FEE STRUCTURE

In accordance with Article 74 of Regulation No 1107/2009 of the European Parliament and of the Council, and S.I. No 159 of 2012, the following fees shall apply from 1st September 2021

Fees relating to Plant Protection Products (PPP)

Description	Fee
PPP authorisation	
Authorisation of a product in the central zone (IE as zRMS)	€40,000 ¹
Art 43 re-authorisation of a product in the central zone (IE as zRMS)	€20,000 ¹
Authorisation of a product (IE as concerned MS) - no additional evaluation required	€2,500
Authorisation of a product (IE as concerned MS) - additional evaluation required	€3,500
Mutual recognition of product authorisation - no additional evaluation required	€2,500
Mutual recognition of product authorisation - additional evaluation required	€3,500
Authorisation of an Art 34 submission (exemption from the submission of studies)	€5,000 ²
Amendment to an authorisation (IE as zRMS)	€2,000 ²
Amendment to an authorisation (IE authorisation only)	€1,000 ²
Extension of authorisation for minor uses	€300
Administrative modification	€300
Notification - adjuvants	€300
Data matching check	€5,0001
Parallel Trade Permit - own use only	€100
Parallel Trade Permit - for sale to Third Parties	€300
Low Risk/ Micro-organism/ Biological PPP authorisation	
Authorisation of a product in the central zone (IE as zRMS)	€12,000 ¹
Art 43 re-authorisation of a product in the central zone (IE as zRMS)	€6,000
Authorisation of a product (IE as concerned MS)	€500
Mutual recognition of product authorisation	€500
Authorisation of an Art 34 submission (exemption from the submission of studies)	€500
Amendment to an authorisation (IE as zRMS)	€300 ²
Amendment to an authorisation (IE authorisation only)	€150 ²
Extension of authorisation for minor uses	€100
Administrative modification	€100
Permit for Trials	L
Approval to use a PPP in a trial(s) [APPPT]	€300
Amendment to approval to use a PPP in a trial(s) [APPPT]	€100
Test Facility Trial Permit [TFTO] (permit for a programme of experiments/trials)	€2,000
Annual Renewal of Test Facility Trial Permit [TFTP]	€300
Extension or variation of Test Facility Trial Permit [TFTP]	€300
A major amendment of late renewal of an existing Test Facility Trial Permit [TFTP]	€1,000
Test Facility Trial Permit for a 'once off/in-house' experiment	€300
Annual Renewal of Test Facility Trial Permit for a 'once off/in-house' experiment	€100
Extension or variation of Test Facility Trial Permit for a 'once off/in-house' experiment	€100
Annual Registration Fees ³	
Annual registration fee (ARF) – professional product	€225 *
Annual registration fee (ARF) – amateur product	€125 *
Re-instating product on the register/Late annual registration fee – professional product	€425 *
Re-instating product on the register/Late annual registration fee – amateur product	€225 *

Certification of Inclusion on the Register (Certificate of free sale)

* €25 will be transferred to the Poisons Information Centre to cover the cost of maintaining the Poisons database

€150

²An increased fee will be levied where expert evaluation is required.

³Fee for maintaining a plant protection product or adjuvant on the register (Regulation 13 of S.I. No 159 of 2012)

Fees for work on behalf of the European Union (Plant Protection Active Substance)

Description	Fee
Approval for a new active substance	
Pre-submission consultation/meeting	€1,000
Complete and sanitised dossier receipt, registry and admissibility check	€1,000
Co-ordination of examination of dossier and editing of DAR	€6,000
Examination of physical/chemical properties and analytical methods and peer review	€30,000
Examination of residues profile and peer review	€30,000
Examination of fate and behaviour in the environment and peer review	€80,000
Examination of ecotoxicological profile and peer review	€100,000
Examination of toxicological profile and peer review	€50,000
Examination of efficacy profile and peer review	€7,000
Endocrine assessment	€10,000
Total	€315,000
Co-Rapporteur Fee – IE as Co-RMS for evaluation or peer review	€50,000
Renewal of approval where IE is RMS	€185,000
Renewal of approval where IE is Co-RMS – for evaluation or peer review	€50,000
Import tolerance / MRL process where IE is RMS	€20,000 ⁴
Confirmatory information in support of EU approval where IE is RMS	€10,000
Assessment of equivalence of different sources of substance – Tier 1	€2,000
Assessment of equivalence of different sources of substance – Tier 1 & Tier 2	€3,500
Assessment of equivalence of different sources of substance – Tier 1 (post renewal of approval where IE is RMS)	€2,000 ⁵
Assessment of equivalence of different sources of substance - Tier 1 & Tier 2 (post renewal of approval where IE is RMS)	€3,500 ⁵

Low risk/ Micro-organisms/ Biological active substance

Approval of a new low risk/ micro-organism/ biological active substance	
Pre-submission consultation/meeting	€1,000
Complete and sanitised dossier receipt, registry and admissibility check	€1,000
Co-ordination of examination of dossier and editing of DAR	€6,000
Examination of physical/chemical properties and analytical methods and peer review	€9,000
Examination of residues profile and peer review	€9,000
Examination of fate and behaviour in the environment and peer review	€25,000
Examination of ecotoxicological profile and peer review	€30,000
Examination of toxicological profile and peer review	€15,000
Examination of efficacy profile and peer review	€4,000
Endocrine assessment	€5,000
Total	€105,000
Co-Rapporteur Fee – IE as Co-RMS for evaluation or peer review	€15,000
Renewal of approval where IE is RMS	€60,000
Renewal of approval where IE is Co-RMS – for evaluation or peer review	€15,000
Import tolerance / MRL process where IE is RMS	€10,000 ⁴
Confirmatory information in support of EU approval where IE is RMS	€3,000
Assessment of equivalence of different sources of substance	€500
Assessment of equivalence of different sources of substance (post renewal of approval where IE is RMS)	No fee