

05 June 2018 EMA/HMPC/444244/2015 Committee on Herbal Medicinal Products (HMPC)

European Union herbal monograph on *Pelargonium* sidoides DC and/or *Pelargonium reniforme* Curt., radix

This document was valid from 05 June 2018 until May 2024. It is now superseded by a <u>new version</u> adopted by the HMPC on 29 May 2024 and published on the EMA website.

Initial assessment	
Discussion in Working Party on Community monographs and Community	September 2010
list (MLWP)	November 2010
	January 2011
	March 2011
Adopted by Committee on Herbal Medicinal Products (HMPC) for release	31 March 2011
for consultation	51 Widi CH 2011
End of consultation (deadline for comments).	15 August 2011
Re-discussion in MLWP	September 2011
	November 2011
	January 2012
	May 2012
Adoption by HMPC	20 November 2012
Monograph (EMEA/HMPC/560961/2010)	
AR (EMEA/HMPC/560962/2010)	
List of references (EMEA/HMPC/560963/2010)	
Overview of comments received during the public consultation	
(EMEA/HMPC/748350/2011)>	
HMPC Opinion (EMEA/HMPC/742263/2012)	
First review	
Discussion in MLWP	September 2014
	July 2015
Adopted by HMPC for release for consultation	29 September 2015
End of consultation (deadline for comments ¹)	31 January 2016
Re-discussion in MLWP	April 2016
	May/June 2016
	July 2016

¹ No comments were received during the period of public consultation. Therefore the final monograph is published together with the final assessment report and list of references, without an 'Overview of comments received during the public consultation'.

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			November 2016
			March 2017
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Adoption by HMP	2C		05 June 2018
Keywords Herbal medicinal products; HMPC; Community herbal monographs; traditional		bal monographs; traditional	
use; Pelargonium sidoides DC and/or Pelargonium reniforme Curt., radix;		reniforme Curt., radix;	
	Pelargonii radix; pelargor	nium root	
BG (bălgarski): Пеларгониум, корен LT (lietuvių kalba): Pelargonijų šaknys		elargonijų šaknys	
CS (čeština): pelargoniový kořen		LV (latviešu valoda):	Pelargonijas saknes
DA (dansk): Pela	irgonierod	MT (malti): għerq tal	-ġeranju
DE (Deutsch) · Pa	alargoniumwurzol	NI (nederlands): Cor	ranium

DE (Deutsch): Pelargoniumwurzel	NL (nederlands): Geranium
EL (elliniká): πελαργονίου ρίζα	PL (polski): Korzeń pelargonii
EN (English): pelargonium root	PT (português): pelargónio, raiz
ES (espanol): pelargonio, raíz de	RO (română): rădăcină de muscata
ET (eesti keel): pelargoonijuur	SK (slovenčina): koreň muškátu
FI (suomi): pelargoni, juuri	SL (slovenščina): korenina pelargonije
FR (français): pélargonium (racine de)	SV (svenska): pelargon, rot
HU (magyar): muskátligyökér	IS (íslenska):
IT (italiano): Pelargonio radice	NO (norsk): pelargoniumrot

European Union herbal monograph on Pelargonium sidoides DC and/or Pelargonium reniforme Curt., radix

1. Name of the medicinal product

To be specified for the individual finished product.

2. Qualitative and quantitative composition 2,3

To be specified for the individual finished product.	
2. Qualitative and quantitative	composition ^{2, 3}
Well-established use	Traditional use
	With regard to the registration application of Article 16d(1) of Directive 2001/83/EC
	Pelargonium sidoides DC and/or Pelargonium reniforme Curt., radix (pelargonium root)
	i) Herbal substance
	Not applicable ii) Herbal preparations
	a) Liquid extract (DER 1:8-10), extraction solvent ethanol 11% (m/m)
	 b) Dry extract, (DER 4-25:1), extraction solvent ethanol 11% (m/m)

3. Pharmaceutical form

Well-established use	Traditional use
	Herbal preparations in liquid or solid dosage forms for oral use. The pharmaceutical form should be described by the European Pharmacopoeia full standard term.

4. Clinical particulars

4.1. Therapeutic indications

Well-established use	Traditional use
	Traditional herbal medicinal product for the symptomatic treatment of common cold.
	The product is a traditional herbal medicinal

 ² The material complies with the Ph. Eur. monograph (ref.: 2264).
 ³ The declaration of the active substance(s) for an individual finished product should be in accordance with relevant herbal quality guidance.

Well-established use	Traditional use
	product for use in the specified indication exclusively based upon long-standing use.

4.2. Posology and method of administration

Well-established use	Traditional use
	Posology
	Single dose
	Adolescents over the age of 12 years, adults and
	elderly
	a) Liquid extract: 1.19-1.25 ml, 3 times daily.
	b) Dry extract: 20 mg, 3 times daily.
	Children between 6-12 years
	a) Liquid extract: 0.79-0.83 ml, 3 times daily.
	b) Dry extract: 20 mg, 2 times daily.
	The use in children under 6 years of age is not
	recommended (see section 4.4 'Special warnings and precautions for use').
	Duration of use
	If the symptoms persist longer than 1 week
	during the use of the medicinal product, a doctor
	or a qualified health care practitioner should be consulted.
	Method of administration
	Oral use

4.3. Contraindications

Well-established use	Traditional use
	Hypersensitivity to the active substance(s).

4.4. Special warnings and precautions for use

Well-established use	Traditional use
	The use in children under 6 years of age has not been established due to lack of adequate data.
	Hepatotoxicity and hepatitis cases were reported in association with the administration of the

Well-established use	Traditional use
	medicinal product. In case signs of hepatotoxicity
	occur, the administration of the medicinal product
	should be stopped immediately and a medical
	doctor should be consulted.
	If the symptoms worsen during the use of the
	medicinal product, a doctor or a qualified health
	care practitioner should be consulted.
	For liquid extracts containing ethanol, the
	appropriate labelling for ethanol, taken from the
	'Guideline on excipients in the label and package
	leaflet of medicinal products for human use', must
	be included.

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

Well-established use	Traditional use
	None reported
4.6. Pregnancy and lactation	

Well-established use	Traditional use
	No fertility data available.
	Safety during pregnancy and lactation has not
	been established. In the absence of sufficient
	data, the use during pregnancy and lactation is
	not recommended.

4.7. Effects on ability to drive and use machines

Well-established use	Traditional use
	No studies on the effect on the ability to drive and use machines have been performed.

4.8. Undesirable effects

Well-established use	Traditional use
	Mild gastrointestinal complaints (diarrhea, epigastric discomfort, nausea or vomiting, dysphagia), mild nasal and gingival bleeding and
	allergic reactions have been reported. The

Well-established use	Traditional use
	frequency was very rare.
	Hepatotoxicity has been reported. The frequency is not known.
	If other adverse reactions not mentioned above occur, a doctor or a qualified health care practitioner should be consulted.

4.9. Overdose

Well-established use	Traditional use
	No case of overdose has been reported.

5. Pharmacological properties

5.1. Pharmacodynamic properties

Well-established use	Traditional use
	Not required as per Article 16c(1)(a)(iii) of Directive 2001/83/EC.

5.2. Pharmacokinetic properties

Well-established use		Traditional use
		Not required as per Article 16c(1)(a)(iii) of Directive 2001/83/EC.

5.3. Preclinical safety data

Well-established use	Traditional use
	Not required as per Article 16c(1)(a)(iii) of Directive 2001/83/EC, unless necessary for the safe use of the product.
	Adequate tests on reproductive toxicity, genotoxicity and carcinogenicity are not available.

6. Pharmaceutical particulars

Well-established use	Traditional use
	Not applicable

7. Date of compilation/last revision

05 June 2018