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BIOASSAY OF

3,3'-IMINOBIS-1-PROPANOL DIMETHANESULFONATE (ESTER) HYDROCHLORIDE (IPD)

FOR POSSIBLE CARCINOGENICITY

CAS No. 3458-22-8

NCI-CG-TR-18

U.S. DEPARTMENT OF HEALTH, EDUCATION, AND WELFARE Public Health Service National Institutes of Health



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

U.S. DEPARTMENT OF HEALTH, EDUCATION, AND WELFARE Public Health Service National Institutes of Health

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Carcinogenesis Testing Program Division of Cancer Cause and Prevention National Cancer Institute National Institutes of Health

<u>CONTRIBUTORS</u>: This report presents the results of the bioassay of 3,3'-iminobis-1-propanol dimethanesulfonate (ester) hydrochloride [IPD] for possible carcinogenicity, conducted for the Carcinogenesis Testing Program, Division of Cancer Cause and Prevention, National Cancer Institute (NCI), Bethesda, Maryland. The bioassay was conducted by Southern Research Institute, Birmingham, Alabama, initially under direct contract to NCI and currently under a subcontract to Tracor Jitco, Inc., prime contractor for the NCI Carcinogenesis Testing Program.

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Animal pathology tables and survival tables were compiled at EG&G Mason Research Institute⁵. The statistical analyses were performed by Dr. J. R. Joiner⁶, using methods selected for the bioassay program by Dr. J. J. Gart⁷. Chemicals used in this bioassay were analyzed under the direction of Dr. W. J. Haggerty, Jr.⁸, and the results of the analyses were reviewed by Dr. S. S. $Olin^{6}$.

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SUMMARY

A bioassay of 3,3'-iminobis-l-propanol dimethanesulfonate (ester) hydrochloride [IPD] for possible carcinogenicity was conducted by administering the test chemical intraperitoneally to Sprague-Dawley rats and B6C3F1 mice.

The IPD was injected three times per week to groups of 35 animals, using doses of 12, 24, or 48 mg/kg for the rats, and 20 or 40 mg/kg for the mice. Rats at 12 mg/kg were treated for 52 weeks. Because of the toxicity of the chemical, administration of IPD for the group receiving 24 mg/kg was discontinued at week 34. Rats receiving 48 mg/kg were treated until all had died at week 23 (males) and week 27 (females). Both groups of mice were treated for 52 weeks. All survivors were killed after postadministration periods that varied among groups.

With rats, untreated and vehicle-control groups, each consisting of 10 males and 10 females, were started with the high- and middose groups and additional untreated and vehicle-control groups of the same size were started with the low-dose groups. With mice, untreated and vehicle-control groups each consisted of 15 males and 15 females.

The toxicity of IPD was associated with lower mean body weights and lower rates of survival of both the rats and mice. The shortened life spans, particularly in the rats, reduced the likelihood of the development of tumors.

In rats, peritonitis and fibrous adhesions, possibly, from direct irritation by the test chemical were observed in most treated rats at necropsy. Sarcoma, fibroma, or fibrosarcoma of the peritoneum occurred in two low-dose male, one mid-dose male, and three mid-dose female rats, but not in any control animals. Because of this low incidence, and because irritation by the test chemical may have been involved in the pathogenesis, these tumors may have been due to local effects of the chemical. In mice, lymphomas were observed at the following incidences (males: controls 0/14, low-dose 0/26, high-dose 3/21; females: controls 1/15, low-dose 2/29, high-dose 6/27). The Tarone test for life-table analysis of the probability of survival without lymphoma indicated a significant positive dose-related increase of lymphomas with a probability level of 0.011 for male mice and 0.003 for female mice.

Squamous-cell carcinoma was noted in the mice (low-dose males 6/26, high-dose females 2/27). Seven of these tumors were observed in subcutaneous tissue in the inguinal region near the sites of injection. Although not statistically significant, this tumor may be associated with administration of IPD.

Tumors of the peritoneum in rats and tumors in the subcutaneous tissue in mice may have been due to local effects related to administration of the test chemical. The lymphomas in mice, although marginally significant, were too few in number to clearly be related to dosing.

Conclusions from this study are limited by early deaths and toxicity, but the appearance of tumors in the peritoneum near the injection sites in both rats and mice indicate the carcinogenic potential of IPD.

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I. INTRODUCTION

3,3'-Iminobis-1-propanol dimethanesulfonate (ester) hydrochloride (CAS 3458-22-8; NCI C01547), hereinafter called IPD, was synthesized from bis(3-hydroxypropyl)amine and methanesulfonic acid anhydride (El-Merzabani and Sakurai, 1965). It was found to have antitumor activity against a number of experimental tumors that were naturally resistant to nitrogen mustard and has been used in Japan for the treatment of myelogenous leukemia (El-Merzabani and Sakurai, 1965; Hirano et al., 1972). IPD was selected for carcinogen bioassay as one agent in a series of anticancer drugs that are administered chronically in the treatment of human cancer.

II. MATERIALS AND METHODS

A. Chemical

The IPD was supplied by the Drug Development Branch, Division of Cancer Treatment (DCT), National Cancer Institute (NCI). It was purchased from Yoshitomi Pharmaceutical Industries, Limited, 35 Hiranomachi 3-chome, Higashi-ku, Osaka, Japan.

Analyses of each of two batches were provided through contracts of the Division of Cancer Treatment, NCI, and showed that the material consisted of > 99% of the designated chemical. The analytical methods included melting point, infrared and nuclear magnetic resonance spectroscopy, thin-layer chromatography (three solvent systems), and elemental analysis. No impurities were detected.

The IPD was stored at -20° C in the original glass container until used in this study.

B. Dosage Preparation

The IPD was prepared in phosphate-buffered saline as a fresh solution immediately prior to use. The actual mixing of the drug and the vehicle was performed in the animal laboratory, in a 10-ml Potter-Elvehjem tissue grinder with a Teflon pestle. The concentrations administered were 0.48%, 0.96%, or 1.92% (w/v) for

rats, and 0.2% or 0.4% (w/v) for mice. The test chemical solution or the vehicle was administered to the treated animals or vehicle controls intraperitoneally, using one needle for each injection group at a constant volume of 0.25 ml/100 g body weight for rats, or 1.0 ml/100 g for mice. Unused solutions of IPD were discarded each day of administration.

C. Animals

For the subchronic studies, female Sprague-Dawley rats and Swiss mice of each sex, obtained from the Charles River Breeding Laboratories, Inc., Wilmington, Massachusetts, through contracts of the Division of Cancer Treatment, NCI, were used. For the chronic studies, Sprague-Dawley rats and B6C3F1 mice, obtained from Charles River Breeding Laboratories, Inc., were used. On arrival at the laboratory, all animals were quarantined for an acclimation period (rats for 5 days, mice for 8 days), assigned to control or treated groups, and then earmarked for individual identification.

D. Animal Maintenance

All animals were housed in temperature- and humidity-controlled rooms. The temperature range was 20-24°C and the relative humidity was maintained at 40-60%. In addition to natural light, illumination was provided by fluorescent light for 9 hours per

day. Wayne[®] Lab Blox animal meal (Allied Mills, Inc., Chicago, Ill.) and water were supplied daily and were made available <u>ad</u> <u>libitum</u>.

Rats were housed five per cage and mice seven per cage in solid-bottom stainless steel cages (Hahn Roofing and Sheet Metal Co., Birmingham, Ala.). The bottoms of the rat cages were lined with Iso-Dri[®] hardwood chips (Carworth, Edison, N. J.), and cage tops were covered with disposable filter bonnets beginning at week 22; mouse cages were provided with Sterolit[®] clay bedding (Englehard Mineral and Chemical Co., New York, N. Y.). Bedding was replaced once per week; cages, water bottles, and feeders were sanitized at 82°C once per week; racks were cleaned once per week.

The rats and mice were housed in separate rooms. Control animals were housed with respective treated animals. Animals treated with IPD were maintained in the same rooms as animals of the same species being treated with the following chemicals:

RATS

Gavage Studies

```
cholesterol (p-(bis(2-chloroethyl)amino)phenyl)acetate
  (phenesterin) (CAS 3546-10-9)
estradiol bis((p-(bis(2-chloroethyl)amino)phenyl)acetate)
  (estradiol mustard) (CAS 22966-79-6)
```

Intraperitoneal Injection Studies

```
4'-(9-acridinylamino)methansulfon-m-aniside monohydrochloride
  (MAAM) (NSC 141549)
acronycine (CAS 7008-42-6)
5-azacytidine (CAS 320-67-2)
beta-2'-deoxy-6-thioguanosine monohydrate (beta-TGdR)
  (CAS 789-61-7)
1,4-butanediol dimethanesulfonate (busulfan) (CAS 55-98-1)
emetine dihydrochloride tetrahydrate (CAS 316-42-7)
(+)-4,4'-(1-methyl-1,2-ethanediyl)bis-2,6-piperazinedione
  (ICRF-159) (CAS 21416-87-5)
N, 3-bis(2-chloroethyl)tetrahydro-2H-1, 3, 2-oxazaphosphorin-2-
  amine-2-oxide (isophosphamide) (CAS 3778-73-2)
N-(2-chloroethyl)-N-(1-methyl-2-phenoxyethyl)benzylamine
  hydrochloride (phenoxybenzamine) (CAS 63-92-3)
N-(1-methylethyl)-4-((2-methylhydrazino)methyl)benzamide
  monohydrochloride (procarbazine) (CAS 366-70-1)
tris(l-aziridinyl)phosphine sulfide (thio-TEPA) (CAS 52-24-4)
```

MICE

Feed Studies

```
4-acetyl-N-((cyclohexylamino)carbonyl)benzenesulfonamide
  (acetohexamide) (CAS 968-81-0)
anthranilic acid (CAS 118-92-3)
1-buty1-3-(p-toly1sulfony1)urea (tolbutamide) (CAS 64-77-7)
4-chloro-N-((propylamino)carbonyl)benzenesulfonamide
  (chlorpropamide) (CAS 94-20-2)
5-(4-chlorophenyl)-6-ethyl-2,4-pyrimidinediamine
  (pyrimethamine) (CAS 58-14-0)
2,6-diamino-3-(phenylazo)pyridine hydrochloride
  (phenazopyridine hydrochloride) (CAS 136-40-3)
L-tryptophan (CAS 73-22-3)
N-9H-fluoren-2-ylacetamide (CAS 53-96-3)
N-(p-toluenesulfonyl)-N'-hexamethyleniminourea
  (tolazamide) (CAS 1156-19-0)
1-phenethylbiguanide hydrochloride (phenformin) (CAS 114-86-3)
pyrazinecarboxamide (pyrazinamide) (CAS 98-96-4)
4,4'-sulfonyldianiline (dapsone) (CAS 80-08-0)
4,4'-thiodianiline (CAS 139-65-1)
ethionamide (CAS 536-33-4)
```

Gavage Studies

cholesterol (p-(bis(2-chloroethyl)amino)phenyl)acetate
 (phenesterin) (CAS 3546-10-9)
estradiol bis((p-(bis(2-chloroethyl)amino)phenyl)acetate)
 (estradiol mustard) (CAS 22966-79-6)

Intraperitoneal Injection Studies

4'-(9-acridinylamino)methansulfon-m-aniside monohydrochloride (MAAM) (NSC 141549) acronycine (CAS 7008-42-6) 5-azacytidine (CAS 320-67-2) beta-2'-deoxy-6-thioguanosine monohydrate (beta-TGdR) (CAS 789-61-7) 1,4-butanediol dimethanesulfonate (busulfan) (CAS 55-98-1) emetine dihydrochloride tetrahydrate (CAS 316-42-7) (+)-4,4'-(1-methyl-1,2-ethanediyl)bis-2,6-piperazinedione (ICRF-159) (CAS 21416-87-5) N, 3-bis(2-chloroethyl)tetrahydro-2H-1, 3, 2-oxazaphosphorin-2amine-2-oxide (isophosphamide) (CAS 3778-73-2) N-(2-chloroethyl)-N-(1-methyl-2-phenoxyethyl)benzylamine hydrochloride (phenoxybenzamine) (CAS 63-92-3) N-(1-methylethyl)-4-((2-methylhydrazino)methyl)benzamide monohydrochloride (procarbazine) (CAS 366-70-1) tris(1-aziridiny1)phosphine sulfide (thio-TEPA) (CAS 52-24-4)

E. <u>Subchronic Studies</u>

Subchronic studies were conducted with female Sprague-Dawley rats and male and female Swiss mice to estimate the maximum tolerated doses of IPD, on the basis of which low and high concentrations (hereinafter referred to as "low doses" and "high doses") were determined for administration in the chronic studies. In these subchronic studies, IPD was administered intraperitoneally three times per week for 45 days to female rats and mice of each sex in twofold increasing concentrations, followed by 45 days of observation. The treated groups consisted of five animals each; the vehicle controls consisted of 10 animals each. The rats received 24, 48, 96, or 192 mg/kg/dose; the mice received 40, 80, 160, or 320 mg/kg/dose.

All rats receiving doses of 96 or 192 mg/kg died. The mortality rates for rats receiving 24 or 48 mg/kg were 20% and 40%, respectively; at these doses there were no differences in mean body weight between the 90-day survivors and the controls.

All mice receiving doses of 160 or 320 mg/kg died. The mortality rate for mice receiving 80 mg/kg was 90%. At 40 mg/kg, all mice survived, but there was a 30% depression in mean body weight, compared with the controls.

Low and high doses for the chronic study were set at 24 and 48 mg/kg, respectively, for rats and 20 and 40 mg/kg for mice.

F. Designs of Chronic Studies

The designs of the chronic studies are shown in tables 1 and 2. Originally, doses of 24 or 48 mg/kg were administered to groups of rats of each sex; however, toxicity resulted at the high dose, and low-dose groups at 12 mg/kg were started on study at week 56.

Table 1.

1. Design of Chronic Studies of IPD in Rats

Sex and	Initial	IPD	Time or	n Study
Test	No. of	Dose ^b	Treated	Untreated
Group	<u>Animals</u> a	(mg/kg)	(weeks)	(weeks)
)(-1-				
Male				
Low-Dose				
Untreated-Control	s ^c 10	0		89
Mid- and High-Dose		-		
Untreated-Control		0		89
Low-Dose				
Vehicle-Controls ^C	10	0 ^d	52	37
Mid- and High-Dose				
Vehicle-Control	10	0d		37
Low-Dose	35	12	52	3
Mid-Dose	35	24	34e	38f
High-Dose	35	48	23g	
Female				
Low-Dose				
Untreated-Control	s ^c 10	0		89
Mid- and High-Dose				
Untreated-Control	10	0		89
Low-Dose				
Vehicle-Controls ^C	10	0q	52	37
Mid- and High-Dose				
Vehicle-Control	10	0q	52	37
Low-Dose	35	12	52	28
Mid-Dose	35	24	34e	37 ^f
High-Dose	35	48	278	

^aHigh- and mid-dose males, with controls, were 35 days of age when placed on study; females with controls were 42 days of age. Lowdose males and females, with controls, were 55 days of age when placed on study.

^bIPD was administered in buffered saline by intraperitoneal injection three times per week at a volume of 0.25 ml/100 g body weight. Doses were based on individual weights. Table 1. Design of Chronic Studies of IPD in Rats

(continued)

- ^cTen controls were started initially with the mid- and high-dose groups, the other 10 were started concurrently with the low- dose group.
- ^dVehicle-control groups received only buffered saline solution, at the same volume as treated rats.
- ^eMid-dose male and female animals were treated only 34 weeks due to the toxicity of the chemical.

^fMid-dose male and female animals were observed only 38 and 37 weeks, respectively, due to the death of all animals.

^gHigh-dose male and female animals were treated only 23 and 27 weeks, respectively, due to the death of all animals.

Table 2.	Design	of	Chronic	Studies	of	IPD	in	Mice	
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Sex and	Initial	IPD	Time or	1 Study
Test	No. of	Dose ^b	Treated	Untreated
Group	<u>Animals</u> a	(mg/kg)	(weeks)	(weeks)
Male				
Untreated-Control	15	0		86
Vehicle-Control	15	0c	52	34
Low-Dose ^d	34	20	52	25
High-Dose	35	40	52	11
Female				
Untreated-Control	15	0		86
Vehicle-Control	15	0c	52	34
Low-Dose ^d	36	20	52	34
High-Dose	35	40	52	25

^aAll animals were 38 days of age when placed on study.

- ^bIPD was administered in buffered saline by intraperitoneal injection three times per week at a volume of 1 m1/100 g body weight. Doses were based on the mean weights of the animals in each cage.
- ^cVehicle-control groups received only buffered saline solution, at the same volume as treated mice.
- ^dThe low-dose group consisted of 34 males and 36 females instead of 35 animals of each sex, because of missexing during the initiation of the study.

G. Clinical and Pathologic Examinations

All animals were observed twice per day for signs of toxicity, and animals that were moribund were killed and necropsied. Rats (mid- and high-dose) and mice were weighed individually each week for the first 8 weeks, once every 2 weeks for the next 72 weeks, and once per month for the remainder of the study. Low-dose rats were weighed once every 2 weeks for 66 weeks and once per month thereafter. Palpation for masses was carried out at each weighing.

The pathologic evaluation consisted of gross and microscopic examination of major tissues, major organs, and all gross lesions from killed animals and from animals found dead. The following tissues were examined microscopically: skin, muscle, lungs and bronchi, trachea, bone and bone marrow, spleen, lymph nodes, thymus, heart, salivary gland, liver, gallbladder and bile duct (mice), pancreas, esophagus, stomach, small intestine, large intestine, kidney, urinary bladder, pituitary, adrenal, thyroid, parathyroid, mammary gland, prostate or uterus, testis or ovary, brain, and sensory organs. Peripheral blood smears were taken from each animal. Occasionally, additional tissues were also examined microscopically. The different tissues were preserved in 10% buffered formalin, embedded in paraffin, sectioned, and

stained with hematoxylin and eosin. Special staining techniques were utilized when indicated for more definitive diagnosis.

A few tissues from some animals were not examined, particularly from those animals that died early. Also, some animals were missing, cannibalized, or judged to be in such an advanced state of autolysis as to preclude histopathologic evaluation. Thus, the number of animals from which particular organs or tissues were examined microscopically varies, and does not necessarily represent the number of animals that were placed on study 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) for testing two groups for equality and Tarone's (1975) extensions of Cox's methods for testing for 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 is 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 appear-

ed 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) was used to compare the tumor incidence of a control group with 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, correction to ensure an overall significance level of 0.05 may be made. The Bonferroni inequality (Miller, 1966) 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), was also used. Under the assumption of a linear trend, this test determines if the slope of the dose-response curve is different from zero at the onetailed 0.05 level of significance. Unless otherwise noted, the direction of the significant trend is a positive dose relation-

ship. 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 an animal died naturally or was 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, two-tailed test) were also noted.

The approximate 95 percent confidence interval for the relative risk of each treated 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% 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 (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. RESULTS - RATS

A. Body Weights and Clinical Signs (Rats)

The mean body weights of rats of each sex treated with IPD were consistently lower than those of the vehicle controls (figure 1). Although the suppression of mean body weights was more marked in the males than in the females, the data indicate dose-related effects for both sexes. The growth rates of the untreated controls, not shown, were similar to those of the vehicle controls. Fluctuations in the growth curve may be due to mortality; as the size of the group diminishes, the mean body weight may be subject to wide variation. As the study progressed, all treated animals developed a poor physical condition; however, no other clinical signs were recorded.

B. Survival (Rats)

The Kaplan and Meier curves estimating the probabilities of survival for male and female rats receiving IPD at the doses used in this study, together with those of the controls, are shown in figure 2. The following table shows the numbers of weeks on study at which 50% and 100% mortality occurred in the treated and control rats.

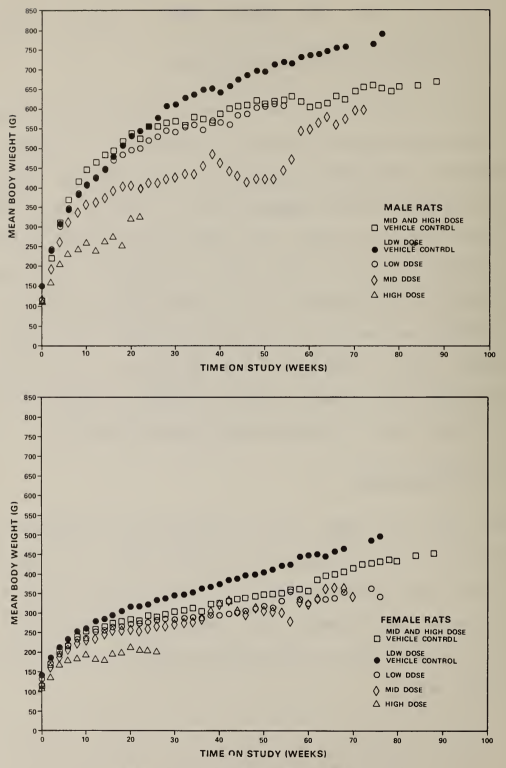
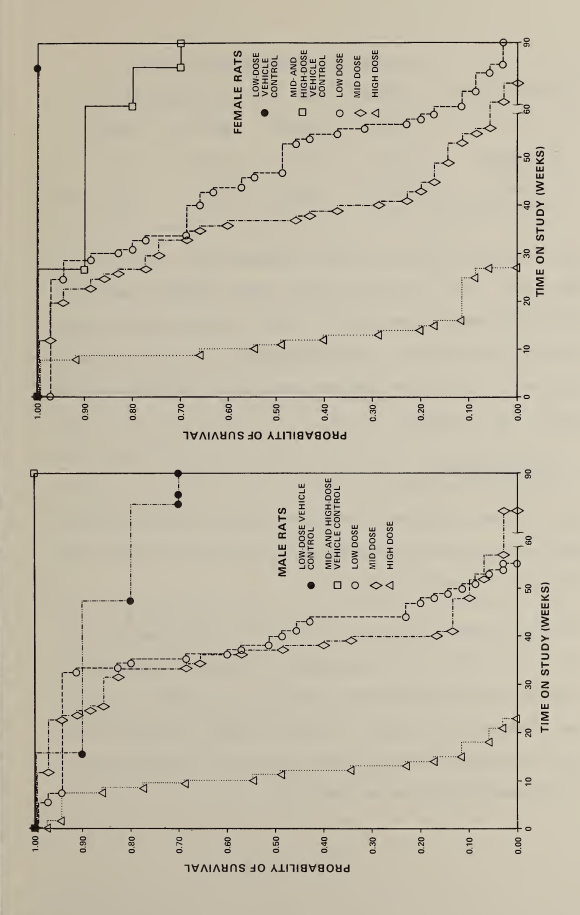


Figure 1. Growth Curves for Rats Treated with IPD



	Time	e to Death	(Weeks on	Study)	
Treated Group	50% Mo1	ctality	100% Mortality		
	Male	<u>Female</u>	Male	Female	
High-Dose	11	11	23	27	
Mid-Dose	37	37	72	71	
Low-Dose	38	47	55	80	
Vehicle-Control	89+	89+	89+	89+	

The data show that as the dose increased, the time to death decreased. The survival curves show highly significant (P < 0.001) dose-related positive trends in mortality, which are not linear (P < 0.001). The nonlinearity is due to the steep declines in survival of the treated groups compared with the vehicle-control groups.

C. Pathology (Rats)

Histopathologic findings on neoplasms in rats are summarized in Appendix A, tables Al-A4; findings on nonneoplastic lesions are summarized in Appendix C, tables Cl-C4.

A variety of tumors occurred in both the control (untreated and vehicle) and chemical-treated groups. Some types of neoplasms occurred only, or with greater frequency, in rats of treated groups compared with those of control groups. These lesions, however, are not uncommon in this strain of rat independent of the administration of any test chemical.

A small number of spindle-cell tumors occurred in the subcutaneous and peritoneal tissues with metastasis to the lungs and mediastinal lymph nodes. The subcutaneous tumors included one fibroma (1/18 [6%] in the combined groups of vehicle-control females) and two fibrosarcomas (1/20 [5%]) in the combined vehicle-control males and 1/33 [3%] in the low-dose females). The peritoneal tumors occurred in six rats: one fibroma (1/28)[4%] in the mid-dose males); two fibrosarcomas (1/28 [4%] in the mid-dose males and 1/31 [3%] in the mid-dose females); and four undifferentiated or pleomorphic spindle-cell sarcomas coded as sarcoma, NOS (not otherwise specified), (2/32 [6%] in the lowdose males and 2/31 [6%] in the mid-dose females). In one middose female rat, a peritoneal fibrosarcoma had metastasized to the mediastinal lymph nodes. In another mid-dose female rat, a peritoneal sarcoma of unspecified type had metastasized to the lungs. No sarcomas were found in untreated control animals.

Both the subcutaneous and the peritoneal spindle-cell tumors varied in the degree of differentiation and had various degrees of collagenous formation. The fibromas had well-differentiated fibroblasts with ample collagen, whereas the fibrosarcomas were more pleomorphic, more anaplastic, and more variable in the amount of collagen deposited. The poorly differentiated spindlecell tumors with little or no production of collagen were classi-

fied as sarcoma, NOS. One of the subcutaneous fibrosarcomas and two of the sarcoma, NOS, lesions were pleomorphic with formation of bizarre multinucleated giant cells. Two of the nonspecified sarcomas had extensively infiltrated the smooth muscle of the digestive tract. The confusing blend of neoplastic and nonneoplastic tissue made further classification of the sarcomas virtually impossible. The possibility of a leiomyosarcoma was not ruled out. Adenocarcinomas were observed in the large intestine in 2/28 (7%) of the mid-dose males. Both lesions were well differentiated, and one had large glandular spaces lined by columnar, cuboidal, and squamous epithelial cells. These spaces were filled by large amounts of mucin. Glands of this tumor had invaded the muscle layers.

In addition to the neoplastic lesions, a number of degenerative, proliferative, and inflammatory changes were also encountered in animals of the treated and control groups (Appendix C). For the most part, these nonneoplastic lesions are commonly seen in aged rats; however, the proliferative nonneoplastic lesions of the connective tissues lining the peritoneum were associated with treated groups. The incidences of these nonneoplastic lesions together with incidences of neoplastic peritoneal lesions, were as follows:

	Untreated <u>Control</u>	Vehicle <u>Control</u>	Low Dose	Mid <u>Dose</u>	High <u>Dose</u>
MALES					
Number of animals					
necropsied	(20)	(20)	(32)	(28)	(30)
Peritoneum					
Chronic Inflammation	0	0	30	23	7
Fibrous Adhesion, NOS	0	0	21	5	0
Fibroma	0	0	0	1	0
Fibrosarcoma	0	0	0	1	0
Sarcoma, NOS					
(spindle-cell)	0	0	2	0	0
FEMALES					
Number of animals					
necropsied	(20)	(18)	(33)	(31)	(31)
Peritoneum					
Chronic Inflammation	0	0	32	28	7
Fibrous Adhesion, NOS	0	0	30	8	0
Fibrosarcoma	0	0	0	1	0
Sarcoma, NOS					
(spindle-cell)	0	0	0	2	0

Chronic peritonitis and needle trauma may have had an important role in the pathogenesis of these peritoneal neoplasms. Needle injuries may also have been a factor in the induction of the subcutaneous fibromas and fibrosarcomas.

The small number of tumors observed may have been influenced by complications of severe chronic peritonitis and bone-marrow atrophy, with resulting decreased life span. With the reduced

period at risk, however, the tumors that were observed may have greater importance.

Injection of rats with IPD resulted in few tumors. The majority of the neoplastic lesions appeared unrelated to administration of the chemical. Adenocarcinomas of the large intestine and peritoneal sarcomas may have significance. The effectiveness of the carcinogenesis bioassay was reduced by an associated decrease in life span resulting from bone-marrow atrophy and severe chronic peritonitis. In the judgment of the pathologist, the results of this study failed to define the carcinogenic activity of IPD in Sprague-Dawley rats.

Histologic features of the tumors referred to above are presented in Appendix G.

D. Statistical Analyses of Results (Rats)

Tables El and E2 in Appendix E contain the statistical analyses of the incidences of those primary tumors that were observed in at least 5% of one or more than one treated group of either sex. No pooled-control groups are used in the statistical analyses, since there are no controls from other studies that are suitable for pooling. The untreated-control groups are not used in the analyses, since the conditions of the vehicle-control groups are more nearly comparable to the conditions of the treated groups.

Two separate analyses using the Cochran-Armitage test for linear trend are included in the tables. The first line in the tables shows the results of the analysis of the incidences of tumors in the four groups of animals (vehicle-control, low-dose, mid-dose, and high-dose groups), while the second line shows the results of the analysis of the incidences seen in the three groups of animals (vehicle-control, low-dose, and mid-dose groups).

In each sex, neither the Cochran-Armitage tests for positive dose-related trend in proportions for the incidence of tumors at any site (using either three doses or two doses) nor any of the Fisher exact tests for the comparison of the incidences of tumors in a treated group with that in the controls in the positive direction is significant at the 0.05 level. The survival was so poor in the treated groups that little reliance can be placed on the negative results of this test. Significant results in the negative direction are observed in the incidences of mammary and pituitary tumors, which may be accounted for by the early mortalities of the treated animals.

In all of the 95% confidence intervals shown in the tables, values of one or less than one are included, indicating the absence of positive statistically significant results. It should also be noted that in each of the intervals with an upper limit greater than one, there is the theoretical possibility of the

induction of that particular tumor by IPD, which could not be detected under the conditions of this test.

Time-adjusted analysis on the proportions of sarcoma, NOS, fibroma, or fibrosarcoma of the peritoneum in male rats is shown in table E2. The results of the Cochran-Armitage test are significant (P = 0.031 when control, low-, mid- and high-dose groups are used, and P = 0.045 when only control, low- and mid-dose groups are used), but the Fisher exact tests are not significant.

The time-adjusted analysis on the incidence of sarcoma, NOS, or fibrosarcoma of the peritoneum in female rats is shown in table E4. The results of the Cochran Armitage test are significant (P < 0.001), but departures from linear trend are observed (P < 0.001). These departures from a linear effect result from the higher proportion observed in the mid-dose group when compared with either the low-dosed or high-dosed groups. The Fisher exact test shows that the incidence in the mid-dose rats is significantly higher (P = 0.003) than that in the matched vehicle controls. These statistical test results suggest the possibility of dose association; however, it should be noted that the sample size used is very small, especially that in the mid-dose group, which is only 4. The zero incidence observed in the high-dose rats is probably due to the severe early mortality of this group of rats.

IV. RESULTS - MICE

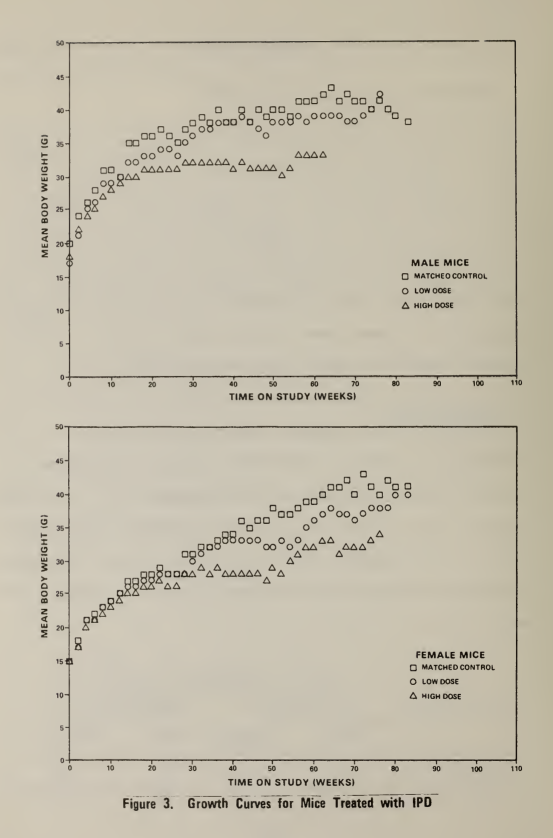
A. Body Weights and Clinical Signs (Mice)

The mean body weights of low- and high-dose male and female groups of mice were consistently lower in a generally doserelated manner than those of the vehicle controls (figure 3). The growth rates of the untreated controls, not shown, were similar to those of the vehicle controls. Fluctuations in the growth curve may be due to mortality; as the size of the group diminishes, the mean body weight may be subject to wide variation. No other clinical signs were recorded.

B. Survival (Mice)

The Kaplan and Meier curves estimating the probabilities of survival for male and female rats receiving IPD at the doses used in this study, together with those of the controls, are shown in figure 4. The following table shows the numbers of weeks that elapsed before 50% and 100% mortality occurred in the treated and control mice.

	Tim	e to Death (Weeks on St	tudy)
Treated Group	<u>50% Mor</u>	tality	100% Mo	rtality
	Male	Female	Male	Female
High-Dose	53	63	63	77
Low-Dose	63	65	77	86
Vehicle-Control	86+	86+	86+	86+



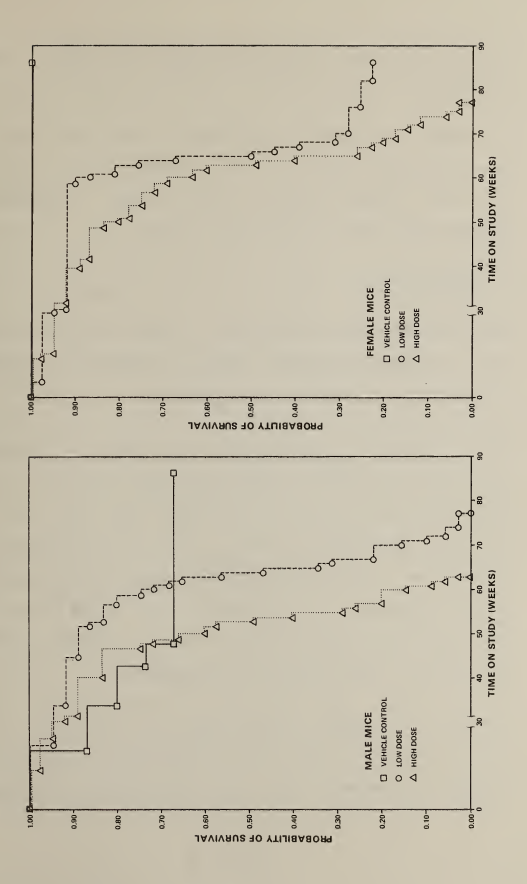


Figure 4. Survival Curves for Mice Treated with IPD

The data show that as the dose increased, the time to death decreased.

As in rats, the results of the Tarone test for mice showed a highly significant dose-related trend in mortality (P < 0.001 for each sex). Neither males nor females showed a significant departure from linear trend in relation to dose.

C. Pathology (Mice)

Histopathologic findings on neoplasms in mice are summarized in Appendix B, tables Bl and B2; findings on nonneoplastic lesions are summarized in Appendix D, tables Dl and D2.

With the exception of squamous-cell carcinomas and malignant lymphomas, the neoplasms listed in Appendix B appeared with approximately equal frequency in the treated and control mice, or appeared in insignificant numbers. Most of the squamous-cell carcinomas were located in the inguinal region and were believed to have originated from the preputial glands. Since the region where these arose was also near the sites of injection, local factors may have been involved in the pathogenesis.

Squamous-cell carcinomas in eight mice were recorded. The incidences were 6/26 (23%) in the low-dose males and 2/27 (7%) in the high-dose females. Seven of the squamous-cell carcinomas,

(in the six low-dose males and in one of the high-dose females) occurred in subcutaneous tissue in the inguinal area near the sites of injections. The presence of cyst walls in organization the tissues suggests that these tumors arose from the of The lesions were similar in epithelium of preputial glands. appearance, with abundant well-differentiated keratin-forming pearls in the center and poorly organized pleomorphic squamous cells in the walls. Some of the lesions were atypical with bizarre giant tumor cells. Several lesions had numerous mitotic figures. One lesion had invaded a perineural lymphatic. The eighth tumor was a squamous-cell carcinoma of the sebaceous glands in the ear canal of a female mouse. Neoplastic squamous replaced the sebaceous glands cells had and resulted in epithelial atypism and hypercellularity. (A tissue mass from another male mouse consisted cnly of keratin and probably represented a ninth squamous-cell carcinoma; however, this lesion was not included in the summary tables.)

Malignant lymphomas involved one or more organs in 13 mice. The majority of the affected mice were female (males: high-dose 3/21 [14%]; females: untreated controls 1/13 [8%], vehicle controls 1/15 [7%], low-dose 2/29 [7%], high-dose 6/27 [22%]). The organs with lymphoid tumors were kidney, liver, spleen, thymus, mesenteric and mediastinal lymph nodes, heart, lungs, stomach,

adrenal, ovary, uterus, urinary bladder, and bone marrow. One malignant lymphoma, grossly identified as an abdominal mass, probably originated in the mesenteric lymph node. Seven mice had disseminated malignant lymphomas involving multiple organs. 0f the 13 malignant lymphomas, nine were of the well-differentiated lymphocytic type; the remaining four included two undifferentiated lymphomas and two types unspecified. Lesions classified as a lymphocytic type had a uniform population of small cells with little cytoplasm and small round to oval nuclei having small inconspicuous nucleoli and coarse, dark chromatin. Cells of the lymphoblastic type were similar to cells of the lymphocytic type, with an increased amount of basophilic cytoplasm and larger, more varied nuclei having finer reticulated chromatin and more distinct nucleoli. The unspecified lymphomas had cellular distortion which prevented further classification.

> In addition to the neoplastic lesions, a number of degenerative, proliferative, and inflammatory changes were also encountered in animals of the treated and control groups (Appendix D). For the most part these nonneoplastic lesions were similar to those commonly observed in aged mice. The chronic fibrous peritonitis that was observed in rats given IPD was also present to a lesser extent in the mice. Two of these mice had peritoneal osseous metaplasia. Both the peritonitis and osseous metaplasia appeared

to be related to intraperitoneal injection, since these lesions also occurred in the vehicle-control group. In addition to the peritonitis, respiratory infections and bone-marrow injury may also have had a role in reducing the life spans of mice during this study. The extent to which reduced life spans influenced the number of tumors observed could not be determined.

In the judgment of the pathologist, the results of this study indicate that IPD given intraperitoneally to B6C3F1 mice was responsible for squamous-cell carcinomas of the inguinal region and an increased frequency of malignant lymphomas.

D. Statistical Analyses of Results (Mice)

Tables F1 and F2 in Appendix F contain the statistical analyses of the incidences of those primary tumors that were observed in 5% of one or more than one treated group of either sex. No pooled-control groups are used in the statistical analyses, since there are no controls from other studies that are suitable for pooling. The untreated-control groups are not used in the analyses, since the conditions of the vehicle controls are more nearly comparable to the conditions of the treated groups.

In male mice, the Cochran-Armitage test for positive dose-related trend in proportions for malignant lymphomas has a probability level of 0.045, with an incidence of 3/21 (14%) in the high-dose

group and none in the low-dose or vehicle-control groups. The female mice also show a larger proportion of this tumor in the high-dose group; however, due to the relatively small numbers in the groups, the results of the Fisher exact test do not show a significant difference between the high-dose and control groups The results of the Cochran-Armitage test on the in either sex. incidence in female mice are not significant. The time-adjusted analyses on the incidence of malignant lymphoma in both male and female mice are shown in tables F2 and F4, respectively. After adjustment, the male mice show an incidence of 3/20 (15%), 0/24, and 0/12 in the high-dose, low-dose, and vehicle-control groups, respectively. In female mice, the time-adjusted incidence becomes 1/15 (7%) in the vehicle-control group, 2/29 (7%) in the low-dose group, and 6/26 (23%) in the high-dose group. The results of the statistical tests using time-adjusted data are not significant. The life-table method was performed, using as an adjustment the week on study at which each malignant lymphoma was observed. Based on these data, figure 5 shows the Kaplan and Meier estimate of the probability of survival without the observation of a tumor. The Tarone test results indicated a significant positive dose-related trend for both males and females with a probability level of 0.011 for male mice and 0.003 for female mice. In neither sex is departure from linear trend indicated.

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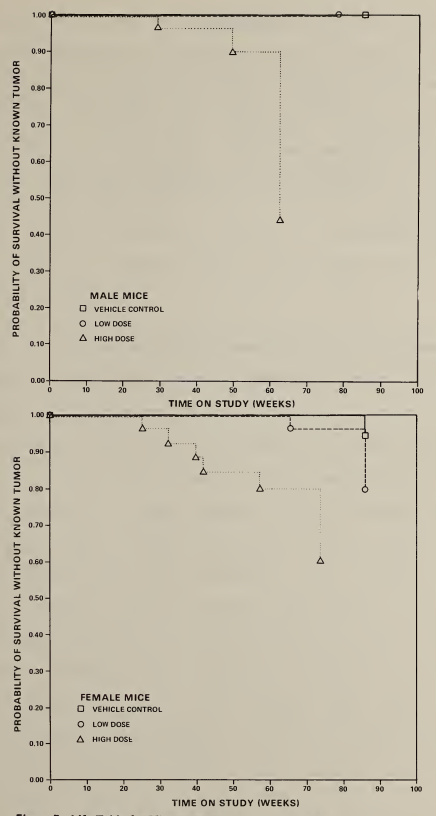


Figure 5. Life Table for Mice Treated with IPD: Malignant Lymphoma

In each of the 95% confidence intervals shown in the statistical tables, the value of one is included; this indicates the absence of positive statistically significant results. It should also be noted that each of the intervals has an upper limit greater than one, indicating the theoretical possibility of the induction of tumors by IPD, which could not be detected under the conditions of this test.

V. DISCUSSION

The doses of IPD used in the bioassay were toxic, as shown by the lowered mean body weights and rates of survival of the treated rats and mice. The shortened life spans, particularly in the rats, reduced the likelihood of the development of tumors. High-dose rats of both sexes had all died by week 27. It should also be noted that animals were treated for a maximum of only 52 weeks, which is a shorter period of time than used in other current bioassays.

In rats, peritonitis and fibrous adhesions of the peritoneum were observed in most of the treated animals at necropsy. Sarcoma, fibroma, or fibrosarcoma of the peritoneum occurred only in treated animals. Metastases to the lung or the lymph nodes occurred in two of the mid-dose females. Using time-adjusted analyses of the incidences of these tumors in mid- and low-dose animals surviving at least 44 weeks (male) and 52 weeks (female), there were significant dose-related trends (P = 0.045 in males, and P =0.001 in females), and the incidence in the mid-dose females was significantly higher (P = 0.003) than that in the vehicle controls. However, these significant results were based on tumors in only six animals (two low-dose males, one mid-dose male, and three mid-dose females). Because of this low incidence and because irritation by the test chemical and chronic

peritonitis may have been involved in the pathogenesis, these tumors may have been due to local effects of the injection of IPD. Therefore, these tumors are not considered as evidence of carcinogenicity of the test chemical.

Atrophy of the bone marrow was observed in all of the high-dose animals of both sexes that were necropsied and examined histopathologically.

In mice, lymphomas were observed at the following incidences (males: controls 0/14, low-dose 0/26, high-dose 3/21; females: controls 1/15, low-dose 2/29, high-dose 6/27). The results of the unadjusted and time adjusted analysis are not significant; however, the Tarone test for life-table analysis of the probability of survival without lymphoma indicated a significant positive dose-related increase of lymphomas with a probability level of 0.011 for male mice and 0.003 for female mice. These significant results are based on tumors in only three male and eight treated female mice.

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Squamous-cell carcinomas occurred in 6/26 low-dose male and 2/27 high-dose female mice, but in no other group. Seven of these carcinomas occurred in subcutaneous tissue in the inguinal area near the sites of injection and probably arose from the epithelium of the preputial glands; the eighth, in a female, was in the

sebaceous glands of the ear canal. None of the statistical tests for these tumors was significant; however, since most tumors arose near the sites of injection, they may have been related to the repeated intraperitoneal injections of the test chemical, irritation by the test chemical, or both.

IPD is an antitumor agent that has immunosuppressive properties, as shown by reduction of leukocytes in Donryu rats (Tsukagoshi and Sakurai, 1970), reduction of spleen and bone-marrow cells in CDF1 mice (Vadlamudi et al., 1971a), and suppression of hemagglutinin synthesis in CDFl mice (Vadlamudi et al., 1971b). These immunosuppressive properties may, in turn, be responsible for the apparent increase in tumors of the lymphoid system in mice, which was observed in the present bioassay. Results from a pulmonary tumor test system have shown that intraperitoneal injections of IPD into strain A mice at doses of 46 mg/kg three times per week for 8 weeks induced statistically significant numbers of pulmonary tumors (Stoner et al., 1973).

Tumors of the peritoneum in rats and tumors in the subcutaneous tissue in mice may have been due to local effects related to administration of the test chemical. The lymphomas in mice, although statistically significant, were too few in number to be clearly related to dosing.

Conclusions from this study are limited by early deaths and toxicity, but the appearance of tumors in the peritoneum near the injection sites in both rats and mice indicate the carcinogenic potential of IPD.

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APPENDIX A

SUMMARY OF THE INCIDENCE OF NEOPLASMS IN RATS GIVEN INTRAPERITONEAL INJECTIONS

OF IPD

TABLE A1

SUMMARY OF THE INCIDENCE OF NEOPLASMS IN MALE RATS **GIVEN INTRAPERITONEAL INJECTIONS OF IPD (CONTROL GROUPS)**

	MID-ANOHIGH-OOSE UNTREATED CONTROL	LOW-DOSE Untreated Control	MID- AND HIGH-DOSE Vehicle Control	LDW-DOSE Vehicle Control
ANIMALS INITIALLY IN STUDY ANIMALS NECROPSIED ANIMALS EXAMINEC HISTOPATHOLOGICAI	10 10 10 LY 10	10 10 10	10 10 10	10 10 10
INTEGUMENTARÝ SYSTEM				
*SKIN KERATOACANTHOMA	(10)	(10)	(10) 1 (10%)	(10)
*SUÉCUT TISSUF FIBROSA FCCMA	(10)	(10)	(10)	(10) 1 (10%
RESPIRATORY SYSTEM				
HEMATOPOIETIC SYSTEM				
NONE				
CIRCULATCRY SYSTEM				
NONE				
DIGESTIVE SYSTEM				
NONE				
URINARY SYSTEM				
NONE				
ENLOCRINE SYSTEM				

NUMBER OF ANIMALS WITH TISSUE EXAMINED MICROSCCPICALLY
 NUMBER OF ANIMAIS NECROPSIED

TABLE A1 CONTROL	MALE RATS:	NEOPLASMS	(CONTINUED)

	MIO AND HIGH OOSE UNTREATED CONTROL	LOW-OOSE UNTREATEO CONTROL	MIO- AND HIGH-DOSE Vehicle Control	LOW-OOSE Vehicle Control
REPPODUCTIVE SYSTEM				
*MAMMARY GLANI ADENGCARCINCHA, NOS FIBROADENCHA	(10)	(10)	(10)	(10) 1 (10% 1 (10%
TESTIS INTERSTITI⊁L-CELL TUMOR	(9)	(10) 1 (10 %)	(10)	(10)
NERVOUS SYSTEM				
NONE				
SPECIAL SENSE CRGANS				
*FAR CANAL SQUAMOUS CFIL CARCINOMA	(10)	(10) 1 (10%)	(10)	(10)
USCULOSKELETAI SYSTEM				
NONE				
OLY CAVITIES				
NONE				
ALL OTHER SYSTEMS				
NONE				
ANIMAL DISPOSITION SUMMARY				
ANIMALS INITIALLY IN STUDY NATURAL DEATHƏ MORIBUND SACPIFICE SCHEDULED SACRIFICE	10 1	10	10	10 1 2
ACCIDENTAIIY KILLED TERMINAL SACRIFICE ANIMAL MISSING	9	10	10	7
INCLUDES AUTCLYZED ANIMALS				

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

TABLE A1 CONTROL MALE RATS: NEOPLASMS (CONTINUED)

		UNTREATEO	MIO- ANO HIGH-OOSE Vehicle Control	VEHICLE
TUMOR SUMMARY				
TOTAL ANIMAIS WITH PRIMARY TUMORS TOTAL PEIMARY TUMORS	* 1	1 2	2 2	3 3
TOTAL ANIMAIS WITH BENIGN TUMORS TOTAL EENIGN TUMORS		1	1	1
TOTAL ANIMALS WITH MALIGNANT TUMO TOTAL MALIGNANT TUMORS	rs 1 1	1	1	2 2
TOTAL ANIMALS WITH SECONDARY TUMO TOTAL SECONDARY TUMORS	RS#			
TOTAL ANIMALS WITH TUMORS UNCERTA Benign or malignant Total uncertain tumors	IN -			
TOTAL ANIMAIS WITH TUMORS UNCERTA Primary or metastatic Total unceftain tumors	IN -			
* FRIMARY TUMORS: ALL TUMORS EXCEPT \$ SECONDARY TUMORS: METASTATIC TUMO			ADJACENI ORGAN	

TABLE A2

SUMMARY OF THE INCIDENCE OF NEOPLASMS IN MALE RATS GIVEN INTRAPERITONEAL INJECTIONS OF IPD (TREATED GROUPS)

LOW DOSE		
35 32 32	35 28 28	35 30 30
(32)	(28) 1 (4 %)	(24)
(32)	(28) 1 (4 %)	(24)
	35 32 32 (32)	35 32 32 32 (32) (28) 1 (4%) (32) (28)

* NUMBER OF ANIMALS NECROPSIED

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TABLE A2 TREATED MALE RATS: NEOPLASMS (CONTINUED)

	LOW DOSE	MIO OOSE	HIGH DOSE
REPRODUCTIVE SYSTEM			
NONE			
NER VOUS SYSTEM			
# ER A IN ASTROCYTCM P	(29) 1 (3%)	(28)	(22) 1 (5%
SPECIAL SENSE CEGANS			
*EAR CANAL KERATOACANTHOMA	(32)	(28) 1 (4%)	(30)
NUSCULOSKELETAI SYSTEM			
NONE			
ECDY CAVITIES			
*PERITCNEUM	(32)	(28)	(30)
SARCOMA, NCS FIBROMA	2 (6%)	1 (4%)	
FI BROS A RC CM A		1 (4%)	
ALL OTHER SYSTEMS			
NONE			
PNIMAL DISPOSITION SUMMARY			
ANIMALS INITIALLY IN STUDY	35	35	35
NATURAL DEATHD MORIBUNE SACRIFICE	17 18	16 19	22 13
SCHETULEI SACRIFICE			
ACCIDENTAILY KILLED TERMINAL SPCRIFICE			
ANIMAL MISSING			
a_INCLUDES_AUICLYZED_ANIMALS			

* NUMBER OF ANIMALS NECROPSIED

TABLE A2 TREATED MALE RATS: NEOPLASMS (CONTINUED)

	LDW DOSE	MID DDSE	HIGH DOSE
TEMOR SUMMARY			
TOTAL ANIMAIS WITH PRIMARY TUMORS* TOTAL PRIMARY TUMOPS	3	4 5	1
TOTAL ANIMAIS WITH BENIGN TUMORS TOTAL FENIGN TUMORS		2 2	
TOTAL ANIMALS WITH MALIGNANT TUMORS TOTAL MALIGNANT TUMORS	3 3	3 3	1
TOTAL ANIMALS WITH SECONDARY TUMORS* TOTAL SECCNEARY TUMORS	ł		
TOTAL ANIMALS WITH TUMORS UNCERTAIN- BENIGN OR MAIIGNANT TOTAL UNCEFTAIN TUMORS			
TOTAL ANIMAIS WITH TUMORS UNCERTAIN- FRIMARY OF MFTASTATIC TOTAL UNCEFTAIN TUMORS			
 FRIMARY TUMORS: ALL TUMORS EXCEPT SE SECONDARY TUMORS: METASTATIC TUMORS 			JACENI ORGAN

TABLE A3

SUMMARY OF THE INCIDENCE OF NEOPLASMS IN FEMALE RATS **GIVEN INTRAPERITONEAL INJECTIONS OF IPD (CONTROL GROUPS)**

	MIO- ANO HIGH-DOSE UNTREATEO CONTROL	LOW-OOSE UNTREATEO CONTROL	MIO- AND HIGH-OOSE VEHICLE CONTROL	LOW-DOSE Vehicle Control
ANIMALS INITIALLY IN STUDY ANIMALS NECECESIEC ANIMALS EXAMINED HISTOPATHOLOGICAI	10 10 1.y 10	10 10 10	 10 8 8	10 10 10
INTEGUMENTARY SYSTEM				
*SUBCUT TISSUF FIBROMA	(10)	(10)	(8) 1 (13%)	(10)
RESPIRATORY SYSTEM				
NONE				
EEMATOPOIFTIC SYSTEM				
NONE				
CIRCULATCRY SYSTEM				
NONE				
CIGESTIVE SYSTEM				
NONE				
URINARY SYSTEM				
NONE				
ENDOCRINE SYSTEM				
*PITUITARY CHRONOPHOEE ACENOMA	(10)	(6) 1 (17 %	(8)) 3 (38%)	(10) 5 (50
#ADRENAL CORTICAL ADENCHA	(10)	(10)	(8) 1 (13%)	(10)

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

TABLE A3 CONTROL FEMALE RATS: NEOPLASMS (CONTINUED)

	MID- AND HIGH-DDSE UNTREATED CDNTRDL	LDW-DDSE UNTREATED CDNTRDL	MID- AND HIGH-DOSE Vehicle Control	LOW-DOSE VEHICLE CDNTRDL
THYROID C-CELL CARCINCMA	(5) 1 (20 %)	(7)	(8)	(10)
REPRODUCTIVE SYSTEM				
*MAMMARY GLANI ADENCHA, NOS ADENCCARCINCMA, NOS PIBROADENCMA		(1D) 3 (3D%)	(8) 3 (38%)	(10) 1 (1DX 1 (1DX 2 (2DX
# OTER US EN DOMETRIAL STROMAL POLYP	(10) 1 (1D%)	(1D)	(8) 1 (13%)	(10)
ERVOUS SYSTER				
#ERAIN ASTROCYTOMA	(1D)	(9)	(8) 1 (13 %)	(1D)
PECIAL SENSE CEGANS				
NONE				
USCULOSKELETAL SYSTEM				
NONE				
COLA CAVITIES				
*MESENTERY LIPOMA	(1D) 1 (10%)	(1D)	(8)	(1D)
ALL OTHER SYSTEMS				
NONE				

TABLE A3 CONTROL FEMALE RATS: NEOPLASMS (CONTINUED)

٨		LDW-DDSE Untreated Control	MID- AND HIGH-DDSE Vehicle Control	LDW-DDSE Vehicle Control
NIMAL DISPOSITION SUMMARY				
ANIMALS INITIALLY IN STUDY	10	10	10	10
NATUFAL DEATH@	1		3	
MORIBUND SACRIFICE	1	1		
SCHELULEL SACRIFICE			1	
ACCIDENTALLY KILLED		_		
TERMINAL SACRIFICE	8	9	6	10
ANIMAL MISSING				
I INCLUDES AUTCLYZED ANIMALS				
LUNOR SUMMARY				
TOTAL ANIMALS WITH PRIMARY TUMORS	5 * 5	4	6	6
TOTAL PRIMARY TUMORS	6	4	10	9
TOTAL ANIMALS WITH BENIGN TUMORS	u	ų	6	6
TOTAL BENIGN TUMORS	4	4	9	8
TOTAL ANIMALS WITH MALIGNANT TUNC	DRS 2		1	1
TOTAL MALIGNANT TUMORS	2		1	1
TOTAL ANIMALS WITH SECONDARY TUMO	RS#			
TOTAL SECCNEARY TUMORS				
TOTAL ANIMALS WITH TUMORS UNCERTA	TN -			
EENIGN OR MAIIGNANT				
TOTAL UNCEFTAIN TUMORS				
TOTAL ANIMAIS WITH TUMORS UNCERTA	IN-			
FRIMARY OR METASTATIC				
TOTAL UNCEFTAIN TUMORS				
PRIMARY TUMORS: ALL TUMORS EXCEPT	SECONDARY TUMO	RS		
SECONDARY TUMORS: METASTATIC TUMO			ADTROPEST OPCAN	

TABLE A4

SUMMARY OF THE INCIDENCE OF NEOPLASMS IN FEMALE RATS GIVEN INTRAPERITONEAL INJECTIONS OF IPD (TREATED GROUPS)

	LOW OOSE	MIO ODSE	HIGH DOSE
NNIMALS INITIAILY IN STUEY Animals necfofsied Animals examined histopathologically	35 33 33	35 31 31	35 31 31
INTEGUMENTARY SYSTEM			
*SUBCUT TISSUE PIBROSARCCYA	(33) 1 (3%)	(31)	(3 1)
FESPIRATORY SYSTEM			
<pre>IDNG SARCCNA, NCS, METASTATIC</pre>	(33)	(31) 1 (3%)	(30)
HEMATOPOIPTIC SYSTEM			
<pre>MEDIASTINAL L.NODE PIBROSARCCKA, MITASTATIC</pre>	(2)	(9) 1 (11%)	(12)
CIRCULATORY SYSTEM			
DIGESTIVE SYSTEM			
*LIVER HEPATOCELLULAR ADENOMA	(33)	(31) 1 (3%)	(31)
DRINARY SYSTEM			
NONE			
ENLOCRINE SYSTEM			
*PITUITARY CHROMOPHOFF_ACTNOMA	(29) 2 (7%)	(27)	(22)

* NUMBER OF ANIMALS NECROPSIEC

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TABLE A4 TREATED FEMALE RATS: NEOPLASMS (CONTINUED)

		MID DOSE	HIGH DOSI
REPRODUCTIVE SYSTEM			
*MAMMARY GLANI Adencha, NCS Adenocarcincma, Nos Fibroadenoma	(33) 2 (6%) 1 (3%) 4 (12%)	(31)	(31)
HE MANGIOMA		1 (3%)	
ERVOUS SYSTEM			
NONE			
PECIAL SENSE CEGANS			
NONE			
USCULOSKELETAI SYSTEM			
NONE			
CEY CAVITIES			
*PERITCNEUM SARCOMA, NCS FIBRCSARCCMA	(33)	(31) 2 (6%) 1 (3%)	(31)
IL OTHER SYSTEMS			
NONE			
NIMAL DISPOSITION SUMMARY			
ANIMALS INITIALLY IN STUDY Natural cfathð	35 8	35 15	35 21
MORIBUNE SACRIFICE SCHEEULEE SACRIFICF	26	20	14
ACCIDENTAILY KILLED TERMINAL SPCRIFICE ANIMAL MISSING	1		
INCLUDES AUTCLYZED ANIMALS			

* NUMBER OF ANIMALS WITH TISSUE * NUMBER OF ANIMALS NECROPSIED

TABLE A4 TREATED FEMALE RATS: NEOPLASMS (CONTINUED)

	LOW DOSE	MID DOSE	HIGH DOSE
TCHOR SUMMARY			
TOTAL ANIMALS WITH PRIMARY TUMORS*	8	3	
TOTAL PRIMARY TUMORS	10	້5	
TOTAL ANIMALS WITH BENIGN TUMORS	7	2	
TOTAL EFNICN TUMORS	8	2	
TOTAL ANIMALS WITH MALIGNANT TUMORS	2	3	
TOTAL MALIGNANT TUMORS	2	3	
TOTAL ANIMALS WITH SECONDARY TUMORS TOTAL SECONDARY TUMORS	•	2	
		-	
TOTAL ANIMALS WITH TUMORS UNCERTAIN BENIGN OR MAIIGNANT	-		
TOTAL UNCEFTAIN TUMORS			
TOTAL ANIMALS WITH TUMORS UNCERTAIN	-		
FRIMARY OF METASTATIC Total Unceftain Tumors			
* PRIMARY TUMORS: ALL TUMORS EXCEPT SI		D.C.	
# SECONDARY TUMORS: METASTATIC TUMORS			DJACENT ORGAN

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APPENDIX B

SUMMARY OF THE INCIDENCE OF NEOPLASMS IN MICE GIVEN INTRAPERITONEAL INJECTIONS

OF IPD

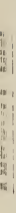


TABLE B1

SUMMARY OF THE INCIDENCE OF NEOPLASMS IN MALE MICE **GIVEN INTRAPERITONEAL INJECTIONS OF IPD**

		VEHICLE Control	LOW DOSE	HIGH OOSE
ANIMALS INITIALLY IN STUDY ANIMALS MISSING	15	15	34	35
ANIMALS NECFORSIFC ANIMALS EXAMINED HISTOPATHOLOGICALLY	14 14	14 14	26 26	21 21
INTEGUMENTAFY SYSTEM				
*SUBCUT TISSUE SQUAMOUS CEIL CARCINOMA	(14)	(14)	(26) 6 (23 %)	(21)
RESPIRATORY SYSTEM				
#LUNG CARCINOMA,NOS		(14)	1 (4%)	(22)
FEMATOPCIETIC SYSTEM				
*MUITIPIE CEGANS MALIG.LYMPHOMA, LYMPHOCYTIC TYPE		(14)	(26)	(21) 1 (5%)
#MEDIASTINAL I.NODE CARCINOMA, NOS, METASTATIC		(2)	(1) 1 (100%)	(3)
*THYMUS MALIG.LYMFHOMA, LYMPHOCYTIC TYPE		(14)	(22)	(20) 2 (10%)
CIRCULATORY SYSTEM				
NONE				
<i>LICESTIVE SYSTEM</i>				
#STOMACH SQUAMOUS CELL PAPILLOMA	(14)	(14)	(23)	(22) 1 (5 %)
URINARY SYSTEM				
NONE				

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

TABLE B1 MALE MICE: NEOPLASMS (CONTINUED)

	UNTREATED CONTROL	VEHICLE Control	LDW DOSE	HIGH DOSE
ENDOCRINE SYSTEM				
#PANCREATIC ISLETS ISLET-CELI ∤CENOMA	(13)	(14) 1 (7%)	(24)	(21)
REPRODUCTIVE SYSTEM				
NONE				
NERVOUS SYSTEM				
NONE				
SFECIAL SENSE CEGANS				
NO N E				
EUSCULOSKELETAI SYSTEM				
NON E				
ECEY CAVITIES				
NO N E				
ALL OTHER SYSTEMS				
NON E				
INIMAL DISPOSITION SUMMARY				
ANIMALS INITIALLY IN STUDY NATURAL DEATHƏ MORIBUNI SACRIFICF SCHEIDLFI SACPIFICF	15 3	15 5	34 20 13	35 26 9
ACCIDENTAILY KILLED TERMINAL SACRIPICF ANIMAL MISSING	12	10	1	
a INCLUDES ACTOLYZED ANIMALS				
<pre>* NUMBER OF ANIMALS WITH TISSUE F * NUMBER OF ANIMAIS NECROPSIEC</pre>	EXAMINED MICROSCO	PICALLY		

TABLE B1 MALE MICE: NEOPLASMS (CONTINUED)

	UNTREATED CONTRDL		LDW DDSE	HIGH DDSE
TUMOR SUNMARY				
TOTAL ANIMALS WITH PRIMARY TUMORS* TOTAL PRIMARY TUMORS		1	6 7	4 4
TOTAL ANIMALS WITH BENIGN TUMORS TOTAL BENIGN TUMORS		1		1
TOTAL ANIMALS WITH MALIGNANT TUMORS TOTAL MALIGNANT TUMORS			6 7	3 3
TOTAL ANIMAIS WITH SECONDARY TUMORS TOTAL SECCNDARY TUMORS	#		1 1	
TOTAL ANIMALS WITH TUMORS UNCERTAIN BENIGN OR MAIIGNANT TOTAL UNCERTAIN TUMORS	-			
TOTAL ANIMAIS WITH TUMORS UNCERTAIN PRIMARY OF MEIASTATIC TOTAL UNCEFTAIN TUMORS	-			
* FRIMARY TUMORS: ALL TUMORS EXCEPT S SECONDARY TUMORS: METASTATIC TUMORS			ADJACENI ORGAN	

TABLE B2

SUMMARY OF THE INCIDENCE OF NEOPLASMS IN FEMALE MICE **GIVEN INTRAPERITONEAL INJECTIONS OF IPD**

	UNTREATEO CONTROL		LOW DOSE	
IMALS INITIALLY IN STUDY	 15 1	15	36	35
NIMALS MISSING NIMALS NECRCESIEC NIMALS EXAMINED HISTOPATHOLOGICAIL	13	15 15	29 25	27 26
TEGUMENTAFY SYSTEM				
SUBCUT TISSUF Squamods CFLI Carcinoma FIBRCSAFCCPA	(13)	(15)	(29)	(27) 1 (49 1 (49
SPIRATCRY SYSTEM				
	(13)	(15)	(29)	(24)
ALVEOLAR/FRENCHIOLAR ADENOMA ALVEOLAR/EFONCHIOLAR CARCINOMA ADENCSQUAECUS CARCINOMA, MFTAST		1 (7%)		2 (8)
SHATOPOIETIC SYSTEM				
MULTIPIE CRGANS MALIGNANT IYMPHOMA, NOS MALIGLYMEHCMA, UNDIPPER-TYPE MALIGLYMEHCMA, LYMPHOCYTIC TYP	(13) E 1 (8%)	(15)	(29)	(27) 2 (7) 2 (7) 1 (4)
AEDOMINAL CAVITY MALIG.LYMFHONA, LYMPHOCYTIC TYP	(13) E	(15)	(29) 1 (3%)	(27)
NLIVER MALIG.LYMPEOMA, LYMPHOCYTIC TYP	(13) E	(15) 1 (7%)	(29)	(24)
IKIDNEY Malig.lymphcma, lymphocytic typ	(13) B	(15)	(28) 1 (4%)	(24)
	(13) E	(15)	(25)	(19) 1 (59

\$ NUMBER OF ANIMALS WITH TISSUE EXAMINED MICROSCCPICALLY * NUMBER OF ANIMAIS NECROPSIED

TABLE B2 FEMALE MICE: NEOPLASMS (CONTINUED)

	UNTREATED Control	VEHICLE Control	LOW DOSE	HIGH DOSE
LIGESTIVE SYSTEM				
NCNE				
ORINARY SYSTEM				
NCNE				
ENDOCRINE SYSTEM				
NCNE				
REPRODUCTIVE SYSTEM				
*MAMMARY GLAND ADENCCARCINCHA, NOS	(13)	(15)	(29)	(27) 1 (4%
OVARY ADENOMA, NCS PAPILLARY ADENOMA	(8)	(14)	(25)	(18) 1 (6% 1 (6%
CYSTADENCRA, NOS Tubular Acencma		1 (7%)	9 (36%) 1 (4%)	
NERVOUS SYSTEM				
NONE				
SPECIAL SENSE CEGANS				
*EAR CANAL Squamous cele carcinoma	(13)	(15)	(29)	(27) 1 (4 %
MUSCULOSKELITAI SYSTEM				
NONE				
ECTY CAVITIES				
*ABDOMINAL CAVITY CARCINONA, NOS	(13)	(15)	(29)	(27)

* NUMBER OF ANIMALS NFCROPSIEL

TABLE B2 FEMALE MICE: NEOPLASMS (CONTINUED)

	UNTREATED CONTRDL	VEHICLE CONTROL	LOW DDSE	HIGH DOS
MUCINOUS ADENOCARCINOMA			1 (3%)	
* PELVIS MESOTHELICFA, MALIGNANT	(13)	(15)	(25) 1 (3%)	(27)
LL OTHER SYSTEMS				
NO N E				
NIMAL DISPOSITION SUBMARY				
ANIMALS INITIALLY IN STUDY NATURAL CEATH@	15	15	36 14	35 24
MORIBUNE SACRIFICE	1		14	11
SCHETULEI SACRIFICE ACCIDENTAIIY KILLED			'	
TERMINAL SFCRIFICE Animal Missing	12 1	15	7	
INCLUDES AUTCLYZED ANIMALS				
UMOR SUMMARY				
TOTAL ANIMALS WITH PRIMARY TUMORS* TOTAL PRIMARY TUMORS	1	3 3	13 16	14 15
TOTAL ANIMALS WITH BENIGN TUMORS TOTAL BENIGN TUMORS		1	1 0 10	4 4
TOTAL ANIMALS WITH MALIGNANT TUMOR TOTAL MALIGNANT TUMORS	ts 1 1	2 2	5 6	10 11
TOTAL ANIMAIS WITH SECONDARY TUMOR TOTAL SECCNDARY TUMORS	₹S#			1
TOTAL ANIMALS WITH TUMORS UNCERTAD EENIGN OR MAIIGNANT TOTAL UNCEFTAIN TUMORS	м -			
TOTAL ANIMAIS WITH TUMORS UNCERTAI FRIMARY OR METASTATIC TOTAL UNCEFTAIN TUMORS	N -			

APPENDIX C

SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC

LESIONS IN RATS GIVEN INTRAPERITONEAL

INJECTIONS OF IPD

TABLE C1

SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN MALE RATS **GIVEN INTRAPERITONEAL INJECTIONS OF IPD (CONTROL GROUPS)**

'	MID- AND HIGH-DOSE UNTREATED CONTROL	LOW-OOSE UNTREATED CDNTROL	MID- AND HIGH-DOSE VEHICLE CONTROL	LOW-DOSE Vehicle Control
ANIMALS INITIALLY IN STUDY	10	10	10	10
ANIMALS NECFOPSIED Animals Examinee Histopathological	10 LY 10	10 10	10 10	10 10
INTEGUMENTARY SYSIEM				
*SKIN	(10)	(10)	(10)	(10)
INFLAMMATICN, CHRONIC SUPPURAT INFLAMMATICN, FOCAL GRANULOMAT			1 (10%)	1 (105
INFLAMMATICN WITH FIBROSIS	00	1 (10%)	1 (10%)	
HYPERKEFAICSIS PABAKERAICSIS		1 (10%) 1 (10%)		
		• •		
*SUBCUT TISSUF EPIDERMAL INCLUSION CYST	(10)	(10)	(10)	(10)
RESPIRATCRY SYSTEM				
#TRACHEA	(9)	(10)	(9)	(10)
INFLAMMATICN, CHRONIC INFLAMMATICN, CHRONIC SUPPURAT	T V	1 (10%)		2 (20)
INFLAMMATICA, CHRONIC SUPPORAT	1 V	1 (10%)		
*LUNG PNEUMONIA, CHRONIC MURINE	(10) 1 (10%)	(10)	(10) 1 (10%)	(10)
EEMATOPOIETIC SYSTEM				
*BONE MARROW	(10)	(10)	(9)	(10)
ATROPHY, NCS		5 (50%)		3 (307
#SPLEEN HEMATOPOIESIS	(10)	(10)	(10)	(10)
CIRCULATORY SYSTEM				
NONE				
LIGESTIVE SYSTEM				
NONE				

TABLE C1 CONTROL MALE RATS: NONNEOPLASTIC LESIONS (CONTINUED)

	MID-AND HIGH-OOSE UNTREATEO CONTROL	LOW-OOSE UNTREATEO CONTROL	MID- ANO HIGH-OOSE VEHICLE CONTROL	LOW-DOSE VEHICLE CONTROL
ORINARY SYSTEM				
#KIDN EY HYDRON EP FFCS IS	(10)	(10) 1 (10%)	(10)	(10)
INFLAMMATICN, INTERSTITIAL INFLAMMATICN, CHRONIC	1 (10%) 3 (30%)	6 (6 0%)	2 (20%)	3 (30%
ENDOCRINE SYSTEM				
THYROID CYSTIC FOLLICLES	(4)	(8)	(10)	(7) 1 (14 %
<pre>#PARATHYROIC HYPERPLASIA, NOS</pre>	(2)	(6) 1 (17%)	(6)	(2)
REPRODUCTIVE SYSTEM				
NONE				
NERVOUS SYSTEM				
NONE				
SPECIAL SENSE CEGANS				
NONE				
NUSCULOSKELETAI SYSTEM				
NON E				
ECDY CAVITIES				
NON E				
ALL OTHER SYSTEMS				
<u>NONE</u>				

TABLE C1 CONTROL MALE RATS: NONNEOPLASTIC LESIONS (CONTINUED)

	MID- AND HIGH-DDSE UNTREATED CONTRDL	LOW-DDSE Untreated Control	MID- ANO HIGH-DDSE VEHICLE CDNTRDL	LDW-DDSE VEHICLE CONTRDL
SPECIAL MORPHOLOGY SUMMARY				
NO LESICN REFERTED	4	2	6	2
<pre># NUMBER OF ANIMALS WITH TISSUE # NUMBER OF ANIMALS NECROPSIED</pre>	EXAMINED MICROSCOP	ICALLY		

TABLE C2

SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN MALE RATS GIVEN INTRAPERITONEAL INJECTIONS OF IPD (TREATED GROUPS)

		MID DOSE	
NIMALS INITIALLY IN STUDY NIMALS NECFOFSIED NIMALS EXAMINED HISTOPATHOLOGICALLY	35 32 32	35 28 28 28	35 30 20
NTEGUMENTARY SYSTEM			
*SUBCUT TISSUE EDEMA, NOS HEMORRHAGE INFLAMMATION, HEMORRHAGIC INPLAMMATICN, CHRONIC	1 (3%)	(28) 1 (4%)	(30) 1 (3%) 1 (3%)
ESPIRATORY SYSTEM			
TRACHEA INFLAMMATICN, ACUTE/CHRCNIC	(30)	(27)	(22) 1 (5%)
#LUNG/BRONCEUS ULCER, NOS	(31)	(28)	(25) 1 (4%)
ILUNG PNEUMONIA, CHRONIC MURINE	(31) 1 (3%)	(28) 7 (25%)	(25) 9 (36%)
EMATOPOIETIC SYSTEM			
#BONE MARROW ATROPHY, NCS	(30) 1 (3%)	(27) 8 (30%)	(29) 28 (97%)
#SPLEEN HEMATOPOIISIS	(32) 2 (6%)	(28) 1 (4%)	(23)
*LYMPH NODE Hyperplasia, plasma cell		(8)	(E) 1 (13%)
#MEDIASTINAL L.NCDF HYPERPLASIP, PLASMA CELL		(8)	(8) 2 (25%)
MESENTERIC I. NCDE		(8)	(8) <u>1 (13%)</u>

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

TABLE C2 TREATED MALE RATS: NONNEOPLASTIC LESIONS (CONTINUED)

	LOW DOSE	MID DOSE	HIGH DOSE
CIRCULATORY SYSTEM			
CIRCULATORI SISTER			
#MYOCARCIUM INPLAMMATICN, INTERSTITIAL	(31)	(28) 1 (4%)	(23)
*CARDIAC VALVE	(31)	(28)	(23)
FIBROSIS DEGENERATICN, NOS	1 (3%) 1 (3%)		
*PULMONARY ARTERY THROMBUS, CFGANIZED	(32)	(28) 1 (4%)	(30)
DIGESTIVE SYSTEM			
#LIVER	(32)	(27)	(27)
HEMORRHAGE Hematcha, Nos			2 (7%) 3 (11%)
NECROSIS, ECCAL			2 (7%)
NECRCSIS, COAGULATIVE		1 (4%)	1 (4%)
#LIVER/CENTRIIOEULAR	(32)	(27)	(27)
DEGENERATICN, NOS NECROSIS, NOS			1 (4%) 1 (4%)
#PANCREAS	(32)	(27)	(23)
INPLAMMATICN, INTERSTITIAL	(32)	1 (4%)	(23)
#COLON	(32)	(28)	(24)
HEMORR HAGE	1 (3%)		1 (4%)
ULCER, NOS	1 (3%)		1 (4%)
#CECUM HEMORR HAGE	(32) 1 (3\$)	(28)	(24)
URINARY SYSTER			
#KIDNEY	(32)	(28)	(24)
HEMORR HAGE		1 (4%)	1 (4%)
PYELCNEEHEITIS, NOS Inflammaticn, Chronic	1 (3%)	(4,4)	
ENDOCRINE SYSTEM			
#ADRENAL COFTEX	(32)	(28)	(23)
HYPERPLASIA NODULAR		2 (7%)	

NUMBER OF ANIMALS WITH TISSUE EXAMINED MICROSCOPICALLY
 NUMBER OF ANIMALS NECROPSIED

	LOW DOSE	MID DOSE	HIGH DOSE
REPRODUCTIVE SYSTEM			
<pre>#PROSTATE INFLAMMATICN, ACUTE SUPFURATIVE</pre>	(32)	(27) 1 (4%)	(23)
NERVOUS SYSTEM			
NONE			
PECIAL SENSE CEGANS			
NONE			
USCULOSKELETAI SYSTEM			
NONE			
ECTY CAVITIES			
*PERITONEUM INFLAMMATICN, SUPPURATIVE	(32) 3 (9%)	(28) 4 (14%)	(30)
INFLAMMATICN, HEMORRHAGIC INFLAMMATICN, CHRONIC	30 (94%)	23 (82%)	1 (3%) 7 (23%
ADHESION, NOS Hyperplasia, mesothelial	21 (66%)	5 (18%) 1 (4%)	
*PLEURA INFLAMMATICN, SUPPURATIVE	(32) 1 (3%)	(28)	(30)
LL OTHER SYSTEMS			
NONE			
SPECIAL MORFHCIOGY SUMMARY			
NO LESION FEFORTED	2	3	
NO NECROPSY PERFORMED AUTOLYSIS/NO NECROPSY	1 2	7	5

TABLE C2 TREATED MALE RATS: NONNEOPLASTIC LESIONS (CONTINUED)

TABLE C3

SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN FEMALE RATS GIVEN INTRAPERITONEAL INJECTIONS OF IPD (CONTROL GROUPS)

	MID- AND HIGH-DDSE UNTREATED CDNTRDL	LOW-DDSE UNTREATED CDNTRDL	MID- AND HIGH-DDSE VEHICLE CONTRDL	LDW-DOSE Vehicle CDNTRDL
ANIMALS INITIALLY IN STUDY ANIMALS NECFCESIEC ANIMALS EXAMINED HISTOPATHOLOGICAI	10 10 LY 10	10 10 10	1C 8 8 .	10 10 10
INTEGUMENTARY SYSTEM NONE				
RESPIRATORY SYSTEM				
<pre>#TRACHEA INFLAMMATICN, ACUTF/CHRCNIC INFLAMMATICN, CHRONIC INFLAMMATICN, CHRONIC SUFPURAT</pre>	(10) 1 (10%)	(10)	(8) 1 (13%)	(10) 1 (10%) 1 (10%)
*LUNG PNEUMONIA, CHRONIC MURINE	(10) 1 (10%)	(10)	(3)	(10)
HEMATOPOIETIC SYSTEM				
#EONE MARROW ATROPHY, NCS	(9)	(10) 4 (40%)	(8)	(9) 4 (44%)
CIRCULATCRY SYSTEM				
NONE				
CIGESTIVE SYSTEM				
<pre>#PANCREAS NECROSIS, NOS</pre>	(10) 1 (10%)	(10)	(8)	(10)
URINARY SYSTEM			•	
#KIDNEY <u>HYDRONEPHROSIS</u>	(9) <u>1 (11%)</u>	(10)	(8)	(10)

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

TABLE C3 CONTROL FEMALE RATS: NONNEOPLASTIC LESIONS (CONTINUED)

	MID- AND HIGH-OOSE UNTREATEO CONTROL	LOW-DOSE UNTREATED CONTROL	MIO-ANO HIGH-OOSE Vehicle Control	LOW-DOSE VEHICLE CONTROL
INFLAMMATICN, CHRONIC				2 (20%)
UFFINARY BLADDER Hyperplasi, fpithelial	(10) 1 (10%)	(10)	(9)	(10)
ENDCCRINE SYSTEM				
<pre>PITUITARY HYPERPLASIA, CHROMOPHOBE-CELL</pre>	(10)	(6) 2 (33%)	(8)	(10) 2 (20%)
NADRENAL ANGIECTASIS	(10)	(10) 2 (20%)	(8)	(10)
REPRODUCTIVE SYSTEM				
<pre>#UTERUS/ENDCMITFIUM INFLAMMATICN, SUPPURATIVE INFLAMMATICN, CHRONIC SUPPURAT</pre>	(10) TIV	(10) 3 (30%) 1 (10%)	(8) 2 (25%)	(10) 1 (10%)
NOVARY INFLAMMATICN, SUPPUPATIVE	(8)	(10)	(8) 1 (13%)	(10)
NONE				
SPECIAL SENST CEGANS				
NONE				
MUSCULOSKELETAI SYSTEM				
NONE				
ECDY CAVITIES				
NONE				
ALL OTHER SYSTEMS				
NONE				

TABLE C3 CONTROL FEMALE RATS: NONNEOPLASTIC LESIONS (CONTINUED)

	MIO- ANO HIGH-DOSE UNTREATED Control	LOW-DOSE UNTREATED CONTROL	MIO- ANO HIGH-OOSE VEHICLE Control	LOW-DOSE VEHICLE CONTROL
SPECIAL MORPHCIOGY SUMMARY				
NO LESICN FEFCRTED AUTOLYSIS/NO NECROPSY	1	2	2	1

* NUMBER OF ANIMALS WITH TISSUE * NUMBER OF ANIMALS NECROPSIED

TABLE C4

SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN FEMALE RATS **GIVEN INTRAPERITONEAL INJECTIONS OF IPD (TREATED GROUPS)**

	LOW DOSE	MID DOSE	HIGH DOSE
FNIMALS INITIALLY IN STUDY	35	35	35
ANIMALS NECFOFSIED ANIMALS EXAMINEE HISTOPATHOLOGICALLY	33	31 31	31 31
INTEGUMENTARY SYSTEM			
*SUBCUT TISSUF	(33)	(31)	(31)
EDEMA, NCS HEMOFRHAGE		1 (3%)	1 (3%)
RESPIRATORY SYSTEM			
#TRACHEA	(32)	(31)	(25)
INFLAMMATICN, ACUTE/CHRCNIC INFLAMMATICN, CHRONIC SUPPURATIV		2 (6%)	
#LUNG	(33)	(31)	(30)
EDEMA, NOS HEMOFRHAGE			2 (7%) 4 (13%
PNEUMCNIA, CHRONIC MURINE		4 (13%)	15 (50%)
EEMATOPOIETIC SYSTEM			
#BONE MARECW	(32)	(31)	(31)
ATROPHY, NCS		2 (6%)	30 (97%
#SPLEEN	(33)	(31)	(31)
PERIARTERITIS Atrophy, NCS			1 (3%) 2 (6%)
HEMATOPOIESIS	5 (15%)		1 (3%)
#MEDIASTINAL I.NCDE	(2)	(9)	(12)
HYPERPLASIA, PLASMA CELL			2 (17%
MESENTERIC I. NODE	(2)	(9)	(12)
HENORR BAGE			1 (8%)

NONE

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

TABLE C4 TREATED FEMALE RATS: NONNEOPLASTIC LESIONS (CONTINUED)

		MID DOSE	
CIGESTIVE SYSTEM			
#LIVER HEMATOMA, NOS NECROSIS, NOS NECRCSIS, FCCAL	(33)	(31)	(31) 3 (10%) 1 (3%) 1 (3%)
NECRCSIS, COAGULATIVE HE MATOFCIESIS	1 (3%)	2 (6%)	2 (6%)
<pre>*LIVER/CENTRIIOEULAR DEGENERATICN, NOS NECROSIS, NOS</pre>	(33)	(31) 1 (3%)	(31) 1 (3%) 3 (10%)
#ILEUM INFLAMMATICN, ACUTE/CHRONIC INFLAMMATICN, CHRONIC	(33)	(28)	(31) 1 (3%) 1 (3%)
URINARY SYSTEM			
#KIDNEY/TUBUIE NEPHROPATEY		(31)	(31) (3%)
ENDOCRINE SYSTEM			
# ADRENAL ANGIECTASIS	(33) 3 (9%)	(31)	(31)
<pre>#THYROID INFLAMMATION, CHRONIC</pre>	(27)	(27) 1 (4%)	(10)
REPRODUCTIVE SYSTEM			
#UTERUS CYST, NOS	(33)	(28) 1 (4%)	(30)
AUTERUS/ENCOMETRIUM INPLAMMATICN, SUPPURATIVE INPLAMMATICN, ACUTE SUPPURATIVE	(33) 1 (3%)	(28) 1 (4%)	(30)
#OVARY CIST, NOS INFLAMMATION, SUPPURATIVE	(33) 2 (6%) 1 (3%)	(22)	(25)

NERVOUS SYSTEM

NONE

NUMBER OF ANIMALS WITH TISSUE EXAMINED MICROSCCPICALLY # NUMBER OF ANIMALS NECROPSIED

		MID DOSE	HIGH DOSE
SPECIAL SENSE CRGANS			
*MIDDLE PAR INPLAMMATICN, CHRONIC SUFFURATIV		(31) 1 (3%)	(31)
MUSCULOS KELETAI SYSTEM			
*JCINT INFLAMMATICN, ACUTE/CHRCNIC INFLAMMATICN, CHRCNIC	(33)		(±1) 1 (3%) 1 (3%)
ECDY CAVITIES			
*PERITCNEUM INFLAMMATICN, SUPPURATIVE INFLAMMATION, CHRONIC ADHESICN, NOS	(33) 1 (3系) 32 (57系) 30 (51系)	(31) 5 (16%) 28 (90%) 8 (26%)	(31) 7 (23%)
* MESENTERY HEMOFRHAGE	(33) 1 (3%)	(3 1)	(31)
ALL OTHER SYSTEMS			
SEFCIAL MOREHCICGY SUMMARY			
NO LESICN FEFORIED AUTOIYSIS/NO NFCROPSY	2	1 4	4

APPENDIX D

SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN MICE GIVEN INTRAPERITONEAL INJECTIONS OF IPD



TABLE D1

SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN MALE MICE GIVEN INTRAPERITONEAL INJECTIONS OF IPD

	UNTREATED CONTROL	VEHICLE CONTROL	LOW OOSE	HIGH DOSE
ANIMALS INITIALLY IN STUDY ANIMALS MISSING	15	15	34 1	35
ANIMALS NECROFSIEL	14	14	26	21
ANIMALS EXAMINED HISTOPATHOLOGICALLY		14	26	21
INTEGUMENTARY SYSTEM				
NCNE				
RESPIRATORY SYSTEM		,		
#L UNG	(14)	(14) 2 (14%)	(26) 2 (8%)	(22)
INFLAMMATICN, INTERSTITIAL BRONCHOFNEUMONIA SUPPURATIVE	5 (36%)	2 (14%) 2 (14%)	2 (8%)	1 (5%)
EEMATOPOIETIC SYSTEM				
#ECNE MARROW Atrophy, NCS	(9)	(13)	(21)	(19) 2 (11)
#SPLEEN HEMATOPOIISIS	(14)	(14)	(24) 2 (8%)	(20)
#MESENTERIC I. NODE HYPERPLASI, IYMPHOID		(2) 1 (50%)	-(1)	(3)
CIRCULATORY SYSTEM				
NONE				
LIGESTIVE SYSTEM				
#LIVER	(14)	(14)	(25)	(20)
NECROSIS, COAGULATIVE HYPEFPLASIA, NODULAR ANGIECTASIS	1 (7%) 1 (7%)	1 (7%)	1 (4%) 1 (4%)	
URINARY SYSTEM				

NUMBER OF ANIMALS WITH TISSUE EXAMINED MICROSCCPICALLY # NUMBER OF ANIMALS NECROPSIED

TABLE D1 MALE MICE: NONNEOPLASTIC LESIONS (CONTINUED)

	UNTREATEO Control	VEHICLE Control	LOW OOSE	HIGH OOSE
NDOCBINE SYSTEM				
NONE				
EPRODUCTIVE SYSTEM				
* SEMINAL VESICLE HYPEFPLASIA, LYMPHOID	`(14)	(14) 1 (7%)	(26)	(21)
TTESTIS Atrophy, Nos	(13)	(14)	(22)	(20) 1 (5%
ERVOUS SYSTEM				
NONE				
PECIAL SENSE CEGANS				
NONE				
USCULOSRELETAL SYSTEM				
NONE				
ODY CAVITIES				
*PERITONEUM INFLAMMATICN, SUPPURATIVE	(14)	(14)	(26) 1 (4%)	(21)
INFLAMMATICN, FIBRINOUS INFLAMMATICN, CHRONIC METAFLASIA, OSSEOUS		1 (7%) 1 (7%)	2 (8%)	1 (59 5 (24
IL OTHER SYSTEMS	•			
NONE				
PECIAL MORFHCICGY SUMMARY				
NO LESION FEFORTED	9	7	13	9

TABLE D1 MALE MICE: NONNEOPLASTIC LESIONS (CONTINUED)

	UNTREATED CDNTRDL	VEHICLE	LDW DDSE	HIGH DDSE
ANIMAL MISSING/NO NECROFSY			1	
NO NECROPSI PERFORMED Autolisis/NC becropsi	1	1	7	1 13

* NUMBER OF ANIMALS NECROPSIED

TABLE D2

SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN FEMALE MICE GIVEN INTRAPERITONEAL INJECTIONS OF IPD

	UNTREATED CONTROL	VEHICLE CONTROL	LOW OOSE	HIGH DOSE
ANIMALS INITIPILY IN STUDY	15	15	36	35
ANIBALS MISSING PNIMALS NECFOFSIFC ANIMALS EXABINED HISTOPATHOLOGICALLY	13	15 15	29 29	27 26
NTEGUMENTARY SYSTEM				
NONE				
FESPIRATORY SYSTEM				
<pre>\$LUNG INFLAMMATICN, INTERSTITIAL BRONCHOPNEDMONIA SUPPURATIVE</pre>	(13) 2 (15%)	(15) 6 (40%)	(29) 3 (10%) 1 (3%)	(24) 3 (13% 2 (8%)
BEBATOPOIETIC SYSTEM				
#SPLEEN HEMATOPOIESIS	(13)	(15)	(26) 1 (4%)	(24)
<pre>#MESENTERIC I. NODE INFLAMMATICN, GRANULOMATOUS</pre>		(3) 1 (33%)		(2)
#THYMUS ATROPHY, NOS		(15)	(25)	(19) 1 (5%)
CIRCULATORY SYSTEM				
NONE				
CIGESTIVE SYSTEM				
<pre>#LIVER HYPERPLASI#, NODULAR</pre>	(13)	(15)	(29) 2 (7%)	(24)
ISTONACE ULCER, NOS			(29)	(25) 1 (4%)
IRINARY SYSTEE				
NONB				

TABLE D2 FEMALE MICE: NONNEOPLASTIC LESIONS (CONTINUED)

	UNTREATEO Control	VEHICLE Control	LOW DOSE	HIGH DOSE
ENDOCRINE SYSTEM				
<pre>#THYROID POLIICLF HYPERPLASI;, PAPILLARY</pre>	(12)	(13) 1 (8%)	(23)	(18)
REPRODUCTIVE SYSTEM				
AUTERUS/ENECKETFIUM INFLAMMATICN, SUPPURATIVE Hypefplasia, Cystic	(13) 9 (69%)	1 (7%)	(29) 10 (34%)	(23)
AOVARY Atrophy, NCS	(8)	(14)	(25)	(18) 1 (6%)
SPECIAL SENSE CEGANS NONE				
NUSCULOS KELETAL SYSTEM				
*FEMUR OSTECPOFOSIS FIBRCUS CSTECDYSTROPHY	(13)		(29) 1 (3%) 1 (3%)	(27)
ECDY CAVITIES				
*PERITONEUM INFLAMMATION, CHRONIC METAFLASIA, CSSEOUS	(13)		(29) 2 (7%) 1 (3%)	(27) 6 (221
ALL OTHER SYSTEMS				
ANIMAL MISSING/NO NECROPSY	1			

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

TABLE D2 FEMALE MICE: NONNEOPLASTIC LESIONS (CONTINUED)

	UNTREATED CONTROL	VEHICLE CONTROL	LOW DOSE	HIGH OOS
FECIAL BORPROIOGY SUBBARY				
NO LESION FEFCRIED	3	3	12	8
NECROFSY FERF/NO HISTO PERFORMED NO NECROFSY FERPORMED	1		1	1
AUTOLISIS/NO NECROPSY	•		6	8

APPENDIX E

ANALYSES OF THE INCIDENCE OF PRIMARY TUMORS IN RATS GIVEN INTRAPERITONEAL INJECTIONS

OF IPD

THE LEVELORY

Mid High Dose <u>Dose</u>	1/28 (4) 0/30 (0)	N.S. N.S.	N.S. N.S.	Infinite 0.039 Infinite	72	week.	percent).	<pre>probability level for the .) is indicated. Beneath the or the Fisher exact tests for the . P < 0.05; otherwise not</pre>	group than in the vehicle-control	P < 0.05 for any comparison.	d group and the vehicle-
Low Dose) 2/32 (6)	N.S.	N.S.	Infinite 0.192 Infinite	44	24, or 48 mg/kg three times per week.	animals examined at site (nors in the vehicle-control group is the probability level for the $P < 0.05$; otherwise, not significant (N.S.) is indicated. Beneath treated group is the probability level for the Fisher exact tests group with the vehicle-control group when $P < 0.05$; otherwise not ated.	lower incidence in a treated group	from linear trend is given when P	of the relative risk between each treated group and the vehicle-
Vehicle Control	0/20 (0)	N.S.	N•S•		1		'number of	in the veh 0.05; othen ated group with the	a lower inc		che relativ
Topography: Morphology	Peritoneum: Sarcoma, NOS, Fibroma, or Fibrosarcoma ^b	P Values (Control, Low, Mid, High Dose) ^{c,d}	P Values (Control, Low, Mid Dose) ^{c,d}	Relative Risk (Vehicle Control) ^f Lower Limit Upper Limit	Weeks to First Observed Tumor	^a Treated groups received doses of 12,	^b Number of tumor-bearing animals/number of animals examined at site (percent).	^c Beneath the incidence of tumors in the vehicle-control group is the probability level Cochran-Armitage test when $P < 0.05$; otherwise, not significant (N.S.) is indicated. incidence of tumors in each treated group is the probability level for the Fisher exac comparison of that treated group with the vehicle-control group when $P < 0.05$; otherwising the significant (N.S.) is indicated.	d _A negative trend (N) indicates a group.	^e The probability level for departure	^f The 95% confidence interval of t control group.

Analyses of the Incidence of Primary Tumors in Male Rats Given Intraperitoneal Injections of IPD^a

Table El.

Table E2. Time-ac	Time-adjusted Analyses of		mors in Male Ra	ß
Gí	Given Intraperitoneal	l Injections of IPD ^a	œ_	
Topography: Morphology	Vehicle Control	Low Dose	Mid Dose	High Dose
Peritoneum: Sarcoma, NOS, Fibroma, or Fibrosarcoma ^b	0/19 (0)	2/14 (14)	1/3 (33)	(0) 0/0
P Values (Control, Low, Mid, High Dose) ^c ,d	P = 0.031	N.S.	N.S.	ł
P Values (Control, Low, Mid Dose) ^{c,d}	P = 0.045	N.S.	N.S.	
Relative Risk (Vehicle Control) ^f Lower Limit Upper Limit		Infinite 0.421 Infinite	Infinite 0.345 Infinite	
-	12, 24, or 48 mg/	doses of 12, 24, or 48 mg/kg, three times per week.	week.	
^D Number of tumor-bearing animals/r at least 44 weeks of study.	number of animals	animals/number of animals examined at site (percent) that survived dy.	ercent) that sum	rvived
^c Beneath the incidence of tumors in the vehicle-control group is the probability level for the Cochran-Armitage test when $P < 0.05$; otherwise, not significant (N.S.) is indicated. Beneath th incidence of tumors in each treated group is the probability level for the Fisher exact tests fo the comparison of that treated group with the vehicle-control group when $P < 0.05$; otherwise not significant (N.S.) is indicated.	in the vehicle-con .05; otherwise, no ted group is the p roup with the vehi	tumors in the vehicle-control group is the probability level for the probability level for the probability of the probability level for the Fisher exact tests ceated group with the vehicle-control group when $P < 0.05$; otherwise r licated.	robability leve.) is indicated. If the Fisher exemption P < 0.05; ot	l for the Beneath the act tests for cherwise not
dA negative trend (N) indicates a group.	lower incidence in a		treated group than in the vehicle-control	cle-control
^e The probability level for departu	ure from linear tr	departure from linear trend is given when P	<pre>< 0.05 for any comparison.</pre>	comparison.

ANVERIE FILE

 f_{T} he 95% confidence interval of the relative risk between each treated group and the vehiclecontrol group.

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n Female Rats	
'y Tumors in Fem	of IPD ^a
of Primary	l Injections of IPD ^a
s of the Incidence	n Intraperitoneal I
Analyses	Given
Table E3.	

Topography: Morphology	Vehicle Control	Low Dose	Mid Dose	High Dose
Peritoneum: Sarcoma, NOS or Fibrosarcoma ^b	0/18 (0)	0/33 (0)	3/31 (10)	0/31 (0)
P Values (Control, Low, Mid, High Dose) ^c ,d	N. S.	N.S.	N.S.	N. S.
P Values (Control, Low, Mid Dose) ^{c,d}	N.S.	N.S.	N.S.	
Relative Risk (Vehicle Control) ^f Lower Limit Upper Limit			Infinite 0.367 Infinite	
Weeks to First Observed Tumor			56	8
Mammary Gland: Fibroadenoma ^b	5/18 (28)	4/33 (12)	0/31 (0)	0/31 (0)
P Values (Control, Low, Mid, High Dose) ^c ,d	P = 0.001(N)	N.S.	P = 0.004(N)	P = 0.004(N)
P Values (Control, Low, Mid Dose) ^{c,d}	P = 0.003(N)	N.S.	P = 0.004(N)	
Relative Risk (Vehicle Control) ^f Lower Limit Upper Limit		0.436 0.102 1.800	0.000 0.000 0.446	0.000 0.000 0.446
Weeks to First Observed Tumor	79	47		8

(continued)		7		
Topography: Morphology	Vehicle Control	Low Dose	Mid Dose	High Dose
Mammary Gland: Adenoma, NOS ^b	1/18 (6)	2/33 (6)	0/31 (0)	0/31 (0)
P Values (Control, Low, Mid, High Dose) ^{C,d}	N. S.	N.S.	N.S.	N. S.
P Values (Control, Low, Mid Dose) ^{c,d}	N.S.	N.S.	N.S.	
Relative Risk (Vehicle Control) ^f Lower Limit Upper Limit		1.091 0.062 62.383	0.000 0.000 10.726	0.000 0.000 10.726
Weeks to First Observed Tumor	80	29	1	-
Mammary Gland: Adenoma, NOS or Fibroadenoma ^b	6/18 (33)	6/33 (18)	0/31 (0)	0/31 (0)
P Values (Control, Low, Mid, High Dose) ^{c,d}	P < 0.001(N)	N.S.	P = 0.001(N)	P = 0.001(N)
P Values (Control, Low, Mid Dose) ^{c,d}	P = 0.001(N)	N.S.	P = 0.001(N)	
Relative Risk (Vehicle Control) ^f Lower Limit Upper Limit		0.545 0.178 1.774	0.000 0.000 0.351	0.000 0.000 0.351
Weeks to First Observed Tumor	79	29	-	-

Analyses of the Incidence of Primary Tumors in Female Rats Given Intraperitoneal Injections of IPD^a Table E3.

(continued)				
Topography: Morphology	Vehicle Control	Low Dose	Mid Dose	High Dose
Pituitary: Chromophobe Adenoma ^b	8/18 (44)	2/29 (7)	0/27 (0)	0/22 (0)
P Values (Control, Low, Mid, High Dose) ^{c,d}	P < 0.001(N)	P = 0.004(N)	P < 0.001(N)	P = 0.001(N)
P Values (Control, Low, Mid Dose) ^{c,d}	P < 0.001(N)	P = 0.004(N)	P < 0.001(N)	
Departure from Linear Trend (Control, Mid, High Dose) ^e	P = 0.001			
Relative Risk (Vehicle Control) ^f Lower Limit Upper Limit		0.155 0.019 0.675	0.000 0.000 0.279	0.000 0.000 0.338
Weeks to First Observed Tumor	80	58	-	
a Treated groups received doses of 12, 24, or 48 mg/kg three times per week.	12, 24, or 48 mg/h	kg three times per	c week.	

Table E3. Analyses of the Incidence of Primary Tumors in Female Rats Given Intraperitoneal Injections of IPD^a

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

Analyses of the Incidence of Primary Tumors in Female Rats	Given Intraperitoneal Injections of IPD ^a
Analyse	Give
Table E3.	

(continued)

Cochran-Armitage test when P < 0.05; otherwise, not significant (N.S.) is indicated. Beneath the incidence of tumors in each treated group is the probability level for the Fisher exact tests for the comparison of that treated group with the vehicle-control group when P < 0.05; otherwise not ^cBeneath the incidence of tumors in the vehicle-control group is the probability level for the significant (N.S.) is indicated.

^dA negative trend (N) indicates a lower incidence in a treated group than in the vehicle-control group. ^eThe probability level for departure from linear trend is given when P < 0.05 for any comparison.

^fThe 95% confidence interval of the relative risk between each treated group and the vehiclecontrol group.

Topography: Morphology	Vehicle Control	Low Dose	Mid Dose	High <u>Dose</u>
Peritoneum: Sarcoma NOS or Fibrosarcoma ^b	0/18 (0)	0/16 (0)	3/4 (75)	(0) 0/0
P Values (Control, Low, Mid, High Dose) ^c ,d	P < 0.001	N.S.	P = 0.003	N.S.
Departure from Linear Trend ^e	P < 0.001			
P Values (Control, Low, Mid Dose) ^{c,d}	P = 0.001	N.S.	P = 0.003	
Departure from Linear Trend ^e	P < 0.001			
Relative Risk (Vehicle Control) ^f Lower Limit Upper Limit			Infinite 3.119 Infinite	
^a Treated groups received doses of	doses of 12. 24. 48 me/ke three times ner week.	hree times ner we	, and the second se	

12, 24, 48 mg/kg three times per week. TO SASOD TACATACAT Sdno 18

^bNumber of tumor-bearing animals/number of animals examined at site (percent) which survived at least 52 weeks of study.

Table E4. Time-adjusted Analyses of Peritoneal Tumors in Female Rats Given Intraperitoneal Injections of IPD^a

Time-adjusted Analyses of Peritoneal Tumors in Female Rats Given Intraperitoneal Injections of IPD^a Table E4.

(continued)

Cochran-Armitage test when P < 0.05; otherwise, not significant (N.S.) is indicated. Beneath the incidence of tumors in each treated group is the probability level for the Fisher exact tests for the comparison of that treated group with the vehicle-control group when P < 0.05; otherwise not ²Beneath the incidence of tumors in the vehicle-control group is the probability level for the significant (N.S.) is indicated.

dA negative trend (N) indicates a lower incidence in a treated group than in the vehicle-control group. ^eThe probability level for departure from linear trend is given when P < 0.05 for any comparison.

 $^{
m fT}$ he 95% confidence interval of the relative risk between each treated group and the vehiclecontrol group.

APPENDIX F

ANALYSES OF THE INCIDENCE OF PRIMARY TUMORS IN MICE GIVEN INTRAPERITONEAL INJECTIONS

OF IPD

Mice	
Male	
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e of Primary Tumors in Male Mice	IPD ^a
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Analyses of the Incidence o	Given Intraperitoneal Injections of IPD ^a
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Table F1	

Topography: Morphology	Vehicle Control	Low Dose	High Dose
Subcutaneous Tissue: Squamous-cell Carcinoma ^b	0/14 (0)	6/26 (24)	0/21 (0)
P Valuesc,d	N.S.	N.S.	N. S .
Departure from Linear Trend ^e	P = 0.003		
Relative Risk (Vehicle Control) ^f Lower Limit Upper Limit		Infinite 0.931 Infinite	111
Weeks to First Observed Tumor	1	64	1
Hematopoietic System: Malignant Lymphoma ^b	0/14 (0)	0/26 (0)	3/21 (14)
P Values ^{c,d}	P = 0.045	N.S.	N.S.
Relative Risk (Vehicle Control) ^f Lower Limit Upper Limit		111	Infinite 0.431 Infinite
Weeks to First Observed Tumor	-	-	29

<pre>(continued) afreated groups received doses of 20 or 40 mg/kg three times per week. afreated groups received doses of 20 or 40 mg/kg three times per week. byumber of tumor-bearing animals/number of animals examined at site (percent). bumber of tumor-bearing animals/number of animals examined at site (percent). Beneath the incidence of tumors in the vehicle-control group is the probability level for the contran-tranizage test when P < 0.05; otherwise, not significant (N.S.) is indicated. Beneath th incidence of tumors in each treated group with the vehicle-control group when P < 0.05; otherwise not significant (N.S.) is indicates a lower incidence in a treated group than in the vehicle-control group. dA negative trend (N) indicates a lower incidence in a treated group than in the vehicle-control group. fThe probability level for departure from linear trend is given when P < 0.05 for any comparison. fThe 95% confidence interval of the relative risk between each treated group and the vehicle- control group.</pre>		Given Intraperitoneal Injections of IPD ^a			
^a Treated groups received doses of 20 or 40 mg/kg three times per week. ^b Number of tumor-bearing animals/number of animals examined at site (percent ^c Beneath the incidence of tumors in the vehicle-control group is the probability ^c CBeneath the incidence of tumors in the vehicle-control group when P ^{control} significant (N.S.) is indicated. ^d A negative trend (N) indicated. ^d A negative trend (N) indicates a lower incidence in a treated group than in group. ^e The probability level for departure from linear trend is given when P < 0.0 ^f The 95% confidence interval of the relative risk between each treated group.		(continued)			
bNumber of tumor-bearing animals/number of animals examined at site (percent CBeneath the incidence of tumors in the vehicle-control group is the probabi Cochran-Armitage test when $P < 0.05$; otherwise, not significant (N.S.) is incidence of tumors in each treated group is the probability level for the the comparison of that treated group with the vehicle-control group when P significant (N.S.) is indicated. dA negative trend (N) indicates a lower incidence in a treated group than in group. FThe probability level for departure from linear trend is given when $P < 0.0$ from probability level for departure from linear trend is given when $P < 0.0$ from probability level for departure from linear trend is given when $P < 0.0$		groups received		s per week.	
^C Beneath the incidence of tumors in the vehicle-control group is the probability cochran-Armitage test when P < 0.05; otherwise, not significant (N.S.) is incidence of tumors in each treated group is the probability level for the the comparison of that treated group with the vehicle-control group when P significant (N.S.) is indicated. ^d A negative trend (N) indicates a lower incidence in a treated group than it group. ^e The probability level for departure from linear trend is given when P < 0.0 ^f The 95% confidence interval of the relative risk between each treated group.			'number of animals examined	at site (percent).	
		^c Beneath the incidence of tumors Cochran-Armitage test when P < 0 incidence of tumors in each trea the comparison of that treated g significant (N.S.) is indicated.	in the vehicle-control grou 0.05; otherwise, not signifitited group is the probabilitit group with the vehicle-contr	up is the probability lev lcant (N.S.) is indicated y level for the Fisher e col group when P < 0.05;	el for the • Beneath kact tests otherwise
⁶ The probability level for departure from linear trend is given when P < 0.05 for any comparis ^f The 95% confidence interval of the relative risk between each treated group and the vehicle- control group.		dA negative trend (N) indicates a group.	a lower incidence in a treat	ced group than in the veh	icle-contr
	104	^e The probability level for depart	cure from linear trend is gi	iven when P < 0.05 for an	y comparis
		^f The 95% confidence interval of t control group.	che relative risk between ea	ich treated group and the	vehicle-

Time-adjusted Analyses of Hematopoietic Tumors in Male Mice Given Intraperitoneal Injections of IPD^a

Table F2.

High	Dose	1/27 (4)	N.S.	Infinite 0.031 Infinite	63	6/27 (22)	N.S.	3.333 0.473 146.288	25
Low	Dose	0/29 (0)	N.S.		1	2/29 (7)	N.S.	1.034 0.060 58.874	65
Vehicle	Control	0/15 (0)	N.S.		-	1/15 (7)	N.S.		86
	Topography: Morphology	Subcutaneous Tissue: Squamous-cell Carcinoma ^b	P Values ^c ,d	Relative Risk (Vehicle Control) ^f Lower Limit Upper Limit	Weeks to First Observed Tumor	Hematopoietic System: Malignant Lymphoma ^b	P Values ^c ,d	Relative Risk (Vehicle Control) ^f Lower Limit Upper Limit	Weeks to First Observed Tumor

Table F3. Analyses of the Incidence of Primary Tumors in Female MiceGiven Intraperitoneal Injections of IPD^a

Mice	
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Table F3.	
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	Vehicle	Low	High
<u>Topography: Morphology</u>	Control	Dose	Dose
Lung: Alveolar/Bronchiolar Adenoma or Carcinoma ^b	1/15 (7)	2/29 (7)	2/24 (8)
P Values ^{c,d}	N • S -	N.S.	N.S.
Relaitve Risk (Vehicle Control) ^f Lower Limit Upper Limit		1.034 0.060 58.874	1.250 0.073 70.551
Weeks to First Observed Tumor	86	84	64
Ovary: Cystadenoma ^b	1/14 (7)	9/25 (36)	0/18 (0)
P Values ^c ,d	N.S.	N.S.	N.S.
Departure from Linear Trend ^e	P = 0.002		
Relative Risk (Vehicle Control) ^f Lower Limit Upper Limit		5.040 0.838 207.589	0.000 0.000 14.053
Weeks to First Observed Tumor	86	65	-

Table F3. Analyses of the Incidence of Primary Tumors in Female Mice Given Intraperitoneal Injections of IPD ^a
(continued) aTreated groups received doses of 20 or 40 mg/kg three times per week.
^b Number of tumor-bearing animals/number of animals examined at site (percent).
^c Beneath the incidence of tumors in the vehicle-control group is the probability level for the Cochran-Armitage test when $P < 0.05$; otherwise, not significant (N.S.) is indicated. Beneath the incidence of tumors in each treated group is the probability level for the Fisher exact tests for the comparison of that treated group with the vehicle-control group when $P < 0.05$; otherwise not significant (N.S.) is indicated.
^d A negative trend (N) indicates a lower incidence in a treated group than in the vehicle-control group.
$^{ m e}$ The probability level for departure from linear trend is given when P < 0.05 for any comparison.
$^{\rm f}{\rm The}$ 95% confidence interval of the relative risk between each treated group and the vehicle-control group.

Table F4. Time-adjusted Ana Given Intrap	Time-adjusted Analyses of Hematopoietic Tumors Given Intraperitoneal Injections of IPD ^a	.c Tumors in Female Mice of IPD ^a	<i>f</i> ice
Topography: Morphology	Vehicle Control	Low Dose	High Dose
Hematopoietic System: Malignant Lymphoma ^b	1/15 (7)	2/29 (7)	6/26 (23)
P Values ^{c,d}	N.S.	N.S.	N.S.
Relative Risk (Vehicle Control) ^f Lower Limit Upper Limit		1.034 0.060 58.874	3.462 0.429 151.557
^a Treated groups received doses of 20 or 40	20 or 40 mg/kg three times per	per week.	
^b Number of tumor-bearing animals/number of animals examined at site (percent) which survived at least 25 weeks of study.	animals examined at s	ite (percent) which	survived
^c Beneath the incidence of tumors in the vehicle-control group is the probability level for the Cochran-Armitage test when $P < 0.05$; otherwise, not significant (N.S.) is indicated. Beneath th incidence of tumors in each treated group is the probability level for the Fisher exact tests fo the comparison of that treated group with the vehicle-control group when $P < 0.05$; otherwise not significant (N.S.) is indicated.	tumors in the vehicle-control group is the probability level for the n P < 0.05 ; otherwise, not significant (N.S.) is indicated. Beneath ch treated group is the probability level for the Fisher exact tests eated group with the vehicle-control group when P < 0.05 ; otherwise icated.	<pre>the probability lev (N.S.) is indicated vel for the Fisher froup when P < 0.05;</pre>	vel for the 1. Beneath the exact tests for otherwise not
d _A negative trend (N) indicates a lower incidence in a treated group than in the vehicle-control group.	cidence in a treated g	croup than in the ver	nicle-control
^e The probability level for departure from linear trend is	linear trend is given	given when P < 0.05 for any comparison.	ny comparison.
fThe 95% confidence interval of the relative risk between each treated group and the vehicle-	ve risk between each t	reated group and the	e vehicle-

control group.

APPENDIX G

HISTOLOGIC FEATURES OF TUMORS OBSERVED IN SPRAGUE-DAWLEY RATS GIVEN INTRAPERITONEAL INJECTIONS OF IPD IN BUFFERED SALINE THREE TIMES PER WEEK FOR ONE YEAR

Appendix G

Histologic Features of Tumors Observed in Sprague-Dawley Rats Given Intraperitoneal Injections of IPD in Buffered Saline Three Times per Week for One Year

<u>Skin and Ear Canal</u> - Three epithelial neoplasms were observed in male rats. Two lesions had diagnoses of keratoacanthomas; one was on the back and the other involved the external ear canal. Another related lesion was a papillomatous squamous-cell carcinoma with extensive keratinization and focal secondary suppurative inflammation. Each tumor occurred in a rat from a different group: the untreated-controls, the vehicle-controls, and the middose group.

<u>Subcutis</u> - Three pelvic mesenchymal tumors occurred in the pelvic tissues of three rats, two females and one male. All were spindle-cell tumors of various degrees of differentiation. One was classified as a well-differentiated fibroma and a second as welldifferentiated fibrosarcoma. The latter had an extensive ulcerative epidermitis over the neoplastic tissue. The third lesion was a pleomorphic fibrosarcoma with bizarre multinucleated giant cells.

<u>Mammary Glands</u> - One mid-dose female had a hemangioma composed of large cavernous blood spaces filled with erythrocytes and lined by low cuboidal or squamoid cells. No glandular tissues were involved. The adenomas, adenocarcinomas, and fibroadenomas of the mammary gland were typical of those previously described by Davis et al. (1956) and Thompson et al. (1961).

<u>Liver</u> - A hepatocellular lesion was observed in one mid-dose female rat. This lesion was termed a hepatocellular adenoma and consisted of a focal area of pleomorphic hepatocytes with great variation in their size and shape, as well as macronucleosis. The hepatocytic plates were disrupted and thickened. Marked fatty changes were observed in the hepatocytes of the affected area. Following the classification of Squire and Levitt (1975), the lesion would be classified as a neoplastic nodule.

Large Intestines - Two adenocarcinomas were observed in the gastrointestinal tracts of two mid-dose male rats. One lesion had large glandular spaces lined by columnar, cuboidal, and squamoid epithelial cells and was filled by large amounts of mucin. Glands of this tumor had invaded the muscular layers.

<u>Peritoneum</u> - Six rats, three males and three females, had eight spindle-cell mesenchymal tumors which were classified as one fibroma, three fibrosarcomas, and sarcomas, type unspecified.

Similar or related lesions have been termed also as peritoneal sarcomas (Dunning and Curtis, 1946), malignant fibrous histiocytomas (Pradham et al., 1974) and malignant mesotheliomas (Dunning and Curtis, 1946; and Robbins, 1967). They occurred only in low- and mid-dose rats. These rats had chronic peritonitis with extensive adhesions involving the abdominal viscera. Two rats had active suppurative inflammation. One rat had three peritoneal masses: one mass classified as a fibroma with well-differentiated fibroblasts and collagen, and two masses as fibrosarcomas. In another rat, multiple well-differentiated spindle-cell tumors (fibrosarcomas) were observed. These lesions involved the liver, stomach, intestines, and pancreas with metastasis to the mediastinal lymph nodes.

The four lesions given the diagnosis of sarcoma, NOS were similar in many respects to the above fibrosarcomas. One rat had multiple abdominal masses. One mass consisted of mature spindle-cells which had infiltrated the muscular wall from the serosa and blended into the small muscle, making identification of the neoplastic cells difficult. Much of this tumor resembled a fibrosarcoma, but a leiomyosarcoma could not be ruled out. Another rat had multifocal areas of spindle-cell proliferation involving the spleen, liver, adrenals, kidneys and pancreas. The muscle layers of the stomach wall were bisected by neoplastic

tissues with a confusing mixture of neoplastic and nonneoplastic cells. Unusual cells, probably regenerative and reactive leiomyocytes, were seen with fibroblasts and collagen. A pleomorphic spindle-cell sarcoma occurred in two animals. In the former, attached to the serosal of the speen, was a mass composed of spindle-cells and bizarre giant cells. The latter had a poorly differentiated sarcoma, with some areas resembling histiocytes; other areas had stromal spindle-cells. The lesions had a mixed population of small and large cells having various amounts of cytoplasm and round to oval nuclei with small nucleoli and delicate chromatin. A few areas had epithelioid cells. The liver, pancreas, large intestine, uterus, and mesenteric lymph nodes were involved. A vascular embolus of large vacuolated histiocytes was present in the lungs. No invasion of the pulmonary parenchyma was seen.

A lipoma was also seen in the mesentery of one rat. This lesion consisted of normal-appearing, well-differentiated lipocytes.

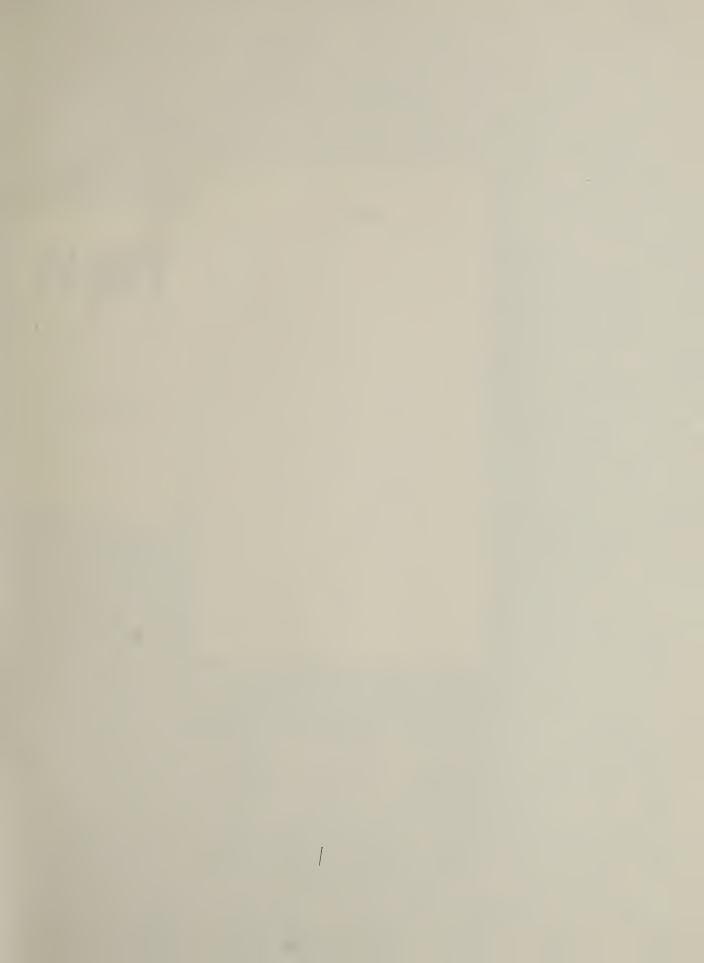
Endocrine glands - Tumors of the pituitary, adrenals, and thyroid were similar to those previously observed in Sprague-Dawley rats (Davis et al., 1956; Thompson et al., 1961; and Prejean et al., 1973).

Reproductive tract - Multiple interstitial-cell adenomas were

seen in the testes of one untreated male. These were typical of those described by Jacobs and Huseby (1967) and by Davey and Moloney (1970). Stromal polyps in the uteri of two rats were similar to those described by Jacobs and Huseby (1967), and Davey and Moloney, 1970.

Brain - Three astrocytomas of the cerebral cortex were observed; one in a vehicle-control female, one in a low-dose male, and one in a high-dose male. Two were classified as astrocytoma, NOS, and one as a gemistocytic astrocytoma. In the latter lesion, a large number of large neuron-like cells were localized around a blood-filled space. The lesion resembled a hyperplastic basal ganglion; however, the number of cells, their size, and orientation around a vascular space supports a diagnosis of gemistocytic astrocytoma. The other lesions were well-differentiated highly cellular neoplasms composed of a homogenous population of small astrocytes having round to oval nuclei with delicate chromatin and small nucleoli. These cells tended to blend into the adjacent neural tissues.

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