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BIOASSAY OF

1,5-NAPHTHALENEDIAMINE

FOR POSSIBLE CARCINOGENICITY

Carcinogenesis Testing Program Division of Cancer Cause and Prevention 20.5. National Cancer Institute '' National Institutes of Health Bethesda, Maryland 20014

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

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REPORT ON THE BIOASSAY OF 1,5-NAPHTHALENEDIAMINE FOR POSSIBLE CARCINOGENICITY

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

FOREWORD: This report presents the results of the bioassay of 1,5-naphthalenediamine conducted for the Carcinogenesis Testing Program, Division of Cancer Cause and Prevention, National Cancer Institute (NCI), National Institutes of Health, Bethesda, Maryland. This is one of a series of experiments designed to determine whether selected chemicals have the capacity to produce cancer in animals. Negative results, in which the test animals do not have a significantly greater incidence of cancer than control animals, do not necessarily mean the test chemical is not a carcinogen because the experiments are conducted under a limited set of circumstances. Positive results demonstrate that the test chemical is carcinogenic for animals under the conditions of the test and indicate a potential risk to man. The actual determination of the risk to man from animal carcinogens requires a wider analysis.

<u>CONTRIBUTORS</u>: This bioassay of 1,5-naphthalenediamine was conducted by Mason Research Institute, Worcester, Massachusetts, initially under direct contract to the NCI and currently under a subcontract to Tracor Jitco, Inc., prime contractor for the NCI Carcinogenesis Testing Program.

The experimental design was determined by the NCI Project Officers, Dr. J. H. Weisburger (1,2) and Dr. E. K. Weisburger (1). The principal investigators for the contract were Dr. E. Smith (3) and Dr. A. Handler (3). Animal treatment and observation were supervised by Mr. G. Wade (3) and Ms. E. Zepp (3). Chemical analysis was performed by Midwest Research Institute (4) and the analytical results were reviewed by Dr. N. Zimmerman (5).

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Compilation of individual animal survival, pathology, and summary tables was performed by EG&G Mason Research Institute (7); the statistical analysis was performed by Mr. W. W. Belew (5,8) and Mr. R. M. Helfand (5), using methods selected for the Carcinogenesis Testing Program by Dr. J. J. Gart (9).

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SUMMARY

A bioassay of 1,5-naphthalenediamine for possible carcinogenicity was conducted using Fischer 344 rats and B6C3F1 mice. 1,5-Naphthalenediamine was administered in the feed, at either of two concentrations, to groups of 50 male and 50 female animals of each species. The high and low dietary concentrations utilized in the chronic bioassay were, respectively, 0.1 and 0.05 percent for rats and 0.2 and 0.1 percent for mice. The compound was administered in the diet for 103 weeks, followed by up to 4 weeks of observation. Fifty mice of each sex and 25 rats of each sex were placed on test as controls. These animals were observed for up to 110 weeks.

There were no significant positive associations between the administered concentrations of 1,5-naphthalenediamine and mortality in either sex of rats or mice. In all groups adequate numbers of animals survived sufficiently long to be at risk from late-developing tumors.

Among dosed female rats, a statistically significant increase in endometrial stromal polyps was observed. Several of these tumors underwent malignant transformation to endometrial stromal sarcomas. The incidence of female rats having either adenoma or carcinoma of the clitoral gland was statistically significant. No neoplasms were observed at significantly increased incidences in dosed male rats. Based on lack of clinical signs or weight loss, the male rats may have been able to withstand a higher dose.

In mice, dose-related increases in thyroid neoplasms were observed in both sexes. The incidence of thyroid C-cell carcinomas was significant for high dose female mice. The combined incidences of papillary adenomas, follicular-cell adenomas and papillary cystadenomas of the thyroid were significant for mice of both sexes. The incidence of hepatocellular carcinomas and the incidence of alveolar/bronchiolar adenomas were each significant for dosed female mice.

Under the conditions of this bioassay, 1,5-naphthalenediamine was carcinogenic in female Fischer 344 rats, causing clitoral and uterine neoplasms. 1,5-Naphthalenediamine was also carcinogenic for B6C3F1 mice, producing thyroid neoplasms in males and neoplasms of the thyroid, liver, and lung in females. Insufficient evidence was provided for the carcinogenicity of the compound in male Fischer 344 rats.

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I. INTRODUCTION

1,5-Naphthalenediamine (Figure 1) (NCI No. CO3021), a bicyclic aromatic amine used in the dye industry, was selected for bioassay by the National Cancer Institute because of the high incidence of bladder cancer reported among dye manufacturing industry workers (Anthony and Thomas, 1970; Wynder et al., 1963). Aromatic amines are one class of chemicals believed to contribute to the increased cancer risk in this industry (Wynder et al., 1963). The structural similarity of 1,5naphthalenediamine to both the human bladder carcinogen 2-naphthylamine (International Agency for Research on Cancer [IARC], 1974) and the suspected carcinogen 1-naphthylamine (IARC, 1974) was an additional factor in its selection for testing.

The Chemical Abstracts Service (CAS) Ninth Collective Index (1977) name for this compound is 1,5-naphthalenediamine.^{*} It is also known as 1,5-diaminonaphthalene.

1,5-Naphthalenediamine can be used as an oxidation base (Colour Index [C.I.] 76595), an intermediate in the synthesis of the dye Naphthylene Red (C.I. 21650) (Society of Dyers and Colourists, 1956), and in the production of a black trisazo dye for cotton (Taube, 1973). 1,5-Naphthalenediamine has also been used as a precursor for 1,5-naphthalenediisocyanate (Hirai and Yamamoto, 1975); as an intermediate in the synthesis of drugs for the symptomatic treatment of asthma or

The CAS registry number is 2243-62-1

¹



FIGURE 1 CHEMICAL STRUCTURE OF 1,5-NAPHTHALENEDIAMINE

rhinitis (Hall, 1976); as a component of piperazine-modified aromatic polyamides (Fujiwara et al., 1974); and as a modifier for phenolic resins used in rapid curing compounds (Freeman et al., 1974); however, these uses appear to be purely experimental.

Specific production data for 1,5-naphthalenediamine are not available; however, the exclusion of this compound from the <u>1977</u> <u>Directory of Chemical Producers, U.S.A</u>. (Stanford Research Institute, 1977) implies that it is not produced in commercial quantities (in excess of 1000 pounds or \$1000 in value annually).

The potential for exposure to 1,5-naphthalenediamine may be greatest for workers in the dye industry and persons engaged in chemical research with this compound.

II. MATERIALS AND METHODS

A. Chemicals

1,5-Naphthalenediamine was purchased from Carroll Products, Wood River Junction, Rhode Island by the NCI for Mason Research Institute. Worcester, Massachusetts, and chemical analysis was performed by Midwest Research Institute, Kansas City, Missouri. The experimentally determined melting point of 190° to 191°C suggested a compound of high purity based on its narrow range and its close proximity to the value (190°C) reported in the literature (Pollock and Stevens, 1965). Elemental analysis was consistent with $C_{10}H_{10}N_{2}$, the molecular formula for 1,5-naphthalenediamine. However, nonaqueous amine group titration was approximately 89 to 90 percent of that expected on a theoretical basis. Vapor-phase chromatography revealed one homogeneous peak, but thin-layer chromatography utilizing two solvent systems (acetone:ammonium hydroxide and methylethylketone:formic acid), each visualized with 254 nm and 367 nm light, indicated the presence of one nonmotile impurity. Nuclear magnetic and infrared analyses were consistent with the structure of the compound. Ultraviolet analysis showed λ_{max} at 232, 328 and 498 nm with ϵ values of 62,800, 10,640 and 9, respectively. The literature (Sadtler Standard Spectra) indicates a λ_{max} at 328.5 nm with $\epsilon = 10,000$ for 1,5-naphthalenediamine. The observed € at 328 nm was 10,640 (6 percent greater than expected).

Throughout this report the term 1,5-naphthalenediamine is used to represent this compound.

B. Dietary Preparation

The basal laboratory diet for both dosed and control animals consisted of Wayne Lab-Blox[®] (Allied Mills, Inc., Chicago, Illinois). 1,5-Naphthalenediamine was administered to the dosed animals as a component of the diet. Under an exhaust hood, proper amounts of the chemical were removed from the stock bottle. The compound was blended in an aluminum bowl with an aliquot of the ground feed. Once visual homogeneity was attained, the mixture was placed into a 6 kg capacity Patterson-Kelley twin-shell stainless steel V-blender, along with the remainder of the meal and blended for 20 minutes. Prepared diets were placed in double plastic bags and stored in the dark at 4°C. The mixture was used for 1 week only.

C. Animals

Two animal species, rats and mice, were used in the carcinogenicity bioassay. Fischer 344 rats and B6C3F1 mice were obtained through contracts of the Division of Cancer Treatment, National Cancer Institute. All animals were obtained from Charles River Breeding Laboratories, Inc., Wilmington, Massachusetts. Dosed and control animals were received in separate shipments. Upon arrival, a sample of animals was examined for parasites and other signs of disease. All animals appeared to have parasites. They were treated with 3.0 gm of piperazine adipate per liter of drinking water, <u>ad libitum</u>, for 3 days, followed by 3 days of plain tapwater and 3 subsequent days of piperazine adipate administration. During this period, new cages

with fresh bedding were provided daily. Animals were held in quarantine by species for 2 weeks prior to initiation of test. Animals were assigned to groups and distributed among cages so that average body weight per cage was approximately equal for a given sex and species.

D. Animal Maintenance

All animals were housed by species in rooms having a temperature range of 23° to 34°C. Incoming air was filtered through Tri-Dek[®] 15/40 denier Dacron[®] filters (Tri-Dim Filter Corp., Hawthorne, New Jersey) providing six changes of room air per hour. Fluorescent lighting was provided on a 12-hour-daily cycle.

Rats were housed five per cage by sex. During quarantine and for the first 14 months of study rats were housed in galvanized-steel wire-mesh cages suspended over newspapers. Newspapers under cages were replaced daily and cages and racks washed weekly. For the remainder of the study, rats were held in suspended polycarbonate cages equipped with disposable nonwoven fiber filter sheets. Clean bedding and cages were provided twice weekly. SAN-I-CEL[®] corncob bedding (Paxton Processing Company, Paxton, Illinois) was used for the first 2 months that rats were housed in polycarbonate cages. For the remainder of the study, Aspen hardwood chip bedding (American Excelsior Company, Baltimore, Maryland) was used. Stainless steel cage racks were cleaned once every 2 weeks, and disposable filters were replaced at that time.

Mice were housed by sex in polycarbonate shoe box type cages. Cages were fitted with perforated stainless steel lids (Lab Products, Inc., Garfield, New Jersey). Nonwoven fiber filter bonnets were used over cage lids. Control mice were housed ten per cage for the first month of study and five per cage thereafter. Dosed mice were held five per cage throughout the study. Clean cages, lids, and bedding were provided twice per week. SAN-I-CEL[®] was used during the first 9 months of study. A second corncob bedding (Bed-o-Cobs[®], The Andersons Cob Division, Maumee, Ohio) was used for the next 8 months. Aspen bedding was used for the remainder of the study. Reusable filter bonnets and pipe racks were sanitized every 2 weeks throughout the study.

Water was available from 250 ml polycarbonate water bottles equipped with rubber stoppers and stainless steel sipper tubes. Bottles were replaced twice weekly and, for rats only, water was supplied as needed between changes. Food and water were available ad libitum.

Wayne Lab-Blox[®] meal was supplied to rats for 12 months and mice for 11 months from Alpine[®] aluminum feed cups (Curtin Matheson Scientific, Inc., Woburn, Massachusetts) containing stainless steel baffles. After that period, meal was supplied from stainless steel gangstyle food hoppers (Scientific Cages, Inc., Bryan, Texas). During the 2-year period of chemical administration, dosed animals were supplied

with meal containing the appropriate concentrations of 1,5-naphthalenediamine. Control animals had untreated meal available. Food hoppers were changed on the same schedule as were cages. Food was replenished daily in Alpine[®] feed cups.

All rats utilized in the 1,5-naphthalenediamine bioassay were housed in a room with other rats receiving diets containing * acetylaminofluorene (53-96-3); sodium nitrite (76-32-00-0); L-arginine glutamate (4320-30-3); N-butylurea (592-31-4); N,N-dimethyl-p-nitrosoaniline (138-89-6); 2,5-toluenediamine sulfate (6369-59-1); 2,4-dinitrotoluene (121-14-2); 4-nitroanthranilic acid (619-17-0); N-(1-naphthyl)ethylenediamine dihydrochloride (1465-25-4); 2-chloro-p-phenylenediamine sulfate (61702-44-1); aniline hydrochloride (142-04-1); and p-anisidine hydrochloride (20265-97-8).

Dosed mice were in a room with mice intubated with m-cresidine (102-50-1); and with other mice receiving diets containing N-(1-naphthyl)ethylenediamine dihydrochloride (1465-25-4) and 1H-benzotriazole (95-14-7). Control mice were in a room with other mice receiving diets containing hydrazobenzene (530-50-7); 2,3,5,6-tetrachloro-4nitroanisole (2438-88-2); tris(2,3-dibromopropyl)phosphate (126-72-7); N-(1-naphthyl)ethylenediamine dihydrochloride (1465-25-4); aniline hydrochloride (142-04-1); and 2-chloro-o-phenylenediamine sulfate.

CAS registry numbers are given in parentheses.

E. Selection of Initial Concentrations

In order to establish the maximum tolerated concentrations of 1,5-naphthalenediamine for administration to dosed animals in the chronic studies, subchronic toxicity studies were conducted with both rats and mice. Animals of each species were distributed among six groups, each consisting of five males and five females. 1,5-Naphthalenediamine was incorporated into the basal laboratory diet and supplied <u>ad libitum</u> to five of the six rat groups and five of the six mouse groups in concentrations of 0.03, 0.1, 0.3, 1.0, and 3.0 percent. The sixth group of each species served as a control group, receiving only the basal laboratory diet. The dosed dietary preparations were administered for 8 weeks.

The highest concentration causing no deaths, no compound-related gross abnormalities, and no mean body weight depression in excess of 20 percent relative to controls was selected as the high concentration for the chronic bioassay.

Deaths were recorded for all groups of rats receiving concentrations of 0.3 percent or more. Mean body weight depression was approximately 19 and 9 percent, respectively, in males and females dosed with 0.1 percent 1,5-naphthalenediamine. The concentration of 1,5-naphthalenediamine selected for administration as the high dose in the rat chronic bioassay was 0.1 percent.

Deaths were recorded for all groups of mice receiving concentrations of 0.3 percent or more and in the group of female mice

receiving 0.03 percent. Mean body weight depression was approximately 22 and 3 percent, respectively, in males and females dosed with 0.3 percent. Males receiving 0.1 percent experienced mean body weight depression of approximately 3 percent, while females receiving the same concentration had a greater mean body weight than the controls. The concentration of 1,5-naphthalenediamine selected for administration as the high dose in the mouse chronic bioassay was 0.2 percent.

F. Experimental Design

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

Rats were all approximately 7 weeks old at the time they were placed on test. Dosed rats were born approximately 1 month earlier . than controls and were started on test 1 month earlier than controls. The dietary concentrations of 1,5-naphthalenediamine administered were 0.10 and 0.05 percent. Throughout this report those rats receiving the former concentration are referred to as the high dose groups and those receiving the latter concentration are referred to as the low dose groups. The dosed rats were supplied with feed containing 1,5-naphthalenediamine for a total of 103 weeks, followed by a 3- to 4-week observation period.

All mice were approximately 7 weeks old at the time they were placed on test. Dosed mice were born approximately 1 month earlier

TABLE 1

DESIGN SUMMARY FOR FISCHER 344 RATS 1,5-NAPHTHALENEDIAMINE FEEDING EXPERIMENT

	INITIAL GROUP SIZE	1,5-NAPHTHALENE- DIAMINE CONCENTRATION (PERCENT)	OBSERVAT TREATED (WEEKS)	ION PERIOD UNTREATED (WEEKS)
MALE				
CONTROL	25	0	0	109
LOW DOSE	50	0.05 0	103	3
HIGH DOSE	50	0.10 0	103	3
FEMALE			<u>, , , , , , , , , , , , , , , , , , , </u>	
CONTROL	25	0	0	110
LOW DOSE	50	0.05 0	103	3
HIGH DOSE	50	0.10 0	103	4

TABLE 2

DESIGN SUMMARY FOR B6C3F1 MICE 1,5-NAPHTHALENEDIAMINE FEEDING EXPERIMENT

	INITIAL GROUP SIZE	l,5-NAPHTHALENE- DIAMINE CONCENTRATION (PERCENT)	OBSERVAT TREATED (WEEKS)	ION PERIOD UNTREATED (WEEKS)
MALE				
CONTROL	50	0	0	109
LOW DOSE	50	0.1 0	103	2
HIGH DOSE	50	0.2 0	103	2
FEMALE				
CONTROL	50	0	0	109
LOW DOSE	50	0.1 0	103	2
HIGH DOSE	50	0.2 0	103	3

than controls and were started on test 1 month earlier than controls. The dietary concentrations of 1,5-naphthalenediamine administered were 0.2 and 0.1 percent. Throughout this report those mice receiving the former concentration are referred to as the high dose groups and those receiving the latter concentration are referred to as the low dose groups. The dosed mice were supplied with feed containing 1,5-naphthalenediamine for a total of 103 weeks, followed by a 2- to 3-week observation period.

G. Clinical and Histopathologic Examinations

Animals were weighed immediately prior to initiation of the experiment. Body weights were recorded twice weekly for the first 12 weeks of the study and at monthly intervals thereafter. From the first day, all animals were inspected twice daily for mortality. Food consumption, for two cages from each group, was monitored for seven consecutive days once a month for the first nine months of the bioassay and for three consecutive days each month thereafter. The presence of tissue masses and lesions was determined by monthly observation and palpation of each animal.

A necropsy was performed on each animal regardless of whether it died, was killed when moribund, or was sacrificed at the end of the bioassay. The animals were euthanized by carbon dioxide inhalation, and were immediately necropsied. The histopathologic examination consisted of gross and microscopic examination of major tissues, organs,

and gross lesions taken from sacrificed animals and, whenever possible, from animals found dead.

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

Slides were prepared from the following tissues: skin, subcutaneous tissue, larynx, lungs and bronchi, trachea, bone marrow, spleen, lymph nodes, thymus, heart, salivary gland, liver, gallbladder (mice), pancreas, esophagus, stomach, small intestine, large intestine, kidney, urinary bladder, pituitary, adrenal, thyroid, parathyroid, ear, brain, testis, prostate, uterus, mammary gland, and ovary.

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

H. Data Recording and Statistical Analyses

Pertinent data on this experiment have been recorded in an automatic data processing system, the Carcinogenesis Bioassay Data System (Linhart et al., 1974). The data elements include descriptive information on the chemicals, animals, experimental design, clinical

observations, survival, body weight, and individual pathologic results, as recommended by the International Union Against Cancer (Berenblum, 1969). Data tables were generated for verification of data transcription and for statistical review.

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

Probabilities of survival were estimated by the product-limit procedure of Kaplan and Meier (1958) and are presented in this report in the form of graphs. Animals were statistically censored as of the time that they died of other than natural causes or were found to be missing; animals dying from natural causes were not statistically censored. Statistical analyses for a possible dose-related effect on survival used the method of Cox (1972) when testing two groups for equality and used Tarone's (1975) extensions of Cox's methods when testing a dose-related trend. One-tailed P-values have been reported for all tests except the departure from linearity test, which is only reported when its two-tailed P-value is less than 0.05.

The incidence of neoplastic or nonneoplastic lesions has been given as the ratio of the number of animals bearing such lesions at a specific anatomic site (numerator) to the number of animals in which that site was examined (denominator). In most instances, the denominators included only those animals for which that site was examined

histologically. However, when macroscopic examination was required to detect lesions prior to histologic sampling (e.g., skin or mammary tumors), or when lesions could have appeared at multiple sites (e.g., lymphomas), the denominators consist of the numbers of animals necropsied.

The purpose of the statistical analyses of tumor incidence is to determine whether animals receiving the test chemical developed a significantly higher proportion of tumors than did the control animals. As a part of these analyses, the one-tailed Fisher exact test (Cox, 1970, pp. 48-52) was used to compare the tumor incidence of a control group to that of a group of treated animals at each dose level. When results for a number of treated groups, k, are compared simultaneously with those for a control group, a correction to ensure an overall significance level of 0.05 may be made. The Bonferroni inequality (Miller, 1966, pp. 6-10) requires that the P-value for any comparison be less than or equal to 0.05/k. In cases where this correction was used, it is discussed in the narrative section. It is not, however, presented in the tables, where the Fisher exact P-values are shown.

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

noted, the direction of the significant trend was a positive dose relationship. This method also provides a two-tailed test of departure from linear trend.

A time-adjusted analysis was applied when numerous early deaths resulted from causes that were not associated with the formation of tumors. In this analysis, deaths that occurred before the first tumor was observed were excluded by basing the statistical tests on animals that survived at least 52 weeks, unless a tumor was found at the anatomic site of interest before week 52. When such an early tumor was found, comparisons were based exclusively on animals that survived at least as long as the animal in which the first tumor was found. Once this reduced set of data was obtained, the standard procedures for analyses of the incidence of tumors (Fisher exact tests, Cochran-Armitage tests, etc.) were followed.

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

relationship. Significant departures from linearity (P < 0.05, two-tailed test) were also noted.

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

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

the lower limit indicates the absence of a significant result while the upper limit indicates that there is a theoretical possibility of the induction of tumors by the test chemical which could not be detected under the conditions of this test.

III. CHRONIC TESTING RESULTS: RATS

A. Body Weights and Clinical Observations

There was no appreciable depression in mean body weight when dosed rats were compared with their respective controls (Figure 2).

Subcutaneous masses were observed in 2 high dose, 3 low dose, and 1 control males, and in 12 high dose, 3 low dose, and 2 control females. Crusted cutaneous masses occurred in 4 high dose males, 1 low dose male, 2 low dose females, and 1 control female, while firm nodular growths were detected in 1 high dose, 2 low dose, and 2 control males, and in 1 low dose female. Swelling of the eyes was exhibited by 2 high dose males, 2 high dose females, and 2 low dose females and swelling of the nose by 1 low dose male. Only 1 control female experienced crusted lesions in the vaginal area while 4 low dose and 9 high dose females were so effected. Alopecia was recorded for 1 low dose female, emaciation was observed in 1 male and 1 female control, and 1 female control exhibited abdominal distention.

B. Survival

The estimated probabilities of survival for male and female rats in the control and 1,5-naphthalenediamine-dosed groups are shown in Figure 3. There was no significant positive association between dosage and mortality for either male or female rats.

Adequate numbers of male rats were at risk from late-developing tumors with 74 percent (37/50) of the high dose, 80 percent (40/50)of the low dose and 68 percent (17/25) of the control surviving on





FIGURE 2 GROWTH CURVES FOR 1,5-NAPHTHALENEDIAMINE CHRONIC STUDY RATS



FIGURE 3 SURVIVAL COMPARISONS OF 1,5-NAPHTHALENEDIAMINE CHRONIC STUDY RATS
test until the termination of the study. No lesions were reported for the 4 control rats that died in week 55.

With 76 percent (38/50) of the high dose, 76 percent (38/50) of the low dose and 64 percent (16/25) of the control rats surviving on test until the termination of the study, adequate numbers of females were at risk from late-developing tumors.

C. Pathology

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

The incidence of liver neoplasms in male and female rats administered 1,5-naphthalenediamine in the diet appeared to be increased relative to controls. In female rats, tumors of the clitoral gland, uterus, and C-cell neoplasms of the thyroid appeared to be related to compound administration. The incidences of these tumors are as follows:

		MALES		F	EMALES	
	Con-	Low	High	Con-	Low	High
	trol	Dose	Dose	trol	Dose	Dose
LIVER (Number of animals with tissues					<i>.</i>	
examined histopathologically)	(25)	(49)	(49)	(24)	(50)	(49)
Neoplastic Nodule	1	3	2	0	3	4
Hepatocellular Carcinoma	0	4	2	0	1	0
PREPUTIAL/CLITORAL GLAND		()	()	()	()	
(Number of animals necropsied)	(25)	(49)	(50)	(24)	(50)	(50)
Carcinoma	0	0	1	1	3	8
Adenoma	0	0	1	0	0	5

		MALES		F	EMALES	
	Con-	Low	High	Con-	Low	High
	<u>trol</u>	Dose	Dose	<u>trol</u>	Dose	Dose
UTERUS AND ENDOMETRIUM (Number of animals with tissues						
examined histopathologically)	-	-	-	(24)	(49)	(48)
Adenocarcinoma Endometrial Stromal Polyp Endometrial Stromal Sarcoma				1 2 1	2 14 2	4 20 2
THYROID						
(Number of animals with tissues examined histopathologically) C-Cell Adenoma C-Cell Carcinoma	(21) 0 2	(47) 2 3	(47) 5 3	(21) 0 1	(49) 7 5	(48) 3 1

Neoplasms of the clitoral (preputial) gland were presented grossly as round, fluctuant cystic subcutaneous lesions in the genital area, which on section were filled with pasty green material. On microscopic examination, the cyst contents consisted of desquamated epithelial cells, frequently mixed with leukocytes from secondary inflammation. The inner portion of the cyst wall was lined by hyperkeratinized squamous epithelium often thrown into papillary folds. Peripheral to this was a zone of large, round glandular cells at least a few of which had coarse, brightly eosinophilic cytoplasmic granules. If the peripheral border appeared smooth and intact, the lesion was classified as an adenoma. If there was disorganization of the glandular structure and invasion into the surrounding stroma, the tumor was called a carcinoma.

Thyroid C-cell tumors were observed in dosed female rats at incidences increased relative to controls (4/48 [8 percent] high dose, 12/49 [24 percent] low dose, 1/21 [5 percent] controls). Ccell adenomas were discrete masses of these cells, often containing small cysts lined by flat epithelium and containing colloid-like material. In C-cell carcinomas, the tumor cells often assumed a spindle shape and tended to invade surrounding tissue.

Uterine horns containing neoplasms were usually grossly enlarged. The neoplasms themselves were varicolored, polypoid, frequently gelatinous masses projecting into the uterine cavity. Endometrial stromal polyps had a fibrous connective tissue core richly supplied with large vessels. The surface of the polyps was covered with welldifferentiated endometrium which often formed glands in the superficial portion of the polyps. These tumors frequently became necrotic at the tip and exhibited hemorrhage and secondary inflammation. In a few rats, the connective tissue stroma of these lesions underwent malignant transformation characterized by increased cellularity, mitoses, and formation of plump, pleomorphic nuclei. Such tumors were classified as stromal sarcomas. A uterine adenocarcinoma was a collection of fairly well-differentiated glands arranged back-to-back with no obvious intervening stroma. Nuclei of the glands were markedly pleomorphic with frequent mitoses. There was invasion into the myometrium and sometimes into extra uterine structures.

There were instances in this study, as noted in the summary tables, where neoplastic lesions occurred only in dosed animals, or with increased frequency when compared to the control group. No pulmonary neoplasms were found in the controls; alveolar/bronchiolar tumors were seen in dosed rats of both sexes. There was only one urinary tract neoplasm in a female control; a few more occurred in dosed rats, both male and female. No gliomas of the brain were seen in controls; a few gliomas were found in dosed rats of both sexes. These neoplasms occurred in such small numbers that a conclusive interpretation as to their significance is not possible.

Rats in all groups exhibited a variety of nonneoplastic inflammatory and degenerative changes, and none were associated with administration of the compound.

Based upon the results of this pathologic examination, 1,5-naphthalenediamine was carcinogenic to female Fischer 344 rats since feeding of the compound was associated with adenomas and carcinomas of the clitoral gland. In addition, 1,5-naphthalenediamine feeding appeared to be associated with increased incidences of thyroid, liver and uterine neoplasms in female rats and liver neoplasms in male rats.

D. Statistical Analyses of Results

The results of the statistical analyses of tumor incidence in rats are summarized in Tables 3 and 4. The analysis is included for

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ANALYSES OF THE INCIDENCE OF PRIMARY TUMORS AT SPECIFIC SITES IN MALE RATS TREATED WITH 1,5-NAPHTHALENEDIAMINE^a

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TOPOGRAPHY : MORPHOLOGY	CONTROL	DOSE	DOSE
Subcutaneous Tissue: Fibroma ^b	1/25(0.04)	3/49(0.06)	2/50(0.04)
P Values ^c	N.S.	N.S.	N.S.
Relative Risk (Control) ^d Lower Limit		1.531 0.133	1.000 0.056
Upper Limit		78.493	56.712
Weeks to First Observed Tumor	66	106	102
Skin: Squamous-Cell Papilloma ^b	2/25(0.08)	1/49(0.02)	1/50(0.02)
P Values ^c	N.S.	N.S.	N.S.
Relative Risk (Control) ^d		0.255	0.250
Lower Limit		0.005	0.004
Upper Limit		4./0/	4*0T0
Weeks to First Observed Tumor	109	106	106
Lung: Alveolar/Bronchiolar Adenoma or Alveolar/Bronchiolar Carcinoma ^b	0/25(0.00)	3/49(0.06)	4/47(0.09)
C			N C
P Values	N. V.	N. J.	N. J.
Relative Risk (Control) ^d		Infinite	Infinite
Lower Limit		0.315	0.508
Upper Limit		Infinite	Infinite
Weeks to First Observed Tumor		104	106

HIGH DOSE	10/50(0.20)	N.S.	5.000 0.787 213.351	26	4/49(0.08)	N.S.	2.041 0.218 96.949	104	11/44(0.25)	N.S.	2.750 0.683 24.081	65
LOW DOSE	10/49(0.20)	N.S.	5.102 0.801 212.137	100	7/49(0.14)	N.S.	3.571 0.503 156.046	106	7/44(0.16)	N.S.	1.750 0.376 16.365	96
CONTROL	1/25(0.04)	N.S.		109	1/25(0.04)	N.S.		109	2/22(0.09)	N.S.		98
TOPOGRAPHY: MORPHOLOGY	Hematopoietic System: Leukemia or Malignant Lymphoma ^b	P Values ^C	Relative Risk (Control) ^d Lower Limit Upper Limit	Weeks to First Observed Tumor	Liver: Hepatocellular Carcinoma or Neoplastic Nodule ^b C	P Values	Relative Risk (Control) ⁻ Lower Limit Upper Limit	Weeks to First Observed Tumor	Pituitary: Adenoma NOS, Chromophobe Adenoma, Acidophil Adenoma, or Basophil Adenoma ^b	P Values ^c	Relative Risk (Control) ^d Lower Limit Upper Limit	Weeks to First Observed Tumor

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TOPOGRAPHY : MORPHOLOGY	CONTROL	LOW DOSE	HIGH DOSE
Adrenal: Pheochromocytoma or Malignant Pheochromocytoma ^b	2/24(0.08)	4/48(0.08)	5/48(0.10)
P Values ^c	N.S.	N.S.	N.S.
Relative Risk (Control) ^d	-	1.000	1.250
Upper Limit		10.563	0.220 12.529
Weeks to First Observed Tumor	109	106	102
Thyroid: C-Cell Carcinoma ^b	2/21(0.10)	3/47(0.06)	3/47(0.06)
P Values ^c	N.S.	N.S.	N.S.
Relative Risk (Control) ^d		0.670	0.670
Lower Limit Upper Limit		0.084 7.650	0.084 7.650
Weeks to First Observed Tumor	97	100	106
Thvroid: C-Cell Adenoma or C-Cell			
Carcinomab	2/21(0.10)	5/47(0.11)	8/47(0.17)
P Values ^c	N.S.	N.S.	N.S.
Relative Risk (Control) ^d		1.117	1.787
Lower Limit	-	0.205	0.405
Upper Limit		11.249	16.445
Weeks to First Observed Tumor	97	100	104

TOPOGRAPHY: MORPHOLOGY	CONTROL	LOW DOSE	HIGH DOSE
<pre>Pancreatic Islets: Islet-Cell Adenoma or Islet-Cell Carcinoma^b</pre>	1/25(0.04)	2/48(0.04)	5/45(0.11)
P Values ^c	N.S.	N.S.	N.S.
Relative Risk (Control) ^d		1.042	2.778
Lower Limit Upper Limit		0.038 60.184	0.340 128.213
Weeks to First Observed Tumor	98	106	104
Testis: Interstitial-Cell Tumor ^b	21/25(0.84)	44/49(0.90)	45/49(0.92)
P Values ^c	N.S.	N.S.	N.S.
Relative Risk (Control) ^d		1.069	1.093
Lower Limit		0.890	0.912
Upper Limit		c25.1	L.324
Weeks to First Observed Tumor	97	94	65
^a Treated groups received doses of 0.05 or	0.10 percent in	feed.	
^b Number of tumor-bearing animals/number o	f animals examir	ed at site (proport	ion).
^C The probability level for the Cochran-Ar the control group when P < 0.05; otherwi	mitage test is g se, not signific	iven beneath the in ant (N.S.) is indic	cidence of tumors in ated. The probability
given beneath the incidence of tumors in cant (N.S.) is indicated. For both Coch	the treated gro	up when P < 0.05; o Fisher exact tests	therwise, not signifi- a negative designa-

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

tion (N) indicates a lower incidence in the treated group(s) than in the control group.

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ANALYSES OF THE INCIDENCE OF PRIMARY TUMORS AT SPECIFIC SITES IN FEMALE RATS TREATED WITH 1,5-NAPHTHALENEDIAMINE^a

I CONTROL I	System: Leukemia or ymphoma ^b 3/24(0.13) 7/50 N.S. h	(Control) ^d Lower Limit (Upper Limit	t Observed Tumor 94	ocellular Carcinoma or Nodule ^b N.S. N.S. ^N .S.	(Control) ^d InfLower Limit(Upper LimitInf	t Observed Tumor	denoma NOS, Chromophobe idophil Adenoma, or Baso-6/21(0.29) 10/50 ^{ab} N.S. N	(Control) ^d ((Lower Limit () Upper Limit 2	+ Observed Tumor 91
OW HIGH OSE DOSE	(0.14) 1/50(0.02) .S. N.S.	.120 0.160 .287 0.003 .292 1.890	76 103	(0.08) 4/49(0.08) .S. N.S.	inite Infinite .458 0.467 inite Infinite	02 106	(0.20) 17/47(0.36) .S. N.S.	.700 1.266 .275 0.577 .090 3.426	98 98

TOPOGRAPHY: MORPHOLOGY	CONTROL	LOW DOSE	HIGH DOSE
Pituitary: Carcinoma NOS, Adenoma NOS, Chromophobe Adenoma, Chromophobe Car- cinoma, Acidophil Adenoma, or Basophil Adenoma ^b	6/21(0.29)	11/50(0.22)	18/47(0.38)
P Values ^c	N.S.	N.S.	N.S.
Relative Risk (Control) ^d		0.770	1.340
Lower Limit Upper Limit		0.312 2.262	0.618 3.606
Weeks to First Observed Tumor	91	88	98
Adrenal: Cortical Adenoma or Cortical Carcinoma ^b	0/24(0.00)	3/50(0.06)	1/49(0.02)
P Values ^c	N.S.	N.S.	N.S.
Relative Risk (Control) ^d Lower Limit Upper Limit		Infinite 0.297 Infinite	Infinite 0.027 Infinite
Weeks to First Observed Tumor		106	106
Adrenal: Pheochromocytoma ^b	1/24(0.04)	0/50(0.00)	3/49(0.06)
P Values ^c	N.S.	N.S.	N.S.
Relative Risk (Control) ^d Lower Limit Unner Limit		0.000 0.000 8.966	1.469 0.127 75.534
Weeks to First Observed Tumor	110		106

TOPOGRAPHY : MORPHOLOGY	CONTROL	LOW DOSE	HIGH DOSE
Thyroid: C-Cell Carcinoma ^b	1/21(0.05)	5/49(0.10)	1/48(0.02)
P Values ^c	N.S.	N.S.	N.S.
Relative Risk (Control) ^d		2.143	0.438
Lower Limit Upper Limit		0.266 99.147	0.006 33.659
Weeks to First Observed Tumor	109	106	106
Thyroid: C-Cell Adenoma or C-Cell			
Carcinoma ^D	1/21(0.05)	12/49(0.24)	4/48(0.08)
P Values ^c	N.S.	P = 0.046	N.S.
Departure from Linear Trend ^e	P = 0.009	-	1
Relative Risk (Control) ^d		5.143	1.750
Lower Limit		0.855	0.192
Upper Limit		215.370	83.548
Weeks to First Observed Tumor	109	104	103
Thyroid: Papillary Carcinoma, Follicular-	1		
Cell Carcinoma, or Fapillary Cystadenocarcinoma NOS ^B	1/21(0.05)	1/49(0.02)	3/48(0.06)
P Values ^c	N.S.	N.S.	N.S.
Relative Risk (Control) ^d	!	0.429	1.313
Lower Limit		0.006	0.115
Upper Limit		32.983	67.452
Weeks to First Observed Tumor	110	106	66

TOPOGRAPHY: MORPHOLOGY	CONTROL	LOW DOSE	HIGH DOSE
Thyroid: Papillary Carcinoma, Follicular- Cell Carcinoma, Papillary Cystadenocar- cinoma NOS, or Papillary Cystadenoma ^b	1/21(0.05)	2/49(0.04)	4/48(0.08)
P Values ^c	N.S.	N.S.	N.S.
Relative Risk (Control) ^d Lower Limit Upper Limit		0.857 0.648 49.555	1.750 0.191 84.310
Weeks to First Observed Tumor	110	106	81
Mammary Gland: Fibroadenoma ^b P Values ^C	4/24(0.17) N S	5/50(0.10) N S	13/50(0.26) N.S
Relative Risk (Control) ^d Lower Limit Upper Limit		0.600 0.145 2.812	1.560 0.556 6.019
Weeks to First Observed Tumor	109	102	98
Mammary Gland: Fibroadenoma, Adenocar- cinoma NOS, or Papillary Adenocarcinoma	4/24(0.17)	5/50(0.10)	14/50(0.28)
P Values	N.S.	N.S.	N.S.
Relative Risk (Control) ^d Lower Limit Upper Limit		0.600 0.145 2.807	1.680 0.609 6.412
Weeks to First Observed Tumor	109	102	98

LOW CONTROL DOSE	1/24(0.04) 3/50(0.06)	N.S. N.S.	1.440 0.125	75.487	. 110 106	or 1/24(0.04) 3/50(0.06)	P = 0.003 N.S.	1.440 0.125	74.077	110 106	Polyp ^b 2/24(0.08) 14/49(0.29)	P = 0.003 $P = 0.043$	3.429 0.892	29.588	102 88
TOPOGRAPHY : MORPHOLOGY	Clitoral Gland: Carcinoma NO	P Values ^c	Relative Risk (Control) ^d Tower Limit	Upper Limit	Weeks to First Observed Tumor	Clitoral Gland: Adenoma NOS (Carcinoma NOS ^b	P Values ^c	Relative Risk (Control) ^d Lower Limit	Upper Limit	Weeks to First Observed Tumor	Uterus: Endometrial Stromal I	P Values ^c	Relative Risk (Control) ^d Lower Limit	Upper Limit	Weeks to First Observed Tumor

TABLI	E 4 (CONCLUDED)		
TOPOGRAPHY:MORPHOLOGY	CONTROL	LOW DOSE	HIGH DOSE
Uterus and Endometrium: Adenocarcinoma NOS ^b	1/24(0.04)	2/49(0.04)	4/48(0.08)
P Values ^c	N.S.	N.S.	N.S.
Relative Risk (Control) ^d Lower Limit Unner Limit		0.980 0.054 56 627	2.000 0.216 96 367
Weeks to First Observed Tumor	110	104	106
Zymbal's Gland: Sebaceous Adenocar- cínoma ^b P Values ^C	0/24(0.00) N.S.	0/50(0.00) N.S.	3/50(0.06) N.S.
Relative Risk (Control) ^d Lower Limit Upper Limit			Infinite 0.296 Infinite
Weeks to First Observed Tumor			89
^a Treated groups received doses of 0.05 or ^b Number of tumor-bearing animals/number o ^c The probability level for the Cochran-Arr the control group when P < 0.05; otherwis level for the Fisher exact test for the given beneath the incidence of tumors in icant (N.S.) is indicated. For both Coch tion (N) indicates a lower incidence in the arrest of the relation	0.10 percent in f f animals examined nitage test is giv se, not significan comparison of a tr the treated group nran-Armitage and the treated group	eed. l at site (proport en beneath the in (t (N.S.) is indic eated group with when P < 0.05; o Fisher exact test (s) than in the co	<pre>ion). cidence of tumors in ated. The probabilit the control group is therwise, not signif- s a negative designa- itrol group.</pre>
דווב אוי ריוב דבדמרי -	דאב דאר ארד בווב רד	eared group to the	control group.

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

every type of tumor in either sex where at least two such tumors were observed in at least one of the control or 1,5-naphthalenediaminedosed groups and where such tumors were observed in at least 5 percent of the group.

For female rats an increased incidence of endometrial stromal polyps was observed in both the high and low dose groups compared to the control group. The Cochran-Armitage test indicated a significant (P = 0.003) positive association between compound administration and tumor incidence. The Fisher exact tests supported this result with a significant (P = 0.003) comparison of the high dose group to the control; for the low dose comparison the probability level was P = 0.043, a marginal result which was not significant under the Bonferroni criterion. Based on these results, the administration of 1,5-naphthalenediamine was associated with an elevated incidence of endometrial stromal polyps in female rats.

A number of adenomas NOS and carcinomas NOS of the clitoral gland were observed in female rats. The Cochran-Armitage test indicated a significant (P = 0.003) positive association between dose and the combined incidence of adenomas NOS or carcinomas NOS of the clitoral gland. The Fisher exact test comparing high dose to control was also significant (P = 0.021). In historical data collected by this laboratory for the NCI Carcinogenesis Testing Program, 4/249 (2 percent) of the untreated female Fischer 344 rats had one of these tumors, compared to the 13/50 (26 percent) observed in the high dose group in

this bioassay. Based upon these statistical results, the administration of 1,5-naphthalenediamine was associated with an elevated incidence of clitoral gland neoplasms in female rats.

For females the Fisher exact test comparing control to low dose for the combined incidence of C-cell adenomas or C-cell carcinomas of the thyroid had a probability level of P = 0.046, a marginal result which was not significant under the Bonferroni criterion.

Based on these statistical tests, it is concluded that 1,5-naphthalenediamine was carcinogenic for female rats, producing tumors of the clitoral gland and uterus.

IV. CHRONIC TESTING RESULTS: MICE

A. Body Weights and Clinical Observations

Mean body weight depression was readily apparent in dosed male mice when compared to controls. A similar but less pronounced trend was evident in dosed females (Figure 4).

One low dose male had a soft subcutaneous mass on the leg and two males in this group had palpable abdominal masses. Firm nodular growths developed in one low dose male and two high dose females. Alopecia was observed in 27 control males, 16 low dose males, 4 high dose males, 25 control females, and 3 low dose females. Two low dose and two high dose males experienced noticeable swelling of the eyes. Abdominal distention was observed in one control male and one control female mouse.

B. Survival

The estimated probabilities of survival for male and female mice in the control and 1,5-naphthalenediamine-dosed groups are shown in Figure 5. There was no significant positive association between dosage and mortality for either male or female mice.

Adequate numbers of male mice were at risk from late-developing tumors with 58 percent (29/50) of the high dose, 78 percent (39/50) of the low dose and 66 percent (33/50) of the controls surviving on test until the termination of the study. The 6 control male mice that died in week 11 were autolyzed, as were 2 of the 4 high dose male mice that died in week 41.





FIGURE 4 GROWTH CURVES FOR 1,5-NAPHTHALENEDIAMINE CHRONIC STUDY MICE



FIGURE 5 SURVIVAL COMPARISONS OF 1,5-NAPHTHALENEDIAMINE CHRONIC STUDY MICE

For female mice, with 68 percent (34/50) of the high dose, 82 percent (41/50) of the low dose and 60 percent (30/50) of the control mice surviving on test until the termination of the study, adequate numbers were at risk from late-developing tumors.

C. Pathology

Histopathologic findings on neoplasms in mice are summarized in Appendix B (Tables B1 and B2); findings on nonneoplastic lesions are summarized in Appendix D (Tables D1 and D2).

Dietary administration of 1,5-naphthalenediamine produced an increase in hepatocellular neoplasms in female mice, and it produced a dose-related increase in thyroid neoplasms and compound-related nonneoplastic thyroid lesions in both sexes. The compound-related lesions are summarized below:

		MALES	;	F	EMALES	
	Con-	Low	High	Con-	Low	High
	<u>trol</u>	Dose	Dose	<u>trol</u>	Dose	Dose
LIVER						
(Number of animals with						
tissues examined histo-						
pathologically)	(39)	(45)	(43)	(46)	(49)	(46)
Hepatocellular Carcinoma	12	10	7	1	25	16
Hepatocellular Adenoma	0	3	6	0	3	11
THYROID						
(Number of animals with						
tissues examined histo-						
pathologically)	(38)	(46)	(43)	(44)	(49)	(45)
Follicular-Cell Adenoma						
(Papillary or Follicular-Cell						
Adenoma, Papillary						
Cystadenoma)	0	8	16	2	17	14
Follicular-Cell Carcinoma	0	1	1	2	0	1
Follicular-Cell Hyperplasia	2	12	9	2	1	4
C-Cell Adenoma	0	2	0	0	1	2
C-Cell Carcinoma	0	0	4	0	1	6

In male mice, dietary administration of the compound did not increase the incidence of hepatocellular neoplasms, whereas dosed females showed a striking increase in hepatocellular carcinomas and hepatocellular adenomas.

Grossly, hepatocellular neoplasms appeared as smooth, nodular, rounded masses distorting the normal shape of the liver. Color varied, many neoplasms appearing pale tan or dark red. Microscopically, hepatocellular carcinomas were expansive masses of hepatocytes exhibiting loss of normal architectural pattern, the cells being arranged in sheets or trabeculae instead of the normal lobules. Nuclei were frequently uniform, although variable amounts of pleomorphism did occur. The cytoplasm was either basophilic or acidophilic, sometimes varying from one region of the tumor to another, and was frequently pale. Lesions classified as hepatocellular adenomas were smaller, usually better differentiated, and were less pleomorphic than the hepatocellular carcinomas.

The criteria for classification of thyroid neoplasms in mice were the same as those used to classify thyroid neoplasms in rats. The nonneoplastic thyroid lesions found in dosed mice were similar to those in the rats but occurred in higher incidences. Hyperplasia of follicular cells (focal, papillary or adenomatous) were found in 2/38 (5 percent) control, 12/46 (26 percent) low dose, and 9/43 (21 percent) high dose male mice. Abundant golden brown pigment was seen in follicular epithelium, colloid, and macrophages. In the mice,

there were frequent foci of lymphocytes in the thyroid parenchyma and occasional cystic areas filled with amorphous material containing long clefts suggesting cholesterol crystals.

Three transitional-cell papillomas occurred in the bladder or urethra of dosed mice (two high dose males and one high dose female), but none occurred in controls.

Based upon the results of this pathologic examination, 1,5naphthalenediamine was carcinogenic to B6C3F1 mice, producing hepatocellular neoplasms in females and thyroid neoplasms in both sexes.

D. Statistical Analyses of Results

The results of the statistical analyses of tumor incidence in mice are summarized in Tables 5 and 6. The analysis is included for every type of tumor in either sex where at least two such tumors were observed in at least one of the control or 1,5-naphthalenediamine-dosed groups and where such tumors were observed in at least 5 percent of the group.

For both male and female mice elevated incidences of thyroid tumors were observed in the dosed groups. In female mice the Cochran-Armitage test indicated a significant (P = 0.005) positive association between dietary concentration and the incidence of C-cell carcinomas. This was supported by a significant (P = 0.014) Fisher exact test for the high dose group. For males the Cochran-Armitage test result was also significant (P = 0.017), but the Fisher exact tests were

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SPECIFIC SITES IN MALE MICE TREATED WITH 1,5-NAPHTHALENEDIAMINE^a ANALYSES OF THE INCIDENCE OF PRIMARY TUMORS AT

1			
TOPOGRAPHY : MORPHOLOGY	CONTROL	LOW DOSE	HIGH DOSE
Lung: Alveolar/Bronchiolar Carcinoma ^b	2/39(0.05)	3/46(0.07)	0/45(0.00)
P Values ^c	N.S.	N.S.	N.S.
Relative Risk (Control) ^d Lower Limit Upper Limit		1.272 0.153 14.686	0.000 0.000 4.478
Weeks to First Observed Tumor	109	82	
Lung: Alveolar/Bronchiolar Adenoma or Alveolar/Bronchiolar Carcinoma ^b	4/39(0.10)	9/46(0.20)	2/45(0.04)
P Values ^c	N.S.	N.S.	N.S.
Departure from Linear Trend ^e	P = 0.037	1	
Relative Risk (Control) ^d Lower Limit		1.908 0.582	0.433 0.041
Upper Limit	!	7.882	2.871
Weeks to First Observed Tumor	109	82	105
Hematopoietic System: Malignant Lymphoma ^b	13/39(0.33)	14/47(0.30)	5/49(0.10)
P Values ^c	P = 0.007(N)	N.S.	P = 0.008(N)
Relative Risk (Control) ^d Lower Limit Upper Limit		0.894 0.448 1.817	0.306 0.094 0.829
Weeks to First Observed Tumor	100	82	95

		TOW	HIGH
TOPOGRAPHY : MORPHOLOGY	CONTROL	DOSE	DOSE
Thyroid: C-Cell Carcinoma or C-Cell Adenoma ^b	0/38(0.00)	2/46(0.04)	4/43(0.09)
P Values ^c	P = 0.044	N.S.	N.S.
Relative Risk (Control) ^d Lower Limit Upper Limit		Infinite 0.246 Infinite	Infinite 0.825 Infinite
Weeks to First Observed Tumor	1	105	105
Thyroid: Papillary Adenoma, Follicular- Cell Adenoma, or Papillary Cystadenoma NOS ^b	0/38(0.00)	8/46(0.17)	16/43(0.37)
P Values ^c	P < 0.001	P = 0.006	P < 0.001
Relative Risk (Control) ^d Lewer Limit Upper Limit	1 1 1	Infinite 1.905 Infinite	Infinite 4.523 Infinite
Weeks to First Observed Tumor		105	80,
^a Treated groups received doses of 0.1 or 0	.2 percent in fee	.b4	
^D Number of tumor-bearing animals/number of	animals examined	l at site (proportio	n).
^c The probability level for the Cochran-Arm	itage test is giv	ren beneath the inci	dence of tumors in

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TABLE 5 (CONCLUDED)

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

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

HIGH DOSE	3/46(0.07)	N.S.	Infinite 0.638 Infinite	91	5/46(0.11)	P = 0.024		Infinite 1.347	Infíníte	91	5/46(0.11)	P = 0.045(N)		0.410	0.124 1.117	105
LOW DOSE	1/48(0.02)	N.S.	Infinite 0.055 Infinite	89	10/48(0.21)	P = 0.001		Infinite 3.037	Infinite	89	19/50(0.38)	N.S.		1.432	0./60 2.781	63
CONTROL	0/49(0.00)	N.S.			0/49(0.00)	N.S.	P = 0.005				13/49(0.27)	N.S.	P = 0.011			57
TOPOGRAPHY : MORPHOLOGY	Lung: Alveolar/Bronchiolar Carcinoma ^b	P Values ^c	Relative Risk (Control) ^d Lower Limit Upper Limit	Weeks to First Observed Tumor	<pre>by Lung: Alveolar/Bronchiolar Adenoma or Alveolar/Bronchiolar Carcinoma^b</pre>	P Values ^c	Departure from Linear Trend ^e	Relative Risk (Control) ^d Lower Limit	Upper Limit	Weeks to First Observed Tumor	Hematopoietic System: Leukemia or Malignant Lymphoma ^b	P Values ^c	Departure from Linear Trend ^e	Relative Risk (Control) ^d	Lower Limit Upper Limit	Weeks to First Observed Tumor

ANALYSES OF THE INCIDENCE OF PRIMARY TUMORS AT SPECIFIC SITES IN FEMALE MICE TREATED WITH 1,5-NAPHTHALENEDIAMINE^a

TABLE 6

TOPOGRAPHY : MORPHOLOGY	CONTROL	LOW DOSE	HIGH DOSE
Liver: Hepatocellular Carcinoma ^b	1/46(0.02)	25/49(0.51)	16/46(0.35)
P Values ^C	P = 0.001	P < 0.001	P < 0.001
Departure from Linear Trend ^e	P < 0.001	-	
Relative Risk (Control) ^d Lower Limit		23.469 4.156	16.000 2.683
Upper Limit	8	906.346	646.516
Weeks to First Observed Tumor	109	74	66
Liver: Hepatocellular Adenoma or Hepatocellular Carcinoma ^b	1/46(0.02)	28/49(0.57)	27/46(0.59)
P Values ^C	P < 0.001	P < 0.001	P < 0.001
Departure from Linear Trend ^e	P = 0.002	1	!
Relative Risk (Control) ^d Lower Limit		26.286 4.741	27.000 4.874
Upper Limit	-	1030.801	1027.943
Weeks to First Observed Tumor	109	74	66
Stomach: Squamous-Cell Papilloma ^b	0/41(0.00)	3/47(0.06)	0/46(0.00)
P Values ^c	N.S.	N.S.	N.S.
Departure from Linear Trend ^e	P = 0.017	1	5
Relative Risk (Control) ^d	-	Infinite	
Upper Limit		U.J.27 Infinite	
Weeks to First Observed Tumor	-	105	

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TOPOGRAPHY : MORPHOLOGY	CONTROL	DOSE	HIGH DOSE
Pituitary: Adenoma NOS, Chromophohe Adenoma or Acidophil Adenoma ^b	3/34(0.09)	4/35(0.11)	1/30(0.03)
P Values ^C	N.S.	N.S.	N.S.
Relative Risk (Control) ^d	-	1.295	0.378
Lower Limit Upper Limit		0.238 8.188	0.007 4.424
Weeks to First Observed Tumor	109	105	106
Adrenal: Pheochromocytoma ^b	3/46(0.07)	0/44(0.00)	0/44(0.00)
P Values ^C	P = 0.040(N)	N.S.	N.S.
Relative Risk (Control) ^d		0.000	0.000
Lower Limit		0.000	0.000
Upper Limit		1.731	1.731
Weeks to First Observed Tumor	68		
Thyroid: C-Cell Carcinoma ^b	0/44(0.00)	1/49(0.02)	6/45(0.13)
P Values ^C	P = 0.005	N.S.	P = 0.014
Relative Risk (Control) ^d		Infinite	Infinite
Lower Limit	-	0.048	1.574
Upper Limit		Infinite	Infinite
Weeks to First Observed Tumor		105	105

TOPOGRAPHY : MORPHOLOGY	CONTROL.	LOW DOSE	HIGH DOSE
Thyroid: C-Cell Adenoma or C-Cell Carcinoma ^b	0/44(0.00)	2/49(0.04)	8/45(0.18)
P Values ^c	P = 0.001	N.S.	P = 0.003
Relative Risk (Control) ^d Lower Limit		Infinite 0.267	Infinite 2.250
Upper Limit		Infinite	Infinite
Weeks to First Cbserved Tumor		105	41
Thyroid: Papillary Adenoma, Follicular-Ce Adenoma, or Papillary Cystadenoma NOS ^b	.11 2/44(0.05)	17/49(0.35)	14/45(0.31)
P Values ^C	F = 0.003	P < 0.001	P = 0.001
Departure frcm Linear Trend ^e	P = 0.025	-	
Relative Risk (Control) ^d		7.633	6.844
Lower Limit Upper Limit		1.9/1 64.662	L./U9 58.827
Weeks to First Observed Tumor	80	105	91
^a Treated groups received doses of 0.1 or 0	.2 percent in f	sed.	
b Nimber of timor-bearing animale/number of	animale evamine	d at site (proporti	(uu)

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TABLE 6 (CONCLUDED)

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given beneath the incidence of tumors in the treated group when P < 0.05; otherwise, not signifithe control group when P < 0.05; otherwise, not significant (N.S.) is indicated. The probability level for the Fisher exact test for the comparison of a treated group with the control group is ^cThe probability level for the Cochran-Armitage test is given beneath the incidence of tumors in cant (N.S.) is indicated. For both Cochran-Armitage and Fisher exact tests a negative designa-^dThe 95% confidence interval on the relative risk of the treated group to the control group. tion (N) indicates a lower incidence in the treated group(s) than in the control group.

^eThe probability level of the test for departure from linear trend is given beneath the control group when P < 0.05 not. When incidences were combined so that the numerator represented mice with either a papillary adenoma, a follicular-cell adenoma, or a papillary cystadenoma of the thyroid, the Cochran-Armitage test indicated a significant positive association between dietary concentration and tumor incidence for both males (P < 0.001) and females (P = 0.003). These were supported by significant ($P \leq 0.006$) Fisher exact test results in each sex for comparisons of each dosed group to the control group. Based on these results, the administration of 1,5-naphthalenediamine was associated with the incidence of thyroid neoplasms in both male and female mice.

For females an increased incidence of hepatocellular carcinomas was also observed among the dosed mice. The Cochran-Armitage test indicated a significant (P = 0.001) positive association between dose and incidence. This was supported by significant (P < 0.001) comparisons of both the high and low dose to the control group using the Fisher exact test. Based on these results the administration of 1,5-naphthalenediamine was associated with the incidence of hepatocellular carcinomas in female mice.

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For female mice, when the incidence of alveolar/bronchiolar adenomas and alveolar/bronchiolar carcinomas were combined, an increased incidence in the dosed groups was noted. The Fisher exact test was significant for both the high (P = 0.024) and low (P = 0.001) dose groups. The departure from linear trend was significant since tumor incidence was increased more in the low dose than in the high

dose group. In historical control data compiled by this laboratory for the NCI Carcinogenesis Testing Program, 17/275 (6 percent) of the untreated female B6C3F1 mice had an alveolar/bronchiolar neoplasm. Based upon these results the administration of 1,5-naphthalenediamine was associated with the incidence of alveolar/bronchiolar neoplasms in female mice.

For females the Fisher exact test comparing the incidence of leukemia or malignant lymphoma in high dose mice with that in the controls had a probability level in the negative direction of P = 0.045, a marginal result which was not significant under the Bonferroni criterion.

Also for females the Cochran-Armitage test showed a significant (P = 0.040) negative association between dose and the incidence of adrenal pheochromocytomas, but the Fisher exact tests were not significant.

In male mice the possibility of a negative association between dose and the incidence of malignant lymphomas or leukemia was noted.

Based upon these statistical results the administration of 1,5naphthalenediamine was associated with the increased incidence of thyroid neoplasms in male mice and of thyroid neoplasms, of hepatocellular carcinomas, and of alveolar/bronchiolar neoplasms in female mice.

V. DISCUSSION

There were no significant positive associations between dietary concentrations of 1,5-naphthalenediamine and mortality in either sex of rats or mice. In all groups adequate numbers of animals survived sufficiently long to be at risk from late-developing tumors.

Several uterine neoplasms occurred in dosed female rats at higher incidences than in corresponding controls. There was a significant positive association between dietary concentration of the compound and the incidences of endometrial stromal polyps in female rats. In addition, the high dose to control Fisher exact comparison was significant. Endometrial stromal sarcomas were observed in two low dose and two high dose female rats, but not in controls. Uterine adenocarcinomas occurred at a higher incidence in the high dose female rat group than in the control group, but the difference in tumor incidence was not statistically significant.

The administration of 1,5-naphthalenediamine was associated with an elevated incidence of clitoral gland neoplasms in female rats. There was a significant positive association between the concentration of the chemical added to the diet and the incidence of either adenomas or carcinomas of the clitoral gland in female rats. The incidence of either of these neoplasms in the high dose female rat group was significant relative to the incidence in the control group.

Elevated incidences of thyroid neoplasms were observed among dosed mice. For mice of both sexes there were significant positive

associations between dietary concentration of 1,5-naphthalenediamine and the incidences of thyroid C-cell carcinomas. For the females the high dose to control Fisher exact comparison supported the finding; this was not true for males. When the mice were grouped so that the numerator of the incidence represented those animals with a papillary adenoma, a follicular-cell adenoma, or a papillary cystadenoma of the thyroid, the Cochran-Armitage test was significantly positive for both males and females and all the Fisher exact comparisons supported the findings.

The incidence of hepatocellular carcinomas in female mice was significantly associated with increased concentration of 1,5-naphthalenediamine. In addition, the high dose to control and the low dose to control Fisher exact comparisons were significant. The incidence of alveolar/bronchiolar adenomas was significant, relative to controls, in both the low dose and the high dose female mouse groups.

Under the conditions of this bioassay, 1,5-naphthalenediamine was carcinogenic in female Fischer 344 rats, causing clitoral and uterine neoplasms. 1,5-Naphthalenediamine was also carcinogenic for B6C3F1 mice, producing thyroid neoplasms in males and neoplasms of the thyroid, liver, and lung in females. Insufficient evidence was provided for the carcinogenicity of the compound in male Fischer 344 rats.

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APPENDIX A

SUMMARY OF THE INCIDENCE OF NEOPLASMS IN RATS TREATED WITH 1,5-NAPHTHALENEDIAMINE

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TABLE AI SUMMARY OF THE INCIDENCE OF NEOPLASMS IN MALE RATS TREATED WITH 1,5-NAPHTHALENEDIAMINE

	CONTROL (UNTR) 01-0330	LOW DOSE 01-0280	HIGH DOSE 01-0285
ANIMALS INITIALLY IN STUDY ANIMALS NECROPSIED ANIMALS EXAMINED HISTOPATHOLOGICALLY**	25 25 4 25	50 49 49	50 50 49
INTEGUMENTARY SYSTEM			
*SKIN SOUAMOUS CELL PAPILICMA SEBACEOUS ADENOCARCINOMA FIBPOUS HISTIOCYTOMA	(25) 2 (8%)	(49) 1 (2%) 1 (2%) 1 (2%)	(50) 1 (2%)
*SUBCUT TISSUE PTBROMA LIPOMA	(25) 1 (4%)	(49) 3 (6%)	(50) 2 (4%) 1 (2%)
RESPIRATORY SYSTEM			
*LARYNX Papiilcma, Nos	(25)	(4 9)	(50) 1 (2 %)
#TRACHEA P&PILLOM&, NOS	(24)	(16)	(13) 1 (8%)
*LUNG ADENOCARCINOMA, NOS, METASTATIC ALVEOLAF/BRONCHIOLAR ADENOMA ALVFOLAR/BRONCHIOLAR CARCINOMA C-CELL CARCINOMA, METASTATIC SFB4CFOUS ADENOCARCINOMA, METAST PHEOCHROMOCYTOMA, METASTATIC	(25) 1 (4%)	(49) 1 (2%) 2 (4%) 1 (2%)	(47) 1 (2%) 4 (9%) 1 (2%)
HEMATOPOIETIC SYSTEM			
*MULTIPLE ORGANS LFUKEMIA,NOS UNDIFERRENTIATED LEUKEMIA	(25)	(49) 10 (20%)	(50) ⁻ 1 (2%) 4 (8%)
MYELONONCYTIC LEUKEMIA LYMPHOCYTIC LEUKEMIA	1 (4%)		1 (2%) 2 (4%)

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

	CONTROL (UNTR) 01-0330	ICW DOSE 01-0280	HIGH DOSE 01-0285
<pre>#MANDIBULAP L. NODE C-CELL CANCINOMA, METASTATIC SEBACEOUS ADENOCARCINCMA, METAST NEUPILEMOMA, METASTATIC</pre>	(24) 1 (4%)	(47) 1 (2%)	(47) 1 (2%)
#MEDIASTINAL L.NODE SEBACEOUS ADENOCARCINOMA, METAST	(24)	(47) 1 (2%)	(47)
*MESENTERIC L. NODE Lymphangiona	(24)	(47)	(47) 1 (2 %)
*LIVEP UNCIPPERENTIATED LEUKEMIA	(25)	(49)	(49) 1 (2%)
*THYMUS THYMOMA MALIG.LYMPHOMA, LYMPHOCYTIC TYPE	(13)	(35)	(36) 1 (3%) 1 (3%)
CIFCULATORY SYSTEM			
NONF			
UIGESTIVE SYSTEM			
*SPIIVARY GLAND ADENOCARCINOMA, NOS FIBPOSAPCOMA NEURILEMOMA, MALIGNANT	(25)	(47) 1 (2%)	(46) 1 (2%) 1 (2%)
*LIVER NEOPLASTIC NODULE HEPAIOCELLULAR CAPCINOMA LYMPHANGIOMA	(25) 1 (4%)	(49) 3 (6%) 4 (8%)	(49) 2 (4%) 2 (4%) 1 (2%)
*STOMACH SQUAMOUS CELL PAPILLOMA	(24)	(47) 1 (2%)	(47)
UFINARY SYSTEM			
*KIDNEY LIPOMS	(25)	(49) 1 (2 %)	(48)
*URINAPY ELADDER TFANSITIONAL-CELL CARCINOMA	(25)	(49)	(48)
A NUMBER OF ANTMALS WITH TISSUE FYAM	INED MICROSCOPIC	ALLY	

* NUMBER OF ANIMALS WITH TISSUE * NUMBER OF ANIMALS NECROPSIED

	CONTROL (UNTR) 01-0330	LOW DOSE 01-0280	HIGH LOSE 01-0285
ENDOCFINE SYSTEM			
*PITUITAPY	(22)	(44)	(44)
CARCINOMA, NOS			1 (2%)
ADENOMA, NOS	2 (9%)	1 (2%)	1 (2%)
CHROMOPHOBE ADENOMA		3 (7%)	/ (16%)
ACIDOPHIL ADENOMA			1 (2%)
RCIDUPHIL CARCINOMA		2 1781	1 (2%)
DESUPHIL ADDIVINA TUMPDOMITINI-OPII MUMOD MEMICUN		5 (10)	2 (3%)
INTERSTITIAL-CELL TOROR, METASTA			1,2%)
*ADRENAL	(24)	(48)	(48)
CORTICAL ADENOMA	()	1 (2%)	
PHEOCHROMOCYTOMA	1 (4%)	3 (6%)	5 (10%)
PHEOCHROMOCYTOMA, MALIGNANT	1 (4%)	1 (2%)	• •
NEUPOBLASTOMA			1 (2%)
	(01)	< 11 T 1	< 11 T 1
*THYROID	(21)	(47)	(47)
FOLLICULAR-CELL ADENOMA	1 (5%)	0 (1) 7 .	1 (2%)
C-CELL ADENOMA	2 (10 %)	2 (4%)	5 (11%)
SERVICEOUS ADENOCARCINONA MEESCE	2 (1.7%)	5 (n7a) 1 (D7a)	3 (670)
DEDICLOUS FDENOCARCINONA, RELASI		, 270)	1 (28)
PREILLERT CISTADENOCRACINONA,NOS			* ,2%)
*PARATHYROID	(13)	(24)	(28)
ADENOMA, NOS	v - y	·- /	1 (4%)
*PANCREATIC ISLETS	(25)	(48)	(45)
ISLET-CELL ADENOMA	1 (4%)	1 (2%)	4 (9%)
ISLET-CELL CARCINOMA		1 (2%)	7 (2%)
REPRODUCTIVE SYSTEM			
*MAMMARY GLAND	(25)	(49)	(50)
ADENOCARCINOMA, NOS		4 (0/)	7 (2%)
FIBRUADENOMA		1 (2%)	
* PREPUTIAL GLAND	(25)	(49)	(50)
CARCINOMA.NOS	()	()	1 (2%)
ADENOMA, NOS			1 (2%)
*TESTIS	(25)	(49)	(49)
INTERSTITIAL-CELL TUMOR	21 (84%)	44 (90%)	45 (92%)
INTERSTITIAL-CELL TUMOR, MALIGNA			1.471

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

	CONTROL (ONTR) • 01-0330	LCW DCSE 01-0280	HIGH DOSE 01-0285
NERVOUS SYSTEM			
APRAIN CAPCINCMA, NCS, METASTATIC GITOMA, NOS	(25)	(49) 1 (2%)	(47) 1 (2%) 1 (2%)
#CEPEBFILNM GLIOMA, NOS	(25)	(49) 1 (2%)	(47)
SPECIAL SENSE OPGANS			
*EAR CANAL SQUAMOUS CELL CAPCINCHA	(25) 1 (4 %)	(49)	(50)
*ZYMBAL'S GIAND SEBACEOUS ADENOCARCINOMA	(25)	(49)	(50) 1 (2 %)
MUSCULOSKELETAL SYSTEM			
NONF			
EOLY CAVITIES			
* BODY CAVITIES	(25)	(49)	(50)
MESOTHELIOMA, NOS MESOTHELIOMA, MALIGNANT		1 (2%)	1 (2%)
*ABEOMINAL CAVITY CSTEOSAFCOMA	(25)	(49)	(50) 1 (2%)
ALL OTHEP SYSTEMS			
TAIL SOUAMOUS CELL PAPILICHA			1
NUMBER OF ANIMALS WITH TISSUE EX NUMBER OF ANIMALS NECECESTED	AMINED MICPOSCOPIC	ALLY	

TABLE AI (CONCLUDED)

	CONTROL (UNTR) 01-0330	LOW DOSE 01-0280	HIGH DOSE 01-0285
PHIMAL DISPOSITION SUMMARY			
ANTMALS INTUTALLY IN STUDY	25	50	50
NATURAL DEATHD	5	5	8
MORIBUND SACPIFICE	3	5	5
SCHEDULFD SACRIFICE			
TERMINAL SECRETICE	17	40	37
PNIMAL MISSING	• '		
I INCLUDES AUTOLYZED ANIMALS			
ELMOR SUMMARY			
TOTAL ANIMALS WITH PPIMARY TUMOPS*	21	47	49
TOTAL FRIMARY TUMORS	35	96 .	110
TOTAL ANIMALS WITH BENIGN TUMORS	21	46	49
TOTAL BENIGN TUMORS	29	67	87
TOTAL ANIMALS WITH MALIGNANT TUMOFS	5	19	24
TOTAL MALIGNANT TUMORS	5	25	30
TOTAL ANIMALS WITH SECONDARY TUMOPS#	2	1 /	5
TOTAL SECONDARY TUMORS	2	4	5
TOTAL ANIMALS WITH TUMORS UNCERTAIN-			
BENIGN OR MAIIGNENT	1	4	2
TOTAL UNCERTAIN TUMCES	1	4	2
TOTAL ANIMALS WITH TUMOPS UNCERTAIN-			
PFIMAPY OF METASTATIC			
TOTAL UNCERTAIN TUMORS			
PRIMARY TUMORS: FLL TUMORS EXCEPT SE	CONDARY TUMORS		
SECONDERY TUMORS: METASTATIC TUMORS	OR TUMORS INVA	STVE INTO AN AD	JACENT OFGAN

TABLE A2 SUMMARY OF THE INCIDENCE OF NEOPLASMS IN FEMALE RATS TREATED WITH 1,5-NAPHTHALENEDIAMINE

	CONTROL (UNTP) 02-0330	104 DOSE 02-0280	HIGH DOSE 02-0285
NIMALS INITIALLY IN STUDY	25	<u>د</u> ٥	÷0
NIMALS NECFO2SIED NIMALS EXAMINED HISTOFATHOLOGTCALLY	24 ** 24	50 50	50
NTEGUMENTARY SYSTEM			
*SKIN SQU&MOUS CELL C&PCINOMA	(24)	(50)	(50) 2 (4%)
FSPIPATORY SYSTEM			
TRACHEN PAPILLONN, NOS	(23)	(16)	(10) 1 (10%)
#LUNG ALVEOLAF/BPONCHIOLAR ADENOMA	(24)	(50) 1 (2%)	(50) 1 (2%)
ALVEDIAP/BPONCHIOLAR CARCINOMA COFFICAL CAPCINOMA, METASTAMIC		1 (2%)	1 (2%)
C-CELL CARCINOMA, METASTATIC ENDOMETRIAL STROMAL SARCOMA, MET		1 (2%) 1 (2%)	
EMATOPOITI'C SYSTEM			
MULTIPLE OPGANS	(24)	(50)	(50)
LEUKEMIA, NOS	((* *)	1 (2%)	4 (25)
MYFLOMONOSYTIC ISUKEMIA	2 (9%)	5 (12%)	1 (2%)
IRCULATORY SYSTEM			
NCNT			
IGESTIVE SYSIEM			
*LIVEP	(24)	(51)	(49)

** EXCLUDES PARTIALLY AUTOLYZED ANIMALS

	CONTFOL (UNTR) 02-0330	IOW DOSE 02-0280	HIGH ECSE 02-0285
HEPAINCEILUIAR CARCANCMA ENDOMEIPIAL SIPOMAL SARCOMA, INV		1 (2%) 1 (2%)	
*SIGMACH CAPCINOMA,NOS SQUAMDUS CELL PAPILIOMA ENDOMETRIAL STROMAL SAPCOMA, THV	(?u)	(5 ⁴) 1 (2¥)	(49) 1 (2종) 1 (2종)
UFINARY SYSTEM			
TIDCM, #KIDxEA	(24)	(5°) 1 (2%)	(49) 1 (?≪)
#KIDNEY/PELVIS Transitionel-Cell Fepilloma	(24)	(51)) 1 (2%)	(49) 1 (7%)
#UFTNAPY BLADDEP TRANSITIONAL-CTIL PAPILIONA TPANSITIONAL-CFLL CARCINOMA	(24) 1 (4%)	(48)	(4°) 1 (2%)
ENDOCRINE SYSTEM			
#PTTUITARY CAPCINOMA,NOS	(21)	(5-7)	(47) 1 (2%)
DEMCMA, NOS CHROMORROBE DENOME CHROMORHOBE CARCINOME ACTDORHIL ADEMOMA RESORATI ADEMOMA	주 (29%)	1 (2%) 7 (14考) 1 (2%) 1 (2%) 1 (2%)	1 (2%) 16 (34%)
PAPTLIARY CYSITDEMOCRECTNOMA, MET		, ,2%)	1 (2%)
#SDRENAL CORMICAL ADENOMA CORMICAL CARCINOMS	(۵۵)	(50) ? (4%) 1 (2%)	(49) 1 (2%)
FHEOCHFOMOCYTOMA LIPOMA	1 (4%)	1 (2%)	3 (6%)
*THYBOTD PARTILARY CARCINOME	(21)	(49)	(48) 1 (2%)
FOLLITULAR-CELL CAFCINOMA C-CELL ADANOMA C-CELL CARCINOMA	1 (5%)	7 (14%)	1 (2条) 3 (6条) 1 (2年)
PAPILLARY CYSIADENOMA, NOS 	· · · · · · · · · · · · · · · · · · ·	1 (23) 1 (2%)	1 (2%) 2 (4%)

NUMBER OF ANIMALS WITH TISSUE EXAMINED MICPOSCOPICALLY * NUMBER OF ANIMALS NUCROESTED

	CONTROL (UNIR) 02-1330	LOW DOSE 12-0280	HIGH DOSE 02-0285
*DINCREATIC ISLETS TRIET-CEIL CARCINOMA	(22) 1 (5考)	(u 9)	(47)
PEPPODUCTIVE SYSTEM			
• MIMMAPY GLAND ADEMCHA, NOS ADEMORAPCINOMA, MOS PARTILARY ADEMORAPCINOMA	(24)	(5 ⁰) 1 (2%) 1 (2%)	(FO) 1 (25) 1 (25) 1 (25)
INTRADUCIAL PARTILOMA		· · · · · · ·	1 (2%)
FIPROIDENOMA	u (17ま)	5 (10%)	13 (26%)
*CLITOFAL GLAND CIECTNONA,NOS ADENONA, JOS	1 (4者) (24)	(50) 3 (6%)	(50) 8 (165) 5 (103)
* Um pe U S	(24)	(49)	(48)
ADENOCARCINOMA, NOS Endometrial sipomai polyp Endometriai sipomai s/RCOMA	1 (4号) 2 (8考) 1 (4号)	14 (29%) 2 (4%)	1 (2%) 20 (42%) 2 (4%)
#UTTPUS/INTCATTPIUM BTENOCITOMA, MOS	(24)	(49) ? (48)	(48) २ (6इ)
SESWOIDEF-CEIT INWOD CEFNOIDEF-CEIT INVOD ROAFBA	(24)	(49) 1 (2%) 1 (2%)	(to)
NERVOUS SYSTEM			
*BB*IM CHPOMORHOBE CATCINCMA, INVASIVE CLIOM*, NOS	(23)	(57) 1 (2巻) 1 (2巻)	(「り) 1 (2年)
CEECTIT REAGE LOCANS			
HIRDERINN GLIND ADENCIARCINOMA, NOS	(24)	(57)	(50) 1 (2)
SONARUAS CAIT CASCOMB	(24) 1 (4%)	(50)	(50)
NZYMPALIS GLAND SEBACFOUS ADENOCRECTNOMA	(24)	(50)	(50) 3 (63)
# NUMPER OF ANIMALS WITH TISSUE EXAM	THED MICPOSCOPIC	2 LLY	

* NUMPER OF ANIMALS NECESSIED

	CONTROL (UNTR) 02-0330	LOW DOSE 02-0380	HIGH DOSE 02-0285
MUSCHLOSKELETAL SYSTEM			
исир			
BODY CAVILLES			
FOLY CAVITIES MESOTHRIIDMA, MAITGNANT	(?4)	(50) 1 (2%)	(50)
*ABROMINAL CAVITY LEICMYOSANCOM?	(24) 1 (4考)	(50)	(
*PERITONEUM ENDOMPTEIAL STROMEL SARCOMA, MET	(24)	(57) 1 (28)	(źŋ)
ALL OTHER SYSTEMS			
TATL SQUEMOUS JELL PAPTILOME		1	
DIAPHPAGM ENDOMETRIAL STROMAL SARCOMA, MET		1	
PNIMAL DISPOSITION SUMMARY			
ANIMALS INTETALLY IN STUDY	זק	Ξ0	50
NATURAL DEATHD	4	<i>ב</i> ز	5
MORIBUND SACRIFICE SCHEDUIED SACRIFICE ACCIDENTALLY KIUED	د.	7	٦
TERMINAL SACRIFICE ANIMAL MISSING	16	39	38
3_INCLUDES_AUCOLYZED ANIMALS.			

* NUMBER OF ANIMALS WITH TISSUE EXAMINED MICROSCOPICALLY • NUMBER OF ANIMALS NRCECESIED

	00Nº HEL (0438) 02-0330	12-0280	HIGH DOSE 02-0285
uider suidderk			
IOLII BEINIDA LANDEC IOLII VAIMUTZ MIWH BEINIDA IAMUEZ+	17 25	4 1 76	49 107
αθωρή βενίαν ύμφοθό μομρή γηματό Μήμκ βενιάν ΦυΝθέδ	10 14	33 ЦЕ	44 70
TOTAL ANIMALS WITH ANTIGNANT TUMOPS TOTAL ANIMALS WITH ANTIGNANT TUMOPS	10 11	22 26	33 5e
TOTAL ANIMALS WITH RECONDARY TUMORS		5 8	1 1
AUDIT DACEDUIL SUNCES BENICA UN MUTICALA DUMOES UNCEEDILA DOULT FAILAITE MULH DUMOES UNCEEDILA.		ц ц	ц ц
TOTAL ANTHALS WITH TUMORS UNCERTAIN- PRIMARY OF METASTRUTH			

SECONDARY TUNDES: METISTATIO TUMOES OR TUMORS INVASIVE INTO AN ADJACENT OFGAN

APPENDIX B

SUMMARY OF THE INCIDENCE OF NEOPLASMS IN MICE TREATED WITH 1,5-NAPHTHALENEDIAMINE





TABLE BI SUMMARY OF THE INCIDENCE OF NEOPLASMS IN MALE MICE TREATED WITH 1,5-NAPHTHALENEDIAMINE

	CONTPOL (UNTR) 05-0330	10W DOSE 05-0285	05-0200 05-0200
ANIMAIS INITIALLY IN STUDY ANIMALS MISSING ANIMALS NECTOPSIED ANIMALS EXAMINED HISTOPAINCLOGICALLY**	50 2 39 39	50 1 47 47	50 49 46
INTEGRMENTARY SYSTEM			
*SURCUT TISSUE FIBROUS HISTIOCYTOMA	(3º)	(47) 1 (2%)	(49)
FESPTRATORY SYSTEM			
<pre>#IMMG #FPATOCTILUIA® CAPOTNOMA, MFTAST %IVFOLAF/BFONCHIOLAR /DENOMA %IVFOLAF/BFONCHIOLAP CARCINGMA C-JPLL CARCINOMA, METISTATIC</pre>	(30) 2 (5系) 2 (5系) 2 (5系)	(46) 2 (4%) 6 (13%) 3 (7%)	(45) 2 (4명) 1 (2명)
HEMATOPOIRTIC SYSTEM			
-MULTIPLE OFGINS MILIGNANT LYMPHOMI, NCS MILIGLLYMPHOMI, LYMPHOCYTIC TYPE MALIGLLYMPHOMI, HISTIOCYTIC TYPE MALIGNANT LYMPHOMI, MIXED TYPE	(39) 11 (28%)	、47) 1 (2%) 3 (6%) 2 (6然) 5 (11%)	(49) 1 (2%) 1 (2%)
*SPLEPN HEMANGIOSASCOMA MAITGNANT LYPHOMA, NOS MALIGNANT LYMPHOMA, MIXED TYPE	(38) 1 (3%)	(45) 1 (2%)	(41) 2 (5努)
#MESENTETIC L. NODE LYMPHANGIOMA M/IIG.IYMPHOMA, HISTIOCYTIC TYPE MALIGNAMI LYMPHONA, MIXEDITYPE	(36) 1 (2%)	(43) 2 (5%)	(35) 1 (38)
#TFYMUS MALTGNANT LYMPHONA, NOS	(13)	(29)	(16) ⁻¹ (6葉)

NONE

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

TABLE B1 (CONTINUED)

	02-0330 О́лшеОГ (ЛИШе)	100 DOSE 05-0285	HIGH DOSE 05-02°0
LIGESTIVE SYSDEM			
HEDYLOCHTRIYS CYJCLAOM7 HEDYLOCHTRIYS YDENOW7 #ILALS	(3≏) 12 (31₹)	(45) 3 (7%) 10 (22%)	(43) 6 (143) 7 (163)
UTTNAFY SYSTEM			
ФЙВИТИВ-СПІТ УДЕИОМЯ ЖКІДИВА	(39)	(47) 1 (2%)	(45)
*UPINARY ELADDER PRANSITIONAL-PELL PAPILICMA	(37)	(45)	(41) 1 (2%)
A DISILOVST-JET. DIDITIONS A DISILOVST-JET. DIDITIONS	(39)	(^{u ~})	(49) 1 (2%)
ENDOCRINE SYSIEM			
BHEUCHAUMOCAIDKs' APTIGNAML RIDEENTI	(36)	(42) 1 (2%)	(⁴ 2)
<pre>#THYPOTD PIPILARY OFFCHOMA DISTLLARY ADENCOME POLLIGHTA-CPLL ADENOMA POLLIGHTA-CPLL ADENOMA C-TPLL ADENOMA C-TPLL ADENOMA DATLLARY CYSTIDENCMA, NOG</pre>	(۹۶)	(46) 1 (2%) 1 (2%) 7 (15%) 1 (2%) 2 (4%) 1 (2%)	(43) 1 (2%) 1 (2%) 14 (33%) 1 (2%) 4 (3%) 2 (5%)
NCAE			
NERVOUS SYSTEM			
NONE SLECITT ZENCE DECENC			
 NUMBER OF ANIMALS WITH TISSUE FYT NUMBER OF ANIMALS NECROPSIED 	WINED WICENSCOPIC	7 11 Y	

TABLE B1 (CONTINUED)

and all the second			
	00NTEOL (UNTR) 05-0330	ICW DOSE 05-0285	HIGH DOSE 05-0290
USCHLOSKELEIAL SYSTEM			
NONF			
ODY CAVITIES			
*BODY CAVITIES	(30)	(47)	(49)
MESOTHELIOMA, NOS		1 (2%)	
IL OTHER SYSIEMS			
NONE			
NIMAL DISPOSITION SUMMARY			
ANTMALS INITIALLY IN STULY	5 Ņ	⊂ 0	51
NATUELL DEATHD	13	Q	17
MORTBUND SACFIFICE	2	1	Ц
SCHEEVIED SACRIFICE			
ICCTDENELT V VIII UD			
ACCTDENTALLY KILLED	3.5	33	20

NUMPER OF ANTMALS WITH TISSUE EXAMINED MTCROSCOPICALLY * NUMBER OF ANTMALS NECROFSIED

TABLE B1 (CONCLUDED)

	CONTROL (UNTR) 05-0330	LOW DOSZ 05-0085	HIGH DOSE 05-0290
IIMCZ SIIMMIFX			
TOTLL ANIMALS WITH PRIMARY TUMOPS*	24	ų٩	2 =
TOTIL PRIMARY TUMORS	29	54	U F
TOTAL ANIMALS WITH BENIGN THMORE	2	12	20
TOWAL BENIGN TUMOPS		21	2.8
TOTAL ANTMALS WITH MELIGNANT TUMORS	22	25	12
TOTEL MALIGNANT THMCRS	2-	32	1.8
TOTAL ANTMALS WITH SECONDARY TUMOPSH	\$ 2	?	1
TOTRE SECONDARY TUMOPS	2	2	1
TOTEL INIMALS WITH TUMOPS UNCERTAIN-			
BENIGN OF MALTGHEFT		1	
TOINI UNCEPTAIN TUMORS		1	
TOTEL ANIMELS WITH THMOPS THCEPTAIN-			
SulWyeA Us Weaffaguru			
TOTAL UNCEPTAIN TUMORS			

* SECONDARY TUROPS: METASTATIC TUMOPS OF TUROPS INVARIAS INTO AN ADJACENT ORGAN

	06-0330 06-0330	LOW DOSE 06-0285	HIGH DOSE OF-0290
ANIMALS INITIALLY IN STUDY ANIMALS MISSING	50	50	دم 1
NNIMALS NECEOPSTED INTMALS EXAMINED HISTOPATHOLOGICALLY**	49 : 49	50 49	46 46
INTEGUMENTARY SYSTEM			
*SKIN UNDIFFERENTIATED CFFCINCM®	(10)	(5)	(46) 1 (2%)
*SUECUT TISSUF MIKED NESLNCHYMAI TUMOR, MAIIGNA	(⁴ °)	(50)	(46) 1 (2%)
FYSDTRATORY SYSTEM			
<pre>#LUNG UNDIFFERENTIATED CAFCINCMA METAS HEP/TOCRILULAP CARCINOMA, METAST ALVEOLAP/BEONCHIOLAP ADENOMA ALVEOLAP/BEONCHIOLAP CAPCINOMA</pre>	(μċ)	(43) 2 (43) 9 (193) 1 (23)	(46) 1 (2葉) 2 (4葉) 3 (7葉)
ARMATOPOIRTIC SYSTEM			
MULTIPLE ORGANS MALIGNANT LYMPHOMA, NOS MALIGNAMTHOMA, LYMPHOCYTIC TYPE	(49) 10 (20%)	(50) 1 (2%) 1 (2%)	(46)
MALIGALYMPHOMA, HISTIOCYTEC TYPP Malignant tymphoma, mixed typp lymphocytec leukemia	3 (2%)	10 (20%) 1 (2%)	3 (7%)
#BONE MARBOW HEPSTOCELLUIAR CARCINOMA, METSSI	(40)	(48) 1 (2%)	(45)
#SPIEEN HEPATOCFILUIAR CARCINONA, METASI HEMANGIOSAFCOMA MALIGMANI LYMPHOMA, MIXFD TYPF	(45) 1 (2%)	(48) 1 (<u>2</u> %) 1 (2%)	(45)
*MFDISSTINSI I.KODE MALISMANT_IYMPRCMA,_MIXED_TYPE	(44)	(45) <u>1 (27)</u>	(40)
* NUMERE OF ANTMALS WITH TISSHE EXAMIN * NUMPER OF ANTMALS MECROPSIES ** Excludes partially autolyzed animals	NED MICFOSCOPIC	NLLY	

TABLE B2 SUMMARY OF THE INCIDENCE OF NEOPLASMS IN FEMALE MICE TREATED WITH 1,5-NAPHTHALENEDIAMINE

B-7

			· · · · · · · · · · · · · · · · · · ·
		108 0028 06-1285	PIGB DOST COCT-31
**************************************	ر (ر غ ز) (ح -)	(25)	(-^)
***STATITITI (**DE **TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT	(22)	(03) 2 (-5) 1 (25) 1 (25)	، (عد) (,)
A.[[3434. [Āmbeuma]' 202 sīlāis	(一三) 「(2系)	(25)	(24)
PORTORNAL LANDRUNG, ALTER LAND	(+2)	(2~)	(C=) 1 (75)
	((()
IIIIIIII SISIA			
avbittettilligb cistadar keure	(19)	:-3)	(
CLEELLAE SASTA			
#15,100201001% 1951260%1 #5512002070012 10660%1 #11660	(===) (23)	(一子) 2 (6年) 35 (日下号)	(+4) 11 (243) 14 (258)
#20%127%7 #20%127%7	(38)	(=*) 1 (25)	(44)
51,00010415 0101041 #20160	(1)	(~) ㅋ (주종) ㅋ (주종)	(<i>z</i> ()
LELAST SISTER			
LESASILLONSE-LEIT ESÈLFICAS ALEIXIEX ELIDEE	(°°)	(++)	(2*)
EXDUCATIVE SAGER			
	(72) 	(35)	(3°) 1 (38)
* Kumpts ob skimple Alum Lickuf site	1978 **CEAR 208 20	· LIT	

Auxidz us a trails descipation

		·	
	04-0337 204 15 (2413)	107 2055 26-0285	HIGE DOST
SUIDOBELI DENOMS CREOMORECEE ADENCMS		· (3%) 3 (3%)	
#1Daidel	(22)	(2.1)	(4-)
BHEDCHBUNGUAGUAR	3 (7 %)	· ,	
BEAUCHBUNDCALONS' ANTIGARAL	* (73)		
*1541010	(22)	(++)	(45)
ENDITINEA PDENCAR		* (2%)	7 (15)
FOLDITOTAS-IELI IDEMONS	2 (= 3)	(11年)	10 (728)
FILLIOUINA- TELL CAPPINDMN	2 (= = = = =)		* (2*)
C-TELL SDENDWS		: (13)	二 (2年)
-1411 033C-N0*3		* (22)	E (132)
FIFTLIARY CYSIADERCMA, MOS		2 (1372)	2 (23)
SEPECOUCCIVE SUCCES			
	1431	121	11-53
301475-1211 032414C43			(23)
* D. 20 JS	(11)	(-==)	(-3)
RADOWERSINE STROWNE BUIND	* (23)		1 (24)
HER NGINA		· (~₹)	* (2#)
*.A.14	(44)	(1)	(41)
FEPATOCELITIAR CONTINUES, METAST		1 (22)	
CENNUICES-CELL TUNCE			1 (23)
TUETILE EDENGME	7 (22)	2 (2%)	
VILAULE SISIEN			
N087			
SEACINT SEASE UNGINE			
*SAPDETIN GINE	(20)	(50)	(48)
CASE SUEAGAS AGE	· · ·	1 22)	. ,
MUSCHINSKTIET, I SYSTEM			
NCAL			
* NUMBER OF SNIMALS WILE HISSNE EXAMI * NUMBER OF ANIMALS WEREPETED		, 11X	

TABLE B2 (CONCLUDED)

	СОМТРОІ (ПМТР) 96-0320	LOW DOSE 06-0285	HIGH DOSE 06-0290
CDY CIVITIES			
PODY CAVITIES MESOTHEFTOM/, NCS	(49)	(50)	(46) 1 (2%)
IL OTHER SYSTEMS			
NONT			
NIMAL DISECSITION SUMMARY			
ANTHRUS INTETALLY IN STURY	50	50	50
WETTIENT DESTRO	17	9	Q
MCPTEUND SACEIFICE	3		ج
SCHEDNIED SACHTEITE			1
TERMINAL SACRIFICE	30	41	24
ADINAL MISSING			1
THOTHNES SHEDIVZED ANTMERS			
CHOF SUMMERY			
TOTEL ANTWELS WITH PRIMERY TUMOPS*	21	41	37
LUGFT BELWERK LUNCES	2.9	88	71
DOMAL SALWARD ATTAL DENICH THREES	0	25	22
TOTAL REALGE THMORS	10	42	25
A AL LL LING LOUGHA	1	~ 2	22
TOTAL ANIMALS WITH MALIGNANT TUMOPS	16	37	?"
TOTAL MALIGNANT TUMORS	18	46	24
TOTAL ANIMALS WITH SECONDARY TUMOPS*	:	3	1
TOTAL SECURDARY TUMORS		10	2
TOTAL ANTWARS STOR DUMORS UNCERPAIN-			
BENTGN OF MATIGNENT			2
TOTAL UNCEPENTN TUMORS			2
TOTAL BRITHER DITER THREES HAGEDDETAL			
LUTAT ANTURTO MITH INDORE OVERALINA			
TOTAL UNCLEMATE TUMORS			
PFIMARY IUMORS: MIL TUMOPS EXCEPT SE	CONDARY TUMORS		
SECONDARY CUMOPS: METESTATIC TUMOPS	OF TUMORS INVA	SIVE INTO AN A	LJACENT OFGAN

APPENDIX C

SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN RATS TREATED WITH 1,5-NAPHTHALENEDIAMINE



	01-0330 1001-01(0413)	10W D05E 01-0280	-1-000F
WTWALS INITIALLY IN STUDY	25		24
SKYNARE EVANTVED UTETDIATUOTOTON	20 * T + 7* - 75	+ 7 	10
SALIPES EARLINED REFE PRIDE STOR			
INIEGHNENIAPI SYSTEM			
*58TN	(25)	(22)	(50)
VEGETABLE FOREIGN PODY	• •	1 (23)	
EPIDEBN'I INCLUSION CIST		1 (2%)	
BACESS, NOS		1 (2%)	
SCAP	* (의 종)		
*SUBOUT TISSUE	()=)	(:===)	(5.)
CYSI, BOS			1 (23)
FESDIRATORY SYSTEM			
+1 3 2AAA	(25)	(14 9)	(51)
INFLAMMNDIAN, SCHDE/CERCNIC		1 (2*)	
INFI PAMATION, CFRONIC		11 (22%)	1 (54)
ATTENCIONAL STREET	12=1	()	(23)
METAPLASTA, NOS	(- <i>1</i>	2 (03)	(-)
47.72			
#103G	(2-)	() ()	(~)
EPONCHOENZHWONTS NOS		- (-s)	2 (0.5)
PPONCHOENSEMONTS NECROTTZING	1 (63)		1
BRONCHOENEUMONTS, ACUTE	(- ()		? *3₹)
AB3CE55, NOS		1 (25)	~ ~ /
PNEUMONTE, CHRONIC MURINE		22 (45%)	≒ (11₹)
ABASTUBUIEILE SAGINA			
*BUKE A72238	(23)	(49)	()
EYPERPISSIN, NOS		っ (6え)	マ (二星)
HABABAIYSIY, HENSIUBULEDIU			」(7字)
1SDIFTY	(25)	(49)	(-3)
CONGESTION, NOS	* (===)	1 (25)	(. /
a a deservative and the field with the form of a second or the ca	- when it is a trivial with a start is a source	- and a course do Tak Sar In an	
* NUMBER OF ANIMALS WITH DISSUE E	XAMINED MICENSCHEIN	1112	
* ADABLE OF FRINTIS AFCSCERIED			
** EXCLUDES PARTIALLY AUTOLYZED ANIMALS			

TABLE C1 SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN MALE RATS TREATED WITH 1.5-NAPHTHALENEDIAMINE

TABLE C1 (CONTINUED)

	01-0330 01-0330	10W DOSE 01-0280	HIGH DOSE 01-0285
HEMOSTELEOSIS		2 (4%)	
MANDIBUISE L. NORE INFISHMATION, CERONIC HYPPEPINCIS, PISSME CELL	(24) 2 (8%)	(47)	(47) 1 (2*)
**************************************	(^{c t})	(35) 1 (3%) 1 (3%)	(76)
CIPCUINTOPY SYDDEM			
RHERBA DEPIRETION, NOS RHERBA	(25)	(49) 1 (2%) 25 (51%)	(47) 7 (15%)
INFLIMMNTION, THEONIC	(25) 1 (4%)	(49)	(47)
DIGESTIVE SYSLEM			
<pre>#LIVER CONGESTION, CHECKIC EXESSIVE CHOINGTOFIPEOSIS DEGENERATION, NOS NECENERATION, NOS NECENERATION, NOS NECENERATION, NOS EACOPHILIS CYTO CHANGE FF CAL CELLUINA CHANGE CIENE-CELL CHANGE HYPEFEISET, POCL NECENERATION.</pre>	(2°) 1 (4%) 1 (4%) 1 (4%) 1 (4%) 1 (4%)	(49) 1 (2%) 4 (9%) 1 (2%) 1 (2%) 3 (6%) 1 (2%)	(49) 2 (4%) 1 (2%) 1 (2%) 2 (6%) 1 (2%)
#LIVEF/CENTPILOBUL/P METROSIS, NOS	(25)	(49)	(49) 1 (2%)
#PSNOFIS, FOCAT BIBOSIS, FOCAT BIBOBHY, FOCAL	(25)	(48) 1 (2%)	(45) 1 (2%)
*SIOK*CH ULCER, ACUTY HYPEFELSELA, BASAL CELL HYPEFELSELA, DASAL CELL	(24)	(47) 1 (2%) 1 (2%)	(47) 1 (?%)

• NUMBER OF ANIMALS AITH TISSUE EXAMINED MICROSCOPICALLY • NUMBER OF ANIMALS NECECRESIED

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TABLE C1 (CONTINUED)

	CONTROL (UNTR) 01-0330	LCW DOSE 01-0280	HIGH DOSE 01-0285
#COLON PARPSTISA	;23)	(45) 5 (11%)	(46)
UFINAPY SYSTEM			
#KIDNEY CYST, NOS PYEIONEPHRITIS, POCAL FYELONEPHRITIS, CHRONIC	(2 ⁵)	(49) 1 (2%) 1 (2%)	(48) 1 (2⊄)
NEPHROSIS, NOS Hyperplasia, spithelial	21 (84%)	42 (86%)	15 (31%) 2 (4%)
*KIENEY/COPTEX MULTILOCULAP CYST	(?5) 1 (4系)	(49)	(48)
*KIDNEY/PELVIS CALCULUS, NOS	(25)	(49)	(48) 1 (2%)
#UPINAPY ELADDEF CPLCULUS, NOS	(25)	(49) 2 (4%)	(48) 1 (2%)
ENDCCRINE SYSTEM			
#EITUITARY Hyperplisin, focal	(22)	(44)	(44) 1 (2%)
#ADPENAL HYPERPLASIA, NOEULPP	(24)	(48)	(48) 1 (2%)
*ADRENAI COPTEX HYPERDI'SIA, NOS HYPERDIASIA, FOCAL	(24) 1 (4系)	(48)	(48) 3 (6%)
#ADRENAL MEDULIA Hyperplasia, NOS Hyperplasia, Focal	(24) 4 (17%)	(48) 1 (2%)	(48) 2 (4%)
*"HYPOID CYST, NOS FOLLICULAR CYST, NOS INFLAMMATION, CHPONIC HYPEPPL'SIA, C-CELL	(21)	(⁴⁷) 1 (2%) 1 (2%) 1 (2%)	(47) 1 (2%) 6 (13%)
#PARATHYROID HYPERPLASIA, NOS	(13)	(24)	(28)

NUMBER OF INIMALS WITH TISSUE EXAMINED MICEOSCOPICALLY * NUMBER OF ANIMALS NECROPSIED

TABLE C1 (CONTINUED)

	CONTERL (UMTE) 01-0330	109 DOSE 01-0280	HIGH DOSE 01-0285
*FRNOPRATIC ISLETS Hypfrpi'sl', Nos	(25)	(48) 1 (2%)	(45)
FEPRODUCTIVE SYSTEM			
ABSCESS, NOS LECTROIDM	(25)	:49) ; (68)	(50) 1 (25)
HEMOBRINGS	(25) 1 (4종)	(49)	(50)
*FROSTATE INFLAMMATION, SUPPUPATIVE TYPLAMMATION, CUTF/CHPCMIC INFLAMMATION, CHRONIC STROFHY, NOS	(25) ? (8%)	(49) 3 (6%) 1 (2%) 10 (20%)	(47) 17 (?6 %)
SEMINAL VESICLE NERREHY, NOS	(25)	(49) 10 (20%)	(50) 17 (34)
*TESTTS PEPTAPTEPITTS ATTOPHY, NOS ATTOPHY, FOCAL SPEEM TOGENTC APTRET HYPOSPERMATOGENESIS	(25) 5 (2つぎ)	(49) 10 (20%) 3 (6%) 1 (2%) 1 (2%)	(49) 1 (2%)
REPTETERNIS NECROSIS, NOS	(25)	(49) 1 (2%)	(50)
APROUS SYSTEM			
#CEREBRUM REMORTFAGE	(25)	(49) 1 (23)	(47)
#STAIN HYPPOCEPHALUS, NCS	(25)	(4°) 2 (4%)	(47) 1 (2*)
*PENIN SIFN HEMORPHAGE	(25)	(49) 1 (2%)	(27)
STEDINT SENSE ORGANS			
* NUMBER OF FNIMALS WITH TISSUE EX	AMINED MICPOSCOPIC	4 L L Y	

* NUMBER OF ANIMALS WITH TISSUE * NUMBER OF ANIMALS NECROPSIED

TABLE C1 (CONCLUDED)

	CONTROL (UNTR) 01-0330	LOW DOSE 01-0280	HIGH DOSE 01-0285
MUSCULOSKEIETAL SYSTEM			
NONP			
CON CAVITIES			
TABDOMINAL CAVITY NFCROSIS, PAI	(25)	(49) 1 (2%)	(57) 2 (4%)
ALL OTHER SYSTEMS			
NONE			
SPECIAL MORPHOLOGY SUMMARY			
NO LESION FEFORIED	4	1	
AUTO/NECPOPSY/NO HISTO AUTOLYSIS/NO NECPOPSY		1	1
* NUMBER OF ANIMALS WITH TISSUE EXA * NUMBER OF ANIMALS NECROPSIED	MINED MICROSCOPIC	ALLY	

TABLE C2 SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN FEMALE RATS TREATED WITH 1,5-NAPHTHALENEDIAMINE

	00NTEOL (UNTR) 02-0330	100 DOSE 02-0280	HIGH DOSE
NUTHILS INITIALLY IN STUDY ANIMALS NECROPSIED ANIMALS EXAMILED HISTOFATHOLOGICALLY	25 24 ** 24	50 50 50	50 50 50
INTEGUMENTARY SYSTEM			
N.CM E			
FESPIFATORY SYSTEM			
NELYMMATION, CHOONIC	(24)	(50) 1 (2%)	(50)
#LUNG/BRONCHIOLF METAPLASTA, NOS	(24)	(5 ⁰) 2 (4秀)	(())
#LUNG EFONCHIECTASIS EFONCHIECTASIS INFLAMMNION, INTERSTITIAL ONTUMNEA, CHPONIC MURINE	(24)	(50) 1 (2%) 19 (38%)	(50) 2 (4%) 1 (2%) 1 (2%) 2 (4%)
HENAMODOTATIO SYSTEM			
#POPE MARFOW HYPERDIASIA, NOS Hyperdiasia, Hemamopoistic	(21)	(49) 1 (2%) 1 (2%)	(49) 1 (2%)
*SPLERN MERRPLASIA, NOS HYPERPLASIA, PEMITOPOIETIC HYPERPLASIA, EPYTHPOID	(23)	(50) 1 (2%) 1 (2%) 2 (5%)	(49) 1 (?%)
EFAL HODDULELLS	1 (4%)	5 (08)	. (2.3)
#LYMEH NODE Hyperplasia, lymphote	(19)	(48) 1 (2%)	(49)
#MP"DIBULAS L. NODE <u>CONGESTION, NOS</u>	(19)	(49)	(49) <u>1 (23)</u>

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

	CONCROL (UNTR) 02-0330	10W DOSE 02-0280	HIGH DOSE 02-0285
HYPERPLISIA, PIASMA CELI	2 (114)		
CIRCULATORY SYSTEM			
#HFABT DEGENERATION, NCS	(24)	(5^) 5 (10%)	(50) 2 (4%)
#MYDCARDIUM Calcification, Focal	(24) 1 (4架)	(50)	(50)
MEDTAL CALCIFICATION	(2 ⁴) 1 (4系)	(50)	(50)
"COSONNEY ARTERY Mediai Calcification Calcification, Nos	(24) 1 (4%)	(50) 1 (2%)	(50)
CTGESTIVE SYSTEM			
#ITVPP CONGESTION, CHPONIC PASSIVE CHOINGICFIEPOSIS NECTOSIS, FOCAL INFARCT, FOCAL METAMORPHOSIS PATTY PASOPHILIC CYTO CHANGE CLFAF-CRLL CHANGE INGIECTASIS	(24) 10 (42系) 1 (4系)	(57) 1 (2%) 1 (2%) 1 (2%) 1 (2%) 2 (4%) 1 (2%)	(49) 1 (2%) 2 (4%) 12 (24%)
HYPERPLASEA, RASOPHILIC #LIVER/CENTFILOPULAP NECROSIS, NOS	(24)	(50) 2 (4%)	. (49)
#PANCPEAS ATFORMY, FOCAL	(22)	(49)	(47) 1 (2%)
#STOMACH Atypin, Nos Hyperpinsia, Basal Ceil	(24)	(50) 1 (2%) 2 (4%)	(49) 3 (6%)
#COION PARASTTISM	(24)	(47) 3 (6%)	(46)
UPINAFY SYSTEM			
*KIDNEY GYSTNOS	(24)	(50) 1. <u>[2%]</u>	(49)

NUMBER OF ANIMALS WITH TISSUE EXAMINED MICROSCOPICALLY
 NUMBER OF ANIMALS NECROPSIED

	CONTROL (UNIR) 02-0330	1CW DCSE 02-0280	HIGH DOSE
GIOMERUIONEPHETTIS, NOS NEPHEOSIS, NOS NECROSIS, MEDULLARY CALCIFICETION, POCAL	8 (33%)	1 (2%) 7 (14%) 1 (2%)	1 (2%) 1 (2%)
*KIDNEY/TUBULE CALCIPICATION, NOS	(24) 1 (4%)	(50)	(49)
DOCTINE SYSIEM			
*EITHITDERY CYST, NOS HEMOSTDERUSIS HYPERPLASI, NOS	(21) 1 (5%)	(50) 1 (2%) 1 (2%)	(47) 1 (2%)
#/DPENAL COPIFX HypfRplasia, Nos	(24)	(50) 1 (2%)	(49) 3 (F%)
#ADPENAL MEDULIA Hyperplasia, Nos	(34)	(51) 1 (2%)	(49)
#THYPOIC POLLICULAR CYST, NGS INFLAMMAINON, CHRONIC HYPEFELASIA, C-CFLL	(21) 1 (5%)	(49) 1 (2%)	(48) 7 (15%)
PPRODUCTIVE SYSTEM			
*MAMMARY GIAND GAINCTCTELE LACTATION	(24)	(5)) 12 (24%)	(50) 3 (6%) 4 (8%)
YNGINA HYPEFKEFAIOSIS	(24)	(50) 1 (2%)	(50)
#UTEPUS HYDPOMETPA EPIDEPMAL INCLUSION CYST THPOMBOSIS, NOS FYOMETRA ATROPHY, NOS	(24)	(49) 1 (2%) 1 (2%) 6 (12%) 3 (5%)	(48) 1 (2%) 3 (6%)
#UT EPUS/ENCOMETRIUM	(24)	(49)	(48)

* NUMBER OF ANIMALS NECROPSIED

	CONTROL (UNTR) 02-0330	LOW DOSE 02-0280	HTGH DOSE 02-0285		
INFLAMMATION, SUPPUPATIVE INFLAMMATION, CHPONIC HYPEFPLASIA, CYSTIC MEMAPLASIA, SQUAMOUS	3 (13%) 1 (4%)	1 (2%) 4 (8%)	1 (2%)		
#OVARY/OVIDUCT INFLAMMATION, SUPPUPATIVE ABSCESS, NOS	(24) 5 (21%)	(49)	(48) 2 (4%)		
HOVAFY CYST, NOS INFLAMMATION, SUPPUPATIVE INFLAMMATION, ACUTE ABSCESS, NOS	(24) 2 (8考) 1 (4考)	(49) 8 (16%) 1 (2%) 1 (2%)	(49)		
NERVOUS SYSTEM					
#BRAIN HYDPCCFEHALUS, NCS	(23)	(50) 1 (2%)	(51)		
SFECIAL SENSE ORGANS					
*EYF TYPLAMMATION, NOS PHTHISIS BULBI	(24)	(50)	(50) 1 (2%) 1 (2%)		
MUSCULOSKELETAL SYSTEM NONE					
EODY CAVITIES					
CABDOMIN®L CAVITY NECPOSIS, FAT	(24)	(5 ⁰) 1 (2%)	("0)		
ALL OTHER SYSTEMS					
CRANIOBUCCAL POUCH CYST, NCS		1			
SFPCIAL MORPHOLOGY SUMMARY					
NO LESION FEPOFIED	1				
# NUMBER OF ANIMALS WITH TISSUE EXAMINED MICROSCOPICALLY * NUMBER OF ANIMALS NECEOFSIED					

TABLE C2 (CONCLUDED)

	CONTROL (UNTR)	LOW DOSE	HIGH DOSE
	02-0330	02-0280	02-0285
AUTOLYSIS/NC NECROPSY	1		
# NUMPER OF ANIMALS WITH TISSUE EXAM: * NUMBER OF ANIMALS NECROPSIED	INED MICHOSCOPIC	A LLY	

APPENDIX D

SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN MICE TREATED WITH 1,5-NAPHTHALENEDIAMINE
	CONTROL (UNTR) 05-0330	LOW DOSE 05-0285	HTGH DOSE 05-0290
PNIMALS INITTALLY IN STUDY PNIMALS MISSING	50 2	50 1	۶Ŋ
INIMALS NECPORSTED INIMALS EXAMINED HISTOPATHOLOGICILLY**	39 * 39	47 47	49 46
IMTEGUMENTAPY SYSTEM			
*SKIN EPIDERMAL INCLUSION CYST INFLAMMATION, CEPOPIC FIBROSIS	(39) 1 (3%) 1 (3%) 1 (3%)	(47)	(49)
*SUBCUT TISSUE Abstess, Nos	(39)	(47) 1 (2%)	(49)
RESPIRATORY SYSTEM			
#LUNG EFONCHOENBUMONTA, NCS Hypefelisia, adenomatous	(3 ⁹)	(46)	(45) 1 (2종) 14 (71종)
HEMATOPOIETTC SYSTEM			
#SPIFEM HYPERPLASIA, LYMPHOID	(38) 1 (3%)	(45)	(41)
HEMATOPOIESIS FFYTHROPOIESIS	1 (3%) 2 (8%)	2 (4%)	
#MANFTEULAP L. NODE Hypprdiasia, plasma Cell	(36) 1 (3%)	(43)	(35)
*MESENTERIC L. NODE INFIAMMATION, GRANULCMATOUS	(36)	(43)	(35) 1 (3%)
HYPEPPLASIA, NOS Hypppplasia, lymphoid	4 (11%) 4 (11%)	1 (2%)	
CIPCULATORY SYSTEM			
#HEAFT PEDIARTPECTS	(39)	(47) 1 (2%)	(44) • us - and the task assessment and to asses

TABLE DI SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN MALE MICE TREATED WITH 1,5-NAPHTHALENEDIAMINE

* NUMPER OF ANIMALS WITH TISSUE EXAMINED MICROSCOPICAILY * NUMPER OF ANIMALS NECEOFSIED **EXCLUDES PARTIALLY AUTOLYZED ANIMALS

TABLE D1 (CONTINUED)

	CON "FOI (ปีพบัR) 05-0330	10W DOSE 05-0285	HIGH DOSE 05-0290
DIGESTIVE SYSIEM			
#LIVEP NECPOSIS, FOCAL CLEAP-CELL CHANGE HYPEEPL'SIA, FOCAL	(39) 3 (8%)	(45) 1 (2%) 1 (2%)	(43)
*PANCEEAFIC DUCT DILATITION, NOS	(36)	(46) 1 (2%)	(40)
*EUGENUM &MYLOTEOSIS	(37)	(47)	(39) 1 (3%)
*JEJUNUM Amyloidosis	(37) 1 (3%)	(47)	(२९) 1 (२६)
#TIEUM BMYLOTDOSIS	(37) 2 (5考)	(47)	(36)
COLON PRASTISM	(37)	(40) 2 (5%)	(36)
UFINERY SYSTEM			
*KYDNEY HYDPONEPHICSIS FYELOMEPHALIIS, FOCAL FYELOMEPHATIS, CHPONIC GLOMEAULOSCLEPOSIS, NOS NFCPOSIS, MEDULIARY INFAECT, KELLED AMYLODOSIS C/LCTFICETION, NOS CALCIFICATION, FOCAL	(39) ? (8%)	(47) 1 (2%) 1 (2%) 1 (2%) 5 (11%) 1 (2%) 1 (2%) 1 (2%)	(45) 2 (47) 1 (25) 4 (97) 3 (75) 12 (273) 1 (25)
*KTENEY/COPTEX SCRF	(39) 1 (3%)	(47)	(45)
CALCIPICATION, NOS	(39)	("")	(45) 5 (11%)
*PERIPENDI TISSUE DBSCF35, NOS	(39)	(47)	(45) 1 (2%)
#KIENEY/GLOMEPULUS MAYLOIDOSIS	(39)	(47)	(45)

NUMPER OF ANIMALS WITH TISSUE EXAMINED MICROSCOPICALLY
 NUMPER OF ANIMALS NITPOFSIED

TABLE D1 (CONTINUED)

		- const 10. 20 count of hit - dealers and a pair	
	СОМТРОЦ (UN TP) 05-0330	LCW DOSE	HIGH DOSE 05-C290
<pre>#UPINARY BIADDEF CALCULUS, NOS PEFTA 374FETTS</pre>	(37)	(45) 1 (2%) 2 (4%)	(41)
HYPPRPLASIA, BPTTHRIIAL			1 (?%)
ENDOCRINE SYSTEM			
#ADRENKI AMYLCIDOSIS	(36) 2 (6%)	(4?) 1 (2%)	(42)
THYPOID	(38) 1 (3%)	(46)	(47)
INFLAMMATION, ACHPP INFLAMMATION, ACHPP INFLAMMATION, ACHTE/CHRONIC AMVICIDOSIS	د (۲۰۰۰) د (۲۰۰۰)		1 (2%) 1 (2%)
HYPERDISIA, PACAL Hyperdisia, paptilary Hyperdisia, adenomatous Hyperdisia, politcular-cell	2 (5%)	1 (2%) 11 (24%)	7 (16%) 2 (5系)
HADLEDFISTS' NOR #5925-443-011	(28) 1 (4系)	(10)	(9)
*PANCEPATTC ISLETS Hyperplisit, Nos	(36) 1 (3%)	(46)	(45)
FEPFODUCTIVE SYSTEM			
CLITORAL GLAND DTLATETICN, NOC	(39)	(47) 1 (2%)	(^{4 °})
NEEVOUS SYSTEM			
#SUBARACHNOID SPACE HEMORPHAGE	(?8)	(47)	(40) 1 (3%)
#3P&TN HFMORPH2GE	(38)	(47)	(40) 1 (3%)
SPECIAL SENSE ORGANS			

* NUMBER OF ANTMAIS WITH TISSUE EXAMINED MICROSCOPICALLY * NUMBER OF ANIMALS NECROFIED

TABLE DI (CONCLUDED)

	0001201 (11012) 0001201 (11012)	LOW DOSE 05-0285	HIGH DOSE 05-0290
RUSCUTURKETEIVI RARIEM			
NONF			
ECDY CRVITIES			
Oúz			
VII OTHER SYSTEMS			
-WATU-DUSIS -WATU-DUSIS	(39) 1 (3%)	(47)	(4 9)
SUSCINE MORPHOLOGY SUMMARY			
NO LEGION REPORTED	e	1	1
NEGEORESY EPPENDE HISTO FERFORMEL NEGEORESY EPPENDE HISTO FERFORMEL NUTO/NECEDESY (CO. MISTO	2	1	1
AUTOLYSIS/NC RECEDESY	Q.	2	1

ng na ng panalana na 100 na 100 na 100 na na 100 na na 100 ma na 100 m			
	06-0330 CONIEOF (DAIS)	LCW DOSE 06-0285	HIGH LOSE
ANTMATS INTTATIY TO STUDY	50	50	5.0
ANTMALS MISSING		···	1
ANIMALS NECFOPSIED	49	50	4 S
INIMALS EXAMINED HISTOPATHCLOGICALLY*	* 49	49	<u>и</u> к
INTEGUMENTARY SYSTEM			
- SKTN	(40)	(50)	(46)
INFLAMMETION, MOUTE	1 (2≯)		
FESPIPATOPY SYSTEM			
#T.UNG	(4°)	(43)	(46)
EFONCHOFNEUMONIA, NOS			1 (2#)
PNEUMONTA, CHRONIC MUPINE		3 (F%)	2 (4%)
MEMAPTASIA, ADENOMATIOS MEMAPTASIA, NOS			2 (45)
UINITADAIITA SVETIM			
HENRYOPPULLUL DIVIZH			
#BONE MARBOW	(40)	(49)	(45)
FIBECSIS		1 (2%)	
HYPERPLASIA, HEMITOPOIETIC		5 (4名)	
#SPLERN	(45)	(48)	(45)
HEMOSIDEPOSIS		1 (2%)	
HYPEFPIISIA, LYMPHOID	1 (?%)		
БелйньОБОТь2.2	4 (9%)		
#LYMPH NOLE OF THOREY	(44)	(45)	(46)
HYPERPL'SIA, NOS	¹ (2%)	() =)	v · · <i>i</i>
#PANCPEATIC L.NODE	(44)	(45)	(40)
HFMATOPOIESIS	1 (2%)	(-)	、 ,
#LUMBSE LYMPH NODE	(44)	(45)	(40)
HYPEPPLASIA, NOS	1 (2%)		
#MESENT FOIC L. MODE	(44)	(45)	(40)
HYPERPI'SIA, NOS	* (27)		

TABLE D2 SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN FEMALE MICE TREATED WITH 1,5-NAPHTHALENEDIAMINE

* NUMBER OF ANIMALS WITH TISSUE EXAMINED MICEOSCOPICALLY * NUMBER OF ANIMALS MECEOPSIED

**EXCLUDES PARTIALLY AUTOLYZED ANIMALS

TABLE D2 (CONTINUED)

	CONTECL (UNTR) D6-0330	LOW DOSE 06-0285	8768 DOSE 06-0290
HYPERDISIS, IYMPHOID Hemaiopoidsis	1 (2号) 1 (2号)		
#SENIL LYMER NOBE HEMAICEDIASIS	(44) 1 (2%)	(45)	(u0)
CTROULITORY SYSTEM			
HEART CALOTRICATION, FOCAL	(49)	(49)	(46) 1 (2%)
DIGESTIVE SYSTEM			
#IIVPP INFIGUT, NCS CIENTROTIC CUNNER	(46)	(43) 1 (2%)	(ué)
HYPPYELSETC NODULE HEMATOPOLESIS	1 (2%) 1 (2%)	(23)	
ADIMORENS Berteiterite Adimorens	(38)	(47)	(40) 1 (2%) 1 (2%)
≠JFJUNUM MYLCIROSIS	(42)	(47)	(45) 1 (2考)
#TUFUM Amyinidosis	(42)	(47) ? (4%)	(45)
ULT VPY SYSTEM			
GIOWERNICNEPRATT'S, NCS	(4 K)	(49) 1 (2%)	(u 6)
ARAIOIDUSIS URANIOSUIFAURIS VOR	2 (4%)	3 (6%)	1 (2%)
ANDARA FISDDER AADEBDISEN, EDIMAETISI ANDIAVEA FISDDER	(u 3)	(46)	(^{[[]}) u (°%) 1 (2%)
ENDOCSTWE SYSCEM			
*PITUITATY HYPEFPINSIA, ECCAL	(34)	(35)	(30)
* NUMBER OF ANIMALS WITH TISSUE EXAM * NUMBER OF ANIMALS FECROPTIED	MIMED MICROSCOPIC	A LLY	

TABLE D2 (CONTINUED)

	CONTECT (UNTE) 16-1330	LCW DCSE 06-0285	HIGH DOSE 06-0390
#PDRFNAL THPCMBUS, ORGANIZED AMYLCIDOSIS	(48)	(44) 1 (2%)	(44) 1 (2%)
#THYPOTE INPEAMMETION, ACUTE IMPLAMMATION, FOUTE FOCAL	(44) 1 (2%)	(→ ^G) 1 (2%)	(45) * (24)
HYDEFPLASIE, NOS HYDEFPLASIE, PEPILIERY HYDERPLASIE, FOLLTODIER-CELL	2 (5%)	1 (2%)	2 (4%) 2 (4%)
#PARAMHYOOTS Hyperplasis, Nos	(?4)	(1?)	(15) 1 (7%)
FFFODUCTIVE SYSTEM			
#UTERUS HYDRCMETRA	(44)	(45)	(⁴³) 5 (12%)
#UTFFUS/ENDOMETFIUM HYPEFPLISI', CYSTIC	(44) 30 (63%)	(45) 도 (11종)	(<i>m</i> , <i>s</i>)
#OVARY/OVIDUUT Absoiss, Hos	(40)	(45)	(43) 1 (?%)
<pre>#OVAFY CYST, NOS HEMOSEHAGIC CYSI ABSCESS, NCS INFLAMMATION, CHEONIC</pre>	(世4) 7 (14で) 4 (9%) 1 (2※) 2 (5%)	(45) 3 (7%)	(غەر) <u>:</u> (غەر) (17)
FRVOUS SYSIRM			
#FRATN/MENINGES INFLAMMATION, CHOONIC	(UF)	(47)	(43) 1 (27)
δεειβομές Τυίδ πόεντά	(4£)	(47)	ا (5 مز) (ج)
RECTAL SENSE OPGANS			
NCVE			

* NUMEEP OF NIMALS NECECOSIED

TABLE D2 (CONCLUDED)

	(UNIP) JOALWOL	LOW DOST	HIGH DOSE
	0e=0330	76-0285)F = r 2 q 0
USCULOSKEIETAL SYSTEM			
*SKELPTAL MUSCLE AFSCISS, NOS	(40) 1 (2考)	(51)	(4 E)
UDA U.ATLIBE			
NECCOSIS, FUT	(uò)	(51) 1 (2%)	(⁴ 6)
LL OTHER SYSTEMS			
DIEUSS JISSUE SIEFTIIS DIEUSS JISSUE	1 1		
PECIAL YOPEHOLOGY SUMMAPY			
NO LESION FEPOPTED Avinal Missing/No Necropesy	3	2	3
NUTO/NETPOPSY/HISTO FERF	ŝ	21	
NUTOIYSTS/NO NECRORSY	1		3

Review of the Bioassay of 1,5-Naphthalenediamine* for Carcinogenicity

by the Data Evaluation/Risk Assessment Subgroup of the Clearinghouse on Environmental Carcinogens

June 29, 1978

The Clearinghouse on Environmental Carcinogens was established in May, 1976, in compliance with DHEW Committee Regulations and the Provisions of the Federal Advisory Committee Act. The purpose of the Clearinghouse is to advise the Director of the National Cancer Institute (NCI) on its bioassay program to identify and to evaluate chemical carcinogens in the environment to which humans may be exposed. The members of the Clearinghouse have been drawn from academia, industry, organized labor, public interest groups, State health officials, and quasi-public health and research organizations. Members have been selected on the basis of their experience in carcinogenesis or related fields and, collectively, provide expertise in chemistry, biochemistry, biostatistics, toxicology, pathology, and epidemiology. Representatives of various Governmental agencies participate as ad hoc members. The Data Evaluation/Risk Assessment Subgroup of the Clearinghouse is charged with the responsibility of providing a peer review of reports prepared on NCI-sponsored bioassays of chemicals studied for carcinogenicity. It is in this context that the below critique is given on the bioassay of 1,5-Naphthalenediamine for carcinogenicity.

The reviewer agreed with the conclusion in the report that 1,5-Naphthalenediamine was carcinogenic in treated female rats and in both sexes of mice. He noted that the study was conducted in a room in which other compounds were under test. Based on the experimental findings, he concluded that 1,5-Naphthalenediamine may pose a carcinogenic risk to humans. The reviewer moved that the report on the bioassay of 1,5-Naphthalenediamine be accepted as written. The motion was approved without objection.

Clearinghouse Members present:

Arnold L. Brown (Chairman), Mayo Clinic
Paul Nettesheim, National Institute of Environmental Health Sciences
Verne Ray, Pfizer Medical Research Laboratory
Verald K. Rowe, Dow Chemical U.S.A.
Michael B. Shimkin, University of California at San Diego
Louise Strong, University of Texas Health Sciences Center

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

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