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BIOASSAY OF DIOXATHION

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Carcinogenesis Testing Program Division of Cancer Cause and Prevention National Cancer Institute National Institutes of Health Bethesda, Maryland 20014

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REPORT ON THE BIOASSAY OF DIOXATHION FOR POSSIBLE CARCINOGENICITY

CARCINOGENESIS TESTING PROGRAM DIVISION OF CANCER CAUSE AND PREVENTION NATIONAL CANCER INSTITUTE, NATIONAL INSTITUTES OF HEALTH

FOREWORD: This report presents the results of the bioassay of dioxathion conducted for the Carcinogenesis Testing Program, Division of Cancer Cause and Prevention, National Cancer Institute (NCI), National Institutes of Health, Bethesda, Maryland. This is one of a series of experiments designed to determine whether selected chemicals have the capacity to produce cancer in animals. Negative results, in which the test animals do not have a significantly greater incidence of cancer than control animals, do not necessarily mean the test chemical is not a carcinogen because the experiments are conducted under a limited set of circumstances. Positive results demonstrate that the test chemical is carcinogenic for animals under the conditions of the test 'and indicate a potential risk to man. The actual determination of the risk to man from animal carcinogens requires a wider analysis.

<u>CONTRIBUTORS</u>: This bioassay of dioxathion was conducted by Hazleton Laboratories America, Inc., Vienna, Virginia, initially under direct contract to the NCI and currently under a subcontract to Tracor Jitco, Inc., prime contractor for the NCI Carcinogenesis Testing Program.

The experimental design was determined by the NCI Project Officers, Dr. J. H. Weisburger (1,2) and Dr. E. K. Weisburger (1). The principal investigators for the contract were Dr. M. B. Powers (3), Dr. R. W. Voelker (3), Dr. W. A. Olson (3,4) and Dr. W. M. Weatherholtz (3). Chemical analysis was performed by Dr. C. L. Guyton (3,5) and the analytical results were reviewed by Dr. N. Zimmerman (6); the technical supervisor of animal treatment and observation was Ms. K. J. Petrovics (3).

Histopathologic examinations were performed by Dr. R. H. Habermann (3) and reviewed by Dr. R. W. Voelker (3) at the Hazleton Laboratories America, Inc., and the diagnoses included in this report represent the interpretation of these pathologists. Histopathology findings and reports were reviewed by Dr. R. L. Schueler (7).

Compilation of individual animal survival, pathology, and summary tables was performed by EG&G Mason Research Institute (8); the statistical analysis was performed by Mr. W. W. Belew (6) and Dr. J. R. Joiner (7), using methods selected for the Carcinogenesis Testing Program by Dr. J. J. Gart (9). This report was prepared at METREK, a Division of The MITRE Corporation (6) under the direction of the NCL. Those responsible for this report at METREK are the project coordinator, Dr. L. W. Thomas (6), task leader Dr. M. R. Kornreich (6), senior biologist Ms. P. Walker (6), biochemist Mr. S. C. Drill (6), and technical editor Ms. P. A. Miller (6). The final report was reviewed by members of the participating organizations.

The following other scientists at the National Cancer Institute were responsible for evaluating the bioassay experiment, interpreting the results, and reporting the findings: Dr. K. C. Chu (1), Dr. C. Cueto, Jr. (1), Dr. J. F. Douglas (1), Dr. D. G. Goodman (1), Dr. R. A. Griesemer (1), Dr. H. A. Milman (1), Dr. T. W. Orme (1), Dr. R. A. Squire (1,10), Dr. J. M. Ward (1), and Dr. C. E. Whitmire (1).

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SUMMARY

A bioassay for possible carcinogenicity of technical-grade dioxathion was conducted using Osborne-Mendel rats and B6C3F1 mice. Dioxathion was administered in the feed, at either of two concentrations, to groups of 50 male and 50 female animals of each species. The high and low time-weighted average concentrations were, respectively, 180 and 90 ppm for male rats and 90 and 45 ppm for female rats. The high and low time-weighted average concentrations for male mice were 567 and 284 ppm, respectively, and for female mice were 935 and 467 ppm, respectively. After a 78-week period of chemical administration, observation of the rats continued for an additional 33 weeks and the mice were observed for an additional 12 to 13 weeks. For rats, 50 animals of each sex were placed on test as controls and fed only the basal diet, while for mice 20 animals of each sex served as controls.

In both species adequate numbers of animals survived long enough to be at risk from late-appearing tumors.

A variety of neoplasms was observed in treated animals of both species; however, none of the neoplasms observed were either histopathologically unusual or in statistically significant incidences.

Under the conditions of this bioassay, dietary administration of dioxathion was not carcinogenic in Osborne-Mendel rats or B6C3F1 mice. P

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I. INTRODUCTION

Dioxathion (NCI No. COO395), an organophosphorous insecticide, was selected for bioassay by the National Cancer Institute because of its widespread use on livestock and edible crops, and a lack of adequate chronic toxicity data.

The Chemical Abstracts Service (CAS) Ninth Collective Index (1977) name for this compound is phosphorodithioic acid, S,S'-pdioxane-2,3-diyl 0,0,0',0'-tetraethyl ester. *t* It is also called 2,3-p-dioxanedithiol S,S-bis(0,0-diethyl phosphorodithioate); phosphorodithioic acid, S,S'-1,4-dioxane-2,3-diyl 0,0,0',0'-tetraethyl ester; and p-dioxane-2,3-dithiol, S,S-diester with 0,0-diethyl phosphorodithioate.

Dioxathion is used as a spray to control mites on cotton, grapes, citrus, ornamentals, and certain other fruits including apples, pears, and quinces; and as a spray or dip on cattle, goats, sheep, and hogs for control of ticks, lice, horn flies, and sheep ked (<u>Farm Chemicals</u> Handbook, 1976).

Specific production figures for dioxathion are not available; however, one U.S. company reported commercial production (in excess of 1000 pounds or \$1000 in value annually) in 1975 (U.S. International Trade Commission, 1977).

The CAS registry number is 78-34-2.

The potential for exposure to dioxathion is greatest for agricultural workers but may also be considerable for workers in plants which produce the compound. The general public may be exposed to dioxathion in house and garden pesticides for evergreens and shrubs (Gosselin et al., 1976) and to airborne dioxathion in agricultural regions following spraying operations. Dioxathion is nonvolatile and relatively stable (<u>Farm Chemicals Handbook</u>, 1976); consequently, exposure via ingestion may occur as a result of persistence and possible accumulation of residues on treated crops and livestock.

Organophosphorous insecticides, which are chemically related to the nerve gases, are among the most toxic pesticides in current use (Gosselin et al., 1976; Hall, 1950). These compounds act as powerful cholinesterase inhibitors throughout the body, and can be absorbed to a dangerous degree through all routes, including the intact skin. Inhalation is considered to be the most dangerous route of exposure, followed by ingestion (Gosselin et al., 1976); oral intake of dioxathion in quantities greater than 0.075 mg/kg produces measurable cholinesterase inhibition in humans (Spencer, 1973).

The initial symptoms of organophosphorous insecticide poisoning vary with the site of absorption: nausea, vomiting, diarrhea, and sialorrhea after ingestion; rhinorrhea and a feeling of tightness of the chest following inhalation; miosis, blurring or dimness of vision, tearing, and ciliary muscle spasms after ocular exposure; and local sweating and twitching following dermal contact (Gosselin et al.,

1976). Progressive loss of muscular control, due to the build-up of acetylcholine at the neuromuscular junctions, occurs regardless of the route of exposure and results in slurring of speech, difficulty in breathing, fasciculations and twitching, and an overall loss of coordination. In severe cases, convulsions, incontinence, random jerky movements, and coma may ensue. When de_th occurs, it is usually due to respiratory failure.

II. MATERIALS AND METHODS

A. Chemicals

Commercial technical-grade dioxathion (Figure 1) (Delnav[®]) was purchased from Hercules, Incorporated as a single lot and analyzed for purity by Hazleton Laboratories America, Inc., Vienna, Virginia. Dioxathion, a liquid existing in the <u>cis</u> and <u>trans</u> geometric forms, is sensitive to both heat and alkali.

The first analysis, one month prior to initiation of the chronic bioassay, was by gas-liquid chromotography (GLC). It was felt that, although dioxathion undergoes thermal decomposition in the chromatograph inlet to produce numerous compounds, a GLC profile would be useful for comparisons from one year to the next. In this way, changes in the chromatogram might indicate any alterations in the composition of the chemical due to storage.

The second analysis, performed approximately one year later, was by both GLC and a cleavage-hydrolysis gravimetric method suggested by the manufacturer. The chromatogram from this second GLC analysis was in general agreement with that of the first year. A mean dioxathion content of approximately 69 percent was indicated using the gravimetric method and this was in accord with the manufacturer's stated analysis of a mean dioxathion content ranging from 68 to 75 percent. It was noted by the manufacturer that, although the gravimetric method does not differentiate the <u>cis</u> and <u>trans</u> isomers, the <u>trans</u> isomer predominates.



FIGURE 1 CHEMICAL STRUCTURE OF DIOXATHION

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The third and last analysis, performed approximately two years after the first one, was by the gravimetric method alone. The mean dioxathion content was indicated to be approximately 56 percent, considerably lower than the mean detected the previous year. It was, therefore, concluded that chemical degradation had occurred.

Throughout this report the term dioxathion is used to represent this commercial technical-grade material.

B. Dietary Preparation

The basal laboratory diet for both treated and control animals consisted of Wayne Lab-Blox[®] (Allied Mills, Inc., Chicago, Illinois) plus 2 percent Duke's[®] corn oil (S. F. Sauer Company, Richmond, Virginia) by weight. Fresh mixtures of dioxathion in corn oil were prepared each week and stored in the dark. The dioxathion mixtures were incorporated into the appropriate amount of laboratory diet in a twin-shell blender fitted with an accelerator bar.

C. <u>Animals</u>

Two animal species, rats and mice, were used in the carcinogenicity bioassay. The Osborne-Mendel rat was selected on the basis of a comparative study of the tumorigenic responsiveness to carbon tetrachloride of five different strains of rats (Reuber and Glover, 1970). The B6C3F1 mouse was selected because it has been used by the NCI for carcinogenesis bioassays and has proved satisfactory in this capacity.

Rats and mice of both sexes were obtained through contracts of the Division of Cancer Treatment, National Cancer Institute. The

Osborne-Mendel rats were procured from the Battelle Memorial Institute, Columbus, Ohio, and the B6C3F1 mice were obtained from the Charles River Breeding Laboratories, Inc., Wilmington, Massachusetts. Upon receipt, animals were quarantined for at least 10 days, observed for visible signs of disease or parasites, and assigned to the various treated and control groups.

D. Animal Maintenance

All animals were housed by species in temperature- and humiditycontrolled rooms. The temperature range was 20° to 24°C and the relative humidity was maintained between 45 and 55 percent. The air conditioning system in the laboratory provided filtered air at a rate of 10 to 15 complete changes of room air per hour. Fluorescent lighting was provided on a 12-hour-daily cycle.

The rats were individually housed in suspended galvanized-steel wire-mesh cages with perforated floors, while mice were housed by sex in groups of ten in solid-bottom polycarbonate cages equipped with filter tops. Sanitized cages with fresh bedding (Sanichips, Pinewood Sawdust Company, Moonachie, New Jersey) were provided once each week for mice. Rats received sanitized cages with no bedding with the same frequency. Food hoppers were changed and heat-sterilized once a week for the first 10 weeks and once a month thereafter, while fresh heat-sterilized glass water bottles and sipper tubes were provided three times a week. Food and water were available <u>ad libitum</u>.

Treated and control rats were housed in the same room with other rats receiving diets containing^{*} trifluralin (1582-09-8); endosulfan (115-29-7); dicofol (115-32-2); nitrofen (1836-75-5); and mexacarbate (315-18-4).

All mice used in the dioxathion study were housed in the same room as other mice receiving diets containing trifluralin (1582-09-8); chlorobenzilate (510-15-6); sulfallate (95-06-7); p,p'-DDT (50-29-3); methoxychlor (72-43-5); p,p'-DDE (72-55-9); p,p'-TDE (72-54-8); dicofol (115-32-2); pentachloronitrobenzene (82-68-8); clonitralid (1420-04-8); nitrofen (1836-75-5); endosulfan (115-29-7); mexacarbate (315-18-4); amitrole (61-82-5); safrole (94-59-7); and acetylaminofluorene (53-96-3).

E. Selection of Initial Concentrations

In order to establish the maximum tolerated concentrations of dioxathion for administration to treated animals in the chronic studies, subchronic toxicity studies were conducted with both rats and mice. Animals of each species were distributed among six groups, each consisting of five males and five females. Dioxathion was premixed with a small amount of corn oil. This mixture was then incorporated into the laboratory diet and fed <u>ad libitum</u> to five of the six rat groups in concentrations of 21, 46, 100, 215, and 464 ppm, and to five of the six mouse groups in concentrations of 251, 398, 631, 1000, and 1590 ppm. The sixth group of each species served

CAS registry numbers are given in parentheses.

as a control group, receiving only the mixture of corn oil and laboratory chow. The dosed dietary preparations were administered for a period of 6 weeks, followed by a 2-week observation period during which all animals were given the basal diet.

A concentration inducing no mortality and resulting in a depression in mean group body weight of approximately 20 percent relative to controls was selected as the initial high concentration for the chronic study. When weight gain criteria were not applicable, mortality data alone were utilized.

All male rats treated with 215 ppm or less survived the entire 8-week period. Mean body weight depression was 10 percent in the males receiving 100 ppm and 27 percent in the males receiving 215 ppm. In the female rat groups, one of the five animals treated with 21 ppm died, no females died at 46 ppm, and two of the five treated with 100 ppm died during the study. Significant mean body weight depression was not observed in the females receiving 100 ppm or less. The initial high concentrations selected for the chronic bioassay were 150 ppm for the male rats and 75 ppm for the female rats.

In the mice one male receiving 398 ppm, four males receiving 1000 ppm, three males receiving 1590 ppm, and one female treated with 1000 ppm died during the study. The initial high concentrations selected for the chronic bioassay were 450 ppm for male mice and 700 ppm for the female mice.

F. Experimental Design

The experimental design parameters for the chronic study (species, sex, group size, actual concentrations administered, duration of treated and untreated observation periods, and the time-weighted average concentrations) are summarized in Tables 1 and 2.

Rats were all approximately 6 weeks old at the time they were placed on test. The concentrations of dioxathion initially utilized for male rats were 150 and 75 ppm. Throughout this report those male rats initially receiving the former concentration are referred to as the high dose group and those initially receiving the latter concentration are referred to as the low dose group. The initial concentrations utilized for the females were 75 and 37 ppm. Throughout this report those female rats initially receiving the former concentration are referred to as the high dose group while those initially receiving the latter concentration are referred to as the low dose group. In week 32 of the study high and low dioxathion levels were increased, respectively, to 200 and 100 ppm for the males, and to 100 and 50 ppm for the females. These concentrations were maintained until the end of the period of compound administration. Final observations on all rats were made 111 weeks after the experiment began.

Mice were all approximately 6 weeks old at the time they were placed on test. The concentrations initially administered to the male mice were 450 and 225 ppm. Throughout this report those male mice initially receiving the former concentration are referred to

TABLE 1

DESIGN SUMMARY FOR OSBORNE-MENDEL RATS DIOXATHION FEEDING EXPERIMENT

	INITIAL GROUP SIZE	DIOXATHION CONCENTRATION ^a	TREATED	ION PERIOD UNTREATED (WEEKS)	TIME-WEIGHTED AVERAGE CONCENTRATION ^D
MALE					
CONTROL	50	0	Ø	111	0
LOW DOSE	50	75 100 0	31 47	33	90
HIGH DOSE	50	150 200 0	31 47	33	180
FEMALE					
CONTROL	50	0	Q	111	C
LOW DOSE	50	37 50 0	31 47	33	45
HIGH DOSE	50	75 100 0	31 47	33	90

a Concentrations given in parts per million.

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^b Time-weighted average concentration = $\frac{\Sigma(\text{concentration X weeks received})}{\Sigma(\text{weeks receiving chemical})}$

TABLE 2

DESIGN SUMMARY FOR B6C3F1 MICE DIOXATHION FEEDING EXPERIMENT

	INITIAL GROUP SIZE	DIOXATHION CONCENTRATION ^a	OBSERVAT TREATED (WEEKS)	ION PERIOD UNTREATED (WEEKS)	TIME-WEIGHTED AVERAGE <u>CONCENTRATION</u> ^b
MALE					
CONTROL	20	0	Q	90	Ø
LOW DOSE	50	225 300 Q	17 61	12	284
HIGH DOSE	50	450 600 0	17 61	13	567
FEMALE				,,, , , , , , , , , , , , , , , , , ,	
CONTROL	20	Q	٥	90	0
LOW DOSE	50	350 500 0	17 61	13	467
HIGH DOSE	50	700 1000 0	17 61	13	935

^aConcentrations given in parts per million.

^b Time-weighted average concentration = $\frac{\Sigma(\text{concentration X weeks received})}{\Sigma(\text{weeks receiving chemical})}$ as the high dose group while those initially receiving the latter concentration are referred to as the low dose group. Female mice initially received concentrations of 700 and 350 ppm. Throughout this report those female mice initially receiving the former concentration are referred to as the high dose group while those initially receiving the latter concentration are referred to as the low dose group. In week 18 of the study, the high and low dosages were increased to 600 and 300 ppm, respectively for the male mice, and to 1000 and 500 ppm, respectively, for the female mice. These concentrations were maintained for the remainder of the dosing period. Final observations on all mice were made 90 to 91 weeks after the experiment began.

G. Clinical and Hispathologic Examinations

Animals were weighed immediately prior to initiation of the experiment. Body weights, food consumption, and data concerning appearance, behavior, signs of toxic effects, and incidence, size, and location of tissue masses were recorded at weekly intervals for the first 10 weeks and at monthly intervals thereafter. From the first day, all animals were inspected daily for mortality. The presence of tissue masses was determined by observation and palpation of each animal.

During the course of this bioassay several pathology protocols were in effect, each for different periods of time. The minimum protocol required that, if possible, certain tissues were to be taken

and examined histopathologically from all control animals, from any animal in which a tumor was observed during gross examination, and from at least 10 grossly normal males and 10 grossly normal females from each treated group. In addition, any tissues showing gross abnormalities were to be taken and examined histopathologically. Under later protocols, some tissues were taken from additional dosed animals. The number of animals in each group from which a tissue was examined is indicated in Appendices A through D.

A necropsy was performed on each animal regardless of whether it died, was killed when moribund, or was sacrificed at the end of the bioassay. The animals were euthanized by exsanguination under sodium pentobarbital anesthesia, and were immediately necropsied. The histopathologic examination consisted of gross and microscopic examination of major tissues, organs, and gross lesions taken from sacrificed animals and, whenever possible, from animals found dead.

Tissues were preserved in 10 percent buffered formalin, embedded in paraffin, sectioned, and stained with hematoxylin and eosin prior to microscopic examination. An occasional section was subjected to special staining techniques for more definitive diagnosis.

Slides were prepared from the following tissues from selected animals: skin, subcutaneous tissue, muscle, lungs and bronchi, trachea, bone marrow, spleen, lymph nodes, thymus, heart, salivary gland, liver, gallbladder (mice), pancreas, esophagus, stomach, small intestine, large intestine, kidney, urinary bladder, pituitary,

adrenal, thyroid, parathyroid, testis, prostate, seminal vesicle, brain, uterus, mammary gland, and ovary.

A few tissues were not examined for some animals, particularly for those that died early. Also, some animals were missing, cannibalized, or judged to be in such an advanced state of autolysis as to preclude histopathologic interpretation. Thus, the number of animals for which particular organs, tissues, or lesions were examined microscopically varies and does not necessarily represent the number of animals that were placed on experiment in each group.

H. Data Recording and Statistical Analyses

Pertinent data on this experiment have been recorded in an automatic data processing system, the Carcinogenesis Bioassay Data System (Linhart et al., 1974). The data elements include descriptive information on the chemicals, animals, experimental design, clinical observations, survival, body weight, and individual pathologic results, as recommended by the International Union Against Cancer (Berenblum, 1969). Data tables were generated for verification of data transcription and for statistical review.

These data were analyzed using the statistical techniques described in this section. Those analyses of the experimental results that bear on the possibility of carcinogenicity are discussed in the statistical narrative sections.

Probabilities of survival were estimated by the product-limit procedure of Kaplan and Meier (1958) and are presented in this report

in the form of graphs. Animals were statistically censored as of the time that they died of other than natural causes or were found to be missing; animals dying from natural causes were not statistically censored. Statistical analyses for a possible dose-related effect on survival used the method of Cox (1972) when testing two groups for equality and used Tarone's (1975) extensions of Cox's methods when testing a dose-related trend. One-tailed P-values have been reported for all tests except the departure from linearity test, which is only reported when its two-tailed P-value is less than 0.05.

The incidence of neoplastic or nonneoplastic lesions has been given as the ratio of the number of animals bearing such lesions at a specific anatomic site (numerator) to the number of animals in which that site was examined (denominator). In most instances, the denominators included only those animals for which that site was examined histologically. However, when macroscopic examination was required to detect lesions prior to histologic sampling (e.g., skin or mammary tumors), or when lesions could have appeared at multiple sites (e.g., lymphomas), the denominators consist of the numbers of animals necropsied.

The purpose of the statistical analyses of tumor incidence is to determine whether animals receiving the test chemical developed a significantly higher proportion of tumors than did the control animals. As a part of these analyses, the one-tailed Fisher exact test (Cox, 1970, pp. 48-52) was used to compare the tumor incidence of a control

group to that of a group of treated animals at each dose level. When results for a number of treated groups, k, are compared simultaneously with those for a control group, a correction to ensure an overall significance level of 0.05 may be made. The Bonferroni inequality (Miller, 1966, pp. 6-10) requires that the P-value for any comparison be less than or equal to 0.05/k. In cases where this correction was used, it is discussed in the narrative section. It is not, however, presented in the tables, where the Fisher exact P-values are shown.

The Cochran-Armitage test for linear trend in proportions, with continuity correction (Armitage, 1971, pp. 362-365), was also used when appropriate. Under the assumption of a linear trend, this test determined if the slope of the dose-response curve is different from zero at the one-tailed 0.05 level of significance. Unless otherwise noted, the direction of the significant trend was a positive dose relationship. This method also provides a two-tailed test of departure from linear trend.

A time-adjusted analysis was applied when numerous early deaths resulted from causes that were not associated with the formation of tumors. In this analysis, deaths that occurred before the first tumor was observed were excluded by basing the statistical tests on animals that survived at least 52 weeks, unless a tumor was found at the anatomic site of interest before week 52. When such an early tumor was found, comparisons were based exclusively on animals that survived at least as long as the animal in which the first tumor was

found. Once this reduced set of data was obtained, the standard procedures for analyses of the incidence of tumors (Fisher exact tests, Cochran-Armitage tests, etc.) were followed.

When appropriate, life-table methods were used to analyze the incidence of tumors. Curves of the proportions surviving without an observed tumor were computed as in Saffiotti et al. (1972). The week during which animals died naturally or were sacrificed was entered as the time point of tumor observation. Cox's methods of comparing these curves were used for two groups; Tarone's extension to testing for linear trend was used for three groups. The statistical tests for the incidence of tumors which used life-table methods were one-tailed and, unless otherwise noted, in the direction of a positive dose relationship. Significant departures from linearity (P < 0.05, twotailed test) were also noted.

The approximate 95 percent confidence interval for the relative risk of each dosed group compared to its control was calculated from the exact interval on the odds ratio (Gart, 1971). The relative risk is defined as p_t/p_c where p_t is the true binomial probability of the incidence of a specific type of tumor in a treated group of animals and p_c is the true probability of the spontaneous incidence of the same type of tumor in a control group. The hypothesis of equality between the true proportion of a specific tumor in a treated group and the proportion in a control group corresponds to a relative risk

of unity. Values in excess of unity represent the condition of a larger proportion in the treated group than in the control.

The lower and upper limits of the confidence interval of the relative risk have been included in the tables of statistical analyses. The interpretation of the limits is that in approximately 95 percent of a large number of identical experiments, the true ratio of the risk in a treated group of animals to that in a control group would be within the interval calculated from the experiment. When the lower limit of the confidence interval is greater than one, it can be inferred that a statistically significant result (a P < 0.025 one-tailed test when the control incidence is not zero, P < 0.050 when the control incidence is zero) has occurred. When the lower limit is less than unity but the upper limit is greater than unity, the lower limit indicates the absence of a significant result while the upper limit indicates that there is a theoretical possibility of the induction of tumors by the test chemical which could not be detected under the conditions of this test.

III. CHRONIC TESTING RESULTS: RATS

A. Body Weights and Clinical Observations

No distinct dose-related mean body weight depression was apparent in either male or female rats (Figure 2).

During the first year of the study the appearance and behavior of the treated rats were generally comparable to that of the controls except that intermittent or occasional hunched appearance, squinted or reddened eyes and urine stains on the abdomen were observed in several treated rats. From week 54 to cessation of dosing in week 78, clinical signs were observed with slightly greater frequency in the treated groups than in the controls, but were noted at comparable rates in treated and control animals during the remainder of the study. Signs commonly associated with aging and observed at comparable rates in control and treated rats included body sores, alopecia, rough or stained fur, palpable nodules or tissue masses and/or bloating. Respiratory signs characterized by labored respiration, wheezing and/or nasal discharge were generally observed at a low incidence in all groups during the study. Isolated, sporadic, and spontaneous observations in one to three treated rats included tremors, head tilt, hyperactivity, apparent hind-limb paralysis and small gonads.

B. Survival

The estimated probabilities of survival for male and female rats in the control and dioxathion-dosed groups are shown in Figure 3.



FIGURE 2 GROWTH CURVES FOR DIOXATHION CHRONIC STUDY RATS 21




For both male and female rats the Tarone test did not indicate a significant association between increased dosage and elevated mortality.

For males adequate numbers of rats were at risk from latedeveloping tumors as 50 percent (25/50) of the high dose, 46 percent (23/50) of the low dose, and 48 percent (24/50) of the control group survived on test until the end of the study. For females the survival was also adequate as 68 percent (34/50) of the high dose, 62 percent (31/50) of the low dose, and 72 percent (36/50) of the control rats survived on test until the end of the study.

C. Pathology

Histopathologic findings on neoplasms in rats are tabulated in Appendix A (Tables Al and A2); findings on nonneoplastic lesions are tabulated in Appendix C (Tables Cl and C2).

A variety of neoplasms was represented among both the treated and control rats. Each of the types of tumors represented has been encountered before as a naturally occurring lesion in the Osborne-Mendel rat and is without apparent relationship to the administration of the chemical.

Inflammatory, degenerative, and proliferative lesions seen in treated and control animals were similar to those naturally occurring in untreated aged rats.

Based upon this histopathologic examination, dioxathion was not toxic or carcinogenic in Osborne-Mendel rats at the doses administered.

D. Statistical Analyses of Results

The results of the statistical analyses of tumor incidence in rats are summarized in Tables 3 and 4. The analysis is included for every type of tumor in either sex where at least two malignant tumors were observed at one site in at least one of the control or dioxathiondosed groups and where such tumors were observed in at least 5 percent of the group.

In male rats the incidence of pituitary chromophobe adenoma was relatively high in the low dose group (30 percent). The Fisher exact test comparing low dose to control, however, had a probability level of P = 0.041, a marginal result which was not significant under the Bonferroni criterion.

No statistical tests indicated a positive association between chemical administration and tumor incidence at any of the other sites tested for male or female rats. Based upon these results there was no convincing evidence of the carcinogenicity of dioxathion in Osborne-Mendel rats.

The possibility of a negative association between dosage and the incidence of hemangiosarcomas of the subcutaneous tissue was observed for female rats. The Fisher exact tests, however, were not significant.

To provide additional insight into the possible carcinogenicity of this compound, 95 percent confidence intervals on the relative risk have been estimated and entered in the tables based upon the

TABLE 3

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TOPOGRAPHY : MORPHOLOGY	CONTROL	LOW DOSE	HIGH DOSE
Subcutaneous Tissue: Hemangiosarcoma	4/49(0.08)	0/50(0.00)	1/50(0.02)
P Values ^C	N.S.	N.S.	N.S.
Relative Risk (Control) ^d		0.000	0.245
Lower Limit Upper Limit		0.000 1.057	0.005 2.362
Weeks to First Observed Tumor	72		77
Spleen: Hemangiosarcoma ^b	4/47(0.09)	0/23(0.00)	2/29(0.07)
P Values ^C	N.S.	N.S.	N.S.
Relative Risk (Control) ^d		0.000	0.810
Lower Limit Upper Limit		0.000 2.136	0.077 5.225
Weeks to First Observed Tumor	90		88
Pituitary: Chromophobe Adenoma ^b	4/41(0.10)	7/23(0.30)	5/27(0.19)
P Values ^C	N.S.	P = 0.041	N.S.
Relative Risk (Control) ^d		3.120	1.898
Lower Limit Upper Limit	 	0.885 12.690	0.448 8.638
Weeks to First Observed Tumor	108	77	94

ANALYSES OF THE INCIDENCE OF PRIMARY TUMORS AT SPECIFIC SITES IN MALE RATS TREATED WITH DIOXATHION⁴

TOPOGRAPHY: MORPHOLOGY	CONTROL	LOW DOSE	HIGH DOSE
Pancreatic Islets: Islet-Cell Adenomab	1/46(0.02)	2/23(0.09)	2/30(0.07)
P Values ^C	N.S.	N.S.	N.S.
Relative Risk (Control) ^d Lower Limit	120	4.000 0.217	3.067 0.166
Upper Limit		225.008	174.643
Weeks to First Observed Tumor	111	111	106
Thyroid: C-Cell Carcinoma ^b	0/48(0.00)	2/49(0.04)	3/49(0.06)
P Values ^C	N.S.	N.S.	N.S.
Relative Risk (Control) ^d Lower Limit Upper Limit	میں سیند میں سیند میں سیند	Infinite 0.290 Infinite	Infinite 0.590 Infinite
Weeks to First Observed Tumor		111	111
Thyroid: C-Cell Adenoma or C-Cell Carcinoma ^b	0/48(0.00)	3/49(0.06)	4/49(0.08)
P Values ^C	N.S.	N.S.	N.S.
Relative Risk (Control) ^d Lower Limit Upper Limit	 سن می می سو می می	Infinite 0.590 Infinite	Infinite 0.909 Infinite
Weeks to First Observed Tumor		111	106

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TABLE 3 (CONTINUED)

TABLE 3 (CONCLUDED)

TOPOGRAPHY : MORPHOLOGY	CONTROL	LOW DOSE	HIGH DOSE
Thyroid: Follicular-Cell Carcinoma ^b	4/48(0.08)	4/49(0.08)	3/49(0.06)
P Values ^C	N.S.	N.S.	N.S.
Departure from Linear Trend ^e	P = 0.038		
Relative Risk (Control) ^d		0.980	0.735
Lower Limit		0.193	0.113
Upper Limit		4.972	4.114
Weeks to First Observed Tumor	106	111	9 4
Thyroid: Follicular-Cell Adenoma or			
Follicular-Cell Carcinoma ^b	5/48(0.10)	8/49(0.16)	7/49(0.14)
P Values ^C	N.S.	N.S.	N.S.
Relative Risk (Control) ^d		1.567	1.371
Lower Limit		0.489	0.403
Upper Limit		5.678	5.119
Weeks to First Observed Tumor	106	99	94

^aTreated groups received time-weighted average doses of 90 or 180 ppm in feed.

^bNumber of tumor-bearing animals/number of animals examined at site (proportion).

^CThe probability level for the Cochran-Armitage test is given beneath the incidence of tumors in the control group when P < 0.05; otherwise, not significant (N.S.) is indicated. The probability level for the Fisher exact test for the comparison of a treated group with the control group is given beneath the incidence of tumors in the treated group when P < 0.05; otherwise, not significant (N.S.) is indicated. For both Cochran-Armitage and Fisher exact tests a negative designation (N) indicates a lower incidence in the treated group(s) than in the control group.

^dThe 95% confidence interval on the relative risk of the treated group to the control group.

^eThe probability level of the test for departure from linear trend is given beneath the control group when P < 0.05.

TABLE 4

ANALYSES OF THE INCIDENCE OF PRIMARY TUMORS AT SPECIFIC SITES IN FEMALE RATS TREATED WITH DIOXATHION^a

TOPOGRAPHY: MORPHOLOGY	CONTROL	LOW DOSE	HIGH DOSE
Subcutaneous Tissue: Hemangiosarcoma	3/50(0.06)	0/50(0.00)	0/50(0.00)
P Values ^C	P = 0.037(N)	N.S.	N.S.
Relative Risk (Control) ^d		0.000	0.000
Lower Limit		0.000	0,000
Upper Limit		1.663	1.663
Weeks to First Observed Tumor	89		
Kidney: Hemangiosarcoma ^b	0/50(0.00)	2/40(0.05)	0/37(0.00)
P Values ^C	N.S.	N.S.	N.S.
Relative Risk (Control) ^d		Infinite	
Lower Limit		0.370	
Upper Limit		Infinite	
Weeks to First Observed Tumor		95	
Pituitary: Chromophobe Adenoma ^b	15/50(0.30)	9/40(0.23)	15/39(0.38)
P Values ^C	N.S.	N.S.	N.S.
Relative Risk (Control) ^d		0.750	1.282
Lower Limit		0.324	0.668
Upper Limit		1.621	2.423
Weeks to First Observed Tumor	100	78	99

TABLE 4 (CONTINUED)

TOPOGRAPHY: MORPHOLOGY	CONTROL	LOW DOSE	HIGH DOSE
Thyroid: C-Cell Carcinoma ^b	0/50(0.00)	2/33(0.06)	2/34(0.06
P Values ^C	N.S.	N.S.	N.S.
Relative Risk (Control) ^d Lower Limit		Infinite 0.449	Infinite 0.433
Upper Limit Weeks to First Observed Tumor		Infinite 95	Infinite 111
Thyroid: C-Cell Adenoma or C-Cell Carcinoma ^b	5/50(0.10)	5/33(0.15)	2/34(0.06
P Values ^C	N.S.	N.S.	N.S.
Relative Risk (Control) ^d Lower Limit Upper Limit		1.515 0.374 6.026	0.588 0.058 3.338
Weeks to First Observed Tumor	111	95	111
Mammary Gland: Fibroadenoma	15/50(0.30)	13/50(0.26)	12/50(0.24
P Values ^C	N.S.	N.S.	N.S.
Relative Risk (Control) ^d Lower Limit Upper Limit		0.867 0.426 1.742	0.800 0.383 1.635
Weeks to First Observed Tumor	87	89	105

TOPOGRAPHY: MORPHOLOGY	CONTROL	LOW HIGH DOSE DOSE 2/34(0.06) 1/31(0.03) N.S. N.S. 0.721 0.395 0.068 0.008 4.696 3.729	
Uterus: Endometrial Stromal Polyp ^b	4/49(0.08)	2/34(0.06)	1/31(0.03)
P Values ^C	N.S.	N.S.	N.S.
Relative Risk (Control) ^d			
Lower Limit Upper Limit			
Weeks to First Observed Tumor	111	106	111

TABLE 4 (CONCLUDED)

^aTreated groups received time-weighted average doses of 45 or 90 ppm in feed.

^bNumber of tumor-bearing animals/number of animals examined at site (proportion).

^CThe probability level for the Cochran-Armitage test is given beneath the incidence of tumors in the control group when P < 0.05; otherwise, not significant (N.S.) is indicated. The probability level for the Fisher exact test for the comparison of a treated group with the control group is given beneath the incidence of tumors in the treated group when P < 0.05; otherwise, not significant (N.S.) is indicated. For both Cochran-Armitage and Fisher exact tests a negative designation (N) indicates a lower incidence in the treated group(s) than in the control group.

^dThe 95% confidence interval on the relative risk of the treated group to the control group.

observed tumor incidence rates. In all of the intervals shown in Tables 3 and 4, the value one is included; this indicates the absence of statistically significant results. It should also be noted that all of the confidence intervals have an upper limit greater than one, indicating the theoretical possibility of tumor induction in rats by dioxathion that could not be established under the conditions of this test.

IV. CHRONIC TESTING RESULTS: MICE

A. Body Weights and Clinical Observations

Distinct, dose-related mean body weight depression was observed among the female mice. This same trend was not, however, apparent in the male mice (Figure 4).

There was no evidence that dioxathion, at the concentrations used in this bioassay, produced any effect on physical appearance or behavior of the treated mice. Signs often observed in group-housed laboratory mice were noted at a comparable rate in control and treated animals, with the incidence increasing gradually as the animals aged. These signs included sores and/or desquamation on parts of the body (more prevalent in males due to fighting), localized alopecia, abdominal urine stains, penile, vulvar or anal irritation, bloated appearance, and palpable nodules and/or tissue masses.

B. <u>Survival</u>

The estimated probabilities of survival for male and female mice in the control and dioxathion-dosed groups are shown in Figure 5. For both male and female mice the Tarone test did not indicate a significant association between increased dosage and elevated mortality.

For males adequate numbers of mice were at risk from latedeveloping tumors as 74 percent (37/50) of the high dose, 70 percent (35/50) of the low dose, and 80 percent (16/20) of the control group survived on test until the end of the study. For females the survival



FIGURE 4 GROWTH CURVES FOR DIOXATHION CHRONIC STUDY MICE



was also adequate as 90 percent (45/50) of the high dose, 92 percent (46/50) of the low dose, and 90 percent (18/20) of the control mice survived on test until the end of the study.

C. Pathology

Histopathologic findings on neoplasms in mice are tabulated in Appendix B (Tables Bl and B2); findings on nonneoplastic lesions are tabulated in Appendix D (Tables Dl and D2).

A variety of neoplasms was present in both the treated and control mice. Each of the types of tumors represented has been encountered previously as a spontaneous lesion in B6C3F1 mice and is apparently unrelated to the administration of dioxathion.

The inflammatory, degenerative, and proliferative lesions that occurred in the control and treated animals were also without appreciable difference from the number and kind of naturally occurring lesions found in untreated aged mice.

Based upon this histopathologic examination, dioxathion was not toxic or carcinogenic in B6C3F1 mice at the doses administered.

D. Statistical Analyses of Results

The results of the statistical analyses of tumor incidence in mice are summarized in Tables 5 and 6. The analysis is included for every type of tumor in either sex where at least two malignant tumors were observed in at least one of the control or dioxathion-dosed groups and where such tumors were observed in at least 5 percent of the group.

TABLE 5

ANALYSES OF THE INCIDENCE OF PRIMARY TUMORS AT SPECIFIC SITES IN MALE MICE TREATED WITH DIOXATHION $^{\rm a}$

TOPOGRAPHY : MORPHOLOGY	CONTROL	LOW DOSE	HIGH DOSE
Subcutaneous Tissue: Fibrosarcoma ^b	1/20(0.05)	5/49(0.10)	3/49(0.06)
P Values ^C	N.S.	N.S.	N.S.
Relative Risk (Control) ^d	**~-	2.041	1.224
Lower Limit Upper Limit		0.254 94.440	0.107 62.958
Weeks to First Observed Tumor	90	73	77
Lung: Alveolar/Bronchiolar Adenoma or Alveolar/Bronchiolar Carcinoma ^b	1/16(0.06)	1/17(0.06)	4/19(0.21)
P Values ^C	N.S.	N.S.	N.S.
Relative Risk (Control) ^d		0.941	3.368
Lower Limit		0.013 69.450	0.387
Upper Limit Weeks to First Observed Tumor	90	90	155.210 90
Liver: Hepatocellular Carcinoma	4/17(0.24)	4/49(0.08)	5/49(0.10)
P Values	N.S.	N.S.	N.S.
Relative Risk (Control) ^d		0.347	0.434
Lower Limit		0.076	0.113
Upper Limit		1.705	2.036
Weeks to First Observed Tumor	90	90	90

TOPOGRAPHY: MORPHOLOGY	CONTROL	LOW DOSE	HIGH DOSE
Liver: Hepatocellular Adenoma or Hepatocellular Carcinoma ^b	4/17(0.24)	4/49(0.08)	6/49(0.12)
P Values ^C	N.S.	N.S.	N.S.
Relative Risk (Control) ^d Lower Limit Upper Limit	 	0.347 0.076 1.705	0.520 0.150 2.331
Weeks to First Observed Tumor	90	90	90

TABLE 5 (CONCLUDED)

^aTreated groups received time-weighted average doses of 284 or 567 ppm in feed.

^bNumber of tumor-bearing animals/number of animals examined at site (proportion).

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^CThe probability level for the Cochran-Armitage test is given beneath the incidence of tumors in the control group when P < 0.05; otherwise, not significant (N.S.) is indicated. The probability level for the Fisher exact test for the comparison of a treated group with the control group is given beneath the incidence of tumors in the treated group when P < 0.05; otherwise, not significant (N.S.) is indicated. For both Cochran-Armitage and Fisher exact tests a negative designation (N) indicates a lower incidence in the treated group(s) than in the control group.

 d The 95% confidence interval on the relative risk of the treated group to the control group.

TABLE 6

ANALYSES OF THE INCIDENCE OF PRIMARY TUMORS AT SPECIFIC SITES IN FEMALE MICE TREATED WITH DIOXATHION^a

TOPOGRAPHY : MORPHOLOGY	CONTROL	LOW DOSE	HIGH DOSE
Hematopoietic System: Malignant Lymphoma ^b	2/19(0.11)	4/50(0.08)	1/49(0.02)
P Values ^C	N.S.	N.S.	N.S.
Relative Risk (Control) ^d		0.760	0.194
Lower Limit		0.122	0.003
Upper Limit		7.931	3.561
Weeks to First Observed Tumor	90	73	91

^aTreated groups received time-weighted average doses of 467 or 935 ppm in feed.

ω 8

^bNumber of tumor-bearing animals/number of animals examined at site (proportion).

^CThe probability level for the Cochran-Armitage test is given beneath the incidence of tumors in the control group when P < 0.05; otherwise, not significant (N.S.) is indicated. The probability level for the Fisher exact test for the comparison of a treated group with the control group is given beneath the incidence of tumors in the treated group when P < 0.05; otherwise, not significant (N.S.) is indicated. For both Cochran-Armitage and Fisher exact tests a negative designation (N) indicates a lower incidence in the treated group(s) than in the control group.

 $^{\rm d}$ The 95% confidence interval on the relative risk of the treated group to the control group.

None of the statistical tests for any site in mice of either sex indicated a significant positive association between the administration of dioxathion and tumor incidence. Thus, at the dose levels used in this experiment there was no convincing statistical evidence that dioxathion was a carcinogen in B6C3F1 mice.

To provide additional insight into the possible carcinogenicity of this compound, 95 percent confidence intervals on the relative risk have been estimated and entered in the tables based upon the observed tumor incidence rates. In all of the intervals shown in Tables 5 and 6, the value one is included; this indicates the absence of statistically significant results. It should also be noted that all of the confidence intervals have an upper limit greater than one, indicating the theoretical possibility of tumor induction in mice by dioxathion that could not be established under the conditions of this test.

V. DISCUSSION

Under the conditions of this bioassay, no significant positive association was established between dietary administration of dioxathion and mortality in Osborne-Mendel rats or B6C3F1 mice of either sex. Adequate numbers of animals in all groups survived sufficiently long to be at risk from late-developing tumors. The possibility that the animals in this bioassay did not receive dosages approximating the maximum tolerated dosages must be considered. Dietary administration of dioxathion had no significant effect on survival in rats or mice of either sex and it affected body weight gain in only the female mice. No particularly unusual clinical observations were reported during the bioassay in either sex of either species.

A variety of neoplasms was observed in treated animals of both species; however, none of the neoplasms observed were either histopathologically unusual or in statistically significant incidences.

Under the conditions of this bioassay, dietary administration of dioxathion was not carcinogenic in Osborne-Mendel rats or B6C3F1 mice.

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Review of the Bioassay of Dioxathion* for Carcinogenicity by the Data Evaluation/Risk Assessment Subgroup of the Clearinghouse on Environmental Carcinogens

April 26, 1978

The Clearinghouse on Environmental Carcinogens was established in May, 1976, in compliance with DHEW Committee Regulations and the Provisions of the Federal Advisory Committee Act. The purpose of the Clearinghouse is to advise the Director of the National Cancer Institute (NCI) on its bioassay program to identify and to evaluate chemical carcinogens in the environment to which humans may be The members of the Clearinghouse have been drawn exposed. from academia, industry, organized labor, public interest groups, State health officials, and guasi-public health and research organizations. Members have been selected on the basis of their experience in carcinogenesis or related fields and, collectively, provide expertise in chemistry, biochemistry, biostatistics, toxicology, pathology, and epidemiology. Representatives of various Governmental agencies participate as ad hoc members. The Data Evaluation/ Risk Assessment Subgroup of the Clearinghouse is charged with the responsibility of providing a peer review of reports prepared on NCI-sponsored bioassays of chemicals studied for carcinogenicity. It is in this context that the below critique is given on the bioassay of Dioxathion for carcinogenicity.

The primary reviewer said that the compound was not carcinogenic in rats or mice, under the conditions of test, but did cause testicular atrophy in male rats. He said the finding was particularly important since other pesticides have been shown to produce testicular damage in exposed chemical or farm workers. The primary reviewer commented on a dose-related increase in the incidence of hyperplastic nodules and nodular hyperplasia of the liver in male mice, as well as on other lesions observed at elevated levels in the treated animals. Given the widespread exposure to Dioxathion, he recommended that the non-neoplastic findings were sufficiently important to be noted in the report's discussion and summary sections. Based on the bioassay, the primary reviewer concluded that Dioxathion does not pose a carcinogenic risk to humans, although it may present a risk for testicular damage.

The secondary reviewer agreed with the conclusion in the report that Dioxathion was not carcinogenic under the conditions of test. He noted that the dose administered was increased during the course of the chronic study to achieve a maximum tolerated dose.

A motion was approved unanimously that the report on the bioassay of Dioxathion be accepted as written.

Members present were

Michael Shimkin (Acting Chairman), University of California at San Diego Joseph Highland, Environmental Defense Fund George Roush, Jr., Monsanto Company Louise Strong, University of Texas Health Sciences Center John Weisburger, American Health Foundation (Sidney Wolfe, Health Research Group, submitted a written review)

* Subsequent to this review, changes may have been made in the bioassay report either as a result of the review or other reasons. Thus, certain comments and criticisms reflected in the review may no longer be appropriate.

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SUMMARY OF THE INCIDENCE OF NEOPLASMS IN RATS TREATED WITH DIOXATHION

APPENDIX A

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TABLE A1 SUMMARY OF THE INCIDENCE OF NEOPLASMS IN MALE RATS TREATED WITH DIOXATHION

	CONTROL (VBH) 01-M001	LOW DOSE 01-M008	HIGH DOSE 01-m009
NIHALS INITIALLY IN STUDY	50		50
NINALS NECROPSIED	49	50	50
NIMALS EXAMINED HISTOPATHOLOGICALLY	** 49	50	49
NTEGUNENTARY SYSTEM			
* SKIN	(49)	(50)	(50)
HEMANGIO FERICITONA, MALIGNANT	1 (2%)	• •	• •
*SUBCUT TISSUE	(49)	(50)	(50)
FIBROMA	1 (2%) 1 (2%)		1 (2\$)
PIBROS ARCOMA	1 (2%)	2 (4%)	
LIPONA			2 (4%)
HENANGIOS ABCONA	4 (8%)		1 (2%)
ESPIRATORY SYSTEM			
#LUNG	(49)	(39)	(40)
SIXED TOMOR, MALIGNANT	1 (2%)		
HENANGIOSARCONA, HETASTATIC	2 (4%)		
ENATOPOIETIC SYSTEM			
*NULTIPLE ORGANS	(49)	(50)	(50)
MALIG.LYMPHONA, LYMPHOCYTIC TYPE	1 (2%)		1 (2%)
LYNPHOCYTIC LBUKEMIA			1 (2%)
*SUBCOT TISSUE/BACK	(49)	(50)	(50)
HALIG. LYHPHONA, HISTIOCYTIC TYPE		1 (2%)	1
#SPLEEN	(47)	(23)	
HENANGIOSARCONA	4 (9%)		2 (7%)
TROULATORY SYSTEM			
€HEART HEMANGIOSARCONA	(47) <u>1 (25)</u>	(24)	(28)

NUMBER OF ANIMALS WITH TISSUE EXAMINED NICROSCOPICALLY * NUMBER OF ANIMALS HECROPSIED **Excludes partially autolyzed animals

TABLE A1 (CONTINUED)

	CONTROL (VEH) 01-8001	LOW DOSE 01-M008	HIGH DOS B 01- <i>M</i> 009
HEMANGIOSARCOMA, NETASTATIC	1 (2%)		
<pre>#ENDOCARDIUM SARCONA, NOS</pre>	(47)	• •	(28) 1 (4%)
JIGESTIVE SYSTEM			
*LIVER HEMANGIOSARCOMA, METASTATIC	(49) 1 (2 %)	(37)	túr):
<pre>PANCREAS ADENOCARCINOMA, NOS, METASTATIC ACINAB-CELL ADENOMA</pre>	(46)	(23) 1 (4%)	(30) 1 (3%)
NSTOMACH Adenocarcinona, nos	(46)	(29) 1 (3%)	(29)
#SMALL INTESTINE Adenocarcinoma, NGS, Metastatic	(46)	(21) 1 (5%)	(28)
#COLON ADBBOCARCINONA, NOS, METASTATIC	(46)	(20) 1 (5%)	(28)
URINARY SYSTEM			
4XIDNEY LIPONA	(47) 2 (4 %)	(38)	(41)
LIPOSARCONA MIXED TUMOB, HALIGNANT HENANGIOSARCONA	t (2 %)	1 (3%)	2 (5%)
#URINARY BLADDER PAPILLOMA, NOS	(46) 3 (7%)	(25)	(26 <u>}</u>
ENDOCRINE SISTEM			
FITUITARY CHROMOPHOBE ADENOMA	(41) 4 (10%)	(23) 7 (30%)	(27) 5 (19%)
#ADRENAL LIPOSARCOMA	(46)	(21) 1 (5%)	(29)
#THYROID FOLLICULAR-CELL ADENOMA	(46) <u>3 (6%)</u>	(49) <u>4 (85)</u>	(49) <u>4 (8%)</u>

* NUMBER OF ANIMALS WITH TISSUE EXAMINED MICROSCOPICALLY * NUMBER OF ANIMALS NECROPSIED

TABLE AT (CONTINUED)

	CONTROL (VEH) 01~0001		HIGH DOSE 01-MCO9
FOLLICULAR-CELL CARCINONA C-CELL ADENONA C-CELL CARCINONA	4 (9%)	4 (8%) 1 (2%) 2 (4%)	3 (6%) 1 (2%) 3 (6%)
#PARATHYROID ADBRONA, NOS	(46) 1 (2%)	(27)	(21)
IPANCREATIC ISLETS ISLET-CELL ADENOMA	(46) 1 (2%)	(23) 2 (9%)	(30) 2 (7 %)
FRODUCTIVE SYSTEM			
*AAMMARY GLAND Adenoma, Nos Adenocarcinoma, Nos Pibroadenona	(49) 2 (4系) 1 (2 系)	(50) 1 (2%) 2 (4%)	(50)
PROSTATE HENANGIOSARCONA, METASTATIC	(34) 1 (3%)	(15)	(8)
SEMINAL VESICLE Henangiosarcoma, metastatic	(49) 1 (2%)	(50)	(50)
BVOUS SYSTEM			
BRAIN GLIONA, NOS	(47) 1 (2 %)	{24)	(28)
ECIAL SENSE ORGANS			
NONE	****		
USCULOSKELETAL SYSTEM			
SKELETAL NUSCLE FIBROSARCOMA	(49) 1 (2%)	(50)	(50)
HUSCLE OF BACK HEMANGIOSARCOMA	(49)	(50)	(50) 1 (2 %)
*MUSCLE OF THORAX BENANGIOSABCOBA	(49) 1 (2%)	(50)	(50)

NUMBER OF ANIMALS WITH TISSUE EXAMINED MICROSCOPICALLY
 NUMBER OF ANIMALS NECROPSIED

TABLE AI (CONCLUDED)

	CONTROL (VEH) 01-M001	LOW DOSE	HIGH DOSE
	01-M001	01-8008	01-8009
DY CAVITIES			
ABDONINAL CAVITY Lipoma	1 (2%)	(50)	(50)
L OTHER SYSTENS			
MULTIPLE ORGANS REMANGIOSARCOMA	1 (2%)	(50)	
IMAL DISPOSITION SUBMARY			
ANIMALS INITIALLY IN STUDY NATURAL DEATHƏ NORIBUND SACRIPICE SCHEDULED SACRIPICE	50 24 2	50 25 2	50 25
ACCIDENTALLY XILLED Terminal sacrifice Avimal bissing	24	23	25
INCLUDES AUTOLYZED ANIMALS		**-**	
MOR SUMMARY			
TOTAL ANIMALS WITH PRIMARY TUNORS* Total Primary Tumors	28 41	23. 29	21 31
TOTAL ANIMALS WITH BENIGN TUNORS TOTAL BENIGN TUMORS	13 17	14 15	14 16
TOTAL ANIMALS WITH MALIGNANT TUBORS TOTAL MALIGNANT TUBORS	1B 24	12 14	12 15
TOTAL ANIMALS WITH SECONDARY TUMORS TOTAL SECONDARY TUMORS	# 2 6	"1 ≆	
TOTAL ANIMALS WITH TUMORS UNCERTAIN BENIGN OR KALIGNANT TOTAL UNCERTAIN TUMORS	-		
TOTAL ANIMALS WITH TUMORS UNCERTAIN- PRIMARY OR BETASTATIC Total Uncertain Tumors	-		
PRIMARY TUMORS: ALL TUMORS BICBPT S SECONDARY TUMORS: NETASTATIC TUMORS			

 TABLE A2
 SUMMARY OF THE INCIDENCE OF NEOPLASMS IN FEMALE RATS TREATED WITH DIOXATHION

	CON TROL (VEH) 01-2001	LOW DOSE 01-F010	HIGH DOSE 01-F011
	50		50
NIMALS NECROPSIED	50	50	50
WINALS EXAMINED HISTOPATHOLOGICALLY		47	48
INTEGUMENTARY SISTEM			
*SUBCUT TISSUE	(50)	(50)	(50)
BASAL-CELL CARCINOMA	1 (2%)		
FIBRONA	2 (4%)	2 (4%)	
FI BROSAR COMA		1 (2%)	1 (2%)
LIPONA	1 (2%)		
HEMANGIOSARCOMA	3 (6%)		
ESPIRATORY SYSTEM			
#LUNG		(39)	
ADENOCARCINOMA, NOS, METASTATIC			1 (2%)
REMATOPOIETIC SISTEM			
ICERVICAL LINPH NODE MALIG.LINPHONA, LYMPHOCYTIC TYPE	(48)	(31)	(28)
MALIG.LIMPHONA, LYMPHOCYTIC TYPE	1 (2%)		
CIRCULATORY SYSTEM			
#PHDOCARDIUN	(50)	(34)	(33)
SARCONA, NOS			2 (6%)
DIGESTIVE SYSTEM			
ŧlişer	(50)	(44)	(40)
NBOPLASTIC NODULE	1 (2%)		
SPANCREAS OSTEOSABCONA	(50)	(35)	(29)

NUMBER OF ANIMALS WITH TISSUE EXAMINED MICROSCOPICALLY
 # NUMBER OF ANIMALS NECROPSIED
 **EXCLUDES PARTIALLY AUTOLYZED ANIMALS

TABLE A2 (CONTINUED)

	CONTROL (VEH) 01-P001	LOW DOSE 01-P010	HIGH DOSE 01-P011
#STONACH OSTEOSARCONA	(50)	[32]	(39) 1 (3%)
É CUODE NUN HEHANGIOS à RCOMÀ OSTEOS à RCOMÀ	(50) 1 (2%)	į 3kį	(32) 1 (3%)
AILEOM OSTBOSARCONA	(50)	(34)	(32) 1 (3%)
RINARY SYSTEM			
*KIDNEY LIPONA Mixed Tuhor, Malignart Hemangiosarcona Hanartona +	(50) 1 (2%) 1 (2%) 1 (2%)	(# 0) 2 (5%)	(37)
SUPINARY BLADDEP Papillona, Nos	[49]	(34)	[32) 1 [3%}
NDOCRINE SYSTEM			
*FITUITARY CHROMOPHOBE ADENONA	(50) 15 (30%)	(40) 9 (23%)	(39) 15 (38%)
*THYROID POLIICULAR-CELL ADENOMA POLLICULAR-CELL CARCINONA	(50) 1 (25)	(33)	(34) 1 (3 %)
C-CELL ADENOMA C-CELL CARCINOMA	5 (10%)	3 (9%) 2 (6%)	2 (6%)
#PANCREATIC ISLETS ISLET-CELL ADENOMA	(50)	(35)	(29) 1 (3%)
EFRODUCTIVE SYSTEM			
*BANNARY GLAND Adbnona, Kos Adenocarcingna, Nos	(50)	(50) 1 (2%)	(50) 2 (4 %)
PIBROADENONA	15 (30%)	13 (265)	

NUMBER OF ANIMALS WITH TISSUE EXAMINED NICROSCOPICALLY
 NUMBER OF ANIMALS NECROPSIED
 THIS IS CONSIDERED TO BE A BENIGN FORM OF THE MALIGNANT MIXED TUMOR OF THE KIDNEY AND CONSISTS OF PROLIFERATIVE LIPOCYTES, TUBULAR STRUCTURES, FIBROBLASTS, AND VASCULAR SPACES IN VARYING PROPORTIONS

TABLE A2 (CONTINUED)

	CONTROL (VEH) 01-P001	LON DOSE 01-F010	HIGH DOSE 01-P011
UTERUS	(49)	(34)	(31)
LEIONYCMA Endometrial stronal polyp	4 (8%)	2 (6%)	1 (3%) 1 (3%)
OVARY	(49)		(31)
CARCINOMA, NOS	1 (2%)	(•••	• •
LUTEOMA GRANULOSA+CBLL TUMOR	1 (2%)		1 (3%)
OUS SYSTEN			
N B			
IAL SENSE ORGANS			
340		····	
ULOSKELETAL SYSTEM			
DNE		_	
Y CAVITIES			
DOMINAL VISCERA HBHANGIOSARCOMA	(50) 1 (2%)	(50)	(50)
OTHER SYSTEMS			
HOBACIC CAVITY FIBROSARCOMA			1
• • • · · · · · · · · · · · · · · · · ·			
AL DISPOSITION SUMMARY			
NIMALS INITIALLY IN STODY	50	50	50
NATURAL DEATHƏ Moribund Sacrifice Scheduled Sacrifice	14	19	15 1
ACCIDENTALLY KILLED TERMINAL SACRIFICE	36	31	34
ANIMAL MISSING		~-	
CLUDES AUTOLYZED ANIHALS			

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TABLE A2 (CONCLUDED)

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		LOW DOSE 01-P010	HIGH DOSE 01-F011
TUNOR SUNMARY			
TOTAL ANIMALS WITH PRIMARY TUMORS*	36	27	29
TOTAL FRIMARY TUNORS	56	35	45
TOTAL ANIMALS WITH BENIGN TUMORS	31	25	26
TOTAL BENIGN TUMORS	44	30	33
TOTAL ANIMALS WITH MALIGBANT TOMORS	9	4	9
TOTAL MALIGNANT TUMORS	10	4	12
TOTAL ANIMALS WITH SECONDARY TUMORS	ŧ		۱
TOTAL SECONDARY TUMORS			1
TOTAL ANIMALS WITH TUMORS UNCERTAIN-	-		
BENIGN OR MALIGNANT	2		
TOTAL UNCERTAIN TUMORS	2.		
TOTAL ANIMALS WITH TUBORS UNCERTAIN-	-		
PRIMARY OR METASTATIC			
TOTAL UNCERTAIN TUMORS			
* PRIMARY TUMORS: ALL TUMORS BACEPT S	ROONDARY THMORS	5	

SUMMARY OF THE INCIDENCE OF NEOPLASMS IN MICE TREATED WITH DIOXATHION

APPENDIX B

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TABLE B1
SUMMARY OF THE INCIDENCE OF NEOPLASMS IN MALE MICE TREATED WITH DIOXATHION

	CONTROL (VER) 02-M011	LON DOSE 02-N012	HIGH DOSB 02-8013
	20	şp	50 1
NIHALS BECROPSIED NIMALS EXAMINED HISTOPATHOLOGICALLY	20 ** 19	49 49	49 49
NTEGUMENTARY SYSTEM			
*SKIN SQUAMOUS CELL CARCINONA	(29)	(49) 1 (2 %)	(49)
*SUBCUT TISSUE PIBROSARCONA OSTEOSARCONA	(20) 1 (5%) 1 (5%)	(49) 5 (10%)	(49) 3 (6 %)
RESPIRATORY SYSTEM			
#LUNG ALVEOLAR/BRONCHIOLAR ADENOMA ALVEOLAB/BBONCHIOLAR CARCINOMA	(16) 1 (6%)	(17) 1 (6 %)	(19) 3 (16%) 1 (5%)
IEMATOPOIETIC SYSTEM			
*MULTIPLE OBGANS GBANULOCYTIC LEUKEMIA	(20)	(49)	(49) 1 (2%)
*LYMPH NODE MALIG.LIMPHOMA, HISTIOCYTIC TYPE	(15)	(13) 1 (8%)	(12)
CIRCULATORY SYSTEM			
NONE			
DIG esti∛e System			
*LIVER HEPATOCELLULAR_ADENONA	(17)	(49)	(49)

B-3

TABLE BI (CONTINUED)

	CONTROL (YEH) 02-0011	A2-8612	HIGH DOSE 02-M013
HEPATOCELLULAR CARCINONA PIBROSARCONA, METASTATIC HEMANGIOSARCONA		• [2 m]	5 (10%) 1 (2%)
URINARY SYSTEM			
NONE			
ENDOCRINE SYSTEM			
NONE			
REPRODUCTIVE SYSTEM			
NONE			
NERVOUS SYSTEM			
NONE			
SPECIAL SENSE ORGANS			
NONE			
NUSCOLOSKELETAL SYSTEM			
*SKELETAL MUSCLE FIBROSARCONA	(20)	(49)	(49) 1 (2%)
BODY CAVITIES			
*ABDOMINAL CAVITY LIPONA	(20)	(49) 1 (2%)	(49) 1 (2%)
ALL OTHER SYSTEMS			
NONE			
# NUMBER OF ANIMALS WITH TISSOR P	IAMINED MICROSCOPIC	ALLY	

* NUMBER OF ANIMALS NECROPSIED

B-4
TABLE B1 (CONCLUDED)

	02-8011	LOW DOSE 02-0012	02-8013
INAL DISPOSITION SUMMARY			
ANIMALS INITIALLY IN STUDY WATURAL DBATHƏ MORIBUND SACBIPICE SCHEDULED SACRIFICE	29 ₄	50 14	50 1) - 1
ACCIDENTALLY KILLED Terminal Sacrifice Animal Missing	Ť6	35 , 1	37 1
INCLUDES AUTOLYZED ANIMALS			
HOR SOUNARY			
TOTAL ANIMALS WITH PRIMARY TUMORS* Total Primary Tumors	7 8	10 1+3	13 17
IOTAL ANIMALS WITH BENIGH TUMORS TOTAL BENIGN TUMORS	2 2	1 1	4 5
TOTAL ANIMALS WITH MALIGNANT TUBORS TOTAL MALIGNART TUBORS	6 6	19 12:	11 12
OTAL ANIMALS WITH SECONDARY TUMORS Total Secondary Tumors	•	¹ 1	
OTAL ANIMALS WITH TUBORS UNCERTAIN ENIGN OR MALIGNANT TOTAL UNCERTAIN TUMORS	-		
NOTAL ANIMALS WITH TUMORS UNCERTAIN FRIMABY OR METASTATIC TOTAL UNCERTAIN TUMORS	-		
PRIMARY TUMORS: ALL TUMORS BRCEPT SI Secondary Tumors: Metastatic Tumors			DJACZNT OBGAN

 TABLE B2

 SUMMARY OF THE INCIDENCE OF NEOPLASMS IN FEMALE MICE TREATED WITH DIOXATHION

	02-2011	LOW EOSE 02-P014	
ANIMALS INITIALLY IN STUDY	20	50	50
NIMALS HISSING	1		1
ANIMALS NECROPSIED ANIMALS EXAMINED HISTOPATHOLOGICALLY	19	50 50	49 49
			+0
NTEGUMENTARY SYSTEM			
*SUBCUT TISSUE	(19)	(50)	(49)
MYXOS ARCOMA			1 (2%)
RESPIRATORY SYSTEM			
	(19)	(15)	(12)
ALVEOLAR/BRONCHIOLAR ADENONA	(18)	1 (75)	1 (8%)
EMATOPOIETIC SYSTEM *HUITIPLE ORGANS MALIG.LYMPHONA, HISTIOCYTIC TYPE		(50)	(49)
*ABDOMINAL CAVITY MALIGNANT LYMPHOMA, NOS	(19)	(50) 1 (2%)	(49)
#MESENTERIC L. NODE MALIGNANT LYMPHOMA, MIXED TYPE	(17) 1 (6%)	(10)	(12)
<pre>#LIVER MALIG.LYMPHONA, LYMPHOCYTIC TYPE MALIG.LYMPHONA, HISTIOCYTIC TYPE</pre>	(18)	(50) 1 (25) 1 (25)	(48) 1 (2%)
#UTERUS MALIG.LYMPHONA, HISTIOCYTIC TYPE	(18)	(16) 1 (6≸)	(21)

CIRCULATORY SYSTEM

NONE_____

* NUMBER OF ANIMALS WITH TISSUE EXAMINED MICROSCOPICALLY * NUMBER OF ANIMALS NECROPSIED

**EXCLUDES PARTIALLY AUTOLYZED ANIMALS

.

TABLE B2 (CONTINUED)

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			* * * * * * * * * * *	
	Control (VEH) 02-F011	LOW DOSE 02-F014	HIGH DOSE 02-7015	
DIGESTIVE SYSTEM			•	
#LIVER HEPATOCRILULAR CARCINONA HEMANGIOSARCOMA	(18)	(50) 1 (2%)	(48) 1 (2 %)	
URINARY SYSTEM				
BONE				
ENDOCRINE SYSTEM				
NONE				
REPRODUCTIVE SYSTEM				
*MAMMARY GLAND ADENGCARCINOMA, NOS	(19)	(50)	(49) 2 (4%)	
NERVOUS SYSTEM				
NONE				
SPECIAL SENSE ORGANS				
NONB				
NUSCULOSKELETAL SYSTEM				
None				
BODY CAVITIES				
NONE				
ALL OTHER SYSTEMS				
NORE				
4 NUMBER OF ANIMALS WITH TISSUE E * NUMBER OF ANIMALS HECROPSIED	XAMINED MICROSCOPI	CALLY		

TABLE B2 (CONCLUDED)

	CONTROL (VER) 02-F011	02-2014	BIGE DOSE 02-F015	
NINAL DISPOSITION SUMMARY				
ANINALS INITIALLY IN STUDY Natural deathg Moribund Sacrifice Scheduled Sacrifice	20 1	50 4	50 4	
ACCIDENTALLY KILLED Terminal sacrifice Animal missing	18 1	46	45 1	
INCLUDES AUTOLYZED ANIMALS				
UNOR SUMMARY				
TOTAL ANIMALS WITH PRIMARY TUMORS* Total primary tumors	22	6 6	6 6	
TOTAL ADIMALS WITH CENIGN TUMORS TOTAL BENIGN TUMORS		1 1	ች - ሺ	
TOTAL ANIMALS WITH MALIGNANT TUMORS TOTAL MALIGNANT TUMORS	2 2	<u>হ</u>	s S	
TOTAL ANIMALS WITH SECONDARY TUNORS TOTAL SECONDARY TUNORS	•			
TOTAL ANIMALS WITH TUMORS UNCERTAIN BENIGN OR NALIGNANI TOTAL UNCERTAIN TUMORS	-			
TOTAL ANIMALS WITH TUMORS UNCERTAIN FRIMABY OR METASIATIC TOTAL UNCERTAIN TUMORS	-			
PRINARY TUMORS: ALL TUMORS EXCEPT S Secondary Tumors: Metastatic Tumors				

APPENDIX C

SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN RATS TREATED WITH DIOXATHION z.

TABLE CI	
SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN MALE RATS TREATED WITH DIOXATHION	

.

	CONTROL (VEN) 01-N001	LOW DOSE 01-8000	HIGH DOSE 01-MD09
ANIMALS INITIALLY IN STUDY Animals necropsied Animals byanined histopathologica	50 49	50 50 50	50 50 49
NTEGONENTARY SYSTEM			
*SKIN	(49)	(50)	(50)
EPIDERMAL INCLUSION CYST Inflammation, nos	1 (2%)	1 (2%)	2 (4%)
*SUBCUT TISSUE Abscess, Nos	(49) 1 (2%)	(50) 1 (2 %)	(50)
RESPIRATORY SYSTEM			
#T₽⊼CH8A	(4)		
INFLAMMATION, NOS	4 (100%)		
#LUNG Atelectasis	(49)	(39) 1 (3%)	(40)
PNEUHONIA, CHRONIC MURINE	20 (41%)	19 (49%)	17 (43\$)
HENATOPOIETIC SYSTEM			
#SPLEEN		(23)	(29)
PIBROSIS Henatopoiesis	1 (2%) 1 (2%)	3 (13%)	2 (7%)
#BESENTERIC L. NODE	(45)	(19)	(21)
CYST, NOS Inplandation, Nos	1 (2%)	1 (6%) 1 (6%)	
		· (4.4)	
CIRCULATORY SYSTEM			
#MYOCARDIUM INFLAMMATION, NOS	(47) <u>14_(30%)</u>	(24) 1 (AS)	(28) <u>1 (45)</u>

TABLE CI (CONTINUED)

	CONTROL (VBH) 01-4001	LON COSE 01-8008	HIGH DOSE 01-8009
PIBROSIS Degeneration, Nos	1 (2%)		3 (11%)
ENDOCARDIUM HYPERPLASIA, NOS	(47)	(24) 1 (4%)	(28) 1 (4 %)
*AORTA PERIARTERITIS ARTERIOSCLEROSIS, NOS	{49} 4 (8%)	(50) 1 (2\$)	(50)
GESTIVE SYSTEM			
LIVER DILATATION, NOS CIST, NOS INFLAMMATION, NOS MBTANOBPHOSIS PATTY HYPERPLASIA, NOS	(49) 2 (9%) 3 (6%) 3 (6%) 5 (10%)	(37) 1 (3%) 2 (5%) 1 (3%) 7 (19%) 1 (3%)	(44) 3 (7%) 3 (7%)
BILE DUCT HYPERPLASIA, NOS	(49) 3 (6%)	(50) 5 (10%)	(50) 1 (2%)
FANCREAS PERIARTERITIS DEGENERATION, NOS ATROPHY, NOS	(46) 5 (11%)	(23) 3 (13%) 1 (4%) 1 (4%)	(30) 1 (3%)
ESOPHAGUS INFLAMMATION, NOS	(1) 1 (100%)		
STONACH INFLAMMATION, NOS ULCER, POCAL CALCIUM DEPOSIT CALCIPICATION, NOS	(46) 1 (2 %)	(29) 1 (3%) 1 (3%) 7 (24%)	(29) 3 (10%) 2 (7%) 1 (3%)
LARGE INTESTINE NEMATODIASIS	(46)	(2ØJ	(28) 1 (4 5)
COLON INFLAMMATION, NOS PARASITISM	(46) 1 (2%)	(20)	(28) 1 (4%)
INARY SYSTEM			
KIDNEY HIDROBEPHROSIS	(47)	(38) <u>1_(3%)</u>	(41)

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NUMBER OF ANIMALS WITH TISSOE BIANINED MICROSCOPICALLY * NUMBER OF ANIMALS NECROPSIED

TABLE C1 (CONTINUED)

	CONTROL (VEH) 01-1001	LON DOSE 01-NO08	HIGH DOSE 01-x009
PYELONEPHEITIS, NOS PYONEPHEOSIS	2 (4%) 1 (2%)	***	1 (2%)
PYONEPHROSIS Abscess, Nos	1 / 281		
INPLAMMATION, CHRONIC	37 (79%)	30 (79%) 3 (8%)	36 (88%)
CALCIUM DEPOSIT		3 (8%)	3 (7%)
#URINARY BLADDER	(46)	(25)	(26)
INFLAMMATION, NOS	1 (2%)	1 (4\$)	1 (4%)
NDOCRINE SYSTEM			
#PITULTARY	(4 1)	(23)	(27)
CYST, NOS	1 (2%)		1 (4%)
#ADRENAL	1461	(21)	(29)
CALCIUM DEPOSIT			± (3≸)
ANGIECTASIS	8 (17%)	5 (24%)	5 (17%)
#THYROID	(48)	(49)	(49)
CIST, NOS	1 (25)	• •	
FOLLICULAR CYST, NOS			2 (4%)
HYPERPLASIA, PAPILLABY			1 (2%)
HTPERPLASIA, C-CELL	1 (2%)	1 (25)	
HYPERPLASIA, POLLICULAR-CELL	4 (8\$)	3 (6%)	3 (6%)
* FA RATHY ROID	(46)	(27)	(21)
HYPERPLASIA, NOS	2 (4%)	7 (26%)	5 (24%)
BPRODUCTIVE SYSTEM			
*MAMMARY GLAND	(49)	(50)	(50)
GALACTOCELE	(49) 1 (2%)	1 (2%)	1 (2%)
CYST, NOS	1 (2%)		
U PROSTATE	(34)	(15)	(8)
INFLAMMATION, NOS	(34) 9 (26%)	1 (7%)	
*SEMINAL VESICLE	(49)	(50)	(50)
INFLAMMATION, NOS	1 (2%)	1 (2%)	1 (2%)
HYPEBPLASIA, NOS	· •,		1 (2%)
#TESTIS	(44)	(20)	(27)
ATROPHY. NOS	9 (20%)	9 (458)	10 (328)

NUMBER OF ANIMALS WITH TISSUE EXAMINED DICROSCOPICALLY * NUMBER OF ANIMALS BECROPSIED

TABLE CI (CONTINUED)

	CONTROL (VEN) 01-8001	LON DOSE 01-0008	HIGH DOSE 01-009	
*EPIDIDINIS Atrophy, Nos	(49)	{50} 1 (2%)	(50) 2 (4\$)	
NERVOUS SYSTEM				
NONE				
SPECIAL SENSE ORGANS				
•EYE Phthisis Bulbi	(49) 1 (2%)	(50)	(50)	
*FYE/LACRIMAL GLAND INFLAMMATION, NOS	(49)	(50) 2 (4%)	(50)	
+HARDERIAN GLAND INPLANMATION, NOS	(49) 1 (2%)	(50)	(55)	
NUSCULOSKELETAL SYSTEM				
*SKULL INFLAMMATION, NOS FIBROSIS	(49)	(50) 1 (2%) 1 (2%)	(50) 1 (2%) 1 (2%)	
*SKELETAL MUSCLE Degeneration, Nos	(49) 1 (2 %)	(50)	(50)	
BODY CAVITIES				
*FERICARDIUM Inflammation, Nos	(49) 5 (10%)	(50) 3 (6 %)	(5 P j	
•MESENTERY PERIARTERITIS	(49) 2 (4%)	(50)	(50)	
ALL OTHER SYSTEMS				
NONE				

NUMBER OF ANIMALS WITH TISSUE BYANIMED MICROSCOPICALLY * NUMBER OF ANIMALS NECROPSIED

TABLE CI (CONCLUDED)

	CONTROL (VBH) 01-0001	LON DOSE 01-8008	HIGH DOSE 01-0009	
SPECIAL MORPHOLOGY SUMMARY				
NO LESION REPORTED Auto/Wechopsy/Histo Perp Auto/Wechopsy/No Histo Auto/Wechopsy/No Wechopsy	1	1	2 1 1	
# NUMBER OF ANIMALS WITH TISSUE ERANI) * NUMBER OF ANIMALS NECROPSIED	NED MICROSCOPIC	ALLY		

TABLE C2 SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN FEMALE RATS TREATED WITH DIOXATHION

	CONTROL (VBH) 01-9001	LOW COSE 01-P010	HIGH DOSE 01-P011
	50 50	50 50	50 50 48
NTEGUMENTARY SYSTEM			
*SKIN EPIDERHAL INCLUSION CIST INFLAMMATION, NOS ACANTHOSIS	(50) 1 (2%)	(50) 1 (2%) 1 (2%)	(50) 1 (2%) 1 (2%)
*SUBCUT TISSUE Abscess, Nos		(50)	(50) 1 (2%)
ESPIRATORY SYSTEM			
TRACHEA INFLAMMATION, NOS	(5) 5 (100%)		
<pre>#LUNG INPLANDATION, NOS PNEUMONIA, CHRONIC MURINE HIPERPLASEA, ADEMONIATOUS</pre>		(39) 19 (49%)	1 (25)
ENATOPOIETIC SYSTEM			
#BONE MARROW NETAHORPHOSIS FATTY	(50) 1 (2%)	(33)	(31)
ISPLEEN HEMORRHAGE REMATOPOIESIS	(50) 5 (10%)	(34) 1 (3%) 2 (6%)	(35) 7 (20 %)
IPCULATORY SYSTEM			
ANYOCARDIUM	(50)		(33)

* NUMBER OF ANIMALS NECROPSIED **EXCLUDES PARTIALLY AUTOLYZED ANIMALS

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TABLE C2 (CONTINUED)

	CONTROL (VEH) 01-P001	LON DOSE 01-F010	HIGH DOSE 01-P011
DEGENERATION, NOS	2 (4%)	4 (12%)	
#ENDOCARDIUM HTPERPLASIA, NOS	(50) 1 (2%)	(34) 1 (3%)	(33)
*AORTA ARTERIOSCLEROSIS, NOS	(50) 1 (2 %)	(50) 1 (2 %)	(50)
DIGESTIVE SYSTEN			
*LIVER INFLAMMATION, NOS METANORPHOSIS PATTY POCAL CELLULAR CHANGE ANGIECTASIS	(50) 4 (8%) 7 (2%) 1 (2%)	(44) 1 (25)	(40) 1 (3%) 1 (3%)
*BILE DUCT DILATATION, NOS HYPERPLASIA, NOS	(50) 1 (2%) 2 (4%)	(50)	(50) 1 (2%) 1 (2%)
♥FANCREAS PERIARTERITIS	(50)	(35) 1 (3%)	(29)
#STONACH INPLAMMATION, NOS ULCEB, POCAL CALCIUM DEPOSIT	(50) 5 (10%) 1 (2%)	(37) 1 (38) 1 (3%)	(39) 1 (3%) 3 (8%)
#LARGE INTESTINE Parasitism	(49) 1 (2%)	(34)	(29)
#COLON PARASITISM	(49)	(34) 4 (12 %)	(29)
URINARY SYSTEM			
*KIDNEY HINERALIZATION CYST, NOS	(50)	(4 0)	(37) 1 (3%) 1 (3%)
PYELONEPHRITIS, NOS INFLAMMATION, SUPPUBATIVE INFLAMMATION, CHRONIC CALCIUM DEPOSIT	1 (2%) 23 (46%)	25 (63%)	1 (3%) 21 (57%) 1 (3%)

NUMBER OF ANIMALS WITH TISSUE EXAMINED MICROSCOPICALLY * NUMBER OF ANIMALS NECROPSIED

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TABLE C2 (CONTINUED)

	CONTROL (VEH) 01-P001	LON DOSE 01-9010	HIGH DOSE 01-P011
KIDNEY/PELVIS CALCIUM DEPOSIT	(50)	(40) 2 (5%)	(37)
AURINARY BLADDER INFLANNATION, NOS	(49)	(34)	(32) 1 (3%)
NDOCRINE SYSTEM			
#FITUITARY Anglectasis	(50) 1 (2≸)	(40)	(39)
#ADRENAL ANGIECTASIS	(50) 17 (34%)	(37) 6 (16%)	(33) 5 (15 %)
THYROID Hyperplasia, Nos Hyperplasta c-cree	(50) ⊃ (#€)	(33) 1 (3%)	(34)
HYPERPLASIA, C-CELL Hyperplasia, Pollicular-Cell	2 (4%)	3 [9%)	
EPRODUCTIVE SYSTEM			
MAMMARY GLAND	(50)	(50)	(59)
GALACTOCELE Hyperplasia, nos	1 (2%)	1 (2%) 1 (2%)	
#UT 5885	(49)	(34)	(31)
HYDROMETRA INFLAMMATION, NOS	(49) 9 (18%) 2 (4%)	3 (9%) 2 (6%)	2 (6%)
#UTERUS/ENDOMETRIUM Hyperplasia, cystic	(49) 3 (6%)	(34) 2 (6%)	(31) 1 (3%)
ACVARY CIST, NOS	(49) 2 (4%)	(34) 1 (3 %)	(31)
REFOUS SYSTEM			
80NB			
PECIAL SENSE ORGANS			
*EYE SYNECHIANOS	(50)	(50)	(50) 1 (2 %)

TABLE C2 (CONCLUDED)

	CONT ROL (VEH) 01-F001	LOW DOSE 01-E010	RIGH DOSE 01-P011
CATARBCT		4 (8%)	4 (8%)
*EYE/CORNEA	(50)	(50)	(50)
PIBBOSIS Cytoplasmic Vacuolization		1 (2%)	1 (2%)
	12.44 Jain		•
*BYE/RBTINA ATROPHY, NOS	(50)	(50) 4 (8%)	(50)
*EYE/CRYSTALLINE LENS	(50)	(50)	(50)
CALCIUM DEPOSIT	(30)	1 (2%)	(30)
*HARDERIAN GLAND	(50)	(50)	(50)
INFLAMMATION, NOS	- ·	1 (2%)	
NONE COY CAVITIES *PERITONEUM		(50)	(50)
INFLAMMATION, NOS	(29)	(30)	1 (2%)
*FERICARDIUM INFLAMMATION, NOS	(50)	(50)	(50) 4 (8%)
LL OTHER SYSTEMS			
NONE			
PECIAL MORPHOLOGY SUMMARY			
NO LESION REPORTED	1	1	4
NECROPSY PERF/NG HISTO PERFORME Auto/Necropsy/No histo	C	2	2

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APPENDIX D

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SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN MICE TREATED WITH DIOXATHION

 TABLE D1

 SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN MALE MICE TREATED WITH DIOXATHION

	CONTROL (VEH) 02-0011	LON DOSE 02-m012	HIGH DOSE 02-N013
NIMALS INITIALLY IN STUDY	20	50 1	50 1
NIMALS NECEOPSIED NIMALS EXAMINED HISTOPATHOLOGICA		49 49	49 49
NTEGUMENTARY SYSTEM			
*SKIN EPIDERMAL INCLUSION CYST	(20) 1 (5%)	(49) 1 (2%)	(49)
INFLAMMATION, NOS INFLAMMATION, CHRONIC		1 (2%)	2 (4%) 1 (2%)
+SUBCUT TISSUE Abscess, Nos	(20)	(49)	(49) 1 (2 %)
ESPIRATORY SYSTEM			
FLUNG INFLAMMATION, FOCAL	(16) 1 (6 %)	(17)	(19)
PNEUMONIA, CHEONIC MURINE		1 (6%)	
EMATOPOIETIC SYSTEM			
*SPLEEN INFLAMMATION, NOS	(16)	(11)	(15)
ANYLOIDOSIS	3 (19%)	1 (9%)	1 (7%) 2 (13%)
HYPERPLASIA, NOS Henatopolesís		1 (9%)	1 (7%) 2 (13%)
LYMPH NODE	1191	(13)	
INFLANNATION, NOS Henatopoiesis			1 (8%) 1 (8%)
<pre>#MESENTERIC L. NODE INFLAMMATION, NOS</pre>	(15)	(13) 2 (15%)	(12) 1 (8%)
HENATOPOIESIS		1 (8%)	· (0%)
IECOLATOBY SYSTEM			
#HEART ANYLOIDOSIS	(16)	(10)	(10) <u>1 (195)</u>

NUMBER OF ANIMALS WITH TISSUE EXAMINED MICROSCOPICALLY
 NUMBER OF ANIMALS NECROPSIED

** EXCLUDES PARTIALLY AUTOLYZED ANIMALS

TABLE DI (CONTINUED)

	CONTBOL (VEH) 02-n011	LOW DOSE 02-N012	HIGH DOSE 02-8013
GESTIVE SYSTEM			
#LIVE R	(17)	(49)	(49)
CYST, NOS			1 (2%)
INFLAMMATION, NOS		1 (2%)	1 (28)
NECROSIS, NOS Anyloidosis	2 (12%)	3 (6%)	1 (2%) 7 (14%)
METANORPHOSIS PATTY	2 (124)	, (c.,,	1 (2%)
HYPERPLASIA, NODULAR		3 (6%)	9 (18%)
HYPEBPLASTIC NODULE	1 (6%)	1 (25)	1 (2%)
HYPERPLASIA, FOCAL	1 (6\$)		2 (4%)
ANGIECTASIS	·		2 (47)
#FANCREAS	(16)	(11)	(10)
INFLAMMATICN, BOS		1 (9%)	
AMYLOIDOSIS	1 (6%)		
#LARGE INTESTINE	(15)	(10)	(10)
NEMATODIASIS	2 (13%)		()
*RECTUM	(20)	(49)	(49) 1 (2%)
INFLAMMATION, NOS			• (2.8)
RINART SYSTEM			
# KIDNEY	(19)	(16)	(18)
HIDRONEPHROSIS			1 (6%)
POLYCYSTIC KIDWEY Pyelonephritis, Nos		2 (13%)	1 (6%)
INPLANMATION, CHRONIC	8 (42%)	2 (13%)	2 (11%)
PERIABTERITIS	1 (5%)		
ANYLOIDOSIS	2 (11%)	3 (195)	3 (17%)
CALCIUM DEPOSIT	1 (5%)		
#URINARY BLADDER	(15)	(10)	(11)
INPLANMATION, CHRONIC	1 (7%)		
·			
NDOCRINE SYSTEM			
#ADRENAL	(16)	(10)	(10)
ANYLOIDOSIS	2 (13%)	• • •	• •
#PARATHIROID ANYLOIDOSIS	(6) 1 (17%)	(10)	(10)

* NUMBER OF ANIMALS WITH TISSUE EXAMINED MICROSCOPICALLY * NUMBER OF ANIMALS NECROPSIED

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D-4

TABLE DI (CONCLUDED)

*	CONTROL (VEH) 02-K011	109 DOSE 02-M012	HIGH DOSE D2-H013	
REPRODUCTIVE SYSTEM				
TESTIS Atrophy, Nos	(16) 1 (6%)	(12)	(10)	
*EPIDIDYMIS GRANGLOMA, SPERMATIC	(20) 1 (5%)	(49)	(49)	
NEFVOUS SYSTEM				
BONE				
SPECIAL SENSE ORGANS				
NONE				
MUSCULOSKELETAL SYSTEM				
NONE				
BODY CAVITIES				
NQMB				
ALL OTHER SYSTEMS				
NONE				
SPECIAL MORPHOLOGY SUMMARY				
NO LESION REPORTED Animal Missing/No Mecropsy Auto/Necropsy/No Misto	14 1	25 1	13 1	
* NUMBER OF ANIMALS WITH TISSUE EX * MUMBER OF ANIMALS NECROPSIED	ABINED MICBOSCOPIC	CALLY		

D-5

	CONTROL (VER) 02-F011	LOW DOSE 02-1014	HIGH DOSE 02-P015
MALS INITIALLY IN STUDY	20	50	50
IMALS MISSING IMALS NECROPSIED	1 19	50	1 49
ALS EXANINED HISTOPATHOLOGICALL	£ *. 18	50	48
GUMENTARY SYSTEM			
NB			
IRATORY SYSTEM			
NG	(18)	(15)	(12)
INFLAMMATION, FOCAL PNEDHOWIA, CHRONIC MURINE	1 (6%)	1 (7%)	
TOPOIETIC SYSTEM			
	(18)	(12)	(11)
IENATOPOIESIS		1 (8 %)	
LATORY SYSTEM			
E			
TIVE SYSTEM			
	(10)	(50)	(48)
INFLAMMATION, POCAL NECROSIS, POCAL		1 (2%)	1 (2%) 1 (2%)
ANGIECTAŠIS	1 (6%)	1 (2%)	2 (4%)
NCREAS ATROPHY, NOS	(17) 1 (6%)	(10)	(10)
GNACH		(11)	(10)
INFLAMMATION, NOS		1_1981	

TABLE D2 SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN FEMALE MICE TREATED WITH DIOXATHION

TABLE D2 (CONTINUED)

	CONTROL (VEH) 02-F011	LOW DOSE 02-F014	HIGH DOSE 02-7015
INARY SYSTEM			
KIDNEY LYMPHOCYTIC INPLANMATORY INFILTR	(18)	(10)	(11) 1 (9\$)
OCRINE SYSTEM			
ADRENAL HYPERPLASIA, NOS	(18)	(10) 1 (10 %)	(10)
RODUCTIVE SYSTEM			
NAMMARY GLAND Cyst, Nos	{19}	(50) 1 (2%)	(49)
UTERUS Hydrometra Inplanmation, Nos	(18)	(16) 2 (13%) 1 (6%)	(21) 2 (10 %)
INFLANMATION, SUPPURATIVE	1 (6%)		
ITERUS/ENDOMETRICH HYPERPLASIA, CYSTIC	(18) 2 (11%)	(16) 3 (19 \$)	(21) 8 (38 %)
VARY/OVIDUCI CYST, NOS INPLANHATION, NOS	(18)	(16)	(21) 1 (5%) 1 (5%)
VART CYST, NOS INFLARMATION, NOS	(18)	(13) 1 (8%) 1 (8%)	(12) 1 (8\$)
INFLAGRATION, SUPPORATIVE	2 (11%)	. (68)	
VOUS SYSTEM			
ON E			
CIAL SENSE ORGANS			
IYE HYPOPLASIA, NOS	(19)	1 (25)	(49)
CULOSKELBTAL SYSTEM			
IONE			

* NUMBER OF ANIMALS SECROPSIED

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TABLE D2 (CONCLUDED)

	CONTROL (VEH) 02-P011	LON DOSE 02-J014	HIGH DOSE 02-F015
BODY CAVITIES			
*PERITONEUM INFLAMMATION, NOS	(19) 1 (5%)		(49)
ALL OTHER SYSTEMS			
NONE			
SPECIAL MORPHOLOGY SUMMARY			
NO LESION REPORTED	9	34	3 f
ANIHAL MISSING/NO NECROPSY NECROPSY PERF/NO HISTO PERFORMED	1		1 1

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