

US EPA ARCHIVE DOCUMENT

2-26-91



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

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OFFICE OF
PESTICIDES AND TOXIC SUBSTANCES

MEMORANDUM

SUBJECT: EPA Reg. No./File Symbol 121-UA Evergreen
Cutter Insect Repellent Formula MMII

FROM: William S. Woodrow WSW 2-6-91
Precautionary Review Section E 2/26/91
Registration Support Branch
Registration Division (H7505C)

TO: Phillip Hutton / Joseph Tavano (PH 17)
Insecticide - Rodenticide Branch
Registration Division (H7505C)

APPLICANT: Miles Inc.
Consumer Household Products Div.
7123 West 65 Street
Chicago, IL 60638

FORMULATION FROM LABEL:

Active Ingredient(s):	by wt.
<u>Deet (N,N-diethyl-m-toluamide)</u>	<u>28.50</u>
<u>other isomers</u>	<u>1.50</u>
_____	_____
_____	_____
<u>Inert Ingredient(s):</u>	<u>70.00</u>
Total	100.00

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1.71

BACKGROUND

Miles, Inc. submitted primary eye and skin and dermal sensitization studies to support registration of Evergreen Cutter Insect Repellent Formula N.N. II (EPA Reg. No. 121-UA) P.R.I.D. NOS. used were 416933-01, 415757-02, and 415757-03. This product is to be sprayed from pressurized cans from 6-to-8 inches away from skin or clothing, keeping nozzle pointed away from the face.

RECOMMENDATION

- 1) The primary eye and skin irritation and the dermal sensitization studies submitted by Miles are acceptable to RSB/PRS.
- 2) A requirement for acute oral, acute dermal, and acute inhalation studies to support ^{121-UA} registration remains. However, the Dert Pesticide Registration Standard published by the Agency in December, 1980, does address acute toxicity data requirements for Dert Ready-to-use and Pressurized Liquid products:

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a. From page 42 of the Deet Standard :
Quote — Acute oral toxicity)

(2) Pressurized Liquid (PrL)

Because technical Deet and the 15% and 30% PrL formulations are in Toxicity Category III, and because the inert ingredients in the PrL formulations are not expected to increase the acute oral toxicity potential, existing PrL products between 12% and 75% concentrations, except those that contain freon propellants, are classified in Toxicity Category III. Toxicity Category III corresponds to a low acute oral toxicity potential. //

b. From page 43 — Deet Standard
Quote " (2) Pressurized Liquid (PrL) — (Acute dermal tox).

Because technical Deet is placed in Toxicity Category III, and because the inert ingredients in PrL products are not expected to increase the acute dermal toxicity potential, all existing PrL products between 12% and 75% concentrations, except those that incorporate freon propellants, are classified in Toxicity Category III. Toxicity Category III corresponds to a low acute dermal toxicity potential. //

c. From page 45 — Deet Standard — Acute
Inhalation Toxicity.
Quote

(2) Pressurized Liquids (PrL)

Because technical Deet is placed in Toxicity Category IV, and because the inert ingredients in PrL products are not expected to increase the acute inhalation toxicity potential, all existing PrL products between 12% and 75% concentrations, except those that incorporate freon propellants, are placed in Toxicity Category IV. Toxicity Category IV, corresponding to a very low acute inhalation Toxicity potential.

Thus, according to the Deet Standard it

will not be necessary to submit additional acute oral, acute dermal, or acute inhalation studies Evergreen Cutter Insect Repellent Formula MM II (pressurized), EPA Reg. No. 121-UA.

3) Current acute toxicity profile for Evergreen Cutter Insect Repellent (EPA Reg. No. 121-UA):

study	MRID/acc. [#]	Tox. Cat.	Grade
Eye irritation	415757-02	III	Guideline
Skin irritation	415757-03	IV	Guideline
Skin sensitization	416953-01	Not a sensitizer	Guideline
Acute oral	(Deet std.)	II	-
Acute dermal	(Deet std.)	III	-
Acute inhalation	(Deet std.)	IV	-

No additional acute toxicity studies for EPA Reg. No. 121-UA are required.

LABELING

1) Change the product signal word from WARNING, to read "CAUTION".

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2) The Precautionary Statements and
statement of Practical Treatment are
acceptable.

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DATA REVIEW FOR ACUTE EYE IRRITATION TESTING (§81-4)

00856

Product Manager: (17)

Reviewer: Woodrow M. Waller

MRID No.: 465755-01

Report Date: 1-2-91

Testing Laboratory: Tox Dept., Miller, Inc.

Report No. MTD0164

Author(s): R.W. Fowlesky, R.E. Hartnagle

Species: Rabbit, M 7 White

Sex: 9 females, 16 mos. old

Weight: 3.5 - 4.7 kgs

Source: Miller Cooperative animal facility

Dosage: 1 - 5 sec spray

Test Material: Evergreen Cutter insect repellent Formula M118

Quality Assurance (40 CFR §160.12): adequate

Summary:

Tox. Category: III Classification: Quadr. lines

Procedure (~~Division from test~~) Eyes tested for defects using Fluorescein

prior to study. Test eyes anesthetized prior to treating; aliquots (by spraying) of approx. 0.023g, 23mg, or 0.27ml were administered as a single burst of 1 second from a

Results: NOTE: washed & unwashed eye results are similar & so scoring has recorded as 3+6 = 9 animals

Observations

	(number "positive"/number tested)							
	Hour	Days						
	1	1	2	3	4	7	14	21
Cornea Opacity	9/9	9/9	9/9	8/9		0/9		
Iris	0/9	0/9	0/9	0/9		0/9		
Conjunctivae Redness	9/9	9/9	9/9	6/9		0/9		
Chemosis	9/9	9/9	9/9	8/9		0/9		
Discharge	9/9	9/9	3/9	4/9		0/9		

1.0 scores

1.0 scores

1/2/3.0 scores

3/2/1.0 scoring

Comments: Distance of 10cm, directly in front of treated eye. Dose estimated by weighing can before and after application. Eye lids held together 1 second - Treated eyes of these animals washed by irrigation = water for 1 minute, beginning 20-30 sec
Corneal involvement or irritation clearing in 7 days or less

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post exposure. Each animal observed daily for up to 7 days, and were graded for infection according to ~~serology and histology~~ ^{Dr. D. H. 27} at 1 hour, 1, 2, 3 and 7 days post exposure, using the scale devised by Davies.

DATA REVIEW FOR SKIN IRRITATION TESTING (1981-5)

Product Manager: (17)
 MRID No.: 415755-03
 Testing Laboratory: Tax Dept. Miles, Inc.
 Author(s): R.L. Kowalski, R.E. Hartnagle
 Species: Rabbit, NZ White
 Age: 10.5 months
 Sex: not given
 Weight: 3.5 - 4.2 kg
 Dosage: 0.5ml
 Test Material: Evergreen Cutter Insect Repellent Formula MARI
 Quality Assurance (40 CFR §160.12): adequate

Reviewer: Woodrow
M. Waller
 Report Date: 1-2-91
 Report No.: MTD 0146

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Summary:

The Primary Irritation Index = 1.37

Toxicity Category: IV

Classification: Guidelines

Procedure (~~Domestic Form 501-57~~): 0.5ml test material applied to clipped, depilated dorsal area of skin of 6 rabbits each, under a Weibull pad, held in place by Blenderm tape. Entire back covered with occlusive Dressing secured with tape. 4 hour skin contact, Dressings removed, no test material evident.
 Results: Each animal observed 3 days; test sites examined & scored according to Orange for erythema/edema.

Results: All six rabbits showed total clearing of irritation by day 10 (for erythema); 2 of the treated animals showed no scores on days 2 and 3, all other erythema scores (to and through day 6) were 1.0. 4/6 animals showed slight (inc) swelling for edema initially, absent by day 2.
 Primary irritation index = (mean)

Special Comments:	time	mean P.I. Scores	
		erythema	edema
	1 hr.	0.83	0.67
	24 hrs	1.0	0.33
	48 hrs	1.33	0.0
	72 hrs	1.33	0.0

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DATA REVIEW FOR SKIN SENSITIZATION TESTING (§81-6)

00867

Product Manager: (17)

Reviewer: Woodrow H. Walter

MRID No.: 416953-01

Report Date: 1-2-71

Testing Laboratory: Tex. Dept. of Males, Inc.

Report No. MTD 0136

Author(s): R.L. Kovalick, R.E. Hartnagle

Species: Guinea Pig, Hartley

Sex: male

Weight: 469-694g

Source: Chickler Russ

Test Material: Evergreen Cutter Insect Repellent Formula MM 21

Positive Control Material: 1-chloro-2,4-dinitrobenzene (DNCB)

Quality Assurance (40 CFR §160.12): adequate

Method: Modified Draeg, intradermal injection

Summary:

1. This product is / is not a dermal sensitizer
2. Classification: Guidelines

Procedure (Deviation From §81-6): A preliminary screening to determine the maximum non-irritation product concentration for use in the main study was conducted (ID test). 5%, 10% (v/v in corn oil), minimal irritation @ 1% v/v in corn oil, 0.05, 0.10, 0.25;

Results: 0.5 and 1.0% v/v in corn oil; 0.5% v/v in corn oil used for main study. These results indicated 0.5% mix gave same results as corn oil alone. Positive control study employed 1-chloro-2,4-dinitrobenzene (DNCB). DNCB formulated as 0.05% w/v ad in 5.0% v/v ethanol in normal saline.

10 males induced to test mat, 5 naive controls. DNCB induced 4 naive control, 5 animals each. (25 animals total).

Induction: 10 test mat. + 5 DNCB animals clipped & depilated 2 days before test. 10 animals injected intradermally = 0.05ml test (0.5% v/v in corn oil) near left pelvic girth, on days 0, 2, 4, 7, 9, 11, 14, 16, 18, 21 over 21 day period. After 1st injection, 0.1ml injected in staggered pattern. 5 DNCB animals induced using 0.05% w/v DNCB in 5.0% v/v EtOH in normal saline; these (DNCB) injected using same repeat pattern for 10 injections. Induced animals rested 2 weeks

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prior to challenge injection.

Challenge - Diptheria not used. Test material injected as 0.1 ml in right (naive site) scapula, and also injected at a naive site with 0.1 ml even oil DNCB animals similarly challenged (induced and naive animals) with 10 injections of 0.1 ml DNCB near right scapula (naive sites), and also challenged at naive sites with 0.1 ml of other oil/saline vehicle. (Test material also administered to 5 naive animals) Each injection and challenge site scored for erythema & edema @ 24 and 48 hours post injection. Body wts. days 0 & 37 (termination).

Note: Tests conducted with product formula - minus the propellant.

Dosing scale 0-3.

Results: All test animals gained weight, all test naive animals gained weight. All induced and non-induced DNCB animals gained weight.

induced

Test (induced): 24 hrs - severity index 0.7-1.0, 48 hrs - severity index 0.8-0.8
DNCB (induced): 24 hrs - severity index 0.4-1.0, 48 hrs - 0.3-0.6
Test (naive): 24 hrs test → 0.7, vehicle → 0.8, 48 hrs test 0.8, vehicle 0.7.

DNCB (naive): 24 hr challenge severity score 0.4, 48 hr 0.5
vehicle 0.0 @ 24 & 48 hrs

DNCB challenged - 24 hrs severity index = 2.6, vehicle = 0.0
48 hrs = 2.6, vehicle = 0.0

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Conclusions:

1. Test material did not sensitize guinea pigs
2. DNCB control (positive control) chemical did sensitize guinea pigs.

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