135 PC NIH LIBRARY 298.5 1155 no.95 itional Cancer Institute 1978 ARCINOGENESIS **Technical Report Series** No. 95 1978 **BIOASSAY OF 3-(CHLOROMETHYL)PYRIDINE HYDROCHLORIDE** FOR POSSIBLE CARCINOGENICITY CAS No. 6959-48-4 NCI-CG-TR-95

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



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DEPARTMENT OF HEALTH, EDUCATION, AND WELFARE

National Institutes of Health

REPORT ON BIOASSAY OF 3- (CHLOROMETHYL) PYRIDINE HYDROCHLORIDE FOR POSSIBLE CARCINOGENICITY

Availability

3-(Chloromethyl)pyridine hydrochloride (CAS 6959-48-4) has been tested for cancer-causing activity with rats and mice in the Bioassay Program, Division of Cancer Cause and Prevention, National Cancer Institute. A report is available to the public.

<u>Summary</u>: A bioassay of 3-(chloromethyl)pyridine hydrochloride for possible carcinogenicity was conducted by administering the test chemical by gavage to Fischer 344 rats and B6C3F1 mice. Applications of the chemical include use in manufacturing of a variety of chemicals.

It is concluded that under the conditions of this bioassay, 3-(chloromethyl)pyridine hydrochloride was carcinogenic in male Fischer 344 rats and in B6C3F1 mice of both sexes, producing papillomas and carcinomas at the site of topical application, the stomach.

Single copies of the report are available from the Office of Cancer Communications, National Cancer Institute, Building 31, Room 10A21, National Institutes of Health, Bethesda, Maryland 20014.

Dated: October 13, 1978

Director National Institutes of Health

(Catalogue of Federal Domestic Assistance Program Number 13.393, Cancer Cause and Prevention Research)

Mational Institutes of Health Building 10 Bethesda Corviand 20014

BIOASSAY OF

3-(CHLOROMETHYL)PYRIDINE HYDROCHLORIDE

FOR POSSIBLE CARCINOGENICITY

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

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

DHEW Publication No. (NIH) 78-1345



BIOASSAY OF 3-(CHLOROMETHYL)PYRIDINE HYDROCHLORIDE 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 3-(chloromethyl)pyridine hydrochloride 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 environmental chemicals have the capacity to produce cancer in animals. Negative results, in which the test animals do not have a greater incidence of cancer than control animals, do not necessarily mean that the test chemical is not a carcinogen, inasmuch as 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 that exposure to the chemical is a potential risk to The actual determination of the risk to man from animal man. carcinogens requires a wider analysis.

CONTRIBUTORS: The bioassay of 3-(chloromethyl)pyridine hydrochloride was conducted by Litton Bionetics, Inc., Kensington, Maryland, initially under direct contract to NCI and currently under a subcontract to Tracor Jitco, Inc., the prime contractor for the NCI Carcinogenesis Testing Program.

The experimental design and doses were chosen by Drs. E. K. Weisburger¹, J. H. Weisburger^{1,2}, and N. P. Page^{1,3}, NCI project officers, and Dr. F. M. Garner⁴, the principal investigator. The administration of the test chemical and the observation of the animals were supervised by Dr. Garner and Mr. S. Johnson, the co-principal investigator, with the technical assistance of Mr. R. Cypher⁴, Mr. H. D. Thornett⁴, and Mr. D. J. Howard⁴. Ms. J. Blalock⁴ was responsible for assembly of data. Histopathologic examination of the rats was performed by Drs. R. J. Montali⁴, H. Seibold⁴, N. J. Wosu⁴, and P. Hildebrandt⁴. Mouse tissues were examined by Drs. Wosu and B. C. Zook⁴. All diagnoses of tumors were reviewed by Dr. Montali, who prepared the interpretive pathology narrative.

Animal pathology tables and survival tables were compiled at EG&G Mason Research Institute⁵. The statistical analyses were performed by Dr. J. R. Joiner⁶ and Ms. P. L. Yong⁶, using methods selected for the bioassay program by Dr. J. J. Gart⁷. Chemicals used in this bioassay were analyzed under the direction of Dr. E. Murrill⁸, dosage solutions were analyzed by Mr. H. Paulin⁴, and the results of the analyses were reviewed by Dr. S. S. Olin⁶.

This report was prepared at Tracor Jitco⁶ under the direction of NCI. Those responsible for the report at Tracor Jitco were Dr. L. A. Campbell, Director of the Bioassay Program; Dr. S. S. Olin, Deputy Director for Science; Dr. J. F. Robens, toxicologist; Dr. R. L. Schueler, pathologist; Dr. G. L. Miller, Ms. L. A. Waitz, and Mr. W. D. Reichardt, bioscience writers; and Dr. E. W. Gunberg, technical editor, assisted by Ms. Y. E. Presley and Ms. P. J. Graboske.

The following other scientists at NCI were responsible for evaluating the bioassay, interpreting the results, and reporting the findings: Dr. Kenneth C. Chu, Dr. Cipriano Cueto, Jr., Dr. J. Fielding Douglas, Dr. Dawn G. Goodman⁹, Dr. Richard A. Griesemer, Dr. Morton H. Levitt, Dr. Harry A. Milman, Dr. Thomas W. Orme, Dr. Robert A. Squire¹⁰, Dr. Sherman F. Stinson, Dr. Jerrold M. Ward, and Dr. Carrie E. Whitmire.

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SUMMARY

A bioassay of 3-(chloromethyl)pyridine hydrochloride for possible carcinogenicity was conducted by administering the test chemical by gavage to Fischer 344 rats and B6C3F1 mice.

Groups of 50 rats and 50 mice of each sex were administered 3-(chloromethyl)pyridine hydrochloride in a vehicle of distilled water three times per week at one of the following doses, either 75 or 150 mg/kg body weight for the rats and either 100 or 200 mg/kg body weight for the mice. The low-dose rats were dosed for 103 weeks and the low-dose mice for 102 weeks. Because of early deaths in the high-dose animals, the high-dose rats were dosed for only 83 weeks and the high-dose mice for only 81 weeks. Controls consisted of groups of 20 rats and 20 mice of each sex which were administered the vehicle only for 104 weeks. All surviving rats and mice were killed at 104 weeks.

Mean body weights of the male and female rats were lower in the dosed groups than in the corresponding control groups, and the depressions in weight were dose related. At the termination of the administration of the test chemical to the high-dose groups of rats, the mean body weights of these groups recovered rapidly. The mean body weights of the male mice were unaffected by the administration of the chemical; those of the females were only slightly affected. Mortality was generally higher in the dosed groups of rats and mice than in the corresponding control groups and was dose related in all tests except those using the female mice; however, sufficient numbers of animals of each species and sex were at risk for the development of late-appearing tumors.

In rats, proliferative squamous-cell lesions of the forestomach were observed in the dosed males (carcinomas: high-dose 1/50; papillomas: low-dose 1/47, high-dose 2/50; hyperplasias: low-dose 1/47, high-dose 2/50) and the dosed females (carcinomas: high-dose 1/48), but not in the male or female vehicle controls. The results of the Fisher exact test were not significant for squamous-cell papillomas or carcinomas. However, comparison of the incidence of these tumors in the high-dose males with that in 99 historical vehicle controls shows that the probability that three or more such tumors did not occur by chance, given that none have been observed in the controls in this laboratory, is P = 0.014.

or carcinomas In mice, squamous-cell papillomas of the forestomach occurred in the low- and high-dose groups of each sex, but not in the corresponding control groups. The incidence in the high-dose males was significantly higher (P = 0.025) than that in the control males (males: vehicle controls 0/19, low-dose 2/43, high-dose 10/47 [21%]; females: vehicle controls 0/19, low-dose 1/45, high-dose 5/48 [10%]). Comparison of the incidences of these tumors in the high-dose males and females with those observed in the corresponding groups of 100 historical vehicle controls of each sex shows that the probability that their occurrence was not due to chance is P < 0.001. Also, a life- table analysis of the incidence in males indicated a significant (P = 0.003) increase in tumors over the period of observation (58 weeks to 104 weeks) in relation to an increase in dose.

Although the incidence of squamous-cell papillomas and carcinomas in male and female rats was significant only in males compared with historical vehicle controls, these tumors are of the same type as those appearing at the same site in male and female mice. Because these tumors are rare and not found in controls, and because they were found in dosed animals of both species, they are considered to be related to administration of the test chemical by gavage.

It is concluded that under the conditions of this bioassay, 3-(chloromethyl)pyridine hydrochloride was carcinogenic in male Fischer 344 rats and in B6C3F1 mice of both sexes, producing papillomas and carcinomas at the site of topical application, the stomach.

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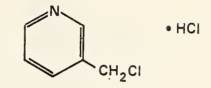
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I. INTRODUCTION

3-(Chloromethyl)pyridine hydrochloride (CAS 6959-48-4; NCI CO3838) is an intermediate that has been proposed for use in the synthesis of agricultural, pharmaceutical, and veterinary

chemicals (Arnall and Clark.



3(chloromethyl)pyridine (hydrochloride)

1966). Its synthesis was achieved in the mid T960's by the Midland Tar Distillers, Ltd., in Great Britain, through a process in which the alpha carbon of the alkyl side chain to the pyridine ring is chlorinated preferentially (Arnall and Clark, 1966). This discovery made it economically feasible for the first time to synthesize large quantities of monochloroalkylpyridine intermediates. This compound is neither manufactured in the United States, nor imported, at the present time (Stanford Research Institute, 1977; USITC, 1977a and 1977b), however, at the time it was selected for bioassay, it was felt that it could become a widely used industrial intermediate.



II. MATERIALS AND METHODS

A. Chemical

Three batches of the test chemical, hereinafter referred to as 3-(chloromethyl)pyridine hydrochloride, were obtained from Columbia Organic Chemicals, Columbia, South Carolina, for these studies. These batches were identified by the date of receipt at Midwest Research Institute, Kansas City, Missouri, as Lot No. CO2-7-73, Lot No. CO12-5-73, and Lot No. CO2-25-75. All batches were used during the chronic studies; Lot No. CO2-7-73 was also used during the subchronic studies.

The identity of each batch was confirmed by infrared, nuclear magnetic resonance, and ultraviolet spectral analyses. Elemental analyses (C, H, N, Cl) for $C_6H_7NCl_2$ were slightly low for chlorine in Lot Nos. C02-7-73 and C02-25-75. Trace impurities were found in all three lots by thin-layer chromatography. High-pressure liquid chromatography (ultraviolet detector, 254nm) indicated the presence of a single impurity, accounting for 0.2% of the total peak area in Lot No. C02-7-73, and of three minor impurities (0.49%) in Lot No. C02-25-75. Lot No. C02-7-73 contained 0.69 \pm 0.04% water, Lot No. C012-5-73, < 0.17%, and Lot No. C02-25-75, 0.82 \pm 0.11%, as determined by Karl Fischer

analysis. Throughout this report the term used to represent this material is 3-(chloromethyl)pyridine hydrochloride.

These batches were stored at 4°C in the original containers.

B. Dosage Preparation

Solutions of 3-(chloromethyl)pyridine hydrochloride were prepared in distilled water (Borden Polar Water Co., Beltsville, Md.) at concentrations of 1 and 2% for mice and 0.75 and 1.50% for rats. These were administered by gavage on the same day on which they were prepared.

C. Animals

Fischer 344 rats and B6C3Fl mice of each sex were obtained from Charles River Breeding Laboratories, Wilmington, Massachusetts, under a contract with the Division of Cancer Treatment, NCI.

The animals were 28 days of age when received at the laboratory and were quarantined for 2 weeks prior to the start of the bioassay. Animals with clinical signs of disease and runts were killed. The remaining animals were segregated into equal weight groups and assigned to control or dosed groups in such a way that the mean weights of animals in each cage within a particular group were approximately the same.

D. Animal Maintenance

All animals were housed in temperature- and humidity-controlled rooms. The temperature was maintained at 21-25°C and the relative humidity at 45-55%. There were 10 changes of room air per hour, and the incoming and exhaust air was filtered through high efficiency particulate air (HEPA) filters (Flanders Filters, McLean, Va.). The animal rooms were positively pressurized with respect to the exit hall and negatively pressurized with respect to the entrance hall. Rooms were illuminated by cool white fluorescent lighting 8 hours per day.

Rats were housed four per cage and mice five per cage in solid polycarbonate cages (Lab Products, Inc., Garfield, N. J.). Each cage was covered with a wire mesh screen and a sheet of filter paper and contained heat-treated hardwood chip bedding (Absorb-Dri[®], Lab Products, Garfield, N.J.) in the bottom. Cages and water bottles were sanitized two times per week, racks were washed each month, and feed hoppers were sanitized once per week at approximately 82°C; bedding was replaced two times per week and filter paper was replaced each month.

The animals were fed Wayne[®] Lab Blox (Allied Mills, Inc., Chicago, Ill.) and received fresh diets three times per week.

Water bottles contained tap water which had been acidified to pH 2.5.

Rats and mice were housed in separate rooms. Control and dosed animals were housed in the same room. Animals administered 3-(chloromethyl)pyridine hydrochloride were housed in the same room with animals administered the following chemicals:

Rats

Feed Studies

(CAS 105-55-5) N,N-diethylthiourea (CAS 99-56-9) 4-nitro-o-phenylenediamine (CAS 89-25-8) 1-phenyl-3-methyl-5-pyrazolone

Mice

Feed Studies

```
(CAS 2735-04-8) 2,4-dimethoxyaniline
(CAS 140-49-8) 4-chloroacetylacetanilide
(CAS 139-94-6) nithiazide
(CAS 624-18-0) p-phenylenediamine dihydrochloride
(CAS 99-56-9) 4-nitro-o-phenylenediamine
(CAS 89-25-8) 1-phenyl-3-methyl-5-pyrazolone
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Gavage Studies

```
(CAS 512-56-1) trimethylphosphate
(CAS 4377-33-7) 2-(chloromethyl)pyridine hydrochloride
(CAS 1955-45-9) pivalolactone
```

E. Subchronic Studies

The LD_{50} for 3-(chloromethyl)pyridine hydrochloride administered orally to either Fischer 344 rats or B6C3F1 mice has been reported as 316 mg/kg (Litton-Bionetics, Inc., 1973). Subchronic studies were conducted with Fischer 344 rats and B6C3F1 mice to estimate the maximum tolerated doses of 3-(chloromethyl)pyridine hydrochloride, on the basis of which two doses (hereinafter called "low" and "high" doses) were determined for the chronic studies. For the subchronic studies, the test chemical was administered by gavage, three times per week, at doses of 68, 100, 147, 215, or 316 mg/kg to rats, and 100, 147, 215, 316, or 464 mg/kg to mice. Five males and five females were tested at each dose, and groups of equal size served as vehicle controls, receiving distilled water only. Following a 7-week period of administration of the test chemical, the animals were observed for l week and then killed and necropsied.

Within the first 3 weeks of the subchronic test, two of the male rats and all of the female rats died at the highest dose, 316 mg/kg. In those male rats that were alive at week 7, mean body weights were depressed to the same extent in all groups, and in no case did this exceed 15% of controls. In the surviving females, there were no apparent effects on body weights.

In the mice, all males and females administered the highest dose, 464 mg/kg, died during the first week. At the end of week 7, there were only small weight depressions in the surviving female groups and no apparent weight depression in the males.

No signs of toxicity were found on gross pathologic examination of the organs taken from rats and mice.

The low and high doses for the chronic studies using rats were set at 75 and 150 mg/kg, and those for the chronic studies using mice were set at 100 and 200 mg/kg, respectively.

F. Chronic Studies

The test groups, doses administered, and times on study of the chronic studies are shown in tables 1 and 2.

G. Clinical and Pathologic Examinations

All animals were observed twice daily for signs of toxicity and were weighed and palpated for masses at regular intervals. Animals that were moribund and those that survived to the termination of the bioassay were killed using CO₂ and necropsied.

The pathologic evaluation consisted of gross and microscopic examination of major tissues, major organs, and all gross lesions from killed animals and from animals found dead. The following tissues were examined microscopically: skin, lymph nodes, mammary gland, salivary gland, bone marrow, trachea, lungs and bronchi, heart, thyroid, parathyroid, esophagus, stomach, small intestine, large intestine, liver, gallbladder (mice), pancreas, spleen, kidney, adrenal, urinary bladder, prostate or uterus, testis or

Sex and Test Group	Initial No. of <u>Animals</u> ^a	3-(Chloromethyl) pyridine Hydrochloride Dose ^b <u>(mg/kg)</u>	Time (Dosed (weeks)	on Study Observed (weeks)
Male				
Vehicle-Control ^c	20	0		104
Low-Dose	50	75	103	1
High-Dose	50	150	83d	21
Female				
Vehicle-Control ^c	20	0		104
Low-Dose	50	75	103	1
High-Dose	50	150	83d	21

Table 1. Chronic Studies of 3-(Chloromethyl)pyridine Hydrochloride in Rats

^aRats were approximately 6 weeks of age when placed on study.

b3-(Chloromethyl)pyridine hydrochloride was administered three times per week in distilled water at a volume of 1 ml per 100 gm body weight. Doses were calculated using the mean body weight of the group and were adjusted every month.

^CVehicle controls received distilled water three times per week at a volume of 1 ml per 100 gm body weight.

^dBecause of early deaths, the high-dose rats were dosed for only 83 weeks.

Sex and Test Group	Initial No. of <u>Animals</u> ^a	3-(Chloromethyl) pyridine Hydrochloride Dose ^b (mg/kg)	Time on Study Dosed Observed (weeks) (weeks)		
Male					
Vehicle-Control ^C	20	0		104	
Low-Dose	50	100	102	2	
High-Dose	50	200	81 ^d	23	
Female					
Vehicle-Control ^C	20	0		104	
Low-Dose	50	100	102	2	
High-Dose	50	200	81 ^d	23	

Table 2. Chronic Studies of 3-(Chloromethyl)pyridine Hydrochloride in Mice

^aMice were approximately 6 weeks of age when placed on study.

- b3-(Chloromethyl)pyridine hydrochloride was administered three times per week in distilled water at a volume of 1 ml per 100 gm body weight. Doses were calculated using the mean body weight of the group and were adjusted every month.
- ^CVehicle controls received distilled water three times per week at a volume of 1 ml per 100 gm body weight.
- ^dBecause of early deaths, the high-dose mice were dosed for only 81 weeks.

ovary, brain, and pituitary. Occasionally, additional tissues were also examined microscopically. The different tissues were preserved in 10% buffered formalin, embedded in paraffin, sectioned, and stained with hematoxylin and eosin. Special staining techniques were utilized when indicated for more definitive diagnosis.

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

H. Data Recording and Statistical Analyses

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

verification of data transcription and for statistical review.

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

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

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

was examined histologically. However, when macroscopic examination was required to detect lesions prior to histologic sampling (e.g., skin or mammary tumors), or when lesions could have 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) was used to compare the tumor incidence of a control group with that of a group of dosed animals at each dose level. When results for a number of dosed 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) requires that the P value for any comparison be less than or equal to 0.05/k. In cases where this correction was used, it is discussed in the narrative section. It is not, however, presented in the tables, where the Fisher exact P values are shown.

The Cochran-Armitage test for linear trend in proportions, with continuity correction (Armitage, 1971), was also used when appropriate. Under the assumption of a linear trend, this test determines 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 is a positive dose relationship. This method also provides a twotailed test of departure from linear trend.

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

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

groups. The statistical tests for the incidence of tumors which used life-table methods were one-tailed and, unless otherwise noted, in the direction of a positive dose relationship. Significant departures from linearity (P < 0.05, two-tailed test) were also noted.

The approximate 95 percent confidence interval for the relative risk of each 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 dosed 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 dosed 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 dosed group than in the control.

The lower and upper limits of the confidence interval of the relative risk have been included in the tables of statistical analyses. The interpretation of the limits is that in approximately 95% of a large number of identical experiments, the true ratio of the risk in a dosed group of animals to that in a control group would be within the interval calculated from the

experiment. When the lower limit of the confidence interval is greater than one, it can be inferred that a statistically significant result (P < 0.025 one-tailed test when the control incidence is not zero, P < 0.050 when the control incidence is zero) has occurred. When the lower limit is less than unity, but the upper limit is greater than unity, the lower limit indicates the absence of a significant result while the upper limit indicates that there is a theoretical possibility of the induction of tumors by the test chemical, which could not be detected under the conditions of this test.

III. RESULTS - RATS

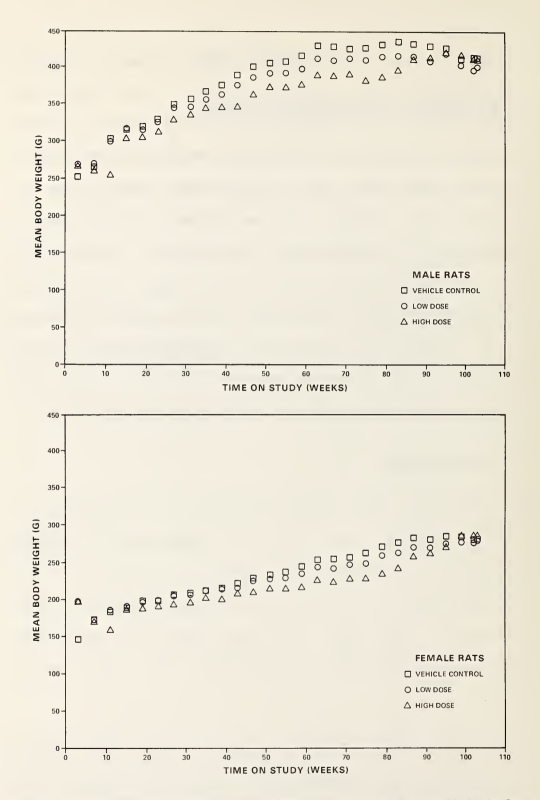
A. Body Weights and Clinical Signs (Rats)

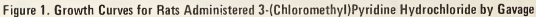
The mean body weights of the male and female rats were lower in the dosed groups than in the corresponding control groups, and the depressions in weight were dose related (figure 1). At the termination of the administration of the test chemical to the high-dose groups of rats (83 weeks), the mean body weights of these groups increased. Fluctuation in the growth curve may be due to mortality; as the size of a group diminishes, the mean body weight may be subject to variation. No other clinical signs related to administration of the test chemical were reported.

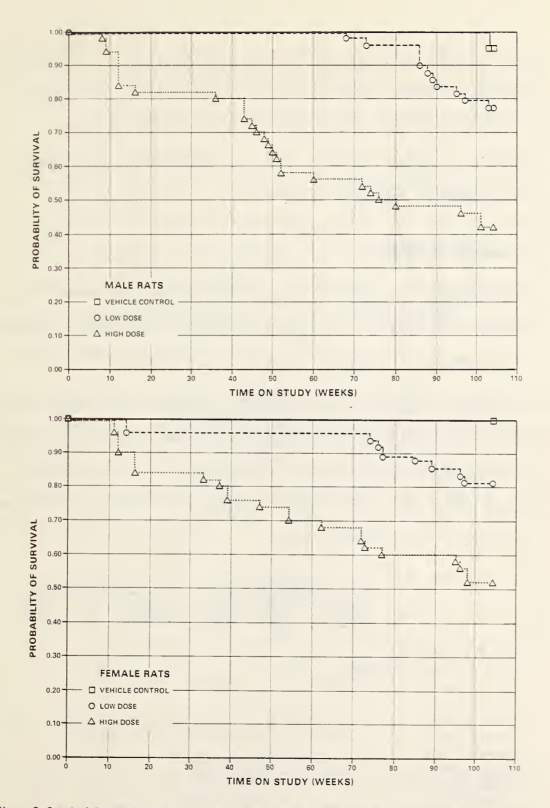
B. Survival (Rats)

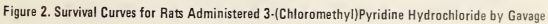
The Kaplan and Meier curves estimating the probabilities of survival for male and female rats administered 3-(chloromethyl)pyridine hydrochloride by gavage at the doses of this bioassay, together with those of the vehicle controls, are shown in figure 2. The result of the Tarone test for positive doserelated trend in mortality is significant (P < 0.001) in each sex.

In male rats, 21/50 (42%) of the high-dose group, 38/50 (76%) of the low-dose group, and 19/20 (95%) of the control group survived









to termination of the study. In females, 26/50 (52%) of the high-dose group, 40/50 (80%) of the low-dose group, and all 20 of the control group lived to termination of the study.

Sufficient numbers of rats of each sex were at risk for the development of late-appearing tumors.

C. Pathology (Rats)

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.

A variety of neoplastic lesions were evident in the control and dosed rats. Except for those of the forestomach, the tumors occurred in a random fashion in all groups. In the stomachs of dosed rats, there were both neoplastic and hyperplastic lesions as noted in the following table:

	Male Rats			Female Rats		
	Vehicle	Low	High	Vehicle	Low	High
	Control	Dose	Dose	<u>Control</u>	Dose	Dose
Number of Animals with Stomach Exami Microscopically	ned 19	47	50	20	45	48
Squamous-cell carcinoma			1(2%)			1(2%)
Squamous-cell			- (/			
papilloma		1(2%)	2(4%)			
Squamous-cell						
hyperplasia		1(2%)	2(4%)			

Grossly, the gastric nodules were described as single, white, pinpoint to 1 mm or 2 mm in diameter, warty growths, limited to the squamous portion of the stomach. Microscopically, the squamous-cell carcinomas were well differentiated and consisted of nests of squamous cells, some with keratin formation, that obliterated or invaded the muscularis mucosa but did not extend below the submucosa. The papillomas consisted of raised areas of spikes of hyperplastic squamous epithelium with hyperkeratotic caps. Chronic inflammation occurred in the submucosa beneath some of these lesions.

In organs other than the stomach, there were some degenerative and inflammatory lesions of the type usually encountered in aged rats, but none of the lesions were attributed to the test chemical. Chemically related lesions were not found in rats dying early in the study.

Based on the histopathologic examination, it was concluded that the gastric squamous-cell neoplastic lesions may be associated with the administration of 3-(chloromethyl)pyridine hydrochloride in rats under the conditions of this bioassay.

D. Statistical Analyses of Results (Rats)

Tables El and E2 in Appendix E contain the statistical analyses of the incidences of those primary tumors that occurred in at

least two animals of one group and at an incidence of at least 5% in one or more than one group. The early deaths in the high-dose group resulted in a shortened dosage period which obscures the meaning of trend analyses. Thus, these analyses have been omitted in the tables.

The results of the Fisher exact tests comparing the incidences of tumors in each of the dosed groups with that in the control group are not significant in the positive direction in either sex. The combined incidence of squamous-cell papillomas or carcinomas of the stomach in male rats was 0/19 in the controls, 1/47 (2%) in the low-dose group, and 3/50 (6%) in the high-dose group. The records of control animals at this laboratory indicate no such tumors occurred in 99 historical gavage vehicle-control male The results of the Fisher exact test of these incidences rats. are not significant, but under the estimate of 1% incidence in male control rats, the binomial probability (Fears, 1977) of three or more such tumors in 50 male rats is significant (P = 0.014). This analysis suggests an association between the occurrence of squamous-cell tumors in the high-dose group and the administration of the chemical. There was one squamous-cell carcinoma in the high-dose group of female rats compared with none in the 100 gavage vehicle-control female rats seen at this laboratory.

The results of the Fisher exact test on incidences of tumors of the pancreas and tumors of the testis in dosed male rats were significant in the negative direction. The higher incidences of tumors in the control group than in the dosed groups may have occurred because the dosed animals did not live as long as the control animals.

In each of the 95% confidence intervals of relative risk, shown in the tables, the value of one or less than one is included; this indicates the absence of significant positive results. It should also be noted that each of the intervals has an upper limit greater than one (except those of the incidences of isletcell tumors and of tumors of the testis in high-dose male rats), indicating the theoretical possibility of the induction of tumors by 3-(chloromethyl)pyridine hydrochloride, which could not be detected under the conditions of this test.

IV. RESULTS - MICE

A. Body Weights and Clinical Signs (Mice)

The mean body weights of the male mice were unaffected by administration of the 3-(chloromethyl)pyridine hydrochloride, and those of the females were only slightly affected (figure 3). Fluctuation in the growth curve may be due to mortality; as the size of a group diminishes, the mean body weight may be subject to variation. No other clinical signs related to administration of the test chemical were reported.

B. Survival (Mice)

The Kaplan and Meier curves estimating the probabilities of survival for male and female mice administered 3-(chloromethyl)pyridine hydrochloride by gavage at the doses of this bioassay, together with those of the vehicle controls, are shown in figure 4. In male mice, the result of the Tarone test for positive dose-related trend in mortality is significant (P = 0.006). In females, the result of the Tarone test is not significant.

There were 23/50 (46%) of the male high-dose group, 30/50 (60%) of the low-dose group, and 15/20 (75%) of the control group surviving to termination of the study. There were 30/50 (60%) of

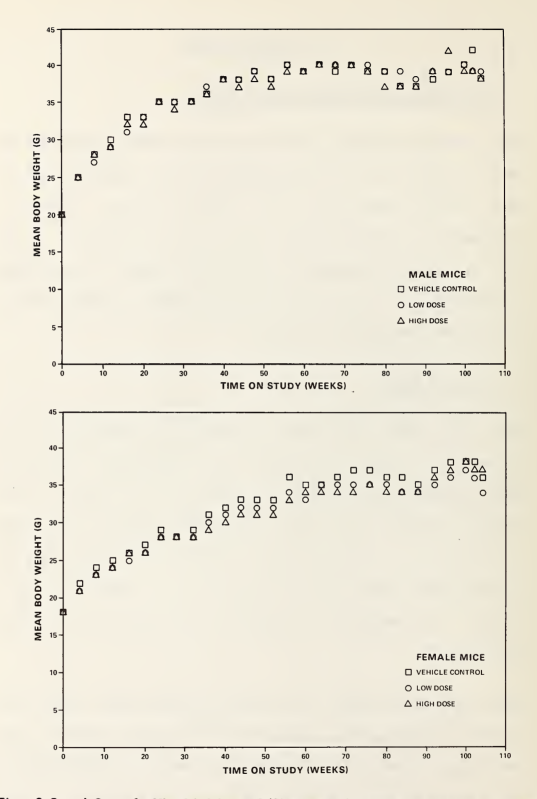
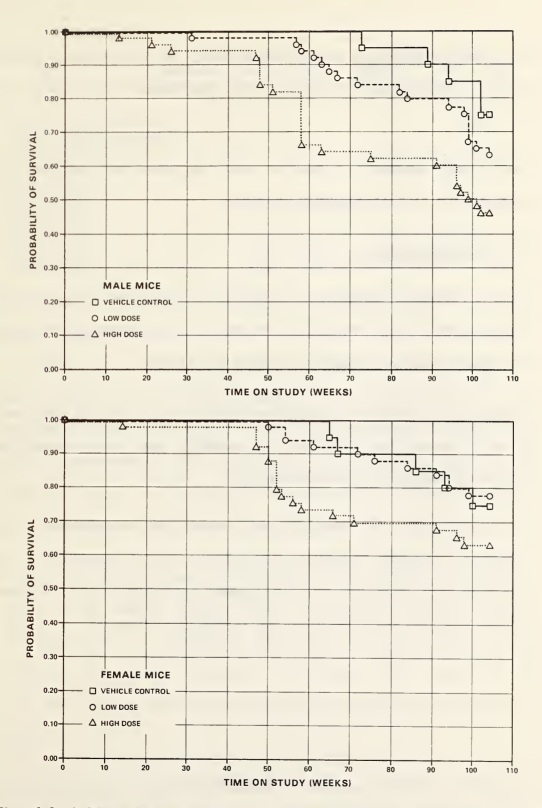
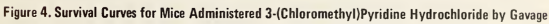


Figure 3. Growth Curves for Mice Administered 3-(Chloromethyl)Pyridine Hydrochloride by Gavage





the female high-dose group, 39/50 (78%) of the low-dose group, and 15/20 (75%) of the control group surviving to termination of the study.

Sufficient numbers of mice of each sex were at risk for the development of late-appearing tumors.

C. Pathology (Mice)

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

A variety of neoplastic lesions were evident in control and dosed mice. Except for those in the stomach, they were spontaneous tumors occurring in a random fashion.

In the stomach of dosed mice, there were hyperplastic and neoplastic lesions that did not occur in the controls. The lesions were limited to the squamous portion of the stomach (forestomach) and occurred at the following incidence:

	М	ale Mic	e	Fe	male M	ice
	Vehicle	Low	High	Vehicle	Low	High
	Control	Dose	Dose	<u>Control</u>	Dose	Dose
Number of Animals with Stomach Exam	ined					
Microscopically	19	43	47	19	45	48
meroscopicariy	17	45	-17	17	15	10
Squamous-cell carcinoma			2(4%)			2(4%)
Squamous-cell						
papilloma		2(5%)	8(17%)		1(2%) 3(6%)
Squamous-cell hyperplasia		1(2%)	2(4%)			3(6%)

One of the squamous-cell carcinomas in a high-dose male mouse was described grossly as a large, hard mass in the stomach. Microscopically, this tumor, which arose from the forestomach, had invaded the liver and pancreas and metastasized to the mesenteric lymph node. The other squamous-cell carcinomas were well differentiated and consisted of epithelial nests that invaded through the muscularis mucosa or into the stalk but did not extend beyond the gastric submucosa.

Grossly, the papillomas were described as single, pinpoint to small white nodules arising from the squamous gastric mucosa. Histologically, in favorable sections, they consisted of tall hyperplastic spikes of squamous epithelium with hyperkeratotic caps that lined connective tissue stalks derived from the stomach wall.

The hyperplastic areas consisted of increased layers of basophi-

lic squamous cells and hyperkeratosis. In some of the affected mice there was mild to moderate chronic inflammation in the submucosa underlying the papillomas and hyperplastic areas.

Other tumors observed were those commonly recorded as spontaneous neoplasms of mice. These included a few hepatocellular and pulmonary tumors and some hematopoietic neoplasms, all with a slightly elevated incidence in the dosed mice. The lowered survival rates of the dosed mice made it difficult to interpret the incidences of these tumors, particularly when the differences in incidences between the dosed and control groups were marginal.

Other nonneoplastic lesions were of the type usually found in aged mice, and none was attributed to the test chemical. Chemical-related lesions did not occur in mice dying early in the study.

Based on the histopathologic examination, it was concluded that the squamous-cell tumors of the forestomach were associated with the administration of 3-(chloromethyl)pyridine hydrochloride in B6C3F1 mice under the conditions of this bioassay.

D. Statistical Analyses of Results (Mice)

Tables Fl and F2 in Appendix F contain the statistical analyses of the incidences of those primary tumors that occurred in at

least two animals of one group and at an incidence of at least 5% in one or more than one group. The early deaths in the high-dose group resulted in a shortened dosage period which obscures the meaning of trend analyses. Thus, these analyses have been omitted in the tables.

The probability level of the Fisher exact test comparing the combined incidence of squamous-cell papillomas or carcinomas of the stomach in the high-dose group of male mice with that in the control group of male mice is at the upper limit (P = 0.025) of that required for significance by the Bonferroni inequality criterion for multiple comparisons. In females, the results of the Fisher exact test are not significant; however, these tumors occurred only in the dosed groups. No such tumors have been observed at this laboratory in any of 100 historical gavage vehicle-control male or female mice. Under the estimate of the binomial parameter (Fears, 1977) of an incidence of 1% in control mice, the binomial probability of 10/47 (21%) or higher incidence in the high-dose male mice is significant at a level less than P = 0.001.

In addition to the analysis described above, a life-table analysis was performed, utilizing the time at which a tumor was observed (see section H, p. 13) and based on the time-weighted

dose calculated over the 104 weeks of the bioassay. This analysis of the incidence in males (controls 0/19, low-dose 2/43 [5%], high-dose 10/47 [21%]) ranging from the first tumor observed in the high-dose group (58 weeks) to the end of the bioassay (104 weeks) indicated a significant (P = 0.003) increase in the observation of tumors over this time period in relation to increase in dose. These analyses indicate an association of squamous-cell papillomas or carcinomas of the stomach with the administration of the test chemical.

V. DISCUSSION

3-(Chloromethyl)pyridine hydrochloride was toxic for Fischer 344 rats and B6C3F1 mice inasmuch as mean body weights were depressed in the dosed rats and mortality was generally higher in both dosed rats and dosed mice than in corresponding control groups. The nature of the toxic effect could not be established histopathologically. The depression in mean body weight was dose related in both sexes of rats, and the mortality was dose related in rats and male mice. In female mice only the survival of the high-dose group was affected. Because of early deaths in the high-dose groups of both the rats and the mice, administration of the test chemical was terminated about 20 weeks earlier for these groups than for the low-dose groups of both species. Sufficient numbers of animals of each species and sex were at risk, however, for the development of late-appearing tumors.

In rats, proliferative squamous-cell lesions of the stomach were observed in the dosed males (carcinomas: high-dose 1/50: papillomas: low-dose 1/47, high-dose 2/50; hyperplasias: low-dose 1/47, high-dose 2/50) and a dosed female (carcinomas: high- dose 1/48), but not in the male or female vehicle controls. The results of the Fisher exact test were not significant for squamous-cell papillomas or carcinomas. However, comparison of the incidence of these tumors in the high-dose males with that in

99 historical gavage vehicle controls shows that the probability that three or more such tumors did not occur by chance, given that none have been observed in the controls in this laboratory, is 0.014.

In mice, squamous-cell papillomas or carcinomas of the stomach occurred in the low- and high-dose groups of each sex, but not in the corresponding control groups of the 100 historical controls of each sex. The incidence in the high-dose males was significantly higher (P = 0.025) than that in the control males (males: vehicle controls 0/19, low-dose 2/43, high-dose 10/47 [21%]; females: vehicle controls 0/19, low-dose 1/45, high-dose 5/48 [10%]). Comparison of the incidences of these tumors in the females with those observed high-dose males and in the corresponding groups of 100 historical vehicle controls of each sex shows that the probability that their occurrence was not due to chance is P < 0.001. Also, a life-table analysis of the incidence in males indicated a significant (P = 0.003) increase in tumors over the period of observation (58 weeks to 104 weeks) in relation to an increase in dose.

Although the incidence of squamous-cell papillomas and carcinomas in male rats was significant only in comparison with historical vehicle controls, these tumors are of the same types as those that appeared at the same site in male and female mice. Because

these tumors are rare and not found in controls, and because they were found in dosed animals of both species, they are considered to be related to administration of the test chemical by gavage in both species.

It is concluded that under the conditions of this bioassay, 3-(chloromethyl)pyridine hydrochloride was carcinogenic in male Fischer 344 rats and B6C3F1 mice of both sexes, producing papillomas and carcinomas of the forestomach. Neoplastic lesions related to chemical administration were restricted to the site of topical application, the stomach.



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APPENDIX A

SUMMARY OF THE INCIDENCE OF NEOPLASMS IN RATS ADMINISTERED 3-(CHLOROMETHYL)PYRIDINE HYDROCHLORIDE

BY GAVAGE



TABLE A1.

SUMMARY OF THE INCIDENCE OF NEOPLASMS IN MALE RATS ADMINISTERED 3(CHLOROMETHYL)PYRIDINE HYDROCHLORIDE BY GAVAGE

	VEHICLE CONTROL	LOW DOSE	HIGH DOSE
ANIMALS INITIALLY IN STUDY ANIMALS NECPOPSIED ANIMALS FXAMINED HISTOPATHOLOGICALLY	20 20 20	50 50 49	50 50 50
INTEGUMENTAPY SYSTEM			
*SUBCUT TISSUE FIBROMA FIEROSAFCOMA NEUROFIBROMA	(20)	(50) 1 (2%) 1 (2%) 1 (2%)	(50) 2 (4系)
RESPIRATORY SYSTEM			
*LUNG ALVECLAR/BRONCHIOLAR CARCINOMA C-CELL CARCINOMA, METASTATIC FIPROSARCOMA, METASTATIC	(20) 2 (10%) 1 (5%)	(49) 3 (6%) 1 (2%)	(49)
HEMATOPOIETIC SYSTEM			
*MULTIPLE ORGANS LEUKEMIA,NOS UNDIFPERENTIATED LEUKEMIA	(20) 1 (5%) 1 (5%)	(50) 1 (2%)	(50) 2 (4%)
CIRCULATORY SYSTEM			
* BLOOD VESSEL C-CFLL CARCINOMA, METASTATIC	(20)	(50) 1 (2%)	(50)
DIGESTIVE SYSTEM			
*LIVER NEOFLASTIC NODULE HEPATOCELLULAR CARCINOMA	(19) 1 (5%)	(48) 1 (2%)	(48)
*STOMACH SQUAMOUS CELL PAPILLONA	(19)	(47) <u>1 (2%)</u>	(50) <u>2 (4%)</u>
* NUMBER OF ANIMALS WITH TISSUE EXAMI * NUMBER OF ANIMALS NECROPSIED	INED MICROSCOP	ICALLY	

TABLE A1. MALE RATS: NEOPLASMS (CONTINUED)

ŧ

	VEHICLE CONTROL	LOW DOSE	HIGH DOSE
SQUAMOUS CELL CARCINOMA			1 (2%)
RINARY SYSTEM			
NONE			
NDOCRINE SYSTEM			
#PITUITARY CHRCMOPHOBE ADENOMA CHROMOPHOBE CARCINCMA	(19) 1 (5%)	(42) 4 (10%) 3 (7%)	(41) 1 (2%) 1 (2%)
#ADRENAL CORTICAL ADENOMA COKTICAL CARCINOMA PHEOCHROMOCYTOMA	(19) 1 (5%) 1 (5%) 2 (11%)	(49) 1 (2%) 2 (4%)	(49) 1 (2%) 1 (2%)
#THYRCIC FOLLICULAR-CELL ADENOMA FOLLICULAR-CELL CARCINOMA C-CELL ADENOMA C-CELL CARCINOMA	(17) 1 (6%) 1 (6%)	(39) 1 (3%) 1 (3%) 1 (3%)	(41) 1 (2%)
*PANCREATIC ISLETS ISIET-CELL ADENOMA ISIET-CELL CARCINOMA	(20) 3 (15%)	(48) 2 (4%) 1 (2%)	(49)
EPRODUCTIVE SYSTEM			
*PREPUTIAL GLAND ADENCMA, NOS	(20) 1 (5%)	(50)	(50)
#TESTIS INTERSTITIAL-CELL TUMOR	(19) 17 (89%)	(46) 38 (83%)	(47) 20 (43%
IERVOUS SYSTEM			
#BRAIN GLICBLASTOMA MULTIFORME	(20)	(46)	(47) 1 (2%)
PECIAL SENSE ORGANS			
NONE			

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

TABLE A1. MALE RATS: NEOPLASMS (CONTINUED)

	VEHICLE CONTROL	LOW DOSE	HIGH DOSE
MUSCULCSKEIETAL SYSTEM			
• MANDIBLE FIERCSARCOMA	(20)	(50)	(50) 1 (2%)
BODY CAVITIES			
• ME SENTERY LIFCMA	(20)	(50) 1 (2%)	(50)
*TUNICA VAGINALIS MESOTHELIOMA, NOS	(20)	(50) 1 (2%)	(50)
ALL OTHER SYSTEMS			
NCNE			
ANIMAL EISPOSITION SUMMARY			
ANIMALS INITIALLY IN STUDY NATUFAL DEATHØ MORIEUND SACRIFICE SCHELULED SACRIFICE	20 1	50 10 1	50 27 2
ACCIDENTALLY KIIIED TERMINAL SACRIFICE ANIMAL MISSING	19	1 38	21
@_INCLUEES_AUTOLYZED_A NI MALS			

NUMBER OF ANIMALS WITH TISSUE EXAMINED MICROSCOPICALLY
NUMBER OF ANIMALS NECROPSIED

TABLE A1. MALE RATS: NEOPLASMS (CONTINUED)

- -

	VEHICLE CONTROL	LOW DOSE	HIGH DOSE
TUMOR SUMMARY			
TOTAL ANIMALS WITH PRIMARY TUMORS*	20	45	24
TOTAL FRIMARY TUMORS	33	65	34
TOTAL ANIMALS WITH BENIGN TUMERS	19	41	21
TOTAL BENIGN TUMORS	25	52	26
TOTAL ANIMALS WITH MALIGNANT TUMORS	7	11	8
TOTAL MALIGNANT TUMORS	7	12	8
TOTAL ANIMALS WITH SECONDARY TUMORS	# 1	2	
TOTAL SECONDARY TUMORS	1	2	
TOTAL ANIMALS WITH TUMORS UNCERTAIN-	_		
BENIGN OR MALIGNANT	1	1	
TOTAL UNCERTAIN TUMORS	1	1	
TOTAL ANIMALS WITH TUMORS UNCERTAIN	-		
FRIMARY CR METASTATIC			
TOTAL UNCERTAIN TUMORS			
* PRIMARY TUMORS: ALL TUMORS EXCEPT S	ECONDARY TU	MORS	
# SECONDARY TUMORS: METASTATIC TUMORS	OR TUMORS	INVASIVE INTO AN	ADJACENT ORGAN

TABLE A2.

SUMMARY OF THE INCIDENCE OF NEOPLASMS IN FEMALE RATS ADMINISTERED 3(CHLOROMETHYL)PYRIDINE HYDROCHLORIDE BY GAVAGE

	VEHICLE CONTROL	LOW DOSE	HIGH DOSE
ANIMALS INITIALLY IN STUDY ANIMALS NECFOPSIED ANIMALS EXAMINED HISTOPATHOLOGICALLY	20 20 20	250 49 49	50 50 50
INTEGUMENTARY SYSIEM			
*SKIN TRICFORPITHELIOMA SEBACEOUS ADENGCARCINOMA KERATOACANTHOMA FIEPOMA	(20)	(49) 1 (2%) 1 (2%) 1 (2%) 1 (2%)	(50)
*SUECUT TISSUE LIPOMA MIXED MESENCHYMAL TUMOR, MALIGNA	(20)	(49) 1 (2%)	(50) 1 (2%)
RESPIRATOFY SYSTEM			
#LUNG ALVECLAF/BRONCHIOLAR CARCINOMA CORTICAL CARCINOMA, METASTATIC	(20) 1 (5%) 1 (5%)	(48)	(50) 1 (2%)
HEMATOPOIETIC SYSTEM			
* EFAIN MALIGNANT RETICULOSIS	(19)	(49)	(50) 1 (2%)
*MULTIFIE ORGANS UNDIFFERENTIATED LEUKEMIA LYMPHOCYTIC LEUKEMIA	(20) 1 (5%)	(49) 2 (4%) 1 (2%)	(50) 2 (4%)
CIRCULATORY SYSTEM			
NONE			
DIGESTIVE SYSTEM			
#STOMACH SQUAMOUS CELL CARCINCMA	(20)	(45)	(48) <u>1_(2%)</u>
 NUMBER OF ANIMALS WITH TISSUE EXAM NUMBER OF ANIMALS NECROFSIED 	INED MICROSCOP	ICALLY	
ο 50 ANIMALS WERE INITIALLY IN THE A MALE IN A FEMALE GROUP.	STUDY, BUT ONE	ANIMAL WAS FOUND	D TO BE

TABLE A2. FEMALE RATS: NEOPLASMS (CONTINUED)

	VEHICLE CONTROL	LOW DOSE	HIGH DOSE
RINARY SYSTEM			
#URINARY FLADDER PAPIILCMA, NOS	(16)	(40) 1 (3%)	(35)
NDOCRINE SYSTEM			
#PITUITARY CHRCMOFHOBE ADENOMA CHROMOPHOBE CARCINCMA	(19) 8 (42%) 1 (5%)	(43) 11 (26%) 1 (2%)	(40) 13 (33%) 1 (3%)
#ADRENAL CORTICAL CARCINOMA PHEOCHROMOCYTOMA	(19) 1 (5%)	(49) 1 (2%)	(50) 2 (4%)
#THYPOID FOLLICULAR-CELL ADENOMA C-CELL ADENOMA C-CELL CARCINOMA	(17)	(45) 1 (2%) 1 (2%)	(38)
*MAMMAPY GLAND SARCCMA, NOS FI EROADENOMA	(20) 2 (10%)	(49) 1 (2%) 4 (8%)	(50) 1 (2%)
SARCCMA, NOS		1 (2%)	• •
*FREPUTIAL GLAND ADENCMA, NOS	(20) 2 (10%)	(49) 1 (2%)	(50)
*VAGINA IEICMYCMA	(29)	(49)	(50) 1 (2%)
#UTERUS PAPILLARY ADENOMA SARCOMA, NOS	(19)	(47) 1 (2%) 1 (2%)	(49)
ENDOMETRIAL STROMAL POLYP HEMANGIOPERICYTOMA, NOS	2 (11%)	8 (17%) 1 (2%)	2 (4%)
#OVARY PAFILLARY ADENOMA	(18)	(45)	(48) 1 (2%)
ERVCUS SYSTEM			
*BRAIN CHRCMOFHOBE_CARCINCMA, INVASIVE	(19)	(49)	(50)

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

8 8 10 A A A

TABLE A2. FEMALE RATS: NEOPLASMS (CONTINUED)

	VEHICLE CONTROL	LOW DOSE	HIGH DOSI
EPENTYMONA		1 (2%)	
SPECIAL SENSE ORGANS			
NCNE			
USCULOSKEIFTAL SYSTEM			
NONE			
OEY CAVITIES			
* MEDIASTINUM MESCTHELIONA, NOS	(20) 1 (5%)	(49)	(50)
* MESENTERY LIFCMA	(20)	(49)	(50) 1 (2%)
LL OTHER SYSTEMS			
SITE UNKNOWN LIFCMA		1	
NIMAL EISPOSITION SUMMARY			
ANIMAIS INITIALLY IN STUDY NATURAL DEATHØ	20	50	50
MATURAL DEATHØ MORIPUND SACRIFICE SCHEDULED SACRIFICE ACCIDENTALLY KILLED		7 2	21 3
TERMINAL SACRIFICE ANIMAL MISSING	20	40	26
ANIMAL DELETED (WRONG SEX) INCLUDES AUTOLYZED ANIMALS		1	

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

TABLE A2. FEMALE RATS: NEOPLASMS (CONTINUED)

8 L

	VEHICLE CONTROL	LOW DOSE	HIGH DOSE	
TUMOR SUMMARY				
TOTAL ANIMALS WITH PRIMARY TUMORS* TOTAL FRIMARY TUMORS	14 19	31 43	23 29	
TOTAL ANIMALS WITH BENIGN TUMCRS TOTAL BENIGN TUMORS	12 14	26 34	18 21	
TOTAL ANIMALS WITH MALIGNANT TUMORS TOTAL MALIGNANT TUMORS	4 4	8 8	5 8	
TOTAL ANIMALS WITH SECONDARY TUMORS* TOTAL SECONDARY TUMORS	2 2			
TOTAL ANIMALS WITH TUMORS UNCERTAIN- BENIGN CR MALIGNANT TOTAL UNCERTAIN TUMORS	1 1	1 1		
TOTAL ANIMALS WITH TUMORS UNCERTAIN- PRIMABY OF METASTATIC TOTAL UNCERTAIN TUMORS				
* PRIMARY TUMORS: ALL TUMORS EXCEPT SEC * SECONDARY TUMORS: METASTATIC TUMORS (JACENT ORGAN	

APPENDIX B

SUMMARY OF THE INCIDENCE OF NEOPLASMS IN MICE ADMINISTERED 3-(CHLOROMETHYL)PYRIDINE HYDROCHLORIDE BY GAVAGE



TABLE B1.

SUMMARY OF THE INCIDENCE OF NEOPLASMS IN MALE MICE ADMINISTERED 3(CHLOROMETHYL)PYRIDINE HYDROCHLORIDE BY GAVAGE

	VEHICLE CONTROL	LOW DOSE	HIGH DOSE
ANIMALS INITIALLY IN STUDY	20	50	50
ANIMALS MISSING	20	2 48	50
AN IM ALS NECROPSIED ANIMAIS EXAMINED HISTOFATHOLOGICALLY		48	50
INTEGUMENTARY SYSTEM			
NONE			
RESFIRATCRY SYSTEM			
# LU NG	(20)	(45)	(48)
HEFATOCELLULAR CARCINOMA, METAST	1 (5%)		
ALVECLAE/DRCNCHIOLAR ADENCMA ALVEOLAP/BRONCHIOLAR CARCINGMA	1 (5%) 1 (5%)	4 (9%)	5 (10%)
OSTEOSAECOMA, METASTATIC	1 (5%)	4 (5%)	5 (10%)
*MULTIPLE ORGANS MALIGNANT LYMPHOMA, NOS	(20) 1 (5%)	(48) 2 (4%)	(50) 2 (4%)
MALIGNANT LYMPHOMA, MIXED TYPE LEUKEMIA,NOS	1 (5%)	1 (2%)	1 (2%) 1 (2%)
UN CIFFEFENTIATED LEUKEMIA	1 (5%)	3 (6%)	3 (6%)
LYMPHOCYTIC LEUKEMIA		1 (2%)	1 (2%)
* ERONCFIAL LYMPH NODE	(16)	(33)	(35)
ALVECLAR/BRONCHIOLAR CA, METASTA			1 (3%)
* MESENTERIC L. NODE	(16)	(33)	(35)
SQUAMOUS CELL CARCINCMA, METASTA MALIGNANT LYMPHOMA, NOS			1 (3%)
ALIGNANT LIMPHOMA, NOS ALIGNANT LYMPHOMA, MIXED TYPE		2 (6%)	1 (3%) 1 (3%)
CIRCULATORY SYSTEM			
* ELOOD VESSEL	(20)	(48)	(50)
ALVECLAR/BRONCHIOLAR CA, METASTA		1 (2%)	

NUMBER OF ANIMALS WITH TISSUE EXAMINED MICROSCOPICALLY

* NUMBER OF ANIMALS NECROPSIED

TABLE B1. MALE MICE: NEOPLASMS (CONTINUED)

	VEHICLE CONTROL	LOW DOSE	HIGH DOSE
DIGESTIVE SYSTEM			
#LIVFR SQUAMOUS CELL CARCINOMA, INVASI HEPATOCELLULAR ADENOMA	1 (5%)	(46)	(49) 1 (2%)
HEPATOCELLULAR CARCINOMA *PANCREAS SQUAMOUS CELL CARCINOMA, INVASI	2 (10%) (18) V	5 (11%) (33)	9 (18% (36) 1 (3%)
#SICMACH SQUAMOUS CELL PAPILLOMA SQUAMOUS CELL CARCINOMA	(19)	(43) 2 (5%)	(47) 8 (17% 2 (4%)
URINARY SYSTEM			
NONE			
ENDOCRINE SYSTEM #THYROID FOLLICULAR-CELL CARCINOMA	(10)	(31)	(35) 1 (3%)
REPRODUCTIVE SYSTEM			
#TESTIS INTEFSTITIAL-CELL TUMOR	(19)	(45) 1 (2%)	(44)
NERVOUS SYSTEM			
NONE			
SPICIAL SENSE ORGANS			
NGNE			

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

-

	VEHICLE CONTROL	LOW DOSE	HIGH DOSE
DDY CAVITIES			
NONE			
LI CTHER SYSTEMS			
SITE UNKNOWN			
SARCCMA, NOS		1	
NIMAL EISPESITION SUMMARY			
ANIMALS INITIALLY IN STUDY	20	50	50
NATURAL DEATHD	4	17	25
MORIBUND SACRIFICE	1	1	2
SCHEDUIED SACRIFICE ACCIDENTALLY KILLED			
TERMINAL SACRIFICE	15	30	23
ANIMAL MISSING		2	20
INCLUDES AUTOLYZED ANIMALS			
JHOR SUMMARY			
TOTAL ANIMALS WITH PRIMARY TUMCRS*	7	20	28
TOTAL PRIMARY TUMORS	8	22	35
TOTAL ANIMALS WITH BENIGN TUMORS	2	3	8
TOTAL BENIGN TUMORS	2	3	8
TOTAL ANIMALS WITH MALIGNANT TUMORS	5	19	24
TOTAL MALIGNANT TUMORS	6	19	27
TOTAL ANIMALS WITH SECONDARY TUMORS#	2	1	2
TOTAL SECONDARY TUMORS	2	1	4
TOTAL ANIMALS WITH TUMORS UNCERTAIN-			
BENIGN OF MALIGNANT			
TOTAL UNCERTAIN TUMORS			
TOTAL ANIMALS WITH TUMORS UNCERTAIN-			
FRIMARY OR METASTATIC			
TOTAL UNCERTAIN TUMORS			
PRIMARY TUMORS: ALL TUMORS EXCEPT SE	CONDARY TUMOR	S	

TABLE B1. MALE MICE: NEOPLASMS (CONTINUED)

TABLE B2.

SUMMARY OF THE INCIDENCE OF NEOPLASMS IN FEMALE MICE ADMINISTERED 3(CHLOROMETHYL)PYRIDINE HYDROCHLORIDE BY GAVAGE

	VEHICLE	LOW DOSE	HIGH DOSE
		LOW DOSE	
NIMALS INITIALLY IN STUDY	20	50	50
NIMALS MISSING NIMALS NECROPSIED	20	50	1 49
NIMALS EXAMINED HISTOFATHOLOGICALLY		50	49
NTEGUMENTAPY SYSTEM			
NONE			
ESPIRATORY SYSTEM			
# I U NG	(18)	(50)	(49)
HEFATOCELLULAR CARCINOMA, METAST ALVECLAR/BRONCHIOLAR ADENOMA		1 (2%)	1 (2%)
ALVEOLAR/BRONCHIOLAR CARCINCMA	1 (6%)		3 (6%)
OSTEOSARCOMA, METASTATIC		1 (2%)	
EMATOPOIETIC SYSTEM			
*MULTIPLE ORGANS	(20)	(50)	(49)
MALIGNANT LYMPHOMA, NOS MALIG.LYMPHOMA, HISTIOCYTIC TYPE	1 (5%)	2 (4%)	1 (2%) 1 (2%)
MALIGNANT LYMPHOMA, MIXED TYPE			2 (4%)
LEUKEMIA,NOS UN DIFFERENTIATED LEUKEMIA		2 (4%)	1 (2%) 2 (4%)
LYMPHOCYTIC LEUKEMIA	1 (5%)	1 (2%)	- (,
#LYMPH NOCE	(16)	(42)	(40)
ALVECLAR/BRONCHIOLAR CA, METASTA			1 (3%)
# MANDIBULAR L. NODE	(16)	(42)	(40)
MALIGNANT LYMPHONA, MIXED TYPE			1 (3%)
* MESENTFRIC L. NODE	(16)	(42)	(40)
MALIGNANT LYMPHOMA, MIXED TYPE		4 (10%)	2 (5%)
IRCULATORY SYSTEM			

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

villa - -

TABLE B2. FEMALE MICE: NEOPLASMS (CONTINUED)

	VEHICLE CONTROL	LOW DOSE	HIGH DOSE
DIGESTIVE SYSTEM			
*IIVEF HEFATOCFLIULAR CARCINOMA	(20) 1 (5%)	(50)	(49) 2 (4%)
#STCMACH SQUAMOUS CELL PAPIIICMA SQUAMOUS CELL CARCINOMA	(19)	(45) 1 (2%)	(48) 3 (6%) 2 (4%)
JAINARY SYSTEM			
NONE			
ENDOCRINE SYSTEM			
*PITUITAFY CHRCMOPHOBE ADENOMA	(9)	(25)	(29) 2 (7 %)
#ADRENAL CORTICAL ADENOMA	(16)	(41) 1 (2%)	(43)
*THYRCIC FCLIICULAR-CELL CARCINOMA	(8)	(15) 1 (7%)	(36)
EPRCDUCTIVF SYSTEM			
*UTERUS ENCCMETRIAL STROMAL FCLYP	(18)	(47)	(47) 1 (2%)
*CVARY FAFILLARY ADENOMA	(6)	(13) 1 (8%)	(33)
ERVCUS SYSTEM			
NONE			
FECIAL SENSE ORGANS			
* EYE/LACFIMAL GLAND ADENOCAPCINOMA, NOS	(20)	(50)	(49)

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

TABLE B2. FEMALE MICE: NEOPLASMS (CONTINUED)

	VEHICLE CONTROL	LOW DOSE	HIGH DOSE
MUSCULOSKEIFTAL SYSTEM	- •	•	
NONE			
BOEY CAVITIES *ABDOMINAL CAVITY SARCCMA, NOS	(20) 1 (5%)	(50)	(49)
ALL OTHER SYSTEMS			
*MUITIFLE ORGANS SARCCMA, NOS	(20)	(50) 1 (2%)	(49)
ANIMAL EISFOSITION SUMMARY			
ANIMAIS INITIALLY IN STUDY NATURAL DEATHƏ MORIBUND SACRIFICE SCHEDUIED SACRIFICE ACCIDENTALLY KILLED	20 4 1	50 11	50 17 2
TERMINAL SACRIFICE Animal Missing	15	39	30 1
<u>@ INCLUDES_AUTOLYZED_ANIMALS</u>			
# NUMBER OF ANIMALS WITH TISSUE EX * NUMBER OF ANIMALS NECROPSIED	AMINED MICROSCOE	CALLY	

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TABLE B2. FEMALE MICE: NEOPLASMS (CONTINUED)

	VEHICLE CONTROL	LOW DOSE	HIGH DOSE
TUMOR SUMMARY			
TCTAL ANIMALS WITH PRIMARY TUMORS* TOTAL FRIMARY TUMORS	5 5	14 15	20 24
TOTAL ANIMALS WITH BENIGN TUMCRS TOTAL BENIGN TUMORS		4 4	6 6
TOTAL ANIMALS WITH MALIGNANT TUMORS	s 5 5	1 1 11	17 18
TCTAL ANIMALS WITH SECONDARY TUMORS TOTAL SECONDARY TUMORS	5#	2 2	2 2
TOTAL ANIMALS WITH TUMORS UNCERTAIN BENIGE OR MALIGNANT TOTAL UNCERTAIN TUMORS	N -		
TOTAL ANIMALS WITH TUMORS UNCERTAIN FRIMARY CP METASTATIC TOTAL UNCERTAIN TUMORS	N -		
* PRIMARY TUMORS: ALL TUMORS EXCEPT S * SECONDARY TUMORS: METASTATIC TUMORS			DJACENT ORGAN



APPENDIX C

SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN RATS ADMINISTERED 3-(CHLOROMETHYL)PYRIDINE HYDROCHLORIDE

BY GAVAGE

TABLE C1.

SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN MALE RATS ADMINISTERED 3(CHLOROMETHYL)PYRIDINE HYDROCHLORIDE BY GAVAGE

	VEHICLE	LOW DOSE	HIGH DOSE
ANIMALS INITIALLY IN STUDY ANIMALS NECROPSIED ANIMALS EXAMINED HISTOPATHOLOGICALLY	20 20 20 20	50 50 49	5 0 50 50
INTEGUMENTARY SYSTEM			
NONE			
RESFIRATORY SYSTEM			
LUNG/ERONCHUS POLYFOID HYPERPLASIA	(20)	(49) 1 (2%)	(49)
*LUNG CONGESTION, NOS EDEMA, NOS	(20)	(49)	(49) 7 (14%) 1 (2%)
HEMORRHAGE PNEUMONIA, CHRONIC MURINE GRANULOMA, FOREIGN EODY HYPERPLASIA, ADENOMATOUS		1 (2%) . 2 (4%)	1 (2%) 1 (2%) 1 (2%)
EMATOPOIETIC SYSTEM			
SPLEEN INFARCT, NOS	(20)	(48) 1 (2%)	(48)
HEMOSIDEROSIS HEM ATOPOIESIS	1 (5%)		1 (2%)
*MESENTERIC L. NODE HISTIOCYTOSIS	(19)	(47) 1 (2%)	(43)
CIRCULATORY SYSTEM			
*HEART CALCIFICATION, NOS	(19)	(48)	(46) 1 (2%)
* MY OC A RDI UM INFIAMMATION, FOCAL	(1 9)	(48)	(46)

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

	VEHICLE CONTROL	LOW DOSE	HIGH DOSE
INFLAMMATION, CHRONIC FOCAL FIEROSIS FIEROSIS, FOCAL	1 (5%) 2 (11%)	6 (13%) 3 (6%) 4 (8%)	2 (4%) 1 (2%) 3 (7%)
*BLOOE VESSEL CALCIFICATION, NOS	(20)	(50)	(50) 1 (2%)
*ARTERY INFLAMMATION, CHRONIC	(20)	(50)	(50) 1 (2%)
*AORTA INFLAMMATION, CHRONIC	(20)	(50) 1 (2%)	(50)
DIGESTIVE SYSTEM			
#SALIVARY GLAND INFLAMMATION, NOS FIBROSIS, DIFFUSE	(19)	(48)	(44) 1 (2%) 1 (2%)
#IIVER CONGESTION, PASSIVE INFIAMMATION, ACUTE FOCAL INFIAMMATION, GRANUIOMATOUS CIRRHOSIS, NUS METAMORPHOSIS FATTY	(19) 1 (5%) 2 (11%)	(48) 1 (2%) 1 (2%)	(48) 2 (4%)
EOSINOPHILIC CYTO CHANGE	1 (5%)		
*HEFATIC CAPSULE HYFFPPIASIA, FOCAL	(19)	(48) 1 (2%)	(48)
*BILE LUCT HYPERPLASIA, NOS	(20) 17 (85%)	(50) 34 (68%)	(50) 22 (44%)
*PANCREAS ATRCPHY, NOS	(20)	(48)	(49) 1 (2%)
#FANCRFATIC ACINUS ATRCFHY, NGS ATROPHY, FOCAL	(20)	(48) 1 (2%) 2 (4%)	(49)
#ESOPHAGUS INFLAMMATION, ACUTE	(20)	(47) 1 (2%)	(44)
#SICMACH DILATATION, NOS	(19)	(47)	(50) <u>1 (2%)</u>

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

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	VEHICLE CONTROL	LOW DOSE	HIGH DOSE
HYFEFPLASIA, EPITHELIAL		1 (2%)	2 (4%)
GASTRIC MUCOSA DILATATION, NOS	(19) 12 (63%)	(47) 23 (49%)	(50) 15 (30%)
COLON PARASITISM	(20) 14 (70%)	(47) 18 (38%)	(48) 10 (21%)
URINARY SYSTEM			
• KIDNEY CONGESTION, NOS GLOMERULONEPHRITIS, NOS IN FLAMMATION, CHRONIC NEPHROSIS, HEMOGLOEINURIC CALCIFICATION, NOS	(20) 12 (60%)	(49) 28 (57%)	(49) 6 (12%) 13 (27%) 1 (2%) 1 (2%) 1 (2%)
<pre>*KIDNEY/TUBULE NECROSIS, NOS NECROSIS, FOCAL</pre>	(20) 1 (5%)	(49)	(49) 1 (2%)
ENDOCRINE SYSTEM			
<pre>#PITUITARY CYST, NOS HYPERPIASIA, CHROMCPHOBE-CELL</pre>	(19)	(42) 1 (2%) 2 (5%)	(41) 1 (2%)
<pre>#ADRENAL CYTCPLASMIC VACUOLIZATICN HYPERPLASIA, NODULAR ANGIECTASIS</pre>	(19) 1 (5%)	(49) 1 (2%) 6 (12%)	(49) 2 (4%) 1 (2%) 2 (4%)
*THYROID CYSTIC FOLLICLES INFLAMMATION, CHRONIC FCCAL HYPERPLASIA, C-CELL	(17) 2 (12%)	(39) 1 (3%)	(41) 1 (2%) 1 (2%) 1 (2%)
* PA RA THYRCID HYPERPLASIA, ADENOMATOUS	(8) 1 (13%)	(19)	(17) -
REPRODUCTIVE SYSTEM			
* MA MMA RY GLAND	(20)	(50) <u>1 (2%)</u>	(50)

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

E I

	VEHICLE		
	CONTROL	LOW DOSE	HIGH DOSE
# FROSTATE	(17)	(43)	(41)
INFLAMMATION, SUPPURATIVE INFLAMMATION, ACUTE FOCAL HYPERPLASIA, NOS	1 (6%)		1 (2%) 1 (2%)
#TESTIS TORSION	(19)	(46) 1 (2%)	(47)
ATROPHY, NOS HYPERPLASIA, INTERSTITIAL CELL	2 (11%)	4 (9%) 1 (2%)	1 (2%)
#TESTIS/TUBULE CALCIFICATION, NOS	(19)	(46) 1 (2%)	(47)
ERVOUS SYSTEM			
NONE			
PECIAL SENSE ORGANS			
NONE			
USCULOSKELETAL SYSTEM			
NCNE			
ODY CAVITIES			
*MESENTERY	(20)	(50)	(50)
PERIARIERITIS	1 (5%)		1 (2%)
NECROSIS, FAT			
LL OTHER SYSTEMS			
LL OTHER SYSTEMS NONE			
LL OTHER SYSTEMS			

	VEHICLE CONTROL	LOW DOSE	HIGH DOSE
AUIC/NECROPSY/HISTC PERF AUIC/NECROPSY/NO HISTO		2 1	1
# NUMBER OF ANIMALS WITH TISSUE EXA	MINED MICROSCOP	ICALLY	

* NUMBER OF ANIMALS NECROPSIED

TABLE C2.

SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN FEMALE RATS ADMINISTERED 3(CHLOROMETHYL)PYRIDINE HYDROCHLORIDE BY GAVAGE

	VEHICLE CONTROL	LOW DOSE	HIGH DOSE
ANIMALS INITIALLY IN STUDY ANIMALS NECROPSIED ANIMALS EXAMINED HISTOFATHOLOGICALLY	20 20 20	a50 49 49	5 0 50 50
NTEGUMENTARY SYSTEM			
*SUBCUT TISSUE EPIDERMAL INCLUSION CYST	(20)	(49) 1 (2%)	(50)
RESPIRATORY SYSTEM			
*TRACHEA INFLAMMATION, CHRONIC HYPERPLASIA, EPITHELIAL METAPLASIA, SQUAMOUS	(17)	(44) 1 (2%) 1 (2%)	(41) 1 (2%)
*LUNG CONGESTION, NOS ABSCESS, NOS PNEUMONIA, CHRONIC MURINE INFLAMMATION, GRANULOMATOUS GRANULOMA, NOS GRANULOMA, FOREIGN BODY HYPERPLASIA, ADENOMATOUS HYPERPLASIA, ALVEOLAR EPITHELIUM HISTIOCYTOSIS LEUKEMOID REACTION	(20) 1 (5%) 1 (5%) 1 (5%)	(48) 1 (2%) 1 (2%) 1 (2%) 1 (2%) 2 (4%) 1 (2%) 2 (4%)	(50) 8 (16% 2 (4%) 1 (2%) 1 (2%) 1 (2%)
HEMATOPOIETIC SYSTEM #SPLEEN HEMATOMA, OKGANIZFD FIEROSIS, FOCAL INFARCT, NOS HEMOSIDFROSIS HEMATOPOIESIS	(20) 1 (5%)	(48) 1 (2%) <u>2 (4%)</u>	(49) 1 (2%) 1 (2%)

* NUMBER OF ANIMALS NECROPSIED

 ω 50 ANIMALS WERE INITIALLY IN THE STULY, BUT ONE ANIMAL WAS FOUND TO BE A MALE IN A FEMALE GROUP.

TABLE C2. FEMALE RAT	: NONNEOPLASTIC LESIONS (CONTINUED)
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		LOW DOSE	HIGH DOSE
		•••••	
IRCULATORY SYSTEM			
*MYOCARDIUM	(20)	(47)	(49)
INFLAMMATION, ACUTE POCAL			1 (2%)
INFLAMMATION, CHRONIC FOCAL FIEROSIS, FOCAL	4 (20%) 1 (5%)	6 (13%) 1 (2%)	8 (16%) 1 (2%)
	(() //)		
IGESTIVE SYSTEM			
*LIVER	(20)	(48)	(49)
CONGESTION, NOS			1 (2%)
CONGESTION, CHRONIC PASSIVE INFLAMMATION, NECROTIZING		1 (2%) 1 (2%)	
GRANULOMA, NOS		1 (2%)	
NECROSIS, FOCAL		1 (2%)	2 (4%)
METAMORPHOSIS FATTY BASOPHILIC CYTO CHANGE		1 (2%)	1 (2%)
* EILE DUCT INFLAMMATION, CHRONIC	(20)	(49) 1 (2%)	(50)
HYPERPLASIA, NOS	5 (25%)	21 (43%)	11 (22%)
* PA NC R EAS	(20)	(46)	(49)
INFLAMMATION, INTERSTITIAL	(20)	1 (2%)	(4))
* PA NC REATIC ACINUS	(20)	(46)	(49)
ATRCFHY, NOS		1 (2%)	1 (2%)
ATROPHY, FOCAL	1 (5%)	1 (2%)	2 (4%)
#GASTRIC MUCOSA	(20)	(45)	(48)
DILATATION, NOS	12 (60%)	27 (60%)	23 (48%)
*COLON	(20)	(46)	(49)
PARASITISM	7 (35%)	17 (37%)	10 (20%)
PINIPY CHOMPH			
RINARY SYSTEM			
# KIDNEY	(19)	(49)	(50)
CONGESTION, NOS GLOMERULONEPHRITIS, NOS	4 (21%)	4 (8%)	7 (14%) 3 (6%)
			5 (07)
#KIINEY/CORTEX	(19)	(49) 1 (2%)	(50)

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

	VEHICLE CONTROL	LOW DOSE	HIGH DOSE
#KIDNEY/TUBULE NECRCSIS, NOS NECROSIS, FOCAL	(19) 1 (5%)	(49) 1 (2%)	(50)
ENDOCRINE SYSTEM			
*PITUITARY CYST, NOS NECROSIS, FOCAL HYPERPLASIA, CHROMOPHOBE-CELL ANGIECTASIS	(19) 1 (5%)	(43) 4 (9%) 1 (2%) 1 (2%) 1 (2%)	(40) 1 (3%)
<pre>#ADRENAL CYTCPLASMIC VACUOLIZATICN ANGIECTASIS HEMATOPOIESIS</pre>	(19) 1 (5%)	(49) 4 (8%) 3 (6%)	(50) 3 (6%) 1 (2%)
<pre>#THYROID INFLAMMATION, CHRONIC FCCAL HYPERPLASIA, C-CELL HYPERPLASIA, FOLLICULAR-CELL</pre>	(17) 1 (6%)	(45) 2 (4%)	(38) 3 (8%) 1 (3%)
REPRODUCTIVE SYSTEM			
*MAMMARY GLAND CYST, NOS CYSTIC DUCTS INFLAMMATION, CHRONIC HYPERPLASIA, NOS	(20) 1 (5%) 1 (5%) 1 (5%)	(49)	(50) 1 (2%)
*MAMMARY DUCT Hyperplasia, NOS	(20)	(49)	(50) 1 (2%)
#UTERUS HY DROMETRA	(19)	(47) 1 (2%)	(49) 2 (4%)
*CERVIX UTERI CYST, NOS	(19) 1 (5%)	(47)	(49)
#UTERUS/ENDOMETRIUM ABSCESS, NOS HYPERPLASIA, CYSTIC	(19)	(47)	(49) 1 (2%) 1 (2%)

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

TABLE C2. FEMALE RATS: NONNEOPLASTIC	LESIONS (CONTINUED)
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	VEHICLE CONTROL	LOW DOSE	HIGH DOSE
OVARY CYST, NOS	(18) 1 (6%)	(45) 3 (7%)	(48) 2 (4%)
NERVOUS SYSTEM			
<pre>#BRAIN HEMCRRHAGE</pre>	(19)	(49) 1 (2%)	(50)
PECIAL SENSE ORGANS			
* EYE HE MOBR HAGE	(20)	(49) 1 (2%)	(50)
USCULOSKELETAL SYSTEM			
NONE			
BODY CAVITIES			
*AB COMINAL CAVITY Abscess, Nos Necrosis, Nos	(20)	(49)	(50) 1 (2%) 1 (2%)
*EPICARDIUM INPLAMMATION, FIBRINOUS	(20)	(49) 1 (2%)	(50)
LL OTHER SYSTEMS			
SITE UNKNOWN NECRCSIS, FAT		1	
PECIAL MORPHOLOGY SUMMARY			
NO LESION REPORTED	1	2	8

* NUMBER OF ANIMALS NECROPSIED



APPENDIX D

SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN MICE ADMINISTERED 3-(CHLOROMETHYL)PYRIDINE HYDROCHLORIDE

BY GAVAGE

TABLE D1.

SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN MALE MICE ADMINISTERED 3(CHLOROMETHYL)PYRIDINE HYDROCHLORIDE BY GAVAGE

	VEHICLE CONTROL	LOW DOSE	HIGH DOSE
ANIMALS INITIALLY IN STUDY	20	50	50
ANIMALS MISSING ANIMALS NECFOPSIED ANIMALS EXAMINED HISTOPATHOLOGICALLY	20 20	2 48 48	50 50
INTEGUMENTARY SYSTEM			
NONE			
RESPIRATCRY SYSTEM			
#IUNG CONGESTION, NOS	(20)	(45) 2 (4%)	(48) 7 (15%)
ERCNCHCPNEUMONIA, ACUTE PNEUMONIA, CHRONIC MURINE PERIVASCULAR CUFFING	1 (5%) 3 (15%) 1 (5%)	2 (4%) 1 (2%)	1 (2%)
HYPERPLASIA, ADENOMATOUS HISTICCYTOSIS	1 (5%)	(2%)	2 (4%) 3 (6%)
LEUKEMOID REACTION HYPERPLASIA, BASOPHILIC		1 (2%)	1 (2%)
* LUNG/ALVEOLI	(20)	(45)	(48)
CRYSTALS, NOS HISTIOCYTOSIS	1 (5%)		1 (2%)
HEMATOPOIETIC SYSTEM			
* SPLEEN	(17)	(41)	(41)
A MYLCICOJIS AN GIECTASIS		1 (2%)	1 (2%)
HEMATOPOIESIS		3 (7%)	1 (2%)
#MESENTERIC L. NODE CONGESTION, NOS	(16)	(33) 1 (3%)	(35)
HEMORR EAGE		4 (12%)	1 (3%)
HEMOSIDEROSIS HISTIOCYTOSIS		1 (3%) 1 (3%)	1 (3%)
MASTOCYTOS IS		<u> </u>	

NUMBER OF ANIMALS WITH TISSUE EXAMINED MICROSCOPICALLY
 NUMBER OF ANIMALS NECROPSIED

	VEHICLE CONTROL	LOW DOSE	HIGH DOSE
CIRCULATORY SYSTEM			
*PORTAL VEIN THROMBOSIS, NOS	(20)	(48) 1 (2%)	(50)
IGESTIVE SYSTEM			
#SALIVARY GLAND PERIVA SCULAR CUFFING	(19) 1 (5%)	(39) 2 (5%)	(44) 2 (5%)
<pre>#LIVER CYST, NOS CONGESTION, NOS HEMATOMA, NOS INFLAMMATION, FOCAL INFLAMMATION, CHRONIC INFLAMMATION, CHRONIC FOCAL GRANULATION, TISSUE THROM BOPHLEBITIS FIEROSIS PERIVASCULAR CUFFING</pre>	(20) 1 (5%) 1 (5%) 1 (5%) 1 (5%) 1 (5%)	(46) 1 (2%) 1 (2%) 1 (2%) 4 (9%) 1 (2%) 1 (2%)	(49) 1 (2 %)
NECROSIS, FOCAL INFARCT, NOS CALCIUM DEPOSIT CALCIFICATION, NOS BASOPHILIC CYTO CHANGE ATROPHY, NOS	1 (5%) 1 (5%)	1 (2%) 2 (4%) 1 (2%) 1 (2%)	1 (2%)
#LIVER/HEPATOCYTES CYTCPLASMIC CHANGE, NOS	(20)	(46) 1 (2%)	(49)
*PANCFEAS INFLAMMATION, INTERSTITIAL	(18)	(33)	(36) 1 (3%)
# FA NC REATIC DUCT NECRCSIS, NOS	(18) 1 (6%)	(33)	(36)
# FA NC REATIC ACINUS NEC RCSIS, NOS ATROPHY, NOS	(18) 1 (6%) 1 (6%)	(33)	(36)
#STCMACH CYST, NOS	(19)	(43)	(47) 1 (2%)

TABLE D1. MALE MICE: NONNEOPLASTIC LESIONS (CONTINUED)

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

Da,

	VEHICLE	LOW DOSE	HIGH DOSE
INFLAMMATION, NOS Hyperplasia, epithelial		1 (2%)	1 (2%) 2 (4%)
PEYERS PATCH HYFEPPIASIA, NOS	(17) 1 (6%)	(42) 2 (5%)	(42)
COLON PARASITI SM	(18) 3 (17%)	(44) 12 (27%)	(47) 13 (28%)
RINARY SYSTEM			
*KIDNEY CONGESTION, NOS PERIVASCULAR CUFFING	(20) 3 (15%)	(46) 2 (4%) 7 (15%)	(47) 7 (15%) 5 (11%)
*KIDNEY/MEDULLA CYST, NOS	(20)	(46)	(47) 1 (2%)
* FERIFENAL TISSUE NECROSIS, FOCAL	(20)	(46) 1 (2%)	(47)
* KI DN EY/TUB UL E HYPERPLASIA, FOCAL	(20) 1 (5%)	(46)	(47)
* KI DN EY/PEL VI S IN FIAMMATION, CHRONIC PERIVA SCULAR CUFFING	(20)	(46) 1 (2%)	(47) 1 (2%)
ULINARY BLADDER MUC OC ELE	(13)	(33) 1 (3%)	(37)
INFLAMMATION, CHRONIC PERIVASCULAR CUFFING		2 (6%)	1 (3%)
NDOCRINE SYSTEM			
NONE			
EPRODUCTIVE SYSTEM			
PROSTATE INFLAMMATION, CHRONIC SUPPURATI	(19) V	(40)	(46) <u>1 (2%)</u>

* NUMBER OF ANIMALS NECROPSIED

	VEHICLE CONTROL	LOW DOSE	HIGH DOSE
NERVOUS SYSTEM			
#BRAIN INFARCT, NOS CALCIFICATION, FOCAL	(20) 6 (30%)	(46) 7 (15%)	1 (2%)
SPECIAL SENSE ORGANS NONE			
MUSCULOSKEIETAL SYSTEM NONE			
BODY CAVITIES NONE			
ALL OTHER SYSTEMS NONE			
SPECIAL MORPHOLOGY SUMMARY			
NO LESION REPORTED ANIFAL MISSING/NO NECROPSY AUTC/NECROPSY/HISTO PERF	3 1	11 2	12
<pre># NUMBER OF ANIMALS WITH TISSUE EX * NUMBER OF ANIMALS NECROPSIED</pre>	AMINED NICROSCOPI	CALLY	

TABLE D2.

SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN FEMALE MICE ADMINISTERED 3(CHLOROMETHYL)PYRIDINE HYDROCHLORIDE BY GAVAGE

	VEHICLE CONTROL	LOW DOSE	HIGH DOSE
ANIMALS INITIALLY IN STUDY ANIMALS MISSING ANIMALS NECROPSIED ANIMALS FXAMINED HISTOFATHOLOGICALLY	20 20 20	50 50 50	50 1 49 49
IN TEGUMENTARY SYSTEM NONE			
RESPIRATORY SYSTEM	(10)	(50)	(#0)
*LUNG/BRCNCHUS CRYSTAIS, NOS	(18)	1 (2%)	(49)
*LUNG AT ELECTASIS CONGESTION, NOS PNEUMONIA, CHRONIC MURINE INFLAMMATION, GRANULOMATOUS PERIVASCULAR CUFFING HISTICCYTOSIS	(18) 1 (6%) 3 (17%)	(50) 1 (2%) 11 (22%)	(49) 8 (16%) 3 (6%) 1 (2%) 1 (2%) 1 (2%)
*IUNG/AIVEOLI CRYSTAIS, NOS PHAGOCYTIC CELL	(18)	(50) 1 (2%) 1 (2%)	(49)
HEMATOPOIETIC SYSTEM			
<pre>#BONE MARROW HYFEPPIASIA, HEMATCFOIETIC</pre>	(9)	(46) 3 (7%)	(46) 1 (2%)
*SPIEEN HYPERPIASIA, RETICULUM CELL HEMATOPOIESIS	(17) 4 (24%)	(46) 6 (13%)	(45) 2 (4%) 2 (4%)
#MANDIBULAR L. NODE INFIAMMATION, GRANULOMATCUS	(16)	(42) 1 (2%)	(40)
*BRCNCHIAL LYMPH NODE HYFERPIASIA, NOS	(16)	(42) <u>1_(2%)</u>	(40)

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

	VEHICLE CONTROL	LOW DOSE	HIGH DOSE
#MESENTERIC L. NODE HEMCFRHAGE INFIAMMATION, GRANULCMATOUS CYTOMEGALY HYPERFLASIA, RETICULUM CEIL HEMATOPOIESIS	(16)	(42) 1 (2%) 1 (2%) 1 (2%) 1 (2%) 1 (2%)	(40) 1 (3%)
CIRCULAIORY SYSTEM			
#HEART CALCIFICATION, FOCAL	(18) 1 (6%)	(48)	(48)
*BLCOE VESSEL INFIAMMATION, CHRONIC THROMBOPHLEBITIS	(20)	(50) 1 (2%)	(49) 1 (2%)
*CCRONARY ARTERY INFLAMMATION, CHRONIC	(20)	(50)	(49) 1 (2%)
*UTERINE VEIN THROMBOSIS, NOS	(20) 1 (5%)	(50)	(49)
#HEFATIC SINUSOID THPCMBOSIS, NOS	(20)	(50)	(49) 1 (2%)
DIGESTIVF SYSTEM			
#SALIVARY GLAND INFIAMMATION, CHRONIC FOCAL PERIVA SCULAR CUFFING	(19) 2 (11%)	(48) 1 (2%) 1 (2%)	(44) 2 (5%)
#LIVER INFIAMMATION, NOS	(20)	(50) 1 (2%)	(49)
INFLAMMATION, CHRONIC FOCAL INFLAMMATION, GRANULOMATOUS	2 (10%)	1 (2%)	1 (2%) 1 (2%)
PERIVASCULAR CUFFING DEGENERATION, NOS	1 (5%) 1 (5%)	3 (6%)	1 (2%)
NECROSIS, FOCAL INFARCT, NOS METAMORPHOSIS FATTY	1 (5%) 1 (5%)	1 (2%) 1 (2%)	1 (2%)
HYPERPLASIA, NOS ANGIECTASIS	1 (5%)		1 (2%)

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

	VEHICLE CONTROL	LOW DOSE	HIGH DOSE
HISTIOCYTOSIS	1 (5%)		
*LIVER/CENTRILOBULAR NECRCSIS, NOS	(20)	(50) 2 (4%)	(49)
*BILE FUCT INFLAMMATION, CHRONIC INFLAMMATION, CHRONIC FOCAL	(20)	(50) 1 (2%) 1 (2%)	(49) 1 (2%)
* PANCREAS CYSTIC DUCTS	(13)	(34) 2 (6%)	(38)
PANCRFATIC ACINUS INFLAMMATION, CHRONIC ATROPHY, NOS ATROPHY, POCAL	(13)	(34) 1 (3%) 1 (3%) 1 (3%)	(38) 1 (3%)
#STOMACH CYST, NOS ABSCESS, NOS CALCIFICATION, FOCAL HYPERPLASIA, EPITHELIAL MASIOCYTOSIS	(19)	(45)	(48) 1 (2%) 1 (2%) 1 (2%) 3 (6%) 1 (2%)
#GASTRIC SUBMUCOSA INFLAMMATION, ACUTE/CHRONIC	(19)	(45)	(48) 1 (2%)
#COLON PARASITISM	(19) 3 (16%)	(45) 7 (16%)	(47) 4 (9%)
URINARY SYSTEM			
<pre># KIDNEY CONGESTION, NOS INFIAMMATION, CHRONIC PERIVASCULAR CUFFING NEPHROSIS, NOS</pre>	(19) 1 (5%)	(49) 2 (4%) 1 (2%) 5 (10%) 1 (2%)	(49) 9 (18%) 1 (2%) 4 (8%)
AMYLOIDOSIS METAPLASIA, OSSEOUS		1 (2%)	1 (2%)
<pre>#KIDNEY/TUBULE CALCIFICATION, NOS ATROPHY, NOS REGENERATION, NOS</pre>	(19)	(49) 1 (2%) 1 (2%) 1 (2%)	(49)

NUMBER OF ANIMALS WITH TISSUE EXAMINED MICROSCOPICALLY
 NUMBER OF ANIMALS NECROPSIED

	VEHICLE CONTROL	LOW DOSE	HIGH DOSE
#URINARY BLADDER	(15)	(29)	(36)
INFLAMMATION, CHRONIC PERIVASCULAR CUFFING	1 (7%)	2 (7%)	1 (3%)
LYMPHOCYTOSIS			1 (3%)
NDOCRINE SYSTEM			
#THYROID	(8)	(15)	(36)
HYPERCHROMATISM		1 (7%)	
EPRODUCTIVF SYSTEM			
#UT ERU S	(18)	(47)	(47)
CYST, NOS Pyometra	4 (22%)	1 (2%) 2 (4%)	3 (6%)
FIONEINA		2 (4%)	
#UTERUS/ENDOMETRIUM CYST, NOS	(18)	(47) 4 (9%)	(47) 1 (2%)
INFLAMMATION, SUPPURATIVE		1 (2%)	1 (2%)
ABSCESS, NOS	1 (6%)		
INFLAMMATION, CHRONIC HYPERPLASIA, CYSTIC	4 (22%)	1 (2%) 9 (19%)	6 (13)
#OVARY/CVIDUCT INFLAMMATION, NOS	(18)	(47) 1 (2%)	(47)
HYPERPLASIA, NOS		1 (2%)	
# OV AR Y	(6)	(13)	(33)
CYST, NOS	1 (17%)	5 (38%)	4 (12)
HEMORRHAGIC CYST			2 (6%)
ERVOUS SYSTEM			
#BRAIN	(18)	(50)	(49)
PERIVA SCULITIS CORPORA AMYLACEA	1 (6%)		1 (2%)
CALCIFICATION, NOS	1 (6%)		1 (2%)
CALCIFICATION, FOCAL	2 (11%)	9 (18%)	7 (149
# MIDBRAIN	(18)	(50)	(49)
CALCIFICATION, FOCAL			1 (2%)

NONE

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

	VEHICLE CONTROL	LOW DOSE	HIGH DOSE
USCULCSKELETAL SYSTEM			
*BONE PIERCUS OSIEODYSTRCPHY OSTEOSCLEROS IS	(20) 7 (35%)	(50) 22 (44系) 1 (2系)	(49) 18 (37%)
ODY CAVITIES			
* MEDIASTINUM INFLAMMATION, CHRONIC	(20) 1 (5落)	(50)	(49)
*PERITCHEAL CAVITY GRANULCHA, NOS	(20)	(50) 1 (2%)	(49)
* FLEUEA IN FLAMMATION, CHEONIC IN FLAMMATION, CHRONIC FOCAL	(20) 1 (5%)	(50)	(49) 1 (2%)
LL OTHER SYSTEMS			
NCNE			
PECIAL MORPHOLOGY SUMMARY			
NO LESION EEPORTED ANIMAL MISSING/NO NECROPSY AUTC/NECROPSY/HISTO PERF	1	2	2 1 1

* NUMBER OF ANIMALS NPCROPSIED



APPENDIX E

ANALYSES OF THE INCIDENCE OF PRIMARY TUMORS IN RATS ADMINISTERED 3-(CHLOROMETHYL)PYRIDINE HYDROCHLORIDE

BY GAVAGE

Table El. Analyses of the Incidence of Primary Tumors in Male Rats Administered 3-(Chloromethyl)pyridine Hydrochloride by Gavage^a

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in Male Rats by Gavage^a

Table El. Analyses Administered 3-(Ch	ble El. Analyses of the Incidence of Primary Tumors Administered 3-(Chloromethyl)pyridine Hydrochloride	Primary Tumors Hydrochloride b
(continued)		
Topography: Morphology	Vehicle Control	Low Dose
Stomach: Squamous-cell Papilloma or Carcinoma ^b	0/19 (0)	1/47 (2)
P Values ^{c,d}		N. S.
Relative Risk ^e Lower Limit Upper Limit		Infinite 0.022 Infinite
Weeks to First Observed Tumor		67
Pituitary: Chromophobe Carcinoma ^b	0/19 (0)	3/42 (7)
P Values ^{c,d}		N. S.
Relative Risk ^e Lower Limit Upper Limit		Infinite 0.284 Infinite
Weeks to First Observed Tumor		89

Infinite 0.238 Infinite

3/50 (6) N.S.

High Dose

N.S. Infinite 0.026 Infinite

104

1/41 (2)

76

		against far anti-strings	
(continued)			
	Vehicle	Low	High
Topography: Morphology	Control	Dose	Dose
Pituitary: Chromophobe Adenoma or Carcinoma ^b	1/19 (5)	7/42 (17)	2/41 (5)
P Values ^{c,d}		N. S.	N • S •
Relative Risk ^e Lower Limit Upper Limit		3.167 0.460 138.815	0.927 0.052 53.355
Weeks to First Observed Tumor	104	86	104
Adrenal: Pheochromocytoma ^b	2/19 (11)	2/49 (4)	1/49 (2)
P Valuesc,d		N.S.	N • S •
Relative Risk ^e Lower Limit Upper Limit		0.388 0.031 5.108	0.194 0.003 3.563
Weeks to First Observed Tumor	104	104	101

Table El. Analyses of the Incidence of Primary Tumors in Male Rats Administered 3-(Chloromethyl)pyridine Hydrochloride by Gavage^a

IADIE EI. ANALYSES Administered 3-(Ch	yses of the incluence of Frimary lumors in Male K 3-(Chloromethyl)pyridine Hydrochloride by Cavage ^a	Analyses of the incluence of rimary lumors in Male Naus red 3-(Chloromethyl)pyridine Hydrochloride by Gavage ^a	re nars 1ge ^a
(continued)	•		
Topography: Morphology	Vehicle Control	Low Dose	High Dose
Adrenal: Cortical Adenoma or Carcinoma ^b	2/19 (11)	1/49 (2)	1/49 (2)
P Values ^{c,d}		N•S•	N.S.
Relative Risk ^e Lower Limit Upper Limit		0.194 0.003 3.563	0.194 0.003 3.563
Weeks to First Observed Tumor	104	104	104
Thyroid: C-cell Adenoma or Carcinoma ^b	1/17 (6)	2/39 (5)	1/41 (2)
P Values ^{c,d}		N•S•	N. S.
Relative Risk ^e Lower Limit Upper Limit		0.872 0.050 50.118	0.415 0.006 31.786
Weeks to First Observed Tumor	104	104	104

Table El. Analyses of the Incidence of Primary Tumors in Male Rats

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THE WILL

(continued)VehicleTopography:WorphologyPancreatic Islets: Islet-cell Adenomab3/20 (15)P valuesc.d3/20 (15)Relative Risk ^e 3/20 (15)	5)	Low <u>Dose</u> 2/48 (4) N.S.	High <u>Dose</u> 0/49 (0) P = 0.022(N)
<u>Morphology</u> Islets: 1 Adenoma ^b sk ^e	5)	Low <u>Dose</u> /48 (4) N.S.	
Islets: 1 Adenomab sk ^e		/48 (4) N.S.	
S	Ŏ	N.S.	
Relative Risk ^e	Ŏ		
Lower Limit Upper Limit	5.0	0.278 0.025 2.278	0.000 0.000 0.673
Weeks to First Observed Tumor 104	104	104	
Pancreatic Islets: Islet-cell Adenoma or Carcinoma ^b 3/20 (15)		3/48 (6)	0/49 (0)
P Values ^c ,d		N • S •	P = 0.022 (N)
Relative Risk ^e Lower Limit Upper Limit		0.417 0.062 2.915	0.000 0.000 0.673
Weeks to First Observed Tumor 104	104	103	-

Table E1. Analyses of the Incidence of Primary Tumors in Male Rats Administered 3-(Chloromethyl)pyridine Hydrochloride by Gavage^a

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Table El. Anslyses of the Incidence of Primary Tumors in Male Rats Administered 3-(Chloromethyl)pyridine Hydrochloride by Gavage^a

	High <u>Dose</u>	20/47 (43)	P < 0.001(N)	0.476 0.394 0.728	76
	Low Dose	38/46 (83)	N • S •	0.923 0.800 1.235	67
	Vehicle <u>Control</u>	17/19 (89)			104
(continued)	Topography: Morphology	Testis: Interstitial-cell Tumor ^b	P Values ^c ,d	Relative Risk ^e Lower Limit Upper Limit	Weeks to First Observed Tumor

^aDosed groups received 75 or 150 mg/kg by gavage.

90

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

^cBeneath the incidence of tumors in a dosed group is the probability level for the Fisher exact test for the comparison of that dosed group with the control group when P < 0.05; otherwise, not significant (N.S.) is indicated.

^dA negative trend (N) indicates a lower incidence in a dosed group than in a control group.

^eThe 95% confidence interval of the relative risk between each dosed group and the control group.

Table E2. Analyses of the Incidence of Primary Tumors in Female Rats Administered 3-(Chloromethyl)pyridine Hydrochloride by Gavage^a i

YUUTISLEEG DJJSTUTUS	טרטווופרוואד/אערזמדוופ ה	ασπιπιςτετεα 3-(υπιστοπειηγι/μγιμμιε πλαισσπιστιαε υγ σαναξε	U
(continued)			
	Vehicle	Low	High
Topography: Morphology	Control	Dose	Dose
Pituitary: Chromophobe Adenoma or Carcinoma ^b	9/19 (47)	12/43 (28)	14/40 (35)
P Values ^c		N • S •	N•S•
Relative Risk ^d		0.589	0.739
Lower Limit Upper Limit		0.293 1.342	0.385 1.619
Weeks to First Observed Tumor	104	85	72
Mammary Gland: Fibroadenoma ^b	2/20 (10)	4/49 (8)	1/50 (2)
P Values ^c		N.S.	N • S •
Relative Risk ^d Tower Timit		0.816 0.131	0.200 0.004
Upper Limit		8.603	3.681
Weeks to First Observed Tumor	104	104	104

Table E2. Analyses of the Incidence of Primary Tumors in Female Rats Administered 3-(Chloromethyl)pyridine Hydrochloride by Gavage^a

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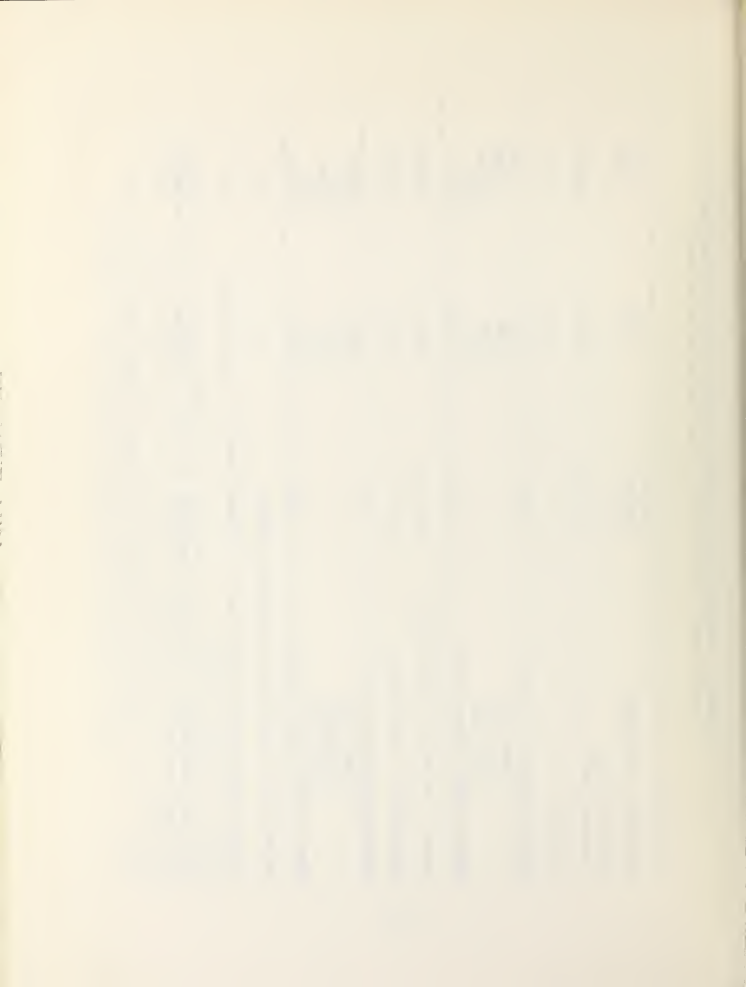
Administered 3-(Chloromethyl)pyridine Hydrochloride	ered 3-(Chloromethyl)pyridine	Hydrochloride by Gavage ^a	gea
(continued)			
Topography: Morphology	Vehicle Control	Low Dose	High Dose
19			
Preputial Gland: Adenoma, NOS ^b	2/20 (10)	1/49 (2)	0/50 (0)
P Values ^c		N • S •	N•S•
Relative Risk ^d Lower Limit Upper Limit		0.204 0.004 3.754	0.000 0.000 1.345
Weeks to First Observed Tumor	104	104	
Uterus: Endometrial Stromal Polyp ^b	2/19 (11)	8/47 (17)	2/49 (4)
P Values ^c		N.S.	N•S•
Relative Risk ^d Lower Limit Upper Limit		1.617 0.370 14.802	0.388 0.031 5.108
Weeks to First Observed Tumor	104	76	104
^a Dosed groups received 75 or 150 mg/kg by gavage.	y gavage.		
^b Number of tumor-bearing animals/number of animals	of animals examined	d at site (percent).	

Analyses of the Incidence of Primary Tumors in Female Rats

Table E2.

^cBeneath the incidence of tumors in a dosed group is the probability level for the Fisher exact test for the comparison of that dosed group with the control group when P < 0.05; otherwise, not significant (N.S.) is indicated.

dThe 95% confidence interval of the relative risk between each dosed group and the control group.



APPENDIX F

ANALYSES OF THE INCIDENCE OF PRIMARY TUMORS IN MICE ADMINISTERED 3-(CHLOROMETHYL)PYRIDINE HYDROCHLORIDE

BY GAVAGE



Topography: Morphology	Vehicle <u>Control</u>	Low Dose	High Dose
Lung: Alveolar/Bronchiolar Carcinoma ^b	1/20 (5)	4/45 (9)	5/48 (10)
P Values ^c		N • S •	N• S•
Relative Risk ^d Lower Limit Upper Limit		1.778 0.195 85.520	2.083 0.259 96.358
Weeks to First Observed Tumor	104	84	102
Lung: Alveolar/Bronchiolar Adenoma or Carcinoma ^b	2/20 (10)	4/45 (9)	5/48 (10)
P Values ^c		N. S.	N• S•
Relative Risk ^d Lower Limit Upper Limit		0.889 0.143 9.340	1.042 0.192 10.410
Weeks to First Observed Tumor	104	84	102

Table Fl. Analyses of the Incidence of Primary Tumors in Male Mice Administered 3-(Chloromethyl)pyridine Hydrochloride by Gavage^a

NOV-C DATATITITINA		ααιμτιτειετεα ο-(ουτοτομειάτ/βλετατμε υλαιοουτοιτα ολ σαναβε.	L D D
(continued)			
Topography: Morphology	Vehicle Control	Low Dose	High Dose
0 4	1/20 (5)	(9) 87/8	3/50 (6)
P Values ^c		N • S •	N•S•
Relative Rigk ^d		1.250	1.200 0.106
Upper Limit		0.110 64.251	0.100 61.724
Weeks to First Observed Tumor	73	94	63
Hematopoietic System: Malignant Lymphoma, Lymphocytic Leukemia or Undifferentiated Leukemiab	3/20 (15)	0//8 (10)	0/50 (18)
P Values ^c		N.S.	N• S•
Relative Risk ^d		1. 250	1.200
LOWER LIMIC Upper Limit		0.501 6.662	0.340 6.408
Weeks to First Observed Tumor	73	61	63

Table Fl. Analyses of the Incidence of Primary Tumors in Male Mice Administered 3-(Chloromethyl)pyridine Hydrochloride by Gavage^a

Administered 3-(Cr	iloromethy1)pyridine	Administered 3-(Ghloromethyl)pyridine Hydrochloride by Gavage ^a	264
(continued)			
Topography: Morphology	Vehicle Control	Low Dose	High Dose
Liver: Hepatocellular Adenoma or Carcinoma ^b	3/20 (15)	5/46 (11)	9/49 (18)
P Values ^c		N. S.	N. S.
Relative Risk ^d Lower Limit Upper Limit		0.725 0.160 4.348	1.224 0.354 6.533
Weeks to First Observed Tumor	94	58	66
Stomach: Squamous-cell Papilloma ^b	(0) 61/0	2/43 (5)	8/47 (17)
P Values ^c		N • S •	N. S.
Relative Risk ^d Lower Limit Upper Limit		Infinite 0.136 Infinite	Infinite 0.966 Infinite
Weeks to First Observed Tumor		61	58

Table Fl. Analyses of the Incidence of Primary Tumors in Male Mice Administered 3-(Chloromethyl)pyridine Hydrochloride by Gayage^a

Administered 3-(Chlor	Administered 3-(Chloromethyl)pyridine Hydrochloride by Gavage ^a	hloride by Gavage ^a	
(continued)			
	Vehicle	Low	High
Topography: Morphology	Control	Dose	Dose
Stomach: Squamous-cell Papilloma or Carcinoma ^b	0/19 (0)	2/43 (5)	10/47 (21)
P Values ^c		N • S •	P = 0.025
Relative Risk ^d Lower Limit Upper Limit		Infinite 0.136 Infinite	Infinite 1.259 Infinite
Weeks to First Observed Tumor		61	58
^a Dosed groups received 100 or 200 mg/kg by gavage.	y gavage.		

Table Fl. Analyses of the Incidence of Primary Tumors in Male Mice

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

^cBeneath the incidence of tumors in a dosed group is the probability level for the Fisher exact test for the comparison of that dosed group with the control group when P < 0.05; otherwise, not significant (N.S.) is indicated.

^dThe 95% confidence interval of the relative risk between each dosed group and the control group.

Administered 3-(Chi	stered 3-(Chloromethyl)pyridine Hydrochloride by Gavage ^a .	drochloride by Gavag	5 D
Topography: Morphology	Vehícle Control	Low Dose	High Dose
Lung: Alveolar/Bronchiolar Carcinoma ^b	1/18 (6)	0/50 (0)	3/49 (6)
P Values ^c		N.S.	N.S.
Relative Risk ^d Lower Limit Upper Limit		0.000 0.000 6.729	1.102 0.098 56.666
Weeks to First Observed Tumor	65		50
Lung: Alveolar/Bronchiolar Adenoma or Carcinoma ^b	1/18 (6)	1/50 (2)	3/49 (6)
P Values ^c		N.S.	N. S.
Relative Risk ^d Lower Limit Upper Limit		0.360 0.005 27.724	1.102 0.098 56.666
Weeks to First Observed Tumor	65	104	50

Table F2. Analyses of the Incidence of Primary Tumors in Female Mice Administered 3-(Chloromethyl)nyridine Hydrochloride by Gayage^a

(continued)			
Topography: Morphology	Vehicle Control	Low Dose	High Dose
Hematopoietic System: Malignant Lymphoma, Lymphocytic Leukemia or Undifferentiated Leukemia ^b	2/20 (10)	9/50 (18)	9/49 (18)
P Values ^c		N • S •	N. S.
Relative Risk ^d Lower Limit Upper Limit		1.800 0.426 16.255	1.837 0.434 16.572
Weeks to First Observed Tumor	93	76	98
Hematopoietic System: All Lymphomas or Leukemias ^b	2/20 (10)	9/50 (18)	10/49 (20)
P Values ^c		N • S •	N•S•
Relative Risk ^d Lower Limit Upper Limit		1.800 0.426 16.255	2.041 0.498 18.154
Weeks to First Observed Tumor	93	76	96

1ge4		High Dose	2/29 (7)	N • S •	Infinite 0.103 Infinite	104	3/48 (6)	N• S•	Infinite 0.248 Infinite	104
Administered 3-(Chloromethyl)pyridine Hydrochloride by Gavage ^a		Low Dose	0/25 (0)	N•S•		1	1/45 (2)	N•S•	Infinite 0.023 Infinite	104
LoromethyL)pyridine		Vehicle Control	(0) 6/0			1	(0) 61/0			-
Administered J-(Ch.	(continued)	Topography: Morphology	Pituitary: Chromophobe Adenoma ^b	P Values ^c	Relative Risk ^d Lower Limit Upper Limit	Weeks to First Observed Tumor	Stomach: Squamous-cell Papilloma ^b	P Values ^c	Relative Risk ^d Lower Limit Upper Limit	Weeks to First Observed Tumor

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Table F2. Analyses of the Incidence of Primary Tumors in Female Mice Administered 3-(Chloromerhyl)nyridine Hydrochloride by Gayage^a

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Table F2. Analys Administered 3	le F2. Analyses of the Incidence of Primary Tumors in Female Mice Administered 3-(Chloromethyl)pyridine Hydrochloride by Gavage ^a	Primary Tumors in Fema e Hydrochloride by Gava	ile Mice 1ge ^a
(continued)			
	Vehicle	Low	High
Topography: Morphology	Control	Dose	Dose
Stomach: Squamous-cell Papilloma or Carcinoma ^b	0/19 (0)	1/45 (2)	5/48 (10)
P Values ^c		N • S •	N _• S _•
Relative Risk ^d Lower Limit Upper Limit		Infinite 0.023 Infinite	Infinite 0.522 Infinite
Weeks to First Observed Tumor	I	104	104
^a Dosed groups received 100 or 200	00 or 200 mg/kg by gavage.		
Mumbou of turnor tooring of a	(according to formation of action (according to the formation of the forma	at of a (account)	

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

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^cBeneath the incidence of tumors in a dosed group is the probability level for the Fisher exact test for the comparison of that dosed group with the control group when P < 0.05; otherwise, not significant (N.S.) is indicated.

dThe 95% confidence interval of the relative risk between each dosed group and the control group.



Review of the Bioassay of 3-(Chloromethyl)pyridine Hydrochloride* 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 3-(Chloromethyl)pyridine Hydrochloride for carcinogenicity.

The reviewer agreed with the conclusion in the report that 3-(Chloromethyl)pyridine Hydrochloride was carcinogenic under the conditions of test. After a brief review of the experimental design, the reviewer said that the study was adequate to support the conclusion on the compound's carcinogenicity. The review of the bioassay of 3-(Chloromethyl) pyridine Hydrochloride was accepted 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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