

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

006057

OFFICE OF CIDES AND TOXIC SUBSTANCES MAY 16 1986

E -127/8

#### **MEMORANDUM**

SUBJECT: EPA File Symbol 39702-G

Muralo Lumber Jacket Stain & Wood Preservative

FROM:

Deloris F. Graham Day 5/37/86 Technical Support Section Fungicide-Herbicide Branch

Registration Division (TS-767C)

TO:

Henry M. Jacoby, PM 21 Fungicide-Herbicide Branch Registration Division (TS-767C)

Applicant:

The Muralo Company, Inc.

148 East 5th Street Bayonne, NJ 07002

Active Ingredients:

Bis(tribulyltin)Oxide Folpet N-(Trichloromethyl) Thiophthalimide Inert Ingredients: .

# Background:

Submitted Acute Oral, Acute Dermal, Acute Inhalation, Eye Irritation, and Primary Dermal Irritation Studies. Studies conducted by Food and Drug Research Laboratories, Inc. Data under Accession Numbers: 259664, 259665, 259666, 259667, and 259668. Method of support not indicated.

# Recommendation:

FHB/TSS finds these data acceptable to support conditional registration of this product.

- 2. A Dermal Sensitization Study was not submitted and one must be submitted or data to support waiver.
- 3. The appropriate signal word is CAUTION.

## Label:

- The signal word "CAUTION" must appear on center front panel of label.
- The word "Combustible" should appear under the heading "Physical and Chemical Hazards" not following signal word.
- Precautionary statements must precede general instructions and directions for use.
- 4. Precautionary statements must be revised similar to the following "CAUTION. Harmful if absorbed through skin, inhaled and if in eyes."
- Upon submission of dermal sensitization data additional labeling may be necessary.

# Review:

(1) Eye Irritation Study: Food and Drug Research Laboratories, Inc.; FDRL Study No. 8645A; August 15, 1985; EPA Accession No. 259664.

### Procedure:

Six rabbits received 0.1 ml of the test material in one eye each. Observations made for 72 hours posttreatment.

# Results:

At 1 hour, 5/6 conjunctive redness (5/6 = 1). At 24 hours, 2/6 redness (2/6 = 1). Irritation had cleared by 48 hours.

Study Classification: Core Guideline Data.

Toxicity Category: III - CAUTION.

(2) Primary Dermal Irritation Study: Food and Drug Research Laboratories, Inc.; FDRL Study No. 8645A; August 15, 1985; EPA Accession No. 259665.

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#### Procedure:

Six rabbits with two intact skin sites each were treated with 0.5 ml of the test material under occlusive wrap for 4-hour exposure period. Observations made for 14 days posttreatment.

## Results:

At 24 hours, 6/6 slight erythema (scores of 1) and edema (scores of 1). At 72 hours, 6/6 slight erythema (scores of 1) and 5/6 edema (scores of 1). Irritation persisted through day 14 in a few animals. Test site reported to appear dry and slightly cracked beginning at day 7. Mean Primary Irritation Score reported to be 1.95.

Study Classification: Core Guideline Data.

Toxicity Category: IV - CAUTION.

(3) Acute Dermal Toxicity Study: Food and Drug Research Laboratories, Inc.; FDRL Study No. 8654A; August 23, 1985; EPA Accession No. 259666.

# Procedure:

Five male and five female rabbits with intact skin sites each were treated with 2.0 g/kg of the test material under occlusive wrap for 24-hour exposure period. Observations made for 15 days posttreatment. Necropsy performed on all animals.

# Results:

No mortalities or abnormalities at necropsy reported. Toxic signs reported included decreased activity, diarrhea, anorexia, and soft stools.  $LD_{50}$  reported to be greater than 2.0 g/kg.

Study Classification: Core Guideline Data.

Toxicity Category: III - CAUTION.

(4) Acute Oral Toxicity Study: Food and Drug Research Laboratories, Inc.; FDRL Study No. 8645A; August 21, 1985; EPA Accessior. No. 259667.

### Procedure:

Five male and five female rats received a single 5.0 g/kg dose of the test material orally. Observations were made for 15 days postdosing. Necropsy performed on all animals.

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## Results:

No mortalities or abnormalities at necropsy reported. Toxic signs reported included wet abdomen in 5/5 F and sores in rectal area in 1/5 M. LD<sub>50</sub> reported to be greater than 5.0 g/kg.

Study Classification: Core Guideline Data.

Toxicity Category: IV - CAUTION.

(5) Acute Inhalation Toxicity Study: Food and Drug Research Laboratories, Inc.; FDRL Study No. 8645A; August 19, 1985; EPA Accession No. 259668.

## Procedure:

Five male and five female rats were exposed whole body for 4 hours to a 0.9 mg/L liquid droplet aerosol determined gravimetrically (nominal concentration = 5.4 mg/L). Mass median aerodynamic diameter 3.8 um with 2.2 geometric standard deviation. Average temperature reported to be 26 °C and relative humidity 64%. Observations made for 15 days postexposure. Necropsy performed on all animals.

## Results:

No mortalities or abnormalities at necropsy reported. Toxic signs reported included labored breathing and/or rales; decreased activity; dried material on fur of test animals; skin sores; alopecia.

Study Classification: Core Guideline Data.

Toxicity Category: III - CAUTION.