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TOXICOLOGY AND CARCINOGENESIS STUDIES OF TOLUENE

(CAS NO. 108-88-3)

IN F344/N RATS AND B6C3F1 MICE

(INHALATION STUDIES)

U.S. DEPARTMENT OF HEALTH AND HUMAN SERVICES
Public Health Service
National Institutes of Health

NTP TECHNICAL REPORT

ON THE

TOXICOLOGY AND CARCINOGENESIS STUDIES OF TOLUENE

(CAS NO. 108-88-3)

IN F344/N RATS AND B6C3F₁ MICE

(INHALATION STUDIES)

James Huff, Ph.D., Study Scientist

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TOLUENE

CAS No. 108-88-3

 C_7H_8

Molecular weight 92.1

Synonyms: methylbenzene, toluol, phenylmethane, tolueen (Dutch), toluen (Czech), tolueno (Spanish), toluolo (Italian)

Trade Name: Methacide

ABSTRACT

Toluene (monomethylbenzene) is used to back-blend gasoline, as a chemical intermediate, and as a solvent; 920 million gallons was produced in the United States in 1988. Toxicology studies were conducted by administering toluene (greater than 99% pure) in corn oil by gavage to groups of F344/N rats and B6C3F₁ mice of each sex for 13 weeks or by whole-body inhalation exposure for 14 or 15 weeks. Toxicology and carcinogenesis studies were conducted by whole-body inhalation exposure of F344/N rats and B6C3F₁ mice of each sex for 15 months or 2 years. Genetic toxicology studies were conducted in Salmonella typhimurium, mouse L5178Y lymphoma cells, and Chinese hamster ovary cells.

Thirteen-Week Gavage Studies: All rats that received the top dose of 5,000 mg/kg died during the first week, and 8/10 male rats that received 2,500 mg/kg died early. The final mean body weight of male rats that received 2,500 mg/kg was 19% lower than that of vehicle controls. Relative liver, kidney, and heart (female only) weights for rats that received the higher doses were greater than those for vehicle controls. Necrosis of the brain and hemorrhage of the urinary bladder were seen at increased incidences in dosed rats.

All mice that received the top dose of 5,000 mg/kg died during the first week, and 40% of those that received 2,500 mg/kg died before the end of the 13-week gavage studies. The final mean body weight of males at 2,500 mg/kg was 16% lower than that of vehicle controls. At the higher doses, relative liver weights were increased for mice.

Fifteen-Week and Fourteen-Week Inhalation Studies: Eight of 10 male rats exposed at the top exposure concentration of 3,000 ppm died during week 2. Final mean body weights of rats exposed at concentrations of 2,500 or 3,000 ppm were 14%-25% lower than that of controls. As in the gavage studies, the relative liver, kidney, and heart weights for rats exposed at the top two concentrations were increased compared with those for controls. No compound-related effects were seen on sperm; no adverse effects on the estrous cycle were observed.

Five of 10 male mice and all female mice exposed at 3,000 ppm and 70% of female mice at 2,500 ppm died during the first 2 weeks. Final mean body weights of all exposed groups were 7%-13% lower than those of controls. Relative liver weights for mice exposed at 625 ppm or higher, relative lung weights for mice exposed at 1,250 ppm or higher, and relative kidney weights for female mice exposed at 1,250 ppm or higher were greater than those for controls. Centrilobular hypertrophy of the liver was

observed in all male mice exposed at 2,500 ppm and 70% of male mice exposed at 3,000 ppm. No effects on sperm or the estrous cycle were observed.

Fifteen-Month and Two-Year Inhalation Studies: Long-term studies were conducted by exposing groups of 60 rats of each sex to 0, 600, or 1,200 ppm toluene by inhalation, 6.5 hours per day, 5 days per week. Groups of 60 mice of each sex were exposed at 0, 120, 600, or 1,200 ppm on the same schedule. Ten animals per group (except male mice) were removed for toxicologic evaluation after being exposed for 15 months. All other animals were exposed to toluene for 103 weeks.

In the 15-month inhalation studies, the incidences and severity of nonneoplastic lesions of the nasal cavity (degeneration of olfactory and respiratory epithelium and goblet cell hyperplasia) were increased in exposed rats. Minimal hyperplasia of the bronchial epithelium was seen in 4/10 female mice at 1,200 ppm. The severity of nephropathy was slightly increased in exposed female rats. No chemical-induced neoplasms were observed.

Body Weight and Survival in the Two-Year Studies: Mean body weights of rats and mice were generally similar (yearly averages within 5%) among groups throughout the 2-year studies. No significant differences in survival were observed among rats or mice of either sex, although survival in all groups of male mice was lower than usual (male rats: control, 30/50; 600 ppm, 28/50; 1,200 ppm, 22/50; female rats: 33/50; 35/50; 30/50; male mice: control, 17/60; 120 ppm, 22/60; 600 ppm, 16/60; 1,200 ppm, 19/60; female mice: 30/50; 33/50; 24/50; 32/50). Scrotal, preputial, and penile lesions observed in the male mice were associated with many of the early deaths and with animals killed in a moribund condition.

Nonneoplastic and Neoplastic Effects in the Two-Year Studies: Nephropathy was seen in almost all rats, and the severity was somewhat increased in exposed rats. A rare renal tubular cell carcinoma in a female rat and an equally uncommon sarcoma of the kidney in another female rat were seen in the 1,200-ppm exposure group. Erosion of the olfactory epithelium and degeneration of the respiratory epithelium were increased in exposed rats. Inflammation of the nasal mucosa and metaplasia of the olfactory epithelium were increased in exposed female rats. A rare squamous cell carcinoma of the nasal mucosa was seen in one female rat at 1,200 ppm. A squamous cell papilloma of the forestomach was observed in one female rat at 1,200 ppm, and a squamous cell carcinoma was observed in a second female rat at 1,200 ppm. No chemically related neoplasms were found in male rats, and the one nasal, two kidney, and two forestomach neoplasms observed in female rats were considered not to be associated with inhalation exposure to toluene.

For mice, no biologically important increases were observed for any nonneoplastic or neoplastic lesions.

Genetic Toxicology: Toluene did not induce gene mutations in S. typhimurium strain TA98, TA100, TA1535, or TA1537 with or without exogenous metabolic activation. In the mouse lymphoma assay, toluene gave an equivocal response with and without exogenous metabolic activation. Toluene did not induce sister chromatid exchanges or chromosomal aberrations in Chinese hamster ovary cells in the presence or absence of exogenous metabolic activation.

Conclusions: Under the conditions of these 2-year inhalation studies, there was no evidence of carcinogenic activity* for male or female F344/N rats exposed to toluene at concentrations of 600 or 1,200 ppm. There was no evidence of carcinogenic activity for male or female B6C3F₁ mice exposed by inhalation to toluene at concentrations of 120, 600, or 1,200 ppm for 2 years.

SUMMARY OF THE TWO-YEAR INHALATION STUDIES OF TOLUENE

Male F344/N Rats	Female F344/N Rats	Male B6C3F ₁ Mice	Female B6C3F ₁ Mice
Exposure concentrations 0, 600, or 1,200 ppm toluene, 6.5 h/d, 5 d/wk	0, 600, or 1,200 ppm toluene, 6.5 h/d, 5 d/wk	0, 120, 600, or 1,200 ppm toluene, 6.5 h/d, 5 d/wk	0, 120, 600, or 1,200 ppm toluene, 6.5 h/d, 5 d/wk
Body weights in the 2-yea	•		
Exposed and controls similar		Exposed and controls similar	Exposed and controls similar
Survival rates in the 2 -yea 30/50; 28/50; 22/50	ar study 33/50; 35/50; 30/50	17/60; 22/60; 16/60; 19/60	30/50; 33/50; 24/50; 32/50
tory epithelium and goblet cel	eneration of respiratory epithe- nmation of nasal mucosa and		
•	Maria	MT	NT
Neoplastic effects None	None	None	None

^{*}Explanation of Levels of Evidence of Carcinogenic Activity is on page 6.
A summary of the Peer Review comments and the public discussion on this Technical Report appears on page 9.

EXPLANATION OF LEVELS OF EVIDENCE OF CARCINOGENIC ACTIVITY

The National Toxicology Program describes the results of individual experiments on a chemical agent and notes the strength of the evidence for conclusions regarding each study. Negative results, in which the study animals do not have a greater incidence of neoplasia than control animals, do not necessarily mean that a chemical is not a carcinogen, inasmuch as the experiments are conducted under a limited set of conditions. Positive results demonstrate that a chemical is carcinogenic for laboratory animals under the conditions of the study and indicate that exposure to the chemical has the potential for hazard to humans. Other organizations, such as the International Agency for Research on Cancer, assign a strength of evidence for conclusions based on an examination of all available evidence including: animal studies such as those conducted by the NTP, epidemiologic studies, and estimates of exposure. Thus, the actual determination of risk to humans from chemicals found to be carcinogenic in laboratory animals requires a wider analysis that extends beyond the purview of these studies.

Five categories of evidence of carcinogenic activity are used in the Technical Report series to summarize the strength of the evidence observed in each experiment: two categories for positive results ("Clear Evidence" and "Some Evidence"); one category for uncertain findings ("Equivocal Evidence"); one category for no observable effects ("No Evidence"); and one category for experiments that because of major flaws cannot be evaluated ("Inadequate Study"). These categories of interpretative conclusions were first adopted in June 1983 and then revised in March 1986 for use in the Technical Reports series to incorporate more specifically the concept of actual weight of evidence of carcinogenic activity. For each separate experiment (male rats, female rats, male mice, female mice), one of the following quintet is selected to describe the findings. These categories refer to the strength of the experimental evidence and not to either potency or mechanism.

- Clear Evidence of Carcinogenic Activity is demonstrated by studies that are interpreted as showing a dose-related (i) increase of malignant neoplasms, (ii) increase of a combination of malignant and benign neoplasms, or (iii) marked increase of benign neoplasms if there is an indication from this or other studies of the ability of such tumors to progress to malignancy.
- Some Evidence of Carcinogenic Activity is demonstrated by studies that are interpreted as showing a chemically related increased incidence of neoplasms (malignant, benign, or combined) in which the strength of the response is less than that required for clear evidence.
- Equivocal Evidence of Carcinogenic Activity is demonstrated by studies that are interpreted as showing a marginal increase of neoplasms that may be chemically related.
- No Evidence of Carcinogenic Activity is demonstrated by studies that are interpreted as showing no chemically related increases in malignant or benign neoplasms.
- Inadequate Study of Carcinogenic Activity is demonstrated by studies that because of major qualitative or quantitative limitations cannot be interpreted as valid for showing either the presence or absence of carcinogenic activity.

When a conclusion statement for a particular experiment is selected, consideration must be given to key factors that would extend the actual boundary of an individual category of evidence. This should allow for incorporation of scientific experience and current understanding of long-term carcinogenesis studies in laboratory animals, especially for those evaluations that may be on the borderline between two adjacent levels. These considerations should include:

- The adequacy of the experimental design and conduct;
- Occurrence of common versus uncommon neoplasia;
- Progression (or lack thereof) from benign to malignant neoplasia as well as from preneoplastic to neoplastic lesions;
- Some benign neoplasms have the capacity to regress but others (of the same morphologic type) progress. At present, it is impossible to identify the difference. Therefore, where progression is known to be a possibility, the most prudent course is to assume that benign neoplasms of those types have the potential to become malignant;
- Combining benign and malignant tumor incidences known or thought to represent stages of progression in the same organ or tissue;
- Latency in tumor induction;
- Multiplicity in site-specific neoplasia;
- Metastases:
- Supporting information from proliferative lesions (hyperplasia) in the same site of neoplasia or in other experiments (same lesion in another sex or species);
- The presence or absence of dose relationships;
- The statistical significance of the observed tumor increase;
- The concurrent control tumor incidence as well as the historical control rate and variability for a specific neoplasm;
- Survival-adjusted analyses and false positive or false negative concerns;
- Structure-activity correlations; and
- In some cases, genetic toxicology.

CONTRIBUTORS

The NTP Technical Report on the Toxicology and Carcinogenesis Studies of Toluene is based on 13week gayage studies that began in May 1981 and ended in August 1981, on 14- and 15-week inhalation studies that began in November 1981 and ended in February 1982, and on 2-year studies that began in September 1982 and ended in October 1984 at International Research and Development Corporation (Mattawan, MI).

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PEER REVIEW PANEL

The members of the Peer Review Panel who evaluated the draft Technical Report on toluene on March 13, 1989, are listed below. Panel members serve as independent scientists, not as representatives of any institution, company, or governmental agency. In this capacity, Panel members have five major responsibilities: (a) to ascertain that all relevant literature data have been adequately cited and interpreted, (b) to determine if the design and conditions of the NTP studies were appropriate, (c) to ensure that the Technical Report presents the experimental results and conclusions fully and clearly, (d) to judge the significance of the experimental results by scientific criteria, and (e) to assess the evaluation of the evidence of carcinogenicity and other observed toxic responses.

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SUMMARY OF PEER REVIEW COMMENTS ON THE TOXICOLOGY AND CARCINOGENESIS STUDIES OF TOLUENE

On March 13, 1989, the draft Technical Report on the toxicology and carcinogenesis studies of toluene received public review by the National Toxicology Program Board of Scientific Counselors' Technical Reports Review Subcommittee and associated Panel of Experts. The review meeting was held at the National Institute of Environmental Health Sciences, Research Triangle Park, NC.

Dr. J. Huff, NIEHS, began the discussion by reviewing the experimental design, results, and proposed conclusions (no evidence of carcinogenic activity for male or female rats, no evidence of carcinogenic activity for male or female mice). Dr. Huff also indicated during his remarks that male and female mice might have been able to endure somewhat higher exposure concentrations without having their health or longevity compromised.

Dr. Gallo, a principal reviewer, agreed with the conclusions. He thought that the dose selection for both rats and mice was correct based on organ weight changes and biologic activity after 14-15 weeks of exposure. He stated that discussion on the comparative metabolism of benzene and the alkylbenzenes was excellent, although some discussion on the area of toluene's (and xylene's) modifying the metabolism of benzene would enhance this section.

Dr. Popp, the second principal reviewer, agreed with the conclusions and considered the dose selection quite appropriate based on the available information. He said that the only question of carcinogenicity based on the original pathology concerned the incidence of kidney tumors; thus, the approach of making additional step-sections of the male rat kidney was appropriate and should be commended. Decreased survival in groups of male mice might best be put in perspective in relation to other inhalation studies [see page 53], and the decision not to kill 10 mice from each group at 15 months was proper.

Dr. Lijinsky, the third principal reviewer, opined that the use of the inhalation route of exposure was inappropriate because it prevented a maximum dose from being given to the animals. He noted that short-term studies had been done by the gavage route and asked why this route was not used for the 2-year studies. As a corollary, he commented that a carcinogenic effect was demonstrated by the gavage route for benzene but not by inhalation exposure. Dr. Huff responded that the 14-15 week studies were done by two routes for comparative purposes and the inhalation route was chosen largely because it was particularly relevant to human exposure and because metabolism patterns in rodents were similar by either route. He further noted that Dr. C. Maltoni in Italy has shown multi-organ carcinogenesis for benzene after inhalation exposure. Dr. Huff stated that he would communicate with Dr. Maltoni and attempt to obtain more details about the gavage studies and especially if any toxic and neoplastic lesions were considered to be related to toluene exposure.

There was some discussion about the usefulness of adding 10 animals to some groups to be killed at 15 months. Dr. Scala noted that this had been a recommendation by the NTP Ad Hoc Panel on Chemical Carcinogenesis Testing and Evaluation in its 1984 report, with the rationale being to obtain a broader look at chronic toxicity unobscured by geriatric changes. Dr. Huff said that this earlier evaluation was also helpful in preparing for evaluations after 2 years by identifying putative target organs early. Dr. Perera suggested that the NTP assess the usefulness of the interim evaluation, and Dr. Huff agreed. Dr. Ashby commented on the finding of five tubular cell adenomas in control male rats after step-sectioning and urged caution in use of data from step-sectioning until there is a fairly large data base.

SUMMARY OF PEER REVIEW COMMENTS (Continued)

Dr. Scala asked if there was more discussion on Dr. Lijinsky's contention that the study was inadequate because a high enough dose was not given. Dr. S. Eustis, NIEHS, said that the evidence of renal toxicity in both male and female rats spoke to there being sufficient exposure. Dr. Ashby thought that the inhalation route was the most appropriate. Dr. R. Griesemer, NIEHS, agreed and said that under the conditions used, the studies were adequately done and reported.

Dr. Gallo moved that the Technical Report on toluene be accepted with the conclusions as written for rats and mice of each sex, no evidence of carcinogenic activity, but with deletion of the statement that "Male and female mice might have been able to endure somewhat higher exposure concentrations without having their health or longevity compromised." Dr. Popp seconded the motion, which was accepted by nine votes and one abstention (Dr. Lijinsky).

I. INTRODUCTION

PROPERTIES, PRODUCTION, AND USE EXPOSURE STANDARDS ENVIRONMENTAL FATE HUMAN EXPOSURE METABOLISM TOXICITY IN ANIMALS

Short-Term Studies
Six-Month to One-Year Studies
Hematologic Effects
Central Nervous System Effects
Fetotoxicity and Teratogenicity
Genetic Toxicology
Carcinogenicity

TOXICITY IN HUMANS

Central Nervous System Effects
Kidney and Liver Effects
Hematologic Effects
Teratogenicity
Carcinogenicity
STUDY RATIONALE



TOLUENE

CAS No. 108-88-3

 C_7H_8

Molecular weight 92.1

Synonyms: methylbenzene, toluol, phenylmethane, tolueen (Dutch), toluen (Czech), tolueno (Spanish), toluolo (Italian)

Trade Name: Methacide

The name toluene derives from a natural resin. balsam of Tolu, named for a small town in Colombia (Kirk-Othmer, 1983). Toluene and other alkyl benzenes are single-ring, aromatic compounds containing one or more aliphatic side chains. The major products of commerce and those to which humans are most probably exposed include monomethylbenzene (toluene), ethylbenzene, 1-methylethylbenzene (cumene), and the three dimethylbenzenes (1.2-, 1.3-, and 1,4-xylene) (Andrews and Snyder, 1986). The parent molecule benzene (NTP, 1986a; Huff et al., 1988, 1989) and xylenes, mixed (NTP, 1986b; Huff et al., 1988) have been studied for carcinogenicity. Ethylbenzene is in the short-term study phase by the National Toxicology Program (NTP), and cumene is not being studied for longterm effects. This Technical Report presents the results and evaluative conclusions of the data collected from the short-term and long-term toxicology and carcinogenesis inhalation studies of toluene.

PROPERTIES, PRODUCTION, AND USE

Toluene is a colorless liquid with a benzene-like odor, a boiling point of 110.6° C, a specific gravity of 0.866, a refractive index of 1.497 at 20° C, and a flash point of 4.4° C (closed cup) (Merck, 1983). Toluene is soluble in alcohol, benzene, and ether but is insoluble in water (Condensed Chemical Dictionary, 1981).

Toluene ranked 25th (1986), 23rd (1987), and

24th (1988) in production volume for chemicals produced in the United States; approximately 7 billion pounds was produced in 1987 and 5.8 billion pounds in 1986 (Chem. Eng. News, 1988, 1989). Among organic chemicals, toluene places 13th. Seven billion pounds of toluene was produced in 1987 by 25 companies (USITC, 1988). It is produced by petroleum-refining processes, from by-products of styrene production, and from by-products of coke oven operation (Syracuse Research Corp., 1983). Purified toluene usually contains less than 0.01% benzene, but the industrial grade may contain up to 25% benzene (IPCS, 1985).

Toluene is blended with gasoline and is used as an intermediate in the synthesis of benzoic acid. benzyl and benzoyl derivatives, saccharin, medicines, dyes, perfumes, toluene diisocyanates (polyurethane resins), TNT, and toluene sulfonate detergents; and as a solvent in paints, lacquers, gums, thinners, adhesives, inks, plant resins, and pharmaceutical and cosmetic products (Condensed Chemical Dictionary, 1981; Merck, 1983; FDA, 1984; Fishbein, 1985; USEPA, 1987). Toluene is used in more than 500 cosmetic products (FDA, 1984), which include nail basecoats and undercoats (32 products), nail polish and enamel (501 products), and other manicuring preparations (22 products). Reported concentrations of toluene in these products range from 10%-25% (448 products) to more than 25%-50% (107 products).

EXPOSURE STANDARDS

The current threshold limit value/time-weighted average (TWA) for an 8-hour workday, 40-hour workweek, in the United States is 100 ppm (375 mg/m³); the short-term exposure limit is 150 ppm (560 mg/m³) (ACGIH, 1987). The Occupational Safety and Health Administration lists 200 ppm (8-hour TWA), whereas the National Institute for Occupational Safety and Health has promulgated a standard of 100 ppm (10-hour TWA) and of 200 ppm with a 10-minute ceiling. The Immediately Dangerous to Life or Health level is 2,000 ppm (NIOSH, 1985). In 1987, the value for an 8-hour workday was reduced to 50 ppm in several countries (IRPTC, 1987).

ENVIRONMENTAL FATE

Toluene is quite stable in air but can be oxidized in the presence of catalysts to yield benzoic acid. In the presence of heat (or a catalyst) and hydrogen, toluene undergoes dealkylation to produce benzene. Under conditions of water chlorination, toluene may be chlorinated and subsequently hydrolyzed to benzaldehyde. Toluene may undergo photo-oxidation (Shepson et al., 1984) and other photochemical reactions (NRC, 1981; Syracuse Research Corp., 1983). Because of the limited number of studies available, the extent of toluene degradation in soil cannot be determined, although studies with pure cultures indicate that a variety of bacteria and fungi can metabolize toluene and that some organisms can use toluene as a sole source of carbon. A toluene half-life of 20-60 minutes was observed in soil containing toluene-degrading bacteria (USEPA. 1983). Toluene is readily biodegraded in water, both in surface water and during wastewater treatment; however, disappearance of toluene from water is mainly through evaporation.

Evaporation of gasoline and automobile exhaust are the largest combined source of toluene (677 million kg per year) in the environment, and industries that use toluene as a solvent are the second largest source (375 million kg per year); these two sources account for 75% of the toluene emitted to the atmosphere (USEPA, 1983). Nonatmospheric release of toluene to the environment (e.g., to water or soil) is comparatively small and is approximately 0.15% of the total amount emitted to the atmosphere.

Toluene is the most prevalent aromatic hydrocarbon in the atmosphere, with average measured levels ranging from 0.14 to 59 µg/liter (USEPA, 1983). Toluene has been detected in surface water and in treated wastewater effluents at levels generally below 10 µg/liter. A concentration of toluene as high as 19 µg/liter has been detected in a drinking water supply. In a study of toluene levels in the tissue of edible aquatic organisms, 95% of the samples contained less than 1 ppm of toluene.

HUMAN EXPOSURE

The estimated intake of toluene by the general public is between a trace and 94 mg per week by inhalation (depending on whether an individual resides in an urban or rural area or near an industry that uses toluene) and 0-0.75 mg per week from food and water (USEPA, 1983). Occupational exposure (up to 18,000 mg per week) or cigarette smoking (0.1 mg per cigarette) adds considerably to an individual's exposure to toluene. Exposure also occurs through deliberate inhalation of solvents found in various preparations, such as glue (sniffing). An estimated 124 million people in the United States are exposed to atmospheric toluene at a concentration greater than 0.27 µg/liter.

METABOLISM

The metabolism of toluene has been extensively reviewed (IPCS, 1985; CTFA, 1986; Wallen, 1986; USEPA, 1987). Toluene is readily absorbed from the respiratory tract of mammals (Nomiyama and Nomiyama, 1974; Sato et al., 1974a,b; Astrand, 1975; Egle and Gochberg, 1976; Sherwood, 1976; Carlsson and Lindqvist, 1977; Sato and Nakajima, 1979; Benignus, 1981a; WHO, 1981; Carlsson, 1982; Rees et al., 1985). In humans, the uptake of toluene is 40%-60% of the amount inhaled (Nomiyama and Nomiyama, 1974). The uptake of toluene through the skin of humans (with respiratory protection) exposed at 600 ppm in air for 3.5 hours was 1% of the theoretical respiratory uptake; the toluene concentration in peripheral venous blood after 1, 2, and 3 hours of exposure was approximately 100 µg/liter (Riihimaki and Pfaffli, 1978). Toluene is almost completely absorbed from the gastrointestinal tract (Smith et al., 1954; El Masry et al., 1956; Cohr and Stokholm, 1979; Syracuse Research Corp., 1983; Slooff and Blokzijl, 1988).

In mice exposed to toluene at 4,000 ppm for 3 hours, the concentration of toluene was 625 mg/kg in liver, 420 mg/kg in brain, and 200 mg/kg in blood (Peterson and Bruckner, 1978; Bruckner and Peterson, 1981a). Immediately after adult male rats were exposed to [methyl-14C]toluene at 550 ppm by inhalation for 1 hour, the amount of radioactive label in adipose tissue was more than two times the amount in any other organ (Carlsson and Lindqvist, 1977). Six hours after exposure, radioactivity was still measurable in liver, kidney, and adipose tissue. In studies with mice exposed to [methyl-14C]toluene by inhalation, high levels of radioactive label were found in adipose tissue, bone marrow, spinal nerves, spinal cord, and white matter of brain (Bergman, 1979, 1983). Radioactivity was no longer detectable in nervous system tissue 1 hour after exposure and was cleared from body fat after 4 hours. Traces of nonvolatile radioactivity were present after 4 hours, but no radioactivity was detectable by 24 hours. Distribution of toluene to tissues after gavage administration is similar to, but slower than, that after inhalation exposure (Pyykko et al., 1977). Studies in adult male Wistar rats exposed to toluene at 300 ppm (6 hours per day for 1-15 weeks) indicated a decrease of toluene in perirenal fat over time, suggesting enhanced activity of drug-metabolizing enzymes in liver and metabolic and functional adaptation after long-term low-level exposure (Elovaara et al., 1979).

After an intraperitoneal injection of [methyl-14C]toluene at 500 µmol to rats, the concentration of radioactivity was highest in the cerebrum (Savolainen, 1978). After an intraperitoneal injection at 0.2 mg/kg to mice, most of the radioactivity in the adipose tissue and brain was volatile (probably unmetabolized toluene), whereas most of the radioactivity in the liver and kidney was nonvolatile (probably a metabolite) (Koga, 1978).

When pregnant CFY rats were exposed to toluene at 370 or 720 ppm for 24 hours on days 10-13 of gestation, the toluene concentration 2 hours after exposure was 6.4 or 1 3.7 mg/liter in

maternal blood and 4.9 or 10.4 mg/liter in fetal blood (Ungvary, 1984).

Toluene is rapidly metabolized primarily in the liver (SRI, 1980; Slooff and Blokzijl, 1988). In rats, rabbits, and humans, 25%-40% of an oral or inhaled dose is excreted unchanged in exhaled air, and 60%-75% of the dose is converted to benzoic acid and excreted in urine, primarily as hippuric acid; smaller amounts are excreted as the sulfate or glucuronide conjugate of benzoic acid (Figure 1) (von Oettingen et al., 1942a; Srbova and Teisinger, 1952; Smith et al., 1954; El Masry et al., 1956; Daly et al., 1968; Bakke and Scheline, 1970; Angerer, 1976; Pfaffli et al., 1979; Toftgard and Gustafsson, 1980; Van Doorn et al., 1980; Woiwode and Drysch, 1981; Baelum et al., 1987). Less than 1% of the absorbed toluene is hydroxylated to o-, m-, or p-cresol, which is then excreted as the sulfate or glucuronide conjugates (DeBruin, 1976; Woiwode et al., 1979; Baelum et al., 1987). The first step in the conversion of toluene to benzoic acid is conversion to benzyl alcohol by an NADH-dependent hydratase or a monooxygenase (Bakke and Scheline, 1970; SRI, 1980). The benzyl alcohol is rapidly converted to benzaldehyde by an NADdependent alcohol dehydrogenase, and the benzaldehyde is converted to benzoic acid by an NAD-dependent aldehyde dehydrogenase (SRI, 1980; IPCS, 1985). o-Cresol or hippuric acid in the urine is an indication of toluene exposure in workers (Angerer, 1979; Ogata et al., 1970; Woiwode and Drysch, 1981; Apostoli et al., 1982; Andersson et al., 1983; Dossing et al., 1983; Hasegawa et al., 1983; Kono et al., 1985; De Rosa et al., 1985, 1986, 1987; Ogata and Taguchi. 1986); however, Baelum et al. (1985a) indicated that urinary o-cresol concentrations give a more specific estimate of toluene exposure than hippuric acid concentrations. o-Cresol excretion is delayed compared with hippuric acid excretion and is more consistent when exposure is accompanied by physical activity (Baelum et al., 1987). In rats exposed to toluene at 5-500 ppm by inhalation, the urinary ratio of hippuric acid to ocresol was constant, but at 2,500 or 3,500 ppm, the amount of o-cresol increased sharply (Inoue et al., 1984). Of four strains of rats exposed to toluene, F344 rats excreted the most o-cresol and Sprague Dawley rats excreted the least; Wistar

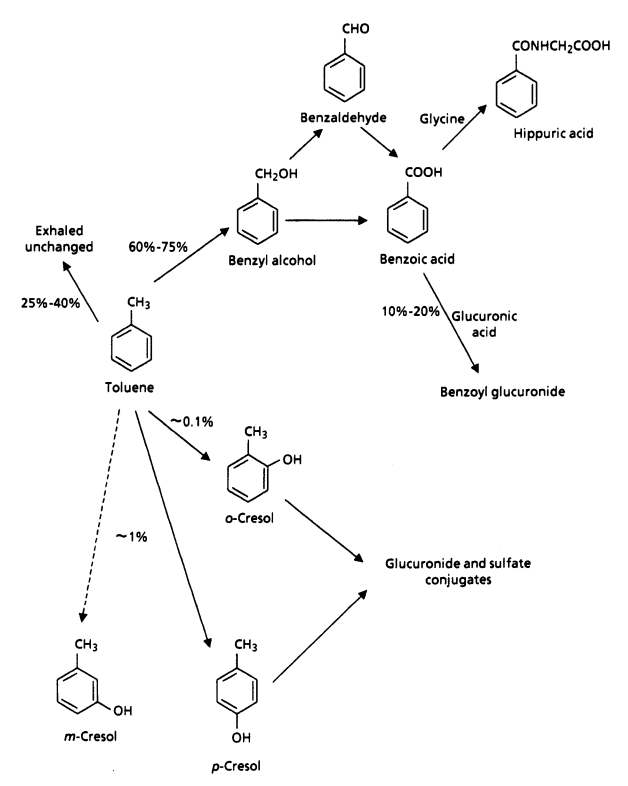


FIGURE 1. METABOLISM OF TOLUENE IN HUMANS AND ANIMALS (taken from IPCS, 1985, and modified)

rats excreted the most *p*-cresol. A biologic exposure index based on the direct analysis of toluene in urine has been suggested by Pezzagno et al. (1985), who found a correlation between the concentration of toluene in air and in urine.

TOXICITY IN ANIMALS

Extensive reviews of the toxicity of toluene are presented in the International Programme on Chemical Safety (IPCS, 1985), Cosmetic, Toiletry and Fragrance Association (CTFA, 1986), Bell et al. (1988), Agency for Toxic Substances and Disease Registry (ATSDR, 1989), and International Agency for Research on Cancer (IARC, 1989).

Short-Term Studies

Oral LD₅₀ values for toluene range from 2.6 to 7.5 g/kg for juvenile to adult rats (Cameron et al., 1938; Kimura et al., 1971; Withey and Hall, 1975; Ungvary et al., 1979). The dermal LD_{50} is 14.1 ml/kg for rabbits (Smyth et al., 1969). The LC₅₀ value for a 6- to 7-hour exposure is 12,200 ppm for rats (Cameron et al., 1938) and 5,300-7,000 ppm for mice (Svirbely et al., 1943; Bonnet et al., 1979). The intraperitoneal LD₅₀ is 1.64 g/kg for female rats (Ikeda and Ohtsuji, 1971). In freshwater organisms, the LC₅₀ for mosquito larvae is 21.5 mg/liter, whereas for fathead minnows, the LC₅₀ is 26 mg/liter for juveniles and 29 mg/liter for day-old fry. The LC₅₀ values for marine organisms include 3.7 mg/liter for bay shrimp and 28 mg/liter for Dungeness crabs (Caldwell et al., 1976; Benville and Korn, 1977; Berry and Brammer, 1977; USEPA, 1980; Devlin et al., 1982).

Adverse effects were observed in animals administered toluene by various routes of exposure. Undiluted toluene was found to be an ocular and skin irritant (Wolf et al., 1956; Guillot et al., 1982a,b). Slight induration at the injection site, decreased body weight, hyperplasia of bone marrow and of malpighian corpuscles in the spleen, marked pigmentation of spleen, focal necrosis of the liver, and slight cloudy swelling in the kidney were seen in rats given subcutaneous injections of toluene at 1 ml/kg for 21 days (Batchelor, 1927). In guinea pigs, toluene given by subcutaneous injection at 0.25 ml per day for 30-

70 days caused necrosis at the injection site, polypnea and convulsions toward the end of the studies, and hemorrhagic, hyperemic, and degenerative changes in the lung, kidney, adrenal gland, liver, and spleen (Sessa, 1948). An increased number of casts was seen in the collecting tubules of the kidney in rats exposed to toluene (99.9% pure) in air at 200-5,000 ppm, 7 hours per day, 5 days per week for 5 or 15 weeks, and in dogs exposed at 200-600 ppm, 8 hours per day for 20 days, followed by exposure for 7 hours per day, 5 days per week for 1 week, and then at 850 ppm for 1 hour (von Oettingen et al., 1942a).

Toluene at near lethal exposure concentrations (up to 66,000 ppm for up to 30 minutes) had no untoward electrocardiographic effects and even appeared to decrease epinephrine-induced ectopic beats in male Wistar rats (220-242 g) (Vidrio et al., 1986).

Six-Month to One-Year Studies

DONRYU male rats were exposed to 0, 100, 200, or 2,000 ppm toluene vapor 8 hours per day, 6 days per week, for 10, 18, or 43 weeks (Matsumoto et al., 1971). The most significant histopathologic change was in the kidney; numerous eosinophilic droplets of various sizes (termed by the authors as "hyaline droplets") were observed in the renal tubular epithelium in each exposed group. Only a few droplets were seen in controls. The longer the duration, the larger and more frequent were the hyaline droplets. In the 2,000ppm toluene group exposed for 43 weeks, the droplets were large and the amounts were increased. Matsumoto and coworkers indicated that the hyaline droplets originated from degenerated microsomes, and although the droplets were seen after proteinuria was evident, the relationship between hyaline droplets and proteinuria was not examined. [Note: In January 1989, at NTP's request, Dr. Matsumoto kindly sent several Kodachrome slides of the kidney sections from his 1971 study; these were examined, and the typical hyaline droplet nephropathy was not observed.

No histopathologic effects were seen in female Wistar rats after gavage administration of toluene in olive oil and gum arabic at 590 mg/kg, 5 days per week for 6 months (Wolf et al., 1956); in

Sprague Dawley rats after inhalation exposure at 1,481 ppm, 6 hours per day, 5 days per week for 6 months (API, 1980); or in OFA rats after inhalation exposure at 1,000 ppm, 6 hours per day, 5 days per week for 6 months (Gradiski et al., 1981).

Hematologic Effects

Leukocytosis, decreased thrombocyte and erythrocyte counts, and bone marrow hypoplasia were observed for mice exposed to toluene (grade unspecified) by inhalation at 1,000 ppm for 20 days or at 4,000 ppm for 8 weeks (Horiguchi and Inoue, 1977; Bruckner and Peterson, 1981b). A transient, slight granulopenia followed by granulocytosis was seen in rabbits administered toluene (grade unspecified) by gavage at 865 mg/kg for 6 days (Braier, 1973). These effects may have been due to the presence of benzene as a contaminant (percent not reported) in the toluene (USEPA, 1987).

No effect on hematologic values was reported for rats exposed to toluene by inhalation at 1,000 ppm for 6 weeks (Jenkins et al., 1970; Bruckner and Peterson, 1981b) or Sprague Dawley rats exposed to toluene by inhalation at 1,000 ppm, 8 hours per day, 7 days per week for 13 weeks (Tahti et al., 1983). No effect on hemoglobin concentration, hematocrit, or leukocyte count was observed for rats, guinea pigs, dogs, or monkeys exposed to toluene by inhalation continuously at 103 ppm or at 1,092 ppm for 8 hours per day, 5 days per week for 6 weeks (Jenkins et al., 1970). No hematologic effects were seen in female Wistar rats after gavage administration of toluene in olive oil and gum arabic at 590 mg/kg, 5 days per week for 6 months (Wolf et al., 1956); in Sprague Dawley rats after inhalation exposure at 1,481 ppm, 6 hours per day, 5 days per week for 6 months (API, 1980); in OFA rats at 1,000 ppm, 6 hours per day, 5 days per week for 6 months (Gradiski et al., 1981); or in F344 rats at 299 ppm, 6 hours per day, 5 days per week for 24 months (Gibson and Hardisty, 1983).

Central Nervous System Effects

The brain is highly vascularized and has a high lipid content. Therefore, the high lipid solubility of toluene indicates the possibility of a wide distribution in the brain following exposure (Benignus, 1981a). The initial uptake of toluene (after a 10-minute inhalation exposure) was greatest in the medulla/pons, followed by midbrain, cerebellum, thalamus, frontal cortex, hippocampus, caudate, and hypothalamus (Gospe and Calaban, 1988). The toluene uptake correlated with the total lipid content of each brain region. In spite of the clinical and epidemiologic data implicating toluene as a neurotoxicant, there are few studies that have systematically studied this problem in animals.

The central nervous system response to toluene is biphasic--an initial excitable phase followed by central nervous system depression (Contreras et al., 1979). At vapor concentrations less than 2,000 ppm or exposure for less than 30 minutes, increased locomotor activity in rats (Yamawaki and Sarai, 1982), operant response rates in rats and mice (Weiss et al., 1979; Glowa, 1981; Moser and Balster, 1981, 1985; Wood et al., 1983; Bushnell et al., 1985), and sensitivity to shock and heat in rats (Contreras and Bowman, 1982) were generally observed. Exposure to toluene at concentrations higher than 2,000 ppm suppressed activity (Cohr and Stokholm, 1979; Moser and Balster, 1981). Exposure of rats to toluene at 1,000 ppm produced excitability, followed by depression of cortical activity which resulted in coma (Contreras et al., 1979). Brief inhalation exposure at 3,500-4,500 ppm for 50 minutes impaired the cognitive and motor abilities in macaque monkeys (Taylor and Evans, 1985). In rats, Ikeda and Miyake (1978) reported impaired learning after repeated toluene exposure at 4,000 ppm, 2 hours per day for 60 days.

Hearing loss was reported for rats exposed to toluene at 7,550 mg/m³ for 8 hours per day for 3 days or 5,660 mg/m³ for 14 hours per day as weanlings or as young adults (Pryor et al., 1984). Toluene given to F344 and Sprague Dawley rats at 620 mg/kg by gavage once per day for 4 weeks was shown to produce hearing loss by damaging the outer hair cells of the inner ear (Sullivan, 1986). Continuous inhalation exposure of male Sprague Dawley rats to toluene at 320 ppm resulted in decreased weight of the whole brain and the cerebral cortex; the phospholipid content of the cerebral cortex was significantly decreased (Kyrklund et al., 1987).

Pryor et al. (1983a) examined the effects of 14week inhalation exposure of weanling rats to toluene and found that toluene had no consistent effect on measures of forelimb and hind limb grip strength, motor activity, startle reactivity (acoustic or air-puff stimuli), or reactivity to a thermal stimulus. n-Hexane, a known neurotoxicant, had marked effects on neuromuscular components of this neurobehavioral test battery. However, toluene-exposed rats were found to acquire a multisensory conditional avoidance response more slowly than controls and had an altered component of the brainstem auditoryevoked response. Toluene-exposed animals were also tested in a tone-intensity discrimination task and found to be deficient. In a subsequent study, Pryor et al. (1983b) reported that weanling rats exposed to toluene were impaired in learning a conditioned avoidance response if the conditioning stimulus was a 20-kHz tone; learning was not affected by toluene exposure if the training cue was visual or somatosensory. These behavioral measurements were made during the course of repeated toluene exposure. In subsequent studies, rats were tested 2.5 months after cessation of exposure; hearing of toluene-exposed animals was unimpaired at 4 kHz, slightly impaired at 8 kHz, and markedly impaired at 12 kHz or above. Rebert et al. (1983), using electrophysiologic techniques, examined the auditory effects of toluene 2.5 months after cessation of exposure: the thresholds for brainstem auditoryevoked responses were increased twofold, and the latency-intensity functions were consistent with the occurrence of sensory loss, i.e., ototoxicity. Therefore, unlike solvents such as n-hexane, there is little evidence that toluene produces peripheral neuropathy. However, if exposure occurs repeatedly in young animals, toluene produces behavioral and electrophysiologic alterations indicating toxicity.

Fetotoxicity and Teratogenicity

Skeletal anomalies were observed in the fetuses of rats and mice exposed to toluene by inhalation or gavage during gestation at doses that were not toxic to the dams. Cleft palates were seen in the fetuses of CD®-1 mice given 1 ml/kg (870 mg/kg) toluene in cottonseed oil by gavage on days 6-15 of gestation (Nawrot and Staples, 1979). At this dose, an increase in embryonic

deaths and reduced fetal weights were also observed. An increase in irregular sternebrae or extra fused ribs was seen in the fetuses of CFY rats continuously exposed to toluene at 400 ppm by inhalation on days 9-14 of gestation (Hudak and Ungvary, 1978). An increase in rudimentary 14th ribs or in extra ribs was seen in the fetuses of ICR mice exposed to toluene at 1,000 ppm by inhalation for 6 hours per day on days 1-17 of gestation (Shigeta et al., 1981, 1982) and in the fetuses of CFY rats exposed to toluene at 266 ppm by inhalation for 24 hours per day on days 7-14 of gestation (Tatrai et al., 1980) or exposed at 1,000 pm for 24 hours per day on days 7-15 (Ungvary, 1985). A significant increase in the number of fetuses with 13 ribs was observed for CD®-1 mice exposed to pesticide-grade toluene at 400 ppm by inhalation for 7 hours per day on days 6-16 of gestation (Courtney et al., 1986). Decreased weights, but no malformations, were observed for the fetuses of CFLP mice continuously exposed to toluene at 133 ppm by inhalation on days 6-13 of gestation (Hudak and Ungvary, 1978). Retarded bone ossification and inhibition of growth, but no teratogenic effects, were observed for the fetuses of Charles River rats exposed to toluene (99.96% pure) at 400 ppm by inhalation for 6 hours per day on days 6-15 of gestation (LBI, 1978a). Deaths, but no teratogenic effects, were observed for the fetuses of New Zealand rabbits continuously exposed to toluene at 266 ppm by inhalation on days 6-20 of gestation (Ungvary and Tatrai, 1985).

Genetic Toxicology

Toluene has been studied extensively for genotoxic effects both in vitro and in vivo, and the overwhelming weight of evidence indicates that the chemical is not genotoxic. A summary of these results is presented in Table 1. The positive responses reported in in vivo studies may have resulted from artifacts of the protocol or possibly from contaminants in the toluene samples. For example, the detection of single-strand breaks reported by Sina et al. (1983) was probably a secondary effect of cell lysis rather than direct interaction of toluene with nuclear DNA because it occurred only when cytotoxicity was greater than 30%. The studies reporting induction of chromosomal aberrations by toluene

TABLE 1. SUMMARY OF THE GENETIC TOXICOLOGY STUDIES OF TOLUENE

Test System/Reference	Endpoint	Dose	Results	
acteria				
Bacillus subtilis				
McCarroll et al., 1981a	Growth inhibition due to DNA damage		Negative	
Escherichia coli				
Fluck et al., 1976	Growth inhibition due to DNA damage	25 μl/plate	Negative	
McCarroll et al., 1981b	Growth inhibition due to DNA damage	0.01.101/51545	Negative Negative	
Mortelmans and Riccio, 1980	Gene mutation	0.01-10 µl/plate	Negative	
Salmonella typhimurium				
Mortelmans and Riccio, 1980	Growth inhibition due to DNA damage	0.001-0.01 μl/plate	Negative	
	Gene mutation	0.01-10 µl/plate	Negative	
Anderson and Styles, 1978	Gene mutation		Negative	
LBI, 1978b	Gene mutation	0.001-5 µl/plate	Negative	
Nestmann et al., 1980	Gene mutation	0.00.00	Negative	
Florin et al., 1980	Gene mutation	0.03-30 µmol/plate	Negative	
Snow et al., 1981	Gene mutation	0.3-100 µl/plate	Negative	
Bos et al., 1981	Gene mutation	0-2,000 µg/plate	Negative	
Spanggord et al., 1982	Gene mutation	0-5 mg/plate	Negative Negative	
Haworth et al., 1983	Gene mutation	0-1,000 µg/plate	Negative	
east				
Saccharomyces cerevisiae				
LBI, 1978b	Mitotic gene conversion	0.001-5 µl/plate	Negative	
Mortelmans and Riccio, 1980	Mitotic gene conversion	0.001%-5%	Negative	
	Mitotic crossing over	0.001%-5%	Negative	
	Gene mutation	0.001%-5%	Negative	
lammalian Cells in Vitro				
Mouse lymphoma L5178Y cells				
LBI, 1978b	Trifluorothymidine resistance	0-0.3 µl/ml	Negative	
McGregor et al., 1988	Trifluorothymidine resistance	6.25-500 μg/ml	Equivoca	
Chinese hamster ovary cells				
Evans and Mitchell, 1980	Sister chromatid exchange	0.0025%-0.04%, 21.4 h	Negative	
	Sister chromatid exchange	0.0125%-0.21%, 2 h	Negative	
Human lymphocytes	Cinton harmania and an alaman	0.1.500()	Namativa	
Gerner-Smidt and Friedrich, 1978	Sister chromatid exchange	0-1,520 µg/ml	Negative	
lammalian Cells in Vivo				
Rat			=	
Dobrokhotov, 1972	Chromosomal aberrations	0.8 g/kg/d for 12 d	(a) Positive	
Lyapkalo, 1973	Chromosomal aberrations	1 g/kg/d for 12 d	(a) Positive	
Dobrokhotov and Enikeev, 1975	Chromosomal aberrations	80 ppm, 4 h/d for 4 mo	(a) Positive	
LBI, 1978b Sina et al., 1983	Chromosomal aberrations DNA single-strand breaks	0-214 mg/kg 0-3 mM	Negative (b) Positive	
·	<u>-</u>			
Mouse Kirkhart, 1980	Micronucleus induction	0-1,000 mg/kg	Negative	
Topham, 1980	Sperm head abnormalities	0-1,5 mg/kg/d	Negative	
10pnam, 1980 LBI, 1981	Dominant lethal mutations	400 ppm, 6 h/d, 5 d/wk	Negative	
•		for 8 wk	J	
Tice et al., 1982	Sister chromatid exchange	0-32.4 mmol/kg	Negative	
Gad-El-Karim et al., 1984	Micronucleus induction	860-1,720 mg/kg	Negative	
	Chromosomal aberrations	860-1,720 mg/kg	Negative	

⁽a) Purity of chemical unspecified; possible contamination with benzene. (b) Greater than 30% cell lethality

(Dobrokhotov, 1972; Lyapkalo, 1973; Dobrokhotov and Enikeev, 1975) were difficult to evaluate because the types of aberrations scored were unclear, cells scored from a group of animals were pooled rather than analyzed individually, and, in one case (Dobrokhotov and Enikeev, 1975), there was no indication of the numbers of cells actually scored. Further, none of these positive studies specified the purity of the toluene sample used. Since nonreagent-grade toluene is frequently contaminated with varying amounts of benzene, it is possible that the increased incidence of chromosomal aberrations reported was due to exposure to benzene, a demonstrated in vivo clastogen. Similar contamination of toluene samples (purity unspecified) evaluated in other in vitro assays (e.g., for mutation induction in bacteria) would not be expected to give positive responses because benzene is negative in these assays (NTP, 1986a; Huff et al., 1989).

Several investigators have examined tolueneexposed factory workers for cytogenetic effects (Forni et al., 1971; Funes-Cravioto et al., 1977; Maki-Paakkanen et al., 1980; Bauchinger et al., 1982). The studies that reported increased levels of chromosomal aberrations or sister chromatid exchanges (SCEs) in exposed workers compared with unexposed populations failed either to adequately document that chemical exposure was to toluene alone (Funes-Cravioto et al., 1977) or to consider the data from smokers separately from those from nonsmokers (Bauchinger et al., 1982). When Bauchinger et al. (1982) reanalyzed the SCE data according to the smoking history of the workers, they still reported a small but significant increase in SCEs in the toluene-exposed groups; however, a similar reanalysis of the chromosomal aberration data (Integrated Criteria Document Toluene) eliminated the reported difference between exposed and nonexposed workers.

The metabolites of toluene for which there are data available are also nongenotoxic. Benzyl alcohol (NTP, 1989a) was negative in bacterial assays for induction of DNA damage (Fluck et al., 1976; Oda et al., 1978) or gene mutation (Florin et al., 1980; Mortelmans et al., 1986) and did not cause DNA single-strand breaks or chromosomal aberrations in human fibroblasts in vitro

(Waters et al., 1982). Benzoic acid was negative in assays for induction of gene mutation in Salmonella (McCann et al., 1975; Anderson and Styles, 1978; Simmon and Kauhanen, 1978; Zeiger et al., 1988), mitotic recombination in yeast (Simmon and Kauhanen, 1978), SCEs in Chinese hamster ovary cells (Oikawa et al., 1980), and SCEs and chromosomal aberrations in cultured human fibroblasts (Tohda et al., 1980: Zhurkov, 1975). Hippuric acid was negative in Salmonella gene mutation assays (Milvy and Garro, 1976; Wiessler et al., 1983). The cresols (m- and p-) were negative for induction of gene mutation in Salmonella (Pool and Lin, 1982: Haworth et al., 1983) and did not induce SCEs in human fibroblasts in vitro or in mouse fibroblasts in vivo (Cheng and Kligerman, 1984). The genotoxicity profile for o-cresol was similar, with the exception of a weakly positive response in the in vitro SCE assay with human fibroblasts in which a significant increase in SCEs was observed at the highest nontoxic dose tested (Cheng and Kligerman, 1984).

Carcinogenicity

A summary of the dermal, gavage, and inhalation carcinogenicity studies with toluene which have been reported in the literature is available (CTFA, 1986; Bell et al., 1988; IARC, 1989). Results for carcinogenicity were uniformly negative, although Lijinsky and Garcia (1972) reported on the occurrence of one papilloma in 30 mice exposed to 16-20 ul toluene (as toluene vehicle controls) by topical application two times per week for 72 weeks and one carcinoma in another mouse; none occurred in the acetone vehicle controls. Toluene was used as a vehicle in numerous dermal initiation/promotion studies in mice (Poel, 1963; Frei and Stephens, 1968; Lijinsky and Garcia, 1972; Vose et al., 1981; Blackburn et al., 1984), directly in dermal application studies (Coombs et al., 1973; Doak et al., 1976; Coombs and Bhatt, 1978), and in a 3month subcutaneous implant study (Purchase and Longstaff, 1978). Application of 40 µl toluene two times per week at the initiation/promotion site on the back reduced the average number of skin tumors per mouse at week 15 for C3H mice initiated with 1 mg benzo[a]pyrene or for CD®-1 mice initiated with 2.5 µg dimethylbenz[a]anthracene followed by promotion with 1-5 µg phorbol-12-myristate-13-acetate two times per week (Weiss et al., 1986).

At week 92, the incidences of neoplasms seen at various sites were not compound related in groups of 40 male and 40 female Sprague Dawley rats given 500 mg/kg toluene (98.3% pure) in olive oil by gavage 4-5 days per week for 2 years (Maltoni et al., 1983). At the end of the study (week 141), hemolymphoreticular neoplasms were reported in 3/37 toluene-exposed males and 7/40 toluene-exposed females compared with 3/45 and 1/49 in vehicle controls (Maltoni et al., 1985). Also reported were the numbers of animals with malignant tumors (olive oil control male, 11/45 vs. toluene-exposed male, 18/40; female, 10/49 vs. 21/40) and the total number of malignancies per group (male, 12/45 vs. 23/40; female, 11/49 vs. 32/40). In another inhalation study using F344 rats (Gibson and Hardisty. 1983), the incidences of neoplasms in groups of 120 male and 120 female F344 rats exposed to air containing toluene at 0, 30, 100, or 300 ppm. 6 hours per day, 5 days per week for 2 years, were not significantly different from those in controls.

Several metabolites of toluene have been or are being evaluated in long-term studies in rodents. No evidence of carcinogenicity of benzyl alcohol was seen in male or female F344/N rats given 0. 200, or 400 mg benzyl alcohol/kg body weight 5 days per week in corn oil for 2 years (NTP. 1989a). At one-half these doses, no evidence of carcinogenicity was found in male or female B6C3F₁ mice. To study the effects of antioxidants on BHA-induced forestomach carcinogenesis, Ito exposed groups of 15 F344 rats to 2% benzoic acid in the diet for 52 weeks with and without 2% butylated hydroxyanisole (BHA) (IARC, 1988a; personal communication from N. Ito, Nagoya City University Medical School, to J. Huff, NTP, December 1988). Benzoic acid did not modify the incidences of BHA-induced forestomach neoplasms, and benzoic acid alone did not cause forestomach hyperplasia. Long-term studies of benzaldehyde are currently being evaluated (NTP, 1989b). F344/N rats and male B6C3F₁ mice were given 0, 200, or 400 mg benzaldehyde/kg body weight in corn oil by gavage, and female mice were given 0, 300, or 600 mg/ kg. Short-term studies have been completed on

o-cresol and mixed m- and p-cresols (personal communication from D. Dietz, NTP, 1989).

TOXICITY IN HUMANS

Central Nervous System Effects

Inhalation of toluene produces symptoms of nervous system dysfunction and signs of neurologic impairment (Longley et al., 1967; Benignus, 1981a), which appear to be reversible except for long-term abusers (Benignus, 1981b). After a single exposure to toluene at 50-1,500 ppm for 3-8 hours, individuals developed fatigue, drowsiness, impaired cognitive function. incoordination, and irritation of the eyes and throat; these effects increased in severity with increases in concentration and progressed to pronounced nausea, staggering gait, confusion, extreme nervousness, muscular fatigue, and insomnia lasting for several days (Ogata et al., 1970; Gamberale and Hultengren, 1972; Carpenter et al., 1976; Winneke, 1982; IPCS, 1985; Baelum et al., 1985b). Narcosis increased after exposure at 4,000-30,000 ppm, with death occurring after exposure at the highest concentrations for from a few minutes to greater than 1 hour (von Oettingen, 1942a,b; IPCS, 1985). Long-term toluene abusers (for at least 1 year) reported disturbed behavior; slow thought and speech; illusionary misinterpretations; tactile, auditory, and visual hallucinations; and delusional ideas (Evans and Raistrick, 1987). Cerebellar dysfunction, mental retardation, abnormal electroencephalograms, brain atrophy, and visual impairment were observed in long-term (6-14 years) abusers of pure toluene (Grabski, 1961; Knox and Nelson, 1966; Sasa et al., 1978; Malm and Lying-Tunnell, 1980; Lewis et al., 1981; Takeuchi et al., 1981; Lazar et al., 1983). Juntunen et al. (1985) reported that long-term occupational exposure (up to 22 years) of 43 male rotogravure printers at an estimated 117 ppm toluene had no clinically significant adverse effects on the nervous system.

Kidney and Liver Effects

Effects of toluene abuse on the kidney (pyuria, hematuria, and proteinuria) have been summarized (IPCS, 1985; USEPA, 1987). Most of the persons with symptoms or signs of toluene

sniffing were also exposed to other solvents. Renal tubule effects, indicated by metabolic acidosis (hypokalemia, hypophosphatemia, and hyperchloremia), have been associated with abusers of toluene-containing solvents (Sokol and Robinson, 1963; Taher et al., 1974; Fischman and Oster, 1979; Bennett and Forman, 1980; Kroeger et al., 1980; Moss et al., 1980; Voigts and Kaufman, 1983; Patel and Benjamin, 1986); the contribution of toluene to these effects is not clear. Nielsen et al. (1985) and Krusell et al. (1985) claim that no causal relationship exists between exposure to toluene alone and renal injury (as measured by excretion of albumin and β_{2u} -globulin). In contrast, increased protein excretion and increased excretion of erythrocytes and leukocytes/tubular epithelial cells were reported for construction workers exposed to toluene (Askergren, 1984). A positive relationship exists between alcohol consumption before exposure to toluene and the urinary excretion rate of albumin (Krusell et al., 1985). Hepatomegaly was noted for 61 airplane painters exposed to toluene at 100-1,115 ppm in air for up to 5 years (Greenberg et al., 1942). In another study, hepatomegaly was observed for 20%-50% of workers exposed to toluene at 53-80 ppm in air for 2-14 years, but biopsies from 22 of the workers indicated no pathologic changes in liver (Szilard et al., 1978). Liver impairment was observed in long-term (6-14 years) abusers of pure toluene (Grabski, 1961; Knox and Nelson, 1966; Takeuchi et al., 1981).

Hematologic Effects

Early reports (generally pre-1950) of occupational exposure ascribed myelotoxic effects to toluene (Ferguson et al., 1933; Greenberg et al., 1942; Wilson, 1943), but most of the recent evidence indicates that toluene does not cause toxic effects in blood or bone marrow (Parmeggiani and Sassi, 1954; Capellini and Alessio, 1971; Matsushita et al., 1975; Tahti et al., 1981; Yin et al., 1987). Eosinophilia, leukocytosis, low hemoglobin concentration, basophilic stippling of erythrocytes, and poikilocytosis, anisocytosis, hypochromia, and polychromasia were observed for sniffers of toluene-based glues (Sokol and

Robinson, 1963). Myelotoxic effects previously attributed to toluene are currently considered to have been the result of concurrent exposure to benzene, typically present as a contaminant in commercially available toluene (USEPA, 1987).

Teratogenicity

No studies linking toluene and birth defects have been reported. All studies or reports were of solvent mixtures containing toluene (Euler, 1967; Syrovadko, 1977; Holmberg, 1979; Hersh et al., 1985).

Carcinogenicity

No published epidemiology studies on toluene were located. At least two studies are underway: one in Sweden and one in the United States (IARC, 1988b). Several epidemiology studies in which workers were exposed to other solvents as well as to toluene are described in IARC (1989).

STUDY RATIONALE

The aromatic six-member hydrocarbon (benzene), the monomethyl derivative (toluene), and the dimethyl derivatives (xylenes) were nominated and selected for toxicology and carcinogenesis characterization because each met several of the eight criteria of the Chemical Selection Principles established by the National Toxicology Program in 1978. These include considerable production volume, widespread occupational and general population exposure, and lack of adequate long-term studies at the time these chemicals were selected and the studies designed. Additionally, long-term studies on these three chemicals would provide some indications of structure-activity associations for benzene and simple alkylbenzenes. For toluene, the shortterm studies were conducted using both the gavage and inhalation routes of exposure so that the two routes could be compared. The 2-year studies used inhalation exposure to better mimic human occupational exposure (although oral and dermal exposure also occur) and to compare the results with those from an earlier study by Gibson and Hardisty (1983).

II. MATERIALS AND METHODS

PROCUREMENT AND CHARACTERIZATION OF TOLUENE
CHARACTERIZATION OF DOSE MIXTURES
GENERATION AND MEASUREMENT OF CHAMBER

CONCENTRATIONS

Vapor Generation System

Vapor Concentration Monitoring

Vapor Concentration Uniformity in Chamber

THIRTEEN-WEEK GAVAGE STUDIES

FOURTEEN-WEEK AND FIFTEEN-WEEK INHALATION STUDIES

FIFTEEN-MONTH AND TWO-YEAR STUDIES

Study Design
Source and Specifications of Animals
Animal Maintenance
Clinical Examinations and Pathology
Statistical Methods

PROCUREMENT AND CHARACTERIZATION OF TOLUENE

Toluene was obtained in one lot (lot no. H-12-19-80) from Exxon Company, USA (Baytown, TX) as a clear, colorless liquid and was received in sixteen 55-gallon drums. Purity and identity analyses were conducted on representative samples at Midwest Research Institute (Kansas City, MO) (Appendix I). The study material was identified as toluene by infrared, ultraviolet/visible, and nuclear magnetic resonance spectroscopy.

The toluene study material was found to be greater than 99% pure, as determined by elemental analysis, Karl Fischer water analysis, and gas chromatography. Gas chromatography by two systems detected three impurities with individual peak areas less than 0.1% of the major peak area. Benzene content of the study material was determined by spiking with benzene and then quantitating against a benzene reference standard and was found by gas chromatography to be present as an impurity at a concentration of 5.7 ppm (v/v). (The calculated benzene concentrations used in these studies were 0.82, 4.1, and 8.2 ppb for the toluene exposures at 120, 600, or 1,200 ppm.)

Periodic analysis of the toluene for purity by gas chromatography and ultraviolet spectroscopy and for identity by infrared spectroscopy indicated no apparent degradation of the study material throughout the studies.

CHARACTERIZATION OF DOSE MIXTURES

Toluene dissolved in corn oil at 20 mg/ml was found to be stable for at least 2 weeks when stored protected from air and light at 5° C and at room temperature. Solutions exposed to air and light for 3 hours were chemically stable, but a 23% loss due to evaporation was observed over the 3-hour period. Dose mixtures were stored at room temperature protected from light in Nalgene® bottles for no longer than 2 weeks throughout the studies. Dose mixtures were analyzed several times during the 13-week studies

and found to be within $\pm 10\%$ of the target concentrations.

GENERATION AND MEASUREMENT OF CHAMBER CONCENTRATIONS

Vapor Generation System

Toluene vapor was generated by delivering liquid toluene to a heated Spraying Systems® atomizer that was operated with nitrogen (Appendix I). Toluene vapor was diluted with chamber ventilation air to produce the desired exposure concentrations in the chambers. The uniformity of the vapor concentrations in each exposure chamber was measured several times during the studies. Generally good chamber distribution of the toluene vapor was observed in these studies.

Vapor Concentration Monitoring

The concentration of toluene in the chambers was measured in sampled chamber air at 3.3 µ by a MIRAN® gas-phase infrared spectrophotometer connected to a Hewlett-Packard Model 3388A laboratory computer. Air from each chamber was sampled and analyzed about 5 minutes every hour. Data were collected, recorded, and reported as weekly mean exposure concentrations (Tables I2 and I3). During the 2-year studies, the time-weighted-average concentrations of toluene for each exposure group were 1.3, 119.9, 593.2, and 1,179 ppm for target concentrations of 0, 120, 600, and 1,200 ppm.

Studies for the detection of toluene aerosol in the 1,200-ppm chamber were conducted with a Sibata® P-5 Digital Dust Indicator (2-year studies) or in the 3,000-ppm chamber with a Model CI-252 (Climet Instrument Co.) aerosol particle counter (14-week studies). Aerosol was not detected in measurable quantitites.

The presence of detectable concentrations (more than 10 ppm) of residual toluene was determined by analyzing the atmosphere in all chambers at various times postexposure. Measurable concentrations occurred by 4 months after the studies began, and, after further evaluation, the animals and/or caging were indicated as the source of the residual toluene.

Vapor Concentration Uniformity in Chamber

The uniformity of the vapor concentration in each exposure chamber was measured five times over a 5-month period during the studies with the same system used to monitor the vapor concentration (used as a reference) and a second system with a different infrared monitor used for comparison with the reference. Four of the five tests that used this combined system indicated good chamber distribution; the range of variation from the reference was 3%-12%. In the fifth test, the range of variation was 26%; this large range was attributed to instrument variance and not to chamber inhomogeneity. In three subsequent tests that used only the second infrared monitor for both reference and comparison, variations from the reference position were 2%, 5%, and 14%.

THIRTEEN-WEEK GAVAGE STUDIES

Thirteen-week gavage studies were conducted to evaluate the cumulative toxic effects of repeated administration of toluene, to identify target organs, and to compare results with the inhalation study findings.

Male and female F344/N rats and B6C3F₁ mice were obtained from Charles River Breeding Laboratories. Animals were observed for 18 days (rats) or 20 days (mice) and then assigned to dose groups. Rats were 6-7 weeks old when placed on study, and mice were 7-8 weeks old.

Groups of 10 rats and mice of each sex were administered 0, 312, 625, 1,250, 2,500, or 5,000 mg/kg toluene in corn oil by gavage, 5 days per week for 13 weeks.

Rats and mice were housed five per cage. Feed and water were available ad libitum. Animals were observed two times per day; moribund animals were killed. Individual animal weights were recorded at the beginning of the studies and once per week thereafter. Further experimental details are summarized in Table 2.

At the end of the studies, animals were fasted overnight in stainless steel metabolism cages

and urine was collected. Blood samples were taken from the orbital sinus. Analyses of blood and urine were performed. Survivors were killed, and a necropsy was performed on all animals. The brain, liver, lung, right kidney, right testis, and thymus were weighed. Histologic examinations were performed on animals that died before the end of the studies, vehicle controls, and animals that received 2,500 or 5,000 mg/kg. Selected tissues of lower dose animals were examined. Tissues and groups examined are listed in Table 2.

FOURTEEN-WEEK AND FIFTEEN-WEEK INHALATION STUDIES

Fourteen- and 15-week studies were conducted to evaluate the cumulative toxic effects of repeated exposure to toluene, to identify target organs, to compare results with the gavage study findings, and to determine the concentrations to be used in the 2-year studies.

Four- to 5-week-old male and female F344/N rats and 6-week-old male and female B6C3F₁ mice were obtained from Charles River Breeding Laboratories. Animals were observed for 16 days, distributed to weight classes, and assigned to groups according to tables of random numbers. Feed was available ad libitum during non-exposure periods; water was available at all times. Further experimental details are summarized in Table 2.

Groups of 10 rats and 10 mice of each sex were exposed to air containing target concentrations of 0 (chamber controls), 100, 625, 1,250, 2,500, or 3,000 ppm toluene, 6.5 hours per day, 5 days per week for 65 exposures. Animals were observed two times per day; moribund animals were killed. Animal weights were recorded once per week.

Sperm morphologic and vaginal cytologic evaluations were performed on all surviving animals exposed at 0, 100, 625, or 1,250 ppm toluene (methods are described in Appendix G). At the end of the studies, blood was collected from the orbital sinus plexus of unfasted animals. Hematologic and biochemical analyses were performed.

TABLE 2. EXPERIMENTAL DESIGN AND MATERIALS AND METHODS IN THE STUDIES OF TOLUENE

Thirteen-Week Gavage Studies	Fourteen-Week and Fifteen-Week Inhalation Studies	Fifteen-Month and Two-Year Inhalation Studies		
EXPERIMENTAL DESIGN				
Size of Study Groups 10 males and 10 females of each species	10 males and 10 females of each species	60 males and 60 females of each species		
Doses 0, 312, 625, 1,250, 2,500, or 5,000 mg/kg toluene in corn oil by gavage; dose vol 10 ml/kg	0, 100, 625, 1,250, 2,500, or 3,000 ppm toluene by inhalation	Rats0, 600, or 1,200 ppm toluene by inhalation; mice0, 120, 600, or 1,200 ppm		
Date of First Dose Rats5/19/81; mice5/21/81	11/12/81	Rats9/27/82; mice11/8/82		
Date of Last Dose Rats8/17/81; mice8/20/81	Rats2/25/82; mice2/18/82	2 y: rats9/14/84; mice10/26/84		
Duration of Dosing 5 d/wk for 13 wk	6.5 h/d, 5 d/wk for 14 wk (mice) or 15 wk (rats)	6.5 h/d, 5 d/wk for 15 mo or 103 wk		
Type and Frequency of Observation Observed $2 \times d$; weighed initially and the $1 \times wk$	n Observed $2 \times d$; weighed initially and then $1 \times wk$	Observed $2 \times d$; weighed $1 \times wk$ for 13 wk , $1 \times 4 \text{ wk}$ to wk 92, and then $1 \times 2 \text{ wk}$		
Method of Animal Kill Carbon dioxide	Intraperitoneal injection of sodium pentobarbital, followed by exsanguination	Intraperitoneal injection of sodium pentobarbital, followed by exsanguination		
Necropsy, Histologic Examinations, a Necropsy performed on all animals; the following tissues examined for vehicle controls, 2,500 and 5,000 mg/kg groups, and all animals dying before the end of the studies: adrenal glands, aorta, brain, cecum, colon, duodenum, esophagus, gallbladder (mice), gross lesions, heart, ileum, jejunum, kidneys, liver, lungs and bronchi, mammary gland, mesenteric lymph nodes, nasal cavity and turbinates, pancreas, parathyroid glands, pituitary gland, preputial or clitoral gland (rats), prostate/testes or ovaries/uterus, rectum, regional lymph nodes (mice), salivary glands, spinal cord, spleen, sternebrae including marrow, stomach, thymus, thyroid gland, tissue masses, trachea, and urinary bladder. Tissues	Necropsy performed on all animals; histologic exams performed on all controls, 2,500- and 3,000-ppm groups, and all animals dying before the end of the studies. Tissues examined include: adrenal glands, aorta, brain, cecum, colon, duodenum, epididymis/prostate/testes or ovaries/uterus, esophagus, femur, gallbladder (mice), heart, ileum,	Necropsy and histologic exams performed on all animals except 3 high dose female mice; the following tissu examined: adrenal glands, brain, cecum, colon, duodenum, epididymis/prostate/testes or ovaries/uterus, esophagus, femur including marrow, gross lesions and tissue masses with regional lymph nodes, heart and aori ileum, jejunum, kidneys, liver, lungs and mainstem bronchi, mammary gland, mandibular lymph nodes, nas cavity and turbinates, pancreas, part thyroid glands, pituitary gland, preptial or clitoral gland (rats), rectum, salivary glands, spleen, stomach, thy mus, thyroid gland, trachea, and urinary bladder. Blood taken for hemat		

ANIMALS AND ANIMAL MAINTENANCE

examined in other groups include brain,

kidneys, liver, and urinary bladder.

Blood and urine collected for analysis

before terminal kill; organs weighed

Strain and Species F344/N rats; B6C3F₁ mice

at necropsy

F344/N rats; B6C3F₁ mice

F344/N rats; B6C3F₁ mice

logic analysis before scheduled kill and

organs weighed at necropsy for 10

male rats, 10 female rats, and 10 fe-

male mice from each group at 15 mo

studies. Blood collected for analysis

before terminal kill; organs weighed

at necropsy. Sperm morphologic and

all surviving animals in the control, 100-, 625-, and 1,250-ppm groups

vaginal cytologic exams performed for

TABLE 2. EXPERIMENTAL DESIGN AND MATERIALS AND METHODS IN THE STUDIES OF TOLUENE (Continued)

Thirteen-Week Fourteen-Week and Fifteen-Week Fifteen-Month and Gavage Studies Inhalation St							
ANIMALS AND ANIMAL MAINTEN	ANCE (Continued)						
Animal Source Charles River Breeding Laboratories	Charles River Breeding Laboratories	Charles River Breeding Laboratories					
Portage, MI)	(Portage, MI)	(Kingston, NY)					
Study Laboratory nternational Research and Development Corporation	International Research and Development Corporation	International Research and Development Corporation					
Method of Animal Identification Ratsear tag; micetoe clip	Ratsear tag; micetoe clip	Ratsear tag; micetoe clip					
Fime Held Before Study Rats18 d; mice20 d	16 d	Rats12 d; mice26 d					
Age When Placed on Study Rats6-7 wk; mice7-8 wk	Rats6-7 wk; mice8 wk	Rats6-7 wk; mice9-10 wk					
Age When Killed Rats19-20 wk; mice20-21 wk	21-22 wk	15 mo: rats72-73 wk; mice75-76 w 2 y: rats110-111 wk; mice113-114 wk					
Necropsy Dates Rats8/18/81; mice8/20/81	Rats2/23/82-2/26/82; mice2/16/82-2/19/82	15 mo: rats12/28/84-12/29/84; femal mice2/7/84; 2 y: rats9/24/84- 9/28/84; mice11/5/84-11/9/84					
Method of Animal Distribution Animals distributed to weight classes and then assigned to cages by one table of ran- dom numbers and to groups by another able of random numbers		Same as 13-wk studies					
Diet NIH 07 Rat and Mouse Ration (Zeigler Bros., Inc., Gardners, PA); available ad ibitum	Same as 13-wk studies, but feed removed during exposure	Same as 14- and 15-wk studies					
Water Automatic watering system (Edstrom Industries, Waterford, WI); available ad libitum	Same as 13-wk studies	Same as 13-wk studies					
Bedding Beta Chips hardwood bedding Northeastern Products, Inc., Warrensburg, NY)	None	None					
C ages Polycarbonate	Stainless steel wire mesh (Unifab, Inc., Portage, MI)	Same as 14- and 15-wk studies					
C age Filters Nonwoven polyester fiber	None	None					
Animals per Cage	1	1					
Other Chemicals on Study in the San	ne Room None	None					

TABLE 2. EXPERIMENTAL DESIGN AND MATERIALS AND METHODS IN THE STUDIES OF TOLUENE (Continued)

Thirteen-Week Gavage Studies

fluorescent light 12 h/d

Fourteen-Week and Fifteen-Week Inhalation Studies

Fifteen-Month and Two-Year Inhalation Studies

ANIMALS AND ANIMAL MAINTENANCE (Continued)

Animal Room or Chamber (for Inhalation Studies) Environment Temp--mean, 72.4° F, range, 63°-82° F; hum--mean, 59.8%, range, 44%-82%;

Temp--74°-80° F during exposure: hum--45%-55% during exposure; fluorescent light 12 h/d

Temp--69°-81° F; hum--23%-75%; fluorescent light 12 h/d; 12-14 room air changes/h

At the end of the 14- and 15-week studies, survivors were anesthetized with sodium pentobarbital and killed by abdominal agrta incision. A necropsy was performed on all animals except those excessively autolyzed or cannibalized. The brain, heart, liver, lungs, right kidney, right testis, and thymus of all animals surviving to the end of the studies were weighed. Histologic examinations were performed on animals that died before the end of the studies, controls, and animals that were exposed at 2,500 and 3,000 ppm. A bone marrow examination was performed on selected animals. Tissues and groups examined are listed in Table 2.

FIFTEEN-MONTH AND TWO-YEAR STUDIES

Study Design

Groups of 60 rats of each sex were exposed to toluene at target concentrations of 0 (chamber controls), 600, or 1,200 ppm, 6.5 hours per day, 5 days a week for 15 months or 103 weeks. Groups of 60 mice of each sex were exposed at 0, 120, 600, or 1,200 ppm on the same schedule.

At 15 months, 10 male and 10 female rats and 10 female mice from each group had blood samples taken from the orbital sinus plexus; the erythrocyte count, total leukocyte count, hemoglobin concentration, hematocrit value, leukocyte differential count, and methemoglobin concentration were determined. The brain, liver, and right kidney were weighed at necropsy, and histologic examinations were performed on controls and animals at 1,200 ppm.

Source and Specifications of Animals

The male and female F344/N rats and B6C3F1 (C57BL/6N, female × C3H/HeN MTV⁻, male) mice used in these studies were produced under strict barrier conditions at Charles River Breeding Laboratories under a contract to the Carcinogenesis Program. Breeding stock for the foundation colonies at the production facility originated at the National Institutes of Health Repository. Animals shipped for study were progeny of defined microflora-associated parents that were transferred from isolators to barriermaintained rooms. Rats were shipped to the study laboratory at 4-5 weeks of age and mice at 5-6 weeks of age. The rats were quarantined at the study laboratory for 2 weeks and the mice for 4 weeks. Thereafter, a complete necropsy was performed on five animals of each sex and species to assess their health status. Rats were placed on study at 6-7 weeks of age and mice at 9-10 weeks of age. The health of the animals was monitored during the course of the studies by serologic analysis of controls at 15 months and 2 years (Appendix E).

Animal Maintenance

Rats and mice were housed individually. Feed was removed during exposure periods; otherwise, feed (Appendix F) and water were available ad libitum. Cages were rotated during these studies. Further details of animal maintenance are given in Table 2.

Clinical Examinations and Pathology

All animals were observed two times per day, and clinical signs were recorded every 4 weeks. Body weights were recorded once per week for the first 13 weeks of the study, once every 4 weeks until week 92, and then once every 2 weeks. Mean body weights were calculated for each group. Animals found moribund and those surviving to the end of the studies were humanely killed. A necropsy was performed on all animals including those found dead (in this study, three high dose female mice were missing after week 70). In some cases, not all samples of a particular organ were saved or some were autolyzed (e.g., mandibular lymph nodes and thymus gland in male rats, clitoral gland in 1,200-ppm female rats, and gallbladder in male and female mice). Thus, the number of animals from which particular organs or tissues were examined microscopically varies and is not necessarily equal to the number of animals that were placed on study.

During necropsy, all organs and tissues were examined for grossly visible lesions. All major tissues were fixed and preserved in 10% neutral buffered formalin, processed and trimmed, embedded in paraffin, sectioned, and stained with hematoxylin and eosin for microscopic evaluation. Tissues examined are listed in Table 2.

When the pathology evaluation was completed by the laboratory pathologist and the pathology data entered into the Toxicology Data Management System, the slides, paraffin blocks, and residual formalin-fixed tissues were sent to the NTP Archives. The slides, blocks, and residual wet tissues were audited for accuracy of labeling and animal identification and for thoroughness of tissue trimming. The slides, individual animal necropsy records, and pathology tables were sent to an independent pathology quality assessment laboratory. The individual animal records and pathology tables were compared for accuracy, slides and tissue counts were verified, and histotechnique was evaluated. All tissues with a tumor diagnosis, all potential target tissues, and all tissues from a randomly selected 10% of the animals were re-evaluated microscopically by a quality assessment pathologist. Nonneoplastic lesions were evaluated for accuracy and consistency of diagnosis in the potential target organs, in the randomly selected 10% of animals, and in tissues with unusual incidence patterns or trends such as the nose and kidney in rats.

The quality assessment report and slides were submitted to a Pathology Working Group (PWG) Chairperson, who reviewed microscopically all potential target tissues and any other tissues for which there was a disagreement in diagnosis between the laboratory and quality assessment pathologists. Representative examples of potential chemical-related nonneoplastic lesions and neoplasms and examples of disagreements in diagnosis between the laboratory and quality assessment pathologists were examined by the PWG. The PWG, which included the quality assessment pathologist and other pathologists experienced in rodent toxicology, examined the tissues without knowledge of dose group or previously rendered diagnoses. When the consensus diagnosis of the PWG differed from that of the laboratory pathologist, the diagnosis was changed to reflect the opinion of the PWG. This procedure has been described, in part, by Maronpot and Boorman (1982) and Boorman et al. (1985). The final pathology data represent a consensus of contractor pathologists and the NTP Pathology Working Group. For subsequent analysis of pathology data, the diagnosed lesions for each tissue type are combined according to the guidelines of McConnell et al. (1986).

Statistical Methods

Survival Analyses: The probability of survival was estimated by the product-limit procedure of Kaplan and Meier (1958) and is presented in the form of graphs. Animals were censored from the survival analyses at the time they were found to be missing or dead from other than natural causes; animals dying from natural causes were not censored. Statistical analyses for a possible dose-related effect on survival used the method of Cox (1972) for testing two groups for equality and Tarone's (1975) life table test for a dose-related trend. All reported P values for the survival analysis are two-sided.

Calculation of Incidence: The incidence of neoplastic or nonneoplastic lesions is given as the ratio of the number of animals bearing such lesions at a specific anatomic site to the number of animals in which that site was examined. In most instances, the denominators include only those animals for which the site was examined histologically. However, when macroscopic examination was required to detect lesions (e.g., skin or mammary tumors) prior to histologic sampling, or when lesions could have appeared at multiple sites (e.g., lymphomas), the denominators consist of the number of animals on which a necropsy was performed.

Analysis of Tumor Incidence: The majority of tumors in this study were considered to be incidental to the cause of death or not rapidly lethal. Thus, the primary statistical method used was a logistic regression analysis, which assumed that the diagnosed tumors were discovered as the result of death from an unrelated cause and thus did not affect the risk of death. In this approach, tumor prevalence was modeled as a logistic function of chemical exposure and time. Both linear and quadratic terms in time were incorporated initially, and the quadratic term was eliminated if it did not significantly enhance the fit of the model. The dosed and control groups were compared on the basis of the likelihood score test for the regression coefficient of dose. This method of adjusting for intercurrent mortality is the prevalence analysis of Dinse and Lagakos (1983), further described and illustrated by Dinse and Haseman (1986). When tumors are incidental, this comparison of the time-specific tumor prevalences also provides a comparison of the time-specific tumor incidences (McKnight and Crowley, 1984).

In addition to logistic regression, alternative methods of statistical analysis were used, and the results of these tests are summarized in the appendixes. These include the life table test (Cox, 1972; Tarone, 1975), appropriate for rapidly lethal tumors, and the Fisher exact test and the Cochran-Armitage trend test (Armitage, 1971; Gart et al., 1979), procedures based on the overall proportion of tumor-bearing animals.

Tests of significance include pairwise comparisons of each dosed group with controls and a test for an overall dose-response trend. Continuity-corrected tests were used in the analysis of tumor incidence, and reported P values are one-sided. The procedures described above also were used to evaluate selected nonneoplastic lesions. (For further discussion of these statistical methods, see Haseman, 1984.)

Historical Control Data: Although the concurrent control group is always the first and most appropriate control group used for evaluation, there are certain instances in which historical control data can be helpful in the overall assessment of tumor incidence. Consequently, control tumor incidences from the NTP historical control data base (Haseman et al., 1984, 1985) are included for those tumors appearing to show compound-related effects.

Analysis of Continuous Variables: For all end points, dosed groups were compared with the control group using the nonparametric multiple comparison test of Dunn (1964) or Shirley (1977). Jonckheere's test (Jonckheere, 1954) was used to assess the significance of the dose response trends and to determine whether Dunn's or Shirley's test was more appropriate for pairwise comparisons.

III. RESULTS

RATS

THIRTEEN-WEEK GAVAGE STUDIES
FIFTEEN-WEEK INHALATION STUDIES
FIFTEEN-MONTH STUDIES
TWO-YEAR STUDIES

Body Weights and Clinical Signs Survival Pathology and Statistical Analyses of Results

MICE

THIRTEEN-WEEK GAVAGE STUDIES
FOURTEEN-WEEK INHALATION STUDIES
FIFTEEN-MONTH STUDIES
TWO-YEAR STUDIES

Body Weights and Clinical Signs
Survival
Pathology and Statistical Analyses of Results

GENETIC TOXICOLOGY

THIRTEEN-WEEK GAVAGE STUDIES

All rats that received 5,000 mg/kg died during the first week, and 8/10 male and 1/10 female rats that received 2,500 mg/kg died before the end of the studies (Table 3). No other compoundrelated deaths occurred. The final mean body weight of males that received 2,500 mg/kg was 19% lower than that of vehicle controls. Clinical signs included prostration, hypoactivity, ataxia, piloerection, lacrimation, and excessive salivation in the 2,500 and 5,000 mg/kg groups and body tremors in the 2,500 mg/kg groups. These signs reflect onset of death. The relative liver and kidney weights for female rats that received 1,250 or 2,500 mg/kg and for males that received 625 or 1,250 mg/kg were increased relative to those for vehicle controls (Table 4). The relative heart weights for female rats that received 1,250 or 2,500 mg/kg were increased compared with

that for vehicle controls. None of the differences in the results of the hematologic or serum chemical analyses (Appendix H) or urinalyses was considered to be biologically meaningful. Several increases and decreases were observed (Table H1). Necrosis of the brain, consisting of neuronal necrosis in the dentate gyrus and Ammons horn of the hippocampus, was seen in male and female rats that received 1,250 or 2,500 mg/kg (Table 5). In addition to the hippocampal lesions, necrosis and/or mineralization was present in the granular cell layer of the cerebellar cortex. Hemorrhage was present in the mucosa, submucosa, or muscularis of the urinary bladder of male and female rats in the two highest dose groups. Kidney sections were examined in particular for the occurrence of hyaline droplets. and there was no evidence of an increase in the proximal tubules of the kidney of exposed rats.

TABLE 3. SURVIVAL AND MEAN BODY WEIGHTS OF RATS IN THE THIRTEEN-WEEK GAVAGE STUDIES OF TOLUENE

		Mean Body Weights (grams)							
Dose (mg/kg)	Survival (a)	Initial (b)	Final	Change (c)	to Vehicle Controls (percent)				
IALE									
0	10/10	127 ± 3	331 ± 6	+204 ± 5					
312	10/10	128 ± 3	344 ± 6	$+216 \pm 5$	104				
625	10/10	126 ± 3	350 ± 6	$+224 \pm 5$	106				
1,250	10/10	127 ± 3	340 ± 6	$+213 \pm 4$	103				
2,500	(d) 2/10	127 ± 3	269 ± 16	$+148 \pm 20$	81				
5,000	(e) 0/10	127 ± 3	(f)	(f)	(f)				
EMALE									
0	10/10	107 ± 2	201 ± 3	+94 ± 2					
312	10/10	107 ± 2	200 ± 4	$+93 \pm 2$	100				
625	10/10	106 ± 2	195 ± 4	$+89 \pm 3$	97				
1,250	10/10	107 ± 2	202 ± 3	$+95 \pm 2$	100				
2,500	(e) 9/10	108 ± 2	200 ± 4	$+91 \pm 3$	100				
5,000	(e) 0/10	107 ± 2	(f)	(f)	(f)				

⁽a) Number surviving/number initially in group

⁽b) Initial group mean body weight \pm standard error of the mean. Subsequent calculations are based on animals surviving to the end of the study.

⁽c) Mean body weight change of the survivors \pm standard error of the mean

⁽d) Week of death: 3,3,6,7,7,8,8,10

⁽e) Week of death: all 1

⁽f) No data are reported due to 100% mortality in this group.

TABLE 4. ANALYSIS OF SELECTED ORGAN WEIGHTS OF RATS IN THE THIRTEEN-WEEK GAVAGE STUDIES OF TOLUENE (a) $\,$

Organ	Vehicle	Control	312 mg	g/kg	625	mg	/kg	1,250	m	g/kg	2,500	n	ng/kg
MALE	·				<u></u> .								
Number weig	hed 10		10			10			10		(b)	2	
Body weight													
(grams)	315 ±	6.2	328 ±	5.8	329	±	5.8	321	±	6.4	238	±	7.5
Brain													
Absolute	1,828 ±	12	1,810 ±	28	1,835	±	17	1,795	±	18	*1,544	±	48
Relative			5.5 ±				0.08			0.10			0.003
Heart		3.00		3.0.	3.0	_		0.0	_		0.0 .	_	3.000
Absolute	1,058 ±	28	1,110 ±	33	1,120	±	38	1,115	±	26	1,114 :	±	10
Relative	3.4 ±		3.4 ±				0.08			0.09	*4.7		
Right kidney				-	3	_		0.0				_	J
Absolute	1,084 ±	14	1,159 ±	34	*1,213	±	39	**1,292	±	34	*1,227	±	114
Relative		0.06	3.5 ±				0.06	**4.0			**5.1		
Liver		J.00	J.0 -	J.V.	3.1	_		1.0	_	00	J.1 .		7.02
Absolute	10,490 ±	360	11,310 ±	300	*11,850	±	390	**14,400	+	480	*14,130	+	1.220
Relative	33.3 ±		34.5 ±		*35.9			**45.0			**59.4		
FEMALE													
Number weig	hed 10		10			10			10			9	
Body weight													
(grams)	183 ±	3.2	182 ±	3.5	175	+	3.8	181	+	27	180	+	3 4
(B. a)	100 =	0.2	102 1	0.0	110	_	0.0	101	_	2.1	100 .	-	0.4
Brain													
Absolute	1,718 ±	19	1,688 ±	30	1,698	±	24	1,693	±	19	**1,625	ŧ	17
Relative		0.22	9.3 ±				0.15			0.12	9.1		
Heart	V.= _	J.22	V.0 ±	J.10	3.1	_	0.10	J. 4	_	J.12	J.1 .	-	V.10
Absolute	693 ±	16	703 ±	27	692	+	25	*753	+	12	**790 :	+	26
Relative		0.08	3.9 ±	0.10	4.0	_	0.11	**4.2	_	0.05	**4.4	_	
Right kidney	0.0 =	J.00	0.0 -	J.20	4.0	_	~	7.4	_	0.00		-	
Absolute	686 ±	12	676 ±	19	652	+	36	*733	+	18	**803	+	26
Relative	3.8 ±		3.7 ±		3.7		0.17	**4.1			**4.5		
Liver	0.0 ±	0.00	J., 1	V.U I	0.1	_	V-11	4.1	_	0.00	4.0 .	_	V.12
Absolute	5,596 ±	119	5,822 ±	177	5,730	4	225	**6,780	+	169	**8,918	+	995
Relative	30.7 ±		31.9 ±		32.7			**37.5			**49.6		1.53
relative	30.7 I	10.0	31.5 I	0.40	32.7	T	0.07	TT37.0	工	0.00	45.0	_	1.00

⁽a) Mean in milligrams per gram necropsy body weight (relative) or milligrams (absolute) unless otherwise specified \pm standard error; P values are vs. the vehicle controls by Dunn's test (Dunn, 1964) or Shirley's test (Shirley, 1977).

 $⁽b) \, Sample \, size \, was \, in adequate \, for \, reliable \, statistical \, comparisons \, with \, vehicle \, controls.$

^{*}P<0.05

TABLE 5. NUMBERS OF RATS WITH SELECTED LESIONS IN THE THIRTEEN-WEEK GAVAGE STUDIES OF TOLUENE (a)

Site/Lesion	Vehicle Control	312 mg/kg	625 mg/kg	1,250 mg/kg	2,500 mg/kg	5,000 mg/kg
MALE						
Brain						
Necrosis	0	0	0	**6	**8	0
Urinary bladder Hemorrhage	0	(b) 0	0	0	2	**6
FEMALE	·	(0) 0	·	•	_	-
Brain Nameria	0	0	0	0	**7	0
Necrosis Urinary bladder	U	U	υ	U	,	U
Hemorrhage	0	0	0	0	0	3

⁽a) Ten animals were examined in each group unless otherwise specified. All rats at 5,000 mg/kg and one female at 2,500 mg/kg died during week 1.

FIFTEEN-WEEK INHALATION STUDIES

Eight of 10 male rats exposed at 3,000 ppm died during week 2 (Table 6). The final mean body weights of rats exposed at 2,500 or 3,000 ppm were 15% or 25% lower than that of controls for males or 15% or 14% lower for females. Clinical signs included dyspnea in all exposed groups, except males exposed at 3,000 ppm and females exposed at 1,250 ppm, and ataxia in rats exposed at 2.500 or 3.000 ppm; other clinical signs observed in the gavage studies were not observed in these inhalation studies. The relative weights of the heart, liver, and kidney for female rats exposed at 2,500 or 3,000 ppm, of the kidney and liver for male rats exposed at 1,250 or 2,500 ppm, and of the heart for male rats exposed at 2,500 ppm were increased compared with those for controls (Table 7). None of the differences in the results of the hematologic or serum chemical analyses was considered to be biologically meaningful (Table H2). Plasma cholinesterase activity

decreased as the exposure concentration increased, and the leukocyte count was decreased for female rats at 1,250 ppm or higher. No compound-related effects were seen on sperm or on the estrous cycle. The toxic lesions seen in animals exposed by gavage (see Table 5) were not observed in animals exposed by inhalation.

Dose Selection Rationale: Because of the decreases in body weights in each sex at 2,500 and 3,000 ppm, deaths in the 3,000-ppm males, and, to a lesser extent, the increases in relative organ weights, inhalation exposure concentrations selected for rats for the 15-month and 2-year studies were 0,600, or 1,200 ppm toluene, 6.5 hours per day, 5 days per week. Also considered useful in the selection of exposure concentrations was the lack of any toxicity or carcinogenicity findings from a previously reported inhalation study using the same strain of rats exposed at 0, 30, 100, or 300 ppm (Gibson and Hardisty, 1983).

⁽b) Nine animals were examined.

^{**}P<0.01 vs. the vehicle controls

TABLE 6. SURVIVAL AND MEAN BODY WEIGHTS OF RATS IN THE FIFTEEN-WEEK INHALATION STUDIES OF TOLUENE

		Mea	n Body V	Veights	(grams)		Final Weight Relative
Concentration (ppm)	Survival (a)	Initial (b)	Fina	l (c)	Chan	ge (d)	to Controls (percent)
MALE							
0	10/10	177 ± 4	356	± 4	+179	± 4	
100	10/10	186 ± 6	366	t 6	+180	± 4	103
625	10/10	187 ± 5	361	£ 6	+174	± 6	101
1,250	10/10	181 ± 5	360	Ŀ 7	+179	± 6	101
2,500	10/10	177 ± 5	302	t 4	+125	± 4	85
3,000	(e) 2/10	152 ± 4	268	t 26	+134	± 25	75
FEMALE							
0	10/10	127 ± 2	211	E 3	+84	± 2	
100	10/10	132 ± 3	210	ե 2	+78	± 3	100
625	10/10	129 ± 3	213	<u> 4</u>	+84	± 4	101
1,250	10/10	127 ± 3	209	£ 3	+82		99
2,500	10/10	126 ± 3	180	E 2	+54	± 3	85
3,000	10/10	116 ± 2	182	E 2	+66	± 1	86

⁽a) Number surviving/number initially in group

⁽b) Initial group mean body weight ± standard error of the mean. Subsequent calculations are based on animals surviving to

the end of the study.

(c) Final body weight data represent weights taken at 14 weeks because, due to the unusually long terminal necropsy period, some animals were killed before the final weighing at 15 weeks.

(d) Mean body weight change of the survivors ± standard error of the mean

⁽e) Week of death: all 2

TABLE 7. ANALYSIS OF ORGAN WEIGHTS OF RATS IN THE FIFTEEN-WEEK INHALATION STUDIES OF TOLUENE (a)

Organ	Contr	ol	100 pp	m	625 pp	om	1,250	ppm	2,500 p	pm	3,000	ppm
MALE			-						, , , , , , , , , , , , , , , , , , , ,			
Number weigh	ed 10		10		10		10)	10		(b) 2	2
Body weight (g												
	356 ±	3.3	367 ±	6.5	362 ±	7.0	362 ±	7.5	**304 ±	4.4	*280 ±	29.5
Brain												
Absolute	1,825 ±	37	1,865 ±	23	1,865 ±	21	1,830 ±	24	1,753 ±	24	1,853 ±	17.5
Relative		0.10		0.10		0.11		0.13	**5.8 ±	0.11	*6.7 ±	
Heart			•	••			V		• • • •	****	V., _	
Absolute	955 ±	16	1,019 ±	25	971 ±	14	990 ±	24	900 ±	21	871 ±	: 87
Relative	2.7 ±	-	•	0.04		0.03	2.7 ±		**3.0 ±	0.05	**3.1 ±	
Right kidney				0.0 -					0.0 -		0,1	
Absolute	1,161 ±	18	1,238 ±	27	1,206 ±	30	1,242 ±	- 28	1,147 ±	28	1,108 ±	100.0
Relative	3.3 ±	0.04		0.05		0.07	*3.4 ±		**3.8 ±		**4.0 ±	
Liver	0.0 _	0.04	0.4 -	0.00	0.0 =	0.01	0.4 _	. 0.00	0.0 _	0.01	4.0 _	. 0.00
Absolute	12,760 ±	260	13,210 ±	370	13,610 ±	360	14,110 ±	420	12,470 ±	300	13,310 ±	1 620
Relative	35.8 ±			0.61		0.55	**38.9 ±		**41.0 ±	0.60	**47.6 ±	
Lung	00.0 =	0.00	00.0 =	0.01	00 =	0.00	00.0 =	. 0.0.		0.00		
Absolute	1,187 ±	22	1,255 ±	18	1,213 ±	21	1,271 ±	- 38	1,139 ±	18	1,087 ±	72
Relative		0.06		0.05		0.05		0.08	**3.8 ±	0.07	**3.9 ±	
Right testis	0.0 ±	0.00	J.4 ±	0.00	J.4 I	0.00	3.0 1	. 0.00	3.0 ±	0.01	J.5 _	. 0.10
Absolute	1,471 ±	25	1,532 ±	20	1,524 ±	21	1.538 ±	. 00	1 421 +	18	1 411 4	: 35
Relative	4.1 ±								1,431 ± **4.7 ±		1,411 ± *5,1 ±	
Cauda	4.1 1	0.05	4.2 _	0.05	4.2 1	0.05	4.3 1	0.08	**4.7 I	0.04	· 5.1 ±	0.41
	143 ±	c	152 ±	0	1 477 ±	4	154 4					
Absolute		6	152 1	8	147 ±	4	154 ±	: 5			•	-
Right epididyn Absolute	284 ±	7	304 ±	7	289 ±	4	289 ±	· 8				
FEMALE		·	• • • • • • • • • • • • • • • • • • • •	•		_						
FEMALE												
Number weigh	ned (c) 10		10		10		10)	10		10)
Body weight (g	rams)											
Dody weight (6	209 ±	3.4	213 ±	2.4	213 ±	3 3	208 ±	. 29	**185 ±	2.2	**188 ±	- 98
	200 1	U. TE	210 1	4.4	210 <u>1</u>	0.0	200 1	. 0.2	100 1	4.4	100 7	. 4.0
Brain												
Absolute	1,729 ±	23	1,750 ±	21	1,729 ±	12	1,739 ±	: 8	1,691 ±	22	1,703 ±	- 97
Relative	8.3 ±	0.12	8.3 ±	0.16	8.2 ±	0.17		0.12	**9.1 ±	0.14	**9.1 ±	
Heart	0.0 I	0.12	ع. د.ه	0.10	0.2 I	0.17	0.4 1	. 0.12	9.1 I	0.14	9.1 3	. 0.43
Absolute	646 ±	8	663 ±	10	653 ±	15	631 ±	: 16	C10 ±	10	C40 4	: 8
Relative	3.1 ±	-		10 0.04	3.1 ±	15 0.05	3.0 ±		613 ± *3.3 ±	10 0.07	643 ± **3.4 ±	
	3.1 I	0.04	0.1 <u>r</u>	V.U4	3.1 I	0.03	J.U 1	. 0.07	'3.3 I	0.07	J.4 1	. 0.03
Right kidney	704 ±	10	700 ±	1.4	717 ±	10	740 4	: 17	710 ±	1.1	717 4	- 17
Absolute		13		14		12	740 ±		710 ±	11	717 ±	
Relative	3.4 ±	0.06	$3.4 \pm$	0.07	3.4 ±	0.07	*3.6 ±	0.05	**3.8 ±	0.05	**3.8 ±	0.05
Liver	77 11 F ±	1	77 4077 ±	010	7 100 ±	101	7 170 -	. 100	# 000 ±	O.F	### COO =	150
Absolute		155	7,407 ±	216	7,188 ±	121	7,172 ±		7,302 ±		**7,698	
Relative	34.1 ±	0.55	$34.8 \pm$	0.85	33.8 ±	0.33	34.6 ±	0.46	**39.5 ±	0.54	**41.1 =	0.54
Lung	(3) 0 4 4 4	0.	070		(1) 000 1	10	0.05	- 01	000 1	00	000	- 10
Absolute Relative	$(d) 944 \pm (3) 45 \pm (4)$	21		15	$(d) 966 \pm (d) 4.5 \pm$	19	967 ±	21 0.08	909 ± **4.9 ±	22	922 ±	
	$(d) 4.5 \pm$	0.07	46 +	0.07	14175 +	0.08	<i>a</i> 7 →	- 11110	771 L) +	0.10	++1 C →	0.08

⁽a) Mean in milligrams per gram of necropsy body weight (relative) or milligrams (absolute) unless otherwise specified ± standard error; P values are vs. the controls by Dunn's test (Dunn, 1964) or Shirley's test (Shirley, 1977).

(b) Sample size was inadequate for reliable statistical comparisons with controls.

(c) Unless otherwise specified

⁽d) Lungs of nine animals were weighed.

^{*}P<0.05

^{**}P<0.01

FIFTEEN-MONTH STUDIES

In the nasal cavity, mild-to-moderate degeneration of the olfactory and respiratory epithelium was more obvious in toluene-exposed rats (male: control, 5/10; 600 ppm, 10/10; 1,200 ppm, 10/10; female: 2/10; 10/10; 9/10) and goblet cell hyperplasia was somewhat increased (male: 3/10; 8/10; 5/10; female: 2/10; 5/10; 6/10), whereas other lesions were seen in a few exposed rats (necrosis: three males and four females; metaplasia: one male and three females), and the incidences and severity of chronic inflammation were greater in exposed females than in controls (5/10; 9/10; 8/10). Hyperplasia of the alveolar and bronchiolar epithelium was found in two males and three females in the 1,200-ppm group and in one control female. The severity of nephropathy was slightly increased in exposed female rats. No other nonneoplastic lesions or any neoplastic lesions were observed which were considered to be related to toluene exposure. No compound-related effects were seen for relative

organ weights (Table H7). Results of hematologic analyses did not suggest any compound-related effects (Table H3).

TWO-YEAR STUDIES

Body Weights and Clinical Signs

The initial mean body weights of rats exposed at 1,200 ppm were 9% greater than those of controls (Table 8 and Figure 2); in the early weeks of the studies, these differences were diminished. Mean body weights of male rats exposed at 1,200 ppm were 4%-8% lower than those of controls from week 72 to the end of the study. Mean body weights of female rats exposed at 1,200 ppm were 4%-7% lower than those of controls from week 92 to the end of the study. An evaluation of mean body weights averaged over the first and second years indicates late decreases of 4%-5% for the 1,200-ppm groups. No compound-related clinical signs were recorded.

TABLE 8. MEAN BODY WEIGHTS OF RATS IN THE TWO-YEAR INHALATION STUDIES OF TOLUENE

Week		r Control		600 ppm			1,200 ppm	
on	Av. Wt.	Number	Av. Wt.	Wt. (percent	Number	Av. Wt.	Wt. (percent	Number
Study	(grams)	Weighed	(grams)	of controls)	Weighed	(grams)	of controls)	Weighed
MALE			•					
1	103	60	109	106	60	112	109	60
2	131	60	133	102	60	140	107	60
3	161	60	165	102	60	170	105	60
4	185	60	189	102	60	193	104	60
5	200	60	210	105	59	212	106	60
6	221	60	225	102	59	222	100	60
7	233	60	236	101	59	235	101	60
8	244	60	246	101	59	245	101	60
9	251	60	255	102	59	258	103	60
10	26 0	60	264	102	59	267	103	60
11	270	60	272	101	59	269	100	60
12	281	60	281	100	59	280	100	60
13	285	60	286	100	59	287	101	60
16	311	60	307	99	59	306	99	60
20	335	60	332	99	59	328	98	60
24	350	60	347	99	59	342	98	60
28	362	60	359	99	59	354	98	60
32	370	60	373	101	59	366	99	59
36	381	60	376	99	59	371	97	59
40	390	60	381	98	59	368	94	(a) 58
44	397	60	390	98	58	383	97	59
48	389	60	394	101	58	375	97	(a) 58
52	404	(a) 53	402	100	58	387	96	(a) 57
56	396	(a) 58	383	97	58	383	97	(a) 54
60	398	59	395	99	58	384	96	(a) 56
64	399	58	400	100	58	395	99	(a) 58
68	403	(b) 48	410	102	(b) 48	3 9 1	97	(b) 49
72	409	46	407	100	47	388	95	(a) 46
76	411	(a) 45	406	99	47	389	95	47
80	405	46	406	100	47	387	96	43
84	412	46	412	100	44	385	93	42
88	408	44	407	100	44	384	94	38
92	408	38	416	102	40	389	95	34
94	412	36	412	100	40	384	93	33
96	405	35	406	100	40	384	95	32
98	407	31	392	96	38	376	93	29
100	401	31	393	98	(a) 30	371	92	(a) 26
102	393	30	405	103	30	373	95	26
104	396	30	400	101	29	381	96	22
Mean for v			222	444 -		****	100 5	
1-13	217.3		220.8	101.6		222.3	102.3	
16-52	368.9		366.1	99.2		358.0	97.0	
56 -104	403.9		403.3	99.9		384.0	95.1	

TABLE 8. MEAN BODY WEIGHTS OF RATS IN THE TWO-YEAR INHALATION STUDIES OF TOLUENE (Continued)

Week	Chambe	r Control		600 ppm			1,200 ppm	
on Study	Av. Wt. (grams)	Number Weighed	Av. Wt. (grams)	Wt. (percent of controls)	Number Weighed	Av. Wt. (grams)	Wt. (percent of controls)	Number Weighed
EMALE								· · · · · · · · · · · · · · · · · · ·
1	85	60	93	109	60	93	109	60
2	99	60	104	105	60	106	107	60
3	113	60	118	104	60	121	108	60
4	124	60	130	105	60	131	106	60
5	132	60	139	105	60	143	108	60
6	143	60	145	101	60	146	102	60
7	149	60	150	101	60	152	102	60
8	153	60	155	101	(a) 59	156	102	60
9	156	60	160	103	60	164	105	60
10	159	60	162	103	60	168	106	60
11	163	60	168	103	60	170	104	60
12	168	60	174	104	60	173	103	60
					• •	177	103	60
13	171	60	175	102	60	184	102	60
16	180	60	183	102	60		99	60
20	192	60	192	100	60	190		
24	198	60	200	101	60	198	100	59
28	203	60	207	102	60	203	100	59
32	207	60	217	105	60	212	102	59
36	218	60	220	101	60	217	100	59
40	222	(a) 59	220	99	60	213	96	(a) 58
44	228	(a) 59	232	102	60	223	98	59
48	233	60	235	101	60	227	97	(a) 58
52	238	(a) 59	239	100	60	229	96	(a) 58
56	236	60	233	99	60	227	96	(a) 58
60	245	60	241	98	5 9	238	97	(a) 58
64	245	59	250	102	59	247	101	58
68	257	(b) 48	259	101	(b) 48	249	97	(b) 47
72	263	48	254	97	(a) 42	256	97	48
76	267	48	262	98	48	258	96	48
80	263	48	262	100	44	256	97	48
84	270	(a) 45	265	98	44	258	96	44
88	272	46	274	101	44	262	97	42
92	280	43	281	100	(a) 43	265	95	40
94	283	41	279	99	42	264	93	36
96	280	40	275	98	41	264	94	(a) 34
98	281	38	277	99	37	267	95	34
100	282	38	281	100	37	261	93	33
102	281	36	279	99	36	266	95	31
104	287	33	284	99	35	274	96	30
Mean for v								
1-13	139.6		144.1	103.2		146.2	104.7	
16-52	211.9		214.5	101.2		209.6	98.9	
56-104	268.3		266.0	99.1		257.0	95.8	

⁽a) The number of animals weighed was lower than the number of animals surviving. (b) Interim kill occurred,

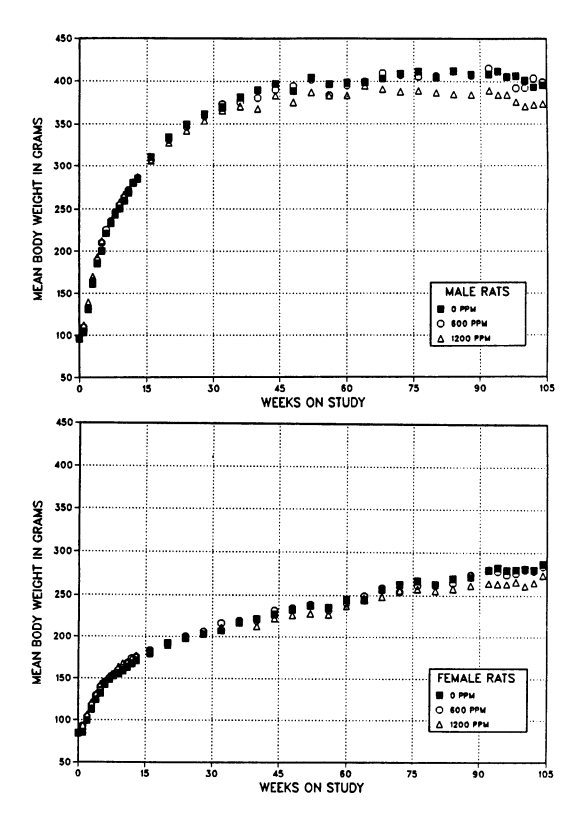


FIGURE 2. GROWTH CURVES FOR RATS EXPOSED TO TOLUENE BY INHALATION FOR TWO YEARS

Survival

Estimates of the probabilities of survival for male and female rats exposed to toluene at the concentrations used in these studies and for controls are shown in Table 9 and in the Kaplan and Meier curves in Figure 3. No significant differences in survival were observed between any groups of either sex.

Pathology and Statistical Analyses of Results

This section describes the statistically significant or biologically noteworthy changes in the incidences of rats with neoplastic or nonneoplastic lesions of the nose, kidney, and forestomach.

Summaries of the incidences of neoplasms and nonneoplastic lesions, individual animal tumor diagnoses, statistical analyses of primary tumors that occurred with an incidence of at least 5% in at least one animal group, and historical control incidences for the neoplasms mentioned in this section are presented in Appendixes A and B for male and female rats, respectively.

TABLE 9. SURVIVAL OF RATS IN THE TWO-YEAR INHALATION STUDIES OF TOLUENE

	Chamber Control	600 ppm	1,200 ppn
MALE (a)			
Animals initially in study	60	60	60
Animals removed at 15 mo	10	10	10
Natural deaths	6	12	5
Moribund kills	14	11	23
Killed accidentally	1	0	0
Animals surviving until study termination	(b) 29	(b) 27	22
Mean survival (days)	641	639	630
Survival P values (c)	0.17	0.99	0.21
FEMALE (a)			
Animals initially in study	60	60	60
Animals removed at 15 mo	10	10	10
Natural deaths	7	7	6
Moribund kills	11	8	14
Animals surviving until study termination	(b) 32	35	30
Mean survival (days)	658	654	643
Survival P values (c)	0.52	0.84	0.57

⁽a) First day of termination period: 729

⁽b) Animals killed at the end of the study; an additional animal died or was killed during the termination period and was combined, for statistical purposes, with those killed at termination.

⁽c) The result of the life table trend test is in the control column, and the results of the life table pairwise comparisons with the controls are in the dosed columns.

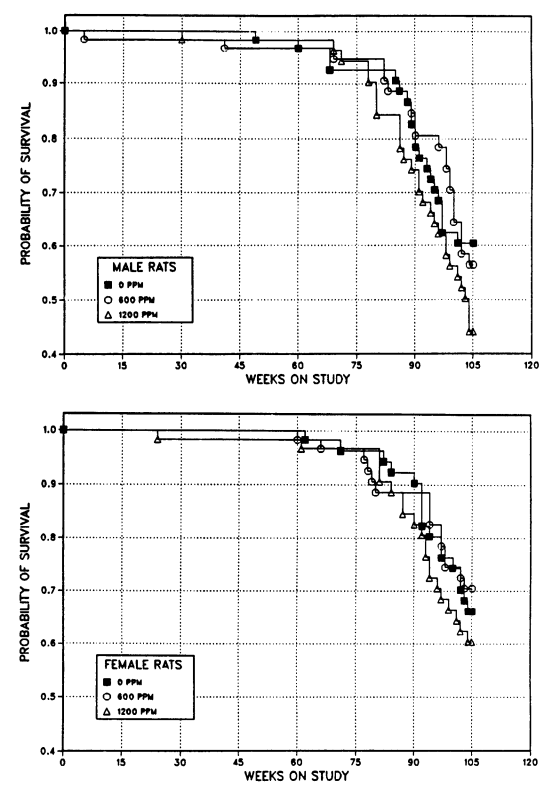


FIGURE 3. KAPLAN-MEIER SURVIVAL CURVES FOR RATS EXPOSED TO TOLUENE BY INHALATION FOR TWO YEARS

Nose: Erosion of the olfactory epithelium and degeneration of the respiratory epithelium were significantly (P<0.05) increased in exposed rats (erosion of the olfactory epithelium--male: control, 0/50; 600 ppm, 3/50; 1,200 ppm, 8/49; female: 2/49; 11/50; 10/50; degeneration of the respiratory epithelium--male: 15/50; 37/50; 31/49; female: 29/49; 45/50; 39/50). Inflammation of the nasal mucosa and respiratory metaplasia of the olfactory epithelium were observed at significantly (P<0.05) increased incidences in exposed female rats (inflammation of the nasal mucosa: 27/49; 42/50; 41/50; metaplasia of the olfactory epithelium: 0/49; 2/50; 6/50). This spectrum of lesions is not unusual in inhalation exposure studies of organic solvents, and the lesions were, for the most part, of mild severity. A squamous cell carcinoma of the mucosa was seen in one female rat at 1,200 ppm. Squamous cell neoplasms of the nose, nares, or nasal cavity have not been observed in 349 chamber control female F344/N rats or in 1.643 untreated controls.

Kidney: The evaluation of the kidney was done in two stages; first, a diagnostic evaluation was made on the single sections typically prepared

for NTP carcinogenesis studies, and then additional sections were made and evaluated for males.

The severity of nephropathy was increased with exposure concentration in male and female rats (Table 10). Renal tubule cysts were somewhat increased in male rats at 1,200 ppm (control, 1/50; 600 ppm, 2/50; 1,200 ppm, 5/50). Renal neoplasms observed in the original evaluation include tubule adenomas in one male rat at 600 ppm and two male rats at 1,200 ppm, a carcinoma of the renal transitional epithelium in one male rat at 600 ppm, a renal tubule carcinoma in one female rat at 1.200 ppm, and a sarcoma in one female rat at 1,200 ppm. The historical incidence of renal tubule adenomas, adenocarcinomas, or carcinomas (combined) is 1/346 (0.3%) in chamber control male F344/N rats and 14/1,590 (0.9%) in untreated controls. The historical incidence of renal tubule adenomas, adenocarcinomas, or carcinomas (combined) is 1/347 (0.3%) in chamber control female F344/N rats and 2/1,639 (0.1%) in untreated controls: the historical incidence of renal sarcomas is 0/347 in chamber control female F344/N rats and 0/1,639 in untreated controls.

TABLE 10. INCIDENCES AND SEVERITY OF NEPHROPATHY IN RATS IN THE TWO-YEAR INHALATION STUDIES OF TOLUENE

		Male		Female				
	Control	600 ppm	1,200 ppm	Control	600 ppm	1,200 ppm		
Incidence	49/50	48/50	48/50	49/50	48/50	49/50		
Severity (a)								
None	1	2	2	1	2	1		
Minimal	1	0	1	3	4	4		
Mild	19	11	10	26	17	17		
Moderate	15	21	11	16	23	14		
Marked	14	16	26	4	4	14		
Mean severity (b)	2.8	3.0	*3.2	2.4	2.5	*2.7		

⁽a) Number of rats with indicated severity

⁽b) 0 = none; 1 = minimal; 2 = mild; 3 = moderate; 4 = marked

^{*}P<0.05 vs. controls

Because tubular cell neoplasms were observed in three exposed male rats and none was observed in chamber controls, the male rat kidneys were evaluated further by a more extensive sampling procedure in order to more accurately assess the actual incidences of tubular cell neoplasms of the kidney. The standard sampling method for microscopic examination involves a single longitudinal section taken from the center of the left and right kidneys, plus additional sections of any grossly visible potential neoplasms. The additional pathology procedure involved embedding the remaining pieces of each kidney which had been retained as part of the wet tissues and step sectioning the embedded tissue every millimeter to yield an average of approximately six additional sections per animal. The results of the original diagnoses, additional tissue review

after eliminating duplicate diagnoses from the original review, and the combined data are presented in Table 11. Based on these data, no chemical-related increases were observed for neoplasms of the kidney.

Forestomach: Ulcers were marginally increased in exposed male rats (control, 4/50; 600 ppm, 7/50; 1,200 ppm, 9/49). A squamous cell papilloma was observed in one female rat at 1,200 ppm, and a squamous cell carcinoma was observed in a second female rat at 1,200 ppm. The historical incidence of squamous cell papillomas or carcinomas (combined) of the forestomach is 0/344 in chamber control female F344/N rats and 3/1,623 (0.2%) in untreated controls. These two neoplasms were considered to be chance occurrences and unrelated to toluene exposure.

TABLE 11. NUMBERS OF MALE RATS WITH RENAL TUBULE LESIONS IN THE FIFTEEN-MONTH AND TWO-YEAR INHALATION STUDIES OF TOLUENE

Lesion	Control	600 ppm	1,200 ppm
Number examined	60	60	60
Original single sections			
Hyperplasia	4	4	0
Adenoma	0	1	2
Carcinoma	0	0	0
Subsequent sections (a)			
Hyperplasia	0	(b) 3	2
Adenoma	5	4	0
Carcinoma	Ö	Ō	Õ
Composite data			
Hyperplasia	4	6	2
Adenoma	5	5	$\tilde{2}$
Carcinoma	ŏ	ŏ	ō

⁽a) Six additional sections per rat

⁽b) Two of the rats with hyperplasia also had adenomas.

THIRTEEN-WEEK GAVAGE STUDIES

All mice that received 5,000 mg/kg died during week 1, and 4/10 male and 4/10 female mice that received 2,500 mg/kg and 1/10 female mice that received 1,250 mg/kg died before the end of the studies (Table 12). The final mean body weight of males at 2,500 mg/kg was 16% lower than that of vehicle controls. Clinical signs included subconvulsive jerking, prostration, impaired grasping reflex, bradypnea, hypothermia, hypoactivity, and ataxia in mice at 2,500 and 5,000 mg/kg. Relative liver weights were increased for male and female mice that received 1,250 or 2,500 mg/kg (Table 13). None of the differences in the results of the hematologic or serum chemical analyses (Table H4) or urinalyses was considered to be biologically meaningful. Myocardial fiber degeneration was observed in 3/10 males and 2/10 females at 5,000 mg/kg; all animals in these groups died during the first week of exposure.

FOURTEEN-WEEK INHALATION STUDIES

Five of 10 male mice and 10/10 female mice at 3,000 ppm died during the first 2 weeks; an additional male at 3,000 ppm, 7/10 female mice at 2,500 ppm, 1/10 female mice at 1,250 ppm, and 1/10 female mice at 625 ppm died before the end of the studies (Table 14). Final mean body weights of all exposed groups were 7%-13% lower than those of controls. Dyspnea was observed primarily at 2,500 and 3,000 ppm. The other clinical signs observed in the gavage studies were not seen in these inhalation studies. The relative liver weights for mice exposed at 625 ppm or higher and lung weights for mice exposed at 1,250 ppm or higher and the relative kidney weights for female mice exposed at 1,250 ppm or higher were greater than those for controls (Table 15). None of the differences in the results of the hematologic or serum chemical analyses was considered to be biologically meaningful (Table H5). Centrilobular hepatocellular

TABLE 12. SURVIVAL AND MEAN BODY WEIGHTS OF MICE IN THE THIRTEEN-WEEK GAVAGE STUDIES OF TOLUENE

		Mea	n Body Weights	(grams)	Final Weight Relative
Dose (mg/kg)	Survival (a)	Initial (b)	Final	Change (c)	to Vehicle Controls (percent)
MALE					
0	10/10	23.0 ± 0.5	32.1 ± 1.1	$+9.1 \pm 0.7$	
312	10/10	23.3 ± 0.4	31.7 ± 0.7	$+8.4 \pm 0.6$	98.8
625	10/10	23.3 ± 0.5	31.5 ± 0.7	$+8.2 \pm 0.4$	98.1
1,250	10/10	23.0 ± 0.4	30.0 ± 0.8	$+7.0 \pm 0.6$	93.5
2,500	(d) 6/10	22.6 ± 0.6	26.8 ± 0.7	$+4.0 \pm 0.4$	83.5
5,000	(e) 0/10	22.8 ± 0.4	(f)	(f)	(f)
FEMALE					
0	10/10	19.1 ± 0.3	24.1 ± 0.4	$+5.0 \pm 0.3$	
312	10/10	19.5 ± 0.4	25.2 ± 0.6	$+5.7 \pm 0.4$	104.6
625	10/10	18.6 ± 0.6	23.9 ± 1.0	$+5.3 \pm 0.4$	99.2
1,250	(g) 9/10	18.6 ± 0.5	24.0 ± 0.7	$+5.4 \pm 0.4$	99.6
2,500	(h) 6/10	18.5 ± 0.3	23.5 ± 0.6	$+5.0 \pm 0.3$	97.5
5,000	(e) 0/10	19.1 ± 0.5	(f)	(f)	(f)

⁽a) Number surviving/number initially in group

⁽b) Initial group mean body weight \pm standard error of the mean. Subsequent calculations are based on animals surviving to the end of the study.

⁽c) Mean body weight change of the survivors ± standard error of the mean

⁽d) Week of death: 2,2,9,12

⁽e) Week of death: all 1

⁽f) No data are reported due to 100% mortality in this group.

⁽g) Week of death: 9

⁽h) Week of death: 1,8,8,10

TABLE 13. ANALYSIS OF ORGAN WEIGHTS OF MICE IN THE THIRTEEN-WEEK GAVAGE STUDIES OF TOLUENE (a)

Organ	Vehicle C	ontrol	312 mg	g/kg	625 n	ng/kg	1,250 m	g/kg	2,500 mg/kg		
MALE											
Number weighed (b)	10		10		10		10		6		
Body weight (grams)	26.4 ±	0.85	26.5 ±	0.62	26.2 ±	0.65	24.7 ±	0.52	**22.7 ±	0.71	
Brain											
Absolute	427 ±	6.0	424 ±	5.6	436 ±	5.3	433 ±	4.5	433 ±	14.4	
Relative	1.6 ±	0.06	1.6 ±	0.04	1.7 ±		*1.8 ±		**1.9 ±		
Heart										0.0.	
Absolute	144 ±	3.4	145 ±	5.1	149 ±	5.9	147 ±	4.8	130 ±	7.2	
Relative		0.13	5.5 ±		5.7 ±		6.0 ±		5.7 ±		
Right kidney			J.U _		J <u>-</u>	~·-V	0.0 4	J.20	V.1 ±	V.01	
Absolute	222 ±	6.0	228 ±	4.9	224 ±	5.8	212 ±	77	**185 ±	8.6	
Relative		0.15				0.13	8.6 ±		8.1 ±		
Liver	0		0.0 ==	J.12	0.0 =	3.10	0.0 ±	٠.22	U.1 ±	0.20	
Absolute	1,035 ±	28	1,071 ±	30	1,079 ±	35	1,073 ±	38	1,128 ±	47	
Relative		0.43	40.5 ±			0.98	**43.4 ±		**49.7 ±		
Right testis	00.0 ±	J. 40	-U.U -	J.U4	71.2 ±	0.00	TU.7 1	1.00	₹0.1 ±	0.07	
Absolute	(c) 117 ±	1 9	118 ±	24	114 ±	2.8	116 ±	3.7	108 ±	3.0	
Relative	(c) 4.4 ±		4.5 ±		4.4 ±		*4.7 ±		**4.8 ±		
100100110	(0, 4.4 1	J.1-1	7.0 ⊥	0.00	4.4 7	0.00	4.1 ⊥	0.11	4.0 ⊥	0.00	
FEMALE											
Number weighed	10		10		10		9		6		
Body weight (grams)	19.4 ±	0.45	20.6 ±	0.43	19.0 ±	0.76	19.3 ±	0.47	19.0 ±	0.45	
Brain											
Absolute	422 ±	6.6	445 ±	5.7	439 ±	6.0	436 ±	7.9	418 ±	12.2	
Relative		0.05	2.2 ±	0.05	2.4 ±	0.11		0.06	2.2 ±		
Heart											
Absolute	114 ±	4.7	115 ±	4.9	119 ±	5.4	118 ±	4.1	122 ±	10.1	
Relative	5.9 ±	0.17	5.6 ±		6.3 ±		6.1 ±		6.4 ±		
Right kidney	-							30	J		
Absolute	155 ±	3.7	169 ±	4.8	159 ±	4.1	160 ±	4.7	164 ±	5.1	
Relative		0.13		0.18	8.5 ±	0.25		0.11	8.7 ±		
Liver			-	30			0.0	J	J., _	0.01	
Absolute	858 ±	38	*975 ±	27	906 ±	36	955 ±	28	**1,083 ±	31	
Relative		1.25	*47.3 ±		*47.8 ±	0.72		0.88	**57.0 ±		

⁽a) Mean in milligrams per gram necropsy body weight (relative) or milligrams (absolute) unless otherwise specified ± standard error; P values are vs. the vehicle controls by Dunn's test (Dunn, 1964) or Shirley's test (Shirley, 1977). (b) Unless otherwise specified (c) Testes of nine animals were weighed.

^{*}P<0.05

^{**}P<0.01

TABLE 14. SURVIVAL AND MEAN BODY WEIGHTS OF MICE IN THE FOURTEEN-WEEK INHALATION STUDIES OF TOLUENE

		Mea	n Body Weights	(grams)	Final Weight Relative
Concentration (ppm)	Survival (a)	Initial (b)	Final	Change (c)	to Controls (percent)
MALE					
0	10/10	20.1 ± 0.6	31.8 ± 0.8	$+11.7 \pm 0.5$	
100	10/10	22.8 ± 0.7	29.4 ± 0.5	$+6.6 \pm 0.3$	92.5
625	10/10	22.6 ± 0.7	29.0 ± 0.9	$+6.4 \pm 0.4$	91.2
1,250	10/10	22.4 ± 0.6	28.9 ± 0.6	$+6.5 \pm 0.4$	90.9
2,500	10/10	21.0 ± 1.1	27.9 ± 0.7	$+6.9 \pm 0.6$	87.7
3,000	(d) 4/10	21.0 ± 0.3	28.8 ± 1.1	$+7.8 \pm 1.0$	90.6
FEMALE					
0	10/10	17.5 ± 0.3	28.6 ± 0.6	$+11.1 \pm 0.4$	
100	10/10	18.8 ± 0.2	24.9 ± 0.5	$+6.1 \pm 0.4$	87.1
625	(e) 9/10	19.4 ± 0.6	25.1 ± 0.8	$+5.7 \pm 0.5$	87.8
1,250	(f) 9/10	19.1 ± 0.4	25.4 ± 0.5	$+6.3 \pm 0.3$	88.8
2,500	(g) 3/10	15.0 ± 0.7	26.7 ± 0.7	$+9.0 \pm 0.0$	93.4
3,000	(h) 0/10	17.0 ± 0.8	(i)	(i)	(i)

⁽a) Number surviving/number initially in group

⁽a) Number surviving/number initially in group

(b) Initial group mean body weight ± standard error of the mean. Subsequent calculations are based on animals surviving to the end of the study.

(c) Mean body weight change of the survivors ± standard error of the mean

(d) Week of death: 1,1,2,2,2,11

(e) Week of death: 8

(f) Week of death: 13

⁽g) Week of death: all 1

⁽h) Week of death: 1,1,1,1,1,2,2,2,2,2 (i) No data are reported due to 100% mortality in this group.

TABLE 15. ANALYSIS OF ORGAN WEIGHTS OF MICE IN THE FOURTEEN-WEEK INHALATION STUDIES OF TOLUENE (a)

Organ	Contr	ol	100 j	ppm	625)	ppm	1,250	ppm	2,500	ppm	3,000) ppm
MALE	-						••••					
Number weigh	ed (b) 10		10		10		10)	10)	4	4
Body weight												
(grams)	29.0 ±	0.70	28.0 ±	0.52	27.8 ±	0.93	27.3 ±	0.47	27.4 ±	0.58	27.3 ±	0.7
Brain												
Absolute	442 ±	6.3	449 ±	6.1	436 ±	4.4	456 ±	7.2	433 ±	5.6	425 ±	10.5
Relative	15.3 ±	0.40	16.1 ±	0.31	15.8 ±	0.53	16.7 ±	0.30	15.8 ±	0.29	15.7 ±	0.7
Heart												
Absolute	155 ±	2.4	150 ±	6.9	150 ±	9.4	139 ±	3.4	157 ±	7.1	153 ±	7.3
Relative	5.4 ±		5.4 ±		5.4 ±			0.13		0.19		0.2
Right kidney	··· =	3,		J. J	··	·						
Absolute	282 ±	10.0	282 ±	9.4	262 ±	11.5	*254 ±	7.3	*253 ±	8.4	248 ±	5.2
Relative		0.24	10.1 ±			0.22		0.19	9.2 ±		_	0.1
Liver	J., ±	J.27	10.1 _	5.20	V.T _	0.22	0.0 1	. 0.20	Ų.J <u>_</u>	VV	0.1 4	
Absolute	1,482 ±	31	1,481 ±	38	1,425 ±	48	1,519 ±	- 38	**2,106 ±	63	**2,026 ±	48
Relative		1.29	52.9 ±	0.80	51.3 ±	0.54	*55.7 ±		**77.0 ±	2.18	**74.4 ±	
Lung	01.0 ±	1.20	02.5 ±	0.00	01.0 ±	0.04	00.7 2	. 1.20	71.0 =	2.10		. 0.0
Absolute	172 ±	30	173 ±	4.5	174 ±	6.7	(c) 173 ±	. 40	173 ±	3.5	183 ±	- 63
Relative		0.09	6.2 ±	0.17	6.3 ±	0.18	(c) 6.3 ±		*6.3 ±		*6.7 ±	
Right testis	0.0 ±	0.03	0.2 1	0.11	0.5 ±	0.10	(0) 0.5 1	. 0.10	0.5 1	0.01	0.7 ±	. 0.4
Absolute	113 ±	5 Q	121 ±	2.8	114 ±	4.2	117 ±	. 20	(c) 100 ±	5.0	104 ±	- 63
Relative	3.9 ±		*4.3 ±	0.05	4.1 ±			0.07	$(c) 3.7 \pm$			0.19
FEMALE												
Number weigh	ed (b) 10		10		9		ç	•	3	3	()
Body weight (g	rama)											
Dody weight (g	25.3 ±	0.58	23.3 ±	0.47	23.8 ±	0.60	23.7 ±	0.33	26.0 ±	0.58		-
Brain												
Abs olute	462 ±		463 ±		468 ±		466 ±		453 ±			
Relative	18.4 ±	0.63	19.9 ±	0.38	19.7 ±	0.43	19.7 ±	0.28	17.4 ±	0.91		•
Heart												
Abs olute		4.4	119 ±	3.1	(d) $125 \pm$	3.5	125 ±		148 ±			
Relative	5.0 ±	0.18	5.1 ±	0.07	(d) $5.3 \pm$	0.11	5.3 ±	0.09	**5.7 ±	0.11		•
Right kidney												
Absolute	182 ±	5.9	176 ±	5.3	183 ±	7.5	183 ±	5.5	208 ±			
Relative	$7.2 \pm$	0.28	7.6 ±	0.16	7.7 ±	0.21	*7.7 ±	0.15	*8.0 ±	0.15		-
Liver												
Absolute	1,293 ±	17	1,251 ±	30	1,300 ±	44	*1,417 ±	35	**2,058 ±	189		
Relative	51.3 ±	1.09	53.7 ±	0.89	*54.6 ±	0.92	**59.9 ±		**78.9 ±	5.57		-
Lung												
	169 ±	5.8	175 ±	4.3	(e) 175 ±	6.8	178 ±	6.5	188 ±	4.3		
Abs olute	100											

⁽a) Mean in milligrams per gram of necropsy body weight (relative) or milligrams (absolute) unless otherwise specified ± standard error; P values are vs. the controls by Dunn's test (Dunn, 1964) or Shirley's test (Shirley, 1977).

(b) Unless otherwise specified

⁽c) Organs of nine animals were weighed.

⁽d) Organs of eight animals were weighed.

⁽e) Organs of seven animals were weighed.

^{*}P<0.05

^{**}P<0.01

hypertrophy was observed in 10/10 male mice at 2,500 ppm and 4/6 male mice at 3,000 ppm. No effects on sperm count or motility or on the estrous cycle were seen.

Dose Selection Rationale: Because of body weight decreases, deaths in the 3,000-ppm group of each sex, deaths in the 2,500-ppm females, dyspnea and liver hypertrophy at 2,500 and 3,000 ppm, and differences in body weights observed for most exposed groups, inhalation exposure concentrations selected for mice for the 15-month and 2-year studies were 0, 120, 600, or 1,200 ppm toluene, 6.5 hours per day, 5 days per week. A top dose of 1,200 ppm was used to match the top dose for rats; three exposure concentrations were chosen to permit an adequate study if the top dose proved to be too high for good health maintenance.

FIFTEEN-MONTH STUDIES

Minimal hyperplasia of the bronchial epithelium was seen in 4/10 female mice at 1,200 ppm, and one female mouse exposed at 1,200 ppm had an adenocarcinoma of the lung. No other lesions were observed which were considered to be related to toluene exposure. No compound-related effects were seen on relative organ weights (Table H8) or on results of hematologic analyses (Table H6).

TWO-YEAR STUDIES

Body Weights and Clinical Signs

The initial mean body weights for all exposed groups of mice were 5%-14% higher than those of controls; these differences diminished rather quickly (Table 16 and Figure 4). Mean body weights of male mice at 1,200 ppm were generally similar to or somewhat higher than those of controls throughout the study. Mean body weights of female mice at 1,200 ppm were 4%-9% lower than those of controls from week 36 to week 76 and from week 88 to week 96. The yearly averages of the mean body weights of males were similar among groups, and those of females in the low and top exposure groups were about 4% lower than those of controls in the second year. No compound-related clinical signs were observed.

TABLE 16. MEAN BODY WEIGHTS OF MICE IN THE TWO-YEAR INHALATION STUDIES OF TOLUENE

Week	Chamber Control			120 ppm			600 ppm		1,200 ppm			
on Study	Av. Wt. (grams)	Number Weighed	Av. Wt. (grams)	Wt. (percent of controls)	Number Weighed	Av. Wt. (grams)	Wt. (percent of controls)	Number Weighed	Av. Wt. (grams)	Wt. (percent of controls)	Number Weigher	
MALE								·				
1	24.1	60	25.4	105.4	60	26.1	108.3	60	26.0	107.9	60	
2	26.5	(a) 59	26.9	101.5	60	27.4	103.4	60	27.3	103.0	60	
3	27.5	60	27.9	101.5	60	28.0	101.8	60	29.0	105.5	60	
4	28.9	60	28.9	100.0	60	29.6	102.4	60	29.5	102.1	60	
5	29.6	60	29.2	98.6	60	28.9	97.6	60	29.0	98.0	60	
6	29.6	60	29.8	100.7	60	29.9	101.0	60	29.8	100.7	59	
7	30.4	60	30.1	99.0	60	30.5	100.3	60	31.0	1 02 .0	59	
8	30 .8	60	30.5	99.0	60	30.8	100.0	60	31.5	102.3	(a) 58	
9	31.5	60	30.8	97.8	60	30.9	98.1	60	31.4	99.7	59	
10	31.5	60	31.3	99.4	60	31.1	98.7	60	31.5	100.0	59	
11	31.8	60	31.4	98.7	60	31.8	100.0	60	32.1	100.9	59	
12	31.9	60	31.7	99.4	60	31.8	99.7	60	32.8	102.8	59	
13	32.3	60	32.1	99.4	60	32.6	100.9	60	33.2	102.8	59	
16	33.0	60	32.3	97.9	60	33.0	100.0	60	33.6	101.8	59	
20	33.3	60	32.5	97.6	60	33.1	99.4	60	33.9	101.8	59	
24	33.7	60	34.0	100.9	59	34.6	102.7	60	34.4	102.1	58	
28	35 .1	60	34.4	98.0	59	34.7	98.9	60	35.2	100.3	58	
32	34.8	60	36.0	103.4	59	36.8	105.7	60	36.0	103.4	58	
36	37.4	60	36.7	98.1	59	37.8	101.1	59	37.4	1 00 .0	(a) 57	
40	37.6	60	38.4	102.1	59	39.4	104.8	58	38.0	101.1	58	
44	38.4	(a) 56	38.1	99.2	59	40.3	104.9	57	39.0	101.6	56	
48	39.2	56	38.1	97.2	(a) 58	39.7	101.3	(a) 52	38.3	97.7	56	
52	38.1	56	38.0	99.7	59	39.3	103.1	53	34.8	91.3	53	
56	36.7	55	37.0	100.8	58	38.1	103.8	(a) 48	36.3	98.9	(a) 50	
60	38.1	(a) 53	38.2	100.3	(a) 54	39.2	102.9	48	38.6	101.3	52	
64	37.9	47	39.2	103.4	53	39.8	105.0	44	38.1	100.5	48	
68	38.7	45	39.5	102.1	52	40.7	105.2	39	37.8	97.7	45	
72	38.3	44	38.4	100.3	51	39.8	103.9	37	38.9	101.6	45	
76	38.1	42	37.4	98.2	46	39.1	102.6	35	38.9	102.1	40	
80	37.4	37	38.0	101.6	43	38.7	103.5	(a) 30	38.7	103.5	(a) 34	
84	38.5	33	38.8	100.8	38	39.9	103.6	26	38.6	100.3	(a) 32	
88	37.9	30	39.0	102.9	33	40.4	106.6	26	39.8	105.0	32	
92	38.8	(a) 22	37.7	97.2	26	39.3	101.3	21	39.6	102.1	28	
94	37.8	22	38.3	101.3	(a) 22	39.0	103.2	21	38.7	102.4	(a) 27	
96	37.4	22	36.7	98.1	25	38.2	102.1	21	39.1	104.5	27	
98	37.1	21	37.4	100.8	25	37.5	101.1	20	38.1	102.7	26	
100	37.1	21	37.6	101.3	24	37.1	100.0	19	37.0	99.7	24	
102 104	36.7 38.0	20 17	37.1 37.6	101.1 98.9	24 22	36.0 36.9	98.1 97.1	19 16	36.7 38.7	100.0 101.8	22 19	
104	JO.V	11	31.0	30.3	22	30.8	31.1	10	36.1	101.0	19	
Mean for v	weeks 29.7		29.7	100.0		00.0	101.0		90.9	1 02 .0		
16-52	36.1		29.7 35.9	100.0 99.4		30.0 36.9	101.0 102.2		30.3 36.1	102.0 100.0		
10.02	30.1		30.5	30.4		30.9	102.2		30.1	100.0		

TABLE 16. MEAN BODY WEIGHTS OF MICE IN THE TWO-YEAR INHALATION STUDIES OF TOLUENE (Continued)

Week	Chamber Control		120 ppm			600 ppm			1,200 ppm		
on Study	Av. Wt.	Number Weighed	Av. Wt.	Wt. (percent of controls)	Number Weighed	Av. Wt.	Wt. (percent of controls)	Number Weighed	Av. Wt.	Wt. (percent of controls)	Number Weigher
,	.9/										
FEMA	LE										
1	19.1	59	20.5	107.3	60	21.8	114.1	60	21.6	113.1	59
2	21.1	59	21.6	102.4	59	22.1	104.7	60	21.8	103.3	59
3	21.8	59	21.4	98.2	59	22.7	104.1	60	22.7	104.1	59
4	22.7	59	23.0	101.3	59	23.5	103.5	60	23.7	104.4	59
5	23.9	59	23.6	98.7	59	23.4	97.9	60	23.8	99.6	59
6	24.2	59	24.2	100.0	59	24.6	101.7	60	24.2	100.0	59
7	24.8	59	24.2	97.6	59	25.1	101.2	60	25.6	103.2	59
8	24.9	59	24.7	99.2	59	25.9	104.0	60	26.1	104.8	59
9	25.7	59	24.7	96.1	59	25.3	98.4	60	26.1	101.6	59
10	25.7	59	25.1	97.7	59	25.6	99.6	60	25.9	100.8	59
11	26.0	59	25.6	98.5	59	26.0	100.0	60	26.6	102.3	59
12	26.4	59	25.9	98.1	59	26.7	101.1	60	27.6	104.5	58
13	26.8	59	26.2	97.8	59	27.1	101.1	60	27.5	102.6	58
16	27.6	59	26.7	96.7	59	28.3	102.5	60	27.9	101.1	58
20	28.1	59	27.0	96.1	59	27.4	97.5	60	29.0	103.2	58
24	29.0	59	28.1	96.9	59	28.3	97.6	59	28.9	99.7	58
28	29.5	59	28.6	96.9	59	28.7	97.3	59	29.6	100.3	58
32	30.4	59	29.1	95.7	59	30.6	100.7	(a) 57	30.5	100.3	58
36	33.4	59	31.0	92.8	59	32.3	96.7	(a) 58	31.6	94.6	56
40	35.7	59	32.8	91.9	59	34.8	97.5	59	32.4	90.8	56
44	36.1	58	34.3	95.0	59	36.6	101.4	59	33.8	93.6	57
48	36.1	58	33.9	93.9	59	36.5	101.1	(a) 58	33.0	91.4	57
52	34.5	58	33.8	98.0	59	34.2	99.1	(a) 58	32.7	94.8	57
56	34.1	58	33.4	97.5	(a) 55	34.9	102.3	(a) 57	32.5	95.3	(a) 55
60	36.0	56	34.8	96.7	56	36.8	102.2	59	34.2	95.0	56
64	37.3	56	35.9	96.2	55	37.1	99.5	(a) 51	34.4	92.2	56
68	36.4	(b) 45	36.0	98.9	(b) 43	38.1	104.7	(b) 49	34.7	95.3	(b) 44
72	36.6	44	34.7	94.8	(a) 41	37.5	102.5	49	35.1	95.9	43
76	37.0	45	34.6	93.5	42	37.4	101.1	46	35.6	96.2	40
80	36.4	45	34.5	94.8	41	36.6	100.5	46	35.3	97.0	41
84	37.5	44	36.5	97.3	41	38.6	102.9	43	35.8	95.5	41
88	38.0	44	36.8	96.8	40	39.5	103.9	40	36.0	94.7	38
92	39.0	40	36.5	93.6	40	39.4	101.0	37	36.8	94.4	37
94	38.5	39	37.0	96.1	38	40.4	104.9	35	36.0	93.5	37
96	37.4	39	34.6	92.5	38	39.2	104.8	32	35.6	95.2	36
98	36.7	38	34.9	95.1	38	38.0	103.5	32	35.6	97.0	36
100	36.1	36	34.7	96.1	34	38.2	105.8	32	35.7	98.9	35
102	35.8	35	35.3	98.6	34	36.8	102.8	27	34.5	96.4	34
104	36.3	30	35.3	97.2	33	36.9	101.7	24	36.5	100.6	32
dean for											
1-13	24.1		23.9	99.2		24.6	102.1		24.9	103.3	
16-52	32.0		30.5	95.3		31.8	99.4		30.9	96.6	
56-104	36.8		35,3	95.9		37.8	102.7		35.3	95.9	

⁽a) The number of animals weighed was lower than the number of animals surviving. (b) Interim kill occurred.

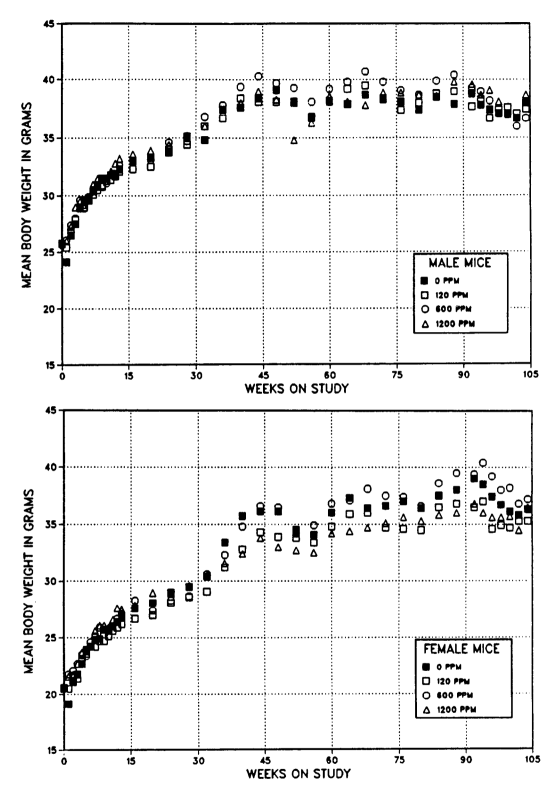


FIGURE 4. GROWTH CURVES FOR MICE EXPOSED TO TOLUENE BY INHALATION FOR TWO YEARS

Survival

Estimates of the probabilities of survival for male and female mice exposed to toluene at the doses used in these studies and for controls are shown in the Kaplan and Meier curves in Table 17 and in Figure 5. No significant differences in survival were observed between any groups of either sex. Survival in all groups of male mice, however, was inexplicably low. The particular cause or causes of early deaths in male mice were not specifically identified or recorded. However, the spectrum of nonneoplastic inflammatory lesions of the urinary and genital systems in all groups indicates that these lesions may have contributed to the early deaths (Table C5). Further, the numbers of male mice with ulcers of the prepuce (13%-23%) and/or scrotum (14%-27%) are greater than expected or than typically seen. Thus, it is reasonable to conclude that these lesions contributed to the cause of death of male mice, especially for the animals in

a moribund condition. As a comparison, survival of chamber control male mice in other studies at this same laboratory is good: mean of 80% (319/400) with a range of 56%-92% at the end of the 2-year studies.

Pathology and Statistical Analyses of Results

This section describes the statistically significant or biologically noteworthy changes in the incidences of mice with neoplastic or nonneoplastic lesions of the pituitary gland, spleen, lung, and hematopoietic system.

Summaries of the incidences of neoplasms and nonneoplastic lesions, individual animal tumor diagnoses, statistical analyses of primary tumors that occurred with an incidence of at least 5% in at least one animal group, and historical control incidences for the neoplasms mentioned in this section are presented in Appendixes C and D for male and female mice, respectively.

TABLE 17. SURVIVAL OF MICE IN THE TWO-YEAR INHALATION STUDIES OF TOLUENE

	Chamber Control	120 ppm	600 ppm	1,200 ppm
MALE (a)				
Animals initially in study	60	60	60	60
Natural deaths	23	20	17	22
Moribund kills	19	19	25	17
Killed accidentally	1	0	2	2
Animals surviving until study termination		(b) 21	16	19
Mean survival (days)	587	615	558	586
urvival P values (c)	0.99	0.36	0.74	0.66
FEMALE (a)				
Animals initially in study	60	60	60	60
Animals removed at 15 mo	10	10	10	10
Vatural deaths	11	6	16	11
Moribund kills	8	8	11	3
Killed accidentally	1	3	0	1
Animals surviving until study termination	30	33	(b) 23	32
Animals missing	0	0	0	3
fean survival (days)	635	625	633	617
Survival P values (c)	0.99	0.53	0.21	0.57

⁽a) First day of termination period: 729

⁽b) Animals killed at the end of the study; an additional animal died or was killed during the termination period and was combined, for statistical purposes, with those killed at termination.

⁽c) The result of the life table trend test is in the control column, and the results of the life table pairwise comparisons with the controls are in the dosed columns.

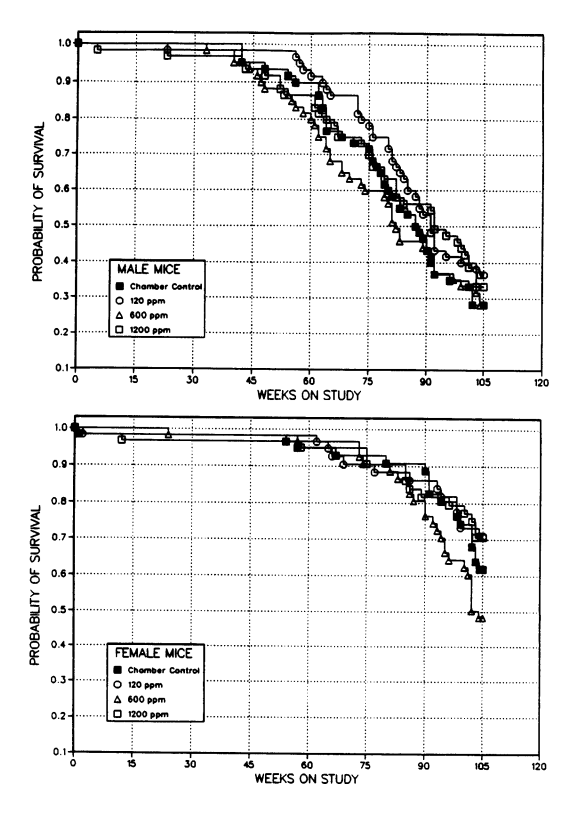


FIGURE 5. KAPLAN-MEIER SURVIVAL CURVES FOR MICE EXPOSED TO TOLUENE BY INHALATION FOR TWO YEARS

Pituitary Gland: The incidence of adenomas of the pars distalis in female mice at 600 ppm was statistically greater than that in controls (male: control, 0/59; 120 ppm, 2/58; 600 ppm, 1/58; 1,200 ppm, 0/56; female: 12/49; 19/48; 21/49; 15/46). The increased incidence in the mid exposure group of females was considered marginal and, together with a lack of supporting hyperplasia and dose response, was not considered biologically meaningful. Adenomas of the pars intermedia were seen in 0/49 control, 1/48 120-ppm, 1/49 600-ppm, and 1/46 1,200-ppm female mice. The historical incidence of neoplasms of the pars intermedia is 1/370 (0.3%) in chamber control female B6C3F₁ mice and 3/1,528 (0.2%) in untreated controls. An adenoma of the pars intermedia was seen in 1/56 1,200-ppm male mice. Although the occurrence of a single adenoma in each of the three exposure groups of females might indicate a possible effect, these neoplasms were not considered to be related to toluene exposure because the incidences did not increase with dose and could have occurred by chance and because only one neoplasm was observed in males.

Spleen: Pigmentation was observed at increased incidences in exposed male mice (male: control, 4/60; 120 ppm, 9/60; 600 ppm, 11/60; 1,200 ppm, 18/59; female: 37/50; 33/50; 34/49; 28/47).

Lung: The incidences of alveolar/bronchiolar adenomas in male mice at 120 ppm and of alveolar/bronchiolar adenomas or carcinomas (combined) at 120 and 600 ppm were lower than those in controls (adenomas: control, 8/60; 120 ppm, 1/60; 600 ppm, 2/60; 1,200 ppm, 8/60; adenomas or carcinomas, combined: 9/60; 1/60; 2/60; 9/60). These decreases are not explainable by survival differences, and because the incidence in the top dose group was not decreased, this effect was not considered to be toluene related.

Hematopoietic System: The incidences of malignant lymphomas in female mice at 120 or 1,200 ppm were lower than those in controls (control, 22/50; 120 ppm, 10/50; 600 ppm, 17/50; 1,200 ppm, 11/47). The lack of a consistent exposure concentration-related decrease precludes considering the decrease at 1,200 ppm to be other than a marginal effect not related to toluene exposure.

III. RESULTS: GENETIC TOXICOLOGY

Toluene, within a dose range of 10-1,000 µg/ plate, did not induce reverse gene mutations in four strains of S. typhimurium (TA98, TA100, TA1535, or TA1537) when tested in a preincubation protocol in the presence or absence of Aroclor 1254-induced male Sprague Dawley rat or Syrian hamster liver S9 (Haworth et al., 1983; Table J1). In the mouse lymphoma assay for induction of trifluorothymidine resistance in L5178Y/TK cells, toluene was positive in trials conducted with and without Aroclor 1254induced male F344 rat liver S9 (McGregor et al., 1988; Table J2); significant responses were noted at doses of 200 µg/ml and above, which, in all but one trial, represented the highest nonlethal dose tested. Despite the statistically

positive, reproducible responses observed in this assay, the overall conclusion was judged to be equivocal because the presence of a toluene/ water emulsion could not be ruled out conclusively, therefore leaving a question of whether acceptable dose levels had been achieved in this assay as per the study criteria set forth in McGregor et al. (1988). In cytogenetic tests with cultured Chinese hamster ovary cells, toluene did not induce sister chromatid exchanges (Table J3) or chromosomal aberrations (Table J4) when tested with doses up to 1,600 µg/ml in the presence or absence of Aroclor 1254-induced male Sprague Dawley rat liver S9; no induction of cell cycle delay, necessitating delayed harvest, was noted at any of the nonlethal doses tested.

IV. DISCUSSION AND CONCLUSIONS

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The alkylbenzenes are single-ring aromatic compounds containing one or more aliphatic side chains. The major products of commerce and those alkylbenzenes to which humans are most probably exposed include monomethylbenzene (toluene), ethylbenzene, isopropylbenzene, and the three dimethylbenzenes (xylenes). The industrial solvent toluene and the closely related benzene, xylenes, and ethylbenzene solvents account for more than 40 billion pounds produced each year in the United States: benzene, $11.7 \times$ 109 lb; ethylbenzene, 9.4×10^9 lb; toluene, 7.0 \times 109 lb; and xylenes, 10.9 \times 109 lb (Chem. Eng. News, 1988; USITC, 1988). The aromatic six-member hydrocarbon (benzene) (NTP. 1986a; Huff et al., 1988, 1989), the monoethyl derivative (ethylbenzene), the monomethyl derivative (toluene), and the dimethyl derivatives (xylenes) (NTP, 1986b; Huff et al., 1988) were nominated and selected for toxicology and carcinogenesis characterization because each met several of the selection criteria established by the National Toxicology Program (NTP, 1986a, 1988). These four solvent congeners have considerable production volume and have widespread occupational and general population exposure. At the time these studies were designed, no adequate long-term study results were available. Additionally, long-term studies on these chemicals would provide some indications of structure-activity associations for benzene and simple alkylbenzenes.

This discussion centers on the toluene studies in F344/N rats and B6C3F₁ mice and compares these findings with those on benzene and xylenes (mixed) previously reported. In early 1982, using the results of the NTP short-term studies together with the preliminary findings from the Chemical Industry Institute of Toxicology long-term studies (Gibson and Hardisty, 1983) showing an apparent lack of any significant toxic effects, the NTP decided to continue with the 2-year studies described in this Technical Report. The toluene inhalation studies were conducted at exposure concentrations up to 1,200 ppm, four times higher than those used by Gibson and Hardisty (1983).

A comparison of the effects on rats and mice exposed to toluene by the gavage and inhalation routes in the current 13- or 15-week studies indicates an apparent similarity in certain clinical

signs (e.g., dyspnea and ataxia) and organ weights (kidney and liver weight increases) and a difference in organs showing some nonneoplastic responses--brain and urinary bladder in the gavage studies in rats and none in particular for the rats and mice exposed by inhalation (liver hypertrophy in male mice). In any event, other than clinical signs of a usually transient toxicity and the brain lesions in the gavage studies, toluene-induced toxic effects were relatively rare by either route of short-term exposure.

For the longer term toluene studies, body weights for male and female rats were similar among the respective groups for both the 15month and 2-year exposure periods (see Tables 8 and H7). The 1,200-ppm group of males in the 15-month study and the 1,200-ppm group of females at the end of 2 years had mean body weights 5%-9% lower than those of the controls. Body weights for male and female mice were within 5% of those of controls among groups in the 15-month and 2-year studies, except that the body weights for the 1,200-ppm group of 10 female mice examined at 15 months were approximately 8% lower than those for controls. These relatively small differences in mean body weights are within the range of normal variations observed for untreated control F344/N rats and B6C3F₁ mice (Haseman et al., 1985) and are considered to be only minimally associated with toluene exposure. When body weights are averaged over the second year (see Tables 8 and 16), the differences become even less meaningful.

Survival for male rats at the end of the study was slightly lower than that for untreated historical control male F344/N rats (Haseman et al., 1984, 1985), especially for the 1,200-ppm group. Survival for female rats and for male and female mice at the end of the 2-year exposure was good and similar within the particular species and gender groups, although the numbers of male mice surviving were unexpectedly low in all groups. Many of the animals dying or killed in a moribund condition had inflammatory and ulcerous lesions of the penis, prepuce, and scrotum; the lesions were considered to be factors contributing to the low survival. Courses of action that were considered but not taken involved either reducing the exposure concentrations to about 900 ppm (the midpoint between the top and mid exposure groups) or altering the frequency of all exposures to three times per week (every other day). Another option was to discontinue exposure at week 64 (survival: control, 47/60; low dose, 53/60; mid dose, 44/60; high dose, 48/60) and consider these studies as 15-month studies. None of these possibilities was adopted, and the studies were continued because the decreases in survival were observed in all groups and were not considered related to toluene exposure.

Considering these body weight and survival observations in isolation from any other effects. one might reasonably conclude that higher exposure concentrations could have been used. For mice, higher exposure concentrations might have been tolerated; yet the findings from the 14-week inhalation studies supported the selected exposure concentrations and forecast that using the top exposure concentration of 1,200 ppm was appropriate. Matsumoto et al. (1971) likewise reported organ weight to body weight increases for liver, kidney, and heart in DONRYU male rats exposed to 2,000 ppm toluene by inhalation for 18 weeks. Thus, these whole-body exposures, up to 1,200 ppm for 6 hours per day, 5 days per week for 2 years, were considered both prospectively and retrospectively to have been an adequate and sufficient exposure challenge for determining the presence or absence of a carcinogenic response.

No chemically related macroscopic or organ weight effects were observed in the groups of 10 male and 10 female rats and 10 female mice killed and examined at 15 months. Because of low survival, none of the male mice was killed for evaluation at 15 months. Microscopically, mild degeneration of the olfactory and respiratory epithelium was evident in toluene-exposed rats; in females, but not males, the severity and incidence of chronic inflammation of the nasal tissue was somewhat greater in both the 600-and 1,200-ppm groups. Necrosis and squamous metaplasia of the nasal cavity were seen in a few animals, as was hyperplasia of alveolar and bronchiolar epithelium.

For rats at the end of the 2-year exposure, no striking macroscopic alterations were observed at necropsy which were considered to be related to toluene exposure. Microscopically, nonneoplastic (toxic) effects likely due to toluene exposure were limited to mild responses in the nasal cavity and were similar to those observed in the 15-month studies. Concentration-dependent increases in the severity of nephropathy was seen in rats. Although Matsumoto et al. (1971) reported finding eosinophilic droplets in the kidney of male DONYRU rats after exposure to toluene, no evidence was found of an increase in hyaline droplets in the proximal tubules of the kidney of exposed male rats in either the 14- or 15-week gavage or inhalation studies of toluene. Other changes that usually accompany the increase in hyaline droplets (a_{2u}-globulin) in the "hydrocarbon nephropathy" syndrome, such as granular casts at the junction of the inner and outer stripe of the outer medulla in short-term studies and linear mineralization of the medulla in long-term studies, also were not found in the current NTP studies. These findings suggest that a_{2u}-globulin did not play a role in the exacerbation of spontaneous renal disease observed in the current NTP studies. The biologic significance of the eosinophilic droplets found by Matsumoto et al. (1971) is uncertain; the droplets could be albumin or other pigment but do not appear to be a_{2u}-globulin, as best as can be determined by an examination of Kodachrome slides of kidney sections from the Matsumoto studies. It is not certain how long after the last exposure the animals were killed, which is particularly important because the α_{2u}-globulin has a relatively rapid disappearance (72-96 hours) after exposure has been stopped. Further, earlier studies were conducted for 43 weeks (rats were approximately 1 year old when killed); synthesis of a_{2u}-globulin is known to decrease with age (Motwani et al., 1984) and seems to parallel the amount of cellular proliferation, being greater at 3 months of age and least at 12 months (Swenberg et al., 1989).

One possible neoplastic finding from these studies was that tubular cell adenomas of the kidney on routine (single) sectioning were found in one low dose and in two high dose male rats, and another low dose male rat had a transitional epithelial carcinoma; one high dose female rat had a renal tubular cell carcinoma, and another had a sarcoma of the kidney. Because these lesions were not accompanied by the occurrence of

tubular cell hyperplasia, because benzene (NTP, 1986a) and xylenes (NTP, 1986b) did not cause kidney neoplasms, although the kidney appears to be a target for other organic solvents, and, more important, because the finding of these few uncommon tumor types were possible signals for potential public health concern, the Program decided to evaluate this organ in extra detail. For each male rat, approximately six additional tissue sections were evaluated. The results of this supplementary evaluation revealed nine microscopic adenomas that were not discovered on routine sectioning--five in controls and four in the low dose group. All tubular cell neoplasms were benign. The combined diagnoses show that toluene did not cause any increases in hyperplasia, benign neoplasms, or malignant neoplasms of the kidney (see Table 11). Likewise, neither benzene nor xylenes were associated with kidney toxicity. The incidence of the five neoplasms (8.3%) found in the control male rats is the highest seen in the four NTP studies evaluated by step-sectioning of the kidney. For the three other studies, additional sections of the kidney of control male rats revealed either no increase (phenylbutazone, 0/50 to 0/50; NTP, 1989c) or an increase of two (furosemide, 1/50 to 3/50; NTP, 1989d) or three (nitrofurantoin, 0/50 to 3/50; NTP, 1989e). Thus, for the four separate control groups of male F344/N rats, the incidences of tubular cell adenomas of the kidney increased from 1/210 (0.5%) by routine sectioning to 11/210 (5.3%) by additional sectioning. This is similar to the experience of Kurokawa et al. (1983, 1986), who compared single sections vs. 15-20 sections of kidneys from F344 rats exposed to potassium bromate; in these studies, males showed an increase from one to three neoplasms in a group of 53 rats, whereas the number of neoplasms in females was zero before and after additional sections.

In contrast to the results reported by Maltoni et al. (1983, 1985) regarding increases in total malignant tumors in toluene-exposed Sprague Dawley rats (see Introduction), none of the study groups of F344/N rats or B6C3F₁ mice in the current studies showed a toluene-associated increase in the numbers of animals with benign or malignant tumors or total benign and/or malignant tumors. However, using the overall number of animals with tumors is not considered

to be appropriate for detecting potential carcinogenic effects of chemicals, except when observed in studies of less than 18 months (IARC, 1980, 1986; Huff et al., 1985; Haseman et al., 1986).

Both rats and mice exposed to benzene exhibited chemically related nonneoplastic and neoplastic effects of the hematopoietic system, Zymbal gland, forestomach, and adrenal gland. Further, neoplasms were induced in the oral cavity in rats and in the lung, liver, harderian gland, preputial gland, ovary, and mammary gland in mice (NTP, 1986a; Huff et al., 1988, 1989). In contrast to benzene (oral intubation at concentrations of 0 and 25-200 mg/kg), no significant changes in the incidences of neoplastic or nonneoplastic lesions in rats or mice were considered to be related to exposure to toluene (other than increased severity of nephropathy in rats) or xylenes (mixed) (NTP, 1986b) for 2 years. Exposure concentrations in the current toluene studies (0 and 120-1,200 ppm) were estimated to be considerably (up to 15 times) higher than those in the benzene studies. In addition, the gavage concentration of xylenes (mixed) (0 and 250-1,000 mg/kg) in the 2-year studies was from 2.5 (rats) to 10 (mice) times greater than those of benzene. Reviews of the xylene literature generally support the results of the NTP short-term studies (NIOSH, 1975; Miller et al., 1976; Mazella et al., 1978), but no reports on 2-year or lifetime studies were found. In two papers on benzene, Maltoni et al. (1983, 1985) reported some incomplete findings from long-term studies in which Sprague Dawley rats were given 500 mg/ kg xylenes in olive oil by gavage for 2 years and survivors were continued without exposure to week 141. They reported an increase in the total number of animals with malignant neoplasms in dosed vs. vehicle control males (14/40, 35% vs. 11/50) and females (22/40, 55% vs. 10/50). By comparison, after F344/N rats in the NTP studies were exposed to xylenes (mixed) for 104 weeks (NTP, 1986b), the total number of females with malignant neoplasms was not statistically increased at 500 mg/kg (16/50) compared with vehicle controls (12/50), and the total number of males with malignant neoplasms was actually decreased at 500 mg/kg (19/50) compared with vehicle controls (32/50), although this decrease was probably due to decreased survival in the dosed group. However, basing any conclusion on the overall proportion of animals with primary neoplasms (or with malignant neoplasms) is not the best approach for deciding potential carcinogenic effects of chemicals (IARC, 1980, 1986; Haseman et al., 1986).

One mechanism often proposed for observed differences in chemically induced toxic responses comes from metabolic studies whereby a chemical may modify its own toxicity or that of other chemicals by altering metabolism through induction of metabolizing enzymes. Pathiratne et al. (1986) investigated the effects of benzene, toluene, and xylenes on liver metabolism in male Sprague Dawley rats. Benzene was more potent at inducing conjugating systems, whereas xylenes were more potent at inducing cytochrome P450-dependent enzymes. Toluene was equipotent at inducing both enzyme types. The addition of methyl groups to the aromatic ring affects not only chemical-specific metabolism but also the inductive pattern of these monocyclic aromatic hydrocarbons. Thus, cytochrome P450 and related enzymes were induced to a greater degree with an increasing number of methyl groups, whereas the conjugating enzymes were affected in the opposite direction.

The key to the differences in responses between benzene and the methyl and dimethyl derivatives convincingly resides in the metabolism and the pattern of metabolic products. Benzene (NTP, 1986a; Huff et al., 1989) undergoes a complex constellation of multiple metabolic pathways leading to varied ring and ring-opened chemicals, some of which have been shown to induce cancer in animals when administered alone (catechol, Hirose et al., 1987, 1988; hydroguinone, NTP, 1989f; phenol, data considered negative or pehaps equivocal, NCI, 1980). To the contrary, toluene (and xylenes, NTP, 1986b) undergoes a simple Phase I oxidation to benzyl alcohol (no evidence of carcinogenicity; NTP, 1989a) on through benzaldehyde (NTP, 1989b) and benzoic acid (no evidence of carcinogenicity; personal communication from N. Ito, Nagoya City University, to J. Huff, NTP, 1989) and then to Phase II conjugation via mainly glycine (hippuric acid) and, to a lesser extent, by way of glucuronic acid (benzoyl glucuronide). Nearly 25% appears to be excreted unchanged, and this overall pattern pertains about equally whether the chemical is given orally or by inhalation (Pyykko et al., 1977). Small amounts of cresols are formed; these or hippurates in urine are used as biomonitors of human exposures. A postulated but never isolated metabolite is toluene oxide (Jerina et al., 1971). Ethylbenzene behaves similarly in that the major urinary metabolite is hippuric acid; following oxidation to benzoic, phenylacetic, and mandelic acids, these are excreted as glycine conjugates.

The results of the 2-year studies on benzene, toluene, and xylenes (mixed) show that methyl and dimethyl substitution on the benzene ring eliminates the carcinogenic influence of this molecule on rodents. Given the varying metabolic disposition patterns of these three congeners, these findings are perhaps not surprising.

Toluene, like benzene and its congeners and metabolites, does not appear to be a mutagen in vitro. The responses with toluene in microbial mutagenicity assays were uniformly negative, including several assays in which protocols specific for volatile chemicals were used. Results of the few in vitro assays conducted with mammalian cells for gene mutations or for cytogenetic damage were also negative. It is uncertain whether toluene, like benzene and its congeners and metabolites, is a clastogen in vivo. Single-exposure studies with toluene of specified purity were negative; mice or rats were exposed to toluene at doses up to 1,000 mg/kg and assayed for induction of chromosomal aberrations and micronucleated erythrocytes in bone marrow, and no chemical-related increases were observed (Kirkhart, 1980). On the other hand, Dobrokhotov (1972) and Lyapkalo (1973) both reported induction of chromosomal aberrations in bone marrow erythrocytes of rats exposed to toluene at 800-1,000 mg/kg per day for 12 days. Since the purity of the toluene was not specified, it is possible that the positive response could have come from benzene contamination. In some studies, benzene has been shown to be more effective when given repeatedly than when given once. Although toluene has not been tested for clastogenicity in somatic cells except in studies using a single exposure, it did not induce dominant lethal mutations in the germ cells of CD®-1 mice exposed by inhalation at concentrations up to 400 ppm for 6 hours per

IV. DISCUSSION AND CONCLUSIONS

day, 5 days per week, for 8 weeks (LBI, 1981). The key to the differences in the genetic toxicity between benzene and its methyl and dimethyl derivatives, like those in other toxicity responses, convincingly resides in the metabolism and the pattern of metabolic products.

The experimental and tabulated data for the NTP Technical Report on toluene were examined for accuracy, consistency, completeness, and compliance with Good Laboratory Practice regulations. As summarized in Appendix K, the audit revealed no major problems with the

conduct of the studies or with collection and documentation of the experimental data. No discrepancies were found that influenced the final interpretation of the results of these studies.

Under the conditions of these 2-year inhalation studies, there was no evidence of carcinogenic activity* for male or female F344/N rats exposed to toluene at concentrations of 600 or 1,200 ppm. There was no evidence of carcinogenic activity for male or female B6C3F₁ mice exposed by inhalation to toluene at concentrations of 120, 600, or 1,200 ppm for 2 years.

^{*}Explanation of Levels of Evidence of Carcinogenic Activity is on page 6. A summary of the Peer Review comments and the public discussion on this Technical Report appears on page 9.

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APPENDIX A

SUMMARY OF LESIONS IN MALE RATS IN THE TWO-YEAR INHALATION STUDY OF TOLUENE

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TABLE A1. SUMMARY OF THE INCIDENCE OF NEOPLASMS IN MALE RATS IN THE TWO-YEAR INHALATION STUDY OF TOLUENE

Chambe	er Control	600 p	pm	1,200	ppm
60		60		60	
60		60		60	
50		50		50	
	-				
(50)				(50)	
			(2%)	(40)	
	(0%)	* (50)			(90)
	(2%)	±(50)			(2%)
	(20%)	(50)			(4%)
	(270)	*(50)			(4270)
	(2%)	(00)		(40)	
	(2/0/	(50)		(49)	
,	(6%)		(6%)		(2%)
(49)		(50)	•	(49)	
	(4%)		(6%)	\/	
		•	•		
(50)	•	(50)		(49)	
,	(2%)	,- ,,		, , ,	
		2	(4%)		
(50)		(50)		(50)	
		1	(2%)		
4	(8%)	2	(4%)		
16	(32%)	25	(50%)		(38%)
					(2%)
* (50)				₹(50)	
			(2%)	(7.0)	
			/d 4 AV \		(OX)
2	(4%)	7	(14%)		(6%)
	(04)			1	(2%)
	(2%)	(50)		(EO)	
(507			(90%)	(50)	
(EO)			(270)	(49)	
	(6%)		(14%)		(2%)
J	(0%)	•	(1470)		(2%)
(50)		(50)			(= ,0)
	(6%)	,	(8%)	,	(6%)
					
(50)		(50)		(50)	
	(2%)	(00)		(00)	
	*	17	(34%)	7	(14%)
(50)		(50)		(50)	
			(2%)		
					(2%)
(50)		(50)		(50)	
	(28%)		(32%)		(20%)
(49)		(50)		(50)	
11	(22%)			10	(20%)
				_	/4 Car :
7	(14%)			6	(12%)
		1	(2%)		
	60 60 50 (50) (50) 1 (50) 3 (49) 2 1 (50) 4 16 *(50) (50) (50) (50) 3 (50) ((50) (50) (50) (1 (2%) (50) (1 (2%) (49) (1 (2%) (50) (3 (6%) (49) (2 (4%) (1 (2%) (50) (4 (8%) (50) (4 (8%) (50) (50) (50) (2 (4%) (50) (50) (50) (50) (50) (50) (50) (50	(50) (50) (50) (50) (50) (50) (50) (50)	(50) (50) (50) (50) (50) (50) (50) (50)	60 60 60 60 60 60 60 50 50 50 50 (50) (50) (50) (50) (50) 1 (2%) (49) 1 (2%) (50) (49) 1 (2%) (50) (49) 1 (2%) (50) (49) 1 (2%) (50) (49) 1 (2%) (50) (49) 2 (4%) 3 (6%) 1 1 (2%) (50) (49) 2 (4%) 3 (6%) (49) 1 (2%) (50) (50) (49) 2 (4%) 3 (6%) (50) (49) 1 (2%) (50) (50) (49) 1 (2%) (50) (50) (50) (50) 1 (2%) (50) (50) (50) 4 (8%) 2 (4%) (50) (50) 4 (8%) 2 (4%) (50) (50) 1 (2%) (50) (50) (50) 2 (4%) 7 (14%) 3 1 (2%) (50) (50) (50) 3 (6%) 7 (14%) 1 (50) (50) (50) (50) (50) 3 (6%) 7 (14%) 1 (50) (50) (50) (50) (49) 3 (6%) 7 (14%) 1 (50) (50) (50) (50) (49) 3 (6%) 7 (14%) 7 (14%) 7 (14%) 7

TABLE A1. SUMMARY OF THE INCIDENCE OF NEOPLASMS IN MALE RATS IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

	Chamb	er Control	600 I	pm	1,200	ppm
ENDOCRINE SYSTEM (Continued)				·······	" H	
Islets, pancreatic	(50)		(50)		(50)	
Adenoma		(4%)		(4%)		(4%)
Carcinoma	_	(2%)	_	(- / - /		(2%)
Pituitary gland	(49)		(50)		(50)	(= ,,,
Pars distalis, adenoma		(45%)		(48%)		(36%)
Pars distalis, carcinoma				(10,11)		(2%)
Pars distalis, leukemia monocytic					_	(2%)
Pars distalis, leukemia mononuclear	3	(6%)	5	(10%)		(6%)
Pars intermedia, leukemia mononuclear				(2%)	•	(,
Pars nervosa, leukemia mononuclear				(2%)		
Thyroid gland	(50)		(50)	, ,	(50)	
Leukemia mononuclear		(2%)	• /	(4%)	(00)	
C-cell, adenoma		(10%)		(14%)	8	(16%)
C-cell, carcinoma	•	(== ,= ,		(2%)	ŭ	(=0,0)
Follicle, adenoma	1	(2%)		(4%)		
GENERAL BODY SYSTEM None						
GENITAL SYSTEM						
Preputial gland	(50)		(50)		(50)	
Adenoma			1	(2%)		
Carcinoma	1	(2%)				
Leukemia mononuclear	2	(4%)	3	(6%)		
Prostate	(50)		(50)		(50)	
Leukemia mononuclear	(00)		, /	(6%)	(00)	
Testes	(50)		(50)		(50)	
Leukemia mononuclear		(2%)		(4%)	,,,,,	
Bilateral, interstitial cell, adenoma		(46%)		(54%)	30	(60%)
Interstitial cell, adenoma		(26%)		(18%)		(20%)
Serosa, mesothelioma malignant		(2%)				(2%)
HEMATOPOIETIC SYSTEM	·-··	***************************************	****			
Blood	*(50)		*(50)		*(50)	
Leukemia mononuclear	(00)		, ,	(2%)		(4%)
Bone marrow	(50)		(50)	(/ - /	(50)	. = ,0,
Leukemia mononuclear		(20%)		(30%)		(24%)
Lymph node	(50)	(30.0)	(50)	(30,0)	(50)	.= = 10)
Carcinoma adenosquamous, metastatic, lung		(2%)	(55)		(55)	
Pibrosarcoma, metastatic, skin	•	/	1	(2%)		
Leukemia mononuclear	1	(2%)	•	, , , , ,	2	(4%)
Iliac, leukemia mononuclear	•	,	1	(2%)	-	,
Mediastinal, leukemia mononuclear	4	(8%)		(10%)	1	(2%)
Mesenteric, leukemia mononuclear		(28%)		(40%)		(34%)
Mesenteric, lymphoma malignant lymphocyti		(2%)	_0		- '	
Renal, carcinoma, metastatic, kidney			1	(2%)		
Renal, leukemia mononuclear	2	(4%)	-	,		
Lymph node, mandibular	(36)	•	(45)		(37)	
Leukemia mononuclear		(6%)		(24%)		(11%)
Spleen	(50)	•	(50)		(50)	
Hemangiosarcoma	/			(2%)	1/	
Leukemia mononuclear	17	(34%)		(50%)	19	(38%)
Lymphoma malignant	•		•	•		(2%)
Capsule, mesothelioma malignant						(2%)

TABLE A1. SUMMARY OF THE INCIDENCE OF NEOPLASMS IN MALE RATS IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

	Chambe	er Control	600 p	pm	1,200	ppm
HEMATOPOIETIC SYSTEM (Continued)						 -
Thymus	(44)		(46)		(46)	
Carcinoma, metastatic, kidney	(==)			(2%)	(40)	
Carcinoma adenosquamous, metastatic, l	ung 1	(2%)				
Leukemia mononuclear		(14%)	9	(20%)	4	(9%)
Osteosarcoma, metastatic, uncertain prin	nary site				1	(2%)
NTEGUMENTARY SYSTEM		•==				
Mammary gland	(50)		(48)		(49)	
Fibroadenoma	1	(2%)				
Skin	(50)		(50)		(50)	
Histiocytic sarcoma, metastatic				(2%)		
Keratoacanthoma			1	(2%)		
Subcutaneous tissue, fibroma	2	(4%)				(2%)
Subcutaneous tissue, fibrosarcoma			2	(4%)		(2%)
Tail, papilloma squamous					1	(2%)
Thoracic, subcutaneous tissue, fibroma	1	(2%)				
MUSCULOSKELETAL SYSTEM						
Skeletal muscle	*(50)		*(50)		*(50)	
Fibrosarcoma					1	(2%)
NERVOUS SYSTEM					·	
Brain	(50)		(50)		(50)	
Leukemia mononuclear	3	(6%)	7	(14%)	3	(6%)
Cerebrum, astrocytoma malignant				(2%)		
RESPIRATORY SYSTEM				· · · · · · · · · · · · · · · · · · ·		
Lung	(50)		(50)		(50)	
Alveolar/bronchiolar carcinoma				(2%)		
Carcinoma, metastatic, kidney			1	(2%)		
Carcinoma adenosquamous	1	(2%)			1	(2%)
Fibrosarcoma, metastatic, skin			1	(2%)		
Fibrous histiocytoma, metastatic, skin					1	(2%)
Histiocytic sarcoma, metastatic			1	(2%)		
Leukemia monocytic		(2%)				
Leukemia mononuclear		(32%)	22	(44%)		(28%)
Osteosarcoma, metastatic, uncertain prin	-		/= A ·		_	(2%)
Trachea	(50)	(04)	(50)		(50)	
Carcinoma adenosquamous, metastatic, l		(2%)	_	(100)		
Leukemia mononuclear	3	(6%)	5	(10%)		
SPECIAL SENSES SYSTEM	_					
Zymbal gland	*(50)		*(50)		*(50)	
Carcinoma	1	(2%)	2	(4%)	1	(2%)
JRINARY SYSTEM						
Kidney	(50)		(50)		(50)	
Histiocytic sarcoma, metastatic				(2%)		
Leukemia mononuclear	3	(6%)		(20%)	4	(8%)
Capsule, fibrosarcoma, metastatic, skin			1	(2%)		
Capsule, mesothelioma malignant						(2%)
Renal tubule, adenoma Transitional epithelium, carcinoma				(2%)	2	(4%)
Observation at a sittle discussion and a second			1	(2%)		

TABLE A1. SUMMARY OF THE INCIDENCE OF NEOPLASMS IN MALE RATS IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

	Chambe	er Control	600 p	ppm	1,200	ppm
URINARY SYSTEM (Continued)						
Urinary bladder	(50)		(50)	(04)	(50)	
Fibrosarcoma, metastatic, skin	_	(100)		(2%)		
Leukemia mononuclear		(10%)	10	(20%)	3	(6%)
Sarcoma	1	(2%)				رم <i>ح</i> ،
Serosa, mesothelioma malignant Transitional epithelium, papilloma						(2%) (2%)
Transitional epitherium, papinoma						(270)
SYSTEMIC LESIONS						
Multiple organs	*(50)		*(50)		*(50)	
Leukemia mononuclear	17	(34%)	26	(52%)	19	(38%)
Lymphoma malignant lymphocytic		(2%)				
Leukemia monocytic		(2%)				(2%)
Mesothelioma malignant	1	(2%)			1	(2%)
Hemangiosarcoma			1	(2%)		
Lymphoma malignant					1	(2%)
ANIMAL DISPOSITION SUMMARY						
Animals initially in study	60		60		60	
Interval sacrifice	10		10		10	
Terminal sacrifice	29		27		22	
Moribund	14		11		23	
Dead	6		9		5	
Accident	1					
Natural death			3			
TUMOR SUMMARY						. -
Total animals with primary neoplasms **	49		48		49	
Total primary neoplasms	108		122		107	
Total animals with benign neoplasms	45		46		47	
Total benign neoplasms	75		77		73	
Total animals with malignant neoplasms	23		31		26	
Total malignant neoplasms	26		37		28	
Total animals with secondary neoplasms ***	ī		3		2	
Total secondary neoplasms	4		12		3	
Total animals with malignant neoplasms					•	
uncertain primary site					1	
Total animals with neoplasms						
uncertain benign or malignant	7		8		6	
Total uncertain neoplasms	7		8		6	

^{*} Number of animals receiving complete necropsy examinations; all gross lesions including masses examined microscopically.

** Primary tumors: all tumors except secondary tumors

*** Secondary tumors: metastatic tumors or tumors invasive into an adjacent organ

TABLE A2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE RATS IN THE TWO-YEAR INHALATION STUDY OF TOLUENE: CHAMBER CONTROL

INHALATI	U-1 K			O	•				E:	01		VID.	EK		UN	TK	O.	4							
WEEKS ON STUDY	0 4 9	0 6 0	0 6 8	0 6 8	0 8 5	0 8 6	0 8 8	0 8 9	0 8 9	0 9 0	0 9 0	0 9 1	0 9 3	0 9 4	0 9 5	0 9 6	0 9 7	0 9 7	0 9 7	1 0 1	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5
CARCASS ID	1 4 1	5 3 1	3 1 1	2 4 1	1 6 1	8	3 2 1	7 1	4 2 1	0	3 7 1	2 7 1	1 0 1	7 1	3 3 1	3 9 1	3 5 1	9	5 2 1	1 3 1	0 3 1	2 5 1	0	5 1 1	5 7 1
ALIMENTARY SYSTEM	·												_				_								
Esophagus Intestine large	+	+	+	+	+	+	+	++	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine large, cecum	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Leukemia mononuclear Intestine large, colon	+	X +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Leukemia mononuclear Intestine large, rectum	М	X	_	_	_	4	_	_	_	_	4	_	_	4	_	_	_	_	_	_	_	_	_	_	+
Leukemia mononuclear		X	Ċ		,		,	,	,			,	,	•	,	,	7	•	,	7	'	,	,		•
Intestine small Intestine small, duodenum	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Leukemia mononuclear				Ċ			Ċ										Ċ	X		Ċ				Ċ	
Intestine small, ileum Leukemia mononuclear	1	+	+	+	+	+	+	+	+	+	+	+	+	M	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant lymphocytic Intestine small, jejunum Adenocarcinoma, mucinous	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Leukemia mononuclear	1.																								
Liver Hepatocellular adenoma	+	+	+	+	+	+	+	+	+	+	+	+	+	*X	+	+	+	+	+	+	+	+	+	+	+
Leukemia mononuclear Mesentery		X				X	X	X		X	X	X	X					X			X				X
Pancreas	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Leukemia mononuclear Acinus, adenoma		X					X												x						
Salivary glands	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Stomach Stomach	+	+	+	+	+	++	+	+	+	++	+	++	++	+	+	+	+	+	+	+	+	+	+	+	++
Leukemia mononuclear	+	X			·		X		·			Ċ	X		Ċ					Ċ			·		
Stomach, glandular Leukemia mononuclear	+	*	+	+	+	+	*	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
CARDIOVASCULAR SYSTEM	-																								
Blood vessel Heart	‡	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Carcinoma adenosquamous, metastatic,	"		_	_	_	Τ.	_	_	+	_	_	_	_	т	_	Τ.	_	_	_	_	Ŧ	_	Τ.	Ŧ	
lung Leukemia mononuclear		X	X			x	x					x	х					х							
ENDOCRINE SYSTEM	·																								
Adrenal gland	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Adrenal gland, cortex Leukemia mononuclear	+	*X	+	+	+	*X	*X	*	+	+	+	*X	X	+	+	+	+	+ X	+	+	*	+	+	+	*
Adrenal gland, medulla Leukemia mononuclear	+	*X	+	+	+	*X	*X	*	+	+	+	x X	* X + X	+	+	+	+	*	+	+	+	+	+	+	+
Pheochromocytoma, NOS	1	Λ.						Х				n.	^					Ŷ							
Islets, pancreatic Adenoma	+	+	+	+	+	+	+	*	+	+	+	+	+	+	+	+	+	+	+	+	+	+	*	+	+
Carcinoma	1.,	.,	.,													.,									
	M	M	M	+	+	M		+	+	+	+	+	M	+	+ X	M + X	M + X	+	+	+	+	+	+	+	+
Parathyroid gland Pituitary gland	+	+	+	+	+	+	+	+	+	+	+	+										Y			
Pituitary gland Pars distalis, adenoma		+	+	*	*	+	x	*	*	+	+	*X	141	*	X	X	X	¥	X			Α.			
Pituitary gland Pars distalis, adenoma Pars distalis, leukemia mononuclear Thyroid gland		+	+	* *	* +	+	x +	* +	* +	+	+	* +	+	X +	X +	X +	X +	X +	+	+	+	+	+	+	+
Pituitary gland Pars distalis, adenoma Pars distalis, leukemia mononuclear Thyroid gland Leukemia mononuclear C-cell, adenoma	+		+	* *	+	+	* *	* *	* +	+	+	* +	+	X +	X +	X +	X +	X +	+	+	+	+	+	+	+
Pitutary gland Pars distalis, adenoma Pars distalis, leukemia mononuclear Thyroid gland	+	+	+	* +	* + X	+	* +	* +	* +	+	+	* +	+	X +	X +	X +	X +	X +	+	+	+	+	+	+	+
Pituitary gland Pars distalis, adenoma Pars distalis, leukemia mononuclear Thyroid gland Leukemia mononuclear C-cell, adenoma	+	+	+	* +	+	+	* +	* +	* +	+	+	* +	+	X +	* +	* +	* +	X +	+	+	+	+	+	+	+
Pituitary gland Pars distalis, adenoma Pars distalis, leukemia mononuclear Thyroid gland Leukemia mononuclear C-cell, adenoma Follicle, adenoma GENERAL BODY SYSTEM None GENITAL SYSTEM	+	+	+	* +	+	+	* +	* +	* +	+	+	* +	+	X +	* +	* +	* +	X +	+	+	+	+	+	+	+
Pituitary gland Pars distalis, adenoma Pars distalis, leukemia mononuclear Thyroid gland Leukemia mononuclear C-cell, adenoma Follicle, adenoma GENERAL BODY SYSTEM None GENITAL SYSTEM Epididymis	+	+	+ + + + + +	* + +	+	+ + + +	* + + + + + + + + + + + + + + + + + + +	* + + + + + + + + + + + + + + + + + + +	* + + + + + + + + + + + + + + + + + + +	+ +	+ + + + + + + + + + + + + + + + + + + +	* + + + + + + + + + + + + + + + + + + +	+ + + +	* + + + + + + + + + + + + + + + + + + +	* + +	+ + +	* + +	+ +	* + + + + + + + + + + + + + + + + + + +	+	+ + + + +	++	+ + +	+ + + +	+
Pituitary gland Pars distalis, adenoma Pars distalis, leukemia mononuclear Thyroid gland Leukemia mononuclear C-cell, adenoma Follicle, adenoma GENERAL BODY SYSTEM None GENITAL SYSTEM Epididymis Preputial gland Carcinoma	+	+	+ + +	* + +	+	+ + +	* + + + + + + + + + + + + + + + + + + +	* + + +	* + + + + + + + + + + + + + + + + + + +	+ + + + + +	+ + + + + + + + + + + + + + + + + + + +	* + + + + + + + + + + + + + + + + + + +	+ + +	* + + + + + + + + + + + + + + + + + + +	+ + +	+ + +	+ + +	* + + + + + + + + + + + + + + + + + + +	++	+++	+ + +	++	+	+ + +	+ + + +
Pituitary gland Pars distalis, adenoma Pars distalis, leukemia mononuclear Thyroid gland Leukemia mononuclear C-cell, adenoma Follicle, adenoma GENERAL BODY SYSTEM None GENITAL SYSTEM Epididymis Preputial gland Carcinoma Leukemia mononuclear	+	* * * * * * * * * * * * * * * * * * *	+ + + +	* + + +	+	+ + + + +	* + + + + + + + + + + + + + + + + + + +	* + + + + + + + + + + + + + + + + + + +	* + + + + + + + + + + + + + + + + + + +	+ + + + + +	+ + + + + +	* + + + + + + + + + + + + + + + + + + +	+ + + +	* + + + + + + + + + + + + + + + + + + +	+ + + +	+ + + +	+ + + +	* + + + + + + + + + + + + + + + + + + +	++++++	+++++	+ + + +	+++++	+	++++++	+ + + +
Pituitary gland Pars distalis, adenoma Pars distalis, leukemia mononuclear Thyroid gland Leukemia mononuclear C-cell, adenoma Folicle, adenoma GENERAL BODY SYSTEM None GENITAL SYSTEM Epididymis Preputial gland Carcinoma Leukemia mononuclear Prostate Testas	+	* * * * * * * * * * * * * * * * * * *	+ + + + + + + + + + + + + + + + + + + +	+ + + + +	+	+ + + + + + + + + + + + + + + + + + + +	* + + + + + + + + + + + + + + + + + + +	+ + + + +	+ + + + +	+ + + + + + + + + + + + + + + + + + + +	+ + + + + + + + + + + + + + + + + + + +	+ + + + +	+ + + + + + + + + + + + + + + + + + + +	+ + + +	+ + + +	+ + + +	+ + + +	+ + + +	* + + + + + + + + + + + + + + + + + + +	+++++	+ + + +	++++	+	++++	+ + + +
Pituitary gland Pars distalis, adenoma Pars distalis, leukemia mononuclear Thyroid gland Leukemia mononuclear C-cell, adenoma Follicle, adenoma GENERAL BODY SYSTEM None GENITAL SYSTEM Epididymis Preputial gland Carcinoma Leukemia mononuclear Prostate	+ + + + + +	+	+ + + + + + + + + + + + + + + + + + + +	+ + + + +	+	+ + + + + + + + + + + + + + + + + + + +	* * * + + + + * * * * * * * * * * * * *	+ + + + + + + + + + + + + + + + + + + +	+ + + + +	+ + + + + + X	+ + + + + + + + + + + + + + + + + + + +	+ + + + +	+ + + + + + X	* + + + + + + + + + + + + + + + + + + +	+ + + *	+ + + + x	+ + + + +	+ + + + x	+++++	+ + + *	+ + + X	+ + + X	+ + + + x	+ + + + x	+ + + + x

^{+:} Tissue examined microscopically
: Not examined
-: Present but not examined microscopically
I: Insufficient tissue

M: Missing
 A: Autolysis precludes examination
 X: Incidence of listed morphology

TABLE A2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE RATS: CHAMBER CONTROL (Continued)

WEEKS ON	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1
STUDY	5	0 5	0 5	0 5	0 5	0 5	0 5	0 5	0 5	0 5	0 5	0 5	0 5	0 5	0 5	1										
CARCASS	-6	0	0	-0	2	4	٠,	,	_	_	4	5	5	0	0	2	3	3	3	0	0	1	1	-2	4	TOTAL
ID	0	2	4	6	1	6	9	2	5	9	3	6	8	7	8	9 1	0	4	8	5	9	1	8	3	4	TUMOR
Y Y & F D & W Y D	1		<u>.</u>	1					1				<u> </u>				<u>.</u>								<u>.</u>	
LIMENTARY SYSTEM	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
ntestine large	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 50
ntestine large, cecum Leukemia mononuclear	"	7	_	_	•		~	*		т	т	•	Τ.	т	•		•	*	_	-	т		,	т	7	1
ntestine large, colon	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Leukemia mononuclear itestine large, rectum	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	49
Leukemia mononuclear ntestine small	+	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	50
ntestine small, duodenum	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	÷	+	+	÷	+	+	+	÷	+	+	50
Leukemia mononuclear ntestine small, ileum	١.											X +	_						_	_		_	_	X +	+	3 49
Leukema mononuclear	-		+		~	_	+	_	_	-	_	x	Τ.	т	т	_	•	_	т.		*	7	Τ.	X	-	2
Lymphoma malignant lymphocytic	Ι.							X																	+	1 50
ntestine small, jejunum Adenocarcinoma, mucinous	+	+	+	+	+	*	+	+	+	+	+	+	_	+	_	+	+	_	_	_	+	x	_	_	Τ.	1
Leukemia mononuclear												X												X		2
aver Hepatocellular adenoma	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	X	+	+	X	+	+	+	X	50 4
Leukemia mononuclear	1						X					X	X					••	X					X		16
lesentery Pancreas	1 +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Leukemia mononuclear	1.	•	•	Ċ		·	•	·									•	•			,					50 2
Acınus, adenoma alıvary glands	1	+	+	+	+	+	_	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	1 50
tomach	+	÷	÷	+	÷	+	÷	÷	÷	÷	+	+	+	+	÷	+	+	÷	+	÷	+	+	+	+	+	50
tomach, forestomach Leukemia mononuclear	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 3
tomach, glandular	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Leukemia mononuclear																								X		3
CARDIOVASCULAR SYSTEM				_		_				_			_													·
Blood vessel Heart	1 +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 50
Carcinoma adenosquamous, metastatic,	'		'	'	•	•			•	,	'	•	·	•			,	•	,	Ċ	•		,	·		
lung Leukemia mononuclear												X												x	x	9
ENDOCRINE SYSTEM																										
Adrenal gland	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
idrenal gland, cortex Leukemia mononuclear	+	+	+	+	+	+	X +	+	+	+	+	X	+	+	+	+	+	+	*	+	+	+	+	+ X	*	50 14
Adrenal gland, medulla	+	M	+	+	+	+	+	+	+	+	+	X	+	+	+	+	+	+	+	+	+	+	+	+	+	49
Leukemia mononuclear Pheochromocytoma, NOS							Х					X	x	x					X		х			X		11 7
slets, pancreatic	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Adenoma Carcinoma	1																								x	2
arathyroid gland	+	+	+	+	+	+	+	+	M	M	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	42
htuitary gland Pars distalis, adenoma	+	+	+	+	+	+	+	+	+	*	+	+	*	*	*	*	*	+	*	+	+	+	+	+	+	49 22
Pars distalis, leukemia mononuclear					^		Λ.		Λ.	^		X	Α.	Λ	Λ	Λ	Λ.		А					X		3
hyroid gland	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Leukemia mononuclear C-cell, adenoma						x				X										X		X				5
Follicle, adenoma	-																								X	1
ENERAL BODY SYSTEM None																										-
ENITAL SYSTEM		—																								-
pididymis	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
	1 +	+	+	+	+	+	+	+	+	+	+	+	*	+	+	+	+	+	+	+	+	+	+	+	+	50
reputial gland	1											w	4.											x		2
reputial gland Carcinoma Leukemia mononuclear												Х														
Preputial gland Carcinoma Leukemia mononuclear Prostate	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	++	+	50
Preputial gland Carcinoma Leukemia mononuclear Prostate Testes Leuke <i>mia mononuc</i> lear	++	++	+	+	+	++	+	+	+	+	+	++	+	++	++	+	+	+	+	+	+	++	+	+	++	50 50 1
Preputial gland Carcinoma Leukemia mononuclear Prostate Testes	+ + x	* + *	+	+ + X	+ + X	+	+ + X	+	+ + X	50 50																

TABLE A2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE RATS: CHAMBER CONTROL (Continued)

					``		****	ucu	,																
WEEKS ON STUDY	0 4 9	0 6 0	0 6 8	0 6 8	0 8 5	0 8 6	0 8 8	0 8 9	0 8 9	0 9 0	0 9 0	9 1	0 9 3	0 9 4	0 9 5	0 9 6	9 7	0 9 7	0 9 7	1 0 1	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5
CARCASS ID	1 4 1	5 3 1	3 1 1	2 4 1	1 6 1	4 8 1	3 2 1	7 1	4 2 1	2 0 1	3 7 1	7 1	1 0 1	1 7 1	3 3 1	3 9 1	3 5 1	4 9 1	5 2 1	1 3 1	0 3 1	2 5 1	4 0 1	5 1 1	5 7 1
HEMATOPOIETIC SYSTEM Blood Bone marrow Leukemia mononuclear Lymph node Carcinoma adenosquamous, metastatic, lung	+++++	* X +	+ + X	+	+	+ X +	* X +	+	+	+	* X +	* X +	+ X +	+	+	+	+	* X +	+	+	+	+	+++	+	++
Leukemia mononuclear Mediastinal, leukemia mononuclear Mesenteric, leukemia mononuclear Mesenteric, lymphoma malignant lymphocytic		x				X X		x			X X	x	X					x			x				x
Renal, leukemia mononuclear Lymph node, mandibular Leukemia mononuclear	М	X M	M	M	M	M	+	M	M	M	M	M	M	M	+	+	+	+ X	+	+	+	+	+	+	+
Spieen Leukemia mononuclear Thymus Carcinoma adenosquamous, metastatic, lung Leukemia mononuclear	+	* X + X	+ + X	+	+	+ X +	* X +	* *	+	* X +	* X +	* X + X	* * *	+ M	+	+ M	+ M	* *	+ M	+	* *	+	+	+	X +
INTEGUMENTARY SYSTEM Mammary gland	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Fibroadenoma Skin Subcutaneous tissue, fibroma Thoracic, subcutaneous tissue, fibroma	+	+	+	+	+	+	+	+	+	*	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
MUSCULOSKELETAL SYSTEM Bone	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
NERVOUS SYSTEM Brain Leukemia mononuclear	+	*	+	+	+	+	* X	+	+	+	+	+	*	+	+	+	+	+	+	+	+	+	+	+	+
RESPIRATORY SYSTEM Lung Carcinoma adenosquamous	+	+	+ X	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Leukemia monocytic Leukemia mononuclear Nose Trachea Carcinoma adenosquamous, metastatic, lung Leukemia mononuclear	+++	x + +	+ + X	+	++	X + +	X + +	X + +	+	X + +	X + +	X + +	X + +	++	++	++	++	X + +	+ +	++	X + +	++	+	++	X + +
SPECIAL SENSES SYSTEM Eye Zymbal gland Carcinoma																		+		+ X					
URINARY SYSTEM Kidney Leukemia mononuclear Unnary bladder Leukemia mononuclear Sarcoma	+ +	* X * X	+	+	+	+ +	+ *	+	+	+	+	+	+ X +	+	+	+	+	+ *	+	+	+	+	+	+	+

TABLE A2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE RATS: CHAMBER CONTROL (Continued)

WEEKS ON STUDY	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	TOTAL.						
CARCASS ID	6 0 1	0 2 1	0 4 1	0 6 1	2 1 1	4 6 1	5 9 1	1 2 1	1 5 1	1 9 1	4 3 1	5 6 1	5 8 1	7 1	0 8 1	2 9 1	3 0 1	3 4 1	3 8 1	0 5 1	9 1	1 1 1	1 8 1	2 3 1	4	TISSUES TUMORS
HEMATOPOIETIC SYSTEM Blood Bone marrow Leukemia mononuclear Lymph node	+	+	+	+	+ +	+	+	+	+	+ +	+	* X +	* X +	+	++++	+ +	+ +	+	+ +	+ + +	+	+	++	+ X +	+ +	4 50 10 50
Carcinoma adenosquamous, metastatic, lung Leukemia mononuclear Mediastinal, leukemia mononuclear Mesenteric, leukemia mononuclear Mesenteric, lymphoma malignant							x					x	x											X X X	x	1 1 4 14
lymphocytic Renal, leukemia mononuclear Lymph node, mandibular Leukemia mononuclear Spleen Leukemia mononuclear	+	+	+	+	+	+	+ *	+ +	+	+	+	+ *	+ *	+	+	+	+	M +	+ *	+	++	+	+	X + X + X	+ + X	1 2 36 2 50 17
Thymus Carcinoma adenosquamous, metastatic, lung Leukemia mononuclear	+	+	M	М	+	+	+	+	+	+	+	x	+	+	+	+	+	+	+	+	+	+	+	×	+	1 6
INTEGUMENTARY SYSTEM Mammary gland Fibroadenoma Skin Subcutaneous tissue, fibroma Thoracic, subcutaneous tissue, fibroma	+	+	+	+	+	+	+	+	+	+	+	+	+ *	+	+	+ + X	+	+	+	* X +	+	+	+	+	+	50 1 50 2 1
MUSCULOSKELETAL SYSTEM Bone	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
NERVOUS SYSTEM Brain Leukemia mononuclear	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
RESPIRATORY SYSTEM Lung Carcinoma adenosquamous Leukemia monocytic	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+ X	+	+	+	+	+	+	50 1 1
Leukema monocytic Leukema mononuclear Nose Trachea Carcinoma adenosquamous, metastatic,	++	++	++	+	++	++	X + +	+	+	+	++	X + +	* + +	+	++	++	++	+	++	++	+	+	++	* + +	X + +	16 50 50
lung Leukemia mononuclear	Ì																							x		3
SPECIAL SENSES SYSTEM Eye Zymbal gland Carcinoma																						+				2 1 1
URINARY SYSTEM Kidney Leukemia mononuclear Urinary bladder Leukemia mononuclear Sarcoma	+	+	+	+	+	+	+	+	+	+	+ + x	+ *	+	+	+	+	+	+	+	+	+	+	+	+ X + X	+	50 3 50 5 1

TABLE A2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE RATS IN THE TWO-YEAR INHALATION STUDY OF TOLUENE: 600 ppm

	IALA			31		<i>,</i> 1	UF		UL					рp											
WEEKS ON STUDY	0 0 5	4	0 6 9	0 8 2	0 8 2	0 8 3	8 8	0 8 9	9	0 9 0	0 9 6	9 8	9	9	9	1 0 0	1 0 0	1 0 0	1 0 2	1 0 2	1 0 2	1 0 4	1 0 5	1 0 5	1 0 5
CARCASS	6	1 2	3	4	3	4	7	3	3	1 2	4	1 2	7	1 2	1 2	4	5	5	4	4	7	7	3	1 4	5
ID	9	2 1	3	1	9	3	3 1	6	5 1	3 1	7 1	1 1	0	9 1	5 1	0 1	3	5 1	4 1	5 1	1	9	1	6 1	1
ALIMENTARY SYSTEM Esophagus	- -			+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Leukemia mononuclear	1:	Ċ					X	Ċ								Ċ		·	Ċ				+		+
Intestine large Intestine large, cecum	‡	+	+	+	+	+	+	+	Ŧ	+	+	+	Ŧ	Ŧ	+	+	Ŧ	+	Ŧ	+	+	+	Ŧ	Ŧ	+
Intestine large, colon Intestine large, rectum	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine small Intestine small, duodenum	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	++	+	+	+	+
Leukemia mononuclear				Ċ			i			i	·	·	X	·	×				_	i	i	X	+		+
Intestine small, ileum Leukemia mononuclear	†	+	+	+	+	+	+	+	X	+	+	+	+	+	+	+	_	_	_	7	_	X	т.	_	Τ.
Intestine small, jejunum Leukemia mononuclear	+	+	+	+	+	+	+	+	+	+	+	+	+	+	*	+	+	+	+	+	+	×	+	+	+
Liver Hepatocellular carcinoma	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Hepatocellular adenoma																									
Histiocytic sarcoma, metastatic Leukemia mononuclear			X		x		x	X	X		X	X	X		X	X	X	X	X			X	X		
Mesentery _ Leukemia mononuclear	+						*																		
Pancreas	+	+	+	+	+	+	+	+	<u>+</u>	+	+	+	±	+	+	+	+	+	+	+	+	+	+	+	+
Leukemia mononuclear Salivary glands	+	+	+	+	+	+	X	+	X +	+	X +	+	X +	+	X +	+	+	+	+	+	+	X +	X +	+	+
Leukemia mononuclear Stomach	1.	_	_	_	_	_	X	_	_	_	_	_	_	_	_	_	_	+	+	4	+	_	+	+	+
Stomach, forestomach	+	+	+	+	+	+	+	+	+	+	+	+	+	+	±	+	÷	+	+	+	+	÷	+	÷	÷
Leukemia mononuclear Stomach, glandular	+	+	+	+	X	+	X	+	+	+	X +	+	+	+	X +	+	+	+	+	+	+	X +	+	+	+
Leukemia mononuclear							X				X				X							X			
CARDIOVASCULAR SYSTEM	-		-																						
Blood vessel Heart	†	++	+	++	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Leukemia mononuclear			X				X	X	X		X				X		X	X	X			X	X		
ENDOCRINE SYSTEM	-																								
Adrenal gland Capsule, fibrosarcoma, metastatic, skin	†	+	+	+	+	+	+	+	+	X	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Adrenal gland, cortex Adenoma	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	*	+	+
Leukemia mononuclear	1.		X				X		X + X		X + X		X + X		X + X		X	X	X		_	X +	X +	+	+
Adrenal gland, medulla Leukemia mononuclear	†	+	X	+	+	+	x	+	X	+	X	+	X	+	X	+	*	*	*	_		X	X	т	Τ.
Pheochromocytoma malignant Pheochromocytoma, NOS											X					x			X			X	X		
Bilateral, pheochromocytoma, NOS Islets, pancreatic			4	_	_	_	_	_	_	_		_	_	_	_		X	_	_	_	_	_	_	+	+
Adenoma				т		_			· ·	•	•			•	7								Ċ	X	
Parathyroid gland Pituitary gland	+	+	+	+	+	+	M +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Pars distalis, adenoma Pars distalis, leukemia mononuclear					X	X	X	X	X		*	*	*	X	¥				X	X		x	X		X
Pars intermedia, leukemia mononuclear							•								X							••			
Pars nervosa, leukemia mononuclear Thyroid gland	+	+	+	+	+	+	+	+	+	+	+	+	+	+	X +	+	+	+	+	+	+	+	+	+	+
Leukemia mononuclear C-cell, adenoma							X						х		X		х								
C-cell, carcinoma													••				••			X					
Follicle, adenoma	_ _																								
GENERAL BODY SYSTEM None																									
GENITAL SYSTEM	- -																								
Epididymis Preputial gland	†	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Adenoma Leukemia mononuclear			Y												¥							x			
Prostate	+	+	+	+	+	+	+	+	+	+	+	+	+	+	х + х	+	+	+	+	+	+	+	+	+	+
Leukemia mononuclear Seminal vesicle		+		+					X						Х										
Testes Leukemia mononuclear	+	+	+	+	+	+	+	+	+	+	+	+	+	+		+	+	+	+	+	+	+ X	+	+	+
Bilateral, interstitial cell, adenoma				_						X	X		X						**	X	X	**		X	X
Interstitial cell, adenoma				X										X	X	X			X			X	X		

TABLE A2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE RATS: 600 ppm (Continued)

WEEKS ON STUDY	1 0	0	0	0	0	0	0	0	0	0	0	0	1 0	0	0	0	0	0	0	0	0	0	0	0	0	
21422	5	Š	5	5	5	5	5	5	5	5	5	Š	5	5	5	5	Š	5	5	5	5	5	5	Š	5	
	1	1	•	1	-	-			1	-	1	-	-	-	•	1	Ψ-	-	-	•	_	-	-	-	1	TOTAL
CARCASS	5	2	3	5	6	7	2	3	5	5	6	ê	2	4	4	Ĝ	Ĝ	8	3	3	4	5	5	6	7	TUMORS
ID	2	6	4	9	6	8	4	8	6	8	0	7	8	2	9	1	3	0	0	7	8	0	4	4	6	
	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	
ALIMENTARY SYSTEM	<u> </u>																									J
Esophagus	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 1
Leukemia mononuclear Intestine large	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Intestine large, cecum	+	+	+	+	+	+	÷	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Intestine large, colon Intestine large, rectum	++	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 50
Intestine small	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	÷	÷	+	+	÷	÷	+	÷	50
Intestine small, duodenum	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 3
Leukemia mononuclear Intestine small, ileum	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Leukemia mononuclear																X										3
Intestine small, jejunum Leukemia mononuclear	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 2
Liver	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Hepatocellular carcinoma	i								Х																	1
Hepatocellular adenoma Histiocytic sarcoma, metastatic		X										X														2
Leukemia mononuclear	X			X			X				X	X		X	X	X				X			X		X	25
Mesentery																										2
Leukemia mononuclear Pancreas	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Leukemia mononuclear	1						•	•	•	•	•			•		,			•	•	•			•		7
Salivary glands Leukemia mononuclear	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 1
Stomach	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Stomach, forestomach	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Leukemia mononuclear Stomach, glandular	1	+	+	+	+	+	+	+	+	+	+	+	_	+	4	X	+	+	+	+	+	+	X	+	+	7 50
Leukemia mononuciear	'		,		•	,	•	•	•	•	,	•	•		т	•	•		•	,	•			•	•	4
CARDIOVASCULAR SYSTEM																										
Blood vessel	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Heart	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Leukemia mononuclear	X			Х												Х				Х			Х		X	17
ENDOCRINE SYSTEM	-																									
Adrenal gland	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Capsule, fibrosarcoma, metastatic, skin Adrenal gland, cortex	1	4		_	+	_	_	_	_	4	4	_	+	4	_	_	4	+	_	+	+	+	+	_	+	50
Adenoma	l '	,				,	•		•			,	•	•			•				•	•	•	•	•	i
Leukemia mononuclear	X			X			X									X							X			16
Adrenal gland, medulla Leukemia mononuclear	X	+	+	*	+	+	*	+	+	+	+	+	+	+	+	*	+	+	+	+	+	+	X	+	+	50 16
Pheochromocytoma malignant	"			••			4									••							••			1
Pheochromocytoma, NOS	1										X				X			X								7
Bilateral, pheochromocytoma, NOS Islets, pancreatic	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	1 50
Adenoma	'				•		•	X	•	•	•	·	•	·	•									·		2
Parathyroid gland	+	M	+	M	+	+	+	+	+	+	+	M	+	+	+	+	+	+	+	+	+	M +	+	+	+	45 50
Pituitary gland Pars distalis, adenoma	+ X	*	•	*	X X		_	-	*	*	•	+	*	~	+	+	X X	*	*	_	-		-	*	•	24
Pars distalis, leukemia mononuclear	"	•••														X							X			5
Pars intermedia, leukemia mononuclear Pars nervosa, leukemia mononuclear																										1 1
Thyroid gland	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Leukemia mononuclear	1									_																2
C-cell, adenoma C cell, carcinoma		х			Х		X			X		X			X											7
Follicle, adenoma	Į.	Λ					X																			2
GENERAL BODY SYSTEM																										-
None SISTEM																										
GENITAL SYSTEM																										-
Epididymis	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Preputial gland	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Adenoma Leukemia mononuclear																		A								1 3
Prostate	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	±	+	+	50
Leukemia mononuclear Seminal vesicle	1.					_												_					X			3 5
Testes	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Leukemia mononuclear		•	•	**		.,	•			**	•	v			40	**	*				v	v	v	•	tr	2
Bilateral, interstitial cell, adenoma Interstitial cell, adenoma	X	X	X	X		X	X	X	X	X	X	X		X	X	X	X	х		X	Х	X	X	X	X	27 9
sound com, additiona								A										^								"

TABLE A2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE RATS: 600 ppm (Continued)

					``	U 11	****		,																
WEEKS ON STUDY	0 0 5	0 4 1	0 6 9	0 8 2	0 8 2	0 8 3	0 8 8	0 8 9	0 9 0	0 9 0	0 9 6	0 9 8	0 9 8	0 9 9	0 9 9	1 0 0	1 0 0	1 0 0	1 0 2	1 0 2	1 0 2	1 0 4	1 0 5	1 0 5	1 0 5
CARCASS ID	1 6 9 1	1 2 2 1	1 3 3 1	1 4 1	1 3 9 1	1 4 3 1	7 3 1	3 6 1	1 3 5 1	1 2 3 1	1 4 7 1	1 2 1	7 0 1	1 2 9 1	1 2 5 1	1 4 0 1	1 5 3 1	1 5 5 1	1 4 4 1	1 4 5 1	1 7 1	1 7 9 1	1 3 1 1	1 4 6 1	1 5 1
HEMATOPOIETIC SYSTEM Blood	-								+							_									
Leukemia mononuclear Bone marrow Leukemia mononuclear Lymph node Fibrosaroma, metastatic, skin Iliac, leukemia mononuclear	+	+	* *	+	+	+	* *	* X +	X + X +	+ *	* X +	+	* *	+	* *	* *	+	+	* *	+	+	* *	* * *	+	+
Mediastinal, leukemia mononuclear Mesenteric, leukemia mononuclear Renal, carcinoma, metastatic, kidney			x		X X		x	X X	X X		x		x		x	x	X X	x	x			x	X		
Lymph node, mandibular Leukemia mononuclear Spleen	M +	+	M +	M +	M +	+	* X +	+	M +	+	* X	+	+	+	* *	+	* X +	* X +	X +	+	+	* X +	* X +	+	+
Hemangosarcoma Leukemia mononuclear Thymus Carcinoma, metastatic, kidney Leukemia mononuclear	+	+	X +	+	X	+	X +	X	Х + Х	+	X + X	+	X	+	X + X	X +	Х + Х	X X +	X + X	+	+	X + X	X	M	+
INTEGUMENTARY SYSTEM Mammary gland Skin Histocytic sarcoma, metastatic Keratoacanthoma Subcutaneous tissue, fibrosarcoma	++	++	+	M +	++	++	M +	++	+	+ + x	++	+ + X	++	++	++	++	++	+ +	+++	+	+	++	+++	+++	++
MUSCULOSKELETAL SYSTEM Bone	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
NERVOUS SYSTEM Brain Leukemia mononuclear Cerebrum, astrocytoma malignast	+	+	+ X	+	+	+	*	*	+	+	+	+	*	+	* X	+	+	+	+	+	+	+ X	+	+	+
RESPIRATORY SYSTEM Lung Alveolar/bronchiolar carcinoma Carcinoma, metastatic, kidney	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+ X	+	+	+	+	+	+
Fibrosarcoma, metastatic, skin Histocytic sarcoma, metastatic Leukemia mononuclear Nose Trachea Leukemia mononuclear	++	÷ +	X + +	++	X + +	++	X + + X	X + +	X + +	* + +	X + + X	x + +	X + +	++	X + + X	X + +	X + + X	X + +	+	++	++	X + +	X + + X	++	++
SPECIAL SENSES SYSTEM Zymbal gland Carcinoma																+ X									
URINARY SYSTEM Kidney Histiocytic sarcoma, metastatic Leukemia mononuclear Capsule, fibrosarcoma, metastatic, skin Renal tubule, adenoma	+	+	+ X	+	+	+	+ X	+ X	+ X	+ x	+ X	*	+ X	+	+ X	+	+	+	+	+	+	+ X	+	+	+
kenai tuouis, adenoma Transitional epithelium, carcinema Urinary bladder Fibrosarcoma, metastatic, skin Leukema mononuclear	+	+	+	+	+	+	+ X	+ X	+	*	+ X	+	+	+	+ X	+	+	+ X	X +	+	+	+ X	+ X	+	+

TABLE A2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE RATS: 600 ppm (Continued)

WEEKS ON	-	1	_	-	_	_		_	_	-	-	-	_	-	7	-		-	-	-	-	_	1	_	1	
STUDY	0 5	0 5	0 5	0 5	0 5	0 5	0 5	0 5	0 5	0 5	0 5	0 5	0 5	0 5	0 5	0 5	0 5	5	0 5	TOTAL						
CARCASS ID	1 5 2 1	1 2 6 1	1 3 4 1	1 5 9	1 6 6	1 7 8 1	1 2 4 1	1 3 8 1	1 5 6 1	1 5 8 1	1 6 0 1	1 6 7 1	1 2 8 1	1 4 2 1	1 4 9 1	1 6 1	1 6 3 1	1 8 0 1	1 3 0 1	1 3 7 1	1 4 8 1	1 5 0 1	1 5 4 1	1 6 4 1	1 7 6 1	TISSUES
HEMATOPOIETIC SYSTEM Blood															-											1
Leukemia mononuclear Bone marrow Leukemia mononuclear	+	+	+	+	+	+	* X	+	+	+	+	+	+	x	+	*	+	+	+	+	+	+	x	+	+	50 15
Lymph node Fibrosarcoma, metastatic, skin Iliac, leukemia mononuclear Mediastinal, leukemia mononuclear	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+ X	+	+	+	+	+	+	+	+	+	50 1 1 5
Mesenteric, leukemia mononuclear Renal, carcinoma, metastatic, kidney	X			X										X		X				X			X		X	20
Lymph node, mandibular Leukemia mononuclear Spleen	+	+	+	+	+	+	+	+	++	+	+	+	+	+	+	* *	+	+	+	* *	+	+	* *	+	+	45 11 50
Hemangiosarcoma Leukemia mononuclear Thymus	X +	+	+	X +	M	X	X +	+	+	+	+	X +	+	X	X	X +	+	M	+	X	+	+	X +	+	X M	25 46
Carcinoma, metastatic, kidney Leukemia mononuclear																x							X			9
INTEGUMENTARY SYSTEM Mammary gland Skin Histocytic sarcoma, metastatic	++	+	++	++	+	++	+	+	++	++	++	+	++	+	++	+	+	+	++	++	+	+	++	+	++	48 50 1
Keratoacanthoma Subcutaneous tissue, fibrosarcoma			X					x																		1 2
MUSCULOSKELETAL SYSTEM Bone	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
NERVOUS SYSTEM Brain Leukemia mononuclear Cerebrum, astrocytoma malignant	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+ X	+	+	+	+	+	+	+	*	+	+	50 7 1
RESPIRATORY SYSTEM Lung Alveolar/bronchiolar carcinoma Carcinoma, metastatic, kidney Fibrosarcoma, metastatic, skin	+	+	+	+	*	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 1 1
Histocytic sarcoma, metastatic Leukemia mononuclear Nose Trachea Leukemia mononuclear	X + +	++	++	X + +	++	++	X + +	++	+++	++	X + +	++	+	X + +	X + +	X + +	++	++	+	X + +	+	+	X + +	+	+ +	1 22 50 50 50
SPECIAL SENSES SYSTEM Zymbai gland Carcinoma							*		-												•					2 2
URINARY SYSTEM Kidney Histrocytic sarcoma, metastatic Leukemia mononuclear Capsule, fibrosarcoma, metastatic, skin	+	+	+	+	+	+	+	+ x	+	+	+	+	+	+	+	+ X	+	+	+	+	+	+	+ X	+	+	50 1 10 1
Renal tubule, adenoma Transitional epithelium, carcinoma Urinary bladder	_	_	_	_	_	+	+	Т	_	_	_	_	_	_	_	_	_	_	4	+	+	+	+	+	+	1 50

TABLE A2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE RATS IN THE TWO-YEAR INHALATION STUDY OF TOLUENE: 1,200 ppm

1.	NIALA.	U	17	21	UD	1 (JF	10	L	EI	Œ:	1,	ZUL	P	biii										
WEEKS ON STUDY	0 3 0	0 6 9	0 7 1	0 7 8	0 7 8	0 8 0	0 8 0	0 8 0	0 8 6	0 8 6	0 8 6	0 8 7	0 8 9	0 9 1	0 9 1	0 9 2	0 9 4	0 9 5	0 9 6	0 9 8	9 8	0 9 9	1 0 1	1 0 2	1 0 3
CARCASS ID	9 6 1	2 8 7 1	9 1 1	2 4 7 1	2 4 8 1	2 8 4 1	7 0 1	2 8 1 1	2 4 9 1	6 2 1	7 2 1	2 5 7 1	9 4 1	2 7 1	5 5 1	9 7 1	6 1 1	2 6 5 1	2 6 3 1	2 5 8 1	9 9 1	2 4 3 1	7 5 1	2 8 3 1	2 8 5 1
ALIMENTARY SYSTEM	[_															
Esophagus	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine large Intestine large, cecum	+	+	+	+ A	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	++
Leukemia mononuclear	1 '	-	•	•	•					•	•	•	,	•		•	X		•		,				•
Intestine lange, colon Leukemia mononuclear	+	+	*	A	+	+	+	+	+	+	+	+	+	+	+	+	*	+	+	+	+	+	+	+	+
Intestine large, rectum	+	+	7	A	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine small	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine small, duodenum Leukemia mononuclear	+	+	+	A	+	+	+	+	+	+	+	+	+	+	+	+	X	+	+	+	+	+	+	+	+
Intestine small, ileum	+	+	+	A	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine small, jejunum Liver	+	+	+	A +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Leukemia mononuclear			X		X	X	т	7	X	т-	X	1	-		•	X	X	χ̈́	•	*		•	X	X	
Serosa, mesothelioma malignant	1	X																							
Mesentery Pancreas	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Leukemia mononuclear			X														X								
Mesothelioma malignant Salivary glands	+	X +	_			4	4	_		4	_	+	_	_	+	4	_	_	+	+	+	+	+	+	+
Stomach	+	+	+	÷	+	+	÷	+	+	+	÷	+	÷	÷	++	+	÷	+	+	+	+	+	+	+	+
Stomach, forestomach	+	+	+	A	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Leukemia mononuclear Serosa, mesothelioma malignant		X															А								
Stomach, glandular Leukemia mononuclear	+	+	+	A	+	+	+	+	*	+	+	+	+	+	+	*	*	+	+	+	+	+	+	+	+
CARDIOVASCULAR SYSTEM	I	_																							
Blood vessel	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Heart Leukemia mononuclear	+	+	+	+	*	+	+	+	X X	+	+	+	+	+	+	*	*	+	+	+	+	+	+	*	+
						**																			
ENDOCRINE SYSTEM Adrenal gland	+	_	_	_	_	_	_	_		_	_		_	_		_	_				_	_	_	_	_
Capsule, mesothelioma malignant		X		~	Τ.	~	7	_	_	~	7	~		~	~	~	7	~	7	~	_	7	-	7	7
Adrenal gland, cortex	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Leukemia mononuclear Adrenal gland, medulla	+	+	X +	+	X +	+	+	+	X	+	X	+	+	+	+	+	X	+	+	X	+	+	X +	X +	+
Leukemia mononuclear			X		X				*		X,						*			*			X	X	
Pheochromocytoma, NOS Islets, pancreatic		+	+	+	_	_	+	+	_	+	4	+	+	+	+	+	+	+	+	+	+	+	+	+	X
Adenoma	1	•			•	,	•	ŕ	•	•	•				•					•			X		
Carcinoma Parathyroid gland		М	M	_		_	М	_	1	_	+		+	X +	_	_		_	_	1	_	_		М	+
Pituitary gland	1 7	+ X	+ X	+	+	+	+ X	+	+	+ X	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Pars distalis, adenoma		X	X		X		X	X		X		X,	x	X	X			X			X		X		X
Pars distalis, carcinoma Pars distalis, leukemia monocytic													А										х		
Pars distalis, leukemia mononuclear	1		X																	X					
Thyroid gland C-cell, adenoma	+	+	+	+	+	+	+	+	+	+	+	*	+	+	+	+	X	X	+	+	+	+	+	+	+
GENERAL BODY SYSTEM None																									
GENITAL SYSTEM			-										-												-
Epididymis Preputial gland	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Prostate	‡	+	+	+	+++++	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Testes	+	+	+	+	+	÷ X	+	÷	X	+	*	+	÷	+	+	+	+	+	+	+ X	+	+ *	+	+	+
Bilateral, interstitual cell, adenoma Interstitual cell, adenoma			x	x		Х	x		X	x	Х				X	x	X	X	X	Х		X	X	x	X
Serosa, mesothelioma malignant		X																							

TABLE A2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE RATS: 1,200 ppm (Continued)

WEEKS ON	11	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	T
STUDY	0	4	4	0 5	0 5	0 5	0 5	0 5	0 5	0 5	0 5	0 5	0 5	0 5	0 5	0 5	0 5	5	0 5	0 5	5	0 5	0 5	0 5	0 5	TOTAL
CARCASS ID	2 4 6 1	2 6 0 1	6 9 1	2 5 4 1	7 4 1	7 7 1	2 8 9	9 8 1	5 3 1	7 3 1	2 9 2 1	2 4 2 1	2 5 6 1	2 5 9 1	9 0 1	9 3 1	2 4 4 1	2 5 0 1	2 5 1	2 5 2 1	2 8 2 1	2 4 1 1	2 7 6 1	2 8 0 1	2 8 8 1	TISSUES
ALIMENTARY SYSTEM	-																									
Esophagus Intestine large	++	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 50
Intestine large, cecum	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	49
Leukemia mononuclear Intestine large, colon	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	49
Leukemia mononuclear	١.																									2
Intestine large, rectum Intestine small	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	49 50
Intestine small, duodenum	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	49
Leukemia mononuclear Intestine small, ileum	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	49
Intestine small, jejunum Liver	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	49 50
Leukemia mononuclear	+	X	+	x ⁺	+	+	+	X	X	+	+	+	+	+	+	*	+	+	X	*	+	+	x	+	+	19
Serosa, mesothelioma malignant																					غر					1 4
Mesentery Pancreas	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Leukemia mononuclear																			X							3
Mesothelioma malignant Salivary glands	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Stomach	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Stomach, forestomach Leukemia mononuclear	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	49
Serosa, mesothelioma malignant																										1
Stomach, glandular Leukemia mononuclear	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	49
CARDIOVASCULAR SYSTEM	.																									
Blood vessel	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Heart Leukemia mononuclear	+	X +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
ENDOCRINE SYSTEM	.																									
Adrenal gland	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Capsule, mesothelioma malignant Adrenal gland, cortex	1 +	_	_		_					_	_	4.	_	_	_	_	_	_	_	_	_	_	_	_	_	50
Leukemia mononuclear	*	X			7		т	Τ.	_	•	т		т	Τ.	_	т	Τ.	7	X					•	-	10
Adrenal gland, medulla Leukemia mononuclear	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	*	+	+	+	+	+	+	50 10
Pheochromocytoma, NOS			X		X				x									X	Λ			X				6
Islets, pancreatic	+	+	+	+	+	+	+	+	X +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 2
Adenoma Carcinoma										А																1
Parathyroid gland	+	+	+	+	+	M	+	+	+	+	+	+	+	+	+	+	+	+	M	+	+	+	+	+	+	44
Pituitary gland Pars distalis, adenoma	T X	+	+	+	X X	+	+	+	+	+	*	+	+	+	+	+	+ X	+	+	X	+	+	+	+	+	50 18
Pars distalis, carcinoma	1																									1
Pars distalis, leukemia monocytic Pars distalis, leukemia mononuclear																			x							1 3
Thyroid gland	+	+	+	+	+	+ X	+	+	+	+	+	+	+	±	+	+	+	+	±	+	+	+	+	+	+	50
C cell, adenoma	-					Х				Х				Х					Х				X			8
GENERAL BODY SYSTEM None		_																								
GENITAL SYSTEM	·																									
Epididymis Preputial gland	+	+	+	+	+	+	+	+	+	+	+	+	++	+	+	+	+	+	+	+	+	+	+	+	+	50 50
Prostate	++++	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+		+	+	+	50
Testes Pulstonel interestinal cell adenome	T + X	+	+	+	+	*	*X	*X	*	*	*X	*	+ X	*	+ X	*	+	Ť	*	*	*	*	*	+	+ X	50 30
Bilateral, interstitial cell, adenoma Interstitial cell, adenoma Serosa, mesothelioma malignant	, x	X	X	X	X	A	Х	A	А	А	А	A	А	A	A	А		А	A	А	Α.	A	Λ.		Λ.	10
																										· 1———

TABLE A2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE RATS: 1,200 ppm (Continued)

					``	··			.,																
WEEKS ON STUDY	0 3 0	0 6 9	0 7 1	0 7 8	0 7 8	0 8 0	0 8 0	0 8 0	0 8 6	0 8 6	0 8 6	0 8 7	0 8 9	0 9 1	0 9 1	0 9 2	0 9 4	0 9 5	0 9 6	0 9 8	0 9 8	0 9 9	1 0 1	1 0 2	1 0 3
CARCASS ID	2 9 6 1	2 8 7	9 1 1	2 4 7 1	2 4 8 1	2 8 4 1	7 0 1	2 8 1 1	2 4 9 1	2 6 2 1	7 2 1	5 7 1	2 9 4 1	7 1 1	2 5 5 1	2 9 7 1	2 6 1	2 6 5 1	2 6 3 1	2 5 8 1	9 9	2 4 3 1	2 7 5 1	2 8 3 1	2 8 5 1
HEMATOPOIETIC SYSTEM																									
Blood Leukemia mononuclear				+		*										*									
Bone marrow	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	*	<u>+</u>	*X	+	±	+	+	<u>+</u>	+	+
Leukemia mononuclear Lymph node		4	X	+	X	X	+	+	X	4	X	+	+	+	+	X +	* * * X	X +	+	X	+	+	X	X	+
Leukemia mononuclear	'	•	'		•		,	•		,	X	•	•	•	•	Ċ	X			•	·	•	·		
Mediastinal, leukemia mononuclear Mesenteric, leukemia mononuclear			X		X	х			x		x					X	х	X		X			X	X	
Lymph node, mandibular	M	+	+	M	M	M	M	M	M	M	M	M	M	M	M	+	+	+	+	+	+	+	+	+	+
Leukemia mononuclear Spleen	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	X +	X	+	+	+	+	+	X +	X	+
Leukemia mononuclear Lymphoma malignant		·	X		X	X			X		X					X	*	X		X			X	X	
Capsule, mesothelioma malignant Thymus	+	X +	+	+	+	+	+	+	*	+	+	+	+	+	+	M	+	+	+	M	+	+	+	+	M
Leukemia mononuclear Osteosarcoma, metastatic, uncertain			X		X	X			X																
primary site	1																		X						
INTEGUMENTARY SYSTEM	_																								
Mammary gland Skin	‡	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Subcutaneous tissue, fibroma											*														
Subcutaneous tissue, fibrosarcoma Tail, papilloma squamous	}																		X						
MUSCULOSKELETAL SYSTEM																								_	
Bone Skeletal muscle	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Fibrosarcoma	x																								
NERVOUS SYSTEM																									
Brain	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Leukemia mononuclear			X		X																				
RESPIRATORY SYSTEM		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Carcinoma adenosquamous			,	·	·	•	•	·		•	•	X	•	·							•				
Fibrous histiocytoma, metastatic, skin Leukemia mononuclear			X		X	X			X		X					X	X	X		X			X	X	
Osteosarcoma, metastatic, uncertain primary site	-																		X						
Nose	+	+	+	A	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Trachea	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
SPECIAL SENSES SYSTEM	_ _	_			-																				
Eye Zymbal gland																									
Carcinoma																									
URINARY SYSTEM			-																						
Kidney Leukemia mononuclear	+	+	+	+	+	+	+	+	+	+	*	+	+	+	+	+	¥	+	+	+	+	+	*	+	+
Capsule, mesothelioma malignant		X									Λ						41	_					^		
Renal tubule, adenoma Urinary bladder	+	_	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	X	+	+	+	+	+	+	+
Leukemia mononuclear	'	_	•			•	,		X	,	•	•	,	,	,	•	,	,					X	•	•
Serosa, mesothelioma malignant Transitional epithelium, papilloma		X																							

TABLE A2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE RATS: 1,200 ppm (Continued)

WEEKS ON STUDY	1 0 4	1 0 4	1 0 4	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	TOTAL
CARCASS ID	2 4 6 1	6 0 1	2 6 9 1	2 5 4 1	7 4 1	2 7 7 1	2 8 9 1	9 8 1	2 5 3 1	7 3 1	2 9 2 1	2 4 2 1	2 5 6 1	2 5 9 1	9 0 1	9 3 1	2 4 4 1	2 5 0 1	5 1 1	5 2 1	2 8 2 1	2 4 1	2 7 6 1	2 8 0 1	2 8 8 1	TISSUES
HEMATOPOIETIC SYSTEM																										
Blood Leukemia mononuclear			+																							4 2
Bone marrow	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 12
Leukemia mononuclear Lymph node	+	X	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Leukemia mononuclear Mediastinal, leukemia mononuclear																										2
Mesenteric, leukemia mononuclear	1	X							X							X			X	X			X			17
Lymph node, mandibular Leukemia mononuclear	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	37 4
Spleen	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Leukemia mononuclear Lymphoma malignant		X		X				X	X							X			X	X			X			19 1
Capsule, mesothelioma malignant	1																									1
Thymus Leukemia mononuclear	+	+	M	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	46 4
Osteosarcoma, metastatic, uncertain primary site																										1
INTEGUMENTARY SYSTEM																										-
Mammary gland Skin	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	M	+	+	+	+	+	+	+	+	49 50
Subcutaneous tissue, fibroma	+	+	+	7		Ψ.	Ŧ	7	+	_	*	+	*	Τ.	7	_	•	_	7	_	т	_		_	T	1
Subcutaneous tissue, fibrosarcoma Tail, papilloma squamous										X																1
MUSCULOSKELETAL SYSTEM	<u> </u>																									-
Bone Skeletal muscle	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Fibrosarcoma																										1
NERVOUS SYSTEM																										
Brain Leukemia mononuclear	+	+ X	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 3
RESPIRATORY SYSTEM	1	+	4	4	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Carcinoma adenosquamous		·				·		·									·			·	·				•	1
Fibrous histiocytoma, metastatic, skin Leukemia mononuclear		X	X																X	X						1 14
Osteosarcoma, metastatic, uncertain primary site																										1
Nose	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	49
Trachea	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
SPECIAL SENSES SYSTEM											,															2
Eye Zymbal gland			+	+											+											1 1
Carcinoma			X																							1
URINARY SYSTEM				_																						-
Kidney Leukemia mononuclear	+	+ ¥	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 4
Capsule, mesothelioma malignant		Λ																								1
Renal tubule, adenoma Unnary bladder	+	_	_	4	+	+	+	4	+	_	+	_	4	+		+	+	+	X	+	+	+	+	+	+	50
Leukemia mononuclear	"	X	7	7	r	r	-	*	т	,	r	Τ'	τ'	1.	r	r	,-	r	,	-	r	,	•	r	'	3
Serosa, mesothelioma malignant Transitional epithelium, papilloma												x														1 1
,	[

TABLE A3. ANALYSIS OF PRIMARY TUMORS IN MALE RATS IN THE TWO-YEAR INHALATION STUDY OF TOLUENE

	Chamber Control	600 ppm	1,200 ppm
Adrenal Gland Medulla: Pheochromocyto	ma		
Overall Rates (a)	7/49 (14%)	8/50 (16%)	6/50 (12%)
Adjusted Rates (b)	21.6%	23.2%	24.6%
Terminal Rates (c)	5/29 (17%)	3/28 (11%)	4/22 (18%)
Day of First Observation	618	670	715
Life Table Tests (d)	P=0.507	P=0.516	P=0.558
Logistic Regression Tests (d)	P = 0.487N	P = 0.548	P = 0.552N
Cochran-Armitage Trend Test (d)	P = 0.427N		
Fisher Exact Test (d)		P = 0.517	P = 0.484N
Adrenal Gland Medulla: Pheochromocyto	ma or Malignant Pheochr	omocytoma	
Overall Rates (a)	7/49 (14%)	9/50 (18%)	6/50 (12%)
Adjusted Rates (b)	21.6%	26.3%	24.6%
Terminal Rates (c)	5/29 (17%)	4/28 (14%)	4/22 (18%)
Day of First Observation	618	670	715
Life Table Tests (d)	P=0.497	P=0.410	P=0.558
Logistic Regression Tests (d)	P = 0.491N	P=0.442	P = 0.552N
Cochran-Armitage Trend Test (d)	P = 0.431N P = 0.427N	1 -0.774	1 -0.00214
Fisher Exact Test (d)	r = 0.42/N	D=0.410	D=0.404NT
risher Exact lest(d)		P = 0.410	P = 0.484N
Pancreatic Islets: Adenoma or Carcinoma		0/50 / 46%	0/50 / 00 \
Overall Rates (a)	3/50 (6%)	2/50 (4%)	3/50 (6%)
Adjusted Rates (b)	8.8%	7.1%	10.4%
Terminal Rates (c)	2/30 (7%)	2/28 (7%)	1/22 (5%)
Day of First Observation	618	72 9	632
Life Table Tests (d)	P = 0.487	P = 0.525N	P = 0.571
Logistic Regression Tests (d)	P = 0.568	P = 0.488N	P = 0.649
Cochran-Armitage Trend Test (d)	P = 0.588N		
Fisher Exact Test (d)		P = 0.500N	P = 0.661 N
Liver: Hepatocellular Adenoma			
Overall Rates (a)	4/50 (8%)	2/50 (4%)	0/50 (0%)
Adjusted Rates (b)	12.4%	7.1%	0.0%
Terminal Rates (c)	3/30 (10%)	2/28 (7%)	0/22 (0%)
Day of First Observation	652	729	0/22 (0 /0)
Life Table Tests (d)	P=0.061N	P = 0.358N	P = 0.101N
Logistic Regression Tests (d)	P = 0.061N P = 0.042N	P = 0.335N P = 0.315N	P = 0.101N P = 0.072N
Cochran-Armitage Trend Test (d)	P = 0.042N P = 0.037N	1 -0.01014	1 -0.01214
Fisher Exact Test (d)	F - 0.03/14	P = 0.339N	P = 0.059N
		r = 0.33314	P = 0.03N
iver: Hepatocellular Adenoma or Carcin Overall Rates (a)	oma 4/50 (8%)	3/50 (6%)	0/50 (0%)
Adjusted Rates (b)	12.4%	10.7%	0.0%
	3/30 (10%)	3/28 (11%)	0.0%
Terminal Rates (c)			0/44 (070)
Day of First Observation	652 P = 0.091 N	729 B = 0.526N	D=0.101N
Life Table Tests (d)	P=0.081N	P = 0.526N	P = 0.101N
Logistic Regression Tests (d)	P=0.056N	P = 0.473N	P = 0.072N
Cochran-Armitage Trend Test (d)	P = 0.049N	m	.
Fisher Exact Test (d)		P = 0.500N	P = 0.059N
Pituitary Gland/Pars Distalis: Adenoma			
Overall Rates (a)	22/49 (45%)	24/50 (48%)	18/50 (36%)
Adjusted Rates (b)	52. 4%	59.7%	45.0%
Terminal Rates (c)	11/30 (37%)	13/28 (46%)	4/22 (18%)
Day of First Observation	473	572	478
Life Table Tests (d)	P = 0.513N	P=0.429	P = 0.511N
Logistic Regression Tests (d)	P=0.212N	P = 0.455	P = 0.221N
			1
Cochran-Armitage Trend Test (d)	P = 0.212N		

TABLE A3. ANALYSIS OF PRIMARY TUMORS IN MALE RATS IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

	Chamber Control	600 ppm	1,200 ppm
Pituitary Gland/Pars Distalis: Adenoma o	r Carcinoma		
Overall Rates (a)	22/49 (45%)	24/50 (48%)	19/50 (38%)
Adjusted Rates (b)	52.4%	59.7%	46.4%
Terminal Rates (c)	11/30 (37%)	13/28 (46%)	4/22 (18%)
Day of First Observation	473	572 D-0 400	478
Life Table Tests (d)	P=0.484	P=0.429	P = 0.551
Logistic Regression Tests (d)	P = 0.274N	P = 0.455	P = 0.284N
Cochran-Armitage Trend Test (d) Fisher Exact Test (d)	P = 0.276N	P = 0.457	P = 0.311N
ubcutaneous Tissue: Fibroma			
Overall Rates (e)	3/50 (6%)	0/50 (0%)	1/50 (2%)
Adjusted Rates (b)	8.9%	0.0%	2.4%
Terminal Rates (c)	2/30 (7%)	0/28 (0%)	0/22 (0%)
		0/28 (0%)	
Day of First Observation	625 D = 0.991 N	D_0 10131	597
Life Table Tests (d)	P=0.221N	P=0.131N	P = 0.382N
Logistic Regression Tests (d)	P = 0.175N	P=0.119N	P = 0.304N
Cochran-Armitage Trend Test (d)	P = 0.176N		
Fisher Exact Test (d)		P = 0.121N	P = 0.309N
ubcutaneous Tissue: Fibroma or Fibrosa		0/50 / 400 :	0(50.44%)
Overall Rates (e)	3/50 (6%)	2/50 (4%)	2/50 (4%)
Adjusted Rates (b)	8.9%	5.9%	6.9%
Terminal Rates (c)	2/30 (7%)	1/28 (4%)	1/22 (5%)
Day of First Observation	625	625	597
Life Table Tests (d)	P = 0.500N	P = 0.519N	P = 0.601 N
Logistic Regression Tests (d)	P = 0.410N	P = 0.499N	P = 0.508N
Cochran-Armitage Trend Test (d)	P = 0.406N		
Fisher Exact Test (d)		P = 0.500N	P = 0.500N
Cestis: Interstitial Cell Adenoma			
Overall Rates (a)	36/50 (72%)	36/50 (72%)	40/50 (80%)
Adjusted Rates (b)	94.6%	92.2%	95.1%
Terminal Rates (c)	28/30 (93%)	25/28 (89%)	20/22 (91%)
Day of First Observation	614	570	495
Life Table Tests (d)	P=0.013	P=0.488	P=0.019
Logistic Regression Tests (d)	P=0.072	P = 0.435N	P=0.096
		1 -0.40011	1 -0.000
Cochran-Armitage Trend Test (d) Fisher Exact Test (d)	P = 0.210	D-0.599N	P = 0.241
		P=0.588N	r = U.241
'hyroid Gland: C-Cell Adenoma Overall Rates (a)	5/50 (10%)	7/50 (14%)	8/50 (16%)
Adjusted Rates (b)	15.2%	22.3%	29.1%
Terminal Rates (c)	4/30 (13%)	5/28 (18%)	5/22 (23%)
Day of First Observation	589	686	606
			P=0.145
Life Table Tests (d)	P=0.113	P=0.360	
Logistic Regression Tests (d)	P=0.194	P = 0.404	P = 0.245
Cochran-Armitage Trend Test (d)	P = 0.231	D 0000	D
Fisher Exact Test (d)		P = 0.380	P = 0.277
hyroid Gland: C-Cell Adenoma or Carcin		9/E0 /16% \	0/E0 (1.0% \
Overall Rates (a)	5/50 (10%)	8/50 (16%)	8/50 (16%)
Adjusted Rates (b)	15.2%	25.7%	29.1%
Terminal Rates (c)	4/30 (13%)	6/28 (21%)	5/22 (23%)
Day of First Observation	589	686	606
Life Table Tests (d)	P = 0.111	P = 0.257	P = 0.145
Logistic Regression Tests (d)	P = 0.195	P = 0.300	P = 0.245
Cochran-Armitage Trend Test (d)	P = 0.236		

TABLE A3. ANALYSIS OF PRIMARY TUMORS IN MALE RATS IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

	Chamber Control	600 ppm	1,200 ppm
lematopoietic System: Mononuclear Le	oukemia		
Overall Rates (e)	17/50 (34%)	26/50 (52%)	19/50 (38%)
Adjusted Rates (b)	41.3%	62.3%	52.1%
Terminal Rates (c)	8/30 (27%)	13/28 (46%)	7/22 (32%)
Day of First Observation	416	478	495
Life Table Tests (d)	P=0.184	P = 0.085	P≈0.231
Logistic Regression Tests (d)	P = 0.374	P = 0.053	P = 0.429
Cochran-Armitage Trend Test (d)	P = 0.380		- +
Fisher Exact Test (d)		P = 0.053	P = 0.418

⁽a) Number of tumor-bearing animals/number of animals examined microscopically at the site

⁽b) Kaplan-Meier estimated tumor incidences at the end of the study after adjusting for intercurrent mortality (c) Observed tumor incidence in animals killed at the end of the study

⁽d) Beneath the control incidence are the P values associated with the trend test. Beneath the dosed group incidence are the P values corresponding to pairwise comparisons between that dosed group and the controls. The life table analysis regards tumors in animals dying prior to terminal kill as being (directly or indirectly) the cause of death. The logistic regression test regards these lesions as nonfatal. The Cochran-Armitage and Fisher exact tests compare directly the overall incidence rates. A negative trend or lower incidence in a dosed group than in controls is indicated by (N).

⁽e) Number of tumor-bearing animals/number of animals examined grossly at the site

TABLE A4. HISTORICAL INCIDENCE OF KIDNEY TUBULAR CELL TUMORS IN MALE F344/N RATS RECEIVING NO TREATMENT (a)

Study	Incidence of Adenomas or Carcinomas in Controls	
Historical Incidence for Chamber Co	ntrols at Battelle Pacific Northwest Laboratories	
Propylene oxide	0/50	
Methyl methacrylate	0/50	
Propylene	0/50	
1,2-Epoxybutane	0/50	
Dichloromethane	0/50	
Tetrachloroethylene	(b) 1/49	
Bromoethane	0/47	
TOTAL	1/346 (0.3%)	
SD(c)	0.77%	
Range (d)		
High	1/49	
Low	0/50	
Overall Historical Incidence for Untr	eated Controls in NTP Studies	
TOTAL	(e) 14/1,590 (0.9%)	
SD(c)	1.68%	
Range (d)		
High	3/50	
Low	0/50	

⁽a) Data as of May 12, 1988, for studies of at least 104 weeks

⁽b) Tubular cell adenoma

⁽c) Standard deviation
(d) Range and SD are presented for groups of 35 or more animals.
(e) Includes 10 tubular cell adenomas, 1 adenoma, NOS, 2 tubular cell adenocarcinomas, and 1 tubular adenocarcinoma

TABLE A5. SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN MALE RATS IN THE TWO-YEAR INHALATION STUDY OF TOLUENE

	Chambe	er Control	600 I	pm	1,200	ppm
nimals initially in study	60		60		60	-
animals removed	60		60		60	
animals examined histopathologically	50		50		50	
LIMENTARY SYSTEM						
Intestine large, cecum	(50)		(50)		(49)	
Erosion			1	(2%)		
Hemorrhage			2	(4%)		
Inflammation, acute	1	(2%)				
Intestine large, colon	(50)		(50)		(49)	
Granuloma					1	(2%)
Parasite metazoan	2	(4%)	4	(8%)	1	(2%)
Intestine large, rectum	(49)		(50)		(49)	
Erosion				(2%)		
Inflammation, acute				-	1	(2%)
Parasite metazoan	2	(4%)	4	(8%)		(14%)
Ulcer	_			(8%)		(6%)
Intestine small, duodenum	(50)		(50)	/	(49)	/- /
Erosion	(50)		(00)			(2%)
Ulcer	1	(2%)			-	- / - /
Intestine small, ileum	(49)	· · · · · ·	(50)		(49)	
Erosion	(-0)		(00)			(2%)
Liver	(50)		(50)		(50)	(2 70)
Angiectasis		(14%)		(22%)		(6%)
Congestion		(6%)	-	(4%)	_	(2%)
Developmental malformation		(2%)	_	(8%)		(4%)
Fatty change		(6%)	_	(4%)		(16%)
Focal cellular change	_	(56%)		(46%)	_	(46%)
Granuloma		(2%)		(2%)		(2%)
Hematopoietic cell proliferation		(270)		(4%)		(270)
Hemorrhage				,		
Necrosis		(40)		(4%)	•	(OM)
	Z	(4%)		(4%)	1	(2%)
Bile duct, dilatation	40	(0.00)		(2%)		
Bile duct, hyperplasia	48	(96%)		(98%)	49	(98%)
Bile duct, inflammation, chronic				(2%)		
Portal, inflammation, chronic	48	(96%)	45	(90%)		(92%)
Serosa, fibrosis	.~.					(2%)
Mesentery	(2)		(2)	/E041	(4)	
Inflammation, acute			1	(50%)	_	
Arteriole, inflammation, chronic						(50%)
Arteriole, inflammation, chronic active	_	(4.00%)				(25%)
Fat, necrosis		(100%)				(25%)
Pancreas	(50)		(50)		(50)	
Inflammation, acute	. -			(2%)	. -	
Acinus, atrophy		(60%)		(68%)		(60%)
Acinus, hyperplasia	1	(2%)	4	(8%)		(2%)
Arteriole, inflammation, chronic active Arteriole, mineralization						(2%) $(2%)$
Artery, inflammation, chronic		(2%)			_	
Salivary glands	(50)		(50)		(50)	
Infiltration cellular, lymphocytic	1	(2%)		(2%)		
Inflammation, chronic				(2%)		
Acinus, atrophy			1	(2%)		(2%)
Arteriole, mineralization						(2%)
Stomach	(50)		(50)		(50)	
Ulcer					· ·	(2%)

TABLE A5. SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN MALE RATS IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

ALIMENTARY SYSTEM (Continued) Stomach, forestomach	1,200	ppm
Stomach, forestomach		
Cyst	(49)	
Erosion	(43)	
Hyperplasia, squamous 2 (4%) 1 (2%) Inflammation, acute 4 (8%) 3 (6%) Inflammation, chronic 1 (2%) Inflammation, chronic 1 (2%) Inflammation, chronic 1 (2%) Inflammation, chronic active Mineralization Ulcer	1	(2%)
Inflammation, acute		(4%)
Inflammation, chronic Inflammation, chronic active Inflammation, chronic active Inflammation, chronic active Mineralization Ulcer Stomach, glandular (50) Inflammation, acute Inflammation, chronic active Mineralization Mucosa, congestion Mucosa, dilatation Mucosa, dilatation Mucosa, erosion Mucosa, erosion Mucosa, granuloma Mucosa, granuloma Mucosa, granuloma Mucosa, inflammation, acute Mucosa, inflammation, acute Mucosa, inflammation, acute Mucosa, inflammation, chronic active Heart Cardiomyopathy, chronic Hemorrhage Mineralization Artery, miner		
Inflammation, chronic active Mineralization Ulcer 4 (8%) 7 (14%) Stomach, glandular (50) (50) (50) Inflammation, acute 1 (2%) Inflammation, chronic active Mineralization Mucosa, congestion 1 (34%) 21 (42%) Mucosa, congestion 1 (34%) 21 (42%) Mucosa, getasia 2 (4%) Mucosa, ectasia 2 (4%) Mucosa, granuloma 1 (2%) Mucosa, granuloma 1 (2%) Mucosa, inflammation, acute 1 (2%) Mucosa, inflammation, acute 2 (4%) 5 (10%) CARDIOVASCULAR SYSTEM Blood vessel (50) (50) Cardiomyopathy, chronic active Heart (50) (50) Cardiomyopathy, chronic 49 (98%) 47 (94%) Hemorrhage 1 (2%) Mineralization Artery, mineralization Artery	Z	(4%)
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Pars distalis, angiectasis 3 (6%) Pars distalis, atrophy 1 (2%) Pars distalis, cyst 3 (6%) 6 (12%) Pars distalis, hemorrhage 1 (2%) Pars distalis, hyperplasia 18 (37%) 22 (44%) Pars intermedia, angiectasis 2 (4%)		
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Pars distalis, cyst 3 (6%) 6 (12%) Pars distalis, hemorrhage 1 (2%) Pars distalis, hyperplasia 18 (37%) 22 (44%) Pars intermedia, angiectasis 2 (4%)		
Pars distalis, hemorrhage 1 (2%) Pars distalis, hyperplasia 18 (37%) 22 (44%) Pars intermedia, angiectasis 2 (4%)	5	(10%)
Pars distalis, hyperplasia 18 (37%) 22 (44%) Pars intermedia, angiectasis 2 (4%)		(2%)
Pars intermedia, angiectasis 2 (4%)		(22%)
		(22%)
	1	(2%)
Pars intermedia, hyperplasia 2 (4%) Pars nervosa, pigmentation 1 (2%)		

TABLE A5. SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN MALE RATS IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

	Chambo	er Control	600 I	pm	1,200	ppm
ENDOCRINE SYSTEM (Continued)						
Thyroid gland	(50)		(50)		(50)	
Infiltration cellular, lymphocytic		(2%)	(00)		,	(2%)
Ultimobranchial cyst	1	(2%)			2	(4%)
C-cell, hyperplasia	12	(24%)	20	(40%)	14	(28%)
Follicle, ectasia	3	(6%)	2	(4%)	3	(6%)
GENERAL BODY SYSTEM None					•	•
GENITAL SYSTEM						
Preputial gland	(50)		(50)		(50)	
Abscess		(6%)	(00)		(00)	
Hyperplasia	ŭ	(370)	1	(2%)	1	(2%)
Inflammation, acute	1	(2%)	•	·-··	1	(2,0)
Inflammation, chronic		(58%)	28	(56%)	22	(44%)
Inflammation, chronic active	10	(20%)		(34%)		(50%
Mineralization					1	(2%)
Duct, ectasia		(2%)				(4%)
Prostate	(50)		(50)		(50)	
Atrophy	_	(2%)	_	.aa :	_	.a~:
Hyperplasia		(20%)		(6%)	-	(6%)
Inflammation, acute		(14%)		(14%)		(10%)
Inflammation, chronic Inflammation, chronic active		(2%) (28%)		(6%) (12%)		(4%) (20%)
Seminal vesicle	14	(20%)	(5)	(12%)	10	(20%)
Dilatation				(20%)		
Testes	(50)		(50)	(20 %)	(50)	
Atrophy		(22%)		(24%)		(18%)
Congestion		(==::,	1	(2%)	_	,
Mineralization			2	(4%)		
Arteriole, inflammation, chronic			2	(4%)	1	(2%)
Interstitial cell, hyperplasia	35	(70%)	46	(92%)	37	(74%)
HEMATOPOIETIC SYSTEM						
Bone marrow	(50)		(50)		(50)	
Myelofibrosis		(6%)		(2%)		(10%)
Lymph node	(50)		(50)	(0%)	(50)	
Congestion Edema	1	(20%)	1	(2%)		
Hyperplasia, lymphoid	-	(2%) (2%)	9	(4%)		
Mediastinal, congestion		(2%) (2%)		(10%)		
Mediastinal, hemorrhage	•	(= <i>/</i> 0 <i>)</i>	J	(-0/0/	1	(2%)
Mediastinal, hyperplasia, lymphoid			1	(2%)		(2%)
Mesenteric, congestion	1	(2%)		(2%)		(8%)
Mesenteric, hyperplasia, lymphoid		(34%)		(40%)		(44%)
Mesenteric, inflammation, acute			1	(2%)		
						(2%)
Popliteal, hyperplasia, lymphoid	(0.0)	(004)	(45)		(37)	
Lymph node, mandibular	(36)		10	(22%)	3	(8%)
Lymph node, mandibular Congestion	10	(28%)		(C ~)		
Lymph node, mandibular Congestion Cyst	10 1	(3%)	1	(2%)		(3%)
Lymph node, mandibular Congestion Cyst Hyperplasia, lymphoid	10 1 32		1 31	(2%) (69%)	30	
Lymph node, mandibular Congestion Cyst Hyperplasia, lymphoid Spleen	10 1 32 (50)	(3%) (89%)	1 31 (50)	(69%)	30 (50)	(81%)
Lymph node, mandibular Congestion Cyst Hyperplasia, lymphoid Spleen Fibrosis	10 1 32 (50) 5	(3%) (89%) (10%)	1 31 (50) 3	(69%) (6%)	30 (50) 4	(81%) (8%)
Lymph node, mandibular Congestion Cyst Hyperplasia, lymphoid Spleen Fibrosis Hematopoietic cell proliferation	10 1 32 (50) 5 42	(3%) (89%) (10%) (84%)	1 31 (50) 3	(69%)	30 (50) 4 40	(81%) (8%) (80%)
Lymph node, mandibular Congestion Cyst Hyperplasia, lymphoid Spleen Fibrosis Hematopoietic cell proliferation Hyperplasia, lymphoid	10 1 32 (50) 5 42	(3%) (89%) (10%)	1 31 (50) 3 42	(69%) (6%) (84%)	30 (50) 4 40	(81%) (8%)
Lymph node, mandibular Congestion Cyst Hyperplasia, lymphoid Spleen Fibrosis Hematopoietic cell proliferation	10 1 32 (50) 5 42 2	(3%) (89%) (10%) (84%)	1 31 (50) 3 42	(69%) (6%)	30 (50) 4 40	(81%) (8%) (80%)

TABLE A5. SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN MALE RATS IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

	Chambe	er Control	600 g	pm	1,200	ppm
HEMATOPOIETIC SYSTEM (Continued)	, ·	<u> </u>				
Thymus	(44)		(46)		(46)	
Congestion	2	(5%)	5	(11%)	1	(2%)
Cyst	4	(9%)	1	(2%)		
Hyperplasia, lymphoid			4	(9%)		
NTEGUMENTARY SYSTEM			-		·····	-
Mammary gland	(50)		(48)		(49)	
Inflammation, acute	2	(4%)				
Inflammation, chronic	1	(2%)				
Acinus, ectasia	34	(68%)	25	(52%)	28	(57%)
Duct, ectasia	10	(20%)	19	(40%)	8	(16%)
Skin	(50)		(50)		(50)	
Abscess				(2%)	1	(2%)
Cyst epithelial inclusion	1	\ = ··· /	2	(4%)		
Epidermis, cyst	1	(2%)				
Foot, ulcer					1	(2%)
Hair follicle, inflammation, chronic			1	(2%)		
Hair follicle, head, hemorrhage		(2%)				
Hair follicle, head, inflammation, acute	1	(2%)				
Head, abscess						(2%)
Head, inflammation, acute		(2%)	1	(2%)	1	(2%)
Head, inflammation, chronic active	1	(2%)				
Subcutaneous tissue, edema				(2%)		
Subcutaneous tissue, hemorrhage			1	(2%)		
Tail, inflammation, chronic active		(2%)				
Tail, subcutaneous tissue, inflammation, act	ute				1	(2%)
MUSCULOSKELETAL SYSTEM	· · · · · · · · · · · · · · · · · · ·					
Bone	(50)		(50)		(50)	
Cranium, abscess			1	(2%)		
Cranium, fibrous osteodystrophy	1	(2%)	1	(2%)	2	(4%)
Cranium, inflammation, chronic					1	(2%)
Femur, fibrous osteodystrophy	1	(2%)	4	(8%)	2	(4%)
Humerus, fracture					1	(2%)
NERVOUS SYSTEM			··-	<u> </u>		
Brain	(50)		(50)		(50)	
Compression	17-7	(10%)	,	(14%)		(12%)
Hemorrhage		(22%)		(16%)		(4%)
Necrosis		(2%)		(2%)	1	(2%)
ESPIRATORY SYSTEM						
Lung	(50)		(50)		(50)	
Congestion		(12%)		(6%)	4	(8%)
Hemorrhage		(16%)		(8%)		(10%)
Infiltration cellular, lymphocytic	50	(100%)	47	(94%)	49	(98%)
Inflammation, chronic						(2%)
Metaplasia, osseous						(2%)
Pigmentation, cholesterol	1	(2%)	2	(4%)	2	(4%)
Alveolar epithelium, hyperplasia	2	(4%)	2	(4%)	3	(6%)
Alveolus, infiltration cellular, histiocytic	26	(52%)	18	(36%)	25	(50%)
Alveolus, mineralization					2	(4%)
Arteriole, mineralization	41	(82%)	40	(80%)	38	(76%)
Bronchiole, inflammation, acute					2	(4%)
Bronchiole, mineralization	1	(2%)				
Interstitium, inflammation, chronic	5	(10%)	6	(12%)		(20%)
Interstitium, inflammation, chronic active Peribronchiolar, inflammation, chronic	5	(10%)		(6%)	2	(4%)

TABLE A5. SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN MALE RATS IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

•	Chamb	er Control	600 j	opm	1,200	ppm
RESPIRATORY SYSTEM			 			
Lung (Continued)	(50)		(50)		(50)	
Peribronchiolar, inflammation, chronic active				(2%)	(30)	
Pleura, inflammation, chronic active		(2%)	1	(270)		
Nose	(50)		(50)		(40)	
Inflammation, chronic active		(2%)	(80)		(49)	
Lumen, degeneration		(270)			1	(90%)
Lumen, foreign body	4	(8%)	5	(10%)		(2%) (4%)
Lumen, hemorrhage		(62%)		(50%)		(71%)
Mucosa, hemorrhage	01	(02 %)		(2%)	30	(1170)
Mucosa, inflammation, acute	30	(60%)	-	(60%)	40	(82%)
Mucosa, inflammation, chronic active		(2%)	00	(00 %)		(02 10)
Nasolacrimal duct, degeneration	•	(270)	1	(2%)		
Nasolacrimal duct, inflammation	1	(2%)	•	(2 %)		
Nasolacrimal duct, inflammation, acute		(8%)	1	(2%)	9	(4%)
Nasopharyngeal duct, inflammation, acute		(2%)	•	(2 %)	2	(470)
Olfactory epithelium, degeneration		(78%)	48	(96%)	49	(86%)
Olfactory epithelium, erosion	00	(1070)		(6%)		(16%)
Olfactory epithelium, metaplasia				(2%)		(2%)
Olfactory epithelium, metaplasia, squamous				(270)		(4%)
Respiratory epithelium, degeneration	15	(30%)	27	(74%)		(43%) (63%)
Respiratory epithelium, erosion		(8%)		(12%)		
Respiratory epithelium, hemorrhage	*	(6%)		(12%) (2%)	ა	(6%)
Respiratory epithelium, metaplasia, squamou	a 1	(2%)				
Respiratory epithelium, ulcer	s i	(270)	1	(2%)	1	(90%)
Vomeronasal organ, inflammation, acute	1	(2%)				(2%)
Trachea	(50)	(270)	(50)			(2%)
Inflammation, acute	(30)			(90%)	(50)	(COL)
Inflammation, chronic active				(2%)		(6%)
				(6%)	1	(2%)
PECIAL SENSES SYSTEM			·			
Eye	(2)				(2)	
Cataract		(50%)				(100%)
Retina, degeneration		(50%)				(50%)
Sclera, mineralization		(50%)				(100%)
RINARY SYSTEM	* 1					
Kidney	(50)		(50)		(50)	
Congestion	(00)			(2%)	(00)	
Infarct				(a ro)	2	(4%)
Infiltration cellular, lymphocytic	1	(2%)			2	(270)
Mineralization	•	(2 //)			1	(2%)
Nephropathy, chronic	40	(98%)	40	(96%)		(96%)
Pelvis, inflammation, acute	70	(00 10)		(2%)		(2%)
Renal tubule, cyst	1	(2%)		(4%)		(10%)
Renal tubule, hyperplasia		(8%)		(8%)	5	(1070)
Renal tubule, pigmentation		(100%)		(96%)	A17	(0.4 %)
Urinary bladder	(50)	(10070)		(3070)		(94%)
Calculus gross observation		(2%)	(50)		(50)	
Calculus micro observation only	1	(270)	1	(2%)		
Hemorrhage	1	(2%)	1	(470)		
Inflammation, acute		(2%) (2%)				
Inflammation, chronic			•	(9%)		
Inflammation, chronic active		(2%) (2%)		(2%)	•	(40%)
Transitional epithelium, hyperplasia	1	(2%)		(2%)	2	(4%)
rransimunai epimenum, nyperpiasia			1	(2%)		

APPENDIX B

SUMMARY OF LESIONS IN FEMALE RATS IN THE TWO-YEAR INHALATION STUDY OF TOLUENE

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TABLE B1. SUMMARY OF THE INCIDENCE OF NEOPLASMS IN FEMALE RATS IN THE TWO-YEAR INHALATION STUDY OF TOLUENE

	Chambe	er Control	600 r	pm	1,200	ppm
Animals initially in study	60		60		60	
Animals removed	60		60		60	
Animals examined histopathologically	50		50		50	
ALIMENTARY SYSTEM						
Esophagus	(49)		(50)		(49)	
Carcinoma, metastatic, thyroid gland		(2%)				
Intestine small, duodenum	(50)		(49)		(49)	
Leukemia mononuclear	(FO)			(4%)	(40)	
Intestine small, ileum	(50)		(49)	(40)	(49)	
Leukemia mononuclear	(50)			(4%)	(40)	
Intestine small, jejunum	(50)		(49)	(00)	(49)	
Leukemia mononuclear	(EA)			(2%)	(EA)	
Liver	(50)	(40%)	(50)	(COL)	(50)	
Hepatocellular adenoma Leukemia mononuclear		(4%) (34%)	-	(6%) (32%)	10	(20%)
	*(50)	(J470)	*(50)	(3270)	*(50)	(2070)
Mesentery Leukemia mononuclear	(00)		(00)			(2%)
Pancreas	(50)		(49)		(50)	(& NO)
Leukemia mononuclear		(6%)	(40)			(2%)
Salivary glands	(50)	(3707	(50)		(50)	(a /v)
Leukemia mononuclear		(2%)		(2%)		(2%)
Stomach, forestomach	(50)	(2 %)	(50)	(2,0)	(50)	(2,0)
Leukemia mononuclear	,,	(2%)		(4%)		(6%)
Papilloma squamous	-	(=,0)	-	(2,0)		(2%)
Squamous cell carcinoma						(2%)
Stomach, glandular	(50)		(50)		(50)	(=,
Leukemia mononuclear		(6%)		(4%)	* * * * * * * * * * * * * * * * * * * *	(6%)
CARDIOVASCULAR SYSTEM			· ·			
Heart	(50)		(50)		(50)	
Leukemia mononuclear	5	(10%)	4	(8%)	2	(4%)
ENDOCRINE SYSTEM						
Adrenal gland, cortex	(50)		(49)		(49)	
Leukemia mononuclear	14	(28%)	9	(18%)	4	(8%)
Adrenal gland, medulla	(49)		(48)		(49)	
Leukemia mononuclear	10	(20%)	6	(13%)	3	(6%)
Pheochromocytoma, NOS		(2%)				(8%)
Islets, pancreatic	(50)		(49)		(50)	
Carcinoma				(4%)		(2%)
Pituitary gland	(50)		(50)		(50)	
Pars distalis, adenoma	31	(62%)		(54%)	31	(62%)
Pars distalis, craniopharyngioma	_		1	(2%)	_	
Pars distalis, leukemia mononuclear		(6%)			2	(4%)
Pars intermedia, adenoma		(2%)	120		/EA\	
Thyroid gland	(50)	(00)	(50)		(50)	
Leukemia mononuclear		(2%)	0	(16%)	9	(6%)
C-cell, adenoma	7	(4%)	•	(16%)	3	(070)

GENERAL BODY SYSTEM

None

TABLE B1. SUMMARY OF THE INCIDENCE OF NEOPLASMS IN FEMALE RATS IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

	Chambe	er Control	600 I	ppm	1,200	ppm
GENITAL SYSTEM						
	(49)		(50)		(44)	
Clitoral gland Adenoma		(4%)		(8%)		(7%)
	_	(470)	_	(070)	_	(170)
Ovary	(50)		(50)		(50)	(40)
Granulosa cell tumor malignant		(0~)		(OW)		(4%)
Leukemia mononuclear Uterus		(8%)		(6%)		(4%)
Adenocarcinoma	(50)		(50)		(50)	(04)
Adenocarcinoma Adenoma		(90)			1	(2%)
		(2%)		(00)	•	(401)
Leukemia mononuclear		(6%)		(2%)		(4%)
Endometrium, polyp stromal Endometrium, sarcoma stromal		(8%) (2%)	5	(10%)		(4%) (2%)
HEMATOPOIETIC SYSTEM Blood	*(50)		*(50)		*(50)	
Leukemia mononuclear	(00)			(2%)	*(50)	
Bone marrow	(40)			(470)	/EA\	
Leukemia mononuclear	(49)	(10%)	(50)	(90%)	(50)	(40)
		(10%)		(8%)	_	(4%)
Lymph node Leukemia mononuclear	(50)	(20)	(50)	(40%)	(50)	
		(2%)		(4%)		
Mediastinal, leukemia mononuclear		(2%)		(4%)		
Mediastinal, lymphoma malignant histio	•	(004)		(2%)	_	
Mesenteric, leukemia mononuclear		(30%)		(28%)		(16%)
Lymph node, mandibular	(47)	(54)	(48)		(42)	
Carcinoma, metastatic, thyroid gland		(2%)				
Leukemia mononuclear	-	(6%)	_	(10%)		(2%)
Spleen	(50)		(50)		(50)	
Leukemia mononuclear	17	(34%)	16	(32%)	10	(20%)
Thymus	(46)		(46)		(46)	
Leukemia mononuclear	6	(13%)	5	(11%)	2	(4%)
Thymoma benign	1	(2%)				
NTEGUMENTARY SYSTEM						
Mammary gland	(50)		(50)		(50)	
Adenocarcinoma		(4%)		(2%)	** * * *	(12%)
Adenoma		(4%)		(2%)		(2%)
Fibroadenoma		(26%)	_	(8%)	_	(14%)
Skin	(50)	(20 %)	(50)	(3 70)	(50)	(1470)
Keratoacanthoma	(30)		(00)		,,	(2%)
Subcutaneous tissue, fibroma						(2%)
Subcutaneous tissue, lipoma					_	(2%)
Subcutaneous tissue, neurofibrosarcoma				(20%)	1	(470)
Subcutaneous cissue, neuronorosarcoma			1	(2%)		
MUSCULOSKELETAL SYSTEM None						
NERVOUS SYSTEM	<u></u>					
	(50)		(50)		(49)	
Brain	(50)					
Brain Astrocytoma malignant	(50) 1	(2%)	(00)		(20)	

TABLE B1. SUMMARY OF THE INCIDENCE OF NEOPLASMS IN FEMALE RATS IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

	Chamb	er Control	600 p	ppm	1,200	ppm
RESPIRATORY SYSTEM						
Lung	(50)		(50)		(50)	
Carcinoma, metastatic, thyroid gland		(2%)	(00)		(00)	
Leukemia mononuclear		(32%)	13	(26%)	6	(12%)
Nose	(49)		(50)	,	(50)	
Mucosa, squamous cell carcinoma	, , ,					(2%)
Trachea	(50)		(50)		(50)	
Carcinoma, metastatic, thyroid gland		(2%)				
Leukemia mononuclear	1	(2%)				
SPECIAL SENSES SYSTEM						
Eye	* (50)		* (50)		*(50)	
Leukemia mononuclear	1	(2%)				
Zymbal gland	*(50)		*(50)		*(50)	
Carcinoma	1	(2%)				
Squamous cell carcinoma			1	(2%)		
URINARY SYSTEM						
Kidney	(50)		(50)		(50)	
Leukemia mononuclear	3	(6%)	4	(8%)	5	(10%)
Sarcoma					1	(2%)
Renal tubule, carcinoma					1	(2%)
Urinary bladder	(50)		(50)		(50)	
Leukemia mononuclear		(12%)	2	(4%)	2	(4%)
Transitional epithelium, papilloma	1	(2%)				
SYSTEMIC LESIONS						
Multiple organs	*(50)		*(50)		*(50)	
Leukemia mononuclear	18	(36%)		(32%)	10	(20%)
Lymphoma malignant histiocytic			1	(2%)		
ANIMAL DISPOSITION SUMMARY					·	
Animals initially in study	60		60		60	
Terminal sacrifice	32		35		30	
Interval sacrifice	10		10		10	
Dead	7		7		6	
Moribund	11		8		14	
TUMOR SUMMARY						
Total animals with primary neoplasms **	46		43		43	
Total primary neoplasms	86		77		84	
Total animals with benign neoplasms	39		37		34	
Total benign neoplasms	60		53		52	
Total animals with malignant neoplasms	21		21		23	
Total malignant neoplasms	25		24		28	
Total animals with secondary neoplasms ***	1					
Total secondary neoplasms	4					
Total animals with neoplasms	_				-	
uncertain benign or malignant	1				4	
Total uncertain neoplasms	1				4	

^{*} Number of animals receiving complete necropsy examinations; all gross lesions including masses examined microscopically.

** Primary tumors: all tumors except secondary tumors

*** Secondary tumors: metastatic tumors or tumors invasive into an adjacent organ

TABLE B2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF FEMALE RATS IN THE TWO-YEAR INHALATION STUDY OF TOLUENE: CHAMBER CONTROL

0 6 2	0 7 1	0 8 2	0 8 4	0 9 0	0 9 2	0 9 2	0 9 2	0 9 2	0 9 4	0 9 7	0 9 7	1 0 0	1 0 2	1 0 2	1 0 3	1 0 4	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5
0 6 9	1 2	1 0 2	1 1 1	0 7 0	0 8 0	9	0 8 6	7 8 1	1 6	0 9 2	1 0 4	0 9 1	7 4	7 3 1	0 7 9	1 9 1	0 6 8	7 7	0 8 1	0 9 7	0 9 8 1	1 4 1	1 1 8	0 6 1
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+: Tissue examined microscopically
: Not examined
-: Present but not examined microscopically
I: Insufficient tissue

M: Missing
A: Autolysis precludes examination
X: Incidence of listed morphology

TABLE B2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF FEMALE RATS: CHAMBER CONTROL (Continued)

								(•	U 111			,														
WEEKS ON STUDY	1 0	1 0	10	1 0	1 0	1 0	1 0	1	1 0	1 0	1	1 0	1 0	1 0	1 0	1 0	1 0	1 0	1 0	1 0	1 0	1 0	1 0	1	1 0	
	5	5	5	5	5	5	5	5	5	5	5	5	5	5	5	5	5	5	5	5	5	5	5	5	5	TOTAL:
CARCASS ID	0 6 5	0 8 7	9	9	1 2 0	0 ⁻ 7 5	8 5	8 8	0 0	1 0 1	1 0 7	0 8	6 2	0 6 6	0 7 1	8	9 5	1 3	6	0 8 2	0 8 4	9	1 0 6	0 9	1 5	TUMORS
	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	
ALIMENTARY SYSTEM Esophagus	+	+	+	+	*	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	М	+	+	+	+	49
Carcinoma, metastatic, thyroid gland Intestine large	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Intestine large, cecum	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Intestine large, colon Intestine large, rectum	+ +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 50
Intestine small		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Intestine small, duodenum	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Intestine small, ileum Intestine small, jejunum	++	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	++	+	+	+	++	+	50 50
Liver	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	÷	÷	÷	+	+	÷	+	+	50
Hepatocellular adenoma Leukemia mononuclear			v		х		x		x		X		v			v										2 17
Mesentery			X		А		х		А				X			X						X				17
Pancreas	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Leukemia mononuclear Salivary glands		_	1		+																					3 50
Leukemia mononuclear		-			X	~	+	_	_	+	+	_	+	+	+	+	+	+	+	+	+	+	+	+	_	1 1
Stomach	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Stomach, forestomach Leukemia mononuclear	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Stomach, glandular Leukemia mononuclear	+	+	+	+	*	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
CARDIOVASCULAR SYSTEM	·				-																					
Blood vessel Heart	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 50
Leukemia mononuclear	İ		_		7		_	_	_	_	_		Ŧ	Ŧ	_	_	*	_	_	_	_	7	_		т.	5
ENDOCRINE SYSTEM	·																									
Adrenal gland Adrenal gland, cortex	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 50
Leukemia mononuclear	'	,		-	X	-	X	т	X	_	т	т	Τ.	т	т	x			_		-	X	т	т	7	14
Adrenal gland, medulla	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	X +	+	+	+	+	+	+	+	+	+	49
Leukemia mononuclear Pheochromocytoma, NOS					Х		Х		X							X										10
Islets, pancreatic	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Parathyroid gland	+	+	+	+	M	+	+	+	+	+	+	+	+	+	+	+	÷	+	+	+	+	+	+	M		42
Pituitary gland Pars distalis, adenoma	+	+	+	*	X	*	+	+	*	*X	+	*	+	*X	+ X	*	+	*	+	+ X	*	*	+	*	+	50 31
Pars distalis, leukemia mononuclear	1			Α.	Ŷ	Λ.			Λ.	^		^		Λ.	^	^		Λ		Λ	Λ	Λ		Λ.		31
Pars intermedia, adenoma																										1
Thyroid gland Leukemia mononuclear	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 1
C-cell, adenoma																			x							2
C-cell, carcinoma					X																					2
GENERAL BODY SYSTEM None	-																									
GENITAL SYSTEM	-											_			-											
Clitoral gland Adenoma	+	+ X	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	*	+	+	+	49
Ovary	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Leukemia mononuclear					X																					4
Uterus Adenoma	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Leukemia mononuclear					X																	Λ				1 3
Endometrium, polyp stromal												X				X									X	4
Endometrium, sarcoma stromal Vagina	Х				+																					1 4
· ag.ma	-				+																					4

TABLE B2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF FEMALE RATS: CHAMBER CONTROL (Continued)

WEEKS ON STUDY	0 6 2	7 1	0 8 2	0 8 4	9 0	0 9 2	0 9 2	0 9 2	0 9 2	0 9 4	0 9 7	0 9 7	0	0 2	1 0 2	1 0 3	0 4	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5
CARCASS ID	0 6 9 1	1 2 1	1 0 2 1	1 1 1 1	0 7 0 1	0 8 0 1	0 9 4 1	0 8 6 1	0 7 8 1	1 6 1	0 9 2 1	1 0 4 1	0 9 1	0 7 4 1	0 7 3 1	0 7 9	1 9 1	0 6 8 1	0 7 7 1	0 8 1 1	9 7 1	0 9 8 1	1 1 4 1	1 1 8 1	6 1 1
HEMATOPOIETIC SYSTEM Bone marrow Leukemia mononuclear Lymph node Leukemia mononuclear	+ +	M +	+	+	+	+	+	+	+	+	+ X +	+	+ X +	+	+	+	+ X +	+	+	+	+	+	+	+	+
Mediastinal, leukemia mononuclear Mesenteric, leukemia mononuclear Lymph node, mandibular Carcinoma, metastatic, thyroid gland Leukemia mononuclear	М	X	+	+	X M	х + х	X M	+	+	X +	X + X	+	X X +	+	+	+	X	X +	X +	+	+	+	+	+	+
Spleen Leukemia mononuclear Thymus Leukemia mononuclear Thymoma benign	++	* * *	+ M	+	* X +	X M	* * * X	+	+	* *	X M	+	* * *	+ M	+	+	* * *	* *	* *	+	+	+	+	+	+
INTEGUMENTARY SYSTEM Mammary gland Adenocarcinoma Adenoma Fibroadenoma Skin	+	+	+	+	+	+	+	+	+	+ X +	+	+	+	+	+ X X +	+	+ X +	+	+	+ X +	+	x x	+	+	+ X X +
MUSCULOSKELETAL SYSTEM Bone	+	M	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
NERVOUS SYSTEM Brain Astrocytoma malignant Leukemia mononuclear	+	+	+	+	+	+	+ X	+	+	+	+ X	+	+	+	+	+	+	+	+ X	+	+	+	+	+	+
RESPIRATORY SYSTEM Lung Carcinoma, metastatic, thyroid gland Leukemia mononuclear Nose Trachea Carcinoma, metastatic, thyroid gland Leukemia mononuclear	+ + +	+ X M +	+ + +	+ + +	+ X + +	+ X + +	+ X + +	+ + +	+ + +	+ X + +	+ X + +	+ + +	+ + +	+ + +	+ + +	+ + +	+ X + +	+ X + +	+ X + +	+++	+++	+ X + +	+ + +	+ + +	+ + +
SPECIAL SENSES SYSTEM Eye Leukemia mononuclear Zymbal gland Carcinoma			+			*	*			-	+		+				•								
URINARY SYSTEM Kidney Leukemia mononuclear Urinary bladder Leukemia mononuclear Transitional epithelium, papilloma	+ +	+	+	+	+	+	* * *	+	+	+ X +	+ *	+	+ *	+	+	+	+ *	+	+	+	+	+	+	+	+ +

TABLE B2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF FEMALE RATS: CHAMBER CONTROL (Continued)

WEEKS ON STUDY	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	TOTAL:												
CARCASS ID	0 6 5 1	0 8 7 1	0 9 0 1	0 9 3 1	1 2 0 1	0 7 5 1	0 8 5 1	0 8 8 1	0 0 1	1 0 1 1	1 0 7 1	1 0 8 1	0 6 2 1	0 6 6	0 7 1 1	0 8 3 1	0 9 5 1	1 3 1	0 6 4 1	0 8 2 1	0 8 4 1	9 9 1	0 6 1	0 9 1	1 1 5 1	TISSUES
HEMATOPOIETIC SYSTEM Bone marrow Leukemia mononuclear Lymph node	+ +	+	++	+	* *	+	+	+	+	+	+	+	+	+	+	* X +	+	+	+	+	+	+	+	+	+	49 5 50
Leukemia mononuclear Mediastinal, leukemia mononuclear Mesenteric, leukemia mononuclear Lymph node, mandibular Carcinoma, metastatic, thyroid gland	+	+	X +	+	X + X	+	X +	+	X	+	+	+	+	+	+	X +	+	+	+	+	+	+	+	+	+	1 15 47 1
Leukemia mononuclear Spleen Leukemia mononuclear Thymus Leukemia mononuclear Thymoma benign	+	+	* X +	+	* * *	+	+ X + X	+	* *	+	+	+	* *	+	+	X + X + X	+	+	+	+	+	+ X +	+	+	+	3 50 17 46 6
INTEGUMENTARY SYSTEM Mammary gland Adenocarcinoma Adenoma	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	*	+	+	+	+	+	+	+	+	+	50 2 2
Fibroadenoma Skin	+	+	+	+	+	X	X +	+	+	+	+	X	X	+	X	+	+	X	+	X	+	+	+	+	+	13 50
MUSCULOSKELETAL SYSTEM Bone	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	49
NERVOUS SYSTEM Brain Astrocytoma malignant Leukemia mononuclear	+	+	+	+	+ X	+	+	+	+	+	+	+	+	+	+ X	+	+	+	+	+	+	+	+	+	+	50 1 4
RESPIRATORY SYSTEM Lung Carcinoma, metastatic, thyroid gland Leukemia mononuclear Nose Trachea Carcinoma, metastatic, thyroid gland Leukemia mononuclear	+ + +	+ + +	+ X + +	+ + +	* X X + + X X	+ + +	+ X + +	+ + +	+ X + +	+ + +	+ + +	+ + +	+ + +	+ + +	+ + +	+ X + +	+ + +	+ + +	+ + +	+ + +	+ + +	+ X + +	+ + +	+ + +	+ + +	50 1 16 49 50 1
SPECIAL SENSES SYSTEM Eye Leukemia mononuclear Zymbal gland Carcinoma						•				•					+					•					_	5 1 1 1
URINARY SYSTEM Kidney Leukemia mononuclear Urinary bladder Leukemia mononuclear Transitional epithelium, papilloma	+ +	+	+	+	* * *	+	+	+	+	+	+	+	+ + x	+	+	+	+	+	+	+	+	+ *	+	+	+	50 3 50 6 1

TABLE B2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF FEMALE RATS IN THE TWO-YEAR INHALATION STUDY OF TOLUENE: 600 ppm

	TIVITAL	47	11,	<i>7</i> 14	5,	. 01	,,	O.		UL	C E	114 1	٠. ٠	,00	PP	1111										
WEEKS ON STUDY	- 1	0 6 0	0 6 6	0 7 7	0 7 8	0 7 9	0 8 0	0 9 4	0 9 4	0 9 4	0 9 7	0 9 7	.9 .8	9 8	1 0 2	0 3	1 0 5	1 0 5	0 5	0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5
CARCASS ID		2 3 1	1 8 2 1	2 1 5 1	2 3 3 1	9 5 1	1 9 2 1	2 0 7 1	2 0 8 1	0 2 1	2 1 9	2 2 3 1	1 8 1	2 2 7 1	2 2 0 1	2 1 1	1 8 5 1	1 9 4 1	9 6 1	2 0 6 1	1 3 1	2 2 6 1	2 2 8 1	1 8 3 1	9 3 1	2 0 1
ALIMENTARY SYSTEM																										—
Esophagus Intestine large		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	++
Intestine large, cecum	1	Ă	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine large, colon		Ą	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine large, rectum Intestine small		+ A	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine small, duodenum		Ā	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	÷	+	+	+	+	+	+
Leukemia mononuclear Intestine small, ileum		Α	4	+	4	_	+	+	_	+	X	_	+	_	_	+	4	_	_	_	4	+	X	+	+	+
Leukemia mononuclear			т		*	-	•	1	-	,	X		-	-	7	,	*	7	,	7	-	,	X	,	•	,
Intestine small, jejunum Leukemia mononuclear	1	A	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Liver		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	X	+	+	+
Hepatocellular adenoma								_			_					_						_	_			
Leukemia mononuclear Mesentery					X			X			X		X		X	X					X	X	X	_		
Pancreas		M	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Salivary glands Leukemia mononuclear		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Stomach	1	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	X	+	+	+
Stomach, forestomach		+	+	+	+	+	+	+	+	÷	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Leukemia mononuclear Stomach, glandular	1	_	_	_	_	_	_	_	_	_	_	_	X	_	_	_	_	_	_	_	_	_	X +	_	+	+
Leukemia mononuclear		,	,	'		-	•		-	,	,	т	*	т.	•	,	,	,	,	-	,	,	X			
Tooth																										
CARDIOVASCULAR SYSTEM		_			_																					
Blood vessel Heart		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Leukemia mononuclear		+	+	+	+	+	+	+	+	+	*	+	*	+	*	+	+	+	+	+	+	+	+	+	+	+
ASSOCIATION ASSOCIATION																										
ENDOCRINE SYSTEM Adrenal gland		+	+	+	+	+	+	+	4	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Adrenal gland, cortex		÷	+	÷	÷	+	+	÷	÷	÷	÷	+	+	+	+	+	+	+	+	+	+	+	÷	+	+	+
Leukemia mononuclear Adrenal gland, medulla		_	_	_	X	_	_	X	_	М	X +	+	X	_	X	_	_	_	_	_	_	X	X +	_	_	+
Leukemia mononuclear	ļ	т.	_	_	7		т	X	-	141	X		X		X	-	_	-	-			X	X	,	-	
Islets, pancreatic Carcinoma	1	M	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Parathyroid gland	i	+	M	+	+	+	+	+	+	+	M	+	+	+	+	+	+	M	+	+	+	+	+	+	+	M
Pituitary gland		+	* X	*	+	+	+	+	+	*	*	+	+	±	+	*	*	+ X	+	+	+	*	+	+ X	+	+
Pars distalis, adenoma Pars distalis, craniopharyngioma	ļ		х	А		X				А	A	X		X		А	А	А			X	А	X	Α		X
Thyroid gland		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	±	+	+	+	+	+	+	+
C-cell, adenoma C-cell, carcinoma						X			X	X								X	X			X				х
	_																									
GENERAL BODY SYSTEM None																										
GENITAL SYSTEM	-																									
Clitoral gland		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Adenoma		•	•			,	X			,	·		•	•	•	•	·		•	•			•		•	
Ovary Leukemia mononuclear		+	+	+	+	+	+	+	+	+	*X	+	+	+	*	+	+	+	+	+	+	+	*	+	+	+
Uterus	j	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Leukemia mononuclear				v									X									v				
Endometrium, polyp stromal Vagina	-			X	X	X				+				+								X		+		
vagina										+	_			+			_									

TABLE B2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF FEMALE RATS: 600 ppm (Continued)

								`-				-/														
WEEKS ON STUDY	1 0 5																									
CARCASS	2	2	2 3	9	1 9	2	2	2 2	2 2	2 3	1 8	1 8	1 9	2	2 2	2 2	3	1 8	1 8	1 9	2	2	2 2	2 3	2 4	TOTAL: TISSUES TUMORS
ID	2	6	5	8	9	3	4	2 1	9 1	2	6	8	1	7 1	ī 1	4	7	4	9 1	7	0	9	5 1	9	0	
ALIMENTARY SYSTEM	-																									
Esophagus Intestine large	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 50
Intestine large Intestine large, cecum	+	+	+	+	+	+	+	+	+	+	<u>+</u>	+	+	+	+	+	+	+	+	+	+	+	+	+	+	49
Intestine large, colon	+	÷	+	+	+	+	+	Ŧ	+	+	+	Ŧ	+	Ŧ	Ŧ	Ŧ	Ŧ	7	+	Ŧ	Ŧ	Ŧ	4	Ŧ	+	49
Intestine large, rectum	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Intestine small	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	49
Intestine small, duodenum	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	49
Leukemia mononuclear Intestine small, ileum		_	_				_																			49
Leukemia mononuclear		_			_	_	~			_	_	+	Τ.	Ŧ	_	+	7	Τ.	Τ.	+	+	7	_	+	+	2
Intestine small, jejunum	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	49
Leukemia mononuclear																•	•							•		1
Liver	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Hepatocellular adenoma Leukemia mononuclear			X			v	X		v		X		•		v											.3
Mesentery			А			X		X	X		X		Х		X											16
Pancreas	-	+	+	_	_	_	4	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	49
Salivary glands	+	+	+	÷	+	+	+	+	+	+	+	÷	÷	+	+	÷	+	+	+	+	+	+	+	+	+	50
Leukemia mononuclear												,				•		•	•		,				•	i
Stomach	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Stomach, forestomach	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Leukemia mononuclear Stomach, glandular																										2
Leukemia mononuclear Tooth		_	_	+	+	+	+	+	+	+	+	+	+	+	+	+	*	+	+	+	+	+	+	+	+	50 2 1
CARDIOVASCULAR SYSTEM	-																									
Blood vessel	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Heart	+	+	+	+	+	+	+	+	+	+	+	+	*	+	+	+	+	+	+	+	+	+	+	+	+	50
Leukemia mononuclear													Х													4
ENDOCRINE SYSTEM	-																									
Adrenal gland	+	+	M	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	49
Adrenal gland, cortex	+	+	M	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	49
Leukemia mononuclear	1								X				X													9
Adrenal gland, medulla Leukemia mononuclear	+	+	M	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	48
Isiets, pancreatic	+	+	_		4	_	_																			6 49
Carcinoma		X	x	т					_	_	_	+	+	Ŧ	_	+	+	*	-	+	+	+	_	+	-	2
Parathyroid gland	+	+	+	+	+	+	+	+	+	+	+	+	+	+	M	+	M	+	+	+	+	+	+	+	+	44
Pituitary gland	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Pars distalis, adenoma		X				X		X	X	X	X			Х		X						X	X	Х	X	27
Pars distalis, craniopharyngioma			X																							1
Thyroid gland C-cell, adenoma	+	+	+	+	X X	+	+	*	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
C-cell, carcinoma					А			А													X					8 2
JENERAL BODY SYSTEM None	-																									
GENITAL SYSTEM	-																									
Clitoral gland		_	_	_		_	_	.4.		_	- 4	_			.4.	.4.			4			_	_	_	4	50
Adenoma	-	7	7	X	x		~	~	т	~	¥	_	_	•	_	7	7	~	~	+	_	Ŧ	т	+	+	4
Ovary	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Leukemia mononuclear													•			•				,	•					3
Uterus	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Leukemia mononuclear																							_			1
Endometrium, polyp stromal Vagina			_																				X			5
ı mêring	1		+																							4
	_																									1

TABLE B2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF FEMALE RATS: 600 ppm (Continued)

WEEKS ON STUDY	0 6 0	0 6 6	0 7 7	0 7 8	0 7 9	0 8 0	0 9 4	0 9 4	0 9 4	0 9 7	0 9 7	0 9 8	0 9 8	1 0 2	1 0 3	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5
CARCASS ID	2 3 1	1 8 2 1	2 1 5 1	2 3 3 1	1 9 5 1	1 9 2 1	2 0 7 1	2 0 8 1	2 0 2 1	2 1 9	2 2 3 1	1 8 1 1	2 2 7 1	2 2 0 1	2 1 1 1	1 8 5 1	9 4 1	9 6 1	2 0 6 1	2 1 3 1	2 2 6 1	2 2 8 1	1 8 3 1	9 3 1	0 1 1
HEMATOPOIETIC SYSTEM Blood Leukemia mononuclear Bone marrow Leukemia mononuclear Lymph node Leukemia mononuclear Mediastinai, leukemia mononuclear Mediastinai, lymphoma malignant	+ +	+ +	+	* X +	+	+	+ X + X X	+	+	* *	+	+ X + X	++	+	+	+	+	+	+	+	+ *	* X +	+ +	+	+ +
histiccytic Mesenteric, leukemia mononuclear Lymph node, mandibular Leukemia mononuclear Spleen Leukemia mononuclear Thymus Leukemia mononuclear	+ + +	M + +	+ + +	X M + X +	+++	+++	+ X +	+ + +	+ + +	* + * * * + * * * * * * * * * * * * * *	+++	X + X + X +	+++++	X + X +	* + * * * * * * * * * * * * * * * * * *	+++	++++	++++	+ + +	+ * * +	X + X + X M	X + X + X	++++	+++	+ + + +
INTEGUMENTARY SYSTEM Mammary gland Adenocarcinoma Adenoma Fibroadenoma Skin Subcutaneous tissue, neurofibrosarcoma	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+ X +	+	+	+	+	+	+ X +	+	+	+	+
MUSCULOSKELETAL SYSTEM Bone	+	+	+	+	+	+	+	+ '	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
NERVOUS SYSTEM Brain Leukemia mononuclear	+	+	+	+	+	+	*	+	+	*	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
RESPIRATORY SYSTEM Lung Leukemia mononuclear Nose Trachea	+ + +	+ + +	+ + +	+ X + +	+++	+++	* X + +	+ + +	+ + +	* X + +	+ + +	* X + +	+ + +	+ X + +	+ + +	+ + +	+ + +	+ + +	+ + +	+ + +	+ X + +	+ X + +	+ + +	+ + +	+ + +
SPECIAL SENSES SYSTEM Eye Zymbal gland Squamous cell carcinoma	+ X														+								+		
URINARY SYSTEM Kidney Leukemia mononuclear Urinary bladder Leukemia mononuclear	+	+	+	+	+	+	+	+	+	+	+	* X +	+	+ X + X	+	+	+	+	+	+	+	+ X + X	+	++	+ +

TABLE B2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF FEMALE RATS: 600 ppm (Continued)

WEEKS ON STUDY	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	
CARCASS ID	2 1 2 1	2 1 6 1	2 3 5 1	1 9 8 1	1 9 9	2 0 3 1	2 1 4 1	2 2 2 1	2 2 9 1	2 3 2 1	1 8 6 1	1 8 8 1	1 9 1 1	2 1 7 1	2 2 1 1	2 2 4 1	2 3 7 1	1 8 4 1	1 8 9 1	1 9 7 1	2 0 0 1	2 0 9	2 2 5 1	2 3 9 1	2 4 0 1	TOTAL TISSUES TUMORS
HEMATOPOIETIC SYSTEM Blood Leukemia mononuclear Bone marrow Leukemia mononuclear Lymph node Leukemia mononuclear Mediastinal, leukemia mononuclear	+	+	+ +	+	+	+	+ +	+	+	+	+	+	+	+ +	+	+	+	+	+	+	+	+	+	+	+ +	1 1 50 4 50 2 2
Mediastinal, lymphoma malignant histocytic Mesenteric, leukemia mononuclear Lymph node, mandibular Leukemia mononuclear Spleen Leukemia mononuclear Thymus Leukemia mononuclear Thymus Leukemia mononuclear	+ +	+ + +	X + X + X + X	+ + M	+ + +	* + * * * + * * * * * * * * * * * * * *	* + + + +	* + * * + * * + * * * * * * * * * * * *	* + * * + * * + * * * * * * * * * * * *	++++	* + * * * * * * * * * * * * * * * * * *	++++	X + X + X + X	+++	X + X + X	+ + +	+ + +	+ + +	+ + +	+ + +	+ + M	+ + +	+ + M	+++	++++	1 14 48 5 50 16 46 5
INTEGUMENTARY SYSTEM Mammary gland Adenocarcinoma Adenoma Fibroadenoma Skin Subcutaneous tissue, neurofibrosarcoma	+	+	+	+	+	+	+	+	+	+	* *	+	+	+	+ *	+	+	+	+	+ X +	+ X +	+	+	+	+ X +	50 1 1 4 50 1
MUSCULOSKELETAL SYSTEM Bone	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
NERVOUS SYSTEM Brain Leukemia mononuclear	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 2
RESPIRATORY SYSTEM Lung Leukemia mononuclear Nose Trachea	+ + +	+ + +	+ X + +	+ + +	+ + +	+ X + +	+ + +	+ + +	+ X + +	+ + +	* X + +	+ + +	+ X + +	+ + +	* X + +	+ + +	+ + +	+ + +	+ + +	+ + +	+ + +	+ + +	+ + +	+ + +	+ + +	50 13 50 50
SPECIAL SENSES SYSTEM Eye Zymbal gland Squamous cell carcinoma											+					+										4 1 1
URINARY SYSTEM Kidney Leukemia mononuclear Urnary bladder Leukemia mononuclear	+ +	+	+	+	+	+	+	+	+	+	+	+	+	+	* X +	+	+	+	+	+	+	+	+	+	+	50 4 50 2

TABLE B2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF FEMALE RATS IN THE TWO-YEAR INHALATION STUDY OF TOLUENE: 1,200 ppm

														•											
WEEKS ON STUDY	2 4	0 6 1	0 8 1	0 8 1	0 8 1	0 8 4	0 8 7	0 8 7	9	0 9 2	9	0 9 3	0 9 4	9	0 9 6	0 9 7	9	1 0 1	1 0 2	1 0 4	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5
	3	3	3	3	3	3	3	3	3	3	3	3	3	3	3	3	3	3	3	3	3	3	3	3	3
CARCASS ID	0 6 1	3 5 1	2 6 1	0 4 1	2 8 1	3 1 1	1 8 1	4 8 1	5 6 1	0 9 1	1 3 1	1 6 1	4 1 1	5 2 1	4 2 1	0 2 1	0 5 1	4 7 1	2 0 1	3 1	0 1 1	1 9 1	4 1	4 6 1	5 5 1
ALIMENTARY SYSTEM							_																		
Esophagus	+	+	+	+	+	+	+	+	+	M	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine large Intestine large, cecum	+	+	+	+	+	+	++	+	+	M M	++	+	++	+	+	+	+	+	+	+	+	+	+	+	+
Intestine large, colon	+	÷	÷	+	÷	÷	÷	÷	+	M	÷	+	÷	+	+	+	+	+	+	+	+	+	÷	+	+
Intestine large, rectum	+	. +	+	M	+	+	+	+	+	M M	+	+	+	+	+	+	+	+	+	+	+	+	+	+	++
Intestine small Intestine small, duodenum	++	+	+	+	+	+	+	+	+	M	+	++	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine small, ileum	+	+	+	+	+	+	+	+	+	M	+	÷	+	+	+	+	+	+	÷	+	+	+	+	+	+
Intestine small, jejunum Liver	+++	+	+	+	+	++	+	+	+	M	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Leukemia mononuclear	+	+	+	+	+	X	+	X	+	*	+	+	*	X	+	+	+	+	+	+	+	+	*	+	x
Mesentery						••		••		+			••	••											••
Leukemia mononuclear Pancreas	١.		,							X															+
Leukemia mononuclear	+	+	+	+	+	+	+	+	+	*	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Salivary glands	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Leukemia mononuclear Stomach	١.							X																	
Stomach, forestomach	+	+	+	+	+	+	+	+ + X	+	++	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Leukemia mononuclear	· .							X		*	•			X											
Papilloma squamous	l																								
Squamous cell carcinoma Stomach, glandular	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	4	+	+	+	+	+
Leukemia mononuclear		,	,	•			,	X		X				X			,	•		•					·
Tooth	ĺ																						+		+
CARDIOVASCULAR SYSTEM	_						-			-															
Blood vessel	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Heart Leukemia mononuclear	+	+	+	+	+	+	+	+	+	*	+	+	+	*	+	+	+	+	+	+	+	+	+	+	+
										Λ.				Λ											
ENDOCRINE SYSTEM			-																						
Adrenal gland Adrenal gland, cortex	+	+	+	+	+	+	+	+	+	+	M M	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Leukemia mononuclear	+	_	_		_	X	_	x	_	_	TAT	_	X	X		Ψ.	~		7	_	_	т			_
Adrenal gland, medulla	+	+	+	+	+	+	+	+	+	+	M	+	*	+	+	+	+	+	+	+	+	+	+	+	+
Leukemia mononuclear Pheochromocytoma, NOS					x	Х							Х	X		X		x							
Islets, pancreatic	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	7	+	+	+	+	+	+	+	+	+
Carcinoma	'																								
Parathyroid gland Pituitary gland	+	M	+	+	+	+	M	+	+	+	+	+	+	+	M +	+	+	+	+	+	+	+	M	+	+
Pars distalis, adenoma	1	X	*	x	_		_	~	X	7	Ŧ	x	x	X	x	x	x	X X	x	x		X X	*		X
Pars distalis, leukemia mononuclear	1												X	X			•••								
Thyroid gland C-cell, adenoma	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	*	+	+	+	+	+
C-cell, carcinoma	1																			Λ					X
GENERAL BODY SYSTEM														· · · ·											
None																									
GENITAL SYSTEM					-																				
Clitoral gland	+	M	M	+	M	+	+	M	+	M	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Adenoma																		X				1			
Ovary Granulosa cell tumor malignant	+	+	+	+	+	+	+	+	+	+	+	+	×	+	+	+	+	+	+	+	+	+	+	+	+
Leukemia mononuclear										X				X											
Uterus Adenocarcinoma	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Leukemia mononuclear										X				X											
Endometrium, polyp stromal			_							-															
Endometrium, sarcoma stromal Vagina	1		X														4		+						
	1																-		,						

TABLE B2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF FEMALE RATS: 1,200 ppm (Continued)

WEEKS ON STUDY	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	TOTAL:
CARCASS ID	3 5 8 1	3 1 0	3 1 2 1	3 2 2 1	3 2 3 1	3 3 6 1	3 6 0	3 0 3 1	3 1 7 1	3 2 4 1	3 2 5 1	3 3 7 1	3 5 0	3 1 4 1	3 3 4	3 8 1	3 4 5 1	3 4 9	3 5 4	3 1 1	3 3 0 1	3 4 0 1	3 5 3 1	3 5 7	3 5 9	TISSUES
ALIMENTARY SYSTEM Esophagus Intestine large, cecum Intestine large, colon Intestine large, colon Intestine small, duodenum Intestine small, leum Intestine small, leum Intestine small, jejunum Liver Leukemia mononuclear Mesentery Leukemia mononuclear Pancreas Leukemia mononuclear Salivary glands Leukemia mononuclear Stomach Stomach, forestomach Leukemia mononuclear Papilloma squamous Squamous cell carcinoma Stomach, glandular	+++++++++++++++++++++++++++++++++++++++	+++++++++++++++++++++++++++++++++++++++	+++++++++++++++++++++++++++++++++++++++	+++++++++++++++++++++++++++++++++++++++	+++++++++++++++++++++++++++++++++++++++	+++++++++++++++++++++++++++++++++++++++	+++++++++++++++++++++++++++++++++++++++	+++++++++ + + + + + + + + + + + + + +	+++++++++++++++++++++++++++++++++++++++	+++++++++++++++++++++++++++++++++++++++	+++++++++++++++++++++++++++++++++++++++	+++++++X + + + + + + + + + + + + + + +	+++++++++++++++++++++++++++++++++++++++	+++++++++++++++++++++++++++++++++++++++	+++++++++++++++++++++++++++++++++++++++	+++++++++++++++++++++++++++++++++++++++	+++++++++++++++++++++++++++++++++++++++	+++++++++++++++++++++++++++++++++++++++	++++++++++++++++++++++++++++++++++++++	+++++++++++++++++++++++++++++++++++++++	+++++++++++++++++++++++++++++++++++++++	++++++++++++++++++++++++++++++++++++++	+++++++++++++++++++++++++++++++++++++++	++++++++++++++++++++++++++++++++++++++	+++++++++++++++++++++++++++++++++++++++	49 49 49 49 49 49 49 50 10 2 1 50 1 50 3 1 1 50
Leukemia mononuclear Tooth CARDIOVASCULAR SYSTEM Blood vessel Heart	++	++	++	+ +	++	++	+ + +	++	++	++	+ +	+++	++	+ + +	++	+ + +	+	+++	+ +	++	++	+++	++	+	+ + +	50 3 2 50 50
Leukemia mononuclear ENDOCRINE SYSTEM Adrenai gland Adrenai gland, cortex Leukemia mononuclear Adrenai gland, meduila Leukemia mononuclear Pheochromocytoma, NOS Islets, pancreatic Carcinoma Parathyroid gland Pituitary gland Pars distalis, adenoma Pars distalis, leukemia mononuclear Thyroid gland C-cell, adenoma C-cell, carcinoma GENERAL BODY SYSTEM None	+ + + + X +	+ + + + + X +	+ + + + X + + X + +	+ + + + + + * * +	++++ ++X X	+ + + + + M + X + X	+ + + + + +	+++++++	+ + + + + +	+ + + + + X +	+ + + + + M +	+ + + + + X +	+ + + + + * * *	++++++++	++++++++	+ + + + + M +	+ + + + + X +	+ + + + + X +	+ + + + + M +	+++++++	+ + + + + X +	+ + + + + X +	+ + + + + * *	+ + + + + X +	+ + + + X + + X	49 49 49 49 3 4 50 31 41 50 31 2
GENITAL SYSTEM Clitoral gland Adenoma Ovary Granulosa cell tumor malignant Leukemia mononuclear Uterus Adenocarcinoma Leukemia mononuclear Endometrium, polyp stromal Endometrium, sarcoma stromal Vagina	+ +	+ + +	+ + x	+ + x	+ +	M + +	+ + +	+ + +	+ + +	+ + +	+ + +	+ X +	+ + +	+ + +	+ + +	+ + X	+ * X +	+ + +	+ + +	+ + +	+ +	* X + + + +	+ + +	+ +	+ + +	44 3 50 2 2 50 1 2 2 1 3

TABLE B2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF FEMALE RATS: 1,200 ppm (Continued)

Some marrow						(0	On	L111	ueu	.,																
CARCASS ID 0 3 2 0 2 3 1 4 5 6 0 0 4 2 3 0 1 1 4 5 6 6 6 4 6 6 6 6 6 8 1 8 8 6 8 9 3 6 1 2 2 2 5 7 0 3 1 9 4 6 5 6 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	WEEKS ON STUDY		6			8	0 8 4	8	8	9	9	ğ	9	9	0 9 4	9	9	9		1 0 2	1 0 4		1 0 5			
		6	3	2	4	2	3 1 1	3 1 8 1	3 4 8 1			3 1 3 1	1	3 4 1 1		2		•	3 4 7 1				3 1 9	3 4 4	3 4 6 1	5
	Lymph node Mesenteric, leukemia mononuclear Lymph node, mandibular	+ + +	+ + +	+ + M	+ + M		*		* X +	+ + M	X		+ + +	+ * M	+ * *	+ + +	+ + +	+ + +	+ + +	+ + +	+ + +	+ + +	+ + +	+ * *	+ + +	+ * *
Mammary gland + + + + + + + + + + + + + + + + + + +	Spleen Leukemia mononuclear Thymus	+	+	+	+	+	* *		*	+	X	+	+	+	+	+	+ M	+	+	+	+	+	+		+ M	
Subcutaneous tissue, fibroma X MUSCULOSKELETAL SYSTEM + + + + + + + + + + + + + + + + + + +	Adenoma Fibroadenoma Skin	+	+	+ X +	* *	+	+	+	+	+	+	+ X +	+	+	+	+	+	+	* *	+	+	+ X +		* *	+	+ X +
	Subcutaneous tissue, fibroma Subcutaneous tissue, lipoma MUSCULOSKELETAL SYSTEM		х —							x																
+ + + + + + + + + + + + + + + + + + +	Bone	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
+ + + + + + + + + + + + + + + + + + +	Brain	+	+	+	+	+	+	+	+	+	+	+	+	+		+	+	+	+	+	+	+	+	+	+	+
Eye + URINARY SYSTEM	Nose	+ + +	+ + +	+ + +	+ + +	+ + +	* * +	+ + +	* * +	+ + + +	+ X +	+ + +	+ + +	* * + +	+ X +	+ + +	+ + +	+ + +	+ + +	+ + +	+ + +	+ + +	+ + +	+ + +	+ + +	+ + +
Mr. S	SPECIAL SENSES SYSTEM Eye												+								•					
Leukemia mononuclear X X X X X X Sarcoma X Renal tubule, carcinoma	Sarcoma Renal tubule, carcinoma	+	+	+	+	+	*	+	*	+	*	+	+	*	*	+	+	+	+	+	+ X	+	+	+	+	+
Urinary bladder + + + + + + + + + + + + + + + + + + +	Urinary bladder	+	+	+	+	+	+	+	*	+	+	+	+	+	*	+	+	+	+	+	+	+	+	+	+	+

TABLE B2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF FEMALE RATS: 1,200 ppm (Continued)

WEEKS ON STUDY	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	0 5	1 0 5	1 0 5	TOTAL:
CARCASS ID	5 8 1	3 1 0 1	3 1 2 1	3 2 2 1	3 2 3 1	3 6 1	3 6 0 1	3 0 3 1	3 1 7 1	3 2 4 1	3 2 5 1	3 7 1	3 5 0 1	3 1 4 1	3 4 1	3 8 1	3 4 5 1	3 4 9 1	3 5 4 1	1 1 1	3 0 1	3 4 0 1	3 5 3 1	3 5 7 1	3 5 9 1	TISSUES
HEMATOPOIETIC SYSTEM Bone marrow Leukemia mononuclear	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 2 50
Lymph node Mesenteric, leukemia mononuclear Lymph node, mandibular Leukemia mononuclear	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	X +	+	8 42 1
Spieen Leukemia mononuclear Thymus Leukemia mononuclear	++	+	+	+	+	+	+	* X +	+	+	+	* * +	+	+	+	+	+	+ M	+	+	+	+	+	* *	+	50 10 46 2
INTEGUMENTARY SYSTEM Mammary gland Adenocarcinoma Adenoma	+ X	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	*	+	+	+	+	+	+	+	+ X	50 6
Fibroadenoma Skin Keratoacanthoma Subcutaneous tissue, fibroma Subcutaneous tissue, lipoma	+	+	+	+	+	X +	+	+	+	+	+	+	X +	+	+	+	+	+	+	+	+	+ X	+	+	+	50 1 1 1
MUSCULOSKELETAL SYSTEM Bone	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
NERVOUS SYSTEM Brain Leukemia mononuclear	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+		+	+	+	49
RESPIRATORY SYSTEM Lung Leukemia mononuclear Nose	+	+	+	+	+	+	+	*	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 6 50
Mucosa, squamous cell carcinoma Trachea	+	+	+	+	+	+	+	+	+	+	+	+	+	X +	+	+	+	+	+	+	+	+	+	+	+	50
SPECIAL SENSES SYSTEM Eye		+														~									+	3
URINARY SYSTEM Kidney Leukemia mononuclear Sarcoma	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 5 1
Renal tubule, carcinoma Urinary bladder Leukemia mononuclear	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	X	+	+	+	+	+	50 2

TABLE B3. ANALYSIS OF PRIMARY TUMORS IN FEMALE RATS IN THE TWO-YEAR INHALATION STUDY OF TOLUENE

	Chamber Control	600 ррш	1,200 ppm
Adrenal Gland Medulla: Pheochromocyto			
Overall Rates (a)	1/49 (2%)	0/48 (0%)	4/49 (8%)
Adjusted Rates (b)	3.0%	0.0%	10.9%
Terminal Rates (c)	1/33 (3%)	0/34 (0%)	1/30 (3%)
Day of First Observation	729	0/04(070)	564
Life Table Tests (d)	P=0.071	P = 0.494N	P=0.157
Logistic Regression Tests (d)	P=0.086	P=0.494N	P=0.186
Cochran-Armitage Trend Test (d)	P=0.082	F - 0.434M	P=0.100
Fisher Exact Test (d)	P=0.082	P = 0.505N	P = 0.181
litoral Gland: Adenoma			
Overall Rates (a)	2/49 (4%)	4/50 (8%)	3/44 (7%)
Adjusted Rates (b)	6.3%	10.6%	9.7%
Terminal Rates (c)	2/32 (6%)	3/35 (9%)	2/29 (7%)
Day of First Observation	729	554	706
Life Table Tests (d)	P=0.374	P=0.367	P=0.454
Logistic Regression Tests (d)	P = 0.362	P = 0.367 P = 0.345	P=0.448
		F - U.040	r = U.448
Cochran-Armitage Trend Test (d) Fisher Exact Test (d)	P = 0.364	D-0040	D. 0.440
risner Exact lest(q)		P = 0.349	P = 0.449
iver: Hepatocellular Adenoma			
Overall Rates (a)	2/50 (4%)	3/50 (6%)	0/50 (0%)
Adjusted Rates (b)	5.7%	8.6%	0.0%
Terminal Rates (c)	1/33 (3%)	3/35 (9%)	0/30(0%)
Day of First Observation	710	729	
Life Table Tests (d)	P = 0.226N	P = 0.524	P = 0.267N
Logistic Regression Tests (d)	P = 0.226N	P=0.506	P = 0.256N
Cochran-Armitage Trend Test (d)	P = 0.202N	. 0.000	
Fisher Exact Test (d)	1 - 0.20211	P = 0.500	P = 0.247N
fammary Gland: Fibroadenoma			
Overall Rates (e)	13/50 (26%)	4/50 (8%)	7/50 (14%)
Adjusted Rates (b)	35.8%	11.4%	20.4%
Terminal Rates (c)	10/33 (30%)	4/35 (11%)	5/30 (17%)
Day of First Observation			
	655	729	561
Life Table Tests (d)	P = 0.094N	P = 0.013N	P = 0.158N
Logistic Regression Tests (d)	P = 0.084N	P = 0.014N	P = 0.133N
Cochran-Armitage Trend Test (d)	P = 0.067N		
Fisher Exact Test (d)		P = 0.016N	P = 0.105N
lammary Gland: Adenoma or Fibroaden			
Overall Rates (e)	13/50 (26%)	5/50 (10%)	8/50 (16%)
Adjusted Rates (b)	35.8%	13.9%	23.6%
Terminal Rates (c)	10/33 (30%)	4/35 (11%)	6/30 (20%)
Day of First Observation	655	717	561
Life Table Tests (d)	P = 0.162N	P = 0.028N	P = 0.231N
Logistic Regression Tests (d)	P = 0.146N	P = 0.030N	P = 0.203N
Cochran-Armitage Trend Test (d)	P = 0.117N		- 0.2001
Fisher Exact Test (d)		P = 0.033N	P = 0.163N
lammary Gland: Adenocarcinoma			
Overall Rates (e)	2/50 (4%)	1/50 (2%)	6/50 (12%)
Adjusted Rates (b)	6.1%	2.9%	17.7%
Terminal Rates (c)	2/33 (6%)	1/35 (3%)	4/30 (13%)
Day of First Observation	729	729	563
	P=0.055	P = 0.479N	P=0.110
Life Table Tests (d)		A U 1 U.A	7 O'T T O
Life Table Tests (d)			P = 0.190
Life Table Tests (d) Logistic Regression Tests (d) Cochran-Armitage Trend Test (d)	P=0.061 P=0.070	P = 0.479N	P = 0.120

TABLE B3. ANALYSIS OF PRIMARY TUMORS IN FEMALE RATS IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

Mammary Gland: Adenoma or Adenocarcinoma	2/50 (4%) 5.6% 1/35 (3%) 717 P=0.317N P=0.? P=0.339N 6/50 (12%) 16.7%	7/50 (14%) 20.9% 5/30 (17%) 563 P = 0.213 P = 0.?
Overall Rates (e) Adjusted Rates (b) 11.6% Terminal Rates (c) Day of First Observation Life Table Tests (d) Cochran-Armitage Trend Test (d) P=0.151 Logistic Regression Tests (d) P=0.187 Fisher Exact Test (d) Mammary Gland: Adenoma, Fibroadenoma, or Adenocarcinoma Overall Rates (e) Adjusted Rates (b) 38.6% Terminal Rates (c) 11/33 (33%) Day of First Observation Life Table Tests (d) P=0.546N Logistic Regression Tests (d) P=0.546N Logistic Regression Tests (d) P=0.545N Fisher Exact Test (d) P=0.452N Fisher Exact Test (d) P=0.452N Fisher Exact Test (d) Pituitary Gland/Pars Distalis: Adenoma Overall Rates (c) Day of First Observation Life Table Tests (d) P=0.362 Logistic Regression Tests (d) P=0.362 Logistic Regression Tests (d) P=0.505 Cochran-Armitage Trend Test (d) Fisher Exact Test (d) Thyroid Gland: C-Cell Adenoma Overall Rates (a) Adjusted Rates (b) Terminal Rates (c) Day of First Observation Life Table Tests (d) P=0.505 Cochran-Armitage Trend Test (d) Fisher Exact Test (d) Thyroid Gland: C-Cell Adenoma Overall Rates (a) Adjusted Rates (b) 5.4% Terminal Rates (c) Day of First Observation Life Table Tests (d) P=0.379 Logistic Regression Tests (d) P=0.416 Cochran-Armitage Trend Test (d) P=0.429 Fisher Exact Test (d) Thyroid Gland: C-Cell Adenoma or Carcinoma Overall Rates (a) Adjusted Rates (b) 10.5% Terminal Rates (c) Day of First Observation Gestic Regression Tests (d) P=0.379 Logistic Regression Tests (d) P=0.379 Logistic Regression Tests (d) P=0.377 Logistic Regression Tests (d) P=0.414 Cochran-Armitage Trend Test (d) P=0.440 Fisher Exact Test (d) Uterus:	5.6% 1/35 (3%) 717 P=0.317N P=0.? P=0.339N 6/50 (12%) 16.7%	20.9% 5/30 (17%) 563 P=0.213
Adjusted Rates (b) 11.6% Terminal Rates (c) 3/33 (9%) Day of First Observation 710 Life Table Tests (d) P=0.151 Logistic Regression Tests (d) P=0.187 Fisher Exact Test (d) P=0.187 Mammary Gland: Adenoma, Fibroadenoma, or Adenocarcinoma Overall Rates (e) 14/50 (28%) Adjusted Rates (b) 38.6% Terminal Rates (c) 11/33 (33%) Day of First Observation 655 Life Table Tests (d) P=0.452N Fisher Exact Test (d) Pituitary Gland/Pars Distalis: Adenoma Overall Rates (a) 31/50 (62%) Adjusted Rates (b) 70.1% Terminal Rates (c) 20/33 (61%) Day of First Observation 430 Life Table Tests (d) P=0.362 Logistic Regression Tests (d) P=0.505 Cochran-Armitage Trend Test (d) P=0.541 Fisher Exact Test (d) Thyroid Gland: C-Cell Adenoma Overall Rates (a) 2/50 (4%) Adjusted Rates (b) 5.4% Terminal Rates (c) 1/33 (3%) Day of First Observation 655 Life Table Tests (d) P=0.379 Logistic Regression Tests (d) P=0.379 Logistic Regression Test (d) P=0.416 Cochran-Armitage Trend Test (d) P=0.429 Fisher Exact Test (d) Thyroid Gland: C-Cell Adenoma or Carcinoma Overall Rates (a) 4/50 (8%) Adjusted Rates (b) 10.5% Terminal Rates (c) 2/33 (6%) Day of First Observation 642 Life Table Tests (d) P=0.377 Logistic Regression Tests (d) P=0.414 Cochran-Armitage Trend Test (d) P=0.414	5.6% 1/35 (3%) 717 P=0.317N P=0.? P=0.339N 6/50 (12%) 16.7%	5/30 (17%) 563 P=0.213
Terminal Rates (c)	1/35 (3%) 717 P=0.317N P=0.? P=0.339N 6/50 (12%) 16.7%	563 P=0.213
Day of First Observation	717 P=0.317N P=0.? P=0.339N 6/50(12%) 16.7%	563 P=0.213
Life Table Tests (d)	P=0.317N P=0.? P=0.339N 6/50(12%) 16.7%	P = 0.213
Logistic Regression Tests (d)	P=0.? P=0.339N 6/50(12%) 16.7%	
Cochran-Armitage Trend Test (d)	P=0.339N 6/50(12%) 16.7%	1 0
Fisher Exact Test (d) Mammary Gland: Adenoma, Fibroadenoma, or Adenocarcinoma Overall Rates (e)	6/50 (12%) 16.7%	
Overall Rates (e) Adjusted Rates (b) 38.6% Terminal Rates (c) 11/33 (33%) Day of First Observation Life Table Tests (d) Cochran-Armitage Trend Test (d) Pelutitary Gland/Pars Distalis: Adenoma Overall Rates (a) Adjusted Rates (b) Terminal Rates (c) Day of First Observation Life Table Tests (d) Pelutitary Gland/Pars Distalis: Adenoma Overall Rates (a) Adjusted Rates (b) Terminal Rates (c) Day of First Observation Life Table Tests (d) Cochran-Armitage Trend Test (d) Pelutitary Gland: C-Cell Adenoma Overall Rates (a) Adjusted Rates (b) Terminal Rates (c) Day of First Observation Overall Rates (a) Adjusted Rates (b) Support First Observation Life Table Tests (d) Pelutitary Gland: C-Cell Adenoma Overall Rates (a) Adjusted Rates (b) Cochran-Armitage Trend Test (d) Pelutitary Gland: C-Cell Adenoma Overall Rates (c) Day of First Observation Life Table Tests (d) Cochran-Armitage Trend Test (d) Pelutic Regression Tests (d) Cochran-Armitage Trend Test (d) Pelutic Table Tests (d) Pelutic Table Tests (d) Pelutic Test Test Test Test Test Test (d) Pelutic Test Test Test Test Test Test Test Test	16.7%	P = 0.262
Overall Rates (e) Adjusted Rates (b) 38.6% Terminal Rates (c) 11/33 (33%) Day of First Observation Life Table Tests (d) Cochran-Armitage Trend Test (d) Pelutitary Gland/Pars Distalis: Adenoma Overall Rates (a) Adjusted Rates (b) Terminal Rates (c) Day of First Observation Life Table Tests (d) Pelutitary Gland/Pars Distalis: Adenoma Overall Rates (a) Adjusted Rates (b) Terminal Rates (c) Day of First Observation Life Table Tests (d) Cochran-Armitage Trend Test (d) Pelutitary Gland: C-Cell Adenoma Overall Rates (a) Adjusted Rates (b) Terminal Rates (c) Day of First Observation Overall Rates (a) Adjusted Rates (b) Support First Observation Life Table Tests (d) Pelutitary Gland: C-Cell Adenoma Overall Rates (a) Adjusted Rates (b) Cochran-Armitage Trend Test (d) Pelutitary Gland: C-Cell Adenoma Overall Rates (c) Day of First Observation Life Table Tests (d) Cochran-Armitage Trend Test (d) Pelutic Regression Tests (d) Cochran-Armitage Trend Test (d) Pelutic Table Tests (d) Pelutic Table Tests (d) Pelutic Test Test Test Test Test Test (d) Pelutic Test Test Test Test Test Test Test Test	16.7%	
Adjusted Rates (b) 38.6% Terminal Rates (c) 11/33 (33%) Day of First Observation 655 Life Table Tests (d) P=0.546N Logistic Regression Tests (d) P=0.515N Cochran-Armitage Trend Test (d) P=0.452N Fisher Exact Test (d) Pituitary Gland/Pars Distalis: Adenoma Overall Rates (a) 31/50 (62%) Adjusted Rates (b) 70.1% Terminal Rates (c) 20/33 (61%) Day of First Observation 430 Life Table Tests (d) P=0.362 Logistic Regression Tests (d) P=0.505 Cochran-Armitage Trend Test (d) P=0.541 Fisher Exact Test (d) Chyroid Gland: C-Cell Adenoma Overall Rates (a) 2/50 (4%) Adjusted Rates (b) 5.4% Terminal Rates (c) 1/33 (3%) Day of First Observation 655 Life Table Tests (d) P=0.379 Logistic Regression Tests (d) P=0.416 Cochran-Armitage Trend Test (d) P=0.429 Fisher Exact Test (d) Chyroid Gland: C-Cell Adenoma or Carcinoma Overall Rates (a) 4/50 (8%) Adjusted Rates (b) 10.5% Terminal Rates (c) 2/33 (6%) Day of First Observation 642 Life Table Tests (d) P=0.377 Logistic Regression Tests (d) P=0.414 Cochran-Armitage Trend Test (d) P=0.414 Cochran-Armitage Trend Test (d) P=0.440 Fisher Exact Test (d) Uterus: Stromal Polyp	16.7%	13/50 (26%)
Terminal Rates (c) Day of First Observation Life Table Tests (d) Logistic Regression Tests (d) Pe 0.515N Cochran-Armitage Trend Test (d) Pituitary Gland/Pars Distalis: Adenoma Overall Rates (a) Adjusted Rates (b) Terminal Rates (c) Day of First Observation Life Table Tests (d) Cochran-Armitage Trend Test (d) Pe 0.33 (61%) Day of First Observation Life Table Tests (d) Cochran-Armitage Trend Test (d) Pe 0.505 Cochran-Armitage Trend Test (d) Pe 0.541 Fisher Exact Test (d) Cochran-Regression Tests (d) Cochran-Regression Tests (d) Cochran-Armitage Trend Test (d) Cochran-Armitage Tests (d) Day of First Observation Cochran-Armitage Tests (d) Day of First Observation Cochran-Armitage Tests (d) Cochran-Armitage Tests (d) Cochran-Armitage Tests (d) Cochran-Armitage Trend Test (d) Pe 0.379 Logistic Regression Tests (d) Pe 0.416 Cochran-Armitage Trend Test (d) Pe 0.429 Fisher Exact Test (d) Cochran-Armitage Trend Test (d) Pe 0.429 Fisher Exact Test (d) Cochran-Armitage Trend Test (d) Pe 0.429 Fisher Exact Test (d) Cochran-Armitage Trend Test (d) Pe 0.377 Logistic Regression Tests (d) Pe 0.377 Logistic Regression Tests (d) Pe 0.414 Cochran-Armitage Trend Test (d) Pe 0.440 Fisher Exact Test (d) Uterus: Stromal Polyp		36.6%
Day of First Observation	5/35 (14%)	9/30 (30%)
Life Table Tests (d) Logistic Regression Tests (d) Cochran-Armitage Trend Test (d) Pituitary Gland/Pars Distalis: Adenoma Overall Rates (a) Adjusted Rates (b) Day of First Observation Cochran-Armitage Trend Test (d) P=0.362 Logistic Regression Tests (d) Cochran-Armitage Trend Test (d) P=0.505 Cochran-Armitage Trend Test (d) P=0.541 Fisher Exact Test (d) Cochran-Regression Tests (d) Day of First Observation Cochran-Armitage Trend Test (d) Cochran-Armitage Trend Test (d) Cochran-Regression Tests (d) Cochran-Armitage Trend Test (d) Cochran-Armitage Trend Test (d) P=0.416 Cochran-Armitage Trend Test (d) P=0.429 Fisher Exact Test (d) Cochran-Regression Tests (d) Cochran-Armitage Trend Test (d) Cochran-Armitage Trend Test (d) Cochran-Armitage Trend Test (d) Cochran-Armitage Trend Test (d) Fisher Exact Test (d) Uterus: Stromal Polyp		
Logistic Regression Tests (d)	717 D = 0.022N	561 B-0 569
Cochran-Armitage Trend Test (d) Fisher Exact Test (d) Pituitary Gland/Pars Distalis: Adenoma Overall Rates (a) 31/50 (62%) Adjusted Rates (b) 70.1% Terminal Rates (c) 20/33 (61%) Day of First Observation 430 Life Table Tests (d) P=0.362 Logistic Regression Tests (d) P=0.505 Cochran-Armitage Trend Test (d) P=0.541 Fisher Exact Test (d) Fisher Exact Test (d) Thyroid Gland: C-Cell Adenoma 2/50 (4%) Adjusted Rates (b) 5.4% Terminal Rates (c) 1/33 (3%) Day of First Observation 655 Life Table Tests (d) P=0.379 Logistic Regression Tests (d) P=0.416 Cochran-Armitage Trend Test (d) P=0.429 Fisher Exact Test (d) P=0.429 Fisher Exact Test (d) P=0.429 Thyroid Gland: C-Cell Adenoma or Carcinoma Overall Rates (a) 4/50 (8%) Adjusted Rates (b) 10.5% Terminal Rates (c) 2/33 (6%) Day of First Observation 642 Life Table Tests (d) P=0.377 Logistic Regression Tests (d) P=0.414 Cochran-Armitage Trend Test (d) P=0.440 Fisher Exact Test (d) P=0.440 Uterus: Stromal Polyp	P=0.032N	P = 0.568
Fisher Exact Test (d) Pituitary Gland/Pars Distalis: Adenoma Overall Rates (a) 31/50 (62%) Adjusted Rates (b) 70.1% Terminal Rates (c) 20/33 (61%) Day of First Observation 430 Life Table Tests (d) P=0.362 Logistic Regression Tests (d) P=0.505 Cochran-Armitage Trend Test (d) P=0.541 Fisher Exact Test (d) P=0.541 Chyroid Gland: C-Cell Adenoma Overall Rates (a) 2/50 (4%) Adjusted Rates (b) 5.4% Terminal Rates (c) 1/33 (3%) Day of First Observation 655 Life Table Tests (d) P=0.379 Logistic Regression Tests (d) P=0.429 Fisher Exact Test (d) P=0.429 Fisher Exact Test (d) Perminal Rates (c) 2/33 (6%) Day of First Observation 642 Life Table Tests (d) P=0.377 Logistic Regression Tests (d) P=0.377 Logistic Regression Tests (d) P=0.414 Cochran-Armitage Trend Test (d)	P = 0.035N	P = 0.565N
Pituitary Gland/Pars Distalis: Adenoma Overall Rates (a) 31/50 (62%) Adjusted Rates (b) 70.1% Terminal Rates (c) 20/33 (61%) Day of First Observation 430 Life Table Tests (d) P=0.362 Logistic Regression Tests (d) P=0.505 Cochran-Armitage Trend Test (d) P=0.541 Fisher Exact Test (d) Thyroid Gland: C-Cell Adenoma Overall Rates (a) 2/50 (4%) Adjusted Rates (b) 5.4% Terminal Rates (c) 1/33 (3%) Day of First Observation 655 Life Table Tests (d) P=0.379 Logistic Regression Tests (d) P=0.416 Cochran-Armitage Trend Test (d) P=0.429 Fisher Exact Test (d) Thyroid Gland: C-Cell Adenoma or Carcinoma Overall Rates (a) 4/50 (8%) Adjusted Rates (b) 10.5% Terminal Rates (c) 2/33 (6%) Day of First Observation 642 Life Table Tests (d) P=0.377 Logistic Regression Tests (d) P=0.377 Logistic Regression Tests (d) P=0.377 Logistic Regression Tests (d) P=0.414 Cochran-Armitage Trend Test (d) P=0.414 Cochran-Armitage Trend Test (d) P=0.440 Fisher Exact Test (d) Uterus: Stromal Polyp		
Overall Rates (a) 31/50 (62%) Adjusted Rates (b) 70.1% Terminal Rates (c) 20/33 (61%) Day of First Observation 430 Life Table Tests (d) P=0.362 Logistic Regression Tests (d) P=0.505 Cochran-Armitage Trend Test (d) P=0.541 Fisher Exact Test (d) P=0.541 Fisher Exact Test (d) 5.4% Terminal Rates (a) 2/50 (4%) Adjusted Rates (b) 5.4% Terminal Rates (c) 1/33 (3%) Day of First Observation 655 Life Table Tests (d) P=0.379 Logistic Regression Tests (d) P=0.429 Fisher Exact Test (d) P=0.429 Fisher Exact Test (d) P=0.429 First Observation 642 Life Table Tests (d) P=0.377 Logistic Regression Tests (d) P=0.377 Logistic Regression Tests (d) P=0.414 Cochran-Armitage Trend Test (d) P=0.440 Fisher Exact Test (d) P=0.440 Uterus: Stromal Polyp	P = 0.039N	P = 0.500N
Adjusted Rates (b) 70.1% Terminal Rates (c) 20/33 (61%) Day of First Observation 430 Life Table Tests (d) P=0.362 Logistic Regression Tests (d) P=0.505 Cochran-Armitage Trend Test (d) P=0.541 Fisher Exact Test (d) Thyroid Gland: C-Cell Adenoma Overall Rates (a) 2/50 (4%) Adjusted Rates (b) 5.4% Terminal Rates (c) 1/33 (3%) Day of First Observation 655 Life Table Tests (d) P=0.379 Logistic Regression Tests (d) P=0.416 Cochran-Armitage Trend Test (d) P=0.429 Fisher Exact Test (d) Thyroid Gland: C-Cell Adenoma or Carcinoma Overall Rates (a) 4/50 (8%) Adjusted Rates (b) 10.5% Terminal Rates (c) 2/33 (6%) Day of First Observation 642 Life Table Tests (d) P=0.377 Logistic Regression Tests (d) P=0.414 Cochran-Armitage Trend Test (d) P=0.440 Fisher Exact Test (d) Uterus: Stromal Polyp	OH (FO / F 1 ~)	04/20/405
Terminal Rates (c) 20/33 (61%) Day of First Observation 430 Life Table Tests (d) P= 0.362 Logistic Regression Tests (d) P= 0.505 Cochran-Armitage Trend Test (d) P= 0.541 Fisher Exact Test (d) Thyroid Gland: C-Cell Adenoma Overall Rates (a) 2/50 (4%) Adjusted Rates (b) 5.4% Terminal Rates (c) 1/33 (3%) Day of First Observation 655 Life Table Tests (d) P= 0.379 Logistic Regression Tests (d) P= 0.416 Cochran-Armitage Trend Test (d) P= 0.429 Fisher Exact Test (d) Thyroid Gland: C-Cell Adenoma or Carcinoma Overall Rates (a) 4/50 (8%) Adjusted Rates (b) 10.5% Terminal Rates (c) 2/33 (6%) Day of First Observation 642 Life Table Tests (d) P= 0.377 Logistic Regression Tests (d) P= 0.377 Logistic Regression Tests (d) P= 0.414 Cochran-Armitage Trend Test (d) P= 0.414 Cochran-Armitage Trend Test (d) P= 0.440 Fisher Exact Test (d) Uterus: Stromal Polyp	27/50 (54%)	31/50 (62%)
Day of First Observation	62.3%	71.8%
Life Table Tests (d) P = 0.362 Logistic Regression Tests (d) P = 0.505 Cochran-Armitage Trend Test (d) P = 0.541 Fisher Exact Test (d) Thyroid Gland: C-Cell Adenoma Overall Rates (a) 2/50 (4%) Adjusted Rates (b) 5.4% Terminal Rates (c) 1/33 (3%) Day of First Observation 655 Life Table Tests (d) P = 0.379 Logistic Regression Tests (d) P = 0.416 Cochran-Armitage Trend Test (d) P = 0.429 Fisher Exact Test (d) Thyroid Gland: C-Cell Adenoma or Carcinoma Overall Rates (a) 4/50 (8%) Adjusted Rates (b) 10.5% Terminal Rates (c) 2/33 (6%) Day of First Observation 642 Life Table Tests (d) P = 0.377 Logistic Regression Tests (d) P = 0.414 Cochran-Armitage Trend Test (d) P = 0.414 Cochran-Armitage Trend Test (d) P = 0.440 Fisher Exact Test (d) Uterus: Stromal Polyp	19/35 (54%)	18/30 (60%)
Logistic Regression Tests (d) P=0.505 Cochran-Armitage Trend Test (d) P=0.541 Fisher Exact Test (d) Thyroid Gland: C-Cell Adenoma Overall Rates (a) 2/50 (4%) Adjusted Rates (b) 5.4% Terminal Rates (c) 1/33 (3%) Day of First Observation 655 Life Table Tests (d) P=0.379 Logistic Regression Tests (d) P=0.416 Cochran-Armitage Trend Test (d) P=0.429 Fisher Exact Test (d) Thyroid Gland: C-Cell Adenoma or Carcinoma Overall Rates (a) 4/50 (8%) Adjusted Rates (b) 10.5% Terminal Rates (c) 2/33 (6%) Day of First Observation 642 Life Table Tests (d) P=0.377 Logistic Regression Tests (d) P=0.414 Cochran-Armitage Trend Test (d) P=0.440 Fisher Exact Test (d) Uterus: Stromal Polyp	457	421
Logistic Regression Tests (d) P = 0.505 Cochran-Armitage Trend Test (d) P = 0.541 Fisher Exact Test (d) Thyroid Gland: C-Cell Adenoma Overall Rates (a) 2/50 (4%) Adjusted Rates (b) 5.4% Terminal Rates (c) 1/33 (3%) Day of First Observation 655 Life Table Tests (d) P = 0.379 Logistic Regression Tests (d) P = 0.416 Cochran-Armitage Trend Test (d) P = 0.429 Fisher Exact Test (d) Thyroid Gland: C-Cell Adenoma or Carcinoma Overall Rates (a) 4/50 (8%) Adjusted Rates (b) 10.5% Terminal Rates (c) 2/33 (6%) Day of First Observation 642 Life Table Tests (d) P = 0.377 Logistic Regression Tests (d) P = 0.414 Cochran-Armitage Trend Test (d) P = 0.440 Fisher Exact Test (d) Uterus: Stromal Polyp	P = 0.243N	P = 0.381
Cochran-Armitage Trend Test (d) Fisher Exact Test (d) Thyroid Gland: C-Cell Adenoma Overall Rates (a) Adjusted Rates (b) Terminal Rates (c) Day of First Observation Life Table Tests (d) Cochran-Armitage Trend Test (d) Fisher Exact Test (d) Coverall Rates (a) Adjusted Rates (b) Thyroid Gland: C-Cell Adenoma or Carcinoma Overall Rates (a) Adjusted Rates (b) Terminal Rates (c) Day of First Observation Life Table Tests (d) Cochran-Armitage Trend Test (d) Fisher Exact Test (d) Cochran-Rates (a) Adjusted Rates (b) Terminal Rates (c) Day of First Observation Life Table Tests (d) Cochran-Armitage Trend Test (d) Fisher Exact Test (d) Cochran-Armitage Trend Test (d) Fisher Exact Test (d) Cochran-Armitage Trend Test (d) Fisher Exact Test (d) Cochran-Stromal Polyp	P = 0.275N	P = 0.540
Fisher Exact Test (d) Thyroid Gland: C-Cell Adenoma Overall Rates (a) 2/50 (4%) Adjusted Rates (b) 5.4% Terminal Rates (c) 1/33 (3%) Day of First Observation 655 Life Table Tests (d) P=0.379 Logistic Regression Tests (d) P=0.416 Cochran-Armitage Trend Test (d) P=0.429 Fisher Exact Test (d) Thyroid Gland: C-Cell Adenoma or Carcinoma Overall Rates (a) 4/50 (8%) Adjusted Rates (b) 10.5% Terminal Rates (c) 2/33 (6%) Day of First Observation 642 Life Table Tests (d) P=0.377 Logistic Regression Tests (d) P=0.414 Cochran-Armitage Trend Test (d) P=0.440 Fisher Exact Test (d) Uterus: Stromal Polyp		
Overall Rates (a) 2/50 (4%) Adjusted Rates (b) 5.4% Terminal Rates (c) 1/33 (3%) Day of First Observation 655 Life Table Tests (d) P=0.379 Logistic Regression Tests (d) P=0.416 Cochran-Armitage Trend Test (d) P=0.429 Fisher Exact Test (d) P=0.429 Chyroid Gland: C-Cell Adenoma or Carcinoma 4/50 (8%) Overall Rates (a) 4/50 (8%) Adjusted Rates (b) 10.5% Terminal Rates (c) 2/33 (6%) Day of First Observation 642 Life Table Tests (d) P=0.377 Logistic Regression Tests (d) P=0.414 Cochran-Armitage Trend Test (d) P=0.440 Fisher Exact Test (d) P=0.440	P = 0.272N	P=0.582N
Overall Rates (a) 2/50 (4%) Adjusted Rates (b) 5.4% Terminal Rates (c) 1/33 (3%) Day of First Observation 655 Life Table Tests (d) P=0.379 Logistic Regression Tests (d) P=0.416 Cochran-Armitage Trend Test (d) P=0.429 Fisher Exact Test (d) P=0.429 Thyroid Gland: C-Cell Adenoma or Carcinoma Overall Rates (a) 4/50 (8%) Adjusted Rates (b) 10.5% Terminal Rates (c) 2/33 (6%) Day of First Observation 642 Life Table Tests (d) P=0.377 Logistic Regression Tests (d) P=0.414 Cochran-Armitage Trend Test (d) P=0.440 Fisher Exact Test (d) P=0.440		
Adjusted Rates (b) 5.4% Terminal Rates (c) 1/33 (3%) Day of First Observation 655 Life Table Tests (d) P=0.379 Logistic Regression Tests (d) P=0.416 Cochran-Armitage Trend Test (d) P=0.429 Fisher Exact Test (d) Thyroid Gland: C-Cell Adenoma or Carcinoma Overall Rates (a) 4/50 (8%) Adjusted Rates (b) 10.5% Terminal Rates (c) 2/33 (6%) Day of First Observation 642 Life Table Tests (d) P=0.377 Logistic Regression Tests (d) P=0.414 Cochran-Armitage Trend Test (d) P=0.440 Fisher Exact Test (d) Uterus: Stromal Polyp	8/50 (16%)	3/50 (6%)
Terminal Rates (c)	20.0%	9.7%
Day of First Observation 655 Life Table Tests (d) P = 0.379 Logistic Regression Tests (d) P = 0.416 Cochran-Armitage Trend Test (d) P = 0.429 Fisher Exact Test (d) P = 0.429 Thyroid Gland: C-Cell Adenoma or Carcinoma Overall Rates (a) 4/50 (8%) Adjusted Rates (b) 10.5% Terminal Rates (c) 2/33 (6%) Day of First Observation 642 Life Table Tests (d) P = 0.377 Logistic Regression Tests (d) P = 0.414 Cochran-Armitage Trend Test (d) P = 0.440 Fisher Exact Test (d) P = 0.440	5/35 (14%)	2/30 (7%)
Life Table Tests (d) P = 0.379 Logistic Regression Tests (d) P = 0.416 Cochran-Armitage Trend Test (d) P = 0.429 Fisher Exact Test (d) Thyroid Gland: C-Cell Adenoma or Carcinoma Overall Rates (a) 4/50 (8%) Adjusted Rates (b) 10.5% Terminal Rates (c) 2/33 (6%) Day of First Observation 642 Life Table Tests (d) P = 0.377 Logistic Regression Tests (d) P = 0.414 Cochran-Armitage Trend Test (d) P = 0.440 Fisher Exact Test (d) Uterus: Stromal Polyp	553	723
Logistic Regression Tests (d) P = 0.416 Cochran-Armitage Trend Test (d) P = 0.429 Fisher Exact Test (d) Phyroid Gland: C-Cell Adenoma or Carcinoma Overall Rates (a) 4/50 (8%) Adjusted Rates (b) 10.5% Terminal Rates (c) 2/33 (6%) Day of First Observation 642 Life Table Tests (d) P = 0.377 Logistic Regression Tests (d) P = 0.414 Cochran-Armitage Trend Test (d) Fisher Exact Test (d) Uterus: Stromal Polyp	P=0.061	P=0.457
Cochran-Armitage Trend Test (d) P=0.429 Fisher Exact Test (d) P=0.429 Thyroid Gland: C-Cell Adenoma or Carcinoma Overall Rates (a) 4/50 (8%) Adjusted Rates (b) 10.5% Terminal Rates (c) 2/33 (6%) Day of First Observation 642 Life Table Tests (d) P=0.377 Logistic Regression Tests (d) P=0.414 Cochran-Armitage Trend Test (d) P=0.440 Fisher Exact Test (d) P=0.440	P = 0.049	P = 0.470
Fisher Exact Test (d) Phyroid Gland: C-Cell Adenoma or Carcinoma Overall Rates (a) 4/50 (8%) Adjusted Rates (b) 10.5% Terminal Rates (c) 2/33 (6%) Day of First Observation 642 Life Table Tests (d) P=0.377 Logistic Regression Tests (d) P=0.414 Cochran-Armitage Trend Test (d) P=0.440 Fisher Exact Test (d) Uterus: Stromal Polyp	F = 0.049	1-0.410
Thyroid Gland: C-Cell Adenoma or Carcinoma	D 0.040	D - 0 500
Overall Rates (a) 4/50 (8%) Adjusted Rates (b) 10.5% Terminal Rates (c) 2/33 (6%) Day of First Observation 642 Life Table Tests (d) P=0.377 Logistic Regression Tests (d) P=0.414 Cochran-Armitage Trend Test (d) P=0.440 Fisher Exact Test (d)	P = 0.046	P = 0.500
Adjusted Rates (b) 10.5% Terminal Rates (c) 2/33 (6%) Day of First Observation 642 Life Table Tests (d) P=0.377 Logistic Regression Tests (d) P=0.414 Cochran-Armitage Trend Test (d) P=0.440 Fisher Exact Test (d)	10/50 (20%)	5/50 (10%)
Terminal Rates (c) 2/33 (6%) Day of First Observation 642 Life Table Tests (d) P = 0.377 Logistic Regression Tests (d) P = 0.414 Cochran-Armitage Trend Test (d) P = 0.440 Fisher Exact Test (d) Uterus: Stromal Polyp		16.1%
Day of First Observation 642 Life Table Tests (d) P=0.377 Logistic Regression Tests (d) P=0.414 Cochran-Armitage Trend Test (d) P=0.440 Fisher Exact Test (d) Uterus: Stromal Polyp	25.3%	
Life Table Tests (d) Logistic Regression Tests (d) Cochran-Armitage Trend Test (d) Fisher Exact Test (d) Uterus: Stromal Polyp	7/35 (20%)	4/30 (13%)
Logistic Regression Tests (d) P=0.414 Cochran-Armitage Trend Test (d) P=0.440 Fisher Exact Test (d) Uterus: Stromal Polyp	553	723
Cochran-Armitage Trend Test (d) P=0.440 Fisher Exact Test (d) Uterus: Stromal Polyp	P=0.097	P = 0.443
Fisher Exact Test (d) Uterus: Stromal Polyp	P = 0.075	P = 0.463
Jterus: Stromał Polyp		
	P = 0.074	P = 0.500
Overall Rates (e) 4/50 (8%)		
	5/50 (10%)	2/50 (4%)
Adjusted Rates (b) 11.4%	11.6%	6.7%
Terminal Rates (c) 3/33 (9%)	2/35 (6%)	2/30 (7%)
Day of First Observation 675	537	729
Life Table Tests (d) P=0.319N	P=0.517	P = 0.387N
Logistic Regression Tests (d) $P = 0.257N$	0.0 . 1	P = 0.378N
Cochran-Armitage Trend Test (d) P=0.283N	P = 0.526	1 -0.01014
Fisher Exact Test (d)	P = 0.526	P = 0.339N

TABLE B3. ANALYSIS OF PRIMARY TUMORS IN FEMALE RATS IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

	Chamber Control	600 ppm	1,200 ppm
Hematopoietic System: Mononuclear I	.eukemia	-	···
Overall Rates (e)	18/50 (36%)	16/50 (32%)	10/50 (20%)
Adjusted Rates (b)	42.6%	38.6%	26.4%
Terminal Rates (c)	10/33 (30%)	10/35 (29%)	5/30 (17%)
Day of First Observation	491	540	584
Life Table Tests (d)	P = 0.105N	P = 0.376N	P = 0.123N
Logistic Regression Tests (d)	P = 0.051N	P = 0.418N	P = 0.056N
Cochran-Armitage Trend Test (d)	P = 0.050N		
Fisher Exact Test (d)	2 2.30011	P = 0.417N	P = 0.059N

⁽a) Number of tumor-bearing animals/number of animals examined microscopically at the site

⁽b) Kaplan-Meier estimated tumor incidences at the end of the study after adjusting for intercurrent mortality

⁽c) Observed tumor incidence in animals killed at the end of the study

⁽d) Beneath the control incidence are the P values associated with the trend test. Beneath the dosed group incidence are the P values corresponding to pairwise comparisons between that dosed group and the controls. The life table analysis regards tumors in animals dying prior to terminal kill as being (directly or indirectly) the cause of death. The logistic regression test regards these lesions as nonfatal. The Cochran-Armitage and Fisher exact tests compare directly the overall incidence rates. A negative trend or lower incidence in a dosed group than in controls is indicated by (N).

⁽e) Number of tumor-bearing animals/number of animals examined grossly at the site

TABLE B4a. HISTORICAL INCIDENCE OF NOSE OR NASAL CAVITY SQUAMOUS CELL TUMORS IN FEMALE F344/N RATS RECEIVING NO TREATMENT (a)

Historical Incidence for Chamber Controls at Battelle Pacific Northwest Laboratories

0/349

Overall Historical Incidence for Untreated Controls in NTP Studies

0/1,643

(a) Data as of May 12, 1988, for studies of at least 104 weeks

TABLE B4b. HISTORICAL INCIDENCE OF KIDNEY SARCOMAS OR TUBULAR CELL TUMORS IN FEMALE F344/N RATS RECEIVING NO TREATMENT (a)

Study	Incidence of Adenomas or Adenocarcinomas in Controls	
Historical Incidence for Chamber C	ontrols at Battelle Pacific Northwest Laboratories	
Propylene oxide	(b) 1/50	
Methyl methacrylate	0/50	
Propylene	0/47	
1,2-Epoxybutane	0/50	
Dichloromethane	0/50	
Tetrachloroethylene	0/50	
Bromoethane	0/50	
TOTAL	1/347 (0.3%)	
SD(c)	0.76%	
Range (d)		
High	1/50	
Low	0/50	
Overall Historical Incidence for Unt	reated Controls in NTP Studies	
TOTAL	(e) 2/1,639 (0.1%)	
SD(c)	0.49%	
Range (d)		
High	1/49	
Low	0/50	

⁽a) Data as of May 12, 1988, for studies of at least 104 weeks

⁽b) Tubular cell adenocarcinoma

⁽c) Standard deviation

⁽d) Range and SD are presented for groups of 35 or more animals.

⁽e) Includes one tubular cell adenoma and one adenocarcinoma, NOS; no renal sarcomas have been observed.

TABLE B4c. HISTORICAL INCIDENCE OF STOMACH SQUAMOUS CELL TUMORS IN FEMALE F344/N RATS RECEIVING NO TREATMENT (a)

Study	Incidence of Papillomas or Carcinomas in Controls	
Historical Incidence for Chamber Con	trols at Battelle Pacific Northwest Laboratories	
Propylene oxide	0/49	
Methyl methacrylate	0/50	
Propylene	0/48	
1,2-Epoxybutane	0/50	
Dichloromethane	0/50	
Tetrachloroethylene	0/49	
Bromoethane	0/48	
TOTAL	0/344	
SD(b)	0.00%	
Range (c)		
High	0/50	
Low	0/50	
Overall Historical Incidence for Untre	ated Controls in NTP Studies	
TOTAL	(d) 3/1,623 (0.2%)	
SD(b)	0.59%	
Range(c)		
High	1/49	
Low	0/50	
 ·		

⁽a) Data as of May 12, 1988, for studies of at least 104 weeks (b) Standard deviation $\,$

⁽c) Range and SD are presented for groups of 35 or more animals.
(d) Includes two squamous cell papillomas and one squamous cell carcinoma

TABLE B5. SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN FEMALE RATS IN THE TWO-YEAR INHALATION STUDY OF TOLUENE

	Chamb	er Control	600 p	opm	1,200	ppm
Animals initially in study	60		60		60	
Animals removed	60		60		60	
animals examined histopathologically	50		50		50	
LLIMENTARY SYSTEM						
Esophagus	(49)		(50)		(49)	
Inflammation, chronic				(2%)		
Intestine large, cecum	(50)		(49)		(49)	
Inflammation, acute	-	(2%)		(4%)		
Intestine large, colon	(50)		(49)	(O~)	(49)	
Inflammation, acute	_	(100)		(2%)	0	(400)
Parasite metazoan Intestine large, rectum	_	(10%)		(2%)		(4%)
Parasite metazoan	(50)	(8%)	(50)	(100)	(48)	
Ulcer	4	(870)		(10%) (2%)	1	(2%)
Intestine small, duodenum	(50)		(49)	(270)	(49)	(270)
Edema	(80)			(2%)	(49)	
Erosion	1	(2%)		(2%)	1	(2%)
Intestine small, ileum	(50)	(270)	(49)	(= ru)	(49)	(2 /0)
Edema	(00)			(2%)	(43)	
Ulcer				(2%)		
Intestine small, jejunum	(50)		(49)	(=)	(49)	
Edema				(2%)	, ,	
Liver	(50)		(50)		(50)	
Angiectasis	2	(4%)	6	(12%)	6	(12%)
Congestion			1	(2%)		
Developmental malformation	6	(12%)	5	(10%)	5	(10%)
Fatty change	3	(6%)	3	(6%)	2	(4%)
Focal cellular change		(58%)	41	(82%)		(82%)
Granuloma		(24%)		(26%)		(20%)
Hematopoletic cell proliferation	1	(2%)		(4%)		(4%)
Hemorrhage				(2%)	1	(2%)
Necrosis		(8%)	1	(2%)		
Arteriole, thrombus	_	(2%)	0.0	(700)	0.1	(00a)
Bile duct, hyperplasia	38	(76%)	36	(72%)		(62%)
Portal inflammation chronic	477	(0.4%)	46	(094)		(2%)
Portal, inflammation, chronic Portal, inflammation, chronic active	41	(94%)	46	(92%)		(80%)
Venule, thrombus	1	(2%)			1	(2%)
Mesentery	(1)	(470)	(1)		(2)	
Fat, necrosis		(100%)		(100%)		(50%)
Pancreas	(50)	,_00,0,	(49)	(200 //)	(50)	(00 /0)
Infiltration cellular, lymphocytic	(30)		(40)			(2%)
Inflammation, chronic active						(2%)
Acinus, atrophy	18	(36%)	24	(49%)		(30%)
Acinus, hyperplasia	1	(2%)		(8%)		
Acinus, vacuolization cytoplasmic						(2%)
Arteriole, inflammation, chronic					1	(2%)
Duct, ectasia		(2%)				
Salivary glands	(50)		(50)		(50)	
Infiltration cellular, lymphocytic						(2%)
Acinus, atrophy	, PA		/ F.A.:			(2%)
Stomach Foreign body	(50)		(50)		(50)	(0.05)
r oreign body Hyperplasia, squamous						(2%)
Stomach, forestomach	(50)		(50)			(2%)
Hyperkeratosis	(00)			(2%)	(50)	
Hyperkerawsis Hyperplasia, squamous	1	(2%)		(4%)	9	(6%)
	1	(470)		(6%)	ა	(070)
Intiammation acute						
Inflammation, acute Inflammation, chronic active	9	(4%)		(2%)	2	(6%)

TABLE B5. SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN FEMALE RATS IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

	Chambe	er Control	600 p	pm	1,200	ppm
ALIMENTARY SYSTEM (Continued)						
Stomach, glandular	(50)		(50)		(50)	
Inflammation, acute	(00)			(2%)		(2%)
Inflammation, chronic active			-	(= /4/		(4%)
Mucosa, dilatation	27	(54%)	26	(52%)	_	(56%)
Mucosa, erosion		(12%)		(8%)		(10%)
Mucosa, mineralization	•	(== /0/	-	(0,0)		(2%)
Mucosa, pigmentation	1	(2%)			-	(= /)
Mucosa, ulcer		(4%)				
Serosa, inflammation, chronic active	-	(1/0)			1	(2%)
Tooth			(1)		(2)	(2 /0 /
Peridontal tissue, inflammation, chronic			(1)			(50%)
Pulp, inflammation, chronic						(50%)
Pulp, inflammation, chronic active			1	(100%)	•	(00 %)
aup, initiatimation, emoine active				(100%)		
CARDIOVASCULAR SYSTEM						
Blood vessel	(50)		(50)		(50)	
Aorta, mineralization			,			(2%)
Heart	(50)		(50)		(50)	• •
Cardiomyopathy, chronic		(96%)	,	(100%)		(94%)
Inflammation, acute		(2%)				
Mineralization		(2%)			1	(2%)
Artery, mineralization	_					(2%)
Atrium, thrombus	2	(4%)	1	(2%)		(2%)
Ventricle, thrombus		(2%)	-	· · · · ·	-	
ENDOCRINE SYSTEM					-	
Adrenal gland, cortex	(50)		(40)		(49)	
Congestion	(50)		(49)			(2%)
	22	(CCM)	0.1	(60a)		
Degeneration, fatty	33	(66%)		(63%)	37	(76%)
Hematopoietic cell proliferation				(2%)		
Hemorrhage	10	(00%)		(2%)	•	(100)
Hyperplasia		(32%)		(39%)	-	(18%)
Hypertrophy	4	(8%)	9	(18%)		(8%)
Inflammation, chronic					_	(2%)
Pigmentation		(100%)		(98%)		(98%)
Adrenal gland, medulla	(49)		(48)		(49)	
Hematopoietic cell proliferation						(2%)
Hyperplasia		(8%)		(8%)	_	(12%)
Islets, pancreatic	(50)		(49)		(50)	
Hyperplasia		(2%)		(2%)		
Parathyroid gland	(42)		(44)		(41)	
Hypertrophy						(2%)
Pituitary gland	(50)		(50)		(50)	
Pars distalis, angiectasis	1	(2%)	3	(6%)		(2%)
Pars distalis, congestion						(2%)
Pars distalis, cyst	10	(20%)	14	(28%)	11	(22%)
Pars distalis, hemorrhage	3	(6%)			1	(2%)
Pars distalis, hyperplasia	7	(14%)	17	(34%)	7	(14%)
Pars distalis, metaplasia, osseous			1	(2%)		
Pars intermedia, angiectasis	1	(2%)		(4%)	2	(4%)
Pars intermedia, cyst		(2%)				
Pars nervosa, cyst		•	1	(2%)		
Thyroid gland	(50)		(50)	•	(50)	
Hemorrhage		(2%)	,		, /	
Inflammation, acute		(2%)				
	-	,	1	(2%)	2	(4%)
Ultimobranchial cvst						/
Ultimobranchial cyst C-cell, hyperplasia	17	(34%)		(46%)	15	(30%)

TABLE B5. SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN FEMALE RATS IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

	Chamb	er Control	600 I	pm	1,200	ppm
GENERAL BODY SYSTEM						
None						
GENITAL SYSTEM						
Clitoral gland	(49)		(50)		(44)	
Cyst	1	(2%)				
Hyperplasia			2	(4%)		(5%)
Inflammation, acute		(2%)			2	(5%)
Inflammation, chronic	21	(43%)	20	(40%)	21	(48%)
Inflammation, chronic active		(20%)	14	(28%)		(16%)
Duct, ectasia	2	(4%)				(2%)
Duct, hyperplasia, squamous						(2%)
Ovary	(50)		(50)		(50)	
Follicle, cyst						(2%)
Periovarian tissue, cyst	_	(4%)		(2%)		(4%)
Uterus	(50)		(50)		(50)	
Ectasia		(4%)	1	(2%)	1	(2%)
Hemorrhage		(2%)				
Prolapse	1	(2%)				
Endometrium, ectasia						(2%)
Endometrium, hyperplasia, cystic	2	(4%)		(4%)		(14%)
Endometrium, inflammation, chronic			1	(2%)		(2%)
Myometrium, inflammation, chronic						(2%)
Vagina	(4)		(4)		(3)	
Inflammation, acute		.054			1	(33%)
Inflammation, chronic active	1	(25%)				
IDMA TO DOLETTIC CACTEDA						
IEMATOPOIETIC SYSTEM	(40)		(50)		(50)	
Bone marrow	(49)		(50)		(50)	/o~ \
Hyperplasia	•	(4.0%)		/4.5a.		(2%)
Myelofibrosis	_	(16%)	9	(18%)	4	(8%)
Myeloid cell, hyperplasia		(2%)	(=0)		(50)	
Lymph node	(50)		(50)		(50)	
Congestion		(2%)		(2%)		(2%)
Hyperplasia, lymphoid		(12%)	3	(6%)		(20%)
Inflammation, acute	1	(2%)			1	(2%)
Pigmentation				(2%)		
Mediastinal, congestion		(O.W.)	1	(2%)		
Mediastinal, hemorrhage	1	(2%)	_	(O~ \	_	
Mediastinal, hyperplasia, lymphoid			1	(2%)		(4%)
Mediastinal, pigmentation		(00)	_	(00)		(2%)
Mesenteric, congestion		(8%)	3	(6%)	4	(8%)
Mesenteric, edema		(2%)	00	(640)	00	/FO~ :
Mesenteric, hyperplasia, lymphoid		(34%)		(64%)	29	(58%)
Mesenteric, inflammation, acute	1	(2%)		(2%)		
Renal, congestion	/45			(2%)	/40	
Lymph node, mandibular	(47)	(110)	(48)	(150)	(42)	(50° \
Congestion		(11%)		(17%)		(7%)
Cyst Hyperplasia lymphaid		(2%)		(2%)		(5%)
Hyperplasia, lymphoid Inflammation, acute		(87%)	42	(88%)		(93%)
Spleen		(2%)	(E0)			(2%)
Fibrosis	(50)		(50)	(6%)	(50)	(60-)
Hematopoietic cell proliferation	41	(82%)		(6%)		(6%)
Hyperplasia, lymphoid				(92%) (2%)		(88%)
Hyperplasia, lympholo Hyperplasia, reticulum cell		(2%) (2%)	1	(2%)	1	(2%)
Infarct	1	(270)	1	(2%)	1	(2%)
			1	(470)	1	(470)
	4.4	(88%)	4.4	/88 <i>0</i> L\	47	1010
Pigmentation Capsule, fibrosis	44	(88%)		(88%) (4%)	47	(94%)

TABLE B5. SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN FEMALE RATS IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

Chamb	er Control	600	opm	1,200	ppm
					
(46)		(46)		(46)	
			(4%)		(2%)
					(7%)
_	(-/-/			_	(2%)
1	(2%)	_	, ,	_	
					
(50)		(50)		(50)	
1	(2%)	, , , ,			
				1	(2%)
1	(2%)				
37	(74%)	34	(68%)	27	(54%)
3	(6%)	1	(2%)		
				1	(2%)
21	(42%)	31	(62%)	_	(46%)
(50)		(50)		(50)	
		,			(2%)
1	(2%)	1	(2%)	_	
		_		1	(2%)
		1	(2%)		
				1	(2%)
				1	(2%)
		1	(2%)		
(49)		(50)		(50)	
1	(2%)			2	(4%)
2	(4%)	1	(2%)		
1	(2%)			4	(8%)
3	(6%)	1	(2%)		
(50)		(50)		(49)	
11	(22%)	6	(12%)	13	(27%)
_				3	(6%)
		2	(4%)		
1	(2%)				
(56)		/FA:			
	(10%)		(100)		(10~)
					(12%)
7	(14%)			4	(8%)
40	(0.8%)			20	(100%)
49	(3070)			50	(100%)
1	(2%)	1	(470)	n	(4%)
1	(470)	1	(9%)		(4%) (4%)
90	(56%)			_	(4%) (72%)
		20	(*U70)	ან	(1270)
		30	(60%)	96	(52%)
					(32%) (4 %)
					(8%)
	(4%)		(2%)	4	(070)
	(= 70 /	1	(470)		
_		E	(100/-)	1	1902
		5	(10%)		(2%)
e		5	(10%)	1	(2%) (2%) (2%)
	(46) 4 2 1 (50) 1 1 37 3 21 (50) 1 2 1 3 (50) 11 1 5 1 1 1 28 1 28 1 28 1 25 17	11 (22%) 1 (2%) 5 (10%) 1 (2%) 1 (2%)	(46) (46) (46) 4 (9%) 2 2 (4%) 2 3 1 (2%) (50) (50) 1 (2%) 31 (50) (50) 1 (2%) 1 1 (2%) 1 (2%	(46) (46) (46) (46) (47) (2 (47) (2 (47) (3 (77)) (50) (50) (50) (50) (1 (27) (27) (27) (27) (27) (27) (27) (27)	(46) (46) (46) (46) (46) (46) (46) (46)

TABLE B5. SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN FEMALE RATS IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

(49)					
(49)					
		(50)		(50)	
				1	(2%)
4	(8%)	1	(2%)	1	(2%)
		34	(68%)		(64%)
	(/		
27	(55%)	41	(82%)	41	(82%)
1	(2%)		, - , - ,		
5	(10%)	2	(4%)	6	(12%)
	,,	1	(2%)	_	
	(90%)	_		47	(94%)
					(20%)
_	(1,0)	••	(52,0)		(2%)
		1	(2%)		(10%)
					(2%)
20	(50%)				(78%)
-			,		(6%)
				ა	(0%)
	(2%)	_	(2%)	(50)	
	(00)	(00)		(50)	
_					
1	(2%)				
		1	(2%)		
(5)		(4)		(3)	
	(20%)		(25%)	(0)	
				9	(67%)
U	(00 /0)			2	(01/0/
			(2070)	1	(33%)
					(33%)
4	(80%)	2	(75%)		(67%)
	, ,			2	(0170)
			(30%)		
(50)		(50)		(50)	
				1	(2%)
1	(2%)				
49	(98%)	48	(96%)	49	(98%)
		-		1	(2%)
1	(2%)	1	(2%)		
		-		1	(2%)
		1	(2%)		
-		_			
49	(98%)	_		48	(96%)
	/-/		,,		
,	(2%)	,		, /	(2%)
				_	(4%)
	27 1 5 5 tive 44 2 29 4 1 (50) 1 1 3 3 4 2 2 (50) 1 49 (50) 1 49 (50)	(50) (50) (1 (2%) (2%) (2%) (40%) (50) (1 (20%) (3 (60%) (50) (4 (80%) (2 (40%) (50) (1 (2%) (40%) (50) (1 (2%) (40%) (4 (80%)	27 (55%) 41 1 (2%) 5 (10%) 2 tive 44 (90%) 48 2 (4%) 11 29 (59%) 45 4 (8%) 1 (50) (50) 1 (2%) 1 (5) (4) 1 (20%) 1 (5) (4) 1 (20%) 2 (50) (50) 1 (2%) 2 (50) (50) 1 (2%) 49 (98%) 48 1 (2%) 1 49 (98%) 48 1 (2%) 1 49 (98%) 50 (50) 1 (2%) 50 (50) 1 (2%) (50)	27 (55%) 1 (2%) 1 (2%) 5 (10%) 2 (4%) 1 (2%) 44 (90%) 2 (4%) 11 (22%) 44 (90%) 2 (4%) 11 (22%) 1 (2%) 1 (2%) 29 (59%) 4 (8%) 1 (2%) 1 (2%) (50) (50) (50) 1 (2%) 1 (2%) 1 (25%) 4 (80%) 2 (50%) 4 (80%) 2 (50%) 4 (80%) 3 (60%) 2 (50%) (50) 1 (2%) (50) (50) (50) 1 (2%) (50) (50) (50) (50) 1 (2%) (50) (50) (50) 1 (2%) (50) (50) (50) 1 (2%) 4 (80%) 2 (50%) (50) (50) 1 (2%) 4 (80%) 3 (75%) 2 (40%) 2 (50%)	1 (2%) 1 (2%) 1 (2%) 5 (10%) 5 (10%) 2 (4%) 44 (90%) 48 (96%) 47 2 (4%) 11 (22%) 10 1 (2%) 1 (2%) 5 (10%) 48 (96%) 47 2 (4%) 11 (22%) 10 1 (2%) 5 (10%) 5 (10%) 1 (2%) 1

APPENDIX C

SUMMARY OF LESIONS IN MALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE

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TABLE C1. SUMMARY OF THE INCIDENCE OF NEOPLASMS IN MALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE

Cha	mber (Control	120 _l	opm	600 1	ppm	1,200 p	pm
Animals initially in study	60		60		60		60	
Animals removed	60		60		60		60	
Animals examined histopathologically	60		60		60		60	
LIMENTARY SYSTEM								
Gallbladder	(59)		(54)		(57)		(59)	
Lymphoma malignant mixed			1	(2%)			_	
Lymphoma malignant undifferentiated cell type			(00)		(00)			(2%)
Intestine large, cecum Lymphoma malignant mixed	(60)	(5%)	(60)		(60)		(59)	
Lymphoma malignant mixed Lymphoma malignant undifferentiated cell type		(370)					1	(2%)
Intestine large, colon	(60)		(60)		(60)		(59)	(270)
Lymphoma malignant lymphocytic	(00)		(00)			(2%)	(00)	
Lymphoma malignant mixed						(3%)		
Intestine small, jejunum	(60)		(60)		(60)	(0 ,0 ,	(60)	
Lymphoma malignant mixed	1	(2%)		(2%)	,		,	
Lymphoma malignant undifferentiated cell type							1	(2%)
Live:	(60)		(60)		(60)		(59)	
Hemangioma					1	(2%)		
Hemangiosarcoma	2	(3%)						(2%)
Hemangiosarcoma, metastatic, spleen							_	(2%)
Hepatocellular carcinoma		(22%)	-	(13%)		(15%)	_	(14%)
Hepatocellular adenoma	7	(12%)	10	(17%)	9	(15%)		(17%)
Hepatocellular adenoma, multiple								(2%)
Histiocytic sarcoma								(2%)
Ito cell tumor malignant			_				1	(2%)
Lymphoma malignant histiocytic	•			(3%)	•	/O.W.		
Lymphoma malignant mixed		(3%)	2	(3%)	Z	(3%)		(O.W.)
Lymphoma malignant undifferentiated cell type Mesentery	*(60)		±(CO)		*(00)			(2%)
Lymphoma malignant mixed		(2%)	*(60)		*(60)	(2%)	*(60)	
Lymphoma malignant mixed Lymphoma malignant undifferentiated cell type	1	(270)			1	(270)	1	(2%)
Pancreas	(60		(60)		(60)		(59)	(270)
Lymphoma malignant lymphocytic	101		(00)			(2%)	(007	
Lymphoma malignant mixed	2	(3%)				(5%)		
Lymphoma malignant undifferentiated cell type		(0,0)			•	(0,0)	1	(2%)
Salivary glands	(60)		(60)		(60)		(59)	(= /0/
Lymphoma malignant mixed	3	(5%)	2	(3%)	3	(5%)		
Stomach, forestomach	(60)		(59)		(60)		(60)	
Lymphoma malignant mixed					1	(2%)		
Papilloma squamous	1	(2%)		(2%)	1	(2%)		
Stomach, glandular	(60)		(60)		(60)		(60)	
Lymphoma malignant mixed			****			(2%)		
Tooth	*(60)		*(60)		*(60)		*(60)	
Pulp, lymphoma malignant undifferentiated cell type							1	(2%)
CARDIOVASCULAR SYSTEM						·		<u> </u>
Heart	(60)		(60)		(60)		(59)	
Hemangiosarcoma	'		'		,			(2%)
Lymphoma malignant histiocytic			1	(2%)			_	
Lymphoma malignant mixed					1	(2%)		
NDOCRINE SYSTEM						····		
Adrenal gland	(60)		(60)		(59)		(60)	
Lymphoma malignant mixed			1	(2%)				
Capsule, spindle cell, adenoma		(2%)						
Adrenal gland, cortex	(60)		(60)		(59)		(59)	
Adenoma					1	(2%)		

TABLE C1. SUMMARY OF THE INCIDENCE OF NEOPLASMS IN MALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

Char	nber (Control	120 p	pm	600 p	pm	1, 200 p	pm
ENDOCRINE SYSTEM (Continued)								
Islets, pancreatic	(60)		(60)		(60)		(58)	
Adenoma					_	(2%)	_	(2%)
Pituitary gland	(59)		(58)		(58)		(56)	
Pars distalis, adenoma			2	(3%)	1	(2%)		
Pars intermedia, adenoma	(00)		(00)		(00)			(2%)
Thyroid gland Follicle, adenoma	(60) 1	(2%)	(60) 2	(3%)	(60)		(59)	
GENERAL BODY SYSTEM None		2.14						- "
GENITAL SYSTEM								
Epididymis	(60)		(60)		(60)		(60)	
Lymphoma malignant histiocytic			2	(3%)		(0 .0)		
Lymphoma malignant mixed	4/44		*/**			(2%)	#/00:	
Preputial gland	* (60)		* (60)		*(60)	(90)	*(60)	
Lymphoma malignant mixed	(EO)		(60)		(59)	(2%)	(59)	
Prostate Lymphoma malignant mixed	(60)	(2%)	(00)			(2%)	(03)	
Lymphoma malignant mixed Lymphoma malignant undifferentiated cell type	1	(470)			•	(470)	1	(2%)
Seminal vesicle	*(60)		*(60)		*(60)		*(60)	/0/
Lymphoma malignant mixed		(2%)	(00)		.007		(00)	
Testes	(60)	(=,	(60)		(60)		(60)	
Lymphoma malignant mixed	1	(2%)			1	(2%)		
Lymphoma malignant undifferentiated cell type							1	(2%)
Interstitial cell, adenoma			1	(2%)			1	(2%)
HEMATOPOIETIC SYSTEM								
Bone marrow	(60)		(60)		(60)		(59)	
Hemangiosarcoma, metastatic, liver	1	(2%)						(0 <i>0</i> ′ \
Hemangiosarcoma, metastatic, spleen				/0 <i>0</i> / \			1	(2%)
Lymphoma malignant histocytic	1	(2%)		(2%) (2%)				
Lymphoma malignant mixed Lymph node	(60)	(270)	(60)	(270)	(59)		(59)	
Hepatocellular carcinoma, metastatic, liver	(00)		(00)			(2%)	(09)	
Histiocytic sarcoma	1	(2%)			•	(~ 10)		
Axillary, lymphoma malignant mixed		(2%)	1	(2%)				
Iliac, lymphoma malignant mixed Iliac, lymphoma malignant undifferentiated		(3%)		(3%)	1	(2%)		
cell type								(2%)
Mediastinal histocytic sarcoma	1	(9%)	0	(20:1	o	(20)	1	(2%)
Mediastinal, lymphoma malignant mixed Mediastinal, lymphoma malignant undifferentiated cell type	1	(2%)	2	(3%)	2	(3%)	1	(2%)
Mesenteric, hemangiosarcoma, metastatic, liver	1	(2%)					•	_ <i>\\</i> /
Mesenteric, histiocytic sarcoma	-		2	(0 <i>0</i> ()			1	(2%)
Mesenteric, lymphoma malignant histiocytic			2	(3%)		(90)		
Mesenteric, lymphoma malignant lymphocytic Mesenteric, lymphoma malignant mixed		(8%)	E	(10%)		(2%) (10%)	2	(5%)
Mesenteric, lymphoma malignant mixed Mesenteric, lymphoma malignant	3	(070)	0	(1070)	v	(1070)	3	10 10)
undifferentiated cell type							1	(2%)
Renal, histiocytic sarcoma								(2%)
Renal, lymphoma malignant mixed	2	(3%)	1	(2%)	1	(2%)	-	,
Renal, lymphoma malignant undifferentiated cel							1	(2%)

TABLE C1. SUMMARY OF THE INCIDENCE OF NEOPLASMS IN MALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

Alveolar/bronchiolar adenoma, multiple Alveolar/bronchiolar carcinoma	Cha	mber (Control	120 լ	ppm	600 1	ppm	1 ,200 p	pm
Lymphon malignant histocytic Lymphons malignant histocytic Lymphons malignant histocytic Lymphons malignant mixed 3 (6%) 2 (4%) 4 (7%) 2 (4%) Lymphons malignant mixed 3 (6%) 2 (4%) 4 (7%) 3 (5%) Lymphons malignant mixed 3 (6%) 2 (4%) 4 (7%) 3 (5%) Lymphons malignant mixed 60 600 660 659 659 660 660 660 660 659	HEMATOPOIETIC SYSTEM (Continued)								·······
Lymphoma malignant histocytic 1 (2%) 1 (2%		(53)		(52)		(54)		(51)	
Lymphoma malignant lymphocytic Lymphoma malignant mixed 1 (2%) 2 (4%) 4 (7%) 2 (4%) 1 (2%) 3 (5%) 1 (2%) 3 (5%) 1 (2%) 3 (5%) 1 (2%) 3 (5%) 1 (2%) 3 (5%) 1 (2%) 3 (5%) 1 (2%) 3 (5%) 1 (2								,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	
Lymphoma malignant undifferentiated cell type Spleen	Lymphoma malignant lymphocytic					1	(2%)		
Lymphoma malignant undifferentiated cell type Spleen 1 (2%) 3 (5%)		3	(6%)	2	(4%)	4	(7%)	2	(4%)
Spien		е						1	(2%)
1 (2%) 1				(60)		(60)		(59)	
1 (2%) 1	Hemangiosarcoma					1	(2%)	3	(5%)
Lymphoma malignant mixed 5 (8%) 5 (8%) 6 (10%) 3 (5%)								1	(2%)
Lymphoma malignant mixed 5 (8%) 5 (8%) 6 (10%) 3 (5%) 1 (2%)				3	(5%)				
Lymphoma malignant undifferentiated cell type		5	(8%)	5	(8%)	6	(10%)	3	(5%)
Thymus							,	1	(2%)
Histocytic sarcoma				(59)		(58)			(,,
Lymphoma malignant mixed 2 (4%) 2 (3%) 3 (5%) 1 (2%) 1				(**/		,,,,			(2%)
Lymphoma malignant mixed 2 (4%) 2 (3%) 3 (5%) 1 (2%) 1		-	,_,,	1	(2%)			_	(= ,0 ,
Lymphoma malignant undifferentiated cell type		2	(4%)			3	(5%)	1	(2%)
NTEGUMENTARY SYSTEM Skin (60) (60) (60) (60) (58)			,	-		·	,		
Skin (60) (60) (60) (58) (58)									
Lymphoma malignant mixed Neck, subcutaneous tissue, hemangioma 1 (2%) 1 (2%)		,						. = *	
Neck, subcutaneous tissue, hemangioma		(60)		(60)			. 0∼∶	(58)	
MUSCULOSKELETAL SYSTEM Go Go Go Go Go Go Go G					(O.O.)	1	(2%)		
Bone	Neck, subcutaneous tissue, nemangioma			1	(2%)				
Hemangiosarcoma, metastatic, spleen *(60) *(60) *(60) *(60) *(60) *(60)	MUSCULOSKELETAL SYSTEM								
Hemangiosarcoma, metastatic, spleen *(60) *(60) *(60) *(60) *(60) *(60)	Bone	(60)		(60)		(60)		(60)	
Skeletal muscle *(60) *(60) *(60) *(60) *(60) (60)	Hemangiosarcoma, metastatic, spleen			,		'			(2%)
Head, lymphoma malignant mixed 1 (2%)	Skeletal muscle	*(60)		*(60)		*(60)			
Brain	Head, lymphoma malignant mixed					1	(2%)		
Brain	NERVOUS SYSTEM								
Lymphoma malignant mixed 1 (2%) 1 (2%)		(60)		(60)		(60)		(60)	
Color		(00)			(2%)		(2%)	(00)	
Lung Adenocarcinoma, metastatic, harderian gland Alveolar/bronchiolar adenoma Alveolar/bronchiolar adenoma, multiple Alveolar/bronchiolar carcinoma Alveolar/bronchiolar adenoma, multiple Alveolar/bronchiolar adenoma, multiple Alveolar/bronchiolar adenoma, multiple Alveolar/bronchiolar adenoma, metastatic, liver Alveolar/bronchiolar adenoma, metastatic, liver Alveolar/bronchiolar adenoma, metastatic, liver Alveolar/bronchiolar adenoma Alveolar/bronchiolar adenoma Alveolar/bronchiolar adenoma Alveolar/bronchiolar adenoma Alveolar/bronchiolar adenoma Alveolar/bronchiolar adenoma, multiple Alveolar/bronchiolar adenoma multiple Alve					(270)		(270)		
Adenocarcinoma, metastatic, harderian gland									
Alveolar/bronchiolar adenoma 8 (13%) 1 (2%) 2 (3%) 7 (12%) Alveolar/bronchiolar adenoma, multiple Alveolar/bronchiolar carcinoma 2 (3%) 1 (2%) 1 (2%) Hepatocellular carcinoma, metastatic, liver 3 (5%) 1 (2%) 1 (2%) Histiocytic sarcoma 1 (2%) Lymphoma malignant histiocytic Lymphoma malignant mixed 2 (3%) Lymphoma malignant undifferentiated cell type Nose (59) (59) (60) (59) Adenocarcinoma, metastatic, harderian gland Mucosa, lymphoma malignant mixed 1 (2%) Submucosa, lymphoma malignant undifferentiated cell type 1 (2%) PECIAL SENSES SYSTEM Harderian gland *(60) *(60) *(60) *(60) *(60) Adenocarcinoma 1 (2%)		(60)		(60)				(60)	
Alveolar/bronchiolar adenoma, multiple Alveolar/bronchiolar carcinoma							,		
Alveolar/bronchiolar carcinoma 2 (3%) 1 (2%) 1 (2%) Hepatocellular carcinoma, metastatic, liver 3 (5%) 1 (2%) 1 (2%) Histiocytic sarcoma 1 (2%) Lymphoma malignant histiocytic 2 (3%) Lymphoma malignant mixed 3 (5%) 3 (5%) 2 (3%) Lymphoma malignant undifferentiated cell type 1 (2%) Nose (59) (59) (60) (59) Adenocarcinoma, metastatic, harderian gland Mucosa, lymphoma malignant mixed 1 (2%) Submucosa, lymphoma malignant undifferentiated cell type 1 (2%) PECIAL SENSES SYSTEM Harderian gland *(60) *(60) *(60) *(60) Adenocarcinoma 1 (2%)		8	(13%)	1	(2%)	2	(3%)		
Hepatocellular carcinoma, metastatic, liver 3 (5%) 1 (2%) 1 (2%) Histiocytic sarcoma 1 (2%) 1 (2%									
Histiocytic sarcoma Lymphoma malignant histiocytic Lymphoma malignant mixed Lymphoma malignant mixed Lymphoma malignant undifferentiated cell type Nose (59) Adenocarcinoma, metastatic, harderian gland Mucosa, lymphoma malignant mixed Submucosa, lymphoma malignant undifferentiated cell type PECIAL SENSES SYSTEM Harderian gland Adenocarcinoma *(60) *(60) *(60) *(60) *(60) *(60) *(60) *(60) *(60) *(60) *(60) *(60) *(60) *(60) *(60) *(60) *(60)		_		1	(2%)				
Lymphoma malignant histocytic Lymphoma malignant mixed Lymphoma malignant mixed Lymphoma malignant undifferentiated cell type Nose (59) Adenocarcinoma, metastatic, harderian gland Mucosa, lymphoma malignant mixed Submucosa, lymphoma malignant undifferentiated cell type PECIAL SENSES SYSTEM Harderian gland Adenocarcinoma *(60) *(60) *(60) *(60) *(60) *(60) *(60) *(60) *(60) *(60) *(60) *(60) *(60) *(60) *(60) *(60) *(60) *(60)		3	(5%)			1	(2%)		
Lymphoma malignant mixed 3 (5%) 3 (5%) 2 (3%) Lymphoma malignant undifferentiated cell type Nose (59) (59) (60) (59) Adenocarcinoma, metastatic, harderian gland Mucosa, lymphoma malignant mixed Submucosa, lymphoma malignant undifferentiated cell type 1 (2%) PECIAL SENSES SYSTEM Harderian gland *(60) *(60) *(60) *(60) Adenocarcinoma 1 (2%)								1	(2%)
Lymphoma malignant undifferentiated cell type Nose (59) Adenocarcinoma, metastatic, harderian gland Mucosa, lymphoma malignant mixed Submucosa, lymphoma malignant undifferentiated cell type PECIAL SENSES SYSTEM Harderian gland Adenocarcinoma 1 (2%) (59) (59) (60) (59) (1 (2%) 1 (2%) 1 (2%) *(60) *(60) *(60) *(60) *(60) *(60) *(60) *(60)									
Nose (59) (59) (60) (59) Adenocarcinoma, metastatic, harderian gland Mucosa, lymphoma malignant mixed 1 (2%) Submucosa, lymphoma malignant undifferentiated cell type 1 (2%) PECIAL SENSES SYSTEM Harderian gland *(60) *(60) *(60) *(60) Adenocarcinoma 1 (2%)			(5%)	3	(5%)	2	(3%)		_
Adenocarcinoma, metastatic, harderian gland Mucosa, lymphoma malignant mixed Submucosa, lymphoma malignant undifferentiated cell type PECIAL SENSES SYSTEM Harderian gland Adenocarcinoma 1 (2%) 1 (2%) 1 (2%)						,			(2%)
Mucosa, lymphoma malignant mixed Submucosa, lymphoma malignant undifferentiated cell type 1 (2%) PECIAL SENSES SYSTEM Harderian gland Adenocarcinoma *(60) *(60) *(60) *(60) *(60) *(60)		(59)		(59)				(59)	
Submucosa, lymphoma malignant									
undifferentiated cell type 1 (2%) PECIAL SENSES SYSTEM Harderian gland *(60) *(60) *(60) *(60) Adenocarcinoma 1 (2%)						1	(2%)		
Harderian gland *(60) *(60) *(60) *(60) Adenocarcinoma 1 (2%)								1	(2%)
Harderian gland *(60) *(60) *(60) *(60) Adenocarcinoma 1 (2%)			·						
Adenocarcinoma 1 (2%)		*(60)		*(<i>&</i> ()		*(60)		*/ <i>C</i> /\	
- \ - \-		(00)		(00)			1206	(00)	
	Adenoma			9	(5%)	1	(470)	n	(30-1

TABLE C1. SUMMARY OF THE INCIDENCE OF NEOPLASMS IN MALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

Chai	mber (Control	120 p	pm	600 p	opm	1,200 p	pm
URINARY SYSTEM			***************************************			·		
Kidney	(60)		(60)		(60)		(59)	
Hepatocellular carcinoma, metastatic, liver					1	(2%)		
Lipoma			1	(2%)				
Lymphoma malignant mixed	4	(7%)	4	(7%)	3	(5%)	1	(2%)
Urinary bladder	(60)		(60)		(60)		(59)	
Lymphoma malignant mixed	2	(3%)	2	(3%)	3	(5%)		
Lymphoma malignant undifferentiated cell type							1	(2%)
SYSTEMIC LESIONS								
Multiple organs	*(60)		*(60)		*(60)		*(60)	
Lymphoma malignant mixed	5	(8%)	, ,	(12%)	, ,	(10%)	1,	(5%)
Hemangiosarcoma	2	(3%)			1	(2%)	5	(8%
Lymphoma malignant histiocytic			3	(5%)				
Hemangioma			1	(2%)	1	(2%)		
Lymphoma malignant lymphocytic					2	(3%)		
Lymphoma malignant undifferentiated cell							1	(2%)
ANIMAL DISPOSITION SUMMARY				·				
Animals initially in study	60		60		60		60	
Moribund	19		19		25		17	
Terminal sacrifice	17		21		16		19	
Dead	23		19		14		21	
Accident	1				2		2	
Natural death			1		3		1	
TUMOR SUMMARY			· · · · · ·					
Total animals with primary neoplasms **	29		29		26		33	
Total primary neoplasms	42		41		35		50	
Total animals with benign neoplasms	16		18		14		19	
Total benign neoplasms	18		22		16		24	
Total animals with malignant neoplasms	20		16		17		17	
Total malignant neoplasms	24		19		19		26	
Total animals with secondary neoplasms ***	4				3		2	
Total secondary neoplasms	5				5		4	

^{*} Number of animals receiving complete necropsy examinations; all gross lesions including masses examined microscopically.

** Primary tumors: all tumors except secondary tumors

*** Secondary tumors: metastatic tumors or tumors invasive into an adjacent organ

TABLE C2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE: CHAMBER CONTROL

	IMIMERITOR	٠.				,,			211	ш.	0.	17.	.,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,			011			_							
WEEKS ON STUDY	1	0 4 2	0 4 2	0 4 2	0 4 8	0 5 4	0 5 6	0 6 2	0 6 2	0 6 3	0 6 3	0 6 4	0 6 4	0 6 4	0 6 4	0 6 8	0 7 1	0 7 5	0 7 6	0 7 6	0 7 7	0 7 8	0 7 9	0 7 9	0 8 0	0 8 1
CARCASS ID		0 6 1	8 1	5 9 1	4 6 1	3 3 1	5 2 1	1 9 1	1 0 1	3 2 1	0 9 1	2 8 1	0 4 1	4 3 1	0 1 1	6 1	4 0 1	3 7 1	1 1	5 8 1	2 2 1	2 4 1	1 1	2 3 1	0 7 1	5 4 1
ALIMENTARY SYSTEM																										
Esophagus		+	M	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Esophagus Galibladder		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	M	+	+	+	+	+
Intestine large		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine large, cecum Lymphoma malignant mixed		+	+	+	-	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	_
Intestine large, colon		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine large, rectum	1	+	+	+	M	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine small Intestine small, duodenum		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine small, ileum		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	÷	+	+	+	+	+	+
Intestine small, jejunum		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed Liver																										+
Hemangiosarcoma		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	X	+	+	+	+
Hepatocellular carcinoma										Х									Х			••	Х	X		
Hepatocellular adenoma															X		Х				Х					
Lymphoma malignant mixed																										
Mesentery Lymphoma malignant mixed																										
Pancreas		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed																										
Salivary glands		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed Stomach																			,		,					
Stomach, forestomach		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Papilloma squamous		,	,	,		•	,				,	,	•		,			,								
Stomach, giandular		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Tooth														+												
CARDIOVASCULAR SYSTEM																										
Blood vessel		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	M	+	+	+	+	+
Heart		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
ENDOCEDNIE SYSTEM																										
ENDOCRINE SYSTEM Adrenal gland		4	_	_	_	_	_	_	_	_	_	_	-1	_	_	_	_	_	_	_	_	_	_	_	_	_
Capsule, spindle cell, adenoma		-	•	_	*		~	-	-	т.	_		-	-	т		7	7	-	-		,	-			-
Adrenal gland, cortex		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Adrenal gland, medulla		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Islets, pancreatic Parathyroid gland	ì	+ M	+	+	+	+	+	+	+	+	, M	+	+	+	+	+ M	+	+	+	+	*	+	+ M	+	+	+
Pituitary gland		+ tar	M +	M +	M +	+	<u> </u>	M +	+	M +	141	+	Ŧ	Ţ	+	+	+	M +	+	+	M +	+	141	- T	M +	+
Thyroid gland		÷	+	+	+	+	+	+	+	+	+	+	÷	+	÷	+	+	+	+	+	+	+	+	+	÷	+
Follicle, adenoma																										
GENERAL BODY SYSTEM None											—				—											
GENITAL SYSTEM																										
Epididymis	1	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Penis							+							+		+	+		+	+	+		+	+		
Preputial gland	1			+	+				+	M				+	+		+	+			+					
Prostate Lymphoma malignant mixed		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Seminal vesicle										+									+							
Lymphoma malignant mixed																										
Testes		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed																										

^{+:} Tissue examined microscopically
: Not examined
-: Present but not examined microscopically
I: Insufficient tissue

M: Missing
 A: Autolysis precludes examination
 X: Incidence of listed morphology

TABLE C2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE MICE: CHAMBER CONTROL (Continued)

WEEKS ON STUDY	8	0 8	0 8	0 8	8	8	8	9	0 9	9	9	9	0	9	1 0	1	1	0	0	0	0	1	0	0	1
STODE	3	3	5	7	7	8	9	0	0	1	1	2	2	6	1	2	2	2	5	5	5	5	5	5	5
CARCASS	-2	5	5	3	-	- 1	3	-5	5		9	_	2	0	,	6	5	5	0	_	-	2	4	3	4
ID	0	3	5	9	3	6	8	1	6	5	7	5	9	8	8	0	0	7	2	2	4	5	9	Ō	4
	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1
ALIMENTARY SYSTEM																									
Esophagus Gallbladder	+	. +	. +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine large					+	+	+	+	+	+	+	+	+	+	+	+	+	÷	÷	+	+	+	+	+	+
Intestine large, cecum	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed Intestine large, colon					+	+	+	+	+	+	+	+	+	+	X	+	+	+	+	+	X	+	+	+	X +
Intestine large, rectum					+	+	+	+	+	+	+	+	+	+	+	+	÷	+	+	+	+	+	÷	+	+
Intestine small	+		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine small, duodenum Intestine small, ileum	+		. +	. +	+	+	+	+	+	+	+	+	±	+	+	+	+	+	+	+	+	+	+	+	+
Intestine small, jejunum		. +	+		+	+	+	+	+	+	+	÷	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed																					X				
Liver Hemangiosarcoma	†	. +	+	. +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	*X	+
Hepatocellular carcinoma					Х							х	X			X	Х						Х	**	
Hepatocellular adenoma										Х		Х													X
Lymphoma malignant mixed															X								_		
Mesentery Lymphoma malignant mixed															X								Ŧ		
Pancreas	4	- +	- +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed																					Х				
Salivary glands Lymphoma malignant mixed	1	- +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	x X	+	+	+	*
Stomach		- +	- +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Stomach, forestomach		- +	- +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Papilloma squamous																						_		_	_
Stomach, glandular Tooth	1		•	•						_	_			Τ.	_	-	т	_		-	_	т	-		-
CARDIOVASCULAR SYSTEM	_																								
Blood vessel	1 4		- 4	. +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Heart	1		- +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
ENDOCRINE SYSTEM																									
Adrenal gland	1.	- 4			+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Capsule, spindle cell, adenoma						·																			
Adrenal gland, cortex	-		- +	- +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Adrenal gland, medulla Isiets, pancreatic			- +	- +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Parathyroid gland	l N					+	+	+	+	+	+	М	+	+	M	M	M	+	M	M	+	M		+	M
Pituitary gland		- 4			+	+	+	+	+	M	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Thyroid gland Follicle, adenoma	1	- +	- +	- +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
GENERAL BODY SYSTEM																									
None																									
GENITAL SYSTEM																									
Epididymis	+	- +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Penis Preputial gland		•	-	+	+	+	+	+	+		+	+	+	+											
Prostate	-	- 4	- 4	- +	+	÷	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed																					X				+
Seminal vesicle Lymphoma malignant mixed	1																								X
Testes			+ +	- +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed																									X

TABLE C2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE MICE: CHAMBER CONTROL (Continued)

	(Continued)	
WEEKS ON STUDY	$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	
CARCASS ID	0 0 2 3 1 4 1 3 3 4 3 5 1 4 7 2 6 1 5 7 1 1 1 1 1 1 1 1 1 1	TOTAL: TISSUES TUMORS
ALIMENTARY SYSTEM		
Esophagus Gallbladder	+ + + + + + + + + + + + + + + + + + + +	. 59 . 59
Intestine large	+ + + + + + + + + + + +	60
Intestine large, cecum	+ + + + + + + + +	60
Lymphoma malignant mixed Intestine large, colon	+ + + + + + + + +	60
Intestine large, rectum	+ + + + + + + + + + + + + + + + + + + +	59 60
Intestine small Intestine small, duodenum	\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	60
Intestine small, ileum	+ + + + + + + + +	60 60
Intestine small, jejunum Lymphoma malignant mixed	+ + + + + + + + +	1
Liver	+ + + + + + + + +	60
Hemangiosarcoma Hepatocellular carcinoma	x x x	13
Hepatocellular adenoma	X	7
Lymphoma malignant mixed Mesentery	X	2 2
Lymphoma malignant mixed		1
Pancreas	+ + + + + + + + + + +	60
Lymphoma malignant mixed Salivary glands	A + + + + + + + + + + + + + + + + + + +	60
Lymphoma malignant mixed Stomach	X	3 60
Stomach, forestomach	\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	60
Papilloma squamous	X X	60
Stomach, glandular Tooth		
CARDIOVASCULAR SYSTEM		
Blood vessel	\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	! 59 60
Heart	+ + + + + + + + +	
ENDOCRINE SYSTEM		60
Adrenal gland Capsule, spindle cell, adenoma	+ + + + + + + + + + X	li
Adrenal gland, cortex	+ + + + + + + + + + + + + + + + + + +	60 60
Adrenai giand, medulla Islets, pancreatic	1 + + + + + + + + + +	60
Parathyroid gland	+ M M M M M + M M +	30 59
Pituitary gland Thyroid gland	+ + + + + + + + + + + + + + + + + + +	60
Follicle, adenoma	X	1
GENERAL BODY SYSTEM None		
GENITAL SYSTEM		
Epididymis Penis	+ + + + + + + + + + + +	60 17
Preputial gland	T T	12
Prostate	+ + + + + + + + +	60
Lymphoma malignant mixed Seminal vesicle	+	4
Lymphoma malignant mixed		1 60
Testes Lymphoma malignant mixed	+ + + + + + + + + + +	1
-1E-our word was many		

TABLE C2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE MICE: CHAMBER CONTROL (Continued)

					(0	V		uec	•/																
WEEKS ON STUDY	0 4 2	0 4 2	0 4 2	0 4 8	0 5 4	0 5 6	0 6 2	0 6 2	0 6 3	0 6 3	0 6 4	0 6 4	0 6 4	0 6 4	0 6 8	0 7 1	0 7 5	0 7 6	0 7 6	0 7 7	0 7 8	0 7 9	0 7 9	0 8 0	0 8 1
CARCASS ID	0 6 1	8 1	5 9 1	4 6 1	3 3 1	5 2 1	9	1 0 1	3 2 1	0 9 1	2 8 1	0 4 1	4 3 1	0 1 1	2 6 1	4 0 1	3 7 1	1 1	5 8 1	2 2 1	2 4 1	1 1	2 3 1	0 7 1	5 4 1
HEMATOPOIETIC SYSTEM Bone marrow Hemangiosarcoma, metastațic, liver	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed Lymph node Histiocytic sarcoma Axillary, lymphoma malignant mixed Iliac, lymphoma malignant mixed Mediastinal, lymphoma malignant mixed Mesenteric, hemangiosarcoma, metastatic, liver Mesenteric, lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	*X	+	+	+	+
Renal, lymphoma malignant mixed Lymph node, mandibular Lymphoma malignant mixed	+	+	+	+	+	M	+	+	+	+	+	+	+	+	+	+	M	+	+	+	M	M	+	+	+
Spieen Lymphoma malignant mixed Thymus Histiocytic sarcoma Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+ M	+ X	+	+	+	+
INTEGUMENTARY SYSTEM Mammary gland Skin	M +	M +	M +	++	M +	M	M	M +	M +	M +															
MUSCULOSKELETAL SYSTEM Bone	_	+		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
NERVOUS SYSTEM Brain	_ _	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
RESPIRATORY SYSTEM Lung Alveolar/bronchiolar adenoma Alveolar/bronchiolar carcinoma Hepatocellular carcinoma, metastatic, liver	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+ X	+	+	+	+	+	+	+
Lymphoma malignant mixed Nose Trachea	++	+	++	M +	+	+	+	+	+	+	+	++	++	+	+	+	+	+	+	+	+	+	+	++	++
SPECIAL SENSES SYSTEM None	_																								
URINARY SYSTEM Kidney Lymphoma malignant mixed Ureter Urethra Urnary bladder Lymphoma malignant mixed	+ + +	+ + +	+	+ + +	+	+	+ + +	+	+	+	+	+	+	+ + +	+	+	+	+ + + +	+ + +	+ + +	+	+ + +	+ + +	+	+

TABLE C2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE MICE: CHAMBER CONTROL (Continued)

WEEKS ON STUDY	0 8 3	0 8 3	0 8 5	0 8 7	0 8 7	0 8 8	0 8 9	0 9 0	9	0 9 1	0 9 1	0 9 2	0 9 2	0 9 6	1 0 1	1 0 2	1 0 2	1 0 2	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5
CARCASS ID	0 1	5 3 1	5 5 1	3 9 1	1 3 1	3 6 1	3 8 1	5 1 1	5 6 1	1 5 1	7 1	4 5 1	9 1	0 8 1	1 8 1	6 0 1	5 0 1	5 7 1	0 2 1	1 2 1	1 4 1	2 5 1	4 9 1	3 0 1	4
HEMATOPOIETIC SYSTEM Bone marrow Hemangiosarcoma, metastatic, liver Lymphoma malignant mixed Lymph node	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	-	+	* X	+ X +
Misticcytic sarcoma Axillary, lymphoma malignant mixed lliac, lymphoma malignant mixed Mediastinal, lymphoma malignant mixed Mesenteric, hemangiosarcoma,		•	•		τ.	r	Ť	7	•			,	,		X X X	,									x
metastatic, liver Mesenteric, lymphoma malignant mixed Renal, lymphoma malignant mixed Lymph node, mandibular Lymphoma malignant mixed Spieen Lymphoma malignant mixed Thymus Histocytic sarcoma Lymphoma malignant mixed	+ +	+ + M	+ +	+ + +	M + +	+ + +	+ + M	+ + +	+ +	+ + +	+ + +	+ +	M + +	+ + +	X X + X + X M	++++	+++	+ + +	+ + +	M + +	X X + X M	+ + +	+ +	+ + +	X X + X + X
INTEGUMENTARY SYSTEM Mammary gland Skin	M +	M +	M +	M +	M +	M +	+ +	M +	M	M +	M +	+	M +	M +	M +	M +	++	M +	M +	M +	M +	M +	M	M +	M +
MUSCULOSKELETAL SYSTEM Bone	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
NERVOUS SYSTEM Brain	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	÷	+
RESPIRATORY SYSTEM Lung Alveolar/bronchiolar adenoma Alveolar/bronchiolar carcinoma Hepatocellular carcinoma, metastatic,	+	+	+	+	+	* X	X X	+	+	+	+	*X	* X	+	, X	+	+ X	+	+	, X	+	+	+	+	+
niver Lymphoma malignant mixed Nose Trachea	+++	+	++	+	++	++	+	+	+	+	+	++	++	++	X + +	++	++	++	+	* + +	++	++	+	++	X + +
SPECIAL SENSES SYSTEM None	-																								
URINARY SYSTEM Kidney Lymphoma malignant mixed Ureter	+	+	+	+	+	+	+	+	+	+	+	+	+	+	*	+	+	+	+	+	, X	+	+	+	+ X
Ürethra Urinary bladder Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	+	+	+	+	*X	+	+	+	+	+	+	+	+	+	*

TABLE C2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE MICE: CHAMBER CONTROL (Continued)

WEEKS ON STUDY	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	TOTAL.
CARCASS ID	0 3 1	0 5 1	2 1 1	3 4 1	1 7 1	4 2 1	1 6 1	3 1 1	3 5 1	7	 TOTAL: TISSUES TUMORS
HEMATOPOIETIC SYSTEM Bone marrow Hemangiosarcoma, metastatic, liver	+	+	+	+	+	+	+	+	+	+	60 1 1
Lymphoma malignant mixed Lymph node Histiccytic sarcoma Axillary, lymphoma malignant mixed Iliac, lymphoma malignant mixed Mediastinal, lymphoma mal. mixed Mesenterc, hemangiosarcoma,	+	+	+	+	+	+	+	+	+	+	60
metastatic, liver Mesenteric, lymphoma malignant mixed Renal, lymphoma malignant mixed Lymph node, mandibular Lymphoma malignant mixed	+	* *	X +	+	+	+	+	+	+	+	1 5 2 53 3
Spieen Lymphoma malignant mixed Thymus Histiocytic sarcoma Lymphoma malignant mixed	+	* X + X	+ X +	+	+	+ M	+	+	+ M	+	60 5 53 1 2
INTEGUMENTARY SYSTEM Mammary gland Skin	++	M +	M +	M +	+	M +	M +	M +	M +	M +	 6 60
MUSCULOSKELETAL SYSTEM Bone	+	+	+	+	+	+	+	+	+	+	60
NERVOUS SYSTEM Brain	+	+	+	+	+	+	+	+	+	+	60
RESPIRATORY SYSTEM Lung Alveolar/bronchiolar adenoma Alveolar/bronchiolar carcinoma	*	+	* X	+	+	+	+	+	+	+	60 8 2
Hepatocellular carcinoma, metastatic, liver Lymphoma malignant mixed Nose Trachea	++	X X + +	++	++	++	++	++	+	++	++	3 3 59 60
SPECIAL SENSES SYSTEM None											
URINARY SYSTEM Kidney Lymphoma malignant mixed Ureter Urethra	+	, X	+	+	+	+	+	+	+	+	60 4 8 5
Urinary bladder Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	60 2

TABLE C2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE: 120 ppm

WEEKS ON STUDY		0 2 3	0 5 6	0 5 7	0 5 8	6	0 6 3	0 6 4	0 6 5	0 7 2	0 7 2	0 7 2	0 7 3	0 7 5	0 7 6	0 7 6	0 8 0	0 8 0	8	0 8 1	0 8 2	8 3	8	0 8 5	0 8 5	0 8 7
CARCASS	-	4	4	3 8	- 4	4	4	3 9	3 8	4	3 7	3 8	3 7	3 9	3 9	4 0	3 6	3 9	3 6	3 7	3 8	3 9	3 6	3 7	4 0	3 9
ID	;	4	9	2	8	3	2 1	1	1	7	6	5	3	7 1	5	5	6	3	2	1	4	4	5	5 1	7 1	0
LIMENTARY SYSTEM		-																								
sophagus allbladder		+	+	+	+	+	+	+	+	+	+	+ M	+ M	+ +	+	+ M	, M	+ M	+	+	+	+	+	+	+	-
Lymphoma malignant mixed testine large		+	_	_		_	_	_	_	_	_	_	_	_	_	_	_	4	_	_	_	_	_	_	_	4
testine large, cecum		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	÷	-
testine large, colon		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	
testine large, rectum testine small		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	M +	
testine small, duodenum		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	
testine small, ileum		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	
testine small, jejunum Jymphoma malignant mixed		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	
ver		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	
Hepatocellular carcinoma Hepatocellular adenoma Lymphoma malignant histiocytic	ĺ											x		X	X		X	х							X	
ymphoma maiignant mixed																		Λ.							X	
ncreas	1	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	
livary glands ymphoma malignant mixed	i	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	
omach		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	
omach, forestomach		+	+	+	M	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	
Papilloma squamous omach, glandular	į	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	
ooth																	+									
ARDIOVASCULAR SYSTEM														_												_
lood vessel eart		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	
eart Lymphoma malignant histiocytic		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	
NDOCRINE SYSTEM	i																									
irenal gland		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	
ymphoma malignant mixed	ļ																									
irenal gland, cortex irenal gland, medulla	i	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	H M	+	+	+	+	+	+	+	
ets, pancreatic		+	+	<u>.</u>	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	
rathyroid gland		+	-	+	M	+	+	+	M	M	M	M	M	M	+	+	+	M	M	M	M	+	M	+	M	
tuitary gland Pars distalis, adenoma		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	M	+	+	M	+	+	+	+	+	
iyroid gland		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	
Follicle, adenoma	İ																									
ENERAL BODY SYSTEM																										_
ENITAL SYSTEM																										-
oididymis Lymphoma malignant histiocytic	1	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	
nis															+							+		+	+	
reputial gland rostate		+		+				+							+											
ostate minal vesicle		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	
estes	j	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	
Interstitial cell, adenoma																	X									

TABLE C2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE MICE: 120 ppm (Continued)

STUDY 8 8 8 9 9 9 9 9 9 9 9 9 9 9 0 0 0 0 0 0																										
CARCASS 0	WEEKS ON STUDY	8	8		9	0 9 1		9	9	9	9	9							1 0 5	1 0 5				0		0
Cappagus		0	1 5	6	$\frac{7}{2}$	7	2	8	1	6	8	7 8	0	0	6 7	8	9	0	6	3 7 9 1	8 7	8	9	6	6	0
Cappagus	ALIMENTARY SYSTEM																							_		
Intestine small, lieum	Esophagus Gailbiadder Lymphoma malignant mixed Intestine large, cecum Intestine large, cecum Intestine large, colon Intestine large, rectum Intestine small	+ + + + + +	+	+ + + + + + + +	+ + + + + + + + +	++++++	++ ++++	+ + + + + + +	+ + + + + + +	+ + + + + + +	+ + + + + + +	+	+ + + + + + +	+	++++++	+	X + +	++++++	++ ++++	++++++	+	,	++ ++++	+ + + + + + +	+ + + + + +	+ + + + +
Repatoee ular carcinoma		+			+++	+++	+ + +	+++	+ + +	+ + +	++++	+ + +	++++	+++	+ + +	+ + +	++++	+++	++++	+++	++++	+++		+ + X	+++++	+
Pancreas Salivary glands Lymphoma malignant mixed Stomach Stomach Stomach, forestomach Papilioms squamous Stomach, glandular Tooth CARDIOVASCULAR SYSTEM Blood vessel Heart Lymphoma malignant histiccytic ENDOCRINE SYSTEM Adrenai gland Lymphoma malignant mixed Adrenai gland Lymphoma malignant mixed Adrenai gland Lymphoma malignant mixed Adrenai gland Lymphoma malignant mixed Adrenai gland Lymphoma malignant mixed Adrenai gland Lymphoma malignant mixed Adrenai gland Lymphoma malignant mixed Adrenai gland Lymphoma malignant mixed Adrenai gland Lymphoma malignant mixed Adrenai gland Lymphoma malignant mixed Adrenai gland Lymphoma malignant mixed Adrenai gland Lymphoma malignant mixed Adrenai gland Lymphoma malignant mixed Adrenai gland Lymphoma malignant mixed Adrenai gland Lymphoma malignant mixed Adrenai gland Lymphoma malignant mixed Adrenai gland Lymphoma malignant mixed Lymphoma malignant mixed Attack the store of	Hepatocellular adenoma Lymphoma malignant histiocytic	+	X	+	+	+	+	+	+	*X	+ X	+		+	x	X	·	+	+	x	+	X X	+	x	+	+
	Pancreas Salivary glands	++	+	+	+	+	++	+	+	+	++	+ +	+ +	+	+	+	+	+	+	+	+	+	++	+ +	+	
CARDIOVASCULAR SYSTEM	Stomach Stomach, forestomach Papilloma squamous	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+		+	+	+	+
Hoart	Tooth	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Adrenal gland	Blood vessel Heart	++	+	+	+	+	++	++	++	+	+	+	+	+	+	++	+	+	+	+	++	+	+	+	+	
Adrenal gland, cortex	ENDOCRINE SYSTEM Adrenal gland Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+ ¥	+	+	+	+	+	+	+	+	+
Pars distalis, adenoma	Adrenai gland, cortex Adrenai gland, medulia Isiets, pancreatic Parathyroid gland	, M	M	+	+ + + +		+ + + +	+ + + +	+ + + +		+++++	M +	+ M	+		++	+ + + M	M		++++				+ + + + +	+ + M +	+
None	Pars distalis, adenoma Thyroid gland	+	X	+	+	X +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Epididymis	GENERAL BODY SYSTEM None						_									_										
Preputial gland Prostate + + + + + + + + + + + + + + + + + + +	GENITAL SYSTEM Epididymis Lymphoma malignant histocytic	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
	Preputial gland Prostate Seminal vesicle	+	+	+	+	+	+	++	+	+	+	+++	+	+	++	+	+	+	+	+	+	+	+	+	+	+
	Testes Interstitial cell, adenoma	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+

TABLE C2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE MICE: 120 ppm (Continued)

WEEKC ON							-,-			<u> </u>	 ,
WEEKS ON STUDY	1 0	0	0	0	0	0	0	0	0	1	
31001	5	5	5	5	5	5	5	5	5	5	
	ľ	U		•	•		•		•		TOTAL
	4	3	3	3	3	4	3	3	3	4	 TISSUE
CARCASS	1	6	6	7	8	0	7	9	9	0	TUMOR
ID	0	1	4	0	6	8	7	2	8	1	ĺ
	1	1	1	1	1	1	1	1	1	1	
ALIMENTARY SYSTEM	-										
Esophagus	+	+	+	+	+	+	+	+	+	+	60
Gallbladder	+	+	+	+	+	+	+	+	+	+	54
Lymphoma malignant mixed											1
Intestine large	+	+	+	+	+	+	+	+	+	+	60 60
Intestine large, cecum Intestine large, colon	1 1	+	+	+	+	+	+	+	+	+	60
Intestine large, rectum	1 1	_ I	+		T .		Ŧ	Ι	I	- I	59
Intestine small	+	+	÷	+	+	+	+	+	+	+	60
Intestine small, duodenum	+	+	+	+	+	+	+	+	+	+	60
Intestine small, ileum	+	+	+	+	+	+	+	+	+	+	60
Intestine small, jejunum	+	+	+	+	+	+	+	+	+	+	60
Lymphoma malignant mixed											1
Liver	+	+	+	+	+	+	+	+	+	+	60
Hepatocellular carcinoma	x			X			v				8 10
Hepatocellular adenoma Lymphoma malignant histiocytic	I X						X		х		10
Lymphoma malignant mixed									А		2
Pancreas	1 +	_	+	_	_	_	4	_	+	_	60
Salivary glands	+	4	+	+	+	+	+	÷	+	+	60
Lymphoma malignant mixed					X						2
Stomach	+	+	+	+	+	+	+	+	+	+	60
Stomach, forestomach	+	+	+	+	+	+	+	+	+	+	59
Papilloma squamous											1
Stomach, glandular Tooth	+	+	+	+	+	+	+	+	+	+	60
100th		+		+				+			8
CARDIOVASCULAR SYSTEM	-										
Blood vessel	+	+	+	+	+	+	+	+	+	+	60
Heart	+	+	+	+	+	+	+	+	+	+	60
Lymphoma malignant histiocytic	- 1			Х							1
ENDOCRINE SYSTEM	-										
Adrenal gland	+	+	+	+	+	+	+	+	+	+	60
Lymphoma malignant mixed	1.	,				,		,			i
Adrenal gland, cortex	+	+	+	+	+	+	+	+	+	+	60
Adrenal gland, medulla	+	+	+	+	+	+	+	+	+	+	58
Islets, pancreatic	+	+	+	+	+	+	+	+	+	+	60
Parathyroid gland	+	M	M	+	+	M	M	+	M		29
Pituitary gland	+	+	+	+	+	+	+	+	+	+	58
Pars distalis, adenoma Thyroid gland	1 .		4			X	+				60
Folitcie, adenoma	+	+	+	+	+	+	+	+	+	X X	2
										A.	"
GENERAL BODY SYSTEM	_										
None	- 1										1
GENITAL SYSTEM	-										
Epididymis	+	+	+	+	+	+	+	+	+	+	60
Lymphoma malignant histiocytic		,		X		,			X		2
Penis		+		••			+	+	••	+	21
Preputial gland			+							+	9
Prostate	+	+	+	+	+	+	+	+	+	+	60
Seminal vesicle											4
Testes Interstitial cell, adenoma	+	+	+	+	+	+	+	+	+	+	60

TABLE C2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE MICE: 120 ppm (Continued)

WEEKS ON STUDY	0 2 3	0 5 6	0 5 7	0 5 8	0 6 0	0 6 3	0 6 4	0 6 5	0 7 2	0 7 2	0 7 2	0 7 3	0 7 5	0 7 6	0 7 6	0 8 0	0 8 0	0 8 1	0 8 1	0 8 2	0 8 3	0 8 4	0 8 5	0 8 5	0 8 7
CARCASS ID	1 4 1	4 1 9 1	3 8 2 1	4 1 8 1	4 1 3 1	4 1 2 1	3 9 1 1	3 8 1 1	4 1 7 1	3 7 6 1	3 8 5 1	3 7 3 1	3 9 7 1	3 9 5 1	4 0 5 1	3 6 6 1	3 9 3 1	3 6 2 1	3 7 1 1	3 8 4 1	3 9 4 1	3 6 5 1	3 7 5 1	4 0 7 1	3 9 0 1
HEMATOPOIETIC SYSTEM Blood Bone marrow Lymphoma malignant histiocytic	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymph onde Lymph node Axillary, lymphoma malignant mixed Iliac, lymphoma malignant mixed Mediastinal, lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Mesenteric, lymphoma mangnant histocytic Mesenteric, lymphoma malignant mixed Renal, lymphoma malignant mixed Lymph node, mandibular	+	М	М	+	M	+	М	+	М	+	+	+	+	+	М	+	X	+	+	+	+	М	+	X +	+
Lymphoma malignant histocytic Lymphoma malignant mixed Spleen Lymphoma malignant histocytic	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+ X	+	+	+	+	+	+	+	+
Lymphoma malignant mixed Thymus Lymphoma malignant histiocytic Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	M
INTEGUMENTARY SYSTEM Mammary gland Skin Neck, subcutaneous tissue, hemangioma	M +	M +	M +	M +	M +	M +	+	M +	M + X																
MUSCULOSKELETAL SYSTEM Bone Skeletal muscie	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	++	+	+	+	+	+	+
NERVOUS SYSTEM Brain Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
RESPIRATORY SYSTEM Lung Alveolar/bronchiolar adenoma Alveolar/bronchiolar carcinoma Lymphoma malignant histocytic	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed Nose Trachea	++	++	++	+	+	++	++	+	+	+	+	++	+	++	+	+	+	+	+	+	+	+	+	X + +	++
SPECIAL SENSES SYSTEM Hardenan gland Adenoma	_			, X				-		-															
URINARY SYSTEM Kidney Lipoma Lymphoma malignant mixed Ureter	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Urethra Urnary bladder Lymphoma malignant mixed	++	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+

TABLE C2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE MICE: 120 ppm (Continued)

					` -			ueu	,																
WEEKS ON STUDY	8 8	0 8 8	0 8 9	0 9 1	0 9 1	0 9 1	0 9 2	0 9 2	0 9 2	0 9 5	0 9 9	1 0 3	1 0 4	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5
CARCASS ID	4 0 9 1	1 5 1	3 6 8 1	3 7 2 1	3 7 4 1	4 2 0 1	3 8 9	1 1 1	3 6 9	3 8 3 1	3 7 8 1	4 0 0 1	4 0 2 1	3 6 7 1	3 8 0 1	3 9 9	4 0 3 1	4 0 6 1	3 7 9	3 8 7 1	3 8 8 1	3 9 6 1	1 6 1	3 6 3 1	4 0 4 1
HEMATOPOIETIC SYSTEM Blood	-											-													
Bone marrow Lymphoma malignant histocytic	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed Lymph node Axillary, lymphoma malignant mixed Iliac, lymphoma malignant mixed Mediastinal, lymphoma malignant mixed Mesenteric, lymphoma malignant	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	X X X X	+	+	+ X	+	+	+	* X	+	+
histocytic Mesenteric, lymphoma malignant mixed Renal, lymphoma malignant mixed Lymph node, mandibular	+	+	+	+	+	M	+	+	+	+	+	+	X +	+	+	X X +	+	+	X +	+	+	+	X	+	+
Lymphoma malignant histocytic Lymphoma malignant mixed Spleen	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	X	+	+	+	+	+	+	X +	+	+
Lymphoma malignant histocytic Lymphoma malignant mixed Thymus	+	+	+	+	+	+	+	+	+	+	+	+	X	+	+	X	+	+	X	+	+	+	X	+	+
Lymphoma malignant histiocytic Lymphoma malignant mixed																x							x		
INTEGUMENTARY SYSTEM Mammary gland Skin Neck, subcutaneous tissue, hemangioma	M +	M +	M +	M +	M +	M +	++	M +	M +	M +	M +	M +	M +	M +	M +	M +	M +	M +	M +	M +	M +	M +	M +	M +	M +
MUSCULOSKELETAL SYSTEM Bone Skeletal muscie	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
NERVOUS SYSTEM Brain Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	*	+	+
RESPIRATORY SYSTEM Lung Alveolar/bronchiolar adenoma Alveolar/bronchiolar carcinoma	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	X X	+	+
Lymphoma malignant histiocytic Lymphoma malignant mixed Nose Trachea	++	+	++	++	+	+	+	++	++	++	++	++	+	++	++	X + +	+	++	+	++	++	+	X + +	+	++
SPECIAL SENSES SYSTEM Harderian gland Adenoma						* X	* X					-					-								
URINARY SYSTEM Kidney	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lipoma Lymphoma malignant mixed Ureter										X X	+					X							X		
Urnary bladder Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	*	+	+

TABLE C2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE MICE: 120 ppm (Continued)

WEEKS ON STUDY	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	TOTAL
CARCASS ID	4 1 0 1	3 6 1 1	3 6 4 1	3 7 0 1	3 8 6 1	0 8 1	3 7 7 1	3 9 2 1	3 9 8 1	0 1 1	TISSUES
HEMATOPOIETIC SYSTEM Blood Bone marrow Lymphoma malignant histocytic Lymphoma malignant mixed Lymph node Axillary, lymphoma malignant mixed Iliac, lymphoma malignant mixed Mediastinal, lymphoma mal mixed	+	+	+	+ X +	+	+	+	+	+	+	1 60 1 1 60 1 2
Mesentenc, lymphoma malignant histocytic Mesentenc, lymphoma malignant mixed Renai, lymphoma malignant mixed Lymph node mandibular Lymphoma malignant histocytic Lymphoma malignant mixed Spleen Lymphoma malignant mixed Thymus Lymphoma malignant histocytic Lymphoma malignant mixed Lymphoma malignant histocytic Lymphoma malignant histocytic Lymphoma malignant mixed	+ +	+ + +	+ + +	* * * * * * * * * * * * * * * * * * *	X + + X +	+ + +	+ + +	+ + +	+ * +	+ + +	2 6 1 52 1 2 60 3 5 59
INTEGUMENTARY SYSTEM Mammary gland Skin Neck, subcutaneous tissue, hemangioma	M +	M +	M +	M +	M +	M +	M +	++	M +	. M	3 60 1
MUSCULOSKELETAL SYSTEM Bone Skeletal muscle	+	+	+	+	+	+	+	+	+	+	60 2
NERVOUS SYSTEM Brain Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	60
RESPIRATORY SYSTEM Lung Alveolar/bronchiolar adenoma Alveolar/bronchiolar carcinoma Lymphoma malignant histocytic Lymphoma malignant mixed Nose Trachea	+ + +	+ + +	+ + +	+ X + +	+ M +	+ + +	+ + +	+ + +	+ X +	+	60 1 1 2 3 59 60
SPECIAL SENSES SYSTEM Harderian gland Adenoma											3 3
URINARY SYSTEM Kidney Lipoma Lymphoma malignant mixed Ureter Uretra	+	+	+	+	+ X	+	+	+	+	+	60 1 4 1 6
Umnary bladder Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	60

TABLE C2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE: 600 ppm

WEEKS ON STUDY	0 3 3	0 4 0	0 4 0	0 4 4	0 4 6	0 4 7	0 4 8	0 5 4	0 5 5	0 5 6	0 5 8	0 6 0	0 6 1	0 6 2	0 6 2	0 6 4	0 6 4	0 6 5	0 6 5	0 6 8	0 6 8	0 7 0	0 7 2	0 7 3	0 7 4
CARCASS ID	1 2 5 1	1 3 6 1	1 3 2 1	1 8 0 1	1 5 7 1	1 4 4 1	1 3 8 1	1 2 6 1	1 2 3 1	1 6 3 1	1 3 9 1	1 4 1 1	1 3 5 1	1 5 4 1	1 5 8 1	1 5 5 1	1 6 5 1	1 5 3 1	1 4 0	1 7 4 1	1 3 7 1	1 2 9 1	1 4 2 1	1 2 2 1	1 7 5 1
ALIMENTARY SYSTEM Esophagus Gailbladder Intestane large, cecum Intestane large, coion	 + + + +	++++	++++	+ + + + +	+ + + + +	+ + + + +	++++	+ + + +	+ + + +	+ + + + +	+ + + + +	+ + + + +	+ + + + +	+ + + + +	+ + + + +	+ + + + +	+ + + +	+ + + + +	+ M + +	++++	+ + + + +	+ + + + +	+ + + + +	+ + + +	+ + + + +
Lymphoma maingnant lymphocytic Lymphoma maingnant mixed Intestine large, rectum Intestine small Intestine small, loudenum Intestine small, ileum Intestine small, jejunum Liver Hemangioma	+ + + + + + + + +	+ + + + +	+ + + + +	+ + + + +	* + + + + + + + + + + + + + + + + + + +	+ + + + +	+ + + + +	+ + + + +	+ + + + +	+ + + + + +	+ + + + + +	+ + + + +	+ + + + +	+ + + + +	+++++	+ + + + +	+ + + + +	+ + + + +	+ + + + +	+ + + + +	+ + + + +	+ + + + + +	+ + + + +	+ + + + +	+ + + + + +
Hepatocellular carcinoma Hepatocellular adenoma Lymphoma malignant mixed Mesentery					x			x					х				X	x							
Lymphoma malignant mixed Pancreas Lymphoma malignant lymphocytic Lymphoma malignant mixed	+	+	+	+	*	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Salivary glands Lymphoma malignant mixed Stomach Stomach, forestomach	+++	+++	++++	+++	+++	+++	+++	+ + +	+++	+++	+++++	+++	+++	+++	++++	+++	+++	+++	++++	+++	+++	++++	++++	+++	+ + + +
Lymphoma malignant mixed Papilloma squamous Stomach, glandular Lymphoma malignant mixed Tooth	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
CARDIOVASCULAR SYSTEM Blood vessel Heart Lymphoma malignant mixed	+	+	+	+	+++	+	+	+ +	+ +	+++	++	+	++	++	+++	++	+	+++	+	+	++	+ +	+++	++	++
ENDOCRINE SYSTEM Adrenal gland Adrenal gland, cortex Adenoma	 ++	+	+	+	+	++	+	++	+	++	+	+	++	+	+	+	+	+	++	+	+	+	+	+++	M M
Adrenal gland, medulla Islets, pancreatic Adenoma Parathyroid gland	++++++	+ + M	+ + M	+	+ + M	+	+	+	+ + M	++	+	++	+ + M	++	+	+ + M	+	+	++	++	+	+	+ + M	+++++++++++++++++++++++++++++++++++++++	M + M
Pituitary gland Pars distalis, adenoma Thyroid gland	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
GENERAL BODY SYSTEM None	 																								
GENITAL SYSTEM Epididymis Lymphoma malignant mixed Penis Preputial gland	+	+	+	+	+	+	+ + +	+	+	+ + +	+	+	+	+	+	+ + +	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed Prostate Lymphoma malignant mixed Seminal vesicle Testes	+	+	+	+	+	+	+	+ +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed	<i>-</i>	т	.	T		T	~		т	T	т	~	7	T	т.	т.	т		*	,,	T	<i>T</i>	,	· ·	

TABLE C2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE MICE: 600 ppm (Continued)

					(0	OII		ueu	,																
WEERS ON STUDY	0 7 9	0 8 0	0 8 0	0 8 1	0 8 1	0 8 1	0 8 2	0 8 3	0 8 3	0 8 9	0 9 1	0 9 2	0 9 2	0 9 2	0 9 7	0 9 9	1 0 3	1 0 4	1 0 4	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5
CARCASS ID	1 5 1	1 6 6 1	1 6 2 1	1 3 0 1	1 5 9	1 7 3 1	1 7 9 1	1 4 3 1	1 3 1 1	1 2 8 1	1 4 5 1	1 3 4 1	1 6 9 1	1 7 1	1 2 7 1	1 2 4 1	1 4 8 1	1 4 7 1	1 7 2 1	1 5 0 1	1 6 7 1	1 7 6 1	1 7 8 1	1 3 3 1	1 6 8 1
ALIMENTARY SYSTEM		_																							
Esophagus Gallbladder	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Galibladder Intestine large	+	+	+	+	M +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine large, cecum	1 +	+	+	+	+	+	+	+	+	+	+	÷	÷	÷	÷	+	÷	+	+	+	+	+	+	+	÷
Intestine large, colon Lymphoma malignant lymphocytic Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+ X	+
Intestine large, rectum	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine small	+ +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine small, duodenum Intestine small, ileum	‡	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine small, jejunum	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Liver Hemangioma	+	+	+	+	+	+	+	+	+	+	+	*X	+	+	+	+	+	+	+	+	+	+	+	+	+
Hepatocellular carcinoma Hepatocellular adenoma	x				X		x		x						X	X		X				X			
Lymphoma malignant mixed Mesentery					+							X													
Lymphoma malignant mixed	+				+																				
Pancreas	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant lymphocytic Lymphoma malignant mixed												X													
Salivary glands	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed Stomach	1.		_	_	_	_	_	_	_	_	_	X +	+		_	_	_	_	_	_	_	_	4	_	_
Stomach, forestomach	Ţ	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed Papilloma squamous																X									
Stomach, glandular Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	7	+
Tooth											+														
CARDIOVASCULAR SYSTEM																		_			_				
Blood vesser Heart	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed	'		•		,	,	7	-	7	-	_	т.	-	т	-	,		'	,	,	,	,			,
ENDOCRINE SYSTEM																									
Adrenal gland	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Adrenal gland, cortex Adenoma	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	X ⁺	+	+	+	+
Adrenal gland, medulla	+	+	+	+	+	+	+	+	M	+	+	+	+	+	+	+	+	+	+	+	A.	+	+	+	+
Islets, pancreatic	+	+	+	-	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Adenoma Parathyroid gland	М	М	М	M	М	M	М	М	+	+	+	+	+	+	М	М	M	М	+	М	M	M	M	M	+
Pituitary gland	+	+	+	+	+	+	+	+	+	M	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Pars distalis, adenoma Thyroid gland		_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_		_	_	+	+	4
· •								-				,	,				'								
GENERAL BODY SYSTEM None																									
GENITAL SYSTEM Epididymis			_						_				_			+			+				+	+	
Lymphoma malignant mixed	1	~	7	_	т	_	Τ.	7	-	7		~	7	7	7	,	*	7	-	ſ	r	,	,	*	
Penis	+		+		+		+		+	+	+		+		+		+								
Preputial gland Lymphoma malignant mixed	i																+	+	+						
Prostate	+	+	+	+	+	+	+	M	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed Seminal vesicle						_																			
Testes	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed																									

TABLE C2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE MICE: 600 ppm (Continued)

								``	•			
WEEKS ON	1	1	1	1	1	1	1	1	1	1		
STUDY	0 5	0 5										
	9	5	Э	Э	Э	Э	Э	Э	Э	Э		TOTAL
	1	1	1	1	1	1	1	1	1	1		TISSUES
CARCASS	7	4	5	6	6	5	6	2	4	7		TUMORS
ID	7	9	6 I	0	1	2	4	1	6	0		j j
	1	1	1	1	ı	T	1	i	1	1		
ALIMENTARY SYSTEM												
Esophagus	+	+	+	+	+	+	+	+	+	+		60
Gallbladder Intestine large	+	+	+	+	M +	+	+	+	+	+		57 60
Intestine large Intestine large, cecum	+	+	+	+	+	+	+	+	+	+		60
Intestine large, colon	+	+	+	+	+	+	+	+	+	+		60
Lymphoma malignant lymphocytic	1											1 2
Lymphoma malignant mixed Intestine large, rectum	+	+	+	+	+	+	4	+	X +	+		60
Intestine small	+	+	+	+	+	+	+	+	+	+		60
Intestine small, duodenum	+	+	+	+	+	+	+	+	+	+		60
Intestine small, ileum Intestine small, jejunum	++	+	+	+	+	+	+	+	+	+		60 60
Liver	+	+	+	+	+	+	+	+	+	+		60
Hemangioma												1
Hepatocellular carcinoma	- 1	Х			17	•		.,		**		9
Hepatocellular adenoma Lymphoma malignant mixed					X	X		Х	X	X		9 2
Mesentery							+		**			2
Lymphoma malignant mixed	1						Х					1
Pancreas	+	+	+	+	+	+	+	+	+	+		60
Lymphoma malignant lymphocytic Lymphoma malignant mixed	x								х			1 3
Salivary glands	+	+	+	+	+	+	+	+	+	+		60
Lymphoma malignant mixed	Ι.						X		X			3
Stomach Stomach, forestomach	+	+	+	+	+	+	+	+	+	+		60 60
Lymphoma malignant mixed	1 '	7	7	,	,	,	7	7	X	7		1
Papilioma squamous												1
Stomach, glandular	+	+	+	+	+	+	+	+	X	+		60
Lymphoma malignant mixed Tooth									^			2
CARDIOVASCULAR SYSTEM Blood vessel												60
Heart	1 7	+	+	+	+	+	+	+	+	+		60 60
Lymphoma malignant mixed									X			1
	-											
ENDOCRINE SYSTEM Adrenal gland	+	+	+	+	4	+	+	+	+	+		59
Adrena: gland, cortex	+	+	+	+	+	+	+	+	+	+		59
Adenoma												1
Adrenal gland, medulla Islets, pancreatic	1 ‡	+	+	+	+	+	+	+	+	+		58 60
Adenoma	"	-				X	τ.			т		1
Parathyroid gland	M	M		M	M	M	M		M			25
Pituitary gland	+	+	+	+	+	+	M	+	+	+		58
Pars distalis, adenoma Thyroid gland	1	_	_	X	+	+	_	+	+	+		60
· -	1 '	•	•			,			,			00
GENERAL BODY SYSTEM												
None												
GENITAL SYSTEM	-		-									
Epididymis	+	+	+	+	+	+	+	+	+	+		60
Lymphoma malignant mixed Penis	- 1								X			17
Penis Preputial gland								+	+			10
Lymphoma malignant mixed									X			1
Prostate	+	+	+	+	+	+	+	+	+	+		59
Lymphoma malignant mixed Seminal vesicle									X			1 4
Testes	+	+	+	+	+	+	+	+	+	+	-	60
Lymphoma malignant mixed									X			i
	_											

TABLE C2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE MICE: 600 ppm (Continued)

WEEKS ON STUDY	0 3 3	0 4 0	0 4 0	0 4 4	0 4 6	0 4 7	0 4 8	0 5 4	0 5 5	0 5 6	0 5 8	0 6 0	0 6 1	0 6 2	0 6 2	0 6 4	0 6 4	0 6 5	0 6 5	0 6 8	0 6 8	0 7 0	0 7 2	0 7 3	0 7 4
CARCASS ID	1 2 5 1	1 3 6 1	1 3 2 1	1 8 0 1	1 5 7 1	1 4 4 1	1 3 8 1	1 2 6 1	1 2 3 1	6 3 1	1 3 9 1	1 4 1	1 3 5 1	1 5 4 1	1 5 8 1	1 5 5	1 6 5 1	1 5 3 1	1 4 0 1	7 4 1	1 3 7 1	1 2 9 1	1 4 2 1	1 2 2 1	7 5 1
HEMATOPOIETIC SYSTEM Blood Bone marrow Lymph node Hepatocellular carcinoma, metastatic, liver Iliac, lymphoma malignant mixed	+ +	+	++	+ +	+ +	+ +	+ +	+ +	++	+ +	++	++	+ +	+ +	+ +	+ +	+ +	++	++	+ +	+	+ +	+ +	+	+ A
Mediastinal, lymphoma malignant mixed Mesenteric, lymphoma malignant lymphocytic Mesenteric, lymphoma malignant mixed Renal, lymphoma malignant mixed Lymph node, mandibular Lymphoma malignant lymphocytic	+	м	М	+	x M	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	A
Lymphoma malignant mixed Spieen Hemangiosarcoma Lymphoma malignant mixed Thymus Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	A +
INTEGUMENTARY SYSTEM Mammary gland Skin Lymphoma malignant mixed	M +	M +	M +	M +	M +	M +	M +	M +	M +	M +	M +	+ +	M +	M +	M +	M +	M +								
MUSCULOSKELETAL SYSTEM Bone Skeletai muscle Head, lymphoma malignant mixed	+	+++	+	+	+	+	+	+	++	+	+	+++	+	+	+	+	+	+	+	+	+	+	+	+	+
NERVOUS SYSTEM Brain Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
RESPIRATORY SYSTEM Lung Adenocarcinoma, metastatic, hardenan gland Alveoiar/bronchiolar adenoma Hepatocellular carcinoma, metastatic, liver	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+ X	+	+	+	+	+	+	+
Lymphoma malignant mixed Nose Adenocarcinoma, metastatic, harderian gland Mucosa, lymphoma malignant mixed Trachea	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
SPECIAL SENSES SYSTEM Hardenan giand Adenocarcinoma																									
URINARY SYSTEM Kidney Hepatocellular carcinoma, metastatic, hver Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Ureter Urethra Urnary bladder Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+

TABLE C2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE MICE: 600 ppm (Continued)

WEEKS ON STUDY	0 7 9	0 8 0	0 8 0	0 8 1	0 8 1	0 8 1	0 8 2	0 8 3	0 8 3	0 8 9	0 9 1	0 9 2	9	0 9 2	0 9 7	9	1 0 3	1 0 4	1 0 4	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5
CARCASS ID	1 5 1 1	6 6 1	1 6 2 1	1 3 0 1	1 5 9 1	1 7 3 1	7 9 1	1 4 3 1	1 3 1	1 2 8 1	1 4 5 1	1 3 4 1	6 9 1	7 1 1	1 2 7 1	1 2 4 1	1 4 8 1	1 4 7 1	1 7 2 1	1 5 0 1	1 6 7 1	1 7 6 1	7 8 1	1 3 3 1	6 8 1
HEMATOPOIETIC SYSTEM Blood Bone marrow Lymph node Hepatocellular carcinoma, metastatic, liver Iliac, lymphoma malignant mixed	+++	+ +	++	+	+	++	+++	+	+	+ +	++	+ +	++	+ +	+ +	+	+ +	+	+ + +	+	+ +	+ + X	++	+++	+ +
Mediastinal, lymphoma malignant mixed Mesenteric, lymphoma malignant lymphocytic Mesenteric, lymphoma malignant mixed Renal, lymphoma malignant mixed Lymph node, mandibular	+	+	+	+	+	+	+	+	+	+	+	x x +	+	М	+	+	+	M	+	+	+	+	+	x	+
Lymphoma malignant lymphocytic Lymphoma malignant mixed Spleen Hemangiosarcoma Lymphoma malignant mixed Thymus	+	+	+	+	+	+ M	+	+	+	+	+	X + X +	+	+	+	+	+	+	+	+	+	* *	+	+ X +	+
Lymphoma malignant mixed INTEGUMENTARY SYSTEM Mammary gland Skin Lymphoma malignant mixed	+++	M +	M +	M +	M +	M +	M +	M +	M +	M +	M +	Х М +	M +	M +	++	M +	M +	Х М +	M +						
MUSCULOSKELETAL SYSTEM Bone Skeletal muscle Head, lymphoma malignant mixed	+	+	+	+	++	+	+	+	+	+	++	+	+	+	+	+	+	+	+	+	+	+	+	+	+
NERVOUS SYSTEM Brain Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
RESPIRATORY SYSTEM Lung Adenocarcinoma, metastatic, harderian gland Alveolar/bronchiolar adenoma Hepatocellular carcinoma, metastatic, liver Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	+	+ X	+	+	+	+	+	+ X	+	+	+	+	+	+	+ x
Nose Adenocarcinoma, metastatic, harderian gland Mucosa, lymphoma malignant mixed Trachea	+	+	+	+	+	+	+	+	+	+	+	+ X +	+	+	+	+	+	+	+	+	+	+	+	+	+ +
SPECIAL SENSES SYSTEM Harderian gland Adenocarcinoma	_											+ X						-							
URINARY SYSTEM Kidney Hepatocellular carcinoma, metastatic, liver Lymphoma malignant mixed Ureter	+	+	+	+	+	+	+	+	+	+	+	+ X	+	+	+	+	+	+	+	+	+	+ X	+	+	+
Urethra Urnary bladder Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	*	+

TABLE C2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE MICE: 600 ppm (Continued)

WEEKS ON STUDY	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	TOTAL.
CARCASS ID	7 7 1	1 4 9 1	1 5 6 1	1 6 0 1	1 6 1	1 5 2 1	1 6 4 1	1 2 1 1	1 4 6 1	1 7 0 1	TISSUES TUMORS
HEMATOPOIETIC SYSTEM											
Blood Bone marrow Lymph node Hepatocellular carcinoma, metastatic,	++	+	+	+	+	+	+	+	+	+	60 59
liver Iliac, lymphoma malignant mixed Mediastinal, lymphoma mali mixed Mesenteric, lymphoma malignant							X X				1 1 2
lymphocytic Mesentenic, lymphoma malignant mixed Renal, lymphoma malignant mixed Lymph node, mandibular Lymphoma malignant lymphocytic	x +	+	+	+	*	+	X X +	x + x	+	+	1 6 1 54 1
Lymphoma malignant mixed Spieen Hemangiosarcoma Lymphoma malignant mixed	+ X	+	+	+	+	+	^ + X	т + Х	х + х	+	60
Thymus Lymphoma malignant mixed	+	+	+	+	+	+	+	M	X	+	58 3
INTEGUMENTARY SYSTEM Mammary gland Skin Lymphoma malignant mixed	M +	M +	M +	M +	M +	M +	M +	M +	M + X	M +	3 60 1
MUSCULOSKELETAL SYSTEM Bone Skeletal muscle Head, lymphoma malignant mixed	+	+	+	+	+	+	+	+	+ + X	+	60 6 1
NERVOUS SYSTEM Brain Lymphoma malignant mixed	+	+	+	+	+	+	+	+	*	+	60
RESPIRATORY SYSTEM Lung Adenocarcinoma, metastatic, harderian gland Alveolar/bronchiolar adenoma	+	+	+	+	+	+	+	+	+	+	 60
Hepatocellular carcinoma, metastatic, liver Lymphoma malignant mixed Nose Adenocarcinoma, metastatic, hardenan	+	+	+	+	+	+	X +	+	X +	+	1 2 60
gland Mucosa, lymphoma malignant mixed Trachea	+	+	+	+	+	+	+	+	X +	+	60
SPECIAL SENSES SYSTEM Hardenan gland Adenocarcinoma											1 1
URINARY SYSTEM Kidney Hepatocellular carcinoma, metastatic,	+	+	+	+	+	+	+	+	+	+	60
liver Lymphoma malignant mixed Ureter Urethra							X		X		1 3 3
Urinary bladder Lymphoma malignant mixed	+	+	+	+	+	+	*	+	*	+	60 3

TABLE C2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE: 1,200 ppm

	 							_					_			-	_			_		^		À	_
WEEKS ON STUDY	0 0 5	0 2 3	0 4 2	0 4 3	0 4 8	0 5 2	0 5 2	0 5 3	0 6 1	0 6 1	0 6 2	0 6 4	0 6 6	0 6 7	0 6 7	0 7 3	0 7 5	0 7 5	0 7 6	0 7 6	0 7 9	0 7 9	0 8 1	0 8 1	8 2
CARCASS	2 6	6	2 4	5	2 4	2 4	2 4	5	5	2 5	7	2 5	2 5	9	2 8	7	2 7	2 9	2 8	9	6	9	8	8	6
ID	3 1	9 1	1	8 1	6 1	2 1	4 1	2 1	4 1	6 1	9 1	9 1	7 1	3	0 1	7 1	6 1	9	8	7 1	1	5 1	1	$\frac{2}{1}$	0 1
ALIMENTARY SYSTEM	 +								_	+	+	+	+			_	_	+	+				_	+	+
Esophagus Galibiadder Lymphoma malignant undifferentiated cell type	+	+	Ŧ	+	+	+	+	+	+	÷	+	÷	+	÷	÷	÷	÷	+	÷	M	+	+	+	+	+
Intestine large Intestine large, cecum Lymphoma malignant undifferentiated	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+ A	+	+	+	+	+
cell type Intestine large, colon	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	A	+	+	+	+	+
Intestine large, rectum Intestine small	+	++	+	+	+	+	++	+	+	+	++	+	++	+	+	++	++	+	+	M +	+	+	+	+	+
ntestine small, duodenum	+++++++++++++++++++++++++++++++++++++++	++	+	++	++	++	+	+	+	++	++	+	+	+	+	+	+	+	+	+	M	+	+	+	+
intestine small, ileum intestine small, jejunum Lymphoma malignant undifferentiated cell type	+	+	+	+	+	+	+	+	+	+	+	+	+	Ŧ	+	+	+	+	÷	+	+	+	+	+	+
liver	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	A	+	+	+	+	+
Hemangiosarcoma Hemangiosarcoma, metastatic, spieen Hepatocellular carcinoma				х				x					x					х					x		х
Hepatocellular adenoma Hepatocellular adenoma, multiple Histocytic sarcoma Ito cell tumor malignant Lymphoma malignant undifferentiated cell type Mesentery																		Α					Α		Α.
Lymphoma malignant undifferentiated cell type Pancreas	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	Δ	+	+	+	+	+
Lymphoma malignant undifferentiated cell type Salivary glands													· •					+		Α	+	+	+	+	+
Stomach Stomach, forestomach	+	+	+	+	++	++	++	++	++	++	+	+	++	+	+	+	+	+	+	+	+	+	+	+	+
stomach, glandular Pooth Pulp, lymphoma malignant undifferentiated cell type	+	Ŧ	+	Ŧ	+	Ŧ	+	+	+	7	+	Ŧ	+	+	+	+	÷	+	+	+	+	+	+	+	+
CARDIOVASCULAR SYSTEM Blood vessel	 																								
Heart Hemangnosarcoma	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	M	+	+	+
ENDOCRINE SYSTEM	 																								
Adrenal gland Adrenal gland, cortex	++	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+ A	+	+	+	+	+
Adrenai gland, medulla	+	+	+	+	+	+	+	++	+	++	+	+	+	+	+	+	+	+	+	A	+	+	+	+	+
Islets, pancreatic Adenoma													_									Ţ,			
Parathyroid gland Pituitary gland	++	M +	M +	M +	+	+	M +	+	M +	M M	M +	+	+	+	M +	+	M +	M +	+	M M	M +	M +	+	+	+
Pars intermedia, adenoma Phyroid gland	+	+	+	+	+	+	+	+	+	M	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
GENERAL BODY SYSTEM None	 														-					•	_				
GENITAL SYSTEM	 <u> </u>						_																		
Epididymis Penis	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	-	+	+			+	_	7	7	+
Preputial gland Prostate Lymphoma malignant undifferentiated	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	M
cell type Seminal vesicle								+			+										+	+			
Testes Lymphoma malignant undifferentiated cell type Interstitial cell, adenoma	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+

TABLE C2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE MICE: 1,200 ppm (Continued)

WEEKS ON STUDY	0 8 2	0 8 2	0 8 4	0 9 1	0 9 2	0 9 2	0 9 2	0 9 5	0 9 8	0 9 9	1 0 0	1 0 1	1 0 1	1 0 3	1 0 3	1 0 3	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5
CARCASS ID	2 8 4 1	2 8 7 1	5 0 1	9 0 1	9 6 1	6 5 1	2 7 2 1	2 4 8 1	2 7 1 1	2 4 7 1	2 9 4 1	2 5 1	2 4 3 1	2 8 3 1	2 7 8 1	2 7 3 1	2 5 5 1	2 6 4 1	7 4 1	2 8 5 1	9 1 1	2 4 5 1	2 4 9 1	2 6 7 1	7 0 1
ALIMENTARY SYSTEM Esophagus Gallbladder	++	++	++	++	++	++	++	M +	++	<i>+</i> +	++	++	M +	+	++	+	++	++	++	+	++	++	++	+	++
Lymphoma malignant undifferentiated cell type Intestine large	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	X +	+	+	+	+
Intestine large, cecum Lymphoma malignant undifferentiated cell type	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+ X	+	+	+	+
Intestine large, colon	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine large, rectum Intestine small	++	+	+	+	+	+	+	+	M +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine small, duodenum	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine small, ileum	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine small, jejunum Lymphoma malignant undifferentiated cell type	+	+		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	X	+	+	+	+
Liver Hemangiosarcoma Hemangiosarcoma, metastatic, spieen	+	+	+	+	X	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	
Hepatocellular carcinoma Hepatocellular adenoma Hepatocellular adenoma, multiple		x	X			X					X	x	X	X	X							Х			X
Histocytic sarcoma to cell tumor malignant Lymphoma malignant undifferentiated cell type Mesentery										X									x		X +				
Lymphoma malignant undifferentiated cell type Pancreas																	1				x				+
Lymphoma malignant undifferentiated cell type	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	X		+	+	
Salivary glands Stomach	1 ‡	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Stomach, forestomach	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Stomach, glandular Tooth Pulp, lymphoma malignant undifferentiated cell type	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+ + X	+	+	+	+
CARDIOVASCULAR SYSTEM																									
Blood vessel	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Heart Hemangiosarcoma	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
ENDOCRINE SYSTEM														· .											+
Adrenal gland Adrenal gland, cortex	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	Ŧ	+	+
Adrenal gland, medulla	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Islets, pancreatic Adenoma	+	+	+	*X	+	+	+	+	+	+	+	+	+	+	M	+	+	+	+	+	+	+	+	+	+
Parathyroid gland	+	+	M	M	+	M	+	M	M	M	M	M	+	+	M	+	M	М	M	M	M		M	+	М
Pituitary gland	+	+	+	+	M	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	M	+	+	+	+
Pars intermedia, adenoma Thyroid gland	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	X +	+	+	+	+	+	+	+	+
GENERAL BODY SYSTEM None			-																						
GENITAL SYSTEM																									
Epididymis Penis	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Preputial gland	7	7	+		+			+	+		7		7	7	*	7									+
Prostate Lymphoma malignant undifferentiated cell type	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+ X	+	+	+	+
Seminal vesicle Testes	1,		_	_	_			_	_	+	_		+		_	_	_	_	_		٠. ـ	4		_	د
Lymphoma malignant undifferentiated	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	X	+	+	+	+

TABLE C2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE MICE: 1,200 ppm (Continued)

								` -			ueu)	
WEEKS ON STUDY	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5		TOTAL
CARCASS ID	2 7 5 1	2 5 3 1	2 8 9 1	2 6 2 1	2 6 6 1	2 6 8 1	9 2 1	2 8 6 1	9 8 1	3 0 0 1		TISSUES TUMORS
ALIMENTARY SYSTEM Esophagus Gallbladder Lymphoma malignant undifferentiated	+ +	+	++	+	+	+	++	+	+ +	+		57 59
cell type Intestine large Intestine large, cecum Lymphoma malignant undifferentiated	+	+	+	+	+	+	+	+	+	++		60 59
cell type Intestine large, colon Intestine large, rectum Intestine small Intestine small, duodenum Intestine small, ileum Intestine small, jejunum Lymphoma malignant undifferentiated cell type	+ + + + + +	+ + + + +	+ + + + +	+ + + + +	+ + + + + +	+ + + + +	+++++	+ + + + + +	+ + + + + +	+ + + + +		1 59 58 60 59 60 60
Liver Hemangiosarcoma Hemangiosarcoma, metastatic, spieen Hepatoceilular carcinoma Hepatoceilular adenoma Hepatoceilular adenoma, multiple Histiocytic sarcoma Ito cell tumor malignant Lymphoma malignant undifferentiated cell type	x	x	*	* X	+	+	+	+	+	+		59 1 1 8 10 1 1 1
Mesentery Lymphoma malignant undifferentiated cell type Pancreas Lymphoma malignant undifferentiated cell type	+	+	+	+	+	+	+	+	+	+		1 59
Salivary glands Stomach Stomach, forestomach Stomach, glandular Tooth Pulp, lymphoma malignant	+++++	+ + + +	+ + + +	+ + + +	++++	+ + + +	++++	+ + + +	+ + + +	+ + + +		59 60 60 60 1
undifferentiated cell type CARDIOVASCULAR SYSTEM Blood vessel Heart Hemangiosarcoma	<u>+</u>	++	++	++	++	++	++	+ + X	+	+ +		60 59 1
ENDOCRINE SYSTEM Adrenal gland, cortex Adrenal gland, medulla Islets, pancreatic Adenoma Parathyroid gland Pituitary gland Pars intermedia, adenoma Thyroid gland	+ + + + + M +	+ + + + + + + + + + + + + + + + + + + +	+ + + + + + M +	++++	+ + + + + +	+ + + + + +	+ + + + + M	+ + + + + + +	+ + + + + + + +	++++++++		60 59 59 58 1 24 56 1 59
GENERAL BODY SYSTEM None	-	_										39
GENITAL SYSTEM Epididymis Penis Preputial gland Prostate Lymphoma malignant undifferentiated cell type Seminal vesicle	+++++++++++++++++++++++++++++++++++++++	+	+	+	+	+	+	+	+ + + +	+		60 21 11 59
Testes Lymphoma malignant undifferentiated cell type Interstitial cell, adenoma	+	+	+	+	+	+	+	+	+	+		1 1

TABLE C2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE MICE: 1,200 ppm (Continued)

					(0	OH	LIKI	uea	.,																
WEEKS ON STUDY	0 0 5	0 2 3	0 4 2	0 4 3	0 4 8	0 5 2	0 5 2	0 5 3	0 6 1	0 6 1	0 6 2	0 6 4	0 6 6	0 6 7	0 6 7	0 7 3	0 7 5	0 7 5	0 7 6	0 7 6	0 7 9	0 7 9	0 8 1	0 8 1	0 8 2
CARCASS ID	6 3 1	2 6 9 1	2 4 1	2 5 8 1	2 4 6 1	2 4 2 1	2 4 4 1	2 5 2 1	2 5 4	2 5 6 1	7 9 1	2 5 9	2 5 7 1	2 9 3 1	2 8 0 1	7 7 1	2 7 6 1	2 9 9	2 8 8 1	9 7 1	2 6 1	9 5 1	2 8 1 1	2 8 2 1	2 6 0 1
HEMATOPOIETIC SYSTEM Bone marrow	- -	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	A	+	+	+	+	+
Hemangosarcoma, metastatic, spleen Lymph node liac, lymphoma malignant undifferentiated cell type Mediastinal, histocytic sarcoma Mediastinal, lymphoma malignant undifferentiated cell type Mesenteric, histocytic sarcoma Mesenteric, lymphoma malignant mixed Mesenteric, lymphoma malignant undifferentiated cell type Renal, histocytic sarcoma Renal, histocytic sarcoma Renal, hymphoma malignant	+	+	+	+	+	+	+	+	A	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
undifferentiated cell type Lymph node, mandibular Lymphoma malignant mixed Lymphoma malignant undifferentiated cell type	М	+	+	M	+	M	+	+	A	M	+	+	+	+	+	+	+	+	+	+	+	M	+	+	+
Spieen Hemangiosarcoma Histiocytic sarcoma Lymphoma malignant mixed Lymphoma malignant undifferentiated cell type	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	A	+	+	+	+	+
Thymus Histocytic sarcoma Lymphoma malignant mixed Lymphoma malignant undifferentiated cell type	+	+	+	+	+	+	+	+	+	M	+	+	+	+	M	+	+	+	+	A	+	+	+	+	M
INTEGUMENTARY SYSTEM Mammary gland Skin	M +	. +	M +	. M	M +	M +	M +	M +	+	M +	M +	M +	M +	M +	M +	M +	M +	M +	M +	M A	M +	M +	M +	M +	M +
MUSCULOSKELETAL SYSTEM Bone Hemangiosarcoma, metastatic, spleen Skeletal muscle	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
NERVOUS SYSTEM Brain	_ -	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
RESPIRATORY SYSTEM Lung Alveolar/bronchiolar adenoma Alveolar/bronchiolar adenoma, multiple Alveolar/bronchiolar carcinoma Hepatocellular carcinoma, metastatic, liver Histocytic sarcoma Lymphoma malignant undifferentiated	+	+	+	+	+	+	+	*X	+	+	+	+	+	+ X	+	+	+	+	+	+	+	+	*X	+	* *
cell type Nose Submucosa, lymphoma malignant	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	A	+	+	+	+	+
undifferentiated cell type Trachea SPECIAL SENSES SYSTEM	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	A	+	+	+	+	+
Harderian gland Adenoma																									
URINARY SYSTEM Kidney Lymphoma malignant mixed Urethra Urnnary bladder Lymphoma malignant undifferentiated cell type	+	+	+	+ +	+	+	+	+	+	+	+	+	+	+	+	+++	+	+	+	A +	+	+	+	+	+

TABLE C2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE MICE: 1,200 ppm (Continued)

				(C	on	uni	nea	()																
0 8 2	0 8 2	0 8 4	0 9 1	0 9 2	0 9 2	0 9 2	0 9 5	0 9 8	0 9 9	1 0 0	1 0 1	1 0 1	1 0 3	1 0 3	1 0 3	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5
2 8 4 1	2 8 7 1	5 0 1	9 0 1	2 9 6 1	6 5 1	7 2 1	2 4 8 1	2 7 1 1	2 4 7 1	2 9 4 1	5 1 1	2 4 3 1	2 8 3 1	2 7 8 1	2 7 3 1	2 5 5 1	2 6 4	7 4 1	2 8 5 1	9 1 1	2 4 5 1	2 4 9 1	6 7 1	7 0 1
-																								
+	+	+	+	+	+	+	+	+	+ + X	+	+	+	+	+	+	+	+	+	+	+ + X	+	+	+	+
									X X											x x	x			x
+	+	+	+	+	+	+	+	+	+	+	М	M	+	+	+	+	+	+	+	X +	+	+	+	* X
+	+	+	+	+	+	+	+	+	+ X	+	+	+	+	+	+	+	+	+	+	X +	+	+	+	+
+	+	+	+	+	+	+	M	+	*	+	+	М	+	+	+	+	+	+	+	X +	+	+	+	* ** ** ** ** ** ** ** ** ** ** ** ** *
_																								
M +	M +	M +	M +	M +	M +	M +	M +	M +	M +	M +	M +	M +	M +	M +	M +	M +	M M		M +	M +	M +	M +	M +	M +
+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
-	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
+	+	+ X	+	+	+	+	* X	+	+	+	+	+	+	+	+	*	+	+	* X	+	+	+	+	+
									x	x										x				
+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+ X +	+	+	+	+
-			+ X							+												<u></u>	* X	
+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+ X
	8 2 2 8 4 1 + + + + + + + + + + + + + + + + + +	8 8 2 2 8 8 8 4 7 1 1 1 + + + + + + + + + + + + + + + +	8 8 8 2 2 4 2 2 8 8 7 0 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	8 8 8 9 2 2 4 1 2 2 2 8 8 5 9 4 7 0 0 0 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	M M M M M M + + + + + + + + + + + + + +	M M M M M M M H + + + + + + + + + + + +	0	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	8 8 8 9 9 9 9 8 8 2 2 2 2 2 2 2 2 2 8 1 1 1 1 1 1 1 1 1 1 1 1 1 + + + + + +	0	0	0	0	0	0	0	0	0	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1 1 1 1	0	0	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1 1 1	0	0

TABLE C2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF MALE MICE: 1,200 ppm (Continued)

	,										
WEEKS ON STUDY	0 5	1 0 5	0 5	0 5	0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	mom a t
CARCASS ID	7 5 1	2 5 3 1	2 8 9	2 6 2 1	2 6 6 1	2 6 8 1	2 9 2 1	2 8 6 1	9 8 1	3 0 0	TOTAL: TISSUES TUMORS
HEMATOPOIETIC SYSTEM Bone marrow Hemangrosarcoma, metastatic, spleen Lymph node Iliac, lymphoma malignant undifferentiated cell type Mediastinal, histocytic sarcoma Mediastinal, lymphoma malignant undifferentiated cell type	+ +	+	+	+ X +	+	+	+ +	+	+	+	59 1 59 1 1
Mesenteric, histiocytic sarcoma Mesenteric, lymphoma malignant mixed Mesenteric, lymphoma malignant undifferentiated cell type Renal, histiocytic sarcoma Renal, lymphoma malignant undifferentiated cell type	x										1 1 1
Lymph node, mandibular Lymphoma malignant mixed Lymphoma malignant undifferentiated cell type	x x	+	+	+	+	+	+	+	+	M	51 2 1
Spleen Hemangiosarcoma Histiocytic sarcoma Lymphoma malignant mixed Lymphoma malignant undifferentiated	+ X	+	*	*	+	+	*X	+	+	+	59 3 1 3
cell type Thymus Histocytic sarcoma Lymphoma malignant mixed Lymphoma malignant undifferentiated cell type	+	+	+	+	+	+	+	+	+	+	1 54 1 1
INTEGUMENTARY SYSTEM Mammary gland Skin	M +	M +	M +	++	M +	M +	M +	M +	M +	M +	3 58
MUSCULOSKELETAL SYSTEM Bone Hemangiosarcoma, metastatic, spieen Skeletal muscle	+	+	+	+ X	+	+	+	+	+	+	60 1 3
NERVOUS SYSTEM Brain	+	+	+	+	+	+	+	+	+	+	60
RESPIRATORY SYSTEM Lung Alveolar/bronchiolar adenoma Alveolar/bronchiolar adenoma, multiple Alveolar/bronchiolar aercinoma Hepatocellular carcinoma, metastatic,	+	+	+	+	+	+	+	+	*	+	60 7 1 1
liver Histocytic sarcoma Lymphoma malignant undifferentiated cell type Nose	+	+	+	+	+	+	+	+	+	+	1 1 59
Submucosa, lymphoma malignant undifferentiated cell type Trachea	+	+	+	+	+	+	+	+	+	+	1 59
SPECIAL SENSES SYSTEM Hardenan gland Adenoma											3 2
URINARY SYSTEM Kidney Lymphoma malignant mixed Urethra Urinary bladder Lymphoma malignant undifferentiated	+	+	+	+	+	+	+	+	+	+	59 1 3 59
cell type											1

TABLE C3. ANALYSIS OF PRIMARY TUMORS IN MALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE

	Chamber Control	120 ppm	600 ppm	1,200 ppm
Harderian Gland: Adenoma				· · · · · · · · · · · · · · · · · · ·
Overall Rates (a)	0/60 (0%)	3/60 (5%)	0/60 (0%)	2/60 (3%)
Adjusted Rates (b)	0.0%	8.1%	0.0%	8.2%
Terminal Rates (c)	0/17 (0%)	0/22 (0%)	0/16 (0%)	1/19 (5%)
Day of First Observation	0/11 (0 /0)	400	0/10(0/0/	633
Life Table Tests (d)	P = 0.491	P=0.152	(e)	P = 0.275
Logistic Regression Tests (d)	P = 0.472	P = 0.102	(e)	P = 0.253
Cochran-Armitage Trend Test (d)	P = 0.477	1 -0.107	(6)	1 -0.200
Fisher Exact Test (d)	1-0.4//	P = 0.122	(e)	P = 0.248
Harderian Gland: Adenoma or Adenoc	arcinoma			
Overall Rates (a)	0/60 (0%)	3/60 (5%)	1/60 (2%)	2/60 (3%)
Adjusted Rates (b)	0.0%	8.1%	4.2%	8.2%
Terminal Rates (c)	0/17 (0%)	0/22 (0%)	0/16 (0%)	1/19 (5%)
Day of First Observation	3,21 (3,0)	400	639	633
Life Table Tests (d)	P = 0.459	P = 0.152	P=0.500	P=0.275
Logistic Regression Tests (d)	P = 0.431	P = 0.102	P = 0.485	P = 0.253
Cochran-Armitage Trend Test (d)	P=0.437	- = 0.201	01400	- 0.200
Fisher Exact Test (d)	1 -0.401	P = 0.122	P = 0.500	P = 0.248
Liver: Hepatocellular Adenoma				
Overall Rates (f)	7/60 (12%)	10/60 (17%)	9/60 (15%)	11/59 (19%)
Adjusted Rates (b)	24.1%	33.2%	34.4%	34.3%
Terminal Rates (c)	2/17 (12%)	5/22 (23%)	4/16 (25%)	3/19 (16%)
Day of First Observation	446	501	319	297
Life Table Tests (d)	P=0.246	P=0.475	P=0.332	P = 0.305
Logistic Regression Tests (d)	P=0.226	P = 0.347	P=0.377	P = 0.212
Cochran-Armitage Trend Test (d)	P = 0.241	1 -0.041	1 -0.017	1 - 0.212
Fisher Exact Test (d)	1 -0.241	P = 0.301	P = 0.395	P = 0.210
Liver: Hepatocellular Carcinoma				
Overall Rates (f)	13/60 (22%)	8/60 (13%)	9/60 (15%)	8/59 (14%)
Adjusted Rates (b)	44.5%	27.1%	32,7%	27.1%
Terminal Rates (c)	4/17 (24%)	4/22 (18%)	2/16 (13%)	2/19 (11%)
Day of First Observation	436	526	425	366
Life Table Tests (d)	P = 0.263N	P = 0.088N	P = 0.315N	P = 0.138N
Logistic Regression Tests (d)	P=0.261N	P = 0.119N	P = 0.293N	P = 0.165N
Cochran-Armitage Trend Test (d)	P = 0.242N	0.11011	1 - 0.20011	1 - 0.10011
Fisher Exact Test (d)	1 - 0.24211	P = 0.168N	P = 0.240 N	P = 0.179N
Liver: Hepatocellular Adenoma or Car	cinoma			
Overall Rates (f)	19/60 (32%)	17/60 (28%)	18/60 (30%)	19/59 (32%)
Adjusted Rates (b)	57.9%	51.0%	57.9%	53.2%
Terminal Rates (c)	6/17 (35%)	8/22 (36%)	6/16 (38%)	5/19 (26%)
Day of First Observation	436	501	319	297
Life Table Tests (d)	P = 0.432	P = 0.210N	P = 0.537	P = 0.457N
Logistic Regression Tests (d)	P = 0.400	P=0.326N	P = 0.573N	P = 0.566
Cochran-Armitage Trend Test (d)	P=0.438			
Fisher Exact Test (d)		$P = 0.421 \mathrm{N}$	P = 0.500N	P = 0.553
Lung: Alveolar/Bronchiolar Adenoma				
Overall Rates (f)	8/60 (13%)	1/60 (2%)	2/60 (3%)	8/60 (13%)
Adjusted Rates (b)	32.9%	4.5%	8.4%	26.2%
Terminal Rates (c)	3/17 (18%)	1/22 (5%)	1/16 (6%)	3/19 (16%)
Day of First Observation	613	729	4 50	366
Life Table Tests (d)	P = 0.228	P = 0.010N	P = 0.068N	P = 0.516N
Logistic Regression Tests (d)	P = 0.210	P = 0.010N	P = 0.062N	P = 0.591 N
Cochran-Armitage Trend Test (d) Fisher Exact Test (d)	P = 0.224	P=0.016N	P = 0.047N	P=0.605N

TABLE C3. ANALYSIS OF PRIMARY TUMORS IN MALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

	Chamber Control	120 ppm	600 ppm	1,200 ppm
Lung: Alveolar/Bronchiolar Adenoma	or Carcinoma			
Overall Rates (f)	9/60 (15%)	1/60 (2%)	2/60 (3%)	9/60 (15%)
Adjusted Rates (b)	36.4%	4.5%	8.4%	28.4%
Terminal Rates (c)	3/17 (18%)	1/22 (5%)	1/16 (6%)	3/19 (16%)
Day of First Observation	613	729	450	366
Life Table Tests (d)	P = 0.205	P = 0.005N	P = 0.042N	P = 0.508N
Logistic Regression Tests (d)	P = 0.187	P = 0.005N	P = 0.037N	P = 0.584N
Cochran-Armitage Trend Test (d)	P = 0.200			
Fisher Exact Test (d)		P = 0.008N	P = 0.027N	P = 0.601 N
Circulatory System: Hemangiosarcon	na			
Overall Rates (a)	2/60 (3%)	0/60 (0%)	1/60 (2%)	5/60 (8%)
Adjusted Rates (b)	8.2%	0.0%	6.3%	23.6%
Terminal Rates (c)	1/17 (6%)	0/22 (0%)	1/16 (6%)	4/19 (21%)
Day of First Observation	543		729	638
Life Table Tests (d)	P = 0.030	P = 0.199N	P = 0.530N	P = 0.264
Logistic Regression Tests (d)	P = 0.032	P = 0.223N	P = 0.531N	P = 0.250
Cochran-Armitage Trend Test (d)	P = 0.031			
Fisher Exact Test (d)		P = 0.248N	P = 0.500N	P = 0.219
Circulatory System: Hemangioma or	Hemangiosarcoma			
Overall Rates (a)	2/60 (3%)	1/60 (2%)	2/60 (3%)	5/60 (8%)
Adjusted Rates (b)	8.2%	2.8%	10.2%	23.6%
Terminal Rates (c)	1/17 (6%)	0/22 (0%)	1/16 (6%)	4/19 (21%)
Day of First Observation	543	607	639	638
Life Table Tests (d)	P = 0.066	P = 0.440N	P = 0.670	P = 0.264
Logistic Regression Tests (d)	P = 0.064	P = 0.484N	P = 0.664	P = 0.250
Cochran-Armitage Trend Test (d)	P = 0.065			
Fisher Exact Test (d)		P=0.500N	P = 0.691 N	P = 0.219
Hematopoietic System: Lymphoma, A	ll Malignant			
Overall Rates (a)	5/60 (8%)	10/60 (17%)	8/60 (13%)	4/60 (7%)
Adjusted Rates (b)	27.2%	36.4%	41.2%	21.1%
Terminal Rates (c)	4/17 (24%)	6/22 (27%)	6/16 (38%)	4/19 (21%)
Day of First Observation	701	556	319	729
Life Table Tests (d)	P = 0.218N	P = 0.262	P = 0.230	P = 0.425N
Logistic Regression Tests (d)	P = 0.198N	P = 0.205	P = 0.210	P = 0.393N
Cochran-Armitage Trend Test (d)	P = 0.211N			
Fisher Exact Test (d)		P = 0.135	P = 0.279	P = 0.500N

⁽a) Number of tumor-bearing animals/number of animals examined grossly at the site

⁽b) Kaplan-Meier estimated tumor incidences at the end of the study after adjusting for intercurrent mortality

⁽c) Observed tumor incidence in animals killed at the end of the study

⁽d) Beneath the control incidence are the P values associated with the trend test. Beneath the dosed group incidence are the P values corresponding to pairwise comparisons between that dosed group and the controls. The life table analysis regards tumors in animals dying prior to terminal kill as being (directly or indirectly) the cause of death. The logistic regression test regards these lesions as nonfatal. The Cochran-Armitage and Fisher exact tests compare directly the overall incidence rates. A negative trend or lower incidence in a dosed group than in controls is indicated by (N).

⁽e) No P value is reported because no tumors were observed in the 600-ppm and control groups.

⁽f) Number of tumor-bearing animals/number of animals examined microscopically at the site

TABLE C4. HISTORICAL INCIDENCE OF ALVEOLAR/BRONCHIOLAR TUMORS IN MALE B6C3F $_1$ MICE RECEIVING NO TREATMENT (a)

		Incidence in Co	ontrols
Study	Adenoma	Carcinoma	Adenoma or Carcinoma
Historical Incidence for C	hamber Controls at Battelle	Pacific Northwest La	boratories
Propylene oxide	14/50	2/50	15/50
Methyl methacrylate	10/50	3/50	11/50
Propylene	7/50	9/50	16/50
1,2-Epoxybutane	7/49	5/49	11/49
Dichloromethane	3/50	2/50	5/50
Ethylene oxide	5/50	6/50	11/50
Bromoethane	5/50	2/50	7/50
Tetrachloroethylene	3/49	4/49	6/49
TOTAL	54/398 (13.6%)	33/398 (8.3%)	82/398 (20.6%)
SD(b)	7.45%	4.96%	8.03%
Range (c)			
High	14/50	9/50	16/50
Low	3/50	2/50	5/50
Overall Historical Incident	ce for Untreated Controls in	n NTP Studies	
TOTAL	204/1,684 (12.1%)	80/1,684 (4.8%)	277/1,684 (16.4%)
SD(b)	6.18%	2.70%	6.91%
Range (c)			
High	14/50	5/49	17/50
Low	1/50	0/49	4/50

⁽a) Data as of May 12, 1988 for studies of at least 104 weeks (b) Standard deviation $\,$

⁽c) Range and SD are presented for groups of 35 or more animals.

TABLE C5. SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN MALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE

	Chamber	Control	120 _I	ppm	600 ₁	ppm	1,200 p	pm
Animals initially in study	60		60		60		60	
Animals minary in study Animals removed	60		60		60		60	
Animals examined histopathologically	60		60		60		60	
ALIMENTARY SYSTEM				· · · · · · · · · · · · · · · · · · ·				
Esophagus	(59)		(60)		(60)		(57)	
Inflammation, chronic			1	(2%)				
Gallbladder	(59)		(54)		(57)		(59)	
Cyst		(2%)						
Infiltration cellular, lymphocytic	-	(10%)		(15%)	_	(14%)	7	(12%)
Inflammation, acute	_	(2%)	1	(2%)	2	(4%)		
Inflammation, chronic	1	(2%)						
Inflammation, chronic active	_			(2%)				
Intestine large	(60)		(60)		(60)		(60)	
Anorectal junction, erosion			1	(2%)	1	(2%)		
Anorectal junction, inflammation, acute					_			(2%)
Anus, erosion						(3%)	1	(2%)
Anus, inflammation, acute						(2%)		
Anus, inflammation, chronic active				(2%)	1			
Anus, ulcer				(3%)	_	(3%)		(5%)
Intestine large, cecum	(60)		(60)		(60)		(59)	
Parasite metazoan					_	(3%)	_	(5%)
Intestine large, colon	(60)		(60)		(60)		(59)	
Parasite metazoan		(2%)		(2%)		(2%)	_	(3%)
Intestine large, rectum	(59)		(59)		(60)		(58)	
Hemorrhage			1	(2%)				
Inflammation, acute						(2%)		
Inflammation, chronic					_	(2%)		
Ulcer	4	(7%)	2	(3%)	1	(2%)		
Anorectal junction, ulcer								(2%)
Intestine small, ileum	(60)		(60)		(60)		(60)	
Amyloid deposition		(2%)		(2%)		(2%)		(2%)
Intestine small, jejunum	(60)		(60)		(60)		(60)	
Hyperplasia, lymphoid		(2%)	(00)					
Liver	(60)		(60)		(60)		(59)	. = ~ .
Basophilic focus	_	(OM)	Z	(3%)			3	(5%)
Bile stasis	1	(2%)				(90)		(90)
Cyst	4	(9%)			1	(2%)	1	(2%)
Focal cellular change Hematopoietic cell proliferation	_	(2%) (35%)	10	(22%)	177	(28%)	177	(29%)
Hyperplasia	21	(30%)		(22%)	17	(4070)		(29%)
Hypertrophy			1	(470)			_	(3%)
Infarct	1	(2%)	1	(2%)	1	(2%)		(3%)
Infiltration cellular, lymphocytic		(3%)		(12%)		(276)		(5%)
Inflammation, acute		(5%)		(2%)		(3%)		(3%)
Inflammation, chronic		(5%)		(2%)		(2%)	4	(0 10)
Necrosis		(8%)		(10%)		(7%)	Q	(15%)
Bile duct, hyperplasia	3	(0 70)	J	(10 %)	•	(1 ~)		(2%)
Caudate lobe, infarct			1	(2%)			1	(2 10)
Centrilobular, fatty change	1	(2%)		(270)				
Periportal, inflammation, chronic		(3%)						
Serosa, inflammation, chronic	2	(0 /0)					1	(2%)
Mesentery	(2)				(2)		(1)	(270)
Artery, inflammation, chronic	(2)					(50%)	(1)	
Artery, inflammation, chronic active	1	(50%)			1	(3070)		
At wry, immanimation, enfonce active	1	(00%)						

TABLE C5. SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN MALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

Cha	mber (Control	120 g	opm	600 p	opm	1,200 p	pm
ALIMENTARY SYSTEM (Continued)								
Pancreas	(60)		(60)		(60)		(59)	
Cyst							1	(2%)
Fibrosis					1	(2%)		
Infiltration cellular, lymphocytic	11	(18%)	14	(23%)	18	(30%)	14	(24%)
Inflammation, acute			1	(2%)				
Inflammation, chronic					1	(2%)		
Inflammation, chronic active			1	(2%)				
Acinus, atrophy	1	(2%)		(5%)	1	(2%)		
Acinus, hyperplasia	1	(2%)	_		1	(2%)		
Salivary glands	(60)	(=,	(60)		(60)		(59)	
Hemorrhage		(2%)	,,,,,		(,			
Hyperplasia, glandular		(2%)						
Infiltration cellular	•	.= .~/	1	(2%)				
Infiltration cellular, lymphocytic	30	(50%)		(60%)	34	(57%)	30	(51%)
Mineralization	00	(00,0)		.00,00	Ų.	,	-	(2%)
Stomach, forestomach	(60)		(59)		(60)		(60)	,
Erosion	(00)		(00)		,00)			(2%)
Hyperkeratosis	1	(2%)			1	(2%)	•	. = 101
Hyperkerawsis Hyperplasia, squamous	1	(270)	7	(12%)		(2%)		
Hyperplasia, squamous Hyperplasia, squamous, focal			,	(1470)	1	(2 /0)	1	(2%)
	1	(2%)	•	(EQ.)	•	(2%)		(2%)
Infiltration cellular, lymphocytic				(5%)	1	(470)		(3%)
Inflammation, acute	4	(7%)	Z	(3%)	1	(90%)	Z	(370)
Inflammation, chronic	•	(0~)				(2%)		
Inflammation, chronic active		(3%)		(O~)		(2%)		
Ulcer		(2%)		(2%)		(2%)	(00)	
Stomach, glandular	(60)		(60)		(60)		(60)	
Edema								(3%)
Erosion		(7%)		(10%)		(10%)		(7%)
Infiltration cellular, lymphocytic		(23%)		(23%)	17	(28%)		(25%)
Inflammation, acute		(3%)		(3%)			3	(5%)
Inflammation, chronic active		(2%)		(2%)				
Mucosa, dilatation	2	(3%)	6	(10%)	5	(8%)	13	(22%)
Mucosa, hyperplasia	1	(2%)						
Submucosa, edema					1	(2%)		
Tooth	(2)		(8)		(2)		(1)	
Peridontal tissue, inflammation, chronic active			2	(25%)				
Pulp, inflammation, acute	2	(100%)	2	(25%)	2	(100%)		
Pulp, inflammation, chronic		, =	1	(13%)				
Pulp, inflammation, chronic active			3	(38%)				
ARDIOVASCULAR SYSTEM								
Blood vessel	(59)		(60)		(60)		(60)	
Aorta, inflammation, chronic	,,		(/		/			(2%)
Aorta, mineralization	1	(2%)						(2%)
Heart	(60)		(60)		(60)		(59)	
Atrophy	(50)		,30/			(2%)	,	
Infiltration cellular, lymphocytic	9	(15%)	13	(22%)		(15%)	8	(14%)
Inflammation, acute	,	.10,0/		(2%)		(2%)		(3%)
Inflammation, chronic	5	(8%)		(7%)	_	(2%)		(3%)
Inflammation, chronic active	J	10 /0/	-	(1,4)		(2%)		(3%)
Mineralization						(2%)	2	.0 101
Atrium, thrombus						(2 10 1	1	(2%)
Ventricle right, thrombus					1	(2%)	•	(2/0)
NDOCRINE SYSTEM								
	(60)		(60)		(59)		(60)	
CNDOCRINE SYSTEM Adrenal gland Capsule, hyperplasia	(60)		(60)		(59) 1	(2%)	(60)	

TABLE C5. SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN MALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

	Chamber Control		120 ppm		600 ppm		1, 200 p	pm
ENDOCRINE SYSTEM (Continued)	 							
Adrenal gland, cortex	(60)		(60)		(59)		(59)	
Hyperplasia	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,			(5%)		(3%)	1=2,	
Hypertrophy	4	(7%)	3	(5%)	3	(5%)	3	(5%)
Pigmentation		(13%)		(12%)	-	(17%)	16	(27%)
Adrenal gland, medulla	(60)		(58)	\·+ <i>,</i>	(58)		(59)	, = ,,
Hyperplasia		(2%)		(2%)	(55)		(007	
Islets, pancreatic	(60)		(60)	(= ,-,	(60)		(58)	
Hyperplasia	(00)			(2%)		(2%)	(00)	
Necrosis			•	(= ///		(2%)		
Pituitary gland	(59)		(58)		(58)		(56)	
Pars distalis, cyst		(2%)		(2%)	(00)		(00)	
Pars distalis, hyperplasia		(3%)		(2%)				
Pars intermedia, hyperplasia	-	(0,0)	•	(2 /0)			1	(2%)
Thyroid gland	(60)		(60)		(60)		(59)	(2 /0)
Infiltration cellular, lymphocytic		(13%)		(8%)		(3%)		(5%)
Inflammation, acute	0	(1070)	3	(0/0)	2	(0 /0)		(2%)
C-cell, hyperplasia			1	(2%)			1	(2 10)
Follicle, cyst	9	(3%)		(2 10)	1	(2%)		
Follicle, dilatation		(5%)	9	(3%)	1	(24/01	2	(5%)
Follicle, hyperplasia		(7%)		(8%)	9	(3%)		(5%)
GENITAL SYSTEM Epididymis	(60)		(60)		(60)		(60)	
Granuloma sperm		(5%)	(00)		(007			(2%)
Infiltration cellular, lymphocytic		(15%)	3	(5%)	R	(13%)		(7%)
Inflammation, acute	J	(10 %)		(2%)		(3%)		(5%)
Inflammation, chronic			•	(2 10)			-	(2%)
Inflammation, chronic active			1	(2%)		(2%)		(7%)
Penis	(17)		(21)	(270)	(17)	(270)	(21)	(170)
Abscess	(11)			(14%)		(6%)	121/	
Concretion	1	(6%)	J	(1470)	•	(0%)		
Inflammation		(6%)	1	(5%)			1	15941
Inflammation, acute		(94%)			10	(76%)		(5%) (57%)
	10	(3470)		(48%)	13	(1070)		
Inflammation, chronic Inflammation, chronic active				(10%)				(5%)
Necrosis				(10%)	4	(0.40)		(5%)
Preputial gland	(10)			(10%)		(24%)	(11)	(19%)
	(12)		(9)	(00% \	(10)	(202)	,	(978)
Abscess		(33%)		(22%) (22%)		(30%)		(27%)
Cyst	3	(25%)				(20%)	Z	(18%)
Infiltration cellular, lymphocytic Inflammation, acute	0	(17%)		(22%)	1	(10%)	٥	(100)
				(11%)	0	(900)	Z	(18%)
Inflammation, chronic		(25%)		(11%)		(20%)	o	(1990)
Inflammation, chronic active Prostate		(8%)		(11%)		(30%)		(18%)
Infiltration cellular, lymphocytic	(60)		(60)	(10%)	(59)	(27%)	(59)	/9.40%×
Infiltration cellular, lymphocytic Inflammation, acute		(13%)		(10%)		(27%)		(24%)
		(23%) (2%)		(22%)	9	(15%)	12	(20%)
Inflammation, chronic		(3%)	2	(3%)	•	(E0)		(00)
Inflammation, chronic active		(3%)	.4.			(5%)		(2%)
Seminal vesicle	(4)		(4)	(DE~ \	(4)	(E0%\	(6)	(150
Dilatation	4	(980°)		(25%)	2	(50%)	1	(17%)
Inflammation, chronic	1	(25%)	1	(25%)			4	(17%)
THE STREET STREET								. i (970)

1 (17%)

Inflammation, chronic active

TABLE C5. SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN MALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

•	Chamber (Control	ontrol 120 ppm		600 j	ppm	1,200 ppm		
GENITAL SYSTEM (Continued)									
Testes	(60)		(60)		(60)		(60)		
Atrophy		(15%)		(20%)		(5%)	,	(5%)	
Granuloma sperm		(3%)		(=0 ,0)	·	(0,0)	·	(0,0)	
Hemorrhage		(2%)							
Infiltration cellular, lymphocytic		(2%)							
Inflammation, acute	-	(= ,0)			1	(2%)	2	(3%)	
Mineralization			1	(2%)	-	(= ,0 ,	_	(0,0)	
Interstitial cell, hyperplasia	1	(2%)		(2%)	2	(3%)			
Tunic, inflammation, chronic active	_	(= , ,	_	(= ///		(2%)			
HEMATOPOIETIC SYSTEM									
Bone marrow	(60)		(60)		(60)		(59)		
Fibrosis	(00)		(00)			(2%)		(2%)	
Inflammation, acute			1	(2%)		(270)	1	(2 70)	
Myeloid cell, hyperplasia	51	(85%)		(80%)	56	(93%)	4.4	(75%)	
Lymph node	(60)		(60)	(0070)	(59)	(0070)	(59)	(1070)	
Congestion		(3%)	(00)			(2%)	(88)		
Hyperplasia, lymphoid		(3%) (7%)	۰	(5%)	_	(3%)	1	(2%)	
Inflammation, acute	4	(170)		(2%)	_	(2%)		(2%)	
Iliac, hyperplasia, lymphoid	10	(170-)		(2%) (7%)		(8%)		(7%)	
Iliac, hyperplasia, reticulum cell	10	(17%)		(1%) (2%)	9	(070)	4	(170)	
Iliac, inflammation, acute				(2%)	•	(2%)			
Inguinal, hyperplasia, lymphoid	9	(3%)	1	(2%)		(5%)	•	(2%)	
Lumbar, hyperplasia, lymphoid	2	(370)			_	(3%)	1	(270)	
Mediastinal, hyperplasia, lymphoid	4	(7%)		(2%)		(10%)		(8%)	
Mesenteric, autolysis	4	(170)	1	(270)	•	(10%)		(2%)	
Mesenteric, autolysis Mesenteric, congestion	90	(220)	1.4	(000)	10	(20%)	_		
	20	(33%)	14	(23%)	12	(20%)		(19%)	
Mesenteric, hematopoietic cell proliferation		(94)	0	(0 or)			1	(2%)	
Mesenteric, hemorrhage Mesenteric, hyperplasia, lymphoid		(2%)		(3%)	0.5	(E0@\	07	(400)	
Mesenteric, hyperplasia, lympholo Mesenteric, inflammation, acute		(53%)		(40%) (8%)		(59%) (5%)		(46%)	
Renal, congestion		(7%) (2%)	э	(8%)	J	(070)	10	(17%)	
Renal, hyperplasia, lymphoid		(3%)			0	(3%)		(3%)	
Lymph node, mandibular	(53)	(3%)	(52)		(54)	(370)	(51)	(370)	
Autolysis	(53)		(32)		(54)		,	(90%)	
Congestion					,	(2%)	1	(2%)	
Hyperplasia, lymphoid	20	(72%)	40	(77%)		(2%) (7 0%)	90	(63%)	
Inflammation, acute			40	(1170)	30	(1070)	32	(0070)	
Necrosis		(2%) (2%)							
Pigmentation	1	(470)			,	(90%)			
Spleen	(60)		(60)			(2%)	(59)		
Angiectasis		(20%)	(00)		(60)	(200)	(59)		
Congestion	1	(2%)	•	(2%)	2	(3%)			
Hematopoietic cell proliferation	E 4	(0.00)			EO	(020)	477	(80%)	
Hyperplasia, lymphoid		(90%) (17%)		(83%)		(93%)		(80%)	
Hyperplasia, lymphold Hyperplasia, reticulum cell	10	(1/70)		(12%)	0	(10%)	ð	(8%)	
Necrosis			1	(2%)			0	(20%)	
Pigmentation	4	(70.)	0	(150)	1.1	(1900)		(3%)	
Capsule, inflammation, chronic active	4	(7%)	9	(15%)		(18%)	18	(31%)	
	/FO		/E01			(2%)	1845		
Thymus	(53)		(59)	(90)	(58)		(54)		
Congestion	^	(Car)	1	(2%)	^	(00)	_	(OC)	
Cyst Fotonia namathymaid gland		(6%)			2	(3%)	1	(2%)	
Ectopic parathyroid gland		(2%)		(0.00)	,	(TO)	_	(46)	
Hyperplasia, lymphoid	2	(4%)		(3%)	4	(7%)	2	(4%)	
Inflammation, chronic active			1	(2%)					

TABLE C5. SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN MALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

Cha	Chamber Control		120 ppm		600 ppm		1,200 ppm	
NTEGUMENTARY SYSTEM								
Skin	(60)		(60)		(60)		(58)	
Alopecia	4	(7%)	5	(8%)	3	(5%)	5	(9%)
Inflammation, acute	1	(2%)	1	(2%)				
Inflammation, chronic		(3%)		(2%)		(2%)		
Ulcer	3	(5%)		(2%)	2	(3%)	4	(7%)
Abdominal, abscess				(2%)				
Abdominal, thoracic, alopecia				(2%)				
Foot, ulcer				(2%)	2	(3%)		(2%)
Head, abscess	_			(2%)			1	(2%)
Head, inflammation, acute		(3%)	1	(2%)				
Head, inflammation, chronic	3	(5%)			1	(2%)		
Head, ulcer							1	(2%)
Inguinal, abscess	1	(2%)						
Inguinal, inflammation, acute		(B.4)	_	.=		(2%)		
Inguinal, ulcer	1	(2%)	4	(7%)		(3%)		
Neck, abscess					1	(2%)	_	
Neck, alopecia	_		_				1	(2%)
Prepuce, abscess	2	(3%)	2	(3%)				/A-4 \
Prepuce, inflammation, acute								(2%)
Prepuce, inflammation, chronic active	_			(2%)	_		-	(2%)
Prepuce, ulcer		(15%)	14	(23%)		(13%)		(17%)
Scrotal, abscess	5	(8%)			1	(2%)	1	(2%)
Scrotal, inflammation, acute			1	(2%)				
Scrotal, inflammation, chronic active		(05%)	_	/4 = ~ \		(2%)		(1.4~)
Scrotal, ulcer Subcutaneous tissue, edema	16	(27%)		(15%)	16	(27%)	-	(14%)
		(0~)		(2%)			1	(2%)
Subcutaneous tissue, inflammation, acute	2		1	(2%)				
Subcutaneous tissue, inflammation, chronic		(2%)	•	(00)	,	(OM)	•	(0 <i>a</i> r.)
Subcutaneous tissue, head, abscess Subcutaneous tissue, head, granuloma	Z	(3%)		(2%) (2%)	1	(2%)	1	(2%)
Subcutaneous tissue, head, inflammation, acute	•	(2%)	1	(270)				
Subcutaneous tissue, head, inflammation,		(270)						
chronic active			1	(2%)				
Tail, inflammation, acute	1	(2%)	1	(270)				
Tail, inflammation, chronic		(270)			1	(2%)		
Tail, necrosis	1	(2%)			•	(2 10)		
Tail, ulcer	•	(270)	1	(2%)	1	(2%)		
Thoracic, alopecia	1	(2%)		(270)	1	(210)		
Ventral, alopecia		(3%)	2	(3%)				
USCULOSKELETAL SYSTEM								
Bone	(60)		(60)		(60)		(60)	
Cranium, inflammation, chronic active	/			(2%)				(2%)
Femur, hyperostosis	1	(2%)	-					(2%)
Tibia, fracture	•	,						(2%)
Skeletal muscle			(2)		(6)		(3)	
Abscess						(17%)		
Inflammation, acute							1	(33%)
Inflammation, chronic								(33%)
Inflammation, chronic active								(33%)
Head, abscess					4	(67%)	_	
Head, inflammation, acute			1	(50%)				
Head, inflammation, chronic active			_		1	(17%)		

TABLE C5. SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN MALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

(Chamber Control		120 ppm		600 ppm		1, 20 0 p	pm
NERVOUS SYSTEM								
Brain	(60)		(60)		(60)		(60)	
Abscess	, 507		,		,			(2%)
Compression					1	(2%)	-	
Hemorrhage	2	(3%)	3	(5%)	2		1	(2%)
Infiltration cellular, lymphocytic		(2%)	·	(0,0)	_	(0,0)	-	(= / • /
Mineralization		(67%)	35	(58%)	41	(68%)	35	(58%)
RESPIRATORY SYSTEM					,,,,			
Lung	(60)		(60)		(60)		(60)	
Abscess			,					(2%)
Congestion	11	(18%)	12	(20%)	5	(8%)		(13%)
Granuloma		(20.0)		(20,0)		(3%)	•	(40.00
Hemorrhage	6	(10%)	11	(18%)		(27%)	3	(5%)
Infiltration cellular, lymphocytic		(83%)		(90%)		(92%)	-	(87%)
Inflammation, acute	50	,00,00,	~~	(00,00		(2%)	~2	, , , , , ,
Mineralization	9	(3%)			•	(- . v /		
Alveolar epithelium, hyperplasia		(2%)			2	(3%)	1	(2%)
Alveolus, infiltration cellular, histiocytic		(5%)	•	(2%)	_	(3%)		(5%)
Arteriole, inflammation, acute	3	(570)	1	(270)		(2%)	3	1070)
Artery, inflammation, acute	-	(00)		(00)	_	(2%)		(70)
Interstitium, inflammation, acute		(8%)	1	(2%)	3	(5%)	4	(7%)
Interstitium, inflammation, chronic		(2%)						
Peribronchiolar, inflammation, acute		(2%)	_					
Peribronchiolar, inflammation, chronic		(3%)	1	(2%)				
Peribronchiolar, inflammation, chronic activ	e						1	(2%)
Perivascular, inflammation, acute				(2%)				
Nose	(59)		(59)		(60)		(59)	
Lumen, hemorrhage	36	(61%)	25	(42%)		(67%)	29	(49%)
Mucosa, inflammation, acute	1	(2%)	3	(5%)	1	(2%)	6	(10%)
Nasolacrimal duct, hemorrhage	2	(3%)	1	(2%)				
Nasolacrimal duct, inflammation, acute		(2%)		(7%)	5	(8%)		
Olfactory epithelium, degeneration		(69%)		(19%)	_	(45%)	30	(51%)
Olfactory epithelium, metaplasia	-21	,00,0,	**	,		/ - /		(2%)
Respiratory epithelium, degeneration	7	(12%)	1	(2%)	Ω	(13%)		(7%)
Respiratory epithelium, inflammation, acute		(1270)		(2%)	0	(1070)	4	(170)
Septum, inflammation, acute	;		ī	(470)		(2%)		
				(90')	1	(270)		
Septum, inflammation, chronic active			1	(2%)				(0¢.
Sinus, inflammation, acute					_	′0 ~ `		(2%)
Turbinate, congestion					1	(2%)	2	(3%)
Turbinate, inflammation, acute	1	(2%)						
Vomeronasal organ, congestion				(2%)				
Vomeronasal organ, inflammation, acute				(2%)		(3%)		
Trachea	(60)		(60)		(60)		(59)	
Inflammation, chronic active					1	(2%)		
Glands, inflammation, acute			3	(5%)				
Mucosa, erosion	1	(2%)	,	/				
PECIAL SENSES SYSTEM						· · · · · · · · · · · · · · · · · · ·		
Harderian gland			(3)		(1)		(3)	
Cyst			,,,		,-,			(33%)
URINARY SYSTEM				,				
Kidney	(60)		(60)		(60)		(59)	
Abscess	(00)			(2%)		(2%)		(2%)
Congestion	1	(2%)	1	(470)	•	(a / v /	1	~ 101
Cyst		(2%)						
Hemorrhage	1	(270)			1	(2%)		
Hydronephrosis				(90%)		(2 /0)		
113 at otte htt rosts			1	(2%)				

TABLE C5. SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN MALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

	Chamber (Control	120 ppm		600 ppm		1,200 ppm	
TRINARY SYSTEM								
Kidney (Continued)	(60)		(60)		(60)		(59)	
Infiltration cellular, lymphocytic		(88%)	\ + - /	(87%)		(87%)	48	(81%)
Inflammation, acute		(7%)		(5%)		(5%)		(5%)
Inflammation, chronic	ī	(2%)	_	(+ /	-	, = ,		,
Inflammation, chronic active			1	(2%)	1	(2%)	1	(2%)
Metaplasia, osseous					1	(2%)		
Capsule, inflammation, chronic							1	(2%)
Cortex, cyst	2	(3%)	2	(3%)			2	(3%)
Pelvis, calculus micro observation only							1	(2%)
Pelvis, dilatation	14	(23%)	10	(17%)	9	(15%)	4	(7%)
Pelvis, hemorrhage			1	(2%)			1	(2%)
Pelvis, inflammation, acute	7	(12%)	3	(5%)	5	(8%)	9	(15%)
Pelvis, inflammation, chronic	1	(2%)						
Pelvis, inflammation, chronic active	1	(2%)					1	(2%)
Renal tubule, casts protein	4	(7%)	6	(10%)	4	(7%)	2	(3%)
Renal tubule, cyst	2	(3%)				(2%)		
Renal tubule, dilatation	13	(22%)	17	(28%)	15	(25%)	17	(29%)
Renal tubule, hyperplasia	1	(2%)	3	(5%)	1	(2%)		
Renal tubule, mineralization	2	(3%)	6	(10%)	1	(2%)		
Renal tubule, necrosis	6	(10%)	12	(20%)	9	(15%)	12	(20%)
Renal tubule, regeneration	36	(60%)		(50%)	29	(48%)	21	(36%)
Ureter	(8)		(1)		(3)			
Dilatation	3	(38%)	1	(100%)	3	(100%)		
Inflammation, acute	-	(13%)	_	, / - /	_	(,		
Urethra	(5)	,	(6)		(1)		(3)	
Calculus micro observation only		(60%)	,	(67%)	/			(33%)
Inflammation, acute	-	(20%)	-	(17%)	1	(100%)	_	(33%)
Inflammation, chronic		(20%)	•	(11,0)	•	(100,0)	•	(00,0)
Bulbourethral gland, inflammation, acute		(20,0)	1	(17%)				
Urinary bladder	(60)		(60)	(1170)	(60)		(59)	
Angiectasis	,	(3%)	(,	(2%)	(00)		(00)	
Calculus gross observation	_	(2%)		(2%)			2	(3%)
Calculus micro observation only	î		•	(2,0)			_	(3%)
Congestion	•	(210)	1	(2%)			_	(0,0)
Ectasia	19	(32%)		(17%)	15	(25%)	13	(22%)
Hemorrhage		(2%)	.0	(21/0)	.0	.20,0,	.0	(22,0)
Infiltration cellular, lymphocytic		(45%)	25	(42%)	36	(60%)	31	(53%)
Inflammation, acute		(17%)		(13%)		(12%)		(19%)
Inflammation, chronic		(2%)		(5%)	•	122701	• •	(10 10)
Inflammation, chronic active		(10%)	J	(3 10)		(2%)	4	(7%)

APPENDIX D

SUMMARY OF LESIONS IN FEMALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE

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TABLE D1. SUMMARY OF THE INCIDENCE OF NEOPLASMS IN FEMALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE

Char	mber (Control	120 _l	ppm	600 j	pm	1,200 p	pm
Animals initially in study	60		60		60		60	
Animals removed	60		60		60		60	
Animals examined histopathologically	50		50		50		47	
ALIMENTARY SYSTEM			-		-			
Gallbladder	(49)		(49)		(50)		(45)	
Lymphoma malignant mixed	4	(8%)	3	(6%)	1	(2%)		
Lymphoma malignant undifferentiated cell type					2	(4%)		
Intestine large, cecum	(49)		(50)		(50)		(47)	
Histiocytic sarcoma					1	(2%)		
Lymphoma malignant histiocytic								(2%)
Lymphoma malignant lymphocytic							1	(2%)
Lymphoma malignant mixed	1	(2%)	1	(2%)				
Intestine large, rectum	(50)		(50)		(50)		(45)	
Lymphoma malignant lymphocytic								(2%)
Intestine small, duodenum	(49)		(50)		(50)		(47)	_
Lymphoma malignant lymphocytic							1	(2%)
Lymphoma malignant mixed						(2%)		
Lymphoma malignant undifferentiated cell type						(4%)		
Intestine small, jejunum	(49)		(50)		(50)		(47)	
Lymphoma malignant mixed	3	(6%)	1	(2%)				_
Lymphoma malignant undifferentiated cell type								(2%)
Liver	(49)		(50)		(50)		(47)	
Hemangioma		(2%)						
Hemangiosarcoma		(2%)						
Hepatocellular carcinoma		(8%)	-	(4%)	_	(4%)		(15%)
Hepatocellular adenoma	3	(6%)	7	(14%)		(12%)		(15%)
Histiocytic sarcoma						(2%)		(2%)
Lymphoma malignant histiocytic					1	(2%)		(2%)
Lymphoma malignant lymphocytic		(4%)					2	(4%)
Lymphoma malignant mixed	8	(16%)	6	(12%)		(16%)		
Lymphoma malignant undifferentiated cell type					3	(6%)	1	(2%)
Mesentery	*(50)		*(50)		*(50)		*(47)	
Hepatocellular carcinoma, metastatic, liver							1	(2%)
Lymphoma malignant lymphocytic	1	(2%)						
Lymphoma malignant					1	(2%)		
Lymphoma malignant mixed	3	(6%)	2	(4%)	2	(4%)		
Osteosarcoma, metastatic, uncertain primary								
site							1	(2%)
Pancreas	(50)		(50)		(50)		(47)	
Lymphoma malignant histiocytic								(2%)
Lymphoma malignant lymphocytic							1	(2%)
Lymphoma malignant mixed	6	(12%)	3	(6%)	3	(6%)		
Lymphoma malignant undifferentiated cell type						(2%)		
Salivary glands	(50)	(5.24)	(50)		(50)		(46)	
Lymphoma malignant histiocytic		(2%)						(0.55
Lymphoma malignant lymphocytic		(4%)		/0 <i>~</i> \			1	(2%)
Lymphoma malignant mixed	4	(8%)	4	(8%)				
Lymphoma malignant undifferentiated cell type	/FA:		/=~			(2%)		
Stomach	(50)		(50)		(50)		(47)	
Histiocytic sarcoma		.00			1	(2%)		
Lymphoma malignant lymphocytic		(2%)						
Stomach, forestomach	(50)		(49)		(50)		(47)	.00
Lymphoma malignant lymphocytic		(CA)		/O# \	_	(40)	1	(2%)
Lymphoma malignant mixed Lymphoma malignant undifferentiated cell type	3	(6%)	1	(2%)		(4%)		
					I	(2%)		
Papilloma squamous	0	(6%)		(2%)	1	(2%)	•	(6%)

TABLE D1. SUMMARY OF THE INCIDENCE OF NEOPLASMS IN FEMALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

Cha	namber Control		120 ppm		600 ppm		1,200 ppm	
ALIMENTARY SYSTEM (Continued)								
Stomach, glandular	(49)		(50)		(50)		(47)	
Lymphoma malignant histiocytic							1	(2%)
Lymphoma malignant lymphocytic							1	(2%)
Lymphoma malignant mixed	6	(12%)	2	(4%)	2	(4%)	1	(2%)
Lymphoma malignant undifferentiated cell typ	е				2	(4%)		
CARDIOVASCULAR SYSTEM						****		
Blood vessel	(50)		(50)		(50)		(47)	
Lymphoma malignant lymphocytic	1	(2%)						
Heart	(50)		(50)		(50)		(47)	
Hemangiosarcoma	,,		, ,				1	(2%)
Hemangiosarcoma, metastatic, ovary							1	(2%)
Histiocytic sarcoma					1	(2%)		
Lymphoma malignant lymphocytic	1	(2%)					1	(2%)
Lymphoma malignant mixed	1	(2%)	3	(6%)	1	(2%)		
ENDOCRINE SYSTEM								
Adrenal gland, cortex	(49)		(50)		(50)		(47)	
Adenoma					1	(2%)	1	(2%)
Lymphoma malignant mixed		(2%)						
Lymphoma malignant undifferentiated cell typ					1	(2%)		
Capsule, lymphoma malignant undifferentiated	ł							
cell type					-	(2%)		
Adrenal gland, medulla	(49)		(50)		(49)		(47)	
Pheochromocytoma malignant				(2%)				
Pheochromocytoma, NOS		(2%)		(2%)	_	(2%)		(4%)
Pituitary gland	(49)		(48)		(49)		(46)	
Pars distalis, adenoma	12	(24%)	19	(40%)	21	(43%)	15	(33%)
Pars distalis, lymphoma malignant mixed			1	(2%)				
Pars intermedia, adenoma			1	(2%)	1	(2%)	1	(2%)
Thyroid gland	(50)		(50)		(50)		(47)	
Lymphoma malignant mixed			1	(2%)				
Follicle, adenocarcinoma	1	(2%)						
Follicle, adenoma					4	(8%)		
GENERAL BODY SYSTEM None								
GENITAL SYSTEM								
Ovary	(50)		(49)		(50)		(47)	
Granulosa cell tumor benign			1	(2%)				
Hemangioma	1	(2%)						(2%)
Hemangiosarcoma								(2%)
Hemangiosarcoma, metastatic								(2%)
Histiocytic sarcoma							1	(2%)
Luteoma	1	(2%)						
Lymphoma malignant histiocytic								(2%)
Lymphoma malignant lymphocytic							1	(2%)
Lymphoma malignant mixed		(8%)	1	(2%)		(4%)		
Lymphoma malignant undifferentiated cell typ						(2%)		
	(EA)		(50)		(50)		(47)	
Uterus	(50)							
Adenocarcinoma	(50)		1	(2%)				
	(50)		1	(2%) $(2%)$				(2%) (2%)

TABLE D1. SUMMARY OF THE INCIDENCE OF NEOPLASMS IN FEMALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

Char		mber Control		120 ppm		600 ppm		pm
GENITAL SYSTEM								
Uterus (Continued)	(50)		(50)		(50)		(47)	
Leiomyoma			1	(2%)				
Lymphoma malignant histiocytic					1	(2%)		
Lymphoma malignant lymphocytic		(2%)					1	(2%)
Lymphoma malignant mixed	2	(4%)	1	(2%)		(2%)		
Lymphoma malignant undifferentiated cell type						(2%)	_	
Endometrium, polyp stromal	3	(6%)	2	(4%)	1	(2%)	2	(4%)
HEMATOPOIETIC SYSTEM								
Blood	* (50)		* (50)		*(50)		*(47)	
Leukemia			1	(2%)				
Bone marrow	(50)		(50)		(50)		(47)	
Hemangioma			1	(2%)				
Hemangiosarcoma, metastatic, ovary							1	(2%)
Hemangiosarcoma, metastatic, skin					1	(2%)		
Hemangiosarcoma, metastatic, spleen								(2%)
Lymphoma malignant lymphocytic	1	(2%)					1	(2%)
Lymphoma malignant mixed	2	(4%)	1	(2%)	1	(2%)		
Lymph node	(48)		(49)		(50)		(46)	
Adenocarcinoma, metastatic, uterus			1	(2%)				
Lymphoma malignant mixed	3	(6%)	2	(4%)	_	(2%)		
Lymphoma malignant undifferentiated cell type					2	(4%)	1	(2%)
Axillary, lymphoma malignant mixed	1	(2%)						
Axillary, lymphoma malignant undifferentiated								
cell type							1	(2%)
Bronchial, lymphoma malignant mixed	1	(2%)						
Iliac, histiocytic sarcoma					1	(2%)		(2%)
Iliac, lymphoma malignant histiocytic								(2%)
Iliac, lymphoma malignant mixed	3	(6%)	3	(6%)	5	(10%)	3	(7%)
Iliac, lymphoma malignant undifferentiated								
cell type					2	(4%)		
Mediastinal, lymphoma malignant histiocytic								(2%)
Mediastinal, lymphoma malignant lymphocytic							_	(2%)
Mediastinal, lymphoma malignant mixed	5	(10%)	2	(4%)	4	(8%)	1	(2%)
Mediastinal, lymphoma malignant								
undifferentiated cell type						(2%)		
Mesenteric, histiocytic sarcoma					2	(4%)		
Mesenteric, lymphoma malignant histiocytic							_	(2%)
Mesenteric, lymphoma malignant lymphocytic		(4%)						(4%)
Mesenteric, lymphoma malignant mixed	14	(29%)	8	(16%)	10	(20%)	6	(13%)
Mesenteric, lymphoma malignant							_	
undifferentiated cell type					2	(4%)	2	(4%)
Mesenteric, osteosarcoma, metastatic, uncertain								
primary site							1	(2%)
Renal, histiocytic sarcoma					2	(4%)		
Renal, lymphoma malignant histiocytic								(2%)
Renal, lymphoma malignant lymphocytic								(2%)
Renal, lymphoma malignant mixed	3	(6%)	3	(6%)	3	(6%)	1	(2%)
Renal, lymphoma malignant undifferentiated								
cell type						(4%)		
Lymph node, mandibular	(47)		(48)		(50)	_	(43)	
Histiocytic sarcoma					1	(2%)		
Lymphoma malignant histiocytic								(2%)
Lymphoma malignant lymphocytic	1	(2%)						(5%)
Lymphoma malignant mixed	11	(23%)	6	(13%)		(10%)		(9%)
Lymphoma malignant undifferentiated cell type					1	(2%)	1	(2%)

TABLE D1. SUMMARY OF THE INCIDENCE OF NEOPLASMS IN FEMALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

Ch	ıamber (amber Control		120 ppm		600 ppm		pm
HEMATOPOIETIC SYSTEM (Continued)								
Spleen	(50)		(50)		(49)		(47)	
Adenocarcinoma, metastatic, uterus	(00)			(2%)				
Hemangiosarcoma	1	(2%)	1	(2%)				(4%)
Hemangiosarcoma, metastatic, uterus								(2%)
Histiocytic sarcoma					2	(4%)		(2%)
Lymphoma malignant histiocytic								(2%)
Lymphoma malignant lymphocytic	1	(2%)						(4%)
Lymphoma malignant mixed		(36%)	9	(18%)		(22%)		(13%)
Lymphoma malignant undifferentiated cell ty	pe				_	(6%)		(4%)
Thymus	(46)		(48)		(48)		(47)	
Lymphoma malignant histiocytic								(2%)
Lymphoma malignant lymphocytic	_	(4%)						(4%)
Lymphoma malignant mixed		(33%)	8	(17%)		(8%)		(2%)
Lymphoma malignant undifferentiated cell ty	pe				1	(2%)	1	(2%)
NTEGUMENTARY SYSTEM								
Mammary gland	(49)		(50)		(48)		(46)	
Adenoacanthoma			1	(2%)				
Adenocarcinoma	2	(4%)			2	(4%)		
Carcinoma								(2%)
Skin	(50)		(50)		(50)		(47)	
Papilloma squamous								(2%)
Subcutaneous tissue, fibrosarcoma					2	(4%)		(2%)
Subcutaneous tissue, hemangioma								(2%)
Subcutaneous tissue, hemangiosarcoma					1	(2%)	1	(2%)
MUSCULOSKELETAL SYSTEM						_		
Skeletal muscle	*(50)		*(50)		*(50)		*(47)	
Head, sarcoma, deep invasion			1	(2%)				
NERVOUS SYSTEM								
Brain	(50)		(50)		(50)		(47)	
Lymphoma malignant lymphocytic	(00)		,,				1	(2%)
Lymphoma malignant mixed	2	(4%)	1	(2%)	2	(4%)		
Choroid plexus, lymphoma malignant mixed	_	, = , ,	1	(2%)				
Meninges, sarcoma, metastatic			1	(2%)				
RESPIRATORY SYSTEM								
Lung	(50)		(50)		(50)		(47)	
Alveolar/bronchiolar adenoma		(10%)	(-0)			(6%)		(9%)
Alveolar/bronchiolar carcinoma	·	(- V /V /	3	(6%)	-	(2%)		(6%)
Carcinoma, metastatic			•	,	-	,		(2%)
Hepatocellular carcinoma, metastatic, liver	1	(2%)						(4%)
Histiocytic sarcoma	•	/-/			1	(2%)		(2%)
Lymphoma malignant histiocytic	1	(2%)						(2%)
Lymphoma malignant lymphocytic		(4%)					2	(4%)
Lymphoma malignant mixed		(20%)	8	(16%)	8	(16%)	1	(2%)
Lymphoma malignant undifferentiated cell ty			-		2	(4%)		
Osteosarcoma, metastatic, uncertain primary					_	•		
							1	(2%)
site		 -						
SPECIAL SENSES SYSTEM	*(50)		*(50)		*(50)		*(47)	
SPECIAL SENSES SYSTEM Harderian gland	*(50)		*(50) 2	(4%)	*(50) 1	(2%)	*(47) 1	(2%)
SPECIAL SENSES SYSTEM	*(50)			(4%)		(2%)	1	(2%) (2%)

TABLE D1. SUMMARY OF THE INCIDENCE OF NEOPLASMS IN FEMALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

Cha	amber (Control	120 p	pm	600 g	pm	1, 200 p	pm
URINARY SYSTEM		-						
Kidney	(50)		(50)		(50)		(47)	
Histiocytic sarcoma					1	(2%)		(2%)
Lymphoma malignant histiocytic			_					(2%)
Lymphoma malignant lymphocytic		(2%)		(2%)	_			(2%)
Lymphoma malignant mixed		(22%)	5	(10%)		(14%)		(6%)
Lymphoma malignant undifferentiated cell typ Osteosarcoma, metastatic, uncertain primary	e				1	(2%)		(2%)
site	(50)		(FA)		(FO)			(2%)
Urinary bladder	(50)		(50)		(50)		(47)	(2%)
Lymphoma malignant histiocytic		(40)						(2%)
Lymphoma malignant lymphocytic		(4%) (20%)		(10%)		(8%)		(2%)
Lymphoma malignant mixed Lymphoma malignant undifferentiated cell typ		(20%)	5	(10%)		(2%)	•	(270)
SYSTEMIC LESIONS								
Multiple organs	*(50)		*(50)		*(50)		*(47)	
Lymphoma malignant mixed	20	(40%)	10	(20%)	13	(26%)	6	(13%)
Hemangiosarcoma	2	(4%)	2	(4%)	1	(2%)	6	(13%)
Lymphoma malignant lymphocytic	2	(4%)	1	(2%)			2	(4%)
Lymphoma malignant histiocytic	2	(4%)			1	(2%)	1	(2%)
Hemangioma	2	(4%)	1	(2%)			2	(4%)
Leukemia			1	(2%)				
Lymphoma malignant					1	(2%)		
Lymphoma malignant undifferentiated cell					3	(6%)	2	(4%)
ANIMAL DISPOSITION SUMMARY				· · · · · · · · · · · · · · · · · · ·				
Animals initially in study	60		60		60		60	
Interval sacrifice	10		10		10		10	
Terminal sacrifice	30		33		23		32	
Dead	11		6		14		9	
Moribund	8		8		11		3	
Accident	1		3				1	
Natural death					2		2	
Missing							3	
FUMOR SUMMARY								
Total animals with primary neoplasms **	33		36		42		38	
Total primary neoplasms	63		59		81		76	
Total animals with benign neoplasms	19		28		29		29	
Total benign neoplasms	29		35		39		37	
Total animals with malignant neoplasms	26		17		27		26	
Total malignant neoplasms	33		23		41		37	
Total animals with secondary neoplasms ***	1		2		1		8	
Total secondary neoplasms	1		3		1		13	
Total animals with malignant neoplasms-							•	
uncertain primary site							1	
Total animals with neoplasms-	,		1		1		n	
uncertain benign or malignant	1		1		1		$\frac{2}{2}$	
Total uncertain neoplasms	Ţ		1		1		Z	

^{*} Number of animals receiving complete necropsy examinations; all gross lesions including masses examined microscopically.

** Primary tumors: all tumors except secondary tumors

*** Secondary tumors: metastatic tumors or tumors invasive into an adjacent organ

TABLE D2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF FEMALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE: CHAMBER CONTROL

WEEKS ON STUDY	0 0 1	0 5 0	0 5 4	0 5 7	0 6 7	0 8 0	0 9 0	0 9 1	9 1	0 9 1	0 9 4	0 9 8	9 8	0 9 9	1 0 2	1 0 2	1 0 2	1 0 3	1 0 3	1 0 4	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5
CARCASS ID	1 1 3 1	1 1 2 1	0 8 3 1	1 1 8 1	1 0 9 1	0 9 8 1	0 6 3	0 8 9	7 7 1	0 9 0	0 9 6	1 1 1 1	1 0 5 1	0 9 9	0 8 8	0 8 4 1	1 0 3 1	0 6 4 1	0 7 0 1	0 7 4	0 6 2 1	0 7 2 1	0 8 1	0 8 5 1	0 9 1
ALIMENTARY SYSTEM	_																								
Esophagus Gallbladder	+	+	+	+	+	+	+	+	+	+	+	+	+	M	+	+	+	+	+	+	+	+	+	+	+
Gallbladder	+	+	+	+	+	+	+	+	M	+	+	+	+	+	+	+	+	+	+ X	+	+	+	+	+	+
Lymphoma malignant mixed Intestine large	+	+	+	+	+	+	+	+	+	+	+	+	X	+	+	+	+	+	+	+	+	+	+	+	+
Intestine large, cecum	+	+	+	+	+	+	+	+	+	+	+	+	+	A	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed Intestine large, colon	+	+	+	+	+	+	+	+	+	4	+	+	+	+	X +	+	+	+	+	+	+	+	+	+	+
Intestine large, rectum	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	÷	+	+	+	+	+
Intestine small	+	+	+	+	+	+	+	+	+	+	+	+	+	A A	+	+	+	+	+	+	+	++	+	+	+
Intestine small, duodenum Intestine small, ileum	+	Ŧ	+	+	+	+	+	+	+	+	+	+	+	Ã	+	+	+	+	+	+	+	+	+	+	+
Intestine small, jejunum	+	+	+	+	+	+	+	+	+	+	+	+	+	A	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed Liver	1 _	+	+	_	4	-	_	_	Δ	_	4	+	+	+	_	+	_	4	+	_	_	+	+	+	X +
Hemangioma Hemangiosarcoma		•							••		·	x		Ů	·				·	Ţ,		·			·
Hepatocellular carcinoma Hepatocellular adenoma	- (X		X		25													x
Lymphoma malignant lymphocytic Lymphoma malignant mixed													х		x				x	x					••
Mesentery			+										••		^				+	+					
Lymphoma malignant lymphocytic Lymphoma malignant mixed																			X	X					
Pancreas Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	+	+	+	+	*	+	+	+	*	+	+	+	+	+	+
Salivary glands	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant histiocytic Lymphoma malignant lymphocytic															_										
Lymphoma malignant mixed Stomach	1 +	+	+		_	+	_	4	_	_	4	_	+	_	X	+	_	+	+	_	+	+	+	+	+
Lymphoma malignant lymphocytic	'	•		,	•				•			,				•				•	•	•	·		•
Stomach, forestomach Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	+	+	+	+	*	+	+	+	+	+	+	+	+	+	+
Papilloma squamous Stomach, glandular Lymphoma malignant mixed	+	+	+	+	+	+	x	+	+	+	+	+	+	A	X	+	+	+	* X	+	+	+	+	+	+
CARDIOVASCULAR SYSTEM Blood vessel								_															_		
Lymphoma malignant lymphocytic	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Heart	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant lymphocytic Lymphoma malignant mixed																									
ENDOCRINE SYSTEM	_																								
Adrenal gland Adrenal gland, cortex	+	+	+	+	+	+	+	M	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed							•												X						
Adrenal gland, medulla Pheochromocytoma, NOS	+	+	+	+	+	+	+	M	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Islets, pancreatic	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Parathyroid gland	M	M	+	M	+	+	+	M	+	+	+	M	M	+	M	+	M	M	M	+	+	+	+	+	M
Pituitary gland Pars distalis, adenoma	+	+	+	+	M	+	+	+	+	+	X	+	+	+	+	X X	+	+	+	+	+	+	+	+	*X
Thyroid gland Follicle, adenocarcinoma	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
GENERAL BODY SYSTEM None								-		·					-										
GENITAL SYSTEM	_																			_					
Clitoral gland																	+								
Ovary Hemangnoma	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Luteoma Lymphoma malignant mixed							v			v					v										
Uterus	+	+	+	+	+	+	X +	+	+	X +	+	+	+	+	X +	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant lymphocytic							•-																		
	1						X																		
Lymphoma malignant mixed Endometrium, polyp stromal	Į.																								

⁺ Tissue examined microscopically
Not examined
- Present but not examined microscopically
I Insufficient tissue

M Missing
A Autolysis precludes examination
X Incidence of listed morphology

TABLE D2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF FEMALE MICE: CHAMBER CONTROL (Continued)

WEEKS ON STUDY	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	TOTA
CARCASS ID	1 0 2 1	1 1 6	1 1 9	0 7 8	0 7 9	0 9 2 1	9 4 1	1 0 4 1	1 0 6	1 0 7 1	0 7 1	1 0 0	1 0 8	1 1 7 1	1 2 0	0 6 5	0 6 6	0 7 3	0 8 6	0 9 3	0 6 8	0 7 6	0 8 2 1	0 8 7 1	1 1 5	TISSU
LIMENTARY SYSTEM	-				_	<u> </u>									_				_		_					-
sophagus allbladder	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	49 49
Lymphoma malignant mixed	1	7			Τ.	X	X	т	т	_	т	т	7	'									•	,	•	4
testine large	+ +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 49
testine large, cecum Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	1
testine large, colon	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
testine large, rectum	++	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 49
testine small, duodenum	17	+	+	+	+	+	+	+	+	+	+	+	+	+	÷	+	+	+	+	+	+	÷	+	+	<i>+</i>	49
itestine small, ileum	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	49
testine small, jejunum Lymphoma malignant mixed	+	+	+	+	+	+ X	+	+	+	+	+	+	*	+	+	+	+	+	+	+	+	+	+	+	+	49
ver	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+ X	49
Hemangioma Hemangiosarcoma	ļ																								Λ	1
Hepatocellular carcinoma	1				X							Х														4
Hepatocellular adenoma Lymphoma malignant lymphocytic						X								х			X	x								3 2
Lymphoma malignant nixed					X	X	X	X										4.								8
lesentery														+							+					5
Lymphoma malignant lymphocytic Lymphoma malignant mixed														Х							х					1 3
ancreas	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Lymphoma malignant mixed								X	X										X		X					6
alivary glands Lymphoma malignant histiocytic	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	*	+	+	+	+	+	50
Lymphoma malignant lymphocytic														X				X		**						2
Lymphoma malignant mixed	1.				X					X									X							4
tomach Lymphoma malignant lymphocytic	+	+	+	+	+	+	+	+	+	+	+	+	+	X	+	+	+	+	+	+	+	*	+	+	+	50
tomach, forestomach	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Lymphoma malignant mixed Papilioma squamous							X												X	Х				¥		3
tomach, glandular	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	49
Lymphoma malignant mixed	-					X				Х									X							6
ARDIOVASCULAR SYSTEM	-																				-					-
lood vessel	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Lymphoma malignant lymphocytic leart	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	A.	+	+	+	+	+	+	+	50
Lymphoma malignant lymphocytic				•			,	,	'	,				X	,											1
Lymphoma malignant mixed										X																1
NDOCRINE SYSTEM	-				_																					-
drenal gland	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	49 49
drenal gland, cortex Lymphoma malignant mixed	*	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	1
idrenal gland, medulla	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	49
Pheochromocytoma, NOS siets, pancreatic	1	_	_	_	_			_	_ــ		_	_	_	X +	+	_	+	_	_	_	4	_	1	4	+	50
arathyroid gland	M	+	+	M	+	M	+	+	+	+	+	M	+	+	M	+	+	M	+	+	+	+	+	+	M	32
ituitary gland	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	49
Pars distalis, adenoma hyroid gland	+	+	+	X	+	+	+	X +	+	X	+	+	X +	+	+	+	X	+	+	X	X +	+	X	+	X +	12 50
Follicle, adenocarcinoma		·		,	x	,						,	•	•	•				•		,	,				1
ENERAL BODY SYSTEM	-			•						•••																-
ENITAL SYSTEM	-																									-
htoral gland																										1
vary	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Hemangioma Lutaoma									v	X																1 1
Luteoma Lymphoma malignant mixed									X																	4
terus	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Lymphoma malignant lymphocytic									x									X								$\frac{1}{2}$
Lymphoma malignant mixed Endometrium, polyp stromal									А		X		Х		X											3
agina	1																									ĭ

TABLE D2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF FEMALE MICE: CHAMBER CONTROL (Continued)

					` -				,																
WEEKS ON STUDY	0 0 1	0 5 0	0 5 4	0 5 7	0 6 7	0 8 0	0 9 0	0 9 1	0 9 1	0 9 1	0 9 4	0 9 8	9 8	0 9 9	1 0 2	1 0 2	1 0 2	1 0 3	1 0 3	1 0 4	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5
CARCASS ID	1 1 3 1	1 1 2 1	0 8 3 1	1 1 8 1	0 9 1	9 8 1	0 6 3 1	0 8 9	7 7 1	0 9 0	0 9 6 1	1 1 1	1 0 5 1	0 9 9	0 8 8	0 8 4 1	1 0 3 1	0 6 4 1	0 7 0 1	0 7 4 1	0 6 2 1	0 7 2 1	0 8 1 1	0 8 5 1	0 9 1
HEMATOPOIETIC SYSTEM	- -																								
Blood Bone marrow	_	_	_	_	_	_	+	_	_	_	_	_	_	_		+	+	_	+	+	+	+	+	+	+
Lymphoma malignant lymphocytic	*	_			7	т.	_	т	т	т		т	Τ.	т	т	,	1.	7	,	,	•	•	,	,	_
Lymphoma malignant mixed Lymph node	М	+	+	+	+	+	+	+	M	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	X
Lymphoma malignant mixed	***		•	•	·		•	,	***	,	•		·	·	X		·				•	•			
Axiliary, lymphoma malignant mixed Bronchial, lymphoma malignant mixed															х					X					
Iliac, lymphoma malignant mixed Mediastinal, lymphoma malignant mixed Mesenteric, lymphoma malignant													X		X					X					X X
lymphocytic Mesenteric, lymphoma malignant mixed										x			х		x				х	x				X	x
Renal, lymphoma malignant mixed	١.,								.,	•-					X					X					X
Lymph node, mandibular Lymphoma maiignant lymphocytic	M	+	+	+	+	+	+	+	M	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed Spleen		_	_	_	_	_	_	_	_	_	_	_	X	_	X	_	_	_	X	_	_		_	X	X
Hemangiosarcoma		-	-	т		-		т	_		Т	,	,	•	-	,	•	•	,	•	,		,	,	,
Lymphoma malignant lymphocytic Lymphoma malignant mixed										x			X		x				x	X				X	х
Thymus	+	+	+	+	M	+	+	+	+	+	+	M	+	M		M	+	+	+	+	+	+	+	+	+
Lymphoma malignant lymphocytic Lymphoma malignant mixed	1						X						X		Х				X	X				X	X
INTEGUMENTARY SYSTEM	—		_														_								
Mammary gland Adenocarcinoma	+	+	+	+	+	+	M	+	+	+	+	+	+	+	+	+	+	*	+	+	+	+	+	+	+
Skin	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
MUSCULOSKELETAL SYSTEM Bone	— -	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
NERVOUS SYSTEM	- -																								
Brain Lymphoma malignant mixed	_ +	+		+	+	+	+	+	+	X	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
RESPIRATORY SYSTEM Lung	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Alveolar/bronchiolar adenoma Hepatocellular carcinoma, metastatic, liver			•			·		·		X	,	X				X									
Lymphoma malignant histiocytic Lymphoma malignant lymphocytic	}									•									v						
Lymphoma malignant mixed Nose	+	+	+	+	+	+	A +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Trachea	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
SPECIAL SENSES SYSTEM None										_														_	
URINARY SYSTEM	_																								
Kidney Lymphoma malignant lymphocytic	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed Urinary bladder			J.	,	J.		,		ن	ر	_		X	_	X	_		د	X	X	د	4	4	X	+
Lymphoma malignant lymphocytic	†	+	*	+	+	+	+	+	+	+	+	+		+	+	+	+	+	+	-	+	+		+	
Lymphoma malignant mixed													X							Х				X	X
	'																				_				

TABLE D2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF FEMALE MICE: CHAMBER CONTROL (Continued)

								``	••••		reu	,														
WEEKS ON STUDY	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	0 5	1 0 5	TOTAL						
CARCASS ID	1 0 2 1	1 1 6 1	1 1 9 1	0 7 8 1	0 7 9	0 9 2 1	0 9 4 1	1 0 4 1	0 6 1	1 0 7 1	7 1 1	1 0 0 1	1 0 8 1	1 1 7 1	1 2 0 1	0 6 5	0 6 6	0 7 3 1	0 8 6 1	0 9 3 1	0 6 8 1	0 7 6 1	0 8 2 1	0 8 7 1	1 1 5 1	TISSUES
HEMATOPOIETIC SYSTEM																										
Blood Bone marrow Lymphoma malignant lymphocytic Lymphoma malignant mixed Lymph node	+	+	+	+	+	+	+	+ X +	+	+	+	+	+	* *	+	+	+	+	+	+	+	+	+	+	+	1 50 1 2 48
Lymphoma malignant mixed Axillary, lymphoma malignant mixed Bronchial, lymphoma malignant mixed Iliac, lymphoma malignant mixed		·		•	•				X			·	·		,		X									3 1 1 3
Mediastinal, lymphoma mal mixed Mesenteric, lymphoma malignant lymphocytic					X									x				x								5 2
Mesenteric, lymphoma malignant mixed Renal, lymphoma malignant mixed						X	X	X	X	X				-					X		X					14
Lymph node, mandibular Lymphoma malignant lymphocytic	+	+	+	+	+	+	+	+	+	+	+	+	+	M	+	+	+	+ X	+	+	+	+	+	+	+	47
Lymphoma malignant mixed Spleen Hemangiosarcoma	+	+	+	+	X +	X +	X +	+	X +	+	+	+	+	+	+	+	+	+	X +	+	Х * Х	+	+	+	+	11 50 1
Lymphoma malignant lymphocytic Lymphoma malignant mixed Thymus Lymphoma malignant lymphocytic	+	+	+	+	X + X	X +	X +	X +	X +	Х + Х	X + X	+	+	x x	+	+	Х + Х	*	Х + Х	X +	Х + Х	+	+	+	+	18 46 2 15
Lymphoma malignant mixed INTEGUMENTARY SYSTEM	_					Х	X 																			
Mammary gland Adenocarcinoma Skin	+	+	+	+	+	+	+	+	+	+	+	X X	+	+	+	+	+	+	+	+	+	+	+	+	+	49 2 50
MUSCULOSKELETAL SYSTEM Bone	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
NERVOUS SYSTEM Brain Lymphoma malignant mixed	+	+	+	+	+	+	+	+	*	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 2
RESPIRATORY SYSTEM Lung Aiveolar/bronchiolar adenoma Hepatocellular carcinoma, metastatic.	+	+	+	+	+	+	+	*	* X	+	+	+	+	+	+	+	*	+	+	+	+	+	+	+	+	50 5
liver Lymphoma malignant histocytic Lymphoma malignant lymphocytic Lymphoma malignant mixed Nose Trachea	++	+	+	<u> </u>	X +	X + +	*	+	X +	X + +	+	++	+	x + +	+	+	+	x +	X + +	++	X + +	+ +	+	+	+ +	1 1 2 10 50 50
SPECIAL SENSES SYSTEM None	-											•														-
URINARY SYSTEM Kidney Lymphoma malignant lymphocytic	+	+	+	+	+	+	+	+	+	+	+	+	+	+ X	+	+	+	+	+	+	+	+	+	+	+	50
Lymphoma malignant mixed Urinary bladder Lymphoma malignant lymphocytic	+	+	+	+	+	X +	+	+	X +	X +	X +	+	+	+ X	+	+	+	*	X +	+	X +	+	+	+	+	11 50 2
Lymphoma malignant mixed						X	X		X	X									X	X						10

TABLE D2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF FEMALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE: 120 ppm

				-			_		_					-	PF											
WEEKS ON STUDY		0 0 2	0 5 6	0 5 6	0 5 6	0 6 2	0 6 5	0 6 6	0 6 9	0 7 7	0 8 6	9 3	0 9 4	9 8	0 9 8	0 9 9	9	1 0 4	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5
CARCASS	-	7	4	7	7	5	4	7	4	4	4	3	7	5	3	7	3	6	4	4 2	4	6	6	4 2	4 2	4
ID		7	9	0 1	1	6	8 1	6	5 1	4	9	3	4	9	6 1	9	5	8	7 1	9 1	1	2	7 1	1	6 1	3
ALIMENTARY SYSTEM																										
Esophagus Gallbladder	1	+	+	+	+	+	+	+	+	+	+	+	+ M	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed		т.	_	_			_	т			_		IAT	_	X	Τ		X	τ.		-		-			-
Intestine large	1	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine large, cecum Lymphoma malignant mixed	1	+	+	+	+	+	+	+	+	+	+	+	+	+	X	+	+	+	+	+	+	+	+	+	+	+
Intestine large, colon	1	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine large, rectum Intestine small	j	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine small, duodenum	1	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine small, ileum		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine small, jejunum Lymphoma malignant mixed	1	+	+	+	+	+	+	+	+	+	X +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Liver		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Hepatocellular carcinoma									X						X											
Hepatoceliular adenoma Lymphoma malignant mixed	- 1													X	X		X	х								
Mesentery	1													X X	+			Α.								
Lymphoma malignant mixed															X											
Pancreas Lymphoma malignant mixed		+	+	+	+	+	+	+	+	+	+	+	+	+	X	+	+	+	+	+	+	+	X	+	+	+
Salivary glands		+	+	+	+	+	+	+	+	+	+	+	+	+	X X	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed														X		4		X					,			
Stomach Stomach, forestomach	1	+	+	+	+	+	+	+	+	+	M	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed																										
Papilloma squamous Glandular, lymphoma malignant mixed	}														х											
Stomach, glandular		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed														*												
Tooth	1																									
CARDIOVASCULAR SYSTEM														_												
Blood vessel		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Heart Lymphoma malignant mixed		+	+	+	+	+	+	+	+	+	+	+	+	*X	X	+	+	X	+	+	+	+	+	+	+	+
ENDOCRINE SYSTEM																										
Adrenal gland		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Adrenal gland, cortex		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Adrenal gland, medulia Pheochromocytoma malignant		+	+	+	+	+	+	+	+	+	*	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Pheochromocytoma, NOS	1										Α.															
Islets, pancreatic		+	+	+	+	+	+	M	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Parathyroid gland Pituitary gland		M	+	M	+	+	M	M	+	M M	M +	M +	M +	M +	M +	M	M	M +	M +	+	M	M	M	+	M +	M +
Pars distalis, adenoma	1	,		,				,	•		•	•	X	X	X	X +		•	X	•		,	,			X
Pars distalis, lymphoma malignant														•												
mixed Pars intermedia, adenoma	}													X												
Thyroid gland	ĺ	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed														X												
GENERAL BODY SYSTEM					_																	_				
None																										
GENITAL SYSTEM						_																				
Clitoral gland	ĺ	+																								
Ovary Granulosa cell tumor benign		+	+	+	+	+	+	+	+	×	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed)									Λ					Х											
Uterus		+	+	+	+	+	+	+	+	+	+	+	+ X	+	+	+	+	+	+	+	+	+	+	+	+	+
Adenocarcinoma Hemangiosarcoma	1												X										х			
Leiomyoma	ļ																						А			
Lymphoma malignant mixed																		X								
Endometrium, polyp stromal																		Х								

TABLE D2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF FEMALE MICE: 120 ppm (Continued)

												•														
WEEKS ON STUDY	1 0 5	0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	0 5	1 0 5	1 0 5	0 5	1 0 5	1 0 5	1 0 5	1 0 5	0 5	1 0 5	1 0 5	1 0 5	TOTAL
CARCASS ID	4 4 1	4 5 0 1	4 6 3 1	7 5 1	4 2 2 1	2 3 1	4 3 1 1	4 4 2 1	4 4 6 1	4 5 5 1	4 6 5 1	3 9 1	4 5 3 1	4 5 7 1	4 6 1	4 6 4 1	7 2 1	4 7 8 1	4 2 4 1	4 2 5 1	4 2 8 1	4 3 0 1	4 5 8 1	4 6 6	8 0 1	TISSUES TUMORS
ALIMENTARY SYSTEM																										
Esophagus Gallbladder	+ +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 49
Lymphoma malignant mixed Intestine large		+	+	4	+	+	4	+	+	+	+	4	+	+	_	+	+	+	+	X	+	+	+	+	+	3 50
Intestine large, cecum	+	+	+	+	+	+	+	÷	+	+	÷	+	+	÷	+	÷	+	+	+	+	+	+	+	+	+	50
Lymphoma malignant mixed Intestine large, colon	1	_	_	_	_	_	_	_	_	_	_	_	_	+	+	_	_	_	_	+	_	_	_	+	+	50
Intestine large, colon Intestine large, rectum	+	+	+	+	+	+	+	+	+	+	+	+	+	+	Ŧ	+	+	+	+	+	+	+	+	+	+	50
Intestine small	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Intestine small, duodenum Intestine small, ileum	++	+	+	+	++	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 50
Intestine small, jejunum Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	÷	+	÷	+	+	+	50
Liver	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 2
Hepatocellular carcinoma Hepatocellular adenoma Lymphoma malignant mixed		X					X		X			x							X	x						7 6
Mesentery Lymphoma malignant mixed																										2 2
Pancreas Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	*	+	+	+	+	+	+	50
Salivary glands	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Lymphoma malignant mixed Stomach	1	_	4	_	_	+	_	_	4	_	_	_	_	_	_	_	_	_	_	X	+	+	+	+	+	50 50
Stomach, forestomach	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	49
Lymphoma malignant mixed Papilloma squamous												x								X						1 1
Glandular, lymphoma malignant mixed	ļ											Λ														1 1
Stomach, glandular	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 2
Lymphoma malignant mixed Tooth																				X +						1
CARDIOVASCULAR SYSTEM												_														
Blood vessel	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Heart Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 3
ENDOCRINE SYSTEM																										
Adrenal gland	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50
Adrenal gland, cortex Adrenal gland, medulla	+	+	+	+	+	+	+	+	+	+	+	++++	+	+	+	+	+	+	+	+	+	+	+	+	+	50 50
Pheochromocytoma malignant Pheochromocytoma, NOS	_	_	_		_	T	*		_	_	_	_	_	т	_	X	_	_	_	_		_	_	т.	_	1 1
Islets, pancreatic	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	49
Parathyroid gland Pituitary gland	M +	M +	M +	M +	+	M	M	M +	M +	+	+	M +	M +	M	M +	M +	M +	M	+	+	M +	M +	+	M +		12 48
Pars distalis, adenoma Pars distalis, lymphoma malignant	'	,	·	X	X	X	*	X	·	*	X	·	·	*	•	·	·	X		X	·	X	X		*	19
mixed Pars intermedia, adenoma	1																	x								1
Thyroid gland Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 1
GENERAL BODY SYSTEM None										-	_								<u>-</u>		-					
GENITAL SYSTEM						_		_																		-
Clitoral gland Ovary	+	+	+	+	+	+	+	+	+	м	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	1 49
Granulosa cell tumor benign	'	•	•	,	•			,			•	,				•	·	•		,	,	•	,		•	1
Lymphoma malignant mixed Uterus	1	_	_	1	_	_	_	4	_	_	_	_	_	_	_		_	_	_		4	4	+	_	4	50
Adenocarcinoma	_	~	_	~	~	-	~	~	~		т		Τ.	-		_	_		Τ'	7	*	7.	*	*	-	1
Hemangiosarcoma				v																						1 1
Leiomyoma Lymphoma malignan; == xed	1			X																						1
Endometrium, polyp s a.								X																		2
	1																									1

TABLE D2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF FEMALE MICE: 120 ppm (Continued)

					•				•																
WEERS ON STUDY	0 0 2	0 5 6	0 5 6	0 5 6	0 6 2	0 6 5	0 6 6	0 6 9	0 7 7	0 8 6	0 9 3	0 9 4	0 9 8	0 9 8	0 9 9	0 9 9	1 0 4	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5
CARCASS ID	7 7 1	6 9 1	7 0 1	4 7 1 1	4 5 6 1	4 4 8 1	7 6 1	4 5 1	4 5 4 1	4 4 9 1	4 3 3 1	4 7 4 1	5 9 1	4 3 6 1	4 7 9 1	3 5 1	4 6 8 1	4 2 7 1	2 9 1	4 4 1 1	4 6 2 1	4 6 7 1	4 2 1 1	4 2 6 1	4 4 3 1
HEMATOPOIETIC SYSTEM Blood Leukemia																									
Bone marrow Hemangioma	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed Lymph node Adenocarcinoma, metastatic, uterus	М	+	+	+	+	+	+	+	+	+	+	x	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed Iliac, lymphoma malignant mixed Mediastinal, lymphoma malignant mixed Messenteric, lymphoma malignant mixed Renal, lymphoma malignant mixed										x			X	X X			X X X					х	X X X		
Lymph node, mandibular Lymphoma malignant mixed Spieen Adenocarcinoma, metastatic, uterus	M +	+	+	+	+	+	+	M +	+	+	+	+ + X	* X +	+ X +	+	+	X +	+	+	+	+	+ X +	+ X +	+	+
Hemangiosarcoma Lymphoma malignant mixed Thymus Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	M	M	X + X	X + X	+	+	х + х	+	+	+	+	Х + Х	X + X	+	+
INTEGUMENTARY SYSTEM Mammary gland Adenoacanthoma	+	+	+	+	+	+	+	+	+	+	* X	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Skin MUSCULOSKELETAL SYSTEM	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Bone Skeletal muscle Head, sarcoma, deep invasion	+	+	+	+	+	+	+	+	+ *	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
NERVOUS SYSTEM Brain Lymphoma malignant mixed Choroid plexus, lymphoma malignant	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
mixed Meninges, sarcoma, metastatic									x					X											
RESPIRATORY SYSTEM Lung Alveolar/bronchiolar carcinoma	+	+	+	+	+	+	+	+	+	+	+	+	+	+ X	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed Nose Trachea	+	+ +	++	+	+	+	++	++	++	++	++	++	X + +	X + +	++	++	X + +	+	++	++	++	X + +	+	++	++
SPECIAL SENSES SYSTEM Eye Hardenan gland Adenoma							· ·	-	+	_		+						-							
URINARY SYSTEM Kidney Lymphoma malignant lymphocytic	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	* X	+	+	+	+	+	+	+	+
Lymphoma malignant mixed Ureter Urnary bladder Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	+	+	* X	X + X	+	+	+	+	+	+	+	X	X + X	+	+

TABLE D2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF FEMALE MICE: 129 ppm (Continued)

WEEKS ON STUDY	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	TOTAL.
CARCASS ID	4 4 4 1	5 0 1	4 6 3 1	7 5 1	4 2 2 1	4 2 3 1	3 1 1	4 4 2 1	4 4 6 1	4 5 5 1	4 6 5 1	3 9 1	5 3 1	5 7 1	6 1 1	4 6 4 1	7 2 1	4 7 8 1	2 4 1	4 2 5 1	4 2 8 1	4 3 0 1	4 5 8 1	4 6 6 1	8 0 1	TISSUES
HEMATOPOIETIC SYSTEM Blood Leukemia Bone marrow Hemangioma Lymphoma mailgnant mixed	*	* X + X	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	1 1 50 1 1
Lymph node Adenocarcinoma, metastatic, uterus Lymphoma malignant mixed Iliac, lymphoma malignant mixed Mediastinal, lymphoma malignant mixed Mesenteric, lymphoma malignant mixed Renal, lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	+	+	+	x	+	+	+	+	x	x x	+	+	+	+	+	49 1 2 3 2 8 3
Lymph node, mandibular Lymphoma malignant mixed Spleen	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	X	+	+	+	+	+	48 6 50
Adenocarcinoma, metastatic, uterus Hemangiosarcoma Lymphoma malignant mixed Thymus Lymphoma malignant mixed	+	X	+	+	+	X +	+	+	+	+	+	+	+	X + X	+	+	+	+	X + X	X + X	+	+	+	+	+	1 1 9 48 8
INTEGUMENTARY SYSTEM Mammary giand Adenoacanthoma Skin	++	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	50 1 50
MUSCULOSKELETAL SYSTEM Bone Skeletal muscle Head, sarcoma, deep invasion	+	+	+	+	+	+	+	+		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	49 1 1
NERVOUS SYSTEM Brain Lymphoma malignant mixed Choroid piexus, lymphoma malignant mixed Meninges, sarcoma, metastatic	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	*	+	+	+	+	+	+	50 1 1 1 1
RESPIRATORY SYSTEM Lung Alveolar/bronchiolar carcinoma Lymphoma malignant mixed Nose Trachea	+ + +	+ X + +	+ X + +	+ + +	+ + +	+ + +	+ + +	+ + +	+ + +	+ + +	+ + +	+ + +	+ +	+ X + +	* * *	+ +	+ + +	+ + +	+ X + +	+ X +	+ + +	+ + +	+ + +	+ + +	+ + +	50 3 8 50 50
SPECIAL SENSES SYSTEM Eye Harderian gland Adenoma										+ X														+ X		1 3 2
URINARY SYSTEM Kidney Lymphoma malignant lymphocytic Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+ X	+ X	+	+	+	+	+	50 1 5
Ureter Urinary bladder Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	*	*	+	+	+	+	+	50 5

TABLE D2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF FEMALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE: 600 ppm

															FF											
WEEKS ON STUDY		0 2 4	0 5 7	0 7 3	0 7 3	0 7 4	0 8 1	0 8 3	0 8 6	8 6	0 8 7	0 9 0	0 9 0	0 9 2	9	9 4	0 9 5	9 5	0 9 6	0 0	0 1	1 0 2	1 0 2	1 0 2	1 0 2	0 2
CARCASS ID		1 8 3 1	9 2 1	2 6 1	2 0 6 1	0 0 1	1 4 1	9 6 1	0 2 1	2 3 1 1	1 8 5 1	2 2 2 1	2 1 8 1	1 2 1	3 3 1	2 3 8 1	2 1 6 1	2 4 0 1	2 1 3 1	2 1 5 1	2 0 7 1	9 9 1	2 2 0 1	2 2 4 1	1 8 4 1	2 3 4 1
ALIMENTARY SYSTEM																										
Esophagus Gallbladder	Ì	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed		Ċ	·		·			·		·						X										
Lymphoma malignant undifferentiated cell type											X	X														
Intestine large Intestine large, cecum		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Histiocytic sarcoma		,	т.	•	_		7	Τ.	•	т	т	7	т		-	-	*		7	'	7	r	-	'		
Intestine large, colon Intestine large, rectum		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	++	+	+	+	+	+	+	+	+	+
Intestine small		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	++	+	+	+	+	+
Intestine small, duodenum Lymphoma malignant mixed Lymphoma malignant undifferentiated		+	+	+	+	+	*	+	+	+	+		+	*	+	+	+	+	_	*	+	+	+	+	+	+
cell type Intestine small, ileum		+	+	+	+	+	+	+	+	+	X +	X +	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine small, jejunum Liver	J	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Hepatocellular carcinoma		7		-		_	7	X	_	•	т	Τ.	_			т	_		•	-	т	*	_	7		т
Hepatocellular adenoma Histiocytic sarcoma Lymphoma malignant histiocytic					X																				х	
Lymphoma malignant mixed Lymphoma malignant undifferentiated										X				X		Х								X		Х
cell type											X	X									X					
Mesentery Lymphoma malignant											*								+							
Lymphoma malignant mixed	į										••															
Pancreas Lymphoma malignant mixed Lymphoma malignant undifferentiated		+	+	+	+	+	+	+	+	+	+	+	+	X	+	+	+	+	+	X	+	+	+	+	+	X
cell type Salivary glands Lymphoma malignant undifferentiated		+	+	+	+	+	+	+	+	+	+	X +	+	+	+	+	+	+	+	+	+	+	+	+	+	+
cell type Stomach		+	+	+	+	+	4	+	+	+	+	X +	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Histiocytic sarcoma													Ċ													
Stomach, forestomach Lymphoma malignant mixed Lymphoma malignant undifferentiated		+	+	+	+	+	+	+	+	+	+	+	+	X	+	+	+	+	+	7	+	+	+	+	+	+
cell type Papilloma squamous												X														
Stomach, glandular		+	+	+	+	+	+	+	+	+	+	+	+	+ X	+	+	+	+	+	+	+	+	+	+	+	*
Lymphoma malignant mixed Lymphoma malignant undifferentiated														A												Α.
cell type												X									X					
CARDIOVASCULAR SYSTEM	i				_									_									_			
Blood vessel Heart		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Histiocytic sarcoma		'	•		,		,	,		,			,	Ċ	•		·		•	,			,			
Lymphoma malignant mixed																								X		
ENDOCRINE SYSTEM																-										
Adrenal gland Adrenal gland-cortex		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Adenoma Lymphoma malignant undifferentiated																	X									
cell type											X															
Capsule, lymphoma malignant undifferentiated cell type												X														
Adrenal gland, medulla Pheochromocytoma NOS		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	M	+	+	+	+
Islets, pancreatic	ļ	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Parathyroid gland Pituitary gland	ļ	++	M +	+	+	M +	+	+	M +	+	M M	+	+	M +	M +	M +	M +	+	+	M +	+	M +	+	M +	M +	M +
Pars distalis, adenoma			X					X	v			X	X				X		X	X	X		X			X
Pars intermedia, adenoma Thyroid gland		+	+	+	+	+	+	+	X +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Follicle adenoma													X				X	X								
GENERAL BODY SYSTEM None							,																			
GENITAL SYSTEM							-4-												-							
Chtoral gland Ovary	,	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant mixed			•				•							X					,							X
Lumphoma malignant infact												X						_								
Lymphoma malignant undifferentiated cell type	1																	-								+
Lymphoma malignant undifferentiated cell type Oviduct Uterus		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	,
Lymphoma malignant undifferentiated cell type Oviduct Uterus Lymphoma malignant histiocytic		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	X	x
Lymphoma malignant undifferentiated cell type Oviduct Uterus Lymphoma malignant histocytic Lymphoma malignant mixed Lymphoma malignant undifferentiated		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	X	x
Lymphoma malignant undifferentiated cell type Oviduct Uterus Lymphoma malignant histiocytic Lymphoma malignant mixed	!	+	+	+	+	+	+	+	+	+	+	x	+	+	+	+	+	+	+	+	+	+	+	+	X	x

TABLE D2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF FEMALE MICE: 600 ppm (Continued)

NEKKS ON	1 1 0 5 5 5 2 2 1 1 3 0 2 1 1 1 1 + + + + + + + + + + + + + + +	TOTA TISSUE TUMOI + 50 1 + 50 1 + 50 1 + 50 + 50 + 50 + 50 + 50 + 50 + 50 + 50
CARCASS 1	1 3 0 2 1 1 1 + + + + + + + + + + + + + + + +	+ 50 + 50 + 50 + 50 + 50 + 50 + 50 + 50
ALIMENTARY SYSTEM Sephagus Isalibiader Lymphoma maignant mixed Lymphoma maignant mixed Lymphoma maignant undifferentiated cell type Intestine large Intestine large, cecum Intestine large, colon Intestine large, colon Intestine large, colon Intestine large, colon Intestine large, colon Intestine large, colon Intestine large, colon Intestine large, colon Intestine large, colon Intestine large, colon Intestine small, duodenum Lymphoma malignant mixed Lymphoma mixed Lymphoma mixed Lymphoma mixed Lymphoma mixed Lymphoma	+ + + + + + + + + + + + + + + + + + +	+ 50 + 50 1 2 + 50 + 50 + 50 + 50 + 50 + 50 + 50 + 50
Sophagus Jalibiadder Lymphoma maingnant mixed Lymphoma maingnant undifferentiated cell type Intestine large, cecum Historytic sarcoma Intestine large, colon Intestine large, colon Intestine large, colon Intestine large, rectum Intestine mail, duodenum Lymphoma mailgnant mixed Lymphoma mailgnant mixed Lymphoma mailgnant mixed Lymphoma mailgnant mixed Lymphoma mailgnant mixed Lymphoma mailgnant mixed Lymphoma mailgnant mixed Lymphoma mailgnant historytic Lymphoma mailgnant mixed Lymphoma mixed Lymphoma mixed Lymphoma mixed Lymphoma mixed Lymphoma mixed Lymphoma mixed	+ + + + + + + + + + + + + + + + + + + +	+ 50 1 2 + 50 + 50 + 50 + 50 + 50 + 50
Albiadder	+ + + + + + + + + + + + + + + + + + + +	1 2 + 50 + 50 + 50 + 50 + 50 + 50
The string large	+ + + + + + + + + + + + + + + + + + + +	+ 50 + 50 + 50 + 50 + 50 + 50 + 50
the state arge, cecum	+ + + + + + + + + + + + + + + + + + + +	+ 50 1 + 50 + 50 + 50 + 50 + 50
Histocytic sarcoma	+ + + + + + + + + + + + + + + + + + + +	+ 50 + 50 + 50 + 50
+ + + + + + + + + + + + + + + + + + +	+ + + + + + + + + + + + + + + + + + + +	+ 50 + 50 + 50
## # # # # # # # # # # # # # # # # # #	+ + + + + + + + + + + + + + + + + + + +	+ 50 + 50
Lymphoma malignant mixed Lymphoma malignant undifferentiated cell type ntestine small, lelum the three small, lelum three sma	+ +	
+ + + + + + + + + + + + + + + + + + +	+ +	2
Hepatocellular carcinoma Hepatocellular adenoma Hepatocellular adenoma Histocytic sarcoma Lymphoma malignant mixed Lymphoma malignant mixed Lymphoma malignant mixed Lymphoma malignant mixed Lymphoma malignant mixed Lymphoma malignant mixed Lymphoma malignant mixed Lymphoma malignant mixed Lymphoma malignant mixed Lymphoma malignant mixed Lymphoma malignant mixed Lymphoma malignant mixed Lymphoma malignant mixed Lymphoma malignant mixed Lymphoma malignant undifferentiated cell type Salivary glands Lymphoma malignant undifferentiated Coll type Stomach Histocytic sarcoma Lymphoma malignant mixed Lymphoma malignant mixed Lymphoma malignant mixed Lymphoma malignant undifferentiated Coll type Lymphoma malignant mixed Lymphoma malignant mixed Lymphoma malignant mixed Lymphoma malignant mixed Lymphoma malignant undifferentiated Coll type Lymphoma malignant mixed Lymp	+ +	+ 50 + 50
Hejatocellular adenoma Kisticcytic sarcoma Lymphoma malignant mixed Lymphoma malignant undifferentiated cell type 4		+ 50
Lymphoma malignant mixed Lymphoma malignant undifferentiated cell type desentery		6 1 1
Lymphoma malignant Lymphoma malignant mixed	х	X 8
Lymphoma malignant mixed X X X X		1
Lymphoma malignant mixed Lymphoma malignant undifferentiated cell type salivary glands Lymphoma malignant undifferentiated cell type stomach Histocytic sarcoma Lymphoma malignant mixed Lymphoma malignant undifferentiated cell type *		+ 50
Lymphoma malignant undifferentiated cell type tomach + + + + + + + + + + + + + + + + + + +	+ +	3
tomach	+ +	+ 50
tomach, forestomach + + + + + + + + + + + + + + + + + + +	+ +	+ 50
Lymphoma malignant mixed Lymphoma malignant undifferentiated cell type	+ +	+ 50
		1
Papilloma squamous X tomach, glandular + + + + + + + + + + + + + + + + + + +	+ +	+ 1 50 2 2
CARDIOVASCULAR SYSTEM		
Blood vessel	+ +	+ 50 + 50
Hastr	+ +	1 1
ENDOCRINE SYSTEM		
Adrenal gland + + + + + + + + + + + + + + + + + +	+ +	+ 50
drenal gland, cortex + + + + + + + + + + + + + + + + + + +	+ +	+ 50
cell type Capsule, lymphoma malignant		1
undifferentiated cell type		+ 49
Adrenal gland, medulla	* *	1
slets, pancreatic	· + + [+ +	+ 50 + 18
Pututary gland	+ +	+ 49
Pars distalis, adenoma X X X X X X X X X X X X X X X X X X X	х х	X 21
+ + + + + + + + + + + + + + + + + + +	+ +	+ 50
ENERAL BODY SYSTEM None		_
ENITAL SYSTEM		
Chtoral gland Ovary		+ 50
Lymphoma malignant undifferentiated cell type	, ,	1
Dviduct	. + +	+ 50
Lymphoma malignant histocytic Lymphoma malignant mixed Lymphoma malignant undifferentiated cell type		1 1
Endometrium, polyp stromal agina		1

TABLE D2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF FEMALE MICE: 600 ppm (Continued)

WEEKS ON STUDY	0	0																							
	4	5 7	0 7 3	7 3	0 7 4	0 8 1	0 8 3	0 8 6	0 8 6	0 8 7	0 9 0	0 9 0	0 9 2	9 3	0 9 4	0 9 5	0 9 5	0 9 6	1 0 0	1 0 1	1 0 2	1 0 2	1 0 2	1 0 2	1 0 2
CARCASS ID	1 8 3 1	1 9 2 1	2 2 6 1	2 0 6 1	2 0 0 1	2 1 4 1	9 6 1	2 0 2 1	2 3 1 1	1 8 5 1	2 2 2 1	2 1 8 1	1 2 1	2 3 3 1	2 3 8 1	2 1 6 1	2 4 0 1	2 1 3 1	2 1 5 1	2 0 7 1	1 9 9	2 2 0 1	2 2 4 1	1 8 4 1	2 3 4 1
HEMATOPOIETIC SYSTEM Blood Bone marrow Hemangrosarcoma, metastatic, skin Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	* X	+	+	+	+
Lymph node Lymphoma malignant mixed Lymphoma malignant undifferentiated cell type	+	+	+	+	+	+	+	+	*	+ X	+	+	+	+	+	+	+	+	+	* X	+	+	+	+	+
Iliac, histiocytic sarcoma Iliac, lymphoma malignant mixed Iliac, lymphoma malignant undifferentiated cell type Mediastual, lymphoma malignant mixed											x		x		x x				X	x					x x
Mediastinal, lymphoma malignant undifferentiated cell type Mesenteric, histocytic sarcoma Mesenteric, lymphoma malignant mixed		x									x		x		x				x						x
Mesentenc, lymphoma malignant undifferentiated cell type Renal, histiocytic sarcoma Renal, lymphoma malignant mixed Renal, lymphoma malignant											x		x						x	x					
undifferentiated cell type Lymph node, mandibular Histocytic sarcoma Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	X +	+	* X	+	+ X	+	+	+	+ X	X +	+	+	+	+	+
Lymphoma malignant undifferentiated cell type Spleen Histocytic sarcoma Lymphoma malignant mixed	+	+ X	+	A	+	+	+	+	+ X	+	X +	+	+ X	+	+	+	+	+	+ X	+	+	+	+ X	+	+ X
Lymphoma malignant undifferentiated cell type Thymus Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	X	X	+	*	+	М	+	+	+	*	X +	+	+	+	+	* X
Lymphoma malignant undifferentiated cell type INTEGUMENTARY SYSTEM Mammary gland	- -										X										M				
Adenocarcinoma Skin Subcutaneous tissue, fibrosarcoma Subcutaneous tissue, hemangiosarcoma	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	X +	+	+	+	+	+ X	X	+	+	+
MUSCULOSKELETAL SYSTEM Bone Skeletal muscle	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
NERVOUS SYSTEM Brain Lymphoma malignant mixed	+	+	+	+	+	+	+	+	*	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
RESPIRATORY SYSTEM Lung Alveolar/bronchiolar adenoma Alveolar/bronchiolar carcinoma	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	*	+	*	+	+	+
Histocytic sarcoma Lymphoma malignant mixed Lymphoma malignant undifferentiated cell type Nose		_	_	_	_	_	_	_	x	×	X	_	x	_	x	_	_	_	X	_	_	_	X	+	x
SPECIAL SENSES SYSTEM Hardenan gland	- -	+	-	-	-	-	-	-	-	-	-	-	-	-	-	+	-	-	+	+	+	+	+	+	-
Adenoma Bilateral, adenocarcinoma URINARY SYSTEM	_																								
Kidney Histocytic sarcoma Lymphoma malignant mixed Lymphoma malignant undifferentiated cell type Urster	+	+	+	+	+	+	+	+	+	+	+ X	+	+ X	+	+ X	+	+	+	+ X	+	+	+	+	+	* X
	1		_		Ĩ	_	+	4	_				_	_	_	4	+	+	_	_		4		+	+

TABLE D2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF FEMALE MICE: 600 ppm (Continued)

								(0	oni		ieu	.,														
WEEKS ON STUDY	1 0 4	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	TOTAL.
CARCASS ID	2 0 1 1	1 9 5 1	2 0 3 1	1 1 1	3 6 1	3 9 1	1 8 1	1 8 2 1	2 0 8 1	2 1 7 1	2 2 3 1	1 8 7 1	1 9 0 1	2 0 4 1	2 2 1 1	2 2 7 1	9 1 1	1 9 4 1	2 9 1	2 3 7 1	1 8 6 1	1 8 9	1 9 3 1	2 1 0 1	2 3 2 1	TISSUES
HEMATOPOIETIC SYSTEM Blood Bone marrow Hemangosarcoma, metastatic, skin Lymphoma malignant mixed Lymph node Lymphoma malignant mixed Lymphoma malignant undifferentiated cell type	+++	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+ X +	+	+	+	+	+	+	+	1 50 1 1 50 1
llac, histocytic sarcoma Iliac, lymphoma malignant mixed Iliac, lymphoma malignant undifferentiated cell type Mediastinal, lymphoma mal. mixed Mediastinal, lymphoma malignant undifferentiated cell type Mesenteric, histocytic sarcoma Mesenteric, lymphoma malignant mixed		x x									x		x				x	x x		x x					X	1 5 2 4 1 2 10
Mesenteric, lymphoma malignant undifferentiated cell type Renal, histiocytic sarcoma Renal, lymphoma malignant mixed Renal, lymphoma malignant											x		x							x						2 2 3
undifferentiated cell type Lymph node, mandibular Histiocytic sarcoma Lymphoma malignant mixed Lymphoma malignant undifferentiated	+	+ X	+	+	+	+	+	+	+	+	+	+	*X	+	+	+	+	+	+	+ X	+	+	+	+	+	50 1 5
cell type Spleen Histiocytic sarcoma Lymphoma malignant mixed Lymphoma malignant undifferentiated	+	+ X	+	+	+	+	+	+	+	+	*	+	*X	+	+	+	+ X	+ X	+	+ X	+	+ X	+	+	+	1 49 2 11
cell type Thymus Lymphoma malignant mixed Lymphoma malignant undifferentiated cell type	+	*	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	M	+	+	+	+	+	3 48 4
INTEGUMENTARY SYSTEM Mammary gland Adenocarcinoma Skin Subcutaneous tissue, fibrosarcoma Subcutaneous tissue, hemangiosarcoma	+ + X	+	+	+	+	+	+	M +	+	+	+	+ *	+	+	+	+	+	+	+	+	+	+	+	+	+	48 2 50 2 1
MUSCULOSKELETAL SYSTEM Bone Skeletal muscle	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	++	+	+	+	+	+	+	50 2
NERVOUS SYSTEM Brain Lymphoma malignant mixed	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	*	+	+	+	+	+	+	+	+	50 2
RESPIRATORY SYSTEM Lung Alveolar/bronchiolar adenoma Alveolar/bronchiolar carcinoma Histocytic sarcoma Lymphoma malignant mixed Lymphoma malignant undifferentiated cell type	+	+ X X	+	+	+	+	+	+	+	+	+	+	+ X	+	*	+	+	+	+	+ x	+	+	+	+	+	50 3 1 1 8
Nose Trachea	++	+	++	+	+	+	+	+	+	++	++	+	++	++	++	+	++	+	+	+	+	++	+	+	+	50 50
SPECIAL SENSES SYSTEM Hardeman gland Adenoma Bilateral, adenocarcinoma					*			+ X																		2 1 1
URINARY SYSTEM Kidney Histocytic sarcoma Lymphoma malignant mixed Lymphoma malignant undifferentiated cell type	+	+	+	+	+	+	+	+	+	+	+	+	* X	+	+	+	+	+ X	+	+ X	+	+ X	+	+	+	50 1 7
Ureter Urnary bladder Lymphoma malignant mixed Lymphoma malignant undifferentiated cell type	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	*	*	+	+	+	+	+	+	+	1 50 4

TABLE D2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF FEMALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE: 1,200 ppm

	IMIIMI			•	31			OF	10				-,			Ьш										
WEEKS ON STUDY		0 0 1	0 1 2	0 3 3	0 5 8	0 7 0	0 7 0	0 7 0	0 7 5	0 7 5	0 8 5	0 8 5	0 8 6	0 8 9	0 9 6	1 0 0	1 0 2	1 0 3	1 0 3	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5
CARCASS ID		3 5 8 1	3 4 1	3 5 3 1	3 1 7 1	3 5 0 1	3 5 1	3 5 2 1	3 4 7 1	3 2 6 1	3 8 1	3 0 7 1	3 4 8 1	3 9 1	3 2 8 1	3 5 7 1	3 5 4 1	3 4 9	3 1 1 1	3 0 4 1	3 0 9	3 1 5 1	3 2 2 1	3 2 7 1	3 7 1	3 4 5 1
ALIMENTARY SYSTEM Esophagus Gallbladder Intestine large		+ + +	+ A +	+ +	+ + +				+ + +	+ + +	+ + +	+ + +	+ + +	+ + +	+++	+ + +	+ + +	+ + +	+ + +	+ + +	+ + +	+ + +	+ + +	+ + +	+ + +	+ + +
Intestine large, cecum Lymphoma malignant histocytic Lymphoma malignant lymphocytic Intestine large, colon		+	+	+	+				+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine large, rectum Lymphoma malignant lymphocytic Intestine small	ļ	M +	M +	+	+				+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Intestine small, duodenum Lymphoma malignant lymphocytic Intestine small, ileum Intestine small, jejunum		+++	+++	+++	+++				++	+++	+++	+++	+++	+++	+ +	+++	+++	+++	+++	+++	+++	+++	+++	++++	+ + +	+++
Lymphoma malignant undifferentiated cell type Liver Hepatocellular carcinoma		+	+	+	+				+	+	+ X	+	+	*X	X	+	+	+	*	+	+	+	+	+	+	+
Hepatocellular adenoma Histiocytic sarcoma Lymphoma malignant histiocytic Lymphoma malignant lymphocytic Lymphoma malignant undifferentiated									х								x	Х					x x			
cell type Mesentery Hepatocellular carcinoma, metastatic, liver											+ X				X	+										
Osteosarcoma, metastatic, uncertain primary site Pancreas Lymphoma malignant histocytic		+	+	+	+				+	+	+	+	+	+	+	X +	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant lymphocytic Salivary glands Lymphoma malignant lymphocytic Stomach		+	+	+	+				+	+	+	+	+	+	•	+	+	+	+	+	+	+	+	M +	+	++
Stomach, forestomach Lymphoma malignant lymphocytic Papilloma squamous Stomach, glandular		+	+	+	+				+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant histocytic Lymphoma malignant lymphocytic Lymphoma malignant mixed																										
CARDIOVASCULAR SYSTEM Blood vessel Heart Hemangiosarcoma Hemangiosarcoma, metastatic, ovary		++	+	+	+				+	+	+	++	+	+ +	+ + X	+	++	+	++	+	+	+	+	+	++	++
Lymphoma malignant lymphocytic ENDOCRINE SYSTEM																										
Adrenal gland Adrenal gland, cortex Adenoma Adrenal gland, medulla		+	+	+	+				+	+	+	+	+	+	+	+	+	+	+	+	+ X +	+	+	+	+	+
Pheochromocytoma, NOS Islets, pancreatic Parathyroid gland Pituitary gland		+ M +	+++++	+++	+++				+ M +	+ M +	+ M +	+ + M	++++	+ M +	++++	+ M +	+++	X + M +	+ M +	+ M +	+ M +	+ M + X	+ M + X	+ M +	+ M +	+ M + X
Pars distalis, adenoma Pars intermedia, adenoma Thyroid gland		+	+	+	+				+	+	+	+	+	+	+	+	+	+	X +	+	+	+	+	+	+	+ -
GENERAL BODY SYSTEM None GENITAL SYSTEM																										
Clitoral gland Ovary Hemangioma Hemangiosarcoma		++	+	+	+				+	+	+	+	+	+	+ X	+	+	+	+	+	+	+	+	+	+	+
Hemangiosarcoma, metastatic Histocytic sarcoma Lymphoma malignant histocytic Lymphoma malignant lymphocytic Uterus		1		,							,	,	X	4	,	,	X				,			J	,	_
Hemangiosarcoma Histocytic sarcoma Lymphoma malignant lymphocytic Endometrium, polyp stromai			_	7	*				т	7	T	т	•	_	•	т	x	+	7	*	X	+	*	7	•	T

TABLE D2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF FEMALE MICE: 1,200 ppm (Continued)

								(0	OII	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	uea	,														
WEEKS ON STUDY	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	TOTAL
CARCASS ID	3 0 2 1	3 0 6 1	3 2 4 1	3 2 5 1	3 5 1	3 1 6 1	3 3 2 1	3 3 3	3 6 1	3 4 0 1	3 4 1 1	3 0 3 1	3 1 4 1	3 1 9	3 3 1 1	3 4 3 1	3 4 4 1	3 5 9	3 0 8 1	3 1 0 1	3 2 0 1	3 2 3 1	3 3 0 1	3 4 6 1	3 6 0 1	TISSUES TUMORS
ALIMENTARY SYSTEM Esophagus Galibladder Intestine large Intestine large, cecum	+ + + +	+ + + +	+ + + +	+ + + +	+ + + +	+ + + +	+ + + +	+ + + +	+ M + +	+ + + +	+ + + +	+ + +	+ + + +	+ + +	+ + + +	+ + + +	+ + + +	+ + + +	+ + + +	+ + + +	+ + + +	+ + +	+ + +	+ + + +	+ + + +	47 45 47 47
Lymphoma malignant histrocytic Lymphoma malignant lymphocytic Intestine large, colon Intestine large, rectum Lymphoma malignant lymphocytic	X + X	+	X + +	++	++	++	++	++	+	++	++	++	++	+	+	+	++	++	++	++	++	++	+	++	++	1 1 47 45 1
Intestine small Intestine small, duodenum Lymphoma malignant lymphocytic Intestine small, ileum	+ X +	+++++++++++++++++++++++++++++++++++++++	+ + +	+++++	+ + + +	+ + +	+++++++++++++++++++++++++++++++++++++++	+++++++++++++++++++++++++++++++++++++++	+++++++++++++++++++++++++++++++++++++++	+++++++++++++++++++++++++++++++++++++++	+ + +	+++++++++++++++++++++++++++++++++++++++	+ + +	++++++	+ + +	+++++++++++++++++++++++++++++++++++++++	+++++++++++++++++++++++++++++++++++++++	+ + +	+ + +	+++++	+++++++++++++++++++++++++++++++++++++++	+++++++++++++++++++++++++++++++++++++++	+++++++++++++++++++++++++++++++++++++++	+ + +	+ + +	47 47 1 47
Intestine small, jejunum Lymphoma malignant undifferentiated cell type Liver Hepatocellular carcinoma	+ *	+ + X	+	+	+	+	+	+	+ *	+	+ *	+	+	+	+	+	+	+	+	+	+	+	+	+	+	47 1 47 7
Hepatocellular adenoma Histocytic sarcoma Lymphoma malignant histocytic Lymphoma malignant lymphocytic Lymphoma malignant undifferentiated cell type Mesentery Hepatocellular carcinoma, metastatic.	x	Λ	x				X	x	â		Λ							X								7 1 1 2 1 2
hver Osteosarcoma, metastatic, uncertain primary site Pancreas Lymphoma malignant histocytic	+	+	+ X	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	1 1 47 1
Lymphoma malignant lymphocytic Salivary glands Lymphoma malignant lymphocytic Stomach	X + X + +	+	+	+	+	+	+	+	+	+	+	+	+	+ +	+	+	+	+	+	+	+	+	+	+	+	1 46 1 47
Stomach, forestomach Lymphoma malignant lymphocytic Papilloma squamous Stomach, glandular Lymphoma malignant histiocytic Lymphoma malignant lymphocytic Lymphoma malignant mixed	* x + x	+ X +	+ X	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+ + X	+	+ X +	* X +	+	+	+	+	47 1 3 47 1 1
CARDIOVASCULAR SYSTEM Blood vessel Heart Hemangiosarcoma Hemangiosarcoma, metastatic, ovary Lymphoma malignant lymphocytic	+ + X	+ +	++	++	++	+	+ + X	++	++	++	++	++	++	++	++	++	++	++	++	++	++	++	++	++	+ +	47 47 1 1
ENDOCRINE SYSTEM Adrenal gland Adrenal gland, cortex Adenoma	++	++	++	++	++	++	++	++	++	+	++	++	++	++	++	++	++	++	++	++	++	+	++	+	++	47 47 1
Adrenal gland, medulia Pheochromocytoma, NOS Islets, pancreatic Parathyroid gland Pituitary gland	+ M +	+ M +	+	+	+ + M +	+ + M +	+ M +	+ + M +	+ + M +	+ M + X	+ + M +	+ M +	+ M + X	+ + M +	+ X + + X	+ + + +	+ + M +	+ + + X	+ + + X	+ M + X	+	+ + M +	+ M +	+ + + +	+ + + +	47 2 47 14 46
Pars distalis, adenoma Pars intermedia, adenoma Thyroid gland GENERAL BODY SYSTEM	+	+	X +	X	+	+	+	+	+	+ 	+	X +	+ +	+	+ -	+	+	+ 	* +	+ -	X +	+	* +	+ -	+	15 1 47
None GENITAL SYSTEM Clitoral gland	-		~	-																_						1
Ovary Hemangioma Hemangiosarcoma Hemangiosarcoma, metastatic Histiocytic sarcoma	+	*	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	47 1 1 1
Lymphoma malignant histiocytic Lymphoma malignant lymphocytic Uterus Hemangiosarcoma Histiocytic sarcoma	X +	+	X +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	1 47 1 1
Lymphoma malignant lymphocytic Endometrium, polyp stromal	Х											х										х				1 2

TABLE D2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF FEMALE MICE: 1,200 ppm (Continued)

WEEKS ON STUDY	0 0 1	0 1 2	0 3 3	0 5 8	0 7 0	0 7 0	0 7 0	0 7 5	0 7 5	0 8 5	0 8 5	0 8 6	0 8 9	0 9 6	1 0 0	1 0 2	1 0 3	1 0 3	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5
CARCASS ID	3 5 8 1	3 4 1	3 5 3 1	3 1 7 1	3 5 0 1	3 5 1	3 5 2 1	3 4 7 1	3 2 6 1	3 8 1	3 0 7 1	3 4 8 1	3 3 9	3 2 8 1	3 5 7 1	3 5 4 1	3 4 9 1	3 1 1 1	3 0 4 1	3 0 9	3 1 5 1	3 2 2 1	3 2 7 1	3 7 1	3 4 5 1
HEMATOPOIETIC SYSTEM Bone marrow Hemangiosarcoma, metastatic, ovary Hemangiosarcoma, metastatic, spleen	+	+	+	+				+	+	+	+	+	+	*	+	+	+	+	+	+	+	+	+	+	+
Lymphoma malignant lymphocytic Lymph node Lymphoma malignant undifferentiated cell type Axillary, lymphoma malignant undifferentiated cell type lhac, histiocytic sarcoma linac, lymphoma malignant histiocytic llac, lymphoma malignant mixed Mediastinal, lymphoma malignant histiocytic Mediastinal, lymphoma malignant lymphocytic Mediastinal, lymphoma malignant lymphocytic Mediastinal, lymphoma malignant mixed Mesenteric, lymphoma malignant histiocytic	M	+	+	+				+	+	+	+	+	+	+ X	+	x	+	+	+	+	+	+	+	+	+
Mesentenc, lymphoma malignant lymphocytic Mesentenc, lymphoma malignant mixed Mesentenc, lymphoma malignant undifferentiated cell type Mesentenc, osteosarcoma, metastatic, uncertain primary site Renal, lymphoma malignant histocytic Renal, lymphoma malignant hymphocytic														x	x				x			X			
Renal, lymphoma malignant mixed Lymph node, mandibular Lymphoma malignant histocytic Lymphoma malignant lymphocytic Lymphoma malignant mixed Lymphoma malignant undifferentiated cell type	М	M	+	+				+	+	+	+	М	+	+	+	+	+	+	+	+	+	+ X	M	+	+
Spleen Hemangiosarcoma Hemangiosarcoma, metastatic, uterus Histiocytic sarcoma Lymphoma malignant histiocytic Lymphoma malignant lymphocytic	+	+	+	+				+	+	+	+	*	+	+	+	+ X	+	+	+	+ X	+	+ X	+	+	+
Lymphoma malignant mixed Lymphoma malignant undifferentiated cell type Thymus Lymphoma malignant histocytic Lymphoma malignant lymphocytic Lymphoma malignant mixed Lymphoma malignant undifferentiated cell type	+	+	+	+				+	+	+	+	+	+	X +	+	+	+	+	X	+	+	+ X	+	+	+

TABLE D2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF FEMALE MICE: 1,200 ppm (Continued)

								`-				-/														
WEEKS ON STUDY	1 0	1 0	1 0	1 0	1	1 0	1	1	1	1 0	0	0	1	1 0	1	1 0	1	1	1 0	1	1	1 0	0	1	1 0	
	5	5	5	5	5	5	5	5	5	5	5	5	5	5	5	5	5	5	5	5	5	5	5	5	5	TOTAL
CARCASS	3	3	3	3	3	3	3	3	3	3	3	3	3	3	3	3	4	5	3	3	3 2	3	3	3	-3 6	TISSUES
ID	2	6	4	5	5	6	2	3	6	ó	ī	3	4	9	ī	3	4	ğ	8	ō	0	3	ō	6	0	101120112
	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	
HEMATOPOIETIC SYSTEM																										47
Bone marrow Hemangiosarcoma, metastatic, ovary	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	47
Hemangiosarcoma, metastatic, spleen	ł																								X	i
Lymphoma malignant lymphocytic	X																									1 1
Lymph node Lymphoma malignant undifferentiated	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	46
cell type	1																									1
Axillary, lymphoma malignant undifferentiated cell type	ł																х									1
Iliac, histiocytic sarcoma																										i
Iliac, lymphoma malignant histiocytic	ŀ		Х																							1
Iliac, lymphoma malignant mixed Mediastinal, lymphoma malignant					X													X						X		3
histiocytic	{		X																							1
Mediastinal, lymphoma malignant lymphocytic	x																									1
Mediastinal, lymphoma mal mixed	^																							х		1 1
Mesenteric, lymphoma malignant																										1
histiocytic Mesenteric, lymphoma malignant			X																							1
lymphocytic	x																									2
Mesenteric, lymphoma malignant mixed				X	X							X						X						X		6
Mesenteric, lymphoma malignant undifferentiated cell type																	x									2
Mesenteric, osteosarcoma, metastatic,	-																Λ									
uncertain primary site	1																									1
Renal, lymphoma malignant histocytic Renal, lymphoma mal lymphocytic	X		X																							1
Renal, lymphoma mai lymphocytic Renal, lymphoma malignant mixed	^																							Х		1 1
Lymph node, mandibular	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	43
Lymphoma malignant histocytic	X		X																							$\frac{1}{2}$
Lymphoma malignant lymphocytic Lymphoma malignant mixed	Α.			x	X													х						X		4
Lymphoma malignant undifferentiated																										
cell type Spleen	١.																X									47
Hemangiosarcoma	-					•			•	7			_					_	_			7	-	т.	*	2
Hemangiosarcoma, metastatic, uterus																										1
Histocytic sarcoma	ļ		x																							1
Lymphoma malignant histiocytic Lymphoma malignant lymphocytic	x		А																							1 2
Lymphoma malignant mixed	"			Х	X							X						X						X		- 6
Lymphoma malignant undifferentiated cell type																	v									2
Thymus	1	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	X	+	+	+	+	4	+	+	+	47
Lymphoma malignant histiocytic	'	•	X	•				*	•									,	•		•	,		•		1
Lymphoma malignant lymphocytic	X				**																					2
Lymphoma malignant mixed Lymphoma malignant undifferentiated					X																					1
cell type																	X									1
	l																									l

TABLE D2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF FEMALE MICE: 1,200 ppm (Continued)

					``			uec	• /																
WEEKS ON STUDY	0 0 1	0 1 2	0 3 3	0 5 8	0 7 0	0 7 0	0 7 0	0 7 5	0 7 5	0 8 5	0 8 5	0 8 6	0 8 9	0 9 6	1 0 0	1 0 2	1 0 3	1 0 3	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5
CARCASS ID	3 5 8 1	3 3 4 1	3 5 3 1	3 1 7 1	3 5 0	3 5 1	3 5 2 1	3 4 7 1	3 2 6 1	3 8 1	3 0 7 1	3 4 8 1	3 9 1	3 2 8 1	3 5 7 1	3 5 4 1	3 4 9 1	3 1 1 1	3 0 4 1	3 0 9	3 1 5 1	3 2 2 1	3 2 7 1	3 7 1	3 4 5 1
INTEGUMENTARY SYSTEM Mammary gland Carrinoma Skin	+	+	+	+				+	+	+	+	+ +	+	+	+	+	+	+	+	+	M +	+	+ +	+	+ +
Papilloma squamous Subcutaneous tissue, fibrosarcoma Subcutaneous tissue, hemangioma Subcutaneous tissue, hemangiosarcoma															x		x								
MUSCULOSKELETAL SYSTEM Bone		+	+	+				+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
NERVOUS SYSTEM Brain Lymphoma malignant lymphocytic	+	+	+	+	_	-		+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+
RESPIRATORY SYSTEM Lung Alveolar/bronchiolar adenoma Alveolar/bronchiolar carcinoma Carcinoma, metastatic	+	+	+	+				+	+ X	+	+ X	*	+	+	+	+	+	+	+	+	+	+ X	+	+	+
Hepatocellular carcinoma, metastatic, liver Histiocytic sarcoma Lymphoma malignant histiocytic Lymphoma malignant lymphocytic Lymphoma malignant mixed										x						x						x			
Osteosarcoma, metastatic, uncertain primary site Nose Trachea	+	++	++	++				+	+	+	++	++	++	++	X + +	++	+	++	++	+	++	++	++	++	+++
SPECIAL SENSES SYSTEM Hardenan gland Adenoma Carcinoma																-	-	+	*			+ X			
URINARY SYSTEM Kidney Histrocytic sarcoma Lymphoma malignant histrocytic Lymphoma malignant lymphocytic	+	+	+	+				+	+	+	+	+	+	+	+	*	+	+	+	+	+	+ X	+	+	+
Lymphoma malignant mixed Lymphoma malignant undifferentiated cell type Osteosartoma, metastatic, uncertain primary site Urinary bladder Lymphoma malignant histocytic Lymphoma malignant lymphocytic Lymphoma malignant mixed	+	+	+	+				+	+	+	+	+	+	x	X +	+	+	+	+	+	+	+	+	+	+

TABLE D2. INDIVIDUAL ANIMAL TUMOR PATHOLOGY OF FEMALE MICE: 1,200 ppm (Continued)

WEEKS ON STUDY	1 0 5	1 0 5	1 0 5	0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	1 0 5	TOTAL.
CARCASS ID	3 0 2 1	3 0 6 1	3 2 4 1	3 2 5 1	3 5 1	3 1 6 1	3 3 2 1	3 3 1	3 6 1	3 4 0 1	3 4 1 1	3 0 3 1	3 1 4 1	3 1 9	3 1 1	3 4 3 1	3 4 4 1	3 5 9	3 0 8 1	3 1 0 1	3 2 0 1	3 2 3 1	3 0 1	3 4 6 1	3 6 0 1	TISSUES TUMORS
INTEGUMENTARY SYSTEM Mammary gland Carcinoma Skin	+ +	+	+	+	+	+	+ X +	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	46 1 47
Papilloma squamous Subcutaneous tissue, fibrosarcoma Subcutaneous tissue, hemangioma Subcutaneous tissue, hemangiosarcoma											Х						х									1 1 1 1
MUSCULOSKELETAL SYSTEM Bone	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	47
NERVOUS SYSTEM Brain Lymphoma malignant lymphocytic	+ X	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	47
RESPIRATORY SYSTEM Lung Alveolar/bronchiolar adenoma Alveolar/bronchiolar carcinoma Carcinoma, metastatic Hepatocellular carcinoma, metastatic,	+	+	+	+	+	+	+ X	*	+	+	+	+	+	+	+	+	*	+	+	+	+	+ X	+	+	+	47 4 3 1
inver Histocytic sarcoma Lymphoma malignant histocytic Lymphoma malignant lymphocytic Lymphoma malignant mixed Osteosarcoma, metastatic, uncertain	x	х	x	x																						2 1 1 2 1
primary site Nose Trachea	+++	+	++	+ +	++	+	++	+ +	+	+ +	++	<i>+</i> +	+ +	+ +	+ +	+ +	+	+	+	+	++	++	+	+	+ +	47 47
SPECIAL SENSES SYSTEM Hardenan gland Adenoma Carcinoma												-							=							3 1 1
URINARY SYSTEM Kidney Histocytic sarcoma Lymphoma malignant histocytic Lymphoma malignant lymphocytic Lymphoma malignant mixed	+	+	+ X	+ X	+ x	+	+	+	+	+	+	+	+	+	+	+	+	+ X	+	+	+	+	+	+	+	47 1 1 1 3
Lymphoma malignant undifferentiated cell type Osteosarcoma, metastatic, uncertain primary site Unnary bladder Lymphoma malignant histiocytic Lymphoma malignant lymphocytic Lymphoma malignant mixed	+ x	+	* X	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+ X	+	+	+	+	+	+	+	1 47 1 1

TABLE D3. ANALYSIS OF PRIMARY TUMORS IN FEMALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE

	Chamber Control	120 ppm	600 ppm	1,200 ppm
Liver: Hepatocellular Adenoma				
Overall Rates (a)	3/49 (6%)	7/50 (14%)	6/50 (12%)	7/47 (15%)
Adjusted Rates (b)	10.0%	19.7%	22.5%	20.0%
Terminal Rates (c)	3/30 (10%)	5/33 (15%)	5/24 (21%)	5/32 (16%)
Day of First Observation	729	683	509	520
Life Table Tests (d)	P = 0.235	P = 0.190	P = 0.153	P = 0.181
Logistic Regression Tests (d)	P = 0.208	P = 0.147	P = 0.225	P = 0.136
Cochran-Armitage Trend Test (d)	P = 0.210			
Fisher Exact Test (d)		P = 0.167	P = 0.254	P = 0.142
Liver: Hepatocellular Carcinoma				
Overall Rates (a)	4/49 (8%)	2/50 (4%)	2/50 (4%)	7/47 (15%)
Adjusted Rates (b)	11.1%	5.0%	6.3%	19.4%
Terminal Rates (c)	2/30 (7%)	0/33 (0%)	1/24 (4%)	4/32 (13%)
Day of First Observation	631	477	578	589
Life Table Tests (d)	P = 0.090	P = 0.337N	P = 0.407N	P = 0.264
Logistic Regression Tests (d)	P = 0.078	P = 0.323N	P = 0.328N	P = 0.232
Cochran-Armitage Trend Test (d)	P = 0.078			
Fisher Exact Test (d)		P = 0.329N	P = 0.329N	P = 0.238
Liver: Hepatocellular Adenoma or Carc	inoma			
Overall Rates (a)	7/49 (14%)	9/50 (18%)	8/50 (16%)	13/47 (28%)
Adjusted Rates (b)	20.6%	23.7%	28.3%	34.5%
Terminal Rates (c)	5/30 (17%)	5/33 (15%)	6/24 (25%)	8/32 (25%)
Day of First Observation	631	477	509	520
Life Table Tests (d)	P = 0.093	P = 0.435	P = 0.358	P = 0.124
Logistic Regression Tests (d)	P = 0.070	P = 0.390	P = 0.503	P = 0.081
Cochran-Armitage Trend Test (d) Fisher Exact Test (d)	P = 0.072	P = 0.410	P = 0.517	P = 0.086
risher Exact Test (d)		F = 0.410	F = 0.517	F = 0.000
Lung: Alveolar/Bronchiolar Adenoma				
Overall Rates (a)	5/50 (10%)	0/50 (0%)	3/50 (6%)	4/47 (9%)
Adjusted Rates (b)	14.8%	0.0%	10.3%	11.7%
Terminal Rates (c)	3/30 (10%)	0/33 (0%)	1/24 (4%)	3/32 (9%)
Day of First Observation	681	5 000437	702	598
Life Table Tests (d)	P = 0.354	P = 0.031N	P = 0.477N	P = 0.489N
Logistic Regression Tests (d)	P=0.335	P = 0.036N	P = 0.409N	P = 0.545N
Cochran-Armitage Trend Test (d)	P = 0.334	D 0.000M	D 0.05531	D 0 7 4037
Fisher Exact Test (d)		P = 0.028N	P = 0.357N	P = 0.540N
Lung: Alveolar/Bronchiolar Carcinoma	0.00.000	0/50/0~>	1/50 /0~ :	0.44 .0 ~ .
Overall Rates (a) Adjusted Rates (b)	0/50 (0%)	3/50 (6%)	1/50 (2%)	3/47 (6%)
•	0.0%	8.6%	4.2% 1/24 (4%)	7.8%
Terminal Rates (c)	0/30 (0%)	2/33 (6%)		1/32 (3%)
Day of First Observation	B-0.999	686 B = 0.134	729 D = 0.455	523 B=0.118
Life Table Tests (d)	P=0.228	P=0.134 P=0.114	P = 0.455	P = 0.118
Logistic Regression Tests (d) Cochran-Armitage Trend Test (d)	P = 0.210	P = 0.114	P = 0.455	P = 0.114
Fisher Exact Test (d)	P = 0.211	P = 0.121	P = 0.500	P = 0.110
Lung: Alveolar/Bronchiolar Adenoma or	Carcinoma			
Overall Rates (a)	5/50 (10%)	3/50 (6%)	4/50 (8%)	7/47 (15%)
Adjusted Rates (b)	14.8%	8.6%	14.2%	18.9%
Terminal Rates (c)	3/30 (10%)	2/33 (6%)	2/24 (8%)	4/32 (13%)
Day of First Observation	681	686	702	523
	P = 0.182	P = 0.331N	P = 0.627N	P=0.386
Life Table Tests (d)				
Life Table Tests (d) Logistic Regression Tests (d)				
Life Table Tests (d) Logistic Regression Tests (d) Cochran-Armitage Trend Test (d)	P=0.156 P=0.157	P = 0.375N	P = 0.567N	P = 0.330

TABLE D3. ANALYSIS OF PRIMARY TUMORS IN FEMALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

	Chamber Control	120 ppm	600 ppm	1,200 ppm
Pituitary Gland/Pars Distalis: Adenom	а			
Overall Rates (a)	12/49 (24%)	19/48 (40%)	21/49 (43%)	15/46 (33%)
Adjusted Rates (b)	36.9%	51.0%	58.0%	46.9%
Terminal Rates (c)	10/30 (33%)	15/33 (45%)	10/24 (42%)	15/32 (47%)
Day of First Observation	653	653	398	729
Life Table Tests (d)	P=0.410	P=0.146	P=0.014	P=0.382
Logistic Regression Tests (d)	P=0.344	P=0.066	P=0.033	P = 0.278
Cochran-Armitage Trend Test (d)	P=0.367	1 -0.000	1 -0.000	1 -0.270
Fisher Exact Test (d)	1 = 0.301	P = 0.084	P = 0.043	P = 0.258
Forestomach: Squamous Papilloma				
Overall Rates (e)	3/50 (6%)	1/50 (2%)	1/50 (2%)	3/47 (6%)
Adjusted Rates (b)	10.0%	3.0%	4.2%	9.4%
Terminal Rates (c)	3/30 (10%)	1/33 (3%)	1/24 (4%)	3/32 (9%)
Day of First Observation	729	729	729	729
Life Table Tests (d)	P = 0.439	P = 0.271N	P = 0.387N	P = 0.634N
Logistic Regression Tests (d)	P = 0.439	P = 0.271N	P = 0.387N	P = 0.634N
Cochran-Armitage Trend Test (d)	P = 0.418			
Fisher Exact Test (d)		P = 0.309N	P = 0.309N	P = 0.631
Thyroid Gland: Follicular Cell Adenor	na			
Overall Rates (a)	0/50 (0%)	0/50 (0%)	4/50 (8%)	0/47 (0%)
Adjusted Rates (b)	0.0%	0.0%	12.0%	0.0%
Terminal Rates (c)	0/30 (0%)	0/33 (0%)	1/24 (4%)	0/32 (0%)
Day of First Observation	0,00 (0,0)	0.00 (0.0)	629	
Life Table Tests (d)	P = 0.405	(f)	P = 0.050	(f)
Logistic Regression Tests (d)	P=0.404	(f)	P = 0.063	(f)
Cochran-Armitage Trend Test (d)	P = 0.404	\- /	1 0.000	1-/
Fisher Exact Test (d)	x = 0.101	(f)	P = 0.059	(f)
Thyroid Gland: Follicular Cell Adenon	na or Carcinoma			
Overall Rates (a)	1/50 (2%)	0/50 (0%)	4/50 (8%)	0/47 (0%)
Adjusted Rates (b)	3.3%	0.0%	12.0%	0.0%
Terminal Rates (c)	1/30 (3%)	0/33 (0%)	1/24 (4%)	0/32 (0%)
Day of First Observation	729		629	
Life Table Tests (d)	P = 0.592	P = 0.481 N	P = 0.140	P = 0.487N
Logistic Regression Tests (d)	P = 0.593	P = 0.481N	P = 0.178	P = 0.487N
Cochran-Armitage Trend Test (d)	P = 0.594			
Fisher Exact Test (d)		P = 0.500N	P = 0.181	P = 0.515N
Uterus: Stromal Polyp				
Overall Rates (e)	3/50 (6%)	2/50 (4%)	1/50 (2%)	2/47 (4%)
Adjusted Rates (b)	10.0%	5.9%	3.8%	6.3%
Terminal Rates (c)	3/30 (10%)	1/33 (3%)	0/24 (0%)	2/32 (6%)
Day of First Observation	729	722	713	729
Life Table Tests (d)	P = 0.422N	P = 0.457N	P = 0.393N	P = 0.470N
Logistic Regression Tests (d)	P = 0.429N	P = 0.490N	P = 0.369N	P = 0.470N
Cochran-Armitage Trend Test (d) Fisher Exact Test (d)	P = 0.435N	P=0.500N	P = 0.309N	P = 0.530N
Circulatory System: Hemangiosarcoma		0/50 (46)	1/50 (97%)	C/AT (100)
Overall Rates (e)	2/50 (4%)	2/50 (4%)	1/50 (2%)	6/47 (13%)
Adjusted Rates (b)	5.8%	6.1%	3.3%	17.0%
Terminal Rates (c)	1/30 (3%)	2/33 (6%)	0/24 (0%)	4/32 (13%)
Day of First Observation	681	729	708	598
Life Table Tests (d)	P = 0.054	P = 0.670N	P = 0.569N	P = 0.142
Logistic Regression Tests (d)	P = 0.044	P = 0.684	P = 0.519N	P = 0.112
Cochran-Armitage Trend Test (d)	P = 0.043		n	
Fisher Exact Test (d)		P = 0.691N	P = 0.500N	P = 0.115

TABLE D3. ANALYSIS OF PRIMARY TUMORS IN FEMALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

	Chamber Control	120 ppm	600 ppm	1,200 ppm
Circulatory System: Hemangioma or	Hemangiosarcoma		ale Affres and a second about	
Overall Rates (e)	4/50 (8%)	3/50 (6%)	1/50 (2%)	8/47 (17%)
Adjusted Rates (b)	12.3%	9.1%	3.3%	22.2%
Terminal Rates (c)	3/30 (10%)	3/33 (9%)	0/24 (0%)	5/32 (16%)
Day of First Observation	681	729	708	598
Life Table Tests (d)	P = 0.078	P = 0.457N	$P = 0.251 \mathrm{N}$	P = 0.194
Logistic Regression Tests (d)	P = 0.063	P = 0.514N	P = 0.208N	P = 0.143
Cochran-Armitage Trend Test (d)	P = 0.062			
Fisher Exact Test (d)		P = 0.500N	P = 0.181N	P = 0.149
Hematopoietic System: Lymphoma, A	ll Malignant			
Overall Rates (e)	22/50 (44%)	10/50 (20%)	17/50 (34%)	11/47 (23%)
Adjusted Rates (b)	60.5%	26.6%	45.9%	33.1%
Terminal Rates (c)	16/30 (53%)	6/33 (18%)	6/24 (25%)	10/32 (31%
Day of First Observation	624	599	398	670
Life Table Tests (d)	P = 0.147N	P = 0.007N	P = 0.473N	P = 0.012N
Logistic Regression Tests (d)	P = 0.138N	P = 0.010N	P = 0.227N	P = 0.022N
Cochran-Armitage Trend Test (d)	P = 0.142N			
Fisher Exact Test (d)		P = 0.009N	P = 0.206N	P = 0.027N

⁽a) Number of tumor-bearing animals/number of animals examined microscopically at the site

⁽b) Kaplan-Meier estimated tumor incidences at the end of the study after adjusting for intercurrent mortality

⁽c) Observed tumor incidence in animals killed at the end of the study

⁽d) Beneath the control incidence are the P values associated with the trend test. Beneath the dosed group incidence are the P values corresponding to pairwise comparisons between that dosed group and the controls. The life table analysis regards tumors in animals dying prior to terminal kill as being (directly or indirectly) the cause of death. The logistic regression test regards these lesions as nonfatal. The Cochran-Armitage and Fisher exact tests compare directly the overall incidence rates. A negative trend or lower incidence in a dosed group than in controls is indicated by (N).

⁽e) Number of tumor-bearing animals/number of animals examined grossly at the site

⁽f) No P value is reported because no tumors were observed in the exposed and control groups.

TABLE D4a. HISTORICAL INCIDENCE OF ANTERIOR PITUITARY GLAND TUMORS IN FEMALE B6C3F₁ MICE RECEIVING NO TREATMENT (a)

		Incidence in Co	ntrols
Study	Adenoma	Carcinoma	Adenoma or Carcinoma
Historical Incidence for Ch	namber Controls at Battelle P	acific Northwest La	boratories
Propylene oxide	8/46	1/46	9/46
Methyl methacrylate	12/49	0/49	12/49
Propylene	(b) 13/41	0/41	(b) 13/41
1,2-Epoxybutane	19/47	3/47	22/47
Dichloromethane	4/46	0/46	4/46
Ethylene oxide	4/48	1/48	5/48
Bromoethane	2/48	0/48	2/48
Fetrachloroethylene	2/45	5/45	7/45
TOTAL	(b) 64/370 (17.3%)	10/370 (2.7%)	(b) 74/370 (20.0%)
SD(c)	13.55%	4.04%	13.97%
Range (d)			
High	19/47	5/45	22/47
Low	2/48	0/49	2/48
Overall Historical Incidenc	e for Untreated Controls in I	NTP Studies	
TOTAL	(e) 244/1,528 (16.0%)	(f) 12/1,528 (0.8%)	(e,f) 256/1,528 (16.8%)
SD(c)	10.80%	1.42%	11.09%
Range (d)			
High	18/49	3/50	19/49
Low	0/48	0/50	0/48

⁽a) Data as of May 12, 1988, for studies of at least 104 weeks (b) Includes 11 chromophobe adenomas

⁽c) Standard deviation
(d) Range and SD are presented for groups of 35 or more animals.

⁽e) Includes four chromophobe adenomas (f) Includes three adenocarcinomas, NOS

TABLE D4b. HISTORICAL INCIDENCE OF INTERMEDIA PITUITARY GLAND TUMORS IN FEMALE B6C3F₁ MICE RECEIVING NO TREATMENT (a)

Study	Incidence of Adenomas in Controls	
Historical Incidence for Chamber (Controls at Battelle Pacific Northwest Laboratories	
Propylene oxide	0/46	
Methyl methacrylate	1/49	
Propylene	0/41	
1,2-Epoxybutane	0/47	
Dichloromethane	0/46	
Ethylene oxide	0/48	
Bromoethane	0/48	
Te trachlo roethylene	0/45	
TOTAL	1/370 (0.3%)	
SD(b)	0.72%	
Range (c)		
High	1/49	
Low	0/48	
Overall Historical Incidence for Un	ntreated Controls in NTP Studies	
TOTAL	3/1,528 (0.2%)	
SD(b)	0.64%	
Range (c)		
High	1/43	
Low	0/50	

⁽a) Data as of May 12, 1988, for studies of at least 104 weeks; no malignant tumors have been observed.

⁽b) Standard deviation
(c) Range and SD are presented for groups of 35 or more animals.

TABLE D4c. HISTORICAL INCIDENCE OF HEMATOPOIETIC SYSTEM TUMORS IN FEMALE B6C3F1 MICE RECEIVING NO TREATMENT (a)

	Incidence in Controls				
Study	Lymphoma	Lymphoma or Leukemia			
istorical Incidence for Chambe	r Controls at Battelle Pacific North	west Laboratories			
copylene oxide	12/50	12/50			
ethyl methacrylate	8/50	8/50			
copylene	16/50	16/50			
-Epoxybutane	13/50	13/50			
ichloromethane	7/50	7/50			
hylene oxide	9/49	9/49			
romoethane	11/50	11/50			
trachloroethylene	8/49	8/49			
TOTAL	84/398 (21.1%)	84/398 (21.1%)			
SD(b)	6.08%	6.08%			
nge (c)					
High	16/50	16/50			
Low	7/50	7/50			
verall Historical Incidence for	Untreated Controls in NTP Studies				
TOTAL	523/1,689 (31.0%)	537/1,689 (31.8%)			
SD(b)	12.73%	12.20%			
inge (c)					
High	37/50	(d) 38/50			
Low	5/50	6/50			

⁽a) Data as of May 12, 1988, for studies of at least 104 weeks (b) Standard deviation $\,$

⁽c) Range and SD are presented for groups of 35 or more animals.
(d) Second highest: 29/50

TABLE D5. SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN FEMALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE

	Chamber (Control	120	ppm	600	ppm	1,200 p	pm
Animals initially in study	60		60	***	60		60	
Animals removed	60		60		60		60	
Animals examined histopathologically	50		50		50		47	
ALIMENTARY SYSTEM						-		
Esophagus	(49)		(50)		(50)		(47)	
Infiltration cellular, lymphocytic			2	(4%)				
Inflammation, acute					1	(2%)		
Gallbladder	(49)		(49)		(50)		(45)	
Hyperplasia, lymphoid		(4%)						
Infiltration cellular, lymphocytic		(29%)		(37%)		(30%)		(36%)
Inflammation, acute	1	(2%)	1	(2%)		(2%)	1	(2%)
Inflammation, chronic						(2%)	_	
Inflammation, chronic active						(2%)		(2%)
Intestine large	(50)		(50)		(50)		(47)	
Anorectal junction, inflammation, acute		(2%)				/0 <i>~</i> \		
Anorectal junction, inflammation, chronic	active				1	(2%)		,o~ .
Anorectal junction, ulcer								(2%)
Anus, erosion Anus, inflammation, acute						(00)		(2%)
					1	(2%)		(2%)
Anus, inflammation, chronic active Anus, ulcer					,	(90()	1	(2%)
Intestine large, cecum	(49)		(50)		(50)	(2%)	(47)	
Inflammation, acute		(2%)	(50)		(50)		(47)	
Intestine large, rectum	(50)	(270)	(50)		(50)		(45)	
Inflammation, acute		(2%)	(00)			(2%)		(2%)
Intestine small, duodenum	(49)	(= ,0 ,	(50)		(50)	(270)	(47)	(= ,0,
Inflammation, chronic active	(/		(00)			(2%)	(,	
Intestine small, ileum	(49)		(50)		(50)	,	(47)	
Amyloid deposition	1	(2%)	2	(4%)	2	(4%)		(2%)
Intestine small, jejunum	(49)		(50)		(50)		(47)	
Diverticulum							1	(2%)
Necrosis							1	(2%)
Liver	(49)		(50)		(50)		(47)	
Basophilic focus	1	(2%)	3	(6%)				
Bile stasis							1	(2%)
Congestion							1	(2%)
Cyst			1	(2%)				
Fatty change		(2%)			_	(2%)		(6%)
Hematopoietic cell proliferation	23	(47%)		(32%)	30	(60%)	26	(55%)
Hyperplasia				(2%)				
Hyperplasia, lymphoid	2	,	1	(2%)				
Infarct		(2%)		(00% \		.00~)	0.1	
Infiltration cellular, lymphocytic		(47%)	16	(32%)	15	(30%)		(45%)
Inflammation, acute Inflammation, chronic	1	(2%)		(00)			1	(2%)
Inflammation, chronic active			1	(2%)				(900)
Necrosis	~	(14%)	4	(80%)	4	(9.0%)		(2%)
Thrombus		(14%) $(2%)$	4	(8%)	4	(8%)	4	(9%)
Vacuolization cytoplasmic	1	(270)			1	(2%)	1	(2%)
Hepatocyte, hypertrophy	1	(2%)			1	(470)	1	(270)
Median lobe, angiectasis	1	(270)	1	(2%)				
Serosa, inflammation, chronic			1	(270)	1	(2%)	,	(2%)
Mesentery	(5)		(2)		(4)	(470)	(2)	(470)
Inflammation, chronic	(3)		(4)			(25%)	(2)	
Fat, necrosis	1				•	(20/0/		

TABLE D5. SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN FEMALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

	Chamber Control		120	120 ppm		600 ppm		pm
ALIMENTARY SYSTEM (Continued)						-		
Pancreas	(50))	(50)		(50)		(47)	
Hyperplasia, lymphoid		(8%)		(2%)	(00)		(=1)	
Infiltration cellular, lymphocytic		(34%)		(56%)	25	(50%)	26	(55%)
Inflammation, acute		(0 1.0)		(2%)		(2%)		(00,0)
Inflammation, chronic active	1	(2%)		` '		,	1	(2%)
Acinus, atrophy	3	(6%)	1	(2%)	2	(4%)	2	(4%)
Duct, cyst	1	(2%)			1	(2%)		
Salivary glands	(50)	1	(50)		(50)		(46)	
Atrophy			1	(2%)				
Hyperplasia, lymphoid	4	(8%)						
Infiltration cellular, lymphocytic	28	(56%)	37	(74%)	33	(66%)	32	(70%)
Stomach	(50)	İ	(50)		(50)		(47)	
Hyperplasia, squamous, focal					1	(2%)		
Stomach, forestomach	(50)		(49)		(50)		(47)	
Angiectasis								(2%)
Erosion		(2%)	1	(2%)	2	(4%)	2	(4%)
Hyperkeratosis	_	(4%)	_	(40~:	_	(O. o		
Hyperplasia, squamous		(4%)		(12%)		(8%)		(9%)
Hyperplasia, squamous, focal		(4%)		(6%)		(4%)	_	(4%)
Infiltration cellular, lymphocytic		(10%)	-	(12%)		(16%)		(11%)
Inflammation, acute	2	(4%)	2	(4%)	6	(12%)		(15%)
Inflammation, chronic				.o~ .	•			(2%)
Inflammation, chronic active		(04)	4	, ,		(6%)	3	(6%)
Ulcer		(2%)	_	(2%)		(8%)		(9%)
Stomach, glandular	(49)		(50)		(50)	.a~\	(47)	
Edema	0	(400)		(OW)		(2%)	•	
Erosion		(4%)	4	(8%)	4	(8%)	2	(4%)
Hyperplasia, lymphoid		(4%)	10	(000)		(440)	00	(400)
Infiltration cellular, lymphocytic Inflammation, acute		(31%) (8%)		(36%) (4%)	_	(44%) (4%)		(49%)
Inflammation, chronic	4	(070)	Z	(4270)		(4 %) (2%)	Z	(4%)
Inflammation, chronic active						(2%)		
Metaplasia, squamous			1	(2%)	-	(270)		
Mineralization				(2%)	9	(4%)	9	(4%)
Ulcer	9	(4%)	•	(2 10)		(4%)		(4/0)
Mucosa, dilatation		(20%)	6	(12%)		(20%)	14	(30%)
Tooth	10	(20%)	(1)	(12,0)		(20 %)	• •	(00 /0)
Pulp, inflammation, acute				(100%)				
ARDIOVASCULAR SYSTEM		-	· · · · · · · · · · · · · · · · · · ·	*******				
Blood vessel	(50)		(50)		(50)		(47)	
Aorta, mineralization				(2%)				
Heart	(50)		(50)		(50)		(47)	
Fibrosis, focal				(2%)				
Infiltration cellular, lymphocytic	10	(20%)	13	(26%)		(40%)	18	(38%)
Inflammation, acute	-					(6%)		
Inflammation, chronic		(4%)	1	(2%)	4	(8%)	1	(2%)
Inflammation, chronic active		(4%)		/O# :				.o~ :
Mineralization	1	(2%)	1	(2%)		/O.W.:	1	(2%)
Atrium, thrombus				/O# :	1	(2%)		
Coronary artery, inflammation, acute				(2%)		(OA)		
Coronary artery, inflammation, chronic Valve, thrombus	1	(2%)	1	(2%)	1	(2%)		
NDOCDINE CYCTEN			 -				<u>, </u>	
NDOCRINE SYSTEM Adrenal gland	(40)		(FA)		(EO)		/ 4171	
Capsule, inflammation, acute	(49)		(50)		(50)	(90%)	(47)	
Capsule, inflammation, acute Capsule, spindle cell, hyperplasia	40	(100%)	EΛ	(100%)		(2%)	AF	(0604)
Capacie, spindle cell, hyperpiasia	49	(100%)	οU	(100%)	49	(98%)	40	(96%)

TABLE D5. SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN FEMALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

	Chamber Control		120	ppm	600 ppm		1,200 ppm	
ENDOCRINE SYSTEM (Continued)								
Adrenal gland, cortex	(49)		(50)		(50)		(47)	
Congestion	(40)		(00)		(00)			(2%)
Cyst	1	(2%)						(2%)
Degeneration, fatty		(2%)	9	(18%)	2	(4%)		(2%)
Hematopoietic cell proliferation	•	(270)	•	(10,0)	_	(470)		(2%)
Hyperplasia	1	(2%)					•	(2,70)
Hypertrophy		(2%)	2	(4%)	2	(4%)	4	(9%)
Hypertrophy, diffuse	•	(270)	-	(470)	-	(470)		(2%)
Necrosis					1	(2%)	•	(2,0)
Pigmentation	35	(71%)	37	(74%)		(90%)	37	(79%)
Thrombus		(2%)	•	((,		(2%)	•	(.0,0)
Spindle cell, hyperplasia	-	(= 10)			-	(= ,0 ,	1	(2%)
Adrenal gland, medulla	(49)		(50)		(49)		(47)	(= ,0)
Hyperplasia		(2%)	(30)			(2%)	(/	
Necrosis	-	(= ,,,				(2%)		
Islets, pancreatic	(50)		(49)		(50)	.= ,0 ,	(47)	
Hyperplasia		(2%)		(2%)		(2%)	(*1)	
Pituitary gland	(49)		(48)		(49)	,	(46)	
Pars distalis, angiectasis		(4%)	(10)			(4%)	(10)	
Pars distalis, cyst	-	(4,0)	3	(6%)		(6%)		
Pars distalis, hemorrhage			·	(0,0)		(2%)		
Pars distalis, hyperplasia	12	(24%)	18	(38%)	_	(29%)	15	(33%)
Thyroid gland	(50)		(50)	(00,0)	(50)	(40 ,0 ,	(47)	(00 /0 /
Hyperplasia, lymphoid		(2%)	(00)		(00)		(=1)	
Infiltration cellular, lymphocytic		(10%)	6	(12%)	8	(16%)	6	(13%)
Inflammation, acute	·	(10,0)	-	(2%)	-	(2%)	-	(2%)
Inflammation, chronic			•	(2 /0 /		(2%)	•	(2/0)
Inflammation, chronic active			1	(2%)	•	(2,70)	3	(6%)
C-cell, hyperplasia			•	(2 10)	1	(2%)		(0,0)
Follicle, dilatation	2	(16%)	5	(10%)		(8%)	ξ.	(11%)
Follicle, hyperplasia		(18%)		(26%)		(14%)		(21%)
1 officie, ny per piasia	3	(10 %)	10	(20%)		(14/0)		(21%)
GENERAL BODY SYSTEM None								
GENITAL SYSTEM								
Clitoral gland	(1)	(100%)	(1)		(1)		(1)	
Abscess	1	(100%)						
Cyst	(20)		(40)			(100%)		
Ovary	(50)		(49)		(50)	(40)	(47)	
Abscess						(4%)		
Hemorrhage	•	(400)			1	(2%)	2	(4%)
Hyperplasia, lymphoid	2	(4%)						(96)
Hyperplasia, adenomatous						(90)	1	(2%)
Infiltration cellular, lymphocytic						(2%)		(00)
Inflammation, acute Mineralization					2	(4%)		(9%)
Pigmentation, cholesterol, hemosiderin					1	(20c)	1	(2%)
						(2%)		
Follicle, cyst Follicle, cyst multilocular						(2%)		
Periovarian tissue, cyst	11	(22%)	o	(16%)		(2%)	-	(150)
Oviduct	11	(4470)	•	(1070)		(16%)	- 1	(15%)
Inflammation, acute					(1)	(1000)		
Uterus	(50)		(50)			(100%)	(47)	
		(2%)	(50)	(4%)	(50)		(47)	(2%)
Angiectasis Endometrium, hyperplasia, cystic					20	(760)		
Endometrium, hyperplasia, cystic Endometrium, inflammation, acute		(90%) (8%)		(86%) (2%)		(76%) (14%)		(81%) $(13%)$

TABLE D5. SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN FEMALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

	Chamber Control 120 p		ppm	600 ppm		1,200 ppm		
IEMATOPOIETIC SYSTEM				-				
Bone marrow	(50)		(50)		(50)		(47)	
Angiectasis	1	(2%)			•			
Fibrosis	25	(50%)	32	(64%)	26	(52%)	21	(45%)
Fibrous osteodystrophy				,		(/		(9%)
Myeloid cell, hyperplasia	33	(66%)	35	(70%)	38	(76%)	34	(72%)
Lymph node	(48)	, ,	(49)		(50)	, - ,	(46)	
Hemorrhage	1	(2%)						
Hyperplasia, lymphoid	1	(2%)	2	(4%)	2	(4%)		
Inflammation, acute	1	(2%)						
Axillary, hyperplasia, lymphoid							1	(2%)
Bronchial, hyperplasia, lymphoid					1	(2%)		
Iliac, hyperplasia, lymphoid	3	(6%)					2	(4%)
Iliac, inflammation, acute							1	(2%)
Mediastinal, congestion							2	(4%)
Mediastinal, hyperplasia, lymphoid	1	(2%)	1	(2%)	2	(4%)		(11%)
Mediastinal, inflammation, acute		•		•	1	(2%)		
Mesenteric, angiectasis	1	(2%)					1	(2%)
Mesenteric, congestion			5	(10%)	6	(12%)		(15%)
Mesenteric, hemorrhage					,	-		(2%)
Mesenteric, hyperplasia, lymphoid	26	(54%)	17	(35%)	21	(42%)		(52%)
Mesenteric, inflammation, acute		(2%)		(4%)		(14%)		(15%)
Renal, congestion	_	• •	_			(2%)	,	,
Renal, hyperplasia, lymphoid	2	(4%)				(2%)	1	(2%)
Renal, inflammation, acute	_					(2%)		(2%)
Lymph node, mandibular	(47)		(48)		(50)		(43)	,,,
Congestion	, ,		(-5)		,,			(2%)
Hyperplasia			1	(2%)	1	(2%)	•	,
Hyperplasia, glandular			•	,_ ,- ,- ,		(2%)		
Hyperplasia, lymphoid	31	(66%)	29	(60%)		(64%)	25	(58%)
Inflammation, acute	0.			(4%)	-	(4%)	-	(2%)
Pigmentation				(2%)		(2%)	-	101
Spleen	(50)		(50)		(49)	, /	(47)	
Angiectasis	,,,,,		`/		/			(2%)
Hematopoietic cell proliferation	41	(82%)	39	(78%)	41	(84%)		(96%)
Hyperplasia, lymphoid		(32%)	-	(32%)		(27%)		(30%)
Infiltration cellular			- •			(2%)		2 ,
Inflammation, chronic	1	(2%)				•		
Necrosis	1	(2%)						
Pigmentation	37	(74%)	33	(66%)	34	(69%)	28	(60%)
Capsule, inflammation, acute			_			(2%)	_	
Thymus	(46)		(48)		(48)		(47)	
Congestion			1	(2%)			1	(2%)
Cyst	2	(4%)	1	(2%)	_	(2%)	2	(4%)
Ectopic thyroid					1	(2%)		
Edema						(2%)		
Hyperplasia, lymphoid	7	(15%)	6	(13%)	10	(21%)	10	(21%)
TEGUMENTARY SYSTEM								·····
Mammary gland	(49)		(50)		(48)		(46)	
Ectasia	16	(33%)		(36%)	18	(38%)		(24%)
Inflammation, acute				(2%)				
Acinus, ectasia						(2%)		
Skin	(50)		(50)		(50)		(47)	
Alopecia		(18%)		(18%)	8	(16%)		(4%)
Inflammation, acute	1	(2%)						
Inflammation, chronic					1	(2%)		
Ulcer	1	(2%)						
Back, ulcer			1	(2%)				
Foot, ulcer					1	(2%)		
Head, ulcer		(2%)				(2%)		

TABLE D5. SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN FEMALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

	Chamber	Control	120 j	ppm	600 į	ppm	1,200 p	pm
INTEGUMENTARY SYSTEM								
Skin (Continued)	(50)		(50)		(50)		(47)	
Inguinal, inflammation, acute	1	(2%)						
Inguinal, ulcer						(2%)		
Subcutaneous tissue, edema					1	(2%)	1	(2%)
Subcutaneous tissue, hemorrhage		(2%)	1	(2%)				
Subcutaneous tissue, inflammation, acute					I	(2%)	1	(2%)
Ventral, inflammation, chronic	1	(2%)						
MUSCULOSKELETAL SYSTEM								
Bone	(50)		(49)		(50)		(47)	
Cranium, fracture							1	(2%)
Cranium, inflammation, chronic active							1	(2%)
Femur, fracture				(2%)		(2%)	1	(2%)
Skeletal muscle			(1)		(2)			
Head, inflammation, acute					2	(100%)		
NERVOUS SYSTEM								
Brain	(50)		(50)		(50)		(47)	
Compression						(6%)	1	(2%)
Hemorrhage		(10%)				(8%)		(4%)
Infiltration cellular, lymphocytic		(8%)	_	(6%)		(8%)		(6%)
Mineralization	35	(70%)	31	(62%)	30	(60%)		(60%)
Hippocampus, cyst				(0 <i>0</i>				(2%)
Meninges, inflammation, acute			1	(2%)		(90%)	1	(2%)
Nerve, inflammation, acute Ventricle, dilatation	1	(2%)			1	(2%)		
RESPIRATORY SYSTEM		······						
Lung	(50)		(50)		(50)		(47)	
Congestion	13	(26%)		(4%)		(18%)		(17%)
Hemorrhage	5	(10%)	8	(16%)		(14%)		(13%)
Hyperplasia, lymphoid	5	(10%)	1	(2%)				
Infiltration cellular, lymphocytic	30	(60%)	42	(84%)	40	(80%)	43	(91%)
Mineralization		(4%)	7	(14%)			1	(2%)
Pigmentation, cholesterol		(2%)						
Alveolar epithelium, hyperplasia		(4%)		(6%)		(6%)		(9%)
Alveolus, infiltration cellular, histiocytic		(4%)	2	(4%)		(6%)		(15%)
Interstitium, inflammation, acute	3	(6%)		(0 <i>a</i> ')		(2%)	1	(2%)
Interstitium, inflammation, chronic		(00)	1	(2%)		(2%)		
Peribronchiolar, inflammation, acute Pleura, inflammation, chronic	1	(2%)				(2%)		(OC)
						(2%)	1	(2%)
Pleura, interstitium, inflammation, acute Nose	(50)		(50)		(50)	(2%)	(47)	
11056		(62%)		(56%)		(56%)	(47)	(53%)
Lumen hemorrhege		(62%) (6%)		(2%)		(8%)		(13%)
Lumen, hemorrhage Mucosa, inflammation, acute		(2%)		(2%) (2%)		(2%)	o	(1370)
Mucosa, inflammation, acute				(2%)		(10%)	9	(4%)
Mucosa, inflammation, acute Nasolacrimal duct, hemorrhage		(270)	1			(1070)		(T 10)
Mucosa, inflammation, acute Nasolacrimal duct, hemorrhage Nasolacrimal duct, inflammation, acute	1					(82%)	31	(66%)
Mucosa, inflammation, acute Nasolacrimal duct, hemorrhage Nasolacrimal duct, inflammation, acute Olfactory epithelium, degeneration	1 48	(96%)	31	(62%)		(82%)		
Mucosa, inflammation, acute Nasolacrimal duct, hemorrhage Nasolacrimal duct, inflammation, acute Olfactory epithelium, degeneration Olfactory epithelium, inflammation, acute	1 48		31			(82%)	3	(6%)
Mucosa, inflammation, acute Nasolacrimal duct, hemorrhage Nasolacrimal duct, inflammation, acute Olfactory epithelium, degeneration Olfactory epithelium, inflammation, acute Olfactory epithelium, metaplasia	1 48	(96%)	31 1	(62%) (2%)	41		3 3	(6%) (6%)
Mucosa, inflammation, acute Nasolacrimal duct, hemorrhage Nasolacrimal duct, inflammation, acute Olfactory epithelium, degeneration Olfactory epithelium, inflammation, acute Olfactory epithelium, metaplasia Respiratory epithelium, degeneration	1 48 18		31 1	(62%)	41	(82%)	3 3 12	(6%) (26%)
Mucosa, inflammation, acute Nasolacrimal duct, hemorrhage Nasolacrimal duct, inflammation, acute Olfactory epithelium, degeneration Olfactory epithelium, inflammation, acute Olfactory epithelium, metaplasia	1 48 18	(96%)	31 1 19	(62%) (2%)	41 14		3 3 12	(6%) (6%)

TABLE D5. SUMMARY OF THE INCIDENCE OF NONNEOPLASTIC LESIONS IN FEMALE MICE IN THE TWO-YEAR INHALATION STUDY OF TOLUENE (Continued)

	Chamber	Control	120	ppm	600 j	ppm	1,200 p	pm
RESPIRATORY SYSTEM (Continued)								
Trachea	(50)		(50)		(50)		(47)	/a.w.\
Hemorrhage						(04)	1	(2%)
Inflammation, acute					1	(2%)		(OW)
Glands, inflammation, acute								(2%)
Lumen, hemorrhage							1	(2%)
SPECIAL SENSES SYSTEM								
Eye			(1)					
Atrophy			1	(100%)				
Harderian gland			(3)		(2)		(3)	
Hyperplasia							1	(33%)
Infiltration cellular, lymphocytic			1	(33%)				
Inflammation, acute								(33%)
Inflammation, chronic active							1	(33%)
URINARY SYSTEM								
Kidney	(50)		(50)		(50)		(47)	
Abscess			1	(2%)	1	(2%)		
Congestion							1	(2%)
Hemorrhage					1	(2%)		
Hyperplasia, lymphoid	6	(12%)	1	(2%)				
Infarct	1	(2%)					1	(2%)
Infiltration cellular, lymphocytic	31	(62%)	42	(84%)	36	(72%)	38	(81%)
Inflammation					1	(2%)		
Inflammation, acute							1	(2%)
Inflammation, chronic					1	(2%)		
Metaplasia, osseous	4	(8%)	1	(2%)	3	(6%)	4	(9%)
Mineralization			1	(2%)				
Regeneration							1	(2%)
Capsule, inflammation, chronic						(2%)		
Pelvis, dilatation	2	(4%)	1	(2%)		(2%)		
Pelvis, inflammation, acute					_	(4%)		
Renal tubule, casts protein		(12%)		(22%)	_	(10%)		(26%)
Renal tubule, dilatation		(48%)		(32%)		(60%)		(51%)
Renal tubule, necrosis	5	(10%)		(14%)	_	(18%)		(13%)
Renal tubule, pigmentation		(0.00)		(2%)	_	(2%)		(2%)
Renal tubule, regeneration	14	(28%)	_	(26%)		(46%)	15	(32%)
Ureter			(1)	/100m3	(1)	/*AA~:		
Dilatation				(100%)	1	(100%)		
Infiltration cellular, lymphocytic	(50)		_	(100%)	(EQ)		(47)	
Urinary bladder Angiectasis	,,	(2%)	(50)		(50)		(47)	(2%)
Ectasia	1	(270)			9	(4%)	1	(470)
Hyperplasia, lymphoid	Ę	(10%)	1	(2%)	2	(%70)		
Infiltration cellular, lymphocytic	-	(62%)	_	(78%)	24	(68%)	25	(74%)

APPENDIX E

RESULTS OF SEROLOGIC ANALYSIS

		PAGE
TABLE E1	MURINE ANTIBODY DETERMINATIONS FOR RATS AND MICE IN THE FIFTEEN- MONTH AND TWO-YEAR INHALATION STUDIES OF TOLUENE	213

APPENDIX E. RESULTS OF SEROLOGIC ANALYSIS

Methods

Rodents used in the Carcinogenesis Program of the National Toxicology Program are produced in optimally clean facilities to eliminate potential pathogens that may affect study results.

Data were collected on 5 F344/N control rats of each sex and 10 female $B6C3F_1$ control mice killed at 15 months and from 5/50 or 5/60 randomly selected control animals of each sex and species that lived to the end of the studies. The blood from each animal was collected and clotted, and the serum was separated. The serum was cooled on ice and shipped to Microbiological Associates' Comprehensive Animal Diagnostic Service for determination of the antibody titers. The following tests were performed:

	Hemagglutination <u>Inhibition</u>	Complement <u>Fixation</u>	<u>ELISA</u>
Mice	PVM (pneumonia virus of mice) (15 mo) Reo 3 (reovirus type 3) (15 mo) GDVII (Theiler's encephalomyelitis virus) (15 mo) Poly (polyoma virus) MVM (minute virus of mice) Ectro (infectious ectromelia) (15 mo) Sendai (15 mo)	M. Ad (mouse adenovirus) LCM (lymphocytic choriomeningitis virus) (15 mo) IFA EDIM (epizootic diarrhea of infant mice) (24 mo)	MHV (mouse hepatitis virus) Reo 3 (24 mo) M. Ad. (24 mo) PVM (24 mo) Sendai (24 mo) Ectro (24 mo) GDVII (24 mo) M. pul. (Mycoplasma pulmonis) (24 mo) M. arth. (Mycoplasma arthritidis) (24 mo)
Rats	PVM (15 mo) KRV (Kilham rat virus) H-1 (Toolan's H-1 virus) Sendai (15 mo)		RCV/SDA (rat coronavirus/sialodacryoadenitis) (15,24 mo) Sendai (24 mo) PVM (24 mo) M. pul. (24 mo) M. arth. (24 mo)

Results

Results are presented in Table E1.

TABLE E1. MURINE ANTIBODY DETERMINATIONS FOR RATS AND MICE IN THE FIFTEEN-MONTH AND TWO-YEAR INHALATION STUDIES OF TOLUENE (a)

Interval (months)	Number of Animals	Positive Serologic Reaction for
тѕ		
15	9/10 3/10	Sendai RCV/SDA
24	10/10 10/10 2/10 1/10 2/10	PVM Sendai RCV/SDA Possibly <i>M. arth.</i> <i>M. pul.</i> (b)
CE 15	8/10	мну
24	9/10 9/9 3/7	PVM MHV EDIM

⁽a) Blood samples were taken at 15 and 24 months from control animals just before they were killed; samples were sent to Microbiological Associates (Bethesda, MD) for determination of antibody titers.

(b) Further evaluation of this assay indicated that it was not specific for M. pulmonis, and these results were considered to be

false positive.

APPENDIX F

INGREDIENTS, NUTRIENT COMPOSITION, AND CONTAMINANT LEVELS IN NIH 07 RAT AND MOUSE RATION

Pelleted Diet: October 1982 to November 1984

(Manufactured by Zeigler Bros., Inc., Gardners, PA)

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TABLE F4	CONTAMINANT LEVELS IN NIH 07 RAT AND MOUSE RATION	218

TABLE F1. INGREDIENTS OF NIH 07 RAT AND MOUSE RATION (a)

Ingredients (b)	Percent by Weight					
Ground #2 yellow shelled corn	24.50					
Ground hard winter wheat	23.00					
Soybean meal (49% protein)	12.00					
Fish meal (60% protein)	10.00					
Wheat middlings	10.00					
Dried skim milk	5.00					
Alfalfa meal (dehydrated, 17% protein)	4.00					
Corn gluten meal (60% protein)	3.00					
Soy oil	2.50					
Dried brewer's yeast	2.00					
Dry molasses	1.50					
Dicalcium phosphate	1.25					
Ground limestone	0.50					
Salt	0.50					
Premixes (vitamin and mineral)	0.25					

TABLE F2. VITAMINS AND MINERALS IN NIH 07 RAT AND MOUSE RATION (a)

	Amount	Source
Vitamins		
A	5,500,000 IU	Stabilized vitamin A palmitate or acetate
D_3	4,600,000 IU	D-activated animal sterol
K ₃	2.8 g	Menadione
d-a-Tocopheryl acetate	20,000 IŬ	
Choline	560.0 g	Choline chloride
Folic acid	2.2 g	
Niacin	30.0 g	
d-Pantothenic acid	18.0 g	d-Calcium pantothenate
Riboflavin	3.4 g	•
Thiamine	10.0 g	Thiamine mononitrate
B ₁₂	4,000 µg	
Pyridoxine	1.7 g	Pyridoxine hydrochloride
Biotin	140.0 mg	d-Biotin
Ainerals		
Iron	120.0 g	Iron sulfate
Manganese	60.0 g	Manganous oxide
Zinc	16.0 g	Zinc oxide
Copper	4.0 g	Copper sulfate
Iodine	1.4 g	Calcium iodate
Cobalt	0.4 g	Cobalt carbonate

⁽a) Per ton (2,000 lb) of finished product

⁽a) NCI, 1976; NIH, 1978
(b) Ingredients ground to pass through a U.S. Standard Screen No. 16 before being mixed

TABLE F3. NUTRIENT COMPOSITION OF NIH 07 RAT AND MOUSE RATION

Nutrients	Mean ± Standard Deviation	Range	Number of Samples
Protein (percent by weight)	23.01 ± 1.07	21.3-26.3	26
Crude fat (percent by weight)	5.25 ± 0.70	3.3-6.5	26
Crude fiber (percent by weight)	3.51 ± 0.51	2.8-5.6	26
Ash (percent by weight)	6.66 ± 0.32	6.1-7.1	26
Amino Acids (percent of total d	iet)		
Arginine	1.32 ± 0.072	1.310-1.390	5
Cystine	0.319 ± 0.088	0.218-0.400	5
Glycine	1.146 ± 0.063	1.060-1.210	5
Histidine	0.571 ± 0.026	0.531-0.603	5
Isoleucine	0.914 ± 0.030	0.881-0.944	5
Leucine	1.946 ± 0.056	1.850-1.990	5
Lysine	1.280 ± 0.067	1.200-1.370	5
Methionine	0.436 ± 0.165	0.306-0.699	5
Phenylalanine	0.938 ± 0.158	0.665-1.05	5
Threonine	0.855 ± 0.035	0.824-0.898	5
Tryptophan	0.277 ± 0.221	0.156-0.671	5
Tyrosine	0.618 ± 0.086	0.564-0.769	5
Valine	1.108 ± 0.043	1.050-1.170	5
Essential Fatty Acids (percent o	f total diet)		
Linoleic	2.290 ± 0.313	1.83-2.52	5
Linolenic	0.258 ± 0.040	0.210-0.308	5
Vitamins			
Vitamin A (IU/kg)	$12,289 \pm 4,640$	4,100-24,000	26
Vitamin D (IU/kg)	$4,450 \pm 1,382$	3,000-6,300	4
a-Tocopherol (ppm)	43.58 ± 6.92	31.1-48.0	5
Thiamine (ppm)	18.42 ± 4.01	12.0-27.0	26
Riboflavin (ppm)	7.6 ± 0.85	6.10-8.2	5
Niacin (ppm)	97.8 ± 31.68	65.0-150.0	5
Pantothenic acid (ppm)	30.06 ± 4.31	23.0-34.0	5
Pyridoxine (ppm)	7.68 ± 1.31	5.60-8.8	5
Folic acid (ppm)	2.62 ± 0.89	1.80-3.7	5
Biotin (ppm)	0.254 ± 0.053	0.19-0.32	5
Vitamin B ₁₂ (ppb)	24.21 ± 12.66	10.6-38.0	5
Choline (ppm)	$3,122 \pm 416.8$	2,400-3,430	5
Minerals			
Calcium (percent)	1.27 ± 0.13	0.95-1.54	26
Phosphorus (percent)	0.97 ± 0.06	0.87-1.10	26
Potassium (percent)	0.900 ± 0.098	0.772-0.971	3
Chloride (percent)	0.513 ± 0.114	0.380-0.635	5
Sodium (percent)	0.323 ± 0.043	0.258-0.371	5
Magnesium (percent)	0.167 ± 0.012	0.151-0.181	5
Sulfur (percent)	0.304 ± 0.064	0.268-0.420	5
Iron (ppm)	410.3 ± 94.04	262.0-523.0	5
Manganese (ppm)	90.29 ± 7.15	81.7-99.4	5
Zinc (ppm)	52.78 ± 4.94	46.1-58.2	5
Copper (ppm)	10.72 ± 2.76	8.09-15.39	5
Iodine (ppm)	2.95 ± 1.05	1.52-3.82	4
Chromium (ppm)	1.85 ± 0.25	1.44-2.09	5
Cobalt (ppm)	0.681 ± 0.14	0.490-0.780	4

TABLE F4. CONTAMINANT LEVELS IN NIH 07 RAT AND MOUSE RATION

Contaminants	Mean ± Standard Deviation	Range	Number of Samples
Arsenic (ppm)	0.54 ± 0.17	0.17-0.77	26
Cadmium (ppm)	<0.10		26
Lead (ppm)	0.58 ± 0.20	0.33-1.27	26
Mercury (ppm) (a)	< 0.05	V.00 I.I.	26
Selenium (ppm)	0.32 ± 0.07	0.13-0.42	26
Aflatoxins (ppb) (a)	< 5.0	0.20 0.02	26
Nitrate nitrogen (ppm) (b)	9.96 ± 4.90	0.10-22.0	26
Nitrite nitrogen (ppm) (b)	1.05 ± 1.61	0.10-7.20	26
BHA (ppm) (c)	3.23 ± 3.95	2.00-17.00	26
BHT (ppm) (c)	2.62 ± 2.40	1.00-12.00	26
Aerobic plate count (CFU/g) (d)	$47,473 \pm 37,556$	7,100-130,000	26
Coliform (MPN/g) (e)	40.69 ± 97.61	3.00-460	26
E. coli (MPN/g)	3.04 ± 0.20	3.00-4.00	26
Total nitrosamines (ppb) (f)	5.64 ± 5.66	1.80-30.90	26
N-Nitrosodimethylamine (ppb) (f)	4.59 ± 5.67	0.80-30.00	26
N-Nitrosopyrrolidine (ppb) (f)	1.05 ± 0.24	0.90-1.70	26
Pesticides (ppm)			
a-BHC (a,g)	< 0.01		26
β-BHC (a)	< 0.02		26
γ-BHC (a)	< 0.01		26
δ-BHC (a)	< 0.01		26
Heptachlor (a)	< 0.01		26
Aldrin (a)	< 0.01		26
Heptachlor epoxide (a)	< 0.01		26
DDE (a)	< 0.01		26
DDD (a)	< 0.01		26
DDT(a)	< 0.01		26
HCB(a)	< 0.01		26
Mirex (a)	< 0.01		26
Methoxychlor (a)	< 0.05		26
Dieldrin (a)	< 0.01		26
Endrin (a)	< 0.01		26
Telodrin (a)	< 0.01		26
Chlordane (a)	< 0.05		26
Toxaphene (a)	< 0.1		26
Estimated PCBs (a)	< 0.2		26
Ronnel (a)	< 0.01		26
Ethion (a)	< 0.02		26
Trithion (a)	< 0.05		26
Diazinon (a)	< 0.1		26
Methyl parathion (a)	< 0.02		26
Ethyl parathion (a)	< 0.02		26
Malathion (h)	0.11 ± 0.09	0.05-0.45	26
Endosulfan I (a)	< 0.01		26
Endosulfan II (a)	< 0.01		26
Endosulfan sulfate (a)	< 0.03		26

⁽a) All values were less than the detection limit, given in the table as the mean.
(b) Source of contamination: alfalfa, grains, and fish meal
(c) Source of contamination: soy oil and fish meal
(d) CFU = colony-forming unit
(e) MPN = most probable number
(f) All values were corrected for percent recovery.
(g) BHC = hexachlorocyclohexane or benzene hexachloride
(h) Fifteen lots contained more than 0.05 ppm.

APPENDIX G

METHODS FOR EVALUATION OF REPRODUCTIVE ORGAN TOXICITY IN THE FOURTEEN-WEEK AND FIFTEEN-WEEK INHALATION STUDIES OF TOLUENE

APPENDIX G. METHODS

The right testis and epididymis of rats and mice were removed and placed in a disposable weigh boat. The epididymis was dissected free of the testis, and excess fat was trimmed away.

The right cauda epididymis was removed, weighed, and placed in a prewarmed (32° C) Petri dish containing 1 ml sterile prewarmed (32° C) Tyrode's solution (mice) or sterile phosphate-buffered saline (rats).

The right cauda was secured with a small forceps, gently chopped with a scalpel, and incubated for 15 minutes (mice) or 5 minutes (rats) to release its contents. The right testis and corpus epididymis were weighed. Immediately before the end of the incubation period, a prewarmed (32° C) standard microscope slide and an American Optical hemocytometer were placed on a microscope for the evaluation of sperm motility and sperm progressive drive range. When the incubation period was completed, the suspension was mixed by gently swirling the Petri dish and was aspirated with a prewarmed (32° C) Pasteur pipet. The same pipet was used to distribute samples for the motility and drive range determinations. Two drops of suspension were placed on the microscope slide and covered with a prewarmed (32° C) coverslip (24×50 mm). One drop of suspension was used to fill the hemocytometer. Only the assays for progressive motility and general motility required live samples. Each sample was evaluated by two viewers, A and B, working independently.

Using the $40 \times$ objective of a microscope, Viewer A randomly selected motile sperm. The time required for each randomly selected sperm to traverse 0.1 mm (two small squares) was noted to the nearest 100th second with a stopwatch. Only sperm moving horizontally or vertically in a straight line were considered. If no progressive motility or fewer than 50 motile sperm were observed, a statement to that effect was entered on the data form. All data recorded for this test were entered by a designated recorder. Viewer A observed and timed the progressive motility of 25 sperm cells.

Viewer B, also using the $40\times$ objective of a microscope, initially determined whether more or less than 50% of the sperm in the viewing field were motile. Viewer B then counted in increments of 5% the percentage of sperm above or below 50% of the sperm that were motile. On completion of the motility estimations for the field, Viewer B recorded the findings on the data sheet. Viewer B determined motility for five separate fields of view. Viewers A and B then changed workstations, and the measurements were repeated.

After the live sperm tests were completed, a sperm count was performed. The Petri dish containing the sperm was gently swirled to resuspend the cells. A 1-ml glass micropipet then was used to pipet 0.5 ml of the sperm suspension into a glass test tube $(15 \times 150 \text{ mm})$ containing 2 ml sterile Tyrode's solution (mice) or 4 ml sterile phosphate-buffered saline (rats). After being agitated with a vortex mixer, the suspension was immersed in a hot water bath for about 1 minute. The sperm mixture then was aspirated with a Pasteur pipet, and a drop was placed in the counting chamber of a hemocytometer. The sperm in five large grid areas were counted, and the results, as well as the initial dilution factor, were recorded on the data form. Two counts were performed, each with a separately aspirated sample.

The suspension remaining in the Petri dish was pipetted with a Pasteur pipet into a disposable culture tube (10×75 mm) containing 1 drop of eosin Y stain (1%) in water and allowed to stand for 45 minutes. At the conclusion of the staining time, the solution was gently aspirated. One drop of the stained suspension was placed on a standard, pencil-labeled microscope slide and was spread by one pass of a coverslip. Six smears were prepared from each suspension. The slides were placed at an angle in a microscope slide box and covered with transparent polyethylene. After drying, the slides were coverslipped.

Smears for evaluation of the vaginal cytology were taken between 7:00 a.m. and 9:00 a.m. from 7-9 (mice) or 12-14 (rats) consecutive days before the animals were killed and also on the morning of the kill. The methods used for taking smears were as described in the protocol provided by the laboratory evaluating sperm morphology and vaginal cytology.

The microscope slides used for making the smears were marked into a grid consisting of seven squares on the back of the slide. The squares were labeled 1 through 7. Since more smears than the seven called for in the original protocol were made, a second slide was prepared for each animal with squares labeled 8 through the last day a smear was taken for that animal. Slides were prepared in duplicate for each animal.

One drop of 0.9% saline solution was placed on the appropriate square of the microscope slide. A medicine dropper was then moistened by aspirating the saline solution. The tip of the moistened medicine dropper was placed in the vagina, and the vaginal fluids were aspirated several times. The contents of the medicine dropper were transferred onto the microscope slide and allowed to air dry. The slides were stored in closed slide boxes between collection days.

After completion of smearing, the slides were loaded into glass racks and were stained as follows:

- 1. Absolute ethanol 1 minute
- 2. 95% ethanol 1 minute
- 3. 95% ethanol 1 minute
- 4. Distilled water 1 minute
- 5. 0.5% toluidine blue in 20% ethanol 45 seconds or less
- 6. Running tap water until tap water ran clear
- 7. Distilled water 1 minute

The stained slides were blotted gently with bibulous paper; care was taken not to use the same paper on more than one slide. The slides were allowed to dry completely and then were coverslipped.

The slides for both the vaginal cytology and sperm morphology evaluations were identified by a coding system, as described in the protocol, and were shipped to the laboratory evaluating sperm morphology and vaginal cytology.

APPENDIX H

HEMATOLOGIC AND SERUM CHEMICAL DATA IN THE THIRTEEN-WEEK GAVAGE AND FOURTEEN-WEEK AND FIFTEEN-WEEK INHALATION STUDIES AND HEMATOLOGIC DATA AND ORGAN WEIGHTS IN THE FIFTEEN-MONTH INHALATION STUDIES OF RATS AND MICE EXPOSED TO TOLUENE

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TABLE H1. HEMATOLOGIC AND SERUM CHEMICAL DATA FOR RATS IN THE THIRTEEN-WEEK GAVAGE STUDIES OF TOLUENE (a) $\,$

Analysis	Vehicle Control	312 mg/kg	625 mg/kg	1,250 mg/kg	2,500 mg/kg		
MALE							
Number examined (b)	10	10	10	10	2		
Eosinophils (103/mm3)	0.04 ± 0.013	0.09 ± 0.021	0.06 ± 0.025	0.05 ± 0.017	(c)		
Hematocrit (percent)	42.3 ± 0.32	43.3 ± 0.62	42.9 ± 0.45	43.1 ± 0.47	43.2 ± 2.10		
Hemoglobin (g/dl)	16.2 ± 0.13	16.7 ± 0.26	16.5 ± 0.16	16.5 ± 0.24	16.7 ± 0.95		
Lymphocytes (103/mm3)	4.5 ± 0.20	4.3 ± 0.36	5.3 ± 0.23	4.5 ± 0.27	(c)		
Mean corpuscular hemoglobin (pg Mean corpuscular hemoglobin		20.4 ± 0.09	**20.0 ± 0.11	**20.0 ± 0.14	21.3 ± 0.70		
concentration (g/dl)	38.4 ± 0.17	38.5 ± 0.12	38.2 ± 0.08	38.4 ± 0.24	38.6 ± 0.35		
Mean cell volume (μ ³)	53.3 ± 0.15	53.0 ± 0.21	**52.2 ± 0.25	**52.2 ± 0.13	55.5 ± 1.50		
Methemoglobin (percent)	2.26 ± 0.722	2.84 ± 0.595	3.83 ± 0.863	3.17 ± 0.934	1.54 ± 1.537		
Monocytes (10 ³ /mm ³)	0.13 ± 0.039	0.09 ± 0.020	0.18 ± 0.029	0.13 ± 0.032	(c)		
Platelets (103/mm3)	583 ± 8.3	575 ± 10.8	591 ± 17.0	**640 ± 11.9	*685 ± 35.0		
Erythrocytes (106/mm ³)	7.94 ± 0.060	$*8.17 \pm 0.111$	**8.25 ± 0.064	**8.27 ± 0.092	7.82 ± 0.190		
Reticulocytes (percent)	2.74 ± 0.167	3.18 ± 0.294	$*3.40 \pm 0.204$	**3.57 ± 0.194	*3.95 ± 0.350		
Segmented neutrophils (103/mm3)		1.27 ± 0.079	1.30 ± 0.143	1.17 ± 0.089	(c)		
Leukocytes (103/mm3)	5.82 ± 0.265	5.70 ± 0.374	6.86 ± 0.283	5.93 ± 0.325	6.70 ± 0.800		
Albumin (g/dl)	3.7 ± 0.05	3.8 ± 0.07	3.8 ± 0.07	**4.0 ± 0.05	4.1 ± 0.30		
Blood urea nitrogen (mg/dl)	10.6 ± 0.27	10.6 ± 0.48	11.3 ± 0.31	11.4 ± 0.69	12.5 ± 1.60		
Calcium (g/dl)	11.0 ± 0.05	11.0 ± 0.14	*11.3 ± 0.08	**11.4 ± 0.09	10.9 ± 0.30		
Lactic dehydrogenase (IU/liter)	423 ± 48.8	433 ± 45.5	478 ± 56.0	478 ± 59.2	488 ± 79.0		
Inorganic phosphorus (mg/dl) Total protein (g/dl)	6.74 ± 0.114 6.96 ± 0.156	7.08 ± 0.230 7.09 ± 0.186	7.04 ± 0.106	**7.69 ± 0.188	*7.70 ± 0.500		
•	0.50 ± 0.150	7.09 ± 0.100	7.14 ± 0.169	$*7.55 \pm 0.195$	7.85 ± 0.450		
FEMALE							
Number examined (b)	10	10	10	10	9		
Eosinophils (103/mm3)	0.07 ± 0.025	0.05 ± 0.022	0.04 ± 0.011	0.08 ± 0.017	(c)		
Hematocrit (percent)	41.6 ± 0.52	42.6 ± 0.27	41.7 ± 0.54	42.2 ± 0.59	40.8 ± 0.43		
Hemoglobin (g/dl)	15.8 ± 0.22	16.1 ± 0.08	15.8 ± 0.19	16.1 ± 0.23	15.4 ± 0.20		
Lymphocytes (103/mm3)	4.1 ± 0.36	4.3 ± 0.34	3.6 ± 0.10	4.4 ± 0.18	(c)		
Mean corpuscular hemoglobin (pg) Mean corpuscular hemoglobin		21.5 ± 0.06	21.4 ± 0.12	*21.4 ± 0.09	21.5 ± 0.10		
concentration (g/dl)	37.9 ± 0.13	37.7 ± 0.15	37.9 ± 0.21	38.1 ± 0.13	37.7 ± 0.18		
Mean cell volume (µ³)	57.5 ± 0.17	56.9 ± 0.23	**56.4 ± 0.31	**56.1 ± 0.28	**56.9 ± 0.11		
Methemoglobin (percent) Monocytes (10 ³ /mm ³)	3.04 ± 0.768 0.04 ± 0.017	2.31 ± 0.589	2.90 ± 0.713	3.20 ± 0.674	2.20 ± 0.697		
Platelets (10 ³ /mm ³)	0.04 ± 0.017 592 ± 20.5	0.05 ± 0.022 597 ± 13.7	*0.10 ± 0.012	*0.12 ± 0.030	(c)		
Erythrocytes (106/mm ³)	7.27 ± 0.086		602 ± 14.1	622 ± 7.1	*654 ± 24.9		
Reticulocytes (percent)	2.92 ± 0.086	7.48 ± 0.045 2.97 ± 0.162	7.37 ± 0.084 2.75 ± 0.312	7.52 ± 0.123 3.02 ± 0.372	7.16 ± 0.068 **4.57 ± 0.356		
Segmented neutrophils (103/mm3)		0.83 ± 0.093	0.81 ± 0.079	3.02 ± 0.372 1.01 ± 0.125	**4.57 ± 0.356 (d) 1.28 ± 0.219		
Leukocytes (103/mm3)	5.29 ± 0.353	5.21 ± 0.292	4.57 ± 0.079	5.61 ± 0.123	$(d) 6.70 \pm 0.219$ $(d) 6.70 \pm 0.800$		
Albumin (g/dl)	3.6 ± 0.07	3.7 ± 0.07	3.7 ± 0.05	*3.8 ± 0.05	*3.8 ± 0.07		
Blood urea nitrogen (mg/dl)	10.5 ± 0.36	11.8 ± 0.48	10.9 ± 0.13	11.5 ± 0.49	10.2 ± 0.57		
Calcium (g/dl)	10.7 ± 0.12	10.7 ± 0.09	10.7 ± 0.08	10.8 ± 0.06	*10.9 ± 0.14		
Lactic dehydrogenase (IU/liter)	351 ± 41.6	368 ± 36.5	406 ± 65.8	392 ± 39.0	390 ± 32.5		
Inorganic phosphorus (mg/dl)	6.08 ± 0.238	6.00 ± 0.203	6.33 ± 0.259	6.43 ± 0.168	**7.30 \pm 0.216		
Total protein (g/dl)	6.53 ± 0.183	6.76 ± 0.208	6.80 ± 0.113	**7.14 ± 0.136	*7.06 ± 0.187		

⁽a) Mean \pm standard error. P values are vs. the vehicle controls by Dunn's test (Dunn, 1964) or Shirley's test (Shirley, 1977). IU = international units.

⁽b) Except as noted
(c) Fewer than two animals were examined.

⁽d) Two animals were examined.

^{*}P<0.05

^{**}P<0.01

TABLE H2. HEMATOLOGIC AND SERUM CHEMICAL DATA FOR RATS IN THE FIFTEEN-WEEK INHALATION STUDIES OF TOLUENE (a) $\,$

walle fumber examined eukocytes (10 ³ /mm ³) symphocytes (percent) egmented neutrophils (percent) fosinophils (percent) fematocrit (percent) fematocrit (percent) fematocrit (percent) fean corpuscular hemoglobin (g/dl) fean corpuscular hemoglobin concentration (g/dl) fean cell volume (cubic microns) fethemoglobin (g/dl) latelets (10 ³ /mm ³)	10 8.19 ± 0.441 77.5 ± 1.86 19.2 ± 1.91 2.7 ± 0.30 0.6 ± 0.27 50.3 ± 1.17 17.3 ± 0.47 21.3 ± 0.21 34.4 ± 0.45 61.8 ± 0.73	10 8.12 ± 0.409 73.7 ± 1.31 23.1 ± 1.52 1.9 ± 0.31 1.3 ± 0.30 51.8 ± 0.51 17.9 ± 0.19 21.3 ± 0.12 34.6 ± 0.40	10 7.43 ± 0.367 73.7 ± 2.13 22.2 ± 2.22 2.6 ± 0.58 1.4 ± 0.27 51.1 ± 0.65 17.6 ± 0.15 21.1 ± 0.17	10 7.19 ± 0.443 73.2 ± 3.01 23.8 ± 2.75 1.8 ± 0.47 1.2 ± 0.25 51.0 ± 0.96 17.4 ± 0.24	10 7.86 ± 0.623 *71.1 ± 2.12 24.5 ± 2.22 3.1 ± 0.53 1.1 ± 0.23 50.6 ± 1.17 17.3 ± 0.55	2 7.70 ± 0.095 71.5 ± 3.50 24.0 ± 2.00 3.5 ± 1.50 1.0 ± 0.00 48.3 ± 2.05
eukocytes (103/mm3) symphocytes (percent) legmented neutrophils (percent) dosnocytes (percent) dosnophils (percent) demadorit (percent) femoglobin (g/dl) fean corpuscular hemoglobin (pg) fean corpuscular hemoglobin concentration (g/dl) fean cell volume (cubic microns) fethemoglobin (g/dl)	8.19 ± 0.441 77.5 ± 1.86 19.2 ± 1.91 2.7 ± 0.30 0.6 ± 0.27 50.3 ± 1.17 17.3 ± 0.47 21.3 ± 0.21 34.4 ± 0.45	8.12 ± 0.409 73.7 ± 1.31 23.1 ± 1.52 1.9 ± 0.31 1.3 ± 0.30 51.8 ± 0.51 17.9 ± 0.19 21.3 ± 0.12	7.43 ± 0.367 73.7 ± 2.13 22.2 ± 2.22 2.6 ± 0.58 1.4 ± 0.27 51.1 ± 0.65 17.6 ± 0.15	7.19 ± 0.443 73.2 ± 3.01 23.8 ± 2.75 1.8 ± 0.47 1.2 ± 0.25 51.0 ± 0.96	7.86 ± 0.623 *71.1 ± 2.12 24.5 ± 2.22 3.1 ± 0.53 1.1 ± 0.23 50.6 ± 1.17	7.70 ± 0.095 71.5 ± 3.50 24.0 ± 2.00 3.5 ± 1.50 1.0 ± 0.00
.ymphocytes (percent) egmented neutrophils (percent) fonocytes (percent) losinophils (percent) lematocrit (percent) lematocrit (percent) lemoglobin (g/dl) fean corpuscular hemoglobin (pg) fean corpuscular hemoglobin concentration (g/dl) fean cell volume (cubic microns) fethemoglobin (g/dl)	77.5 ± 1.86 19.2 ± 1.91 2.7 ± 0.30 0.6 ± 0.27 50.3 ± 1.17 17.3 ± 0.47 21.3 ± 0.21 34.4 ± 0.45	73.7 ± 1.31 23.1 ± 1.52 1.9 ± 0.31 1.3 ± 0.30 51.8 ± 0.51 17.9 ± 0.19 21.3 ± 0.12	73.7 ± 2.13 22.2 ± 2.22 2.6 ± 0.58 1.4 ± 0.27 51.1 ± 0.65 17.6 ± 0.15	73.2 ± 3.01 23.8 ± 2.75 1.8 ± 0.47 1.2 ± 0.25 51.0 ± 0.96	*71.1 ± 2.12 24.5 ± 2.22 3.1 ± 0.53 1.1 ± 0.23 50.6 ± 1.17	71.5 ± 3.50 24.0 ± 2.00 3.5 ± 1.50 1.0 ± 0.00
egmented neutrophils (percent) Aonocytes (percent) Cosinophils (percent) Eematocrit (percent) Eematocrit (percent) Eematocrit (percent) Eematocrit (grdl) Aean corpuscular hemoglobin (pg) Eean corpuscular hemoglobin concentration (grdl) Aean cell volume (cubic microns) Aethemoglobin (grdl)	19.2 ± 1.91 2.7 ± 0.30 0.6 ± 0.27 50.3 ± 1.17 17.3 ± 0.47 21.3 ± 0.21 34.4 ± 0.45	23.1 ± 1.52 1.9 ± 0.31 1.3 ± 0.30 51.8 ± 0.51 17.9 ± 0.19 21.3 ± 0.12	22.2 ± 2.22 2.6 ± 0.58 1.4 ± 0.27 51.1 ± 0.65 17.6 ± 0.15	23.8 ± 2.75 1.8 ± 0.47 1.2 ± 0.25 51.0 ± 0.96	24.5 ± 2.22 3.1 ± 0.53 1.1 ± 0.28 50.6 ± 1.17	24.0 ± 2.00 3.5 ± 1.50 1.0 ± 0.00
(percent) Aoncytes (percent) Costinophils (percent) Tematocrit (percent) Temoglobin (g/dl) Aean corpuscular Temoglobin (pg) Aean corpuscular hemoglobin Concentration (g/dl) Aean cell volume (cubic microns) Aethemoglobin (g/dl)	2.7 ± 0.30 0.6 ± 0.27 50.3 ± 1.17 17.3 ± 0.47 21.3 ± 0.21 34.4 ± 0.45	$ \begin{array}{cccc} 1.9 & \pm & 0.31 \\ 1.3 & \pm & 0.30 \\ 51.8 & \pm & 0.51 \\ 17.9 & \pm & 0.19 \end{array} $ $ 21.3 & \pm & 0.12 $	2.6 ± 0.58 1.4 ± 0.27 51.1 ± 0.65 17.6 ± 0.15	$\begin{array}{cccc} 1.8 & \pm & 0.47 \\ 1.2 & \pm & 0.25 \\ 51.0 & \pm & 0.96 \end{array}$	3.1 ± 0.53 1.1 ± 0.28 50.6 ± 1.17	3.5 ± 1.50 1.0 ± 0.00
Aonocytes (percent) losinophils (percent) lematocrit (percent) lemoglobin (g/dl) lean corpuscular hemoglobin (pg) lean corpuscular hemoglobin concentration (g/dl) lean cell volume (cubic microns) lethemoglobin (g/dl)	2.7 ± 0.30 0.6 ± 0.27 50.3 ± 1.17 17.3 ± 0.47 21.3 ± 0.21 34.4 ± 0.45	$ \begin{array}{cccc} 1.9 & \pm & 0.31 \\ 1.3 & \pm & 0.30 \\ 51.8 & \pm & 0.51 \\ 17.9 & \pm & 0.19 \end{array} $ $ 21.3 & \pm & 0.12 $	2.6 ± 0.58 1.4 ± 0.27 51.1 ± 0.65 17.6 ± 0.15	$\begin{array}{cccc} 1.8 & \pm & 0.47 \\ 1.2 & \pm & 0.25 \\ 51.0 & \pm & 0.96 \end{array}$	3.1 ± 0.53 1.1 ± 0.28 50.6 ± 1.17	3.5 ± 1.50 1.0 ± 0.00
losinophils (percent) Iematocrit (percent) Iematocrit (percent) Iemaglobin (g/dl) Iema corpuscular Iemaglobin (pg) Ieman corpuscular Iemaglobin Ieman contentration (g/dl) Ieman cell volume Ieubic microns) Iethemoglobin (g/dl)	0.6 ± 0.27 50.3 ± 1.17 17.3 ± 0.47 21.3 ± 0.21 34.4 ± 0.45	$ \begin{array}{rcl} 1.3 & \pm & 0.30 \\ 51.8 & \pm & 0.51 \\ 17.9 & \pm & 0.19 \end{array} $ $ 21.3 & \pm & 0.12 $	$\begin{array}{ccc} 1.4 & \pm & 0.27 \\ 51.1 & \pm & 0.65 \\ 17.6 & \pm & 0.15 \end{array}$	1.2 ± 0.25 51.0 ± 0.96	1.1 ± 0.23 50.6 ± 1.17	1.0 ± 0.00
Tematocrit (percent) Temoglobin (g/dl) Tean corpuscular Themoglobin (pg) Tean corpuscular hemoglobin Concentration (g/dl) Tean cell volume (cubic microns) Tethemoglobin (g/dl)	50.3 ± 1.17 17.3 ± 0.47 21.3 ± 0.21 34.4 ± 0.45	51.8 ± 0.51 17.9 ± 0.19 21.3 ± 0.12	51.1 ± 0.65 17.6 ± 0.15	51.0 ± 0.96	50.6 ± 1.17	
Hemoglobin (g/dl) Hean corpuscular Hemoglobin (pg) Hean corpuscular hemoglobin concentration (g/dl) Hean coll volume (cubic microns) Hethemoglobin (g/dl)	17.3 ± 0.47 21.3 ± 0.21 34.4 ± 0.45	17.9 ± 0.19 21.3 ± 0.12	17.6 ± 0.15			
Mean corpuscular hemoglobin (pg) Mean corpuscular hemoglobin concentration (g/dl) Mean cell volume (cubic microns) Methemoglobin (g/dl)	21.3 ± 0.21 34.4 ± 0.45		21.1 ± 0.17			17.0 ± 0.10
Mean corpuscular hemoglobin concentration (g/dl) Mean cell volume (cubic microns) Methemoglobin (g/dl)	34.4 ± 0.45		21.1 ± 0.17			
concentration (g/dl) Mean cell volume (cubic microns) Methemoglobin (g/dl)		346 + 040		20.8 ± 0.16	21.0 ± 0.28	22.1 ± 0.10
Aean cell volume (cubic microns) Aethemoglobin (g/dl)		346 + 040				
(cubic microns) Aethemoglobin (g/dl)	61.8 ± 0.73	34.0 I U.40	34.4 ± 0.60	34.2 ± 0.47	34.1 ± 0.67	35.3 ± 1.30
Aethemoglobin (g/dl)	61.8 T 0.73	01.77 + 0.70	61.2 ± 0.73	60.9 ± 0.69	61.6 ± 0.56	63.0 ± 2.00
	1.19 ± 0.263	61.7 ± 0.52 1.35 ± 0.221	61.2 ± 0.73 1.28 ± 0.204	1.58 ± 0.333	0.91 ± 0.270	1.12 ± 0.64
TOTAL TO THE !	555 ± 29.0	542 ± 18.5	520 ± 13.9	547 ± 11.8	548 ± 20.1	564 ± 27.0
Crythrocytes (10 ⁶ /mm ³)	8.13 ± 0.199	8.39 ± 0.086	8.35 ± 0.065	8.36 ± 0.094	8.22 ± 0.196	7.70 ± 0.09
leticulocytes (10 ⁶ /mm ³)	3.22 ± 0.286	3.18 ± 0.252	3.18 ± 0.364	2.86 ± 0.367	3.38 ± 0.256	4.10 ± 0.00
lbumin/globulin ratio	1.03 ± 0.021	1.04 ± 0.022	1.06 ± 0.016	1.05 ± 0.027	**1.12 ± 0.020	*1.15 ± 0.05
lbumin (g/dl)	3.75 ± 0.040	3.77 ± 0.047	3.70 ± 0.015	3.69 ± 0.046	3.83 ± 0.063	3.75 ± 0.05
Jrea nitrogen (mg/dl)	18.5 ± 0.68	18.7 ± 1.03	18.6 ± 0.64	18.3 ± 0.92	17.1 ± 0.78	19.6 ± 0.20
Calcium (mg/dl)	11.0 ± 0.17	11.1 ± 0.18	10.8 ± 0.09	11.0 ± 0.14	11.1 ± 0.16	11.0 ± 0.20
Chloride (meq/liter)	106 ± 1.1	105 ± 1.1	104 ± 1.1	104 ± 1.2	104 ± 1.1	109 ± 6.5
holinesterase (IU/liter)	714 ± 16	712 ± 18	682 ± 15	671 ± 14	**630 ± 19	*595 ± 9
reatinine (mg/dl)	0.54 ± 0.043	0.61 ± 0.038	0.50 ± 0.021 1.2 ± 0.13	0.48 ± 0.025	0.51 ± 0.035 1.3 ± 0.15	0.45 ± 0.05 1.5 ± 0.50
GT (IU/liter) norganic phosphorus (mg/dl)	1.1 ± 0.10 7.12 ± 0.213	1.1 ± 0.10 7.01 ± 0.333	1.2 ± 0.13 6.39 ± 0.133	1.0 ± 0.00 6.97 ± 0.294	7.46 ± 0.501	7.35 ± 0.05
organic phosphorus (mg/di)	5.75 ± 0.174	5.75 ± 0.201	5.36 ± 0.136	5.75 ± 0.190	6.29 ± 0.192	6.45 ± 0.65
lucose (mg/dl)	144 ± 5.8	147 ± 5.2	148 ± 4.0	148 ± 3.5	144 ± 4.4	138 ± 0.0
odium (meq/liter)	147 ± 1.1	147 ± 1.2	147 ± 0.7	146 ± 1.2	148 ± 1.4	152 ± 7.0
otal bilirubin (mg/dl)	0.26 ± 0.027	0.22 ± 0.013	0.27 ± 0.033	0.26 ± 0.037	0.25 ± 0.022	0.15 ± 0.05
otal protein (g/dl)	7.42 ± 0.088	7.45 ± 0.073	7.24 ± 0.045	7.25 ± 0.109	7.31 ± 0.167	7.05 ± 0.15
FEMALE						
Number examined (b)	10	10	10	10	10	10
eukocytes (10 ³ /mm ³)	7.26 ± 0.392	6.75 ± 0.387	6.48 ± 0.480	*5.98 ± 0.352	**5.64 ± 0.342	*6.39 ± 0.38
ymphocytes (percent)	76.9 ± 1.91	72.2 ± 2.39	74.8 ± 1.82	77.0 ± 1.98	76.3 ± 1.80	77.5 ± 1.76
egmented neutrophils						
(percent)	20.4 ± 1.79	24.7 ± 2.25	21.9 ± 1.46	19.7 ± 2.04	21.4 ± 1.83	19.2 ± 1.60
(onocytes (percent)	2.2 ± 0.33	2.4 ± 0.43	2.2 ± 0.36	2.6 ± 0.45	1.0 ± 0.26	2.7 ± 0.26
osinophils (percent)	0.4 ± 0.22	0.6 ± 0.22	1.1 ± 0.38	0.6 ± 0.22	1.1 ± 0.35	0.6 ± 0.31
[ematocrit (percent)	48.4 ± 1.33	49.4 ± 0.66	49.5 ± 0.93	49.8 ± 0.97	48.5 ± 1.01	49.5 ± 0.70
[emoglobin (g/dl)	16.4 ± 0.36	16.9 ± 0.21	16.8 ± 0.20	17.0 ± 0.32	16.2 ± 0.44	16.7 ± 0.20
fean corpuscular hemoglobin (pg)	22.8 ± 0.08	22.8 ± 0.13	22.6 ± 0.14	22.5 ± 0.13	*22.0 ± 0.38	*22.4 ± 0.13
dean corpuscular hemoglobin	22.0 ± 0.00	22.0 1 0.10	22.0 1 0.14	22.0 1 0.10	22.0 1 0.00	22.4 _ 0.20
concentration (g/dl)	33.9 ± 0.32	34.1 ± 0.43	33.9 ± 0.39	34.1 ± 0.45	33.4 ± 0.73	33.8 ± 0.35
fean cell volume	33.5 - 3.55	**** =				
(cubic microns)	67.3 ± 0.45	66.8 ± 0.57	66.7 ± 0.40	66.1 ± 0.53	66.1 ± 0.48	66.3 ± 0.47
iethemoglobin (g/dl)	1.19 ± 0.300	1.28 ± 0.301	0.68 ± 0.187	1.15 ± 0.318	0.91 ± 0.232	0.79 ± 0.24
latelets (103/mm ³)	577 ± 22.6	573 ± 32.9	541 ± 19.3	(c) 582 ± 19.1	544 ± 12.7	564 ± 20.3
rythrocytes (10 ⁶ /mm ³)	7.18 ± 0.164	7.40 ± 0.078	7.42 ± 0.110	7.53 ± 0.124	7.35 ± 0.140	7.46 ± 0.07
leticulocytes (10 ⁶ /mm ³)	3.10 ± 0.343	3.97 ± 0.295	3.04 ± 0.387 1.12 ± 0.025	2.96 ± 0.240	3.72 ± 0.321 1.16 ± 0.022	3.38 ± 0.25
dbumin/globulin ratio dbumin (g/dl)	1.11 ± 0.018 3.68 ± 0.060	1,12 ± 0.039 3.74 ± 0.040	1.12 ± 0.025 3.64 ± 0.040	1.11 ± 0.018 3.69 ± 0.046	$\begin{array}{cccc} 1.16 & \pm & 0.022 \\ 3.50 & \pm & 0.052 \end{array}$	1.15 ± 0.02 3.53 ± 0.03
rea nitrogen (mg/dl)	17.8 ± 1.07	18.7 ± 0.65	16.5 ± 0.50	16.3 ± 0.49	16.3 ± 0.80	16.2 ± 0.91
alcium (mg/dl)	10.5 ± 0.11	10.8 ± 0.11	10.6 ± 0.05	10.7 ± 0.12	10.2 ± 0.14	10.4 ± 0.11
Chloride (meq/liter)	106 ± 1.5	105 ± 1.1	106 ± 1.1	106 ± 1.4	106 ± 0.9	106 ± 1.1
holinesterase (IU/liter)	3,701 ± 83	3,644 ± 78	*3,166 ± 209	**2,997 ± 205	**1,798 ± 162	**1,702 ± 163
reatinine (mg/dl)	0.47 ± 0.015	0.51 ± 0.028	0.46 ± 0.027	0.48 ± 0.029	0.47 ± 0.026	0.50 ± 0.02
GT (IU/liter)	1.4 ± 0.16	1.5 ± 0.17	1.4 ± 0.22	1.6 ± 0.22	1.3 ± 0.15	1.5 ± 0.17
norganic phosphorus (mg/dl)	6.31 ± 0.198	6.38 ± 0.118	5.84 ± 0.236	6.43 ± 0.216	6.08 ± 0.243	6.43 ± 0.22 5.48 ± 0.16
otassium (meq/liter)	5.21 ± 0.084	5.29 ± 0.091	5.20 ± 0.098	5.46 ± 0.127	5.14 ± 0.064 133 ± 3.0	5.48 ± 0.16 134 ± 3.0
lucose (mg/dl)	131 ± 2.0	134 ± 2.7 145 ± 0.8	$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	134 ± 3.2 146 ± 1.3	133 ± 3.0 145 ± 0.9	146 ± 1.1
odium (meq/liter) otal bilirubin (mg/dl)	144 ± 1.1 0.15 ± 0.017	145 ± 0.8 0.19 ± 0.018	0.15 ± 0.022	0.17 ± 0.021	0.13 ± 0.015	0.15 ± 0.02
otal protein (g/dl)	6.98 ± 0.107	7.09 ± 0.087	6.92 ± 0.090	7.02 ± 0.079	**6.53 ± 0.110	*6.61 ± 0.07

⁽a) Mean ± standard error. P values are vs. the controls by Dunn's test (Dunn, 1964) or Shirley's test (Shirley, 1977). GGT = Y-glutamyl transferase; cholinesterase activity was measured in plasma; IU = international units.
(b) Except as noted
(c) Nine animals were examined.
*P<0.05
**P<0.01

TABLE H3. HEMATOLOGIC DATA FOR RATS IN THE FIFTEEN-MONTH INHALATION STUDIES OF TOLUENE (a) $\,$

Analysis	C	Control			600 ppm			1,200 ppm			
MALE											
Leukocytes (1,000/µl)	5.9	±	0.43	6.2	±	0.39	6.9	±	0.36		
Lymphocytes (1,000/μl)	3.4	±	0.31	3.9	±	0.27	3.8	±	0.32		
Segmented neutrophils (1,000/μl)	2.2	±	0.27	2.1	±	0.22	2.8	±	0.40		
Monocytes (1,000/µl)	(b) 0.11	±	0.011	(b) 0.13	±	0.024	(c) 0.20		0.053		
Eosinophils (1,000/µl)	0.16	±		(d) 0.18	±	0.025	(e) 0.22		0.040		
Hematocrit (percent)	49.6	±		50.4		0.88	49 .0				
lemoglobin (g/dl)	18.3	±	0.23	18.6		0.32	18.2		0.49		
Methemoglobin (g/dl)	0.35	±	0.031	(b) 0.30	_	0.032	0.35		0.044		
Mean corpuscular hemoglobin (pg)	20.0	±	0.18	19.8		0.10	19.4		0.30		
Mean corpuscular hemoglobin concentration (g/dl)	37.0	±	0.25	36.8		0.10	37.2	_	0.25		
Mean cell volume (µ³)	54.1	±	0.64	53.4		0.34	*52.2		0.66		
Nucleated erythrocytes (1,000/µl)	0.08	±	0.034	0.03		0.017	0.01		0.005		
Erythrocytes (106/µl)	9.2	±	0.08	9.4	±	0.20	9.4	±	0.19		
FEMALE											
-eukocytes (1,000/μl)	3.9	±	0.39	3.5	±	0.21	3.7	±	0.21		
ymphocytes (1,000/μl)	2.5	±	0.17	2.4	±	0.16	2.5	±	0.14		
Segmented neutrophils (1,000/µl)	1.4	±	0.32	1.0	±	0.08	1.0	±	0.10		
Monocytes (1,000/µl)	(e) 0.10	±	0.00	(f) 0.10	±	0.00	(d) 0.10	±	0.00		
Cosinophils (1,000/µl)	(d) 0.10	±	0.00	(c) 0.10	±	0.00	(e) 0.18	±	0.040		
Iematocrit (percent)	48.6	±	0.69	47.7	±	0.34	48.4	±	0.58		
Hemoglobin (g/dl)	18.1	±	0.23	17.7	±	0.16	17.9	±	0.21		
fethemoglobin (g/dl)	0.26	±	0.015	0.32	±	0.042	0.30	±	0.030		
Mean corpuscular hemoglobin (pg)	21.8	±	0.13	21.8	±	0.09	21.6	±	0.11		
Mean corpuscular hemoglobin concentration (g/dl)	37.1	±	0.14	37.0	±	0.17	37.1	±	0.15		
Mean cell volume (µ3)	58.8	±	0.29	58.7	±	0.30	58.4	±	0.27		
Nucleated erythrocytes (1,000/µl)	0.06	±	0.016	0.06	±	0.017	0.03	±	0.011		
Crythrocytes (106/µl)	8.3	±	0.12	8.1	\pm	0.09	8.3	±	0.10		

⁽a) Mean ± standard error for groups of 10 animals unless otherwise specified; P values vs. the controls by Dunn's test (Dunn, 1964) or Shirley's test (Shirley, 1977).
(b) Nine animals were examined.

⁽c) Seven animals were examined.
(d) Eight animals were examined.

⁽e) Six animals were examined.

⁽f) Five animals were examined. *P<0.05

TABLE H4. HEMATOLOGIC AND SERUM CHEMICAL DATA FOR MICE IN THE THIRTEEN-WEEK GAVAGE STUDIES OF TOLUENE (a)

Analysis	Vehicle Co	ontrol	312 m	ıg/kg	625 m	ng/kg	1,250 1	ng/kg	2,500 m	g/kg
MALE						- "				
Number examined (b)	10		10		10		10		6	
Eosinophils (10 ³ /mm ³)	0.04 ±	0.019	0.09 ±	0.043	0.05 ±	0.019	0.01 ±	0.008	(c) 0.08 ±	0.055
Hematocrit (percent)	38.4 ±	0.41	36.9 ±	0.68	37.8 ±	0.39	38.0 ±	0.32	37.4 ±	0.64
Hemoglobin (g/dl)	14.9 ±	0.22	14.4 ±	0.25	14.9 ±	0.15	14.9 ±	0.19	14.8 ±	0.20
Lymphocytes (10 ³ /mm ³)	2.75 ±	0.321	2.98 ±	0.668	3.11 ±	0.346	2.50 ±	0.464	2.89 ±	0.382
Mean corpuscular hemoglobin (pg) Mean corpuscular hemoglobin	18.9 ±	0.23	19.1 ±	0.16	19.2 ±	0.12	19.3 ±	0.13	19.2 ±	0.04
concentration (g/dl) Mean cell volume	38.8 ±	0.40	39.1 ±	0.32	39.5 ±	0.37	39.2 ±	0.42	39.6 ±	0.34
(cubic microns)	48.6 ±	0.27	48.8 ±	0.53	48.7 ±	0.30	49.5 ±	0.40	48.3 ±	0.56
Methemoglobin (percent)	7.57 ±	1.205	12.12 ±	1.303	6.24 ±	1.301	6.21 ±	1.379	(d) 6.16 ±	2.997
Monocytes 10 ³ /mm ³)	0.07 ±	0.017	0.07 ±	0.022	0.07 ±	0.014	0.07 ±	0.017	(e) 0.11 ±	0.033
Platelets (10 ³ /mm ³)	864 ±	20.6	865 ±	45.8	798 ±	36.8	863 ±	54.1	809 ±	74.0
Erythrocytes (10 ⁶ /mm ³)	7.89 ±	0.062	*7.56 ±	0.096	7.76 ±	0.064	7.71 ±	0.076	7.72 ±	0.110
Reticulocytes (percent)	(e) $4.57 \pm$	0.570	(c) 3.38 ±	0.410	(f) 4.11 ±	0.275	(g) 3.70 ±	0.473	(c) 3.88 ±	0.265
Segmented neutrophils (103/mm3)	0.75 ±	0.083	0.80 ±	0.200	0.87 ±	0.097	0.50 ±	0.082	0.93 ±	0.090
Leukocytes (10 ³ /mm ³)	3.61 ±	0.382	3.95 ±	0.893	4.11 ±	0.391	3.08 ±	0.557	4.02 ±	0.450
Albumin (g/dl)	(f) 3.58 ±	0.073	(f) 3.56 ±	0.094	3.52 ±	0.047	3.52 ±	0.036	(c) 3.78 ±	0.162
Blood urea nitrogen (mg/dl)		1.32	(f) 23.3 ±	3.88	19.6 ±	0.69	19.0 ±	0.47	(c) 18.6 ±	1.70
Calcium (mg/dl)	(g) $10.8 \pm$	0.23	(e) 11.1 ±	0.30	(f) $10.8 \pm$	0.22	(f) 11.3 ±		*(h) 14.2 ±	1.95
Lactic dehydrogenase (IU/liter)	(f) 928 ±	89.9	(f) 908 ±	101.8	754 ±	104.7	837 ±	102.7	(c) 990 ±	167.1
Inorganic phosphorus (mg/dl) Total protein (g/dl)	(e) 9.78 ± (g) 6.91 ±	0.535 0.108	(e) 10.33 ± (f) 7.05 ±	0.868 0.304	(f) 9.90 ± 7.10 ±	0.387 0.136	(f) 9.99 ± (f) 7.13 ±	0.753 0.164	(h) 9.15 ± (h) 7.50 ±	3.050 0.100
FEMALE										
Number examined (b)	10		10		10		9		6	
Eosinophils (10 ³ /mm ³)	0.03 ±	0.018	0.06 ±	0.015	0.02 ±	0.010	(f) 0.06 ±	0.021	*(c) 0.07 ±	0.018
Hematocrit (percent)		0.38	37.7 ±	0.56	38.4 ±	0.59	37.6 ±	0.71	37.9 ±	0.30
Hemoglobin (g/dl)	14.8 ±	0.14	14.5 ±	0.11	14.8 ±	0.14	14.6 ±	0.23	14.9 ±	0.09
Lymphocytes (10 ³ /mm ³)		1,420	2.45 ±	0.225	2.20 ±	0.436	(f) 2.34 ±	0.236	(c) 3.19 ±	0.355
Mean corpuscular hemoglobin (pg)		0.11	19.1 ±	0.09	19.3 ±	0.04	19.3 ±	0.12	*19.4 ±	0.14
Mean corpuscular										
hemoglobin concentration (g/dl) Mean cell volume	38.6 ±	0.36	38.5 ±	0.33	38.6 ±	0.35	38.7 ±	0.37	39.3 ±	0.25
(cubic microns)	49.8 ±	0.51	49.5 ±	0.52	50.0 ±	0.47	49.9 ±	0.26	49.7 ±	0.33
Methemoglobin (percent)		1.532	10.05 ±	1.777	7.51 ±	1.628	(f) 6.53 ±	1.301	(c) 5.88 ±	1.946
Monocytes 10 ³ /mm ³)	0.04 ±	0.012	0.06 ±	0.011	0.07 ±	0.020	0.07 ±	0.017	(c) 0.07 ±	0.017
Platelets (10 ³ /mm ³)	756 ±	25.2	801 ±	27.0	795 ±	30.6	739 ±	55.4	*838 ±	36.9
Erythrocytes (10 ⁶ /mm ³)	7.75 ±	0.077	7.58 ±	0.057	7.69 ±	0.068	7.56 ±	0.123	7.67 ±	0.073
Reticulocytes (percent)	(e) 3.05 ±	0.593	(e) 2.72 ±	0.634	(g) $4.07 \pm$	0.275	(g) $4.04 \pm$	0.428	3.40 ±	0.465
Segmented neutrophils (10 ³ /mm ³)	0.78 ±	0.150	0.83 ±	0.128	0.83 ±	0.092	0.73 ±	0.053	(c) 1.06 ±	0.140
Leukocytes (10 ³ /mm ³)		1.569	3.39 ±	0.265	$3.12 \pm$	0.486	3.23 ±	0.286	4.18 ±	0.452
Albumin (g/dl)	3.58 ±	0.083	3.56 ±	0.034	(i) 3.69 ±	0.070	*(f) 3.80 ±	0.060	3.80 ±	0.086
Blood urea nitrogen (mg/dl)		0.72	**15.9 ±	0.42	**(i) 17.5 ±	0.65		1.28	*16.7 ±	0.39
Calcium (mg/dl)	(i) 10.7 \pm	0.15	(i) $10.3 \pm$	0.22	(g) $10.5 \pm$	0.22	(e) 11.0 ±	0.19	**11.9 ±	0.18
Lactic dehydrogenase (TU/liter)	628 ±	48.8	580 ±	31.5	(f) 683 ±	58.7	(g) 649 ±	81.4	694 ±	99.3
norganic phosphorus (mg/dl)	(i) 9.64 ±	0.701	9.34 ±	0.568	(g) $8.56 \pm$	0.241	(e) 9.22 ±	0.601	10.87 ±	0.650
Total protein (g/dl)	(i) 6.37 ±	0.177	6.30 ±	0.087	(f) 6.75 ±	0.217	(e) 6.63 ±	0.079	*7.12 ±	0.218

⁽a) Mean \pm standard error. P values are vs. the vehicle controls by Dunn's test (Dunn, 1964) or Shirley's test (Shirley, 1977). IU = international units.

⁽a) Mean ± standard error. P value (b) Except as noted (c) Five animals were examined. (d) Four animals were examined. (e) Six animals were examined. (f) Eight animals were examined. (g) Seven animals were examined. (h) Two animals were examined. (i) Nine animals were examined. *P<0.05

TABLE H5. HEMATOLOGIC AND SERUM CHEMICAL DATA FOR MICE IN THE FOURTEEN-WEEK INHALATION STUDIES OF TOLUENE (a)

Analysis	Con	trol	100	ppm	625	ppm	1,250	ppm	2,500	ppm	3,000	ppn
MALE			-									
Number examined (b)	10		10		10		10		10		4	
Leukocytes (10 ³ /mm ³)	5.17 ±	0.791	4.93 ±	0.593	3.96 ±	0.454	5.31 ±	0.495	5.10 ±	0.673	4.73 ±	1.01
Lymphocytes (percent)	77.9 ±	1.72	75.9 ±	3.66	76.9 ±	4.19	78.7 ±	2.95	73.2 ±	2.14	78.8 ±	3.09
Segmented neutrophils (percent)		1.74	19.1 ±	3.53	20.2 ±	4.38	16.5 ±	2.73	22.8 ±	2.31	17.0 ±	3.6
Monocytes (percent)	2.3 ±	0.34	2.6 ±	0.50	1.6 ±	0.31	2.2 ±	0.47	2.1 ±	0.53	2.0 ±	1.0
Eosinophils (percent)	2.4 ±	0.45	2.2 ±	0.59	1.0 ±	0.26	2.4 ±	0.40	1.7 ±	0.30	2.3 ±	1,1
Hematocrit (percent)	42.1 ±	0.56 0.18	41.9 ±	0.35	41.8 ±	0.64 0.26	42.2 ±	0.59 0.14	42.3 ±	0.47 0.23	41.9 ± 16.6 ±	0.7
Hemoglobin (g/dl)	17.1 ±	0.16	16.9 ±	0.19	16.9 ±	0.26	16.7 ±	0.14	17.0 ±	0.23	10.0 I	0.3
Mean corpuscular hemoglobin (pg)	20.0 ±	0.10	19.7 ±	0.17	20.0 ±	0.18	19.8 ±	0.21	20.2 ±	0.17	20.0 ±	0.2
Mean corpuscular hemoglobin	40.77	0.40	40.0 ±	0.40			00.0 (0.50	40.0	0.40	20.2.4	
concentration (g/dl) Mean cell volume	40.7 ±	0.43	40.3 ±	0.49	40.4 ±	0.52	39.8 ±	0.56	40.2 ±	0.48	39.6 ±	0.5
(cubic microns)	49.1 ±	0.41	49.0 ±	0.30	49.4 ±	0.34	49.9 ±	0.38	*50.3 ±	0.30	50.5 ±	0.6
Methemoglobin (g/dl)	0.76 ±	0.187	0.45 ±	0.115	0.73 ±		0.65 ±		0.53 ±	0.131	0.64 ±	0.2
Platelets (10 ³ /mm ³)	565 ±	49.8	600 ±	26.4	664 ±		666 ±	27.0	661 ±	33.6	507 ±	170
Erythrocytes (10 ⁶ /mm ³)	$8.60 \pm$	0.100	8.55 ±	0.070	8.47 ±	0.130	8.47 ±	0.109	8.42 ±	0.105	8.31 ±	0.0
Reticulocytes (106/mm ³)	4.30 ±	0.402	5.04 ±	0.488	4.18 ±	0.383	4.68 ±	0.421	4.50 ±	0.287	3.98 ±	0.7
Albumin/globulin ratio	1.18 ±	0.025	1.20 ±	0.030	1.17 ±	0.026	1.16 ±	0.031	*1.10 ±	0.015	*1.08 ±	0.0
Albumin (g/dl)	3.37 ±	0.063	3.35 ±	0.045	3.34 ±	0.048	3.16 ±	0.064	3.35 ±	0.069	3.40 ±	0.0
Urea nitrogen (mg/dl)	24.1 ±	1.36	24.5 ±	1.42	24.8 ±	1.88	23.9 ±	1.25	23.1 ±	1.15	(c) 23.3 ±	1.6
Calcium (mg/dl)	9.98 ± 0.15 ±	0.131 0.031	10.03 ±	0.097 0.027	9.99 ±	0.085 0.010	9.74 ±	0.183	*10.63 ±	0.184	*(c) 11.03 ±	0.2
Creatinine (mg/dl) GGT (IU/liter)	1.30 ±	0.031	0.15 ± 1.10 ±	0.027	0.11 ± 1.10 ±	0.010	0.15 ± 1.30 ±	$0.027 \\ 0.213$	0.15 ± 1.00 ±	0.027	(c) 0.10 ±	0.00
norganic phosphorus (mg/dl)	10.0 ±	0.300	10.1 ±	0.100	10.3 ±		9.3 ±	0.213	1.00 ±	0.40	1.25 ± (c) 11.3 ±	0.23
Glucose (mg/dl)	159 ±	5.0	143 ±	7.1	153 ±	7.6	145 ±	3.7	(d) 144 ±	6.5	(c) 152 ±	11.2
Total bilirubin (mg/dl)	0.30 ±	0.037	0.31 ±	0.018	0.33 ±	0.030	0.32 ±	0.025	(e) 0.42 ±	0.039	(c) 0.40 ±	0.05
Total protein (g/dl)	6.21 ±	0.087	6.15 ±	0.093	6.20 ±		5.89 ±	0.126	6.42 ±	0.117	6.63 ±	0.14
FEMALE												
Number examined (b)	10		10		9		9		3		0	
Leukocytes (10 ³ /mm ³)	3.63 ±	0.447	3.40 ±	0.315	3.09 ±	0.551	3.07 ±	0.376	3.50 ±	1.185		
Lymphocytes (percent)	78.7 ±	2.44	(d) 81.8 ±	2.40	80.3 ±	1.67	82.4 ±	1.20	78.7 ±	2.40		
Segmented neutrophils			(4) 01.0 1	4.40	00.0 1	1.01	02.7 1	1.20	70.1 1	2.40		
(percent)	17.2 ±	2.31	(d) 15.4 ±	2.09	16.9 ±	1.74	14.9 ±	1.29	16.3 ±	2.96		
Monocytes (percent)	2.4 ±	0.34	1.9 ±	0.35	1.8 ±	0.47	1.4 ±	0.38	2.7 ±	0.33		
Cosinophils (percent)	$1.7 \pm$	0.37	*0.5 ±	0.40	1.0 ±	0.29	(e) 1.1 ±	0.30	2.3 ±	1.33		
Hematocrit (percent)	41.9 ±	0.34	42.0 ±	0.41	42.5 ±	0.45	42.4 ±	0.56	41.1 ±	0.59		
Hemoglobin (g/dl)	16.7 ±	0.14	16.9 ±	0.13	17.0 ±	0.14	*17.1 ±	0.12	16.4 ±	0.27		
Mean corpuscular hemoglobin (pg)	20.0 ±	0.18	20.2 ±	0.21	20.3 ±	0.20	20.6 ±	0.24	20.3 ±	0.54	**	
Mean corpuscular hemoglobin	20.0 1	3.20	=V.2 1	~. .	40.0 ±	3.20	20.0 I	J. 4/7	20.0 ±	3.04	"	
concentration (g/dl)	39.8 ±	0.48	40.3 ±	0.54	39.9 ±	0.55	40.4 ±	0.57	40.1 ±	1.23		
Mean cell volume												
(cubic microns)	50.2 ±	0.33	50.1 ±	0.43	51.0 ±	0.37	50.9 ±	0.45	50.7 ±	0.33		
Methemoglobin (g/dl)	0.54 ±	0.149	0.74 ±	0.149	0.39 ±	0.112	0.65 ±	0.145	1.11 ±	0.244		
Platelets (10 ³ /mm ³)	663 ± 8.35 ±	14.2 0.063	638 ± 8.37 ±	26.7 0.070	663 ± 8.34 ±	33.6 0.047	676 ± 8.34 ±	29.2 0.094	667 ± 8.12 ±	50.0 0.082		
Crythrocytes (10 ⁶ /mm ³) leticulocytes (10 ⁶ /mm ³)	4.86 ±	0.063	6.37 ±	0.070	6.34 ±	0.448	8.34 I 4.04 ±	0.094	8.12 ± 4.33 ±	0.636		
Albumin/globulin ratio		0.413	1.41 ±	0.038	1.37 ±	0.448	1.29 ±	0.045	4.33 ±	0.067		
lbumin (g/dl)	3.46 ±		3.53 ±	0.054	3.41 ±	0.039	3.40 ±	0.075	3.37 ±	0.120		
Jrea nitrogen (mg/dl)		1.08	21.8 ±	1.02	20.6 ±	1.37	19.8 ±	1.54	21.9 ±	2.24		
Calcium (mg/dl)	10.02 ±		10.02 ±	0.090	9.82 ±	0.090	(e) 10.06 \pm		10.93 ±	0.190		
reatinine (mg/dl)	$0.13 \pm$		0.13 ±	0.015	$0.20 \pm$	0.041	0.13 ±		0.20 ±	0.100		
GT (IU/liter)	1.00 ±		1.00 ±	0.000	1.00 ±	0.000	1.11 ±		1.00 ±	0.000		
norganic phosphorus (mg/dl)	9.8 ±		10.5 ±	0.57	9.9 ±	0.26	10.3 ±	0.41	10.9 ±	0.77		
lucose (mg/dl)	138 ±	6.2	143 ±	4.7	144 ±	7.9	(e) 154 ±	7.7	132 ±	10.0		
Total bilirubin (mg/dl)	0.28 ±	0.025	0.24 ±	0.027	0.23 ±	0.017	0.24 ±	0.018	0.37 ±	0.067		
Fotal protein (g/dl)	5.99 ±	0.071	6.07 ±	0.084	5.94 ±	0.060	6.03 ±	0.099	6.17 ±	0.088		

⁽a) Mean \pm standard error. P values are vs. the controls by Dunn's test (Dunn, 1964) or Shirley's test (Shirley, 1977). GGT = γ -glutamyl transferase; IU = international (a) Mean I standard error. P valuants.
(b) Except as noted
(c) Three animals were examined.
(d) Nine animals were examined.
(e) Eight animals were examined.

TABLE H6. HEMATOLOGIC DATA FOR FEMALE MICE IN THE FIFTEEN-MONTH INHALATION STUDY OF TOLUENE (a)

Analysis	Control 10			120 ppm			600 ppm 9			1,200 ppm		
Number examined (b)												
Leukocytes (1,000/µl)	3.6	±	0.41	3.9	±	0.32	3.5	±	0.32	4.0	±	0.51
Lymphocytes (1,000/µl)	2.5	±	0.32	2.7	±	0.13	2.4	±	0.24	2.7	±	0.30
Segmented neutrophils (1,000/µl)	0.94	±	0.129	1.10	±	0.256	0.96	±	0.113	1.18	±	0.207
Monocytes (1,000/µl)	(c) 0.18	±	0.049	(d) 0.11	±	0.014	(c) 0.11	±	0.012	(c)0.15	±	0.038
Eosinophils (1,000/µl)	(e) 0.10	±	0.000	(f) 0.15	±	0.029	(e) 0.12	±	0.017	(g) 0.10	±	0.000
Hematocrit (percent)	45.1	±	2.32	42.6	±	0.58	43.8	±	0.60	45.1	±	0.78
Hemoglobin (g/dl)	15.8	±	0.45	15.3	±	0.11	15.6	±	0.10	15.7	±	0.16
Methemoglobin (g/dl)	0.22	±	0.062	0.36	±	0.175	0.35	±	0.063	0.34	±	0.086
Mean corpuscular hemoglobin (pg)	19.7	±	0.20	19.3	±	0.14	19.6	±	0.18	19.5	±	0.34
Mean corpuscular hemoglobin												
concentration (g/dl)	36.4	±	0.52	36.0	+	0.39	35.6	+	0.41	34.9	±	0.57
Mean cell volume (µ³)	54.1	Ŧ	0.64	53.6	Ŧ		55.1	+	0.70	56.0	±	1.61
Erythrocytes (106/µl)	8.3	±	0.45	7.9	±		8.0	±	0.11	8.1	±	0.23

⁽a) Mean \pm standard error; no significant differences vs. the controls were observed by Dunn's test (Dunn, 1964) or Shirley's

test (Shirley, 1977). (b) Except as noted

⁽c) Eight animals were examined.
(d) Seven animals were examined.

⁽e) Six animals were examined.

⁽f) Four animals were examined.

⁽g) Five animals were examined.

TABLE H7. ORGAN WEIGHTS OF RATS IN THE FIFTEEN-MONTH INHALATION STUDIES OF TOLUENE (a)

Organ	Co	ntr	ol	600	600 ppm 1,200 pp		pm		
MALE			· · · · · · · · · · · · · · · · · · ·	- , , , , , , , , , , , , , , , , , , ,					
Body weight (grams)	400	±	13.4	397	±	11.4	364	±	11.9
Right kidney									
Absolute	1,327	±	60	1,378	±	37	1,283		49
Relative	3.3	±	0.10	3.5	±	0.09	3.5	±	0.08
iver									
Absolute	(b) 13,770	±	440	13,320		620	12,280		530
Relative	34.7	±	1.18	33.9	±	1.82	33.7	±	0.99
3rain									
Absolute	1,944			1,973			1,873		
Relative	5.0	±	0.18	5.0	±	0.16	5.2	±	0.16
FEMALE									
Body weight (grams)	242	±	3.1	266	±	9.0	238	±	9.0
Right kidney									
Absolute	860	±	23	880	±	23	(b) 854	±	27
Relative	3.6	±	0.08	3.3	±	0.08	(b) 3.6	±	0.09
Liver									
Absolute	8,062	±	340	*10,409			8,486	±	283
Relative	33.3	±	1.19	40.1	±	4.83	35.8		0.72
Brain									
Absolute	1,754	±	27	1,852	±	57	1,783	±	23
Relative	7.3	±	0.12	7.0	±	0.28	7.6		0.27

⁽a) Mean ± standard error in milligrams per gram (relative) or milligrams (absolute) for groups of 10 animals unless otherwise specified; P values vs. the controls by Dunn's test (Dunn, 1964) or Shirley's test (Shirley, 1977).

(b) Nine organs were weighed.

TABLE H8. ORGAN WEIGHTS OF FEMALE MICE IN THE FIFTEEN-MONTH INHALATION STUDY OF TOLUENE (a)

Organ	Co	ntr	ol	12	20 1	ppm	66)O j	ppm	1,2	200	ppm
Body weight (grams)	34.9	±	1.83	33.4	±	1.83	35.1	±	1.88	32.0	±	0.68
Brain												
Absolute	497	±	5.1	487	±	10.5	498	±	9.9	492	±	11.8
Relative	14.5	±	0.61	15.0	±	0.87	14.5	±	0.67	15.5	±	0.54
Right kidney												
Absolute	285	±	12.7	272	±	13.9	285	±	10.4	280	±	16.8
Relative	8.2	±	0.34	8.2	±	0.30	8.2	±	0.32	8.8	±	0.61
Liver												
Absolute	1,854	±	99	1,903	±	89	1,920	±	70	1,920	±	105
Relative	54.3	±	3.89	57.5	±	2.19	55.2	±	1.67	60.4	±	3.81

⁽a) Mean \pm standard error in milligrams per gram (relative) or milligrams (absolute) for groups of 10 animals; no significant differences vs. the controls were observed by Dunn's test (Dunn, 1964).

^{*}P<0.05

APPENDIX I

CHEMICAL CHARACTERIZATION, ANALYSIS, AND GENERATION OF CHAMBER CONCENTRATIONS OF TOLUENE FOR THE TOXICOLOGY STUDIES

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APPENDIX I. CHEMICAL CHARACTERIZATION

Procurement and Characterization of Toluene

Toluene was obtained in one lot (lot no. H-12-19-80) from Exxon Company, USA (Baytown, TX) as a clear, colorless liquid and was received in sixteen 55-gallon drums. Purity and identity analyses were conducted on representative samples at Midwest Research Institute (MRI) (Kansas City, MO). MRI reports on the analyses performed in support of the toluene studies are on file at the National Institute of Environmental Health Sciences.

The study material was identified as toluene by spectroscopic analyses. The infrared (Figure I1), ultraviolet/visible, and nuclear magnetic resonance (Figure I2) spectra were consistent with the literature spectra (Sadtler Standard Spectra) and with those expected for the structure.

The purity of toluene was determined by elemental analysis, Karl Fischer water analysis, and gas chromatography. Gas chromatographic analysis was performed with flame ionization detection and a nitrogen carrier with a flow rate of 70 ml/minute with either a 0.1% SP1000 (system 1) or a 20% SP2100/0.1% Carbowax 1500 (system 2) column. Benzene was identified as an impurity by spiking and was quantitated against standard benzene solutions with gas chromatographic system 2. The results of elemental analyses for carbon and hydrogen were in agreement with the theoretical values. Karl Fischer analysis indicated the presence of 0.047% water. Gas chromatography by both systems detected three impurities with individual peak areas less than 0.1% of the major peak area. Benzene was present as an impurity at 5.7 ppm (v/v). The data indicated that lot no. H-12-19-80 was greater than 99% pure.

Stability studies performed by gas chromatography with system 1 (with chlorobenzene as an internal standard) indicated that toluene was stable as a bulk chemical when stored for 2 weeks protected from light at temperatures up to 60° C.

Periodic analysis of lot no. H-12-19-80 for purity by gas chromatography and ultraviolet spectroscopy and for identity by infrared spectroscopy indicated no apparent degradation of the study material throughout the studies.

Preparation and Characterization of Dose Mixtures

The appropriate amounts of toluene and corn oil were mixed (w/v) to give the desired concentrations. The stability of toluene in corn oil was determined after the sample was extracted with methanol by gas chromatography with flame ionization detection and with the same column as system 2 but at a flow rate of 30 ml/minute and with nonane as an internal standard. Toluene dissolved in corn oil at 20 mg/ml was found to be stable at 5° C and at room temperature in the dark for 2 weeks. Solutions exposed to air and light for 3 hours were chemically stable, but a 23% loss due to evaporation was observed over the 3-hour period. Dose mixtures were stored at room temperature protected from light in Nalgene® bottles for no longer than 2 weeks throughout the studies. Dose mixtures were analyzed several times during the 13-week studies, and concentrations ranged from 91% to 107% of the target concentrations (Table II).

Generation and Measurement of Chamber Concentrations

Vapor Generation System

Liquid toluene was delivered by a pump from a stainless steel safety can through Teflon® tubing to a Spraying Systems® atomizer (Figure I3) that was operated with nitrogen. Nitrogen and the atomizer were heated to approximately 80°C with a 400-W cartridge heater. The toluene was sprayed into a

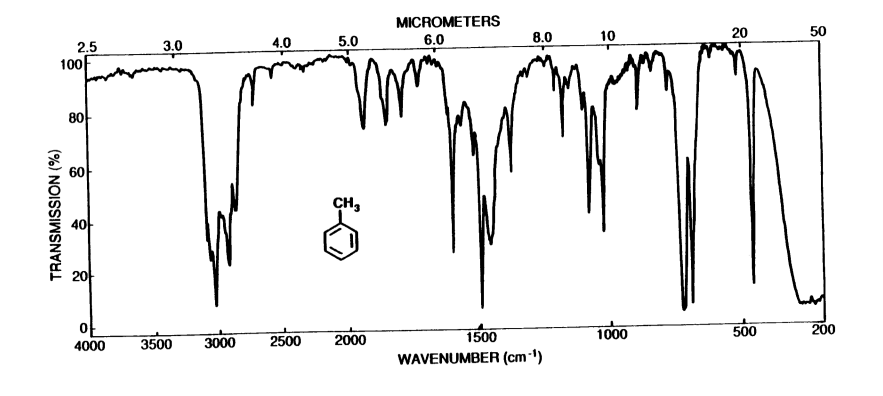


FIGURE 11. INFRARED ABSORPTION SPECTRUM OF TOLUENE (LOT NO. H-12-19-80)

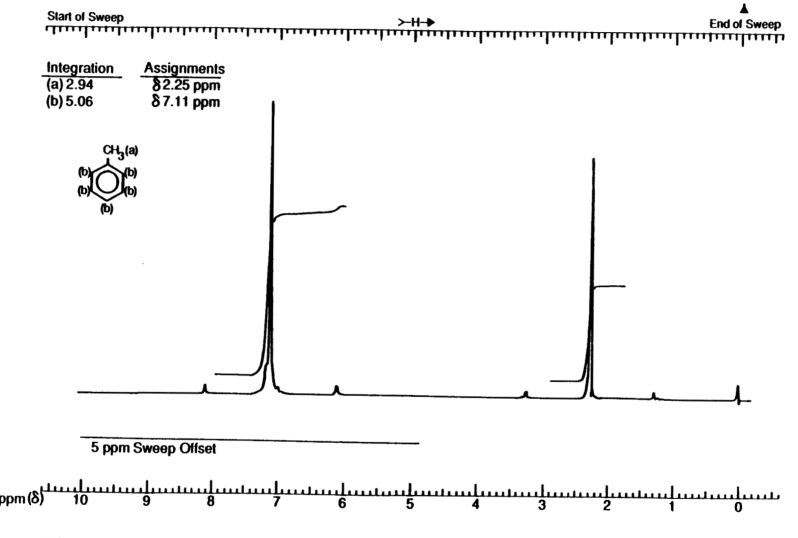


FIGURE 12. NUCLEAR MAGNETIC RESONANCE SPECTRUM OF TOLUENE (LOT NO. H-12-19-80)

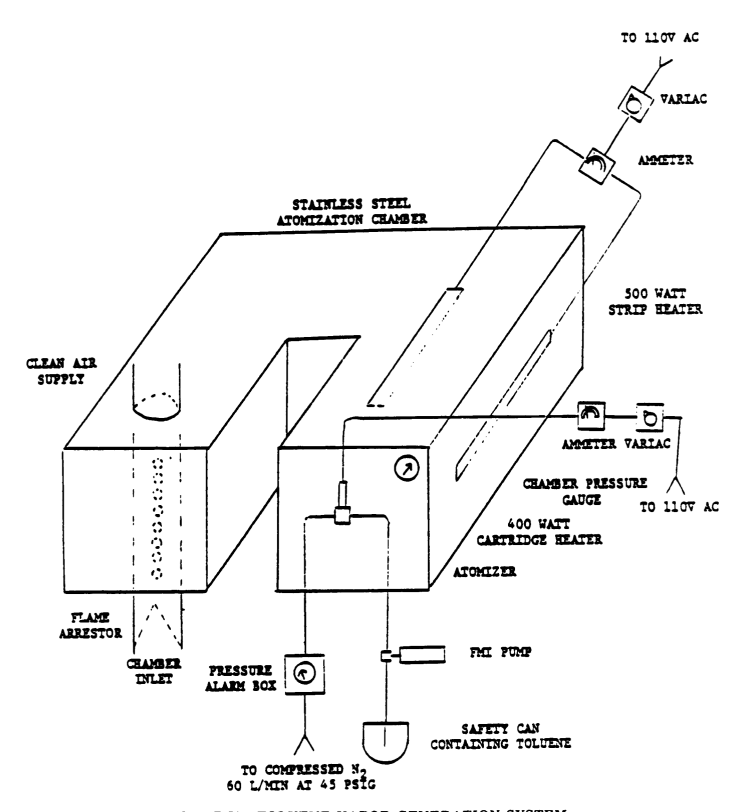


FIGURE 13. TOLUENE VAPOR GENERATION SYSTEM

TABLE II. RESULTS OF ANALYSIS OF DOSE MIXTURES IN THE THIRTEEN-WEEK GAVAGE STUDIES OF TOLUENE

Date Mixed	Concentration of T Target	Oluene in Corn Oil (mg/ml) Determined (a)	Determined as a Percent of Target
05/18/81	31.2	28.4	91
	62.5	57.2	92
	125	116	93
	250	239	96
	500	488	98
05/20/81	31.2	29.5	95
	62.5	59.4	96
	125	118	94
	250	240	96
	500	489	98
06/29/81	62.5	57.7	92
	125	119	95
	250	237	95
06/30/81	31.2	28.6	92
07/01/81	31.2	32.7	105
	62.5	65.2	104
	125	134	107
	250	(b) 260	104
07/13/81	31.2	30.0	96

⁽a) Results of duplicate analysis, except as noted

U-shaped stainless steel atomization chamber. For high dose groups, the atomization chamber walls were also heated by four 500-W strip heaters to a surface temperature of 85° C to increase the rate of vaporization. Toluene vapors flowed into a pipe extending through one end of the atomization chamber and were diluted with chamber ventilation air to produce the desired exposure concentrations in the chambers.

Vapor Concentration Monitoring

The concentration of toluene in the chambers was measured in sampled chamber air at 3.3 μ by a MIRAN® gas-phase infrared spectrophotometer connected to a Hewlett-Packard Model 3388A laboratory computer. Air from each chamber was sampled and analyzed about 5 minutes every hour. Weekly mean exposure concentrations for the 14- and 15-week and 2-year studies are presented in Tables I2 and I3. Toluene aerosol, measured in the 1,200-ppm chamber with a Sibata® P-5 Digital Dust Indicator (2-year studies) or in the 3,000-ppm chamber with a Model CI-252 (Climet Instrument Co.) aerosol particle counter (14-week studies), was not detected in measurable quantitites.

The presence of detectable concentrations (more than 10 ppm) of toluene was determined by analyzing the atmosphere in all chambers postexposure at various times. Measurable concentrations occurred by 4 months after the studies began, and, after further evaluation, the animals and/or caging were indicated as the source of the residual toluene.

⁽b) Average of two duplicate analyses

TABLE 12. MEAN CHAMBER CONCENTRATIONS IN THE FOURTEEN-WEEK AND FIFTEEN-WEEK INHALATION STUDIES OF TOLUENE

Week on		Week	ly Chamber Co	oncentration (p	pm) (a)	
Study	0	100	625	1,250	2,500	3,000
1	5	89	697	584	1,859	3,123
2	2	99	641	732	2,113	3,282
3	7	89	548	600	2,383	2,677
4	1	93	601	913	2,246	2,940
5	1	81	604	1,120	2,116	2,757
6	8	95	605	1,182	2,353	2,992
7	12	97	624	1,276	2,458	2,878
8	16	105	659	1,200	2,592	2,923
9	15	107	688	1,208	2,398	2,980
10	20	95	627	1,205	2,432	2,941
11	22	88	596	1,094	2,323	2,848
12	21	90	562	1,087	2,584	2,865
13	27	91	577	1,113	2,316	2,788
14	25	95	577	1,110	2,366	2,932
(b) 15 (mice)	24	90	573	1,073	2,175	2,760
15 (rats)	23	85	565	1,026	2,175	2,760
(c) 16 (rats)	20	74	491	1,025	2,220	3,013

⁽a) Calculated as the mean of the actual hourly concentration (b) One exposure during week 15 (c) One exposure during week 16

TABLE 13. MEAN CHAMBER CONCENTRATIONS IN THE TWO-YEAR INHALATION STUDIES OF TOLUENE

	n Study		Weekly Mean Chamber	Concentration (ppm	·/
Rats	Mice	0	120	600	1,200
1		0		614	1,211
2		Ŏ		581	1,167
3		Ŏ		563	1,173
4		ĭ		573	1,136
5		ō		624	1,188
6		ŏ	••	621	1,214
7	1	ĭ	122	621	1,210
8	2	i	124	618	1,144
9	3	Ö	118	576	1,230
10	4	ĭ	118	577	1,13
11	5	ō	120	614	1,183
12	6	ŏ	124	617	1,14
13	ž	ĭ	120	618	1,158
14	8	ō	110	580	1,203
15	9	ŏ	115	582	1,093
16	10	ĭ	104	570	1,046
17	11	2	114	559	1,166
18	12	2	113	563	1,186
19	13	1	115	525	1,218
20	14	ō	117	520	1,202
21	15	ĭ	121	499	1,156
22	16	ī	131	614	1,214
23	17	3	122	595	1,178
24	18	8	124	628	1,194
25	19	Ö	118	597	1,20
26	20	Ŏ	121	616	1,226
27	21	Ō	120	579	1,18
28	22	Ö	118	598	1,214
29	23	Ö	123	592	1,200
30	24	Ö	123	583	1,230
31	25	ĺ	119	588	1,20
32	26	ī	117	580	1,186
33	27	Ō	122	584	1,217
34	28	Ö	118	591	1,188
35	29	Ö	124	620	1,171
36	30	Ŏ	124	599	1,149
37	31	Ŏ	123	612	1,117
38	32	Ö	128	618	1,134
39	33	Ö	119	612	1,100
40	34	4	123	609	1,138
41	35	0	121	633	1,16
42	36	$\overset{\circ}{2}$	120	634	1,219
43	37	5	118	604	1,23
44	38	ĺ	125	599	1,24
45	39	1	118	601	1,16
46	40	0	120	585	1,192
47	41	1	120	603	1,198
48	42	0	123	600	1,248
49	43	1	121	598	1,177
50	44	2	123	624	1,290
51	45	3	112	598	1,158
52	46	0	112	549	1,177
53	47	1	118	593	1,230
54	48	2	115	565	1,183
55	49	0	117	596	1,179
56	50	1	111	620	1,159
57	51	7	114	615	1,167
58	52	2	119	582	1,094
59	53	6	122	609	1,164
60	54	8	114	610	1,156

TABLE 13. MEAN CHAMBER CONCENTRATIONS IN THE TWO-YEAR INHALATION STUDIES OF TOLUENE (Continued)

<u>Week</u> o	n Study		Weekly Mean Chamber	r Concentration (ppm	1)
Rats	Mice	0	120	600	1,200
62	56	8	117	568	1,188
63	57	5	122	580	1,196
64	58	8	115	590	1,169
65	59	8 2	118	600	1,149
66	60	0	118	592	1,151
67	61	Ö	121	582	1,158
68	62	Ö	126	608	1,141
69	63	Ŏ	119	609	1,181
70	64	Ö	124	564	1,194
71	65	Ŏ	123	564	1,185
72	66	Ö	128	544	1,182
73	67	ŏ	125	593	1,147
74	68	Ö	127	598	1,149
75	69	Ö	123	601	1,165
76	70	ŏ	121	591	1,179
77	71	ŏ	118	613	1,179
78	72	Ŏ	117	596	1,106
79	73	Ö	119	590	1,201
80	74	Ö	122	597	1,210
81	75	0	120	598	1,210
82	76	4	121	604	1,195
83	77	3	120	590	
84	78	ა 5	120	595	1,150
85	79	5 4	122		1,220
86	80			597	1,208
87	81	4 3	123 118	599 596	1,205
88	82			607	1,209 1,200
	83	3	119		
89		1	119	597	1,189
90	84	2	119	606	1,180
91	85	1	123	592	1,201
92	86	2	119	605	1,197
93	87	2	119	595	1,175
94	88	1	122	606	1,178
95	89	2	121	597	1,212
96	90	1	118	599	1,210
97	91	0	123	598	1,218
98	92	0	116	587	1,184
99	93	0	120	580	1,207
100	94	0	123	576	1,159
101	95	0	121	596	1,210
102	96	0	120	587	1,160
103	97	0	117	585	1,169
	98	0	126	588	1,114
	99	0	118	594	1,018
	100	0	123	607	1,162
	101	0	128	582	1,171
	102	1	124	582	1,117
	103	0	115	592	1,264

APPENDIX I. CHEMICAL CHARACTERIZATION

Vapor Concentration Uniformity in Chamber

The uniformity of the vapor concentration in each exposure chamber was measured at intervals over a 5-month period during the studies with the same system used to monitor the vapor concentration (used as a reference at the site normally used for analytical measurements during the studies) and a second system with a different infrared monitor used as a comparison with the reference. Between each hourly sample for the main analytical system (reference site), the probe on the second system was moved to predetermined test sites that encompassed the entire animal exposure zone; in four of the five tests that used this combined system, the range of variation from the reference was 3%-12%, whereas in the fifth test, it was 26% (77%-103% that of the reference). In the three tests that used only the second infrared monitor (used as both reference and comparison monitors by serially sampling the reference and six comparison sites, beginning and ending with the reference site), variations from the reference position were 2%, 5%, and 14%.

APPENDIX J

GENETIC TOXICOLOGY OF TOLUENE

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APPENDIX J. GENETIC TOXICOLOGY

METHODS

Salmonella Protocol: Testing was performed as reported by Ames et al. (1975) with modifications listed below; both data and detailed protocol are included in Haworth et al. (1983). Chemicals were sent to the laboratories as coded aliquots from Radian Corporation (Austin, TX). The study chemical was incubated with the Salmonella typhimurium tester strains (TA98, TA100, TA1535, and TA1537) either in buffer or S9 mix (metabolic activation enzymes and cofactors from Aroclor 1254-induced male Sprague Dawley rat or Syrian hamster liver) for 20 minutes at 37° C before the addition of soft agar supplemented with L-histidine and D-biotin and subsequent plating on minimal glucose agar plates. Incubation was continued for an additional 48 hours.

Chemicals were tested in four strains; all trials were repeated. Each test consisted of triplicate plates of concurrent positive and negative controls and of at least five doses of the study chemical. The high dose was limited by toxicity or solubility but did not exceed 1 mg/plate. All negative assays were repeated, and all positive assays were repeated under the conditions that elicited the positive response.

A positive response was defined as a reproducible, dose-related increase in histidine-independent (revertant) colonies in any one strain/activation combination. An equivocal response was defined as an increase in revertants which was not dose related, not reproducible, or of insufficient magnitude to support a determination of mutagenicity. A response was considered negative when no increase in revertant colonies was observed after chemical treatment.

Mouse Lymphoma Protocol: The experimental protocol is presented in detail by McGregor et al. (1988) and follows the basic format of Clive et al. (1979). All study chemicals were supplied as coded aliquots from Radian Corporation (Austin, TX). The highest dose of the study compound was determined by solubility or toxicity and did not exceed 5 mg/ml. Mouse L5178Y/TK lymphoma cells were maintained at 37° C as suspension cultures in Fischer's medium supplemented with 2 mM L-glutamine, 110 µg/ml sodium pyruvate, 0.05% pluronic F68, antibiotics, and heat-inactivated horse serum; normal cycling time was about 10 hours. To reduce the number of spontaneously occurring trifluorothymidine (Tft)-resistant cells, subcultures were exposed once to medium containing thymidine, hypoxanthine, methotrexate, and glycine for 1 day, to thymidine, hypoxanthine, and glycine for 1 day, and to normal medium for 3-5 days. For cloning, horse serum content was increased and Noble agar was added. Freshly prepared S9 metabolic activation factors were obtained from the liver of either Aroclor 1254-induced or noninduced male F344 rats.

All doses within an experiment, including concurrent positive and solvent controls, were replicated. Treated cultures contained 6×10^6 cells in 10 ml of medium. This volume included the S9 fraction in those experiments performed with metabolic activation. Incubation with the study chemical continued for 4 hours, after which time the medium plus chemical was removed and the cells were resuspended in 20 ml of fresh medium and incubated for an additional 2 days to express the mutant phenotype. Cell density was monitored so that log phase growth was maintained. After the 48-hour expression period, 3×10^6 cells were plated in medium and soft agar supplemented with Tft for selection of Tft-resistant cells (TK $^{+/+}$), and 600 cells were plated in nonselective medium and soft agar to determine cloning efficiency. Plates were incubated at 37° C under 5% carbon dioxide for 10-12 days. All data were evaluated statistically for both trend and peak response. Both responses had to be significant (P < 0.05) for a chemical to be considered capable of inducing Tft resistance; a single significant response led to an "equivocal" conclusion, and the absence of both a trend and a peak response resulted in a "negative" call.

Minimum criteria for accepting an experiment as valid and a detailed description of the statistical analysis and data evaluation are presented in Myhr et al. (1985). This assay was initially performed without S9; if a clearly positive response was not obtained, the experiment was repeated with induced S9.

APPENDIX J. GENETIC TOXICOLOGY

Chinese Hamster Ovary Cytogenetics Assays: Testing was performed as reported by Galloway et al. (1985, 1987) and is described briefly below. Chemicals were sent to the laboratories as coded aliquots from Radian Corporation (Austin, TX). Chemicals were tested in cultured Chinese hamster ovary (CHO) cells for induction of sister chromatid exchanges (SCEs) and chromosomal aberrations both in the presence and absence of Aroclor 1254-induced male Sprague Dawley rat liver S9 and cofactor mix. Cultures were handled under gold lights to prevent photolysis of bromodeoxyuridine (BrdU)-substituted DNA. Each test consisted of concurrent solvent and positive controls and of at least three doses of the study chemical; the high dose was limited by toxicity or solubility but did not exceed 5 mg/ml.

In the SCE test without S9, CHO cells were incubated for 26 hours with the study chemical in McCoy's 5A medium supplemented with 10% fetal bovine serum, L-glutamine (2 mM), and antibiotics. BrdU was added 2 hours after culture initiation. After 26 hours, the medium containing the study chemical was removed and replaced with fresh medium plus BrdU and colcemid, and incubation was continued for 2 more hours. Cells were then harvested by mitotic shake-off, fixed, and stained with Hoechst 33258 and Giemsa. In the SCE test with S9, cells were incubated with the chemical, serum-free medium, and S9 for 2 hours. The medium was then removed and replaced with medium containing BrdU and no study chemical; incubation proceeded for an additional 26 hours, with colcemid present for the final 2 hours. Harvesting and staining were the same as for cells treated without S9.

In the chromosomal aberration test without S9, cells were incubated in McCoy's 5A medium with the study chemical for 8 hours; colcemid was added, and incubation was continued for 2 hours. The cells were then harvested by mitotic shake-off, fixed, and stained with Giemsa. For the chromosomal aberration test with S9, cells were treated with the study chemical and S9 for 2 hours, after which the treatment medium was removed and the cells were incubated for 10 hours in fresh medium, with colcemid present for the final 2 hours. Cells were harvested in the same manner as for the treatment without S9.

For the SCE test, if significant chemical-induced cell cycle delay was seen, incubation time was lengthened to ensure a sufficient number of scorable cells. The harvest time for the chromosomal aberration test was based on the cell cycle information obtained in the SCE test; if cell cycle delay was anticipated, the incubation period was extended approximately 5 hours.

Cells were selected for scoring on the basis of good morphology and completeness of karyotype (21 ± 2 chromosomes). All slides were scored blind, and those from a single test were read by the same person. For the SCE test, 50 second-division metaphase cells were usually scored for frequency of SCEs per cell from each dose; 100 first-division metaphase cells were scored at each dose for the chromosomal aberration test. Classes of aberrations included simple (breaks and terminal deletions), complex (rearrangements and translocations), and other (pulverized cells, despiralized chromosomes, and cells containing 10 or more aberrations).

Statistical analyses were conducted on both the slopes of the dose-response curves and the individual dose points. An SCE frequency 20% above the concurrent solvent control value was chosen as a statistically conservative positive response. The probability of this level of difference occurring by chance at one dose point is less than 0.01; the probability for such a chance occurrence at two dose points is less than 0.001. Chromosomal aberration data are presented as percentage of cells with aberrations. As with SCEs, both the dose-response curve and individual dose points were statistically analyzed. A statistically significant (P < 0.003) trend test or a significantly increased dose point (P < 0.05) was sufficient to indicate a chemical effect.

APPENDIX J. GENETIC TOXICOLOGY

RESULTS

Toluene, within a dose range of 10-1,000 µg/plate, did not induce reverse gene mutations in four strains of S. typhimurium (TA98, TA100, TA1535, or TA1537) when tested in a preincubation protocol in the presence or absence of Aroclor 1254-induced male Sprague Dawley rat or Syrian hamster liver S9 (Haworth et al., 1983; Table J1). In the mouse lymphoma assay for induction of Tft resistance in L5178Y/TK cells, toluene was positive in trials conducted with and without Aroclor 1254-induced male F344 rat liver S9 (McGregor et al., 1988; Table J2); significant responses were noted at doses of 200 µg/ml and above, which, in all but one trial, represented the highest nonlethal dose tested. Despite the statistically positive, reproducible responses observed in this assay, the overall conclusion was judged to be equivocal because the presence of a toluene/water emulsion could not be ruled out conclusively, therefore leaving a question of whether acceptable dose levels had been achieved in this assay as per the study criteria set forth in McGregor et al. (1988). In cytogenetic tests with cultured CHO cells, toluene did not induce SCEs (Table J3) or chromosomal aberrations (Table J4) when tested with doses up to 1,600 µg/ml in the presence or absence of Aroclor 1254-induced male Sprague Dawley rat liver S9; no induction of cell cycle delay, necessitating delayed harvest, was noted at any of the nonlethal doses tested.

TABLE J1. MUTAGENICITY OF TOLUENE IN SALMONELLA TYPHIMURIUM (a)

Revertants/Plate (b) Strain Dose (µg/plate) + 10% S9 (hamster) - 89 +10% S9 (rat) Trial 1 Trial 2 Trial 3 Trial I Trial 2 Trial 1 Trial 2 **TA100** 0 86 ± 6.8 99 ± 5.2 89 ± 5.5 104 ± 5.3 115 ± 8.7 92 ± 3.8 113 ± 2.0 96 ± 88 ± 10 98 ± 11.2 88 ± 4.3 47 ± 3.8 85 ± 5.0 6.0 2.4 108 ± 2.5 99 ± 12.7 53 ± 1.8 90 ± 117 ± 12.5 33.3 110 ± 23.8 99 ± 4.6 112 ± 4.5 6.8 102 ± 7.5 167 ± 51.1 58 ± 3.2 118 ± 15.6 84 ± 2.3 124 ± 5.5 100 94 ± 6.7 226 ± 52.3 114 ± 18.3 333.3 93 ± 4.2 60 ± 4.7 94 ± 2.7 100 ± 1.9 96 ± 7.4 1,000 (c) $71 \pm$ $2.1 (c) 90 \pm$ $3.8 (c) 51 \pm 3.2$ 86 ± 9.3 69 ± 10.8 103 ± 10.4 (c) 95 ± 14.2 Trial summary Negative Positive Negative Negative Negative Negative Negative 460 ± 15.3 Positive control (d) 519 ± 14.7 625 ± 28.8 494 ± 8.4 1.507 ± 41.2 963 ± 104.3 604 ± 39.5 - S9 +10% S9 (hamster) +10% S9 (rat) Trial 1 Trial 2 Trial 1 Trial 2 Trial 1 Trial 2 **TA1535** 0 20 ± 2.6 20 ± 13 ± 1.0 10 ± 0.3 6 ± 1.3 13 ± 10 24 ± 17 ± 12 ± 11 ± 9 ± 0.6 3 1 2.7 8 ± 4.3 3.2 2.0 33.3 22 ± 3.9 $22 \pm$ 3.9 8 ± 0.9 11 ± 2.5 9 ± 2.0 10 ± 1.5 100 16 ± 2.3 14 ± 2.2 10 ± 2.5 10 ± 1.2 10 ± 3.1 11 ± 1.5 21 ± 2.6 12 ± 9 ± 8 ± 333.3 19 ± 8 ± 2.6 2.2 4.4 3.5 1.5 1,000 (c) $12 \pm$ 0.9 (c) $14 \pm$ 1.7 11 ± 3.5 $(c)7 \pm$ 0.9 8 ± 2.7 $(c) 9 \pm$ 1.8 Trial summary Negative Negative Negative Negative Negative Negative Positive control (d) 512 ± 18.9 444 ± 23.0 374 ± 16.0 268 ± 34.8 389 ± 11.4 305 ± 3.2 7 ± TA1537 n 1.0 8 ± 1.0 16 ± 2.2 12 ± 2.7 14 ± 1.2 14 ± 1.9 10 7 ± 8 ± 11 ± 13 ± 1.0 0.9 19 ± 4.0 1.9 16 ± 4.8 2.0 12 ± 33.3 6 ± 8 + 19 ± 1.9 9 + 3.0 14 ± 0.3 2.9 1.2 3.5 100 10 ± 9 ± 3.0 20 ± 2.7 14 ± 3.2 7 ± 1.0 10 ± 1.2 1.3 333.3 7 ± 9 ± 2.0 15 ± 18 ± 5.5 7 ± 1.2 11 ± 2.0 1.0 1.5 1,000 7 ± 1.9 $(c)6\pm$ 2.3 17 ± 1.8 $(c) 2 \pm$ 1.0 5 ± 0.9 (c)9 ± 2.2 Negative Trial summary Negative Negative Negative Negative Negative Positive control (d) 227 ± 43.8 710 ± 64.7 556 ± 24.3 244 ± 5.8 365 ± 19.7 158 ± 10.6 **TA98** 0 17 +2.4 27 ± 3.1 $25 \pm$ 0.9 $37 \pm$ 4.7 27 ± 3.5 35 ± 2.3 10 22 ± 3.0 28 ± 0.3 26 ± 5.9 33 ± 1.3 24 ± 1.8 35 ± 3.4 41 ± 2.9 21 ± 25 ± 23 ± 37 ± 33.3 3.4 0.3 29 ± 6.4 1.8 3.7 19 ± 24 ± 34 ± 23 ± 28 ± 100 5.2 5.9 28 ± 0.71.5 3.8 1.5 26 ± 333.3 22 ± 3.3 25 ± 3.4 4.1 39 ± 4.1 17 ± 0.7 32 ± 4.7 1,000 (c) 20 ± 2.6 23 ± (c) 29 ± 16 ± 0.7 (c) $19 \pm$ 2.9 29 ± 1.5 2.2 0.3 Negative Negative Negative Trial summary Negative Negative Negative $1,292 \pm 27.6$ $1,258 \pm 130.5$ 288 ± 3.0 Positive control (d) 645 ± 7.4 636 ± 3.4 428 ± 24.2

⁽a) Study performed at SRI International. The detailed protocol is presented in Haworth et al. (1983). Cells and study compound or solvent (dimethyl sulfoxide) were incubated in the absence of exogenous metabolic activation (-S9) or with Aroclor 1254-induced S9 from male Syrian hamster liver or male Sprague Dawley rat liver. High dose was limited by toxicity or solubility but did not exceed 10 mg/plate; 0 µg/plate dose is the solvent control.

⁽b) Revertants are presented as mean \pm standard error from three plates.

⁽c) Slight toxicity

⁽d) Positive control; 2-aminoanthracene was used on all strains in the presence of S9. In the absence of metabolic activation, 4-nitro-o-phenylenediamine was used with TA98, sodium azide was used with TA100 and TA1535, and 9-aminoacridine was used with TA1537.

TABLE J2. INDUCTION OF TRIFLUOROTHYMIDINE RESISTANCE BY TOLUENE IN MOUSE L5178Y LYMPHOMA CELLS (a,b)

Compound	Concentration (µg/ml)	Cloning Efficiency (percent)	Relative Total Growth (percent)	Tft-Resistant Cells	Mutant Fraction (c)
- S9					
Trial 1					
Dimethyl sulfoxide (d)		65.0 ± 5.4	100.0 ± 14.8	120.8 ± 12.9	62.5 ± 4.7
Toluene	31.25 62.5 125 250 500	$\begin{array}{ccc} 69.0 \pm & 3.0 \\ 59.5 \pm & 2.5 \\ 66.0 \pm & 3.0 \\ 66.5 \pm & 6.5 \\ \text{Lethal} \end{array}$	$74.5 \pm 0.5 \\ 66.5 \pm 0.5 \\ 91.0 \pm 20.0 \\ 28.0 \pm 0.0$	129.0 ± 18.0 122.0 ± 28.0 124.5 ± 7.5 228.0 ± 30.0	62.0 ± 6.0 68.0 ± 13.0 63.0 ± 1.0 $(e) 114.5 \pm 3.5$
Methyl methanesulfonat	e 15	37.5 ± 3.5	38.5 ± 9.5	446.5 ± 29.5	(e) 401.0 ± 14.0
Trial 2					
Dimethyl sulfoxide (d)		74.8 ± 1.3	100.3 ± 7.8	126.0 ± 15.0	56.3 ± 6.6
Toluene (f)	50 100 200 300	75.7 ± 9.8 75.7 ± 3.3 82.0 ± 11.5 Lethal	$ \begin{array}{r} 104.3 \pm & 7.5 \\ 95.7 \pm & 9.7 \\ 60.3 \pm & 3.7 \end{array} $	138.3 ± 10.9 130.7 ± 18.5 209.0 ± 22.9	61.7 ± 4.7 57.3 ± 6.8 (e) 86.7 ± 6.8
Methyl methanesulfonat	e 15	22.0 ± 2.0	13.5 ± 1.5	304.0 ± 17.0	(e) 462.5 ± 19.5
Trial 3					
Dimethyl sulfoxide (d)		67.8 ± 5.2	100.3 ± 4.8	119.3 ± 4.9	59.5 ± 3.4
Toluene	150 175 200 225 250 275	68.5 ± 11.5 70.0 ± 2.0 80.0 ± 4.0 62.5 ± 5.5 58.5 ± 5.5 Lethal	$\begin{array}{cccc} 66.5 \pm & 10.5 \\ 64.0 \pm & 11.0 \\ 41.0 \pm & 9.0 \\ 21.0 \pm & 5.0 \\ 13.5 \pm & 5.5 \end{array}$	157.0 ± 20.0 168.0 ± 38.0 243.0 ± 33.0 347.5 ± 80.5 471.5 ± 76.5	77.0 ± 3.0 81.0 ± 21.0 (e) 103.0 ± 19.0 (e) 184.0 ± 27.0 (e) 274.5 ± 67.5
Methyl methanesulfonat	e 15	23.5 ± 3.5	16.5 ± 2.5	303.5 ± 48.5	(e) 428.0 ± 7.0
S9 (g)					
Trial 1					
Dimethyl sulfoxide (d)		95.5 ± 8.2	100.0 ± 1.7	118.0 ± 3.9	41.8 ± 2.4
Toluen <i>e</i>	6.25 12.5 25 50 100 200	$\begin{array}{cccc} 91.0 \pm & 6.0 \\ 92.5 \pm & 3.5 \\ 84.0 \pm & 6.0 \\ 94.0 \pm & 17.0 \\ 88.0 \pm & 3.0 \\ 74.5 \pm & 3.5 \end{array}$	$\begin{array}{cccc} 100.0 \pm & 8.0 \\ 107.0 \pm & 5.0 \\ 82.5 \pm & 4.5 \\ 98.5 \pm & 5.5 \\ 92.0 \pm & 9.0 \\ 40.5 \pm & 1.5 \end{array}$	130.0 ± 15.0 140.5 ± 13.5 127.0 ± 5.0 126.5 ± 15.5 123.0 ± 17.0 162.0 ± 12.0	48.0 ± 9.0 50.5 ± 2.5 50.5 ± 1.5 45.5 ± 2.5 47.0 ± 8.0 (e) 72.5 ± 8.5
Methylcholanthrene	2.5	69.0 ± 10.0	83.5 ± 6.5	347.0 ± 15.0	(e) 169.5 ± 17.5

TABLE J2. INDUCTION OF TRIFLUOROTHYMIDINE RESISTANCE BY TOLUENE IN MOUSE L5178Y LYMPHOMA CELLS (Continued)

Compound	Concentration (µg/ml)			Tft-Resistant Cells	Mutant Fraction (c)		
S9 (Continued)	<u> </u>						
Trial 2							
Dimethyl sulfoxide		63.0 ± 2.0	100.0 ± 4.0	128.5 ± 9.5	67.5 ± 2.5		
Toluene	125 150 175 200 (h) 225 250	65.5 ± 0.5 62.0 ± 2.0 53.0 ± 3.0 47.5 ± 4.5 45.5 ± 4.5 Lethal	$76.5 \pm 0.5 73.5 \pm 2.5 58.5 \pm 0.5 35.5 \pm 1.5 18.0 \pm 0.0 $	$\begin{array}{cccc} 134.5 \pm & 16.5 \\ 127.0 \pm & 5.0 \\ 118.5 \pm & 12.5 \\ 137.5 \pm & 1.5 \\ 189.5 \pm & 27.5 \\ \end{array}$	68.5 ± 7.8 68.0 ± 0.0 75.0 ± 4.0 98.5 ± 8.8 (e) 138.0 \pm 6.0		
Methylcholanthrene	2.5	30.0 ± 3.0	27.0 ± 1.0	570.5 ± 22.5	(e) 635.0 ± 42.0		

⁽a) Study performed at Inveresk Research International. The experimental protocol is presented in detail by McGregor et al. (1988) and follows the basic format of Clive et al. (1979). The highest dose of study compound is determined by solubility or toxicity and may not exceed 5 mg/ml. All doses are tested in duplicate, unless otherwise indicated; the average for the tests is presented in the table. Cells $(6 \times 10^5/\text{ml})$ were treated for 4 hours at 37° C in medium, washed, resuspended in medium, and incubated for 48 hours at 37° C. After expression, 3×10^6 cells were plated in medium and soft agar supplemented with trifluorothymidine (Tft) for selection of Tft-resistant cells, and 600 cells were plated in nonselective medium and soft agar to determine the cloning efficiency.

⁽b) Mean \pm standard error from replicate trials of approximately 1×10^6 cells each. All data are evaluated statistically for both trend and peak response (P<0.05 for at least one of the three highest dose sets). Both responses must be significantly (P<0.05) positive for a chemical to be considered capable of inducing Tft resistance. If only one of these responses is significant, the call is "equivocal"; the absence of both trend and peak response results in a "negative" call.

⁽c) Mutant fraction (frequency) is a ratio of the Tft-resistant cells to the cloning efficiency, divided by 3 (to arrive at MF per 1×10^6 cells treated); MF = mutant fraction.

⁽d) Data presented are the results of four tests.

⁽e) Significant positive response; occurs when the relative mutant fraction (average MF of treated culture/average MF of solvent control) is greater than or equal to 1.6.

⁽f) Data presented are the results of three tests.

⁽g) Tests conducted with metabolic activation were performed as described in (a) except that S9, prepared from the liver of Aroclor 1254-induced F344 rats, was added at the same time as the study chemical and/or solvent.

⁽h) Precipitation occurred at this concentration.

TABLE J3. INDUCTION OF SISTER CHROMATID EXCHANGES IN CHINESE HAMSTER OVARY CELLS BY TOLUENE (a)

Compound	Dose (µg/ml)	Total Cells	No. of Chromo- somes	No. of SCEs	SCEs/ Chromo- some	SCEs/ Cell	Hours in BrdU	Relative SCEs/Cell (percent) (b)
-S9 (c)								
Trial 1Summary: Nega	tive							
Dimethyl sulfoxide		50	1,051	490	0.47	9.8	27.0	
Toluene	50 160 500 1,600 5,000	50 50 50 0	1,050 1,045 1,034	423 492 496	0.40 0.47 0.48	8.5 9.8 9.9	27.0 27.0 27.0	86.7 100.0 101.0
Mitomycin C	0 0.01	50 10	1,050 210	753 572	0.72 2.72	15.1 57.2	27.0 27.0	154.1 583.7
Trial 2Summary: Nega	tive							
Dimethyl sulfoxide		50	1,052	436	0.41	8.7	26.5	
Toluene	100 200 300 400 4,000 5,000	50 50 50 0 0	1,047 1,044 1,038	460 426 474	0.44 0.41 0.46	9.2 8.5 9.5	26.5 26.5 26.5	105.7 97.7 109.2
Mitomycin C	0 0.01	50 10	1,047 210	632 446	0.60 2.12	12.6 44.6	26.0 26.5	144.8 512.6
+ S9 (d)								
Trial 1Summary: Nega	tive							
Dimethyl sulfoxide		50	1,051	436	0.41	8.7	26.5	
Toluene	16 50 160 500 1,600	50 50 50 22 0	1,046 1,045 1,045 459	433 416 457 205	0.41 0.40 0.44 0.45	8.7 8.3 9.1 9.3	26.5 26.5 26.5 26.5	100.0 95.4 104.6 106.9
Cyclophosphamide	0.3 2	50 10	1,042 211	759 480	0.73 2.27	15.2 48.0	26.5 26.5	174.7 551.7
Trial 2Summary: Nega	tive							
Dimethyl sulfoxide		50	1,044	413	0.4	8.3	26.0	
Toluene	50 100 250 500	50 50 50 0	1,044 1,046 1,049	396 413 413	0.38 0.39 0.39	7.9 8.3 8.3	26.0 26.0 26.0	95.2 100.0 100.0
Cyclophosphamide	0.3 2	50 10	1,050 209	605 414	0.58 1.98	12.1 41.4	26.0 26.0	145.8 498.8

TABLE J3. INDUCTION OF SISTER CHROMATID EXCHANGES IN CHINESE HAMSTER OVARY CELLS BY TOLUENE (Continued)

(a) Study performed at Environmental Health Research and Testing, Inc. SCE = sister chromatid exchange; BrdU = bromode-oxyuridine. A detailed description of the SCE protocol is presented by Galloway et al. (1985, 1987). Briefly, Chinese hamster ovary cells were incubated with study compound or solvent as described in (c) and (d) below and cultured for sufficient time to reach second metaphase division. Cells were then collected by mitotic shake-off, fixed, air dried, and stained.

(b) SCEs/cell of culture exposed to study chemical relative to those of culture exposed to solvent

(d) In the presence of S9, cells were incubated with study compound or solvent for 2 hours at 37°C. Cells were then washed, and medium containing BrdU was added. Cells were incubated for a further 26 hours, with colcemid present for the final 2-3 hours. S9 was from the liver of Aroclor 1254-induced male Sprague Dawley rats.

⁽c) In the absence of S9, Chinese hamster ovary cells were incubated with study compound or solvent for 2 hours at 37° C. Then BrdU was added, and incubation was continued for 24 hours. Cells were washed, fresh medium containing BrdU and colcemid was added, and incubation was continued for 2-3 hours.

TABLE J4. INDUCTION OF CHROMOSOMAL ABERRATIONS IN CHINESE HAMSTER OVARY CELLS BY TOLUENE (a)

			-S9 (b)					+ S9 (c)		
	Oose g/ml)	Total Cells	No. of Abs	Abs/ Cell	Percent Cells with Abs	Dose (µg/ml)	Total Celis	No. of Abs	Abs/ Cell	Percent Cells with Abs
Harvest ti	ime: 12 h	1				Harvest time: 13	3.3 h			
Dimetl	hyl sulfor	tide				Dimethyl sul	foxide			
	•	100	0	0	0.0	•	100	1	0.01	1.0
Toluer	ne					Toluene				
	50	100	1	0.01	1.0	50	100	3	0.03	3.0
1	160	100	2	0.02	2.1	60	100	2	0.02	2.0
5	500	100	1	0.01	1.0	500	100	1	0.01	1.0
1,6	600	100	3	0.03	3.0	1,600	100	4	0.04	4.0
	Sumr	nary: Ne	gative			Sur	nmary: Ne	gative		
Mitom	nyein C					Cyclophospha	amide			
	0.125	100	5	0.05	5.0	15	100	10	0.10	10.0
	0.25	100	27	0.27	19.0	50	50	28	0.56	48.0

⁽a) Study performed at Environmental Health Research and Testing, Inc. Abs = aberrations. A detailed presentation of the technique for detecting chromosomal aberrations is found in Galloway et al. (1985, 1987). Briefly, Chinese hamster ovary cells were incubated with study compound or solvent as indicated in (b) and (c). Cells were arrested in first metaphase by addition of colcemid and harvested by mitotic shake-off, fixed, and stained in 6% Giemsa.

⁽b) In the absence of S9, cells were incubated with study compound or solvent for 8-10 hours at 37° C. Cells were then washed, and fresh medium containing colcemid was added for an additional 2-3 hours followed by harvest.

⁽c) In the presence of S9, cells were incubated with study compound or solvent for 2 hours at 37°C. Cells were then washed, medium was added, and incubation was continued for 8-10 hours. Colcemid was added for the last 2-3 hours of incubation before harvest. S9 was from the liver of Aroclor 1254-induced male Sprague Dawley rats.

APPENDIX K

AUDIT SUMMARY

APPENDIX K. AUDIT SUMMARY

The pathology specimens, experimental data, study documents, and the draft NTP Technical Report No. 371 for the 2-year studies of toluene in rats and mice were audited for the National Institute of Environmental Health Sciences (NIEHS) at the National Toxicology Program (NTP) Archives by quality assurance, resource-support contractors. The audit included review of:

- (1) All records concerning animal receipt, quarantine, randomization, and disposition prior to study start.
- (2) All inlife records including protocol, correspondence, animal husbandry, room and exposurechamber environmental conditions, dosing, external masses, mortality, animal identification, and serology.
- (3) Body weight and clinical observation data; all data were scanned before individual data for a random 10% sample of animals in each study group were reviewed in detail.
- (4) All chemistry records.
- (5) All postmortem records for individual animals concerning date of death, disposition code, condition code, tissue accountability, correlation of masses or clinical signs recorded at or near the last inlife observation with gross observations and microscopic diagnoses, data entry discrepancies on necropsy record forms, and correlation between gross observations and microscopic diagnoses.
- (6) All wet tissue bags for inventory and residual wet tissues from a random 20% sample of animals in each study group, plus other relevant cases, to evaluate the integrity of individual animal identity and the thoroughness of necropsy and trimming procedure performance.
- (7) Blocks and slides of tissues from a random 20% sample of animals from each study group, plus animals with less than complete or correct identification, to examine for proper match, inventory, and preservation.
- (8) All microscopic diagnoses for a random 20% sample of animals, plus 100% of the changes in diagnoses made to preliminary pathology tables, to verify their incorporation into the final pathology tables.
- (9) The extent of correlation between the data, factual information, and procedures for the 2-year studies as presented in the draft Technical Report and the study records available at the NTP Archives.

Procedures and events for the exposure phase of the studies were documented adequately by the archival records, with the exception that some or all records for method of randomization, disposition of surplus animals and study chemical, frequency of feeding, frequency of rack changes and cleaning of inner chamber surfaces, room light cycle, last day of dosing for interim-kill animals, and chemical use log were not available at the Archives. Records documented that exposure concentrations were generated, monitored, and administered properly. Some body weight fluctuations possibly occurred when animal numbers and weight data were confused for 51/53 high dose male rats on one occasion (week 52) and entered into the computer, resulting in an apparent weight loss for these animals. Recalculation of approximately 10% of the group mean body weight values in the Technical Report showed 21/25 for rats and 22/24 for mice to be correct; differences ranged from 1.9% to 8.8%. All external masses observed inlife were correlated with masses noted at necropsy for both rats and mice. The disposition code and date of death recorded at necropsy for each unscheduled-death animal (185 rats and 73 mice) had matching entries in the inlife records, except for the dates of death for 2 mice, which had no effect on survival values given in the Technical Report.

Individual animal identifiers (ear tags for rats and toe clips for mice) were present and correct in the residual tissue bags for 62/69 rats and 59/70 mice examined. Review of the entire data trail for the 7 rats and 11 mice with less than complete and correct identifiers indicated that the integrity of their individual animal identity had been maintained, but the absence of ear tags and toe clips had not been documented. A total of five untrimmed potential lesions were found in the wet tissues of 69 rats examined and nine in the wet tissues of 70 mice; none involved target organs. Intestinal segments

were not completely opened for 24/69 rats and 38/70 mice, and the stomach was partially opened in 13 rats; however, no potential lesions were evident by external examination. Gross observations made at necropsy were correlated with microscopic diagnoses. Tissue blocks and slides matched each other properly. All post-Pathology Working Group changes in diagnoses had been incorporated into the final pathology tables.

Full details about these and other audit findings are presented in the audit reports that are on file at NIEHS. This summary describes the extent to which the data and factual information presented in the Technical Report for the 2-year inhalation studies of toluene are supported by the records at the NTP Archives.