

Primary and Secondary New Product Applications, Submitted at the Same Time
(PC = Product Chemistry Data)
FY'25-FY'26 Fees

	Primary Application ¹		Secondary Application ²			
Description	PRIA Code	Application Fee (\$)	Where the only data submitted with a secondary application is product chemistry data or where the secondary application is a 100% repack of the primary		Where the secondary application contains more data than just PC such as efficacy and/or acute toxicity data	
			Agency Code ³	Expected Fee ⁴ (\$)	Agency Code ³	Expected Fee ⁴ (\$)
Registration Division						
New end-use or manufacturing-use product with registered source(s) of active ingredient(s); includes products containing two or more registered active ingredients previously combined in other registered products; excludes products requiring or citing an animal safety study; requires review of data package within RD only; includes data and/or waivers of data for only: <ul style="list-style-type: none">• product chemistry and/or• acute toxicity and/or• child resistant packaging and/or• pest(s) requiring efficacy – for up to 3 target pests	R310	10,990	R310.1	2,748	R310.2	10,990
New end use product containing up to three registered active ingredients never before registered as this combination in a formulated product; new product label is identical or substantially similar to the labels of currently registered products which separately contain the respective component active ingredients; excludes products requiring or citing an animal safety study; requires review of data package within RD only; includes data and/or waivers of data for only: <ul style="list-style-type: none">• product chemistry and/or• acute toxicity and/or• child resistant packaging and/or• pest(s) requiring efficacy for up to three target pests	R314	12,983	R314.1	3,246	R314.2	10,990

¹ Each new product application is subject to a PRIA fee. Where one set of data or data waivers apply to two or more new product applications that are submitted at the same time, the Agency refers to the first product application containing the data or data waivers as the primary application.

² Additional new product applications that rely on data or data waivers that were submitted with the primary product application are referred to as secondary applications.

³ EPA will assign a tracking code to alert reviewers to the relationship between primary and secondary applications. These codes are internal EPA tracking codes only.

⁴ Based on previous years of experience, EPA expects that it can grant a discretionary refund that will likely result in a reduced fee equal to the amount indicated in this column. This expected fee is based on either the fee for an identical/substantially similar product with no data review for the type of product (i.e. conventional, antimicrobial, or biopesticide) or 25% of the fee for the primary, whichever is greater and rounded up to the nearest whole dollar. In accordance with FIFRA 33(b)(2)(C), payment of at least 25% of the fee for the applicable PRIA category accompanied by a request for a refund of all or part of the remaining fee would allow this application to go forward into review. **Where this chart indicates the expected fee is more than 25%, EPA recommends submitting the amount of the expected fee as listed in this column along with a request for a refund to avoid delays in processing applications for which a complete fee has not been received.**

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			Agency Code ³	Expected Fee ⁴ (\$)	Agency Code ³	Expected Fee ⁴ (\$)
New end-use product containing up to three registered active ingredients never before registered as this combination in a formulated product; new product label is identical or substantially similar to the labels of currently registered products which separately contain the respective component active ingredients; excludes products requiring or citing an animal safety study; requires review of data package within RD only; includes data and/or waivers of data for only: <ul style="list-style-type: none"> • product chemistry and/or • acute toxicity and/or • child resistant packaging and/or • pest(s) requiring efficacy for 4 to 7 target pests 	R319	19,002	R319.1	4,751	R319.2	10,990
New end-use product containing four or more registered active ingredients never before registered as this combination in a formulated product; new product label is identical or substantially similar to the labels of currently registered products which separately contain the respective component active ingredients; excludes products requiring or citing an animal safety study; requires review of data package within RD only; includes data and/or waivers of data for only: Only: <ul style="list-style-type: none"> • product chemistry and/or • acute toxicity and/or • child resistant packaging and/or • pest(s) requiring efficacy – for up to 3 target pests 	R318	19,994	R318.1	4,986	R318.2	10,990

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			Agency Code ³	Expected Fee ⁴ (\$)	Agency Code ³	Expected Fee ⁴ (\$)
New end-use product containing four or more registered active ingredient(s) never before registered as this combination in a formulated product; new product label is identical or substantially similar to the labels of currently registered products which separately contain the respective component active ingredients; excludes products requiring or citing an animal safety study; and requires review of data and/or waivers for only: <ul style="list-style-type: none"> • product chemistry and/or • acute toxicity and/or • child resistant packaging and/or • pest(s) requiring efficacy – for 4 to 7 target pests 	R321	25,964	R321.1	6,491	R321.2	10,990
New end-use on-animal product, registered source of active ingredient(s) with submission of data and/or waivers for only: <ul style="list-style-type: none"> • animal safety and • pest(s) requiring efficacy and/or • product chemistry and/or • acute toxicity and/or • child resistant packaging 	R315	14,779	R315.1	3,695	R315.2	10,990
New end-use or manufacturing-use product with registered source(s) of active ingredient(s) including products containing two or more registered active ingredient previously combined in other registered products; excludes products requiring or citing an animal safety study; and requires review of data and/or waivers for only: <ul style="list-style-type: none"> • product chemistry and/or • acute toxicity and/or • child resistant packaging and/or • pest(s) requiring efficacy for 4 to 7 target pests 	R316	17,009	R316.1	4,253	R316.2	10,990

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			Agency Code ³	Expected Fee ⁴ (\$)	Agency Code ³	Expected Fee ⁴ (\$)
New end-use or manufacturing-use product with registered source(s) of active ingredient(s) including products containing two or more registered active ingredient previously combined in other registered products; excludes products requiring or citing an animal safety study; and requires review of data and/or waivers for only: <ul style="list-style-type: none"> • product chemistry and/or • acute toxicity and/or • child resistant packaging and/or • pest(s) requiring efficacy – for greater than 7 target pests 	R317	23,029	R317.1	5,758	R317.2	10,990
New product; new physical form; requires data review in science divisions	R320	19,906	R320.1	4,977	R320.2	10,990
New product; repack of identical registered end-use product as a manufacturing-use product; same registered uses only	R331	3,809	R331	N/A	R331	N/A
New manufacturing-use product; registered active ingredient; unregistered source of active ingredient; submission of completely new generic data package; registered uses only; requires review in RD and science divisions	R332	426,215	R332.1	106,554	R332.2	106,554
New product; manufacturing-use product or end-use product with unregistered source of active ingredient; requires science data review; new physical form; etc. Cite-all or selective data citation where applicant owns all required data	R333	29,856	R333.1	7,464	R333.2	10,990
New product; manufacturing-use product or end-use product with unregistered source of the active ingredient; requires science data review; new physical form; etc. Selective data citation.	R334	34,764	R334.1	8,691	R334.2	10,990

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			Agency Code ³	Expected Fee ⁴ (\$)	Agency Code ³	Expected Fee ⁴ (\$)
New end-use product containing up to three registered active ingredients never before registered as this combination in a formulated product; new product label is identical or substantially similar to the labels of the currently registered products which separately contain the respective component active ingredients; excludes products requiring or citing an animal safety study; requires review of data package within RD only; includes data and/or waivers of data for only: <ul style="list-style-type: none"> • product chemistry and/or • acute toxicity and/or • child resistant packaging and/or • pest(s) requiring efficacy – for more than 7 target pests 	R361	24,570	R361.1	6,143	R361.2	10,990
New end-use product containing four or more registered active ingredients never before registered as this combination in a formulated product; new product label is identical or substantially similar to the labels of the currently registered products which separately contain the respective component active ingredients; excludes products requiring or citing an animal safety study; requires review of data package within RD only; includes data and/or waivers of data for only: <ul style="list-style-type: none"> • product chemistry and/or • acute toxicity and/or • child resistant packaging and/or • pest(s) requiring efficacy – for more than 7 target pests 	R362	26,618	R362.1	6,655	R362.2	10,990
New product; repack of identical registered manufacturing-use product as an end-use product; same registered uses only, with no additional data.	R363	8,190	R363.1	N/A	R363.2	N/A

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Description	PRIA Code	Application Fee (\$)	Where the only data submitted with a secondary application is product chemistry data or where the secondary application does not require confirmatory efficacy data		Where the secondary application contains more data than just PC such as efficacy and/or acute toxicity data	
			Agency Code ³	Expected Fee ⁴ (\$)	Agency Code ³	Expected Fee ⁴ (\$)
Antimicrobials Division						
New end use product; FIFRA §2(mm) uses only; 0 to 10 public health organisms	A460	7,689	A460.1	1,923	A460.2	7, 689
New end use product; FIFRA §2(mm) uses only; 11 to 20 public health organisms	A461	10,666	A461.1	2,667	A461.2	7, 689
New end use product; FIFRA §2(mm) uses only; 21 to 30 public health organisms	A462	13,645	A462.1	3,412	A462.2	7, 689
New end use product; FIFRA §2(mm) uses only; 31 to 40 public health organisms	A463	16,623	A463.1	4,156	A463.2	7, 689
New end use product; FIFRA §2(mm) uses only; 41 to 50 public health organisms	A464	19,602	A464.1	4,901	A464.2	7, 689
New end use product; FIFRA §2(mm) uses only; 51 or more public health organisms	A465	22,581	A465.1	5,646	A465.2	7, 689
New end-use product; uses other than FIFRA §2(mm); non-FQPA product	A550 A550	13,266 19,906	A550.1 A550.1	3,307 4,977	A550.2 A550.2	5,107 7, 689
New manufacturing-use product; registered active ingredient; selective data citation	A560	18,957	A560.1	4,740	A560.2	7, 689
New manufacturing-use product; registered active ingredient; unregistered source of active ingredient; submission of new generic data package; registered uses only; requires science review	A565	27,442	A565.1	6,861	A565.2	7, 689
New Product or amendment requiring data review for risk assessment by Science Branch (e.g., changes to Restricted Entry Interval, or Personal Protective Equipment, or use rate)	A572	19,906	A572.1	4,977	A572.2	7, 689

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Biopesticides and Pollution Prevention Division						
New product; registered source of active ingredient(s); no change in an established tolerance or tolerance exemption; (including non-food); must address product specific data requirements.	B670	7,689	B670.1	1,923	B670.2	7,689
New product; unregistered source of at least one active ingredient (or registered source with new generic data package); no change to an established tolerance or tolerance exemption (including non-food); must address product specific and generic data requirements.	B672	13,723	B672.1	3,431	B672.2	7,689
New product; unregistered source of active ingredient(s); citation of Technical Grade Active Ingredient (TGAI) data previously reviewed and accepted by the Agency; requires an Agency determination that the cited data support the new product.	B673	7,689	B673.1	1,923	B673.2	7,689
New end-use non-food animal product with submission of two or more target animal safety studies; includes data and/or waivers of data for only: <ul style="list-style-type: none">– product chemistry and/or– acute toxicity and/or– public health pest efficacy and/or– animal safety studies and/or– child resistant packaging	B677	13,276	B677.1	3,319	B677.2	7,689
SCLP; new product; unregistered source of active ingredient; no change in an established tolerance or tolerance exemption (including non-food); must address product specific data requirements.	B721	4,028	B721.1	1,007	B721.2	4,028
Registration application; registered PIP; new product or new terms of registration; additional data submitted; no petition since a permanent tolerance/tolerance exemption is already established for the active ingredient(s).	B880	48,024	B880.1	12,006	B880.2	12,006
Registration application; registered PIP, seed increase; breeding stack of previously approved PIPs, same crop; no petition since a permanent tolerance/tolerance exemption is already established for the active ingredient(s).	B885	48,024	B885.1	12,006	B885.2	12,006
Registration application; new product, registered active ingredient; exogenous applications of RNA to elicit the RNA interference pathway in pests; no petition since a permanent tolerance/tolerance exemption is already Established for the active ingredient(s).	B929	7,689	B929.1	1,923	B929.2	7,689

Example

A company submits 3 new antimicrobial registration applications. The 3 applications are: one 20% concentrate, a 15% concentrate and a 10% concentrate. The package consists of chemistry data for each application, one set of acute toxicity studies using the 20% concentrate and one set of efficacy data generated at the use dilution (the use dilution is the same for all three products). All three products will rely on the same efficacy data because all three products will be diluted to the same concentration and the difference in the inert ingredients is water.

Description of action	Expected Fee (\$)	Tracking Code
New product conc 20%	7, 689	A460
New product conc 15%	1,923	A460.1
New product conc 10%	1,923	A460.1
Total Fee	11,535	