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**COMMITTEE ON HERBAL MEDICINAL PRODUCTS
(HMPC)**

DRAFT

COMMUNITY HERBAL MONOGRAPH ON *RUSCUS ACULEATUS* L., RHIZOMA

DISCUSSION IN WORKING PARTY ON COMMUNITY MONOGRAPHS AND COMMUNITY LIST (MLWP)	July 2007 September 2007
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REDISCUSSION IN WORKING PARTY ON COMMUNITY MONOGRAPHS AND COMMUNITY LIST (MLWP)	
ADOPTION BY HMPC	

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KEYWORDS

Herbal medicinal products; HMPC; Community herbal monographs; traditional use; *Ruscus aculeatus* L.; Rusci rhizoma; butcher's broom

COMMUNITY HERBAL MONOGRAPH ON *RUSCUS ACULEATUS* L., RHIZOMA

1. NAME OF THE MEDICINAL PRODUCT

To be specified for the individual finished product.

2. QUALITATIVE AND QUANTITATIVE COMPOSITION^{1, 2}

<u>Well-established use</u>	<u>Traditional use</u>
	<p>With regard to the registration application of Article 16d(1) of Directive 2001/83/EC as amended</p> <p>i) Herbal substance <i>Ruscus aculeatus</i> L., rhizoma (butcher's broom)</p> <p>ii) Herbal preparations³ Dried powdered root Dry extract (4.5-6.5 : 1 ; water) Dry extract (5-8.5 : 1 ; 80% V/V ethanol) Dry extract (6-9 : 1 ; 96 % V/V ethanol) Dry extract (15-20 :1; 60% V/V methanol)</p>

3. PHARMACEUTICAL FORM

<u>Well-established use</u>	<u>Traditional use</u>
	<p>Herbal substance or herbal preparation in solid dosage forms for oral use.</p> <p>The pharmaceutical form should be described by the European Pharmacopoeia full standard term.</p>

¹ The material complies with the Ph. Eur. monograph (ref. 01/2005 : 1847)

² The declaration of the active substance(s) for an individual finished product should be in accordance with relevant herbal quality guidance.

³ Quantified for ruscogenins as determined by the total amount of ruscogenin and neoruscogenin

4. CLINICAL PARTICULARS

4.1. Therapeutic indications

<u>Well-established use</u>	<u>Traditional use</u> Herbal medicinal product traditionally used to relieve symptoms of heavy legs. The product is a traditional herbal medicinal product for use in specified indication exclusively based upon long-standing use.
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4.2. Posology and method of administration

<u>Well-established use</u>	<u>Traditional use</u> Posology <i>Adults, elderly</i> Dried powdered root: 350 mg 3 times daily Dry extract (4.5-6.5 :1 ; water): 200 mg 2 times daily Dry extract (5-8.5 :1 ; 80 % V/V ethanol): 86 mg 1 to 2 times daily Dry extract (6-9 : 1 ; 96 % V/V ethanol): 45 mg 2 times daily Dry extract (15-20:1; 60% V/V methanol): 37 mg 2 times daily Recommendations given for dried powdered root or dry extracts (7-11 mg daily) of quantified ruscogenins as determined by the total amount of ruscogenin and neoruscogenin. <i>Children, adolescents</i> There is no relevant indication for children and adolescents. Duration of use 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. Method of administration Oral use.
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4.3. Contraindications

<u>Well-established use</u>	<u>Traditional use</u> Hypersensitivity to the active substance.
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4.4. Special warnings and precautions for use

<u>Well-established use</u>	<u>Traditional use</u> If there is inflammation of the skin or subcutaneous induration, ulcers, sudden swelling of one or both legs, cardiac or renal insufficiency, a doctor should be consulted.
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4.5. Interactions with other medicinal products and other forms of interaction

<u>Well-established use</u>	<u>Traditional use</u> None reported.
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4.6. Pregnancy and lactation

<u>Well-established use</u>	<u>Traditional use</u> Safety during pregnancy and lactation has not been established. In the absence of sufficient data, the use during pregnancy and lactation is not recommended.
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4.7. Effects on ability to drive and use machines

<u>Well-established use</u>	<u>Traditional use</u> No studies on the effect on the ability to drive and use machines have been performed.
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4.8. Undesirable effects

<u>Well-established use</u>	<u>Traditional use</u> Nausea, gastrointestinal complaints, diarrhea, lymphocytic colitis may occur. The frequency is not known. If other adverse reactions not mentioned above occur, a doctor or a qualified health care practitioner should be consulted.
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4.9. Overdose

<u>Well-established use</u>	<u>Traditional use</u> No case of overdose has been reported.
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5. PHARMACOLOGICAL PROPERTIES

5.1. Pharmacodynamic properties

<u>Well-established use</u>	<u>Traditional use</u> Not required as per Article 16c(1)(a)(iii) of Directive 2001/83/EC as amended.
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5.2. Pharmacokinetic properties

<u>Well-established use</u>	<u>Traditional use</u> Not required as per Article 16c(1)(a)(iii) of Directive 2001/83/EC as amended.
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5.3. Preclinical safety data

<u>Well-established use</u>	<u>Traditional use</u> Not required as per Article 16c(1)(a)(iii) of Directive 2001/83/EC as amended, unless necessary for the safe use of the product. Tests on genotoxicity, carcinogenicity, and reproductive toxicity have not been performed.
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6. PHARMACEUTICAL PARTICULARS

<u>Well-established use</u>	<u>Traditional use</u> Not applicable.
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7. DATE OF COMPILATION/LAST REVISION

7 September 2007