

Proposed Registration Decision

PRD2016-03

Mandestrobin

(publié aussi en français)

29 January 2016

This document is published by the Health Canada Pest Management Regulatory Agency. For further information, please contact:

Publications Pest Management Regulatory Agency Health Canada 2720 Riverside Drive A.L. 6607 D Ottawa, Ontario K1A 0K9

Internet:

pmra.publications@hc-sc.gc.ca healthcanada.gc.ca/pmra Facsimile: 613-736-3758 Information Service: 1-800-267-6315 or 613-736-3799 pmra.infoserv@hc-sc.gc.ca



ISSN: 1925-0878 (print) 1925-0886 (online)

Catalogue number: H113-9/2016-03E (print version) H113-9/2016-03E-PDF (PDF version)

© Her Majesty the Queen in Right of Canada, represented by the Minister of Health Canada, 2016

All rights reserved. No part of this information (publication or product) may be reproduced or transmitted in any form or by any means, electronic, mechanical, photocopying, recording or otherwise, or stored in a retrieval system, without prior written permission of the Minister of Public Works and Government Services Canada, Ottawa, Ontario K1A 0S5.

Table of Contents

Overview		1
Proposed Reg	gistration Decision for Mandestrobin	1
What Does H	lealth Canada Consider When Making a Registration Decision?	1
	destrobin?	
Health Consi	derations	2
Environment	al Considerations	4
Value Consid	lerations	5
Measures to I	Minimize Risk	5
Next Steps		6
Other Inform	ation	6
Science Evalua	tion	7
1.0	The Active Ingredient, Its Properties and Uses	7
1.1	Identity of the Active Ingredient	
1.2	Physical and Chemical Properties of the Active Ingredient and End-Use	
	Products	8
1.3	Directions for Use	9
1.4	Mode of Action	. 10
2.0	Methods of Analysis	. 10
2.1	Methods for Analysis of the Active Ingredient	. 10
2.2	Method for Formulation Analysis	
2.3	Methods for Residue Analysis	. 10
3.0	Impact on Human and Animal Health	. 10
3.1	Toxicology Summary	
3.1.1	Pest Control Products Act Hazard Characterization	. 14
3.2	Acute Reference Dose (ARfD)	. 15
3.3	Acceptable Daily Intake	. 15
3.4	Occupational and Residential Risk Assessment	. 16
3.4.1	Toxicological Endpoints	. 16
3.4.2	Occupational Exposure and Risk	. 17
3.4.3	Residential Exposure and Risk Assessment	. 23
3.5	Food Residues Exposure Assessment	. 25
3.5.1	Residues in Plant and Animal Foodstuffs	
3.5.2	Concentrations in Drinking Water	. 26
3.5.3	Dietary Risk Assessment	. 27
3.5.4	Aggregate Exposure and Risk	. 28
3.5.5	Maximum Residue Limits	. 28
4.0	Impact on the Environment	. 29
4.1	Fate and Behaviour in the Environment	. 29
4.2	Environmental Risk Characterization	. 30
4.2.1	Risks to Terrestrial Organisms	. 30
4.2.2	Risks to Aquatic Organisms	
4.2.3	Further characterization of risk to aquatic organisms	. 34

5.0	Value	. 35
5.1	Consideration of Benefits	. 35
5.2	Effectiveness Against Pests	. 35
5.3	Non-Safety Adverse Effects	. 36
5.4	Supported Uses	. 36
6.0	Pest Control Product Policy Considerations	. 36
6.1	Toxic Substances Management Policy Considerations	
6.2	Formulants and Contaminants of Health or Environmental Concern	
7.0	Summary	. 37
7.1	Human Health and Safety	
7.2	Environmental Risk	. 38
7.3	Value	. 39
8.0	Proposed Regulatory Decision	. 39
List of Abbrev	viations	
Appendix I	Tables and Figures	. 47
Table 1a	Residue Analysis for Environmental Media	. 47
Table 1b	Residue Analysis for Crops	
Table 2	Chemical Names of Isomers and Metabolites of Mandestrobin	. 48
Table 3	Toxicity Profile of S-2200 3.2 FS Fungicide	. 48
Table 4	Toxicity Profile of S-2200 4 SC Fungicide, S-2200 4 SC AG Fungicide and	
	2200 4 SC VPP Fungicide	
Table 5	Toxicity Profile of Technical Mandestrobin	
Table 6	Toxicity Profile of Metabolites of Mandestrobin	
Table 7	Toxicology Endpoints for Use in Health Risk Assessment for Mandestrobin.	
Table 8	Integrated Food Residue Chemistry Summary	
Table 9	Food Residue Chemistry Overview of Metabolism Studies and Risk	
	Assessment	. 69
Table 10	Fate and Behaviour in the Environment	. 70
Table 11	Major Transformation Products Formed in the Environment	. 77
Table 12	Toxicity of Mandestrobin and Transformation Products to Non-Target	
	Terrestrial Species	. 79
Table 13	Screening Level and Refined Risk Assessment of Mandestrobin for Non-Tar	get
	Species, Other than Birds and Mammals	. 80
Table 14	Screening Level Risk Assessment of Foliar Application of Mandestrobin for	
	Birds and Mammals	. 81
Table 15	Screening Level Assessment of Seed Treatment with Mandestrobin for Birds	S
	and Mammals	
Table 16	Toxicity of Mandestrobin, S-2200-R-Isomer, S-2200-S-Isomer and	
	Transformation Products to Non-Target Aquatic Species	. 82
Table 17	Screening Level Risk Assessment of Mandestrobin to Aquatic Organisms	
Table 18	Screening Level Risk Assessment of Mandestrobin Isomers and	
	Transformation Products for Terrestrial and Aquatic Organisms	. 85
Table 19	Refined Risk Assessment of Potential Risk from Drift of Mandestrobin, S-22	
	R-isomer and the Transformation Product S-2200-ORC to Aquatic Organism	
	1 0	

Table 20	Risk Quotients for Aquatic Organisms Determined for Runoff of	
	Mandestrobin, S-2200- <i>R</i> -Isomer and S-2200-ORC in Water Bodies	87
Table 21	Toxic Substances Management Policy Considerations-Comparison to TSMP	
	Track 1 Criteria	87
Table 22	Fungicide Resistance Action Committee modes of action groups of currently	
	registered alternative products (as of June 2015)	88
Table 23	List of Supported Uses	89
Appendix II	Supplemental Maximum Residue Limit Information—International Situation	l
	and Trade Implications	91
Table 1	Comparison of Canadian MRLs, American Tolerances and Codex MRLs	91
References		93

Overview

Proposed Registration Decision for Mandestrobin

Health Canada's Pest Management Regulatory Agency (PMRA), under the authority of the *Pest Control Products Act* and Regulations, is proposing full registration for the sale and use of Mandestrobin Technical (previously known as S-2200 Fungicide Technical) and the associated end-use products: S-2200 4 SC Fungicide, Intuity Fungicide (previously known as S-2200 4 SC Ag Fungicide), Pinpoint Fungicide (previously known as S-2200 4 SC VPP Fungicide), and S-2200 3.2 FS Fungicide, containing the technical grade active ingredient mandestrobin, for the management of various fungal diseases in canola and other oilseed crops, corn, grape, legume vegetables, strawberry and other low growing berries, as well as turfgrass.

An evaluation of available scientific information found that, under the approved conditions of use, the product has value and does not present an unacceptable risk to human health or the environment.

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 S-2200 Fungicide Technical, S-2200 4 SC Fungicide, S-2200 4 SC Ag Fungicide, S-2200 4 SC VPP Fungicide, and S-2200 3.2 FS Fungicide.

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 acceptable¹ 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 value² when used according to the label directions. Conditions of registration may include special precautionary measures on the product label to further reduce risk.

¹ "Acceptable risks" as defined by subsection 2(2) of the *Pest Control Products Act*.

² "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."

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, the assessment process and risk-reduction programs, 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 mandestrobin, the PMRA will consider any comments received from the public in response to this consultation document.³ The PMRA will then publish a Registration Decision⁴ on mandestrobin, which will include the decision, the reasons for it, a summary of comments received on the proposed 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 Mandestrobin?

Mandestrobin is the active ingredient in the following fungicide products that are being proposed for registration in Canada: S-2200 4 SC Fungicide, S-2200 4 SC Ag Fungicide, S-2200 4 SC VPP Fungicide, and S-2200 3.2 FS Fungicide. These products are formulated for either foliar or seed applications and are intended for the management of various fungal diseases in canola and other oilseed crops, corn, grape, legume vegetables, strawberry and other low growing berries, as well as turfgrass. Mandestrobin has preventative and systemic properties and acts by interfering with the cellular mechanisms in susceptible fungal pathogens.

Health Considerations

Can Approved Uses of Mandestrobin Affect Human Health?

Products containing mandestrobin are unlikely to affect your health when used according to label directions.

Potential exposure to mandestrobin may occur through the diet (food and water) or when handling and applying the products. When assessing health risks, two key factors are considered: the levels where no health effects occur 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). Only uses for which the exposure is well below levels that cause no effects in animal testing are considered acceptable for registration.

³ "Consultation statement" as required by subsection 28(2) of the *Pest Control Products Act*.

⁴ "Decision statement" as required by subsection 28(5) of the *Pest Control Products Act*.

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 products are used according to label directions.

In laboratory animals, the technical grade active ingredient mandestrobin was of low acute toxicity by the oral, dermal and inhalation routes. Mandestrobin was minimally irritating to the eye and non-irritating to the skin. Mandestrobin did not cause allergic skin reactions.

The acute toxicity of the end-use products was low via the oral, dermal and inhalation routes of exposure. The products were non-irritating to the skin and minimally irritating to the eyes. They did not cause allergic skin reactions. Consequently, no hazard signal words are required on their labels.

Applicant-supplied short- and long-term (lifetime) animal toxicity tests were assessed for the potential of mandestrobin to cause neurotoxicity, immunotoxicity, chronic toxicity, cancer, reproductive and developmental toxicity, and various other effects. The most sensitive endpoints for risk assessment were adverse effects noted on growth and in the liver, bile duct, and kidneys. There was a low level of concern for sensitivity of the young animal. The risk assessment protects against the finding noted above as well as any other potential effects by ensuring that the level of exposure to humans is well below the lowest dose at which these effects occurred 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 mandestrobin relative to body weight, are expected to be exposed to less than 10% of the acceptable daily intake. Based on these estimates, the chronic dietary risk from mandestrobin is not of health concern for all population subgroups.

Animal studies revealed no acute health effects. Consequently, a single dose of mandestrobin is not likely to cause acute health effects in the general population (including infants and children). Mandestrobin is not carcinogenic; therefore, a cancer dietary risk assessment is not required.

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 Canada (and the United States) using mandestrobin on rapeseed, corn, grapes, strawberries, and soybeans are acceptable. The MRLs for this active ingredient can be found in the Science Evaluation of this consultation document.

Occupational Risks From Handling S-2200 4 SC Fungicide, S-2200 3.2 FS Fungicide, S-2200 4 SC Ag Fungicide, and S-2200 4 SC VPP Fungicide

Occupational risks are not of concern when S-2200 4 SC Fungicide, S-2200 3.2 FS Fungicide, S-2200 4 SC Ag Fungicide and S-2200 4 SC VPP Fungicide are used according to the label directions, which include protective measures.

Farmers, custom applicators, seed treaters and planters who mix, load, apply or treat seeds with S-2200 4 SC Fungicide, S-2200 3.2 FS Fungicide, S-2200 4 SC Ag Fungicide, and S-2200 4 SC VPP Fungicide, as well as field workers re-entering freshly treated fields, can come in direct contact with residues on the skin. Therefore, the label specifies that anyone mixing/loading and applying S-2200 4 SC Fungicide, S-2200 4 SC Ag Fungicide, and S-2200 4 SC VPP Fungicide must wear a long-sleeved shirt, long pants, chemical-resistant gloves, shoes and socks. The field crew and the mixer/loaders for aerial applications of S-2200 4 SC Ag Fungicide must wear coveralls over a long-sleeved shirt, long pants, chemical-resistant gloves, shoes and socks, and goggles or face shield during mixing/loading, cleanup and repair. The label also requires that workers do not enter treated fields for 12 hours after application. Workers treating seeds with S-2200 3.2 FS Fungicide must use closed transfer systems in commercial facilities (excluding mobile treaters), and must wear long pants, a long-sleeved shirt and chemical-resistant gloves, shoes and socks during mixing, loading, treating, bagging, sewing or stacking of bagged treated seed, handling treated seed, and planting treated seed. In addition, seed treatment workers performing cleaning, maintenance, and repair of seed treatment equipment must wear chemicalresistant coveralls. As well, standard label statements to protect against drift during application are on the labels. Taking into consideration these label statements, the use pattern, and the duration of exposure for handlers and workers, risks to these individuals are not a concern.

For bystanders, exposure is expected to be much less than that for workers and is considered negligible. Therefore, health risks to bystanders are not of concern.

Environmental Considerations

What Happens When Mandestrobin Is Introduced Into the Environment?

When used according to label directions mandestrobin is not expected to pose an unacceptable risk to the environment.

Mandestrobin enters the environment when applied as a foliar spray, seed treatment or chemigation. Mandestrobin can break down in the presence of microbes in terrestrial systems. Laboratory studies indicate mandestrobin has the potential to be persistent in certain soils, whereas field studies indicate that mandestrobin is less likely to persist in the environment. The properties of mandestrobin and its transformation products, 5-COOH-S-2200 and 2-COOH-S-2200, indicate some potential for downward movement through the soil. However, field studies and modelling indicate that levels of mandestrobin and its transformation products that may reach groundwater are low. In aquatic systems, mandestrobin will move out of the water column and into sediments where it has the potential to persist.

Mandestrobin does not break down by reacting with water, but it can break down rapidly in the presence of sunlight, especially in clear shallow waters. It is not expected to accumulate in the tissues of aquatic organisms. Mandestrobin is not expected to enter the atmosphere nor be transported long distances from where it was applied.

When used according to label directions, mandestrobin is expected to pose a negligible risk to earthworms, bees, beneficial arthropods, birds and small mammals. If exposed to high enough concentrations, mandestrobin may pose a risk to non-target aquatic organisms and terrestrial plants. Risks to non-target aquatic organisms and terrestrial plants can be mitigated with label statements and spray buffer zones to protect sensitive aquatic and terrestrial habitats. Label statements are required on the product labels to inform the users of the potential risks.

Value Considerations

What Is the Value of S-2200 4 SC Fungicide, S-2200 4 SC Ag Fungicide, S-2200 4 SC VPP Fungicide, and S-2200 3.2 FS Fungicide?

These products have demonstrated good efficacy and will provide growers with additional product options for the management of a broad range of common diseases in economically important crops in Canada.

S-2200 4 SC Fungicide, S-2200 4 SC Ag Fungicide, and S-2200 4 SC VPP Fungicide are formulated for foliar spray applications to control or suppress a range of diseases on oil seed crops, grape, low growing berries including strawberry, and turfgrass. S-2200 3.2 FS Fungicide is applied as a seed treatment to control various common fungi that cause seed rots in corn, legume vegetables, and oil seed crops. Mandestrobin is most effective when applied preventatively or at the early stages of disease development. Appropriate use of these products will help growers maximize the quality and yield of their crops.

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 S-2200 4 SC Fungicide, S-2200 3.2 FS Fungicide, S-2200 4 SC Ag Fungicide, and S-2200 4 SC VPP Fungicide 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 S-2200 4 SC Fungicide, S-2200 3.2 FS Fungicide, S-2200 4 SC Ag Fungicide, and S-2200 4 SC VPP Fungicide on the skin or through inhalation of spray mists, anyone mixing, loading and applying S-2200 4 SC Fungicide, S-2200 4 SC Ag Fungicide, and S-2200 4 SC VPP Fungicide must wear a long-

sleeved shirt, long pants, chemical-resistant gloves, shoes and socks. The field crew and the mixer/loaders for aerial applications of S-2200 4 SC Ag Fungicide must wear coveralls over a long-sleeved shirt, long pants, chemical-resistant gloves, shoes and socks, and goggles or face shield during mixing/loading, cleanup and repair. The label also requires that workers do not enter treated fields for 12 hours after application.

Workers treating seeds with S-2200 3.2 FS Fungicide must use closed transfer systems in commercial facilities (excluding mobile treaters), and must wear long pants, a long-sleeved shirt and chemical-resistant gloves, shoes and socks during mixing, loading, treating, bagging, sewing or stacking of bagged treated seed, handling treated seed, and planting treated seed. In addition, seed treatment workers performing cleaning, maintenance, and repair of seed treatment equipment must wear chemical-resistant coveralls. Standard label statements to protect against drift during application are also on the labels.

Environment

Mandestrobin can pose a risk to non-target aquatic organisms and terrestrial plants. To mitigate potential exposures to mandestrobin via spray drift, spray buffer zones of 0 to 15 metres are required to protect sensitive terrestrial and aquatic habitats, depending on the method of application. These spray buffer zones are to be specified on the product labels.

Next Steps

Before making a final registration decision on mandestrobin, 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 note that, to comply with Canada's international trade obligations, consultation on the proposed MRLs will also be conducted internationally via a notification to the World Trade Organization. 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 mandestrobin (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

Mandestrobin

1.0 The Active Ingredient, Its Properties and Uses

1.1 Identity of the Active Ingredient

Active substance	Mandestrobin
Function	Fungicide
Chemical name	
	Union $rac-(2R)-2-\{2-[(2,5-dimethylphenoxy)methyl]phenyl\}-2-$ pplied methoxy- <i>N</i> -methylacetamide PAC)
2. Chemical Abs Service (CAS)	tracts 2-[(2,5-dimethylphenoxy)methyl]-α-methoxy- <i>N</i> -methylbenzeneacetamide
CAS number	173662-97-0
Molecular formul	$\mathbf{a} \qquad \mathbf{C}_{19}\mathbf{H}_{23}\mathbf{NO}_{3}$
Molecular weight	313.4
Structural formul	a H ₃ C CH ₃ OCH ₃ CONHCH ₃
Purity of the activ	re 88.8%

ingredient

1.2 Physical and Chemical Properties of the Active Ingredient and End-Use Products

Property	Result							
Colour and physical state	White solid							
Odour	Odourless							
Melting point	102°C							
Boiling point	296°C							
Relative density	1.2015							
Vapour pressure at 20°C	3.36×10^{-8} Pa (extrap	olated)						
Ultraviolet (UV)-visible spectrum	Solution	λmax (nm)	Molar absorptivity, (ϵ) (mol/L) ⁻¹ cm ⁻¹					
	Methanol	273	2140					
	Acidic	273	1920					
	Basic	273	1880					
	Neutral	273	1740					
Solubility in water at 20°C	15.8 mg/L							
Solubility in organic solvents at								
20°C	Dichloromethane	395						
	Acetone	332						
	Ethyl acetate	274						
	Methanol	182						
	Toluene	147						
	n-Octanol	63.6						
	Hexane	2.00						
<i>n</i> -Octanol-water partition coefficient (K_{ow})	$Log K_{ow} = 3.51 at 25$	°C						
Dissociation constant (pK_a)	No dissociative activ	ity in the pH ra	ange 2-10					
Stability (temperature, metal)	Stable for 14 days when exposed to normal and elevated (54°C) temperatures and metals and metal ions (iron, nickel, iron (II) acetate and nickel (II) acetate).							

Technical Product—S-2200 Fungicide Technical

End-Use Product—S-2200 3.2 FS Fungicide

Property	Result
Colour	White, opaque
Odour	Faint, paint-like odour
Physical state	Liquid
Formulation type	Suspension
Guarantee	383 g/L

Property	Result
Container material and description	High-density polyethylene (HDPE)
Density at 20°C	1.091 g/mL
pH of 1% dispersion in water	5.7 at 25°C
Oxidizing or reducing action	No oxidizing or reducing action
Storage stability	In progress
Corrosion characteristics	In progress
Explodability	Not explosive

End-Use Product—S-2200 4 SC Fungicide, S-2200 4 SC Ag Fungicide and S-2200 4 SC VPP Fungicide

Property	Result
Colour	White, opaque
Odour	Sweet odour
Physical state	Liquid
Formulation type	Suspension
Guarantee	43.4%
Container material and description	High-density polyethylene (HDPE)
Density at 20°C	1.095 g/mL
pH of 1% dispersion in water	6.8 at 25°C
Oxidizing or reducing action	No oxidizing or reducing action
Storage stability	In progress
Corrosion characteristics	In progress
Explodability	Not explosive

1.3 Directions for Use

S-2200 4 SC Fungicide, S-2200 4 SC Ag Fungicide, and S-2200 4 SC VPP Fungicide are to be applied by foliar application to labelled crops on a preventative basis or at the earliest signs of infection. The application rates range from 439 to 986 mL/ha. When repeated treatments are required and permitted, reapplication intervals range from seven to 28 days.

S-2200 3.2 FS Fungicide is applied once, directly to seed before planting. Application rates per 100 kg of seed range from 15.6 to 26 mL depending on the treated crop. Because S-2200 3.2 FS Fungicide does not contain a colourant, the product must be applied in combination with an appropriate colourant.

1.4 Mode of Action

Mandestrobin belongs to the quinone outside inhibitors class of fungicides. These fungicides provide control of target pathogens by interfering with their ability to produce energy, which in turn inhibits development of spores and mycelia. Mandestrobin is a broad spectrum fungicide with preventive and systemic properties. It is most effective when applied prior to infection.

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 impurities in the technical product have been validated and assessed to be acceptable for the determinations.

2.2 Method for Formulation Analysis

The method provided for the analysis of the active ingredient in the formulations has been validated and assessed to be acceptable for use as an enforcement analytical method.

2.3 Methods for Residue Analysis

For environmental media, 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 environmental media. Methods for residue analysis of environmental media are summarized in Appendix I, Table 1a.

For crops, high performance liquid chromatography methods with tandem mass spectrometric detection (HPLC-MS/MS; Method RM-48C-2A) in plant matrices was developed and proposed for data gathering and enforcement purposes for the determination of mandestrobin residues. The method fulfilled the requirements with regards to specificity, accuracy and precision at the respective method limit of quantitation (0.02 ppm). Acceptable recoveries (70–120%) were obtained in plant matrices. The proposed enforcement method was successfully validated in plant matrices by an independent laboratory. In addition the extraction solvents used in the method were similar to those used in the metabolism studies. Method RM-48M-1 was developed and proposed for enforcement purposes for poultry muscle only. Methods for residue analysis of crops are summarized in Appendix I, Table 1b.

3.0 Impact on Human and Animal Health

3.1 Toxicology Summary

A detailed review of the toxicological database for mandestrobin was conducted. The database is complete, consisting of the full array of toxicity studies required for hazard assessment purposes. The studies were carried out in accordance with accepted international testing protocols and Good Laboratory Practices. The scientific quality of the data is high and the database is

considered adequate to define the majority of the toxic effects that may result from exposure to mandestrobin. Mechanistic studies were also provided to support a proposed mode of action (MOA) for the thyroid effects in rats, and to investigate a possible effect of mandestrobin and its major metabolites on human estrogen and androgen receptors, as well as a possible effect of mandestrobin on the steroidogenesis of testosterone and estrogen. Acute oral toxicity and mutagenicity assays were also provided for five of the metabolites of mandestrobin observed in the rat.

Mandestrobin (S-2200TG, RS-Mandestrobin) is a racemic mixture of isomers R-mandestrobin (S-2167) and S-mandestrobin (S-2354). The metabolism and pharmacokinetics of mandestrobin were characterised after single oral gavage administration (racemic mixture at low and high doses and separate isomers at a low dose) and multiple oral gavage administration (14-day treatment at a low dose). All forms of mandestrobin, radiolabelled on either benzyl (mandestrobin, R-isomer and S-isomer) or phenoxy (mandestrobin) ring, were rapidly absorbed after a single gavage dose in rats. Area under the curve was proportionally higher at the low dose than at the high dose, indicating saturation of oral absorption at the high dose. Peak plasma levels were reached by 1.2 to 2.6 hours after an oral low dose, and 7.0 to 9.1 hours after a high dose. The labeled compound was widely distributed throughout tissues, but was detected primarily in the gastrointestinal (GI) tract, liver and kidney. Pancreas, uterus and ovaries also had higher levels than most other tissues. Mandestrobin isomers had similar tissue distribution, except the Sisomer was more prevalent in the GI tract than the R-isomer. There were no sex differences in tissue distribution, with the exception of the repeat-dose study in which females retained a larger portion of the dose in the cecum/large intestine/intestinal contents, and reached peak levels in tissues outside the GI tract later than males.

There was no evidence of accumulation of mandestrobin or its metabolites in tissues following repeated dosing. Clearance of a single dose of mandestrobin from plasma was almost complete by 120 hours post-dose. Clearance of the S-isomer was less rapid than the R-isomer. In the repeated-dose study, the majority of the administered dose was excreted by 14 days after last dose. Bile-cannulated rats excreted the radiolabelled compound more quickly than non-cannulated rats, suggesting enterohepatic recirculation. Excretion was consistent with all mandestrobin treatments. Fecal excretion via bile was the primary route, and renal excretion was also important. Excretion in expired air was negligible.

Mandestrobin was almost completely metabolized in rats via (1) oxidation followed by glucuronidation, or (2) demethylation followed by oxidation, or (3) oxidation followed by demethylation. There was no cleavage of benzyl and phenoxy rings. The major metabolites excreted were: 5-CA-S-2200-NHM and 4-OH-S-2200 in feces; 4-OH-S-2200-Glucuronide A in bile; and 5-CA-S-2200-NHM in urine. The same metabolites were identified for both isomers however at different proportions. Treatment with the R-isomer resulted in the following fecal/urinary metabolites in decreasing order of magnitude: 5-CA-S-2200-NHM, 5-CA-MCBX-NDM, 5-CA-2-HM-S-2200-NHM and 5-CA-S-2200-NDM; while S-isomer produced mostly 4-OH-S-2200 and 5-COOH-S-2200. In the multiple-dose study, fecal metabolites found at slightly higher levels in males than females included 5-CA-S-2200-NHM, 5-COOH-S-2200, 5-CA-2-HM-S-2200-NHM, 5-CA-2-HM-MCBX and 5-CA-MCBX-NDM. Results from cannulated rats suggest that 4-OH-S-2200 and its A glucuronide (GlucA) underwent enterohepatic circulation.

In acute toxicity testing, mandestrobin was demonstrated to be of low toxicity via the oral, dermal and inhalation routes in rats. Mandestrobin was minimally irritating to the eye and non-irritating to the skin of rabbits. Mandestrobin was not a dermal sensitizer in guinea pigs (Maximization Test).

The acute toxicity of the end-use products containing mandestrobin was low via the oral, dermal and inhalation routes in rats. The products were non-irritating to the skin and minimally irritating to the eyes of rabbits. They did not elicit a sensitization reaction in mice in the Local Lymph Node Assay.

In a short-term dermal toxicity study in rats, no adverse effects were noted at the limit dose. In repeated-dose dietary short-term studies, the liver and bile duct were the target organs of toxicity; the thyroid gland was also affected. Hepatic effects were noted in dogs and rats including hepatocellular hypertrophy accompanied by increased liver weight. At the lowest observed adverse effect level (LOAEL), increased incidences and severity of hepatocyte hypertrophy accompanied by centrilobular degeneration, liver agonal congestion and/or hemorrhage, large and dark liver, bile duct brown pigmentation and periductular inflammation, increased total cholesterol and serum liver enzymes and increased incidence and severity of thyroid follicular hypertrophy were observed. It was concluded based on the results of special mechanistic studies that, the effects on the thyroid gland were secondary to the liver effects, and were not observed without liver hypertrophy. The liver and thyroid effects were not observed in mice.

Rats and mice received a repeated dose of the test substance in the diet over most of their life span. Liver and thyroid gland effects were noted in rats at a lower dose than what was observed in short-term studies. Liver vacuolation was observed at the high dose in male rats and at the mid-high dose in female rats. Mice seemed less sensitive to the liver effects with adverse effects noted at the highest dose tested only. In long-term studies, the kidneys were a target organ of toxicity in male mice and female rats, which showed an increased incidence of corticomedullary mineralisation. According to scientific publications, the clinical significance of this observation is unclear.

However, this effect was observed in two species and different sexes in a dose-response manner, it was considered treatment-related and adverse. In female rats, renal corticomedullary mineralisation was accompanied by body weight effects, but liver and thyroid effects were only apparent at a higher dose.

MOA studies were provided to explain the liver and thyroid changes observed mostly in rats and dogs. Seven- and 14-day dietary treatments with mandestrobin caused an early induction of liver enzymes at the low dose, and, at mid-high and high dose, increased liver and thyroid gland weight and hepatocyte DNA synthesis, diffuse hepatocellular hypertrophy and diffuse thyroid follicular cell hypertrophy. In addition, effects on thyroxine, triiodothyronine, thyroid-stimulating hormone levels and T4-UDP-glucuronosyltransferase activity consistent with increased clearance of thyroxine were observed.

To evaluate reversibility of these findings, a 7-day dietary treatment followed by a 7-day recovery period was performed. The results demonstrated that most of these effects were transient and reversible. Taken together, the liver MOA studies demonstrated that the thyroid gland effects were secondary to the liver effects.

There was an increased incidence of benign ovary sex cord-stromal tumours in female rats in the long term toxicity study. Although this effect showed a dose-response relationship, it was considered as equivocal evidence of oncogenicity as this type of tumour is commonly observed in aging rats. The first tumour was observed late in the study in the control group (week 91) and later in treatment groups (week 98). No malignant tumors of this type were observed. The concern for this finding was tempered by its occurrence at a relatively high dose.

MOA studies were provided to investigate the ovary sex-cord stromal tumours observed in female rats. A reporter gene assay performed on HeLa cells derived from human uterine cervix carcinoma suggested that mandestrobin and its major metabolites (5-COOH-S-2200, 4-OH-S-2200, 5-CH2OH-S-2200 and 5-CA-S-2200-NHM) did not have agonistic or antagonistic effects on the human estrogen receptor alpha (hERa) and the human androgen receptor (hAR). Confidence in these assays was limited given that they were performed only once. The effect of mandestrobin on the production of testosterone and estradiol was evaluated in a well conducted steroidogenesis assay and the results were negative.

Although these mechanistic studies provide useful input to the mandestrobin toxicology database, they do not constitute an acceptable MOA to explain the incidence of ovary sex cord-stromal tumors in female rats.

Mandestrobin tested negative for genotoxicity in several assays, including bacterial reverse mutation assay, a forward mutation assay in mammalian cells, a chromosomal aberration assay, and an in vivo micronucleus assay.

In a two-generation dietary reproductive toxicity study in rats, there was no treatment-related effect on reproductive performance. Effects observed in parental animals were consistent with those reported in other repeated-dose dietary studies in rats and included increased liver weight and increased incidence of hypertrophy, brown pigmentation of the bile duct and periductular inflammatory cell infiltration. At a higher dose level, offspring of F_1 and F_2 generations exhibited reduced body weights beginning on postnatal day seven. In this study, there was no evidence of sensitivity of the young.

In a gavage developmental toxicity study in rats, developmental effects included increased incidences of litters with distended ureter or delayed skull ossification. These findings occurred at the limit dose of testing in the absence of maternal toxicity, thus providing evidence of sensitivity of the young. The incidences in control animals for these variations were on the low end of the historical control range. Also, clear signs of toxicity were observed at a lower dose in adult animals in other short-term studies. For these reasons the level of concern for these variations was low. No adverse effects were observed in rabbit dams exposed to mandestrobin via gavage or in their fetuses.

In an acute gavage neurotoxicity study in rats, decreased overall locomotor activity was noted in both sexes at 0-30 minutes on study day zero at the highest dose tested. This effect was considered an indicator of general toxicity rather that of specific neurotoxicity. In a 90-day dietary neurotoxicity study, decreased body weight and food consumption were noted, but no signs of neurotoxicity were noted at any time during the study. Overall, the evidence did not suggest that mandestrobin was a frank neurotoxicant.

In a 28-day dietary immunotoxicity study conducted in rats, there was an increase in spleen weight at a dose exceeding the limit dose of testing. Mandestrobin was not considered immunotoxic.

Oral acute studies and mutagenicity studies were also provided for some of the metabolites of mandestrobin. From the results of these studies, it was concluded that the metabolites 2-CH2OH-S-2200, 2-COOH-S-2200, 4-OH-S-2200, 5-COOH-S-2200 and De-Xy-S-2200 should be considered of comparable toxicity to the parent.

The code names for mandestrobin isomers and metabolites can be found in Appendix I, Table 2. Results of the toxicology studies conducted on laboratory animals with mandestrobin and its associated end-use products are summarized in Appendix I, Tables 3, 4, 5 and 6. The toxicology endpoints for use in the human health risk assessment are summarized in Appendix I, Table 7.

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. Mandestrobin is a new active ingredient pending registration for use in Canada. No human or domestic animal incidents involving the active ingredient mandestrobin have been reported to the PMRA and the applicant did not submit any additional data.

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 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 database contains the standard complement of required studies including gavage developmental toxicity studies in rats and rabbits and a two-generation dietary reproductive toxicity study in rats.

With respect to potential prenatal and postnatal toxicity, there was no indication of increased susceptibility of fetuses or offspring compared to parental animals in the reproductive toxicity study in rats and prenatal developmental toxicity study in rabbits. Minor developmental variations (increased incidence of delayed skull ossification and distended ureters) were observed in the rat developmental toxicity study in the absence of maternal toxicity. The concern

for this finding was low in view of the fact that it occurred at the limit dose of testing and toxicity was evident at this dose in other short-term studies (body weight effects, liver and thyroid effects). Furthermore, the incidences of these variations just slightly exceeded the laboratory historical data range.

Overall, the database is adequate for determining the sensitivity of the young. Concern for sensitivity was low and there were no serious endpoints were noted. On the basis of this information, the 10-fold *Pest Control Products Act* factor was reduced to 1-fold.

3.2 Acute Reference Dose (ARfD)

No acute endpoints of concern were identified in the toxicology database; therefore, an ARfD was not established.

3.3 Acceptable Daily Intake

To estimate risk of repeated dietary exposure, a no observed adverse effect level (NOAEL) of 27 mg/kg bw/day from the combined dietary chronic toxicity/carcinogenicity study in rats was selected for risk assessment. At the LOAEL of 135 mg/kg bw/day, decreased body weight and body weight gain and increased incidence of renal corticomedullary mineralization were observed in female rats. This study provides the lowest NOAEL in the database. Standard uncertainty factors of 10-fold for interspecies extrapolation and 10-fold for intraspecies variability were applied. As discussed in the *Pest Control Products Act* Hazard Characterization section, the *Pest Control Products Act* factor was reduced to 1-fold. The composite assessment factor (CAF) is thus 100.

The Acceptable Daily Intake (ADI) is calculated according to the following formula:

$$ADI = \frac{NOAEL}{CAF} = \frac{27 \text{ mg/kg bw/day}}{100} = 0.3 \text{ mg/kg bw/day of mandestrobin}$$

This ADI provides a margin of greater than 1500 to the dose resulting in increased incidence of benign ovary sex cord-stromal tumours in female rats and greater than 3300 to the dose resulting in increased incidence of fetal variations in the rat.

Cancer Assessment

As previously discussed, an increase in benign sex cord-stromal tumours in females in the rat oncogenicity study with mandestrobin was considered equivocal based on the weight of evidence. Overall, the endpoints selected for the non-cancer risk assessment are protective of these equivocal findings.

3.4 Occupational and Residential Risk Assessment

3.4.1 Toxicological Endpoints

Occupational exposures to S-2200 4 SC Fungicide, S-2200 4 SC Ag Fungicide, S-2200 4 SC VPP Fungicide, and S-2200 3.2 FS Fungicide are characterized as short- to intermediate-term in duration and are predominantly by the dermal and inhalation routes.

3.4.1.1 Short- and Intermediate-term Dermal

For short- and intermediate-term dermal risk assessment, the 28-day dermal toxicity study in rats was selected. No adverse effects were observed up to the limit dose of testing. A NOAEL of 1000 mg/kg bw/day was established.

The target margin of exposure (MOE) for these scenarios is 100, which includes uncertainty factors of 10-fold for interspecies extrapolation and 10-fold for intraspecies variability. The selection of this study and MOE is considered to be protective of all populations, including nursing infants and the unborn children of exposed female workers.

3.4.1.2 Short- and Intermediate-term Inhalation

For short- and intermediate-term inhalation scenarios, the reproductive toxicity study in rat was selected. For the inhalation scenarios, no repeat-dose inhalation toxicity study was available. Since an oral NOAEL was selected, an inhalation absorption factor of 100% was used in a route-to-route extrapolation. At the LOAEL of 166 mg/kg bw/day, increased brown pigmentation of the bile duct and periductular inflammatory cell infiltration in F_1 parental animals were observed. A NOAEL of 56 mg/kg bw/day was established.

The target MOE for these scenarios is 100, which includes uncertainty factors of 10-fold for interspecies extrapolation and 10-fold for intraspecies variability. The selection of this study and MOE is considered to be protective of all populations, including nursing infants and the unborn children of exposed female workers. A *Pest Control Products Act* factor of 1-fold was applied for the reasons outlined in the *Pest Control Products Act* Hazard Characterization section.

3.4.1.3 Non-dietary Oral Ingestion

For short-term incidental oral scenarios, the reproductive toxicity study in rat was selected. At the LOAEL of 166 mg/kg bw/day, increased brown pigmentation of the bile duct and periductular inflammatory cell infiltration in F_1 parental animals were observed. A NOAEL of 56 mg/kg bw/day was established.

The target MOE for this scenario is 100, which includes uncertainty factors of 10-fold for interspecies extrapolation and 10-fold for intraspecies variability. The selection of this study and MOE is considered to be protective of all populations, including nursing infants and the unborn children of exposed female workers. A *Pest Control Products Act* factor of 1-fold was applied for the reasons outlined in the *Pest Control Products Act* Hazard Characterization section.

3.4.1.4 Dermal Absorption

No in vivo or in vitro, rat or human, dermal absorption studies and/or data were submitted.

A dermal absorption value was not required for route-to-route extrapolation, as the NOAEL from the short-term 28-day dermal toxicity study is appropriate for assessing short- to intermediate-term exposures (no effects up to the limit dose of 1000 mg/kg bw/day).

3.4.2 Occupational Exposure and Risk

3.4.2.1 Mixer/loader/applicator Exposure and Risk Assessment

Individuals have potential for exposure to S-2200 4 SC Fungicide, S-2200 4 SC Ag Fungicide, and S-2200 4 SC VPP Fungicide during mixing, loading and application (ground and aerial), and exposure to S-2200 3.2 FS Fungicide during treatment of seeds (including mixing, loading, treating, and handling of treated seeds including bagging, sewing and stacking). Dermal and inhalation exposure estimates for workers were generated from PHED and surrogate worker exposure studies and data belonging to exposure task forces (AHETF, ARTF, SEEDTROPEX, and ORETF).

Exposures to workers mixing, loading and applying S-2200 4 SC Fungicide, S-2200 4 SC Ag Fungicide, S-2200 4 SC VPP Fungicide, and S-2200 3.2 FS Fungicide are expected to be short-to intermediate-term duration and to occur primarily by the dermal and inhalation routes. Exposure estimates were derived for mixers/loaders/applicators applying S-2200 4 SC Fungicide or S-2200 4 SC Ag Fungicide to canola (representing Crop Group 20A (Rapeseed Subgroup)) using groundboom (open-cab), aerial, and sprinkler chemigation equipment; to grape (representing Crop Group 13-07F (Small fruit vine climbing subgroup except fuzzy kiwifruit)) using airblast (open-cab) and sprinkler chemigation equipment; and to strawberry (representing Crop Group 13-07G (Low growing berry subgroup)) using groundboom and sprinkler chemigation.

Exposure estimates were derived for mixers/loaders/applicators applying S-2200 4 SC Fungicide or S-2200 4 SC VPP Fungicide to turfgrass (golf courses, lawns and landscape areas around residential, institutional, public, commercial and industrial buildings, recreational areas and to sod (sod farms)) using groundboom, backpack, manually- and mechanically-pressurized handwands, and turf handgun equipment. The exposure estimates are based on mixers/loaders/applicators wearing a long-sleeved shirt, long pants, and chemical-resistant gloves for foliar and turf uses. In addition, mixers and loaders for aerial application wear coveralls.

Exposure estimates were derived for seed treaters (including treaters, baggers, and cleaners) applying S-2200 3.2 FS Fungicide to canola seeds (representing Crop Group 20A (Rapeseed Subgroup)) and corn seeds (field corn, sweet corn, and popcorn) using closed transfer commercial treating equipment; and to legume vegetable seeds (represented by soybean and pea seeds) using closed transfer commercial or open transfer on-farm treating equipment. The exposure estimates are based on treaters wearing a long-sleeved shirt, long pants, and chemical-resistant gloves with seed treatment equipment cleaners also wearing chemical-resistant coveralls.

Chemical-specific data for assessing human exposures during pesticide handling activities were not submitted.

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

Exposure estimates were compared to the toxicological endpoints (NOAELs) to obtain the MOE; the target MOE is 100 for each of the dermal and inhalation routes. Routes of exposure cannot be combined since the toxic effects are different.

Table 3.4-1 Estimated Exposures of Mixers, Loaders, and Applicators From Foliar Applications of S-2200 4 SC Fungicide to Canola (Subgroup 20A), Grapes (Subgroup 13-07F), Strawberry (Subgroup 13-07G), and Turfgrass and Sod

Scenario	Application rate (g a.i./ha)	ATPD (ha/day)	Amount of a.i. handled per day 1 (kg a.i./day)	PHED (unless otherwise stated) dermal (μg a.i./kg a.i. handled)	Dermal Exposure 2 (mg a.i./kg bw/day)	Dermal MOE ³	PHED (unless otherwise stated) Inhalation (μg a.i./kg a.i. handled)	Inhalation Exposure ² (mg a.i./kg bw/day)	Inhalation MOE ⁴
GROUND App	olications								
Canola (covers long sleeves, an	strawberry (26 l nd no gloves)	na/day)): All	Liquids Op	en M/L ⁵ (long	pants, long sl	eeves, and g	loves); Open ca	ab groundboon	ı (long pants,
Farmer M/L/A		107	44.94	84.12	0.04725	21164	2.56	0.001438	38943
Custom M/L/A	420	360	151.2	84.12	0.15899	6290	2.56	0.004838	11575
Grapes: All Lie	quids Open M/L	(long pants, l	long sleeves	, and gloves); o	pen-cab airbl	ast (long pa	nts, long sleeve	s, and gloves)	
Farmer, Custom M/L/A	420	20	8.4	3820.44 ^b	0.40115	2493	10.68 ^b	0.001121	49955
Turfgrass and	sod: Backpack e	quipment; Al	ll Liquids O	pen M/L/A (lo	ng pants, long	sleeves, and	d gloves)		
M/L/A	472	0.1875	0.0885	5445.85	0.00602	166113	62.1 ^a	0.000069	782609

Scenario	Application rate (g a.i./ha)	ATPD (ha/day)	Amount of a.i. handled per day 1 (kg a.i./day)	PHED (unless otherwise stated) dermal (µg a.i./kg a.i. handled)	Dermal Exposure ² (mg a.i./kg bw/day)	Dermal MOE ³	PHED (unless otherwise stated) Inhalation (μg a.i./kg a.i. handled)	Inhalation Exposure ² (mg a.i./kg bw/day)	Inhalation MOE ⁴
Turfgrass and	sod: Turf handg	un; All Liqui	ds Open M	/L/A (long pant	s, long sleeve	s, and glove	5)		
Custom M/L/A	472	2	0.944	785 °	0.00926	107991	4 °	0.000047	1148936
Turfgrass and gloves)	sod: All Liquids	Open M/L (l	ong pants, l	ong sleeves, and	l gloves); Op	en cab grou	ndboom (long p	oants, long sleev	ves, and no
Golf course M/L/A	472	16	7.552	84.12	0.00794	125945	2.56	0.000242	223140
Sod farm M/L/A		30	14.16	84.12	0.01489	67159	2.56	0.000453	119205
Turfgrass and	sod: Mechanical	ly-pressurize	d handgun;	All Liquids Op	en M/L/A (lo	ong pants, lo	ng sleeves, and	gloves)	
Custom M/L/A	472	4.75 ^d	2.242	5585.49	0.15653	6389	151	0.004232	12760
AERIAL (can	ola only): All Liq	uids Open M	/L (coverall	ls, long pants, lo glove		nd gloves);	Applicator (lon	g pants, long sl	eeves, and no
Farmer, Custom M/L	420	400	168	32.77	0.06882	14531	1.6	0.003360	16667
Farmer, Custom A		400	168	9.66	0.02029	49285	0.07	0.000147	380952
CHEMIGATI	ON (canola, grap	es, strawber	ry): All Liqu	uids Open M/L	(long pants, l	ong sleeves,	and gloves)		
Sprinkler Chemigation M/L	420 mount of a.i. hand	140	58.8	51.14	0.03759	26603	1.6	0.001176	47619

. Amount of a.i. handled per day calculated using the maximum application rate \times ATPD

2. Exposure was calculated using the amount of a.i. handled per day × unit exposure value/body weight (80 kg) (dermal or inhalation). No dermal absorption value required; assumed 100% inhalation systemic absorption

3. Estimates of dermal exposure for M/L and A were compared to a dermal NOAEL of 1000 mg/kg bw/day, target MOE = 100.

4. Estimates of inhalation exposure for M/L and A were compared to a NOAEL of 56 mg/kg bw/day, target MOE = 100.

5. Farmer mixer/loader is sufficient to represent the expected exposures of workers for chemigation

a. Moderate inhalation rate

b. Revised open-cab airblast unit exposures

c. ORETF (commercial) professional low pressure nozzle gun sprayer. 3800L/day default volume per day \div (min. 8 L spray volume/100 m² × $10000 \text{ m}^2/\text{ha}$)

Table 3.4-2 Risk Estimates for Workers Treating Seeds in Commercial Facilities with S-2200 3.2 FS Fungicide

Washen to sh	Unit exposure $(\mu g/kg a.i. handled)^1$		Appl. rate	Seed treated	Dermal Exposure	Dermal	Inhalation	Inhalation	
Worker task	Dermal	Inhalation	(kg a.i./ kg seed)	(kg seed/ day) ³	(mg/kg bw/day) ⁴	MOE ⁵	(mg/kg bw/day) ⁴	MOE ⁵	
		Close	ed mix, load,	transfer comm	ercial facilities				
			Corn (fi	eld, sweet, and	pop)				
Treater	0.88	0.016	0.00006	125,000	8.25E-05	1.21E+07	1.50E-06	3.73E+07	
Bagger	17.67	0.89	0.00006	125,000	1.66E-03	6.02E+05	8.34E-05	6.71E+05	
Cleaner (normalized)*	17.1	5.1	0.00006	-	1.29E-03	7.75E+05	3.83E-04	1.46E+05	
Treater + Cleaner †	-	-	0.00006	-	1.37E-03	7.30E+05	3.85E-04	1.45E+05	
Canola									
Treater	0.88	0.016	0.0001	67,000	7.37E-05	1.36E+07	1.34E-06	4.18E+07	

Worker task	Unit exposure $(\mu g/kg a.i. handled)^1$		Appl. rate	Seed treated	Dermal Exposure	Dermal	Inhalation	Inhalation
	Dermal	Inhalation	(kg a.i./ kg seed)	(kg seed/ day) ³	(mg/kg bw/day) ⁴	MOE ⁵	(mg/kg bw/day) ⁴	MOE ⁵
Bagger	17.67	0.89	0.0001	67,000	1.48E-03	6.76E+05	7.45E-05	7.52E+05
Cleaner (normalized)*	17.1	5.1	0.0001	-	2.15E-03	4.65E+05	6.38E-04	8.78E+04
Treater + Cleaner †	-	-	0.0001	-	2.22E-03	4.50E+05	6.39E-04	8.76E+04
				Soybean				
Treater	0.88	0.016	0.0001	63,000	6.93E-05	1.44E+07	1.26E-06	4.44E+07
Bagger	17.67	0.89	0.0001	63,000	1.39E-03	7.19E+05	7.01E-05	7.99E+05
Cleaner (normalized)*	17.1	5.1	0.0001	-	2.15E-03	4.65E+05	6.38E-04	8.78E+04
Treater + Cleaner †	-	-	0.0001	-	2.22E-03	4.50E+05	6.39E-04	8.76E+04
			Legumes	(other than soy	bean)			
Treater	0.88	0.016	0.0001	216,000	2.38E-04	4.20E+06	4.32E-06	1.30E+07
Bagger	17.67	0.89	0.0001	216,000	4.77E-03	2.10E+05	2.40E-04	2.33E+05
Cleaner (normalized)*	17.1	5.1	0.0001	-	2.15E-03	4.65E+05	6.38E-04	8.78E+04
Treater + Cleaner †	-	-	0.0001	-	2.39E-03	4.18E+05	6.42E-04	8.72E+04

* Normalized Cleaner unit exposure ($\mu g/g a.i./100kg seed/day$)

= 240.02 μ g/day / 14.04 g a.i./100 kg seed, application rate in the exposure study

† Assuming that a worker both treats and cleans in the same workday; using normalized cleaner, and considered conservative

1. For closed transfer commercial facilities, the arithmetic mean values were used from the surrogate exposure study; long-sleeved shirt and long pants (plus nitrile gloves for mixer/loader/calibrators); plus Tyvek coveralls and nitrile gloves for cleaners.

2. Highest PMRA-supported application rate on the label

3. Default seed treatment throughput for beans (PMRA seed treatment database)

4. Exposure (mg/kg bw/day) = unit exposure (μ g/kg a.i. handled) × application rate (g a.i./100 kg seed) × seeds treated (kg seed/day) × 0.001(mg/ μ g) / body weight (kg bw)

Cleaner Normalized Exposure (mg/kg bw/day) = (unit exposure (μ g/g a.i./100 kg seed) × (0.0001 or 0.00006) kg a.i./kg seed × 100 kg seed × 1000 g/kg)/(80 kg bw × 1000 μ g/mg)

5. Margin of Exposure (MOE) = NOAEL/Exposure; NOAEL = 1000 mg/kg bw/day, target MOE = 100; inhalation NOAEL = 56 mg/kg bw/day, target MOE = 100

Table 3.4-3 Risk Estimates for On-Farm Treatment (Mobile Treaters) of Legume Seeds with S-2200 3.2 FS Fungicide

Unit exposure values ^{1,2} (µg/kg a.i.)		Amount of a.i.	Dermal	Devel	Inhalation	
Dermal	Inhalation	handled per day	Exposure ^{4,5} (mg/kg	Dermal MOE ⁶	Exposure ^{4,5} (mg/kg	Inhalation MOE ⁶
90 th percentile	90 th percentile	(kg a.i./day)	bw/day)		bw/day)	
141.9	7.825	1.9	0.00337	296736	0.00019	284211

1. Surrogate mixer/loader/applicator study

2. Workers wearing a single layer (a long-sleeved shirt, long pants, and chemical-resistant gloves)

3. Amount a.i. handled per day based on peas (except cowpea and field pea; representing legumes, including soybean) = application rate (0.0001 kg a.i./kg seed) \times seeding rate (190 kg seed/ha) \times area planted per day (100 ha/day).

4. No adjustment for dermal absorption required; both dermal and inhalation exposures considered 100% systemically available

5. Daily exposure (mg/kg bw/day)

= (Unit exposure \times Amount of a.i. handled/day) / (80 kg bw \times 1000 µg/mg)

6. Margin of Exposure (MOE) = NOAEL (route-specific)/Exposure; short- to intermediate term durations, target MOE = 100

Table 3.4-4 Exposure and Risk Estimates for Farmers Treating and Planting LegumeSeeds with S-2200 3.2 FS Fungicide

	PPE: single layer + chemical-resistant gloves									
Cross	Unit exposure (µg/kg bw/day) ¹		Seed treated and	Rate	Dermal	Dermal MOE	Inhalation	Inhalation MOE		
Сгор	Dermal	Inhalation	Planted ² (kg seed/day)	(kg a.i./kg seed)	Exposure ³ (mg/kg bw/day)	(target = 100) ⁴	Exposure ³ (mg/kg bw/day)	(target = 100) ⁴		
Legume vegetable seeds	407.34	223.03	19000	0.0001	0.0096743	103366	0.005297	10572		

1 Unit exposures in the surrogate seed treatment study for workers wearing a long-sleeved shirt, long pants, and chemical-resistant gloves 2 From the Seed Treated/Planted Per Day tables (2009)

3 Exposure = (Unit exposure × Seed treated/planted × Rate)/(1000 μ g/mg × 80 kg bw)

4 Dermal NOAEL = 1000 mg/kg bw/day, target MOE= 100; inhalation NOAEL = 56 mg/kg bw/day, target MOE = 100

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 S-2200 4 SC Fungicide, S-2200 4 SC Ag Fungicide, S-2200 4 SC VPP Fungicide, and workers planting seeds treated with S-2200 3.2 FS Fungicide. Given the nature of activities performed, the primary route of exposure for workers re-entering treated areas would be through the dermal route, and through the dermal and inhalation routes for planting treated seeds. The durations of exposure are considered to be short-term in duration when entering treated crops, turf, and handling and planting seeds.

Dermal exposures to workers entering treated areas (Table 3.4-5) are estimated by coupling dislodgeable foliar residue values for foliar treatments, or turf transferable residue values for turf, with activity-specific transfer coefficients. Chemical-specific dislodgeable foliar residue data were not submitted. Therefore, a default dislodgeable foliar residue value of 25%, or default turf transferable residue value of 1% of the application rate on the day of the final treatment, was used in the exposure assessment.

Dermal and inhalation exposures of workers planting treated seeds (Table 3.4-6) are estimated by coupling unit exposures for planting treated seeds with the amount of seeds planted, and size of areas planted.

No dermal absorption value was required since the dermal endpoint was based on a dermal study. Inhalation absorption is considered 100%. Exposures were normalized to mg/kg bw/day by using 80 kg adult body weight.

Exposure estimates were compared to the toxicological end point to obtain the MOE; the target MOE is 100.

Table 3.4-5 Postapplication Occupational Exposure and Risk Estimates for Re-Entry TasksPerformed in Crops Treated with S-2200 4 SC Fungicide

Сгор	Tasks	Maximum Application Rate (µg/cm ²)	Number of Applications at Maximum Rate	Minimum Application Interval (days)	Dislodgeable Foliar Residue or Turf Transferable Residue (µg/cm ²) ^A ; DALA = 0	Highest of Crop Group Transfer Coefficients ^B (cm ² /hour)	Dermal Exposure (mg/kg bw/day)	MOE
Canola	Scouting, solid stand	4.2	1		1.050	1100	0.1155	8658
	Girdling, turning	4.2	3	10	1.544	19300	2.9799	336
Grape	Hand harvesting, leaf pulling, tying and training	4.2	3	10	1.544	8500	1.3124	762
Strawberry	Hand harvesting	4.2	4	7	1.907	1100	0.2098	4766
Turfgrass/ Sod	Golf course maintenance: transplanting; sod slab harvesting					6700	0.0409	24450
	Golf course/Sod mowing, watering, etc.	4.72	4	14	0.061	3500	0.0214	46729
	Aerating, scouting, fertilizing, hand pruning, mechanical weeding, seeding					1000	0.0061	163934

DALA = Days after last application

^A Use default dislodgeable foliar residue for canola, grapes, and strawberry; turf transferable residue for turfgrass

^B PMRA Transfer Coefficients based on ARTF database (2012)

^C Dermal NOAEL = 1000 mg/kg bw/day, target MOE= 100

Table 3.4-6 Exposure and Risk Estimates for Planting Canola, Corn, Soybean, and Pea Seeds Treated with S-2200 3.2 FS Fungicide

Scenario	Unit exposure (µg/kg a.i. handled) ¹		Seeds planted per day ² (kg)	Appl. rate (kg a.i./kg seed)	Amount of a.i. handled per day ³ (kg a.i./day)	Exposure ⁴ (mg/kg bw/day)		MOE ⁵	MOE ⁵
Planting	Dermal	Inhalation	(ng)			Dermal	Inhalation	Dermal	Inhalation
canola			600	0.0001	0.06	0.0011	0.00006	909091	933333
corn			1350	0.00006	0.081	0.0015	0.00008	666667	700000
Legumes (excluding soybean)	1515	82.83	19000	0.0001	1.9	0.0360	0.00197	27778	28426
soybean			9000	0.0001	0.9	0.0170	0.00093	58824	60215

Note: Exposure estimates based on passive dosimetry study during planting of treated maize corn.

¹ Unit exposure values for planters wearing a single layer + chemical-resistant gloves and using closed-cab planting equipment

² Seed Treated Planted Per Day Table

³ Kg a.i. handled per day = kg seed treated per day \times application rate (kg a.i./kg seed)

⁴ Exposure (mg/kg bw/day) = <u>Unit exposure (μ g/kg a.i. handled per day) × kg a.i. handled per day × 0.001 mg/ μ g</u>

80 kg bw

⁵Dermal NOAEL = 1000 mg/kg bw/day, target MOE= 100; inhalation NOAEL = 56 mg/kg bw/day, target MOE = 100

Risks to workers entering treated crops, turfgrass and sod, are not of concern. The default restricted entry interval (REI) of 12 hours is adequate to protect workers. Risks to workers planting treated seeds are not of concern when following the treated seed label precautions.

3.4.3 Residential Exposure and Risk Assessment

3.4.3.1 Handler Exposure and Risk

No homeowner applicator scenario is proposed.

3.4.3.2 Postapplication Exposure and Risk

3.4.3.2.1 Postapplication Dermal Exposures

Section 3, Lawns and Turf, of the 2012 United States Environmental Protection Agency (USEPA) Residential Standard Operating Procedure (SOP) was used to determine postapplication exposures to people re-entering treated lawns in residential areas.

The duration of exposure is considered to be short-term (up to 30 days) for golfing and for residential turf activities. No product-specific turf transferable residue (TTR) study was provided; therefore, the current default peak (day 0) DFR value of 1% of the application rate, transfer coefficients, exposure time, and algorithms were based on the SOP for turf. Body weights used were 80 kg for adults, 57 kg for youth, 32 kg for children (6<11years old), and 11 kg for children (1<2 years old).

Lifestage (years of age)	$\frac{TTR_t}{(\mu g/cm^2)}$	Transfer Coefficient (cm ² /hour)	Hours of Exposure (hour/day)	Exposure (mg/day)	Absorbed Dose (mg/kg/day)	Dermal MOE
High Contact Lav	wn Activities					
Adult 18+		180,000	1.5	12.74	0.1593	6277
Youth 11 <16	0.05	148,000	1.3	9.08	0.1593	6277
Children 1 <2		49,000	1.5	3.47	0.3154	3171
Mowing Turf						
Adult 18+		5,500	1	0.2596	0.003245	308166
Youth 11 <16	0.05	4,500	1	0.2124	0.003726	268362
Golfing (Treated	greens, tees, a	and fairways)			-	<u>.</u>
Adult 18+		5,300	4	1.0006	0.012508	79949
Youth 11 <16	0.05	4,400	4	0.8307	0.01457	68615
Children 6 <11		2,900	4	0.5475	0.01711	58445

Table 3.4-7 Postapplication Dermal Exposure and Risk Estimates for Re-Entry onto TurfTreated with S-2200 4 SC Fungicide

3.4.3.2.2 Child Incidental Oral Ingestion

A child may place a hand in his/her mouth a number of times, as well as place an object in their mouth a number of times during a certain period of time. Each of these events could result in a potential transfer of residue, but could also result in a soil ingestion event as soil may be present on the hand or object during mouthing (USEPA Residential SOP 2012). Table 3.4-8 shows estimates of the exposures for each scenario. Risks from each of the three non-dietary oral ingestion scenarios are considered not to be of concern (greater than the target MOE of 100).

Table 3.4-8 Postapplication Residential Child Oral Exposure and Risk Assessment for Entry onto Turfgrass Treated with S-2200 4 SC Fungicide

Lifestage (years of age)	Route of ingestion	Residue loading (mg/cm ²)	Oral Exposure (mg/kg bw/day)	MOE
	Hand-to-mouth	0.00069	0.0065	8660
Children 1 <2	Object-to-mouth	0.04720	0.00020	285171
	Incidental soil ingestion	3.1624 (µg/g)	1.4E-05	3.9E+06

3.4.3.2.3 Residential Aggregate Assessment

Adults and youth are considered to conduct multiple postapplication activities on treated turf. The public (including children) may enter treated recreational turf areas on the day of treatment. Table 3.4-9 shows the activities that are considered to co-occur when adults and youth are active on treated turf. Golfing is not part of the aggregate assessment. Risks for adults and youth are not considered to be of concern (greater than the target MOE of 100).

Table 3.4-9 Residential Dermal Aggregate Risks of Entry onto Turfgrass Treated withS-2200 4 SC Fungicide (or S-2200 4 VPP Fungicide)

	Scenar Absorbed Dose (mg/kg by	Dermal MOE	
Lifestage (years of age)	High Contact Lawn Activities ¹	Mowing Turf ¹	Aggregate ²
Adult 18+	0.1593	0.003245	6154
Youth 11 <16	0.1593	0.003726	6135

1. Exposures taken from Table 3.4-7

2. Based on SPN2003-04; target MOE is 100

Postapplication exposure scenarios that are likely to co-occur over a short term are the dermal and hand-to-mouth scenarios (2012, USEPA Residential SOP, Turf and Lawns) (Table 3.4-10). However, there is no common toxic effect for aggregation of dermal and oral exposures; therefore, the dermal route is not included. The PMRA also includes dietary exposure in this aggregate because these exposures are by the oral route. Aggregate oral exposure risks to children are not considered to be of concern.

Table 3.4-10Aggregate Oral (Non-Dietary and Dietary) Ingestion of MandestrobinResidues by Children 1<2 yrs</td>

Route of ingestion	Oral Exposure (mg/kg bw/day)
Hand-to-mouth ¹	0.0065
Chronic dietary exposure ²	0.027593
Aggregate MOE 3 (target = 100)	1644

1. From Table 3.4-8; hand-to-mouth activity is not expected to under-estimate non-dietary intake of children 1<2 years of age

2. From Dietary Exposure Assessment (food + water)

3. Using PMRA SPN2003-04

3.4.3.2.4 Pick-Your-Own Acute Aggregate Exposure Assessment

Strawberry and lowbush blueberry are considered to be pick your own (PYO) crops. The public may be exposed to residues of S-2200 4 SC Fungicide on these crops at PYO operations. This exposure is expected to be acute and to occur by the dermal and oral routes. The acute dermal exposures for adults, youth, and children in PYO facilities are addressed by the dermal exposures of workers re-entering treated fields for hand harvesting, and not considered to be of concern. Exposure by the oral route is addressed by the dietary exposure assessment. Therefore, the public picking strawberries and lowbush blueberries at PYO facilities is acceptable at the pre-harvest interval on the label.

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 Residues in Plant and Animal Foodstuffs

The residue definition for risk assessment and enforcement in plant products is mandestrobin. The data gathering/enforcement analytical method is valid for the quantitation of mandestrobin (S-2200) residues in crops. The residue definition for risk assessment and enforcement in animal commodities will be determined once a livestock feeding study and enforcement method are submitted to the Agency.

The residues of mandestrobin are stable in rapeseed (seed, oil and meal), lettuce, barley (grain, straw), soybean seed, and corn (grain, forage, K+CWHR) for up to 12 months; in corn stover for up to 9 months; in grapes for up to 8 months; in grape juice for up to 7 months; and in strawberries and raisins for up to 5 months, when stored in a freezer at -18°C. Freezer storage stability studies are on-going for rapeseed (seed, oil, and meal), strawberries and grapes (fruit, raisins). Mandestrobin residues concentrated in the following processed commodities: raisin (1.9-fold), and grape juice (1.4-fold). Quantifiable residues are not expected to occur in livestock

matrices with the current use pattern. Crop field trials conducted throughout Canada and the United States using end-use products containing mandestrobin at approved rates in or on corn, grapes, rapeseed, soybeans, and strawberries are sufficient to support the proposed maximum residue limits.

3.5.2 Concentrations in Drinking Water

Estimated environmental concentrations (EECs) of mandestrobin combined residue in potential drinking water sources (groundwater and surface water) were generated using computer simulation models. An overview of how the EECs are estimated is provided in the PMRA's Science Policy Notice SPN2004-01, *Estimating the Water Component of a Dietary Exposure Assessment*. EECs of mandestrobin 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 average concentrations in the top one metre of the water table. EECs of mandestrobin combined residue in surface water were calculated using the SWCC model, which simulates 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 vulnerable drinking water source, a small reservoir.

Drinking water modelling of mandestrobin includes four transformation products, 5-COOH-S-2200, MCBX, S-2200-OR and S-2200-ORC as well as mandestrobin itself. Degradation rates for drinking water modelling were calculated from the overall degradation of these five compounds, and for groundwater modelling, the more conservative K_d from 5-COOH-S-2200 was used.

A Level 1 drinking water assessment was conducted using conservative assumptions with respect to environmental fate, application rate and timing, and geographic scenario. The Level 1 EEC estimate is expected to allow for future use expansion into other crops at this application rate. Table 3.5-1 lists the application information and main environmental fate characteristics used in the simulations. Twenty-seven initial application dates between May and September were modelled. The model was run for 50 years for all scenarios. The largest EECs of all selected runs are reported in Table 3.5-2 below.

Type of Input	Parameter	Value
Application Information	Crop(s) to be treated	Beans, Blueberries, Canola, Chick peas, Corn, Flaxseed, Gooseberries, Grapes, Lentils, Lupins, Mustard seed, Peas, Radish, Soybeans, Strawberries, Turf
	Maximum allowable application rate per year (g a.i./ha)	1890
	Maximum rate each application (g a.i./ha)	472 (turf) and 420 (for other crops)
	Maximum number of applications per year	4

Table 3.5-1 Major groundwater and surface water model inputs for Level 1 assessment of mandestrobin

Type of Input	Parameter	Value
	Minimum interval between applications (days)	14 (turf) and 7 (for other crops)
	Method of application	Field sprayer or aerial application equipment
Environmental Fate	Hydrolysis half-life at pH 7 (days)	stable
Characteristics	Photolysis half-life in water (days)	4.5 days (mandestrobin only) 10.7 days (mandestrobin with four transformation products)
	Adsorption K _{OC} or K _d (mL/g)	Mandestrobin K_{OC} of 365 used for surface water modelling; 5-COOH-S2200 K_d of 1.24 used for groundwater modelling
	Aerobic soil biotransformation half-life (days)	264 days @ 20°C for eco 306 days @ 20°C for drinking water
	Aerobic aquatic biotransformation half-life (days)	693 days @ 20°C for eco 4320 days @ 20°C for drinking water
	Anaerobic aquatic biotransformation half-life (days)	2345 days @ 20°C for eco 6335 days @ 20°C for drinking water

Table 3.5-2 Level 1 estimated environmental concentrations of mandestrobin in potential drinking water sources

-	Groundwater EEC (μg a.i./L)		Surface Water EEC (µg a.i./L) Reservoir				
	Daily ¹	Yearly ²	Daily ³	Yearly ⁴			
Mandestrobin							
combined residue	225	225	82	21			
90^{th} perce 90^{th} perce							

3.5.3 Dietary Risk Assessment

Chronic (non-cancer) dietary risk assessments were conducted using the Dietary Exposure Evaluation Model (DEEM–FCIDTM NHANES, Version 4.02), which incorporates food consumption data from the National Health and Nutritional Examination Survey, What We Eat in America (NHANES/ WWEIA) dietary survey for the years 2003-2008 available through CDC's National Center for Health Statistics (NCHS).

3.5.3.1 Chronic Dietary Exposure Results and Characterization

The following criteria were applied to the basic chronic non-cancer analysis for mandestrobin: 100% crop treated, default processing factors and proposed MRLs of rapeseed (CSG 20A), grape (CSG 13-07F), strawberry (CSG 13-07G, except cranberry), soybean (CG 6, except cowpea and field pea) and corn. The basic chronic dietary exposure from all supported mandestrobin food uses (alone) for the total population, including infants and children, and all representative population subgroups is less than 1.3% of the acceptable daily intake (ADI). Aggregate exposure from food and drinking water is considered acceptable. The PMRA estimates that chronic dietary exposure to mandestrobin from food and drinking water is 2.8% (0.0084 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 9.2% (0.028 mg/kg bw/day) of the ADI.

3.5.3.2 Acute Dietary Exposure Results and Characterization

No appropriate endpoint attributable to a single dose for the general population (including children and infants) was identified.

3.5.4 Aggregate Exposure and Risk

The aggregate risk for mandestrobin consists of exposure from food and drinking water sources.

3.5.5 Maximum Residue Limits

Table 3.5-3 Proposed	Maximum	Residue	Limits
----------------------	---------	---------	--------

Commodity	Recommended Maximum Residue Limit (ppm)
Raisins	7.0
Small fruit vine climbing (Crop Subgroup 13-07F, except fuzzy kiwifruit)	5.0
Low growing berry (Crop Subgroup 13-07G, except cranberry)	3.0
Rapeseed (Crop Subgroup 20A)	0.5
Legume vegetables (succulent or dried) (Crop Group 6, except cowpea and field pea), corn (field, popcorn, sweet)	0.02

Maximum Residue Limits (MRLs) are proposed for each commodity included in the listed crop groupings in accordance with the Residue Chemistry Crop Groups webpage in the Pesticides and Pest Management section of Health Canada's website.

For additional information on maximum residue limits in terms of the international situation and trade implications, refer to Appendix II.

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, 8 and 9.

4.0 Impact on the Environment

The active substance mandestrobin is a racemic mixture of the two isomers (S-2200 *R*-isomer and S-2200 *S*-isomer) in a ratio of 50:50. The results presented in this section are for the mixture unless stated otherwise.

4.1 Fate and Behaviour in the Environment

The fate and behaviour of mandestrobin and its major transformation products are summarized in Appendix I, Table 10. The chemical name and structure of transformation products formed in the environment, as well as a summary of their occurrence in environmental fate studies, are presented in Appendix I, Table 11.

Mandestrobin is introduced in the environment when it is applied as a seed treatment, foliar spray and chemigation to a variety of seeds, field crops and non-crops. Mandestrobin could reach the soil upon application or through wash-off from the leaves. Abiotic transformation processes are not expected to contribute significantly to the dissipation of mandestrobin in soil as this compound is stable to hydrolysis and phototransformation. Biotransformation may be an important route of dissipation for mandestrobin in the terrestrial environment.

In both aerobic and anaerobic soils, mandestrobin transformed to the major products 5-COOH-S-2200, DX-CA-S-2200, 2-CONH₂-S-2200, 5-CONH₂-S-2200, MCBX and De-Xy-S-2200. Mandestrobin is slightly persistent to persistent in aerobic soil and will also persist in anaerobic soil.

Based on results from adsorption studies, mandestrobin exhibited medium to low mobility in soil, transformation products 5-COOH-S-2200 and 2-COOH-S-2200 exhibited very high to low mobility. The groundwater ubiquity scores (GUS) calculated for mandestrobin and its transformation products 5-COOH-S-2200 and 2-COOH-S-2200 based on their persistence and mobility indicate that mandestrobin is a borderline leacher and that 5-COOH-S-2200 and 2-COOH-S-2200 are probable leachers. In field dissipation studies, no residues of mandestrobin and its transformation products were detected beyond a 30 cm soil depth. Mandestrobin will not carry over to the next growing season.

Mandestrobin could reach surface water through spray drift and runoff. Once in the aquatic environment, mandestrobin is not expected to hydrolyze, but will undergo rapid phototransformation in clear shallow water to the major transformation products S-2200-OR, S-2200-ORC, S-2200-PR and CO₂. Biotransformation is not a primary route of transformation for mandestrobin in water and large amounts of mandestrobin were shown to partition into sediment. MCBX and 5-COOH-S-2200 were the only major biotransformation products identified in aquatic systems. Mandestrobin does not accumulate to a large degree in fish and depuration occurs rapidly.

Residues of mandestrobin and its transformation products are not expected to be found in air. Mandestrobin exhibits low volatility based on its low vapour pressure.

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 concentrations are concentrations of pesticide in various environmental media, such as food, water, soil and air. The EECs are derived using standard models which 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 (in other words, 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 in most cases except for bees and certain beneficial arthropods where the level of concern is 0.4 and 2, respectively). 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 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.

4.2.1 Risks to Terrestrial Organisms

A risk assessment for mandestrobin and its transformation products was undertaken for terrestrial organisms based on available toxicity data. A summary of terrestrial toxicity data is presented in Appendix I, Table 12. Results of the accompanying risk assessment are presented in Appendix I, Tables 13-15.

Earthworms: Earthworms could be exposed to mandestrobin when this compound reaches the soil upon application. The expected environmental concentration is, therefore, calculated based on a direct application of mandestrobin to bare soil at the maximum seasonal labelled application rate.

At levels higher than those expected in the Canadian environment, the acute exposure of earthworms to mandestrobin has been shown to cause mortality and decreased weight. Chronic exposure to mandestrobin also resulted in significant reduction in reproduction and body weight. However, given the low expected environmental concentration, risk quotients calculated for acute and chronic exposure to mandestrobin do not exceed the level of concern. The 5-COOH-S-2200 and 2-COOH-S-2200 transformation products were not acutely toxic to earthworms. There is therefore no concern from acute exposure to these transformation products.

Bees (pollinators): Foraging bees could be exposed directly to mandestrobin spray droplets during application or to mandestrobin residues found on the surface of leaves (contact exposure). Foraging bees could also be exposed to mandestrobin through the ingestion of pollen and nectar contaminated from direct spray or the systemic movement in the plant (oral exposure). In addition, brood may be exposed to mandestrobin and its transformation products as foraging bees bring contaminated pollen and nectar back to the hive.

In laboratory tests, mandestrobin was non-toxic to adult honey bees when applied directly on bees or through diet consumption. When honey bee larvae were incubated with a diet treated with mandestrobin, no mortality was observed at the highest dose tested. All risk quotients calculated for mandestrobin did not exceed the level of concern.

Beneficial arthropods: The risk assessment for beneficial arthropods considers that the main route of exposure for these non-target organisms is from contact with treated plant material both on the treated area (from direct spray on the crop) and at the margins of the treated field (from spray drift). The expected concentration of mandestrobin residues on foliage within the treated field is calculated as the cumulative application rate, which takes into account the maximum labelled application rate, the application interval and the dissipation of the compound on the surface of the leaves.

In laboratory tests carried out with freshly dried residues on a glass plate, S-2200 25SC Fungicide caused no statistically significant adverse acute effects on the parasitic wasp and predatory mite. Significant effects on the reproduction of the parasitic wasp were observed, but at levels higher than those expected in the environment. The screening level risk quotients calculated for both the predatory mite and the parasitic wasp were below the level of concern.

Birds and mammals: At the highest dose, acute oral exposure to mandestrobin caused no mortality in bobwhite quail and canary. When mandestrobin was administered in the diet no adverse effects to bobwhite quail and mallard duck were observed. In reproductive tests, no subchronic or reproductive effects were observed at the highest concentrations tested for either bobwhite quail or mallard duck.

Laboratory studies indicated that mandestrobin was not acutely toxic to rats. Among chronic effects observed in a two-generation dietary reproduction study with rats, was a decreased body weight in offspring.

For birds and mammals, risk quotients calculated at the screening level for mandestrobin did not exceed the level of concern on an acute or reproductive basis for both foliar application (up to 4 times a year for a maximum total rate of 1888 g a.i./ha/year) and seed treatment (up to 10 g a.i./100 kg seeds or 0. 9 g ai/ha).

Non-target plants: For the risk assessment, the cumulative application rate is compared to plant toxicity endpoints. The cumulative application rate takes into account the maximum labelled application rate, the application interval and the dissipation of the compound on the surface of the leaves. For the off-field assessment, the rate is adjusted according to the projected drift deposition at one metre downwind from the site of application.

The toxicity of S-2200 4 SC Fungicide to non-target plants was determined through vegetative vigour and seedling emergence assays using standard crop species. No significant adverse effects were observed in any plant species at the highest application rate tested (560 g a.i./ha). However, given that the study was conducted at the one-time application rate of 560 g a.i./ha, whereas the seasonal maximum application rate (4×472 g a.i./ha) is 1888 g a.i./ha, the screening risk quotient exceeded the level of concern. The risk to terrestrial plants was, thus, further assessed.

For an ASAE (American Society of Agricultural Engineers) 'coarse' droplet size, the maximum spray drift deposition at one metre downwind from the point of application is 17% for aerial application. Based on the risk quotients using the off-field EECs from drift, the level of concern for terrestrial vascular plants is not exceeded.

There is still uncertainty with regards to potential effects on the vegetative vigour and seedling emergence of plants after exposure to multiple applications. Due to this uncertainty, it was determined that drift mitigation measures are necessary.

4.2.2 Risks to Aquatic Organisms

A risk assessment of mandestrobin, its *R*- and *S*-isomers and transformation products 2-COOH-S-2200, 5-COOH-S-2200, S-2200-OR and S-2200-ORC was undertaken for freshwater and marine organisms based on available toxicity data. A summary of aquatic toxicity data is presented in Appendix I, Table 16. The accompanying risk assessment is presented in Appendix I, Tables 17-20.

Freshwater invertebrates: Mandestrobin was moderately toxic and S-2200 *R*-isomer was highly toxic to daphnids on an acute basis. Acute exposure to S-2200 *S*-isomer and other transformation products did not affect daphnids. In general, the *S*-isomer was shown to be of lower toxicity to aquatic organisms than mandestrobin, while the *R*-isomer was shown to be of comparable toxicity. Chronic exposure to mandestrobin reduced parental survival, reproduction and growth in daphnids, reduced the survival in chironomids, and reduced body length in amphipods.

The screening level risk quotients for freshwater invertebrates from acute and chronic exposure to mandestrobin do not exceed the level of concern. The risk quotients for daphnids from acute exposure to S-2200 *R* and *S*-isomers and the transformation products, 2-COOH-S-2200, 5-COOH-S-2200, S-2200-OR and S-2200-ORC do not exceed the level of concern at the screening level.

Freshwater fish: Mandestrobin was demonstrated to be of moderate toxicity to bluegill sunfish and sheepshead minnow, but showed high toxicity to rainbow trout and fathead minnow. The S-2200 *R*-isomer was highly toxic to rainbow trout on an acute basis. Acute exposure to S-2200 *S*-isomer and other transformation products did not affect rainbow trout.

The risk quotients for rainbow trout resulting from acute exposure to S-2200 *S*-isomer and transformation products 2-COOH-S-2200, 5-COOH-S-2200, S-2200-OR did not exceed the level of concern at the screening level. The screening level risk quotients for freshwater fish resulting from acute and chronic exposure to mandestrobin, and from acute exposure to S-2200 *R*-isomer and S-2200-ORC exceeded the level of concern. The risk to freshwater fish from acute and chronic exposure to mandestrobin, S-2200 *R*-isomer and S-2200-ORC was further assessed (see Section 4.2.3).

Amphibians: To assess the risk to amphibians, fish toxicity endpoints are used as surrogate data, when amphibian data are not available, to represent aquatic life-stages of amphibians. The difference between fish and amphibian risk assessments is related to the water depth used for the estimated environmental concentrations (water depth of 15 cm for amphibians). The screening level risk quotients for acute and chronic exposures of amphibians to mandestrobin exceeded the level of concern. The risk to amphibians was thus further assessed (see Section 4.2.3).

Freshwater algae and vascular plants: Mandestrobin showed adverse effects on cell density, growth rate and yield in green algae, blue-green algae and diatoms. S-2200 *R*-isomer and 2-COOH-S-2200 reduced the growth rate and biomass in test with green algae. S-2200 *S*-isomer and other transformation products had no adverse affects on green algae. Mandestrobin reduced the frond number and biomass of duckweed.

The risk quotient for freshwater algae resulting from acute exposure to mandestrobin and the S-2200 *R*-isomer exceeded the level of concern at the screening level. The risk quotients for freshwater green alga from exposure to S-2200 *S*-isomer and the transformation products 2-COOH-S-2200, 5-COOH-S-2200, S-2200-OR and S-2200-ORC did not exceed the level of concern at the screening level. The risk quotient for duckweed from exposure to mandestrobin did not exceed the level of concern. The risk to freshwater algae from acute exposure to mandestrobin and S-2200 *R*-isomer was thus further assessed (see Section 4.2.3).

Marine/estuarine invertebrates: Mandestrobin was acutely toxic to the mysid shrimp and moderately toxic to the eastern oyster. Exposure to mandestrobin for 36 days affected the first generation growth and survival of the mysid shrimp. Mandestrobin had no chronic effects on the sediment dwelling estuarine amphipod.

The risk quotient for marine invertebrates (mysid shrimp) resulting from acute and chronic exposure to mandestrobin exceeded the level of concern at screening level. The risk to marine invertebrates was, thus, further assessed (see Section 4.2.3).

Marine/estuarine fish: Acute exposure to mandestrobin had no adverse effects to the sheepshead minnow at the highest test concentration. Exposure to mandestrobin during early life stages of sheepshead minnows resulted in significant reductions in total length, wet weight and dry weight.

The screening level risk quotient for marine fish resulting from acute exposure to mandestrobin exceeded the level of concern. The risk quotients for marine fish resulting from chronic exposure to mandestrobin did not exceed the level of concern at the screening level. The acute risk to marine fish was, thus, further assessed (see Section 4.2.3).

Marine/estuarine algae: Mandestrobin exhibited acute adverse effects on cell density, growth rate and yield of the marine diatom.

The risk quotient for marine diatom resulting from acute exposure to mandestrobin did not exceed the level of concern at the screening level.

4.2.3 Further characterization of risk to aquatic organisms

4.2.3.1 Assessment of potential risk from spray drift

To further characterize the risk to fish, amphibians, crustaceans and algae, refined EECs for aerial application were calculated using a maximum drift deposition percent at one metre downwind from the point of application. The maximum percent drift deposition for aerial and ground application and an ASAE 'coarse' droplet size (as specified on the product labels) is 17% and 3% of the application rate, respectively. The EECs were calculated for water bodies 15-cm and 80-cm deep.

The refined risk quotients for fish, crustacean (mysid shrimp) and marine diatom indicate that the level of concern from exposure to mandestrobin, S-2200 *R*-isomer and S-2200-ORC due to spray drift is not exceeded. The refined risk quotients for amphibians and freshwater algae indicate that the level of concern from mandestrobin exposure due to spray drift is exceeded for aerial application, but not exceeded for ground application. It was determined that drift mitigation measures in the form of spray buffer zones were necessary.

4.2.3.2 Assessment of potential risk from run-off

The risk from exposure to run-off into a body of water directly adjacent to the application field was determined using the run-off 90th percentile of the EECs predicted by PRZM-EXAMS. The risk quotients for exposure to mandestrobin, S-2200 *R*-isomer and the transformation product S-2200-ORC were calculated using toxicity endpoints and EECs representing the 90th percentile of 96-hour concentration.

The risk quotients for aquatic organisms resulting from exposure to S-2200 *R*-isomer and S-2200-ORC through runoff do not exceed the level of concern. However, the risk quotients from exposure of mandestrobin to amphibians, freshwater algae and marine crustaceans through runoff marginally exceeded the level of concern ($RQ \le 2.5$). Standard precautionary label statements are required on the product label to inform users of best practices to minimize potential run-off from treated fields.

5.0 Value

5.1 Consideration of Benefits

Registration of mandestrobin, and the end-use products that contain this new active ingredient, will provide Canadian growers and turf managers with additional products on the market with which to address major disease problems.

A number of fungicides, including some from the same mode of action group, are registered on the subject crops to control or suppress the plant diseases indicated on the mandestrobin product labels. Refer to Appendix I, Table 22 for further information on the currently available alternatives grouped according to their modes of action.

With broad-ranging efficacy and two methods of application, mandestrobin represents a valuable addition to an effective integrated pest management approach for various cropping systems. As multiple alternative fungicides from different mode of action groups are currently registered for most diseases appearing on the foliar product and seed treatment labels, appropriate resistance management strategies can be implemented. The use patterns being registered for mandestrobin will allow application of this active ingredient in combination with good agricultural practices, including cultural methods that help to lower disease pressure.

5.2 Effectiveness Against Pests

S-2200 4 SC Fungicide; S-2200 4 SC AG Fungicide; S-2200 4 VPP Fungicide (foliar products)

Efficacy data from a total of 33 small scale trials conducted between 2006 and 2012 were reviewed to support the value of nine different disease claims on the foliar-applied mandestrobin products. A complete list of supported uses with additional details is provided in Appendix I, Table 23. Most trials were located either in Canada or northern United States. Depending on the crop disease combinations, the levels of efficacy demonstrated across the different trials were consistent with performance standards expected from claims of either control or suppression. In addition, low water spray volumes were used for ground applications in six of the trials provided with the intention of simulating aerial applications. No reduction in efficacy was observed from these treatments, thereby supporting directions for aerial applications on canola.

S-2200 3.2 FS Fungicide (seed treatment product)

Twenty small scale field and greenhouse efficacy trials, conducted in Canada between 2011 and 2013, were reviewed in support of the claims on the mandestrobin-containing seed treatment product S-2200 3.2 FS Fungicide. Evidence of efficacy from four bioassays was also considered in support of the product's label. Mandestrobin was demonstrated to be effective in reducing the severity and incidence of seed decay and seedling diseases caused by various seed- and soilborne pathogens, including *Fusarium* and *Rhizoctonia*. In the majority of tested crop/pathogen combinations, mandestrobin was effective in controlling the diseases in question. However, when assessing seed decay caused by the pathogen *Phomopsis* in soybean, mandestrobin seed treatments resulted in disease suppression rather than control.

5.3 Non-Safety Adverse Effects

Phytotoxicity resulting from applications of mandestrobin-containing products was not reported in any of the available trial data. There is no indication that non-safety adverse effects would result from S-2200 4 SC Fungicide, S-2200 4 SC AG Fungicide, S-2200 4 SC VPP Fungicide or S-2200 3.2 FS Fungicide when applied to crops in accordance with label directions and restrictions.

5.4 Supported Uses

A complete list of supported uses is provided in Appendix I, Table 23.

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: in other words, 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, mandestrobin and its transformation products were assessed in accordance with the PMRA Regulatory Directive DIR99-03⁵ and evaluated against the Track 1 criteria. The PMRA has reached the following conclusions:

• Mandestrobin does not meet all Track 1 criteria, and is not considered a Track 1 substance. See Appendix I, Table 21 for comparison with Track 1 criteria.

⁵ DIR99-03, The Pest Management Regulatory Agency's Strategy for Implementing the Toxic Substances Management Policy

• Mandestrobin does not form any transformation products that meet all Track 1 criteria. Major transformation products S-2200-OR, S-2200-ORC and S-2200-PR are formed only by aqueous phototransformation. They are not expected to be formed in important quantities in the environment as aqueous phototransformation is restricted to the upper layer of clear water bodies.

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*.⁶ The list is used as described in the PMRA Notice of Intent NOI2005-01⁷ and is based on existing policies and regulations including: DIR99-03; and DIR2006-02,⁸ 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 conclusion:

• Technical grade mandestrobin (S-2200 Fungicide Technical) and the end-use products S-2200 4 SC Fungicide, S-2200 3.2 FS Fungicide, S-2200 4 SC Ag Fungicide and S-2200 4 SC VPP Fungicide do 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.

7.0 Summary

7.1 Human Health and Safety

The toxicology database submitted for mandestrobin is adequate to define the majority of toxic effects that may result from exposure. The main targets were the liver (mice, rats, dogs), thyroid gland (rats) and kidneys (mice, rats) in several short- and long-term studies. There was evidence of increased susceptibility of the young in the toxicity studies submitted, but the level of concern was low. There was equivocal evidence of oncogenicity in rats following longer-term dosing. The risk assessment protects against the toxic effects noted by ensuring that the level of human exposure is well below the lowest dose at which these effects occurred in animal tests.

⁶ 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.

⁷ NOI2005-01, *List of Pest Control Product Formulants and Contaminants of Health or Environmental Concern* under the *New Pest Control Products Act.*

⁸ DIR2006-02, Formulants Policy and Implementation Guidance Document.

Mixers, loaders, and applicators handling S-2200 4 SC Fungicide, S-2200 4 SC Ag Fungicide, S-2200 4 SC VPP Fungicide, and S-2200 3.2 FS Fungicide, and workers re-entering treated areas of rapeseed, climbing vine fruit, low-growing berries, turfgrass and sod, and planting treated corn, legumes, and rapeseed seeds are not expected to be exposed to levels of mandestrobin that will result in an unacceptable risk when these products are used according to label directions. The personal protective equipment and restricted entry interval on the product labels are adequate to protect workers. The S-2200 3.2 FS Fungicide label also includes the requirement to treat seeds in commercial facilities using closed mix, load, and transfer seed treating equipment.

Residential exposures to individuals contacting treated turf, including golfing, are not expected to result in risks of concern when S-2200 4 SC VPP Fungicide is used according to label directions. Exposure to the public entering pick-your-own strawberry and lowbush blueberry farms, following treatment with S-2200 4 SC Ag Fungicide, is also not expected to result in risks of concern.

The nature of the residues in plants is adequately understood. The residue definition for enforcement and risk assessment is mandestrobin in plant matrices. The proposed use of mandestrobin on rapeseed (canola), corn, grapes, strawberries, and soybeans does not constitute a risk of concern from chronic 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 recommend MRLs. The PMRA recommends that the following MRLs be specified for residues of mandestrobin.

Commodity	Recommended MRL (ppm)
Raisins	7.0
Small fruit vine climbing (Crop Subgroup 13-07F, except fuzzy kiwifruit)	5.0
Low growing berry (Crop Subgroup 13-07G, except cranberry)	3.0
Rapeseed (Crop Subgroup 20A)	0.5
Legume vegetables (Crop Group 6, except cowpea and field pea), corn (field, popcorn, sweet)	0.02

7.2 Environmental Risk

To mitigate risks to non-target organisms, spray buffer zones to protect sensitive aquatic and terrestrial habitats from spray drift and label statements to inform users of potential risks to the environment are required. With these measures in place, risks to the environment from the use of mandestrobin are considered to be acceptable.

7.3 Value

The value information submitted was sufficient to support the registration of 16 fungal disease claims for four new end-use products containing the new fungicidal active ingredient mandestrobin. Intended either for foliar applications or seed treatments, the supported uses cover a broad range of diseases in economically important field and horticultural crops including turfgrass. Registration of these products will provide Canadian growers and producers additional disease management products with a new active ingredient from a mode of action group with a well-established efficacy profile.

8.0 Proposed Regulatory Decision

Health Canada's PMRA, under the authority of the *Pest Control Products Act* and Regulations, is proposing full registration for the sale and use of Mandestrobin Technical (previously known as S-2200 Fungicide Technical) and the associated end-use products: S-2200 4 SC Fungicide, Intuity Fungicide (previously known as S-2200 4 SC Ag Fungicide), Pinpoint Fungicide (previously known as S-2200 4 SC VPP Fungicide), and S-2200 3.2 FS Fungicide, containing the technical grade active ingredient mandestrobin, for the management of various fungal diseases in canola and other oilseed crops, corn, grape, legume vegetables, strawberry and other low growing berries, as well as turfgrass.

An evaluation of available scientific information found that, under the approved conditions of use, the product has value and does not present an unacceptable risk to human health or the environment.

List of Abbreviations

°C	degrees Celsius
\mathbf{C}	female
+ 1	male
♀ ? ↓	
	increasing
	decreasing
3	emittance
λ	wavelength
μg	microgram(s)
μM	micromolar(s)
1/n	exponent for the Freundlich isotherm
a.i.	active ingredient
abs	absolute
AD	administered dose
ADI	acceptable daily intake
ADME	absorption, distribution, metabolism and excretion
AHETF	Agricultural Handlers Exposure Task Force
ALP	alkaline phosphatase
ALT	alanine aminotransferase
APTT	activated partial thromboplastin time
appl.	application
AR	applied radioactivity
ARfD	acute reference dose
ARTF	Agricultural Re-entry Task Force
ASAE	American Society of Agricultural Engineers
atm	atmosphere
ATPD	area treated per day
BAF	bioaccumulation factor
BBCH	Biologishe Bundesanstalt, Bundessortenamt and Chemical industry
BCF	bioconcentration factor
bw	body weight
bwg	body weight gain
Bz	benzyl
C _{max}	maximum plasma concentration
CAF	composite assessment factor
CAS	Chemical Abstracts Service
CDC	Center for Disease Control
CDN	Canadian
CEPA	Canadian Environmental Protection Act
CG	Crop Group
CHL	chinese hamster lung cell line
cm	centimetre(s)
cm ²	centimetre(s) squared
CO_2	carbon dioxide
conj.	conjugated
CSG	Crop Sub-Group
	1 ··· - · · · · · · · · · · · · · · · ·

СҮР	cytochrome P450
d	day(s)
DALA	days after last application
DALA DAT	days after treatment
DEEM-FCID	•
	Dietary Exposure Evaluation Model – Food Commodity Intake Database
DFOP	double first-order in parallel
DFR	dislodgeable foliar residue Directive
DIR	
DNA	deoxyribonucleic acid
DT_{50}	dissipation time 50% (the dose required to observe a 50% decline in
ЪШ	concentration)
DT_{90}	dissipation time 90% (the dose required to observe a 90% decline in
	concentration)
dw	dry weight
E_bC_{50}	EC ₅₀ in terms of algal biomass
EC_{25}	effective concentration on 25% of the population
EC_{50}	effective concentration on 50% of the population
E. coli	Escherichia coli
EDE	estimated daily exposure
EEC	estimated environmental concentration
ER_{25}	effective rate on 25% of the population
ER_{50}	effective rate on 50% of the population
F_1	first generation
F_2	second generation
fc	food consumption
FDA	Food and Drugs Act
FIR	food ingestion rate
FRAC	Fungicide Resistance Action Committee
FS	flowable suspension
g	gram(s)
GAP	Good Agricultural Practice
GC-MS	gas chromatography with mass spectrometry
GI	gastrointestinal
GUS	groundwater ubiquity score
h	hour(s)
H295R	human adrenocarcinoma cell line
ha	hectare(s)
HAFT	highest average field trial value
hAR	human androgen receptor
HCl	hydrochloric acid
HeLa	Henrietta Lacks
hERa	human oestrogen receptor alpha
HDPE	High-density polyethylene
HPLC	high performance liquid chromatography
HPLC-MS/MS	high performance liquid chromatography with tandem mass spectrometry
HPRT	hypoxanthine phosphorybosyl transferase
ID	identification
	nontricution

ILV	independent laboratory validation
IORE	indeterminate order rate equation
IUPAC	International Union of Pure and Applied Chemistry
K+CWHR	kernels and cobs with husk removed
kg V	kilogram(s)
K _d	soil-water partition coefficient
K _F	Freundlich adsorption coefficient
K _{FOC}	Freundlich adsorption coefficient normalized to organic carbon
K _{oc}	organic-carbon partition coefficient
$K_{ m ow}$	<i>n</i> -octanol-water partition coefficient
	litre(s)
LAFT	lowest average field trial value
LC_{50}	lethal concentration 50%
LC-MS/MS	liquid chromatography with tandem mass spectrometry
LD_{50}	lethal dose 50%
LLNA	local lymph node assay
LOAEL	lowest observed adverse effect level
LOC	level of concern
LOQ	limit of quantitation
LR ₅₀	lethal rate 50%
LSC	liquid scintillation counting
m_{2}^{2}	square metre(s)
m ³	cubed metre(s)
Max.	maximum
mg	milligram(s)
Min.	minimum
mL	millilitre(s)
M/L/A	mixer/loader/applicator
MAS	maximum average score
MIS	maximum irritation score
MOA	mode of action
MOE	margin of exposure
mol	mole
MQL	minimum quantification limit
MRL	maximum residue limit
m/z	mass-to-charge ratio of an ion
n	number of field trials
N/A or n/a	not applicable
NAFTA	North American Free Trade Agreement
NaOH	sodium hydroxide
NC	not classified
NCHS	National Center for Health Statistics
NHANES/ WWEIA	National Health and Nutritional Examination Survey, What We Eat in
	America
nm	nanometre(s)
NOAEL	no observed adverse effect level
NOEC	no observed effect concentration

NOF	
NOEL	no observed effect level
NZW	New Zealand white
OC	organic carbon content
ORETF	Outdoor Residential Exposure Task Force
Pa	Pascal
PBI	plantback interval
PES	post-extraction solids
Ph	phenoxy
PHED	Pesticide Handlers Exposure Database
PHI	preharvest interval
p <i>K</i> a	dissociation constant
PLT	platelet
PMRA	Pest Management Regulatory Agency
PND	postnatal day
ppb	parts per billion
PPE	
	personal protective equipment
ppm	parts per million
PT	prothrombin time
PYO	pick your own
RAC	raw agricultural commodity
REI	restricted entry interval
rel	relative
RQ	risk quotient
R/S	enantiomeric ratio
SC	soluble concentrate
SD	standard deviation
SEEDTROPEX	Seed Treatment Operator Exposure Task Force
SFO	single first order
SOP	Standard Operating Procedure
SPN	Science Policy Notice
S. typhimurium	Salmonella typhimurium
t _{1/2}	total elimination half-life
T3	tri-iodothyronine
T4	thyroxine
TLC	thin layer chromatography
	time to reach maximum plasma concentration
t _{max} TG	
	technical grade
t _{R IORE}	representative half-life dissipation in soil calculated with the IORE model
TRR	total radioactive residue
TRT.	treatment
TSH	thyroid-stimulating hormone
TSMP	Toxic Substances Management Policy
TTR	turf transferable residue
UDP	uridine 5'-diphospho-
UF	uncertainty factor
UGT	UDP-glucuronosyltransfase
UR	unextracted residues

US USEPA UV	United States United States Environmental Protection Agency ultraviolet
VS.	verses
v/v	volume per volume dilution
v/v/v	volume per volume per volume dilution
W	week(s)
wt(s)	weight(s)

Appendix I Tables and Figures

Matrix	Method ID	Analyte	Method Type	LOQ	Reference
Soil	CLE 8213772-	S-2200 R isomer	LC-MS/MS m/z 314 \rightarrow 192	0.0025 mg/kg	2377924
	01V	S-2200 S isomer	<i>m</i> / <i>z</i> 314→192		
		De-Xy-S-2200	<i>m/z</i> 210→192	0.005 mg/kg	
		2-COOH-S-2200	<i>m</i> / <i>z</i> 344→192		
		5-COOH-S-2200	<i>m</i> / <i>z</i> 344→160		
Soil	RM-48S-3	S-2200	HPLC-MS/MS m/z 314 \rightarrow 192	0.02 mg/kg	2377958 and 2377849
		DX-CA-S-2200	<i>m/z</i> 224→146		
		2-COOH-S-2200	<i>m</i> / <i>z</i> 344→192		
		5-COOH-S-2200	<i>m</i> / <i>z</i> 344→192		
		2-CONH2-S-2200	<i>m</i> / <i>z</i> 343→192		
	5-CONH2	5-CONH2-S-2200	<i>m</i> / <i>z</i> 343→192		
Water	Not stated	S-2200	HPLC-MS/MS m/z 314 \rightarrow 192	0.10 μg/L	2377854 and 2377857

Table 1a Residue Analysis for Environmental Media

Table 1bResidue Analysis for Crops

Matrix	Method ID	Analyte	Method Type	LOQ (ppm)	Reference
Rapeseed Rotational Crops: Wheat (forage, grain, straw), Lettuce, Beet (roots, leaves)	RM-48C-2A (precedent RM-48C-2) (Enforcement Method- Plant Commodities)	S-2200	LC-MS/MS	0.02	2377885
Rapeseed	RM-48C-2A (ILV of enforcement method – plant)	S-2200	LC-MS/MS	0.02	2377899
	was monitored for S-2200 od for the analysis of S-220		wever DFG-S1	9 method can be used	as a
Rapeseed (seeds)	DFG-S19 (Multiresidue method)	S-2200	LC-MS/MS	0.005: <i>R</i> and <i>S</i> -isomer	2377889
Barley (grain, straw), lettuce	DFG-S19 (Validation of multiresidue method)	S-2200	LC-MS/MS	0.005: <i>R</i> and <i>S</i> -isomer	2377892
Radiovalidation (plant commodities)	Similar solvents were used for the metabolism studies (acetone/water; 80:20) and the enforcement method (acetone/water; 70:30). In general, extraction efficiencies were similar for both extraction methods (Tissumizer or shaking) with both solvent systems in rapessed forage (80.3-90.9% of the TRR). Seed samples were also extracted with a Tissumizer and a solvent solution of acetone:water, 70:30, v/v, which yielded an average extraction efficiency of 108.1% of the TRR.			2377887	
Rapeseed, Corn (Forage and Stover)	RM-48C-1 (Data-Gathering Method – Plant	S-2200	GC-MS	0.02	2377926; 2377153; 2377152

Matrix	Method ID	Analyte	Method Type	LOQ (ppm)	Reference
Corn (Grain and K+CWHR), Soybean	Commodities)			0.01	
Strawberry, Grape	RM-48G (Data-Gathering Method – Plant Commodities)	S-2200	LC-MS/MS	0.02	2377917; 2377911; 2377920; 2377913
Poultry muscle	RM-48M-1 (ILV of Proposed Enforcement Method – Livestock Commodities)	S-2200	LC-MS/MS	0.02	2377859

If livestock feed items increase the dietary burden in cattle and poultry, a validated enforcement method using cattle fat, meat and meat byproducts, milk, and eggs will be necessary, including ILV and radiovalidation.

For further use expansions involving cereal grains (wheat), the applicant is requested to also measure for the metabolite De-Xy-S-2200 in the crop field trials (along with mandestrobin) as it was a major metabolite found in the metabolism studies for wheat grain and straw. The residue definition of mandestrobin may be revised as a result.

Table 2 Chemical Names of Isomers and Metabolites of Mandestrobin

Code/Trivial name	Chemical name
S-2167 (R-isomer of	(R)-2-methoxy-N-methyl-2-[α-(2,5-xylyloxy)-o-tolyl]acetamide
mandestrobin)	
S-2354 (S-isomer of	(S)-2-methoxy-N-methyl-2-[α-(2,5-xylyloxy)-o-tolyl] acetamide
mandestrobin)	
2-CH ₂ OH-S-2200	(2RS)-2-[2-(2-hydroxymethyl-5-methylphenoxymethyl)phenyl]-2-methoxy-N-
	methylacetamide
2-COOH-S-2200	2-({2-[(1RS)-1-methoxy-2-(methylamino)-2-oxoethyl]benzyl}oxy)-4-
	methylbenzoic acid
4-OH-S-2200	(2RS)-2-[2-(4-hydroxy-2,5-dimethylphenoxymethyl)phenyl)-2-methoxy-N-
	methylacetamide
5-COOH-S-2200	3-({2-[(1RS)-1-methoxy-2-(methylamino)-2-oxoethyl]benzyl}oxy)-4-
	methylbenzoic acid
De-XY-S-2200	(2RS)-2-(2-hydroxymethylphenyl)-2-methoxy-N-methylacetamid

Table 3Toxicity Profile of S-2200 3.2 FS Fungicide

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

Study Type/Animal/PMRA #	Study Results
Acute Oral Toxicity	$LD_{50^{\circ}} = 3129 \text{ mg/kg bw}$
	Low Toxicity
Sprague-Dawley Rat	
PMRA # 2378126	
Acute Dermal Toxicity	$LD_{50\Im/2} > 5000 \text{ mg/kg bw}$
	Low Toxicity
Sprague-Dawley Rat	
PMRA # 2378127	

Study Type/Animal/PMRA #	Study Results
Acute Inhalation	$LC_{50\Im/2} > 2.04 \text{ mg/L}$
(nose-only)	Low Toxicity
Sprague-Dawley Rat	
PMRA # 2378128	
Eye Irritation	MIS= 8.0/110
	MAS (24, 48, 72 hour)= 0.22/110
NZW rabbit	Minimally irritating
PMRA # 2378129	
Dermal Irritation	MIS= 1/8
	MAS (24, 48, 72 hour)= 0/8
NZW Rabbit	Non-irritating
PMRA # 2378130	
Local Lymph Node Assay	Non-sensitizer
(LLNA)	
CBA/J Mouse	
PMRA # 2378131	

Table 4Toxicity Profile of S-2200 4 SC Fungicide, S-2200 4 SC AG Fungicide and S-
2200 4 SC VPP Fungicide

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

Study Results
$LD_{50\circ} = 3129 \text{ mg/kg bw}$
Low Toxicity
$LD_{50,0/2} > 5000 \text{ mg/kg bw}$
Low Toxicity
$LC_{50\Im/\wp} > 2.04 \text{ mg/L}$
Low Toxicity

Study Type/Animal/PMRA #	Study Results
Eye Irritation	MIS= 3.33/110
	MAS (24, 48, 72 hour)= 0.22/110
NZW rabbit	Minimally irritating
PMRA # 2377866	
Dermal Irritation	MIS= 0/8
	MAS (24, 48, 72 hour)= 0/8
NZW Rabbit	Non-irritating
PMRA # 2377868	
Local Lymph Node Assay	Non-sensitizer
(LLNA)	
CBA/J Mouse	
PMRA # 2377870	

Table 5 Toxicity Profile of Technical Mandestrobin

(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) Effects seen above the LOAEL(s) have not been reported in this table for most studies for reasons of brevity.

Study Type/Animal/PMRA #	Study Results				
ADME	Mandestrobin (benzyl- ¹⁴ C-S-2200TG and phenoxy- ¹⁴ C-S-2200TG, racemic mixtures), R-				
(single administration [S-	isomer (benzyl- ¹⁴ C-S-2167), S-isomer (benzyl- ¹⁴ C-S-2354). All forms of mandestrobin were				
2200TG, S-2167, S-2354],	rapidly absorbed (\geq 95% administered single dose (AD); 73-81% by 78 hours). Systemic				
multiple administration [S-	exposure was proportionally higher at the low dose than at the high dose, indicating saturation				
2200TG])	of oral absorption at high dose. Peak plasma levels (C _{max}) were reached by 1.2-2.6 hours (t _{max})				
	after an oral dose of 5 mg/kg, and 7.0-9.1 hours after a dose of 1000 mg/kg.				
Wistar Rat	The labeled compound was widely distributed throughout tissues, but was detected primarily in				
	the GI tract, liver and kidney. Pancreas, uterus and ovaries also had higher levels than most				
	other tissues. Isomers had similar tissue distribution except the S-isomer was more prevalent in				
PMRA #2377990, 2377989,	GI tract than the R-isomer. There was no sex difference in distribution into tissues, except				
2377987	during the multiple dosing period, females kept a larger portion of the dose in the cecum/large				
	intestine/intestinal contents, and reached peak levels in tissues outside the GI tract later than				
	males.				
	There was no evidence of long-term accumulation of S-2200TG or its metabolites in tissues. Clearance of a single dose of S-2200TG from plasma was almost complete by 120 hours post-				
	dose ($t_{1/2} = 18-23$ hours at the low dose; $t_{1/2} = 25-29$ hours at the high dose). Clearance of the S-				
	isomer was less rapid than the R-isomer. In the repeated dose study, 85-89% of the AD was				
	excreted by 14 days after last dose. Bile cannulated rats excreted S-2200TG faster than non-				
	cannulated rats (97-99% vs. 49-51% of a single dose by 24 hours), suggesting enterohepatic				
	recirculation. Excretion was consistent with all S-2200TG treatments. Fecal excretion (60-75%)				
	of AD) via bile was the primary excretion route, and renal excretion (15-21% of AD) was also				
	important. Excretion in expired air was negligible.				
	S-2200TG is almost completely metabolized in rats via (1) oxidation followed by				
	glucuronidation, (2) demethylation followed by oxidation, or (3) oxidation followed by				
	demethylation. The molecule core remains essentially intact, with no cleavage of benzyl and				
	phenoxy rings. The major metabolites excreted are: 5-CA-S-2200-NHM and 4-OH-S-2200 in				
	feces; 4-OH-S-2200-GlucA in bile; and 5-CA-S-2200-NHM in urine. The same metabolites				
	were identified for both isomers however at different proportions. In order of magnitude, R-				
	isomer produced mostly 5-CA-S-2200-NHM, 5-CA-MCBX-NDM, 5-CA-2-HM-S-2200-NHM				

	and 5 CA S 2200 NDM, while S isomer produced models 4 OH S 2200 and 5 COOH S 2200
	and 5-CA-S-2200-NDM; while S-isomer produced mostly 4-OH-S-2200 and 5-COOH-S-2200. In the multiple dose study, fecal metabolites found at slightly higher levels in male than females were 5-CA-S-2200-NHM, 5-COOH-S-2200, 5-CA-2-HM-S-2200-NHM, 5-CA-2-HM- MCBX and 5-CA-MCBX-NDM.
	Results from cannulated rats suggest that 4-OH-S-2200 and its A glucuronide (GlucA) undergo enterohepatic circulation.
Acute Oral Toxicity	$LD_{50^{\circ}} > 2000 \text{ mg/kg bw}$ Low Toxicity
Wistar Rat	
PMRA # 2377929	
Acute Dermal Toxicity	LD ₅₀₃₉ > 2000 mg/kg bw
Wistar Rat	Low Toxicity
PMRA # 2377936	
Acute Inhalation (nose-only)	$LC_{50\Im \circ} > 4.96 \text{ mg/L}$ Low Toxicity
Wistar Rat	
PMRA # 2377937	
Eye Irritation	MIS = 10.3/110 $MAS (24, 48, 72 hour) = 4.0/110$
NZW rabbit	MAS (24, 48, 72 hour)= 4.0/110 Minimally irritating
PMRA # 2377939	
Dermal Irritation	MIS= 0/8
NZW Rabbit	MAS (24, 48, 72 hour)= 0/8 Non-irritating
PMRA # 2377940	
Sensitization Study	Non-sensitizer
(Maximization Test)	
Guinea Pig	
PMRA # 2377941	
28-Day dermal toxicity	NOAEL = 1000 mg/kg bw/day LOAEL was not determined due to absence of effects up to the limit dose.
Wistar Rat	Ĩ
PMRA # 2377957	
90-Day oral toxicity (diet)	NOAEL= 807/529 mg/kg bw/day in $3/2$ LOAEL was not determined in 3
CD-1 Mouse	LOAEL \Im = 1111 mg/kg bw/day; effects included decreased bw and lower bwg
PMRA # 2377946	
90-Day oral toxicity (diet)	NOAEL= 283/320 mg/kg in $3/2$
Wistar Rat	LOAEL= 743/789 mg/kg bw/day in \Im/\Im ; effects included \uparrow incidence hepatocyte hypertrophy and severity, \uparrow total cholesterol, \uparrow incidence of thyroid follicular cell hypertrophy; \uparrow liver wts, \uparrow incidence large liver, \uparrow incidence liver agonal congestion/ hemorrhage, \uparrow severity thyroid
PMRA # 2377948	follicular hypertrophy (\Im); \uparrow rel liver wt (\Im)
90-Day oral toxicity (diet)	NOAEL= 91/103 mg/kg bw/day in \Im/\Im
Beagle Dog	LOAEL= 268/304 mg/kg bw/day in \Im/\Im ; effects included \uparrow incidence liver centrilobular degeneration and severity, \uparrow incidence dark liver; \uparrow incidence large liver (\Im); \uparrow ALP (\Im)

PMRA # 2377954					
1-Year oral toxicity (diet)	NOAEL= 92 mg/kg bw/day				
	LOAEL= 181/226 mg/kg bw/day in ∂/φ ; effects included \downarrow bw, \uparrow liver wt, \uparrow incidence liver				
Beagle Dog	centrilobular degeneration, agonal congestion/hemorrhage, hepatocyte hypertrophy and				
5 6	severity, hepatocyte pigment, \uparrow ALP, \uparrow ALT, \uparrow PLT; \downarrow bwg (\checkmark); \downarrow PT and APTT, thin				
PMRA # 2377952	appearance and reduced muscle tone $(1/4)$ (\bigcirc)				
78- week Oncogenicity	NOAEL = 239/994 mg/kg bw/day in \Im/\Im				
(diet)	LOAEL not determined in \mathcal{L}				
	LOAEL = 824 mg/kg bw/day in 3 ; effects included \uparrow incidence of corticomedullary				
CD-1 Mouse	mineralization (δ)				
PMRA #2377960	No evidence of oncogenicity				
2-Year combined oral chronic	NOAEL = $130/27$ mg/kg bw/day in $\sqrt[3]{9}$				
toxicity /carcinogenicity	LOAEL = 449/135 mg/kg bw/day in \Im/\Im ; effects included \uparrow liver wt, liver hypertrophy,				
(diet)	thyroid follicular cell hypertrophy (δ); \downarrow bw, \downarrow bwg, rel liver wt, hepatocyte hypertrophy, \uparrow				
	incidence of corticomedullary mineralization (\mathbb{Q})				
Wistar rat	(+)				
	Equivocal evidence of oncogenicity based on a treatment-related increase in ovary sex cord-				
PMRA # 2377961	stromal benign tumors in female rats $(2, 0, 1, 4, 6)$.				
2-generation reproductive	NOAEL (parental) = 1000 ppm (56/63 mg/kg bw/day in $3^{1/2}$)				
toxicity study	LOAEL (parental) = 3000 ppm (166/195 mg/kg bw/day in ∂/φ); effects included \uparrow bile duct				
	brown pigmentation and periductular inflammatory cell infiltration (with liver weight and				
BrlHan:WIST@Jcl	hypertrophy at the same dose)				
(GALAS) Rats					
	NOAEL (offspring) = 3000 ppm (195 mg/kg bw/day)				
PMRA # 2377964	LOAEL (offspring) = 10000 ppm (628 mg/kg bw/day); effects included \downarrow bw F1/F2 from PND				
	7				
	NOAEL (reproductive) = 10000 ppm (629 mg/kg bw/day)				
	LOAEL (reproductive) \geq 10000 ppm (629 mg/kg bw/day)				
	No evidence of sensitivity of the young				
Developmental Toxicity Study in	Maternal NOAEL = 1000 mg/kg bw/day				
Rats	Maternal LOAEL > 1000 mg/kg bw/day				
Culture) Data	Developmental NO AEL = 200 mg/kg hu/day				
Crl:WI(Han) Rats	Developmental NOAEL = 300 mg/kg bw/day Developmental LOAEL = 1000 mg/kg bw/day; effects included ↑ increased incidences of				
PMRA # 2377967	litters containing fetuses with distended ureter or delayed skull ossification				
F WIRA # 2377907	inters containing refuses with distended dreter of delayed skull ossification				
	Evidence of sensitivity of the young				
Developmental Toxicity Study in	Maternal NOAEL = $1000 \text{ mg/kg bw/day}$				
Rabbits	Maternal LOAEL > 1000 mg/kg bw/day				
Hsd:IfNZW Rabbits	Developmental NOAEL = $1000 \text{ mg/kg bw/day}$				
	Developmental LOAEL > 1000 mg/kg bw/day				
PMRA # 2377970					
	No evidence of sensitivity of the young				
Bacterial Reverse Mutation	Negative in S. typhimurium strains (TA 98, TA100, TA1535, TA1537 and TA1538) and E. coli				
Assay (Ames test)	WP2uvrA in the presence and absence of metabolic activation.				
• • •					
PMRA # 2377971					
In vitro Mammalian Cell Assay	Negative in V79-HPRT cells				
(forward gene mutation)					

In vitro Mammalian	Negative in Chinese hamster lung cells (CHL/IU)		
Clastogenicity Assay			
(chromosomal aberration)			
PMRA#2377984			
Micronucleus Assay	Negative in BDF ₁ [SPF] mice		
	No mortality, clinical signs of toxicity and no differences in body weight compared to controls.		
PMRA #2377985			
Acute neurotoxicity (gavage)	NOAEL = 1000 mg/kg bw in $3/2$		
	LOAEL = 2000 mg/kg bw in \Im/\Im ; effects included \downarrow overall locomotor activity (total and/or		
Wistar Rat	ambulatory counts) on study day 0 (0-30 min)		
PMRA # 2377992			
90-Day neurotoxicity study	NOAEL = 338/1223 mg/kg bw/day in ∂/Q		
	LOAEL not determined in \bigcirc		
Wistar rat	LOAEL = 1024 mg/kg bw/day in 3 ; effects included \downarrow bw and lower bwg, \downarrow fc (3)		
PMRA # 2377994	No evidence of neurotoxicity		
28-Day Immunotoxicity (diet)	NOAEL = 471 mg/kg bw/day in \Im		
20-Day minutotoxicity (dict)	LOAEL = 1419 mg/kg bw/day in \Im ; effects included \uparrow spleen wt (\Im)		
Wistar rat	1 + 1 + 1 + 1 + 1 + 1 + 1 + 1 + 1 + 1 +		
PMRA # 2378005			
Special study	Neither S-2200TG nor its metabolites (5-COOH-S-2200, 4-OH-S-2200, 5-CH ₂ OH-S-2200 and		
	5-CA-S-2200-NHM) showed agonistic or antagonistic effects on hERα and hAR from HeLa		
Ovary: Reporter Genes Assay	cells derived from human uterine cervix carcinoma.		
(hER α and hAR assays)			
	Note:Test substances were assayed only once.		
PMRA # 2378010			
Ovary: Steroidogenesis Assay	S-2200TG did not modulate testosterone or estradiol production under the conditions of this		
DMD A # 227800C	study up to 30 μ M (viability \ge 80%) in H295R cells.		
PMRA # 2378006 Liver and thyroid changes	7 dou tractment		
(diet)	7-day treatment: ≥23/26 mg/kg bw/day in ♂/♀:↑T4-UGT (♂)		
(diet)	\geq 116/131 mg/kg bw/day in $\Im/2$: \uparrow CYP2B; \uparrow rel liver wt (\Im)		
Wistar Rat	\geq 379/420 mg/kg bw/day in $\Im/2$: \downarrow fc, diffuse hepatocellular hypertrophy, \uparrow DNA synthesis; \downarrow		
i istal rat	bw, \downarrow bwg, \uparrow rel liver wt (\Diamond); \uparrow thyroid wt, diffuse thyroid follicular cell hypertrophy (\bigcirc)		
PMRA # 2378008	744/812 mg/kg bw/day in \mathcal{O}/\mathcal{P} : enlarged liver, bile duct brown pigment, peribiliary		
	inflammation, $\downarrow T4$ (\Diamond); $\downarrow bw$, $\downarrow bwg$, \uparrow abs liver wt, $\uparrow T4$ -UGT, $\uparrow TSH$ (\bigcirc)		
	14-day treatment:		
	796/952 mg/kg bw/day in \mathcal{O}/\mathcal{Q} : \downarrow bwg, \uparrow liver wt, \uparrow thyroid wt, diffuse hepatocellular		
	hypertrophy ($\mathcal{O}^{\mathbb{Q}}$) with \uparrow severity versus 7-day treatment (\mathcal{O}), \downarrow T4, \uparrow TSH ; \uparrow DNA synthesis		
	(\circlearrowleft); \downarrow bw , diffuse thyroid follicular cell hypertrophy, \downarrow T3 (\updownarrow)		
	7-day treatment + recovery:		
	bw and bwg recoveries, fc recovery, liver wt still significantly \uparrow in ∂ , recovery for thyroid wt,		
	no remarkable pathology findings at necropsy, \uparrow CYP4A, recovery for thyroid hormone levels except for T4 in \Im		

Study Type/Animal/PMRA #Study ResultsAcute Oral Toxicity $LD_{50\circ} > 2000 \text{ mg/kg bw}$ 2-CH2OH-S-2200Low ToxicityWistar RatPMRA # 2377935Bacterial Reverse Mutation Assay (Ames test)Negative in <i>S. typhimurium</i> strains (TA 98, TA100, TA1535, TA15 TA1538) and <i>E. coli</i> WP2uvrA in the presence and absence of meta activation.PMRA#2377976 Acute Oral ToxicityLD _{50\overline{2}} > 2000 mg/kg bw Low Toxicity	
2-CH2OH-S-2200Low ToxicityWistar RatPMRA # 2377935Bacterial Reverse Mutation Assay (Ames test) 2-CH2OH-S-2200Negative in S. typhimurium strains (TA 98, TA100, TA1535, TA15 TA1538) and E. coli WP2uvrA in the presence and absence of meta activation.PMRA#2377976 Acute Oral ToxicityLD $_{50\circ} > 2000 \text{ mg/kg bw}$	
Wistar RatPMRA # 2377935Bacterial Reverse Mutation Assay (Ames test) 2-CH2OH-S-2200PMRA#2377976 Acute Oral ToxicityLD50 \circ 2000 mg/kg bw	
PMRA # 2377935Bacterial Reverse Mutation Assay (Ames test) 2-CH2OH-S-2200Negative in S. typhimurium strains (TA 98, TA100, TA1535, TA15 TA1538) and E. coli WP2uvrA in the presence and absence of meta activation.PMRA#2377976 Acute Oral Toxicity $LD_{50\circ} > 2000 \text{ mg/kg bw}$	
PMRA # 2377935Bacterial Reverse Mutation Assay (Ames test) 2-CH2OH-S-2200Negative in S. typhimurium strains (TA 98, TA100, TA1535, TA15 TA1538) and E. coli WP2uvrA in the presence and absence of meta activation.PMRA#2377976 Acute Oral Toxicity $LD_{50\circ} > 2000 \text{ mg/kg bw}$	
Bacterial Reverse Mutation Assay (Ames test)Negative in S. typhimurium strains (TA 98, TA100, TA1535, TA15 TA1538) and E. coli WP2uvrA in the presence and absence of meta activation.PMRA#2377976 Acute Oral ToxicityLD_ $50\circ$ > 2000 mg/kg bw	
Bacterial Reverse Mutation Assay (Ames test)Negative in S. typhimurium strains (TA 98, TA100, TA1535, TA15 TA1538) and E. coli WP2uvrA in the presence and absence of meta activation.PMRA#2377976 Acute Oral ToxicityLD_ $50\circ$ > 2000 mg/kg bw	
Assay (Ames test)TA1538) and <i>E. coli</i> WP2uvrA in the presence and absence of meta activation.PMRA#2377976 $LD_{50\circ} > 2000 \text{ mg/kg bw}$	
2-CH2OH-S-2200activation.PMRA#2377976Acute Oral ToxicityLD50 \circ > 2000 mg/kg bw	37 and
2-CH2OH-S-2200activation.PMRA#2377976Acute Oral ToxicityLD50 \circ > 2000 mg/kg bw	
Acute Oral Toxicity $LD_{50\circ} > 2000 \text{ mg/kg bw}$	
Acute Oral Toxicity $LD_{50\circ} > 2000 \text{ mg/kg bw}$	
2-COOH-S-2200 Low Toxicity	
Wistar Rat	
PMRA # 2377932	
Bacterial Reverse Mutation Negative in <i>S. typhimurium</i> strains (TA 98, TA100, TA1535, TA15	
Assay (Ames test) TA1538) and <i>E. coli</i> WP2uvrA in the presence and absence of meta	abolic
2-COOH-S-2200 activation.	
DMD A #2277072	
PMRA#2377973	nation (nalative
In vitro Mammalian Positive at $2200 \ \mu g/mL$ after 24 hours in absence of metabolic active contraction and the second	ation (relative
Clastogenicity Assay cell growth 48.8%) in Chinese hamster lung cells (CHL/IU).	
(chromosomal aberration)	
2-COOH-S-2200	
PMRA#2377983	
Micronucleus Assay Negative in BDF ₁ [SPF] mice.	
2-COOH-S-2200 No mortality, clinical signs of toxicity and no differences in body w	veight
compared to controls.	
PMRA #2377986	
Acute Oral Toxicity $LD_{50\circ} > 2000 \text{ mg/kg bw}$	
4-OH-S-2200 Low Toxicity	
Wistar Rat	
DMD A # 2277044	
PMRA # 2377944 Pasterial Payerse Mutation Negative in S. tunkingurium strains (TA 08, TA 100, TA 1525, TA 15	27 and
Bacterial Reverse Mutation Negative in S. typhimurium strains (TA 98, TA100, TA1535, TA15 $TA1538$) and E. coli WP2uurA in presence and absence of metabol	
Assay (Ames test) 4-OH-S-2200 TA1538) and <i>E. coli</i> WP2uvrA in presence and absence of metabol	ic activation.
+-011-3-2200	
PMRA#2377977	
Acute Oral Toxicity $300 \text{ mg/kg bw} < \text{LD}_{502} < 2000 \text{ mg/kg bw}$	
5-COOH-S-2200 High Toxicity $F = \frac{1}{2000} + \frac{1}{2000}$	
Wistar Rat All the animals dosed at 2000 mg/kg bw died within 24 hours.	
PMRA # 2377933 2000 mg/kg bw: the animals presented retention of a white fluid in	the stomach,
yellowish-white fluid content in the small intestine and foamy fluid	in the trachea

Bacterial Reverse Mutation	Negative in S. typhimurium strains (TA 98, TA100, TA1535, TA1537 and
Assay (Ames test)	TA1538) and E. coli WP2uvrA in presence or absence of metabolic activation.
5-COOH-S-2200	
PMRA#2377975	
In vitro Mammalian Cell Assay	Negative in V79-HPRT cells in presence and absence of metabolic activation.
(forward gene mutation)	
5-COOH-S-2200	
PMRA#2377981	
In vitro Mammalian	Negative in Chinese hamster lung cells (CHL/IU) in presence and absence of
Clastogenicity Assay	metabolic activation.
(chromosomal aberration)	
5-COOH-S-2200	
PMRA#2377984	
Acute Oral Toxicity	$LD_{50^{\circ}} > 2000 \text{ mg/kg bw}$
De-Xy-S-2200	Low Toxicity
Wistar Rat	
PMRA # 2377931	
Bacterial Reverse Mutation	Negative in S. typhimurium strains (TA 98, TA100, TA1535, TA1537 and
Assay (Ames test)	TA1538) and <i>E. coli</i> WP2uvrA in presence and absence of metabolic activation.
De-XY-S-2200	
PMRA#2377972	

Table 7Toxicology Endpoints for Use in Health Risk Assessment for Mandestrobin

Exposure Scenario	Study	Point of Departure and Endpoint	CAF ¹ or Target MOE
Acute dietary general population	An ARfD is not required		
Repeated dietary	Rat chronic toixicity/oncogenicity study	NOAEL= 27 mg/kg bw/day; effects included reduced body weight/body weight gains as well as liver and kidney effects	100
	ADI = 0.3 mg/kg bw/day		
Short-term dermal	Rat 28-day dermal toxicity	NOAEL= 1000 mg/kg bw/day	
Intermediate-term dermal	study	LOAEL was not determined due to absence of effects up to the limit dose.	100
Short-term inhalation ² Intermediate-term inhalation ²	Rat reproductive toxicity study	NOAEL= 56 mg/kg bw/day; effects included increased bile duct brown pigmentation and periductular inflammatory cell infiltration in F_1 parental generation.	100
Non-dietary oral ingestion (short- term)	Rat reproductive toxicity study	NOAEL= 56 mg/kg bw/day; effects included increased bile duct brown pigmentation and periductular inflammatory cell infiltration in F_1 parental generation.	100
Cancer	An increase in benign sex cord-stromal tumours in females in the rat oncogenicity study was considered equivocal based on the weight of evidence. The endpoints selected for the non-cancer risk assessment are protective of these equivocal findings.		

¹CAF (composite assessment factor) refers to a total of uncertainty and *Pest Control Products Act* factors for dietary assessments; MOE refers to a target MOE for occupational and residential assessments ² Since an oral NOAEL was selected, an inhalation absorption factor of 100% (default value) was used in route-to-

route extrapolation.

Table 8	Integrated Food Residue Chemistry Summary	
---------	---	--

NATURE OF THE RESIDUE IN LETTUC		CE PMRA # 2377840		77840	
Radiolabel Position	[Phenoxy- ¹⁴ C]-S-2200; [Ph- ¹⁴ C]-S-2200 \downarrow CH _ξ $H_{\xi}C$ \downarrow CCH _ξ CCH_{ξ} CONHCH _ξ		00 [Benzyl- ¹⁴ C	$[Benzyl-^{14}C]-S-2200; [Bz-^{14}C]-S-2200$ $H_{C}CH;$ $CCH_{\zeta}CCH_{\zeta}$	
In all studies, the <i>R/S</i> ratio of	S-2200 remain	ed approximately 1:1, ind	icating no R/S isom	erization.	
Test Variety		Lactuca sativa (Buttercr	unch)		
Test Site		In individual pots in gree	enhouse		
Formulation		The test material was for (SC) with an approximat			
Treatment/Pata 1 foliar application o		1 foliar application of 80	00 g a.i./ha at BBCH 43 followed by a 2 nd /ha at BBCH 48 for a total of 1600 g a.i./ha		
PIII (deve)		Immature lettuce leaves: 5 days after the 1 st application Mature lettuce leaves: 5 days after the 2 nd application			
Analytical Method for Over		LSC and combustion			
Identification & Characterization		HPLC and TLC co-chromatography			
Extraction Solvents		Acetonitrile (surface wash only); acetone/water (80:20; v/v);			
Post-Extraction Solids (PES)		acetone/water/hydrochloric acid (80:20:1; v/v/v) Enzymatic hydrolysis with Driselase (enzyme mixture of fungal carbohydrolases), mild acid hydrolysis (0.1M HCl, 40°C, overnight) and mild base hydrolysis (0.1M NaOH, 40°C, overnight)			
Storage Stability		Representative samples from both radiolabels were re-extracted and analysed to verify the stability of the analytes during freezer storage (5 months). Similar profiles were obtained, which indicated stability of [¹⁴ C]-S-2200 metabolites in lettuce samples.			
Matrices	Total Rate	PHI	[Ph- ¹⁴ C]-S-220	$0 \qquad [Bz-^{14}C]-S-2200$	
	(g a.i./ha)	(days)	TRR (ppm, i	n S-2200 equivalents)	
Immature Lettuce Leaves	800	5 (after 1 st appl.)	35.11	27.94	
Mature Lettuce Leaves	1600 5 (after 2^{nd} appl.)		43.14	41.59	
Metabolites Identified		Major Metabol	ites (>10% of the TR	RR)	
Radiolabel Position	[Ph- ¹⁴ C]-S-2200		[Bz	[Bz- ¹⁴ C]-S-2200	
Lettuce – Immature leaves	S-2200			S-2200	
Lettuce - Mature leaves	S-2200			S-2200	
Most of the radioactivity was recovered in the surface washes (78.5-88.4% of the TRR). The metabolites De-Xy-S2200, MCBX, and conjugates of 2-CH ₂ OH-S-2200, 4-OH-S-2200, 5-CH ₂ OH-S-2200 and 5-COOH-S-2200 were detected along with several other minor metabolites.					

NATURE OF THE RESIDUE	N RAPES	EED			PMRA # 2377845		
Radiolabel Position			[Ph- ¹⁴ C]-S-2200		[Bz- ¹⁴ (C]-S-2200	
Test Variety		Bra	ussica napus L. (Phoe	nix L	iberty Link)		
Test Site		mat	ndividual pots mainta turity)		-	-	
Formulation			e test material was for () with an approximat		200 R:S isomer rati	o of 50:50	
Treatment/Rate			<u>T # 1</u> : 1 foliar lication of 400 g a.i./ BBCH 55-61	ha	<u>TRT # 2</u> : 1 foliar a a.i./ha at BBCH 66 application 14-day total of 800 g a.i./h	s after the 1 st for a	
PHI (days)		54	(seeds)		14 (forage); 40 (se	eds)	
Analytical Method for Overall Identification & Characterizati			C and combustion LC and TLC co-chro	mato	graphy		
Extraction Solvents			etonitrile (forage surf tone/water (80:20; v/ /v)				
Post-Extraction Solids (PES)			Sequential enzyme hydrolysis with amylase and protease followed by weak acid hydrolysis (1M HCl, 40°C, overnight), strong acid hydrolysis (6M HCl, 80°C, 4 hours), weak base hydrolysis (0.1M NaOH, 40°C, overnight) and strong base hydrolysis (6M NaOH, 80°C, overnight)				
Storage Stability			Representative samples from both radiolabels were re-extracted and analysed to verify the stability of the analytes during freezer storage (7-11 months). Similar profiles were obtained, which indicated stability of [¹⁴ C]-S-2200 metabolites in rapeseed samples.				
Matrices	Total Ra			[Ph- ¹⁴ C]-S-2200	[Bz- ¹⁴ C]-S-2200	
Wathces	(g a.i./ha	a)	Growth Stage		TRR (ppm, in S-22	200 equivalents)	
Rapeseed – Seeds (TRT #1)	400		54 (BBCH 89)		0.05	0.11	
Rapeseed – Seeds (TRT #2)	800		40 (BBCH 89)		0.47	0.64	
Rapeseed – Forage (TRT #2)	800		14 (BBCH 55-61)		3.99	3.44	
Metabolites Identified		Major Metabolites (>10% of the TRR)					
Radiolabel Position	[]	Ph- ¹⁴ C]-S-2200			[Bz- ¹⁴ C]-S-2200		
Rapeseed (Seeds) – TRT #1			None	None			
Rapeseed (Seeds) – TRT #2	S-2200,	4-O	0H-S-2200 (conj.)		S-2200, 4-OH-S-2200 (conj.)		
Rapeseed (Forage) – TRT #2	(conj.),	00, 2-CH ₂ OH-S-2200 S , 4-OH-S-2200 (conj.)			S-2200, 2-CH ₂ OH-S-2200 (conj.), 4-OH-S- 2200 (conj.)		
For forage, the radioactivity in the the TRR was present in the extract extractable fractions from each tre MCBX and a conjugate of 5-CH ₂	table fraction	ons. oup. 1	For seed, approximat	tely 8	1 to 99% of the TRI	R was present in	
NATURE OF THE RESIDUE I	N WHEAT	Г			PMRA # 2377842		
Radiolabel Position			[Ph- ¹⁴ C]-S-2200		[B z- ¹⁴ (C]-S-2200	
Test Variety			ticum L. (Promontory				
Test Site		mat	ndividual pots mainta turity)		-	~	
Formulation		(SC		te S-2	200 R:S isomer rati	o of 50:50	
Formulation Treatment/Rate			(SC) with an approximate S-2200 R:S isomer ratio of 50:50 A single foliar application of 300 g a.i./ha at BBCH 32 (104 days before final harvest)				

Extraction Solventsacetone/water/hydred Sequential enzyme (0.1M HCl, 40°C or 40°C overnight)Post-Extraction Solids (PES)Representative same analysed to verify to months). Similar price [14C]-S-2200 metableMatricesTotal Rate (g a.i./ha)PHI (days) Growth StageWheat Forage3007 (BBCH 37)Wheat straw300104 (BBCH 97)Metabolites IdentifiedMajor MetalRadiolabel Position[Ph-14C]-S-2200 (conj.), 4) 2200 (conj.)Wheat strawSolo 200, 2-CH2OH-S-2200 (conj.), 4) 2200 (conj.)Wheat strawNone	o-chromatographyce wash only); acetone/water (80:20; v/v); rochloric acid (80:20:1; v/v/v)e hydrolysis with Driselase, mild acid hydrolysis overnight) and mild base hydrolysis (0.1M NaOH, mples from both radiolabels were re-extracted and the stability of the analytes during freezer storage rofiles were obtained, which indicated stability of bolites in wheat samples)/[Ph-14C]-S-2200[Bz-14C]-S-2200geTRR (ppm, in S-2200 equivalents)7)11.1410.4437)6.219.0492)0.010.0992)1.852.49abolites (>10% of the TRR)[Bz-14C]-S-2200				
Identification & CharacterizationHPLC and TLC co- Acetonitrile (surfac acetone/water/hydra Sequential enzyme (0.1M HCl, 40°C or 40°C overnight)Post-Extraction Solids (PES)Sequential enzyme (0.1M HCl, 40°C or 40°C overnight)Storage StabilityRepresentative sam analysed to verify timonths). Similar pr [¹⁴ C]-S-2200 metabMatricesTotal Rate (g a.i./ha)PHI (days) Growth StagWheat Forage3007 (BBCH 37Wheat straw300104 (BBCH 9Metabolites IdentifiedMajor MetalRadiolabel Position $[-1^{4}C]$ -S-2200 (con)Wheat forageS-2200, 2-CH2OH-S-2200 (con)Wheat forageS-2200, 2-CH2OH-S	ce wash only); acetone/water (80:20; v/v);rochloric acid (80:20:1; v/v/v)rochloric acid (80:20:1; v/v/v)e hydrolysis with Driselase, mild acid hydrolysisovernight) and mild base hydrolysis (0.1M NaOH,nples from both radiolabels were re-extracted andthe stability of the analytes during freezer storageTRR (ppm, in S-2200Bz-14C]-S-2200geTRR (ppm, in S-2200 equivalents)7)11.1437)6.2192)0.0192)1.852.49abolites (>10% of the TRR)				
Extraction Solventsacetone/water/hydroPost-Extraction Solids (PES)Sequential enzyme (0.1M HCl, 40°C o 40°C overnight)Storage StabilityRepresentative sam analysed to verify ti months). Similar pr [14C]-S-2200 metablicationMatricesTotal Rate (g a.i./ha)PHI (days) Growth StagWheat Forage3007 (BBCH 37)Wheat grain300104 (BBCH 9)Wheat straw300104 (BBCH 9)Wheat forageS-2200, 2-CH2OH-S-2200 (con)Wheat forage <t< th=""><th>rochloric acid (80:20:1; $v/v/v$)e hydrolysis with Driselase, mild acid hydrolysise hydrolysis with Driselase, mild acid hydrolysisovernight) and mild base hydrolysis (0.1M NaOH,nples from both radiolabels were re-extracted andthe stability of the analytes during freezer storageorofiles were obtained, which indicated stability ofbolites in wheat samples)/[Ph-14C]-S-2200(Bz-14C]-S-2200geTRR (ppm, in S-2200 equivalents)7)11.1437)6.2192)0.010.0992)1.852.49abolites (>10% of the TRR)</th></t<>	rochloric acid (80:20:1; $v/v/v$)e hydrolysis with Driselase, mild acid hydrolysise hydrolysis with Driselase, mild acid hydrolysisovernight) and mild base hydrolysis (0.1M NaOH,nples from both radiolabels were re-extracted andthe stability of the analytes during freezer storageorofiles were obtained, which indicated stability ofbolites in wheat samples)/[Ph-14C]-S-2200(Bz-14C]-S-2200geTRR (ppm, in S-2200 equivalents)7)11.1437)6.2192)0.010.0992)1.852.49abolites (>10% of the TRR)				
Post-Extraction Solids (PES)(0.1M HCl, 40°C or 40°C overnight)Representative sam analysed to verify to months). Similar propriation proprision propriation propriation proprision propriation	overnight) and mild base hydrolysis (0.1M NaOH,mples from both radiolabels were re-extracted andthe stability of the analytes during freezer storageorfiles were obtained, which indicated stability ofbolites in wheat samples)/[Ph-14C]-S-2200(Bz-14C]-S-2200[Bz-14C]-S-2200geTRR (ppm, in S-2200 equivalents)7)11.1410.4437)6.2192)0.010.0992)1.852.49abolites (>10% of the TRR)[Bz-14C]-S-2200				
Storage Stabilityanalysed to verify to months). Similar prosted to the second seco	The stability of the analytes during freezer storage profiles were obtained, which indicated stability of bolites in wheat samples (/ [Ph- ¹⁴ C]-S-2200 [Bz- ¹⁴ C]-S-2200 ge TRR (ppm, in S-2200 equivalents) 7) 11.14 10.44 87) 6.21 9.04 92) 0.01 0.09 92) 1.85 2.49 abolites (>10% of the TRR)				
Matrices(g a.i./ha)Growth StageWheat Forage 300 7 (BBCH 37)Wheat hay 300 14 (BBCH 37)Wheat grain 300 104 (BBCH 97)Wheat straw 300 104 (BBCH 97)Metabolites IdentifiedMajor MetabolitesRadiolabel Position[Ph-14C]-S-2200Wheat forageS-2200, 2-CH ₂ OH-S-2200 (conj.), 4 2200 (conj.)Wheat strawNone	Image TRR (ppm, in S-2200 equivalents) 7) 11.14 10.44 87) 6.21 9.04 92) 0.01 0.09 92) 1.85 2.49 abolites (>10% of the TRR) [Bz- ¹⁴ C]-S-2200				
Matrices(g a.i./ha)Growth StageWheat Forage 300 7 (BBCH 37)Wheat hay 300 14 (BBCH 37)Wheat grain 300 104 (BBCH 97)Wheat straw 300 104 (BBCH 97)Metabolites IdentifiedMajor MetabolitesRadiolabel Position[Ph-14C]-S-2200Wheat forageS-2200, 2-CH ₂ OH-S-2200 (conj.), 4 2200 (conj.))Wheat strawNone	Image TRR (ppm, in S-2200 equivalents) 7) 11.14 10.44 87) 6.21 9.04 92) 0.01 0.09 92) 1.85 2.49 abolites (>10% of the TRR) [Bz- ¹⁴ C]-S-2200				
Wheat Forage 300 7 (BBCH 37) Wheat hay 300 14 (BBCH 37) Wheat grain 300 14 (BBCH 97) Wheat grain 300 104 (BBCH 97) Wheat straw 300 104 (BBCH 97) Metabolites Identified Major Metabolites Radiolabel Position [Ph- ¹⁴ C]-S-2200 Wheat forage S-2200, 2-CH ₂ OH-S-2200 (conj.), 4 2200 (conj.) Wheat straw None	(11) (12) 7) 11.14 10.44 37) 6.21 9.04 92) 0.01 0.09 92) 1.85 2.49 Bolites (>10% of the TRR)				
Wheat hay 300 $14 (BBCH 37)$ Wheat grain 300 $104 (BBCH 97)$ Wheat straw 300 $104 (BBCH 97)$ Wheat straw 300 $104 (BBCH 97)$ Metabolites Identified Major Metal Radiolabel Position [Ph- ¹⁴ C]-S-2200 Wheat forage S-2200, 2-CH ₂ OH-S-2200 (conj.), 4 Wheat hay S-2200, 2-CH ₂ OH-S-2200 (conj.), 4 Wheat straw None	37) 6.21 9.04 92) 0.01 0.09 92) 1.85 2.49 abolites (>10% of the TRR) [Bz- ¹⁴ C]-S-2200				
Wheat grain 300 104 (BBCH 9) Wheat straw 300 104 (BBCH 9) Metabolites Identified Major Metal Radiolabel Position [Ph- ¹⁴ C]-S-2200 Wheat forage S-2200, 2-CH ₂ OH-S-2200 (conj.), 4 Wheat straw S-2200, 2-CH ₂ OH-S-2200 (conj.), 4 Wheat straw None	92) 0.01 0.09 92) 1.85 2.49 abolites (>10% of the TRR) [Bz- ¹⁴ C]-S-2200				
Wheat straw 300 $104 (BBCH 9)$ Metabolites IdentifiedMajor MetalRadiolabel Position $[Ph-^{14}C]-S-2200$ Wheat forageS-2200, 2-CH ₂ OH-S-2200 (conj.), 4 2200 (conj.))Wheat strawS-2200, 2-CH ₂ OH-S-2200 (conj.), 4 2200 (conj.))	92) 1.85 2.49 abolites (>10% of the TRR) [Bz- ¹⁴ C]-S-2200				
Metabolites IdentifiedMajor MetabolitesRadiolabel Position[Ph-14C]-S-2200Wheat forageS-2200, 2-CH2OH-S-2200 (conWheat hayS-2200, 2-CH2OH-S-2200 (conj.), 4 2200 (conj.)Wheat strawNone	abolites (>10% of the TRR) [Bz- ¹⁴ C]-S-2200				
Radiolabel Position[Ph-14C]-S-2200Wheat forageS-2200, 2-CH2OH-S-2200 (conWheat hayS-2200, 2-CH2OH-S-2200 (conj.), 4 2200 (conj.)Wheat strawNone	[Bz- ¹⁴ C]-S-2200				
Wheat forage S-2200, 2-CH ₂ OH-S-2200 (con Wheat hay S-2200, 2-CH ₂ OH-S-2200 (conj.), 4 Wheat straw None					
Wheat hayS-2200, 2-CH2OH-S-2200 (conj.), 4 2200 (conj.)Wheat strawNone					
Wheat hayS-2200, 2-CH2OH-S-2200 (conj.), 4 2200 (conj.)Wheat strawNone	onj.) S-2200				
Wheat straw None					
	De-Xy-S-2200				
Wheat grain None	De-Xy-S-2200				
The radioactivity in the surface rinse ranged from $19 - 41\%$ approximately 3% TRR in the straw samples from both treatmer present in the extractable fraction of the forage, hay, straw and g CH ₂ OH-S-2200 (free), 4-OH-S-2200 (free), 5-CH ₂ OH-S-2200 were considered minor metabolites. S-2200 was not detected in TRR was very low in these samples. De-Xy-S-2200 accounted forage, hay and straw samples and accounted for 61% of the TRI	ent groups. Approximately 53 - 73% of the TRR v grain from both wheat treatment groups. MCBX, (free and conjugated), and 5-COOH-S-2200 (fr the wheat grain from either treatment group but 1 for 2 to 12% of the TRR (0.14 - 0.33 ppm) in				
S-2200 treatment group.	DMD A # 227915(/2279157				
Nature of the Residue in Soybeans and Corn (Radiotracer)	PMRA # 2378156/2378157				
Test VarietySoybean: PioneerFormulationFormulated product1.20 MBq/mg a					
Treatment/Rate[Ph-14C]-S-2:Soybean: Seed/9.33Corn: 11.03 g/1	2200 [Bz- ¹⁴ C]-S-2200 3 g/100 kg Soybean: Seed/10.72 g/100 kg (100 kg Corn: 11.55 g/100 kg				
Analytical Method for Overall TRR	LSC and combustion				
	PLC and TLC co-chromatography				
Radiolabel Position[Ph- ¹⁴ C]-S-2MatrixIPD					
MatricesTRR (ppnSoybean forage0.027, 0.03					
, ,					
Soybean hay 0.030, 0.02 Soybean pod with seed <0.005, <0.0					
Soybean pod with seed <0.005, <0.0					
Soybean mature seed<0.005, <0.0Corn kernels plus cob<0.005, <0.0					
Corn forage <0.005, <0.0					
Corn stover <0.005, <0.0					

Corn grain	1		<0.0	05 <0.005		<0.005, <0.0	005			
		ie in treated R	<0.005, <0.005<0.005, <0.005AC was below the minimum quantification limit (MQL) of 0.005 ppm in all							
RACs, wit	th the exception	of soybean (for	rage and hay), a	nd corn (stover)	. Samples of so	ybean forage a	nd hay grown			
from seeds	s treated with [Pl	$n^{-14}C$]-S-2200	and [Bz-14C]-S-2	2200, and corn	stover from see	ds treated with	[Bz- ¹⁴ C]-S-			
	extracted with a									
	t S-2200 was det									
	ce amounts of 2-									
	<5.1% of the TR					00, which was	observed in			
	orage (12.3% of 1									
Confined Accumulation in Rotational Crops – Lettuce, wheat, carrotPMRA # 2377934										
Test Site			Each plot consi container main			c sheet-lined w	ooded			
Formulati	ion		The test materia with an approx			1	ntrate (SC)			
	l Method for O		LSC and comb							
Identification	tion & Charact	erization	HPLC and TLC							
Extraction	n Solvents		Acetone/water $v/v/v$)	(80:20; v/v); ac	etone/water/hyo	drochloric acid	(80:20:1;			
			Sequential enzy	yme hydrolysis	with Driselase	or amylase, mil	d acid			
Post-Fytr	action Solids (P	FS)	hydrolysis (0.1							
1 051-12411	action Solius (1	L3)	NaOH, 40°C or							
			hours) and strop							
			Samples were s							
a, a			Representative							
Storage St	tability		both radiolabel							
			the analytes during freezer storage. Similar profiles were obtained, which indicated stability of $[^{14}C]$ -S-2200 metabolites in the tested samples.							
Radiolabe	Desition		[Ph- ¹⁴ C]-S-2200 metabolites in the tested samples.							
	op/Crop	Rate	PBI G 11 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1							
	p/Variety	(g a.i./ha)	(days)	Growth stag	ge at harvest	Harvest	ed RAC			
	ace/Leafy	1600	30, 120, 365		50% mature size Immature heads					
vegetabl	e/Salad Bowl			Mat	•	Mature heads				
Whe	eat/Cereal	1,000	20 120 255		n elongation	Forage (immature)				
	lanca Royale	1600	30, 120, 365		to soft dough	Нау				
				Mat	urity	Grain a	nd straw			
	oot vegetable/ Half Long 126	1600	30, 120, 365	Mat	urity	Tops (foliag	e) and roots			
	<u> </u>			Overall TRR (I	opm, in S-2200	equivalents)				
	otational		[Ph- ¹⁴ C]-S-220			Bz- ¹⁴ C]-S-220	0			
cro	p sample	30 DAT	120 DAT	365 DAT	30 DAT	120 DAT	365 DAT			
	Forage	2.73	0.14	0.10	2.54	0.31	0.26			
W/L	Hay	1.56	0.29	0.35	4.54	0.72	0.74			
Wheat	Straw 1.32 0.34 0.31		0.82	0.59	0.38					
	Grain	0.04	0.04	< 0.01	0.12	0.20	< 0.01			
Lettuce	Immature	0.33	0.03	0.07	0.32	0.08	0.07			
Lenuce	Mature	0.08	0.02	0.02	0.22	0.05	0.02			
	Mature roots	0.05	0.03	< 0.01	0.04	0.04	< 0.01			
Carrot	Mature									

Metabolites	Identified		Major Metabolites (>10% of the TRR)					
Matrices	PBI (days)		[Ph- ¹⁴ C]-S-2200	[Bz- ¹⁴ C]-S-2200				
	30	5-CH ₂ OH-	-S-2200 (conj.), 4-OH-S-2200 (conj.)	5-CH ₂ OH-S-2200 (conj.), 4-OH-S- 2200 (conj.)				
Immature Lettuce	120		None	4-OH-S-2200 (conj.)				
Lettuce	365	S-2200, 4-0	H-S-2200 (conj.), 5-CH ₂ OH-S- 2200 (conj.)	S-2200, 4-OH-S-2200 (conj.)				
	30	S-2200, 4-0	0H-S-2200 (conj.), 5-CH ₂ OH-S- 2200 (conj.)	4-OH-S-2200 (conj.), 5-CH ₂ OH-S- 2200 (conj.)				
Mature Lettuce	120		None	None				
Lettuce	365	4-OH-S-22	200 (conj.), 5-CH ₂ OH-S-2200 (conj.)	None				
XXII - C	30	4-OH-S-22	CH ₂ OH-S-2200 (conj.), 200 (conj.), 5-CH ₂ OH-S-2200 (conj.)	5-CH ₂ OH-S-2200 (conj.), 4-OH-S-2200 (conj.)				
Wheat forage	120	4-OH-S-22	200 (conj.), 5-CH ₂ OH-S-2200 (conj.)	None				
	365	5-0	CH ₂ OH-S-2200 (conj.)	None				
	30		200 (conj.), 5-CH ₂ OH-S-2200 nj.), 2-CH ₂ OH-S-2200	4-OH-S-2200 (conj.)				
Wheat hay	120	4-OH-S-22	200 (conj.), 5-CH ₂ OH-S-2200 (conj.)	None				
	365		None	None				
W/h a at atmost	30/120		None	None				
Wheat straw	365		None	4-OH-S-2200 (conj.)				
Wheat grain	30/120/365		Not Profiled ¹	Not Profiled ¹				
	30		None	None				
Carrot tops	120		None	S-2200				
	365		None	5-CH ₂ OH-S-2200 (conj.)				
	30		S-2200	S-2200				
Carrot roots	120		S-2200 2-CH ₂ OH-S-2200	S-2200				
	365		Not profiled ¹	Not profiled ¹				
from both treatm radioactive resid followed by ext metabolite De-2 benzyl-labelled	50 - 90% of the nent groups. The dues in rotation ensive metabol Xy-S-2200, aris 30 DAT imma	e TRR was pre ne results gene al crops seede ism to polar m ing from cleav ture and matur	sent in the extractable fraction of rated in this study demonstrated th d in soil 30, 120, and 365 days aft etabolites and incorporation into t vage of the ether linkage, was foun	er treatment with ¹⁴ C-S-2200, the constituents of the plant. The and at very low levels only in the included MCBX, 5-COOH-S-2200,				
NATURE OF 7	THE RESIDU	E IN LAYING	G HEN	PMRA # 2377838				
ppm in the diet)	, via gelatin ca	psule, once dai		ppm and [Bz- ¹⁴ C]-S-2200 at 13.15 a were collected daily. Samples of istration of the final dose.				
Form	ulated Test Su	bstance	The approximate S-2200 R:S isomer ratio was 50:50.					
Analytical Met Identification &			LSC and combustion Radio-HPLC using co-chromatography; LC-MS/MS					

Extraction Solvents		Eggs, fat, and muscle: Sequential extraction with hexane, ethyl acetate, acetonitrile and 1% formic acid in acetonitrile Liver and skin: Sequential extraction with hexane, ethyl acetate, acetrontrile, 1% formic acid in acetonitrile, water, 1M HCl, and 1M ammonia solution Liver: Sequential hydrolysis using protease digestion for 18					
Post-Extraction Solids (PES		hours at 37°C, acid hydrolysis with 10M HCl, base hydrolysis with 10M NaOH under reflux					
Storage Stability		Samples were stored and ana	ored and analyzed within 6 months.				
N7 / ·	[]	Ph- ¹⁴ C]-S-2200	[Bz-	¹⁴ C]-S-2200			
Matrices	TRR (ppm)	% of AD	TRR (ppm)	% of AD			
Excreta	-	83.37	-	98.36			
Cage wash	-	1.33	0	0.99			
Muscle (breast)	0.0127	0.007	0.025	0.014			
Muscle (thigh)	0.0144	0.003	0.023	0.005			
Fat (peritoneal)	0.033	0.003	0.032	0.005			
Skin	0.0478	0.003	0.0543	0.003			
Liver	0.295	0.055	0.299	0.063			
Eggs (Day 2-14)	0.051-0.113	0.21	0.050-0.081	0.18			

The majority of the administered dose was excreted, with 85% of the total dose of $[Ph-^{14}C]$ -S-2200 and 99% of the total dose of $[Bz-^{14}C]$ -S-2200 being recovered in the excreta and cage wash. The highest residues were detected in liver and eggs. Following administration of $[Ph-^{14}C]$ -S-2200 to hens, total radioactive residues in eggs reached a maximum on day 11 (0.113 ppm). Similarly, following administration of $[Bz-^{14}C]$ -S-2200 to hens, total radioactive residues in eggs reached a maximum on day 7 (0.081 ppm). The major residue from laying hen dosed with $[Ph-^{14}C]$ -S-2200 and $[Bz-^{14}C]$ -S-2200 was S-2200 in fat (33.9-49.5% of the TRR; 0.011-0.016 ppm), and eggs (33.1-51.2% of the TRR; 0.025-0.058 ppm). Another major residue from laying hen dosed with $[Ph-^{14}C]$ -S-2200 was 4-OH-S-2200 in liver (13.6% of the TRR; 0.040 ppm). Only S-2200 was confirmed by LC-MS.

All other minor metabolites were only tentatively identified based on chromatographic retention times of metabolite standards. The metabolism of $[^{14}C]$ -S-2200 was considered extensive as some tissues contained up to 21 peaks, including diffuse regions of interest. The unextractable radioactivity in liver, remaining after acid, alkaline, and enzymatic hydrolysis (PES), was presumed to be associated mostly with endogenous material, or more polar multi-components.

Metabolites identified	Major Metabolites (>10% of the TRRs)					
Radiolabel Position	[Ph- ¹⁴ C]-S-2200	[Bz- ¹⁴ C]-S-2200				
Egg (Day 11)	S-2200	S-2200				
Liver	4-OH-S-2200	None				
Muscle	None	None				
Skin	None	None				
Fat	S-2200	S-2200				
NATURE OF THE RESIDUE IN L	PMRA # 2377836					

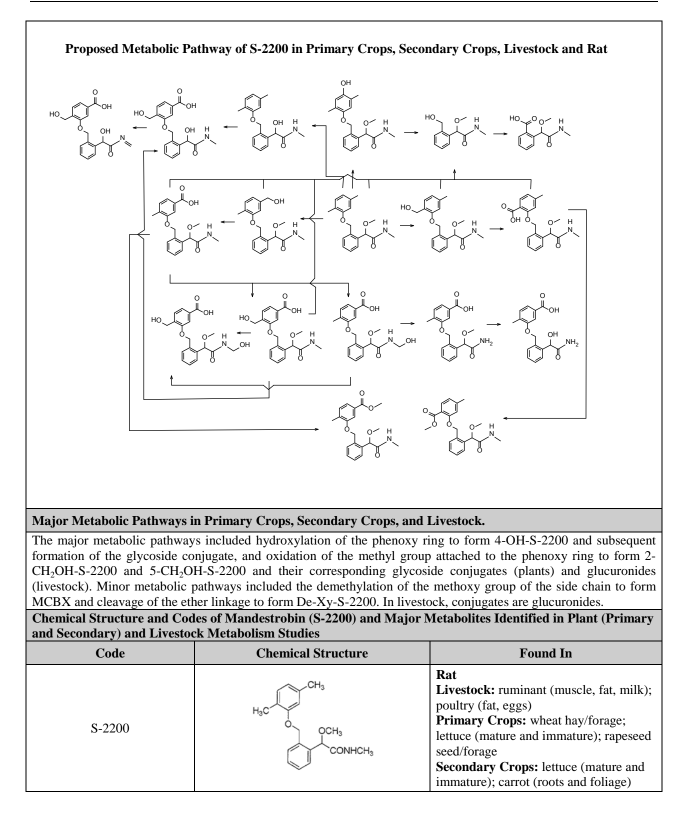
Two lactating goats were dosed orally with [Ph¹⁴C]-S-2200 at 12.65 ppm and [Bz-¹⁴C]-S-2200 at 14.33 ppm in the diet), via gelatin capsule, once daily for 7 days. Samples of excreta were collected once daily and milk was collected twice daily. The goats were euthanized 6 hours after administration of the final dose.

Formulated Test Substance	The approximate S-2200 R:S isomer ratio was 50:50.
Analytical Method for Overall TRR	LSC and combustion
Identification & Characterization	Radio-HPLC using co-chromatography; LC-MS/MS
Extraction Solvents	Fat, milk fat, muscle: Sequential extraction with hexane, ethyl acetate, acetonitrile and 1% formic acid in acetonitrile Liver and kidney: Sequential extraction with hexane, ethyl acetate, acetrontrile, 1% formic acid in acetonitrile, water, 1M HCl, and 1M ammonia solution

Post-Extraction Solids (PE	2S) 37°C, a under r	cid hydrolysis with eflux	10M HCl, base hydrol	stion for 18 hours at ysis with 10M NaOH
Storage Stability	-		alyzed within 6 month	
	[Ph- ¹⁴ C]-	-S-2200	[Bz- ¹⁴ C	C]-S-2200
Matrices	TRR (ppm)	% of AD	TRR (ppm)	% of AD
Urine	-	35.22	-	39.73
Cage wash	-	1.19	-	0.68
Feces	-	- 42.49		38.07
Muscle (flank)	0.012	0.012 0.005		0.003
Muscle (loin)	0.008	0.001	0.014	0.001
Fat (omental)	0.012	0.006	0.028	0.002
Fat (renal)	0.013	0.008	0.034	0.004
Fat (subcutaneous)	0.010	0.001	0.033	< 0.001
Kidney	0.170	0.022	0.412	0.031
Liver	0.319	0.225	0.613	0.289
Milk – fat fraction	0.008-0.033	0.008-0.033 0.002		0.005
Milk – aqueous fraction	0.004-0.010	0.024	0.006-0.018	0.073

S-2200 was extensively metabolised and the majority of the administered dose (79% of [Ph-¹⁴C]-S-2200 and 78% of [Bz-¹⁴C]-S-2200) was excreted in approximately equal proportions in the urine and faeces. Following administration of [Ph-¹⁴C]-S-2200 to goats, total radioactive residues in the aqueous fraction of milk reached a maximum on day 6 (0.01 ppm), and on day 7 in the milk fat (0.033 ppm). Similarly, following administration of [Bz-¹⁴C]-S-2200 to goats, total radioactive residues in the aqueous fraction of milk reached a maximum on day 3 (0.018 ppm) and on day 5 in the milk fat (0.035 ppm). Residues in the tissues, with the exception of the liver, were readily extractable with organic solvents or polar solvents most notably following administration of [Ph-¹⁴C]-S-2200. The major residue from lactating goat dosed with [Ph-¹⁴C]-S-2200 and [Bz-¹⁴C]-S-2200 was S-2200 in fat (22.9-49.6% of the TRR; 0.006-0.007 ppm), muscle (18.2-23.0% of the TRR; 0.002-0.003 ppm), and milk fat fraction (32.7-35.3% of the TRR; 0.011-0.012 ppm); 4-OH-S-2200 (conj.) in kidney (13.3-14.9% of the TRR; 0.025-0.055 ppm); and 5-COOH-S-2200 in kidney (20.2-25.0% of the TRR; 0.043-0.083 ppm), and liver (10.6-19.3% of the TRR; 0.062-0.065 ppm). Other major residues from lactating goat dosed with [Bz-14C]-S-2200 were 2-CH2OH-S-2200 in muscle (10.1% of the TRR; 0.002 ppm), and 5-CA-S-2200-NHM in milk (14.7% of the TRR; 0.003 ppm). LC-MS was used to confirm the identity of S-2200 (milk fat, muscle and fat), 2-CH₂OH-S-2200 (muscle), 5-COOH-S-2200 (liver and kidney) and 4-OH-S-2200 glucuronide (kidney). All other minor metabolites were only tentatively identified based on chromatographic retention times of metabolite standards. The metabolism of $[^{14}C]$ -S-2200 was considered extensive as some tissues contained up to 25 peaks, including diffuse regions of interest. The unextractable radioactivity in liver, remaining after acid, alkaline, and enzymatic hydrolysis (PES), was presumed to be associated mostly with endogenous material, or more polar multi-components.

Metabolites identified	Major Metabolites (>10% of the TRR)
Radiolabel Position	[Ph- ¹⁴ C]-S-2200	[Bz- ¹⁴ C]-S-2200
Milk – fat fraction	S-2200	S-2200
Milk – aqueous fraction	Not analysed	5-CA-S-2200-NHM
Liver	5-COOH-S-2200	5-COOH-S-2200
Kidney	5-COOH-S-2200, 4-OH-S-2200 (conj.)	5-COOH-S-2200, 4-OH-S-2200 (conj.)
Muscle	S-2200	S-2200, 2-CH ₂ OH-S-2200
Fat	S-2200	S-2200



2-CH ₂ -OH-S-2200	HOH ₂ C CH ₃ OCH ₃ CONHCH ₃	Rat Primary Crops: wheat hay/forage Secondary Crops: carrot roots
4-OH-S-2200		Rat Livestock: poultry (liver)
De-Xy-S-2200	HO OCH ₃ CONHCH ₃	Rat Primary Crops: wheat grain/straw; soybean forage (seed treatment) Secondary Crops: lettuce (mature and immature); carrot (roots and foliage)
5-COOH-S-2200	CH ₅ CH ₅ CCOCH ₃ CCNHCH ₅	Rat Livestock: ruminant (kidney, liver)
4-OH-S-2200 (Bound/ Conjugated/ glycosides)		Rat Livestock: ruminant (kidney) Primary Crops: rapeseed seed/forage Secondary Crops: wheat hay/forage/straw; lettuce (mature and immature)
2-CH ₂ -OH-S-2200 (conjugated/glycosides)		Primary Crops: wheat hay/forage; rapeseed forage Secondary Crops: wheat forage
5-CH ₂ -OH-S-2200 (conjugated/glycosides)		Secondary Crops: wheat hay/forage; lettuce (mature and immature); carrot foliage
Freezer Storage Stability		PMRA # 2377901, 2377902
Matrix	Demonstrated storage interval at -18°C (months)	Actual storage interval (months)
Storage stability studies		
Rapeseed (seed)	12 and on-going*	38
Rapeseed (oil)	12	10
Rapeseed (meal)	12	9
Lettuce Barley (grain)	12 12	No associated residue trial
Barley (gram)	12	
Darley (Suaw)	12	

Concurrent stor	age stab	oility studies									
Strawberries	~			nd on-	going'	*		1	9		
Grapes					going'			16			
Grape (juice)				7					7		
Grape (raisins)			5 ai	nd on-	going	*		ç)		
Soybean (seed)				12				8	.5		
Corn forage				12				1	2		
Corn K+CWHR				12	, ,			1	1		
Corn grain				12				ç			
Corn stover			14	9				Ç			
In addition, storagy years, and no sigr * Additional stora CROP FIELD T	nificant c age stabi RIALS	lecline of S- lity studies t	2200 was o be subm	obser nitted	ved. by app	licant at a la	ater date.		-		
Residue trials wer S-2200 4 SC, and rapeseed. Adjuva ±25% of GAP we the case of rapese Crop Field Trial	S-2200 nts were ere included eds, the s on Str	3.2 FS Fung added to ead ded herein. F residues dec rawberries (ticide) con ch spray r Residues c lined in o Foliar)	ntainir nixtur of S-22	ng man e. Only 200 de	destrobin of y trial value cline with ir greas residue	n corn, grape s that were concreasing PH	s, soybeans, onducted at Is in strawbe han LOQ in	strawberrie GAP or wit erries and g the other tr	es, and hin rapes. In ial.	
Number and Loo	cation of	f Field Tria	s								
Region		1	2	2		3	5	10	12	Tota	
CDN Required*		1	(0		0	3	0	1	5	
US Required*		1	0			1	1	2	1	6	
Submitted		1	1			1	3	3	1	10	
*As per DIR98-02	2 and US	SEPA Residu	ue Chemis	stry Te	est Gu	idelines (cro	op group redu	uction).			
Use Pattern											
Approved Use Pa	attern (GAP)				Use Patter	n				
Single foliar appl # of appl.: 4-5 apj Maximum rate: 1 PHI: 0 day	pl. per ye	ear at 7-14 d		ıls –	# of ap Maxim PHI: 0 US tri Single # of ap Maxim PHI: 0	e foliar appl. ppl.: 4 appl. num rate: 16) day als: e foliar appl. ppl.: 4 appl. num rate: 16) day	574-1734 g a. rate: ~420 g	i./ha/season a.i./ha + adj a.i./ha/seaso	uvant		
Total Rate	PHI				S-2200 Residue Levels (ppm)						
(g a.i./ha)	(days)	n	Min. [#]	M	ax. #	LAFT *	HAFT [*]	Median [*]	Mean *	SD *	
US trials											
1675-1702	0	8	0.33	2	.12	0.45	2.05	0.84	0.97	0.52	
				L		1	1	I	1		
Canadian trials							1			1	
1674-1734	0	2	0.6	1	.07	0.62	0.995	0.81	0.81	0.27	
Canadian trials 1674-1734 Combined trials	0	2	0.6	1	.07	0.62	0.995	0.81	0.81	0.27	
1674-1734	0	2	0.6	I	.07	0.62	0.995 2.05	0.81	0.81	0.27	

Crop Field Trials of	on Grap	es (Folia	r)		I	PMRA #	#s 237791	3 (CDN)	and 23	877920 (US	5)
Number and Locat	tion of F	ield Tria	als								
Region		1			5		10		11		Total
CDN Required*		0			4		0		1		5
US Required*		2			0		5		2		9
Submitted		2			3		7		3		15
*As per DIR98-02 a	and USE	PA Resi	due Chem	istry Test	Guide	lines (c	rop group	reduction).		
Approved Use Patt	tern (GA	P)		-	Stud	y Use P	attern				
Single foliar appl. ra # of appl.: 3-4 appl. Maximum seasonal PHI: 10 days	per year	at 10-14	day inter	rvals	Singl # of a Maxi PHI: US ti Singl # of a Maxi PHI:	appl.: 3 imum ra 9-11 da rials: le foliar appl.: 3 imum ra 10 days	appl. per y tte: 1249-: tys appl. rate appl. per y tte: 1240-:	/ear 324 g a.i. : ~420 g a /ear 295 g a.i.	/ha/sea i./ha + ./ha/sea	- adjuvant	
Total Rate	РН	Decline Trials: 0, 3, 7, 10 and 14 da S-2200 Residue Levels (ppm)									
(g a.i./ha)	(days		n Min.	# Max.		AFT *	HAFT			Mean *	SD *
US trials											
1240-1295	10	1	0 0.68	3.74		0.74	3.46	1.4	0	1.56	0.83
Canadian trials		I									
1249-1324	9-11	4	0.44	1.13		0.47 1.08		0.8	0.86 0.8		0.29
Combined trials					I		1				
1240-1324	9-11	1	4 0.44	3.74		0.47	3.46	1.1	9	1.37	0.79
[#] Values based on to	otal numb	per of sa	mples; * V	alues bas	ed on	per-trial	averages	n = numl	ber of i	independen	t trials;
Crop Field Trials of										d 2377926	
Number and Locat	-	,							.,		
Region	1		2	5		7	1	11		14	Total
CDN Required*	1		0	1		0		0	-	14	16
US Required*	0		1	2		1		2		0	6
Submitted	0		0	4		2		3	-	14	23
*As per DIR98-02 a	and USE	PA Resid	due Chem	istry Test	Guide	lines (c	rop group	reduction		I	-
*				Study U			10 1		,		
Approved Use Pattern (GAP) # of appl.: 1 appl. per year Maximum foliar seasonal rate: 420 g a.i./ha PHI: 35 days			CDN trials: Maximum foliar rate: 406-442 g a.i./ha/season + adjuvant # of appl.: 3 appl. per year PHI: 31-44 days Decline Trial: 28, 33, 37 and 41 days US trials: Maximum foliar rate: 402-458 g a.i./ha/season + adjuvant # of appl.: 1 appl. per year PHI: 34-36 days Decline Trial: 26, 31, 36 and 41 days								

		PHI	S-2200 Residue Levels (ppm)												
		(days)	n	n Min. #		Max	• #	LAFT *		AFT *	Median [*]		Mean *	SD *	
US trials									•						
402-458 34-		34-36	12	<0	.02	0.13	3	< 0.02	0	.125	< 0.02		0.03	0.04	
Canadian t	rials														
406-442		31-44	8	< 0.02		0.54	4	< 0.02	0	.508	0.03		0.09	0.15	
Combined trials										•					
402-458 3		31-44	20	< 0.02		0.544	4	< 0.02	0	.508	0.02		0.07	0.12	
[#] Values based on tota		otal nur	nber of	samples; *		Values b	ased on	on per-trial a		verages; n = nun		mber of inde		pendent trials	
Crop Field	l Trials	on Fiel	d Corn	(Seed	Trea	tment)				P	MRA # 2	378	153		
Number a	nd Loca	tion of	Field T	rials		,									
Three (3) the results					rated	l rates in	n Region	5 in NA	FTA r	epresenta	tive grow	ing	regions bas	sed on	
Approved							Use Patt								
Maximum seed	seasona	g a.i./10)0 kg	a.i./100	aximum seasonal rate: 3 trials at 10 g a.i./100 kg seed and at 50 g i./100 kg seed; Samples were collected between 71-168 days after ed planting.										
Crop	Tota		РНІ				S-2200 Residue Levels (ppm					n)			
Matrix	Rate (g a.i./	e (d	lays)	n Min.		# M	Iax. #	LAFT	*	HAFT *	Median	*	Mean *	SD*	
K+CWHR		7	1-78	3	<0.01	. <	0.01	< 0.01		< 0.01	< 0.01		< 0.01		
Forage	- 11.6-1 (~50	0	8-92	3 <0.		2 <	0.02	< 0.02		< 0.02	< 0.02		< 0.02		
Grain	ain a.i./ 100 kg seed)		8-168	3 <0.0			0.01	< 0.01		< 0.01	< 0.01		< 0.01		
Stover			8-168	3 <0.02		2 <	0.02	< 0.02		< 0.02	< 0.02		< 0.02		
Residues fr were all <1 independer	LOQ. [#]	Values	based o	n total	num	ber of s	were not samples;	* Values	d as r s base	esidues fi d on per-	-trial aver	ages	s; $n = num$	on rate ber of	
Crop Field Trials on Soybeans (Seed Treatment)								PMRA # 2378152							
Number a							D .		1	1 7 (2)					
Three (3) the growing re								s 4 (1 tria	al) and	15 (2 tria	ls) in NAI	ΓA	representa	tive	
Approved	×		13 01 11		Study Use Pattern										
Maximum seasonal rate: 10 g a.i./100 kg seed							Maximum seasonal rate: 3 trials at 10 g a.i./100 kg seed and at 50 g a.i./100 kg seed; Samples were collected between 116-136 days after seed planting.								
Crop Matrix	Total R (g a.i./l		PHI (days)	S-7700 Residue Levels (1						opm)					
Soybean	21.7-29 (~50	σ	16 126	n	М	[in. [#]	Max. #	LAF	Т*	HAFT	* Medi	an	Mean *	SD *	
seed	a.i./100 seed	кg	16-136	3	<	0.01	< 0.01	<0.0)1	< 0.01	<0.0	1	<0.01		
Residues fr were all <1 independer	LOQ. [#]														
Residue Data in Field Accumulation – Spring							at, lettuce, garden beets PMRA # 2377938, 2377942								
Primary	Crop	Study Use Pattern				ı	Rotational Crop (PBIs)								
Leaf Let	1 trial in Region 10 (US) - 4 foliar applications of S-2200 on leaf lettuce for a total of 1696 g a.i./ha				JS) - of	Wheat, spinach, garden beets: (101 DAT/3.8 months) Sorghum, spinach, garden beets: (253 DAT/8.4 months) Wheat, spinach, garden beets: (356 DAT/11.8 months)									

	Total			S-2200 Residue Levels (ppm)						
Commodity	Rate (g a.i./ha)	PBI (days)	n	Min. #	Max. #	LAFT *	HAFT *	Median *	Mean *	SD *
Wheat (forage, hay, straw & grain)			1	< 0.01	< 0.01	< 0.01	< 0.01	< 0.01	< 0.01	
Spinach		101 & 356	1	< 0.01	< 0.01	< 0.01	< 0.01	< 0.01	< 0.01	
Garden beets (roots and leaves)	1.000	330	1	< 0.01	< 0.01	< 0.01	< 0.01	<0.01	< 0.01	
Sorghum (forage, stover & grain)	1696		1	< 0.01	< 0.01	< 0.01	< 0.01	<0.01	< 0.01	
Spinach		253	1	< 0.01	< 0.01	< 0.01	< 0.01	< 0.01	< 0.01	
Garden beets (roots and leaves)			1	< 0.01	< 0.01	< 0.01	< 0.01	<0.01	< 0.01	
Primary Crop		red Use pat (GAP)	ttern	Stu	ıdy Use P	attern	Rot	tational Cr	op (PBIs)	
Rapeseed	Maximum foliar rate: 420 g a.i./ha/season (ground and air) Timing: 20-50% bloon # of appl.: 1 appl./year PHI: 35 days			on rapeseed at 415 g a.i./ha				Spring wheat (286 DAT/9.5 months) Garden beets, lettuce (304 DAT/10 months)		
	Total					S-2200 Res	idue Levels (ppm)		
Commodity	Rate (g a.i./ha)	PBI (days)	n	Min. #	Max. #	LAFT *	HAFT *	Median *	Mean *	SD *
Spring Wheat (forage, hay, straw, grain)			1	<0.01	<0.01	< 0.01	<0.01	<0.01	<0.01	
Lettuce	415	286 & 304	1	< 0.01	< 0.01	< 0.01	< 0.01	< 0.01	< 0.01	
Garden beets (roots and leaves)			1	< 0.01	< 0.01	< 0.01	<0.01	< 0.01	< 0.01	
[#] Values based on		-			-		-		-	
Based on the result plant-back for labe			ılatioı	n study, a	plant-bacl	c interval of	f 4 months is	recommen	led. Imme	diate
Processed Food a	nd Feed -	Grapes					PMRA	# 2377920		
Test Site						Growing I	Region 10			
Treatment					t foliar app					
Rate					./ha (5-fol	d GAP)				
End-use product/	tormulati	on		S-2200 4	SC					
PHI (days) Processed Commo	dition			10	ogo Dogid	lues (ppm)		Duogoaciu	a Footon	
rrocesseu Commo	RAC			Aver	age Kesit 11.6			Processin	g ractor	
Grape	Juice				16.2			1.	4	
Grupe	Raising	s			22.4			1.		
Processed Food a	nd Feed -	Rapeseed					PMRA# 2	377926		
Test Site						Growing I	Region 5			
Treatment					t foliar app					
Rate					./ha (5-fol	d GAP)				
End-use product/formulation				S-2200 4	SC					
PHI (days)	1.			34				D •	T	
Processed Comme				Aver		lues (ppm)		Processin	•	
	RAC		0.23 0.02				0.09			
Rapeseed		Refined oil Meal			0.02 0.09 0.09					

Residues in Livestock Studies Residues in Livestock Studies were not conducted as the only potential livestock feed items from the petitioned uses were rapeseed meal, corn (seed treatment), and rotational crops (sorghum, and wheat), and as such, there was no expectation of quantifiable residues in meat, milk or eggs. In the absence of livestock feeding studies, the metabolism studies (laying hen, lactating goat) can confirm that no finite residues of S-2200 are expected in meat, milk, and eggs. Dietary Burden and Anticipated Residues PMRA # 2523426 Dietary burdens of 0.03 ppm (dairy cattle) and 0.008 ppm (poultry) were estimated based on potential livestock feeding items. The anticipated residues in poultry tissues and eggs were calculated to be less than 0.00004 ppm

Dietary burdens of 0.03 ppm (dairy cattle) and 0.008 ppm (poultry) were estimated based on potential livestock feeding items. The anticipated residues in poultry tissues and eggs were calculated to be less than 0.00004 ppm based on the residue-to-feed ratio from the laying hen metabolism study. The anticipated residues in livestock tissues and milk were calculated to be less than 0.00006 ppm based on the residue-to-feed ratio from the lactating goat metabolism study.

Table 9Food Residue Chemistry Overview of Metabolism Studies and Risk
Assessment

Plant Studies					
Residue Definition for Enforcement Residue Definition for Risk Assessment Primary crops (wheat, lettuce, rapeseed)	Parent (S-2200)				
Rotational crops (wheat, sorghum, lettuce, spinach, beets) Metabolic Profile in Diverse Crops	Plant metabolism studies have been conducted covering the crop categories of cereals, oilseeds and leafy vegetables (except <i>brassica</i>). A confined rotational crop study has also investigated the metabolism of residues taken up by wheat, lettuce and carrot. The route of metabolism of S-2200 has been shown to be similar, with the extent of metabolism being greater in commodities harvested at longer pre-harvest intervals. In the primary crop metabolism studies, measurement of the ratio of <i>R</i> - and <i>S</i> -isomers of S-2200 at the 2-position of the acetamide moiety.				
Animal Stu	dies				
Animals	Ruminant and Poultry				
Residue Definition for Enforcement	Not being proposed				

Residue Definition for Risk Assessment		In the absence of a livestock feeding study and an enforcement method for animal matrices, and considering that there is no expectation of measurable residues in meat, milk and eggs, based on the anticipated residues, (calculated from the metabolism studies), a residue definition for animal matrices is not being proposed. However, this will be reassessed in the event that there is an expansion of use to a livestock feed item, that contributes significantly to the dietary burden, and livestock feeding studies and enforcement methods are submitted to the Agency.			
Metabolic Profile in Animals (goat, hen)		Mandestrobin is extensively metabolized in livestock. Metabolism studies in laying hens and lactating goats showed that the metabolic pathways in livestock were similar to that found in the rat.			
Fat Soluble Residue		Yes			
Dietary Risk From Food and Water					
	Population	Estimated Risk % of Acceptable Daily Intake (ADI)			

	Population	Estimated Risk % of Acceptable Daily Intake (ADI)		
		Food Alone	Food and Water	
	All infants < 1 year	2.1	7.7	
Basic chronic non-cancer dietary	Children 1–2 years	7.1	9.2	
exposure analysis	Children 3 to 5 years	4.2	5.9	
ADI = 0.3 mg/kg bw/day	Children 6–12 years	1.7	3.0	
Estimated chronic drinking water	Youth 13–19 years	0.7	1.8	
concentration = $225 \ \mu g/L$	Adults 20–49 years	0.8	2.3	
	Adults 50+ years	1.0	2.5	
	Females 13-49 years	0.9	2.4	
	Total population	1.3	2.8	

Table 10Fate and Behaviour in the Environment

Property	Test substance	Value ¹	Transformation products	Comments	Reference
		Abiotic transformat	ion		•
Hydrolysis at 50°C	[Benzyl- ¹⁴ C]-S-2200 <i>R</i> -isomer, [Benzyl- ¹⁴ C]-S-2200 <i>S</i> -isomer	pH 4: stable pH 7: stable pH 9: stable	none	Not a route of transformation in the environment	2377861 2377863
Phototransformation on soil	[Benzyl- ¹⁴ C]-S-2200 <i>R</i> -isomer, [Benzyl- ¹⁴ C]-S-2200 <i>S</i> -isomer and [Phenoxy- ¹⁴ C]-S-	$DT_{50} (irradiated) = 52.3-63.8 d;DT_{90} (irradiated) =173.7-211.8 d;DT_{50} (dark) = 71.7-82.9$	<u>Major:</u> UR* <u>Minor</u> : DX-CA-S-2200; De-Xy-S-2200; 2-COOH-S-2200; 5-	Not a route of transformation in the environment; no	2377867 2377864

Property	Test substance	Value ¹	Transformation	Comments	Reference
	2200 <i>R</i> -isomer	d;	productsCOOH-S-2200;	isomerization	
	2200 K-Isoliki	$DT_{90} (dark) = 238.1-$ 275.4 d (SFO)	MCBX; S-2200-OR; CO ₂	occurred	
Phototransformation in water	[Benzyl- ¹⁴ C]-S-2200 and	$\frac{\text{S-2200 } R\text{-isomer}}{\text{DT}_{50}=4.39 \text{ d};}$	Minor (Benzyl label only:	Non-persistent	2377871
25±1°C	[Phenoxy- ¹⁴ C]-S- 2200 <i>R</i> -isomer (combined labels)	DT ₉₀ =14.6 d (SFO) (combined labels) Dark: none	De-Xy-S-2200; CO2 Major: S-2200-OR; S-2200- ORC; 2200-PR and	Phototransfor mation is expected to be an important	
		$\frac{S-2200-OR}{DT_{50}=9.25 \text{ d};}$ DT ₉₀ =30.7 d (SFO)	CO_2 . (CO_2 for phenoxy label, only)	route of dissipation	
	[Benzyl- ¹⁴ C]-S-2200 S-isomer	Irradiated: <u>S-2200 S-isomer</u> DT_{50} =4.59 d;	<u>Major:</u> S-2200-OR; S-2200- ORC; De-Xy-S-2200	Non-persistent Phototransfor	2377869
		DT ₉₀ =15.3 d (SFO) Dark: none <u>S-2200-OR</u>	<u>Minor</u> : S-2200-PR; CO ₂	mation is expected to be an important route of	
	Mandestrobin	DT ₅₀ = 8.23 d; DT ₉₀ =27.3 d (SFO)		dissipation	
Phototransfor- mation in air	Mandestrobin	Mandestrobin is not exp conditions based on va constant.			
		Biotransformation	n	· · · · · · · · · · · · · · · · · · ·	·
Biotransformation in aerobic soil	[Benzyl- ¹⁴ C]-S-2200 <i>R</i> -isomer,	<u>California Sand</u> DT ₅₀ : 295 d; DT ₉₀ : 980 d (SFO)	<u>Major</u> : 5-COOH-S-2200; CO ₂ ; UR <u>Minor</u> : DX-CA-S-2200; 2- COOH-S-2200; 2- CONH ₂ -S-2200; 5-CONH ₂ -S-2200; MCBX; De-Xy-S- 2200	Biotransforma tion in aerobic soil is a main route of dissipation for mandestrobin	2377894
		$\frac{\text{Mississippi Silt loam}}{\text{DT}_{50}: 393 \text{ d};}$ $\text{DT}_{90}: 1867 \text{ d} (\text{DFOP})$ $\text{Slow } t_{1/2} = 635 \text{ d}$	<u>Major:</u> CO ₂ ; UR <u>Minor:</u> DX-CA-S-2200; 2-COOH-S-2200; 5-COOH-S-2200; 2-CONH ₂ -S-2200; 5-CONH ₂ -S-2200; MCBX; De-Xy-S- 2200	Mandestrobin is moderately persistent to persistent	
		$\frac{\text{North Dakota Sandy}}{\text{DT}_{50}: 334 \text{ d};}$ $\text{DT}_{90}: 6785 \text{ d} (\text{IORE})$ $t_{\text{R IORE}} = 2040 \text{ d}$	<u>Major:</u> 5-COOH-S-2200; CO ₂ <u>Minor:</u> DX-CA-S-2200; 2- COOH-S-2200; 2- CONH ₂ -S-2200; 5-CONH ₂ -S-2200; MCBX; De-Xy-S- 2200		
	[Benzyl- ¹⁴ C]-S-2200 <i>R</i> -isomer	$\frac{\text{Sandy loam (Speyer}}{5M)}$ $DT_{50}: 66.5 \text{ d};$	<u>Major</u> : 5-COOH-S-2200; CO ₂ ; UR	Mandestrobin is moderately persistent to	2377881

Property	Test substance	Value ¹	Transformation products	Comments	Reference
		DT ₉₀ : 221 d (SFO)	<u>Minor</u> : DX-CA-S-2200; MCBX; 2-COOH-S- 2200; 2,5-DMP	persistent	
		<u>Loamy sand (Speyer</u> <u>2.2)</u> DT_{50} : 238 d; DT_{90} : 920 d (DFOP) Slow $t_{1/2}$ = 294 d	<u>Major:</u> none <u>Minor</u> : 5-COOH-S-2200; 2-COOH-S-2200; MCBX; DX-CA-S- 2200;CO ₂		
		$\frac{\text{Clay loam (SK920191)}}{\text{DT}_{50}: 43.3 \text{ d};}$ DT ₉₀ : 205 d (DFOP) Slow t _{1/2} = 70 d	<u>Major:</u> 5-COOH-S-2200; CO ₂ ; UR <u>Minor:</u> 2-COOH-S-2200; MCBX; De-Xy-S- 2200;DX-CA-S-2200		
		$\frac{\text{Silt loam (Chelmorton)}}{\text{DT}_{50}: 96.5 \text{ d};}$ DT ₉₀ : 631 d (DFOP) Slow t _{1/2} = 230 d	<u>Major</u> : CO ₂ ; UR <u>Minor</u> : 5-COOH-S-2200; 2-COOH-S-2200; MCBX; DX-CA-S- 2200		
	[Benzyl- ¹⁴ C]-S-2200 <i>R</i> -isomer	$\frac{\text{Loam (Aschard)}}{\text{DT}_{50}: 43.9 \text{ d};}$ $\text{DT}_{90}: 213 \text{ d} (\text{DFOP})$ $\text{Slow } t_{1/2} = 77.4 \text{ d}$	<u>Major</u> : 5-COOH-S-2200;	Slightly persistent	2377886
		$\frac{\text{Silty clay loam}}{(\text{Monteil})}$ $DT_{50}: 37.5 \text{ d};$ $DT_{90}: 152 \text{ d} (\text{IORE})$ $t_{\text{R IORE}} = 45.8 \text{ d}$	CO ₂ ; UR <u>Minor</u> : 2-COOH-S-2200; MCBX; De-Xy-S- 2200;DX-CA-S-220		
	[Benzyl- ¹⁴ C]- and [Phenoxy- ¹⁴ C]- S- 2200 <i>R</i> -isomer	$\frac{L_{ORE} = 43.8 \text{ d}}{L_{ORE} = 43.8 \text{ d}}$ $\frac{L_{Oam} (\text{New Jersey})}{\text{DT}_{50}: 110 \text{ d};}$ $DT_{90}: 930 \text{ d} (\text{DFOP})$ Slow $t_{1/2} = 397 \text{ d}$ (combined labels)	Major: DX-CA-S-2200; 5-COOH-S-2200; CONH ₂ -S-2200; 5-CONH ₂ -S-2200; CO ₂ ; UR Minor: 2-COOH-S-2200; MCBX; De-Xy-S-2200;	Mandestrobin is moderately persistent	2377890
	[Benzyl- ¹⁴ C]-S-2200 S-isomer	$\frac{\text{Loam (New Jersey)}}{\text{DT}_{50}: 118 \text{ d};}$ DT ₉₀ : 1494 d (IORE) $t_{\text{R IORE}} = 450 \text{ d}$	Major: DX-CA-S-2200; 2-CONH ₂ -S-2200; 5-CONH ₂ -S-2200; CO ₂ ; UR Minor: 2-COOH-S-2200; 5-COOH-S-2200; MCBX; De-Xy-S-2200		

Property	Test substance	Value ¹	Transformation products	Comments	Reference
	[Benzyl- ¹⁴ C]-S-2200 S-isomer	<u>Sandy loam (Speyer</u> <u>5M)</u> DT ₅₀ : 85.4 d; DT ₉₀ : 284 d (SFO)	Major: 5-COOH-S-2200; CO ₂ ; UR Minor: 2-COOH-S-2200; MCBX; De-Xy-S- 2200;DX-CA-S-220	Mandestrobin is moderately persistent to persistent	2377883
		<u>Loamy sand (Speyer</u> <u>2.2)</u> DT_{50} : 22305 d; DT_{90} : 6599792944837 d (IORE) $t_{R IORE} = 1.99e+12$ d	<u>Major:</u> none <u>Minor</u> : 5-COOH-S-2200; 2-COOH-S-2200; MCBX; DX-CA-S- 2200; CO ₂		
		$\frac{\text{Clay loam (SK920191)}}{\text{DT}_{50}: 90.2 \text{ d};}$ $\text{DT}_{90}: 367 \text{ d (DFOP)}$ $\text{Slow } t_{1/2} = 119 \text{ d}$	<u>Major:</u> 5-COOH-S-2200; CO ₂ ; UR <u>Minor:</u> 2-COOH-S-2200; MCBX		
		$\frac{\text{Silt loam (Chelmorton)})}{\text{DT}_{50}: 124 \text{ d};}$ $\text{DT}_{90}: 730 \text{ d} (\text{DFOP})$ Slow $t_{1/2} = 261 \text{ d}$	<u>Major:</u> CO ₂ ; UR <u>Minor:</u> 5-COOH-S-2200; 2-COOH-S-2200; MCBX ;DX-CA-S- 2200		
	[Benzyl- ¹⁴ C]-S-2200 S-isomer	$\frac{\text{Loam (Aschard)}}{\text{DT}_{50}: 72.2 \text{ d};}$ DT ₉₀ : 266 d (DFOP) Slow t _{1/2} = 83.3 d	<u>Major:</u> 5-COOH-S-2200; CO ₂ ; UR <u>Minor:</u> 2-COOH-S-2200; MCBX; De-Xy-S- 2200; DX-CA-S-2200	Moderately persistent	2377888
		$\frac{\text{Silty clay loam}}{\text{(Monteil)}}$ $DT_{50}: 56.5 \text{ d};$ $DT_{90}: 269 \text{ d} \text{(IORE)}$ $t_{R \text{ IORE}} = 81 \text{ d}$	<u>Major:</u> CO ₂ ; UR <u>Minor:</u> 5-COOH-S-2200; 2-COOH-S-2200; MCBX; De-Xy-S- 2200; DX-CA-S-2200		
	[Benzyl- ¹⁴ C]5- COOH-S-2200	$\frac{\text{Silt loam (SK104691)}}{\text{DT}_{50}: 20.3 \text{ d};}$ $\text{DT}_{90}: 89.7 \text{ d (IORE)}$ $t_{\text{R IORE}} = 27 \text{ d}$	<u>Major:</u> CO ₂ ; UR <u>Minor</u> : DX-CA-S-2200	Slightly to moderately persistent	2377879
		$\frac{\text{Clay loam (SK920191)}}{\text{DT}_{50}: 24.6 \text{ d};}$ $\text{DT}_{90}: 165 \text{ d (IORE)}$ $t_{\text{R IORE}} = 49.8 \text{ d}$	<u>Major:</u> CO ₂ ; UR <u>Minor</u> : none <u>Major</u> :		
		$\frac{\text{Sandy loam (Speyer}}{5M)} \\ DT_{50}: 39.1 \text{ d}; \\ DT_{90}: 161 \text{ d (IORE)} \\ t_{R \text{ IORE}} = 48.5 \text{ d}$	CO ₂ ; UR <u>Minor</u> : none		

Property	Test substance	Value ¹	Transformation products	Comments	Reference
	[Benzyl- ¹⁴ C]2- COOH-S-2200	<u>Silt loam (SK104691)</u> DT ₅₀ : 18.1 d; DT ₉₀ : 60 d (SFO)	<u>Major:</u> CO ₂ ; UR <u>Minor:</u> DX-CA-S-2200	Slightly to moderately persistent	2377877
		$\frac{\text{Clay loam (SK920191)}}{\text{DT}_{50}: 19.7 \text{ d};}$ $\text{DT}_{90}: 73 \text{ d (IORE)}$ $t_{\text{R IORE}} = 22 \text{ d}$	<u>Major</u> : CO ₂ ; UR <u>Minor</u> : DX-CA-S-2200		
		$\frac{\text{Sandy loam (Speyer}}{5M)}{\text{DT}_{50}: 27.2 \text{ d};}$ $\text{DT}_{90}: 76.9 \text{ d (IORE)}$ $t_{\text{R IORE}} = 23.2 \text{ d}$	<u>Major:</u> CO ₂ ; UR <u>Minor:</u> DX-CA-S-2200 De-Xy-S-2200		
Biotransformation in anaerobic soil	[Benzyl- ¹⁴ C]- and [Phenoxy- ¹⁴ C]- S- 2200 <i>R</i> -isomer	Water: loam soilsystem (New Jersey) DT_{50} : 3801 d; DT_{90} : 12625 d (SFO)(combined labels)	<u>Major:</u> UR (New Jersey) <u>Minor</u> : DX-CA-S-2200; 2- COOH-S-2200; 5-	Not a route of dissipation for mandestrobin	2377897
	[Benzyl- ¹⁴ C] S-2200 <i>R</i> -isomer	DT ₅₀ : 1795 d; DT ₉₀ : 5963 d (SFO)	COOH-S-2200; De- Xy-S-2200; MCBX; CO ₂		
	[Benzyl- ¹⁴ C] S-2200 (1:1, <i>R:S</i> isomer ratio)	Water: loamy sand soilsystem (California) DT_{50} : 12,772 d; DT_{90} : 42426 d (SFO)Water: sandy loam soil	<u>Major:</u> none (California); Unknown A (North Dakota and Mississippi)	Not a route of dissipation for mandestrobin	2377900
		$\frac{\text{system (North Dakota)}}{\text{DT}_{50}: 861 \text{ d};}$ $\text{DT}_{90}: 2860 \text{ d} (\text{SFO})$ $\frac{\text{Water: silt loam soil}}{\text{Water: silt loam soil}}$	<u>Minor</u> : 5-COOH-S-2200; 2- COOH-S-2200; DX- CA-S-2200; MCBX; De-Xy-S-2200; CO ₂		
		<u>system (Mississippi)</u> DT ₅₀ : 1,723 d; DT ₉₀ : 5722 d (SFO)	<i>De Ny 5 2200, CO</i> ₂		
Biotransformation in aerobic water- sediment systems (20±2°C)	[Benzyl- ¹⁴ C]-S-2200 <i>R</i> -isomer; [Phenoxy- ¹⁴ C]-S- 2200 <i>R</i> -isomer (combined labels)	<u>Lake water:silt loam</u> <u>sediment</u> $DT_{50}=322 \text{ d}$ $DT_{90}=1069 \text{ d} \text{ (SFO)}$	$\begin{tabular}{l} \hline \underline{Major:} \\ 5-COOH-S-2200 \\ \hline \underline{Minor:} \\ 2-COOH-S-2200, \\ \hline MCBX; CO_2 \end{tabular}$	Partition: 77.9% (at 62 d) and 74.1% in sediment at end of study	2377903
		Lake water:loamy sand sediment DT_{50} : 781 d; DT_{90} : 2803 d (DFOP) Slow $t_{1/2} = 870$	<u>Major</u> : none <u>Minor:</u> 5-COOH-S-2200; 2- COOH-S-2200, MCBX; CO ₂	Partition: 67.8% in sediment at end of study Mandestrobin	
	[Benzyl- ¹⁴ C]-S-2200 S-isomer	$\label{eq:calwich_Abbey_Lake} \hline \\ \hline \\ \underline{Calwich_Abbey_Lake}\\ water: silt loam}\\ \underline{sediment}\\ DT_{50}{=}161 \ d\\ DT_{90}{=}535 \ d \ (SFO) \\ \hline \end{array}$	Major: MCBX; UR* <u>Minor</u> : 5-COOH-S-2200; CO ₂	is persistent Partition: 69.6% (at 29 d) and 58.2% in sediment at end of study	2377905
		Swiss Lake water:	<u>Major</u> : none		

Property	Test substance	Value ¹	Transformation	Comments	Reference
		<u>loamy sand sediment</u> DT ₅₀ : 733 d; DT ₉₀ : 2435 d (SFO)	products Minor: 5-COOH-S-2200; 2- COOH-S-2200, MCBX; CO2	Partition: 64.0% in sediment at end of study Mandestrobin is moderately persistent to	
Biotransformation in anaerobic water- sediment (25°C)	Benzyl- ¹⁴ C]-S-2200 <i>R</i> -isomer + [Phenoxy- ¹⁴ C]-S- 2200 <i>R</i> -isomer (combined) [Benzyl- ¹⁴ C]-S-2200 <i>S</i> -isomer	Water:clay sediment Bosket Lake DT ₅₀ : 2,917 d; DT ₉₀ : 9691 d (SFO) (combined label)	R-isomer Major: UR Minor: MCBX; 5-COOH-S-2200; 2,5-DMP; 2-COOH-S-2200; DX-CA-S-2200; DX-CA-S-2200; CO2	persistent Partition: 87.09 % in sediment at end of study	2377907
		DT ₅₀ : 458 d; DT ₉₀ : 1523 d (SFO)	S-isomer <u>Major</u> : MCBX <u>Minor</u> : 5-COOH-S-2200; 2-COOH-S-2200; DC- Xy-S-2200; DPMBA; CO ₂	Partition: 68.47% (at 120 d) and 50.77 % in sediment at end of study	
		$\frac{\text{Water:sand sediment}}{\text{Golden Lake}}$ $DT_{50}: 8,822 \text{ d};$ $DT_{90}: 29307 \text{ d (SFO)}$ (combined label) $DT_{50}: 859 \text{ d};$ $DT_{90}: 2854 \text{ d (SFO)}$	<i>R</i> -isomer <u>Major</u> : none <u>Minor</u> : MCBX; 5-COOH-S- 2200; 2,5-DMP; 2-COOH- S-2200; De-Xy-S-2200; CO ₂	Partition: 88.39 % in sediment at end of study	
			S-isomer Major: MCBX Minor: 5-COOH-S-2200; 2-COOH-S-2200; DPMBA; CO2	Partition: 69.58 % in sediment at end of study Mandestrobin is persistent	
Mobility	[Dammel ¹⁴ C] 9 2000	$V = 2.05 \pm 19.2 (1 / 1_{})^{11}$	<u>∖-l/n</u>	Low to	2277010
Adsorption / desorption in soil (5+2 soils)	[Benzyl- ¹⁴ C] S-2200 (1:1, <i>R</i> : <i>S</i> isomer ratio)	$\begin{split} K_F &= 2.05\text{-}18.2 \text{ (L/kg-soil}\\ K_{FOC} &= 287\text{-}1104 \text{ (L/kg-C)}\\ 1/n &= 0.882\text{-}0.962 \end{split}$		Low to moderately mobile	2377910 2377919
(3 + 3 soils)	[Benzyl- ¹⁴ C]-2- COOH-S-2200	$\begin{split} K_F &= 0.27\text{-}2.94 \text{ (L/kg-soil)}^{-l/n} \\ K_{FOC} &= 6\text{-}226 \text{ (L/kg-OC)}^{-l/n} \\ 1/n &= 0.850\text{-}0.922 \end{split}$		Moderate to very high mobility	2377912 2377915
(3 + 3 soils)	[Benzyl- ¹⁴ C]-5- COOH-S-2200;	$\begin{split} K_F &= 1.26\text{-}8.89 \ (L/kg\text{-soil})^{\text{-l/n}} \\ K_{FOC} &= 29\text{-}684 \ (L/kg\text{-OC})^{\text{-l/n}} \\ 1/n &= 0.853\text{-}1.038 \end{split}$		Low to very high mobility	2377914 2377916
Adsoption (HPLC)	S-2200	K _{OC} = 1780		<u>Mobilty:</u> Low	2377918

Property	Test substance	Value ¹	Transformation products	Comments	Reference
	2-COOH-S-2200 5-COOH-S-2200 2-CONH ₂ -S-2200 5-CONH ₂ -S-2200			Very high Very high High Moderate	
4-years lysimeter study (2 lysimeters)	Dx-CA-S-2200 [Benzyl- ¹⁴ C]-S-2200 25SC	K _{OC} < 18 Silty sand soil: 0.26% and 0.43% AR recovered in leachate	De-Xy-S-2200; DX-CA-S-2200; 2-COOH-S-2200; 5-COOH-S-2200; MCBX	Very high Low leaching / mobility potential	2377921
Volatilization	Not required based or $(6.5 \times 10^{-12} \text{ atm m}^3/\text{m})$	n the low vapour pressure nol).		C) and Henry's	law constan
Field studies		,			
Saskatchewan Bare soil	S-2200 4SC	$\label{eq:loss} \begin{array}{ c c c } \underline{Loam}: \\ DT_{50} & 1.04 & d; \\ DT_{90}: & 367 & d & (DFOP) \\ Slow & t_{1/2} = 163 & d \end{array}$	Minor: DX-CA-S-2200; 2-COOH S-2200 5-COOH S-2200	No residues beyond 30 cm soil depth Mandestrobin is non- persistent in soil	2377953
North Dakota Bare ground	S-2200 4SC	$\label{eq:loss} \begin{array}{ c c c } \hline Loam: & \\ DT_{50}: 4.89 \text{ d}; \\ DT_{90}: 2015 \text{ d} (IORE) \\ t_{R \ IORE} = 606 \text{ d or } 142 \\ (excluding \ outlier \ at \\ 303 \text{ d}) \end{array}$	Minor: DX-CA-S-2200; 2-COOH S-2200 5-COOH S-2200	No residues beyond 15 cm soil depth. Mandestrobin is non- persistent in soil	2377968
Ontario, Canada Bare ground	S-2200 4SC	$\frac{\text{Sandy loam:}}{\text{DT}_{50}=83.9 \text{ d};} \\ \text{DT}_{90}: 486 \text{ d} \text{ (DFOP)} \\ \text{Slow } t_{1/2} = 173 \text{ d}$	Minor: DX-CA-S-2200; 2-COOH S-2200 5-COOH S-2200	No residues beyond 15 cm soil depth soil Mandestrobin is moderately persistent in soil	2495782
Field dissipation Established Turfgrass in Ontario, Canada	S-2200 4SC	$\frac{\text{Turf/Thatch:}}{\text{DT}_{50}: 8.43 \text{ d};}$ $\text{DT}_{90}: 28 \text{ d} \text{ (SFO)}$ $\frac{\text{Sandy loam soil:}}{\text{DT}_{50}: 43.3 \text{ d};}$ $\text{DT}_{90}: 380 \text{ d} \text{ (IORE)}$ $t_{\text{R IORE}} = 114 \text{ d}$	Minor: DeXy- S-2200; Minor: 2-COOH S-2200 5-COOH S-2200	No residues beyond 30 cm soil depth. Mandestrobin is non- persistent to slightly persistent	2377963
Bioconcentration/bioa	ccumulation			Francis	
Bioconcentration and Metabolism with Bluegill Sunfish (Lepomis macrochirus)	[Benzyl- ¹⁴ C]S-2200	Whole body steady state Bioconcentration factor = 25-26 Kinetics for S-2200 was not calculated	Transformation products formed by hydroxylation then conjugated with sulfate and glucuronic acid	depuration half-life for the total radioactive residues ~2 days	2378049

~ -			<i>a.</i> -			References
Code	Chemical name	Chemical structure	Study	max %AR (day)	%AR at Study End (study length)	
			PARENT			
S-2200	(RS)-2- methoxy-N- methyl-2- $[\alpha$ - (2,5-xylyloxy)- o- tolyl]acetamide	↓ ↓ ↓ ↓ ↓ ↓ ↓				
		MAJOR (>10%) TRA	NSFORMATION I	PRODUCTS		
DX-CA- S-2200		HO O O H	Aerobic soil Anaerobic soil Soil photolysis Aqueous photolysis Hydrolysis	11.8 (131) 6.2 (266) 6.7 (17)	8.4 (362) 6 (363) 6.3 (30) -	2377890 2377897 2377864 -
		\sim	Aerobic aquatic Anaerobic aquatic Field studies	- 0.37 (120) 3.3 (59)	- 0.16 (357) 0 (420)	- - 2377907 2377953
5- COOH- S-2200	(<i>RS</i>)-3-{2-[1- methoxy-1-(<i>N</i> - methylcarbamo yl) methyl]benzylo xy}-4- methylbenzoic acid		Aerobic soil Anaerobic soil Soil photolysis Aqueous photolysis Hydrolysis Aerobic aquatic Anaerobic aquatic Field studies	18 (59) 8.4 (63) - - 11.9 (62) 1.34 (240) 6.8 (28)	12.6 (120) 7.9 (363) - - 2 (101) 0 (365) 0 (656)	2377881 2377897 - - 2377903 2377907 2377963
MCBX	(<i>RS</i>)-2-hydroxy- <i>N</i> -methyl-2-[α- (2,5- xylyloxy)- <i>o</i> - tolyl]acetamide	OH HI	Aerobic soil Anaerobic soil Soil photolysis Aqueous photolysis Hydrolysis Aerobic aquatic Anaerobic aquatic Field studies	2.8 (15) 7.5 (314) 0.9 (17) - 18.1 (102) 32.04 (357)	0.7 (364) 7.5 (314) 0.6 (30) - 18.1 (102) 32.04 (357)	2377894 2377890 2377864 - - 2377905 2377907
S-2200- OR	(<i>RS</i>)-2-[2-(2- hydroxy-3,6- dimethylbenzyl) phenyl]-2- methoxy- <i>N</i> - methylacetamid e		Aerobic soil Anaerobic soil Soil photolysis Aqueous photolysis Hydrolysis Aerobic aquatic Anaerobic aquatic Field studies	- 1.8 (30) 24.5 (7) - -	- 1.8 (30) 8.5 (30) - - -	- 2377864 2377871 - - -
S-2200-	(<i>RS</i>)- <i>N</i> ,1,4-		Aerobic soil	-	-	-

Table 11 Major Transformation Products Formed in the Environment
--

Code	Chemical name	Chemical structure	Study	max %AR (day)	%AR at Study End (study length)	References	
ORC	trimethyl-6,11-		Anaerobic soil	-	-	-	
	dihydrodibenzo [b,e]oxepine-6-	\rightarrow	Soil photolysis	-	-	-	
	carboxamide	ί μ Ι Ι Ν	Aqueous photolysis	19.8 (14)	12.8 (30)	2377871	
			Hydrolysis	-	-	-	
		l l	Aerobic aquatic	-	-	-	
			Anaerobic aquatic	-	-	-	
			Field studies				
		HO.	Aerobic soil	-	-	-	
S-2200-	(<i>RS</i>)-2-[2-(4-		Anaerobic soil	-	-	-	
PR	hydroxy-2,5- dimethylbenzyl)	ОНН	Soil photolysis	-	-	-	
	phenyl]-2-		Aqueous photolysis	10.2 (4)	1.7 (30)	2377871	
	methoxy-N-	8	Hydrolysis	-	-	-	
	methylacetamid e		Aerobic aquatic	-	-	-	
			Anaerobic aquatic	-	-	-	
			Field studies	-	-	-	
	2-({2-[1-		Aerobic soil	14.1 (362)	14.1 (362)	2377890	
2- CONH2-	methoxy-2- (methylamino)-		Anaerobic soil	-	-	-	
S-2200			Soil photolysis	-	-	-	
5				Aqueous photolysis	-	-	-
			Hydrolysis	-	-	-	
		HN	Aerobic aquatic	-	-	-	
		∽ «н,	Anaerobic aquatic	-	-	-	
			Field studies				
5-	3-({2-[1-		Aerobic soil	13.7 (362)	13.7 (362)	2377890	
CONH ₂ - S-2200	methoxy-2-	°	Anaerobic soil	-	-	-	
5-2200	(methylamino)- 2-	nino)-	Soil photolysis	-	-	-	
	oxoethyl]benzyl	н,с	Aqueous photolysis	-	-	-	
	}oxy)-4-	о сн,	Hydrolysis	-	-	-	
	methylbenzamid		Aerobic aquatic	-	-	-	
	e		Anaerobic aquatic	-	-	-	
		Сн.	Field studies	-	-	-	
De-Xy-S-	(<i>RS</i>)-2-(2-	10	Aerobic soil	3 (0)	0 (362)	2377890	
2200	hydroxymethylp henyl)-2-	HO O	Anaerobic soil	3.6 (48)	0 (363)	2377897	
	henyl)-2- methoxy- <i>N</i> - methylacetamid		Soil photolysis	3.3 (30)	3.3 (30)	237864	
			Aqueous photolysis	9.6 (30)	9.6 (30)	2377869	
	e		Hydrolysis	-	-	-	
			Aerobic aquatic	-	-	-	
			Anaerobic aquatic	3.73 (7)	0.58 (357)	2377907	
			Field studies	-	-	-	

Organism Exposure **Test substance Endpoint value Degree of** Reference toxicity¹ Invertebrates 14-d Acute S-2200 TG $LC_{50} = 168 \text{ mg a.i./kg}$ 2378014 Earthworm, n/a soil dw Eisenia fetida 2-COOH-S-14-d LC₅₀>1000 mg n/a 2378015 /kg soil dw (highest 2200 concentration tested) 5-COOH-S- $LC_{50}>1000 \text{ mg/kg}$ 2378016 n/a 2200 soil dw (highest concentration tested) 56-d Chronic S-2200 TG NOEC = 7.5 mg2378021 n/a a.i./kg soil dw LD₅₀ >110.71 µg Honeybee, Apis S-2200 TG Practically non-2378017 Acute oral mellifera a.i./bee (highest dose toxic tested) 2378017 Acute contact S-2200 TG LD₅₀ >100 µg a.i./bee Practically non-(highest dose tested) toxic S-2200 TG LD₅₀ >100 µg 2378018 Honeybee, Apis Acute n/a a.i./larva (highest mellifera Larva dose tested) LR₅₀ > 1000 g a.i./ha Predatory mite, 7-d Contact, S-2200 25SC 2378019 n/a **Typhlodromus** ER_{50} (reproduction) > Glass plates pyri Scheuten 1000 g a.i./ha (highest rate tested) Parasitoid, 48h-Contact, S-2200 25SC $48-h LR_{50} > 1000 g$ n/a 2378020 Aphidius Glass plates a.i./ha (highest rate tested) rhopalosiphi 48 h-ER₅₀ (reproduction) =757.2 g a.i./ha Birds S-2200 TG Bobwhite quail, 14-d LD₅₀ >2250 mg Practically non-2378050 Acute Oral Colinus a.i./kg bw (highest toxic concentration tested) virginianus Canary, S-2200 TG 14-d LD₅₀ >1000 mg non-toxic at the 2378051 a.i./kg bw highest dose Serinus canaria with no regurgitation Bobwhite quail, S-2200 TG 8-d LC₅₀ >5620 mg Practically non-2378052 5-d Dietary a.i./kg diet (>1136 mg Colinus toxic a.i./kg bw/day) virginianus (highest concentration tested) 8-d LC₅₀ >5620 Mallard duck, S-2200 TG Practically non-2378053 Anas a.i./kg diet toxic platyrhynchos (>2460 mg a.i./kg bw/day) (highest concentration tested) Bobwhite quail 21-w S-2200 TG 21 -w NOEC = 10002378055 n/a Colinus Reproduction mg a.i./kg diet (91.1 virginianus mg a.i./kg bw/day)

Table 12Toxicity of Mandestrobin and Transformation Products to Non-Target
Terrestrial Species

Organism	Exposure	Test substance	Endpoint value	Degree of toxicity ¹	Reference
Mallard duck,	20-w	S-2200 TG	20-w NOEC = 1000	n/a	2378057
Anas	Reproduction		mg a.i./kg diet (129.1		
platyrhynchos			mg a.i./kg bw/day)		
Mammals					
Rat	Acute	S-2200 TG	LD ₅₀ >2000 mg	Practically non-	2377929
			a.i./kg bw (highest	toxic	
			concentration tested)		
	2-generation	S-2200 TG	NOAEL (offspring) =	n/a	2377964
	reproduction		3000 ppm (166/195		2525908
	-		mg/kg bw/d $\sqrt[3]{\uparrow}$		
Vascular plants					
Four monocots:	Seedling	S-2200 4 SC	EC ₂₅ >560 g a.i./ha	n/a	2378070
onion, ryegrass,	Emergence		NOEC = 560 g a.i./ha		
wheat and corn.	-		(highest rate tested)		
Six dicots:	Vegetative Vigor	S-2200 4 SC	EC ₂₅ >560 g a.i./ha	n/a	2378071
sugarbeet, oilseed			NOEC = 560 g a.i./ha		
rape, cabbage,			(highest rate tested)		
soybean, lettuce					
and tomato.					
¹ Atkins et al. (1981) for	bees and USEPA classification	ation for others, where a	pplicable; n/a, not applicable		

Table 13Screening Level and Refined Risk Assessment of Mandestrobin for Non-
Target Species, Other than Birds and Mammals

Organism	Exposure	Endpoint Value	EEC	RQ	Level of Concern
Invertebrates					
Earthworm	Acute	LC ₅₀ /2: 84 mg a.i./kg soil	0.844 mg a.i./kg soil	0.01	Not exceeded
Bee	Contact	LD ₅₀ : > 100 µg a.e./bee	0. 472 kg a.i/ha \times 2.4 µg a.i./bee per kg/ha = 1.13 µg a.i./bee	< 0.01	Not exceeded
	Oral	LD ₅₀ : > 110.71 µg a.i./bee	0. 472 kg a.i./ha \times 29 µg a.i./bee per kg/ha = 13.7 µg a.i./bee	< 0.1	Not exceeded
	Larva acute	LD ₅₀ : > 100 µg a.i./larva	0. 472 kg a.i./ha × 29 µg a.i./bee per kg/ha = 13.7 µg a.i./larva	<0.1	Not exceeded
Predatory arthropod, <i>Typhlodromus</i> <i>pyri</i>	Contact, glass plate	LR ₅₀ : > 1000 g a.i./ha	In-field: 472 g a.i./ha	In-field:	Not exceeded
Parasitoid arthropod, Aphidius rhopalosiphi	Contact, glass plate	LR ₅₀ : > 1000 g a.i./ha	Cumulative rate of 744.4 g a.i./ha	< 0.7	

Organism	Exposure	Endpoint Value	EEC	RQ	Level of Concern
Vascular plants		, and			concern
Vascular plant	Seedling emergence; Vegetative vigour	ER ₂₅ : >560 g a.i./ha	In-field: 472 g a.i./ha Cumulative rate of 744.4 g a.i./ha Off-field (aerial appl.,	In-field: <1.3 Off-field	Exceeded ¹ Not exceeded
			17% drift): 80.24 g a.i./ha; cumulative rate of 126.55 g a.i./ha	(aerial): <0.2	

Table 14Screening Level Risk Assessment of Foliar Application of Mandestrobin for
Birds and Mammals

	Toxicity (mg a.i./kg bw/d)	Feeding Guild (food item)	EDE* (mg a.i./kg bw)	RQ	Level of Concern				
Small Bird (0.02 kg	g)								
Acute	>100.00	Insectivore	60.59	<0.6	Not exceeded				
Reproduction	91.10	Insectivore	60.59	0.7	Not exceeded				
Medium Sized Bird (0.1 kg)									
Acute	>100.00	Insectivore	47.28	< 0.5	Not exceeded				
Reproduction	91.10	Insectivore	47.28	0.5	Not exceeded				
Large Sized Bird (1 kg)								
Acute	>100.00	Herbivore (short grass)	30.54	<0.3	Not exceeded				
Reproduction	91.10	Herbivore (short grass)	30.54	0.3	Not exceeded				
Small Mammal (0.	015 kg)								
Acute	>200.00	Insectivore	34.85	< 0.2	Not exceeded				
Reproduction	166	Insectivore	34.85	0.2	Not exceeded				
Medium Sized Ma	mmal (0.035 kg)								
Acute	>200.00	Herbivore (short grass)	67.59	< 0.3	Not exceeded				
Reproduction	166	Herbivore (short grass)	67.59	0.4	Not exceeded				
Large Sized Mam	nal (1 kg)				-				
Acute	>200.00	Herbivore (short grass)	36.12	< 0.2	Not exceeded				
Reproduction	166	Herbivore (short grass)	36.12	0.2	Not exceeded				
FIR: Food Ingestion Rate birds with body weight g Passerine Equation (body All birds Equation (body For mammals, the "all m bw: Generic Body Weigl	e. For generic birds wi greater than 200 g, the y weight <or 200="" =="" g):<br="">y weight >200 g): FIR ammals" equation was ht</or>	ted using the following formula: (F th body weight less than or equal to "all birds" equation was used: FIR (g dry weight/day) = 0.398 (b (g dry weight/day) = 0.648 (bw in g s used: FIR (g dry weight/day) = 0.	o 200 g, the "passerine w in g) ^{0.850} g) 0.651 235 (bw in g) ^{0.822}	" equation was	-				

EEC: Concentration of pesticide on food item. At the screening level, relevant food items representing the most conservative EEC for each feeding guild are used.

Table 15Screening Level Assessment of Seed Treatment with Mandestrobin for Birds
and Mammals

	Study Endpoint (mg a.i./kg bw/day/UF)	EDE* (mg a.i./kg bw/day)	RQ	Level of concern
Small bird (0.02 kg				
Acute	100.00	25.394	0.3	Not exceeded
Reproduction	91.10	25.394	0.3	Not exceeded
Medium bird (0.10 kg)	·			·
Acute	100.00	19.947	0.2	Not exceeded
Reproduction	91.10	19.947	0.2	Not exceeded
Large bird (1.00 kg)				
Acute	100.00	5.815	0.1	Not exceeded
Reproduction	91.10	5.815	0.1	Not exceeded
Small mammals (0.015 kg)			
Acute	200.00	14.512	0.1	Not exceeded
Reproduction	166	14.512	0.3	Not exceeded
Medium mammals (0.035	kg)			
Acute	200.00	12.480	0.1	Not exceeded
Reproduction	166	12.480	0.2	Not exceeded
Large mammals (1.00 kg)				
Acute	200.00	6.872	0.03	Not exceeded
Reproduction	166	6.872	0.1	Not exceeded
FIR: Food Ingestion Rate. For gen with body weight greater than 200	e; is calculated using the following formula: (FIR/b eric birds with body weight less than or equal to 200 g, the "all birds" equation was used: cor = 200 g): FIR (g dry weight/day) = 0.398 (bw in) g, the "passerine" equ	ation was us	sed; for generic birds

All birds Equation (body weight >200 g): FIR (g dry weight/day) = 0.598 (bw in g) 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. At the screening level, relevant food items representing the most conservative EEC for each feeding guild are used.

Table 16Toxicity of Mandestrobin, S-2200-*R*-Isomer, S-2200-S-Isomer and
Transformation Products to Non-Target Aquatic Species

Organism	Exposure	Test substance	Endpoint value	Degree of toxicity ¹	Reference
Freshwater species					
Water flea,	48-h Acute	S-2200 TG	$EC_{50} = 1.2 \text{ mg a.i./L}$	Moderately toxic	2378022
Daphnia magna		S-2200 (R-	$EC_{50} = 0.92 \text{ mg/L}$	Highly toxic	2378026
		Isomer)			
		S-2200 (S-	$EC_{50} > 14 \text{ mg/L}$	Slightly toxic at the	2378027
		Isomer)		highest	
				concentration tested	
		2-COOH-S-	$EC_{50} > 100 \text{ mg/L}$	Practically non-	2378023
		2200		toxic at the highest	
				concentration tested	
		5-COOH-S-	$EC_{50} > 100 \text{ mg/L}$	Practically non-	2378024
		2200		toxic at the highest	

Organism	Exposure	re Test substance Endpoint value		Degree of toxicity ¹	Reference	
0	-			concentration tested		
		S-2200 -OR	EC ₅₀ > 14 mg	Slightly toxic at the highest concentration tested	2378028	
		S-2200 -ORC	$EC_{50} > 2.5 \text{ mg/L}$	Moderately toxic up to the limit of solubility of the test	2378025	
	21-d Chronic	S-2200 TG	NOEC = 0.56 mg a.i./L	n/a	2378029	
Sediment dwelling invertebrate, <i>Chironomus dilutus</i>	65-d Chronic	S-2200 TG	NOEC = 3.0 mg a.i./kg dw sediment, 4.93 mg a.i./L porewater	n/a	2378036	
Sediment dwelling invertebrate, <i>Chironomus</i> <i>riparius</i>	28-d Chronic	S-2200 TG	NOEC = 8.1 mg a.i./L	n/a	2378033	
Amphipod, Hyalella azteca	42-d Life cycle	S-2200 TG	NOEC = 5.0 mg a.i./kg dw sediment and 6.38 mg a.i./L in pore water	n/a	2378035	
Rainbow trout,	96-h Acute	S-2200 TG	$LC_{50} = 0.93 \text{ mg a.i./L}$	Highly toxic	2378037	
Oncorhynchus mykiss		S-2200 (<i>R</i> - Isomer)	$LC_{50} = 0.84 \text{ mg/L}$	Highly toxic	2378038	
		S-2200 (<i>S</i> - Isomer)	$LC_{50} > 12 \text{ mg/L}$ (highest concentration tested)	Slightly toxic	2378039	
		2-COOH-S- 2200	LC ₅₀ >89 mg/L (highest concentration tested)	Slightly toxic	2378040	
		5-COOH-S- 2200	$LC_{50} > 100 \text{ mg/L}$ (highest concentration tested)	Practically non toxic	2378041	
		S-2200 -OR	LC ₅₀ >9.0 mg/L (highest concentration tested)	Moderately toxic	2378042	
		S-2200 -ORC	$LC_{50} > 1.4 \text{ mg/L}$ (highest concentration with on precipitate)	Moderately toxic	2378043	
Bluegill sunfish, Lepomis macrochirus)	96-h Acute	S-2200 TG	$LC_{50} = 2.4 \text{ mg a.i./L}$	Moderately toxic	2378044	
Fathead minnow, Pimephales promelas	96-h Acute	S-2200 TG	$LC_{50} = 1.0 \text{ mg a.i./L}$	Highly toxic	2378045	
Fathead minnow, (<i>Pimephales</i> promelas)	28-d Chronic	S-2200 TG	NOEC = 0.15 mg a.i./L (28 days post-hatch)	n/a	2378047	
Green algae, Pseudokirchneriell	96-h Acute	S-2200 TG	$\frac{\text{EC}_{50} = 0.39 \text{ mg a.i./L}}{\text{(inhibition)}}$	n/a	2378060	
a subcapitata	72-h Acute	S-2200 (<i>R</i> - Isomer)	$E_b C_{50} = 0.38 \text{ mg/L}$	n/a	2378067	

Organism	Exposure Test substance		Endpoint value	Degree of toxicity ¹	Reference	
		S-2200	E _b C ₅₀ >12 mg/L	n/a	2378068	
		(S-Isomer)	(highest			
			concentration tested)			
		2-COOH-S-	$E_bC_{50} = 58 \text{ mg/L}$	n/a	2378061	
		2200				
		5-COOH-S-	E _b C ₅₀ >54 mg/L	n/a	2378062	
		2200	(highest			
			concentration tested)			
		S-2200 -OR	$E_bC_{50} > 9.9 \text{ mg/L}$	n/a	2378063	
			(highest			
			concentration tested)			
		S-2200 -ORC	$E_bC_{50} > 5.0 \text{ mg/L}$	n/a	2378059	
			(highest			
			concentration tested)			
Blue-green algae,	96-h Acute	S-2200 TG	$EC_{50} = 0.065 \text{ mg}$	n/a	2378065	
Anabaena flos-			a.i./L (yield)			
aquae			NOEC = 0.023 mg			
			a.i./L (algistatic)			
Diatom, Navicula	96-h Acute	S-2200 TG	$EC_{50} = 1.7 \text{ mg a.i./L}$	n/a	2378066	
pelliculosa			NOEC = 1.1 mg			
			a.i./L			
Monocot vascular	7-d	S-2200 TG	$EC_{50} > 2.3 \text{ mg a.i./L}$	n/a	2378072	
plant, duckweed,	Dissolved		NOEC = 0.32 mg			
Lemna gibba			a.i./L (Frond number)			
Marine/estuarine sp	oecies					
Crustacean,	96-h Acute	S-2200 TG	$LC_{50} = 0.43 \text{ mg a.i./L}$	Highly toxic	2378031	
mysid shrimp,	36-d	S-2200 TG	NOEC = 0.049 mg	n/a	2378032	
Americamysis	Chronic		a.i./L			
bahia						
Sediment-dwelling	28-d	S-2200 TG	NOEC = 10.3 mg	n/a	2378034	
amphipod,	Chronic		a.i./kg dw sediment,			
Leptocheirus			1.56 mg a.i./L			
plumulosus			porewater			
Mollusk, Eastern	96-h Acute	S-2200 TG	$EC_{50} = 2.0 \text{ mg a.i./L}$	Moderately toxic	2378030	
oyster, Crassostrea						
virginica						
Sheepshead	96-h Acute	S-2200 TG	LC ₅₀ >2.2 mg a.i./L	Moderately toxic	2378046	
minnow,	28-d	S-2200 TG	NOEC = 0.64 mg	n/a	2378048	
Cyprinodon	Chronic		a.i./L			
variegatus			(28 days post-hatch)			
Marine diatom,	96-h Acute	S-2200 TG	$EC_{50} = 0.5 \text{ mg a.i./L}$	n/a	2378069	
Skeletonema			NOEC = 0.18 mg			
costatum			a.i./L (yield)			

Organism		Exposure	Endpoint Value (mg a.i./L)	EEC (mg a.i./L)	RQ	Level of Concern
Freshv	vater species	·	· · · · · · · · · · · · · · · · · · ·			
Inverte	brates (Daphnia	Acute	EC ₅₀ /2: 0. 6	0.24	0.4	Not exceeded
magna)	Chronic	NOEC: 0.56	0.24	0.4	Not exceeded
	ent Invertebrate nomus riparius/ dilutes)	Chronic	NOEC: 4.93 (porewater)	0.057*	0.01	Not exceeded
Fish	Oncorhynchus mykiss	Acute	LC ₅₀ /10: 0.093	0.24	3.0	Exceeded
	Pimephales promelas	Chronic	NOEC: 0.15	0.24	1.9	Exceeded
Amphi	bians	Acute,	LC ₅₀ /10: 0.093	1.26	13.5	Exceeded
(fish en	nd-points)	Chronic	NOEC: 0.15	1.26	8.4	Exceeded
Algae ((Anabaena flosaquae)	Acute	EC ₅₀ /2: 0.0325	0.24	7.4	Exceeded
Vascular plants (monocot, <i>Lemna gibba</i>)		Dissolved	EC ₅₀ /2: 1.15	0.24	0.2	Not exceeded
Marin	e species					
Crustac	cean (Americamysis	Acute	LC ₅₀ /2: 0.215	0.24	1.1	Exceeded
bahia)		Chronic	NOEC: 0.049	0.24	4.9	Exceeded
Mollusk (Crassostrea virginica)		Acute	EC ₅₀ /2: 1	0.24	0.2	Not exceeded
Fish (C	Cyprinodon variegatus)	Acute	$LC_{50}/10: > 0.22$	0.24	<1.1	Exceeded
		Early-life stage	NOEC: 0.64	0.24	0.4	Not exceeded
Algae ((Skeletonema costatum)	Acute	EC ₅₀ /2: 0.25	0.24	0.96	Not exceeded
Sediment Invertebrate (Leptocheirus plumulosus)		Chronic	NOEC: 1.56 (porewater)	0.057*	0.04	Not Exceeded
· •	EEC in sediment pore wa	ater from aquatic ed	coscenario modelling		•	·

Table 17 Screening Level Risk Assessment of Mandestrobin to Aquatic Organisms

Table 18Screening Level Risk Assessment of Mandestrobin Isomers and
Transformation Products for Terrestrial and Aquatic Organisms

Organism (exposure)	Compounds	Endpoint Value	EEC	RQ	Level of
		(mg/L)	(mg/L)		Concern
Earthworms	2-COOH-S-2200	LC ₅₀ /2:>500*	0.92*	< 0.1	Not exceeded
Eisenia foetida (acute)	5-COOH-S-2200	LC ₅₀ /2:>500*	0.92*	< 0.1	Not exceeded
Invertebrates	S-2200 R-isomer	LC ₅₀ /2: 0.46	0.24	0.5	Not exceeded
Daphnia magna (acute)	S-2200 S-isomer	$LC_{50}/2:>7$	0.24	< 0.1	Not exceeded
	2-COOH-S-2200	$LC_{50}/2:>50$	0.26	< 0.1	Not exceeded
	5-COOH-S-2200	$LC_{50}/2:>50$	0.26	< 0.1	Not exceeded
	S-2200-OR	LC ₅₀ /2:>7	0.24	< 0.1	Not exceeded
	S-2200-ORC	LC ₅₀ /2: >1.25	0.20	< 0.2	Not exceeded
Fish	S-2200 R-isomer	LC ₅₀ /10: 0.084	0.24	2.9	Exceeded
Oncorhynchus mykiss	S-2200 S-isomer	$LC_{50}/10: > 1.2$	0.24	< 0.2	Not exceeded
(acute)	2-COOH-S-2200	$LC_{50}/10: > 8.9$	0.26	< 0.1	Not exceeded
	5-COOH-S-2200	$LC_{50}/10: > 10$	0.26	< 0.1	Not exceeded
	S-2200-OR	$LC_{50}/10: > 0.9$	0.24	< 0.1	Not exceeded
	S-2200-ORC	$LC_{50}/10: > 0.14$	0.20	<1.4	Exceeded

Organism (exposure)	Compounds	Endpoint Value	EEC	RQ	Level of
		(mg/L)	(mg/L)		Concern
Algae	S-2200 R-isomer	EC ₅₀ /2: 0.19	0.24	1.3	Exceeded
Pseudokirchneriella	S-2200 S-isomer	$EC_{50}/2:>6$	0.24	< 0.1	Not exceeded
subcapitata (acute)	2-COOH-S-2200	EC ₅₀ /2: 29	0.26	0.1	Not exceeded
	5-COOH-S-2200	EC ₅₀ /2: >27	0.26	< 0.1	Not exceeded
	S-2200-OR	EC ₅₀ /2: >4.95	0.24	< 0.1	Not exceeded
	S-2200-ORC	EC ₅₀ /2: >2.5	0.20	< 0.1	Not exceeded
*mg/kg dw soil					

Table 19Refined Risk Assessment of Potential Risk from Drift of Mandestrobin, S-
2200 R-isomer and the Transformation Product S-2200-ORC to Aquatic
Organisms

Organism	Exposure	Endpoint value	Refined EEC	RQ	Level of Concern
Mandestrobin					
Fish Oncorhynchus mykiss	Acute	LC ₅₀ /10: 0.093 mg a.i./L	Aerial appl. (17% drift): 0.0401 mg a.i./L	0.4	Not exceeded
Fish Pimephales promelas	Chronic	NOEC: 0.15 mg a.i./L	Aerial appl. (17% drift): 0.0401 mg a.i./L	0.3	Not exceeded
Amphibians (fish end-points)	Acute,	LC ₅₀ /10: 0.093 mg a.i./L	Aerial appl. (17% drift): 0.214 mg a.i./L	2.3	Exceeded
			Ground appl. (3% drift): 0.038 mg a.i./L	0.4	Not exceeded
	Chronic	NOEC: 0.15 mg a.i./L	Aerial appl. (17% drift): 0.214 mg a.i./L	1.4	Exceeded
			Ground appl. (3% drift): 0.038 mg a.i./L	0.3	Not exceeded
Algae Anabaena flos-aquae	Acute	EC ₅₀ /2: 0.0325 mg a.i./L	Aerial appl. (17% drift): 0.0401 mg a.i./L	1.2	Exceeded
			Ground appl. (3% drift): 0.007 mg a.i./L	0.2	Not exceeded
Crustacean Mysid shrimp	Acute	LC ₅₀ /2: 0.215 mg a.i./L	Aerial appl. (17% drift): 0.0401 mg a.i./L	0.2	Not exceeded
Americamysis bahia	Chronic	NOEC: 0.049 mg a.i./L	Aerial appl. (17% drift): 0.0401 mg a.i./L	0.8	Not exceeded
Fish Pimephales promelas	Acute	LC ₅₀ /10: > 0.133 mg a.i./L	Aerial appl. (17% drift): 0.0401 mg a.i./L	< 0.3	Not exceeded
S-2200 R-isomer					
Fish Oncorhynchus mykiss	acute	LC ₅₀ /10: 0.084 mg/L	Aerial appl. (17% drift): 0.0401 mg/L	0.5	Not exceeded
Algae Pseudokirchneriella subcapitata	Acute	EC ₅₀ /2: 0.19 mg/L	Aerial appl. (17% drift): 0.0401 mg/L	0.2	Not exceeded
S-2200-ORC	•	•	•		•
Fish Oncorhynchus mykiss	Acute	$\begin{array}{c} LC_{50} / 10: > 0.14 \\ mg/L \end{array}$	Aerial appl. (17% drift): 0.034 mg/L	<0.2	Not exceeded

Table 20	Risk Quotients for Aquatic Organisms Determined for Runoff of
	Mandestrobin, S-2200-R-Isomer and S-2200-ORC in Water Bodies

Organism	Exposure	Endpoint value	Refined EEC	RQ	Level of
		(mg/L)	(mg/L)		Concern
Mandestrobin					
Oncorhynchus mykiss	Acute	LC ₅₀ /10: 0.093	0.0821	0.9	Not exceeded
Pimephales promelas	Chronic	NOEC: 0.15	0.0821	0.5	Not exceeded
Amphibians	Acute	LC ₅₀ /10: 0.093	0.223	2.4	Exceeded
	Chronic	NOEC: 0.15	0.223	1.5	Exceeded
Anabaena flosaquae	Acute	EC ₅₀ /2: 0.0325	0.0821	2.5	Exceeded
Americamysis bahia	Acute	LC ₅₀ /2: 0.215	0.0821	0.4	Not exceeded
	Chronic	NOEC: 0.049	0.0821	1.7	Exceeded
Cyprinodon variegatus	Acute	$LC_{50}/10: > 0.22$	0.0821	< 0.4	Not exceeded
S-2200 R-isomer					
Oncorhynchus mykiss	Acute	LC ₅₀ /10: 0.084	0.0821	0.98	Not exceeded
Pseudokirchneriella	Acute	EC ₅₀ /2: 0.19	0.0821	0.4	Not exceeded
subcapitata					
S-2200-ORC					
Oncorhynchus mykiss	Acute	$LC_{50}/10: > 0.14$	0.0821	<0.6	Not exceeded

Table 21Toxic Substances Management Policy Considerations-Comparison to TSMP
Track 1 Criteria

TSMP Track 1 Criteria	TSMP Track 1 Criterion value		Mandestrobin Endpoints
Toxic or toxic equivalent as defined by the <i>Canadian Environmental</i> <i>Protection Act</i> ¹	Yes		Yes
Predominantly anthropogenic ²	Yes		Yes
Persistence ³ :	Soil	Half-life ≥ 182 days	Laboratory studies: DT_{50} of 37.5 to 22,305 days in aerobic soil and 5 to 35 years in anaerobic soil Field studies: DT_{50} of 114 - 173 days
	Water	Half-life $\geq 182 \text{ days}$	DT_{50} of 161 to 781 days in aquatic aerobic system and 458 days to 24 years in anaerobic aquatic systems.
	Sediment	Half-life \geq 365 days	Total system DT_{50} values range from 161 to 8822 days in aerobic and anaerobic water-sediment systems.
	Air	Half-life ≥ 2 days or evidence of long range transport	Volatilisation is not an important route of dissipation and long-range atmospheric transport is unlikely to occur based on the vapour pressure $(3.36 \times 10^{-8} \text{ Pa at } 20^{\circ}\text{C})$ and Henry's law constant $(6.5 \times 10^{-12} \text{ atm m}^3/\text{mol at } 20^{\circ}\text{C})$.

Bioaccumulation ⁴	$\text{Log } K_{\text{OW}} \ge 5$	3.51 at 25°C; Criteria not met	
	bioconcentration factor	25-26	
	\geq 5000		
	bioaccumulation factor	Not available	
	\geq 5000		
Is the chemical a TSMP Tr	ack 1 substance (all four	No, does not meet TSMP Track 1 criteria.	
criteria must be met)?			
¹ All pesticides will be considered	¹ All pesticides will be considered toxic or toxic equivalent for the purpose of initially assessing a pesticide against the TSMP criteria. Assessme		
of the toxicity criteria may be refi	of the toxicity criteria may be refined if required (in other words, all other TSMP criteria are met).		
² The policy considers a substance "predominantly anthropogenic" if, based on expert judgement, its concentration in the environment medium			
largely due to human activity, rather than to natural sources or releases.			
³ If the pesticide and/or the transformation product(s) meet one persistence criterion identified for one media (soil, water, sediment or air) than			
criterion for percistence is considered to be met			

criterion for persistence is considered to be met. ⁴Field data (for example, bioaccumulation factors) are preferred over laboratory data (for example, bioconcentration factors) which, in turn, are preferred over chemical properties (for example, log K_{ow}).

Table 22Fungicide Resistance Action Committee modes of action groups of currently
registered alternative products (as of June 2015)

Сгор	Disease	Fungicide Resistance Action Committee Mode of Action groups of registered alternatives
Foliar applied pr	oducts	
Canola and crop subgroup 20A	White mold/sclerotinia rot	2; 3; 7; 7+11; 9+12; 11; 44; NC* (<i>Coniothyrium minitans</i> strain CON/M/91-08)
Grape and crop subgroup 13-07F	Botrytis bunch rot / gray mold	2; 7; 7+11; 7+9; 9; 9+12; 17; 44; P5: NC* (<i>Aureobasidium pullulans</i> DSM 14940 and DSM 14941; BLAD polypeptide)
	Powdery mildew	3; 7; 7+9; 7+11; 11; 13; 29+M2; 44; 46; M2; M4; P5; U8; NC* (mineral oil; potassium bicarbonate; <i>Streptomyces lydicus</i> strain WYEC 108; Garlic powder; BLAD polypeptide)
Strawberry and crop subgroup 13- 07G	Botrytis gray mold	1; 2; 7; 7+11; 9; 9+12; 17; 44 M2; M3; M5; P5; NC* (<i>Trichoderma harzianum</i> Rifai strain KRL-AG2; BLAD polypeptide; <i>Streptomyces lydicus</i> strain WYEC 108)
Turfgrass	Dollar Spot	1; 2; 3; 3+11; 3+ M5; 7; 11; 44; M5; NC* (mineral oil; <i>Trichoderma harzianum</i> Rifai strain KRL-AG2; <i>T. harzianum</i> Rifai strain T-22)
	Brown Patch	1; 23; 3+11; 3+M; 7; 11; 12; 14; 44; M5 ; NC* (mineral oil; hydrogen peroxide)
	Fairy Ring	3+11; 7+11; 11
	Rust	11
	Take-all Patch	3; 11
Seed treatment prod		
Corn	Rhizoctonia solani seed rot	3; 3+4+7; 7; 7+11; 11
T	Fusarium seed rot	3; 3+4; 3+4+7; 4+11; 7; 7+11; 11
Legume	Rhizoctonia solani seed rot	1+4+12; 3+4+7; 4+11; 4+11+12; 4+12; 7; 7+11; 7+M3;11; 44
vegetables and crop group 6	Fusarium seed rot	1+4+12; 3; 3+4; 3+4+7; 4+11; 4+11+12; 4+12; 7; 7+11; 7+M3; 11; 44
	Phomopsis seed rot	3; 3+4+7; 3+7; 4+12; 4+11; 4+11+12; 7+11; 7+M3; 11
Canola and crop	Rhizoctonia solani seed rot	3; 3+4+12; 3+4+7+12; 3+7; 4+7+11; 7; 7+11; 7+M3; 11
subgroup 20A	Fusarium seed rot	3; 3+4; 3+7; 3+4+12; 3+4+7+12; 4+7+11; 7; 7+11; 11;44

*NC: not classified

Table 23List of Supported Uses

S-2200 4 SC Fungicide; S-2200 4 SC AG Fungicide; S-2200 4 VPP Fungicide (foliar-applied products)

Supported claim	Conclusion and comment from value
	assessment
Control of white mold/sclerotinia rot (Sclerotinia sclerotiorum) on canola	Supported as proposed
(crop subgroup 20A) with one foliar application of 439 – 877 mL/ha	
applied when the crop is at 20-50% bloom.	
Control of botrytis bunch rot / gray mold (Botrytis cinerea) on grape (crop	Supported as proposed
subgroup 13-07F) with three to four foliar applications of 439 - 877 mL/ha	
(seasonal max. 2631 mL/ha) prior to infection during early bloom, bunch	
pre-closure and veraison up to 10 days before harvest with an interval of	
10 days for sequential applications.	
Suppression of powdery mildew (Uncinula necator) on grape (crop	Supported as proposed; claim limited
subgroup 13-07F) with three to four foliar applications of 439 - 877 mL/ha	to susceptible crops only (grape and
(seasonal max. 2631 mL/ha) prior to infection at bud break with 10 to 14	Amur grape)
intervals.	
Control of botrytis gray mold (Botrytis cinerea) on strawberry (crop	Supported as proposed
subgroup 13-07G) with four to five foliar applications of 439 – 877 mL/ha	
(seasonal max. 3508 mL/ha) starting at 10% bloom, or prior to infection	
up to 0 days before harvest with reapplication intervals of 7 to 14 day.	
Control of dollar spot (Sclerotinia homoeocarpa) on turfgrass with	Supported as proposed
applications of 540-986 mL/ha (seasonal max. 3944 mL/ha) with 14 to 28	
day intervals starting when conditions favor disease development.	
Control of brown Patch (Rhizoctonia solani) on turfgrass with applications	Supported as proposed
of 986 mL/ha (seasonal max. 3944 mL/ha) with 14 day intervals starting	
when conditions favor disease development or when the disease first	
appears.	
Control of fairy ring (various basidiomycetes) on turfgrass with	Supported with the claim specified to
applications of 986 mL/ha (seasonal max. 3944 mL/ha) with 14 day	Agaricus campestris rather than
intervals starting when conditions favor disease development or when the	"various basidiomycetes" as the causal
disease first appears.	pathogen
Suppression of rust diseases (Puccinia spp.) on turfgrass with applications	Supported with the claim specified to
of 986 mL/ha (seasonal max. 3944 mL/ha) with 14 day intervals starting	'Rust (Puccinia graminis)' rather than
when conditions favor disease development or when the disease first	the general claim against 'Rust diseases
appears.	Puccinia spp.'
Control of take-all patch (Gaeumannomyces graminis) on turfgrass with	Supported as proposed
applications of 986 mL/ha (seasonal max. 3944 mL/ha) with 14 day	
intervals starting when conditions favor disease development or when the	
disease first appears.	

S-2200 3.2 FS Fungicide (seed treatment product)

Supported claim	Conclusion and comment from value assessment
Control of seed decay caused by <i>Rhizoctonia solani</i> in on corn (field corn, sweet corn, and popcorn) with one application seed of 15.6 mL/ 100 kg seed.	Supported as proposed
Control of seed decay caused by <i>Fusarium</i> spp. in on corn (field corn, sweet corn, and popcorn) with one application seed of 15.6 mL/ 100 kg seed.	Supported as proposed
Control of seed decay caused by <i>Rhizoctonia solani</i> in legume vegetables (crop group 6) with one seed application of 26 mL/ 100 kg seed.	Supported as proposed

Control of seed decay caused by <i>Fusarium</i> spp. in legume vegetables (crop group 6) with one seed application of 26 mL/ 100 kg seed.	Supported as proposed
Suppression of seed decay caused by <i>Phomopsis</i> spp. in legume vegetables (crop group 6) with one seed application of 26 mL/ 100 kg seed.	The causal pathogen for the claim is specified to <i>Phomopsis longicolla</i> .
Control of seed decay caused by <i>Rhizoctonia solani</i> in canola (crop subgroup 20A) with one seed application of 26 mL/100 kg seed.	Supported as proposed
Control of seed decay caused by <i>Fusarium</i> spp. in canola (crop subgroup 20A) with one seed application of 26 mL/100 kg seed.	Supported as proposed

Appendix II Supplemental Maximum Residue Limit Information— International Situation and Trade Implications

Mandestrobin is a new active ingredient which is concurrently being registered in Canada and the United States. The MRLs proposed for mandestrobin in Canada are the same as corresponding tolerances to be promulgated in the United States.

Once established, the American tolerances for mandestrobin will be listed in the Electronic Code of Federal Regulations, 40 CFR Part 180, by pesticide.

Currently, there are no Codex MRLs⁹ listed for mandestrobin in or on any commodity on the Codex Alimentarius Pesticide Residues in Food website.

{Table 1 compares the MRLs proposed for mandestrobin in Canada with corresponding American tolerances and Codex MRLs¹⁰. American tolerances are listed in the Electronic Code of Federal Regulations, 40 CFR Part 180, by pesticide. A listing of established Codex MRLs is available on the Codex Alimentarius Pesticide Residues in Food website, by pesticide or commodity}.

Food Commodity	Canadian MRL (ppm)	American Tolerance (ppm)	Codex MRL (ppm)
Raisins	7.0	7.0	Not established
Small fruit vine climbing (Crop Subgroup 13-07F, except fuzzy kiwifruit)	5.0	5.0	Not established
Low growing berry (Crop Subgroup 13-07G, except cranberry)	3.0	3.0	Not established
Rapeseed (Crop Subgroup 20A)	0.5	0.5	Not established
Legume vegetables (Crop Group 6, except cowpea and field pea), corn (field, popcorn, sweet)	0.02	0.02	Not established

Table 1 Comparison of Canadian MRLs, American Tolerances and Codex MRLs

MRLs may vary from one country to another for a number of reasons, including differences in pesticide use patterns and the locations of the field crop trials used to generate residue chemistry data. For animal commodities, differences in MRLs can be due to different livestock feed items and practices.

⁹ The <u>Codex Alimentarius Commission</u> is an international organization under the auspices of the United Nations that develops international food standards, including MRLs.

¹⁰ The <u>Codex Alimentarius Commission</u> is an international organization under the auspices of the United Nations that develops international food standards, including MRLs.

References

A. List of Studies/Information Submitted by Registrant

1.0 Chemistry

PMRA Document Number	Reference
2377784	2012, S-2200 Technical: Product Identity and Composition, Description of Materials Used to Produce the Product, Description of Production Process and Discussion of Formation of Impurities, DACO: 2.11.1, 2.11.2, 2.11.3, 2.11.4
2377785	2012, S-2200 Technical: Product Identity and Composition, Description of Materials Used to Produce the Product, Description of Production Process and Discussion of Formation of Impurities, DACO: 2.11.1, 2.11.2, 2.11.3, 2.11.4 CBI
2377786	2013, S-2200 Fungicide Technical: Product Identity and Composition; Certified Limits; Enforcement Analytical Method, DACO: 2.12.1, 2.13.1, 2.4, 2.5, 2.6, 2.7, 2.8, 2.9
2377787	2013, S-2200 Fungicide Technical: Product Identity and Composition; Certified Limits; Enforcement Analytical Method, DACO: 2.12.1, 2.13.1, 2.4, 2.5, 2.6, 2.7, 2.8, 2.9 CBI
2377788	2012, Preliminary Analysis and Enforcement Analytical Method of S-2200 Technical Grade for Confirmation of Its Racemic Composition, DACO: 2.13.1
2377789	2012, Preliminary Analysis and Enforcement Analytical Method of S-2200 Technical Grade for Confirmation of Its Racemic Composition, DACO: 2.13.1 CBI
2377790	2012, Enforcement Analytical Methods of S-2200 Technical Grade, DACO: 2.13.1
2377791	2012, Enforcement Analytical Methods of S-2200 Technical Grade, DACO: 2.13.1 CBI
2377792	2013, Analytical Method for the Determination of the Contents of S-2200 on Dry and Wet Weight Basis, DACO: 2.13.1
2377793	2012, Preliminary Analysis of S-2200 Technical Grade, DACO: 2.13.1, 2.13.2, 2.13.3
2377794	2012, Preliminary Analysis of S-2200 Technical Grade, DACO: 2.13.1, 2.13.2, 2.13.3 CBI
2377795	2012, S-2200TG: Determination of Physical State, Colour and Odour and Relative Density, DACO: 2.14.1, 2.14.2, 2.14.3, 2.14.6

	-
2377796	2010, Product Chemistry Testing for S-2200 PAI, DACO: 2.14.1, 2.14.12, 2.14.2, 2.14.3, 2.14.4, 2.14.5, 2.14.6, 8.2.1
2377797	2010, Product Chemistry Testing for S-2167 PAI, DACO: 2.14.1, 2.14.12, 2.14.2, 2.14.3, 2.14.4, 2.14.5, 2.14.6
2377798	2010, Product Chemistry Testing for S-2354 PAI, DACO: 2.14.1, 2.14.12, 2.14.2, 2.14.3, 2.14.4, 2.14.5, 2.14.6, 8.2.1
2377799	2009, Determination of the Water Solubility of S-2200 PAI, DACO: 2.14.7, 8.2.1
2377800	2009, Determination of the Water Solubility of S-2167 PAI, DACO: 2.14.7, 8.2.1
2377801	2009, Determination of the Water Solubility of S-2354 PAI, DACO: 2.14.7, 8.2.1
2377802	2011, S-2200 PAI - Determination of the Solvent Solubility, DACO: 2.14.8
2377803	2011, S-2167 PAI - Determination of the Solvent Solubility, DACO: 2.14.8
2377804	2011, S-2167 PAI - Determination of the Solvent Solubility, DACO: 2.14.8
2377805	2013, S-2200TG: Determination of the Solvent Solubility, DACO: 2.14.8
2377807	2013, S-2200 Dissociation Constant Waiver Request, DACO: 2.14.10
2377808	2011, S-2200 PAI - Determination of Vapour Pressure, DACO: 2.14.9, 8.2.1
2377810	2011, S-2167 PAI - Determination of Vapour Pressure, DACO: 2.14.9, 8.2.1
2377811	2010, Determination of the Partition Coefficient (n-Octanol/Water) S-2167 PAI, DACO: 2.14.11, 8.2.1
2377812	2010, Determination of the Partition Coefficient (n-Octanol/Water) S-2200 PAI, DACO: 2.14.11, 8.2.1
2377813	2012, Stability of S-2200 Technical Grade to Normal and Elevated Temperatures, Metals and Metal Ions, DACO: 2.14.13
2377814	2013, Storage Stability and Corrosion Characteristics of S-2200 Technical Grade, DACO: 2.14.14 ,2.16
2377816	2010, Determination of Oxidation/Reduction Properties S-2200 TGAI, DACO: 2.16
2377818	2012, S-2200 TG: Determination of Flammability, DACO: 2.16
2377820	2013, S-2200 - Henrys Law Constant, DACO: 2.16
2377822	2010, Determination of the Impact Explodability of S-2200 TGAI, DACO: 2.16
2377824	2011, Determination of the Thermal Explodability of S-2200 TGAI, DACO: 2.16

2377826	2012, Analytical Determination of Active Ingredient and Impurities in S-2200 Technical Grade, DACO: 2.16
2377828	2012, Analytical Determination of Active Ingredient and Impurities in S-2200 Technical Grade, DACO: 2.16 CBI
2377830	2012, Confirmation of Identification of Ingredients in S-2200 Technical Grade, DACO: 2.16
2377832	2012, Confirmation of Identification of Ingredients in S-2200 Technical Grade, DACO: 2.16 CBI
2377834	2013, pH of S-2200 Fungicide Technical, DACO: 2.16
2377861	2010, [14C]S-2167 (S-2200 R-isomer): Hydrolytic Stability, DACO: 8.2.3.2
2377863	2010, [14C]S-2354 (S-2200 S-isomer): Hydrolytic Stability, DACO: 8.2.3.2
2377869	2010, [14C]S-2354 (S-2200 S-isomer): Photodegradation and Quantum Yield in Sterile, Aqueous Solution, DACO: 8.2.3.3.2
2377871	2010, [14C]S-2167 (S-2200 R-isomer): Photodegradation and Quantum Yield in Sterile, Aqueous Solution, DACO: 8.2.3.3.2
2377875	2013, S-2200 - Stability in Air, DACO: 8.6
2443917	2011, Characterization of S-2200 Technical Grade and Its Related Substances, DACO: 2.13.2
2443918	2011, Characterization of S-2200 Technical Grade and Its Related Substances, DACO: 2.13.2 CBI
2377849	2013, S-2200: Independent Laboratory Validation of Valent Method RM-48S- 3, Determination of S-2200, DACO: 8.2.2.1, 8.2.2.2
2377854	2011, Validation of an Analytical Method for the Determination of S-2200 in Surface Water for Post-Registration Control and Monitoring Purpose, DACO: 8.2.2.3
2377857	2013, S-2200: Independent Laboratory Validation of the Analytical Method Validation of an Analytical Method for the Determination of S-2200 in Surface Water for Post Registration Control and Monitoring Purposes, DACO: 8.2.2.3
2377924	2011, S-2200 and Metabolites: Storage Stability of Residues in EU Soil Stored Deep Frozen, DACO: 8.6
2377958	2013, Calculation of Laboratory Soil Kinetics for S-2200 and its Major Metabolites According to FOCUS (2006) Guidance, DACO: 8.6
2377851	2013, S-2200 4 SC Ag Fungicide: Product Identity, Composition, and Analysis, DACO: 3.2.1, 3.2.2, 3.2.3, 3.3.1, 3.4.1, 3.4.2
2377853	2013, S-2200 4 SC Ag Fungicide: Product Identity, Composition, and Analysis, DACO: 3.2.1, 3.2.2, 3.2.3, 3.3.1, 3.4.1, 3.4.2 CBI

2377856	2013, Physical and Chemical Properties of S-2200 4 SC Fungicide, VC-1902, DACO: 3.5.1, 3.5.10, 3.5.11, 3.5.12, 3.5.13, 3.5.14, 3.5.15, 3.5.2, 3.5.3, 3.5.6, 3.5.7, 3.5.8, 3.5.9
2378123	2013, S-2200 3.2 FS Fungicide: Product Identity, Composition, and Analysis, DACO: 3.2.1, 3.2.2, 3.2.3, 3.3.1, 3.4.1, 3.4.2
2378124	2013, S-2200 3.2 FS Fungicide: Product Identity, Composition, and Analysis, DACO: 3.2.1, 3.2.2, 3.2.3, 3.3.1, 3.4.1, 3.4.2 CBI
2378125	2013, Physical and Chemical Properties of S-2200 3.2 FS Fungicide, VC- 1929, DACO: 3.5.1, 3.5.10, 3.5.11, 3.5.12, 3.5.13, 3.5.14, 3.5.15, 3.5.2, 3.5.3, 3.5.6, 3.5.7, 3.5.8, 3.5.9

2.0 Human and Animal Health

PMRA	Reference
Document Number	
2377929	2010, Acute Oral Toxicity Study of S-2200 TG in Rats, DACO: 4.2.1
2377931	2011, Acute Oral Toxicity Study of De-Xy-S-2200 in Rats, DACO: 4.2.1
2377932	2012, Acute Oral Toxicity Study of 2-COOH-S-2200 in Rats, DACO: 4.2.1
2377933	2012, Acute Oral Toxicity Study of 5-COOH-S-2200 in Rats, DACO: 4.2.1
2377935	2012, Acute Oral Toxicity Study of 2-CH2OH-S-2200 in Rats, DACO: 4.2.1
2377936	2010, Acute Dermal Toxicity Study of S-2200 TG in Rats, DACO: 4.2.2
2377937	2010, Acute Inhalation Toxicity Study of S-2200 TG in Rats, DACO: 4.2.3
2377939	2010, Primary Eye Irritation Test of S-2200 TG in Rabbits, DACO: 4.2.4
2377940	2010, Primary Skin Irritation Test of S-2200 TG in Rabbits, DACO: 4.2.5
2377941	2010, Skin Sensitization Test of S-2200 TG in Guinea Pigs (Maximization Test), DACO: 4.2.6
2377943	2009, S-2200: 14 Day Dietary Administration Range-Finding Study in the Mouse, DACO: 4.3.1
2377944	2012, Acute Oral Toxicity Study of 4-OH-S-2200 in Rats, DACO: 4.2.1
2377946	2011, S-2200 Technical Grade: 13 Week Oral (Dietary) Administration Toxicity Study in the Mouse, DACO: 4.3.1
2377948	2011, S-2200 Technical Grade: 13 Week Oral (Dietary) Administration Toxicity Study in the Rat, DACO: 4.3.1
2377951	2008, S-2200: 1 Week Palatability Study in the Dog, DACO: 4.3.2
2377952	2012, S-2200 Technical Grade: 52 Week Oral (Dietary) Administration Toxicity Study in the Dog, DACO: 4.3.2
2377954	2012, S-2200 Technical Grade: 13 Week Oral (Dietary) Administration Toxicity Study in the Dog, DACO: 4.3.2
2377956	2009, S-2200: 4 Week Dietary Study in the Dog, DACO: 4.3.3
2377957	2011, A 28-Day Repeated Dose Dermal Toxicity Study of S-2200 Technical Grade in Rats, DACO: 4.3.5
2377959	2013, Waiver Request: S-2200 Fungicide Technical Subchronic (28-Day) Rat Inhalation Study, DACO: 4.3.6

2377960	2012, S-2200 Technical Grade: 78 Week Oral (Dietary) Administration Carcinogenicity Study in the Mouse, DACO: 4.4.3
2377961	2012, S-2200 Technical Grade: 104 Week Oral (Dietary) Administration
	Combined Toxicity/Carcinogenicity Study in the Rat, DACO: 4.4.4
 	2010, Dose Range-Finding Study for Two-Generation Reproduction Study of
2377962	
	S-2200 TG in Rats, DACO: 4.5.1
2377964	2012, Two-Generation Reproduction Toxicity Study of S-2200 TG in Rats,
	DACO: 4.5.1
2377965	2009, S-2200 TG: Oral (Gavage) Range-Finding Study of Prenatal
2377703	Development in the Rat, DACO: 4.5.2
2277067	2012, S-2200 TG: Oral (Gavage) Prenatal Development Toxicity Study in the
2377967	Rat, DACO: 4.5.2
22550 60	2009, S-2200 TG: Oral (Gavage) Range-Finding Study of Prenatal
2377969	Development in the Rabbit, DACO: 4.5.3
	2012, S-2200 TG: Oral (Gavage) Prenatal Development Toxicity Study in the
2377970	Rabbit, DACO: 4.5.3
	2010, Reverse Mutation Test of S-2200 TG in Bacterial Systems, DACO:
2377971	4.5.4
2377972	2011, Reverse Mutation Test of De-XY-S-2200 in Bacterial Systems, DACO:
	4.5.4
2377973	2012, Reverse Mutation Test of 2-COOH-S-2200 in Bacterial Systems,
	DACO: 4.5.4
2377975	2012, Reverse Mutation Test of 5-COOH-S-2200 in Bacterial Systems,
2311713	DACO: 4.5.4
2377976	2012, Reverse Mutation Test of 2-CH2OH-S-2200 in Bacterial Systems,
2311910	DACO: 4.5.4
000000	2012, Reverse Mutation Test of 4-OH-S-2200 in Bacterial Systems, DACO:
2377977	4.5.4
2255050	2010, Gene Mutation Assay in Chinese Hamster V79 Cells In vitro (V79 /
2377978	HPRT) with S-2200 TG, DACO: 4.5.5
	2011, Gene Mutation Assay in Chinese Hamster V79 Cells In vitro
2377980	(V79/HPRT) with 2-COOH-S- 2200, DACO: 4.5.5
	2011, Gene Mutation Assay in Chinese Hamster V79 Cells <i>In vitro</i>
2377981	(V79/HPRT) with 5-COOH-S-2200, DACO: 4.5.5
	2010, <i>In vitro</i> Chromosomal Aberration Test on S-2200 TG in Chinese
2377982	
	Hamster Lung Cells (CHL/IU), DACO: 4.5.6
2377983	2012, <i>In vitro</i> Chromosomal Aberration Test on 2-COOH-S-2200 in Chinese
	Hamster Lung Cells (CHL/IU), DACO: 4.5.6
2377984	2012, <i>In vitro</i> Chromosomal Aberration Test on 5-COOH-S-2200 in Chinese
	Hamster Lung Cells (CHL/IU), DACO: 4.5.6
2377985	2010, Micronucleus Test on S-2200 TG in CD-1 Mice, DACO: 4.5.7
2377986	2012, Micronucleus Assay of 2-COOH-S-2200 in Mice, DACO: 4.5.7
2377987	2011, Metabolism of S-2200 R-isomer (S-2167) and S-2200 S-isomer (S2354)
	in Rats, DACO: 4.5.9
2377989	2012, [14C]S-2200: Absorption, Distribution, Metabolism and Excretion
	Following Repeat Oral Administration to the Rat, DACO: 4.5.9

2377990	2012, [14C]S-2200: Absorption, Distribution, Metabolism and Excretion
	Following Single Oral Administration to the Rat, DACO: 4.5.9
2377991	2011, An Oral (Gavage) Dose Range-Finding Acute Neurotoxicity Study of S-
	2200 TG in Wistar Rats, DACO: 4.5.12
2377992	2011, An Oral (Gavage) Acute Neurotoxicity Study of S-2200TG in Wistar
	Rats, DACO: 4.5.12
2377994	2012, A 90-Day Oral Dietary Neurotoxicity Study of S-2200 TG in Wistar
2311774	Rats, DACO: 4.5.13
2377995	2010, Validation for the Determination of S-2200 in Dietary Formulations
2311))3	Using High Performance Liquid Chromatography (HPLC), DACO: 4.8
2377997	2010, Validation for the Determination of S-2200 in Dietary Formulations
2311991	using High Performance Liquid Chromatography (HPLC), DACO: 4.8
2378000	2010, Validation for the Determination of S-2200 in Canine Diet using High
2378000	Performance Liquid Chromatography (HPLC), DACO: 4.8
2378001	2011, S-2200 TG: Analytical Validation and Stability Study of Analyte
2578001	Concentration in Dietary and Gavage Formulations, DACO: 4.8
2279004	2011, S-2200 TG: A 28-Day Dietary Dose Range-Finding Study in Wistar
2378004	Han Rats, DACO: 4.8
2279005	2011, S-2200 TG: A 28 Day Oral (Dietary) Immunotoxicity Study in Female
2378005	Wistar Han Rats, DACO: 4.8(B)
2270006	2012, In vitro Steroidogenesis Assay of S-2200 TG in H295R Cells, DACO:
2378006	4.8
2279007	2012, Short-Term Study for Mode of Action Analysis for Mouse Liver
2378007	Findings by S-2200 TG, DACO: 4.8
	2012, Short-Term Study for Mode of Action Analysis for Rat Liver and
2378008	Thyroid Findings by S- 2200 TG; -Dose Response, Time-Course, and
	Reversibility-, DACO: 4.8
	2012, The Toxicological Relevance of the Liver and Thyroid Alterations
2378009	Observed in Rats Treated with S-2200 TG Based on Mode of Action, DACO:
	4.8
	2012, Evaluation of Effects of S-2200 TG and its Metabolites on Human
2378010	Estrogen Receptor Alpha and Human Androgen Receptor Using In vitro
	Reporter Gene Assays, DACO: 4.8
	2013, Assessment of Toxicological Significance: Higher Incidence of Ovarian
2378011	Sex-Cord Stromal Tumour in Female Rats Treated with S-2200 TG in a 2-
	Year Carcinogenicity Study, DACO: 4.8
2377860	2013, S-2200 4 SC: Acute Oral Toxicity Up and Down Procedure in Rats,
	DACO: 4.6.1
2377862	2013, S-2200 4 SC: Acute Dermal Toxicity Study in Rats, DACO: 4.6.2
2377865	2013, S-2200 4 SC: Acute Inhalation Toxicity Study in Rats, DACO: 4.6.3
2377866	2013, S-2200 4 SC: Primary Eye Irritation Study in Rabbits, DACO: 4.6.4
2377868	2013, S-2200 4 SC: Primary Skin Irritation Study in Rabbits, DACO: 4.6.5
2377870	2013, S-2200 4 SC: Local Lymph Node Assay (LLNA) in Mice, DACO: 4.6.6
	2012, S-2200 3.2 FS: Acute Oral Toxicity Up and Down Procedure in Rats,
2378126	DACO: 4.6.1

2378127 2012, S-2200 3.2 FS: Acute Dermal Toxicity Study in Rats Limit Test, DACO: 4.6.2 2378128 2012, S-2200 3.2 FS: Acute Inhalation Toxicity Study in Rats, DACO: 4.6.4 2378130 2012, S-2200 3.2 FS: Primary Eye Irritation Study in Rabbits, DACO: 4.6.4 2378131 2012, S-2200 3.2 FS: Local Lymph Node Assay (LLNA) in Mice, DACO: 4.6.6 2377872 2013, Summary of Occupational Risk Assessments for the Use of S-2200 4 SC Fungicide on Canola, Grapes, Strawberries and Turfgrass, DACO: 5.1 2013, Occupational and Residential Exposure and Risk Assessments for S- 2200 4 SC Fungicide on Canola (Crop Subgroup 13-07G), and Turf, DACO: 5.1 2377874 2013, Use Description and Scenario (Mixer/Loader/Applicator and Postapplication) for S-2200 4 SC Fungicide, DACO: 5.2 2013, Pesticide Handlers Exposure Database Assessment in Support of Spray Applications of S-2200 4 SC Fungicide to Canola, Grapes, Strawberries, and Turfgrass, DACO: 5.3 2377880 2013, Postapplication Assessment of S-2200 4 SC Fungicide Following Treatment to Canola, Grapes, Strawberries, and Turf, DACO: 5.6 2377813 2013, Waiver Request: Foliar Disologeable and Turf Transferable Residue Study for S-2200 4 SC Ag and VPP Fungicides, DACO: 5.1 2378132 2013, Summary of Occupational Risk Assessments for the Seed Treatment Use of S-2200 3.2 FS Fungicide on Canola, Corn, and Soybean, DACO: 5.1 2378133 2013, Occupational Exposure and Risk Assessments for S-2200 3.2 FS Fungicide, DACO: 5.14 2378134 <th></th> <th>-</th>		-
 2378129 2012, S-2200 3.2 FS: Primary Eye Irritation Study in Rabbits, DACO: 4.6.4 2378130 2012, S-2200 3.2 FS: Local Lymph Node Assay (LLNA) in Mice, DACO: 4.6.6 2378131 2013, Summary of Occupational Risk Assessments for the Use of S-2200 4 SC Fungicide on Canola, Grapes, Strawberries and Turfgrass, DACO: 5.1 2013, Occupational and Residential Exposure and Risk Assessments for S-2200 4 SC Fungicide on Canola (Crop Subgroup 20A), Grapes (Crop Subgroup 13-07F), Strawberries (Crop Subgroup 20A), Grapes (Crop Subgroup 13-07F), Strawberries (Crop Subgroup 13-07G), and Turf, DACO: 5.1 2377876 2013, Use Description and Scenario (Mixer/Loader/Applicator and Postapflication) for S-2200 4 SC Fungicide, DACO: 5.2 2013, Pesticide Handlers Exposure Database Assessment in Support of Spray Applications of S-2200 4 SC Fungicide to Canola, Grapes, Strawberries, and Turfgrass, DACO: 5.3 2377880 2013, Postapplication Assessment of S-2200 4 SC Fungicide Following Treatment to Canola, Grapes, Strawberries, and Turf Transfrable Residue 2013, Waiver Request: Foliar Dislodgeable and Turf Transfreable Residue 2013, Summary of Occupational Risk Assessments for the Seed Treatment Use of S-2200 3.2 FS Fungicide on Canola, Corn, and Soybean, DACO: 5.1 2378132 2013, Summary of Occupational Risk Assessments for the Seed Treatment Use of S-200 3.2 FS Fungicide on Canola, Corn, and Soybean, DACO: 5.1 2378134 2013, Dust Off Study in Support of the Seed Treatment Use of S-2200 3.2 FS Fungicide reations for Canola, Corn, and Soybean, Sorghum, and Canola (Rapeseed Subgroup 20A), DACO: 5.1 2378134 2013, Dust Off Study in Support of the Seed Treatment Use of S-2200 3.2 FS Fungicide, DACO: 5.4 2378135 2013, Mixer/Loader/Applicator Passive Dosimetry Study in Support of On-Farm Seed Treatment with S-2200 3.2 FS Fungicide, DACO: 5.2 2013, Mixer/Loader/Applicator Passive Dosi	2378127	2012, S-2200 3.2 FS: Acute Dermal Toxicity Study in Rats - Limit Test, DACO: 4.6.2
 2378129 2012, S-2200 3.2 FS: Primary Eye Irritation Study in Rabbits, DACO: 4.6.4 2378130 2012, S-2200 3.2 FS: Local Lymph Node Assay (LLNA) in Mice, DACO: 4.6.6 2378131 2013, Summary of Occupational Risk Assessments for the Use of S-2200 4 SC Fungicide on Canola, Grapes, Strawberries and Turfgrass, DACO: 5.1 2013, Occupational and Residential Exposure and Risk Assessments for S-2200 4 SC Fungicide on Canola (Crop Subgroup 20A), Grapes (Crop Subgroup 13-07F), Strawberries (Crop Subgroup 20A), Grapes (Crop Subgroup 13-07F), Strawberries (Crop Subgroup 13-07G), and Turf, DACO: 5.1 2377876 2013, Use Description and Scenario (Mixer/Loader/Applicator and Postapflication) for S-2200 4 SC Fungicide, DACO: 5.2 2013, Pesticide Handlers Exposure Database Assessment in Support of Spray Applications of S-2200 4 SC Fungicide to Canola, Grapes, Strawberries, and Turfgrass, DACO: 5.3 2377880 2013, Postapplication Assessment of S-2200 4 SC Fungicide Following Treatment to Canola, Grapes, Strawberries, and Turf Transfrable Residue 2013, Waiver Request: Foliar Dislodgeable and Turf Transfreable Residue 2013, Summary of Occupational Risk Assessments for the Seed Treatment Use of S-2200 3.2 FS Fungicide on Canola, Corn, and Soybean, DACO: 5.1 2378132 2013, Summary of Occupational Risk Assessments for the Seed Treatment Use of S-200 3.2 FS Fungicide on Canola, Corn, and Soybean, DACO: 5.1 2378134 2013, Dust Off Study in Support of the Seed Treatment Use of S-2200 3.2 FS Fungicide reations for Canola, Corn, and Soybean, Sorghum, and Canola (Rapeseed Subgroup 20A), DACO: 5.1 2378134 2013, Dust Off Study in Support of the Seed Treatment Use of S-2200 3.2 FS Fungicide, DACO: 5.4 2378135 2013, Mixer/Loader/Applicator Passive Dosimetry Study in Support of On-Farm Seed Treatment with S-2200 3.2 FS Fungicide, DACO: 5.2 2013, Mixer/Loader/Applicator Passive Dosi	2378128	2012, S-2200 3.2 FS: Acute Inhalation Toxicity Study in Rats, DACO: 4.6.3
 2378130 2012, S-2200 3.2 FS: Primary Skin Irritation Study in Rabbits, DACO: 4.6.5 2378131 2012, S-2200 3.2 FS: Local Lymph Node Assay (LLNA) in Mice, DACO: 4.6.6 2377872 2013, Summary of Occupational Risk Assessments for the Use of S-2200 4 SC Fungicide on Canola, Grapes, Strawberries and Turfgrass, DACO: 5.1 2013, Occupational and Residential Exposure and Risk Assessments for S-2200 4 SC Fungicide on Canola (Crop Subgroup 13-07G), and Turf, DACO: 5.1 2013, Use Description and Scenario (Mixer/Loader/Applicator and Postapplication) for S-2200 4 SC Fungicide, DACO: 5.2 2013, Pesticide Handlers Exposure Database Assessment in Support of Spray Applications of S-2200 4 SC Fungicide to Canola, Grapes, Strawberries, and Turfgrass, DACO: 5.3 2377878 2013, Postapplication Assessment of S-2200 4 SC Fungicide Following Treatment to Canola, Grapes, Strawberries, and Turf, DACO: 5.6 2013, Waiver Request: Foliar Dislodgeable and Turf Transferable Residue Study for S-2200 4 SC Ag and VPP Fungicides, DACO: 5.9(A), 5.9(B) 2378132 2013, Summary of Occupational Risk Assessments for the Seed Treatment Use of S-2200 3.2 FS Fungicide on Canola, Corn, and Soybean, DACO: 5.1 2013, Occupational Exposure and Risk Assessments for S-2200 3.2 FS Fungicide, DACO: 5.14 2013, Use Description and Exposure Scenarios for Canola, Corn, and Soybean, Sorghum, and Canola (Rapesed Subgroup 20A), DACO: 5.1 2378134 2013, Use Description and Exposure Scenarios for Canola, Corn, and Soybean Seed Treatment wit S-2200 3.2 FS Fungicide, DACO: 5.4 2013, Use Description and Exposure Scenarios for Canola, Corn, and Soybean Seed Treatment wit S-2200 3.2 FS Fungicide, DACO: 5.14 2013, Use Description and Exposure Scenarios for Canola, Corn, and Soybean Seed Treatment wit S-2200 3.2 FS Fungicide, DACO: 5.4<	2378129	
2378131 2012, S-2200 3.2 FS: Local Lymph Node Assay (LLNA) in Mice, DACO: 2377872 2013, Summary of Occupational Risk Assessments for the Use of S-2200 4 SC 2377874 2013, Occupational and Residential Exposure and Risk Assessments for S-2200 4 SC Fungicide on Canola (Crop Subgroup 20A), Grapes (Crop Subgroup 13-07F), Strawberries (Crop Subgroup 20A), Grapes (Crop Subgroup 13-07F), Strawberries (Crop Subgroup 13-07G), and Turf, DACO: 2377876 2013, Use Description and Scenario (Mixer/Loader/Applicator and Postapplication) for S-2200 4 SC Fungicide, DACO: 5.2 2377878 2013, Pesticide Handlers Exposure Database Assessment in Support of Spray Applications of S-2200 4 SC Fungicide to Canola, Grapes, Strawberries, and Turfgrass, DACO: 5.3 2377880 2013, Postapplication Assessment of S-2200 4 SC Fungicide Following Treatment to Canola, Grapes, Strawberries, and Turf, DACO: 5.6 2377882 2013, Waiver Request: Foliar Dislodgeable and Turf Transferable Residue Study for S-2200 4 SC Ag and VPP Fungicides, DACO: 5.9(A), 5.9(B) 2378132 2013, Summary of Occupational Risk Assessments for the Seed Treatment Use of S-2200 3.2 FS Fungicide Seed Treatment of Corn, Legume Vegetables (Succulent or Dried) Except Cowpea and Field Pea (Crop Group 6 except Cowpea and Field Pea), Sorghum, and Canola (Rapeseed Subgroup 20A), DACO: 5.1 2378135 2013, Use Description and Exposure Scenarios for Canola, Corn, and Soybean Seed Treatment with S-2200 3.2 FS Fungicide, DACO: 5.2 2378136 2013, Mixer/Loader/Applicator Passive Dosimetry Study in Sup		
2378131 4.6.6 2377872 2013, Summary of Occupational Risk Assessments for the Use of S-2200 4 SC Fungicide on Canola, Grapes, Strawberries and Turfgrass, DACO: 5.1 2377874 2013, Occupational and Residential Exposure and Risk Assessments for S- 2200 4 SC Fungicide on Canola (Crop Subgroup 20A), Grapes (Crop Subgroup 13-07F), Strawberries (Crop Subgroup 20A), Grapes (Crop Subgroup 13-07F), Strawberries (Crop Subgroup 13-07G), and Turf, DACO: 5.1 2377876 2013, Use Description and Scenario (Mixer/Loader/Applicator and PostapPlication) for S-2200 4 SC Fungicide, DACO: 5.2 2013, Pesticide Handlers Exposure Database Assessment in Support of Spray Applications of S-2200 4 SC Fungicide to Canola, Grapes, Strawberries, and Turfgrass, DACO: 5.3 2377880 2013, PostapPlication Assessment of S-2200 4 SC Fungicide Following Treatment to Canola, Grapes, Strawberries, and Turf, DACO: 5.6 2377882 2013, Waiver Request: Foliar Dislodgeable and Turf Transferable Residue Study for S-2200 3.2 FS Fungicide on Canola, Corn, and Soybean, DACO: 5.1 2378132 2013, Summary of Occupational Risk Assessments for S-2200 3.2 FS Fungicide Seed Treatment of Cron, Legume Vegetables (Succulent or Dried) Except Cowpea and Field Pea (Crop Group 6 except Cowpea and Field Pea), Sorghum, and Canola (Rapeseed Subgroup 20A), DACO: 5.1 2378135 2013, Use Description and Exposure Scenarios for Canola, Corn, and Soybean Seed Treatment with S-2200 3.2 FS Fungicide, DACO: 5.2 2378136 2013, Juse Description and Exposure Scenarios for Canola, Corn, and Soybean Seed Treatment		
237/872 Fungicide on Canola, Grapes, Strawberries and Turfgrass, DACO: 5.1 2377874 2013, Occupational and Residential Exposure and Risk Assessments for S- 2200 4 SC Fungicide on Canola (Crop Subgroup 20A), Grapes (Crop Subgroup 13-07F), Strawberries (Crop Subgroup 13-07G), and Turf, DACO: 5.1 2377876 2013, Use Description and Scenario (Mixer/Loader/Applicator and Postapplication) for S-2200 4 SC Fungicide, DACO: 5.2 2013, Pesticide Handlers Exposure Database Assessment in Support of Spray Applications of S-2200 4 SC Fungicide to Canola, Grapes, Strawberries, and Turfgrass, DACO: 5.3 2377880 2013, Postapplication Assessment of S-2200 4 SC Fungicide Following Treatment to Canola, Grapes, Strawberries, and Turf, DACO: 5.6 2377882 2013, Waiver Request: Foliar Dislodgeable and Turf Transferable Residue Study for S-2200 4 SC Ag and VPP Fungicides, DACO: 5.9(A), 5.9(B) 2378132 2013, Occupational Exposure and Risk Assessments for the Seed Treatment Use of S-2200 3.2 FS Fungicide on Canola, Corn, and Soybean, DACO: 5.1 2378133 2013, Outpational Exposure and Risk Assessments for S-2200 3.2 FS Fungicide Seed Treatment of Corn, Legume Vegetables (Succulent or Dried) Except Cowpea and Field Pea (Crop Group 6 except Cowpea and Field Pea), Sorghum, and Canola (Rapeseed Subgroup 20A), DACO: 5.1 2378134 2013, Dust Off Study in Support of the Seed Treatment Use of S-2200 3.2 FS Fungicide, DACO: 5.14 2378135 2013, Mixer/Loader/Applicator Passive Dosimetry Study in Support of Commercial Seed Treatment of Canola, Corn, and Soybean with S-220	2378131	4.6.6
2377874 2200 4 SC Fungicide on Canola (Crop Subgroup 20A), Grapes (Crop Subgroup 13-07F), Strawberries (Crop Subgroup 13-07G), and Turf, DACO: 5.1 2377876 2013, Use Description and Scenario (Mixer/Loader/Applicator and Postapplication) for S-2200 4 SC Fungicide, DACO: 5.2 2377878 2013, Pesticide Handlers Exposure Database Assessment in Support of Spray Applications of S-2200 4 SC Fungicide to Canola, Grapes, Strawberries, and Turfgrass, DACO: 5.3 2377880 2013, Postapplication Assessment of S-2200 4 SC Fungicide Following Treatment to Canola, Grapes, Strawberries, and Turf, DACO: 5.6 2377880 2013, Waiver Request: Foliar Dislodgeable and Turf Transferable Residue Study for S-2200 4 SC Ag and VPP Fungicides, DACO: 5.9(A), 5.9(B) 2378132 2013, Summary of Occupational Risk Assessments for the Seed Treatment Use of S-2200 3.2 FS Fungicide on Canola, Corn, and Soybean, DACO: 5.1 2378133 2013, Occupational Exposure and Risk Assessments for S-2200 3.2 FS Fungicide, DACO: 5.14 2013, Dust Off Study in Support of the Seed Treatment Use of S-2200 3.2 FS 2378134 2013, Use Description and Exposure Scenarios for Canola, Corn, and Soybean Seed Treatment with S-2200 3.2 FS Fungicide, DACO: 5.14 2378135 2013, Use Description and Exposure Scenarios for Canola, Corn, and Soybean Seed Treatment with S-2200 3.2 FS Fungicide, DACO: 5.4 2378136 2013, Use Description and Exposure Scenarios for Canola, Corn, and Soybean Seed Treatment with S-2200 3.2 FS Fungicide, DACO: 5.4 2378	2377872	
2317876 Postapplication) for S-2200 4 SC Fungicide, DACO: 5.2 2013, Pesticide Handlers Exposure Database Assessment in Support of Spray Applications of S-2200 4 SC Fungicide to Canola, Grapes, Strawberries, and Turfgrass, DACO: 5.3 2377880 2013, Postapplication Assessment of S-2200 4 SC Fungicide Following Treatment to Canola, Grapes, Strawberries, and Turf, DACO: 5.6 2377882 2013, Waiver Request: Foliar Dislodgeable and Turf Transferable Residue Study for S-2200 4 SC Ag and VPP Fungicides, DACO: 5.9(A), 5.9(B) 2378132 2013, Summary of Occupational Risk Assessments for the Seed Treatment Use of S-2200 3.2 FS Fungicide on Canola, Corn, and Soybean, DACO: 5.1 2013, Occupational Exposure and Risk Assessments for S-2200 3.2 FS Fungicide Seed Treatment of Corn, Legume Vegetables (Succulent or Dried) Except Cowpea and Field Pea (Crop Group 6 except Cowpea and Field Pea), Sorghum, and Canola (Rapeseed Subgroup 20A), DACO: 5.1 2378134 2013, Use Description and Exposure Scenarios for Canola, Corn, and Soybean Seed Treatment with S-2200 3.2 FS Fungicide, DACO: 5.2 2013, Mixer/Loader/Applicator Passive Dosimetry Study in Support of Commercial Seed Treatment of Canola, Corn, and Soybean with S-2200 3.2 FS Fungicide, DACO: 5.4 2378137 2013, Mixer/Loader/Applicator Passive Dosimetry Study in Support of On- Farm Seed Treatment of Soybean with S-2200 3.2 FS Fungicide, DACO: 5.4 2378137 2013, Mixer/Loader/Applicator Passive Dosimetry Study in Support of On- Farm Seed Treatment of Soybean Treated with S-2200 3.2 FS Fungicide, DACO: 5.6	2377874	2200 4 SC Fungicide on Canola (Crop Subgroup 20A), Grapes (Crop Subgroup 13-07F), Strawberries (Crop Subgroup 13-07G), and Turf, DACO:
2377878Applications of S-2200 4 SC Fungicide to Canola, Grapes, Strawberries, and Turfgrass, DACO: 5.323778802013, Postapplication Assessment of S-2200 4 SC Fungicide Following Treatment to Canola, Grapes, Strawberries, and Turf, DACO: 5.623778822013, Waiver Request: Foliar Dislodgeable and Turf Transferable Residue Study for S-2200 4 SC Ag and VPP Fungicides, DACO: 5.9(A), 5.9(B)23781322013, Summary of Occupational Risk Assessments for the Seed Treatment Use of S-2200 3.2 FS Fungicide on Canola, Corn, and Soybean, DACO: 5.123781332013, Occupational Exposure and Risk Assessments for S-2200 3.2 FS Fungicide Seed Treatment of Corn, Legume Vegetables (Succulent or Dried) Except Cowpea and Field Pea (Crop Group 6 except Cowpea and Field Pea), Sorghum, and Canola (Rapeseed Subgroup 20A), DACO: 5.123781342013, Dust Off Study in Support of the Seed Treatment Use of S-2200 3.2 FS Fungicide, DACO: 5.1423781352013, Use Description and Exposure Scenarios for Canola, Corn, and Soybean Seed Treatment with S-2200 3.2 FS Fungicide, DACO: 5.223781362013, Mixer/Loader/Applicator Passive Dosimetry Study in Support of Commercial Seed Treatment of Canola, Corn, and Soybean with S-2200 3.2 FS Fungicide, DACO: 5.423781372013, Mixer/Loader/Applicator Passive Dosimetry Study in Support of On- Farm Seed Treatment of Soybean with S-2200 3.2 FS Fungicide, DACO: 5.423781382013, Postapplication: Seed Planter Passive Dosimetry Study in Support of Planting Canola, Corn, and Soybean Treated with S-2200 3.2 FS Fungicide, DACO: 5.423781372012, [14C]-S-2200: Absorption, Distribution, Metabolism and Excretion Following Repeated Oral Administration to the Laying Hen, DACO 6.2237783	2377876	
2377880Treatment to Canola, Grapes, Strawberries, and Turf, DACO: 5.623778822013, Waiver Request: Foliar Dislodgeable and Turf Transferable Residue Study for S-2200 4 SC Ag and VPP Fungicides, DACO: 5.9(A), 5.9(B)23781322013, Summary of Occupational Risk Assessments for the Seed Treatment Use of S-2200 3.2 FS Fungicide on Canola, Corn, and Soybean, DACO: 5.12013, Occupational Exposure and Risk Assessments for S-2200 3.2 FS Fungicide Seed Treatment of Corn, Legume Vegetables (Succulent or Dried) Except Cowpea and Field Pea (Crop Group 6 except Cowpea and Field Pea), Sorghum, and Canola (Rapeseed Subgroup 20A), DACO: 5.123781342013, Dust Off Study in Support of the Seed Treatment Use of S-2200 3.2 FS Fungicide, DACO: 5.1423781352013, Use Description and Exposure Scenarios for Canola, Corn, and Soybean Seed Treatment with S-2200 3.2 FS Fungicide, DACO: 5.223781362013, Mixer/Loader/Applicator Passive Dosimetry Study in Support of Commercial Seed Treatment of Canola, Corn, and Soybean with S-2200 3.2 FS Fungicide, DACO: 5.423781372013, Mixer/Loader/Applicator Passive Dosimetry Study in Support of On- Farm Seed Treatment of Soybean with S-2200 3.2 FS Fungicide, DACO: 5.423781382013, Postapplication: Seed Planter Passive Dosimetry Study in Support of Planting Canola, Corn, and Soybean Treated with S-2200 3.2 FS Fungicide, DACO: 5.623778362012, [14C]-S-2200: Absorption, Distribution, Metabolism and Excretion Following Repeated Oral Administration to the Laying Hen, DACO 6.223778382012, [14C]-S-2200: Absorption, Distribution, Metabolism and Excretion Following Repeated Oral Administration to the Laying Hen, DACO 6.2	2377878	Applications of S-2200 4 SC Fungicide to Canola, Grapes, Strawberries, and
23778822013, Waiver Request: Foliar Dislodgeable and Turf Transferable Residue Study for S-2200 4 SC Ag and VPP Fungicides, DACO: 5.9(A), 5.9(B)23781322013, Summary of Occupational Risk Assessments for the Seed Treatment Use of S-2200 3.2 FS Fungicide on Canola, Corn, and Soybean, DACO: 5.123781332013, Occupational Exposure and Risk Assessments for S-2200 3.2 FS Fungicide Seed Treatment of Corn, Legume Vegetables (Succulent or Dried) Except Cowpea and Field Pea (Crop Group 6 except Cowpea and Field Pea), Sorghum, and Canola (Rapeseed Subgroup 20A), DACO: 5.123781342013, Dust Off Study in Support of the Seed Treatment Use of S-2200 3.2 FS Fungicide, DACO: 5.1423781352013, Use Description and Exposure Scenarios for Canola, Corn, and Soybean Seed Treatment with S-2200 3.2 FS Fungicide, DACO: 5.223781362013, Mixer/Loader/Applicator Passive Dosimetry Study in Support of Commercial Seed Treatment of Canola, Corn, and Soybean with S-2200 3.2 FS Fungicide, DACO: 5.423781372013, Mixer/Loader/Applicator Passive Dosimetry Study in Support of On- Farm Seed Treatment of Soybean with S-2200 3.2 FS Fungicide, DACO: 5.423781382013, Postapplication: Seed Planter Passive Dosimetry Study in Support of Planting Canola, Corn, and Soybean Treated with S-2200 3.2 FS Fungicide, DACO: 5.623778362012, [14C]-S-2200: Absorption, Distribution, Metabolism and Excretion Following Repeated Oral Administration to the Lactating Ruminant, DACO 6.223778382012, [14C]-S-2200: Absorption, Distribution, Metabolism and Excretion Following Repeated Oral Administration to the Laying Hen, DACO 6.2	2377880	
23781322013, Summary of Occupational Risk Assessments for the Seed Treatment Use of S-2200 3.2 FS Fungicide on Canola, Corn, and Soybean, DACO: 5.123781332013, Occupational Exposure and Risk Assessments for S-2200 3.2 FS Fungicide Seed Treatment of Corn, Legume Vegetables (Succulent or Dried) Except Cowpea and Field Pea (Crop Group 6 except Cowpea and Field Pea), Sorghum, and Canola (Rapeseed Subgroup 20A), DACO: 5.123781342013, Dust Off Study in Support of the Seed Treatment Use of S-2200 3.2 FS Fungicide, DACO: 5.1423781352013, Use Description and Exposure Scenarios for Canola, Corn, and Soybean Seed Treatment with S-2200 3.2 FS Fungicide, DACO: 5.223781362013, Mixer/Loader/Applicator Passive Dosimetry Study in Support of Commercial Seed Treatment of Canola, Corn, and Soybean with S-2200 3.2 FS Fungicide, DACO: 5.423781372013, Mixer/Loader/Applicator Passive Dosimetry Study in Support of On- Farm Seed Treatment of Soybean with S-2200 3.2 FS Fungicide, DACO: 5.423781382013, Mixer/Loader/Applicator Passive Dosimetry Study in Support of On- Farm Seed Treatment of Soybean with S-2200 3.2 FS Fungicide, DACO: 5.423781382013, Postapplication: Seed Planter Passive Dosimetry Study in Support of Planting Canola, Corn, and Soybean Treated with S-2200 3.2 FS Fungicide, DACO: 5.623778362012, [14C]-S-2200: Absorption, Distribution, Metabolism and Excretion Following Repeated Oral Administration to the Lactating Ruminant, DACO 6.223778382012, [14C]-S-2200: Absorption, Distribution, Metabolism and Excretion Following Repeated Oral Administration to the Laying Hen, DACO 6.2	2377882	2013, Waiver Request: Foliar Dislodgeable and Turf Transferable Residue
23781332013, Occupational Exposure and Risk Assessments for S-2200 3.2 FS Fungicide Seed Treatment of Corn, Legume Vegetables (Succulent or Dried) Except Cowpea and Field Pea (Crop Group 6 except Cowpea and Field Pea), Sorghum, and Canola (Rapeseed Subgroup 20A), DACO: 5.123781342013, Dust Off Study in Support of the Seed Treatment Use of S-2200 3.2 FS Fungicide, DACO: 5.1423781352013, Use Description and Exposure Scenarios for Canola, Corn, and Soybean Seed Treatment with S-2200 3.2 FS Fungicide, DACO: 5.223781362013, Mixer/Loader/Applicator Passive Dosimetry Study in Support of Commercial Seed Treatment of Canola, Corn, and Soybean with S-2200 3.2 FS Fungicide, DACO: 5.423781372013, Mixer/Loader/Applicator Passive Dosimetry Study in Support of On- Farm Seed Treatment of Soybean with S-2200 3.2 FS Fungicide, DACO: 5.423781382013, Postapplication: Seed Planter Passive Dosimetry Study in Support of Planting Canola, Corn, and Soybean Treated with S-2200 3.2 FS Fungicide, DACO: 5.623778362012, [14C]-S-2200: Absorption, Distribution, Metabolism and Excretion Following Repeated Oral Administration to the Laying Hen, DACO 6.223778382012, [14C]-S-2200: Absorption, Distribution, Metabolism and Excretion Following Repeated Oral Administration to the Laying Hen, DACO 6.2	2378132	
23781342013, Dust Off Study in Support of the Seed Treatment Use of S-2200 3.2 FS Fungicide, DACO: 5.1423781352013, Use Description and Exposure Scenarios for Canola, Corn, and Soybean Seed Treatment with S-2200 3.2 FS Fungicide, DACO: 5.22013, Mixer/Loader/Applicator Passive Dosimetry Study in Support of Commercial Seed Treatment of Canola, Corn, and Soybean with S-2200 3.2 FS Fungicide, DACO: 5.423781362013, Mixer/Loader/Applicator Passive Dosimetry Study in Support of Commercial Seed Treatment of Canola, Corn, and Soybean with S-2200 3.2 FS Fungicide, DACO: 5.423781372013, Mixer/Loader/Applicator Passive Dosimetry Study in Support of On- Farm Seed Treatment of Soybean with S-2200 3.2 FS Fungicide, DACO: 5.423781382013, Postapplication: Seed Planter Passive Dosimetry Study in Support of Planting Canola, Corn, and Soybean Treated with S-2200 3.2 FS Fungicide, DACO: 5.623778362012, [14C]-S-2200: Absorption, Distribution, Metabolism and Excretion Following Repeated Oral Administration to the Lactating Ruminant, DACO 6.223778382012, [14C]-S-2200: Absorption, Distribution, Metabolism and Excretion Following Repeated Oral Administration to the Laying Hen, DACO 6.2	2378133	2013, Occupational Exposure and Risk Assessments for S-2200 3.2 FS Fungicide Seed Treatment of Corn, Legume Vegetables (Succulent or Dried) Except Cowpea and Field Pea (Crop Group 6 except Cowpea and Field Pea),
23781352013, Use Description and Exposure Scenarios for Canola, Corn, and Soybean Seed Treatment with S-2200 3.2 FS Fungicide, DACO: 5.223781362013, Mixer/Loader/Applicator Passive Dosimetry Study in Support of Commercial Seed Treatment of Canola, Corn, and Soybean with S-2200 3.2 FS Fungicide, DACO: 5.423781372013, Mixer/Loader/Applicator Passive Dosimetry Study in Support of On- Farm Seed Treatment of Soybean with S-2200 3.2 FS Fungicide, DACO: 5.423781382013, Mixer/Loader/Applicator Passive Dosimetry Study in Support of On- Farm Seed Treatment of Soybean with S-2200 3.2 FS Fungicide, DACO: 5.423781382013, Postapplication: Seed Planter Passive Dosimetry Study in Support of Planting Canola, Corn, and Soybean Treated with S-2200 3.2 FS Fungicide, 	2378134	2013, Dust Off Study in Support of the Seed Treatment Use of S-2200 3.2 FS
2378136Commercial Seed Treatment of Canola, Corn, and Soybean with S-2200 3.2 FS Fungicide, DACO: 5.423781372013, Mixer/Loader/Applicator Passive Dosimetry Study in Support of On- Farm Seed Treatment of Soybean with S-2200 3.2 FS Fungicide, DACO: 5.423781382013, Postapplication: Seed Planter Passive Dosimetry Study in Support of Planting Canola, Corn, and Soybean Treated with S-2200 3.2 FS Fungicide, DACO: 5.623778362012, [14C]-S-2200: Absorption, Distribution, Metabolism and Excretion Following Repeated Oral Administration to the Lactating Ruminant, DACO 6.223778382012, [14C]-S-2200: Absorption, Distribution, Metabolism and Excretion Following Repeated Oral Administration to the Laying Hen, DACO 6.2	2378135	2013, Use Description and Exposure Scenarios for Canola, Corn, and Soybean
23781372013, Mixer/Loader/Applicator Passive Dosimetry Study in Support of On- Farm Seed Treatment of Soybean with S-2200 3.2 FS Fungicide, DACO: 5.42013, Postapplication: Seed Planter Passive Dosimetry Study in Support of Planting Canola, Corn, and Soybean Treated with S-2200 3.2 FS Fungicide, DACO: 5.62378362012, [14C]-S-2200: Absorption, Distribution, Metabolism and Excretion Following Repeated Oral Administration to the Lactating Ruminant, DACO 6.223778382012, [14C]-S-2200: Absorption, Distribution, Metabolism and Excretion Following Repeated Oral Administration to the Laying Hen, DACO 6.2	2378136	Commercial Seed Treatment of Canola, Corn, and Soybean with S-2200 3.2
2013, Postapplication: Seed Planter Passive Dosimetry Study in Support of Planting Canola, Corn, and Soybean Treated with S-2200 3.2 FS Fungicide, DACO: 5.623778362012, [14C]-S-2200: Absorption, Distribution, Metabolism and Excretion Following Repeated Oral Administration to the Lactating Ruminant, DACO 6.223778382012, [14C]-S-2200: Absorption, Distribution, Metabolism and Excretion Following Repeated Oral Administration to the Lactating Ruminant, DACO 6.223778382012, [14C]-S-2200: Absorption, Distribution, Metabolism and Excretion Following Repeated Oral Administration to the Laying Hen, DACO 6.2	2378137	
2377836Following Repeated Oral Administration to the Lactating Ruminant, DACO 6.223778382012, [14C]-S-2200: Absorption, Distribution, Metabolism and Excretion Following Repeated Oral Administration to the Laying Hen, DACO 6.2	2378138	2013, Postapplication: Seed Planter Passive Dosimetry Study in Support of Planting Canola, Corn, and Soybean Treated with S-2200 3.2 FS Fungicide,
Following Repeated Oral Administration to the Laying Hen, DACO 6.2	2377836	Following Repeated Oral Administration to the Lactating Ruminant, DACO 6.2
	2377838	•
	2377840	

-	
2377842	2010, Metabolism of [14C]-S-2200 in Wheat, DACO 6.3
2377845	2011, Metabolism of [14C]-S-2200 in Rapeseed Plants, DACO 6.3
2378156	2013, Residues in Crops Grown from Seeds Treated with RS-[Benzyl-14C] S-2200 and RS- [Phenoxy-14C]-S-2200, DACO 7.8
2378157	2013, Characterization of Radioactive Residues of [Benzyl-14C]S-2200 and [Phenoxy-14C]S-2200 in Corn and Soybean Samples from Study VP-37088, DACO 7.8
2377859	2013, S-2200: Independent Laboratory Validation of Valent Method RM- 48M-1 for the Determination of S-2200 in Tissue Samples, DACO 7.2.2, 7.2.3A, 8.2.2.4
2377885	2013, Determination of S-2200 and De-Xy-S-2200 In Crops, DACO 7.2.1
2377887	2013, Radiovalidation of Residue Methods for S-2200 and its Metabolites, DACO 7.2.3B
2377893	2011, Validation of an Analytical Method for Determination of S-2200 Metabolite, De-Xy-S-2200, in Seeds of Oilseed Rape, Barley (grain and straw) and Lettuce (head), DACO 7.2.2,7.2.3A
2377894	2011, Validation of an Analytical Method for Determination of S-2200 Metabolites, 4-OH-S-2200 and its Conjugates, in Seeds of Oilseed Rape, Barley (grain and straw) and Lettuce (head), DACO 7.2.2,7.2.3A
2377896	2011, Validation of an Analytical Method for Determination of S-2200 Metabolites, 2-CH2OH-S- 2200 and its Conjugates, in Seeds of Oilseed Rape, Barley (grain and straw) and Lettuce (head), DACO 7.2.2,7.2.3A
2377898	2011, Validation of an Analytical Method for Determination of S-2200 Metabolites, 5-CH2OH-S- 2200 and its Conjugates, in Barley (grain and straw) and Lettuce (head), DACO 7.2.2,7.2.3A
2377899	2013, Independent Laboratory Validation for the Determination of S-2200 and De-Xy-S-2200 in Grapes and Canola Seed Using LC-MS/MS, DACO 7.2.2, 7.2.3A
2377889	2010, Adaptation and Validation of Multi-Method DFG S 19 for the Determination of Residues of S-2200 in Seeds of Oilseed Rape, DACO 7.2.2, 7.2.3A
2377892	2010, Validation of a Method based on Multi-Method DFG S 19 for the Determination of Residues of S-2200 in High-Water and Dry Crops, DACO 7.2.2, 7.2.3A
2377901	2011, Freezer Storage Stability Study of S-2200 (its Optical Isomers of S-2167 (R-Isomer) and S-2354 (S-Isomer) in Seeds of Oilseed Rape, DACO 7.3
2377902	2011, Freezer Storage Stability Study of S-2200 (its Optical Isomers of S-2167 (R-isomer) and S-2354 (S-Isomer) in/on High-Water and Dry Crops over 12 Months, DACO 7.3
2377904	2012, Freezer Storage Stability Study of S-2200 Metabolite, De-Xy-S2200, in Lettuce (Head), Seeds of Oilseed Rape and Barley (Grain and Straw) over 12 Months, DACO 7.3
2377906	2012, Freezer Storage Stability Study of S-2200 Metabolite, 4-OH-S-2200, in Lettuce (Head), Seeds of Oilseed Rape and Barley (Grain and Straw) over 12 Months, DACO 7.3

2012, Freezer Storage Stability Study of S-2200 Metabolite, 2-CH2OH-S- 2200, in Lettuce (Head), Seeds of Oilseed Rape and Barley (Grain and Straw) over 12 Months, DACO 7.32012, Freezer Storage Stability Study of S-2200 Metabolite, 5-CH2OH-S- 2200, in Lettuce (Head) and Barley (Grain and Straw) over 12 Months, DACO 7.32013, S-2200: Magnitude of the Residue on Strawberries Grown in Canada, DACO 7.4.1, 7.323779112013, S-2200: Magnitude of the Residue on Strawberries, DACO 7.4.1, 7.323779172013, S-2200: Magnitude of the Residue on Grapes Grown in Canada, DACO 7.4.1, 7.323779202013, S-2200: Magnitude of the Residue on Grapes Grown in Canada, DACO 7.4.1, 7.323779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779302013, S-2200: Magnitude of the Residue on Canola Grown in Canada, DACO 7.4.1, 7.323779262013, S-2200: Magnitude of the Residue on Canola Grown in Canada, DACO 7.4.1, 7.323779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523781532013, S-2200: Magnitude of the Residue in Corn Following Seed Treatment at an Exaggerated Rate. DACO 7.4.1, 7.323781522013, Magnitude of the Residue of S-2200 in Soybeans Following Seed Treatment at an Exaggerated Rate, DACO 7.4.123779202013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779202013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779202013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.52377920	-	
2012, Freezer Storage Stability Study of S-2200 Metabolite, 5-CH2OH-S- 2200, in Lettuce (Head) and Barley (Grain and Straw) over 12 Months, DACO 7.323779112013, S-2200: Magnitude of the Residue on Strawberries Grown in Canada, DACO 7.4.1, 7.323779172013, S-2200: Magnitude of the Residue on Strawberries, DACO 7.4.1, 7.323779182013, S-2200: Magnitude of the Residue on Grapes Grown in Canada, DACO 7.4.1, 7.323779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779302013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779202013, S-2200: Magnitude of the Residue on Canola Grown in Canada, DACO 7.4.1, 7.323779262013, S-2200: Magnitude of the Residue on Canola Grown in Canada, DACO 7.4.1, 7.323779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523781522013, Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779342011, Confined Rotational Crop Study with [14C]-S-2200, DACO 7.4.323779382013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Leaf Lettuce, DACO 7.4.423779422013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Canola, DACO 7.4.42013, S-2200 Fungicide: Ann	2377908	2200, in Lettuce (Head), Seeds of Oilseed Rape and Barley (Grain and Straw)
23779092200, in Lettuce (Head) and Barley (Grain and Straw) over 12 Months, DACO 7.323779112013, S-2200: Magnitude of the Residue on Strawberries Grown in Canada, DACO 7.4.1, 7.323779172013, S-2200: Magnitude of the Residue on Strawberries, DACO 7.4.1, 7.323779182013, S-2200: Magnitude of the Residue on Grapes Grown in Canada, DACO 7.4.1, 7.323779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779302013, S-2200: Magnitude of the Residue on Canola Grown in Canada, DACO 7.4.1, 7.323779262013, S-2200: Magnitude of the Residue on Canola Grown in Canada, DACO 7.4.1, 7.323779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523781532013, S-2200: Magnitude of the Residue in Corn Following Seed Treatment at an Exaggerated Rate. DACO 7.4.1, 7.323781522013, Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779342011, Confined Rotational Crop Study with [14C]-S-2200, DACO 7.4.323779382013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Leaf Lettuce, DACO 7.4.423779422013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Canola, DACO 7.4.423779422013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Canola, DACO 7.4.423780812013, S-2200 Fungicide: A		over 12 Months, DACO 7.3
23779092200, in Lettuce (Head) and Barley (Grain and Straw) over 12 Months, DACO 7.323779112013, S-2200: Magnitude of the Residue on Strawberries Grown in Canada, DACO 7.4.1, 7.323779172013, S-2200: Magnitude of the Residue on Strawberries, DACO 7.4.1, 7.323779182013, S-2200: Magnitude of the Residue on Grapes Grown in Canada, DACO 7.4.1, 7.323779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779302013, S-2200: Magnitude of the Residue on Canola Grown in Canada, DACO 7.4.1, 7.323779262013, S-2200: Magnitude of the Residue on Canola Grown in Canada, DACO 7.4.1, 7.323779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523781532013, S-2200: Magnitude of the Residue in Corn Following Seed Treatment at an Exaggerated Rate. DACO 7.4.1, 7.323781522013, Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779342011, Confined Rotational Crop Study with [14C]-S-2200, DACO 7.4.323779382013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Leaf Lettuce, DACO 7.4.423779422013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Canola, DACO 7.4.423779422013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Canola, DACO 7.4.423780812013, S-2200 Fungicide: A		2012, Freezer Storage Stability Study of S-2200 Metabolite, 5-CH2OH-S-
7.323779112013, S-2200: Magnitude of the Residue on Strawberries Grown in Canada, DACO 7.4.1, 7.323779172013, S-2200: Magnitude of the Residue on Strawberries, DACO 7.4.1, 7.323779132013, S-2200: Magnitude of the Residue on Grapes Grown in Canada, DACO 7.4.1, 7.323779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779302013, S-2200: Magnitude of the Residue on Canola Grown in Canada, DACO 7.4.1, 7.323779262013, S-2200: Magnitude of the Residue on Canola Grown in Canada, DACO 7.4.1, 7.323779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523781532013, S-2200: Magnitude of the Residue in Corn Following Seed Treatment at an Exaggerated Rate. DACO 7.4.1, 7.323781522013, Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779342011, Confined Rotational Crop Study with [14C]-S-2200, DACO 7.4.323779382013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Leaf Lettuce, DACO 7.4.423779422013, S-2200 Fungicide: Annex IIA Tier II Summary, Residues in or on Treated Products, Food or Feed (Nature of the Residue, Level of Residue and	2377909	2200, in Lettuce (Head) and Barley (Grain and Straw) over 12 Months, DACO
2377911DACO 7.4.1, 7.323779172013, S-2200: Magnitude of the Residue on Strawberries, DACO 7.4.1, 7.323779132013, S-2200: Magnitude of the Residue on Grapes Grown in Canada, DACO 7.4.1, 7.323779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779302013, S-2200: Magnitude of the Residue on Canola Grown in Canada, DACO 7.4.1, 7.323779262013, S-2200: Magnitude of the Residue on Canola Grown in Canada, DACO 7.4.1, 7.323779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523781532013, S-2200: Magnitude of the Residue in Corn Following Seed Treatment at an Exaggerated Rate. DACO 7.4.1, 7.323781522013, S-2200: Magnitude of the Residue in Corn Following Seed Treatment at an Exaggerated Rate, DACO 7.4.123779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779342011, Confined Rotational Crop Study with [14C]-S-2200, DACO 7.4.323779382013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Canola, DACO 7.4.423779422013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Canola, DACO 7.4.42378081Treated Products, Food or Feed (Nature of the Residue, Level of Residue and		
2377911DACO 7.4.1, 7.323779172013, S-2200: Magnitude of the Residue on Strawberries, DACO 7.4.1, 7.323779132013, S-2200: Magnitude of the Residue on Grapes Grown in Canada, DACO 7.4.1, 7.323779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779302013, S-2200: Magnitude of the Residue on Canola Grown in Canada, DACO 7.4.1, 7.323779262013, S-2200: Magnitude of the Residue on Canola Grown in Canada, DACO 7.4.1, 7.323779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523781532013, S-2200: Magnitude of the Residue in Corn Following Seed Treatment at an Exaggerated Rate. DACO 7.4.1, 7.323781522013, S-2200: Magnitude of the Residue in Corn Following Seed Treatment at an Exaggerated Rate, DACO 7.4.123779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779342011, Confined Rotational Crop Study with [14C]-S-2200, DACO 7.4.323779382013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Canola, DACO 7.4.423779422013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Canola, DACO 7.4.42378081Treated Products, Food or Feed (Nature of the Residue, Level of Residue and	0077011	2013, S-2200: Magnitude of the Residue on Strawberries Grown in Canada,
23779132013, S-2200: Magnitude of the Residue on Grapes Grown in Canada, DACO 7.4.1, 7.323779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779302013, S-2200: Magnitude of the Residue on Canola Grown in Canada, DACO 7.4.1, 7.323779262013, S-2200: Magnitude of the Residue on Canola Grown in Canada, DACO 7.4.1, 7.323779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523781532013, S-2200: Magnitude of the Residue in Corn Following Seed Treatment at an Exaggerated Rate. DACO 7.4.1, 7.323781522013, Magnitude of the Residue of S-2200 in Soybeans Following Seed Treatment at an Exaggerated Rate, DACO 7.4.123779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779342011, Confined Rotational Crop Study with [14C]-S-2200, DACO 7.4.323779382013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Leaf Lettuce, DACO 7.4.423779422013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Canola, DACO 7.4.42378081Treated Products, Food or Feed (Nature of the Residue, Level of Residue and	2377911	
23779137.4.1, 7.323779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779302013, S-2200: Magnitude of the Residue on Canola Grown in Canada, DACO 7.4.1, 7.323779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523781532013. S-2200: Magnitude of the Residue in Corn Following Seed Treatment at an Exaggerated Rate. DACO 7.4.1, 7.323781522013, Magnitude of the Residue of S-2200 in Soybeans Following Seed Treatment at an Exaggerated Rate, DACO 7.4.123779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779202013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779342011, Confined Rotational Crop Study with [14C]-S-2200, DACO 7.4.323779382013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Leaf Lettuce, DACO 7.4.423779422013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Canola, DACO 7.4.423780817reated Products, Food or Feed (Nature of the Residue, Level of Residue and	2377917	2013, S-2200: Magnitude of the Residue on Strawberries, DACO 7.4.1, 7.3
7.4.1, 7.323779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779302013, S-2200: Magnitude of the Residue on Canola Grown in Canada, DACO 7.4.1, 7.323779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523781532013. S-2200: Magnitude of the Residue in Corn Following Seed Treatment at an Exaggerated Rate. DACO 7.4.1, 7.323781522013, Magnitude of the Residue of S-2200 in Soybeans Following Seed Treatment at an Exaggerated Rate, DACO 7.4.123779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.323781522013, Magnitude of the Residue of S-2200 in Soybeans Following Seed Treatment at an Exaggerated Rate, DACO 7.4.123779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779342011, Confined Rotational Crop Study with [14C]-S-2200, DACO 7.4.323779382013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Leaf Lettuce, DACO 7.4.423779422013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Canola, DACO 7.4.423780817013, S-2200 Fungicide: Annex IIA Tier II Summary, Residues in or on Treated Products, Food or Feed (Nature of the Residue, Level of Residue and	0077010	2013, S-2200: Magnitude of the Residue on Grapes Grown in Canada, DACO
23779302013, S-2200: Magnitude of the Residue on Canola Grown in Canada, DACO 7.4.1, 7.323779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523781532013. S-2200: Magnitude of the Residue in Corn Following Seed Treatment at an Exaggerated Rate. DACO 7.4.1, 7.323781522013, Magnitude of the Residue of S-2200 in Soybeans Following Seed Treatment at an Exaggerated Rate, DACO 7.4.123779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779342011, Confined Rotational Crop Study with [14C]-S-2200, DACO 7.4.323779382013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Leaf Lettuce, DACO 7.4.423779422013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Canola, DACO 7.4.423780817reated Products, Food or Feed (Nature of the Residue, Level of Residue and	2377913	7.4.1, 7.3
23779302013, S-2200: Magnitude of the Residue on Canola Grown in Canada, DACO 7.4.1, 7.323779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523781532013. S-2200: Magnitude of the Residue in Corn Following Seed Treatment at an Exaggerated Rate. DACO 7.4.1, 7.323781522013, Magnitude of the Residue of S-2200 in Soybeans Following Seed Treatment at an Exaggerated Rate, DACO 7.4.123779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779342011, Confined Rotational Crop Study with [14C]-S-2200, DACO 7.4.323779382013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Leaf Lettuce, DACO 7.4.423779422013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Canola, DACO 7.4.423780817reated Products, Food or Feed (Nature of the Residue, Level of Residue and	2377920	2013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.5
23779307.4.1, 7.323779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523781532013. S-2200: Magnitude of the Residue in Corn Following Seed Treatment at an Exaggerated Rate. DACO 7.4.1, 7.323781522013, Magnitude of the Residue of S-2200 in Soybeans Following Seed Treatment at an Exaggerated Rate, DACO 7.4.123779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779262013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779342011, Confined Rotational Crop Study with [14C]-S-2200, DACO 7.4.323779382013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Leaf Lettuce, DACO 7.4.423779422013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Canola, DACO 7.4.423780812013, S-2200 Fungicide: Annex IIA Tier II Summary, Residues in or on Treated Products, Food or Feed (Nature of the Residue, Level of Residue and	2277020	
2377926States, DACO 7.4.1, 7.3, 7.4.523781532013. S-2200: Magnitude of the Residue in Corn Following Seed Treatment at an Exaggerated Rate. DACO 7.4.1, 7.323781522013, Magnitude of the Residue of S-2200 in Soybeans Following Seed Treatment at an Exaggerated Rate, DACO 7.4.123779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779342011, Confined Rotational Crop Study with [14C]-S-2200, DACO 7.4.323779382013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Leaf Lettuce, DACO 7.4.423779422013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Canola, DACO 7.4.42378081Treated Products, Food or Feed (Nature of the Residue, Level of Residue and	2377930	-
States, DACO 7.4.1, 7.3, 7.4.523781532013. S-2200: Magnitude of the Residue in Corn Following Seed Treatment at an Exaggerated Rate. DACO 7.4.1, 7.323781522013, Magnitude of the Residue of S-2200 in Soybeans Following Seed Treatment at an Exaggerated Rate, DACO 7.4.123779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779342011, Confined Rotational Crop Study with [14C]-S-2200, DACO 7.4.323779382013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Leaf Lettuce, DACO 7.4.423779422013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Canola, DACO 7.4.42378081Treated Products, Food or Feed (Nature of the Residue, Level of Residue and	2277026	2013, S-2200: Magnitude of the Residue on Canola Grown in the United
2378153an Exaggerated Rate. DACO 7.4.1, 7.323781522013, Magnitude of the Residue of S-2200 in Soybeans Following Seed Treatment at an Exaggerated Rate, DACO 7.4.123779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779342011, Confined Rotational Crop Study with [14C]-S-2200, DACO 7.4.323779382013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Leaf Lettuce, DACO 7.4.423779422013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Canola, DACO 7.4.423780812013, S-2200 Fungicide: Annex IIA Tier II Summary, Residues in or on Treated Products, Food or Feed (Nature of the Residue, Level of Residue and	2377920	States, DACO 7.4.1, 7.3, 7.4.5
an Exaggerated Rate. DACO 7.4.1, 7.323781522013, Magnitude of the Residue of S-2200 in Soybeans Following Seed Treatment at an Exaggerated Rate, DACO 7.4.123779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779342011, Confined Rotational Crop Study with [14C]-S-2200, DACO 7.4.323779382013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Leaf Lettuce, DACO 7.4.423779422013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Canola, DACO 7.4.423780812013, S-2200 Fungicide: Annex IIA Tier II Summary, Residues in or on Treated Products, Food or Feed (Nature of the Residue, Level of Residue and	0270152	2013. S-2200: Magnitude of the Residue in Corn Following Seed Treatment at
2378152Treatment at an Exaggerated Rate, DACO 7.4.123779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779342011, Confined Rotational Crop Study with [14C]-S-2200, DACO 7.4.323779382013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Leaf Lettuce, DACO 7.4.423779422013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Canola, DACO 7.4.423779422013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Canola, DACO 7.4.42378081Treated Products, Food or Feed (Nature of the Residue, Level of Residue and	2378133	an Exaggerated Rate. DACO 7.4.1, 7.3
Treatment at an Exaggerated Rate, DACO 7.4.123779262013, S-2200: Magnitude of the Residue on Canola Grown in the United States, DACO 7.4.1, 7.3, 7.4.523779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779342011, Confined Rotational Crop Study with [14C]-S-2200, DACO 7.4.323779382013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Leaf Lettuce, DACO 7.4.423779422013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Canola, DACO 7.4.423780812013, S-2200 Fungicide: Annex IIA Tier II Summary, Residues in or on Treated Products, Food or Feed (Nature of the Residue, Level of Residue and	2278152	2013, Magnitude of the Residue of S-2200 in Soybeans Following Seed
2377920States, DACO 7.4.1, 7.3, 7.4.523779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779342011, Confined Rotational Crop Study with [14C]-S-2200, DACO 7.4.323779382013, S-2200: Residues in Rotational Crops Following Application of S-22004 SC to Leaf Lettuce, DACO 7.4.42013, S-2200: Residues in Rotational Crops Following Application of S-22004 SC to Canola, DACO 7.4.42013, S-2200: Residues in Rotational Crops Following Application of S-22002378081Treated Products, Food or Feed (Nature of the Residue, Level of Residue and	2378132	Treatment at an Exaggerated Rate, DACO 7.4.1
States, DACO 7.4.1, 7.3, 7.4.523779202013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.523779342011, Confined Rotational Crop Study with [14C]-S-2200, DACO 7.4.323779382013, S-2200: Residues in Rotational Crops Following Application of S-22004 SC to Leaf Lettuce, DACO 7.4.423779422013, S-2200: Residues in Rotational Crops Following Application of S-22004 SC to Canola, DACO 7.4.423780812013, S-2200 Fungicide: Annex IIA Tier II Summary, Residues in or on Treated Products, Food or Feed (Nature of the Residue, Level of Residue and	2277026	2013, S-2200: Magnitude of the Residue on Canola Grown in the United
23779342011, Confined Rotational Crop Study with [14C]-S-2200, DACO 7.4.323779382013, S-2200: Residues in Rotational Crops Following Application of S-22004 SC to Leaf Lettuce, DACO 7.4.423779422013, S-2200: Residues in Rotational Crops Following Application of S-22004 SC to Canola, DACO 7.4.42013, S-2200 Fungicide: Annex IIA Tier II Summary, Residues in or on2378081Treated Products, Food or Feed (Nature of the Residue, Level of Residue and	2377920	
23779382013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Leaf Lettuce, DACO 7.4.423779422013, S-2200: Residues in Rotational Crops Following Application of S-2200 4 SC to Canola, DACO 7.4.423780812013, S-2200 Fungicide: Annex IIA Tier II Summary, Residues in or on Treated Products, Food or Feed (Nature of the Residue, Level of Residue and	2377920	2013, S-2200: Magnitude of the Residue on Grapes, DACO 7.4.1, 7.3, 7.4.5
23779384 SC to Leaf Lettuce, DACO 7.4.423779422013, S-2200: Residues in Rotational Crops Following Application of S-22004 SC to Canola, DACO 7.4.42013, S-2200 Fungicide: Annex IIA Tier II Summary, Residues in or on Treated Products, Food or Feed (Nature of the Residue, Level of Residue and	2377934	
4 SC to Leaf Lettuce, DACO 7.4.423779422013, S-2200: Residues in Rotational Crops Following Application of S-22004 SC to Canola, DACO 7.4.42013, S-2200 Fungicide: Annex IIA Tier II Summary, Residues in or on2378081Treated Products, Food or Feed (Nature of the Residue, Level of Residue and	2377938	2013, S-2200: Residues in Rotational Crops Following Application of S-2200
23779424 SC to Canola, DACO 7.4.42013, S-2200 Fungicide: Annex IIA Tier II Summary, Residues in or on Treated Products, Food or Feed (Nature of the Residue, Level of Residue and		4 SC to Leaf Lettuce, DACO 7.4.4
4 SC to Canola, DACO 7.4.42013, S-2200 Fungicide: Annex IIA Tier II Summary, Residues in or on2378081Treated Products, Food or Feed (Nature of the Residue, Level of Residue and	2377942	2013, S-2200: Residues in Rotational Crops Following Application of S-2200
2378081 Treated Products, Food or Feed (Nature of the Residue, Level of Residue and		,
	2378081	
Consumer Risk Assessment), DACO M12.7, 6.1, 7.1		Consumer Risk Assessment), DACO M12.7, 6.1, 7.1

3.0 Environment

PMRA Document	
Number	Reference
2377847	2011, S-2200 and Metabolites: Validation of an Analytical Method for
	Residues in EU Soil, DACO: 8.2.2.1, 8.2.2.2
2377849	2013, S-2200: Independent Laboratory Validation of Valent Method RM-
	48S-3, Determination of S-2200 and Metabolites in Soil, DACO: 8.2.2.1,
	8.2.2.2
2377852	2013, Analytical Method Verification for the Determination of S-2200 in
	Sediment, DACO: 8.2.2.2

2377854	2011, Validation of an Analytical Method for the Determination of S-2200 in Surface Water for Post-Registration Control and Monitoring Purpose, DACO: 8.2.2.3
2377855	2012, Analytical Method Verification for the Determination of S-2200 in Freshwater and Saltwater, DACO: 8.2.2.3
2377857	2013, S-2200: Independent Laboratory Validation of the Analytical Method Validation of an Analytical Method for the Determination of S-2200 in Surface Water for Post Registration Control and Monitoring Purposes, DACO: 8.2.2.3
2377861	2010, [¹⁴ C]S-2167 (S-2200 <i>R</i> -isomer): Hydrolytic Stability, DACO: 8.2.3.2
2377863	2010, [¹⁴ C]S-2354 (S-2200 S-isomer): Hydrolytic Stability, DACO: 8.2.3.2
2377864	2010, [¹⁴ C]S-2354 (S-2200 S-isomer): Photodegradation on a Soil Surface, DACO: 8.2.3.3.1
2377867	2011, [¹⁴ C]S-2167 (S-2200 <i>R</i> -isomer): Photodegradation on a Soil Surface, DACO: 8.2.3.3.1
2377869	2010, [¹⁴ C]S-2354 (S-2200 <i>S</i> -isomer): Photodegradation and Quantum Yield in Sterile, Aqueous Solution, DACO: 8.2.3.3.2
2377871	2010, [¹⁴ C]S-2167 (S-2200 R-isomer): Photodegradation and Quantum Yield in Sterile, Aqueous Solution, DACO: 8.2.3.3.2
2377873	2013, S-2200: Waiver for Photodegradation in Air, DACO: 8.2.3.3.3
2377877	2010, [¹⁴ C]2-COOH-S-2200: Aerobic Degradation in Three Soils, DACO: 8.2.3.4.2
2377879	2010, [¹⁴ C]5-COOH-S-2200: Aerobic Degradation in Three Soils, DACO: 8.2.3.4.2
2377881	2011, [¹⁴ C]S-2167 (S-2200 <i>R</i> -isomer): Aerobic Soil Metabolism and Degradation, DACO: 8.2.3.4.2
2377883	2011, [¹⁴ C]S-2354 (S-2200 <i>S</i> -isomer): Aerobic Soil Metabolism and Degradation, DACO: 8.2.3.4.2
2377886	2011, [¹⁴ C]S-2167 (S-2200 <i>R</i> -isomer): Aerobic Degradation in Two Soils, DACO: 8.2.3.4.2
2377888	2011, [¹⁴ C]S-2354 (S-2200 <i>S</i> -isomer): Aerobic Soil Degradation in Two Soils, DACO: 8.2.3.4.2
2377890	2013, S-2200: Degradation Under Aerobic Conditions in Soil, DACO: 8.2.3.4.2
2377894	2013, S-2200: Degradation Under Aerobic Conditions in Soil - Rate Studies, DACO: 8.2.3.4.2
2377897	2013, S-2200: Anaerobic Soil Metabolism Study, DACO: 8.2.3.4.4
2377900	2013, S-2200: Degradation Under Anaerobic Conditions in Soil - Rate Studies, DACO: 8.2.3.4.4
2377903	2011, [¹⁴ C]S-2167 (S-2200 <i>R</i> -Isomer): Degradation in Water-Sediment Systems under Aerobic Conditions, DACO: 8.2.3.5.4
2377905	2011, [¹⁴ C]S-2354 (S-2200 <i>S</i> -Isomer): Degradation in Water-Sediment Systems under Aerobic Conditions, DACO: 8.2.3.5.4
2377907	2013, S-2200: Degradation under Anaerobic Aquatic Conditions, DACO: 8.2.3.5.6

	14
2377910	2010, [¹⁴ C]S-2200: Adsorption/Desorption in Five Soils, DACO: 8.2.4.2
2377912	2010, [¹⁴ C]2-COOH-S-2200:Adsorption/Desorption in Three Soils, DACO: 8.2.4.2
2377914	2010, [¹⁴ C]5-COOH-S-2200:Adsorption/Desorption in Three Soils, DACO: 8.2.4.2
2377915	2012, [¹⁴ C]2-COOH-S-2200: Adsorption/Desorption in Three Soils, DACO: 8.2.4.2
2377916	2012, [¹⁴ C]5-COOH-S-2200: Adsorption/Desorption in Three Soils, DACO: 8.2.4.2
2377918	2013, S-2200: Estimation of Adsorption Coefficient (Koc) of 2-CONH ₂ -S- 2200, 5-CONH ₂ -S- 2200 and Dx-CA-S-2200 by High Performance Liquid Chromatography, DACO: 8.2.4.2
2377919	2013, S-2200: Soil Adsorption/Desorption, DACO: 8.2.4.2
2377921	2012, Leaching of S-2200 and its Major Metabolites in Two Outdoor Lysimeters, DACO: 8.2.4.3
2377922	2013, Summary of Storage, Disposal, and Decontamination for S-2200 Fungicide Technical, DACO: 8.4.1
2377923	2011, Calculation of S-2200 Sediment Water Kinetics According to FOCUS (2006) Guidance, DACO: 8.6
2377924	2011, S-2200 and Metabolites: Storage Stability of Residues in EU Soil Stored Deep Frozen, DACO: 8.6
2377925	2013, Calculation of Laboratory Soil Kinetics for S-2200 and its Major Metabolites According to FOCUS (2006) Guidance, DACO: 8.6
2377927	2013, S-2200: Classification of Soils Used in Foreign Environmental Laboratory Studies and Their Representation Within North America, DACO: 8.6
2377928	2013, Ecological Exposure of Estimated Environmental Concentrations in Surface Water Following Application of S-2200 in the United States and Canada, DACO: 8.6
2377929	2010, Acute Oral Toxicity Study of S-2200 TG in Rats, DACO 4.2.1
2377950	2013, S-2200 4 SC Ag and VPP Fungicide: Annex IIIA Tier II Summary, Fate and Behavior in the Environment, DACO: 8.1, 8.2.1, 8.2.3.1, 8.2.4.1, 8.3.1, M12.7
2377953	2013, S-2200: Terrestrial Field Soil Dissipation on Bare Soil in Saskatchewan, Canada, DACO: 8.3.2.1
2377958	2013, S-2200: Terrestrial Field Soil Dissipation on Bare Ground in Ontario, Canada, DACO: 8.3.2.1
2377963	2013, S-2200: Terrestrial Field Soil Dissipation on Established Turfgrass in Ontario, Canada, DACO: 8.3.2.1
2377964	2012, Two-Generation Reproduction Toxicity Study of S-2200 TG in Rats, DACO: 4.5.1
2377968	2013, S-2200: Terrestrial Field Soil Dissipation on Bare Ground in North Dakota, DACO: 8.3.2.2
2377974	2013, S-2200: Terrestrial Field Soil Dissipation on Bare Soil in California, DACO: 8.3.2.3

2377979	2013, S-2200: Terrestrial Field Soil Dissipation on Established Turfgrass in California, DACO: 8.3.2.3
2377988	2013, S-2200: Terrestrial Field Soil Dissipation on Established Turfgrass in Mississippi, DACO: 8.3.2.3
2377993	2013, S-2200: Terrestrial Field Soil Dissipation on Bare Ground in Georgia, DACO: 8.3.2.3
2378014	2009, Acute Toxicity of S-2200 TG on Earthworms, <i>Eisenia fetida</i> Using an Artificial Soil Test, DACO: 9.2.3.1
2378015	2011, Acute Toxicity of 2-COOH-S-2200 on Earthworms, <i>Eisenia fetida</i> Using an Artificial Soil Test, DACO: 9.2.3.1
2378016	2011, Acute Toxicity of 5-COOH-S-2200 on Earthworms, <i>Eisenia fetida</i> Using an Artificial Soil Test, DACO: 9.2.3.1
2378017	2009, S-2200 TG: Acute Oral and Contact Toxicity to the Honeybee <i>Apis</i> <i>mellifera</i> L. in the Laboratory, DACO: 9.2.4.1
2378018	2013, S-2200 Technical Grade: 72-hour Acute Toxicity of Honey bee Larvae, <i>Apis mellifera</i> L., During an In Vitro Exposure, DACO: 9.2.4.2
2378019	2010, S-2200 25 SC: Toxicity to the Predatory Mite, <i>Typhlodromus pyri</i> Scheuten (Acari, Phytoseiidae) in Laboratory (Rate Response Test), DACO: 9.2.5
2378020	2010, S-2200 25 SC: Toxicity to the Aphid Parasitoid, <i>Aphidius</i> <i>rhopalosiphi</i> De Stefani Perez (Hymenoptera, Braconidae) in the Laboratory (Rate Response Test), DACO: 9.2.6
2378021	2013, Sublethal Toxicity of S-2200 TG to the Earthworm <i>Eisenia fetida</i> in Artificial Soil, DACO: 9.2.7
2378022	2010, S-2200 Technical Grade - Acute Toxicity to Water Fleas, (<i>Daphnia magna</i>), Under Static Conditions, Following OECD Guideline #202, OPPTS Draft Guideline 850.1010, The Official Journal of the European Communities L383A, Method C.2 and JMAFF 12 NohSan, No. 8147 Daphnia Acute Immobilization Test (2-7-2-1) and JMAFF 13 SeiSan No. 3986, DACO: 9.3.2
2378023	2012, 2-COOH-S-2200: Acute Toxicity to Water Fleas, (<i>Daphnia magna</i>) Under Static Conditions, Following OECD Guideline # 202 and The Official Journal of the European Communities L 142/456 Method C.2, DACO: 9.3.2
2378024	2012, 5-COOH-S-2200: Acute Toxicity to Water Fleas, (<i>Daphnia magna</i>) Under Static Conditions, Following OECD Guideline # 202 and The Official Journal of the European Communities L 142/456 Method C.2, DACO: 9.3.2
2378025	2012, S-2200-ORC: Acute Toxicity to Water Fleas, (<i>Daphnia magna</i>) Under Static Conditions, Following OECD Guideline # 202 and The Official Journal of the European Communities L 142/456 Method C.2, DACO: 9.3.2
2378026	2012, S-2167 (<i>R</i> -Isomer of S-2200) - Acute Toxicity to Water Fleas, (<i>Daphnia magna</i>) Under Static Conditions, Following OECD Guideline #202 and The Official Journal of the European Communities L 142/456, Method C.2, DACO: 9.3.2

2378027	2012, S-2354 (S-Isomer of S-2200) Acute Toxicity to Water Fleas,
	(Daphnia magna) Under Static Conditions, Following OECD Guideline
	#202 and The Official Journal of the European Communities L 142/456,
2250020	Method C.2, DACO: 9.3.2
2378028	2012, S-2200-OR - Acute Toxicity to Water Fleas, (<i>Daphnia magna</i>) Under
	Static Conditions, Following OECD Guideline #202 and The Official
	Journal of the European Communities L 142/456, Method C.2, DACO:
2279020	9.3.2 2010 S 2200 Technical Crade Full Life Crude Terricity Tect with Weter
2378029	2010, S-2200 Technical Grade - Full Life Cycle Toxicity Test with Water Fleas (<i>Daphnia magna</i>) Under Static Renewal Conditions, Following
	OPPTS Draft Guideline 850.1300, OECD Guideline #211, The Official
	Journal of the European Communities L225, Method C.20 JMAFF 12
	NohSan, No. 8147 <i>Daphnia spp</i> . Reproduction Toxicity Studies (2-7-2-3)
	and JMAFF 13 SeiSan No. 3986, DACO: 9.3.3
2378030	2012, S-2200: A 96-Hour Shell Deposition Test with the Eastern Oyster
	(<i>Crassostrea virginica</i>), DACO: 9.4.2
2378031	2012, S-2200: A 96-Hour Flow-Through Acute Toxicity Test with the
	Saltwater Mysid (<i>Americanysis bahia</i>), DACO: 9.4.2
2378032	2012, S-2200: A Flow-Through Life-Cycle Toxicity Test with the Saltwater
	Mysid (<i>Americanysis bahia</i>), DACO: 9.4.5
2378033	2012, S-2200 - Toxicity Test with Sediment-Dwelling Midges (Chironomus
	<i>riparius</i>) Under Static Conditions, Following OECD Guideline 219, DACO:
	9.4.5
2378034	2013, S-2200: A Life Cycle Toxicity Test with the Marine Amphipod
	(Leptocheirus plumulosus) Using Spiked Sediment, DACO: 9.4.5
2378035	2013, S-2200: A Life Cycle Toxicity Test with the Freshwater Amphipod
	(Hyalella azteca) Using Spiked Sediment, DACO: 9.3.4
2378036	2013, S-2200: A Life Cycle Toxicity Test with Chironomus dilutus Using
	Spiked Sediment, DACO: 9.4.5
2378037	2009, S-2200 Technical Grade - Acute Toxicity to Rainbow Trout
	(Oncorhynchus mykiss) Under Static Conditions, Following OECD
	Guideline #203, EC Guideline L383A, Method C.1 and OPPTS Draft
2250020	Guideline 850.1075, DACO: 9.5.2.1
2378038	2009, S-2167 (R-Isomer of S-2200) - Acute Toxicity to Rainbow Trout
	(Oncorhynchus mykiss) Under Static Conditions, Following OECD
	Guideline #203, EC Guideline L383A, Method C.1 and OPPTS Draft
2279020	Guideline 850.1075, DACO: 9.5.2.1
2378039	2009, S-2354 (S-Isomer of S-2200) - Acute Toxicity to Rainbow Trout (<i>Oncorhynchus mykiss</i>) Under Static Conditions, Following OECD
	Guideline #203, EC Guideline L383A, Method C.1 and OPPTS Draft
	Guideline #205, EC Guideline ES85A, Method C.1 and OPP 15 Draft Guideline 850.1075, DACO: 9.5.2.1
2378040	2012, 2-COOH-S-2200: Acute Toxicity to Rainbow Trout (<i>Oncorhynchus</i>
2370040	<i>mykiss</i>) Under Static Conditions, Following OECD Guideline # 203 and
	The Official Journal of the European Communities L 142/446 Method C.1,
	DACO: 9.5.2.1
	2110017101211

2378041	2012, 5-COOH-S-2200: Acute Toxicity to Rainbow Trout (<i>Oncorhynchus mykiss</i>) Under Static Conditions, Following OECD Guideline # 203 and
	The Official Journal of the European Communities L 142/446 Method C.1, DACO: 9.5.2.1
2378042	2012, S-2200-OR: Acute Toxicity to Rainbow Trout (Oncorhynchus
	mykiss) Under Static Conditions, Following OECD Guideline # 203 and
	The Official Journal of the European Communities L 142/446 Method C.1,
	DACO: 9.5.2.1
2378043	2012, S-2200-ORC: Acute Toxicity to Rainbow Trout (Oncorhynchus
	mykiss) Under Static Conditions, Following OECD Guideline # 203 and
	The Official Journal of the European Communities L 142/446 Method C.1,
	DACO: 9.5.2.1
2378044	2009, S-2200 Technical Grade - Acute Toxicity to Bluegill Sunfish
	(Lepomis macrochirus) Under Static Conditions, Following OECD
	Guideline #203, EC Guideline L383A, Method C.1 and OPPTS Draft
	Guideline 850.1075, DACO: 9.5.2.2
2378045	2009, S-2200 Technical Grade - Acute Toxicity to Fathead Minnow
	(<i>Pimephales promelas</i>) Under Static Conditions, Following OECD
	Guideline #203, EC Guideline L383A, Method C.1 and OPPTS Draft
2279046	Guideline 850.1075, DACO: 9.5.2.2
2378046	2012, S-2200: A 96-Hour Flow-Through Acute Toxicity Test with the
2279047	Sheepshead Minnow (<i>Cyprinodon variegatus</i>), DACO: 9.5.2.4
2378047	2010, S-2200 Technical Grade - Early Life-Stage Toxicity Test with
	Fathead Minnow, <i>Pimephales promelas</i> , Following OECD Guideline #210
2279049	and OPPTS Draft Guideline 850.1400, DACO: 9.5.3.1
2378048	2012, S-2200: An Early Life-Stage Toxicity Test with the Sheepshead
2378049	Minnow (<i>Cyprinodon variegatus</i>), DACO: 9.5.3.1
2378049	2010, Flow-Through Bioconcentration and Metabolism Study of [¹⁴ C]S-
2378050	2200 with Bluegill Sunfish (<i>Lepomis macrochirus</i>), DACO: 9.5.6
2578030	2009, S-2200 TG: An Acute Oral Toxicity Study with the Northern Bobwhite, DACO: 9.6.2.1
2378051	
2378031	2011, S-2200: An Acute Oral Toxicity Study with the Canary (<i>Serinus canaria</i>), DACO: 9.6.2.3
2378052	2009, S-2200 TG: A Dietary LC ₅₀ Study With the Northern Bobwhite,
2370032	DACO: $9.6.2.4$
2378053	2009, S-2200 TG: A Dietary LC_{50} Study With the Mallard, DACO: 9.6.2.5
2378053	2009, S-2200 FG. A Dietary LC ₅₀ Study with the Manaid, DACO. 9.0.2.5 2010, S-2200: A Pilot Reproduction Study with the Northern Bobwhite,
2570054	DACO: 9.6.3.1
2378055	2011, S-2200: A Reproduction Study with the Northern Bobwhite, DACO:
2370033	9.6.3.1
2378056	2010, S-2200: A Pilot Reproduction Study with the Mallard, DACO: 9.6.3.2
2378057	2010, S 2200: A Phot Reproduction Study with the Mallard, DACO: 9.6.3.2 2011, S-2200: A Reproduction Study with the Mallard, DACO: 9.6.3.2
2378059	2011, S-2200. A Reproduction Study with the Manard, DACO. 9.0.3.2 2012, S-2200-ORC: 72-Hour Toxicity Test with the Freshwater Green
2310037	Alga, <i>Pseudokirchneriella subcapitata</i> , DACO: 9.8.2
	1115a, 1 seauoni enienena suocupitata, DACO. 7.0.2

2378060	2010, S-2200 Technical Grade - 96-Hour Toxicity Test with the Freshwater
	Green Alga, <i>Pseudokirchneriella subcapitata</i> , Following OPPTS Draft Guideline 850.5400, DACO: 9.8.2
2378061	2012, 2-COOH-S-2200: 72-Hour Toxicity Test with the Freshwater Green Alga, <i>Pseudokirchneriella subcapitata</i> , Following The Official Journal of the European Communities L383A, Method C.3, JMAFF 12 Nohsan, No. 8147 Alga, Growth Inhibition Test 2-7-7 and JMAFF 13 Seisan No. 3986, DACO: 9.8.2
2378062	2012, 5-COOH-S-2200: 72-Hour Toxicity Test with the Freshwater Green Alga, <i>Pseudokirchneriella subcapitata</i> , Following The Official Journal of the European Communities L383A, Method C.3, JMAFF 12 Nohsan, No. 8147 Alga, Growth Inhibition Test 2-7-7 and JMAFF 13 Seisan No. 3986, DACO: 9.8.2
2378063	2012, S-2200-OR: 72-Hour Toxicity Test with the Freshwater Green Alga, <i>Pseudokirchneriella subcapitata</i> , Following The Official Journal of the European Communities L383A, Method C.3, JMAFF 12 Nohsan, No. 8147 Alga, Growth Inhibition Test 2-7-7 and JMAFF 13 Seisan No. 3986, DACO: 9.8.2
2378065	2012, S-2200: A 96-Hour Toxicity Test with the Freshwater Alga (<i>Anabaena flos-aquae</i>), DACO: 9.8.2
2378066	2012, S-2200: A 96-Hour Toxicity Test with the Freshwater Diatom (<i>Navicula pelliculosa</i>), DACO: 9.8.2
2378067	2013, S-2167 (R-Isomer of S-2200) - 72-Hour Toxicity Test with the Freshwater Green Alga, <i>Pseudokirchneriella subcapitata</i> , Following the Official Journal of the European Communities L383A, Method C.3, DACO: 9.8.2
2378068	2013, S-2354 (S-Isomer of S-2200) - 72-Hour Acute Toxicity Test with Freshwater Green Alga, <i>Pseudokirchneriella subcapitata</i> , Following the Official Journal of the European Communities L383A, Method C.3, DACO: 9.8.2
2378069	2012, S-2200: A 96-Hour Toxicity Test with the Marine Diatom (<i>Skeletonema costatum</i>), DACO: 9.8.3
2378070	2012, S-2200: A Toxicity Test to Determine the Effects of the Test Substance on Seedling Emergence of Ten Species of Plants, DACO: 9.8.4
2378071	2012, S-2200: A Toxicity Test to Determine the Effects of the Test Substance on Vegetative Vigor of Ten Species of Plants, DACO: 9.8.4
2378072	2012, S-2200: A 7-Day Static-Renewal Toxicity Test with Duckweed (<i>Lemna gibba</i> G3), DACO: 9.8.5
2378073	2009, Analytical Method Verification for the Determination of S-2200 TG in Avian Diet, DACO: 9.9
2378074	2012, Method Verification for the Determination of S-2200 on Drift Cards Used to Confirm Application During Non-Target Plant Tests, DACO: 9.9
2378075	2011, Effects of S-2200 TG on the Activity of the Soil Microflora, DACO: 9.9
2420267	2012, Analytical Method Verification for the Determination of S-2200 in Freshwater and Saltwater, DACO: 8.2.2.3

2420268	2012, Analytical Method Verification for the Determination of S-2200 in Freshwater and Saltwater, DACO: 8.2.2.3
2420269	2013, Analytical Method Verification for the Determination of S-2200 in
2420270	Sediment, DACO: 8.2.2.2
2420270	2013, S-2200 Technical Grade - 72-Hour Acute Toxicity of Honey Bee
2420271	Larvae, <i>Apis mellifera</i> L., during an In Vitro Exposure , DACO: 9.2.4.2
2420271	2013, S-2200: A Life Cycle Toxicity Test with the Freshwater Amphipod
2420273	(<i>Hyalella azteca</i>) Using Spiked Sediment, DACO: 9.3.4
2420273	2013, S-2200: A Life Cycle Toxicity Test with the Freshwater Amphipod (<i>Hyalella azteca</i>) Using Spiked Sediment, DACO: 9.3.4
2420275	
2420273	2013, S-2200: A Life Cycle Toxicity Test with the Freshwater Amphipod (Hyglalla astag) Using Spiked Sediment DACO: 0.3.4
2420276	(<i>Hyalella azteca</i>) Using Spiked Sediment , DACO: 9.3.4
2420270	2013, S-2200: A Life Cycle Toxicity Test with the Freshwater Amphipod
2420277	(<i>Hyalella azteca</i>) Using Spiked Sediment , DACO: 9.3.4
2420277	2013, S-2200: A Life Cycle Toxicity Test with the Freshwater Amphipod
2420279	(<i>Hyalella azteca</i>) Using Spiked Sediment, DACO: 9.3.4
2420278	2013, S-2200: A Life Cycle Toxicity Test with the Freshwater Amphipod
2420270	(Hyalella azteca) Using Spiked Sediment, DACO: 9.3.4
2420279	2012, S-2200: A 96-Hour Shell Deposition Test with the Eastern Oyster
2 4 2 2 2 2 2	(Crassostrea virginica), DACO: 9.4.2
2420280	2012, S-2200: A 96-Hour Flow-Through Acute Toxicity Test with the
	Saltwater Mysid (Americamysis bahia), DACO: 9.4.2
2420281	2013, S-2200: A Life Cycle Toxicity Test with the Marine Amphipod
	(Leptocheirus plumulosus) Using Spiked Sediment, DACO: 9.4.5
2420283	2013, S-2200: A Life Cycle Toxicity Test with the Marine Amphipod
	(Leptocheirus plumulosus) Using Spiked Sediment, DACO: 9.4.5
2420284	2012, S-2200: A Flow-Through Life-Cycle Toxicity Test with the Saltwater
	Mysid (Americamysis bahia), DACO: 9.4.5
2420285	2013, S-2200: A Life Cycle Toxicity Test with Chironomus dilutus Using
	Spiked Sediment, DACO: 9.4.5
2420286	2013, S-2200: A Life Cycle Toxicity Test with Chironomus dilutus Using
	Spiked Sediment, DACO: 9.4.5
2420287	2012, S-2200: A 96-Hour Flow-Through Acute Toxicity Test with the
	Sheepshead Minnow (Cyprinodon variegatus), DACO: 9.5.2.4
2420288	2012, S-2200: An Early Life-Stage Toxicity Test with the Sheepshead
	Minnow (Cyprinodon variegatus), DACO: 9.5.3.1
2420289	2012, S-2200: A 96 Hour Toxicity Test with the Freshwater Alga
	(Anabaena flos-aquae), DACO: 9.8.2
2420290	2012, S-2200: A 96 Hour Toxicity Test with the Freshwater Diatom
	(Navicula pelliculosa), DACO: 9.8.2
2420291	2012, S-2200: A 96 Hour Toxicity Test with the Marine Diatom
	(Skeletonema costatum), DACO: 9.8.3
2420292	2012, S-2200: A Toxicity Test to Determine the Effects of the Test
	Substance on Seedling Emergence of Ten Species of Plants, DACO: 9.8.4

2420293	2012, S-2200: A Toxicity Test to Determine the Effects of the Test
	Substance on Vegetative Vigor of Ten Species of Plants, DACO: 9.8.4
2420294	2012, S-2200: A 7-Day Static-Renewal Toxicity Test with Duckweed
	(Lemna gibba G3), DACO: 9.8.5
2420295	2012, Method Verification For the Determination of S-2200 on Drift Cards
	Used to Confirm Application During Non-Target Plant Tests, DACO: 9.9

4.0 Value

PMRA	Reference
Document	
Number	
2377846	2013. Appendix 1 - Trial Reports for "S-2200 Fungicide: Annex IIA Tier II
	Summary, Efficacy Data and Information on S-2200 4 SC Fungicide,
	Containing Mandestrobin, for Use on Canola, Grape, Strawberry and
	Turfgrass". DACO 10.1, 10.2.1, 10.2.2, 10.2.3.1, 10.2.3.3, 10.3.1, 10.3.2, 10.4,
	10.5, 10.5.1, 10.5.2, 10.5.3, 10.5.4
2377844	2013. S-2200 Fungicide: Annex IIA Tier II Summary, Efficacy Data and
	Information on S-2200 4 SC Fungicide, Containing Mandestrobin, for Use on
	Canola, Grape, Strawberry and Turfgrass. DACO 10.1, 10.2.1, 10.2.2, 10.2.3.1,
	10.2.3.3, 10.3.1, 10.3.2, 10.4, 10.5, 10.5.1, 10.5.2, 10.5.3, 10.5.4, M12.7
2378119	2013. S-2200 Fungicide: Annex IIA Tier II Summary, Efficacy Data and
	Information on S-2200 3.2 FS Fungicide, a Seed Protectant Containing
	Mandestrobin, for Control of Seed and Seedling Diseases of Canola, Corn and
	Soybeans. DACO 10.1, 10.2.1, 10.2.2, 10.2.3.1, 10.2.3.2, 10.2.3.3, 10.3, 10.3.2,
	10.3.3, 10.4, 10.5, 10.5.1, 10.5.2, 10.5.3, 10.5.4, M12.7
2378120	2013. Appendix 1: Trials Reports for "S-2200 Fungicide: Annex IIA Tier II
	Summary, Efficacy Data and Information on S-2200 3.2 FS Fungicide, a Seed
	Protectant Containing Mandestrobin, for Control of Seed and Seedling Diseases
	of Canola, Corn and Soybeans". DACO 10.1, 10.2.1, 10.2.2, 10.2.3.1, 10.2.3.2,
	10.2.3.3, 10.3, 10.3.3, 10.4, 10.5, 10.5.1, 10.5.2, 10.5.3, 10.5.4, M12.7

B. Additional Information Considered

i) Published Information

1.0 Human and Animal Health

PMRA Document Number	Reference
2409268	U.S. EPA. 2012b. Standard Operating Procedures for Residential Pesticide Exposure Assessment. DACO 12.5.5
2528612	Clapp, M.J.L, Wade, J.D. and D.M. Samules. 1982. Control of Nephrocalcinosis by Manipulating the Calcium:Phosphorus Ratio in Commercial Rodent Diets, Laboratory Animals, 16:130-132. DACO: 4.8

2528613	Cockell, K.A. and B. Belonje. 2004. Nephrocalcinosis Caused by Dietary Calcium:Phosphorus Imbalance in Female Rats Develops Rapidly and Is
	Irreversible, The Journal of Nutrition, 134:637-640. DACO: 4.8
2528614	Frazier, K.S., Seely, J.C., Hard, G.C., Betton, G., Burnett, R., Nakatsuji, S.,
	Nishikawa, A., Durchfeld-Meyer, B. and A. Bube. 2012. Proliferative and
	Nonproliferative Lesions of the Rat and Mouse Urinary System, Toxicologic
	Pathology, 40: 14S-86S. DACO: 4.8
2528616	Ritskes-Hoitinga, J., Lemmens, A.G., Danse, L.H.J.C. and A.C. Beynen. 1989.
	Phosphorus-Induced Nephrocalcinosis and Kidney Function in Female Rats, The
	Journal of Nutrition, 119:1423-1431. DACO: 4.8
2528615	Woo, D.C. and R.M. Hoar. 1972, "Apparent Hydronephrosis" as a Normal
	Aspect of Renal Development in Late Gestation of Rats: The Effect of Methyl
	Salicylate, Teratology, 6:191-196. DACO: 4.8