

# STATUTORY INSTRUMENTS.

S.I. No. 551 of 2009

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IRISH MEDICINES BOARD (FEES) REGULATIONS 2009

### IRISH MEDICINES BOARD (FEES) REGULATIONS 2009

- I, MARY HARNEY, Minister for Health and Children, in exercise of the powers conferred on me by sections 13 and 32 of the Irish Medicines Board Act 1995 (No. 29 of 1995), as adapted by the Health (Alteration of Name of Department and Title of Minister) Order 1997, (S.I. No. 308 of 1997) and of all other powers enabling me in that behalf, hereby make the following Regulations:
- 1. These Regulations may be cited as the Irish Medicines Board (Fees) Regulations 2009.
  - 2. These Regulations shall come into force on the 1st day of January 2010.
  - 3. In these Regulations—

"Board" means the Irish Medicines Board;

"manufacturing licence" means a licence granted pursuant to the Medical Preparations (Licensing of Manufacture) Regulations 1993 to 1996;

"product authorisation" means an authorisation granted pursuant to the Medical Preparations (Licensing and Sale) Regulations 1998 (S.I. No. 142 of 1998);

"wholesale licence" means a licence granted pursuant to the Medical Preparations (Wholesale Licences) Regulations 1993 to 1996.

- 4. These Regulations shall apply to the fees that may be charged by the Board, in pursuit of its statutory duties, in relation to applications for the grant or renewal of manufacturing licences, wholesale licences and product authorisations in respect of medicinal products for human use.
- 5. Subject to Regulation 6 hereof, there shall be paid to the Board in respect of each and every matter set out in column 1 of the schedule hereto the fee as set out in column 2 of the said schedule.
- 6. The Board may, in circumstances where it considers it appropriate to do so, waive, remit or refund, either in whole or in part, any fee that would otherwise be payable to it under Regulation 5 hereof.
- 7. The Irish Medicines Board (Fees) Regulations 2008 (S.I. No. 542 of 2008) are hereby revoked.

# SCHEDULE

COLUMN 1	COLUMN 2
Fees for National Applications for Product Authorisations	€
National application — complex dossier, new active substance Each additional form (same time) Each additional strength (same time) Additional drug master file submitted with any of the above National application — reduced complex Each additional form (same time) Each additional strength (same time) Additional drug master file submitted with any of the above National application — reduced dossier standard Each additional form (same time) Each additional strength (same time) Additional drug master file submitted with any of the above Subsequent extension applications — first additional form Each additional form (same time) First additional strength (existing form) Each additional strength (same time) Additional drug master file submitted with any of the above	15,211 5,090 656 3,251 11,329 5,090 656 3,251 7,658 5,090 6,56 3,251 7,658 5,090 2,756 656 3,251
Fees for Applications for Product Authorisations using European Mutual Recognition procedure	
Mutual recognition incoming — complex dossier, new active substance Each additional form (same time) Each additional strength (same time) Outgoing mutual recognition supplement Decentralised Incoming Decentralised Outgoing Each additional form (same time) Each additional strength (same time) Mutual recognition incoming — reduced complex Each additional form (same time) Each additional strength (same time) Outgoing mutual recognition supplement Decentralised Incoming Decentralised Outgoing Each additional form (same time) Each additional strength (same time) Mutual recognition incoming — reduced dossier standard Each additional form (same time) Outgoing mutual recognition supplement Decentralised Incoming Decentralised Incoming Decentralised Incoming Decentralised Incoming Decentralised Outgoing Each additional form (same time) Each additional form (same time)	10,647 3,660 656 10,962 15,211 40,000 5,090 656 8,077 3,251 656 10,962 11,329 30,000 5,090 656 5,350 2,859 656 7,126 7,658 20,000 5,090 656
Subsequent extension applications  — mutual recognition incoming (first additional form)  — mutual recognition incoming (first additional strength)  — mutual recognition incoming (subsequent additional strength)  Outgoing mutual recognition/decentralised supplement (additional form)  Outgoing mutual recognition/decentralised supplement (additional strength)  Decentralised Incoming first additional form  Decentralised Outgoing — first additional form  Each additional form (same time)  First additional strength (existing form)  Each additional strength (same time)  Switching Applications	5,350 1,929 656 2,859 656 7,658 20,000 5,090 2,756 656 5,000

### **Fees for Parallel Product Authorisations**

Application fee — per country at the same time or by variation Each additional strength per country Each additional form per country Dual pack registration of parallel imports Each additional strength or form Parallel Imports where the originator is not on the Irish Market Change of Ownership	1,662 495 495 831 495 5,000 525
Fees for Variations to Product Authorisations that are nationally licensed	
Type IB variation Type IB variation — reduced rate Type II complex variation Type II standard variation Type II standard variation — reduced rate Notifications made under Article 61 (3) of Directive 2001/83/EC Notifications made under Article 61 (3) of Directive 2001/83/EC — reduced rate Bulk variation to multiple changes to the SPC (per product range) Bulk variation for multiple changes to the same document Introduction of standard statements from PHV working party — 1 to 5 licences Introduction of standard statements from PHV working party — 6 to 10 licences Introduction of standard statements from PHV working party — 21 to 40 licences Introduction of standard statements from PHV working party — 41 to 100 licences Introduction of standard statements from PHV working party — 41 to 100 licences Introduction of standard statements from PHV working party — 41 to 100 licences Introduction of standard statements from PHV working party — 101 and	520 260 3,251 633 316 495 248 7,200 5,400 1,500 3,000 6,000 12,000
above	30,000
Fees for Variations to Product Authorisations licensed under European Mutual Recognition procedure	
Type IB variation outgoing mutual recognition/decentralised supplement Type IB variation — mutual recognition incoming Type IB variation — mutual recognition incoming — reduced rate Type II complex variation — outgoing mutual recognition/decentralised supplement Type II complex variation — mutual recognition incoming Type II standard variation — outgoing mutual recognition/decentralised supplement Notifications made under Article 61 (3) of Directive 2001/83/EC Notifications made under Article 61 (3) of Directive 2001/83/EC — reduced rate	383 422 218 656 2,246 422 218 422 495
Fees for the granting of a Product Authorisation on transfer to another company	
Change of ownership — related company — per form Change of ownership — related company — per strength Change of ownership — non-related company — per form Change of ownership — non-related company — per strength	1,123 321 1,645 321
Other fees relating to the granting of Product Authorisations	
Service item	612
Fees for Wholesale Licences Application fee Annual fee — large site Annual fee — medium site Annual fee — minor site Variation to licence — minor site technical	555 2,771 1,576 400 400

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Variation to licence — administrative Variation to licence — technical	219 603
Fees for Manufacturing Licences	
Application fee Annual fee — major site (more than 250 employees) Annual fee — large site (150 - 250 employees) Annual fee — medium site (50 - 149 employees) Annual fee — small site (less than 50 employees) Variation to licence — administrative Variation to licence — technical	1,853 16,669 11,112 7,409 3,703 274 768
Fees for Blood and Tissue Establishments	
Application fee Annual fee — major site (more than 250 employees) Annual fee — large site (150 - 250 employees) Annual fee — medium site (50 - 149 employees) Annual fee — small site (less than 50 employees) Variation to licence — administrative Variation to licence — technical Appeal to amend/revoke a Tissue Establishment	1,853 16,669 11,112 7,409 3,703 274 768 500
Fees for Laboratory Approvals Application fee Annual fee — minor site Variation to licence — administrative Variation to licence — technical	555 555 219 603
Fees for the granting of a Manufacturing Licence or a Wholesale Licence on transfer to another company	
Manufacture — related companies Manufacture — unrelated companies Wholesale — related companies Wholesale — unrelated companies	1,107 1,853 365 555
Fees for Notifications of Exempt Medicinal Products	
Per notification Cap on total notifications	10,000
Fees for Medical Devices	
Certificates of free sale Certificates of free sale issued within 2 days (4 certs per request) Certificates of free sale issued within 1 day (4 certs per request) Additional certificates (available at the time of the initial request)	147 277 23
Registration of Devices  Electronic registration — Admin Fee  Annual Verification Fee — up to 5 employees  Annual Verification Fee — between 6-20 employees  Annual Verification Fee — between 21-100 employees  Annual Verification Fee — Over 100 employees	136 153 384 768 1,645
Clinical Investigations Clinical Investigations — active implatable medical devices Clinical Investigations — Class III and class IIb medical devices Clinical Investigations — Class IIa and class I medical device Clinical Investigations — Technical amendment to a previously approved clinical investigation Clinical Investigation — Administrative amendment to a previously approved clinical investigation  Audits/Inspections	3,837 3,837 1,645 1,129 219
Audits/Inspections (including Notified Body) per day Audits/Inspections (including Notified Body) per hour	1,489 213

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Classifications Classification of a product (1 product per request)	250
Classification of additional products (available at the time of the intial request)  Appeal of a classification decision	200 250
Designation Fee for a Notified Body	
Designation Fee	3,672
Extension to the scope (per extension)	1,836
Medicinal Product/Medical Device — Drug Consultation fees	
New active substance	41,960
Established active in new therapeutic area	15,000
Established active and therapeutic area	12,000
Variations — Minor	839
Variations — Major	3,776
Requests for Information	
Search fee of Medical Devices data base.	58



Given under my Official Seal, 22 December 2009

MARY HARNEY,

Minister for Health and Children.

### **EXPLANATORY NOTE**

(This note is not part of the instrument and does not purport to be a legal instrument.)

The purpose of these Regulations is to provide for the revision of fees payable to the Irish Medicines Board pursuant to Section 13 of the Irish Medicines Board Act 1995.

## BAILE ÁTHA CLIATH ARNA FHOILSIÚ AG OIFIG AN tSOLÁTHAIR

Le ceannach díreach ón

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