Primary Eye Irritation Potential of DIGL-RP Solid Propellant in Rabbits

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MAMMALIAN TOXICOLOGY BRANCH
DIVISION OF TOXICOLOGY

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[Signature]
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The potential for DIGL-RP Solid Propellant to produce primary eye irritation was evaluated in male New Zealand White rabbits by using a modified Draize method. DIGL-RP produced slight conjunctival vasodilation (indicative of mild inflammation), and slight chemosis of the conjunctival membranes. However, these responses were not sufficiently severe to be classified as a positive response. Therefore, the results indicate that DIGL-RP Solid Propellant is not a primary eye irritant under conditions of this study.

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ABSTRACT

The potential for DIGL-RP Solid Propellant to produce primary eye irritation was evaluated in male New Zealand White rabbits by using a modified Draize method. DIGL-RP produced slight conjunctival vasodilation (indicative of mild inflammation), and slight chemosis of the conjunctival membranes. However, these responses were not sufficiently severe to be classified as a positive response. Therefore, the results indicate that DIGL-RP Solid Propellant is not a primary eye irritant under conditions of this study.

Key Words: DIGL-RP Solid Propellant, Ocular Irritation, Mammalian Toxicology, Rabbit, Diethylene glycol Dinitrate, Munition
TYPE REPORT: Primary Eye Irritation GLP Study Report

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Letterman Army Institute of Research
Presidio of San Francisco, CA 94129-6800

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US Army Biomedical Research and Development Laboratory
Fort Detrick, MD 21701-5010
Project Officer: Gunda Reddy, PhD

PROJECT/WORK UNIT/APC: 3E162720A835/180/TLB0

GLP STUDY NUMBER: 85023

STUDY DIRECTOR: LTC Don W. Korte, Jr., PhD, MSC
Diplomate, American Board of Toxicology

PRINCIPAL INVESTIGATOR: Gerald F.S. Hiatt, PhD

CO-PRINCIPAL INVESTIGATOR: SSG James D. Justus, MPA

PATHOLOGIST: MAJ G. Tracy Makovec, DVM, VC, Diplomate
American College of Veterinary Pathologists

REPORT AND DATA MANAGEMENT:
A copy of the final report, study protocol, retired SOPs, raw data,
analytical, stability, and purity data of the test compound, and an aliquot of the
test compound will be retained in the LAIR Archives.

TEST SUBSTANCE: DIGL-RP Solid Propellant

INCLUSIVE STUDY DATES: 14 Nov 1985 - 13 Dec 1985

OBJECTIVE:
The objective of this study was to determine the primary eye irritation
potential of DIGL-RP in male New Zealand White rabbits.
ACKNOWLEDGMENTS

MAJ Larry D. Brown, DVM, and SP4 Gayle A. Omer, BS, provided technical assistance. SP4 James J. Fisher, SP4 Scott L. Schwebe, SP4 Theresa L. Polk, Obie B. Goodrich, and Diane Arevalo provided care for the animals. Colleen Kamiyama provided administrative and clerical support during the performance of this study and preparation of the report.
SIGNATURES OF PRINCIPAL SCIENTISTS INVOLVED IN THE STUDY

We, the undersigned, declare that GLP Study 85023 was performed under our supervision, according to the procedures described herein, and that this report is an accurate record of the results obtained.

DON W. KORTE, JR., PhD / DATE
LTC, MSC
Study Director

GERALD F.S. HIATT, PhD / DATE
DAC
Principal Investigator

JAMES D. JUSTUS, MPA / DATE
SSG, USA
Co-Principal Investigator

CONRAD R. WHEELER, PhD / DATE
DAC
Analytical Chemist
MEMORANDUM FOR RECORD

SUBJECT: GLP Compliance for GLP Study 85023

1. This is to certify that the protocol for LAIR GLP Study 85023 was reviewed on 10 May 1985.

2. The institute report entitled "Primary Eye Irritation Potential of DIGL-RP Solid Propellant in Rabbits," Toxicology Series 165, was audited on 2 October 1989.

Carolyn M. Lewis
CAROLYN M. LEWIS
Diplomate, American Board of Toxicology
Quality Assurance Auditor
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INTRODUCTION

The Department of Defense is considering the use of diethylene glycol dinitrate (DEGDN), triethylene glycol dinitrate (TEGDN), or trimethyleneolane trinitrate (TMETN) as a replacement for nitroglycerin in new propellant formulations. However, considerable gaps in the toxicology data of the compounds were identified during a review of their health effects (1) conducted for the US Army Biomedical Research and Development Laboratory (USABRDL). Consequently, USABRDL has tasked the Division of Toxicology, Letterman Army Institute of Research (LAIR), to conduct an initial health effects evaluation of the proposed replacement nitrate esters. This initial evaluation of DEGDN, TMETN, TEGDN, and two DEGDN-based propellants, JA-2 and DIGL-RP, includes the Ames mutagenicity assay, acute oral toxicity tests in rats and mice, acute dermal toxicity in rabbits, dermal and ocular irritation studies in rabbits, and dermal sensitization studies in guinea pigs.

Objective of Study

The objective of this study was to determine the primary eye irritation potential of DIGL-RP in male New Zealand White rabbits.

MATERIALS

Test Substance

Chemical Name: DIGL-RP Solid Propellant

LAIR Code Number: TP57
Hiatt et al.–2

Physical State: Solid black cylinders (stick configuration)
Lot No.: RAD83M001S169
Other test substance information is presented in Appendix A.

Animal Data

Six male New Zealand White rabbits (Elkhorn Rabbitry, Watsonville, CA) were identified individually with ear tattoos numbered 85F309 to 85F314 inclusive. Animal weights on dosing day ranged from 2.6 to 2.9 kg. Additional animal data appear in Appendix B.

Husbandry

The rabbits were housed individually in stainless steel, screen-bottomed, battery-type cages with automatically flushing dump tanks. The diet consisted of approximately 150 g/day of Certified Purina Chow® Diet 5322 (Ralston Purina Company, Checkerboard Square, St. Louis, MO); water was provided by continuous drip from a central line. The animal room temperature was maintained at 16.2°C to 20.0°C and relative humidity ranged from 51% to 69%, except for occasional humidity spikes as high as 78% (room washing). The photoperiod was 12 hours of light per day.

METHODS

Conduct of this study was in accordance with the LAIR Standard Operating Procedure OP-STX-33, "Primary Eye Irritation Study," and guidelines promulgated by the EPA for ocular irritation testing (2,3).

Group Assignment/Acclimation

Study rabbits were assigned by numerical sequence to two dose groups of 3 males each. These animals were quarantined in the Division of Animal Care and Services for 14 days and acclimated for 7 days in the GLP Suite before dosing. During these periods they were observed daily for signs of illness.
Dose Levels and Administration

Approximately 83 mg (0.1 ml) of DIGL-RP was administered one time to one eye of each rabbit by gently pulling the lower lid away from the conjunctival cul-de-sac to form a cup into which the compound was instilled. Upper and lower lids were then held gently together for one second to prevent loss of material. Group 1 was dosed on 3 Dec 85 and Group 2 was dosed on 10 Dec 85.

Compound Preparation

DIGL-RP was received as pellets and was ground in a liquid nitrogen freezer mill (Spex Industries, Edison, NJ) to a fine gray powder which required no further preparation.

Test Procedures

On 2 Dec 85, both eyes of each Group 1 animal were examined for any preexisting abnormalities, by the procedure detailed below. For each animal, the eye with the nearest normal appearance was designated for treatment, the contralateral eye serving as an untreated control. On 3 Dec 85, approximately 83 mg of DIGL-RP was placed in the designated eye of each rabbit in this group. Group 2 rabbits underwent the same procedures on 9 and 10 Dec 85, respectively.

Ocular Examination/Grading

Initially each eye was observed unaided in a darkened room with focal illuminant (pen light). Structures examined included: the lids and surrounding fur, the conjunctiva (semilunar, palpebral, and bulbar), the cornea, and the iris. Grading of the cornea, iris, and conjunctiva was performed according to Table 1 (4). During the observations, each eye was also examined with a slit lamp. Special attention was given to integrity of the corneal surface, thickness of the corneal stroma, clarity of anterior chamber fluid, iridial morphology, clarity of the lens, and lenticular surface morphology (5). Additionally, any areas appearing grossly abnormal were examined.
TABLE 1: Grades for Ocular Lesions

<table>
<thead>
<tr>
<th>CORNEA</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Opacity: degree of density (area of greatest density taken for reading)</td>
<td></td>
</tr>
<tr>
<td>No ulceration or opacity</td>
<td>0</td>
</tr>
<tr>
<td>Scattered or diffuse areas of opacity (other than slight dulling of normal luster) details of iris clearly visible</td>
<td>1†</td>
</tr>
<tr>
<td>Easily discernible translucent areas, details of iris slightly obscured</td>
<td>2</td>
</tr>
<tr>
<td>Nacreous areas, no details of iris visible, size of pupil barely discernible</td>
<td>3</td>
</tr>
<tr>
<td>Opaque cornea, iris not discernible through opacity</td>
<td>4</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>IRIS</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal</td>
<td>0</td>
</tr>
<tr>
<td>Markedly deepened rugae, congestion, swelling, moderate circumjacent hyperemia or injection, any of these or any combination thereof, iris still reacting to light (sluggish reaction is positive)</td>
<td>1†</td>
</tr>
<tr>
<td>No reaction to light, hemorrhage, gross destruction (any or all of these)</td>
<td>2</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>CONJUNCTIVA</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Redness: (refers to palpebral and bulbar conjunctiva, excluding cornea and iris)</td>
<td></td>
</tr>
<tr>
<td>Blood vessels normal</td>
<td>0</td>
</tr>
<tr>
<td>Some blood vessels definitely hyperemic (injected)</td>
<td>1</td>
</tr>
<tr>
<td>Diffuse, crimson color, individual vessels not easily discernible</td>
<td>2†</td>
</tr>
<tr>
<td>Diffuse, beefy red</td>
<td>3</td>
</tr>
<tr>
<td>Chemoals: (lids and/or nictitating membranes)</td>
<td></td>
</tr>
<tr>
<td>No swelling</td>
<td>0</td>
</tr>
<tr>
<td>Any swelling above normal including nictitating membranes</td>
<td>1</td>
</tr>
<tr>
<td>Obvious swelling with partial eversion of lids</td>
<td>2†</td>
</tr>
<tr>
<td>Swelling with lids about half-closed</td>
<td>3</td>
</tr>
<tr>
<td>Swelling with lids more than half-closed</td>
<td>4</td>
</tr>
</tbody>
</table>

* Adapted from Table 6 in Draize et al. (4).
† Indicates minimum level for a positive response.
under high magnification. All observations, including normal appearance, were
detailed on the grading sheet. Following this, fluorescein dye (Fluor-I-Strips,
Ayerst Laboratories Inc., New York, NY) was introduced into the eye, which
was then observed under ultraviolet light. Any corneal areas reacting with the
dye (a sign of discontinuity of the corneal epithelium) were described with
respect to area and intensity of fluorescence. Examination and grading of
ocular reactions were performed in this fashion at 1, 4, 24, 48, and 72 hours
after dosing. Fluorescein staining was omitted from the 1- and 4-hour
observations. Due to an almost total lack of reaction during the 72 hours after
dosing, the study was terminated after this observation in accordance with the
protocol. All animals were submitted for necropsy. Therefore, no scoring or
observations were performed at 7, 14, or 21 days.

Duration of Study

Appendix C is a complete listing of historical events.

Changes/Deviations

Slit lamp examination was added to the standard observation
procedures. The slit lamp enables one to detect subtle reactions not grossly
observable and to evaluate more thoroughly those abnormalities which are
grossly observable. Color photographic documentation was not performed due
to lack of significant response to the test compound. Animal 85F312 was
removed from the study after sustaining a broken back on 9 Dec 85. With
these exceptions, this study was completed in accordance with the
appropriate protocol and addenda. It is believed that none of these
changes/deviations affected the performance of the study or the validity of
the results.

Storage of Raw Data and Final Report

A copy of the final report, study protocols, raw data, retired SOPs and
an aliquot of the test compound will be retained in the LAIR Archives.
RESULTS

Tabulation of the Draize-type ocular grading results is presented in Appendix D and a summary of the ocular observations in Appendix E.

Comea

DIGL-RP produced no grossly observable effects in the cornea. All treated eyes were assigned zero scores for both opacity and area involvement at all observations after dosing.

Slit lamp examination revealed no corneal reactions attributable to the test compound. Slit lamp observations revealed corneas of normal thickness, indicating lack of edema, and smooth surfaces, indicating epithelial integrity. No staining of corneal epithelium was observed at any of the fluorescein examinations in the treated eyes.

Iris/Anterior Chamber

No grossly observable reactions were produced in the iris by DIGL-RP. Iridial scores were consistently zero at all observation times.

No iridial abnormalities were detected by slit lamp examination of the treated eyes. Circumiridial vessels and surface morphology were normal at all times after dosing. Close examination of anterior chamber fluid revealed no evidence of the presence of protein or cells (signs of iridial inflammation).

Lens

The lens was not scored under the Draize-type grading system because of the difficulty in making unaided observations. At all times after dosing, the lens appeared normal during slit lamp examination. No changes were observed in clarity or surface morphology.

Conjunctiva

In this study, DIGL-RP produced only two grossly observable responses: slight conjunctival redness and chemosis. At 1 and 4 hours after dosing, all
of the treated eyes exhibited slight vasodilatation in the bulbar (sclera) or semilunar (nictitating membrane) conjunctiva. Conjunctival redness scores of 1 were assigned to 5 of 5 treated eyes and slit lamp examination confirmed the presence of dilated vessels within the outer layers of the sclera and the nictitating membrane. Chemosis scores of 1 were assigned to 4 of 5 treated eyes. Animal 85F309 continued to have conjunctival redness at 24 hours. Chemosis was not present in any treated eyes at 24 hours.

**Control Eyes**

At no time during the study did the contralateral untreated eyes exhibit any change from their normal condition on the day of dosing.

**Pathology Report**

Lesions observed were considered incidental and in no way related to the treatment. The pathologist's report is presented in Appendix F.

**DISCUSSION**

The primary goal of ocular toxicity testing is to determine the potential for ocular damage resulting from accidental contact of the test compound with the eye. For this purpose the Draize-type irritation test, used in the present study, is especially well-suited. An important feature of this test is that the route and type of exposure (ocular instillation followed by a forced blink) closely mimics potential human exposures.

Consumer Product Safety Commission guidelines, which the EPA recommends for ocular irritation testing, state that an animal has exhibited a positive reaction if the test substance produces one or more of the following signs: ulceration of the cornea (other than a fine stippling); opacity of the cornea (other than a slight dulling of the normal luster); inflammation of the iris (other than a slight deepening of the rugae or a slight hyperemia of the circumcorneal blood vessels); an obvious swelling in the conjunctiva with partial eversion of the lids; or a diffuse crimson-red coloration in the conjunctiva with individual vessels not easily discernible (2).
Guidelines for classification of chemicals as ocular irritants or nonirritants have been published and form the basis for evaluation in the present study (6). These Interagency Regulatory Liaison Group (IRLG) guidelines state: "[a] test result is considered positive if four or more animals exhibit a positive reaction. If only one animal exhibits a positive reaction, the test result is regarded as negative."

In this study, DIGL-RP produced no positive reactions, as defined by the IRLG. Slight conjunctival redness and slight chemosis were the only responses observed. These reactions, although scorable, did not achieve sufficient severity to warrant consideration as a "positive response." Due to this lack of positive response, DIGL-RP is classified as a nonirritant by the results of the present study.

CONCLUSION

DIGL-RP exhibited minimal potential to produce ocular irritation under conditions of this study.
REFERENCES


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Appendix A: CHEMICAL DATA

Chemical Name: DIGL-RP Solid Propellant

LAIR Code Number: TP57

Physical State: Solid black cylinders (stick configuration)

Preparation of test substance for dosing: The cylinders of DIGL-RP were ground under liquid nitrogen using a Spex freezer mill. After grinding, the powder was sieved through an 80-mesh screen.

Chemical analysis:

DEGDN was the only major component of DIGL which could be easily analyzed. For analysis, samples of DIGL powder were added to individual 100 ml volumetric flasks. After dilution to volume with 90% ethanol, a second 1:100 dilution was performed. These solutions were analyzed by HPLC. Standards consisted of solutions of DEGDN in ethanol, ranging in concentration from 164.5 to 670.5 μg/ml. Analysis of DEGDN by HPLC was performed under the following conditions: column, Brownlee RP-18 (4.6 x 250 mm, Brownlee Labs, Inc., Santa Clara, CA); solvent system, 40% water - 60% acetonitrile; flow rate, 0.9 ml/min; wavelength monitored, 210 nm. Under these conditions, DEGDN eluted with a retention time of approximately 5.4 min. The results from the analysis of standards and DIGL powder samples are presented in Tables 1 and 2.

Table 1. Analysis of Standards

<table>
<thead>
<tr>
<th>Concentration of Standard (μg/ml)</th>
<th>Peak Area* (x 10^-7)</th>
</tr>
</thead>
<tbody>
<tr>
<td>164.5</td>
<td>0.94</td>
</tr>
<tr>
<td>191.0</td>
<td>1.09</td>
</tr>
<tr>
<td>275.5</td>
<td>1.60</td>
</tr>
<tr>
<td>299.4</td>
<td>1.74</td>
</tr>
<tr>
<td>362.0</td>
<td>2.08</td>
</tr>
<tr>
<td>399.6</td>
<td>2.31</td>
</tr>
<tr>
<td>444.4</td>
<td>2.52</td>
</tr>
<tr>
<td>539.8</td>
<td>3.07</td>
</tr>
<tr>
<td>585.0</td>
<td>3.32</td>
</tr>
<tr>
<td>670.5</td>
<td>3.79</td>
</tr>
</tbody>
</table>

*Average of 2 determinations

Equation for line by linear regression analysis:

\[ Y = 5.62 \times 10^4 X + 3.51 \times 10^5, r^2 = 0.9999 \]
Appendix A (cont.): CHEMICAL DATA

Table 2. Analysis of DIGL Powder

<table>
<thead>
<tr>
<th>Weight of DIGL Analyzed (mg)</th>
<th>Dilution Factor</th>
<th>Peak Area (x 10^-7)</th>
<th>Conc. of DEGDN in DIGL (weight %)</th>
</tr>
</thead>
<tbody>
<tr>
<td>111.7</td>
<td>100</td>
<td>2.45</td>
<td>38.5</td>
</tr>
<tr>
<td>112.6</td>
<td>100</td>
<td>2.46</td>
<td>38.3</td>
</tr>
<tr>
<td>100.1</td>
<td>100</td>
<td>2.21</td>
<td>38.7</td>
</tr>
</tbody>
</table>

*Calculated using the equation for the standard curve as follows:

\[
\text{Conc. of DEGDN} = \frac{\text{Peak Area} - 3.51 \times 10^5}{5.62 \times 10^4} + \frac{\text{wgt DIGL (mg)}}{10}.\]

The average value for the concentration of DEGDN in DIGL was 38.5% and this agrees closely with the value of 36.70 ± 1.50 reported in the manufacturer’s data sheet.

Source: Radford Army Ammunition Plant, Radford, Virginia
(Prime contractor: Hercules, Inc., Wilmington, Delaware)

Lot No.: RAD83M001S169

1 Wheeler CW. Toxicity Testing of Propellents. Laboratory Notebook #85-12-023, p. 51-61. Letterman Army Institute of Research, Presidio of San Francisco, CA.

Appendix A (cont.): CHEMICAL DATA

CHEMICAL ANALYSIS FOR DIGL-RP
(Information from the Manufacturer's Data Sheet)

<table>
<thead>
<tr>
<th>Ingredients</th>
<th>Finished Propellant Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nitrocellulose</td>
<td></td>
</tr>
<tr>
<td>(13.05 ±0.05% Nitrogen)</td>
<td>62.5 ±2.00</td>
</tr>
<tr>
<td>(6-12 seconds viscosity)</td>
<td></td>
</tr>
<tr>
<td>Diethylene glycol Dinitrate (DEGDN)</td>
<td>36.70 ±1.50</td>
</tr>
<tr>
<td>Ethyl Centralite (EC)</td>
<td>0.25 ±0.05</td>
</tr>
<tr>
<td>Akardit II</td>
<td>0.25</td>
</tr>
<tr>
<td>Magnesium Oxide</td>
<td>0.45 ±0.15</td>
</tr>
<tr>
<td>Graphite (Chg 5)</td>
<td>0.05 Max</td>
</tr>
<tr>
<td><strong>TOTAL</strong></td>
<td><strong>100.0</strong></td>
</tr>
</tbody>
</table>
Appendix B: ANIMAL DATA

Species: *Oryctolagus cuniculus*

Strain: New Zealand White (albino)

Source: Elkhorn Rabbitry
5265 Starr Way
Watsonville, CA 95076

Sex: Male

Age: Young adults

Animals in each group: 3 males

Condition of animals at start of study: Normal

Body weight range at dosing: 2.6 - 2.9 kg

Identification procedures:

Ear tattoo: numbers 85F309, 85F310, 85F311, 85F312, 85F313, 85F314.

Pretest conditioning:

1. Quarantine from 14 Nov 85 - 27 Nov 85.
2. Acclimation from 28 Nov 85 - 3 Dec 85.
3. Animal eyes were examined 24 hours before dosing using slit lamp, fluorescein dye and ultraviolet light.

Justification:

Laboratory rabbits are a proven sensitive animal model for ocular testing.
### Appendix C: HISTORICAL LISTING OF STUDY EVENTS

<table>
<thead>
<tr>
<th>Date</th>
<th>Event</th>
</tr>
</thead>
<tbody>
<tr>
<td>14 Nov 85</td>
<td>Animals arrived at LAIR. They were examined for illness and placed under a two-week quarantine.</td>
</tr>
<tr>
<td>14 - 27 Nov 85</td>
<td>Animals were checked daily by Division of Animal Care and Services personnel.</td>
</tr>
<tr>
<td>15,27 Nov, 3,13 Dec 85</td>
<td>Animals were weighed.</td>
</tr>
<tr>
<td>18 Nov 85</td>
<td>They were tattooed and given one application of Canex®/mineral oil into ears for earmite prevention.</td>
</tr>
<tr>
<td>27 Nov 85</td>
<td>Rabbits were certified healthy by a staff veterinarian removed from quarantine, and assigned to test groups.</td>
</tr>
<tr>
<td>2 Dec 85</td>
<td>Animals were checked for preexisting ocular injury (Group 1).</td>
</tr>
<tr>
<td>3 Dec 85</td>
<td>Group 1 rabbits were dosed. Eyes were scored 1 and 4 hours after exposure.</td>
</tr>
<tr>
<td>4 Dec 85</td>
<td>Eyes were scored 24 hours after exposure (Group 1).</td>
</tr>
<tr>
<td>5 Dec 85</td>
<td>Eyes were scored 48 hours after exposure (Group 1).</td>
</tr>
<tr>
<td>6 Dec 85</td>
<td>Eyes were scored 72 hours after exposure. Study was terminated (Group 1).</td>
</tr>
<tr>
<td>9 Dec 85</td>
<td>Animals were checked for preexisting ocular injury (Group 2). Group 1 animals were weighed and submitted for necropsy.</td>
</tr>
<tr>
<td>10 Dec 85</td>
<td>Group 2 rabbits were dosed and weighed. Eyes were scored 1 and 4 hours after exposure.</td>
</tr>
<tr>
<td>11 Dec 85</td>
<td>Eyes were scored 24 hours after exposure (Group 2).</td>
</tr>
<tr>
<td>12 Dec 85</td>
<td>Eyes were scored 48 hours after exposure (Group 2).</td>
</tr>
<tr>
<td>13 Dec 85</td>
<td>Eyes were scored 72 hours after exposure. Study was terminated and animals were submitted for necropsy (Group 2).</td>
</tr>
</tbody>
</table>
### Appendix D: TABULATED OCULAR DATA

#### CORNEAL OPACITY
(score by animal)

<table>
<thead>
<tr>
<th>Rabbit Number</th>
<th>Baseline</th>
<th>1 hr</th>
<th>4 hr</th>
<th>24 hr</th>
<th>48 hr</th>
<th>72 hr</th>
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</thead>
<tbody>
<tr>
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</tbody>
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#### IRIS
(score by animal)

<table>
<thead>
<tr>
<th>Rabbit Number</th>
<th>Baseline</th>
<th>1 hr</th>
<th>4 hr</th>
<th>24 hr</th>
<th>48 hr</th>
<th>72 hr</th>
</tr>
</thead>
<tbody>
<tr>
<td>85F309</td>
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<td>0</td>
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<tr>
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<tr>
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### Appendix D (cont.): TABULATED OCULAR DATA

#### CONJUNCTIVA (CHEMOSIS)
(score by animal)

<table>
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<th>Rabbit Number</th>
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<th>4 hr</th>
<th>24 hr</th>
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<th>72 hr</th>
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</thead>
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<td>0</td>
</tr>
<tr>
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<td>0</td>
</tr>
<tr>
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<td>0</td>
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</table>

#### CONJUNCTIVA (REDNESS)
(score by animal)

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<th>Baseline</th>
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<th>4 hr</th>
<th>24 hr</th>
<th>48 hr</th>
<th>72 hr</th>
</tr>
</thead>
<tbody>
<tr>
<td>85F309</td>
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<td>0</td>
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<tr>
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</tbody>
</table>
Appendix E: SUMMARY OF OCULAR OBSERVATIONS

One Hour After Dosing

Slight hyperemia was present in all 5 test rabbits. This hyperemia was confined to the lower bulbar and palpebral conjunctiva and the nictitating membrane. Chemosis was present in 4 of the 5 test rabbits in the lower conjunctiva and the nictitating membrane. Both the vasodilatation and swelling were visible with the unaided eye. All other structures appeared normal.

Four Hours After Dosing

Slight hyperemia was present in the conjunctiva of all 5 rabbits. Chemosis was present in 4 of the 5 rabbits. All other structures appeared normal.

Twenty-four Hours After Dosing

Slight hyperemia persisted in 1 rabbit. The chemosis was no longer present. All other structures appeared normal.

Forty-eight Hours After Dosing

All structures examined by slit lamp appeared normal and no fluorescein staining was present.

Seventy-two Hours After Dosing

All structures examined by slit lamp appeared normal and no fluorescein staining was present.
Appendix F: PATHOLOGY REPORT

LAIR Gross Pathology Report
GLP Study 85023

Test: Eye Irritation

Investigator: Dr. Hiatt, Toxicology Branch

Test Substance: DIGL-RP


Gross findings:

<table>
<thead>
<tr>
<th>ANIMAL ID</th>
<th>LAIR accession</th>
<th>FINDINGS</th>
</tr>
</thead>
<tbody>
<tr>
<td>85F309</td>
<td>38672</td>
<td>Not remarkable (NR)</td>
</tr>
<tr>
<td>85F310</td>
<td>38673</td>
<td>Pinworm - cecum</td>
</tr>
<tr>
<td>85F311</td>
<td>38674</td>
<td>NR</td>
</tr>
<tr>
<td>85F313</td>
<td>38679</td>
<td>Pinworm - cecum</td>
</tr>
<tr>
<td>85F314</td>
<td>38680</td>
<td>Pinworm - cecum</td>
</tr>
</tbody>
</table>

Comment: The lesions noted were incidental and not related to the treatment.

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Comparative Pathology Branch

G. Tracy Manovec, DM
CPT, VC
Diplomate, ACVP
Comparative Pathology Branch

23 December 1985
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