

Subject: EPH Registration Number - 7501-42
Gustafson Vernon - The Good Protestant

From: Delius, J. Triakam

THE/ISS

E 8/1/P4

005195

To: Henry Jacoby
Product Manager (21)

Applicant: Gustafson, John
P.O. Box 660065
Dallas, Texas 75266-0065

Arthur Ingham

the tetracycl: N-(2,6-dimethylphenyl)-
-N-(methoxycarbonyl) vaniline methyl
ester

25.35%

most important

71.65%

Exel generated & submitted by top 10 for
accuracy of acute stimulation data and LOSO
and 75% confidence intervals for acute renal
clearance. Data were recorded on October 20, 1983.
Method of support not indicated.

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Reinhold

① Based on the 2001 information submitted for Great Bear River, the appropriate treaty category for this species is III CHITWAN.

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(2) Based on the information attached to letter dated March 9, 1984 from the Company and information received on acute inhalation October 20, 1983, it is concluded that there would be minimal exposure through inhalation route.

005195

Label:

(1) No additional labeling comments.

- References:
1. Acute Oral Toxicity Study: Product Safety Labs; Report No. T-3030; May 6, 1983.

Procedure: Five male and five female rats weighing between 200 and 300 grams received a single dose of 5.0 g/kg. Observations were daily for 14 days after treatment. Necropsy performed on all animals.

Results: 2/5 M and 5/5 F died. Lethargy was the only toxic sign noted in the 3/5 M survivors. Necropsy revealed discolored liver and spleen; intestinal hemorrhage; pulmonary hemorrhage. ~~LD50 less than 5.0 g/kg.~~

LD50 expected to be 2.9 g/kg with 95% confidence limits between 1.8 and 4.6 g/kg.

Study Classification: Carcinogenic Data.

Toxicity Category: III - (MUTAGEN)

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METALAXYL

Page 3 is not included in this copy.

Pages _____ through _____ are not included.

The material not included contains the following type of information:

- ☐ Identity of product inert ingredients.
 - ☐ Identity of product impurities.
 - ☐ Description of the product manufacturing process.
 - ☐ Description of quality control procedures.
 - ☐ Identity of the source of product ingredients.
 - ☐ Sales or other commercial/financial information.
 - ☒ A draft product label.
 - ☐ The product confidential statement of formula.
 - ☐ Information about a pending registration action.
 - ☐ FIFRA registration data.
 - ☐ The document is a duplicate of page(s) _____.
 - ☐ The document is not responsive to the request.
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The information not included is generally considered confidential by product registrants. If you have any questions, please contact the individual who prepared the response to your request.
