



EUROPEAN MEDICINES AGENCY  
SCIENCE MEDICINES HEALTH

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Executive Director

## Explanatory note on fees payable to the European Medicines Agency

The fees, fee exemptions and definitions described in this Explanatory Note apply as of 1 April 2013.

The latest amendments to the Rules for the implementation of Regulation (EC) No 297/95 on fees payable to the European Medicines Agency and other measures<sup>1</sup> were last adopted by the Agency's Management Board on 21 March 2013.

### Disclaimer:

This Explanatory Note is meant as a guidance note only. In case of discrepancies between the text and amounts of fees payable to the Agency quoted in the Explanatory Note and the provisions of the Fee Regulation (EC) No 297/95 and its Implementing Rules adopted by the Agency's Management Board, the latter documents prevail.

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<sup>1</sup> Article 11(2) of Regulation (EC) No 297/95.



### ***Changes introduced in this version (1 April 2013)***

- Increase in level of fees and charges throughout the document in line with the inflationary increase of 2.6% as per Commission Regulation (EU) 220/2013 of 13.03.2013<sup>2</sup> and as per the amendments to the Rules for the implementation of Regulation (EC) No 297/95 on fees payable to the European Medicines Agency of 21.03.2013.
- Revision of section 1.1.3.1 (level II fee) and section 2.1.3.1 (level II and III fees) to clarify the definitions of quality extensions of marketing authorisations by including references to non-clinical data.
- Correction in section 1.1.4.3 (level II fee) and section 2.1.4.3 (level II fee) to add reference to non-clinical data in relation to previously submitted data.
- Revision of section 1.3.1 to clarify the fees applicable by analogy to scientific opinions pursuant to Article 58 of Regulation (EC) No 726/2004 and the applicability of SME incentives.
- Addition of new basic fee level in section 1.4.2 for grouped amendments to the documentation on an ancillary medicinal substance or blood derivative incorporated in a medical device.
- Updating of sections 1.4.2, 1.5.2 and 1.5.5 to add references to Article 7(2)(c) of Commission Regulation (EC) No 1234/2008.
- Revision of section 3.3 on notification of parallel distribution following the revision of the Rules for the implementation of the fee regulation.

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<sup>2</sup> OJ L 70, 14.3.2013, p. 1

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# 1. Medicinal products for human use

Note: This section on fees for medicinal products for human use should be read in conjunction with the section on fee exemptions (Section 5) and the annex.

## 1.1. Centralised procedure

### 1.1.1. Definitions

*Pharmaceutical form:* According to the 'Standard Terms' published by the Council of Europe.

*Strength:* See [definition](#) used in the pre-submission guidance.

*Presentation:* Each unit/entity of a certain strength and form of a pharmaceutical product which will be individually authorised and eventually marketed (= each individual sub-number).

### 1.1.2. Application for a marketing authorisation

#### 1.1.2.1. Application for which a full dossier needs to be presented

Basic fee	274 400 EURO For a single strength associated with one pharmaceutical form and one presentation.
Additional fee	+ 27 500 EURO For each additional strength or pharmaceutical form including one presentation, submitted at the same time as the initial application for authorisation.
	+ 6 900 EURO For each additional presentation of the same strength and pharmaceutical form, submitted at the same time as the initial application for authorisation.

#### 1.1.2.2. Application for which a full dossier need not be presented

Basic fee	177 300 EURO For an application for a marketing authorisation pursuant to Article 10(4) of Directive 2001/83/EC. This fee is for a single strength associated with one pharmaceutical form and one presentation.
Additional fee	+ 10 600 EURO For each additional strength or pharmaceutical form including one presentation submitted at the same time as the initial application for authorisation.
	+ 6 900 EURO For each additional presentation of the same strength and pharmaceutical form, submitted at the same time as the initial application for authorisation.

Basic fee	106 500 EURO For applications for a marketing authorisation pursuant to Article 10(1), Article 10(3) and Article 10c of Directive 2001/83/EC. This fee is for a single strength associated with one pharmaceutical form and one presentation.
Additional fee	+ 10 600 EURO For each additional strength or pharmaceutical form including one presentation submitted at the same time as the initial application for authorisation.
	+ 6 900 EURO For each additional presentation of the same strength and pharmaceutical form, submitted at the same time as the initial application for authorisation.

**Note:** Refer to section [A.1.1.2.](#) in the annex for examples of the determination of fees for applications for marketing authorisation.

### 1.1.3. Extension of a marketing authorisation

#### 1.1.3.1. Extension of a marketing authorisation under Annex I to Commission Regulation (EC) No 1234/2008

Basic fee (Level I)	82 400 EURO For each extension of a marketing authorisation within the meaning of Annex I to Commission Regulation (EC) No 1234/2008 concerning the examination of variations to the terms of a marketing authorisation for medicinal products for human use. This fee is for a single strength/potency associated with one pharmaceutical form and one presentation.
Additional fee	+ 20 600 EURO For each additional strength/potency of the same pharmaceutical form of the same extension including one presentation submitted at the time of the extension application.
Additional fee	+ 6 900 EURO For each additional presentation of the same strength/potency of the same pharmaceutical form of the same extension submitted at the time of the extension application.

Basic fee (Level II)	61 800 EURO For all quality extensions (i.e. requiring chemical, pharmaceutical and biological documentation) for which no clinical or non-clinical data are submitted or no cross-references to previously submitted clinical or non-clinical data are made by the MAH. This fee is for a single strength/potency associated with one pharmaceutical form and one presentation. (Note: Bioequivalence data qualify as clinical data. Biowaiver dossiers are not considered as clinical data.)
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Additional fee	+20 600 EURO For each additional strength/potency of the same pharmaceutical form of the same extension including one presentation submitted at the time of the extension application.
Additional fee	+ 6 900 EURO For each additional presentation of the same strength/potency of the same pharmaceutical form of the same extension submitted at the time of the initial extension application.

**1.1.3.2. Extension of a marketing authorisation for use in the paediatric population under Article 29 of Regulation (EC) No 1901/2006**

Basic fee (Level III)	82 400 EURO For each extension of a marketing authorisation made under Article 29 of Regulation (EC) No 1901/2006 on medicinal products for paediatric use. This fee is for a single strength/potency associated with one pharmaceutical form and one presentation.
Additional fee	+ 20 600 EURO For each additional strength/potency of the same pharmaceutical form of the same extension including one presentation submitted at the time of the extension application.
Additional fee	+ 6 900 EURO For each additional presentation of the same strength/potency of the same pharmaceutical form of the same extension submitted at the time of the initial extension application.

**Note:** Refer to section [A.1.1.3.](#) in the annex for examples of the determination of fees for extensions of marketing authorisation.

**1.1.4. Variation to a marketing authorisation**

The following fees apply to variations that are notified or applied for individually in a notification or application.

**1.1.4.1. Type-IA variation**

Basic fee	3 000 EURO For a minor variation to a marketing authorisation, as defined in Article 2(2) of Commission Regulation (EC) No 1234/2008.
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#### 1.1.4.2. Type-IB variation

Basic fee	6 900 EURO For a minor variation to a marketing authorisation, as defined in Article 2(5) of Commission Regulation (EC) No 1234/2008.
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- In the event of the same variation being introduced, the fee will cover all authorised strengths, pharmaceutical forms and presentations of a given medicinal product (= per main authorisation number).

#### 1.1.4.3. Type-II variation

Basic fee (Level I)	82 400 EURO For a major variation to a marketing authorisation, as defined in Article 2(3) of Commission Regulation (EC) No 1234/2008.
Basic fee (Level II)	61 800 EURO For a quality variation (i.e. amendments to the chemical, pharmaceutical and biological documentation) for which no clinical or non-clinical data are submitted or no cross-references to previously submitted clinical or non-clinical data are made by the MAH. (Note: Bioequivalence data qualify as clinical data. Biowaiver dossiers are not considered as clinical data.)
Basic fee (Level III)	20 600 EURO For each of the third and subsequent type II variations that are grouped in a single application made under the terms of Article 7(2) of Commission Regulation (EC) No 1234/2008. For each of the third and subsequent type II variations to the centralised marketing authorisation or to one of the centralised marketing authorisations, where there is more than one centralised marketing authorisation, in the case of a worksharing application made under the terms of Article 20 of Commission Regulation (EC) No 1234/2008.

### 1.1.5. Grouping and worksharing procedures for variations

#### 1.1.5.1. Grouping of extensions and/or variations notified or submitted under the terms of Article 7(2) of Commission Regulation (EC) No 1234/2008

- The applicable fee specified in sections 1.1.3 and 1.1.4 shall be payable for each individual extension and/or variation to a marketing authorisation that is grouped in a single notification or a single application.
- The applicable level I and level II basic fees specified in sub-section 1.1.4.3 above are payable for the first and second type II variation respectively when both levels of fees are applicable to variations in the same grouping.
- Consequential variations in a grouping shall be similarly charged the applicable fees as specified above.
- In the case of grouping of the same Type IA variations to the terms of several marketing authorisations owned by the same holder (as set out in Article 7(2)(a) of Commission Regulation



(EC) No 1234/2008), the applicable fee shall be payable for each individual Type IA variation and for each marketing authorisation in the grouping.

- The same marketing authorisation holder also means several marketing authorisation holders that are linked through a parent company. The fee for the grouping shall be payable by the marketing authorisation holder applying for the grouping procedure.
- Where any extensions/variations included in a grouping are found not to be valid and the remainder are validated positively, the applicable fees as specified above shall be payable for each of the positively validated extensions/variations.

**1.1.5.2. Worksharing procedure under the terms of Article 20 of Commission Regulation (EC) No 1234/2008**

- The applicable fee specified in section 1.1.4 is payable for each individual variation to one of the centralised marketing authorisations owned by the same holder, where more than one centralised marketing authorisation is included in the worksharing application, or to the single centralised marketing authorisation included in the worksharing application, as applicable.
- The applicable level I and level II basic fees specified in 1.1.4.3 above are payable for the first and second type II variation respectively when both levels of fees are applicable to variations in the same worksharing procedure.
- The administrative fee shown in the table below is additionally payable for each individual variation to each of the other centralised marketing authorisation(s) owned by the same holder included in the same worksharing application, if applicable.
- No fee is payable to the Agency for any national marketing authorisations included in the same worksharing application.
- The same marketing authorisation holder also means several marketing authorisation holders that are linked through a parent company. The fee for the worksharing procedure shall be payable by the marketing authorisation holder applying for the procedure.

Variation type	Worksharing fees for one centralised marketing authorisation	Worksharing administrative fees	
		Other centralised marketing authorisations (excluding multiple applications on usage patent grounds)	Other centralised marketing authorisations (multiple applications on usage patent grounds <sup>3</sup> )
Type II (Level I)	82 400 EURO	6 870 EURO	3 940 EURO
Type II (Level II)	61 800 EURO		
Type II (Level III)	20 600 EURO		
Type IB	6 900 EURO	1 140 EURO	1 140 EURO
Type IA	3 000 EURO	570 EURO	570 EURO

- Where any variations included in a worksharing procedure are found not to be valid and the remainder are validated positively, the applicable fees as specified above shall be payable for each of the positively validated variations.

<sup>3</sup> These fees apply to generic medicinal product applications, hybrid applications and similar biological medicinal product applications and are applicable for as long as the concerned marketing authorisation is affected by usage patent(s) pertaining to indication(s) and/or dosage form(s). Refer to section 4.3 for further details on fee exemptions.

### 1.1.6. Renewal of a marketing authorisation

Basic fee	13 600 EURO For each strength associated with a pharmaceutical form for which renewal is sought.
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**Note:** Refer to section [A.1.1.6.](#) in the annex for examples of the determination of fees for renewals of marketing authorisation.

### 1.1.7. Inspection

Basic fee (Level I)	20 600 EURO For each inspection inside or outside the European Union; for inspections outside the European Union, travel expenses shall be charged extra on the basis of actual cost.
Basic fee (Level II)	10 300 EURO For each consecutive distinct plasma master file (PMF) inspection performed in conjunction with an inspection that attracts the level I fee, provided that such consecutive inspection concerns the same PMF application, the same inspection team and is conducted in the same PMF inspection tour.

**Note:** Refer to section [A.1.1.7.](#) in the annex for examples of the determination of fees for good manufacturing practice (GMP) inspections.

### 1.1.8. Transfer of a marketing authorisation

Basic fee	6 900 EURO For all authorised presentations of the medicinal product.
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### 1.1.9. Maintenance of a marketing authorisation – Annual fee

Basic fee (Level I)	98 400 EURO For each marketing authorisation of a medicinal product. This fee covers all authorised presentations of the medicinal product.
Basic fee (Level II)	49 200 EURO For each marketing authorisation of a biosimilar medicinal product (Article 10(4) of Directive 2001/83/EC). This fee covers all authorised presentations of the medicinal product.
Basic fee (Level III)	24 500 EURO For each marketing authorisation of a generic, hybrid or informed consent medicinal product (Articles 10(1), 10(3) and 10c of Directive 2001/83/EC). This fee covers all authorised presentations of the medicinal product.

### 1.1.10. Referral

Basic fee	68 400 EURO For procedures laid down in Article 30(1) and 31 of Directive 2001/83/EC that are initiated by the applicant of a marketing authorisation or the holder of an existing marketing authorisation.
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**Note:** Where more than one applicant for marketing authorisation or holder of existing marketing authorisations are concerned by the procedures referred to in the first subparagraph, the applicants or holders may be grouped for the purpose of the payment of one single referral fee. If, however, the same procedure concerns more than 10 different applicants or holders, the fee will be charged by the application of the above mentioned referral fee.

## 1.2. Scientific advice

### 1.2.1. Definitions

*Quality development:* Chemical, pharmaceutical and biological testing.

*Safety development:* Toxicological and pharmacological tests.

*Clinical development:* Studies in human subjects, whether patients or non-patient volunteers, including clinical pharmacological trials designed to determine the efficacy and safety of the product.

*Qualification advice:* Advice on the acceptability of a specific use of the proposed method (e.g. use of a biomarker) in a research and development (R&D) context (non-clinical or clinical studies), based on the assessment of submitted data.

*Initial request:* First request for scientific advice or protocol assistance introduced in relation to the submission of an application for marketing authorisation or a variation, whatever the authorisation phase (pre- or post-authorisation).

*Follow-up to initial request:* Any subsequent request falling within the same therapeutic indication and initial area(s) as the initial request, (area meaning quality, preclinical and/or clinical development, including pharmacovigilance/risk management aspects).

### 1.2.2. Initial request for scientific advice

Basic fee (Level III)	82 400 EURO For <b>initial</b> requests for scientific advice on: <ul style="list-style-type: none"><li>• Quality, and safety and clinical development, or</li><li>• quality and clinical development, or</li><li>• safety and clinical development, or</li><li>• qualification advice.</li></ul>
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<p>Basic fee (Level II)</p>	<p>61 800 EURO</p> <p>For <b>initial</b> requests on:</p> <ul style="list-style-type: none"> <li>• clinical development, or</li> <li>• quality and safety development, or</li> <li>• quality and bioequivalence studies for generic medicinal products.</li> </ul>
<p>Basic fee (Level I)</p>	<p>41 100 EURO</p> <p>For <b>initial</b> requests on:</p> <ul style="list-style-type: none"> <li>• quality development or</li> <li>• safety development, or</li> <li>• bioequivalence studies for generic medicinal products.</li> </ul>

### 1.2.3. Follow-up request for scientific advice

<p>Basic fee (Level III)</p>	<p>41 100 EURO</p> <p>For <b>follow-up</b> to the initial request on:</p> <ul style="list-style-type: none"> <li>• quality and safety and clinical development, or</li> <li>• quality and clinical development, or</li> <li>• safety and clinical development, or</li> <li>• qualification advice.</li> </ul>
<p>Basic fee (Level II)</p>	<p>30 900 EURO</p> <p>For <b>follow-up</b> to the initial request on:</p> <ul style="list-style-type: none"> <li>• clinical development, or</li> <li>• quality and safety development, or</li> <li>• quality and bioequivalence studies for generic medicinal products.</li> </ul>
<p>Basic fee (Level I)</p>	<p>20 600 EURO</p> <p>For <b>follow-up</b> to the initial request on:</p> <ul style="list-style-type: none"> <li>• quality development, or</li> <li>• safety development, or</li> <li>• bioequivalence studies for generic medicinal products.</li> </ul>

### **1.3. Scientific services**

A scientific service fee shall apply where an application is made for any scientific advice or opinion by a scientific Committee, which is not covered by Articles 3 to 7 or by Article 8(1).

#### **1.3.1. Scientific opinions pursuant to Article 58 of Regulation (EC) No 726/2004**

The basic and additional fees specified in section 1.1.2 apply by analogy for a scientific opinion for the evaluation of medicinal products for human use intended exclusively for markets outside the European Union. Likewise the inspection fees in section 1.1.7 apply by analogy to any inspection undertaken for the purpose of assessment prior to an opinion.

Fees for post-opinion services and annual fees are charged according to the corresponding fees for centrally authorised products. Therefore the fees specified in sections 1.1.3, 1.1.4, 1.1.5, 1.1.7 and 1.1.9 apply by analogy.

The fee incentives for micro, small and medium-sized enterprises apply to scientific services described in section 5.1.2. However fee deferrals and conditional fee exemptions do not apply to services in relation to scientific opinions pursuant to Article 58 of Regulation (EC) No 726/2004.

#### **1.3.2. Compassionate use**

Basic fee	137 300 EURO For any opinion on medicinal products for compassionate use
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#### **1.3.3. Herbal medicinal products**

Basic fee (Level I)	20 600 EURO For requests for scientific support and advice by the Committee on Herbal Medicinal Products (HMPC) on multiple areas related to traditional herbal medicinal products
Basic fee (Level II)	13 600 EURO For requests for scientific support and advice by the HMPC on single areas related to traditional herbal medicinal products, for example: <ul style="list-style-type: none"><li>• quality, or</li><li>• safety, or</li><li>• long-standing use.</li></ul>

#### **1.3.4. Certification of quality and non-clinical data relating to advanced therapy medicinal products developed by micro, small and medium-sized enterprises**

Basic fee (Level I)	61 800 EURO Evaluation of an application relating to quality and non-clinical data.
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Basic fee (Level II)	41 100 EURO Evaluation of an application relating to quality data.
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- The above fees are subject to the fee reduction for scientific services applicable to micro, small and medium-sized enterprises (SMEs) in accordance with section [5.1.2](#).

#### **1.4. Consultation on ancillary substances including blood derivatives incorporated in medical devices**

##### **1.4.1. Initial request**

Basic fee (Level I)	82 400 EURO For consultation on an ancillary medicinal substance or blood derivative new to the centralised procedure. This fee applies where the substance/derivative from the specified manufacturer has not been evaluated by the Agency in connection with a previous marketing authorisation and/or through a previous successful notified body consultation.
Basic fee (Level II)	61 800 EURO For consultation on a known ancillary blood derivative from a known source. This fee applies where the blood derivative from the specified manufacturer has been evaluated by the Agency in connection with a previous marketing authorisation and/or through a previous successful notified body consultation.
Basic fee (Level III)	41 100 EURO For consultation on a known ancillary medicinal substance from a known source. This fee applies where the substance from the specified manufacturer has been evaluated by the Agency in connection with a previous marketing authorisation and/or through a previous successful notified body consultation.

##### **1.4.2. Follow-up to the initial request**

Basic fee	20 600 EURO For consultation on a known ancillary medicinal substance from a known source. This fee applies where the substance from the specified manufacturer has been evaluated by the Agency in connection with a previous marketing authorisation and/or through a previous successful notified body consultation. In this case a further consultation is requested by a notified body after a first consultation, i.e. when additional data are submitted to the Agency for evaluation in response to a list of deficiencies notified in an initial Agency report.
Basic fee	41 100 EURO For consultation on an amendment to the documentation on an ancillary medicinal substance or blood derivative already evaluated by the Agency (The amendments should be classified by analogy to Annex I of Commission Regulation (EC) No 1234/2008).

Basic fee	41 100 EURO For consultation on a major amendment to the documentation on an ancillary medicinal substance or blood derivative already evaluated by the Agency (The amendment should be equivalent to a Type II variation as classified by analogy to Commission Regulation (EC) No 1234/2008).
Basic fee	41 100 EURO For consultation on two or more amendments to the documentation on an ancillary medicinal substance or blood derivative already evaluated by the Agency that are grouped in a single application by analogy to the terms of Articles 7(2)(b) or 7(2)(c) of Commission Regulation (EC) No 1234/2008 where at least one of the amendments is classified by analogy to Annex I of Commission Regulation (EC) No 1234/2008 or as a major amendment.
Basic fee	6 900 EURO For consultation on a minor amendment to the documentation on an ancillary medicinal substance or blood derivative already evaluated by the Agency. (The amendment should be equivalent to a Type IB variation as classified by analogy to Commission Regulation (EC) No 1234/2008).
Basic fee	3 000 EURO For consultation on a minor amendment to the documentation on an ancillary medicinal substance or blood derivative already evaluated by the Agency. (The amendment should be equivalent to a Type IA variation as classified by analogy to Commission Regulation (EC) No 1234/2008).

## **1.5. Certification of compliance with European Union legislation for plasma master files (PMF) and vaccine antigen master files (VAMF)**

### **1.5.1. Application for PMF certification (initial certification)**

#### **1.5.1.1. Not submitted simultaneously with a new application under the centralised procedure**

Basic fee (Level I)	68 400 EURO For the review of the PMF and its initial certification where the data contained in the PMF have not been previously evaluated within the centralised procedure.
Basic fee (Level II)	61 800 EURO For the review of the PMF and its initial certification where the PMF applicant has included change(s) to the data previously evaluated within the centralised procedure and which are now part of the PMF application.
Basic fee (Level III)	20 600 EURO For the review of the PMF and its initial certification where the data contained in the PMF have been previously evaluated under the centralised procedure and no changes have been included.

### **1.5.1.2. Submitted simultaneously with a new application under the centralised procedure**

Basic fee	6 900 EURO  For the review of the PMF and its certification when it is submitted in parallel and within the submission of a new application within the centralised procedure. The PMF documentation will be evaluated by the Agency simultaneously with a centralised marketing authorisation application.
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### **1.5.2. Variation to a certified PMF**

Basic fee	61 800 EURO  For the review and certification of a major variation to the PMF (in accordance with Commission Regulation (EC) No 1234/2008).
Basic fee	61 800 EURO  For the review and certification of two or more variations that are grouped in a single application made under the terms of Articles 7(2)(b) or 7(2)(c) of Commission Regulation (EC) No 1234/2008 where at least one of the variations is a major variation.
Basic fee	6 900 EURO  For the review and certification of a minor variation of type IB to the PMF (in accordance with Commission Regulation (EC) No 1234/2008).
Basic fee	3 000 EURO  For the review and certification of a minor variation of type IA to the PMF (in accordance with Commission Regulation (EC) No 1234/2008).

### **1.5.3. Annual re-certification of PMF**

Basic fee	61 800 EURO  For the review and annual re-certification of the PMF under this scheme where one or more variations are included in the submitted documentation and at least one of the variations is a major variation.
Basic fee	13 600 EURO  For the review and annual re-certification of the PMF under this scheme where no major variations are included in the submitted documentation.  This fee is increased by the applicable fee for each minor variation of type IA or type IB included in the submitted documentation, up to a maximum total fee of 61 800 EURO.



## 1.5.4. Application for a VAMF certification (initial certification)

### 1.5.4.1. Not submitted simultaneously with a new application under the centralised procedure

Basic fee	68 400 EURO For the review of the VAMF and its certification where the data contained in the vaccine antigen master file have not been previously evaluated within the centralised procedure. In the case of a group of antigens aimed at preventing a single infectious disease, e.g. inactivated polio serotypes 1, 2 and 3, the VAMF application for one antigen will be charged at 68 400 EURO.
Additional fee	+ 6 900 EURO For each VAMF application submitted simultaneously for antigens from the same group, up to a maximum total fee of 82 400 EURO.

Basic fee	61 800 EURO For the review of the VAMF and its certification where the initial data have been previously evaluated within the centralised procedure but where the VAMF applicant has included changes or harmonisation as part of the VAMF certification scheme. In the case of a group of antigens aimed at preventing a single infectious disease e.g. inactivated polio serotypes 1, 2 and 3, the VAMF application for one antigen will be charged at 61 800 EURO.
Additional fee	+ 6 900 EURO For each VAMF application submitted simultaneously for antigens from the same group, up to a maximum total fee of 82 400 EURO.

Basic fee	20 600 EURO For the review of the VAMF and its initial certification where the data contained in the vaccine antigen master file have been previously evaluated under the centralised procedure and where no changes or harmonisation have been included. In the case of a group of antigens aimed at preventing a single infectious disease e.g. Inactivated Polio serotypes 1, 2 and 3, the VAMF application for one antigen will be charged at 20 600 EURO.
Additional fee	+ 6 900 EURO For each VAMF application submitted simultaneously for antigens from the same group, up to a maximum total fee of 82 400 EURO.

**1.5.4.2. Submitted simultaneously with a new application under the centralised procedure**

Basic fee	6 900 EURO For the review of the VAMF and its certification when it is submitted in parallel and within the submission of a new application containing the named antigen within the centralised procedure. In the case of a group of antigens aimed at preventing a single infectious disease e.g. inactivated polio serotypes 1, 2 and 3, the VAMF application for one antigen will be charged at 6 900 EURO.
Additional fee	+ 6 900 EURO For each VAMF application submitted simultaneously for antigens from the same group, up to a maximum total fee of 82 400 EURO.

**1.5.5. Variation to a certified VAMF**

Basic fee	61 800 EURO For the review and certification of a major variation to the VAMF (in accordance with Commission Regulation (EC) No 1234/2008). In the case of a group of antigens aimed at preventing a single infectious disease e.g. inactivated polio serotypes 1, 2 and 3, the VAMF application for one antigen will be charged at 61 800 EURO.
Additional fee	+ 6 900 EURO For each VAMF application submitted simultaneously for antigens from the same group, up to a maximum total fee of 82 400 EURO.

Basic fee	61 800 EURO For the review and certification of two or more variations that are grouped in a single application made under the terms of Articles 7(2)(b) or 7(2)(c) of Commission Regulation (EC) No 1234/2008 where at least one of the variations is a major variation. In the case of a group of antigens aimed at preventing a single infectious disease e.g. inactivated polio serotypes 1, 2 and 3, the grouped variations application for one antigen will be charged at 61 800 EURO.
Additional fee	+ 6 900 EURO For each VAMF grouping application submitted simultaneously for antigens from the same group, up to a maximum total fee of 82 400 EURO.

Basic fee	<p>6 900 EURO</p> <p>For the review and certification of a minor variation of type IB to the VAMF (in accordance with Commission Regulation (EC) No 1234/2008).</p> <p>In the case of a group of antigens aimed at preventing a single infectious disease e.g. inactivated polio serotypes 1, 2 and 3, the VAMF application for one antigen will be charged at 6 900 EURO.</p>
Additional fee	<p>+ 6 900 EURO</p> <p>For each VAMF applications submitted simultaneously for antigens from the same group, up to a maximum total fee of 41 100 EURO.</p>

Basic fee	<p>3 000 EURO</p> <p>For the review and certification of a minor variation of type IA to the VAMF (in accordance with Commission Regulation (EC) No 1234/2008).</p> <p>In the case of a group of antigens aimed at preventing a single infectious disease e.g. inactivated polio serotypes 1, 2 and 3, the VAMF application for one antigen will be charged at 3 000 EURO.</p>
Additional fee	<p>+ 3 000 EURO</p> <p>For each VAMF application submitted simultaneously for antigens from the same group, up to a maximum total fee of 20 600 EURO.</p>

## 2. Medicinal products for veterinary use

Note: This section on fees for veterinary medicinal products should be read in conjunction with the section on fee exemptions (Section 5) and the annex.

### 2.1. Centralised procedure

#### 2.1.1. Definitions

*Pharmaceutical form:* According to the 'Standard Terms' published by the Council of Europe.

*Strength:* See [definition](#) used in the pre-submission guidance.

*Presentation:* Each unit/entity of a certain strength and form of a pharmaceutical product which will be individually authorised and eventually marketed (= each individual sub-number).

#### 2.1.2. Application for a marketing authorisation

##### 2.1.2.1. Full fee

Basic fee	137 300 EURO For a single strength associated with one pharmaceutical form and one presentation.
Additional fee	+ 13 600 EURO For each additional strength or pharmaceutical form including one presentation, submitted at the same time as the initial application for authorisation.
	+ 6 900 EURO For each additional presentation of the same strength and pharmaceutical form, submitted at the same time as the initial application for authorisation.

- Number of target species applied for does not impact on the fee.

##### 2.1.2.2. Full fee – Immunologicals

Basic fee	68 400 EURO For a single strength associated with one pharmaceutical form and one presentation.
Additional fee (I)	+ 6 900 EURO For each additional strength or pharmaceutical form including one presentation submitted at the same time as the initial application for authorisation.
	+ 6 900 EURO For each additional presentation of the same strength and pharmaceutical form, submitted at the same time as the initial application for authorisation.

Additional fee (II)	<p>+ 6 900 EURO</p> <p>For each multi-strain additional presentation of the same application submitted at the time of the initial application as described in the guideline on data requirements for multi-strain applications for inactivated vaccines against avian influenza, Bluetongue and Foot-and-Mouth Disease<sup>4</sup> up to a maximum total fee of 137 300 EURO.</p> <p>In this context, each combination of strain identified in the application represents a presentation.</p>
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- The number of target species applied for does not impact on the fee.

### 2.1.2.3. Reduced fee

Basic fee	<p>68 400 EURO</p> <p>For applications for marketing authorisation pursuant to Article 13(1), Article 13(3) and Article 13c of Directive 2001/82/EC. This fee is for a single strength associated with one pharmaceutical form and one presentation.</p>
Additional fee	<p>+ 13 600 EURO</p> <p>For each additional strength or pharmaceutical form <u>including one presentation</u> submitted at the same time as the initial application for authorisation.</p>
	<p>+ 6 900 EURO</p> <p>For each additional presentation of the same strength and pharmaceutical form, submitted at the same time as the initial application for authorisation.</p>
Basic fee	<p>116 000 EURO</p> <p>For applications for a marketing authorisation pursuant to Article 13(4) of Directive 2001/82/EC. This fee is for a single strength associated with one pharmaceutical form and one presentation.</p>
Additional fee	<p>+ 13 600 EURO</p> <p>For each additional strength or pharmaceutical form <u>including one presentation</u> submitted at the same time as the initial application for authorisation.</p>
	<p>+ 6 900 EURO</p> <p>For each additional presentation of the same strength and pharmaceutical form, submitted at the same time as the initial application for authorisation.</p>

- The number of target species applied for does not impact on the fee.

<sup>4</sup> Commission Directive 2009/9/EC amending Directive 2001/82 EC of the European Parliament and of the Council on the Community code relating to medicinal products for veterinary use introduces under Title IV B the concept of multi-strain veterinary immunological products for vaccines against avian influenza (AI), Bluetongue (BT) and Foot-and-mouth disease (FMD)

#### 2.1.2.4. Reduced fee – Immunologicals

Basic fee	34 300 EURO For a single strength associated with one pharmaceutical form and one presentation.
Additional fee	+ 6 900 EURO For each additional strength or pharmaceutical form and one presentation submitted at the same time as the initial application.

- The number of target species applied for does not impact on the fee.

**Note:** Refer to section [A.2.1.2.](#) in the annex for examples of the determination of fees for applications for marketing authorisation.

### 2.1.3. Extension of a marketing authorisation

#### 2.1.3.1. Extension of a marketing authorisation under Annex I to Commission Regulation (EC) No 1234/2008

Basic fee (Level I)	34 300 EURO For each extension of a marketing authorisation within the meaning of Annex I to Commission Regulation (EC) No 1234/2008 concerning the examination of variations to the terms of a marketing authorisation for veterinary medicinal products. This fee is for a single strength/potency associated with one pharmaceutical form and one presentation. This fee shall also cover one or more target species associated with that pharmaceutical form.
Additional fee	+ 8 600 EURO For each additional strength/potency of the same pharmaceutical form of the same extension including one presentation submitted at the time of the extension application. This fee shall also cover one or more target species associated with that pharmaceutical form.
Additional fee	+ 6 900 EURO For each additional presentation of the same strength/potency of the same pharmaceutical form of the same extension submitted at the same time of the extension application.

Basic fee (Level II)	30 900 EURO For all quality extensions (i.e. requiring chemical, pharmaceutical and biological documentation) for which no clinical or non-clinical data are submitted or no cross-references to previously submitted clinical or non-clinical data are made by the MAH. This fee is for a single strength/potency associated with one pharmaceutical form and one presentation.
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Additional fee	+ 8 600 EURO For each additional strength/potency of the same pharmaceutical form of the same extension including one presentation submitted at the time of the extension application. This fee shall also cover one or more target species associated with that pharmaceutical form.
Additional fee	+ 6 900 EURO For each additional presentation of the same strength/potency of the same pharmaceutical form of the same extension submitted at the same time of the extension application.

Basic fee (Level III)	8 600 EURO For all quality extensions (i.e. requiring chemical, pharmaceutical and biological documentation) of an immunological veterinary medicinal product for which no clinical or non-clinical data are submitted or no cross-references to previously submitted clinical or non-clinical data are made by the MAH. This fee is for a single strength/potency associated with one pharmaceutical form and one presentation.
Additional fee	+ 8 600 EURO For each additional strength/potency of the same pharmaceutical form of the same extension including one presentation submitted at the time of the extension application. This fee shall also cover one or more target species associated with that pharmaceutical form.
Additional fee	+ 6 900 EURO For each additional presentation of the same strength/potency of the same pharmaceutical form of the same extension submitted at the same time of the extension application.

Note: Refer to section [A.2.1.3.](#) in the annex for examples of the determination of fees for extensions of marketing authorisation.

## 2.1.4. Variation to a marketing authorisation

The following fees apply to variations that are notified or applied for individually in a notification or application.

### 2.1.4.1. Type-IA variations

Basic fee	3 000 EURO For a minor variation to a marketing authorisation, as defined in Article 2(2) of Commission Regulation (EC) No 1234/2008.
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### 2.1.4.2. Type-IB variations

Basic Fee	6 900 EURO For a minor variation to a marketing authorisation, as defined in Article 2(5) of Commission Regulation (EC) No 1234/2008.
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- In the event of the same variation being introduced, the fee will cover all authorised strengths, pharmaceutical forms and presentations of a given medicinal product (= per main authorisation number).

### 2.1.4.3. Type-II variations

Basic fee (Level I)	41 100 EURO For a major variation to a marketing authorisation, as defined in Article 2(3) of Commission Regulation (EC) No 1234/2008.
Basic fee (Level II)	30 900 EURO For a quality variation (i.e. amendments to the chemical, pharmaceutical and biological documentation) for which no clinical or non-clinical data are submitted or no cross-references to previously submitted clinical or non-clinical data are made by the MAH. (Note: bioequivalence data qualifies as clinical data.)
Basic fee (Level III)	10 300 EURO For each of the third and subsequent type II variations that are grouped in a single application made under the terms of Article 7(2) of Commission Regulation (EC) No 1234/2008.  For each of the third and subsequent type II variations to the centralised marketing authorisation or to one of the centralised marketing authorisations, where there is more than one centralised marketing authorisation, in the case of a worksharing application made under the terms of Article 20 of Commission Regulation (EC) No 1234/2008,
Basic fee (Level IV)	6 900 EURO For each variation to a marketing authorisation for an immunological veterinary medicinal product.



## **2.1.5. Grouping and worksharing procedures for variations**

### ***2.1.5.1. Grouping of extensions and/or variations notified or submitted under the terms of Article 7(2) of Commission Regulation (EC) No 1234/2008***

- The applicable fee specified in sections 2.1.3 and 2.1.4 shall be payable for each individual extension and/or variation to a marketing authorisation that is grouped in a single notification or a single application.
- The applicable level I and level II basic fees specified in sub-section 2.1.4.3 above are payable for the first and second type II variation respectively when both levels of fees are applicable to variations in the same grouping.
- Consequential variations in a grouping shall be similarly charged the applicable fees as specified above.
- In the case of grouping of the same Type IA variations to the terms of several marketing authorisations owned by the same holder (as set out in Article 7(2)(a) of Commission Regulation (EC) No 1234/2008), the applicable fee shall be payable for each individual Type IA variation and for each marketing authorisation in the grouping.
- The same marketing authorisation holder also means several marketing authorisation holders that are linked through a parent company. The fee for the grouping shall be payable by the marketing authorisation holder applying for the grouping procedure.
- Where any extensions/variations included in a grouping are found not to be valid and the remainder are validated positively, the applicable fees as specified above shall be payable for each of the positively validated extensions/variations.

### ***2.1.5.2. Worksharing procedure under the terms of Article 20 of Commission Regulation (EC) No 1234/2008***

- The applicable fee specified in section 2.1.4 is payable for each individual variation to one of the centralised marketing authorisations owned by the same holder, where more than one centralised marketing authorisation is included in the worksharing application, or to the single centralised marketing authorisation included in the worksharing application, as applicable.
- The applicable level I and level II basic fees specified in 2.1.4.3 above are payable for the first and second type II variations respectively, when both levels of fees are applicable to variations in the same worksharing procedure.
- The administrative fee shown in the table below is additionally payable for each individual variation to each of the other centralised marketing authorisation(s) owned by the same holder included in the same worksharing application, if applicable.
- No fee is payable to the Agency for any national marketing authorisations included in the same worksharing application.
- The same marketing authorisation holder also means several marketing authorisation holders that are linked through a parent company. The fee for the worksharing procedure shall be payable by the marketing authorisation holder applying for the procedure.

Variation type	Worksharing fees for one centralised marketing authorisation	Worksharing administrative fees
Type II (Level I)	41 100 EURO	3 400 EURO
Type II (Level II)	30 900 EURO	
Type II (Level III)	10 300 EURO	
Type IB	6 900 EURO	1 140 EURO
Type IA	3 000 EURO	570 EURO

- Where any variations included in a worksharing procedure are found not to be valid and the remainder are validated positively, the applicable fees as specified above shall be payable for each of the positively validated variations.

### 2.1.6. Renewal of a marketing authorisation

Basic fee	6 900 EURO For each strength associated with a pharmaceutical form.
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**Note:** Refer to section [A.2.1.6.](#) in the annex for examples of the determination of fees for renewals of marketing authorisation.

### 2.1.7. Inspection

Basic fee (Level I)	20 600 EURO For inspections outside the European Union, travel expenses shall be charged extra on the basis of actual cost.
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**Note:** Refer to section [A.1.1.7.](#) in the annex for examples of the determination of fees for good manufacturing practice (GMP) inspections.

### 2.1.8. Transfer of a marketing authorisation

Basic fee	6 900 EURO For all authorised presentations of the medicinal product.
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### 2.1.9. Maintenance of a marketing authorisation – Annual fee

Basic fee (Level I)	32 800 EURO For each marketing authorisation of a medicinal product. This fee covers all authorised presentations of the medicinal product.
Basic fee (Level II)	16 300 EURO For each marketing authorisation of a biosimilar medicinal product (Article 13(4) of Directive 2001/82/EC). This fee covers all authorised presentations of the medicinal product.

Basic fee (Level III)	8 200 EURO  For each marketing authorisation of a generic, hybrid or informed consent medicinal product (Articles 13(1), 13(3) and 13c of Directive 2001/82/EC). This fee covers all authorised presentations of the medicinal product.
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### 2.1.10. Referral

Basic fee	41 100 EURO  For the procedures laid down in Article 34(1) and 35 of Directive 2001/83/EC initiated by the applicant of a marketing authorisation or the holder of an existing marketing authorisation.
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**Note:** Where more than one applicant of marketing authorisations or holder of existing marketing authorisations are concerned by the procedures referred to in the first subparagraph, the applicants or holders may be grouped for the purpose of the payment of one single referral fee. If, however, the same procedure concerns more than 10 different applicants or holders, the fee will be charged by the application of the above mentioned referral fee.

### 2.2. Maximum residue limits (MRLs)

Basic fee	68 400 EURO  For an application to set an initial MRL for a given substance intended to be used in veterinary medicinal products or in biocidal products used in animal husbandry.
	20 600 EURO  For each application to modify or extend an existing MRL, as included in Table 1 of the Annex to Regulation (EEC) No 470/2009.

- MRL fees shall be deducted from the fee payable for an application for marketing authorisation or an application to extend a marketing authorisation for the medicinal product containing the substance for which an MRL has been set where such applications are submitted by the same applicant. However, this deduction may in total be no more than one half of the fee to which it applies.

## 2.3. Scientific advice

### 2.3.1. Definitions

The following definitions shall apply for the determination of fees for scientific advice requests.

*Quality development:* Chemical, pharmaceutical and biological testing.

*Safety development:* Toxicological and pharmacological tests.

*Clinical development:* Studies in animal patients, including clinical pharmacological trials designed to determine the efficacy and safety of the product.

*Initial request:* First request for scientific advice or protocol assistance introduced in relation to the submission of an application for marketing authorisation or a variation, whatever the authorisation phase (pre- or post-authorisation).

*Follow-up to initial request:* Any subsequent request falling within the same therapeutic indication and initial area(s) as the initial request, (area meaning quality, preclinical and/or clinical development including pharmacovigilance/risk management aspects).

### 2.3.2. Initial request for scientific advice

Basic fee (Level I)	41 100 EURO  This fee corresponds to an <b>initial</b> request for scientific advice (SA) on: <ul style="list-style-type: none"><li>• quality and safety and clinical development, or</li><li>• quality and clinical development, or</li><li>• safety and clinical development.</li></ul>
Basic fee (Level II)	20 600 EURO  For <b>initial</b> requests on: <ul style="list-style-type: none"><li>• clinical development, or</li><li>• quality and safety development, or</li><li>• quality and bioequivalence studies for generic medicinal products.</li></ul>
Basic fee (Level III)	13 600 EURO  For <b>initial</b> requests on: <ul style="list-style-type: none"><li>• quality development, or</li><li>• safety development, or</li><li>• bioequivalence studies for generic medicinal products, or</li><li>• new MRL.</li></ul>

### 2.3.3. Follow-up request for scientific advice

Basic fee (Level I)	20 600 EURO For <b>follow-up</b> to the initial request on: <ul style="list-style-type: none"><li>• quality and safety and clinical development, or</li><li>• quality and clinical development, or</li><li>• safety and clinical development.</li></ul>
Basic fee (Level II)	13 600 EURO For <b>follow-up</b> to the initial request on: <ul style="list-style-type: none"><li>• clinical development, or</li><li>• quality and safety development, or</li><li>• quality and bioequivalence studies for generic medicinal products.</li></ul>
Basic fee (Level III)	10 300 EURO For <b>follow-up</b> to the initial request on: <ul style="list-style-type: none"><li>• quality development, or</li><li>• safety development</li><li>• bioequivalence studies for generic medicinal products, or</li><li>• new MRL.</li></ul>

### 2.3.4. Scientific advice in relation to products classified by the CVMP

Basic fee	10 300 EURO For assessing compliance of a proposed data package with relevant guidelines on data requirements for veterinary medicinal products intended for minor uses or minor species.
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### 2.4. Fee for scientific services

A scientific service fee shall apply where an application is made for any scientific advice or opinion by a scientific committee, which is not covered by Articles 3 to 7 or by Article 8(1).

Basic fee (Level I)	137 300 EURO When it concerns veterinary medicinal products.
Basic fee (Level II)	34 300 EURO E.g. vaccine antigen master file.
Basic fee (Level III)	6 900 EURO Variations to a vaccine antigen master file.

### 3. Administrative fees

#### 3.1. Negative validation

Basic fee	2 980 EURO For an application that has been found not to be valid, an administrative fee for the validation of the application shall be charged. This fee is for all negative validations except for grouping or worksharing applications.
Basic fee	2 980 EURO For a grouping or worksharing application where all extensions/variations in the application have been found not to be valid. If individual extensions/variations in an application are found not to be valid but the remainder are validated positively, no administrative fee shall be charged for the invalid extensions/variations.

#### 3.2. Certificate of a medicinal product

##### 3.2.1. Definitions

*A set of certificates:* a maximum of six identical original certificates for a medicinal product with a distinct marketing authorisation number, addressed to the same importing country, issued in the same official language of the European Union and having identical annexes.

*Standard procedure:* procedure for issuing certificates of medicinal product within 10 working days.

*Urgent procedure:* procedure for issuing certificates of medicinal product within 2 working days.

##### 3.2.2. Standard procedure

Basic fee (Level I)	280 EURO For each request for certificates, including one set of certificates.
Additional fee	140 EURO For each additional set of certificates included in the same request.

##### 3.2.3. Urgent procedure

Basic fee (Level II)	840 EURO For each request for certificates, including one set of certificates.
Additional fee	420 EURO For each additional set of certificates included in the same request.

### 3.2.4. Withdrawal of request for certificates

Basic fee	280 EURO When a request for certificates is withdrawn by the requester following confirmation by the Agency of the start of the procedure.
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**Note:** Refer to section [A.3.2.](#) in the annex for examples of the determination of fees for certificates of medicinal product

### 3.3. Notification of parallel distribution

Initial notification of parallel distribution	2 980 EURO For each EU presentation of a medicinal product for one Member State of destination having one or more official languages or for several Member States of destination having the same official language. This fee covers any subsequent safety update notification relating to the initial notification triggered by a safety update adopted by the Committee for Medicinal Products for Human Use or the Committee for Medicinal Products for Veterinary Use, which is identified and communicated by the Agency to the parallel distributor.
Annual update notification, manual check	570 EURO For all changes made in one year from the anniversary of the initial notification to all the EU presentations belonging to the same pharmaceutical form of the same medicinal product for one Member State of destination having one or more official languages, or for several Member States of destination having the same official language. This fee applies when a report generated by text comparison software has not been provided or has not met the specific conditions laid down by the Agency for the acceptance of the report.
Annual update notification, automated check	280 EURO For all changes made in one year from the anniversary of the initial notification to all the EU presentations belonging to the same pharmaceutical form of the same medicinal product for one Member State of destination having one or more official languages, or for several Member States of destination having the same official language. This fee applies when a report generated by text comparison software has been provided and has met the specific conditions laid down by the Agency for the acceptance of the report.

<p>Notification of a change, <b>manual</b> check</p>	<p>570 EURO</p> <p>For each notification of a change that is not submitted as part of the annual update notification and is not a safety update.</p> <p>This fee applies for each EU presentation of a medicinal product for one Member State of destination having one or more official languages, or for several Member States of destination having the same official language.</p> <p>This fee applies when a report generated by text comparison software has not been provided or has not met the specific conditions laid down by the Agency for the acceptance of the report.</p>
<p>Notification of a change, <b>automated</b> check</p>	<p>280 EURO</p> <p>For each notification of a change that is not submitted as part of the annual update notification and is not a safety update.</p> <p>This fee applies for each EU presentation of a medicinal product for one Member State of destination having one or more official languages, or for several Member States of destination having the same official language.</p> <p>This fee applies when a report generated by text comparison software has been provided and has met the specific conditions laid down by the Agency for the acceptance of the report.</p>

### ***3.4. Worksharing procedures for variations to marketing authorisations***

Refer to section 1.1.5 and 2.1.5 for details of administrative charges applicable to worksharing procedures under the terms of Article 20 of Commission Regulation (EC) No 1234/2008.



## 4. Fee determination and payment

### 4.1. Legal requirements

Article 10 of Council Regulation (EC) No 297/95 stipulates the dates when fees are due and when they are payable.

#### Fee due date

The fee due date is the date when the Agency determines the total fee amount that is due for an application or service. The applicant's or marketing authorisation holder's obligation to pay that fee starts on that date.

The fee due date corresponds to:

- The date of administrative validation of an application; or
- The date of the start of a procedure (when there is no administrative validation), e.g. for type IA variation fees, transfer fees, Article 30(1) or 31 referrals; or
- The date of the anniversary of the notification of the marketing authorisation decision, for annual fees; or
- The date of the start of an inspection, for inspection fees.

#### Payment of fees

The Agency's terms and conditions for payment of fees stipulate that an invoice is payable 30 days from the invoice date. That represents the deadline for the applicant to settle the payment.

In summary, the procedure shown in Figure 1 applies.

**Figure 1.** Procedure for fee determination and payment



#### Deferred fees

When the applicant is a small or medium-sized enterprise (SME) payment of the fee for a marketing authorisation (MA) application and the fee for inspections undertaken as part of the assessment of a MA application is deferred until the final decision on the MA is issued or the application is withdrawn. The invoice issued at the time the fee is due informs the applicant that the fee is deferred. A reminder is issued by the Agency as soon as there is a decision or withdrawal; the fee is payable within 45 days of the date of the notification of the final decision on the MA or of the withdrawal of the application. The procedure is summarised in Figure 2.

**Figure 2.** Procedure for fee determination and deferral of payment by SME



The payment of the fee for an application for a MA of a medicinal product to be used in a human pandemic situation, so-called core dossier submission, is deferred until the pandemic situation is duly recognised, either by the World Health Organisation or the European Union, or up to 5 years from the due date, whichever comes first. Nevertheless the deferral can come to an end earlier on withdrawal prior to an opinion on the MA application, or if a negative Commission Decision is issued. A reminder is issued by the Agency as soon as the deferral comes to an end and the fee is payable immediately.

#### **4.2. Criteria for fee determination**

The criteria that are taken into account by the Agency when determining the fee that is due are the ones that apply by the fee due date. These include:

- The applicant's status (e.g. valid SME status assigned by the Agency);
- The product type (e.g. advanced therapy medicinal product, designated orphan product, veterinary product for minor uses and minor species (MUMS)/limited markets);
- The product status (e.g. in the first year following grant of a MA); and
- The procedure type.

Any change to the applicable criteria after the fee due date is not taken into account and will not influence the total fee due that has been determined.

#### **4.3. Established fee exemptions**

An applicant qualifies for fee exemptions and incentives specified in Chapter 5 provided that the required criteria apply on the fee due date. Any criterion that is no longer valid on the fee due date, e.g. expired SME status, cannot be taken into consideration by the Agency.

Applicants should ensure that they meet all the required criteria at the time of submission of an application or a request for a service.

#### **4.4. Request for ad hoc fee reduction**

Applicants may request the Executive Director to grant an ad hoc fee reduction under the provisions of the first paragraph of Article 9 of Council Regulation (EC) No 297/95.

The applicant is required to provide sufficient justification to prove that the request is made (1) in exceptional circumstances and (2) for imperative reasons of public or animal health. The request will be considered by the Executive Director who will consult the competent scientific committee and will decide each request on a case by case basis.

In view of the administrative procedure that has to be followed and the mandatory scientific consultation, applicants are required to make their request in a letter to the Executive Director at least two months before the date of submission of the relevant application. The applicant should cite Article 9, paragraph 1 of Council Regulation (EC) No 297/95 and provide details of the product, procedure type and applicable fee, and the reason(s) for the request that justify exceptional circumstances and imperative reasons of public or animal health.

Applicants are advised that late requests may not be processed in time by the fee due date and may not be taken into consideration when determining the fee. Requests received after the fee due date will not be considered.

Further information on fees and fee exemptions is available on the Agency's [website](#).

## 5. Fee exemptions

Where an applicant could, in respect of the same fee, benefit from more than one category of fee reduction or incentive (e.g. advanced therapy medicinal product, orphan medicinal product, small and medium sized enterprises) the provisions which are the most favourable to the applicant would apply. The applicant can contact the Agency prior to submission of the application for confirmation of the applicable fee.

### 5.1. Micro, small or medium-sized enterprise (SMEs)

#### 5.1.1. Definitions

Pursuant to Article 70(2) of Regulation (EC) No 726/2004 of 31 March 2004, applicants that meet the definition of a micro, small or medium-sized enterprise are eligible for fee incentives from the Agency. Companies developing medicinal products for human and/or veterinary use can benefit from fee reductions, exemptions and/or deferrals, as applicable, under Articles 5 to 9 of Regulation (EC) No 2049/2005. Applicants must be established in the EEA and fulfil the definition of an SME as set out in Commission Recommendation 2003/361/EC of 6 May 2003.

#### 5.1.2. Fee incentives

Scientific advice	90% reduction to the total applicable fee for non-orphan medicinal products
	100% reduction to the total applicable fee for designated orphan medicinal products (see section 5.2.3)
Inspection (pre-authorisation)	90% reduction to the total applicable fee
	Deferral of total applicable fee
Application for a marketing authorisation	Deferral of total applicable fee
	Conditional fee exemption
Scientific services	90% reduction to the total applicable fee for non-orphan medicinal products
	100% reduction to the total applicable fee for designated orphan medicinal products (see section 5.2.3)
Establishment of maximum residue limit for a veterinary medicinal product	90% reduction to the total applicable fee
Administrative services (excluding parallel distribution)	100% reduction to the total applicable fee
Inspection (post-authorisation)	90% reduction to the total applicable fee

## **5.2. Orphan medicinal products**

### **5.2.1. Definitions**

Total or partial exemptions from the payment of fees for applications for designated orphan medicinal products for human use shall be granted as laid down in a decision of the Executive Director on the use of the special contribution from the European Union, provided for by Article 7(2) of Regulation (EC) No 141/2000, reflecting the advice of the Committee for Orphan Medicinal Products.<sup>5</sup>

### **5.2.2. Fee incentives for applicants other than micro, small and medium sized enterprises**

Protocol assistance (non-paediatric-related*)	40% reduction to the total applicable fee
Protocol assistance (paediatric-related*)	100% reduction to the total applicable fee

\* Paediatric-related protocol assistance is restricted to the development of an orphan medicinal product for the paediatric population, where the advice requested does not include the adult population.

### **5.2.3. Fee incentives for micro, small and medium sized enterprises**

Protocol assistance	100% reduction to the total applicable fee
Scientific services**	100% reduction to the total applicable fee (see section 5.1.2)
Inspection (pre-authorisation)	100% reduction to the total applicable fee
Application for a marketing authorisation	100% reduction to the total applicable fee
Post-authorisation activities, including annual fees, during the first year after marketing authorisation	100% reduction to the total applicable fee
Inspection (post-authorisation)**	90% reduction to the total applicable fee (see section 5.1.2)

\*\* Fee reductions for scientific services and post-authorisation inspections are not funded by the special contribution from the European Union for designated orphan medicinal products but are provided for by Article 7 of Regulation (EC) No 2049/2005 on SMEs.

## **5.3. Multiple applications on usage patent grounds**

### **5.3.1. Definitions**

The full or partial exemptions from payment of fees described below are applicable for as long as the concerned marketing authorisation is affected by usage patent(s) pertaining to indication(s) and/or dosage form(s).

<sup>5</sup> Executive Decision of 6 December 2012 applicable from 1 January 2013 (EMA/663496/2012)

The following ranges and classification shall apply for fees for:

- generic medicinal product applications submitted under Article 10(1) of Directive 2001/83/EC;
- hybrid applications submitted under Article 10(3) of Directive 2001/83/EC;
- similar biological medicinal product applications submitted under Article 10(4) of Directive 2001/83/EC.

### 5.3.2. Fee incentives for an application for a marketing authorisation

#### 5.3.2.1. Reduced fee

Second and each subsequent multiple application submitted under Articles 10(1) and 10(3) of Directive 2001/83/EC	20 400 EURO
Second and each subsequent multiple application submitted under Article 10(4) of Directive 2001/83/EC	34 000 EURO
Additional strengths, pharmaceutical forms and presentations submitted at the same time as the aforementioned applications	100% reduction to the total applicable fee

### 5.3.3. Fee incentives for post-authorisation activities for a second and for each subsequent multiple application

#### 5.3.3.1. Reduced fee

Extension of a marketing authorisation	19 700 EURO
Additional strengths and presentations of the same pharmaceutical form submitted at the same time as the aforementioned application	100% reduction to the total applicable fee

Type-IA variation to a marketing authorisation. This fee shall only apply in the case of grouping of the same type-IA variations to the terms of multiple marketing authorisations on usage patent grounds owned by the same holder (as defined in Article 7(2)(a) of Commission Regulation (EC) No 1234/2008). The applicable fee shall be payable for each individual type-IA variation relating to the second and each of the subsequent multiple marketing authorisations in the grouping	570 EURO
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Worksharing procedure under the terms of Article 20 of Commission Regulation (EC) No 1234/2008. Administrative fees for variations to multiple centralised marketing authorisations on usage patent grounds	Refer to section 1.1.5.2
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Renewal of a marketing authorisation	2 620 EURO
Additional strengths associated with a pharmaceutical form submitted at the same time as the aforementioned application	100% reduction to the total applicable fee

Annual fee for a marketing authorisation granted under Articles 10(1) and 10 (3) of Directive 2001/83/EC	4 600 EURO
Annual fee for a marketing authorisation granted under Article 10(4) of Directive 2001/83/EC	9 400 EURO

## **5.4. Medicinal products for paediatric use**

### **5.4.1. Definitions**

A partial exemption from the payment of the fees laid down in the fee regulation is granted for paediatric use marketing authorisation applications (PUMAs) submitted under Article 30 of Regulation (EC) No 1901/2006 on medicinal products for paediatric use.

A total exemption from the payment of the fees laid down in the fee regulation is granted for scientific advice provided by the Agency to sponsors developing medicinal products for the paediatric population as laid down in Regulation (EC) No 1901/2006 on medicinal products for paediatric use.

### **5.4.2. Fee incentives for paediatric use marketing authorisations (PUMAs)**

Application for a paediatric use marketing authorisation	50% reduction to the total applicable fee
Inspection (pre-authorisation)	
During the first year after marketing authorisation for: <ul style="list-style-type: none"> <li>• extension of a marketing authorisation;</li> <li>• type-IA, type-IB and type-II variations;</li> <li>• annual fee;</li> <li>• inspection (post-authorisation).</li> </ul>	

### **5.4.3. Fee incentives for scientific advice**

Scientific advice on the development of a medicinal product for the paediatric population (when the advice requested does not include the adult population)	100% reduction to the total applicable fee
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## **5.5. Advanced therapy medicinal products**

### **5.5.1. Definitions**

Total or partial exemptions from the payment of fees for applications for advanced therapy medicinal products for human use are granted as laid down in Articles 16(2) of Regulation (EC) No 1394/2007.

Fee reductions for applications for marketing authorisation and for post-authorisation activities during the first year from granting of a marketing authorisation ceased to apply after 30 December 2011 in the case of advanced therapy medicinal products other than tissue-engineered products and ceased to apply after 30 December 2012 in the case of tissue-engineered products.

### 5.5.2. Fee incentives for applicants other than micro, small and medium-sized enterprises

Scientific advice	65% reduction to the total applicable fee
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### 5.5.3. Fee incentives for micro, small and medium-sized enterprises

Scientific advice	90% reduction to the total applicable fee (see section 5.1.2)
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## 5.6. Core dossier medicinal products to be used in a human pandemic situation

### 5.6.1. Definitions

A total exemption from the payment of the fees laid down in the fee regulation is granted for the regulatory activities specified below within the framework of the submission of a core dossier for a pandemic influenza vaccine and the follow-up submission of a pandemic variation, as described in the 'Guideline on Dossier Structure and Content for Pandemic Influenza Vaccine Marketing Authorisation Application' ([EMEA/CPMP/VEG/4717/03](#)).

The following total exemptions apply until the type II pandemic variation, submitted once the human pandemic situation is duly recognised, has been authorised by the European Union but, in any case, do not apply after the five-year period from the date of administrative validation of the marketing authorisation application for the core dossier has elapsed.

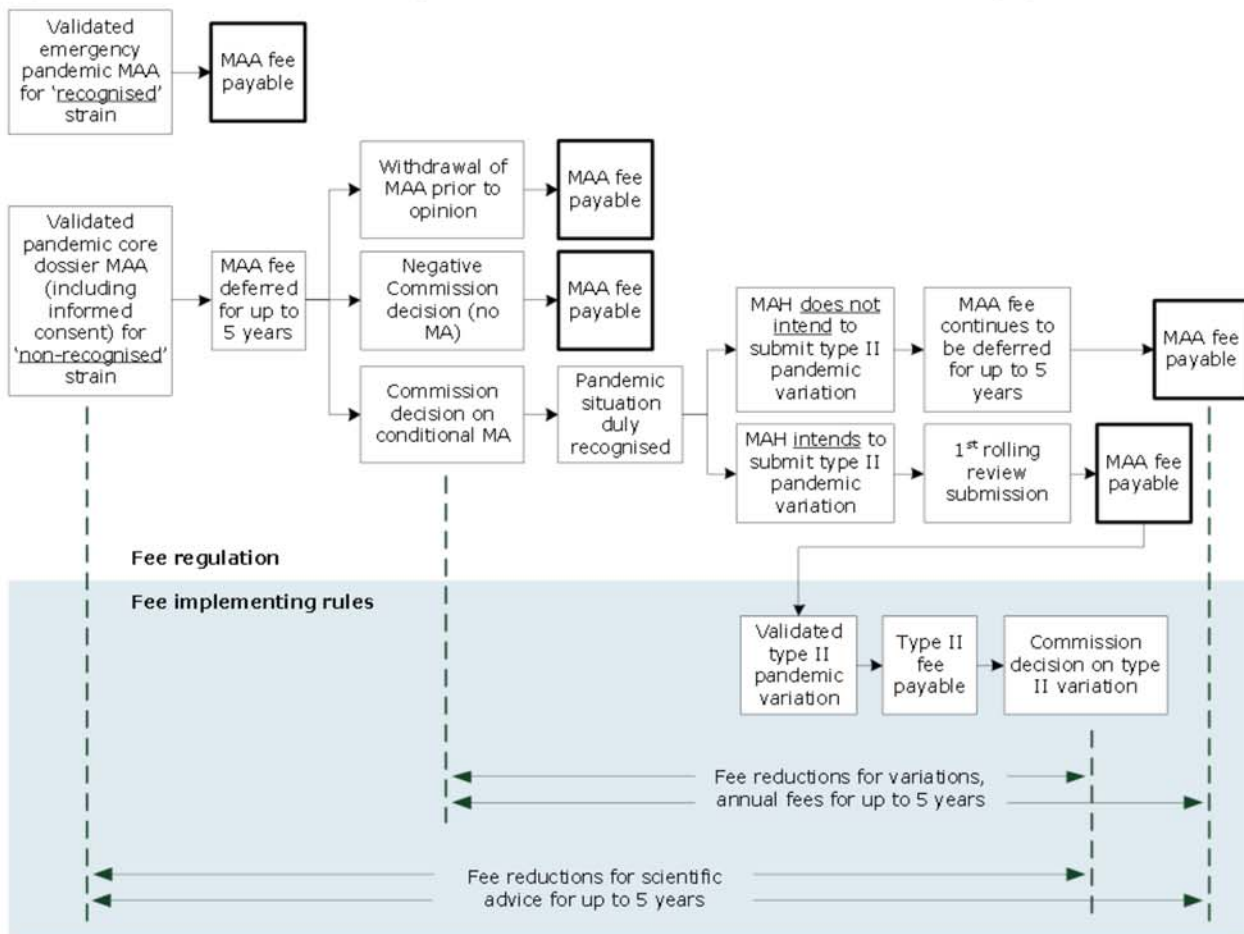
### 5.6.2. Fee incentives related to pandemic core dossier (including informed consent) for 'non-recognised' strain

Scientific advice	100% reduction to the total applicable fee
Post-authorisation activities including type-IA, type-IB, type-II variations (but excluding the type-II pandemic variation) and annual fee	100% reduction to the total applicable fee
Negative validation of a type-IB or type-II variation (but excluding the type-II pandemic variation)	100% reduction to the total applicable fee

- The fee for a validated pandemic core dossier marketing authorisation application (MAA), including informed consent application, for a pandemic strain that has not yet been "duly recognised, either by the World Health Organisation or by the Community" is deferred.
- If the core dossier MAA is withdrawn prior to opinion or receives a negative Commission Decision, the fee becomes payable and the deferral comes to an end.
- If the core dossier receives a positive Commission Decision, the fee shall continue to be deferred. The MAH also receives fee reductions on variations and annual fees.
- When the pandemic situation becomes duly recognised, the marketing authorisation holder (MAH) may choose not to submit a Type II pandemic variation. In this case the MAA fee continues to be deferred for up to 5 years from the MAA validation date. The MAH continues to receive fee reductions on variations and annual fees for up to 5 years from the MAA validation date.

- If the MAH chooses to submit a Type II pandemic variation, the initial MAA fee becomes payable on the first submission made as part of the rolling review. When the Type II pandemic variation is validated the applicable Type II fee becomes payable. The MAH continues to receive fee reductions on variations and annual fees until the date of the Type II Commission Decision.
- Fee reductions for scientific advice are applicable to requests in the context of a pandemic core dossier and at any time up to 5 years from the MAA validation date or the date of the Type II Commission Decision, whichever comes first.
- The fee for an emergency pandemic MAA submitted when the pandemic situation has been duly recognised becomes payable on validation.

**Figure 3. Schematic overview of pandemic vaccine fee reductions, deferrals and payments**





## **5.7. Medicinal products for minor uses and minor species (MUMS)/limited markets**

### **5.7.1. Definitions**

Exemptions and reductions are granted on fees relating to applications for products classified by the Committee for Medicinal Products for Veterinary Use (CVMP) as indicated for minor use and/or minor species or for which the market is confirmed by the Committee as 'limited' and for which no alternative products are authorised in the EU.

### **5.7.2. Fee incentives**

Scientific advice	100% exemption to the total applicable fee
Administrative fee for negative validation	100% exemption to the total applicable fee

Establishment or extension of an MRL for a substance with respect to an relevant indication	50% reduction to the total applicable fee
Application for a marketing authorisation	50% reduction to the total applicable fee
Extension of a marketing authorisation for a product indicated for a food-producing species to add a minor species	50% reduction to the total applicable fee
Type II variation to a marketing authorisation for a non-food-producing species to add a minor species.	50% reduction to the total applicable fee
Extrapolations of existing MRLs to relevant minor species for which no data are required and therefore no assessment is performed	100% reduction to the total applicable fee
Annual fee for a product authorised exclusively for indications classified by the CVMP	75% reduction to the total applicable fee

## Annex

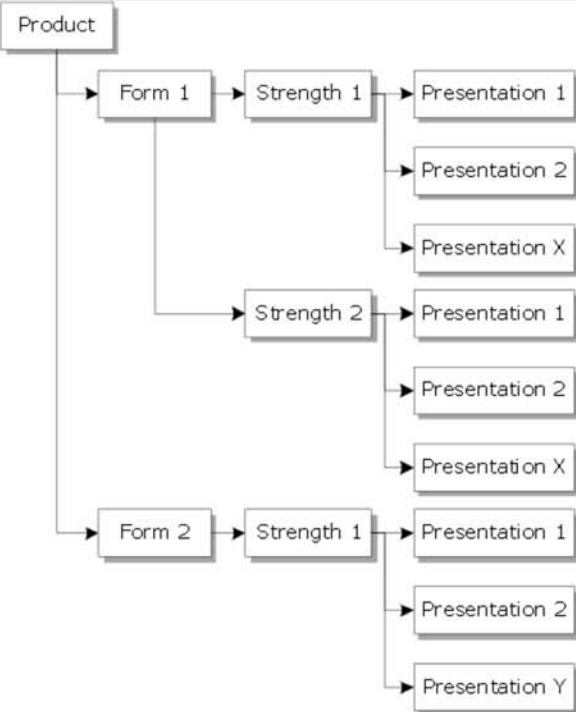
The additional information in this annex is listed using the same numbering as the corresponding sections in chapters 1 to 4. For example, section A.1.1.2. in this annex relates to section 1.1.2. in Chapter 1.

### A.1. Medicinal products for human use

#### A.1.1.2. Examples of the determination of fees for applications for marketing authorisation

It should be noted that the calculation of the total fee for a marketing authorisation application is driven by the pharmaceutical forms, the associated strengths and the associated presentations. The examples shown below do not represent an exhaustive list.

Scenario		Fee payable
1. Full dossier application for a medicinal product having one pharmaceutical form with one strength and X presentations	<pre>graph LR; Product --&gt; Form1[Form 1]; Form1 --&gt; Strength1[Strength 1]; Strength1 --&gt; Presentation1[Presentation 1]; Strength1 --&gt; Presentation2[Presentation 2]; Strength1 --&gt; PresentationX[Presentation X];</pre>	274 400 EURO (basic fee including one pharmaceutical form and one associated strength and one presentation)  + (X-1)*6 900 EURO (additional presentations associated with the single strength)

Scenario		Fee payable
<p>2. Full dossier application for a medicinal product having two pharmaceutical forms with two strengths and X presentations/ strength associated with the first form and one strength and Y presentations associated with the second form</p>	 <pre> graph LR     Product --&gt; Form1[Form 1]     Product --&gt; Form2[Form 2]     Form1 --&gt; Strength1_1[Strength 1]     Form1 --&gt; Strength2_1[Strength 2]     Form2 --&gt; Strength1_2[Strength 1]     Strength1_1 --&gt; P1_1[Presentation 1]     Strength1_1 --&gt; P2_1[Presentation 2]     Strength1_1 --&gt; PX_1[Presentation X]     Strength2_1 --&gt; P1_2[Presentation 1]     Strength2_1 --&gt; P2_2[Presentation 2]     Strength2_1 --&gt; PX_2[Presentation X]     Strength1_2 --&gt; P1_3[Presentation 1]     Strength1_2 --&gt; P2_3[Presentation 2]     Strength1_2 --&gt; PY_3[Presentation Y] </pre>	<p>274 400 EURO (basic fee including one pharmaceutical form and one associated strength and one presentation)</p> <p>+ (X-1)*6 900 EURO (additional presentations associated with the first form and strength)</p> <p>+ 27 500 EURO (second strength associated with the first form including one presentation)</p> <p>+ (X-1)*6 900 EURO (additional presentations associated with the first form and second strength)</p> <p>+ 27 500 EURO (second form including its associated strength and one presentation)</p> <p>+ (Y-1)*6 900 EURO (additional presentations associated with the second form and its strength)</p>

Scenario		Fee payable
<p>3. Full dossier application for an insulin product having two pharmaceutical forms with six strengths (consisting of two sets of one uncombined preparation and two combination preparations (having different proportions of insulin) with insulin amounts corresponding to A I.U. and B I.U.) and X presentations/strength associated with the first form; and two strengths (of uncombined preparations with insulin amounts corresponding to A I.U. and B I.U.) and Y presentations/strength associated with the second form</p>	<p>(u) = un-combined insulin preparation (c) = combination insulin preparation (refer to section 1.1.1 for definition of strength of insulin products)</p>	<p>274 400 EURO (basic fee including one pharmaceutical form and one associated strength and one presentation)</p> <p>+ (X-1)*6 900 EURO (additional presentations associated with the first form and strength)</p> <p>+ 5*27 500 EURO (for second to sixth strengths associated with the first form including one presentation for each strength)</p> <p>+ 5*(X-1)*6 900 EURO (additional presentations associated with the second to sixth strengths of the first form)</p> <p>+ 27 500 EURO (second form including one associated strength and one presentation)</p> <p>+ (Y-1)*6 900 EURO (additional presentations associated with the second form and first strength)</p> <p>+ 27 500 EURO (second strength associated with the second form including one presentation)</p> <p>+ (X-1)*6 900 EURO (additional presentations associated with the second form and second strength)</p>

### A.1.1.3. Examples of the determination of fees for extensions of marketing authorisation

It should be noted that the basic fee for an extension of a marketing authorisation is driven by the pharmaceutical form. The examples shown below do not represent an exhaustive list.

Scenario	Extension application	Fee payable
<p>1. New pharmaceutical form with two strengths and X presentations/strength, for authorised or new route of administration</p> <p>(with submitted/cross-referenced clinical data)</p>	<p>One pharmaceutical form, first strength and X presentations</p> <p>Second strength (of same new pharmaceutical form) and X presentations</p>	<p>82 400 EURO (basic fee for extension)</p> <p>+ (X-1)*6 900 EURO (additional presentation fees)</p> <p>+ 20 600 EURO (additional strength fee)</p> <p>+ (X-1)*6 900 EURO (additional presentation fees)</p>
<p>2. New route of administration for authorised pharmaceutical form with two authorised strengths and X presentations/strength</p> <p>(with submitted/cross-referenced clinical data)</p>	<p>Route of administration for authorised pharmaceutical form, first strength and X presentations</p> <p>Second strength (same new route of administration for same authorised pharmaceutical form) and X presentations</p>	<p>82 400 EURO (basic fee for extension)</p> <p>+ (X-1)*6 900 EURO (additional presentation fees)</p> <p>+ 20 600 EURO (additional strength fee)</p> <p>+ (X-1)*6 900 EURO (additional presentation fees)</p>
<p>3. Two new strengths of same authorised pharmaceutical form and X presentations/strength</p> <p>(without submitted/cross-referenced clinical data)</p>	<p>First new strength and X presentations</p> <p>Second new strength (of same authorised pharmaceutical form) and X presentations</p>	<p>61 800 EURO (basic fee for extension)</p> <p>+ (X-1)*6 900 EURO (additional presentation fees)</p> <p>+ 20 600 EURO (additional strength fee)</p> <p>+ (X-1)*6 900 EURO (additional presentation fees)</p>
<p>4. One new strength of each of two authorised pharmaceutical forms and X presentations/strength</p> <p>(without submitted/cross-referenced clinical data)</p> <p><i>THESE SHOULD BE SUBMITTED AS TWO EXTENSION APPLICATIONS</i></p>	<p>New strength (of first authorised pharmaceutical form) and X presentations</p> <p>New strength (of second authorised pharmaceutical form) and X presentations</p>	<p>61 800 EURO (basic fee for extension)</p> <p>+ (X-1)*6 900 EURO (additional presentation fees)</p> <p>61 800 EURO (basic fee for extension)</p> <p>+ (X-1)*6 900 EURO (additional presentation fees)</p>

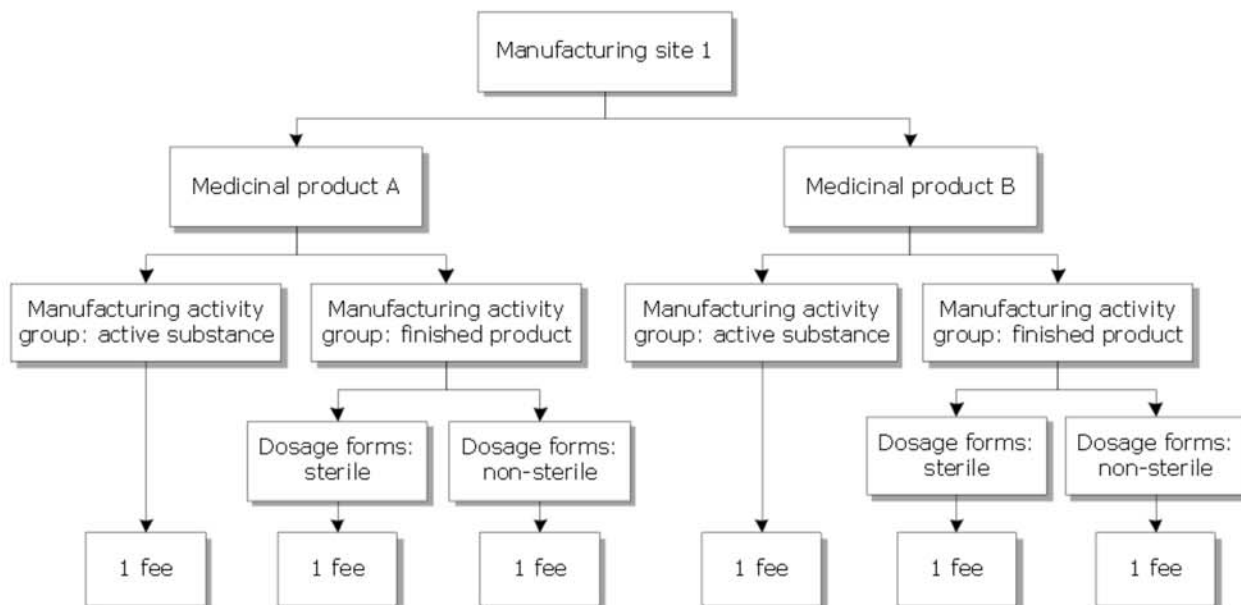
### A.1.1.6. Examples of the determination of fees for renewals of marketing authorisation

It should be noted that the number of renewal fees charged for a medicinal product depends on the number of strengths associated with each pharmaceutical form determined as shown in section A.1.1.2. The number of presentations is not taken into consideration for the calculation of the renewal fee. The examples shown below do not represent an exhaustive list.

Scenario	Strengths associated with a pharmaceutical form	Fee payable
1. Full dossier application for a medicinal product having one pharmaceutical form with one strength and X presentations	One strength associated with one pharmaceutical form	13 600 EURO (basic fee for renewal)
2. Full dossier application for a medicinal product having two pharmaceutical forms with two strengths and X presentations/ strength associated with the first form and one strength and Y presentations associated with the second form	Two strengths associated with first pharmaceutical form  One strength associated with second pharmaceutical form	2*13 600 EURO (basic fee for renewal)  + 13 600 EURO (basic fee for renewal)
3. Full dossier application for an insulin product having two pharmaceutical forms with six strengths (consisting of two sets of one un-combined preparation and two combination preparations (having different proportions of insulin) with insulin amounts corresponding to A I.U. and B I.U.) and X presentations/ strength associated with the first form; and two strengths (of un-combined preparations with insulin amounts corresponding to A I.U. and B I.U.) and Y presentations/strength associated with the second form	Six strengths associated with first pharmaceutical form  Two strengths associated with second pharmaceutical form	6*13 600 EURO (basic fee for renewal)  + 2*13 600 EURO (basic fee for renewal)

### A.1.1.7. Examples of the determination of fees for GMP inspections

In accordance with Annex IV to the "Rules for the implementation of Council Regulation (EC) No 297/95 on fees payable to the European Medicines Agency and other measures", applicants are liable for more than one inspection fee on the basis of the following flowchart.



The examples shown below do not represent an exhaustive list. They apply to good manufacturing practice (GMP) inspections in relation to medicines for human use and to medicines for veterinary use.

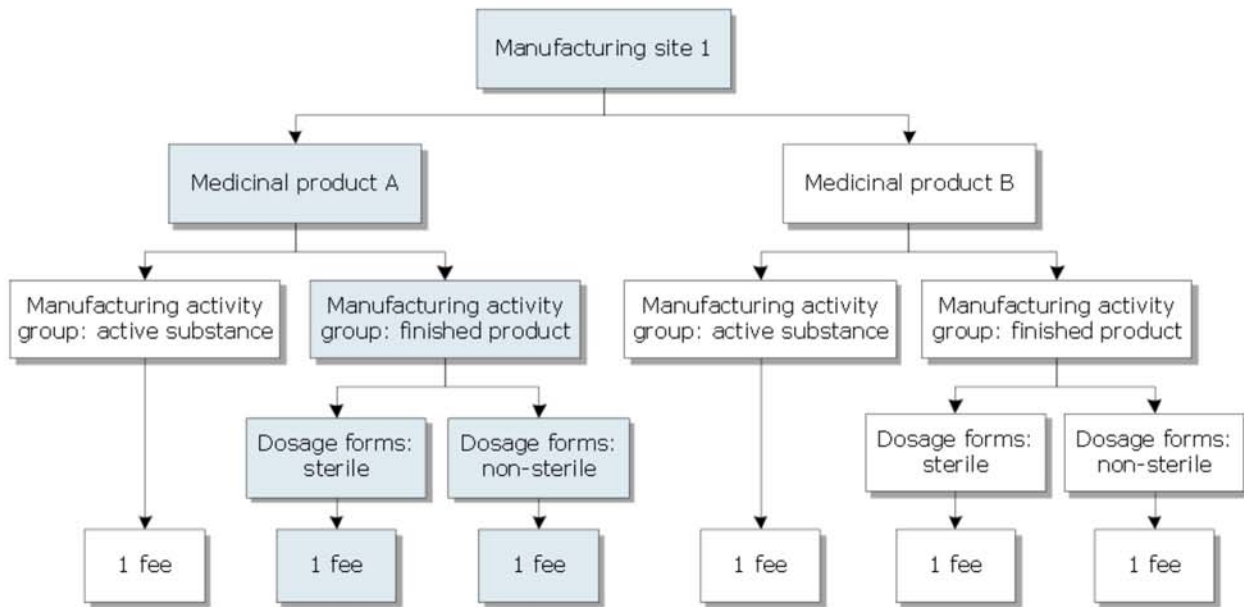
**Scenario 1:**

GMP inspection of manufacturing site 1 for one medicinal product A and involving two pharmaceutical forms: capsules (non-sterile) and solution for injection (sterile). The manufacturing activity for the two pharmaceutical forms is the same, i.e. manufacture of the finished product.

Fee payable: 2 basic fees (level I), i.e. 20 600 EURO + 20 600 EURO = 41 200 EURO

Rationale: there are two types of dosages forms (sterile and non-sterile) and each one attracts a basic fee (Level I).

The applicable fees to be paid are represented by the grey boxes in the flowchart below.





## Scenario 2:

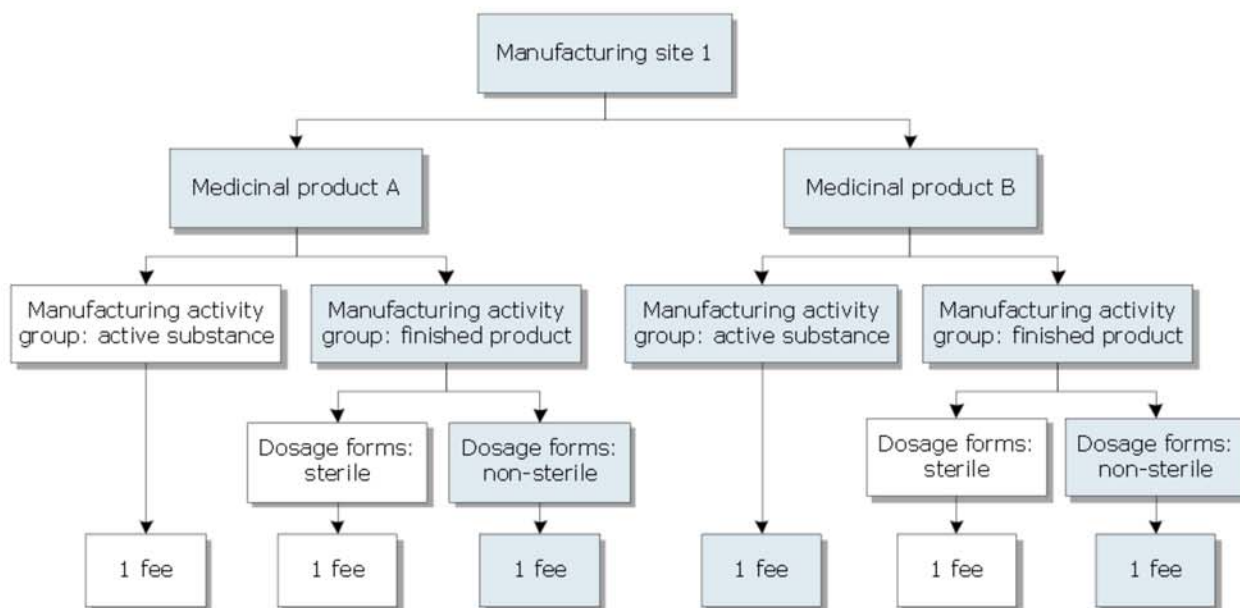
GMP inspection of manufacturing site 1 for two medicinal products (A and B). Product A involves only one pharmaceutical form (capsules) and one pharmaceutical activity (primary packaging). Product B also involves one pharmaceutical form (tablets) and four manufacturing activities (manufacture of the active substance, quality control of the active substance, manufacture of the finished product and primary packaging).

Fee payable: 3 basic fees (Level I), i.e. 20 600 EURO + 20 600 EURO + 20 600 EURO = 61 800 EURO

Rationale: Product A attracts only one fee because there is only one group of manufacturing activities (i.e. finished product) and one dosage form (non-sterile). Product B attracts two fees because there are manufacturing activities related to each group as follows:

- Group Active Substance: manufacture of the active substance and quality control of the active substance
- Group Finished Product: manufacture of the finished product and Primary Packaging

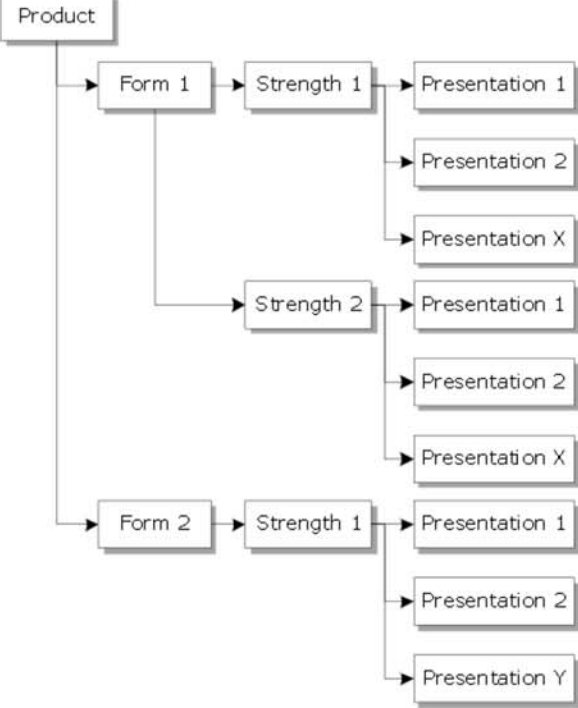
The applicable fees to be paid are represented by the grey boxes in the flowchart below.



## A.2. Medicinal products for veterinary use

### A.2.1.2. Examples of the determination of fees for applications for marketing authorisation

It should be noted that the calculation of the total fee for a marketing authorisation application is driven by the pharmaceutical forms, the associated strengths and the associated presentations. The number of target species is irrelevant. The examples shown below do not represent an exhaustive list.

Scenario	Diagram	Fee payable
<p>1. Full dossier application for a medicinal product having one pharmaceutical form with one strength and X presentations</p>	 <pre> graph LR     Product[Product] --&gt; Form1[Form 1]     Form1 --&gt; Strength1[Strength 1]     Strength1 --&gt; P1[Presentation 1]     Strength1 --&gt; P2[Presentation 2]     Strength1 --&gt; PX[Presentation X]         </pre>	<p>137 300 EURO (basic fee including one pharmaceutical form and one associated strength and one presentation)</p> <p>+ (X-1)*6 900 EURO (additional presentations associated with the single strength)</p>
<p>2. Full dossier application for a medicinal product having two pharmaceutical forms with two strengths and X presentations/ strength associated with the first form and one strength and Y presentations associated with the second form</p>	 <pre> graph LR     Product[Product] --&gt; Form1[Form 1]     Product --&gt; Form2[Form 2]     Form1 --&gt; S1_1[Strength 1]     Form1 --&gt; S2_1[Strength 2]     Form2 --&gt; S1_2[Strength 1]     S1_1 --&gt; P1_1[Presentation 1]     S1_1 --&gt; P2_1[Presentation 2]     S1_1 --&gt; PX_1[Presentation X]     S2_1 --&gt; P1_2[Presentation 1]     S2_1 --&gt; P2_2[Presentation 2]     S2_1 --&gt; PX_2[Presentation X]     S1_2 --&gt; P1_3[Presentation 1]     S1_2 --&gt; P2_3[Presentation 2]     S1_2 --&gt; PY_3[Presentation Y]         </pre>	<p>137 300 EURO (basic fee including one pharmaceutical form and one associated strength and one presentation)</p> <p>+ (X-1)*6 900 EURO (additional presentations associated with the first form and strength)</p> <p>+ 13 600 EURO (second strength associated with the first form including one presentation)</p> <p>+ (X-1)*6 900 EURO (additional presentations associated with the first form and second strength)</p> <p>+ 13 600 EURO (second form including its associated strength and one presentation)</p> <p>+ (Y-1)*6 900 EURO (additional presentations associated with the second form and its strength)</p>

### A.2.1.3. Examples of the determination of fees for extensions of marketing authorisation

It should be noted that the basic fee for an extension of a marketing authorisation is driven by the pharmaceutical form. The examples shown below do not represent an exhaustive list.

Scenario	Extension application	Fee payable
<p>1. New pharmaceutical form with two strengths and X presentations/strength, for authorised/new route of administration</p> <p>(with submitted/cross-referenced clinical data)</p>	<p>One pharmaceutical form, first strength and X presentations</p> <p>Second strength (of same new pharmaceutical form) and X presentations</p>	<p>34 300 EURO (basic fee for extension)</p> <p>+ (X-1)* 6 900 EURO (additional presentation fees)</p> <p>+ 8 600 EURO (additional strength fee)</p> <p>+ (X-1)* 6 900 EURO (additional presentation fees)</p>
<p>2. New route of administration for authorised pharmaceutical form with two authorised strengths and X presentations/strength</p> <p>(with submitted/cross-referenced clinical data)</p>	<p>Route of administration for authorised pharmaceutical form, first strength and X presentations</p> <p>Second strength (same new route of administration for same authorised pharmaceutical form) and X presentations</p>	<p>34 300 EURO (basic fee for extension)</p> <p>+ (X-1)*6 900 EURO (additional presentation fees)</p> <p>+ 8 600 EURO (additional strength fee)</p> <p>+ (X-1)*6 900 EURO (additional presentation fees)</p>
<p>3. Two new strengths of same authorised pharmaceutical form and X presentations/strength</p> <p>(without submitted/cross-referenced clinical data)</p>	<p>First new strength and X presentations</p> <p>Second new strength (of same authorised pharmaceutical form) and X presentations</p>	<p>30 900 EURO (basic fee for extension)</p> <p>+ (X-1)*6 900 EURO (additional presentation fees)</p> <p>+ 8 600 EURO (additional strength fee)</p> <p>+ (X-1)*6 900 EURO (additional presentation fees)</p>
<p>4. One new strength of each of two authorised pharmaceutical forms and X presentations/strength</p> <p>(without submitted/cross-referenced clinical data)</p> <p>THESE SHOULD BE SUBMITTED AS TWO EXTENSION</p>	<p>New strength (of first authorised pharmaceutical form) and X presentations</p> <p>New strength (of second authorised pharmaceutical form) and X presentations</p>	<p>30 900 EURO (basic fee for extension)</p> <p>+ (X-1)*6 900 EURO (additional presentation fees)</p> <p>30 900 EURO (basic fee for extension)</p> <p>+ (X-1)*6 900 EURO (additional presentation fees)</p>

Scenario	Extension application	Fee payable
APPLICATIONS	New pharmaceutical form for new target species, first strength and X presentations	34 300 EURO (basic fee for extension) + (X-1)*6 900 EURO (additional presentation fees)
	Second strength (of same new pharmaceutical form for same new target species) and X presentations	+ 8 600 EURO (additional strength fee) + (X-1)*6 900 EURO (additional presentation fees)
	New pharmaceutical form for a new target species, one strength and X presentations	34 300 EURO (basic fee for extension) + (X-1)*6 900 EURO (additional presentation fees)
	Second new target species of authorised pharmaceutical form, new strength and X presentations	34 300 EURO + (X-1)*6 900 EURO (additional presentation fees) +8 600 EURO (additional strength fee)
	Second strength (for same new target species of authorised pharmaceutical form) and X presentations	+ (X-1)*6 900 EURO (additional presentation fees)

### A.2.1.6. Examples of the determination of fees for renewals of marketing authorisation

It should be noted that the number of renewal fees charged for a medicinal product depends on the number of strengths associated with each pharmaceutical form determined as shown in section A.2.1.2. The number of presentations is not taken into consideration for the calculation of the renewal fee. The examples shown below do not represent an exhaustive list.

Scenario	Strengths associated with a pharmaceutical form	Fee payable
1. Full dossier application for a medicinal product having one pharmaceutical form with one strength and X number of presentations	One strength associated with one pharmaceutical form	6 900 EURO (basic fee for renewal)
2. Full dossier application for a medicinal product having two pharmaceutical forms with two strengths and X number of presentations/strength associated with the first form and one strength and Y number of presentations associated with the second form	Two strengths associated with first pharmaceutical form	2*6 900 EURO (basic fee for renewal)
	One strength associated with second pharmaceutical form	+ 6 900 EURO (basic fee for renewal)

### A.3. Administrative fees

#### A.3.2. Examples of the determination of fees for certificates of medicinal product

In accordance with Annex III of the "Rules for the implementation of Council Regulation (EC) No 297/95 on fees payable to the European Medicines Agency and other measures", requesters are liable for more than one fee depending on the number of medicinal products, importing countries, languages and annexes.

The examples shown below do not represent an exhaustive list.

##### Scenario 1:

One request for certificates for one medicinal product, as follows:

- Addressed to country 1: 5 certificates with annex 1
- Addressed to country 2: 10 certificates with annex 1

Fee payable using the standard procedure:

1 basic fee for the first set of 5 certificates for country 1 = 280 EURO

1 additional fee for the second set of maximum 6 out of 10 certificates for country 2 = 140 EURO

1 additional fee for the third set of 4 out of 10 certificates for country 2 = 140 EURO

Total fee: 280 EURO + 140 EURO + 140 EURO = 560 EURO

Fee payable using the urgent procedure:

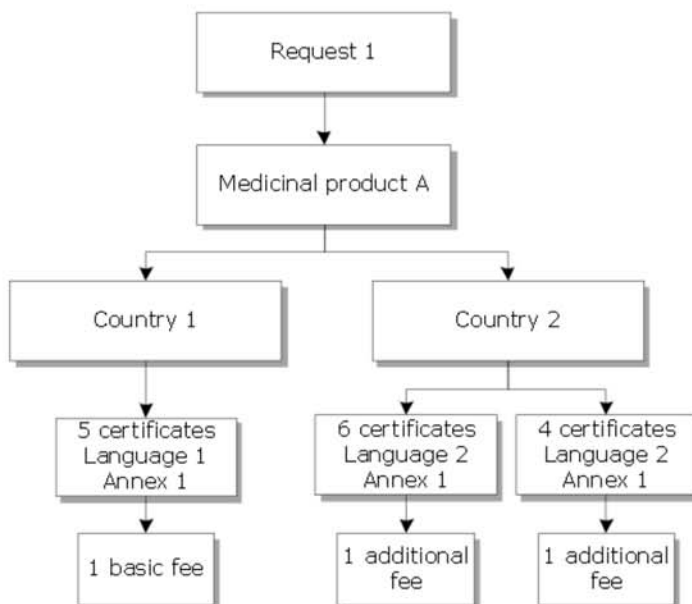
1 basic fee for the first set of 5 certificates for country 1 = 840 EURO

1 additional fee for the second set of maximum 6 out of 10 certificates for country 2 = 420 EURO

1 additional fee for the third set of 4 out of 10 certificates for country 2 = 420 EURO

Total fee: 840 EURO + 420 EURO + 420 EURO = 1 680 EURO

The applicable fees to be paid using the standard procedure are represented in the flowchart below.



## Scenario 2:

One request for certificates for two medicinal products, as follows:

- Medicinal product A:
  - Addressed to country 1: 5 certificates with annex 1
  - Addressed to country 2: 6 certificates with Annex 1, 9 certificates with annexes 1 and 2
- Medicinal product B:
  - Addressed to country 1: 15 certificates with annex 1
  - Addressed to country 2: 6 certificates with annex 1

Fee payable using the standard procedure:

1 basic fee for the first set of 5 certificates for medicinal product A for country 1 = 280 EURO

1 additional fee for the second set of 6 certificates for medicinal product A for country 2 = 140 EURO

1 additional fee for the third set of maximum 6 out of 9 certificates for medicinal product A for country 2 = 140 EURO

1 additional fee for the fourth set of 3 out of 9 certificates for medicinal product A for country 2 = 140 EURO

1 additional fee for the fifth set of maximum 6 out of 15 certificates for medicinal product B for country 1 = 140 EURO

1 additional fee for the sixth set of maximum 6 out of 15 certificates for medicinal product B for country 1 = 140 EURO

1 additional fee for the seventh set of 3 out of 15 certificates for medicinal product B for country 1 = 140 EURO

1 additional fee for the eighth set of 6 certificates for medicinal product B for country 2 = 140 EURO

Total fee: 280 EURO + (140 EURO \* 7) = 1 260 EURO

Fee payable using the urgent procedure: applying the same rationale as for the standard procedure, one basic fee and seven additional fees are payable.

Total fee = 840 EURO (420 EURO\* 7) = 3 780 EURO

The applicable fees to be paid using the standard procedure are represented in the flowchart below.

