

BIOASSAY OF

ALLYL CHLORIDE

FOR POSSIBLE CARCINOGENICITY

Carcinogenesis Testing Program Division of Cancer Cause and Prevention US, 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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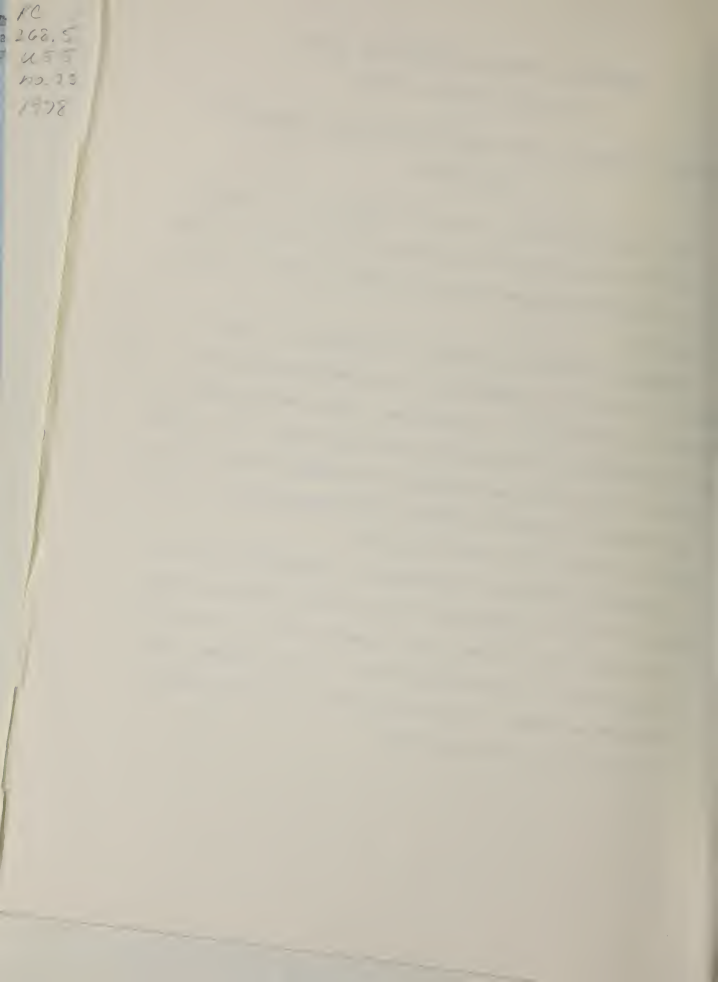
DEPARTMENT OF HEALTH, EDUCATION, AND WELFARE National Institutes of Health

REPORT ON BIOASSAY OF ALLYL CHLORIDE FOR POSSIBLE CARCINOGENICITY Availability

Allyl chloride (CAS 107-05-1) 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 for possible carcinogenicity of technicalgrade allyl chloride (3-chloropropene) was conducted using Osborne-Mendel rats and B6C3F1 mice. Applications of the chemical include use as an intermediate in the synthesis of other chemicals. Allyl chloride in corn oil was administered by gavage to two groups of each species for 5 days a week for 78 weeks, followed by observation periods of 30 to 33 weeks for the rats and 14 weeks for the mice.

Under the conditions of this bioassay no convincing evidence was presented for the carcinogenicity of allyl chloride in Osborne-Mendel rats of either sex. The results are suggestive that allyl chloride is carcinogenic in male and female B6C3F1 mice since the compound, when administered by gavage, caused a low incidence of neoplastic and nonneoplastic lesions of the forestomach.

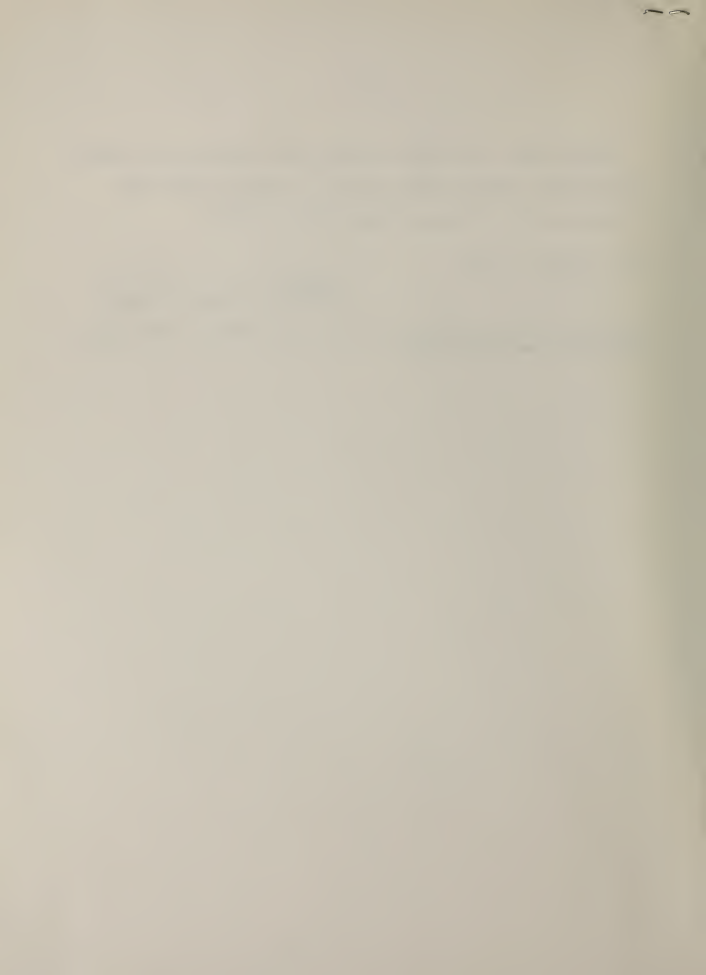


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REPORT ON THE BIOASSAY OF ALLYL CHLORIDE FOR POSSIBLE CARCINOGENICITY

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 allyl chloride 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 allyl chloride was conducted by Hazleton Laboratories America, Inc., Vienna, Virginia, initially under direct contract to the NCI and currently under a subcontract to Tracor Jitco, Inc., prime contractor for the NCI Carcinogenesis Testing Program.

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

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

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

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

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SUMMARY

A bioassay for possible carcinogenicity of technical-grade allyl chloride (3-chloropropene) was conducted using Osborne-Mendel rats and B6C3F1 mice. At initiation of the study the rats were approximately 6 weeks old and the mice approximately 5 weeks old. Allyl chloride in corn oil was administered by gavage to two groups of each species for 5 days a week for 78 weeks, followed by observation periods of 30 to 33 weeks for the rats and 14 weeks for the mice. The time-weighted average dosages were, respectively, 77 and 57 mg/kg/day for high and low dose male rats; 73 and 55 mg/kg/day for high and low dose female rats; 199 and 172 mg/kg/day for high and low dose male mice; and 258 and 129 mg/kg/day for high and low dose female mice.

For each species, 20 animals of each sex were placed on test as vehicle controls. These animals were intubated with corn oil at the same time that dosed animals were gavaged with allyl chloride in corn oil. Twenty animals of each sex were placed on test as untreated controls for each species. These animals were not intubated.

Survival of high dose male mice and high dose rats of both sexes was extremely poor. Fifty percent of the high dose male mice were dead by week 27; the 10 members of this group that survived past week 48 were sacrificed in week 56. Among the high dose rats, 50 percent of the males had died by week 14 and 50 percent of the females had died by week 38. Because of early mortality in these groups, the number of animals surviving long enough to be at risk from late-developing tumors was not adequate for meaningful statistical analysis.

In this bioassay, squamous-cell carcinomas of the forestomach in male and female mice and squamous-cell papillomas of the forestomach in female mice occurred in incidences that were higher than in historical controls. No other neoplasms occurred in statistically significant increased incidences in dosed rats or mice.

Under the conditions of this bioassay no convincing evidence was presented for the carcinogenicity of allyl chloride in Osborne-Mendel rats of either sex. The results are suggestive that allyl chloride is carcinogenic in male and female B6C3F1 mice since the compound, when administered by gavage, caused a low incidence of neoplastic and nonneoplastic lesions of the forestomach.

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WEEKS

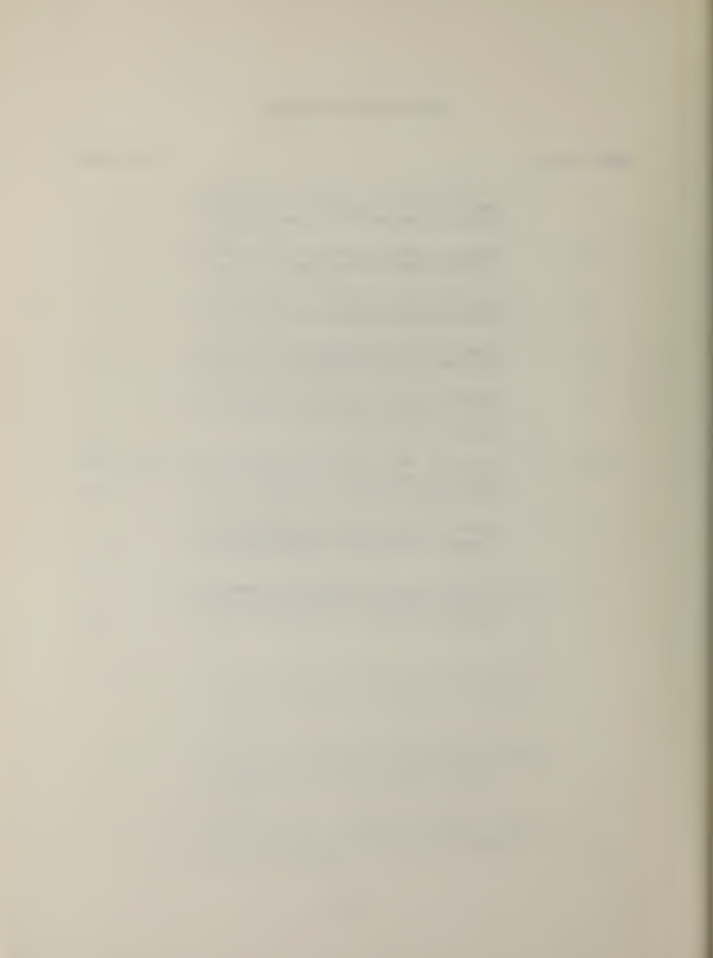
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I. INTRODUCTION

Allyl chloride (NCI No. CO4615) is one of a group of halogenated chemical intermediates selected for carcinogenesis bioassay by the National Cancer Institute. Chemicals were selected on the basis of large-scale production, extensive use, and lack of adequate chronic toxicity data.

The Chemical Abstracts Service (CAS) Ninth Collective Index (1977) name for this compound is 3-chloro-l-propene. It is also called 3-chloropropene or chloropropylene.

Allyl chloride is one of the most commercially important allyl compounds. Commercial-scale production of allyl chloride began in 1945 and increased to 295 million pounds annually in 1972 (Stanford Research Institute, 1975). It is an extremely useful chemical intermediate since it can react both as an organic halide and as an olefin (Pilorz, 1964). Most derivatives of allyl chloride do not reach an end-use market themselves, but are part of further syntheses. Important "first generation" derivatives of allyl chloride include glycerol, epichlorohydrin, and allyl alcohol (Pilorz, 1964). Other derivatives include medicinals, such as barbiturates, diuretics (Pilorz, 1964), and herbicides (Kuwahara et al., 1973).

The National Institute for Occupational Safety and Health (1976) estimates that approximately 5000 workers in the United States are

The CAS registry number is 107-05-1.

potentially exposed to allyl chloride annually. Human exposure to allyl chloride occurs principally by vapor inhalation in the working areas of industrial plants employing this compound for syntheses. Liver damage was reported in employees of the plastics industry after exposure to air concentrations varying from 1 to 113 ppm of allyl chloride for 16 months (Hausler and Lenich, 1968). Vapor exposure also produces eye and lung damage (Pilorz, 1964). Allyl chloride can be absorbed rapidly through the skin (Pilorz, 1964). Observations of industrial exposure indicate that liquid allyl chloride is a skin irritant which can cause dermatitis, damage to underlying tissues of the skin, chemical burns, and deep-seated pain (National Institute for Occupational Safety and Health, 1976).

II. MATERIALS AND METHODS

A. Chemicals

One batch of technical-grade allyl chloride (Figure 1) was purchased by Hazleton Laboratories America, Inc., Vienna, Virginia, from Aldrich Chemical Company, Inc. The purity of the compound was initially determined by gas-liquid chromatography (GLC) total-area analysis and by infrared spectroscopy. Six peaks were revealed; the fourth peak accounted for 98 percent of the total area and was presumed to be allyl chloride. One minor peak accounted for about 1.4 percent of the total area and the other four peaks totaled less than 1 percent of the area. This indication of purity in the range of 98 percent was consistent with the purity noted by the supplier. The infrared spectrum of the allyl chloride was consistent with that expected from the structure of the compound.

Second and third purity determinations were conducted approximately 19 and 26 months, respectively, after the original analysis in order to establish the stability of allyl chloride after storage. The second and third purity determinations, using GLC, showed the major peak to be approximately 98 and 99 percent, respectively, of the total area. The infrared spectra obtained in both of these analyses were consistent with the pattern shown in the first analysis. Therefore it was assumed that this batch of allyl chloride remained stable during the storage period of approximately 2 years.

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FIGURE 1 CHEMICAL STRUCTURE OF ALLYL CHLORIDE Throughout this report the term allyl chloride is used to represent this technical-grade material.

B. Dosage Preparation

Fresh solutions of allyl chloride in Duke's[®] corn oil (S. F. Sauer Company, Richmond, Virginia) were prepared weekly, sealed, and stored in dark bottles at 1°C. The concentration of allyl chloride in corn oil varied from 5.5 to 7.0 percent for the rat chronic bioassay and from 2.0 to 5.0 percent for the mouse chronic bioassay. These allyl chloride solutions were considered generally stable for 10 days under the indicated storage conditions.

C. Animals

Two animal species, rats and mice, were used in the carcinogenicity bioassay. The Osborne-Mendel rat was selected on the basis of a comparative study of the tumorigenic responsiveness to carbon tetrachloride of five different strains of rats (Reuber and Glover, 1970). The B6C3Fl mouse was selected because it has been used by the NCI for carcinogenesis bioassays and has proved satisfactory in this capacity.

Rats and mice of both sexes were obtained through contracts with the Division of Cancer Treatment, National Cancer Institute. The Osborne-Mendel rats were procured from the Battelle Memorial Institute, Columbus, Ohio, and the B6C3F1 mice from the Charles River Breeding Laboratories, Inc., Wilmington, Massachusetts. Upon receipt, animals were quarantined for at least 10 days, observed for visible

signs of disease or parasites, and assigned to the various treated and control groups.

D. Animal Maintenance

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

The rats were individually housed in suspended galvanized-steel wire-mesh cages with perforated floors. The mice were housed by sex in groups of 10 in solid-bottom polypropylene cages equipped with filter tops. Sanitized cages with fresh bedding (Sanichips[®], Pinewood Sawdust Company, Moonachie, New Jersey) were provided once each week for mice. Rats received sanitized cages with no bedding with the same frequency. Food hoppers were changed and heat-sterilized once a week for the first 10 weeks and once a month thereafter. Fresh heatsterilized glass water bottles and sipper tubes were provided three times a week. Food (Wayne Lab-Blox[®] meal, Allied Mills, Inc., Chicago, Illinois) and water were available ad libitum.

The rats dosed with allyl chloride and the untreated and vehicle control rats were housed in the same room with rats intubated with^{*} chloroform (67-66-3); carbon tetrachloride (56-23-5); 1,1,2,2-tetrachloroethane (79-34-5); and 1,2-dibromoethane (106-93-4).

^{*}CAS registry numbers are given in parentheses.

The mice dosed with allyl chloride and the untreated and vehicle control mice were housed in the same room as mice intubated with 1,1,2,2-tetrachloroethane (7934-5); chloroform (67-66-3); 3-sulfolene (77-79-2); chloropicrin (76-06-2); trichlorofluoromethane (75-69-4); dibromochloropropane (96-12-8); iodoform (75-47-8); methylchloroform (71-55-6); 1,2-dibromoethane (106-93-4); 1,2-dichloroethane (107-06-2): 1,1-dichloroethane (75-34-3); trichloroethylene (79-01-6); carbon tetrachloride (56-23-5); 1,1,2-trichloroethane (79-00-5); hexachloroethane (67-72-1) carbon disulfide (75-15-0); and tetrachloroethylene (127-18-4).

E. Gastric Intubation

Intubation was performed for five consecutive days per week on a mg/kg body weight basis, utilizing the most recently observed group mean body weight as a guide for determining the dose. Mean body weights for each group were recorded at weekly intervals for the first 10 weeks and at monthly intervals thereafter. All animals of one sex within a treated group received the same dose. Animals were gavaged with allyl chloride solutions under a hood to minimize extraneous exposure of other animals and laboratory personnel to the chemical.

F. Selection of Initial Dose Levels

In order to estimate the maximum tolerated dosages of allyl chloride for administration to treated animals in the chronic studies, subchronic toxicity tests were conducted with both rats and mice.

Animals of each species were distributed among six groups, each consisting of five males and five females. Intubation was performed 5 days per week for 6 weeks, followed by a 2-week observation period to detect any delayed toxicity. Allyl chloride, dissolved in corn oil, was introduced by gavage to five of the six rat groups at dosages of 56, 100, 178, 316, and 562 mg/kg/day and to five of the six mouse groups at dosages of 178, 316, 562, 1000, and 1780 mg/kg/day. The sixth group of each species served as a vehicle control group, receiving only corn oil.

Based on observations during the subchronic toxicity tests, the initial high dosage selected for the chronic bioassay was 110 mg/kg/ day for rats of both sexes.

At a level of 562 mg/kg/day or less only one mouse died during the 8-week study (a female treated with 562 mg/kg/day). No retardation in body weight gain was observed in either sex at 562 mg/kg/day or less. The initial high dosages selected for the mouse chronic bioassay were 400 mg/kg/day for males and 300 mg/kg/day for females.

G. Experimental Design

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

A chronic bioassay was initiated using dosages of 110 and 55 mg/ kg/day for rats. Due to lack of toxicity, dosage levels were raised

DESIGN SUMMARY FOR OSBORNE-MENDEL RATS ALLYL CHLORIDE GAVAGE EXPERIMENT

	INITIAL GROUP SIZE	ALLYL CHLORIDE DOSAGE ^a	OBSERVAT TREATED (WEEKS)	ION PERIOD UNTREATED (WEEKS)	TIME-WEIGHTED AVERAGE DOSAGE ^b
MALE					
UNTREATED CONTROL	20	0	0	110	0
VEHICLE CONTROL	20	0	78	32	0
LOW DOSE	50	70 55 0	10 68	32	57
HIGH DOSE	50	140 110 55 0	10 16 52	30	77
FEMALE					
UNTREATED CONTROL	20	0	0	110	0
VEHICLE CONTROL	20	0	78	33	0
LOW DOSE	50	55 0	78	32	55
HIGH DOSE	50	110 55 0	26 52	32	73

a Dosages, given in mg/kg body weight, were administered by gavage 5 consecutive days per week.

^bTime-weighted average dosage = $\frac{\sum (\text{dosage X weeks received})}{\sum (\text{weeks receiving chemical})}$

TABLE 2

DESIGN SUMMARY FOR B6C3F1 MICE ALLYL CHLORIDE GAVAGE EXPERIMENT

	INITIAL GROUP SIZE	ALLYL CHLORIDE DOSAGE ^a	OBSERVAT TREATED (WEEKS)	ION PERIOD UNTREATED (WEEKS)	TIME-WEIGHTED AVERAGE DOSAGE OVER A 78-WEEK PERIOD ^b
MALE					
UNTREATED					
CONTROL	20	0	0	90	0
VEHICLE					
CONTROL	20	0	78	13	0
LOW DOSE	50	200	15		172
		250	1		
		200	9		
		200 [°]	42	11	
	1	0		13	
HIGH DOSE	a 50	400	15		199
		500	1		
		400	9		
		400 [°]	3	1	
		200 ^c	21	6	
FEMALE					
UNTREATED					
CONTROL	20	0	0	90	0
VEHICLE					
CONTROL	20	0	78	13	0
LOW DOSE	50	150	25		129
		150 ^c	42	11	
		0		14	
HIGH DOSE		300	25		258
	50	300 [°]	42	11	
		0		14	

^aDosages, given in mg/kg body weight, were administered by gavage 5 consecutive days per week.

^bTime-weighted average dosage = $\frac{\sum (\text{dosage X weeks received})}{78 \text{ weeks}}$

^CThese dosages were cyclically administered with a pattern of 1 dosefree week followed by 4 weeks (5 days per week) of chemical administration at the dosage level indicated.

^dTerminated in week 56.

twice. Effective week 6, the dosages were raised to 140 and 70 mg/kg/day for both sexes and in week 12, the dosage levels for male rats were raised again, this time to 180 and 90 mg/kg/day. Because of excessive mortality after week 12, this bioassay was terminated during week 31 and the animals were discarded. Based on observations during this bioassay, a new bioassay of allyl chloride was initiated at the following levels: 140 and 70 mg/kg/day for male rats and 110 and 55 mg/kg/day for female rats.

At initiation of this study the treated and untreated control rats were approximately 6 weeks old. The vehicle control rats were approximately 7 weeks old when they were started on test; however, they were placed on test approximately 3 months before the untreated controls and the dosed groups.

Gavage was performed five consecutive days per week. The initial dosages utilized for male rats were 140 and 70 mg/kg/day. Throughout this report the male rats initially receiving the former dosage are referred to as the high dose group and those initially receiving the latter dosage are referred to as the low dose group. In week 11 the high and low dosages were reduced to 110 and 55 mg/kg/day, respectively. After 16 weeks the dosage administered to the high dose males was decreased to 55 mg/kg/day, the same dosage received by the low dose group. This dosage was maintained for the remainder of the compound administration period. Initially, the female rats received dosages of 110 and 55 mg/kg/day. Throughout this report the female

rats initially receiving the former dosage are referred to as the high dose group and those initially receiving the latter dosage are referred to as the low dose group. In week 27, because of toxic effects, the dosage level for high dose females was lowered to 55 mg/ kg/day. The vehicle control rats received corn oil in volumes equal to those administered to the high dose groups. Untreated control rats received no intubations. After the 78-week dosing period, rats were observed for 30 to 33 weeks.

At the initiation of the study the vehicle control and treated mice were approximately 5 weeks old. The untreated control mice had a median birth date approximately 2 weeks later than the other mice, and were placed on test a corresponding 2 weeks later.

Throughout this report the male mice receiving initial dosages of 400 mg/kg/day are referred to as the high dose and those receiving initial dosages of 200 mg/kg/day are referred to as the low dose. The female mice intubated with 300 and 150 mg/kg/day are referred to, respectively, as the high and low dose groups.

The dosages utilized for high dose male mice were 400 mg/kg/day for the first 15 weeks, 500 mg/kg/day the next week, 400 mg/kg/day from week 17 through week 29, and 200 mg/kg/day from week 30 until week 56 when all surviving animals in this group were sacrificed. The dosage utilized for low dose males was 200 mg/kg/day except during week 16, when they received 250 mg/kg/day. The dosages used for high and low dose female mice were 300 and 150 mg/kg/day,

respectively, throughout the 78 weeks of the experiment. In order to decrease total intake of allyl chloride, in week 26 intubation ceased for all mice for 1 week and was followed by 4 weeks of intubation at the previous dose levels. This pattern of cyclic administration continued for the remainder of the dosing period.

After the 78-week dosing period the surviving groups were observed for up to 14 weeks.

H. Clinical and Histopathologic Examinations

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

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

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

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

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

I. 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 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.

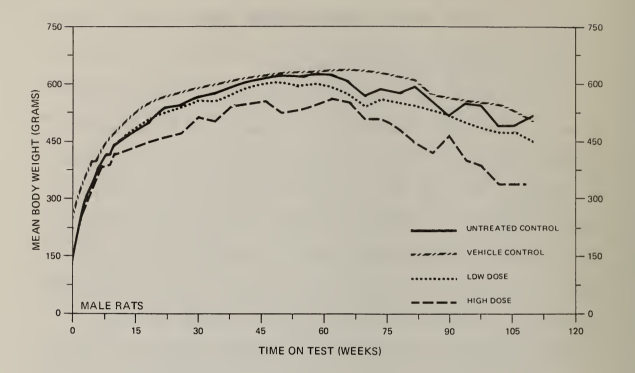
III. CHRONIC TESTING RESULTS: RATS

A. Body Weights and Clinical Observations

All groups of rats gained weight consistently during the first 46 weeks of the experiment (Figure 2). Between week 46 and week 50, the mean body weight of high dose male rats decreased from 557 grams to 528 grams. The high dose male rats continued to lose weight so that at the end of the dosing period the mean body weight of animals in this group had dropped to 487 grams. Low dose male rats experienced no appreciable mean body weight depression relative to controls. Throughout the bioassay, male and female rats treated with allyl chloride experienced consistent mean body weight depression relative to controls. Fluctuations in the growth curve may be due to mortality; as the size of the group diminishes, the mean body weight may be subject to wide variations.

Hunched appearance and abdominal urine stains were the predominant clinical signs observed during the study. Abdominal urine stains were noted in an increasing number of treated rats as the experiment progressed. By week 42 approximately 40 percent of the high dose males and high and low dose females had this condition. Abdominal urine stains were infrequently noted in the controls until the last 6 months of the study when the observation was noted in approximately 30 percent of the control rats.

A few rats showed hunched posture during the first few weeks of the study, but their appearance and behavior were otherwise normal.



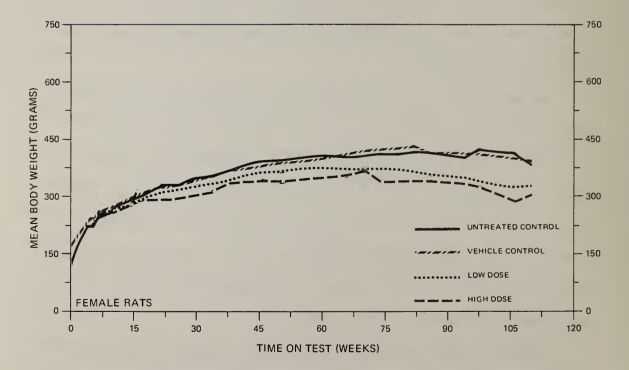


FIGURE 2 GROWTH CURVES FOR ALLYL CHLORIDE CHRONIC STUDY RATS

By week 10 more rats, particularly the females, exhibited a hunched appearance. By week 26, 30 percent of the treated males and 50 percent of the treated females had a hunched appearance. As the study progressed more animals in all groups including the controls showed hunched appearance and at the end of the study (in week 110), most or all survivors had a hunched appearance.

Respiratory signs, characterized by labored respiration, wheezing, and/or nasal discharge, were observed at a low or moderate incidence in all groups including controls during the latter part of the first year, increasing gradually as the rats aged. Other signs commonly associated with aging in the laboratory rat were observed at a comparable rate in control and treated animals during the second year of the study. These common signs included alopecia, sores on the body and/or extremities, reddish discharge or crust around the body orifices, and palpable subcutaneous masses or nodules.

B. Survival

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

For both male and female rats the Tarone test indicated a significant association (P < 0.001) between increased dosage and accelerated mortality. For both sexes the departure from linearity was significant (P < 0.001), primarily because of the extremely poor survival among the high dose groups.

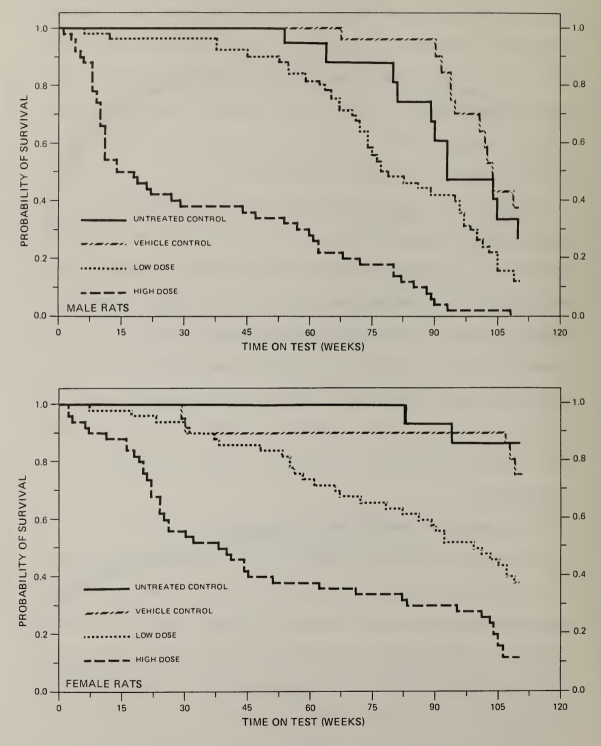


FIGURE 3 SURVIVAL COMPARISONS OF ALLYL CHLORIDE CHRONIC STUDY RATS

Fifty percent (25/50) of the low dose males survived until week 77 and 50 percent (25/50) of the low dose females survived until week 99. As a result, adequate numbers of low dose rats were at risk from late-developing tumors.

For the high dose groups, however, 50 percent (25/50) of the males had died by week 14 and 50 percent (25/50) of the females had died by week 38. Only 34 percent (17/50) of the high dose males survived one year; none survived until the end of the study. Only 12 percent (6/50) of the high dose females survived until the end of the study. These unusually early deaths were not associated with observed tumors. The small numbers of high dose males and females that survived long enough to be at risk from late-developing tumors precluded meaningful analysis of the incidence of these types of tumors for these groups.

C. Pathology

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

A variety of neoplasms were observed among both treated and control rats. Each of the types of tumors observed had been encountered historically as a spontaneous lesion in the Osborne-Mendel rat. No difference in the frequency of neoplasms or nonneoplastic lesions were noted in this test between the control and treated animals.

Results of this histopathologic examination present no evidence that allyl chloride is carcinogenic in Osborne-Mendel rats of either sex.

D. Statistical Analyses of Results

The results of the statistical analyses of tumor incidence in rats are summarized in Tables 3 and 4. Due to the high early mortality in both male and female high dose rats, many rats may have died before they were at risk from late-developing tumors. To partially compensate for this, these analyses were performed based solely upon those rats that survived at least 52 weeks or, if the tumor of interest was observed earlier than 52 weeks, at least until the first tumor of that type was observed. The analysis for every type of tumor that was observed in more than 5 percent of any of the allyl chloride-dosed groups of either sex is included.

For all analyses neither the Cochran-Armitage tests nor the Fisher exact tests indicated any statistically significant increase in the proportion of tumors found in dosed rats over that found in control rats for any tumor type for either sex. These results, therefore, provided no conclusive evidence of the carcinogencity of allyl chloride. In the high dose groups of both sexes these results must be considered statistically inconclusive due to elevated mortality. In the low dose groups, however, adequate numbers of rats were at risk for meaningful statistical analyses.

TABLE 3	
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ANALYSES OF THE INCIDENCE OF PRIMARY TUMORS AT SPECIFIC SITES IN MALE RATS TREATED WITH ALLYL CHLORIDE^A AND SURVIVING AT LEAST 52 WEEKS

	UNTREATED	VEHICLE	TOW	HJIH
TOPOGRAPHY: MORPHOLOGY	CONTROL	CONTROL	DOSE	DOSE
Subcutaneous Tissue: Fibroma ^b	0/20(0.00)	0/19(0.00)	3/45(0.07)	0/17(0.00)
P Values ^c	N.S.	N.S.	N.S.	N.S.
Relative Risk (Untreated Control) ^d		}	Infinite	
Lower Limit		-	0.283	
Upper Limit	!		Infinite	
Relative Risk (Vehicle Control) ^d			Infinite	
Lower Limit			0.272	
Upper Limit			Infinite	
Weeks to First Observed Tumor	1		77	
Subcutaneous Tissue: Fibrosarcoma ^b	1/20(0.05)	0/19(0.00)	1/45(0.02)	0/17(0.00)
P Values ^c	N.S.	N.S.	N.S.	N.S.
Relative Risk (Untreated Control) ^d			0.444	0.000
Lower Limit			0.006	0.000
Upper Limit			34.903	21.164
Relative Risk (Vehicle Control) ^d	!		Infinite	
Lower Limit			0.024	
Upper Limit			Infinite	
Weeks to First Observed Tumor	54		105	

	UNTREATED	VEH I CLE	MOT	HIGH
Pituitary: Chromophobe Adenoma	2/20(0.10)	0/16(0.00)	1/45(0.02)	0/17(0.00)
P Values ^C	N.S.	N.S.	N.S.	N.S.
Relative Risk (Untreated Control) ^d			0.222	0.000
Lower Limit Upper Limit			0.004 4.167	0.000 3.778
Relative Risk (Vehicle Control) ^d			Infinite	
Lower Limit		-	0.020	
Upper Limit			Infinite	
Weeks to First Observed Tumor	06		110	-
Thyroid: Follicular-Cell Carcinoma ^b	1/19(0.05)	2/19(0.10)	5/44(0.11)	1/17(0.06)
P Values ^C	N.S.	N.S.	N.S.	N.S.
Relative Risk (Untreated Control) ^d			2.159	1.118
Lower Limit	8	1	0.277	0.015
Upper Limit		-	101.862	82.445
Relative Risk (Vehicle Control) ^d			1.080	0.559
Lower Limit			0.205	0.010
Upper Limit			10.982	9.702
Weeks to First Observed Tumor	06	104	76	88

TABLE 3 (Continued)

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TOPOGRAPHY: MORPHOLOGY	UNTREATED CONTROL	VEHICLE CONTROL	LOW DOSE	HIGH DOSE
Thyroid: Follicular-Cell Adenoma or Follicular-Cell Carcinoma ^b	1/19(0.05)	3/19(0.16)	6/44(0.14)	1/17(0.06)
P Values ^c	N.S.	N.S.	N.S.	N.S.
Relative Risk (Untreated Control) ^d	-		2.651	1.118
Lower Limit	-	1	0.361	0.015
Upper Limit		1	118.816	82.445
Relative Risk (Vehicle Control) ^d	-		0.884	0.373
Lower Limit		-	0.218	0.008
Upper Limit	-		5.055	4.101
Weeks to First Observed Tumor	06	103	65	88
Thyroid: C-Cell Adenoma ^b	0/19(0.00)	1/19(0.05)	0/44(0.00)	0/17(0.00)
P Values ^c	N.S.	N.S.	N.S.	N.S.
Relative Risk (Untreated Control) ^d				
Lower Limit				
Upper Limit				
Relative Risk (Vehicle Control) ^d		1	0.000	0.000
Lower Limit			0.000	0.000
Upper Limit			7.800	19.052
Weeks to First Observed Tumor		110	1	1

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TABLE 3 (Concluded)

 $^{\rm a}{\rm T}{\rm reated}$ groups received time-weighted average doses of 57 and 77 mg/kg by gavage. ^bNumber of tumor-bearing animals/number of animals examined at site (proportion).

cated. The probability level for the Fisher exact test for the comparison of a treated group incidence of tumors in that 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) indi-^cThe probability level for the Cochran-Armitage test is given beneath the incidence of tumors in the corresponding control group when P < 0.05; otherwise, not significant (N.S.) is indiwith the untreated control group (*) or the vehicle control group (**) is given beneath the

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

cates a lower incidence in the treated group(s) than in the control group.

ANALYSES OF THE INCIDENCE OF PRIMARY TUMORS AT SPECIFIC SITES IN FEMALE RATS TREATED WITH ALLYL CHLORIDE ^A AND SURVIVING AT LEAST 52 WEEKS
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	UNTREATED	VEHICLE	TOW	HIGH
TOPOGRAPHY: MORPHOLOGY	CONTROL	CONTROL	DOSE	DOSE
Subcutaneous Tissue: Fibroma ^b	0/20(0.00)	0/18(0.00)	1/42(0.02)	2/19(0.011)
P Values ^c	N.S.	N.S.	N.S.	N.S.
Relative Risk (Untreated Control) ^d	1	1	Infinite	Infinite
Lower Limit			0.027	0.326
Upper Limit	-		Infinite	Infinite
Relative Risk (Vehicle Control) ^d			Infinite	Infinite
Lower Limit			0.024	0.295
Upper Limit	-		Infinite	Infinite
Weeks to First Observed Tumor			78	101
Subcutaneous Tissue: Fibrosarcoma ^b	0/20(0.00)	0/18(0.00)	2/42(0.05)	1/19(0.05)
P Values ^c	N.S.	N.S.	N.S.	N.S.
Relative Risk (Untreated Control) ^d	-	-	Infinite	Infinite
Lower Limit		1	0.149	0.058
Upper Limit	-		Infinite	Infinite
Relative Risk (Vehicle Control) ^d			Infinite	Infinite
Lower Limit			0.136	0.052
Upper Limit	1	ļ	Infinite	Infinite
Weeks to First Observed Tumor			92	95

TABLE 4

тороскарну. Мокрногосу	UNTREATED CONTROL	VEHI CLE CONTROL	DOSE	HIGH DOSE
Pituitary: Chromophobe Adenoma ^b	6/19(0.32)	6/18(0.33)	6/42(0.14)	1/19(0.05)
P Values ^c	P = 0.027(N)	P = 0.021(N)	N.S.	P = 0.045*(N) $P = 0.037**(N)$
Relative Risk (Untreated Control) ^d Lower Limit Upper Limit			0.452 0.145 1.502	0.167 0.004 1.189
Relative Risk (Vehicle Control) ^d Lower Limit Upper Limit			0.429 0.139 1.418	0.158 0.004 1.124
Weeks to First Observed Tumor	110	108	89	103
Thyroid: Follicular-Cell Carcinoma ^b P Values ^c	0/20(0.00) N.S.	0/17(0.00) N.S.	1/42(0.02) N.S.	1/19(0.05) N.S.
Relative Risk (Untreated Control) ^d Lower Limit Upper Limit			Infinite 0.026 Infinite	Infinite 0.058 Infinite
Relative Risk (Vehicle Control) ^d Lower Limit Upper Limit			Infinite 0.023 Infinite	Infinite 0.050 Infinite
Weeks to First Observed Tumor	1	1	92	110

TABLE 4 (Continued)

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	UNTREATED	VEHI CLE	TOW	HIGH
TOPOGRAPHY: MORPHOLOGY	CONTROL	CONTROL	DOSE	DOSE
Thyroid: Follicular-Cell Adenoma or Follicular-Cell Carcinoma ^b	0/20(0.00)	1/17(0.05)	1/42(0.02)	1/19(0.05)
P Values ^c	N.S.	N.S.	N.S.	N.S.
Relative Risk (Untreated Control) ^d	-		Infinite	Infinite
			0.026	0.058
Upper Limit			Infinite	Infinite
Relative Risk (Vehicle Control) ^d			0.405	0.895
Lower Limit			0.006	0.012
Upper Limit			31.046	66.483
Weeks to First Observed Tumor		110	92	110
Thyroid: C-Cell Carcinoma ^b	2/20(0.10)	0/17(0.00)	1/42(0.02)	0/19(0.00)
P Values ^c	N.S.	N.S.	N.S.	N.S.
Relative Risk (Untreated Control) ^d			0.238	0.000
Lower Limit			0.004	0.000
Upper Limit			4.359	3.408
Relative Risk (Vehicle Control) ^d	-		Infinite	
Lower Limit		-	0.023	
Upper Limit			Infinite	
Weeks to First Observed Tumor	110		103	-

	UNTREATED	VEHICLE	TOW	HIGH
TOPOGRAPHY: MORPHOLOGY	CONTROL	CONTROL	DOSE	DOSE
Thyroid: C-Cell Adenoma or C-Cell				
Čarcinoma ^b	4/20(0.20)	0/17(0.00)	1/42(0.02)	0/19(0.00)
P Values ^c	P = 0.007(N)	N.S.	P = 0.034*(N)	N.S.
Relative Risk (Untreated Control) ^d	1		0.119	0.000
Lower Limit			0.003	0.000
Upper Limit			1.117	1.077
Relative Risk (Vehicle Control) ^d			Infinite	
Lower Limit			0.023	
Upper Limit		-	Infinite	
Weeks to First Observed Tumor	61	-	103	1
Mammary Gland: Adenocarcinoma NOS ^b	2/20(0.10)	0/18(0.00)	0/42(0.00)	1/19(0.05)
P Values ^c	N.S.	N.S.	N.S.	N.S.
Relative Risk (Untreated Control) ^d			0.000	0.526
Lower Limit		-	0.000	0.009
Upper Limit		-	1.595	9.234
Relative Risk (Vehicle Control) ^d		-		Infinite
Lower Limit	1			0.052
Upper Limit	-		-	Infinite
Weeks to First Observed Tumor	110			83

TABLE 4 (Continued)

	UNTREATED	VEHICLE	LOW	HIGH
TO POGRAPHY: MORPHOLOGY	CONTROL	CONTROL	DOSE	DOSE
Mammary Gland: Fibroadenoma ^b	2/20(0.10)	7/18(0.39)	13/42(0.31)	4/19(0.21)
P Values ^c	N.S.	N.S.	N.S.	N.S.
Relative Risk (Untreated Control) ^d	-		3.095	2.105
Lower Limit		!	0.809	0.344
Upper Limit		-	26.440	20.839
Relative Risk (Vehicle Control) ^d			0.796	0.541
Lower Limit			0.373	0.142
Upper Limit		!	2.018	1.750
Weeks to First Observed Tumor	61	109	82	83
	1			

T.1.00	83	0/19(0.00)	N.S.				!	1	-		
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010.2	82	3/42(0.07)	N.S.	Infinite	0.297	Infinite	Infinite	0.269	[nfinite	110	
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	15		щ	щ			щ			2	

TABLE 4 (Continued)

TABLE 4 (Concluded)

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

^eFor sites where the first tumor of interest was observed earlier than 52 weeks, the analyses were based upon all animals that survived until or past the date that the first tumor was observed.

The possibility of negative associations between administration of the chemical and the incidence of pituitary chromophobe adenomas and thyroid C-cell neoplasms was observed in female rats. Mortality, however, was greater in the high dose group as only 6/50 (12 percent) high dose females survived until the end of the study compared to 15/50 (30 percent) vehicle control and 13/20 (65 percent) untreated control rats.

To provide additional insight into the possible carcinogenicity of this compound, 95 percent confidence intervals on the relative risk have been estimated and entered in the tables based upon the observed tumor incidence rates. In all of the intervals shown in Tables 3 and 4 the value one is included; this indicates the absence of statistically significant results. It should also be noted that all of the confidence intervals have an upper limit greater than one, indicating the theoretical possibility of tumor induction in rats by allyl chloride that could not be established under the conditions of this test.

IV. CHRONIC TESTING RESULTS: MICE

A. Body Weights and Clinical Observations

No significant depression in mean body weight was observed for allyl chloride-treated male mice (Figure 4). Among female mice a slight but consistent mean body weight depression was observed after week 10 for the high dose group and after week 20 for the low dose group.

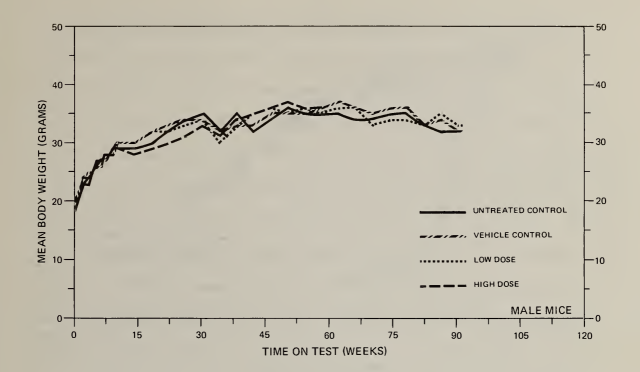
Throughout the study, signs often observed in group-housed laboratory mice were noted at a comparable rate among control and treated mice. These signs included: sores on the body (more prevalent in males because of fighting), penile, anal, or vulvar irritation, anal prolapse, reddened or squinted eyes, hunched posture, soft feces, palpable nodules, and alopecia. The incidence of these signs generally increased in all groups during the last 6 months of the study.

The only symptoms likely to be attributable to allyl chloride toxicity were observed in the 10 high dose male mice surviving beyond week 48. An apparent loss of equilibrium was observed in 8 of the 10, and abdominal distension was observed in all 10 of these animals. These signs were not noted in any of the other groups.

B. Survival

The estimated probabilities of survival for male and female mice in the control and allyl chloride-dosed groups are shown in Figure 5.

In male mice the Tarone test for a positive dose-related trend in mortality was significant (P < 0.001). There was a significant



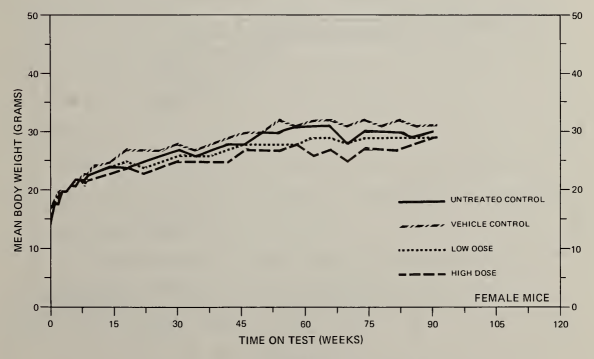
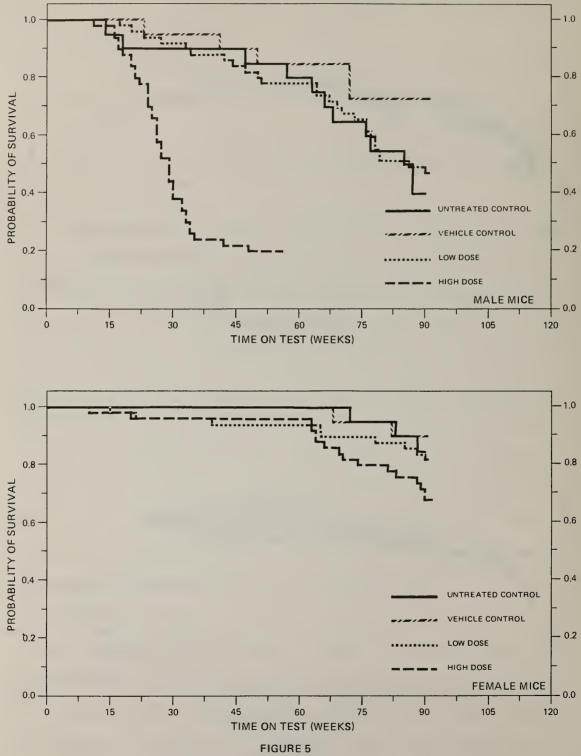


FIGURE 4 GROWTH CURVES FOR ALLYL CHLORIDE CHRONIC STUDY MICE



SURVIVAL COMPARISONS OF ALLYL CHLORIDE CHRONIC STUDY MICE

departure from linear trend (P < 0.001), primarily because of the extremely poor survival among the high dose group. Forty-eight percent (24/50) of the high dose group were dead by week 27; the 10 members of this group that survived past week 48 were sacrificed in week 56. At the same time, 10 of the 20 vehicle control mice were sacrificed. There was no indication of an association between the early deaths of high dose male mice and observed tumors. There were not adequate numbers of high dose male mice at risk from latedeveloping tumors. Survival of low dose male mice, however, was adequate for meaningful statistical analysis, with 50 percent (25/50) living at least 86 weeks.

In female mice the Tarone test also indicated a positive doserelated trend in mortality (P = 0.022). However, with 68 percent (34/50) of the high dose group and 80 percent (40/50) of the low dose group surviving to the end of the experiment, adequate numbers of female mice were at risk from late-developing tumors.

C. Pathology

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

Increased incidences of stomach lesions, both neoplastic and nonneoplastic, were observed in treated male and female mice, as shown in the following table:

	Untreated Control	Vehicle Control	Low Dose	High Dose
Males				
Number of Animals Necropsied	(18)	(20)	(46)	(50)
Squamous-Cell Carcinoma	0	0	2	0
Squamous-Cell Papilloma	0	0	0	0
Leiomyosarcoma	0	0	1	0
Acanthosis	0	0	9	19
Hyperkeratosis	0	0	9	19
Females				
Number of Animals Necropsied	(20)	(19)	(48)	(45)
Squamous-Cell Carcinoma	0	0	2	0
Squamous-Cell Papilloma	0	0	1	3
Leiomyosarcoma	0	0	0	0
Acanthosis	0	0	17	25
Hyperkeratosis	0	0	17	25

Squamous-cell carcinomas of the forestomach were observed in four treated mice. Metastases of this lesion occurred in the two low dose males, but not in the two low dose females. These tumors were not observed in the control animals and are infrequently observed in B6C3F1 mice. Microscopically, early squamous-cell carcinoma of the stomach showed acanthosis of the squamous epithelium. The surface was covered with squames of irregular needle-like structures of keratin that projected into the lumen. At the base of the epithelial layer there were papillary cords and nests of anaplastic squamous epithelial cells, supported with dense bands of fibrous connective tissue invading and replacing the lamina propria and muscularis

In well-differentiated lesions, nests of basophilic cells mucosa. with intercellular spines surrounding central areas of keratin (epithelial pearls) were seen. The more undifferentiated squamous cells had large nuclei of varying shapes, contained coarse, irregular chromatin, and had one or more nucleoli. Mitotic figures were frequently seen. In advanced lesions the cords and nests of anaplastic squamous epithelial cells invaded the muscular layers and serosa, and extended to the glandular portion of the stomach and other organs. The tumor masses in the abdominal cavity consisted of squames of keratin enclosed in nests of anaplastic squamous epithelial cells, fibrinous mats, and necrotic tissues infiltrated with inflammatory cells. Squamous-cell papillomas of the stomach were present in one low dose female and three high dose females but not in any controls. Leiomyosarcoma of the stomach was present in a single treated male mouse. Acanthosis and hyperkeratosis of the forestomach occurred with increased incidence in the treated mice of both sexes.

Hepatocellular carcinomas occurred in increased numbers in the low dose male group (8/46 [17 percent] versus 2/20 [10 percent] in the vehicle controls; and 1/49 [2 percent] in the high dose males) but was not in excess of the incidence occasionally seen in control groups. Other proliferative, inflammatory, and degenerative lesions were seen in the control and treated animals without apparent relationship to the administration of the chemical.

Oral administration of allyl chloride was associated with squamous-cell carcinomas in the stomachs of two male and two female treated mice and papillomas in the stomachs of four females, and with proliferative lesions in the forestomach of male and female mice at both dose levels.

D. Statistical Analyses of Results

The results of the statistical analyses of tumor incidence in mice are summarized in Tables 5 and 6. Due to the high early mortality in high dose male mice, many may have died before they were at risk from late-developing tumors. To partially compensate for this, the analyses for males were performed based solely upon those male mice that survived at least 52 weeks or, in the event that the tumor of interest was observed earlier than 52 weeks, upon those males which survived at least until that tumor was detected. For both males and females the analysis for every type of tumor that was observed in more than 5 percent of any of the allyl chloride-dosed groups of either sex is included.

Neither the Cochran-Armitage tests nor the Fisher exact tests indicated any statistically significant increase in the proportion of tumors found in dosed mice over that found in control mice for any tumor type for either sex. For male mice the incidence of tumors at most sites was greater in the low dose than in the high dose group; probably due to the longer survival of the low dose mice.

ANALYSES OF THE INCIDENCE OF PRIMARY TUMORS AT SPECIFIC SITES IN MALE MICE TREATED WITH ALLYL CHLORIDE ^A AND SURVIVING AT LEAST 52 WEEKS	E INCIDENCE OF WITH ALLYL CHI	OF PRIMARY TUMORS AT CHLORIDE ^a AND SURVIV	AT /IVING AT LEAS	r 52 weeks
TOPOGRAPHY: MORPHOLOGY	UNTREA TED CONTROL	VEHI CLE CONTROL	LOW DOSE	HIGH DOSE
Lung: Alveolar/Bronchiolar Adenoma ^b	0/12(0.00)	3/17(0.18)	6/35(0.17)	0/10(0.00)
P Values ^c	N.S.	N.S.	N.S.	N.S.
Relative Risk (Untreated Control) ^d			Infinite	
Lower Limit			0.600	
Upper Limit			Infinite	
Relative Risk (Vehicle Control) ^d	1		0.971	0.000
Lower Limit			0.245	0.000
Upper Limit			5.468	2.523
Weeks to First Observed Tumor	 	56	79	
Liver: Hepatocellular Carcinoma ^b	1/12(0.08)	2/17(0.12)	8/36(0.22)	1/10(0.10)
P Values ^C	N.S.	N.S.	N.S.	N.S.
Relative Risk (Untreated Control) ^d	-		2.667	1.200
Lower Limit		-	0.433	0.017
Upper Limit		-	114.336	84.143
Relative Risk (Vehicle Control) ^d			1.889	0.850
Lower Limit			0.442	0.015
Upper Limit	-		17.011	13.725
Weeks to First Observed Tumor	06	56	06	56

TABLE 5

TOPOGRAPHY: MORPHOLOGY	UNTREATED CONTROL	VEHICLE CONTROL	LOW DOSE	HIGH DOSE
Stomach: Squamous-Cell Carcinoma ^b	0/12(0.00)	0/17(0.00)	2/36(0.06)	0/10(0.00)
P Values ^c	N.S.	N.S.	N.S.	N.S.
Relative Risk (Untreated Control) ^d			Infinite	-
Lower Limit			0.107	-
Upper Limit			Infinite	
Relative Risk (Vehicle Control) ^d			Infinite	
Lower Limit		-	0.146	
Upper Limit			Infinite	!
Weeks to First Observed Tumor		-	76	
^a Treated groups received time-weighted average doses of 172 and 199 mg/kg by gavage.	average doses o	of 172 and 199	mg/kg by gavag	
"Number of tumor-bearing animals/number of animals examined at site (proportion).	r of animals exa	amined at site	(proportion).	
^c The probability level for the Cochran-Armitage test is given beneath the incidence of tumors in the corresponding control group when $P < 0.005$; otherwise, not significant (N.S.) is indi- cated. The probability level for the Fisher exact test for the comparison of a treated group with the untreated control group (*) or the vehicle control group (**) is given beneath the incidence of tumors in that 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.	-Armitage test i en P < 0.005; ot Fisher exact te or the vehicle o roup when P < 0. and Fisher exac	is given beneat therwise, not s est for the con control group (.05; otherwise, ct tests a nega than in the con	<pre>the incidence ignificant (N. parison of a t **) is given b not significa thive designati throl group.</pre>	ce of tumors S.) is indi- treated group beneath the int (N.S.) is ion (N)
d The 95% confidence interval on the relative rick of the treated aroun to the control aroun	lative rick of t	the treated and	un to the cont	rol aroun

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

TABLE 5 (Concluded)

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

TOPO GRA PHY : MORPHOLO GY	UNT REATED CONTROL	VEHICLE CONTROL	DOSE	HI GH DOSE
Lung: Alveolar/Bronchiolar Adenoma ^b	1/20(0.05)	1/19(0.05)	5/48(0.10)	4/45(0.09)
P Values ^c	N.S.	N.S.	N.S.	N.S.
Relative Risk (Untreated Control) ^d	-	-	2.083	1.778
Lower Limit			0.260	0.194
Upper Limit		!	96.358	85.520
Relative Risk (Vehicle Control) ^d			1.979	1.689
Lower Limit			0.247	0.187
Upper Limit			91.529	81.255
Weeks to First Observed Tumor	06	91	91	92
Hematopoietic System: _h				
Malignant Lymphoma ^D	3/20(0.15)	1/19(0.05)	6/48(0.13)	8/45(0.18)
P Values ^C	N.S.	N.S.	N.S.	N.S.
Relative Risk (Untreated Control) ^d			0.833	1.185
Lower Limit	1	1	0.205	0.330
Upper Limit			4.799	6.425
Relative Risk (Vehicle Control) ^d			2.375	3.378
Lower Limit			0.325	0.511
Upper Limit			106.788	145.991
Weeks to First Observed Tumor	87	91	39	81

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Treated groups received time-weighted average doses of 129 and 258 mg/kg by gavage.

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

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

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

Rare stomach tumors--squamous-cell papillomas and squamous-cell carcinomas--were observed in 2/46 (4 percent) low dose males, 3/47 (6 percent) low dose females, and 3/45 (7 percent) high dose females. In historical vehicle control data tabulated by this laboratory for the NCI Carcinogenesis Testing Program, 1/180 male and 1/180 female B6C3F1 mice had either a squamous-cell papilloma or a squamous-cell carcinoma of the stomach. Assuming a binomial distribution with a probability of 1/180 of a spontaneous tumor, the probability of observing 2 or more tumors out of 46 males was P < 0.029. For a spontaneous tumor rate of 1/180, the probabilities of 3 such tumors occurring by chance in a sample of either 47 or 45 females is very small (P < 0.003).

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

V. DISCUSSION

Because of excessive early mortality of high dose male rats, high dose female rats, and high dose male mice (50 percent of each group died by week 14, week 38, and week 27, respectively) the majority of animals in these groups did not survive long enough to be at risk from late-developing tumors. Any conclusions derived from this bioassay are, then, based on observations of the remaining groups.

Although a compound-related reduction in mortality was also observed among low dose rats, 50 percent of the low dose males survived until week 77 and 50 percent of the low dose females survived until week 99. This survival was considered adequate for meaningful statistical analysis of tumor incidence. Survival of low dose male mice (50 percent living at least 86 weeks) was also considered adequate for statistical analysis. Female mice, with 68 percent of the high dose group and 80 percent of the low dose group surviving until the end of the experiment, were the only species and sex for which animals in all groups could be considered to have lived long enough to be at risk from late-developing tumors.

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Among the groups in which an adequate number of mice had survived long enough to be at risk from late-appearing tumors, proliferative nonneoplastic stomach lesions (i.e., acanthosis and hyperkeratosis) were observed in 20 percent of the low dose males, 39 percent of the high dose males, 36 percent of the low dose females, and 56 percent

of the high dose females, but in none of the control mice. In addition, squamous-cell carcinomas of the forestomach were detected in 2/46 low dose males and 2/47 low dose females. Metastases of this lesion occurred in the two low dose males. Squamous-cell papillomas of the forestomach were observed in 1/47 low dose females and 3/45high dose females. These neoplasms were not observed in any other treated or control mice. The historical data for vehicle control B6C3F1 mice at this laboratory for the NCI Carcinogenesis Testing Program indicate that 1/180 males and 1/180 females had either a squamous-cell papilloma or a squamous-cell carcinoma of the forestom-The occurrence of these neoplasms at the incidences observed in ach. this bioassay was statistically and significantly higher than in the historical incidences. The proliferative nonneoplastic stomach lesions, squamous-cell papillomas and squamous-cell carcinomas, may all represent progressive stages in a neoplastic process. When the probable pathogenesis of this tumor is coupled with the known chemical reactivity of the compound and the statistical evidence for the rare occurrence of this tumor, the results are strongly suggestive of the carcinogenic action of allyl chloride in mice.

There were no other neoplasms in rats or mice for which statistical significance could be attributed to differences in incidence between control and treated groups.

Under the conditions of this bioassay no convincing evidence was presented for the carcinogenicity of allyl chloride in Osborne-Mendel

rats of either sex. The results are strongly suggestive that allyl chloride is carcinogenic in male and female B6C3F1 mice since the compound, when administered by gavage, is associated with neoplastic and nonneoplastic lesions of the forestomach.

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APPENDIX A

SUMMARY OF THE INCIDENCE OF NEOPLASMS IN RATS TREATED WITH ALLYL CHLORIDE

	CONTROL (UNTR) 01-031M	CONTROL (VEH) 01-021M	LOW DOSE 01-032m	HIGH DOSI 01-0330
NIMALS INITIALLY IN STUDY NIMALS MISSING	20	20 1	50	50
NIMALS NECROPSIED	20 * 20	19 19	50 50	50 50
NTEGUMENTARY SYSTEM				
*SUBCUT TISSUE FIBROMA	(20)	(19)	(50) 3 (6 %)	(50)
PIBROSARCOMA PIBROUS HISTIOCYTOMA FIBROUS HISTIOCYTOMA, MALIGNANT	1 (5%)	1 (5%) 1 (5%)	1 (23)	
LIPOMA HEMANGIOSARCOMA			1 (2%) 1 (2%)	1 (2%)
ESPILATORY SYSTEM				
NONE				
IEMATOPOIETIC SYSTEM				
*MULTIPLE ORGANS LYMPHOCYTIC LEUKEMIA	(20) 1 (5%)	(19)	(50)	(50)
*SPLEEN HEMANGIOSARCOMA	(20)	(17) 1 (6%)	(50) 1 (2 %)	(50) 1 (2%)
#MESENTERIC L. NODE HEMANGIOMA	(19)	(17)	(50)	(49) 1 (2%)
IRCULATORY SYSTEM				
NUNE				
IGESTIVE SYSTEM				
#SALIVARY GLAND CARCINOMA, NOS	(14) <u>1 (78)</u>	(17)	(41)	(18)
NUMBER OF ANIMALS WITH TISSUE EXAMI NUMBER OF ANIMALS NECROPSIED **EXCLUDES PARTIALLY AUTOLYZED ANIMALS	NED MICROSCOPIC	ALLY		

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

TABLE A1 (CONTINUED)

. <u></u>	CONTROL (UNTR) 01-031H	CONTROL (VEH) 01-0214	LOW DOSE 01-032M	HIGH DOSE 01-033M
PANCREAS OSTEOSARCOMA, METASTATIC	(20)	(16)	(50) 1 (2 %)	(50)
*STOBACH USTEUSARCOMA, METASTATIC	(20)	(19)	(50) 1 (2 %)	(50)
*SMALL INTESTIBE FIBEGSARCOMA	(20) 1 (5%)	(19)	(49)	(50)
KINAKY SYSTEM				
*KIDNEY HAMARTOMA +	(20)	(19)	(50) 1 (2 %)	(50)
NDOCRINE SYSTEM				
PITUITARY CHRONOPHOBE ADENOMA	(20) 2 (10 %)	(16)	(50) 1 (2 %)	(50)
⇒ADRENAL CURTICAL CARCINOMA PHEOCHROMOCYTOMA HEMANGIOSARCOMA	(20) 2 (10%)	(19) 1 (5%) 7 (5%)	(49) 1 (2 %)	(50)
*THYROID FOLLICULAR-CELL ADENOMA	(19)	(19) 1 (5%)	(49) 1 (2 %)	(49)
FOLLICULAR-CELL CARCINOMA C-CELL ADENOMA	1 (5%)	2 (11%) 1 (5%)	5 (10%)	1 (2\$)
*FANCHEATIC ISLETS ISLET-CELL ADENOMA	(20)	(16) 1 (6 %)	(50) 1 (2 %)	(50)
EPHODUCTIVE SISTEM				
*MAMBARY GLAND Adlaocarcinoma, NOS Pibroadenoma	(20) 1 (5%) 1 (5%)	(19) 1 (5%)	(50)	(50)
ERVOUS SYSTEM				
NONE				

NONE

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

TABLE A1 (CONTINUED)

	CONTROL (UNTR) 01-0318	CONTROL (VEH) 01-021H	LOW DOSE 01-032M	HIGH DOSE 01-033M
MUSCULOSKELETAL SYSTEM				
*SKELETAL MUSCLE PIBROUS HISTIOCYTOMA, MALIGNANT		(19) 1 (5%)	(50)	(50)
BODY CAVITIES				
BODI CAVILLES				
*MESENTERY OSTEOSARCOMA, METASTATIC	(20)	(19)	(50) 1 (2%)	(50)
ALL OTHER SYSTEMS				
NONE				
ANIMAL DISPOSITION SUMMARY				
ANIMALS INITIALLY IN STUDY	20	20	50	50
NATURAL DEATHD	11	12	42	50
MORIBUND SACRIFICE			1	
SCHEDULED SACRIFICE	5			
ACCIDENTALLY KILLED		_		
TERMINAL SACRIFICE	4	7 1	7	
ANIMAL MISSING		1		
D INCLUDES AUTOLYZED ANIMALS				

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

TABLE A1 (CONCLUDED)

		CONTROL (VEH) 01-021H		HIGH DOSI 01-0338
UNOR SUMMARY				
TOTAL ANIMALS WITH PRIMARY TUMORS* TOTAL PRIMARY TUMORS	6 11	9 12	14 . 17	46 16
TOTAL ANIMALS WITH BENIGN TUMORS TOTAL BENIGN TUMOKS	2 3	5 5	8 8	1
TOTAL ANIMALS WITH MALIGMANT TUBORS TOTAL MALIGMANT TUBORS	6 8	47	9 9	3 3
TOTAL ANIMALS WITH SECONDARY TUMORS TOTAL SECONDARY TUMORS			1 3	
TOTAL ANIMALS WITH TUMORS UNCERTAIN- Benign or Malignant Total uncertain tumors				
TOTAL ANIMALS WITH TUMORS UNCERTAIN- PRIMARY OR METASTATIC TOTAL UNCERTAIN TUMORS				

* SECONDARY TUMORS: METASTATIC TUMORS OR TUMORS INVASIVE INTO AN ADJACENT ORGAN

	IABLE A2
SUMMARY OF THE INCIDENCE OF NEOPLASMS	IN FEMALE RATS TREATED WITH ALLYL CHLORIDE

	CONTROL (UNTR) 01-031P	CONTROL (VEH) 01-021P	LOW DOSE 01-034P	BIGH DOSE 01-035P
NIMALS INITIALLY IN STUDY NIMALS NECROPSIED NIMALS EXAMINED HISTOPATHOLOGICALLY**	20 20 20	20 20 20	50 50 50	50 50 50
NTEGUMENTARY SYSTEM				
*SKIN SEBACEOUS ADENOMA	(20)	(20)	(50)	(50) 1 (2%)
*SUBCUT TISSUE SQUAMOUS CELL CARCINOMA, METASTA PIBROMA PIBROSARCOMA HEMANGIOSARCOMA	(20)	(20)	(50) 1 (2%) 1 (2%) 2 (4%) 2 (4%)	(50) 2 (4%) 1 (2%)
ESPIRATORY SYSTEM				
LUNG SQUAMOUS CELL CARCINGMA, METASTA ADENOCARCINOMA, NOS, METASTATIC FILROSARCOMA	(20)	(20)	(50) 1 (2%) 1 (2%)	(50) 1 (2%)
EMATOPOIETIC SYSTEM				
*MULTIPLE ORGANS Malig.lymphoma, Histiocytic type	(∠0)	(20)	(50) 1 (2%)	(50)
#SPLEEN HEMANGIOSARCOMA	(20)	(20)	(50) 1 (2%)	(49) 1 (2 %)
*LYMPH NODE ADENOCARCINOMA, NOS, METASTATIC	(20)	(20)	(50)	(49) 1 (2%)
*THYMUS SQUANOUS CELL CARCINOMA ADENOCARCINOMA, NOS, METASTATIC	(15)	(19)	(39) 1 (3 %)	(37) 1 (3%)
IRCULATORY SYSTEM				
#HEART MIXED TUMOR, METASTATIC	(20) 1 (5%)	(20)	(50)	(50)

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

TABLE A2 (CONTINUED)

	CONTROL (UNTR) 01-031P	CONTROL (VEH) 01-021P	LOW DOSE 01-034P	HIGH DOSE 01-035P
IGESTIVE SYSTEM				
#LIVER	(20)	(20)	(50)	(50)
NEOPLASTIC NODULE EEPATOCELLULAR CARCINOMA	1 (5%)	2 (10%)		
#PANCREAS LIPOMA	(20)	(20)	(50)	(50) 1 (2%)
*ESOPHAGUS Squamous cell carcinoma, metasta	(15)	(14)	(50) 1 (2%)	(50)
RINARY SYSTEM				
*KIDNEY	(20)	(20)	(50)	(50)
TUBULAR-CELL ADENOMA MIXED TUMOR, MALIGNANT HAMAKTONA +	1 (5%) 2 (10%)		1 (2 %) 1 (2 %)	1 (2 %) 1 (2 %)
NUOCRIAE SYSTEM				
*PITUITARY CHROMOPHOBE ADENOMA	(19) 6 (32\$)	(20) 6 (30%)	(50) 6 (12 %)	(50) 1 (2 %)
*ADRENAL CORTICAL ADENOMA	(20)	(20) 1 (5%)	(50)	(50)
FHEOCHROMOCYTOME		1 (5%)		
*THYKOID FOLLICULAR-CELL ADENOMA	(20)	(19) 1 (5 %)	(50)	(48)
FOLLICULAR-CELL CARCINOMA C-CELL ADENOMA	2 (10%)	(3%)	1 (2%)	1 (2%)
C-CELL CARCINOMA	2 (10%)		1 (2%)	
*FANCREATIC ISLETS ISLEF-CELL ADENOMA ISLET-CELL CARCINOMA	(20) 1 (5%)	(20) 1 (5%) 1 (5%)	(50)	(50)
EPRODUCTIVE SYSTEM				
*MAMMARY GLAND	(20)	(20)	(50)	(50)

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

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TABLE A2 (CONTINUED)

	CONTROL (UNTR) 01-031F	CONTROL (VEB) 01-021P	LOW DOSE 01-034P	HIGH DOSE 01-035P
PIBROSARCOMA FIBROADENOMA	2 (10%)	1 (5%) 7 (35%)	13 (26%)	4 (8%)
#OTERUS	(20)	(20)	(50)	(49)
SQUAMOUS CELL CARCINOMA Endometrial stromal polyp Hemangioma	1 (5%)	1 (5%)	3 (6%)	
#OVARY CYSTADENOCARCINOMA, NOS	(20) 1 (5%)	(20)	(50)	(49) 1 (2%)
ERVOUS SYSTEM				
NONE				
PECIAL SENSE ORGANS				
NONE				
NONE ODY CAVITIES NONE				
LL OTHER SYSTEMS				
DIAPHRAGM SQUAMOUS CELL CARCINOMA, HETASTA ADENOCARCINOMA, NOS, BETASTATIC			1	1
NIMAL DISPOSITION SUMMARY				
ANIMALS INITIALLY IN STUDY Natural deathg Moribund Sacrifice	20 2	20 5	50 31	50 42 2
SCHEDULED SACRIFICE ACCIDENTALLY KILLED TERMINAL SACRIFICE	5 13	15	19	6
ANIMAL MISSING				

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

TABLE A2 (CONCLUDED)

	01-031F	01-021P	01-034P	01-035P
DE SUMMARY				
DTAL ANIMALS WITH PRIMARY TUMORS	⊧ 1 4	12	26	11
TOTAL PRIMARY TOMORS	21	22	35	18
OTAL ANIMALS WITH BENIGN TOMORS	12	12	21	8
TOTAL BENIGN TUMORS	14	17	24	12
DTAL ANIMALS WITH MALIGNANT TUMOR	RS 6	2	9	4
TOTAL MALIGNANT TUMORS	7	3	11	6
JTAL ANIMALS WITH SECONDARY TUMOR	RS# 1		1	1
TOTAL SECONDARY TUMORS	1		4	4
OTAL ANIMALS WITH TUMORS UNCERTAD	[N-			
ENIGN OR MALIGNANT		2		
TOTAL UNCERTAIN TUBORS		2		
OTAL ANIMALS WITH TUMORS UNCERTAD	LN -			
RIMARY OR METASTATIC				
TOTAL UNCERTAIN TUMOES				

APPENDIX B

SUMMARY OF THE INCIDENCE OF NEOPLASMS IN MICE TREATED WITH ALLYL CHLORIDE



	CONTROL (VEH) 02-M031		HIGH DOSE 02-E033
20 18 * 15	20 20 20	50 46 45	50 50 49
(18)	(20)	(46) 1 (2%)	(50)
(15)	(20) 3 (15%)	(45) 1 (2%) 6 (13%)	(49)
(18)	(20) 1 (5%)	(46) 1 (2%) 1 (2%) 2 (4%)	(50) 2 (4%)
(15)	(19)	(39) 1 (3%)	(28)
			·
		(46) 1 (2%) 8 (17%) 1 (2%) 1 (2%)	(49) 1 (2%)
	20 18 (18) (15) (15) (15)	20 20 18 20 (18) (20) (15) (20) 3 (15%) (18) (20) 1 (5%)	$\begin{array}{cccccccccccccccccccccccccccccccccccc$

TABLE BI SUMMARY OF THE INCIDENCE OF NEOPLASMS IN MALE MICE TREATED WITH ALLYL CHLORIDE

TABLE B1 (CONTINUED)

	CONTROL (UNTR) 02-M041	CONTROL (VEH) 02-m031	LOW DOSE 02-H032	HIGH DOSE 02-E033
«PANCKEAS SQUAROUS CELL CARCINOMA, METASTA	(14)	(20)	(45) 1 (2%)	(43)
STUBACH SQUAHOUS CELL CARCINOMA LEIOAYOSARCOMA	(15)	(20)	(46) 2 (4%) 1 (2%)	(49)
JRIMARY SYSTEM				
NONF				
ENDUCRINE SYSTEM				
NONE				
REPRODUCTIVE SYSTEM				
*SPIDIDYMIS SQUAHOUS CELL CARCINOMA, METASTA		(20)	(46) 1 (2 %)	(50)
NERVOUS SYSTEM				
AONE				
SPECIAL SENSE ORGANS				
NONE				
NUSCULOSKELETAL SYSTEM				
NONE				
budi CAVITIES				
NONE				
ALL OTHER SYSTEMS				
NONE				

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

TABLE B1 (CONCLUDED)

	CONTROL (UNTR) 02-M041	CONTROL (VEH) 02-M031	LOW DOSE 02-M032	HIGH DOSE 02-m033
NIMAL DISPOSITION SUMMARY				
ANIMALS INITIALLY IN STUDY NATURAL DEATHO MORIBUND SACRIFICE SCHEDDLED SACRIFICE	20 12	20 4 10	50 27 1	50 40
ACCIDENTALLY KILLED TERMINAL SACRIPICE ANIMAL MISSING	8	6	1 21	10
INCLUDES AUFOLYZED ANIMALS				
UNOR SUMMARY				
IOTAL ANIMALS WITH PRIMARY TUMORS* TOTAL PRIMARY TUMORS	1	5 6	19 24	3 3
TOTAL ANIMALS WITH BENIGN TUMORS TOTAL BENIGN TUMORS		3 3	8 8	
TOTAL ANIMALS WITH MALIGNANT TUMORS TOTAL MALIGNANT TUMORS	1	3 3	12 16	3 3
TOTAL ANIMALS WITH SECONDARY TUMORS FOFAL SECONDARY TOMORS	•		2 5	
TOTAL ANIMALS WITH 1UMORS UNCERTAIN- BENIGN OR MALIGNANT TOTAL UNCERTAIN TUMORS	-			
TOTAL ANIMALS WITH TUMORS UNCERTAIN PRIMARY OR METASTATIC TOTAL UNCERTAIN TUMORS	-			
PRIMARY TUMORS: ALL TUMORS EXCEPT S SECONDARY TUMORS: METASTATIC TUMORS				

	02-P041	CONTROL (VEH) 02-F031	02-2034	02-2035
ANIMALS INITIALLY IN STUDY	20		50	50
ANIMALS NECROPSIED ANIMALS BXAMINED HISTOPATHOLOGICALLY**	20	19 19	48 48	45 44
			*0	
INTEGUNENTARY SYSTEM				
*SUBCUT TISSUE	(20)	(19)	(48)	(45)
FIBROSARCOMA LEIJHYOSARCOMA	1 (5%)		1 (2%)	
RESPIRATORY SYSTEM				
#LUNG	(20)	(19)	(48)	(45)
ADENOCARCINOMA, NOS, METASTATIC ALVEOLAR/ERONCHIOLAR ADENOMA	1 (58)	1 (54)	1 (2%)	4 (9%)
OSTEOSARCOMA, METASTATIC	(3%)	(3,4)	1 (2%)	4 (54)
NERATOPOIETIC SYSTER *IULTIPLE ORGANS	(20)	(19)	(48)	(45)
MALIG.LYMPHOMA, LYMPHOCYTIC TYPE MALIG.LYMPHOMA, HISTIOCYTIC TYPE	1 (5%)		1 (2%)	• •
MALIG.LYMPHOMA, HISTIOCYTIC TYPE MALIGNANT LYMPHOMA, MIXED TYPE	2 (10%)	1 (5%)	4 (8\$)	6 (13% 1 (2%)
*SPLEEN	(20)	(19)	(48)	(45)
LEIONYOSARCOMA, METASTATIC	1 (5%)			
#MESENTERIC L. NODE	(20) 1 (5%)	(19)	(45)	(42)
LEIONYOSARCONA, METASTATIC MALIG.LYMPHONA, HISTIOCYTIC TYPE	(5%)		1 (2%)	
*STALL INTESTINE	(20)	(19)	(47)	(44)
MALIG.LYMPHOMA, HISTIOCYTIC TYPE				1 (2%)

TABLE B2 SUMMARY OF THE INCIDENCE OF NEOPLASMS IN FEMALE MICE TREATED WITH ALLYL CHLORIDE

NONE

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

TABLE B2 (CONTINUED)

	CONÍROL (UNTR) 02-F041	CONTROL (VEH) 02-P031	LOW DOSE 02-F034	BIGH DOSE 02-P035
IGESTIVE SYSTEM				
#LIVER HEPATOCELLULAR CARCINOMA FIBROSARCOMA, METASTATIC	(20) 1 (5%)	(19)	(48) 1 (2 %) 1 (2 %)	(45) 1 (2%)
FSTOMACH SQUAMOUS CELL PAPILLOMA SQUAMOUS CELL CARCINOMA	(20)	(19)	(47) 1 (2\$) 2 (4\$)	(45) 3 (7%)
RIMARY SYSPEN				
NDOCRINE SYSTEM				
*PITUITARY CHROMOPHOBE ADENOMA	(17) 1 (6%)	(18)	(43)	(41)
*ADRENAL CORTICAL CARCINONA	(20)	(19)	(47)	(45) 1 (2%)
EPRODUCTIVE SYSTEM				
*MAMMARY GLAND ADENOCARCINOMA, NOS	(20)	(19)	(48) 1 (2 %)	(45)
#OVAKY CYSTADENOMA, NOS	(20)	(19) 1 (5%)	(48)	(43)
HEMANGIOSARCOMA			1 (2%)	
ERVOUS SYSTEM				
NONE				
SPECIAL SENSE ORGANS				
*HARDZEIAA GLAND ADENOMA, NUS	(20)	(19) 1 (5%)	(48) 1 (2 %)	(4 5)

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

B-7

TABLE B2 (CONTINUED)

		CONTROL (VEH) 02-P031		HIGH DOSI 02-P035
NUSCULOSKELETAL SYSTER				
*FEAUA USTEOSARCOMA	(20)	(19)	(48) 1 (2 %)	(45)
BODY CAVITIES				
NONE				
ALL OTHER SYSTEMS				
ANIMAL DISPOSITION SUMMARY				
ANIMALS INITIALLY IN STUDY NATURAL DEATHG MORIBUND SACRIPICE	20 3	20 2	50 9	50 15 1
SCHEDULED SACRIPICE ACCIDENTALLY KILLED	1	18	1 40	34

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

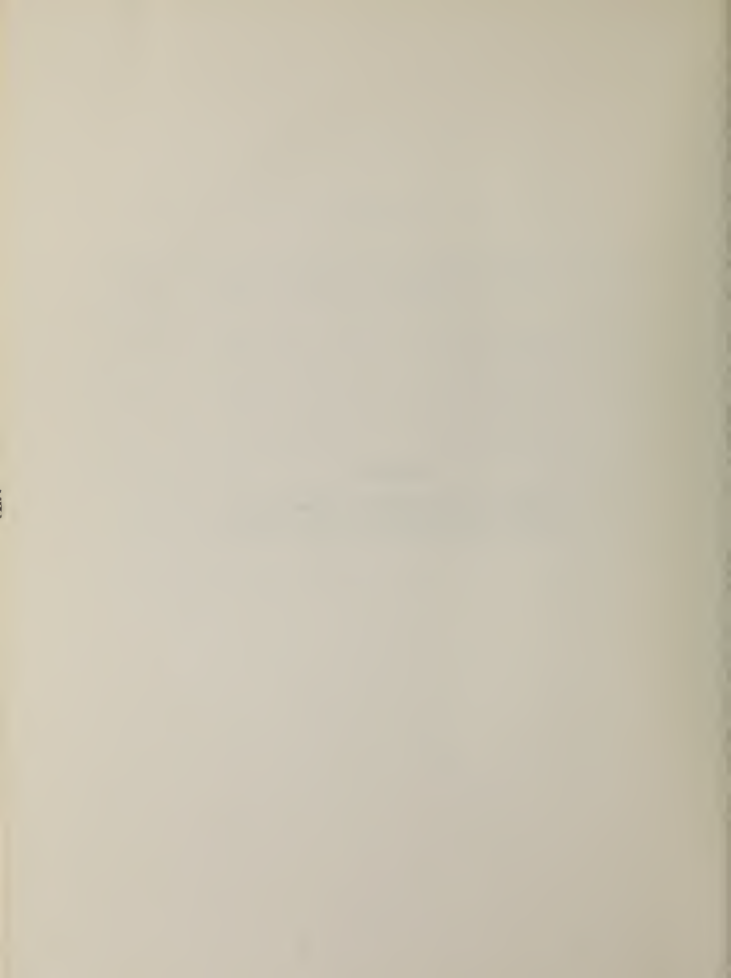
TABLE B2 (CONCLUDED)

		CONTROL (VEH) 02-P031		
JNOK SUMMARY				
TOTAL ANIMALS WITH PRIMARY TUMORS* TOTAL PRIMARY TUMORS	7 7	4 4	18 20	15 17
TOTAL ANIMALS WITH BENIGN TUMORS TOTAL BENIGN TOMORS	2 2	3 3	7 7	7 7
TOTAL ANIMALS WITH MALIGMANT TOMORS TOTAL MALIGMANT TUMORS	5 5	1	11 13	9 10
TOTAL ANIMALS WITH SECONDARY TUBORS TOTAL SECONDARY TUBORS	1 2		2 3	
TOTAL ANIMALS WITH TUNORS UNCERTAIN- BEAIGN OR MALIGNAMT TOTAL UNCERTAIN TUMORS				
TOTAL ANIMALS WITH TUMORS UNCERTAIN- PRIMARY OR METASTATIC TOTAL UNCERTAIN TUMORS	-			
PRIMARY TUMORS: ALL TUMORS ZICEPT SH SECONDARY TUMORS: METASTATIC TUMORS				



APPENDIX C

SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN RATS TREATED WITH ALLYL CHLORIDE



		CONTROL (VEH) 01-021M	LOW DOSE 01-0321	BIGH DOSE 01-03JB
ANIMALS INITIALLY IN STODY ANIMALS HISSING	20	20	50	50
NIMALS NECROPSIED NIMALS EXAMINED HISTOPATHOLOGICALLY**	20 20	19 19	50 50	50 50
NTEGUNENTARY SYSTEM				
*SKIN EPIDERMAL INCLUSION CYST	(20)	(19)	(50)	(50)
INFLAMMATION, NOS INFLAMMATION, SUPPURATIVE	1 (5%)	1 (5%)		1 (2%) 1 (2%)
ESPIRATORY SYSTEM				
*NASAL CAVITY INFLAMMATION, NOS	(20)	(19)	(50)	(50) 1 (2 %)
#TRACHEA INPLANMATION, NOS INPLANMATION, SUPPURATIVE	(15)	(19) 3 (16%) 3 (16%)	(50)	(50)
*LUNG/BRONCHIOLE INPLAEMATION, SUPPURATIVE	(20)	(19) 2 (11 %)	(50)	(50)
ALUNG PNEUMONIA, CHRONIC MURINE CALCIUM DEPOSIT	(20) 16 (80%) 1 (5%)	(19) 13 (68 %)	(50) 43 (86%)	(50) 45 (90%) 2 (4%)
ALVEOLAR MACROPHAGES	1 (5%)	1 (5%)		2 (4)
ENATOPOIETIC SYSTEM				
#BONE MARROW MERAMORPHOSIS PATTY	(3)	(19)	(50)	(50) 1 (2%)
* SPLEEN HEMATOPOIESIS	(20) 1 (5%)	(17) 1 (6%)	(50) 1 (2 %)	(50)
CERVICAL LIMPH NODE	(19) 1 (5%)	(17)	(50) 1 (2%)	(49)

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

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

**EXCLUDES PARTIALLY AUTOLYZED ANIMALS

TABLE C1 (CONTINUED)

	CONTROL (UNTR) 01-031M	CONTROL (VEH) 01-021H	LON DOSE 01-0328	HIGH DOSE 01-0338
*THYMUS Angiectasis	(13)	(2)	(40) 1 (3 %)	(26)
IRCULATORY SYSTEM				
#HEART PIEROSIS CALCIUM DEPOSIT	(20) 1 (5%)	(19)	(50) 1 (2%) 7 (14%)	(50) 2 (4%)
<pre>#MYOCANDIOM INPLAMMATION, NOS INPLAMMATION, INTERSTITIAL</pre>	(20) 2 (10%)	(19) 3 (16%) 2 (11%)	(50)	(50) 1 (2%)
FIBROSIS DEGENERATION, NOS CALCIUM DEPOSIT CALCIPICATION, NOS	1 (5%)	13 (68%)	5 (10%) 4 (8%) 7 (2%)	1 (2%) 1 (2%)
*ENDOCARDIUM HYPERPLASIA, NOS	(20) 1 (5%)	(19)	(50) 1 (2%)	(50)
*AORTA MEDIAL CALCIPICATION CALCIPICATION, BOS	(20) 2 (10 ∜)	(19) 3 (16%)	(50) 8 (16%)	(50) 4 (8 %)
*CORONARY ARTERY BEDIAL CALCIPICATION CALCIPICATION, NOS	(20)	(19) 1 (5 %)	(50) 1 (2%)	(50) 1 (2%)
*PULMONARY ARTERY CALCIFICATION, FOCAL	(20)	(19) 1 (5%)	(50)	(50)
*ASSENTERIC ARTERY PERIARTERITIS MEDIAL CALCIPICATION	(20) 1 (5%)	(19) 2 (11 %)	(50) 5 (10 %)	(50) 4 (8 %)
*PROSTATIC ARTERY MEDIAL CALCIPICATION	(20)	(19)	(50)	(50) 1 (2 %)
DIGESTIVE SYSTEM				
#LIVER THROMBUS, ORGANIZED INFLAMMATION, NOS	(20) 1 (5%)	(19)	(50) 1 (2%) 1 (2%)	(49) 1 (2 3)

* NUMBER OF ANIMALS WITH TISSUE EXAMINED HICROSCOPICALLY * NUMBER OF ANIMALS NECKOPSIED

.

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TABLE C1 (CONTINUED)

	CONTROL (UNTR) 01-031M	CONTROL (VEH) 01-021M	LOW DOSE 01-032M	HIGH DOSE 01-033H
FIEROSIS CIRKHOSIS, NOS FELIOSIS HEPATIS			1 (2%) 4 (8%) 2 (4%)	8 (16%)
NECROSIS, NOS METAHORPHOSIS PATTY Cytoplashic vacuolization Hepatocytomegaly	2 (10%)	3 (16%) 7 (37%) 7 (37%)	1 (2%) 12 (24%)	4 (8%)
ANGIECTASIS	3 (15%)	, (3, M)	8 (16%)	4 (8%)
LIVER/CENTRILOBULAR AECROSIS, NOS	(20)	(19)	(50)	(49) 1 (2 %)
*BILE DUCT INFLAMMATION, NOS FIBROSIS	(20)	(19) 8 (42%) 5 (26%)	(50)	(50)
HIPERPLASIA, NOS	4 (20%)	11 (58%)		
PANCREAS PIBROSIS, PUCAL	(20)	(16) 3 (19 %)	(50)	(50)
PERIARTERITIS	4 (20%)	- (1 <i>)</i> , (1)	8 (16%)	2 (4%)
STONACH ULCER, FOCAL	(20)	(19)	(50) 1 (2 %)	(50) 2 (4%)
CALCIUM DEPOSIT Hyperkeratosis Acanthosis	2 (10%)		6 (12\$) 1 (2\$) 1 (2\$)	4 (8%)
GASTRIC MUCOSA CALCIFICATION, NOS	(20)	(19) 2 (11%)	(50)	(50)
COLON NEMATODIASIS	(19) 1 (5%)	(19)	(49)	;50)
ALNARY SYSTEM				
KIDNEY HYDKONEPHROSIS	(20)	(19)	(50) 2 (4%)	(50)
CYST, NOS PYELOMEPHRITIS, NOS ABSCESS, NOS	1 (5%)	1 (5%)	1 (2%) 3 (6%)	1 (2%) 4 (8%)
INFLAMMATION, CHRONIC CALCIUM DEPOSIT	15 (75%) 1 (5%)	19 (100%)	42 (84%) 7 (14%)	14 (28% 5 (10%
*KIDNEY/MEDULLA CALCIFICATION, NOS	(20)	(19) 2 (11 %)	(50)	(50)

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

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TABLE C1 (CONTINUED)

	CONTROL (UNTR) 01-031M	CONTROL (VEB) 01-0218	LOW DOSE 01-0322	HIGH DOSE 01-0338
<pre>#KID#EY/PEL#IS HYPERPLASIA, EPITHELIAL</pre>	(20)	(19) 2 (11%)	(50)	(50)
UKINAKY BLADDZR ISPLAGMATIOS, BUS ISPLAGMATION, SUPPURATIVE HYPERPLASIA, EPITHELIAL	(19) 1 (5%)	(18) 1 (6 %) 1 (6 %)	(49) 3 (6 %)	(49) 4 (8 %)
NDOCKINE SYSTEM				
*PITUITARY ANGIECTASIS	(20) 1 (5 %)	(16)	(50)	(50)
*ADRENAL CORTEX DEGENERATION, NOS CYTOMEGALY ANGIECTASIS	(20) 1 (5%)	(19) 1 (5%)	(49) 1 (2%) 3 (6%)	(50)
*THYROID ULTIMJBRANCHIAL CYST POLLICULAR CYST, NOS HYPERPLASIA, C-CELL HYPERPLASIA, FOLLICULAR-CELL	(19) 2 (11%) 1 (5%) 1 (5%)	(19) 1 (5%) 4 (21%) 1 (5%)	(49) 4 (8 %)	(49) 1 (2 %)
*PARATHYKOID HYPERPLASIA, NOS	(3) 2 (67%)	(18) 4 (22%)	(23) 5 (22%)	(26) 3 (12%)
EPRODUCTIVE SYSTEM				
<pre>PROSTATE INFLAMATION, NOS INFLAMATION, SUPPORATIVE HYPERPLASIA, EPITHELIAL METAPLASIA, SQUABOUS</pre>	(20) 5 (25%)	(18) 8 (44%) 1 (6%) 4 (22%)	(43) 4 (9%)	(33) 5 (15%)
*SEMINAL VESICLE INFLAMMATION, NOS	(20) 1 (5 %)	(19)	(50) 2 (4%)	(50) 3 (6 %)
FESTIS GRANULOMA, SPERMATIC PERIARTERITIS	(20) 1 (5%)	(18) 2 (11%)	(49)	(49)
CALCIUM DEPOSIT ATROPHY, NOS	11 (55%)	8 (44%)	20 (41%)	1 (2%) 9 (18%)

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

TABLE CI (CONCLUDED)

	CONTROL (UNTR) 01-031M	CONTROL (VEH) 01-021H	LOW DOSE 01-0328	HIGH DOSE 01-0338
*2PIDIDYMIS NECROSIS, PAT ATROPHY, NOS	(20) 1 (5%) 3 (15%)	(19)	(50)	(50)
HYPERPLASIA, EPITHELIAL		1 (5%)		
ERVOUS SYSTEM				
ломк 				
SPECIAL SENSE ORGANS				
*EYE/LACRIMAL GLAND IMPLAMMATION, NOS	(20) 1 (5%)	(19)	(50)	(50)
NUSCULOSKELETAL SYSTEM				
*BONE OSTEOPOROSIS	(20)	(19) 1 (5%)	(50)	(50)
SODY CAVITIES				
*PERITONEUM INPLAMMATION, NOS	(20) 1 (5 %)	(19)	(50)	(50)
*PERICARDIUM INPLAMMATION, NOS	(20) 2 (10%)	(19)	(50) 1 (2%)	(50) 2 (4%)
*AESENTERY PERIARTERITIS NECKOSIS, PAT	(20) 4 (20%)	(19)	(50) 5 (10%) 1 (2%)	(50) 1 (2%)
ALL OTHER SYSTEMS				
NONE				
SPECIAL MORPHOLOGY SUMMARY				
NO LESION REPORTED ANIMAL MISSING/NO NECKOPSY			1	

* NUMBER OF ANIMALS NECROPSIED

C-7

	CONTROL (ONTR) 01-031P	CONTROL (VEH) 01-021P	LOW DOSE 01-034P	HIGH DOSE 01-035P
NIMALS INITIALLY IN STUDY	20	20	50	50
NIMALS NECROPSIED	20	20	50	50
NIMALS EXAMINED HISTOPATHOLOGICALLY*	* 20	20	50	50
NTEGUMENTARY SYSTEM				
*SUBCUT TISSUE	(20)	(20)	(50)	(50)
ABSC255, NOS				1 (2%)
ESPINATORY SYSTER				
*TRACHEA	(15)	(20)	(50)	(50)
INFLAMMATION, NOS		5 (25%)	1 (2%)	
INFLAMMATION, SUPPURATIVE		1 (5%)		
BETAPLASIA, SQUAMOUS		1 (5%)		
*LUNG/BRONCHIOLE	(20)	(20)	(50)	(50)
INFLAMMATION, SUPPORATIVE		4 (20%)		
*LUNG	(20)	(20)	(50)	(50)
FOREIGN BODY, NOS		1 (5%)		
CONGESTION, NOS HEMORKHAGE		1 (5%) 2 (10%)		
	18 (90%)	13 (65%)	44 (88%)	43 (86%
ALVEOLAR MACKOPHAGES	10 (2014)	5 (30%)	(,	
HYPERPLASIA, ALVEOLAR EPITHELIUE		1 (5%)		
ENATOPOLETIC SYSTEM				
*SPLEXN	(20)	(20)	(50)	(49)
HEMORRHAGE		1 (5%) 1 (5%)		
FIBROSIS, FOCAL HEMATOPOIESIS		4 (20%)	6 (12%)	1 (2%)
#CERVICAL LYMPH NODE	(20)	(20)	(50)	(49)
INFLAMMATION, NOS				1 (2%)
ANGIZCTASIS			3 (6%)	1 (2%)
HYPERPLASIA, LYMPHOID		2 (10%)		

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

• NUMBER OF ANIMALS WITH TISSUE EXAMINED BICROSCOPICALLY • NUMBER OF ANIMALS NECROPSIED **EXCLUDES PARTIALLY AUTOLYZED ANIMALS

TABLE C2 (CONTINUED)

	CONTROL (UNTR) 01-031P	CONTROL (VEH) 01-021F	LOW DOSE 01-034P	HIGH DOSE 01-035P
*THYOUS ANGIECTASIS	(15)	(19)	(39)	(37) 1 (3%)
IKCULATORY SYSTEM				
*AYOCARDIUM	(20)	(20)	(50)	(50)
INFLAMMATION, SUPPURATIVE FISHOSIS	1 (5%)	1 (5%) 1 (5%)		1 (2%)
#ENDUCARDIUM Inflammation, NOS	(20)	(20)	(50) 1 (2%)	(50)
*AORTA MEDIAL CALCIPICATION	(20) 1 (5%)	(20)	(50)	(50)
*MESENTERIC ARTERY PERIARTERITIS	(20)	(20) 3 (15%)	(50)	(50)
IGESTIVE SYSTEM #LIVER INFLAMMATION, NOS FIBROSIS	(20)	(20)	(50) 3 (6%)	(50) 1 (2%) 1 (2%)
PELIOSIS HEPATIS NECROSIS, NOS METAMORPHOSIS PATTY CYTOPLASMIC VACUOLIZATION FOCAL CELLULAR CHANGE	1 (5%)	6 (30 %)	2 (4%) 4 (8%)	1 (2%) 1 (2%) 1 (2%)
HEPATOCYTONEGALY ANGLECTASIS		6 (30%) 2 (10%)	2 (4%)	2 (4%)
#LIVER/CENTRILOBULAR DEGEMERATION, NOS	(20)	(20)	(50)	(50) 1 (2 %)
NECROSIS, NOS	1 (5%)		1 (2%)	
*BILE DUCT INFLAMMATION, NOS Flekosis Hypekplasia, Nos	(20)	(20) 8 (40%) 3 (15%) 14 (70%)	(50)	(50)
*PANCREAS PIBROSIS PERIARTERITIS	(20)	(20) 1 (5%) 1 (5%)	(50)	(50)

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

TABLE C2 (CONTINUED)

	CONTROL (UNTR) 01-031P	CONTROL (VEH) 01-021P	LOW DOSE 01-034P	HIGH DOSE 01-035F
#PANCKEATIC ACINUS Hypekplasia, focal	(20)	(20) 1 (5%)	(50)	(50)
VESOPHAGUS GRANULOMA, POREIGN BODY	(15)	(14) 1 (7 %)	(50)	(50)
<pre>#STONACH INPLAMMATION, NOS ULCER, POCAL HYPERKERATOSIS ACANTHOSIS</pre>	(20) 1 (5%) 1 (5%) 1 (5%) 1 (5%)	(20)	(50) 1 (2 %)	(50)
JRINARY SYSTEM				
<pre>#KIDNEY HIDNEPHROSIS PYELOMEPHROSIS INFLAMMATION, CHRONIC</pre>	(20) 9 (45%)	(20) 14 (70 %)	(50) 2 (4%) 16 (32%)	(50) 1 (2\$) 2 (4\$) 1 (2\$)
CALCIUM DEPOSIT *KIUNEY/PELVIS HYPERPLASIA, EPITHELIAL	1 (5%) (20)	(20) 4 (20%)	1 (2 %) (50)	1 (2%) (50)
FURIARY BLADDER INFLAMMATION, NOS	(19)	(20)	(50)	(48) 2 (4%)
ENDOCRINE SYSTEM				
<pre>#PITUITARY ANGLECTASIS</pre>	(19)	(20)	(50) 1 (2%)	(50) 1 (2%)
*ADRENAL CORTEX DEGENERATION, NOS	(20)	(20) 4 (20 %)	(50) 1 (2 %)	(50) 1 (2%)
CYTONEGALY ANGIECTASIS	3 (15%)	4 (20A)	10 (20\$)	2 (4%)
#ADRENAL MEDULLA Hypekplasia, Pocal	(20)	(20) 1 (5%)	(50)	(50)
<pre>#THYROID ULTINUBRAACHIAL CYST HYPERPLASIA, C-CELL</pre>	(20) 4 (20 %)	(19) 2 (11%) 7 (37%)	(50)	(48)
#PARATHYROID HYPERPLASIA, NOS	(1)	(16) <u>3 (19%)</u>	(8)	(6)

NUBBER OF ANIMALS WITH TISSUE EXAMINED MICROSCOPICALLY * AUNBER OF ANIMALS MECKOPSIED

TABLE C? (CONTINUED)

	CONTROL (UNTR) 01-031P	CONTROL (VEH) 01-021F	LOW DOSE 01-034F	HIGH DOSE 01-035P
REPRODUCTIVE SYSTEM				
*VAGINA INPLAMMATION, NOS	(20)	(20)	(50) . 4 (8 %)	(50) 3 (6%)
#UTERUS Hydrometra	(20) 4 (20%)	(20)	(50) 4 (8 %)	(49) 4 (8%)
*UTERUS/ENDOMETRIUM CYST, NOS INFLAMMATION, NOS INFLAMMATION, SUPPURATIVE HYPLAPLASIA, CYSTIC	(20) 1 (5%) 1 (5%)	(20) 1 (5%) 2 (10%) 1 (5%)	(50) 3 (6 %) 1 (2%)	(49) 1 (2%) 1 (2%)
*OVARY CYST, NOS	(20)	(20) 1 (5 %)	(50)	(49)
NERVOUS SYSTEM None				
SPECIAL SENSE ORGANS			*************	
*EYE/CORNEA INFLAMMATION, NOS	(20)	(20)	(50) 1 (25)	(50) .
MUSCULOSRELETAL SYSTEM				
*SKELETAL MUSCLE GRANULOMA, NOS	(20)	(20) 2 (10%)	(50)	(50)
BODY CAVITIES				
■BELIASTINUM GRANULOMA, FOREIGN BODY	(20)	(20) 1 (5%)	(50)	(50)
*PLEURA Plbkosis, Pocal	(20)	(20) 1 (5%)	(50)	(50)
*PERICARDIUM INFLAMMATION, NOS	(20)	(20)	(50) 3 (6%)	(50)

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

TABLE C2 (CONCLUDED)

		CONTROL (VEH) 01-021F		HIGH DOSE 01-035P
METAPLASIA, OSSEOUS			1 (2%)	
*EPICARDIUM INFLAMMATION, SUPPURATIVE	(20)	(20) 2 (10%)	(50)	(50)
*MASENTERY PERIARTERITIS AECROSIS, PAT	(20)	(20)	(50) 1 (2%) 1 (2%)	(50)
LL OTHER SYSTEMS				
BOAL				
PECIAL BORPHOLOGY SUBMARY				
NG LESION REPORTED				3

* NUMBER OF ANIMALS NECROPSIED

APPENDIX D

SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN MICE TREATED WITH ALLYL CHLORIDE

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SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN MALE MICE TREATED WITH ALLYL CHLORIDE

		CONTROL (VEH) 02-M031		
NIMALS INITIALLY IS STUDY	20	20	50	50
NIMALS NECROPSIED NIMALS EXAMINED HISTOPATHOLOGICALLY**	18	20 20	46 45	50 49
NTEGUMENTARY SYSTEM				
*SKIN	(18)	(20)	(46) 2 (4 %)	(50) 1 (2%)
INFLAMMATION, NGS		2 (10%)	2 (4%)	1 (2%)
*SUBCUT TISSUE	(18)	(20)	(46)	(50)
ABSCESS, NOS	1 (6%)	1 (5%)	3 (7%)	1 (2%)
LSPIRAFORY SYSTEM				
*LUNG	(15)	(20)	(45)	(49)
PREUMONIA, CHRONIC MURINE HYPERPLASIA, ALVEOLAR EPITHELIUM	((/	4 (9%) 1 (2%)	(49) 3 (6%)
EMATOPOIETIC SYSTEM	(15)	(20)	(46)	(48)
AMYLOIDOSIS HEMATOPOIESIS	7 (47%)	2 (10%) 3 (15%)	6 (13%) 1 (2%)	
#LYMPH NODE INFLAMMATION, NOS	(15)	(19)	(39) 1 (3 %)	(28)
#CERVICAL LIMPH NODE INFLAEMATION, NOS	(15)	(19) 1 (5%)	(39)	(28)
*MLSENTERIC L. NODE	(15)	(19)	(39)	(28)
INPLANMATION, NOS	1 (7%)	7 (37%)	10 (26%)	(20)
ANGIECTASIS			1 (3%)	
IRCULATORY SYSTEM				
нылыг.	(15)	(20)	(46)	(49)
PIBROSIS			1 (2%)	

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

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**EXCLUDES PARTIALLY AUTOLYZED ANIMALS

TABLE D1 (CONTINUED)

	CONTROL (UNTR) 02-M041	CONTROL (VEH) 02-M031	LOW DOSE 02-M032	HIGH DOSE 02-6033
MYOCARDIUM DEGENERATION, NOS	(15)	(20)	(46) 1 (2%)	(49)
IGESTIVE SYSTEM				
FLIVEN THROMBUS, ORGANIZED AMYLOIDOSIS BETAMORPHOSIS PATTY	(15)	(20)	(46) 1 (2%) 1 (2%) 1 (2%)	(49) 2 (4%)
*PANCREAS CYST, NOS INPLAMBATION, NUS	(14) 1 (7%)	(20)	(45) 1 (2%)	(43)
*STOMACH ULCER, NOS CALCIUM DEPOSIT HYPERKERATOSIS ACANTHOSIS	(15)	(20)	(46) 1 (2%) 9 (20%) 9 (20%)	(49) 1 (2%) 19 (39%) 19 (39%)
*COLON NENATODIASIS	(13)	(20)	(44) 1 (2%)	(46)
KINARY SYSTEM				
<pre>#KIDNEY HYDRONEPHROSIS CYST, NUS PYELONEPHRITIS, NOS INFLAMMATION, CHRONIC AMYLOIDOSIS</pre>	(15) 12 (60%) 7 (47%)	(20) 1 (5%) 2 (10%) 2 (10%) 2 (10%)	(46) 1 (2%) 5 (11%) 10 (22%) 5 (11%)	(49) 5 (10%) 3 (6%) 2 (4%)
#UKINARY BLADDER CALCULUS, NOS INFLAMMATION, NOS HYPERPLASIA, EPITHELIAL	(15)	(19) 2 (11%)	(44) 1 (2%)	(47) 1 (2%) 1 (2%) 1 (2%)
NDOCRINE SYSTEM #FHYROID FOLLICULAR CYST, NOS	(10) 1 (10%)	(20) 1 (5%)	(38)	(37)

JUMBER OF ANIMALS WITH TISSUE EXAMINED MICROSCOPICALLY * NUABER OF ANIMALS NECROFSIED

TABLE D1 (CONTINUED)

	CONTROL (UNTR) 02-8041	CONTROL (VEH) 02-H031	LOW DOSE 02-M032	HIGH DOSI 02-11033
REPRODUCTIVE SYSTEM				
*PROSTATE INFLAMMATION, NOS	(15)	(12)	(40) 1 (3%)	(43) 4 (9%)
*SEMINAL VESICLE INFLAMMATION, NOS	(18)	(20)	(46) 1 (2%)	(50) 1 (2%)
*TLSTIS ATROPHY, NOS	(15) 1 (7%)	(20)	(46)	(49)
*EPIDIDYMIS INPLAMMATION, NOS GRANULOMA, SPERMATIC	(18)	(20)	(46) 1 (2 %)	(50) 1 (2%) 1 (2%)
NERVOUS SYSTEM				
AONE				
SPECIAL SENSE ORGANS				
*HARDERIAN GLAND INFLAMMATION, NOS HYPERPLASIA, NOS	(18)	(20)	(46) 1 (2%) 1 (2%)	(50)
NUSCULOSKELETAL SYSTEM				
NONE				
BUDY CAVITIES				
NONE				
ALL OTHER SYSTEMS				
NUNE				
SPECIAL MORPHOLOGY SUMMARY				
NO LESION REPORTED	3			18
#O LESION REPORTED # NUMBER OF ANIMALS WITH TISSUE * WUMBER OF ANIMALS MECKOPSIED			55)

TABLE D1 (CONCLUDED)

AUTO/NECROPSY/HISTO 1 AUTO/NECROPSY/NO 3 1 1 AUTO/NECROPSY/NO 3 1 1		CONTROL (UNTR) 02-H041	CONTROL (VEH) 02-M031	LOW DOSE 02-M032	HIGH DOSI 02-n033
	AUTO/NECHOPSY/HISTO PERF				1
	AUTO/NECROPSY/NO HISTO	3		1	1
AUTOLISIS/NO HECKOPSI 2 4	AUTOLYSIS/NO NECROPSY	2		4	

D-6

 TABLE D2

 SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN FEMALE MICE TREATED WITH ALLYL CHLORIDE

	CONTROL (USTR) 02-P041	CONTROL (VEH) 02-F031	LOW DOSE 02-P034	HIGH DOSE 02-P035
ANIMALS INITIALLY IN STUDY ANIMALS NECROPSIED ANIMALS EXAMINED HISTOPATHOLOGICALLY ^{**}	20 20 20	20 19 19	50 48 48	50 45 44
INTEGUMENTARY SYSTEM				
*SK1 M INPLANMATION, GRANULOMATOUS	(20)	(19)	(48) 1 (2%)	(45)
*SUBCUT TISSUE ABSCESS, NOS	(20)	(19)	(48)	(45) 1 (2 %)
RESPIRATORY SYSTEM				
*LUNG PHEUMONIA, CHRONIC MURINE	(20) 1 (5%)	(19) 1 (5%)	(48) 3 (6%)	(45) 5 (11%)
ILMATOPOIETIC SYSTEM				
#SPLZEN HZMATOPOIESIS	(20)	(19)	(48) 1 (2 %)	(45) 3 (7%
BLIGPH NODE INFLAMATION, NOS	(20) 1 (5%)	(19)	(45) 1 (2%)	(42)
*SKONCHIAL LYMPH NODE IMPLAMMATION, NOS	(20)	(19)	(45)	(42) 1 (2\$)
*MESENTERIC L. NODE INPLAMMATION, NOS ANGIECTASIS	(20)	(19) 1 (5%) 1 (5%)	(45) 1 (2 %)	(42) 1 (2%)
CLACULATORY SYSTEM				
#ENDOCARDIUM HYPERPLASIA, NOS	(20)	(19)	(48)	(45) <u>1 (2%)</u>

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

**EXCLUDES PARTIALLY AUTOLYZED ANIMALS

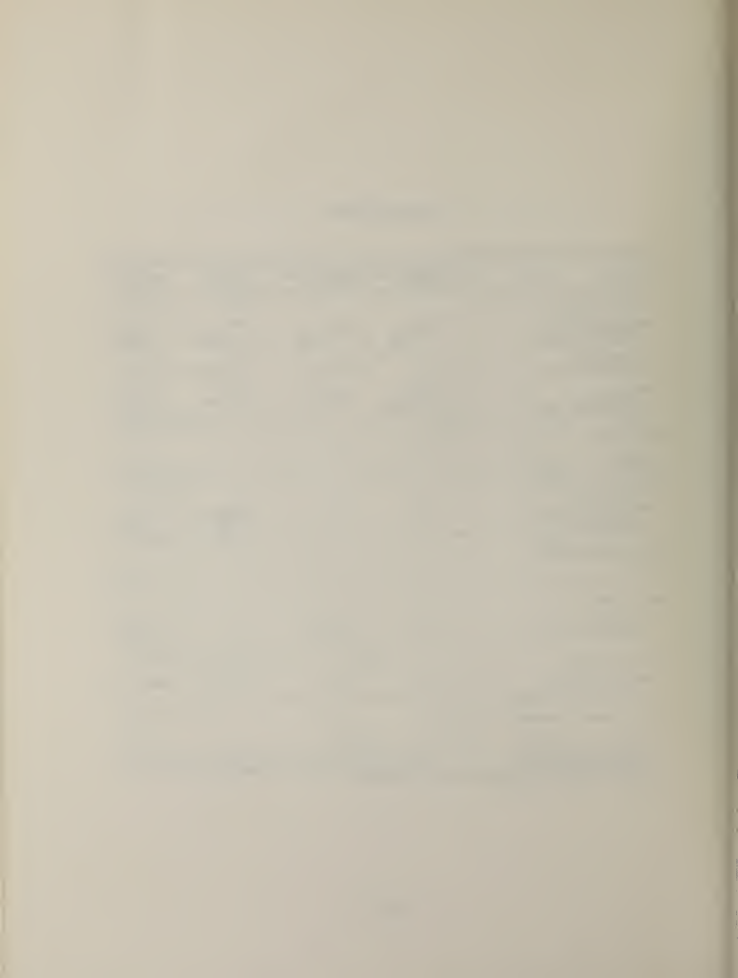
	CONTROL (UNTR) 02-P041	CONTROL (VEH) 02-P031	LOW DOSE 02-F034	HIGH DOSE 02-P035
IGESTIVE SYSTEM				
*LIVER LEUKEMOID REACTION	(20) 1 (5%)	(19)	(48)	(45)
*BILE DUCT DILATATION, NOS HYPERPLASIA, NOS	(20) 1 (5%)	(19) 1 (5 %)	(48)	(45)
#PANCREAS CYST, NOS	(20)	(19) 1 (5 %)	(46)	(45)
CYSTIC DUCTS LAPLAMMATION, NOS ATROPHY, NOS		1 (5%)	2 (4%) 2 (4%)	1 (2%) 2 (4%) 1 (2%)
*STOMACH ULCER, NOS HYPERKERATOSIS ACANTHOSIS	(20)	(19)	(47) 1 (2%) 17 (36%) 17 (36%)	(45) 1 (2%) 25 (56%) 25 (56%)
*SMALL INTESTINE HYPERPLASIA, LYMPHOID	(20)	(19)	(47) 2 (4 %)	(44) 2 (5%)
RPEYERS PATCH Hyperplasia, Lymphoid	(20)	(19)	(47) 1 (2 %)	(44)
RINAKY SYSTEM Nong				
NDOCKINE SYSTEM				
*THYRGID POLLICULAR CYST, NOS HYPERPLASIA, POLLICULAR-CELL	(16) 1 (6%) 1 (6%)	(16)	(43)	(4 1)
EPRODUCTIVE SYSTEM				
*VAGINA Laplanmation, Nos	(20)	(19) 1 (5 %)	(48)	(45)
#UTERUS HYDROMETRA	(20) 5 (25%)	(19) 2 (115)	(48) 15 (3 1%)	(43) 12 (28%)

TABLE D2 (CONTINUED)

* NUMBER OF ANIMALS WITH TISSUE EXAMINED MICROSCOPICALLY * JUMBER OF ANIMALS RECROPSIED

TABLE D2 (CONCLUDED)

	CONTROL (UNTR) 02-P041	CONTROL (VEH) 02-F031	LOW DOSE 02-F034	HIGH DOSE 02-P035
INFLAMMATION, NOS			1 (2%)	
FUTERUS/ENDOMETRIUM	(20)	(19)	(48)	(43)
INFLAMMATION, NOS	(20) 3 (15%) 6 (30%)	2 (11%)	19 (40%)	5 (12%
HYPERPLASIA, CYSTIC	6 (30%)	9 (47%)	19 (40%)	13 (30%
OVARY/OVIDUCT	(20)	(19)	(48)	(43)
INFLAMMATION, NOS		1 (5%)	1 (2%)	2 (5%)
#OVARY	(20)	(19)	(48)	(43)
CIST, NOS INFLAMMATION, NOS	12 (60%) 5 (25%)	4 (21%) 4 (21%)	11 (23%)	11 (26% 3 (7%)
ERVOUS SYSTEM				
NONE				
PECIAL SENSE ORGANS				
*HARDERIAN GLAND	(20)	(19)	(48)	(45)
INFLAMMATION, NOS	• •	• •	1 (2%)	
HYPERPLASIA, MOS			2 (4%)	4 (9%)
USCOLOSKELETAL SYSTEM				
NONE				
UDY CAVITIES				
*PERITO#EUM	(20)	(19)	(48)	(45)
INFLAMMATION, NOS		1 (5%)		1 (2%)
LL OTHER SYSTEMS				
*MULTIPLE ORGANS	(20)	(19)	(48)	(45)
LEUKENOID REACTION			1 (2%)	
PECIAL MORPHOLOGY SUMMARY				
NO LESION REPORTED	1	3	2	2
AUTO/NECROPSY/NO HISTO			2	1 5



Review of the Bioassay of Allyl Chloride* 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 Allyl Chloride for carcinogenicity.

The reviewer said that the study should be considered inadequate for drawing any conclusion about the compound's carcinogenicity. After a brief description of the experimental design, the reviewer said that the poor survival in the high dose treatment groups precluded an evaluation of the carcinogenicity of Allyl Chloride. A Program staff member said that the compound would be considered by the Chemical Selection Working Group to determine if it should be retested. The reviewer moved that the bioassay of Allyl Chloride was inadequate for drawing any conclusion on its carcinogenicity. 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

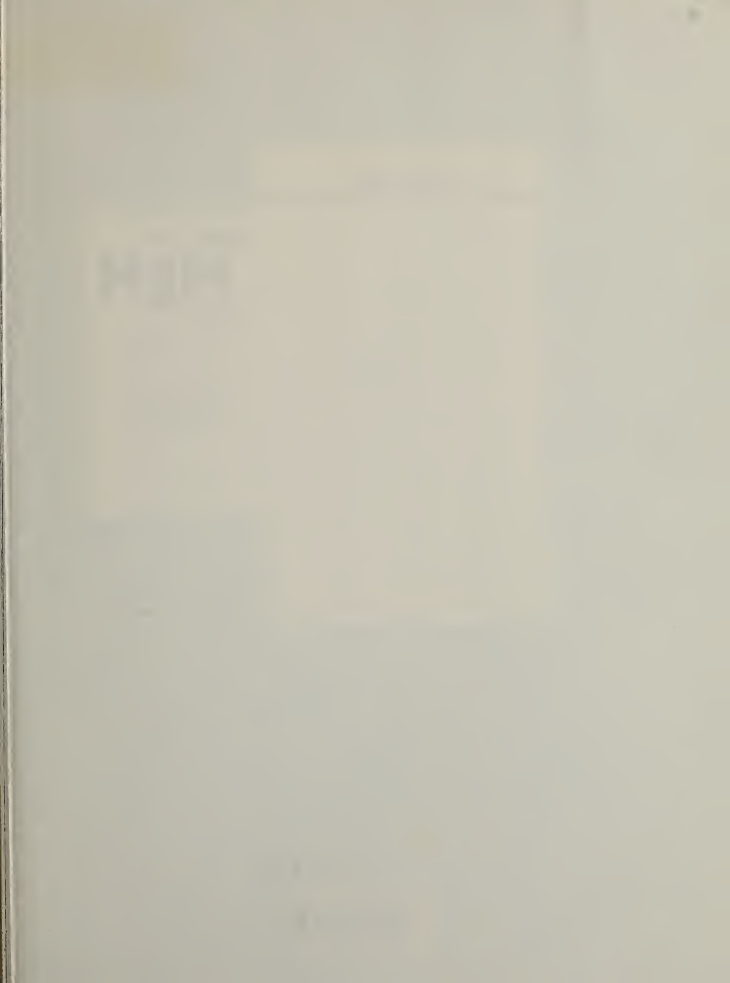
^{*} Subsequent to this review, changes may have been made in the bioassay report either as a result of the review or other reasons. Thus, certain comments and criticisms reflected in the review may no longer be appropriate.











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