Assessment report on *Polygonum aviculare* L., herba

Final

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

<table>
<thead>
<tr>
<th>Herbal substance(s) (binomial scientific name of the plant, including plant part)</th>
<th><em>Polygonum aviculare</em> L., herba</th>
</tr>
</thead>
<tbody>
<tr>
<td>Herbal preparation(s)</td>
<td>Comminuted herbal substance</td>
</tr>
</tbody>
</table>
| Pharmaceutical form(s) | Comminuted herbal substance as herbal tea for oral use.  
Comminuted herbal substance for decoction preparation for oromucosal use. |
| Rapporteur(s) | B. Jansone |
| Peer-reviewer | J. Viguet Poupelloz |
5.2. Patient exposure ........................................................................................................ 29
5.3. Adverse events, serious adverse events and deaths .............................................. 29
5.4. Laboratory findings .................................................................................................. 29
5.5. Safety in special populations and situations ......................................................... 29
  5.5.1. Use in children and adolescents ................................................................. 29
5.5.2. Contraindications .............................................................................................. 29
5.5.3. Special warnings and precautions for use ........................................................ 29
5.5.4. Drug interactions and other forms of interaction ............................................ 29
5.5.5. Fertility, pregnancy and lactation ................................................................. 29
5.5.6. Overdose ........................................................................................................... 29
5.5.7. Effects on ability to drive or operate machinery or impairment of mental ability ......................................................... 29
5.5.8. Safety in other special situations ................................................................. 30
5.6. Overall conclusions on clinical safety ..................................................................... 30
6. Overall conclusions (benefit-risk assessment) .......................................................... 30
Annex .............................................................................................................................. 31
1. Introduction

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

- Herbal substance(s)

In accordance with the European Pharmacopoeia (8th ed., 07/2013:1885) knotgrass (Polygonum aviculare L. s.l.) consists of whole or fragmented dried flowering aerial parts of Polygonum aviculare L. The minimum content of flavonoids in dried drug: 0.30%, expressed as hyperoside (C_{21}H_{20}O_{12}; M_r 464.4) (dried drug).

In other pharmacopoeias and text books Polygonum aviculare is defined as follows:

- the whole or cut dried flowering aerial parts of Polygonum aviculare L. (WHO monographs plants on medicinal plants commonly used in the Newly Independent States [NIS] 2010),
- the dried herb, occasionally containing roots of Polygonum aviculare L. gathered during flowering (Blumenthal et al., 1998; Österreichisches Arzneibuch 1981),
- whole herb of Polygonum aviculare L. (Poradnik Terapeutyczny 1975; Weiss and Fintelman V 2000; Fintelman and Weiss 2006),
- whole herb of Polygonum aviculare L. collected during flowering (Produse farmaceutice românești 1970; Pētersone 1961; Rubine et al., 1977; Rubine and Šriste, 1999),
- whole herb of Polygonum aviculare L., sometimes with the root, collected during the flowering season and dried, as well as the fresh aerial parts collected during the flowering season (Gruenwald et al., 2000; 2004; 2007; Wyk and Wink 2005).

Description and origin of the plant

The Polygonaceae family consists of about 800 species and the Polygonum genus comprises about 300 species distributed throughout the world - in Europe (31 species), Asia, North Africa, North America and is also introduced into South America. Polygonum aviculare L. is native to Europe and Asia (China, Korea etc). In European Union (EU), it is mainly collected in Eastern Europe. It is a sturdy annual herbaceous plant and widely distributed mainly in fields or roadsides (Światek et al., 1986; Karlsson 2000; Sawicka et al., 2002; Costea and Tardif 2005, Wyk and Wink 2005 Narasimhulu et al., 2014; Styles 1962).

Knotgrass is procumbent and spreads along the ground. The stem is slender (diameter 0.5–2 mm), branched, with nodes, cylindrical and longitudinally striated. Average length of the stem is 30 cm. Leaves are small, narrow, sessile and glabrous entire. The characteristic of the plant is the sheath-like stipules (ochrea) which have form of a lacerated membrane in silver and brown colour at the base. Very small flowers have 5 greenish-white perianth segments, the tips of which are often coloured red, generally flowered from June-September. The fruits (2–4 mm) are brown to black ovate, triangular nuts, often punctate or striate (European Pharmacopoeia 8th ed., 07/2013:1885; Österreichisches Arzneibuch 1981; Wichtl 2004; Wyk and Wink 2005).

Polygoni avicularis herba is known under the following other names: knotgrass (English); Vogelknoterichkraut (German); maura sūrene (Latvian); takažolių žolė (Lithuanian); Ziele rdestu ptasiego (Polish); nat rdesna ptačího (Czech); madárkereserű virágos hajtás (Hungarian); Stavikrvová vňať (Slovak language); Vejpileurt (Danish); Vogelknoterichkraut (German).
Constituents
The following constituents of knotgrass (Polygonum aviculare herba) according to existing references are reported in the literature:

**Flavonoids** (0.1-1%; rarely 2.5-3%): derivatives of kaempferol, quercetin and myricetin, particularly avicularin (quercetin-3-O-arabinoside approximately 0.2%), juglanin (kaempferol-3-O-arabinoside), hyperoside, quercitrin, quercetin-3-galactoside, as well as vitexin, isovitexin, rhamnetin-3-O-galactoside, rhamnazin hydrogen sulphate, myricetin-3-O-rhamnoside, rutin, astragalin, isoquercitrin, miquelianin, spiraeoside, orientin, myricitrin, desmanthin-1, luteolin, betmidin, taxifolin, isorhamnetin, apigenin (Poradnik Terapeutyczny 1975, Granica et al., 2013b; Gruenwald et al., 2000; 2004; 2007; Haverland 1963; Kim et al., 1994; Malinowska 2013; Nugroho et al., 2014; Smolarz 2002 (a,b); Wichtl 2004; Wyk and Wink 2005; WHO monographs NIS 2010; Zhang and Xu 1989). Recently some new flavonoids were isolated: liquiritin and cinaroside (Yunuskhodzhaeva and Eshbakova 2010), 5,3',5,7-tetrahydroxy-4'-O-angeloyxyflavone-7-O-β-D-glucopyranoside (Sun et al., 2002). Five new flavones were isolated: 5,7-dihydroxy-6-methoxyflavane, 5,7-dihydroxy flavane, 5,7-dimethoxy-4'-hydroxyflavane, morin-7-O-β-D-glucoside and 5-hydroxy-3'-methoxyflavanone-7-O-rutinoside (Zheng et al., 1999). New dimeric procyanidin glucoside: catechin 3-O-acetate-(4α→8)-catechin 3-O-acetate-3′-O-β-D-glucopyranoside was isolated in 2012 (Cong et al., 2012).

**Tannins** (3.5-4%): rhatannin, gallo- and catechol tannins (Gruenwald et al., 2000; 2004; 2007; Wyk and Wink 2005; WHO monographs NIS 2010)

**Phenolic carboxylic acids:** caffeic, chlorogenic, gallic and protocatechuic acids (Sawicka et al., 2002; Światek et al., 1986; Wichtl 2004; WHO monographs NIS 2010)

**Naphthoquinone:** 6-methoxyplumbagin (Al-Hazimi and Haque 2002)

**Hydroxycomarins:** umbelliferone, scopoletin (Gruenwald et al., 2000; 2004; 2007)

**Lignans:** lignin glycoside, aviculin (Gruenwald et al., 2000; 2004; 2007; WHO monographs NIS 2010)

**Sterols:** mainly β-sitosterol (Narasimhulu et al., 2014)

**Saponins:** triterpenoid saponins, mostly oleanolic acid (Narasimhulu et al., 2014)

**Anthraquinones:** emodin (Cai et al., 2004)

**Silicic, tartaric and formic acids** (1%): present as water-soluble silicates (Poradnik Terapeutyczny 1975; Gruenwald et al., 2000; 2004; 2007; WHO monographs NIS 2010)

**Carbohydrates:** glucose, galactose, arabinose, sucrose, rhamnose, galacturonic acid (Yakovlev et al., 1985; WHO monographs NIS 2010)

**Other constituents:** essential oils, carotene, vitamins C and K, (Petersone 1961; Rubine et al., 1977; Yunuskhodzaeva and Eshbakova 2010; Yunuskhodzaeva and Abdullabekova VH 2012)

- Herbal preparation(s)

Comminuted herbal substance.
• 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.

Knotgrass is also available as a component of combination medicinal products with many other herbal substances/herbal preparations (for instance: Foeniculi fructus, Myrtilli herba, Galegae herba, Rubi fruticosi folium, Salviae officinalis folium and herba, Taraxaci radix cum herba, Urticae herba etc).

The following combination medicinal herbal products containing Polygoni avicularis herba are currently registered/marketed and available on the EU market. This information was provided by the National Competent Authorities in the overview of the marketed products:

• Czech Republic: herbal teas with 8-9 substances
• Germany: herbal tea with 8 substances
• Hungary: combination product with 4 substances
• Latvia: paste for oral use with 9 substances
• Poland: paste for oral use with 10 substances and combination product for bath preparation with 6 substances
• Slovak Republic: combination product with 10 substances
• Austria, Denmark, Latvia and United Kingdom: combination product in a form of tablet or capsules with more than 20 substances

This assessment report and the European Union herbal monograph refer exclusively to Polygoni avicularis herba as a single ingredient. The European Union herbal monograph describes the use of the comminuted herbal substance for tea preparations and for preparation of decoction for oromucosal or oral use

Vitamin(s): not applicable
Mineral(s): not applicable

1.2. Search and assessment methodology

Books, Book chapters, articles and letters in Journals, Medical press reviews, Acts of law and regulations, electronic databases and other sources were used to assess information available on traditional use, pharmaceutical, non-clinical, clinical data and current indications on Polygoni avicularis herba.

Search engines used: Google, Google Scholar; key words: Polygonum aviculare L., Polygoni avicularis herba, knotgrass, Vogelknöterichkraut

Scientific databases: PubMed, ScienceDirect, Scopus, Scifinder, Web of Science, EMBASE, EBSCO, Dawsonera; search period 1.06.2014-1.03.2015; key words: Polygonum aviculare L., Polygoni avicularis herba, knotgrass, Vogelknöterichkraut

Medical databases: UpToDate, Cochrane library, PubMed; search period 1.06.2014-1.03.2015; key words: Polygonum aviculare L., Polygoni avicularis herba, knotgrass, Vogelknöterichkraut

Toxicological databases: Toxnet; search period 1.06.2014-1.03.2015; key words: Polygonum aviculare L., Polygoni avicularis herba, knotgrass, Vogelknöterichkraut
Other resources:

- Libraries: hand searches in handbooks, textbooks and Pharmacopoeias on Polygoni avicularis herba and knotgrass at the various libraries were performed, namely, EMA, University of Latvia, Rigas Stradinu University, The State Agency of Medicines of Latvia, Central library in Riga

- Other databases: The World Health Organization, National Centre for Complementary and Alternative Medicine (NCCAM), The division for Complementary and Alternative Medicines of the Federal Institute for Drugs and Medical Devices (BfArM)

References in Latvian, English, German, Polish and Russian were assessed. Only references found to be relevant for assessment are included in the list of references.

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

According to the information provided by the National Competent Authorities in the overview of the marketed products, monocomponent medicinal products containing *Polygonum aviculare* L., herba are available on the markets of Lithuania and Poland.

Table 1: Overview of data obtained from marketed medicinal products

<table>
<thead>
<tr>
<th>Active substance</th>
<th>Indication</th>
<th>Pharmaceutical form</th>
<th>Regulatory Status</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polygoni avicularis herba; comminuted herbal substance</td>
<td>a) Traditionally used in mild urinary tract inflammation</td>
<td>Comminuted herbal substance <em>Adolescents, adults and elderly</em> a) Single dose: decoction of 3g of herbal substance in 200 mL for oral use 2 times daily between meals. Method of administration: oral use Duration: 2 weeks</td>
<td>Poland TUR since 19. 05. 2013</td>
</tr>
<tr>
<td></td>
<td>b) Traditionally used in mild inflammation of the mouth and throat.</td>
<td>b) Single dose: 3g (2 teaspoons) in glass of water (200-250 mL) as freshly prepared decoction for rinsing or gargling between meals 2 times a day. Method of administration: oromucosal use Duration: 1 week</td>
<td></td>
</tr>
<tr>
<td>Active substance</td>
<td>Indication</td>
<td>Pharmaceutical form</td>
<td>Regulatory Status</td>
</tr>
<tr>
<td>------------------</td>
<td>------------</td>
<td>---------------------</td>
<td>-----------------</td>
</tr>
<tr>
<td><em>Polygonum aviculare</em> L., herba; comminuted herbal substance for tea preparation</td>
<td>Traditional herbal medicinal product used for the relief of symptoms of common cold.</td>
<td>Comminuted herbal substance for tea preparation <em>Adolescents, adults and elderly</em> Single dose: Herbal tea: 1.5-2 g of the comminuted herbal substance ad to 150 mL cold water, then bring to boil. As a herbal infusion use 3–4 times daily, 30 minutes before meal. Daily dose: 5-8 g Duration: 1 week</td>
<td>Lithuania TUR since 05. 07. 1994</td>
</tr>
</tbody>
</table>

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**

Combination products containing *Polygonum aviculare* L., herba with only a few as well as up to 20 other herbal substances/herbal preparations (for instance: *Salviae officinalis* herba, *Thymi* herba, *Taraxaci radix cum herba* etc.) are available in Austria, Czech Republic, Denmark, Germany, Hungary, Latvia, Poland, Slovak Republic, United Kingdom. For more detailed description see below.

**Czech Republic**

a) Pharmaceutical form: Herbal tea containing (in 1 tea bag): 170 mg *Phaseoli fructus* sine semine, 150 mg *Myrtilli herba*, 150 mg *Salviae officinalis* herba, 120 mg *Galegae herba*, 100 mg *Polygoni avicularis* herba, 80 mg *Taraxaci radix cum herba*, 80 mg *Rubi fruticosi folium*, 80 mg *Foeniculi fructus*, 20 mg *Bardanae radix*  
Indication: as adjuvant in diabetes  
Posology: 1 tea bag /250 mL of boiling water 3 times daily  
On the market since 1969

b) Pharmaceutical form: Herbal tea containing (in 1 tea bag): 225 mg *Salviae officinalis* folium, 225 mg *Althaeae radix*, 225 mg *Polygoni avicularis* herba, 225 mg *Thymi herba*, 225 mg *Urticae herba*, 150 mg *Foeniculi fructus*, 75 mg *Sambuci flos*, 75 mg *Plantaginis folium*, 75 mg *Liquiritiae radix*  
Indication: as adjuvant for treatment of acute and chronic upper respiratory tract catarrhs  
Posology: 1 tea bag /250 mL of boiling water 3 times daily  
On the market since 1971

c) Pharmaceutical form: Herbal tea containing (in 1 tea bag): 413 mg *Betulae folium*, 337 mg *Uvae ursi folium*, 150 mg *Ononis radix*, 150 mg *Petroselini radix*, 165 mg *Polygoni avicularis* herba, 150 mg *Urticae herba*, 105 mg *Millefolii herba*, 30 mg *Sambuci flos*  
Indication: as adjuvant for treatment of symptoms of mild lower urinary tract infections such as burning sensation during urination and frequent urination

Assessment report on *Polygonum aviculare* L., herba  
EMA/HMPC/143659/2015  
Page 8/31
Posology: 1 tea bag/250 mL of boiling water 3-5 times daily
On the market since 1969

Germany
Pharmaceutical form: herbal tea containing (in 1 sachet (1.6 g)): 240 mg Foeniculi amari fructus, 208 mg Thymi herba, 192 mg Tiliae flos, 192 mg Polygoni avicularis herba, 176 mg Lichen islandicus, 96 mg Primulae flos, 64 mg Lamium album flos, 64 mg Verbascum thapsus flos
Indication: traditional herbal medicinal product to liquefy mucus. (Traditionell angewendet zur Unterstützung der Schleimlösung im Bereich der Atemwege)
Posology: 4-6 times daily 1 cup of tea (1 sachet/cup)
On the market since: at least 1978

Hungary
Combination product containing: 450 mg Graminis rhizoma pulv., 375 mg Agrimoniae herba pulv., 375 mg Taraxaci radix pulv., 225 mg Polygoni avicularis herba pulv. /filter (1.5 g)
Recommended indication: to promote the elimination of excreta from the body

Latvia
Pharmaceutical form: Oral paste. 100 g of paste contains 67.2 g of composed extract: Extractum hydroalcoholicum compositum (1:1.3-1.6) ex: Petroselinum sativum Hoffm., radix; Trigonella foenum-graecum L., semen; Polygonum aviculare L., herba; Agropyron repens (L.) Beauv., rhizoma; Betula pendula Roth or/and Betula pubescens Ehrh., folium; Equisetum arvense L., herba; Levisticum officinale Koch., radix; Solidago virgaurea L., herba; Allium cepa L., squama (7/6/6/5/4/4/4/2/2)
Indication: in cases of infections of urinary tract, kidney and bladder, for treatment and prevention of nephrolithiasis
Posology: Adults 3-4 times daily teaspoon (approximately 5 g) paste, dissolved in 1/2 cup of warm boiled water
On the market since 2000

Poland
a) Pharmaceutical form: Oral paste containing in 100 g of products 67.2 g of composed extract:
Extractum compositum (1:1.3-1.6) ex: 12.5 parts Agropyri rhizomate, 5.0 parts Allii cepae squamata, 10.0 parts Betulae folio, 15.0 parts Foenugraeci semine, 17.5 parts Petroselini radice, 5.0 parts Solidaginis herba, 10.0 parts Equiseti herba, 10.0 parts Levistici radice, 15.0 parts Polygoni avicularis herba
Indication: traditional herbal medicinal product used adjunctively in infections and inflammatory conditions of the urinary tract, urolithiasis (in fine streaks, the so-called kidney gravel) and prophylactically nephrolithiasis
Posology: Adults 3-4 times daily teaspoon (approximately 5 g) paste, dissolved in 1/4 cup of warm boiled water
On the market since 06/01/1961

b) Pharmaceutical form: Herbal mixture for preparation a bath containing: 25.0 g Quercus cortex, 20.0 g Matricariae flos, 17.5 g Polygoni avicularis herba, 17.5 g Salviae folium, 17.5 g Urticae folium, 2.5 g Calendulae flos
Indication: traditional herbal medicinal product used to wash of the area of external female genital organs in mild inflammation states
Posology: 15-20 g of herbal mixture (6 spoons) pour with 1.5 L of hot water, boil 10 minutes in
covered vessel, let stand 15 minutes, strain. Used for washing.

On the market since 1956

Slovak Republic

Contain: Polygoni avicularis herba, Phaseoli fructus s.sem, Myrtilli herba, Salviae herba, Galegae herba, Taraxaci radix c. herba, Rubi fruticosi folium, Foeniculi fructus, Bardanae radix, Liquiritiae radix

Austria, Denmark, Latvia and United Kingdom

Combination medicinal product containing Polygoni avicularis herba together with more than 20 active substances in a form of tablet or capsules is available on these markets.

Indications mentioned: a traditional herbal medicinal product used to relieve the symptoms of Raynaud’s syndrome and for the relief of symptoms associated with minor venous circulatory disturbances, such as tired heavy legs, pain, swelling, and for calf cramps; herbal medicinal product for the relief of walking provoked pains in the legs due to insufficient blood circulation (claudication intermittence).

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

Czech Republic

The herbal substance and preparations thereof are available on the market as food supplements as well.

Italy

Polygoni avicularis herba is included in the list of herbal substances and herbal preparations allowed in food supplements, published on the website of the Italian Ministry of Health, with the following indication: ’Regulation of intestinal transit; Counteraction of disturbances of menstrual cycle; Functionality of digestive system; Functionality of upper airways.’

Latvia

The herbal substance and preparations thereof are available on the market as food supplements (mono and combination products).

**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**

Knotgrass herb (Polygoni avicularis herba) is included in many handbooks, compendia, monograph such as Hagers Handbuch (Frerichs et al., 1927), Lehrbuch der Biologischen Heilmittel (Madaus, 1938, 1976), Drogenkunde (Hoppe, 1942), Farmakopea Polska (1970), Gift- and Arzneipflanzen von Mitteleuropa (Gessner and Orzechowski, 1974), Braun H (1974), Poradnik Terapeutyczny (1975), Hagers Handbuch (List & Hörhammer, 1977), Teedrogen (Wichtl, 1984), Kommission E Monographie, 1987, The Complete German Commission E Monographs (Blumenthal et al., 1998), Herbal drugs and Phytopharmaceuticals (Wichtl, 2004), Rational Phytotherapy (Schulz et al., 2004), PDR for Herbal medicines (Gruenwald et al., 2004 and 2007), Medicinal plants of the world (Wyk and Wink, 2005), A Compendium 'Medicinal Herbs' (Gehrmann et al., 2005), Herbal Medicine (Weiss and Fintelmann, 2000), Lehrbuch der Phytotherapie (Fintelmann and Weiss, 2006), WHO Monographs on medicinal
plants commonly used in the newly independent states (WHO monographs NIS 2010), in the list of herbals used by herbal industry in Poland (Ożarowski, 1978) and mentioned in the ordinance of the Minister of Health of Poland (Rozporządzenie Ministra Opieki Społecznej Poz 454, 1938).

The herbal substance is also mentioned in well-known handbooks as an expectorant and secretolytic to treat cough, bronchial catarrh and is also recommended for use in inflammation of the mouth and upper respiratory tract (Hagers Handbuch (Frerichs et al., 1927), Madaus, 1938 and 1976; Braun, 1974; Poradnik Terapeutyczny, 1975; Wichtl, 1984; Kommission E Monographie, 1987; Blumenthal et al., 1998; Weiss and Fintelmann, 2000; Wichtl, 2004; Gruenwald et al., 2000, 2004 and 2007; Gehrmann et al., 2005; Wyk and Wink, 2005). Several monographs and handbooks described the usage of knotgrass for the treatment of urinary tract infection with difficult painful urination, kidney stones and gallstone, gout and diuretic effect (Madaus, 1938 and 1976; Hoppe, 1942; Haverland, Roeske, 1955; 1963; Poradnik Terapeutyczny, 1975; Ożarowskiego, 1976, Wichtl, 1984; Mashkovsky, 1985 and 1993; Pētersone, 1961; Rubine et al., 1977; Skutelis and Pakalns, 2005 Sokolov and Zamotaev, 1984).

Diuretic activity of Polygoni avicularis herba was described also in the Pharmacopoeia of the former Soviet Union (Mashkovsky, 1985). In Lithuania, before 1990, Polygoni avicularis herba was available as medicinal product with indication for the treatment of urinary tract disorders. Also in Latvia, from 1999 until 2006, a Polygoni avicularis herba containing medicinal product was used to increase the amount of urine to achieve flushing of the urinary tract and in case of kidney stones. In Poland, knotgrass has been traditionally used as diuretic with evidence in the literature going back at least to the middle of the last century (Roeske, 1955; Farmakopea Polska, 1970; Poradnik Terapeutyczny, 1975; Ożarowskiego, 1976; Ożarowski, 1978). Currently one medicinal product as single ingredient herbal tea is available on the Polish market for the treatment of mild urinary tract inflammation.

In Madaus (1938, 1976), a broad spectrum of traditional usage is listed, for example, in bleedings (uterus, stomach, intestine, lung); diseases of the gastro-intestinal tract (ulcus ventriculi et duodeni, diarrhoea, gastroenteritis); catarrh of the lungs, tuberculosis, irritating cough, asthma, hoarseness; diseases of kidney and bladder, kidney stones and gravel.

Usage based on tradition in folk medicine indicated the external usage of the knotgrass for skin disorders such as eczema and vulval itching, skin wounds and ulcers as well as internal use in articular pain, rheumatism, as hemostatic in cases of haemorrhage and in choledolithiasis (Gruenwald et al., 2000, 2004 and 2007; Hoppe, 1942; Muszyński, 1954; Rubine et al., 1977; List & Hörhammer, 1977; Turowa et al., 1987; Wichtl, 1984; Rios et al., 1987; Novaretti and Lemordant 1990; Wichtl, 2004; Wyk and Wink, 2005).

The Chinese phonetic name for knotgrass is ‘Bian Xu’. In China, knotgrass has been used in traditional medicine since ancient times as diuretic, for throat and bronchial ailments as well as to treat skin problems (Gruenwald et al., 2000, 2004 and 2007; WHO monographs NIS 2010). Other usages in Chinese medicine are reported including gonorrhoea, jaundice or tapeworm in children (Gruenwald et al., 2000, 2004 and 2007).
Table 2: Overview of historical data

<table>
<thead>
<tr>
<th>Herbal preparation</th>
<th>Documented use / Traditional use</th>
<th>Pharmaceutical form</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Comminuted herbal substance or equivalent preparations</td>
<td>Mild catarrhs of the respiratory tract. Inflammatory changes to the oral and pharyngeal mucosa.</td>
<td>Daily dosage: 4-6 g as herbal tea and other galenical preparations for internal use and local application</td>
<td>Blumenthal et al., 1998</td>
</tr>
<tr>
<td>Comminuted herbal substance or equivalent preparations</td>
<td>Cough and bronchial catarrh. Inflammation of the mouth and upper respiratory tract.</td>
<td>Single dose: 1.5 g to be taken 3-5 times daily</td>
<td>Wyk and Wink, 2005</td>
</tr>
<tr>
<td>Comminuted herbal substance or equivalent preparations</td>
<td>Mild catarrhs of the respiratory tract. Inflammatory changes to the oral and pharyngeal mucosa.</td>
<td>Single dose: 1.5 g as infusion and other galenical preparations for internal use and local application taken 3-5 times daily Daily dosage: 4-6 g</td>
<td>Wichtl, 2004</td>
</tr>
<tr>
<td>Comminuted herbal substance or equivalent preparations</td>
<td>Cough/bronchitis. Inflammations of the mouth and pharynx.</td>
<td>Single dose: 1.5 g as infusion and other galenical preparations for internal use taken 3-5 times daily Daily dosage: 4-6 g Infusion for external use: daily dose of 5 g for local application</td>
<td>Gruenwald et al., 2000, 2004 and 2007</td>
</tr>
<tr>
<td>Comminuted herbal substance or equivalent preparations</td>
<td>Catarrhs of the respiratory tract. Inflammatory of mouth, throat and pharyngeal mucosa.</td>
<td>Single dose: 1.5 g/150mL as cold maceration for internal and local (gargle and cleansing) use, 3-5 times daily Daily dosage: 4-6 g Duration of use: if the acute symptoms persist longer than 1 week or illness is recurring, a medical practitioner should be consulted</td>
<td>Gehrmann et al., 2005</td>
</tr>
<tr>
<td>Comminuted herbal substance or equivalent preparations</td>
<td>For treatment of gingivitis, For the relief of cough and cold symptoms</td>
<td>Single dose: 1.5 g as infusion for oral use taken 3-5 times daily Daily dosage: 4-6 g</td>
<td>WHO monographs NIS 2010</td>
</tr>
<tr>
<td>Comminuted herbal substance or equivalent preparations</td>
<td>Catarrh of the respiratory tract. Inflammatory changes to the oral mucosal membrane</td>
<td>No information</td>
<td>Weiss and Fintelmann, 2000</td>
</tr>
<tr>
<td>Comminuted herbal substance or equivalent preparations</td>
<td>as adjuvant in coughs and bronchial catarrh, astringent gargle for the symptomatic treatment in the mouth and throat</td>
<td>Single dose: 1.5 g as decoction use 3-5 times daily</td>
<td>Wichtl, 1984</td>
</tr>
<tr>
<td>Comminuted herbal substance</td>
<td>For the treatment of urinary tract disorders</td>
<td>Single dose: 2 g in glass of water for oral use as decoction, 2-3 times daily</td>
<td>Ożarowski, 1978;</td>
</tr>
<tr>
<td>Herbal preparation</td>
<td>Documented use / Traditional use</td>
<td>Pharmaceutical form</td>
<td>Reference</td>
</tr>
<tr>
<td>--------------------</td>
<td>----------------------------------</td>
<td>----------------------</td>
<td>-----------</td>
</tr>
</tbody>
</table>
| Comminuted herbal substance or equivalent preparations | As diuretic  
In gout, bad metabolism, pulmonary diseases | About 5–10 g in glass of water for oral use, taken ¼-1/2 of the glass 2-3 times daily | Poradnik Terapeutyczny, 1975 |
| Comminuted herbal substance | As diuretic, bad metabolism, as astringent | Single dose: 10-20 g in 2 glasses of water (400-500 mL) for oral use as decoction, 3 times daily | Ożarowskiego, 1976 |
| Comminuted herbal substance | As an adjuvant pulmonary therapy, astringent gargle for the symptomatic treatment in the mouth and throat, in gout, diuretic | 1.5 g in about 150 mL water as infusion or decoction  
Daily dosage: 4.8-7.2 g | Hagers Handbuch (List & Hörhammer1977), |
| Comminuted herbal substance | catarrh of the bladder, cough, kidney stones and gallstone, | Infusion for oral use of 20 g of herbal substance in 200 mL, take 1 tablespoon 3 times daily | Rubine et al., 1977 |
| Comminuted herbal substance | cough and bronchial catarrh, gout as diuretic | 1 tablespoon 3 times daily | Pētersone, 1961 |
| Comminuted herbal substance | As diuretic | No information | Haverland, 1963 |
| Comminuted herbal substance | As diuretic  
in nephrolithiasis | Daily dosage: 2 glasses (400-500mL) of 10% Infusion, usage divided in several portions | Roeske, 1955 |
| Comminuted herbal substance | As adjuvant therapy in the lung disorders, for gout as diuretic | No information | Hoppe, 1942 |
| Comminuted herbal substance | Catarrh of the lungs, irritating cough, diseases of kidney and bladder, kidney stones and grave in bleedings, gastro-intestinal tract disorders | Daily dosage: 4.8-7.2 g (2-3 teaspoons) as infusion | Madaus, 1938, 1976 |
| Comminuted herbal substance | throat and bronchial catarrh | No information | Hagers Handbuch (Frerichs et al., 1927) |
## 2.3. Overall conclusions on medicinal use

Table 3: Overview of evidence on period of medicinal use

<table>
<thead>
<tr>
<th>Herbal preparation</th>
<th>Indication</th>
<th>Posology, Strength</th>
<th>Period of medicinal use</th>
</tr>
</thead>
</table>
| Comminuted herbal substance for tea preparation | For the relief of symptoms of common cold | Adolescents, adults and elderly  
Single dose: Herbal tea: 1.5-2 g of the comminuted herbal substance in 150 mL of water 30 minutes before meal 3–4 times daily  
Daily dosage: 4.5-8 g  
Method of administration: oral use  
Duration of use: 1 week | TUR use in Lithuania since 05. 07. 1994  
Posology data from literature since 1938.  
Madaus, 1938, 1976;  
Pētersone, 1961;  
Poradnik Terapeutyczny, 1975;  
Hagers Handbuch (List & Hörhammer, 1977);  
Wichtl, 1984;  
Blumenthal et al., 1998;  
WHO monographs NIS 2010;  
Gruenwald et al., 2000, 2004 and 2007;  
Wichtl, 2004;  
Wyk and Wink, 2005;  
Gehrman et al., 2005. |
| Comminuted herbal substance | Mild inflammation of the mouth and throat | Single dose: 3 g (2 teaspoons) in glass of water (200-250 mL) as freshly prepared decoction for rinsing or gargling between meals 2 times a day.  
Method of administration: oromucosal use  
Duration: 1 week | TUR use in Poland since 19. 05. 2013.  
Posology data from literature since 1975.  
Poradnik Terapeutyczny, 1975;  
Hagers Handbuch (List & Hörhammer, 1977);  
Wichtl, 1984;  
Blumenthal et al., 1998;  
Gruenwald et al., 2000, 2004 and 2007;  
Wichtl, 2004;  
Wyk and Wink, 2005;  
Gehrman et al., 2005. |
| Comminuted herbal substance | Mild urinary tract inflammation | Adults  
Single dose: decoction of 3 g of herbal substance in 200 mL for oral use 2 times daily, between meals.  
Daily dose: 6 g  
Method of administration: oral use  
Duration: 2 weeks | TUR use in Poland since 19. 05. 2013.  
Posology data from literature since 1938.  
Madaus, 1938, 1976;  
Pētersone, 1961;  
Poradnik Terapeutyczny, 1975;  
Ożarowskiego, 1976;  
Hagers Handbuch (List & Hörhammer, 1977);  
Rubine et al., 1977;  
Ozarowski, 1978;  
Wyk and Wink, 2005. |
The most frequently cited indications that can be based on the traditional medicinal use of Polygoni aviculare herba refers to the mild catarrhs of the respiratory tract, inflammation of the mouth and upper respiratory tract and renal elimination of water. For those indications the traditional use of Polygoni aviculare herba for more than 30 years in European Union in the forms of infusion and decoction is supported by the literature as well as by the medicinal products of *Polygonum aviculare* L., herba that are available in Lithuania and Poland.

The wording 'traditional herbal medicinal product to increase the amount of urine to achieve flushing of the urinary tract as an adjuvant in minor urinary complaints' for the therapeutic indication of Polygoni aviculare herba related to urinary tract disorders is in accordance with other approved European Union monographs for traditional herbal medicinal products, which have similar properties on urinary tract (e.g. *Equisetum arvense* L., herba). The use of the standard wording in the monograph is preferred, although several historic variations exist for the use in minor urinary tract disorders.

There is no posology specified in the literature and no preparations known to be available on the EU market for the topical usage of Polygoni aviculare herba for skin disorders. Therefore this indication is not proposed in the monograph. Other indications (e.g. in bleedings of the uterus, stomach, intestine, lung) of Polygoni aviculare herba mentioned in older books are not acceptable for a traditional herbal medicinal product. Based on the available data, the requirements for the traditional medicinal use according to Directive 2001/83/EC is fulfilled and included in the European Union Monograph for the following indications:

**Indication 1:**
For symptomatic treatment of minor inflammations in the mouth or the throat
*In adolescents, adults and elderly:*
Single dose: 1.5 g; rinsing or gargling between meals, 4-5 times daily.
Duration of use: 1 week.

**Indication 2:**
For the relief of symptoms of common cold
*In adolescents, adults and elderly:*
Single dose: 1.5-2 g; 3-4 times daily.
Duration of use: 1 week.

**Indication 3:**
To increase the amount of urine to achieve flushing of the urinary tract as an adjuvant in minor urinary complaints
*In adolescents, adults and elderly:*
Single dose: 3 g; 2 times daily.
Duration of use: 2 weeks.
The use of knotgrass in children and adolescents under 12 years of age is not recommended due to lack of data.

**3. Non-Clinical Data**

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

Anti-inflammatory activity

In vitro

Aqueous extracts

Inhibitory effect on the prostaglandin biosynthesis and platelet activating factor (PAF)-induced exocytosis of Polygoni avicularis herba were studied among other 52 plants of Swedish traditional medicine used to treat inflammatory diseases and/or wounds (Tunón et al., 1995). The dried plant material was extracted twice with water (1:20 and 1:10) for 48 hours at room temperature and then lyophilised. The extract of Polygoni avicularis herba induced high (83%) inhibition level on platelet activating factor (PAF)-induced exocytosis and moderate (52%) inhibition effect on prostaglandin biosynthesis indicating an anti-inflammatory activity of this herbal substance (Tunón et al., 1995).

Isolated constituents

The inhibition of elastase release by 11 flavonol glucuronides (myricetin 3-O-β-D-glucuronide, mearsetin 3-O-β-D-glucuronide, quercetin 3-O-β-D-glucuronide, isorhamnetin 3-O-β-D-glucuronide, kaempferide 3-O-β-D-glucuronide, kaempferol 3-O-β-(2″-O-acetyl-β-D-glucuronide), isorhamnetin 3-O-β-(2″-O-acetyl-β-D-glucuronide), quercetin 3-O-β-(2″-O-acetyl-β-D-glucuronide), quercetin 3-O-β-(3″-O-acetyl-β-D-glucuronide), kaempferol 3-O-β-(3″-O-acetyl-β-D-glucuronide)) isolated from Polygonum aviculare L. was studied using the model of elastase release inhibition in human neutrophils (Granica et al., 2013a). All 11 compounds were obtained from Polygonum aviculare L., herba (1000 g), which was powdered and extracted with solvents of rising polarity (petroleum ether, chloroform, ethyl acetate, acetone and water). All compounds with exception of kaempferol 3-O-β-D-glucuronide and quercetin 3-O-β-(2″-O-acetyl-β-D-glucuronide) at 1 μM concentration significantly decreased the release of elastase by neutrophils compared to stimulated control. Myricetin 3-O-β-D-glucuronide, mearsetin 3-O-β-D-glucuronide, quercetin 3-O-β-D-glucuronide and quercetin 3-O-β-(3″-O-acetyl-β-D-glucuronide) exerted similar inhibitory capability as quercetin (served as a control). According to the authors the data demonstrated the anti-inflammatory activity of Polygoni avicularis herba (Granica et al., 2013a).

Antibacterial activity

Aqueous extracts

Knotgrass conventional extract and (pressure-assisted) water extract were tested for the antimicrobial effects against Salmonella Typhimurium, Listeria monocytogenes and Propionibacterium acnes together with extracts from Lygodium japonicum, Dianthus superbus and Sophora flavescens giving heterogeneous results. The results have shown as the most important antimicrobial effect the one against Salmonella Typhimurin, indicated by the significantly decreased MIC99 (the minimum inhibitory concentration) value for the Lygodium japonicum (3,33 mg/mL) and for the Polygonum aviculare (9.04 mg/mL) water extracts. Knotgrass water extract had not much effect in Listeria monocytogenes and in Propionibacterium acnes compared to Lygodium japonicum and Dianthus superbus extracts (Gou et al., 2011).

Aqueous and other extracts

Organic and aqueous solvent extracts (the yield of powder was 41% wt/wt from water extracts, 27% wt/wt from acetone, 17% wt/wt from ethanol and 15% wt/wt from chloroform extracts for the stem, the respective values of 32%, 30%, 20% and 18% wt/wt were obtained for the leaves) of Polygonum aviculare, herba were evaluated for antimicrobial activities on gram-negative bacteria (Escherichia coli,
Proteus mirabilis, Pseudomonas aeruginosa, Salmonella typhi, S. paratyphi and Shigella flexneri) and gram-positive bacteria (Staphylococcus aureus, Bacillus subtilis, and Streptococcus pyogenes) by the paper disc diffusion method using cotrimoxazole and chloramphenicol as standard antimicrobials for comparison of the plant effect. Extracts were used in various concentrations: 0.5 mL of 20.0, 18.0, 15.0, 10.0, 8.0, 5.0, 1.0, 0.5, 0.05 and 0.005 mg/mL. As well as a different concentrations of isolated panicudine dissolved in chloroform (400, 40, 4, 0.4 mg/g) were tested for antimicrobial activity by measurement of inhibition zone (Salama and Marraiki, 2010). The data show that the highest average activity of four plant extracts against the tested organisms had chloroform (e.g. diameter of inhibition zone 28 mm against Proteus mirabilis) followed by water extracts (e.g. diameter of inhibition zone 25 mm against Bacillus subtilis), ethanol extracts (e.g. diameter of inhibition zone 23 mm against Salmonella typhi and Proteus mirabilis) and finally the lowest activity was demonstrated by the acetone extracts (diameter of inhibition zone 2 mm against S. aureus). The results revealed that all of the plant extracts were effective against most of the tested microorganisms but the leaf extracts generally had lower activity against the test bacteria compared to the stem extracts. Antimicrobial activity of the extracts ws unaffected after exposure to heat, but was reduced at alkaline pH. Panicudine expressed very good antimicrobial activity against all Gram-negative and Gram-positive bacteria tested bacteria. In generally low minimum inhibitory concentration (MIC) and minimum bactericidal concentration (MBC) values of chloroform extracts of Polygonum aviculare, herba indicated good efficacy towards tested bacteria. The lowest MIC and the MBC were demonstrated against Salmonella paratyphi, Bacillus subtilis and Salmonella typhi. The only exception was higher MIC and MBC values obtained with Staphylococcus aureus (Salama and Marraiki, 2010).

Polygonum aviculare, herba was extracted with 95% V/V ethanol. As dried plant extracts were prepared three different concentrations (1.00, 0.10 and 0.01 mg/mL) for the testing against gram-negative (Acinetobacter baumannii and Pseudomonas aeruginosa from MEDINA’s Culture Collection) and gram-positive bacteria (Staphylococcus aureus). The extract was considered to show anti-bacterial activity if its percentage of inhibition was more than 60%. Results of this study indicated that Polygonum aviculare, herba exhibited anti-bacterial activity: inhibition of Acinetobacter baumannii was 81±3% for the dose 1.00 mg/mL and inhibition of Pseudomonas aeruginosa was 74±6% also for the higher dose 1.00 mg/mL. Gram-positive bacteria (Staphylococcus aureus) was inhibited 100±0% by Polygonum aviculare, herba at the dose of 1.00 mg/mL (Zhang et al., 2013).

Table 4: Overview of the main non-clinical data/conclusions

<table>
<thead>
<tr>
<th>Herbal preparation tested</th>
<th>Posology</th>
<th>Experimental model</th>
<th>Reference</th>
<th>Main non-clinical conclusions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aqueous extracts of Polygonum aviculare L., herba</td>
<td>Concentrations of 0-500 mg/mL</td>
<td>In vitro: extracted with the pressure-assisted water extraction (PAWE) technique, assessed measuring of minimum of inhibitory concentration against Salmonella Typhimurin</td>
<td>Gou et al., 2011</td>
<td>antibacterial activity produced</td>
</tr>
<tr>
<td>Aqueous extracts of Polygonum aviculare L., herba</td>
<td>1-0.2 mg/mL</td>
<td>In vitro: evaluation of the inhibitory effect on prostaglandin biosynthesis and platelet activating factor (PAF)-induced exocytosis</td>
<td>Tunón et al., 1995</td>
<td>anti-inflammatory activity demonstrated</td>
</tr>
<tr>
<td>Herbal preparation tested</td>
<td>Posology</td>
<td>Experimental model</td>
<td>Reference</td>
<td>Main non-clinical conclusions</td>
</tr>
<tr>
<td>---------------------------</td>
<td>----------</td>
<td>--------------------</td>
<td>-----------</td>
<td>-------------------------------</td>
</tr>
<tr>
<td>Aqueous-acetone, ethanolic, chloroform extract of <em>Polygonum aviculare</em> L., herba</td>
<td>concentrations of the extracts: 20.0, 18.0, 15.0, 10.0, 8.0, 5.0, 1.0, 0.5, 0.05 and 0.005 mg/mL</td>
<td><em>In vitro</em>: low minimum inhibitory concentration (MIC) and minimum bactericidal concentration (MBC) was estimated by the paper disc diffusion method</td>
<td>Salama and Marraiki, 2010</td>
<td>antibacterial activity against gram-negative and gram-positive bacteria demonstrated with highest average activity of four plant extracts against the tested organisms had chloroform (e.g. diameter of inhibition zone 28 mm against <em>Proteus mirabilis</em>) followed by water extracts (e.g. diameter of inhibition zone 25 mm against <em>Bacillus subtilis</em>)</td>
</tr>
<tr>
<td>Ethanolic extract of <em>Polygonum aviculare</em> L., herba</td>
<td>1.00, 0.10 and 0.01 mg/mL</td>
<td><em>In vitro</em>: if its percentage of inhibition of gram-negative (<em>Acinetobacter baumannii</em> and <em>Pseudomonas aeruginosa</em> from MEDINA’s Culture Collection) and gram-positive bacteria (<em>Staphylococcus aureus</em>) was more than 60% antibacterial activity was considered as significant</td>
<td>Zhang et al., 2013</td>
<td>anti-bacterial activity was exhibited at the higher dose (1.00 mg/mL) tested: inhibition of <em>Acinetobacter baumannii</em> was 81±3% and for the <em>Pseudomonas aeruginosa</em> was 74±6%. Gram-positive bacteria (<em>Staphylococcus aureus</em>) were inhibited 100±0%.</td>
</tr>
<tr>
<td>11 flavonol glucuronides of <em>Polygonum aviculare</em> L., herba, extraction solvent: petroleum ether, chloroform, ethyl acetate, aceton and water</td>
<td>0.5, 1 and 10 μM</td>
<td><em>In vitro</em>: influence of the release of elastase by neutrophils</td>
<td>Granica et al., 2013a</td>
<td>anti-inflammatory activity demonstrated</td>
</tr>
<tr>
<td>panicudine isolated from <em>Polygonum aviculare</em> L., herba extract, extraction chloroform</td>
<td>0.4, 4, 40, 400, mg/mL</td>
<td><em>In vitro</em>: measurement of inhibition zone was estimated</td>
<td>Salama and Marraiki, 2010</td>
<td>antibacterial activity against both gram-negative and gram-positive bacteria demonstrated</td>
</tr>
</tbody>
</table>

### 3.1.2. Secondary pharmacodynamics

#### Antioxidative effects

*In vitro*
Ethanolic extracts

The free radical scavenging activity of extract powder was determined using free radical scavenging, lipid peroxidation, superoxide radical scavenging assays and hydroxyl radical-induced DNA strand scission assays. In the Hsu (2006) study the *Polygonum aviculare* L., herba powder was extracted in 50% ethanol solution and then was lyophilised and the resulting powder extract was used in the experiments. One mg of this extract powder was dissolved in 1 mL of 50% ethanol solution to obtain 1000 μg/mL sample solution. 1000 μg/mL solutions were diluted into series of 1 μg/mL, 5 μg/mL, 10 μg/mL, 20 μg/mL, 50 μg/mL, 100 μg/mL, 200 μg/mL, 500 μg/mL, and 1000 μg/mL with 50% ethanol. Well-known antioxidant compounds such as 1,1-Diphenyl-2-picrylhydrazyl (DPPH) assay, (+)-catechin and L-ascorbic acid were used as controls. The results show that the IC$_{50}$ (the concentration required to inhibit radical formation by 50%) value of *Polygonum aviculare* L. extract in free radical scavenging assays is 50 μg/mL, while the IC$_{50}$ values of (+)-catechin and ascorbic acid were 35 μg/mL and 50 μg/mL, respectively, indicating that all have inhibitory effect on the DPPH radical. The IC$_{50}$ value in superoxide radical scavenging assays for *Polygonum aviculare* L. extract was 0.8 μg/mL but for (+)-catechin it was 40 μg/mL. Most of superoxide radicals were inhibited by 10 μg/mL *Polygonum aviculare* L. extract indicating that the superoxide scavenging activity of *Polygonum aviculare* L. extract is higher than that of (+)-catechin. The data obtained from the lipid peroxidation suppressing activity measured by TBA (2-thiobarbituric acid) assay shown that the IC$_{50}$ value of *Polygonum aviculare* L. extract was 16 μg/mL exhibiting similar curve effect of antioxidant activity to (+)-catechin (IC$_{50}$ value 17 μg/mL). The results of hydroxyl radical-induced DNA strand scission assay had shown the DNA protective effect of *Polygonum aviculare* L. extract. The total phenolics and flavonoid content of extract were 677.4 ± 62.7 mg/g and 112.7 ± 13 mg/g. The data of this study have demonstrated that *Polygonum aviculare* L. herba extracts have high phenolics and flavonoid content, and produces valuable antioxidant activity (Hsu, 2006).

Aqueous extracts

The antioxidant activity of *Polygonum aviculare* L., herba was examined by spectrophotometric method using DPPH free radical (Gou et al., 2011). The pressure-assisted water extraction technique was used to obtain higher extraction yields of plant and absorbance was measured using microplate reader. The efficient concentration (EC$_{50}$) was expressed as the concentration (mg/mL) required for 50% reduction of DPPH free radical. The high antioxidant activity was observed at *Polygonum aviculare* L., herba extracted by the pressure-assisted water extraction technique, showing the lowest EC$_{50}$ value (0.09 mg/mL). For the standard curve gallic acid was used. The total phenolic amount was expressed as gallic acid equivalent of plant extract (mg GAE/g). The free radical scavenging activity correlated with the content of total phenols (71.72 mg GAE/g) of *Polygonum aviculare* L., herba (Gou et al., 2011).

Other extracts

Dried *Polygonum aviculare* L., herba plant material extracted with 80% methanol was estimated among other 32 selected herbs for their antioxidant capacity by using the following methods: ABTS$^+$ (2,20 azinobis-(3-ethylbenzthiazoline-6-sulfonic acid)), DPPH (1,1-diphenyl-2-picrylhydrazyl radical) and ferric reducing/antioxidant power (FRAP) expressed as TEAC (total equivalent antioxidant capacities). The data were expressed as μM trolox per 100 g dry weight (dw). The TEAC values were 19.2 μM trolox/100 g dw for ABTS$^+$, 141 μM trolox/100 g dw for DPPH and 161 μM trolox/100 g dw for FRAP. The total phenolic amount was expressed as gallic acid equivalent of plant extract (mg GAE/g). The total phenolics were 11.2 mg of GAE/100g of dry weight for the *Polygonum aviculare* L., herba measured by Folin–Ciocalteu colorimetric method. High levels of phenolics in *Polygonum aviculare* L., herba indicate remarkable antioxidant activity (Wojdyło et al., 2007).
Isolated constituents

Granica et al., (2013a) investigated the antioxidant activity of 11 flavonol glucuronides (myricetin 3-O-β-D-glucuronide, mearsetin 3-O-β-D-glucuronide, quercetin 3-O-β-D-glucuronide, isorhamnetin 3-O-β-D-glucuronide, kaempferide 3-O-β-D-glucuronide, kaempferol 3-O-β-D-glucuronide, kaempferol 3-O-β-(2″-O-acetyl-β-D-glucuronide), isorhamnetin 3-O-β-(2″-O-acetyl-β-D-glucuronide), quercetin 3-O-β-(3″-O-acetyl-β-D-glucuronide), kaempferol 3-O-β-(3″-O-acetyl-β-D-glucuronide)) isolated from Polygonum aviculare L., herba. All studied compounds at concentrations of 1 and 10 μM significantly inhibited the production of reactive oxygen species (ROS). Quercetin 3-O-β-D-glucuronide, quercetin 3-O-β-(2″-O-acetyl-β-D-glucuronide) and quercetin 3-O-β-(3″-O-acetyl-β-D-glucuronide) all at the concentration of 10 μM strongly inhibited production of ROS with the effect comparable to quercetin that served as a control (Granica et al., 2013a).

Vasorelaxant activity

Ex vivo

The vasorelaxant activity of hexane and n-butanol (n-BuOH) extracts of Polygonum aviculare, herba was investigated on phenylephrine (3×10⁻⁶ M) pre-contracted rat aortic tissues. The maximum relaxation obtained by hexane extract was 62.6% and for n-BuOH extract was 89.7%. The expressed vasorelaxant activity on endothelium –intact aortic tissues was in a dose-dependent manner. Removal of functional endothelium or pre-treatment of the inhibitor of nitric oxide synthase (aortic tissues with NG-nitro-l-arginine methyl ester (l-NAME)) suppressed this effect indicating that relaxation of the vascular smooth muscle is realized via endothelium-dependent nitric oxide mechanism (Yin et al., 2005).

Antifungal activity

In vitro

The antifungal activity of Polygoni avicularis herba extracts (water, acetone, ethanol and chloroform) was tested on fungal isolates (Aspergillus flavus, Aspergillus fumigatus, Aspergillus niger and the yeast C. albicans) by the paper disc diffusion method using streptomycin as standard were used for comparison (Salama and Marraiki, 2010). Also a different concentrations of isolated panicudine dissolved in chloroform (400, 40, 4, 0.4 mg/g) were tested for antifungal activity by measurement of inhibition zone (Salama and Marraiki, 2010). The measurement of inhibition zone around each paper disc demonstrated that chloroform extract show a good antimicrobial activity against all tested fungi (diameter of inhibition zone - 17 mm against Aspergillus flavus, 18 mm against Aspergillus fumigatus, 14 mm against Aspergillus niger) except C. albicans (diameter of inhibition zone 0 mm). The next potent antimicrobial activity was expressed by the water extracts (diameter of inhibition zone - 14 mm against Aspergillus flavus, 15 mm against Aspergillus fumigatus, 13 mm against Aspergillus niger, 0 mm against Candida albicans), then followed by the ethanol extracts and, finally by the acetone extracts (Salama and Marraiki, 2010).

Anti-obesity activity

In vitro

Ethanol extract of Polygonum aviculare, herba was tested in vitro for enzymatic lipase activity in pancreatic lipase inhibition assay. The results showed that inhibition of the lipid accumulation in 3T3-L1 adipocytes in vitro was 21.2%, indicating anti-obesity activity for Polygonum aviculare, herba (Roh and Jung, 2012).
In vivo

The antiobesity effects of ethanol extract of *Polygonum aviculare*, herba was explored in high-fat diet-(HFD-) induced obese mice. The results indicated that administration of *Polygonum aviculare*, herba extract to HFD-induced obese mice significantly decreased adipose tissue weight, body weight gain, adipocyte size, and decreased lipogenic gene expression. The levels of serum triglyceride, leptin, and MDA were lowered as well as 3T3-L1 adipocyte differentiation was inhibited. The data obtained suggests that exerted antiobesity effects in HFD-induced obese mice is via the suppression of lipogenesis in adipose tissue and increased antioxidant activity (Sung *et al.*, 2013)

Anti-atherosclerotic effects

In vitro and in vivo

The ethanol extract of *Polygonum aviculare* L., herba was studied in atherosclerotic (ApoE KO) mice that were fed with a Western diet (WD) alone or with *Polygonum aviculare* L. ethanol extract (50 and 100 mg/kg) or a statin (10 mg/kg) for 12 weeks. Results show that *Polygonum aviculare* L. ethanol extract and statin intake induced less body weight, lowered serum lipid levels and blood pressures and had less adipose tissue than the WD group mice.

Examination of a staining of the aorta and adipose tissue, expression levels of adhesion molecules, and the effect of the NF-κB activity demonstrated that aorta ICAM-1, VCAM-1 and NF-κB levels were decreased by *Polygonum aviculare* L. ethanol extract in a dose-dependent manner. Also *Polygonum aviculare* L. ethanol extract and statin reduced atherosclerotic plaque and adipocyte size compared to the Western diet group and decreased phosphorylation of MAPK pathway components in the aorta of *Polygonum aviculare* L.-treated mice, suggesting that anti-atherosclerotic effects of the ethanol extract of *Polygonum aviculare* L., herba are mediated via a MAPK pathway dependent mechanism (Park *et al.*, 2014).

Anticancer effects

In vitro

Anticancer effects of *Polygonum aviculare* herbal methanol extracts using different concentrations (50, 100, 150, 200, 250, 300,350 400 ng/μL) was investigated using the MCF-7 cell line (human breast cancer cell line). Cell proliferation and apoptosis were evaluate by MTT (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl-tetrazoliumbromide) assay and Flow Cytometry, respectively. It was found that *Polygonum aviculare* herbal methanol extract induced cytotoxicity in MCF-7 cell line at concentrations higher than 300 ng/μL and it was corresponding to the highest rate of cell death measured by Trypan Blue and MTT assays (Habibi *et al.*, 2011).

The same authors further carried out RT-PCR (real-time polymerase chain reaction test) in order to evaluate the expression of apoptotic genes. The RT-PCR results showed up-regulation of P53 and down-regulation of Bcl-2 proteins which indicated the ability of *Polygonum aviculare* to induce apoptosis in MCF-7 cells and confirmed its potential anticancer properties (Habibi *et al.*, 2011).

Methanol extracts of *Polygonum aviculare*, herba of various concentration (50, 100, 250, 500, 1000 mg/mL) was tested on Cell lines nalm-6 (pre B-cell leukemia). Trypan blue method for assessing the viability of the cells and to evaluate the cytotoxic effect of the MTT assay (3-(4, 5 Dimethylthiazol-2-yl) was used. The results of the MTT assay demonstrated that with increasing concentrations cells increased death rate indicating that methanolic extract of *Polygonum aviculare*, herba has cytotoxic effect on nalm-6 cell line (Kazemivash and Jamali 2013).
3.1.3. Safety pharmacology

No data available.

3.1.4. Pharmacodynamic interactions

No data available.

3.1.5. Conclusions

Significant anti-inflammatory, antioxidative and antibacterial effects of *Polygonum aviculare* L., herba extracts (water, acetone, ethanol and chloroform) and isolated constituents (panicudine, flavonol glucuronides, avicularin) have been demonstrated in the experimental models *in vitro, ex vivo* and *in vivo*.

The aqueous extracts of *Polygonum aviculare* L., herba as well as fractions and isolated individual substances in some studies have shown less potent activity than chloroform extracts but higher activity than ethanol and acetone extracts (Salama and Marraiki, 2010). It has been pointed out in the several studies that the high phenolics and flavonoid content present in the leaves and stem of the *Polygonum aviculare* L. is responsible for the most of important pharmacological effects of *Polygonum aviculare* L., herba.

The historical medicinal use of *Polygonum aviculare* L., herba described in several monographs and well-known handbooks as well as indications used for traditional herbal products available on the EU market correspond with the effects obtained in the non-clinical studies and are plausible supporting the traditional use of the herbal substance and preparation in the following indications:

- for symptomatic treatment of minor inflammations in the mouth or the throat;
- for the symptomatic treatment of common cold;
- to increase the amount of urine to achieve flushing of the urinary tract as an adjuvant in minor urinary complaints.

Other effects not related to the indications proposed for the *Polygonum aviculare* L., herba in the monograph but demonstrated in the non-clinical studies are anti-obesity activity, antifungal, vasorelaxant, anti-atherosclerotic and anticancer effects.

None of the reported non-clinical pharmacological studies described any cause for safety concerns.

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

Tissue (heart, liver, spleen, lung, kidney and bladder) distribution of avicularin isolated from Polygoni avicularis herba after a single i.v. administration in rat was measured by HPLC method. The concentration of avicularin in above mentioned tissues showed significant diversity within 60 minutes. Determination of avicularin indicated that kidney and bladder showed the highest concentrations of this compound (Xu et al., 2012).
3.3. **Overview of available toxicological data regarding the herbal substance(s)/herbal preparation(s) and constituents thereof**

3.3.1. **Single dose toxicity**

No data available.

3.3.2. **Repeat dose toxicity**

Female Wistar rats were gavaged with *Polygonum aviculare*, herba extract (1 mL) for 10 days. After 10 days rats were sacrificed and samples of brain - hypotalamus, adrenal glands, liver, kidney and small intestine (duodenum) were collected for further histological analysis. Using hematoxylin – eosine staining emphasizing the neuro – endocrine – pituitary – adrenal axis reaction as well as activity of enzymes (lactate dehydrogenase, succinate-dehydrogenase, Mg-dependent adenosine triphosphatase, cytochromoxidase) were investigated. The obtained results indicated that *Polygonum aviculare*, herba extract did not produce any pathological effects at the neuro-endocrine axis level and in the metabolism of the studied organs (Roman et al., 2008).

3.3.3. **Genotoxicity**

No data available.

3.3.4. **Carcinogenicity**

No data available.

3.3.5. **Reproductive and developmental toxicity**

No data available.

3.3.6. **Local tolerance**

No data available.

3.3.7. **Other special studies**

**Cytotoxicity studies**

Dose-dependent protective effect of *Polygonum aviculare* L., herba aqueous extracts was quantitatively evaluated on acetaminophen-induced cytotoxicity in human embryonic kidney HEK293 cells using an MTS (cell viability) assay. *Polygonum aviculare*, herba extracts had significant recovery effect on acetaminophen-induced damage in HEK293 cells. Dose response non-linear regression analysis demonstrated that *Polygonum aviculare*, herba extracts had a strong recovery effect rate (98%) on acetaminophen-induced damage in HEK293 cells. The EC50 value was 0.1 ng/mL (Sohn et al., 2009).

The cytotoxic effect of 11 flavonol glucuronides (myricetin 3-O-β-D-glucuronide, mearsetin 3-O-β-D-glucuronide, isorhamnetin 3-O-β-D-glucuronide, kaempferide 3-O-β-D-glucuronide, kaempferol 3-O-β-D-glucuronide, kaempferol 3-O-β-(2″-O-acetyl-β-D-glucuronide), isorhamnetin 3-O-β-(2″-O-acetyl-β-D-glucuronide), quercetin 3-O-β-(2″-O-acetyl-β-D-glucuronide), quercetin 3-O-β-(3″-O-acetyl-β-D-glucuronide), kaempferol 3-O-β-(3″-O-acetyl-β-D-glucuronide)) isolated from *Polygonum aviculare* L., herba was investigated on human neutrophils by flow cytometry with propidium iodide assay.
The data show that these compounds did not influence the viability of neutrophils at concentrations up to 30 μM. The camptothecin at the concentration of 10 μM was used as a positive control (Granica et al., 2013a).

### 3.3.8. Conclusions

The data on toxicology of *Polygonum aviculare*, herba and relevant preparations are limited. The repeated dose study *in vivo* consuming *Polygonum aviculare*, herba extract for 10 days did not produce any pathological effects in brain - hypotalamus, adrenal glands, liver, kidney and small intestine (duodenum).

Oral administration of *Polygonum aviculare*, herba can be regarded as safe at traditionally used doses. Due to the lack of data on genotoxicity a list entry cannot be proposed.

### 3.4. Overall conclusions on non-clinical data

Non-clinical data on *Polygonum aviculare*, herba support the traditional use of the herbal substance and preparation the proposed indication.

Pharmacokinetics and interactions have not been fully evaluated. Limited data available indicated that the highest concentrations of avicularin, isolated from Polygoni avicularis herba, were found in the kidney and the bladder after 24 hours of administration (Xu et al., 2012).

There are no data available regarding single dose toxicity and non-clinical information on the safety of Polygoni avicularis herba.

Other specific studies demonstrated protective effect on acetaminophen-induced cytotoxicity in human embryonic kidney cells (Sohn et al., 2009) and no cytotoxic effect on human neutrophils (Granica et al., 2013a).

Tests on genotoxicity and carcinogenicity have not been performed.

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

The long-term widespread use in the European Union and available data indicate no toxicological concern and potential risk associated with Polygoni avicularis herba use.

Oral and oromucosal administration of Polygoni avicularis herba can be regarded as safe at traditionally used doses.

### 4. Clinical Data

#### 4.1. Clinical pharmacology

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

No data available.

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

No data available.
4.2. **Clinical efficacy**

No data available.

4.2.1. **Dose response studies**

No data available.

4.2.2. **Clinical studies (case studies and clinical trials)**

The effectiveness of *Polygonum aviculare* L. extract (a natural Mexican Sanguinaria extract, extraction solvent ethanol) against gingivitis has been studied in 60 male volunteers with gingivitis aged 18 to 25 years. However, the methodology of the study is weak; for instance, parts of the *Polygonum aviculare* L. used in this study are not clearly specified. The following inclusion criteria were used: at least 24 natural teeth and gingivitis. The following exclusion criteria were introduced: current illness, a history of oral mucosal tissue reaction or known sensitivity to oral rinses, orthodontic appliances, periodontitis, continuous antibiotic usage and unwillingness to use the oral rinse and to adhere strictly to the examination procedures. Over a period of 2 weeks no tooth-brushing was allowed instead volunteers of this study use *Polygonum aviculare* L. extract (1 mg/ml) as oral rinse twice daily. The Löe and Silness Gingival Index was recorded at day 0, the study population had a gingival index mean of 1.056 (S.D.±0.074) with a maximum value of χ=1.290 and a minimum value of χ=0.920. The O'Leary Plaque Index had an arithmetic mean of χ=91.389 (S.D.±9.168) with a maximum value of χ=100 and a minimum value of χ=62.50. During the treatment the prevalence of gingivitis decreased, demonstrating χ=1.017 (S.D.±0.032) on day 7, an χ=1.012 (S.D.±0.32) on day 11, and an χ=1.011 (S.D.±0.032) on day 14 (P≤0.05). The obtained results indicated the usefulness of the *Polygonum aviculare* L. as supportive in the therapy of gingivitis in contrast, the results of the plaque index shown the significant increase from day 0 (χ=91.38) to day 14 (χ=98.69) (P≤0.05). However, the consistency of this plaque permitted its mechanical flushing easily (González Begné et al., 2001).
Table 5: Clinical studies on humans, in gingivitis

<table>
<thead>
<tr>
<th>Type</th>
<th>Study</th>
<th>Test Product(s)</th>
<th>Number of subjects</th>
<th>Type of subjects</th>
<th>Outcomes</th>
<th>Statistical analysis</th>
<th>Clinical relevance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Treatment of gingivitis using <em>Polygonum aviculare</em> L.</td>
<td>open study</td>
<td>Treatment: extract of <em>Polygonum aviculare</em> L. (1 mg/mL) as oral rinse twice daily for 2 weeks</td>
<td>60 patients with gingivitis the Löe and Silness Gingival Index had a mean of 1.056 (S.D.=0.074), a maximum value was $\chi=1.290$, a minimum value was $\chi=0.920$ the O'Leary Plaque Index had an arithmetic mean of $\chi=91.389$ (S.D.=9.168) with a maximum value of $\chi=100$ and a minimum value of $\chi=62.50$. age: 18 to 25 sex: male 9 drop-out (15%) reason: unknown</td>
<td>Patients with gingivitis inclusion criteria: at least 24 natural teeth and gingivitis exclusion criteria: current illness, a history of oral mucosal tissue reaction or known sensitivity to oral rinses, orthodontic appliances, periodontitis, continuous antibiotic usage and unwillingness to use the oral rinse and to adhere strictly to the examination procedures required for the study</td>
<td>Improvement: the prevalence of gingivitis decreased, demonstrating $\chi=1.017$ (S.D.=0.032) on day 7, an $\chi=1.012$ (S.D.=0.32) on day 11, and an $\chi=1.011$ (S.D.=0.32) on day 14. whereas the plaque index increased significantly from day 0 ($\chi=91.38$) to day 14 ($\chi=98.69$) ($P\leq0.05$) Good tolerability, no adverse effects</td>
<td>Wilcoxon Matched-Pairs at day Wilcoxon Matched-Pairs Signed-Ranks test ($P\leq0.05$) (SPSS: PC package)</td>
<td>Methodology of the study is week, part of the plant studied is not clear</td>
</tr>
<tr>
<td>Efficacy study</td>
<td>O'Leary Plaque Index and the Löe and Silness Gingival Index were recorded at day 0 duration: 2 weeks evaluation performed on days 7, 11 and 14 no tooth-brushing was allowed</td>
<td>Treatment: extract of <em>Polygonum aviculare</em> L. (1 mg/mL) as oral rinse twice daily for 2 weeks</td>
<td>60 patients with gingivitis the Löe and Silness Gingival Index had a mean of 1.056 (S.D.=0.074), a maximum value was $\chi=1.290$, a minimum value was $\chi=0.920$ the O'Leary Plaque Index had an arithmetic mean of $\chi=91.389$ (S.D.=9.168) with a maximum value of $\chi=100$ and a minimum value of $\chi=62.50$. age: 18 to 25 sex: male 9 drop-out (15%) reason: unknown</td>
<td>Patients with gingivitis inclusion criteria: at least 24 natural teeth and gingivitis exclusion criteria: current illness, a history of oral mucosal tissue reaction or known sensitivity to oral rinses, orthodontic appliances, periodontitis, continuous antibiotic usage and unwillingness to use the oral rinse and to adhere strictly to the examination procedures required for the study</td>
<td>Improvement: the prevalence of gingivitis decreased, demonstrating $\chi=1.017$ (S.D.=0.032) on day 7, an $\chi=1.012$ (S.D.=0.32) on day 11, and an $\chi=1.011$ (S.D.=0.32) on day 14. whereas the plaque index increased significantly from day 0 ($\chi=91.38$) to day 14 ($\chi=98.69$) ($P\leq0.05$) Good tolerability, no adverse effects</td>
<td>Wilcoxon Matched-Pairs at day Wilcoxon Matched-Pairs Signed-Ranks test ($P\leq0.05$) (SPSS: PC package)</td>
<td>Methodology of the study is week, part of the plant studied is not clear</td>
</tr>
</tbody>
</table>
4.3. Clinical studies in special populations (e.g. elderly and children)

No data available.

4.4. Overall conclusions on clinical pharmacology and efficacy

One clinical study has been reported to investigate the efficacy of *Polygonum aviculare* L. in gingivitis. The obtained results have demonstrated the significant decrease in prevalence of gingivitis during the two weeks twice daily usage of *Polygonum aviculare* L. extract. However, the methodology of the study is weak, not clearly indicating the parts of the plant used in the study. Furthermore there are currently no preparations for treatment of gingivitis available on the EU market; neither there are strong literature data supporting it. Therefore, the indication of *Polygonum aviculare* L. in gingivitis is not currently supported in the monograph.

For the indications proposed in the monograph, no data from clinical studies are available on pharmacodynamics, pharmacokinetic, dose-response relationship or special populations, such as elderly and children related to the indications of traditional use. Therefore, in accordance with Directive 2001/83/EC, the well-established use cannot be supported.

In contrast, the plausibility of efficacy for the traditional use of *Polygonum aviculare* L., herba for the symptomatic treatment of minor inflammations of the mouth or the throat, the relief of symptoms in coughs and colds and to increase the amount of urine to achieve flushing of the urinary tract as an adjuvant in minor urinary complaints is based on long-standing use and availability in the EU. It is also sufficiently supported and well documented by a number of well-known handbooks and monographs (Hoppe 1942; Haverland, 1963; Pētersone, 1961; Rubine *et al.*, 1977; Blumenthal *et al.*, 1998; Weiss and Fintelmann, 2000; Wichtl, 2004; Gehrmann *et al.*, 2005; Wyk and Wink, 2005; WHO monographs NIS 2010 etc). The traditional use is also supported by pharmacological data, see section 3.

Overall, the medicinal use of *Polygonum aviculare* L., herba has to be regarded as traditional and allows the development of a European Union herbal monograph on the traditional use.

5. Clinical Safety/Pharmacovigilance

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

There is only one clinical study on *Polygonum aviculare* L. use in gingivitis for two weeks available (González Begné *et al.*, 2001). Besides the efficacy data, a good tolerability and no adverse effects were mentioned in the publication. However, in general the methodology of this study is weak, not clearly indicating the parts of the plant used in the study. Gingivitis is not the indication proposed in the monograph.
### Table 6: Clinical safety data from clinical trials

<table>
<thead>
<tr>
<th>Type</th>
<th>Study</th>
<th>Test Product(s)</th>
<th>Number of Subjects</th>
<th>Type of Subjects</th>
<th>Adverse reactions</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Treatment of gingivitis using <em>Polygonum aviculare</em> L.</td>
<td>open study: O’Leary Plaque Index and the Löe and Silness Gingival Index were recorded at day 0</td>
<td>Treatment: extract of <em>Polygonum aviculare</em> L. (1 mg/mL) as oral rinse twice daily for 2 weeks</td>
<td>60 patients with gingivitis: the Löe and Silness Gingival Index had a mean of 1.056 (S.D.±0.074), a maximum value was $\chi=1.290$, a minimum value was $\chi=0.920$ the O’Leary Plaque Index had an arithmetic mean of $\chi=91.389$ (S.D.=9.168) with a maximum value of $\chi=100$ and a minimum value of $\chi=62.50$. age: 18 to 25 sex: male</td>
<td>Patients with gingivitis inclusion criteria: at least 24 natural teeth and gingivitis exclusion criteria: current illness, a history of oral mucosal tissue reaction or known sensitivity to oral rinses, orthodontic appliances, periodontitis, continuous antibiotic usage and unwillingness to use the oral rinse and to adhere strictly to the examination procedures required for the study</td>
<td>No adverse effects Good tolerability Methodology of the study is weak, part of the plant studied is not clear</td>
<td></td>
</tr>
</tbody>
</table>
5.2. Patient exposure

No exact data on patient exposure are available. On the basis of the use in some Member States over a period of more than 30 years a significant exposure can be assumed.

5.3. Adverse events, serious adverse events and deaths

Any kinds of adverse events or deaths have not been reported so far.

5.4. Laboratory findings

No data available.

5.5. Safety in special populations and situations

No data available.

5.5.1. Use in children and adolescents

The use in children has not been investigated therefore the use in children under 12 years of age is not recommended. For the adolescents the use is supported through the traditional use.

5.5.2. Contraindications

Hypersensitivity to the active substance.

5.5.3. Special warnings and precautions for use

The use in children under 12 years of age has not been established due to lack of adequate data.

If the symptoms worsen during the use of the medicinal product, a doctor or a qualified health care practitioner should be consulted.

5.5.4. Drug interactions and other forms of interaction

No data available.

5.5.5. Fertility, pregnancy and lactation

No data on fertility is available. As well as no data are supporting the safety use of Polygonum aviculare L., herba during pregnancy and lactation. Therefore, it should not be recommended during pregnancy and lactation.

5.5.6. Overdose

No data available.

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

No data available.
5.5.8. Safety in other special situations

No data available.

5.6. Overall conclusions on clinical safety

No adverse events or deaths have been reported so far. Conventional clinical safety data are absent. There is only one clinical study on the efficacy of *Polygonum aviculare* L. on gingivitis demonstrating a good safety profile. However, the methodology of the study is weak, not clearly indicating the parts of the plant used in the study.

For the indications proposed in the monograph, there are no clinical safety data for *Polygonum aviculare* L., herba available. Although clinical data on use in children or adolescents are not available, there are products used in EU market with the indications: a) traditional herbal medicinal product used for the relief of symptoms of common cold and b) traditional herbal medicinal product used in mild urinary tract inflammation cases. In both indications, the use is intended also for adolescents.

There are no data on reproductive and developmental toxicity therefore the use during pregnancy cannot be recommended.

The duration of use is limited:

- to one week for the symptomatic treatment of minor inflammations in the mouth or the throat and used the relief of symptoms of common cold;
- two weeks in order to increase the amount of urine to achieve flushing of the urinary tract as an adjuvant in minor urinary complaints.

A long-standing medicinal use and experience of *Polygonum aviculare* L., herba is documented within the EU in accordance with Directive 2004/24/EC. The available information on present use and historical data indicate that there are no safety concerns of usage of *Polygonum aviculare* L., herba in humans if used in accordance with the specified indications and doses. The oral and oromucosal administration of *Polygonum aviculare* L., herba can be regarded as safe.

6. Overall conclusions (benefit-risk assessment)

Well-established use as defined in Directive 2001/83/EC cannot be accepted for *Polygonum aviculare* L., herba due to the lack of data from clinical studies to conclude on recognised clinical efficacy.

Based on the references available and the information provided by the National Competent Authorities the presence of medicinal products on the European market throughout a period more than 30 years, including at least 15 years within the EU is substantiated. The traditional medicinal use of *Polygonum aviculare* L., herba according to Directive 2004/24/EC is fulfilled.

The efficacy of *Polygonum aviculare* L., herba for the traditional medicinal use in the European Union monograph is plausible on the basis of long-standing use and experience for the following indications:

*For oromucosal use*

‘Traditional herbal medicinal product for symptomatic treatment of minor inflammations in the mouth or the throat’.
In adolescents, adults and elderly:

Single dose: 1.5 g in glass of water (200-250 mL) as freshly prepared decoction for rinsing or gargling between meals; 4-5 times daily; Daily dosage: 6–7.5 g; Duration of use: 1 week

For oral use:

‘Traditional herbal medicinal product used for the relief of symptoms of common cold”.

In adolescents, adults and elderly:

Single dose: 1.5-2 g of the comminuted herbal substance in 150 mL of boiling water as a herbal tea for oral use; 3-4 times daily; Daily dosage: 4.5-8 g; Duration of use: 1 week.

For oral use:

‘Traditional herbal medicinal product to increase the amount of urine to achieve flushing of the urinary tract as an adjuvant in minor urinary complaints’.

In adolescents, adults and elderly:

Single dose: Herbal tea: 3 g of the comminuted herbal substance in 200 mL of water as a decoction for oral use; 2 times daily; Daily dosage: 6 g; Duration of use: 2 weeks

The indications fulfil the requirements for traditional use in so far that they are suitable for self-medication.

Due to lack of adequate data, Polygonum aviculare L., herba cannot be recommended for oral use in children under 12 years of age.

Studies on genotoxicity, carcinogenicity, reproductive and developmental toxicology of Polygonum aviculare L., herba have not been conducted. A European Union list entry is not supported due to lack of data on genotoxicity for Polygonum aviculare L., herba.

There is no information on reproductive and developmental toxicity of Polygonum aviculare L., herba available, therefore the use during pregnancy and lactation due to insufficient data are not recommended. No data on fertility are available.

No adverse effects have been reported in the documentation of medicinal use. The history of long-term and present use in humans in the European Union and the currently available information indicate no safety concerns associated with the oral or oromucosal use of Polygonum aviculare L., herba in recommended dosages under the conditions specified in the monograph.

The therapeutic areas for browse search on the EMA website are ‘Mouth and throat disorders’, ‘Cough and cold’ and ‘Urinary tract and gynaecology disorders’.

No constituents with known therapeutic activity are identified by the HMPC.

Annex

List of references