TEXT SEARCHABLE DOCUMENT

Data Evaluation Report on the Acute Dietary Toxicity of AMPA to Avian Species, Colinus virginianus

PMRA Submission Number {.....}

EPA MRID Number 43334710

Data Requirement:

PMRA Data Code EPA DP Barcode OECD Data Point EPA MRID **EPA** Guideline

{.....} Not Provided {.....} 43334710 OPPTS 850.2200 (71-2a)

Purity: 87.8%

Aminomethyl Phosphonic Acid Test material: (Glyphosate Degradate) Common name: AMPA Chemical name: IUPAC: Not Reported CAS name: Not Reported CAS No. Not Reported

Synonyms: None Reported

Primary Reviewer: John Marton Staff Scientist, Cambridge Environmental Inc.

Secondary Reviewer: Teri S. Myers Senior Scientist, Cambridge Environmental Inc.

Primary Reviewer: Stephen Carey EPA Biologist, OPP, EFED, ERBIII

{EPA/OECD/PMRA}

Reference/Submission No.: {......}

Company Code {.....} [For PMRA] Active Code {.....} [For PMRA] Use Site Category: [For PMRA] {.....} EPA PC Code 417300 (Parent Compound) & 207800 (Degradate Compound)

Date Evaluation Completed: July 18 2007

CITATION: Long, R.D., G.J. Smith, J.B. Beavers and S.P. Lynn. 1994. AMPA: A Dietary LC₅₀ Study with the Northern Bobwhite. Unpublished study performed by Wildlife International Ltd., Easton, MD. Laboratory report number 139-275. Study sponsored by Monsanto Agricultural Company, St. Louis, Missouri. Study completed October 28, 1991.

DISCLAIMER: This document provides guidance for EPA and PMRA reviewers on how to complete a data evaluation record after reviewing a scientific study concerning the acute dietary toxicity of a pesticide to avian species. It is not intended to prescribe conditions to any external party for conducting this study nor to establish absolute criteria regarding the assessment of whether the study is scientifically sound and whether the study satisfies any applicable data requirements. Reviewers are expected to review and to determine for each study, on a case-bycase basis, whether it is scientifically sound and provides sufficient information to satisfy applicable data requirements. Studies that fail to meet any of the conditions may be accepted, if appropriate; similarly, studies that meet all of the conditions may be rejected, if appropriate. In sum, the reviewer is to take into account the totality of factors related to the test methodology and results in determining the acceptability of the study.

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Signature: Date: 1/12/07

Signature: Date: 2/12/07

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Date: 7/18/07

PMRA Submission Number {.....}

EPA MRID Number 43334710

EXECUTIVE SUMMARY:

The subacute dietary toxicity of AMPA (Glyphosate Degradate) to 10-d-old Northern Bobwhite Quail (*Colinus virginianus*) was assessed over 8 days. AMPA was administered to the birds in the diet at 0 (vehicle control), 493, 878, 1563, 2774 and 4934 mg ai/kg dw of diet; verification of these dose levels was provided in MRID 43334712 and showed that all values were within 93-109% of target (see Reviewer's Comments section for details). The 8-day acute dietary LC_{50} was >4934 mg ai/kg diet. The 8-day NOAEC of AMPA based on the lack of treatment-related mortality and sub-lethal effects was 4934 mg ai/kg diet. According to the US EPA classification, AMPA (Glyphosate Degradate) would be classified as practically non-toxic to Northern bobwhite quail on a subacute dietary basis at a nominal concentration of 4934 mg ai/kg diet.

One mortality was observed on Day 2 in the 493 mg ai/kg diet treatment group; however, this was not considered to be treatment related. Furthermore, the test material did not significantly impact weight gain or food consumption.

This toxicity study is classified as scientifically sound and does satisfy the guideline requirement for a subacute dietary toxicity study with Northern bobwhite quail.

Results Synopsis

Test Organism Size/Age(Mean Weight): 10 Days; 20 (18-22) g

 LC_{50} : >4934 mg ai/kg diet Probit slope: N/A NOAEC: 4934 mg ai/kg diet Endpoint(s) affected: None 95% C.I.: N/A 95% C.I.: N/A

PMRA Submission Number {.....}

EPA MRID Number 43334710

I. MATERIALS AND METHODS:

GUIDELINE FOLLOWED:

This study was conducted following guidelines outlined in Section 71-2 of the Environmental Protection Agency Registration Guidelines, *Pesticide Assessment Guidelines, FIFRA Subdivision E, Hazard Evaluation, Wildlife and Aquatic Organisms*; and upon ASTM Standard E857-87, "Standard Practice for Conducting Subacute Dietary Toxicity Tests with Avian Species." The following deviation from OPPTS 850.2200 was noted:

The physiochemical properties of the test material were not reported.

This deviation did not impact the acceptability of the study.

COMPLIANCE:

Signed and dated No Data Confidentiality, GLP and Quality Assurance statements were provided. This study was conducted in compliance with GLP standards as published by the U.S. EPA in 40 CFR, Part 160; OECD, ISBN 92-84-12367-9; and Japan MAFF, 59 NohSan, Notification No. 3850, Agricultural Production Bureau.

A. MATERIALS:

1. Test Material	Aminomethyl Phosphonic Acid (Glyphosate Degradate)				
Description:	White Powder				
Lot No./Batch No. :	PIT-9008-2407T				
Purity:	87.8%				

Stability of Compound Under Test Conditions:

Samples of the test diets were taken to verify the test concentrations administered and to confirm the stability and homogeneity of the test substance in the diets. Samples were frozen and transferred to Monsanto Environmental Health Laboratories for analysis. The results of this analysis are provided in a separate report, MRID 43334712. They revealed that the test material levels taken on days 0 and 5 ranged from 88-105% of target.

Storage Conditions of Test Chemicals:

Stored at room temperature.

PMRA Submission Number {.....}

EPA MRID Number 43334710

Physicochemical properties of AMPA.

Parameter	Values	Comments
Water solubility at 20EC	Not Reported	
Vapor pressure	Not Reported	· .
UV absorption	Not Reported	
pKa	Not Reported	
Kow	Not Reported	

(OECD recommends water solubility, stability in water and light, pKa, Pow, and vapor pressure of test compound)

2. Test organism:

Species (common and scientific names): Northern bobwhite quail (*Colinus virginianus*) (*EPA recommends using either bobwhite quail or mallard duck.*)

Age at study initiation: 10 Days (EPA recommends: 10-14 days old)

Weight at study initiation (mean and range): 20 (18-22) g

Source: On-site Production Flock.

B. STUDY DESIGN:

1. Experimental Conditions

a. Range-finding Study: No range-finding studies were reported.

b. Definitive Study:

Table 1: Experimental Parameters

Parameter	Details	Remarks		
		Criteria		
Acclimation Period: Conditions: (same as test or not) Feeding: Health: (any mortality observed)	Continuous Same as test Game bird ration formulated to Wildlife International Ltd.'s specifications Not reported; however, birds exhibiting abnormal behavior or physical injury were not used in the definitive test.	Food and water from the town of Easton public water supply were provided <i>ad libitum</i> .		

PMRA Submission Number {.....}

EPA MRID Number 43334710

Parameter	Details	Remarks		
	Details	Criteria		
Pen size and construction materials	Thermostatically controlled brooding pens had floors that measured approximately 72 x 90 cm and ceiling height was approximately 23 cm. External walls, ceilings and floors were constructed of galvanized steel wire and sheeting.	Recommended pen size is about 35 x 100 x 24 cm		
Test duration	5 days with treated feed and 3 days with untreated feed	Recommended test duration is 5 days with treated feed and at least 3 days observation with "clean" feed.		
Test concentrations nominal:	0 (vehicle control), 493, 878, 1563, 2774 and 4934 mg ai/kg dw of diet	The reviewer corrected the nominal concentrations for the purity of the test material (87.8%).		
measured:	Measured samples of dose levels showed them to be 93-109% of	The results from the analysis of the test material in the feed were provided in MRID 43334712.		
	target.	Five or six test concentrations should be used in a geometric scale, unless the $LC_{50} > 5000$ mg ai/kg diet.		
Solvent/vehicle, if used type: amount:	Corn Oil 2%	Recommended solvents include distilled water, corn oil, propylene glycol, 1% carboxymethylcellulose, or gum arabic. The solvent should not be more than 2%.		
Diet preparation and feeding	Test diets were prepared by mixing the test substance into the diet with corn oil. Mixing was done with a Hobart mixer. Diets were prepared on the day of test initiation and sufficient feed was prepared for the duration of the treated feed period.	The control was treated with 2% corn oil only. The control group should be tested with a diet containing the maximum amount of vehicle used in treated diets.		
Feed withholding period	None reported			
Stability and homogeneity of test material in the diet determined (Yes/No)	Samples were shipped to Monsanto Environmental Health Laboratories for analysis; the results of these analyses are described in another study report, MRID 43334712 (see Reviewer's Comments section for			

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PMRA Submission Number {.....}

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EPA MRID Number 43334710

Parameter	Details	Remarks <i>Criteria</i>	
	details).		
Number of birds per replicate/groups for negative control: for vehicle control: for treated:	N/A 10 10	The recommended number of birds per replicate is a minimum of ten.	
Number of replicates/group (if used) for negative control: for vehicle control: for treated:	N/A 3 1		
<u>Test conditions</u> temperature: relative humidity(%): photoperiod:	Brooding Compartment 35±2°C Room 25±2°C 41±1% 16L:8D; 130 lux	Recommended brooder temperature is about 35°C (95°F) Recommended room temperature is 22-27°C (71-81°F) Recommended relative humidity is 30-80% Recommended photoperiod is a minimum of 14 hours of light.	
Reference chemical, if used	N/A; a reference chemical was not used		

2. Observations: Table 2: Observations

Parameters	Details	Remarks		
Parameters measured (mortality/body weight/ mean feed consumption/ others)	-Mortality -Average Weight Gain -Feed Consumption	Feed consumption was reported as an estimate due to the unavoidable wastage of the by the birds.		
Indicate the stability and homogeneity of test chemical in the diet	Not reported			
Indicate if the test material was regurgitated	No regurgitation was reported			
Treatments on which necropsies were performed	No necropsies were performed			
Observation intervals	Daily			
Were raw data included?	Sufficient summarized data tables			

PMRA Submission Number {.....}

EPA MRID Number 43334710

Parameters	Details	Remarks
	were provided.	

II. RESULTS AND DISCUSSION:

A. MORTALITY:

Throughout the duration of the test, a single mortality occurred in the nominal 493 mg ai/kg diet treatment group on Day 2. This one mortality was not attributed to the test material. Therefore, the resulting NOAEC and LC_{50} values were 4934 and >4934 mg ai/kg diet, respectively.

Table 3: Effect of AMPA on Mortality of Colinus virginianus

Treatment (mg ai/kg diet)		No. of	Cumulative Mortality				
		Birds	Day 1	Day 2	Day 3	Day 5	Day 8
Vehic	le Control	30	0	0	0	0	0
	493	10	0	1	· 1	-1	1
	878	10	0	0	0	0	0
1563		10	0	0	0	0	0
2774		10	0	0	0	0	0
4934		10	0	0	0	0	0
NOAEC 4934 mg ai/			kg diet	•			
LC ₅₀		>4934 mg ai/kg diet				· · · ·	
Reference	mortality	N/A	N/A N/A N/A N/A N/A				
chemical	LC ₅₀	N/A					
	NOAEC	N/A					

N/A- Not Applicable

B. SUB-LETHAL TOXICITY ENDPOINTS:

No differences in food consumption of body weight gain were noted at any treatment level. Furthermore, no behavioral abnormalities were noted. The resulting NOAEC and EC_{50} values were 4934 and >4934 mg ai/kg diet, respectively.

PMRA Submission Number {.....}

EPA MRID Number 43334710

				Observation					
Treatment (mg ai/kg diet)		Body Weight (g)				Food Consumption (g/bird/day)			
		Day 0	Day	5 Day	y 8	Days 0-5	Days 6-8		
Vehicle	Control	18	30	39)	8	10		
49	93	20	30	39)	8	10		
878		20	32	42	2	8	11		
1563		22	35	4	5	9	10		
2774		21	34	43	3	7	8		
4934		19	33	4	1	10	9		
NOAEC	-	4934 mg ai/kg diet			· · · · · · · · · · · · · · · · · · ·				
EC ₅₀	· · · ·	>4934 mg ai/kg diet							
Reference chemical	effect	N/A	N/A N/A N/A N/A N/A						
	NOAEL	N/A	N/A						
	LC ₅₀	N/A	N/A						

Table 4: Sub-lethal Effect of AMPA on Colinus virginianus

N/A- Not Applicable

C. REPORTED STATISTICS:

All toxicity values were determined by visual inspection of the data.

D. VERIFICATION OF STATISTICAL RESULTS:

Statistical Method(s): The lack of treatment-related mortality precluded the statistical analysis of the data. As values for weight gain and food consumption at all treatment levels were similar to the control values, the reviewer visually determined the toxicity values. All values were determined based on the nominal concentrations which the reviewer corrected for the purity of the test material (87.8%).

LC_{50} : >5620 mg ai/kg diet	95% C.I.: N/A
NOAEC: 5620 mg ai/kg diet	
Probit Slope: N/A	95% C.I.: N/A

Adjusted for active ingredient: (Optional if over 80% ai)

LC ₅₀ : >4934 mg ai/kg diet	95% C.I.: N/A
NOAEC: 4934 mg ai/kg diet	
Probit Slope: N/A	95% C.I.: N/A

PMRA Submission Number {.....}

EPA MRID Number 43334710

E. STUDY DEFICIENCIES:

There were no study deficiencies.

F. REVIEWER'S COMMENTS:

The reviewer's results were identical to those of the study author.

Samples of the test diets were taken to verify the test concentrations administered and to confirm the stability and homogeneity of the test substance in the diets. Samples were frozen upon transfer to and for storage at Monsanto Environmental Health Laboratories. Results of these analyses are described in a separate report:

MRID 43334712. Schermes, S. and K.W. McCune. (1994). Results of the Analyses of Avian Diet Samples for AMPA (aminomethyl phosphonic acid). Guideline 71-2. Submitted by The Agricultural Group of Monsanto Company, St. Louis, MO.

Data from this report showed that mixing was uniform (coefficient of variations <3%), as determined by high and low level mixtures of avian diet and test material. Stability of AMPA in avian diet over 5 days at room temperature was also shown (day 0 and 5 samples ranged from 88-105% of target). Adequate diet homogeneity was observed and concentrations of AMPA in avian diet were shown to accurately reflect target levels (93-109%). The analytical report additionally noted a positive diet interference problem (resulting in higher recoveries), where co-extracted amino acids from the diet matrix were possibly reacting with the derivitization agent used in the LC-UV process. To minimize this interference, some sample recovery values were corrected using the concurrently analyzed mean QC sample recoveries.

The in-life portion of the definitive toxicity test was conducted from September 6 to September 14, 1990.

G. CONCLUSIONS:

This study is scientifically sound and does satisfy the guideline requirements for a subacute dietary toxicity test with the Northern bobwhite quail. The NOAEC and LC_{50} values were 4934 and >4934 mg ai/kg diet, respectively.

LC₅₀: >4934 mg ai/kg diet NOAEC: 4934 mg ai/kg diet Endpoint(s) affected: None 95% C.I.: N/A

III. REFERENCES:

Environmental Protection Agency. 1982. Pesticide Assessment Guidelines, FIFRA Subdivision E, Hazard Evaluation: Wildlife and Aquatic Organisms, Subsection 71-2. Office of Pesticide Programs. Washington, DC. 86 pp.

ASTM Standard E857-81. 1982 "Standard Practice for Conducting Subacute Dietary Toxicity Tests with Avian Species". American Society for Testing and Materials.

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Page 9 of 10

PMRA Submission Number {.....}

EPA MRID Number 43334710

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Thompson, W.R. 1947. Bacteriological Reviews. Vol II, 2:115-145.

Stephan, C.E. 1977. Methods for Calculating an LC50. <u>Aquatic Toxicology and Hazard Evaluations</u>, Amer. Soc. Test Mat., Pub. No. STP 634: 65-84.