Proposed Registration Decision

Santé

Canada

PRD2017-11

Bifenthrin and Capture 240 EC

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Overview

Proposed Registration Decision for Bifenthrin

Health Canada's Pest Management Regulatory Agency (PMRA), under the authority of the Pest Control Products Act and Regulations, is proposing cancellation of the registration of Bifenthrin Technical Insecticide and Capture 240 EC, containing the technical grade active ingredient bifenthrin. The use of bifenthrin on potatoes to control wireworm will be cancelled as of 31 December 2017. As the use on raspberries in British Columbia has been identified as a critical need in Canada, the PMRA is proposing registration for the sale and use of Bifenthrin Technical Insecticide and Capture 240 EC, for a period of three years to allow for the phase out of this use.

Bifenthrin Technical Insecticide (Registration Number 31395) and Capture 240 EC (Registration Number 31396) are conditionally registered in Canada. At the time of the original registration, the evaluation of the scientific information demonstrated that Capture 240 EC had value and that human health and environmental risk was acceptable. However, there remained some uncertainty with respect to the criteria for Track 1 classification under Canada's Toxic Substances Management Policy (TSMP). As such, additional information was required and a conditional registration was granted, but was limited to uses on potato and raspberry only, which were identified as critical needs by Canadian growers. The current applications were submitted to address the requested information and to convert Bifenthrin Technical Insecticide and Capture 240 EC from conditional registration to full registration.

The evaluation of the additional environmental information found that bifenthrin meets the criteria for Track 1 substances under the TSMP. The PMRA's implementation of the TSMP is outlined in Regulatory Directive DIR99-03. This directive describes how Track 1 substances will be managed by the PMRA, and calls for the virtual elimination of Track 1 substances. For use of Capture 240 EC for control of weevils on raspberries, a phase-out period of three years is proposed. Since no suitable alternative exists for raspberries, the PMRA has determined that bifenthrin meets the critical need criteria as outlined in DIR99-03, which would allow for its use on a time-limited basis with the imposition of restrictions on the registration designed to minimize the risks associated with its use. Specific risk mitigation measures designed to minimize the risks include: limiting the use to a small geographical location (raspberries in British Columbia), buffer zones to minimize spray drift, vegetative filter strips to reduce runoff, and prohibiting application during crop blooming period. Therefore, registration for the use of Capture 240 EC on raspberries is granted for use only in British Columbia for three years beginning 1 January 2018 until 31 December 2020. The use of Capture 240 EC for control of wireworm on potato will be cancelled as of 31 December 2017.

This Overview describes the key points of the evaluation, while the Science Evaluation provides detailed technical information on the human health, environmental and value assessments of bifenthrin and Capture 240 EC.

Health Canada. DIR99-03, The Pest Management Regulatory Agency's Strategy for Implementing the Toxic Substances Management Policy (TSMP).

What Does Health Canada Consider When Making a Registration Decision?

The key objective of the *Pest Control Products Act* is to prevent unacceptable risks to people and the environment from the use of pest control products. Health or environmental risk is considered acceptable² if there is reasonable certainty that no harm to human health, future generations or the environment will result from use or exposure to the product under its proposed conditions of registration. The Act also requires that products have value³ when used according to the label directions. Conditions of registration may include special precautionary measures on the product label to further reduce risk.

To reach its decisions, the PMRA applies modern, rigorous risk-assessment methods and policies. These methods consider the unique characteristics of sensitive subpopulations in humans (for example, children) as well as organisms in the environment. These methods and policies also consider the nature of the effects observed and the uncertainties when predicting the impact of pesticides. For more information on how the PMRA regulates pesticides and the assessment process, please visit the Pesticides and Pest Management portion of Health Canada's website at healthcanada.gc.ca/pmra.

Before making a final registration decision on bifenthrin, the PMRA will consider any comments received from the public in response to this consultation document.⁴ The PMRA will then publish a Registration Decision⁵ on bifenthrin, which will include the decision, the reasons for it, a summary of comments received on the proposed final registration decision and the PMRA's response to these comments.

For more details on the information presented in this Overview, please refer to the Science Evaluation of this consultation document.

What Is Bifenthrin?

Bifenthrin is an insecticide found in the commercial class product Capture 240 EC that is used to control wireworms in potatoes and several pests in raspberries, particularly those that are present at the time of harvest.

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² "Acceptable risks" as defined by subsection 2(2) of the *Pest Control Products Act*.

[&]quot;Value" as defined by subsection 2(1) of the *Pest Control Products Act*: "... the product's actual or potential contribution to pest management, taking into account its conditions or proposed conditions of registration, and includes the product's (a) efficacy; (b) effect on host organisms in connection with which it is intended to be used; and (c) health, safety and environmental benefits and social and economic impact."

⁴ "Consultation statement" as required by subsection 28(2) of the *Pest Control Products Act*.

⁵ "Decision statement" as required by subsection 28(5) of the *Pest Control Products Act*.

Health Considerations

Can Approved Uses of Bifenthrin Affect Human Health?

Capture 240 EC, containing bifenthrin, is unlikely to affect your health when used according to label directions.

Potential exposure to bifenthrin containing products may occur through the diet (food and water) or when handling, applying or entering treated areas. When assessing health risks, two key factors are considered:

- the levels where no health effects occur in animal testing, and
- the levels to which people may be exposed.

The dose levels used to assess risks are established to protect the most sensitive human population (for example, children and nursing mothers). As such, sex and gender are taken into account in the risk assessment. Only uses for which the exposure is well below levels that cause no effects in animal testing are considered acceptable for registration.

Toxicology studies in laboratory animals describe potential health effects from varying levels of exposure to a chemical and identify the dose where no effects are observed. The health effects noted in animals occur at doses more than 100-times higher (and often much higher) than levels to which humans are normally exposed when pesticide-containing products are used according to label directions.

In laboratory animals, the technical grade active ingredient, bifenthrin, was of high acute toxicity by the oral route; consequently, the signal word and hazard statement "DANGER – POISON" are required on the label. It was of low toxicity via the dermal route and slightly toxic via the inhalation route. Bifenthrin was non-irritating to the eyes and skin. Bifenthrin caused an allergic skin reaction.

The end-use product Capture 240 EC was of high acute toxicity via the oral route; consequently, the signal word and hazard statement "DANGER - POISON" are required on the label. It was of low toxicity via the dermal route, and of slight toxicity via the inhalation route. Capture 240 EC was minimally irritating to the eyes, slightly irritating to skin, and a potential dermal sensitizer based on a lack of acceptable sensitization test. Consequently, the signal work and hazard statement "POTENTIAL SKIN SENSITIZER", are required on the label.

Registrant-supplied short, and long term (lifetime) animal toxicity tests, as well as information from published scientific literature were collectively assessed for the potential of bifenthrin to cause neurotoxicity, immunotoxicity, chronic toxicity, cancer, reproductive and developmental toxicity, and various other effects. The most sensitive endpoint used for risk assessment was neurotoxicity. There is some concern for increased sensitivity of the young exposed to bifenthrin. Data suggest that immature detoxification mechanisms in the young may lead to the accumulation of bifenthrin and increased toxicity at high levels of exposure.

The risk assessment protects against these and any other potential effects by ensuring that the level of human exposure is well below the lowest dose at which these effects occur in animal tests.

Residues in Water and Food

Dietary risks from food and drinking water are not of health concern.

Aggregate dietary intake estimates (food plus drinking water) revealed that the general population and children 1-2 years old, the subpopulation which would ingest the most bifenthrin relative to body weight, are expected to be exposed to less than 34% of the acceptable daily intake. Based on these estimates, the chronic non-cancer dietary risks from bifenthrin are not of health concern for all population subgroups.

Acute dietary (food plus drinking water) intake estimates for the general population and all population subgroups were less than 75% of the acute reference dose, and are not of health concern. The highest exposed subpopulation was children 1-2 years old.

The *Food and Drugs Act* prohibits the sale of adulterated food, that is, food containing a pesticide residue that exceeds the established maximum residue limit (MRL). Pesticide MRLs are established for *Food and Drugs Act* purposes through the evaluation of scientific data under the *Pest Control Products Act*. Food containing a pesticide residue that does not exceed the established MRL does not pose an unacceptable health risk.

Residue trials conducted throughout the United States using bifenthrin on various crops and trials conducted on tea in India are acceptable.

For the listing of MRLs for this active ingredient on crop commodities, please refer to the Maximum Residue Limit Database in the Pesticides and Pest Management section of Health Canada's website.

Occupational Risks from Handling Bifenthrin

Occupational risks are not of concern when bifenthrin is used according to the label directions, which include protective measures.

Farmers and custom applicators who mix, load or apply Capture 240 EC as well as field workers re-entering freshly treated fields can come in direct contact with bifenthrin residues on the skin and/or through inhalation. Therefore, the label specifies that a long-sleeved shirt, long pants, chemical resistant gloves, shoes plus socks and goggles must be worn during mixing, loading, application, clean up and repair. The label also requires that workers not re-enter treated fields for 12 hours after application except for hand harvesting of raspberries where workers cannot reenter for 3 days. Taking into consideration these label statements, the precautionary measures, the number of applications and the expectation of the exposure period for handlers and workers, it was determined that the risks to these individuals are not a concern.

Risks in Residential and Other Non-Occupational Environments

For bystanders, exposure is expected to be much less than that for workers and the potential for drift from agricultural areas is expected to be minimal. Therefore, health risks to bystanders are not of concern. Applications are limited to agricultural crops only when there is low risk of drift to areas of human habitation or activity, such as houses, cottages, schools and recreational areas, taking into consideration wind speed, wind direction, temperature inversions, application equipment and sprayer settings.

Environmental Considerations

What Happens When Bifenthrin Is Introduced into the Environment?

Bifenthrin meets the Government of Canada's criteria for a Track 1 Substance under TSMP. These criteria determine if a substance is toxic, takes a long time to break down and accumulates in living organisms. Because of these environmental concerns it is proposed that the registered use of bifenthrin on potatoes and raspberries be cancelled by 31 December 2017 and 2020, respectively. Interim risk reduction measures are proposed to further minimize environmental exposure during the phase out period.

Bifenthrin enters into the environment when applied as a foliar spray to raspberries and an infurrow planting treatment to potatoes.

When bifenthrin is released into the environment, it can enter soil and surface water where it can persist under certain conditions. In soil, bifenthrin can be broken down slowly by soil bacteria. It binds strongly to soil particles, making it unlikely to move downward in the soil and reach groundwater. In aquatic environments, bifenthrin moves rapidly out of water and into sediment where it can persist. The vapour pressure and Henry's law constant of bifenthrin suggest that bifenthrin has low potential to volatilize from water and moist soil.

In the terrestrial environment, bifenthrin does not pose a risk to earthworms, birds, and plants; however, small mammals, beneficial insects, including bees, could be at risk if they come into direct contact or residues on plants. Precautionary label statements are required to inform users of the potential hazards to mammals and bees. The risk to bees can be reduced by restricting or prohibiting the application of bifenthrin during the crop blooming period.

In the aquatic environment aquatic plants are not at risk; however freshwater and marine fish and aquatic invertebrates are at potential risk on an acute and chronic basis. Spray buffer zones are required to protect aquatic organisms from spray drift. To reduce the risks of bifenthrin being carried in runoff to aquatic environments, in addition to precautionary label statements, a mandatory requirement for the construction and maintenance of a 10-metre vegetative filter strip between the area of application and waterbodies is required.

The persistency and bioaccumulation characteristics of bifenthrin were both above and below the cut-off values for the criteria for Track 1 substances under the TSMP. Therefore, PMRA requested additional environmental data to examine accumulation of bifenthrin by aquatic organisms through diets and to further characterize bifenthrin's behaviour and risks in the environment. The PMRA also requested other environmental data identified by the European Commission (2012).

Bifenthrin has the potential to accumulate in the tissues of organisms through various routes of exposures to levels that could cause effects. Characteristics such as persistence, accumulation in animal tissues and effects suggest that bifenthrin may be of concern if animals are exposed over a long period of time. Based on the available environmental information, PMRA concluded that bifenthrin meets the Government of Canada's criteria for a Track 1 Substance under the TSMP adopted in 1995. These criteria are also outlined in the *Persistence and Bioaccumulation* Regulations (SOR/2000-107) which are used to define persistent or bioaccumulative under sections of the Canadian Environmental Protection Act (CEPA, 1999). Under TSMP, substances meeting these criteria (Track 1) are persistent, bioaccumulative, toxic and primarily the result of human activity. Track 1 substances are targeted for vitual elimination from the environment, but may be registered for a limited time in exceptional circumstances (for example: critical need). The critical need for bifenthrin on raspberries for pre-harvest control in British Columbia continues to exist as there are no suitable alternatives. The need for bifenthrin on potatoes is not considered critical as alternatives are now registered. It is proposed that the registered uses of bifenthrin on potatoes and raspberries be cancelled by 31 December 2017 and 2020, respectively, because of these environmental concerns. Interim risk reduction measures are proposed to minimize environmental exposure during the phase out period.

Value Considerations

What Is the Value of Capture 240 EC?

Capture 240 EC controls wireworms, which are a major pest of potatoes, and several insects that are harvest contaminants in raspberries.

When Capture 240 EC was registered in 2014, Canadian growers identified a soil application at the time of planting to control wireworms in potato as a high priority. Since then, a new product containing phorate has provided growers with an additional management tool for wireworms.

Canadian growers identified a need for pre-harvest control of weevils and lepidopteran larvae in raspberries at the time that Capture 240 EC was first registered. There continues to be zero tolerance for insect harvest contaminants in raspberries and no suitable alternatives available to control weevils, the most prevalent contaminant. As such, foliar application of Capture 240 EC prior to harvest is considered to be critical for the management of insect contaminants of raspberries in British Columbia.

Measures to Minimize Risk

Labels of registered pesticide products include specific instructions for use. Directions include risk-reduction measures to protect human and environmental health. These directions must be followed by law.

The key risk-reduction measures being proposed on the label of Capture 240 EC to address the potential risks identified in this assessment are as follows.

Key Risk-Reduction Measures

Human Health

Because there is a concern with users coming into direct contact with bifenthrin on the skin or through inhalation of spray mists, anyone mixing, loading and applying bifenthrin and performing cleaning and repair activities must wear long-sleeved shirt, long pants, chemical resistant gloves, shoes plus socks and goggles. In addition, standard label statements to protect against drift during application are present on the label.

Environment

- Environmental hazard statements for bees, beneficial insects, small mammals and aquatic organisms are required.
- To reduce risk to pollinators, application is prohibited during the crop blooming period.
- Spray buffer zones ranging from 10 to 75 metres for non-target aquatic habitats are required. For all ground field sprayer use restrictions include the use of low drift air induction nozzles only, a minimum ASAE (American Society of Agricultural Engineers) medium spray quality and a wind speed restriction at the time of application (<8 km/hr).
- To reduce the potential for runoff of bifenthrin to adjacent aquatic habitats, precautionary statements for sites with characteristics that may be conducive to runoff and when heavy rain is forecasted, are required. In addition, a minimum 10-metre vegetative filter strip between the treatment area and the edge of a water body is required to reduce runoff of bifenthrin to aquatic environments.

Value

Limit the use of Capture 240 EC to raspberries grown in British Columbia for a period of three years.

Next Steps

Before making a final registration decision on bifenthrin, the PMRA will consider any comments received from the public in response to this consultation document. The PMRA will accept written comments on this proposal up to 45 days from the date of publication of this document. Please forward all comments to Publications (contact information on the cover page of this document).

The PMRA will then publish a Registration Decision, which will include its decision, the reasons for it, a summary of comments received on the proposed final decision and the Agency's response to these comments.

Other Information

When the PMRA makes its registration decision, it will publish a Registration Decision on bifenthrin (based on the Science Evaluation of this consultation document). In addition, the test data referenced in this consultation document will be available for public inspection, upon application, in the PMRA's Reading Room (located in Ottawa).

Science Evaluation

Bifenthrin

1.0 The Active Ingredient, Its Properties and Uses

1.1 **Identity of the Active Ingredient**

Active substances Bifenthrin and its trans isomer

Function Miticide/Insecticide

Chemical name

1. International bifenthrin:

Union of Pure (2-methylbiphenyl-3-yl)methyl (1RS,3RS)-3-[(1Z)-2-chloro-3,3,3trifluoroprop-1-en-1-yl]-2,2-dimethylcyclopropanecarboxylate and Applied **Chemistry**

(IUPAC) trans isomer:

> (2-methylbiphenyl-3-yl)methyl (1S,3R)-3-[(1Z)-2-chloro-3,3,3trifluoroprop-1-en-1-yl]-2,2-dimethylcyclopropanecarboxylate or

> (2-methylbiphenyl-3-yl)methyl (1*R*,3*S*)-3-[(1*Z*)-2-chloro-3,3,3trifluoroprop-1-en-1-yl]-2,2-dimethylcyclopropanecarboxylate

2. Chemical bifenthrin:

Abstracts Service cyclopropanecarboxylic acid, 3-[(1Z)-2-chloro-3,3,3-trifluoro-1propen-1-yl]-2,2-dimethyl-, (2-methyl[1,1'-biphenyl]-3-yl)methyl (CAS)

ester, (1RS,3RS)-

trans isomer (double bond geometry unknown):

cyclopropanecarboxylic acid, 3-(2-chloro-3,3,3-trifluoro-1propenyl)-2,2-dimethyl-, (2-methyl[1,1'-biphenyl]-3-yl)methyl

ester, trans- (9Cl)

CAS number 82657-04-3 (bifenthrin)

83322-02-5 (trans isomer)

Molecular formula C₂₃H₂₂ClF₃O₂

Molecular weight 422.88 Structural formula bifenthrin:

Purity of the active 95.5% ingredients

1.2 Physical and Chemical Properties of the Active Ingredients and End-Use Product

Technical Product—Bifenthrin Technical

Property	Result
Colour and physical state	Off-white solid
Odour	Fluoro-chloro like smell
Melting range	66.6-69.0°C
Boiling point or range	N/A
Density	1.316 g/cm ³
Vapour pressure at 25°C	$2.413 \times 10^{-5} \text{ Pa}$
Henry's law constant at 20°C	1.01×10^{-4} atm m ³ /mol
	$1/H = 9901 \text{ mol/ atm m}^3$
Ultraviolet (UV)-visible spectrum	$\lambda \text{max} = 250 \text{ nm}, \ \epsilon = 3282.9$
	no absorption was observed above 300 nm.
Solubility in water at 20°C	<u>pH</u> Solubility (μg/L)
	4.05 <1
	7.18 <1
	9.20 3.76
Solubility in organic solvents at	Solvent Solubility
20°C (g/L)	Methanol 48.0±0.7
	Xylene 556.3±29.3
	Acetone 735.7±47.2
	1,2-dichloroethane 743.2±16.4
	Ethyl acetate 579.8±47.2
	n-heptane 144.5±9.0
<i>n</i> -Octanol-water partition	$\log K_{\rm ow} = 8.0$ at 20° C
coefficient (K_{ow})	
Dissociation constant (pK_a)	Practically insoluble in water and no dissociable groups
Stability	Thermally stable at >200°C. No corrosion to phenoxy resin
(temperature, metal)	lined HDPE or steel. It is subject to photo-degradation in an
	aqueous environment. Isomer conversion and ester cleavage
	are very slow under natural sunlight without a
	photosensitizer.

End-Use Product—Capture 240 EC

Property	Result
Colour	Straw yellow
Odour	Mild naphthenic odour

Property	Result
Physical state	Liquid
Formulation type	Emulsifiable concentrate
Guarantee	240 g/L
Container material and description	Fluorinated HDPE bottle, 2.5-5 L
Density	0.954 g/ml
pH of 1% dispersion in water	4.6
Oxidizing or reducing action	The product does not contain any oxidants or reductants.
Storage stability	The product is stable to fluorinated HDPE bottles for 2 years under ambient temperature.
Corrosion characteristics	The emulsifiable concentrate is compatible with glass and fluorinated plastic containers.
Explodability	Non-explosive

1.3 Directions for Use

Potato

One in-furrow application of Capture 240 EC applied at planting at a rate range of 2.0–3.4 g a.i./100 m row (222–337 g a.i./ha) controls wireworms in potatoes, with low application rates for use under light to moderate pest pressure and high application rates for heavy pest pressure.

Raspberry

Foliar application of Capture 240 EC at a rate of 112 g a.i./ha controls obliquebanded leafroller, orange tortrix, black vine weevil, obscure root weevil, clay coloured weevil and earwigs in raspberries. One application may be made pre-bloom and a second may be made post-bloom with a minimum re-application interval of 30 days. A maximum of 224 g a.i./ha may be applied per year.

1.4 Mode of Action

Bifenthrin is a pyrethroid in the Insecticide Resistance Action Committee Mode of Action (MOA) Group 3A. Bifenthrin interferes with sodium channels within nerve cell membranes of insects, causing paralysis and death. This active ingredient acts by contact or ingestion.

2.0 Methods of Analysis

2.1 Methods for Analysis of the Active Ingredient

The methods provided for the analysis of the active ingredient and the impurities in Bifenthrin Technical have been validated and assessed to be acceptable for the determinations.

2.2 Method for Formulation Analysis

The methods provided for the analysis of the active ingredient in the formulation have been validated and assessed to be acceptable for use enforcement analytical methods.

2.3 Methods for Residue Analysis

High-performance liquid chromatography methods with tandem mass spectrometry (HPLC-MS/MS) were developed and proposed for data generation and enforcement purposes. These methods fulfilled the requirements with regards to selectivity, accuracy and precision at the respective method limit of quantitation. Acceptable recoveries (70–120%) were obtained in plant and animal matrices and environmental media.

Gas chromatography methods with electron capture or mass spectrometric detection (GC-ECD; GC-MSD; Methods P-0757/P-1073 in plant matrices and Methods P-1031 (Revised), R-1843M (Revised) and RAN-0204M in animal matrices) were developed and proposed for data generation and enforcement purposes. These methods fulfilled the requirements with regards to specificity, accuracy and precision at the respective method limit of quantitation. Acceptable recoveries (70-120%) were obtained in plant and animal matrices. The proposed enforcement methods were successfully validated in plant and animal matrices by an independent laboratory. Extraction solvents used in the method were similar to those used in the metabolism studies; thus, further demonstration of extraction efficiency with radiolabelled crops was not required for the enforcement method.

Methods for residue analysis are summarized in Appendix I, Table 1.

3.0 Impact on Human and Animal Health

3.1 Toxicology Summary

Bifenthrin is a synthetic Type I pyrethroid insecticide. Synthetic pyrethroids induce neurotoxic effects primarily by binding to voltage-dependant sodium channels in neurons. Binding to the receptor delays closing of sodium channels, causing the depolarization of neurons. This affects action potentials and results in repetitive activity. Type I pyrethroids, such as bifenthrin, typically induce "T-syndrome" which is characterized by rapid onset of aggressive behavior, increased sensitivity to external stimuli, fine tremor, prostration with coarse whole body tremor, elevated body temperature, coma, and death.

A detailed review of the toxicological database for bifenthrin was conducted. The database is complete, consisting of the full array of toxicity studies currently required for hazard assessment purposes. The majority of the submitted studies were carried out in accordance with accepted international testing protocols and Good Laboratory Practices. The scientific quality of the data is acceptable and the database is considered adequate to define the majority of toxic effects that may result from exposure to bifenthrin. The database was supplemented by published scientific literature studies.

Metabolism and toxicokinetics were investigated using radiolabelled bifenthrin in standard single

low- and high-dose, as well as repeated low- and high-dose oral gavage studies in rats. In addition to these studies, a longer term study was conducted in which female rats were dosed with bifenthrin for 70 days, followed by an 85 day period without dosing. Absorption was low. Most of the administered dose was recovered within 48-72 hours of dosing, with negligible amounts of radioactivity in expired air. The majority of test material was excreted in the feces (25-96%) as unchanged bifenthrin and metabolites. In bile duct cannulated rats, biliary excretion was 19-30%. A lower proportion of radioactivity was found in the urine (13-25%) and excreted primarily as metabolites of hydrolysis or oxidation processes including: 4'-OH BP acid, BP acid, 4'-OH BP alcohol and dimethoxy BP alcohol, TFP acid, cis-OH-methyl TFP acid and trans-OHmethyl TFP acid (<0.01% of the administered dose was unchanged bifenthrin). Approximately half of the radioactivity excreted in the feces was as metabolites and included: hydroxymethylbifenthrin, 4'-OH-bifenthrin, 3'-OH-hydroxymethyl-bifenthrin, 4'-OH-hydroxy-methylbifenthrin, 3'and 4'-monomethyl-catechol-bifenthrin, dimethoxy-bifenthrin and 4'-methoxybifenthrin. Hydrolytic products related to mono- and dihydroxylated bifenthrin were also detected, which included 4'-OH-biphenyl alcohol, dimethoxy-biphenyl acid, dimethoxy-biphenyl alcohol, 4'-methoxy biphenyl alcohol and biphenyl alcohol, TFP acid, and cis and transhydroxymethyl-TFP acid.

Levels of radioactivity in plasma and whole blood were relatively low and elimination from plasma was faster than elimination from whole blood, decreasing to 1/6 of the peak concentration at days 78 and 99, respectively. Peak radioactivity levels in the blood occurred between 4 and 6 hours. Repeated dosing resulted in a 2-fold increase in peak blood radioactivity levels. Unchanged bifenthrin was detected in plasma along with major metabolites, biphenyl acid and biphenyl alcohol. The highest concentrations of radioactivity were found in fat and skin, in particular, the fat of females. Organs with higher fat concentrations had higher peak radioactivity concentrations and longer half-lives (for example: sciatic nerve and ovaries) while other organs had lower peak concentrations and shorter half-lives (for example, liver and kidney).

In vitro study results of radiolabeled bifenthrin in rat and mouse hepatic S9 microsomes indicated that metabolism was not extensive in the cells tested. The identified metabolites 4-OH bifenthrin and BP acid were formed by ring hydroxylation and scission of bifenthrin, respectively. Metabolic activity among species was male mouse > female mouse > male rat. In vitro treatment of rat and human hepatic microsomes indicated differences in P450 isoforms involved in bifenthrin metabolism, and rat microsomes also metabolized bifenthrin faster than human microsomes.

In acute toxicity studies, bifenthrin was highly toxic to rats and mice via the oral route, of low toxicity via the dermal route in rats and rabbits, and slightly toxic via the inhalation route in rats. Bifenthrin was non-irritating to the eyes and skin of rabbits, and was negative in a guinea pig sensitization test by the Buehler method. However, bifenthrin is considered a potential dermal sensitizer based on positive findings in a guinea pig Maximization test.

In the absence of acute toxicity testing with Capture 240 EC, acute toxicity studies with Capture 2EC, a similar end-use product also containing bifenthrin, were considered acceptable surrogates. On the basis of these data, Capture 240 EC is considered highly toxic to rats via the oral route, of low toxicity to rabbits via the dermal route, and of slight toxicity to rats via the

inhalation route. It was minimally irritating to the eyes of rabbits, slightly irritating to rabbit skin, and a potential dermal sensitizer based on a lack of acceptable sensitization test.

Based on dietary (rat, mouse) or capsule (dog) repeat-dose studies conducted by the oral route, the most sensitive indicators of toxicity were signs of neurotoxicity (tremors, clonic convulsions, ataxia, abnormal gait, hypersensitivity to sound), which increased in frequency and intensity with increasing dose. In range-finding studies, mortality was observed at higher doses. In addition, effects on body weight, food consumption, organ weights, clinical chemistry, and haematology were also observed, but were not consistent in magnitude or direction of change among studies. Additional effects following repeat-dosing with bifenthrin included a delay of first estrous in the 1-year dog (capsule) study, retinal atrophy in rats from the chronic/oncogenicity study, increased incidence of submucosal lesions of the bladder at the highest dose tested, and bilateral germinal epithelial degeneration in testis in the mouse oncogenicity study at all dose levels tested. There was no indication that toxicity increased with increasing duration of exposure. No sex-related differences in sensitivity were noted, despite the greater deposition of bifenthrin in the adipose tissue of female rats in toxicokinetic studies, compared to males.

In a 21-day dermal toxicity study in rats, neurotoxicity, evident as paraesthesia, and skin irritation (desquamation, eschar, erythrema, ulceration, hyperplasia) were observed and increased in severity and incidence with dose.

Bifenthrin and several impurities were tested in a battery of in vitro and in vivo genotoxicity studies. Of the 17 studies using bifenthrin and nine studies using bifenthrin impurities, only the mouse lymphoma assay for bifenthrin yielded a positive result. On the basis of the overall findings, bifenthrin and the tested impurities are not considered genotoxic.

There was no evidence of treatment-related tumors in the dietary rat chronic/oncogenicity study. In the dietary mouse oncogenicity study, an increased incidence of lung and liver tumors was observed. In female mice, a statistically significant increase in the combined incidence of bronchioalveolar adenocarcinomas and adenomas was observed in the treated dose groups, compared to control. In comparison to historical controls, the tumor incidence in concurrent controls appeared low, likely contributing to the finding of statistical significance in the treatment groups. This study was considered to provide equivocal evidence for lung tumorigenicity as there was a lack of a dose response; the incidences were within the historical control range for the testing laboratory, the genotoxicity results were negative, and no treatmentrelated tumors were observed in rats. The overall weight of evidence suggested a low level of concern for the lung tumor findings in female mice. In high-dose male mice, a slight increase in the incidence of combined hepatocellular adenomas and adenocarcinomas was observed. Preneoplastic changes that are normally present in chemically-induced tumors were not observed. The incidence of tumors did not reach statistical significance in any treated group; however, a trend test was statistically significant for the incidences of hepatocellular adenocarcinomas and combined hepatocellular adenomas and carcinomas. The combined incidence in the high dose group exceeded the historical control incidence based on a single, relevant historical control reference study.

This study was considered to provide equivocal evidence for liver tumorigenicity, as there were no pre-neoplastic changes, the genotoxicity results were negative, and there was no similar response in female mice or in rats. The overall weight of evidence suggested a low level of concern for the liver tumor findings in male mice.

Bifenthrin did not affect reproductive indices in a two-generation dietary toxicity study in rats. Systemic effects in female parental animals included tremors, clonic convulsions, and decreased body weight and bodyweight gain. A minimal increase in stillbirths observed in the second generation pups believed to be caused by a heating failure in the animal room. As such, this finding was considered equivocal and of low concern. No effects were noted in offspring at the highest dose tested.

In rat dietary and oral gavage developmental toxicity studies, systemic toxicity in dams included tremors, clonic convulsions, splayed hindlimbs, hypersensitivity to sound, piloerection, twitching, and loss of muscle control. There was no evidence of developmental toxicity in the rat dietary study. A 10-fold lower dose in the gavage study produced a treatment-related increased incidence of a variation, hydroureter without hydronephrosis, at a dose that produced tremors in the dam. In an oral gavage rabbit developmental toxicity study, maternal animals showed evidence of neurotoxicity (tremors). No developmental toxicity was evident. Developmental and reproductive toxicity studies did not identify evidence of increased sensitivity of the young.

Several neurotoxicity studies, including gavage delayed neurotoxicity studies in hens and rats, as well as a gavage acute rat neurotoxicity study, a dietary 90-day rat neurotoxicity study, and a dietary rat developmental neurotoxicity (DNT) study, were available.

Bifenthrin did not cause delayed acute neurotoxicity in either the hen or the rat. In hens, oral (gavage) dosing with bifenthrin caused various neurotoxic effects (including unsteadiness, inability to walk/stand, wing-dropping, twitching of head/neck, jerking head movements, trembling), shortly after dosing. These effects reversed four to seven days after cessation of dosing. There were no neuropathology findings. In the rat tilting plane test, oral (gavage) dosing caused a slight increase in the mean angle of slip, stereotyped grooming and greasy appearance of fur, mortality, and decreased bodyweight.

In studies with rats, neurotoxic effects included tremors and twitching, reduced grip strength, increased arousal, increased landing foot splay, increased grooming, and clonic convulsions. In the acute gavage neurotoxicity study, neurotoxic effects occurred rapidly following dosing, with peak effects occurring in about 4-8 hours, and sooner at higher doses. In these studies, clinical signs of neurotoxicity were reversible following cessation of dosing. In the 90-day dietary neurotoxicity study, and in the maternal animals of the dietary DNT study, neurotoxic effects occurred from one week to several weeks after initiation of dosing, depending on dose level. Offspring in the DNT study exhibited increased grooming, reduced ambulatory and total motor activity, and increased acoustic startle at the dose causing maternal tremors and increased grooming. Neuropathology was not observed in toxicity studies except at the highest dose tested in the DNT study, where female offspring had a treatment related increased incidence of minimal axonal degeneration of the lumbar dorsal root fibres. Interpretation of this finding was complicated by the lack of examination of the low- and mid-dose groups; however, the incidence

of this finding was outside historical control range. The results of the DNT study did not suggest increased sensitivity of the young relative to maternal animals.

In general, pyrethroid neurotoxicity is correlated with peak concentrations of the unchanged pyrethroid in blood, with bolus dosing resulting in larger internal doses and greater toxicity, compared to dietary administration. As the design of the DNT study does not consider the time of peak effect and may miss the window of peak toxicity for the pyrethroids, neurobehavioural assessments of the young in the DNT study may not be particularly informative. It is known that the metabolic clearance of pyrethroids in rats increases during maturation, primarily due to increased hepatic enzyme activity. Incomplete maturation of the enzyme systems in the liver which detoxify pyrethroids may result in increased pyrethroid concentrations in target tissues (for example: brain) and increased susceptibility of the young to toxicity, compared to adults receiving the same oral dose. Given the limitations of the DNT study in this regard, an adequate comparison of the sensitivity of the young animal is currently not available. A comparative oral gavage neurotoxicity study conducted in pups, weanling and adults, which considers the time of peak effect, could address this uncertainty. In the interim, this uncertainty has been reflected in the form of a database uncertainty factor.

Results of the toxicity studies conducted with bifenthrin, and its associated end-use product, are summarized in Appendix I, Tables 2 and 3. The toxicology endpoints for use in the human health risk assessment are summarized in Appendix I, Table 4.

Incident Reports

Since 26 April 2007, registrants have been required by law to report incidents to the PMRA, including adverse effects to Canadian health or the environment. Bifenthrin and Capture 240 EC, which contains bifenthrin, were first registered for use in Canada on 23 May 2014. As of 3 May 2017, the PMRA received 1 human and 11 domestic animal incidents involving bifenthrin.

The human incident classified as major occurred in the United States and involved a child that accidentally ingested a small amount of a concentrated American termiticide product containing bifenthrin (7.9%). The child was hospitalized for several days and had symptoms of stomach pain, vomiting, lethargy, headache, hyperthermia, sweating, tingling skin, muscle tremors, unstable gait, nystagmus, and tachycardia. The incident was considered related to the reported pesticide exposure. Bifenthrin is only registered in one Canadian end-use product, Capture 240 EC. The label of the Canadian product indicates that it is a poison, to keep out of reach of children and that it must be stored locked. The label statements on the Canadian product address the circumstances of exposure reported in the United States incident. No additional mitigation measures are necessary.

The eleven domestic animal incident reports involving bifenthrin also occurred in the United States and were classified as animal death. In eight incidents, dogs were exposed as a result of coming in contact with an area treated with bifenthrin and other active ingredients, ingesting a bifenthrin product, or being treated directly with a bifenthrin product. Fish mortality as result of drift from an application site was reported in two incidents. Also, one incident involved a cat that

was potentially exposed after an area inside the home was treated with a bifenthrin product. Although serious and fatal animal outcomes were described in American incidents, the reported exposure scenarios are rare and varied, and no Canadian domestic animal effects have been reported. No additional mitigation measures are necessary.

The incident report data was incorporated into the evaluation of bifenthrin.

3.1.1 Pest Control Products Act Hazard Characterization

For assessing risks from potential residues in food or from products used in or around homes or schools, the *Pest Control Products Act* requires the application of an additional 10-fold factor to threshold effects to take into account the completeness of the data with respect to the exposure of, and toxicity to, infants and children, and potential prenatal and postnatal toxicity. A different factor may be determined to be appropriate on the basis of reliable scientific data.

With respect to the completeness of the toxicity database as it pertains to the toxicity to infants and children, the standard complement of required studies, including developmental toxicity studies in the rabbit and rat, and a reproductive toxicity study in the rat, were available. In addition, a developmental neurotoxicity study in the rat was available.

With respect to potential prenatal and postnatal toxicity, there was no indication of increased susceptibility of fetuses or young animals compared to parental animals in the rat reproductive study, in the rat dietary, rat gavage, and rabbit gavage developmental toxicity studies, or in the DNT study. In the 2-generation reproductive toxicity study, at a dose causing neurotoxicity (tremors, clonic convulsions) and bodyweight effects in dams, an equivocal increase in fetal stillbirths was observed while no effects were observed in offspring. In the rat gavage developmental toxicity study, an increased incidence of a variation, hydroureter without hydronephrosis, occurred at the same dose at which maternal toxicity (tremors) was observed. No developmental effects occurred in a gavage rabbit study at doses causing tremors, convulsions and lack of muscle control in the dams.

Young animals have incomplete maturation of the enzyme systems that detoxify pyrethroids and thus may be more susceptible due to higher and prolonged brain concentrations, compared to adults. Due to the lack of a comparative oral neurotoxicity study, an adequate assessment of sensitivity of the young is currently not available and residual uncertainty remains concerning susceptibility of the young to potential neurotoxic effects. This concern was reflected through the use of a 3-fold database uncertainty factor in the risk assessment. Since these concerns were addressed with a database uncertainty factor, the *Pest Control Products Act* factor was reduced to 1-fold.

3.2 Determination of Acute Reference Dose – all populations

To estimate acute dietary risk, the $BMDL_{20}$ of 2.6 mg/kg bw from an acute oral neurotoxicity study conducted with bifenthrin was selected, based on reduced motor activity in adult rats. Reduced motor activity was considered the critical endpoint since it is a sensitive neurobehavioral endpoint that is relevant to pyrethroid toxicity and is derived by a relevant route and duration of exposure. The $BMDL_{20}$ was specifically selected based on the reported

variability of motor activity in control rats in the literature. The rabbit developmental toxicity study was considered a co-critical study for acute dietary risk. The NOAEL for the dams was 2.67 mg/kg bw/day based on neurotoxic signs (head and forelimb twitching) that were observed within two days of dosing at the LOAEL.

For the reasons outlined in the *Pest Control Products Act* Hazard Characterization Section, a 3-fold UF_{DB} was applied and the *Pest Control Products Act* factor reduced to 1-fold. Standard uncertainty factors of 10-fold for interspecies extrapolation and 10-fold for intraspecies variability were also applied, resulting in a composite assessment factor (CAF) of 300.

The acute reference dose (ARfD) is calculated according to the following formula:

$$ARfD = NOAEL = 2.6 \text{ mg/kg bw} = 0.009 \text{ mg/kg bw of bifenthrin}$$

$$CAF = 300$$

3.3 Determination of Acceptable Daily Intake – all populations

To estimate risk from repeated dietary exposure, a NOAEL of 1.0 mg/kg bw/day was selected based on the results of the following co-critical toxicity studies: the oral NOAEL of 1.5 mg/kg bw/day in the 1-year dog study and the NOAEL of 1.0 mg/kg bw/day in the gavage developmental toxicity study in rats. In the rat developmental toxicity study, tremors were observed in the dams at the LOAEL of 2.0 mg/kg bw/day; a dose causing hydroureter without hydronephrosis in the fetuses. At the LOAEL of 3.0 mg/kg bw/day in the 1-year dog study, tremors and delayed first estrous were observed.

For the reasons outlined in the *Pest Control Products Act* Hazard Characterization Section, a 3-fold UF_{DB} was applied and the *Pest Control Products Act* factor reduced to 1-fold. Standard uncertainty factors of 10-fold for interspecies extrapolation and 10-fold for intraspecies variability were also applied, resulting in a CAF of 300.

The acceptable daily intake (ADI) is calculated according to the following formula:

$$ADI = \frac{NOAEL}{CAF} = \frac{1.0 \text{ mg/kg bw/day}}{300} = 0.003 \text{ mg/kg bw/day of bifenthrin}$$

The ADI provides a margin of 1200 to the NOAEL based on increased grooming counts, reduced motor activity, increased acoustic startle response, and neuropathology in female offspring at the LOAEL in the rat DNT study. Note that the neuropathology was observed in high dose female offspring, however, neither the low- or mid-dose groups were examined for neuropathology. Therefore, the margin was calculated to the low-dose group dose of 3.6 mg/kg bw/day. Additionally, the ADI provides a margin of >2500 to the LOAEL for bilateral germinal epithelial degeneration of the testes in male mice in the mouse oncogenicity study, to the dose showing an equivocal increase in lung tumors in female mice and to the equivocal liver tumors in male mice in the mouse oncogenicity study.

Cancer Assessment

As previously discussed, the lung and liver tumors noted in the mouse oncogenicity study were considered equivocal or not relevant to humans based on the weight of evidence. Overall, the endpoints selected for the non-cancer risk assessment are protective of these findings.

3.4 Occupational and Residential Risk Assessment

3.4.1 Toxicological Endpoints

Occupational exposure to bifenthrin is characterized as short- to intermediate-term in duration and is predominantly by the dermal and inhalation route for mixers, loaders, and applicators and by the dermal route for post-application re-entry workers.

Short- and Intermediate-term Dermal

For short- and intermediate-term occupational exposures via the dermal route, a NOAEL of 50 mg/kg bw/day was selected from the 21-day dermal toxicity study in adult rats. At a dose of 100 mg/kg bw/day, staggered gait, exaggerated hind-limb flexion, and reduced tail flick latency were observed. This study is representative of the route of exposure, and was considered relevant for the short- and intermediate-term scenarios since there was no pronounced evidence of increased toxicity following increased duration of dosing in rats. The target Margin of Exposure (MOE) is 300. This includes uncertainty factors of 10-fold for interspecies extrapolation, 10-fold for intraspecies variability, and a 3-fold database uncertainty factor for concerns related to potential sensitivity of the young.

Since the endpoint selected in the dermal study was systemic neurotoxicity, it was considered protective of the tremors noted in dams in the developmental toxicity study, and the tremors and delayed estrous noted in the 1-year dog study.

Short- and Intermediate-term Inhalation

For short-, and intermediate-term occupational exposure via the inhalation route, a NOAEL of 1.0 mg/kg bw/day was selected from the gavage developmental toxicity study in rats. At a dose of 2.0 mg/kg bw/day maternal tremors were observed. No inhalation toxicity study was available, therefore, an oral endpoint was selected. The selection of this study was considered appropriate as it addresses the most sensitive toxic effect, neurotoxicity, and was considered relevant for the short- and intermediate-term scenarios, since there was no pronounced evidence of increased toxicity following increased duration of dosing in rats. In addition, worker populations could include pregnant or lactating women and, therefore, this endpoint is considered appropriate for the occupational risk assessment.

The target MOE is 300. This includes uncertainty factors of 10-fold for interspecies extrapolation, 10-fold for intraspecies variability, and a 3-fold database uncertainty factor for concerns related to potential sensitivity of the young.

3.4.1.1 Dermal Absorption

3.4.2 Occupational Exposure and Risk

3.4.2.1 Mixer/loader/applicator Exposure and Risk Assessment

Individuals have potential for exposure to bifenthrin during mixing, loading, application, clean up and repair. Dermal and inhalation exposure estimates for workers were generated from the PHED or AHETF databases and Agency default area treated per day values (ATPD).

Exposure to workers mixing, loading, applying, cleaning up and repairing equipment is expected to be short- to intermediate-term in duration and to occur primarily by the dermal and inhalation route. Chemical-specific data for assessing human exposures during pesticide handling activities were not submitted. Exposure estimates were derived for workers applying bifenthrin to raspberries using airblast equipment and handheld sprayers and in-furrow to potatoes using groundboom application equipment. The exposure estimates are based on mixers/loaders/applicators wearing long sleeved shirt and long pants with gloves except for groundboom applicators for which non-gloved data was used.

Dermal exposure was estimated by coupling the unit exposure values with the amount of product handled per day. Inhalation exposure was estimated by coupling the unit exposure values with the amount of product handled per day with 100% inhalation absorption. Exposure was normalized to mg/kg bw/day by using 80 kg adult body weight.

Exposure estimates were compared to the toxicological end points (no observed adverse effects levels) to obtain the MOE; the target MOE is 300 (Appendix I, Tables 5 and 6).

3.4.2.2 Exposure and Risk Assessment for Workers Entering Treated Areas

There is potential for exposure to workers re-entering areas treated with bifenthrin to complete tasks such as establishing irrigation lines, hand harvesting, scouting and tying/training. Given the nature of activities performed, dermal contact will be the primary route of exposure. Inhalation exposure is not expected to be of concern as bifenthrin is considered non-volatile with a vapour pressure of 2.4×10^{-5} Pa which is less than the NAFTA criteria for a non-volatile product for outdoor uses $[1 \times 10^{-4} \text{ kPa} (7.5 \times 10^{-4} \text{ mm Hg}) \text{ at } 20\text{-}30^{\circ}\text{C}]$. The duration of exposure is considered to be short- to intermediate-term.

Exposure to workers re-entering treated potato fields was not calculated as exposure to residues from in-furrow application at planting is expected to be negligible.

Dermal exposure to workers entering treated raspberry fields is estimated by coupling dislodgeable foliar residue values with activity-specific transfer coefficients. Activity transfer coefficients are based on ARTF data. Chemical-specific dislodgeable foliar residue data were not

submitted for raspberries. As such, a default dislodgeable foliar residue value of 25% of the application rate and a 10% dissipation per day were used in the exposure assessment.

Exposure estimates were compared to the toxicological end point to obtain the MOE; the target MOE is 300. Only exposure and risk to the activities with the highest TCs are presented as all activities exceed the target MOE of 300 (Appendix I, Table 7).

3.4.3 Residential Exposure and Risk Assessment

3.4.3.3 Bystander Exposure and Risk

Bystander exposure should be negligible since the potential for drift is expected to be minimal. Application is limited to agricultural crops only when there is low risk of drift to areas of human habitation or activity such as houses, cottages, schools and recreational areas, taking into consideration wind speed, wind direction, temperature inversions, application equipment and sprayer settings.

3.5 Food Residues Exposure Assessment

3.5.1 Exposure from Drinking Water

3.5.1.1 Concentrations in Drinking Water

Estimated environmental concentrations (EECs) of bifenthrin in potential drinking water sources (groundwater and surface water) were generated using computer simulation models. EECs of bifenthrin in groundwater were calculated using the PRZM-GW model to simulate leaching through a layered soil profile over a 50-year period. The concentrations calculated using PRZM-GW are based on the flux, or movement, of pesticide into shallow groundwater with time. EECs of bifenthrin in surface water were calculated using the PRZM/EXAMS models, which simulate pesticide runoff from a treated field into an adjacent water body and the fate of a pesticide within that water body. Pesticide concentrations in surface water were estimated in a small reservoir.

A Level 1 drinking water assessment was conducted using conservative assumptions with respect to environmental fate, application rate and timing, and geographic scenario. Appendix I, Table 8 lists the application information and main environmental fate characteristics used in the simulations. Fifteen initial application dates between February and September were modelled. The model was run for 50 years for all scenarios. The largest EECs of all selected runs are reported in Appendix I, Table 9. In this case, both the modelled EECs and the limit of solubility are reported. For groundwater modelling, the potato rate was modelled. Higher leaching potential would be expected from an in-furrow application at 337 g a.i./ha compared to the foliar method of application for raspberries at a lower rate (2 × 112 g a.i./ha). For surface water modelling, comparative modelling (with in-furrow and t-band applications relevant to the use pattern for potato in-furrow, and foliar applications for raspberries) indicated lower EECs from the potato rate compared to the raspberry rate. Although the potato rate is higher, the application method for potato includes application of the pesticide below the soil surface, which reduces the availability of the pesticide for runoff. Therefore, the more conservative EEC values are reported for raspberries. Details of water modelling inputs and calculations are available upon request.

3.5.2 Residues in Plant and Animal Foodstuffs

The residue definition for risk assessment and enforcement in plant products and animal commodities is bifenthrin. The data gathering/enforcement analytical methods are valid for the quantitation of bifenthrin residues in crop and livestock matrices. The residues of bifenthrin are stable in representative matrices from five crop categories (high water, high oil, high protein, high starch and high acid content) and processed commodities for 6-49 months when stored at -38 to 0°C. Therefore, bifenthrin residues are considered stable in all frozen crop matrices and processed crop fractions for the duration of storage. The raw agricultural commodities (potato; soybean; tomato; pear) were processed and bifenthrin residues only concentrated in the following processed commodities: soybean aspirated grain fractions (AGF) (83-fold); pear wet pomace (peeled) (2.8-fold) and pear wet pomace (ground) (14.6-fold). Adequate feeding studies were carried out to assess the anticipated residues in livestock matrices resulting from the current uses. Crop field trials conducted throughout the United States and India (tea only) using end-use products containing bifenthrin at approved or exaggerated rates in or on the various petitioned commodities are sufficient to support the proposed maximum residue limits (MRLs).

3.5.3 Dietary Risk Assessment

Acute and chronic (cancer and non-cancer) dietary risk assessments were conducted using the Dietary Exposure Evaluation Model (DEEM–FCIDTM, Version 2.16), which uses updated food consumption data from the United States Department of Agriculture's Continuing Surveys of Food Intakes by Individuals, 1994–1996 and 1998.

3.5.3.1 Chronic Dietary Exposure Results and Characterization

The following criteria were applied to the refined chronic cancer and non-cancer analysis for bifenthrin: projected percent crop treated data for potato and raspberry; American percent crop treated data for crops grown in the United States (where available); default and experimental processing factors (where available), residues of bifenthrin in or on carrot; potato; head lettuce; spinach; mustard greens; broccoli; cauliflower; cabbage; soybean; edible-podded pea and bean; succulent shelled pea and bean; tomato; pepper (bell and nonbell); eggplant; cantaloupe; cucumber; summer squash; pear; mayhaw; raspberry; blackberry; celery and tea based on supervised trial median residue (STMdR) values and anticipated residues for all animal commodities. The refined chronic dietary exposure from all supported bifenthrin food uses (alone) for the total population, including infants and children, and all representative population subgroups is less than 16.5% of the ADI. Aggregate exposure from food and drinking water is considered acceptable. The PMRA estimates that chronic dietary exposure to bifenthrin from food and drinking water is 16.7% (0.000500 mg/kg bw/day) of the ADI for the total population. The highest exposure and risk estimate is for children 1-2 years old at 33.9% (0.001017 mg/kg bw/day) of the ADI.

3.5.3.2 Acute Dietary Exposure Results and Characterization

The following assumptions were applied in the refined acute analysis for bifenthrin: United States monitoring data (where available); default and experimental processing factors (where available), residues of bifenthrin in or on carrot; potato; head lettuce; spinach; mustard greens; broccoli; cauliflower; cabbage; soybean; edible-podded pea and bean; succulent shelled pea and bean; tomato; pepper (bell and nonbell); eggplant; cantaloupe; cucumber; summer squash; pear; mayhaw; raspberry; blackberry; celery and tea based on maximum values and anticipated residues for all animal commodities. The refined acute dietary exposure (food alone) for all supported bifenthrin registered commodities is estimated to be 33.8% (0.003045 mg/kg bw/day) of the ARfD for the general population (95th percentile, deterministic). Aggregate exposure from food and drinking water is considered acceptable: 35.4% of the ARfD for the general population.

3.5.4 Aggregate Exposure and Risk

3.5.5 Maximum Residue Limits

For the MRLs for this active ingredient, please refer to the Maximum Residue Limit Database on the Maximum Residue Limits for Pesticides webpage in the Pesticides and Pest Management section of Health Canada's website.

The nature of the residues in animal and plant matrices, analytical methodologies, field trial data, and acute and chronic dietary risk estimates are summarized in Appendix I, Tables 1, 10 and 11.

4.0 Impact on the Environment

4.1 Fate and Behaviour in the Environment

The PMRA requested additional Section 12 data during the conditional registration period which included the following:

- Aerobic soil and water/sediment studies examining the transformation of the isomers for bifenthrin, 4'-OH bifenthrin and TFP acid as requested by the European Commission,
- A biomonitoring program to assess whether risks to organisms remain low under conditions of use under Canadian conditions,
- A study which determines the residual toxicity for non-target arthropods and the potential for re-colonisation,
- Tissue residue-based acute and chronic fish ecotoxicity studies,
- Additional biomagnification and bioconcentration studies for fish and aquatic invertebrates, and
- Terrestrial field dissipation study conducted in an ecoregion equivalent to Canada use sites.

In addition, the bioaccumulation assessment was updated using the most recent OECD guidelines and recommended calculation methods.

Based on submitted information, bifenthrin is classified as practically insoluble in water. The vapour pressure and Henry's law constant suggest that bifenthrin has low potential to volatilize from water and moist soil. An atmospheric model predicted an atmospheric half-life of less than one day. However, there is a high amount of uncertainty around this estimate as a high fraction of bifenthrin is expected to sorb to airborne particles which will decrease the availability of bifenthrin to be degraded by atmospheric oxidation.

Technical bifenthrin primarily consists of the cis-isomer with trace amounts of the trans-isomer. When bifenthrin treated soil or water was irradiated, it underwent isomerization and formed a higher ratio of trans-bifenthrin. The enantiomer ratios of two stereoisomers of cis-bifenthrin (R-cis-bifenthrin and S-cis-bifenthrin), 4'-OH bifenthrin and TFP acid remained relatively the same under laboratory aerobic conditions in absence of light. Therefore, the enantiometric ratios of the R- and S-cis-bifenthrin, 4'-OH bifenthrin and TFP acid are not expected to change significantly in the aerobic environment over time. Slight preferential enantiometric transformation between the R-cis-bifenthrin and S-cis-bifenthrin was observed under laboratory anaerobic soil conditions; however, it is not expected to impact the ratio of these enantiomers significantly in the natural environment as all enantiomers of bifenthrin are very persistent under anaerobic condition.

Bifenthrin has a high soil adsorption coefficient ($K_d = 26,013 - 154,299$), is highly absorptive to soil and exhibits very limited mobility. Bifenthrin is moderately persistent to persistent in aerobic soils with DT_{50} values ranging from 78.7 to 203 days and persistent in anaerobic (flooded) soils with DT_{50} values greater than 1000 days under laboratory conditions. The primary dissipation route of bifenthrin in terrestrial ecosystems is biotransformation; however, the rate varies depending on the climate and soil characteristics. The results of terrestrial field studies conducted in Canadian equivalent ecoregions confirm the laboratory findings. Under field conditions, bifenthrin was moderately persistent to persistent with DT_{50} values ranging from 80 to 215 days with 11-27% of residues carried over to the next growing season. Bifenthrin was mainly detected in the 0-30 cm soil layer without any evidence of movement below 30 cm soil depth in field studies conducted in United States, Germany, Italy and France. Bifenthrin is not expected to leach to ground water.

No major transformation products were detected in the terrestrial laboratory and field studies. The minor transformation products identified were 4'-OH bifenthrin, TFP acid, biphenyl acid, biphenyl alcohol and biphenyl aldehyde. The mobility of TFP acid and 4'-OH bifenthrin was tested in a variety of soils. TFP acid is classified as either moderately mobile or very highly mobile depending on soil. The mobility of TFP acid is strongly dependent on the pH and the organic carbon content of the soil. It is less mobile in soils with low pH and high organic carbon content. 4'-OH bifenthrin is less mobile than TFP acid and is classified as either low or immobile depending on the soils. The transformation products are not expected to form in appreciable amount in the natural environment and are unlikely to leach to ground water in a significant amount on a seasonal basis.

Bifenthrin can enter the aquatic environment through spray drift and runoff from the application site. Bifenthrin is highly insoluble in water and hydrolysis is not an important route of transformation. Photolysis and biotransformation can be important processes of degradation.

In the laboratory photolysis study, bifenthrin underwent isomerization from cis-bifenthrin to trans-bifenthrin and transformed to bifenthrin alcohol and TFP acid that reached maximums of 19% and 10.2% of applied, respectively. In aerobic water-sediment biotransformation studies, bifenthrin partitioned from the water into sediment after a few days and remained moderately persistent to persistent with whole system DT_{50} 's ranging from 92.9 days to 276 days. 4'OH-bifenthrin was the only major transformation product detected in the sediment of the aerobic water/sediment system. Minor transformation products detected were TFP acid, biphenyl acid and biphenyl alcohol. These minor transformation products are not expected to be formed in high quantities in the environment. Under anaerobic conditions, bifenthrin is expected to be persistent based on the results from the flooded soil study ($DT_{50}>1000$ days).

Under aquatic field conditions, bifenthrin is expected to quickly partition from the water to the sediment. Microbial biotransformation is expected to be the most important route of transformation in aquatic systems. In an aquatic field study conducted in Alabama, bifenthrin was much more persistent than predicted by the laboratory studies. The estimated DT₅₀ of bifenthrin residues in pond water of the aquatic field study was 609 days. Bifenthrin residues in the pond sediment declined slowly until study termination, two years after the last application. Although half-lives could not be estimated, the mean concentration in sediment samples collected 737 days after the final application were approximately 21% of the highest mean observed on day 57 after the last treatment. This was possibly due to its persistence in soil and the continuous input into the pond from runoff for the two years following the last application.

The majority of the environmental fate endpoints from the open literature, while highly variable, confirm the persistence and immobility characteristics of bifenthrin. Some reported half-lives were less than one year and were comparable to those obtained from the submitted environmental fate studies. A few reported half-lives were greater than one year and were significantly greater than those obtained from the submitted studies.

A review of the published literature indicated the presence of bifenthrin in freshwater sediment, urban estuarine sediment, irrigation and storm runoff in the United States. While the available data from the United States Geological Survey's National Water-Quality Assessment Program, the United States Environmental Protection Agency's (UESPA) Storage and Retrieval Data Warehouse, and the California Department of Pesticide Regulation reported the low frequency of bifenthrin detections, several recent articles reported bifenthrin to be one of the most frequently detected pyrethroids in water samples collected from runoff and urban or agricultural creeks. Bifenthrin is also a common contaminant in sediment samples at concentrations up to 430 μ g/kg sediment.

Based on a log $K_{\rm ow}$ greater than six, bifenthrin has a high potential to bioaccumulate in aquatic organisms. At the time of original registration, the PMRA determined that the measured bioconcentration and bioaccumulation factors were highly variable and were both above and below bioaccumulation cut-off values for TSMP. A review of bioaccumulation of bifenthrin in both the terrestrial and aquatic environment reported in the European Commission's Additional Report by France (2010) which included a fish dietary biomagnification study, field data and modelling information, indicated that bifenthrin is found in each level of the food chain, but biomagnification in upper trophic organisms was not observed. It also found that despite the

availability of several studies, the bioaccumulation of bifenthrin is not fully characterised as the phenomenon appears to depend on species, life stage and exposure. PMRA requested studies reported in foreign reviews that were not originally submitted to the PMRA for review. PMRA reviewed these studies and integrated this information into the assessment. When all relevant information was integrated together, the weight of evidence indicated that the TSMP criteria were indeed met. In particular, it was concluded that the field bioaccumulation factor (BAF) values were a better indicator of bioaccumulation than the single dietary laboratory study conducted on one species. The BAFs from the field study considered multiple species and pathways of exposure under environmentally relevant conditions.

Calculations for bioconcentration and bioaccumulation factors were updated using the most recent OECD recommended calculation methods correcting for lipid content and growth. Results of the laboratory bioaccumulation studies showed that bifenthrin bioconcentrates significantly in some species. Bioconcentration factor (BCF) values in carp (Cyprinus carpio) (kinetic BCF) corrected for growth rate and lipid content: 1265 – 1861) are low compared to those for bluegill sunfish (kinetic BCF: 3400-12850) and adult fathead minnow (Pimephales promelas) (steady state BCF: 21000-30000). The kinetic BCF for *Daphnia magna* corrected for growth rate is 6273. Bifenthrin did not biomagnify in bluegill sunfish, the biomagnification factor (BMF) was <1.0; however, assimilation efficiency (α) of bifenthrin in bluegill sunfish was very low (3.9% to 5.8%). In comparison, PCBs that are known to have BMFs of \geq 1 have an α of 40-60%. Low assimilation efficiency can be a result of some limitation to the uptake via the gut due to steric effects or bound residues that cannot partition off the food or a combination of these factors. Under field conditions which consider all exposure pathways, estimated bioaccumulation factors (BAFs) indicate significant bioaccumulation. Results of the aquatic field study conducted in Alabama suggested that BAFs >5000 were sustained in several fish species longer than 150 days after the last application. Although there are uncertainties and variability (spatial and temporal) associated with field studies, the field BAF values are considered to offer a reasonable characterization of the exposure history of fish to bifenthrin and were found to be consistent with BCF values obtained under controlled laboratory conditions.

A field monitoring study of dolphins from the Brazilian coast detected bifenthrin residues in liver, breast milk and placental samples (Alonsa, et al. 2012). The results of this study provide evidence of exposure and accumulation of bifenthrin in marine mammals and maternal transfer by both gestational and lactation pathways in non-agricultural areas far from the source of release. Marine mammals are at the top of the marine food chain. Occurance of bifenthrin in their liver samples provides further evidence of bioaccumulation of persistent chemicals in upper trophic level of food chain.

The requested biomonitoring program under Canadian conditions of use was not submitted. A biomonitoring study provided additional residue information on bifenthrin in earthworms and terrestrial small mammals and aquatic biota in a treated field and adjacent ponds at two different locations in Germany at three different time periods. The treatment rate of these studies was 1/10th of the Canadian rate (Canadian rate: 112 g a.i./ha). The results of these European studies are considered to be of limited value in terms of assessing bioaccumulation in aquatic biota because in most cases, residues in water and sediment were undetectable, very close to or below the limit of quantitation (LOQ) and could not confirm exposure which precluded calculating a

bioaccumulation ratio under field conditions. The lack of detections in aquatic environment under the European conditions cannot be interpreted as a lack of exposure under Canadian use conditions. Bifenthrin residues were detected in the earthworm and small rodent samples collected from treated fields. Bifenthrin residues in earthworms were highly variable. In general, the shallow burrowing earthworm had lower mean bifenthrin residues than *L. terrestris*, deep burrowing earthworms. The mean bifenthrin residue in *L. terrestris* peaked at 0.32 mg a.i./kg between 7 to 21 days before declining and remained detectable 119 days after the last application. Bifenthrin residues in small mammal samples were low and highly variable. The majority of samples were under the level of detection or quantification. The highest mean bifenthrin residue in small mammals was 0.0142 mg a.i./kg from Day 6 after the last treatment in Autumn 2014. Specimens containing detectable bifenthrin residues were captured in the treated field but not from the edge of the field and surrounding area. The bifenthrin residues were lower in the small mammal gut samples and gut contents than in the small mammal samples without gut or gut content. This indicated that small mammals had assimilated and accumulated bifenthrin and retained the residues in the body.

A summary of environmental fate data is presented in Appendix I, Tables 12, 13, and 14.

4.2 Environmental Risk Characterization

The environmental risk assessment integrates the environmental exposure and ecotoxicology information to estimate the potential for adverse effects on non-target species. This integration is achieved by comparing exposure concentrations with concentrations at which adverse effects occur. Estimated environmental exposure concentrations (EECs) are concentrations of pesticide in various environmental media, such as food, water, soil and air. The EECs are estimated using standard models that take into consideration the application rate(s), chemical properties and environmental fate properties, including the dissipation of the pesticide between applications. Ecotoxicology information includes acute and chronic toxicity data for various organisms or groups of organisms from both terrestrial and aquatic habitats including invertebrates, vertebrates, and plants. Toxicity endpoints used in risk assessments may be adjusted to account for potential differences in species sensitivity as well as varying protection goals (i.e., protection at the community, population, or individual level).

Initially, a screening level risk assessment is performed to identify pesticides and/or specific uses that do not pose a risk to non-target organisms, and to identify those groups of organisms for which there may be a potential risk. The screening level risk assessment uses simple methods, conservative exposure scenarios (for example, direct application at a maximum cumulative application rate) and sensitive toxicity endpoints. A risk quotient (RQ) is calculated by dividing the exposure estimate by an appropriate toxicity value (RQ = exposure/toxicity), and the risk quotient is then compared to the level of concern (LOC = 1). If the screening level risk quotient is below the level of concern, the risk is considered negligible and no further risk characterization is necessary. If the screening level risk quotient is equal to or greater than the level of concern, then a refined risk assessment is performed to further characterize the risk. A refined assessment takes into consideration more realistic exposure scenarios (such as drift to non-target habitats) and might consider different toxicity endpoints. Refinements may include further characterization of risk based on exposure modelling, monitoring data, results from field

or mesocosm studies, and probabilistic risk assessment methods. Refinements to the risk assessment may continue until the risk is adequately characterized or no further refinements are possible.

There are many published articles examining the toxicity of bifenthrin to pollinators, non-target arthropods and aquatic organisms. The results of up-to-date published journal articles confirm the overall conclusion of bifenthrin's toxicity to pollinators, non-target arthropods and aquatic organisms. The reported endpoints in the open literature do not change the outcome of the risk assessment; therefore, the risk assessment is primarily based on the submitted data.

4.2.1 Risks to Terrestrial Organisms

A risk assessment of bifenthrin to terrestrial organisms was based upon evaluation of toxicity data for earthworms (acute and chronic exposure), bees (acute oral and contact exposure and field aged residue), non-target beneficial arthropods (acute contact laboratory studies, aged residue and extended laboratory study), birds (acute oral, dietary and reproduction), mammals (acute oral and reproduction) and terrestrial plants (effects on seedling emergence).

A summary of terrestrial toxicity data for bifenthrin, formulated products and its transformation products is presented in Appendix I, Table 15. Terrestrial toxicity data for Talstar 8 SC, an enduse product containing nominal 80 g a.i./L, were used as surrogate data when toxicity data for bifenthrin or Capture 240 EC were not available. The screening and refined risk assessment for terrestrial organisms other than birds and mammals are presented in Appendix I, Tables 16 and Table 17, respectively.

No major transformation product was identified or detected in the terrestrial field dissipation studies or laboratory soil studies. Therefore, bifenthrin is expected to be the only residue of concern in the terrestrial habitat.

Earthworms: The screening level risk assessment of bifenthrin indicates that the LOC for earthworms was not exceeded for either acute or chronic exposure. As the LOC was not exceeded for bifenthrin, RQ was not calculated for its transformation product, 4'OH-bifenthrin, which is less toxic and has a lower EEC than bifenthrin.

Bees (pollinators): Pollinators can be exposed to bifenthrin from contact and/or feeding on contaminated parts of plants, for example, pollen and nectar. In-hive bees, including immature bees, can be exposed via contaminated plant materials brought back by foraging bees. The screening level risk assessment of bifenthrin concluded that the LOC for bees was exceeded (RQ = 2.1 for acute oral and 99 for acute contact). According to the results of a field-aged residue study, residues on plants after a foliar spray could remain at toxic levels for several days. Potential exposure and risk to bees will be further mitigated through label statements which prohibit application during the crop blooming period and require off-field drift to be minimized.

Beneficial arthropods: The toxicity of bifenthrin was determined for five different beneficial arthropod species: ladybird beetle (*Coccinella septempunctata* L.), parasitic wasps (*Aphidius rhopalosiphi*), ground beetles (*Poecilus cupreus* L.), green lacewing (*Chrysoperia carnea* Steph.) and predatory mites (*Typhlodromus pyri*). The risk to non-target arthropods was assessed using

maximum cumulative in-field and off-field EECs on plant surfaces, calculated from a direct spray on a field. The acute contact test exposing adult *A. rhopalosiphi* to glass plates or maize leaves treated with 7.5 g a.i./ha of Talstar 8 SC had 100% mortality. Therefore, the screening level RQs from in-field and off-field exposures to bifenthrin exceeded the LOC for uses on raspberry as the treatment rate was significantly lower than the single application rate for raspberry. The results of extended laboratory tests on natural substrate were used for all five species for screening risk assessment purposes. The in-field RQs were 1500, 15, 25 and 1115 for *C. septempunctata* L. (ladybird beetle), *A. rhopalosiphi* (aphid parasitoid), *C. carnea* (green lacewing) and *T. pyri* (predatory mite), respectively. The corresponding off-field RQs were 885, 9, 14 and 658, respectively. The RQs of these four species exceeded the LOC. The RQ for *P. cupreus* L was 0.84 and did not exceed the LOC.

The bifenthrin EEC values for beneficial predatory and parasitic arthropods were refined to consider foliar interception. The exposure estimates are assuming deposition to a 2-dimensional structure. Therefore, the values can be corrected to take into account the 3-dimensional structure where a certain fraction is intercepted by the crop (for in-field exposure) or the off-field vegetation (for off-field exposure). For the in-field EEC, crop-specific foliar interception factors are applied to the application rate. No factors are proposed for raspberries. As raspberry plants at fruiting stage have similar foliage coverage as vines during flowing and fruiting stage, the deposition fractions for vines (Fint = 0.8 and Fsoil = 0.2) are used as surrogates. For the off-field EEC, a vegetation distribution factor of 0.1 is applied to the application drift rate.

The refined in-field RQs were 1202, 12, 20 and 894 for *C. septempunctat* L., *A. rhopalosiphi*, *C. carnea* and *T. pyri*, respectively, and still exceeded the LOC. The refined off-field RQs determined based on the uses on raspberries were 88.5, 1.4 and 7.43 for *C. septempunctat* L., *C. carnea* and *T. pyri*, respectively and exceeded the LOC. As the use of bifenthrin on raspberries is expected to pose potential risks to non-target arthropods including those used in Integrated Pest Management, precautionary statements and directions to minimize off-field drift are required on the label.

Terrestrial plants: The toxicity of Talstar 8 SC to six non-target terrestrial plant species was determined through a seedling emergence test. No treatment-related adverse effects (i.e., >25% effect) were observed in any plant species up to 0.08 mg a.i./kg dw soil, which was the highest test concentration. Therefore, the NOEC and EC₂₅ were 0.08 mg a.i./kg dw soil and > 0.08 mg a.i./kg dw soil, respectively. The soil EEC values based on the highest application rates for potato and raspberries were 0.150 mg a.i./kg dw soil and 0.094 mg a.i./kg dw soil, respectively. The RQ values based on the no-effect endpoints for use on potato and raspberries were <1.9 and <1.2, respectively. Further testing was not completed to determine the level at which effects would be observed as such the true no-effects level may be higher than 0.08 mg ai/kg dw soil. Bifenthrin is a pyrethroid insecticide and the mode of action is not expected to pose adverse effect on plants. Therefore, the use of bifenthrin is not expected to pose a risk to terrestrial plants and mitigation measures are not required.

Terrestrial vertebrates: Birds and mammals may be exposed to bifenthrin following the ingestion of plant materials and insects sprayed with bifenthrin during foliar application. The screening level risk assessment for Capture 240 EC is conducted for direct on-field exposure,

assuming exposure occurs entirely through the consumption of food sources contaminated with bifenthrin at the maximum nomogram residue levels, the most conservative scenario. Concentrations of bifenthrin on different food guilds are calculated based on the highest rate for foliar application (i.e., 2×112 g a.i./ha) with a 30-day interval and a foliar half-life of 10 days.

<u>Birds</u>: Bifenthrin is slightly toxic to Northern bobwhite (*Colinus virginianus*) and is practically non-toxic to mallard duck (*Anas platyrhynchos*). Reproduction studies did not detect any effects or dose response relationship in any measured reproductive parameter, including number of eggs laid, egg weights, egg shell thickness, number of infertile eggs, early and late embryonic death, hatching, chick health, chick body weight and number of 14-day survivors, up to the highest treatment level of 75 mg a.i./kg dw diet. The risk quotients (RQs<1) for acute and reproductive exposure to birds at the screening level risk assessment do not exceed the LOC for small, medium or large birds (Appendix I, Table7). Therefore, the use of bifenthrin on raspberries is not expected to pose a risk to birds.

<u>Mammals</u>: The laboratory toxicity of bifenthrin to mice and rats was used to assess risk to small terrestrial mammals. The results of the acute toxicity test suggest that bifenthrin is highly toxic to mice and is not gender specific (Appendix I, Table 15). In the rat reproduction study, reduced body weight gain in parent and offspring and increased litter incidence of stillborn pups were observed. The risk assessment for mammals are conducted using these two endpoints (acute LD_{50} : 43 mg a.i./kg bw/day and NOAEL for reproduction: 3 mg a.i./kg bw/day) and are presented in Appendix I, Table 18 and Table 19, respectively.

The screening RQs for acute exposures to medium- and large-sized mammals exceeded the LOC but not for small-sized mammals. The screening RQs for reproductive exposures to all three sizes of mammals exceeded the LOC. Therefore, further mammalian risk characterization was conducted. To further characterize the risk, the assessment was expanded to all relevant food guilds, with concentrations of bifenthrin on food items to cover both maximum and mean residue values from the nomogram. Both in- and off-field exposure estimates are considered in this assessment. When considering the maximum residues, the acute on-field RQs (1.32-2.66) exceeded the LOC for medium herbivores and large herbivores, except large herbivores in the long grass food guild. The acute off-field RQs exceeded the LOC for medium herbivores feeding on short grass and forage crops. The reproductive risk from on-field exposure exceeded the LOC for small insectivores, medium herbivores and large herbivores. The reproductive risk from offfield exposure exceeded the LOC for medium and large herbivores, except large herbivores feeding on long grass. When considering the mean residues, only the on-field RQs for medium sized herbivorous mammals feeding on short grass and forage crops exceeded the LOC slightly. The reproductive risk from off-field exposure did not exceed the LOC for any size of mammals. Although the LOC was exceeded for medium sized mammals for on-field exposures, the identified risk is a result of mammals feeding solely on contaminated food, a scenario that would not normally occur. Consequently, the on-field risk to mammals is unlikely to be realized.

As bifenthrin is highly toxic to mammals, a statement informing users of this is required on the label for the end-use product, Capture 240 EC.

4.2.2 Risks to Aquatic Organisms

A summary of aquatic toxicity data for bifenthrin, its formulated products and transformation products is presented in Appendix I, Table 20.

4.2.2.1 Screening Level Assessment

To assess the potential for adverse effects, conservative screening level EECs in the aquatic environment based on a direct application to water were used as the exposure estimates. A risk assessment of bifenthrin was undertaken for freshwater and marine aquatic organisms based on available toxicity data to invertebrates (acute and chronic), fish (acute and chronic), amphibians (using fish as surrogate data), and algae (acute). Aquatic toxicity data for other formulations were used as surrogate data when data for neither bifenthrin nor Capture 240 EC were submitted.

For acute toxicity studies, uncertainty factors of 1/2 and 1/10 the EC₅₀ (LC₅₀) are typically used in modifying the toxicity values for aquatic plants and invertebrates, and fish species, respectively when calculating RQs. No uncertainty factors were applied to chronic NOEC endpoints. The calculated RQs for bifenthrin are summarized in Appendix I, Table 21 (screening level). A risk assessment for 4'OH-bifenthrin was not conducted for the following reasons: it was less toxic than bifenthrin for the species tested and it was not expected to form at concentrations higher than bifenthrin.

Species sensitivity distribution (SSD) analysis was conducted when sufficient acute toxicity data were available to determine hazardous concentration to five percent of species (HC₅) using the software program ETX 2.1. The HC₅ is the concentration which is theoretically protective for 95% of species. At the HC₅ exposure level, five percent of all species may be exposed to a concentration which exceeds the toxicity value. The variability around the fraction of species affected (FA value) is indicated by the lower and upper confidence limits (90% CI), which indicates the minimum and maximum percent of species that may be affected at the HC₅ value.

Freshwater invertebrates: Bifenthrin is toxic to aquatic invertebrates such as *Daphnia magna* and *Chironomus riparius*. Talstar 80g/L Flowable was less acutely toxic to *D. magna* than the bifenthrin technical, but is still toxic to freshwater invertebrates. 4'-OH bifenthrin was much less acutely toxic than the parent to *C. riparius*. A total of 10 acute toxicity endpoints for freshwater invertebrate species compiled from open literature search and applicant data are available for SSD analysis. The median HC₅ value for bifenthrin for acute effects on freshwater invertebrates was determined to be 0.009 μg a.i./L (90% CI: 0.0006 to 0.044 μg a.i./L). The fraction of species affected (expressed as a percentage of all species) at the HC₅ value ranges from 0.61 to 20.1% (90% CI).

Chronic exposure to bifenthrin resulted in reduced reproduction and growth of adult daphnids and declined emergence rate of the fresh water midge *C. riparius*. As the aquatic risk assessment is mainly based on EECs in surface water, the *C. riparius* toxicity endpoint determined from the water-spiked toxicity test was used for the chronic aquatic invertebrate risk assessment.

The screening level RQs for acute and chronic exposure of freshwater invertebrate to bifenthrin exceeded the LOC. Therefore, further refinement of the aquatic risk assessment was required.

Freshwater fish: Bifenthrin is very highly toxic to rainbow trout, bluegill sunfish, fathead minnow, medaka, common carp, and zebra fish. A total of six acute toxicity endpoints for freshwater fish species based on submitted studies were available for SSD analysis. The median HC_5 value of bifenthrin for acute effects on freshwater fish was determined to be 0.008 μ g a.i./L (90% CI: 0.009 to 0.02 μ g a.i./L). The fraction of species affected (expressed as a percentage of all species) at the HC_5 value ranges from 0.25 to 27.7% (90% CI).

For the chronic assessment on fish, fry survival was the most sensitive endpoint for the full life cycle test conducted with fathead minnow.

The screening level RQs for acute and chronic exposure of freshwater fish to bifenthrin exceeded the LOC. Therefore, further refinement of the aquatic risk assessment was required

Freshwater amphibians: In absence of any actual amphibian data, acute freshwater fish HC₅ and fathead minnow full life cycle NOEC were used as surrogates for acute and chronic amphibian endpoints, respectively. The risk for amphibians was characterized at the screening level by comparing EECs in 15 cm water depth with fish toxicity endpoints as surrogates for aquatic life-stages of amphibians. The screening level RQs exceeded the LOC.

Freshwater algae: A 72-hour limit test conducted with green alga *Desmodesmus subspicatus* reported that bifenthrin did not exert any toxic effect on *D. subspicatus* at 6.3 mg a.i./L. The screening level RQ did not exceed the LOC.

Marine/estuarine species: In laboratory studies, bifenthrin was acutely toxic to mysid shrimp (Mysidopsis bahia), eastern oyster (Crassostrea virginica) and sheepshead minnow (Cyprinodon variegatus). Chronic exposure to bifenthrin for 28 days resulted in reduced survival, growth and reproduction of mysid shrimp. The screening RQs exceeded the LOC for mysid shrimp and sheepshead minnows; therefore a more refined assessment is needed.

The endpoints for acute embryo survival and acute shell deposition of eastern oyster differed by two orders of magnitude. Therefore, the screening RQ for acute shell deposition exceeded the LOC while the screening RQ for acute embryo did not. As the EC_{50} for acute shell deposition endpoint was greater than the highest treatment level, the screening RQ may be overestimated.

4.2.2.2 Refined Risk Assessment

Aquatic field studies

Higher tier studies confirm that the most sensitive aquatic organisms are invertebrates. Mesocosm systems show that exposure to bifenthrin decreases species diversity and abundance of the invertebrate community – pelagic zooplankton and benthic invertebrate populations (for example, calanoids, chironomid, oligochaeta, chaoboridae), at environmentally relevant concentrations. Phytoplankton and macrophyte communities showed no direct effects from bifenthrin exposure; in some cases transient effects (increase or decrease in adundance of phytoplankton) were observed, however, these effects were not considered to be treatment related. The NOEC values based on decreased abundance of zooplankton and macroinvertebrate communities were 0.001 µg a.i./L and 0.005 µg a.i./L, respectively. Although effects were

observed to the zooplankton and macroinvertebrate communities at concentrations as low as 0.005 and 0.015 ug/L (LOEC), the taxonomic diversity of the community remained unchanged at 0.015 ug a.i./L. In addition, the invertebrate populations adversely affected at the 0.015 ug a.i/L treatment concentration level were shown to recover within seven days (i.e., rotatoria and some crustacea – for example, copepods) and 14 days (i.e., chaoboridae). Based on the short recovery period observed, consideration of the NOEC of 0.015 ug a.i./L (taxonomic richness) for the risk assessment would be expected to be sufficiently protective of aquatic ecosystems.

Similar effects to the aquatic invertebrate community were observed in the Alabama aquatic field study. In this study, biomonitoring was conducted in a pond for up to two years after bifenthrin was applied aerially to adjacent cotton fields. There were clear effects on various biota in the pond due to bifenthrin treatments:

- 1. Immediate and prolonged elimination (for at least one year after treatment) of calanoid copepods.
- 2. Immediate and prolonged extremely low abundance (for at least one year after treatment) of the mayfly, *Caenis* sp.
- 3. Disappearance of the damselfly *Enallagma* after treatment for at least one year
- 4. Severe reduction of chironomid populations.
- 5. Reduction in survival and reproductive potential of *Daphnia* and snails.

Effects observed in this pond study were shown to persist for a longer period of time than was observed in the mesocosm study. The major limiting factor of the pond is the lack of multiple exposure concentrations where an NOEC (community, abundance etc.) could be determined. Although the pond study clearly shows immediate and prolonged effects to specific aquatic species, it is not amenable for quantitative use in the risk assessment because an NOAEC cannot be determined.

The multiple dose design of the mesocosm study does allow for determination of an NOAEC endpoint for quantitative use in the risk assessment; however, a number of deficiencies and uncertainties limit its accuracy, therefore a safety factor is required to ensure it is protective. These deficiencies are:

- The exposure concentrations used over the test period are uncertain. Four of the seven nominal test concentrations were below the analytical detection limit. In addition, measured concentrations of bifenthrin in water decreased to below the limit of quantitation between days 3 and 7 after treatment in all treatments. Therefore, the reported NOEC, based on nominal test concentrations, likely represents an underestimate of the true NOEC.
- Effects observed in the multi-year field pond study were observed at lower measured concentrations than those observed in the mesocosm study based on nominal test concentrations. The NOEC from the mesocosm study is based on the nominal test concentration instead of the measured concentration. Therefore, the NOEC likely represents an underestimate of effects.

Based on the deficiencies above, the PMRA has determined that a safety factor of three be applied to the NOEC of $0.015~\mu g$ a.i./L so that the significance of these uncertainties is reduced and the chosen NOEC is protective of the community structure and individual aquatic organisms. Note that this safety factor was also used by the European Food Safety Authority (EFSA).

Spray drift and runoff refinements

Given the conservative assumption made in the screening level assessment (i.e., direct over spray to a water body), refined assessments were conducted to better characterize the risk to aquatic organisms. For groups where the screening LOC is exceeded (i.e., $RQ \ge 1$), a refined assessment was conducted to determine if the risks are acceptable from spray drift and runoff separately. In addition to spray drift and runoff refinement, the results of higher tier studies such mesocosm and field studies were considered and compared in refined risk assessment.

Spray drift refinement: The risk to aquatic organisms was further characterized by taking into consideration the concentrations of bifenthrin that could be deposited in off-field aquatic habitats that are downwind and directly adjacent to the treated field through spray drift. The maximum amount of spray that is expected to drift one metre downwind from the application site during spraying using late growth stage airblast application is 59% of the application rate. The estimated EEC values were 16 µg a.i./L and 85 µg a.i./L for permanent water bodies (80 cm deep) and non-permanent/shallow water bodies (15 cm deep), respectively. The risks to aquatic organisms resulting from spray drift are summarized in Appendix I, Table 22. The risk quotients show that the LOC is exceeded for all organisms on an acute exposure basis and chronic exposure basis. For acute exposure, the refined RQs for freshwater invertebrate and fish were 1778 and 205, respectively when considering the acute SSD endpoint. The refined RQ for amphibians was 1090. The refined RQs for the most sensitive marine invertebrate endpoint and marine fish are 8060 and 9, respectively.

For the chronic exposure, the refined RQ for freshwater invertebrates determined using the freshwater mesocosm endpoint was 3000. The refined RQ for freshwater fish based on the chronic fathead minnow endpoint was 400. The refined RQ for amphibian was 2125. The refined RQ for marine invertebrates was 13,333. As the majority of bifenthrin partitions to the sediment, the chronic exposure scenario can be viewed as conservative to pelagic organisms.

For airblast sprayers, calculated buffer zones ranged from 50 to 75 metres and did not require additional spray mitiation measures.

However, initial spray buffer zones for field sprayers calculated based on fine ASAE spray quality were large and did not fully mitigate the risk to aquatic organisms. Therefore spray buffer zones were refined by setting restrictions on various spray application parameters: a minimum ASAE medium spray quality, an 8 km/h wind speed restriction and the requirement to use drift-reducing air induction nozzles. These restrictions are required on the product label. Calculated buffer zones for field sprayer adjusted according to windspeed and low drift nozzle modifiers ranged from 10 to 55 metres.

Runoff refinement: Exposure to runoff into a body of water directly adjacent to the application field was determined using the runoff 90^{th} percentile of the EECs predicted by PRZM-EXAMS for an appropriate time-frame. The EECs were calculated from the model output from each run as follows. For each year of the simulation, PRZM/EXAMS calculates peak (or daily maximum) and time-averaged concentrations. The time-averaged concentrations were calculated by averaging the daily concentrations over five time periods (96-hour, 21-day, 60-day, 90-day, and one year). The 90^{th} percentiles over each averaging period are reported as the EECs for that period. The Level 1 EECs for bifenthrin in water bodies with 15 cm and 80 cm depth were predicted by PRZM-EXAMS for five standard regional scenarios. The EECs used for calculation of the RQs were the highest values at the appropriate depth and appropriate time-frame. The highest detection (5.2 μ g/L) in water monitoring data is within the range of the peak concentration predicted by modeling; consequently, this value was used along with the modelling numbers in the acute assessment for aquatic organisms (both 15 cm and 80 cm depths).

The RQs for exposure to bifenthrin through runoff are provided in Appendix I, Table 23. The RQs exceeded the LOC for majority of freshwater species (freshwater RQ = 5-2,500, marine RQ = 207-208), except *C. riparius*, Eastern oyster or sheepshead minnow. Therefore, freshwater invertebrates, fish, amphibians, and marine invertebrates in aquatic habitats directly adjacent to the application field are potentially at risk from exposure to runoff from a treated field. To reduce the risks of bifenthrin being carried in runoff to aquatic environment, in addition to precautionary label statements, a label statement for the construction and maintenance of a 10 metre vegetative filter strip between the area of application and waterbodies is required.

Other risks to aquatic organisms: Biological effects from pesticides can be a result of immediate direct toxic effects and also from effects that may be asserted after bioaccumulation; therefore, accumulated bifenthrin could potentially impact fitness and survival in biota. In the Alabama pond study discussed in the environmental fate section, bifenthrin residues were detected in different fish species and mussels more than one year after the last application to the adjacent field. A significant die-off of gizzard shads (approximately 1600 fish) was noted between mid-November 1986 and early March 1987. Gizzard shad samples collected during this period had bifenthrin residues ranging from 30 to 487 µg a.i./kg fish (average 312 µg a.i./kg fish). The mean bifenthrin residues of rainbow trout and bluegill sunfish sampled from highest treatment level with zero mortality were 137 µg a.i./kg and 196 µg a.i./kg, respectively. The mean bifenthrin residues of fish sampled from the treatment levels with 80 to 100% mortality were 546 µg a.i./kg and 625 µg a.i./kg, respectively. Bifenthrin residues in the field samples were comparible to residues observed in fish sampled from high mortality groups.

4.2.3 Incident reports

Environmental incident reports are obtained from two main sources, the Canadian pesticide incident reporting system (including both mandatory reporting from the registrant and voluntary reporting from the public and other government departments) and the United States Environmental Protection Agency (USEPA) Ecological Incident Information System (EIIS).

Specific information regarding the mandatory reporting system regulations that came into force 26 April 2007 under the *Pest Control Products Act* can be found at https://www.canada.ca/en/health-canada/services/consumer-product-safety/pesticides-pest-management/public/protecting-your-health-environment/report-pesticide-incident.html.

There are no environmental incidents for bifenthrin in the PMRA database. As bifenthrin is registered in the United States, a review of the latest version of EIIS was completed. As of 5 October 2015, the EIIS database contained 38 incidents involving bifenthrin reported between 1992 and 2015. Six of these incidents are categorized as "unlikely" associated with exposure to bifenthrin. Of the remaining incidents, reports for various organisms including fish, plants, pollinators and birds were located in the USEPA EIIS database.

There were five incidents associated with bifenthrin involving fish mortality as a result of runoff. Three of these were classified as probable and occurred between 1994 and 1996. In 2005 and 2006 there were two fish mortality incidents where it was considered highly probable that the observed effects were a result of runoff of bifenthrin. A dead bird was also noted in the 2006 incident and was considered to be a result of exposure to granular formulated bifenthrin product for use on turf and residential sites. No incident involving birds from agricultural uses has been reported. These reported incidents support what was concluded in the risk assessment for bifenthrin and the need for mitigation.

The USEPA EIIS database included 19 incident reports of effects on plants occurring between 1999 and 2012. The 1999 incident was classified as probable that the effects observed were a result of application of bifenthrin to a rose plant. The remaining 18 plant incidents were classified as possible. Fourteen of the remaining plant incidents occurred in 2012 following application to plants of the same product which contained two active ingredients and a high percentage of formulants. Therefore, the formulants and other active ingredient have contributed to these fourteen incidents. No plant incidents were noted since 2012. As bifenthrin is still registered for direct application on various plants, such as garden vegetables, roses, ornamentals and agricultural crops in the United States without new incidents being reported in recent years, other unknown factors may have contributed to the historical plant incidents.

Seven pollinator incidents were reported in the USEPA EIIS associated with various bee species between 1992 and 2013. Four of the incidents were classified as either probable or high probable that the mortality observed resulted from exposure to bifenthrin.

For the remaining three incidents it was considered possible that bifenthrin contributed to the effects observed; however, other toxic active ingredients were also identified in the incident report. The observation of bee mortality incidents supports the conclusions of the current risk assessment and the need for mitigation measures to protect pollinators.

The PMRA concluded that the information from the incident reports is consistent with the known toxicity hazard of bifenthrin to fish and pollinators.

4.2.4 Use of Vegetative Filter Strips for Reducing Runoff to Aquatic Habitats

To reduce movement of bifenthrin into aquatic habitats via runoff, the PMRA is requiring that 10 metre vegetative filter strips be mandatory in areas of use.

Since 2008, the USEPA has required statements on all pyrethroid agricultural product labels requiring a 3.05 metre (10 ft) vegetative filter strip (VFS) composed of grass or other permanent vegetation between the field edge and aquatic habitats. No fish kill incidents have been reported in the United States since the implementation of these requirements. The absence of further incidences in relation to adherence/implementation of the VFS is uncertain, however, VFS's have been shown to reduce movement of contaminants, excess nutrients, soil and other detrimental components into aquatic systems.

In 2000, the province of Prince Edward Island (PEI) introduced buffer legislation which mandated vegetative filter strips for various land uses, including agricultural crops. The legislation required all agricultural fields that border water courses to maintain a 10 metre vegetative filter strip along the water edge. The minimum buffer width was increased to 15 metre in a 2008 amendment to PEI's Environmental Protection Act. Fields with steeper slopes (i.e. >5%) within 50 metres of the upland boundary of the 10 metre buffer and having no other mitigating management practices in place are required to have a 20 metre vegetative filter strip.

The EU has also adopted the use of VFS for sustainable use of pesticides. EFSA has proposed 20 metre VFS's for a number of crop protection products.

5.0 Value

5.1 Consideration of Benefits

5.1.1 Potatoes

Capture 240 EC controls wireworms which are important pests of potatoes that live in the soil and damage potato tubers. Growers identified the use of bifenthrin to control wireworms in potatoes as a high priority, both through the Canadian Grower Priority Database and the Canadian Minor Use Priority Setting Workshop.

At the time Capture 240 EC was registered in 2014, the last date of use for Thimet 15G, a product containing phorate that was used to control wireworms on potato, was in August 2015. Thimet 20-G, a new product containing this active ingredient, was registered in July 2015. As a result, bifenthrin to control wireworm on potato is no longer considered as a critical need due to the availability of a suitable alternative management tool for wireworms. Registered active ingredients for management of wireworms in potato are phorate and chlorpyrifos in mode of action group 1B and clothianidin in mode of action group 4A.

5.1.2 Raspberry

Capture 240 EC controls obliquebanded leafroller, orange tortrix, black vine weevil, obscure root weevil, clay coloured weevil and earwigs in raspberries. The pre-bloom application targets clay coloured weevil adults, which feed on raspberry buds in early spring. The post-bloom application targets black vine weevil, obscure root weevil, earwigs, obliquebanded leafroller and orange tortrix, which are contaminants of mechanically harvested raspberries.

Clay coloured weevils occur in the spring when it is too cold to use the one registered alternative, malathion (mode of action group 1B). Viable alternatives are available to control obliquebanded leafroller, including spinosad and spinetoram (mode of action group 5), *Bacillus thuringiensis* subspecies *kurstaki* (mode of action group 11) and methoxyfenozide (mode of action group 18). Weevils that occur at harvest, black vine weevil and obscure root weevil, are the most abundant harvest contaminants. Thiamethoxam is the only alternative for control of these pests. Though there are limited or no alternatives for earwigs and orange tortrix, these are not considered to be of significant concern because of the potential to register alternative active ingredients.

Considering the alternatives and impact caused by the insects, a critical need for bifenthrin remains to provide control of insect contaminants in raspberries, specifically obscure root weevil and black vine weevil, throughout the harvest season. As well, bifenthrin contributes to resistance management because it is a new mode of action (mode of action group 3A) for use against insect pests in raspberries.

5.2 Effectiveness Against Pests

Nine efficacy trials on potatoes were reviewed and demonstrated control of wireworms in potatoes at rates of 2.0–3.4 g a.i./100 m row.

Four field trials and three laboratory trials were reviewed to support control of listed pests in raspberries. The data demonstrated control of black vine weevil, obscure root weevil, clay coloured weevil and earwigs in raspberries at a rate of 112 g a.i./ha. Control of obliquebanded leafroller and orange tortrix in raspberries at a rate of 112 g a.i./ha was supported based on extrapolation from data provided on other lepidopteran pests. Application timing to control obliquebanded leafroller and orange tortrix activity in raspberries is similar to that of earwigs, black vine weevil and obscure root weevil.

5.3 Non-Safety Adverse Effects

No phytotoxicity was observed in any of the reviewed trials.

5.4 Supported Uses

Value information provided for the registration of Capture 240 EC supported control of wireworms in potato applied in furrow at the time of planting. However, there is presently not a critical need for this use because of the availability of other products to manage this pest. Therefore, this use is no longer supported.

Value information provided for the registration of Capture 240 EC supported control of clay coloured weevils, obliquebanded leafroller, orange tortrix, black vine weevil, obscure root weevil, and earwigs by foliar application in raspberries. Critical need has been established for foliar applications on raspberries in British Columbia to control clay coloured weevil and insects that are present as harvest contaminants, particularly the black vine and obscure root weevils.

6.0 Pest Control Product Policy Considerations

6.1 Toxic Substances Management Policy Considerations

The Toxic Substances Management Policy (TSMP) is a federal government policy developed to provide direction on the management of substances of concern that are released into the environment. The TSMP calls for the virtual elimination of Track 1 substances [those that meet all four criteria outlined in the policy, i.e., persistent (in air, soil, water and/or sediment), bio-accumulative, primarily a result of human activity and toxic as defined by the *Canadian Environmental Protection Act*.

During the review process, bifenthrin and its transformation products were assessed in accordance with the PMRA Regulatory Directive DIR99-03⁶ and evaluated against the Track 1 criteria. The PMRA has reached the following conclusions:

- The PMRA considers bifenthrin to be persistent, bioaccumulative and toxic (PBT) and it meets all the key criteria for a Track 1 substance under Canada's TSMP policy:
 - o Bifenthrin is CEPA-Toxic equivalent.
 - o Bifenthrin is persistent in the environment under most circumstances. Bifenthrin meets the TSMP Track 1 criteria for soil and aquatic systems.
 - o Laboratory BCF values and the field BAF estimates exceeded the TSMP criteria of greater than 5000 for several fish species. One study showed that bifenthrin does not biomagnify in fish when only considering the diet under laboratory conditions (BMFs are <1.0). Under field conditions, however, sufficient information was provided to show that bifenthrin BAFs > 5000 are sustained long after applications. The field BAF values are a better representation of bioaccumulation under environmentally relevant conditions and consider multiple pathways of exposure.

Bifenthrin meets the Government of Canada's criteria for a Track 1 Substance under the Toxic Substances Management Policy adopted in 1995. These criteria are also outlined in the Persistence and Bioaccumulation Regulations (SOR/2000-107) which are used to define persistent or bioaccumulative under sections of the CEPA, 1999.

DIR99-03, The Pest Management Regulatory Agency's Strategy for Implementing the Toxic Substances Management Policy

The PMRA's implementation of the TSMP is outlined in DIR99-03 describes how Track 1 substances will be managed. As this policy calls for the virtual elimination of Track 1 substances, the PMRA is proposing to phase out the use of bifenthrin over three years. The use of Capture 240 EC for control of wireworm on potato will be cancelled immediately, as of 31 December 2017 as an additional pest management tool has become available since the original registration. For the use of Capture 240 EC for the control of weevils on raspberries, a phaseout period of 3 years is proposed. Since no suitable alternative exists for raspberries, the PMRA has determined that bifenthrin meets the critical need criteria as outlined in DIR99-03 which would allow for its use on a time-limited basis with the imposition of restrictions on the registration designed to minimize the risks associated with its use. Specific risk mitigation measures designed to minimize the risks include: limiting the use to a small geographic location (raspberries in British Columbia), buffer zones to minimize spray drift, vegetative filter strip to reduce runoff, and prohibiting application during crop blooming period.

6.2 Formulants and Contaminants of Health or Environmental Concern

During the review process, contaminants in the technical and formulants and contaminants in the end-use products are compared against the *List of Pest Control Product Formulants and Contaminants of Health or Environmental Concern* maintained in the *Canada Gazette*. The list is used as described in the PMRA Notice of Intent NOI2005-01⁸ and is based on existing policies and regulations including DIR99-03 and DIR2006-02⁹, and taking into consideration the Ozone-depleting Substance Regulations, 1998, of the *Canadian Environmental Protection Act* (substances designated under the Montreal Protocol). The PMRA has reached the following conclusions:

• Technical grade Bifenthrin Technical Insecticide does not contain any formulants or contaminants of health or environmental concern identified in the *Canada Gazette*.

The use of formulants in registered pest control products is assessed on an ongoing basis through PMRA formulant initiatives and Regulatory Directive DIR2006-02.

Canada Gazette, Part II, Volume 139, Number 24, SI/2005-114 (2005-11-30) pages 2641–2643: List of Pest Control Product Formulants and Contaminants of Health or Environmental Concern and in the order amending this list in the Canada Gazette, Part II, Volume 142, Number 13, SI/2008-67 (2008-06-25) pages 1611-1613. Part 1 Formulants of Health or Environmental Concern, Part 2 Formulants of Health or Environmental Concern that are Allergens Known to Cause Anaphylactic-Type Reactions and Part 3 Contaminants of Health or Environmental Concern.

NOI2005-01, List of Pest Control Product Formulants and Contaminants of Health or Environmental Concern under the New Pest Control Products Act.

⁹ DIR2006-02, Formulants Policy and Implementation Guidance Document.

7.0 Summary

7.1 Human Health and Safety

The toxicology database submitted for bifenthrin is adequate to define the majority of toxic effects that may result from exposure. In short-term and chronic studies on laboratory animals, the primary target of toxicity was the neurological system. There was no evidence of carcinogenicity in rats after longer-term dosing, and lung tumors in female mice and liver tumors in male mice were considered equivocal. Bifenthrin did not damage genetic material. Bifenthrin was not teratogenic in rats or rabbits, and did not cause any adverse effects on reproduction in rats. There was a low level of concern for increased sensitivity of the offspring; however, residual uncertainty remains regarding susceptibility of the young to the effects of pyrethroids. The risk assessment protects against the toxic effects noted above by ensuring that the level of human exposure is well below the lowest dose at which these effects occurred in animal tests.

Mixer, loader, applicators handling bifenthrin and workers re-entering treated areas are not expected to be exposed to levels of bifenthrin that will result in an unacceptable risk when bifenthrin is used according to label directions. The personal protective equipment on the product label is long-sleeved shirt, long pants, chemical resistant gloves, shoes plus socks and goggles during mixing, loading, application, clean up and repair.

The nature of the residues in plants and animals is adequately understood. The residue definition for enforcement is bifenthrin in plant products and in animal matrices. The use of bifenthrin on potatoes and raspberries does not constitute a risk of concern for chronic (cancer and non-cancer) or acute dietary exposure (food and drinking water) to any segment of the population, including infants, children, adults and seniors. Sufficient crop residue data have been reviewed to support the established MRLs for this active ingredient, as found in the Maximum Residue Limit Database on the Maximum Residue Limits for Pesticides webpage for MRLs

7.2 Environmental Risk

Bifenthrin meets the Government of Canada's criteria for a Track 1 Substance under the Toxic Substances Management Policy adopted in 1995. These criteria are also outlined in the Persistence and Bioaccumulation Regulations (SOR/2000-107) which are used to define persistent or bioaccumulative under sections of the *Canadian Environmental Protection Act* (CEPA), 1999. Under TSMP, substances meeting these criteria (Track 1) are persistent, bioaccumulative, toxic and primarily the result of human activity. Track 1 substances that cannot be managed successfully throughout their life cycle are targeted for vitual elimination from the environment, i.e., phase out of generation and uses.

Bifenthrin is moderately persistent to persistent in the terrestrial environment. The rate of transformation in the environment varies with environmental conditions such as soil type and climate. It is immobile and has a limited potential to leach to groundwater.

Bifenthrin may reach aquatic environments through spray drift and surface runoff. In the aquatic environment, it is expected to partition from the water layer and persist in the sediment. Therefore, continuous repetitive applications will increase the likelyhood of chronic risk to benthic communities as bifenthrin residues in the sediment increase overtime.

Bifenthrin is bioaccumulative and can accumulate in organisms under field conditions. There is some evidence that accumulated bifenthrin could potentially impact fitness and survival in aquatic organisms as observed in the field aquatic study.

Bifenthrin may pose a potential risk to pollinators, beneficial predatory and parasitic arthropods, small wild mammals, aquatic invertebrates, fish and amphibians.

Interim risk reduction measures are proposed to minimize environmental exposure during the phase out period and include the following: no-spray buffer zones to protect sensitive aquatic habitats from spray drift, and through the use of mitigative label statements to inform users of potential risks to the environment and indicate appropriate risk reduction measures.

7.3 Value

Value information demonstrated that Capture 240 EC controls wireworms in potatoes; however, bifenthrin is not a critical need because of the availability of products to manage this pest. Therefore, this use is no longer supported.

Value information provided for the registration of Capture 240 EC supported control of clay coloured weevils, obliquebanded leafroller, orange tortrix, black vine weevil, obscure root weevil, and earwigs by foliar application in raspberries. On raspberries, there is zero tolerance for insect harvest contaminants that affect the marketability of the fruit. Capture 240 EC is considered a critical need to manage harvest contaminants of raspberries in British Columbia because of the lack of suitable alternative products. As well, bifenthrin is a new mode of action for resistance management of these pests in raspberries.

8.0 Proposed Regulatory Decision

Health Canada's PMRA, under the authority of the *Pest Control Products Act* and Regulations, is proposing a time limited registration for the sale and use of Bifenthrin Technical Insecticide and Capture 240 EC, containing the technical grade active ingredient bifenthrin, to control certain pests in raspberries.

The evaluation of the additional environmental information found that bifenthrin meets the criteria for Track 1 substances under the TSMP. The PMRA's implementation of the TSMP is outlined in DIR99-03. This directive describes how Track 1 substances will be managed by the PMRA, and calls for the virtual elimination of Track 1 substances. For use of Capture 240 EC for control of weevils on raspberries, a phase out period of three years is proposed. Since no suitable alternative exists for raspberries, the PMRA has determined that bifenthrin meets the critical need criteria as outlined in DIR99-03 which would allow for its use on a time-limited basis with the imposition of restrictions on the registration designed to minimize the risks associated with its use. Specific risk mitigation measures designed to minimize the risks include:

limiting the use to a small geographical location (raspberries in British Columbia), buffer zones to minimize spray drift, vegetative filter strips to reduce runoff, and prohibiting application during crop blooming period. Therefore, registration for the use of Capture 240 EC on raspberries is granted for use only in British Columbia for three years beginning 1 January 2018 until 31 December 2020. The use of Capture 240 EC for control of wireworm on potato will be cancelled as of 31 December 2017

List of Abbreviations

°C degrees Celsius

equalgreater thanless than

 \geq greater than or equal to

± plus or minus

↑ increase

↓ decrease

% percent

♀ female

male

λ wavelength

 λ wavelength ϵ emittance

α assimilation efficiency

μg microgram(s)μgL microliter(s)a.i. active ingredient

abs. absolute

ADI acceptable daily intake

AHETF Agricultural Handler Exposure Task Force

AGF aspirated grain fraction

AOPWIN model that estimates the gas-phase reaction rate for the reaction between the most

prevalent atmospheric oxidant, hydroxyl radicals, and a chemical and calculates

atmospheric half-lives for chemicals

app application appl. Application

AR applied radioactivity
ARfD acute reference dose

ARTF Agricultural Re-entry Task Force

ASAE American Society of Agricultural Engineers

ATPD Area treated per day
BAF bioaccumulation factor
BCF bioconcentration factor

BCF_K kinetic BCF

BCF_{K.G} kinetic BCF corrected for growth

BCF_{K.G.L} kinetic BCF corrected for growth and lipid content

BCF_{SS} steady state BCF BMD benchmark dose

BMDL₂₀ benchmark dose limit estimated for 20% effect

BMF biomagnification factor

BMF_K kinetic BMF

BMF_{KG} kinetic BMF corrected for growth

BMF_{K G.L.} kinetic BMF corrected for growth and lipid content

BMF_{SS} steady state BMF

BP biphenyl

BW generic body weight

bw body weight bwg body weight gain

C_{max} maximum concentration CAF composite assessment factor CAS Chemical Abstracts Service

CDN Canadian

CEPA Canadian Environmental Protection Act

CG crop group

CI confidence interval

cm centimetre(s)

cm² centimetre(s) squared cm³ centimetre(s) cubed

CP cyclopropyl CSG crop sub-group

d day(s)

DAA day(s) after application

DACO data code

DFR Dislodgeable foliar residue

DIR directive

DNA deoxyribonucleic acid DNT developmental neurotoxicity

DT₅₀ dissipation time 50% (the dose required to observe a 50% decline in

concentration)

DT₉₀ dissipation time 90% (the dose required to observe a 90% decline in

concentration)

dw dry weight

EC emulsifiable concentrate or European Commission EC_{50} effective concentration on 50% of the population

EDE estimated daily exposure

EEC estimated environment concentrations EFSA European Food Safety Authority

EIIS USEPA Ecological Incident Information System

 $\begin{array}{ccc} EP & end-use \ product \\ ERC & evaluation \ report \\ F_0 & parental \ generation \\ F_1 & first \ generation \end{array}$

FA fraction of species affected

FC food consumption
FDA Food and Drugs Act
FIR food ingestion rate
FMC FMC Corporation

FOB functional observation battery

g gram(s)
G granule
gal gallon(s)

GAP good agricultural practice

GC-ECD gas chromatography with electron capture detection

GC-MS gas chromatography mass spectroscopy

ha hectare(s)

HAFT highest average field trial

hazardous concentration to five percent of species HC₅

HDPE high density polyethylene

mercury Hg

HPLC high performance liquid chromatography

hr

IUPAC International Union of Pure and Applied Chemistry

kilogram(s) kg

 $K_{\rm d}$ soil-water partition coefficient

kilometre(s) km

organic-carbon partition coefficient $K_{\rm oc}$ *n*–octanol-water partition coefficient K_{ow}

KiloPascals kPa L litre(s)

LAFT lowest average field trial

lb pound(s)

lethal concentration 50% LC_{50}

LC/MS/MS high performance liquid chromatography with tandem mass spectrometry

LD lactation day lethal dose 50% LD_{50}

lowest observed adverse effect level LOAEL

LOC level of concern

LOEC low observed effect concentration

limit of quantitation LOO lethal rate 50% LR_{50}

meter(s) m m^3 cubic metre(s)

MAS maximum average score

maximum Max. milligram(s) mg minute(s) min. Min. minimum

MIS maximum irritation score

ml millilitre(s)

M/L/A mixer/loader/applicator

millimeter(s) mm MOE margin of exposure MRL maximum residue limit mass spectrometry MS

MSD mass selective detection or mass spectrometric detection

n number NA not analysed not applicable N/A

NAFTA North American Free Trade Agreement

US Geological Survey's National Water-Quality Assessment Program NAWQA

National Institute for Occupational Safety and Health **NIOSH**

nanometre(s) nm NM not modelled

NMR nuclear magnetic resonance

no observed adverse effect concentration NOAEC

no observed adverse effect level NOAEL **NOEC** no observed effect concentration

NOEL no observed effect level **NOER** no observed effect rate

NOI Notice of Intent **NZW** New Zealand white

OECD Organisation for Economic Co-operation and Development

OH hydroxy

parental generation P

Pascal(s) Pa

plantback interval **PBI**

polychlorinated biphenyl PCB **PCPA** Pest Control Product Act Prince Edward Island PEI Pf processing factor

PH phenyl

Pesticide Handlers Exposure Database **PHED**

preharvest interval PHI dissociation constant pK_a

PMRA Pest Management Regulatory Agency

postnatal day **PND** parts per billion ppb parts per million ppm

EFSA'a Pesticide Risk Assessment Peer Review unit **PRAPeR**

raw agricultural commodity **RAC** REI restricted-entry interval

relative rel.

ROLD repeat oral low dose

RO risk quotient

 RT_{25} residual time to 25% bee mortality

RTI retreatment interval SC soluble concentrate SD standard deviation **SOHD** single oral high dose SOLD single oral low dose

SSD species sensitivity distribution supervised trial median residue STMdR

STORET EPA Storage and Retrieval Data Warehouse

total applied radioactivity TAR TC Transfer coefficient **TFP** trifluoro-1-propenyl-

TFP acid cis-3-(2-chloro-3,3,3-trifluoroprop-1-en-1-yl)-2,2-

dimethylcyclopropanecarboxylic acid

toxicity exposure ratio **TER**

 T_{max} time to peak blood concentration

TP transformation product TRR total radioactive residue

TSMP Toxic Substances Management Policy

US United States of America

USDA United States Department of Agriculture

USEPA United States Environmental Protection Agency

USGS U.S. Geological Survey

UV ultraviolet

VFS vegetative filter strip

vs. verses wk week(s)

WP wettable powder WSB water soluble bag

wt weight

w/w weight per weight dilution

- 1	ict	∩t.	Λ hh	rovi.	ations

Appendix I Tables and Figures

Table 1 Residue Analysis

Matrix	Method ID	Analyte	Method Type	LOQ		Reference
Fish ¹	N/A	bifenthrin	GC-ECD	1 ppb		1755330
Soil	N/A	bifenthrin	GC-MS	10 ppb		1755362
	N/A	4'-hydroxy bifenthrin	LC/MS/MS	10 ppb		1755361
	N/A	TFP acid	LC/MS/MS	10 ppb		1755361
Sediment ²	N/A	bifenthrin	GC-MS	0.1 ppb		1755356
Surface water	N/A	bifenthrin	GC-MS	0.1 ppb		1755362
	N/A	4'-hydroxy bifenthrin	LC/MS/MS	0.01 ppb		1755361
	N/A	TFP acid	LC/MS/MS	0.1 ppb		1755361
Air ³	N/A	bifenthrin	GC-MS	$0.5 \mu\text{g/m}^3$		1755346
	P-0757, P-1073	Bifenthrin	Enforcement: GC-ECD	0.05 ppm	Apples, strawberries	1762180; 1762256
	P-2132M	Bifenthrin	Data-gathering: GC-ECD	0.5 ppm	Apples, cottonseed, field corn (silage, stover, grain)	1762165
Plant	P-2281M	Bifenthrin	Data-gathering: GC-ECD	0.01 ppm	Field corn (grain, meal, flour, oil, starch, grits)	1762171
	P-2550M	Bifenthrin	Data-gathering: GC-ECD	0.1 ppm	Field corn grain Field corn silage Field corn stover	1762179
	P-2763 (Revised 4/99)	Bifenthrin	Data-gathering: GC-ECD	0.05 ppm	Walnut (nutmeat), peanut (nutmeat, soapstock)	1762181
	FCC 0596	Bifenthrin	Data-gathering: GC-ECD or MSD	0.05 ppm	Tea (fresh, green, black)	1828905
	RAN-0140	Bifenthrin	Data-gathering: GC-ECD		Cottonseed	1762182
	P-1031 (Revised)	Bifenthrin	Enforcement: GC-ECD	0.02 ppm 0.10 ppm 0.05 ppm		1762177
	P-1843M (Revised)	Bifenthrin BP alcohol	Enforcement: GC-ECD or MSD	0.02 ppm		1762178
	RAN-0204M	Bifenthrin	Enforcement: GC-ECD	0.01 ppm	Eggs	1762176
Animal (except fish)	P-1703M	Bifenthrin	Data-gathering: GC-ECD	0.2 ppm	Milk fat	1762170
	P-2533M	4'-OH- bifenthrin	Data-gathering: GC-ECD	0.5 ppm	Fat	1762163
	P-1704M	BP alcohol	Data-gathering: GC-MSD	0.02 ppm 0.05 ppm	Milk Liver, kidney, muscle, fat	1762172
	BP acid		0.05 ppm	Liver, kidney, muscle, fat		
	RAN-0203M	BP alcohol	Data-gathering: GC-MSD	0.01 ppm	Eggs	1762167
	P-1883M	TFP acid	Data-gatheringGC-MSD	0.05 ppm	Liver	1762173

For 4'-hydroxy bifenthrin and TFP acid, the corresponding methods for cow fat can be used.

For 4'-hydroxy bifenthrin and TFP acid, the corresponding methods for soil can be used.

Bifenthrin is also found in air, for which the method was provided under this DACO since there is no distinct DACO for air.

Table 2 Toxicity Profile of Capture 240 EC Containing Bifenthrin

(Effects are known or assumed to occur in both sexes unless otherwise noted; in such cases, sexspecific effects are separated by semi-colons.)

Study Type/Animal/ PMRA #	Study Results
Acute Toxicity Studies (Capture 2EC fo	ormulation used as surrogate for listed toxicity studies)
Acute oral toxicity	LD ₅₀ : (undiluted)
Sprague-Dawley rats	Q = 159 mg/kg bw
	Combined = 167 mg/kg bw
PMRA #1762133	
	High acute toxicity
Acute dermal toxicity	$LD_{50} > 2000 \text{ mg/kg bw}$
NEWY 111	
NZW rabbits	T
DIAD A #17/20104	Low acute toxicity
PMRA #1762134	
Acute inhalation toxicity (whole body)	LC ₅₀ :
Cama ana Davidan mata	0 = 1.6 mg/L
Sprague-Dawley rats	$\bigcirc = 2.3 \text{ mg/L}$
PMRA #1762135	Combined = 1.9 mg/L
PMRA #1/02133	Clicht coute toxicity
Dermal Irritation	Slight acute toxicity MAS = 1.3
	MIS = 1.4 (24h)
NZW rabbits	11113 – 1.4 (2411)
142 W Tabbits	
PMRA #1762137	Slightly irritating
Eye Irritation	MAS = 3.9
	MIS = 9.7 (1h)
NZW rabbits) (III)
1,2,7,1,000,100	
PMRA #1762136	Minimally irritating
Dermal sensitization	Sensitizer (inadequate study)
(Buehler test)	
Hartley guinea pigs	
PMRA #1762138	

Table 3 Toxicity Profile of Technical Bifenthrin

(Effects are known or assumed to occur in both sexes unless otherwise noted; in such cases, sexspecific effects are separated by semi-colons. Organ weight effects reflect both absolute organ weights and relative organ to bodyweights unless otherwise noted)

	Study Results
Toxicokinetic Studies	
Metabolism Excretion (gavage)	Bile duct-cannulated rats were dosed with single oral doses of bifenthrin. Excreta and tissues were assayed to determine route of excretion and the nature of metabolites.
Sprague-Dawley rats	or metalogical and the matare of metalogical
(bile duct-cannulated) PMRA #1755390	Total oral absorption was estimated to be about 50% or 36% in female and male rats respectively. Feces excretion: $25\% \ 3/49\% \ 2$; biliary excretion: $19\% \ 3/30\% \ 2$; urinary excretion: $11\% \ 3/15\% \ 2$. Gut
	microflora had minimal impact on bifenthrin degradation. Bifenthrin metabolites in feces were the result of biliary excretion. The primary metabolites were OH-Me-bifenthrin, 4'-OH-bifenthrin and polar metabolites.
Distribution	Male and female rats were given, by oral gavage, a single radiolabelled
Excretion (gavage)	bifenthrin. Excreta were monitored for 7 days. Seven days after dosing, rats were killed and tissues assayed for radioactivity.
Sprague-Dawley rats	
PMRA #1755391	Minimal amounts found in tissues, greatest amount found in fat. \bigcirc exposed to the alcohol labeled ¹⁴ C-bifenthrin had the greatest amount in fat (twice as much as \bigcirc and animals exposed to the acid labeled ¹⁴ C-bifenthrin).
	The primary route of excretion was via feces (76-84%). Most of the radioactivity was excreted in feces and urine by 48 hours. Excretion times were similar for all groups (majority excreted 8-48 hours post dose), with the exception of $\[\]$ exposed to the alcohol labeled ¹⁴ C-bifenthrin (majority excreted 12-48 hours post dose).
	Unchanged bifenthrin was the major product found in feces and urine (37-46% ♂/28-47% ♀). Unidentified metabolites amounted to 24-27% ♂/22-40% ♀. Amount metabolised was similar between labeled moieties, however ♀ exposed to the alcohol labeled ¹⁴C-bifenthrin had a larger amount metabolised (40% vs 22-27% in other groups) and less recovered as unchanged bifenthrin (28% vs 37-47% in other groups).
	Supplemental: Urinary metabolites not characterized, metabolites not identified.
Distribution	Animals were dosed with radiolabeled bifenthrin as a single low or
Metabolism Excretion (gavage)	high dose, or dosed with a low dose for two weeks.
Sprague-Dawley rats	Minimal amounts found in tissues, largest administered dose found in fat of single high-dose \mathcal{L} .

	Study Results
PMRA #1755392	Feces was the primary route of excretion (68.9-82.8% $3/94.0-96.2\%$
	♀). Most of the administered dose was recovered by 72 hours.
	Unchanged bifenthrin was the major product found in feces (17.2 - 44.2% of the administered dose). Twelve other metabolites, derived from hydrolysis and oxidation of bifenthrin were also detected. These included 4'-OH-hydroxy-methyl-bifenthrin; 3'-OH-hydroxy-methyl-bifenthrin, hydroxy-methyl-bifenthrin, 4'-OH-bifenthrin, 3'-monomethyl-catechol-bifenthrin and 4'-monomethyl-catechol-bifenthrin along with the hydrolytic and oxidative-hydrolytic products of bifenthrin including biphenyl alcohol, 4'-OH-biphenyl alcohol, biphenyl aldehyde, biphenyl acid, TFP acid hydroxymethyl-TFP acid. Unknown metabolites (three) found in the feces ranged between 0.2 - 3.6% of the total administered dose.
	Total organosoluble residues of the urine ranged from 10.6 - 17.6% of the administered dose. Unchanged bifenthrin was a negligible product (0.1% possibly due to faecal contamination). The urinary metabolites from hydrolysis and hydrolysis-oxidation products of bifenthrin included TFP acid, hydroxymethyl-TFP acid, biphenyl alcohol, 4'-OH-biphenyl alcohol, 3'-OH-biphenyl alcohol, 3'-monomethyl-catechol-biphenyl alcohol, 4'-monomethyl-catechol-biphenyl alcohol, biphenyl acid, 4'-OH-biphenyl acid and 4'-hydroxy-biphenyl acid methyl ester. Four unidentified metabolites combined did not exceed 1% of the dose. A maximum level of polar, water soluble degradates amounted to 5.5% of the dose.
Metabolism	Groups of male rats were given either a low (Group A, B) or high dose (Group C, D) and either blood samples taken at intervals (A, C) or
Sprague-Dawley rats	animals were killed at intervals (B, C) and tissues sampled.
(gavage)	Radioactivity levels in the blood of Group A animals peaked at 4 hours $(0.66 \pm 0.13 \mu\text{g/ml})$. Levels peaked at 6 hours $(3.29 \pm 1.62 \mu\text{g/ml})$ in
PMRA #1755393 and 1755394	Group C animals. Radioactivity was still detected in both groups at 72 hours. For the sacrificed animals, the highest radioactivity levels were observed at 4 hours (Group B) and 6 hours (Group D). Radioactivity levels within the low dose groups (Groups A and B) were comparable with the exception of the peak levels at 4 hours. The levels observed in Group B was more than 2-fold greater than that observed in Group A ($1.88 \pm 1.09 \mu\text{g/ml}$ in Group B vs. $0.66 \pm 0.13 \mu\text{g/ml}$ in Group A). There was a greater difference in radioactivity levels between the high dose groups (Group C and D). Radioactivity at 24 hours post dose was comparable, but there was large difference in levels observed at 3, 6 and 10 hours. Particularly at 6 hours when peak levels were observed ($8.78 \pm 2.89 \mu\text{g/ml}$ in Group C vs. $3.29 \pm 1.62 \mu\text{g/ml}$ in Group D).
	In Group B, a large amount of unchanged bifenthrin was detected in the plasma (0.103 ppm and 0.677 ppm at 2 and 4 hours post dose, respectively. At 10 hours, plasma levels decreased to 0.128 ppm. In Group D, unchanged bifenthrin was also detected, but to a lesser extent

	Study Results
	than Group B (0.734 ppm at 3 hours post dose, peaking to 3.29 ppm at
	6 hours and declining to 0.13 ppm by 24 hours). The major metabolites were biphenyl acid (0.037-0.317 ppm in Group B and 0.502-1.3 ppm in Group D) and biphenyl alcohol (0.091-0.615 ppm in Group B and 0.134-2.86 ppm in Group D). Unchanged bifenthrin and its metabolites declined in plasma over time with a corresponding increase in protein-bound residues of the total radiolabel.
Kinetics	Female rats were given a low dose of radiolabelled bifenthrin for 70
Metabolism Sprague-Dawley Rat	days. Animals were killed and tissues analysed at intervals after the initiation of dosing. Tissue samples were collected for up to 85 days after the last dose of bifenthrin. Sciatic nerve tissue was sampled in a
(gavage)	subset of these intervals (day 56 to termination of study).
PMRA #1755395	Radioactivity levels in plasma and whole blood was relatively low, starting at 0.01 $\mu g/ml$ and 0.01 $\mu g/g$, respectively, and peaking at 0.06 $\mu g/ml$ (day 70) and 0.06 $\mu g/g$ (days 49-70), respectively. Elimination from the plasma was relatively rapid with the concentration of ^{14}C -bifenthrin dropping to 0.02 and 0.01 $\mu g/ml$ at days 73 and 78, respectively. Elimination from whole blood was slower with the concentration of ^{14}C -bifenthrin dropping to 0.03 and 0.01 $\mu g/g$ on days 78 and 99, respectively. Radioactivity concentrations reached non-detectable levels after day 78 and 113 for plasma and whole blood, respectively. The highest concentrations of radioactivity were found in the fat (9.62 $\mu g/g$ at day 70) and skin (2.06 $\mu g/g$ at day 73) with half-lives of 51 and 50 days, respectively. The organs with higher fat concentrations (sciatic nerve and ovaries) had longer half-lives (42 and 37 days, respectively) and higher peak radioactivity concentrations (3.25 and 1.69 $\mu g/g$, respectively) while other organs (liver and kidney) had shorter half-lives (19 and 28 days, respectively) and lower peak concentrations of radioactivity (0.4 and 0.32 $\mu g/g$, respectively).
	Radioactivity detected in fat was predominantly unchanged bifenthrin (72%-85% of total radioactivity between days 7 and 155) with 3 metabolites making up the rest of the radioactivity detected. One metabolite was found in higher concentrations than the others, and percent of radioactivity detected increased steadily up until study termination, possibly indicating that metabolic processes continue to occur in the fat to facilitate excretion (metabolite is slightly more polar than unchanged bifenthrin).
	Potential for bioaccumulation
Metabolism and	Groups of male and female rats were gavage dosed with either a single
excretion	low bifenthrin dose or with a high dose of bifenthrin. Groups of rats were also dosed with low and high doses for 14 days of a high dose of
Sprague-Dawley Rat (gavage)	bifenthrin.

	Study Results
PMRA #1755399 and 1755403	Less than 1% of administered ¹⁴ C was detected in expired air in preliminary study, therefore, expired air not monitored in main study.
	Primary route of excretion was via feces (71.15-83.5%) and urine (9.37-14.46%). Ranges are in both sexes and all doses after 168 hours. Majority of radioactivity excreted within the first 48 hours in all dose groups (66.9-80.6% in feces, 7.26-11.27% in urine). Feces elimination slightly slower in SOHD animals (68.79% and 66.09% at 48 hours in 3 and 3 , respectively).
	Highest concentrations seen in fat and skin. Fat: SOLD-1.12 and 1.18 ppm; ROLD-1.44 and 1.27; SOHD-7.66 and 15.60 for \lozenge and \lozenge , respectively. Skin: SOLD-0.14 and 0.18; ROLD-0.19 and 0.21; SOHD-0.73 and 2.16 for \lozenge and \lozenge , respectively. Percent of administered dose in tissue ranged from 0.05-0.10% in fat, and <0.01-0.03% in skin.
	Feces metabolites excreted primarily as unchanged bifenthrin (21.80-39.22% of administered dose) and non-conjugates (including: hydroxymethyl-bifenthrin, 4'-OH-bifenthrin, 3'-OH-hydroxymethyl-bifenthrin, 4'-OH-hydroxy-methyl-bifenthrin 3'-monomethyl-catechol-bifenthrin and 4'-monomethyl-catechol-bifenthrin, dimethoxy-bifenthrin and 4'-methoxy-bifenthrin. Hydrolytic products related to mono- and dihydroxylated bifenthrin were also detected which included 4'-OH-biphenyl alcohol, 4'-OH-biphenyl alcohol, dimethoxy-biphenyl acid, dimethoxy-biphenyl alcohol, 4'-methoxy biphenyl alcohol and biphenyl alcohol, TFP acid, cis and trans-hydroxymethyl-TFP acid).
	In urine, no radioactivity was detected as unchanged bifenthrin in $\ \ \ $ while very little (0.01% of administered dose) was detected in SOHD $\ \ \ \ \ $ only. The major metabolites in alcohol labeled samples ($\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \$
Absorption	In a preliminary study, rats were dosed with radiolabeled bifenthrin and
Distribution	expired air collected. Very low levels of radioactivity were collected in
Excretion (gavage)	expired air. Based on this finding, expired air was not collected in the main study.
Sprague-Dawley Rat	In the main study, male and female rats were dosed with radiolabelled
PMRA #1755405	bifenthrin, either as a single low dose, a repeated (14 days) low dose, or

	Study Results
	a single high dose. The majority of radioactivity was collected in feces
	and urine. There were no difference seen between sexes and dose levels. After 168 hours 73-83% of administered dose excreted in feces versus 13-20% in urine (SOLD). For ROLD the range of excretion was 66-73% (feces) and 18.36-25.01% (urine), and for SOHD it was 69-71% (feces) and 22% (urine). Most of the radioactivity was excreted within 48 hours, but was slower in the SOHD. SOLD: 64-78% (feces), 11-17% (urine); ROLD: 50-69% (feces), 14-21% (urine); SOHD: 43-56% (feces), 18-15% (urine).
	Tissue distribution: SOLD – individual tissue residues were relatively low (0.004-1.502 ppm) and similar between sexes. Tissues with the most residue were fat > skin > ovaries > pancreas > lungs (\circlearrowleft) > liver. Total residues in all tissue ranged from 1.76-8.35% of administered dose (AD). ROLD – similar residues and distribution compared to SOLD. Residues for individual tissues ranged from 0.009-2.532 ppm while total residues ranged from 1.64-5.45% of AD. Tissue distribution was fat > pancreas > ovaries > skin > liver in \circlearrowleft and fat > pancreas > prostate > liver > skin in \circlearrowleft . SOHD – tissue residues were higher in \backsim . Individual tissue residues ranged from 0.087-23.895 ppm in \backsim and 0.036-4.380 ppm in \circlearrowleft . Total tissue residues were similar between sexes and ranged from 1.02-6.65% of AD. Tissue distribution was fat > skin > ovaries > pancreas in \backsim and fat > skin > liver > prostate in \circlearrowleft .
Kinetics - in vitro	In vitro metabolic rate constants in rat and human pooled hepatic microsomes were determined using several concentrations of
Long-Evans rat human	bifenthrin.
	Km and Vmax in rat hepatic microsomes:
	$Km = 5.42 \mu M (3.25-8.52 \mu M)$
PMRA# 2501803	Vmax = 0.64 nmol/min/mg (0.56-0.74 nmol/min/mg) Vmax/Km = 0.12 ml/min/mg
	Intrinsic clearance from rat and human liver microsomes : $CL_{int} \ (rat) = 224 \pm 20 \ ml/min/kg \ bw$ $CL_{int} \ (human) = 20 \pm 6 \ ml/min/kg \ bw$
	In the rat, the following P450 isoforms are involved in the metabolism of bifenthrin: CYP1A1, CYP1A2, CYP2B1, CYP2C6, CYP2C11, CYP2C12, CYP3A1, CYP3A2.
	In the human, the following P450 isoforms are involved in the metabolism of bifenthrin: CYP2C8, CYP2C9*1, CYP2C9*2, CYP2C9*3, CYP2C19.
In vitro comparative metabolism	To examine the relative metabolic capabilities of mice and rats, liver microsomes were prepared from male and female mice and male rats. Microsomes were incubated with bifenthrin for 0, 15, 30, or 60

	Study Results
Swiss-Webster Mouse	minutes.
Sprague-Dawley Rat	
PMRA #1755402	Recovery of radioactivity: Incubation time had no effect on radioactivity recovered in supernatant or pellet washes in all samples. Radioactivity recovered in the pellet increased from 0.1-1.2 % of the administered dose at 0 minutes to 2.1-5.2% of administered dose at 60 minutes, in a time dependent manner in all samples.
	Metabolism was not extensive. At 0 minutes, 78.1-88.6% administered radioactivity was unchanged bifenthrin, <15% were metabolites. At 60 minutes, 60.7-80.8% administered radioactivity was unchanged bifenthrin, <19% were metabolites. Identified metabolites 4-OH bifenthrin (0.55-3.44% and 2.25-5.95% for 0 and 60 minute incubation times, respectively) and BP acid (0-3.29% and 0-3.27% for 0 and 60 minute incubation times, respectively) were formed by ring hydroxylation and scission of bifenthrin, respectively.
	Metabolic activity between species was believed to be in the order male mouse > female mouse > male rat. Total identified and unidentified metabolites, respectively: cyclopropyl ring labelled bifenthrin-male mouse (8.02 and 15.97%), female mouse (3.48 and 2.17%), male rat (1.81 and 11.11%); phenyl-ring labelled bifenthrin-male mouse (3.84 and 23.86%), female mouse (5.58 and 5.79%), male rat (0.14 and 6.63%).
Acute Toxicity Studies	,
Acute Toxicity Studies Acute oral toxicity	LD ₅₀ : (in corn oil)
Acute oral toxicity	$\beta = 55.5 \text{ mg/kg bw}$
Sprague-Dawley Rat	Q = 53.4 mg/kg bw Combined = 54.5 mg/kg bw
PMRA #1755500	
	High toxicity
Acute oral toxicity	LD ₅₀ : (in corn oil)
Swiss Webster Mouse	
PMRA #1755501	Combined – 45.0 mg/kg ow
1 1411(1 11 1 1 1 1 1 1 1 1 1 1 1 1 1 1	High toxicity
Acute oral toxicity	LD ₅₀ : (in corn oil)
	$\delta = 70.1 \text{ mg/kg bw}$
Sprague-Dawley Rat	Q = 53.8 mg/kg bw
	Combined = 56.7 mg/kg bw
PMRA #1755502	
	High toxicity
Acute oral toxicity	LD ₅₀ : (undiluted)
	3 = 168.4 mg/kg bw
Sprague-Dawley Rat	Q = 210.4 mg/kg bw

	Study Results
	Combined = 186.1 mg/kg bw
PMRA #1755508	comemon room mg ng e w
	High toxicity
Acute dermal toxicity	LD ₅₀ :
	$\frac{3}{3}$ > 2000 mg/kg bw
Sprague-Dawley Rat	$\varphi > 2000 \text{ mg/kg bw}$
Sprague 2 amiej rau	Combined > 2000 mg/kg bw
PMRA #1755499	
	Low toxicity
Acute dermal toxicity	LD ₅₀ :
	$\delta > 2000 \text{ mg/kg bw}$
New Zealand White	$\varphi > 2000 \text{ mg/kg bw}$
Rabbit	Combined > 2000 mg/kg bw
114001	comemon 2000 mg/ng ovi
PMRA #1755498	Dermal effects: Irritation, desquamation
A	Low toxicity
Acute inhalation	LC_{50} :
toxicity (nose-only)	$\delta = 1.1 \text{ mg/L}$
	Q = 0.8 mg/L
Sprague-Dawley Rat	Combined = 1.01 mg/L
PMRA #1755497	
1 1/11(1 11 17 33 17 7	Slight acute toxicity
Eye Irritation	Non-irritating
Lyc minution	1 ton minuming
New Zealand White	
Rabbit	
PMRA #1755496	
Skin Irritation	No dermal irritation observed.
New Zealand White	Non-irritating Non-irritating
Rabbit	
DMD A #1755405	
PMRA #1755495	Canaitinan
Skin sensitization -	Sensitizer
Maximization Test	
Guinea Pig	
PMRA #1755492	
Skin sensitization -	Non-sensitizer
Buehler Method	
Hartley Guinea Pig	

	Study Results
DMD A #1755404	
PMRA #1755494 Short-Term Toxicity S	tudies
28-day oral toxicity	Study 1:
(diet) Range-Finding	$\geq 21.6/28.3 \text{ mg/kg bw/day} - (\circlearrowleft, \circlearrowleft)$: \downarrow abs. brain wt., \downarrow rel. brain wt.
Swiss Webster Mouse	Study 2: ≥108.0/120.4 mg/kg bw/day – (\circlearrowleft ; \circlearrowleft): tremors; \downarrow FC (\circlearrowleft , wk 1-3)
PMRA #1755478	\geq 112.8/186.7 mg/kg bw/day – (\updownarrow): mortality, \downarrow FC, \uparrow rel. liver wt.
	\geq 142.8/217.3 mg/kg bw/day – (\circlearrowleft): \uparrow bwg; (\updownarrow) overall \downarrow bwg, \downarrow FC
	164.1/120.5 mg/kg bw/day – (♂/ \updownarrow): ↓FC; (♂) mortality; (\updownarrow): ↓bw, overall ↑bwg
	Supplementary: Range-finding
28-day oral toxicity (diet) Range-Finding	\geq 5.38/5.42 mg/kg bw/day – (\circlearrowleft): \uparrow relative brain and relative kidney wt
	\geq 21.9/21.6 mg/kg bw/day – (\circlearrowleft / \updownarrow): tremors (\geq day 2); (\circlearrowleft): \downarrow bw and
Sprague-Dawley Rat	bwg; (♀): ↓FC
PMRA #1755477	34.5/32.6 mg/kg bw/day $-(\Im/\Im)$: mortality, clonic convulsions, \downarrow bw and bwg, \downarrow FC; $(\Im$ s): \downarrow adrenal wt, \downarrow testicular wt, \uparrow relative adrenal wt, brain and kidney wts; (\Im) : \uparrow relative brain, kidney, and liver wt
	Supplementary: Range-finding
90-day oral toxicity	NOAEL = 3.8/4.3 mg/kg bw/day
(diet)	LOAEL = 7.5/8.5 mg/kg bw/day; based on (\circlearrowleft / \updownarrow): tremors (days 3 to 5)
Sprague-Dawley Rat	Recovery period: tremors ceased within three days of the cessation of dosing
PMRA #1755483	
90-day oral toxicity	NOAEL = 2.5 mg/kg bw/day
(capsule)	LOAEL = 5 mg/kg bw/day; based on (∂/Q) : tremors; (Q) : ataxia, cyclic activity with signs of estrous
Beagle Dog	
PMRA #1755479, 1923291, 1923290, 1923289	
12-month oral toxicity	NOAEL = 1.5 mg/kg bw/day
(capsule)	LOAEL = 3 mg/kg bw/day; based on (\circlearrowleft / \updownarrow): tremors; (\circlearrowleft): \uparrow serum Na ⁺ , delayed estrous
Beagle Dog	

	Study Results
PMRA #1755481,	Study Results
19293292	
	NOAEL = 50 mg/kg bw/day
	Systemic LOAEL = 100 mg/kg bw/day; based on (\lozenge / \diamondsuit): tremors,
Sprague-Dawley Rat	staggered gait; (\lozenge): erythema; (\lozenge): skin ulceration, exaggerated
	hindlimb flexion, \tail flick latency
PMRA #1755474	, · · · · · · · · · · · · · · · · · · ·
	Dermal effects were noted at all doses, with severity increasing with
	dose;
	(3/2): desquamation, hyperplasia, hyperkeratosis, eschar, clinical
	signs of cutaneous paraesthesia (vocalization, thrashing in cage, lying
	on back)
21-day dermal toxicity	≥25 mg kg bw/day: Erythema
New Zealand White	500 mg/kg bw/day: (2/0). Tramore loss of muscle goordination:
Kauut	(♀): + abs. and ref. fiver wt
PMRA #1755475	Supplementary: Lack of historathology on nervous tissue (collected but
111111111111111111111111111111111111111	1 22
Chronic Toxicity/Oncos	
, , ,	
Swiss Webster Mouse	
	adenocarcinomas and adenomas
PMRA #1755453,	
1755387, 1755461,	Equivocal evidence of carcinogenicity
1755462, 1755463,	
1755464, 2376129	
	NOAEL = 2.3 mg/kg bw/day
(diet)	
	wt.; (\mathcal{P}) : tremors
Sprague-Dawley Rat	
D) (D) 1 114555 150	Non-carcinogenic
1	
·	
· ·	NT
	negauve
Iviutation Assay	
Salmonella typhimurium	
TA98, TA100, TA1535,	
TA1537, TA 1538	
PMRA #1755423	
Bacterial Reverse	Negative
New Zealand White Rabbit PMRA #1755475 Chronic Toxicity/Oncogonic Toxicity/Oncogonic Toxicity/Oncogonic Toxicity/Oncogonic Toxicity (diet) Swiss Webster Mouse PMRA #1755453, 1755461, 1755462, 1755463, 1755464, 2376129 Chronic/Oncogonicity (diet) Sprague-Dawley Rat PMRA #1755450, 175545, 1755452, 1755465, 1755452, 1755465, 1755449 Genotoxicity Studies Bacterial Reverse Mutation Assay Salmonella typhimurium TA98, TA100, TA1535, TA1537, TA 1538 PMRA #1755423	on back) ≥25 mg kg bw/day: Erythema 500 mg/kg bw/day: — (♂/♀): Tremors, loss of muscle coordination; (♀): ↑ abs. and rel. liver wt Supplementary: Lack of histopathology on nervous tissue (collected I not examined). Hematology, clinical chemistry, gross necropsy, and pathology data not reported. genicity Studies NOAEL not observed LOAEL = 7.6/10 mg/kg bw/day; based on (♂): tremors, bilateral germinal epithelial degeneration of the testes; (♀): bronchioalveolar adenocarcinomas and adenomas Equivocal evidence of carcinogenicity NOAEL = 2.3 mg/kg bw/day LOAEL = 4.7 mg/kg bw/day; based on ↑bw, ↑rel. liver, ↑rel. kidney wt.; (♀): tremors Non-carcinogenic

	Study Results
Mutation Assay	
Salmonella typhimurium	
TA98, TA100, TA1535,	
TA1537 and E. coli	
WP2 uvrA	
PMRA #1755428	
Bacterial Reverse	Negative
Mutation Assay	
Salmonella typhimurium	
TA98, TA1535 and	
TA1537	
PMRA #1755429	
Chromosomal	Negative
Aberrations	
CHO cells	
PMRA #1755415	
Chromosomal	Negative (non-activated)
Aberrations	
	Inconclusive (activated) – s.s. positive response at 20µg/ml but not at
CHO cells	higher doses. No dose response.
DIED A HARREALC	
PMRA #1755416	
Morphological	Negative
Transformation	
D A I D /2/72	
BALB/3T3 mouse	
embryo cells	
DMD A #1755417	
PMRA #1755417	No sodino
Fluctuation Assay	Negative
Mouse lymphoma	
L5178Y cells	
LJ1/01 CEIIS	
PMRA #1755418	
Mouse Lymphoma	Negative
Mutagenesis Assay	
2000	
L5178Y TK+/-	
L	

	Study Results
DMD A #1755421	Study Results
PMRA #1755421 Unscheduled DNA	Negative
Synthesis	Inegative
5 yittiesis	
Sprague-Dawley Rat	
hepatocytes	
DMD A #17772007	
PMRA #1755396	
Unscheduled DNA	Equivocal @ 2.0 <u>u</u> l/ml
Synthesis	
Sprague-Dawley Rat	
hepatocytes	
PMRA #1755397	
Mouse Lymphoma	Positive ± S9
Mutagenesis Assay	
L5178Y TK+/-	
L31701 TK17	
PMRA #1755413	
Sister Chromatid	Negative
Exchange Assay	
CHO II	
CHO cells	
PMRA #1755414	
In vivo Cytogenetics	Negative
Assay (Chromosome	
Aberration)	
Sprague-Dawley rat bone marrow	
bone marrow	
PMRA #1755408	
Mammalian Erythrocyte	Negative
Micronucleus Test	
TOP :	
ICR mouse bone	
marrow	
PMRA #1755410	
In vivo Unscheduled	Negative
DNA Synthesis Test –	
gavage	

	C4-, J-, D 4-,
G D 1	Study Results
Sprague-Dawley rat	
hepatocytes	
PMRA #1755406	
Gene Mutation Assay	Negative
CHO cells	1 togati vo
PMRA #1755388	
Unscheduled DNA	Negative
Synthesis	
Adult & Wistar/WU rat	
hepatocytes	
DMD A #1755200	
PMRA #1755389	4' . TD - '-'4 . C4 . 1'
Developmental/Reprod	
Multi-generation	Parental Toxicity NOAEL = 2 mg/kg bw/day
reproductive toxicity	NOAEL = 3 mg/kg bw/day
(diet)	LOAEL = 5 mg/kg bw/day; based on (\mathcal{P}): tremors P, F1dams, clonic
Spragua Davilay rat	convulsions; (\circlearrowleft): \downarrow FC (F1), \downarrow bw (P), \downarrow bwg (P, F1), \uparrow rel. brain weight
Sprague-Dawley rat	(P), ↓bw in P and F1 dams
PMRA #1755448	Reproductive Toxicity
	NOAEL = 5 mg/kg bw/day
	LOAEL = Not observed; (\mathcal{P}) : equivocal \uparrow litter incidence of stillborn
	pups possibly related to heating failure
	Offspring Toxicity
	NOAEL = 5 mg/kg bw/day
	LOAEL = Not observed
	Maternal Toxicity
(diet)	NOAEL = 7.1 mg/kg bw/day
	LOAEL = 15.5 mg/kg bw/day; based on (\updownarrow): tremors, splayed
Sprague-Dawley rat	hindlimbs, hypersensitivity to sound, piloerection, ↓bwg, ↓rel. FC
PMRA #1755431	Developmental Toxicity
1 1/11/21 1 / 25 + 31	NOAEL = 15.5 mg/kg bw/day
	LOAEL not observed
	EOTED Not 66561 (64
	No evidence for sensitivity of the young or malformations
Developmental toxicity	Maternal Toxicity
(gavage)	NOAEL = 1.0 mg/kg bw/day
	LOAEL = 2.0 mg/kg bw/day ; based on (\updownarrow): tremors
Sprague-Dawley Rat	
PMRA #1755432,	Developmental Toxicity
2376130	NOAEL = 1.0 mg/kg bw/day

	Study Results
	LOAEL = 2.0 mg/kg bw/day; based on $(3/2)$: hydroureter without
	hydronephrosis (variation)
	nydronephrosis (variation)
	No evidence for sensitivity of the young or malformations
Developmental toxicity	Maternal Toxicity
(gavage)	NOAEL 2.67 mg/kg bw/day
	LOAEL 4.0 mg/kg bw/day; based on (♀): head and forelimb twitching
New Zealand White	
Rabbit	Developmental Toxicity
	$NOAEL \ge 8.0 \text{ mg/kg bw/day}$
PMRA #1755430	LOAEL not observed
	No evidence for sensitivity of the young
Neurotoxicity Studies	
Acute-Delayed	LD ₅₀ Determination:
Neurotoxicity (gavage)	$LD_{50} > 5000 \text{ mg/kg bw (in corn oil)}$
Hen	\geq 1250 mg/kg bw: $-()$: unsteadiness, inability to walk, wing-
	dropping, twitching of head and neck. All birds recovered after 72
DMD A #1755 A 47	hours post-dose.
PMRA #1755447	
	Neurotoxicity Determination:
	Initial dose (Day 0) – (\updownarrow): unsteadiness, difficulty standing, jerking
	head movements (all animals in test groups). All birds recovered by day 4 except 2 in last dose group. \(\psi FC \)
	4 except 2 in last dose group. FC
	Second dose (Day 21) – (\updownarrow): mortality, \downarrow bw, unsteadiness, difficulty
	standing, jerking head movements, violent head and leg movements
	and trembling (all treated animals 4 days after repeat dose). Surviving
	birds recovered by day 28. \$\\$FC\$ more marked effect after repeat dose.
	onds recovered by day 20. \$1 0 more marked effect after repeat dose.
	No evidence of delayed neurotoxicity. There were no neuropathology
	findings, clinical effects were transient.
	Supplementary: limited statistical analysis, lack of neurochemical
	assessment
Acute-Delayed	Irwin dose range-finding study (in corn oil)
Neurotoxicity (Tilting-	10 mg/kg bw − (♂): moderately abnormal body carriage (hunched
Plane Test) (gavage)	posture) and slightly abnormal gait, tremors, apathy, and reduced
(non-guideline)	grooming
COBS/Wistar Rat	30 mg/kg bw − (♂): tremors, abnormal body carriage and gait, and
DMD A #17757444	respiratory depression, mortality, CNS depression and increased
PMRA #1755444	salivation, respiratory depression, clonic convulsions, abnormal body
	carriage, ptosis, greasy looking fur, paralysis, severe tremors and
	twitches.

	Study Results
	Didy Results
	Minimum effective dose eliciting severe neurological signs = 30 mg/kg/day (dose selected for tilting-plane test). No neurohistopathology was conducted.
	Tilting-plane test (in corn oil) 30 mg/kg bw – (\circlearrowleft / \updownarrow): slight increase in the mean angle of slip on day 2, mortality; (\updownarrow): stereotyped grooming and greasy appearance of fur, \downarrow bw.
	No neurohistopathology was conducted.
	No evidence of delayed neurotoxicity.
Acute Oral	4 hours post exposure (time of peak effect) with 100% (Z, 1R cis)
Neurotoxicity (gavage)	isomer:
	\geq 3 mg/kg bw − (δ): \downarrow mean total motor activity (in corn oil).
Non-Guideline Motor	
Activity	Using an exponential dose-response model, the following benchmark dose (BMD) values for motor activity were derived:
Long-Evans Rats	dose (BMD) values for motor activity were derived.
Long Lvans Rats	$BMD_{20} = 4.1 \text{ mg/kg bw}$
PMRA#2007554	$BMDL_{20} = 2.6 \text{ mg/kg bw}$
	Note: The BMDL ₂₀ was selected based on the normal variability of motor activity in historical control rats (i.e., 9.6% to 26%) (PMRA#2351167).
Acute oral neurotoxicity	NOAEL = 35 mg/kg bw
(gavage)	LOAEL = 75 mg/kg bw; based on clinical findings – (\circlearrowleft): decreased feces; (\updownarrow): mortality, twitching, abdominogenital staining, clonic
Sprague-Dawley Rats	convulsions, chromorhinorrhea, tremors. All clinical signs resolved
PMRA #1755440	after day 2. FOB findings – @ 6-8 hours post dose – $(3/2)$: whole body tremors, tense/rigid during handling; (3) : abnormal mobile posture, uncoordinated movement/ataxia; splayed hindlimbs; (2) : convulsions, unusual posture. @ end of day 0 – $(3/2)$: whole body tremors; (3) : localized spasms/twitching, staggered gait, abnormal posture, uncoordinated movement/ataxia, splayed hindlimbs, \downarrow landing foot-splay values; (2) : increased activity; decreased activity, convulsions, walking on toes, unusual immobile posture. Motor activity – (3) : decreased; (2) : increased.
28-day neurotoxicity	$100 \text{ ppm} - (\stackrel{\bigcirc}{+})$: tremors
Range finding (diet)	
Sprague-Dawley Rat	200 ppm – ($\circlearrowleft/$?): tremors; (\circlearrowleft): twitching, unthriftiness, dehydration.
PMRA #1755443	300 ppm – (\circlearrowleft / \hookrightarrow): tremors; (\circlearrowleft): twitching, unthriftiness, dehydration, \downarrow bw; (\hookrightarrow): mortality, \downarrow bwg, chromorhinorrhea, hypersensitivity to sound, splayed hindlimbs, abdominogenital staining

	Study Results
90-day Neurotoxicity	NOAEL 2.9 mg/kg bw/day
(diet)	LOAEL 6.0 mg/kg bw/day; based on $(3/2)$: tremors, twitching. FOB – (2) : tremors, \downarrow hindlimb grip strength
Sprague-Dawley Rat	(1). 1 1 3, 4 1 1 3 F 1 1 3.
PMRA 1755438	
Dietary feasibility and Range-Finding for DNT study	9.3/22.5 mg/kg bw/day: (♀): slight to moderate whole body tremors Cmax(milk) on LD 11 for all groups. Residue levels increased in
Sprague-Dawley Rat PMRA# 1755433	somewhat linear fashion with increased dietary concentration. Maternal plasma levels similar on LD 4 and 22. Plasma levels increased more than milk levels with increasing dose. Offspring plasma levels, dose response and time course profiles similar to maternal levels.
Developmental	Maternal Toxicity
Neurotoxicity (diet)	NOAEL = 3.6 mg/kg bw/day LOAEL = 7.2 mg/kg bw/day; based on (\mathcal{P}): tremors, \mathcal{T} grooming,
Sprague-Dawley Rat	clonic convulsions.
PMRA #1755435, 2376131	Offspring Toxicity NOAEL = 3.6 mg/kg bw/day LOAEL = 7.2 mg/kg bw/day; based on \uparrow mean grooming counts (\updownarrow at PND 21), \downarrow total motor activity, \uparrow T _{max} acoustic startle (\updownarrow at PND 20)
Impurity Studies	
Bacterial Reverse	Negative
Mutation Assay	
FMC 102032	
(Bifenthrin impurity)	
Salmonella typhimurium TA98, TA100, TA1535,	
TA1537 and TA1538	
PMRA #1755424/1755425	
Bacterial Reverse Mutation Assay FMC 78162 (Bifenthrin impurity)	Negative
Salmonella typhimurium TA98, TA100, TA1535, TA1537 and TA1538 PMRA #1755426	
Bacterial Reverse Mutation Assay FMC 78161 (Bifenthrin impurity)	Negative

	Study Results
	Study Results
Salmonella typhimurium	
TA98, TA100, TA1535,	
TA1537 and TA1538	
PMRA #1755427	
CHO Mutation Assay	Negative
FMC 102032	
(Bifenthrin impurity)	
DMD A #1555410	
PMRA #1755419	NT ('
Chromosomal	Negative
Aberrations	
FMC 102032	
(Bifenthrin impurity)	
CHO cells	
PMRA #1755420	
Mouse Lymphoma	Negative
Mutagenesis Assay	
FMC 78161 (Bifenthrin	
impurity)	
L5178Y TK+/-	
PMRA #1755422	
Unscheduled DNA	Negative
Synthesis	
FMC 102032 (bifenthrin	
impurity)	
PMRA #1755398	
Micronucleus	Negative
Cytogenetic Assay	
FMC 78161 (Bifenthrin	
impurity)	
ICR mouse bone	
marrow	
PMRA #1755412	

Table 4 Toxicology Endpoints for Use in Health Risk Assessment for Bifenthrin

Exposure	Study	Point of Departure and Endpoint	CAF ¹ or
Scenario			target
			MOE
Acute Dietary	Acute oral neurotoxicity	$BMDL_{20} = 2.6 \text{ mg/kg bw}$; based on	300
all populations	in rats	decreased motor activity in the rat; in the	
		co-critical rabbit developmental toxicity	
		study based on head and forelimb	
		twitching	
	ARfD = 0.009 mg/kg bw	1	
Chronic Dietary	Developmental toxicity	NOAEL = 1.0 mg/kg bw/day; based on	300
all populations	study in rats; co-critical	maternal tremors in rat; in the co-critical	
	1-year dog study	dog study, tremors and delayed estrous	
	ADI = 0.003 mg/kg bw/s	l day	
Short-,	21 day dermal toxicity	NOAEL = 50 mg/kg bw/day;	300
Intermediate-Term	in rats	based on staggered gait, exaggerated	
Dermal		hind-limb flexion, reduced tail flick	
all populations		latency	
Short-,	Developmental toxicity	NOAEL = 1.0 mg/kg bw/day; based on	300
Intermediate-and	study in rats	maternal tremors	
Long-Term			
Inhalation			
all populations			
Cancer	Cancer risk for equivoca	l lung and liver tumors was addressed thro	ugh the
	selected toxicology endp	points.	

¹ CAF (Composite assessment factor) refers to the total uncertainty and *Pest Control Products Act* factors for dietary and residential risk assessment; MOE refers to the target margin of exposure for occupational assessment

 Table 5
 Mixer/Loader/Applicator Dermal Exposure Estimates and MOEs

Application Method	Maximum Single Application Rate (kg a.i./ha) 1	ATPD ²	Dermal Unit Exposure Value (µg/kg a.i. handled) ³	Dermal Exposure (mg/kg bw/day) 4	Dermal MOE ⁵
Groundboom (M/L/A)	0.337	20 ha/day	84.12	0.007087	7060
Airblast (M/L/A)	0.112	20 ha/day	3820.44	0.10697	467
Manually Pressurized Handwand	0.112	150 L/day	943.37	0.0003962	126000
Backpack	0.112	150 L/day	5445.85	0.002287	21900

¹ Supported rate

² ATPD for potatoes using groundboom equipment (farmer) is based on the largest area that can be planted in a single day because it is application at-plant only.

³ Dermal unit exposure values for mixer/loader liquid open pour, groundboom applicator, low pressure handwand (M/L/A) and backpack (M/L/A) were taken from PHED. Airblast applicator dermal unit exposures were derived from AHETF. PHED M/L liquid open pour and AHETF airblast applicator unit exposure values were combined for total airblast M/L/A exposure.

⁴ Dermal Exposure (mg/kg bw/day) = Application Rate (kg a.i./ha) × ATPD × Unit Exposure (μg/kg a.i. handled) × Unit Conversion (mg/ 1000 μg) ÷ 80 kg bw. For sprayers, the ATPD was divided by the spray volume, 500 L/ha, provided by the

applicant.

Dermal MOE = NOAEL of 50 mg/kg bw/day ÷ Exposure; Target = 300

Table 6 Mixer/Loader/Applicator Inhalation and Combined Exposure Estimates and MOEs

Application Method	Maximum Single Applicatio n Rate (kg a.i./ha) ¹	ATPD ²	Inhalatio n Unit Exposure Value (µg/kg a.i. handled) ³	Inhalation Exposure (mg/kg bw/day) 4	Inhalatio n MOE ⁵	Combine d MOE ⁶
Groundboom (M/L/A)	0.337	20 ha/day	2.56	0.00021568	4640	2800
Airblast (M/L/A)	0.112	20 ha/day	10.68	0.00029904	3340	410
Manually Pressurized		150		0.00001898	52700	37200
Handwand	0.112	L/day	45.2	4	32700	
		150		0.00002608	38300	13900
Backpack	0.112	L/day	62.1	2	38300	

¹ Supported rate

 Table 7
 Postapplication Margin of Exposure on Raspberries

Re-entry activity	Peak DFR (μg/cm²) ¹	Transfer coefficient (cm²/hr)²	Dermal exposure (mg/kg bw/day) ³	MOE ⁴	REI ⁵
Irrigation (hand-set)	0.2919	1750	0.0511	979	0
Tying/training (max foliage)	0.2919	1400	0.0409	1220	0
Hand harvesting	0.2128	1400	0.0298	1679	PHI = 3

¹ Calculated using the default 25% dislodgeable on the day of application and 10% dissipation per day

Table 8 Major groundwater and surface water model inputs for Level 1 assessment of bifenthrin

Type of Input	Parameter	Value
Application	Crop(s) to be treated	Raspberries, potatoes
Information	Maximum allowable application rate per year (g	224, 337
	a.i./ha)	
	Maximum rate each application (g a.i./ha)	112, 337
	Maximum number of applications per year	2, 1
	Minimum interval between applications (days)	30
	Method of application	airblast, in-furrow or T-band

² ATPD for potatoes using groundboom equipment (farmer) is based on the largest area that can be planted in a single day because it is application at-plant only.

³ Inhalation unit exposure values for mixer/loader liquid open pour, groundboom applicator, low pressure handwand (M/L/A) and backpack (M/L/A) were taken from PHED. Airblast applicator inhalation unit exposures were derived from AHETF. PHED M/L liquid open pour and AHETF airblast applicator unit exposure values were combined for total airblast M/L/A exposure.

⁴ Inhalation Exposure (mg/kg bw/day) = Application Rate (kg a.i./ha) \times ATPD \times Unit Exposure (µg/kg a.i. handled) \times Unit Conversion (mg/ 1000 µg) \div 80 kg bw. For sprayers, the ATPD was divided by the spray volume, 500 L/ha, provided by the applicant.

⁵ Inhalation MOE = NOAEL of 1 mg/kg bw/day ÷ Exposure; Target = 300

⁶ Combined MOE = 1 ÷ [(1/Dermal MOE) + (1/Inhalation MOE)]; Target = 300

² Transfer coefficients obtained from ARTF data.

³ Exposure = (Peak DFR [μ g/cm²] × TC [cm²/hr] × 8 hours / (80 kg bw × 1000 μ g/mg)

⁴ Based on a NOAEL of 50 mg/kg bw/day, target MOE = 300 (see Table 3)

⁵ Minimum REI is 12 hours to allow residues to dry but MOEs were calculated assuming 0 days for REI. A PHI of 3 days has been established for hand harvesting of raspberries

Type of Input	Parameter	Value
Environmental	Hydrolysis half-life at pH 7 (days)	stable
Fate	Photolysis half-life in water (days)	41.7
Characteristics	Adsorption K_{oc} (ml/g)	72490 (20 th percentile of four
		$K_{\rm oc}$ values for "bifenthrin")
	Aerobic soil biotransformation half-life (days)	167 (90 th percentile
		confidence bound on mean of
		four half-life values adjusted
		to 25°C)
	Aerobic aquatic biotransformation half-life	276 (longest of two half-lives)
	(days)	
	Anaerobic aquatic biotransformation half-life	0 (only value available)
	(days)	

Table 9 Level 1 estimated environmental concentrations of bifenthrin in potential drinking water sources

	Groundwate	r (ug o i /I)	Surface Water (µg a.i./L) Reservoir				
Crop	Groundwate	1 (μg a.i./L)					
	Daily ¹	Yearly ²	Daily ³	Yearly ⁴	Simulation Average ⁵		
Raspberries	NM	NM	1.5*	0.29	0.25		
Potatoes	0	0	NM	NM	NM		

NM Not modelled

Table 10 Integrated Food Residue Chemistry Summary

NATURE OF THE RESIDUE – A _I	ton, Corn			RA # 1755 376; 1755	5373; 1755 5377	374; 175	5375;		
APPLES	PMRA # 1755373	755373 1 or 3 applications on surface of leaves with a pipet at					ipet at tota	al rates of	
	PMRA # 1755377	7 Tr							
Radiolabel Position	[Cy	[Cyclopropyl (CP) Ring] [Pher				[Phenyl (I	henyl (PH) Ring]		
Crop/Fraction		Apple/W	hole Fruit	,	Apple/Whole Fruit				
Harvest Interval (days)	0	7	14	21	0	7	14	21	
Overall TRR (mg/kg)	0.72	0.59	0.43	0.59	0.81	0.74	0.64	0.61	
Major Metabolites (> 10% of TRR)		Bifer	thrin			Bifen	thrin		
Minor Metabolites (< 10% of TRR)									
Crop/Fraction		Apple/	Leaves			Apple/	Leaves		
Harvest Interval (days)		2	8		28				
Overall TRR (mg/kg)	Not reported			Not reported					
Major Metabolites (> 10% of TRR)		Bifer	thrin	•	Bifenthrin				
Minor Metabolites (< 10% of TRR)			-			BP	acid		

¹ 90th percentile of daily average concentrations ² 90th percentile of yearly average concentrations ³ 90th percentile of yearly peak concentrations ⁴ 90th percentile of yearly average concentrations

s average of yearly average concentrations
The limit of solubility in pH 7 buffered water is 1 μg a.i./L

NATURE OF THE RESIDUE – A	pples, Cot	ton, Cor	'n				5374; 175	5375;	
					376; 175				
				op surface					
				otal rates o	t 37.2 μg/	ieat (**C-	CP label)	and 25.2	
			f (¹⁴ C-PH 1		.401 0		1 .	*.1	
	PMRA#	1 appl	ication to s	soil surface	, at 8-leaf	cotton gro	owth stage	, with a	
COTTON	1755375		syringe at t /ha (¹⁴ C-PI	otal rates o	1 2.3 Kg a.	1./na (C	-CP label)	and 2.7	
COTTON				n iabei) est substanc	ec were a	nnlied to	soil and or	าไร	
				.e. no sam					
				orted herei				, 010,	
	PMRA#			cotton seed				te of 1.3	
	1755376		/seed (¹⁴ C-		5 (6 5 55 u 5		· u totul lu		
Radiolabel Position	[C:		oyl (CP) R			[Phenyl (PH) Ring]	
Crop/Fraction				<u> </u>			n/Seeds	_	
Harvest Interval (days)					0	1	14	28	
Overall TRR (µCi)					0.37	0	.31	0.22	
Major Metabolites (> 10% of TRR)						Bife	nthrin		
Minor Metabolites (< 10% of TRR)		-							
Crop/Fraction			n/Leaves				/Leaves		
Harvest Interval (days)	0	14	28	maturity	0	14		maturity	
Overall TRR (mg/kg)	14.92	9.52	6.66	29.28	15.05	8.37	7.08	6.47	
Major Metabolites (> 10% of TRR)			enthrin				nthrin		
Minor Metabolites (< 10% of TRR)		TF	P acid				BP alcoho		
	Foliar treatment 2 applications at the 2				2 feet stage and post tassel stage (RTI				
	of 21 days) at total ra				ates of 0.48 kg a.i./ha (¹⁴ C-CP label)				
	and 0.43 kg a.i./ha (¹⁴ C-PH label)								
				s 30 days prior to silage stage at total a (¹⁴ C-CP label) and 0.48 kg a.i./ha					
CODN	rates of 0.53 kg a.1./f				E: corn plants had received 2 foliar				
CORN	husks				E: corn pi	ants nad r	eceived 2	Toliar	
	treatments) 3 applications at the 2 feet stage, post tassel stage and 30							nd 20	
	days prior to silege stage (PTIs of 21 and 30 days) at t								
	Soil treati	ment			a (14C-CP label) and 2.26 kg a.i./ha				
			(¹⁴ C-PH la		a (C C1 10001) and 2.20 kg a.i./ild				
Radiolabel Position	[C:	ycloprop	oyl (CP) R	ing]	[Phenyl (PH) Ring]				
Crop/Fraction			/Leaves		Corn/Leaves				
Type of Treatment		Foliar	to leaves			Foliar t	to leaves		
Harvest Interval (days)	0	7	14	30	0	7	14	30	
Overall TRR (mg/kg) – treated	29.51	19.89	20.68	20.48	29.11	25.87	26.00	25.39	
samples	27.31	17.09	20.00	20.40	۵۶.11	23.07	20.00	23.33	
Overall TRR (mg/kg) – control	0.12	0.16	0.24	0.22	0.13	0.19	0.23	0.21	
samples					0.10			J.21	
Major Metabolites (> 10% of TRR)			ifenthrin	C .1 .	5.7		fenthrin	.1	
Minor Metabolites (< 10% of TRR)	trans-B		n, 4′-OH-bi P acid	tenthrin,		,	'-OH-bifer ol, BP ald	,	
Crop/Fraction		Cor	n/Grain				/Grain		
Type of Treatment	Foliar to	o leaves		to leaves	Foliar t	o leaves		o leaves	
				husks			and	husks	
Harvest Interval (days)	mati			turity		urity		urity	
Overall TRR (mg/kg)	control	treated	_	treated	control	treated	control	treated	
- · · · · · · · · · · · · · · · · · · ·	0.053	0.057	0.053	0.063	0.056	0.056	0.056	0.069	
Major Metabolites (> 10% of TRR) Minor Metabolites (< 10% of TRR)			analysed analysed				nalysed nalysed		

NATURE OF THE RESIDUE – App	ples, Cotton, Corn		PMRA # 1755373; 1755376; 1755377	1755374; 1755375;	
Crop/Fraction	Corn/Wh	ole Plant	Corr	/Whole Plant	
Type of Treatment	Sc	oil		Soil	
Harvest Interval	At silag	ge stage	At	silage stage	
Overall TRR (mg/kg)	control	treated	control	treated	
Overall TKK (llig/kg)	0.21	0.06	0.04	0.06	
Major Metabolites (> 10% of TRR)	Not an	alysed	N	ot analysed	
Minor Metabolites (< 10% of TRR)	Not an	alysed	N	ot analysed	
Crop/Fraction	Corn/Stalks	and Leaves	Corn/S	talks and Leaves	
Type of Treatment	Soil			Soil	
Harvest Interval	At ma	turity	A	At maturity	
Organil TDD (ma/lsa)	control	treated	control	treated	
Overall TRR (mg/kg)	0.10	0.30	0.25	0.15	
Major Metabolites (> 10% of TRR)	Not an	alysed	N	ot analysed	
Minor Metabolites (< 10% of TRR)	Not an	alysed	N	ot analysed	
Crop/Fraction	Corn/	Husks	C	orn/Husks	
Type of Treatment	Sc	oil		Soil	
Harvest Interval	At ma	turity	A	At maturity	
Overall TDD (ma/lsa)	control	treated	control	treated	
Overall TRR (mg/kg)	0.17	0.07	0.19	0.24	
Major Metabolites (> 10% of TRR)	Not an	alysed	Not analysed		
Minor Metabolites (< 10% of TRR)	Not an	alysed	N	ot analysed	

CONFINED ACCUMULATION IN ROTATIONAL CROPS – PMRA # 1762331
Lettuce, sugar beet and wheat

Sandy loam soil was treated at a rate of 0.56 kg a.i./ha. Seeds of lettuce, sugar beet and wheat were sown at three plant-back intervals of 1, 2 and 4 months. Samples of wheat straw were analysed for identification/characterization of radioactive residues. Samples of sugar beet (roots, foliage), lettuce and wheat grain were not analysed further for identification/ characterization of radioactive residues although TRR levels in these matrices were above the trigger level of 0.01 ppm as per OECD Guidance Document on Overview of Residue Chemistry Studies (ENV/JM/MONO(2009)31) and OECD test guidelines 502. However, bifenthrin is not expected to be a predominant residue in/on sugar beet roots; sugar beet foliage, wheat grain and lettuce based on the results of the characterization/identification performed on wheat straw.

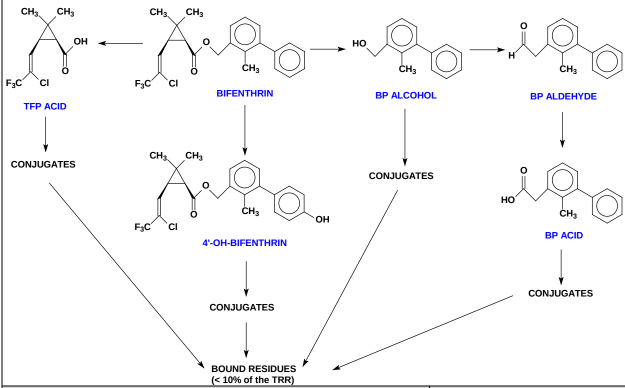
Radiolabel Position	n	[Cyclo	propyl (CP)	Ring]	[Phenyl (PH) Ring]			
PBI (months)		1	2	4	1	2	4	
Lettuce	Overall TRR (mg/kg)	0.014	0.029	0.017	0.012	0.021	0.014	
Sugar beet foliage	Overall TRR (mg/kg)	0.031	0.023	0.017	0.009	0.007	0.004	
Sugar beet roots	Overall TRR (mg/kg)	0.021	0.019	0.008	0.009	0.008	0.005	
Wheat grain	Overall TRR (mg/kg)	0.035	0.042	0.049	0.016	0.025	0.032	
Wheat straw	Overall TRR (mg/kg)	0.247	0.247	0.312	0.094	0.160	0.193	
Major Metabolites (> 10% of TRR)					Bifenthrin (2-month PBI only)			
Minor Metabolites (< 10% of TRR)		4'-OH-bifenthrin, TFP acid			4'-OH-bifenthrin, BP acid, BP alcohol, BP aldehyde			

CONFINED ACCUMULATION IN ROTATIONAL CROPS – PMRA # 1762330 Wheat

Sandy loam soil was treated at a rate of 0.56 kg a.i./ha. Seeds of wheat were sown at four plant-back intervals of 1, 4, 7 and 12 months. Samples of wheat straw and forage were analysed for identification/characterization of radioactive residues.

Radiolab	el Position	[0	Cyclopropyl	(CP) Ring	g] [Phenyl (PH) Ring]				
PBI (mor	nths)	1	4	7	12	2 1 4 7 1			12
Wheat forage Overall TRR (mg/kg)		0.283	0.116 0.089 0.034		0.192	0.054	0.010		
Major Metabolites (> 10% of TRR)		N/A	Bifenthrin (4-month PBI only); TFP acid			N/A	Bifenthrin (4 only); BP		N/A

NATUR	NATURE OF THE RESIDUE – Apples, Cotton, Corn PMRA # 1755373; 1755374; 1755375; 1755376; 1755377									
Minor M	etabolites (< 10% of TRR)	N/A Bifenthrin (7- and 12-mont PBIs); 4'-OH-bifenthrin				N/A	Bifenthrin (7 only); 4'-OH		N/A	
Wheat straw	Overall TRR (mg/kg)	0.371	0.371 0.335 0.151 0.17			0.326	0.100	0.047	0.077	
Major M	etabolites (> 10% of TRR)	Bifenthrin (1-month PBI only); TFP acid				Bifenthrin (1- and 7-month PBIs)				
Minor M	etabolites (< 10% of TRR)	Bifenthrin (4-, 7- and 12-month PBIs); 4'-OH-bifenthrin OH-bifenthrin; BP a								
Proposed	Proposed Metabolic Scheme in Plants (Primary and Secondary Crops)									
CH ₃	CH ₃ CH ₃ CH ₃ O									



NATURE OF THE RESIDUE – Laying Hens

PMRA # 1755380; 1755381; 1755382; 1755383

Forty laying hens were dosed orally with [\frac{14}{C}-cyclopropyl (CP) ring]-bifenthrin and [\frac{14}{C}-phenyl (PH) ring]-bifenthrin at doses of 4 mg a.i./bird/day (corresponding to 30.8-31.5 ppm in feed) by gelatin capsule once daily for 10 consecutive days. Samples of excreta were collected on Study Days 8, 9 and 10. Samples of eggs were collected once daily in the morning. The hens were euthanized within 24 hours after administration of the final dose.

www.mini.com	worming word of the film door.											
	[14C-Cyclop	oropyl (CP) Ring]	[¹⁴ C-Phenyl (PH) Ring]									
Matrices	TRR (mg/kg)	% of Administered Dose	TRR (mg/kg)	% of Administered Dose								
Excreta (Day 10)	55.0	Not reported	48.2	Not reported								
Muscle (adductor)	0.144	Not reported	0.102	Not reported								
Muscle (pectoral)	0.063	Not reported	0.035	Not reported								
Fat (abdominal)	2.09	Not reported	2.17	Not reported								
Liver	2.17	Not reported	1.36	Not reported								
Egg Yolk (Day 10)	3.20	Not reported	3.28	Not reported								
Egg White (Day 10)	0.042	Not reported	0.015	Not reported								

NATURE OF THE RES	IDUE – Apples, Cotton, Corn	PMRA # 1755373; 1755374; 1755375; 1755376; 1755377							
Metabolites identified	,	Minor Metabolites (<10% of the TRR)							
Radiolabel Position	[14C-CP Ring] [14C-PH Ring]	[14C-CP Ring]	[14C- PH Ring]						
Muscle (adductor)	Bifenthrin; combined OH-methyl bifenthrin and BP alcohol (PH label only)	4'-OH-bifenthrin; fatty acid conjugates of OH- methyl-bifenthrin; OH- methyl-bifenthrin	Fatty acid conjugates of OH-methyl-bifenthrin						
Fat (abdominal)	Bifenthrin; fatty acid conjugates of OH- methyl-bifenthrin	OH-methyl-bifenthrin; TFP acid; OH-methyl- TFP acid	OH-methyl-bifenthrin; BP alcohol						
Liver	Fatty acid conjugates of OH-methyl- bifenthrin; TFP acid (CP label only); OH-methyl-bifenthrin	OH-methyl-TFP acid; bifenthrin; TFP acid lactone	Bifenthrin; BP alcohol; 3'-4'-dimethoxy-BP alcohol; 4'-methoxy-BP alcohol; BP aldehyde; 3'-4'-dimethoxy-BP acid; 4'-methoxy-BP acid; BP acid						
Egg Yolk (Day 10)	Bifenthrin; fatty acid conjugates of OH- methyl-bifenthrin	OH-methyl-bifenthrin; TFP acid	OH-methyl-bifenthrin; BP alcohol; 4'-OH-BP acid; 4'-OH-BP alcohol						
NATURE OF THE RES	NATURE OF THE RESIDUE – Lactating Goats PMRA # 1755378; 1755379; 1755384; 1755385								

Four lactating goats were dosed orally with [14C-cyclopropyl (CP) ring]-bifenthrin and [14C-phenyl (PH) ring]-bifenthrin at doses of 2.3 mg/kg bw (corresponding to 79 ppm in feed) by gelatin capsule twice daily for seven consecutive days. Samples of excreta were collected once daily and milk was collected twice daily. The goats were

euthanized approximately 15 hours after administration of the final dose.

	[14C-Cyclopro]	pyl (CP) Ring]	[¹⁴ C-Phenyl (PH) Ring]			
Matrices	TRR (mg/kg)	% of Administered Dose	TRR (mg/kg)	% of Administered Dose		
Feces	Not reported	46	Not reported	48		
Urine	Not reported	12	Not reported	12		
Milk	Not reported	1.3	Not reported	1.2		
Fat (omental; perirenal; subcutaneous)	0.86-2.1	0.97	0.97-2.3	0.97		
Muscle (deltoid; flank; quadriceps)	0.25-0.38	0.42	0.24-0.29	0.38		
Heart muscle	0.58	< 0.2	0.48	< 0.2		
Kidney	0.39	< 0.2	0.77	< 0.2		
Liver	2.1	0.24	3.6	0.46		
Metabolites identified	Major Metabolites (Minor Metabolites (<10% of the TRRs)			
Radiolabel Position	[14C-CP Ring]	[14C- PH Ring]	[14C-CP Ring]	[¹⁴ C- PH Ring]		
Milk	Bifer	nthrin	OH-methyl-bifenthrin	4'-OH-bifenthrin; OH- methyl-bifenthrin; BP acid; BP alcohol		
Fat (perirenal)	Bifer	nthrin	4'-OH-bifenthrin; OH- methyl-bifenthrin; TFP acid	4'-OH-bifenthrin; OH- methyl-bifenthrin; BP		
Muscle (quadriceps)	Bifer	nthrin	OH-methyl-bifenthrin	4'-OH-bifenthrin; OH- methyl-bifenthrin; BP alcohol		
Heart muscle	Bifer	nthrin	4'-OH-bifenthrin; OH- methyl-bifenthrin; TFP acid	4'-OH-bifenthrin; OH- methyl-bifenthrin; BP alcohol		

NATURE OF THE RES	IDUE – Apples, Cotton,	Corn	PMRA # 1755373; 1755376; 1755377	; 1755374; 1755375;
Kidney	Bifenthrin	Bifenthrin; BP acid	4'-OH-bifenthrin; OH- methyl-bifenthrin; TFP acid; OH-methyl-TFP acid	BP alcohol; 4'-OH-BP alcohol
Liver		Di acid	OH-methyl-bifenthrin; TFP acid; OH-methyl- TFP acid	OH-methyl-bifenthrin; BP alcohol
Proposed Metabolic Scho	eme in Livestock			
CH ₃ CH ₃ O CH 4'-OH-BIFEN CH ₃ CONJUGATES 4'-OH-BP ALCO	BP AL OHOL	CH ₃ ENTHRIN CH ₃ CH ₃ ACID	CH ₃ CH ₂ OH O OH-METHYL-BI CH ₃ CH ₃ OH TFP ACID CONJUGATES	FATTY ACID CONJUGATES CH ₃ CH ₂ CH ₂ OH OH OH OH OH OH OH OH OH O
HO' CH ₃	ОН			

	OII BI AOID											
List of Metabolite	List of Metabolites Identified in Plant and Livestock Metabolism Studies											
Common name (Company code)	Chemical name	Chemical structure	Found in metabolism studies									
Bifenthrin (FMC 54800)	[2-methyl-(1,1'-biphenyl)-3-yl]-methyl-cis,trans-3-(2-chloro-3,3,3-trifluoro-1-propenyl)-2,2-dimethyl) cyclopropanecarboxylate	CI CH ₃ CH ₃ CH ₃	Apple; cotton; corn; wheat; hen; goat									
4'-OH-bifenthrin (FMC 78128)	3-(4'-hydroxyphenyl)-2- methylbenzyl(±)cis-3-(2- chloro-3,3,3-trifluoro-1- propenyl)-2,2- dimethylcyclopropane- carboxylate	F F O CH ₃	Corn; wheat; hen; goat									

NATURE OF TH	E RESIDUE – Apples, Cotton,	Corn PMRA # 175 1755376; 175	5373; 1755374; 1755375; 5377
Hydroxymethyl- bifenthrin (OH-methyl- bifenthrin; FMC 108561)	2-Methyl-[1,1'biphenyl]-3- yl)- methyl-cis-3-(2-chloro- 3,3,3- trifluoro-1-propenyl) trans-2- hydroxymethyl2- methylcyclopropane- carboxylate	HO——CH ₃ O H ₃ C	Hen; goat
TFP acid (FMC 53997)	cis, trans-3-(2-chloro-3,3,3- trifluoro-1-propenyl)-2,2- dimethyl- cyclopropanecarboxylic acid	F ₃ C CH ₃ CO ₂ H	Cotton; corn; wheat; hen; goat
Biphenyl acid (BP acid; FMC 65328)	2-methyl-3-phenylbenzoic acid	HO ₂ C CH ₃	Cotton; corn; wheat; hen; goat
Biphenyl alcohol (BP alcohol; FMC 56789)	2-methyl-3-phenylbenzyl alcohol	HOH ₂ C CH ₃	Cotton; corn; wheat; hen; goat
Biphenyl aldehyde (BP aldehyde)	2-methyl-3- phenylbenzylaldehyde	H CH ₃	corn

FREEZER STORAGE STABILITY – Plant Matrices PMRA # 1762217; 1762218; 1762212; 1762211; 1762215; 1762216; 1762219; 1762220; 1762221

Samples were stored at -38°C to 0°C. The data can be extended to crop field trials and processing studies to cover the maximum storage intervals observed.

Commodity Categories	Representative Commodities	Demonstrated Duration of Stability
	Head lettuce	183 days [6 months]
High weter content	Apple fruits	1490 days [49 months]
High water content	Field corn silage, stover	1490 days [49 months]
	Bananas	730 days [24 months]
High oil content	Cottonseed	730 days [24 months]
High oil content	Pecan	1095 days [36 months]
High protein content	Dried shelled peas	440 days [14.5 months]
High stouch content	Potato tubers	1095 days [36 months]
High starch content	Field corn grain	1034 days [34 months]
High acid content	Orange	548 days [18 months]
Processed fractions	Orange (dried pulp, juice, oil)	548 days [18 months]
riocessed fractions	Field corn (flour, meal, starch, refined oil)	365 days [12 months]

FREEZER STORAGE STABILITY – Animal Matrices PMRA # 1762220; 1762221

Samples were stored in freezer at -18°C. The demonstrated duration of stability covers the maximum storage interval observed in the livestock feeding studies.

Species	Commodity	Demonstrated Duration of Stability
Cattle	Fat	36 months
	Liver	36 months
	Muscle	36 months
	Milk	36 months
Poultry	Eggs	36 months

CROP FIELD	CROP FIELD TRIALS – Potatoes and Raspberries PMRA # 1762335; 1762236											
		D	OMEST	IC RE	GISTRAT	ION AND IME	ORTED (COMMODIT	IES			
Potatoes and ra								and corm	vegetables	(CSG 1C)	and	
caneberry (CS	G 13-()7A) are	petition	ed for			T					
Crops:	_				Potat			Raspberries and Blackberries				
Number of Tria					12			4 and 1				
Trial Locations	:		2 11	. ,		Conducted in					(4.07777)	
Formulation Ty	ype:		2 lb a.i./gal Emulsifiable Concentrate (2EC) and 1.15 lb a.i. Granular (1.15G)				10% a.i. by weight Wettable Powder (10WP) or Water-Soluble Bag (WSB)					
						row at plantin	_					
Application Ty	pe:			-	(2EC or	1.15 G)			directed sprand during ma			
			Seco	nd/thire		oadcast foliar	(2EC)	oloom ui			berries	
Adjuvant Use:					Nor				Non			
Residue Decline					Nor				ue-decline s			
X-fold Approve			(DN: 1.	8	US: 1.0		CDN:		US:	1.0	
Commodity]	al App. Rate a.i./ha]	PHI (days)	,		Bii	enthrin k	Residue Leve	els (ppm)			
	1-8	<u>-</u>		n	Min.	Max.	LAFT	HAFT	Median	Mean	SD	
Potatoes	(1.1	6-0.61 5 G and EEC)	21	24	< 0.05	< 0.05	< 0.05	< 0.05	0.05	0.05	0	
	0.56-0.61 (2EC)		21	24	< 0.05	< 0.05	< 0.05	< 0.05	0.05	0.05	0	
Raspberries	(W	0.224 VSP or OWP)	3	8	0.08	0.54	0.08	0.44	0.35	0.32	0.17	
Blackberries	0.224	4 (WSB)	2	2	0.68	0.80	0.74	0.74	0.74	NA	NA	
CROP FIELD beet; head let				radish	garden			1762248; 1 1828919;			762243;	
,			-		IMPOR'	гер Соммо		,	· · · · · · · · · · · · · · · · · · ·			
CG/CSG		1A/				4-13A	4-13A	22B				
Crops:		Carı	ot	Ra	dish	Garden Be	et Hea	d Lettuce	Spinacl	1	Celery	
Number of Tria	als:	10)		6	6		10	8		12	
Trial Locations	:			Tria	ls were co	onducted in ap	propriate l	NAFTA Gro	wing Region	ıs.		
Formulation Ty	ype:		2	lb a.i./g		fiable Concen a.i. by weigh				1.15G);		
Application Ty	pe:		foliar (directed or broadcast) applications to carrot and garden beet; 1 in-furrow at planting + 2 directed or broadcast) applications to radish; foliar (directed or broadcast) applications to head lettuce and celery (5-6 appl.); foliar ground spray to spinach (1 or 4 appl.)									
Adjuvant Use:							None					
Residue Decline Trend:	e	No 1	esidue d	lecline		ved in carrot r nrin residues in				ofile obser	ved for	
X-fold Register GAP	ed	1.0)	().4	1.0		1.0	1.0		1.0	
Total App.			PHI			Bi	fenthrin I	Residue Leve	els (ppm)	ls (ppm)		
Стор		Rate a.i./ha]	(days)) n	Min.	Max.	LAFT	HAFT	Median	Mean	SD	
Carrot roots	0.55	0-0.575	20-22	20	< 0.05	< 0.05	0.05	0.05	0.05	0.05	0	
Radish roots	0.22	1-0.241	6-8	12	< 0.05		0.05	0.06	0.05	0.05	0.01	
Radish tops Garden beet	0.44	8-0.460	1	12	< 0.05	2.25 5 0.28	0.63	0.28	0.06	0.11	0.58	
Jaruen beet	0.448-0.460		1	12	< 0.03	0.28	0.03	0.28	0.00	0.11	0.09	

NATURE OF	THE	RESIDU	JE – Ap	ple	es, C	Cotton, C	orn			MRA # 17: 55376; 17	55373; 175 55377	55374	; 175	5375;
roots														
Garden beet tops					12	4.80	12.2		5.0	11.9	6.35	7.2	29	2.55
Head lettuce	0.5	6-0.67	6-8		8	< 0.05	1.91		0.05	1.76	0.51	0.6	67	0.59
Spinach	0.44	8-0.467	36-41		14	< 0.05	0.16	(0.05	0.15	0.05	0.0)7	0.04
Celery	0.49	7-0.566	6-8		32	0.06	1.26	(0.10	1.16	0.52	0.5	57	0.35
CROP FIELD cauliflower; cand bean; suc shelled pea an	abbag culent	e; soybea t shelled	an; edil pea and	ole-	pod ean;	ded pea dried	PMRA 176231	7; 1′	76223		3; 1762264 5; 1762234			
aguaga		1 12	n				ED COMMO	DDIT	1	() /D				0.004
Crops:	Crops: Brassica Brassica Green		Leafy	a	nd S	a Head Stem	6 Soybeans	·	6A/B Edible-Podded and Succulent Shelled Pea and Bean		6C Dried Shelled Pea and Bean			8-09A omatoes
Number of Trials: 7		4 (broc 7 (cabb 4 (caulif	age)		(mu gree	istard ens)	15			for each nmodity	6 (pea) 9 (bean			16
Trial Locations	:			Τ	rials		ducted in ap					ıs.		
Formulation Ty Application Ty	Foliar (directed or broadcast) spray to <i>Brassica</i> head and stem vegetables (5-11 appl. to brocc													
Adjuvant Use:								No	one					
Residue Decline Trend:	2	No res	sidue dec	cline	was	s observed	in cabbage,	soyl	beans,	dried shelled	d peas and b	eans,	and to	omatoes.
X-fold Register GAP	ed	0.8-1	.1		1.	.0	1.0		1.0		1.0		0.8-1.0	
		al App.	PHI				Bi	Bifenthrin Residue Levels (ppm)						
Crop		Rate a.i./ha]	(days)		n	Min.	Max.	L	AFT HAFT		Median	Mean		SD
Mustard greens	_	5-0.479	6-7		16	0.05	2.05	(0.07	2.01	1.04	1.0)8	0.71
Broccoli	C	0.560	6-7		10	< 0.05	0.56	(0.06	0.44	0.16	0.1	19	0.16
Cauliflower	C	0.560	6-8		10	< 0.05	0.19	(0.05	0.18	0.07	0.1	10	0.05
Cabbage		0.560	6-7	_	5	0.44	3.09	(0.57	3.09	1.45	1.4	14	1.03
Soybean	0.33	4-0.336	17-18		28	< 0.05	0.18	(0.05	0.18	0.05	0.0)6	0.03
Edible-podded pea	C	0.224	3		11	0.16	0.50	(0.17	0.49	0.19	0.2	27	0.14
Edible-podded bean	C	0.224	2-4		12	< 0.05	0.15	(0.05	0.14	0.05	0.0)8	0.04
Succulent shelled pea	C	0.224	3		11	< 0.05	< 0.05	(0.05	0.05	0.05	0.0)5	0
Succulent shelled bean	0.22	3-0.225	2-4		14	< 0.05	< 0.05	(0.05	0.05	0.05	0.0)5	0
Dried shelled pea	C	0.224	14-15		12	< 0.05	< 0.05	(0.05	0.05	0.05	0.0)5	0
Dried shelled bean	C	0.336	13-15		18	< 0.05	0.10	(0.05	0.10	0.05	0.0)6	0.02
Tomato	0.34	7-0.364	0		4	< 0.05	0.10	-	0.07	0.09	0.08	0.0)8	0.02
Tomato 0.347-0		. 5.507	3		4	< 0.05	0.11	(0.06	0.10	0.08	0.0)8	0.03

NATURE OF T	THE I	RESIDUE	– App	les, C	Cotton, C	orn		PM	RA # 175	55373; 175	55374; 175	55375;
						ī			5376; 17:	55377		
			4	4	0.06	0.09	0.0	07	0.08	0.07	0.07	0.02
			5	20	< 0.05	0.09	0.0	05	0.09	0.06	0.06	0.01
			6	4	< 0.05	< 0.05	0.0	05	< 0.05	0.05	0.05	0
			7	4	< 0.05	0.10	0.0	07	0.08	0.07	0.07	0.02
			9	4	< 0.05	0.05	0.0	05	0.05	0.05	0.05	0
CROP FIELD	TRIA	LS – Rell	nennei	r. noi	nhell	PMRA	# 1762	246. 1	762247•	1762266;	1762228	1762229
pepper; eggplar										1762311;		
squash; pear; n				,						1828910;		
	·	,				182891			ĺ	<i>,</i>	<i>'</i>	ĺ
					IMPORTE	ED COM	MODITII	ES				
CG/CSG		8-09B/			9	11	-09		11-09		N/A	
Crops:		Pepper (I Nonbel Eggpla	l);		curbit etables	Pe	ear]	Mayhaw		Tea	
		5 (bell per	pper)		ntaloupe)	3 (at II	S GAP)					
Number of Trials	s:	7 (nonb		•	cumber)		t at US		3		3	
		pepper 3 (eggpla		,	ummer uash)	· G	AP)					
		3 (eggpit	1111 <i>)</i>	sq	uasii)	<u>. </u>				Trial	s conducted	l in India
Trial Locations:		Trials	were co	nducte	ed in appro	priate NA	AFTA Gr	owing	Regions.		ern and No	th-Eastern
											regions	
Formulation Typ	e:									t Wettable l		
Application Type										naw (2 appl. t spray to cu		
Application Type	:	broadcas	t) spray	s to ai		appl.); sp				t spray to ct	icuibit vege	tables (5
Adjuvant Use:						шрриј, гр	No		(1 upp11)			
Residue Decline		No mosiduo	daalina	muofil	a waa ahaa	amuad in a	a.uhit v	ra a a ta b	laa A maai	dua daalina	rrios obsomi	ad in maona
Trend:		No residue	decime	prom	e was obse	ervea in c	ucurbit v	egetab	ies. A resi	due decline		
X-fold Registered	i	0.55.1		1.0			2.0	0 1.0			0.38 (Japa	
GAP		0.75-1.	0	1.0		0.4-2.0		1.0		1	1.0 (Korea); 1.1 (China, Taiwan)	
	Tota	l App. Rate	PHI			<u> </u>	Rifenth	rin Re	sidue Lev		i (Cillia, 1	iiwaii)
Crop		g a.i./ha]	(days)	n	Min.	Max.	LAFT	HA			1 9	SD
Bell pepper		223-0.227	6-7	10	< 0.055	0.24	0.06	0.1				.06
Nonbell pepper	_	168-0.227	6-7	14	< 0.05	0.31	0.05	0.2				.08
Eggplant		0.224	7	6	< 0.05	< 0.05	0.05	0.0	5 0.03	5 0.05		0
Cantaloupe		0.336	3	14	< 0.10	0.35	0.10	0.3				.08
Cucumber		0.336	3	18	<0.10	0.24	0.10	0.2				.04
Summer squash	_	0.336	3	20	< 0.10	0.18	0.10	0.1	5 0.10	0.10	0	.02
		560-0.571 WP; HVS)	14	6	0.10	0.38	0.12	0.3	3 0.18	8 0.21	0	.10
Pear		560-0.571 WP; LVS)	14	6	0.12	0.43	0.21	0.3	5 0.25	5 0.28	0	.13
i cai		.673-1.12 (10WP)	14	20	0.07	0.039	0.08	0.3	7 0.20	6 0.25	0	.09
	0.	.673-1.12 (2EC)	14	6	0.07	0.56	0.09	0.5	0.20	0.28	0	.21
Mayhaw	0.3	224-0.227	28-29	6	0.24	0.78	0.26	0.7	5 0.39	9 0.47	0	.23
Tea (fresh)		0.060	7	9	0.66	5.05	0.81	4.8			2	.03
Tea (black)				9	0.39	5.85	0.40	5.7			2	.47
RESIDUE DAT									ARA # 24			
Seven field trials	s for l	oifenthrin o	n wint	er wh	eat as a ro	otational	crop we	ere coi	nducted in	n the Unite	d States	

Seven field trials for bifenthrin on winter wheat as a rotational crop were conducted in the United States encompassing NAFTA Growing Regions 4 [1 trial], 5 [4 trials] and 8 [2 trials] during the 1999-2000 growing season. Wheat was planted as a rotational crop 30-32 days after the last application of bifenthrin to primary crops (i.e. cotton, field corn, sweet corn). Samples of wheat forage, hay, straw and grain were collected at maturity.

** /					PMRA # 1 1755376; 1	1755373; 1 1755377	.755374; 1	755375;		
	Total	PBI		Residue Levels (ppm)						
Commodity	Application Rate (kg a.i./ha)	(days)	n	Min. #	Max. #	LAFT *	HAFT *	Median *	Mean *	SD *
Wheat forage			14	< 0.05	< 0.05	< 0.05	< 0.05	0.05	0.05	0
Wheat hay	0.559-0.560	30-32	14	< 0.05	< 0.05	< 0.05	< 0.05	0.05	0.05	0
Wheat straw	0.339-0.300		14	< 0.05	< 0.05	< 0.05	< 0.05	0.05	0.05	0
Wheat grain			14	< 0.05	< 0.05	< 0.05	< 0.05	0.05	0.05	0

^{*} Values based on total number of samples.

Based on the results of the field accumulation study, a plant-back interval of 30 days is required for wheat. All the other crops not appearing on the label will have a plant-back interval of 365 days.

PROCESSED FOOD AND FEED – Potatoes; PMRA #

PMRA # 1762320; 1762335; 1762317; 1762225; 1762340

Soybeans; Tomatoes; Pears

Processing studies were conducted using samples from crop field trials conducted in NAFTA growing regions. The at plant band application (i.e. first application) to potatoes was made with a granular (G) formulation. The ground or broadcast foliar spray applications were made with an emulsifiable concentrate (2EC) or a wettable powder (10WP) formulation. As per DIR98-02, Section 10.6.3, the residues in processed potato and soybean fractions are corrected for the degree of exaggeration: $Pf = [Residues in processed fraction \div Degree of exaggeration] \div LOQ in RAC.$

RAC	Processed Commodity	Total Rate (kg a.i./ha)	X-Fold GAP	PHI (days)	Average Residues (ppm)	Processing Factor (Pf)	RAC HAFT	Anticipated Residue (ppm)
	Tuber (RAC)				< 0.05			
	Granule				< 0.05	1		0.05
Potato	Chip	1.68	5	21	< 0.05	1	< 0.05	0.05
	Wet peel				< 0.05	1		0.05
	Dry peel				0.094	0.4		0.02
	RAC				< 0.05			
	Meal				< 0.05	1		0.18
Soybean	Hulls	0.78	2.3	18	0.07	0.6	0.18	0.11
	Refined oil				< 0.05	1		0.18
	AGFs				9.51	83		15
	RAC		0.6 (3EC)		0.075		0.10	
Tomato	Purée	0.361	0.8 (2EC) or 1 (WSB)		< 0.05	0.67	(at 3-day PHI)	0.07
	Paste		or r (wsb)		< 0.05	0.67		0.07
	RAC				0.633			
	Wet pomace (peeled)				1.776	2.81		0.97
	Wet pomace (ground)				9.223	14.6		5.0
Pear	Canned pear (peeled)	0.560	1	14	< 0.01	0.016	0.35	0.005
	Purée (peeled)				< 0.01	0.016		0.005
	Purée (ground)				0.014	0.022		0.008
	Nectar (peeled)				< 0.01	0.016		0.005
	Nectar (ground)				0.011	0.017		0.006

LIVESTOCK FEEDING – Dairy cattle

PMRA # 1762350; 1762353; 1762355; 1762356; 1792357; 1762346; 1762351

Lactating dairy cows were administered orally twice daily for 28 consecutive days with gelatin capsules containing bifenthrin at dose levels of 5 ppm (45-fold), 15 ppm (136-fold), and 50 ppm (455-fold) of the estimated dietary burden for dairy cattle. Only the values from the feeding level closest to the estimated dietary burden are reported. Some cattle matrices were also analysed for metabolites of bifenthrin. As these metabolites are not part of the residue definition, the results are not reported herein.

^{*} Values based on per-trial averages. LAFT = Lowest Average Field Trial, HAFT = Highest Average Field Trial, SD = Standard Deviation. For computation of the LAFT, HAFT, median, mean and standard deviation, values < LOQ are assumed to be at the LOQ. n = number of field trials.

NATURE OF TI	HE RESIDUE -		PMRA # 1755373; 1755374; 1755375; 1755376; 1755377				
Commodity	Feeding Level	Bifenthrin Residues (ppm)		Dietary Burden (DB) (ppm)		Anticipated Bifenthrin Residues (ppm)	
	(ppm)	mean	max	Beef cattle	mean	max	
Muscle (adductor)		< 0.10	< 0.10		0.0022	0.0022	
Muscle (pectoral)		< 0.10	< 0.10		0.0022	0.0022	
Muscle (cardial)		< 0.10	< 0.10		0.0022	0.0022	
Liver		< 0.10	< 0.10		0.0022	0.0022	
Kidney	5	< 0.10	< 0.10	0.11	0.0022	0.0022	
Fat (subcutaneous)		0.50	0.77		0.0109	0.0169	
Fat (peritoneal)		1.22	1.82		0.027	0.040	
Whole milk		0.08	0.12		0.0018	0.0026	
Milk fat		0.53	0.62		0.012	0.014	
LIVESTOCK FI	EEDING – Lay	ing hen		PMRA # 1762347;	1762348; 17623 1762352	349; 1762354;	

Laying hens were administered orally once daily for 28 consecutive days with gelatin capsules containing bifenthrin at dose levels of 1.96 ppb (0.05-fold), 21.0 ppb (0.5-fold) and 216 ppb (5.4-fold) of the estimated dietary burden for laying hens. Only the values from the feeding level closest to the estimated dietary burden are reported. Some poultry matrices were also analysed for metabolites of bifenthrin. As these metabolites are not part of the residue definition, the results are not reported herein.

Commodity	Feeding Level		Residues om)	Dietary Burden (DB) (ppm)	- ,	fenthrin Residues opm)
	(ppm)	mean	max	Laying hen	mean	max
Muscle (thigh)		< 0.02	< 0.02		< 0.02	< 0.02
Muscle (breast)		< 0.02	< 0.02		< 0.02	< 0.02
Fat (subcutaneous)	0.216	< 0.05	< 0.05	0.04	< 0.05	< 0.05
Liver		< 0.05	< 0.05		< 0.05	< 0.05
Eggs (Day 28)		< 0.01	< 0.01		< 0.01	< 0.01

Table 11 Food Residue Chemistry Overview of Metabolism Studies and Risk Assessment

	PLANT STU	DIES		
RESIDUE DEFINITION FOR ENF	ORCEMENT			
Primary crops (apples; cotton; corn		Bifenthrin		
Rotational crops (wheat)				
RESIDUE DEFINITION FOR RISI	K ASSESSMENT			
Primary crops (apples; cotton; corn		Bifen	thrin	
Rotational crops (wheat)				
METABOLIC PROFILE IN DIVE	RSE CROPS	Similar in apples, cot	ton, corn and wheat.	
	ANIMAL STU	DIES		
ANIMALS		Ruminant a	and Poultry	
RESIDUE DEFINITION FOR ENF	ORCEMENT	Bifenthrin		
RESIDUE DEFINITION FOR RIS	K ASSESSMENT	Bifenthrin		
METABOLIC PROFILE IN ANIM	ALS	The profile in similar in the investigated animals.		
(goat, hen, rat)				
FAT SOLUBLE RE	SIDUE	Yes		
DIETARY RISK FROM FOOD AN	ID WATER			
- M		ESTIMAT	ED RISK	
Refined chronic (cancer and non-	POPULATION	% of ACCEPTABLE D	OAILY INTAKE (ADI)	
cancer) dietary exposure analysis		Food Alone	Food and Water	
ADI = 0.003 mg/kg bw/day	All infants < 1 year	18.1	18.8	
ADI = 0.003 mg/kg bw/day	Children 1–2 years	33.6	33.9	
Estimated chronic drinking water	Children 3 to 5 years	28.9	29.2	
concentration = 0.29 µg a.i./L	Children 6–12 years	18.9	19.1	
υπετιστιστιστιστιστιστιστιστιστιστιστιστιστ	Youth 13-19 years	13.0	13.2	

	PLANT STUI	DIES		
	Adults 20–49 years	14.9	15.0	
	Adults 50+ years	15.2	15.4	
	Females 13-49 years	14.1	14.3	
	Total population	16.5	16.7	
		ESTIMAT	·-	
Refined acute dietary exposure	POPULATION	% of ACUTE REFERENCE DOSE (ARfD		
analysis, 95 th percentile		Food Alone	Food and Water	
	All infants < 1 year	41.9	47.1	
ARfD = 0.009 mg/kg bw	Children 1–2 years	71.8	75.2	
Estimated couts deinbing water	Children 3 to 5 years	59.5	61.4	
Estimated acute drinking water concentration = 1.5 µg a.i./L	Children 6–12 years	38.6	39.6	
concentration – 1.5 μg a.i./L	Youth 13-19 years	27.9	29.0	
Monitoring data (highest detection)	Adults 20–49 years	28.2	29.6	
= 5.2 µg a.i./L (Note: this value was	Adults 50+ years	27.7	28.9	
used in estimated risk)	Females 13-49 years	25.7	27.0	
	Total population	33.8	35.4	

Table 12 Fate and Behaviour in the Environment

Study	Test substance	Value	Comments	Reference
Abiotic transformation	1			
Hydrolysis (25°C)	Bifenthrin	Stable	Not expected to be a route of dissipation	1924822
Phototransformation on soil (20°C)	Bifenthrin	Based on 12 hours of daylight DT ₅₀ : 115 d, DT ₉₀ : 382 d	Not expected to be a route of dissipation (half-life >7 days)	1755325
Phototransformation in water (25°C)	Bifenthrin	Based on 12 hours of daylight DT ₅₀ : 31.6 to 51.8 d, DT ₉₀ : 105-173 d	Not expected to be a route of dissipation (half-life >7 days)	1924824
air	potential to vo bifenthrin rele However, giv	ressure and Henry's law constant of bifer blatilize from water and moist soil. The A cased to the air would have an atmospher en the adsorption properties of bifenthring of dissipation in air.	Atkinson method predict ric half-life less than one	ed that any day.
Biotransformation	1			_
Biotransformation in	Bifenthrin	Bifenthrin (combined labels)		1755314
aerobic soil (20°C)		Hagertown silt clay $DT_{50} = 112 \text{ d}, DT_{90} = 371 \text{ d}$	Persistent	1755315 1755318
		Cosad sandy loam $DT_{50} = 89.4 \text{ d}, DT_{90} = 1399 \text{ d}$	Moderately persistent	1755319 1755321
		Dunkirk silt loam $DT_{50} = 203 \text{ d}, DT_{90} = 674 \text{ d}$	Persistent	1755322
		Georgetown silt loam $DT_{50} = 78.7 \text{ d}, DT_{90} = 261 \text{ d}$	Moderately persistent	
		Arlington DT ₅₀ for <i>R-cis</i> : 277days DT ₅₀ for <i>S-cis</i> : 330 days	Persistent	Qin <i>et al</i> 2006
Biotransformation in	Bifenthrin	$DT_{50} > 1000$	Persistent	1924829
anaerobic soil		Arlington DT_{50} for R - cis : 770 days DT_{50} for S - cis : 495 days	Persistent	Qin <i>et al</i> 2006

Study	Test	Value	Comments	Reference
<u> </u>	substance			
Biotransformation in	Bifenthrin	Calwich lake	Persistent	1755309
aerobic water-sediment		Water		
systems (20°C)		$DT_{50} = 1.86 \text{ d}, DT_{90} = 6.16 \text{ d}$		
		Whole system		
		$DT_{50} = 276 \text{ d}, DT_{90} = 918 \text{ d}$		
		Swiss lake	Moderately persistent	
		Water	7 1	
		$DT_{50} = 1.7 \text{ d}, DT_{90} = 5.66 \text{ d}$		
		Whole system		
		$DT_{50} = 92.9 \text{ d}, DT_{90} = 308 \text{ d}$		
Biotransformation in	Study not requ	aired as the anaerobic (flooded) soil stud	v addressed this	1755308
	requirement.	ince as the unacrosse (modeca) son stac	y addressed tims	1,0000
sediment systems	requirement.			
Mobility	<u> </u>			
Adsorption / desorption	Rifenthrin	Leon fine sand	Immobile	1755306
in soil	Висисии	$K_d = 2005 \text{ mL/g}, K_{OC} = 115245 \text{ mL/g}$	minoone	1924831
111 5011		Cosad sandy loam		1921031
		$K_d = 2685 \text{ mL/g}, K_{OC} = 154299 \text{ mL/g}$		
		Dunkirk silt loam		
		$K_d = 1800 \text{ mL/g}, K_{OC} = 103474 \text{ mL/g}$		
		Hagerstown clay loam		
	TED 11	$K_d = 453 \text{ mL/g}, K_{OC} = 26013 \text{ mL/g}$	N/ 1 . 1 1 1 1 1	1755205
	TFP acid	M sandy loam	Moderately mobile	1755305
		$K_d = 118 \text{ mL/g}, K_{OC} = 411 \text{ mL/g}$		4
		J1 loamy clay	Very highly mobile	
		$K_d = 0.742 \text{ mL/g}, K_{OC} = 15.6 \text{ mL/g}$		
		A1 sand	Moderately mobile	
		$K_d = 5.49 \text{ mL/g}, K_{OC} = 499 \text{ mL/g}$		
		E3 silty loam	Moderately mobile	
		$K_d = 7.94 \text{ mL/g}, K_{OC} = 241 \text{ mL/g}$		
		Horn sandy loam	Very highly mobile	
		$K_d = 0.573 \text{ mL/g}, K_{OC} = 24.6 \text{ mL/g}$		
	4'-OH	K _{OC} : 3,043-397,253 mL/g	Low mobility to	2533219
	bifenthrin		immobile	
Volatilization from soil	Bifenthrin	< 2% TAR of bifenthrin detected in	No classification	1755299
		volatile traps.		1755300
Bioconcentration/Bioac	ccumulation	1		
	Bifenthrin	Carp:		1755224
fish		BCF _{ss} : 709 – 1170		
		BCF _K : 815-1200		
		BCF _{K.G} : 809-1191		
				1755215
				1733213
				1755010
				1755218
				1755216
1				
		BCF _K : 12850		
		$\begin{array}{l} BCF_{K,G,L}\colon 1265\text{ - }1861\\ Bluegill \ sunfish:\\ BCF_{SS}\colon 1584\text{-}1649\\ BCF_{SS}\colon 5\% \ lipid \ normalized: 2507\text{ - }2820\\ BCF_{K}\colon 2117\text{-}2147\\ BCF_{K,G}\colon 2251\text{-}2325\\ BCF_{K,G,L}\colon 3400\text{-}3511\\ Bluegill \ sunfish:\\ Based \ on \ measured \ bifenthrin \ in \ water:\\ BCF_{SS}\colon 6090\\ BCF_{K}\colon 12850\\ \end{array}$		1755

Study	Test substance	Value	Comments	Reference
		Based on nominal bifenthrin in water: BCF _{SS} : 2107 BCF _K : 5250		
		Fathead minnow: (not corrected for growth dilution or lipid content) F ₀ adult BCF _{SS} : 21,000-30,000		1755227
Bioconcentration in <i>D.</i> magna	Bifenthrin	$BCF_{K} = 4750$ $BCF_{K,G} = 6273$		2533225
Dietary biomagnification in <i>D. magna</i>	Bifenthrin	$\begin{split} BMF_{SS} &= 0.11 \\ BMF_{K} &= 0.11 \\ BMF_{K,G} &: 0.11 \\ BMF_{K,G,L} &: 0.11 \end{split}$	The study indicates that bifenthrin did not biomagnify in this study as the reported BMFs are <1.0.	2533226
Dietary biomagnification in bluegill sunfish	Bifenthrin	Bluegill sunfish: $BMF_{K,=} 0.08$ $BMF_{K,G,=} 0.13$ $BMF_{K,G,L} = 0.28$	The study indicates that bifenthrin did not biomagnify in this study as the reported BMFs are <1.0.	2533236
Bioaccumulation in biota sampled in the aquatic field	Capture 2EC (240 g bifenthrin/L formulation)	Estimated ranged of BAF:Catfish: 134 – 5385 Channel catfish: 77 – 12682 Gizzard shad: 499 – 12458 Threadfin shad: 182 – 1855 Redear sunfish: 51 – 3844 Spotted sucker: 535 – 11564 Bluegill sunfish: 11 – 7430 White crappie: 11 – 3430 Largemouth bass: 116 - 8715		1755966 1762382
Food web bioaccumulation modelling and risk assessment	Modelling	The risk via bioaccumulation was assessed with new data and modelling by EFSA. A food web bioaccumulation model, evaluated against field data for fish measured in agricultural settings after extensive bifenthrin applications, was applied using evaluative bioaccumulation and exposure assumptions. Time-dependent bioaccumulation and exposure calculations were determined by linking the output from a five-year FOCUS model scenario as input for the food web predictions. Bifenthrin is found in each level of the	A summary of the information reviewed by European Commission indicates that although bifenthrin is found in each level of the food chain, no biomagnification has been observed. (EC, 2010) However, concerns raised by the PRAPeR 87 experts' meeting indicated that the high risk from bioaccumulation through the food chain for aquatic organisms could not be excluded (EFSA, 2011).	2422597

Study	Test substance	Value		Comments	Reference
		the trigger when all acute fish toxicity endpoints were compared to estimates of aggregate fish residues based on combining all potential routes of exposure including dietary and environmental water sources.			
Field studies					
Field dissipation in an ecoregions not representative of Canadian conditions (Supplemental studies)	Capture 2EC (240 g bifenthrin/L formulation)	US Champaign Illinois	DT ₅₀ : 215 d DT ₉₀ : 714 d	Persistent	1762370
(Supplemental studies)	Brigade 10WP (10% bifenthrin w/w)		DT ₅₀ : 86.7 d DT ₉₀ : 288 d	Moderately persistent	1762369
	Talstar SC (7.8% bifenthrin w/w)	European: Goch Germany	DT ₅₀ : 207 d DT ₉₀ : 687 d	Persistent	1755908
	,	Meynes France	DT ₅₀ : 153 d DT ₉₀ : 507 d	Persistent	
		Budrio Italy	DT ₅₀ : 80 d DT ₉₀ : 264 d	Moderately persistent	
Terrestrial field dissipation study in Canada	Bridgade 2EC (25.1% bifenthrin w/w)	Prince Edward Island	DT ₅₀ : 108 days DT ₉₀ : 288 days	Moderately persistent This study is classified as unreliable and does not fulfill the data requirement.	2533221
Field dissipation – cottonfield and outdoor pond study	Capture 2EC (240 g bifenthrin/L formulation)	US Dallas, Alabama	Cotton field DT ₅₀ : 197 d DT ₉₀ : 645 d Pont water DT ₅₀ : 609 days DT ₉₀ : not calculated	Persistent	1755966 1762383 1762384 1762387 1762385 1762380

Table 13 Name and chemical structure of environmental transformation products of bifenthrin

Code Name/ Synonym	Chemical Name	Chemical Structure
	Major transformation product (>10%)
aqueous photolysis study	cyclopropanecarboxylic acid, 3-(2-chloro-3,3,3-trifluoro-1-propenyl)-2,2-dimethyl-, (2-methyl[1,1'-biphenyl]-3-yl)methyl ester, trans- (9Cl)	F F O CH ₃

	<u> </u>	OH
Biphenyl alcohol	2-methyl-3-phenylbenzyl alcohol	OH /
BP-alcohol in		⟨ CH₃
	2-methyl-3-biphenyl methanol	\rightarrow
study		()—()
4'-hydroxy	(4'-hydroxy-2-methyl-3-biphenyl)methyl	F, F
bifenthrin	(1RS, 3RS)-3-[(1Z)-2-chloro-3,3,3-trifluoro-	F CI CH ₃ OH
4'-OH bifenthrin	1-propen-1-yl]-2,2-	
in laboratory	dimethylcyclopropanecarboxylate	
aerobic water-		H ₃ C
sediment study		H-C
		E F
		F.X
		CI CH ₃
		OH OH
		H ₃ C O
TED soid (Cir. or. 1	(1DC 2DC) 2 [(17) 2 allows 2 2 2 millions	H₃ć HO
TFP acid (Cis and	(1RS, 3RS)-3-[(1Z)-2-chloro-3,3,3-trifluoro-	F CI \ _
Trans) in aquatic photolysis study	1-propen-1-yl]-2,2- dimethylcyclopropanecarboxylic acid	— 0
photorysis study		
		F —
		H₃C CH₃
		F CI
		[\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \
		F—
		F \\
		\times
		H ₃ C´ CH ₃
	Minor transformation products	
Hydroxyl-methyl	(2-methyl-3-biphenyl)methyl (1RS, 3RS)-3-	F, F
bifenthrin	[(1Z)-2-chloro-3,3,3-trifuoro-1-propen-1-yl]-	F CI CH ₃
OH-methyl	2-(hydroxymethyl)-2-	
bifenthrin	methylcyclopropanecarboxylate	
		H ₃ C \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \
Rinhenyl aldehyde	2-methyl-3-phenylbenzyl aldehyde	О
BP-aldehyde	2 monty 5 phonytoenzy andenyde	//
	2-methyl-3-biphenylcarbaldehyde	CH ₃
	, , , , , , , , , , , , , , , , , , ,	
		\ \ \ \ \ \ \

Code Name/ Synonym	Chemical Name	Chemical Structure
Biphenyl acid BP-acid	2-methyl-3-phenylbenzoic acid 2-methyl-3-biphenylcarboxylic acid	HO—CH ₃

Table 14 Summary of formation of transformation products (% applied radioactivity) formed in the submitted environmental studies

Study type		Maximum % AR (day)	Final %AR (study length)	Reference		
Trans bifent	thrin			, , , , , , , , , , , , , , , , , , , ,	(Starting Language)	
Soil photolys		Cyclopro	pyl label	2.3 (30)	2.3 (30)	1755325
1 3		Phenyl la	* *	3.1 (30)	3.1 (30)	1
Aqueous pho	otolysis	Cyclopro		36.7 (262)	36.7 (262)	1924824
1	3	Phenyl la		48.9 (165)	30.4 (310)	1
BP-alcohol				, ,	, ,	
Soil photolys	sis	Phenyl la	ibel	1.6 (30)	1.6 (30)	1755325
Aqueous pho				19.0 (211)	15.2 (310)	1924824
Aerobic soil				0.2% (120)	0.2% (120)	1755314
	Sandy loam			0.4% (120)	0.4% (120)	1755318
	Silt loam			0.4% (120)	0.4% (120)	1755319
	(Dunkirk)	_				1755321
	Silt loam			0.4% (62-90)	0.2% (126)	1755315
	(Georgetown)			, ,	, ,	1755322
Aerobic	Calwich	Phenyl	Water	1.1% (2)	NA (99)	1755309
water-		label	Sediment	0.6% (14-30)	0.1% (99)	7
sediment	Swiss lake		Water	1.3% (0)	NA (99)	1
			Sediment	1.1 (99)	1.1 (99)	7
4'-OH bifen	thrin			(-2)	()	
Soil photolys		Cyclopro	pyl label	0.5 (14)	0.3 (30)	1755325
1 3		Phenyl la		0.6 (14)	0.5 (30)	
Aerobic soil	Silty clay	Cyclopro	pyl label	1.0% (180)	1.0% (180)	1755314
		Phenyl la		3.3% (120)	3.3% (120)	1755318
	Sandy loam	Cyclopro	pyl label	5.0% (180)	5.0% (180)	1755319
		Phenyl la		4.1% (120)	4.1% (120)	1755321
	Silt loam	Cyclopro	pyl label	3.7% (180)	3.7% (180)	
	(Dunkirk)	Phenyl la	ibel	8.2% (120)	8.2% (120)	7
	Silt loam	Cyclopro	pyl label	3.8% (30)	2.5% (126)	1755315
	(Georgetown)	Phenyl la	ibel	3.6% (30)	3.6% (30)	1755322
Aerobic	Calwich	Cyclo-	Water	5.4% (0)	0 (1)	1755309
water- sediment		propyl Label	Sediment	4.4% (99)	4.4% (99)	
		Phenyl	Water	1.9% (9)	NA (99)	7
		Label	Sediment	5.6% (99)	5.6% (99)	7
	Swiss Lake	Cyclo-	Water	0.2% (14-30)	NA (99)	7

Study type			Maximum % AR (day)	Final %AR (study length)	Reference	
		propyl	Sediment	9.2% (99)	9.2% (99)	
		Label				
		Phenyl	Water	0.3% (0)	NA (99)	7
		Label	Sediment	11.1% (99)	11.1% (99)	
TFP acid	•	l .	- 1		-	
Soil photo	lysis	Cyclopro	pyl label	3.8 (30)	3.8 (30)	1755325
Aqueous p	hotolysis			12.2 (262)	10.2 (262)	1924824
Aerobic	Silty clay			3.7% (180)	3.7% (180)	1755314
soil	Sandy loam			0.2% (180)	0.2% (180)	1755318
	Silt loam			1.6% (180)	1.6% (180)	1755319
(Dunkirk)						1755321
	Silt loam			0.8% (62)	0.5% (126)	1755315
	(Georgetown)					1755322
Aerobic	Calwich	Cyclo-	Water	0.8% (30)	NA (99)	1755309
water-		propyl	Sediment	0.8% (14)	0.6% (99)	
sediment	Swiss lake	Label	Water	1.5% (30)	NA (99)	
			Sediment	0.9% (60)	0.3% (99)	
BP-aldehy						
Soil photo	•			1.3 (30)	1.3 (30)	1755325
Aerobic	Silty clay	Phenyl label		0.2% (120)	0.2% (120)	1755314
soil						1755318
						1755319
						1755321
BP-acid				T	1	1
Soil photo	•			1.4 (14)	1.4 (30)	1755325
Aerobic	Silty clay	Phenyl la	ıbel	0.6% (120)	0.6% (120)	1755314
soil	Sandy loam			1.7% (120)	1.7% (120)	1755318
	Silt loam			0.5% (120)	0.5% (120)	1755319
	(Dunkirk)					1755321
	Silt loam			0.7% (126)	0.7% (126)	1755315
	(Georgetown)		1			1755322
Aerobic	Calwich	Phenyl	Water	1.5% (28)	NA (99)	1755309
water-		label	Sediment	0.9 (30)	0.0 (99)	
sediment	Swiss lake		Water	1.4% (14)	NA (99)	
			Sediment	0.9% (30)	0.0 (99)	
	d radioactivity					
NA not an	alyzed					

Table 15 Toxicity of bifenthrin, a formulated product and transformation products to Non-Target terrestrial Species

Organism	Exposure	Test substance	Endpoint value	Degree of toxicity	Reference
Invertebrates					
Earthworm (E. fetida)	14-day	Bifenthrin	1- 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 -	No classification	1755289
		Talstar 8 SC	14-d NOEC = 7.89 mg a.i./kg dw soil	No	1755918

Organism	Exposure	Test substance		Degree of toxicity	Reference	
			14-d LOEC = 14.0 mg a.i./kg dw soil	classification		
			14-d LC ₅₀ >78.9 mg a.i./kg dw soil			
	56-day	Talstar 8 SC	NOEC = 2.13 mg a.i./kg dw soil	No	1755920	
	reproduction		LOEC = 4.26 mg a.i./kg dw soil	classification		
		4'-OH	NOEC = 178 mg TP/kg dw soil	No	1755186	
		bifenthrin	LOEC = 316 mg TP/kg dw soil	classification		
		TFP acid	NOEC = 17.8 mg TP/kg dw soil	No		
		111 4014	LOEC = 31.6 mg TP/kg dw soil	classification		
Bee	Acute Oral	Talstar 8 SC	NOEC < 0.09 µg a.i./bee	Highly toxic	1755922	
Всс	ricute oran	Taistai o Be	LD ₅₀ =0.13 μg a.i./bee	inginy toxic	1733722	
	Acute		NOEC = $0.05 \mu g \text{ a.i./bee}$	Highly toxic		
	Contact		LD ₅₀ =0.07 µg a.i./bee	inginy toxic		
	Field aged	Capture 2		No	1755286	
	residue	EC		classification	1733280	
	residue	EC	3.2-4.2 DAA for 112 g a.i./ha	Classification		
[A == 4 == 1.1	T-1-4 0.00	>5.2 DAA for 224 g a.i./ha	NI.	1755017	
Ladybird beetle	Agea residue	1 aistar 8 SC	% mortality in the 7.87 g a.i./ha × 2	No	1755917	
(C. septempunctat			treatment group was not statistically	classification		
L)			different from the control when			
			exposed to 21-day old residue.			
			% Mortality in the 50 g a.i./ha \times 2			
			treatment group was not statistically			
			different from the control when			
			exposed to 35-day aged residue.			
	Extended		Exposure to treated apple leaves	No	1755925	
	laboratory		$LR_{50} = 0.084 \text{ g a.i./ha} (95\% \text{ CI } 0.055\text{-}$	classification		
	study		0.105 g a.i./ha)			
A. rhopalosiphi	Acute contact	Talstar 8 SC	24 hours old adults exposed to glass	No	1755919	
			plates or maize leaves treated with 7.5	classification		
			g a.i./ha had 100% mortality			
			2.2.1			
			2-3 days old mummified aphids treated			
			with 7.5 g a.i./ha had comparable			
			emergence rate and reproductive			
		_	success as the control.			
	Extended			No	1755926	
	laboratory		, ,	classification		
	study]	10.805 g a.i./ha)			
	Aged residue		% mortality in the 1.6 g a.i./ha × 2	No	1755921	
			treatment group was not statistically	classification	1755927	
			different from the control when			
			exposed to 7-day old residue.			
			O/ Manufalitaria da C 1 a ci (ha c 2			
			% Mortality in the 6.1 g a.i./ha \times 2			
			treatment group was not statistically			
			different from the control when			
			exposed to 14-day aged residue.			
			% Mortality in the 50 g a.i./ha × 2			
			treatment group was not statistically			
			different from the control when			
			exposed to 28-day aged residue.			
Ground beetle	Extended	Taletar & SC	NOER based on mortality and food	No	1755923	
	LAWINGU	Taistai o SC	process based on mortality and 1000		1133743	
(P. cupreus L)	laboratory		consumption was 0.1066 mg a.i./kg	classification		

Organism	Exposure	Test substance	Endpoint value	Degree of toxicity	Reference
	Limit test				
Green lacewing	Extended	Talstar 8 SC	Exposure to treated bean leaves	No	1755924
(C. carnea)	laboratory		NOER = 2.279 g a.i./ha	classification	
	study		LR ₅₀ = 5.132 g a.i./ha (95% CI 1.952-		
			7.151 g a.i./ha)		
Predatory mite	Extended	Talstar 8 SC	Exposure to treated apple leaves	No	1755930
(T. pyri)	laboratory		NOER = 0.0009 g a.i./ha	classification	
	study		$LR_{50} = 0.113 \text{ g a.i./ha} (95\% \text{ CI } 0.091 -$		
			0.143 g a.i./ha)		
Birds					
Bobwhite quail	Acute oral	Bifenthrin	NOEL = 464 mg a.i./kg bw	Slightly toxic	1755213
			$LD_{50} = 1800 \text{ mg a.i./kg bw}$		
	5-d Dietary		NOEC = 2500 mg a.i./kg diet	Slightly toxic	1755205
			(NOEL = 393 mg a.i./kg bw/day)		
			$LC_{50} = 4450 \text{ mg a.i./kg diet}$		
			$(LD_{50} = 597 \text{ mg a.i./kg bw/day})$		
	24-week		NOEC = 75 mg a.i./kg dw diet	No	1755200
	Reproduction			classification	
Mallard duck	Acute oral		NOEL = 2150 mg a.i./kg bw	Practically	1755212
			$LD_{50} \ge 2150 \text{ mg a.i./kg bw}$	non-toxic	
	5-d Dietary		NOEC <312 mg a.i./kg diet	Slightly toxic	1755203
			(NOEL < 104 mg a.i./kg bw/day)		
			$LC_{50} = 1222 \text{ mg a.i./kg diet}$		
			LD ₅₀ was not calculated because of		
			inconsistent food consumption		
	22-week		NOEC = 75 mg a.i./kg dw diet	No	1755199
	Reproduction			classification	
Mammals	Ta	lmia i i	V.D. (050) GY	Ixx. 11 ·	Lagrani
Mouse	Acute oral	Bifenthrin	LD ₅₀ (95% CI):	Highly toxic	1755501
Swiss Webster			3 = 43.5 mg/kg bw (36.2-50.7 mg/kg)		
			bw)		
			$\bigcirc = 42.5 \text{ mg/kg bw } (37.1-47.9 \text{ mg/kg})$		
			bw)		
Dat	Multi-	Bifenthrin	Combined = 43.0 mg/kg bw	No	1755440
Rat		Bilenthrin	Repro LOAEL = 5 mg/kg bw/day	No classification	1755448
Sprague-Dawley	generation		Repro NOAEL = 3 mg/kg bw/day	ciassification	
Vacanlar la4-	Reproduction				1
Vascular plants	214 Candline	Tolotor 9 CC	NOEC = 0.09 g a i /lra dur sail	No	1755192
Vascular plant	_	Taistar 8 SC	NOEC = 0.08 g a.i./kg dw soil	No classification	1/33192
	emergence	1		crassification	

Table 16 Screening level risk assessment for bifenthrin and transformation products on non-target terrestrial species other than birds and mammals

Organism	Exposure	Endpoint value	EEC	RQ	Level of
					concern
Invertebrates					
Earthworm	Acute	$LC_{50}/2 = 9.5 \text{ mg a.i./kg soil}$	0.150 mg a.i./kg dw	0.02	Not
			soil		exceeded
	Chronic	NOEC = 2.13 mg a.i./kg dw	0.150 mg a.i./kg dw	0.07	Not
		soil	soil		exceeded
Bee	Acute oral	$LD_{50} = 0.13 \mu g a.i./bee$	0.269 μg a.i./bee	2.1	Exceeded
1	Acute	$LD_{50} = 0.07 \mu g a.i./bee$	6.96 µg a.i./bee	99	Exceeded

Organism	Exposure	Endpoint value	EEC	RQ	Level of concern
	contact				
A. rhopalosiphi	Acute	LR ₅₀ < 7.5 g a.i./ha	In-field:	In-field:	Exceeded
(aphid parasitoid)	contact		126 g a.i./ha	>17	
Foliar dwelling parasite			Off-field:	Off-	Exceeded
			74.3 g a.i./ha	field: >10	
C. septempunctat L.	Extended	$LR_{50} = 0.084 \text{ g a.i./ha}$	In-field:	In-field:	Exceeded
(ladybird beetle)	laboratory		126 g a.i./ha	1500	
Foliar dwelling predator	study		Off-field:	Off-	Exceeded
	Contact		74.3 g a.i./ha	field: 885	
A. rhopalosiphi		$LR_{50} = 8.145 \text{ g a.i./ha}$	In-field:	In-field:	Exceeded
(aphid parasitoid)		50 8	126 g a.i./ha	15	
Foliar dwelling parasite			Off-field:	Off-	Exceeded
21			74.3 g a.i./ha	field:	
P. cupreus L		NOER = 0.1066 mg a.i./kg	In-field:	In-field:	Not
(carabid beetle)		dw soil	0.09 mg a.i./kg dw soil	0.84	exceeded
Soil dwelling predator					
C. carnea		$LR_{50} = 5.132 \text{ g a.i./ha}$	In-field:	In-field:	Exceeded
(green lacewing)		50 2	126 g a.i./ha	25	
Foliar dwelling predator			Off-field:	Off-	Exceeded
21			74.3 g a.i./ha	field: 14	
T. pyri		$LR_{50} = 0.113 \text{ g a.i./ha}$	In-field:	In-field:	Exceeded
(predatory mite)			126 g a.i./ha	1115	
Foliar dwelling predator			Off-field:	Off-	Exceeded
			74.3 g a.i./ha	field: 658	
Vascular plants	•			•	
Vascular plant	Seedling	NOEC = 0.08 mg a.i./kg dw	Potato: 0.150 mg	< 1.9	Exceeded*
_	emergence	soil	a.i./kg dw soil		
		EC25 > 0.08 mg a.i./kg dw		< 1.175	
		soil	Raspberry: 0.094 mg a.i./kg dw soil		

^{*} Even though the LOC is slightly exceeded, effects on terrestrial plants are not expected given that no adverse effects were observed in six species of non-target terrestrial plants at the highest dose tested.

Table 17 Refined risk assessments on terrestrial non-target arthropods

Organism	Endpoint value	EEC	RQ	Level of concern
C. septempunctat L.	$LR_{50} = 0.084 \text{ g a.i./ha}$	In-field: 101 g a.i./ha	In-field: 1202	Exceeded
(ladybird beetle)		Off field: 7.43 g a.i./ha	Off-field: 88.5	Exceeded
Foliar dwelling predator				
A. rhopalosiphi	$LR_{50} = 8.145 \text{ g a.i./ha}$	In-field: 101 g a.i./ha	In-field: 12	Exceeded
(aphid parasitoid)		Off-field: 7.43 g a.i./ha	Off-field: 0.9	Not exceeded
Foliar dwelling parasite		_		
C. carnea	$LR_{50} = 5.132 \text{ g a.i./ha}$	In-field: 101 g a.i./ha	In-field: 20	Exceeded
(green lacewing)		Off-field: 7.43 g a.i./ha	Off-field: 1.4	Exceeded
Foliar dwelling predator				
T. pyri	$LR_{50} = 0.113 \text{ g a.i./ha}$	In-field: 101 g a.i./ha	In-field: 894	Exceeded
(predatory mite)		Off-field: 7.43 g a.i./ha	Off-field: 7.43	Exceeded
Foliar dwelling predator				

Table 18 Screening level risk assessment on birds and mammals

Organism	Toxicity (mg a.i./kg bw/d) ¹	Feeding Guild (food item)	EDE (mg a.i./kg bw) ²	RQ	Level of Concern
Small Bird (0.0	02 kg)				
Acute	180	Insectivore (small insects)	6.35	0.04	Not exceeded
Reproduction	6.71	Insectivore (small insects)	6.35	0.95	Not exceeded
Medium-Sized	Bird (0.1 kg)				
Acute	180	Insectivore (small insects)	4.95	0.03	Not exceeded
Reproduction	6.71	Insectivore (small insects)	4.95	0.74	Not exceeded
Large-Sized B	ird (1 kg)				
Acute	180	Herbivore (short grass)	5.17	0.03	Not exceeded
Reproduction	6.71	Herbivore (short grass)	5.17	0.77	Not exceeded
Small Mamma	l (0.015 kg)				
Acute	4.30	Insectivore (small insects)	3.65	0.85	Not exceeded
Reproduction	3	Insectivore (small insects)	3.65	1.22	Exceeded
Medium-Sized	Mammal (0.035 kg)				
Acute	4.3	Herbivore (short grass)	11.44	2.66	Exceeded
Reproduction	3	Herbivore (short grass)	11.44	3.81	Exceeded
Large-Sized M	lammal (1 kg)				
Acute	4.3	Herbivore (short grass)	6.11	1.42	Exceeded
Reproduction	3	Herbivore (short grass)	6.11	2.04	Exceeded

¹ Toxicity values for the acute exposure have been adjusted with an uncertainty factor of 0.1.

FIR: Food Ingestion Rate (Nagy, 1987). For generic birds with body weight less than or equal to 200 g, the "passerine" equation was used; for generic birds with body weight greater than 200 g, the "all birds" equation was used:

Passerine Equation (body weight < or =200 g): FIR (g dry weight/day) = 0.398 (BW in g) $^{0.850}$ All birds Equation (body weight > 200 g): FIR (g dry weight/day) = 0.648 (BW in g) $^{0.651}$.

For mammals, the "all mammals" equation was used: FIR (g dry weight/day) = 0.235(BW in g) 0.822

BW: Generic Body Weight

EEC: Concentration of pesticide on food item based on Hoerger and Kenaga (1972) and Kenaga (1973) and modified according to Fletcher et al. (1994). At the screening level, relevant food items representing the most conservative EEC for each feeding guild are used.

Table 19 Expanded risk assessment on mammals

Organism/ Toxicity	Food Guild (food item)	Maximum nomogram residues				Mean nome residues	Mean nomogram residues	
		On-field		Off-Field		On-field		
		EDE	RQ	EDE	RQ	EDE	RQ	
		(mg a.i./kg		(mg a.i./kg		(mg a.i./kg		
		bw)		bw)		bw)		
Small Mammal	(0.015 kg)							
Reproduction	Insectivore (small insects)	3.65	1.2	2.15	0.72	2.04	0.68	
3.00	Granivore (grain and	0.91	0.30	0.54	0.18	0.44	0.15	
mg a.i./kg bw/d	seeds)							
	Frugivore (fruit)	1.83	0.61	1.08	0.36	0.87	0.29	
Medium Sized N	Tammal (0.035 kg)							
Acute	Insectivore (small insects)	3.20	0.74	1.89	0.43	1.79	0.42	
4.30	Insectivore (large insects)	0.80	0.19	0.47	0.11	0.38	0.09	
mg a.i./kg bw/d	Granivore (grain and	0.80	0.19	0.47	0.11	0.38	0.09	
	seeds)							
	Frugivore (fruit)	1.60	0.37	0.94	0.22	0.76	0.18	
	Herbivore (short grass)	11.44	2.66	6.75	1.57	4.06	0.94	

 $^{^{2}}$ EDE = Estimated dietary exposure; is calculated using the following formula: (FIR/BW) × EEC, where:

Organism/ Toxicity	Food Guild (food item)	Maximum nomogram residues				Mean nomogram residues	
		On-field		Off-Field		On-field	
			RQ	EDE	RQ	EDE	RQ
		(mg a.i./kg bw)		(mg a.i./kg bw)		(mg a.i./kg bw)	
	Herbivore (long grass)	6.99	1.62	4.12	0.96	2.28	0.53
	Herbivore (forage crops)	10.59	2.46	6.25	1.45	3.50	0.81
Reproduction	Insectivore (small insects)	3.20	1.07	1.89	0.63	1.79	0.60
3.00	Insectivore (large insects)	0.80	0.27	0.47	0.16	0.38	0.13
mg a.i./kg bw/d	Granivore (grain and seeds)	0.80	0.27	0.47	0.16	0.38	0.13
	Frugivore (fruit)	1.60	0.53	0.94	0.31	0.76	0.25
	Herbivore (short grass)	11.44	3.81	6.75	2.25	4.06	1.35
	Herbivore (long grass)	6.99	2.33	4.12	1.38	2.28	0.76
	Herbivore (forage crops)	10.59	3.53	6.25	2.08	3.50	1.17
Large Sized Ma	mmal (1 kg)						
Acute	Insectivore (small insects)	1.71	0.40	1.01	0.23	0.95	0.22
4.30	Insectivore (large insects)	0.43	0.10	0.25	0.06	0.20	0.05
mg a.i./kg bw/d	Granivore (grain and seeds)	0.43	0.10	0.25	0.06	0.20	0.05
	Frugivore (fruit)	0.86	0.20	0.50	0.12	0.41	0.09
	Herbivore (short grass)	6.11	1.42	3.61	0.84	2.17	0.50
	Herbivore (long grass)	3.73	0.87	2.20	0.51	1.22	0.28
	Herbivore (forage crops)	5.66	1.32	3.34	0.78	1.87	0.43
Reproduction	Insectivore (small insects)	1.71	0.57	1.01	0.34	0.95	0.32
3.00	Insectivore (large insects)	0.43	0.14	0.25	0.08	0.20	0.07
mg a.i./kg bw/d	Granivore (grain and seeds)	0.43	0.14	0.25	0.08	0.20	0.07
	Frugivore (fruit)	0.86	0.29	0.50	0.17	0.41	0.14
	Herbivore (short grass)	6.11	2.04	3.61	1.20	2.17	0.72
	Herbivore (long grass)	3.73	1.24	2.20	0.73	1.22	0.41
	Herbivore (forage crops)	5.66	1.89	3.34	1.11	1.87	0.62

Table 20 Toxicity of bifenthrin, formulated products and transformation products to aquatic species

Organism	Exposure	Test substance		Degree of toxicity ¹	Reference
Freshwater sp	ecies				
D. magna	48hr-Acute	Bifenthrin	NOEC < 0.025 μg a.i./L	Very highly	1755275
			$LOEC = 0.025 \mu g \text{ a.i./L}$	toxic	
			$LC_{50} = 0.118 \mu g a.i./L$		
		Talstar 80g/l	$NOEC = 3.2 \mu g \text{ a.i./L}$	Very highly	1762399
		flowable	$LOEC = 5.6 \mu g \text{ a.i./L}$	toxic	
			$LC_{50} = 5.7 \mu g a.i./L$		
	21d-Chronic	Bifenthrin	NOEC =0.0013 μg a.i./L	No	1755269
			$LOEC = 0.0029 \mu g a.i./L$	classification	
C. riparius	Water spiked	Bifenthrin	NOEC = $0.32 \mu g a.i./L$	Very highly	1755267
			$LOEC = 1.0 \mu g \text{ a.i./L}$	toxic	
			$EC_{50} = 3.96 \mu g a.i./L$		
	Sediment spiked		NOEC = $40 \mu g \text{ a.i./kg dw sediment}$		1755279
			$LOEC = 126 \mu g \text{ a.i./kg dw sediment}$	classification	
			$EC_{50} = 345.5 \mu g a.i./kg dw$		
			sediment		

Organism	Exposure	Test substance	Endpoint value	Degree of toxicity ¹	Reference
	Sediment spiked	4'-OH	NOEC = 1.581 mg TP/kg dw	No	1755277
		bifenthrin	sediment	classification	
			LOEC = 5.000 mg TP/kg dw		
			sediment		
			$EC_{50} = 3.593 \text{ mg TP/kg dw}$		
D:1	0.61	D:C 41:	sediment	37 1:11	1775051
Rainbow trout	96hr-Acute	Bifenthrin	NOEC = $0.03 \mu g a.i./L$	Very highly	1755251
(O. mykiss)			LOEC = $0.08 \mu g \text{ a.i./L}$	toxic	
			$LC_{50} = 0.10 \mu g \text{ a.i./L}$ $NOEC = 0.0948 \mu g \text{ a.i./L}$	Very highly	2533229
			LC ₅₀ = 0.256 μg a.i./L	toxic	2333229
			$LC_{50} = 0.230 \mu g a.i./L$	toxic	
			Bifenthrin residues in fish sampled	N/A	2533230
			from the 0.0948 µg a.i./L treatment		
			group: 137±18.1 µg a.i./kg		
			Bifenthrin residues in fish sampled		
			from 0.322 µg a.i./L treatment		
			group: 546 ± 9.76 μg a.i./kg		
		4'-OH	NOEC = $0.805 \mu g TP/L$	Very highly	1755252
		bifenthrin	$LC_{50} = 3.9 \mu g \text{ TP/L}$	toxic	155551
		TFP acid	NOEC = 2.8 mg TP/L	Highly toxic	1755254
			LOEC = 6.2 mg TP/L		
		TD 1 .	$LC_{50} = 24.5 \text{ mg TP/L}$	37 1:11	1750122
		Talstar	NOEC = 0.01 mg a.i./L	Very highly	1759123
		80g/L Flowable	LOEC = 0.018 mg a.i./L	toxic	
Bluegill sunfish	96hr-Acute	Bifenthrin	$LC_{50} = 0.030 \text{ mg a.i./L}$ NOEC = 0.10 µg a.i./L	Very highly	1755246
(L. macrochiru)	90III-Acute	Biteitiiiii	LOEC = $0.10 \mu g a.i./L$	toxic	1733240
(L. macrochiru)			$LC_{50} = 0.26 \mu g \text{ a.i./L}$	toxic	
			NOEC = 0.209 μg a.i./L	Very highly	2533231
			$LC_{50} = 0.269 \mu g a.i./L$	toxic	2333231
			Bifenthrin residues in fish sampled	toxic	2533230
			from the 0.209 µg a.i./L treatment		2333230
			group: $196 \pm 5.74 \mu g \text{ a.i./kg}$		
			group. 170 = 3.7 · µg a.i., ng		
			Bifenthrin residues in fish sampled		
			from 0.346 µg a.i./L treatment		
			group: 625 ± 121 μg a.i./kg		
Fathead minnow	96hr-Acute	Bifenthrin	NOEC =0.083µg a.i./L	Very highly	1755227
			$LOEC = 0.17 \mu g \text{ a.i./L}$	toxic	
			$LC_{50} = 0.21 \mu g \text{ a.i./L}$		
			$LC_{50} = 0.234 \mu g \text{ a.i./L}$	Very highly	2533232
				toxic	
	Full life cycle		NOEC =0.04 µg a.i./L	No	1755227
			$LOEC = 0.09 \mu g \text{ a.i./L}$	classification	
Medaka	96hr-Acute	Bifenthrin	$LC_{50} = 1.77 \text{ g a.i./L}$	Very highly	2533233
(O. latipes)				toxic	
Common carp	96hr-Acute	Bifenthrin	$LC_{50} = 0.635 \ \mu g \ a.i./L$	Very highly	2533234
(C. carpio)				toxic	
Zebra fish	96hr-Acute	Bifenthrin	$LC_{50} = 1.965 \ \mu g \ a.i./L$	Very highly	2533235
(B. rerio)				toxic	
Green alga	72hr-limit test	Talstar 8 SC		No	1755945
(D. subspicatus)			(100 mg EP/L)	classification	

Organism	Exposure	Test substance	Endpoint value	Degree of toxicity ¹	Reference
			$LC_{50} > 6.3 \text{ mg a.i./L}$		
Mesocosm	Phytoplankton	Talstar 80	NOEC =0.935 μg a.i./L ⁺	No	1755962
	Macrophyte	Flowable	NOEC =0.935 μg a.i./L*	classification	
	Zooplankton		NOEC =0.001 μg a.i./L*		
			$LOEC = 0.005 \mu g \text{ a.i./L*}$		
	Macroinvertebrate		NOEC =0.005 μg a.i./L*		
			$LOEC = 0.015 \mu g a.i./L*$		
	Taxonomic richness		NOEC =0.015 μg a.i./L*		
			$LOEC = 0.037 \mu g a.i./L*$		
Marine species					
Mysids	96hr-Acute	Bifenthrin	NOEC <0.00248 µg a.i./L	Very highly	1755262
(A. bahia)			$LOEC = 0.00248 \ \mu g \ a.i./L$	toxic	
			$LC_{50} = 0.00397 \mu g \text{ a.i./L}$		
	28d-Chronic	Bifenthrin	$NOEC = 0.0012 \mu g a.i./L$	No	1755258
			$LOEC = 0.0013 \mu g a.i./L$	classification	
			$LC_{50} = 0.00226 \mu g a.i./L$		
Eastern Oyster	48hr-Acute embryo	Bifenthrin	NOEC = 0.126g mg a.i./L	Highly toxic	1755261
(C. virginica)			LOEC = 0.448 mg a.i./L		
			$EC_{50} = 0.280 \text{ mg a.i./L}$		
	96hr-Acute shell	Bifenthrin	$NOEC = 1.09 \mu g a.i./L$	Very highly	1755260
	deposition		$LOEC = 2.15 \mu g a.i./L$	toxic	
			$EC_{50} > 2.15 \mu g \text{ a.i./L}$		
Sheepshead	96hr-Acute	Bifenthrin	NOEC = $10.9 \mu g a.i./L$	Very highly	1755242
minnow			$LC_{50} = 17.53 \mu g a.i./L$	toxic	
(C. variegatus)					

¹ USEPA classification, where applicable

Table 21 Screening level risk assessment of bifenthrin to aquatic organisms

Organism	Exposure	Endpoint value	EEC***	RQ	LOC
Freshwater species	S				
D. magna	Acute	$LC_{50}/2 = 0.059 \mu g a.i./L$	27 μg a.i./L	458	Exceeded
	Chronic	NOEC = $0.0013 \mu g a.i./L$		20,770	
C. riparius	Water spiked	$EC_{50}/2 = 1.98 \mu g a.i./L$		14	
Freshwater invertebrates SSD	Acute	$HC_5 = 0.009 \ \mu g \ a.i./L$		3000	
Rainbow trout	Acute	$LC_{50}/10 = 0.01 - 0.0256 \ \mu g$ a.i./L		1055 - 2700	
Bluegill sunfish	Acute	$LC_{50}/10 = 0.026 - 0.027 \ \mu g$ a.i./L		1000 - 1038	
Fathead minnow	Acute	$LC_{50}/10 = 0.021 - 0.023 \ \mu g$ a.i./L		1174 - 1286	
	Chronic	NOEC = $0.04 \mu g \text{ a.i./L}$	1	675	7
Medaka (O. latipes)	Acute	$LC_{50}/10 = 0.177 \mu g \text{ a.i./L}$		153	
Common carp	Acute	$LC_{50}/10 = 0.064 \mu g a.i./L$		422	

^{*} Based on nominal treatment concentration of one application.

 $^{^{+}}$ Increased abundance was observed at 0.037 μg a.i./L and was not considered as adverse effect. Therefore, the highest treatment level was the NOEC.

Organism	Exposure	Endpoint value	EEC***	RQ	LOC
(C. carpio)					
Zebra fish	Acute	$LC_{50}/10 = 0.197 \mu g a.i./L$		137	
(B. rerio)					
Freshwater fish	Acute	$HC_5 = 0.078 \mu g a.i./L$		346	
SSD					
Green algae	Acute	EC ₅₀ /2>3150 µg a.i./L	27 μg a.i./L	< 0.01	Not exceeded
Amphibian	Acute*	Fish $HC_{05} = 0.078 \mu g a.i./L$	144 μg a.i./L	1846	Exceeded
	Chronic**	NOEC = $0.04 \mu g a.i./L$		3600	
Marine species					
Mysid A. bahia	Acute	$LC_{50}/2 = 0.00199 \mu g a.i./L$	27 μg a.i./L	13602	Exceeded
	Chronic	NOEC = $0.0012 \mu g a.i./L$		22500	
Eastern oyster	Acute embryo	$LC_{50}/2 = 140 \mu g a.i./L$	27 μg a.i./L	0.2	Not exceeded
	Acute shell deposition	$EC_{50}/2 > 1.08 \mu g a.i./L$	27 μg a.i./L	<25	Exceeded
Sheepshead minnow	Acute	$LC_{50}/10 = 1.753 \mu g a.i./L$	27 μg a.i./L	15	Exceeded

^{*}Rainbow trout data was used as surrogate.

Table 22 Off-field risk assessment of bifenthrin to aquatic organisms from spray drift

Organism	Exposure	Endpoint value	EEC***	RQ	LOC
Freshwater species					
Freshwater	Acute	$HC_5 = 0.009 \mu g a.i./L$	16 μg a.i./L	1778	Exceeded
invertebrates SSD					
Freshwater	Chronic	$NOEC/3 = 0.005 \mu g \text{ a.i./L}$		3000	
Invertebrates					
mesocosm –					
Taxonomic richness					
Fathead minnow	Chronic	NOEC = $0.04 \mu g \text{ a.i./L}$		400	
Freshwater fish SSD	Acute	Fish $HC_5 = 0.078 \mu g a.i./L$]	205	
Amphibian	Acute	Fish $HC_5 = 0.078 \mu g a.i./L$	95	1090	
	Chronic	NOEC = $0.04 \mu g \text{ a.i./L}$	85 µg a.i./L	2125	
Marine species					
Mysid A. bahia	Acute	$LC_{50}/2 = 0.001985 \mu g$	16 μg a.i./L	8060	Exceeded
		a.i./L			
	Chronic	NOEC = $0.0012 \mu g a.i./L$		13333	
Eastern oyster	Acute shell deposition	$EC_{50}/2 = 1.08 \mu g a.i./L$		15	
Sheepshead minnow	Acute	$LC_{50}/10 = 1.753 \mu g a.i./L$		9	

^{***}The EEC values were based on a 15 cm water depth for amphibians and a 80 cm water depth for all other aquatic organisms. The spray drift exposure was determined by assuming 59% drift from a later season airblast application on raspberries. The estimated EEC values were higher than the theoretical solubility. Bolded values indicates an exceedance of the level of concern (RQ = 1).

^{**}Fathead minnow full life cycle data was used as surrogate.

[#]The NOEC was the highest treatment concentration.
***EEC based on a 15 cm water body depth for amphibians and an 80 cm water depth for all other aquatic organisms. Bolded values indicates an exceedance of the level of concern (RQ = 1).

Table 23 Off-field risk assessment of bifenthrin to aquatic organisms from predicted runoff excluding spray drift

Organism	Exposure	Endpoint value	EEC (µg a.i./L)	RQ	LOC
Freshwater species					
Freshwater invertebrates	Acute	$HC_5 = 0.009 \mu g a.i./L$	5.21	578	Exceeded
SSD					
Fathead minnow	Chronic	NOEC = $0.04 \mu g a.i./L$	0.19^{5}	5	Exceeded
Freshwater fish SSD	Acute	Fish $HC_5 = 0.078 \mu g a.i./L$	0.414	5	
Amphibian	Acute	Fish $HC_5 = 0.078 \mu g a.i./L$	1.46	18	
	Chronic	NOEC = $0.04 \mu g a.i./L$	0.217	5]
Mesocosm – Taxonomic	Chronic	$NOEC/3 = 0.005 \mu g a.i./L$	0.19^{5}	38	Exceeded
richness					
Marine species					
Mysid A. bahia	Acute	$LC_{50}/2 = 0.001985 \mu g a.i./L$	0.41^4	207	Exceeded
	Chronic	NOEC = $0.0012 \mu g a.i./L$	0.25^{2}	208	
Eastern oyster	Acute shell	$EC_{50}/2 = 1.08 \mu g a.i./L$	0.41^4	0.38	Not exceeded
	deposition				
Sheepshead minnow	Acute	$LC_{50}/10 = 1.753 \mu g a.i./L$		0.23	

¹ Acute EEC for *D. magna* was based on the highest detection of bifenthrin out of all surface water samples collected.

Table 24 Toxic Substances Management Policy Considerations-Comparison to TSMP **Track 1 Criteria**

TSMP Track 1 Criteria	TSMP Track 1 Criterion value	Bifenthrin Assessment against criteria
CEPA toxic or	A substance is CEPA toxic (or	Yes: Bifenthrin meets the criteria under
CEPA toxic	CEPA toxic equivalent) if it is	paragraph 64 (a) of CEPA and should be
equivalent	entering or may enter the	considered CEPA-toxic equivalent.
	environment in a quantity or concentration or under conditions	
	that	
	(a) have or may have an	
	immediate or long-term harmful	
	effect on the environment or its	
	biological diversity;	
	(b) constitute or may constitute a danger to the environment on	
	which life depends; or	
	(c) constitute or may constitute a	
	danger in Canada to human life or	
	health.	
Predominantly	The policy considers a substance	Yes: All releases to the environment are
anthropogenic	"predominantly anthropogenic" if,	anthropogenic.
	based on expert judgment, its	
	concentration in the environment medium is largely due to human	
	medium is largery due to numan	

² Chronic EEC for *D. magna* was based on the nignest detection of of offentifin out of an surface water samples confected.

² Chronic EEC for *D. magna* and *A. bahia* was based on the 90th percentile of the 21 day average for 80 cm deep water body.

⁴ Acute EEC for fish and marine species was based on the 90th percentile of the 96 hour average for 80 cm deep water body.

⁵ Chronic EEC for fathead minnow was based on the 90th percentile of the yearly average for 80 cm deep water body.

⁶ Acute EEC for amphibian was based on the 90th percentile of the 96 hour average for 15 cm deep water body.

⁷ Chronic EEC for amphibian was based on the 90th percentile of the yearly average for 15 cm deep water body.

TSMP Track 1 Criteria	TSMP Track 1 Criterion value		Bifenthrin Assessment against criteria		
	activity, rather th	an to natural			
	sources or release	es.			
Persistence ¹ :	Soil	Half-life ≥ 182 days	Yes: Laboratory aerobic soil half-life: 78.7 to 203 days		
			Field dissipation study half-life: 80-215 days		
	Whole system (Water +	Half-life ≥ 182 days	Yes: Aerobic water sediment whole system half-life: 92.9 to 276 days		
	sediment)	(water) Half-life			
		≥ 365 days (sediment)			
	Air	Half-life ≥ 2	Not determined		
		days or evidence of			
		long range transport			
	Other	In an aquatic field	d study conducted in Alabama, bifenthrin residues		
		in the pond sediment persisted for two years after the last application. This was possibly due to its persistence in soil and continuous input into the pond from runoff for the two years following the last application. Variability in sediment and water			
		concentrations wa	as high at each sample date, but this is common in		
			should not be considered as detrimental to rall lack of decline of residues over time in the		
			n addition, the monitoring data collected for and relevance to the persistence observed in the		
		laboratory study i	results, as bifenthrin was frequently detected in		
		NAWQA, PMRA	s collected in agricultural areas in the US (USGS \(\text{2398587} \)).		
	systems. The env is acknowledged	ironmental fate and that the range in ha owever under most	on is met in: soil, water, sediment and aquatic d persistence of bifenthrin is well characterized. It alf-lives includes values below and above the environmental conditions, bifenthrin is		
	Long-range atmo	spheric transport c	annot be determined at this time. The AOPWIN		
			e atmospheric half-life of bifenthrin given the to airborne particles.		
Bioaccumulation ²	Log K _{ow} ≥ 5		20°C (PMRA 1755525) EPA 2010, cited from Laskowski 2002) EFSA 2011)		
	$BCF^3 \ge 5000$	Bluegill BCF _K of Carp: B0	tted laboratory studies: sunfish: BCF _{K,G,L} of 3400 - 3511; 5 5250 to 12,850 CF _K of 1265 - 1861		
		• Fathead	minnow adult: BCF _{SS} : 21,000 to 30,000		

TSMP Track 1 Criteria	TSMP Track 1 C	Criterion value	Bifenthrin Assessment against criteria		
	$BMF^{4, 5} \ge 1$	Diotomy bioma	ification study (bluesill sunfish)		
	BMF ^{3,2} ≥ 1	BMF _{K,G,L} : 0.28	ification study (bluegill sunfish):		
		Half-life in fish: 20 days Assimilation efficiency, α : 0.039			
			tes that dietary uptake of bifenthrin did not icantly to biomagnification as the reported BMF		
	BAF ≥ 5000	Field studies:			
		Under field conditions when biota are exposed through multiple pathways, bioaccumulation occurs as was observed in various fissispecies under field conditions in the Alabama pond study: Catfish: BAF of 134 – 5385 Channel catfish: BAF of 77 – 12682 Gizzard shad: BAF of 499 – 12458 Threadfin shad: BAF of 182 – 1855 Redear sunfish: BAF of 51 – 3844			
			BAF of 535 – 11564		
		White crappie: B	BAF of 11 – 7430 AF of 11 – 3430 S: BAF of 116 - 8715		
	Other	detected bifenthr samples (Alonso evidence of expo mammals and m	ng study of dolphins from the Brazilian coast in residues in liver, breast milk, and placental et al., 2012). The results of this study provide sure and accumulation of bifenthrin in marine aternal transfer by both gestational and lactation agricultural areas.		
	Conclusion: The h		otential of bifenthrin is well characterized based		
	on the information		Schilar of offending is well characterized based		
	fish species. One sconsidering the di	study showed that et under laborator er, sufficient info	to show that bifenthrin BCFs are > 5000 in some bifenthrin does not biomagnify in fish when only y conditions (BMFs are <1.0). Under field rmation was provided to show that bifenthrin er applications.		
I If the masticide of	monograph along BCF/BAFs. The f	s have been integrated into the bioaccumulation summary in the g with all other lines of evidence. BMF <1 should not supersede field BAF values are a better representation of bioaccumulation ntally relevant conditions and consider multiple pathways of			

¹ If the pesticide and/or the transformation product(s) meet one persistence criterion identified for one media (soil, water, sediment or air) than the criterion for persistence is considered to be met.

² Field data (e.g., BAFs) are preferred over laboratory data (e.g., BCFs) which, in turn, are preferred over

TSMP Track 1	TSMP Track 1 Criterion value	Bifenthrin Assessment against criteria
Criteria		

chemical properties (e.g., log K_{OW}).

 $^{^3}$ BCF_K: kinetic BCF; BCF_{K,G,L}: kinetic BCF corrected for growth and lipid content; BCF_{SS}: steady state BCF

⁴BMF_{K,G,L}: kinetic BMF corrected for growth and lipid content

⁵ Biomagnification results from the processes of bioaccumulation and bioconcentration by which tissue concentrations of bioaccumulated substances increase as they are passed up through two or more trophic levels. The term implies an efficient transfer from food to consumer, so that residue concentrations increase systematically from one trophic level to the next. The extent of biomagnification is also influenced by several factors – the substance being accumulated, the number of stages in the food web, the kind of organisms in the food web, the ability of these organisms to metabolize the substance, the concentration of the substance at each level of the web, etc. (Government of Canada, 1995)

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1.0 Chemistry		
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