



EUROPEAN MEDICINES AGENCY
SCIENCE MEDICINES HEALTH

20 November 2018
EMA/HMPC/188805/2017
Committee on Herbal Medicinal Products (HMPC)

Assessment report on *Ruscus aculeatus* L. rhizoma

Based on Article 16d(1), Article 16f and Article 16h of Directive 2001/83/EC (traditional use)

Final

Herbal substance(s) (binomial scientific name of the plant, including plant part)	<i>Ruscus aculeatus</i> L., rhizoma
Herbal preparation(s)	a) Powdered herbal substance b) Dry extract DER (2.5-6.5:1); extraction solvent: water c) Dry extract DER (5-8.5:1) ; extraction solvent: ethanol 80% (V/V) d) Dry extract DER (6-9:1) ; extraction solvent: ethanol 96% (V/V)
Pharmaceutical form(s)	Herbal substance or preparation in solid dosage forms for oral use
Rapporteur(s)	First version: Antoine Sawaya & Jacqueline Viguet Poupelloz Revision: Carlos Cavaleiro
Peer-reviewer	G. Laekeman



Table of contents

Table of contents	2
1. Introduction	4
1.1. Description of the herbal substance(s), herbal preparation(s) or combinations thereof ..	4
1.2. Search and assessment methodology	5
2. Data on medicinal use	5
2.1. Information about products on the market	5
2.1.1. Information about products on the market in the EU/EEA Member States	5
2.1.2. Information on products on the market outside the EU/EEA	9
2.2. Information on documented medicinal use and historical data from literature	9
2.3. Overall conclusions on medicinal use	11
3. Non-Clinical Data	13
3.1. Overview of available pharmacological data regarding the herbal substance(s), herbal preparation(s) and relevant constituents thereof.....	13
3.1.1. Primary pharmacodynamics	13
3.1.1.1. Contractile effect on veins.....	13
3.1.1.2. Contractile effect on lymphatic vessels	22
3.1.1.3. Effect on vascular permeability	23
3.1.1.4. Effect on hypoxia-induced activation of endothelial cells	25
3.1.2. Secondary pharmacodynamics	29
3.1.3. Safety pharmacology	29
3.1.4. Pharmacodynamic interactions	29
3.1.5. Conclusions	29
3.2. Overview of available pharmacokinetic data regarding the herbal substance(s), herbal preparation(s) and relevant constituents thereof.....	30
3.3. Overview of available toxicological data regarding the herbal substance(s)/herbal preparation(s) and constituents thereof	33
3.3.1. Single dose toxicity.....	33
3.3.2. Repeat dose toxicity.....	34
3.3.3. Genotoxicity	34
3.3.4. Carcinogenicity.....	34
3.3.5. Reproductive and developmental toxicity	35
3.3.6. Local tolerance	35
3.3.7. Other special studies.....	35
3.3.8. Conclusions	35
3.4. Overall conclusions on non-clinical data	36
4. Clinical Data	36
4.1. Clinical pharmacology	36
4.1.1. Overview of pharmacodynamic data regarding the herbal substance(s)/preparation(s) including data on relevant constituents.....	36
4.1.2. Overview of pharmacokinetic data regarding the herbal substance(s)/preparation(s) including data on relevant constituents.....	37
4.2. Clinical efficacy	37
4.2.1. Dose response studies.....	37
4.2.2. Clinical studies (case studies and clinical trials)	38

4.2.2.1. Chronic venous insufficiency.....	38
4.2.2.2. Haemorrhoids.....	41
4.2.2.3. Orthostatic hypotension.....	42
4.2.2.4. Diabetic retinopathy.....	42
4.3. Clinical studies in special populations (e.g. elderly and children)	47
4.4. Overall conclusions on clinical pharmacology and efficacy.....	47
5. Clinical Safety/Pharmacovigilance.....	47
5.1. Overview of toxicological/safety data from clinical trials in humans.....	47
5.2. Patient exposure	47
5.3. Adverse events, serious adverse events and deaths.....	47
5.4. Laboratory findings.....	49
5.5. Safety in special populations and situations	49
5.5.1. Use in children and adolescents.....	49
5.5.2. Contraindications.....	49
5.5.3. Special Warnings and precautions for use.....	49
5.5.4. Drug interactions and other forms of interaction.....	49
5.5.5. Fertility, pregnancy and lactation.....	49
5.5.6. Overdose.....	50
5.5.7. Effects on ability to drive or operate machinery or impairment of mental ability	50
5.5.8. Safety in other special situations	50
5.6. Overall conclusions on clinical safety.....	50
6. Overall conclusions (benefit-risk assessment).....	51
Annex	54

1. Introduction

1.1. Description of the herbal substance(s), herbal preparation(s) or combinations thereof

- Herbal substance(s)

Ruscus aculeatus L. (*Liliaceae*) is a widely distributed European plant native from Western Europe. This plant has numerous appellations. The English name-Butcher's Broom-derives from the use by the European butchers of the stems to clean their cutting board not only because of their stiffness and solidity, but also because of the essential oil which was credited with antibacterial properties. *Ruscus aculeatus* is a small, clump-forming shrub with erect shoots bearing stiff, ovate, leaf-like phylloclade. Tiny green flowers appear in late winter and spring on the phylloclade. Both root and stem are used in preparations.

According European Pharmacopoeia VIII (2014), the herbal substance (*Rusci rhizoma*) consists of the dried, whole or fragmented underground parts of *Ruscus aculeatus* L containing not less than 1% of total saponins expressed as ruscogenins (mixture of neoruscogenin and ruscogenin) (dried drug).

- Herbal preparation(s)

Around the middle of the twentieth century spirostanol saponins (based on the aglycones ruscogenin and neoruscogenin) were first isolated from the rhizome. Several of these compounds have been then identified, as, ruscin, neoruscin, deglucoruscin and ruscocide, with their various sulphated and acetylated derivatives, ruscozepines A and B, aculeosides A and B (De Marino *et al.*, 2012; Masullo *et al.*, 2016). Both the above ground and below ground parts of the plant contains spirostanol saponins, although concentrations are higher in the below ground parts of the plant (Nikolov *et al.*, 1976). Besides spirostanol saponins, the extract of the rhizome contains also several other minor compounds such as other steroidal saponins and saponins, sterols, triterpenes, flavonoids, coumarins, including esculin and esculetin, sparteine, tyramine and glycolic acid (Dunouau *et al.*, 1996; Mimaki *et al.*, 1998). Barbic *et al.* (2013) suggested that the active substances in the rhizome are the saponins ruscin and deglucoruscin together with the coumarin glucoside esculin. Upon hydrolysis, saponins yield their aglycones, namely ruscogenin and neoruscogenin that for practical reasons are used to standardize the crude extracts.

Besides the dried powdered rhizome, herbal preparations are mainly based on its aqueous and aqueous-alcoholic extracts.

- Combinations of herbal substance(s) and/or herbal preparation(s) including a description of vitamin(s) and/or mineral(s) as ingredients of traditional combination herbal medicinal products assessed, where applicable.

Ruscus aculeatus extracts are also in commercial products combined with other products such as trimethylhesperidin chalcone (TMHC), ascorbic acid or *Melilotus officinalis* extract.

1.2. Search and assessment methodology

The Assessment Report on *Ruscus aculeatus* rhizome was revised considering the updates from relevant data published in the literature between 2008 and 2016.

Search engines used: Google, Google Scholar

Scientific databases: Web of Science; PubMed; Science Direct; Clinical Key; Cochrane Database of Systematic Reviews

Medical or Toxicological databases: Toxline

Pharmacovigilance resources: Data from EU and non-EU regulatory authorities, European database for suspected adverse drug reaction reports.

Thirty-four new records were retrieved under the search string [*Ruscus* or *Butcher's broom*]. Four were considered relevant, two reporting new phytochemical data, the others reporting new scientific pre-clinical data.

No information was received from the call of scientific data or from the interested parties.

2. Data on medicinal use

2.1. Information about products on the market

2.1.1. Information about products on the market in the EU/EEA Member States

Information on medicinal products marketed in the EU/EEA

Table 1 abridges medicinal products on the market, according information provided by the National Competent Authorities. In Belgium, Croatia, Denmark, Finland, Greece, Latvia, Netherlands, Poland, Portugal and Sweden there are no authorised or registered medicinal products on market.

Table 1: Overview of data obtained from marketed medicinal products

Active substance	Indication	Pharmaceutical form Strength Posology Duration of use	Regulatory Status (date, Member State)
Dried powdered herbal substance	a) Traditional herbal medicinal product to relieve symptoms of discomfort and heaviness of legs related to minor venous circulatory disturbances b) Traditional herbal medicinal product for symptomatic relief of itching and burning associated with	Hard capsule containing 350 mg of dried powdered herbal substance Posology: Adults: 3 times daily 1 capsule If symptoms persist for more than 2 weeks a doctor has to be consulted	THMP acc. to Article 16a of Directive 2001/83/EC (24.1.2013, Austria)

Active substance	Indication	Pharmaceutical form Strength Posology Duration of use	Regulatory Status (date, Member State)
	haemorrhoids		
Dried powdered herbal substance	<p>a) Traditional herbal medicinal product used to relieve symptoms of discomfort and heaviness of the legs related to minor venous circulatory disturbances, based on traditional use only</p> <p>b) Traditional herbal medicinal product used to relieve symptoms of itching and burning associated with haemorrhoids, based on traditional use only</p>	<p>Hard capsule containing 350 mg of dried powdered herbal substance</p> <p>Posology: Adults & the elderly: 1 capsule, 3 times daily during meals</p> <p>If the symptoms persist for more than 2 weeks during the use of the medicinal product, a doctor or a qualified healthcare practitioner should be consulted</p>	2011, TUR, United Kingdom
Dried powdered herbal substance	<p>Traditionally used:</p> <p>a) in subjective signs of venous insufficiency, such as heavy legs</p> <p>b) in haemorrhoidal symptoms</p>	<p>Hard capsules containing 350 mg of dried powdered herbal substance</p> <p>Posology: Adults: 1 hard capsule three times daily (till 6 if necessary)</p>	1986, TUR, France
Dried powdered herbal substance	<p>a) THMP to relieve symptoms of discomfort and heaviness of legs related to minor venous circulatory disturbances.</p> <p>b) THMP for symptomatic relief of itching and burning associated with haemorrhoids</p>	<p>Hard capsules containing 350 mg of dried powdered herbal substance</p> <p>Posology: Adults and adolescents: Single dose: 350 mg, 3 times per day.</p> <p>Duration of use: 2 weeks</p>	(1994), 2011, Spain
Dry extract (DER 2.5-6.5:1); extraction solvent: water	a) Traditional herbal medicinal product to relieve symptoms of discomfort and heaviness of legs	<p>Hard capsule containing 200 mg of extract</p> <p>Posology: Adults: Single dose: 200 mg, 2</p>	2000, TUR (2012), France

Active substance	Indication	Pharmaceutical form Strength Posology Duration of use	Regulatory Status (date, Member State)
	<p>related to minor venous circulatory disturbances</p> <p>b) Traditional herbal medicinal product for symptomatic relief of itching and burning associated with haemorrhoids</p>	<p>times day per day</p> <p>Duration of use: 4 weeks</p>	
<p>Dry extract (DER 4.5-6.5: 1), extraction solvent: water</p>	<p>a) Traditional herbal medicinal product to relieve symptoms of discomfort and heaviness of legs related to minor venous circulatory disturbances</p> <p>b) Traditional herbal medicinal product for symptomatic relief of itching and burning associated with haemorrhoids</p>	<p>Hard capsules containing 150 mg of dry extract</p> <p>Posology: Adults: single dose: 150 mg, 2-3 times daily</p> <p>If the symptoms persist for more than 2 weeks during the use of the medicinal product, a doctor or a qualified health care practitioner should be consulted</p>	<p>2010, TUR, Germany</p>
<p>Dry extract (DER 5.0-8.5: 1), extraction solvent: ethanol 80% (V/V)</p>	<p>a) Traditional herbal medicinal product to relieve symptoms of discomfort and heaviness of legs related to minor venous circulatory disturbances</p> <p>b) Traditional herbal medicinal product for symptomatic relief of itching and burning associated with haemorrhoids</p>	<p>Film-coated tablet containing 86 mg dry extract</p> <p>Adults: Single dose: 86 mg, 1-2 times daily</p> <p>If the symptoms persist for more than 2 weeks during the use of the medicinal product, a doctor or a qualified health care practitioner should be consulted</p>	<p>At least since 1976, TUR, Germany</p>
<p>Dry extract (DER: 6-9: 1), extraction solvent: ethanol 96% (V/V)</p>	<p>a) Traditional herbal medicinal product to relieve symptoms of discomfort and heaviness of legs related to minor venous</p>	<p>Soft capsules, containing 45 mg of dry extract</p> <p>Posology: Adults: single dose: 45 mg, 2 times daily</p>	<p>At least since 1976, TUR, Germany</p>

Active substance	Indication	Pharmaceutical form Strength Posology Duration of use	Regulatory Status (date, Member State)
	circulatory disturbances	If the symptoms persist for more than 2 weeks during the use of the medicinal product, a doctor or a qualified health care practitioner should be consulted	
Dry extract (DER 15-20: 1, extraction solvent: methanol 60% V/V)	<p>a) Traditional herbal medicinal product to relieve symptoms of discomfort and heaviness of legs related to minor venous circulatory disturbances</p> <p>b) Traditional herbal medicinal product for symptomatic relief of itching and burning associated with haemorrhoids</p>	<p>Soft capsules, containing 36.9 mg of dry extract</p> <p>Posology: Adults: 1 capsule, 2 times daily</p>	On market from 1993 to 2003

This overview is not exhaustive. It is provided for information only and reflects the situation at the time when it was established.

Information on relevant combination medicinal products marketed in the EU/EEA

Combinations of *Ruscus aculeatus L.* rhizome with other active ingredients are marketed in several EU member states:

- Capsule, hard containing 150 mg dry extract of *Ruscus aculeatus L.* rhizoma, 150 mg hesperidin methylchalkone, 100 mg ascorbic acid.

On the market in Slovak-Republic (1994) with the following indications:

- symptoms associated with hepatic venous and lymphatic vessels (a feeling of heaviness in the legs, pain, syndrome "restless legs")
- uterine bleeding caused contraception using low doses of progestin or IUD, after a physical examination and laboratory tests, symptoms associated with haemorrhoidal crisis.

Posology: Phlebology and gynaecology (treatment of symptoms associated with with impaired lymphatic vessels and for treatment of uterine bleeding) 2 to 3 capsules daily.

Proctology: (In the treatment of symptoms associated with varicose rectal) applied at a rate of 4 to 5 capsules daily.

On the market in Czech Republic (1996) with the following indications:

- a) for relief of symptoms of lymphatic insufficiency such as heavy legs, syndrome of “restless legs”, legs pain, cramps in the calves
- b) for symptomatic treatment of haemorrhoids

Posology: indication a) 1 capsule 2–3 times daily

indication b) 2-3 capsules twice daily, for long term use 1 capsule twice daily

On the market in Greece

- Cream (1 g) containing 16 mg of *Ruscus aculeatus* dry extract (DER 2.5-6.5:1; extraction solvent: water) and 20 mg of *Mellilotus officinalis* liquid extract (DER 0.07-0.20:1; extraction solvent: ethanol 30% V/V)

On the market in France since 1971 (TU 2013) and Poland with the following indication

Indication: Traditional herbal medicinal product to relieve symptoms of discomfort and heaviness of legs related to minor venous circulatory disturbances

Posology: Adults: Apply two times a day. Duration of use 4 weeks

Information on other products marketed in the EU/EEA (where relevant)

Not applicable

2.1.2. Information on products on the market outside the EU/EEA

Not applicable

2.2. Information on documented medicinal use and historical data from literature

Table 2: Overview of historical data

Herbal preparation	Documented use / Traditional use	Pharmaceutical form Strength Posology Duration of use	Reference
Dried powdered root	Traditional use: Subjective symptoms of chronic insufficiency such as sensation of heavy legs and in haemorrhoids symptoms	Capsules for oral use Adults: Daily dose: amount equivalent to ca. 10 mg of total ruscogenins	French Monograph : cahiers de l'Agence n° 3 (Afssaps, 1998)
Dry aqueous extracts	Traditional use: Subjective symptoms of chronic insufficiency such as sensation of heavy legs and in	Capsules for oral use Adults: Daily dose: amount equivalent to ca. 10 mg of total ruscogenins	French Monograph : cahiers de l'Agence n° 3 (Afssaps, 1998)

Herbal preparation	Documented use / Traditional use	Pharmaceutical form Strength Posology Duration of use	Reference
	haemorrhoids symptoms		
Solid or liquid extracts	Supportive therapy for symptoms of chronic venous insufficiency, such as painful, tired and heavy legs, tingling and swelling Supportive therapy for symptoms of haemorrhoids, such as itching and burning	Oral administration Adults: Single dose: amounts corresponding to 7-11 mg of total ruscogenins, once a day	ESCOP Monograph (ESCOP, 2003) It has to be noted that the ESCOP Monograph takes only into account the available data on <i>Ruscus aculeatus</i> alone
Alcoholic extracts of the whole plant	Supportive therapy for discomforts of chronic venous insufficiency, such as pain and heaviness, as well as cramps in the legs, itching and swelling Supportive therapy for complaints of haemorrhoids, such as itching and burning	Capsules for oral administration Adults: Amount standardised for 7-11 mg of ruscogenins as determined by the total amount of ruscogenin and neoruscogenin	Commission E Monograph (Commission E Monographs, 2001) It has to be noted that the Commission E Monograph takes into account the whole data available on <i>Ruscus aculeatus</i> including non-clinical and clinical data referring to combinations.
Dry extract (DER 5-8.5:1); extraction solvent: ethanol 80% (V/V)	Traditional herbal medicinal product to ease/soothe the feeling/sensation of heavy legs The product is a traditional herbal product for use in specified indications exclusively based on long-standing use	Oral administration Coated tablets or soft capsules Daily dosage and corresponding daily amount of total ruscogenins are not given	Bfarm (BfArM information, 2007)
Dry extract (DER: 6-9:1), extraction solvent: ethanol 96% (V/V)	a) Traditional herbal medicinal product to relieve symptoms of discomfort and	Soft capsules, containing 45 mg of dry extract	At least since 1976, TUR, Germany

Herbal preparation	Documented use / Traditional use	Pharmaceutical form Strength Posology Duration of use	Reference
	heaviness of legs related to minor venous circulatory disturbances	Posology: Adults: single dose: 45 mg, 2 times daily	

2.3. Overall conclusions on medicinal use

Ruscus aculeatus rhizome (dried powder herbal substance, ethanol dry extracts and water dry extracts) has been use in several EU member states in chronic venous insufficiency, haemorrhoids and varicose veins, linked to the vasoconstrictive effect of the ruscogenin and neoruscogenin.

This use is full documented, at least since 1976, for the following indications:

- a) relieve symptoms of discomfort and heaviness of legs related to minor venous circulatory disturbances
- b) symptomatic relief of itching and burning associated with haemorrhoids

Table 3: Overview of evidence on period of medicinal use

Herbal preparation Pharmaceutical form	Indication	Strength Posology	Period of medicinal use
a) Dried powdered herbal substance	Traditional herbal medicinal product used: a) to relieve symptoms of discomfort and heaviness of legs related to minor venous circulatory disturbances b) symptomatic relief of itching and burning associated with haemorrhoids	Powdered herbal substance for oral use. Posology: Adults and adolescents: 350 mg, 3 times per day Duration of use: 2 weeks	Since 1986, TUR France, Since 1994, TUR, Spain
b) Dry extract (DER 2.5-6.5: 1); extraction solvent: water ¹ (Dry aqueous extracts)	a) Traditional herbal medicinal product to relieve symptoms of discomfort and heaviness of legs related to minor venous circulatory	Dry extract for oral use Adults: Single dose: 150-200 mg, 2-3 times daily	French Monograph: cahiers de l'Agence n° 3 (AFSSAPS, 1998) 2000, TUR (2012), France & 2010, TUR, Germany

¹ The first list "Medicaments a Base De Plantes-Avis Aux Fabricants" issued by AFSSAPS in 1984, compiling ancient French bibliographical data, includes *Ruscus aculeatus* rhizome and its therapeutic indications. Although data concern to herbal tea, in the first Community herbal monograph on *Ruscus aculeatus* L., rhizome (EMA/HMPC/261938/2007), HMPC adopted the traditional use of the dry extract (DER:2.5-6.5:1; extraction solvent: water) in the basis that water extracts are comparable to herbal teas.

Herbal preparation Pharmaceutical form	Indication	Strength Posology	Period of medicinal use
	<p>disturbances</p> <p>b) Traditional herbal medicinal product for symptomatic relief of itching and burning associated with haemorrhoids</p>	<p>Daily dose: 450 mg</p> <p>If the symptoms persist for more than 2 weeks during the use of the medicinal product, a doctor or a qualified health care practitioner should be consulted</p>	
<p>c) Dry extract (DER 5.0-8.5:1); extraction solvent: ethanol 80% (V/V)</p>	<p>a) Traditional herbal medicinal product to relieve symptoms of discomfort and heaviness of legs related to minor venous circulatory disturbances</p> <p>b) Traditional herbal medicinal product for symptomatic relief of itching and burning associated with hemorrhoids</p>	<p>Dry extract for oral use</p> <p>Adults: Single dose: 86 mg, 1-2 times per day</p> <p>If the symptoms persist for more than 2 weeks during the use of the medicinal product, a doctor or a qualified health care practitioner should be consulted</p>	<p>At least since 1976, TUR, Germany</p>
<p>d) Dry extract (DER 6-9:1); extraction solvent: ethanol 96% (V/V)</p>	<p>a) Traditional herbal medicinal product to relieve symptoms of discomfort and heaviness of legs related to minor venous circulatory disturbances</p>	<p>Dry extract for oral use</p> <p>Posology: Adults: Single dose 45 mg, 2 times per day</p> <p>If the symptoms persist for more than 2 weeks during the use of the medicinal product, a doctor or a qualified health care practitioner should be consulted</p>	<p>At least since 1976, TUR, Germany</p>

Rapporteur comments:

1. The Community herbal monograph on *Ruscus aculeatus* L., rhizoma (EMA/HMPC/261938/2007) included the preparation 'dry extract (DER 6-9:1), primary solvent ethanol 96% (V/V) followed by water'. During the revision, no information was found anymore to support the use of water as secondary solvent. For this reason only the dry extract (DER 6-9:1), extraction solvent: ethanol 96% (V/V) should be considered.
2. During the revision, no base was found anymore to support the TU of the 'dry extract (DER 15-20:1), extraction solvent: methanol 60% (V/V)' considered in the Community herbal monograph on *Ruscus aculeatus* L., rhizoma (EMA/HMPC/261938/2007) since this preparation doesn't seem to fulfil 30 years of traditional use in the EU. Furthermore, there are no products marketed in EU containing this preparation, since 2003.

3. Non-Clinical Data

Non-clinical data presented in this assessment report should be taken into account according to the kind of preparations registered in the European Union. The request of information concerning *R. aculeatus* in medicinal products approved in the Member States showed that marketed products in France and Germany contain as preparations hydro-alcoholic extracts, aqueous extracts, and plant powder. However, in most of the non-clinical studies the characteristics of the extract used (mode of extraction, content in active substances, etc.) were not provided.

3.1. Overview of available pharmacological data regarding the herbal substance(s), herbal preparation(s) and relevant constituents thereof

3.1.1. Primary pharmacodynamics

Ruscus aculeatus being reported to be used in against venous insufficiency-related feeling/sensation of heavy leg, several *in vitro/in vivo* studies aimed at demonstrating its contractile properties on veins or lymphatic vessels as well as its capacity to improve vascular permeability.

Assessor's comment

It should be noted that studies conducted with the combination Ruscus extract and hesperidine methylchalcone were not taken into consideration. Indeed, the potential beneficial effect of the flavonoids or of the combination should not be attributed to Ruscus extract taken alone.

3.1.1.1. Contractile effect on veins

***In vitro* studies**

Studies performed on rings of canine saphenous veins

This model was first described Marcelon *et al.*, 1983b and Marcelon and Vanhoutte, 1984. Rings of canine saphenous veins were mounted in organ chambers filled with a physiological salt solution and connected to a force transducer for continuous recording of isometric tension. Prior to experimentation with *R. aculeatus* extract, the segments were placed at the optimal point of their length-tension relationship using standard electrical stimulation. After equilibration *R. aculeatus* extract was applied and the tension measured. Concentrations $\geq 3 \cdot 10^{-5}$ g/ml of *R. aculeatus* extract caused a dose-dependent increase in tension.

The effects of the following various pharmacological agents on the contractile effect induced by *R. aculeatus* were studied in an attempt to determine the mechanism of action underlying this contractile effect:

- Phentolamine ($3 \cdot 10^{-6}$ M), an α -adrenolytic agent, nearly abolished the contractile response to *Ruscus*;
- Cocaine ($3 \cdot 10^{-5}$ M), a blocker of the recapture of norepinephrine in the inter-synaptic gap, reduced the contractile response to *R. aculeatus*;
- 6-hydroxydopamine (10^{-6} M), inducing chemical denervation, reduced the contractile response to *R. aculeatus* similarly to cocaine;
- Tetrodotoxin (10^{-7} M), atropine (10^{-8} M), methysergide (10^{-7} M), indomethacin ($3 \cdot 10^{-5}$ M) did not impact on the contractile response to *R. aculeatus*;
- Adenosine ($2 \cdot 10^{-5}$ M) and verapamil ($2 \cdot 10^{-6}$ M) produced relaxation of the rings contracted by *R. aculeatus* (-37% and -50%, respectively);
- Acetylcholine (10^{-7} M, 10^{-6} M) caused further increase in tension (+23% and +37%, respectively).

In parallel, the same authors examined the effect of *R. aculeatus* extract on helical strips of canine saphenous veins connected to a force transducer and incubated in a moist tunnel-shaped chamber superfused with a Krebs-Ringer solution containing 7-1- $[^3\text{H}]$ -norepinephrine. *R. aculeatus* extract ($5 \cdot 10^{-4}$ g/ml) caused an increase in tension. In addition, the release of intact $[^3\text{H}]$ -norepinephrine and the overflow of all metabolites (dihydroxyphenylglycol, dihydroxymandelic acid, monohydroxyphenylglycol, normetanephrine) except vanillylmandelic acid were increased.

It was concluded that the vasoconstrictor response to *R. aculeatus* extract is not due to the activation of cholinergic, serotonergic, or prostaglandinergic receptors. The effect being inhibited by phentolamine but persisting after chemical denervation with 6-hydroxydopamine, a direct effect on postjunctional α -adrenergic receptors of the smooth muscle cells is suggested. However, it seems that the release of norepinephrine stored in adrenergic nerve endings is also involved in the contractile effect of *R. aculeatus* extract in view of the responses obtained with 6-hydroxydopamine, and cocaine, and in view of the increased overflow of $[^3\text{H}]$ -norepinephrine in veins previously incubated with this neurotransmitter.

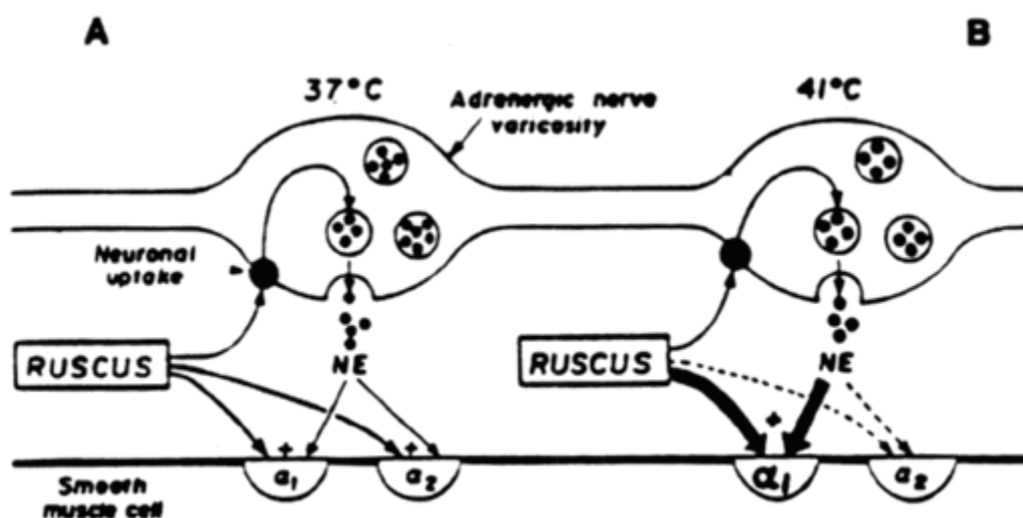
The release of norepinephrine from adrenergic nerve endings does not involve the initiation of spike electrogenesis, since tetrodotoxin had no influence on the effect provoked by *R. aculeatus* extract. The authors considered that the effects of adenosine, verapamil and acetylcholine reflected interference with the postjunctional effect rather than with its action on the adrenergic nerve endings. However, the effect by acetylcholine may be artificially biased by damaging of the endothelium in the experimental model; in a non de-endothelialised vessel a relaxing effect by NO release would be expected.

In another study, Rubanyi *et al.* (1984) confirmed that *R. aculeatus* extract could provoke the contraction of rings from canine saphenous veins. The tension increased as *R. aculeatus* extract concentration increased from 10^{-4} to 10^{-3} g/ml; the maximal contraction averaged 80% of the response to 10^{-4} M norepinephrine. It should be mentioned that a factor 10 within a dose relationship from basic tension to maximum has to be considered as very narrow in pharmacological terms. However, the contractile effect of *R. aculeatus* was depressed by prazosin or rauwolscine (α_1 - and α_2 -antagonist, respectively) from $5 \cdot 10^{-8}$ M and in a dose-dependent fashion, while it was abolished when both substances were used concomitantly. The same response was observed when phentolamine, a non-selective α -antagonist was used in the experiment described above (Marcelon *et al.*, 1983b;

Marcelon and Vanhoutte, 1984). This suggests that the contractile response of *R. aculeatus* in this model is due to α -adrenergic activation only (Rubanyi *et al.*, 1984).

At $5 \cdot 10^{-6}$ M, the inhibitory effect of rauwolscine was higher than that of prazosin. This difference was not observed when prazosin or rauwolscine was used with cocaine, both combinations depressing the contractile activity of *R. aculeatus* in the same extent. According to the authors, this suggests that norepinephrine released from adrenergic nerve endings activates preferentially postjunctional α_2 -adrenoceptors (Rubanyi *et al.*, 1984).

The same authors reiterated the study using *R. aculeatus* extract ($2 \cdot 10^{-4}$ g/ml), prazosin and rauwolscine ($5 \cdot 10^{-6}$ M) at 24°C, 37°C, and 41°C. Compared to the results obtained at 37°C, cooling decreased while warming increased the contractile response to *R. aculeatus* extract in rings of canine saphenous veins. These results were explicated in details by other authors. They acknowledged that these differences are due to the preferential modulation by temperature of the α_1 -adrenergic component of the response. At 37°C, the α_1 - and α_2 -adrenergic components of the response to *R. aculeatus* are equivalent (Rubanyi *et al.*, 1984; Marcelon and Vanhoutte, 1988).



Abbreviations: U1=neuronal uptake; NE=norepinephrine; +=activation

Figure 1: Proposed mechanism of action for *Ruscus* and effect of temperature (Marcelon and Vanhoutte, 1988)

To study the influence of the hormonal status of the animals on the contractile effect obtained with *R. aculeatus* extract, Miller *et al.* (1991a) treated ovariectomized female dogs with subcutaneous pellets containing a carrier substance (untreated), 17β -estradiol, progesterone, or 17β -estradiol and progesterone for 16-25 days. After that treatment period, the animals underwent surgery. The serum levels of hormones were measured and the carotid arteries and lateral saphenous veins removed and cut into rings after having been cleaned of connective tissue. The endothelium was removed in all rings. Thereafter, isometric tension was determined in absence or presence of a combination of prazosin or rauwolscine (10^{-7} M) following *in vitro* exposure to norepinephrine (10^{-8} to 10^{-4} M), tyramine (10^{-8} to 10^{-4} M), or *R. aculeatus* extract (10^{-6} to 10^{-3} g/ml) (Miller *et al.*, 1991a). Serum hormone levels are shown in the table 4.

Table 4: Oestrogen and progesterone measured in serum levels of ovariectomised dogs, according to their hormonal supplementation (Miller *et al.*, 1991a)

Group	Oestrogen level (ng/100 ml)	Progesterone level (ng/100 ml)
Untreated	1	80
Estrogen	30	400
Progesterone	–	750
Estrogen+Progesterone	30	2300

The contractile effect induced by electrical stimulation, norepinephrine and tyramine (indirect sympathomimetic substance, releases norepinephrine from adrenergic nerve endings) were not influenced by hormonal treatment. On the contrary, the contractile effect observed with *R. aculeatus* tended to be augmented in the progesterone group compared to the untreated group, the difference being significant compared to the group treated with estrogen and progesterone. Subsequent assays showed no effect of adrenergic blockade (prazosin plus rauwolscine) on the contractile effect of *R. aculeatus* in rings from untreated animals, whereas this effect was decreased in progesterone group and increased in estrogen and estrogen plus progesterone groups (Miller *et al.*, 1991a).

The same authors published further work performed on the same animals, which consisted in studying the influence of hormonal status and endothelium on the contractile effect of *R. aculeatus* (Miller *et al.*, 1991b).

In coronary arteries, *R. aculeatus* extract was able to initiate the release of endothelium-derived factors whose action is inhibitory to contractions initiated by the extract itself, PGF₂ α and norepinephrine. The release of this factor could involve the stimulation of muscarinic receptors because the relaxing effect was inhibited by atropine. Considering the nature of this relaxing factor, it is not probably prostacyclin because the experiment was performed in presence of indomethacin in the medium. Similarities with nitric oxide are evoked because the relaxing effect attributed to this relaxing factor is inhibited by haemoglobin (which inactivates NO) and by methylene blue (which inactivates guanylate cyclase). The hormonal status of the animals had no effect on the results obtained.

In femoral veins, the constrictor effect of *R. aculeatus* was influenced by the integrity of the endothelial cells and by the hormonal status of the animals. Indeed, the contractile effect obtained with *R. aculeatus* was greater in veins without endothelium, and greater in rings from animals receiving oestrogen plus progesterone. This result is contradictory to previous results obtained by these authors and described above [see (Miller *et al.*, 1991a)]. In varicose veins, the endothelium may be dysfunctional. Therefore, excitatory effect of the extract would prevail in this situation. Elevated oestrogen and progesterone serum levels would potentate this effect (Miller *et al.*, 1991b).

Study performed on rings of saphenous veins taken from rabbits

Harker *et al.* (1988) studied the influence of the hormonal status of female rabbits on the contractile effect obtained with *R. aculeatus* extract, using ovariectomized rabbits treated with placebo or 17 β -oestradiol (subcutaneous implants) for 14 to 21 days. After that treatment period, the animals underwent surgery. The serum level of oestradiol was measured and the lateral saphenous veins removed and cut into rings after having been cleaned of connective tissue. The endothelium was removed in all rings which were then suspended in organ chambers to measure isometric tension after

application of various agents. The effect of α -adrenergic antagonists and of temperature (24°C, 37°C, 41°C) on the contractile effect of *R. aculeatus* extract was also studied (Harker *et al.*, 1988).

The serum oestradiol level reported by the authors amounted to 1.0 ng/ml or less in control animals, while it reached 54 ng/dl in oestradiol treated animals. It should be noted that these figures may be wrong as 1.0 ng/ml is equivalent to 100 ng/dl.

In both groups, a dose-dependent contractile effect of *R. aculeatus* was observed. The involvement of postjunctional adrenoceptors was dependent upon the hormonal status of the animals. Indeed, contractions were not affected by adrenergic blockade in control animals, and partially inhibited by prazosin and rauwolscine in oestradiol-treated animals suggesting that hormone permits the expression of postjunctional α -adrenergic effect of *R. aculeatus*.

Cooling increased tension mediated by α_2 -adrenoceptors (rauwolscine) after application of *R. aculeatus* extract on rings of control rabbits, but not on rings of estradiol-treated animals. Moreover, it was concluded that warming did not affect the contractile response to *R. aculeatus* extract (Harker *et al.*, 1988).

Studies performed on rings of human saphenous and varicose veins

The contractile effect of *R. aculeatus* extract was investigated on rings of human varicose veins collected from females undergoing varicectomy (Marcelon *et al.*, 1988b). To study the influence of the hormonal status, three groups were considered:

- Rings from women at the end of menstrual cycle (14th to 25th day): n=3;
- Rings from women at the beginning of cycle (1st to 6th day): n=4;
- Rings from post-menopausal women: n=6.

As previously described, isometric tension was measured in organ chambers (at 37°C). The contractile effect of *R. aculeatus* extract (10^{-5} to 10^{-3} g/ml) was compared to that obtained with noradrenaline (10^{-4} M). In these conditions, the variation of hormone levels occurring during the menstrual cycle or in post-menopausal women did not influence the contractile effect of *R. aculeatus* extract which reached 43 to 52% of the contractile effect obtained with norepinephrine (Marcelon *et al.*, 1988b).

In another study, the role of endothelium in the contractile effect of *R. aculeatus* extract (10^{-6} to 10^{-3} g/ml) was evaluated on rings of varicose and saphenous veins taken from patients undergoing surgery for primary varicose veins (Miller *et al.*, 1994). In this study, *R. aculeatus* caused a concentration-dependent contractile effect to which varicose veins were more sensitive than saphenous veins. Additionally, contractions to *R. aculeatus* were not affected by removal of the endothelium. It was also confirmed in human veins that a major component of the contraction results from activation of adrenergic receptors because the blockade of adrenergic receptors significantly reduced that contractile effect. It is unlikely that *R. aculeatus* extract stimulates contraction by endothelin-A receptors in veins of these patients as the selective antagonist of these receptors (BQ-123) did not reduce the contractions either in absence or presence of adrenergic blockade. It is also unlikely that endothelin-B receptors are stimulated by *R. aculeatus* extract as varicose tributaries do not contract to sarafotoxin S6c (selective endothelin-B agonist). Therefore, the authors concluded that contractions to *R. aculeatus* in human varicose veins are independent of the endothelium and mediated by activation of adrenergic (but not endothelin-A) receptors on the smooth muscle (Miller *et al.*, 1994).

Studies performed on segments of canine and human saphenous/varicose veins

Branco and Osswald (1988) studied the influence of *R. aculeatus* extract on the uptake and metabolism of norepinephrine in segments of canine lateral saphenous veins, and of human

varicose/saphenous veins. In these tissues, the endogenous norepinephrine and dihydroxymandelic acid (DOMA) contents were determined. Thereafter, removal, accumulation and metabolism of [3H]-norepinephrine were studied without and with application of *R. aculeatus* extract (10^{-5} , 10^{-4} , 10^{-3} g/ml).

Compared to the normal canine vein, the normal human veins appeared sparsely innervated by the sympathetic system as shown by 10-fold lower norepinephrine content. However, it should not be concluded that it is not endowed with efficient adrenergic mechanism. Indeed, its capacity to metabolize norepinephrine is rather high and its high reactivity to adrenergic agonists is well known; in this study, the capacity of human vein to remove, accumulate and metabolise norepinephrine was one-half of that exhibited by the canine vein. Normal canine and human veins also differed in the pattern in which noradrenaline was metabolised. In human veins, O-methylation by catechol-O-methyltransferase (COMT) was of lesser importance than in canine veins. From this finding, the authors concluded that in what concerns disposition of noradrenaline, extrapolation from vessels of experimental animals to those of humans is not permissible.

The comparison of human normal and varicose veins showed that varicose veins contain and accumulate less norepinephrine. They have a high endogenous content and a raised rate of formation of DOMA.

Concerning the effect of *R. aculeatus* extract in human normal veins, it produced a concentration-dependent reduction of [3H]-norepinephrine accumulation (-50% at 10^{-3} g/ml, compared to control). Changes were less marked in the varicose veins (-50% at 10^{-3} g/ml, compared to control). The authors concluded that the highest concentration of *R. aculeatus* extract only may have released norepinephrine. However, all three concentrations affected the metabolism of norepinephrine: a concentration-dependent reduction of the formation of all norepinephrine metabolites was observed, and was similar in normal and varicose veins (DOMA formation was more affected in varicose veins).

Overall, quantitative but not qualitative differences were noted between human normal and varicose veins concerning the effect of *R. aculeatus*. It provoked a depression of the metabolism of norepinephrine and reduced its accumulation. This led to an increase in the concentration of norepinephrine in the biophase. This could explain the potentiation of the action of norepinephrine (Branco and Osswald, 1988).

Assessor's comments

The contractile effect of R. aculeatus extract was investigated on rings of saphenous veins taken from dogs and rabbits, and on human saphenous and varicose veins by recording of the isometric tension. Studies on the canine model showed that a concentration-dependent contractile effect was obtained within the concentration range of 10^{-5} - 10^{-3} g/ml for R. aculeatus extract. This was confirmed in the human model.

To determine the mechanism underlying this effect, further experimentations with various pharmacological agents and a study of tritiated norepinephrine metabolism were performed. They showed that in the canine venous rings, this effect was mediated by direct activation of postjunctional α_1 - and α_2 -adrenergic receptors, and by stimulation of the release of norepinephrine from adrenergic nerve endings. It should be noted that α -adrenergic activation only was involved. The involvement of adrenergic receptors in the contractile effect of R. aculeatus extract was confirmed in human veins (saphenous, varicose).

These results are in accordance with those obtained on segments of human normal and varicose veins, where R. aculeatus extract was shown to reduce norepinephrine accumulation and metabolism.

Although less marked in varicose veins, the reduction of norepinephrine accumulation was significant from 10^{-4} g/ml, a concentration compatible with those inducing a contractile effect on venous rings.

The influence of temperature was evaluated. In canine venous rings, cooling decreased while warming increased the contractile response to *R. aculeatus* extract obtained at 37°C. It was hypothesised that this differences is due to the preferential modulation by temperature of the α_1 -adrenergic component of the response. However, in venous rings taken from rabbits, cooling increased the contraction induced by *R. aculeatus* depending on the hormonal treatment, while warming had no effect.

Regarding the influence of the hormonal status on the contractile effect induced by *R. aculeatus* extract, contradictory results were obtained in animals. In one study performed with rings of varicose veins collected from women, the hormonal status did not impact on the contractile effect obtained. However, the number of samples tested was too low and the grade of varicosis of the veins was not studied, so that no definitive conclusion can be drawn. Similarly, conflicting results were obtained about a possible role of endothelium in the *R. aculeatus* effect in canine venous rings. In a human model, it was shown that "contractions to *R. aculeatus* in human varicose veins are independent of the endothelium and mediated by activation of adrenergic [...] receptors on the smooth muscle" (Miller et al., 1994).

Rings of human varicose veins were more sensitive than saphenous veins to the contractile effect obtained with *R. aculeatus* extract.

In vivo studies

Studies performed in dogs

To confirm the adrenergic mechanism of action of *R. aculeatus* extract, this drug was administered intravenously to anaesthetised dogs at doses ranging from 1 to 10 mg/kg (Marcelon et al., 1983a). The lateral saphenous vein was transilluminated and its diameter measured by a photoelectric cell. *R. aculeatus* caused a dose-dependent constriction of the saphenous vein and potentiated the dose-response curve to local norepinephrine. These responses were antagonized by phentolamine. In the conditions of the study, the venoconstriction caused by *R. aculeatus* was the result of α -adrenergic activation.

Studies performed on the hamster cheek pouch

A team of the University of Lund, Sweden, aimed at examining the effects of *R. aculeatus* extract on the microcirculation of the hamster cheek pouch and at gaining knowledge about the mechanism(s) underlying these effects. They recognised that "the usefulness of the healthy hamster cheek pouch preparation to study chronic venous insufficiency is debatable, but *in vivo* animal models for such study are clearly lacking" (Bouskela et al., 1994).

In the first experiment, six male hamsters were treated with *R. aculeatus* extract solution for 28 days by oral route at 0 and 150 mg/kg per day. At the end of the treatment period, animals were anaesthetised and an area of the cheek pouch isolated for measurement of vascular diameter (arterioles and venules) by means of a videotape recording device. The measurements were made on the same region in every animal. In these conditions, no significant difference was noted between control and treated animals regarding the body weight and mean arterial blood pressure. In the group receiving *R. aculeatus* extract, constriction of venules (diameter decreased by 30% compared to controls) and dilation of arterioles (diameter increased by 37% compared to controls) were observed (Bouskela, 1991).

IV injection (5 mg/kg) of *R. aculeatus* extract caused a venular constriction but did not significantly affect either the arteriolar diameter or the mean arterial blood pressure. It should be noted that the

measurement of vascular diameters were made before and after the injection of *R. aculeatus* extract. When the extract was added to the superfusate (topical application), venular constriction and arteriolar dilation were reported. Cooling to 25°C induced dilation of both types of vessels, whereas warming to 40°C produced the opposite effect (constriction) at 50.10⁻³ mg/ml and higher (Bouskela, 1991).

In a similar study, arteriolar and venular diameters were measured before (3 times, separated by 10 minutes intervals) and after (every 10 minutes for 60 minutes) intravenous injection of *R. aculeatus* extract at the dose of 5 mg/kg to 7 male hamsters. No effect on mean arterial blood pressure was detected. In these animals, venular but not arteriolar injection was observed at the cheek pouch level. When applied topically to 18 male hamsters, the *R. aculeatus* extract produced venular constriction and arteriolar dilation at the cheek pouch level. The effect of temperature was similar to what was reported above by Bouskela (1991) (Bouskela *et al.*, 1993b).

In an attempt to further characterize the mechanism of action of *R. aculeatus* extract on the hamster cheek pouch, another study was performed with male animals. *R. aculeatus* extract was used at concentrations ranging from 5 to 1000 µg/ml per minute applied topically (in the superfusate). At 50 µg/ml per minute and above, *R. aculeatus* extract application induced venular but not arteriolar constriction. The venular constriction amounted to approximately 10%. At the concentration of 0.2 mg/ml per minute, the venular constriction remained for 120 minutes (determined every 10 minutes) whereas the internal diameter of arterioles was not modified. Therefore, the concentration of 0.2 mg/ml per minute was used in further explorations. The venular constriction evoked by *R. aculeatus* was blocked by low concentrations (10⁻⁹ M) of prazosin and diltiazem (these drugs did not induce significant venular dilation at this concentration), and only by high concentrations (10⁻⁶ M) of rauwolscine. The authors postulated that at high concentration, rauwolscine may not be selective only for α₂-adrenoceptors. It was concluded that venular constriction in the hamster cheek pouch was mediated by calcium and preferentially by α₁-adrenoreceptors. Additionally, the authors hypothesised that the lack of effect on arterioles could be explained by augmented liberation of endothelium-derived relaxing factors on the arteriolar side which probably overrides its constrictor properties (Bouskela and Cyrino, 1994; Bouskela *et al.*, 1994).

Complementary mechanistic studies

A team of the Faculdade de Medicina, Oporto tested the effect of *R. aculeatus* extract on segments of veins. They compared at the microscopic level human varicose and normal veins, saphenous vein taken from dogs aged 4 months to 7 years and above, and canine saphenous vein obtained after surgical denervation. Human varicose vein and canine denervated saphenous veins shared common characteristics: thickening of the vessel wall due to increase in extracellular material and smooth muscle cells hypertrophy, abnormalities of smooth muscle cells (signs of increased protein synthesis). In dogs, impact of surgical sympathetic denervation on the structure of the vein was independent of the age of the animals. Additionally, the [³H]-norepinephrine content was determined in vessels of dogs (4 months to 7 years and above), and in human varicose/normal veins. The authors showed that sympathetic innervation was lowered in veins taken from aged dogs, and in human varicose veins (compared to normal veins). Therefore, it was concluded that human varicose vein behaves like a partially denervated vessel and many of its structural and biochemical peculiarities appear to be linked to the reduced sympathetic supply of the vein. The authors considered that the denervated canine vein represents an interesting model for the study of the human varicose vein (Azevedo *et al.*, 1991).

Teixeira and Osswald (1988) produced denervation of lateral saphenous vein in anaesthetised dogs by using artery clamps applied down- and upstream of the segment for 5 min. During the surgical intervention, osmotic minipumps were implanted in the subcutaneous tissue of dogs' leg and connected by a catheter to the plantar branch of the denervated vein. These minipumps allowed IV injection of saline (containing Na₂-EDTA+heparin) ± *R. aculeatus* extract (50 µg/kg per hour). Animals recovered

for 5 days and were then re-operated to remove the segment of the denervated vein and of the contralateral vein (to serve as control). This allowed to study the effect of denervation (injection of saline), of *R. aculeatus* on denervated vein (injection of *R. aculeatus* and saline). For the morphological study of the vein, the dimension of smooth muscle cells was taken as the most important criterion of morphological alterations caused by denervation (at the extraneuronal level). In collected segments, the endogenous norepinephrine content was measured by HPLC; denervation was considered successful and complete when the norepinephrine content was decreased by 95%. Additionally, the O-methylating capacity of these tissues was reflected by the total amount of O-methylated metabolite (OMI), determined after incubation of vein strips with ³H-7-(±)-isoprenaline for 30 minutes (Texeira and Osswald, 1988).

This study showed that intravenous infusion of *R. aculeatus* extract protected the extraneuronal component of the venous tissue against the deleterious consequences of denervation. Indeed, it induced prevention of increase in smooth muscle cell diameter, and partial prevention of the impairment of O-methylating capacity of tissues. However, *R. aculeatus* extract administration had no effect on the denervation process itself (Nad depletion of at least 95% in denervated veins) (Texeira and Osswald, 1988).

Assessor's comments

The venoconstricting property of R. aculeatus extract shown in vitro was confirmed in in vivo models. In anaesthetised dogs, a dose-dependent contractile effect was shown for R. aculeatus extract doses ranging from 1 to 10 mg/kg given intravenously, and attributed to result of α-adrenergic activation only.

The IV administration of R. aculeatus extract (5 mg/kg) to hamsters induced venular constriction at the level of cheek pouch microcirculation, without any impact on the diameter of arterioles or mean arterial blood pressure. However, topical application of the extract resulted in venular constriction and arteriolar dilation. Cooling was shown to induce dilation of both types of vessels, whereas warming produced their constriction. Studies conducted with pharmacological agents showed that venular constriction in the hamster cheek pouch was mediated by calcium and preferentially by α1-adrenoreceptors. The dilation of arterioles was considered to result from augmented liberation of endothelium-derived relaxing factors on the arteriolar side.

The oral route was also investigated in this experimental model; the dose of 150 mg extract per kg per day for 28 days induced venular constriction (internal diameter decreased by 30%) and arteriolar dilation (internal diameter increased by 37%) which was also attributed to liberation of endothelium-derived relaxing factors on the arteriolar side.

Briefly, it can be concluded that the contractile effect of R. aculeatus extract on veins was shown in two in vivo experimental models by intravenous route (anaesthetised models, hamster cheek pouch), topical application (hamster cheek pouch), and oral route (hamster cheek pouch, 150 mg extract per kg per day). The involvement of the α-adrenergic system was confirmed. Arteriolar dilation was noted in some of these studies, and particularly when R. aculeatus extract was administered orally, but no effect on the mean arterial pressure was detected. It could result from induction of liberation of endothelium-derived relaxing factors on the arteriolar side. Additionally, in a canine model of human varicose vein, IV injection of R. aculeatus extract for 5 days prevented against the occurrence morphological alteration of smooth muscle cells, and against the impairment of O-methylating capacity of tissues.

The animal models used to demonstrate the venoconstricting activity of R. aculeatus extract are acceptable, taking into consideration the lack of experimental models reproducing the physiopathological complexity of the chronic venous insufficiency. These models are those commonly

used for the evaluation of drugs belonging to this therapeutic class, even if the use of functional exploration methods (e.g. Doppler) could have been used to complete the weight of evidence.

3.1.1.2. Contractile effect on lymphatic vessels

In vitro studies

Study performed on rings of canine lymphatic vessels

Thoracic ducts were removed from dogs, cleaned of connective tissue, and cut into rings. Those rings were put in organ chambers filled with a physiological salt solution and connected to a force transducer for continuous recording of isometric tension. The substances tested, directly added to the bath solution, and were norepinephrine (10^{-8} to 10^{-4} M) and *R. aculeatus* extract (10^{-5} to $2 \cdot 10^{-3}$ g/ml). To clarify the mechanism of action of *R. aculeatus* extract, some α -adrenergic antagonists were used: phentolamine ($3 \cdot 10^{-6}$ M), prazosin ($3 \cdot 10^{-7}$ M) and rauwolscine ($3 \cdot 10^{-7}$ M) (Marcelon *et al.*, 1988a).

In these experimental conditions, it was shown that both norepinephrine and *R. aculeatus* extract caused a concentration-dependent contraction of the lymphatic thoracic duct rings. The contractile activity of *R. aculeatus* was partially inhibited by prazosin or rauwolscine, and completely eliminated by phentolamine. Taking into account that venous smooth muscle cells have been isolated in mesenteric and thoracic lymphatics, the importance of the presence of adrenergic nerve endings at this level for the control of lymphatic pumping, and the results obtained in this study, the authors concluded that *R. aculeatus* causes a similar adrenergic activation of both lymphatic collectors and cutaneous veins. The contractile effect obtained on lymphatic rings was obtained at similar concentrations than those inducing contraction of venous rings (Marcelon *et al.*, 1988a).

Study performed on bovine lymphatic vessels

In order to determine the mechanism of noradrenaline action on lymphatic vessels, various studies were performed using electric stimulation on bovine mesenteric lymphatic vessels. Among those studies, one was performed to gain knowledge about the desensitisation of adrenoceptors after application of the endogenous transmitter (norepinephrine) or *R. aculeatus* extract (McHale, 1991).

While application of norepinephrine (10^{-6} M) induced an increase of contraction frequency which returned rapidly to control values as the result of receptors desensitisation, application of *R. aculeatus* extract (30 μ g/ml, i.e. $3 \cdot 10^{-5}$ g/ml) had similar excitatory effect which was sustained for the duration of drug perfusion. According to the author, this might suggest that *R. aculeatus* is acting on different receptors in these vessels from those through which norepinephrine is acting (McHale, 1991).

Assessor's comment

R. aculeatus extract was shown to exert contractile effect in vitro on dog thoracic and bovine mesenteric lymphatic vessels. The results concerning the mechanism involved are contradictory between both experiments. Indeed, results obtained with canine tissues suggest that the mechanism of action is similar to the one described at the venous level, i.e. α -adrenergic activation. According to its authors, the study performed on bovine tissue suggests that pathways involved differ from those of norepinephrine. However, they did not further discuss on the role of chemical stability to explain their results (norepinephrine is probably less chemically stable than *R. aculeatus*).

In vivo studies

A study was conducted in anaesthetised dogs, after isolation of a lymph vessel parallel to the saphenous vein at the ankle level and ligation of other vessels (Pouget *et al.*, 1991). After *R. aculeatus* extract intravenous injection, the total protein concentration (Pt) was measured by UV-spectrometry in lymph collected at 10 minutes intervals. This allowed the calculation of oncotic pressure (Ponc) using

the following formula: $Ponc=0.45 Pt+0.078 Pt$. The doses of *R. aculeatus* extract administered were 1, 2 and 5 mg/kg because they were in the range of the doses reported as veinotonic (Marcelon *et al.*, 1983a).

In some animals, the activity of *R. aculeatus* was studied in presence of calcium antagonism by nifedipine. *R. aculeatus* extract (5 mg/kg) was administered and its activity followed for 1 hour. Then, nifedipine was administered intravenously (0.1 mg/kg, the activity of this dose was maintained for more than 1 hour). After 30 minutes, a second injection of *R. aculeatus* extract was performed, and its activity again followed for 1 hour.

At 2 and 5 mg/kg, *R. aculeatus* extract caused a rise in lymph flow without affecting lymph pressure. This reflected the augmentation of the contractility of lymph vessels. This is in agreement with the results of an *in vitro* study performed on peripheral lymphatic vessels (McHale, 1991). It was also shown that *R. aculeatus* caused contraction of the isolated thoracic duct of the dog *in vitro* (Marcelon *et al.*, 1988a). This adds weight to the hypothesis of an enhancement of the "pumping system" suggested by the results obtained *in vivo* in this study, with enhanced propulsion of peripheral lymph toward the heart.

R. aculeatus also caused a rise in oncotic pressure which suggests that it induces a favourable effect on edema.

The activity of *R. aculeatus* extract remained unchanged after nifedipine injection. This suggests that the mechanism underlying the effect of *R. aculeatus* on peripheral lymphatics does not involve the opening of voltage operated calcium channels.

Assessor's comment

In anaesthetised dogs, administration of R. aculeatus extract improved the contractility of peripheral lymph vessels. Taken together with in vitro results, this suggests that R. aculeatus extract enhances the lymphatic pumping system thus favouring a better return of peripheral lymph to the heart. Additionally, a rise in oncotic pressure was detected, which could suggest a favourable effect on oedema.

However, it should be noted that no study was conducted by the intended therapeutic route of administration, i.e. the oral route. It is questionable whether active substance(s) involved in this effect would 1) be absorbed by oral route and 2) undergo significant hepatic metabolism impairing its (their) activity at the lymphatic level.

3.1.1.3. Effect on vascular permeability

***In vitro* studies**

An experiment was conducted using segment of the lateral ear vein of the pig cannulated on both sides and put into an organ bath of Krebs solution. *R. aculeatus* extract (0.05%) was applied intraluminally for 15 minutes. Then, ethacrynic acid (0.1%) was added to damage the endothelium (2.5 minutes). Finally, the permeability to low- and high molecular substances was measured. To quantify the water edema, an aqueous solution of the dye Evansblue was applied, and the protein edema was measured with bovine serum marked by the dye, each being applied under a pressure of 30 cm H₂O for 30 minutes. The content of Evansblue was then determined by photometry (Hönig and Felix, 1989).

The results obtained showed that *R. aculeatus* extract reduced the water- and protein permeability induced by ethacrynic acid. The drug had to be present when the damaging agent was applied; if it was washed 1 minute before, no protective effect was observed. The authors also showed that no chemical binding occurred between the drug and the damaging agent.

To elucidate this oedema-protective mechanism, saponins from *R. aculeatus* were applied for 15 minutes and then washed. The permeability of these vessels was increased compared to controls. According to the authors, the surface-active saponins so strongly bind to the endothelium that they leave holes when being removed.

Assessor's comment

In vitro on ear vein of pig, R. aculeatus extract had protective effects against ethacrynic-induced oedema. Saponins may play an important role by anchoring in the endothelium and taking part to the membrane structure.

Several spirostanol saponins and esculin from *Rusci rhizoma* were assessed by Barbic *et al.* (2013) concerning the effects on the permeability of endothelial cells measuring their ability to reduce the thrombin-induced hyperpermeability of endothelial cells. The highest activities were observed for the spirostanol saponins deglucoruscin and ruscin and for esculin at 10 µM, resulting a significant reduction of the thrombin-induced hyperpermeability (in 41.9%, 42.6% and 53.3%, respectively). Results provide insight into the pharmacological mechanism by which spirostanol saponins and esculin can contribute to the efficacy of Butcher's broom against chronic venous disorders (Barbic *et al.*, 2013).

In vivo studies

Study performed on the hamster cheek pouch

The number of leakage sites on hamster cheek pouch 4 was measured by fluorescence microscopy after IV injection of fluorescein-labelled dextran. The number of leakage sites (diameter of the fluorescent spot > 100 µm) was determined 2, 5, 7, 10, 15, 20, and 20 minutes after the beginning of each topical application of histamine. Each histamine application lasted 5 minutes with a minimum interval of 40 minutes between each application. The first application of histamine, after injection of FITC-dextran, was made prior to any drug treatment and thus served as a positive control. *R. aculeatus* extract was then applied topically in the following concentrations: 0.002, 0.02, 0.2 and 2 mg/ml per minute before the subsequent application of histamine. In the group dosed at 0.2 mg/ml per minute, the influence of prazosin, rauwolscine and diltiazem (10^{-9} M to 10^{-5} M) on *R. aculeatus* effect was studied (Bouskela *et al.*, 1994).

In this experimental model, histamine increased the number of fluorescent vascular leakage sites from postcapillary venules, which evidence an increase in macromolecular permeability (quantified as the number of leaky sites in the prepared area). The topical application of *R. aculeatus* extract induced a dose-dependent inhibition of the macromolecular permeability-increasing effect of histamine (-25% at 0.002 mg/ml per minute, almost -50% at 0.2 mg/ml per minute). This effect was blocked by prazosin and diltiazem, but not by rauwolscine. Overall, it was concluded that *R. aculeatus* extract inhibited the microvascular permeability induced by histamine. This effect would be mediated by calcium and preferentially by α 1-adrenoreceptors (Bouskela *et al.*, 1994).

These findings confirm the results previously obtained in a similar study which showed that topical application of *R. aculeatus* extract has a protective effect against leakage of FITC-dextran in the cheek pouch of hamsters after administration of various permeability-increasing substances, i.e. bradykinin, histamine, and leukotriene B4 (Bouskela *et al.*, 1993a).

Study performed on a feline model of edema

Edema was induced in anaesthetised cats by IV injection of ethacrynic acid. *R. aculeatus* extract was administered one hour before edema induction. Its antiedema effect was evaluated in hindlegs by measuring the water and protein content of the edema (Felix *et al.*, 1984).

Compared to control animals, the protein content of the oedema was decreased in *R. aculeatus* pre-treated cats. Water content was not affected, which would suggest that *R. aculeatus* extract only influences the passage of plasma protein in the interstitium. However, in *R. aculeatus* pre-treated animals, it was demonstrated that waters flows into the tissues more slowly than in control animals. Indeed, the capillary filtration coefficient increased to a smaller degree in *R. aculeatus* pretreated animals, compared to control ones. Therefore, it was concluded that *R. aculeatus* extract inhibited the destruction of the endothelium by ethacrynic acid without totally suppressing it.

The effective dosage was 20 mg/kg by IV route, 10-20 times higher by oral route (administration 4 hours before induction of edema). By oral route, the effective dose decreased after subchronic administration and reached 20-40 mg/kg (4-6 days).

It is also reported that a protective effect was observed with ruscogenin at 4 mg/kg by IV route. This effect was weaker than the effect reported with the whole extract at 20 mg/kg. This last contained 2.5% ruscogenin, leading to a ruscogenin dose of 0.5 mg/kg (Felix *et al.*, 1984).

Assessor's comment

The antiedema effect reported for R. aculeatus extract in vitro was confirmed in two in vivo studies. After IV administration, R. aculeatus extract inhibited the microvascular permeability induced by histamine on hamster cheek pouch, an effect suggested to be mediated by calcium and preferentially by α_1 -adrenoreceptors. In a feline model of ethacrynic acid-induced edema, IV administration of 20 mg R. aculeatus extract/kg decreased the protein content of the edema and slowed the water flow into the tissues. This is in agreement with the increase in oncotic pressure reported by Pouget et al (1991) in lymphatic vessels of anaesthetised dogs administered 2 and 5 mg/kg R. aculeatus extract intravenously.

However, none of these studies was performed by the oral route. In the study performed by Felix et al. (1984), the anti-edema effect of R. aculeatus extract was investigated by oral route. It can be concluded that the oral effective dose approximates 200-400 mg/kg, or 20-40 mg/kg per day after repeated administrations for 4-6 days (the criteria used by the authors to decrease the dose were not detailed). Additionally, it should also be noted that this effect is not only due to ruscogenin, but also to other components of the extract.

3.1.1.4. Effect on hypoxia-induced activation of endothelial cells

Endothelium plays a role in the development of varicose veins. Hypoxic conditions which develop during blood stasis are able to activate endothelial cells to release inflammatory mediators and growth factors. Inflammatory mediators induce neutrophil adherence and activation. Those activated neutrophils may then infiltrate and induce damage in the subendothelial layers. On the other hand, the growth factors trigger smooth muscle cell proliferation, leading to a thickening and disorganisation of the media as observed in the wall of varicose veins. Therefore, a study was performed on human umbilical vein endothelial cells (HUVEC) to determine whether *R. aculeatus* extract could prevent endothelial cell activation by hypoxia (Bouaziz *et al.*, 1999).

R. aculeatus extract was shown to prevent the activation of endothelial cells (HUVEC) incubated in hypoxia conditions for 2 hours. This was demonstrated by measuring the ATP content of control and treated cells. The hypoxia-induced decrease in ATP was concentration-dependently inhibited by *R. aculeatus* extract, the effect being maximal at 50 $\mu\text{g/ml}$.

Hypoxia strongly increased phospholipase A2 (PLA2) activity, more than 2-fold the value obtained in normoxic conditions (\downarrow ATP \rightarrow cytosolic calcium \rightarrow PLA2 activation). When HUVEC cells were incubated with *R. aculeatus* extract under hypoxia, PLA2 activation was inhibited (50% inhibition at 0.05 $\mu\text{g/ml}$).

Additionally, *R. aculeatus* extract inhibited neutrophil adherence to HUVEC (PLA2 activity → prostaglandin, platelet activating factor → ↑ in HUVEC adhesiveness for neutrophils).

Table 5: Overview of the main non-clinical data/conclusions

Herbal preparation tested	Posology	Experimental model	Reference	Main non-clinical conclusions
<i>R. aculeatus</i> extract (extract details not given)	3.10 ⁻⁵ g/ml 5.10 ⁻⁴ g/ml	<i>In vitro</i> Rings and helical strips of canine saphenous veins	Marcelon <i>et al.</i> , 1983b and Marcelon and Vanhoutte, 1984	<i>R. aculeatus</i> extract caused a dose-dependent increase in isometric tension
<i>R. aculeatus</i> extract (extract details not given)	5.10 ⁻⁸ M	<i>In vitro</i> Contraction of rings from canine saphenous veins	Rubanyi <i>et al.</i> , 1984	The tension increased as <i>R. aculeatus</i> extract concentration increased from 10 ⁻⁴ to 10 ⁻³ g/ml; the contractile response of <i>R. aculeatus</i> is due to α-adrenergic activation
<i>R. aculeatus</i> extract (extract details not given)	10 ⁻⁶ to 10 ⁻³ g/ml	<i>Ex-vivo</i> Ovari-ectomised female dogs with subcutaneous pellets containing 17β-estradiol, progesterone, or 17β-estradiol and progesterone	Miller <i>et al.</i> , 1991a	Contractile effect observed with <i>R. aculeatus</i> tended to be augmented by progesterone
<i>R. aculeatus</i> extract (extract details not given)	Not available	<i>In vivo</i> Female ovariectomised rabbits treated with 17β-oestradiol (subcutaneous implants) and <i>Ruscus</i> extract	Harker <i>et al.</i> , 1988	The contractile effect of <i>Ruscus</i> was dependent upon the hormonal status of the animals but not upon warming
<i>R. aculeatus</i> extract (extract details not given)	10 ⁻⁵ to 10 ⁻³ g/ml	<i>In vitro</i> Human varicose veins collected from females undergoing varicectomy at different stages of the	Marcelon <i>et al.</i> , 1988b	Variation of hormone levels occurring during the menstrual cycle or in post-menopausal women did not influence the contractile effect induced by <i>the</i>

Herbal preparation tested	Posology	Experimental model	Reference	Main non-clinical conclusions
		menstrual cycle		extract
<i>R. aculeatus</i> extract (extract details not given)	10 ⁻⁶ to 10 ⁻³ g/ml	<i>In vitro</i> Rings of varicose and saphenous veins taken from patients undergoing surgery for primary varicose	Miller <i>et al.</i> , 1994	<i>R. aculeatus</i> caused a concentration-dependent contractile effect to which varicose veins were more sensitive than saphenous veins
<i>R. aculeatus</i> extract (extract details not given)	<i>R. aculeatus</i> extract (10 ⁻⁵ , 10 ⁻⁴ , 10 ⁻³ g/ml)	<i>In vitro</i> Segments of canine lateral saphenous veins and of segments of human varicose/saphenous veins	Branco and Osswald (1988)	<i>R. aculeatus</i> exert a depression of the metabolism of norepinephrine reducing its accumulation. Changes less marked in the varicose veins
<i>R. aculeatus</i> extract	Intravenous from 1 to 10 mg/kg	<i>In vivo</i> Anaesthetised dogs for measurement of the diameter of saphenous vein	Marcelon <i>et al.</i> , 1983a	Dose-dependent constriction of the saphenous vein result of α -adrenergic activation
<i>R. aculeatus</i> extract	<i>R. aculeatus</i> extract (10 ⁻⁵ to 2.10 ⁻³ g/ml).	<i>In vitro</i> Thoracic lymphatic ducts from dogs	Marcelon <i>et al.</i> , 1988a	<i>R. aculeatus</i> extract caused a concentration dependent contraction of the lymphatic thoracic duct rings <i>R. aculeatus</i> causes a similar adrenergic activation of both lymphatic collectors and cutaneous veins
<i>R. aculeatus</i> extract	<i>R. aculeatus</i> extract (0.05%)	<i>In vitro</i> Pig segment of the lateral ear vein	Hönig and Felix, 1989	<i>R. aculeatus</i> extract reduced the water and protein permeability induced by ethacrynic acid
<i>R. aculeatus</i> extract (extract details not given)	Topical application 0.002, 0.02, 0.2 and 2 mg/ml per minute, for 10	<i>In vivo</i> Hamster cheek pouch	Bouskela <i>et al.</i> , 1994	<i>R. aculeatus</i> extract inhibited the microvascular permeability induced by histamine

Herbal preparation tested	Posology	Experimental model	Reference	Main non-clinical conclusions
	minutes			
<i>R. aculeatus</i> extract	Oral route, 20-40 mg/kg repeated administration for 4-6 days	<i>In vivo</i> : cats	Felix <i>et al.</i> , 1984	Oedema was decreased in <i>R. aculeatus</i> pre-treated cats. Extract inhibited the destruction of endothelium induced by the ethacrynic acid
<i>R. aculeatus</i> extract (extract details not given)	50 µg/ml	<i>In vitro</i> Human umbilical vein endothelial cells incubated in hypoxia conditions. Measure of ATP	Bouaziz <i>et al.</i> , 1999	Decrease in ATP was concentration-dependently inhibited by <i>R. aculeatus</i> extract
<i>R. aculeatus</i> extract (extract details not given)	Intravenous injection of 1, 2 and 5 mg/kg	<i>In vivo</i> Dogs, after isolation of a lymph vessel in presence and absence of calcium antagonism by nifedipine Evaluation of total protein concentration by UV-spectrometry on oncotic pressure	Pouget <i>et al.</i> , 1991	<i>R. aculeatus</i> caused a rise in oncotic pressure which suggests the induction of a favourable effect on edema
<i>R. aculeatus</i> extract (extract details not given)	Oral route at 0 and 150 mg/kg per day	<i>In vivo</i> Male cheek pouch hamsters for measurement of arterioles and venules	Bouskela <i>et al.</i> , 1994	In the group receiving the extract orally the constriction of venules (diameter decreased by 30% compared to controls) and dilation of arterioles (diameter increased by 37%) were observed
<i>R. aculeatus</i> extract (extract details not given)	Intravenous; topical application; oral route	<i>In vivo</i> Contractile effect of <i>R. aculeatus</i> extract on veins anaesthetised models hamster cheek pouch	Texeira and Osswald, 1988	Arteriolar dilation was noted in some of these studies, and particularly when <i>R. aculeatus</i> extract was administered orally

Herbal preparation tested	Posology	Experimental model	Reference	Main non-clinical conclusions
Isolated compounds: Spirostanol saponins: deglucoruscin and ruscin; Esculin	10 µM	<i>In vitro</i> of human microvascular endothelial cells (CDC/EU.HMEC-1)	Barbic <i>et al.</i> , 2013	Significant reduction of the thrombin-induced hyperpermeability of endothelial cells

3.1.2. Secondary pharmacodynamics

No data available.

3.1.3. Safety pharmacology

No study available.

Assessor's comment

Considering the pharmacological profile of R. aculeatus extract, i.e. stimulation of α -adrenergic system, the lack of a safety pharmacology study evaluating its potential effects on the cardiovascular function must be taken into consideration when evaluating undesirable effects in clinical studies. No toxicology study evaluating this endpoint is available.

In the studies performed on the hamster cheek pouch, the mean arterial blood pressure was not modified after IV administration of 5 mg per kg R. aculeatus extract, and oral administration at the dose of 150 mg extract per kg. The need of such a safety pharmacology study addressing cardiovascular aspects should be discussed in light of clinical safety data.

3.1.4. Pharmacodynamic interactions

No study available.

Assessor's comment

R. aculeatus extract contains substances having α -adrenergic stimulating properties. Moreover, it was shown to displace norepinephrine from adrenergic nerve endings. Therefore, pharmacodynamic drug-drug interactions regarding drugs potentiating or antagonising the α -adrenergic are plausible.

3.1.5. Conclusions

Primary pharmacodynamics studies performed *in vitro* and *in vivo* using various experimental models showed that *R. aculeatus* extract possess a contractile activity on veins. This activity is mediated by stimulation of the α -adrenergic system. *In vitro* mechanistic studies showed that direct activation of postjunctional α 1- and α 2-adrenergic receptors, and stimulation of the release of norepinephrine from adrenergic nerve endings were involved. Although this effect does not appear to be clearly influenced by the hormonal status (estrogens, progesterone), it seems potentiated by temperature.

In *in vivo* studies, this vasoconstricting activity was shown after intravenous and oral routes; in the hamster cheek pouch model, local application of the extract (i.e. in the superfusate) was also effective. It should be noted that only one study was conducted by the oral route: at the level of hamster cheek

pouch microcirculation, the dose of 150 mg extract per kg per day administered for 28 days induced venular constriction (internal diameter decreased by 30%) and arteriolar dilation (internal diameter increased by 37%) without any impact on the mean arterial blood pressure, the latter effect being attributed to liberation of endothelium-derived relaxing factors on the arteriolar side.

Similarly, other primary pharmacodynamics studies showed that *R. aculeatus* extract exerts a contractile effect on lymphatic vessels in anaesthetised dogs at 2 and 5 mg/kg administered intravenously. A rise in oncotic pressure suggested a favourable effect on edema. This was confirmed in a feline model of ethacrynic acid-induced edema. The effective dosage amounted to 20 mg/kg by intravenous route, and 10-20 times higher by oral route. However, after subchronic administration (4-6 days), the oral effective dosage decreased to reach 20-40 mg/kg per day. The same study showed that ruscogenin was also effective, but that other components of the extract were involved to obtain maximal activity.

Due to the mechanisms underlying the effect of *R. aculeatus* extract, pharmacodynamic drug-drug interactions are plausible with drugs potentiating or antagonising the α -adrenergic system. However no reports on drug-drug interaction were found. Considering the pharmacological profile of *R. aculeatus* extract, i.e. stimulation of α -adrenergic system, the lack of a safety pharmacology study evaluating its potential effects on the cardiovascular function must be taken into consideration when evaluating clinical studies. No toxicology study evaluating this endpoint is available. In the studies performed in the hamster cheek pouch model, the mean arterial blood pressure was not modified after IV administration of 5 mg/kg *R. aculeatus* extract, and oral administration at the dose of 150 mg extract per kg. No reports on cardiovascular effects were found.

3.2. Overview of available pharmacokinetic data regarding the herbal substance(s), herbal preparation(s) and relevant constituents thereof

Three studies were performed in rats and dogs to gain knowledge in the pharmacokinetics of *R. aculeatus* extract (Chanal *et al.*, 1978; Chanal *et al.*, 1981; Bernard *et al.*, 1985). In each study, the extract was tritiated according to the Wilzbach technique. Briefly, this labelling method involved an incubation period with tritium gas (5-7 days) followed by the elimination of labile tritium. Analysis by TLC was then performed and showed each time that most of the radioactivity was associated with the sapogenins (80 to 94%, depending on the study). According to the authors, other labelling methods could not be applied:

- Carbon-14 labelling was not appropriate because the extract contains a multiplicity of sapogenins
- Labelling by biosynthesis would have required years of growth (the extract is made from the rhizome)
- These sapogenins cannot be chemically synthesized because of the lack of any precursor

Assessor's comment

*As indicated by other authors, the Wilzbach method used to label the extract is controversial (Bernard et al., 1985). Tritium labelling of a complex mixture such as the *R. aculeatus* extract used in this study presents some disadvantages which preclude a full confidence in the results obtained. In particular, the stability of the labelling is not justified (possible ^3H - ^1H exchanges after the labelling, etc.).*

*Additionally, no quantification of $^3\text{H}_2\text{O}$ was performed in any sample of the studies. After ^3H - ^1H exchange or metabolism, $^3\text{H}_2\text{O}$ mix with the body water pool. The elimination of tritiated water from the organism being particularly slow, the monitoring of 3 hours radioactivity to study the pharmacokinetics of *R. aculeatus* extract's compounds is likely to produce biased results. Overall, these*

studies should not be taken into account for regulatory purposes (however main results will be described below).

Absorption

Blood kinetics of radioactivity were determined in a study involving 2 male Wistar rats administered the labelled-extract orally (Chanal *et al.*, 1978). Blood samples were collected at 0.25, 0.5, 1, 2, 3.5, 5, 6, 8 and 24 hours after administration. Radioactivity level, expressed as % radioactivity per 10 ml of blood in relation to the dose administered, was measured by Liquid Scintillation Counting (LSC).

In both rats, blood radioactivity reached a level close to the maximum 2 hours after the administration of the extract. Thereafter, radioactivity level remained practically constant over the remaining 22 hours. The authors concluded that the half-lives of the tritiated compounds were relatively long, on the order of several days.

In another study involving oral administration of the labelled extract to 6 male Wistar rats, blood samples were collected at 0.25, 0.75, 1.75, 2.25, 3.5, 5, 6.5, 8, 24 and 48 hours after administration (Chanal *et al.*, 1981). Radioactivity level was expressed as % radioactivity per 10 ml of blood in relation to the dose administered, and the figures given are the mean for the 6 rats. Relatively high radioactivity levels were reached from 3.5 hours and the Tmax amounted to 8 hours. Blood levels declined slowly and remained high after 24 hours.

Assessor's comment

Two studies performed on a limited number of rats after oral administration of a labelled R. aculeatus extract showed that radioactivity was detected in the blood for more than 24 hours. The authors suggest that the triturated compounds of the extract would have relatively long half-lives. However, such a conclusion is speculative because the triturated water, which is slowly eliminated, was not quantified in the samples collected. Therefore, no conclusion can be drawn from these studies.

*It should be noted that the results obtained in the studies performed by Chanal *et al.* in 1978 and in 1981 are contradictory; in the first study, blood radioactivity remained stable over 25 hours, whereas in the second one, a decrease of blood radioactivity starts after 8 hours.*

Distribution

Tissue distribution was investigated by whole body autoradiography (WBA) in macaque monkeys administered a labelled extract of *R. aculeatus* by either intravenous (n=1, extract dissolved in sodium chloride 0.9%), or oral (n=4, extract dissolved in the content of one oral ampoule) routes (Bernard *et al.*, 1985).

WBA performed 24 hours after IV administration showed intense labelling in the bile, the contents of the distal digestive tract and to a lesser extent the urine. Moderate label was found in the circulating blood, and relatively substantial label in the renal and hepatic parenchyma, spleen, bone marrow, and adrenals (cortex). The intense binding of radioactivity at the hepatic and renal levels was confirmed by LSC.

Two hours after oral administration, highest radioactivity levels were detected in the bile, digestive and urinary contents; liver and kidney were labelled uniformly. Blood activity was slightly lower than after IV administration. The radioactivity levels decreased the following 5 hours except for digestive organs (urine, bile, faeces). After 24 hours, non-negligible radioactivity levels persisted in circulating blood, renal and hepatic parenchyma, bone marrow, spleen, adrenal cortex, contents of the distal digestive tract, bile and urine.

Assessor's comment

Radioactivity was mainly found at the hepatic and renal levels 24 hours after both intravenous and oral administration of the labelled R. aculeatus extract. It should be noted that intense radioactivity was found in the bile. Again, it remains unknown if the radioactivity arises from any of the compounds of the extract, or from triturated water. Therefore, no conclusion can be drawn.

However, deep distribution of radioactivity in the bone marrow after IV and oral administrations was shown, thus underlying the need of genotoxicity studies.

Metabolism

To determine the nature of plasma radioactivity, blood samples collected 2 hours after oral administration of a radio-labelled extract to 2 rats were extracted with methanol, and analysed by TLC (Thin Layer Chromatography). Results indicate that 39% of the radioactivity deposited corresponded to the sapogenin spot. The authors conclude that there was a "considerable fraction of the unchanged product" in the plasma after 2 hours (Chanal *et al.*, 1978).

Assessor's comment

This experiment does not allow drawing any conclusion on the metabolism of R. aculeatus extract due to concerns already expressed on labelling method and experimental procedures, and to the poor separating capacity of the method employed (TLC).

Elimination

Ninety six hours after oral administration of the labelled extract to 2 rats, approximately 18% and 29% of radioactivity was recovered in the urine and faeces, respectively. It should be noted that excretion occurred mainly during the first 24 hours. A non-negligible part of the fraction recovered in the faeces arised from notable biliary excretion (approximately 10% of the administered radioactivity) (Chanal *et al.*, 1978).

Urinary excretion amounted to 32-35% in Wistar and Atrichis rats administered a labelled extract orally, while the corresponding figure for faecal excretion was slightly higher, i.e. 39-45%. Most of the radioactivity was excreted within the first 24 hours. The authors report that the existence of an enterohepatic cycle was confirmed in a study conducted by intravenous administration (one third of total radioactivity eliminated in the faeces) (Chanal *et al.*, 1981).

In monkeys, radioactivity was excreted in the urine (26%) and in the faeces (6.5%) 24 hours after IV administration (n=1). Those figures amounted to 20% and 23% for urine and faeces, respectively, 24 hours after oral administration. The authors suggested the existence of an enterohepatic cycle considering the extent of faecal elimination and bile activity (Bernard *et al.*, 1985).

Assessor's comment

First, it should be noted that these figures were obtained from a limited number of animals. Additionally, it seems that the amount of radioactivity recovered from urine and faeces appears to be insufficient in all studies (less than 50% to 80% of the administered dose). Taking into consideration the relatively long half-lives of the labelled compounds which was hypothesised from blood kinetics, an incomplete radioactivity collection after either 24 hours or 96 hours seems coherent.

However, as most of the studies showed that excretion occurred mainly during the first 24 hours, and in view of the stable blood radioactivity levels after 24 hours, accumulation of radioactivity could occur after repeated administrations.

Radioactivity was excreted in urine and faeces, with faecal elimination slightly above. The authors suggest that *R. aculeatus* extract labelled components undergo an enterohepatic cycle. However, the excretion of a compound in the bile and a high amount of faecal elimination are not sufficient to prove the existence of an enterohepatic cycle. This suggestion should be rather regarded as a hypothesis to be further investigated.

This experiment does not allow to draw any conclusion on the metabolism of *R. aculeatus* extract due to concerns already expressed on labelling method and experimental procedures.

Assessor's overall conclusions on pharmacokinetics

The pharmacokinetics of *R. aculeatus* extract was investigated in rat (2 studies) and in monkey (1 study). In each study, the extract was triturated according to the Wilzbach technique. Briefly, this labelling method involved an incubation period with tritium gas (5-7 days) followed by the elimination of labile tritium. Afterwards, the extract could be administered to animals. However, this labelling method presents some disadvantages. For example, the stability of the labelling remains unknown (possible $3\text{H}-1\text{H}$ exchanges after the labelling, etc.).

Additionally, the authors did not perform a quantification of triturated water in the samples collected. This represents a major bias because the radioactivity measured in samples cannot be attributed without any doubt to the compounds of the *R. aculeatus* extract. Indeed, $^3\text{H}_2\text{O}$ can mix with the body water pool after $^3\text{H}-1\text{H}$ exchange or metabolism, and the elimination of triturated water from the organism is particularly slow.

Overall, it is concluded that these studies should not be taken into account for regulatory purposes because they are endowed with major bias precluding a full confidence in the results obtained.

One concern arises from the monkey distribution study, where in-depth distribution of radioactivity was reported at the bone marrow level. While it remains unknown if radioactivity is due to any component of the extract or to triturated water, the worst-case scenario should be considered. Therefore, this finding strongly underlies the need of genotoxicity studies with *R. aculeatus* extract.

3.3. Overview of available toxicological data regarding the herbal substance(s)/herbal preparation(s) and constituents thereof

3.3.1. Single dose toxicity

The acute toxicity of an ethanolic extract of *R. aculeatus* was investigated in dogs and guinea pigs (Caujolle *et al.*, 1953; ESCOP, 2003):

- In 6 male and female dogs, death occurred within 1 hour following intravenous infusion of the extract at doses ranging from 0.83 g/kg to 1.8 g/kg. The frequency of cardiac contractions was progressively decreased but was not attributed by the authors to a toxic effect on the myocardium because the hearts of treated dogs could react normally to epinephrine. Additionally, the blood pressure was decreased. The monitoring of respiratory function showed that at toxic doses, tachypnoea occurred and was sometimes associated with rhythm perturbation. At lethal doses, hyperventilation was followed by fatal apnoea. The authors considered that at high doses, the *R. aculeatus* ethanolic extract that they administered to dogs produced cardiovascular and respiratory reactions. The respiratory centres were deeply affected and apnoea always preceded the cardiac arrest so that death was attributed to respiratory alteration. Moreover, at high doses, hyperglycaemia was reported (1.82 to 2.84 g/l).

- In 8 male guinea pigs, the intraperitoneal injection of the ethanolic extract induced no toxic symptoms at doses lower than 1.5 g/kg. Animals receiving 2g/kg and above died.

Estimation of the oral and intraperitoneal LD₅₀ values of an ethanolic fluid extract of *R. aculeatus* in rats and mice revealed differences depending on the harvest time of the plant, the route of administration, and the use of roots or rhizomes. The oral LD₅₀ of the rhizome extract could not be determined in rats because administration of doses inducing 100% mortality could not be reached. In mice, it amounted to 24.69-33.73 ml/kg in mice. After intraperitoneal administration, the LD₅₀ of the rhizome extract reached 1.15-1.70 ml/kg in mice, and 2.07-2.39 ml/kg in rats. Root extract was found to be more toxic than rhizome extract in both species. The observed symptoms of intoxication were convulsion, paralysis and gastro-intestinal inflammation with dysentery. Animals died following respiratory failure. Autopsies revealed pronounced irritation of the mucosa and strong visceral congestion (Boucard *et al.*, 1967; ESCOP, 2003).

Assessor's comment

Extracts used in both studies were obtained by ethanolic extraction but contents in ruscogenin and neoruscogenin are unknown.

*The first study in dogs reported a mean LD₀ of 1.20 g/kg (0.83–1.8 g/kg) by intravenous route. The authors attribute the cardiovascular findings observed at high dose (decreased frequency of cardiac contractions, decreased blood pressure) to be secondary to alteration of respiratory centres. However, this is not sufficiently established, particularly if the α -adrenergic activity of *R. aculeatus* components is taken into consideration. In the second study, the oral LD₅₀ of mice reached 25-34 ml/kg. Death occurred by respiratory failure in rats and mice treated by oral and intraperitoneal routes.*

3.3.2. Repeat dose toxicity

The ESCOP monograph reports the findings of a 26-week toxicity studies performed in male rabbits by administration in the diet (Roux, 1969; ESCOP, 2003). A *R. aculeatus* extract was administered to 17 animals at 2 g/kg, and to 19 animals at 5 g/kg. Five animals served as controls. It is stated that body weight and blood counts did not reveal any difference between treated animals and controls.

Assessor's comment

The lack of information available on this (unpublished) study precludes its use in safety evaluation. Indeed, no precision on the extract administered to the animals is given (mode of extraction, content in active substances, etc.), and the choice of the rabbit as a toxicology species is remarkable. Usually, rat or rabbit are used as rodent, and dog or monkey as non-rodent. The parameters monitored in treated animals are not indicated, except body weight and blood count which is insufficient.

3.3.3. Genotoxicity

No data available.

Assessor's comment

*The lack of any genotoxicity study precludes the listing of *R. aculeatus* (EU list entry). Additionally, as no long-term study is available, the carcinogenic risk cannot be estimated.*

3.3.4. Carcinogenicity

No data available.

3.3.5. Reproductive and developmental toxicity

The ESCOP monograph reports the findings of an unpublished study conducted in female pregnant rat by administration of a preparation containing ethanolic *R. aculeatus* extract, trimethylhesperidin methylchalcone, methyl-4-esculetol and ascorbic acid (Lapie, unpublished ; ESCOP, 2003). Twenty female rats received a daily dose of 2.4 ml of the preparation corresponding to 0.24 ml of *R. aculeatus* extract and equivalent to 25 times the recommended dose for humans. Twenty animals served as control. Treatment started one week before conception and continued until delivery. No sign of intoxication was noted in treated animals. The fertility of females in the treatment group was comparable to that in the control group and this offspring did not show any teratogenic sign.

Assessor's comment

The lack of information available on this (unpublished) study precludes its use in safety evaluation (route of administration, number of resorptions, viable fetuses, number of fetuses examined for evaluation of visceral/skeletal abnormalities, etc.). Only one dose level was tested, which is not acceptable. The study was performed on an association of a mixture (R. aculeatus extract) with other components; a conclusion on the teratogenic risk of R. aculeatus extract only cannot be drawn.

It is considered that the embryo-fetotoxic risk of R. aculeatus extract is unknown; adequately conducted reproduction toxicity studies are lacking.

3.3.6. Local tolerance

No data available.

3.3.7. Other special studies

No data available.

3.3.8. Conclusions

Extracts used in two acute toxicity studies were obtained by ethanolic extraction. Their characteristics (content in various compounds, e.g. ruscogenin and neoruscogenin) remain unknown. Potential variability compared to the extract(s) intended for therapeutic cannot be evaluated. In dogs, a mean intravenous LD₀ of 1.20 g/kg was measured, while the oral LD₅₀ of mice reached 25-34 ml/kg with another extract. The authors attribute the cardiovascular findings observed in dogs at high doses (decreased frequency of cardiac contractions, decreased blood pressure) to be secondary to alteration of respiratory centres. In rats and mice, death occurred by respiratory failure too. No safety pharmacology study is available on cardiovascular and respiratory systems. Considering the α -adrenergic activity of *R. aculeatus* components, the need of such studies should be discussed in light of clinical safety data.

No other toxicity studies are available. The ESCOP monograph reports repeat-dose toxicity and reprotoxicity studies performed in rabbits and rats, respectively. However, these studies remain unpublished so that the information available is very sparse. Therefore, they cannot be taken into consideration for safety evaluation in this assessment report.

The lack of adequately conducted genotoxicity and embryo-fetal toxicity studies precludes the listing of *R. aculeatus*. Additionally, as no long-term study is available, the carcinogenic risk cannot be estimated.

3.4. Overall conclusions on non-clinical data

Non-clinical data on *Ruscus aculeatus* rhizome activity supports the traditional use as medicinal product to relieve symptoms of discomfort and heaviness of legs related to minor venous circulatory disturbances as well as for relief of itching and burning associated with haemorrhoids.

The reported pharmacological effects are not considered contradictory to the traditional uses.

Specific data on pharmacokinetics and interactions are not available.

Non-clinical information on the safety of *Ruscus aculeatus* rhizome is scarce.

As there is no information on reproductive and developmental toxicity, the use during pregnancy and lactation cannot be recommended.

Oral administration of *Ruscus aculeatus* rhizome can be regarded as safe at traditionally used doses with the exception of patients with severe renal or cardiac disease e.g. renal and heart failure.

Tests on reproductive toxicity, genotoxicity and carcinogenicity have not been performed.

4. Clinical Data

Clinical data on efficacy and safety of *R. aculeatus* alone are very limited.

4.1. Clinical pharmacology

4.1.1. Overview of pharmacodynamic data regarding the herbal substance(s)/preparation(s) including data on relevant constituents

Clinical pharmacology on *R. aculeatus* is not well documented. Two publications have been identified.

In a randomised, placebo-controlled, double-blind, crossover, 4-arm study, 20 healthy volunteers (11 men and 9 women aged between 20 and 43 years) took a single dose of four different treatments immediately before the first measurements: 450 mg *R. aculeatus* extract, 450 mg trimethylhesperidin chalcone (TMHC), 900 mg of a combination of the two substances or a placebo. A 1-week wash-out phase was provided between the treatments. The venous function was determined by plethysmography. Volumetric measurements were performed in orthostatic conditions with normal blood flow (foot volume) and after a pronounced ischemia (tissue volume). The difference between foot volume and tissue volume corresponds to the blood volume. Hemodynamic and volumetric reactions were monitored before intake and after 70, 90, 120 and 150 minutes. *R. aculeatus* extract caused a significant decrease in venous capacity and venous outflow. *R. aculeatus* extract also significantly reduced tissue volume compared to placebo whereas the decrease in blood volume was not significant (Rudofsky, 1991).

Assessor's comments:

Details on the R. aculeatus extract were not specified.

A long term study was also performed with 141 patients, with chronic venous insufficiency (CVI), who were recruited to a randomised, double-blind, multicentre study. The cause of late CVI was either primary varicosis or post-thrombotic syndrome (PTS). After a two weeks washout, they were given 4 weeks of treatment with 3x2 and then 8 weeks with 2x2 capsules of R. aculeatus extract or placebo.

The patients venous pump function during toe-stand exercises were also investigated by plethysmography.

In CVI patients there was a continuous decrease in the foot and ankle volume after a 12-week treatment with active substance whereas the volume increased under placebo. The tissue volume was reduced by the same degree as the foot volume in PTS patients. In varicosis patients, the reduction in leg swelling was due to a decrease in tissue volume and volume of blood stored in the veins during orthostasis (Rudofsky, 1991).

Assessor's overall conclusions on pharmacodynamics

On the basis of publications, the quality of the two available pharmacodynamic studies cannot be evaluated. For example, the characteristics of the patients are incomplete as well as the design of the studies. The statistical analysis is not given. Moreover, the characteristics of the *R. aculeatus* extract are not specified. However, the findings corroborate the preclinical pharmacological properties described in section II.2.1 the alpha-adrenergic effect and thus a venous vasoconstrictive effect that account for a reduction in volume of blood stored in the veins and for a stimulating effect on the lymphatic drainage. These two pharmacological properties assume a positive effect in patients suffering from chronic venous insufficiency.

Dose-effect studies are missing which preclude from justifying the dosage regimen used in the clinical studies.

Other methods of functional exploration could have been used to evaluate the effect on veins (e.g. Doppler).

4.1.2. Overview of pharmacokinetic data regarding the herbal substance(s)/preparation(s) including data on relevant constituents

During a pilot study involving three volunteers, the major spirostanol glycosides of *R. aculeatus* (degluconeoruscin and degluکورuscin) were detected in human plasma after an oral administration of 1g of *R. aculeatus* extract (Rauwald and Grunwidl, 1991).

Assessor's comments:

*The only conclusion that can be drawn from this study is that degluconeoruscin and degluکورuscin seems to be absorbed after *R. aculeatus* extract oral administration. Of note, the characteristics of the extract are not given. The publication wasn't detailed enough to determine the grade of CVI, the dose administered, the statistical analysis and the Doppler procedure.*

*Available pharmacokinetics data are too scarce. The pharmacokinetics of *R. aculeatus* extract should further studied.*

4.2. Clinical efficacy

4.2.1. Dose response studies

According to the provided literature no dose-finding studies have been conducted with *R. aculeatus* extract alone neither in patients with chronic venous insufficiency nor in patients suffering from haemorrhoids. Monographs dosage recommendations are empirical.

With respect to the ESCOP and the Commission E monographs an adult daily dose should be equivalent to an amount of 7 to 11 mg of total ruscogenins. The French herbal preparations containing *R. aculeatus* alone recommend a daily dose equivalent to an amount of around 10 mg of total ruscogenins with regard to a traditional use in subjective symptoms of chronic insufficiency such as sensation of heavy legs.

4.2.2. Clinical studies (case studies and clinical trials)

The efficacy of *R. aculeatus* has been proposed in the different following indications. A review of the literature for each indication has been performed.

4.2.2.1. Chronic venous insufficiency

Chronic Venous Insufficiency (CVI): symptoms and therapeutic measures Chronic venous insufficiency is a common disease which is characterised by symptoms due to disorders of the venous return in the lower leg and in foot. The situation leads to an increased pressure in the veins and lack of blood flow to the legs and feet. The main symptoms of chronic venous insufficiency are: leg heaviness, aching, dilated or unsightly veins and oedema (swelling). In the more severe cases, patients can have skin colour changes, recurrent skin infections and chronic ulcers.

Two main options to treat symptoms due to chronic venous insufficiency are firstly compression therapy which has demonstrated an efficacy to reduce leg volume, to hinder progression and to reduce symptoms. This therapy can be completed or replaced by the use of systemic veno-active drugs. Physical activity such as for example walking or swimming can also be suitable.

Clinical efficacy and safety data:

Only one clinical study, with *R. aculeatus* extract alone, performed in patients with chronic venous insufficiency (CVI) has been found in the literature Vanscheidt *et al.* (2002).

Methodology:

This multicenter double-blind, randomised, placebo-controlled clinical study has been performed with a *R. aculeatus* extract alone and published (Vanscheidt *et al.*, 2002). The aim of this trial was to confirm the efficacy and safety of a *R. aculeatus* dry extract (15-20:1) extraction solvent, methanol 60% (V/V) according to the latest scientific standards.

Design: The study enrolled women suffering from chronic venous insufficiency (Widmer classification grades I and II, CEAP classification 3-5) and was conducted at 10 different centres in Germany. Randomisation was carried out after a 2-week placebo run-in phase. The treatment phase lasted 12 weeks. Checkpoints were scheduled after 4, 8, and 12 weeks of treatment.

Treatments: Active treatment and placebo were taken twice daily (morning and evening) orally with some fluid. One capsule active treatment contained 36.0-37.5 mg dry extract from *Ruscus aculeatus* rhizome with a drug extract ratio of 15-20:1 (excipient methanol 60%) corresponding to an amount of around 4.5 mg of total ruscogenins.

Outcomes:

- The primary variable of the study was the change in foot and lower leg volume, measured as the area under the curve of volume changes over the 12-weeks treatment-time of (AUB0-12, area under baseline).
- Among the secondary variables "changes in leg volume" and "changes in subjective symptoms" were evaluated after 12 weeks of treatments.

All measurements were carried out at the same time of the respective days in the late afternoon or early evening. Before the measurements the patients underwent a 45 minutes temperature and cardiovascular equilibrium period in a sitting position. The leg volume was determined by water displacement. The ankle and lower leg circumference were measured using a measuring tape. The

measurements were carried out at the lateral and medial ankle and the middle of the middle of the lower leg.

The subjective symptoms: tiredness and heaviness, sensation of tension, tingling sensation, and pain were assessed at each visit before the volume measurements. The subjective symptoms were evaluated by a 10 cm Visual Analog Scale (VAS) where 0 was equivalent to “no complaints” and 10 to “strongest complaints”.

The quality of life was investigated at visits 1 and 5 by a disease specific and validated questionnaire on the quality of life which included subjective symptoms and life style judgement (Launois *et al.*, 1996). It can be compared with FLQA (Freiburger Life Quality Assessment). Global efficacy was assessed by the investigator with a four point scale (very good, good, moderate and bad).

Results: Overall, 167 patients were screened, of whom 166 were randomised and included in the study. Eighteen had insufficient data and were excluded from the efficacy analysis.

Efficacy on leg volume: For the AUB₀₋₁₂, the median values (min/max) were -656 [ml per day] (-13972/5908) for the *R. aculeatus* extract and 175 [ml per day] (-22795/4970) for placebo which reflected a decrease of leg volume in the *Ruscus* group but an increase in the placebo group. Statistical analysis revealed a significant treatment contrast ($p < 0.001$). Moreover, the median values of all parameters showed a decrease over time reflecting an increasing reduction of leg volume, ankle circumference, lower leg circumference, and subjective symptoms in the *R. aculeatus* extract group. With the placebo treatment generally, the baseline values were maintained for all parameters. Significant differences between the treatment groups were seen for the volume changes of the lower leg after 8 and 12 weeks, ankle circumference after 4, 8 and 12 weeks, lower leg circumference for 8 and 12 weeks, and finally after 12 weeks for the subjective symptom 1 (heaviness and tiredness) and subjective symptom 2 (sensation of tension).

Evaluation of subjective symptoms: The global efficacy of the *R. aculeatus* extract was evaluated by the investigator as very good and good more frequently, whereas placebo was more frequently assessed as moderate and bad ($p = 0.0498$). A significantly positive correlation coefficient was calculated for the changes of each of the subjective symptoms 1 (heaviness and tiredness), 2 (sensation of tension) and 3 (tingling tension) and leg volume changes.

Quality of life: The Quality of life did not reveal any changes after 12 weeks of treatment for both groups.

Assessor's comments:

The study of Vanscheidt et al. (2002) is deserving of being the first and the only one clinical study performed with a Ruscus extract alone. Nevertheless, it is always rather difficult to appreciate the real quality of a study through a publication. Several methodological insufficiencies remain unsolved and information is missing such as: lack of complete protocol with modalities of randomisation, sample calculation and power of the study.

Moreover, the respect of the double-blind procedure cannot be evaluated.

Eighteen patients have been excluded from the analysis for insufficient data of which no information is given. At 12 weeks, all missing data have been replaced by a LOCF value (last observation carried forward). However, we do not know how many data are missing.

Lastly, the study has been conducted in 10 centres in Germany; however, the homogeneity between centres has not been evaluated in terms of recruitment or clinical practices. Taking into account these overall comments, efficacy results should be interpreted with caution.

According to the author, the study was designed in accordance with the guidelines for testing drugs for chronic venous insufficiency (CVI), i.e. study design with oedema reduction as the primary variable. It has to be noted that these guidelines were written by the author himself and published in "Phlebologie" after the beginning of this study i.e. April 1999 (Vanscheidt et al., 2000) and that oedema is not considered as a cardiovascular risk factor in the recommendations made by the European Society of Hypertension and the European Society of Cardiology in 2007.

Although we can agree with most of the proposed design of this study (e.g. inclusion and exclusion criteria, duration of the study), the clinical relevance of the primary criterion is debatable. Even if oedema reduction as primary variable can be considered a reliable quantitative primary end-point to evaluate one of the pharmacological effects of *Ruscus aculeatus*, the clinical relevance of this primary variable is questionable. According to the assessor's opinion, improvement in subjective symptoms such as sensation of heaviness or tiredness, tingling or pain should be of more clinical relevance. As stated by the author himself, any reduction of oedema is only regarded as clinically relevant if it is accompanied by an improvement in patient's quality of life.

Thus, despite significant results regarding the primary variable and the positive correlation shown between leg oedema and all the subjective symptoms except pain, the clinical relevance of these results remains questionable. Indeed, the positive effect relative to the subjective symptoms is very limited. The difference between both groups for the subjective symptoms "tingling sensation" and "pain" are not statistically significant and the significance of the difference for the two other subjective symptoms "heaviness and tiredness" and "sensation of tension" is debatable taking into account the multiplicity of the analyses. Moreover, the treatment response measured by the disease specific questionnaire on the quality of life appeared negative at the end of this study as results on this questionnaire did not reveal any changes in both arms).

In conclusion, the credit that we can attribute to this study is to have tested a *Ruscus* extract alone. The posology of the extract used in the study is in adequacy with the one recommended by the available monographs i.e. a daily dose equivalent to an amount of 7 to 11 mg of total ruscogenins. However, the level of evidence of *R. aculeatus* extract efficacy in relieving symptoms of chronic venous insufficiency given by this study is low. It has to be noted that the efficacy was not evaluated in men. Other studies to confirm these results are deemed necessary. Evaluation of a sustained efficacy over a longer period (up to 1 year) has not been studied.

Due the lack of consistency of this single clinical study, the Rapporteur considers that the requirements for considering the well-established use of the *Ruscus aculeatus* rhizome 'dry extract (15-20:1) extraction solvent, methanol 60% (V/V)', are not fulfilled.

Other Clinical Efficacy Data

- Human pharmacological studies

Additional studies (Rudofsky and Nobbe, 1982; Rudofsky et al., 1989) were provided in addition to the only clinical study to sustain efficacy results. Accessors are of the opinion that results, despite the fact that they demonstrate an increase of the tonus of the venous wall, are of limited interest as *Ruscus aculeatus* extract was not tested alone but in combination with other substances. It is therefore difficult to differentiate a specific effect of *Ruscus aculeatus* extract from the effect of the combinations tested.

- Meta-analysis

Clinical data for *Ruscus aculeatus* have also been published in a meta-analysis that included 20 placebo-controlled randomised double-blind studies. However, the conclusions of the meta-analysis by Boyle et al. (2003) cannot be taken into account. Indeed, this publication is related to a preparation

which contains *Ruscus aculeatus* extract (150 mg per capsule), hesperidine chalcone (150 mg) and ascorbic acid (100 mg) and not to *R. aculeatus* extract alone; this demonstration of the clinical efficacy cannot be attributed to *Ruscus*.

- - International Consensus Symposium

The International Consensus symposium, held during the 13th Congress of European Society for Clinical Hemorheology (ESCH) in Siena (Italy), 2005, concluded that vaso-active drugs (VAD) are effective and may be applied in chronic venous disease (CVD), when symptomatic, at any class of CEAP. The Consensus statement of the international experts also declares, that "in some cases VAD may replace compression and/or complement its effects". The experts classified, based on the available data, both the horse chestnut extract and *R. aculeatus* extract to "Grade B" of their recommendation explained in the statement published by Ramelet *et al.* (2005) Despite this conclusion, assessors consider that the efficacy of *Ruscus aculeatus* to relieve symptoms of chronic venous insufficiency is not demonstrated due to the lack of relevance, for this procedure, of the study and the meta-analysis.

- Comparative study with another herbal:

Lancet 1996; 347:292-4.

Comparison with horse chestnut extract, study "Comparison of leg compression stocking and oral horse-chestnut seed extract therapy in patients with chronic venous insufficiency".

Results of a randomised placebo-controlled three-armed study performed with horse chestnut extract were published by Diehm *et al.* (1996). This study demonstrated that on the criterion oedema protection (leg volume reduction), results are comparable between horse chestnut extract, and leg compression stockings.

However, this extrapolation of results to *Ruscus aculeatus* extract is not adequate. This indirect comparison is hazardous and cannot be accepted as no direct comparison has been performed between *Ruscus aculeatus* extract and leg compression stockings.

4.2.2.2. Haemorrhoids

No clinical studies are available in the treatment of haemorrhoids with *Ruscus aculeatus* alone. Most of the studies are performed with a *Ruscus* combination with various flavonoids. One cannot rule out a positive synergistic effect of the combinations. Thus the proper efficacy of *R. aculeatus* is difficult to assess (Abascal and Yarnall, 2005).

Experiments on single cases: patients treated with a 10% hydro-alcoholic (alcohol 30°) *Ruscus*. extract given *per os* were reported (Caujolle *et al.*, 1953). Over 11 cases followed during a few months, 10 cases showed an improvement of the symptoms whereas in one case *R. aculeatus* extract was ineffective. However, this later patient was suffering from haemorrhoids associated with haemorrhage. Another successful case corresponding to a pregnant women treated locally with suppository was reported.

In addition, the author reported 13 out of 15 observations from rural physicians corresponding to patients treated either *per os* or locally using suppository.

Assessor's comments:

Despite the identified R. aculeatus pharmacological effects i.e. enhancement effect on venotonicity, not well-conducted studies are available with Ruscus aculeatus alone.

Even if there is evidence to suggest that R. aculeatus extract is effective in relieving symptoms of haemorrhoids, this evidence is of very low level. Further researches in that field are needed. However, due to the long-term use of R. aculeatus extract-containing products in this indication, R. aculeatus can be considered as traditional herbal medicinal product for symptomatic relief of itching and burning associated with haemorrhoids.

4.2.2.3. Orthostatic hypotension

One case of profound refractory orthostatic hypotension treated for longer than 2 years has been reported (Redman, 2000). The author recommends one "standard" 470 mg capsule of *Ruscus aculeatus* every hour from waking till evening until blood pressure is high enough that it is no longer needed. This corresponds to doubling the normal "recommended" dose of two capsules three times daily. It has to be noted that the author recommends even so combining the intake of *Ruscus* with non-pharmacological measures and other natural products medicines.

Assessor's comments:

According to the author, with regard to its identified pharmacological properties R. aculeatus application and extension to the treatment of orthostatic hypotension is obvious. In the assessor's opinion this application should be clearly better documented. To date, with regard to the available data, no recommendation can be given.

4.2.2.4. Diabetic retinopathy

One study involving *R. aculeatus* alone has been found in the literature (Archimowicz-Cyrylowska *et al.*, 1996).

The study was carried out with 60 patients (32 women, 28 men), aged from 20-75 years, mostly suffering from non-insulin dependent diabetes mellitus (type II) for 1-27 years, characterised by non-proliferative diabetic retinopathy. They were randomly assigned to three equal groups: Group I-treated with troxerutin (Venoruton Zygma, GmbH) 1 tablet containing 0.5 g of O-(beta-hydroxyethyl)-rutoside 2 times a day; Group II-treated with 1 capsule containing 0.0375 g of *R. aculeatus* extract 2 times a day, and Group III-treated with 2 tablets of pressed buckwheat herb (each tablet containing 0.5 g *Fagopyrum esculentum* herb and 0.03 troxerutin, 3 times a day. During the study period of 3 months, all subjects remained on stable diabetic diet, unchanged hypoglycaemic medication for the period of treatment. If administered earlier, any hypolipaemic medication was withdrawn at least 4 weeks before the onset of the study.

At the beginning, as well as on the last day of the investigation each patient was subjected to an ophthalmological examination and clinical biochemistry. The oscillating potentials of the electroretinogram were also recorded.

Results:

Group I, medicated with troxerutin, was characterised by a decrease in amplitude of oscillating potentials by 21% considering both eyes. In contrast, in group II and group III, treated with *R. aculeatus* and buckwheat herb, an increase in amplitude of oscillating potentials (by 15% and by 18% respectively) was observed. The changes in the amplitude detected were statistically insignificant when compared with the initial values.

In all patients treated for 3 months with troxerutin, *Ruscus* and buckwheat herb preparations, a slight statistically insignificant increase in visual acuity was observed.

Examination of the anterior segment of the eyeball after 3 months of pharmacotherapy did not show any differences when compared with the initial picture in all the groups evaluated.

A regression of changes located in fundus of eye was demonstrated in 27.8% while a progression in 5.6% of patients treated with troxerutin was observed. Evaluation of the fundus of the eye in group II (*R. aculeatus* extract) revealed a quite distinct improvement in 23.1% of patients and no cases with progression, while in patients receiving buckwheat herb (group III) an improvement was demonstrated in 26.7% and a deterioration in 3.3% of the examined diabetics.

Mean blood serum concentrations of glucose significantly decreased by 12.7% in the troxerutin treated group, by 10.6% in the *Ruscus* group and by 15.1% in subjects medicated with buckwheat herb. Similarly, concentrations of glycosylated haemoglobin were lower after the 3-month period of treatment in all groups studied.

Assessor's comments:

The design of the study is not acceptable for many reasons, such as:

- *The patients are not adequately defined: the stage of the non-proliferative diabetic retinopathy is not given; no reference is made to an approved European diabetic retinopathy classification such as the ETDRS classification; neither baseline blood pressures nor baseline glycosylated haemoglobin concentrations are given.*
- *With regard to the pathology, the duration of the study is too short.*
- *Only 20 patients were enrolled in each group of treatment.*
- *With regard to the pathology, the choice of the comparators is not justified and cannot be considered as relevant. Moreover, there is no comparison with a placebo.*

Thus, the design of this study is not relevant and cannot be taken into account.

Table 6: Clinical studies on humans, in chronic venous insufficiency

Type	Study	Test Product(s)	Number of subjects	Type of subjects	Outcomes	Statistical analysis	Clinical relevance
Efficacy and safety of a <i>R. aculeatus</i> extract in chronic venous insufficiency Vanscheidt <i>et al.</i> , 2002	Multicenter double-blind, randomised, placebo-controlled	<i>Ruscus</i> extract Active treatment and placebo were taken twice daily (morning and evening) orally with some fluid One capsule active treatment contained 36.0-37.5 mg dry extract from <i>Ruscus aculeatus</i> rhizome with a drug extract ratio of 15-20: 1 (solvent methanol 60%) corresponding to an amount of around 4.5 mg of total ruscogenins 12 weeks	167 women screened, of whom 166 were randomised and included in the study	166 women suffering chronic venous insufficiency (Widmer classification grades I and II, CEAP 18 had insufficient data and were excluded from the efficacy analysis 3-5)	Primary variable: change in foot and lower leg volume, measured as the area under the curve of volume-changes over the 12-weeks treatment-time of (AUB0-12, area under baseline) Secondary variables: changes in leg volume; changes in subjective symptoms Evaluated after 12 weeks of treatments The subjective symptoms: tiredness and heaviness, sensation of tension, tingling sensation, and pain were assessed at each visit before the volume measurements	AUB0-12, median values: 656 ml per day for the <i>Ruscus</i> extract group; and 175 ml per day for placebo Significant treatment contrast (p<0.001) Subjective symptoms: (p=0.0498).	The level of evidence of efficacy in relieving symptoms of chronic venous insufficiency is low. It has to be noted that the efficacy was not evaluated in men. Evaluation of a sustained efficacy over a longer period has not been studied

Table 7: Clinical studies on humans, in chronic diabetic retinopathy

Type	Study	Test Product(s)	Number of subjects	Type of subjects	Outcomes	Statistical analysis	Clinical relevance
Efficacy and safety of a <i>Ruscus extract</i> in chronic diabetic retinopathy (Archimowicz-Cyrylowska <i>et al.</i> , 1996)	3 arms randomised	Group I - treated with troxerutin 1 tablet containing 0.5 g of 0-(beta-hydroxyethyl)-rutoside, 2 times a day Group II - treated with 1 capsule containing 0.0375 g of <i>Ruscus extract</i> (Fagorutin- <i>Ruscus</i> , Fink GmbH), 2 times a day Group III – treated with 2 tablets of pressed buckwheat herb (each tablet containing 0.5g <i>Fagopyrum esculentum</i> herb and 0.03 troxerutin, 3 times a day During the study period of 3 months, all subjects remained on stable diabetic diet, unchanged hypoglycaemic medication for the period of treatment	60 patients (32 women, 28 men), aged from 20-75 years, mostly suffering from non-insulin dependent diabetes mellitus (type II) for 1-27 years, characterised by non-proliferative diabetic retinopathy		Ophthalmologic examination Clinical biochemistry Oscillating potentials of the electro-retinogram	Group I, decrease in amplitude of oscillating potentials by 21% considering both eyes Regression of changes in fundus of eye in 27.8%; Progression in 5.6% Group II increase in amplitude of oscillating potentials (by 15%) Improvement of the fundus of the eye in 23.1% of patients; no cases with progression Group III, increase in amplitude of oscillating potentials (by 18%) improvements demonstrated in 26.7% and a deterioration in 3.3% The changes in the amplitude of oscillating potentials detected were	The design of the study is not acceptable: The patients are not adequately defined: the stage of the non-proliferative diabetic retinopathy is not given; no reference is made to an approved European diabetic retinopathy classification such as the ETDRS classification; neither baseline blood pressures nor baseline glycosylated haemoglobin concentrations are given. With regard to the pathology, the duration of the study is too short. Only 20 patients were

Type	Study	Test Product(s)	Number of subjects	Type of subjects	Outcomes	Statistical analysis	Clinical relevance
						statistically insignificant	<p>enrolled in each group</p> <p>With regard to the pathology, the choice of the comparators is not justified and cannot be considered as relevant</p> <p>There is no comparison with a placebo</p>

4.3. Clinical studies in special populations (e.g. elderly and children)

Elderly

According to the provided literature, no clinical studies have been conducted with *Ruscus* extract alone in elderly.

Children

According to the provided literature, no clinical studies have been conducted with *Ruscus* extract alone in children.

Pregnancy

The use of *Ruscus* extract alone has not been evaluated in pregnant women.

4.4. Overall conclusions on clinical pharmacology and efficacy

To date, the clinical data on *Ruscus aculeatus* extract that can be taken into consideration are limited to only one publication of a randomised placebo-controlled study performed in patients with chronic venous insufficiency (Vanscheidt *et al.*, 2002). The results obtained from this clinical study suggest an efficacy in the relief of symptoms such as "heaviness and tiredness" and "sensation of tension" in patients suffering from chronic venous insufficiency. However, in the opinion of the assessor, the provided evidence is insufficient to implement the *Ruscus aculeatus* monograph for a well-established use in relieving symptoms of chronic venous insufficiency.

In the treatment of haemorrhoids, no clinical data are available with *Ruscus aculeatus*; only pharmacological effects, data provided by studies with *Ruscus aculeatus* in combination and the long-term use suggest that *Ruscus* extract is effective to relieve symptoms of haemorrhoids.

There is no sufficient data to sustain the indication of *Ruscus aculeatus* extract in orthostatic hypotension and in diabetic retinopathy.

5. Clinical Safety/Pharmacovigilance

5.1. Overview of toxicological/safety data from clinical trials in humans

No data available.

5.2. Patient exposure

Aside from market presence and data from studies, there are no concrete data concerning patient exposure.

5.3. Adverse events, serious adverse events and deaths

Results from the study published by Vanscheidt *et al.* (2002)

This study is a multi-centre, double-blind, randomised, placebo-controlled trial with women suffering from chronic venous insufficiency to investigate the efficacy and safety of an extract of *Ruscus aculeatus* rhizome. Randomisation was carried out after a 2-week run-in phase at visit 2. During the run-in period, all patients received placebo. The following treatment phase with either *R. aculeatus* extract or placebo in the two parallel groups lasted 12 weeks. Checkpoints were scheduled after 4, 8 and 12 weeks of treatment. The daily dosage of the *R. aculeatus* extract (72-75 mg dry extract from

butcher's broom rhizome) was chosen according to the German monograph. 166 Patients from 30 to 89 years-old were included. 37 patients experienced one or more treatment emergent adverse events:

- 17 patients experienced 22 adverse events in the *R. aculeatus* extract group, including 2 cases of calf cramps;
- 20 patients experienced 26 adverse events in the placebo group, including 4 cases of calf cramps.

The tolerability was assessed as very good in 76.8% in the *R. aculeatus* extract group versus 78.8% in the placebo group, good in 23.2% versus 20.0% and moderate in 0% versus 1.3%.

Assessor's comments:

37 patients experienced adverse effects, of which 17 were in the R. aculeatus extract group. No information is available on these cases regarding the nature and the seriousness, except for 6 of them (2 in the Ruscus extract group and 4 in the placebo group), which were calf cramps. No conclusion with regards to the safety of this extract can be drawn from these data.

Data from the literature

Three publications and one abstract presented during a French conference regarding the safety of *Ruscus aculeatus* have been identified.

Valnet-Rabier *et al.*, 2004. The abstract reports one case of collagenous colitis which occurred in one 52 years-old female patient. This patient initiated *R. aculeatus* therapy about 10 months before the diagnosis, which has been histologically confirmed. At the onset of the colitis, other fluid extracts were taken by the patient, but none had a known colorectal toxicity.

Assessor's comments:

Rapporteur's comment: Due the lack of details no consequent conclusions could be drawn from this report.

Landa *et al.*, 1990. This publication reports one case of papulo-erythematous eruption of both legs, that spread within a few days to the entire skin, with oedema of the eyelids in a 30 years-old pregnant female patient, after the application of a vasoconstrictor cream for the treatment of varices. Patch tests revealed positive results to *R. aculeatus* and thimoresal, 2 ingredients of the cream.

Elbadir *et al.*, 1998. Eight cases of contact allergy were collected from 1986 to 1995, in patients receiving ruscogenins containing medications. Ruscogenins are components of *R. aculeatus*. These 8 cases involve 6 women and 2 men aged from 28 to 55 years-old who experienced eczema at the application site, after the use of a topic medication (7 cases) or a cosmetic cream (1 case). Prick tests or patch tests were performed in 7 patients and all were positive for ruscogenins or *R. aculeatus*.

Ramirez-Hernandez, 2006. One case of perianal eczema in one 34 years-old patient following the local application of an antihemorrhoidal cream. The outcome was favourable after therapy withdrawal. Several months later, the patient developed a generalised eczematous cutaneous eruption one day after the application of an anticellulitis product on the lower limbs. Patch tests for both creams revealed a positive reaction to ruscogenins.

Assessor's comments:

According to these data, the local application of R. aculeatus or ruscogenins has been associated with allergic reactions, mainly represented by contact eczema. In one case, the event spread to the entire skin. In all cases, the outcome was favourable after therapy withdrawal and administration of corticoids.

Sadarmin and Timperley, 2013. One case of diabetic ketoacidosis in a 39-year-old diabetic woman with poor glucose control and glycosylated hemoglobin of 11.9% was reported following the consumption (for five days) of an herbal medicine containing *Ruscus aculeatus* (no details on the preparation or posology).

Rapporteur's comment:

Due the lack of details no consequent conclusions could be drawn from this report.

Rapporteur's overall comment:

*In case of hypersensitivity to ruscogenins or to the plant the use of *R. aculeatus* rhizome preparations is contraindicated.*

5.4. Laboratory findings

The review of the PSURs (Periodic safety update reports) regarding medicinal products containing *R. aculeatus* (oral and cutaneous use) did not allow identifying other safety issues. *Ruscus* containing creams may potentially be associated with allergic reactions, mainly eczemas; medicinal products for oral use containing *R. aculeatus* in combination with other substances may cause diarrhoea and lymphocytic colitis.

5.5. Safety in special populations and situations

No data are available.

5.5.1. Use in children and adolescents

The use in children and adolescents is not recommended due the lack of data.

5.5.2. Contraindications

No data are available.

Hypersensitivity to the active substance.

5.5.3. Special Warnings and precautions for use

If some conditions, inherent to venous circulatory disturbances, such as inflammation of the skin or subcutaneous induration, ulcers, sudden swelling of one or both legs, cardiac or renal insufficiency, occur a doctor should be consulted.

If rectal bleeding occurs, associated with haemorrhoids, a doctor should be consulted.

5.5.4. Drug interactions and other forms of interaction

None reported

5.5.5. Fertility, pregnancy and lactation

In an open study involving 9 pregnant women, 3 of the pregnant women applied 2 to 3 g of a *Ruscus* containing cream (100 mg of cream contains 1.6 g *R. aculeatus* extract and 1.6 g *Mellilot* extract) twice daily during the 3rd trimester of pregnancy. No embryotoxic effects were noted by the author (Berg, 1991).

Assessor's comments:

The study was performed with a combination of R. aculeatus extract and another product, thus a conclusion on the safety of the R. aculeatus extract alone cannot be drawn. Furthermore, as the combination has been administered during the third trimester of pregnancy, the embryotoxicity cannot be ruled out, only the foetotoxicity is addressed.

In an open study, 20 pregnant women have been enrolled (Baudet *et al.*, 1991). The women have taken two capsules per day of a combination of R. aculeatus extract 150 mg, trimethylhesperidin chalcone 150 mg and ascorbic acid 100 mg after 21 to 24 weeks of amenorrhea. Foetal development was measured through the pulse Doppler method of the cord. The authors conclude that this test shows "an absolute harmlessness for the infant, assessed with the usual clinical and ultrasonographic criteria of pregnancy surveillance, with Doppler's velocimetry at the level of the umbilical artery and with the state of the infant and the anatomopathologic aspect of the placenta after birth."

Assessor's comments:

As for the prior study, this study was performed with a combination of R. aculeatus extract and other products. The combination administered from the second trimester of pregnancy has shown neither foetotoxic effect nor harmful effect for the new born.

In a review of the herbal treatments for haemorrhoids, some authors concluded that the available studies in pregnant women treated with a *R. aculeatus* combination do not establish the safety of *R. aculeatus* in pregnancy conclusively (Abascal and Yarnall, 2005).

Assessor's comments:

No data are available with R. aculeatus extract alone. Even considering R. aculeatus combinations, data are too limited to allow any recommendations.

Lactation

As there are no clinical or animal data available on the use of *R. aculeatus* extract during lactation and due to the potential harmful effect on the breast fed new born, particularly with regard to gastrointestinal disorders, the use of *R. aculeatus* extract should be avoided during the lactation.

There are no data on fertility available.

5.5.6. Overdose

No data are available.

5.5.7. Effects on ability to drive or operate machinery or impairment of mental ability

No data are available.

5.5.8. Safety in other special situations

No data are available.

5.6. Overall conclusions on clinical safety

No conclusion with regards to the safety of a *Ruscus* extract can be drawn from the study published by Vanscheidt *et al.* (2002). The data are too scarce and insufficient. No information is available on the

adverse effects regarding the nature and the seriousness, excepted for six of them (2 in the *R. aculeatus* extract group and 4 in the placebo group), which were calf cramps.

Data from the literature highlight two kinds of adverse effects which have been associated with the intake of *R. aculeatus* or ruscogenins containing products. The topical forms have been associated with contact dermatitis. Although the patient received medicinal products containing multiple substances, prick tests and patch tests allowed confirming, for the topical use, the responsibility of *Ruscus/ruscogenins* in the occurrence of the allergic reaction.

The second well identified risk concerns to the development of diarrhoea/lymphocytic colitis after oral administration of medicinal products containing *Ruscus aculeatus*. The literature data seem to be supported by spontaneous reports regarding different medicinal products and collected data in the PSURs. However in these medicinal products are combinations of *Ruscus aculeatus* and other substances and no definitive conclusion can be drawn regarding the effects of *Ruscus aculeatus* single preparations.

From the available studies, a conclusion on the safety of the use of *R. aculeatus* extract alone during pregnancy cannot be drawn. The women were exposed to a mixture containing *R. aculeatus* extract from the second trimester of pregnancy. Thus, the only conclusions which can be drawn relates to the foetotoxicity or the new born effects. For a very limited number of pregnant women (23) no foetotoxic effect appeared to date but complementary data are necessary to conclusively establish the safety of the *R. aculeatus* extract alone during the latter pregnancy. No data on exposure during the first trimester of pregnancy are available. So, no conclusion can be drawn about the teratogenic potential of the *R. aculeatus* extract alone.

As there are no clinical or animal data available on the use of *R. aculeatus* extract during lactation. The use of *R. aculeatus* extract during lactation is not recommended.

6. Overall conclusions (benefit-risk assessment)

Non-clinical aspects

Pharmacology

Primary pharmacodynamics studies performed *in vitro* and *in vivo* using various experimental models showed that *R. aculeatus* extract possess a contractile activity on veins. This activity is mediated by stimulation of the α -adrenergic system. *In vitro* mechanistic studies showed that direct activation of post-junctional α 1- and α 2-adrenergic receptors, and stimulation of the release of norepinephrine from adrenergic nerve endings were involved. Although this effect does not appear to be clearly influenced by the hormonal status (oestrogens, progesterone), it seems potentiated by temperature.

In *in vivo* studies, this vasoconstricting activity was shown after intravenous and oral routes; in the hamster cheek pouch model, local application of the extract (i.e. in the superfusate) was also effective. It should be noted that only one study was conducted by the oral route: at the level of hamster cheek pouch microcirculation, the dose of 150 mg extract per kg per day administered for 28 days induced venular constriction (internal diameter decreased by 30%) and arteriolar dilation (internal diameter increased by 37%) without any impact on the mean arterial blood pressure, the latter effect being attributed to liberation of endothelium-derived relaxing factors on the arteriolar side.

Similarly, other primary pharmacodynamics studies showed that *R. aculeatus* extract exerts a contractile effect on lymphatic vessels in anaesthetised dogs at 2 and 5 mg/kg administered intravenously. A rise in oncotic pressure suggested a favourable effect on oedema. This was confirmed in a feline model of ethacrynic acid-induced oedema. The effective dosage amounted to 20

mg/kg by intravenous route; and 10-20 times higher by oral route. However, after subchronic administration (4-6 days), the oral effective dosage decreased to reach 20-40 mg/kg per day. The same study showed that ruscogenin was also effective, but that other components of the extract were involved to obtain maximal activity.

Due to the mechanisms underlying the effect of *R. aculeatus* extract, pharmacodynamic drug-drug interactions with any drug potentiating or antagonising the α -adrenergic system are plausible. However, reports on cases of drug interactions were not found.

Considering the pharmacological profile of *R. aculeatus* extract, i.e. stimulation of α -adrenergic system, the lack of a safety pharmacology study evaluating its potential effects on the cardiovascular function gives cause for concern. No toxicology study evaluating this endpoint is available. In the studies performed in the hamster cheek pouch model, the mean arterial blood pressure was not modified after IV administration of 5 mg/kg *R. aculeatus* extract, and oral administration at the dose of 150 mg extract perkg.

Pharmacokinetics

Available pharmacokinetics studies should not be taken into account for regulatory purposes because they are endowed with major bias precluding a full confidence in the results obtained.

Toxicology

Extracts used in two acute toxicity studies were obtained by ethanolic extraction. Their characteristics (content in various compounds, e.g. ruscogenin and neoruscogenin) remain unknown. Potential variability compared to the extract(s) intended for therapeutic cannot be evaluated. In dogs, a mean intravenous LD₀ of 1.20 g/kg was measured, while the oral LD₅₀ of mice reached 25-34 ml/kg with another extract. The authors attribute the cardiovascular findings observed in dogs at high doses (decreased frequency of cardiac contractions, decreased blood pressure) to be secondary to alteration of respiratory centres. In rats and mice, death occurred by respiratory failure too. No safety pharmacology study is available on cardiovascular and respiratory systems.

No other toxicity studies are available. The ESCOP monograph reports repeat-dose toxicity and reprotoxicity studies performed in rabbits and rats, respectively. However, these studies remain unpublished so that the information available is very sparse. Therefore, they cannot be taken into consideration for safety evaluation.

The lack of adequately conducted genotoxicity and embryo-foetal toxicity studies precludes the listing of *R. aculeatus*. Additionally, as no long-term study is available, the carcinogenic risk cannot be estimated.

Clinical aspects

Pharmacology

On the basis of publications, the quality of the two available pharmacodynamics studies cannot be evaluated. For example, the characteristics of the patients are incomplete as well as the design of the studies. The statistical analysis is not given. Moreover, the characteristics of the *R. aculeatus* extract are not specified. However, the findings corroborate the preclinical pharmacological properties that acknowledge to *R. aculeatus* extract an alpha-adrenergic effect and thus a venous vasoconstrictive effect that account for a reduction in volume of blood stored in the veins and for a stimulating effect on the lymphatic drainage. These two pharmacological properties assume a positive effect in patients suffering from chronic venous insufficiency.

Dose-effect studies are missing which preclude from justifying the dosage regimen used in the clinical studies.

Other methods of functional exploration could have been used to evaluate the effect on veins (e.g. Doppler).

Pharmacokinetics

Available pharmacokinetics data are too scarce.

Efficacy

The results obtained from Vanscheidt *et al.*, (2002) suggest efficacy in the relief of symptoms such as “heaviness and tiredness” and “sensation of tension” in patients suffering from chronic venous insufficiency. As stated by the author himself, any reduction of oedema is only regarded as clinically relevant if it is accompanied by an improvement in patient’s quality of life. However, the treatment response measured by the disease specific questionnaire on the quality of life appeared negative at the end of this study. Finally, the provided evidence is insufficient to accept *R. aculeatus* for a well-established use monograph in relieving symptoms of chronic venous insufficiency.

Moreover, the French National Authority for Health reassessed the benefit of all ‘veinotonics’ in the treatment of chronic venous insufficiency. All the ‘veinotonics’ with marketing authorisation in France such as *Ruscus extract and hesperidine methylchalcone*, diosmin, troxerutin had been studied. The conclusions of the Authority were that the efficacy of all the medicines was minor and the evidence given to demonstrate the efficacy was poor.

In the treatment of haemorrhoids, no clinical data are available with *R. aculeatus* alone; only pharmacological effects, data provided by studies with *Ruscus* in combination and the long-term use suggest that *R. aculeatus* extract is effective to relieve symptoms of haemorrhoids.

Based on the available data, the monograph information should remain limited to the traditional use in subjective symptoms of chronic venous insufficiency such as sensation of heavy legs and in symptomatic relief of itching and burning associated with haemorrhoids.

The daily dosage recommended for the *Ruscus aculeatus* containing products available on the market is considered as acceptable as it is in line with dosages reported in literature.

The use in children and adolescents has not been established due to lack of adequate data.

Safety during pregnancy and lactation has not been established. In the absence of sufficient data, the use during pregnancy and lactation is not recommended.

Safety

No conclusion with regards to the safety of *Ruscus aculeatus* can be drawn since data are too scarce and insufficient. Data from the literature highlight two adverse effects which have been associated with the intake of *R. aculeatus* or ruscogenins containing products. *R. aculeatus* containing products for external use have been associated with contact dermatitis. Although the patient received medicinal products containing multiple substances, prick tests and patch tests allowed confirming, for the topical use, the responsibility of *Ruscus*/ruscogenins in the occurrence of the allergic reaction. However, the topical use is not considered in the EU monograph.

The second identified risk concerns the oral administration of some *Ruscus aculeatus* containing products associated with gastro-intestinal complaints, diarrhea/lymphocytic colitis. The literature data seem to be supported by spontaneous reports and collected data in the PSURs, however respecting to

combinations of *Ruscus aculeatus* and other substances. Therefore, no definitive conclusion can be drawn regarding the effects of *Ruscus aculeatus* single preparations.

The relevance of preclinical data on the cardiovascular system has not been confirmed by clinical data.

From the available studies, a conclusion on the safety of the use of *R. aculeatus* extract alone during pregnancy cannot be drawn. No conclusion can be drawn about the teratogenic potential of the *R. aculeatus* extract alone. The use of *R. aculeatus* extract should be not recommended during pregnancy.

As there are no clinical or animal data available on the use of *R. aculeatus* extract during lactation and due to the potential harmful effect on the breast fed new born, particularly with regard to gastrointestinal disorders, the use of *R. aculeatus* extract should be avoided during the lactation.

Rapporteurs overall conclusions

Based on the available data, the monograph information should remain limited to the following indications and herbal preparations:

Indication 1)

Traditional herbal medicinal product to relieve symptoms of discomfort and heaviness of legs related to minor venous circulatory disturbances.

- a) Powdered herbal substance
- b) Dry extract DER (2.5-6.5:1); extraction solvent: water)
- c) Dry extract (DER 5.0-8.5:1), extraction solvent, ethanol 80% (V/V)
- d) Dry extract DER (6-9:1), extraction solvent, ethanol 96% (V/V)

Indication 2)

Traditional herbal medicinal product for symptomatic relief of itching and burning associated with haemorrhoids, after serious conditions have been excluded by a medical doctor.

- a) Powdered herbal substance
- b) Dry extract DER (2.5-6.5:1); extraction solvent: water)
- c) Dry extract (DER 5.0-8.5:1), extraction solvent, ethanol 80% (V/V)

For both indications if the symptoms persist for more than 2 weeks during the use of the medicinal product, a doctor or a qualified health care practitioner should be consulted.

The use should be limited to adults.

Safety during pregnancy and lactation has not been established. In the absence of sufficient data, the use during pregnancy and lactation is not recommended.

Typical analytical marker(s) are ruscogenins (mixture of neoruscogenin and ruscogenin).

A European Union list entry is not supported due to lack of adequate data on genotoxicity.

Annex

List of references