

1.2.2024

### **Rules on reduction of fees according to Tariff no. 1554/2023**

According to Article 17 of Tariff no. 1554/2023 for marketing authorisations, annual fees and other licence fees for medicinal products, collected by the Icelandic Medicines Agency, the Agency may reduce tariff fees under special circumstances. The Agency shall lay down rules regarding prerequisites and arrangements concerning this matter, to be published on the Icelandic Medicines Agency's website.

#### **1) Discount for RMS registrations with no CMS and the medicinal product is marketed**

When a marketing authorisation is granted and Iceland is registered as RMS and no CMS is registered in the process, the Icelandic Medicines Agency collects a registration fee as in the case of a nationally registered medicinal product.

Granting of a discount in accordance with this provision is conditional on the relevant medicinal product or products being marketed in this country. A condition for this is that this is stated in the cover letter when applying for changes in variations for these marketing authorisations.

#### **2) Increase in the number of medicinal products marketed in Iceland**

The Icelandic Medicines Agency has been working for several years to increase the number of marketed medicinal products in Iceland, for both human and animal use. The aim has primarily been to fill gaps in the pharmaceutical market and reduce the risk of supply problems while increasing competition.

The Icelandic Medicines Agency provides the following resources, that applicants can apply to receive, to support this:

##### ***0-day process:***

A request for a simpler application process for a medicinal product can be submitted. Each request is examined separately with regard to whether the medicinal product in question is approved to undergo a simpler application process and a possible reduction in fees. The request shall be sent to [lyfjastofnun@lyfjastofnun.is](mailto:lyfjastofnun@lyfjastofnun.is)

##### ***Fee for marketing authorisation for a medicinal product in a 0-day process:***

If the medicinal product that is the subject of the application is listed in the tariff as an exempt medicinal product (or a comparable medicinal product; same active ingredient, pharmaceutical form and strength (formulation)) and has been approved for marketing authorisation through a 0-day process, a discount of 30% is granted on expense item 2.17 which applies to a 0-day process in the tariff.

***Fee for changes in variations of a medicinal product that has been granted marketing authorisation in a 0-day process:*** Fees for changes in variations for medicinal products that are granted marketing authorisation in a 0-day process and have been in use in the exemption system will be independent of

the number of strengths within the marketing authorisation, i.e. only one fee will be collected for the change even if it has an effect on more than one strength of the marketing authorisation. This applies while the medicinal product is the only one of its kind on the market. If another comparable medicinal product is put on the market this ceases to apply.

The 0-day process fee only applies to the strengths of the medicinal products that are marketed.

***Administrative/shortened renewal: Generic medicinal products***

In cases where an application is made for a shorter renewal process and all conditions regarding such process are met, a 20% discount on fees may be requested. Such request is submitted by writing "For this administrative renewal we apply for 20% discount" in the free text field in the cover letter with the application.

***Annual fees:***

It is possible to apply for a medicinal product to be exempt from the annual fee if it meets the following conditions:

- Information about the medicinal product must have been included in the pharmacopoeia on the previous 31 December.
- No medicinal product marketed in Iceland can replace the medicinal product in question.
- The total turnover (wholesale price w/o VAT) of all pharmaceutical forms/strengths of the medicinal product in the previous year is less than ISK 1,800,000.
- The pharmaceutical form/strength for which a reduction of the annual fee is requested was on the market (available) in this country for the entire previous year.

To apply for the above resource, applicants must fill out the form: "Application for reduction of annual fee".

A completed form together with the applicant's cover letter must be sent to the Icelandic Medicines Agency between 1 January and 20 January each year. No exemption from the annual fee will be granted for applications received by the Icelandic Medicines Agency after that time.

Each cover letter may be accompanied by application forms for medicinal products from the same marketing authorisation holder. Each form may cover all strengths of a single pharmaceutical form. The application must include information on the turnover of the medicinal product in the previous 2 years. Turnover means the total wholesale value of all packaging within the marketing authorisation number of the relevant medicinal product, VAT excluded.