

European Medicines Agency Evaluation of Medicines for Human Use

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2	This document was valid from 07 September 2006 until September 2019. It is now superseded by a <u>new version</u> adopted by the HMPC on 25 September 2019 and published on the EMA website.			
	COMMUNITY HERBAL MONOGRAPH ON RHAMNUS FRA	NGULA L., CORTEX		
	DISCUSSION IN THE SAFETY AND EFFICACY DRAFTING GROUP / WORKING PARTY ON COMMUNITY	January 2006		
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Herbal medicinal products; HMPC; Community herbal monograph; wellestablished use; frangula bark; *Rhamnus frangula* L.

KEYWORDS

¹ Changes introduced in sections 4.9 and 5.1

COMMUNITY HERBAL MONOGRAPH ON RHAMNUS FRANGULA L., CORTEX

1. NAME OF THE MEDICINAL PRODUCT

To be specified for the individual finished product.

QUALITATIVE AND QUANTITATIVE COMPOSITION^{2, 3} 2.

Well-established use	Traditional use
With regard to the marketing authorisation application of Article 10(a) of Directive 2001/83/EC, as amended	With regard to the registration application of Article 16d(1) of Directive 2001/83/EC, as amended
Rhamnus frangula L. (Frangula alnus Miller), cortex (frangula bark)	
• Herbal substance dried, whole or fragmented bark of the stems and branches, standardised	
• Herbal preparation standardised herbal preparations thereof	
3. PHARMACEUTICAL FORM	
Well-established use	Traditional use

Standardised herbal substance or herbal	
preparation for oral use in solid or liquid dosage forms.	
The pharmaceutical form should be described by	
the European Pharmacopoeia full standard term.	

CLINICAL PARTICULARS 4.

Therapeutic indications 4.1.

Well-established use	Traditional use
Herbal medicinal product for short-term use in cases of occasional constipation.	

 ² The material complies with the Ph. Eur. monographs.
 ³ The declaration of the active substance(s) should be in accordance with relevant herbal quality guidance.

4.2. Posology and method of administration

Well-established use	Traditional use
wen-established use	
Posology	
The maximum daily dose of hydroxyanthracene	
glycosides is 30 mg. This is equivalent to(dose	
of the preparation).	
The correct individual dose is the smallest	
required to produce a comfortable soft-formed	
motion.	
Adolescents over 12 years of age, adults, elderly	
Herbal substance/preparation equivalent to $10 - 30$	
mg hydroxyanthracene derivatives, calculated as	
glucofrangulin A, to be taken once daily at night.	
Normally it is sufficient to take this medicinal	
product up to two to three times a week.	
Not recommended for use in children under 12	
years of age (see section 4.3 Contraindications).	
years of uge (see section 1.5 contraincreations).	
The pharmaceutical form must allow lower	
dosages.	
-	
Method of administration	
As described in the package leaflet corresponding	
to the pharmaceutical form.	
Duration of use	
Use for more than 1 - 2 weeks requires medical	
supervision.	
If the symptoms persist during the use of the	
medicinal product, a doctor or a pharmacist should	r i i i i i i i i i i i i i i i i i i i
be consulted.	
See also section 4.4 Special warnings and	
precautions for use.	
4.3. Contraindications	

4.3. Contraindications

Well-established use	Traditional use
Known hypersensitivity to the active substance.	
Cases of intestinal obstructions and stenosis, atony, appendicitis, inflammatory colon diseases	
(e.g. Crohn's disease, ulcerative colitis), abdominal pain of unknown origin, severe	
dehydration state with water and electrolyte depletion.	
Children under 12 years of age.	

4.4. Special warnings and precautions for use

Well-established use

Patients taking cardiac glycosides, antiarrhythmic medicinal products, medicinal products inducing QT-prolongation, diuretics, adrenocorticosteroids or liquorice root, have to consult a doctor before taking frangula bark concomitantly.

Like all laxatives, frangula bark should not be taken by patients suffering from faecal impaction and undiagnosed, acute or persistent gastrointestinal complaints, e.g. abdominal pain, nausea and vomiting unless advised by a doctor because these symptoms can be signs of potential or existing intestinal blockage (ileus).

If laxatives are needed every day the cause of the constipation should be investigated. Long-term use of laxatives should be avoided.

If stimulant laxatives are taken for longer than a brief period of treatment, this may lead to impaired function of the intestine and dependence on laxatives. Frangula bark preparation should only be used if a therapeutic effect cannot be achieved by a change of diet or the administration of bulk forming agents.

When frangula bark preparations are administered to incontinent adults, pads should be changed more frequently to prevent extended skin contact with faeces.

Patients with kidney disorders should be aware of possible electrolyte imbalance.

4.5. Interactions with other medicinal products and other forms of interaction

Well-established use	Traditional use
Hypokalaemia (resulting from long-term laxative	
abuse) potentiates the action of cardiac glycosides	
and interacts with antiarrhythmic medicinal	
products, with medicinal products, which induce	
reversion to sinus rhythm (e.g. quinidine) and with	
medicinal products inducing QT-prolongation.	
Concomitant use with other medicinal products	
inducing hypokalaemia (e.g. diuretics,	
adrenocorticosteroids and liquorice root) may	
enhance electrolyte imbalance.	

Traditional use

4.6. Pregnancy and lactation

Well-established use	Traditional use
Pregnancy	
There are no reports of undesirable or damaging	
effects during pregnancy and on the foetus when	
used at the recommended dosage.	
However, as a consequence of experimental data	
concerning a genotoxic risk of several anthranoids,	
e.g. emodin, frangulin, chrysophanol and	
physcion, use is not recommended during	
pregnancy.	
Lactation	
Use during breastfeeding is not recommended as	
there are insufficient data on the excretion of	
metabolites in breast milk.	
After administration of other anthranoids, active	
metabolites, such as rhein, are excreted in breast	
milk in small amounts. A laxative effect in breast	
fed babies has not been reported.	
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4.7. Effects on ability to drive and use machines	
Well-established use	Traditional use
Not relevant.	
Not relevant.	
Not relevant. 4.8. Undesirable effects	
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	<u>Traditional use</u>
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4.9. Overdose

Well-established use	Traditional use
The major symptome of everlage/abuge are	
The major symptoms of overdose/abuse are griping pain and severe diarrhoea with consequent	
losses of fluid and electrolytes, which should be	
replaced. Diarrhoea may cause potassium	
depletion, in particular. Potassium depletion may	
lead to cardiac disorders and muscular asthenia,	
particularly where cardiac glycosides, diuretics,	
adrenocorticosteroids or liquorice root are being	
taken at the same time. Treatment should be	
supportive with generous amounts of fluid.	
Electrolytes, especially potassium, should be	
monitored. This is especially important in the	
elderly.	
Chronic ingested overdoses of anthranoid	
containing medicinal products may lead to toxic	
hepatitis.	
5. PHARMACOLOGICAL PROPERTIES	
5. THARMACOLOGICAL TROTERTIES	
5.1. Pharmacodynamic properties	
citi i nur nucouynume properties	
Well-established use	
wen-established use	Traditional use:
Pharmaco-therapeutic group: contact laxatives	Not required as per Article 16c(1)(a)(iii) of
Pharmaco-therapeutic group: contact laxatives ATC-code: A 06 AB	Not required as per Article 16c(1)(a)(iii) of
Pharmaco-therapeutic group: contact laxatives ATC-code: A 06 AB 1,8-dihydroxyanthracene derivatives possess a	Not required as per Article 16c(1)(a)(iii) of
Pharmaco-therapeutic group: contact laxatives ATC-code: A 06 AB 1,8-dihydroxyanthracene derivatives possess a laxative effect.	Not required as per Article 16c(1)(a)(iii) of
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Pharmaco-therapeutic group: contact laxatives ATC-code: A 06 AB 1,8-dihydroxyanthracene derivatives possess a laxative effect. Glucofrangulins and frangulins are respectively 0- diglycosides and 0-monoglycosides, which are	Not required as per Article 16c(1)(a)(iii) of
Pharmaco-therapeutic group: contact laxatives ATC-code: A 06 AB 1,8-dihydroxyanthracene derivatives possess a laxative effect. Glucofrangulins and frangulins are respectively 0- diglycosides and 0-monoglycosides, which are largely (all β-0-glycosides) not split by human	Not required as per Article 16c(1)(a)(iii) of
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2. influence on secretion processes by two concomitant mechanisms viz. inhibition of absorption of water and electrolytes (Na⁺, Cl⁻) into the colonic epithelial cells (antiabsorptive effect) and increase of the leakiness of the tight junctions and stimulation of secretion of water and electrolytes into the lumen of the colon (secretagogue effect) resulting in enhanced concentrations of fluid and electrolytes in the lumen of the colon. The motility effects are mediated by direct stimulation of colonic neurons and possibly by prostaglandins.

Defaecation takes place after a delay of 8 - 12 hours due to the time taken for transport to the colon and metabolisation into the active compound.

5.2 Pharmacokinetic properties

Well-established use	Traditional use
The β-0-linked glycosides are not split by human digestive enzymes and therefore not absorbed in the upper gut to a large extent. They are converted by the bacteria of the large intestine into the active metabolite (emodin-9-anthrone). Mainly anthraquinone aglycones are absorbed and transformed into their corresponding glucuronides and sulphate derivatives. After oral administration of frangula bark extract, rhein, emodin and traces of chrysophanol are found in human urine. After administration of other anthranoids, active metabolites, such as rhein, pass in small amounts into breast milk. Animal experiments demonstrated that placental-passage of rhein is low.	Directive 2001/83/EC as amended.

5.3 Preclinical safety data

Well-established use	Traditional use
There are no studies on single dose toxicity, on repeated dose toxicity, on reproductive toxicity or	Not required as per Article 16c(1)(a)(iii) of Directive 2001/83/EC as amended, unless
on carcinogenicity.	necessary for the safe use of the product.
Experimental data, mainly in vitro tests showed a	
genotoxic risk of several anthranoids in the	
Salmonella microsome assay, emodin,	
chrysophanol and physcion were weakly	
mutagenic. No mutagenic effects were observed in	
the V79-HGPRT mutation assay and in the	
unscheduled DNA synthesis (UDS) assay for	
chrysophanol and physcion. Emodin was highly	
mutagenic in the V79-HGPRT mutation assay. In	
the UDS assay emodin was a string inducer of	

UDS in primary hepatocytes. Emodin was also tested with respect to its transforming activity in C3H/M2 mouse fibroblasts *in vitro*. In the *in vitro* salmonella/microsome mutagen test and the deoxyribonucleic acid (DNA) repair test of primary rat hepatocytes emodin and frangulin, an alcoholic extract of "Rhamnus frangula", and a commercial frangula bark preparation showed a dose-dependent increase in the mutation rate or the induction of DNA repair.

However, *in vivo* studies of other anthranoidcontaining herbal substance (senna) in rat hepatocytes (chromosome aberration test, mouse spot test, *in vivo/in vitro* UDS (unscheduled DNA synthesis) showed no evidence of any genetic effects.

Further 2-year studies on male and female rats and mice with emodin gave no evidence of carcinogenic activity for male rats and female mice, and equivocal evidence for female rats and male mice.

Laxative use as a risk factor in colorectal cancer (CRC) was investigated in some clinical trials. Some studies revealed a risk for CRC associated with the use of anthraquinone-containing laxatives, some studies did not. However, a risk was also revealed for constipation itself and underlying dietary habits. Further investigations are needed to assess the carcinogenic risk definitely.

6. PHARMACEUTICAL PARTICULARS

Well-established use	Traditional use
Not applicable.	

7. DATE OF COMPILATION/LAST REVISION

26 October 2006