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Guidance document
Authorisation of Homeopathics, anthroposophics
and other complementary medicinal products HMV4

1 Definitions, terms, abbreviations

1.1 Abbreviations

Ann. Annex
B.Hom.P. British Homeopathic Pharmacopoeia
HAS List List of homeopathic and anthroposophic substances (Annex 6 KPTPO)
HAB Deutsches homöopathisches Arzneibuch [German Homeopathic Pharmacopoeia]
HMP Human Medicinal Products
KPTPO Ordinance of 7 September 2018 of the Swiss Agency for Therapeutic Products on the
Simplified Licensing of Complementary and Phytotherapeutic Products
(Complementary and Phytotherapeutic Products Ordinance, KPTPO; SR 812.212.24)

Let. Letter
Para. Paragraph
Ph.Eur. Pharmacopoeia Europaea
Ph. F. Pharmacopée Française [French Pharmacopoeia]
SC List List of Schüssler salts (Annex 7 KPTPO)
Sec. Section
TPLRO Ordinance of the Swiss Agency for Therapeutic Products of 9 November 2001 on the
Licensing Requirements for Therapeutic Products (SR 812.212.22)
TPA Federal Act of 15 December 2000 on Medicinal Products and Medical Devices
(Therapeutic Products Act, SR 812.21)
TPLO Ordinance of the Swiss Agency for Therapeutic Products of 22 June 2006 on the
Simplified Licensing of Therapeutic Products and the Licensing of Therapeutic
Products by the Notification Procedure (SR 812.212.23)
VMP Veterinary medicinal products

2 Introduction and objective

This guidance document describes the requirements relating to the documentation for the submission
and simplified authorisation of homeopathic and anthroposophic medicinal products (MP) according to
Chapter 4, section 5 KPTPO, as well as complementary medicines used in alternative treatments
according to Chapter 6 KPTPO. As an Administrative Ordinance document aimed at the
administrative bodies, it does not directly specify the rights and obligations of private individuals.
Swissmedic uses this guidance document first and foremost as a resource for applying the legal
provisions on authorisation in a uniform and equitable manner. The intention of publishing this
guidance document is to show third parties what requirements have to be fulfilled, according to
Swissmedic practice, to ensure that corresponding applications can be processed as quickly and
efficiently as possible.

3 Scope

This guidance document applies to the simplified authorisation of homeopathic and anthroposophic
human and veterinary medicinal products (HMP and VMP) according to Chapter 4, section 5 KPTPO,
and other complementary medicines according to Chapter 6 KPTPO. It applies to new authorisation
applications for the corresponding MP.

The guidance document also applies to the requirements relating to the quality documentation for
applications for the authorisation of homeopathic and anthroposophic HMP and VMP according to Art.
14 para. 1 abis – a²quater TPA and Art. 11 TPA. As regards the rest of the requirements for the
corresponding applications, the guidance document Authorisation of human medicinal product with
new active substance HMV4 and the guidance document Authorisation in accordance with Art. 14
para. 1 let. a²bis-quater TPA HMV4 are applicable.

The guidance document does not apply to the authorisation of homeopathic and anthroposophic
medicinal products and other complementary medicines by the notification procedure according to
Chapter 7 KPTPO or to the authorisation of Asian medicinal products according to Chapter 5 KPTPO.
The guidance document Authorisation of Homeopathics, anthroposophics and other complementary medicinal products HMV4 and the guidance document Authorisation of Asian medicinal products HMV4 apply to these medicinal products.

4 Legal framework

The procedures for authorising homeopathic and anthroposophic HMP and VMP according to Chapter 4, section 5 KPTPO and complementary medicines for alternative treatments according to Chapter 6 KPTPO are based primarily on the following legislative texts:

**TPA**
- Art. 9 Marketing authorisation
- Art. 10 Conditions for granting a marketing authorisation
- Art. 11 Application for a marketing authorisation
- Art. 14 Simplified authorisation procedures

**TPLRO**
- Art. 2 General preconditions
- Art. 3 Documentation on the analytical, chemical and pharmaceutical tests
- Art. 4 Documentation on the pharmacological and toxicology tests
- Art. 5 Documentation on clinical trials
- Art. 6 Special requirements for fixed-dose combination medicinal products
- Art. 7 Documentation on the analytical, chemical and pharmaceutical tests (veterinary medicinal products)
- Art. 8 Documentation on innocuousness (veterinary medicinal products)
- Art. 9 Additional documentation on innocuousness and residues for tests on livestock
- Art. 10 Admissibility of pharmacologically active substances and proposed withdrawal periods
- Art. 11 Documentation on preclinical and clinical trials (veterinary medicinal products)
- Ann. 1 Information and texts on containers and packaging materials for human medicinal products
- Ann. 2 Information and text on containers and packaging materials for homeopathic and anthroposophic medicinal products without indication and medicinal products for gemmotherapy without indication
- Ann. 3 Requirements for the declaration of active substances and pharmaceutical excipients in human medicinal products
- Ann. 3a List of excipients subject to declaration
- Ann. 4 Requirements for the Information for healthcare professionals for human medicinal products
- Ann. 5.2 Requirements for the Patient information for homeopathic and anthroposophic medicinal products

**KPTPO**
- Art. 5 Principle of simplified authorisation
- Art. 6 Documentation on the pharmacological and toxicology tests
- Art. 7 Proof of therapeutic efficacy and safety
- Art. 9 Analytical, chemical and pharmaceutical documentation
- Art. 10 Toxicological and pharmacological documentation
- Art. 11 Clinical documentation

5 Description/General requirements/Assessment principles

5.1 Selection of the appropriate authorisation procedure

Homeopathic and anthroposophic medicinal products and other complementary medicines can be authorised under various procedures, depending on the composition of the medicinal product in...
question and its intended use. The annex to this guidance document gives an overview of these procedures and also serves as a decision tree that can be used for identifying the procedure to be employed for a particular preparation.

5.2 Formal requirements

The formal requirements are based on the guidance document Formal requirements HMV4 and the corresponding directory overview of documents to be submitted HMV4.

5.3 Declaration requirements

a) Full declaration form

The following information is required in the sections as shown below:

Active substances:

The active substances in medicinal products with and without indication should be listed under this heading in accordance with Ann. 1a no. 1, para. 1 let. e, no. 1 and 2, and Ann. 1a no. 1 para. 2 and 3 TPLRO.

If a monograph exists for the active substance in a recognised homeopathic pharmacopoeia or in the “Homeopathic preparations and substances for homeopathic preparations” chapter of the European Pharmacopoeia, the pharmacopoeia must be referenced immediately after the substance name.

e.g.

Aralia racemosa (HAB) D6
Belladonna (Ph.Eur.Hom.) D6

If a monograph exists for the active substance in the homeopathic section of Ph.F. or Ph.Eur., but the monograph does not specify manufacturing instructions, the Ph.Eur. manufacturing instructions used must be added after the potency.

e.g.

Ficus carica (Ph.F.) CH6 (Ph.Eur.Hom. 1.1.10)
Crocus (Ph.Eur.Hom.) D6 (Ph.Eur.Hom. 1.1.8)

If no monograph exists for the active substance in a recognised homeopathic pharmacopoeia, it has to be specified in greater detail in accordance with Annex 1a number 1 paragraph 3 letter c TPLRO. This means that the following have to be provided for mother tinctures and potencies of plant or animal origin:

– Primary plant(s) or animal(s) with genus and species
– Plant/animal parts used; indication of their condition (fresh/frozen). If the condition is defined in the manufacturing instructions, the additional indication can be omitted.
– Manufacturing instructions and the pharmacopoeia containing the instructions. This information should be inserted after the potency.
– Any manufacturing method-related labelling required by the HAB (e.g. “spag. Zimpel”, “Rh”) must also be inserted.

e.g.

Calendula officinalis e floribus D4 (HAB 2a) or
Calendula officinalis e floribus D4 (Ph.Eur.Hom. 1.1.3)

Aranea diadema ex animale toto D8 (HAB 4b) or
Aranea diadema ex animale toto D8 (Ph.Eur.Hom. 1.1.9)
Melissa officinalis ex herba Rh D6 (HAB 21)

If Ph. Eur. includes a monograph for the starting material but not the active substance, this monograph is binding for quality and must also be referenced in the form Full declaration HMV4.

e.g.
Magnesium asparticum (Magnesii aspartas dihydricus Ph.Eur.) D6 (HAB 5a)
Ipecacuanha e radice (Ipeccacuanha radix Ph.Eur.) D6 (HAB 4a)

Anthroposophic active substances that are derived from plant-based starting materials but not manufactured and potentised according to a homeopathic process or for which the homeopathic section of a recognised pharmacopoeia contains no manufacturing instructions are subject to the requirements for herbal substances and preparations stated in the guidance document Authorisation of phytotherapeutic products HMV4.

Active substances derived from Schüssler salts carry the number of the salt at the beginning of the active substance name. The active substance names given in the SC List apply. The remaining requirements are the same as those for homeopathic active substances.

e.g.
№ 11 Silicea (HAB) D12
№ 12 Calcium sulfuricum (HAB) D3

Contents from manufacture/potentisation:
The quantities of vehicles or excipients used during manufacture/potentisation (e.g. ethanol 96%, water, lactose monohydrate, glycerol) and accounting for at least 1 % of the finished product or necessary for the information required by Annex 3a TPLRO should be listed under this heading. The quantities of excipients of particular interest according to Annex 3a TPLRO should be listed. The constituents of excipient mixtures such as ethanol/water mixtures or sodium chloride solution must be listed separately. The quality of the excipients (generally Ph. Eur.) must be stated.

Excipients for the pharmaceutical form:
The quantity and quality of all excipients that are used, in addition to the active substance, in the manufacture of the pharmaceutical form should be listed under this heading. The function of the excipients should be stated. The quality of the excipients (generally Ph. Eur.) must be stated.

For pharmaceutical forms that contain ethanol, the ethanol content in the finished product should also be stated as a percentage by volume.

The total sodium content of medicinal products that contain sodium (e.g. solutions for injection containing sodium chloride) must be listed separately.

For liquid pharmaceutical forms, the required figures should be stated in ml.

If the medicinal product is dosed in drops, the drop equivalent should also be stated, as should the number of spray pumps per ml for sprays and the number of globules per gram for globules.

b) Folding carton and patient information

- Active substances manufactured according to a homeopathic manufacturing process
Active substances for which a monograph exists in a recognised homeopathic pharmacopoeia or in the “Homeopathic preparations and substances for homeopathic preparations” chapter of the European Pharmacopoeia should be stated in accordance with the form Full declaration.

e.g.
Aralia racemosa (HAB) D6
Crocus (Ph.Eur.Hom.) D6 (Ph.Eur.Hom. 1.1.8)

If a monograph for the starting material exists in the non-homeopathic part of the pharmacopoeia, it is not mandatory – contrary to the information in the form Full declaration – to specify the quality monograph for the starting material.

e.g.
Magnesium asparticum D6 (HAB 5a)

However, if it is easier to declare the starting material by referencing the Ph.Eur monograph – particularly if there are several possible primary plants – this can be done as follows using the Ph.Eur monograph names.

e.g.
Sennae folium (Ph.Eur) D6 (HAB 4a)
Ipecacuanhae radix (Ph.Eur.) D12 (Ph.Eur.Hom 1.1.8)

In addition to the scientific name, the Latin name commonly used in the respective school of therapy can optionally be stated.

e.g.
Atropa belladonna (Belladonna) (HAB) D4

Although a translation of the Latin active substance names commonly used in the school of therapy into German (or French/Italian) is unusual for potentised active substances, this may be included as additional information in the Patient information, provided the active substances are unambiguously and fully characterised in the translation. Only the translations approved by Swissmedic may be used. If a medicinal product without indication is being authorised with a reduced dossier or by the notification procedure, and Swissmedic has not reviewed or approved the packaging elements, translations into the Swiss national languages of Latin active substance names must not be added.

- Active substances in anthroposophic medicinal products

The information above applies to active substances manufactured using a homeopathic manufacturing process.

For anthroposophic active substances from plant-based starting materials that are not manufactured according to a homeopathic process, the requirements for herbal substances and preparations apply. In addition to the plant drugs used, the botanical name of the primary plant and the plant part used and, for extracts, the type of extract (e.g. dry extract, liquid extract), the drug-extract ratio and the extractant should be listed. Further details concerning these requirements can be found in the guidance document Authorisation of phytotherapeutic products HMV4.

- Excipients
Excipients should be listed in accordance with Annex 3 and 3a TPLRO. If a medicinal product requires neither Information for healthcare professionals nor Patient information, the requirements for the Information for healthcare professionals apply to the full declaration of the excipients on the external packaging or, if there is no packaging, the container (quantities of excipients of particular interest should be specified; all other excipients should be stated qualitatively).

If vehicles or excipients are used in the manufacture/potentisation of the active substances and account for at least 1 % of the finished product or are necessary for the information required by Annex 3a TPLRO, these should be listed among the excipients. The optional mention of other excipients used during manufacture is permitted only if these have also been listed accordingly on the Full declaration form.

For medicinal products containing ingredients of animal origin as described in chapter 5.2.8 of Ph.Eur., the requirements relating to the declaration of the relevant substances stated in the guidance document Minimising the risk of TSE HMV4 should also be observed.

5.4 Documentation requirements

The requirements relating to the documentation of the procedures described in this guidance document are described in Chapters 7 and 8. Evidence must also be provided to show that medicinal products for livestock only contain active substances listed as permitted pharmacologically active substances in foodstuffs legislation. If necessary, withdrawal periods should be proposed and substantiated accordingly.

5.5 Document protection

No document protection is granted for authorisations of complementary medicines according to the procedures for simplified authorisation described in this guidance document. Such protection is possible only for authorisations according to Art. 11 TPA (see guidance document Authorisation of human medicinal product with a new active substance HMV4).

5.6 Requirements concerning the investigation of the medicinal product in specific age groups

5.6.1 Children and adolescents

5.6.1.1 General requirements relating to the dosage instruction in the product information

If the dosage instruction distinguishes between different age categories, the respective ages with specific figures should be stated. The age group classification is based on the definitions in the ICH Guideline “Clinical Investigation of Medicinal Products in the Pediatric Population E11”:

- **Preterm newborn infants**: < 36th week of pregnancy
- **Term newborn infants**: 0 – 27 days
- **Infants and toddlers**: 28 days – 23 months
- **Children**: 2 – 11 years
- **Adolescents**: 12 – 18 years
- **Adults**: from 18 years

Age groups excluded from the dosage instruction must be listed both in the Contraindications (“Do not take/use … :) and Dosage/Administration (“How to take/use …”) sections.

5.6.1.2 Homeopathic and anthroposophic medicinal products according to Chapter 4 KPTPO

According to Ann. 2 no. 5 KPTPO, the following requirements apply to homeopathic and anthroposophic medicinal products:
- Data on children and adolescents must be submitted if dosage recommendations are requested for these age groups. Otherwise, use in these age groups should be ruled out with
the standard texts provided in Ann. 5.2 no. 5, 6 and 8 TPLRO, or permitted on a prescription-only basis.

- A Paediatric Investigation Plan according to Art. 54a TPA is not required for the simplified authorisation procedures described in this guidance document.

5.6.1.3 Complementary medicines used in alternative treatments according to Chapter 6 KPTPO

As regards children, medicines used in gemmotherapy and Schüßler therapy are subject to the same requirements as homeopathic and anthroposophic medicinal products.

For complementary medicines used for alternative treatments, the Agency specifies what documents are needed on a case-by-case basis.

5.6.2 Elderly people (from 65 years)

Any data available for this group of people should be described. As with the provisions for children, any missing data should be mentioned.

5.7 Pharmacovigilance Plan

5.7.1 Homeopathic and anthroposophic medicinal products according to Chapter 4 KPTPO

A Pharmacovigilance Plan according to Art. 11 para. 2. let. a no. 5 TPA is not required for the simplified authorisation procedures described in this guidance document.

5.7.2 Complementary medicines used in alternative treatments according to Chapter 6 KPTPO

As regards a Pharmacovigilance Plan, the same requirements apply as for homeopathic and anthroposophic medicinal products.

5.8 Time limits

The time limits are based on the guidance document Time limits for authorisation applications TPO4.

5.9 Fees

The fees specified in the Ordinance on Fees Levied by the Swiss Agency for Therapeutic Products (FeeO-Swissmedic; SR 812.214.5) apply.

6 Homeopathic and anthroposophic medicinal products with indication (Art. 24 para. 1 KPTPO)

6.1 Precondition for the application of the procedure

According to Art. 24 para. 1 KPTPO, a homeopathic or anthroposophic medicinal product with indication can be authorised if it fully satisfies the definition of a medicinal product in the corresponding school of therapy and if the quality, safety, innocuousness and therapeutic effect are adequately documented.

The following points in particular must be fulfilled:

- The starting materials correspond with Chapter 4, section 2 KPTPO
- All active substances are manufactured according to a homeopathic, anthroposophic or spagyric process (sections 3 and 4 KPTPO).
- The pharmaceutical form is known in the corresponding school of therapy (Art. 23 para. 2 KPTPO).
- Adequate safety is documented for the active substances and their use.
- The indications are substantiated according to the therapeutic principle in question.
6.2 Application documentation requirements

The principles concerning the requirements and scope of documentation are defined in Ann. 2 KPTPO.

6.3 Documentation on quality

6.3.1 Documentation on the quality of the active substance (Module 3.2.S)

Starting materials, active substances

Starting materials

The starting materials used must comply with the Pharmacopoeia monographs applicable to homeopathics and anthroposophics and satisfy the general requirements for starting materials stated in the Pharmacopoeia, HAB, Ph.F. and B.Hom.P.

If an individual monograph exists in a recognised pharmacopoeia for a starting material, the requirements documented therein must be fulfilled. If a corresponding monograph does not exist for a starting material, the company must prepare its own monograph. Quality criteria, as defined in similar monographs of recognised pharmacopoeias, must be specified taking into account the nature of the relevant starting material (for example, herbal, animal, mineral or human origin). Reasons must be given for the choice of test and the methods must be validated.

As regards testing for contaminants, the requirements of the general Ph. Eur. monograph Herbal Drugs for Homoeopathic Preparations/Plantae medicinales ad praeparationes homoeopathicas must be taken into account.

In the case of herbal starting materials, precise details of the primary plant/primary plants and the plant parts used must be provided.

Active substances

For active substances, both the quality of the starting materials used and the quality of the active substances themselves must be documented.

If the preparations are potentised, the process of potentisation must also be documented (see also Batch documentation for finished product). Compliance with the manufacturing instructions as described in the Pharmacopoeia and in HAB, Ph.F. and B.Hom.P. is essential.

Organ preparations

In the case of organ preparations, documentation showing how the requirements relating to organ preparations are observed must also be provided (Art. 18 KPTPO).

Nosodes

In the case of nosodes, documentation showing how the requirements relating to nosodes are observed must also be provided (Art. 19 KPTPO).

Mother tinctures/Solutions/First triturations

If an individual monograph exists in a recognised pharmacopoeia for a mother tincture, solution or first trituration, the requirements documented in the monograph must be fulfilled.

If a corresponding monograph does not exist for a mother tincture, solution or first trituration, the company must prepare its own monograph characterising the quality of the preparation. The structure and content of the company's monograph must be based on existing monographs included in the recognised pharmacopoeias.

Where applicable, assays must be carried out and limits specified (for defined constituents, including inorganic substances, pharmacologically active substances such as alkaloids, cardiac glycosides, etc.).
The description of the manufacture of the mother tinctures, solutions and first triturations must prove that the manufacturing instructions, as described in Ph. Eur. or – where applicable – in other recognised pharmacopoeias, are followed.

**Stability documents**

If they do not immediately undergo further processing, the stability of the mother tinctures, solutions and first triturations must be tested. The recorded data – including colour illustrations of the fingerprints obtained by thin-layer chromatography and/or GC/HPLC – as well as a critical evaluation of the data must be submitted. A substantiated retest period must be requested.

**Container for the active substance**

If the active substances are stored, documentation on the primary container used – particularly specifications and design drawings – and on the suitability of the materials used must be submitted. The required declarations of conformity must likewise be submitted. Particularly for liquid active substances, potential interactions with the container materials must be discussed.

**Excipients**

Specifications and test methods must be documented. If excipients are the subject of a monograph in the pharmacopoeia, the documented requirements must be satisfied; in these cases, a reference to the monograph is sufficient.

The same requirements apply to excipients used in the manufacture of the finished product (see CTD Module 3.2.P).

**Batch documentation**

Sample batch documentation must be submitted for each mother tincture, solution or first trituration used as an active substance, whether directly or after potentisation. The documentation must include all the records from the testing of the starting material through to the testing of the mother tincture, solution or first trituration.

**6.3.2 Documentation on the quality of the finished product (Module 3.2.P)**

**Composition of the finished product**

The full (qualitative and quantitative) composition of the finished product must be documented. Where applicable, the active substances must be designated in accordance with the Pharmacopoeia, HAB, Ph.F. or B.Hom.P. For pharmaceutical forms whose composition (particularly the choice of excipients) is not defined in HAB, Ph.F., in the recognised manufacturing instructions in B.Hom.P or in the Pharmacopoeia, the choice of excipients must be justified.

**Manufacture of the finished product**

The manufacture of the finished product must be described in both narrative and schematic form. The implemented in-process controls must be documented (specifications, analytical methods and test frequencies, if possible in tabular form). A standard batch size and/or batch size range must be defined.

The documentation must prove that the homeopathic or anthroposophic manufacturing instructions, as documented in Ph. Eur. or in other recognised pharmacopoeias, are followed.

The manufacturing process must be validated and the corresponding validation report submitted.

If validation is omitted, the reason for omitting it must be justified, taking into account the pharmaceutical form.

Procedures designed to achieve sterility must be described in detail.

Validation documents must be submitted for the corresponding manufacturing steps.

**Control of the finished product**
The specifications and analytical methods must be submitted, as must the validation documents, where applicable. The requirements of the respective Ph. Eur. monograph for the pharmaceutical form must be taken into account during the preparation of the specifications.

The following parameters – depending on the composition and pharmaceutical form – form part of the finished product specification:

- organoleptic aspects
- physical parameters
- identity tests (particularly chromatographic procedures for lower potencies and mother tinctures contained in the finished product)
- assays and/or limit tests for inorganic substances and herbal components with pharmacologically active/toxicologically relevant constituents
- dry residue or loss on drying
- alcohol content
- content of preservative(s)

The respective tests relating specifically to the pharmaceutical form must also be carried out.

**Container for the finished product**

Documentation on the primary container must be submitted. This includes specifications, test methods and design drawings, plus documentation on the materials used and their suitability for the intended use. The required declarations of conformity must likewise be submitted. Particularly for liquid and semi-solid pharmaceutical forms, potential interactions with the container materials must be discussed.

**Stability documentation for the finished product**

The stability of the finished product must be investigated in accordance with ICH and EMA guidelines, and reasons for the proposed shelf life must be provided.

A shelf-life specification must be submitted.

The stability tests must be carried out on at least two batches and cover the entire shelf life. At least one of the two batches must be a production batch.

At the time of submission, the stability data covering a storage period of at least 6 months for at least 2 pilot batches must be available.

The recorded data must be documented clearly in tabular form. For the stability batches, the manufacturing date, batch size and associated primary container must be stated. If applicable, the colour illustrations of the fingerprints obtained at the individual test times by thin-layer chromatography and/or GC/HPLC must be enclosed.

The submitted data must be discussed: *Out of specification* results, tendencies and statistically significant deviations, for example changes in the fingerprints, must be considered.

Stability after opening should be tested if applicable and depending on the pharmaceutical form. A use-by period backed by appropriate reasons must be requested.

**6.4 Non-clinical documentation**

**6.4.1 Type and scope of documentation for toxicologically known starting materials, active substances or excipients**

The familiarity of a starting material, active substance or excipient should be verified according to Annex 2 no. 3.4 KPTPO.

The toxicological innocuousness of the starting material or the proposed potency/dilution should be demonstrated by submitting scientific findings.
6.4.2 Type and scope of documentation for toxicologically new starting materials, active substances or excipients

For toxicologically new starting materials, active substances and excipients, all aspects of safety and possible pharmacokinetic substance interactions should be addressed. These include information on general toxicity, genotoxicity, reproductive toxicity, carcinogenicity, immunotoxicity or local tolerance (e.g. allergenic potential). This information can be based on publications or experimental data generated with a starting material, active substance or excipient that is comparable with the substance to be authorised. If possible, preference should be given to the use of validated/qualified alternative methods rather than animal studies. The documentation should be discussed in the Nonclinical Overview (CTD Module 2.4), for example; experimental data should be prepared accordingly in summary form (CTD Module 2.6). All documents should be submitted in full (CTD Module 4).

The following should also be taken into account for homeopathic and anthroposophic medicinal products:

- Toxicological tests on animals are not required for new starting materials, active substances or excipients in dilutions or concentrations that exclude a possible risk of allergy or toxicity.
- Specifically, documents and studies on the allergenic potential of new starting materials, active substances or excipients are required only if these substances are present in the medicinal product in potencies up to, and including, D7. Any omission of such documentation must be justified.
- Specifically, documents on the interaction potential of new starting materials, active substances or excipients are required only if these are present in the medicinal product as active substances in a final dilution below D4. Any omission of such documentation must be justified.

6.5 Clinical documentation

6.5.1 Bibliographic clinical documentation

- Bibliographic documentation should demonstrate that:
  a) the composition can be sufficiently justified by use in the respective school of therapy, and familiarity in homeopathy or anthroposophic medicine can be proved for the requested indication;
  b) sufficient knowledge exists concerning possible adverse effects;
- If submitting bibliographic documentation to demonstrate therapeutic benefit, literature on the indication in the proposed school of therapy must be submitted for each individual active substance. In the case of books, at least the relevant chapter, the title page and publication details must be submitted. The literature must be assessed and referenced in the Summary (CTD Module 2.5.).
- The recognised sources for demonstrating therapeutic benefit are the monographs published by the German Commission D for homeopathic medicinal products and the monographs of the German Commission C for anthroposophic medicinal products, as well as the primary literature of the respective school of therapy. The Agency will verify on a case-by-case basis whether other literature sources can be recognised.
- Company publications, non-fiction books, articles not issued by specialist publishers, and books and publications that are not expressly concerned with the respective school of therapy will not be recognised as a matter of course.

6.5.2 Requirements for clinical documentation relating to use

- Apart from literature references, the evidence can also consist, for example, of scientifically prepared documentation on case reports, open trials or controlled clinical studies.
- For new homeopathic active substances, analyses of new homeopathic medicinal products can also be submitted.
The case reports, open trials or controlled studies must involve the corresponding school of therapy, and the results must be clearly attributable to the corresponding active substance or medicinal product.

Results of clinical trials (open trials/controlled clinical studies) are required in the case of a new use that is not adequately documented in the specialist literature for the respective school of therapy.

Demonstration by means of case reports or open trials requires at least 50 reports/open trials from several investigating physicians. Trials must be prepared, conducted and assessed in accordance with the requirements stated in Art. 14 paragraph 4 of the Ordinance on Clinical Trials (ClinO). Furthermore, trials must be conducted and assessed by qualified medical study investigators.

If the notified medicinal product is directly comparable with a medicinal product that has already been authorised, reference can be made to the published results of observational studies of the authorised medicinal product.

Sales figures on the documentation of patient exposure will not be accepted as proof for an indication, but they will be used for the purpose of calculating the incidence of adverse effects.

6.5.3 Requirements for the indication wording

Indications are recognised only if adequate evidence has been obtained from the corresponding school of therapy.

Requested indications may not be derived from the findings of other schools of therapy, nor may they be mixed with findings from phytotherapy or findings from allopathy.

Formulations on indications always start with “According to homeopathic pharmacology…” or “According to anthroposophical knowledge of humans and nature…” etc.

Indications that are considered to be obsolete on the basis of the latest medical and therapeutic findings for a complementary medicine cannot be recognised as such (e.g. immune disorders, defined psychiatric illnesses, cancers).

If a medicinal product is intended to be used in addition to another treatment for a particular indication, this should be specified accordingly (e.g. “in addition to a medically required treatment for…”)

6.5.4 Requirements for documentation on tolerability

6.5.4.1 Basic information

Any adverse effects that occur during investigations designed to demonstrate therapeutic benefit should be prepared and reported to the Agency in accordance with Instruction BW101 22 001e MB Safety relating to clinical trials – Compulsory notification.

Any adverse effects of the medicinal product or one of its constituents that are known from the literature or postmarketing experience should be described and assessed.

This requirement also applies if new, previously unknown adverse effects or risks emerge for substances and potencies included in the HAS List.

Swissmedic can determine, in individual cases, that documentation on tolerability does not need to be submitted if the medicinal product contains sufficiently diluted active substances that are known in homeopathy or anthroposophic medicine. In this case, substances included in the HAS List for the respective method of administration can be taken into account in the dilutions stated under “Notification procedure as of” or higher dilutions, as can substances not on this list in potencies from D12/C6, provided the HAS List accords with the latest scientific and technical findings for the corresponding substance.
and only contains known excipients, or carrier substances commonly used in homeopathic or anthroposophic medicine.

- Swissmedic decides whether documentation is not required on the basis of corresponding reasons.

### 6.5.4.2 Scope of documentation

For preparations with substances and potencies that are not included on the HAS List for the respective method of administration, or whose familiarity and efficacy cannot be adequately demonstrated in the corresponding school of therapy, the following documents relating to safety are expected:

- Provided the medicinal product is marketed in another country, the findings obtained in that country from postmarketing experience must be documented in a Safety Update Report and a risk assessment undertaken based on this report:
  - sales figures (packages or units sold per country and year)
  - information on how long the preparation has been on the market in the country in question and whether (and if so, what and why) quality-related changes have been made
  - known side effects, contraindications and interactions, as well as documentation of a corresponding systematic literature search
  - submission of the current or most recent valid medicinal product information from the country in question
  - submission of existing Periodic Safety Update Reports (PSUR).

- Provided the substances concerned are known from their use in other areas (e.g. medical device, foods, cosmetics), the findings obtained in these areas must be documented and a risk assessment undertaken based on these findings.

- If sufficient data and findings on the use of the substances or medicinal product are not available, the following documents must be submitted:
  - Medicinal products for topical use on the skin or mucous membranes require single and repeated testing for local tolerance and sensitising properties. To this end, at least 50 open trials by several investigating physicians are required as proof of clinical tolerance.
  - For medicinal products intended for parenteral administration, safety data in the form of clinical trials (open trials) with humans or the proposed target animal species and relating to the respective method of administration must be submitted.
  - For nosodes and organ preparations, safety in humans or in the proposed target animal species must also be investigated in addition to the testing of local tolerance in animals (for preparations for local or parenteral administration), taking account of immunological parameters for determining the risk of sensitisation.

### 6.5.5 Requirements for the summary of the clinical documentation (Clinical Expert Statement, e.g. in a CTD Module 2.5)

- This part should give a clear picture of the therapeutic benefit and clinical tolerability of the notified medicinal product. The name of the medicinal product, pharmaceutical form, method of administration, dosage and therapeutic use must be stated in an introduction. The summary should include a clear overview of the assessment of the clinical documentation with reference to all claimed uses and should be adequately referenced. It should present all results from clinical trials and the systematic literature search – including positive and negative findings – and assess the possible benefit and risks.
- If clinical trials (e.g. open trials) are submitted, the corresponding Study Reports must be submitted and a summary evaluation carried out. The author of the Expert Statement should be an expert who is qualified and trained in the relevant subject area.

### 6.6 Medicinal product information and package texts

#### 6.6.1 Special situations

In addition to the requirements of Ann. 2 no. 1 KPTPO, the requirements of the guidance document *Product information for human medicinal products HMV4*, the guidance document *Product information for veterinary medicinal products HMV4*, the guidance document *Packaging for human medicinal products HMV4* and the guidance document *Packaging texts for veterinary medicinal products HMV4*, as well as any Swissmedic publications that apply to the product information and package texts of complementary medicines, also apply.

### 7 Authorisation of homeopathic and anthroposophic medicinal products without indication

#### 7.1 Authorisation without indication with complete documentation (Art. 25 para. 2 KPTPO)

The statements on the authorisation of medicinal products with indication contained in Art. 24 KPTPO apply to the documentation to be submitted on quality and safety. The labelling and medicinal product information should be based on the requirements of Art. 26 para. 1 KPTPO.

#### 7.2 Authorisation with Reduced Dossier (Art. 25 para. 1 KPTPO)

##### 7.2.1 Preconditions for the application of the procedure

The preconditions set out in Art. 25 para. 1 KPTPO apply.

##### 7.2.2 Required documentation

Only the documentation specified in Ann. 3 KPTPO needs to be submitted for an application with a reduced dossier. Applications containing additional documents do not satisfy the formal requirements.
The following list is intended to facilitate the compilation of the required documents:

- Application for authorisation with reduced dossier with all documents per Ann. 3 no. 1, no. 1.1 KPTPO and per VZ Table of documents to be submitted.
- Substances in the MP are not included on the HAS List.
- Documents on familiarity should be submitted in accordance with Ann. 3 no. 2 KPTPO.
- The potency/dilution is below D12/C6.
- The potency/dilution is below D12/C6 and the active substance is not suitable for self-medication.
- The active substance exists in a potency/dilution that is not suitable for self-medication.
- Furthermore, active substance not monographed in the Pharmacopoeia, HAB, Ph.Eur., nor are they already approved by Swissmedic.
- The substances are manufactured using material of animal or human origin.
- Form "Substances of animal origin", if applicable, including the required documentation on TSE safety.
- For the substance on the HAS List, an * was entered next to the potency in the notification procedure, or the application concerns a substance for which a reduced dossier is required below D24/C12.
- The application concerns parenterals or MP for use on/in the eye.
- The application concerns parenterals or MP for use on/in the eye and the substance is subject to the Narcotics Control Ordinance and exists in a potency of D8/C4 or lower.
- Evidence of corresponding approval.
- Documents on safety and innocuousness should be submitted according to Ann. 3 no. 3 KPTPO.
- Documents on safety and innocuousness should be submitted according to Ann. 3 no. 3 KPTPO and the AS is more concentrated than the potency/dilution stated in the HAS List in the column "reduced dossier without documents on safety and innocuousness".
- Quality monograph corresponding to a pharmacopoeia monograph for a homeopathic substance.
- Company documentation on the HSs (**).
- Documents on tolerability should be submitted in accordance with Ann. 3 no. 4 KPTPO.
- Master Dossier on the manufacture of the pharmaceutical form per Ann. 3 no. 5 and on tolerability per Annex 3 no. 4 KPTPO.
- Documents on the general requirements for starting materials of animal or human origin (**).

Note: The documents listed in the oval boxes must be submitted with the application for authorisation with a reduced dossier.

** If applicable for several products, a template in the form of Master Dossiers is possible. For Master Dossiers the corresponding information stated in the WL Notification procedure for homeopathics and anthroposophics should be observed. It is also possible to submit Master Dossiers that apply both to products in the notification procedure and those with a reduced dossier. A reference to other valid Master Dossiers should be included in the documentation.
Authorisation with a reduced dossier is also possible for substances not included in the HAS List and for substances to be authorised at a lower potency than envisaged for a notification procedure.

As part of an application with reduced dossier, it is also possible to apply for a dispensing category other than that stated in the HAS List.

An application for authorisation with a reduced dossier can be accompanied by a simultaneous request to add the substances or potencies contained in the proposed preparation to the HAS List. (see guidance document Authorisation for homeopathic and anthroposophic medicinal products and medicinal products for gemmotherapy without indication in the notification procedure HMV4)

7.2.3 Information and texts on containers and packaging materials

Ann. 1a TPLRO applies. The following should be noted in particular:

- only scientific names are permitted as product names
- no indications or dosages may be stated. Instead, the following standard text should be stated: “For individual therapy, use and dosage as directed by the advising professional”
- all restrictions on use and warnings included in the HAS List for the respective substances, and any other known restrictions on use and warnings, must be listed.
- No other information beyond that specified in Annex 1a TPLRO or approved by Swissmedic is permissible.

According to Art. 26 para. 1 TPO, the information on the packaging must be drafted in at least two of Switzerland’s official languages.

The scientific name of the medicinal product approved by Swissmedic must not be modified and translations into the Swiss national languages must not be added.

If the space available on the container/packaging material is insufficient for all the information required by Ann. 1a TPLRO, the information stated in Ann.1a no. 1 para 1 let. h and j TPLRO can be omitted from the packaging texts. In this case, the missing information should be summarised in a package leaflet for enclosure with the medicinal product in the form of an information sheet in two of Switzerland's official languages (Art. 44 para. 2 KPTPO). However, this information may not contain information or figures that go beyond the scope of Annex 1a or that are ordered by Swissmedic.

As well as Annex 1a no. 1 para 2 and 3 TPLRO, the detailed information in section 5.3 of this guidance document should also be observed when declaring active substances. Examples can be found in Annex 8.2.

If the composition of the preparation is repeated on an enclosed information leaflet, a translation of the active substance names commonly used in the relevant school of therapy into the national languages may be omitted.

In addition to the active substances, all excipients contained in the preparation should also be declared in qualitative terms, regardless of their quantity. Furthermore, the vehicles used during manufacture/potentisation and other excipients should be listed as excipients if they account for at least 1% of the finished product or are necessary for the information required by Annex 3a TPLRO. The quantities of excipients of particular interest according to Annex 3a TPLRO should be listed. The constituents of excipient mixtures such as ethanol/water mixtures or sodium chloride solution must be listed separately. The total quantity of sodium in medicinal products that contain sodium (e.g. solutions for injection containing sodium chloride) must be listed separately.

If the composition is repeated on an enclosed information sheet, the excipients may be listed in the official languages on the information sheet only and in Latin on the packaging materials.
For liquid pharmaceutical forms, the required figures should be stated in ml.

If the medicinal product is dosed in drops, the drop equivalent should also be stated, as should the quantity delivered per spray pump for sprays and the number of globules per gram for globules.

Negative declarations (e.g. “lactose-free”) are not permitted. For excipients of particular interest according to Annex 3a TPLRO, the specified warnings should be adopted. Since a dosage is not stated for medicinal products without an indication, the information on ethanol required by Annex 3a TPLRO should be stated as a reference value per ml rather than as a dose (x mg per ml).

The labelling of medicinal products containing narcotics in dilutions up to, and including, D8/C4, must display the text “Subject to the Federal Act on Narcotics and Psychotropic Substances”. If an information leaflet is provided, this text should be placed immediately after the product name.

The marketing authorisation holder is responsible for the design and correct content of the packaging materials and information leaflet, if one is provided. The texts should be submitted as part of the authorisation application only in response to a specific request from Swissmedic.

The basic aspects of packaging materials design are subject to the requirements of the Guidance document *Packaging materials for human medicinal products HMV4* or the Guidance document *Packaging texts for veterinary medicinal products HMV4*. 
8 Annex

8.1 Decision tree for the application type for the authorisation of a complementary medicine

Part A: Complementary medicine with indication

Key:
- MP: Medicinal product
- CM: Complementary medicine
- Key features of the product
- Requirements

Application for authorisation of a complementary medicine

Is this an application for authorisation of a homeopathic, anthroposophic or spagyric CM or a Schüßler salt per Art. 4 para. 2 KPTPO?

- NO: Authorisation per Chapter 5 or 6 KPTPO

The specific requirements for CMs other than Asian CMs or those used in gemmothrapy should be clarified during a Pre-Submission Meeting.

- YES: Authorisation per Chapter 5 or 6 KPTPO

Is the MP to be placed on the market with an indication?

- NO: See Part B: CM without indication on next page

- YES: Part A: CM with indication

Have efficacy and safety been proven in clinical trials and will document protection be requested?

- YES: Authorisation per Art. 11 TPA

- NO: Variant A

Authorisation per Art. 24 para. 1 KPTPO in conjunction with Ann. 2 KPTPO

- Stated as 'homeopathic medicinal product' or 'anthroposophic medicinal product', etc.
- Indication according to therapeutic principle
- Dosage stated

- Requirements according to Chapter 6 of this WL

Requirements according to Chapter 6 of this WL

- Stated as 'homeopathic medicinal product' or 'anthroposophic medicinal product', etc.
- Indication according to therapeutic principle
- Dosage stated

Authorisation applications per Art. 14 para. 1 let. a bis-quater TPA

- Requirements according to WL: Authorisation applications per Art. 14 para. 1 let. a bis-quater TPA
- for quality documents, see Chapter 6.3 of this WL

- Stated as 'homeopathic medicinal product' or 'anthroposophic medicinal product', etc.
- Indication according to therapeutic principle
- Dosage stated
- Additional note on the type of authorisation and the testing for efficacy and safety in the medicinal product information per WL

- Stated as 'homeopathic medicinal product' or 'anthroposophic medicinal product', etc.
- Indication according to therapeutic principle
- Dosage stated
- Additional note on the type of authorisation and the testing for efficacy and safety in the medicinal product information per WL

Authorisation applications per Art. 14 para. 1 let. a bis-quater TPA

- Requirements according to WL: Authorisation applications per Art. 14 para. 1 let. a bis-quater TPA
- for quality documents, see Chapter 6.3 of this WL

- Stated as 'homeopathic medicinal product' or 'anthroposophic medicinal product', etc.
- Indication according to therapeutic principle
- Dosage stated
- Additional note on the type of authorisation and the testing for efficacy and safety in the medicinal product information per WL

- Stated as 'homeopathic medicinal product' or 'anthroposophic medicinal product', etc.
- Indication according to therapeutic principle
- Dosage stated
- Additional note on the type of authorisation and the testing for efficacy and safety in the medicinal product information per WL
Part B: Complementary medicine without indication

Is this application for the authorisation of an MP with dosage recommendation and/or with invented name?

- Simplified authorisation per Art. 25 para. 2 in conjunction with Ann. 2 no. 1, 2, 3, 5 KPTPO
- Preparation differs from an MP authorised per Art. 24 KPTPO only in the lack of indications
- Stated as "homeopathic medicinal product" or "anthroposophic medicinal product", etc.
- Standard text concerning individual therapy per Ann. 1a, para. 1 no. 1 let. h TPLRO

Are the substances included in the HAS List and do they fulfil the criteria stated in the "Notification procedure column relating to the minimal dilution/potency, or are substance and potency included in the SC List?

- Authorisation in notification procedure per Chapter 7 KPTPO
- Are the criteria for familiarity per Ann. 3 no. 2 KPTPO and for safety per Ann. 3 no. 3 KPTPO fulfilled?
- Simplified authorisation with reduced dossier per Art. 25 para. 1 in conjunction with Ann. 3 KPTPO
- Authorisation not possible

Requirements and documents, see WL Notification procedure for homeopathics and anthroposophics

- Scientific name as product name
- State school of therapy
- No PI/IHP
- Standard text concerning individual therapy per Ann. 1a, para. 1 no. 1 let. h TPLRO

Requirements and documents, see Chapter 7 of this WL

- Scientific name as product name
- State school of therapy
- No PI/IHP
- Standard text concerning individual therapy per Ann. 1a, para. 1 no. 1 let. h TPLRO
8.2 Sample declarations for medicinal products with and without indication

1. Medicinal product with indication and medicinal products without indication in the simplified authorisation procedure (Art. 25 para. 2 KPTPO)

A. Drops containing 3 active ingredients and additional excipients

- Form Full declaration HMV4:

  1 ml contains:
  
  - Atropa belladonna (Ph.Eur. Hom.) D4 0.1 ml
  - Arnica montana ex planta tota (HAB) D3 0.1 ml
  - Calcium fluoratum (HAB) D6 0.3 ml

  Contents from manufacture/potentisation:
  
  - Ethanol 96 % (Ph.Eur.) x mg
  - Purified water (Ph.Eur.) q.s
  - Lactose monohydrate (Ph.Eur.) x mg

  Excipients:
  
  - Ethanol 96 % (Ph.Eur.) y mg
  - Purified water (Ph.Eur.) y mg

  corresp. to Ethanolum xxx % (V/V)

  1 ml is equivalent to xxx drops.

- Medicinal product information (3 official languages)

  Composition
  
  1 ml (xxx drops) contains:
  
  Active substances:
  
  - Atropa belladonna (Ph.Eur. Hom.) D4 0.1 ml
  - Arnica montana ex planta tota (HAB) D3 0.1 ml
  - Calcium fluoratum (HAB) D6 0.3 ml

  Excipients: Ethanol, purified water, lactose monohydrate x mg. Contains xxx vol% alcohol.

  Warning for ethanol and lactose monohydrate in accordance with Annex 3a TPLRO.

- Packaging materials to match (2 official languages, Latin names permissible for excipients) but without warning in accordance with Annex 3a TPLRO.

B. Solution for injection containing 2 active ingredients and additional excipients

- Form Full declaration HMV4:

  One 1 ml ampoule contains:
  
  Active substances:
  
  - Colchicum autumnale (HAB) D12 0.6 ml
  - Crotalus horridus (HAB) D6 0.2 ml

  Contents from manufacture/potentisation:
  
  Sodium chloride (Ph.Eur.) x mg
Guidance document
Authorisation of Homeopathics, anthroposophics
and other complementary medicinal products HMV4

Water for injection purposes (Ph.Eur.) y ml
Glycerol (Ph.Eur.) z ml

Excipients:
Water for injection purposes (Ph.Eur.) 0.2 ml
Sodium chloride (Ph.Eur.) xx mg (isotonising agent)

Equivalent to sodium x mg

- Medicinal product information (3 official languages)

Composition:
1 ampoule contains:
Active substances:
- Colchicum autumnale (HAB) D12 0.6 ml
- Crotalus horridus (HAB) D6 0.2 ml
Excipients: Water for injection purposes, sodium chloride, glycerol.

Warning for sodium in accordance with Annex 3a TPLRO.

- Packaging materials to match (2 official languages, Latin names permissible for excipients) but without warning in accordance with Annex 3a TPLRO.

C. Globules containing 2 active substances

- Form Full declaration HMV4:

1 g of globules contains, in processed form:
Active substances:
- Apocynum cannabinum (HAB) D12 5 mg (or 0.005 g)
- Calcium sulfuricum (HAB) D12 5 mg
Excipients
- Sucrose (Ph.Eur.) 1 g (vehicle)

1 g is equivalent to 110 – 130 globules.

It is no longer possible to state percentages in full declarations.

- Medicinal product information (3 official languages)

1 g of globules contains, in processed form: Active substances: Apocynum cannabinum (HAB) D12 10 mg, Calcium sulfuricum (HAB) D12 5 mg
Excipient: 1 g sucrose

1 g is equivalent to 110 – 130 globules.

It is no longer possible to state percentages in full declarations.

Warning for sucrose in accordance with Annex 3a TPLRO.

- Packaging materials to match (2 official languages, Latin names permissible for excipients) but without warning in accordance with Annex 3a TPLRO.
2. **Medicinal product without indication authorised with reduced dossier (Art. 25 para. 1 KPTPO) or by the notification procedure (chap. 7 KPTPO)**

Principles: As regards requirements for active substance names, the requirements and examples listed under 5.3 should be observed in addition to Ann. 1a para. 1 no. 2 and 3 TPLRO.

If the active substance name commonly used in the school of therapy is to be listed as well, only the names or synonyms listed in the HAS List (Annex 6 KPTPO) and gemmotherapy list (Annex 8 KPTPO) are permissible (cf. sample packaging texts B and C). Names must not be translated into the official languages.

Example for homeopathy:

Atropa belladonna (Ph.Eur. Hom.) D6 or
Atropa belladonna (Belladonna) (Ph. Eur. Hom.) D6

Example for gemmotherapy:

Castanea sativa e gemma D1 (Ph. Eur. Hom. 2.1.3) or
Castanea sativa e gemma (Castanea vesca e gemma) D1 (Ph. Eur. Hom. 2.1.3)

A. **Homeopathic medicinal product with 4 active substances and excipients from manufacturing, drops**

- Declaration on packaging elements (in 2 official languages)
  
  Zusammensetzung/Composition:
  
  1 ml (xxx Tropfen) enthält/contient Wirkstoffe/principes actif:
  
  Atropa belladonna (Ph.Eur. Hom.) D6 0.3 ml
  
  Arnica montana ex planta tota (HAB) D8 0.3 ml
  
  Calcium fluoratum (HAB) D12 0.2 ml
  
  Sennae folium (Ph.Eur.) D8 (Ph. Eur.Hom. 1.1.8) 0.2 ml
  
  Hilfsstoffe/excipients: Ethanol 96 %/d'éthanol, Gereinigtes Wasser/de l'eau purifiée
  
  Enthält/contient: … Vol.-% Alkohol/d'alcool
  
  1 ml is equivalent to xxx drops. or 1 ml is equivalent to xxx spray pumps.

  Warning for ethanol in accordance with Annex 3a TPLRO, reference value per ml.

B. **Homeopathic medicinal product with 1 active substance and excipients from manufacturing, globules**

- Declaration on packaging elements (in 2 official languages, Art. 26 para. 1 TPO)
  
  Composition:
  
  1 g of globules contains, in processed form:
  
  Active substance: Belladonna (Ph. Eur. Hom.) D6 10 mg
  
  Excipient: Sucrose 1g
  
  1 g is equivalent to 110 – 130 globules.

  Composition:
  
  1 g of globules contains, in processed form/1 g de globules contient :
  
  Principe actif: Belladonna (Ph. Eur. Hom.) D6 10 mg
Excipient: Sucrose 1g
1 g is equivalent to 110 – 130 globules.

Warning for sucrose in accordance with Annex 3a TPLRO.

C. Medicinal product for use in gemmotherapy containing 1 active substance and excipients from manufacturing, spray for use in the buccal cavity

- Declaration on packaging elements (in 2 official languages, Art. 26 para. 1 TPO)

Zusammensetzung/Composition:
1 ml Flüssigkeit enthält/contient Wirkstoff/principe actif:
Ribes nigrum e gemma D1 (Ph. Eur.Hom. 2.1.3) 1 ml
Hilfsstoffe/excipients: Ethanol/d’éthanol, Glycerin/de glycérol, Gereinigtes Wasser/de l’eau purifiée
Enthält/contient: … Vol.-% Alkohol/d’alcool
1 ml is equivalent to xxx spray pumps.

Warning for ethanol in accordance with Annex 3a TPLRO, reference value per ml.

D. Medicinal product for use in Schüssler therapy with 1 active substance and excipients from manufacturing, tablet

- Declaration on packaging elements (in 2 official languages, Art. 26 para. 1 TPO)

Zusammensetzung/Composition:
One xxx mg contains in processed form/1 comprimé à xxx mg contient:
Wirkstoff/Principe actif:
№ 11 Silicea (HAB) D12 250 mg;
Hilfsstoffe/Excipients: Lactose-Monohydrat/Lactose monohydraté 250 mg,
Magnesiumstearat/stéarate de magnésium, Kartoffelstärke/amidon de pomme de terre

Warning for sucrose in accordance with Annex 3a TPLRO, reference value per g.