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## UNITED STATES ENVIRONMENTAL PROTECTION AGENCY WASHINGTON, D.C. 20460

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OFFICE OF
PESTICIDES AND TOXIC SUBSTANCES

MEMORANDUM

SUBJECT:

Review of Acute Toxicology Data in Support of the Registration of Tempo" 20% Wettable Powder for Use in Food Handling Establishments

EPA No. 3125-GTT, 6H-5515 Record No. 183991, 181433 Project No. 7-0205 Tox. Chem. No. 266E

TO:

George LaRocca (PM Team #15) Registration Division (TS-767c)

FROM:

John E. Whalan, D.A.B.T., Toxicologist

Section II, Toxicology Branch

Hazard Evaluation Division (TS-769c)

THRU:

Edwin R. Budd, Section Head Section II, Toxicology Branch

Hazard Evaluation Division (TS-769c)

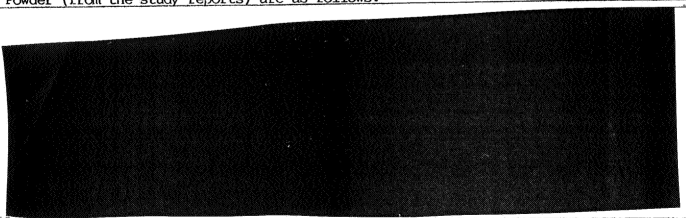
John Shalon 1-21-87

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Mobay Chemical Corporation has requested the registration of Tempo 20% Wettable Powder, "for use by pest control operators and professional applicators for pest control in buildings and structures including food areas of food handling establishments." The proposed tolerance is 0.05 ppm for cyfluthrin (cyano(4-fluoro-3-phenoxyphenyl)methyl 3-(2,2-dichloroethenyl)-2,2-dimethyl-cyclopropanecarboxylate) in or on all food items in food handling establishments where food or food products are held, processed, or prepared. It is to be applied as a general, spot, and crack and crevice treatment. Food should be removed or covered. The product should not be applied directly to food or food handling surfaces.

Tempo™ 20% Wettable Powder is reportedly

Powder (from the study reports) are as follows:



In support of this action, the following studies were submitted:

- 1. Acute Oral Toxicity of Baythroid 20% Wettable Powder in Albino Rats
- 2. Acute Dermal Toxicity of Baythroid 20% Wettable Powder in Albino Rabbits
- 3. Acute Inhalation Toxicity Study with \*Baythroid 20% Wettable Powder Dust in Rats
- 4. Primary Eye Irritation of ®Baythroid 20% Wettable Powder in Albino Rabbits
- 5. Primary Dermal Irritation of Baythroid 20% Wettable Powder in Albino Rabbits
- 6. Dermal Sensitization of \*Baythroid 20% Wettable Powder in Male Guinea Pigs

These studies have been reviewed and found to be acceptable. Reviews of these studies follow.

The ADI was recently reevaluated and established as 0.025 mg/kg/day (based on a 2-year rat feeding study with a NOEL of 2.5 mg/kg/day). Using this ADI, the current commitment for Toxicology Branch approved tolerances is 34.38% of the ADI, with a TMRC of 0.0086 mg/kg/day (60 kg body weight, 1.5 kg diet). Granting these tolerances will increase the % ADI to 39.38%, and the TMRC to 0.0098 mg/kg/day (see attachment). The Toxicology Branch has no objection to granting these tolerances, RCB considerations permitting. The inert ingredients have been cleared. The submitted label is acceptable.

An 8-Point Summary is attached.

#### ACUTE ORAL TOXICITY STUDY OF BAYTHROID 20% WP IN RATS

Mobay Corporation; Report No. 756; June 11, 1986; Accession No. 264524

PROTOCOL: Male (280-356 g) and female (190-222 g) Sprague-Dawley rats were randomly assigned to groups of 5 rats/sex. Baythroid 20% WP (20.8% cyfluthrin) was formulated in deionized water. The rats were fasted overnight, then dosed by gavage with the test article (dose volume of 10 ml/kg) as follows:

Doses Male	(mg/kg) <u>Female</u>
500	500
1000	1000
	1500
2000	2000
3000	
4000	

The rats were observed at least once daily for clinical signs, and weighed weekly during the 14-day study. Food and water were available ad libitum. Terminal weights were measured for all animals which died or were sacrificed moribund. All rats were necropsied and examined grossly.

RESULTS: There were no deaths in males and females dosed at 500 and 1000  $\rm mg/kg$ . The females were nearly twice as sensitive to the test article as the males. The LD50 values were calculated to be 3084 (2160-4459)  $\rm mg/kg$  for males, and 1733 (1491-2014)  $\rm mg/kg$  for females. The slopes of the dose-response curves were 4.5 and 13.6, respectively. There were no clinical signs in the 500  $\rm mg/kg$  males. The occurrence of clinical signs in the other groups was as follows:

		s (mg 2000	/kg) 3000	4000		males 1000	(mg/ 1500	
Salivation	Х	Х	X	X	X	X	X	X
Lacrimation		X		X			X	Х
Red nasal discharge				X				
Diarrhea		Х		X			X	X
Urine stained fur		Х	Х	Χ			Х	X
Decreased activity		X	X	X	Х	Х	X	X
-			X		X			
Tremors		Х	X	Х	х		Х	X
Ataxia Writhing		X	X	X		X	X	X

Clinical signs (not otherwise specified) were observed immediately after dosing, and had reversed by day 5 in the survivors. There were no control animals against which to compare body weights, but there was clearly a doserelated decrease in male weight gain on day 7. Dose-related gross lesions seen in rats which died during the study included discharge and staining of the fur, reddened cervical lymph nodes, reddened lungs, salivation, fluid-filled stomach, and reduced amount of stomach ingesta. There were no compound-related gross lesions in rats sacrificed on day 14.

STUDY CLASSIFICATION: This study is CORE MINIMUM, Toxicity Category III. There was a great deal of disparity between the LD50 values, slopes, and clinical signs for the two sexes. It is very likely that some groups were dosed with improper or nonhomogeneous formulations. If in fact there was a genuine sex difference, it would probably be due to the inert ingredients used in the formulation in which case, there should have been vehicle controls. In the absence of analytical information, the data presented in this report are suspect. This study received Quality Assurance Review. It was submitted to the EPA although the Quality Assurance Officer declared it to be in violation of the Good Laboratory Practice requirements for analysis of test article homogeneity, stability, and concentration.

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### ACUTE DERMAL TOXICITY STUDY OF BAYTHROID 20% WP IN RABBITS

Mobay Corporation; Report No. 743; April 23, 1986; Accession No. 264524

PROTOCOL: Five male (2.85-3.06 kg) and five female (3.02-23.42 kg) New Zealand White rabbits were given the limit dose (2000 mg/kg) of Baythroid 20% WP (20.8% cyfluthrin). The test article was formulated as a paste (using an unspecified vehicle) and applied to a 240 cm² area on the shaved backs of each rabbit. The doses were occluded with gauze, hypoallergenic tape, sheet plastic, and an elastic bandage. They were also fitted with plastic collars. The doses were removed after 24 hours by wiping the dosing sites with water moistened paper towels.

The rabbits were observed at least once daily for clinical signs, and weighed weekly during the 14-day study. Food and water were available ad libitum. All rabbits were necropsied and examined grossly.

RESULTS: There were no deaths or clinical signs during this study. Thus, the LD50 in both sexes was >2000 mg/kg (the limit dose). Body weight gain appeared normal for both sexes. The only compound-related gross lesion was a red zone on the back of one male, presumably due to local irritation.

STUDY CLASSIFICATION: This study is CORE MINIMUM, Toxicity Category III.

There was no mention of the vehicle used to formulate a paste. This study received Quality Assurance Review, but the Quality Assurance Officer failed to note that it was in violation of the Good Laboratory Practice requirements to define the test article formulation, and analyze the test article for homogeneity, stability, and concentration.

## ACUTE INHALATION TOXICITY STUDY OF BAYTHROID 20% WP IN RATS

Mobay Corporation; Report No. 758; June 18, 1986; Accession No. 264524

PROTOCOL: Male (189-262 g) and female (180-216 g) Sprague-Dawley rats were randomly assigned to groups of 10 rats/sex. They were dynamically exposed "head-only" to either air (nontreated controls) or Baythroid 20% WP (20.8% cyfluthrin) aerosol in a 60-liter cylindrical chamber. The test article was generated with a Wright Dust Feed at a nominal chamber concentration of 12.052 mg/l (maximum attainable concentration). The aerosol was generated 30 minutes before introducing the rats into the chamber in order to reach a state of equilibrium. Exposure duration was 4 hours. Aerosol particle size distribution was assessed at four intervals with a TSI Aerodynamic Particle Sizer. Gravimetric concentration measurements were made at four intervals by passing the aerosol through 0.5 u Millipore PVC-5 filters placed near the rats' breathing zone.

The rats were observed for clinical signs during and after the exposure, and at least once daily during the 14-day study. They were weighed prior to exposure, and on days 3, 7, and 14. Food and water were available ad libitum, except during exposure. All rats were necropsied and examined grossly.

RESULTS: The gravimetric analyses of the chamber atmosphere indicated a mean chamber concentration of 1.180 mg/l in the breathing zone of the rats. The chamber concentration was sufficiently consistent throughout the exposure. This was reportedly a maximum attainable concentration. The mass median aerodynamic diameter was calculated to be 3.8 u (particle size distribution was not reported).

There were no deaths in any group. Thus the  $IC_{50}$  was >1.180 mg/l. Because this was reportedly the maximum attainable concentration, the limit test was satisfied and no other dose groups were needed. Compound-related clinical signs included salivation, lacrimation, ocular and nasal irritation, dry crusty skin about the eyes and ears, decreased activity, and periorbital alopecia. All of these signs reversed within three days except for the skin lesions and alopecia, which had nearly reversed by day 14. Weight gain was similar in the dosed and control groups. Gross findings of dark red lungs were reported in 4/10 males and 2/10 females. These lesions could have been caused by either the test article or  $CO_2$  euthanasia.

Particle size distribution was measured but was not reported, so it was impossible to assess the aerosol's dispersion, or what portion of the aerosol was respirable. Nevertheless, only mild toxicity was seen at this maximum attainable concentration. This study received Quality Assurance review. The Quality Assurance Officer stated that this study was in violation of the Good Laboratory Practice regulations since, "The mixture of the test substance with the carrier was not analyzed for homogeneity, stability, or concentration of the test substance." Since a carrier (vehicle) was not used by the laboratory, these analyses are not necessary and the study did not violate the Good Laboratory Practice regulations. The test article, a formulated powder, was used as prepared by the manufacturer. Thus, the stability of the powder was known, and homogeneity and concentration should not have varied appreciably.

## PRIMARY EYE IRRITATION STUDY OF BAYTHROID 20% WP IN RABBITS

Mobay Corporation; Report No. 735; April 14, 1986; Accession No. 264524

PROTOCOL: Six New Zealand White rabbits were dosed by placing 100 mg of Baythroid 20% WP (20.8% cyfluthrin) into the conjunctival sac of the left eye. The eyelids were then held together for 1 second. None of the eyes were rinsed. The right eyes served as controls. The eyes were evaluated for irritation at 1, 24, 48, and 72 hours, and at 7 days. No other observations were made. Food and water were available ad libitum.

RESULTS: No corneal lesions were seen in any of the rabbits. Sluggish irises were seen in two rabbits at 24 hours; these lesions had reversed by the 48 hour evaluation. Conjunctival lesions were seen after 1 hour in all rabbits and reversed by 72 hours in one rabbit, and by day 7 in the other rabbits. These lesions included injected to beefy red vessels, chemosis which ranged from slight to lids swollen half closed, and slight to considerable discharge.

STUDY CLASSIFICATION: This study is CORE MINIMUM, Toxicity Category III.

There was no mention of the age, weight, or sex of the rabbits used. This study received Quality Assurance review.

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## PRIMARY DERMAL IRRITATION STUDY OF BAYTHROID 20% WP IN RABBITS

Mobay Corporation; Report No. 742; April 14, 1986; Accession No. 264524

PROTOCOL: Six New Zealand White rabbits were each dosed with 500 mg of Baythroid 20% WP (20.8% cyfluthrin) applied to a 6 cm² area on their shaved backs and sides. The doses were occluded with gauze, hypoallergenic tape, sheet plastic, and an elastic bandage. They were also fitted with plastic collars. The doses were removed after 4 hours by wiping the dosing sites with water moistened paper towels.

The dosing sites were evaluated for irritation 0.5, 1, 24, 48, and 72 hours after dose removal. Food and water were available ad libitum.

RESULTS: One rabbit had very slight erythema 24 hours after dose removal. No other irritation was seen. Thus, the test article was a mild irritant.

STUDY CLASSIFICATION: This study is CORE MINIMUM, Toxicity Category IV.

There was no mention of the age, weight, or sex of the rabbits used. This study received Quality Assurance review.

# DERMAL SENSITIZATION [BUEHLER TOPICAL CLOSED-PATCH] STUDY OF BAYTHROID 20% WP IN GUINEA PIGS

Mobay Corporation; Report No. 755; June 9, 1986; Accession No. 264524

PROTOCOL: Twenty male Hartley guinea pigs were used in this study. Five of these served as noninduced controls, and the remaining fifteen were given induction treatments with aqueous solutions of Baythroid 20% WP. [NOTE: The report described using Baythroid 50% WP on page 8, but the report abstract mentioned the use of Baythroid 20% WP. Presumably, the page 8 reference was a typographical error which was overlooked by all editors.] The induced animals were given 0.5 ml topical doses of a 50% solution on day 0, and a 25% solution on days 7 and 14. The doses were applied onto the shaved skin of the left flank, and occluded for 6 hours with gauze, plastic sheeting, and an elastic bandage. These animals then had a 2-week rest period. Food and water were available ad libitum.

On day 28, the induced and noninduced control animals were challenged with the topical application of a 25% test article solution on the left flank. An equal volume of deionized water was applied to the right flank. The dosing sites were occluded for 24 hours. The animals were evaluated for skin irritation 24 and 48 hours after application of the induction doses, and 48 and 72 hours after application of the challenge doses.

RESULTS: None of the 15 guinea pigs had any skin irritation following the three induction doses. When these same animals were challenged, they also had no reaction. None of the noninduced controls had any reaction to the challenge doses. Thus, Baythroid 20% WP did not cause a sensitizing reaction in this study. Body weight gain in the induced and control animals was similar.

STUDY CLASSIFICATION: This study is CORE MINIMUM. This study received Quality Assurance Review. It was submitted to the EPA although the Quality Assurance Officer declared it to be in violation of the Good Laboratory Practice requirements for analysis of test article homogeneity, stability, and concentration.

TOXICOLOGY BRANCH ADI PRINTOUT

Cyfluthrin (Baythroid) 2yr feeding- rat

ADI = 0.025000 mg/kg/day

Date: 01/15/87

NOEL = 2.5000 mg/kgSafety Factor = 100

Caswell #266E CFR No. 180. LEL = 7.5000 mg/kg

Status: TOX complete 3/14/86. ORD verified 4/8/86.

RESIDUE CONTRIBUTION OF PUBLISHED TOLERANCES

FOOD PETITION TOLERANCE

FACTOR MG/DAY NUMBER (PPM) CROP

No published tolerances listed in file.

## RESIDUE CONTRIBUTION OF TOX-APPROVED TOLERAN

	TOLERANCE	PETITION	FOOD	
CROP	(PPM)	NUMBER	FACTOR	MG/DAY
Apples	2.000	6G3307	2.53	0.075900
	3.000	5G3193	0.10	0.004500
	1.000	5G3193	0.03	0.000450
	1.000	5 <b>G</b> 3193	0.74	0.011100
	2.000	5G3193	0.07	0.002100
	0.040	4G3126	1.43	0.000858
	2.000	4F3046	0.15	0.004500
	0.040	4G3126	1.00	0.000600
	0.050	4F3046	10.81	0.008108
· · · · · · · · · · · · · · · · · · ·	1.950	4G3126	10.81	0.316192
•	0.000	6G3307	10.81	0.000000
	0.010	4F3046	28.62	0.004293
	0.190	4G3126	28.62	0.081567
	0.000	6G3307	28.62	0.000000
Pears	1.000	6G3307	0.26	0.003900
Potatoes	0.020	4G3126	5.43	0.001629
	Apples Broccoli Brussel sprouts Cabbage, sauerkraut Cauliflower Corn, sweet Cottonseed (oil) Corn, grain (field corn) Meat, red Meat, red Meat, red Milk and dairy products Milk and dairy products Milk and dairy products Pears	CROP       (PPM)         Apples       2.000         Broccoli       3.000         Brussel sprouts       1.000         Cabbage, sauerkraut       1.000         Cauliflower       2.000         Corn, sweet       0.040         Cottonseed (oil)       2.000         Corn, grain (field corn)       0.040         Meat, red       0.050         Meat, red       1.950         Meat, red       0.000         Milk and dairy products       0.010         Milk and dairy products       0.190         Milk and dairy products       0.000         Pears       1.000	Apples 2.000 6G3307 Broccoli 3.000 5G3193 Brussel sprouts 1.000 5G3193 Cabbage, sauerkraut 1.000 5G3193 Cauliflower 2.000 5G3193 Corn, sweet 0.040 4G3126 Cottonseed (oil) 2.000 4F3046 Corn, grain (field corn) 0.040 4G3126 Meat, red 0.050 4F3046 Meat, red 1.950 4G3126 Meat, red 0.000 6G3307 Milk and dairy products 0.010 4F3046 Milk and dairy products 0.190 4G3126 Milk and dairy products 0.190 4G3126 Milk and dairy products 0.000 6G3307 Pears 1.000 6G3307	CROP         (PPM)         NUMBER         FACTOR           Apples         2.000         6G3307         2.53           Broccoli         3.000         5G3193         0.10           Brussel sprouts         1.000         5G3193         0.03           Cabbage, sauerkraut         1.000         5G3193         0.74           Cauliflower         2.000         5G3193         0.07           Corn, sweet         0.040         4G3126         1.43           Cottonseed (oil)         2.000         4F3046         0.15           Corn, grain (field corn)         0.040         4G3126         1.00           Meat, red         0.050         4F3046         10.81           Meat, red         0.000         6G3307         10.81           Meat, red         0.000         6G3307         10.81           Milk and dairy products         0.010         4F3046         28.62           Milk and dairy products         0.190         4G3126         28.62           Milk and dairy products         0.000         6G3307         28.62           Pears         1.000         6G3307         0.26

TMRC 0.008595 mg/kg/day (60kg BW, 1.5kg diet)

% ADI 34.379800

### RESIDUE CONTRIBUTION OF NEW (PENDING) TOLERANCES

CROP	TOLERANCE (PPM)	PETITION NUMBER	FOOD FACTOR	MG/DAY
197 All foods	0.050	6H5515	100.00	0.075000000

TMRC 0.009845 mg/kg/day (60kg BW, 1.5kg diet)

% ADI 39.379800

SUMMARY OF CYFLUTHRIN TOXICITY DATA and EIGHT POINT FREE-STANDING SUMMARY

1. Summary of selected toxicology data considered for these actions:

VIII MAN	RESTITING.	TOXICITY	CLASSIFICATION
Slubi Technical Cyfluthrin Data:			
Acute Oral LD50, Rat	Depending on the vehicle used, LD50 values ranged from 16.2 mg/kg (cremophor/water) to 1271 mg/kg (PEG 400)	II-I	Minimum
Acute Dermal LD50, Rat	${ m LD}_{50}$ >5,000 mg/kg, males and females	III	Minimum
Acute Inhalation ${\rm IC}_{50}$ , Rat	LC <sub>50</sub> >0.735 mg/l, males (4-hour) LC <sub>50</sub> 0.200-0.735 mg/l, females (4-hour)	II	Minimum
Primary Eye Irritation, Rabbit	Mild irritation	III	Minimum
Primary Dermal Irriatation, Rabbit	No irritation	VI	Minimum
Dermal Sensitization, Guinea Pig	Not a sensitizer by: the Draize Test the Maximization Test		Minimum Guideline
Teratology, Rat	<pre>Maternal NOEL = 3 mg/kg/day Maternal LEL = 10 mg/kg/day (behavioral changes in gait and coordination) Fetotoxic NOEL &gt;30 mg/kg/day (HDT) Teratogenic NOEL &gt;30 mg/kg/day (HDT)</pre>		Minimum
Teratology, Rabbit	Maternal NOEL = 15 mg/kg/day Maternal LEL = 45 mg/kg/day (abortion, resorption) Fetotoxic NOEL >45 mg/kg/day (HDT) Teratogenic NOEL >45 mg/kg/day (HDT)	6	Minimum

STUDY	TOXICITY RESULTS CATEGORY	CLASSIFICATION
Mutagenicity Studies:		·
A. Gene Mutation Test:		
CHO/HGPRT Mutation	Negative	Acceptable
B. Structural Chromosome Aberration Test:		
Sister Chromatic Exchange	Negative	Acceptable
<pre>C. Tests for Other Genotoxic Effects:</pre>		
Unscheduled DNA Synthesis	Negative	Acceptable
Metabolism	Blood levels of cyfluthrin isomers are higher and peak more quickly when cyfluthrin is administered in cremophor/distilled water than when administered in polyethylene glycol [sic].	Minimum
Chronic Feeding/Oncogenicity, Rat	Oncogenic NOEL >22.5 mg/kg/day (HDT) Systemic NOEL = 2.5 mg/kg/day Systemic LEL = 7.5 mg/kg/day (decreased body weights in males, inflammatory foci in kidneys of females).	Minimum
Chronic Feeding, Dog	NOEL = 4 mg/kg/day LEL = 16 mg/kg/day (slight ataxia, increased vomiting, diarrhea, and decreased male body weights).	Minimum
Oncogenicity, Mouse	Oncogenic NOEL >120 mg/kg/day (HDT) Systemic NOEL <7.5 mg/kg/day (LDT, increased alkaline phosphatates activity in males) [sic].	Supplementary for chronic feeding Minimum for oncogenici
	-2-	5723

SIUDY	TOXICITY RESULTS CATEGORY	CLASSIFICATION
3-Generation Reproduction, Rat	Reproductive NOEL = 2.5 mg/kg/day Reproductive LEL = 7.5 mg/kg/day (decreased viability) Systemic NOEL = 2.5 mg/kg/day Systemic LEL = 7.5 mg/kg/day (decreased pup body weights)	Minimum
Neurotoxicity, Hen	<ol> <li>Delayed Neurotoxicity Study Cyfluthrin was mildly neurotoxic at 4300 mg/kg/day X2, but did not cause the classic delayed neurotoxic signs seen in hens dosed with TOCP.</li> <li>Neurotoxic Esterase Activity NTE activity in hens dosed with 4300 mg/kg/day X1 of cyfluthrin resembled that of the vehicle controls.</li> </ol>	Minimum
Neurotoxicity, Rat	Wistar Bor:WISW rats given 14 oral doses of 50 or 60 mg/kg/day had non-specific disturbed behavior, rolling, tremors, stretched gait, uncoordinated gait, salivation, phonation, weight loss (males), and death. Histopathologic lesions included slight brain hemorrhages and necrosis of the skeletal muscle fibers.	Guideline
Neurotoxicity, Rat	Male SD rats given oral doses of 80 mg/kg/day for 5 days, then 40 mg/kg/day for 9 days had straddled gait, slow leg movement, titubation, salivation, red tears, and reduced weight gain. Histopathologic lesions included axonal degeneration of the sciatic nerve (light microscopy); and microtubular dilatations with proliferation of neurofilaments and mitochondria degeneration in the sciatic and femoral nerves (electron microscopy).	Guideline 002

Summary of Data Considered Desirable but Lacking for This Action: 2

None

3. Action Being Taken to Obtain the Lacking Information or Other Additionally Needed Information:

Not applicable.

4. A Summary of Other Permanent Tolerances Granted for This Herbicide:

None

5. The current commitment for Toxicology Branch approved tolerances is 34.38% of the ADI with a TMRC of 0.0086 mg/kg/day (60 kg body weight, 1.5 kg diet). Granting these tolerances will increase the % ADI to 39.38%, and mg/kg/day (60 kg body weight, 1.5 kg diet). the TMRC to 0.0098 mg/kg/day.

The 2-year chronic feeding/oncogenicity study in rats with a NOEL of 2.5 mg/kg/day (50 ppm) and a safety factor of 100 were used to set the ADI (0.025 mg/kg/day). •

7. There are at this writing no pending regulatory actions against the registration of this pesticide.

8. Other Relevant Considerations in Setting These Tolerances:

None.

250

120H