MEMORANDUM

SUBJECT: Simazine - Mutagenicity Data Submitted under MRID Nos. 414429-01 and -02
EPA ID #100-541

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Health Effects Division (H7509C)

Registrant: Ciba-Geigy Corporation, Greenboro, NC

Request

Review and evaluate the following (two) mutagenicity studies, submitted as additional data for the reregistration of simazine, and both performed in the genetic toxicology laboratories of Ciba-Geigy Ltd., Basle (Switzerland):

1. Simazine Technical: Structural Chromosomal Aberration Test - Mouse Micronucleus Test (MT)


TB Conclusions:

[Detailed reviews are appended to this memorandum].

<table>
<thead>
<tr>
<th>Study</th>
<th>Reported Results</th>
<th>TB Evaluation</th>
</tr>
</thead>
<tbody>
<tr>
<td>(1) MT</td>
<td>Negative for inducing micronuclei in bone marrow cells of mice treated by acute oral gavage up to 5000 mg/kg.</td>
<td>ACCEPTABLE</td>
</tr>
<tr>
<td>(2) DNA</td>
<td>Although reported as negative in repeat assays at levels up to 170 ug/ml, no evidence was presented that a cytotoxic or solubility limit was tested.</td>
<td>UNACCEPTABLE</td>
</tr>
</tbody>
</table>

ATTACHMENTS (DERs)
DATA EVALUATION REPORT

I. SUMMARY

MRID (ACC) No.: 41442901
ID No.: 100-541
RD Record No.: 262.860
Caswell No.: 740
Project No.: 0-1092

Study Type: Mutagenicity - Chromosomal aberrations in vivo (micronucleus)

Chemical: Simazine

Synonyms: G-27-692 Technical

Sponsor: Ciba-Geigy, Basle

Testing Facility: Ciba-Geigy, Basle

Title of Report: Structural Chromosomal Aberration Test - Micronucleus Test, Mouse

Author: Carla Ceresa

Study No.: 880.89

Date of Issue: September 15, 1988

TR Conclusions:

Negative for inducing micronuclei in bone marrow cells of mice treated by acute oral gavage up to 5000 mg/kg.

Classification (Core-Grade): ACCEPTABLE
II. DETAILED REVIEW

A. Test Material - G-27-692 technical (simazine)

Description: (Not stated)
Batch (Lot): 2001SP
Purity (%): 99.6
Solvent/carrier/diluent: Carboxymethylcellulose, 0.5% (CMC)

B. Test Organism - Rodent

Species: Mouse
Strain: Tif: NAG F/SPF (NMRI-derived)
Age: (Adult)
Weights - males: 27 to 38 g
females: 22 to 33 g
Source: Ciba-Geigy Tierfarm, Sisseln (Switzerland)

C. Study Design (Protocol) - This study was designed to assess the clastogenic potential of simazine when administered by oral gavage to mice, according to internationally-accepted test guidelines (OECD/EEC/FPA).

A statement affirming compliance with Agency GLPs was provided.

A Statement of Quality Assurance measures (inspection/audits) was also provided.

D. Procedures/Methods of Analysis - Following preliminary toxicity testing, the test article was administered by oral gavage to groups of male and female mice according to two schedules: (i) once at 5000 mg/kg, and animals sacrificed 6, 24, and 48 hours later; (ii) once, at doses of 1250, 2500, and 5000 mg/kg, and animals sacrificed 24 hours later. In addition to concurrent solvent controls for each one of the sacrifice times (0.5% CMC), groups of 8 males: 8 females received the clastogen, cyclophosphamide (CP, 64 mg/kg) and were sacrificed 24 hours later.

At the scheduled sacrifice times, bone marrow was collected from both femurs of each animal, and prepared for microscopic examination by conventional cytological techniques. Coded slides from 5 animals/sex/experimental group were examined under oil-immersion, and 1000 polychromatic erythrocytes (PCE) per animal scored for the presence of micronuclei; in addition, ratios of PCE to normochromatic erythrocytes (NCF) were determined for each animal.
The data were analyzed by Chi-square, with the level of significance set at $p \leq 0.05$.

E. Results - In neither experiment were any significant differences in incidences of micronuclei between simazine-treated (0.02 to 0.08%) and solvent control (0.06 to 0.08%) animals recorded (Report Table 2 to 8, appended to this DER). By contrast, highly significant increases ($p < 0.05$) were registered in both CP-treated positive controls (1.81 and 2.02%). No cytotoxicity by simazine treatment was found in bone marrow cells, since the ratios of PCE to NCE were unaffected (Tables 2 to 8); finally, no clinical effects of treatment were reported.

F. TR Evaluation - Acceptable. This study appears to have been properly conducted according to current testing guidelines under conditions ensuring appropriate testing of the test substance for clastogenic potential when administered acutely by oral gavage up to the limit dose (5000 mcg/kg).

ATTACHMENTS (Data Tables)
ATTACHMENT I

Data Tables
SIMAZINE

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DATA EVALUATION REPORT

I. SUMMARY

MRID (ACC) No.: 41442902
ID No.: 100-541
RD Record No.: 262,860
Caswell No.: 740
Project No.: 0-1092

Study Type: Mutagenicity - DNA damage/repair in vitro (HFC/UDS)

Chemical: Trimazine

Synonyms: G-27-692 Technical

Sponsor: Ciba-Geigy

Testing Facility: Ciba-Geigy, Basle (Switzerland)

Title of Report: Tests for Other Genotoxic Effects - Auto-radiographic DNA Repair Test on Rat Hepatocytes.

Author: Thomas Hertner

Study No.: 891412

Date of Issue: December 7, 1989

TB Conclusions:

Although reported as negative for inducing unscheduled DNA synthesis (UDS) in rat hepatocytes treated up to 170 μg/mL, no evidence was presented that this represented a cytotoxic or solubility limiting concentration.

Classification (Core-Grade): UNACCEPTABLE
II. DETAILED REVIEW

A. Test Material - G-27-692 technical (simazine)

Description: (Not stated)
Batch (Lot): PL-850614
Purity (%): 96.9
Solvent/carrier/diluent: Dimethylsulfoxide (DMSO)

R. Test Organism - Rodent hepatocytes

Species: Rat
Strain: Tif:RAF (SPF)
Age: (Adult)
Weights - males (only): 170 to 350 g
Source: Ciba-Geigy Tierform, Sisseln (Switzerland)

C. Study Design (Protocol) - This study was designed to
assess the DNA-damaging potential of simazine as
determined by unscheduled DNA synthesis when administered
in vivo to primary rat hepatocyte cultures, according
to Internationally accepted test guidelines (OPCD, OECD,
EPA).

A statement affirming compliance with Agency SLPs was
provided.

A Statement of Quality Assurance measures (inspections/
audits) was also provided.

D. Procedures/Methods of Analysis - Hepatocytes were isolated
from adult male rats by established procedures, and
allowed to attach to coverslips immersed in multi-well
culture vessels containing appropriate culture medium.
Following preliminary cytotoxicity testing for dose
selection, cultures of attached cells are treated in
quadruplicate for 16 to 18 hours with test substance (at
least 6 preselected concentrations), or with vehicle
(DMSO), or with the mutagen 2-acetylaminofluorene (AAF,
45 μM); as well, untreated (culture medium only)
negative controls were run concurrently.

All cultures were treated concurrently with tritiated
thymidine (8 μCi ³H-TdR, spec. act. = 25 C/mmol)
following which the cells were prepared for autoradiography
with Ilford K-5 photographic emulsion on microscope
slides (under darkroom conditions) by standard procedures.
Slides were then placed in air- and light-tight boxes at
4 °C for 4 days, following which slides were stained
'hematoxylin-eosin) and made permanent.
Developed silver grains were counted over the nuclei and cytoplasm of 50 cells per slide (150 per treatment), by means of an electronic counter attached to a universal binocular microscope (magnification = 2000X). Net grain counts were calculated by subtracting the average grain count over three nuclear-sized cytoplasmic areas from the nuclear grain count, and all calculations summarized for each treatment.

Grain count data were analyzed statistically by Dunnett's one-tailed t-test, with significance set at \( p \leq 0.01 \). The following criteria were applied for determining the quality of response:

For a positive:

"The mean gross number and the mean net number of silver grains per nucleus in relation to their respective vehicle controls are significantly different at any concentration, and the mean net value is at least 2.0."

For a negative:

"The mean gross number and the mean number of silver grains per nucleus in relation to the vehicle control are not statistically different at any concentration and no concentration dependence can be seen."

The entire assay was repeated once ("confirmatory test").

V. Results - In the preliminary toxicity testing, the highest concentration considered "usable" was 170 μg/ml (Report Table 1, appended here).

In the initial repair test carried out at doses of 1.57, 4.72, 14.17, 42.50, 85, and 170 μg/L, none of the mean gross or net grain counts in treated cultures were statistically different from vehicle controls (Report Table 3, 7, 9, and 9, appended here). However, in comparison with the medium control, a slight shift to higher distribution values was noted for the three highest concentrations (Table 4, attached). In the confirmatory trial (using the same dose schedule), none of the grain values in simazine-treated cultures differed from vehicle or medium controls (Tables 5, 8, 10, and 11, also appended here).

*No explanation or definition was provided for this term.
In both trials, the positive control (AAF) responded as expected, with highly significant increased net silver grain counts 14 and 23 times solvent values.

The author concluded that the test substance gave no indication of inducing DNA damage in rat hepatocytes (as measured by the induction of UDS).

F. **TB Evaluation - UNACCEPTABLE.** Although apparently conducted with adequate controls under accepted test guidelines, no indication was given in the entire report that the HDT, 170 μg/mL represented a cytotoxic or solubility limit for the test substance, as evident by the preliminary cytotoxicity testing reported (Tables 1 and 2).

**ATTACHMENTS (Data Tables)**
ATTACHMENT I

Data Tables
SIMAZINE

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