sulfhydryl groups on specific enzymes of proteins that are necessary for platelet aggregation and secretion.\(^{(19)}\) A similar mode of action has been proposed for the inhibitory action of feverfew on polymorphonucleocyte secretion.\(^{(20)}\) In addition, feverfew extracts have been reported to produce a dose-dependent inhibition of anti-IgE-induced histamine release from mast cells.\(^{(21)}\) The authors concluded that the mechanism of action of the feverfew extract was different to that of both cromoglycate and quercetin.

Parthenolide markedly interfered with contractile and relaxant mechanisms in blood vessels.\(^{(G52)}\) An aqueous extract of feverfew administered intravenously significantly inhibited collagen-induced bronchoconstriction in guinea-pigs.\(^{(G52)}\)

The presence of large numbers of lymphocytes and monocytes in the synovium is considered to be of significance in rheumatoid arthritis.\(^{(22)}\) Feverfew extract and parthenolide have been documented to inhibit mitogen-induced proliferation of human peripheral blood mononuclear cells and mitogen-induced prostaglandin E\(_2\) (PGE\(_2\)) production by synovial cells.\(^{(22)}\) The feverfew extract and parthenolide also proved to be cytotoxic to
mitogen–treated peripheral blood mononuclear cells and the authors considered that this cytotoxicity was responsible for the actions observed.\(^{(22)}\) *In vitro* studies using crude feverfew extracts and parthenolide have documented other activities that may contribute to the reported anti–inflammatory effects of feverfew. Pretreatment of human synovial fibroblasts with feverfew extract and with purified parthenolide inhibited cytokine–induced expression of intercellular adhesion molecule 1 (ICAM-1) expression.\(^{(23)}\) A reduction in T cell adhesion to the treated fibroblasts also occurred. In other *in vitro* studies, parthenolide inhibited lipopolysaccharide–induced interleukin-12 (IL-12) production by mouse macrophages in a concentration–dependent manner.\(^{(24)}\) Parthenolide has also been shown to inhibit promoter activity of the inducible nitric oxide synthase gene in a human monocyte cell line, THP-1, in a concentration–dependent manner.\(^{(25)}\) (Excessive nitric oxide production in inflammatory cells is thought to be a causative factor in cellular injury in inflammatory disease.) Anti–inflammatory activity of feverfew has also been attributed to the presence of flavonoids, e.g. santonin.\(^{(26)}\)

Anti–inflammatory properties have also been documented for feverfew extract and parthenolide *in vivo*. Oral administration of feverfew extract (10, 20 and 40 mg/kg) reduced carrageenan–induced oedema in rat paw in a dose–dependent manner.\(^{(27)}\) Intraperitoneal parthenolide (1 and 2 mg/kg) also demonstrated anti–inflammatory effects in this model.

Parthenolide has been documented to have cytotoxic activity in Eagle’s 9KB carcinoma of the nasopharynx cell culture system, the activity being associated with the presence of an α-methylene-γ-lactone moiety in the molecule.\(^{(28)}\) *In vitro*, parthenolide has been shown to inhibit growth of mouse fibrosarcoma (MN-11) and human lymphoma (TK6) cell lines.\(^{(29)}\) The effect appeared to be reversible.

Antinociceptive properties have been reported for feverfew and parthenolide *in vivo*. Oral administration of feverfew extract (10, 20 and 40 mg/kg) and intraperitoneal administration of parthenolide (1 and 2 mg/kg) led to reductions in acetic acid–induced writhing in mice.\(^{(27)}\)

Antimicrobial properties against Gram–positive bacteria, yeasts and filamentous fungi *in vitro* have been documented for parthenolide.\(^{(30)}\) Gram–negative bacteria were not affected.

**Clinical studies**

**Migraine**
Several placebo–controlled clinical trials have assessed the effects of preparations of feverfew in the prevention of migraine.\(^{(7-9,31)}\)

A randomised, double–blind, placebo–controlled trial involved 17 patients who had been successfully controlling their migraine by eating raw feverfew leaves for at least three months.\(^{(8)}\) Patients either continued to receive feverfew (50 mg daily) or were given placebo for six periods of four weeks. The authors reported that the placebo group experienced a significant increase in the frequency and severity of headache. Those given feverfew showed no change. It was suggested that the placebo group was in fact suffering withdrawal symptoms from feverfew and a ‘post-feverfew syndrome’ was described (see Side–effects, Toxicity).

Another study, a randomised double–blind, placebo–controlled, crossover trial involved 72 adults who had experienced migraine for more than two years and who had at least one attack per month.\(^{(31)}\) The only concurrent medication allowed was the oral contraceptive pill. Patients completed a one–month, single–blind, placebo run–in phase, followed by four months‘ administration of placebo/active and four months‘ crossover. It was reported that patients experienced a 24% reduction in the number of attacks during feverfew treatment (one capsule daily; 70–114 mg feverfew equivalent to 2.19 μg parthenolide) although the duration of each individual attack was not significantly affected. Patients allocated to the active and then placebo group did not experience the withdrawal symptoms documented in another study,\(^{(8)}\) although patients involved in the previous study had used feverfew over a longer period of time.

In a randomised, double–blind, placebo–controlled trial, 57 patients received capsules of dried, powdered feverfew leaves (parthenolide 0.2%) 100 mg daily for 60 days (open–label phase), followed by randomisation to feverfew or placebo (ground parsley) for 30 days then crossover to the other arm for 30 days.\(^{(9)}\) There was no washout between crossover. At the end of the open–label phase (i.e. during which all participants received feverfew), there was a significant reduction in pain intensity and symptoms, such as vomiting and sensitivity to light, compared with baseline values \((p < 0.001)\). At the end of the double–blind, crossover phase, it was reported that pain intensity was significantly lower during feverfew administration, compared with placebo administration \((p < 0.01)\).

Thus, these three studies reported beneficial effects for feverfew, as demonstrated by fewer and/or less severe migraine episodes and/or reductions in pain intensity, compared with placebo.\(^{(8,9,31)}\) However, one double–blind, placebo–controlled trial involving 50 feverfew-naïve patients
who experienced migraine attacks at least once a month reported no difference in the number of migraine attacks between placebo recipients and participants who received capsules containing a dried alcoholic extract of feverfew equivalent to 0.5 mg parthenolide daily for nine months.\(^7\) Another randomised, double-blind, placebo-controlled, crossover trial involving 20 patients with migraine assessed the effects of feverfew 100 mg daily for two months on serotonin uptake and platelet activity.\(^{32}\) This trial found no effect for feverfew in the prevention of migraine attacks and also reported that feverfew administration had no effect on the uptake of serotonin by platelets.

The authors of a Cochrane systematic review of six randomised, double-blind, placebo-controlled trials (the five studies mentioned above, plus one another) concluded that although data suggest that feverfew preparations are superior to placebo in preventing migraine, further well-designed clinical trials are required to establish the beneficial effects of feverfew for migraine prophylaxis.\(^{33}\)

**Rheumatoid arthritis**

A double-blind, placebo-controlled, non-crossover trial studying the use of feverfew in rheumatoid arthritis has also been documented.\(^{34}\) Forty-one female patients with inflammatory joint symptoms inadequately controlled by non-steroidal anti-inflammatory drugs were given either one feverfew capsule (70–86 mg equivalent to 2–3 μmol parthenolide) daily, or one placebo capsule, for six weeks. Current non-steroidal therapy was maintained. It was concluded that patients in the trial had experienced no additional benefit from feverfew.\(^{34}\) The authors commented that while concomitant non-steroidal anti-inflammatory drug therapy has been stated to reduce the effectiveness of feverfew, the majority of rheumatoid arthritis sufferers will use feverfew to supplement existing therapy.
Side–effects, Toxicity

Randomised, double–blind, placebo–controlled trials have documented the following adverse effects during feverfew administration, although most effects were also reported (sometimes more frequently) during placebo administration: mouth ulcers (reported more frequently during placebo administration in one study\(^\text{(31)}\)), sore mouth, abdominal pain and indigestion, diarrhoea, flatulence, nausea, dizziness and skin rash.\(^\text{(7,8,31)}\) On balance, adverse effects reported for feverfew are mild and transient, are similar to those reported during placebo administration and occur with a similar frequency.

A ‘post-feverfew syndrome’ has been described on stopping feverfew administration\(^\text{(8)}\) (see Clinical studies) with symptoms such as nervousness, tension headaches, insomnia, stiffness/pain in joints and tiredness.

The onset of side–effects with feverfew is reported to vary, with symptoms becoming apparent within the first week of treatment, or appearing gradually over the first two months.

Sesquiterpene lactones that contain an \(\alpha\)-methyl ene butyrolactone ring are known to cause allergic reactions.\(^\text{(35,G51)}\) Compounds with this structure are present in feverfew and reports of contact dermatitis have been documented.\(^\text{(36–39)}\) No documented allergic reactions following oral ingestion were located.

An LD\(_{50}\) value for feverfew has not been estimated. No adverse effects were reported for rats and guinea–pigs receiving feverfew at doses 100 and 150 times the human daily dose, respectively.\(^\text{(40)}\) No chronic toxicity studies have been reported. However, detailed haematological analysis of 60 feverfew users, some of whom had used feverfew for more than one year, did not show any significant differences when compared with analysis of controls.\(^\text{(40)}\) A human toxicity study has investigated whether the sesquiterpene lactones in feverfew induce chromosomal or other changes in normal human cells of individuals who have taken the herb.\(^\text{(41)}\) The study compared 30 chronic female feverfew users (leaves, tablets or capsules taken daily for more than 11 consecutive months) with matched non–users. The results of lymphocyte cultures established from blood samples taken over a period of several months were stated to indicate that feverfew affects neither the frequency of chromosomal aberrations nor the frequency of sister chromatid exchanges in the circulating peripheral lymphocytes.
Contra-indications, Warnings

Feverfew is contra-indicated in individuals with a known hypersensitivity to other members of the family Compositae (Asteraceae), such as chamomile, ragweed and yarrow. Feverfew should not be ingested by individuals who develop a rash on contact with the plant.

Feverfew should only be considered as a treatment for migraine that has proved unresponsive to conventional forms of medication. Although traditionally recommended as a remedy for rheumatic conditions, self-medication with feverfew should not be undertaken without first consulting a doctor.

Pregnancy and lactation

Feverfew is contra-indicated during pregnancy. It is reputed to be an abortifacient and to affect the menstrual cycle. It is documented to modify menstrual flow, cause abortion in cattle and induce uterine contraction in full-term women. (G30)
Feverfew is characterised by the sesquiterpene lactone constituents, in particular by parthenolide which is thought to be the main active component. *In vitro* studies provide some evidence to support the reputation of feverfew as a herb used to treat migraine and arthritis. Clinical studies have suggested that feverfew may be a useful prophylactic remedy against migraine,\(^{(42, 43)}\) although further research is deemed necessary to establish the benefits.\(^{(33)}\) It has been recommended that feverfew should only be used by sufferers who have proved unresponsive to conventional forms of migraine treatment. Those using feverfew as a remedy for migraine should preferably do so under medical supervision.

Results of a study that investigated the usefulness of feverfew in treating rheumatoid arthritis were less encouraging: feverfew provided no additional benefit when added to existing non-steroidal anti-inflammatory treatment. Feverfew products currently available are unlicensed and vary in their recommended daily doses.\(^{(44)}\) Furthermore, variation between the stated and actual amount of feverfew in commercial products (based on their ability to inhibit platelet secretion) has been reported.\(^{(16)}\)
References


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Species (Family)

*Scrophularia nodosa* L. (Scrophulariaceae)
Synonym(s)
Common Figwort, Scrophularia
Part(s) Used

Herb
Pharmacopoeial and Other Monographs

BHP 1983\(^{(G7)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

Figwort is not included in the GSL.\(^{(G37)}\)
Constituents

See General References G62 G64.

**Amino acids**
Alanine, isoleucine, leucine, lysine, phenylalanine, threonine, tyrosine and valine.\(^{(1)}\)

**Flavonoids**
Diosmetin, diosmin and acacetin rhamnoside.\(^{(2)}\)

**Iridoids**
Aucubin, acetylharpagide, harpagide, harpagoside, isoharpagoside, procumbid and a catalpol glycoside.\(^{(3–5)}\) Figwort is stated to have the same qualitative iridoid composition as devil’s claw, but about half the content of harpagoside.

**Acids**
Various acids including caffeic acid, cinnamic acid, ferulic acid, sinapic acid and vanillic acid, present as both esters and glycosides.\(^{(6,7)}\)
Food Use

Figwort is not used in foods.
Herbal Use

Figwort is stated to act as a dermatological agent and a mild diuretic, and to increase myocardial contraction. Traditionally, it has been used for chronic skin disease, and specifically for eczema, psoriasis and pruritus. (G7 G64)
Dosage

**Dried herb**
2–8 g by infusion.\(^{(G7)}\)

**Liquid extract**
2–8 mL (1 : 1 in 25% alcohol).\(^{(G7)}\)

**Tincture**
2–4 mL (1 : 10 in 45% alcohol).\(^{(G7)}\)
Pharmacological Actions

*In vitro and animal studies*

The iridoid glycosides aucubin and catalpol have been documented to exert a purgative action in mice.\(^8\) Cardioactive properties and anti-inflammatory activity have been claimed for harpagide and the other iridoid constituents (see Devil’s Claw).\(^{G62}\)

*Clinical studies*

None documented. The iridoids are stated to be bitter principles.\(^{G62}\)
Side-effects, Toxicity

None documented.
Contra-indications, Warnings

Figwort should be avoided in ventricular tachycardia.\(^{(G7)}\)

Pregnancy and lactation
The safety of figwort has not been established. In view of the lack of pharmacological and toxicity data, use of figwort during pregnancy and lactation should be avoided.
Pharmaceutical Comment

The chemistry of figwort is well studied and it is stated to be an acceptable substitute for devil’s claw (*Harpagophytum procumbens*) with the same qualitative composition of bitter principles but half the content of harpagoside.\(^{G62}\) Little scientific evidence was located to justify the herbal uses. In view of the lack of toxicity data and possible cardioactive properties, excessive use of figwort should be avoided.
See also General References G7 G31 G36 G37 G62 G64.

Frangula
Species (Family)

*Rhamnus frangula* L. (Rhamnaceae)
Synonym(s)
Alder Buckthorn, *Frangula alnus* Mill.
Part(s) Used

Bark
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)
BHP 1996\(^{(G9)}\)
BP 2002\(^{(G71)}\)
Complete German Commission E (Buckthorn)\(^{(G3)}\)
ESCOP 1997\(^{(G52)}\)
Martindale 33rd edition\(^{(G67)}\)
PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Ph Eur 2004\(^{(G72)}\)
Legal Category (Licensed Products)

GSL(G37)
Constituents

**Anthraquinones**
3–7%. Frangulosides as major components including frangulin A and B (emodin glycosides) and glucofrangulin A and B (emodin diglycosides); emodin derivatives including emodin dianthrone and its monorhamnoside, palmidin C (see Rhubarb) and its monorhamnoside, emodin glycoside; also glycosides of chrysophanol and physcione, and various free aglycones.

**Other constituents**
Flavonoids and tannins.
Food Use

Frangula is listed by the Council of Europe as a natural source of food flavouring (category N4). While this category recognises the use of frangula as a flavouring agent, it indicates that there is insufficient information available to classify it further into categories N1, N2, or N3. (G16)
Herbal Use

Frangula is stated to possess mild purgative properties and has been used traditionally for constipation.\(^{(G2\ G6\ G7\ G8\ G64)}\)

The Committee on Proprietary Medicinal Products (CPMP) has adopted a core SPC (Summary of Product Characteristics) for frangula. The core SPC includes indications for short-term use of frangula in cases of occasional constipation.\(^{(1)}\)
Dosage

*Dried bark*
0.5–2.5 g.(G6)

*Liquid extract*
2–5 mL (1 : 1 in 25% alcohol) three times daily.(G7)
Pharmacological Actions

The pharmacological activity of frangula can be attributed to the anthraquinone glycoside constituents. The laxative action of these compounds is well recognised (see Senna).
See Senna for side-effects and toxicity associated with anthraquinones.

The CPMP core SPC for frangula includes the following information.\(^1\) There are no studies on single dose toxicity, on repeated dose toxicity, on reproductive toxicity or on carcinogenicity. Different frangula extracts were shown to be genotoxic in several \textit{in vitro} systems (bacterial mutation, chromosomal aberration and DNA-repair in mammalian cells). No increases in mutations were observed in a gene mutation assay with mammalian cells. For emodin, the main laxative principle of frangula, signs of a genotoxic potential were observed in several systems (bacteria and mammalian cells \textit{in vitro}). Other anthraquinone constituents also gave positive results in limited experiments.
Contra-indications, Warnings

See Senna for contra-indications and warnings associated with anthraquinones.

The CPMP core SPC for frangula states the following contraindications and warnings.(1)

**Contra-indications**

Not to be used in cases of intestinal obstruction and stenosis, atony, inflammatory colon diseases (e.g. Crohn’s disease, ulcerative colitis), appendicitis, abdominal pain of unknown origin, severe dehydration states with water and electrolyte depletion.

**Precautions**

As with all laxatives, frangula bark should not be given when any undiagnosed acute or persistent abdominal symptoms are present. If laxatives are needed every day, the cause of the constipation should be investigated. Long-term use of laxatives should be avoided. Use for more than two weeks requires medical supervision. Chronic use may cause pigmentation of the colon (pseudomelanosis coli) which is harmless and reversible after drug discontinuation.

Abuse, with diarrhoea and consequent fluid and electrolyte losses, may cause: dependence, with possible need for increased dosages, disturbance of the water and electrolyte (mainly hypokalaemia) balance, an atonic colon with impaired function. Intake of anthranoid containing laxatives exceeding short-term use may result in an aggravation of constipation.

Hypokalaemia can result in cardiac and neuromuscular dysfunction, especially if cardiac glycosides, diuretics or corticosteroids are taken. Chronic use may result in albuminuria and haematuria.

In chronic constipation, stimulant laxatives are not an acceptable alternative to a changed diet.

**Interaction with other medicaments and other forms of interaction.**

Hypokalaemia (resulting from long-term laxative abuse) potentiates the action of cardiac glycosides and interacts with antiarrhythmic drugs and drugs which induce reversion to sinus rhythm (e.g. quinidine). Concomitant use with other drugs inducing hypokalaemia (e.g. thiazide diuretics, adrenocorticosteroids and liquorice root) may enhance electrolyte imbalance.

**Pregnancy and lactation**
Pregnancy and lactation

The use of stimulant laxatives, particularly unstandardised preparations, is not generally recommended during pregnancy (see Senna).

The CPMP core SPC for frangula includes the following information on use during pregnancy and lactation.

**Pregnancy** Frangula is not recommended during pregnancy.\(^1\)

There are no reports of undesirable or damaging effects during pregnancy and on the foetus when used at the recommended dosage schedule. However, experimental data concerning a genotoxic risk of several anthranoids, e.g. emodine and physcione, and frangula extract are not counterbalanced by sufficient studies to eliminate a possible risk.\(^1\)

**Lactation** Frangula is not recommended during breast feeding, as there are insufficient data on the excretion of its metabolites in breast milk. Excretion of the active principles of frangula in breast milk has not been investigated. However, small amounts of active metabolites (e.g. rhein) from other anthranoids are known to be excreted in breast milk. A laxative effect in breastfed babies has not been reported.\(^1\)
Pharmaceutical Comment

The chemistry of frangula is characterised by the anthraquinone glycoside constituents. The laxative action of these compounds is well recognised and supports the herbal use of frangula as a laxative. The use of non-standardised anthraquinone-containing preparations should be avoided, since their pharmacological effect will be variable and unpredictable. In particular, the use of products containing combinations of anthraquinone laxatives should be avoided.
References


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Fucus
Species (Family)

*Fucus vesiculosus* L. and other *Fucus* species (Fucaceae)
Synonym(s)
Black Tang, Bladderwrack, Kelp, Kelpware, Rockweed, Seawrack

Brown seaweeds refer to species of *Fucus, Ascophyllum, Laminaria* and *Macrocystis*. ‘Kelps’ refer to species of *Laminaria* and *Macrocystis*, although kelp is often used in reference to species of *Fucus*. 
Part(s) Used

Thallus (whole plant)
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

BP 2002\(^{(G71)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)

Ph Eur 2004\(^{(G72)}\)
Legal Category (Licensed Products)

GSL$^{(G37)}$
Constituents
See General References G2 G6 G64.

Carbohydrates
Polysaccharides: alginic acid (algin) as the major component; fucoidan and laminarin (sulfated polysaccharide esters).\(^{(1)}\)

Iodine
Content of various *Laminaria* species has been reported as 0.07–0.76% of dry weight.\(^{(2)}\)

Other constituents
Various vitamins and minerals, particularly ascorbic acid (vitamin C) (0.013–0.077% of fresh material).\(^{(2)}\)
Food Use

Seaweeds are commonly included in the diet of certain populations. The gelling properties of alginic acid, the major polysaccharide in brown seaweeds, including fucus, are extensively utilised in the dairy and baking industries to improve texture, body and smoothness of products.\(^{(1)}\) Fucus is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that fucus can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\(^{(G16)}\)
Herbal Use

Fucus is stated to possess antihypothyroid, anti-obesic and antirheumatic properties. Traditionally, it has been used for lymphadenoid goitre, myxoedema, obesity, arthritis and rheumatism.\(^G2\ G6\ G7\ G8\ G64\)
Dosage

*Dried thallus*
5–10 g or infusion three times daily.\(^{(G6 \ G7)}\)

*Liquid extract*
4–8 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G6 \ G7)}\)
Pharmacological Actions

There is a paucity of information documented specifically for *Fucus vesiculosus*, although pharmacological activities are recognised for individual constituents and other brown seaweed species.

Alginic acid is a hydrophilic colloidal substance that swells to approximately 25–35 times its original bulk in an alkaline environment and as such exerts a bulk laxative action. It is stated to compare favourably with the carboxylic type of cation exchange resins. The colloidal properties of alginates have been utilised in wound dressings and skin grafts.

Anticoagulant properties have been documented for brown seaweeds. The glucose polymer laminarin has been identified as the anticoagulant principle in a Laminaria species. A fucoidan fraction has been isolated from *Fucus vesiculosus* with 40–50% blood anticoagulant activity of heparin.

The iodine content of seaweeds is well recognised. The low incidence of goitre amongst maritime people has been attributed to the inclusion of seaweeds in their diet. Similarly, the traditional use of *Fucus vesiculosus* in ‘slimming teas’ is thought to be attributable to the effect of iodine on hypothyroidism.

Extracts of various brown seaweeds including *Ascophyllum nodosum* and *Fucus vesiculosus* have been reported to exhibit a high *in vitro* inhibitory activity towards mammalian digestive enzymes (α-amylase, trypsin and lipase) isolated from the porcine pancreas. Activity was attributed to high molecular weight (30 000–100 000) polyphenols.

Inhibitory effects of laminarin sulfate on lipid aemia and atherosclerosis (*in vivo*, rabbit) have been partially attributed to the *in vitro* inhibition of lipid synthesis observed in cultured chick aortic cells.

Hypotensive activity observed in rats intravenously administered extracts of commercial seaweed (*Laminaria* species) preparations has been attributed to their histamine content. However, histamine concentrations varied considerably between preparations, and authentic specimens of the *Laminaria* species were devoid of histamine.

Kelp extracts have antiviral activity and laminarin is reported to have exhibited some tumour–inhibiting actions.
Side-effects, Toxicity

Hyperthyroidism has been associated with the ingestion of kelp and is attributable to the iodine content in the plant.\(^{(10,11)}\) Typical symptoms of hyperthyroidism (weight loss, sweating, fatigue, frequent soft stools) were exhibited by a 72-year-old woman following ingestion of a commercial kelp product for six months.\(^{(10)}\) Laboratory tests confirmed the hyperthyroidism although no pre-existing evidence of thyroid disease was found and the condition resolved in six months following discontinuation of the tablets. Analysis of the kelp tablets reported an iodine content of 0.7 mg/tablet representing a daily intake of 2.8–4.2 mg iodine.\(^{(10)}\) Clinically evident hyperthyroidism developed in an otherwise healthy woman following the daily ingestion of six 200-mg kelp tablets.\(^{(11)}\) Symptoms gradually resolved on cessation of therapy.

The association between halogen salts and acneiform eruptions is well established.\(^{(12)}\) Ingestion of kelp products has been associated with the worsening of pre-existing acne and the development of acneiform eruptions, which improved following withdrawal of the tablets.\(^{(12)}\)

The ability of marine plants to accumulate heavy metals and other toxic elements is recognised, and the uptake of various radioactive compounds by seaweeds has been reported.\(^{(3,13,14)}\) Fifteen samples of kelp-containing dietary supplements have been analysed for their iodine and arsenic contents.\(^{(15)}\) The levels of arsenic were low in all but one product. The iodine levels varied widely, even between different samples of the same product, and in some products the iodine levels were high in relation to safe daily intake.

Brown algae (\textit{Ascophyllum nodosum} and \textit{Fucus vesiculosus}) have been found to be capable of synthesising volatile halogenated organic compounds (VHOCs).\(^{(16)}\) VHOCs are considered to be troublesome pollutants because land plants and animals have difficulty in degrading the compounds which consequently persist in terrestrial ecosystems.\(^{(16)}\) VHOCs released into the seawater predominantly contain bromine with iodine-containing compounds showing a slower rate of turnover.\(^{(16)}\) Concentration of iron by brown seaweeds has been attributed to fucoidan, and alginic acid exhibits a high specificity for the binding of strontium.\(^{(13)}\) Elevated urinary arsenic concentrations (138 and 293 μg/24 hour) in two female patients have been associated with the ingestion of kelp tablets. Subsequent analysis of the arsenic content of various kelp preparations revealed concentrations ranging from 16 to 58 μg/g product.\(^{(17,18)}\) The botanical source of the kelp in the products was not stated.\(^{(18)}\)
Ascophyllum nodosum is commonly added to animal foodstuffs as a source of vitamin and minerals, with beneficial results reported for dairy cattle, sheep, pigs and poultry.\(^{(13)}\) Feeding studies using A. nodosum have highlighted an atypical toxic response for rabbits compared with that of rats and pigs.\(^{(13,19)}\) Addition of A. nodosum to the diet of rabbits (at 5–10\%) caused a severe drop in haemoglobin content, serum iron concentrations and packed cell volume, leading to weight loss and death in two-thirds of the animals.\(^{(13)}\) No differences in renal and liver function, and in lipid metabolism were found between test and control animals.\(^{(13)}\) Similar, but much milder, toxicity has also been observed in rabbits fed Fucus serratus.\(^{(19)}\) Subsequent studies incorporating A. nodosum into the feed of rats and pigs failed to demonstrate the toxic effects observed in rabbits.\(^{(19)}\) The toxic components in A. nodosum have been reported to be non-extractable with chloroform, ethanol, water and 20\% sodium carbonate solution, remaining in the insoluble residue.\(^{(19)}\)
Contra-indications, Warnings

The iodine content in kelp may cause hyper- or hypothyroidism and may interfere with existing treatment for abnormal thyroid function. In view of this, ingestion of kelp preparations by children is inadvisable. The iodine content in kelp has also been associated with acneiform eruptions and aggravation of pre-existing acne. In general, brown seaweeds are known to concentrate various heavy metals and other toxic elements. Elevated urinary arsenic concentrations have been traced to the ingestion of kelp tablets. Prolonged ingestion of kelp may reduce gastrointestinal iron absorption (binding properties of fucoidan), resulting in a slow reduction in haemoglobin, packed cell volume and serum iron concentrations. Prolonged ingestion may also affect absorption of sodium and potassium ions (alginic acid) and cause diarrhoea.

Pregnancy and lactation

The safe use of kelp products during pregnancy and lactation has not been established. In view of the potential actions on the thyroid gland and possible contamination with toxic elements, the use of kelp should be avoided.
Kelp is a generic term that strictly speaking refers to *Laminaria* and *Macrocystis* species of brown seaweeds, although in practice it may be used in reference to other species of brown algae including *Nereocystis* and *Fucus*. The species *Fucus vesiculosus* is reported to be commonly used in the preparation of kelp products.\(^{(G60)}\) The principal constituents of seaweeds are polysaccharides. For brown seaweeds the major polysaccharide is alginic acid (algin). Fucoidan, present in all brown algae, is thought to refer to a number of related polysaccharide esters whose main sugar component is fucose. The traditional uses of kelp in obesity and goitre are presumably attributable to the iodine content, although the self-diagnosis and treatment of these conditions with a herbal remedy is not suitable. There have been no documented studies supporting the traditional use of kelp in rheumatic conditions. In view of the iodine content and potential accumulation of toxic elements, excessive ingestion of kelp is inadvisable. Doubt over the quality of commercial seaweed preparations has been reported.\(^{(10)}\)
References


18. Walkiw O, Douglas DE. Health food supplements prepared from kelp – a


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Fumitory
Species (Family)

*Fumaria officinalis* L. (Fumariaceae)
Synonym(s)
Fumitory
Part(s) Used

Herb
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

Complete German Commission E\(^{(G3)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents

See General References G2 G6 G40 G64.

Alkaloids
Isoquinoline–type. Protopines including protopine (fumarine) as the major alkaloid and cryptopine,\(^1,2\) protoberberines including aurotensine, stylopine, sinactine and \(N\)-methylsinactine,\(^3\) spirobenzylisoquinolines including fumaritine, fumaricine and fumariline,\(^4,5\) benzophenanthridines including sanguinarine,\(^6\) and indenobenzaz-epines including fumaritridine and fumaritrine.\(^6,7\)

Flavonoids
Glycosides of quercetin including isoquercitrin, rutin and quercetrin–3,7–diglucoside–3–arabinoglucoside.\(^8,9\)

Acids
Chlorogenic, caffeic and fumaric acids.\(^8\)

Other constituents
Bitter principles, mucilage and resin.
Food Use

Fumitory is listed by the Council of Europe as a natural source of food flavouring (category N3). This category indicates that fumitory can be added to foodstuffs in the traditionally accepted manner, although there is insufficient information available for an adequate assessment of potential toxicity.\textsuperscript{(G16)}
Herbal Use

Fumitory is stated to possess weak diuretic and laxative properties and to act as a cholagogue. Traditionally, it has been used to treat cutaneous eruptions, conjunctivitis (as an eye lotion) and, specifically, chronic eczema. (G2 G6 G7 G8 G64)
**Dosage**

*Herb*
2–4 g or by infusion three times daily.\(^{(G6 \ G7)}\)

*Liquid extract*
2–4 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G6 \ G7)}\)

*Tincture*
1–4 mL (1 : 5 in 45% alcohol) three times daily.\(^{(G6 \ G7)}\)
Pharmacological Actions

*In vitro* and animal studies

The herb had no effect on normal cholesresis but it modified bile flow which was artificially increased or decreased.\(^{10}\) Antispasmodic activity on smooth muscle has been reported.\(^{11}\) Extracts inhibited formation of gall-bladder calculi in animals.\(^{12}\) The major alkaloid protopine has antihistaminic,\(^{13}\) hypotensive, bradycardic and sedative activities in small doses,\(^{14}\) whereas larger doses cause excitation and convulsions.\(^{14}\) Bactericidal activity against the Gram-positive organisms *Bacillus anthracis* and *Staphylococcus* have been reported.\(^{14}\)

Clinical studies

Clinical studies involving 105 patients with biliary disorders claimed favourable results.\(^{15}\)
Side–effects, Toxicity

No reported side–effects or documented toxicity studies were located, although possible adverse effects include raised intraocular pressure and oedema.\(^{(16)}\)
Contra–indications, Warnings

Hypotensive actions have been documented in animal studies.

Pregnancy and lactation
The safety of fumitory during pregnancy and lactation has not been established. In view of lack of pharmacological and toxicity data, the use of fumitory during pregnancy and lactation should be avoided.
Pharmaceutical Comment

Fumitory is characterised by isoquinoline alkaloids which represent the principal active ingredients. Animal studies support some of the traditional uses, but it should not be used in home-made ophthalmic preparations. In view of the active constituents and the lack of safety data, excessive ingestion of fumitory should be avoided.
References

See also General References G2 G3 G6 G9 G16 G31 G36 G37 G43 G56 G64.


Garlic
Species (Family)

*Allium sativum* L. (Amaryllidaceae/Liliaceae)
Synonym(s)
Ajo, Allium
Part(s) Used

Bulb (clove)
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)
BP 2002\(^{(G71)}\)
BPC 1949\(^{(G11)}\)
Complete German Commission E\(^{(G3)}\)
ESCOP 1997\(^{(G52)}\)
Martindale 33rd edition\(^{(G67)}\)
Mills and Bone\(^{(G50)}\)
PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Ph Eur 2004\(^{(G72)}\)
USP26/NF21\(^{(G73)}\)
WHO year volume 1\(^{(G63)}\)
Legal Category (Licensed Products)

GSL\textsuperscript{(G37)}
Constituents
See References 1–3, and General References G6 G41 G52 G56 G64.

Enzymes
Allinase, peroxidases, myrosinase and others (e.g. catalases, superoxide dismutases, arginases, lipases).\(^{(2,3)}\)

Volatile oils
0.1–0.36%. Sulfur–containing compounds including alliin, compounds produced enzymatically from alliin including allicin (diallyl thiosulfinate), allylpropyl disulfide, diallyl disulfide, diallyl trisulfide; ajoene and vinylthiines (secondary products of alliin produced non–enzymatically from allicin); S-allylmercaptocysteine (ASSC) and S-methylmercaptocysteine (MSSC); terpenes include citral, geraniol, linalool, α- and β-phellandrene.

Other constituents
Proteins (e.g. glutamyl peptides), amino acids (e.g. arginine, glutamic acid, asparagic acid, methionine, threonine), minerals, vitamins, trace elements, lipids, prostaglandins (A\(_2\), D\(_2\), E\(_2\), F\(_{1\alpha}\), F\(_2\)).\(^{(2,4)}\)

Allicin and other sulfur–containing compounds are formed from alliin by the enzyme alliinase when garlic is crushed or chopped. (Alliin and alliinase are separated while the cells of a garlic bulb are intact, but crushing and chopping damage the cells of the bulb, allowing alliin and alliinase to come into contact with each other.\(^{(G56)}\)) It is considered that 1 mg alliin is equivalent to 0.45 mg allicin.\(^{(G52)}\) Commercial garlic preparations are often standardised on content of sulfur–containing constituents, particularly to alliin, or on allicin yield.

Garlic powder contains not less than 0.45% allicin calculated with reference to the dried drug.\(^{(G28)}\)
Garlic is used extensively as a food and as an ingredient in foods. It is listed by the Council of Europe as a natural source of food flavouring (category N1). This category indicates that there are no restrictions on the use of garlic in foods. (G16) In the USA, garlic is listed as GRAS (Generally Recognised As Safe). (G41)
Herbal Use

Garlic is stated to possess diaphoretic, expectorant, antispasmodic, antiseptic, bacteriostatic, antiviral, hypotensive and anthelmintic properties, and to be a promoter of leukocytosis. Traditionally, it has been used to treat chronic bronchitis, respiratory catarrh, recurrent colds, whooping cough, bronchitic asthma, influenza and chronic bronchitis. Modern use of garlic and garlic preparations is focused on their reputed antihypertensive, anti-atherogenic, antithrombotic, antimicrobial, fibrinolytic, cancer preventive and lipid-lowering effects.
**Dosage**

**Dried bulb**
2–4 g three times daily;\(^{(G6)}\) fresh garlic 4 g daily.\(^{(G3)}\)

**Tincture**
2–4 mL (1 : 5 in 45% alcohol) three times daily.\(^{(G6)}\)

**Oil**
0.03–0.12 mL three times daily.\(^{(G6)}\)

**Juice of Garlic**
(BPC 1949) 2–4 mL.\(^{(G11)}\)

**Syrup of Garlic**
(BPC 1949) 2–8 mL.\(^{(G11)}\)

Clinical trials assessing the effects of garlic powder tablets on various parameters, including total serum cholesterol concentrations, triglyceride concentrations, blood pressure, platelet aggregation, vascular resistance, fibrinolysis and measures of peripheral arterial occlusive disease, have generally involved the administration of doses of 600–900 mg daily for 4–24 weeks.\(^{(G56)}\) For prophylaxis of atherosclerosis, ESCOP (European Scientific Co-operative on Phytotherapy) states a dosage 0.5–1.0 g dried garlic powder daily (approximately equivalent to alliin 6–10 mg and allicin 3–5 mg).\(^{(G52)}\)
Pharmacological Actions

**In vitro and animal studies**

Many pharmacological properties have been documented for garlic and its constituents *in vitro* and *in vivo* (animals), including antihypertensive, lipid-lowering, anti-atherogenic, antithrombotic, fibrinolytic, antioxidant, anticarcinogenic, antitumorogenic, immunomodulatory and antimicrobial activities. The pharmacological properties of garlic are attributed mainly to its sulfur-containing compounds. An extensive review of the pharmacological properties of garlic and its constituents is beyond the scope of this monograph, although several studies are described in brief below. The pharmacological activities of garlic and its constituents have been summarised in many reviews.\(^{(3,5–21,G5 G56)}\)

**Pharmacokinetics**

The available literature on the metabolism and pharmacokinetics of the constituents of garlic in animals has been reviewed.\(^{(3,G18)}\)

In an *ex vivo* study, allicin showed a marked first-pass clearance effect in isolated perfused rat liver.\(^{(3)}\) In rats, alliin and allicin were administered orally at doses of 8 mg/kg.\(^{(22)}\) Absorption of alliin and allicin was complete after 10 minutes and 30–60 minutes, respectively. The mean total urinary and faecal excretion of allicin after 72 hours was 85.5% of the dose. No unchanged alliin or allicin was detected in urine, suggesting rapid and extensive metabolism of these constituents.\(^{(3)}\) Pharmacokinetic studies of the garlic constituent S-allyl-L-cysteine administered orally to rats at doses of 12.5, 25 and 50 mg/kg have reported bioavailability of 64%, 77% and 98%, respectively.\(^{(23)}\) Peak plasma concentrations of S-allyl-L-cysteine occurred at one hour, and the half-life of S-allyl-L-cysteine was 2.33 hours following oral administration of 50 mg/kg to rats.

**Anti-atherosclerotic and cholesterol- and lipid-lowering effects**

The effects of garlic and its constituents on cholesterol biosynthesis *in vitro* and in animal models of hypercholesterolaemia are well documented.\(^{(3)}\)

Several *in vitro* studies have shown that garlic and its sulfur-containing constituents inhibit cholesterol biosynthesis in cultured hepatocytes.\(^{(24–28)}\) In other *in vitro* studies, garlic extracts were shown to inhibit fatty acid and triglyceride synthesis.\(^{(29,30)}\)

The step(s) in the cholesterol biosynthetic pathway inhibited by garlic, and the constituents of garlic causing inhibition have not been definitively
established. Several mechanisms of action for the effects of garlic constituents on cholesterol and lipid synthesis have been proposed, including inhibition of hydroxy methylglutaryl-CoA (HMG-CoA) reductase activity and other enzymes, such as lanosterol–14–demethylase, involved in cholesterol biosynthesis.\(^3\) Other proposed mechanisms include reduction in triacylglycerol biosynthesis via a reduction in tissue concentrations of NADPH, increase in hydrolysis of triacylglycerols via increased lipase activity and inactivation of enzymes involved in lipid synthesis via an interaction with enzyme thiol groups.\(^6,8,31\) More recently, fresh garlic extract and the constituents \(S\)-allylcysteine, diallyl trisulfide and diallyl disulfide were shown to inhibit human squalene mono–oxygen ase, an enzyme catalysing a step in cholesterol biosynthesis.\(^32\) Another \textit{in vitro} study reported that \(S\)-allylcysteine, \(S\)-propylcysteine and \(S\)-ethylcysteine inhibit triglyceride biosynthesis in part by decreasing \textit{de novo} fatty acid synthesis via inhibition of fatty acid synthase.\(^30\)

The anti–atherogenic, anti–atherosclerotic and cholesterol- and lipid–lowering effects of garlic and its constituents have been documented in several animal models (e.g. rabbits, rats, chickens, pigs) of atherosclerosis, hypercholesterolaemia and hyper lipidaemia.\(^3\) For example, a reduction in both blood and tissue lipid concentrations in hypercholesterolaemic animals fed a diet supplemented with dried garlic powder, garlic oil, or allicin has been documented.\(^6,33\) Garlic has also been reported to reduce hepatic triglyceride and cholesterol concentrations in rats, and to reduce aortic lipid deposition and atheromatous lesions in rabbits fed a high–fat diet.\(^6\) Several studies have reported hypolipidaemic effects for garlic oil following administration to rats and rabbits fed a fat–rich diet to induce hyperlipidaemia.\(^3\) Administration of aged garlic extract to rabbits fed a 1% cholesterol–enriched diet for six weeks reduced the surface area of the thoracic aorta covered by fatty streaks (atherosclerosis) and significantly reduced aortic arch cholesterol, although plasma cholesterol concentrations were not reduced.\(^34\) Allicin administration has been reported to reduce significantly the formation of fatty streaks in mice fed a cholesterol–rich diet, compared with control mice (no allicin treatment).\(^35\)

The cholesterol–lowering effect of garlic is thought to be dose–related; proposed mechanisms of action include inhibition of lipid synthesis and increased excretion of neutral and acidic sterols.\(^6,8\) An \textit{in vitro} study reported that aged garlic extract may exert its anti–atherogenic effects via inhibition of smooth muscle proliferation and phenotypic change, and by an effect on lipid accumulation in the artery wall.\(^34,\)

\textbf{Antithrombotic and fibrinolytic activities}
Antithrombotic activity is well documented for garlic in both *in vitro* and *in vivo* (animal) studies. Antithrombotic effects have been documented for fresh garlic, garlic powder and garlic oils.

Increased serum fibrinogen concentrations together with a decrease in blood coagulation time and fibrinolytic activity are associated with a high–fat diet and enhance thrombosis. Garlic has been shown to have a beneficial effect on all of these parameters. Garlic has been shown to inhibit platelet aggregation caused by several inducers such as ADP, collagen, arachidonic acid, adrenaline (epinephrine) and calcium ionophore A23187. Antiplatelet activity has been documented for garlic in *in vitro* studies using human platelets.

Several mechanisms have been proposed by which garlic is thought to exert an anti–aggregatory action. These include inhibition of thromboxane synthesis via cyclooxygenase and lipoxygenase inhibition, inhibition of membrane phospholipase activity and incorporation of arachidonic acid into platelet membrane phospholipids, intraplatelet mobilisation of calcium uptake and inhibition of calcium uptake into platelets. Garlic oil has been reported to reduce artificial surface adhesion of platelets *in vitro*. Certain garlic constituents also affect processes preceding platelet aggregation, such as activation of platelets.

Garlic is thought to contain more than one inhibitor of platelet aggregation and release; allicin is considered to be the major inhibitor. Other studies have investigated the role of ajoene (a secondary degradation product of alliin) as an inhibitor of platelet aggregation and release. Ajoene inhibits platelet aggregation caused by various inducers. Its action is noted to be dose–dependent and reversible both *in vitro* and *in vivo*. It has been suggested that this latter feature may be of clinical significance in instances where a rapid inhibition of platelet aggregation is required with subsequent reversal, such as chronic haemodialysis and coronary bypass surgery. It has been proposed that ajoene exerts its anti–aggregatory effect by altering the platelet membrane via an interaction with sulfhydryl groups. The inhibitory action of ajoene on granule release from platelets is thought to involve alteration of the microviscosity in the inner part of the plasma membrane. Ajoene is reported to synergistically potentiate the anti–aggregatory action of prostacyclin, forskolin, indometacin and dipyridamole, and to potentiate the inhibitory action of prostaglandin I$_2$ (PGI$_2$) on platelet aggregation. Approximately 96% inhibition of prostaglandin synthetase and 100% inhibition of lipoxygenase has been described for ajoene *in vitro*. Structure–activity investigations suggested
that an allylic structure in the open disulfide ring is required for activity.\(^{(40)}\)

**Antioxidant effects**

Antioxidant properties have been documented for garlic in vitro and in vivo (animals).\(^{(3)}\) Garlic constituents inhibit the formation of free radicals, support endogenous radical-scavenging mechanisms, enhance cellular antioxidant enzymes (e.g. superoxide dismutase, catalase, glutathione peroxidase), protect low-density lipoprotein from oxidation by free radicals, and inhibit the activation of the oxidant-induced transcription factor nuclear factor kappa B (NF-κB).\(^{(3,18)}\)

Garlic powder was reported to inhibit the production of superoxide by phorbol ester-activated human granulocytes in vitro (IC\(_{50}\) 390 μg/mL),\(^{(49)}\) whereas alliin did not inhibit superoxide production in this model. It was suggested that allicin may be the constituent of garlic responsible for the observed oxygen-radical scavenging properties. In vitro, aged garlic extract and S-allylcysteine inhibited low-density lipoprotein oxidation and protected pulmonary artery endothelial cells against injury induced by oxidised low-density lipoprotein.\(^{(50)}\) In subsequent studies using bovine pulmonary artery endothelial cells and murine macrophages, it was shown that aged garlic extract inhibited oxidised low-density lipoprotein–induced release of peroxides.\(^{(51)}\) In vivo studies have reported reductions in liver lipid peroxidation and inhibition of ethanol-induced mitochondrial lipid peroxidation in rats fed garlic oil.\(^{(3)}\)

The antioxidant properties are of interest in relation to the antiarteriosclerotic, antihepatotoxic and anticancer effects of garlic and its constituents. For example, oxidation of low-density lipoprotein plays an important role in the initiation and progression of atherosclerosis.\(^{(50)}\)

**Antihypertensive effects**

Several studies involving animal models (e.g. dogs, rats) of hypertension have reported hypotensive effects of garlic preparations.\(^{(3)}\) A hypotensive effect in dogs administered garlic extract has been documented; prior administration of antagonists to known endogenous hypotensive substances such as histamine, acetylcholine, serotonin and kinins did not affect the hypotensive effect.\(^{(52)}\) Spontaneously hypertensive rats fed standardised dry garlic powder 1 mg/kg for nine months exhibited lower blood pressure than control rats (150 versus 205 mmHg, respectively).\(^{(53)}\) By contrast, an ethanolic extract of garlic (1–2 g and 4–8 g daily) fed to spontaneously hypertensive rats did not lead to a reduction in blood pressure.\(^{(54)}\)

**Anticarcinogenic and antitumorigenic activities**
Many in vitro and animal studies have documented anticancer activities of garlic and its constituents.\(^{(3,14,15)}\) These studies indicate that allicin, allicin–derived compounds and other compounds unrelated to allicin contribute to the anticancer effects of garlic.

In several animal models, garlic has been shown to inhibit carcinogenesis and to protect against the development of experimentally induced tumours.\(^{(3,14,15)}\) For example, aged garlic extract significantly inhibited the growth of Sarcoma–180 and LL/2 lung carcinoma cells transplanted into mice.\(^{(55)}\) Garlic powder and its constituents S-allylcysteine and diallyl disulfide inhibited \(N\)-methyl-\(N\)-nitrosourea–induced mammary carcinogenesis in rats,\(^{(56)}\) and fresh garlic (250 mg/kg orally, three times weekly) suppressed 4–nitroquinoline–1–oxide–induced carcinogenesis in rat tongue.\(^{(57)}\) Inhibition of benzo[a]pyrene–induced neoplasia of the fore stomach and lung in female mice has been documented for four allyl group–containing derivatives in garlic.\(^{(58)}\) Structure–activity requirements underlined the importance of the unsaturated allyl groups for activity. Saturated analogues containing propyl instead of allyl groups were devoid of activity.

In vitro studies using human tumour cell lines have reported that garlic powder and garlic extract inhibited the growth of a human lymphatic leukaemia cell line (CCRF CEM) in a concentration–dependent manner at concentrations down to 30 μg/mL.\(^{(59)}\) Also, a combination of garlic extract and garlic powder inhibited the growth of human hepatoma (HepG02) cells and human colorectal carcinoma (Caco2) cells in a concentration–dependent manner, although no activity was observed on these tumour cell lines with garlic extract or powder alone.\(^{(59)}\) Synthetic diallyl disulfide inhibited tumour cell growth in four human breast cancer cell lines.\(^{(60)}\) Growth inhibition occurred regardless of oestrogen receptor status.

Evidence indicates that there are several mechanisms by which garlic and its constituents may exert anticancer effects, such as inhibition of carcinogen formation, modulation of carcinogen metabolism, inhibition of mutagenesis and genotoxicity, increased apoptosis and inhibition of angiogenesis.\(^{(20)}\)

Garlic has been shown to inhibit the synthesis of \(N\)-nitroso compounds (there is a view that \(N\)-nitroso compounds are possible carcinogens for humans).\(^{(14)}\) Also, in rats pretreated with dimethylbenz[a]anthracene (DMBA) and fed a diet supplemented with garlic powder, the occurrence of DNA adducts in mammary tissue was significantly inhibited, compared with control.\(^{(61)}\) (DMBA initiates and promotes cancer, and alkylation of DNA is thought to be an important step in carcinogenesis.) Dietary garlic has also been shown to
Another possible explanation is that garlic constituents may modify drug-metabolising enzymes, which would have the effect of altering the bioactivation of carcinogens.\(^{(14)}\) Glutathione-S-transferase activity has been shown to increase in rat and mouse tissues after administration of garlic powder or its sulfur-containing constituents.\(^{(3,14,15)}\) Certain garlic constituents, e.g. diallyl sulfide, may depress the activity of some hepatic cytochrome P450 (CYP) enzymes, such as CYP2E1 and CYP2A6,\(^{(3,14,15,63)}\) although other studies have shown that garlic constituents induce the activity of other CYP enzymes.\(^{(3,17)}\) In rats, the antimitogenic properties of sulfur-containing compounds from garlic, e.g. diallyl disulfide and diallyl sulfide, against the carcinogens styrene oxide and 4-nitroquinoline-1-oxide and a benzo[a]pyrene compound have been shown to be associated with induction of phase II enzymes.\(^{(64)}\)

A study in mice with transitional cell carcinoma (TCC) of the bladder reported that injection of liquid extract of garlic at the site of tumour transplantation led to a significant reduction in the incidence of TCC in this model.\(^{(65)}\) Furthermore, garlic extract together with suicide-gene therapy significantly inhibited tumour growth (as determined by evidence of apoptosis following histomorphological and immuno histochemical studies) compared with control (no gene therapy).

Effects of garlic constituents on the immune system have been documented in vitro and in vivo; these effects may contribute, at least in part, to the anticancer effects of garlic (see Immunomodulatory activity).

**Immunomodulatory activity**

Immunostimulant activity has been described for a high molecular weight protein fraction obtained from an aged garlic extract.\(^{(66)}\) The fraction was found to strongly stimulate mice peritoneal macrophages in vitro, and to stimulate carbon clearance in mice in vivo. It has been suggested that garlic may suppress tumour cell growth by the stimulation of immunoresponder cells.\(^{(55,66,67)}\)

In vitro and/or in vivo (animal) studies have found that garlic has several immune-enhancing effects, such as stimulation of lymphocyte proliferation and macrophage phagocytosis, induction of macrophage- and lymphocyte-infiltration into transplanted tumours, and stimulation of interferon-γ release.\(^{(67)}\) Other effects on the immune system documented for garlic and/or its constituents include increased natural killer cell activity and increased interleukin-2 production by garlic fractions in vitro,\(^{(68)}\) and increased
numbers of antibody-forming cells in mice spleens following administration of standardised garlic powder.\(^{(3)}\) Other studies demonstrated that, in vitro, aged garlic extract, compared with control, enhanced the proliferation of spleen cells in a concentration-dependent manner, increased production of cytokines (including interleukin-2 and tumour necrosis factor α) and enhanced natural killer cell activity of a T cell fraction of mouse splenic cells against YAC-1 after incubation for 24 hours.\(^{(55)}\) Also, compared with control, aged garlic extract significantly inhibited the growth of sarcoma-180 and LL/2 lung carcinoma cells transplanted into mice, and significant increases in natural killer cell activity of spleen were observed in splenic cells from sarcoma-bearing mice treated with aged garlic extract, compared with those from control mice.\(^{(55, 69)}\)

**Antimicrobial activity**

Antimicrobial activity (including antibacterial, antiviral, antifungal, antiprotozoal and antiparasitic activities) is well documented for garlic.\(^{(3, 7)}\) The *in vitro* antimicrobial activity of garlic is considered to be mainly due to allicin.\(^{(3)}\)

*In vitro* studies have demonstrated that bacteria sensitive to garlic include species from *Staphylococcus*, *Escherichia*, *Proteus*, *Salmonella*, *Providencia*, *Citrobacter*, *Klebsiella*, *Hafnia*, *Aeromonas*, *Vibrio* and *Bacillus* genera.\(^{(7, 70)}\) In these studies, *Pseudomonas aeruginosa* was found not to be sensitive to garlic.\(^{(7, 70)}\) *In vitro* studies have also shown that allicin has significant antibacterial activity against several species, including *Bacillus subtilis*, *Staphylococcus aureus*, *Staphylococcus faecalis*, *Escherichia coli*, *Proteus mirabilis*, *Salmonella typhi* and *Vibrio cholerae*.\(^{(70)}\)

In other *in vitro* studies, garlic oil and four diallyl sulfide constituents, including diallyl disulfide, showed activity against antibiotic-resistant *Pseudomonas aeruginosa* and *Klebsiella pneumoniae*,\(^{(71)}\) and against *S. aureus*, methicillin-resistant *S. aureus*, *Candida* spp. and *Aspergillus* spp.\(^{(72)}\)

Garlic has also been documented to inhibit growth in 30 strains (consisting of 17 species) of mycobacteria, including *Mycobacterium tuberculosis*.\(^{(73)}\) *In vitro*, both aqueous garlic extract and ethanolic garlic extract inhibited the growth of *M. avium* complex (MAC) strains isolated from patients with or without acquired immune deficiency syndrome (AIDS).\(^{(74)}\) Aqueous garlic extract at concentrations of 2–5 mg/mL inhibited the growth of clinical isolates of *Helicobacter pylori* from patients with chronic gastritis or duodenal ulcer.\(^{(75)}\) The minimum inhibitory concentration to inhibit 90% of growth (MIC\(_{90}\)) was 5 mg/mL. Sulfur-containing compounds from garlic (diallyl sulfide and diallyl disulfide, produced from alliin) were shown to decrease growth of *H. pylori* isolates from patients with peptic ulcer.\(^{(76)}\) It has been
proposed that garlic inhibits bacterial cell growth by primarily inhibiting RNA synthesis.\(^{(77)}\)

Broad–spectrum activity against fungi has been documented for garlic including the genera *Microsporum, Epidermophyton, Trichophyton, Rhodotorula, Torulopsis, Trichosporon, Cryptococcus neoformans* and *Candida*, including *Candida albicans*\(^{(7)}\).

Garlic extract has been reported to be more effective than nystatin against pathogenic yeasts, especially *Candida albicans*\(^{(7)}\). Inhibition of lipid synthesis is thought to be an important factor in the anticandidal activity of garlic, with a disulfide–containing component such as allicin thought to be the main active component.\(^{(78)}\) Garlic has been found to inhibit the growth and toxin production of *Aspergillus parasiticus*\(^{(79)}\).

Allicin produced from synthetic alliin with alliinase isolated from garlic cloves inhibited the destruction of baby hamster kidney cells by trophozoites of the protozoan parasite *Entamoeba histolytica in vitro*\(^{(80)}\). Allicin also inhibited the cysteine proteinase activities of intact *E. histolytica* trophozoites. *In vitro* activity against *Giardia intestinalis* has also been documented for whole garlic extract (IC\(_{50}\) 0.3 mg/mL) and for several of its constituents, particularly allyl alcohol and allyl mercaptan (IC\(_{50}\) 7 μg/mL and 37 μg/mL, respectively)\(^{(81)}\).

*In vitro* antiviral activity against parainfluenza type 3, herpes simplex type 1 and influenza B has been documented\(^{(82,83)}\). Activity was attributed to allicin or an allicin derivative. Garlic was reported to be ineffective towards coxsackie B1 virus\(^{(84)}\).

**Antihepatotoxic effects**

Antihepatotoxic activity *in vitro* and *in vivo* has been reported for garlic and its constituents\(^{(3)}\). Garlic oil\(^{(85)}\) and some of its constituents, namely alliin, S-allylmercaptoctysine (ASSC) and S-methylmercaptopcysteine (MSSC) reduced carbon tetrachloride (CCl\(_4\))- and galactosamine–induced hepatotoxicity *in vitro*\(^{(85)}\). Other *in vitro* studies have shown that S-allylcysteine, S-propylcysteine and S-allylmercaptoctysine neutralised CCl\(_4\)-induced hepatotoxicity, and that S-allylcysteine and S-allylmercaptoctysine prevent liver damage induced by hepatotoxins in acute hepatitis in mice\(^{(3,86)}\).

An *in vitro* study in rat hepatocytes found that diallyl sulfide (0.5 and 2 mmol/L) and diallyl disulfide (0.5 and 1 mmol/L) protected against DNA damage induced by aflatoxin B\(_1\), compared with control\(^{(87)}\). In this model, diallyl sulfide and diallyl disulfide appeared to exert a hepatoprotective effect
via increased activity of glutathione-S-transferase and glutathione peroxidase activity.

Other activities

Garlic oil and juice have been reported to protect against isoprenaline–induced myocardial necrosis in rats.\(^{(88)}\) Oral administration of garlic extract 100, 200 or 400 mg/kg to rats given oral lead acetate 5 mg/kg daily for six weeks was found to reduce tissue lead concentrations, compared with those in control rats.\(^{(89)}\) A diet containing 2% aged garlic extract was reported to protect against intestinal damage induced by oral methotrexate and 5–fluorouracil administered to rats for 4–5 days, compared with a control diet.\(^{(90)}\)

A study in senescence–accelerated mice found that S-allylcysteine, present in aged garlic extract, administered in the diet for eight months (40 mg/kg/diet daily) significantly attenuated the decrease in the conditioned avoidance response, compared with a diet lacking S-allylcysteine.\(^{(91)}\) It was suggested that the findings indicate that dietary supplementation with S-allylcysteine may reduce age–related learning disabilities and cognitive disorders in senescence–accelerated mice.

Hypoglycaemic activity has been documented for an alcoholic garlic extract following oral administration to rabbits (dose equivalent to 50 g dry garlic powder). Fifty–nine per cent activity compared to that of 500 mg tolbutamide was observed.\(^{(92)}\)

Garlic has been documented to cause both smooth muscle relaxation and contraction.\(^{(37,52,84)}\) Garlic oil has been reported to depress gastrointestinal movements induced by charcoal meal and castor oil.\(^{(84)}\) In mice, garlic has also inhibited acetylcholine- and PGE\(_2\)-induced contraction of the rat gastric fundus, with the most active components exhibiting the weakest antiplatelet aggregatory activity.\(^{(37)}\) Garlic has also elicited contractions on the rat uterus and the guinea–pig ileum in vitro.\(^{(52)}\) Both actions were blocked by flufenamic acid, but not by atropine or cyproheptadine, indicating a prostaglandin–like mode of action.

In vitro, ajoene was found to inhibit the release of lipopolysaccharide–induced prostaglandin E\(_2\) in macrophages in a concentration–dependent manner.\(^{(93)}\) This effect was reported to be due to inhibition of cyclooxygenase 2 (COX-2) activity by ajoene.

Clinical studies
Pharmacokinetics
The available literature on the metabolism and pharmacokinetics of the constituents of garlic in humans has been reviewed.\(^{(3,94,G18)}\)

Addition of allicin to fresh whole blood results in conversion of allicin to allyl mercaptan and other compounds produced from allicin, such as diallyl trisulfide and ajoene, have also been shown to form allyl mercaptan in blood.\(^{(3)}\) Sulfur–containing compounds, such as diallyl disulfide, diallyl sulfide, dimethyl sulfide and mercapturic acids, have been isolated and identified in human urine following the ingestion of garlic.\(^{(3,95)}\) A subsequent study detected \(N\)-acetyl-\(S\)-allyl-L-cysteine (allylmercapturic acid) in the urine of volunteers \((n = 6)\) who had ingested two garlic tablets containing 100 mg garlic extract (Kwai).\(^{(96)}\) The mean (standard deviation) elimination half–life of allylmercapturic acid was estimated to be 6 (1.3) hours.

It has been reported that the flavour of human breast milk is altered when lactating women consume foods containing sulfur–containing compounds, such as garlic (see Contra–indications, Warnings; Pregnancy and lactation).\(^{(97)}\) Also, garlic ingestion by pregnant women significantly alters the odour of their amniotic fluid (see Contra–indications, Warnings; Pregnancy and lactation),\(^{(98)}\) suggesting that the odorous components of garlic are present. Evidence for this comes from a placebo–controlled study involving 10 healthy pregnant women undergoing routine amniocentesis. The odour of samples of amniotic fluid from women who ingested capsules containing garlic extract was judged to be ‘stronger’ or more ‘garlic-like’ than that of samples from women who had ingested placebo capsules.

Pharmacodynamics
Several of the pharmacological activities documented for garlic and its constituents \textit{in vitro} and \textit{in vivo} (animals) have also been reported in clinical studies (see Clinical studies, Therapeutic effects).

Numerous studies have assessed the effects of the administration of garlic preparations in hypercholesterolaemia.\(^{(3)}\) Many, but not all, of these studies have documented the effects of garlic administration in lowering serum cholesterol and triglyceride concentrations. Clinical studies have also documented fibrinolytic activity associated with garlic administration and effects on platelet function.\(^{(99,100)}\)

Therapeutic effects

\textbf{Anti–atherosclerotic and cholesterol- and lipid– lowering effects}
Numerous studies have investigated the effects of garlic preparations in
lowering raised serum cholesterol concentrations, and the findings of these studies have been reviewed in several meta–analyses.\(^{101–104}\)

A meta–analysis of five randomised, placebo–controlled trials involving mostly patients with serum cholesterol concentrations greater than 5.17 mmol/L who received preparations of garlic extract at doses of 600–1000 mg daily for 8–24 weeks reported that garlic significantly reduced total serum cholesterol concentrations by about 9% (net reduction over placebo), compared with placebo \((p < 0.001)\).\(^{101}\) Another meta–analysis included 16 trials involving patients with a range of disorders (such as hyperlipidaemia, coronary heart disease and hypertension) as well as healthy volunteers, and compared garlic preparations (e.g. fresh garlic, garlic oil, garlic extract, dried garlic powder) with garlic-free diet, placebo or other agents (two studies used bezafibrate or a reserpine/diuretic combination as the comparator treatment).\(^{102}\) This analysis reported a mean difference of \(-0.77\) mmol/L (95% confidence interval (CI) \(-0.65, -0.89\)) in reduction of total serum cholesterol concentrations between garlic recipients and those receiving placebo or a garlic-free diet (net reduction over placebo: 12%). Analysis of data from the eight trials that assessed garlic powder preparations indicated that garlic powder administration significantly reduced serum triglyceride concentrations, compared with placebo (net reduction: 13%). It was stated, however, that several trials had methodological flaws, and that there was not enough evidence to recommend garlic as an effective lipid–lowering agent for routine use. The results of a subsequent randomised placebo–controlled trial involving 115 patients with moderate hyperlipidaemia (which showed no difference between garlic and placebo)\(^{103}\) were included in a re–analysis\(^{103}\) of the meta–analysis described above.\(^{102}\) This analysis showed that the effect of garlic in reducing serum cholesterol concentrations remained statistically significant, compared with placebo, but that the size of the effect was reduced.

Several randomised, double–blind, placebo– controlled trials\(^{105–108}\) have been published since the meta–analyses described above. One of these studies\(^{105}\) has been criticised for its choice of garlic preparation.\(^{109}\)

Several of these trials\(^{105–107}\) were included in the most recent meta–analysis of randomised clinical trials of garlic preparations involving patients with hypercholesterolaemia.\(^{104}\) This meta–analysis included 13 randomised, double–blind, placebo–controlled trials of garlic monopreparations involving 796 patients with coronary heart disease \((n = 1\) trial), hyperlipoproteinaemia \((2)\), hypercholesterolaemia \((7)\), hypertension \((1)\), familial hyperlipidaemia in children \((1)\) and healthy volunteers \((1)\). Ten of the trials assessed the effects of a standardised garlic powder preparation (Kwai) at doses of 600–900 mg
daily for 8–24 weeks; the other three trials tested garlic oil or spray dried powder. Ten trials reported differences favouring garlic over placebo in the reduction of total serum cholesterol concentrations, although these differences were statistically significant in only three studies. Overall, meta-analysis indicated a significant difference in the reduction of total cholesterol concentrations favouring garlic over placebo (−0.41 mmol/L; 95% CI −0.66 mmol/L, −0.15 mmol/L; \( p < 0.01 \)), equivalent to a net reduction in total cholesterol concentrations of 5.8%. It was stated that although these findings indicated that garlic is more beneficial than placebo in reducing serum cholesterol concentrations, the size of the effect is small. Furthermore, several studies had methodological limitations.\(^{104}\)

Another randomised, double-blind, placebo-controlled trial has been published since the meta-analysis described above. This study assessed the effects of garlic powder (tablets) 500 mg and 1000 mg daily, or placebo, for 12 weeks in 53 patients with moderate hypercholesterolaemia (baseline low-density lipoprotein cholesterol (LDL-C) 130–190 mg/dL).\(^{108}\) At the end of the study there were no significant differences in the absolute mean change in LDL-C between the three groups (mean (SD) values were: 0.0 (4.3) mg/dL, 1.4 (4.8) mg/dL and −10.1 (6.8) mg/dL for the placebo, garlic powder 500 mg and garlic powder 1000 mg groups, respectively).

Another meta-analysis, which aimed to summarise the evidence for the effects of garlic on several cardiovascular-related factors, considered 45 randomised controlled trials of at least four weeks’ duration.\(^{110}\) It was reported that after one and three months, garlic treatment may lead to small reductions in total cholesterol concentrations (0.03–0.45 mmol/L and 0.32–0.66 mmol/L, respectively). However, no effect was noted for pooled six-month data. Changes in cholesterol concentrations were paralleled by changes in low-density lipoprotein and triglyceride concentrations.

A randomised, double-blind, placebo-controlled trial explored the anti-atherosclerotic effect of garlic powder 900 mg daily for 48 months in 280 patients with advanced atherosclerotic plaques and an established risk factor for arteriosclerosis (e.g. high systolic blood pressure, hypercholesterolaemia, diabetes mellitus, smoking).\(^{111}\) It was reported that continuous garlic intake significantly reduced the increase in arteriosclerotic plaque volume, compared with placebo. However, the robustness of the findings of this study is difficult to assess independently as no exact \( p \)-value is given.

A Cochrane systematic review of garlic for the treatment of peripheral arterial occlusive disease identified only one eligible randomised, placebo-controlled trial.\(^{112}\) The trial involved 78 participants with peripheral arterial occlusive
disease (lower limb atherosclerosis) who received garlic, or placebo, for 12 weeks. At the end of the study, the difference in the increase in pain-free walking distance between the two groups was found to be statistically non-significant.

An open study involved 101 healthy adults aged 50–80 years who had taken a standardised garlic powder preparation at a dose of at least 300 mg daily for at least two years and 101 age- and sex-matched subjects. Measures of the elastic properties of the aorta were compared for the two groups. Pulse-wave velocity and elastic vascular resistance were reported to be reduced significantly in the garlic group, compared with the control group. These findings suggest that long-term use of garlic powder to attenuate age-related increases in aortic stiffness is worth further study.

**Antithrombotic and fibrinolytic effects**

Several placebo-controlled studies have documented fibrinolytic effects for garlic preparations in clinical studies involving patients with coronary heart disease, hyperlipidaemia and hypercholesterolaemia, and healthy volunteers. Several studies involved the administration of ether-extracted garlic oil 20 mg daily for up to 90 days, whereas several others involved the administration of garlic powder 600–1500 mg daily for up to 28 days. Most, but not all, of these studies reported significant increases in fibrinolytic activity in garlic recipients, compared with placebo recipients.

An open, uncontrolled study explored the effects of garlic consumption (one fresh chopped clove daily for 16 weeks) in eight healthy male volunteers. After 16 weeks, garlic consumption was reported to reduce significantly serum thromboxane B$_2$ and cholesterol concentrations, compared with baseline values.

**Antioxidant effects**

Blood samples from 31 individuals who participated in a randomised, placebo-controlled trial involving 115 patients with moderate hyperlipidaemia who received a standardised garlic powder preparation 900 mg daily for six months (which showed no difference between garlic and placebo) were analysed to explore the effects of garlic treatment on the resistance of low-density lipoprotein to oxidation. There were no significant differences between garlic and placebo recipients in low-density lipoprotein composition. Thus, garlic administration did not reduce the susceptibility of low-density lipoprotein to oxidation. This finding contrasts with some of the results of a double-blind, placebo-controlled study involving 23 patients with coronary artery disease who received a standardised garlic powder preparation (Kwai tablets) 300 mg three times daily, or placebo, for four weeks. The study
reported that garlic powder administration reduced the atherogenicity of low-density lipoprotein. At the end of the study, the ability of low-density lipoprotein to induce intracellular cholesterol accumulation was decreased by 38%, compared with baseline values. Decreases in the susceptibility of low-density lipoprotein to oxidation and in low-density lipoprotein–stimulated cell proliferation (an indicator of low-density lipoprotein atherogenicity) were also documented.

The effects of standardised garlic powder tablets (Sapec; alliin 1.3%, allicin 0.6%) 900 mg daily for two months on oxidative stress status were explored in an open, uncontrolled study involving 25 healthy volunteers. At the end of the study, a reduction in serum malondialdehyde concentrations was observed, compared with baseline values. It was stated that this finding indicates that standardised garlic powder may have antioxidant activity in humans.

**Antihypertensive effects**
A meta-analysis of randomised controlled trials of garlic preparations assessed the evidence for the effects of garlic on blood pressure. Eight trials involving 415 participants were included in the review, all of which had tested the effects of an allicin–standardised garlic powder preparation (Kwai tablets) at doses of 600–900 mg daily for 4–52 weeks. Overall, the absolute change in mean systolic blood pressure was 7.7 mmHg greater for the garlic group, compared with the placebo group (95\% CI 4.3–11.0). However, only three trials specifically involved subjects with hypertension and, as reported by other meta-analyses of trials of garlic preparations, several trials had methodological limitations. Thus, it was stated that there was insufficient evidence to recommend garlic treatment for routine management of hypertension.

Another overview of trials reported that the effects of garlic treatment on blood pressure are ‘insignificant’.

**Anticancer effects**
The protective effects of garlic consumption against various different cancers (including colon, stomach, larynx, breast and endometrial) have been explored in several epidemiological studies, and their findings have been summarised. Most, but not all, of these studies suggest that garlic consumption may have a protective effect, particularly against cancers of the gastrointestinal tract. However, the findings should be interpreted cautiously, as bias and/or confounding cannot be excluded, and there are other methodological issues, for example, most studies did not distinguish between consumption of raw or cooked garlic.
An epidemiological study comprising a dietary interview and measurement of serum *H. pylori* antibodies was conducted among 214 adults in a low-risk area of Shandong Province in China.\(^{(120)}\) The findings suggested a protective effect of garlic consumption against *H. pylori* infection.

An open, uncontrolled study involving 34 patients with tinea pedis explored the effectiveness of ajoene cream (0.4% w/w).\(^{(121)}\) After seven days’ treatment, complete cure of the infection was recorded for 27 (79%) participants. The remaining seven patients experienced complete cure after a further seven days’ treatment. All patients were evaluated for recurrence of infection 90 days after the end of treatment; all found to be infection free as determined by negative cultures for the fungus.

**Other effects**

A reduction in blood sugar concentrations and an increase in insulin have been observed following allylpropyl disulfide administration to normal volunteers, whereas another study reported that garlic exhibits hypoglycaemic actions in diabetic patients but not in controls.\(^{(5)}\) It has also been reported that garlic can prevent tolbutamide- and adrenaline–induced hyperglycaemia.\(^{(5)}\)

A randomised, placebo–controlled, crossover trial involving 100 Swedish participants working in a tick–endemic area found that administration of garlic powder 1200 mg daily for eight weeks resulted in fewer tick bites than did placebo administration.\(^{(122)}\)

An open, uncontrolled study involving 15 patients with hepatopulmonary syndrome explored the effects of treatment with capsules containing a standardised garlic powder preparation for at least six months.\(^{(123)}\) Improvements in arterial oxygenation and symptoms were documented for several participants. The effects of garlic powder treatment in patients with hepatopulmonary syndrome require further study.
Side-effects

Garlic is generally considered to be non-toxic. Adverse effects that have been documented in humans include a burning sensation in the mouth and gastrointestinal tract, nausea, diarrhoea and vomiting.

A meta-analysis of 13 randomised, double-blind, placebo-controlled trials of garlic monopreparations, 10 of which assessed the effects of a standardised garlic powder preparation (Kwai) at doses of 600–900 mg daily for 8–24 weeks (see Clinical studies, Therapeutic effects, Anti-atherosclerotic and cholesterol- and lipid-lowering effects) reported that few adverse events were documented in the included trials. The frequency and nature of adverse events reported for garlic were similar to those for placebo. The most common adverse events reported were ‘garlic breath’, body odour and gastrointestinal symptoms.

The allergenic potential of garlic is well recognised, and allergens have been identified as diallyl disulfide, allylpropyl sulfide and allicin (the latter may be an irritant). A garlic antigen in the serum of affected patients has also been identified. Cases of contact dermatitis resulting from occupational exposure to garlic have been reported. A case of garlic allergy associated with ingestion of raw or cooked garlic has been documented. There is an isolated report of multifaceted dermatitis artefacta associated with local application of garlic by a 19-year-old individual. Garlic burns following local application of garlic have also been documented.

Garlic may enhance existing anticoagulant therapy; a potential interaction between garlic and warfarin has been documented. Case reports have suggested that garlic supplementation may increase the risk of bleeding in patients undergoing surgery.

Toxicity

Erratic pulse rates, abnormal ECGs, weight loss, lethargy and weakness, soft faeces, dehydration and tender skin on fore and hindlimbs have been observed in spontaneously hypertensive rats administered garlic extract at 0.25 and 0.5 mL/kg every 6 hours for 28 days. The effects were most pronounced in animals receiving doses two or three times a day. Conversely, acute toxicity studies for garlic extract in mice and rats have reported LD$_{50}$ values for various routes of administration (by mouth, intraperitoneal injection, intravenous injection) as all greater than 30 mL/kg. Early studies, in 1944, reported LD$_{50}$ values for allicin in mice as 120 mg/kg.
(subcutaneous injection) and 60 mg/kg (intravenous injection). Results of chronic toxicity studies are stated to be conflicting. High doses are reported to cause anaemia due to both decreased haemoglobin synthesis and haemolysis. A chronic toxicity study in rats given a garlic extract (2 g/kg) five times a week for six months, reported no toxic symptoms. High doses were found to decrease food consumption slightly, but did not inhibit weight gain. There were no significant differences in urinary, haematological or serological examinations, and no toxic symptoms in histopathological examinations. Genotoxicity studies using the micronucleus test have reported both positive and negative findings. No evidence of mutagenicity has been reported when assessed using the Ames and Ree assay.

Slight cytotoxic signs have been observed at high doses in Hep2 and Chinese hamster embryo primary cultured cells.

The literature relating to the toxicity of garlic has been reviewed.
Contra-indications, Warnings

In view of the pharmacological actions documented for garlic, therapeutic doses of garlic may interfere with existing hypoglycaemic and anticoagulant therapies. There may be an increased risk of bleeding with use of garlic supplements in patients undergoing surgery. Garlic may potentiate the antithrombotic effects of anti-inflammatory drugs such as aspirin, and may be synergistic with eicosapentaenoic acid (EPA) in fish oils. Gastrointestinal irritation may occur particularly if the clove is eaten raw by individuals not accustomed to ingesting garlic.

A study involving healthy volunteers detected N-acetyl-S-allyl-L-cysteine (allylmercapturic acid) in their urine following ingestion of garlic tablets (see Clinical studies, Pharmacokinetics). As allylmercapturic acid is used as a biomarker for monitoring human exposure to allylhalides and other chemicals leading to allylmercapturic acid excretion, it was suggested that garlic consumption may interfere with and confound this monitoring process.

Pregnancy and lactation
Garlic is reputed to act as an abortifacient and to affect the menstrual cycle, and is also reported to be utero-active. In vitro uterine contraction has been documented.

Studies have shown that consumption of garlic by lactating women alters the odour of their breast milk and the suckling behaviour of their infants. Further evidence for this comes from a blinded, placebo-controlled study involving 30 nursing women. The results indicated that infants who had no prior exposure to garlic odour in their mothers’ milk spent more time breast feeding after their mothers ingested garlic capsules than did infants whose mothers had repeatedly consumed garlic. Findings from a placebo-controlled study involving 10 healthy pregnant women undergoing routine amniocentesis indicate that the odorous components of garlic can be found in amniotic fluid following garlic consumption. The odour of samples of amniotic fluid from women who ingested capsules containing garlic extract was judged to be ‘stronger’ or more ‘garlic-like’ than that of samples from women who had ingested placebo capsules. The effects of in utero exposure to garlic odour and on the neonate’s behaviour towards exposure to garlic-flavoured human breast milk are not known.

There are no experimental or clinical reports on adverse effects during pregnancy or lactation. In view of this, doses of garlic greatly exceeding amounts used in foods should not be taken during pregnancy and lactation.
Pharmaceutical Comment

There is a vast scientific literature on the chemistry, pharmacology and clinical properties of garlic. Experimental studies have focused mainly on the cardiovascular and anticancer effects of garlic and its constituents, as well as its antimicrobial properties. Clinical studies have investigated mainly the anti-atherosclerotic and cholesterol- and lipid-lowering effects of garlic preparations. Generally, these studies report beneficial results for garlic, although the evidence at present is insufficient to recommend garlic as routine treatment for hypercholesterolaemia. *In vitro* and animal studies provide supporting evidence for some of the clinical properties of garlic and its constituents.

Garlic is characterised by its sulfur-containing constituents. Pharmacological activities documented for garlic are also associated with these compounds. It is recognised that allicin, the unstable compound formed by enzymatic action of allinase on alliin when the garlic clove is crushed, is required for the antimicrobial activity that has been demonstrated by garlic. However, serum concentrations of allicin achieved in humans following oral ingestion of garlic are unclear. The hypolipidaemic and antithrombotic actions documented for garlic have been attributed to many of the degradation products of alliin.

One of the difficulties in comparing studies that have investigated the efficacy of garlic, is establishing the concentration of active principles present in the garlic preparations used. It has been reported that the percentage of active constituents in fresh garlic may vary by a factor of 10. Many commercial garlic preparations are standardised on content of sulfur-containing constituents, particularly to alliin, or on allicin yield. Dried garlic powder contains both alliin and allinase and therefore has an allicin-releasing potential. Garlic preparations produced by heat or solvent extraction processes are stated to contain alliin but to be devoid of allinase and therefore have no allicin releasing potential. Garlic oil macerates and steam distillation products are rich in secondary alliin metabolites, such as ajoene. However, it is unclear to what extent these secondary compounds are formed in the body following the ingestion of garlic and whether, therefore, these products exhibit the pharmacological actions of fresh garlic.

Fermented garlic preparations are considered to be practically devoid of the active sulfur-containing compounds. Many ‘odourless’ garlic preparations are available: obviously one should establish if these products are odourless due to the formulation of the product or because they are devoid of the odoriferous, active principles. Further randomised controlled clinical trials with standardised preparations are required to establish the true usefulness.
of garlic in reducing serum lipids, blood pressure, platelet aggregation and exerting an antimicrobial effect. Therapeutic doses of garlic should not be given to those whose blood clots slowly and caution is recommended for patients on anticoagulant therapy. (G58)
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Gentian
Species (Family)

*Gentiana lutea* L. (Gentianaceae)
Part(s) Used

Rhizome, root
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

BP 2002\(^{(G71)}\)

Complete German Commission E\(^{(G3)}\)

ESCOP 1997\(^{(G52)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)

Ph Eur 2004\(^{(G72)}\)
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents
See General References G2 G6 G22 G41 G60 G62 G64.

Alkaloids
Pyridine–type. Gentianine 0.6–0.8%, gentialutine.

Bitters
Major component is secoiridoid glycoside gentiopicroside (also known as gentiamarin and gentiopicrin) 2%, with lesser amounts of amarogentin (0.01–0.04%) and swertiamarine.\(^1\) Gentianose (a trisaccharide bitter principle). The glycosides amaropanin and amaroswerin are reported to be present in the related species Gentiana pannonica, Gentiana punctata and Gentiana purpurea, but are absent from Gentiana lutea.

Xanthones
Gentisein, gentisin (gentianin), isogentisin and 1,3,7-trimethoxyxanthone.

Other constituents
Carbohydrates (e.g. gentiobiose, sucrose and other common sugars), pectin, tannin (unspecified), triterpenes (e.g. β-amyrin, lupeol) and volatile oil (trace).
Food Use

Gentian (root, herbs and preparations) is listed by the Council of Europe as a natural source of food flavouring (category 4, with limits on xanthones) (see Appendix 23).\(^{(G17)}\) In the USA, gentian is approved for food use.\(^{(G41)}\)
Herbal Use

Gentian is stated to possess bitter, gastric stimulant, sialogogue and cholagogue properties. Traditionally, it has been used for anorexia, atonic dyspepsia, gastrointestinal atony, and specifically for dyspepsia with anorexia. The German Commission E approved use for digestive disorders such as loss of appetite, fullness and flatulence. (G3) Gentian is used in combination with angelica root and caraway fruit or with ginger and wormwood for loss of appetite and peptic discomfort. (G3)
Dosage

Dried rhizome/root
0.6–2 g or by infusion or decoction three times daily.\(^{(G6)}\)

Tincture
1–4 mL (1 : 5 in 45% alcohol) three times daily.\(^{(G6)}\)
Pharmacological Actions

**In vitro and animal studies**

The pharmacological activities of gentian root have been reviewed.\(^{(G52)}\) A summary of this information is provided below.

Root extracts have antifungal activity, and are reported to stimulate phagocytic activity of human lymphocytes, indicating immunostimulant activity.\(^{(G52)}\) Choleretic properties have been documented for gentian,\(^{(G41)}\) and gentianine has been reported to possess anti–inflammatory activity.\(^{(G22)}\)

The bitter principles stimulate secretion of gastric juices and bile, thus aiding appetite and digestion. Elevation of gastric secretion by up to 30% has been reported following the administration of gentian tincture to dogs. An infusion given orally to sheep as a single daily dose (5 g) stimulated enzyme secretion in the small intestine. A root extract (12 mg/kg/day) applied by gavage to rats for three days elevated bronchosecretion. A standardised extract perfused into the stomachs of anaesthetised rats increased gastric secretion in a dose–dependent manner. Lower doses caused no changes in gastric pH, whereas higher doses increased pH from 4.25 to 4.85. A dose of 0.5 mL/kg did not affect the incidence of gastric ulceration in rats.

**Clinical studies**

In an open, uncontrolled study, a single dose of an alcoholic extract of gentian (equivalent to 0.2 g), given to 10 healthy volunteers, was reported to result in a stimulation of gastric juice secretion.\(^{(2)}\) Gall–bladder emptying was increased and prolonged whilst protein and fat digestion was enhanced.

Nineteen patients with inflammatory conditions of the gastrointestinal tract (colitis, Crohn’s disease, non–specific inflammation) and elevated secretory immunoglobulin A (IgA) concentrations and eight healthy individuals were treated with gentian tincture (3 × 20 drops/day) for eight days.\(^{(G52)}\) IgA concentrations decreased in both groups.\(^{(G52)}\)
Side-effects, Toxicity

Extracts of gentian are considered to be non-toxic, and are generally well-tolerated.\(^{(G52)}\)

An acute oral LD\(_{50}\) value in mice was reported to be 25 mL/kg of extract (37% ethanol, bitterness value: 200 Swiss Pharmacopoeia units/g), and was the same as that of 37% ethanol. Rabbits treated with gentian extract (12.6 mg/day for three days) showed no toxic or abnormal concentration of serum parameters, with the exception of slightly higher erythrocyte concentrations in treated animals. Gentian may occasionally cause headache in some individuals.\(^{(G3)}\) Mutagenic activity in the Ames test (\textit{Salmonella typhimurium} TA100 with S9 mix) has been documented for gentian, with gentisin and isogentisin identified as mutagenic components.\(^{(3)}\) Gentian root 100 g was reported to yield approximately 100 mg total mutagenic compounds, of which gentisin and isogentisin comprised approximately 76 mg.\(^{(3)}\)
Contra–indications, Warnings

Gentian is stated to be contra–indicated in individuals with high blood pressure,\(^{(G60)}\) although no rationale is given for this statement, and in individuals with hyperacidity, gastric or duodenal ulcers.\(^{(G52 \ G3)}\).

**Pregnancy and lactation**

Gentian is reputed to affect the menstrual cycle,\(^{(G22 \ G60)}\) and it has been stated that gentian should not be used in pregnancy.\(^{(G60)}\) In view of this and the documented mutagenic activity, gentian is best avoided in pregnancy and lactation.
Pharmaceutical Comment

The major constituents of pharmacological importance in gentian are the bitter principles; limited information is available on the other compounds present. The herbal uses of gentian are supported by the known properties of the bitter principles present in the root. Excessive doses should be avoided in view of the lack of toxicity data.
References


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Ginger
Species (Family)

*Zingiber officinale* Roscoe (Zingiberaceae)
Synonym(s)

Zingiber
Part(s) Used

Rhizome
Pharmacopoeial and Other Monographs

BHC 1992 (G6)

BHP 1996 (G9)

BP 2002 (G71)

BPC 1973 (G12)

ESCOP 1996 (G52)

Martindale 33rd edition (G67)

Mills and Bone (G50)

PDR for Herbal Medicines 2nd edition (G36)

Ph Eur 2004 (G72)

USP26/NF21 (G73)

WHO 1999 volume 1 (G63)
Legal Category (Licensed Products)

GSL$^{(G37)}$
Constituents
See Reference 1, and General References G2 G6 G37 G41 G64.

**Carbohydrates**
Starch (major constituent, up to 50%).

**Lipids**
6–8%. Free fatty acids (e.g. palmitic acid, oleic acid, linoleic acid, caprylic acid, capric acid, lauric acid, myristic acid, pentadecanoic acid, heptadecanoic acid, stearic acid, linolenic acid, arachidic acid);\(^{(2)}\) triglycerides, phosphatidic acid, lecithins; gingerglycolipids A, B and C.\(^{(3)}\)

**Oleo–resin**
Gingerol homologues (major, about 33%) including derivatives with a methyl side–chain,\(^{(4)}\) shogaol homologues (dehydration products of gingerols), zingerone (degradation product of gingerols), 1–dehydrogingerdione,\(^{(5)}\) 6–gingesulfonic acid\(^{(3)}\) and volatile oils.

**Volatile oils**
1–3%. Complex, predominately hydrocarbons. β-Bisabolene and zingiberene (major); other sesquiterpenes include zingiberol, zingiberenol, ar-curcumene, β-sesquiphellandrene, β-sesquiphellandrol (cis and trans); numerous monoterpenene hydrocarbons, alcohols and aldehydes (e.g. phellandrene, camphene, geraniol, neral, linalool, \(d\)-nerol).

**Other constituents**
Amino acids (e.g. arginine, aspartic acid, cysteine, glycine, isoleucine, leucine, serine, threonine and valine), protein (about 9%), resins, diterpenes (galanolactone\(^{(6)}\)), vitamins (especially nicotinic acid (niacin) and vitamin A), minerals.\(^{(2)}\)

The material contains not less than 4.5% of alcohol (90%)-soluble extractive and not less than 10% of water–soluble extractive.\(^{(G15)}\)
Food Use

Ginger is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that ginger can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\(^{(G16)}\) It is used widely in foods as a spice. In the USA, ginger is listed as GRAS (Generally Recognised As Safe).\(^{(G41)}\)
Herbal Use

Ginger is stated to possess carminative, diaphoretic and antispasmodic properties. Traditionally, it has been used for colic, flatulent dyspepsia, and specifically for flatulent intestinal colic.\(^{(7,G2,G6,G32,G64)}\) Modern interest in ginger is focused on its use in the prevention of nausea and vomiting, particularly motion (travel) sickness, as a digestive aid, and as an adjunctive treatment for inflammatory conditions, such as osteoarthritis and rheumatoid arthritis.
Dosage

Anti-emetic

*Powdered rhizome*
Single dose of 1–2 g,\(^{(G6)}\) 30 minutes before travel for prevention of motion sickness,\(^{(G52)}\) or 0.5 g, two to four times daily.\(^{(G63)}\)
Other uses

**Powdered rhizome**
0.25–1 g, three times daily.\(^{(G6)}\)

**Tincture**
1.5–3 mL (1 : 5) three times daily,\(^{(G6)}\) 1.7–5 mL daily.\(^{(G50)}\)
Pharmacological Actions

Several pharmacological activities, including anti-emetic, antithrombotic, antimicrobial, anticancer, antioxidant and anti-inflammatory properties, have been documented for preparations of ginger in *in vitro* and/or animal studies. Also, ginger has been reported to have hypoglycaemic, hypo- and hypertensive, cardiac, prostaglandin and platelet aggregation inhibition, antihypercholesterolaemic, cholagogic and stomachic properties.

Clinical studies have focused mainly on the effects of ginger in the prevention of nausea and vomiting.

**In vitro and animal studies**

*In vitro* studies have demonstrated that constituents of ginger, such as 6-, 8- and 10–gingerols and galanolactone, have antiserotonergic activity.\(^6,8\)

**Anti-emetic activity and effects on gastrointestinal motility**

The older literature contains examples of studies documenting the antiemetic effects of ginger extract *in vivo* (e.g. dogs).\(^9\) Oral administration of constituents of ginger (certain shogaols and gingerols at doses of 100 mg/kg body weight) inhibited emesis induced by oral administration of copper sulfate in leopard and ranid frogs.\(^10\) Emetic latency was reported to be prolonged by over 150% by a trichloro methane extract of ginger at a dose of 1 g/kg body weight.

The anti-emetic activity of ginger extracts has also been assessed in dogs.\(^11\) Acetone and ethanolic extracts of ginger, administered intragastrically at doses of 25, 50, 100 and 200 mg/kg, protected against cisplatin–induced emesis (3 mg/kg administered intravenously 30 minutes before ginger extract), compared with control. However, ginger extracts were less effective in preventing emesis than the 5-HT\(_3\) receptor antagonist granisetron, and were ineffective against apomorphine–induced emesis.

Compared with control, an acetone extract of ginger at doses of 200 and 500 mg/kg administered orally reversed the delay in gastric emptying induced by intraperitoneal cisplatin 10 mg/kg in rats.\(^12\) Ginger juice (2 and 4 mL/kg) had a similar effect. A 50% ethanolic extract of ginger also reversed the cisplatin–induced delay in gastric emptying, although only at a dose of 500 mg/kg. In mice, oral administration of an acetone extract of ginger (75 mg/kg), 6–shogaol (2.5 mg/kg) and 6-, 8- and 10–gingerol (5 mg/kg) enhanced the transportation of a charcoal meal, indicating enhancement of gastrointestinal motility.\(^13\)
**Anti-ulcer activity**
The effect of ginger (acetone extract) and zingiberene on hydrochloric acid/ethanol–induced gastric lesions in rats has been examined. \(^{(14)}\) (6)-Gingerol and zingiberene, both 100 mg/kg body weight by mouth, significantly inhibited gastric lesions by 54.5% and 53.6%, respectively. The total extract inhibited lesions by 97.5% at 1 g/kg. Oral administration of both aqueous and methanol ginger extracts to rabbits has been reported to reduce gastric secretions (gastric juice volume, acid and pepsin output). \(^{(15)}\) Both extracts were found to be comparable with cimetidine (50 mg/kg) with respect to gastric juice volume; the aqueous extract was comparable with cimetidine and superior to the methanol extract for pepsin output, and the methanol extract superior to both the aqueous extract and comparable to cimetidine for acid output. In rats, 6–gingerol, 6–shogaol and 6–gingesulfonic acid at doses of 150 mg/kg protected against hydrochloric acid/ethanol–induced gastric lesions, compared with control. \(^{(3)}\) 6-Gingesulfonic acid 300 mg/kg provided almost 100% protection against gastric lesions in this model. Other studies in rats found that oral administration of an ethanolic extract of ginger (500 mg/kg) inhibited gastric lesions induced by ethanol (80%), hydrochloric acid (0.6 mol/L), sodium hydroxide (0.2 mol/L), and 25% sodium chloride, compared with control. \(^{(16)}\) The same dose of extract protected against gastric mucosal damage induced by the non–steroidal anti–inflammatory drugs (NSAIDs) indometacin and aspirin in rats. In pylorus–ligated rats, oral administration of acetone and ethanol extracts of ginger inhibited gastric secretion. \(^{(17)}\) These extracts, at doses of 62 mg/kg, also protected against the development of stress–induced lesions, although to a lesser extent than cimetidine.

**Antiplatelet activity**
(6)-Gingerol, (6)- and (10)-dehydrogingerdione, (6)- and (10)-gingerdione have been reported to be potent inhibitors of prostaglandin biosynthesis (PG synthetase) *in vitro*, with the latter four compounds stated to be more potent than indometacin. \(^{(18)}\) Dose–dependent inhibition of platelet aggregation, *in vitro*, induced by ADP, adrenaline, collagen and arachidonic acid has been described for an aqueous ginger extract. \(^{(19)}\) Ginger was also found to reduce platelet synthesis of prostaglandin–endo peroxides, thromboxane and prostaglandins. A good correlation was reported between concentrations of the extract required to inhibit platelet aggregation and concentrations necessary to inhibit platelet thromboxane synthesis. \(^{(19)}\)

**Anti–atherosclerotic and antioxidant activity**
Ginger oleo–resin, by intragastric administration, has been reported to inhibit elevation in serum and hepatic cholesterol concentrations in rats by impairing
cholesterol absorption. Antihyperchol esterolaemic activity has also been documented for dried ginger rhizome when given to both rats fed a cholesterol–rich diet and those with existing hyperchol esterolaemia. Fresh ginger juice was not found to have an effect on serum cholesterol concentrations within 4 hours of administration. In addition, serum cholesterol concentrations were not greatly increased within 4 hours of cholesterol administration.

An ethanol (50%) extract of ginger administered orally at a dose of 500 mg/kg to hyperlipidaemic rabbits led to a significant reduction in blood serum cholesterol concentrations, compared with those in control rabbits. In a study in rabbits fed cholesterol for 10 weeks, administration of an ethanolic extract of ginger (200 mg/kg orally) decreased raised serum and tissue concentrations of cholesterol, serum triglycerides and serum lipoproteins.

An ethanolic ginger extract, standardised to contain 40 mg/g gingerols, shogaols and zingerone, and 90 mg/g total polyphenols, was reported to inhibit low–density lipoprotein oxidation and to reduce the development of atherosclerosis in atherosclerotic mice, when compared with control. In rats fed a high–fat diet for 10 weeks, an aqueous preparation of ginger powder administered orally at doses of 35 and 70 mg/kg demonstrated antioxidant activity, as measured by raised tissue concentrations of thiobarbituric acid reactive substances and hydroperoxides, and reduced activities of superoxide dismutase and catalase.

The antioxidant activity of ginger constituents has been documented in vitro.

**Anti–inflammatory activity**

Constituents of ginger have been shown to have anti–inflammatory activity in vitro. In a study in intact human airway epithelial cells (A549 cells), 8–paradol and 8–shogaol inhibited cyclooxygenase 2 (COX-2) enzyme activity in a concentration–dependent manner (IC$_{50}$ values ranged from 1 to 25 μmol/L). In other studies, an acetone extract of ginger inhibited inflammation of the chorioallantoic membrane of fertilised hen’s eggs in a concentration–dependent manner. In another assay, the extract exhibited anti–inflammatory properties by inhibiting the release of nitric oxide in a concentration–dependent manner. Ginger oil has demonstrated anti–inflammatory activity in a study in rats with severe chronic adjuvant arthritis induced by injection of 0.05 mL of a suspension of dead *Mycobacterium tuberculosis* bacilli. Ginger oil 33 mg/kg administered orally for 26 days caused a significant suppression of paw and joint swelling, compared with
control (no ginger oil).

Other studies documenting anti-inflammatory activity for ginger constituents have been summarised.  

**Antimicrobial activity**

*In vitro* activity against rhinovirus IB has been reported for sesquiterpenes isolated from ginger rhizomes.\(^1\) The most active compound was β-sesquiphellandrene (IC\(_{50}\) 0.44 μmol/L). *In vitro* anthelmintic activity against *Ascaridia galli* Schrank has been documented for the volatile oil of *Zingiber purpureum* Roxb.\(^{30}\) Activity exceeding that of piperazine citrate was exhibited by the oxygenated compounds fractionated from the volatile oil.

**Anticancer activity**

Extracts of ginger or constituents of ginger have been shown to have cancer chemopreventive and cytotoxic or cytostatic activity *in vitro* and *in vivo* (animals). Application of an ethanolic extract of fresh ginger in a mouse skin tumorigenesis model (SENCAR mice) resulted in significant inhibition of 12-O-tetradecanoylphorbol-13-acetate (TPA)-induced induction of epidermal ornithine decarboxylase, cyclooxygenase and lipoxygenase activities in a concentration-dependent manner.\(^{31}\) Preapplication of ginger extract also inhibited TPA-induced epidermal oedema and hyperplasia. Application of ginger extract 30 minutes before application of two tumour inducers to the skin of SENCAR mice protected against skin tumour incidence, compared with control. In another mouse model, topical application of 6-gingerol or 6-paradol before application of tumour inducers attenuated skin papillomagenesis.\(^{32}\) Other studies documenting the cancer chemopreventive potential of ginger and its constituents have been summarised.\(^{26}\)

*In vitro*, incubation of 6-gingerol with human promyelocytic leukaemia (HL-60) cells resulted in inhibitory effects on cell viability and DNA synthesis.\(^{33}\) Microscopic examination of the incubated cells provided evidence of the induction of apoptosis by 6-gingerol.

**Other activities**

In rats, the anxiolytic effects of pretreatment with a combination preparation of standardised extracts of ginger and *Ginkgo biloba* administered intragastrically at doses between 0.5 and 100 mg/kg were assessed in the elevated plus-maze test.\(^{34}\) The combination was found to have an anxiolytic effect at lower doses, but appeared to have an anxiogenic effect at higher doses.

A hypoglycaemic effect in both non-diabetic and alloxan-induced diabetic
rabbits and rats has been documented for fresh ginger juice administered orally. The effect was stated to be significant in the diabetic animals.\footnote{35}

The pharmacological actions of (6)-shogaol and capsaicin have been compared.\footnote{36} Both compounds caused rapid hypotension followed by a marked pressor response, bradycardia, and apnoea in rats after intravenous administration. The pressor response was thought to be a centrally acting mechanism. Contractile responses in isolated guinea–pig trachea with both compounds, and positive inotropic and chronotropic responses in isolated rat atria with (6)-shogaol were thought to involve the release of an unknown active substance from nerve endings.\footnote{36} A potent, positive inotropic action on isolated guinea–pig atria has been documented and gingerols were identified as the cardiotonic principles.\footnote{37}

A cholagogic action in rats has been described for an acetone extract of ginger administered intra duodenally.\footnote{38} (6)-Gingerol and (10)-gingerol were reported to be the active components, the former more potent with a significant increase in bile secretion still apparent 4 hours after administration.

Utero–activity has been described for a phenolic compound isolated from \textit{Zingiber cassumunar} Roxb.\footnote{39} The compound was found to exhibit a dose–related relaxant effect on the non–pregnant rat uterus \textit{in situ}; the uterine response from pregnant rats was stated to vary with the stage of pregnancy, the post–implantation period being the most sensitive. The compound was thought to act by a similar mechanism to that of papaverine.\footnote{39}

\section*{Clinical studies}

Clinical trials of ginger have focused mainly on its effects on the prevention and treatment of nausea and vomiting of various causes. Other clinical studies have assessed the effects of ginger preparations on gastrointestinal motility and on platelet function, and in vertigo and inflammatory conditions, such as osteo arthritis. Several of these studies are described below.

\subsection*{Nausea and vomiting and effects on gastrointestinal motility}

Ginger has been reported to be effective as a prophylactic against seasickness.\footnote{40,41} Ingestion of powdered ginger root 1 g was found to significantly reduce the tendency to vomit and experience cold sweating in 40 naval cadets, compared with 39 cadets who received placebo.\footnote{40} Powdered ginger root 1.88 g has been reported to be superior to dimenhydrinate 100 mg in preventing the gastrointestinal symptoms of motion sickness induced by a rotating chair.\footnote{41} However, a second study reported ginger
(500 mg powdered, 1 g powdered/fresh) to be ineffective in the prevention of motion sickness induced by a rotating chair.\(^{(42)}\) The study concluded hyoscine 600 μg and dexamfetamine 10 mg to be the most effective combination, with dimenhydrinate 50 mg as the over-the-counter motion sickness medication of choice.\(^{(42)}\)

A systematic review of 6 randomised controlled trials of ginger preparations included three trials involving patients with post-operative nausea and vomiting, and three further trials in patients with seasickness (motion sickness), morning sickness (emesis of pregnancy) and cancer chemotherapy-induced nausea (one trial in each condition).\(^{(43)}\) Two of the three studies assessing the effects of ginger in post-operative nausea and vomiting found that ginger was more effective than placebo and as effective as metoclopramide in reducing nausea. However, when the data from the three studies were pooled, the difference between the ginger and placebo groups was statistically non-significant.\(^{(43)}\)

A randomised, double-blind, crossover trial involving women with nausea of pregnancy assessed the effects of capsules of powdered ginger root 250 mg, or placebo, administered orally four times daily for four days.\(^{(44)}\) It was reported that symptom relief was significantly greater during treatment with ginger than with placebo, and that significantly more women stated a preference for ginger treatment than for placebo (as later disclosed). A more recent randomised, double-blind trial involving 70 women with nausea and vomiting of pregnancy assessed the effectiveness of capsules of powdered fresh ginger root 250 mg four times daily, or placebo, for four days.\(^{(45)}\) At the end of the study, ginger recipients had significantly lower scores for nausea and fewer vomiting episodes than did the placebo group.

Studies involving healthy volunteers have investigated the effects of ginger on gastric emptying as a possible mechanism for the anti-emetic effects of ginger. A randomised, double-blind, placebo-controlled, crossover trial involving 16 volunteers assessed the effects of capsules containing powdered ginger 1 g for one week, followed by a one-week washout period before crossing over to the opposite arm of the study.\(^{(46)}\) Gastric emptying was measured using a paracetamol absorption technique by comparing the effects of ginger administration on mean and peak plasma paracetamol concentrations. The results indicated that the rate of absorption of oral paracetamol was not affected by simultaneous ingestion of ginger. Another randomised, double-blind, placebo-controlled trial involving 12 healthy volunteers assessed the effects of ginger rhizome extract on fasting and postprandial gastroduodenal motility.\(^{(47)}\) The results of this study indicated that oral administration of ginger improved gastroduodenal motility in both
the fasting state and after a test meal.

A randomised, double-blind, placebo-controlled, crossover trial involving eight healthy volunteers tested the effects of powdered ginger root 1 g on experimentally induced vertigo. One hour after ginger or placebo administration, participants’ vestibular system was stimulated by water irrigation of the left ear. It was reported that ginger significantly reduced vertigo, when compared with placebo.

**Other effects**

In a randomised, double-blind, placebo-controlled, crossover trial involving 75 patients with osteoarthritis of the knee or hip, the effects of capsules of ginger extract 170 mg three times daily were compared with those of ibuprofen 400 mg three times daily, or placebo, for three weeks with a one-week washout period between each treatment period. At the end of the study, data for the 56 evaluable participants indicated that there was no strong evidence of an effect for ginger extract over that of placebo on parameters of pain.

A reduction in joint pain and improvement in joint movement in seven rheumatoid arthritis sufferers has been documented for ginger, with a dual inhibition of cyclooxygenase and lipoxygenase pathways reported as a suggested mechanism of action. Patients took either fresh ginger in amounts ranging from 5 to 50 g or powdered ginger 0.1–1.0 g daily.

A placebo-controlled study assessed the effects of two doses of ginger powder (4 g daily for three months, and 10 g as a single dose) on platelet aggregation and fibrinolytic activity in patients with coronary artery disease (CAD). The results indicated that long-term administration of ginger powder did not affect ADP- and epinephrine (adrenaline)-induced platelet aggregation and had no effects on fibrinolytic activity or fibrinogen concentrations, compared with placebo administration. By contrast, administration of a single dose of ginger powder to 10 patients with CAD produced a significant reduction in platelet aggregation, compared with placebo administration (n = 10 patients with CAD).

In a study involving seven women, oral raw ginger 5 g reduced thromboxane B₂ concentrations in serum collected after clotting, thus indicating a reduction in eicosanoid synthesis (associated with platelet aggregation).
Side–effects, Toxicity

None documented for ginger. Ginger oil is stated to be non–irritating and non–sensitising although dermatitis may be precipitated in hypersensitive individuals. Phototoxicity is not considered to be of significance.(53) Ginger oil is stated to be of low toxicity(G58) with acute LD₅₀ values (rat, by mouth; rabbit, dermal) reported to exceed 5 g/kg.(53)

Mutagenic activity has been documented for an ethanolic ginger extract, gingerol and shogaol in Salmonella typhimurium strains TA100 and TA1535 in the presence of metabolic activation (S9 mix) but not in TA98 or TA1538 with or without S9 mix.(54) Zingerone was found to be non–mutagenic in all four strains with or without S9 mix, and was reported to suppress mutagenic activity of gingerol and shogaol. Ginger juice has been reported to exhibit antimutagenic activity, whereas mutagenic activity has been described for (6)-gingerol in the presence of known chemical mutagens.(55) It was suggested that certain mutagens may activate the mutagenic activity of (6)-gingerol so that it is not suppressed by antimutagenic components present in the juice.(55)
Contra–indications, Warnings

Ginger has been reported to possess both cardiotonic and antiplatelet activity \textit{in vitro} and hypoglycaemic activity \textit{in vivo} studies. Excessive doses may therefore interfere with existing cardiac, antidiabetic or anticoagulant therapy. An oleo–resin component, (6)-shogaol has been reported to affect blood pressure (initially decrease then increase) \textit{in vivo}.

\textbf{Pregnancy and lactation}

Ginger is reputed to be an abortifacient\textsuperscript{(G30)} and uteroactivity has been documented for a related species. Doses of ginger that greatly exceed the amounts used in foods should not be taken during pregnancy or lactation.
Pharmaceutical Comment

The chemistry of ginger is well documented with respect to the oleo–resin and volatile oil. Oleo–resin components are considered to be the main active principles in ginger and documented pharmacological actions generally support the traditional uses. In addition, a number of other pharmacological activities have been documented, including hypoglycaemic, antihypercholesterolaemic, anti–ulcer and inhibition of prostaglandin synthesis, all of which require further investigation. The use of ginger as a prophylactic remedy against motion sickness is contentious. It seems likely that ginger may act by a local action on the gastro–intestinal tract, rather than by a centrally mediated mechanism.
References


50. Srivastava KC. Effect of onion and ginger consumption on platelet thromboxane production in humans. *Prostaglandins Leukot Essent Fatty...*


Ginkgo
Species (Family)

Ginkgo biloba L. (Ginkgoaceae)
Synonym(s)
Fossil Tree, Kew Tree, Maidenhair Tree
Part(s) Used

Leaf
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)
BP 2002\(^{(G71)}\)
Complete German Commission E\(^{(G3)}\)
Martindale 33rd edition\(^{(G67)}\)
Mills and Bone\(^{(G50)}\)
PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Ph Eur 2004\(^{(G72)}\)
USP26/NF21\(^{(G73)}\)
WHO volume 1 1999\(^{(G63)}\)
Legal Category (Licensed Products)

Ginkgo is not included in the GSL.\(^{(G37)}\)
Constituents
See References 1 and 2, and General Reference G64.

Leaf

**Amino acids**
6-Hydroxykynurenic acid (2–carboxy-4–one–6–hydroxyquinoline), a metabolite of tryptophan.\(^{(3-5)}\)

**Flavonoids**
Dimeric flavones (e.g. amentoflavone, bilobetin, ginkgetin, isoginkgetin, sciadopitysin);\(^{(6)}\) flavonols (e.g. quercetin, kaempferol) and their glycosides\(^{(3,7)}\) and coumaroyl esters.

**Proanthocyanidins**
Terpenoids Sesquiterpenes (e.g. bilobalide), diterpenes (e.g. ginkgolides A, B, C, J, M, which are unique cage molecules,\(^{(8,9,G48)}\) and triterpenes (e.g. sterols).

**Other constituents**
Benzoic acid, allergenic ginkgolic acids, 2–hexenal, polyprenols (e.g. di-trans-poly-cis-octadecaprenol), sugars, waxes,\(^{(1)}\) a peptide.\(^{(10)}\)

Seeds

**Alkaloids**
Ginkgotoxin (4-O-methylpyridoxine).\(^{(11)}\)

**Amino acids**
Cyanogenetic glycosides Allergenic ginkgolic acids.

Ginkbilobin.\(^{(12)}\)

Standardised extracts of *G. biloba* leaves are standardised on the content of ginkgo flavonoid glycosides (22–27%; determined as quercetin, kaempferol andisorhamnetin), and terpene lactones (5–7%; comprising around 2.8–3.4% ginkgolides A, B and C, and 2.6–3.2% bilobalide, and less than 5 ppm ginkgolic acids).\(^{(G3 G56)}\)
Food Use

*Ginkgo biloba* is not used in foods.
Herbal Use

Ginkgo has a long history of medicinal use, dating back to 2800BC. Traditional Chinese medicine used the seeds (kernel/nuts) for therapeutic purposes. The seed is used in China as an antitussive, expectorant and anti–asthmatic, and in bladder inflammation.\(^{(1,11,G50)}\) In China, the leaves of *Ginkgo biloba* were also used in asthma and in cardiovascular disorders,\(^{(1)}\) although the leaves have little history of traditional use in the West. Today, standardised concentrated extracts of *G. biloba* leaves are marketed in several European countries, and are used in cognitive deficiency, intermittent claudication (generally resulting from peripheral arterial occlusive disease), and vertigo and tinnitus of vascular origin (see Pharmacological Actions, Clinical studies).\(^{(G3 G32 G56 G63)}\)
Dosage

Cognitive deficiency

*Leaf extract*
120–240 mg dry extract orally in two or three divided doses.\(^{(G3)}\)

Peripheral arterial occlusive disease and vertigo/tinnitus

*Leaf extract*
120–160 mg dry extract orally in two or three divided doses.\(^{(G3)}\)

Clinical trials of standardised extracts of *G. biloba* leaves (EGb 761, Willmar Schwabe GmbH and LI 1370, Lichtwer Pharma GmbH) in patients with cognitive deficiency have generally used oral doses ranging from 120 to 240 mg daily, usually for 8–12 weeks, although some studies have continued treatment for up to 24 or 52 weeks.\(^{(G56)}\) Clinical trials in peripheral arterial occlusive disease used oral doses of 120–160 mg extract daily for 3–6 months.\(^{(G56)}\)
Pharmacological Actions

In vitro and animal studies

There is a vast literature describing basic scientific research relating to the effects of ginkgo. Several pharmacological activities have been documented for ginkgo leaf extracts and/or their constituents. These include effects on behaviour, learning and memory, cardiovascular activities, effects on blood flow and antioxidant activity. The most important active principles of ginkgo extract include the ginkgo flavonoid glycosides and the terpene lactones.\(^1\)

Ginkgo has been described as having polyvalent action, i.e. the combined activity of several of its constituents is likely to be responsible for its effects.\(^13\)

The pharmacological activities of ginkgo have been reviewed,\(^1,8,9,13–15\) and other texts bring together several studies in specific areas, e.g. neuroprotective effects.\(^16\) A summary of some of the literature on the in vitro and in vivo (animals) effects of ginkgo leaf is given below.

**Effects on behaviour, learning and memory**
The effects of a standardised extract of ginkgo leaf (EGb 761) on learning and memory, and on behaviour in relation to ageing and in recovery from brain injury, have been well studied.\(^13\) Animal models (rats and mice) designed to test aspects of learning and memory (e.g. acquisition and retention) have documented improvements in animals treated with oral, intraperitoneal or subcutaneous EGb 761, compared with controls.\(^13\) Studies involving rats reported improvements in acquisition and retention in older (24-month-old), but not younger (eight-month-old) rats. Other experiments involving rats of different ages have found that older rats (12- and 18-months old) showed improved performance in an eight-arm radial maze test following oral administration of EGb 761 30 or 60 mg/kg/day, whereas performance was stable among young rats (eight weeks old) following EGb 761 administration.\(^13\) EGb 761 200 mg/kg administered orally to rats aged more than 26 months old led to significant improvements in aspects of cognitive behaviour.\(^17\) In vivo studies have also shown that oral administration of EGb 761 (50 or 100 mg/kg/day for three weeks) to rats prevented the short-term memory-impairing effects of scopolamine administered intraperitoneally (0.125 mg/kg).\(^13\)

The anxiolytic effects of a range of doses (0.01–10 mg/kg) of combination preparations containing different mixture ratios of standardised extracts of ginkgo leaf and ginger root have been tested in rats using the elevated plus-maze test.\(^18\) Compared with controls, rats treated with the combination
preparation (mixture ratio of ginger extract to ginkgo extract, 2.5 : 1; 1 mg/kg, intragastrically) spent increased amounts of time in the open arms of the maze, whereas the behaviour of rats treated with preparations of a mixture ratio of 1 : 1 and 1 : 2.5 did not change.

Several studies have reported that treatment with EGB 761, compared with control, aids recovery of function following brain injury, as demonstrated by behavioural tests in rats who had undergone bilateral frontal lobotomy or septohippocampal deafferentation, and in rat models of cortical hemiplegia.\(^{13}\)

It has been suggested that the effects of EGB 761 in the experimental animal models described above may involve aspects of neuronal plasticity, e.g. neuronal regeneration.\(^{13}\) Studies in rats have investigated, for example, the effects of EGB 761 administration on expression of neurotrophins and apolipoprotein E, and on behavioural recovery, following entorhinal cortex lesions, and on regeneration of primary olfactory neurons following olfactory bulbectomy. Research investigating the effects of EGB 761 on neuronal plasticity has been summarised.\(^{1,13,19}\)

**Cardiovascular and haemorheological activities**

Studies investigating the molecular mechanisms that may contribute to the vasoregulatory (vasodilatation and vasoconstriction) effects of standardised ginkgo leaf extract (EGB 761) have been described.\(^{13,20}\) *In vitro* experiments using isolated rabbit aorta suggested that possible mechanisms include effects on cyclic-GMP phosphodiesterase, prostaglandin I\(_2\) and nitric oxide (NO). *Ex vivo* studies using isolated guinea–pig heart showed that EGB 761 led to a concentration–dependent increase in coronary blood flow. In studies involving isolated rat heart and in anaesthetised rabbits, EGB 761 administration has been reported to protect against myocardial ischaemia–reperfusion injury; the antioxidant and free–radical scavenging effects of EGB 761 (see below) may be important in this regard.\(^{13}\) One study using isolated rat hearts suggested that the cardioprotective effects were due to the terpenoid constituents of EGB 761, and that the mechanism was independent of direct free radical–scavenging activity.\(^{21}\) An *in vitro* study with endothelial cells suggested that the anti–ischaemic activity of EGB 761 may be due partly to the effects of the constituent bilobalide in protecting mitochondrial activity.\(^{22}\)

The effects of ginkgo leaf extract have been studied in normal rats and those with ischaemic brain damage with middle cerebral artery occlusion.\(^{23}\) Oral administration of ginkgo extract 100 mg/kg was reported to increase cerebral blood flow in normal rats, but the increase was less marked in rats with
cerebral artery occlusion.

*In vitro* studies using human blood cells have documented effects of EGb 761 on several haemo rheological parameters.\(^{(13)}\) For example, *in vitro*, EGb 761 normalised changes in erythrocyte viscosity and in the viscoelastic properties of the erythrocyte membrane induced by standard metabolic challenge (pH 6.8; 380 mosmol/L) in six human blood donors. In other *in vitro* experiments, EGb 761 protected against hydrogen peroxide–induced damage in human erythrocytes. In studies using blood from patients with circulatory disorders, incubation with EGb 761 was reported to decrease erythrocyte aggregation. *In vitro* experiments using human neutrophils have found that EGb 761 at a concentration of 10 μmol/L inhibits release of hydrogen peroxide from these cells. The effects of EGb 761 on inhibition of human platelet aggregation elicited by substances such as thrombin and collagen have been documented.\(^{(13)}\) A study using blood donated by healthy volunteers (\(n = 35\)) reported that a standardised extract of ginkgo leaf inhibited ADP- and collagen–induced platelet aggregation in platelet–rich plasma, gel–filtered platelets and in whole blood in a concentration–dependent manner.\(^{(24)}\)

**Platelet–activating factor antagonism**

Ginkgolides have been reported to competitively inhibit the binding of platelet–activating factor (PAF) to its membrane receptor.\(^{(8,9,25)}\) Ginkgolide B antagonises thrombus formation induced by PAF and, in guinea–pigs, it also induces a rapid curative thrombolysis. A protective effect is exerted by ginkgolides on PAF-induced broncho constriction and airway hyperactivity in immuno–anaphylaxis and in antigen–induced bronchial provocation tests. Oral or intravenous injection of ginkgolide B antagonises cardiovascular impairments and bronchoconstriction induced by PAF. Ginkgolide B does not appear to interfere with cyclooxygenase, but at an earlier step involving PAF receptors and phospholipase activation. Eosinophil infiltration occurs in asthma and in allergic reactions, the number of eosinophils increasing during late phase. Since PAF is a potent activator of eosinophil function, it has been argued that ginkgolide B may interfere with the late–phase response.\(^{(25)}\)

Pre–administration of ginkgolide B (1–5 mg/kg, intravenous) to rats has been reported to reduce PAF-induced decreases in diastolic and systolic arterial blood pressure in anaesthetised normotensive rats; this effect has also been reported in this animal model when ginkgolide B is administered shortly after PAF administration.\(^{(13)}\)

Ginkgolide B has also been documented to have some beneficial effects in
endotoxic shock; PAF is believed by some to be implicated in shock states. In anaesthetised guinea–pigs, intravenous administration of ginkgolide B (1 or 6 mg/kg) prior to injection of Salmonella typhimurium endotoxin reduced the initial rapid decrease in blood pressure, and intravenous administration of ginkgolide B during the prolonged phase of shock (1 hour after endotoxin administration) immediately and dose dependently reversed the decrease in blood pressure.(13) Other studies have found that ginkgolide B reduced arterial blood pressure in the secondary, but not the early, phase following administration of Escherichia coli endotoxin.(13)

**Antioxidant activity**
The free radical–scavenging effects and antioxidant activity of EGb 761 *in vitro* are well documented.(13) EGb 761 scavenges several reactive oxygen species, including hydroxyl, superoxide and peroxyl radicals.(13,26,27) In rat cerebellar neurons and cerebellar granule cells, ginkgo extract was reported to protect against oxidative stress induced by hydrogen peroxide, another reactive oxygen species.(28,29) In cultures of rat hippocampal cells, incubation with EGb 761 protected against cell death induced by β-amyloid, protected against toxicity induced by hydrogen peroxide, and blocked β-amyloid–induced events, such as accumulation of reactive oxygen species.(30) Bilobalide has also been documented to protect neurons against oxidative stress induced by reactive oxygen species *in vitro*. (31) Experiments in gerbils have suggested that the neuroprotective effects of gingko extract may be due to inhibition of nitric oxide formation.(32)

Other studies which have described neuroprotective effects of EGb 761 have suggested that anti oxidant activity may be involved.(16) *In vitro*, a standardised ginkgo leaf extract was found to inhibit photo–induced formation of cholesterol oxides in a concentration–dependent manner.(33)

Antioxidant activity has been documented for EGb 761 *in vivo*. In rats, treatment with EGb 761 increased the concentrations of circulating and cellular poly unsaturated fatty acids, and reduced erythrocyte cell lysis induced by hydrogen peroxide.(34) Also in rats, oral administration of EGb 761 was reported to increase activity of the enzymes catalase and superoxide dismutase in the hippocampus, striatum and substantia nigrum.(35) Other data collected in this study suggested a decrease in lipid peroxidation in rat hippocampus in EGb 761–treated rats. In another study in rats, EGb 761 (200 mg/kg/day for four weeks) protected against carbon tetrachloride–induced (1.5 mL/kg) liver damage, as determined by malondialdehyde concentrations (a breakdown product of lipid peroxidation).(36)

**Other activities**
In vivo studies have suggested that EGb 761 may protect against chemically induced carcinogenesis. In mice, oral administration of EGb 761 (150 mg/kg daily for two weeks), compared with control, was reported to reduce tumour multiplicity; however, the inhibitory effect was not statistically significant.(37) It was also reported that EGb 761–treatment reduced the cardiotoxicity of doxorubicin.

EGb 761 and ginkgolide B have been shown to inhibit peripheral–type benzodiazepine receptor (PBR) expression and cell proliferation in the human breast cancer cell line MDA-231, which is known to be rich in PBR.(38) By contrast, the proliferation of MCF-7 breast cancer cells, which are low in PBR, was not affected.

In rats, oral administration of standardised ginkgo leaf extract 300 mg/kg was shown to ameliorate nephrotoxicity induced by administration of gentamicin 80 mg/kg.(39)

Aqueous extracts of dried ginkgo leaves have been reported to inhibit monoamine oxidases (MAO) A and B.(40) A study investigating the effects of bilobalide on gamma–aminobutyric acid (GABA) concentrations and on glutamic acid decarboxylase activity in mouse brain found that GABA concentrations and glutamic acid decarboxylase activity were significantly higher in animals treated orally with bilobalide 30 mg/kg daily for four days.(41) However, there were no differences between treated and control mice with regard to glutamate concentrations.

Several in vivo studies have documented adaptive effects for EGb 761.(20)

A peptide isolated from the leaves of Ginkgo biloba has been reported to have antifungal activity against several fungi, including Pellicularia sasakii and Alternaria alternata.(10)

Three long–chain phenols, anacardic acid, bilobol and cardanol, isolated from seeds of G. biloba are active against Sarcoma 180 ascites in mice.(42)

Clinical studies

Pharmacokinetics

Data on the pharmacokinetics of standardised extracts of ginkgo leaf have been summarised.(1,15,G18 G21) Mean bio availabilities of ginkgolide A, ginkgolide B and bilobalide following oral administration of ginkgo extract 120 mg to fasting healthy volunteers were 80%, 88% and 79%, respectively. Food intake increased the time taken to reach peak concentration (suggesting slower absorption), but did not affect bioavailability.(1,G18) Peak
concentrations of ginkgolides A and B and bilobalide observed in fasting volunteers ranged from 16.5 to 33.3 ng/mL, and from 11.5 to 21.1 ng/mL in volunteers who had consumed food.\(^1\) Urinary excretion of ginkgolides A and B, and bilobalide, is around 70%, 50% and 30%, respectively, of the dose administered orally.\((G18)\)

**Therapeutic effects**

Most clinical trials of ginkgo have explored its effects in the treatment of cognitive deficiency or cerebral insufficiency,\(^{43,G56}\) a term used to describe a collection of symptoms thought to arise from an age–related reduction in cerebral blood flow. These symptoms include forgetfulness, poor concentration, poor perception, debilitation, dizziness, fatigue, sleep disturbances, listlessness, depressed mood, headache, mood swings, restlessness, tinnitus, anxiety, hearing loss and disorientation.\(^{43,G56}\) Several studies have tested the effects of standardised ginkgo leaf extracts on cognitive function in patients with Alzheimer’s disease\(^{44}\) and/or multi–infarct dementia.\(^{45}\) Both are conditions which share several symptoms (e.g. memory impairment) with cerebral insufficiency. Several other trials have explored the effects of ginkgo extracts on cognitive ability in individuals with no history of significant cognitive impairment. A few studies have explored the effects of ginkgo on tinnitus alone.\(^{46}\)

Clinical research with ginkgo extracts has also focused on effects in improving pain–free walking distance in patients with intermittent claudication/peripheral arterial occlusive disease.\(^{47,G56}\) Other studies have explored the effects of ginkgo in patients with chronic venous insufficiency, antidepressant–related sexual dysfunction, seasonal affective disorder (SAD), and symptoms of depression.

Almost all clinical trials of ginkgo have investigated the effects of the standardised ginkgo leaf extracts EGb 761 and LI 1370.

**Cognitive deficiency, dementia in Alzheimer’s disease, multi–infarct dementia**

A review of controlled clinical trials of ginkgo in patients with cerebral insufficiency identified 40 studies.\(^{43}\) Generally, trials tested oral doses of standardised extracts of ginkgo leaf of 120 mg daily administered for at least four to six weeks. Most trials reported significant results or positive (but not statistically significant) trends in favour of ginkgo, compared with control. However, it was reported that most trials were of poor methodological quality; only eight studies were considered to be well–conducted. All of these eight studies reported statistically significant results for ginkgo, compared with placebo. Nevertheless, further randomised, double–blind, controlled
trials involving larger numbers of patients were deemed necessary.\(^{(43)}\) Details of 39 controlled studies of the ginkgo extracts EGb 761 and LI 1370 have been summarised.\(^{(G56)}\) All but two\(^{(48,49)}\) of these studies were conducted before new guidelines for testing the efficacy of nootropic drugs were developed.\(^{(G56)}\) Details of the two studies that did meet the methodological criteria described in the guidelines are given below.

In a randomised, double-blind, placebo-controlled trial, after a four-week placebo run-in period, 216 patients with mild-to-moderate primary degenerative dementia of the Alzheimer type, or multi-infarct dementia, received standardised ginkgo leaf extract (EGb 761) 120 mg orally twice daily, or placebo, for 24 weeks.\(^{(48)}\) At the end of the study, data for 156 patients were eligible for analysis. There were significantly more responders to treatment (defined as a response to at least two of the three primary outcome measures – a psychopathological assessment, an assessment of cognitive performance and a behavioural assessment of activities of daily life) in the ginkgo group, compared with the placebo group (28% versus 10% of ginkgo and placebo recipients, respectively; \(p = 0.005\)). The difference was also statistically significant in an intention-to-treat analysis (23% versus 10% of ginkgo and placebo recipients, respectively; \(p = 0.005\)).

A randomised, double-blind, placebo-controlled study involved 327 patients with mild-to-severe dementia related to Alzheimer’s disease or multi-infarct dementia.\(^{(49)}\) Participants received standardised ginkgo leaf extract (EGb 761) 40 mg orally three times daily \((n = 166)\), or placebo \((n = 161)\), for 52 weeks, and underwent a battery of assessments at 12, 26 and 52 weeks. The primary outcome measures were the Alzheimer’s Disease Assessment Scale Cognitive Subscale (Adas-Cog), the Geriatric Evaluation by Relative’s Rating Instrument (GERRI) and the Clinical Global Impression of Change (CGIC). In an intention-to-treat analysis \((n = 309)\), ginkgo recipients scored significantly better than did placebo recipients on the Adas-Cog and the GERRI \((p = 0.04\) and \(p = 0.004\), respectively). A slight worsening on the CGIC was observed for both groups. The average end-points for the intention-to-treat analysis were 38.6 and 34.6 weeks for the ginkgo and placebo groups, respectively.

A systematic review of randomised, double-blind, placebo-controlled trials assessing the effects of standardised ginkgo leaf extracts on cognitive function in patients with Alzheimer’s disease, characterised according to recognised criteria, included four studies.\(^{(44)}\) These involved oral administration of ginkgo extract 120 or 240 mg daily for 12–26 weeks, and involved a total of 212 patients each in the ginkgo and placebo groups. A meta-analysis of the results of the four studies indicated a modest effect for
ginkgo, compared with placebo (difference of 3% on the Adas-Cog).

Another systematic review included nine randomised, double-blind, placebo-controlled trials of standardised ginkgo leaf extracts in patients with dementia of the Alzheimer type and/or multi-infarct dementia. The review included two studies described above. Studies generally involved the administration of oral doses of ginkgo extract 120 or 240 mg daily for 6–12 weeks, although two studies involved a 24-week or 52-week administration period. One study involved the administration of intravenous infusions of ginkgo extract 200 mg four times per week for four weeks. It was reported that, overall, the studies provided evidence to support the efficacy of standardised ginkgo leaf extracts in the symptomatic treatment of dementia. However, methodological limitations of several of the included studies (e.g. poorly defined inclusion and exclusion criteria and method of randomisation, treatment period less than six months, small sample sizes) were also emphasised. It was concluded that further studies are required to establish the benefits of ginkgo in dementia.

In a randomised, double-blind, placebo-controlled study, 60 elderly volunteers with mild-to-moderate, age-related cognitive dysfunction received oral ginkgo extract (GB-8; no further details provided) 40 mg, 80 mg or placebo, three times daily for three months. At the end of the study, for the 54 patients who completed, it was reported that memory function (as assessed by the Wechsler Memory Scale) improved significantly in the low-dose ginkgo group, compared with baseline values ($p = 0.016$), but that there was no significant improvement in the placebo or high-dose ginkgo groups, compared with baseline values. A significant decrease in diastolic blood pressure, compared with baseline values, was also reported for the low-dose ginkgo group ($p = 0.04$). This study, however, had methodological limitations (e.g. small sample size), and the report of the study did not include statistical analyses between groups.

A more methodologically rigorous randomised, double-blind, placebo-controlled trial involving patients with age-related impairment of memory and/or concentration assessed the effects of an alcohol/water extract of fresh leaves of ginkgo (drug extract ratio 1 : 4; total flavonoid glycosides 0.20 mg/mL, total ginkgolides 0.34 mg/mL). Participants received undiluted ginkgo extract ($n = 77$), diluted ginkgo extract (1 : 1 with placebo) ($n = 82$), or placebo ($n = 82$), 40 drops (1.9 mL) three times daily for 24 weeks. At the end of the treatment period, a check for blinding indicated that participants were unable to identify the treatment they received. There were no statistically significant differences between the three groups in subjective perceptions of memory and concentration, and in the following objective
measures: the Expended Mental Control Test (a measure of attention and concentration), and Rey test parts 1 and 2 (which measure short-term memory and learning curve, and long-term memory and recognition, respectively). However, a significant difference between groups was observed in the Benton test of visual retention–revised (a measure of short-term visual memory) – increases in baseline scores of 18, 26 and 11% were recorded for the high-dose ginkgo, low-dose ginkgo and placebo groups, respectively \((p = 0.0076)\).

In an open study, 18 elderly patients with ‘possible or probable’ Alzheimer’s disease were randomised to receive a single oral dose of tacrine 40 mg or a standardised extract of ginkgo leaf (SeGb; no further details provided) 240 mg in two separate sessions within three- to seven-day intervals.\(^{(52)}\) It was reported that both interventions induced pharmacological effects in the central nervous system (CNS), as assessed by quantitative pharmaco-electroencephalogram measurements. It should be noted that this was an uncontrolled study.

**Cognitive enhancement in healthy volunteers**

Ginkgo has been tested for its cognitive enhancing effects in healthy (i.e. cognitively intact) individuals in addition to investigations into its effects in patients with cognitive deficiency.

In a double-blind, placebo-controlled, cross-over study, 20 healthy volunteers aged 19–24 years received a standardised extract of ginkgo leaf (GK501) at doses of 120 mg, 240 mg and 360 mg.\(^{(53)}\) A battery of tests used to assess cognitive performance was carried out immediately before and at 1, 2.5, 4 and 6 hours after ginkgo administration. It was reported that with doses of ginkgo extract of 240 and 360 mg, there was a statistically significant improvement in ‘speed of attention’ (a measure of reaction time) from 2.5 hours up to 6 hours (the last measurement point) after ginkgo administration.

In a randomised, double-blind, placebo-controlled, crossover study, eight healthy female volunteers (mean age 32 years) were given a standardised extract of *G. biloba* leaf at doses of 120, 240 and 600 mg.\(^{(54)}\) One hour after treatment, volunteers undertook a series of psychological tests. Memory was found to be significantly improved with *G. biloba* leaf 600 mg, compared with placebo.

A randomised, double-blind, placebo-controlled, crossover trial involving 31 volunteers aged 30–59 years tested the effects of a standardised extract of ginkgo leaf (LI 1370) 50 mg three times daily, 100 mg three times daily,
120 mg each morning, 240 mg each morning, and placebo, each taken for two days followed by a washout period of at least five days.\(^{(55)}\) A battery of tests to assess memory and cognitive and psychomotor performance was carried out 30 minutes before ginkgo administration and then hourly for 12 hours. It was reported that there was a ‘marginally significant’ effect of treatment, compared with placebo, in a test assessing short–term memory, although the \(p\)-value given for this was greater than 0.05 (\(p = 0.053\)). Post–hoc analyses suggested that ginkgo extract 120 mg each morning was associated with better performance in this test than other doses of ginkgo extract (including ginkgo extract 50 mg three times daily) and placebo. There were no statistically significant effects of treatment on immediate and delayed word recall and choice reaction time.

Other studies have assessed the cognitive–enhancing effects of ginkgo extracts in older volunteers. In a randomised, double–blind, placebo–controlled trial, 48 cognitively intact individuals aged over 55 years received a standardised ginkgo leaf extract (EGb 761) 60 mg three times daily, or placebo, for six weeks.\(^{(56)}\) A battery of neuropsychological tests was carried out before treatment and at the end of the study. Ginkgo extract recipients experienced a significant improvement in tests assessing speed of processing abilities, compared with placebo recipients (\(p < 0.03\)). However, no statistically significant differences between the ginkgo extract and placebo groups were evident for tests assessing memory.

In a questionnaire survey, the effects of administration of a standardised ginkgo leaf extract (LI 1370) 120 mg daily for four months on the activities of daily living were assessed in volunteers aged 32–97 years (mean (SD) 68.9 (8.4) years).\(^{(57)}\) Volunteers were recruited via editorial in a magazine. Of 8557 initial respondents, 5028 were eligible for the survey. In total, 1000 volunteers (who were not currently using any ginkgo products) were said to be randomly allocated to receive ginkgo extract; all other respondents were allocated to the no–treatment control group, unless they had stated that they only wished to receive ginkgo. It was reported that ginkgo extract recipients achieved significantly better scores than the control group on a scale assessing ability to perform activities of daily living, self–assessment of ability to cope, and visual analogue scales for mood and sleep. However, for several reasons, the results of this study should be interpreted cautiously. For example, the study was carried out by post, therefore investigators did not meet participants at any time, the study was not truly randomised, the study was open, the control group did not receive placebo tablets, and there was no check on compliance.

**Tinnitus and hearing loss**
Tinnitus and hearing loss are two of the symptoms of dementia. Several studies have assessed the effects of ginkgo on these conditions alone.

A systematic review of randomised, controlled trials of ginkgo extracts in tinnitus included five studies – four studies compared ginkgo extracts with placebo, and one study compared ginkgo extract with conventional drugs. Three trials tested the standardised ginkgo leaf extract EGb 761; full details of other extracts tested in the other studies are not given in the review. The review concluded that, overall, the studies identified provided evidence to support ginkgo extracts as a treatment for tinnitus, but that further investigation was required to fully establish the benefits. Typically, at least two of the studies had methodological flaws.

A double-blind, controlled trial, published since the systematic review, tested the effects of a standardised ginkgo leaf extract (LI 1370) 50 mg, or placebo, three times daily for 12 weeks in 1121 individuals, aged 18–70 years, with tinnitus who were otherwise healthy. Participants were recruited via advertisements placed in the UK national press and in a British Tinnitus Association’s publication. The main outcome measure was participants’ self-assessment of tinnitus (loudness and ‘how troublesome’) before, during and after treatment, carried out via postal questionnaires and telephone calls. Participants were paired where possible (489 pairs, i.e. 978 of 1121 participants were matched) and then randomly allocated to active or placebo. At the end of the study, the results indicated that ginkgo extract (LI 1370) 50 mg three times daily was ‘no more effective than placebo in treating tinnitus’. The design of this study has been criticised. For example, participants did not have face-to-face contact with an investigator at any time during the study.

In a randomised, controlled trial, 28 patients with untreated sudden loss of hearing received intravenous infusions of 6% hydroxyethyl starch (HES), or intravenous and oral ginkgo extract, for 10 days. There were no statistically significant differences between the two groups in improvements in hearing. Further studies involving larger numbers of participants are required.

Peripheral arterial occlusive disease/intermittent claudication
The effects of standardised extracts of ginkgo have been investigated in patients with Fontaine stage II peripheral arterial occlusive disease. This condition is characterised by the onset of pain, as a result of oxygen deficit in the leg muscles, on walking distances greater than around 30–300 metres. The rationale for using ginkgo in this condition is for its effects in improving blood flow.
A meta-analysis of randomised, double-blind, placebo-controlled trials of ginkgo extract for the treatment of intermittent claudication included eight studies that assessed effects on walking distance.\(^{(47)}\) The trials involved a total of 415 patients who received a standardised extract of ginkgo leaf at doses of 120 or 160 mg daily, or placebo, for 6, 12 (one trial each) or 24 weeks. The pooled results from all trials indicated a statistically significant increase in pain-free walking distance for ginkgo-treated patients, compared with placebo recipients (weighted mean difference (WMD): 34 metres; 95% confidence intervals (CI): 26–43 metres). A similar result was obtained when results for the six studies of good methodological quality were pooled (WMD: 37 metres; 95% CI: 26–47 metres). It is questionable whether the extent of these increases in pain-free walking distance is clinically relevant.\(^{(47)}\)

**Chronic venous insufficiency and venous ulcers**

A randomised, double-blind, placebo-controlled trial assessed the protective effects of a combination preparation (Ginkgor Forte) containing a standardised extract of ginkgo (2.3%), troxerutine (48.85%) and heptaminol (48.85%) against venous wall injury in 48 female patients with chronic venous insufficiency.\(^{(61)}\) Ginkgor Forte 625 mg daily, or placebo, was given for four weeks. In total, 42 patients completed the study, but only 28 were included in the final analysis because of protocol violations. Circulating endothelial cell (CEC) count was used as a measure of injury to the vascular endothelium (CEC counts are raised in patients with chronic venous insufficiency.\(^{(61)}\) After four weeks’ treatment, CEC counts decreased significantly in both the treatment and placebo groups (by 14.5% and 8.4%, respectively), compared with baseline values \((p = 0.0021\) and \(p = 0.0146\), respectively). The mean change in CEC count after four weeks’ treatment was reported to be significantly greater for the treatment group, compared with the placebo group \((p = 0.039)\).

In another double-blind trial, 213 patients with chronic venous or mixed ulcers located at the malleolus (rounded protuberance on ankle joint) received ginkgo extract 160 mg daily, or placebo, together with standard care (elastic stockings, local dressings and cleansing of ulcers) for 12 weeks.\(^{(1)}\) At the end of the study, ginkgo extract recipients, compared with placebo recipients, showed a significant reduction in ulcer area.

**Asthma, PAF antagonism**

Intradermal injections of PAF induce a biphasic inflammatory response similar to that observed in sensitised individuals subjected to moderate doses of allergen. A single dose of a mixture of ginkgolides has been reported to antagonise this response.\(^{(62)}\) Oral administration of ginkgolides resulted in a reduction of eosinophil infiltration in atopic patients given intracutaneous
In a randomised, double-blind, crossover study, 80 and 120 mg capsules containing a standardised mixture of ginkgolides A, B and C (ratio of 40 : 40 : 20) were given as a single oral dose 2 hours before challenge by intradermal PAF/histamine. Both dose ranges inhibited flare which was maximal after 5 minutes. Within 15–30 minutes wheal volume was reduced, with the greatest effect being observed for the higher dose treatments. The protection was still present 8 hours after oral dosing. Similar inhibition of PAF was observed for platelet aggregation with single oral doses of 80 and 120 mg extract which were given 2 hours before blood withdrawal. The ginkgolide mixture given orally also blocked PAF-induced airway hyper-responsiveness.

Antagonism of the effects of PAF by a standardised mixture of ginkgolides was assessed in a double-blind, placebo-controlled crossover study in six healthy subjects aged 25–35 years. Wheal and flare responses to PAF examined 2 hours after ingestion of 80 mg and 120 mg of ginkgolide mixture were inhibited in a dose-related manner. Both doses significantly inhibited PAF-induced platelet aggregation in platelet-rich plasma.

A randomised, double-blind, crossover study involved patients with atopic asthma who were challenged with their specific dust or pollen antigen. After 6.5 hours, participants were subjected to a provocation test with acetylcholine so that the treatment of later stages of an asthma attack could be assessed. Mixed ginkgolide standardised extract, 40 mg three times daily, or placebo, were given during the three days before the test and a final single dose of 120 mg of extract was given 2 hours before the challenge. The results suggested that ginkgolides were effective in both the early phase and the late phase of airway hyperactivity.

A study involving six patients with a history of exercise-induced asthma assessed the effects of a specific PAF antagonist (BN 52063, a standardised mixture of ginkgolides A, B and C, ratio of 40 : 40 : 20) on the response to isocapnic hyperventilation with dry cold air. Participants were randomised to receive BN 52063 240 mg orally 2 hours before cold air challenge, 2.4 mg by metered dose inhaler 30 minutes before cold air challenge, or placebo. It was reported that oral BN 52063 did not reduce bronchoconstriction during challenge. A significant increase in airways resistance was observed after inhalation of BN 52063. In another study, six patients with a history of exercise-induced asthma received BN 52063 120 mg orally twice daily, 1 mg by spinhaler three times daily, or placebo, for three days, then, on the test day, 240 mg orally 3 hours before exercise challenge, 5 mg by spinhaler 1
hour before challenge, or placebo, respectively. With oral treatment, the prolonged reduction in peak expiratory flow was significantly attenuated ($p < 0.05$). (64)

**Antidepressant–related sexual dysfunction**

Two open, uncontrolled studies have explored the effects of ginkgo extract in sexual dysfunction associated with treatment with antidepressant drugs. (65,66) One of these studies involved 63 men and women who were receiving treatment with selective serotonin reuptake inhibitors (SSRIs), venlafaxine, nefazodone, bupropion, phenelzine or protriptyline, and experiencing sexual dysfunction, including decreased libido, erectile difficulties, delayed or inhibited orgasm and/or ejaculatory failure. (65) All participants received ginkgo extract 40 or 60 mg twice daily, titrated up to a maximum of 120 mg twice daily (average daily dose 207 mg), for four weeks. It was reported that sexual dysfunction was relieved, as assessed by clinical interview and self-report, in 91% and 76% of female and male participants, respectively. An open, prospective pilot study involved 14 patients with sexual dysfunction (severe or complete loss of libido, with or without inability to achieve or maintain erection, failure or delay in ejaculation, anorgasmia) associated with current treatment with antidepressant drugs, mainly SSRIs, but also hypericum (St. John’s wort) extract, amitriptyline and clomipramine. (66) Participants received a standardised extract of ginkgo 240 mg daily for six weeks. Assessment included an eight–item ‘sexual stress’ score, ‘sleep problems’ score, and the Hamilton Anxiety and Depression Scale scores. Among the 12 individuals who completed the study statistically significant improvements in scores were observed for anxiety at week 6 ($p < 0.05$) and sexual stress at weeks 3 and 6 ($p < 0.01$ for both), all compared with baseline values.

One of these studies (65) has been criticised for its methodological limitations and poor quality of the report. (67) Owing to their open, uncontrolled designs, neither study provides reliable evidence of the effects of gingko extract in antidepressant–related sexual dysfunction, as the observed effects cannot be reliably attributed to treatment with ginkgo.

**Other conditions**

A Cochrane review of the effects of ginkgo extract in age–related macular degeneration identified one randomised, controlled trial involving 20 patients. (68) The study design did not include blinding of the assessment of outcome. The review concluded that the effects of ginkgo extract in preventing progression of age–related macular degeneration had not yet been adequately assessed.
In a randomised, double-blind, controlled trial, 27 patients with seasonal affective disorder (SAD) received standardised extract of ginkgo leaf (Bio-Biloba, containing flavone glycosides 24 mg and terpene lactones 6 mg) one tablet twice daily \((n = 15)\), or placebo \((n = 12)\), for 10 weeks or until development of depression requiring treatment.\(^{(69)}\) All participants began the trial during October to December; assessments were carried out at baseline and on termination of study medication. Six of the 15 ginkgo recipients and two of the 12 placebo recipients terminated the study treatment because of emerging symptoms of SAD ('winter depression'). It was reported that this difference between groups was not statistically significant, according to Fisher’s exact test. Also, there were no statistically significant differences between groups on the Montgomery–Asberg Depression Rating Scale, the ATYP scale (for symptoms of hypersomnia, hyperphagia and carbohydrate craving) and on self-assessed symptoms (energy, tiredness, appetite, carbohydrate craving, depressed mood). The results of this study cannot be considered definitive because of the small sample size of the study and other limitations.\(^{(69)}\)

An open pilot study explored the effects of a standardised dried extract of ginkgo leaf (LI 1370) on cognitive performance in patients with major depression who were receiving trimipramine 200 mg daily.\(^{(70)}\) Eight participants received trimipramine for six weeks, and ginkgo extract 120 mg twice daily from day 8 to day 35 of the study. The data for these individuals were compared with those for eight age- and sex–matched controls who received trimipramine only for six weeks. At the end of the study, there was a significantly higher response rate (determined by a reduction of 50% or more in baseline Hamilton Depression Rating Scale scores) among patients who received trimipramine only, compared with those who received trimipramine plus ginkgo extract \((p \leq 0.05)\). However, duration of illness before enrolment in the study was reported to be significantly longer for ginkgo extract recipients than for controls \((p \leq 0.05)\). It was claimed that the ginkgo extract group, compared with the control group (trimipramine only), had improved sleep efficiency and augmented non-REM (random eye movement) sleep, although \(p\)-values were not given. Furthermore, by contrast, sleep time was found to be significantly prolonged in participants receiving trimipramine only. The design of the study, together with conflicting results, do not allow any definitive conclusions regarding the effects on sleep of ginkgo extract in addition to trimipramine treatment.

A placebo-controlled trial involving 165 women with premenstrual syndrome explored the effects of standardised ginkgo leaf extract (EGb 761) 160 mg daily, or placebo, taken from day 16 of the menstrual cycle to day 5 of the following cycle, for two cycles.\(^{(71)}\) Both groups experienced improvements in
symptoms, compared with baseline values, although ginkgo recipients, compared with placebo recipients, were reported to experience significantly greater improvements in breast tenderness (as evaluated by the physician).

The effects of ginkgo extract have been explored in patients with schizophrenia. The rationale for this is based on the theory that excess free radical formation may occur in patients with schizophrenia, since superoxide dismutase (SOD) concentrations have been reported to be higher in certain tissues in such patients. (SOD is an enzyme that detoxifies superoxide radicals.) In a double-blind, placebo-controlled trial, 82 inpatients with chronic schizophrenia (illness for at least five years) were ‘divided randomly’ to receive haloperidol 0.25 mg/kg daily, with or without ginkgo extract 360 mg daily, for 12 weeks. At the end of the study, mean SOD concentrations (expressed as ng/mL haemoglobin) were reported to be significantly lower, compared with baseline values, for ginkgo recipients (mean (SD): 815.8 (697.8) and 596.7 (148.3), respectively; \( p = 0.021 \)). In participants who received haloperidol only, mean (SD) SOD concentrations fell from 780.4 (605.4) at baseline to 617.6 (189.7) at the end of the study, although this decrease was reported to be statistically non-significant. A between-group comparison was not reported. Mean (SD) SOD concentrations in a group of 30 age- and sex-matched healthy volunteers were 515.8 (70.4).

The effects of a preparation comprising *Ginkgo biloba* dimeric flavonoids in a 1:2 complex with phosphatidylcholine (GBDF-Phytosome) on the microcirculation of the skin have been investigated using various techniques including, infrared photo-pulse plethysmography, laser doppler flowmetry, high-performance contact thermography and computerised videothermography. In a controlled study, small numbers of healthy individuals and volunteers with acrocyanosis or cellulitis were treated with 0.5 mL of a cream (oil-in-water emulsion) containing 3% GBDF-Phytosome or an oil-in-water emulsion of 2% phosphatidylcholine (control). Participants who received the GBDF-Phytosome preparation were reported to experience significant increases in capillary blood flow and skin temperature, compared with baseline values, whereas no significant changes were observed for the control group. No between-group comparisons were reported. These preliminary findings suggest that the effects of this preparation on skin microcirculation may deserve further investigation.

A phase II (open, uncontrolled) study explored the effects of standardised ginkgo leaf extract (EGb 761) given in combination with 5-fluorouracil (5-FU) in 44 patients with advanced progressive colorectal cancer who had previously received 5-FU. The rationale for including ginkgo extract in the
regimen was based on its reputed ability to increase local blood flow. Thus, it was hypothesised that ginkgo extract might ‘enhance local tumour blood flow and thus improve the distribution of 5-FU’. In the study, participants received ginkgo extract 350 mg in 250 mL saline intravenously over 30 minutes on days 1–6, and 5-FU 500 mg/m² in 250 mL saline intravenously over 30 minutes on days 2–6. The regimen was repeated every three weeks until recurrence of tumour progression. Data from 32 patients who had received at least two courses of treatment were eligible for analysis. Of these patients, 69% experienced progression of disease, 25% experienced no change, and 6.3% (n = 2) were in partial remission. (74)
Side-effects, Toxicity

The safety and toxicity of ginkgo have been reviewed. Available data indicate that standardised extracts of ginkgo leaf are well tolerated when used at recommended doses. Adverse effects are uncommon. A postmarketing surveillance study involving 10,815 patients who received a standardised extract (LI 1370) of ginkgo leaf reported that the frequency of adverse effects was 1.7%. Adverse effects reported with standardised extracts of ginkgo leaf are generally mild, and include nausea, headache, gastrointestinal upset and diarrhoea; allergic skin reactions occur rarely.

A systematic review of nine randomised, double-blind, placebo-controlled trials of standardised ginkgo leaf extracts in patients with dementia of the Alzheimer type and/or multi-infarct dementia concluded that, overall, the frequency of adverse effects reported for ginkgo was not markedly different than that for placebo. The largest trial included in this review involved 327 patients with mild-to-severe dementia related to Alzheimer’s disease or multi-infarct dementia who received standardised ginkgo leaf extract (EGb 761) 40 mg orally three times daily (n = 166), or placebo (n = 161), for 52 weeks. It was reported that there was no statistically significant differences between ginkgo and placebo in the number of participants reporting adverse events, or in the frequency and severity of adverse events. Of 188 adverse events reported during the study, 97 were reported by ginkgo recipients and 91 by placebo recipients. However, clinical trials generally only have the statistical power to detect common, acute adverse effects. Similar findings were reported in another systematic review/meta-analysis which included eight randomised, double-blind, placebo-controlled trials of ginkgo extract for the treatment of intermittent claudication, involving a total of 415 patients who received standardised extract of ginkgo leaf at doses of 120 or 160 mg daily, or placebo, for up to 24 weeks. Five of the eight studies included reported (rarely) mild, transient adverse events occurring in ginkgo recipients; the remaining three studies, comprising almost 50% of the total number of patients, did not report any adverse events.

There are isolated reports of bleeding associated with ingestion of Ginkgo biloba extract. One report describes a 70-year-old man who experienced spontaneous bleeding from the iris into the anterior chamber of the eye one week after he began taking standardised ginkgo extract 80 mg daily. A 61-year-old man who had taken ginkgo extract 120 mg or 160 mg daily for six months experienced a subarachnoid haemorrhage. Another report describes a 33-year-old woman who began experiencing increasingly severe headaches, as well as double vision and nausea and vomiting, over several
months. During the course of investigations, it was revealed that she had been consuming standardised ginkgo extract 120 mg daily for two years. Her symptoms improved, although her headaches were not entirely relieved, after evacuation of bilateral subdural haematomas which were identified following an MRI scan. On stopping ginkgo extract, her prolonged bleeding time was reduced, and on follow-up she was symptom-free. A causal relationship between ginkgo ingestion and bleeding in these cases has not been definitively established.

There is a report of acute myoglobinuria in a 29-year-old man who was a regular weight-trainer and who had been taking a combination preparation containing extracts of ginkgo (200 mg), guarana (Paullinia cupana, 500 mg) and kava (Piper methysticum, 100 mg). The man was admitted to an intensive care unit with severe muscle pain and blood creatine kinase and myoglobin concentrations of 100 500 IU/L (normal values: 0–195) and 10 000 ng/mL (normal values: 0–90), respectively. Signs and symptoms subsided within six weeks. The relevance, if any, of ginkgo ingestion to the man’s condition, is unclear.

Contact or ingestion of the fruit pulp has produced severe allergic reactions including erythema, oedema, blisters and itching. The seed contains the toxin 4-O-methylpyridoxine which is reported to be responsible for ‘gin-nan’ food poisoning in Japan and China. The main symptoms are convulsion and loss of consciousness and lethality is estimated in about 27% of cases in Japan, infants being particularly vulnerable.
Contra–indications, Warnings

The fruit pulp has produced severe allergic reactions and should not be handled or ingested. The seed causes severe adverse effects when ingested.

Ginkgo extract should only be used with caution in patients taking anticoagulant or antiplatelet agents.

Pregnancy and lactation

No studies appear to have been reported on the effects of *G. biloba* leaf extracts or ginkgolides in pregnant or lactating women. In view of the many pharmacological actions documented and the lack of toxicity data, use of ginkgo during pregnancy and lactation should be avoided.
Pharmaceutical Comment

There is a vast scientific literature describing the pharmacological effects of ginkgo leaf extracts and their constituents. These data provide some supporting evidence for the modern clinical uses of standardised ginkgo leaf extracts. Also, standardised ginkgo leaf extracts are among the herbal preparations that have undergone most extensive clinical investigation. The effects of ginkgo extracts in dementia have been tested clinically mostly in trials involving patients with cognitive deficiency, Alzheimer’s disease and/or multi-infarct dementia. Some high-quality studies involving patients with dementia have reported significant beneficial effects for standardised ginkgo leaf extracts. However, systematic reviews/meta-analysis of all relevant randomised, double-blind, placebo-controlled trials have reported modest effects for ginkgo extract, compared with placebo, and have concluded that further high-quality studies are required to establish the benefits of ginkgo in dementia. Small randomised, double-blind, placebo-controlled trials investigating the cognitive enhancing effects of ginkgo extracts in healthy volunteers have reported conflicting results. Further study is required to determine whether ginkgo extracts are of value in cognitively intact individuals. The effects of ginkgo extract in patients with tinnitus have not been definitively established by trials carried out to date. A meta-analysis of trials of standardised ginkgo leaf extract in peripheral arterial occlusive disease found that ginkgo significantly improved pain-free walking distance, although the clinical relevance of the extent of improvement is questionable.

Generally, the intended uses of ginkgo are not suitable for self-medication.
References

See also General References G3 G5 G18 G21 G29 G31 G32 G36 G43 G50 G54 G56 G63 G64.

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Species (Family)

Eleutherococcus senticosus (Rupr. & Maxim.) Maxim. (Araliaceae)
Synonym(s)

_Acanthopanax senticosus_, Devil’s Shrub, Eleuthero, _Hedera senticosa_, Siberian Ginseng, Touch-Me-Not, Wild Pepper
Part(s) Used

Root
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

BP 2002\(^{(G71)}\)

Complete German Commission E\(^{(G3)}\)

Martindale 33rd edition\(^{(G67)}\)

Mills and Bone\(^{(G50)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)

Ph Eur 2004\(^{(G72)}\)
Eleutherooccus ginseng is not included in the GSL.\(^{(G37)}\)
Constituents

See References 1 and 2, and General Reference G6.

Eleutherosides A–M
Heterogeneous group of compounds including sterol(A), phenylpropanoid(B), coumarin (B1, B3), monosaccharide(C), and lignan(B4,D,E) structural types; many present as glycosides. Characterised eleutherosides include daucosterol(A), syringin(B), isofraxidin glucoside(B1), (−)-sesamin(B4), methyl-α-D-galactoside(C), (−)-syringaresinol glucoside(D), acanthoside D(E) and hedera–saponin B(M).

Carbohydrates
Polysaccharides (glycans); some have been referred to as eleutherans. Galactose, glucose, maltose, sucrose.

Some of the additional documented constituents represent aglycones of the eleutherosides, namely β-sitosterol, isofraxidin, (−)-syringaresinol and sinapyl alcohol.

Phenylpropanoids
Caffeic acid and ester, coniferyl aldehyde.

Terpenoids
Oleanolic acid.

Volatile oils
0.8%. Individual components not documented.
Food Use

Eleutherococcus ginseng is not used in foods.
Eleutherococcus ginseng does not have a traditional herbal use in the UK, although it has been used for many years in the former Soviet Union. Like Panax ginseng, Eleutherococcus ginseng is claimed to be an adaptogen in that it increases the body’s resistance to stress and builds up general vitality.
Dosage

Dry root 0.6–3 g daily for up to one month has been recommended.\textsuperscript{(G6 G49)} Russian studies in healthy human subjects have involved the administration of an ethanolic extract in doses ranging from 2 to 16 mL one to three times daily, for up to 60 consecutive days.

Doses in non–healthy individuals ranged from 0.5 to 6.0 mL one to three times daily for up to 35 days. In both groups, multiple dosing regimens were separated by an extract–free period of two to three weeks.\textsuperscript{(1)}
Pharmacological Actions

The adaptogenic properties of Eleutherococcus ginseng have been extensively investigated in the countries of the former USSR. Pharmacological studies on extracts of Eleutherococcus ginseng started in the 1950s and have been primarily reported by two groups of Russian scientists. In 1962, a 33% ethanolic extract of *Eleutherococcus senticosus* was approved for human use by the Pharmacological Committee of USSR Ministry of Health, and in 1976 it was estimated that some three million people were regularly using this extract.¹

A review by Farnsworth et al.¹ describes the chemistry and toxicity of *Eleutherococcus ginseng* and documents results of *in vitro*, *in vivo* and human studies involving the oral administration of an ethanolic extract.

The majority of literature on *Eleutherococcus ginseng* has been published in the Russian language and therefore great difficulty is encountered in obtaining translations.¹ This monograph will draw mainly on data included in the Farnsworth review as well as on more recent papers that have been published in English. When used in this monograph, ‘ginseng’ will refer to *Eleutherococcus ginseng* unless indicated otherwise.

**In vitro and animal studies**

**Hypo/hyperglycaemic activity**
Hypo/hyperglycaemic activity has been documented in both normal animals and in those with induced hyperglycaemia (rabbit, mouse), but with little effect on alloxan–induced hyperglycaemia (rat).¹⁴ Hypoglycaemic activity (mice, intraperitoneal injection) of an aqueous ginseng extract has been attributed to polysaccharide components termed eleutherans A–G.⁵

**Central nervous system effects**
Sedative actions (rat, mouse), CNS-stimulant effects (intravenous/subcutaneous injection, rabbit), and a decrease/increase in barbiturate sleeping time has been reported.¹⁶

**Immunostimulant, antitoxic actions**
Increased resistance to induced listeriosis infection (mouse, rabbit) with prophylactic ginseng administration and reduced resistance with simultaneous administration, stimulation of specific antiviral immunity (guinea–pig, mouse), regulation of complement titre and lysozyme activity post immunisation have been documented.¹ In addition, protection against cardiac glycoside (intravenous injection, frog), diethylglycolic acid (mouse)
and alloxan (rat) toxicity has also been described.\(^{(1)}\) Immune stimulant effects have been reported for polysaccharide components, together with an ability to lessen thioacetamide, phytohaemagglutinin and X-ray toxicity, and to exhibit antitumour effects.\(^{(5)}\) Immunostimulant activity *in vitro* (using granulocyte, carbon clearance and lymphocyte–transformation tests) has been documented for high molecular weight polysaccharide components.\(^{(7,8)}\)

**Effects on overall performance**
A beneficial action on parameters indicative of stress (rat) and on overall work capacity (mouse) has been reported,\(^{(1)}\) although a lack of adaptogenic response has also been reported in mice receiving various ginseng infusions (Siberian, Korean and American).\(^{(9,10)}\) In one study, mice receiving a commercial concentrated extract of eleutherococcus ginseng were noted to exhibit significantly more aggressive behaviour.\(^{(9)}\) Ginseng is claimed to result in a more economical utilisation of glycogen and high–energy phosphorus compounds, and in a more intense metabolism of lactic and pyruvic acids during stress.\(^{(1)}\) It has been claimed that the adaptogenic effect of ginseng involves regulation of energy, nucleic acid, and protein metabolism in tissues.\(^{(1)}\)

**Steroidal activity**
Gonadotrophic activity in immature male mice (intraperitoneal injection), oestrogenic activity in immature female mice, and an anabolic effect in immature rats (intraperitoneal injection) has been reported.\(^{(1)}\) *In vitro* studies have reported that ginseng extracts bind to progestin, mineralocorticoid, glucocorticoid and oestrogen receptors.\(^{(1)}\)

**Cardiovascular activity**
3,4-Dihydroxybenzoic acid (DBA) has been identified as an anti–aggregatory component in eleutherococcus ginseng.\(^{(3)}\) Compared with aspirin, activity of DBA was comparable versus collagen- and ADP-induced platelet aggregation, but less potent versus arachidonic acid–induced platelet aggregation.\(^{(3)}\) Anti–oedema and anti–inflammatory actions (intravenous injection, mouse), have also been described.\(^{(1)}\)

**Effect on reproductive capacity**
Ginseng has been reported to improve the reproductive capacity of bulls and cows, and to have no adverse effects on the various blood parameters (haemoglobin, total plasma protein, albumin and globulin, protein coefficient) measured.\(^{(1)}\)

Other actions documented for ginseng include the stimulation of liver
regeneration in partially hepatectomised mice,\(^{(1)}\) an increase in catecholamine concentrations in the brain, adrenal gland and urine,\(^{(1)}\) a variable effect on induced hypothermia (rabbit, rat, mouse),\(^{(1)}\) and \textit{in vitro} inhibition (66%) of hexobarbitone metabolism.\(^{(6)}\)

**Clinical studies**

In Russia, ginseng extract has been administered orally to more than 4300 human subjects in studies involving either healthy or non-healthy individuals.\(^{(1)}\)

**Administration to healthy subjects**

These studies were designed to investigate the adaptogenic effects of ginseng and measured parameters such as the ability of humans to withstand adverse conditions (heat, noise, motion, workload increase, exercise, decompression), improvement in auditory disturbances, quality of work under stress conditions and in athletic performance, and increase in mental alertness and work output.\(^{(1)}\) The studies involved more than 2100 subjects and included both male and female subjects ranging in age from 19 to 72 years. Doses ranged from 2 to 16 mL of an ethanolic extract (33%), administered orally one to three times daily, for periods of up to 60 consecutive days. Multiple dosing regimens usually involved a two- to three-week interval between courses.\(^{(1)}\) For many of the studies, it is unclear whether ginseng had a beneficial effect. However, ginseng was found to exert favourable effects in a number of situations including ability to perform physical labour, quality of proofreading, adaptation to a high-temperature environment, speed and quality of work by radiotelegraphers in noisy conditions, resistance to hypoxaemia and physical burdens in skiers, ability to withstand conditions designed to induce motion sickness, capillary resistance, haematological parameters in blood donors, and number of days lost to sickness amongst factory workers. Ginseng was also reported to increase excretion of vitamins B\(_1\), B\(_2\) and C given concurrently with ginseng. On its own, ginseng did not affect the excretion of water-soluble vitamins.

**Administration to non-healthy subjects**

These studies involved more than 2200 subjects with various ailments and included both males and females ranging in age from 19 to 60 years. Ginseng doses ranging from 0.5 to 6 mL were administered orally between one to three times daily for up to 35 days, with as many as eight courses employed. Multiple dosing regimens involved a two- to three-week ginseng-free interval in between courses.\(^{(1)}\) A favourable effect was noted in atherosclerosis (although treatment was stated to be less effective in patients with high blood pressure), acute pyelonephritis, various forms of diabetes mellitus (although...
no marked effect was noted in another study), hypertension and hypotension (tendency to normalisation), acute craniocerebral trauma, various types of neuroses, rheumatic heart disease (reduced blood coagulation properties), chronic bronchitis, and in children with abating forms of pulmonary tuberculosis.\(^1\) An increase in the working capacity of six males, in a single blind crossover study using placebo and no treatment as comparators, has been reported for a 33\% ethanolic ginseng extract.\(^11\) The observed increase in working capacity was partially attributed to an improvement in bodily oxygen metabolism, reflected by the increase in all four measured parameters (oxygen uptake, oxygen pulse, total work and exhaustion time).\(^12\)

**Immunostimulant activity**

A strong immunomodulatory effect has been documented for an ethanolic extract of ginseng, in a placebo–controlled double–blind study using healthy volunteers.\(^13\) A significant increase in the total lymphocyte count, especially in the T lymphocyte cells, was noted in the ginseng-treated group who received a daily dose of 30–40 mL extract (eleutheroside B 0.2\% w/v). Specificity of action on the lymphocytes was confirmed by the fact that neither granulocyte or monocyte levels were significantly altered.\(^13\)
Side–effects, Toxicity

No side–effects were documented from Russian studies involving more than 2100 healthy subjects.\(^1\) Studies involving patients with various ailments have reported a few side–effects: insomnia, shifts in heart rhythm, tachycardia, extrasystole and hypertonia in some atherosclerotic patients; headaches, pericardial pain, palpitations, and elevated blood pressure in 2 of 55 patients (at high dose level) with rheumatic heart disease; insomnia, irritability, melancholy and anxiety in hypochondriac patients receiving higher doses of extract; hypersensitivity reaction (symptoms unspecified) in stressed individuals.\(^1\) Hypertension and mastalgia have been documented as side–effects of ginseng (species unknown).\(^{11}\)

Results of various animal toxicity studies have indicated ginseng to be non–toxic.\(^1\) Many species have been exposed to extracts including mice, rats, rabbits, dogs, minks, deer, lambs, and piglets.\(^1\) Documented acute oral LD\(_{50}\) values for various preparations include: 23 mL/kg and 14.5 g/kg (mice), and greater than 20 mL/kg (dogs) for a 33% ethanolic extract\(^{1,4}\); 31 g/kg (mice) for the powdered root; greater than 3 g/kg (mice) for an aqueous aqueous (equivalent to 25 g dried roots/kg).\(^4\) No deaths occurred in mice administered single 3 g/kg doses of a freeze–dried aqueous extract.\(^{12}\) Symptoms observed in dogs receiving 7.1 mL/kg doses of the ethanolic extract (sedation, ataxia, loss of righting reflex, hypopnoea, tremors, increased salivation and vomiting) were attributed to the ethanol content of the extract.\(^1\) A chronic toxicity study reported no toxic manifestations or deaths in rats fed 5 mL/kg ethanolic extract for 320 days.\(^1\)

Teratogenicity studies in male and female rats, pregnant minks, rabbits and lambs have reported no abnormalities in the offspring and no adverse effects in the animals administered the extracts. Premature death in parent female rabbits fed 13.5 mL/kg ethanolic extract daily was attributed to ethanol intoxication.\(^1\)

Mutagenicity studies using *Salmonella typhimurium* TA100 and TA98, and the micronucleus test in mice have reported no activity for ginseng.\(^{14}\) Differences in various serum biochemical parameters have been reported between test (ginseng) and control groups.\(^{14}\) Parameters affected included alkaline phosphatase and gamma-glutamyl transferase enzymes (increased), serum triglycerides (decreased), and creatinine and blood urea nitrogen (increased).\(^{14}\) No pathological changes were found in rats receiving a ginseng extract.\(^{14}\)
Contraindications, Warnings

It has been stated that ginseng should be avoided by individuals who are highly energetic, nervous, tense, hysterical, manic or schizophrenic, and that it should not be taken with stimulants, including coffee, antipsychotic drugs or during treatment with hormones.\(^{(11)}\) In view of documented pharmacological actions, ginseng may interfere with a number of therapies including cardiac, anticoagulant, hypoglycaemic and hypo/hypertensive. Ginseng is stated to be unsuitable for individuals with high blood pressure (180/90 mmHg or greater)\(^{(1)}\) and has been advised to be avoided by premenopausal women.\(^{(11)}\)

Russian recommendations advise that healthy people under the age of 40 should not use ginseng and that middle-aged people can be treated with small doses of ginseng on a daily basis.\(^{(11)}\) Individuals considered suitable to use ginseng are recommended to abstain from alcoholic beverages, sexual activity, bitter substances and spicy foods.\(^{(11)}\)

In general, long-term use of ginseng is not recommended and one author has documented that the main side-effect of prolonged use manifests as an inflamed nerve, frequently the sciatic, which then causes muscle spasm in the affected area.\(^{(11)}\) Human studies involving long-term administration of ginseng have involved ginseng-free periods of 2–3 weeks every 30–60 days.

Pregnancy and lactation

Teratogenicity studies in various animal species have not reported any teratogenic effects for ginseng. However, in view of the many pharmacological actions documented for ginseng, and the general recommendation that it should not be used by premenopausal women, the use of ginseng during both pregnancy and lactation should be avoided. It is unknown whether the pharmacologically active constituents in ginseng are secreted in the breast milk.
Pharmaceutical Comment

Phytochemical studies have revealed that there is no one constituent type that is characteristic of Eleutherococcus ginseng. Studies have shown that components thought to represent the main active constituents ('eleutherosides') consist of a heterogeneous mixture of common plant constituents. Since the 1950s, many studies (animal and human) have been carried out in Russia, and more recently in Western countries, to investigate the reputed adaptogen properties of Eleutherococcus ginseng. An adaptogen is a substance that is defined as having three characteristics, namely lack of toxicity, non-specific action, and a normalising action.\(^1\) Results of numerous studies in animals and humans seem to support these three criteria for Eleutherococcus ginseng, although pharmacological explanations for the observed actions are less well understood.\(^1\) As with Panax ginseng, Eleutherococcus ginseng has been shown to possess a wide range of pharmacological activities. Consequently, it should be used with appropriate regard to traditional guidelines that have been drawn up in China and Russia.
References


Species (Family)

Various *Panax* species (Araliaceae) including:

i. *Panax ginseng* Meyer

ii. *Panax quinquefolius* L.

iii. *Panax notoginseng* (Burkh.) Hoo & Tseng
Synonym(s)


ii. American Ginseng, Sanchi Ginseng and Tienchi Ginseng

iii. American Ginseng, Five-Fingers, Sang and Western Ginseng
Part(s) Used

Root. White ginseng represents the peeled and sun-dried root whilst red ginseng is unpeeled, steamed and dried.
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

BP 2002\(^{(G71)}\)

Complete German Commission E\(^{(G3)}\)

Martindale 33rd edition\(^{(G67)}\)

Mills and Bone\(^{(G50)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)

Ph Eur 2004\(^{(G72)}\)

USP26/NF21\(^{(G73)}\)
Legal Category (Licensed Products)

GSL\((G37)\)
Constituents
See General References G2 G6 G41 G64.

Terpenoids
Complex mixture of compounds (gin senosides or panaxosides) involves three aglycone structural types – two tetracyclic dammarane–type sapogenins (protopanaxadiol and protopanaxatriol) and a pentacyclic triterpene oleanolic acid–type. Different naming conventions have been used for these compounds. In Japan, they are known as ginsenosides and are represented by \( R_x \) where ‘x’ indicates a particular saponin. For example, \( R_a, R_{b-1}, R_c, R_d, R_{g-1} \). In Russia, the saponins are referred to as panaxosides and are represented as panaxoside X where ‘X’ can be A–F. The suffixes in the two systems are not equivalent and thus panaxoside A does not equal \( R_a \) but \( R_{g-1} \).\(^1\)

The saponin content varies between different *Panax* species. For example, in *P. ginseng* the major ginsenosides are \( R_{b-1}, R_c \) and \( R_{g-1} \) whereas in *P. quinquefolis* \( R_{b-1} \) is the only major ginsenoside.\(^1\)

Other constituents
Volatile oil (trace) mainly consisting of sesquiterpenes including panacene, limonene, terpineol, eucalyptol, \( \alpha \)-phellandrene and citral,\(^2\) sesquiterpene alcohols including the panasinsanols A and B, and ginsenol,\(^3,4\) polyacetylenes,\(^5,6\) sterols, polysaccharides (mainly pectins and glucans),\(^7\) starch (8–32%), \( \beta \)-amylase,\(^8\) free sugars, vitamins (\( B_1, B_2, B_{12}, \) panthotenic acid, biotin), choline (0.1–0.2%), fats, minerals.

The sesquiterpene alcohols are stated to be characteristic components of *Panax ginseng* in that they are absent from the volatile oils of other *Panax* species.\(^4\)
Ginseng is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that ginseng can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product. (G16)
**Herbal Use**

Ginseng is stated to possess thymoleptic, sedative, demulcent and stomachic properties, and is reputed to be an aphrodisiac. Traditionally, it has been used for neurasthenia, neuralgia, insomnia, hypotonia, and specifically for depressive states associated with sexual inadequacy.\(^{(G2 \ G6 \ G8 \ G64)}\)

Ginseng has been used traditionally in Chinese medicine for many thousands of years as a stimulant, tonic, diuretic and stomachic.\(^{(9)}\) Traditionally, ginseng use has been divided into two categories: short–term – to improve stamina, concentration, healing process, stress resistance, vigilance and work efficiency in healthy individuals, and long–term – to improve well–being in debilitated and degenerative conditions especially those associated with old age.
Dosage

Traditionally, dosage recommendations differ between the short-term use in healthy individuals and the long-term use in elderly or debilitated persons.

**Short-term (for the young and healthy)**
0.5–1.0 g root daily, as two divided doses, for a course generally lasting 15–20 days and with a root-free period of approximately two weeks between consecutive courses. Doses are recommended to be taken in the morning, 2 hours before a meal, and in the evening, not less than 2 hours after a meal.\(^9\)

**Long-term (for the old and sick)**
0.4–0.8 g root daily. Doses may be taken continuously.\(^1\)
Pharmacological Actions

In the 1950s, early studies on ginseng reported its ability to improve both physical endurance and mental ability in animals and humans.\(^{10}\) In addition, the ‘tonic’ properties of ginseng were confirmed by the observation that doses taken for a prolonged period of time increased the overall well-being of an individual, measured by various parameters such as appetite, sleep and absence of moodiness, resulting in an increased work efficiency. Furthermore, these effects were felt for some time after cessation of ginseng treatment.\(^{10}\)

In addition, gonadotrophic activity, slight anti-inflammatory activity and an effect on carbohydrate metabolism were noted.\(^{10}\) Since then, numerous studies have investigated the complex pharmacology of ginseng in both animals and humans. The saponin glycosides (ginsenosides/panaxosides) are generally recognised as the main active constituents in ginseng, although pharmacological activities have also been associated with non-saponin components.

The following sections on animal and human studies are intended to give an indication of the type of research that has been published for ginseng rather than to provide a comprehensive bibliography of ginseng research papers.

**In vitro and animal studies**

**Corticosteroid-like activity**

Many of the activities exhibited by Panax ginseng have been compared to corticosteroid-like actions and results of endocrinological studies have suggested that the ginsenosides may primarily augment adrenal steroidogenesis via an indirect action on the pituitary gland.\(^{11}\) Ginsenosides have increased adrenal cAMP in intact but not in hypophysectomised rats and dexamethasone, a synthetic glucocorticoid that provides positive feedback at the level of the pituitary gland, has blocked the effect of ginsenosides on pituitary corticotropin and adrenal corticosterone secretion.\(^{11}\) Hormones produced by the pituitary and adrenal glands are known to play a significant role in the adaptation capabilities of the body.\(^{12}\)

Working capacity is one of the indices used to measure adaptation ability and ginseng has been shown to increase the working capacity of rats following single (132%) and seven-day (179%) administration (intraperitoneal). Furthermore, seven-day administration of ginseng decreased the reduction seen in working capacity when the pituitary-adrenocortical system is blocked by prior administration of hydrocortisone.\(^{12}\)

**Hypoglycaemic activity**

Hypoglycaemic activity has been documented for ginseng and attributed to
both saponin and polysaccharide constituents. In vitro studies using isolated rat pancreatic islets have shown that ginsenosides promote an insulin release which is independent of extracellular calcium and which utilises a different mechanism to that of glucose.\(^{(13)}\) In addition, in vivo studies in rats have reported that a ginseng extract increases the number of insulin receptors in bone marrow and reduces the number of glucocorticoid receptors in rat brain homogenate.\(^{(14)}\) Both of these actions are thought to contribute to the antidiabetic action of ginseng, in view of the known diabetogenic action of adrenal corticoids and the knowledge that the number of insulin receptors generally decreases with ageing.\(^{(14)}\)

Hypoglycaemic activity observed in both normal and alloxan–induced hyperglycaemic mice administered ginseng (intraperitoneal) has also been attributed to non–saponin but uncharacterised principles\(^{(15-18)}\) and to glycan (polysaccharide) components, Panaxans A–E and Q–U.\(^{(19-23)}\) Glycans isolated from Korean ginseng or Chinese ginseng (A–E) were found to possess stronger hypoglycaemic activity than those isolated from Japanese ginseng (Q–U).\(^{(23)}\) Proposed mechanisms of action have included elevated plasma insulin concentration due to an increase of insulin secretion from pancreatic islets, and enhancement of insulin sensitivity.\(^{(21)}\) However, these mechanisms do not explain the total hypoglycaemic activity that has been exhibited by the polysaccharides and further mechanisms are under investigation.\(^{(21)}\)

The effect of panaxans A and B on the activities of key enzymes participating in carbohydrate metabolism has been studied.\(^{(18)}\) DPG-3–2, a non–saponin component isolated from ginseng, has been shown to stimulate insulin biosynthesis in pancreatic preparations from various hyperglycaemic (but not normoglycaemic) animals; ginsenosides Rb\(_1\) and Rg\(_1\) were found to decrease islet insulin concentrations to an undetectable level.\(^{(16)}\)

**Cardiovascular activity**

Individual saponins have been reported to have different actions on cardiac haemodynamics.\(^{(24)}\) For instance R\(_g\), R\(_{g-1}\) and total flower saponins have increased cardiac performance whilst R\(_b\) and total leaf saponins have decreased it; calcium antagonist activity has been reported for R\(_b\) but not for R\(_g\); R\(_b\) but not R\(_g\) has produced a protective effect on experimental myocardial infarction in rabbits.\(^{(24)}\) Negative chronotropic and inotropic effects in vitro have been observed for ginseng saponins and a mechanism of action similar to that of verapamil has been suggested.\(^{(25)}\) In vitro studies on the isolated rabbit heart have reported an increase in coronary blood flow together with a positive inotropic effect.\(^{(26)}\) Anti–arrhythmic action on aconitine and barium chloride (rat) and adrenaline (rabbit)-induced arrhythmias, and prolongation
of RR, PR and QT$_c$ intervals (rat), have been documented for saponins R$_{c-1}$ and R$_{d-1}$. The mode of action was thought to be similar to that of amiodarone. (27) Ginsenosides (i.p.) have been reported to protect mice against metabolic disturbances and myocardial damage associated with conditions of severe anoxia. (28)

Ginseng has produced a marked hypotensive response together with bradycardia following intravenous administration to rats. The dose–related effect was blocked by many antagonists suggesting multi–site activity. (26) Higher doses of ginseng were found to cause vasoconstriction rather than vasodilation in renal, mesenteric and femoral arteries. (26)

The total ginseng saponin fraction has been reported to be devoid of haemolytic activity. However, individual ginsenosides have been found to exhibit either haemolytic or protective activities. Protective ginsenosides include R$_c$, R$_{b-2}$ and R$_e$, whereas haemolytic saponins have included R$_g$, R$_h$ and R$_f$. (29) The number and position of sugars attached to the sapogenin moiety was thought to determine activity. (29) Haemostatic activity has also been documented for ginseng. (30)

Oral administration of ginseng to rats fed a high cholesterol diet reduced serum cholesterol and triglycerides, increased high–density lipoprotein (HDL) cholesterol, decreased platelet adhesiveness, and decreased fatty changes to the liver. (31) Ginseng has also been reported to reduce blood coagulation and enhance fibrinolysis. (32) Panaxynol and the ginsenosides R$_o$, R$_{g-1}$ and R$_{g-2}$ have been documented as the main antiplatelet components in ginseng inhibiting aggregation, release reaction and thromboxane formation in vitro. (32) Anti–inflammatory activity and inhibition of thromboxane B$_2$ have previously been described for panaxynol. (32) Anticomplementary activity in vitro (human serum) has been documented for ginseng polysaccharides with highest activity observed in strongly acidic polysaccharide fractions. (7)

**Effects on neurotransmitters**
Studies in rats have shown that a standardised ginseng extract (G115) inhibits the development of morphine tolerance and physical dependence, of a decrease in hepatic glutathione concentrations, and of dopamine receptor sensitivity without antagonising morphine analgesia, as previously documented for the individual saponins. (33) The inhibition of tolerance was thought to be associated with a reduction in morphinone production, a toxic metabolite which irreversibly blocks the opiate receptor sites, and with the activation of morphinone–glutathione conjugation, a detoxication process. The mechanism of inhibition of physical dependence was unclear but thought
to be associated with changed ratios of adrenaline, noradrenaline, dopamine and serotonin in the brain.\(^{33}\)

A total ginsenoside fraction has been reported to inhibit the uptake of various neurotransmitters into rat brain synaptosomes in descending order of gamma–aminobutyrate and noradrenaline, dopamine, glutamate and serotonin.\(^{34\text{-}36}\) The fraction containing ginsenoside \(R_d\) was most effective. Uptake of metabolic substrates 2–deoxy-D-glucose and leucine was only slightly affected and therefore it was proposed that the ginseng extracts were acting centrally rather than locally as surface active agents.

Studies in rats have indicated that the increase in dopaminergic receptors in the brain observed under conditions of stress is prevented by pretreatment with ginseng.\(^{37}\)

**Hepatoprotective activity**

Antioxidant and detoxifying activities have been documented for ginseng.\(^{38}\) Protection against carbon tetrachloride- and galactos amine–induced hepatotoxicity has been observed in cultured rat hepatocytes for specific ginsenosides (oleanolic acid and dammarane series).\(^{38,39}\) However, at higher doses certain ginsenosides from both series were found to exhibit simultaneous cytotoxic activity.\(^{36}\)

**Cytotoxic and antitumour activity**

Cytotoxic activity (ED\(_{50}\) 0.5 μg/mL) versus L1210 has been documented for polyacetylenes isolated from the root.\(^{5,6,40}\) The antitumour effect of ginseng polysaccharides in tumour–bearing mice has been associated with an immunological mechanism of action.\(^{41}\) Ginseng polysaccharides have been reported to increase the lifespan of tumour–bearing mice and to inhibit the growth of tumour cells \textit{in vivo}, although cytocidal action was not seen \textit{in vitro}.\(^{41}\) Antitumour activity \textit{in vitro} versus several tumour cell lines has been documented for a polyacetylene, panaxytriol.\(^{42}\)

**Antiviral activity**

Antiviral activity (versus Semliki forest virus; 34–40% protection) has been documented for ginseng extract (G115, Pharmaton) administered orally to rats.\(^{37}\) The ginseng extract also enhanced the level of protection afforded by 6-MFA, an interferon–inducing agent of fungal origin.\(^{43}\) Ginseng has been found to induce \textit{in vitro} and \textit{in vivo} production of interferon and to augment the natural killer and antibody dependent cytotoxic activities in human peripheral lymphocytes.\(^{43,44}\) In addition, ginseng enhances the antibody–forming cell response to sheep red blood cells in mice and stimulates cell
mediated immunity both \textit{in vitro} and \textit{in vivo}.\cite{43,44} In view of these observations, it has been proposed that the antiviral activity of ginseng may be immunologically mediated.\cite{43,44}

**Clinical studies**

Improvements in serum total cholesterol, HDL cholesterol, triglycerides, non-essential fatty acids and lipoperoxides have been observed in 67 hyperlipidaemic patients administered 2.7 g/day red ginseng.\cite{27} The addition of ginseng (3 g/65 kg body weight) to alcohol consumption (72 g/65 kg body weight of 25% ethanol) has been reported to enhance blood alcohol clearance by 32–51\%.\cite{45}

A preparation containing ginseng extract with multivitamins and trace elements has been shown to modify some indices of metabolic and liver function in elderly patients with chronic hepatotoxicity induced by alcohol and drugs.\cite{46} Patients who received ginseng exhibited an increase in bromosulphthalein excretion (which is related to hepatic detoxification) and improved serum zinc concentrations.\cite{46}

A favourable effect on various tests of psycho motor performance (attention, processing, integrated sensory motor function and auditory reaction time) in healthy individuals receiving a ginseng extract (200 mg daily for 12 weeks) has been documented in a double-blind placebo-controlled study.\cite{47} No difference was observed between ginseng and placebo groups in tests of pure motor function, recognition and visual reaction time.\cite{47}

Ginseng has been reported to improve the overall control of status asthmaticus when added to conventional steroid, bronchodilator and antibiotic therapies.\cite{48}

Ginseng has been shown to reduce blood sugar concentrations in both diabetics and non-diabetics,\cite{1} such that in one study insulin therapy was no longer required in a proportion of the patients investigated.\cite{1}

Ginseng has also been reported to normalise both high and low blood pressure states.\cite{1}

Ginseng has been found to affect concentrations of corticosteroids such as adrenocorticotrophic hormone (ACTH) and cortisol and noradrenaline.\cite{1}

Ginseng has been reported to successfully treat cases of diabetic polyneuropathy, reactive depression, psychogenic impotence, enuresis and various child psychiatric disorders.\cite{49}
In Japan, ginseng (panax) has been given to more than 500 individuals over the course of two studies with no side-effects experienced.\(^{1}\) However, suspected adverse events associated with ginseng treatment have been documented although it is often difficult to assess individual cases due to a lack of information concerning dose, duration of treatment, species of ginseng used, and concurrent medication.\(^{1}\) Nevertheless, symptoms documented include hypertension\(^{1}\) (ginseng species unspecified), diarrhoea,\(^{1}\) insomnia\(^{1}\) (as a result of over stimulation), mastalgia,\(^{1}\) skin eruptions\(^{1}\) and vaginal bleeding\(^{1}\). A case of vaginal bleeding in a postmenopausal woman has been associated with the use of a ginseng face cream.\(^{50}\) In 1979, two studies referred to a ginseng abuse syndrome (GAS) which emphasised that most side-effects documented for ginseng were associated with the ingestion of large doses of ginseng together with other psychomotor stimulants, including tea and coffee.\(^{1}\) GAS was defined as diarrhoea, hypertension, nervousness, skin eruptions and sleeplessness. Other symptoms occasionally observed included amenorrhoea, decreased appetite, depression, euphoria, hypotension and oedema. However, these two studies have been widely criticised over the variety of ginseng and other preparations used, and over the lack of authentication of the ginseng species ingested.\(^{1}\)

Elsewhere, symptoms of overdose have been described as those exhibited by individuals allergic to ginseng, namely palpitations, insomnia and pruritus, together with heart pain, decrease in sexual potency, vomiting, haemorrhagic diathesis, headache and epistaxis; ingestion of very large doses have even been reported to be fatal.\(^{9}\)

Two cases of a suspected interaction between ginseng and phenelzine have been documented.\(^{51}\) Symptoms of headache and tremulousness in one 64-year-old woman and of manic-like symptoms in a 42-year-old woman were described.\(^{51}\)

Results documented for toxicity studies carried out in a number of animal species using standardised extracts (SE) indicate ginseng to be of low toxicity.\(^{52-56}\)

**Acute toxicity**

Single doses of up to 2 g SE have been administered to mice and rats with no toxic effects observed.\(^{55}\) LD\(_{50}\) values (p.o.) in mice and rats have been estimated at 2 g/kg and greater than 5 g/kg.\(^{52}\) In addition, LD\(_{50}\) values (i.p., mice) have been estimated for individual ginsenosides as 305 mg/kg (R\(_{b-2}\)), 324 mg/kg (R\(_d\)), 405 mg/kg (R\(_e\)), 410 mg/kg (R\(_c\)), 1110 mg/kg (R\(_{b-1}\)),
1250 mg/kg ($R_{g-1}$), and 1340 mg/kg ($R_f$); an LD$_{50}$ (i.v., mice) of 3806 mg/kg has been estimated for the saponins $R_{c-1}$ and $R_{d-1}$.$^{(52)}$

**Subacute toxicity**
Doses of approximately 720 mg of a ginseng extract (G115) have been administered orally to rats for 20 days with no side-effects documented.$^{(56)}$

**Chronic toxicity**
Daily doses of up to 15 mg G115/kg body weight have been administered orally to dogs for 90 days with no toxic effects documented. An initial increase in excitability which disappeared after two to three weeks was the only observation reported in rats fed 200 mg G115/kg body weight for 25 weeks.$^{(55)}$

*P. quinquefolis* has been reported to be devoid of mutagenic potential when investigated versus *Salmonella typhimurium* strain TM677.$^{(57)}$
Contra-indications, Warnings

Ginseng may potentiate the action of monoamine oxidase inhibitors (MAOIs) (inhibits uptake of various neurotransmitter substances)\(^{(34)}\) and two cases of suspected ginseng interaction with phenelzine have been documented.\(^{(51)}\) The use of ginseng has been contraindicated during acute illness, any form of haemorrhage and during the acute period of coronary thrombosis\(^{(1)}\). It has been recommended that ginseng should be avoided by individuals who are highly energetic, nervous, tense, hysterical, manic or schizophrenic, and that it should not be taken with stimulants, including coffee, antipsychotic drugs or during treatment with hormones.\(^{(1,G49)}\)

In view of documented pharmacological actions and side-effects, ginseng should also be used with caution in the following circumstances: cardiac disorders, diabetes, hyper- and hypotensive disorders, and with all steroid therapy. Women may experience oestrogenic side-effects.

In Russia, it is recommended that healthy people under the age of 40 should not use ginseng and that middle-aged people can be treated with small doses of ginseng on a daily basis. In general, long-term use of ginseng is not recommended and one author has documented that the main side-effect of prolonged use manifests as an inflamed nerve, frequently the sciatic, which then causes muscle spasm in the affected area.\(^{(1)}\) In Russia, those individuals considered suitable to use ginseng are recommended to abstain from alcoholic beverages, sexual activity, bitter substances and spicy foods.\(^{(9)}\) Patients allergic to ginseng may exhibit symptoms of palpitation, insomnia, and pruritus.\(^{(9)}\)

Pregnancy and lactation

No fetal abnormalities have been observed in rats and rabbits administered a standardised extract (40 mg/kg, p.o.) from day 1 to day 15 of pregnancy.\(^{(55)}\) Ginseng has also been fed to two successive generations of rats in doses of up to 15 mg G115/kg body weight/day (equivalent to approximately 2700 mg ginseng extract) with no teratogenic effects observed.\(^{(52)}\) However, the safety of ginseng during pregnancy has not been established in humans and therefore its use should be avoided. Similarly, there are no published data concerning the secretion of pharmacologically active constituents from ginseng into the breast milk and use of ginseng during lactation is therefore best avoided.
Phytochemical studies on panax ginseng are well documented and have initially concentrated on the saponin components (ginsenosides) which are generally considered to be the main active constituents. More recently, pharmacological actions documented for the non-saponin components, principally polysaccharides, have stimulated research into identifying non-saponin active constituents. Many of the pharmacological actions documented for ginseng directly oppose one another and this has been attributed to the actions of the individual ginsenosides. For example, ginsenoside R\textsubscript{b1} exhibits CNS-depressant, hypotensive and tranquillising actions whilst ginsenoside R\textsubscript{g1} exhibits CNS-stimulant, hypertensive and anti-fatigue actions. These opposing actions are thought to explain the ‘adaptogenic’ reputation of ginseng, that is the ability to increase the overall resistance of the body to stress and to balance bodily functions.

In summary, ginseng has been shown to possess a wide range of pharmacological activities and it should consequently be used with appropriate regard to the traditional guidelines drawn up in China, Japan and Russia, to the health of the individual and to any concomitant therapies. When used appropriately, ginseng appears to be relatively non-toxic and most documented side-effects are associated with inappropriate use when compared with traditional warnings and guidelines.
References


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Golden Seal
Species (Family)

*Hydrastis canadensis* L. (Ranunculaceae)
Synonym(s)

Yellow Root

Yellow root also refers to *Xanthorhiza simplicissima* Marsh, which is also a member of the Ranunculaceae family and contains berberine as the major alkaloid constituent.
Part(s) Used

Rhizome, root
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)
BHP 1996\(^{(G9)}\)
Martindale 33rd edition\(^{(G67)}\)
Mills and Bone (Hydrastis root)\(^{(G50)}\)
PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
USP26/NF21\(^{(G73)}\)
Legal Category (Licensed Products)

GSL(G37)
Constituents

See General References G6 G22 G40 G41 G62 G64.

Alkaloids
Isoquinoline–type. 2.5–6.0%. Hydrastine (major, 1.5–4.0%), berberine (0.5–6.0%), berberastine (2–3%), and canadine (1%), with lesser amounts of related alkaloids including candaline and canadaline.\(^{(1–3)}\)

Other constituents
Chlorogenic acid, carbohydrates, fatty acids (75% saturated, 25% unsaturated), volatile oil (trace), resin, meconin (meconinic acid lactone).
Food Use

Golden seal is not used in foods, although it is reported to be used in herbal teas.\(^{(G41)}\) The concentration of berberine permitted in foods is limited to 0.1 mg/kg, and 10 mg/kg in alcoholic beverages.\(^{(G16)}\)
Herbal Use

Golden seal is stated to be a stimulant to involuntary muscle, and to possess stomachic, oxytocic, anti haemorrhagic and laxative properties. Traditionally it has been used for digestive disorders, gastritis, peptic ulceration, colitis, anorexia, upper respiratory catarrh, menorrhagia, post-partum haemorrhage, dysmenorrhoea, topically for eczema, pruritus, otorrhoea, catarrhal deafness and tinnitus, conjunctivitis, and specifically for atonic dyspepsia with hepatic symptoms.\(^{(G6 G7 G8)}\)
Dosage

*Dried rhizome*
0.5–1.0 g or by decoction three times daily.\(^{G6 \ G7}\)

*Liquid Extract of Hydrastis*
(BPC 1949) 0.3–1.0 mL.

*Tincture of Hydrastis*
(BPC 1949) 2–4 mL.
Pharmacological Actions

The pharmacological activity of golden seal is attributed to the isoquinoline alkaloid constituents, primarily hydrastine and berberine,\(^{3,4}\) which are reported to have similar properties.\(^{G41}\) Antibiotic, immunostimulant, anticonvulsant, sedative, hypotensive, uterotonic, choleretic and carminative activities have been described for berberine.\(^{3}\)

**In vitro and animal studies**

Limited work has been documented for golden seal, although the pharmacology of berberine and hydrastine is well studied.

The total alkaloid fraction of golden seal has been reported to exhibit anticonvulsant activity in smooth muscle preparations (e.g. mouse intestine, uterus).\(^{5}\) However, *in vitro*, canadine is reported to exhibit uterine stimulation in guinea-pig and rabbit tissues.\(^{4}\) Berberine, canadine and hydrastine are all stated to exhibit utero–activity.\(^{G30}\)

Berberine and hydrastine have produced a hypotensive effect in laboratory animals following intravenous administration.\(^{6,7,G41}\) High doses of hydrastine are documented to produce an increase in blood pressure.\(^{7}\) *In vitro*, berberine has been reported to decrease the anticoagulant action of heparin in canine and human blood.\(^{7}\)

Berberine is reported to exert a stimulant action on the heart and to increase coronary blood flow, although higher doses are stated to inhibit cardiac activity.\(^{7}\)

Antimuscarinic and antihistamine actions have been documented for berberine.\(^{7}\)

In rats, berberine has exhibited antipyretic activity three times as effective as aspirin.\(^{3}\)

Berberine potentiated barbiturate sleeping time, but did not exhibit any analgesic or tranquillising effects.\(^{7}\)

A broad spectrum of antimicrobial activity against bacteria, fungi, and protozoa has been reported for berberine. Sensitive organisms include *Staphylococcus* spp., *Streptococcus* spp., *Chlamydia* *aureus*, *Corynebacterium diphtheriae*, *Salmonella typhi*, *Diplococcus pneumoniae*, *Pseudomonas aeruginosa*, *Shigella dysenteriae*, *Trichomonas vaginalis*, *Neisseria gonorrhoeae*, *Neisseria meningitidis*, *Treponema pallidum*, *Giardia lamblia* and...
Leishmania donovani.\(^{(3)}\) Berberine is reported to be effective against diarrhoeas caused by enterotoxins such as *Vibrio cholerae* and *Escherichia coli*.\(^{(7)}\) In *vivo* and *in vitro* studies in hamsters and rats have reported significant activity for berberine against *Entamoeba histolytica*.\(^{(3)}\)

Anticancer activity has been reported for berberine in B1, KB and PS tumour systems.\(^{(G22)}\) In addition, berberine sulfate was found to inhibit the action of teleocidin, a known tumour promoter, on the formation of mouse skin tumours initiated with 7,12–dimethylbenz[a]anthracene.\(^{(5)}\)

**Clinical studies**

None documented for golden seal. Berberine is stated to have shown significant success in the treatment of acute diarrhoea in several clinical studies.\(^{(3)}\) It has been found effective against diarrhoeas caused by *Escherichia coli, Shigella dysenteriae, Salmonella paratyphi* B, *Klebsiella, Giardia lamblia* and *Vibrio cholerae*.\(^{(3)}\) Berberine has been used to treat trachoma, an infectious ocular disease caused by *Chlamydia trachomatis*, which is a major cause of blindness and impaired vision in developing countries.\(^{(3)}\)

Clinical studies have shown berberine to stimulate bile and bilirubin secretion and to improve symptoms of chronic cholecystitis, and to correct raised levels of tyramine in patients with liver cirrhosis.\(^{(3)}\)
Side-effects, Toxicity

Berberine and berberine–containing plants are considered to be non–toxic.\(^{(3)}\) However, the alkaloid constituents are potentially toxic and symptoms of golden seal poisoning include stomach upset, nervous symptoms and depression; large quantities may even be fatal.\(^{(8)}\) High doses of hydrastine are reported to cause exaggerated reflexes, convulsions, paralysis and death from respiratory failure.\(^{(4)}\) The root may cause contact ulceration of mucosal surfaces.
Contra-indications, Warnings

Golden seal is contra-indicated in individuals with raised blood pressure.\(^{(G7 G22 G49)}\) Prolonged use of golden seal may decrease vitamin B absorption.\(^{(G22)}\) Coagulant activity opposing the action of heparin, and cardiac stimulant activity have been documented for berberine. The use of golden seal as a douche should be avoided because of the potential ulcerative side-effects.\(^{(G22)}\) The alkaloid constituents of golden seal are potentially toxic and excessive use should be avoided.

**Pregnancy and lactation**

Golden seal is contra-indicated for use during pregnancy.\(^{(3, G7 G49)}\) Berberine, canadine, hydrastine and hydrastinine have all been reported to produce uterine stimulant activity.\(^{(G30)}\) It is not known whether the alkaloids are excreted in breast milk. The use of golden seal during lactation should be avoided.
Pharmaceutical Comment

Golden seal is characterised by the isoquinoline alkaloid constituents. These compounds, primarily hydrastine and berberine, represent the main active components of golden seal. Numerous activities have been documented many of which support the traditional herbal uses of the root. However, in view of the pharmacological properties of the alkaloid constituents, excessive use of golden seal should be avoided.
References


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Species (Family)

Eupatorium purpureum L. (Asteraceae/Compositae)
Synonym(s)

Joe-Pye Weed, Kydney Root, Purple Boneset, Queen of the Meadow
Part(s) Used

Rhizome, root
Pharmacopoeial and Other Monographs

BHP 1983\(^{(G7)}\)

Martindale 33rd edition\(^{(G67)}\)
Legal Category (Licensed Products)

GSL(G³)
Constituents

See General References G20 G40 G42 G48 G49 G64.

Little information is available on the chemistry of gravel root. It is stated to contain euparin (a benzofuran compound), eupatorin (a flavonoid), resin and volatile oil.

Other plant parts

The herb is reported to contain echinatine, an unsaturated pyrrolizidine alkaloid.\(^{(1)}\)
Food Use

Gravel root is not used in foods.
Herbal Use

Gravel root is stated to possess antilithic, diuretic and antirheumatic properties. Traditionally, it has been used for urinary calculus, cystitis, dysuria, urethritis, prostatitis, rheumatism, gout, and specifically for renal or vesicular calculi. \(^{(G7 \text{ G64})}\)
Dosage

*Dried rhizome/root*
2–4 g or by decoction three times daily.\(^{(G7)}\)

*Liquid extract*
2–4 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)

*Tincture*
1–2 mL (1 : 5 in 40% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

None documented.
Side–effects, Toxicity\(^{(G20)}\)

None documented for gravel root although pyrrolizidine alkaloids are constituents of many species of *Eupatorium*.\(^{(1,G20)}\) Pyrrolizidine alkaloids with an unsaturated pyrrolizidine nucleus are reported to be hepatotoxic in both animals and humans (see Comfrey). An unsaturated pyrrolizidine alkaloid, echinatine, has been reported for the aerial parts of gravel root.
Contra-indications, Warnings

None documented.

**Pregnancy and lactation**
The safety of gravel root has not been established. In view of the lack of phytochemical, pharmacological and toxicological information the use of gravel root during pregnancy and lactation should be avoided.
The chemistry of gravel root is poorly studied and no scientific evidence was located to justify the herbal uses. Excessive use of gravel root should be avoided.
See also General References G7 G20 G31 G37 G40 G42 G48 G49 G64.


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Ground Ivy
Species (Family)

*Nepeta hederacea* (L.) Trev. (Labiatae)
Synonym(s)

Glechoma hederacea L.
Part(s) Used

Herb
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents
See General References G6 G22 G48 G64.

Amino acids
Asparagic acid, glutamic acid, proline, tyrosine and valine.

Flavonoids
Flavonol glycosides (e.g. hyperoside, isoquercitrin, rutin) and flavone glycosides (e.g. luteolin diglucoside, cosmosyin).\(^{(1)}\)

Steroids
β-Sitosterol.

Terpenoids
Oleanolic acid, α-ursolic acid, β-ursolic acid.\(^{(2)}\)

Volatile oils
0.03–0.06%. Various terpenoid components including \(p\)-cymene, linalool, limonene, menthone, α-pinene, β-pinene, pinocamphone, pulegone and terpineol; glechomafuran (a sesquiterpene).\(^{(3)}\)

Other constituents
Palmitic acid, rosmarinic acid, succinic acid, bitter principle (glechomin), choline, gum, diterpene lactone (marrubiin), saponin, tannin and wax.
Food Use

Ground ivy is listed by the Council of Europe as a natural source of food flavouring (category N3). This category indicates that ground ivy can be added to foodstuffs in the traditionally accepted manner, although there is insufficient information available for an adequate assessment of potential toxicity.\(^{(G16)}\)
Herbal Use

Ground ivy is stated to possess mild expectorant, anticatarrhal, astringent, vulnerary, diuretic and stomachic properties. Traditionally, it has been used for bronchitis, tinnitus, diarrhoea, haemorrhoids, cystitis, gastritis, and specifically for chronic bronchial catarrh.
Dosage

*Dried herb*

2–4 g or by infusion three times daily.\(^{G6\ G7}\)

*liquid extract*

2–4 mL (1:1 in 25% alcohol) three times daily.\(^{G6\ G7}\)
Pharmacological Actions

In vitro and animal studies

In vivo anti-inflammatory activity has been reported for an ethanolic extract of ground ivy, which was stated to exhibit a moderate inhibition (27%) of carrageenan-induced rat paw oedema.\(^4\)

Ursolic acid analogues, 2α- and 2β-hydroxyursolic acid, have been documented to provide significant ulcer-protective activity in mice.\(^5\)

The astringent activity documented for ground ivy has been attributed to rosmarinic acid, a polyphenolic acid.\(^6\)

Glechomin and marrubiin are stated to be bitter principles, and α-terpineol is known to be an antiseptic component of volatile oils.\(^{G48 \text{ G49}}\)

Anti-inflammatory and astringent properties are generally associated with flavonoids and tannins, respectively. Anti-inflammatory properties have been documented for rosmarinic acid (see Rosemary).

In vitro antiviral activity against the Epstein–Barr virus has been documented for ursolic acid.\(^7\) Both oleanolic and ursolic acids were found to inhibit tumour production by TPA in mouse skin, with activity comparable to that of retinoic acid, a known tumour-promoter inhibitor.\(^7\)

Significant cytotoxic activity has also been reported for ursolic acid in lymphocytic leukaemia (P-388, L-1210) and human lung carcinoma (A-549), and marginal activity in KB cells, human colon (HCT-8), and mammary (MCF-7) tumour cells.\(^8\)
Side–effects, Toxicity

Poisoning in cattle and horses has been documented in Eastern Europe.\(^{(9)}\) Symptoms include accelerated weak pulse, difficulty in breathing, conjunctival haemorrhage, elevated temperature, dizziness, spleen enlargement, dilation of the caecum, and gastroenteritis revealed at post–mortem. Anti–tumour and cytotoxic activities have been reported for oleanolic and ursolic acids (see *In vitro* and animal studies).

Ground ivy volatile oil contains many terpenoids and terpene–rich volatile oils are irritant to the gastrointestinal tract and kidneys. Pulegone is an irritant, hepatotoxic, and abortifacient principle of the volatile oil of pennyroyal. However, in comparison with pennyroyal the overall yield of volatile oil is much less (0.03–0.06% in ground ivy and 1–2% in pennyroyal).
Contra–indications, Warnings

Ground ivy is contra–indicated in epilepsy \(^{(G7)}\) although no rationale for this statement has been found. Excessive doses may be irritant to the gastrointestinal mucosa and should be avoided by individuals with existing renal disease.

**Pregnancy and lactation**

The safety of ground ivy has not been established. In view of the lack of toxicity data and the possible irritant and abortifacient action of the volatile oil, the use of ground ivy during pregnancy and lactation should be avoided.
Pharmaceutical Comment

The chemistry of ground ivy is well studied. Documented pharmacological activities support some of the herbal uses, although no references to human studies were located. In view of the lack of toxicity data and the reported cytotoxic activity of ursolic acid, excessive use of ground ivy should be avoided.
References

See also General References G6 G9 G16 G22 G31 G36 G37 G48 G49 G64.


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Species (Family)

i. *Guaiacum officinale* L. (Zygophyllaceae)

ii. *Guaiacum sanctum* L.
Synonym(s)
Guaiac, Guajacum, Lignum Vitae
Part(s) Used
Resin obtained from the heartwood
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

Complete German Commission E\(^{(G3)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents
See General References G6 G48 G64.

Resins
15–20%. Guaiaretic acid, dehydroguaiaretic acid, guaiac, isoguaiac, α-guaiaconic acid (lignans), furoguaiacin and its monomethyl ether, furoguaiacidin, tetrahydrofuroguaiacin-A and tetrahydrofuroguaiacin-B (furano–lignans), furoguaia oxidin (enedione lignan).\(^{1-4}\)

Steroids
β-Sitosterol.

Terpenoids
Saponins, oleanolic acid.\(^{5,6}\)
Food Use

Guaiacum is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that guaiacum can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\textsuperscript{(G16)}
Herbal Use

Guaiacum is stated to possess antirheumatic, anti-inflammatory, diuretic, mild laxative and diaphoretic properties. Traditionally, it has been used for subacute rheumatism, prophylaxis against gout, and specifically for chronic rheumatism and rheumatoid arthritis. \(^{(G6\ G7\ G8\ G64)}\)
Dosage

**Dried wood**
1–2 g or by decoction three times daily.\(^{(G6 \ G7)}\)

**Liquid extract**
1–2 mL (1 : 1 in 80% alcohol) three times daily.\(^{(G6 \ G7)}\)

**Tincture of Guaiacum**
(BPC 1934) 2–4 mL.
Pharmacological Actions

*In vitro and animal studies*

None documented. Antimicrobial properties are associated with lignans and much has been documented for nordihydroguaiaretic acid, the principal lignan constituent in chaparral (see Chaparral).
Side–effects, Toxicity

Guaiacum resin has been reported to cause contact dermatitis.\textsuperscript{(G51)} The resin is documented to be of low toxicity; the oral LD\textsubscript{50} in rats is greater than 5 g/kg body weight.\textsuperscript{(G48)}
Contra-indications, Warnings

It is recommended that guaiacum is avoided by individuals with hypersensitive, allergic or acute inflammatory conditions.\(^{(G49)}\)

**Pregnancy and lactation**

The safety of guaiacum during pregnancy has not been established. In view of this, and the overall lack of pharmacological and toxicological data, the use of guaiacum during pregnancy and lactation should be avoided.
Pharmaceutical Comment

Guaiacum is characterised by the resin fraction of the heartwood and much has been documented on the constituents (principally lignans) of the resin, although little is known regarding other constituents. No scientific information was found to justify the herbal use of guaiacum as an antirheumatic or anti-inflammatory agent. In view of the lack of toxicity data, excessive use of guaiacum should be avoided.


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Hawthorn
Species (Family)

i. *Crataegus laevigata* (Pois) DC. (Rosaceae)

ii. *Crataegus monogyna* Jacq.
Synonym(s)

Whitethorn
Part(s) Used

Fruit
Pharmacopoeial and Other Monographs

American Herbal Pharmacopoeia\textsuperscript{(G1)}

BHP 1996\textsuperscript{(G9)}

BP 2002\textsuperscript{(G71)}

Complete German Commission E\textsuperscript{(G3)}

ESCOP 1999\textsuperscript{(G52)}

Martindale 33rd edition\textsuperscript{(G67)}

Mills and Bone\textsuperscript{(G50)}

PDR for Herbal Medicines 2nd edition\textsuperscript{(G36)}

Ph Eur 2004\textsuperscript{(G72)}
Legal Category (Licensed Products)

Hawthorn is not included in the GSL.\(^{G37}\)
Constituents
See General References G1 G2 G22 G62 G64.

**Amines**
Phenylethylamine, O-methoxyphenethyl amine and tyramine.\(^1\)

**Flavonoids**
Flavonol (e.g. kaempferol, quercetin) and flavone (e.g. apigenin, luteolin) derivatives, rutin, hyperoside, vitexin glycosides, orientin glycosides\(^{2-5}\) and procyanidins.\(^6,7\)

**Tannins**
Pharmacopoeial standard not less than 1.0% procyanidins, condensed (proanthocyanidins).

**Other constituents**
Cyanogenetic glycosides and saponins.
Food Use

Hawthorn is not commonly used in foods. It is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that hawthorn can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\(G^{16}\)
Hawthorn is stated to possess cardiotonic, coronary vasodilator and hypotensive properties. Traditionally, it has been used for cardiac failure, myocardial weakness, paroxysmal tachycardia, hypertension, arteriosclerosis and Buerger’s disease. The German Commission E did not approve therapeutic use.
Dosage

*Dried fruit*
0.3–1.0 g or by infusion three times daily.\(^{(G7)}\)

*Liquid extract*
0.5–1.0 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)

*Tincture*
1–2 mL (1 : 5 in 45% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

In vitro and animal studies

Cardiovascular activity has been documented for hawthorn and attributed to the flavonoid components, in particular the procyanidins. Hawthorn extracts have been documented to increase coronary blood flow both in vitro (in the guinea–pig heart) and in vivo (in the cat, dog and rabbit), reduce blood pressure in vivo (in the cat, dog, rabbit and rat), increase (head, skeletal muscle and kidney) and reduce (skin, gastrointestinal tract) peripheral blood flow in vivo (in the dog) and reduce peripheral resistance in vivo (in the dog).\(^{(6,8–13)}\) The hypotensive activity of hawthorn has been attributed to a vasodilation action rather than via adrenergic, muscarinic or histaminergic receptors.\(^{(12)}\) Beta–adrenoceptor blocking activity (versus adrenaline–induced tachycardia) has been exhibited in vivo in the dog and in vitro in the frog heart using flower, leaf and fruit extracts standardised on their procyanidin content.\(^{(6)}\) The authors reported a direct relationship between the concentration of procyanidin and observed actions.

Negative chronotropic and positive inotropic actions have been observed in vitro using the guinea–pig heart and attributed to flavonoid and proanthocyanidin fractions.\(^{(14)}\) A positive inotropic effect has also been exhibited by amine constituents in vitro using guinea–pig papillary muscle.\(^{(1)}\)

Hawthorn extracts have also been reported to lack any effect on the heart rate and muscle contractility in studies that have observed an effect on blood pressure in the dog and rat.\(^{(11,12)}\) Hawthorn extracts have exhibited some prophylactic anti–arrhythmic activity in rabbits administered intravenous aconitine.\(^{(15)}\) Extracts infused after aconitine did not affect the induced arrhythmias. In vitro, vitexin rhamnoside has been reported to have no effect on the action of ouabain and aconitine.\(^{(15)}\) A crude extract of *Crataegus pinnatifida* Bge. var *major* N.E.Br. and the flavonoid vitexin rhamnoside have been reported to exert a protective action on experimental ischaemic myocardium in anaesthetised dogs.\(^{(16)}\) The extracts were observed to decrease left ventricular work, decrease the consumption of oxygen index, and increase coronary sinus blood oxygen concentrations, resulting in a decrease in oxygen consumption and balance of oxygen metabolism. In contrast to other studies, an increase in coronary blood flow was not observed. The authors attributed these opposing results to the variation in concentrations of active constituents between the different plant parts. In vitro, vitexin rhamnoside has been reported to exert a protective action towards cardiac cells deprived of oxygen and glucose.\(^{(17)}\)
A mild CNS-depressant effect has been documented in mice that received oral administration of hawthorn flower extracts.(18) An increase in barbiturate sleeping time and a decrease in spontaneous basal motility were the most noticeable effects.

Free radicals have been linked with the ageing process. When fed to mice, a hawthorn fruit (C. pinnatifidia) extract has been reported to enhance the action of superoxide dismutase (SOD), which promotes the scavenging of free radicals.(19) An inhibition of lipid peroxidation, which can be caused by highly reactive free radicals was also documented.(19) The pharmacological actions of leaf with flowers include increase in cardiac contractility, increase in coronary blood flow and myocardial circulation, protection from ischaemic damage and decrease of peripheral vascular resistance.(G52)

Clinical studies

The effects of a commercial preparation containing 30 mg hawthorn extract standardised to 1 mg procyanidins were assessed in a double-blind, placebo-controlled study involving 80 patients.(20) Hawthorn extract was reported to exhibit greater overall improvement of cardiac function and of subjective symptoms, such as dyspnoea and palpitations, compared with placebo. Improvements in ECG recordings were not found to differ between the two groups.

A commercial product containing hawthorn, valerian, camphor and cereus was given to 2243 patients with functional cardiovascular disorders and/or hypotension or meteorosensitivity in an open multicentre study.(21) An improvement in 84% of treated individuals was reported.

In a randomised, double-blind, controlled trial involving 60 patients with stable angina, a commercial hawthorn preparation (60 mg three times daily) was reported to increase coronary perfusion and to economise myocardial oxygen consumption.(22)

A commercial hawthorn/passionflower extract (standardised on flavone and proanthocyanidin content) 6 mL daily for 42 days has been assessed in a randomised, double-blind, placebo-controlled trial involving 40 patients with chronic heart failure.(23) Significant improvements were noted for the active group, compared with placebo, in exercise capacity, heart rate at rest, diastolic blood pressure at rest, and concentrations of total plasma cholesterol and low-density lipids. Non-significant improvements were noted in the active group for maximum exercise capacity, breathlessness, and physical performance. The authors commented that a higher dose of the extract administered over a longer period was necessary for further investigation of
the observed improvements.\(^{(23)}\) Clinical studies with standardised extracts of leaf with flowers have demonstrated beneficial effects in patients with cardiac insufficiency.\(^{(G52)}\)
Side–effects, Toxicity

Nausea\(^{(20)}\) and fatigue, sweating and rash on the hands\(^{(23)}\) have been reported as side–effects in clinical trials using commercial preparations of hawthorn.\(^{(20)}\)

General symptoms of acute toxicity observed in a number of animal models (e.g. guinea–pig, frog, tortoise, cat, rabbit, rat) have been documented as bradycardia and respiratory depression leading to cardiac arrest and respiratory paralysis.\(^{(8–10)}\) Acute toxicity (LD\(_{50}\)) of isolated constituents (mainly flavonoids) has been documented as 50–2600 mg/ kg (by intravenous injection) and 6 g/kg (by mouth) in various animal preparations.\(^{(8–10)}\) The documented acute toxicity of commercial hawthorn preparations has also been reviewed.\(^{(8–10)}\)
Contra–indications, Warnings

Hawthorn has been reported to exhibit many cardiovascular activities and as such may affect the existing therapy of patients with various cardiovascular disorders such as hypertension, hypotension and cardiac disorders. These patient groups are likely to be most susceptible to the pharmacological actions of hawthorn.

Pregnancy and lactation

In vivo and in vitro utero–activity (reduction in tone and motility) has been documented for hawthorn extracts.\(^{(8–10)}\) In view of the pharmacological activities described for hawthorn, it should not be taken during pregnancy and lactation.
Pharmaceutical Comment

Hawthorn is characterised by its phenolic constituents, in particular the flavonoid components to which many of the pharmacological properties associated with hawthorn have been attributed. Pharmacological actions documented in both animal and human studies support the traditional actions of hawthorn and include cardioactive, hypotensive and coronary vasodilator.\(^{(24)}\) Separate monographs for the fruit (berries) and leaf with flowers appear in the British Pharmacopoeia and European Pharmacopoeia.\(^{(G15 \ G28)}\) The German Commission E did not approve the use of fruit for therapeutic purposes on the grounds of insufficient evidence. There is some evidence from clinical trials to support the use of the standardised extracts of leaf with flower for cardiac insufficiency. In view of the nature of the actions documented for hawthorn, it is not suitable for self-medications.
References

See also General References G1 G2 G3 G9 G15 G16 G18 G22 G28 G31 G32 G36 G43 G50 G52 G56 G62 G64.

15. Thompson EB et al. Preliminary study of potential antiarrhythmic effects


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Holy Thistle
Species (Family)

*Cnicus benedictus* L. (Asteraceae/Compositae)
Synonym(s)
Blessed Thistle, Carbenia Benedicta, Carduus Benedictus, Cnicus
Part(s) Used

Herb
Pharmacopoeial and Other Monographs

BHC 1992(G6)

BHP 1996(G9)

Complete German Commission E(G3)

Martindale 33rd edition(G67)

PDR for Herbal Medicines 2nd edition(G36)
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents
See General References G2 G6 G30 G40 G62 G64.

Lignans
Arctigenin, nortracheloside, 2–acetyl nortracheloside and trachelogenin.\(^{(1)}\)

Polyenes
Several polyacetylenes.\(^{(2)}\)

Steroids
Phytosterols (e.g. n-nonacosan, sitosterol, sitosteryl glycoside, stigmasterol).\(^{(3)}\)

Tannins
Type unspecified (8%).

Terpenoids
Sesquiterpenes including cnicin 0.2–0.7\%,\(^{(4)}\) yielding salonitenolide as aglycone,\(^{(5)}\) and artemisiifolin. Shoot and flowering head are reported to be devoid of cnicin.\(^{(4)}\) Triterpenoids including α-amyrenone, α-amyrin acetate, α-amyrine, multiflorenol, multiflorenol acetate and oleanolic acid.\(^{(3)}\)

Volatile oils
Many components, mainly hydrocarbons.\(^{(6)}\)

Other constituents
Lithospermic acid, mucilage, nicotinic acid and nicotinamide complex, resin.
Food Use

Holy thistle is listed by the Council of Europe as natural source of food flavouring (category N2). This category indicates that holy thistle can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\(^{(G16)}\) In the USA, holy thistle is permitted for use in alcoholic beverages.\(^{(G65)}\)
Holy thistle is stated to possess bitter stomachic, antidiarrhoeal, antihaemorrhagic, febrifuge, expectorant, antibiotic, bacteriostatic, vulnerary and antiseptic properties. Traditionally, it has been used for anorexia, flatulent dyspepsia, bronchial catarrh, topically for gangrenous and indolent ulcers, and specifically for atonic dyspepsia, and enteropathy with flatulent colic.
Dosage

*Dried flowering tops*
1.5–3.0 g or by infusion three times daily.\(^{G6 \ G7}\)

*liquid extract*
1.5–3.0 mL (1 : 1 in 25% alcohol) three times daily.\(^{G6 \ G7}\)
Pharmacological Actions

In vitro and animal studies

Antibacterial activity has been reported for an aqueous extract of the herb, for cnicin, and for the volatile oil.\(^{6–9}\) Activity has been documented against *Bacillus subtilis*, *Brucella abortus*, *Brucella bronchiseptica*, *Escherichia coli*, *Proteus* species, *Pseudomonas aeruginosa*, *Staphylococcus aureus* and *Streptococcus faecalis*. The antimicrobial activity of holy thistle has been attributed to cnicin and to the polyacetylene constituents.\(^{9}\)

Cnicin has exhibited *in vivo* anti-inflammatory activity (carrageenan-induced rat-paw oedema test) virtually equipotent to indometacin.\(^{4}\) Antitumour activity has been documented in mice against sarcoma 180 for the whole herb,\(^{8}\) and against lymphoid leukaemia for cnicin;\(^{8}\) cnicin has also been reported to exhibit *in vitro* activity against KB cells.\(^{8}\) An \(\alpha\)-methylene-\(\gamma\)-lactone moiety is thought to be necessary for the antibacterial and antitumour activities of cnicin.\(^{8}\)

Lithospermic acid is thought to be responsible for the antigonadotrophic activity documented for holy thistle.\(^{G30}\) The sesquiterpene lactone constituents are stated to be bitter principles.\(^{G62}\)

Tannins are generally known to possess astringent properties.
Side–effects, Toxicity

None documented for holy thistle. The toxicity of cnicin has been studied in mice: the acute oral LD$_{50}$ was stated to be 1.6–3.2 mmol/kg body weight and intraperitoneal administration was reported to cause irritation of tissue. In the writhing test, cnicin was found to cause abdominal pain with an ED$_{50}$ estimated as 6.2 mmol/kg.$^{(4)}$

Antitumour activity has been documented for the whole herb and for cnicin (see *In vitro* and animal studies).
Contra–indications, Warnings

None documented for holy thistle. Plants containing sesquiterpene lactones with an α-methylene-γ-lactone moiety are generally considered to be allergenic, although no documented hypersensitivity reactions to holy thistle were located. Holy thistle may cause an allergic reaction in individuals with a known hypersensitivity to other members of the Compositae (e.g. chamomile, ragwort, tansy).

Pregnancy and lactation
The safety of holy thistle has not been established. In view of the lack of toxicity data, excessive use of holy thistle during pregnancy and lactation should be avoided.
Pharmaceutical Comment

The chemistry of holy thistle is well documented and the available pharmacological data support most of the stated herbal uses, although no references to human studies were located. In view of the lack of toxicity data, excessive use of holy thistle should be avoided.
References

See also General References G2 G3 G6 G9 G16 G30 G31 G36 G37 G40 G43 G56 G62 G64.


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Species (Family)

*Humulus lupulus* L. (Cannabinaceae/Moraceae)
Synonym(s)
Humulus, Lupulus
Part(s) Used
Strobile
Pharmacopoeial and Other Monographs

BHC 1992\textsuperscript{(G6)}

BHP 1996\textsuperscript{(G9)}

BP 2002\textsuperscript{(G71)}

Complete German Commission E\textsuperscript{(G3)}

ESCOP 1997\textsuperscript{(G52)}

Martindale 33rd edition\textsuperscript{(G67)}

PDR for Herbal Medicines 2nd edition\textsuperscript{(G36)}

Ph Eur 2004\textsuperscript{(G72)}
Legal Category (Licensed Products)

GSL\textsuperscript{(G37)}
Constituents

**Flavonoids**
Astragalin, kaempferol, quercetin, quercitrin and rutin.

**Chalcones**
Isoxanthohumol, xanthohumol, 6-isopentenylnaringenin, 3′-(isoprenyl)-2′,4-dihydroxy-4′,6′-dimethoxychalcone, 2′,6′-dimethoxy-4,4′-dihydroxychalcone.\(^1\)

**Oleo–resin**
15–30%. Bitter principles (acylphloroglucides) in a soft and hard resin. The lipophilic soft resin consists mainly of α-acids (e.g. humulone, cohumulone, adhumulone, prehumulone, posthumulone), β-acids (e.g. lupulone, colupulone, adlupulone), and their oxidative degradation products including 2-methyl-3-buten-2-ol\(^2,3,G52\) The hard resin contains a hydrophilic δ-resin and χ-resin.

**Tannins**
2–4%. Condensed; galloatechin identified.\(^4\)

**Volatile oils**
0.3–1.0%. More than 100 terpenoid components identified; primarily (at least 90%) β-caryophyllene, farnesene and humulene (sesquiterpenes), and myrcene (monoterpene).

**Other constituents**
Amino acids, phenolic acids, gamma-linoleic acids, lipids and oestrogenic substances (disputed).\(^5\)

It has been stated that only low amounts of 2-methyl-3-buten-2-ol, the sedative principle identified in hops, are present in sedative tablets containing hops.\(^2\) However, it is thought that 2-methyl-3-buten-2-ol is formed *in vivo* by metabolism of the α-bitter acids and, therefore, the low amount of 2-methyl-3-buten-2-ol in a preparation may not indicate low sedative activity.\(^6\) Interestingly, relatively high concentrations of 2-methyl-3-buten-2-ol were found in bath preparations, suggesting that high concentrations of 2-methyl-3-buten-2-ol may be achieved in both tea and bath products containing hops.\(^2\)
Food Use

Hops are listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that hops can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\(^{(G16)}\) In the USA, hops is listed as GRAS (Generally Recognised As Safe).\(^{(G65)}\)
Hops are stated to possess sedative, hypnotic and topical bactericidal properties. Traditionally, they have been used for neuralgia, insomnia, excitability, priapism, mucous colitis, topically for crural ulcers, and specifically for restlessness associated with nervous tension headache and/or indigestion. The German Commission E approved use for mood disturbances such as restlessness and anxiety as well as sleep disturbances. Hops are used in combination with valerian root for nervous sleeping disorders and conditions of unrest.
Dosage

*Dried strobile*

0.5–1.0 g or by infusion; 1–2 g as a hypnotic.

*Liquid extract*

0.5–2.0 mL (1 : 1 in 45% alcohol).

*Tincture*

1–2 mL (1 : 5 in 60% alcohol).
Pharmacological Actions

*In vitro* and animal studies

Antibacterial activity, mainly against Gram–positive bacteria, has been documented for hops, and attributed to the humulone and lupulone constituents.(7) The activity of the bitter acids against Gram–positive bacteria is thought to involve primary membrane leakage. Resistance of Gram–negative bacteria to the resin acids is attributed to the presence of a phospholipid–containing outer membrane, as lupulone and humulone are inactivated by serum phospholipids.(7) Structure–activity studies have indicated the requirement of a hydrophobic molecule and a six–membered central ring for such activity.(8)

The humulones and lupulones are thought to possess little activity towards fungi or yeasts. However, antifungal activity has been documented for the bitter acids towards *Trichophyton*, *Candida*, *Fusarium* and *Mucor* species.(9) Flavonone constituents have also been documented to possess antifungal activity towards *Trichophyton* and *Mucor* species, and antibacterial activity towards *Staphylococcus aureus*. (10)

Antispasmodic activity has been documented for an alcoholic hop extract on various isolated smooth muscle preparations.(11) Hops have been reported to exhibit hypnotic and sedative properties.(G41) 2-Methyl–3–buten–2–ol, a bitter acid degradation product, has been identified as a sedative principle in hops.(2,3) 2-Methyl–3–buten–2–ol has been shown to possess narcotic properties in mice and motility depressant activity in rats, with the latter not attributable to a muscle–relaxant effect.(12) It has also been suggested that isovaleric acid residues present in hops may contribute towards the sedative action. In mice, hops extract administered intraperitoneally (100, 250, 500 mg/kg) 30 minutes prior to a series of behavioural tests, resulted in a dose–dependent suppression of spontaneous locomotor at doses of 250 mg/kg for up to one hour.(13) The time for mice to be able to remain on a rota rod was decreased by 59 and 65% at doses of 250 mg/kg and 500 mg/kg, respectively. The time of onset of convulsions and survival time after administration of pentylentetrazole (100 mg/kg) was significantly lengthened. Hops extract (35 mg/kg, intraperitoneal administration) produced a dose–dependent increase in sleeping time in mice treated with pentobarbitol. An antinociceptive effect was noted by increased latency of licking forepaws in hotplate tests and hypothermic activity observed from a time–dependent fall of rectal temperature at a dose of 500 mg/kg.(13)

Hops have previously been reported to possess oestrogenic constituents.(5)
However, when a number of purified components, including the volatile oil and the bitter acids, were examined using the uterine weight assay in immature female mice, no oestrogenic activity was found.\(^5\)

**Clinical studies**

Clinical studies have generally assessed hops given in combination with one or more additional herbs. For example, hops has been reported to improve sleep disturbances when given in combination with valerian.\(^{14}\)

Hops, in combination with chicory and peppermint, has been documented to relieve pain in patients with chronic cholecystitis (calculous and non-calculous).\(^{15}\) A herbal product containing a mixture of plant extracts, including hops and uva-ursi, and alpha-tocopherol acetate was reported to improve irritable bladder and urinary incontinence.\(^{16}\)
Side–effects, Toxicity

Respiratory allergy caused by the handling of hop cones have been documented;\(^{(17)}\) a subsequent patch test using dried, crushed flowerheads proved negative. Positive patch test reactions have been documented for fresh hop oil, humulone, and lupulone. Myrcene, present in the fresh oil but readily oxidised, was concluded to be the sensitising agent in the hop oil.\(^{(G51)}\) Contact dermatitis to hops has long been recognised\(^{(G51)}\) and is attributed to the pollen.\(^{(G41)}\) Small doses of hops are stated to be non–toxic.\(^{(G42)}\) Large doses administered to animals by injection have resulted in a soporific effect followed by death, with chronic administration resulting in weight loss before death.\(^{(G39)}\)
Contra-indications, Warnings

It has been stated that hops should not be taken by individuals suffering from depressive illness, as the sedative effect may accentuate symptoms.\(^{(G45 \ G49)}\) The sedative action may potentiate the effects of existing sedative therapy and alcohol. Allergic reactions have been reported for hops, although only following external contact with the herb and oil. Reports of oestrogenic activity are inconclusive.\(^{(G52)}\) It is claimed that hops are not oestrogenic,\(^{(G56)}\) but hop flower is said to have oestrogenic–binding activity and physiological oestrogenic effects.\(^{(18)}\) Concern has been expressed that herbs with oestrogenic effects, including hops, may stimulate breast cancer growth and oppose action of competitive oestrogen receptor antagonists such as tamoxifen.\(^{(18)}\)

**Pregnancy and lactation**

*In vitro* antispasmodic activity on the uterus has been documented. In view of this and the lack of toxicity data, the excessive use of hops during pregnancy and lactation should be avoided.
Pharmaceutical Comment

The chemistry of hops is well documented and is characterised by the bitter acid components of the oleo–resin. Documented pharmacological activities justify the herbal uses, although excessive use should be avoided in view of the limited toxicity data.
References


7. Teuber M, Schmalreck AF. Membrane leakage in *Bacillus subtilis* 168 induced by the hop constituents lupulone, humulone, isohumulone and humulinic acid. *Arch Mikrobiol* 1973; **94**: 159–171. (PubMed)


Horehound, Black
Species (Family)

*Ballota nigra* L. (Labiatae)
Synonym(s)

Ballota
Part(s) Used

Herb
Pharmacopoeial and Other Monographs

BHP 1996$^\text{G9}$

BP 2002$^\text{G71}$

PDR for Herbal Medicines 2nd edition$^\text{G36}$
Legal Category (Licensed Products)

Black horehound is not included in the GSL.\textsuperscript{(G37)}
Constituents

See General References G49 G64.

Limited chemical information is available for black horehound. Documented constituents include diterpenes (e.g. ballotenol, balltinone, preleosibirin),\(^1\) flavonoids, and volatile oil.
Food Use

Black horehound is listed by the Council of Europe as a natural source of food flavouring (category N3). This category indicates that black horehound can be added to foodstuffs in the traditionally accepted manner, although insufficient information is available for an adequate assessment of potential toxicity.\(^{(G16)}\)
Herbal Use

Black horehound is stated to possess anti-emetic, sedative and mild astringent properties. Traditionally, it has been used for nausea, vomiting, nervous dyspepsia, and specifically for vomiting of central origin.\(^{(G7G64)}\)
**Dosage**

**Dried herb**
2–4 g or by infusion three times daily.\(^{(G7)}\)

**Liquid extract**
1–3 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)

**Tincture**
1–2 mL (1 : 10 in 45% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

None documented.
Side-effects, Toxicity

None documented.
Contra–indications, Warnings

None documented.

Pregnancy and lactation
Black horehound is reputed to affect the menstrual cycle.\(^{(G30)}\) In view of the lack of phytochemical, pharmacological and toxicity data, the use of black horehound during pregnancy and lactation should be avoided.
Pharmaceutical Comment

Limited phytochemical or pharmacological information is available for black horehound to justify its use as a herbal remedy. In view of the lack of toxicity data, excessive use should be avoided.
Reference

See also General References G9 G16 G30 G36 G37 G49 G64.

Horehound, White
Species (Family)

*Marrubium vulgare* L. (Labiatae)
Synonym(s)
Common Hoarhound, Hoarhound, Horehound, Marrubium
Part(s) Used

Flower, leaf
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

Complete German Commission E\(^{(G3)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL (G37)
Constituents

See General References G2 G6 G41 G62 G64.

**Alkaloids**

Pyrrolidine–type. Betonicine 0.3%, the cis-isomer turicine.

**Flavonoids**

Apigenin, luteolin, quercetin, and their glycosides.\(^{(1)}\)

**Terpenoids**

Diterpenes including marrubiin 0.3–1.0%, a lactone, as the main component with lesser amounts of various alcohols (e.g. marrubenol, marrubiol, peregrinol and vulgarol). Marrubiin has also been stated to be an artefact formed from a precursor, premarrubiin, during extraction.\(^{(2)}\)

**Volatile oils**

Trace. Bisabolol, camphene, \(p\)-cymene, limonene, \(\beta\)-pinene, sabinene and others,\(^{(2)}\) a sesquiterpene (unspecified).

**Other constituents**

Choline, saponin (unspecified), \(\beta\)-sitosterol (a phytosterol), waxes (C\(_{26}\)-C\(_{34}\) alkanes).
Food Use

White horehound is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that white horehound can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\textsuperscript{(G16)} In the USA, white horehound is listed as GRAS (Generally Recognised As Safe).\textsuperscript{(G65)}
Herbal Use

White horehound is stated to possess expectorant and antispasmodic properties. Traditionally, it has been used for acute or chronic bronchitis, whooping cough, and specifically for bronchitis with non-productive cough. (G2 G6 G7 G8 G64)
Dosage

Dried herb
1–2 g or by infusion three times daily.\(^{(G6 \ G7)}\)

Liquid extract
2–4 mL (1 : 1 in 20\% alcohol) three times daily.\(^{(G6 \ G7)}\)
Pharmacological Actions

**In vitro and animal studies**

Aqueous extracts have been reported to exhibit an antagonistic effect towards hydroxytryptamine *in vivo* in mice, and *in vitro* in guinea–pig ileum and rat uterus tissue.\(^{(3)}\) Expectorant and vasodilative properties have been documented for the volatile oil.\(^{(4)}\) However, the main active expectorant principle in white horehound is reported to be marrubiin, which is stated to stimulate secretions of the bronchial mucosa.\(^{(G60)}\) Marrubiin has also been stated to be cardioactive, possessing anti–arrhythmic properties, although higher doses are reported to cause arrhythmias.\(^{(G60)}\) Marrubin acid (obtained from the saponification of marrubiin) has been documented to stimulate bile secretion in rats, whereas marrubiin was found to be inactive.\(^{(5)}\) White horehound is stated to possess bitter properties (BI 65 000 compared to gentian BI 10 000–30 000) with marrubiin as the main active component.\(^{(G62)}\)

Large doses of white horehound are purgative.\(^{(G10 \ G60)}\) The volatile oil has antischistosomal activity.\(^{(6)}\)
Side–effects, Toxicity

The plant juice of white horehound is stated to contain an irritant principle, which can cause contact dermatitis.\(^{(G51)}\) No documented toxicity studies were located for the whole plant, although an LD\(_{50}\) (rat, by mouth) value for marrubin acid is reported as 370 mg/kg body weight.\(^{(5)}\) The volatile oil is documented to be highly toxic to the flukes \textit{Schisto soma mansoni} and \textit{Schistosoma haematobium}.\(^{(6)}\)
Contra–indications, Warnings

None documented. Cardioactive properties and an antagonism of 5–hydroxytryptamine have been documented in animals.

Pregnancy and lactation

White horehound is reputed to be an abortifacient and to affect the menstrual cycle.\(^\text{G30}\) Uterine stimulant activity in animals has been documented.\(^\text{G30}\) In view of this and the lack of safety data, the use of white horehound during pregnancy should be avoided. Excessive use during lactation should be avoided.
The chemistry of white horehound is well documented. Limited pharmacological information is available, although expectorant properties have been reported which support some of the herbal uses. In view of the lack of toxicity data and suggested cardioactive properties, white horehound should not be taken in excessive doses.
References


Horse-chestnut
Species (Family)

*Aesculus hippocastanum* L. (Hippocastanaceae)
Synonym(s)

Aesculus
Part(s) Used

Seed
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)

ESCOP 1999\(^{(G52)}\)

Martindale 33rd edition\(^{(G67)}\)

Mills and Bone\(^{(G50)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL (for external use only)\(^{(G37)}\)
Constituents
See General References G22 G48 G52 G59 G62 G64.

**Coumarins**
Aesculetin, fraxin (fraxetin glucoside), scopolin (scopoletin glucoside).

**Flavonoids**
Flavonol (kaempferol, quercetin) glycosides including astragalin, isoquercetrin, rutin; leucocyanidin (quercetin derivative).

**Saponins**
French pharmacopoeial standard, not less than 3% aescin. A mixture of saponins collectively referred to as ‘aescin’ (3–10%); α- and β-escin as major glycosides.

**Tannins**
Type unspecified but likely to be condensed in view of the epicatechin content (formed during hydrolysis of condensed tannins).

**Other constituents**
Allantoin, amino acids (adenine, adenosine, guanine), choline, citric acid, phytosterol.
Food Use

Horse-chestnut is not used in foods.
Traditionally, horse-chestnut has been used for the treatment of varicose veins, haemorrhoids, phlebitis, diarrhoea, fever and enlargement of the prostate gland. The German Commission E approved use for treatment of chronic venous insufficiency in the legs.
Dosage

**Fruit**
0.2–1.0 g three times daily.\(^{(G49)}\)

**Preparations**
Extracts equivalent to 50–150 mg triterpenes calculated as aescin.\(^{(G52)}\)
Pharmacological Actions

Documented studies have concentrated on the actions of the saponins, in particular, aescin.

**In vitro and animal studies**

**Anti–inflammatory and anti–oedema effects**

Anti–inflammatory activity in rats has been documented for both a fruit extract and the saponin fraction.\(^1\)–\(^4\) Anti–inflammatory activity in the rat has been reported to be greater for a total horse–chestnut extract compared to aescin. In addition, an extract excluding aescin also exhibited activity, suggesting that horse–chestnut contains anti–inflammatory agents other than aescin.\(^5\) No difference in activity was noted when the horse–chestnut extracts were administered prior to and after dextran (inflammatory agent). It has been proposed that aescin affects the initial phase of inflammation by exerting a ‘sealing’ effect on capillaries and by reducing the number and/or diameter of capillary pores.\(^3\)

**Effects on venous tone**

Horse–chestnut extract (16% aescin, 0.2 mg/mL) and also aescin (0.1 mg/mL) induced contractions in isolated bovine and human veins.\(^6\) Concentration–dependent contractions of isolated canine veins were observed with a horse–chestnut extract (16% aescin, 5 × 10\(^{-4}\) mg/mL).\(^7\) A standardised extract (16% aescin, 50 mg, given intravenously) increased femoral venous pressure in anaesthetised dogs, and decreased cutaneous capillary hyperpermeability in rats (200 mg/kg, given orally).\(^8\)

In addition, the saponin fraction has been reported to exhibit analgesic and antigranulation activities in rats,\(^3\) to reduce capillary permeability,\(^6\) and to produce an initial hypotension followed by a longer lasting hypertension in anaesthetised animals.\(^4\) Prostaglandin production by venous tissue is thought to be involved in the regulation of vascular reactivity.\(^7\)

Prostaglandins of the E series are known to cause relaxation of venous tissues whereas those of the \(F_\alpha\) series produce contraction. Increased venous tone induced by aescin \emph{in vitro} was found to be associated with an increased \(PGF_{2\alpha}\) synthesis in the venous tissue.

**Other activities**

\emph{In vitro}, aescin has been documented to inhibit hyaluronidase activity (IC\(_{50}\) 150 μmol/L).\(^9\)
A saponin fraction of horse-chestnut has been reported to contract isolated rabbit ileum.\(^3\)

Antiviral activity \textit{in vitro} against influenza virus (A\(_2\)/Japan 305) has been described for aescin.\(^8\)

Metabolism studies of aescin in the rat have concluded that aescin toxicity is reduced by hepatic metabolism.\(^9\)

Flavonoids and tannins are generally recognised as having anti-inflammatory and astringent properties, respectively.

**Clinical studies**

**Chronic venous insufficiency**

Several studies have assessed the effects of horse-chestnut seed extract in patients with chronic venous insufficiency, a common condition which causes oedema of the lower leg.

A systematic review of randomised, double-blind, controlled trials of horse-chestnut seed extract in chronic venous insufficiency included 13 studies (eight placebo-controlled trials and five studies comparing horse-chestnut seed extract with reference medication or compression therapy).\(^{10}\)

Generally, trials involved the administration of horse-chestnut seed extract 100 or 150 mg daily for 3–12 weeks. The results of all the placebo-controlled studies indicated that horse-chestnut seed extract was superior. Four comparative studies indicated that horse-chestnut seed extract was as effective as \(O\)-(β-hydroxyethyl)-rutosides in relieving symptoms of chronic venous insufficiency; one study suggested that horse-chestnut seed extract was as effective as compression therapy. It was concluded that horse-chestnut seed extract is effective as a symptomatic, short-term treatment for chronic venous insufficiency, but that further well-designed clinical trials are required to confirm this.\(^{10}\)

**Other effects**

Glycosaminoglycan hydrolysases are enzymes involved in the breakdown of substances (proteoglycans) that determine capillary rigidity and pore size (thus influencing the passage of macromolecules into the surrounding tissue). Proteoglycans also interact with collagen, stabilising the fibres and regulating their correct biosynthesis.\(^{11}\) The activity of these enzymes was found to be raised in patients with varicosis, compared with healthy patients. In a study involving 15 patients with varicosis treated with horse-chestnut extract (900 mg daily) for 12 days, the activity of these enzymes was significantly
reduced.\textsuperscript{(11)} It was proposed that horse–chestnut may act at the site of enzyme release, exerting a stabilising effect on the lysosomal membrane.\textsuperscript{(11)}

In a randomised, double-blind, placebo-controlled study involving 70 healthy individuals with haematomas, a topical gel (2\% aescin) reduced sensitivity to pressure on affected areas.\textsuperscript{(G52)}

The cosmetic applications of horse–chestnut have been reviewed;\textsuperscript{(12)} these effects are attributed to properties associated with the saponin constituents.
Side-effects, Toxicity

Two incidences of toxic nephropathy have been reported and were stated as probably secondary to the ingestion of high doses of aescin.\(^{13}\) In Japan, where horse-chestnut has been used as an anti-inflammatory drug after surgery or trauma, hepatic injury has been described in a male patient who received an intramuscular injection of a proprietary product containing horse-chestnut.\(^{14}\) Liver function tests showed a mild abnormality and a diagnosis of giant cell tumour of bone (grade 2) by bone biopsy was made. Other side-effects stated to have been reported for the product include shock, spasm, mild nausea, vomiting and urticaria.\(^{14}\)

The effect of aescin, both free and albumin-bound, on renal tubular transport processes has been studied in the isolated, artificially perfused frog kidney.\(^{15}\) Aescin was found to primarily affect tubular, rather than glomerular, epithelium and it was noted that binding to plasma protein (approximately 50\%) protects against this nephrotoxicity. Aescin was thought to be neither secreted nor reabsorbed in the tubules, and the concentration of unbound aescin filtered through the kidney (13\%) was considered to be too low to have toxic effects. The authors commented that the symptoms of acute renal failure in humans are caused primarily by interference with glomeruli and in view of this, the nephrotoxic potential of aescin is probably only relevant when the kidneys are already damaged and also if the aescin is displaced from its binding to plasma protein.\(^{15}\)

A proprietary product containing horse-chestnut (together with phenopyrazone and cardiac glycoside-containing plant extracts) has been associated with the development of a drug-induced auto-immune disease called ‘pseudolupus syndrome’ in Germany and Switzerland.\(^{16,17}\) The individual component in the product responsible for the syndrome was not established.

It has been noted that death occurs rapidly in animals given large doses of aescin, due to massive haemolysis. Death is more prolonged in animals given smaller doses of aescin.\(^{4}\)

LD\(_{50}\) values for aescin have been estimated in mice, rats and guinea-pigs and range from 134 to 720 mg/kg (by mouth) and from 1.4 to 15.2 mg/kg (intravenous injection).\(^{G49}\) The total saponin fraction has been reported to be less toxic in mice (intraperitoneal injection) compared to the isolated aescin mixture (LD\(_{50}\) 46.5 mg/kg and 9.5 mg/kg, respectively).\(^{3}\) The haemolytic index of horse-chestnut is documented as being 6000, compared with 9500 to 12 500 for aescin.\(^{G62}\) Daily doses in rats (100 mg/kg, orally) of
a standardised extract of horse–chestnut (16% aescin) did not produce teratogenic effects, and the extract was negative in the Ames test with *Salmonella typhimurium* TA98 without actuation.\(^{G52}\)
Contra–indications, Warnings

Horse–chestnut may be irritant to the gastrointestinal tract due to the saponin constituents. Saponins are generally recognised to possess haemolytic properties, but are not usually absorbed from the gastrointestinal tract following oral administration. Horse–chestnut may interfere with anticoagulant/coagulant therapy (coumarin constituents). Aescin, the main saponin component in horse–chestnut, binds to plasma protein and may affect the binding of other drugs. Horse–chestnut should be avoided by patients with existing renal or hepatic impairment.

Pregnancy and lactation
The safety of horse–chestnut during pregnancy and lactation has not been established. In view of the pharmacologically active constituents present in horse–chestnut, use during pregnancy and lactation is best avoided.
Horse–chestnut is traditionally characterised by its saponin components, in particular aescin which represents a mixture of compounds. However, horse–chestnut also contains other pharmacologically active constituents including coumarins and flavonoids. The traditional use of horse–chestnut in peripheral vascular disorders has largely been substantiated by studies in animals and humans, in which anti–inflammatory and capillary stabilising effects have been observed. There is evidence from randomised, double–blind, controlled clinical trials to support the use of horse–chestnut seed extract in the treatment of symptoms of chronic venous insufficiency.

Many of the documented activities can probably be attributed to the saponin and flavonoid constituents in horse–chestnut.
References

See also General References G2 G9 G16 G22 G31 G36 G37 G43 G48 G49 G50 G52 G56 G59 G62 G64.

1. Farnsworth NR, Cordell GA. A review of some biologically active compounds isolated from plants as reported in the 1974–75 literature. *Lloydia* 1976; **39**: 420–455. ([PubMed](#))


15. Rothkopf M et al. Animal experiments on the question of the renal


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Horseradish
Species (Family)

*Radicula armoracia* (L.) Robinson (*Brassicaceae/ Cruciferae*)
Synonym(s)

Armoracia lopathifolia Gilib., A. rusticana (Gaertn.) Mey & Scherb., Cochlearia armoracia L., Nasturtium armoracia Fries, Roripa armoracia Hitch.
Part(s) Used

Root
Pharmacopoeial and Other Monographs

Complete German Commission E\(^{(G3)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL\textsuperscript{(G37)}
Constituents
See General References G40 G48 G49 G57 G58 G62 G64.

**Coumari**ns
Aesculetin, scopoletin.$^1$

**Phenols**
Caffeic acid derivatives and lesser amounts of hydroxycinnamic acid derivatives. Concentrations of acids are reported to be much lower in the root than in the leaf.$^1$

**Volatile oils**
Glucosinolates (mustard oil glycosides) gluconasturtiin and sinigrin ($S$-glucosides), yielding phenylethylisothiocyanate and allylisothiocyanate after hydrolysis. Isothiocyanate content estimated as 12.2–20.4 mg/g freeze dried root.$^{2,3}$ Other isothiocyanate types include isopropyl, 3–butenyl, 4–pentenyl, phenyl, 3–methylthiopropyl and benzyl derivatives.$^4$

**Other constituents**
Ascorbic acid, asparagin, peroxidase enzymes, resin, starch and sugar.

**Other plant parts**
Kaempferol and quercetin have been documented for the leaf.
Food Use

Horseradish is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that horseradish can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\(^{(G16)}\) In the USA, horseradish is listed as GRAS (Generally Recognised As Safe).\(^{(G57)}\) Horseradish is commonly used as a food flavouring.
Herbal Use

Horseradish is stated to possess antiseptic, circulatory and digestive stimulant, diuretic and vulnerary properties.\(^{(G42 G49 G64)}\) Traditionally, it has been used for pulmonary and urinary infection, urinary stones, oedematous conditions, and externally for application to inflamed joints or tissues.\(^{(G49)}\)
Dosage

*Root (fresh)*
2–4 g before meals.\(^{\text{G49}}\)
Pharmacological Actions

**In vitro and animal studies**

A marked hypotensive effect in cats has been documented for horseradish peroxidase, following intravenous administration.\(^{(5)}\) The effect was completely blocked by aspirin and indometacin, but was not affected by antihistamines. It was concluded that horseradish peroxidase acts by stimulating the synthesis of arachidonic acid metabolites.
Side–effects, Toxicity

Isothiocyanates are reported to have irritant effects on the skin and also to be allergenic.\(^{(G51 G58)}\) Animal poisoning has been documented for horseradish. Symptoms described include inflammation of the stomach or rumen, and excitement followed by collapse.\(^{(G33)}\)
Contra–indications, Warnings

It is stated that horseradish may depress thyroid function, and should be avoided by individuals with hypothyroidism or by those receiving thyroxine.\(^{(G42 G49)}\) No rationale for this statement is included except that this action is common to all members of the cabbage and mustard family.

**Pregnancy and lactation**

Allylisothiocyanate is extremely toxic and a violent irritant to mucous membranes.\(^{(G58)}\) Its use should be avoided during pregnancy and lactation.
Pharmaceutical Comment

The chemistry of horseradish is well established and it is recognised as one of the richest plant sources of peroxidase enzymes.\(^{G48}\) Little pharmacological information was located, although the isothiocyanates and peroxidases probably account for the reputed circulatory stimulant and wound-healing actions, respectively. The oil is one of the most hazardous of all essential oils and it is not recommended for either external or internal use.\(^{G58}\) Horseradish should not be ingested in amounts exceeding those used in foods.
References

See also General References G3 G10 G16 G31 G36 G40 G42 G48 G49 G51 G57 G58 G62 G64.


Hydrangea
Species (Family)

_Hydrangea arborescens_ L. (Saxifragaceae)
Synonym(s)
Mountain Hydrangea, Seven Barks, Smooth Hydrangea, Wild Hydrangea
Part(s) Used
Rhizome, Root
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL\textsuperscript{(G37)}
Constituents

See General References G22 G40 G41 G48 G64.

Limited information is available on the chemistry of hydrangea. It is stated to contain carbohydrates (e.g. gum, starch, sugars), flavonoids (e.g. kaempferol, quercetin, rutin), resin, saponins, hydrangin and hydrangenol, a stilbenoid,\(^1\) and to be free from tannins.
Food Use

Hydrangea is not used in foods. In the USA, hydrangea is listed as a ‘Herb of Undefined Safety’.(G22)
Herbal Use

Hydrangea is stated to possess diuretic and antilithic properties. Traditionally, it has been used for cystitis, urethritis, urinary calculi, prostatitis, enlarged prostate gland, and specifically for urinary calculi with gravel and cystitis.\(^{G7 G64}\)
Dosage

*Dried rhizome/root*
2–4 g or by decoction three times daily.\(^{(G7)}\)

*Liquid extract*
2–4 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)

*Tincture*
2–10 mL (1 : 5 in 45% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

*In vitro and animal studies*

None documented for hydrangea. Synthesised hydrangenol derivatives have been reported to possess anti-allergic properties, exhibiting a strong inhibitory action towards hyaluronidase activity and histamine release.\(^{(2)}\)
Side–effects, Toxicity

Hydrangea has been reported to cause contact dermatitis, (G51) and it is stated that hydrangin may cause gastroenteritis. (G22) Symptoms of overdose are described as vertigo and a feeling of tightness in the chest. (G22) An extract has been reported to be non–toxic in animals. (G64)
Contra–indications, Warnings

None documented.

Pregnancy and lactation
The safety of hydrangea has not been established. In view of the lack of phytochemical, pharmacological and toxicity data, the use of hydrangea during pregnancy and lactation should be avoided.
Pharmaceutical Comment

Limited information is available on the chemistry of hydrangea, although related species have been investigated more thoroughly.\(^{(G41)}\) No scientific evidence was located to justify the herbal uses. In view of the lack of toxicity data, excessive use of hydrangea should be avoided.

Hydrocotyle
Species (Family)

*Centella asiatica* (L.) Urban (Umbelliferae)
Synonym(s)

Centella, Gotu Kola, *Hydrocotyle asiatica*, Indian Pennywort, Indian Water Navelwort
Part(s) Used

Herb
Pharmacopoeial and Other Monographs

BHP 1983\(^{(G7)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)

WHO volume 1 1999\(^{(G63)}\)
Legal Category (Licensed Products)

GSL (for external use only)$^{(G37)}$
Constituents

See General References G22 G44 G60 G64.

**Amino acids**
Alanine and serine (major components), aminobutyrate, aspartate, glutamate, histidine, lysine and threonine.\(^{(1)}\) The root contains greater quantities than the herb.\(^{(1)}\)

**Flavonoids**
Quercetin, kaempferol and various glycosides.\(^{(2-4)}\)

**Terpenoids**
Triterpenes, asiaticoside, centelloside, madecassoside, brahmoside and brahminoside (saponin glycosides). Aglycones are referred to as hydrocotylegenin A–E;\(^{(5)}\) compounds A–D are reported to be esters of the triterpene alcohol \(R_1\)-barrigenol.\(^{(5,6)}\) Asiaticentoic acid, centellic acid, centoic acid and madecassic acid.

**Volatile oils**
Various terpenoids including β-caryophyllene, \(\text{trans-β-farnesene}\) and germacrene D (sesquiterpenes) as major components, \(\alpha\)-pinene and \(\beta\)-pinene. The major terpenoid is stated to be unidentified.

**Other constituents**
Hydrocotylin (an alkaloid), vallerine (a bitter principle), fatty acids (e.g. linoleic acid, linolenic acid, lignocene, oleic acid, palmitic acid, stearic acid), phytosterols (e.g. campesterol, sitosterol, stigmasterol),\(^{(7)}\) resin and tannin.

The underground plant parts of hydrocotyle have been reported to contain small quantities of at least 14 different polyacetylenes.\(^{(8-10)}\)
Food Use

Hydrocotyle is not used in foods.
Herbal Use

Hydrocotyle is stated to possess mild diuretic, antirheumatic, dermatological, peripheral vasodilator and vulnerary properties. Traditionally it has been used for rheumatic conditions, cutaneous affections, and by topical application, for indolent wounds, leprous ulcers, and cicatrisation after surgery. (G7 G64)
Dosage

*Dried leaf*
0.6 g or by infusion three times daily.\(^G_7\)
Pharmacological Actions

In vitro and animal studies

The triterpenoids are regarded as the active principles in hydrocotyle.\(^\text{(7)}\) Asiaticoside is reported to possess wound-healing ability, by having a stimulating effect on the epidermis and promoting keratinisation.\(^\text{(11)}\) Asiaticoside is thought to act by an inhibitory action on the synthesis of collagen and mucopolysaccharides in connective tissue.\(^\text{(11)}\)

Both asiaticoside and madecassoside are documented to be anti-inflammatory, and the total saponin fraction is reported to be active in the carrageenan rat paw oedema test.\(^\text{(12)}\)

In vivo studies in rats have shown that asiaticoside exhibits a protective action against stress-induced gastric ulcers, following subcutaneous administration,\(^\text{(13)}\) and accelerates the healing of chemical-induced duodenal ulcers, after oral administration.\(^\text{(14)}\) It was thought that asiaticoside acted by increasing the ability of the rats to cope with a stressful situation, rather than via a local effect on the mucosa.\(^\text{(13)}\)

In vivo studies in mice and rats using brahmoside and brahminoside, by intraperitoneal injection, have shown a CNS-depressant effect.\(^\text{(15)}\) The compounds were found to decrease motor activity, increase hexobarbitone sleeping time, slightly decrease body temperature, and were thought to act via a cholinergic mechanism.\(^\text{(15)}\) A hypertensive effect in rats was also observed, but only following large doses.\(^\text{(15)}\) In vitro studies with brahmoside and brahminoside indicated a relaxant effect on the rabbit duodenum and rat uterus, and an initial increase, followed by a decrease, in the amplitude and rate of contraction of the isolated rabbit heart.\(^\text{(15)}\) Higher doses were found to cause cardiac arrest, although subsequent intra venous administration in dogs caused no marked change in an ECG.\(^\text{(15)}\)

In vitro antifertility activity against human and rat sperm has been described for the total saponin fraction.\(^\text{(16)}\) Asiaticoside and brahminoside are thought to be the active components, although no spermicidal or spermostatic action could be demonstrated for the pure saponins.\(^\text{(16)}\) A crude hydrocotyle extract has been reported to significantly reduce the fertility of female mice when administered orally.\(^\text{(10)}\) No mechanism of action was investigated.

Teratogenicity studies in the rabbit have reported negative findings for a hydrocotyle extract containing asiatic acid, madecassic acid, madasiatic acid and asiaticoside.\(^\text{(17)}\)
Fresh plant juice is reported to be devoid of antibacterial activity,\(^{(18)}\) although asiaticoside has been reported to be active versus *Mycobacterium tuberculosis*, *Bacillus leprae* and *Entamoeba histolytica*, and oxyasiaticoside was documented to be active against tubercle bacillus.\(^{(16,18)}\) The fresh plant juice is also stated not to exhibit antitumour or antiviral activities, but to possess a moderate cytotoxic action in human ascites tumour cells.\(^{(18)}\)

**Clinical studies**

Several studies describing the use of hydrocotyle to treat wounds and various skin disorders have been documented. A cream containing a hydrocotyle extract was found to be successful in the treatment of psoriasis in seven patients to whom it was applied.\(^{(19)}\) An aerosol preparation, containing a hydrocotyle extract, was reported to improve the healing in 19 of 25 wounds that had proved refractory to other forms of treatment.\(^{(11)}\) A hydrocotyle extract containing asiaticoside (40%), asiatic acid (29–30%), madecassic acid (29–30%) and madasiatic acid (1%) was stated to be successful as both a preventive and curative treatment, when given to 227 patients with keloids or hypertrophic scars.\(^{(17)}\) The effective dose in adults was reported to be between 60 and 90 mg. It was proposed that the triterpene constituents in the hydrocotyle extract act in a similar manner to cortisone, with respect to wound healing, and interfere with the metabolism of abnormal collagen.\(^{(17)}\)

The triterpene constituents are reported to be metabolised primarily in the faeces in a period of 24–76 hours, with a small percentage metabolised via the kidneys.\(^{(17)}\) An extract containing asiatic acid, madecassic acid, madasiatic acid and asiaticoside reached peak plasma concentrations in 2–4 hours, irrespective of whether it is administered in tablet, oily injection or ointment formulations.\(^{(17)}\)

Hydrocotyle has been used in the treatment of patients with chronic lesions such as cutaneous ulcers, surgical wounds, fistulas and gynaecological wounds.\(^{(G44 G45)}\) Hydrocotyle has also been reported to improve the blood circulation in the lower limbs. Stimulation of collagen synthesis in the vein wall resulted in an increase in vein tonicity and a reduction in the capacity of the vein to distend.\(^{(10)}\)

The juice of the leaves or the whole plant is documented to be effective for relieving the itching associated with prickly heat.\(^{(G51)}\)

Asiaticoside has also been documented to improve the general ability and behavioural pattern of 30 mentally retarded children, when given over a period of 12 weeks. It also increased the mean concentrations of blood sugar,
serum cholesterol and total protein, and lowered blood urea and serum acid phosphatase concentrations, in 43 adults. Vital capacity was also increased.
Side–effects, Toxicity

A burning sensation was reported by 4 of 20 patients during the period of application of an aerosol preparation containing hydrocotyle.\(^{(11)}\) However, it is not clear whether other components in the formulation contributed to this reaction. Ingestion of hydrocotyle is stated to have produced pruritus over the whole body.\(^{(G51)}\)
Contra-indications, Warnings

It is stated that hydrocotyle may produce photosensitisation.\textsuperscript{(G7)} Excessive doses may interfere with existing hypoglycaemic therapy and increase serum cholesterol concentrations; hyperglycaemic and hypercholesterolaemic activities have been reported for asiaticoside in humans. Brahmoside and brahminoside have been reported to exert a CNS-depressant action in animal studies.\textsuperscript{(15)}

Pregnancy and lactation

Hydrocotyle is reputed to be an abortifacient and to affect the menstrual cycle.\textsuperscript{(G30)} Relaxation of the isolated rat uterus has been documented for brahmoside and brahminoside.\textsuperscript{(15)} Triterpene constituents have been reported to lack any teratological effects in rabbits.\textsuperscript{(17)} In view of the lack of toxicity data, the use of hydrocotyle during pregnancy should be avoided. Excessive use should be avoided during lactation.
Pharmaceutical Comment

The chemistry of hydrocotyle is well studied and its pharmacological activity seems to be associated with the triterpenoid constituents. Documented clinical and animal data support the herbal use of hydrocotyle as a dermatological agent, and warrants further research into the potential role of hydrocotyle in wound management. In view of the lack of toxicity data, excessive ingestion of hydrocotyle should be avoided.
References

See also General References G7 G22 G30 G31 G36 G37 G43 G51 G60 G63 G64.

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Ispaghula
Species (Family)

*Plantago ovata* Forsk. (Plantaginaceae)
Synonym(s)
Blond Psyllium, Indian Plantago, Ispagol, Pale Psyllium, Spogel
Part(s) Used

Seed, husk
Pharmacopoeial and Other Monographs

BHC 1992(G6)
BHP 1996(G9)
BP 2002(G71)
Complete German Commission E (Psyllium, Blonde)(G3)
ESCOP 1996(G52)
Martindale 33rd edition(G67)
PDR for Herbal Medicines 2nd edition(G36)
Ph Eur 2004(G72)
WHO volume 1 1999(G63)
Legal Category (Licensed Products)

GSL\((G37)\)
Constituents

See General References G2 G6 G41 G52 G59 G64.

Alkaloids
Monoterpen–type. (+)-Boschniakine (indicaine), (+)-boschniakinic acid (plantagonine) and indicainine.

Mucilages
10–30%. Mucopolysaccharide consisting mainly of a highly branched arabininoxylan with a xylan backbone and branches of arabinose, xylose and 2-O-(galacturonic)-rhamnose moieties. Present mainly in the seed husk.

Other constituents
Aucubin (iridoid glucoside), sugars (fructose, glucose, sucrose), planteose (trisaccharide), protein, sterols (campesterol, β-sitosterol, stigmasterol), triterpenes (α- and β-amyrin), fatty acids (e.g. linoleic, oleic, palmitic, stearic), tannins.
Food Use

In food manufacture, ispaghula may be used as a thickener or stabiliser.\(^{(G41)}\)
Ispaghula is stated to possess demulcent and laxative properties. Traditionally, ispaghula has been used in the treatment of chronic constipation, dysentery, diarrhoea and cystitis. Topically, a poultice has been used for furunculosis. The German Commission E approved use for chronic constipation and disorders in which bowel movements with loose stool are desirable, e.g. patients with anal fistulas, haemorrhoids, pregnancy, secondary medication in the treatment of various forms of diarrhoea and in the treatment of irritable bowel syndrome.\(^{(G3)}\)

The European Medicine Evaluation Agency (EMEA) Herbal Medicinal Products Working Group (HMPWG) has proposed a core SPC (Summary of Product Characteristics) for ispaghula.\(^{(G23)}\) The core SPC includes the following indications: (a) treatment of habitual constipation; conditions in which easy defecation with soft stools is desirable, e.g. in cases of painful defecation after rectal or anal surgery; (b) adjuvant symptomatic therapy in cases of diarrhoea from various causes; (c) conditions which need an increased daily fibre intake, e.g. as an adjuvant in irritable bowel syndrome.
Dosage

**Seeds**
5–10 g (3 g in children) three times daily;\(^{(G6 \ G7)}\) 12–40 g per day, husk 4–20 g;\(^{(G3)}\) 3–5 g.\(^{(G43)}\) Children 6–12 years, half adult dose. Children under 6 years, treat only under medical supervision.\(^{(G52)}\) Seeds should be soaked in warm water for several hours before taking.

*Liquid extract*
2–4 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G6 \ G7)}\)

**Husk**
3–5 g.\(^{(G46)}\) Seeds and husk should be soaked in warm water for several hours before administration.

7–11 g in one to three doses for indication (a) and (c); 7–20 g in one to three doses for indication (b).\(^{(G23)}\)
Pharmacological Actions

The principal pharmacological actions of ispaghula can be attributed to the mucilage component.

In vitro and animal studies

An alcoholic extract lowered the blood pressure of anaesthetised cats and dogs, inhibited isolated rabbit and frog hearts, and stimulated rabbit, rat and guinea–pig ileum.\(^{(G41)}\) The extract exhibited cholinergic activity.\(^{(G41)}\) A mild laxative action has also been reported in mice administered iridoid glycosides, including aucubin.\(^{(1)}\) Four–week supplementation of a fibre–free diet with isphagula seeds (100 or 200 g/kg) was compared with that of the husks and wheat bran in rats.\(^{(2)}\) The seeds increased faecal fresh weight by up to 100% and faecal dry weight by up to 50%. Total faecal bile acid secretion was stimulated, and β-glucuronidase activity reduced, by ispaghula. The study concluded that ispaghula acts as a partly fermentable, dietary fibre supplement increasing stool bulk, and that it probably has metabolic and mucosa–protective effects.

Ispaghula husk depressed the growth of chickens by 15% when added to their diet at 2%.\(^{(G41)}\)

Ispaghula seed powder is stated to have strongly counteracted the deleterious effects of adding sodium cyclamate (2%), FD & C Red No. 2 (2%), and polyoxyethylene sorbitan monostearate (4%) to the diet of rats.\(^{(G41)}\)

Clinical studies

Ispaghula is used as a bulk laxative.\(^{(G3 G43 G52)}\) The swelling properties of the mucilage enable it to absorb water in the gastrointestinal tract, thereby increasing the volume of the faeces and promoting peristalsis. Bulk laxatives are often used for the treatment of chronic constipation and when excessive straining must be avoided following ano rectal surgery or in the management of haemorrhoids. Ispaghula is also used in the management of diarrhoea and for adjusting faecal consistency in patients with colostomies and in patients with diverticular disease or irritable bowel syndrome.

Laxative effect

Ispaghula increases water content of stools and total stool weight in patients,\(^{(3)}\) thus promoting peristalsis and reducing mouth–to–rectum transit time.\(^{(4)}\) In a short–term study, 42 adults with constipation (≤3 bowel movements per week) received either ispaghula (7.2 g/day) or ispaghula plus
Both treatments increased defecation frequency, and wet and dry stool weights, improved stool consistency, and gave subjective relief.

A randomised, double-blind, double-dummy, multicentre study involved 170 subjects with chronic idiopathic constipation. The study included a two-week baseline (placebo) phase, followed by two weeks’ treatment with ispaghula (Metamucil) 5.1 g, twice daily or docusate sodium 100 mg twice daily. Compared with docusate, ispaghula significantly increased stool water content (0.01% versus 2.33% for docusate and ispaghula, respectively; \( p = 0.007 \)), and total stool output (271.9 g/week versus 359.9 g/week for docusate and ispaghula, respectively; \( p = 0.005 \)). Furthermore, bowel movement frequency was significantly greater for ispaghula, compared with docusate. It was concluded that ispaghula has greater overall laxative efficacy than docusate in patients with chronic constipation.

**Antidiarrhoeal effect**

An open, randomised, crossover trial involving 25 patients with diarrhoea compared the effects of loperamide with those of ispaghula and calcium. Nineteen patients completed both periods of treatment. The results indicated that both treatments halved stool frequency. Ispaghula and calcium were reported to be significantly better than loperamide with regard to urgency and stool consistency.

Nine volunteers with phenolphthalein-induced diarrhoea were treated in random sequence with placebo, ispaghula (Konsyl), calcium polycarbophyl or wheat bran. Wheat bran and calcium polycarbophyl had no effect on faecal consistency or on faecal viscosity. By contrast, ispaghula made stools firmer and increased faecal viscosity. In a dose-response study involving six subjects, 9, 18 and 30 g ispaghula per day caused a near-linear increase in faecal viscosity.

The effects of ispaghula have been explored in children. In an open, uncontrolled study, 23 children with chronic non-specific diarrhoea were treated with an unrestricted diet for one week, and then treated with ispaghula (Metamucil) for two weeks (one tablespoonful twice daily). Seven patients responded to the unrestricted diet and 13 were said to respond to ispaghula treatment.

**Hypocholesterolaemic effects**

In a double-blind, placebo-controlled, parallel-group study, 26 men with mild-to-moderate hypercholesterolaemia (serum cholesterol concentration: 4.86–8.12 mmol/L) received ispaghula (Metamucil) 3.4 g, or cellulose...
placebo, three times daily at meal times for eight weeks. At the end of the study, serum cholesterol concentrations were reduced by 14.8% in the treated group, low-density lipoprotein (LDL) cholesterol by 20.2% and ratio of LDL to high-density lipoprotein (HDL) cholesterol by 14.8%, compared with baseline values. There were no significant changes in serum lipid concentrations with placebo treatment, compared with baseline values. Differences in serum cholesterol concentrations between the two groups were statistically significant after four weeks (p-value not reported).

A double-blind, placebo-controlled, parallel trial study compared the effects of ispaghula (Metamucil 5.1 g, daily) and placebo in 118 patients (aged 21–70 years old) with primary hypercholesterolaemia (total serum cholesterol ≥5.7 mmol/L). Thirty-seven participants maintained a high-fat diet and 81 a low-fat diet. Treated patients in both low- and high-fat diet groups showed small significant decreases (p < 0.05) in total cholesterol and LDL cholesterol levels (5.8 and 7.2%, respectively, for high-fat diets; 4.2% and 6.4%, respectively, for low-fat diets). No significant differences were seen in LDL cholesterol response for treated patients on either diet.

In a randomised, double-blind, crossover study, 20 males (mean (SD) age 44 (4) years) with moderate hypercholesterolaemia (mean (SD) total cholesterol concentration 265 (17) mg/dL, LDL 184 (15) mg/dL) were randomised to receive a 40-day course of ispaghula (Metamucil) 15 g daily, or placebo (cellulose). There was a wash-out period of more than 10 days between treatments. Ispaghula lowered LDL cholesterol (168 mg/dL) more than did cellulose placebo (179 mg/dL), decreased relative cholesterol absorption, and increased the fractional turnover of both chenodeoxycholic acid and cholic acid. Bile acid synthesis increased in subjects whose LDL cholesterol was lowered by more than 10%. It was concluded that ispaghula lowers LDL cholesterol primarily by stimulation of bile acid synthesis.

A meta-analysis of eight published and four unpublished studies carried out in four countries reviewed the effect of consumption of ispaghula-enriched cereal products on blood cholesterol, and LDL and HDL cholesterol concentrations. Overall, the trials included 404 adults with mild-to-moderate hypercholesterolaemia (5.17–7.8 mmol/L) who consumed low-fat diets. The meta-analysis indicated that subjects who consumed ispaghula cereal had lower total cholesterol and LDL cholesterol than subjects who ate control cereal concentrations (differences of 0.31 mmol/L (5%) and 0.35 mmol/L (9%), respectively). HDL cholesterol concentrations were not affected in subjects eating ispaghula cereal.

Another meta-analysis included eight studies involving a total of 384 patients
with hypercholesterolaemia who received ispaghula and 272 subjects who received cellulose placebo. Compared with placebo, consumption of 10.2 g ispaghula per day for ≥8 weeks lowered serum total cholesterol concentrations by 4% ($p < 0.0001$) and LDL cholesterol by 7% ($p < 0.0001$), but did not affect serum HDL cholesterol or triacyl glycerol concentrations. The ratio of apolipoprotein (apo) B to apo A-1 was lowered by 6% ($p < 0.05$), relative to placebo, in subjects consuming a low–fat diet. It was concluded that ispaghula is a useful adjunct to a low–fat diet in individuals with mild–to–moderate hypercholesterolaemia.

A randomised, placebo–controlled, multicentre study evaluated the long–term effectiveness of ispaghula husk as an adjunct to diet in treatment of primary hypercholesterolaemia. Men and women with hypercholesterolaemia followed the American Heart Association Step 1 diet for eight weeks prior to treatment. Individuals with LDL cholesterol concentrations between 3.36 and 4.91 mmol/L were randomly assigned to receive either ispaghula (Metamucil 5.1 g) or cellulose placebo twice daily for 26 weeks whilst continuing diet therapy. Overall, 163 participants completed the full protocol, 133 receiving ispaghula and 30 receiving cellulose placebo. Serum total and LDL cholesterol concentrations were 4.7% and 6.7% lower, respectively, in the ispaghula group than in the placebo group after 24–26 weeks ($p < 0.001$).

A randomised, double–blind, placebo–controlled crossover trial assessed the effects of ispaghula in lowering elevated LDL cholesterol concentrations in 20 children (aged 5–17 years). Children with LDL cholesterol concentrations of >2.84 mmol/L after three months on a low–fat, low–cholesterol diet received five weeks’ treatment with a ready–to–eat cereal containing water–soluble ispaghula husk (6 g/day) or placebo. The results indicated that there were no significant differences in total cholesterol, LDL cholesterol or HDL cholesterol concentrations between the two groups.

In a similar 12–week study, 50 children (aged 2–11 years) with LDL cholesterol concentrations ≥110 mg/dL received either cereal enriched with ispaghula (3.2 g soluble fibre per day) or plain cereal whilst maintaining a low–fat diet. Total cholesterol decreased by 21 mg/dL for the ispaghula group in comparison with 11.5 mg/dL for the control group ($p < 0.001$). LDL cholesterol also decreased by 23 mg/dL for the treated group in comparison with 8.5 mg/dL for the placebo group ($p < 0.01$).

The effect of adding water–soluble fibre to a diet low in total fats, saturated fat and cholesterol to treat hypercholesterolaemic children and adolescents has been reviewed. The review summarised that reductions in LDL cholesterol concentrations ranged from 0 to 23%. This wide range may be
related to dietary intervention and to clinical trial conditions. It was proposed that additional trials with larger numbers of well-defined subjects are needed.

**Hypoglycaemic effect**

Several studies have shown that ispaghula husk lowered blood glucose concentrations due to delayed intestinal absorption.\(^{G52}\) In one crossover study, 18 patients with non-insulin-dependent diabetes received ispaghula (Metamucil) or placebo twice (immediately before breakfast and dinner) during each 15-hour crossover phase.\(^{19}\) For meals eaten immediately after ispaghula ingestion, maximum postprandial glucose elevation was reduced by 14% at breakfast and 20% at dinner, relative to placebo. Postprandial serum insulin concentrations measured after breakfast were reduced by 12%, relative to placebo. Second-meal effects after lunch showed a 31% reduction in postprandial glucose elevation, relative to placebo. No significant differences in effects were noted between patients whose diabetes was controlled by diet alone and those whose diabetes was controlled by oral hypoglycaemic drugs. It was concluded that the results indicate that ispaghula as a meal supplement reduces proximate and second-meal postprandial glucose and insulin in non-insulin dependent diabetics.\(^{19}\)

**Other effects**

Ispaghula husk has been used to treat small numbers of patients with left-sided diverticular disease.\(^{4}\) Marked motility was observed for the right colon, but was not as pronounced for the left colon. The effects of ispaghula in this condition may be worth further investigation.

In an open, randomised, multicentre trial, 102 patients with ulcerative colitis (three months in remission, salicylate-treated, colitis over 20 cm) received ispaghula (10 g twice daily; \(n = 35\)), oral mesalazine (500 mg three times daily; \(n = 37\)) or ispaghula plus mesalazine (\(n = 30\)) for one year.\(^{20}\) Assessment, including endoscopy, was carried out at 3, 6, 9 and 12 months. The results suggested that ispaghula may be equivalent to mesalazine in maintaining remission in ulcerative colitis. However, this requires further investigation in a randomised, double-blind study.

In China, the seeds of related *Plantago* species have been used to treat hypertension.\(^{G41}\)
Side–effects, Toxicity

In common with all bulk laxatives, ispaghula may temporarily increase flatulence and abdominal distension, and may cause intestinal obstruction. If swallowed dry, ispaghula may cause oesophageal obstruction. In rare cases, allergic reactions may occur.\(^{G3\ G23}\)
Contra-indications, Warnings

In common with all bulk laxatives, ispaghula should not be given to patients with intestinal obstruction or conditions that may lead to intestinal obstruction, such as spastic bowel conditions. Ispaghula should always be taken with plenty of fluid to avoid oesophageal obstruction or faecal impaction. Bulk laxatives lower the transit time through the gastrointestinal tract and therefore may affect the absorption of other drugs. Absorption of currently administered drugs may be delayed. There may be a need to reduce insulin dosage in diabetics who are insulin dependent.

The EMEA HMPWG proposed core SPC for ispaghula includes the following information. Ispaghula husk is not to be used by patients with faecal impaction and undiagnosed abdominal symptoms, abdominal pain, nausea and vomiting (unless advised by a doctor), a sudden change in bowel habit that persists for more than two weeks, rectal bleeding, and failure to defecate following the use of a laxative. Ispaghula husk is also not to be used by patients suffering from abnormal constrictions in the gastrointestinal tract, diseases of the oesophagus and cardia, potential or existing intestinal blockage (ileus), or megacolon, diabetes mellitus which is difficult to regulate, or by patients with known hypersensitivity to ispaghula or any other constituents of the product. The husk should be taken with at least 150 mL of water or other fluid. Taking this product without adequate fluid may cause it to swell and block the throat or oesophagus and may cause choking. Intestinal obstructions may occur if an adequate fluid intake is not maintained. Ispaghula should not be taken by anyone who has had difficulty in swallowing or any throat problems. If chest pain, vomiting or difficulty in swallowing or breathing is experienced after taking the product, immediate medical attention should be sought. The treatment of the debilitated requires medical supervision. The treatment of elderly patients should be supervised. In the case of diarrhoea, sufficient intake of water and electrolytes is important.

Interaction with other medicinal products and other forms of interaction

Enteral absorption of concomitantly administered medicines such as minerals (e.g. calcium, iron, lithium, zinc), vitamins (B₁₂), cardiac glycosides and coumarin derivatives may be delayed. For this reason the product should not be taken 0.5–1 hour before, or after, intake of other drugs. If the product is taken together with meals in the case of insulin–dependent diabetics, it may be necessary to reduce the insulin dose.

Pregnancy and lactation
Ispaghula may be used during pregnancy and lactation.
Pharmaceutical Comment

The characteristic component of ispaghula is the mucilage which provides it with its bulk laxative action. Many of the herbal uses are therefore supported although no published information was located to justify the use of ispaghula in cystitis or infective skin conditions. Adverse effects and precautions generally associated with bulk laxatives apply to ispaghula. Clinical evidence exists for hypocholesterolaemia effects but it has been recommended that reduction in dietary fat intake is preferable to food supplements.\(^{(21)}\)
References


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Jamaica Dogwood
Species (Family)

*Piscidia erythrina* L. (Leguminosae)
Synonym(s)

Part(s) Used

Root bark
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL$^{(G37)}$
Constituents

See General References G6 G22 G40 G41 G64.

**Acids**
Piscidic acid (p-hydroxybenzyltartaric) and its mono and diethyl esters,\(^1\) fukiic acid and the 3’-O-methyl derivative; malic acid, succinic acid, and tartaric acid.

**Isoflavonoids**
Ichthynone, jamaicin, piscerythrone, piscidone and others. Milletone, isomillettone, dehydromillettone, rotenone and sumatrol (rotenoids), and lisetin.\(^2\)–\(^5\)

**Glycosides**
Piscidin, reported to be a mixture of two compounds, saponin glycoside (unidentified).\(^6\)

**Other constituents**
Alkaloid (unidentified, reported to be from the stem), resin, volatile oil 0.01%, β-sitosterol, tannin (unspecified).\(^6\)
Food Use

Jamaica dogwood is stated by the Council of Europe to be toxicologically unacceptable for use as a natural food flavouring.\textsuperscript{(G16)}
Herbal Use

Jamaica dogwood is stated to possess sedative and anodyne properties. Traditionally, it has been used for neuralgia, migraine, insomnia, dysmenorrhoea, and specifically for insomnia due to neuralgia or nervous tension. (G6 G7 G8 G64)
Dosage

Dried root bark
1–2 g or by decoction three times daily.\(^{(G6 \ G7)}\)

Liquid extract
1–2 mL (1 : 1 in 30\% alcohol) three times daily.\(^{(G6 \ G7)}\)

Liquid Extract of Piscidia
(BPC 1934) 2–8 mL.
Pharmacological Actions

In vitro and animal studies

Results of early studies reported Jamaica dogwood to possess weak cannabinoid and sedative activities in the mouse, guinea–pig and cat.\(^{(6–8)}\) In addition, *in vitro* antispasmodic activity on rabbit intestine, and guinea–pig and rat uterine muscle\(^{(6,9,10)}\) were noted and *in vivo* utero–activity in the cat and monkey were documented.\(^{(6,7,10,11)}\) In some instances, *in vitro* antispasmodic activity was found to be comparable to, or greater than, that observed for papaverine.

More recent work has supported these findings and reported that the antispasmodic activity of Jamaica dogwood on uterine smooth muscle is attributable to two isoflavone constituents, one being equipotent to papaverine.\(^{(11)}\)

Jamaica dogwood extracts have also been documented to exhibit antitussive, antipyretic, and anti–inflammatory activities in various experimental animals.\(^{(7)}\)

Rotenone is an insecticide that has been used in agriculture for the control of lice, fleas, and as a larvicide.\(^{(G45)}\) Jamaica dogwood has been used extensively throughout Central and South America as a fish poison;\(^{(6)}\) the wood contains two piscicidal principles, rotenone and ichthynone. Rotenone is relatively harmless to warm–blooded animals.\(^{(12)}\)

Rotenone has reported exhibited anticancer activity towards lymphocytic leukaemia and human epidermoid carcinoma of the nasopharynx.\(^{(G22)}\) It is also documented to be carcinogenic.\(^{(G22)}\)
Side-effects, Toxicity

Symptoms of overdose are stated to include numbness, tremors, salivation and sweating.\(^{G22}\) Jamaica dogwood has been found to be toxic when administered parenterally to rats and rabbits, but non-toxic when given orally, with doses exceeding 90 g dried extract/kg tolerated.\(^{6}\) An LD\(_{50}\) (mice, intravenous injection) of an unidentified saponin constituent has been reported as 75 μg/kg body weight.\(^{9}\) Oral doses of up to 1.5 mg/kg were stated to have no effect.\(^{6}\)

Jamaica dogwood is stated to be irritant and toxic to humans.\(^{G51}\)
Contra-indications, Warnings

It is recommended that Jamaica dogwood should be used with great care, and only by trained practitioners. Jamaica dogwood may potentiate sedative effects of existing therapy.

**Pregnancy and lactation**

Jamaica dogwood has been reported to exhibit a potent depressant action on the uterus both *in vitro* and *in vivo*. In view of this and the general warnings regarding the use of Jamaica dogwood, it should not be used during pregnancy and lactation.
Pharmaceutical Comment

Jamaica dogwood is characterised by various isoflavone constituents, to which the antispasmodic properties described for the wood have been attributed. In addition, sedative and narcotic activities have been documented that justify the reputed herbal uses. Although Jamaica dogwood is reported to be of low toxicity in various animal species, it is also documented as toxic to humans\(^{(G51)}\) and is recommended to be used with great care.\(^{(G49)}\) In view of this, excessive use of Jamaica dogwood should be avoided.
References

See also General References G6 G9 G16 G22 G31 G36 G37 G40 G41 G45 G49 G51 G64.


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Species (Family)

*Orthosiphon stamineus* Benth. (Lamiaceae)
Synonym(s)

Kumis Kucing (Indonesian, Malay), Orthosiphon aristatus Miq., Orthosiphon spicatus (Thundb.) Bak.
Part(s) Used

Fragmented dried leaves, tops of stems
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)

BP 2002\(^{(G71)}\)

Complete German Commission E\(^{(G3)}\)

ESCOP 1997\(^{(G52)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)

Ph Eur 2004\(^{(G72)}\)
Legal category (Licensed Products)

Java tea is not included in the GSL.
Constituents

See General References G2 G52.

**Benzochromenes**
Orthochromene A,\(^{(1)}\) methylripariochromene A\(^{(2)}\) and acetovanillogenone.\(^{(1)}\)

**Diterpenes**
Isopimarane–type diterpenes (orthosiphonones A and B,\(^{(1)}\) orthosiphols A and B,\(^{(3)}\) orthosiphols F, G, H and I\(^{(4)}\)), pimarane–type diterpenes (neoorthosiphols A and B)\(^{(5)}\) and staminol A.\(^{(4)}\)

**Essential oil**
0.02–0.7%. Various compounds including β-elemene, β-caryophyllene, α-humulene, β-caryophyllene oxide, can–2–one and palmitic acid.\(^{(6)}\)

**Flavonoids**
Sinensetin, tetramethylscutellarein and other tetramethoxyflavones, eupatorin, salvigenin, cirsimaritin, pilloin, rhamnazin, trimethylapigenin and tetramethyluteolin.\(^{(7–11)}\) These lipophilic flavonoids are present in concentrations of approximately 0.2–0.3%;\(^{(10)}\) flavonoid glycosides are also present.

**Other constituents**
Caffeic acid and derivatives (e.g. rosmarinic acid), inositol, phytosterols (e.g. β-sitosterol)\(^{(11,12)}\) and potassium salts.
Food Use

Java tea is not used in foods.
Java tea has traditionally been used in Java for the treatment of hypertension and diabetes.\(^{(1,5,13)}\) It has also been used in folk medicine for bladder and kidney disorders, gallstones, gout and rheumatism. Java tea is stated to have diuretic properties.\(^{(14)}\)
Dosage

Dried material
2–3 g in 150 mL water two to three times daily as an infusion.\(^{(G52)}\)
Pharmacological Actions

In vitro and animal studies

Diuretic effects
Several studies in rats have reported diuretic activity of extracts of *O. stamineus* and *O. aristatus*\(^{(14–16)}\) and of flavonoids (sinensetin and a tetramethoxyflavone) isolated from *O. aristatus*.\(^{(17)}\) Intraperitoneal administration of a hydroalcoholic extract of *O. stamineus* to rats caused a significant diuresis over the following 2–24 hours compared with controls.\(^{(14)}\) The effect was similar to that observed following intraperitoneal administration of hydrochlorothiazide (10 mg/kg).\(^{(14)}\) Oral administration of an aqueous extract of *O. aristatus* increased ion excretion to a similar extent as did furosemide, although no diuretic action was noted.\(^{(16)}\)

Oral administration of methylripariochromene A (100 mg/kg) has been shown to increase urinary volume in fasted rats for three hours after oral administration; the increase in urine volume was similar to that observed with oral administration of hydrochlorothiazide (25 mg/kg).\(^{(13)}\) Sodium, potassium and chloride ion excretion was increased with methylripariochromene A (100 mg/kg), although urinary sodium ion excretion did not increase. A mechanism for the diuretic action of methylripariochromene A has not yet been elucidated, although it appears to have a different mode of action to that of hydrochlorothiazide.\(^{(13)}\)

Hypoglycaemic effects
In normoglycaemic rats, oral administration of an aqueous extract of *O. stamineus* (0.5 g/kg) had no significant effect on fasting blood glucose concentrations over a 7-hour period, although administration of 1 g/kg produced a significant decrease in blood glucose concentration compared with that in a control group.\(^{(18)}\) A hypoglycaemic effect was also observed following administration of *O. stamineus* extract (1 g/kg) to rats loaded with glucose (1.5 g/kg) and in streptozotocin–induced diabetic rats; the effect of *O. stamineus* extract in streptozotocin–induced diabetic rats was similar to that observed with glibenclamide (10 mg/kg).\(^{(18)}\)

Antihypertensive effects
Methylripariochromene A has been reported to have several pharmacological actions related to antihypertensive activity.

In stroke–prone, spontaneously hypertensive rats, subcutaneous administration of methylripariochromene A (100 mg/kg) produced a continuous reduction in systolic blood pressure and a decrease in heart rate.
Methylripariochromene A also suppressed agonist–induced contractions in the rat thoracic aorta and decreased the contractile force in isolated guinea–pig atria without significantly affecting the beating (heart) rate. The mechanism of action for these antihypertensive effects of methylripariochromene A is, however, unclear.\(^{(13)}\)

Migrated pimarane–type diterpenes (neoorthosiphols A and B), isopimarane–type diterpenes (orthosiphols A and B, orthosiphoonones A and B), benzochromenes (methylripariochromene, aceto vanillochromene, orthochromene A) and flavones (tetramethylscutellarein, sinensetin) isolated from \textit{O. aristatus} have been reported to exhibit a suppressive effect on contractile responses in the rat thoracic aorta.\(^{(19)}\)

\textbf{Cytostatic effects}\n
Sinensetin and tetramethylscutellarein have been reported to demonstrate \textit{in vitro} cytostatic activity towards Ehrlich ascites tumour cells.\(^{(10)}\) Growth inhibition appears to be dose dependent, with 50\% inhibition occurring at concentrations of approximately 30 and 15 μg/ml for sinensetin and tetramethylscutellarein, respectively. Orthosiphols A and B have been reported to inhibit inflammation induced by the tumour promoter 12-O-tetradecanoylphorbol–13–acetate (TPA) on mouse ears.\(^{(3)}\)

Fractions of \textit{O. stamineus} leaves have been reported to have activity against a melanoma cell line \textit{in vitro}.\(^{(20)}\)

\textbf{Antimicrobial effects}\n
An aqueous extract of \textit{O. aristatus} has demonstrated antibacterial activity against two serotypes of \textit{Streptococcus mutans} (MIC 7.8–23.4 mg/mL).\(^{(21)}\) Other \textit{in vitro} studies have reported a lack of antibacterial activity for flavonoids (sinensetin, tetramethylscutellarein and a tetramethoxyflavone in concentrations of 10 and 100 μg/mL) isolated from \textit{O. aristatus} leaves against \textit{Escherichia coli}, \textit{Proteus mirabilis}, \textit{Pseudomonas aeruginosa}, \textit{Staphylococcus aureus} and \textit{Enterococcus}.\(^{(17)}\)

\textit{O. stamineus} extract has also been shown to inhibit spore germination in six out of nine fungal species tested: \textit{Saccharomyces pastorianus}, \textit{Candida albicans}, \textit{Rhizopus nigricans}, \textit{Penicillium digitatum}, \textit{Fusarium oxysporum} and \textit{Trichophyton mentagrophytes}.\(^{(22)}\)

\textbf{Other effects}\n
\textit{In vitro}, \textit{O. spicatus} has been shown to inhibit 15–lipoxygenase, an enzyme thought to be involved in the development of atherosclerosis.\(^{(11)}\) Furthermore, the flavonoids sinensetin and tetramethylscutellarein
demonstrate dose–dependent inhibition with IC\textsubscript{50} values of 114 ± 5 and 110 ± 3 μmol/L, respectively, although other flavonoids from \textit{O. spicatus} appear to be less efficient inhibitors of 15–lipoxygenase. The inhibitory activity of the whole extract was greater than could be expected from the activities of each of its flavonoid constituents, and it has been suggested that synergism may be occurring.\textsuperscript{(11)} More recent \textit{in vitro} studies have shown that flavonoids from \textit{O. spicatus} prevent oxidative inactivation of 15–lipoxygenase, with trimethylapigenin, eupatorin and tetramethylluteolin showing the strongest enzyme–stabilising effects.\textsuperscript{(23)} However, there was no correlation between enzyme stabilisation and enzyme inhibition.\textsuperscript{(23)}

**Clinical studies**

Early studies reported increases in diuresis in subjects following the oral administration of extracts of \textit{Orthosiphon}.\textsuperscript{(G52)} A randomised, double–blind, placebo–controlled, crossover study reported no effect on 12- and 24–hour urine output or on sodium excretion in 40 healthy volunteers who received 600 mL of an infusion of \textit{Orthosiphon} leaves daily (equivalent to 10 g dried leaves) for four days.\textsuperscript{(24)} A study involving six healthy volunteers who drank \textit{Orthosiphon} tea (250 mL) every 6 hours for one day reported an increase in urine acidity 6 hours after ingestion.\textsuperscript{(25)}

A study involving 67 patients with uratic diathesis who received Java tea for three months reported that no effects were observed on diuresis, glomerular filtration, osmotic concentration, urinary pH, plasma content and excretion of calcium, inorganic phosphorus and uric acid.\textsuperscript{(26)}
Side-effects, Toxicity

None documented.
Contra–indications, Warnings

None known. In view of the lack of clinical data on the use of Java tea, excessive or long–term use should be avoided. Adequate fluid intake (2 L or more per day) should be ensured whilst using Java tea.\(^\text{G35}\)

**Pregnancy and lactation**

There are no data available on the use of Java tea in pregnancy and lactation. In view of the lack of toxicity data, use of Java tea during pregnancy and lactation should be avoided.
The reported pharmacological activities of Java tea are mainly associated with the lipophilic flavonoids, benzochromene and, to a lesser extent, diterpene constituents.

Documented scientific evidence from *in vitro* and animal studies provides some supportive evidence for some of the traditional uses of Java tea. However, there is a lack of clinical data and well-designed, controlled clinical trials involving adequate numbers of patients are required. Furthermore, studies investigating the active principles responsible for specific pharmacological activities and their mechanisms of action are necessary.

There have been reports of adulteration/botanical substitution occurring with *Orthosiphon*.(27,28,G2)

In view of the lack of toxicity and safety data, excessive use of Java tea should be avoided.
References

See also General References G2 G3 G9 G15 G28 G36 G43 G52.

16. Englert J, Harnischfeger G. Diuretic action of aqueous Orthosiphon extract


Species (Family)

*Juniperus communis* L. (Pinaceae)
Synonym(s)
Baccae Juniperi, Genièvre, Wacholderbeeren, Zimbro
Part(s) Used

Fruit (berry)
Pharmacopoeial and Other Monographs

BHP 1996$^{(G9)}$

BP 2002$^{(G71)}$

Complete German Commission E$^{(G3)}$

ESCOP 1997$^{(G52)}$

Martindale 33rd edition$^{(G67)}$

PDR for Herbal Medicines 2nd edition$^{(G36)}$

Ph Eur 2004$^{(G72)}$
Legal Category (Licensed Products)

GSL\(^{G37}\)
Constituents
See General References G2 G22 G41 G53 G58 G62 G64.

**Acids**
Diterpene acids, ascorbic acid and glucuronic acid.

**Flavonoids**
Amentoflavone,\(^1\) quercetin, isoquercitrin, apigenin and various glycosides.

**Tannins**
Proanthocyanidins (condensed), galallocatechin and epigallocatechin.\(^2\)

**Volatile oils**
0.2–3.42%. Primarily monoterpenes (about 58%) including α-pinene, myrcene and sabinene (major), and camphene, camphor, 1,4-cineole, p-cymene, α- and γ-cadinene, limonene, β-pinene, γ-terpinene, terpinen-4-ol, terpinyl acetate, α-thujene, borneol; sesquiterpenes including caryophyllene, epoxydihydrocaryophyllene and β-elemem-7α-ol.\(^3,4\)

**Other constituents**
Geijerone (C\(_{12}\) terpenoid), junionone (monocyclic cyclobutane monoterpenoid),\(^5\) desoxypodophyllotoxin (lignan),\(^6\) resins and sugars.
Food Use

Juniper berries are widely used as a flavouring component in gin. Juniper is listed by the Council of Europe as a natural source of food flavouring (fruit N2, leaf and wood N3). Category N2 indicates that the berries can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product. Category N3 indicates that there is insufficient information available for an adequate assessment of potential toxicity to be made.\(^{G16}\) In the USA, extracts and oils of juniper are permitted for food use.\(^{G65}\)
The German Commission E approved use for dyspepsia. Juniper is stated to possess diuretic, antiseptic, carminative, stomachic and antirheumatic properties. Traditionally, it has been used for cystitis, flatulence, colic, and applied topically for rheumatic pains in joints or muscles.
Dosage

**Dried ripe fruits**
100 mL as an infusion (1 : 20 in boiling water) three times daily. *(G7)*

**Fruit**
1–2 g or equivalent three times daily; 2–10 g (equivalent to 20–100 mg of volatile oil). *(G3)*

**Liquid extract**
2–4 mL (1 : 1 in 25% alcohol) three times daily. *(G7)*

**Tincture**
1–2 mL (1 : 5 in 45% alcohol) three times daily. *(G7)*

**Oil**
0.03–0.2 mL (1 : 5 in 45% alcohol) three times daily.
Pharmacological Actions

Pharmacological actions that have been documented for juniper are primarily associated with the volatile oil components.

**In vitro and animal studies**

The volatile oil is documented to possess diuretic, gastrointestinal antiseptic and irritant properties.\(^{(G41)}\)

The diuretic activity of juniper has been attributed to the volatile oil component, terpinen–4–ol, which is reported to increase the glomerular filtration rate.\(^{(G60)}\) Terpinen–4–ol is also stated to be irritant to the kidneys although in a later review by the same author there is no such statement and the oil is stated to represent no hazards.\(^{(G58)}\)

An antifertility effect has been described for a juniper extract, administered to rats (300/500 mg, by mouth) on days 1–7 of pregnancy.\(^{(7)}\) An abortifacient effect was also noted at both dose levels when the extract was administered on days 14–16.\(^{(7)}\) No evidence of teratogenicity was reported. Anti–implantation activity has been reported as 60 to 70%\(^{(8)}\) and as dose dependent.\(^{(7)}\) Juniper is reported to have both a significant\(^{(9)}\) and no\(^{(8)}\) antifertility effect. A uterine stimulant activity has been documented for the volatile oil.\(^{(G30)}\)

A potent and non–toxic inhibition of the cytopathogenic effects of herpes simplex virus type 1 in primary human amnion cell culture has been described for a juniper extract.\(^{(6,10)}\) The active component isolated from the active fraction was identified as a lignan, desoxypodophyllotoxin.\(^{(6)}\) Antiviral activities documented for the volatile oil have also been partly attributed to the flavonoid amentoflavone.\(^{(1)}\)

Anti–inflammatory activity of 60% compared to 45% for the indometacin control has been reported for juniper berry extract.\(^{(11)}\) Both test and control were administered orally to rats (100 mg/kg and 5 mg/kg respectively) one hour before eliciting foot oedema.

A transient hypertensive effect followed by a more prolonged hypotensive effect has been reported for a juniper extract in rats (25 mg/kg, intravenous injection).\(^{(12)}\)

A fungicidal effect against *Penicillium notatum* has been documented.\(^{(13)}\)

Astringent activity is generally associated with tannins, which have been
documented as components of juniper. An aqueous decoction of the berries has a hypoglycaemic effect in rats.\textsuperscript{(14)} In rats, oral administration of an aqueous infusion (5 mL) increased chloride ion secretion by 119\% and by 45\% in similar experiments with rabbits.\textsuperscript{(G52)}
Side-effects, Toxicity

The volatile oil is reported to be generally non-sensitising and non-phototoxic, although slightly irritant when applied externally to human and animal skin.\(^{(G41,G58)}\) Excessive doses of terpinen-4-ol, the diuretic principle in the volatile oil, may cause kidney irritation.\(^{(G22)}\)

Dermatitic reactions have been recognised with juniper and positive patch test reactions have been documented.\(^{(15,G51)}\) The latter are attributed to the irritant nature of the juniper extract.\(^{(15)}\)

Symptoms of poisoning following external application of the essential oil are described as burning, erythema, inflammation with blisters and oedema.\(^{(G22)}\) Internally, symptoms from overdose are documented as pain in or near the kidneys, strong diuresis, albuminuria, haematuria, purplish urine, tachycardia, hypertension, and rarely convulsions, metrorrhagia and abortion.\(^{(G22)}\)

The acute toxicity of juniper has been investigated in rats who were administered extracts for seven days.\(^{(11)}\) An oral dose of 2.5 g/kg was tolerated with no mortalities or side-effects noted. A dose of 3 g/kg induced hypothermia and mild diarrhoea in 10–30% of animals.\(^{(11)}\) An \(LD_{50}\) value (mice, intraperitoneal injection) has been stated as 3 g/kg.\(^{(4)}\)
Contra–indications, Warnings

Juniper is contra–indicated in individuals with existing renal disease.\(\text{(G7 G42 G49 G52)}\) The internal use of the oil should be restricted to professionals.\(\text{(G42)}\) External application of the oil may cause an irritant reaction. However, this has been refuted and the oil is stated to have no hazards and is not contra–indicated.\(\text{(G58)}\) Juniper has been confused with savin (Juniperus sabina) in the literature and this may be the reason for believing that the oil is toxic.\(\text{(G58)}\) Juniper may potentiate existing hypoglycaemic and diuretic therapies; prolonged use may result in hypokalaemia.

**Pregnancy and lactation**

Juniper is contra–indicated in pregnancy.\(\text{(G7 G22 G49)}\) It is reputed to be an abortifacient and to affect the menstrual cycle.\(\text{(G30)}\)

A juniper fruit extract has exhibited abortifacient, antifertility and anti–implantation activities (see *In vitro* and animal studies).
Pharmaceutical Comment

Many of the traditional uses documented for juniper can be supported by documented pharmacological actions or known constituents. There is evidence that the berries are abortifacient and since this is believed not to be due to the oil there must be other toxic constituents present. It is recommended that use should not exceed levels specified in food legislation.
References


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Kava
Species (Family)

*Piper methysticum* Forst. f. (Piperaceae)

Fourteen different varieties are used throughout Oceania (Polynesia, Melanesia, Micronesia).\(^{(1)}\)

**Related species**

Cultivars of *P. methysticum* have been developed in some Pacific Islands from *Piper wichmanni* C.DC (syn: *Piper erectum* C.DC, *Piper schlecteri* C.DC, *Piper arbuscula* Trelease). *Piper sanctum* (Miqu.) Schlect. is native to Mexico.\(^{(2,3)}\)
Synonym(s)
Intoxicating Pepper, Kava-kava, Kawa, Kawa-kawa
Part(s) Used
Peeled dry rhizome (sometimes referred to incorrectly as the root)
Pharmacopoeial and Other Monographs

BHMA 2003\(^{(G66)}\)

BHP 1996\(^{(G9)}\)

Complete German Commission E 1998\(^{(G3)}\)

Expanded German Commission E 2000\(^{(G4)}\)

Martindale 33rd edition 2002\(^{(G67)}\)

Mills and Bone 2000\(^{(G50)}\)

PDR for Herbal Medicine 2nd edition 2000\(^{(G36)}\)

WHO volume 2 2002\(^{(G70)}\)
Legal Category (Licensed Products)

Prohibited in unlicensed medicines in the UK.\textsuperscript{(4)}
Constituents

Kavalactones
Kavalactones, kavapyrones, 2-pyrones, δ-lactones with styryl or
dihydrostyryl substituents.\(^{(1–3,5–9, G56)}\) Dried rhizomes should contain at least
3.5% kavalactones\(^{(G56)}\) and good-quality material 5.5–8.3%.\(^{(10)}\) Ethanol–
water extracts contain 30% kavalactones, whereas acetone–water extracts
contain 70%.\(^{(G56)}\) The kavalactones occur as a complex mixture of at least 18
compounds,\(^{(5)}\) which are of three main types: styryl enolide pyrones (e.g.
kawain (= kavain), dimethoxykawain, methysticin), styryl dienolide pyrones
(e.g. yangonin, desmethoxyyangonin), and dihydrostyryl enolide pyrones
(e.g. dihydrokawain, dimethoxydihydrokawain, dihydromethysticin). The four
major kavalactones of the rhizome are kawain (1–2%), dihydrokawain (0.6–
1%), methysticin (1.2–2%) and dihydromethysticin (0.5–0.8%).\(^{(G56)}\) Smaller
quantities (<0.1%) of dimeric kavalactones (e.g. trux-yangonins I, II, III)
have also been isolated.\(^{(2,3)}\)

Alkaloids/amides
Cepharadione A (aporphine-type) is a minor component (4 kg yielded
1 mg).\(^{(11)}\) Small quantities of \(N\)-cinnamoylpyrrolidine and its \(o\)-methoxy
analogue are also present.\(^{(6,12,13)}\)

Chalcones
Flavokawains A, B and C.\(^{(6,9,14)}\)

Flavonoids
Pinostrobin, 5,7-dimethoxyflavanone.\(^{(14)}\)

Steroids
Sitosterol, stigmasterol, stigmastanol.\(^{(6,13)}\)

Esters
Bornyl cinnamate\(^{(13)}\) and bornyl 3,4-methylenedioxyxycinnamate.\(^{(14)}\)

Aliphatic alcohols
Docosan-1-ol, dodecan-1-ol, eicosan-1-ol, hexacosan-1-ol, hexadecan-1-ol,
octadecan-1-ol, \(n\)-tetradecanol, transphytol.\(^{(6)}\)

Other constituents
Cinnamylideneacetone,\(^{(5)}\) long-chain fatty acids.\(^{(6)}\)

Other parts of the plant\(^{(15)}\)
Stem peelings may be included as raw material in kava commerce due to the high demand for the rhizome; leaves and branches are used in folk medicine. Pipermethystine (a piperidone amide) is present in stem peelings (traces to 0.85%). 3α,4α-Epoxy-5β-pipermethysticin (0.93%) was isolated from stem peelings of one cultivar, but was absent from 10 other cultivars, and the related alkaloid awaine was present in the unopened leaves of 11 cultivars (0.16–2.67%). 7,8-Dihydrokawain, 7,8-dihydromethysticin and 5,6,7,8-tetrahydroyangonin are present in stem peelings.

Other species

Kavalactones occur in *P. wichmanni* and *P. sanctum*. The latter species contains several cinnamoyl butenolides (piperolides), e.g. methylenedioxypiperolide and 7,8-epoxypiperolide.\(^2,3\)
Food Use

Kava is used as an intoxicant drink, on either informal or ceremonial occasions, by Pacific Islanders e.g. from Fiji, Samoa and Tonga.\(^{(1,7)}\) Some claim that it has a pleasant, cooling, aromatic taste with numbing on the tongue and is stimulating, while others refer to great bitterness with a burning sensation in the mouth. It is reputed to reduce fatigue, allay anxieties and produce a cheerful and sociable attitude. Unpleasant effects reported include dizziness, sleeping disorders, stomach pains, lethargy and skin reactions. These reported effects are taken from a wide geographical area and any differences may be due to a number of reasons including plant varieties or growing conditions.
Herbal Use

In many parts of the Pacific it is believed that kava is beneficial to health by soothing nervous conditions, inducing relaxation and sleep, counteracting fatigue, and reducing weight. Medicinal uses also include treatment of urinary tract infections, asthma, rheumatism, headache, fever, gonorrhoea and syphilis, and use as a diuretic and stomachic.\(^{(1,7,G34)}\) The medicinal use of kava is now widespread, e.g. across Europe, North America and Australia, where it is used to treat anxiety, nervous tension, restlessness, mild depression and menopausal symptoms.\(^{(G9, G32, G50, G56, G60, G67)}\) It has also been adopted by the Aboriginal community in parts of Australia as an intoxicating drink.\(^{(16)}\)

The German Commission E recommended kava for the treatment of nervous anxiety, stress and restlessness.\(^{(G3)}\) Traditional uses listed for kava rhizome in other standard herbal and pharmaceutical reference texts include cystitis, urethritis, infection or inflammation of the genitourinary tract, rheumatism and, topically, for joint pains.\(^{(G66)}\)
Dosage

For treatment of anxiety (adults) by oral administration.

**Dried rhizome**
1.5–3 g per day. \(^{(G50)}\) Equivalent to 60–120 mg kavalactones per day. \(^{(G3)}\)

**Liquid extract**
3–6 mL per day (1 : 2 liquid extract, unspecified solvent). \(^{(G50)}\)

**Standardised preparations**
100–200 mg kavalactones per day. \(^{(G50)}\) 60 mg kavalactones 2–4 times per day in tablet form. \(^{(G50)}\) 60–120 mg kavalactones per day. \(^{(G3)}\)

Kava exists in numerous varieties of differing potency \(^{(7)}\) and only preparations with standardised kavalactone content should be used for medicinal purposes. Medicinal extracts prepared with ethanol–water yield dry extracts with about 30% kavalactones content, whereas acetone–water prepared dry extracts contain about 70% kavalactones. \(^{(G56)}\)

Dosages (adults) used in clinical trials have varied widely, but typically are those equivalent to 60–240 mg kavalactones daily by oral administration in divided doses (see Clinical studies). Duration of use of kava extracts generally should not exceed three months. \(^{(G3, G4, G56)}\)
Pharmacological Actions

Kava has been investigated mostly for its anxiolytic effects, although other central nervous system activities, such as anticonvulsant and analgesic properties, and other effects have been documented following preclinical studies. The kavalactones are believed to be the major active constituents of kava.

In vitro and animal studies

Pharmacokinetics

Uptake of the kavalactones kavain, dihydrokavain, yangonin and desmethoxyyangonin into brain tissue has been documented following intraperitoneal administration of each of these compounds at a dose of 100 mg/kg to mice.\(^\text{17}\) Maximum concentrations of kavain and dihydrokavain were noted 5 minutes after administration, and these compounds were rapidly eliminated. In contrast, yangonin and desmethoxyyangonin were eliminated more slowly. All four compounds were also detected in mouse brain tissue following intraperitoneal administration of kava resin 120 mg/kg (containing kavain 36.7\%, dihydrokavain 19.2\%, yangonin 15\% and desmethoxyyangonin 13.3\%), although the concentrations of kavain and yangonin were higher than was noted following individual administration of these constituents.

Central nervous system activities

Anxiolytic properties for kava extract and isolated kavalactones have been documented in an experimental model of anxiety, the chick social separation-stress procedure. In a series of experiments, kava extract (containing 30% kavalactones; 30 mg/mL per kg body weight), dihydrokavain (30 mg/mL per kg body weight) and chlordiazepoxide (5 mg/mL per kg body weight) administered intraperitoneally 30 minutes before testing significantly reduced the separation-stress effect \( (p < 0.05 \text{ for each substance}) \).\(^\text{18}\) However, the isolated kavalactones kavain, methysticin, dihydromethysticin, yangonin and desmethoxyyangonin administered according to the same regimen did not have a statistically significant effect. Further work using the same experimental model confirmed these findings and found that total kavalactone content was not predictive of outcome, but that a dihydrokavain content of at least 15\% was necessary for anxiolytic activity.\(^\text{19}\) In this study, the kava samples and fractions that demonstrated anxiolytic activity were reported to be without sedative effects.

In contrast, previous studies have reported sedative effects for kava extract. In mice, kava extract (containing 7\% kavalactones) at doses of at least...
50 mg/kg body weight (by intraperitoneal injection) reduced spontaneous motility to a greater extent than did control.\(^{(20)}\) The effect was enhanced by the addition of (±)-kavain (ratio of kava extract to (±)-kavain, 1 : 0.12), although this compound had no sedative effect when administered alone. In another experimental model, kava extract 100 mg/kg body weight and (±)-kavain 12 mg/kg body weight, each administered alone, had no sedative effect, whereas a combination of the two substances significantly reduced amphetamine (5 mg/kg body weight subcutaneously)-induced hypermotility. Sedative effects have also been documented for an ethanolic extract of kava rhizome (containing 50% kavalactones) 100 mg/kg body weight administered by gastric tube and 200 mg/kg body weight intraperitoneally in the amphetamine-induced hypermotility test and barbiturate-induced sleeping time, respectively.\(^{(21)}\)

In studies utilising the conditioned avoidance response test in rats, an experimental model used to test for antipsychotic activity, aqueous (kavalactone-free) kava extract 30–500 mg/kg body weight intraperitoneally had no statistically significant effect.\(^{(22)}\) However, administration of kava resin at doses of 125 mg/kg intraperitoneally significantly inhibited the conditioned avoidance response, although to a lesser extent than did chlorpromazine and haloperidol.

In cats, a kava extract in arachis oil (50–100 mg kavalactones per kg body weight intraperitoneally) and the individual kavalactone (±)-kavain (10–50 mg per kg body weight intraperitoneally) were reported to be active in the amygdala complex region of the brain.\(^{(23)}\)

Receptor binding studies with kava extracts and individual kavalactones have reported conflicting results. One series of experiments found that kava resin and individual kavalactones displayed only weak activity on GABA\(_A\)- and no activity on GABA\(_B\)-binding sites in rat brain membranes \textit{in vitro}, and that there was no significant effect on benzodiazepine receptors following intraperitoneal administration of kava resin 150 mg/kg body weight to mice.\(^{(24)}\) A marked effect of kavain on GABA has also been stated to be unlikely.\(^{(25)}\) By contrast, a kavalactone-enriched ethanol/aqueous extract of kava rhizome (containing 58% kavalactones and 42% other lipid-soluble compounds) increased the density of GABA-binding sites in certain brain regions.\(^{(26)}\) Other experiments have shown concentration- and structure-dependent effects of kavalactones on binding of bicuculline methochloride (BMC) to GABA\(_A\) receptors from rat cortex preparations.\(^{(27)}\) (+)-Kavain, (+)-methysticin and (+)-dihydromethysticin enhanced BMC binding by 18–28% at a concentration of 0.1 \(\mu\text{mol/L}\), whereas (+)-dihydrokavain did so only at a concentration of 10 \(\mu\text{mol/L}\), and yangonin at a concentration of 1 \(\mu\text{mol/L}\);
desmethoxyyangonin had no effect. Further radioreceptor assays demonstrated that these six kavalactones had no effect on the binding of flunitrazepam to benzodiazepine receptors in rat cortex preparations, indicating that the influence of kavalactones on GABA<sub>A</sub> receptors was not based upon an interaction with benzodiazepine receptors.<sup>(27)</sup>

Other <i>in vitro</i> studies have investigated the effects of kava extracts and individual kavalactones on other transmitters in the central nervous system (CNS). A kavalactone-rich kava rhizome extract (containing 68% kavalactones) was a reversible inhibitor of monoamine-oxidase B (MAO-B) in intact and disrupted platelets (inhibitory concentration IC<sub>50</sub> 24 μmol/L and 1.2 μmol/L, respectively), although there were differences in MAO-B inhibition among the different synthetic kavalactones with desmethoxyyangonin and (±)-methysticin being the most potent inhibitors.<sup>(28)</sup> Differences between kavalactones in inhibition of noradrenaline (norepinephrine) uptake in synaptosomes prepared from rat cerebral cortex and hippocampus have also been documented: (±)- and (+)-kavain gave approximately equal values and both were more potent inhibitors than (+)-methysticin, although none of the compounds inhibited serotonin uptake.<sup>(29)</sup>

It has been suggested, following <i>in vitro</i> studies involving ipsapirone (a serotonin-1<sub>A</sub> receptor agonist)-induced field potential changes in guinea-pig hippocampal slices, that kavain and dihydromethysticin may modulate serotonin-1<sub>A</sub> receptor activity, although further work is needed to identify the precise mechanism for this.<sup>(30)</sup>

<i>In vivo</i> studies in rats administered a single oral dose of (+)-dihydromethysticin 100 mg/kg body weight or fed (±)-kavain in the diet over a 78-day period showed that neither kavalactone regimen affected brain tissue concentrations of dopamine and serotonin, although since extracellular neurotransmitter concentrations were not measured in this study, receptor-mediated effects of kavalactones on dopaminergic and serotonergic neurons could not be excluded.<sup>(31)</sup> <i>In vivo</i> (rats), kava extract 20 and 120 mg/kg body weight intraperitoneally increased dopamine concentrations in the nucleus accumbens, although a dose of 220 mg/kg body weight led to an initial decrease followed by an increase above baseline values.<sup>(32)</sup> It was suggested that this ceiling effect may be due to yangonin which may have dopamine antagonist activity.

The development of physiological tolerance to an aqueous extract of kava administered intraperitoneally to mice has been documented, although there was no clear evidence of development of physiological or learned tolerance to kava resin.<sup>(33)</sup>
Anticonvulsant and neuroprotective activities

Studies described in the older literature have documented anticonvulsant effects for kavalactones in several experimental models.\(^{34,50}\) The anticonvulsant properties of (+)-methysticin \textit{in vitro} may arise from a direct membrane action on the excitability of neurons,\(^{34}\) and \textit{in vitro} assays have shown that (+)-methysticin,\(^{35}\) (+)-kavain\(^{36}\) and the synthetic kavalactone (±)-kavain\(^{34-36}\) appear to interact with voltage-dependent sodium channels, and that (±)-kavain also interacts with voltage-dependent calcium channels.\(^{37}\) Inhibition by (±)-kavain of veratridine-activated voltage-dependent sodium ion channels in synaptosomes from rat cerebral cortex,\(^{38}\) and veratridine-induced increase in intracellular calcium ion concentrations has been described following \textit{in vitro} studies utilising rat cerebrocortical synaptosomes.\(^{39}\) Reduction in veratridine-induced glutamate release following (±)-kavain administration has been reported both \textit{in vitro}\(^{39}\) and \textit{in vivo} in freely moving rats.\(^{30}\) Substances which reduce extracellular glutamate concentrations are of interest for their potential as anticonvulsant agents.

Some of the mechanisms described above documented for certain kavalactones may also be important in neuroprotective effects reported for the synthetic kavalactone (±)-kavain. For example, the role of sodium-ion channel blockade in the neuroprotective effect of (±)-kavain against anoxia \textit{in vitro} has been described,\(^{41}\) and (±)-kavain (50, 100 or 200 mg/kg intraperitoneally) has been shown to protect nigrostriatal dopaminergic neurons against 1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine (MPTP)-induced toxicity in mice, an experimental model of Parkinson’s disease.\(^{42}\) A neuroprotective effect against ischaemic brain damage in mice and rats has been demonstrated for kava extract (WS-1490 containing 70% kavalactones) and the individual kavalactones methysticin and dihydromethysticin, but not for kavain, dihydrokavain and yangonin. Kava extract 150 mg/kg given orally as an emulsion (polyethyleneglycol 400 and water; 20 : 80) 1 hour before experimentally induced ischaemia, and methysticin and dihydromethysticin (both 10 and 30 mg/kg intraperitoneally 15 minutes before induction of ischaemia), compared with control, significantly reduced the size of the infarct area in mice brains (\(p < 0.05\)).\(^{43}\) In rats, kava extract administered according to the same regimen as used in mice significantly reduced infarct volume compared with control (\(p < 0.05\)).

Analgesic activity

Antinociceptive activity \textit{in vivo} (mice) in the tail immersion test has been documented for kava resin (150 mg/kg intraperitoneally) and for the individual kavalactones dihydrokawain, dihydromethysticin, kavain and methysticin at doses of 150, 275, 300 and 360 mg/kg (intraperitoneally),
respectively, compared with controls. Yangonin, tetrahydroyangonin, desmethoxyyangonin and dehydroyangonin had no or only a weak effect. Both kava resin (200 mg/kg, orally) and aqueous kava extract (250 mg/kg, intraperitoneally) displayed antinociceptive activity in the acetic acid-induced writhing test, also in mice. In further tests using both models, naloxone failed to reverse the antinociceptive effects of kava resin or aqueous kava extract, indicating that analgesic activity of kava is achieved via non-opiate pathways. Analgesic activity of dihydrokawain and dihydromethysticin has been reported previously.

Other activities
Kava extract has been reported to have a muscle relaxant effect in isolated frog muscles, thought to be due to a direct effect on muscle contractility. Reductions in contractions of isolated guinea-pig ileum induced by carbachol and by raised extracellular potassium ion concentrations have been documented for the synthetic kavalactone (±)-kavain, although the compound had no effect on caffeine-induced contractions of ileum strips or on calcium ion-induced contractions of skinned muscles. (±)-Kavain has also been reported to relax maximally contracted murine airway smooth muscle and to reduce carbachol- and potassium chloride-induced airway smooth muscle contraction. Further investigation indicated that nitric oxide and cyclooxygenase-mediated events did not play a role in kavain-induced relaxation.

In contrast, a previous study found that (+)-kavain inhibited human platelet aggregation in a dose-dependent manner in vitro. The formation of prostaglandin E$_2$ and thromboxane B$_2$ was also inhibited in a dose-dependent manner, suggesting that (+)-kavain is an inhibitor of cyclooxygenase.

An ethanol extract of kava and the isolated kavalactones dehydrokavain, dihydrokavain, kavain, yangonin and methysticin inhibited tumour necrosis factor alpha (TNFα) release in vitro from BALB/3T3 cells incubated with okadaic acid and suppressed lipopolysaccharide-induced TNFα production in vivo in diabetic mice following intraperitoneal administration.

Antifungal activity against several microorganisms, including Candida albicans, has been described for a 10% aqueous kava extract.

Clinical studies
Clinical trials of kava preparations have focused on investigating anxiolytic effects in various patient groups. Several trials have assessed effects in healthy volunteers, and others have explored the effects of kavain, a
A Cochrane systematic review of monopreparations of kava for the treatment of anxiety included 11 randomised, double-blind, placebo-controlled trials involving a total of 645 participants.\(^{(52)}\) All these trials tested the effects of a standardised (70% kavalactones) preparation of kava rhizome (WS-1490) at various dosages but typically equivalent to 60–240 mg kavalactones daily for four weeks. Only two trials were conducted for longer than four weeks; both used a dose equivalent to 210 mg kavalactones daily given for eight weeks in one study\(^{(53)}\) and 24 weeks in the other.\(^{(54)}\)

Six of the 11 trials, involving a total of 345 participants, used the total score on the Hamilton Anxiety Scale as their primary outcome measure, and their data were entered into a meta-analysis. It was reported that, overall, this showed a reduction in anxiety scale scores in kava recipients, compared with placebo recipients (weighted mean difference: 5.0; 95% confidence interval (CI), 1.1–8.8; \(p = 0.01\)). All except one of these trials included participants with non-psychotic anxiety; one study involved women with anxiety associated with the climacteric (perimenopausal period). Removing this trial from the meta-analysis reduced, but did not remove, the effect (weighted mean difference: 3.4; 95% CI, 0.5–6.4; \(p = 0.02\)).\(^{(52)}\)

Four of the studies not included in the meta-analysis reported statistically significant improvements for kava recipients, compared with placebo recipients, on outcomes (e.g. response rates, reduction in scores on various anxiety scales).\(^{(52)}\) The remaining study reported that the kava group, but not the placebo group, experienced a significant reduction in anxiety score, compared with baseline values; however, statistical analysis of the difference in reduction between the two groups was not reported. These five studies were heterogeneous in that they involved different patient groups, such as women with anxiety associated with the perimenopausal period, individuals with preoperative anxiety, and outpatients with neurotic anxiety. Consequently, dosage regimens of kava varied widely (e.g. equivalent to kavalactones 60 mg in the evening and 1 hour preoperatively, to 140 mg kavalactones daily for four weeks).

The conclusions of the review were that kava extract is an effective symptomatic treatment for anxiety, but that limitations of the studies included meant that further rigorous trials were needed.\(^{(52)}\) Some new research has been published since, but has added little to the evidence base because of methodological issues. For example, a study involving 68 perimenopausal women reported that kava extract 100 mg (containing 55% kavain) or 200 mg daily improved anxiety compared with no treatment, but
Results from some clinical studies have suggested that kava extracts may be as effective as certain standard anxiolytic agents, although this requires further investigation and confirmation. In a six-week, randomised, double-blind trial involving 172 patients with non-psychotic anxiety, the standardised kava extract WS-1490 (containing 70% kavalactones) 100 mg three times daily was as effective as oxazepam 5 mg and bromazepam 3 mg, each taken three times daily. Another randomised, double-blind, multicentre trial, involving 129 outpatients with generalised anxiety disorder, reported that the kava extract LI-150 400 mg (standardised to 30% kavapyrones = 120 mg) each morning for eight weeks was as effective as buspirone 5 mg twice daily and opipramol (a tricyclic antidepressant) 50 mg twice daily.

A randomised, placebo-controlled study involving 40 postmenopausal women with anxiety assessed the effects of a kava extract 100 mg daily (containing 55% kavain) given in addition to hormone replacement therapy (oestrogens plus progestogens or oestrogens alone). It was reported that women who received kava showed greater reductions in anxiety scores than women who received placebo. However, the study had various methodological limitations.

In a randomised, controlled study involving 54 healthy volunteers, kava extract (LI-150, equivalent to 120 mg kavalactones; \( n = 18 \)) and valerian extract 600 mg (LI-156; \( n = 18 \)), taken daily for one week, significantly reduced systolic blood pressure following mental stress tests, compared with baseline values, whereas no such reduction was observed in the no-treatment control group (\( p < 0.001 \) for both kava and valerian). No effect on diastolic blood pressure was recorded for either herbal preparation, and valerian, but not kava, appeared to reduce heart rate following mental stress tests. These findings require confirmation in placebo-controlled studies, and their relevance to everyday stress needs to be investigated.

Several other randomised, double-blind, controlled trials involving patients with anxiety have compared the effects of the synthetic kavalactone, (±)-kavain, administered at a dose of 200 mg three times daily for 3–4 weeks, with those of placebo, or benzodiazepines, such as oxazepam. One trial assessed the effects of kavain, or placebo, 200 mg three times daily for three weeks in 83 outpatients who had been treated with benzodiazepines for at least six weeks and who were undergoing benzodiazepine withdrawal. Generally, these studies have reported beneficial effects for kavain, but typically have involved only small numbers of patients.

**Pharmacokinetics**
Little is known about the clinical pharmacokinetics of kava preparations. Several metabolites of kavalactones have been detected and identified in human urine following ingestion of around 1 L of kava (prepared by the traditional method of aqueous extraction of kava rhizome) over 1 hour by healthy male volunteers before sleeping.\(^{(63)}\) Urine samples were collected before sleeping and on rising in the morning. Kawain, dihydrokawain, desmethoxyyangonin, tetrahydroyangonin, dihydromethysticin, 11-methoxytetrahydroyangonin, yangonin, methysticin and dehydromethysticin were detected unchanged in human urine; metabolic transformations observed included reduction of the 3,4-double bond and/or demethylation of the 4-methoxyl group of the kavalactone ring. The C\(_{12}\) hydroxy analogue of yangonin (12-hydroxy-12-desmethoxyyangonin) was also detected, and it may have been formed by demethylation of yangonin and/or C\(_{12}\) hydroxylation of desmethoxyyangonin. Dihydroxylated metabolites of the kavalactones and products from ring opening of the kavalactone ring were not detected.\(^{(63)}\)
Side-effects, Toxicity

In randomised, placebo-controlled trials involving different patient groups with anxiety, kava extracts generally have been well tolerated; adverse events reported, and their frequencies, are similar to those reported for placebo. However, clinical trials have the statistical power only to detect common, acute adverse effects.

Spontaneous reports of suspected adverse drug reactions associated with kava preparations have raised concerns over hepatotoxic reactions (see Hepatotoxicity).

A systematic review of eight placebo-controlled trials of kava extracts administered at doses equivalent to 55–240 mg kavalactones daily for two days to 24 weeks found that adverse events reported for both kava and placebo were most commonly gastrointestinal symptoms, tiredness, restlessness, tremor and headache.\(^{(64)}\) Three trials included in the review, one of which tested kava extract 100 mg daily (equivalent to 55 mg kavalactones) for 24 weeks,\(^{(58)}\) reported that adverse events were not observed in either the kava or placebo groups.

A similar finding was reported by a more recent Cochrane systematic review of monopreparations of kava for the treatment of anxiety (see Clinical studies).\(^{(52)}\) This review comprised seven of the eight placebo-controlled trials from the earlier review\(^{(64)}\) (the other trial was excluded from the Cochrane review because it tested kava extract in addition to hormone replacement therapy)\(^{(58)}\) and four new trials. One of the new trials (unpublished) reported that adverse events were not observed during four weeks’ treatment with kava extract 200 mg daily (equivalent to 140 mg kavalactones).\(^{(52)}\)

Randomised, double-blind clinical trials comparing kava extracts with certain benzodiazepines and other anxiolytic agents have also found kava to be well tolerated. In a six-week trial involving 172 patients with non-psychotic anxiety, gastrointestinal disturbances occurred in one of 57 participants who received WS-1490 100 mg three times daily (equivalent to 210 mg kavalactones daily), whereas tiredness, vertigo and pruritus occurred in seven of the remaining 115 participants who received oxazepam 5 mg or bromazepam 3 mg, both taken three times daily.\(^{(56)}\) In an eight-week trial involving 129 outpatients with generalised anxiety disorder, 14 of 43 (33%) participants who received the kava extract LI-150 400 mg (standardised to 30% kavalactones = 120 mg) each morning experienced adverse events, compared with 10 (24%) and 11 (26%) participants who received buspirone 5 mg twice daily and opipramol 50 mg twice daily, respectively.\(^{(57)}\) A total of
27 adverse events was reported in the kava group, compared with 16 and 14 for buspirone and opipramol, respectively. Adverse events reported for kava included upper respiratory tract infections, gastrointestinal disorders, weight changes, skin reactions and tachycardia, all of which were also reported for buspirone and/or opipramol.

These clinical trials and systematic reviews, however, provide only limited evidence to support the safety of kava extracts since they involved only small numbers of participants, involved different patient groups, tested different doses of kava extract (typically equivalent to 60–240 mg kavalactones daily), and most were of relatively short duration, usually around four weeks. Further, most trials investigated preparations of the kava extract WS-1490, and other standardised extracts of kava and kava preparations supplied by herbal medicine practitioners have undergone considerably less assessment.

Two post-marketing surveillance studies published in the early to mid-1990s involving patients treated in one study with WS-1490 150 mg daily (equivalent to 105 mg kavalactones; \( n = 4049 \)) and in the other with Antares 120 (equivalent to 120 mg kavalactones; \( n = 3029 \)) reported that the frequencies of adverse events were 1.5% and 2.3%, respectively.\(^{(64)}\) In both studies, adverse events commonly reported were mild gastrointestinal disorders and allergic reactions and, in the latter study, headaches and vertigo, which stopped when kava treatment was discontinued. A rather higher frequency of adverse events was reported during a post-marketing surveillance study carried out in Brazil.\(^{(65)}\) Among 850 participants with anxiety who received WS-1490 100 mg three times daily (equivalent to 210 mg kavalactones daily), 16.7% reported adverse events, most commonly fatigue/tiredness, nausea, confusion and gastrointestinal upset.

**Hepatotoxicity**

None of the clinical trials and post-marketing surveillance studies described above reported hepatotoxicity as an observed adverse event, although not all studies carried out liver function tests on participants. Six of the 11 trials included in the Cochrane review of the kava extract WS-1490 for the treatment of anxiety (see Clinical studies) did involve monitoring participants’ liver function (e.g. serum aspartate transaminase and alanine transaminase concentrations) and did not report any abnormalities in values obtained.\(^{(52)}\)

Over the years 2000 and 2001, a safety concern arose regarding cases of hepatotoxicity reported in association with the use of kava extracts. The signal first emerged in Switzerland, following a cluster of spontaneous reports to the medicines’ regulatory authority, and was strengthened a year or so later following further spontaneous reports from Switzerland and Germany. By July
2002, a total of 68 reports of liver toxicity associated with use of kava had been received by regulatory authorities in Canada, France, the UK and USA, as well as in Switzerland and Germany.\(^{(66)}\) The severity of the liver damage described in the reports varied from abnormal liver function test results to irreversible liver failure and death; six patients received liver transplants, one of whom, as well as two other individuals, subsequently died.

Three cases were reported in the UK, two of which described raised liver function test values in men aged 40 and 48 who had taken unspecified kava preparations for three months and eight years, respectively.\(^{(67)}\) Both stopped taking kava and their liver function test values normalised. The third UK case related to a woman (age not stated) who had taken kava 150 mg three times daily for two months, in addition to fluoxetine, and who experienced jaundice and raised liver function test values and was hospitalised for seven weeks.

In the UK, the Committee on Safety of Medicines (CSM) considered evidence for both the benefits and risks of kava preparations and considered whether only certain types of kava preparations might be associated with liver toxicity. However, there appeared to be no relationship between the method of processing/type of extract, strength or dose, and the reported adverse reactions. Therefore, on the basis of the data available, the CSM advised that the possible benefits of kava preparations do not outweigh the risks, and that kava has the potential to cause hepatotoxicity which could be serious in nature. The risk of serious liver toxicity was thought to be low, but idiosyncratic and so risk factors could not be identified. The CSM advised that kava should be prohibited in unlicensed medicines, and on 13 January 2003, a statutory order came into effect in the UK prohibiting the sale, supply and import of unlicensed medicines containing kava.\(^{(4)}\) Licensed kava products are required to remove kava as a herbal ingredient. Similar regulatory action was taken in Germany and several other countries. In the UK, evidence relating to the hepatotoxicity associated with kava will be reviewed after the prohibition has been in place for two years.

Reviews of German data relating to hepatotoxicity associated with kava have produced conflicting opinions on causality. One review emphasised that there was no dose–response relationship for kava-associated hepatotoxicity, and that crude estimates of incidence based on primary care data suggest that any risk of hepatotoxicity is similar to that of benzodiazepines.\(^{(68)}\) However, this conclusion is questionable since estimates of this nature can be inaccurate and misleading.

By contrast, a review of seven previously published and 29 unpublished case reports of kava-associated hepatotoxicity concluded that these data clearly
showed the potential for severe, unpredictable kava-related hepatotoxicity.\(^{(69)}\) Cases included nine individuals who developed fulminant hepatic failure, of whom six underwent successful liver transplantation (one only after retransplantation), two died after transplantation due to postoperative infectious complications, and one who was too old to undergo transplantation also died. All other cases, which comprised mostly cholestatic or necrotising hepatitis, underwent full recovery after withdrawal of kava treatment. Among these 36 reports, the relationship between kava ingestion and hepatotoxicity was considered ‘certain’ in three cases and ‘probable’ in 21.\(^{(69)}\) Most individuals were concurrently using other medication and several were regular consumers of alcohol.

A case report from Australia describes a 56-year-old woman who developed fatigue, nausea and jaundice after taking a preparation named ‘Kava 1800 Plus’ one tablet three times daily for around 10 weeks.\(^{(70)}\) Each tablet was stated to contain kavalactones 60 mg, \textit{Passiflora incarnata} 50 mg and \textit{Scutellaria lateriflora} 100 mg, although the latter ingredient was not identified in the product, so the precise composition of the product is unknown. She presented two weeks after first experiencing these symptoms and was hospitalised. Five days later, a biopsy revealed non-specific severe acute hepatitis with pan-acinar necrosis and collapse of hepatic lobules. She underwent liver transplantation on day 17 after admission, but the procedure was complicated and she died from progressive blood loss and circulatory failure. Subsequent examinations confirmed massive hepatic necrosis.

Inhibition of the cytochrome P450 drug metabolising enzyme CYP3A4 has been shown \textit{in vitro} for kava extracts and individual kavalactones (see Contra-indications, Warnings).\(^{(71)}\) The relevance of this for the hepatotoxic effects described for kava is not known; further work is needed to determine whether the inhibition of CYP3A4 by kava can lead to raised plasma concentrations of concurrently ingested drugs with hepatotoxic effects.

**Skin reactions**

An ichthyosiform (scaly, non-inflammatory), usually yellowish or whitish, skin condition termed kava dermopathy has been documented among kava users in Polynesia, Micronesia and Melanesia where powdered kava rhizome is prepared as a drink with cold water or coconut milk.\(^{(72)}\) The condition is reversible on stopping kava. Initially, it was thought that the condition was related to niacin deficiency, but this hypothesis was rejected following a small randomised, placebo-controlled trial of nicotinamide 100 mg daily for three weeks which showed no difference between groups.\(^{(73)}\)

Measures of health among 39 users of kava (prepared as a cold water
infusion of powdered kava rhizome) were compared with those of 34 age-matched non-users of kava in an Aboriginal community in the Northern Territory in Australia. Most (n = 35) were ‘heavy’ or ‘very heavy’ users of kava (310 g or more per week). It was reported that kava users were more likely to complain of poor health, and to have a scaly skin rash. However, the study had several methodological limitations (e.g. no correction for multiple statistical tests) and potential biases.

Several cases of allergic skin reactions have been reported in association with kava use. One case described a man who presented with oedema and severe non-pruritic erythema involving his upper body, head and neck, the morning after drinking several cups of ‘kava tea’. It was reported that the man had previously had a similar reaction to kava tea three months earlier whilst overseas and for which he was hospitalised and treated with intravenous corticosteroids. A case of systemic contact-type dermatitis following several weeks’ use of kava extract (Antares), chlorprothixene (an antipsychotic agent with properties similar to those of chlorpromazine) and diazepam has been described. Two further cases described a 70-year-old man and a 52-year-old woman who experienced skin eruptions (erythematous plaques and/or papules) in sebaceous gland-rich areas after using kava extract (no further details provided) for two to three weeks. Both patients were reported to display reactions to kava in diagnostic allergy or skin patch tests. Generalised erythema and papules with severe itching were described in a 36-year-old woman who had taken kava extract (Antares) 120 mg daily for three weeks. The rash, but not the itching, responded to short-term treatment with systemic corticosteroids, and six weeks later, patch test results for Antares were positive one day after application.

Central nervous system effects

Four cases of involuntary movements and dyskinesia associated with use of kava extracts have been reported, although causality has not been established; it has been stated that these symptoms suggest that constituents of kava may have antagonistic effects on central dopaminergic pathways. In three cases, involuntary movements involving the neck, head and/or trunk, and involuntary oral and lingual dyskinesia began within a few minutes to 4 hours after ingestion of kava extracts (Laitan 100 mg or Kavasporal forte 150 mg) for anxiety. One of these cases involved a 28-year-old man who had previously experienced three episodes of acute dystonic reactions following exposure to promethazine and fluspirilene, although he denied having used these medicines in relation to the current episode. The fourth case report described a 76-year-old woman being treated with levodopa 500 mg and benserazide 125 mg for Parkinson’s disease and who experienced an increase...
in the duration and frequency of her ‘off’ periods 10 days after starting Kavasporal forte 150 mg twice daily, prescribed by her physician for tension. (The ‘on–off’ phenomenon – sudden swings in mobility–immobility – occurs with long-term use of levodopa.) In all four cases, symptoms resolved on stopping kava or following treatment with biperiden administered intravenously.

Two other cases describe neurological symptoms following excessive use of traditional preparations of kava, i.e. as a beverage. A 27-year-old Aboriginal Australian man experienced generalised severe choreoathetosis (characterised by chorea and athetosis, a form of dyskinesia) without impairment of consciousness on three occasions after drinking large amounts of kava (precise quantity not specified). Routine investigations were normal, apart from raised liver function test values (serum alkaline phosphatase 162 IU/L, normal range 35–135 IU/L; gamma-glutamyltransferase 426 IU/L, normal range <60 IU/L). His symptoms responded to treatment with diazepam administered intravenously. Disorientation was reported in a 34-year-old Tongan man, a heavy user of kava (40 bowls daily for 14 years), who had ingested further excessive amounts of kava over the previous 12 hours. The man was treated in hospital with Plasmalyte intravenously and intramuscular thiamine and 5 hours after admission his symptoms had resolved.

A case of hypokalaemic renal tubular acidosis due to Sjögren’s syndrome (a symptom complex of unknown aetiology, marked by keratoconjunctivitis sicca, xerostamia, with or without lachrymal and salivary gland enlargement, respectively, and presence of connective tissue disease, usually rheumatoid arthritis, but sometimes systemic lupus erythematosus, scleroderma or polymyositis) has been reported in a 36-year-old woman. She was stated to have begun taking kava, echinacea and St John’s wort two weeks before becoming ill, but the report does not provide any further details of the echinacea species contained in the product(s), nor of the types of preparations, formulations, dosages and routes of administration of any of the herbal medicines listed. The woman was hospitalised with severe generalised muscle weakness and tests revealed she had a serum potassium ion concentration of 1.3 mEq/L. She was given electrolyte replacement for four days after which the muscle weakness resolved, and was started on hydroxychloroquine 200 mg daily for ‘probable’ Sjögren’s syndrome. The authors suggested that ingestion of echinacea may have aggravated an autoimmune disorder (see Echinacea, Side-effects, Toxicity) although causality has not been established.

Effects on mental performance
The effects of kava extracts and the synthetic kavalactone (±)-kavain on mental performance have been explored in studies involving healthy volunteers. Preliminary studies involving small numbers of volunteers have suggested that kava extract (WS-1490 200 mg three times daily for five days) did not appear to impair memory as assessed by certain tests (e.g. word recognition) carried out under laboratory conditions.\(^{(83)}\) In another series of tests, designed to assess mental alertness, volunteers received kava extract, Antares 120 (standardised to 120 mg kavalactones per tablet), one tablet daily, diazepam 10 mg daily, or placebo.\(^{(84)}\) It was reported that the experiments provided evidence that kava did not cause drowsiness or lack of concentration, for example, reaction time was reduced in placebo recipients, but not kava recipients. Other research involving volunteers found that a single dose of kava extract 600 mg (LI-158; drug–extract ratio, 12.5 : 1) led to a ‘moderate’ increase in tiredness, compared with placebo, and as assessed using visual analogue scale scores, although statistical analysis was not reported.\(^{(85)}\) Confirmation of these findings is required. In a battery of psychometric and other tests following administration of a range of single doses of the synthetic kavalactone (±)-kavain (200, 400 and 600 mg) and clobazam 30 mg to healthy volunteers, (±)-kavain appeared to have a sedative effect which was stated to be different to that observed with clobazam.\(^{(86)}\) Compared with placebo, (±)-kavain, but not clobazam, improved intellectual performance, attention, concentration and reaction time.

**Other reactions**

There are isolated reports of myopathy and myoglobinuria associated with the use of kava preparations, although causality in these cases has not been established.

One report described dermatomyositis associated with use of kava for anxiety by a 47-year-old woman.\(^{(87)}\) The woman, who had also been taking valproic acid for 18 months and sertraline occasionally over two years for bipolar disorder, developed a rash involving her back, neck and face, as well as muscle weakness, two weeks after taking kava (dosage not specified). She improved initially following treatment with methylprednisolone, but then developed a fever which prompted her to attend a hospital emergency department. Investigations revealed a raised serum creatine kinase concentration (8654 U/L, normal values stated as 24–170 U/L) and myopathic patterns in various muscles, and biopsy samples showed changes indicative of dermatomyositis. The woman was treated initially with parenteral prednisone, after which her creatine kinase concentration returned to normal, and also received methotrexate for five months and
hydroxychloroquine. Prednisone treatment was reduced over the following year, and at one year of follow-up the woman remained symptom-free.

Another isolated report describes a 29-year-old man who experienced severe muscle pain and passed dark urine one morning a few hours after having taken a herbal product said to contain kava 100 mg, *Ginkgo biloba* extract 200 mg and guarana (which contains methylxanthines) 500 mg (daily dosage was not stated), for the first time.\(^{(88)}\) The man was admitted to an intensive care unit and was found to have highly elevated serum creatine kinase (100 500 IU/L, normal range given as 0–195 IU/L) and myoglobin (10 000 ng/mL, normal range stated as 0–90 ng/mL) concentrations, but no renal complications. Investigations excluded metabolic myopathy as a possible cause; his signs and symptoms subsided over six weeks.

Disturbances of visual function have been reported following a study involving a 30-year-old kava-naïve male volunteer who ingested 600 mL of aqueous extract of pulverised kava ‘root’ (rhizome).\(^{(89)}\) Measurements involving the man’s right eye only indicated reductions in near point of accommodation and convergence, an increase in pupil diameter, and disturbance of oculomotor balance, but no effects on visual or stereoacuity or ocular refractive error. The experiment was not carried out according to a double-blind, controlled design and, therefore, the findings require further investigation.

**Toxicology**

A study involving small numbers of rats administered an aqueous (water) extract of kava ‘root’ equivalent to kavalactones 200 or 500 mg/kg/day for 2–4 weeks found that serum concentrations of the enzymes alanine aminotransferase, aspartate aminotransferase, alkaline phosphatase and lactate dehydrogenase were not elevated following kava administration, compared with control.\(^{(90)}\) The clinical relevance of these findings is not known.

In other toxicological studies, LD\(_{50}\) values for a standardised kava extract containing 70% kavalactones have been reported as 370 mg/kg and 16 g/kg for intraperitoneal and oral administration, respectively, in rats, and 380 mg/kg and 1.8 g/kg for intraperitoneal and oral administration, respectively, in mice.\(^{(G50)}\)
Contra-indications, Warnings

It has been stated that kava is contra-indicated in endogenous depression.\(^{G3, G4}\) Even when administered in accordance with recommended dosage regimens, kava may adversely affect motor reflexes, and may affect ability to drive and/or operate machinery.\(^{G3, G4}\)

It has been reported that there is no evidence that use of kava extracts has the potential for physical or psychological dependency to develop.\(^{G56}\) However, as most clinical studies of kava extracts have been of short duration, typically around four weeks (maximum 24 weeks) and/or usually have involved only small numbers of participants, further study is required before definitive statements are made on the potential for dependency with kava.

Interactions

There is an isolated report of a 54-year-old man who was taking alprazolam, cimetidine and terazosin and who became lethargic and disoriented three days after he began taking kava purchased from a health-food store (no further details of the kava preparation were provided).\(^{91}\) The man was hospitalised and his symptoms resolved after several hours. He tested negatively for alcohol, and positively for benzodiazepines; the man stated he had not taken overdoses of either alprazolam or kava. The clinical importance and role of kava in this reaction is not known, although there is a view that concurrent use of kava and substances with central nervous system effects could lead to enhanced activity.\(^{G3, G4}\)

The effects on performance of a kava extract given in combination with bromazepam have been explored in a randomised, double-blind, controlled crossover trial involving 18 healthy volunteers. Participants received a kava extract (Antares) equivalent to 120 mg kavalactones twice daily, or bromazepam 4.5 mg twice daily, or both agents, for 14 days.\(^{92}\) Significant reductions in indicators of performance, such as motor coordination, were reported for recipients of both kava and bromazepam, compared with recipients of kava alone, but there was no difference between kava plus bromazepam compared with bromazepam alone.

Isolated case reports of extrapyramidal symptoms associated with use of kava extracts have led to the suggestion that constituents of kava may have dopamine antagonist effects (see Side-effects, Toxicity, Central nervous system effects).\(^{79}\) On this basis, the potential for kava to interact with dopamine agonists or antagonists should be considered.
There are conflicting results from *in vitro* studies regarding the effects of the kavalactone (+)-kavain on cyclooxygenase activity.\(^{(48,49)}\) One study reported that (+)-kavain inhibited human platelet aggregation *in vitro* (see *In vitro* and animal studies, Other activities], although the clinical relevance of this, if any is not known. At present, there is insufficient evidence to warn against the concurrent use of kava preparations and antiplatelet agents.

Inhibition of the cytochrome P450 drug metabolising enzyme CYP3A4 by kava extracts and individual kavalactones has been described following *in vitro* studies. Methanolic, acetone and ethyl acetate extracts of kava rhizome significantly inhibited CYP3A4 activity, compared with control, at concentrations as low as 10 μg/mL (ethyl acetate extract).\(^{(71)}\) In other *in vitro* experiments, several individual kavalactones were tested for their effects on the activities of CYP1A2, CYP2C9, CYP2C19 and CYP2D6, as well as CYP3A4. Desmethoxyyangonin, dihydromethysticin and methysticin produced a dose-dependent inhibition of one or more of the CYP isoforms at concentrations of <10 μmol/L, considered as ‘potent’ inhibition (e.g. IC\(_{50}\) values for desmethoxyyangonin, dihydromethysticin and methysticin for CYP2C19 were 0.51, 0.43 and 0.93 μmol/L, respectively, and for dihydromethysticin and methysticin for CYP3A4 under certain assay conditions were 2.49 and 1.49 μmol/L, respectively).\(^{(93)}\) In several cases, this degree of inhibition was greater than that shown by positive controls which are known to produce clinically significant drug interactions.

The clinical relevance of these findings is not known, although the potential for kava extracts to interact with concurrently administered drugs metabolised mainly by CYP3A4 should be considered.

**ALCOHOL** The effects of concurrent use of kava extract and alcohol have undergone some investigation. In a randomised, double-blind, controlled trial, 20 healthy participants received kava extract (WS-1490; Laitan) 300 mg daily (equivalent to 210 mg kavalactones), or placebo, for eight days.\(^{(94)}\) Alcohol was ingested on days one, four and eight in quantities sufficient to achieve a blood alcohol concentration of 50 mg%; participants underwent a series of tests designed to assess psychomotor performance before and after alcohol consumption. The results indicated that there was no difference in performance between the kava and placebo groups, apart from one test (concentration) in which the kava group was reported to be superior to the placebo group.\(^{(94)}\)

A small study involving 40 healthy participants found that the concurrent ingestion of a kava beverage (350 mL of aqueous extract of Fijian kava) and alcohol 0.75 g/kg led to a greater reduction in cognitive performance,
as assessed by a series of tests, compared with that observed with ingestion of alcohol alone; ingestion of kava alone did not affect cognitive performance.\(^{95}\)

Studies in mice given ethanol (3.5 and 4 g/kg, intraperitoneally) and kava resin 200 or 300 mg/kg orally have demonstrated a prolongation of hypnotic effects.\(^{96}\)

**Pregnancy and lactation**

There is a lack of information on the use of kava preparations during pregnancy and breastfeeding. Given the lack of data, kava should be avoided during these periods.
The chemistry of kava is well documented (see Constituents) and there is strong evidence that the kavalactone constituents are responsible for the observed pharmacological activities.

Randomised, double-blind, placebo-controlled clinical trials of certain standardised kava preparations have shown beneficial effects on measures of anxiety, although because of methodological limitations of some studies, further well-designed trials are required to confirm the anxiolytic effects. Also, most trials have been carried out with one particular standardised kava extract (containing 70% kavalactones) and it cannot be assumed that the effects shown in these studies will be produced by other kava extracts. Clinical trials involving patients with anxiety have also compared well-defined standardised kava preparations with certain standard anxiolytic agents. While these studies have suggested that the kava extracts tested may be as effective as certain standard anxiolytic agents, further investigation is necessary. Data from pharmacological studies provide supporting evidence for the anxiolytic effects of kava, although many of the other traditional uses of kava (see Herbal use) have not been tested scientifically. Many pharmacological studies involving individual kavalactones have investigated the effects of the synthetic kavalactone (±)-kavain, rather than the natural compound (+)-kavain. Some studies have used both the natural compound and the synthetic racemate and have reported a lack of stereospecific effect.\(^{(29,36)}\)

In placebo-controlled clinical trials, standardised kava extracts generally have been well tolerated; reported adverse events have been mild and transient and similar in nature and frequency to those reported for placebo. Clinical trials, however, can provide only limited information on the safety profile of a medicine. Spontaneous reports of hepatotoxicity associated with the use of kava preparations have arisen since the year 2000. Although the risk of serious liver toxicity is thought to be low, the reaction is idiosyncratic. Against this background, kava was prohibited in unlicensed medicines in the UK in 2003,\(^{(4)}\) and in the EU, all licensed kava products were removed from the market. Regulatory action has also been taken in Canada and Australia (voluntary recall), and in the USA, consumers were warned of the risk of liver toxicity with use of kava-containing products.

Other adverse reactions documented for kava preparations include an ichthyosiform (scaly, non-inflammatory) skin condition, termed ‘kava dermopathy’, usually associated with the traditional method of preparing and ingesting kava (see Side-effects, Toxicity, Skin reactions).
Although kava is prohibited in the UK and several other countries, individuals may obtain kava preparations over the Internet. Healthcare professionals should be aware that patients may be taking herbal medicinal products containing kava, and in view of the reported inhibitory activity against certain cytochrome P450 drug metabolising enzymes, be vigilant to the potential for drug interactions (see Contra-indications, Warnings, Interactions). Healthcare professionals should enquire about use of kava in patients presenting with symptoms of hepatotoxicity (see Side-effects, Toxicity, Hepatotoxicity). Adverse reactions have been reported in association with use of ‘herbal ecstasy’ tablets, which often contain ephedrine alkaloids, although healthcare professionals should be aware that some products have been stated to contain kava.\(^{(97)}\)
References

See also General References G3, G4, G9, G32, G50, G56, G60, G66, G67, G69 and G70.

32. Baum SS et al. Effect of kava extract and individual kavapyrones on neurotransmitter levels in the nucleus accumbens of rats. *Prog Neuro-


17. Medicines Control Agency. Consultation MLX 286: Proposals to prohibit


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Lady’s Slipper
Species (Family)

*Cypripedium pubescens* Willd. (Orchidaceae) and other related species
Synonym(s)

American Valerian, Cypripedium, *Cypripedium calceolus* var.*pubescens* R.Br., Nerve Root

Related species also referred to as Lady’s Slipper include *Calypso bulbosa* (L.) Oakes (*Cypripedium bulbosum* L.) and *Cypripedium parviflorum* Salish
Part(s) Used

Rhizome, root
Pharmacopoeial and Other Monographs

BHP 1983\textsuperscript{(G7)}

PDR for Herbal Medicines 2nd edition (Nerve Root)\textsuperscript{(G36)}
Legal Category (Licensed Products)

GSL (Cypripedium)(G37)
Constituents

See General References G22 G48 G64.

Little chemical information has been documented. Lady’s slipper is stated to contain glycosides, resin, tannic and gallic acids (usually associated with hydrolysable tannins), tannins and a volatile oil.

Several quinones have been reported including cypripedin, stated to belong to a group of rare non-terpenoid phenanthraquinones and not previously isolated from natural sources.\(^1\)
Food Use

Lady’s slipper is not used in foods.
Herbal Use

Lady’s slipper is stated to possess sedative, mild hypnotic, antispasmodic and thymoleptic properties. Traditionally, it has been used for insomnia, hysteria, emotional tension, anxiety states, and specifically for anxiety states with insomnia.\(^{[G7 G64]}\)
Dosage

*Dried rhizome/root*
2–4 g or by infusion three times daily.\(^\text{G7}\)

*Liquid extract*
2–4 mL (1 : 1 in 45% alcohol) three times daily.\(^\text{G7}\)
Pharmacological Actions

None documented.
Side–effects, Toxicity

It has been stated that the roots may cause psychedelic reactions and large doses may result in giddiness, restlessness, headache, mental excitement and visual hallucinations.\(^{(G22)}\) Lady’s slipper is stated to be allergenic and contact dermatitis has been documented.\(^{(G51)}\) The sensitising property of lady’s slipper has been attributed to the quinone constituents.\(^{(1)}\)
Contra–indications, Warnings

Lady’s slipper may cause an allergic reaction in sensitive individuals.

**Pregnancy and lactation**
The safety of lady’s slipper has not been established. In view of the lack of phytochemical, pharmacological and toxicological information the use of lady’s slipper during pregnancy and lactation should be avoided.
Pharmaceutical Comment

Virtually no phytochemical or pharmacological data are available for lady’s slipper to justify its use as a herbal remedy. In view of the lack of toxicity data, excessive use should be avoided.
See also General References G7 G22 G31 G36 G37 G48 G51 G64.

Lemon Verbena
Species (Family)

*Aloysia triphylla* (L’Her.) Britton (Verbenaceae)
Synonym(s)

Aloysia citriodora (Cav.) Ort., Lippia citriodora (Ort.) HBK, Verbena citriodora Cav., Verbena triphylla L’Her.
Part(s) Used

Flowering top, leaf
Pharmacopoeial and Other Monographs

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Lemon verbena is not included in the GSL.\textsuperscript{(G37)}
Constituents
See General References G22 G34 G57 G64.

Flavonoids
Flavones including apigenin, chryso eriol, cirsimaritin, diosmetin, eupafolin, eupatorin, hispidulin, luteolin and derivatives, pectolinarigenin and salvigenin.\(^{(1)}\)

Volatile oils
Terpene components include borneol, cineol, citral, citronellal, cymol, eugenol, geraniol, limonene, linalool, β-pinene, nerol, and terpineol (monoterpenes), and α-caryophyllene, β-caryophyllene, myrcenene, pyrollic acid and isovalerianic acid (sesquiterpenes).\(^{(2)}\)
Food Use

In the USA, lemon verbena is listed as GRAS (Generally Recognised As Safe) for human consumption in alcoholic beverages. Lemon verbena is also used in herbal teas. (G57)
Herbal Use

Lemon verbena is reputed to possess antispasmodic, antipyretic, sedative and stomachic properties. It has been used for the treatment of asthma, cold, fever, flatulence, colic, diarrhoea and indigestion. (G38 G57 G64)
Dosage

*Decoction*
45 mL taken several times daily. (G34)
Pharmacological Actions

None documented.
Side-effects, Toxicity

None documented for lemon verbena. Terpene-rich volatile oils are generally regarded as irritant and may cause kidney irritation during excretion.
Contra-indications, Warnings

Individuals with existing renal disease should avoid excessive doses of lemon verbena in view of the possible irritant nature of the volatile oil.

**Pregnancy and lactation**

In view of the lack of pharmacological and toxicity data, and the potential irritant nature of the volatile oil, excessive doses of lemon verbena are best avoided during pregnancy and lactation.
Pharmaceutical Comment

Limited information is available on lemon verbena. The traditional uses are probably attributable to the volatile oil, for which many components have been identified, and to the flavone constituents. In the UK, lemon verbena is mainly used as an ingredient of herbal teas.
References

See also General References G22 G34 G36 G38 G57 G64.


Liferoot
Species (Family)

Senecio aureus L. (Asteraceae/Compositae)
Synonym(s)

Golden Ragwort, Golden Senecio, Squaw Weed
Part(s) Used

Herb
Pharmacopoeial and Other Monographs

BHP 1983\(^{(G7)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

Liferoot is not included in the GSL.\(^{(G37)}\)
Constituents

See General References G19 G64.

Limited information is documented regarding the constituents of liferoot, although it is well recognised that *Senecio* species contain pyrrolizidine alkaloids.

**Pyrrolizidine alkaloids**
Floridanine, florosenine, otosenine, senecionine.\(^{(1,2)}\)

The volatile oil composition of various *Senecio* species (but not *Senecio aureus*) has been investigated.\(^{(3)}\)
Food Use

Liferoot is not used as a food, although many *Senecio* species are used as a form of spinach in South Africa.
Herbal Use

Liferoot is stated to possess uterine tonic, diuretic and mild expectorant properties. Traditionally, it has been used in the treatment of functional amenorrhoea, menopausal neurosis and leucorrhoea (as a douche).\(^{(G7 \ G64)}\)
Dosage

**Herb**
1–4 g or by infusion three times daily.\(^{(G7)}\)

**Liquid extract**
14 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

No documented studies were located.
Side-effects, Toxicity

Liferoot contains pyrrolizidine alkaloids. The toxicity, primarily hepatic, of this class of compounds is well recognised in both animals and humans\(^{(G19)}\) (see Comfrey).
Contra–indications, Warnings

In view of the hepatotoxic pyrrolizidine alkaloid constituents, liferoot should not be ingested.\(^{(G19)}\)

**Pregnancy and lactation**

In view of the toxic constituents, liferoot is contraindicated during pregnancy and lactation.\(^{(G49)}\) Furthermore, liferoot is traditionally reputed to be an abortifacient, emmenagogue, and uterine tonic.\(^{(G7 G22)}\) In animals, placental transfer and secretion into breast milk\(^{(4)}\) has been documented for unsaturated pyrrolizidine alkaloids.
Pharmaceutical Comment

Little information is documented for liferoot. No pharmacological studies were found to substantiate the traditional uses. The *Senecio* genus is characterised by unsaturated pyrrolizidine alkaloid constituents and the hepatotoxicity of this class of compounds is well recognised (see Comfrey). In view of this, liferoot is not suitable for use as a herbal remedy.
References

See also General References G7 G22 G31 G32 G36 G43 G49 G64.


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Lime Flower
Species (Family)

i. *Tilia cordata* Mill. (Tiliaceae)

ii. *Tilia platyphyllos* Scop.

iii. *Tilia ×europaea* – hybrid of (i) and (ii)
Synonym(s)

Lime Tree, Linden Tree
Part(s) Used

Flowerheads
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

BP 2002\(^{(G71)}\)

Complete German Commission E (Linden)\(^{(G3)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition (Linden)\(^{(G36)}\)

Ph Eur 2004\(^{(G72)}\)
Constituents

See General References G2 G6 G22 G49 G62 G64.

**Acids**
Caffeic acid, chlorogenic acid and \( p \)-coumaric acid.

**Amino acids**
Alanine, cysteine, cystine, isoleucine, leucine, phenylalanine and serine.

**Carbohydrates**
Mucilage polysaccharides (3%). Five fractions identified yielding arabinose, galactose, rhamnose, with lesser amounts of glucose, mannose, and xylose; galacturonic and glucuronic acids;\(^{(1)}\) gum.

**Flavonoids**
Kaempferol, quercetin, myricetin and their glycosides.

**Volatile oil**
Many components including alkanes, phenolic alcohols and esters, and terpenes including citral, citronellal, citronellol, eugenol, limonene, nerol, \( \alpha \)-pinene and terpineol (monoterpenes), and farnesol (sesquiterpene).

**Other constituents**
Saponin (unspecified), tannin (condensed) and tocopherol (phytosterol).
Food Use

Lime flower is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that lime flower can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product. In the USA, lime flower is listed as GRAS (Generally Recognised As Safe).
Herbal Use

Lime flower is stated to possess sedative, antispasmodic, diaphoretic, diuretic and mild astringent properties. Traditionally it has been used for migraine, hysteria, arteriosclerotic hypertension, feverish colds, and specifically for raised arterial pressure associated with arteriosclerosis and nervous tension. (G2 G6 G7 G8 G64)
Dosage

*Flowerhead*
2–4 g by infusion.

*Liquid extract*
2–4 mL (1 : 1 in 25% alcohol).

*Tincture*
1–2 mL (1 : 5 in 45% alcohol).
Pharmacological Actions

In vitro and animal studies

In vitro, lime flower has been reported to exhibit antispasmodic activity followed by a spasmogenic effect on rat duodenum.\(^2\) The actions were inhibited by atropine and papaverine, and reinforced by acetylcholine. The diaphoretic and antispasmodic properties claimed for lime flower have been attributed to \(p\)-coumaric acid and the flavonoids.\(^{G39\text{--}G60}\) In addition, a number of actions have been associated with volatile oils including diuretic, sedative and antispasmodic effects, which may also account for some of the reputed uses of lime flower.\(^{3\text{--}5}\) Volatile oils are not thought to possess any true diuretic activity, but to act as a result of certain terpenoid components having an irritant action on the kidneys during renal excretion.

Lime flower has been documented to possess a restricted range of antifungal activity.\(^6\)
Side–effects, Toxicity

Excessive use of lime flower tea may result in cardiac toxicity.\(^{(G60)}\) However, the rationale for this statement is not included by the author.
Contra–indications, Warnings

It is advised that lime flower should be avoided by individuals with an existing cardiac disorder. (G22 G39 G60)

Pregnancy and lactation
The safety of lime flower has not been established. In view of the lack of toxicological data, excessive use of lime flower during pregnancy and lactation should be avoided.
Pharmaceutical Comment

The chemistry of lime flower is well documented. Little scientific information was located to justify the reputed herbal uses of lime flower, although some correlation can be made with the known pharmacological activities of the reported constituents. The lack of toxicological data, together with a warning concerning cardiac toxicity, indicates that excessive use of lime flower should be avoided.
References


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Liquorice
Species (Family)

*Glycyrrhiza glabra* L. (Leguminosae)
Synonym(s)
Licorice
Part(s) Used

Root, stolon
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

BP 2002\(^{(G71)}\)

Complete German Commission E\(^{(G3)}\)

Martindale 33rd edition\(^{(G67)}\)

Mills and Bone\(^{(G50)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)

Ph Eur 2004\(^{(G72)}\)

WHO volume 1 1999\(^{(G63)}\)
Legal Category (Licensed Products)

GSL$_{G37}$
Constituents

See General References G2 G6 G41 G48 G64.

Coumarins
Glycyrin, heniarin, liqcoumarin, umbelliferone, GU-7 (3-arylcoumarin derivative).\(^{(1)}\)

Flavonoids
Flavonols and isoflavones including formononetin, glabrin, glabrol, glabrone, glyzarin, glycyrol, glabridin and derivatives, kumatakenin, licoflavonol, licoisoflavones A and B, licoisoflavanone, licoricone, liquiritin and derivatives, phaseollininisoflavan;\(^{(2)}\) chalcones including isoliquiritigenin, licuraside, echinatin, licochalcones A and B, neo licurosides.\(^{(3)}\)

Terpenoids
Glycyrrhizin glycoside (1–24%) also known as glycyrrhizic or glycyrrhizinic acid yielding glycyrrhetinic (or glycyrrhetic) acid and glucuronic acid following hydrolysis;\(^{(4)}\) glycyrrhetol, glabrolide, licoric acid, liquiritic acid and β-amyrin.

Volatile oils
0.047%.\(^{(5)}\) More than 80 components identified including anethole, benzaldehyde, butyrolactone, cumic alcohol, eugenol, fenchone, furfuryl alcohol, hexanol, indole, linalool, γ-nonanalactone, oestragole, propionic acid, α-terpineol and thujone.\(^{(5)}\)

Other constituents
Amino acids, amines, gums, lignin, starch, sterols (β-sitosterol, stigmasterol), sugars and wax.

Other plant parts
Components documented for the leaves of *G. glabra* include flavonoids (kaempferol and derivatives, isoquercetin, quercetin and derivatives, phytoalexins), coumarins (bergapten, xanthotoxin), phytoestrogen, β-sitosterol and saponaretin.\(^{(6)}\)
Food Use

Liquorice is widely used in foods as a flavouring agent. Liquorice root is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that liquorice can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\(^{(G16)}\) In the USA, liquorice is listed as GRAS (Generally Recognised As Safe).\(^{(G41)}\)

\(^{(G16)}\): Reference to the Council of Europe's classification of liquorice as a natural source of food flavouring.

\(^{(G41)}\): Reference to the GRAS status of liquorice in the USA.
Herbal Use

Liquorice is stated to possess expectorant, demulcent, antispasmodic, anti-inflammatory and laxative properties. Traditionally, it is also reported to affect the adrenal glands. It has been used for bronchial catarrh, bronchitis, chronic gastritis, peptic ulcer, colic and primary adrenocortical insufficiency.
Dosage

*Powdered root*
1–4 g or by decoction three times daily. (G6 G7)

*Liquorice Extract*
(BPC 1973) 0.6–2.0 g.
Pharmacological Actions

The pharmacological actions of liquorice have been reviewed.\(^{(7,8)}\)

**In vitro and animal studies**

Much has been documented regarding the steroid–type actions of liquorice (see Side–effects, Toxicity). Both glycyrrhizin and glycyrrhetinic acid (GA) have been reported to bind to glucocorticoid and mineralocorticoid receptors with moderate affinity, and to oestrogen receptors, sex hormone–binding globulin and corticosteroid–binding globulin with very weak affinity.\(^{(9–11)}\) It has been suggested that glycyrrhizin and glycyrrhetinic acid may influence endogenous steroid activity via a receptor mechanism, with displacement of corticosteroids or other endogenous steroids.\(^{(9)}\)

The anti–oestrogenic action documented for glycyrrhizin at relatively high concentrations has been associated with a blocking effect that would be caused by glycyrrhizin binding at oestrogen receptors.\(^{(9)}\) However, oestrogenic activity has also been documented for liquorice and attributed to the isoflavone constituents.\(^{(8)}\) Liquorice exhibits an alternative action on oestrogen metabolism, causing inhibition if oestrogen concentrations are high and potentiation when concentrations are low.\(^{(8)}\)

The relatively low affinity of glycyrrhizin and glycyrrhetinic acid for binding to mineralocorticoid receptors, together with the fact that liquorice does not exert its mineralocorticoid activity in adrenalectomised animals, indicates that a direct action at mineralocorticoid receptors is not the predominant mode of action.\(^{(12)}\) It has been suggested that glycyrrhizin and glycyrrhetinic acid may exert their mineralocorticoid effect via an inhibition of 11β-hydroxysteroid dehydrogenase (11β-OHSD).\(^{(12)}\) 11β-OHSD is a microsomal enzyme complex found predominantly in the liver and kidneys which catalyses the conversion of cortisol (potent mineralocorticoid activity) to the inactive cortisone. Deficiency of 11β-OHSD results in increased concentrations of urinary free cortisol and cortisol metabolites. Glycyrrhetinic acid has been shown to inhibit renal 11β-OHSD in rats.\(^{(12)}\) It has also been proposed that glycyrrhizin and glycyrrhetinic acid may displace cortisol from binding to transcortin.\(^{(13)}\)

Antiplatelet activity *in vitro* has been documented for a 3–arylcoumarin derivative, GU–7, isolated from liquorice.\(^{(1)}\) GU–7 was thought to inhibit platelet aggregation by increasing intraplatelet cyclic AMP concentration.

Isoliquiritigenin has been reported to inhibit aldose reductase, the first enzyme in the polyol pathway which reduces glucose to sorbitol.\(^{(14)}\)
Isoliquiritigenin was subsequently found to inhibit sorbitol accumulation in human red blood cells \textit{in vitro}, and in red blood cells, the sciatic nerve and the lens of diabetic rats administered isoliquiritigenin intragastrically.\textsuperscript{(14,15)} Many diabetic complications, such as cataracts, peripheral neuropathy, retinopathy and nephropathy have been associated with the polyol pathway and have shown improvement with inhibitors of aldose reductase.\textsuperscript{(14,15)}

Significant anti-inflammatory action is exhibited by glycyrrhetinic acid against UV erythema.\textsuperscript{(16)} 18\alpha-Glycyrrhetinic acid has exhibited stronger anti-inflammatory action compared to its stereoisomer 18\beta-glycyrrhetinic acid.\textsuperscript{(17)} Chalcones isolated from \textit{G. inflata} Bat. have been reported to inhibit leukotriene production and increase cyclic AMP concentrations in human polymorphonuclear neutrophils \textit{in vitro}.\textsuperscript{(18)} Glycyrrheticin acid derivatives, but not glycyrrhetinic acid, have exhibited inhibitory effects on writhing and vascular permeability tests and on type IV allergy in mice.\textsuperscript{(19)} The dihemiphthalate derivatives were especially active with respect to the two former activities and have previously been found to inhibit lipoxygenase and cyclooxygenase activities, and to prevent formation of gastric ulcer.\textsuperscript{(19)}

Glycyrrhetinic acid is known to inhibit Epstein–Barr virus activation by tumour promotors.\textsuperscript{(20)}

Antimicrobial activity versus \textit{Staphylococcus aureus}, \textit{Mycobacterium smegmatis} and \textit{Candida albicans} has been documented for liquorice and attributed to isoflavonoid constituents (glabridin, glabrol and their derivatives).\textsuperscript{(2)} Antiviral activity has been described for glycyrrhetinic acid, which interacts with virus structures producing different effects according to the viral stage affected.\textsuperscript{(21)} Activity was observed against vaccinia, herpes simplex 1, Newcastle disease and vesicular stomatitis viruses, with no activity demonstrated towards poliovirus 1.\textsuperscript{(21)}

\textit{In vitro} hepatoprotective activity against CCl\textsubscript{4}-induced toxicity has been reported to be greater for glycyrrhetinic acid compared to glycyrrhizin.\textsuperscript{(22)} Glycyrrhetinic acid is thought to act by inhibition of the cytochrome P450 system required for the metabolism of CCl\textsubscript{4} to the highly reactive radical CCl\textsubscript{3}.\textsuperscript{(22)} \textit{Glycyrrhiza uralensis} Fisch is used to treat hepatitis B in China, with a success rate reported to be greater than 70\%.\textsuperscript{(23)} Other activities documented for \textit{G. uralensis} are anti-inflammatory and anti-allergic, treatment of jaundice, inhibition of fibrosis of the liver, corticosteroid–like immunosuppressing effect and a detoxifying effect.\textsuperscript{(23)}

Screening of several plant extracts for antifertility activity reported liquorice to be ineffective following oral administration to rats in days 1–7 of
pregnancy. (24)

Clinical studies

Carbenoxolone, an ester derivative of glycyrrhetinic acid, has been used in the treatment of gastric and oesophageal ulcers. It is thought to exhibit a mucosal–protecting effect by beneficially interfering with gastric prostanoid synthesis, and increasing mucous production and mucosal blood flow. (25)

Liquorice is thought to exert its mineralocorticoid effect by inhibition of the enzyme 11β-OHSD, which catalyses the conversion of cortisol to the inactive cortisone (see In vitro and animal studies). Administration of liquorice to healthy volunteers has resulted in a disturbance of cortisol metabolism and a significant rise in urinary free cortisol, despite there being no change in plasma concentrations. These changes are consistent with this hypothesis, being indicative of 11β-OHSD deficiency. (12) Liquorice has also been found to suppress both plasma renin activity and aldosterone secretion. (26–28)

The pharmacokinetic profile of glycyrrhizin in rats has been found to be similar to that observed in humans. (29) Glycyrrhizin is primarily (80%) excreted into the bile from the liver against a concentration gradient. (29) This process is saturable and can therefore affect the excretion rate of glycyrrhizin. In addition, enterohepatic recycling occurs with reabsorption of bile–excreted glycyrrhizin from the intestinal tract. (29) Subjects consuming 100–200 g liquorice/day have been reported to achieve plasma glycyrrhetic acid concentrations of 80–480 ng/mL. (12)
Side-effects, Toxicity

Apart from confectionery, liquorice can also be ingested from infusions and by chewing tobaccos. Excessive or prolonged liquorice ingestion has resulted in symptoms typical of primary hyperaldosteronism, namely hypertension, sodium, chloride and water retention, hypokalaemia and weight gain, but also in low levels of plasma renin activity, aldosterone and antidiuretic hormone.\(^{(13,26,30)}\)

Raised concentrations of atrial natriuretic peptide (ANP), which is secreted in response to atrial stretch and has vasodilating, natriuretic and diuretic properties, have also been observed in healthy subjects following the ingestion of liquorice.\(^{(13)}\) Individuals consuming between 10–45 g liquorice/day have exhibited raised blood pressure, together with a block of the aldosterone/renin axis and electrocardiogram changes, which resolved one month after withdrawal of liquorice.\(^{(31)}\) Individuals consuming vastly differing amounts of liquorice have exhibited similar side-effect symptoms, indicating that the mineralocorticoid effect of liquorice is not dose dependent and is a saturable process.\(^{(31)}\)

Hypokalaemic myopathy has also been associated with liquorice ingestion.\(^{(32–36)}\) Severe hypokalaemia with rhabdomyolysis has been documented in a male patient following the ingestion of an alcohol-free beverage containing only small amounts of glycyrrhetic acid (0.35 g/day).\(^{(32)}\) The patient had known liver cirrhosis due to alcohol consumption and it was suggested that cirrhotic patients may be more susceptible to the mineralocorticoid side-effects of liquorice.\(^{(32)}\) In one case,\(^{(34)}\) the myoglobinaemia led to glomerulopathy and tubulopathy but with no clinical evidence of acute renal failure (ARF). The latter was attributed to the volume expansion also caused by the liquorice ingestion.

Rhabdomyolysis without myoglobinuria has been described.\(^{(37)}\) In addition, severe congestive heart failure and pulmonary oedema have been reported in a previously healthy man who had ingested 700 g liquorice over eight days.\(^{(30)}\) Liquorice extract given orally has been reported to have a similar but longer lasting action to intravenous deoxycortone and it has been noted that sodium, chloride and water retention do not have to be accompanied by clinical oedema.\(^{(38)}\) Amenorrhoea has been associated with liquorice ingestion (anti-oestrogenic action), with the menstrual cycle re-appearing following the withdrawal of liquorice.\(^{(31)}\)

It has been noted that symptoms of hyperaldosteronism often resolve quickly, within a few days to two weeks, following the withdrawal of liquorice, even in
individuals who have ingested the substance for many years.\textsuperscript{(28)}

A case has been described where a patient presented with symptoms related to hyperglycaemia and myopathy secondary to liquorice-induced hypokalaemia. An inverse relationship was observed between the concentrations of fasting serum glucose and serum potassium.\textsuperscript{(39)} Interestingly, animal studies have indicated that liquorice may reduce diabetic complications associated with intracellular accumulation of sorbitol.\textsuperscript{(19)}
Contra–indications, Warnings

Numerous instances have been documented where liquorice ingestion has resulted in symptoms of primary hyperaldosteronism, such as water and sodium retention and hypokalaemia. Liquorice should therefore be avoided completely by individuals with an existing cardiovascular–related disorder, and ingested in moderation by other individuals. Hypokalaemia is known to aggravate glucose intolerance and liquorice ingestion may therefore interfere with existing hypoglycaemic therapy. Liquorice may interfere with existing hormonal therapy (oestrogens and antioestrogenic activities documented \textit{in vivo}).

\textbf{Pregnancy and lactation}

In view of the oestrogenic and steroid effects associated with liquorice, which may exacerbate pregnancy–related hypertension, excessive ingestion during pregnancy and lactation should be avoided. In addition, liquorice has exhibited a uterine stimulant activity in animal studies, and is traditionally reputed to be an abortifacient and to affect the menstrual cycle (emmenagogue).\textsuperscript{(G30)}
Pharmaceutical Comment

The phytochemistry is well documented for liquorice and it is particularly characterised by triterpenoid components. Many of the traditional uses of liquorice are supported by documented pharmacological data although limited evidence of antispasmodic activity was found. Carbenoxolone, an ester derivative of a triterpenoid constituent in liquorice, is well known for its use in ulcer therapy. Much has been written concerning the steroid-type adverse effects associated with liquorice ingestion. Liquorice ingestion should therefore be avoided by individuals with an existing cardiovascular disorder and moderate consumption should be observed by other individuals.
References


22. Piette AM et al. Hypokaliémie majeure avec rhabdomyolase secondaire à


Lobelia
Species (Family)

*Lobelia inflata* L. (Campanulaceae)
Synonym(s)

Indian Tobacco
Part(s) Used

Herb
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL\(\text{G}^{37}\)
Constituents
See General References G6 G22 G41 G64.

Alkaloids
Piperidine-type. 0.48%. Lobeline (major); others include lobelanine, lobelanidine, norlobelanine, lelobanidine, norlelobanidine, norlobelanidine and lobinine.

Other constituents
Bitter glycoside (lobelacrin), chelidonic acid, fats, gum, resin and volatile oil.
Food Use

Lobelia is not generally used as a food.
Herbal Use

Lobelia is stated to possess respiratory stimulant, antasthmatic, antispasmodic, expectorant, and emetic properties. Traditionally, it has been used for bronchitic asthma, chronic bronchitis, and specifically for spasmodic asthma with secondary bronchitis. It has also been used topically for myositis and rheumatic nodules.
Dosage

Dried herb
0.2–0.6 g or by infusion or decoction three times daily.\(^{(G6 \ G7)}\)

Liquid extract
0.2–0.6 mL (1 : 1 in 50% alcohol) three times daily.\(^{(G6 \ G7)}\)

Simple Tincture of Lobelia
(BPC 1949) 0.6–2.0 mL.

Tincture Lobelia Acid
1–4 mL (1 : 10 in dilute acetic acid) three times daily.\(^{(G6 \ G7)}\)
Pharmacological Actions

The pharmacological activity of lobelia can be attributed to the alkaloid constituents, principally lobeline. Lobeline has peripheral and central effects similar to those of nicotine, but is less potent. Hence, lobeline initially causes CNS stimulation followed by respiratory depression. Lobeline is also reported to possess expectorant properties.
Side–effects, Toxicity

Side–effects of lobeline and lobelia are similar to those of nicotine and include nausea and vomiting, diarrhoea, coughing, tremors and dizziness. Symptoms of overdosage are reported to include profuse diaphoresis, tachycardia, convulsions, hypothermia, hypotension and coma, and may be fatal.\(^{G45}\)
Contra–indications, Warnings

The pharmacological actions of lobeline are similar to those of nicotine.

Pregnancy and lactation
Lobelia should not be used during pregnancy or lactation.
Pharmaceutical Comment

The principal constituent of lobelia is lobeline, an alkaloid with similar pharmacological properties to nicotine. Lobelia has previously been used in herbal preparations for the treatment of asthma and bronchitis, and in anti-smoking preparations aimed to lessen nicotine withdrawal symptoms. However, in view of its potent alkaloid constituents, excessive use of lobelia is not recommended.
References


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Marshmallow
Species (Family)

*Althaea officinalis* L. (Malvaceae)
Synonym(s)
Althaea
Part(s) Used
Leaf, root
Pharmacopoeial and Other Monographs

BHC 1992\textsuperscript{(G6)}

BHP 1996\textsuperscript{(G9)}

BP 2002 (root)\textsuperscript{(G71)}

Complete German Commission E\textsuperscript{(G3)}

ESCOP 1996\textsuperscript{(G52)}

Martindale 33rd edition\textsuperscript{(G67)}

PDR for Herbal Medicines 2nd edition\textsuperscript{(G36)}

Ph Eur 2004 (root and leaf)\textsuperscript{(G72)}
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents
See General References G2 G6 G41 G48 G52 G60 G64.

**Polysaccharides**
Mucilage polysaccharides (5–10%), consisting of galacturono–rhamnans, arabinans, glucans, arabinogalactans.\(^1,G52\)

**Flavonoids**
Hyolaetin 8–glucoside, isoscutellarin 4′-methylether–8–glucoside–2′′-sulfate.\(^2\)

**Phenolic acids**
Caffeic, \(p\)-coumaric, ferulic, \(p\)-hydroxybenzoic and syringic.

**Other constituents**
Asparagine 2%, calcium oxalate, coumarins (scopoletin), pectin, starch and tannin.
Food Use

Marshmallow is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that marshmallow can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product. In the USA, marshmallow is approved for use in foods.
Marshmallow is stated to possess demulcent, expectorant, emollient, diuretic, antilithic and vulnerary properties. Traditionally, it has been used internally for the treatment of respiratory catarrh and cough, peptic ulceration, inflammation of the mouth and pharynx, enteritis, cystitis, urethritis and urinary calculus, and topically for abscesses, boils and varicose and thrombotic ulcers. The German Commission E approved use of root and leaf for irritation of oral and pharyngeal mucosa and associated dry cough and root for mild inflammation of gastric mucosa.\(^{(G3)}\) Marshmallow root is used in combination with anise fruit, eucalyptus oil, liquorice and with anise fruit, liquorice and primrose root and with anise fruit and primrose root for catarrh of the upper respiratory tract and resulting dry cough.\(^{(G3)}\)
Dosage

*Dried leaf*
2–5 g or by infusion three times daily;\(^{(G6 \ G7)}\) 5 g.\(^{(G3)}\)

*Leaf, liquid extract*
2–5 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G6 \ G7)}\)

*Ointment*
5% Powdered althaea leaf in usual ointment base three times daily.\(^{(G6 \ G7)}\)

*Dried root*
2–5 g or by cold extraction three times daily;\(^{(G6 \ G7)}\) 6 g.\(^{(G3)}\)

*Root, liquid extract*
2–5 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G6 \ G7)}\)

*Syrup of Althaea*
(BPC 1949) 2–10 mL three times daily.\(^{(G6 \ G7)}\)
Pharmacological Actions

In vitro and animal studies

Antimicrobial activity against *Pseudomonas aeruginosa*, *Proteus vulgaris* and *Staphylococcus aureus* has been documented for marshmallow.\(^3\)

The mucilage has demonstrated considerable hypoglycaemic activity in non-diabetic mice.\(^4\)

Inhibition (17%) of mucociliary transport in ciliated epithelium isolated from frog oesophagus was observed with 200 μL of cold macerate of marshmallow root (6.4 g/140 mL).\(^G52\)

Marshmallow root extract is reported to stimulate phagocytosis, and to release oxygen radicals and leukotrienes from human neutrophils.\(^G52\) In addition, release of cytokines, interleukin 6 and tumour necrosis factor from monocytes occurs, demonstrating potential anti-inflammatory and immunomodulatory effects. In mice, intraperitoneal administration of isolated polysaccharide (10 mg/kg) resulted in activity of macrophages in a carbon clearance test, and was indicative of non-specific immunomodulation.\(^G52\) A lack of anti-inflammatory activity has been observed for marshmallow in the carrageenan-induced rat paw oedema test.\(^5\) The anti-inflammatory effect of an ointment containing 0.05% dexamethasone was enhanced by addition of aqueous extract of marshmallow (20%) as assessed in a rabbit ear irritancy test using UV irradiation or furfuryl alcohol.\(^G52\)

A total extract of root and isolated polysaccharide (100 and 50 mg/kg, respectively) have been tested for their antitussive activity in unanaesthetised cats.\(^6\) The polysaccharide gave a statistically significant decrease in the number of cough efforts from laryngopharyngeal and tracheobronchial areas. The root extract was less effective than the isolated polysaccharide.

A polysaccharide enriched extract showed moderate concentration-dependent adhesive properties in porcine buccal membranes *ex vivo*.\(^7\)
Side-effects, Toxicity

None documented.
Contra–indications, Warnings
Marshmallow may interfere with existing hypoglycaemic therapy and the absorption of other drugs taken simultaneously may be retarded.\textsuperscript{(G52)}

\textbf{Pregnancy and lactation}
There are no known problems with the use of marshmallow during pregnancy or lactation.
Pharmaceutical Comment

*In vitro* and animal studies provide some supporting evidence for the use of marshmallow in the treatment of cough, irritation of the throat and gastric inflammation. Antibacterial and anti-inflammatory activities, effects on mucociliary transport, adhesion of polysaccharide to buccal membranes and reduction of cough are reported. However, there is a lack of clinical studies investigating the effects of marshmallow. Although no toxicity data were located, the chemistry of marshmallow and its use in foods indicate that there should not be any reason for concern regarding safety.
References

See also General References G2 G3 G6 G9 G10 G16 G31 G36 G37 G43 G48 G52 G54 G60 G64


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Maté
Species (Family)

*Ilex paraguariensis* St. Hil. (Aquifoliaceae)
Synonym(s)
Ilex, Jesuit’s Brazil Tea, Paraguay Tea, St. Bartholomew’s Tea, Yerba Maté
Part(s) Used

Leaf
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)

Complete German Commission E\(^{(G3)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL($G^{37}$)
Constituents
See General References G2 G22 G45 G48 G64.

**Alkaloids**
Xanthine–type. Caffeine 0.2–2.0%, theobromine 0.1–0.2%, theophylline 0.05%.

**Flavonoids**
Kaempferol, quercetin, and their glycosides, including rutin.(1)

**Tannins**
4–16%.

**Terpenoids**
Ursolic acid (major), β-amyrin, ilexoside A, ilexoside B methyl ester.(2)

**Other constituents**
Choline and trigonellin (amines), amino acids,(1) riboflavin (vitamin B$_2$), pyridoxine (vitamin B$_6$), niacin, pantothenic acid, vitamin C and resins.

**Other Ilex species**
Triterpenoid saponins termed ilexsaponins B$_1$, B$_2$, and B$_3$ have been isolated from *Ilex pubescens* Hook. & Arn.(3)

A cyanogenetic glucoside has been isolated from *Ilex aquifolium*.\(^{(4)}\)
Food Use

Maté is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that maté can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product. Maté is commonly consumed as a beverage. It is stated to be less astringent than tea. In the USA, maté is listed as GRAS (Generally Recognised As Safe).
Herbal Use

Maté is stated to possess CNS-stimulant, thymoleptic, diuretic, antirheumatic and mild analgesic properties. Traditionally, it has been used for psychogenic headache and fatigue, nervous depression, rheumatic pains, and specifically for headache associated with fatigue. (G2 G7 G8 G64)
Dosage

*Dried leaf*
2–4 g or by infusion three times daily. (G6 G7)

*Liquid extract*
2–4 mL (1:1 in 25% alcohol) three times daily. (G6 G7)
Pharmacological Actions

In vitro and animal studies

*In vivo* hypotensive activity in rats has been reported for an aqueous extract of *Ilex pubescens* (commonly referred to as maodong qing or MDQ). It was concluded that intravenous administration of MDQ releases histamine.\(^5\)

Clinical studies

The xanthine constituents, in particular caffeine, are the active principles in maté. The pharmacological actions of caffeine are well documented and include stimulation of the CNS, respiration and skeletal muscle, in addition to cardiac stimulation, coronary dilation, smooth muscle relaxation and diuresis.\(^{41}\) Reduction of appetite has been documented for maté.\(^1\)

In China, MDQ is used parenterally for the treatment of cardiovascular diseases (hypotensive action).\(^1\)
Side–effects, Toxicity

Side–effects generally associated with xanthine– containing beverages include sleeplessness, anxiety, tremor, palpitations and withdrawal headache.

Veno–occlusive disease of the liver in a young woman has been attributed to the consumption of large quantities of mate over a number of years.\(^{(G45)}\) The association between consumption of mate infusions and oesophageal cancer has been investigated in Uruguay, where oesophageal cancer constitutes a major public health problem.\(^{(6,7)}\) Heavy consumption was reported to elevate the relative risk of oesophageal cancer by 6.5 and 34.6 in men and women, respectively.

The fatal dose of caffeine in humans is stated to be 10 g.\(^{(G41)}\)
Contra-indications, Warnings

Warnings generally associated with caffeine are applicable, such as restricted intake by individuals with hypertension or a cardiac disorder.

Pregnancy and lactation
It is generally recommended that caffeine consumption should be restricted during pregnancy, although conflicting results have been documented concerning the association between birth defects and caffeine consumption. In view of this, excessive consumption of maté during pregnancy should be avoided. Caffeine is excreted in breast milk, but at concentrations too low to represent a hazard to breastfed infants. As with all xanthine–containing beverages, excessive consumption of maté by breastfeeding mothers should be avoided.
Pharmaceutical Comment

Maté is characterised by the xanthine constituents, which also represent the active principles. The herbal uses of maté can be attributed to the pharmacological actions of caffeine, which are well documented. Side–effects and warnings associated with other xanthine–containing beverages, such as tea and coffee, are applicable to maté.
References

See also General References G2 G3 G9 G10 G16 G22 G31 G36 G37 G41 G43 G48 G64.

Species (Family)

*Filipendula ulmaria* (L.) Maxim. (Rosaceae)
Synonym(s)
Dropwort, Filipendula, Queen of the Meadow, *Spiraea ulmaria* L.
Part(s) Used
Herb
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

Complete German Commission E\(^{(G3)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL$^{(G37)}$
Constituents
See General References G2 G6 G22 G42 G62 G64.

Flavonoids
Flavonols, flavones, flavanones and chalcone derivatives (e.g. hyperoside\(^{(1)}\) and spireoside,\(^{(2)}\) kaempferol glucoside\(^{(3)}\) and avicularin.\(^{(4)}\)

Salicylates
Main components of the volatile oil including salicylaldehyde (major, up to 70%), gaultherin, isosalicin, methyl salicylate, monotropitin, salicin, salicylic acid and spirein.\(^{(5–8)}\)

Tannins
1% (alcoholic extract), 12.5% (aqueous extract).\(^{(5)}\) Hydrolysable type;\(^{(9)}\) leaf extracts have also yielded catechols,\(^{(1)}\) compounds normally associated with condensed tannins.

Volatile oils
Many phenolic components including salicylates (see above), benzyl alcohol, benzaldehyde, ethyl benzoate, heliotropin, phenylacetate, vanillin.\(^{(4, 5)}\)

Other constituents
Coumarin (trace),\(^{(1)}\) mucilage, carbohydrates and ascorbic acid (vitamin C).
Food Use

Meadowsweet is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that meadowsweet can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\(^{(G16)}\) In the USA, meadowsweet is listed by the Food and Drugs Administration (FDA) as a Herb of Undefined Safety.\(^{(G22)}\)
Herbal Use

Meadowsweet is stated to possess stomachic, mild urinary antiseptic, antirheumatic, astringent and antacid properties. Traditionally, it has been used for atonic dyspepsia with heartburn and hyperacidity, acute catarrhal cystitis, rheumatic muscle and joint pains, diarrhoea in children, and specifically for the prophylaxis and treatment of peptic ulcer. (G2 G6 G7 G8 G64)
Dosage

**Dried herb**
4–6 g or by infusion three times daily.\(^{(G6 G7)}\)

**Liquid extract**
1.5–6.0 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G6 G7)}\)

**Tincture**
2–4 mL (1 : 5 in 45% alcohol) three times daily.\(^{(G6 G7)}\)
Pharmacological Actions

*In vitro and animal studies*

Lowering of motor activity and rectal temperature, myorelaxation and potentiation of narcotic action have been documented for meadowsweet.\(^5\) In addition, flower extracts have been reported to prolong life expectancy of mice, lower vascular permeability and prevent the development of stomach ulcers in rats and mice.\(^{5,10,11}\) However, meadowsweet has also been reported to potentiate the ulcerogenic properties of histamine in the guinea–pig.\(^{10}\) The anti–ulcer action documented for meadowsweet is associated with the aqueous extract and greatest activity has been observed with the flowers.\(^{9,11}\) Meadowsweet has been reported to increase bronchial tone in the cat\(^9\) and to potentiate the bronchospastic properties of histamine in the guinea–pig.\(^9\) *In vitro*, meadowsweet has been reported to increase intestinal tone in the guinea–pig and uterine tone in the rabbit.\(^9\)

Bacteriostatic activity against *Staphylococcus aureus*, *Staphylococcus epidermidis*, *Escherichia coli*, *Proteus vulgaris* and *Pseudomonas aeruginosa* has been documented for flower extracts.\(^{12}\)

Tannins are generally considered to possess astringent properties and have been reported as constituents of meadowsweet. Meadowsweet is stated to promote uric acid excretion.\(^{G42}\)
Side-effects, Toxicity

None documented.

**Contra-indications, Warnings**
Salicylate constituents have been documented and therefore the usual precautions recommended for salicylates are relevant for meadowsweet (see Willow). Meadowsweet is stated to be used for the treatment of diarrhoea in children but in view of the salicylate constituents, this is not advisable.

Bronchospastic activity has been documented and meadowsweet should therefore be used with caution by asthmatics.

Aqueous extracts have been reported to contain high tannin concentrations and excessive consumption should therefore be avoided.

**Pregnancy and lactation**
*In vitro* utero–activity has been documented for meadowsweet. In view of the salicylate constituents and the lack of toxicity data, the use of meadowsweet during pregnancy and lactation should be avoided.
Pharmaceutical Comment

The chemistry of meadowsweet is characterised by a number of phenolic constituents including flavonoids, salicylates and tannins. Documented scientific evidence justifies some of the antiseptic, antirheumatic and astringent actions, although no human data were available. No documented toxicity data were located for meadowsweet and in view of this, excessive use should be avoided.
References


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Species (Family)

*Melissa officinalis* L. (Labiatae)
Synonym(s)
Balm, Honeyplant, Lemon Balm, Sweet Balm
Part(s) Used
Dried leaves and flowering tops
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)

BP 2002\(^{(G71)}\)

Complete German Commission E (Lemon Balm)\(^{(G3)}\)

ESCOP 1996\(^{(G52)}\)

Martindale 33rd edition\(^{(G67)}\)

Mills and Bone (Lemon Balm)\(^{(G50)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)

Ph Eur 2004\(^{(G72)}\)
Legal Category (Licensed Products)

GSL$^{(G37)}$
Constituents

See General References G2 G52 G64.

**Volatile oil**
0.06–0.375% v/m (volume in mass).\(^{(1,G52)}\) Contains at least 70 components,\(^{(G2)}\) including: *Monoterpenes* >60%. Mainly aldehydes, including citronellal, geranial, neral; also citronellol, geraniol, nerol, β-ocimene.\(^{(2,3)}\) *Sesquiterpenes* >35%. β-Caryophyllene, germacrene D.

**Flavonoids**
0.5%. Including glycosides of luteolin (e.g. luteolin 3’-O-β-D-glucuronide\(^{(4)}\)), quercetin, apigenin and kaempferol.

**Polyphenols**
Protocatechuic acid, hydroxycinnamic acid derivatives,\(^{(2)}\) caffeic acid, chlorogenic acid, rosmarinic acid,\(^{(2)}\) 2-(3’,4’-dihydroxyphenyl)-1,3-benzodioxole-5-aldehyde.\(^{(5)}\)
Food Use

Lemon balm is used to give fragrance to wine, tea and beer. Lemon balm (herb, flowers, flower tips) is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that lemon balm can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\(^{(G16)}\) In the USA, lemon balm is listed as GRAS (Generally Recognised As Safe).\(^{(G65)}\)
Herbal Use

Lemon balm has been used traditionally for its sedative, spasmylic and antibacterial properties.\(^{(G54)}\) It is also stated to be a carminative, diaphoretic and a febrifuge,\(^{(G64)}\) and has been used for headaches, gastrointestinal disorders, nervousness and rheumatism.\(^{(5)}\) Current interest is focused on its use as a sedative, and topically in herpes simplex labialis as a result of infection with herpes simplex virus type 1 (HSV-1). The German Commission E monographs state that lemon balm can be used for nervous sleeping disorders and functional gastrointestinal complaints.\(^{(G4)}\)
Dosage

**Dried herb**
1.5–4.5 g as an infusion in 150 mL water several times daily.\(^{(G4)}\)

**Topical application**
Cream containing 1% of a lyophilised aqueous extract of dried leaves of *Melissa officinalis* (70 : 1) two to four times daily.\(^{(G52)}\)
Pharmacological Actions

In vitro and animal studies

Antiviral activity
Aqueous extracts of *Melissa officinalis* have been reported to inhibit the development of several viruses.\(^{6-8,G52}\) The virucidal effect of several aqueous extracts of *M. officinalis* against HSV-1 has been demonstrated in a rabbit kidney cell line.\(^9\) However, the extracts appeared to have no activity against experimental HSV-1 infection in the eyes of rabbits.\(^9\)

Anti-human immunodeficiency virus type 1 (HIV-1) activity has been reported for an aqueous extract of *M. officinalis* in *in vitro* studies using MT-4 cells; the ED\(_{50}\) (50% effective dose for inhibition of HIV-1-induced cytopathogenicity) was found to be 16 μg/mL.\(^{10}\) Furthermore, the aqueous extract demonstrated potent inhibitory activity (ED\(_{50} = 62 \mu g/mL\) against HIV-1 replication (KK-1 strain, freshly isolated from a patient with acquired immune deficiency syndrome (AIDS). In other *in vitro* studies, an aqueous extract of *M. officinalis* inhibited giant cell formation in co–cultures of MOLT-4 cells with and without HIV-1 infection, and showed inhibitory activity against HIV-1 reverse transcriptase (ED\(_{50} = 1.6 \mu g/mL\).\(^{10}\)

Aqueous extracts of *M. officinalis* have been reported to inhibit protein biosynthesis in a cell–free system from rat liver cells, and it has been suggested that this effect may be due to caffeic acid and a component isolated from the glycoside fraction of the extract.\(^{11}\) The latter component appears to block the binding of the elongation factor EF-2 to ribosomes, thus terminating peptide elongation.\(^{11}\)

Antimicrobial activity
Antimicrobial activity of essential oil extracted from *M. officinalis* by steam distillation, determined using a micro–atmospheric technique, has been reported against the yeasts *Candida albicans* and *Saccharomyces cerevisiae*, and against *Pseudomonas putida*, *Staphylococcus aureus*, *Micrococcus luteus*, *Mycobacterium smegmatis*, *Proteus vulgaris*, *Shigella sonnei* and *Escherichia coli*.\(^{12}\)

Other activity
In studies in mice, a hydroalcoholic extract of *M. officinalis* leaves administered intraperitoneally significantly reduced behavioural activity in two tests, compared with control, suggesting that the extract has sedative effects.\(^{13}\) In both tests, the effect was maximum at 25 mg/kg. The same
extract demonstrated peripheral analgesic activity by reducing acetic acid–induced writhing and stretching in mice when administered intraperitoneally at doses of 25–1600 mg/kg 30 minutes after intraperitoneal administration of 1.2% acetic acid solution.\(^{(13)}\) However, no analgesic effects were observed on heat–induced pain (hotplate test) which suggests a lack of central analgesic activity. In other tests, low doses (3 and 6 mg/kg) of a hydroalcoholic extract of \textit{M. officinalis} leaves administered intraperitoneally induced sleep in mice given an infrahypnotic dose of pentobarbital.\(^{(13)}\) By contrast, in the same battery of tests, essential oil obtained from \textit{M. officinalis} by distillation did not demonstrate sedative or sleep–inducing effects.\(^{(13)}\)

A 30% alcoholic extract of \textit{M. officinalis} demonstrated an antispasmodic effect on rat duodenum \textit{in vitro}.\(^{(14)}\)

Aqueous methanolic extracts of the aerial parts of \textit{M. officinalis} demonstrated inhibition of lipid peroxidation \textit{in vitro} in both enzyme–dependent and enzyme–independent systems.\(^{(15)}\) The same tests carried out on the main known phenolic components of \textit{M. officinalis} revealed that rosmarinic acid, caffeic acid, luteolin and luteolin–7-O-glucoside were more potent inhibitors of enzyme–dependent lipid peroxidation than enzyme–independent lipid peroxidation.

**Clinical studies**

**Antiviral effects**

The effects of a topical preparation of a standardised aqueous extract of \textit{M. officinalis} leaves (drug/extract 70 : 1) have been investigated in herpes simplex virus (HSV) infection. In an open, multicentre study, 115 patients with HSV infection of the skin or transitional mucosa applied lemon balm leaf extract five times daily for a maximum of 14 days; complete healing of lesions was achieved after eight days of treatment in 96% of participants.\(^{(16,17)}\) Subsequently, a randomised, double–blind, placebo–controlled trial involving 116 patients with HSV infection of the skin or transitional mucosa reported statistically significant differences between the treatment (applied locally two to four times daily over 5–10 days) and placebo groups for some (including redness, physician’s assessment, patient’s assessment), but not all, outcome measures (e.g. extent of scabbing, vesication, pain).\(^{(17)}\) Another randomised, double–blind trial involved 66 patients with an acute episode of recurrent (at least four episodes per year) herpes simplex labialis compared verum cream (applied on the affected area four times daily over five days) with placebo.\(^{(18)}\) There was a significant difference in the primary outcome measure – symptom score after two days’ treatment – between the two groups \((p = 0.042)\). However, further
Sedative effects
The acute sedative effects of several plant extracts, including a preparation of *M. officinalis* leaves, were explored in a randomised, double-blind, placebo-controlled, crossover study involving 12 healthy volunteers.\(^{(19)}\) *M. officinalis* extract 1200 mg was administered orally as a single dose about 2 hours before administration of caffeine 100 mg. Melissa extract was one of the extracts tested that showed least effects on increasing tiredness (i.e. it was no different than placebo) as measured using a visual analogue scale score for alertness.

Several other studies have investigated the sedative effects of combination preparations containing extracts of lemon balm and valerian (*Valeriana officinalis*). A randomised, double-blind trial involving healthy volunteers who received Songha Night (*V. officinalis* root extract 120 mg and *M. officinalis* leaf extract 80 mg) three tablets daily taken as one dose 30 minutes before bedtime for 30 days \((n = 66)\), or placebo \((n = 32)\), found that the proportion of participants reporting an improvement in sleep quality was significantly greater for the treatment group, compared with the placebo group (33.3% versus 9.4%, respectively; \(p = 0.04\)).\(^{(20)}\) However, analysis of visual analogue scale scores revealed only a slight but statistically non-significant improvement in sleep quality in both groups over the treatment period. Another double-blind, placebo-controlled trial involving patients with insomnia who received Euvegal forte (*valerian* extract 160 mg and lemon balm extract 80 mg) two tablets daily for 2 weeks reported significant improvements in sleep quality in recipients of the herbal preparation, compared with placebo recipients.\(^{(21)}\) A placebo-controlled study involving ‘poor sleepers’ who received Euvegal forte reported significant improvements in sleep efficiency and in sleep stages 3 and 4 in the treatment group, compared with placebo recipients.\(^{(22)}\)

Other studies have investigated the sedative effects of combination preparations of extracts of lemon balm, valerian and hops (*Humulus lupulus*). In an open, uncontrolled, multicentre study, 225 individuals who were experiencing nervous agitation and/or difficulties falling asleep and achieving uninterrupted sleep were treated for two weeks with a combination preparation containing extracts of valerian root, hop grains and lemon balm leaves.\(^{(23)}\) Significant improvements in the severity and frequency of symptoms were reported, compared with the pretreatment period. Difficulties falling asleep, difficulties sleeping through the night, and nervous agitation were improved in 89%, 80% and 82% of participants, respectively.
Small-scale, short-term (two weeks' duration) studies investigating the sedative effects of oral combination preparations containing lemon balm extract indicate that these preparations are well-tolerated and do not appear to induce a 'hangover effect'. In an open, uncontrolled, multicentre study, 225 individuals who were experiencing nervous agitation and/or difficulties falling asleep and achieving uninterrupted sleep were treated for two weeks with a combination preparation containing extracts of valerian root, hop grains and lemon balm leaves.\(^{[23]}\) The tolerability of the preparation was rated as 'good' or 'very good' by 97% of physicians and 96% of patients. In a randomised, double-blind, placebo-controlled trial involving healthy volunteers who received Songha Night (\(V. \text{officinalis}\) root 120 mg and \(M. \text{officinalis}\) leaf extract 80 mg) three tablets daily for 30 days \((n = 66)\), or placebo \((n = 32)\), the proportion of volunteers reporting adverse events was similar in both groups (around 28%).\(^{[20]}\) Sleep disturbances and tiredness were the most common adverse events reported during the study. (N.B. the study was designed to assess the effects of the preparation on sleep quality.) No severe adverse events were reported. A randomised, double-blind, placebo-controlled study involving 48 adults assessed the adverse effects of 2 weeks' treatment with a combination preparation (valerian root extract 95 mg, hops extract 15 mg and lemon balm leaf extract 85 mg) taken alone or with alcohol.\(^{[24]}\) Compared with placebo, the herbal combination preparation did not have adverse effects on performance (e.g. concentration, vigilance). Furthermore, co-administration of the combination preparation with alcohol did not have potentiating effects on performance parameters.\(^{[24]}\) No serious adverse events were observed during the study.

A randomised, double-blind, placebo-controlled trial of a topical preparation containing 1% dried extract of \(M. \text{officinalis}\) leaves (drug/extract 70 : 1) involving 116 patients with HSV infection of the skin or transitional mucosa reported that there were no statistically significant differences between the treatment and placebo groups with regard to the frequency of adverse effects.\(^{[17]}\) Adverse events reported were minor (irritation, burning sensation); there were no reports of allergic contact reactions. However, skin sensitisation may occur with melissa.\(^{[G58]}\)
Contra-indications, Warnings

None documented.

**Pregnancy and lactation**
In view of the lack of toxicity data, oral administration of lemon balm during pregnancy and lactation should be avoided. Topical use of lemon balm during pregnancy and lactation is unlikely to be problematic.
Randomised clinical trials have suggested that topical lemon balm extract may have some effects on healing cutaneous lesions resulting from HSV-1 virus infection,\(^{(17,18)}\) although further rigorous studies are required to determine whether there is any effect on recurrence of infection.

In the German Commission E monograph, lemon balm is indicated for nervous disturbance of sleep and functional gastrointestinal complaints.\(^{(G4)}\) While there is some evidence from randomised controlled trials of combination preparations containing lemon balm leaf extract to support the efficacy of such products in individuals with minor sleep disorders, there has been little investigation of the effects of lemon balm extract alone on sleep quality. Further studies are required to determine the effects of preparations of lemon balm leaf extract in individuals with sleep disorders. Supporting evidence for the use of lemon balm for gastrointestinal complaints is limited to \textit{in vitro} work and requires clinical investigation.

Small-scale, short-term studies indicate that oral combination preparations containing lemon balm extract and topical preparations of lemon balm extract are well tolerated.\(^{(17,20)}\) However, there is a lack of research investigating the safety of long-term administration of lemon balm.
References

See also General References G2 G4 G9 G15 G16 G28 G36 G43 G50 G52 G54 G58 G64.


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Milk Thistle
Species (Family)

*Silybum marianum* (L.) Gaertn. (Asteraceae/Compositae)
Synonym(s)

*Carduus marianus* L., Lady’s Thistle, Marian Thistle, Mediterranean Milk Thistle, St. Mary’s Thistle.
Part(s) Used

Fruits (often referred to as ‘seeds’), herb
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)

Complete German Commission E\(^{(G3)}\)

Martindale 33rd edition\(^{(G67)}\)

Mills and Bone\(^{(G50)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)

Ph Eur 2004\(^{(G72)}\)

USP26/NF21\(^{(G73)}\)
Legal Category (Licensed Products)

Milk thistle is not included in the GSL.\(^{(G37)}\)
Constituents

See General References G2 G43.

Fruit

Flavolignans
1.5–3% silymarin, a mixture containing approximately 50% silibinin (= silybin, silybinin), silichristin and silidianin, as well as silimonin, isosilichristin, isosilibinin, silandrin, silhermin, neosili hermins A and B, 2,3-dehydrosilibinin and tri- to pentamers of silibinin (silybinomers).\(^{(1)}\)

Flavonoids
Quercetin, taxifolin and dehydrokaempferol.\(^{(1)}\)

Lipids
20–30%. Linoleic acid, oleic acid and palmitic acid.

Sterols
Cholesterol, campesterol and stigmasterol.

Other constituents
Mucilages, sugars (arabinose, rhamnose, xylose, glucose), amines and saponins.\(^{(1)}\)

Leaves

Flavonoids
Apigenin, luteolin and kaempferol and their glycosides.\(^{(1)}\)

Other constituents
β-Sitosterol and its glucoside, and a triterpene acetate.\(^{(1)}\)

Silymarin is not found in the leaves.
Food Use

Milk thistle is not used in foods.
Traditionally, milk thistle fruits have been used for disorders of the liver, spleen and gall bladder such as jaundice and gall bladder colic. Milk thistle has also been used for nursing mothers for stimulating milk production, as a bitter tonic, for haemorrhoids, for dyspeptic complaints and as a demulcent in catarrah and pleurisy. It is stated to possess hepatoprotective, antioxidant and choleretic properties.\(^1,2\)

Current interest is focused on the hepatoprotective activity of milk thistle and its use in the prophylaxis and treatment of liver damage and disease.

The leaves have also been used for the treatment of liver, spleen and gall bladder disorders and as an antimalarial, emmenagogue and for uterine complaints. Milk thistle leaf preparations are available today, although most research has been conducted with preparations of the fruit since the leaf does not contain the pharmacologically active component silymarin.
Dosage

**Fruit**
Crude drug 12–15 g daily in divided doses (equivalent to silymarin 200–400 mg daily). (G3)

**Herb**
Approximately 1.5 g of finely chopped material as a tea, two or three cups daily.

The doses of silymarin used in clinical trials have ranged from 280 to 800 mg/day (equivalent to milk thistle extract 400–1140 mg/day standardised to contain 70% silibinin). (3) For hepatic disorders, doses of up to 140 mg (equivalent to 60 mg silibinin) two or three times daily have been suggested. (G43)

In Germany, the recommended regimen for treatment of *Amanita phalloides* poisoning with a standardised silymarin preparation (Legalon) is a total dose of silibinin (as the disodium dihemisuccinate) (20 mg/kg body weight) over 24 hours, divided into four intravenous infusions each given over a 2–hour period. (G43 G55)
Pharmacological Actions

Several pharmacological activities have been documented for milk thistle fruit, including hepatoprotective, antioxidant, anti-inflammatory, antifibrotic and antitumour properties, as well as inhibition of lipid peroxidation, stimulation of protein biosynthesis and acceleration of liver regeneration. Silymarin (an isomer mixture comprising mainly silibinin, silichristin and silidianin) is the pharmacologically active component of milk thistle fruit; silibinin is the main component of silymarin. There is an extensive literature on the pharmacological effects of silymarin and silibinin, particularly with regard to their hepatoprotective activity which provides supporting evidence for the clinical uses. The pharmacology and clinical efficacy of milk thistle have been reviewed. \(^{(1-3, G50 \ G55)}\) The following represents a summary of selected publications on this subject.

There is a lack of research investigating the pharmacological effects of preparations of milk thistle leaf. \(^{(G2 \ G32 \ G35)}\)

**In vitro and animal studies**

**Antioxidant activity**

Silymarin and silibinin (silybin) are antioxidants that react with free radicals (e.g. reactive oxygen species) transforming them into more stable and less reactive compounds. \(^{(1,4-6)}\) Silymarin and silybin have been reported to inhibit lipid peroxidation induced by iron–linked systems in rat liver microsomes \(^{(7,8)}\) and protect against phenylhydrazine–induced lipid peroxidation in rat erythrocytes. \(^{(1)}\) Furthermore, in rats, intraperitoneal silymarin has been shown to increase total glutathione in the liver, intestine and stomach and to improve the reduced glutathione to oxidised glutathione ratio. \(^{(9)}\) Silymarin has been shown to inhibit copper–induced oxidation of human low-density lipoprotein (LDL) \textit{in vitro} in a concentration–dependent manner. \(^{(10)}\) Silybin appears to be the constituent of silymarin responsible for the LDL antioxidant effect. In contrast, silichristin and silydianin appeared to act as pro–oxidants, but without significantly reducing the total LDL antioxidant capacity of silymarin.

Free radicals are recognised as having an important role in several pathological processes, including inflammation, necrosis, fibrosis, atherosclerosis, carcinogenesis and ageing and in the hepatotoxic mechanisms of various substances. The antioxidant activity of silymarin is thought to contribute to its hepatoprotective properties. \(^{(11, G55)}\)

**Hepatoprotective properties**
*In vitro* studies using isolated hepatocytes have documented the protective activity of silymarin and several of its components against cell damage induced by various cytotoxic substances.\(^{(1)}\)

*In vivo* studies in rats and mice have demonstrated the hepatoprotective activity of silymarin and silybin in acute liver toxicity induced by various toxic agents with different mechanisms of action, including carbon tetrachloride, galactosamine, thioacetamide, ethanol, paracetamol (acetaminophen), thallium, phalloidin and α-amanitin (the main toxic constituents of the mushroom *A. phalloides*).\(^{(1)}\) Experimental studies in chronic liver toxicity induced by repeated administration of carbon tetrachloride, heavy metals, thioacetamide and several drugs, including azathioprine and indometacin, have also demonstrated that administration of silymarin and silybin protects against damage.\(^{(1)}\) Other studies have reported protective effects of silymarin against liver injury induced by ischaemia\(^{(12)}\) and gamma irradiation.\(^{(13)}\)

Studies in rabbits fed a high–fat diet for 12 weeks have shown that histopathological alterations were least advanced in animals which also received a silymarin–phospholipid complex.\(^{(14)}\) In rats, silymarin inhibited the development of diet–induced hypercholesterolaemia.\(^{(15)}\) The hypocholesterolaemic effects of silymarin may be due to the effects of silymarin on lipoprotein metabolism.\(^{(16)}\)

The effects of silymarin on biliary bile salt secretion have been seen in studies in rats.\(^{(17)}\) Intraperitoneal silymarin (25, 50, 100 and 150 mg/kg/day) for five days induced a dose–dependent increase in bile flow and bile salt secretion. Stimulation of bile salt secretion was mainly accounted for by an increase in the biliary secretion of the hepatoprotective bile salts β-muricholate and ursodeoxycholate.

**Nephroprotective properties**

Silibinin injected into rats prior to administration of cisplatin afforded protection of glomerular and proximal tubular function.\(^{(18,19)}\) Silibinin does not affect the cytotoxic activity of cisplatin.\(^{(19)}\) Intraperitoneal silibinin (5 mg/kg) administered to rats 30 minutes before ciclosporin decreased ciclosporin–induced lipid peroxidation but produced no protective effect on the glomerular filtration rate.\(^{(20)}\)

**Anticancer activity**

Silybin at concentrations of 0.1–20 μmol/L inhibited the growth of drug–resistant ovarian cancer cells and doxorubicin–resistant breast cancer cells *in vitro*.\(^{(21)}\) Furthermore, silybin in the range of 0.1–1.0 μmol/L potentiated the effect of cisplatin and doxorubicin in experimental tumour cell lines. When
applied to the skin of SENCAR mice, silymarin gave protection against the effects of the tumour promoters 12-O-tetradecanoylphorbol (TPA) and okaidic acid (OA).\(^{(22)}\) Topical application of silymarin prior to that of TPA and OA completely inhibited induction of tumour necrosis factor α (TNFα) mRNA expression in the epidermis. Substantial protection from photocarcinogenesis in mice treated with phorbol ester or 7,12-dimethylbenz(a)anthracene has been demonstrated.\(^{(23)}\) The antitumour effect is primarily at stage 1 tumour promotion and silymarin acts by inhibiting cyclo oxygenase 2 (COX-2) and interleukin 1α (IL-1α).\(^{(24)}\) Such effects may involve inhibition of promoter-induced oedema, hyperplasia, the proliferation index and oxidant state.\(^{(25)}\)

Treatment of serum–starved human prostate carcinoma DU145 cells with silymarin resulted in significant inhibition of transforming growth factor α (TGFα)-mediated activation of the epidermal growth factor receptor erbB1.\(^{(26)}\) There was also a decrease in tyrosine phosphorylation of an immediate downstream target, the adaptor protein SHC, together with a decrease in binding to erbB1. In the silymarin–treated cell lines there was a significant induction of the cyclin–dependent kinase inhibitors (CDKIs) Cip1/p21 and Kip/p27 concomitant with a significant decrease in CDK4 expression, but no changes in the levels of CDK2 and CDK6 and their associated cyclins E and D1, respectively. Additional experiments showed that there was a significant inhibition of constitutive tyrosine phosphorylation of both erbB1 and SHC, but no changes in their protein levels. The results indicated that silymarin may exert a strong anticarcinogenic effect against prostate cancer and that this effect is likely to involve impairment of the erbB1–SHC-mediated signalling pathway, induction of CDKIs and resultant G\(_1\) arrest.\(^{(26)}\)

There was a significant inhibition of mitogen–activated protein kinase (MAPK) ERK activity at lower doses in epidermal A431 cells treated with silymarin, whereas higher doses activated MAPK/JNK1.\(^{(27)}\) Silymarin exerted a strong anticarcinogenic effect against human breast carcinoma cells MDA-MB468 with G\(_1\) arrest in cell cycle progression and also induction in protein expression of the CDKI Cip/p21.\(^{(28)}\)

The cancer chemoprevention and anticarcinogenic effects of silymarin have been shown to be due to its major constituent silibinin.\(^{(29)}\) Silibinin decreases prostate–specific antigen (PSA) in hormone–refractory human prostate carcinoma LNCaP cells and inhibits cell growth via G\(_1\) arrest.\(^{(30)}\)

Silibinin was fed orally to SENCAR mice and its tissue distribution investigated.\(^{(31)}\) Free silibinin mainly accumulated in the liver, although it was also distributed in other organs. Increases in glutathione-S-transferase and quinone reductase activities in the liver, lung, stomach, skin and small
bowl were observed. The results demonstrated the bioavailability of and phase II enzyme induction of silibinin in different tissues where silymarin has been shown to be a strong cancer chemopreventive agent.

**Anti-inflammatory activity**
Silymarin administered orally reduced foot-pad abscesses in a dose-dependent manner in the carrageenan rat paw oedema test ($ED_{50} = 62.4$ mg/kg).\(^{(32)}\) In the xylene–induced inflammation test, topically applied silymarin was comparable with indomethacin.\(^{(32)}\) Silymarin given intraperitoneally to mice resulted in inhibition of leukocyte accumulation in inflammatory exudates and reduced the neutrophil count.\(^{(32)}\) Activation of NF-κB induced by TNF, phorbol ester, okaidic acid and ceramide was blocked by silymarin in a dose–dependent manner.\(^{(33)}\) Silymarin also inhibited TNF-induced activation of mitogen–activated protein kinase and c-Jun N-terminal kinase.\(^{(33)}\) The inhibition of activation of NF-κB and the kinases may provide part of the molecular basis for the anti-inflammatory and anticarcinogenic effects of silymarin.\(^{(33)}\) Silymarin potently suppressed both NF-κB–DNA binding activity and its dependent gene expression induced by okaidic acid in the hepatoma cell line Hep G2.\(^{(34)}\) In addition, silymarin inhibits COX-2 and IL-1α.\(^{(24)}\)

**Gastric ulcer protective effects**
Oral administration of silymarin to rats prevented gastric ulceration induced by cold–restraint stress.\(^{(35)}\) Gastric secretion volume and acidity were not affected, but histamine concentration was significantly decreased. It was suggested that the anti-ulcerogenic effect of silymarin may be related to inhibition of enzymic peroxidation by the lipoxygenase pathway.\(^{(35)}\) The protective effect of silymarin on gastric injury induced in rats by ischaemia–reperfusion and its effects on mucosal myeloperoxidase has been compared with that of allopurinol.\(^{(36)}\) The mean ulcer indexes (4.75, 4.50 and 3.63 ui, respectively) of rats treated with 25, 50 and 100 mg/kg silymarin were significantly lower than in control rats, although allopurinol was considerably more potent (2.3 ui; 100 mg/kg).\(^{(36)}\)

**Other effects**
Silymarin has been shown to prevent alloxan–induced diabetes mellitus in rats, possibly due to its antioxidant activity and increases in plasma and pancreatic glutathione concentrations.\(^{(37)}\)

It has been reported that *Silybum marianum* and silymarin beneficially affect skin elasticity. A phospholipid–silymarin complex (Silymarin–Phytosome) evaluated for its topical effects against croton oil dermatitis in mice and UV-

### Clinical studies

Clinical trials with milk thistle preparations have focused on their use in alcoholic liver disease, cirrhosis and acute viral and chronic hepatitis. However, several trials have included patients with liver disease of different aetiology, e.g. alcoholic and non-alcoholic cirrhosis. There is also interest in the use of silymarin in toxin- and drug–induced hepatitis, for example following ingestion of the death cap mushroom \textit{A. phalloides}.

A randomised, double–blind, placebo–controlled, multicentre trial involving 200 alcoholic patients with histologically or laparoscopically proven liver cirrhosis investigated the effects of administration of silymarin (150 mg) three times daily.\(^{(41)}\) The results indicated that silymarin had no effect on survival and the clinical course in these patients: 125 patients (silymarin \(n = 57\) and placebo \(n = 68\)) completed the two–year study period, during which 29 died (\(n = 15\) and 14 for silymarin and placebo, respectively; no statistically significant difference).

In a randomised, double–blind, placebo–controlled trial, patients with alcoholic (\(n = 91\)) or non–alcoholic (\(n = 79\)) cirrhosis received silymarin (140 mg) three times daily or placebo for two years.\(^{(42)}\) The four–year survival rate was significantly higher in silymarin–treated patients than in placebo recipients (58 versus 39\%, respectively, \(p = 0.036\)). Subgroup analysis indicated that the effect of silymarin on mortality was more pronounced in those patients with alcoholic cirrhosis.

Another randomised, double–blind, placebo–controlled trial carried out over four years reported a significantly higher survival rate in patients with alcoholic cirrhosis treated with silymarin (420 mg) daily compared with placebo recipients, although the effect in patients with non–alcoholic cirrhosis was less marked.\(^{(43)}\)

Other controlled trials have investigated the effects of silymarin in patients with alcohol–related liver damage. Several of these,\(^{(44,45)}\) but not all,\(^{(46)}\) reported statistically significant benefits with silymarin, e.g. on serum transaminases, compared with placebo.\(^{(1)}\)
In a randomised controlled trial, 60 patients with diabetes caused by alcoholic liver cirrhosis received silymarin (600 mg/day) or no silymarin treatment for six months. At the end of the study period, the mean values for fasting blood glucose, daily blood glucose, daily glycosuria, glycosylated haemoglobin, daily insulin requirement, malondialdehyde and glucagon–stimulated C peptide were significantly lower in silymarin–treated patients than in those who did not receive silymarin treatment.

A pilot study involving 20 patients with chronic active hepatitis randomised to receive a silybin–phosphatidylcholine complex preparation (IdB1016; Silipide) (240 mg) twice daily or placebo for seven days reported significant reductions in the mean serum concentrations of aspartate aminotransferase (AST), alanine aminotransferase (ALT), γ-glutamyltranspeptidase (GGT) and total bilirubin in silybin complex–treated patients compared with values in placebo recipients. The same preparation has been reported to reduce serum concentrations of liver enzymes (AST and ALT) in 65 patients with chronic persistent hepatitis in a randomised placebo–controlled trial.

The hepatoprotective effects of silymarin in 222 de novo tacrine–treated patients with mild to moderate dementia of the Alzheimer type were investigated in a randomised, double–blind (for silymarin), placebo–controlled, multicentre, 12–week trial. Patients received tacrine plus silymarin (420 mg/day) (n = 110) or tacrine plus placebo (n = 112); silymarin (and placebo) were initiated one week before tacrine (40 mg/day for six weeks then 80 mg/day for six weeks). An intention–to–treat analysis indicated that there was no difference in serum ALT concentrations between the two groups, but that silymarin–treated patients experienced significantly fewer gastrointestinal and cholinergic side–effects without any impact on cognitive status than did placebo recipients.

The effects of silymarin in preventing psychotropic drug–induced hepatic damage have been investigated in a randomised, double–blind, placebo–controlled trial. Sixty women aged 40–60 years who had been taking phenothiazines or butyrophenones for at least five years and who had AST and ALT activity twice normal values were randomised to continued treatment with psychotropic agents or suspension of treatment and to silymarin (800 mg/day) or placebo for 90 days. The findings indicated that treatment with silymarin reduced the lipoperoxidative hepatic damage associated with prolonged administration of butyrophenones and phenothiazines and that the protective effect was greater when treatment with these psychotropic agents was suspended for three months.

There have been numerous case reports, many of which report favourable
outcomes, on the therapeutic use of silymarin and silibinin, usually given in combination with standard treatment, in poisoning caused by ingestion of the death cap mushroom *A. phalloides*, although there are no controlled trials in this indication.\(^\text{(1,51–54,G55)}\) Silibinin is usually given intravenously and case reports have indicated that early administration appears to be important.\(^\text{(55,56)}\)

**Pharmacokinetics**

Studies of the pharmacokinetics of silymarin and its components and of a silibinin–phosphatidylcholine complex preparation (IdB 1016; Silipide) in both healthy volunteers and patients with cirrhosis and those who have undergone cholecystectomy have been reviewed.\(^\text{(1,G50 G55)}\) Approximately 20–50% of silymarin is absorbed following oral administration and approximately 80% of the dose, whether administered orally or intravenously, is excreted in the bile.\(^\text{(57)}\) Studies in healthy volunteers have reported an elimination half-life of approximately 6 hours following administration of single doses of silymarin corresponding to approximately 240 mg silibinin.\(^\text{(58,59)}\) Other studies have compared the pharmacokinetics of different silymarin preparations and shown statistically significant differences in bioavailability.\(^\text{(60,61)}\)

The bioavailability of a silybin–phosphatidylcholine complex preparation (IdB 1016) has been shown to be several times greater than that of silymarin in single-dose studies involving healthy volunteers\(^\text{(62)}\) and patients with hepatic cirrhosis.\(^\text{(63)}\)
Side-effects, Toxicity

No adverse events were noted in a pharmacokinetic study involving healthy male volunteers following single oral doses of silymarin corresponding to up to 254 mg silibinin.\(^{(59)}\)

Clinical trials involving patients with liver disorders of various origin and who received oral silymarin at doses of up to 600–800 mg/day for up to six months have reported that no adverse effects were observed.\(^{(42,47,51)}\)

Data from drug monitoring studies involving more than 3500 patients, including one study involving 2637 patients with various types of chronic liver disease treated with silymarin (Legalon) (560 mg/day) for eight weeks, have indicated that the frequency of adverse effects with silymarin is approximately 1%. Adverse effects are mainly transient, non-serious, gastrointestinal complaints.\(^{(4,64,G55)}\) It is stated that silymarin may occasionally produce a mild laxative effect.\(^{(G3)}\)

A case report from Australia described a reaction associated with a preparation of milk thistle. The symptoms included episodes of severe sweating, abdominal cramping, nausea, vomiting, diarrhoea and weakness and these were verified by rechallenge.\(^{(65)}\) Another report described a case of anaphylactic shock in a 54-year-old man with immediate-type allergy to kiwi fruit.\(^{(66)}\) He experienced facial oedema, swelling of the oral mucosa, bronchospasm, respiratory distress and decreased blood pressure after taking a preparation of milk thistle; a skin-prick test of an extract of milk thistle fruit elicited an immediate-type reaction.

The acute toxicity of oral and intravenous silymarin and silibinin has been investigated in various animal species (mice, rats, rabbits and dogs).\(^{(1)}\) Oral silymarin administered to mice and dogs at doses of 20 and 1 g/kg, respectively, did not cause adverse effects or mortality. Long-term oral administration of silymarin (100 mg/kg/day) to rats for 16 or 22 weeks did not reveal any adverse effects.
Contra–indications, Warnings

None documented. In view of the lack of long–term safety data, excessive use of milk thistle should be avoided (except where its use may help to prevent toxicity caused by other substances).

Milk thistle is contra–indicated for individuals with hypersensitivity to species of Asteraceae.

Pregnancy and lactation

In view of the lack of toxicity data, use of milk thistle preparations during pregnancy and lactation should be avoided unless the expected benefit is thought to outweigh any unknown risks to the fetus.
Pharmaceutical Comment

The chemistry of milk thistle is well-documented and there is good evidence that silymarin and its components, particularly silibinin, are responsible for the pharmacological effects.

Documented scientific evidence from *in vitro* and animal studies provides supportive evidence for some of the uses of milk thistle, particularly those relating to hepatoprotective properties.

There have been several controlled clinical trials investigating the effects of milk thistle in a range of liver disorders, including acute viral hepatitis, chronic hepatitis, alcoholic liver disease, cirrhosis and toxic liver damage. The results of these studies are not entirely consistent or conclusive. In addition, some trials have methodological shortcomings, for example the inclusion of patients with different liver disorders, small numbers of patients and failure to control or monitor alcohol intake.\(^{(67)}\) Further, well-designed, clinical trials in clearly defined patient groups are required in order to establish the efficacy of milk thistle and its components in different liver disorders. In Germany, milk thistle is approved for the treatment of toxic liver disorders and as a supportive treatment in chronic inflammatory liver disease and hepatic cirrhosis.\(^{(G3)}\)

Teas prepared from milk thistle fruits or herb are not commonly used as only a small proportion of silymarin gets into the aqueous extract such that pharmacologically active doses are not attained.\(^{(68)}\) For this reason, in Germany teas are recommended only as supportive treatment in functional gall bladder disorders and not for antihepatotoxic effects.\(^{(G2)}\) In Germany, milk thistle fruit (3–5 g) as an infusion three or four times daily is also indicated for mild digestive disorders.\(^{(G2)}\)

There are some toxicity and safety data for milk thistle which, together with data on the adverse effects reported in clinical trials, provide good evidence for the safety of milk thistle when used at recommended doses in the short term. However, further data on the long-term safety of milk thistle use are required.

Patients wishing to use milk thistle should be advised to consult a pharmacist, doctor or other suitably trained healthcare professional for advice.
References

See General References G2 G3 G9 G34 G36 G43 G50 G64.


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2727. (PubMed)


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Mistletoe
Species (Family)

*Viscum album* L. (Loranthaceae)
Synonym(s)

Viscum
Part(s) Used

Leaf, fruit (berry), twig
Pharmacopoeial and Other Monographs

BHMA 2003\(^{(G66)}\)

BHP 1996\(^{(G9)}\)

Complete German Commission E\(^{(G3)}\)

Martindale 33rd edition\(^{(G43)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

Mistletoe is not included in the GSL. (G37)
Constituents
See references 1, 2, and General reference G2

Acids
Fatty acids (C_{12}–C_{22}), 80% oleic and palmitic, myrsitic;\(^{(3)}\) phenolic acids, e.g. caffeic, p-coumaric, gentisic, p-hydroxybenzoic, p-hydroxyphenylacetic, protocatechuic, vanillic;\(^{(3,4)}\) anisic, quinic and shikimic.\(^{(3,4)}\)

Alkaloids
It has been suggested that alkaloids can be passed on from hosts to parasitic plants such as mistletoe (e.g. nicotine alkaloids have been isolated from mistletoe growing on Solanaceae shrubs).\(^{(5)}\)

Amines
Acetylcholine, choline, β-phenylethylamine, histamine, propionylcholine and tyramine.\(^{(6)}\)

Flavonoids
Chalcones, e.g. 2′-hydroxy-4′, 6′-dimethoxy-chalcone-4-O-glucoside, 2′-hydroxy-3, 4′, 6′-trimethoxychalcone-4-O-glucoside, 2′-hydroxy-4′, 6′-dimethoxychalcone-4-O-(apiosyl-1→2)-glucoside.\(^{(1,7)}\)

Flavanones, e.g. (2R)-5,7-dimethoxyflavanone-4′-O-glucoside, (2S)-3′, 5, 7-trimethoxyflavanone-4′-O-glucoside, homoeriodictyol (4′, 5, 7-trihydroxy-5′-methoxyflavanone.\(^{(1,7)}\)

Flavones, e.g. quercetin, isorhamnetin, sakuranetin, rhamnazin (3,4′,5-trihydroxy-3′, 7-dimethoxyflavone; quercetin 3′, 7-dimethyl ether) and its glycoside rhamnazin-3, 4′-di-O-glucoside and other quercetin methyl ethers.\(^{(1,7)}\)

Lectins
Lectins are heterodimeric glycoproteins (mol. wt between 55 kDa and 63 kDa),\(^{(1)}\) which belong to a group of type 2 ribosome-inactivating proteins (RIPs) and are structurally similar to ricin. They are composed of two distinct subunits, one B-subunit (B-chain, the galactose-binding site; 34 kDa) and one toxophoric A-subunit (A-chain, the cytotoxic-binding site, RNA-N-glycosidase; 29 kDa). The structure of the B-chain of mistletoe lectins is based on 264 amino acids. Seven cysteine residues and three N-linked carbohydrate chains are included. Mistletoe lectins (V. album agglutinin-1, viscumin (mol. wt 60 000)) are galactoside-specific plant lectins (63 kDa). Three different
Mistletoe lectins have been isolated: ML-1 (main component; broad range of affinity for α/β-linked galactopyranosyl residues), ML-2 (affinity for d-galactose and N-acetyl-d-galactosamine) and ML-3 (affinity for N-acetyl-d-galactosamine). The chitin-binding lectin (chitin-binding-agglutinin, visalb-CBA or visalb-CBL), which is distinct from ML-1, ML-2 and ML-3, is a homodimer lectin with two identical subunits (10.8 kDa). Its amino acid composition is similar to that of other chitin-binding hololectins.\(^{(1)}\) Reported yields of ML-1, ML-2 and ML-3 from leaves and stems are 3–170 mg/100 g dry weight, 0.1–35 mg/100 g dry weight, and 0–67 mg/100 g dry weight, respectively.\(^{(1)}\)

ML-1, ML-2 and ML-3 have been isolated from Iscador Qu Special and Iscador M Special. Iscador P contains almost no mistletoe lectins (R Dierdorf, personal communication, September 2003).\(^{(8)}\)

**Terpenoids**
An acyclic monoterpene glucoside, 2,6-dimethylocta-2,7-diene-1,6-diol-6-O-[6′-β-d-apiofuranosyl]-β-d-glucopyranoside, has been isolated from leaves and stems.\(^{(9)}\) Phytosterols, including β-sitosterol, stigmasterol and glycosides, together with pentacyclic triterpenes β-amyrin and its acetate, betulinic acid, oleandrin and oleanolic acid are present.\(^{(2,10)}\)

**Viscotoxins**
Viscotoxins (mol. wt 5 kDa) have been isolated from leaf homogenates. Viscotoxins are amphipathic, strongly basic polypeptides that are highly enriched with cysteine residues; they belong to the family of α- and β-thionins.\(^{(1)}\) Viscotoxins consist of 46 amino acid residues with three disulphide bonds, which stabilise the conformation of the molecule. Several different viscotoxins have been isolated, the main ones being viscotoxins A2, A3 and B.

**Other constituents**

**Lignans** Syringaresinol 4,4′-O-diglucoside (eleutheroside E).\(^{(1)}\)

**Phenylpropanoids** Syringenin-4′-O-glucoside (syringin), syringenin-4′-O-apiosyl-1→2-glucoside (syringoside).\(^{(1)}\)

**Polyalcohols** 1-d-1-O-Methyl-muco-inositol (a derivative of O-methyl-inositol),\(^{(G2)}\) mannitol, pinitol, quebrachitol and viscumitol.\(^{(1)}\)

**Polysaccharides** Mainly a methylester of 1→4α-galacturonic acid in the leaves and rhamnogalacturanrs in the berries. Pectin (mol. wt 42 kDa), arabinogalactan (mol. wt 110 kDa) are present in leaves, and a
rhamnogalacturan (mol. wt 700 kDa) is found in the berries. The berries contain acidic (mol. wt 1340 kDa) and neutral (mol. wt 30 kDa) polysaccharides. The acidic polysaccharides interact with ML-1. Simple sugars and tannins are also present.\(^{(1)}\)
Food Use

Mistletoe is not generally used as a food. The branches and berries of mistletoe are listed by the Council of Europe as natural sources of food flavouring (category N3).\(^{(G16)}\) This category indicates that mistletoe may be added to foodstuffs in the traditionally accepted manner, although there is insufficient information available for an adequate assessment of potential toxicity.
Herbal Use

Mistletoe is stated to possess hypotensive, cardiac-depressant and sedative properties. Traditionally, it has been used for high blood pressure, arteriosclerosis, nervous tachycardia, hypertensive headache, chorea and hysteria. (G2, G7, G66, G69)

Modern use of mistletoe preparations is focused on use as a treatment and as an adjuvant treatment in cancer. Clinical studies of mistletoe preparations have assessed mistletoe preparations as a treatment, or as an adjunctive treatment, in patients with different types of cancers. A small number of other clinical trials have been conducted involving patients with chronic hepatitis C infection, human immunodeficiency virus (HIV) infection and respiratory infections. (see Clinical studies)
Dosage

Dosages for oral administration (adults) recommended in older standard herbal reference texts for the traditional uses are given below.

**Dried leaves**
2–6 g or by infusion three times daily.\(^{(G7)}\)

**Liquid extract**
1–3 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)

**Tincture**
0.5 mL (1 : 5 in 45% alcohol) three times daily.\(^{(G7)}\)

**Infusion**
40–120 mL (1 : 20 in cold water) daily.\(^{(G7)}\)

**Soft extract**
0.3–0.6 mL (1 : 8 infusion or tincture) three times daily.\(^{(G7)}\)

The administration of mistletoe preparations as adjunctive treatments in cancer is mostly by subcutaneous injection, and the manufacturer’s recommendations for the specific mistletoe preparations should be followed. Dosages used in clinical trials assessing different mistletoe preparations, therefore, vary widely and are often complex; dosage regimens also vary depending on the type of cancer being treated. Trials involving ML-1 standardised mistletoe preparations usually describe the dose of mistletoe extract in terms of the equivalent amount of ML-1. For example, in a trial involving patients with head and neck squamous cell carcinoma, an ML-1 standardised mistletoe extract (Eurixor) equivalent to ML-1 1 ng/kg body weight twice weekly was given by subcutaneous injection over a 60-week period in treatment cycles of 12 weeks followed by a break in treatment after every four weeks.\(^{(11)}\)
Pharmacological Actions

There is a vast scientific literature relating to the biochemical and pharmacological effects of mistletoe and its constituents, particularly the lectin and viscotoxin constituents, which are believed to be the active principles.(12) There is also a substantial literature on clinical investigations with mistletoe extracts and purified components, and a recent scientific meeting focused solely on research into mistletoe in cancer therapy.(13)

In vitro and animal studies

Preclinical research has focused on investigating the cytotoxic and immunomodulatory activities of mistletoe and its constituents, although other effects, such as agglutinating and hypotensive properties have also been documented. These activities have been summarised in several comprehensive reviews.(5,12,14–18) A brief summary of some of the scientific literature on the properties of mistletoe and its constituents is given below. Evaluation, interpretation and comparison of preclinical studies is not straightforward as studies have investigated the effects of different types of mistletoe extracts and commercial products, some of which are not characterised or their composition is not adequately described in reports of studies; a number of other studies have used purified mistletoe lectins and/or viscotoxins.

Cytotoxic and anticancer activities

The lectin and, to a lesser extent, viscotoxin constituents of mistletoe are the main cytotoxic components.(12) Cytotoxic activity in vitro has been documented for mistletoe extracts and for purified mistletoe lectins and viscotoxins. In one study, aqueous extracts (ABNOBAviscum) of mistletoe sourced from two host trees and purified mistletoe lectins II and III inhibited cell growth for several tumour cell lines, including murine B cell hybridomas, P815 (murine mastocytoma), EL-4 (murine thymoma), MOLT-4 (human T cell acute lymphoblastic leukaemia) and U937 (human histiocytoma).(19) For example, the mistletoe extract sourced from apple trees at a dilution of 1 : 10 (corresponding to aqueous extract 0.15 mg/mL) resulted in 98% inhibition of the growth of the MOLT-4 tumour cell line. The effects of another mistletoe extract (Isorel M), also obtained from mistletoe grown on apple trees, on normal and tumour cell growth were compared with those of its low and high molecular weight components and mistletoe lectins in vitro using melanoma B16 cell cultures. At the highest concentration tested (2 μg/mL), all preparations significantly inhibited [³H]thymidine incorporation into melanoma cells, compared with control.(20) However, separately, the low and high molecular weight components achieved less inhibition than the total
mistletoe extract, suggesting that the constituents of both fractions are needed for optimal activity.

In another *in vitro* study, mistletoe lectin I and purified viscotoxins (ratio of viscotoxins A2 : A3 : B was 1:1.6:0.6) showed cytotoxic activity against MOLT-4 cells as well as Yoshida sarcoma cells and K562 human myeloid cells in a concentration-dependent manner.\(^{(21)}\) Mistletoe lectins I, II and III showed cytotoxic activity against sensitive and resistant human colon cancer cell lines in a concentration-dependent manner *in vitro*.\(^{(22)}\)

The mechanism of growth arrest demonstrated with mistletoe extracts in *in vitro* studies using human and murine tumour cell lines was shown to be due to the induction by mistletoe constituents of programmed cell death (apoptosis).\(^{(19)}\) Induction of apoptosis by mistletoe extracts (various Iscador preparations) has also been demonstrated in endothelial cells.\(^{(23)}\) The apoptosis-inducing properties and hence cytotoxicity of *V. album* extracts depend on the content of active compounds, mainly the mistletoe lectins, the content of which in turn is dependent on factors such as the host tree and the manufacturing process.\(^{(24,25)}\) Furthermore, other compounds present in mistletoe extract may modulate the cytotoxic activity.

It has recently been reported following *in vitro* studies in hepatocarcinoma cells that mistletoe lectin II from Korean mistletoe induces apoptosis by inhibiting telomerase, an enzyme that maintains protein–nucleic acid complexes (telomeres) at the ends of chromosomes.\(^{(26)}\) This effect appeared to be mediated via a pathway independent of p53.

Antitumour activity in animal models has been demonstrated for several different commercial preparations of mistletoe extract, and for mistletoe lectins. In one study, mice bearing a hind-limb melanoma B16F10 tumour were given a single dose (100 mg/kg body weight, intraperitoneally) of a mistletoe extract (Isorel M) obtained from mistletoe growing on apple trees. Compared with a control group (which received a saline injection), mice treated with mistletoe showed a statistically significant reduction in tumour volume measured two days after mistletoe administration.\(^{(27)}\) Inhibition of tumour growth was documented for an aqueous mistletoe extract (Lektinol, containing mistletoe lectins 405 μg/mL), administered intraperitoneally or subcutaneously at doses equivalent to mistletoe lectins 30 and 300 ng/mL/kg per day for up to four weeks, in mice with Renca (renal cell carcinoma), C8 colon 38 carcinoma and F9 testicular carcinoma, when compared with control.\(^{(28)}\) No such inhibitory effect was observed in mice with Lewis lung carcinoma and B16 melanoma under the same conditions.
Anticarcinogenic activities have been described for a bacterially fermented mistletoe extract (Iscador M; mistletoe grown on apple trees).\(^{(29)}\) Methylcholanthrene-induced sarcoma formation was inhibited in mice treated with Iscador M (1.66 mg twice weekly for 15 weeks, intraperitoneally), compared with a control group. In separate experiments, mistletoe was reported to markedly inhibit the development of lung metastases when co-administered with metastatic B16F10 melanoma cells through the lateral tail vein;\(^{(29)}\) however, the results from this experiment were not subjected to statistical testing. Antimetastatic activity has also been documented for other commercial mistletoe preparations (e.g. Lektinol, Helixor) standardised for mistletoe lectin content in other in vivo studies (mice).\(^{(30–32)}\)

**Immunomodulatory activity**

Certain types of immune cells, such as natural killer (NK) cells, can lyse tumour cells; this can occur with or without activation by cytokines.\(^{(12)}\) Extracts of mistletoe have been shown to enhance the cytolytic activity of NK cells in studies using human peripheral blood mononuclear cells and a human cancer cell line (K562 leukaemia).\(^{(33)}\) Several other studies have described the enhancing effects of mistletoe extracts and their isolated components on natural killer activity of peripheral blood mononuclear cells in vitro and in vivo.\(^{(17)}\)

Components of mistletoe have also been reported to induce the proliferation of human peripheral blood mononuclear cells in vitro.\(^{(9,34)}\) In one study, of six mistletoe extracts tested, an extract (Iscador P) derived from mistletoe grown on pine trees induced the strongest proliferative response on peripheral blood mononuclear cells from non-treated individuals.\(^{(9)}\) This extract is almost devoid of lectins, suggesting that a non-lectin-associated antigen from this extract is responsible for the observed effects.

Stimulation of cytokine release (e.g. interleukin 1 (IL-1), IL-2, IL-6, IL-12, tumour necrosis factor α (TNFα)) by mistletoe extracts or mistletoe lectins has been demonstrated in various experimental studies in vitro.\(^{(12,35–37)}\)

Immunostimulant activity in mice, demonstrated for example by an increase in the phagocytic activity of peritoneal macrophages, has been documented for mistletoe lectin 1 and for mistletoe extracts standardised for mistletoe lectin 1 content.\(^{(12)}\) In vivo immunostimulant activity in mice (humoral and cellular), demonstrated by an enhancement of delayed hypersensitivity and antibody formation to sheep red blood cells, has been documented for the crude plant juice, Iscador and for a polysaccharide fraction isolated from the berries.\(^{(38)}\) Activity was attributed to stimulation of the monophagocytic system and to induction of inflammation. The results indicated that the
immunostimulant property of mistletoe is not solely attributable to the polysaccharides found in the berries (plant juice also active) or to the lactobacilli content of the fermented plant juice (crude extract also active). Non-specific immunological effects with mistletoe extracts are reported to be dependent on the frequency and quantity of the applied extract.\(^{(39)}\)

**Agglutinating activity**
Agglutinating activity that is preferential towards tumour cells over erythrocytes has been exhibited by Iscador and by a lectin fraction.\(^{(40–42)}\) The lectins have been shown to bind to a number of cells including erythrocytes (non-specific to blood type),\(^{(43,44)}\) lymphocytes, leukocytes, macrophages, glycoproteins and plasma proteins.\(^{(44,45)}\) Binding has been found to be stereospecific towards units containing a d-galactose molecule,\(^{(42,45,46)}\) although d-galactose units with unmodified hydroxyl groups at C\(_2\), C\(_3\) and C\(_4\) inhibit erythrocyte agglutination.\(^{(47)}\) Tyrosine residues are also thought to be involved in the agglutination process.\(^{(46)}\) Plasma proteins compete for the lectin receptor site and, therefore, decrease the agglutination of erythrocytes and tumour cells.\(^{(48)}\) Unlike many other sugars, lactose units have also been found to inhibit erythrocyte agglutination.\(^{(47)}\)

Mistletoe lectins have been reported to prevent viscotoxin- and allergen-stimulated histamine release from human leukocytes.\(^{(49)}\)

**Hypotensive effect**
The hypotensive effect documented for mistletoe has been attributed to various biologically active constituents such as acetylcholine, histamine, gamma aminobutyric acid (GABA), tyramine and flavones.\(^{(G24)}\) The exact nature of the hypotensive effect of mistletoe seems unclear: it has been reported that activity is mainly due to an inhibitory action on the excitability of the vasomotor centre in the medulla oblongata.\(^{(50)}\) However, it has also been stated that the hypotensive action of mistletoe is mainly of a reflex character, exerting a normalising effect on both hypertensive and hypotensive states.\(^{(50)}\) The effect of different mistletoe plant parts and host plant on the hypotensive activity has been studied with highest activity reported for mistletoe leaves parasitising on willow.\(^{(50)}\)

**Antimicrobial activity**
Extracts of the *V. album* subspecies *album*, *abietis*, *austriacum* and *pallasiana* have been tested for activity against *Mycobacterium tuberculosis* H37A \textit{in vitro}. Ether extracts of the *V. album* subspecies *album*, *abietis* and *austriacum* and petroleum ether extracts of the subspecies *abietis* and *austriacum* were active against *M. tuberculosis* (minimum inhibitory concentration, MIC:
200 μg/mL for each). From these results, it is not possible to determine whether polar or non-polar compounds are responsible for the observed antimycobacterial activity.

**Clinical studies**

Numerous clinical studies of mistletoe preparations have been carried out, mostly in the area of oncology, although few used rigorous randomised controlled clinical trial methodology and meet today's good clinical practice standards for the conduct of clinical trials. Collectively, the results of these studies are difficult to interpret as they involved patients with different types of cancer, who were or were not receiving different types of conventional cancer treatment (surgery, radiotherapy and/or cancer chemotherapy), and investigated the effects of different mistletoe preparations which vary markedly in their qualitative and quantitative composition. Hence, results regarding efficacy (and safety) need to be considered for each individual product.\(^{(56)}\)

Nevertheless, several narrative reviews and systematic reviews of clinical trials of mistletoe preparations involving patients with cancer have been published. In short, there is insufficient evidence from well-designed clinical trials that mistletoe preparations, or their isolated components, are effective in the treatment of cancer or as adjunctive treatments in cancer. Although a small number of reasonably well-designed trials have reported statistically significant results for mistletoe, compared with controls, at present, there is insufficient evidence to support the use of any specific mistletoe preparation in the treatment of any specific type of cancer. Further research using rigorous clinical trial methodology to test the effects of well-characterised mistletoe extracts is warranted. A summary of the clinical trial evidence relating to mistletoe preparations is given below.

**Pharmacokinetics**

There is a lack of clinical pharmacokinetic data for the constituents of mistletoe preparations.

**Therapeutic effects**

Cancer A systematic review of prospective, controlled clinical trials of mistletoe preparations in the treatment of cancers included 23 studies, of which 16 involved random allocation to treatment, two involved quasi-randomisation and the remaining five were non-randomised studies; three of the studies were nested within an epidemiological cohort study.\(^{(52)}\) In total, the trials involved over 3500 patients with different types of cancer (and at different stages), including breast (\(n = 4\) studies), lung (4),
colorectal (3), melanoma (2), stomach, kidney, bladder, glioma, and head and neck (1 each); several trials studied more than one cancer type. Most trials tested the anthroposophical mistletoe preparations Iscador and Helixor \((n = 14\) and 3, respectively), whereas in six the intervention was Eurixor. In most trials, the mistletoe intervention was administered in addition to conventional treatment (cancer chemotherapy, radiotherapy, corticosteroids and/or surgery) and compared with conventional treatment alone; other trials involved a no-treatment control group, or compared the mistletoe preparation directly with conventional treatment. Only two studies included a placebo control group, and blinding of the intervention was not undertaken in any of the trials (reliable blinding is hard to achieve as parenteral administration of mistletoe can cause local skin reactions at the injection site). An assessment of the methodological quality of the included clinical trials revealed that many of the studies had methodological limitations to their design and/or analysis and that most did not meet contemporary standards in these respects.

The results of this group of clinical trials need to be considered critically because of the poor methodological quality of the studies. Furthermore, the heterogeneity of the studies precluded a quantitative assessment of effect size. Of the 23 trials, 12 recorded statistically significant results in favour of the mistletoe intervention, compared with control, in at least one clinically relevant outcome measure (e.g. mean/median survival time, five/six-year survival rate). Of the five trials that incorporated a quality-of-life outcome measure, three reported statistically significant results in favour of mistletoe, compared with control, one reported no effect and one result was omitted from the original publication of the study. Three other trials reported statistically significant results for mistletoe, compared with control, in reducing the frequency of adverse effects of concomitantly administered conventional cancer treatments. In summary, although several trials reported positive results for certain mistletoe preparations, the data are inconclusive because of the studies’ methodological limitations, and further research using well-designed and properly conducted trials is necessary.

Another systematic review of prospective controlled clinical trials of mistletoe preparations involving patients with cancer employed stricter inclusion/exclusion criteria: only randomised trials were considered. The review included 10 studies, eight of which were included in the systematic review described above; the other two studies were available only as a conference abstract and unpublished data from a manufacturer of a mistletoe preparation, which prevented their thorough critical analysis. Of the eight trials for which a full assessment of methodological quality was possible, the best five scored 3 points from a maximum of 5; as with the
previous review,\(^{(53)}\) a quantitative analysis of data from the included studies was not possible because of the heterogeneity of the trials. From this review, it was concluded that the evidence does not provide strong support for the efficacy of mistletoe preparations as a treatment for cancer, or as supportive therapy in cancer.\(^{(53)}\)

The two reviews\(^{(52,53)}\) described above have superseded a previous systematic review\(^{(54)}\) of controlled clinical trials of mistletoe preparations in cancer, although there is little difference in the respective conclusions of the more recent and older reviews. The earlier review included 11 controlled trials, only one of which was judged to be of high methodological quality, although even this did not involve masking (blinding) of the intervention. This trial was the only study included in the review that did not show a beneficial effect on outcome for a mistletoe preparation. The review concluded that mistletoe extracts could not be recommended in the treatment of patients with cancer, except in the setting of clinical trials.\(^{(54)}\)

One other review (which does not fully describe the methods used so must be described as non-systematic) included only those trials which had evaluated the effects of well-characterised mistletoe extracts standardised for their content of mistletoe lectins I, II and III. The conclusions of the review concur with those of systematic reviews in that mistletoe preparations do not have an established place in oncology and that further rigorous clinical trials are necessary.\(^{(55)}\) This review did not involve a formal assessment of the methodological quality of included studies, but did highlight that two trials\(^{(11,56)}\) only were conducted using contemporary rigorous clinical trial methodology and in accordance with the principles of good clinical practice. These two studies were included in the two recent systematic reviews\(^{(52,53)}\) described above, but at the time, data from one study\(^{(56)}\) were available only in promotional literature produced by a manufacturer of a mistletoe preparation.

One of these studies was a randomised controlled trial of an ML-1 standardised mistletoe extract (Eurixor) involving 477 patients with head and neck squamous cell carcinoma. Participants were stratified into two groups – surgery only, or surgery followed by radiotherapy groups – and randomised to receive additional treatment with the mistletoe extract.\(^{(11)}\) The dosage regimen for mistletoe extract was equivalent to ML-1 1 ng/kg body weight twice weekly by subcutaneous injection over a 60-week period in treatment cycles of 12 weeks followed by a break in treatment after every four weeks. Mistletoe treatment was commenced 1–4 days before surgery in 72% of participants; the remainder (for whom early administration was not possible) first received mistletoe shortly after
undergoing surgery.

In total, 200 (42%) of the 477 participants experienced a relapse during the follow-up period (median four years for surviving participants); there was no statistically significant difference between the mistletoe and control groups in incidence of relapse and in the development of metastases and second primary tumours.\(^{(11)}\) In an intention-to-treat analysis, there was no statistically significant difference between the two groups in disease-free survival, the primary efficacy parameter (adjusted hazard ratio, 95% confidence interval (CI): 0.959, 0.725–1.268). There were also no statistically significant differences between the treatment and control groups in secondary outcome measures (immune parameters, quality-of-life score) assessed at different time points throughout the study. The conclusion of the study was that the ML-1 standardised mistletoe extract tested (Eurixor) is not effective as an adjuvant treatment for patients with head and neck cancer.\(^{(11)}\)

The second of these studies (now published as an abstract so full details are still not available) was a randomised, double-blind, multicentre trial involving 272 patients with breast cancer who had undergone surgery and were receiving treatment with cyclophosphamide, methotrexate and 5-fluorouracil. In addition, they received one of three doses of mistletoe extract (Lektinol) equivalent to 5, 15 or 35 ng ML-1, or placebo, twice weekly by subcutaneous injection for 15 weeks. This was the first study to incorporate double-blinding into the study design, although it is not clear from the published information how this was achieved and whether or not a check on the success of blinding was carried out. Participants who received the intermediate dose of mistletoe extract had statistically significant improvements in quality-of-life scores, compared with those in the placebo group; this result was not observed with the lower and higher doses of mistletoe extract.\(^{(56)}\)

This result is difficult to interpret from a dose–response perspective. However, the assumption that the optimal dose of a herbal medicine is the highest tolerated dose has been challenged previously, particularly since some herbal medicines, including mistletoe, appear to act at least in part through immunostimulatory or immunomodulatory mechanisms.\(^{(57)}\) This view is borne out by the results of the present study\(^{(56)}\) since the apparently effective dose of mistletoe extract (equivalent to ML-1 15 ng, twice weekly) was not the highest dose tested (mistletoe extract equivalent to ML-1 35 ng, twice weekly). Furthermore, the effective dose was much lower than that administered in a previous study\(^{(11)}\) (equivalent to ML-1 1 ng/kg body weight, albeit of a different ML-1 standardised mistletoe
The results of another prospective randomised controlled trial, which have now been published in more detail since systematic reviews were conducted, have echoed previous findings that mistletoe extract (in this case, Iscador M) is not effective as an adjuvant treatment in cancer. Patients involved in this study had either high-risk stage II (thickness > 3 mm) or stage III (regional lymph node metastases) melanoma without distant metastases. The study, carried out by the German Cancer Society (DKG), at present is published only as a joint report with another randomised controlled trial set up by the European Organisation for Research and Treatment of Cancer (EORTC) to assess the value of low doses of the recombinant interferons (rIFNs) rIFN-α2β and rIFN-γ; the DKG study contributed patients randomised to the rIFNs and control groups. With respect to mistletoe, the report provides analysis only for the Iscador M group versus the control group, and not for Iscador M versus rIFNs.

The DKG study randomised 407 patients to receive Iscador M in escalating doses as recommended (n = 102), rIFN-α2β 1 MU subcutaneously every other day (n = 101), rIFN-γ 0.2 mg subcutaneously every other day (n = 102), or control (n = 102), for one year, or until tumour progression. The Iscador M regimen was as follows: escalating doses from 0.01 to 1.0 mg/mL (volume administered not stated) every other day over two weeks, resumed after three treatment-free days for 14 doses of 20 mg/mL every other day, followed by seven treatment-free days, and repeated over one year (or until tumour progression).

In an intention-to-treat analysis, there was no statistically significant difference between the mistletoe group and the control group with respect to the primary end-point of the study, the disease-free interval, and duration of survival (estimated hazard ratio (95% CI): 1.32 (0.93–1.87), \( p = 0.12 \) and 1.21 (0.84–1.75), \( p = 0.31 \) for disease-free interval and survival, respectively). Thus, adjuvant treatment with Iscador M in patients with melanoma did not lead to a more favourable outcome, when compared with control. The findings raised the hypothesis that mistletoe treatment potentially may have an adverse effect in patients with melanoma by stimulating melanoma cell proliferation.

Another prospective, randomised, controlled, multicentre trial, conducted in China, compared the effects of a mistletoe preparation (Helixor A) with those of a polysaccharide biological response modifier (Lentinan; 4 mg by
intramuscular injection, daily) on quality of life in 233 patients with breast
(\(n = 68\)), ovarian (\(n = 71\)) and non-small-cell lung cancer (\(n = 94\)) who
also received conventional cancer chemotherapy.\(^{59}\) Participants
randomised to the mistletoe group received escalating doses, administered
three times weekly by subcutaneous injection and starting with 1 mg up to
a maximum of 200 mg (although it is not stated in a report of the study,
this probably refers to the amount of mistletoe extract, rather than the
mistletoe lectin content). The duration of treatment for standard and
adjuvant therapy was not stated in a report of the trial.

A statistically significant improvement in the Karnofsky Performance Index
(which measures physical condition) was reported for the mistletoe group,
compared with the placebo group (proportion of participants reporting an
improvement: 50.4 and 32.4% for Helixor A and Lentinan, respectively; \(p = 0.002\)).\(^{59}\) However, it is not known to what extent the lack of blinding of
treatment allocation could have influenced the results and, in view of this
and other methodological limitations (e.g. no sample size calculation), the
results of this trial cannot be considered conclusive.

A new instrument has been developed to measure specifically the quality of
life of patients with cancer treated with mistletoe preparations in the
clinical trial setting.\(^{60}\)

**Effects on Immunological Parameters** Immunomodulatory effects do not necessarily
translate into clinical efficacy, and the influence of some immune
parameters, for example, interleukin-1 and -6, may be to stimulate the
growth of certain types of tumour cells.\(^{15}\) Other factors, such as resistance
to changes in immune parameters can occur in advanced stages in some
types of cancer, thus the responses of different tumour types towards
immunomodulatory effects need to be considered separately.

Many studies involving patients with cancer have investigated the effects of
mistletoe preparations on immune parameters, such as lymphocytes,
natural killer cells and cytokines. Most, but not all, of these studies describe
effects for different mistletoe preparations. The results of several studies
have been summarised in a non-systematic, yet comprehensive, review of
clinical trials of well-characterised, lectin-standardised mistletoe extracts in
patients with different types of cancer.\(^{55}\)

The two most methodologically rigorous randomised controlled trials (see
Therapeutic effects, Cancer), which assessed the effects of mistletoe
extracts on immune parameters (as well as on clinical outcome measures),
reported conflicting results. In a randomised controlled trial involving 477
patients with head and neck squamous cell carcinoma who received an ML-1
standardised mistletoe extract (Eurixor) in addition to conventional treatment, there were no statistically significant differences between the treatment and control groups in blood concentrations of the following lymphocyte subsets when assessed at different time points throughout the study: CD3+, CD4+, CD8+, CD19+, CD16+53+3+, CD16+53+3, CD25+ and CD3+DR. In contrast, a randomised, double-blind, multi-centre trial involving 272 patients with breast cancer who received mistletoe extract (Lektinol) equivalent to 5, 15 or 35 ng ML-1, or placebo, in addition to conventional treatment found a statistically significant increase in blood concentrations of CD4+ cells in the mistletoe group, compared with the placebo group.

Other studies discussed in the review had methodological flaws and their results cannot be considered definitive. In one of these studies – a randomised, controlled trial involving 35 patients with malignant stage III/IV glioma – statistically significant increases in peripheral blood CD3+, CD4+ and CD8+ cell counts were observed after three months’ treatment with mistletoe extract (Eurixor) equivalent to 1 ng/kg body weight twice weekly, compared with values for the control group. Another trial, in which 47 patients with advanced breast cancer were randomised to receive mistletoe extract (Eurixor) equivalent to 0.5 or 1 ng/kg body weight twice weekly by subcutaneous injection for eight weeks followed by a four-week break, for at least two cycles, reported increases in plasma β-endorphin concentrations and enhanced activity of natural killer cells and T-lymphocytes, compared with pretreatment values. This trial supported the findings of an earlier similar study, which also reported increased in vitro release of cytokines (IL-2, IFN-γ and TNFα) by mononuclear cells from patients with breast cancer treated with mistletoe extract (Eurixor).

Other preliminary studies have described the production of anti-ML-1 antibodies in patients with cancer who were treated with mistletoe extracts. The antibodies produced were typically of the immunoglobulin G type, although in some instances, antibodies of the immunoglobulin E and other types were produced. There is some evidence that the antibody response differs depending on the type of mistletoe preparation administered (i.e. the profile of antigens within the extract). In association with this work, in vitro studies showed that anti-ML-1 antibodies neutralised the cytotoxic effect of mistletoe lectins on peripheral blood mononuclear cells and tumour cell lines.

In a small (n = 14), uncontrolled study, an increase in the incorporation of [3H]thymidine into the DNA of unstimulated lymphocytes, as a marker for DNA repair, was observed in patients with breast cancer who received
mistletoe extract (Iscador M; single intravenous infusion then, after two
days, subcutaneous injections daily for seven days).\textsuperscript{(69)} The effect was
greatest on days 7–9 of treatment when values were 2.7 times higher than
pretreatment values. This finding raised several hypotheses which require
further testing.

Mistletoe preparations have also been documented to induce immunological
responses in healthy volunteers. In a randomised, double-blind, placebo-
controlled study involving 48 healthy volunteers, Iscador Q (in increasing
doses twice weekly by subcutaneous injection, over 12 weeks), but not
Iscador P (in increasing doses twice weekly by subcutaneous injection over
12 weeks), induced eosinophilia after five weeks’ treatment which persisted
until the end of treatment (week 12), when compared with placebo.\textsuperscript{(70)} At
the end of the study, anti-ML-1 antibodies were detected in all 16
volunteers in the Iscador Q group, but in less than half those exposed to
Iscador P, whereas anti-viscotoxin A2 immunoglobulin G antibodies were
found in all mistletoe-exposed participants.\textsuperscript{(71)} In a preliminary study
involving healthy volunteers who received Iscador Q by subcutaneous
injection, significant increases in the microcirculation were observed, both
local to the injection site and in other target tissues investigated (e.g.
gingival, rectal). Increased transmigration of white blood cells was also
noted both locally and in other target tissues, a phenomenon which is
believed to be important in the immunological response.\textsuperscript{(72)}

In other studies, T-lymphocytes were obtained from blood samples from
patients with cancer who had received subcutaneous injections three times
weekly of aqueous mistletoe extracts prepared from mistletoe grown on
apple and pine trees to examine their response to subsequent incubation
with the mistletoe extracts \textit{in vitro}.\textsuperscript{(73,74)} Proliferation of T-lymphocytes
from mistletoe-treated patients occurred, but was not observed for T-
lymphocytes obtained from untreated control subjects. Further investigation
revealed that the responding cells were the CD4+ T-cell subset.\textsuperscript{(73,74)} A
small uncontrolled study described increases in the release of cytokines,
such as TNFα and IL-6, measured in the supernatants of cultured peripheral
blood mononuclear cells obtained from patients with breast cancer who
were treated with mistletoe extract (ABNOBAViscum, 0.2 mg fresh plant
material/30 kg body weight by subcutaneous injection twice weekly for 16
weeks).\textsuperscript{(66)}

\textbf{Other conditions} Clinical investigation of the effects of mistletoe preparations
in conditions other than cancer is limited. Several preliminary
investigations in patients with chronic hepatitis C infection,\textsuperscript{(75)} respiratory
infections,\textsuperscript{(76,77)} and HIV infection\textsuperscript{(78)} have been undertaken. Some of these
studies have reported improvements in clinical outcomes and immunological parameters in mistletoe recipients, compared with pretreatment values or with placebo or other control group values. However, due to the methodological limitations of these studies, the results are inconclusive.
Side-effects, Toxicity

Only limited data are available on the safety of mistletoe preparations. Many clinical trials have not provided an assessment of safety or have reported little information on safety aspects, and there is a paucity of well-designed postmarketing surveillance studies of mistletoe preparations.

The UK Medicines and Healthcare products Regulatory Agency’s (MHRA) national spontaneous reporting scheme for suspected adverse drug reactions has received a total of four reports of suspected adverse drug reactions (ADRs) associated with Iscador preparations. These reports described 12 suspected ADRs, including hepatitis and abnormal liver function (two reports), myalgia, dizziness, wheezing, cough, hoarseness, abdominal pain, pyrexia, urticaria (one report) and a suspected interaction (one report). The MHRA has received one further suspected ADR report of arteritis associated with a multi-ingredient product containing mistletoe (no further details available at the time of writing). Causality in these cases has not been established.

Reviews of clinical trials which have provided an assessment of the safety of mistletoe, generally have reported mistletoe preparations to be well tolerated. The most frequently reported adverse events in trials using ML-standardised mistletoe preparations were local reactions, such as urticaria and/or erythema, at the injection site which usually resolved within 48 hours. The frequency of occurrence of such reactions in patients with cancer who received mistletoe preparations by injection has been described as being up to 48%.

This information, however, comes mostly from small-scale clinical trials involving patients with different types of cancer who were also receiving various different conventional treatments (surgery, radiotherapy and/or cancer chemotherapeutic agents) in addition to mistletoe preparations. Furthermore, studies have assessed the effects of different types of mistletoe preparations, which vary both qualitatively and quantitatively in their composition, administered according to different dosage regimens. Against this background, the safety of each individual mistletoe preparation needs to be considered.

In a randomised controlled trial involving 477 patients with head and neck squamous cell carcinoma receiving conventional treatment, an ML-1 standardised mistletoe extract (Eurixor) was administered in doses equivalent to ML-1 1 ng/kg body weight twice weekly by subcutaneous injection over a 60-week period in treatment cycles of 12 weeks followed by a break in treatment after every four weeks; the control group did not receive any
additional treatment.\(^{(11)}\) In total, 43% of mistletoe recipients developed local and/or systemic adverse events at the onset of mistletoe treatment, although this fell to around 8% by week 32 or later of treatment. The most common local reactions occurring at the onset of treatment were rubor, prurigo and indurations (hard spots); vesiculation was observed in 1–3% of mistletoe recipients over the treatment period. Other reactions (melalgia, fever, sleeplessness, tiredness, cold or heat sensations and sneezing) occurred in 1–4% of the mistletoe group. In total, around 18% of mistletoe recipients refused further injections during the study (after a median of 12 weeks) because of adverse events believed to be induced by mistletoe.\(^{(11)}\)

In a randomised controlled trial involving 407 patients with melanoma who received mistletoe extract (Iscador M in escalating doses as recommended), rIFN-α2β, rIFN-γ, or control, for one year or until tumour progression, study withdrawals due to anorexia, malaise, depressive moods, fever and inflammation at the injection site occurred in 4.9, 4.6 and 7.8% of the mistletoe-, rIFN-α2β- and rIFN-γ recipients, respectively.\(^{(58)}\) Apart from the statement that no organ toxicity was observed during the study, no further information on safety aspects was provided.

Several small, non-randomised, controlled studies have assessed the tolerability of mistletoe preparations in immunocompromised HIV-positive and healthy individuals. In two studies in which participants received mistletoe extracts (Iscador Qu FrF or Iscador Qu Special) by subcutaneous injection twice weekly in increasing doses for up to 68 weeks, erythema at the injection site occurred in all but one participant.\(^{(80,81)}\)

Other adverse events rated to be possibly or probably associated with mistletoe extract (Iscador Qu FrF) were headache, fatigue, fever and inflammation; diarrhoea and nausea occurred in small numbers of participants, but were not thought to be associated with mistletoe treatment.\(^{(80,81)}\) Significant increases in serum urea nitrogen and creatinine concentrations were observed for HIV-positive participants after mistletoe treatment, compared with baseline values (statistical comparisons with the control group were not undertaken), but concentrations remained within the normal range; no other statistically significant changes in biochemical parameters, including liver function test values, were seen.

Aspects of the local skin reaction that occurs following subcutaneous injection of mistletoe extracts (Iscador QFrF or Qu Spezial) have been described following studies involving healthy volunteers.\(^{(82,83)}\) The reaction appears to involve a subepidermal lymphomonocytic infiltrate, induced by mistletoe extract, at the injection site.
Hepatitis has been documented in a woman who had ingested a herbal preparation containing kelp, motherwort, scullcap and mistletoe. Mistletoe was assumed by the authors of the report to be the causal factor since it was the only ingredient of the remedy known to contain toxic constituents (although these had not previously been linked with hepatotoxicity). No other instances of hepatotoxicity have been associated with mistletoe ingestion and, since the case report was published, hepatitis has been documented with scullcap (see Scullcap).

_V. album_ should not be confused with American mistletoe (_Phoradendron leucarpum, P. serotinus, P. flavescens_). The safety of American mistletoe has been reviewed.\(^{(G21,85-87)}\)

**Anaphylactic reactions**

There are isolated reports of anaphylactic reactions occurring in individuals who have received parenteral treatment with mistletoe extracts. One report described three such cases involving patients who were already undergoing treatment with mistletoe, but did not provide full details of the mistletoe preparations administered, the reactions and outcomes, and did not undertake a causality assessment.\(^{(88)}\) One case involved a non-atopic, 44-year-old man who was admitted to an intensive care unit with hypotension, urticaria and loss of consciousness which had developed within 5 minutes of an injection of _V. album L._ (Quercus) 10 mg (route of injection not specified). The man was treated with intravenous fluids and corticosteroids. A similar case involved a 70-year-old non-atopic woman after receiving her 25th dose (0.1 mg) of mistletoe extract; one month later, she displayed a positive result (which included generalised pruritus, pharyngeal oedema and malaise) following an intradermal challenge with _V. album_ at one-tenth dilution.\(^{(88)}\) The third case involved a 42-year-old man with multiple sclerosis who received a maintenance dose of _V. album_ (Quercus) 10 mg and who immediately developed pruritus, oedema, asthma and hypotension.

**Preclinical studies**

Data from an _in vitro_ study involving human melanoma cell lines indicate that clinically relevant low concentrations of mistletoe lectin may stimulate tumour cell proliferation.\(^{(89)}\) In contrast, other _in vitro_ experiments did not demonstrate that an aqueous mistletoe extract (containing galactoside-specific mistletoe lectin 265 μg/mL) stimulated tumour cell proliferation.\(^{(90)}\)

Toxicity in animals has been documented for mistletoe lectins and viscotoxins. Intravenous administration of viscotoxin to cats (35 μg/kg) resulted in a negative inotropic effect on cardiac muscle, reflex bradycardia and hypotension.\(^{(91)}\) Viscotoxins A3 and B have also caused muscle contracture
and progressive depolarisation in isolated smooth, skeletal and cardiac muscle preparations (rabbit, frog).\(^{(92)}\) The mode of action was thought to involve the displacement of calcium from cell membrane-bound sites. The viscotoxins precipitate histamine release from human leukocytes in an irritant manner without destroying the cells.\(^{(49)}\) Viscotoxin is toxic on parenteral administration and an LD\(_{50}\) value (mice, intraperitoneal injection) has been estimated as 0.7 mg/kg.\(^{(93)}\)

Mistletoe lectins inhibit protein synthesis in both cells and cell-free systems.\(^{(94)}\) In common with other known toxic lectins (e.g. ricin), mistletoe lectins bind to plasma proteins, are specific towards d-galactose, possess some cytotoxic activity and have caused macroscopic lesions in rats (e.g. ascites, congested intestine, pancreatic haemorrhages).\(^{(38)}\) An LD\(_{50}\) (mice) value for mistletoe lectin fraction is reported as 80 μg/kg compared with 3 μg/kg for ricin.\(^{(94)}\)

Documented LD\(_{50}\) values (mice, intraperitoneal injection) are greater than 2.25 mg for the polysaccharide fraction from the berries, 32 mg for the crude plant juice, and 276 mg for Iscador.\(^{(38)}\)
Contra-indications, Warnings

Mistletoe is contra-indicated in cases of known allergy to mistletoe preparations, acute inflammatory conditions and high fever, and chronic progressive conditions, such as tuberculosis.\(^{G56}\) Increases in body temperature can occur following mistletoe injection.

**Interactions**

In view of the documented pharmacological activities, it should be considered whether or not there is potential for clinically important interactions between mistletoe and other medicines with similar or opposing effects, such as existing cardiac, immunosuppressant, hypo/hypertensive, antidepressant and anticoagulant/coagulant therapies.

**Pregnancy and lactation**

The use of mistletoe is contra-indicated in pregnancy and breastfeeding in view of the toxic constituents.
The chemistry of mistletoe is well-documented (see Constituents). The profile of constituents is, to some extent, thought to be dependent on the host plant on which mistletoe is a parasite. The lectin and viscotoxin constituents are considered to be the main active principles, although there is some evidence that certain activities, e.g. cytotoxicity, can be modulated by other constituents. Mistletoe is reputed to be a cardiac depressant, although cardioactive constituents are not generally recognised as constituents of mistletoe; however, this may depend on the host plant on which mistletoe has grown.

Scientific investigation into the effects of mistletoe has centred primarily on the pharmacological and toxicological properties of mistletoe extracts and the lectin and viscotoxin constituents. Preclinical research has focused on the cytotoxic and immunomodulatory activities, and provides a large body of supporting evidence with regard to clinical use of mistletoe in cancer and other conditions involving the immune system.

The results of clinical trials of various commercial mistletoe preparations and purified components have, however, been less convincing. Furthermore, different mistletoe preparations vary markedly in their qualitative and quantitative composition. Hence, results regarding efficacy (and safety) need to be considered for each individual product.\textsuperscript{G56}

Numerous clinical studies of mistletoe preparations have been conducted, although few used rigorous randomised controlled clinical trial methodology. Several systematic reviews of clinical trials of mistletoe preparations involving patients with cancer have concluded that there is insufficient evidence from well-designed clinical trials that mistletoe preparations, or their isolated components, are effective in the treatment of cancer or as adjunctive treatments in cancer. Further research using rigorous clinical trial methodology to test the effects of well-characterised mistletoe extracts is warranted.

Reviews of clinical trials which have provided an assessment of the safety of mistletoe, generally have reported mistletoe preparations to be well tolerated.\textsuperscript{52,55} The most frequently reported adverse events in trials using lectin-standardised mistletoe preparations are local reactions, such as urticaria and/or erythema at the injection site, which usually resolve within 48 hours.\textsuperscript{55}

The toxic nature of the mistletoe constituents (e.g. lectins, viscotoxins) indicates that mistletoe is unsuitable for self-medication. Mistletoe berries are...
poisonous and, in the UK, sale/supply is restricted to pharmacies only and by or under the supervision of a pharmacist.\(^{(95)}\)
References

See also General References G2, G3, G9, G56, G67.


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Species (Family)

*Leonurus cardiaca* L. and various other *Leonurus* species (Labiatae)
Synonym(s)

Leonurus
Part(s) Used
Herb
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

Complete German Commission E\(^{(G3)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)

Ph Eur 2004\(^{(G72)}\)
Legal Category (Licensed Products)

GSL(G37)
Constituents

See General References G6 G22 G40 G49 G64.

**Alkaloids**
0.35%. Stachydrine (a pyrrolidine–type alkaloid), betonicine and turicin (stereoisomers of 4–hydroxystachydrine), leonurine 0.0068% (a guanidine derivative),\(^{(1)}\) leonuridin, leonurinine. The presence of leonurine in *L. cardiaca* has been disputed, although it has been documented for other *Leonurus* species.

**Flavonoids**
Glycosides of apigenin, kaempferol, and quercetin (e.g. hyperoside, kaempferol–3-D-glucoside, genkwanin, quinqueloside, quercitrin and rutin).\(^{(2,3)}\)

**Iridoids**
Ajugol, ajugoside, galiridoside, leonurid and three or four more unidentified glycosides.\(^{(4)}\)

**Tannins**
2–8%. Type not specified. Pseudotannins (e.g. pyrogallol, catechins).

**Terpenoids**
Volatile oil 0.05%, resin, wax, ursolic acid, leocardin (a labdane diterpene)\(^{(5)}\) as an epimeric mixture, and a diterpene lactone similar to marrubiin.\(^{(2)}\) Cardiac glycosides (bufadienolide/bufanolide type) have been documented although their presence in motherwort has not been confirmed.

**Other constituents**
Citric acid, malic acid, oleic acid, bitter principles,\(^{(6,7)}\) carbohydrates 2.89%, choline and a phenolic glycoside (caffeic acid 4–rutinoside).\(^{(8)}\)

A *Cad*-specific lectin has been isolated from the seeds.\(^{(9)}\)
Food Use

Motherwort is not used in foods. In the USA, motherwort is listed by the Food and Drugs Administration (FDA) as a Herb of Undefined Safety. (G22)
Herbal Use
Motherwort is stated to possess sedative and antispasmodic properties. Traditionally, it has been used for cardiac debility, simple tachycardia, effort syndrome, amenorrhoea, and specifically for cardiac symptoms associated with neurosis. (G6 G7 G8 G64)
Dosage

*Dried herb*
2–4 g or by infusion three times daily.\(^{(G6 G7)}\)

*Liquid extract*
2–4 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G6 G7)}\)

*Tincture*
2–6 mL (1 : 5 in 45% alcohol) three times daily.\(^{(G6 G7)}\)
Pharmacological Actions

**In vitro and animal studies**

The uterotonic principle in motherwort is unclear, although leonurine is reported to be the utero-active constituent in various *Leonurus* species. In addition, oxytocic activity documented for *L. cardiaca* has been attributed to another alkaloid constituent, stachydrine.\(^{[G30]}\) Uterotonic activity has been reported for leonurine in various *in vitro* preparations including human myometrial strips and isolated rat uterus.\(^{[10,11]}\)

*In vitro* cardioactivity has been documented for motherwort.\(^{[12]}\) An alcoholic extract was found to have a direct inhibitory effect on myocardial cells: antagonistic action towards calcium chloride (provided that the extract was administered before calcium chloride), and towards both α- and β-adrenoceptor stimulation was observed. No significant effect on the cardiac activity of the isolated guinea-pig heart was noted for caffeic acid 4–rutinoside.\(^{[8]}\)

A related species, *Leonurus heterophyllus*, has been stated to prevent platelet aggregation, although no such documented action was located for motherwort.\(^{[13]}\)

Ursolic acid has been reported to possess antiviral, tumour-inhibitory and cytotoxic activities.\(^{[14,15]}\) Ursolic acid was found to inhibit the Epstein–Barr virus *in vitro* and to inhibit tumour production by 12-O-tetradecanoyl phorbol (TPA) in mouse skin, with activity comparable to that of retinoic acid, a known tumour–promoter inhibitor.\(^{[15]}\) *In vitro* cytotoxicity was documented in lymphocytic leukaemia (P-388, L-1210), human lung carcinoma (A-549), KB cells, human colon (HCT-8) and mammary tumour (MCF-7).\(^{[14]}\)
Side-effects, Toxicity

It has been stated that the leaves of motherwort may cause contact dermatitis and that the lemon-scented oil may result in photosensitisation.\(^{(G51)}\) No documented toxicity studies were located. Cytotoxic activities have been reported for ursolic acid (see *In vitro* and animal studies).
Contra-indications, Warnings

Excessive use may interfere with existing therapy for a cardiac disorder (cardiac glycoside constituents, \textit{in vitro} activity). Sensitive individuals may experience an allergic reaction.

\textit{Pregnancy and lactation}
Motherwort is reputed to affect the menstrual cycle.\textsuperscript{(G22)} In view of the lack of toxicity data and the documented \textit{in vitro} uterotonic activity,\textsuperscript{(G30)} the use of motherwort during pregnancy and lactation should be avoided.
Pharmaceutical Comment

The common name motherwort may be applied to one of many *Leonurus* species. *L. cardiaca* is the typical European species utilised, whereas *Leonurus artemisia* is commonly used in traditional Chinese medicine. Other species referred to as motherwort include *Leonurus sibirious* and *L. heterophyllus*. The chemistry of *L. cardiaca* is well studied although the presence of the uterotonic principle leonurine has been disputed. Cardioactive properties in animals have been reported for motherwort (*L. cardiaca*), which thus support some of the stated herbal uses. However, any symptoms of cardiac disorder are not suitable for self-diagnosis and treatment with a herbal remedy. In view of the lack of toxicity data and possible cardioactivity, excessive use of motherwort should be avoided.
References

See also General References G3 G6 G9 G22 G30 G31 G36 G37 G40 G43 G49 G51 G64.

Myrrh
Species (Family)

i. *Commiphora molmol* Engl. (Bursuraceae)


iii. Other *Commiphora* species
Synonym(s)

i. African Myrrh, Balsamodendron Myrrha, Commiphora, Commiphora myrrha (Nees) Engl., Somali Myrrh

ii. Arabian Myrrh, Yemen Myrrh
Part(s) Used

Oleo–gum–resin
Pharmacopoeial and Other Monographs

BHC 1992\textsuperscript{(G6)}

BHP 1996\textsuperscript{(G9)}

BP 2002\textsuperscript{(G71)}

Complete German Commission E\textsuperscript{(G3)}

ESCOP 1999\textsuperscript{(G52)}

Martindale 33rd edition\textsuperscript{(G67)}

PDR for Herbal Medicines 2nd edition\textsuperscript{(G36)}

Ph Eur 2004\textsuperscript{(G72)}

USP26/NF21\textsuperscript{(G73)}
Legal Category (Licensed Products)

GSL$^{(G37)}$
Constituents
See General References G2 G6 G41 G48 G52 G64.

Carbohydrates
Up to 60% gum yielding arabinose, galactose, xylose, and 4-O-methylglucuronic acid following hydrolysis.

Resins
Up to 40% (average 20%) consisting of α-, β- and γ-commiphoric acids, commiphorinic acid, α- and β-heerabomyrrhols, heeraboresene and commiferin.

Steroids
Campesterol, cholesterol and β-sitosterol.

Terpenoids
α-Amyrin. Furanosesquiterpenes, including furaneudesma–1,3–diene (major), furaneudesma–1,4–diene–6–one, lindestrine, curzerenone, furanodiene, 2-methoxyfuranodiene and 4,5-dihydrofuranodiene–6–one.\(^1,G52\)

Volatile oils
1.5–17%. Main constituents are furanosesquiterpenes. Dipentene, cadinene, heerabolene, limonene, pinene, eugenol, \(m\)-cresol, cinnamaldehyde, cuminaldehyde, cuminic alcohol and others.
Food Use

Myrrh is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that myrrh can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product. 

In the USA, myrrh is permitted for use in alcoholic beverages.
Herbal Use

Myrrh is stated to possess antimicrobial, astringent, carminative, expectorant, anticatarrhal, antiseptic and vulnerary properties. Traditionally, it has been used for aphthous ulcers, pharyngitis, respiratory catarrh, common cold, furunculosis, wounds and abrasions, and specifically for mouth ulcers, gingivitis and pharyngitis. The German Commission E approved topical use for mild inflammation of the oral and pharyngeal mucosa.
Dosage

*Myrrh Tincture*
(BPC 1973) 2.5–5.0 mL; in a glass of water several times daily as a gargle or a mouthwash. For skin, undiluted or diluted.\(^{G52}\)

*Tincture Myrrh Co*
(Thompsons) (1 part Capsicum Tincture BPC 1973 to 4 parts Myrrh Tincture BPC 1973) 1.0–2.5 mL.
Pharmacological Actions

In vitro and animal studies

**Anti–inflammatory activity**
Anti–inflammatory (carrageenan–induced inflammation and cotton pellet granuloma)\(^{(2)}\) and antipyretic activities in mice\(^{(2,3)}\) have been documented for *C. molmol*.

**Hypoglycaemic activity**
Hypoglycaemic activity in both normal and diabetic rats has been reported for a myrrh extract.\(^{(4,5)}\) Together with an aloe gum extract, myrrh was found to be an active component of a multi–plant extract that exhibited antidiabetic activity. The mode of action was thought to involve a decrease in gluconeogenesis and an increase in peripheral utilisation of glucose in diabetic rats.

Myrrh is stated to have astringent properties on mucous membranes\(^{(G45)}\) and to have antimicrobial activities *in vitro*.\(^{(G41)}\)

**Anti–inflammatory activity**
Anti–inflammatory activities have been reported for an Indian plant, *Commiphora mukul*, commonly known as guggulipid. Anti–inflammatory activity was described for a crystalline steroidal fraction of guggulipid in both acute (carrageenan–induced rat paw oedema test) and chronic (adjuvant arthritis) models of inflammation.\(^{(6)}\)

**Lipid–lowering effects**
A ketosteroid has been identified as the active hypocholesterolaemic principle in guggulipid.\(^{(7)}\) In some animal species, thyroid suppression is required as well as cholesterol administration in order to achieve experimental hypercholesterolaemia. Results of studies in chicks administered a thyroid suppressant and cholesterol indicated that guggulipid prevents endogenous hypercholesterolaemia via stimulation of the thyroid gland.\(^{(7)}\) When fed to rabbits, guggulipid has been found to reverse the decrease in catecholamine concentrations and dopamine-β-decarboxylase activity that are associated with hyperlipidaemia.\(^{(8)}\)

**Stimulation of phagocytosis**
Stimulation of phagocytosis has been documented in mice inoculated with *Escherichia coli* and given with extracts of myrrh by intraperitoneal injection.\(^{(G52)}\)
**Cytoprotective activity**
An aqueous suspension of myrrh administered to rats at oral doses of 250–1000 mg/kg gave significant and dose–dependent protection to gastric mucosa against various ulcerogenic agents.\(^{(G52)}\)

**Analgesic activity**
In mice, powdered myrrh (1 mg/kg, orally) had significant analgesic activity in the hotplate test.\(^{(G52)}\) Isolated furanoeudesma–1,3–dione (50 mg/kg, orally) was significantly more effective than control \((p < 0.01)\) in the mouse writhing test, and the effective dose was reversed by naloxone (1 mg/kg).\(^{(G52)}\)

**Anti–tumour and cytotoxic activities**
In mice with Ehrlich solid tumours, an aqueous suspension of myrrh (250 or 500 mg/kg, orally) produced significant decreases in tumour weight \((p < 0.05)\) after 25 days.\(^{(G52)}\) Aqueous suspension of myrrh increased survival time in mice with Erlich ascite tumours.

**Clinical studies**
Well–designed clinical studies of myrrh are lacking. Guggulipid has been reported to lower the concentration of total serum lipids, serum cholesterol, serum triglycerides, serum phospholipids and β-lipoproteins in 20 patients.\(^{(9)}\) This effect was reported to be comparable to that of two other known lipid–lowering drugs also used in the study.
Side–effects, Toxicity

No reported side–effects were located for *C. molmol* or *C. abyssinica*. Hiccup,\(^9\) diarrhoea,\(^7\) restlessness and apprehension,\(^9\) were documented as side–effects for guggulipid when administered to 20 patients.\(^9\) Myrrh has been reported to be non–irritating, non–sensitising and non–phototoxic to human and animal skins.\(^{G41}\)
Contra-indications, Warnings

Myrrh may interfere with existing antidiabetic therapy, as hypoglycaemic properties have been documented. Thyroid stimulation and lipid lowering properties have been documented for the related species, *Commiphora mukul*.

**Pregnancy and lactation**
Myrrh is reputed to affect the menstrual cycle (G41) and the safety of myrrh taken during pregnancy has not been established. Excessive use of myrrh during pregnancy should be avoided.
The volatile oil, gum and resin components of myrrh are well documented. The anti-inflammatory and antipyretic activities documented in animals support some of the traditional uses. Phenol components of the volatile oil may account for the antimicrobial properties of myrrh, although no documented studies were located. Lipid-lowering properties via a stimulant action on the thyroid gland have been documented for *C. mukul* in both animals and humans. In view of the lack of toxicity data, excessive use of myrrh should be avoided.
References


Nettle
Species (Family)

*Urtica dioica* L. (Urticaceae)
Synonym(s)
Stinging Nettle, Urtica
Part(s) Used

Herb
Pharmacopoeial and Other Monographs

BHC 1992\textsuperscript{(G6)}

BHP 1996\textsuperscript{(G9)}

Complete German Commission E\textsuperscript{(G3)}

ESCOP 1996 and 1997\textsuperscript{(G52)}

Martindale 33rd edition\textsuperscript{(G67)}

Mills and Bone\textsuperscript{(G50)}

PDR for Herbal Medicines 2nd edition\textsuperscript{(G36)}
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents

See General References G6 G22 G52 G64.

**Acids**
Carbonic, caffeic, caffeoylmalic, chlorogenic, formic, silicic, citric, fumaric, glyceric, malic, oxalic, phosphoric, quinic, succinic, threonic and threono–1,4–lactone.\(^{(1)}\)

**Amines**
Acetylcholine, betaine, choline, lecithin, histamine, serotonin\(^{(2)}\) and a glycoprotein.\(^{(3)}\)

**Flavonoids**
Flavonol glycosides (e.g. isorhamnetin, kaempferol, quercetin).\(^{(4)}\)

**Inorganics**
Up to 20% minerals, including calcium, potassium and silicon.

**Lignans**
Several lignans, including (−)-secoisolariciresinol.

**Other constituents**
Choline acetyltransferase,\(^{(5)}\) scopoletin,\(^{(4)}\) β-sitosterol and tannin

**Other plant parts**
The rhizome contains lectin (*Urtica dioica* agglutinin) composed of six isolectins,\(^{(6,7)}\) coumarin (scopoletin), triterpenes (β-sitosterol, its glucoside, and six stearyl derivatives),\(^{(8,9)}\) two phenylpropane derivatives, and six lignans.\(^{(10)}\)
Food Use

Nettle (herbs and leaves) is listed by the Council of Europe as a natural source of food flavouring (category 1) (see Appendix 23).\(^{(G17)}\) Nettle is used in soups and herbal teas. In the USA, nettle is listed by the Food and Drugs Administration (FDA) as a Herb of Undefined Safety.\(^{(G22)}\)
Nettle is stated to possess antihaemorrhagic and hypoglycaemic properties. Traditionally, it has been used for uterine haemorrhage, cutaneous eruption, infantile and psychogenic eczema, epistaxis, melaena and specifically for nervous eczema. The German Commission E approved internal use of nettle leaf as supportive therapy for rheumatic ailments and as irrigation therapy for inflammatory disease of the lower urinary tract and prevention of kidney gravel; internal and external use for rheumatic ailments. The root is approved for difficulty in urination from benign prostatic hyperplasia.
**Dosage**

**Dried herb**
2–4 g or by infusion three times daily;\(^{(G6\ G7)}\) 8–12 g daily;\(^{(G3)}\) fresh juice 10–15 mL three times daily.\(^{(G52)}\)

**Liquid extract**
3–4 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G6\ G7)}\)

**Tincture**
2–6 mL (1 : 5 in 45% alcohol) three times daily.\(^{(G6\ G7)}\)
Pharmacological Actions

*In vitro* and animal studies

The pharmacological properties of nettle have been reviewed. Information from these reviews is summarised below.

**Anti–inflammatory activity**

An aqueous ethanol extract and also isolated caffeoylglycine partially inhibited the biosynthesis of arachidonic acid *in vitro*. Nettle extract (0.1 mg/mL) and isolated acid (1 mg/mL) inhibited 5–lipoxygenase–derived biosynthesis of leukotriene B4 by 20.8% and 68.2%, respectively, and inhibited synthesis of cyclooxygenase–derived prostaglandins (IC50 92 μg/mL and 38 μg/mL, respectively). The same extract significantly reduced tumour-necrosis-factor-α (TNFα) and interleukin 1β (IL-1β) concentrations after lipopoly saccharide (LPS)-stimulated secretion of these proinflammatory cytokines in human blood. An aqueous ethanol extract (0.25 mg/mL) inhibited platelet–activating factor (PAF)-induced exocytosis of elastase from human neutrophils by 93%, but failed to inhibit biosynthesis of prostaglandins from [14C]arachidonic acid.

*In vitro* addition of a commercial preparation of nettle leaf (IDS-23) to whole human blood resulted in an inhibition of LPS-stimulated TNFα and IL-1β secretion, correlating with drug ingestion. The same preparation inhibited phytohaemagglutinin–stimulated production of T helper cell 1 (Th1)-specific interleukin-2 (IL-2) and interferon-γ (IFNγ) in culture in a dose–dependent manner up to 50% and 74%, respectively. By contrast, T helper cell 2 (Th2)-specific interleukin-4 (IL-4) production was stimulated. The results suggested that the nettle leaf extract acts by mediating a switch in T helper cell–derived cytokine patterns and may inhibit the inflammatory cascade in autoimmune diseases such as rheumatoid arthritis. The transcription factor NF-κB is elevated in several chronic inflammatory diseases and is responsible for the enhanced expression of some proinflammatory gene products. A nettle leaf extract (IDS 23) potently inhibited NF-κB activation in a number of cells, including human T cells, macrophages, epithelial cells and mouse L929 fibrosarcoma cells *in vitro*. It was proposed that part of the anti–inflammatory effects of nettle may be due to its inhibitory effect on NF-κB activity.

**Benign prostatic hyperplasia activity**

Several lignans and their metabolites reduce binding activity of human sex hormone–binding globulin (SHBG) *in vitro*. Lignans from nettle are
competitive inhibitors of the interaction between SHBG and 5α-dihydrotestosterone. An aqueous extract of nettle root led to a concentration–dependent (0.6–10 mg/mL) inhibition of SHBG interaction with its receptor on human prostatic membranes. A 20% methanol extract of root inhibited binding capacity of SHBG after preincubation in human serum.

Subfractions of an aqueous methanol extract of nettle root inhibited cellular proliferation in benign prostatic hyperplasia (BPH) tissue. A root extract had a specific and concentration–dependent inhibition of human leukocyte elastase (HLE) activity in vitro. (HLE is an important marker in clinically silent genitourinary tract infection and inflammation.) Root extracts inhibited alternative and classic complementary pathways and significantly inhibited prostate growth in mice with induced BPH (by 51%, compared with control; \( p < 0.003 \)).

**Other activities**

CNS-depressant activity has been documented for nettle. It has been shown to produce a reduction in spontaneous activity in rats and mice, inhibition of drug–induced convulsions, and a lowering of body temperature in rats. Nettle has been reported to have no effect on the blood pressure of mice, whereas in cats it has produced a marked hypotensive effect and bradycardia. Atropine was reported to have no effect on these latter actions and a mode of action via \( \alpha \)-adrenoceptors was suggested.

Nettle is stated to contain both hypoglycaemic and hyperglycaemic principles. The hypoglycaemic component has been termed ‘urticin’ and nettle has been reported to lower the blood sugar concentration in hyperglycaemic rabbits.

An 80% ethanolic and an aqueous extract of nettle administered to mice at a dose of 25 mg/kg orally prior to glucose load, led to hypoglycaemia effects. No diuretic or ion excretion effects were observed in rats after oral administration of an aqueous extract of nettle (1 g/kg). Dried nettle had a potassium ion to sodium ion ratio of 63 : 1, whereas an aqueous decoction had a corresponding ratio of 448 : 1. It was suggested that the high potassium ion concentration in aqueous decoctions may contribute to their diuretic activity.

Uterine–activity has been documented for nettle in pregnant and non-pregnant mice; betaine and serotonin were stated to be the active constituents. A nettle extract was reported to be devoid of antifertility activity following oral administration to mice (250 mg/kg). Analgesic activity in mice has been documented. Administration of an aqueous extract (1200 mg/kg) to mice
showed resistance to stimulation in the hotplate test at 55°C with a 190% increase in reaction time.\(^{(14)}\) Conversely, no analgesic activity was noted in the hotplate test on rats given an ethanolic extract, but the same extract did reduce the writhing response to phenylquinone after oral (1 g/kg) and intraperitoneal (500 mg/kg) treatment.\(^{(18)}\)

The isolectins isolated from the rhizome are reported to cause nonspecific agglutination of erythrocytes, to induce the synthesis of interferon by human lymphocytes,\(^{(6,7)}\) and have carbohydrate–binding properties.\(^{(6,7)}\)

An extract of nettle at a concentration of 1.2 mg/mL has been reported to be active against L-1210 leukaemic cells in mice.\(^{(22)}\)

**Clinical studies**

**Diuretic effect**

In an open, uncontrolled study, 32 patients with myocardial or chronic venous insufficiency were treated with 15 mL of nettle juice three times daily for two weeks.\(^{(G52)}\) A significant increase in daily volume of urine was observed throughout the study, the volume by day 2 being 9.2% \((p < 0.0005)\) higher than the baseline value in patients with myocardial insufficiency and 23.9% higher than the baseline value \((p < 0.0005)\) in those with chronic venous insufficiency. It has been proposed that the diuretic activity of aqueous extracts of nettle may be attributed to the high potassium content.\(^{(19)}\) The reputed diuretic effects of nettle require further investigation.

**Arthritis and rheumatism**

An open, uncontrolled multicentre study involving 152 patients with various, mainly degenerative, rheumatic conditions reported that 70% of participants experienced symptom relief by the end of the three–week treatment period.\(^{(G52)}\) In an open, randomised pilot study involving 37 patients with acute arthritis, diclofenac 50 mg plus stewed nettle herb 50 g was compared with diclofenac 200 mg.\(^{(23)}\) Assessment was based on the decrease in elevated acute phase C-reactive protein serum concentrations, and clinical signs of acute arthritis. Clinical improvement was observed in both groups to a similar extent. On the basis of the findings, it was suggested that nettle herb administration may enhance the effectiveness of diclofenac in rheumatic conditions. However, this requires further investigation.

Postmarketing surveillance studies involving a total of almost 2000 patients with rheumatoid arthritis treated for three weeks with nettle leaf extract (IDS-23) administered as an adjuvant to non–steroidal anti–inflammatory drugs (NSAIDs), or as monotherapy, have reported that the extract was well–
In a randomised, double-blind, crossover study, 27 patients with osteoarthritis pain at the base of the thumb and index finger, received stinging nettle leaf (applied for 30 seconds daily for one week to the painful area) or white dead nettle (*Lamium album*) as placebo, followed by a five-week wash-out period before crossing to the other arm of the study. The results indicated that reductions in visual analogue scale scores for pain and in a health assessment questionnaire score for disability were significantly better for the stinging nettle group, compared with the placebo group ($p = 0.026$ and $p = 0.0027$ for pain and disability, respectively).

**Benign prostatic hyperplasia**

Clinical studies of nettle preparations in the treatment of symptoms of benign prostatic hyperplasia (BPH) have been reviewed. Information from this review is summarised below.

Several uncontrolled trials have reported improvements in urological symptoms, compared with baseline values, following administration of nettle root extract (5:1) 600–1200 mg daily for three weeks to 20 months. Large observational studies involving patients with BPH who received nettle root extract for two to three months have reported improvements in various symptoms, such as urinary frequency, urinary flow and nocturia. These studies provide justification for further, rigorous investigation of the effects of nettle in BPH.

A placebo-controlled trial involving 79 patients with BPH assessed the effects of nettle root extract 600 mg daily for six to eight weeks. Compared with placebo, nettle root extract administration resulted in greater improvements in urinary flow and urine volume and residual volume. Another placebo-controlled trial of nettle root extract 600 mg daily for nine weeks in men with BPH ($n = 50$) reported a significant decrease in SHBG concentrations and significant improvement in micturition volume and maximum urinary flow.

**Rhinitis**

A randomised, double-blind, placebo-controlled study assessed the effects of a freeze-dried preparation of nettle herb in individuals with allergic rhinitis. Participants received nettle herb 600 mg, or placebo, at the onset of symptoms over a one-week period. Assessment was based on daily symptom diaries and global responses recorded at follow-up visits after one week of therapy. Nettle herb was rated more highly than placebo in the global assessment, but was rated less highly on the basis of data from the symptom
diaries. It was concluded that there should be further investigation with a larger sample size and involving a longer treatment period.
Side-effects, Toxicity

Consumption of nettle tea has caused gastric irritation, a burning sensation of the skin, oedema and oliguria.\(^{(G22)}\) The leaves are extremely irritant in view of their acetylcholine- and histamine–containing glandular hairs. An LD\(_{50}\) in mice following intraperitoneal administration of nettle has been reported as 3.625 g/kg.\(^{(12)}\) The LD\(_{50}\) for intravenous infusion of nettle leaf in mice has been documented as 1.92 g/kg, and the LD\(_{50}\) for chronic administration in rats has been stated as 1.31 g/kg.\(^{(G50)}\) An ethanolic extract of nettle (plant part unspecified) showed low toxicity in rats and mice after oral and intraperitoneal administration at doses equivalent to 2 g/kg.\(^{(18)}\)
Contra-indications, Warnings

In view of the documented pharmacological actions for nettle, excessive use may interact with concurrent therapy for diabetes, high or low blood pressure, and may potentiate drugs with CNS-depressant actions. Gastrointestinal irritation has been documented.

Pregnancy and lactation
Nettle is reputed to be an abortifacient and to affect the menstrual cycle.\(^{(G30)}\) Utero–activity has been documented in animal studies. In view of this, the use of nettle during pregnancy should be avoided. Excessive use is best avoided during lactation.
The chemistry of nettle is well documented. Limited pharmacological data are available to support the traditional herbal uses although hypoglycaemic activity *in vivo* has been reported. A number of clinical trials have provided some evidence to support the diuretic and anti-inflammatory effects of nettle, and for the effects of nettle in relief of symptoms of allergic rhinitis. Clinical evidence exists to support the efficacy of root extracts in the treatment of benign prostatic hyperplasia. However, further well-designed clinical trials of nettle involving large numbers of patients are required to establish the benefits. Irritant properties have been documented for nettle and excessive use should be avoided.
References

See also General References G2 G3 G5 G6 G9 G16 G22 G30 G31 G32 G36 G37 G43 G50 G52 G54 G56 G64.

16. Oliver-Bever B, Zahland GR. Plants with oral hypoglycaemic activity. *Q J


Species (Family)

Petroselinum crispum (Mill.) Nyman (Apiaceae/Umbelliferae)
Synonym(s)

*Apium petroselinum* L., *Carum petroselinum* (L.) Benth., *Petroselinum sativum* Hoffm.
Part(s) Used

Leaf, root, seed
Pharmacopoeial and Other Monographs

BHC 1992\textsuperscript{(G6)}

BHP 1996\textsuperscript{(G9)}

Complete German Commission E\textsuperscript{(G3)}

Martindale 33rd edition\textsuperscript{(G67)}

PDR for Herbal Medicines 2nd edition\textsuperscript{(G36)}
Legal Category (Licensed Products)

GSL$^{(G37)}$
Constituents


**Flavonoids**
Glycosides of apigenin, luteolin (e.g. apiin, luteolin–7–apiosyl–glucoside, apigenin–7–glucoside (leaf only), luteolin–7–diglucoside (leaf only)).

**Furanocoumarins**
Bergapten and oxypeucedanin as major constituents (up to 0.02% and 0.01% respectively); also 8–methoxypsoralen, imperatorin, iso imperatorin, isopimpinellin, psoralen, xanthotoxin (up to 0.003%). \(^{(1)}\)

**Volatile oils**
2–7% in seed, 0.05% in leaf. The seed contains apiol, myristicin, tetramethoxyallylbenzene, various terpene aldehydes, ketones, and alcohols. The leaf contains myristicin (up to 85%), apiol, 1,3,8-p-menthatriene, 1–methyl-4–isopropenylbenzene, methyl disulfide, monoterpenes (e.g. α- and β-pinene, β-myrcene, β-ocimene, β-phellandrene, p-terpinene, α-terpineol), sesquiterpenes (e.g. α-copaene, carotol, caryophyllene).

**Other constituents**
Fixed oil, oleo–resin, proteins, carbohydrates, and vitamins (especially vitamins A and C).

A detailed vitamin and mineral analysis is given elsewhere. \(^{(G22)}\)
Food Use

Parsley is listed by the Council of Europe as natural source of food flavouring (category N2). This category indicates that parsley can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\textsuperscript{(G16)} Parsley is commonly used in foods. In the USA, parsley is listed as GRAS (Generally Recognised As Safe).\textsuperscript{(G65)}
Herbal Use

Parsley is stated to possess carminative, antispasmodic, diuretic, emmenagogue, expectorant, antirheumatic and antimicrobial properties. Traditionally, it has been used for flatulent dyspepsia, colic, cystitis, dysuria, bronchitic cough in the elderly, dysmenorrhoea, functional amenorrhoea, myalgia and specifically for flatulent dyspepsia with intestinal colic. (G2 G6 G7 G8 G64)
Dosage

*Leaf/root*
2–4 g or by infusion.

*Seed*
1–2 g.

*Dried root*
2–4 g or by infusion three times daily.\(^{(G6 \ G7)}\)

*Liquid extract*
2–4 mL (1:1 in 25% alcohol) three times daily.\(^{(G6 \ G7)}\)
Pharmacological Actions

In vitro and animal studies

Parsley extract (0.25–1.0 mL/kg, by intravenous injection) has been reported to lower the blood pressure of cats by more than 40%,\(^{(2)}\) and to decrease both respiratory movements and blood pressure in anaesthetised dogs.\(^{(3)}\) Parsley exhibits a tonic effect on both intestinal and uterine muscle.\(^{(3)}\) This uterine effect has been attributed to the apiol content,\(^{(G30)}\) but has also been observed with apiole–free aqueous extracts.\(^{(3)}\) An aqueous extract of parsley has been documented to contain an antithiamine substance which was unaffected by cooking or contact with gastric juice.\(^{(3)}\) Myristicin and apiol are both effective insecticides.\(^{(4)}\)

Parsley seed oil has been reported to stimulate hepatic regeneration.\(^{(5)}\)

Clinical studies

Myristicin is the hallucinogenic principle present in nutmeg seed. It has been hypothesised that myristicin is converted in the body to amfetamine, to which it is structurally related.\(^{(4)}\) Myristicin has a structural similarity with sympathomimetic amines and it is thought that it may compete for monoamine oxidase enzymes, thereby exhibiting a monoamine oxidase inhibitor (MAOI)-like action.\(^{(6)}\) Parsley oil has been included in the diet of pregnant women and is reported to increase diuresis, and plasma protein and plasma calcium concentrations.\(^{(4)}\)

The diuretic effect associated with the consumption of parsley is probably attributable to the pharmacological activities of myristicin (sympatho mimetic action) and apiol (irritant effect).
Side-effects, Toxicity

Chronic and excessive consumption of fresh parsley (170 g daily for 30 years) has been associated with generalised itching and pigmentation of the lower legs in a 70-year-old woman. The symptoms were attributed to excessive ingestion of parsley in the presence of chronic liver disease. The aetiology of the chronic hepatitis was unknown, but considered possibly related to the chronic exposure to the psoralen constituents in parsley. Apiole and myristicin are also documented to be hepatotoxic.

The ingestion of approximately 10 g apiole has been reported to cause acute haemolytic anaemia, thrombocytopenia purpura, nephrosis and hepatic dysfunction. However, ingestion of 10 g of apiole would require a dose of more than 200 g parsley. The amount of apiole ingested as a result of normal dietary consumption of parsley is not hazardous. Myristicin has been documented to cause giddiness, deafness, hypotension, decrease in pulse rate, and paralysis, followed by fatty degeneration of the liver and kidney. In addition, myristicin is known to possess hallucinogenic properties. However, when compared to nutmeg, parsley contains a relatively low concentration of myristicin (less than 0.05% in parsley leaf, about 0.4–0.89% in nutmeg); parsley seed is potentially hazardous in view of its higher volatile oil content (about 2–7%) which contains apiole and myristicin.

Parsley contains phototoxic furanocoumarins (see Celery). However, photodermatitis resulting from the oral ingestion of parsley is thought to be unlikely. The ingestion of 50 g parsley provides negligible amounts of bergapten (0.5–0.8 g). The concentration of oxypeucedanin provided was not mentioned. However, a photoactive reaction from topical contact with parsley is possible.

Apiole is an irritant component of the volatile oil and may cause irritation of the kidneys during excretion.

Parsley seed oil has been reported to stimulate hepatic regeneration. Myristicin and apiole are documented to have a similar chemical structure and acute toxicity to safrole, which is known to be carcinogenic and hepatotoxic (see Sassafras). The carcinogenic potential of apiole and myristicin has not been evaluated.

LD$_{50}$ (mice, intravenous injection) values for apiole and myristicin have been documented as 50 mg/kg and 200 mg/kg body weight, respectively.
Contra-indications, Warnings

Parsley should not be ingested in excessive amounts in view of the documented toxicities of apiol and myristicin. Parsley may cause a photoactive reaction, especially following external contact, may aggravate existing renal disease, and may potentiate existing MAOI therapy.

Pregnancy and lactation

Parsley is reputed to affect the menstrual cycle.\(^{(G7)}\) Utero-activity has been documented in humans and animals,\(^{(G30)}\) and parsley is stated to be contra-indicated during pregnancy.\(^{(G49 G58)}\) Myristicin has been reported to cross the placenta and can lead to foetal tachycardia.\(^{(8)}\) In view of this, parsley should not be taken during pregnancy and lactation in doses that greatly exceed the amounts used in foods.
Pharmaceutical Comment

Parsley is commonly consumed as part of the diet. The pharmacological and toxicological properties of parsley are primarily associated with the volatile oil, particularly the apiol, myristicin and furano coumarin constituents. Most of the reported uses of parsley are probably due to the volatile oil; no documented information was located regarding antirheumatic and antimicrobial properties. Parsley should not be consumed in doses that greatly exceed the amounts used in foods, as excessive ingestion may result in apiol and myristicin toxicity.
References


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Species (Family)

*Aphanes arvensis* L. (Rosaceae)
Synonym(s)

*Alchemilla arvensis* Scop., *Aphanes*
Part(s) Used
Herb
Pharmacopoeial and Other Monographs

BHP 1983\(^{(G7)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL$^{(G37)}$
**Constituents**

See General References G7 G34 G64.

Limited information is available. A related species *Alchemilla vulgaris* (lady’s mantle) is reported to contain 6–8% tannins (hydrolysable–type);\(^{(G41)}\) none have been documented for parsley piert, although it is stated to contain an astringent principle.\(^{(G6)}\)
Food Use

Parsley piert is not used in foods.
Herbal Use

Parsley piert is stated to possess diuretic and demulcent properties, and to dissolve urinary deposits. Traditionally, it has been used for kidney and bladder calculi, dysuria, strangury, oedema of renal and hepatic origin, and specifically for renal calculus. (G7 G64)
**Dosage**

**Dried herb**
2–4 g or by infusion three times daily.\(^{(G7)}\)

**Liquid extract**
2–4 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)

**Tincture**
2–10 mL (1 : 5 in 45% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

None documented.
Side-effects, Toxicity

None documented.
Contra-indications, Warnings

None documented.

**Pregnancy and lactation**

In view of the lack of phytochemical, pharmacological, and toxicity information, the use of parsley piert during pregnancy and lactation should be avoided.
Pharmaceutical Comment

Little chemical information is available on parsley piert. No scientific evidence was found to justify the herbal uses. Parsley piert may exhibit astringent actions. In view of the lack of toxicity data, excessive use of parsley piert should be avoided.
References

See General References G7 G31 G34 G36 G37 G64.
Passionflower
Species (Family)

Passiflora incarnata L. (Passifloraceae)
Synonym(s)

Apricot Vine, Grenadille, Maypop, Passiflora, Passion Vine
Part(s) Used

Herb
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

BP 2002\(^{(G71)}\)

Complete German Commission E\(^{(G3)}\)

ESCOP 1997\(^{(G52)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)

Ph Eur 2004\(^{(G72)}\)
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents

See General References G2 G6 G22 G41 G64.

Alkaloids
Indole-type. Harman (major), harmaline, harmalol, harmine and harmol have been reported. Of 17 different samples examined, only one contained alkaloids with a possible harmine content of 0.1 ppm.\(^{(1)}\)

Flavonoids
Pharmacopoeial standard, not less than 1.5\%.\(^{(2-7)}\) Vitexin, isovitexin and their C-glycosides, apigenin, luteolin glycosides (e.g. orientin, homoorientin and lucenin); kaempferol, quercetin and rutin.

Other constituents
Maltol and ethylmaltol (γ-pyrone derivatives), passicol (a polyacetylene), fatty acids (e.g. linoleic acid, linolenic acid, myristic acid, palmitic acid and oleic acid), formic acid, butyric acid, sitosterol, stigmasterol, sugars and gum.

Other plant parts
Coumarins (scopoletin and umbelliferone) are found in the root.

Other Passiflora species
Cyanogenetic glycosides passibiflorin, epipassibiflorin and passitrifasciatin (Passiflora biflora, Passiflora talamancensis, Passiflora trifasciata), linamarin and lotaustralbin (Passiflora lutea)\(^{(10)}\) and prunasin (Passiflora edulis).\(^{(11)}\)
Passionflower is listed by the Council of Europe as a natural source of food flavouring (category N3). This category indicates that passionflower can be added to foodstuffs in the traditionally accepted manner, but that there is insufficient information available for an adequate assessment of potential toxicity. In the USA, passionflower is permitted for use in food.
Passionflower is stated to possess sedative, hypnotic, antispasmodic and anodyne properties. Traditionally, it has been used for neuralgia, generalised seizures, hysteria, nervous tachycardia, spasmodic asthma, and specifically for insomnia. The German Commission E approved internal use for nervous restlessness. Passionflower is used in combination with valerian root and lemon balm for conditions of unrest, difficulty in falling asleep due to nervousness. Passionflower is used extensively in homeopathy.
Dosage

**Dried herb**
0.25–1.0 g or by infusion three times daily; (G6 G7) 0.5–2 g three or four times daily; (G52) 4–8 g daily. (G3)

**Liquid extract**
0.5–1.0 mL (1 : 1 in 25% alcohol) three times daily. (G6 G7)

**Tincture**
0.5–2.0 mL (1 : 8 in 45% alcohol) three times daily. (G6 G7)
Pharmacological Actions

**In vitro and animal studies**

CNS sedation, potentiation of hexobarbitone–induced sleeping time, anti-convulsant activity (at high doses) and a reduction in spontaneous motor activity (at low doses) have been documented for maltol and ethylmaltol in mice.\(^{12,13}\) Subsequent research documenting similar activities in mice was unable to attribute the observed activities to either flavonoid or alkaloid components present in the extract tested.\(^{14}\) An aqueous ethanolic extract of passionflower administered intraperitoneally to rats (160 mg/kg) prolonged sleeping time induced by pentobarbital (50–4000 mg/kg) and reduced spontaneous locomotor activity.\(^{14}\) The same extract (160 mg/kg, intraperitoneal or oral administration) raised the threshold to nociceptive stimuli in tail flick and hotplate tests.\(^{14}\)

Rats showed reduced activity in a one–arm radial maze test after one week of daily oral administration of an aqueous ethanolic extract of passionflower (10 mg/kg).\(^{G52}\) In mice, a 70% ethanol extract (1000 mg/kg, intraperitoneally) administered 10 minutes prior to sodium pentobarbital (40 mg/kg, intraperitoneally) resulted in a significant prolongation (40%) of sleeping time.\(^{15}\) When given by gastric tube 1 hour prior to amfetamine (5 mg/kg, subcutaneous), the same extract (500 mg/kg) caused significant reduction in hypermotility. Oral treatment of mice with an ethanol extract of passionflower (25 and 50 mg/kg) reduced exploratory and spontaneous motor activities, prolonged sleeping time induced by pentobarbital, and inhibited aggressiveness and restlessness caused by amfetamines.\(^{G52}\) The sedative activity was comparable with that of meprobamate (250 mg/kg), and greater than that of diazepam (10 mg/kg) and chlordiazepoxide (10 mg/kg). In another study in mice, sedative action, as assessed by prolongation of hexobarbital–induced sleeping time, was decreased by a 30% aqueous ethanol extract of passionflower (1.75 mg/kg, orally).\(^{16}\)

A dry extract of passionflower (800 mg/kg, orally) containing 2.6% flavonoids resulted in a significant \((p < 0.01)\) anxiolytic effect, as assessed by prolongation of hexobarbital–induced sleeping time, whereas locomotor activity remained unaffected.\(^{G52}\)

It has been suggested that the sedative effects of maltol and ethylmaltol mask the stimulant actions of harman alkaloids.\(^{12}\) The CNS-depressant effect exhibited by *P. edulis* has been attributed to alkaloid and flavonoid compounds\(^{17}\) and to a protein–like substance.\(^{18}\) The pharmacological evidence generally supports sedative and anxiolytic effects of passionflower,
although there are conflicting results. It is not clear which constituents are the active principles, and clinical data are lacking.\textsuperscript{(19)} Maltol is reportedly an artefact and not a relevant constituent.\textsuperscript{(19)}

Passicol exhibits antimicrobial activity towards a wide variety of moulds, yeasts and bacteria.\textsuperscript{(8)} Group A haemolytic streptococci are stated to be more susceptible than \textit{Staphylococcus aureus}, with \textit{Candida albicans} of intermediate susceptibility.\textsuperscript{(8)}
In mice, the acute toxicity of a fluid extract of passionflower (intraperitoneal injection) was stated as being greater than 900 mg/kg.\(^{(14)}\) In rats, subacute oral treatment with an aqueous ethanol extract of passionflower 10 mg/kg for 21 days showed no changes in weight, rectal temperature or motor coordination.\(^{(G52)}\)

Cyanogenic glycosides have been documented for related *Passiflora* species.
Contra-indications, Warnings

Excessive doses of passionflower may cause sedation.

**Pregnancy and lactation**
No other data regarding the use of passionflower during pregnancy or lactation were located. In view of this, excessive use of passionflower during pregnancy and lactation should be avoided.
Pharmaceutical Comment

The active constituents have not been clearly identified. CNS-sedative properties have been documented in animals, thus providing some data to support some of the traditional uses of passionflower. However, well–designed clinical trials assessing the reputed sedative properties of passionflower are lacking. In view of the lack of toxicity data, excessive use of passionflower should be avoided.
References


15. Capasso A, Pinto A. Experimental investigations of the synergistic–


Pennyroyal
Species (Family)

i. *Mentha pulegium* L. (Labiatae)

Synonym(s)

Pulegium

i. European Pennyroyal

ii. American Pennyroyal
Part(s) Used
Herb
Pharmacopoeial and Other Monographs

BHP 1983\(^{(G7)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

Pennyroyal is not included in the GSL.\(^{(G37)}\)
Constituents

See General References G22 G48 G64 G58.

**Volatile oils**

1–2%. Pulegone is the principal component (60–90%); others include menthone, *iso*-menthone, 3-octanol, piperitenone and *trans*-iso-pulegone.
Food Use

Pennyroyal is not commonly used in foods. It is listed by the Council of Europe as a natural source of food flavouring (category N3). This category indicates that there is insufficient information available for an adequate assessment of toxicity (but see Side-effects, Toxicity). In the USA, pennyroyal is permitted for use in foods.
Herbal Use

Pennyroyal is stated to possess carminative, antispasmodic, diaphoretic and emmenagogue properties, and has been used topically as a refrigerant, antiseptic and insect repellent. Traditionally, it has been used for flatulent dyspepsia, intestinal colic, common cold, delayed menstruation, and topically for cutaneous eruptions, formication and gout.\(^{(G7)}\)
Dosage

*Herb*
1–4 g or as infusion three times daily.\(^{(G7)}\)

*Liquid extract*
1–4 mL (1:1 in 45% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

None documented.
The toxicity of pennyroyal oil is well recognised and human fatalities following its ingestion as an abortifacient have been reported.\(^1–3\) Symptoms reported following ingestion of the oil include abdominal pain, nausea, vomiting, diarrhoea, lethargy and agitation, pyrexia, raised blood pressure and pulse rate, and generalised urticarial rash. Generally, doses required for an abortifacient effect are also toxic and fatalities have involved both nephrotoxicity and hepatotoxicity.\(^2–4\) Doses of one ounce and 30 mL\(^1–3\) have proved fatal, whereas individuals have recovered following unsuccessful abortion attempts involving the ingestion of 7.5 mL oil.\(^3\) The mechanism of hepatotoxicity for pennyroyal is not known.\(^2\) A direct hepatoxic action has been suggested for the ketone component, pulegone.\(^2\) Alternatively, metabolic conversion of pulegone to a reactive inter mediate, a furan or epoxide, has been proposed.\(^2\)

Acute \(LD_{50}\) values for pennyroyal oil are documented as 0.4 g/kg (oral, rats) and 4.2 g/kg (dermal, rabbits).\(^4\) The oil is non- or moderately irritating, non–sensitising and non–phototoxic.\(^4\) Acute \(LD_{50}\) values documented for pulegone, the principal oil component, are, not suprisingly, similar to those for the oil: 0.47 g/kg (oral, rats), 3.09 g/kg (dermal, rabbits).\(^5\) Steroid (pregnenolone–16\(\alpha\)-carbonitrile) treatment has reduced hepatotoxicity observed in female rats fed pulegone, whereas triamcinolone has increased it.\(^5\) Toxicity of pulegone is unaffected by partial hepatectomy or ligation of the common bile duct, while partial nephrectomy intensified toxicity.\(^5\)
Contra–indications, Warnings

Pennyroyal oil is irritant and instances of hepato toxicity and nephrotoxicity have been documented following its ingestion. Both the internal and external use of pennyroyal oil has been contra–indicated. (G58)

Pregnancy and lactation

Pennyroyal is contra– indicated in pregnancy. (G7) Traditionally, it has been employed as an abortifacient, this use probably resulting from the irritant action of the oil on the genito–urinary tract. Fatalities have resulted from the doses of oil required to exert an abortifacient effect.
Pharmaceutical Comment

Interest in pennyroyal has focused on the toxicity associated with the volatile oil. No documented reports of the pharmacological actions exhibited by the herb were located. Pennyroyal herb teas have been reported to be used without side-effects,\(^{(2)}\) presumably due to lower amounts of oil ingested. In view of its potential toxicity, excessive ingestion of the oil should be avoided. Pennyroyal oil is not suitable for internal or external use.
References

See also General References G7 G19 G22 G31 G32 G36 G37 G43 G48 G58 G64.


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Species (Family)

*Ranunculus ficaria* L. (Ranunculaceae)
Synonym(s)

Ficaria, *Ficaria ranunculoides Moench.*, Lesser Celandine, Ranunculus
Part(s) Used

Herb
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL$^{(G37)}$
Constituents
See General References G40 G42 G64.

**Lactones**
Anemonin (dimer), protoanemonin (precursor to anemonin).

**Triterpenoids**
Glycosides based on the sapogenins hederagenin and oleanolic acid, with arabinose, glucose and rhamnose, as sugar moieties.\(^1\)

**Other constituents**
Tannin and ascorbic acid (vitamin C).
Food Use

Pilewort is not used in foods.
Herbal Use

Pilewort is stated to possess astringent and demulcent properties. Traditionally, it has been used for haemorrhoids, and specifically for internal or prolapsed piles with or without haemorrhage, by topical application as an ointment or a suppository.\textsuperscript{G7 G64}
Dosage

*Dried herb*
2–5 g or by infusion three times daily.\(^{(G7)}\)

*Liquid extract*
2–5 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)

*Ointment*
3% in a suitable basis.

*Pilewort Ointment*
(BPC 1934) 30% fresh herb in benzoinated lard.
**Pharmacological Actions**

*In vitro* and animal studies

Local antihaemorrhoidal activity has been documented for the saponin constituents.\(^1\) Antibacterial and antifungal properties have been documented for both anemonin and protoanemonin, although anemonin is reported to exhibit much weaker activity.\(^{G33\, G48}\)

The reported presence of tannin constituents\(^{G42}\) supports the reputed astringent activity of pilewort, although no pharmacological studies were located.
Side–effects, Toxicity

The sap of pilewort is stated to be irritant.\(^{(G51)}\) Protoanemonin is stated to be an acrid skin irritant, although it is readily converted into the inactive dimer anemonin.\(^{(G33)}\) Protoanemonin is stated to have a marked ability to combine with sulfhydryl (-SH) groups and it is thought that the toxic subdermal properties of protoanemonin may depend on the inactivation of enzymes containing -SH groups.\(^{(G33)}\) An LD\(_{50}\) value (mice, intraperitoneal injection) for anemonin has been reported as 150 mg/kg body weight.\(^{(G48)}\)
Contra–indications, Warnings

Pilewort is not recommended for internal consumption.\(^{(G49)}\) Topical use of pilewort may cause irritant skin reactions.

**Pregnancy and lactation**
The safety of pilewort has not been established. It is not recommended for internal consumption;\(^{(G49)}\) in view of this and the potential irritant action, the use of pilewort during pregnancy and lactation is best avoided.
Pharmaceutical Comment

Limited information is available on the chemistry of pilewort. Little scientific information was located to justify the herbal uses, although antihaemorrhoidal activity has been documented for the saponin constituents. In view of the toxic and irritant properties stated for protoanemonin, the excessive use of pilewort is not advisable.
References

See also General References G9 G10 G33 G36 G37 G40 G42 G48 G49 G57 G64.


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Plantain
Species (Family)

Plantago major L. (Plantaginaceae)
Synonym(s)
Common Plantain, General Plantain, Greater Plantain
Part(s) Used

Leaf
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)

Complete German Commission E\(^{(G3)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

Plantain is not included in the GSL.\(^{(G37)}\)
Constituents
See General References G2 G22 G40 G51 G62 G64.

Acids
Benzoic acid, caffeic acid, chlorogenic acid, cinnamic acid, \( p \)-coumaric acid, ferulic acid, fumaric acid, gentisic acid, \( p \)-hydroxybenzoic acid, neochlorogenic acid, salicylic acid, syringic acid, ursolic acid, vanillic acid;\(^1\)\(^2\) oleanolic acid and ascorbic acid.

Alkaloids
Trace (unspecified),\(^3\)\(^4\) boschniakine and the methyl ester of boschniakinic acid\(^5\)

Amino acids
DL-\( \alpha \)-Alanine, asparagine, L-histidine, DL-lysine, DL-leucine, serine and tryptophan.\(^6\)

Carbohydrates
L-Fructose, D-glucose, planteose, saccharose, stachyose, \( d \)-xylose, sorbitol, tyrosol, mucilage and gum.\(^7\)

Flavonoids
Apigenin, baicalein, scutellarein, baicalin, homoplantaginin, nepitrin, luteolin, hispidulin and plantagoside.\(^8\)\(^9\)\(^10\)

Iridoids
Aucubin, aucubin derivatives, plantarenaloside, aucuboside and melitoside.\(^5\)\(^11\)\(^12\)

Tannins
4\%. Unspecified.

Other constituents
Choline, allantoin, invertin and emulsin (enzymes), fat 10–20\%, resin, saponins, steroids\(^13\) and thioglucoside.
Food Use

Plantain leaf is not used in foods. A related species, *Plantago lanceolata* L., is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that *P. lanceolata* can be added to foodstuffs in small quantities, with a possible limitation of an active constituent (as yet unspecified) in the final product. In the USA, plantain is listed by the Food and Drugs Administration (FDA) as a Herb of Undefined Safety.
Herbal Use

Plantain is stated to possess diuretic and antihaemorrhagic properties. Traditionally, it has been used for cystitis with haematuria, and specifically for haemorrhoids with bleeding and irritation. (G2 G7 G42 G64)
Dosage

*Dried leaf*
2–4 g or by infusion three times daily.\(^{(G7)}\)

*Liquid extract*
2–4 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)

*Tincture*
2–4 mL (1 : 5 in 45% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

In vitro and animal studies

An aqueous extract has been reported to possess bronchodilatory activity in guinea–pigs. It was more effective against acetylcholine–induced contraction, than towards constriction induced by histamine or serotonin.\(^{(14)}\) The bronchodilatory activity of plantain in guinea–pigs has been reported to be less active and of shorter duration compared to salbutamol or atropine.\(^{(15)}\)

Hypotensive activity in normotensive, anaesthetised dogs has been documented; 125 mg/kg extract was found to decrease arterial blood pressure by 20–40 mmHg.\(^{(16)}\)

An aqueous extract, reported to contain flavonoids, saponins, steroids and alkaloids, was shown to possess anti–inflammatory activity in the rat using various models of inflammation, and a strengthening of capillary vessels has also been documented.\(^{(13)}\) However, an extract was found to exhibit minimal (11%) inhibition of carrageenan–induced rat paw oedema.\(^{(17)}\) Leaf extracts in hexane have shown potent wound–healing activity in rabbits; the effect was primarily attributed to C\(_{26}\)–C\(_{30}\) alcohols present in the extract.\(^{(18)}\) Both the anti–inflammatory and wound–healing activities of plantain have been attributed to the high content of chlorogenic and neochlorogenic acids.\(^{(2)}\)

Aucubin and a haemolytic saponin fraction have exhibited antibiotic activity towards *Micrococcus flavus* and *Staphylococcus aureus* (aucubin only).\(^{(19)}\) Antibacterial activity towards *Bacillus subtilis* has been documented for the fresh plant juice, which was also found to lack activity towards Gram–positive organisms and fungi.\(^{(20)}\) A negative response to cytotoxic, antitumour and antiviral activity was also reported for the plant juice.\(^{(20)}\)

A mild laxative action has been reported in mice administered iridoid glycosides, including aucubin.\(^{(21)}\) Plantain seed is sometimes used as a substitute for ispaghula (a bulk laxative).\(^{(G45)}\)

Plantain has been documented to lower concentrations of total plasma lipids, cholesterol, \(\beta\)-lipoproteins and triglycerides in rabbits with experimental atherosclerosis.\(^{(22)}\) Plantain has been reported to be useful in lowering plasma cholesterol concentrations.\(^{(23)}\)

A tonus–raising effect on isolated guinea–pig and rabbit uterus tissue has been documented for an aqueous extract at a dose of 1–2 mg/cm\(^3\).\(^{(24)}\)
Aucubin has been stated to be the active principle responsible for a hepatoprotective effect documented for plantain.\(^{(25)}\)

**Clinical studies**

Plantain has been reported to be effective in the treatment of chronic bronchitis of a spastic or non–spastic nature.\(^{(14,26,27)}\) A pronounced improvement in both subjective and objective symptoms of the common cold following treatment with plantain has also been reported.\(^{(28)}\) Plantain, in combination with agrimony, German chamomile, peppermint and St. John’s wort, has been documented to provide pain relief in patients with chronic gastroduodenitis.\(^{(29)}\) Following treatment, previously diagnosed erosions and haemorrhagic mucous changes were stated to have disappeared.
Side–effects, Toxicity

Allergic contact dermatitis to plantain has been reported. The green parts of the plant are thought to yield a mustard oil–type of thioglucoside, which releases an irritant principle (isothiocyanate) upon enzymatic hydrolysis. The seed may also cause sensitisation and dermatitis. Plantain is reported to be of low toxicity with LD$_{50}$ values in the rat documented as 1 g/kg (intraperitoneal injection) and greater than 4 g/kg (by mouth).
Contra–indications, Warnings

Plantain may cause a contact allergic reaction; it induces the formation of IgE antibodies, which may cross–react to psyllium.\(^{30}\) Excessive doses may exert a laxative effect and a hypotensive effect.

**Pregnancy and lactation**

*In vitro* uterotonic activity has been documented for plantain. In view of this, excessive use of plantain, which may also exert a laxative effect, should be avoided during pregnancy.
Pharmaceutical Comment

The constituents of plantain are well documented and the reputed antihaemorrhagic properties are probably attributable to the tannin constituents. In addition, bronchospastic activity has been documented in both animal and human studies, and may warrant further research. The toxicity of plantain is reported to be low but excessive ingestion should be avoided. The bulk laxative ispaghula consists of the dried seeds of related species *Plantago psyllium*, *P. ovata* and *P. indica.*

(G45)
References

See also General References G2 G3 G9 G16 G22 G31 G36 G37 G40 G42 G43 G51 G62 G64.


Species (Family)

Asclepias tuberosa L. (Asclepiadaceae)
Synonym(s)

Asclepias
Part(s) Used

Root
Pharmacopoeial and Other Monographs

BHP 1983\(^{G7}\)

Martindale 33rd edition\(^{G67}\)

PDR for Herbal Medicines 2nd edition\(^{G36}\)
Legal Category (Licensed Products)

GSL**(G37)**
Constituents

See General References G48 G64.

Little chemical information is available for pleurisy root. Cardiac glycosides of the cardenolide type (e.g. afroside, asclepin, calactin, calotropin, gomphoside, syriogenin, syrioside, uscharidin, uscharin, uscharin and uzarigenin) have been documented for many *Asclepias* species,\(^{(1-4)}\) including *A. tuberosa*.\(^{(5)}\) Concentrations of cardiac glycosides are reported to vary between *Asclepias* species\(^{(1)}\) and individual plant parts,\(^{(4)}\) in descending order of latex, stem, leaf and root.\(^{(6)}\)

No other data regarding constituents of the root were located.

**Other plant parts**

Constituents documented for the herb include flavonols (e.g. kaempferol and quercetin) and flavonol glycosides (e.g. rutin and isorhamnetin), amino acids, caffeic acid, chlorogenic acid, choline, carbohydrates (e.g. glucose, fructose and sucrose), β-sitosterol, triterpenes (e.g. α-amyrin and β-amyrin, lupeol, friedelin, viburnitol), volatile oil and resin.\(^{(7,8,G48)}\)
Food Use

Pleurisy root is not used in foods.
Herbal Use

Pleurisy root is stated to possess diaphoretic, expectorant, antispasmodic and carminative properties. It has been used for bronchitis, pneumonitis, influenza, and specifically for pleurisy. (G7 G42 G64)
Dosage

**Dried root**
1–4 g or by infusion three times daily.\(^{(G7)}\)

**Liquid extract**
1–4 mL (1:1 in 45% alcohol) three times daily.\(^{(G7)}\)

**Tincture**
1–5 mL (1 : 10 in 45% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

In vitro and animal studies

Low doses of extracts of *Asclepias* species including *A. tuberosa* have been documented to cause uterine contractions (*in vivo*) and to exhibit oestrogenic effects.\(^{5,9,10,G30}\) No effect was observed on blood pressure or respiration (*in vivo*), or on the isolated heart (frog, turtle).\(^{9}\) Various activities have been reported for related *Asclepias* species. A positive inotropic action (*in vivo* and *in vitro*) has been reported for asclepin (*Asclepias curassavica*), which was found to be more potent, longer acting and with a wider safety margin when compared with other cardiac glycosides (including digoxin).\(^{11–13}\) Asclepin was also reported to exhibit a more powerful activity towards weak cardiac muscle.\(^{13}\) Plant extracts of *A. curassavica, Asclepias engelmanniana* and *Asclepias glaucescens* have exhibited a stimulatory effect on the mammalian CNS, causing an increase in serotonin and noradrenaline concentrations.\(^{14}\)

Antitumour/cytotoxic activities have been documented for *A. albicans* and were attributed to various cardenolide constituents.\(^{15}\)
Side-effects, Toxicity

Pleurisy root and other *Asclepias* species have been documented to cause dermatitis; the milky latex is reported to be irritant.\(^{\text{G51}}\) Large doses may cause nausea, vomiting and diarrhoea.\(^{\text{G7 G42}}\) Various *Asclepias* species, including *A. tuberosa*, are known to be toxic to livestock, with cardenolides implicated as the toxic constituents.\(^{1,5}\) Toxic effects on the lungs, gastrointestinal tract, kidneys, brain and spinal cord have been observed in rats and rabbits following intravenous administration of an alcoholic extract.\(^{10}\)

Toxicity studies involving related *Asclepias* species have also been documented. The cardenolide fraction of *Asclepias eriocarpa* is reported to contain toxic principles. The whole plant, plant extracts, an isolated and purified cardenolide (labriformin) and digoxin were all found to show qualitatively similar signs of toxicity and gross pathology in sheep and guinea–pigs.\(^{16}\) LD\(_{50}\) values (mice, intraperitoneal injection) for cardenolides obtained from *A. curassavica* and *A. eriocarpa* were all estimated at less than 50 mg/kg body weight. Asclepin (*A. curassavica*) was reported to be safe following a three–month toxicity study in rats, using doses of 0.8, 8 and 20 mg/kg (route unspecified).\(^{13}\) Asclepin has also been documented to have a wider margin of safety than digoxin\(^{11–13}\) (see *In vitro* and animal studies).

Studies in cats have reported asclepin to be less cumulative compared to digoxin.\(^{13}\)
Contra–indications, Warnings

Pleurisy root may interfere with existing cardiac drug therapy. Excessive doses of pleurisy root may interfere with drug therapies that affect amine concentrations in the brain (e.g. antidepressants) and with hormonal therapy.

Pregnancy and lactation

Uterotonic activity (*in vivo*) has been reported for pleurisy root.\(^{(5,G30)}\) In view of this and the potential toxicity of pleurisy root, it is best avoided during pregnancy or lactation.
Pharmaceutical Comment

The chemistry of pleurisy root is poorly documented, but phytochemical studies on pleurisy root and related *Asclepias* species have identified many cardiac glycoside constituents. No scientific evidence was found to justify the herbal uses. In view of the potential toxicity of pleurisy root, excessive use is not recommended.
References

See also General References G7 G30 G36 G37 G42 G48 G51 G64.

16. Benson JM et al. Comparative toxicology of cardiac glycosides from the

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Pokeroo
Species (Family)

*Phytolacca americana* L. (Phytolaccaceae)
Synonym(s)

*Phytolacca decandra* L., Pocan, Pokeweed, Red Plant
Part(s) Used

Root
Pharmacopoeial and Other Monographs

BHP 1996\textsuperscript{(G9)}

Martindale 33rd edition\textsuperscript{(G67)}

Mills and Bone\textsuperscript{(G50)}

PDR for Herbal Medicines 2nd edition\textsuperscript{(G36)}
Legal Category (Licensed Products)

GSL (G37)
Constituents

See General References G22 G48 G64.

Alkaloids
Betalain-type. Betanidine, betanine, iso betanine, isobetanidine, isoprebetanine, phytolaccine and prebetanine.

Lectins
Pokeweed mitogen (PWM) consisting of five glycoproteins Pa$^{-1}$ to Pa$^{-5}$.

Saponins
Triterpenes – phytolaccosides A-1, D$_2$, and O,$^{(1-3)}$ aglycones include phytolaccagenin, jalogonic acid, phytolaccagenic acid, aesculentic acid,$^{(2,4-6)}$ acinosolic acid methyl ester;$^{(5)}$ monodesmosidic and bidesmosidic compounds with oleanolic acid and phytolaccagenic acids as aglycone in P. dodecandra.$^{(7)}$

Other constituents
Isoamericanin A (neo-lignan),$^{(8)}$PAP, pokeweed antiviral protein)$^{(9)}$, α-spinasterol,$^{(5)}$, histamine and gamma aminobutyric acid (GABA).$^{(10)}$
Food Use

Pokeroot is not commonly used in foods. In the USA, the Herb Trade Association has recommended that pokeroot should not be sold as a herbal beverage or food.\(^{(11)}\)
Herbal Use

Pokeroot is stated to possess antirheumatic, anticatarrhal, mild anodyne, emetic, purgative, parasiticidal and fungicidal properties. Traditionally, it has been used for rheumatism, respiratory catarrh, tonsillitis, laryngitis, adenitis, mumps, skin infections (e.g. scabies, tinea, sycosis, acne), mammary abscesses and mastitis.\(^{(G7 \ G64)}\)
Dosage

*Dried root*
0.06–0.3 g or by decoction three times daily.\(^{(G7)}\)

*Liquid extract*
0.1–0.5 mL (1:1 in 45% alcohol) three times daily.\(^{(G7)}\)

*Tincture*
(BPC 1923) 0.2–0.6 mL.
Pharmacological Actions

**In vitro and animal studies**

Anti-inflammatory activity has been documented for saponin fractions isolated from *P. americana*. Activity comparable or greater than that of cortisone acetate was observed in the carrageenan rat paw oedema test when the extract was administered by intraperitoneal injection. The major aglycone, phytolaccagenin, was reported to exhibit greater activity than glycyrrhetic acid and oleanolic acid, which are both known to be effective in acute inflammation. Oral administration required a six-fold increase in dose for comparable activity. Potency of the saponin extract was reduced to one-eighth of that of cortisone when tested against chronic inflammation (granuloma pouch method). The ED$_{50}$ for saponin and phytolaccagenin fractions against carrageenan-induced oedema in the rat (intraperitoneal injection) has been determined as 15.1 and 26 mg/kg respectively.

Isoamericanin A (a neo-lignan) isolated from the seeds of *P. americana* has been reported to increase prostaglandin I$_2$ (PGI$_2$) production from the rat aorta by up to about 150% at a concentration of $10^{-5}$ and to elicit a moderate inductive effect on the *in vivo* release of PGI$_2$.

Hypotensive properties have been described for a pokeroot extract with the activity attributed to histamine and GABA.

A diuretic effect has been described in rats administered pokeroot extract orally at a dose of 500 mg/kg. The effect was reported to be significantly greater than that observed in the saline-treated group of rats, but less than in the furosemide-treated (150 mg/kg) group.

*In vitro* contraction of the guinea-pig ileum has been described for pokeroor extracts. Activity was attributed to a single active constituent that proved to be heat resistant.

The properties of pokeweed antiviral protein have been reviewed.

Molluscicidal activity against schistosomiasis-transmitting snails and spermicidal activity have been documented for saponin components obtained from the fruits of the related species, *P. dodecandra*. An enzyme located in the seeds has been found to be necessary for molluscicidal activity of *P. dodecandra*. Crushing the seeds to release the enzyme is critical for activity. The enzyme is inactivated by heat or alcohol and a cold water extraction of the finely ground fruits was found to provide the greatest...
molluscicidal activity. The saponin–containing extract of *P. dodecandra* is commonly referred to as ‘Endod’.\(^{(19)}\) Fruits of *P. americana* also possess molluscicidal properties.\(^{(G44)}\)

Abortifacient activity in mice has been exhibited by a related species *P. acinosa* Roxb. with activity strongest in the seed and weakest in the leaf. Activity in the various extracts was destroyed by heat and pepsin suggesting a protein to be the active principle.\(^{(20)}\)
Side–effects, Toxicity

Haematological aberrations have been observed in human peripheral blood following oral ingestion of the berries or exposure of broken skin/conjunctival membrane to the berry juice.\(^{21–23}\) Analysis of peripheral blood revealed plasmacytoid cells, dividing cells and mature plasmacytes. Eosinophilia was also noted. The mitogenic principles in pokeroo, lectins, are reported to be a mixture of agglutinating and non–agglutinating glycoproteins affecting both T cell and B cell lymphocytes.\(^{24}\)

Pokeroo leaf extracts have been reported to be agglutinating, but lacking in mitogenic activity.\(^{25}\)

A 43–year–old woman suffered the following symptoms 30 minutes after drinking a cup of herbal tea prepared from half a teaspoon of powdered pokeroo: nausea, vomiting, cramping, generalised abdominal pain followed by profound watery diarrhoea, weakness, haematemesis and bloody diarrhoea, hypotension and tachycardia.\(^{26}\) Chewing the root for the relief of a sore throat and cough has resulted in severe abdominal cramps, protracted vomiting and profuse watery diarrhoea.\(^{27}\) Additional symptoms of poisoning that have been documented for pokeroo include difficulty with breathing, spasms, severe convulsions and death.\(^{28}\)

The clinical symptoms of pokeroo poisoning have been reviewed.\(^{27}\)

All parts of the pokeroo plant are considered as potentially toxic, with the root generally recognised as the most toxic part.\(^{27}\) Toxicity is reported to increase with plant maturity although the young green berries are more toxic compared to the more mature red fruits.\(^{27}\)

High doses of saponin extracts have produced thymolytic effects in rats.\(^{12}\)

LD\(_{50}\) values for the saponin fraction (intraperitoneal injection) have been determined as 181 mg/kg in mice and 208 mg/kg in rats.\(^{12}\) In contrast, no deaths were observed in rats administered phytolaccagenin intraperitoneal injection up to a dose of 2 g/kg.\(^{12}\) Oral doses of saponin up to 1.5 g/kg did not produce any mortalities in treated rats.\(^{12}\)

The mutagenic potential of \(P.\) americana and \(P.\) dodecandra fruit extracts has been tested using \(Salmonella\ typhimurium\) strain TM677.\(^{19}\) No activity was found for any of the extracts tested.
Contra-indications, Warnings

Fresh pokeroott is poisonous and the dried root emetic and cathartic.\textsuperscript{(G42)} The toxic effects documented following the ingestion of pokeroott make it unsuitable for internal ingestion. In addition, external contact with the berry juice should be avoided: systemic symptoms of toxicity have occurred following exposure of broken skin and conjunctival membranes to the juice.

In 1979, the American Herb Trade Association declared that pokeroott should no longer be sold as a herbal beverage or food.\textsuperscript{(11)} It further recommended that all packages containing pokeroott carry an appropriate warning regarding the potential toxicity of pokeroott when taken internally. In the UK, manufacturers of licensed medicinal products are permitted to include pokeroott provided that the dose is restricted and that suitable evidence is given to demonstrate the absence of the toxic protein constituents.

\textit{Pregnancy and lactation}

Pokeroott is reputed to affect the menstrual cycle and is documented to exhibit uterine stimulant activity in animals.\textsuperscript{(G30)}
Pharmaceutical Comment

Apart from its traditional use as a herbal remedy, pokerooot is also known to possess molluscicidal properties. Anti-inflammatory activity documented in animal studies support the traditional use of pokerooot in rheumatism. However, pokerooot is also recognised as a toxic plant. The effects of pokerooot intoxication arise from the ingestion of any or all plant parts, liquid preparations of plant extracts such as herbal teas, or through skin contact with the plant. The main toxic agents are the pokeweed mitogen (lectins) and the glycoside saponins. The toxic properties of these two classes of compounds, mitogenic and irritant respectively, are well recognised. Excessive use of pokerooot cannot be supported in the light of these known toxicities.
References

See also General References G9 G10 G18 G20 G22 G30 G32 G36 G37 G42 G43 G48 G50 G64.


Poplar
Species (Family)

*Populus tremuloides* Michx. (Salicaceae)
Synonym(s)
Populus Alba, Quaking Aspen, White Poplar
Part(s) Used

Bark
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)

Complete German Commission E\(^{(G3)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL$^{(G37)}$
Constituents

See General References G6 G39 G40 G64.

**Glycosides**
Salicin (about 2.4%), salicortin, sali reposide and various benzoate derivatives including populin (salicin–6–benzoate), tremuloidin (salicin–2–benzoate) and tremulacin (salicortin–2–benzoate).

**Other constituents**
Tannins (unspecified), triterpenes including α-amyrin and β-amyrin, carbohydrates including glucose, fructose and various trisaccharides, fats, waxes.

**Food Use**
Poplar is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that poplar can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\(^{(G16)}\) In the USA, poplar is permitted for use in foods.\(^{(G65)}\)
Herbal Use

Poplar is stated to possess antirheumatic, anti-inflammatory, antiseptic, astringent, anodyne and chola gogue properties. Traditionally, it has been used for muscular and arthrodial rheumatism, cystitis, diarrhoea, anorexia with stomach or liver disorders, common cold, and specifically for rheumatoid arthritis.\(^{(G6\ G7\ G64)}\)

The buds of *Populus tremula* (European white poplar, aspen) and *Populus nigra* (black poplar) are used, reputedly as expectorant and circulatory stimulant remedies, for upper respiratory tract infections and rheumatic conditions.\(^{(G49)}\)
**Dosage**

*Dried bark*
1–4 g or by decoction three times daily.\(^{[G6 G7]}\)

*Liquid extract*
1–4 mL (1 : 1 in 25% alcohol) three times daily.\(^{[G6 G7]}\)
Pharmacological Actions

*In vitro and animal studies*

None documented for poplar. See Willow for the pharmacological actions associated with salicylates.

*Clinical studies*

None documented for poplar. The pharmacological actions of salicylates in humans are well documented and are applicable to poplar. Salicin is a prodrug that is metabolised to saligenin in the gastrointestinal tract and to salicylic acid following absorption.
Side-effects, Toxicity

None documented. See Willow for side-effects and toxicity associated with salicylates.
Contra-indications, Warnings

See Willow for contra-indications and warnings associated with salicylates.

Pregnancy and lactation
The safety of poplar taken during pregnancy has not been established. See Willow for contra-indications and warnings regarding the use of salicylates during pregnancy and lactation.
Pharmaceutical Comment

The chemistry of poplar is characterised by the phenolic glycoside components, which support some of the reputed herbal uses. The usual precautions associated with other salicylate-containing drugs are applicable to poplar.
References

Prickly Ash, Northern
Species (Family)

*Zanthoxylum americanum* Miller (Rutaceae)
Synonym(s)

Toothache Bark, Xanthoxylum, Zanthoxylum
Part(s) Used

Bark, berry
Pharmacopoeial and Other Monographs

BHP 1983\(^{(G7)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

Northern prickly ash is not included in the GSL.\textsuperscript{[G37]}
Constituents
See General References G6 G41 G64.

**Alkaloids**
Isoquinoline–type. Lauriflorine and nitidine (major constituents), candinicine, chelerythrine, magnoflorine and tembetarine.

**Coumarins**
Xanthyletin, xanthoxyletin, alloxanth oxyletin and 8-(3,3–dimethylallyl)alloxanthoxyletin.

**Other constituents**
Resins, tannins and acrid volatile oil.

**Other plant parts**
Two furoquinoline alkaloids (γ-fagarine and skimmianine) have been isolated from the leaves.
Food Use

Prickly ash is listed by the Council of Europe as a natural source of food flavouring (category N3). This category indicates that prickly ash can be added to foodstuffs in the traditionally accepted manner, but that there is insufficient information available for an adequate assessment of potential toxicity. In the USA, prickly ash is listed as GRAS (Generally Recognised As Safe).
Herbal Use

Prickly ash is stated to possess circulatory stimulant, diaphoretic, antirheumatic, carminative and sialagogue properties. Traditionally, it has been used for cramps, intermittent claudication, Raynaud’s syndrome, chronic rheumatic conditions, and specifically for peripheral circulatory insufficiency associated with rheumatic symptoms. The berries are stated to be therapeutically more active in circulatory disorders. (G6 G7 G64)
Dosage

**Dried bark**
1–3 g or by decoction three times daily. (G6 G7)

**Bark, liquid extract**
1–3 mL (1 : 1 in 45% alcohol) three times daily. (G6 G7)

**Bark, tincture**
2–5 mL (1 : 5 in 45% alcohol) three times daily. (G6 G7)

**Dried berry**
0.5–1.5 g. (G6 G7)

**Berry, liquid extract**
0.5–1.5 mL (1 : 1 in 45% alcohol). (G6 G7)
Pharmacological Actions

*In vitro* and animal studies

None documented for northern prickly ash. See Prickly Ash, Southern for activities of alkaloid constituents (e.g. chelerythrine and nitidine).
Side-effects, Toxicity

The alkaloid constituents are potentially toxic (see Prickly Ash, Southern ).
Contra-indications, Warnings

Excessive ingestion may interfere with anticoagulant therapy in view of the coumarin constituents (see Prickly Ash, Southern).

Pregnancy and lactation

The safety of northern prickly ash has not been established. In view of the pharmacologically active constituents the use of northern prickly ash during pregnancy and lactation should be avoided.
Pharmaceutical Comment

Northern prickly ash contains similar alkaloid constituents to the southern species but varies with respect to other documented components. No pharmacological studies documented specifically for northern prickly ash were located. However, activities have been reported for individual alkaloid constituents and the monograph for southern prickly ash should be consulted. There is limited scientific evidence to support the traditional herbal uses. In view of the pharmacologically active constituents and potential toxicity associated with the alkaloids, excessive use of northern prickly ash should be avoided.
See General References G6 G7 G10 G16 G31 G36 G37 G41 G43 G64.
Prickly Ash, Southern
Species (Family)

Zanthoxylum clava-herculis L. (Rutaceae)
Synonym(s)
Toothache Bark, Xanthoxylum, Zanthoxylum
Part(s) Used

Bark, berry
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

Martindale 33rd edition\(^{(G67)}\)
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents

See General References G6 G41 G45 G64.

**Alkaloids**
Isoquinoline–type. Chelerythrine and magno florine (major constituents), candicine, lauriflorine, nitidine, \(N\)-acetylanonaine\(^{(1)}\) and tembetarine.

**Amides**
Cinnamamide, herculin and neoherculin.

**Lignans**
\((-\)-Asarinin, \((-\)-sesamin, \(\gamma,\gamma\)-dimethylallyl ether of \((-\)-pluviatilol\(^{(1)}\)

**Other constituents**
Resins, tannins and an acrid volatile oil (about 3.3%).
Food Use

Southern prickly ash is listed by the Council of Europe as a natural source of food flavouring (category N3). This category indicates that prickly ash can be added to foodstuffs in the traditionally accepted manner, but that there is insufficient information available for an adequate assessment of potential toxicity.\(^{(G16)}\)
Herbal Use

Southern prickly ash is stated to possess circulatory stimulant, diaphoretic, antirheumatic, carminative and sialogogue properties. Traditionally, it has been used for cramps, intermittent claudication, Raynaud’s syndrome, chronic rheumatic conditions, and specifically for peripheral circulatory insufficiency associated with rheumatic symptoms. The berries are stated to be therapeutically more active in circulatory disorders. (G6 G7 G8 G64)
Dosage

*Dried bark*
1–3 g or by decoction three times daily.\(^{(G6 \ G7)}\)

*Bark, liquid extract*
1–3 mL (1 : 1 in 45% alcohol) three times daily.\(^{(G6 \ G7)}\)

*Bark, tincture*
2–5 mL (1 : 5 in 45% alcohol) three times daily.\(^{(G6 \ G7)}\)

*Dried berry*
0.5–1.5 g.\(^{(G6 \ G7)}\)

*Berry, liquid extract*
0.5–1.5 mL (1 : 1 in 45% alcohol).\(^{(G6 \ G7)}\)
Pharmacological Actions

In vitro and animal studies

Southern prickly ash has been reported to act as a reversible neuromuscular blocking agent. Activity was associated with a neutral fraction of the bark that was thought to act primarily by blockade of endplate receptors.\(^{(2)}\)

Various activities have been documented for the benzophenanthridine alkaloids (e.g. chelerythrine, nitidine) present in southern prickly ash. Hypotensive properties in mice have been documented for nitidine chloride, a single dose of 2 mg/kg body weight lowered the blood pressure by 20% within 90 minutes and persisted for 6 hours.\(^{(3)}\) Nitidine was also found to antagonise the effects of angiotensin–induced hypertension.\(^{(3)}\) Antileukaemic activity has been documented for nitidine, although preclinical toxicity prevented further investigations.\(^{(4,5)}\)

Anti–inflammatory activity in rats has been documented for chelerythrine (10 mg/kg by mouth) comparable to that achieved with indomethacin (5 mg/kg by mouth).\(^{(6)}\) Chelerythrine has also been reported to potentiate the analgesic effect of morphine, prolong barbiturate–induced sleep, and cause temporary hypertension followed by hypotension in cats, mice and rabbits.\(^{(7)}\)

Significant antimicrobial activity towards Gram–positive bacteria and *Candida albicans* has been documented for chelerythrine, although conflicting activities have been reported regarding Gram–negative bacteria.\(^{(6)}\) Chelerythrine has been shown to interact with Na\(^+\)K\(^+\) ATPase and to inhibit hepatic L-alanine and L-aspartate aminotransferases in the rat, while nitidine has been reported to inhibit tRNA methyltransferase and catechol-O-methyltransferase.\(^{(5)}\)

The lignan component, asarinin, has been reported to possess antitubercular activity.\(^{(G41)}\) Neoherculin is reported to possess insecticidal and sialogogic properties.\(^{(1)}\)

Pharmacological activities, including anti–inflammatory, cardiovascular and antibacterial properties have been documented for various other *Zanthoxylum* species (or *Fagara/Xanthoxylum* species).\(^{(5)}\) For example, the root of *Zanthoxylum zanthoxyloides*, a Nigerian species, is commonly used as a chewing stick. These sticks are believed to possess antimicrobial properties and extracts were found to exhibit anti microbial activity towards more than 20 organisms, including Gram–positive and Gram–negative bacteria, and
*Candida* species.\(^{(5)}\) Anti-inflammatory activity (carrageenan rat paw oedema test) has been described for fagaramide (piperonyl-4-acrylic isobutylamide), isolated from *Z. zanthoxyloides*.\(^{(8)}\) The activity, approximately 20 times less potent than indometacin, was thought to be partially mediated by inhibition of prostaglandin synthesis.\(^{(8)}\)

The essential oil obtained from the Indian species *Zanthoxylum limonella* has been reported to exhibit *in vitro* anthelmintic activity against earthworms, tapeworms and hookworms that was stated to be superior to that of piperazine phosphate.\(^{(9)}\)
Side-effects, Toxicity

None documented in humans. Ingestion of southern prickly ash by cattle, chicken and fish has proved lethal. This was attributed to the neuromuscular blocking properties of the bark.\(^{(2)}\) Neoherculin is reported to be the major ichthyotoxic principle in an extract of southern prickly ash bark.

The acute and chronic toxicity of chelerythrine in mice is reported to be low.\(^{(4)}\) LD\(_{50}\) values were stated as 18.5 mg/kg body weight (intravenous injection) and 95 mg/kg (subcutaneous injection). Oral administration of 10 mg/kg for three days followed by 5 mg/kg for seven days produced no adverse effects.
Contra–indications, Warnings

None documented for southern prickly ash. Chelerythrine has been reported to interact with Na\(^+\)K\(^+\) ATPase which may interfere with cardiac glycoside therapy. However the clinical relevance of this with respect to prickly ash is unknown. Hypotensive and sedative activities have been documented in animals. Both chelerythrine and nittidine have been reported to inhibit various hepatic enzymes (see In vitro and animal studies). The alkaloid constituents in southern prickly ash are potentially toxic.

Pregnancy and lactation
The safety of southern prickly ash has not been established. In view of this and the pharmacologically active compounds, the use of southern prickly ash during pregnancy and lactation is best avoided.
Pharmaceutical Comment

The chemistry of southern prickly ash is well documented and particularly characterised by the alkaloid constituents. Limited pharmacological information has been documented for southern prickly ash, although several properties have been described for individual constituents. With the exception of anti-inflammatory and analgesic properties few data have been documented that support the herbal uses. Limited toxicity data are available and some benzophenanthridine alkaloids are associated with cytotoxicity. In view of this, excessive use of prickly ash should be avoided. Northern prickly ash has been used for similar herbal uses but has a different chemical composition compared to the southern species (see Prickly Ash, Northern).
References

See also General References G6 G9 G10 G16 G31 G37 G41 G43 G64.


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Species (Family)

i. *Anemone pulsatilla* L. (Ranunculaceae)

ii. *Anemone pratensis* L.

iii. *Anemone patens* L.
Synonym(s)

Pasque Flower, *Pulsatilla nigrans*
Part(s) Used

Herb
Pharmacopoeial and Other Monographs

BHC 1992\textsuperscript{(G6)}

BHP 1996\textsuperscript{(G9)}

Martindale 33rd edition\textsuperscript{(G67)}

PDR for Herbal Medicines 2nd edition\textsuperscript{(G36)}
Legal Category (Licensed Products)

GSL^{G37}
Constituents

See General References G6 G22 G48 G64.

**Flavonoids**
Delphinidin and pelargonidin glycosides.

**Saponins**
Hederagenin (as the aglycone).

**Volatile oils**
Ranunculin (a glycoside); enzymatic hydrolysis yields the unstable lactone protoanemonin which readily dimerises to anemonin.

**Other constituents**
Carbohydrates (e.g. arabinose, fructose, galactose, glucose, rhamnose), triterpenes (e.g. β-amyrin) and β-sitosterol.
Food Use

Pulsatilla is not used in foods.
Herbal Use

Pulsatilla is stated to possess sedative, analgesic, antispasmodic and bactericidal properties. Traditionally, it has been used for dysmenorrhea, orchitis, ovaralgia, epididymitis, tension headache, hyperactive states, insomnia, boils, skin eruptions associated with bacterial infection, asthma and pulmonary disease, earache, and specifically for painful conditions of the male or female reproductive system. Pulsatilla is widely used in homeopathic preparations as well as in herbal medicine.
Dosage

**Dried herb**
0.12–0.3 g by infusion or decoction three times daily. (G6 G7)

**Liquid extract**
0.12–0.3 mL (1 : 1 in 25% alcohol) three times daily. (G6 G7)

**Tincture**
0.3–1.0 mL (1 : 10 in 40% alcohol) three times daily. (G6 G7)
Pharmacological Actions

In vitro and animal studies

Utero–activity (stimulant and depressant) has been documented for pulsatilla.\(^{(1,2,G^{30})}\) In vivo sedative and antipyretic properties in rodents have been documented for anemonin and protoanemonin.\(^{(3)}\)

Cytotoxicity (KB tumour system) has been reported for anemonin.\(^{(G^{22})}\)
Side-effects, Toxicity

Fresh pulsatilla is poisonous because of the toxic volatile oil component, protoanemonin. Protoanemonin rapidly degrades to the non-toxic anemonin. Inhalation of vapour from the volatile oil may cause irritation of the nasal mucosa and conjunctiva.\(^{(G51)}\) Allergic reactions to pulsatilla have been documented and patch tests have produced vesicular reactions with hyperpigmentation.\(^{(G51)}\) Cytotoxicity has been documented for anemonin (see *In vitro* and animal studies).
Contra–indications, Warnings

Fresh pulsatilla is poisonous and should not be ingested. External contact with the fresh plant should be avoided. The toxic principle, protoanemonin, rapidly degrades to the non–toxic anemonin during drying of the plant material. Individuals may experience an allergic reaction to pulsatilla, especially those with an existing hypersensitivity.

**Pregnancy and lactation**
Pulsatilla is reputed to affect the menstrual cycle.\(^{(G22)}\) Utero–activity has been documented for pulsatilla (*see In vitro* and animal studies). In view of this, the use of pulsatilla during pregnancy should be avoided. Excessive ingestion is best avoided during lactation.
Pharmaceutical Comment

Pulsatilla is widely used in both herbal and homeopathic preparations, although little documented chemical and pharmacological information is available to assess its true benefit. The fresh plant is known to be irritant; it contains a toxic principle (protoanemonin) and should not be ingested. The dried plant material is not considered to be toxic.
References

See also General References G6 G9 G10 G22 G30 G31 G36 G37 G43 G48 G51 G64.

1. Pilcher JM et al. The action of the so-called female remedies on the excised uterus of the guinea-pig. *Arch Intern Med* 1916; **18**: 557–583.


Species (Family)

i. *Picrasma excelsa* (Sw.) Planch. (Simaroubaceae)

ii. *Quassia amara* L.
Synonym(s)

Bitterwood, Picrasma


ii. Surinam Quassia
Part(s) Used

Stem wood
Pharmacopoeial and Other Monographs

BHC 1992\(^{G6}\)

BHP 1996\(^{G9}\)

Martindale 33rd edition\(^{G67}\)

PDR for Herbal Medicines 2nd edition\(^{G36}\)
Legal Category (Licensed Products)

GSL\textsuperscript{(G37)}
Constituents

See General References G2 G6 G22 G41 G64.

Alkaloids
Indole-type. Canthin-6-one, 5-methoxycanthin-6-one, 4-methoxy-5-hydroxycanthin-6-one, \( N \)-methoxy-1-vinyl-\( \beta \)-carboline.\(^{(1,2)}\)

Terpenoids
Isoquassin (picrasmin) in \( P. \ excelsa \), quassin 0.2%, quassinol, quassimarin,\(^{(3)}\) 18-hydroxy quassin, neoquassin, a dihydronorneoquassin\(^{(4)}\) and simalikalactone D in \( Q. \ amara \).

Coumarins
Scopoletin.\(^{(1)}\)

Other constituents
\( \beta \)-Sitosterol, \( \beta \)-sitostenone; thiamine 1.8% (in \( P. \ excelsa \)).
Food Use

Quassia is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that quassia can be added to foodstuffs in small quantities, although the concentration of quassin must not exceed 5 mg/kg; a concentration of 50 mg/kg is permitted in alcoholic beverages and 10 mg/kg in pastilles and lozenges. In the USA, quassia is regarded by the Food and Drugs Administration (FDA) as GRAS (Generally Regarded As Safe).
Herbal Use

Quassia is stated to possess bitter, orexigenic, sialogogue, gastric stimulant and anthelmintic properties. Traditionally, it has been used for anorexia, dyspepsia, nematode infestation (by oral or rectal administration), pediculosis (by topical application), and specifically for atonic dyspepsia with loss of appetite. (G2 G6 G7 G8 G64)
Dosage

*Dried wood*
0.3–0.6 g or by cold infusion three times daily.\(^{(G6 \ G7)}\)

*Concentrated Quassia Infusion*
(BPC 1959) 2–4 mL. Quassia Infusion is prepared by diluting one volume of Concentrated Quassia Infusion to eight volumes with water.

*Tincture of Quassia*
(BP 1948) 2–4 mL.

*Enema*
150 mL per rectum (infusion with cold water, 1 in 20) on three successive mornings together with 16 g magnesium sulfate by mouth.
Pharmacological Actions

The quassinoids are reported to possess bitter properties 50 times greater than quinine.\textsuperscript{(G22)}

\textbf{In vitro and animal studies}

The β-carboline alkaloids have exhibited positive inotropic activity \textit{in vitro}.\textsuperscript{(1)} Canthin-6-one is reported to possess antibacterial and antifungal activity. Cytotoxic and amoebicidal activities (assessed against guinea–pig keratinocyte and \textit{Entamoeba histolytica} test systems, respectively) have been documented for canthin-6-one and quassin (\textit{P. excelsa}).\textsuperscript{(5)} However, later studies have disputed any amoebicidal action. Quassin is reported to be inactive against P388 leukaemia and 9KB test systems. Significant antitumour activity in mice against the P388 lymphatic leukaemia and \textit{in vitro} against human carcinoma of the nasopharynx (KB) has been documented.\textsuperscript{(3)} Quassimarin and simalikalactone were both isolated from the active extract.

\textbf{Clinical studies}

The successful treatment of 454 patients with headlice has been documented for quassia tincture.\textsuperscript{(6)} Quassia has been used as an enema to expel threadworms.\textsuperscript{(G44)}
Side–effects, Toxicity

No side–effects have been reported in 454 patients who used quassia tincture as a scalp lotion to treat headlice.\(^6\) Large doses of quassia may irritate the stomach and cause vomiting.\(^{G6}\)
Contra–indications, Warnings

Excessive doses may interfere with existing cardiac and anticoagulant therapies. However, the coumarin concentrations in quassia are not thought to pose a hazard. In addition, large doses of quassia are emetic and therefore excessive consumption is self–limiting.

Pregnancy and lactation
In view of the reported cytotoxic and emetic activities, the use of quassia during pregnancy and lactation is best avoided.
Pharmaceutical Comment

The chemistry of quassia is well studied and is characterised by bitter terpenoids (quassinoids) and β-carboline indole alkaloids. Limited data have been documented to justify the traditional herbal uses although the bitter principles support the use of quassia as an appetite stimulant in anorexia. However, in view of the documented cytotoxic activities and limited toxicological data, quassia in herbal remedies should not be taken in amounts greatly exceeding those used in foods.
References

See also General References G2 G6 G9 G12 G16 G22 G29 G36 G37 G41 G43 G56 G64.


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Queen’s Delight
Species (Family)

*Stillingia sylvatica* L. (Euphorbiaceae)
Synonym(s)
Queen’s Root, Stillingia, *Stillingia treculeana* (Muell. Arg.) Johnst., Yaw Root
Part(s) Used

Root
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL^{G37}
**Constituents**

See General References G41 G48 G64.

**Terpenoids**

Eight compounds, termed stillingia factors $S_1$–$S_8$, have been isolated and identified as daphnane-type and tigliane-type esters carrying saturated, polyunsaturated or hydroxylated fatty acids.\(^{(1)}\)

**Other constituents**

Volatile oil 3–4%, fixed oil, acrid resin (sylvacrol), resinic acid, stillingine (a glycoside) and tannin.

**Other plant parts**

Hydrocyanic acid (leaf and stem).\(^{(1)}\)
Food Use
Queen’s delight is not used in foods.
Herbal Use

Queen’s delight is stated to possess sialogogue, expectorant, diaphoretic, dermatological, astringent, antispasmodic and, in large doses, cathartic properties. Traditionally, it has been used for bronchitis, laryngitis, laryngismus stridulus, cutaneous eruptions, haemorrhoids, constipation and specifically for exudative skin eruption with irritation and lymphatic involvement, and laryngismus stridulus.\textsuperscript{G7 G64}
Dosage

**Dried root**
1–2 g or by decoction three times daily.\(^{(G7)}\)

**Liquid extract**
0.5–2.0 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)

**Tincture**
1–4 mL (1 : 5 in 45% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions
None documented.
Side-effects, Toxicity

Overdose of queen’s delight is reported to cause vertigo, burning sensation of the mouth, throat and gastrointestinal tract, diarrhoea, nausea and vomiting, dysuria, aches and pains, pruritus and skin eruptions, cough, depression, fatigue and perspiration.\(^{(G22)}\) The diterpene esters are toxic irritant principles known to cause swelling and inflammation of the skin and mucous membranes.\(^{(1,G33)}\)

The leaves and stem are documented to be toxic to sheep because of the hydrocyanic acid content.\(^{(2)}\)
Contra-indications, Warnings

In view of the irritant nature of the diterpene esters, queen’s delight may cause irritation to the mucous membranes. It is stated that queen’s delight should be used with care, and never taken in large doses.\(^\text{G49}\) It is recommended that the root should not be used after two years of storage.\(^\text{G49}\)

Pregnancy and lactation

In view of the irritant and potentially toxic constituents, the use of queen’s delight during pregnancy and lactation should be avoided.
The Euphorbiaceae plant family is characterised by the diterpene esters. These compounds, known as phorbol, ingenane or daphnane esters depending on their skeleton type, have been investigated as constituents of genera such as *Euphorbia* and *Croton*, and some of them have been found to be co-carcinogenic and highly irritant to mucous membranes.\(^{G33}\) No scientific evidence was found to justify the reputed herbal uses. In view of this and the potential toxicity of queen’s delight excessive use is not recommended.
References

See also General References G9 G10 G22 G33 G36 G37 G41 G48 G49 G64.

Raspberry
Species (Family)

Rubus idaeus L. (Rosaceae)
Synonym(s)

Rubus
Part(s) Used

Leaf
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents

See General References G2 G59 G62 G64.

Limited phytochemical information is available for raspberry. Documented constituents include acids, polypeptides, tannins and flavonoids (e.g. rutin). (1)
Food Use

Both the leaf and fruit are listed by the Council of Europe as natural sources of food flavouring (categories N2 and N1, respectively). Category N2 allows the addition of the leaf to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product. Category N1 indicates that no restrictions apply to the fruit. (G16) Raspberry fruit is commonly used in foods.
Herbal Use

Raspberry is stated to possess astringent and *partus praeparator* properties. Traditionally, it has been used for diarrhoea, pregnancy, stomatitis, tonsillitis (as a mouthwash), conjunctivitis (as an eye lotion), and specifically to facilitate parturition. (G2 G7 G64)
Dosage

*Dried leaf*
4–8 g or by infusion three times daily.\(^{(G7)}\)

*Liquid extract*
4–8 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

**In vitro and animal studies**

Utero-activity has been documented for a leaf infusion in both pregnant and non-pregnant rat and human uteri.\(^{(2)}\) The extract was reported to have little or no effect on the uterine strips from non-pregnant rats, but inhibited contractions of those from pregnant rats. Similarly, the extract had no effect on strips from non-pregnant human uteri, but initiated contractions in strips from human uteri at 10–16 weeks of pregnancy. The intrinsic rhythm of the uteri in which a pharmacological effect was observed (pregnant rat and human uteri) was reported to become more regular, with contractions, in most cases, less frequent.\(^{(2)}\) Aqueous extracts of raspberry leaves have been reported to contain a number of active constituents, including a smooth muscle stimulant, an anticholinesterase, and an antispasmodic that antagonised the stimulant actions of the two previous fractions. The smooth muscle stimulant fraction was more potent towards uterine muscle.\(^{(3)}\)

Hypoglycaemic activity has been documented for a related species, *Rubus fructicosus* L., in both non-diabetic and diabetic (glucose-induced and alloxan-induced) rabbits.\(^{(4)}\) Greatest activity was observed in the glucose-induced diabetic rabbits. The authors concluded that *R. fructicosus* possesses slight hypoglycaemic activity, which, in part, results from an increase in the liberation of insulin. Tannins are known to possess astringent properties.

*In vitro* antiviral activity documented for raspberry fruit extract has been attributed to the phenolic constituents, in particular to tannic acid.\(^{(5)}\)
Side-effects, Toxicity

None documented.
Contra–indications, Warnings

The excessive ingestion of tannins is not recommended. Hypoglycaemic activity *in vivo* has been documented for a related species.

**Pregnancy and lactation**
Raspberry is traditionally recommended for use during labour to help ease parturition. Animal studies (*in vitro*) have reported that raspberry can reduce and initiate uterine contractions. In view of this, raspberry should not be used during pregnancy and, if taken during labour, should only be done so under medical supervision.
Pharmaceutical Comment

Limited phytochemical information is available for raspberry leaf. However, the documented presence of tannin constituents supports some of the reputed herbal uses, although it is unsuitable to use as a herbal remedy to treat eye infections such as conjunctivitis. Raspberry leaf is widely recommended to be taken during pregnancy to help facilitate easier parturition. Utero-activity has been documented for raspberry leaf and in view of this it should not be taken during pregnancy, unless under medical supervision.
References

See also General References G2 G9 G11 G16 G31 G32 G36 G37 G43 G59 G62 G64.

Red Clover
Species (Family)

*Trifolium pratense* L. (Leguminosae)
Synonym(s)
Cow Clover, Meadow Clover, Purple Clover, Trefoil
Part(s) Used

Flowerhead
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents

See General References G6 G22 G41 G64.

**Carbohydrates**
Arabinose, glucose, glucuronic acid, rhamnose, xylose (following hydrolysis of saponin glycosides); polysaccharide (a galactoglucomannan).

**Coumarins**
Coumarin, medicagol.

**Isoflavonoids**
Biochanin A, daidzein, formononetin, genistein, pratensin, trifoside, calycosine galactoside\(^1\) and pectolinarin.

**Flavonoids**
Isorhamnetin, kaempferol, quercetin, and their glycosides.\(^2\)

**Saponins**
Soyasapogenols B–F (C–F artefacts) and carbohydrates (see above) yielded by acid hydrolysis.\(^3\)

**Other constituents**
Coumaric acid, phaseolic acid, salicylic acid, *trans*- and *cis*-clovamide (L-dopa conjugated with *trans*- and *cis*-caffeic acids), resin, volatile oil (containing furfural),\(^4\) fats, vitamins and minerals. Cyanogenic glycosides have been documented for a related species, *Trifolium repens*.\(^{G33}\)
Food Use

Red clover is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that it can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\textsuperscript{(G16)} In the USA, red clover is listed as GRAS (Generally Recognised As Safe).\textsuperscript{(G65)}
Red clover is stated to act as a dermatological agent, and to possess mildly antispasmodic and expectorant properties. Tannins are known to possess astringent properties. Traditionally red clover has been used for chronic skin disease, whooping cough, and specifically for eczema and psoriasis. (G6 G7 G8 G64)
Dosage

**Dried flowerhead**
4 g or by infusion three times daily.\(^{(G6 G7)}\)

**Liquid extract**
1.5–3.0 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G6 G7)}\)

**Tincture**
1–2 mL (1 : 10 in 45% alcohol) three times daily.\(^{(G6 G7)}\)
Pharmacological Actions

*In vitro* and animal studies

Biochanin A, formononetin, and genistein (isoflavones) are known to possess oestrogenic properties.\(^5\) The saponin constituents are reported to lack any haemolytic or fungistatic activity.\(^3\) A possible chemoprotective effect has been documented for biochanin A, which has been reported to inhibit carcinogenic activity in cell culture.\(^6\)
Side–effects, Toxicity

Urticarial reactions have been documented.\textsuperscript{(G51)} Infertility and growth disorders have been reported in grazing animals.\textsuperscript{(G33)} These effects have been attributed to the oestrogenic isoflavone constituents, in particular to formononetin.\textsuperscript{(5)}
Contra–indications, Warnings

In view of the oestrogenic constituents, excessive ingestion should be avoided. Large doses may interfere with anticoagulant and hormonal therapies (coumarin and isoflavonoid constituents).

**Pregnancy and lactation**

In view of the oestrogenic components the use of red clover during pregnancy and lactation should be avoided.
The chemistry of red clover is well documented. Limited information is available on the pharmacological properties and no documented scientific evidence was found to justify the herbal uses. Reported oestrogenic side-effects in grazing animals have been attributed to the isoflavone constituents. Little toxicity data are available for red clover. In view of this and the isoflavone and coumarin components, excessive ingestion should be avoided.
References

See also General References G6 G9 G16 G22 G31 G32 G33 G36 G41 G43 G51 G64.


Rhodiola
Species (Family)

*Rhodiola rosea* L. (Crassulaceae)
**Synonym(s)**


*Rhodiola rosea* and several other *Rhodiola* species, such as *Rhodiola quadrifida* (Pall.) Fisch. et Mey., are used in Traditional Chinese Medicine and are given the generic name Rhodiolae Radix (meaning rhodiola root).\(^{(1)}\)
Part(s) Used

Rhizome and root
Pharmacopoeial and Other Monographs

None
Legal category (Licensed Products)

Rhodiola is not on the GSL. There are no licensed products containing rhodiola available in the UK.
Constituents

Flavonoids
Herbacetin, gossypetin, kaempferol and their glycosides rhodionin, rhodionidin, rhodiolgin, rhodolgadin, rhodalin, rhodaldin, rhodiosin and kaempferol-7-O-α-L-rhamnopyranoside.\(^{(2)}\)

Phenylethanoids
Hydroxyphenylethyl tyrosol (p-tyrosol) and its glycoside salidroside (rhodioloside; p-hydroxyphenylethyl-O-β-d-glucopyranoside).\(^{(3)}\)

Phenylpropanoids
Rosin (cinnamyl-O-β-d-glucopyranoside), rosarin (cinnamyl-(6′-O-α-L-arabinofuranosyl)-O-β-d-glucopyranoside), rosavin (cinnamyl-(6′-O-α-L-arabinopyranosyl)-O-β-d-glucopyranoside),\(^{(4)}\) sachaliside 1 (4-hydroxy-cinnamyl-O-β-d-glucopyranoside), vimalin (4-methoxy-cinnamyl-O-β-d-glucopyranoside), cinnamyl-(6′-O-β-xylopyranosyl)-O-β-glucopyranoside, 4-methoxy-cinnamyl-(6′-O-α-arabinopyranosyl)-O-β-glucopyranoside\(^{(4)}\) and cinnamyl alcohol.\(^{(2)}\)

Volatile oils
0.05% (1% has been documented for Russian material) of dry weight of crude rhizome, containing monoterpene hydrocarbons, monoterpane alcohols and straight-chain aliphatic alcohols as major components; major compounds (>3%) include geraniol, 1,4-p-menthadien-7-ol, limonene, α-pinene (monoterpenes), and decanol and dodecanol (aliphatic alcohols).\(^{(5)}\) The monoterpenes rosiridol\(^{(2)}\) and rosiridin\(^{(6)}\) are also present.

Other constituents
Picein ((4-O-β-d-glucopyranosyl)-acetophenone), benzyl-O-β-d-glucopyranoside,\(^{(4)}\) sterols (β-sitosterol, daucosterol),\(^{(2)}\) tannins,\(^{(7)}\) gallic acid and its esters.\(^{(8)}\)

Quality
As with other herbal medicines, there can be qualitative and quantitative variation in the profile of constituents present in crude rhodiola material and in marketed products. Content of salidroside and tyrosol in samples of *R. rosea* root obtained from different outlets in the Yanbian area of China has been reported to vary up to ten-fold (range: 1.3–11.1 and 0.3–2.2 mg/g for salidroside and tyrosol, respectively).\(^{(9)}\)

Standardisation of commercial extracts of *R. rosea* may be for content of
‘salidrosides’ and ‘rosavins’, although the precise composition of these groups may not always be stated.\(^6\) For example, one commercial product available in the UK is described as containing 3% ‘rosavins’, 1% ‘salidrosides’ and 24% polyphenols per 100 mg capsule of rhodiola. Several other rhodiola products are stated to contain 1% ‘salidrozid’ and 40% polyphenols. The quality of these products is not assured as they are not licensed products.

An extract of \(R.\ rosea\) (SHR-5) tested in recent clinical trials was standardised for salidroside content (170 mg\(^{10}\) or 185 mg\(^{11}\) extract provided approximately 4.5 mg salidroside).\(^{10,11}\)

Analysis of 19 samples of dried rhizome from 10 \(Rhodiola\) species other than \(R.\ rosea\), obtained in the east of Qinghai province in China, found that all species contained salidroside, although only five species had a content greater than 0.3\%.\(^{12}\) Five species also contained lotaustralain, a cyanoglucoside which is toxic to humans following oral administration.
Food Use

Rhodiola root and rhizome are not used in foods, although the aerial parts of *R. rosea* are used as a food ingredient.\(^{(10)}\)
Herbal Use

Rhodiola has a long history of use as a medicinal plant in several traditional systems. It is reported to have been used as a ‘brain tonic’, to treat headache and lung disorders,\(^{(10)}\) and to eliminate fatigue and improve work capacity.\(^{(13)}\) It is also stated to have stimulant properties, and to prevent stress.\(^{(10)}\)

Modern interest in rhodiola is focused on its adaptogenic properties. (Adaptogens increase an organism’s resistance to physical, chemical and biological stressors, and have a normalizing influence on bodily systems.)
Dosage

Authoritative guidance on dosages of *R. rosea* preparations for adults and children hitherto is lacking. Given the intended uses of *R. rosea*, it is not suitable for use in children.

In clinical trials involving young adults, dosages used have ranged from *R. rosea* root extract (SHR-5) 50 mg twice daily (salidroside content not stated)\(^{(14)}\) to 170 mg daily (equivalent to approximately 4.5 mg salidroside) for two weeks.\(^{(10)}\) In another study investigating the effects of *R. rosea* on fatigue and stress, participants took a single dose of two or three capsules (total daily doses of 370 mg and 555 mg *R. rosea* extract, respectively).\(^{(11)}\)

Other dosages have been suggested, although data supporting these regimens are not available. For example, for long-term use as an adaptogen (up to four months) daily doses of *R. rosea* extract standardised for 1% rosavin of 360–600 mg (equivalent to 3.6–6 mg rosavin daily), initiated several weeks before an anticipated period of increased stress and continued for its duration, have been stated.\(^{(15)}\) For use as an adaptogen in acute stressful situations, such as taking an examination, a dose three times that suggested for longer term administration, taken as a single dose (i.e. 1080–1800 mg *R. rosea* extract standardised for 1% rosavin, equivalent to 10.8–18 mg rosavin) has been advised.\(^{(15)}\)
Pharmacological Actions

A substantial body of research investigating the pharmacological and clinical properties of *R. rosea* has been undertaken, although much of this work has been published in the Russian scientific literature, making it difficult to access. The following represents a summary of some of this literature, together with that published in English. Where papers published in Russian or Ukrainian are cited, information has been taken from the English abstract and/or previous authoritative reviews only, thus the data and methods have not been scrutinised here. The information is included here to guide the interested reader to the original literature, and should not be taken as confirmation of the effects described.

Several activities, including various adaptogenic effects, as well as anti-arrhythmic, cardioprotective, antimutagenic and antitumour properties, have been described for *R. rosea* following preclinical and/or clinical studies. Several of these more specific activities are often considered as part of the overall profile of adaptogenic activity of *R. rosea*. The precise chemical constituents responsible for the documented pharmacological activities of *R. rosea* root and rhizome are not fully understood, although tyrosol and salidroside (rhodioloside) and the cinnamyl (phenylpropanoid) glycoside constituents (rosin, rosavin, rosarin) (see Constituents) are believed to be important for certain activities.\(^3,4,16\)

**In vitro and animal studies**

**Adaptogenic effects**

Several adaptogenic properties have been described for *R. rosea* extracts following preclinical studies.

The effects of *R. rosea* extract on erythropoiesis and granulocytopoiesis following paradoxical sleep deprivation (which leads to behavioural disorders and changes in haematopoiesis, for example, suppression of bone marrow erythropoiesis and activation of granulocytopoiesis) have been explored in experiments in mice.\(^17,18\) In one study, mice were given *R. rosea* extract (no further details given) 1 ml/kg once daily by gastric lavage for five days before sleep deprivation.\(^17\) Measurement of peripheral blood reticulocytes over days 1–7 following treatment indicated that *R. rosea*, compared with control, stimulated erythropoiesis during the early stage (days 1–3) but inhibited accumulation of erythroid cells in the bone marrow during days 4–7. In contrast, in a similar experiment using the same dosage regimen, administration of *R. rosea* before sleep deprivation did not modulate granulocytopoiesis as demonstrated by bone marrow content of neutrophilic
A protective effect against various stress conditions was shown for an aqueous extract of *R. rosea* (SHR-5; containing rosavin 3.6%, salidroside 1.6% and *p*-tyrosol <0.1%) when applied to preparations of three-day-old larvae of the freshwater snail *Lymnaea stagnalis* before they were exposed to stressors. Preincubation of larvae with increasing concentrations of *R. rosea* extract (ranging from 4.05 to 40.5 μg/mL) for 20 hours resulted in a concentration-dependent protective effect against exposure to lethal heat shock (43°C for 4 minutes) for concentrations above 4.05 μg/mL. A protective effect of *R. rosea* was also demonstrated against other stressors (menadione, copper and cadmium) although the effect was less marked than against lethal heat shock. Further investigations, involving incubation of larvae with *R. rosea* extract at a concentration of 40.5 μg/mL for 20 hours, ruled out induction of the synthesis of heat shock proteins by *R. rosea* extract as a possible mechanism of action for its protective effect against stressors.

**Cognitive effects**

The effects of a 40% ethanol extract of *R. rosea* root on learning and memory were investigated in rats in a battery of tests, including those involving passive avoidance and active avoidance with negative or positive reinforcement. The results of these studies were conflicting. A single oral dose of *R. rosea* root extract of 0.1 mL given 30 minutes before maze training significantly facilitated learning and improved memory at 24 hours after administration (*p* < 0.01 compared with control), whereas doses of 0.02 and 1 mL had no demonstrable effect. In other tests using the staircase method (which trains animals to retrieve food from the top stair), oral administration of *R. rosea* extract 0.1 mL daily for 10 days before training resulted in a significantly greater proportion of trained rats, compared with control (92.3% versus 61.5%, respectively; *p* < 0.05). However, in other tests (including passive avoidance and active avoidance with negative reinforcement methods), this dosage regimen of *R. rosea* root extract had no significant effects on learning and memory. Oral administration of a 40% ethanol extract of *R. rosea* rhizome 0.1 mL daily for 10 days before training also had no effect on electroconvulsive shock-impaired learning and memory in rats.

The CNS depressant effects of a 50% ethanol extract (said to be of *Sedum rosea*; plant part not stated) at doses of 10, 30 and 100 mg/kg body weight intraperitoneally were investigated in mice. Animals were given a single dose of sodium pentobarbital (50 mg/kg body weight, intraperitoneally) 30 minutes after *S. rosea*, and the narcosis time (time between loss and recovery of the righting reflex; the righting reflex is any one of the
neuromuscular responses that restore the body to its upright position when it has been displaced) was measured by a blinded observer. It was reported that S. rosea extract potentiated the effects of pentobarbital, since the time to recovery of the righting reflex was significantly longer for treated mice than for controls. However, this finding is questionable since numerical data and a precise $p$-value were not reported (the $p$-value was given as $p \leq 0.05$).

**Antitumour activities**
Antitumour and antimetastatic effects have been reported for an extract of R. rosea (no further details given) following *in vivo* experiments. In one study, mice transplanted with Ehrlich adenocarcinoma were treated with R. rosea extract 0.5 mL/kg body weight orally daily from day 4 post-transplantation until day 13 or 15 post-transplantation when the animals were sacrificed.\(^{22}\) It was reported that tumour growth was significantly inhibited in treated mice, compared with controls. However, the findings of this study are questionable since the $p$-value given for this finding was $p = 0.05$, and the number of animals involved in the experiment was not stated. A similar experiment involved rats transplanted with metastasising Pliss lymphosarcoma and treated with R. rosea extract according to the same regimen. In this study, the extent of metastasis in treated rats was reported to be 50% that in control animals ($p < 0.01$) and the mass of metastases was significantly less than that in control animals (mean (standard deviation): 142.5 (23.0) and 203.3 (27.0), for treated and control groups, respectively; $p < 0.05$).\(^{22}\) However, again, the sample size for this study was not stated.

In another *in vivo* study, an extract of R. rosea root was tested for its effects on the haematotoxicity of cyclophosphamide in mice transplanted with Ehrlich ascites tumour and Lewis lung carcinoma. After transplantation, mice were treated with cyclophosphamide (100 mg/kg body weight), R. rosea root extract (0.5 mL/kg orally on days 2–8 following transplantation), or both substances.\(^{23}\) Cyclophosphamide, but not R. rosea extract, reduced the numbers of leukocytes and myelokaryocytes to 40–50%, compared with control ($p < 0.05$); when the two substances were given in combination, the numbers of leukocytes and myelokaryocytes were reported to increase by 30% and 16–18%, respectively, although only the change in the former parameter was reported to be statistically significant ($p < 0.05$). R. rosea extract was also reported to have no effect on the colony-forming activity of myelokaryocytes, whereas cyclophosphamide inhibited the proliferation of these cells. In mice given a combination of the two substances, no inhibition of colony-forming activity was observed; this was stated to be statistically significant, compared with values for mice treated with cyclophosphamide alone, although no $p$-value was given.\(^{23}\)
Antimutagenic activities
Antimutagenic activity has been described for extracts of *R. rosea*. Ethanol extracts (20 and 40%) of *R. rosea* were reported to counteract gene mutations induced by various chemical mutagens in the Ames test (*Salmonella typhimurium*).(24) In another *in vitro* experiment, *R. rosea* extract (no further details given) was reported to reduce the yield of bone marrow cells with chromosome aberrations induced *in vivo* (mice) by cyclophosphamide. It is postulated in abstracts of the Russian and Ukrainian literature that *R. rosea* acts as an antimutagen by increasing the efficiency of intracellular DNA repair mechanisms.(25) Since this information is taken from abstracts, confirmation is required.

Cardiovascular activities
Inotropic, anti-arrhythmic and other cardioprotective activities have been described in abstracts of papers published in Russian for *R. rosea* extracts. *R. rosea* extract (no further details given), administered orally, was reported to have an anti-arrhythmic effect in epinephrine (adrenaline)-induced arrhythmia in rat heart,(26) and to prevent the reperfusion-induced decrease in contraction amplitude in isolated perfused rat heart.(27) Since both these effects were reversed by naloxone administered by intravenous infusion, the authors postulated that the observed cardioprotective effects of *R. rosea* extract may be related to stimulation of the endogenous opioid system. This information is taken only from abstracts, so confirmation is required.

Clinical studies
Mental performance
Several clinical trials, with reports published in the English scientific literature, have investigated the effects of preparations containing a dry extract of *R. rosea* rhizome (SHR-5; Swedish Herbal Institute) on stress-induced fatigue and on mental work capacity.

In a randomised, double-blind, placebo-controlled, parallel-group trial, 40 healthy male Indian medical students took tablets containing *R. rosea* dry extract 50 mg, orally twice daily, or placebo, for 20 days during an examination period.(14) At the end of the study, statistically significant improvements in self-assessed mental fatigue, self-assessed general well-being and in one of the psychomotor tests (accuracy of movement versus speed in a maze test) were reported for the treated group, compared with the placebo group (*p* < 0.01, 0.05 and 0.01, respectively). Statistically non-significant differences were reported in several other tests, including another psychomotor test (tapping test) and two mental capacity tests. The findings of the study are limited since no predefined primary outcome measure was
stated, the sample size was small, and the method of randomisation of participants was not adequately described. It was stated in a report of the study that students who received R. rosea extract achieved a higher average mark in the examinations than did placebo recipients (3.47 and 3.20, respectively) and that this indicated the ‘usefulness’ of R. rosea. However, this is a post-hoc outcome measure and was not subjected to statistical testing, therefore the conclusion is not necessarily valid.

A similar study, using a randomised, double-blind, placebo-controlled, crossover design, assessed the effects of R. rosea dry extract 170 mg (containing approximately 4.5 mg salidroside) daily for 14 days on mental performance among 56 healthy Armenian physicians aged 24–35 years undertaking night duty. Participants underwent a battery of tests before and after the 14-day treatment period, at the end of the 14-day wash-out period and, after crossing over to the other study arm for 14 days. It was reported that participants who received R. rosea during the first 14-day period experienced statistically significant improvements in tests for fatigue, compared with placebo recipients, but that there was no difference between the two groups at the end of the crossover period. The investigators use this as evidence of the beneficial effects of R. rosea extract, but this analysis and, therefore, the conclusion is flawed since it takes no account of the second treatment period. Crossover studies are used as a means of increasing the statistical power of a study without increasing the sample size, so ignoring the second treatment period in effect halves the sample size of the study.

As the two studies described above had used doses of R. rosea that were considered to be low, a subsequent randomised, double-blind, placebo-controlled, parallel-group study assessed the effects of two different doses of R. rosea extract on capacity for mental work against a background of fatigue. In this study, 121 healthy male cadets aged 19–21 years received two ($n=41$) or three ($n=20$) capsules containing R. rosea extract (SHR-5, 185 mg contains 4.5 mg salidroside); equivalent to a total of 370 mg (9 mg salidroside) and 555 mg extract (13.5 mg salidroside), respectively, or placebo ($n=40$), as a single dose; a further 20 participants were allocated to a no-treatment control group. Participants underwent a battery of tests at 5 pm before undertaking night duties, and took their allocated study medication at 4 am the next morning, 1 hour before assessment in the test battery a second time.

Participants who received higher dose R. rosea extract performed significantly better than did placebo recipients in all five measures of capacity for mental work in the test battery (scanning for pre-assigned symbols, number of errors
in this test, recall of digit sequences, arrangement of numbers in a grid and number of errors in this test; \( p < 0.05 \) in each test). Statistically significant improvements with lower dose \( R. \) rosea extract, compared with placebo, were observed in only two of the five tests (\( p < 0.01 \) for each). Both doses of \( R. \) rosea extract, compared with placebo, were reported to achieve significantly better scores on an Antifatigue Index (mean (standard deviation): 1.04 (0.29), 1.02 (0.21) and 0.90 (0.32) for lower and higher dose \( R. \) rosea extract and placebo, respectively; \( p < 0.0001 \) for dose of \( R. \) rosea extract versus placebo).\(^{(11)}\)

This study, however, has several methodological flaws. For example, a formal sample size calculation was not carried out, the study had no predefined primary outcome measure, and in the analysis, no adjustment appears to have been made for multiple statistical tests. Further, although the study was said to include random allocation to treatment, the randomisation process as described in a report of the study is inadequate and, additionally, the no-treatment control group was not randomly allocated at all but simply comprised the last 20 cadets to be enrolled into the study.

**Other conditions**

In a double-blind, placebo-controlled, crossover study designed to assess the effects of \( R. \) rosea on hypoxia and oxidative stress, 15 healthy volunteers aged 20–33 years received capsules each containing \( R. \) rosea 447 mg (no further details given), four daily for seven days, or placebo, before undergoing hypoxic exposure (to simulate conditions at an elevation of 4600 m).\(^{(28)}\) (The study also assessed the effects of a supplement claimed to contain dissolved oxygen, although these results are not reported here.) Fourteen participants completed the study. There were no statistically significant differences between \( R. \) rosea and placebo recipients in any of the outcome measures, including arterial capillary blood oxygen (concentration – \( P_cO_2 \)), blood oxyhaemoglobin saturation as assessed using a pulse oximeter on an index finger, and serum lipid peroxide and urine malondialdehyde concentrations (as markers of oxidative stress).\(^{(28)}\) However, given the small sample size for this study, further investigation into the effects of \( R. \) rosea preparations in hypoxia is warranted.

Preliminary studies conducted in China and involving men living and working at high altitude have compared the effects of a multiherbal product containing the related species \( Rhodiola kirilowii \) (Regel) Maxim. with those of acetazolamide.\(^{(29,30)}\)
Side-effects, Toxicity

There is a lack of reliable, accessible information relating to the safety and toxicity of preparations of *R. rosea*. Data from clinical trials of *R. rosea* are extremely limited, and no postmarketing surveillance or other pharmacoepidemiological studies have been identified.

Clinical trials involving small numbers of healthy young (<35 years) volunteers who received *R. rosea* extract (SHR-5) 100 mg for 20 days\(^{(14)}\) or 170 mg for 14 days\(^{(10)}\) (see Clinical studies for further details), have reported that no adverse effects or events were observed during the studies.\(^{(10,14)}\)

Another study in which healthy young (19–21 years) volunteers received *R. rosea* extract (SHR-5) 370 mg (containing 9 mg salidroside) or 555 mg (containing 13.5 mg salidroside) as a single dose also reported that no adverse effects or events were observed.\(^{(11)}\)

*In vitro* experiments have investigated the effects of incubation of three-day-old larvae of the freshwater snail *Lymnaea stagnalis* with different concentrations of an aqueous extract of *R. rosea* (SHR-5; containing rosavin 3.6%, salidroside 1.6% and \(p\)-tyrosol <0.1%) for 24 hours. At a concentration of *R. rosea* extract of 1.35 mg/mL, all larvae were killed; at 405 \(\mu\)g/mL, around 80% were killed.\(^{(19)}\) With a longer period of exposure (up to four days), *R. rosea* extract 81.2 \(\mu\)g/mL did not induce death of exposed larvae, but their development appeared to be retarded: specimens hatched later and were smaller than control specimens, yet no deformations or other abnormalities were observed. Exposure of larvae to *R. rosea* 40.5 \(\mu\)g/mL for up to four days was not associated with any slowing down of growth or development.

Abstracts of papers published in Russian or Ukrainian have described antimutagenic activity for *R. rosea* extracts, although confirmation of these data is required. Antimutagenic activity has been reported for ethanol extracts (20% and 40%) of *R. rosea* in the Ames test (*Salmonella typhimurium*).\(^{(24)}\) A reduction in the yield of bone marrow cells with chromosome aberrations induced *in vivo* (mice) by cyclophosphamide has also been described for an extract of *R. rosea*.\(^{(25)}\)
Contra-indications, Warnings

None documented. In view of the lack of safety data, use of *R. rosea* extracts at dosages higher than those recommended (see Dosage) and/or for longer periods should be avoided.

**Interactions**

No interactions have been documented for *R. rosea*. However, in view of the documented pharmacological effects, whether or not there is potential for clinically important interactions with other medicines with similar or opposing effects should be considered.

**Pregnancy and lactation**

The safety of *R. rosea* has not been established. In view of the lack of information on the use of *R. rosea* during pregnancy and lactation, its use should be avoided during these periods.
The chemistry of *R. rosea* root and rhizome is well documented. The precise chemical constituents responsible for the documented pharmacological activities are not fully understood, although tyrosol and salidroside (rhodioloside) and the phenylpropanoid glycoside constituents (rosin, rosavin, rosarin) (see Constituents) are believed to be important for certain activities.

Over 200 other *Rhodiola* species have been described, and several of these, including *Rhodiola crenulata*, *Rhodiola quadrifida* (Pall.) Fisch. et Mey., *Rhodiola sacra* and *Rhodiola sachalinensis* A. Bor. are used in traditional Asian medical systems, such as Traditional Chinese Medicine.\(^1,31\) Certain constituents of *R. rosea* have also been reported for other *Rhodiola* species. For example, salidroside has been documented for extracts of *R. crenulata*,\(^32\) *R. quadrifida*\(^1\) and *R. sachalinensis* root.\(^33\)

A substantial body of research investigating the pharmacological and clinical properties of *R. rosea* has been undertaken, although much this work has been published in the Russian scientific literature, making it difficult to access. A limited number of reports of *in vitro* and animal studies published in the English scientific literature has described certain adaptogenic and other effects, such as antitumour and antimetastatic activities, for extracts of *R. rosea*, although further investigation is required to confirm these findings. Well-designed clinical trials of *R. rosea* root/rhizome extracts published in the English (or other European language) scientific literature are lacking. To date, clinical investigations are limited to a small number of single-dose or short-term trials (lasting less than three weeks) involving small numbers of healthy volunteers.

In view of the lack of safety and toxicity data, excessive use (higher than recommended dosages and/or for long periods of time) of *R. rosea* should be avoided.

Pharmacists and other healthcare professionals should be aware that herbal products containing rhodiola are readily available over the internet and from retail outlets; such products are often promoted as being beneficial in supporting mental and physical performance, emotional balance, cardiovascular health, resistance to infection, male sexual function and in increasing ability to cope with stressful situations. There are no licensed rhodiola products available in the UK, so the quality of commercially available products is not assured.
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Species (Family)

*Rheum officinale* Baill. and *R. palmatum* L. (Polygonaceae)
Synonym(s)
Chinese Rhubarb, other *Rheum* species, e.g. *Rheum tanguticum* Maxim. & Reg., *Rheum emodi* Wall. (Indian Rhubarb) and *Rheum rhaponticum* L. (Garden Rhubarb)
Part(s) Used
Rhizome, root
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)
BHP 1996\(^{(G9)}\)
BP 2002\(^{(G71)}\)
Complete German Commission E\(^{(G3)}\)
ESCOP 1999\(^{(G52)}\)
Martindale 33rd edition\(^{(G67)}\)
PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Ph Eur 2004\(^{(G72)}\)
WHO volume 1 1999\(^{(G63)}\)
Legal Category (Licensed Products)

GSL$^{(G37)}$
Constituents


**Hydroxyanthracenes**
Primarily anthraquinone O-glycosides (anthraglycosides) of aloe–emodin, emodin, chrysophanol and physcion; dianthrone glycosides of rhein (sennosides A and B) and their oxalates; heterodianthrones including palmidin A (aloe–emodin, emodin), palmidin B (aloe–emodin, chrysophanol), palmidin C (chrysophanol, emodin), sennidin C (rhein, aloe–emodin), rheidin B (rhein, chrysophanol), and reidin C (rhein, physcion); free anthraquinones mainly aloe–emodin, chrysophanol, emodin, physcion and rhein.

**Tannins**
Hydrolysable and condensed including glucogallin, free gallic acid, (−)-epicatechin gallate and catechin.

**Other constituents**
Calcium oxalate, fatty acids, rutin, resins, starch (about 16%), stilbene glycosides, carbohydrates, volatile oil (trace) with more than 100 components.
Rhubarb is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that it can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\(^{(G16)}\) Rhubarb stems are commonly eaten as a food. In the USA, rhubarb is permitted for food use.\(^{(G65)}\)
Herbal Use

Rhubarb has been used traditionally both as a laxative and an antidiarrhoeal agent.
Dosage

*Rhizome/root*
0.2–1.0 g.
Pharmacological Actions

The laxative action of anthraquinone derivatives is well recognised (see Senna). Rhubarb also contains tannins, which exert an astringent action. At low doses, rhubarb is stated to act as an antidiarrhoeal because of the tannin components, whereas at higher doses it exerts a cathartic action.\(^{(G42)}\)
Side–effects, Toxicity

See Senna for side–effects and toxicity associated with anthraquinone–containing drugs. Rhubarb leaves are toxic because of the oxalic acid content and should not be ingested. A case of anaphylaxis following rhubarb ingestion has been documented.\(^{(G51)}\)
Contra–indications, Warnings\textsuperscript{(G20)}

See Senna for contra–indications and warnings associated with anthraquinone–containing drugs. The astringent effect of rhubarb may exacerbate, rather than relieve, symptoms of constipation.\textsuperscript{(1)} It has been stated that rhubarb should be avoided by individuals suffering from arthritis, kidney disease or urinary problems.\textsuperscript{(G42)}

\textbf{Pregnancy and lactation}
It is stated that rhubarb should be avoided during pregnancy.\textsuperscript{(G42)} See Senna for contra–indications and warnings regarding the use of stimulant laxatives during pregnancy and lactation.
Pharmaceutical Comment

The chemistry of rhubarb is characterised by the anthraquinone derivatives. The laxative action of these compounds is well recognised and justifies the use of rhubarb as a laxative. As with all anthraquinone-containing preparations, the use of non-standardised products should be avoided because their pharmacological effect will be variable and unpredictable.


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Rosemary
Species (Family)

Rosmarinus officinalis L. (Labiatae)
Synonym(s)
Part(s) Used

Leaf, twig
Pharmacopoeial and Other Monographs

BP 2002\(^{(G71)}\)

BHP 1996\(^{(G9)}\)

Complete German Commission E\(^{(G3)}\)

ESCOP 1997\(^{(G52)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)

Ph Eur 2004\(^{(G72)}\)
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents
See General References G2 G41 G52 G58.

**Flavonoids**
Include diosmetin, diosmin, genkwanin and derivatives, luteolin and derivatives, hispidulin, nepetin, nepitrin and apigenin.

**Phenols**
Caffeic, chlorogenic, labiatic, neochlorogenic and rosmarinic acids.

**Volatile oil**
1–25%. Components vary according to chemotype. Composed mainly of monoterpene hydrocarbons including α- and β-pinenes, camphene and limonene, together with 1,8-cineole, borneol, camphor (20–50% of the oil), linalool, verbinol, terpineol, 3-octanone and isobornyl acetate.

**Terpenoids**
Carnosol, carnosolic acid, rosmanol (diterpenes);\(^1\) oleanolic and ursolic acids (triterpenes).
Food Use

Rosemary herb and oil are commonly used as flavouring agents in foods. Rosemary is listed by the Council of Europe as a source of natural food flavouring (category N2). This category indicates that rosemary can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\(^{(G16)}\) In the USA, rosemary is listed as GRAS (Generally Recognised As Safe).\(^{(G65)}\)
Herbal Use\(^{(G2\ G4\ G7\ G32\ G52)}\)

Rosemary is stated to act as a carminative, spasmolytic, thymoleptic, sedative, diuretic and antimicrobial.\(^{(G7)}\) Topically, rubefacient, mild analgesic and parasiticide properties are documented.\(^{(G7)}\) Traditionally rosemary is indicated for flatulent dyspepsia, headache, and topically for myalgia, sciatica, and intercostal neuralgia. The German Commission E approved internal use for dyspeptic complaints and external use as supporting therapy for rheumatic diseases and circulatory problems.\(^{(G3)}\)
Dosage

*Dried leaf/twig*
2–4 g or by infusion three times daily;\(^{(G7)}\) 4–6 g daily; external use 50 g for one bath.\(^{(G3)}\)

*Liquid extract*
2–4 mL (1 : 1 in 45% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

In vitro and animal studies

Antimicrobial activity
Antibacterial and antifungal activities in vitro have been reported for rosemary oil.\(^2\) Rosemary herb is an effective antimicrobial agent against *Staphylococcus aureus* in meat and against a wide range of bacteria in laboratory media.\(^1\) Antimicrobial activity has been documented for the oil towards moulds, and Gram–positive and Gram–negative bacteria\(^1\) including *S. aureus*, *S. albus*, *Vibrio cholerae*, *Escherichia coli* and corynebacteria.\(^3\) Carnosol and ursolic acid have inhibited a range of food spoilage microbes (*S. aureus*, *E. coli*, *Lactobacillus brevis*, *Pseudomonas fluorescens*, *Rhodotorula glutinis* and *Kluyveromyces bulgaricus*). Activity was comparable to that of known antioxidants butylated hydroxyanisole (BHA) and butylated hydroxytoluene (BHT), and correlated with the respective antioxidant properties of the two compounds (carnosol > ursolic acid).\(^1\)

Antiviral activity
A dried 95% ethanol extract of rosemary (2–100 μg/mL) inhibited in vitro formation of herpes simplex virus type 2 plaques from 2 to 100% in a concentration–dependent manner. Carnosolic acid had activity against human immunodeficiency virus type 1 (HIV-1) protease (IC\(_{50}\) value 0.08 μg/mL) when assayed against HIV-1 virus replication (IC\(_{90}\) value 0.32 μg/mL).\(^4,5\) Carnosolic acid was cytotoxic to lymphocytes with a TC\(_{90}\) on H9 lymphocytes of 0.36 μg/mL.

Antitumour activity
An extract of rosemary (precipitate from aqueous phase of 70% alcohol extract) inhibited KB cells by 87% when applied at a concentration of 50 μg/mL.\(^5\) The volatile oil (1.2–300 μg/mL) was reported to be toxic to L-1210 leukaemia cells.\(^6\) Topical administration of a methanol extract 5 minutes prior to application of carcinogens to the dorsal surface of CD-1 mice reduced the irritation and promotion of tumours. Application of rosemary extract (1.2 mg and 3.6 mg) prior to \(^{3}\)H]-benzo(a)pyrene reduced the formation of metabolite–DNA adducts by 30% and 54%, respectively.\(^7\) In rats, dietary supplementation with 1% rosemary extract for 21 weeks reduced the development of dimethylbenz(a)anthracene mammary carcinoma in the treated group, compared with the control group (40% versus 75%, respectively).\(^8\)

Antispasmodic and anticonvulsant activities
Rosemary oil, 1,8-cineole, and bornyl acetate have exerted a spasmolytic action in both smooth muscle (guinea-pig ileum) and cardiac muscle (guinea-pig atria) preparations, with the latter more sensitive. In smooth muscle this spasmolytic effect has been attributed to antagonism of acetylcholine, with borneol considered the most active component of the oil. The spasmolytic action of rosemary oil is preceded by a contractile action, which is attributed to the pinene components. α-Pinenes and β-pinenes have exhibited a spasmogenic activity towards smooth muscle, with no effect on cardiac muscle.

Spasmolytic action in vivo (guinea-pigs) has been demonstrated by rosemary oil (administered intra venously) via a relaxant action on Oddi’s sphincter contracted by morphine. Activity increased with incremental doses of oil until an optimum dose was reached (25 mg/kg) at which the unblocking effect was immediate. Further increases in dose reintroduced a delayed response time. Smooth muscle–stimulant and analgesic actions have been documented for a rosmaricine derivative.

The volatile oil of rosemary inhibited contractions of rabbit tracheal smooth muscle induced by acetylcholine, and inhibited contraction of guinea-pig tracheal smooth muscle induced by histamine. The oil also inhibited contractions in both preparations induced by high potassium concentrations. Contractions of rabbit and guinea-pig tracheal smooth muscle induced by acetylcholine and histamine, respectively, were inhibited by rosemary oil in calcium ion–free solution. It was suggested that the oil has calcium antagonist activity. A 30% ethanol extract of rosemary produced a spasmolytic effect on guinea-pig ileum, as demonstrated by measuring the increase in the ED$_{50}$ of acetylcholine (4.9 μg/L after addition of 2.5 mL extract and 25.1 μg/L after addition of 10 mL extract). An increase in the ED$_{50}$ for histamine from 8.1 μg/L to 44.6 μg/L, respectively, was noted for the same doses of extract.

Noradrenaline (norepinephrine)- and potassium ion–induced contractions of rabbit aortic rings were significantly reduced by rosemary oil 0.48 mg/mL and 0.64 mg/mL, respectively. It was proposed that action was by a direct vascular smooth muscle effect.

**Anti-inflammatory activity**

Complement activation and subsequent triggering of the arachidonic acid cascade are thought to play an important role in the early phase of shock. An intact complement system is required for the formation of vasoactive prostanoids (prostacyclin, thromboxane A$_2$), arterial hypotension and
The effect of rosmarinic acid on endotoxin–induced haemodynamic and haematological changes has been studied in a rabbit model of circulatory shock. Rosmarinic acid (20 mg/kg, intravenous) was found to suppress the endotoxin–induced activation of complement, formation of prostacyclin, hypotension, thrombocytopenia, and the release of thromboxane $A_2$. Unlike non–steroidal anti–inflammatory drugs (NSAIDs), the mode of action by which rosmarinic acid suppresses prostaglandin formation does not involve interference with cyclooxygenase activity or prostacyclin synthetase. Activity has been attributed to inhibition of complement factor C3 conversion to activated complement components, which mediate the inflammatory process. Rosmarinic acid has inhibited carrageenan–induced rat paw oedema, and passive cutaneous anaphylaxis, also in rats (ID$_{50}$ 1 mg/kg, intravenously; 10 mg/kg, intramuscularly).

Topical application of rosmarinic acid (5%) to rhesus monkeys reduced gingival plaque indices when compared with placebo. A methanol extract of herb (3.6 mg) applied topically to CD-1 mice twice daily for four days inhibited skin inflammation and hyperplasia caused by 12-O-tetradecanoylphorbol–13-acetate (TPA). A similar extract inhibited both TPA- and arachidonic acid–induced inflammation as well as TPA-induced hyperplasia.

**Anti–hepatotoxic activity**

A lyophilised aqueous extract of rosemary significantly reduced hepatotoxicity of t-butylperoxide to rat hepatocytes in vitro, significantly decreasing malonaldehyde formation, release of lactic acid dehydrogenase and aspartate aminotransferase. Pretreatment of rats with an aqueous extract (1 mg of lyophilisate equivalent to 7 mg young shoots) 30 minutes prior to exposure to carbon tetrachloride, resulted in a 72% decrease in plasma glutamic–pyruvic transaminase. Rosemary extract supplementation in the diet of rats enhanced the activity of GSH-transferase and NAD(P)H-quinone reductase.

**Cholagogic activity**

A lyophilised ethanolic extract (1 mg) of young shoots at doses of 0.1, 1.0 and 2.0 g/kg was injected into the jugular vein of common bile duct–cannulated Sprague–Dawley rats infused with sodium taurocholate. A significant, rapid increase in bile flow (114%) was achieved with maximum effect in 30 minutes. The extract of young shoots was significantly more active in stimulating bile flow than a similar extract of whole plant. A rapid increase in bile secretion was observed (138% in 40 minutes) in cannulated guinea–pigs given an aqueous–ethanol extract (15%).
**Antioxidant activity**

A number of extracts and constituents of rosemary have been shown to have antioxidant activity.\(^{52}\) An antioxidant action, demonstrated by inhibition of chemiluminescence and hydrogen peroxide generation from human granulocytes, has been reported for rosmarinic acid.\(^{15}\)

Lipophilic and hydrophobic fractions of rosemary showed activity which was attributed to the diterpenes carnosol, carnosolic acid and rosmanol inhibiting superoxide anion production in the xanthine/xanthine oxidase system.\(^{19}\) These diterpenes at concentrations of 3–30 μmol/L also completely inhibit mitochondrial and microsomal lipid peroxidation induced by NADPH or NADPH oxidation.\(^{19}\)

The complement–inhibiting and antioxidant properties of rosmarinic acid are not thought to adversely affect the chemotaxic, phagocytic and enzymatic properties of polymorphonuclear leukocytes.\(^{20}\)

**Other activities**

A hyperglycaemic effect was observed in glucose–loaded rats treated with a solution of rosemary oil (925 mg/kg, intramuscular).\(^{23}\) In rabbits with alloxan–induced diabetes given rosemary oil (25 mg/kg, intramuscular) 6 hours after fasting, plasma glucose concentrations increased by 17% 6 hours later.

Pretreatment with rosmarinic acid (20 mg/kg and 10 mg/kg, intravenously) has been reported to inhibit the development of adult respiratory distress syndrome (ARDS) in a rabbit model.\(^{20}\) This action can be attributed to both the antioxidant and anticomplement activities of rosmarinic acid.\(^{20}\)

The ability to reduce capillary permeability has been described for diosmin.\(^{41}\) Activity reportedly exceeds that exhibited by rutin.\(^{41}\)

An increase in locomotor activity has been observed in mice following either inhalation or oral administration of rosemary oil.\(^{21}\) The increase in activity paralleled a dose-related increase in serum 1,8–cineole level. Biphasic elimination of 1,8–cineole from the blood was observed \(t_{1/2} = 6\) minutes, \(t_{1/2} = 45\) minutes).\(^{21}\)

In rats, antigonadotrophic activity has been documented for oxidation products of rosmarinic acid administered intramuscularly.\(^{22}\) Activity was determined by suppression of pregnant mares’ serum–induced increase in ovarian and uterine weights. Concentrations of \(10^{-7}\) mol/L of the flavonoids nepitrin and nepetin inhibited aldose–reductase activity in homogenised rat
eye lenses by 31\%.^G52
Rosemary oil is stated to be non–irritating and non–sensitising when applied to human skin, but moderately irritating when applied undiluted to rabbit skin. Bath preparations, cosmetics and toiletries containing rosemary oil may cause erythema and dermatitis in hypersensitive individuals. Photosensitivity has been associated with the oil.

Rosmarinic acid exhibits low toxicity (an LD$_{50}$ in mice is stated as 561 mg/kg for intravenous administration) and is rapidly eliminated from the circulation ($t_{1/2} = 9$ minutes following intravenous administration). Transient cardiovascular actions become pronounced at intravenous doses exceeding 50 mg/kg. Acute LD$_{50}$ values quoted include 5 mL/kg (rat, oral) and >10 mL/kg (rabbit, dermal).

Diosmin is reportedly less toxic than rutin. No mortality was seen in Wistar rats and Swiss mice given single intraperitoneal doses of 2 g/kg of aqueous alcoholic rosemary extract (15%).
Contra–indications, Warnings

Topical preparations containing rosemary oil should be used with caution by hypersensitive individuals. Rosemary oil contains 20–50% camphor; orally, camphor readily causes epileptiform convulsions if taken in sufficient quantity.\(^{(G58)}\)

**Pregnancy and lactation**

Rosemary is reputed to be an abortifacient\(^{(G30)}\) and to affect the menstrual cycle (emmenagogue).\(^{(G48)}\) In view of its common culinary use, rosemary should not be ingested in amounts greatly exceeding those normally encountered in foods.
In addition to the well-known culinary uses of rosemary, various medicinal properties are also associated with the herb. Documented antibacterial, anti-inflammatory and spasmolytic actions, which support the traditional uses of the herb, are attributable to the essential oil. Anticomplement and antioxidant activities documented for rosmarinic acid have generated considerable interest in a potential preventative use against endotoxin shock and adult respiratory distress syndrome. A method for the isolation (TLC, thin-layer chromatography) and subsequent identification (HPLC, high-performance liquid chromatography) of rosmarinic acid has been proposed. Rosemary should not be used by epileptic patients in doses greatly exceeding amounts used in food.
References


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Species (Family)

*Salvia officinalis* L. (Labiatae)
Synonym(s)
Dalmatian Sage, Garden Sage, True Sage
Red Sage refers to *Salvia haematodes* Wall.
Greek Sage refers to *Salvia triloba* (1)
Spanish Sage refers to *Salvia lavundaefolia*
Part(s) Used

Leaf
Pharmacopoeial and Other Monographs

BHP 1996\textsuperscript{(G9)}

BP 2002\textsuperscript{(G71)}

Complete German Commission E\textsuperscript{(G3)}

ESCOP 1996\textsuperscript{(G52)}

Martindale 33rd edition\textsuperscript{(G67)}

PDR for Herbal Medicines 2nd edition\textsuperscript{(G36)}

Ph Eur 2004\textsuperscript{(G72)}
Legal Category (Licensed Products)

GSL\textsuperscript{(G37)}
Constituents
See General References G2 G22 G41 G52 G58 G62 G64.

**Acids**
Phenolic – caffeic, chlorogenic, ellagic, ferulic, gallic and rosmarinic.\(^{(2)}\)

**Flavonoids**
5-Methoxysalvigenin.

**Terpenes**
Monoterpene glycosides. Diterpenes, abietanes including carnosic acid and derivatives, e.g. carnosol. Triterpenes, oleanolic acid and derivatives.

**Tannins**
3–8%. Hydrolysable and condensed.\(^{(2,3)}\)

**Volatile oil**
1–2.8%. Pharmacopoeial standard not less than 1.0% cut herb.\(^{(G15 \ G28)}\)
Major components are α- and β-thujones (35–50%, mainly α). Others include 1,8-cineole, borneol, camphor, caryophyllene, linalyl acetate and various terpenes.\(^{(4,5)}\)

It has been noted that commercial sage may be substituted with *Salvia triloba*.\(^{(1)}\) In contrast to *S. officinalis*, the principal volatile oil component of *S. triloba* is 1,8-cineole, with α-thujone only accounting for 1–5%.\(^{(1)}\) Compared to *S. officinalis*, volatile oil yield of various *Salvia* species is lower, with lower total ketone content and higher total alcohol content.\(^{(6)}\)
Food Use

Sage is commonly used as a culinary herb. It is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that sage can be added to foodstuffs providing the concentration of thujones (α and β) present in the final product does not exceed 0.5 mg/kg, with the exceptions of alcoholic beverages (10 mg/kg), bitters (35 mg/kg), food containing sage (25 mg/kg) and sage stuffing (250 mg/kg). In the USA, sage is listed as GRAS (Generally Recognised As Safe).
Sage is stated to possess carminative, antispasmodic, antiseptic, astringent and antihidrotic properties. Traditionally, it has been used to treat flatulent dyspepsia, pharyngitis, uvulitis, stomatitis, gingivitis, glossitis (internally or as a gargle/mouthwash), hyperhidrosis, and galactorrhoea. The herbals of Gerard, Culpeper and Hill credit sage with the ability to enhance memory.\(^7\) The German Commission E approved internal use for dyspeptic symptoms and excessive perspiration, and external use for inflammation of mucous membranes of mouth and throat.\(^3\)
Dosage

**Leaf**
1–4 g or by infusion three times daily;\(^{(G7)}\) 4–6 g daily. \(^{(G3)}\)

**Liquid extract**
1–4 mL (1:1 in 45% alcohol) three times daily.\(^{(G7)}\)

**Gargles, rinses**
2.5 g/100 mL water.\(^{(G3)}\)
Pharmacological Actions

In vitro and animal studies

Hypotensive activity in anaesthetised cats, CNS-depressant action (prolonged barbiturate sleep) in anaesthetised mice, and an antispasmodic action in vitro (guinea-pig ileum) have been reported for a sage extract\(^8\) and for the essential oil.\(^9\)

Antispasmodic activity

Inhibition of contractions induced by acetylcholine, histamine, serotonin and barium chloride by 60–80% has been noted for a total sage extract, with lesser activity exhibited by a total flavonoid extract.\(^8\) An initial spasmogenic action exhibited by low doses of sage oil has been attributed to the pinene content.\(^9\) Antispasmodic activity in vivo (guinea-pigs) has been reported for sage oil administered intravenously, which released contraction of Oddi’s sphincter induced by intravenous morphine.\(^5\)

Anticholinesterase activity

Early herbals claim that sage enhances the memory.\(^7\) The anticholinesterase activity of several Salvia species and their constituents have been investigated in the search for new drugs for the treatment of Alzheimer’s disease. The inhibition of anticholinesterase in vitro by an ethanolic extract of S. officinalis (2.5 mg/mL) was 68%, and by oils of S. officinalis and S. lavandulaefolia (0.1 μg/mL) was 52% and 63%, respectively.\(^10\) The IC\(_{50}\) value of S. lavandulaefolia oil is reportedly 0.03 μg/mL.\(^11\) The monoterpenes 1,8-cineole and α-pinene from the oil have been identified as the inhibitors of acetylcholin esterase with IC\(_{50}\) values of 0.67 and 0.63 mmol/L, respectively.\(^11\) Rats given S. lavandulaefolia oil (20 μL or 50 μL for five days) were sacrificed, and acetylcholinesterase activity assessed for striatum, cortex and hippocampus of brain left hemisphere.\(^12\) At the lower dose, there was a decrease in acetyl cholinesterase activity in the striatum, but not in the hippocampus or cortex of treated rats. At the higher dose, there was a decrease in striatal acetylcholinesterase activity. It was concluded that the oil inhibited acetylcholinesterase in selective areas of the brain.

Hypoglycaemic activity

Hypoglycaemic activity in vivo (rabbits) has been reported for S. lavandulaefolia.\(^13\) and for mixed phytotherapy preparations containing various Salvia species, including S. officinalis.\(^14\) Activity in normoglycaemic, hypoglycaemic and in alloxan–diabetic rabbits was observed, although no change in insulin concentrations was noted.\(^13\)
Antimicrobial and antiviral activity
Antimicrobial activity of the volatile oil has been attributed to the thujone content.\(^4\) Antimicrobial activity \textit{in vitro} was noted against \textit{Escherichia coli}, \textit{Shigella sonnei}, \textit{Salmonella} species, \textit{Klebsiella ozanae} (Gram–negative), \textit{Bacillus subtilis} (Gram–positive), and against various fungi (\textit{Candida albicans}, \textit{C. krusei}, \textit{C. pseudotropicalis}, \textit{Torulopsis glabrata}, \textit{Cryptococcus neoformans}).\(^{15}\) No activity was observed versus \textit{Pseudomonas aeruginosa}.\(^4\)

Microencapsulation of sage oil into gelatin–acacia capsules introduced a lagtime with respect to antibacterial activity and inhibited antifungal activity.\(^4\) Diterpene constituents of \textit{S. officinalis} are reported to be active against vesicular stomatitis virus.\(^{G52}\)

Other activities
An aqueous ethanolic extract of sage (50%) strongly inhibited collagenolytic activity of \textit{Porphyromonas gingivitis}.\(^{G52}\) In addition to anti cholinesterase activity, other biological activities have relevance in the treatment of Alzheimer’s disease. In this context, \textit{S. lavanadulaefolia} and its individual constituents have been assessed for antioxidant, anti–inflammatory and oestrogenic activities.\(^{11}\) An ethanolic extract of dried herb (5 mg/mL) and the monoterpenes α- and β-pinene and 1,8–cineole (0.1 mol/L) inhibited bovine brain liposome peroxidate activity. Anti–inflammatory activity was demonstrated by weak inhibition of thromboxane B\(_2\) and leukotriene B\(_4\) synthesis, and possible oestrogenic activity of sage oil (0.01 mg/mL) and geraniol (0.1–2 mmol/L), demonstrated by induction of β-galactosidase in yeast cells.

Other species
Various activities in rats, mice and rabbits have been reported for a related species, \textit{S. haematodes} Wall. (commonly known as red sage), including wound–healing, anti–inflammatory, analgesic, anticonvulsant and hypotensive, and positive inotropic and chronotropic actions \textit{(in vitro)}.\(^{16,17}\) \textit{In vivo} studies have indicated different activities for \textit{S. triloba} and \textit{S. verbenaca}, compared with \textit{S. officinalis}.\(^8\)

Clinical studies
Excessive sweat induced by pilocarpine was inhibited by a dialysate of an aqueous extract of fresh sage.\(^{G52}\) In an open study, 40 patients were given dried aqueous extract of sage (440 mg, equivalent to 2.6 g herbs) and 40 were given infusion of sage (4.5 g herb daily). Reduction of sweat (less than 50%) was achieved in both groups of patients with idiopathic hyperhidriosis.\(^{G52}\) It should be noted, however, that this study did not include a control group.
A double-blind, placebo-controlled, crossover study involving 20 healthy volunteers compared the effects of 50 μL, 100 μL and 150 μL of *S. lavandulaefolia* oil and sunflower oil.\(^{(12)}\) Cognitive assessment indicated improvements in both immediate and delayed word recall scores, coupled with decrements in accuracy and speed of attention, with sage oil 50 μL. At this dose, self-related alertness at 2.5 hours and calmness at 4 hours and 6 hours were reported to be reduced. The results suggest that the effects of sage oil in modulating mood and cognition are worth further investigation.
Side-effects, Toxicity

A case of human poisoning has been documented following ingestion of sage oil for acne.\(^{(18)}\) Convulsant activity in both humans and animals has been documented for sage oil.\(^{(19,20)}\) In rats, the subclinical, clinical and lethal doses for convulsant action of sage oil are estimated as 0.3, 0.5, and 3.2 g/kg.\(^{(19)}\) This toxicity has been attributed to the ketone terpenoids in the volatile oil, namely camphor and thujone. Acute LD\(_{50}\) values for sage oil are documented as 2.6 g/kg in rats for oral administration and 5 g/kg in rabbits for intradermal administration.\(^{(21)}\) \textit{S. officianalis} has no mutagenic or DNA-damaging activity in either the Ames test or \textit{Bacillus} rec-assay.\(^{(G52)}\)

Sage oil is reported to be a moderate skin irritant\(^{(21)}\) and is not recommended for aromatherapy.\(^{(G58)}\)
Contra-indications, Warnings

Sage oil is toxic (due to the thujone content) and should not be ingested. *S. lavandulaefolia* oil has a much lower content of thujone than *S. officinalis* oil.\(^{(12)}\) In view of the toxicity of the essential oil, sage extracts should be used with caution and not ingested in large amounts. Sage may interfere with existing hypoglycaemic and anticonvulsant therapies, and may potentiate sedative effects of other drugs.

**Pregnancy and lactation**

Sage is contra-indicated during pregnancy. Traditionally, it is reputed to be an abortifacient and to affect the menstrual cycle.\(^{(G30)}\) The volatile oil contains a high proportion of α- and β-thujones, which are known to be abortifacient and emmenagogic.
Pharmaceutical Comment

The characteristic components of sage to which its traditional uses can be attributed are the volatile oil and tannins. However, the oil contains high concentrations of thujone, a toxic ketone and should not be ingested. Sage is commonly used as a culinary herb and presents no hazard when ingested in amounts normally encountered in foods. However, extracts of the herb should be used with caution and should not be ingested in large amounts or over prolonged periods. *S. lavandulaefolia* oil is being investigated for symptomatic treatment of Alzheimer’s disease.\(^{(11)}\) However, at present, there is a lack of well-designed clinical studies investigating the reputed effects of sage.
12. Houghton PJ. Personal communication.
19. Millet Y. Experimental study of the toxic convulsant properties of commercial preparations of essences of sage and hyssop. *Electroencephal

Sarsaparilla
Species (Family)

*Smilax* species (Liliaceae) including

i. *Smilax aristolochiifolia* Mill.

ii. *Smilax regelii* Killip & Morton

iii. *Smilax ornata* Hook. f.

iv. *Smilax febrifuga* Kunth
Synonym(s)

Ecuadorian Sarsaparilla, Sarsa, Smilax

i. Mexican Sarsaparilla

ii. Honduras Sarsaparilla

iii. Jamaican Sarsaparilla

iv. Ecuadorian Sarsaparilla
Part(s) Used

Rhizome, root
Pharmacopoeial and Other Monographs

BHC 1992\(^{G6}\)

BHP 1996\(^{G9}\)

Martindale 33rd edition\(^{G67}\)

PDR for Herbal Medicines 2nd edition\(^{G36}\)
Legal Category (Licensed Products)

GSL\textsuperscript{(G37)}
Constituents

Saponins
About 2%. Sarsasapogenin (parigenin), smilagenin, diosgenin, tigogenin, asperagenin, laxogenin from various species,\(^1\) sarsasaponin (parillin), smilasaponin (smilacin) and sarsaparilloside.

Other constituents
Caffeoylshikimic acid, ferulic acid, shikimic acid, kaempferol, quercetin, phyto sterols (e.g. β-sitosterol, stigmasterol, pollinastanol), resin, starch, volatile oil (trace) and cetyl alcohol.
Food Use

Sarsaparilla is listed by the Council of Europe as a natural source of food flavouring (category N4). This category indicates that the use of sarsaparilla as a flavouring agent is recognised but that there is insufficient information available to further classify it into categories N1, N2 or N3.\(^{(G16)}\) Sarsaparilla has been used as a vehicle and flavouring agent for medicaments,\(^{(G45)}\) and is widely employed in the manufacture of non-alcoholic beverages.\(^{(G59)}\) In the USA, sarsaparilla is permitted for food use.
Herbal Use

Sarsaparilla is stated to possess antirheumatic, antiseptic and antipruritic properties. Traditionally, it has been used for psoriasis and other cutaneous conditions, chronic rheumatism, rheumatoid arthritis, as an adjunct to other treatments for leprosy, and specifically for psoriasis. (G6 G7 G8 G64)
Dosage

**Dried root**
1–4 g or by decoction three times daily.\(^{G6}\)

**Sarsaparilla Liquid Extract**
(BP 1898) 8–15 mL (1 : 1 in 20% alcohol, 10% glycerol).
Pharmacological Actions

*In vitro* and animal studies

Anti-inflammatory\(^2\) and hepatoprotective\(^3\) effects have been shown in rats.

Clinical studies

Improvement of appetite and digestion\(^4\) as well as a diuretic\(^4,5\) action have been reported. Limited clinical data utilising extracts indicate improvement in psoriasis;\(^6\) the extract has also been used as an adjuvant for the treatment of leprosy.\(^7\)
None documented for sarsaparilla. Large doses of saponins are reported to cause gastrointestinal irritation resulting in diarrhoea and vomiting. Although haemolytic activity has been documented for the saponins,\(^{(G62)}\) they are not harmful when taken by mouth and are only highly toxic if injected into the bloodstream.\(^{(G59)}\)
Contra-indications, Warnings

None documented for sarsaparilla. In view of the possible irritant nature of the saponin constituents, excessive ingestion should be avoided.

Pregnancy and lactation

There are no known problems with the use of sarsaparilla during pregnancy and lactation. However, in view of the possible irritant nature of the saponin components, excessive ingestion should be avoided.
Pharmaceutical Comment

Phytochemical studies on sarsaparilla have focused on the nature of the steroidal saponin constituents, with limited information available regarding additional constituents. No documented scientific evidence was found to justify the herbal uses. No toxicity data were located, although large doses may be irritant to the gastrointestinal mucosa and should, therefore, be avoided.

Sarsaparilla saponins have been used in the partial synthesis of cortisone and other steroids. Several related *Smilax* species native to China are used to treat various skin disorders.\(^{(G41)}\)
References


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Sassafras
Species (Family)

*Sassafras albidum* (Nutt.) Nees (Lauraceae)
Synonym(s)
Ague Tree, Cinnamon Wood, Saloop, *Sassafras varifolium* (Salisb.) Kuntze, *Sassafras officinale* Nees & Eberm., Saxifrax
Part(s) Used

Inner root bark
Pharmacopoeial and Other Monographs

BHP 1983\textsuperscript{(G7)}

Martindale 33rd edition\textsuperscript{(G67)}

PDR for Herbal Medicines 2nd edition\textsuperscript{(G36)}
Legal Category (Licensed Products)

Sassafras is not permitted for use in medicinal products.
Constituents
See General References G2 G22 G41 G48 G64.

**Alkaloids**
Isoquinoline–type about 0.02%. Boldine, isoboldine, norboldine, cinnamolaurine, norcinnamolaurine and reticuline.

**Volatile oils**
5–9%. Safrole as major component (80–90%), others include anethole, apiol, asarone, camphor, caryophyllene, coniferaldehyde, copaene, elemicin, eugenol, 5-methoxyeugenol, menthone, myristicin, α-pinene, α- and β-phellandrene, piperonylacrolein and thujone.

**Other constituents**
Gum, mucilage, lignans (sesamin, desmethoxyaschantin), resin, sitosterol, starch, tannins and wax.
Food Use

Sassafras oil was formerly used as flavouring agent in beverages including root beer.\(^{(G58)}\) However, in the 1960s safrole, the major component of the volatile oil, was reported to be carcinogenic.\(^{(G58)}\) The use of safrole in foods is now banned, and its use in toilet preparations controlled.\(^{(G45)}\) In the USA, safrole–free sassafras extract, leaf and leaf extract are approved for food use. In 1976, the US Food and Drugs Administration (FDA) banned interstate marketing of sassafras for sassafras tea.\(^{(G22)}\)
Herbal Use

Sassafras is stated to possess carminative, diaphoretic, diuretic, dermatologic and antirheumatic properties. Traditionally, it has been used for cutaneous eruptions, gout and rheumatic pains. (G2 G7 G64)
Dosage

*Bark*
2–4 g or by infusion three times daily.\(^{(G7)}\)

*Liquid extract*
2–4 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

Studies have concentrated on investigating the toxicity associated with the bark. However, aqueous and alcoholic extracts have been reported to elicit ataxia, hypersensitivity to touch, CNS depression and hypothermia in mice.\(^{(1)}\) Both inhibition and induction of hepatic microsomal enzymes have been documented for safrole.\(^{(2,3)}\) Enzyme–inducing activity was found to be a transient phenomenon, with activity falling after the onset of hepatic toxicity (see Side–effects, Toxicity).\(^{(2)}\) Safrole is reported to induce both cytochrome P488 and P450 activities. Sassafras oil has been used as a topical antiseptic, pediculicide and carminative.\(^{(4)}\)
The toxicity of sassafras is attributable to the volatile oil, and in particular to the safrole content. It is estimated that a few drops of sassafras oil are sufficient to kill a toddler and as little as one teaspoonful has proved fatal in an adult. Symptoms of poisoning are described as vomiting, stupor and collapse. High doses may cause spasm followed by paralysis. Large amounts of the oil are reported to be psychoactive with the hallucinogenic effects lasting for several days. One of the components of the oil is myristicin, the hallucinogenic principle in nutmeg. Sassafras has traditionally been used as an ingredient of beverages. To put the potential toxicity of sassafras into perspective, the following estimation has been made. Extrapolation of results from animal toxicity studies indicate that 0.66 mg/kg may prove hazardous in humans. By comparison, a cup of sassafras tea, prepared from a 2.5 g teabag, may provide up to 200 mg safrole, representing approximately 3 mg/kg.

Safrole, the principal component of the volatile oil, was first recognised to be a hepatocarcinogen in the 1960s and many animal studies have been documented concerning this toxicity. Both benign and malignant tumours have developed in laboratory animals, depending on the dose of safrole administered.

Both human and animal studies have shown that safrole gives rise to a large number of metabolites. A sulfate ester (formed via a hydroxylated metabolite) has been established as the ultimate carcinogen for safrole with tumour incidence paralleling the rate of conversion to the ester. Induction of cytochrome P450 activity has been associated with mutagenic and carcinogenic activity of the inducing agent. The inducing effect of safrole on certain metabolising enzymes is thought to play a role in the carcinogenic activity of safrole. The liver has a high level of cytochrome P450 activity and is therefore susceptible to induction.

Acute oral LD₅₀ values for safrole have been reported as 1.95 g/kg (rats) and 2.35 g/kg (mice). Major symptoms of toxicity are stated as ataxia, depression, diarrhoea, followed by death within 4 hours to seven days. Rats fed safrole in their diet at concentrations of 0.25, 0.5 and 1.0% exhibited reduction in growth, stomach and testicular atrophy, liver necrosis, biliary proliferation and primary hepatomas. Animals have also developed tumours when fed safrole–free extracts.

Conflicting results have been reported from studies investigating the mutagenicity of safrole, using the Ames test and DNA repair test. Purity
of the safrole, test system employed, type of metabolic activation mix, and toxicity of the test system have been suggested as reasons for the observed variations.\(^{(12)}\)
Contra–indications, Warnings

Sassafras should not be used internally or externally. Safrole, the major component in the volatile oil of sassafras, is hepatotoxic and even safrole–free extracts have been reported to produce tumours in animals. Sassafras essential oil is contra–indicated in internal and external use.\(^{(g58)}\) Sassafras has been reported to inhibit and induce microsomal enzymes.

**Pregnancy and lactation**
Sassafras is contra–indicated during pregnancy and lactation. The oil is reported to be abortifacient.\(^{(5)}\)
Pharmaceutical Comment

In addition to its traditional herbal use for treating dermatological and rheumatic ailments, sassafras also used to be a common flavouring ingredient in beverages, in particular root beer. However, animal studies have revealed the carcinogenic and hepatotoxic potential of safrole, the major component of sassafras volatile oil. Consequently, the use of safrole is no longer permitted in foods and sassafras is not permitted as a constituent of licensed medicinal products.

Antiseptic and diuretic properties claimed for sassafras are probably attributable to the volatile oil, although no documented studies were found supporting the antirheumatic claims. Sassafras should not be used as a herbal remedy, either internally or externally.
References

See also General References G2 G7 G11 G18 G21 G22 G31 G32 G36 G41 G43 G48 G58 G64.


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Saw Palmetto
Species (Family)

_Serenoa serrulata_ Hook., F. (Arecales/Palmae)
Synonym(s)

Part(s) Used

Fruit
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)

BPC 1934\(^{(G10)}\)

Complete German Commission E\(^{(G3)}\)

Martindale 33rd edition\(^{(G67)}\)

Mills and Bone\(^{(G50)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)

Ph Eur 2004\(^{(G72)}\)

USP26/NF21\(^{(G73)}\)
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents

See General References G22 G64.

**Carbohydrates**
Invert sugar 28.2%, mannitol, high molecular weight polysaccharides (e.g. MW 100 000) with galactose, arabinose and uronic acid\(^1\) identified as main sugar components for one.

**Fixed oils**
26.7%. Many free fatty acids and their glycerides. Monoacylglycerides (1–monolaurin, 1–monomyristicin).\(^2\) Oleic acid (unsaturated) and capric acid, caproic acid, caprylic acid, lauric acid, myristic acid, palmitic acid and stearic acid (saturated).

**Steroids**
β-Sitosterol, campesterol, stigmasterol and other compounds.\(^3–5\)

**Other constituents**
Flavonoids (e.g. rutin, isoquercitrin, kaempferol),\(^5\) pigment (carotene), resin, tannin and volatile oil 1.5%.

Most commercial preparations of saw palmetto contain lipophilic extracts.\(^G56\)
Food Use

Saw palmetto is not used in foods. In the USA, saw palmetto is listed by the Food and Drugs Administration (FDA) as a Herb of Undefined Safety.\(^{(G41)}\)
Herbal Use

Saw palmetto is stated to possess diuretic, urinary antiseptic, endocrinological and anabolic properties. Traditionally, it has been used for chronic or subacute cystitis, catarrh of the genitourinary tract, testicular atrophy, sex hormone disorders and specifically for prostatic enlargement.\textsuperscript{(G7 G32 G64)} Modern interest in saw palmetto is focused on its use in the treatment of symptoms of benign prostatic hyperplasia (BPH).
Dosage

*Dried fruit*
0.5–1.0 g or by decoction three times daily.\(^{(G7)}\)

*Extract*
320 mg lipophilic ingredients extracted with lipophilic solvents (hexane or ethanol 90% \(v/v\)).\(^{(G3)}\)

Clinical trials have assessed the effects of lipophilic extracts (containing lipids and sterols) of saw palmetto usually at a dosage of 160 mg twice daily.
Pharmacological Actions

Several pharmacological activities have been documented for saw palmetto *in vitro* and *in vivo* (animals). Several of these properties, such as inhibition of 5-α-reductase activity, inhibition of androgen binding and spasmolytic activity, are thought to explain, at least in part, the effects of saw palmetto in BPH. However, the clinical significance of the *in vitro* inhibition of 5-α-reductase activity by saw palmetto has not been clearly established (see Clinical studies). 5-α-Reductase is the enzyme that catalyses the conversion of testosterone to 5-α-dihydrotestosterone (DHT) in androgen target tissues, including the prostate. DHT is more potent than testosterone, and is thought to be implicated in the development of BPH. There is evidence that 5-α-reductase activity is higher in cells obtained from BPH tissue than from normal prostate tissue.

**In vitro and animal studies**

A lipidic (liposterolic) extract of saw palmetto was found to inhibit 5-α-reductase–mediated conversion of testosterone to dihydrotestosterone, and 3–ketosteroid reductase–mediated conversion of dihydrotestosterone to an androgen derivative. Other *in vitro* studies have shown that an ethanolic extract of saw palmetto (IDS-89) inhibited 5-α-reductase activity in the epithelium and stroma of human BPH tissue in a concentration–dependent manner. The IC$_{50}$ was around 2.2 mg/mL. This study also demonstrated that the inhibitory effect of IDS-89 was mainly due to the fatty acid constituents of a saponifiable subfraction of the extract, as non–saponifiable and hydrophilic subfractions showed little or no inhibition of 5-α-reductase activity. Inhibition of 5-α-reductase by a liposterolic extract of saw palmetto has also been documented in porcine prostatic microsomes.

There are at least two isoenzymes of 5-α-reductase (5-α-reductase types I and II), and several studies have documented that a liposterolic extract (Permixon) of saw palmetto inhibits both isoenzymes in prostate epithelial cells and fibroblast cells. Several other studies have documented inhibition of 5-α-reductase activity by liposterolic extracts of saw palmetto *in vitro*; these studies have been summarised elsewhere. Permixon was reported to inhibit 5-α-reductase activity without affecting the secretion of prostate–specific antigen (PSA) by epithelial cells, suggesting that use of saw palmetto extract should not interfere with PSA measurements for prostate–cancer screening.

Anti–androgenic activity has been documented for a hexane liposterolic extract (Permixon) of saw palmetto. *In vitro* studies in rat prostate tissue and
human foreskin fibroblasts indicated that this extract competitively inhibited the binding of dihydrotestosterone to cytosolic and nuclear androgen receptor sites.\(^{(6,11)}\) By contrast, an alcoholic extract of saw palmetto appeared to be without androgen receptor–binding activity.\(^{(12)}\)

Liposterolic extracts of saw palmetto have also been investigated in animal models of BPH. A liposterolic extract (Permixon) of saw palmetto 50 mg/kg body weight administered for 30 days to castrated rats with estradiol/testosterone–induced prostate enlargement resulted in significant reductions in the wet weight of the dorsal region of the prostate, compared with control.\(^{(13)}\) Another study in rats compared the effects of a liposterolic extract of saw palmetto with those of the 5-α-reductase inhibitor finasteride in rat prostate hyperplasia induced by hyperprolactinaemia.\(^{(14)}\) It was reported that the liposterolic extract of saw palmetto inhibited rat prostate hyperplasia in the lateral lobe induced by hyperprolactinaemia, and that finasteride did not antagonise the action of prolactin. By contrast, a study in dogs with BPH reported a lack of effect for saw palmetto extract on prostatic weight, prostatic volume, prostatic histologic scores, prostatic ultrasonographs and serum testosterone concentrations.\(^{(15)}\) In the study, 20 dogs with BPH, determined by raised prostatic volume and prostatic volume per kilogram body weight, received saw palmetto extract (type of extract not specified) 1500 mg daily in meatballs \((n = 8)\), 300 mg daily in meatballs \((n = 6)\) or unmedicated meatballs \((n = 6)\), for 91 days. Dogs included in this study did not have clinical signs of BPH (i.e. decreased urinary flow and residual urine volume) that often occur in human males with BPH. Dogs did not appear to be randomly assigned to treatment, and the mean prostatic volume in the control group was higher than that in the active treatment groups before treatment, although it was stated that this was not statistically significant. Assessments and data analysis were carried out by blinded investigators.

In a study using human prostate tissue, a liposterolic extract of saw palmetto (Permixon) 30 μg/mL significantly inhibited basic fibroblast growth factor–induced proliferation of human prostate cell cultures, compared with control, although the extract did not affect basal prostate cell proliferation.\(^{(16)}\) An unsaponified fraction of the extract also markedly inhibited basic fibroblast growth factor–induced cell proliferation, but had only a minimal effect on basal cell proliferation. In a study using stromal and epithelial tissue from normal prostate and from patients with BPH, cell numbers and proliferative indices were found to be higher in BPH tissue than in tissue from normal prostates.\(^{(17)}\) In tissue from patients with BPH who had been treated with a liposterolic extract of saw palmetto (Permixon), there was significant induction of apoptosis and inhibition of cell proliferation, compared with tissue from patients with BPH who had not received saw palmetto extract. In
Another *in vitro* study, incubation with Permixon 10 μg/mL also increased the apoptotic index for prostate epithelial cells by 35%.\(^{10}\)

Other *in vitro* studies have explored the effects of saw palmetto extract and its constituents on human prostatic cancer cells and other tumour cell lines. An extract of saw palmetto fruit, prepared by supercritical fluid extraction with carbon dioxide, induced cell death in LNCaP cells (a hormonal therapy–resistant prostatic cancer cell line) in a concentration–dependent manner.\(^{18}\) This confirms the findings of previous studies demonstrating the effect of a liposterolic extract of saw palmetto (Permixon) on the mortality rate of LNCaP cells: increased mortality was observed with saw palmetto extract 50 μg/mL, compared with control.\(^{19}\) Further investigation identified myristoleic acid as a component of saw palmetto extract that caused cell death. The EC\(_{50}\) for both the extract and myristoleic acid was around 100 μg/mL.\(^{18}\) Following incubation of LNCaP cells with saw palmetto extract 130 μg/mL or myristoleic acid 100 μg/mL, the proportions of apoptotic and necrotic cells were 16.5% and 46.8%, respectively, for the extract, and 8.8% and 81.8%, respectively, for myristoleic acid. An extract of saw palmetto obtained by supercritical extraction with carbon dioxide inhibited the invasion of PC-3 cells (derived from human adenocarcinoma of the prostate) into Matrigel *in vitro* in a concentration–dependent manner at concentrations in the range 1–10 μg/mL.\(^{20}\) However, LNCaP cells and SKRC-1 cells (derived from human renal carcinoma) were unaffected by the extract. The extract was also shown to inhibit the activity of urokinase–type plasminogen activator, a protease enzyme that is necessary for tumour–cell invasion into basement membranes. The monoacylglycerides 1–monolaurin and 1–monomyristicin isolated from saw palmetto demonstrated *in vitro* activity against renal (A-498) and pancreatic (PACA-2) human tumour cells (EC\(_{50}\) for 1–monolaurin: 3.77 μg/mL and 2.33 μg/mL, respectively; EC\(_{50}\) for 1–monomyristicin: 3.58 μg/mL and 1.87 μg/mL, respectively).\(^{2}\) However, only borderline cytotoxicity was observed against PC-3 cells (EC\(_{50}\) for 1–monomyristicin: 8.84 μg/mL).\(^{2}\)

Spasmolytic activity has also been documented for saw palmetto and it has been suggested that this may contribute to the herb’s effects in BPH. An ethanolic lipidic extract was reported to produce a concentration–dependent relaxation on rat uterus tonic contraction induced by vanadate (EC\(_{50}\) 11.41 μg/mL).\(^{21}\) Further investigation suggested that a mechanism for the observed effect could be interference with intracellular calcium mobilisation, possibly mediated via cyclic AMP. Other *in vitro* studies demonstrated that a lipophilic ethanolic extract of saw palmetto 0.3–0.75 mg/mL reduced norepinephrine (noradrenaline)-induced contractions in rat deferential duct.
Further study indicated that the relaxant effect of saw palmetto extract results from either α-adrenoceptor blockade or from calcium-blocking activity.\(^{(22)}\)

**Other activity**

*In vivo* oestrogenic activity in the rat has also been documented for an alcoholic extract.\(^{(23)}\) Activity was attributed to the high content of β-sitosterol, a known oestrogenic agent, present in saw palmetto.

*In vivo* anti-oedema activity in the rat has been documented for a hexane extract of saw palmetto, acting by inhibition of histamine–induced increase in capillary permeability.\(^{(24)}\) Low doses of an aqueous extract were effective in carrageenan–induced paw oedema and pellet tests in the rat, although the extract was not found to influence the proliferative stage of inflammation.\(^{(1,25)}\) The observed anti-inflammatory activity was attributed to a high molecular weight polysaccharide (approximately 100 000). Polysaccharides possessing immunostimulating activity have also been documented for saw palmetto and were stated to contain a high content of glucuronic acid.\(^{(1,25)}\)

An extract (SG-291) prepared from saw palmetto fruits by supercritical fluid extraction with carbon dioxide was reported to inhibit both cyclooxygenase and 5–lipoxygenase *in vitro* (IC\(_{50}\) 28.1 μg/mL and 18.0 μg/mL, respectively).\(^{(26)}\) Further study indicated that the component(s) of saw palmetto extract that inhibits these enzymes must be within the acidic lipophilic fraction. Subsequent studies have documented that a liposterolic extract of saw palmetto (Permixon) significantly inhibited the production of 5–lipoxygenase metabolites, including leukotriene B\(_4\), by human polymorphonuclear neutrophils at concentrations of saw palmetto extract of 5 μg/mL and above.\(^{(27)}\)

**Clinical studies**

**Pharmacokinetics**

Some data on the pharmacokinetics of saw palmetto extracts in healthy male volunteers (\(n = 12\)) come from an open, randomised, single-dose bioequivalence study of a 320–mg capsule of a liposterolic extract of saw palmetto compared with two capsules of saw palmetto extract 160 mg as the reference preparation.\(^{(28)}\) The plasma concentration–time curves were reported to be almost identical for both preparations. The maximum concentration (\(C_{\text{max}}\)) for saw palmetto extract 320–mg capsule and 2 × 160–mg capsules was 2.54–2.61 μg/mL and 2.57–2.67 μg/mL, respectively, and
time to $C_{\text{max}} \left( T_{\text{max}} \right)$ was 1.58 and 1.5 hours for the 320–mg capsule and 2 × 160–mg capsules, respectively. Another study explored the bioavailability and pharmacokinetic profile of a rectal formulation of saw palmetto extract 640 mg in healthy male volunteers ($n = 12$). The mean maximum plasma concentration of the second component of saw palmetto was almost 2.6 μg/mL at around 3 hours after drug administration.

**Pharmacodynamics**

The inhibitory effects of saw palmetto extract on 5-α-reductase activity documented *in vitro* (see Pharmacological Actions, *In vitro* and animal studies) have been confirmed in some studies in humans, and refuted by others.

In one study, 25 men with symptomatic, established BPH were randomised to receive either a liposterolic extract of saw palmetto (Permixon) 320 mg/day for three months ($n = 10$), or no treatment ($n = 15$). At the end of the treatment period, analysis of samples of BPH tissue, obtained by suprapubic prostatectomy, showed that dihydrotestosterone concentrations were significantly reduced and that testosterone concentrations were significantly higher in the treatment group, compared with the control group ($p < 0.001$ for both). A significant reduction in concentrations of epidermal growth factor in total BPH tissue was also observed in the treatment group, compared with the control group ($p < 0.01$). The reported biochemical effects were most evident in BPH tissue from the periurethral region.

In another study, biopsy specimens of the prostate were taken from 44 men with symptomatic BPH participating in a randomised, placebo–controlled trial of a herbal combination preparation containing saw palmetto lipoidal extract 106 mg together with nettle root extract, pumpkin seed oil, lemon bioflavonoid extract and vitamin A. There were no statistically significant differences in median tissue dihydrotestosterone and testosterone concentrations between the treatment and placebo groups at baseline. At the end of the study, mean tissue dihydrotestosterone concentrations decreased significantly in the treatment group, compared with baseline values ($p = 0.005$), whereas there was no significant change in dihydrotestosterone concentrations in the placebo group. However, in a separate analysis, it was reported that the median change in tissue dihydrotestosterone concentrations for the treatment group (1.38 ng/g) did not differ significantly from the corresponding change in the placebo group (0.87 ng/g). The findings of this study should be interpreted cautiously as it is possible there are other explanations for the observed effect.

Another randomised, double–blind trial involving 18 men with BPH compared
saw palmetto extract (IDS-89; Strogen) 640 mg three times daily (i.e. six times the normal dose) for three months with placebo.\(^{33}\) This high dose of saw palmetto extract achieved only a moderate decrease in 5-α-reductase activity.

An open, randomised, placebo–controlled study involving 32 healthy male volunteers compared the effects of a liposterolic extract of saw palmetto (Permixon) 80 mg twice daily for seven days with those of finasteride 5 mg daily for seven days on inhibition of 5-α-reductase activity.\(^{34}\) Serum dihydrotestosterone concentrations were reported to decrease significantly with finasteride, compared with baseline values, but no significant changes were observed for the saw palmetto and placebo groups. Thus, this study did not support a mechanism of action for saw palmetto in BPH by inhibition of 5-α-reductase activity.

Alpha–adrenoceptor blocking activity has also been documented for saw palmetto extract \textit{in vitro},\(^{22}\) although this has been refuted in a study involving healthy volunteers.\(^{35}\) In a double–blind, placebo–controlled, four–way, crossover study, 12 healthy male volunteers received three different saw palmetto extract preparations (Prostagutt uno, Prostess uno, Talso uno) 320 mg daily for eight days each, separated by wash–out phases of at least two weeks. It was reported that none of the study medications showed signs of α\textsubscript{1}-adrenoceptor subtype occupancy as determined by a radioreceptor assay.

\textit{Therapeutic effects}

Numerous clinical studies have investigated the effects of saw palmetto in men with BPH.

A systematic review and meta–analysis included 18 randomised clinical trials (16 of which were double–blind) of saw palmetto extracts involving a total of 2939 men with BPH.\(^{36}\) This work has also been published as a Cochrane systematic review.\(^{37}\) The review included 10 studies which compared saw palmetto extracts alone with placebo, three comparing saw palmetto extracts in combination with other herbals with placebo, two comparing saw palmetto extracts alone with an active control, one comparing saw palmetto extracts in combination with other herbals with an active control, one comparing saw palmetto extract with another herb and with placebo, and one comparing oral saw palmetto extract with a rectal formulation of saw palmetto extract. The mean duration of the included studies was nine weeks (range 4–48 weeks).

Compared with placebo, saw palmetto extracts led to a decrease in urinary symptom scores and nocturia, and improvements in self–rating of urinary
symptoms and peak urine flow.\(^{(37)}\) Compared with finasteride, saw palmetto extracts achieved similar improvements in urinary symptom scores and peak urine flow. This systematic review was considered to have provided good evidence that saw palmetto is effective in men with symptoms of BPH, although there is scope for further trials.\(^{(37,38)}\)

Several other clinical studies of saw palmetto extracts in BPH have now been published since the Cochrane systematic review, although few have comprised rigorous study design capable of testing efficacy, and several have investigated combination preparations of saw palmetto with other herbs.

A short report describes a randomised, double-blind, placebo-controlled trial of saw palmetto extract (LG-166S) 160 mg twice daily for six months in 101 men with BPH.\(^{(39)}\) This study reported statistically significant differences in symptom scores between the treatment group and the placebo group at the end of the study \((p < 0.001)\).

In a study involving 75 men with mild/moderate BPH according to their International Prostate Symptom Score (IPSS), participants received a liposterolic extract of saw palmetto (Permixon) 160 mg twice daily for nine weeks \((n = 57)\).\(^{(40)}\) A control group \((n = 18)\) did not receive any medical treatment for BPH, and there was no random allocation to treatment, although it was stated that baseline parameters were comparable between the two groups. It was reported that, at the end of the study, IPSS and quality-of-life scores, compared with baseline values, significantly improved in Permixon-treated men \((p < 0.001)\). There were no significant differences in these parameters, compared with baseline values, for the control group.

Two randomised studies involving men with symptomatic BPH have compared the effects of different regimens of saw palmetto extract.\(^{(41,42)}\) A multicentre, randomised, single-blind trial involving 132 men with BPH compared the effects of saw palmetto extract (Prostaserene) 320 mg once daily with 160 mg twice daily for one year.\(^{(41)}\) Another study compared a liposterolic extract of saw palmetto (Permixon) 320 mg daily with 160 mg twice daily for three months in 100 men with symptomatic BPH.\(^{(42)}\) For each regimen, both studies reported significant improvements in the mean IPSS, maximum and mean urinary flow rates and residual urine volume, at the end of the studies, compared with baseline values. However, as these studies did not include a placebo-control group, the possibility that the observed effects are placebo effects cannot be excluded.

Several other open, uncontrolled studies of saw palmetto extracts (alone or in combination with other herbs), several of which were drug-monitoring studies which also assessed effectiveness, have reported improvements in
symptoms of BPH at the end of treatment, compared with baseline values.\(^{43-47}\) Doses assessed in these studies were usually 160 mg two or three times daily for up to three years. These studies are discussed in more detail later (see Side-effects, Toxicity).

The effects of a combination herbal preparation containing saw palmetto lipoidal extract 106 mg together with nettle root extract, pumpkin seed oil, lemon bioflavonoid extract and vitamin A, were assessed in a six-month, randomised, double-blind, placebo-controlled trial involving 44 men with symptomatic BPH.\(^{32}\) At the end of the treatment period, a slight decrease in symptom score and an increase in urinary flow were observed for both groups, compared with baseline values. These changes were greater in the treatment group, compared with the placebo group, but this difference was not statistically significant. In another randomised, double-blind, controlled trial involving 543 men with BPH, participants received a combination of saw palmetto extract 160 mg and nettle root extract 120 mg (Prostagutt forte) daily, or finasteride 5 mg daily, for 48 weeks.\(^{48}\) Data from a subgroup of 431 participants with ultrasonographic measurements were analysed. Mean maximum urinary flow and IPSS improved in both groups, compared with baseline values; there were no statistically significant differences between the two groups.

Saw palmetto is one of the eight herbal ingredients contained in a commercial preparation known as PC-SPES; the other herbal ingredients are chrysanthemum, isatis, licorice, \textit{Ganoderma lucidum}, \textit{Panax pseudoginseng}, \textit{Rabdosia rubescens} and \textit{Scutellaria} (scullcap). The combination preparation has been investigated for oestrogenic activity, and is currently of interest for its potential effects in the treatment of hormone-sensitive prostate cancer.\(^{49}\)
Side-effects, Toxicity

A systematic review and meta-analysis of 18 randomised clinical trials of saw palmetto extracts (see Pharmacological Actions, Clinical studies, Therapeutic effects) reported that adverse effects with saw palmetto were generally mild and comparable to those with placebo.\(^{(37)}\) Gastrointestinal effects were reported in 1.3% of men taking saw palmetto extracts, placebo (0.9%) and finasteride (1.5%). Study withdrawal rates for men taking saw palmetto, placebo and finasteride were 9.1%, 7.0% and 11.2%, respectively. The authors of the review concluded that saw palmetto extracts are associated with fewer adverse treatment effects than is finasteride, but that little is known about the long-term safety of saw palmetto extracts.

In a drug-monitoring study involving 1334 men with BPH, the tolerability of saw palmetto extract 160 mg twice daily for 12 weeks was reported to be ‘good’ or ‘excellent’ by more than 95% of participants.\(^{(46)}\) This is similar to a finding from a three-year prospective, uncontrolled study involving 435 men with BPH, in which the tolerability of saw palmetto extract (IDS-89) 160 mg twice daily was classified as ‘good’ or ‘very good’ by both physicians and patients for 98% of participants.\(^{(45)}\) A total of 46 adverse events was reported in 34 patients. Of these, 30% were gastrointestinal disturbances. The withdrawal rate from the study was 1.8%, mostly because of digestive disturbances ($n = 3$) and tumours ($n = 3$). Non-serious adverse effects (4.95–6.63%), mainly minor gastrointestinal effects, such as gastralgia, nausea, diarrhoea, constipation and anorexia, as well as vertigo, headache, dry mouth and pruritus, were reported in an open study involving 413 men with BPH who received saw palmetto extract 160 mg twice daily for three months.\(^{(44)}\) An observational study involving 2080 patients with BPH who received a combination of saw palmetto extract (WS-1473) and nettle root extract (WS-1031) reported that the tolerability of the preparation was classified by physicians to be ‘good’ or ‘very good’ for the majority of participants.\(^{(47)}\) Mild adverse effects were reported in 15 patients (0.72%).

Studies assessing the equivalence of two different regimens of saw palmetto extract (320 mg once daily and 160 mg twice daily) report that adverse events occurred with a similar frequency in both groups.\(^{(41,42)}\) Most events were deemed to be unrelated or unlikely to be related to treatment with saw palmetto extract.

**Toxicity** Incubation of high concentrations of saw palmetto extract (Permixon) 9.0 mg/mL for 48 hours inhibited sperm motility, compared with control.\(^{(50)}\)
Contra-indications, Warnings

In view of the reported anti-androgen and oestrogenic activities, saw palmetto may affect existing hormonal therapy, including the oral contraceptive pill and hormone replacement therapy.

Pregnancy and lactation
The safety of saw palmetto has not been established. In view of the lack of toxicity data and the documented hormonal activity, the use of saw palmetto during pregnancy and lactation should be avoided.
Several pharmacological activities have been documented for saw palmetto in vitro and in vivo (animals). Some of these properties, such as inhibition of 5-α-reductase activity, inhibition of androgen binding and spasmolytic activity, are thought to explain, at least in part, the effects of saw palmetto in benign prostatic hyperplasia (BPH). However, some experimental and clinical studies report conflicting results, particularly with regard to the inhibition of 5-α-reductase activity and α-adrenoceptor blocking activity by saw palmetto extracts. Thus, the mechanism(s) of action of saw palmetto extracts in BPH remain unclear. This is not surprising, given that, at present, the exact cause of BPH is unknown. In addition to the effects of saw palmetto in experimental models of BPH, immunostimulant and anti-inflammatory activities have been documented in laboratory studies.

Results of clinical studies indicate that saw palmetto is a potential agent for the treatment of BPH. However, this is not an indication suitable for self-diagnosis and self-treatment, and over-the-counter use of saw palmetto extract for BPH should be under medical supervision. Data from randomised clinical trials and drug-monitoring studies indicate that, generally, saw palmetto is well-tolerated; adverse events are mild and relate mainly to gastrointestinal symptoms. However, in view of the lack of toxicity data and the documented pharmacological actions of saw palmetto, excessive use should be avoided.
References

See also General References G3 G5 G9 G10 G22 G31 G32 G36 G41 G50 G56 G64.


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Species (Family)

*Scutellaria lateriflora* L., *S. baicalensis* Georgi and other *Scutellaria* species (Labiatae)

*S. baicalensis* Georgi is a species commonly referred to as scullcap in Chinese herbal medicine.
Synonym(s)
Helmet Flower, Hoodwort, Quaker Bonnet, Scutellaria, *Scutellaria galericulata* L., Skullcap
Part(s) Used
Herb
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL\textsuperscript{(G37)}
Constituents

See General References G20 G22 G48 G60 G64.

Limited information has been documented regarding the constituents of *S. lateriflora*, although various related *Scutellaria* species have been investigated.

**Flavonoids**
Apigenin, hispidulin, luteolin, scutellarein, scutellarin (bitter glycoside).

**Iridoids**
Catalpol.

**Volatile oils**
Limonene, terpineol (monoterpenes); *d*-cadinene, caryophyllene, *trans*-β-farnesene, β-humulene (sesquiterpenes).

**Other constituents**
Lignin, resin and tannin.

**Other Scutellaria species**
The related species *S. baicalensis* is reported to contain baicalein, baicalin, chrysin, oroxylin A, skullcapflavone II and wogonin.\(^{1-3}\)

*S. galericulata* is stated to contain apigenin, baicalein, baicalin, apigenin-7-glucoside and galeroside (baicalein-β-L-rhamnofuranoside).\(^{4}\)
Food Use

Scullcap is not used in foods. In the USA, scullcap is listed by the Food and Drugs Administration (FDA) as a Herb of Undefined Safety.\(^{(G22)}\)
Herbal Use

Scullcap is stated to possess anticonvulsant and sedative properties.\(^{(G34 \ G64)}\)
Traditionally, it has been used for epilepsy, chorea, hysteria, nervous tension states, and specifically for grand mal epilepsy.\(^{(G7)}\) In Chinese herbal medicine, the roots of *S. baicalensis* Georgi have been used traditionally as a remedy for inflammation, suppurative dermatitis, allergic diseases, hyperlipidaemia and atherosclerosis.
Dosage

**Dried herb**
1–2 g or by infusion three times daily.\(^{(G7)}\)

**Liquid extract**
2–4 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)

**Tincture**
1–2 mL (1:5 in 45% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

In vitro and animal studies

None documented for *Scutellaria lateriflora*.

Many investigations have been undertaken to study the pharmacological actions of *S. baicalensis* root. Documented actions have primarily been attributed to the various flavonoid constituents and include: *in vitro* inhibition of mast cell histamine release comparable to disodium cromoglicate for some flavonoids; *(1)* *in vitro* cytotoxicity of scullcap flavone II; *(5)* *in vivo* and *in vitro* inhibition of lipid peroxidation; *(6–8)* *in vitro* inhibition of lipoxygenase and cyclooxygenase pathways; *(9)* hypocholesterolaemic activity in rats. *(10)* This *in vivo* effect has been linked to *in vitro* actions documented for various flavonoids, including prevention of ethanol–induced hyper lipidaemia, *(11)* catecholamine–induced lipolysis *(10,11)* and lipogenesis in adipose tissue; *(10,11)* there is no pronounced effect on blood pressure in cats and rabbits. *(12)* In addition, the latter study found no CNS-depressant and no antispasmodic activity. However, it did find marked antibacterial activity against various Gram–positive bacteria (e.g. *Bacillus subtilis*, *Escherichia coli*, *Sarcina lutea* and *Staphylococcus aureus*). *(13)*

Clinical studies

Clinical investigation of scutellarin involving 634 cases of cerebral thrombosis, cerebral embolism, and paralysis caused by stroke has been undertaken. An overall effective rate of more than 88% was reported following intramuscular, intravenous or oral administration. *(14)*
Side-effects, Toxicity

Symptoms caused by overdosage of scullcap tincture include giddiness, stupor, confusion and seizures. Hepatotoxic reactions have been reported after ingestion of scullcap-containing preparations. Adulteration of scullcap herb by *Teucrium* is recognised. Several cases of hepatitis have been associated with germander (*Teucrium chamaedrys*).
Contra-indications, Warnings

None documented. In view of the possible hepatotoxicity associated with scullcap, its use is best avoided.

Pregnancy and lactation
Scullcap is stated to have been used traditionally to eliminate a mother’s afterbirth and to promote menstruation.\(^{(G22)}\) Limited information is known regarding the pharmacological activity and toxicity of scullcap. In view of this and concerns over hepatotoxicity, scullcap should not be taken during pregnancy and lactation.
Pharmaceutical Comment

Limited information has been documented regarding the chemistry of scullcap. Most of the pharmacological activities reported for other *Scutellaria* species have been attributed to the flavonoid constituents. Despite the traditional uses of scullcap as a sedative and anticonvulsant, there are no documented scientific data to support these uses. Commercial scullcap is commonly recognised to be adulterated with *Teucrium* species, notably *Teucrium canadense*. Herbal preparations stated to contain scullcap may therefore contain a *Teucrium* species. Few pharmacological studies have been undertaken for *Teucrium* species. Hepatitis has been associated with germander (*Teucrium chamaedrys*). Hepatotoxicity has resulted in humans taking commercially available remedies in the UK which are stated to contain scullcap. It would seem advisable to avoid ingestion of scullcap.
References

See also General References G5 G9 G10 G18 G20 G22 G31 G32 G34 G36 G37 G48 G60 G64.


Species (Family)

Polygala senega L. (Polygalaceae) and other closely related species cultivated in western Canada and Japan.
Synonym(s)
Northern Senega (Canada), Polygala, *Polygala senega* var. *latifolia* (Japan), Rattlesnake Root, Snake Root
Part(s) Used

Root, rootstock
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)
BHP 1996\(^{(G9)}\)
BP 2002\(^{(G71)}\)
Complete German Commission E\(^{(G3)}\)
ESCOP 1997\(^{(G52)}\)
Martindale 33rd edition\(^{(G67)}\)
PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Ph Eur 2004\(^{(G72)}\)
Legal Category (Licensed Products)

GSL(G37)
Constituents

See Reference 1, and General References G2 G6 G20 G40 G48 G52 G59 G62 G64.

Acids
Salicylic acid and its methyl ester 0.1–0.2%; hydroxycinnamic acids (e.g. caffeic acid, ferulic acid, sinapic acid) free or esterified with saponins.\(^{(2)}\)

Carbohydrates
Arabinose, fructose, glucose, melibiose, raffinose, saccharose, stachyose, sucrose; 1,5-anhydro-D-glucitol and other D-glucitol derivatives;\(^{(3,4)}\) trisaccharides; mucilage, pectin. A series of oligosaccharide esters, senegoses A–O, containing acetic, benzoic, trans- and cis-ferulic acid moieties linked to glucose and fructose.\(^{(5,6)}\) Five acylated sucrose glycosides, tenuifolisides A–E, have been isolated from *P. tenuifolia*.\(^{(7,8)}\) The esterifying acids are 3,4,5-trimethoxycinnamic, *p*-hydroxybenzoic, sinapic and ferulic.

Terpenoids
A complex mixture of bidesmosidic triterpene saponins (6–10%) based on the aglycone presenegin. The total saponin mixture may be referred to as senegin. The saponins of *P. senega* var. *latifolia* are 3–glucosides of presenegin with tetra-, penta- or hexa–glucosyl groups linked at C-28 and including 4′′-methoxy–cinnamoyl or 3′′,4′′-dimethoxy cinnamoyl fucosyl resulting in *E*– and *Z*-cinnamoyl isomers of each saponin.\(^{(9–11)}\) Senegins I–IV were the first saponins to be characterised and were *E*-isomers.\(^{(12,13)}\) *P. tenuifolia* contains similar saponins named onjisaponins A–G.\(^{(14,15)}\)

Xanthones
A number of xanthones have been isolated from *P. tenuifolia* including 4-C-[β-D-apiofuranosyl-(1→6)-β-D-glucopyranosyl]-1,3,6-trihydroxy-7-methoxyxanthone.\(^{(8)}\)

Other constituents
Fat, resin, sterols and valeric acid ester.

Other *Polygala* species
*Polygala paniculata* contains coumarins (aurapten, murrangatin, phebalosin and 7–methoxy-8-(1,4–dihydroxy–3–methyl–2–butenyl) coumarin,\(^{(16)}\) pyranocoumarin).\(^{(17)}\) *Polygala chamaebuxus* (European species) contains hydroxycinnamic acid esters involving acetic, ferulic and sinapic acids as the ester moieties, saponins, tenuifolin (prosapogenin), rutin (flavonoid...
glycoside), coniferin and syringen (phenolic glycosides).\(^{(2)}\)

Other European species (e.g. *Polygala alpestris*, *Polygala comosa*, *Polygala vayredae*) contain complex mixtures of bidesmosidic saponins, tenuifolin (prosapogenin), hydroxycinnimic acid esters similar to those reported for *P. chamaebuxus*.\(^{(18)}\) *Polygala triphylla* contains B-ring oxygen–free trioxygenated- and glucosyloxy–xanthones.\(^{(19)}\) *Polygala polygama* contains podophyllotoxin and demethylpodophyllotoxin (lignans).\(^{(20)}\)
Food Use

Senega is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that senega can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product.\(^{(G16)}\)
Senega is stated to possess expectorant, diaphoretic, sialogogue and emetic properties. Traditionally, it has been used for bronchitic asthma, chronic bronchitis, as a gargle for pharyngitis, and specifically for chronic bronchitis.
**Dosage**

**Dried root**
0.5–1.0 g or by infusion three times daily.\(^{6,7}\)

**Senega Liquid Extract**
(BPC 1968) 0.3–1.0 mL.

**Senega Tincture**
(BPC 1968) 2.5–5.0 mL.
Pharmacological Actions

In vitro and animal studies

Mucosal secretion
Polygalic acid and senegin are stated to be irritant to the gastrointestinal mucosa, and to cause a reflex secretion of mucus in the bronchioles.\(^{(1,G6 G44 G52)}\) A fluid extract of senega increased respiratory tract fluid secretion in guinea-pig, cat and dog, but not in rabbit.\(^{(G52)}\)

CNS-depressant activity
CNS-depressant properties in mice (e.g. reduction in spontaneous activity, inhibition of amphetamine stimulation, potentiation of barbiturate-induced sleeping time, and decrease in rectal temperature) have been documented for *Polygala microphylla*.\(^{(21)}\) Similar properties have been reported for *Polygala tenuifolia* and have been attributed to the saponin constituents. A methanolic extract of *P. tenuifolia*, various fractions and pure onjisaponins B, F and G prolonged hexobarbital sleeping time in mice.\(^{(G52)}\) Onjisaponin F produced sleep times in mice of 33 and 35 minutes for doses of 5 and 20 mg/kg, respectively, compared with 24 minutes for control and 42 minutes for chlorpromazine hydrochloride (2 mg/kg).

Inhibition of alcohol absorption
*E,Z*-senegin II and *E,Z*-senagasaponins a and b from *P. senega* var. *latifolia* have potent inhibitory effects on alcohol absorption in rats. *E,Z*-senagasaponins a or b (100 mg/kg) administered orally to rats 1 hour after 20% aqueous ethanol (5 mL/kg, orally) reduced blood alcohol concentrations after 1 hour from 0.5 mg/mL to 0.02 mg/mL.\(^{(10)}\) Under similar test conditions, *E,Z*-senegen II administration led to a blood ethanol concentration of 0.09 mg/mL.

Hypoglycaemic activity
Senegin II and *E,Z*-senagasaponins a and b have significant hypoglycaemic effects in rodents.\(^{(22)}\) Senegin II (2.5 mg/kg, intraperitoneally) reduced blood glucose concentrations in normal mice from 220 mg/dL to 131 mg/dL 4 hours after administration and also significantly lowered blood glucose concentrations in KK-Ay mice from 434 mg/dL to 142 mg/dL under similar test conditions (\(p < 0.001\), compared with control, for both studies). In glucose tolerance tests in rats, administration of *E,Z*-senagasaponins a and b (100 mg/kg, orally) resulted in glucose concentrations of 107–123 mg/mL after 30 minutes compared with 156 mg/mL in control animals (\(p < 0.01\)).\(^{(11)}\)
Hypolipidaemic activity

Seven hours after administration of an n-butanol fraction of a methanolic extract of *P. senega* var. *latifolia* containing senegin II (5 mg/kg, intraperitoneally), the mean (standard deviation) blood triglyceride concentration was 65 (9) mg/100 mL, compared with 152 (17) mg/mL in control animals (*p* < 0.05). The blood triglyceride concentration in cholesterol–fed mice was also significantly reduced (*p* < 0.05) under the similar test conditions. Pure senegin II at a dose of 5 mg/kg was also reported to lower blood triglyceride concentrations in mice.\(^{(23)}\)

**Other activities**

Guinea–pig serum taken 2 hours after administration of lyophilised aqueous extract of *P. tenuifolia* (600 mg, intraperitoneally) inhibited the growth of herpes simplex virus type 1 (HSV-1) in Vero cells.\(^{(G52)}\) An unspecified senegin from *P. senega* produced a 34% inhibition of influenza virus (A2/Japan 305) at a concentration of 12.5 μg/mL.\(^{(G52)}\) An ethanolic extract of *P. senega* has been reported to inhibit growth of a range of fungi.\(^{(G52)}\)

*Polygala erioptera* and *P. paniculata* have exhibited molluscicidal activity, and *P. paniculata* is reported to possess antifungal activity.\(^{(17)}\) A butanol extract of *P. tanuifolia* containing onjisaponins (100 μg/mL) inhibited cyclic adenosine mono phosphate (cAMP) diesterase by 73%.\(^{(G52)}\) Isolated onjisaponins E, F and G inhibited cAMP phosphodiesterase, with IC\(_{50}\) values of 3.1, 2.9, and 3.7 × 10\(^{-5}\) mol/L, respectively, being similar in action to papaverine. A total saponin concentration of *P. senega* var. *latifolia* increased rat plasma concentrations of adrenocorticotropic hormone (ACTH), corticosterone and glucose 30 minutes after intraperitoneal administration (25 mg/kg). Single doses of a dried methanol (50%) extract of *P. senega* var. *latifolia* and *P. tanuifolia* administered orally (2 g/kg) to rats produced 62% and 100% inhibition, respectively, of congestive oedema.\(^{(G52)}\) Under the same conditions, furosemide 100 mg/kg produced 100% inhibition of congestive oedema.

**Clinical studies**

A fluid extract of senega root was reported to reduce the viscosity of sputum in patients with bronchiectasis.\(^{(G52)}\) A French patent has stated that a triterpenic acid extracted from senega possesses anti–inflammatory activity and is effective against graft rejection, eczema, psoriasis and multiple sclerosis.\(^{(24)}\)
Side-effects, Toxicity\textsuperscript{G20}

Saponins are generally regarded as irritant to the gastrointestinal mucosa, and irritant properties have been documented for senega plant and for related Polygala species.\textsuperscript{G51} Large doses of senega are reported to cause vomiting and purging.\textsuperscript{G60}

The haemolytic index (HI) of senega saponins is stated to be between 2500 and 4500.\textsuperscript{G62} Haemolytic saponins are toxic to mammals when administered intravenously, but have a low toxicity when given orally because they do not cross the gastrointestinal mucosa.\textsuperscript{25} Contact with damaged mucosal areas may cause a problem. Toxicity associated with chronic exposure of the gastrointestinal mucosa to haemolytic saponins has not been established. It has been stated that the suitability of saponins for nutritional and pharmacological use requires further investigation: free saponins in the gastrointestinal tract may interact with the mucosal cells, causing a transient increase in the permeability of the small intestine to intraluminal solutes and inhibiting active nutrient absorption.\textsuperscript{25} This action may consequently facilitate the entry of antigens and biologically active food peptides into the blood circulation, with adverse systemic effects.\textsuperscript{25} Aqueous and methanol extracts of *P. senega* and *P. tenuifolia* were negative in the rec–assay with *Bacillus subtilis* and in the reversion assay with Ames strains TA98 and TA100 of *Salmonella typhimurium*.\textsuperscript{G52} A mixture of senegins given to rats (i.p.) gave an LD\textsubscript{50} value of 3 mg/kg and inhibited the growth of Walker carcinoma in rats with an ED\textsubscript{50} value of 1.5 mg/kg.\textsuperscript{G52}

Cytotoxic lignans have been documented as constituents of a related species, *P. polygama*.\textsuperscript{10}
Contra-indications, Warnings

Senega may exacerbate existing gastrointestinal inflammation and excessive doses may cause vomiting. Senega has hypoglycaemic activity and is contraindicated in diabetic patients.

Pregnancy and lactation

Limited information is available on the chemistry, pharmacology and toxicity of senega. In view of this, and the potential irritant properties of senega, its use during pregnancy and lactation should be avoided.
Pharmaceutical Comment

The chemistry and pharmacology of senega has been extensively investigated but there is only limited clinical data. The activity of the saponins in animals supports the herbal use for bronchitis. In view of the lack of toxicity data and uncertainty regarding the risk associated with chronic ingestion of haemolytic saponins, excessive use of senega should be avoided.
References


14. Sakuma S, Shoji J. Studies on the constituents of the root of *Polygala*...


Species (Family)

i. *Cassia senna* L.

ii. *Cassia angustifolia* Vahl. (Leguminosae)
Synonym(s)

i. Alexandrian Senna, *Cassia acutifolia* Delile, Khartoum Senna

ii. Indian Senna, Tinnevelly Senna
Part(s) Used

Fruit (pod), leaf
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

BP 2002\(^{(G71)}\)

Complete German Commission E\(^{(G3)}\)

ESCOP 1997\(^{(G52)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)

Ph Eur 2004\(^{(G72)}\)

USP26/NF21\(^{(G73)}\)

WHO volume 1 1999\(^{(G63)}\)
Legal Category (Licensed Products)

GSL($G^{37}$)
Constituents

See General References G2 G6 G7 G8 G20 G22 G41 G48 G52 G62 G64.

**Hydroxyanthracenes**
Pharmacopoeial standards not less than 2.5% for leaf, 3.5% for *C. senna* fruit and 2.2% for *C. angustifolia* fruit.\(^{(G15 G28)}\) Dianthrone glycosides (1.5–3% leaf; 2–5% fruit), primarily sennosides A and B (rhein dianthrenes) with sennosides C and D (rhein aloe–emodin heterodianthrenes), aloe–emodin dianthrone. Sennosides A and B yield sennidin A and B respectively. Free anthraquinones including aloe–emodin, chrysophanol and rhein with their glycosides.

**Carbohydrates**
Polysaccharides (about 2.5%)\(^{(1)}\) including mucilage (arabinose, galactose, galacturonic acid, rhamnose) and a galactomannan (galactose, mannose);\(^{(2)}\) free sugars (e.g. fructose, glucose, pinitol, sucrose).

**Flavonoids**
Flavonols including isorhamnetin and kaempferol.

**Glycosides**
6-Hydroxymusizin and tinnevellin glycosides.

**Other constituents**
Chrysophanic acid, salicylic acid, saponin, resin, volatile oil (trace).
Food Use

Senna is listed by the Council of Europe as a natural source of food flavouring (Tinnevelly category N2, Alexandrian category N3). Category N2 indicates that senna can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product. Category N3 indicates that there is insufficient information available about Alexandrian senna, for an adequate assessment of potential toxicity.\(^{(G16)}\) In the USA, senna is permitted for food use.
Herbal Use

Senna is stated to possess cathartic properties (leaf greater than fruit) and has been used traditionally for constipation. The German Commission E approved use for constipation. Senna is also used in combination with ispaghula for constipation. The Committee on Proprietary Medicinal Products (CPMP) has adopted core SPCs (Summary of Product Characteristics) for senna leaf and senna fruit (C. angustifolia and C. acutifolia) with indications for short-term use in cases of occasional constipation.
Dosage

**Dried pods**
3–6 pods (Alexandrian) or 4–12 pods (Tinnevelly) steeped in 150 mL warm water for 6–12 hours;\(^{(G6 \ G7)}\) 0.6–0.2 g (equivalent to 20–30 mg hydroxyanthracene glycosides calculated as sennosides B).\(^{(G52)}\)

**Dried leaflets**
0.5–2.0 g\(^{(G6 \ G7)}\) (equivalent to 20–30 mg hydroxyanthracene glycosides calculated as sennoside B).\(^{(G52)}\)

**Leaf, liquid extract**
0.5–2.0 mL (1 : 1 in 25% alcohol).\(^{(G6 \ G7)}\)

**Senna Liquid Extract**
(BPC 1973) 0.5–2.0 mL.

**Herbal drug preparations**
Equivalent to 15–30 mg hydroxyanthracene derivatives (calculated as sennoside B) to be taken at night.\(^{(3)}\)
Pharmacological Actions

The cathartic action of hydroxyanthracene–containing drugs is well recognised and they have been used as laxatives for many years. However, there is still some uncertainty as to the exact mode of action of the hydroxyanthracenes.

It is thought that hydroxyanthracene glycosides are absorbed from the gastrointestinal tract, the aglycones liberated during metabolism and excreted into the colon resulting in stimulation and an increase in peristalsis. However, it has also been suggested that the purgative action of senna is due to the action of intestinal bacteria.\(^{(4)}\) Using human intestinal flora, it was found that sennoside A is reduced to 8–glucosyl rheinanthrone, hydrolysed to rheinanthrone and oxidised to sennidin A. The active principle causing peristaltic movements of the large intestine was thought to be rheinanthrone.\(^{(4)}\)

\textit{In vitro} and animal studies

Sennosides A and B, and their natural metabolites sennidins A and B, have been reported to act specifically on the large intestine in the rat with the acceleration of colonic transport the major component of their laxative effect.\(^{(5)}\) Sennosides A and B have also been reported to induce fluid secretion exclusively in the colon, following oral administration of the glycosides to rats.\(^{(6)}\)

It has been suggested that the laxative action of the sennosides involves prostaglandins. Indomethacin has been found to partly inhibit the action of sennosides A and B, although a bolus injection of prostaglandins into the caecal lumen was stated to neither influence transit time nor to induce diarrhoea.\(^{(5)}\) Pretreatment of mice with indometacin and a prostaglandin E (PGE) antagonist has been documented to prevent diarrhoea caused by intracaecal administration of rhein, which stimulates the production of PGE-like material specifically in the colon.\(^{(7)}\) Indomethacin was found to depress the large intestinal propulsive activity of rhein, but did not suppress PGE\(_2\)-induced diarrhoea. The authors suggest that the action of rhein is mediated by prostaglandin biosynthesis and release.\(^{(7)}\)

Antihepatotoxic activity has been documented for naphtho-\(\alpha\)-pyrrole and naphtho-\(\gamma\)-pyrrole glycosides, and for the hydroxyanthracene glycosides isolated from a related species \textit{Cassia tora}.\(^{(8)}\) Greatest activity was documented for the naphtho-\(\gamma\)-pyrrole glycosides.
Significant inhibitory activity in mice against leukaemia P388 has been documented for aloe–emodin. (G41)

**Clinical studies**

In a randomised, controlled trial, 91 patients with terminal cancer received senna 12 mg daily, or lactulose, for 27 days. (9) At the end of the study, no differences were found between the two groups in defecation–free intervals, or in days with defecation. The general health of each group was also reported to be similar.

A randomised, double–blind, double–dummy, multicentre, controlled, crossover study involving 77 hospitalised elderly patients with a history of chronic constipation compared the effects of a senna–fibre combination (senna 12.4%, ispaghula 54.2%, 10 mL daily) and lactulose (15 mL twice daily) for two 14–day periods with a three- to five–day wash–out period. (10) Assessments included stool frequency and consistency, ease of evacuation, adverse effects and costs of treatment. The senna–fibre combination was reported to be significantly more effective than lactulose. (10)

Commercial preparations containing senna and ispaghula have been reported to be equally effective for the treatment of constipation in small clinical studies involving elderly hospitalised patients and/or residents in nursing homes. (11)
Side-effects, Toxicity

Senna may cause mild abdominal discomfort such as colic or cramps. Prolonged use or overdosage can result in diarrhoea with excessive loss of potassium, albuminuria and haematuria. Potassium deficiency may lead to disorders of the heart and muscular weakness especially with concurrent use of cardiac glycosides, diuretics or corticosteroids. An atonic non-functioning colon may also develop. Excessive use and abuse of senna has been associated with finger clubbing and with the development of cachexia and reduced serum globulin concentrations.

Sennosides A and B are reported to be most potent with respect to laxative action, but to be the least toxic compared with other hydroxyanthracene fractions in senna. Similarly, fractions with a low laxative activity (e.g. rhein–8–glucoside) are reported to have the highest acute toxicity. LD\textsubscript{50} values in mice following intravenous injection of sennosides A and B and of rhein–8–glycoside are reported to be 4.1 g/kg and 400 g/kg, respectively. The acute oral toxicity of all senna fractions in mice has been reported to be greater than 5 g/kg, although all of the animals were stated to have died by the following week. The toxicity of total senna extracts is greater than that of the individual sennosides and it has been proposed that the laxative and toxic components of senna could be separated.

\textit{In vitro} carcinogenicity testing has reported certain anthraquinones, including aloe–emodin, to be active in more than one strain of \textit{Salmonella typhimurium}. Aglycones were documented to exhibit genotoxic activity in a mammalian cell assay.

Sensitising properties have been documented for emodin (see Aloes).

The CPMP core SPCs for senna include the following information. There are no new, systematic preclinical tests for senna leaf or preparations thereof. Most data refer to extracts of senna fruit containing 1.4–3.5% of hydroxyanthracenes, corresponding to 0.9–2.35% of potential rhein, 0.05–0.15% potential aloe–emodin and 0.001–0.006% of potential emodin, or to isolated active constituents, e.g. rhein or sennosides A and B. The acute toxicity of senna fruit and specified extracts thereof, as well as of sennosides in rats and mice was low after oral treatment. As a result of investigations with parenteral application in mice, extracts are supposed to possess a higher toxicity than purified glucosides, possibly due to the content of aglycones.

Sennosides displayed no specific toxicity when tested at doses up to 500 g/kg in dogs for four weeks and up to 100 g/kg in rats for six months. Data for
herbal drug preparations are not available. There was no evidence of any embryolethal, teratogenic or fetotoxic actions in rats or rabbits after oral treatment with sennosides. Furthermore, there was no effect on the postnatal development of young rats, on rearing behaviour of dams or on male and female fertility in rats. Data on herbal drug preparations are not available.

An extract and aloe–emodin were mutagenic in \textit{in vitro} tests; sennosides A and B and rhein gave negative results. \textit{In vivo} examinations of a defined extract of senna pods were negative. A specified senna extract given orally for two years was not carcinogenic in male or female rats. The extract investigated contained approximately 40.8\% of hydroxyanthracenes from which 35\% were sennosides, corresponding to about 25.2\% of potential rhein, 2.3\% of potential aloe–emodin and 0.007\% of potential emodin, and 142 ppm free aloe–emodin and 9 ppm free emodin.
Contra-indications, Warnings

It is recommended that senna should not be given to patients with intestinal obstruction and stenosis, atony, inflammatory colon diseases (e.g. Crohn’s disease, ulcerative colitis), appendicitis, with undiagnosed abdominal symptoms; severe dehydration states with water and electrolyte depletion. Prolonged use should be avoided.\(^{(3,G20 G45 G52)}\)

The CPMP core SPCs for senna include the following information.\(^{(3)}\)

As with all laxatives, senna should not be given when any undiagnosed acute or persistent abdominal symptoms are present. If laxatives are needed every day the cause of the constipation should be investigated. Long–term use of laxatives should be avoided. Use for more than two weeks requires medical supervision. Chronic use may cause pigmentation of the colon (pseudomelanosis coli) which is harmless and reversible after drug discontinuation.

Abuse, with diarrhoea and consequent fluid and electrolyte losses, may cause dependence, with possible need for increased dosages, disturbance of water and electrolyte (mainly hypokalaemia) balance, atonic colon with impaired function. Intake of anthranoid containing laxatives exceeding short–term use may result in an aggravation of constipation.

Hypokalaemia can result in cardiac and neuromuscular dysfunction, especially if cardiac glycosides, diuretics or corticosteroids are also taken. Chronic use may result in albuminuria and haematuria.

In chronic constipation, stimulant laxatives are not an acceptable alternative to a changed diet.

**Interaction with other medicaments and other forms of interaction**\(^{(3)}\)

Hypokalaemia (resulting from long–term use of senna) may potentiate the action of cardiac glycosides and interacts with antiarrhythmic drugs, with drugs which induce reversion to sinus rhythm (e.g. quinidine). Concomitant use with other drugs inducing hypokalaemia (e.g. thiazide diuretics, adreno corticosteroids and liquorice root) may enhance electrolyte imbalance. Abdominal spasms and pain may occur, in particular in patients with irritable colon.\(^{(3)}\)

Anthraquinones cause discoloration of the urine which may interfere with diagnostic tests.\(^{(G45)}\)

**Pregnancy and lactation**
Non-standardised hydroxyanthracene containing laxative preparations should not be taken during pregnancy or lactation since their pharmacological action is unpredictable. Although hydroxyanthracene derivatives may be excreted in the breast milk, following normal dosage their concentration is usually insufficient to affect the nursing infant.\(^{(G45)}\)

The CPMP core SPCs for senna include the following information.\(^{(3)}\)

**Pregnancy** Not recommended during pregnancy. There are no reports of undesirable or damaging effects during pregnancy and on the fetus when used at the recommended dosage schedule. However, experimental data concerning a genotoxic risk of several anthranoids (e.g. emodine and physcione) and senna are not counterbalanced by sufficient studies to eliminate a possible risk.\(^{(3)}\)

**Lactation** Breastfeeding is not recommended as there are insufficient data on the excretion of metabolites in breast milk. Excretion of active principles in breast milk has not been investigated. However, small amounts of active metabolites (e.g. rhein) from other anthranoids are known to be excreted in breast milk. A laxative effect in breastfed babies has not been reported.\(^{(3)}\)
Pharmaceutical Comment

The chemistry of senna is characterised by the hydroxyanthracene derivatives. The laxative action of these compounds is well recognised and supports the herbal use of senna as a laxative for the treatment of constipation. However, the use of non-standardised hydroxyanthracene-containing preparations should be avoided since their pharmacological effect will be variable and unpredictable. The sennoside content of many licensed senna products is standardised and generally calculated as sennoside B. Clinical investigations have concluded that senna with ispaghula is more effective than lactulose as a laxative (see Clinical studies).
References


Shepherd’s Purse
Species (Family)

Capsella bursa-pastoris (L.) Medic (Cruciferae)
Synonym(s)
Capsella
Part(s) Used
Herb
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)

Complete German Commission E\(^{(G3)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL$^{(G37)}$
Constituents
See General References G2 G7 G40 G41 G64.

Amines
Acetylcholine, choline, amino acids 2.33% (major component proline), histamine, tyramine and unidentified crystalline alkaloids.\(^1\)

Flavonoids
Quercetin, diosmetin, luteolin, hesper etin and their glycosides (e.g. rutin, diosmin, hesperidin).\(^2\)

Volatile oils
0.02%. Camphor (major); at least 74 components identified.\(^3,4\)

Other constituents
Carotenoids, fumaric acid, sinigrin (mustard oil glucoside), ascorbic acid (vitamin C) and vitamin K.\(^4,5,G2\)
**Food Use**

Shepherd’s purse is not used in foods.
Herbal Use

Shepherd’s purse is stated to possess antihaemorrhagic and urinary antiseptic properties. Traditionally, it has been used for menorrhagia, haematemesis, haematuria, diarrhoea and acute catarrhal cystitis. (G2 G7 G64)
**Dosage**

*Dried herb*
1–4 g or by infusion three times daily\(^{(G7)}\)

*Liquid extract*
1–4 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

In vitro and animal studies

A variety of actions have been documented for an ethanolic extract of shepherd’s purse in various animal models.\(^{(6–9)}\) Anti-inflammatory activity has been exhibited versus carrageenan–induced and dextran–induced rat paw oedema.\(^{(7)}\) A reduction in capillary permeability in the guinea–pig, induced by histamine and serotonin, has also been observed,\(^{(7)}\) and flavonoid components isolated from shepherd’s purse have been reported to reduce blood vessel permeability in mice.\(^{(2)}\) Anti–ulcer activity has been documented in rats following intraperitoneal injection. The extract did not affect gastric secretion, but accelerated recovery from stress–induced ulcers.\(^{(7)}\) A hypotensive effect observed in cats, dogs, rabbits, and rats, following intravenous injection, was inhibited by a β-adrenoceptor blocker but not by atropine, thus dismissing earlier reports that this action was attributable to cholinergic compounds present in shepherd’s purse.\(^{(8,9)}\)

Diuresis has been reported in mice, following oral or intraperitoneal administration of shepherd’s purse. The mode of action was stated to involve an increase in the glomerular filtration rate.\(^{(7)}\)

Documented cardiac actions include increased coronary blood flow in dogs following intra–arterial administration, and a slight inhibitory effect on ouabain–induced ventricular fibrillation in the rat following intraperitoneal injection, together with a negative chronotropic effect.\(^{(9)}\) Studies on the isolated heart have reported negative chronotropic and inotropic actions in the guinea–pig and rabbit and coronary vasodilatation.\(^{(9)}\)

A CNS-depressant action in mice has been demonstrated (potentiation of barbiturate–induced sleeping time).\(^{(9)}\)

Weak antibacterial activity mainly towards Gram–positive organisms has been reported.\(^{(10)}\)

Antineoplastic activity in rats has been documented for fumaric acid, which prevented the development of hepatic neoplasms when co–administered with the carcinogen 3-MeDAB.\(^{(11)}\)

Shepherd’s purse seeds are stated to possess rubefacient and vesicant properties because of their isothiocyanate–yielding components.\(^{(G51)}\)

In vitro studies have documented stimulatory action in various smooth muscle tissues. Induced contractions of the small intestine in the guinea–pig were
reported to be unaffected by atropine and diphenhydramine, but were inhibited by papaverine.\(^{(8,9)}\) Induced utero–activity in the rat, equivalent to the effect of oxytocin 0.1 i.u., was unaffected by atropine, but inhibited by competitive inhibitors of oxytocin.\(^{(8)}\) Two unidentified alkaloid components of shepherd’s purse have also been stated to elicit a physiological activity on the uterus.\(^{(1)}\) Induced tracheal contractions in the guinea–pig were unaffected by adrenaline, which did inhibit acetylcholine–induced contractions.\(^{(9)}\) These studies concluded that the active substance(s) in shepherd’s purse responsible for the observed actions on smooth muscle were neither acetylcholine nor histamine.\(^{(8,9)}\)
Shepherd’s purse extracts have been reported to exhibit low toxicity in mice. LD$_{50}$ values reported are 1.5 g/kg body weight (mice, intraperitoneal injection) and 31.5 g/kg (mice, subcutaneous injection).\(^{(9)}\) Signs of toxicity were described as sedation, enlargement of pupils, paralysis of hind limbs, difficulty in respiration, and death by respiratory paralysis.\(^{(9)}\) Following hydrolysis, the constituent sinigrin yields allyl isothiocyanate which is an extremely powerful irritant and produces blisters on the skin.\(^{(G41)}\) Isothiocyanates have been implicated in endemic goitre (hypothyroidism with thyroid enlargement) and have been reported to produce goitre in experimental animals.\(^{(G41)}\)
Contra-indications, Warnings

Prolonged or excessive use of the herb may interfere with existing therapy for hyper- or hypotension, thyroid dysfunction or cardiac disorder, and may potentiate sedative actions.

**Pregnancy and lactation**

Shepherd’s purse is reputed to act as an abortifacient and to affect the menstrual cycle, and tyramine is documented as a utero-active constituent.\(^{G30}\) In view of this and the reported oxytocin-like activity, the use of shepherd’s purse during pregnancy should be avoided. Excessive use should be avoided during lactation.
The chemistry of shepherd’s purse is well documented and although a number of actions affecting the circulatory system have been observed in animal studies, these actions do not relate to the traditional herbal uses. Limited toxicity data are available. In view of this together with the demonstrated pharmacological activity of the herb, excessive use of shepherd’s purse should be avoided.
References

See also General References G2 G3 G9 G30 G31 G36 G37 G40 G41 G51 G64.


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Skunk Cabbage
Species (Family)

*Symplocarpus foetidus* (L.) Salisb. (Araceae)
Synonym(s)

*Dracontium foetidum* L., Skunkweed
Part(s) Used
Rhizome, root
Pharmacopoeial and Other Monographs

BHP 1983\(^{(G7)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL\textsuperscript{(G37)}
Constituents

See General References G22 G64.

Reported constituents include starch, gum–sugar, fixed and volatile oils, resin, tannin, an acrid principle and iron.

Other plant parts
Large amounts of alkaloids (unspecified), phenolic compounds and glycosides have been isolated from all plant parts of skunk cabbage.\(^{(1)}\) The leaves are reported to contain hydroxytryptamine;\(^{(G22)}\) three anthocyanin pigments have been isolated from the flowers, namely cyanidin–3–monoglucoside, cyanidin–3–rutinoside and peonidin–3- rutinoside.\(^{(2)}\)
Food Use

Skunk cabbage is not used in foods.
Herbal Use

Skunk cabbage is stated to possess expectorant, antispasmodic and mild sedative properties. Traditionally, it has been used for bronchitis, whooping cough, asthma and specifically for bronchitic asthma. (G7 G64)
Dosage

**Powdered rhizome/root**
0.5–1.0 g in honey or by infusion or decoction three times daily.\(^{(G7)}\)

**Liquid extract**
0.5–1.0 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)

**Tincture**
2–4 mL (1 : 10 in 45% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

*In vitro and animal studies*

None documented for the rhizome/root. The leaf extract has haemolytic properties.\(^{(G22)}\)

*Clinical studies*

None documented.
Side-effects, Toxicity

The root is reported to be bitter and acrid, with a disagreeable odour. Severe itching and inflammation of the skin has been documented.\textsuperscript{G51} No published toxicity studies were located.
Contra-indications, Warnings

It has been stated that the fresh plant can cause blistering.\(^{G42}\) In view of the acrid principle thought to be present in both the dried and fresh root,\(^{G51}\) skunk cabbage should be used with caution.

**Pregnancy and lactation**

Skunk cabbage is reputed to affect the menstrual cycle.\(^{G22}\) In view of the lack of phytochemical, pharmacological, and toxicological information, and the irritant properties, the use of skunk cabbage during pregnancy and lactation should be avoided.
Pharmaceutical Comment

Little is known about the constituents, pharmacological activities or safety of skunk cabbage (even though citings as early as 1817 reported its irritant properties).\(^\text{(G51)}\) No documented evidence was found to justify the herbal uses. In view of the documented irritant properties, excessive use is not recommended.
References

See also General References G7 G22 G31 G36 G37 G42 G51 G64.


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Slippery Elm
Species (Family)

*Ulmus fulva* Michaux (Ulmaceae)
Synonym(s)

*Ulmus rubra* Muhl.
Part(s) Used

Bark (inner)
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL$^{(G37)}$
Constituents

See General References G6 G59 G64.

**Carbohydrates**
Mucilage (major constituent) consisting of hexoses, pentoses, methylpentoses, at least two polyuronides, and yielding on hydrolysis galactose, glucose and fructose (trace), galacturonic acid, L-rhamnose and D-galactose.

**Other constituents**
Tannin 3.0–6.5% (type unspecified), phytosterols (β-sitosterol, citrostadienol, dolichol), sesquiterpenes, calcium oxalate and cholesterol.
Food Use

It has been recommended by the FACC (Food Additives and Contaminants Committee) that the use of slippery elm as a flavouring agent in foods should be prohibited.\(^{(G44)}\) Slippery elm is listed by the Council of Europe as a natural source of food flavouring (category N3). This category indicates that there is insufficient information available to make an adequate assessment of potential toxicity.\(^{(G16)}\)
Herbal Use

Slippery elm is stated to possess demulcent, emollient, nutrient and antitussive properties. Traditionally, it has been used for inflammation or ulceration of the stomach or duodenum, convalescence, colitis, diarrhoea and locally for abscesses, boils and ulcers (as a poultice). (G6 G7 G8 G64)
Dosage

**Powdered bark**
4–16 mL (1 : 8 as a decoction) three times daily. (G6 G7)

**Powdered bark**
4 g in 500 mL boiling water as a nutritional supplement three times daily. (G6 G7)

**Coarse powdered bark**
With boiling water as a poultice. (G6 G7)

**Liquid extract**
5 mL (1 : 1 in 60% alcohol) three times daily. (G6 G7)
Pharmacological Actions

Mucilages are known to have demulcent and emollient properties. Mucilage is the principal constituent of slippery elm. Tannins are known to possess astringent properties.
Side-effects, Toxicity

None documented. In view of the known constituents of slippery elm it would appear to be non-toxic.
Contra-indications, Warnings

Whole bark has been used to procure abortions.

**Pregnancy and lactation**

There are no known problems with the use of powdered slippery elm during pregnancy.
Pharmaceutical Comment

The primary constituent in slippery elm is mucilage, thereby justifying the herbal use of the remedy as a demulcent, emollient and antitussive. There are no known problems regarding toxicity of slippery elm, although its use as a food flavouring agent has not been recommended. The supply of whole bark is controlled by regulations.\(^{(1)}\)
See also General References G5 G6 G9 G11 G22 G31 G36 G37 G43 G59 G64.


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Squill
Species (Family)

*Drimia maritima* (L.) Stearn (Liliaceae)
Synonym(s)
Scilla, Urginea, *Urginea maritima* (L.) Baker, *Urginea scilla* Steinh., White Squill
Part(s) Used

Bulb (red and white varieties)
Pharmacopoeial and Other Monographs

BHC 1992\(^{G6}\)

BHP 1996\(^{G9}\)

BP 2002\(^{G71}\)

Complete German Commission E\(^{G3}\)

Martindale 33rd edition\(^{G67}\)

PDR for Herbal Medicines 2nd edition\(^{G36}\)
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents

See References 1 and 2, and General References G6 G22 G41 G48 G62 G64.

**Cardiac glycosides**
Scillaren A and proscillaridin A (major constituents); others include glucoscillaren A, scillaridin A, scillicyanoside, scilliglaucoside, scilliphaeoside, scillicoeloside, scillazurosode and scillicryptoside. Scillaren B represents a mixture of the squill glycosides.

**Flavonoids**
Apigenin, dihydroquercetin, isovitexin, iso–orientin, luteolin, orientin, quercetin, taxifolin and vitexin.

**Other constituents**
Stigmasterol, tannin, volatile and fixed oils.
Food Use

The Food Additives and Contaminants Committee (FACC) has recommended that squill be prohibited as a food flavouring.\(^{(G45)}\)
Herbal Use

Squill is stated to possess expectorant, cathartic, emetic, cardioactive and diuretic properties. Traditionally, it has been used for chronic bronchitis, asthma with bronchitis, whooping cough, and specifically for chronic bronchitis with scanty sputum.
Dosage

*Dried bulb*
60–200 mg or by infusion three times daily.\(^{G6, G7}\)

*Squill Liquid Extract*
(BPC 1973) 0.06–0.2 mL.

*Squill Tincture*
(BPC 1973) 0.3–2.0 mL.

*Squill Vinegar*
(BPC 1973) 0.6–2.0 mL.
Pharmacological Actions

The aglycone components of the cardiac glycoside constituents possess digitalis–like cardiotonic properties.\(^{(G41)}\) However, the squill aglycones are poorly absorbed from the gastrointestinal tract and are less potent than digitalis cardiac glycosides.\(^{(1,2)}\)

Expectorant, emetic and diuretic properties have been documented for white squill.\(^{(G41)}\) Squill is reported to induce vomiting by both a central action and local gastric irritation.\(^{(1,2)}\) Subemetic or near–emetic doses of squill appear to exhibit an expectorant effect, causing an increase in the flow of gastric secretions.\(^{(1,2)}\)

Antiseborrhoeic properties have been documented for methanol extracts of red squill which have been employed as hair tonics for the treatment of chronic seborrhoea and dandruff.\(^{(G41)}\)

Squill extracts have been reported to exhibit peripheral vasodilatation and bradycardia in anaesthetised rabbits.\(^{(1,2)}\)
Side-effects, Toxicity

Excessive use of squill is potentially toxic because of the cardiotonic constituents. However, squill is also a gastric irritant and large doses will stimulate a vomiting reflex. Red squill is toxic to rats and is mainly used as a rodenticide, causing death by a centrally induced convulsant action.\(^{(1,2)}\) A squill soft mass (crude extract) has been stated to be toxic in guinea-pigs at a dose of 270 mg/kg body weight. A fatal dose for Indian squill (\textit{Urginea indica} Kunth.) is documented as 36 mg/kg.
**Contra-indications, Warnings**

Squill may cause gastric irritation and should be avoided by individuals with a cardiac disorder. In view of the cardiotonic constituents, precautions applied to digoxin therapy should be considered for squill.

**Pregnancy and lactation**

Squill is reputed to be an abortifacient and to affect the menstrual cycle.\(^{(G30)}\) In addition, cardioactive and gastrointestinal irritant properties have been documented. The use of squill during pregnancy should be avoided; excessive use should be avoided during lactation.
Squill is characterised by its cardiac glycoside components and unusual flavonoid constituents. The reputed actions of squill as an expectorant, emetic and cathartic can be attributed to the cardioactive components and squill has been used as an expectorant for many years. However, in view of the documented cardioactive and emetic properties of the aglycones, excessive should be avoided. Red squill is primarily used as a rodenticide.
References


Species (Family)

Collinsonia canadensis L. (Labiatae)
Synonym(s)
Heal-All, Knob Root
Part(s) Used
Rhizome, root
Pharmacopoeial and Other Monographs

BHP 1983\(^{(G7)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Constituents

See General References G40 G48 G49 G64.

Stone root is stated to contain an unidentified alkaloid, mucilage, resin, saponin glycosides, tannins and volatile oil.
Food Use

Stone root is not used in foods.
Herbal Use

Stone root is stated to possess antilithic, litholytic, mild diaphoretic and diuretic properties. Traditionally, it has been used for renal calculus, lithuria, and specifically for urinary calculus. (G7 G64)
Dosage

**Dried root**
1–4 g or by decoction three times daily.\(^{(G7)}\)

**Liquid extract**
1–4 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)

**Tincture**
2–8 mL (1 : 5 in 40% alcohol) three times daily.\(^{(G7)}\)

**Tincture of Collinsonia**
(BPC 1934) 2–8 mL.
Pharmacological Actions

None documented.
Side-effects, Toxicity

None documented.
Contra-indications, Warnings

None documented.

**Pregnancy and lactation**
The safety of stone root has not been established. In view of the lack of phytochemical, pharmacological and toxicological information, the use of stone root during pregnancy and lactation should be avoided.
Pharmaceutical Comment

Information available on the chemistry of stone root is limited and no documented scientific evidence was located to justify the herbal uses. In view of the lack of toxicity data, excessive use of stone root should be avoided.
See General References G7 G10 G31 G36 G37 G40 G48 G49 G64.

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Tansy
Species (Family)

Tanacetum vulgare L. (Asteraceae/Compositae)
Synonym(s)

*Chrysanthemum vulgare* (L.) Bernh., Tanacetum
Part(s) Used

Herb
Pharmacopoeial and Other Monographs

BHP 1983\(^{(G7)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

Tansy is not included in the GSL.\(^{(G37)}\)
Constituents
See General References G22 G51 G64.

**Steroids**
β-Sitosterol (major), campesterol, cholesterol, stigmasterol and taraxasterol.\(^{(1)}\)

**Terpenoids**
a-Amyrin (major), β-amyrin, sesquiterpene lactones including arbusculin-A, tanacetin, germacrene D, crispolide;\(^{(2,3)}\) tanacetols A and B.\(^{(4,5)}\)

**Volatile oils**
0.12–0.18%. Major components as β-thujone (up to 95%) and camphor, others include α-pinene, borneol, 1,8–cineole, umbellone and sabinene. At least ten different chemotypes have been identified in which camphor was the most frequently occurring main component and thujone second.\(^{(4)}\)

**Other constituents**
Gum, mucilage, resin and tannins.
Food Use

Tansy is listed by the Council of Europe as a natural source of food flavouring (Category N3). This category indicates that tansy can be added to foodstuffs in the traditionally accepted manner, although there is insufficient information for an adequate assessment of potential toxicity. In addition, the Council of Europe recommends that the concentration of thujones present in food products is restricted to 0.5 mg/kg.\(^{(G16)}\) Tansy oil is prohibited from use as a food flavouring by the Food Additives and Contaminants Committee (FACC) in view of the thujone content.\(^{(G44)}\)

In the USA, tansy is prohibited from sale by botanical dealers or by mail order as the dried herb.\(^{(G22)}\)
Herbal Use

Tansy is stated to possess anthelmintic, carminative and antispasmodic properties and to act as a stimulant to abdominal viscera. Traditionally, it has been used for nematode infestation, topically for scabies (as a decoction) and pruritus ani (as an ointment), and specifically for roundworm or threadworm infestation in children.\(^\text{G7}\)
Dosage

*Dried herb*
1–2 g or by infusion three times daily.\(^{(G7)}\)

*Liquid extract*
1–2 mL (1:1 in 25% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

**In vitro and animal studies**

*In vitro* antispasmodic activity on rabbit intestine, and *in vivo* choleretic activity in the dog have been documented for tansy extracts.\(^6\) It was suggested that the choleretic action might be attributable to caffeic acid, a known bile stimulant that is present in tansy.\(^6\) Anthelmintic activity in dogs has been described for tansy oil, an ether extract of the oil, and for β-thujone.\(^6\) Daily intragastric doses of a tansy extract given to rabbits have been found to reduce serum lipid concentrations and inhibit further development of hypercholesterol aemia.\(^6\) In addition, it was noted that recovery of blood sugar concentrations was inhibited in animals given twice daily doses. *In vitro* antifungal activity in 15 pathogenic and non-pathogenic fungi has been reported.\(^6\)

**Clinical studies**

Aqueous infusions and alcoholic extracts have been reported to be clinically effective bile stimulants in patients with liver and gall bladder disorders.\(^6\) The treatment alleviated pain and increased appetite and digestion.
Tansy oil contains the toxic ketone β-thujone. Symptoms of tansy oil poisoning are attributable to the thujone content and include rapid and weak pulse, severe gastritis, violent spasms and convulsions. Documented fatalities have mainly been associated with ingestion of the oil, although fatal cases of poisoning have occurred with infusions and powders. An oral LD$_{50}$ value for tansy oil is stated as 1.15 g/kg body weight. The ratio of toxic to therapeutic dose has been reported as 2.5 : 1 and it was noted that all tansy preparations should be administered with castor oil. Tansy yields potentially allergenic sesquiterpene lactones which have been implicated in the aetiology of contact dermatitis. Instances of contact dermatitis to tansy have been documented. In vitro and in vivo antitumour activity has been documented for tansy.
Contra-indications, Warnings

Tansy oil is toxic and should not be used internally or externally. Fatalities have been reported following ingestion of infusions and extracts. Tansy contains allergenic sesquiterpene lactones and may cause an allergic reaction. Tansy has been reported to affect blood sugar concentrations in animals and may interfere with hypoglycaemic therapy.

Pregnancy and lactation

Tansy is contra-indicated in pregnancy and lactation. Tansy is reputed to affect the menstrual cycle and uteroactivity has been documented in animal studies. The volatile oil contains β-thujone, a known hepatotoxin.
Pharmaceutical Comment

Pharmacological activities documented for tansy have been associated with the sterol and triterpene constituents. Tansy yields an extremely toxic volatile oil, which should not be used internally or externally.\(^{(G58)}\) In view of this, the use of tansy as a herbal remedy is not justified even though documented studies have supported the traditional uses of the herb as a choleretic and anthelmintic agent.
References

See also General References G7 G16 G22 G31 G32 G36 G37 G44 G51 G58 G64.


Thyme
Species (Family)

*Thymus vulgaris* L., *Thymus zygis* L. (Labiatae)
Synonym(s)
Common Thyme, French Thyme, Garden Thyme, Rubbed Thyme
Part(s) Used

Flowering top, leaf
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)
BP 2002\(^{(G71)}\)
Complete German Commission E\(^{(G3)}\)
ESCOP 1996\(^{(G52)}\)
Martindale 33rd edition\(^{(G67)}\)
Mills and Bone\(^{(G50)}\)
PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Ph Eur 2004\(^{(G72)}\)
WHO volume 1 1999\(^{(G63)}\)
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents

See General References G2 G22 G41 G52 G58 G64.

Volatile oils
0.8–2.6%. Pharmacopoeial standard, not less than 1.2%.(G15) Phenols as major components (20–80%) primarily thymol and carvacrol; others include p-cymene and γ-terpinene (mono terpenes), linalool, α-terpineol, and thujan-4-ol (alcohols); biphenyl compounds of monoterpenic origin.(G52) A detailed analysis of the volatile oil components is given elsewhere.(G22)

Flavonoids
Cirsilineol, 8–methoxycirsilineol, thymonin and eriodictyol.

Other constituents
Caffeic acid, oleanolic acid, ursolic acid, rosmarinic acid, resins, saponins and tannins.
Food Use

Thyme is commonly used as a culinary herb, and thyme oil is used in food flavouring. In the USA, thyme is listed as GRAS (Generally Recognised As Safe).\(^{(G65)}\)
Thyme is stated to possess carminative, antispasmodic, antitussive, expectorant, secretomotor, bactericidal, anthelmintic and astringent properties. Traditionally, it has been used for dyspepsia, chronic gastritis, asthma, diarrhoea in children, enuresis in children, laryngitis, tonsillitis (as a gargle), and specifically for pertussis and bronchitis. The German Commission E approved internal use for treating symptoms of bronchitis, whooping cough and catarrh of the upper respiratory tract. Thyme is used in various combinations with anise oil, eucalyptus oil, fennel oil, fennel fruit, Iceland moss, lime flower, liquorice root, marshmallow root, primrose root and star anise fruit for catarrh and diseases of the upper respiratory tract.
Dosage

*Dried herb*
1–4 g or by infusion three times daily;\(^{(G7)}\) 1–2 g.\(^{(G3 \ G52)}\)

*Liquid Extract of Thyme*
(BPC 1949) 0.6–4.0 mL.

*Elixir of Thyme*
(BPC 1949) 4–8 mL.

*Tincture*
2–6 mL (1 : 5 in 45% alcohol) three times daily,\(^{(G7)}\) four drops.\(^{(G3 \ G52)}\)
Pharmacological Actions

In vitro and animal studies

Antitussive, expectorant and antispasmodic actions are considered to be the major pharmacological properties of thyme,\(^1\) and have been associated with the volatile oils (e.g. thymol, carvacrol) and flavonoid constituents. Thyme oil has produced hypotensive and respiratory stimulant effects in rabbits following oral or intramuscular administration, and in cats following intravenous injection;\(^{G41}\) an increase in rhythmic heart contraction was also observed in rabbits.\(^{G41}\) Hypotensive activity in rats has been reported for *Thymus orospedanus*; this action was attributed to adrenaline (epinephrine) antagonism.\(^2\)

*In vitro* antispasmodic activity of thyme and related *Thymus* species has been associated with the phenolic components of the volatile oil\(^3\) and with the flavonoid constituents; their mode of action is thought to involve calcium–channel blockage.\(^{1,4,5}\) The flavonoids thymonin, circilineol and 8–methoxycircilineol have potent spasmolytic activity in guinea pig trachea preparations *in vitro*.\(^{G52}\)

Analgesic and antipyretic properties in mice have been reported for a thyme extract.\(^6\)

Thymol possesses anthelmintic (especially hookworms), antibacterial, and antifungal properties.\(^{G41}\) The antibacterial activity of thymol and thyme oil have been reviewed.\(^{G50}\) Thymol, carvacrol and thyme oil have antifungal activity against a range of organisms.\(^{G50}\)

Thyme oil inhibits prostaglandin synthesis; rosmarinic acid has anti–inflammatory activity, inhibiting complement in rats and some of the functions of polymorphonucleocytes.\(^{G52}\) Rosmarinic acid reduced oedema produced by cobra venom factor in rats, and inhibited passive cutaneous anaphylaxis and impairment of *in vivo* activation of mouse macrophages by heat killed *Corynebacterium parvum*.\(^{G52}\) Activity may relate to complement inactivation.\(^{G50}\)

Clinical studies

Generally, well–designed clinical studies assessing the effects of thyme are lacking. A randomised, double–blind, controlled trial involving 60 patients with productive cough compared syrup of thyme and bromhexine over a five–day period. Both groups were similar in self–reported symptom relief.\(^{G50}\)
Thyme oil has been used for the treatment of enuresis in children.\textsuperscript{(G44)}
Side-effects, Toxicity\textsuperscript{(G58)}

Thyme oil is a dermal and mucous membrane irritant.\textsuperscript{(G58)} Toxic symptoms documented for thymol include nausea, vomiting, gastric pain, headache, dizziness, convulsions, coma, and cardiac and respiratory arrest.\textsuperscript{(G22)} Thymol is present in some toothpaste preparations, and has been reported to cause cheilitis and glossitis. Hyperaemia and severe inflammation have been described for thyme oil used in bath preparations.\textsuperscript{(G51)}

A concentrated extract of thyme decreased locomotor activity and caused a slight slowing down of respiration in mice following oral administration of doses of 0.5–3.0 g/kg, equivalent to 4.3–26.0 g dried plant material.\textsuperscript{(G52)} In rats, oral LD\textsubscript{50} values stated for thyme oil include 2.84 g/kg\textsuperscript{(G52)} and 4.7 g/kg in rats, and >5 g/kg following dermal administration.\textsuperscript{(7)} In mice, oral administration of a concentrated ethanol extract of herb in subacute toxicity tests resulted in increased weights of liver and testes. Also in mice, a dose of 0.9 g daily for three months resulted in mortality rates of 30% and 10% in males and females, respectively. Thyme oil had no mutagenic or DNA-damaging activity in either the Ames test or \textit{Bacillus subtilis} rec–assay.\textsuperscript{(G52)}
Contra-indications, Warnings

Thyme oil is toxic and should be used with considerable caution. It should not be taken internally and only applied externally if diluted in a suitable carrier oil.

Pregnancy and lactation
There are no known problems with the use of thyme during pregnancy and lactation, provided that doses do not greatly exceed the amounts used in foods. Traditionally, thyme is reputed to affect the menstrual cycle and, therefore, large amounts should not be ingested.
Pharmaceutical Comment

Thyme is commonly used as a culinary herb and is characterised by its volatile oil. Documented pharmacological actions support some of the traditional medicinal uses, which have been principally attributed to the volatile oil and flavonoid constituents. However, the oil is also toxic and should not be ingested and only applied externally if diluted in a suitable carrier oil. It has been suggested that standardised thyme extracts based on the phenolic volatile components may not be appropriate because antispasmodic actions previously attributed to these compounds may be attributable to other constituents.\(^3\)

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Species (Family)

*Arctostaphylos uva-ursi* (L.) Spreng (Ericaceae)
Synonym(s)

Bearberry
Part(s) Used

Leaf
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

BP 2002\(^{(G71)}\)

Complete German Commission E\(^{(G3)}\)

ESCOP 1997\(^{(G52)}\)

Martindale 33rd edition\(^{(G67)}\)

Mills and Bone\(^{(G50)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)

Ph Eur 2004\(^{(G72)}\)
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents


**Flavonoids**
Flavonols (e.g. myricetin, quercetin) and their glycosides including hyperin, isoquercitrin, myricitrin and quercitrin.

**Iridoids**
Asperuloside (disputed), monotropein.\(^{(1)}\)

**Quinones**
Total content at least 6%, mainly arbutin (5–15%) and methyl–arbutin (glycosides), with lesser amounts of piceoside\(^{(2)}\) (a glycoside), free hydroquinone and free \(p\)-methoxyphenol.\(^{(3)}\)

**Tannins**
6–7% (range 6–40%). Hydrolysable–type (e.g. corilagin pyranoside); ellagic and gallic acids (usually associated with hydrolysable tannins).

**Terpenoids**
\(\alpha\)-Amyrin, \(\alpha\)-amyrin acetate, \(\beta\)-amyrin, lupeol, uvaol, ursolic acid, and a mixture of mono- and di–ketonic \(\alpha\)-amyrin derivatives.\(^{(4,5)}\)

**Other constituents**
Acids (malic, quinic), allantoin, resin (e.g. ursone), volatile oil (trace) and wax.

**Other plant parts**
The root is reported to contain unedoside (iridoid glucoside).\(^{(6)}\)
Food Use

Uva-ursi is not used in foods.
Herbal Use

Uva-ursi is stated to possess diuretic, urinary anti septic, and astringent properties. Traditionally, it has been used for cystitis, urethritis, dysuria, pyelitis, lithuria, and specifically for acute catarrhal cystitis with dysuria and highly acidic urine. (G2 G6 G7 G8 G64)
Dosage

**Dried leaves**
1.5–4.0 g or by infusion three times daily. (G6 G7)

**Liquid extract**
1.5–4.0 mL (1 : 1 in 25% alcohol) three times daily. (G6 G7)

**Concentrated Infusion of Bearberry**
(BPC 1934) 2–4 mL.

**Fresh Infusion of Bearberry**
(BPC 1934) 15–30 mL.
Pharmacological Actions

In vitro and animal studies

Uva-ursi has exhibited antimicrobial activity towards a variety of organisms including Staphylococcus aureus, Bacillus subtilis, Escherichia coli, Mycobacterium smegmatis, Shigella sonnei and Shigella flexneri.\(^7\) The antimicrobial activity of arbutin towards bacteria implicated in producing urinary tract infections, has been found to be directly dependent on the β-glucosidase activity of the infective organism.\(^8\) Highest enzymatic activity was shown by Enterobacter, Klebsiella and Streptococcus genera, and lowest by Escherichia coli.\(^8\) The minimum inhibitory concentration for arbutin is reported to be 0.4–0.8% depending on the micro-organism.\(^8\) Aqueous and methanolic extracts have demonstrated molluscicidal activity against Biomphalaria glabrata, at a concentration of 50 ppm.\(^9\) The activity was attributed to the tannin constituents (condensed and hydrolysable).

Anti-inflammatory activity (rat paw oedema tests) has been documented for uva-ursi against a variety of chemical inducers such as carrageenan, histamine and prostaglandins.\(^10\)

Uva-ursi failed to exhibit any in vitro uterotonic action when tested on rabbit and guinea-pig uteri.\(^11\)

Hydroquinone has been reported to show a dose-dependent cytotoxic activity on cultured rat hepatoma cells (HTC line); arbutin was not found to inhibit growth of the HTC cells.\(^12\) It was stated that hydroquinone appeared to have greater cytotoxic activity towards rat hepatoma cells than agents like azauridin or colchicine, but less than valtrate from valerian (Valeriana officinalis). The cytotoxicity of hydroquinone has also been tested on L1210, CA-755 and S-180 tumour systems.\(^12\)

Clinical studies

A herbal preparation, whose ingredients included uva-ursi, hops and peppermint, has been used to treat patients suffering from compulsive strangury, enuresis and painful micturition.\(^13\) Of 915 patients treated for six weeks, success was reported in about 70%. The antiseptic and diuretic properties claimed for uva-ursi can be attributed to the hydroquinone derivatives, especially arbutin. The latter is absorbed from the gastrointestinal tract virtually unchanged and during renal excretion is hydrolysed to yield the active principle, hydroquinone, which exerts an antiseptic and astringent action on the urinary mucous membranes.\(^14,15\) The crude extract
is reported to be more effective than isolated arbutin as an astringent and antiseptic.\textsuperscript{(G48)} This may be due to the other hydroquinone derivatives, in addition to arbutin, that are present in the crude extract and which will also yield hydroquinone. Furthermore, it has been stated that the presence of gallic acid in the crude extract may prevent β-glucosidase cleavage of arbutin in the gastrointestinal tract before absorption, thereby increasing the amount of hydroquinone released during renal excretion.\textsuperscript{(G48)}
Side-effects, Toxicity

No reported side-effects were located. Hydroquinone is reported to be toxic if ingested in large quantities: 1 g (equivalent to 6–20 g plant material) has caused tinnitus, nausea and vomiting, sense of suffocation, shortness of breath, cyanosis, convulsions, delirium and collapse.\(^{(G48)}\) A dose of 5 g (equivalent to 30–100 g of plant material) has proved fatal.\(^{(G48)}\) In view of the high tannin content, prolonged use of uva-ursi may cause chronic liver impairment.\(^{(G41)}\)

Cytotoxic activity has been documented for hydroquinone (see \textit{In vitro} and animal studies).

Uva-ursi herb can sometimes be adulterated with box leaves (\textit{Buxus sempervirens}), which contain toxic steroidal alkaloids. However, no cases of poisoning as a result of such adulteration have been reported.\(^{(G33)}\)
Contra-indications, Warnings

Uva-ursi requires an alkaline urine for it to be effective as a urinary antiseptic; an alkaline reaction is needed to yield hydroquinone from the inactive esters such as arbutin.¹⁴ Patients have been advised to avoid eating highly acidic foods, such as acidic fruits and their juices.¹⁴ The presence of hydroquinone may impart a greenish-brown colour to the urine, which darkens following exposure to air due to oxidation of hydroquinone.

Excessive use of uva-ursi should be avoided in view of the high tannin content and potential toxicity of hydroquinone.

Prolonged use of uva-ursi to treat a urinary tract infection is not advisable. Patients in whom symptoms persist for longer than 48 hours should consult their doctor.

Pregnancy and lactation
Large doses of uva-ursi are reported to be oxytocic,⁵²² although in vitro studies have reported a lack of utero-activity. In view of the potential toxicity of hydroquinone, the use of uva-ursi during pregnancy and lactation is best avoided.
Pharmaceutical Comment

The chemistry of uva–ursi is well documented with hydroquinone derivatives, especially arbutin, identified as the major active constituents. Documented pharmacological actions justify the herbal use of uva–ursi as a urinary antiseptic. However, clinical information is lacking and further studies are required to determine the true usefulness of uva–ursi in the treatment of urinary tract infections. Although hydroquinone has been reported to be toxic in large amounts, concentrations provided by the ingestion of therapeutic doses of uva–ursi are not thought to represent a risk to human health.\(^{(G42)}\)
References


Species (Family)

*Valeriana officinalis* L.s.l. (Valerianaceae)
Synonym(s)
All-Heal, Belgian Valerian, Common Valerian, Fragrant Valerian, Garden Valerian
Part(s) Used

Rhizome, root
Pharmacopoeial and Other Monographs

American Herbal Pharmacopoeia\(^{1,G1}\)

BHC 1992\(^{G6}\)

BHP 1996\(^{G9}\)

BP 2002\(^{G71}\)

Complete German Commission E\(^{G3}\)

EMEA HMPWG proposed core SPC\(^{G23}\)

ESCOP 1997\(^{G52}\)

Martindale 33rd edition\(^{G67}\)

Mills and Bone\(^{G50}\)

PDR for Herbal Medicines 2nd edition\(^{G36}\)

Ph Eur 2004\(^{G72}\)

USP26/NF21\(^{G73}\)

WHO volume 1 1999\(^{G63}\)
Legal Category (Licensed Products)

GSL\textsuperscript{(G37)}
Constituents

Alkaloids
Pyridine type. Actinidine, chatinine, skyanthine, valerianine and valerine.

Iridoids (valepotriates)
Valtrates (e.g. valtrate, valtrate isovaleroxyhydrin, acevaltrate, valechlorine), didrovaltrates (e.g. didrovaltrate, homodidrovaltrate, deoxydidrovaltrate, homodeoxydidrovaltrate, isovaleroxyhydroxydidrovaltrate) and isovaltrates (e.g. isovaltrate, 7-epideacetylisovaltrate). Valtrate and didrovaltrate are documented as the major components. Valerosidate (iridoid glucoside).\(^{(6)}\) The valepotriates are unstable and decompose on storage or processing; the main degradation products are baldrinal and homobaldrinal. The baldrinals may react further and are unlikely to be present in finished products.

Volatile oils
0.5–2%. Not less than 3 mL/kg of essential oil for the cut drug, both calculated with reference to the dried drug.\(^{(G28)}\)

Numerous identified components include monoterpenes (e.g. α- and β-pinene, camphene, borneol, eugenol, isoeugenol) present mainly as esters, sesquiterpenes (e.g. β-bisabolene, caryophyllene, valeranone, ledol, pacifigorgiol, patchouli alcohol, valerianol, valerenol and a series of valerenyl esters, valerenal, valerenic acid with acetoxy and hydroxy derivatives).\(^{(7–10)}\)

Other constituents
Amino acids (e.g. arginine, γ-aminobutyric acid (GABA), glutamine, tyrosine),\(^{(1,11,G21)}\) caffeic and chlorogenic acids (polyphenolic), β-sitosterol, methyl 2-pyrrolketone, choline, tannins (type unspecified), gum and resin.

As with other plants, there can be variation in the content of active compounds (e.g. valerenic acid derivatives and valepotriates) found in valerian rhizomes and roots.\(^{(12)}\)
Food Use

Valerian is not generally used as a food. Valerian is listed by the Council of Europe as a natural source of food flavouring (root: category 5) (see Appendix 23). In the USA, valerian is permitted for use in food.
Valerian is stated to possess sedative, mild anodyne, hypnotic, antispasmodic, carminative and hypotensive properties. Traditionally, it has been used for hysterical states, excitability, insomnia, hypochondriasis, migraine, cramp, intestinal colic, rheumatic pains, dysmenorrhoea, and specifically for conditions presenting nervous excitability. Modern interest in valerian is focused on its use as a sedative and hypnotic.

A core Summary of Product Characteristics (SPC) proposed by the European Medicines Evaluation Agency Herbal Medicinal Product Working Group (EMEA HMPWG) states the following indications: the relief of temporary mild nervous tension and temporary difficulty in falling asleep.
Dosage

**Dried rhizome/root**
1–3 g by infusion or decoction up to three times daily. (G6)

**Tincture**
3–5 mL (1 : 5; 70% ethanol) up to three times daily; (G6 G50) 1–3 mL, once to several times daily. (G3)

**Extracts**
Amount equivalent to 2–3 g drug, once to several times daily; (G3) 2–6 mL of 1 : 2 liquid extract daily. (G50)

Doses given in older texts vary. For example: Valerian Liquid Extract (BPC 1963) 0.3–1.0 mL; Simple Tincture of Valerian (BPC 1949) 4–8 mL; Concentrated Valerian Infusion (BPC 1963) 2–4 mL.

Clinical trials investigating the effects of valerian extracts on sleep parameters have used varying dosages, for example, valerian extract 400 mg/day (drug : extract ratio of 3 : 1)(13) to 1215 mg/day (drug : extract ratio of 5 to 6 : 1).(14)
Pharmacological Actions

It remains unclear precisely which of the constituents of valerian are responsible for its sedative properties.\(^{(5)}\) Attention had focused on the volatile oil, and then the valepotriates and their degradation products, as the constituents responsible. However, it appeared that the effects of the volatile oil could not account for the whole action of the drug, and the valepotriates, which degrade rapidly, are unlikely to be present in finished products in significant concentrations. Current thinking is that the overall effect of valerian is due to several different groups of constituents and their varying mechanisms of action. Therefore, the activity of different valerian preparations will depend on their content and concentrations of several types of constituent.\(^{(4)}\) One mechanism of action is likely to involve increased concentrations of the inhibitory transmitter GABA in the brain. Increased concentrations of GABA are associated with a decrease in CNS activity and this action may, therefore, be involved in the reported sedative activity.

*In vitro* and animal studies

Sedative properties have been documented for valerian and have been attributed to both the volatile oil and valepotriate fractions.\(^{(15,16)}\) Screening of the volatile oil components for sedative activity concluded valerenal and valerenic acid to be the most active compounds, causing ataxia in mice at a dose of 50 mg/kg by intraperitoneal injection.\(^{(15)}\) Further studies in mice described valerenic acid as a general CNS depressant similar to pentobarbitone, requiring high doses (100 mg/kg by intraperitoneal injection) for activity.\(^{(17)}\) A dose of 400 mg/kg resulted in muscle spasms, convulsions and death.\(^{(17)}\) Valerenic acid was also reported to prolong pentobarbitone-induced sleep in mice, resulting in a hangover effect. Biochemical studies have documented that valerenic acid inhibits the enzyme system responsible for the central catabolism of GABA.\(^{(18)}\) An aqueous extract of roots and rhizomes of *V. officinalis* (standardised to 55 mg valerenic acids per 100 g extract) inhibited the uptake and stimulated the release of radiolabelled GABA in isolated synaptosomes from rat brain cortex.\(^{(19,20)}\) Further work suggested that this aqueous extract of valerian induces the release of GABA by reversal of the GABA carrier, and that the mechanism is Na\(^+\) dependent and Ca\(^{2+}\) independent.\(^{(20)}\) The extract contained a high concentration of GABA (about 5 mmol/L) which was shown to be sufficient to induce the release of radiolabelled GABA by this type of mechanism.\(^{(21)}\) Aqueous and hydroalcoholic (ethanol) extracts of valerian root displaced radiolabelled muscimol binding to synaptic membranes (a measure of the influence of drugs on GABA\(_A\) receptors). However, valerenic acid (0.1 mmol/L) did not displace radiolabelled muscimol in this model.\(^{(22)}\) Other *in vitro* studies using...
rat brain tissue have shown that hydroalcoholic and aqueous total extracts of *V. officinalis* root, and an aqueous fraction derived from the hydroalcoholic extract, show affinity for GABA<sub>A</sub> receptors, although far lower than that of the neurotransmitter itself.<sup>(23)</sup> However, a lipophilic fraction of the hydroalcoholic extract, hydroxyvalerenic acid and dihydrovaltrate did not show any affinity for the GABA<sub>A</sub> receptor in this model.

The effects of valerian extracts on benzodiazepine binding to rat cortical membranes have also been explored. Very low concentrations of ethanolic extract of *V. officinalis* had no effect on radiolabelled flunitrazepam binding in this model, although concentrations of 10<sup>−10</sup> to 10<sup>−8</sup> mg/mL increased radiolabelled flunitrazepam binding with an EC<sub>50</sub> of 4.13 × 10<sup>−10</sup> mg/mL.<sup>(24)</sup> However, flunitrazepam binding was inhibited at higher concentrations (0.5–7.0 mg/mL) of valerian extract (IC<sub>50</sub> 4.82 × 10<sup>−1</sup> mg/mL). In other investigations, valerian extract potentiated radiolabelled GABA release from rat hippocampal slices, and inhibited synaptosomal GABA uptake, confirming the effects of valerian extract on GABA<sub>A</sub> receptors.<sup>(24)</sup>

CNS-depressant activities in mice following intraperitoneal injection have been documented for the valepotriates and for their degradation products, although activity was found to be greatly reduced following oral administration.<sup>(25)</sup> A study explored the effects of a mixture of valepotriates on the behaviour of diazepam–withdrawn male Wistar rats in the elevated plus–maze test (a measure of the anxiolytic or anxiogenic properties of drugs).<sup>(26)</sup> Rats were given diazepam (up to 5 mg/kg for 28 days) then vehicle only for three days to induce a withdrawal syndrome. Rats given diazepam or a mixture of valepotriates (dihydrovaltrate 80%, valtrate 15% and acevaltrate 5%) administered intraperitoneally (12 mg/kg) spent a significantly greater proportion of time in the ‘open’ arms of the maze than did those in the control group.

Another specific valepotriate fraction, Vpt<sub>2</sub>, has been documented to exhibit tranquillising, central myorelaxant, anticonvulsant, coronaro–dilating and anti–arrhythmic actions in mice, rabbits, and cats.<sup>(27,28)</sup> The fraction was reported to prevent arrhythmias induced by Pituitrin vasopressin and barium chloride, and to exhibit moderate positive inotropic and negative chronotropic effects.

Antispasmodic activity on intact and isolated guinea–pig ileum has been documented for isovaltrate, valtrate and valeranone.<sup>(29)</sup> This activity was attributed to a direct action on the smooth muscle receptors rather than ganglion receptors. Valerian oil has been reported to exhibit antispasmodic
activity on isolated guinea-pig uterine muscle, but proved inactive when tested in vivo.

*In vitro* inactivation of complement activation has been reported for the valepotriates.

*In vitro* cytotoxicity (inhibition of DNA and protein synthesis, and potent alkylating activity) has been documented for the valepotriates, with valtrate stated to be the most toxic compound. Valepotriates (valtrate and didrovaltrate) isolated from the related species *Valeriana wallichii*, and baldrinal (a degradation product of valtrate) have been tested for their cytotoxic activity *in vitro* using cultured rat hepatoma cells. Valtrate was the most active compound in this system, leading to a 100% mortality of hepatoma cells after 24 hours’ incubation at a concentration of 33 μg/mL. More detailed studies using the same system showed that didrovaltrate demonstrated cytotoxic activity when incubated at concentrations higher than 8 μg/mL of culture (1.5 × 10⁻⁵ mol/L) and led to 100% cellular mortality with 24 hours of incubation at a concentration of 66 μg/mL. The cytotoxic effect of didrovaltrate was irreversible within 2 hours of incubation with hepatoma cells. In mice, administration of intraperitoneal didrovaltrate led to a regression of Krebs II ascitic tumours, compared with control. A subsequent *in vivo* study, in which valtrate was administered to mice (by intraperitoneal injection and by mouth), did not report any toxic effects on haematopoietic precursor cells when compared with control groups. The valepotriates are known to be unstable compounds in both acidic and alkaline media and it has been suggested that their *in vivo* toxicity is limited due to poor absorption and/or distribution. Baldrinal and homobaldrinal, decomposition products of valtrate and isovaltrate respectively, have exhibited direct mutagenic activity against various *Salmonella* strains *in vitro*.

**Clinical studies**

Numerous studies have explored the effects of valerian preparations on subjective and/or objective sleep parameters. Collectively, the findings of these studies are difficult to interpret, as different studies have assessed different valerian preparations and different dosages, and some have involved healthy volunteers whereas others have involved patients with diagnosed sleep disorders. In addition, other studies have used different subjective and/or objective outcome measures, and some have been conducted in sleep laboratories, whereas others have assessed participants receiving valerian whilst sleeping at home. Overall, several, but not all, studies have documented a hypnotic effect for valerian preparations with
regard to subjective measures of sleep quality, and some have documented effects on objective measures of sleep structure. There is a view that subjective measures of sleep quality may be the most appropriate assessment.\textsuperscript{(45)}

A systematic review of randomised, double-blind, placebo-controlled trials of valerian preparations included nine studies.\textsuperscript{(46)} The review concluded that the evidence for valerian as a treatment for insomnia is inconclusive and that there is a need for further rigorous trials. Several of the studies included in the review, and other studies, are discussed in more detail below.

A placebo-controlled study involving 128 volunteers explored the effects of an aqueous extract of valerian root (400 mg) and a proprietary preparation of valerian and hops (Hova) on subjective measures of sleep quality. Each participant took each of the three preparations at night for three non-consecutive nights.\textsuperscript{(13)} On the basis of participants’ self-assessment, valerian significantly reduced sleep latency (time to onset of sleep) and improved sleep quality, compared with placebo ($p < 0.05$). Subgroup analysis suggested that the effects of valerian were most marked among participants who described themselves as ‘poor’ or ‘irregular’ sleepers.\textsuperscript{(13)} It was reported that Hova did not significantly affect sleep latency or sleep quality, compared with placebo, only that Hova administration was associated with an increase in the number of reports of ‘feeling more sleepy than usual the next morning’ (i.e. a ‘hangover’ effect). The authors were unable to explain this discrepancy in the results for the two preparations.

In a subsequent study, eight volunteers with mild insomnia each received aqueous valerian extract 450 mg, 900 mg or placebo, in a random-order experimental design over almost three weeks.\textsuperscript{(37)} The time to the first period of 5 consecutive minutes without movement, measured using wrist-worn activity meters, was used as an objective measure of sleep latency. For this parameter, valerian 450 mg significantly reduced the mean sleep latency, compared with placebo, although there was no further reduction in sleep latency with valerian 900 mg. Subjective assessments indicated that participants were more likely to experience a ‘hangover’ effect with valerian 900 mg.\textsuperscript{(37)}

The same dosages of aqueous valerian extract were tested for their effects on sleep latency and wake time after sleep onset in healthy volunteers who were either sleeping at home or in a sleep laboratory.\textsuperscript{(38)} Each participant sleeping at home took valerian 450 mg, 900 mg, or placebo, for two consecutive nights on a double-blind, crossover schedule. Participants sleeping under laboratory conditions were randomly assigned to receive valerian 900 mg on
the second or third night of the four nights of the study; placebo was taken on the other nights. Under home conditions, valerian 450 mg and 900 mg significantly reduced subjectively measured sleep latency, compared with placebo. Under laboratory conditions, there were no statistically significant differences between valerian 900 mg and placebo on subjective or objective sleep parameters. It was suggested that the ‘more stressful’ sleep environment of the laboratory may have masked the hypnotic effects of valerian. (38)

A randomised, double-blind, placebo-controlled, crossover study involving 16 patients with previously established psychophysiological insomnia according to International Classification of Sleep Disorders (ICSD) criteria and confirmed by polysomnography assessed the effects of single-dose and longer term administration of valerian root extract on objective parameters of sleep structure and subjective parameters of sleep quality. (39) Participants received valerian root extract (Sedonium; drug : extract ratio 5 : 1) 600 mg, or placebo, 1 hour before bedtime for 14 days, followed by a wash-out period of 13 days, before crossing over to the other arm of the study. There were no statistically significant effects on objective and subjective parameters of sleep following single-dose valerian administration. After long-term treatment, sleep efficiency (ratio of time spent asleep to time spent in bed) improved in both the valerian and placebo groups, compared with baseline values, although there were no significant differences between groups. There was a statistically significant difference with valerian on parameters of slow-wave sleep, compared with baseline values, which did not occur with placebo. However, it is not clear if this difference was significantly different for valerian, compared with placebo, as no p-value was given.

In a randomised, double-blind, pilot study, 14 elderly women who were poor sleepers received valerian aqueous extract (Valdispert forte; drug : extract ratio 5 to 6 : 1), or placebo, for eight consecutive days. (14) Valerian 405 mg was administered one hour before sleep for one night in the laboratory, then taken three times daily for the following seven days. There was no difference in sleep parameters between valerian extract and placebo after acute administration. Valerian recipients showed an increase in slow-wave sleep, compared with baseline values. However, valerian had no effect on sleep onset time, rapid eye movement (REM) sleep or on self-rated sleep quality.

Aqueous ethanolic valerian extract (Sedonium) was compared with placebo in a randomised, double-blind trial involving 121 patients with insomnia not due to organic causes. (40) Participants received valerian extract, or placebo, 600 mg one hour before bedtime for 28 days. At the end of the study, valerian extract achieved a significantly higher clinical global impression
score than did placebo. Sleep quality improved in both groups, compared with baseline values.

The effects of valerian extracts on sleep parameters have been compared with those of the benzodiazepine oxazepam.\textsuperscript{(41)} This randomised, double-blind trial involving people with non-organic and non-psychiatric insomnia compared valerian root extract 600 mg with oxazepam 10 mg; treatment was taken 30 minutes before going to bed for 28 days. At the end of the treatment period, sleep quality had improved significantly ($p < 0.001$) in both groups, compared with baseline values. There was no difference between the two groups with regard to sleep quality.

An open, uncontrolled, multicentre study assessed the effects of a valerian extract (Baldrian-Dispert) 45 mg daily in 11 168 patients with sleep disorders. Valerian was rated as ‘good’ or ‘very good’ in 72% of cases of sleep disturbances, 76% of cases of discontinuous sleep, and in 72% of cases of restlessness and tension.\textsuperscript{(42)}

Several other studies have assessed the effects of valerian extract in combination with other herb extracts, such as hops (\textit{Humulus lupulus}) and/or melissa (\textit{Melissa officinalis}), on measures of sleep.\textsuperscript{(47–50)} A randomised, double-blind trial involving healthy volunteers who received Songha Night (\textit{V. officinalis} root extract 120 mg and \textit{M. officinalis} leaf extract 80 mg) three tablets daily taken as one dose 30 minutes before bedtime for 30 days ($n = 66$), or placebo ($n = 32$), found that the proportion of participants reporting an improvement in sleep quality was significantly greater for the treatment group, compared with the placebo group (33.3% versus 9.4%, respectively; $p = 0.04$).\textsuperscript{(48)} However, analysis of visual analogue scale scores revealed only a slight, but statistically non-significant, improvement in sleep quality in both groups over the treatment period. Another double-blind, placebo-controlled trial involving patients with insomnia who received Euvegal forte (valerian extract 160 mg and lemon balm extract 80 mg) two tablets daily for two weeks reported significant improvements in sleep quality in recipients of the herbal preparation, compared with placebo recipients.\textsuperscript{(49)} A placebo-controlled study involving ‘poor sleepers’ who received Euvegal forte reported significant improvements in sleep efficiency and in sleep stages 3 and 4 in the treatment group, compared with placebo recipients.\textsuperscript{(50)}

Some studies assessing combination valerian preparations have compared the effects with those of benzodiazepines.\textsuperscript{(51,52)} A three-week, randomised, double-blind trial reported that a combination of valerian and hops (200 mg and 45.5 mg dry extract, respectively) was equivalent to bromazepam 3 mg with regard to sleep quality in patients with ‘environmental’ sleep disorders.
A study assessing the ‘hangover’ effects of valerian preparations (valerian syrup and valerian–hops tablets) (see Side–effects, Toxicity) reported that subjective measures of sleep quality improved in both valerian groups, compared with placebo.\(^{(52)}\)

In an open, uncontrolled, multicentre study, 225 individuals who were experiencing nervous agitation and/or difficulties falling asleep and achieving uninterrupted sleep were treated for two weeks with a combination preparation containing extracts of valerian root, hop grains and lemon balm leaves.\(^{(53)}\) Significant improvements in the severity and frequency of symptoms were reported, compared with the pretreatment period. Difficulties falling asleep, difficulties sleeping through the night, and nervous agitation were improved in 89, 80 and 82% of participants, respectively.

In a single–blind, placebo–controlled, crossover study involving 12 healthy volunteers, two different single doses of valerian-hops (valerian 500 mg, hops 120 mg; valerian 1500 mg, hops 360 mg) were assessed for their effects on EEG recordings.\(^{(54)}\) Some slight effects on the quantitative EEG were documented following administration of higher dose valerian–hops, indicating effects on the central nervous system.

The effects of valerian extract 100 mg (no further details of preparation given) on activation and performance of 48 healthy volunteers under experimental social stress conditions have been assessed, with or without propranolol 20 mg, in a randomised double–blind, placebo–controlled study.\(^{(55)}\) Valerian was reported to have no influence on physiological activation and to lead to less intensive subjective feelings of somatic arousal.

The effects of a hydroalcoholic extract of valerian have been assessed in a randomised study involving 40 patients with minor symptoms of anxiety and emotional tension.\(^{(56)}\) Participants received valerian extract 100 mg three times daily, or placebo, for 21 days. It was reported that valerian was superior to placebo.

Several other studies have assessed the effects of combinations of valerian and St. John’s wort (Hypericum perforatum) in patients with anxiety or depression.\(^{(57–59)}\) In a randomised, double–blind study involving 100 patients with anxiety, a combination of valerian and St. John’s wort was reported to be significantly more effective than diazepam according to a physician’s rating scale and a patient’s self–rating scale.\(^{(57)}\) In a randomised, double–blind trial involving 162 patients with dysthymic disorders, the effects of a valerian and St. John’s wort combination (Sedariston) were compared with those of amitriptyline 75–150 mg.\(^{(58)}\) Another randomised, double–blind trial,
involving 100 patients with mild–to–moderate depression compared Sedariston with desipramine 100–150 mg.\textsuperscript{(59)} Pooling the results of these two studies indicated there were 88 (68%) treatment responders in the Sedariston group and 66 (50%) in the group that received standard antidepressants.\textsuperscript{(60)} This difference was not statistically significant.

Several studies have assessed the effects of valerian, or herbal combination products containing valerian, on performance the morning after treatment (see Side–effects, Toxicity).\textsuperscript{(52,61)}
Side-effects, Toxicity

Data relating to the safety of valerian have been reviewed.\(^{(G21)}\)

Studies assessing the effects of valerian on measures of performance suggest that there may be slight impairment for a few hours following valerian ingestion. However, studies have shown that ‘hangover’ effects (impairment of performance the morning following valerian treatment) do not appear to be a concern.

In a randomised, double-blind trial involving 102 healthy volunteers, the effects of single-dose valerian extract (Sedonium) 600 mg on reaction time, alertness and concentration were compared with those of flunitrazepam 1 mg and placebo.\(^{(61)}\) The treatment was administered in the evening and psychometric tests were carried out the next morning. After a one-week wash-out period, 91 volunteers continued with the second phase of the study, which comprised 14 days’ administration of valerian extract 600 mg or placebo. Single-dose valerian extract administration did not impair reaction time, concentration or co-ordination. A ‘hangover’ effect was reported by 59% of flunitrazepam recipients, compared with 32% and 30% of placebo and valerian recipients, respectively \((p < 0.05)\). At the end of the 14-day study, there was no statistically significant difference \((p = 0.45)\) between valerian extract and placebo on mean reaction time (a measure of performance), and valerian recipients showed a trend towards improved sleep quality.

A randomised, double-blind study involving 80 volunteers assessed the ‘hangover’ effects of tablets containing valerian and hops, and a syrup containing valerian only, given as a single dose, against both placebo and active control (flunitrazepam 1 mg).\(^{(52)}\) Performance the morning after treatment, measured both objectively and subjectively, was reported to be impaired only in the flunitrazepam group. Side effects occurred more frequently in the flunitrazepam group (50%), compared with the valerian and placebo groups (10%). A further battery of cognitive psychomotor tests was carried out in another study involving 36 volunteers who received either valerian syrup, valerian–hops tablets, or placebo; tests were conducted 1–2 hours after drug administration to assess acute effects. Compared with placebo, there was a slight, but statistically significant, impairment in vigilance with valerian syrup and impairment in the processing of complex information with valerian–hops tablets.\(^{(52)}\)

Few controlled clinical trials of valerian preparations have provided detailed information on safety. Where adverse event data were provided, randomised,
placebo–controlled trials involving healthy volunteers or patients with diagnosed insomnia reported that adverse events with valerian were mild and transient, and that the types and frequency of adverse events reported for valerian were similar to those for placebo.\(^{46,61}\) One study involving small numbers of patients reported a lower frequency of adverse events with valerian than with placebo; the authors did not suggest an explanation for this.\(^{39}\) Studies comparing valerian preparations with benzodiazepines have reported that valerian root extract (LI-156) 600 mg daily for 14 days\(^{61}\) or 28 days\(^{41}\) had a more favourable adverse effect profile than flunitrazepam 1 mg daily for 14 days\(^{61}\) and oxazepam 10 mg daily for 28 days,\(^{41}\) respectively.

There is an isolated report of cardiac complications and delirium associated with valerian root extract withdrawal in a 58–year–old man with a history of coronary artery disease, hypertension and congestive heart failure.\(^{62}\) The man had been taking valerian root extract (530 mg to 2 g, five times daily). It was hypothesised that given the effects of valerian on GABA, withdrawal of valerian root might produce a benzodiazepine–like withdrawal syndrome. However, the man was taking multiple medications and had undergone surgery, and a causal link with valerian could not be made. There have also been isolated reports of hepatotoxic reactions following the use of combination products containing valerian, although these products contained other herbal ingredients, such as scullcap and chaparral, which could have been responsible.\(^{63,G18 G21}\) Several other reports document hepatotoxic reactions with single–ingredient valerian products, although it is possible that these were idiosyncratic reactions.\(^{64}\) There is a lack of data on the safety of the long–term use of valerian, and such studies are required.\(^{G21}\)

Cases of individuals who had taken overdoses of valerian or valerian-containing products have been documented. One case involved an 18–year–old female who ingested 40–50 capsules of powdered valerian root 470 mg, approximately 20 times therapeutic doses.\(^{65}\) The patient presented 3 hours after ingestion with fatigue, crampy abdominal pain, chest tightness, tremor and lightheadedness. Liver function tests were normal; a urine screen tested positive for tetrahydrocannabinol. The patient was treated with activated charcoal, and symptoms resolved within 24 hours. Several cases (\(n = 47\)) have been documented of overdose with a combination valerian-containing product (‘Sleep-Qik’; valerian dry extract 75 mg, hyoscine hydrobromide 0.25 mg, cyproheptadine hydrochloride 2 mg).\(^{66,67}\) Individuals had ingested tablets equivalent to 0.5–12 g valerian. Liver function tests were carried out for most patients, all of which were normal.

Toxicological studies documented in the older literature have reported an
LD$_{50}$ of 3.3 mg/kg for an ethanolic extract of valerian administered intraperitoneally in rats, and that daily doses of 400–600 mg/kg, administered intraperitoneally for 45 days, did not lead to any changes in weight, blood or urine measurements, compared with controls.\(^{(1)}\) Literature cited in a review of the safety of valerian describes an LD$_{50}$ of 64 mg/kg for valtrate, 125 mg/kg for didrovaltrate and 150 mg/kg for acevaltrate in mice after intraperitoneal injection.\(^{(G21)}\) Another study in mice reported that valerenic acid 150 mg/kg, given by intraperitoneal injection, caused muscle spasms and that 400 mg/kg caused heavy convulsions.\(^{(17)}\) The latter dose was lethal to six of seven mice.

*In vitro* cytotoxicity and mutagenicity have been documented for the valepotriates. The clinical significance of this is unclear, since the valepotriates are known to be highly unstable and, therefore, probably degrade when taken orally. Also, they are unlikely to be present in high concentrations in finished products. The volatile oil is unlikely to present any hazard in aromatherapy.\(^{(G58)}\)

The EMEA HMPWG proposed core SPC states that the total exposure to valepotriates should not exceed the maximum exposure with herbal tea.\(^{(G23)}\) Alkylating and cytotoxic properties of valepotriates are not relevant for finished products as valepotriates decompose rapidly and only traces of valepotriates or their degradation products (in part, baldrinals) are found.
Contra-indications, Warnings

The documented CNS-depressant activity of valerian may potentiate existing sedative therapy.

According to the EMEA HMPWG proposed core SPC, patients should seek medical advice if symptoms persist for more than two weeks, or worsen. Intake of valerian preparations immediately (up to 2 hours) before driving a car or operating machinery is not recommended. The effect of valerian preparations may be enhanced by consumption of alcohol.\(^{(G23)}\)

Pregnancy and lactation

The safety of valerian during pregnancy and lactation has not been established and should, therefore, be avoided.\(^{(68)}\)

A study in rats involved the administration of valepotriates (6, 12 and 24 mg/kg administered orally) during pregnancy up to the 19th day when animals were sacrificed.\(^{(69)}\) There were no differences between valepotriate-treated rats and control rats as determined by fetotoxicity and external examination studies, although the two highest doses of valepotriates were associated with an increase in retarded ossification evident on internal examination.

The EMEA HMPWG proposed core SPC states that as data on the use of valerian during pregnancy are not available, use is not recommended as a general precaution. No adverse effects have been reported from the common use of valerian root as a medicinal product, but experimental data are lacking.\(^{(G23)}\)
Pharmaceutical Comment

The traditional use of valerian as a mild sedative and hypnotic has been supported by actions documented in studies involving both animals and humans.\(^{(G62)}\)

The sedative activity of valerian has been attributed to both the volatile oil and iridoid valepotriate fractions, but it is still unclear whether other constituents in valerian represent the active components. The valepotriate compounds are highly unstable and, therefore, are unlikely to be present in significant concentrations in finished products and probably degrade when taken orally. In view of this, the clinical significance of both the sedative and cytotoxic/mutagenic activities of valepotriates documented \textit{in vitro} is unclear.

The acute toxicity of valerian is considered to be very low.\(^{(G21)}\) There are isolated reports of adverse effects, mainly hepatotoxic reactions, associated with the use of single-ingredient and combination valerian-containing products. However, causal relationships for these reports could not be established as the cases involved other factors which could have been responsible for the observed effects. Some studies have compared valerian with certain benzodiazepines; the data available appear to suggest that valerian may have a more favourable tolerability profile, particularly in view of its apparent lack of ‘hangover’ effects. The safety of valerian in comparison with benzodiazepines requires further investigation and documentation.
References


17. Hendriks H et al. Central nervous depressant activity of valerenic acid in...


Species (Family)

Verbena officinalis L. (Verbenaceae)
Synonym(s)

Verbena
Part(s) Used
Herb
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL$^{(G37)}$
Constituents
See General References G2 G22 G40 G64.

**Glycosides**
Iridoid glycosides: hastatoside, verbenalin (verbanaloside), verbenin (aucubin). Phenyl propanoid glycosides: acetoside (verbascoside) and eukovoside.\(^{(1,2)}\)

**Volatile oils**
Monoterpene components include citral, geraniol, limonene and verbenone.

**Other constituents**
Adenosibe, alkaloid (unspecified), bitters, carbohydrates (stachyose, mucilage), β-carotene, invertin (sucrose hydrolytic enzymes), saponin and tannic acid.
Vervain is listed by the Council of Europe as a natural source of food flavouring (category N2). This category indicates that vervain can be added to foodstuffs in small quantities, with a possible limitation of an active principle (as yet unspecified) in the final product. In the USA, vervain is listed by the Food and Drugs Administration (FDA) as a Herb of Undefined Safety.
Herbal Use

Vervain is stated to possess sedative, thymoleptic, antispasmodic, mild diaphoretic and, reputedly, galactogogue properties. Traditionally, it has been used for depression, melancholia, hysteria, generalised seizures, cholecystalgia, jaundice, early stages of fever, and specifically for depression and debility of convalescence after fevers, especially influenza. (G2 G7 G64)
**Dosage**

*Dried herb*
2–4 g or by infusion three times daily.\(^{(G7)}\)

*Liquid extract*
2–4 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)

*Tincture*
5–10 mL (1 : 1 in 40% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

In vitro and animal studies

Galactogogue properties have been documented for vervain and attributed to aucubin.\(^3\) A luteinising action has also been reported, and attributed to inhibition of the gonadotrophic action of the posterior lobe of the pituitary gland.\(^3\) Extracts of vervain fruit have been used to treat dysmenorrhoea and to stimulate lactation.\(^3\) Vervain has been documented to possess weak parasympathetic properties, causing slight contraction of the uterus.\(^3\) Verbenalin has been reported to exhibit uterine stimulant activity.\(^{G30}\) Sympathetic activity has also been documented: in small doses verbenin has been reported to act as an agonist at sympathetic nerve endings, whereas larger doses result in antagonism.\(^{G22}\) Verbascoside reportedly acts as an agonist to the antitremor action of levodopa, and as an antihypertensive and analgesic.\(^3\) A slight laxative action in mice has been documented for iridoid glycosides.\(^4\)
Side-effects, Toxicity

None documented for vervain. High doses of verbenalin are stated to paralyse the CNS, resulting in stupor and convulsions. \(^{(G22)}\)
Contra-indications, Warnings

None documented. Excessive doses of vervain may interfere with existing hypo- or hypertensive and hormone therapies.

**Pregnancy and lactation**

Vervain is reputed to act as an abortifacient and oxytocic agent\(^{(G30)}\) with *in vivo* utero–activity documented (*see In vitro* and animal studies). In view of this, vervain should not be taken during pregnancy.

Vervain may affect lactation in view of the reported galactogogue properties.\(^{(3)}\)
Pharmaceutical Comment

Limited chemical, pharmacological and toxicity data are available for vervain. Documented scientific information does not justify the herbal uses, although galactogogue properties have been reported. No human data were located. In view of the lack of toxicity data and documented pharmacological actions in animals, excessive use of vervain should be avoided.
References

See also General References G2 G9 G16 G22 G30 G31 G36 G37 G40 G64.


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Species (Family)

*Daucus carota* L. subsp. *carota* (Umbelliferae)
Synonym(s)

Daucus, Queen Anne’s Lace
Part(s) Used

Herb
Pharmacopoeial and Other Monographs

BHC 1992$^{(G6)}$

BHP 1996$^{(G9)}$

Martindale 33rd edition$^{(G67)}$

PDR for Herbal Medicines 2nd edition$^{(G36)}$
Legal Category (Licensed Products)

GSL($^{G37}$)
Constituents

See General References G6 G41 G64.

Documented constituents refer to the fruit or seeds obtained from the dried fruit unless stated.

**Flavonoids**
Flavones (e.g. apigenin, chrysin, luteolin), flavonols (e.g. kaempferol, quercetin) and various glycosides.\(^{(1)}\)

**Furanocoumarin**
8-Methoxypsoralen and 5-methoxypsoralen (0.01–0.02 μg/g fresh weight) in fresh plant. Concentrations increased in the diseased plant.\(^{(2)}\)

**Volatile oils**
0.66–1.65%.\(^{(3)}\) Many components identified; relative composition varies between different cultivars.\(^{(3)}\) Various components include α-pinene, β-pinene, geraniol, geranyl acetate, limonene, α-terpinen, p-terpinen, α-terpineol, terpinen–4–ol, p-decanolactone (monoterpenes); β-bisabolene, β-elemene, caryophyllene, caryophyllene oxide, carotol, daucol (sesquiterpenes); asarone (phenylpropanoid derivative).\(^{(3)}\)

**Other constituents**
Choline,\(^{(4)}\) daucine (alkaloid), a tertiary base (uncharacterised),\(^{(5)}\) fatty acids (butyric, palmitic), coumarin, xylitol (polyol).
Food Use

Wild carrot should not be confused with the common cultivated carrot, *D. carota* L. subsp. *sativus* (Hoffm.), which has the familiar fleshy orange–red edible root. Wild carrot has an inedible tough whitish root. Wild carrot is listed by the Council of Europe as a natural source of food flavouring (category N1, N3). Category N1 indicates that for the roots there are no restrictions on use, whereas category N3 indicates that there is insufficient information available for an adequate assessment of potential toxicity.
Herbal Use

Wild carrot is stated to possess diuretic, antilithic, and carminative properties. Traditionally, it has been used for urinary calculus, lithuria, cystitis, gout, and specifically for urinary gravel or calculus. (G6 G7 G8 G64)
Dosage

*Dried herb*
2–4 g or by infusion three times daily.\(^{G6 G7}\)

*Liquid extract*
2–4 mL (1 : 1 in 25% alcohol) three times daily.\(^{G6 G7}\)
Pharmacological Actions

In vitro and animal studies

Significant antifertility activity (60%) in rats has been reported for wild carrot.\(^{(6)}\) In contrast, insignificant antifertility activity was observed in pregnant rats fed oral doses of up to 4.5 g/kg body weight from day 1 to day 10 of pregnancy.\(^{(7)}\) Aqueous, alcoholic and petrol extracts were reported to exhibit 20%, 40% and 10% activities respectively. Weak oestrogenic activity\(^{(6,8,9)}\) and inhibition of implantation\(^{(6,9)}\) has been documented for seed extracts.\(^{(8)}\) Oestrogenic activity, demonstrated by the inhibition of ovarian hypertrophy in hemicastrated rats, has been attributed to the known constituent coumarin (a weak phytooestrogen).\(^{(10)}\)

Central effects similar to those of barbiturates have been documented for the seed oil obtained from \emph{D. carota} var. \emph{sativa}.\(^{(11)}\) The oil was reported to elicit CNS hypnotic effects in the rat, hypotension in the dog\(^{(4)}\) leading to respiratory depression at higher doses, anticonvulsant activity in the frog, \emph{in vitro} smooth muscle relaxant activity reducing acetylcholine–induced contractions (ileum/uterus, rabbit/rat), antagonism of acetylcholine in isolated frog skeletal muscle, direct depressant effect on cardiac muscle in the dog.\(^{(4,11)}\) \emph{In vitro} cardiotonic activity\(^{(4)}\) and vasodilation of coronary vessels of the isolated cat heart has been reported.\(^{(12)}\) Papaverine–like antispasmodic activity has been documented for a tertiary base isolated from wild carrot seeds.\(^{(5)}\) Activity of approximately one–tenth that of papaverine was noted in a number of isolated preparations: ileum, uterus, blood vessels and trachea.\(^{(5)}\) Cholinergic–type actions have also been reported for wild carrot with \emph{in vitro} spasmodic actions noted in both smooth and skeletal muscle.\(^{(4)}\) This cholinergic activity has been attributed to choline.\(^{(13)}\) The identity of a second quaternary base isolated was not established.

Terpinen–4–ol is a documented component of the seed oil. This constituent is considered to be the diuretic principle in juniper, exerting its effect by causing renal irritation (see Juniper).

Increased resistance to carbon tetrachloride–induced hepatotoxicity has been reported in rats fed wild carrot.\(^{(14)}\)

Limited antifungal activity has been documented, with activity exhibited against only one (\emph{Botrytis cinerea}) out of nine fungi tested.\(^{(15)}\)

Agglutination of \emph{Streptococcus mutans} cells has been described for wild carrot. The agglutinin, found to be heat and trypsin stable but sensitive to
dextranose, was thought to be a dextran.\textsuperscript{(16)}
Side-effects, Toxicity

The oil is reported to be non-toxic.\(^{(G41 G58)}\) Acute LD\(_{50}\) values in mice (oral) and guinea-pigs (dermal) are reported to exceed 5 g/kg.\(^{(17)}\)

The oil contains terpinen-4-ol, which is the component associated with the renal irritancy of juniper oil.

The oil is reported to be generally non-irritating and non-sensitising.\(^{(12)}\) However, hypersensitivity reactions, occupational dermatitis and positive patch tests have been reported for wild carrot.\(^{(2, G51)}\) Wild carrot is reported to have a slight photosensitising effect.\(^{(2)}\) Furanocoumarins are known photosensitisers.
Contra-indications, Warnings

Fruit extracts may cause sensitivity reactions similar to those seen with celery.\(^{(2)}\) Excessive doses of the oil may cause renal irritation in view of the terpinen-4-ol content (see Juniper). Excessive doses may affect existing hypo- and hypertensive, cardiac and hormone therapies.

**Pregnancy and lactation**
The safety of wild carrot has not been established. Both spasmodic and spasmolytic actions on smooth muscle *in vitro* have been reported. In view of this, the documented mild oestrogenic activity and potentially irritant volatile oil, excessive doses of wild carrot during pregnancy and lactation should be avoided.
Pharmaceutical Comment

Phytochemical studies documented for wild carrot concentrate on the composition of the volatile oil obtained from both the fresh and dried fruits (seeds). The composition of the oil varies between different cultivars. Animal studies have documented a variety of pharmacological actions including CNS-depressant, spasmodic and antispasmodic, hypotensive and cardiac-depressant activities. However, the majority of these actions were observed in *in vitro* preparations. The principal traditional use of wild carrot is as a diuretic. This activity has not been documented in animal studies, but the seed oil of wild carrot does contain terpinen-4-ol, the diuretic principle documented for juniper. Toxicity data only refer to the oil and indicate low toxicity. However, in view of the documented mild oestrogenic activity and potential for internal irritation by the oil, excessive ingestion should be avoided.
References

See also General References G6 G9 G16 G31 G36 G37 G41 G43 G51 G58 G64.

16. Ramstorp M et al. Isolation and partial characterization of a substance

Wild Lettuce
Species (Family)

*Lactuca virosa* L. (Asteraceae/Compositae)
Synonym(s)

Bitter Lettuce, Lettuce Opium

Related *Lactuca* species include *Lactuca sativa* (Garden Lettuce), *Lactuca scariola* (Prickly Lettuce), *Lactuca altissima* and *Lactuca canadensis* (Wild Lettuce of America)
Part(s) Used

Leaf, latex
Pharmacopoeial and Other Monographs

BHC 1992\(^{G6}\)

BHP 1996\(^{G9}\)

Martindale 33rd edition\(^{G67}\)

PDR for Herbal Medicines 2nd edition\(^{G36}\)
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents

See General References G6 G22 G48 G60 G64.

All parts of the plant contain a milky, white latex (sap) which, when collected and dried, forms the drug known as lactucarium.\(^{(G33)}\)

**Acids**
Citric, malic and oxalic (up to 1%) acids; cichoric acid (phenolic).\(^{(1)}\)

**Alkaloids**
Hyoscyamine, later disputed.\(^{(2,G33)}\) \(N\)-methyl-\(\beta\)-phenethylamine, also disputed.\(^{(2)}\)

**Coumarins**
Aesculin, cichoriin.\(^{(1)}\)

**Flavonoids**
Flavones (e.g. apigenin, luteolin), flavonols (e.g. quercetin) and their glycosides.\(^{(1)}\)

**Terpenoids**
Bitter principles including the sesquiterpene lactones lactucin and lactupicrin (lactucopicrin); \(\beta\)-amyrin, germanicol, and lactucone (lactucerin). Lactucone is a mixture of \(\alpha\)- and \(\beta\)-lactucerol acetates, \(\beta\)-lactucerol being identical to taraxasterol.

**Other constituents**
Mannitol, proteins, resins and sugars.
Wild lettuce is not used in foods, although the related species *L. sativa* is commonly used as a salad ingredient.
Herbal Use

Wild lettuce is stated to possess mild sedative, anodyne and hypnotic properties. Traditionally, it has been used for insomnia, restlessness and excitability in children, pertussis, irritable cough, priapism, dysmenorrhoea, nymphomania, muscular or articular pains, and specifically for irritable cough and insomnia."
Dosage

**Dried leaves**
0.5–3.0 g or by infusion three times daily.\(^{(G6)}\)

**Liquid extract**
0.5–3.0 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G6)}\)

**Lactucarium (dried latex extract)**
(BPC 1934) 0.3–1.0 g three times daily.

**Soft extract**
(BPC 1934) 0.3–1.0 g three times daily.
Pharmacological Actions

In vitro and animal studies

Lactucarium has been noted to induce mydriasis.\(^{(G6)}\) This effect may be attributable to hyoscyamine, although the dried sap is reportedly devoid of this alkaloid.

An alcoholic extract of a related species, \(L.\ sativa\), has exhibited a sedative effect in toads, causing a reduction in motor activity and behaviour.\(^{(3)}\) Higher doses resulted in flaccid paralysis. In addition, an antispasmodic action on isolated smooth and striated muscle, and \textit{in vitro} negative chronotropic and inotropic effects on normal and stressed (tachycardic) hearts were observed. The antispasmodic action was noted to be antagonised by calcium.

Lactucin, lactupicrin and hyoscyamine have all been proposed as the sedative components in wild lettuce. However in the above study,\(^{(3)}\) the active component was uncharacterised and acted mainly peripherally, not readily crossing the blood–brain barrier. The suggested mode of action was via interference with basic excitatory processes common to neural and muscular functions, and not via a neuromuscular block.

Low amounts (nanograms) of morphine have been detected in \textit{Lactuca} species, although the concentrations involved are considered too low to exert any obvious pharmacological effect.\(^{(G60)}\)
Side-effects, Toxicity

None documented for *L. virosa*. Wild lettuce contains sesquiterpene lactones which are potentially allergenic.\(^{(G19)}\) Occupational dermatitis has been documented for *L. sativa* together with an urticarial eruption after ingestion of the leaves.\(^{(4-6,G51)}\) The milky sap of *L. sativa* is reported to be irritant.\(^{(G51)}\)

The toxicity of wild lettuce is stated to be low.

Consumption of large amounts of *L. scariola* has caused poisoning in cattle, who developed pulmonary emphysema, severe dyspnoea, and weakness.\(^{(7)}\) Only the immature plants were reported to be toxic.

*L. sativa* has been reported to produce only negative responses when tested for mutagenicity using the Ames test (*Salmonella typhimurium* TA98, TA100).\(^{(8)}\)
Contra–indications, Warnings

Overdosage may produce poisoning\(^{(G42)}\) involving stupor, depressed respiration, coma and even death. Wild lettuce may cause an allergic reaction in sensitive individuals, in particular those with an existing sensitivity to other members of the Asteraceae/Compositae family.

**Pregnancy and lactation**

The safety of wild lettuce has not been established. In view of the lack of toxicity data and the possibility of allergic reactions, excessive use of wild lettuce during pregnancy and lactation should be avoided.
Pharmaceutical Comment

The chemistry of wild lettuce is well documented although it is not clear which constituents represent the active components. Early reports of hyoscyamine as a constituent have not been substantiated by subsequent study. No published information was found to support the traditional herbal uses of wild lettuce, although a sedative action in toads has been reported for a related species \textit{L. sativa}. In view of the potential allergenicity of wild lettuce and the lack of toxicity data, excessive use should be avoided.
References

See also General References G6 G9 G10 G19 G22 G31 G33 G36 G37 G42 G43 G48 G51 G60 G64.


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Willow
Species (Family)

Salix species including *Salix alba* L., *Salix fragilis* L., *Salix pentandra* L., *Salix purpurea* L. (Salicaceae)
Synonym(s)

Salix
Part(s) Used

Bark
Pharmacopoeial and Other Monographs

American Herbal Pharmacopoeia\(^{(G1)}\)

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

BP 2002\(^{(G71)}\)

Complete German Commission E\(^{(G3)}\)

ESCOP 1997\(^{(G52)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)

Ph Eur 2004\(^{(G72)}\)
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents
See General References G1 G2 G6 G49 G52 G62 G64.

**Glycosides (phenolic)**
Various phenolic glycosides including salicin, salicortin, tremulacin, salireposide, picein and triandrin.\(^1\) Acetylated salicin, salicortin, salireposide, and esters of salicylic acid and salicyl alcohol may also occur.

**Salicylates (calculated as salicin)**
Vary between species, e.g. 0.5% in *S. alba*, 1–10% in *S. fragilis*, 3–9% in *S. purpurea*.\(^2\)

**Flavonoids**
Flavanones, eriodictoyl–7–glucoside; naringenin–5–glucoside; chalcone; isosalipurposide; catechin.\(^2,G52\)

**Tannins**
Condensed.

**Other constituents**
Catechins.

There is reported to be no difference between the phenolic glycoside pattern of the bark and leaf. The latter is also reported to contain flavonoids, catechins and condensed tannins.\(^{2,3}\)
Food Use

Willow is not used in foods.
Willow is stated to possess anti-inflammatory, antirheumatic, antipyretic, antihidrotic, analgesic, antiseptic and astringent properties. Traditionally it has been used for muscular and arthrodial rheumatism with inflammation and pain, influenza, respiratory catarrh, gouty arthritis, ankylosing spondylitis, and specifically for rheumatoid arthritis and other systemic connective tissue disorders characterised by inflammatory changes. The German Commission E approved internal use for diseases accompanied by fever, rheumatic ailments and headaches. \(^{(G3)}\)
Dosage

Dry bark
1–3 g or by decoction three times daily\(^{(G6 \ G7)}\) corresponding to 60–120 mg total salicin daily.\(^{(G3)}\)

Liquid extract
1–3 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G6 \ G7)}\)
Pharmacological Actions

**In vitro and animal studies**

Pharmacological actions documented for salicylates include anti-inflammatory, antipyretic, hyperglycaemic/hypoglycaemic and uricosuric/antiuricosuric activities, and increased blood–clotting time and plasma albumin binding.\(^{(G46)}\) Anti-inflammatory activity for salicin and tremulacin (isolated from *Populus* spp.) has been assessed in the hen’s egg choriollantoic test.\(^{\left(4, G52\right)}\) The results indicate that the activity may be due to the metabolites of these compounds.\(^{\left(4\right)}\) Salicin is probably the most active anti-inflammatory compound in willow; it is metabolised to salicylic acid.\(^{(5)}\) The enzymatic degradation of salicin, salicortin and tremulacin by β-glucosidase and by esterase has been investigated.\(^{(6)}\)

Tannins are known to have astringent properties.

**Clinical studies**

Willow bark extract (equivalent to 240 mg salicin/day) was compared with placebo in a two–week, randomised, double–blind, controlled trial involving 78 patients with osteoarthritis.\(^{(7)}\) A difference in pain dimension in the treated group, compared with placebo, just reached statistical significance (\(p = 0.047\)). It was concluded that willow bark extract had a moderate analgesic effect in osteoarthritis, and that it was well tolerated.

The pharmacological actions of salicylates in humans are well documented, and are applicable to willow. Salicin is a prodrug which is metabolised to saligenin in the gastrointestinal tract and to salicylic acid after absorption.\(^{(2)}\)
Side-effects, Toxicity

Side-effects and signs of toxicity normally associated with salicylates, such as gastric and renal irritation, hypersensitivity, blood in the stools, tinnitus, nausea and vomiting, may occur. Salicin is documented to cause skin rashes.\(^{(G44)}\)
Contra-indications, Warnings

Minor adverse effects including stomach ache, nausea, dizziness, sweating and rash have been reported in a small percentage of individuals.\(^{G52}\) Precautions associated with salicylate therapy are also applicable to willow. Therefore individuals with known hypersensitivity to aspirin, asthma, active peptic ulceration, diabetes, gout, haemophilia, hypo prothrombinaemia, kidney or liver disease should be aware of the possible risks associated with the ingestion of willow.\(^{8,G46}\) Irritant effects of salicylates on the gastrointestinal tract may be enhanced by alcohol, and barbiturates and oral sedatives have been documented to enhance salicylate toxicity as well as masking the symptoms of overdosage.\(^{G46}\) Concurrent administration of willow with other salicylate–containing products, such as aspirin, should be avoided. Drug interactions listed for salicylates are also applicable to willow and include oral anticoagulants, methotrexate, metoclopramide, phenytoin, probenecid, spironolactone and valproate.

Pregnancy and lactation

The safety of willow has not been established. Conflicting reports have been documented concerning the safety of aspirin taken during pregnancy. In view of this, the use of willow during pregnancy should be avoided. Salicylates excreted in breast milk have been reported to cause macular rashes in breastfed babies.\(^{G46}\)
Pharmaceutical Comment

Willow is rich in phenolic constituents, such as flavonoids, tannins and salicylates. Pharmacological actions normally associated with salicylates are also applicable to willow which support most of the herbal uses, although no studies were located specifically for willow. In view of the lack of toxicity data on willow, the usual precautions taken with other salicylate-containing drugs are applicable. Products containing willow should preferably be standardised on their salicin content, in view of the considerable variation in salicylate concentrations between different *Salix* species.
References

See also General References G1 G2 G3 G5 G6 G9 G10 G31 G36 G37 G43 G49 G52 G54 G56 G62 G64.


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Witch Hazel
Species (Family)

*Hamamelis virginiana* L. (Hamamelidaceae)
Synonym(s)
Hamamelis, Witchazel
Part(s) Used

Bark, leaf
Pharmacopoeial and Other Monographs

BHP 1996\(^{(G9)}\)

BP 2002\(^{(G71)}\)

Complete German Commission E\(^{(G3)}\)

ESCOP 1997\(^{(G52)}\)

Martindale 33rd edition\(^{(G67)}\)

Mills and Bone\(^{(G50)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)

Ph Eur 2004\(^{(G72)}\)

USP26/NF21\(^{(G73)}\)
Legal Category (Licensed Products)

GSL$^{G37}$
Constituents
See Reference 1, and General References G2 G22 G41 G48 G50 G52 G64.

Flavonoids (leaf)
Flavonols (e.g. kaempferol, quercetin) and their glycosides including astragalin, quercitrin, afzelin and myricitrin.

Tannins, catechins
Pharmacopoeial standard, not less than 3%. Hamamelitannin (hydrolysable), lesser amounts of condensed tannins (bark). (+)-catechin, (+)-gallocatechin, (−)-epicatechin gallate, (−)-epigallocatechingallate, proanthocyanidin oligomers of cyanidin and delphinidin type.

Volatile oils
About 0.5%. Hexen–2–ol, hexenol, α- and β-ionones, eugenol, safrole and sesquiterpenes.

Other constituents
Fixed oil (about 0.6%), resin (hamamelin, hamamamelitannin), wax, saponins, choline, free gallic acid and free hamamelose.
Food Use

Witch hazel is listed by the Council of Europe as a natural source of food flavouring (category N3). This category indicates that there is insufficient information available for an adequate assessment of potential toxicity. (G16)
Witch hazel is stated to possess astringent, antihaemorrhagic and anti-inflammatory properties. Traditionally, it has been used for diarrhoea, mucous colitis, haemorrhoids, haematemesis, haemoptysis, and externally for external haemorrhoids, bruises and localised inflamed swellings. The German Commission E approved use for minor skin injuries, local inflammation of skin and mucous membranes, haemorrhoids and varicose veins.
Dosage

*Dried leaves*
2 g or by infusion three times daily.\(^{(G6)}\)

**Hamamelis Liquid Extract**
(BPC 1973) 2–4 mL (1 : 1 in 45% alcohol) three times daily.\(^{(G6)}\)

**Hamamelis Water**
(BPC 1973) for local application, undiluted or 1 : 3 dilution for external use.\(^{(G3)}\)

**Decoction**
5–10 g in 250 mL water for compresses.\(^{(G3)}\)
Pharmacological Actions

The pharmacological properties of witch hazel have been reviewed.\(^1\,\text{G}50\,\text{G}52\)

**In vitro and animal studies**

Witch hazel is known to possess astringent and haemostatic properties, which have been attributed to the tannin constituents. Vasoconstriction was reduced in the hindquarters of rabbits when arteries were perfused with aqueous or ethanolic extracts of witch hazel leaf. A 70% ethanolic extract of leaf (1 : 5, 200 mg/kg, administered orally) significantly inhibited the chronic phase of carrageenan–induced rat paw oedema over a period of 19 days, compared with control (\(p < 0.05\)).\(^2\) An aqueous ethanolic extract of witch hazel bark yielded a fraction rich in polymeric proanthocyanins after ultracentrifugation.\(^3\) This fraction was significantly active against herpes simplex virus type 1 (HSV-1). It also showed radical scavenging properties, inhibited \(\beta\)-glucosidase and human leukocyte elastase activity, and was active in the croton oil ear oedema test in mice. In other studies, \(3-O\)-galloyl–epicatechin-(4\(\beta\),8)-catechin, a catechin oligomer and hamamelitannin isolated from witch hazel bark had IC\(_{50}\) values of 6.6, 8.8 and 1.0 \(\mu\)mol/L, respectively, for inhibition of 5–lipoxygenase.\(^4\) The oligomer was active in the microsomal lyso-PAF:acetyl-CoA-acetyltransferase inhibition assay, with an IC\(_{50}\) value of 9.4 \(\mu\)mol/L, whereas hamamelitannin was inactive.\(^4\)

**Clinical studies**

**Haemorrhoids**

Uncontrolled studies have suggested that witch hazel bark distillate (5%) and a salve containing witch hazel bark may be effective in the treatment of haemorrhoids.\(^\text{G}50\) A double–blind, controlled trial involving 90 patients with haemorrhoids compared the effects of witch hazel bark salve with those of other salves. Witch hazel was reported to be superior in relief of symptoms.

**Dermatology**

In a study involving 30 volunteers who received topical applications of a hydroglycolic extract of witch hazel leaf, skin temperature was significantly reduced, compared with baseline values. This was interpreted as a possible vasoconstrictor effect of witch hazel.\(^\text{G}52\) The effects of an after–sun lotion containing 10% hamamelis distillate were explored in 30 healthy volunteers using a modified UV-B erythema test for inflammation.\(^5\) It was reported that erythema suppression ranged from 20% at 7 hours to 27% at 48 hours.

Witch hazel leaf extract incorporated into a cream formulation was applied
twice daily for two weeks to seven children suffering from dermatitis atopica of the feet (chilblains) and to five children with eczema. Improvements in these conditions were reported.\(^{\text{G52}}\)

In a two–week, randomised, double–blind trial, 72 patients with moderately severe eczema were treated with either a hamamelis distillate cream (5.35 g distillate with 0.64 g ketone/100 g), hydrocortisone cream 0.5%, or drug–free cream.\(^6\) All three treatments significantly reduced itching, erythema and scaling after one week. Hydrocortisone cream was more effective than hamamelis cream.

Several clinical studies of witch hazel in the treatment of eczema have been reviewed.\(^{\text{G50}}\) An uncontrolled study involving 37 patients treated with a witch hazel leaf cream twice daily for two weeks reported improvements in eczema and neurodermatitis. A double–blind, placebo–controlled trial of witch hazel salve (25% water distillate from leaf) involving 80 patients with toxic and degenerative eczema and 31 patients with endogenous eczema found that atopic dermatitis responded to the treatment, but that there was no significant effect on primary irritant contact dermatitis. An uncontrolled study involving 22 patients with atopic eczema who were treated with witch hazel (4 g leaf provided 25 mL distillate/100 g salve) applied to affected arms over a three–week period reported improvements symptoms, compared with baseline values.\(^{\text{G50}}\)
Side-effects, Toxicity

The volatile oil contains safrole, a known carcinogen (see Sassafras), but in amounts too small to cause concern. Stomach irritation may occur in susceptible patients after oral treatment. Four of 1032 patients tested reacted to an ointment containing 25% witch hazel extract, but two of these patients were sensitive to wool fat in the ointment base. (G50)
Contra-indications, Warnings

None documented for witch hazel. In view of the tannin constituents, excessive ingestion of witch hazel is not recommended.

Pregnancy and lactation
There are no known problems with the use of witch hazel during pregnancy, although excessive ingestion should be avoided in view of the tannin content.
Pharmaceutical Comment

Witch hazel is characterised by its tannin constituents and astringent properties. The documented herbal uses are related to these astringent properties. There is some evidence to indicate that witch hazel is effective in the treatment of haemorrhoids and venous tone, but its use in the treatment of eczema and dermatitis is more controversial.
References


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Species (Family)

Achillea millefolium L. (Asteraceae/Compositae)
Synonym(s)
Milfoil, Millefolium
Part(s) Used

Flowerhead
Pharmacopoeial and Other Monographs

BHC 1992\(^{(G6)}\)

BHP 1996\(^{(G9)}\)

BP 2002\(^{(G71)}\)

Complete German Commission E\(^{(G3)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)

Ph Eur 2004\(^{(G72)}\)
Legal Category (Licensed Products)

GSL\textsuperscript{(G37)}
Constituents
See General References G2 G6 G22 G41 G64.

**Acids**
Amino acids (e.g. alanine, aspartic acid, glutamic acid, histidine, leucine, lysine, proline, valine), fatty acids (e.g. linoleic, myristic, oleic, palmitic, stearic), and others including ascorbic acid, caffeic acid, folic acid, salicylic acid and succinic acid.

**Alkaloids/bases**
Betonicine and stachydrine (pyrrolidine), trigonelline (pyridine), betaine and choline (bases). Uncharacterised alkaloids include achiceine, achilleine (possible synonym for L-betonicine), which is stated to yield achilletine on alkaline hydrolysis, and moscatine/moschatine, stated to be an ill-defined glucoalkaloid.

**Flavonoids**
Predominantly flavone glycosides apigenin- and luteolin–7–glycosides, with lesser quantities of artemetin, casticin, 5–hydroxy–3,6,7,4-tetramethoxyflavone and isorhamnetin. Rutin (a flavonol glycoside).

**Tannins**
Condensed and hydrolysable, with glucose as the carbohydrate component of the latter

**Volatile oils**
Numerous identified components include borneol, bornyl acetate (trace), camphor, 1,8–cineole, eucalyptol, limonene, sabinene, terpinen–4–ol, terpineol and α-thujone (monoterpenes), caryophyllene (a sesquiterpene), achillicin, achillin, millefin and millefolide (sesquiterpene lactones), azulene and chamazulene (sesquiterpene lactone-derived) and isoartemisia ketone. The relative composition of the components varies greatly between Achillea species, especially the azulene content. Azulene has been reported as the major component. However, true yarrow (A. millefolium) is thought to be hexaploid and azulene-free, whereas closely related species, such as Achillea lanulosa Nutt. and Achillea collina Becker, are tetraploid and contain up to 50% azulene in their volatile oil. The tetraploid species may be supplied for A. millefolium. The azulenes are not present in the fresh herb: they are formed as artefacts during steam distillation of the oil, from unstable precursors called proazulenes (e.g. achillin and achillicin), via equally unstable azulene-carboxylic acid intermediates.
Other constituents
Unknown cyanogenetic compound,\(^{(13)}\) sugars including arabinose, galactose, dextrose, dulcitol, glucose, inositol, maltose, mannitol and sucrose.\(^{(1,2)}\)

The constituents of yarrow have been reviewed in detail.\(^{(5)}\)
Food Use

Yarrow is listed by the Council of Europe as a natural source of food flavouring (herb, flowers, essential oil and other preparations: category 4, with limits on camphor, eucalyptol and thujone) (see Appendix 23). In the USA, yarrow is only approved for use in alcoholic beverages, and the finished product must be thujone free.
Yarrow is stated to possess diaphoretic, antipyretic, hypotensive, astringent, diuretic and urinary anti septic properties. Traditionally, it has been used for bruises, swellings, strains, fevers, common cold, essential hypertension, amenorrhoea, dysentery, diarrhoea, and specifically for thrombotic conditions with hypertension, including cerebral and coronary thromboses.
Dosage

*Dried herb*
2–4 g or by infusion three times daily.\(^{(G6 G7)}\)

*Liquid extract*
2–4 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G6 G7)}\)

*Tincture*
2–4 mL (1 : 5 in 45% alcohol) three times daily.\(^{(G6 G7)}\)
Pharmacological Actions

Some activities documented for yarrow are associated with the azulene constituents, although it is now thought that azulene is absent from true yarrow (see Constituents). Presumably some of the documented pharmacological studies have used Achillea species other than A. millefolium.

In vitro and animal studies

Anti–inflammatory activity has been documented for an aqueous extract of yarrow using mouse\(^\text{(15)}\) and rat\(^\text{(16)}\) paw oedema models, with inflammation induced by yeast\(^\text{(15)}\) and various inflammatory substances,\(^\text{(16)}\) including histamine, carrageenan and prostaglandin. In mouse studies, the active fraction was reported as a series of protein–carbohydrate complexes. Topical anti–inflammatory activity in rabbits has also been documented for the aqueous extract.\(^\text{(15)}\) In general, anti–inflammatory properties are associated with azulenes (see Chamomile, German). Anti–inflammatory activity has been described for the azulene components documented for the volatile oil of yarrow.\(^\text{(5)}\)

A diuretic effect was also noted in mice administered an aqueous extract of yarrow,\(^\text{(15)}\) but only at a dose more than double that required for an anti–inflammatory effect.\(^\text{(15)}\) Terpinen–4–ol, the diuretic principle in juniper, has been reported as a component of yarrow volatile oil.

CNS-depressant activity has been documented for the volatile oil: a dose of 300 mg/kg decreased the spontaneous activity of mice and lowered the body temperature of rats. In addition, 300–600 mg/kg doses inhibited pentetrazole–induced convulsions and prolonged sleep induced by a barbiturate preparation.\(^\text{(17)}\)

Moderate antibacterial activity has been documented for an ethanolic extract of the herb against *Staphylococcus aureus*, *Bacillus subtilis*, *Mycobacterium smegmatis*, *Escherichia coli*, *Shigella sonnei* and *Shigella flexneri*.\(^\text{(18)}\)

Antimicrobial properties have been documented for the sesquiterpene lactone fraction.\(^\text{(5)}\)

Achilleine 0.5 g/kg by intravenous injection has been noted to decrease the blood clotting time in rabbits by 32%.\(^\text{(8)}\) The haemostatic action persisted for 45 minutes with no observable toxic effects.

Antispasmodic activity on the isolated rabbit intestine has been documented for a flavonoid–containing fraction of yarrow.\(^\text{(9)}\) Antispasmodic activity is generally associated with azulene constituents (see Chamomile, German).
Antipyretic and hypotensive actions have been reported for the basic fraction (alkaloid/base);\(^{(G41)}\) the sesquiterpene lactone fraction is stated to possess cytotoxic activities,\(^{(5)}\) although no further details were located. Tannins are known to possess astringent activity.
Side–effects, Toxicity

Allergic reactions to yarrow (e.g. dermatitis) have been documented, and positive patch tests have been produced in individuals sensitised to other plants.\(^{(5,G33,G51)}\) An instance of yarrow tea causing a generalised eruption in a sensitised individual was reported in 1929. The allergenic properties of some sesquiterpene lactones are well documented, although none of those present in yarrow are recognised sensitisers.\(^{(G51)}\) Yarrow has been suspected of being a photosensitiser, although extracts have been reported to lack phototoxicity and to be devoid of psoralens, compounds with known photosensitising properties.\(^{(G51)}\)

Yarrow is considered to be non–toxic. In mice LD\(_{50}\) values have been reported of up to 3.65 g/kg (by mouth), 3.1 g/kg (by intraperitoneal injection), and greater than or equal to 1 g/kg (by subcutaneous injection).\(^{(15,17)}\) In rats, an LD\(_{50}\) (subcutaneous injection) has been recorded as 16.86 g/kg, with corresponding LD\(_{0}\) and LD\(_{100}\) values reported as 12 and 20 g/kg, respectively.\(^{(16)}\) By comparison, an ED\(_{25}\) for anti–inflammatory activity has been estimated as about 0.43 g/kg.\(^{(16)}\)

Terpenoid–rich volatile oils often possess irritant properties. Terpinen–4–ol, documented as a component of yarrow volatile oil, is thought to represent the diuretic principal of juniper as a result of its irritant action on the kidneys (see Juniper).\(^{(12)}\) The known toxic principle thujone has been documented as a minor component of yarrow volatile oil, although concentrations present are probably too low to represent a risk to human health.

A single report of animal poisoning has been documented for yarrow in which a calf died following the ingestion of a single plant.\(^{(5)}\) No additional reports of animal toxicity were located.
Contra-indications, Warnings

Yarrow may cause an allergic reaction in sensitive individuals, especially those with an existing hypersensitivity to other members of the Asteraceae/Compositae.\(^{(19)}\) Individuals with such a known hypersensitivity should avoid drinking herbal teas containing yarrow.\(^{(G60)}\) Excessive doses may interfere with existing anticoagulant and hypo- and hypertensive therapies, and may have sedative and diuretic effects.

**Pregnancy and lactation**

Yarrow should not be taken during pregnancy. It is reputed to be an abortifacient and to affect the menstrual cycle,\(^{(G30)}\) and the volatile oil contains trace amounts (0.3%) of the abortifacient principle thujone. Excessive use should be avoided during lactation.
Pharmaceutical Comment

The chemistry of yarrow is well documented although there has been some disagreement over the major component in the volatile oil. Various pharmacological actions have been reported in animal studies which support many of the reputed herbal uses although no human data were located. Yarrow is considered to be relatively non–toxic although allergic reactions in susceptible individuals have been documented. The volatile oil is contra–indicated in pregnancy and yarrow should be used with caution in patients with epilepsy. (G58)
References


1. Ivanov Ch, Yankov L. Composition of Achillea millefolium. I. Preparation of the total extracts and composition of the part of the alcoholic extracts soluble in alcohol and water. God Vissh Khimikotekhnol Inst Sofia 1967; 14: 195–222.


Species (Family)

*Rumex crispus* L. (Polygonaceae)
Synonym(s)

Curled Dock
Part(s) Used

Root
Pharmacopoeial and Other Monographs

BHP 1983\(^{(G7)}\)

Martindale 33rd edition\(^{(G67)}\)

PDR for Herbal Medicines 2nd edition\(^{(G36)}\)
Legal Category (Licensed Products)

GSL\(^{(G37)}\)
Constituents

See General References G22 G48 G64.

**Anthraquinones**
2–4%. Chrysophanol, emodin, nepodin, physcion (aglycones).\(^{(1-3)}\)

**Tannins**
Catechol 5% (condensed–type).

**Other plants parts**
The plant constituents documented include oxalic acid, oxalates, chrysophanic acid, emodin, tannin, and a complex volatile oil (more than 60 components identified).\(^{(4, G51)}\)
Food Use

Yellow dock is not used in foods.
Herbal Use

Yellow dock is stated to possess gentle purgative and cholagogue properties. Traditionally, it has been used for chronic skin disease, obstructive jaundice, constipation, and specifically for psoriasis with constipation.\(^{(G7\ G64)}\)
Dosage

Dried root
2–4 g or by decoction three times daily.\(^{(G7)}\)

Liquid extract
2–4 mL (1 : 1 in 25% alcohol) three times daily.\(^{(G7)}\)

Tincture
1–2 mL (1 : 5 in 45% alcohol) three times daily.\(^{(G7)}\)
Pharmacological Actions

*In vitro and animal studies*

None documented for the root. Slight antibacterial activity has been reported for herb extracts, which exhibited activity towards both Gram–positive (*Staphylococcus aureus, Mycobacterium smegmatis*) and Gram–negative (*Escherichia coli, Shigella sonnei, Shigella flexneri*) organisms.\(^4\)
Side–effects, Toxicity

None documented for yellow dock. In view of the documented anthraquinone constituents, side–effects generally associated with laxatives are also applicable to yellow dock. Overuse may cause abdominal cramps and diarrhoea, and prolonged use may lead to intestinal atrophy and hypokalaemia.

Dermatitis has been reported in livestock following the ingestion of plant material in large quantities.\(^{(G51)}\) Oxalic acid is known to be a toxic plant acid that forms insoluble calcium salts which cause a disturbance in calcium concentrations and hence affect the blood coagulation mechanism.\(^{(G33)}\)
Contra-indications, Warnings

Warnings generally associated with stimulant laxatives are also applicable to yellow dock. Therefore, yellow dock should not be taken when there is existing intestinal obstruction, and excessive use should be avoided (see Side-effects, Toxicity).

Pregnancy and lactation
In general, unstandardised stimulant laxatives are not recommended for use during pregnancy. The use of yellow dock should therefore be avoided in favour of a standardised preparation that is recommended for the treatment of constipation during pregnancy. The use of yellow dock by breastfeeding women should also be avoided, since it has been documented that anthraquinones can be secreted into the breast milk (see Senna).
Limited chemical, pharmacological, and toxicity information is available for yellow dock. Documented anthraquinone constituents justify the reputed purgative action. Although the purgative effect of yellow dock is reputed to be gentle, the use of unstandardised anthraquinone–containing preparations should be avoided since their pharmacological effect is unpredictable and may cause abdominal cramp and diarrhoea.
References

See also General References G7 G22 G31 G32 G33 G36 G37 G48 G51 G64.

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