# 20000 814 148

AD

ELECTE

G

# RESEARCH AND DEVELOPMENT ON INHALATION TOXICOLOGIC EVALUATION OF RED THOSPHORUS/BUTYL RUBBER COMBUSTION PRODUCTS

PHASE II REPORT

CATHERINE ARAMYI

DEC.MBER 1983

AD-A158 32

## Supported by

U.S. ARMY MEDICAL RESEARCH AND DEVELOPMENT COMMAND Fort Detrick, Frederick, Maryland 21701

Contract No. DA1017-82-C-2121

IIT Research Institute, Lie Sciences Research Division 10 West 35th Street, Chicago, Illinois 60616

Contracting Officer's Technics: Representative

MARY C. HENRY, Ph.D.

Health Effects Research Division U.S. Army Medical Bloengineering Research and Development Laboratory Fort Detrick, Frederick, Maryland 21701

Approved for public release; distribution unlimited.

The findings in this report are not to be construed as an official Department of the Army position unless so designated by other - authorized documents.

8.5

REPORT DOCUMEN	TATION PAGE	READ INSTRUCT	
REPORT NUMBER	AD-ALLS	BEFORE COMPLETI	
Research and Development on Evaluation of Red Phosphorus		5. TYPE OF REPORT & PERI Phase II: March 19 July 198	ic3 to
Combustion Products		6. PERFORMING ORG. REPO L06139 Phase 11	
AUTHOR(2)	وه به ۲۹۹۳ می با این از این	B. CONTRACT OR GRANT NU	MBER(a)
Catherine Aranyi		DAMD17-82-C-2121	
PERFORMING ORGANIZATION NAME AND IIT Research Institute 10 West 35th Street Chicago, 11!inois 60616	DADDRESS	10. PROGRAM ELEMENT. PRO AREA & WORK UNIT NUM 62777 A-3E1627774878.CA	
CONTROLLING OFFICE NAME AND ADD	DRESS	12. REPORT DATE	·
U.S. Army Medical Research	and Development Command		
Development Laboratory	21701	13. NUMBER OF PAGES	
Fort Detrick, Frederick, MD 4 MonitoRing agency name a abbres US Army Medical Bioenginner		15. SECURITY CLASS (of the	e report)
Development Laboratory Fort Detrick, Frederick, MD	21701	U 15. DECLASSIFICATION DO SCHEDULE	WNGRADING
5. DISTRIBUTION STATEMENT (of this Rep		1	
pproved for public release;		•	
	distribution unlimited		
percyed for public release; 7. DISTRIBUTION STATEMENT (of the ober	distribution unlimited		
	distribution unlimited		
7. DISTRIBUTION STATEMENT (of the ober	distribution unlimited		
7. DISTRIBUTION STATEMENT (of the ober	distribution unlimited	om Report)	
7. DISTRIBUTION STATEMENT (of the ober 8. SUPPLEMENTARY NOTES 9. KEY WORDS (Continue on a Ademosine trichospic	distribution unlimited rent entered in Block 20, it different fro estern and identify by block number ' countsy inh	om Report)	
7. DISTRIBUTION STATEMENT (of the ober 8. SUPPLEMENTARY NOTES 9. KEY WORDS (Continue on o Adenosine trichospic Aerosol Alveolar macrophage Bactericidal acti	distribution unlimited rere entered in Block 20, it different to excerv and identify by block number ' countsy Inh ular proteiny LC5 bustion products >Nec Osure duration Obs	) alation toxicology 0 concentration ropsy, curant	
7. DISTRIBUTION STATEMENT (of the ober 8. SUPPLEMENTARY NOTES	distribution unlimited rere entered in Block 20, it different to excerv and identify by block number ' countsy Inh ular proteiny LC5 bustion products >Nec Osure duration Obs	alation toxicology 0 concentration ropsy, curant, sphoric acid (co	nt inued)
7. DISTRIBUTION STATEMENT (of the ober 8. SUPPLEMENTARY NOTES Adenosine trichospic Aerosol Alveolar macrophage Bactericidal acti Body weight 0. ABSTRACT (Co Exploratory centrations showed highl haled 35S-K in the pulmc	estern and identify by block number rent entered in Block 20, if different fro rent entered in Block 20, if	<pre>&gt;&gt;&gt; Report) &gt;&gt;&gt; alation toxicology alation toxicology 0 concentration ropsy, curant sphoric acid (co -Dawley rats exposed 4 mg/l for 1 to 3 ha actericidal activity ecreases in total co chial lavage. Subse</pre>	to con- periods to in- il counts quent
7. DISTRIBUTION STATEMENT (of the ober 8. SUPPLEMENTARY NOTES Adenosine triphospic Aerosol Alveolar macrophage Bactericidal acti Body weight 0. ABSTRACI (EG Exploratory centrations showed highl haled 35S-K	esterv and identify by block number rent antered in Block 20, if different fro rent antered in Block 20, if different fro rent antered in Block 20, if different fro rent and identify by block number i countsy Inh ular proteiny LC5 bustion products >Nec bustion produc	alation toxicology alation toxicology 0 concentration ropsy, curant sphoric acid (co -Dawley rats exposed 4 mg/l for 1 to 3 he actericidal activity ecreases in total co chial lavage. Subse concentration is the d demonstrated that	to con- periods to in- ll counts quent deter-

#### UNCLASSIFIED

#### SECURITY CLASSIFICATION OF THIS PAGE (When Date Entered)

BLOCK 19; KEY WORDS (concluded)

Pulmonary free cells Radiolabeled bacteria Red phosphorus/butyl rubber Smcke Standard toxicology Statistical evaluation

# BLOCK 20: ABSTRACT (concluded)

studies with single exposures, as required in the contract, were not feasible in the range of aerosol concentrations physically attainable with the generators at the air flow rates required in our inhalation exposure system. When, in a follow-up multiple exposure study, rats innaled RP/BR aerosols for 1 hr/ day for 5 days at concentrations from 1.56 to 3.05 mg/l mortality ranged from 5 to 90 percent and the estimated LC50 value was 2.32 mg/l. When these data were contrasted with observations from two studies with 5 daily 4-hr, exposures to 0.35 or 0.99 mg/l of RP/BR the importance of exposure concentration over duration in terms of producing more pronounced effects at similar CxT values was evident again. Results of the definitive range finding studies designed on the basis of these exploratory experiments demonstrated that inhalation of 0.5 mg/1 RP/BR aerosol for 1 or 3.5 hr daily for 4 days, or for 3.5 hr 4 times weekly for 4 weeks produced decreases in body weights relative to control animals, but the exposures did not result in gross pathologic changes and the microscopic changes observed in lung tissue could not be considered compound-related. The changes found in pulmonary lavage parameters after 4 exposures were no longer present after the recovery period.



UNCLASSIFIED

SECURITY CLISSIFICATION OF THIS PAGE(When Ders Entered)

# RESEARCH AND DEVELOPMENT ON INHALATION TOXICOLOGIC EVALUATION OF RED PHOSPHORUS/BUTYL RUBBER COMBUSTION PRODUCTS

# PHASE II REPORT

CATHERINE ARANYI

# DECEMBER 1983

#### Supported by

# U.S. ARMY MEDICAL RESEARCH AND DEVELOPMENT COMMAND Fort Detrick, Frederick, Maryland 21701

Contract No. DAMD17-82-C-2121

IIT Research Institute, Life Sciences Research Division 10 West 35th Street, Chicago, Illinois 60616

Contracting Officer's Technical Representative

MARY C. HENRY, Ph.D.

Health Effects Research Division U.S. Army Medical Bloengineering Research and Development Laboratory Fort Detrick, Frederick, Maryland 21701

Approved for public release; distribution unlimited.

The findings in this report are not to be construed as an official Department of the Army position unless so designated by other authorized documents.

AD

# EXECUTIVE SUMMARY

The objective of these studies is to develop a data base for health hazard assessment of the effects produced by inhalation of combustion products from red phosphorus/hutyl rubber used as an obscurant smoke for troops and vehicles in factical and training environments. Laboratory rats exposed in inhalation chambers are used to provide a comprehensive definition of the biologic effects of red phosphorus smoke to mammalian systems under conditions which approximate the potential troop exposure. This report summarizes the Phase II range find: g studies.

In early exploratory experiments male and female Sprague-Dawley rats were exposed to various concentrations of RP/BR aerosols ranging from 0.5 to 3 mg/l for durations of 1- to 4-hr. Highly significant decreases in pulmonary bactericidal activity to inhaled  $^{-5}S-K$ .pneumoniae were found in the exposed rats relative to controls after single as well as multiple exposures. The higher exposure concentrations also produced significant decreases in total cell counts in the pulmonary free cells obtained by tracheobronchial lavage from the exposed rats relative to those collected from controls.

In subsequent range finding mortality studies male and female rats we a given single 1-hr exposures to 2.00, 2.22, 2.62, 3.09 and 3.15 mg/l of RP/BR aerosols and observed for 14 days. Only the exposures to  $\geq 2.62$  mg/l caused deaths. The maximum mortality resulting from a single 1-hr exposure to approximately 3 mg/l of RP/BR aerosol (maximum generator capacity) was 20 to 25 percent, while 2.62 mg/l resulted in 6 percent deaths. A single 4-hr exposure to 0.88 mg/1 with a CxT value similar to those in the 3.09 and 3.15 mg/1 1-hr studies caused no deaths thereby suggesting that exposure concentration is the determining factor rather than duration. These experiment: demonstrated that LC studies with sligle exposures at logarithmically spaced concentrations, were not feasible in the range of aerosol concentrations levels physically attainable with the generators at the air flow rates required in our inhalation exposure system.

In a follow-up multiple exposure study rats inhaled RP/BR aerosols for 1 hr daily on 5 consecutive days at concentrations of 1.56, 1.99, 2.49 and 3.05 mg/l. Mortality rates, mean survival times, body weights and overall clinical observations were made during the 5 exposure days and for 14-day observation period. Necropsies were done on all animals that died during the study and on survivors on Day 19. Mortality ranged from 5 to 90 percent decreases in survival times. The estimated LC : value was 2.32 mg/l with 1.99 and 2.73 mg/l confidence limits.

When rats received five daily 4-hr exposures to 0.35 or 0.99 mg/l of RP/BR aerosol only one died. Comparison of body weights and of clinical observations for the 1- and 4-hr studies showed markedly

adverse effects after the 1-hr and practically negligible ones after the 4-hr exposures. Thus the greater influence of exposure concentration on toxicity over duration was again demonstrated.

Based on the results of these range finding experiments the definitive studies were conducted on male and female rats using four daily exposures to 0.5 mg/l of RP/BR aerosol and comparing 1.0 and 3.5-hr exposure durations. Pulmonary free cell parameters and structural changes, as evaluated by gross necropsy and histopathology, were tested immediately following the last exposure and after a 14-day recovery period. In an additional study rats were exposed to 0.5 mg/l of RP/BR for 3.5-hr daily 4 days per week for 4 weeks. Gross pathologic observations were made after the last exposure and 2 weeks following the last exposure and regular body weight determinations.

No deaths occurred in any of the studies and generally no treatment-related clinical observations were seen. Statistical analysis of body weights showed that exposure duration was not a significant factor. When body weight changes were analyzed over the entire period of exposure and recovery, the 4-day as well as 4- week exposures produced weight decreases in male and female rats relative to controls that did not return to the normal control level within the recovery period.

Exposure duration did not have a significant effect on the pulmonary free cell parameters. Total and differential cell counts in the pulmonary lavage were not affected. Cellular ATP levels in the lavaged cells were significantly decreased when examined immediately after the fourth exposure, however recovery was complete after 14 days.

No compound-related gross pathologic lesions were observed during necropsies immediately following the last exposure or after a 14-day recovery period in either the 4-day or the 4-week exposure group.

Microscopic examination of the tissues immediately following the 4-day exposure and after a 2-week recovery period did not reveal any treatment- related changes in kidneys, trachea and nasal turbinates. However, lungs of several animals had focal areas of interstitial reaction with alveolar macrophages which may be treatment-related. These animals, however, also had moderate to marked lymphoid hyperplasia of pulmonary lymph nodes which suggests an infectious agent may have produced the pulmonary lesions.

# FOREWORD

This report, IITRI No. L06139, Phase ti Report describes studies conducted by the Life Sciences Division, IIT Research Institute for the Health Effects Research Division, U.S. Army Medical Bioengineering Research and Development Laboratory during the period of February through July 1983. The studies were carried out under Contract No. DAMD17-82-C-2121.

Catherine Aranyi served as Principal Investigator and James Fenters was Co-Investigator. Principal professional associates were Stanley Vana and Jeannie Bradof. Mr. Vana was responsible for the exposure facilities and the aerosol generation and Inhalation monitoring throughout the studies. Ms. Bradof was in charge of all pulmonary response studies and overall toxicologic observations. collection preparation and tissue for Necropsv procedures, histopathologic evaluation were under the supervision of Vladislava Rac and Carol Thompson. Percent phosphoric acid analysis in the filter-collected aerosol samples was performed under the supervision of Alan Snelson. Robert Gibbons, Consultant Blostatistician performed statistical analysis of the experimental data. Histopathologic evaluation of the collected tissue samples was performed by W. D. lverson, consultant pathologist from Experimental Pathology Laboratories, Herndon, VA.

Animal experiments were conducted according to the "Guide for the Care and Use of Laboratory Animals" (1978), DHEW Publication No. (NIH) 78-23 prepared by the Committee on Care and Use of Laboratory Animals of the Institute of Laboratory Animal Resources, National Research Council: regulations and standards of the Department of Agriculture and Public Law 91-579, "Laboratory Animal Weifare Act" (1970).

Citation of commercial organizations and trade names in this report does not constitute an official Department of the Army endorsement or approval of the products or services of these organizations.

# QUALITY ASSURANCE STATEMENT

Laboratory operations were inspected on May 16, 18, 19 and 24, June 3 and September 28, 1983. Final draft reports were audited on August 8 and December 9 to 14, 1983. Inspection and audits were performed by Josephine M. Reed, Julie McPhilips and Kirit Parikh.

The study was found to conform to lITRI Life Sciences Quality Assurance criteria developed to meet FDA Good Laboratory Practice Regulations (Fed. Reg. CFR, Part 58, 1978). Animal experiments were conducted according to the "Guide for the Care and Use of Laboratory Animals" (1978), DHEW Publication No. (NIH) 78-23 prepared by the Committee on Care and Use of Laboratory Animals of the Institute of Laboratory Animal Resources, National Research Council; regulations and standards of the Department of Agriculture and Public Law 91-579, "Laboratory Animal Welfare Act" (1970).

Raw data generated during the course of the study will be retained in the IITRI Life Sciences Archives.

/Joséphine M. Reed Supervisor, Quality Assurance

# TABLE OF CONTENTS

	Exec	u+1.	ve	Su	m m :	arv			_			_			_	-			_			-						1
																												1
	Forev Qual	1+1	۵۵	•	• r = 1			2 + -	• • +	•·		•+	•	. •	•	•	•	•	•	•	•	•	•	•	•	•	•	3
	Tabl	; i y o o o	4 (		+ a :	nte nte		211	01	Ch	.01		•	•	•	•	٠	•	•	•	•	•	٠	•	•	•	•	4
	List																											,
	List	or	10	יוטב	es	- '	•	•	•	•	•	٠	•'	•	٠	•	٠	•	٠	•	•	•	•	• •	• •	٠	•	7
	LIST	OT	ן <b>ד</b> ו	gu	r e	5.	•		•	•	•	٠	٠	•	•	•	•	٠	•	٠	٠	٠	•	•	•	•	•	8
1.	INTR	GDU	CT		•					_				•	_												*	0
••		000				• •		•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	9
11.	MATE	RIAL	LS	AN	DI	MET	ГНО	DDS	S	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•		•	•	à
	'A. Ai	nima	als	5	•				•	•	•	•			•	•	•	•	•		•			•	•	•	•	9
	B. Ge	enei	r a f	110	n a	anc	1 N	101	n I	to	r l	l n	q	of	11	า่อ่	Cł	n an	nbe	ər	At	h m d	osi	ph e	er	в.		10
	1.	. R	eđ	Ph	os	pho	orι	ıs,	/В	u†	v I		Řu	bbe	ər	(F	RP/	/BF	( {	•	•		•	·		•	•	10
,		. 11																										10
		. A																										11
	C. B																											11
		. Pi																	•	•	•	•	•	•		•	•	
	• •	• • •	-3,5	S-1																								11
	2	. Pi	u t e																		,							13
		. S																										-
	D. 51	• _0 : + = + !	i ai tei		al	Mo	/~ 1 + h		- 1 1 c	vy	ÿ	•	٠	•	•	•	•	•	•	•	*	•	•	•	. •	•	•	14
			וכין			en e		101	4 3		•	•	.•	•	٠	•	•	٠	•	. •	•	•	•	٠	•.	٠		15
11.	RESUL	LTS	•		•					•		•			•				•'	•			•			•	. •	16
	A. Pr	rel	l m 1	Inai	r v	St	ินต์	11.	es											•	•							16
•	1	. Pi	uln	non	arv	v E	lac	:+ (	er	í C	1.0	ta	Ē	Act	-İ v	11	ťv.						ż					16
	2	. Pi			arv	, - , F	re		°c	۰Ĩ	1	. –	•					-		Ţ								18
	B. R.		5 F	-	41.	7 '	Fy	, n 4	۰r	Im	Ar	+	້	•	•	•	•	•	•	•	•	•	•	*		•	•	10
	1	3 9 C	1 .	n i a	ج	"9 707		1 P 4	91 96	4 11		•••														ļ	•	18
,	2			115 F1 a	با د ا	~¥~	/ 3 U		3.3			•	•	•	•	•	. •	•	•	•	.*	•	•	•	•	•	•	10
	C D.	, Mi			ι <del>σ</del>	5.2 		131	11 11	a >	• .	•		٠	٠	٠	٠	۲	•	٠	٠	4	٠		•	•	•,	21
	C. De	9111 e.	111			9 T U		6.63	5	*	*	•	٠	*	٠	•	۰,	•	*	•	, •	٠	•	٠.	•,	•	٠	28
		. <u>ב</u> ,	кре	91 11	161	пта		Ue	35	Ϊâ	11 • • •	•	•		. •	٠	٠	• '	۰	٠	٠	٠	٠	٠	٠	•	٠	29
	- 4	. E)	крс	su		ΤC		( , ,	<b>B</b>	ĸ	Ae		05	01	٠	٠		•	•	٠	٠	٠	٠	٠	٠	٠	٠	
		Mo	r	ra I	1 T )	y a	1 T C		1	п	10	:8		UDS	ser	r V a	ITI	QI	15	٠	٠	٠	٠	٠	٠	٠	٠	31
	4.	Bo	ody	/ W(	elg	ghī	5	4	•	٠.	•	•	<b>'</b> •	•	•	٠	٠	٠	٠	•	٠	٠	٠	٠	٠	٠	·• .	
	5.	. Pi	11 1	nona	ary	y F	re	••	Ç	e I	T	Pa	ar:	ame	)†e	er s		•	٠	٠	٠	٠	٠	٠	٠	•	٠.	. 38
	5.	Gr																										
		. (	Dbs	ier \	va:	† I G	ns	<b>ک</b> ر ا	•		•	٠	٠		٠	•	٠	•	۰.	٠	٠	٠	۰.	•	٠	•	• .	38
17.	CONCL	1151		19												_											•	1. 0
• • •					• •	• •	•		•	•	•	•	•	• .	•	•	٠	٠	•	•	•	•	.*	•	•	•	•	42
	1 1750			: r		FD'									•					÷								1. *

Page

# TABLE OF CONTENTS (continued)

والتروي والمراجع والمراجع

	Page
APPENDIX	• 46 • 47
A-1. Mean Body Weights of Male and Female Rats Exposed One Time to RP/BR Aerosols and	• €
Monitored for 14 Days	. 48-49
Exposed Five Times to RP/BR Aerosol's and Monitored for 14 Days	• 50-52
During Four Daily Exposures to RP/BR Aerosol A-4. Mean Body Weights of Male and Female Rats	• 53 *
During Four Daily Exposures to RP/BR Aerosol and Followed by 14 Days of Recovery A-5. Mean Body Weights of Male and Female Rats	• 54-55
During Four Weeks of Exposure Four Days/Week tc RP/BR Aerosol and Two Weeks of Recovery	- 56 4
Pathology Report. Part 1: Gross Observations(IITRI) Four-day Inhalation Toxicity Study of RP/BR in Rats:	• 57
Pathology and Pulmonary Lavage (Study 76/77) Four-week inhalation Toxicity Study of RP/BR	· -
In Rats (Study 78)	• 67 • 70
Pathology and Pulmonary Lavage (Study 76/77) PERSONNEL SUPPORTED BY THIS PROJECT DURING THE	• 71
PHASE II STUDIES	• 112 3
DISTRIBUTION LIST	• 113

# LIST OF TABLES

Iable		
No.		<u>Page</u>
1.	Effect of Exposure to RP/BR Aerosol on the Pulmonary	
	Bactericidal Activity of Rats	17
2.	Effects of Exposure to RP/BR Aerosol on Free Cells	
7	Lavaged from the Lungs of Rats	19
<b>3</b> .	Mortality and Mean Survival Time of Rats after Single Exposures to RP/BR Aerosol	• •
4.		20
7.	Male and Female Rats after 5 Dally Exposures to	
	RP/BR Aerosol	-23
5.	RP/BR Aerosol Exposure Group Allocation for Study Nos. 76, 77	
•	and 78	30
6.	the second	
. 7	H <sub>3</sub> PO <sub>4</sub> for Study Nos. 76, 77 and 78	32
7.	Mean Body Weights of Male and Female Rats during Four Days of Exposure to RP/BR Aerosols (Study Nos. 76A	
		34
8.		. ۳
	RP/BR Aerosci Exposure and Recovery Days (Study Nos.	
	76B and 77B)	35-36
9.		•
	Four Weeks of Exposure Four Days/Week to RP/BR	
10.	Aerosols and Two Weeks of Recovery	37
	Pulmonary Free Cells Lavaged from the Lungs of Male	
	and Female Rats	39-40
11.	Overall Mean Values for Endpoint Parameters Heasured	<i></i>
	on Pulmonary Free Cells Lavaged from the Lungs of Rats	
	atter Exposure to RP/BR Aerosols	41

# LIST OF FIGURES

#### Elgure. Page 1. Mean Body Weights of Male and Female Rats Before and after a Single 1-Hour Exposure on Day 1 to RP/BR Aerosols 2.0 mg/l, 3.09 mg/l, or 3.15 mg/l. . . . 22 Mean Body Weight Changes in Male and Female Rats Ζ. During and after 5 Daily 1-hr Exposures to RP/BR Aerosols 1.99 mg/l or 3.05 mg/l . . . 25 3. Mean Body Weight Changes in Male and Female Rats During and after 5 Daily 1-hr Exposures to RP/BR Aerosols 1.56 mg/l, or 2.49 mg/l, or to Filtered Alr. 26 Mean Body Weight Changes in Male and Female Rats 4. During and after 5 Daily 4-hr Exposures to RP/BR Aerosols 0.99 mg/l, or 0.35 mg/l, or to Filtered Air. 27

# LA INTRODUCTION

As part of an oversil concern for personnel health and safety, U.S. Army Medical Research and Development Command Is seeking the to evaluate the effects produced by inhalation of combustion products from red phosphorus/buty! rubber used as an obscurant smoke for troops and vehicles in tactical and araining Laboratory rats, exposed in chambers will be used to environments. provide a comprehensive definition of the biologic effects of red phosphorus smoke to mammalian systems andur conditions which approximate the potential troop exposure. The approach to this research includes range finding acute studies to determine lethal concentrations and influence of exposure duration on mortality; studies to define time-concentration repeated exposure relationships as well as threshold levels, healing, and adaptation in biologic , eactions; and a subchreate exposure study with a recovery and observation period after the experimental exposure. The principal biologic response criter a to be monitored include overt toxic signs, clinical chemistry and hematology, histopathology, alveolar macrophage defense functions, pulmonary bactericidal activity and neurobehavioral activity. The research project is set up to proceed in a phased manner. The Phase II biologic range finding studies summarized in this report were to form the basis for the selection of exposure conducted concentration levels, durations and frequencies to be used in the Phase III Intermediate-term multiple exposures.

# 11. MATERIALS AND METHODS

#### A. /.nimais

Male and female Sprague Dawley rats, 3- to 4-weeks-old were obtained from arlan/Sprague-Dawley, Inc., Madison, WI. The rats were housed 3 per cage during the quarantine period in plastic cages with water bottles. Following test animal selection the rats were housed individually in stainless steel inhalation cages containing four compartments, each compartment measuring 18.4 x 16.5 x 15.9 cm. The cages were used for holding as well as exporte throughout the studies. When attached to the racks the cages are equipped with an automatic drinking water distribution system and are suspended over excrement pans. For exposures the cages are removed from the racks. The animals were transferred weekly to clean cages and the deotized absorbing cage boards were changed twice per week.

The animal rooms were maintained on 12 - hr light/dark cycle, Temperature and relative humidity were regulated in such a manner that extreme fluctuations between the animal room and exposure chamber conditions would be avoided. Purina Certified Rodent Chow 5002 and water were available to the rats <u>ad libitum</u> except during the exposures.

Animals were randomized by sex to treatment groups using a constrained random process, stratified by weight, such that all groups tested were comparable in pretest body weight, but assignment of individual animal to groups was random. Each test animal was identified with a unique number attached through ear tags.

# B. Generation and Monitoring of the Chamber Atmosphere

1. <u>Red Phosphorus/Butyl Rubber (RP/BR)</u>

The test article, RP/BR softened with hexane and prepackaged in 0.75 in diameter 4.5 in long stainless steel feed cylinders (billets) with end caps was supplied by the Sponsor through Oak Ridge National Laboratories (ORNL) and was stored at ambient temperature. A record of the test article was maintained which included date of receipt with identification numbers of each cylinder and the date and study number for which it was used.

#### 2. Inhalation Exposure Eacility

1

ľ

i

The inhalation exposure facilities and methods have been described in detail in the Phase I Report (1). Its major components are the inhalation exposure chambers with air flow and differential pressure controls; the conditioned air supply and chamber air exhaust systems; the red phosphorus/butyl rubber combustion generators and the aerosol monitoring systems.

Supply air to the inhalation exposure laboratory was preconditioned by passing through prefilters, charcoal filters and an air conditioning unit. Automatically controlled heating and humidifying units built into the supply air system maintained the air temperature and relative humidity (RH) at the specified ranges of 24 to 27°C and 40 to 60 percent RH. Prior to entering the exposure chambers the conditioned air was further filtered through a fibergiass coarse filter, a charcoal bed and HEPA filter.

The combined exhaust air from the exposure chambers was filtered through a single-housing, 30-element coalescent filter and exhausted above the roof of the building. The control chamber exhaust was independent from that of the experimental test chambers to avoid potential contamination from the test aerosol. The negative pressure and airflow rate in the chambers and exhaust filter loading were monitored by differential pressure gauges.

The aerosol was generated by specially designed hydraulic extrusion-combustion generators provided by the Govornment through CRNL. The generator operates by exerting hydraulic pressure on the RP/BR forcing it to extrude from an orifice into a burn chamber where it is ignited. The aerosol generated from the combustion products is transported directly into the exposure chamber inlet port. At a constant chamber airflow rate the concentration of the aerosol is a function of the extrusion rate of the RP/BR which is controlled by a precision hydraulic metering pump.

The rats were exposed to the test atmosphere in identical 1-m<sup>3</sup>sized stainless steel inhalation chambers operating at a total airflow rate of 500 L/min. Aerosol exposure and filtered air control chambers were located in separate laboratories.

# 3. <u>Aerosol Monitoring</u>

The RP/BR aerosol was monitored within each exposure chamber for mass concentration by gravimetric filter collection three times during the 1-hr exposures and four times during 3.5-hr exposure durations. In addition, mass concentration was monitored and output continuously recorded with light scattering photosensors. An integrated average of the photosensors was recorded simultaneously with the gravimetric filter sample collection. was recorded Aerosol particle size was determined with a Quartz Crystal Microbalance cascade impactor. The particle size was monitored in each chamber once on each exposure day. Determination of total phosphorus was conducted by spectrophotometric analys's of the collected filter samples on one filter per chamber per exposure week. All collected fliters were recorded and stored in sealed containers until submitted for chemical analysis. Oxygen levels in the chambers were spot checked during each experiment and were consistently 21 percent.

# C. Biological Endpoints

# 1. Pulmonary Bactericidal Activity to:355-K.oneumoniae

Aerosols of  $\lfloor^{35}S \rfloor Klebslella pneumonlag disseminated with a$ Retec X-70 disposable nebulizer were used for the bactericidalactivity assay, using a method previously described for mice and $adapted for rats (2). Radiolabeled K_ <u>pneumonlag</u> were grown in a$ medium in which the sulfate requirement of the bacteria was $provided by <math>\lfloor^{35}S \rfloor$  sodium sulfate. Before aerosolization, the bacteria were washed repeatedly and centrifuged for removal of unattached radiolabel. Bacterial counts were determined in a Petroff-Hausser counting chamber by dark-field microscopy and by the culture plate technique. Radioactive counts were measured in a Mark ill Liquid Scintillation System (Tracor Inc.). The ra In a glove for exposure airlocks the airlock proving impingers. to laborator passed first and subseque

3

2

Ŋ

Pu!mona the indivic simultaneous bacteria. radioactive which bacter erial aerosol exposure chamber installed of an 87-liter main compartment (adequate inat was accessible through appropriate e animals were moved. An additional small of the glove box for the nebulizers and radioactive and pathogenic exposure hazard all air exhausted from the chamber was n absolute filter placed within the glove box gh a HEPA filter located on the outside.

icidal activity was determined in the lungs of is from both exposed and control groups that aerosols of the viable radiolabeled if the viable bacterial counts to the h animal's lungs provided the rate at yed 3 hr after infection. Thus, where

ericidal Activity =  $\frac{1-R_3}{K_2}$  100

R<sup>3</sup> is lungs of i from the samu inhaling thu studies. o of bacterial to radioactive counts in the lual rats at 3 hr, and KO is an average determined os in the lungs of rats killed immediately after teria. This assay was used only in preliminary

# 2. Pulmonary Free Cells

Within 4 hr (0 hr) or 14 days after the last exposure, alveolar macrophages (AM) were obtained from designated animals by tracheobronchial lavage. The rats were weighed, killed with an overdose of sodium pentobarbital by intraperitoneal injection, and the lungs were lavaged through a blunted 18-gauge needle inserted inte an incision in the trachea with 9 consecutive 6-ml infusions of warm saline. The AM were collected from the lavage fluids by centrifugation and resuspended in Hanks' balanced salt solution (HBSS) after the lavage fluids were collected for protein Total cell counts were made in a hemocytometer. determination. For determination of the cellular distribution (i.e., percent of AM, polymorphonuclear leukocytes and lymphocytes), differential counts were made on cytocentrifuge preparations of cells fixed inmethanol and stained with Wright's stain.

Cellular adenosine triphosphate (ATP) levels were determined using a DuPont 760 Luminescence Blometer with the procedure recommended for the Instrument. The assay is based on the principle that when a microsample containing ATP is injected into a suitably buffered reaction mixture of luciferase and luciferin, the peak intensity of the resulting light flash is directly proportional to the concentration of ATP. ATP was extracted from the cells by dimethyl suifoxide (DMSO) from allquots of the cell suspension. The DMSO extracts were diluted with a specially prepared 0.01 M morpholinopropane suifonic acid (MOPS) buffer to overcome the quench effect of the high concentration of DMSO in the aqueous extract for the luciferase-luciferir reaction.

For determination of total cellular protein contert, allquots of the cell suspensions were treated with 1 percent sodium deoxycholate (SDC) and assayed by the Lowry method (3). Lavage fluids were assayed without SDC treatment. Total cellular protein values were given per  $10^{\circ}$  AM. ATP levels were expressed in femtograms (fg) of ATP per ug of total protein (experimentally determined), or per  $10^{\circ}$  AM (calculated from initial cell counts).

# 3. <u>Standard Toxicology</u>

<u>Mortality</u> was recorded daily and expressed as the percent of animals dying out of the total number of animals on test.

<u>Mean survival time</u> was estimated by the method of Horsfall (4), from the following equation:

$$MST = \frac{\Sigma(A \times B) + (d \times L)}{\Delta B}$$

where, A was the last day on which any individual rat was alive, B was the number of rats surviving A days, d was the last day of observation, L was the number of rats alive on day d, and n was the number in the experimental group.

<u>Clinical observations</u>: All animals were observed daily for survival, physical appearance, behavior and any pharmacologic and/or toxicologic signs. All observations were recorded on an individual test animal basis.

<u>Body weight:</u> Each animal was weighed using a Mettler PE 1600 balance with a special animal weighing mode at the Initiation of the study and on exposure days daily before exposure. Initial preexposure weights were recorded as Study Day 1. Subsequent study days were numbered consecutively through the recovery period.

<u>Necropsy and Histopathology:</u> Rats were killed with carbon dioxide either within four hours or 14 days after the last exposure, bled from the dorsal aorta, and necropsied under the supervision of the pathologist. The necropsy procedure included a thorough, systemic examination and dissection of the animal viscora and carcass and collection and fixation of all major tissues.

Ail tissues and/or organs were examined <u>in situ</u> before dissection from the carcass for individual examination. Tongue, trachea, lungs and pulmonary lymph nodes were removed intact. Following weighing the lungs were infused via the endotracheal route with 10 percent neutral buffered formalin (NBF) before immersion in NBF. The head was removed and NBF was infused into nasal passages prior to submersion in NBF. All tissues were fixed not less than 24 hours prior to trimming.

Tissue trimming (wet sectioning) was performed at IITRI under the supervision and direction of a veterinary pathologist. Organs were trimmed to allow the largest surface area possible for examination. Each lung lobe was sectioned along its main bronchus. Both a cross section and a longitudinal section of trachea was made.

The following tissues and/or organs were microscopically examined: The entire trachea, pulmonary lymph nodes, each lobe of the lungs, and two transverse sections of the skull through the nasal turbinates. (The first section was at the incisor's level. The second section was taken at the palatal ridge level). Since gross examination of the kidneys revealed mottling, these organs were also subjected to histological examination.

Tissues in paraffin blocks were shipped to Experimental Pathology Laboratories inc. (EPL), Herndon, Virginia where hematoxylin and eosin stained slides were propared and examined.

# D. Statistical Methods

The LC<sub>50</sub> concentration was calculated according to the method of Miller and Tainter by linear regression analysis (5). Comparisons made on the preliminary puimonary lavage and bactericidal activity data were done using Student's t tests.

rer Study Nos. 76/77 a four-factor mixed-model ANOVA was used to assess the effects of sex, exposure duration, treatment and replication on lavage parameters (cell count, cellular protein and ATP, and ATP per protein). A multivariate ANOVA for repeated measurements (growth curve model) was used to determine the effects of sex, exposure duration and treatment on repeated body weight measurements over time in Study Nos. 76/77 and 78. These models used in conjunction with this experimental design allow us to separate the effects of sex, exposure duration and treatment. In light of this, overall differences between males and females in body weight, for example, do not confound our estimation of treatment or duration effects. Furthermore, we also evaluated higher order interactions so that we may determine if, for example, a treatment effect is restricted to one sex group (i.e. sex by treatment interactions).

For the purpose of interpretation, individual post-hoc comparisons were performed only when a significant main effect on interaction was found. This allowed us to control the faise positive rate and not be subject to problems inherent in multiple comparisons. In terms of sufficient statistics, means and standard deviations for each experimental condition were provided. In addition, combined mean values were provided for margins that were not statistically significant. For example, if exposure duration was not a significant effect (i.e. no duration by treatment interaction), combined means averaging over duration conditions were provided. Combined standard deviations were not provided because they are not particularly relevant and may be misleading.

# III. <u>RESULTS</u>

# A. Preliminary Studies

In order to obtain preliminary information on the type and magnitude of the biologic responses produced by inhalation of RP/BR aerosols rats were placed into the exposure chambers during some of the exploratory aerosol generation studies of Phase I. These animals were examined for the effects of the exposures on In vivo pulmonary bactericidal activity and for changes in the pulmonary free cells obtained from their lungs by tracheobronchial lavage. Since the results of these experiments served as guidelines in the planning of the exposure conditions for the Phase II studies they have been included into this, instead of the Phase I report. Some of the rats used in these first exploratory experiments were kept In guarantine for evaluation of the automatic drinking water supply In the newly purchased cages and were older than usual. Subsequently 6-to 8-week-old rats were used in the remaining studies as required.

## 1. Pulmonary Bactericidal Activity

to RP/BR aerosol immediately following exposure the experimental and control rats were simultaneously challenged with an aerosol of <sup>35</sup>S-K. <u>pneumonlae</u>. Pulmonary bactericidal activity determined 3 hr after inhalation of the bacteria was significantly depressed in the exposed compared to the control group in all but one of the exploratory experiments (Table 1). Highly significant decreases in bactericidal activity were found in lungs of 12- to 20-week-old male and female rats exposed to 1.88 mg/i of RP/BR aerosol for 1.5 hr, as well as in the lungs of 5-week-old male rats exposed to 1.08 mg/l RP/BR for 2.3 hr. The results after inhalation of approximately 0.5 mg/l of RP/BR in 6-week-old rats varied. Female rats exposed for 2.8 hr to 0.52 mg/l showed significant reduction in bactericidal activity. However when male and female rats were exposed to RP/BR at 0.63 mg/1 for 3.0-hr pulmonary bactericidal activity, although reduced In both sexes, was significantly depressed in male rats only. In another exploratory experiment 7-week-old female rats were exposed 1.0 hr/day for 4 days to RP/BR ac osol at 0.54 mg/l. After 3 days rest, they were exposed for 1.0 hr/day for 3 additional days followed by aerosol challenge with <sup>55</sup>S-K, pneumoniae after the last exposure. Pulmonary bactericidal activity was significantly decreased in the exposed rats. Thus, repeated 1-hr inhalation exposures to approximately 0.5 mg/l of RP/BR aerosol, even when

EFFECT OF EXPOSURE TO RP/BR AEROSOL ON THE PULNONARY BACTERICIDAL ACTIVITY OF RATS Table 1.

		AP/ "	KP/BR Aerosol Exposure a Durati	<pre><pre>sposure Duration</pre></pre>		Rats	315	. % of	% of Inhaled	ç
Experiment No.	Conc, Mean	Conc, mg/l Mean ± SD	No. of Exposures	hr per day	CXT	Sex	Age weeks	Killed Mean	<pre>X: preumonicae Killed in 3 hr ean ± SE</pre>	
ے ہے ق	0 1.88	0,05		1.5	2.5	۲. ۲	12-20	86.8 41.5***	2.6	
, 	0 1.08	0.05		2.3	2.5	×	ъ	88.5 15.4***	2.1	л <u>о</u>
A.	0.52	0.01	-	2.8	1.5	<b>L4</b>	v <b>o</b>	64.0 47.6*	4.4	8
	0 0.63	0.06	~	3.0	٥. -	لغ	6	70.5 64.5	5.2 6.5	<b>ω</b> σ
	0.63	0.06		3.0	6.1	Σ	ڡ	80.2 59.3*	3.5	
	0.54	0.08	7c	1.0	3.8	ليــ	7	80.4 73.3*	1.9 2.7	

Determined gravimetrically from multiple filter-collected aerosol samples. There was a 30 and 42 min interval between phases a and b of Experiments 1 and 3 respectively to recharge the generator with RP/BR. Four exposures, 3 days rest, and 3 exposures.

J

Difference from control: \* p <0.05 \*\*\*\* p <0.001

いたいまたから ちょうしき

「たたたまでものです。「たたたい」ではないでは、「こうないたい」で

Interrupted with recovery days, resulted in significantly depressed pulmonary bactericidal activity.

# 2. Pulmonary Free Calls

ļ

ļ

In other exploratory experiments 11-to 13-week old male and/or female rats were exposed one time for 1 or 4 hr to RP/BR aerosols at 0.88, 2.22, or 3.18 mg/1, or for 4 daily 1-hr periods to 0.55 Within 2 hr after the exposure the animals were killed with ma/l. sodium pentobarbital, and pulmonary free cells were collected by in situ tracheobronchial lavage. Total and differential cell counts were made and total cellular protein levels were determined. The results summarized in Table 2 show consistently decreased total free cell yields and generally increased cellular protein levels in all groups of RP/BR-exposed animals compared to controls exposed to filtered air. The changes were statistically significant at the higher aerosol concentrations with 1 hr exposure durations (2.22 and 3.18 mg/l) and also in male rats exposed 4 times to daily 1 hr concentrations of 0.55 mg/l. Differential counts demonstrated that 96 to 100 percent of the cells were macrophages with occasional lymphocytes (not shown in the Table) present. No significant effects of RP/BR inhalation could be detected l n cellular. distribution.

# B. Range-Finding Experiments

Range-finding experiments were conducted to provide the basis for the choice of the exposure concentrations to be tested in the definitive inhalation studies of Phase II. Mortality, survival time, body weights and in some axperiments clinical observations were used to monitor the effects of inhalation of the RP/BR aerosols.

#### 1. <u>Single Exposures</u>

In the first series of experiments male and female rats ranging in age from 6 to 13 weeks were given single exposures for approximately 1 hr to RP/BR aerosols ranging in concentration from 2.00 to 3.15 mg/l and mortality, survival time and body weight data were recorded over a 14-day period. As shown in Table 3, only exposure to 3.09 mg/l for over 1 hr caused deaths in male rats. Female rats died after single exposures to 2.62, 3.09 and 3.15 mg/l RP/BR. Mortality increased from 11 percent to 40 percent while mean survival time decreased from a 14.1 to 10.4 days with increasing RP/BR aerosol concentration. The maximum mortality resulting from a single 1+hr exposure to approximately 3 mg/l of RP/BR aerosol (maximum generator capacity) was 30 to 40 percent in female rats and 20 to 25 percent for both sexes combined. Results of a single 4-hr e posure to 0.88 mg/l with a CxT value similar to those in the 3.09 and 3.15 mg/l 1-hr studies caused no deaths in either sex, thereby suggesting that exposure concentration, rather

8

EFFECTS OF EXPOSURE TO RP/BR AEROSOL ON FREE CELLS LAVAGED FROM THE LUNGS OF RATS<sup>a</sup> Table 2.

			No.of		æ	Rats							
No.	Conc, mg/1 Maan ± SD5	hr/day	dai ly exp	CXT	Age wk	Sex	_	Total cells x 106	al x 106	macro	% macrophage	Prote µg/105	Protein /105 celis
33	0 0.88 ± 0.02	00.4		0 3.6	12	£	-4 00	0.11	± 1.5 ± 2.1	98.3 97.6	± 0.8 ± 1.2	15.8 23.9	± 1.0 ± 1.7
	6 0.88 ± 0.02	0.4		3.6		. <b>LL</b>	-1 00	9.8 8.0	± 1.4 ± 1.1	96.5 96.5	± 2.5 ± 2.8	19.2 23.1*	+ + - + +
A .	36A 0 2.22 ± 0.00	0.1		2.2	<u> </u>	<b>E</b> .	စာဗ	12.6 7.8**	+ 1.3 + 0.8	98.8 98.8	± 0.8 ± 0.6	20.0 23.8**	+ +
<b>1</b> 9 9 9 9	0 3.18 ± 0.40	0.1		0 3.2	Ξ	£	é é	6.0 2.3**	± 1.0 ± 0.7	not	done	20.9 37.3**	+ 1.0 + 4.0
	0 3.18 ± 0.40	0.1	هوارم . ۲	0.5	,	44	υ, φ	3.7 ± 1.3***±	± 0.4 ÷ 0.3	not	done	19.5	± 2.6 ± 2.7
72	0 0.55 ± 0.03	0.1	ৰ ব	0 2.2	13	Σ	- <b>4</b> IV	11.6 8.5 <sup>†</sup>	± 1.7 ± 0.8	58.5 98.2	± 0.2 ± 0.1	18.5 23.6**	± 7.5 ± 0.8
·	0 0.55 ± 0.03	0.1		0.2.2	13	لد	-4 M	1.1 5.9	± 0.9 ± 0.7	98.0 99.2	± 0.7	22.3 22.3	+ + 

All data are expressed as mean t S.E. for n number of rats. Determined gravimetrically from multiple filter-collected aerosol samples.

۵

ы

8 rats were used for protein assay. -

in this experiment only lungs were excised before lavage resulting in much lower cell recovery. † p <0.10 \* p <0.05 Significant difference from control:

\*\* p <0.01 \*\*\* p <0.001

Table 3. MORTALITY AND MEAN SURVIVAL TIME<sup>a</sup> of RATS AFTER SINGLE EXPOSURES TO RP/BR AEROSOL

ļ

5

. . . . . .

		. ,	÷.	Age									
.'	RP/BR	RP/BR Aerosol Ext	Exposure		ŗ		Mortali	ty			X	lean Surviv	ali
	Conc.	Conc., mg/l	Duration,		Mal	e	Fem	ale	M+F			Time <sup>a</sup> , <sup>D</sup> ay	5
Exp. No.	Mean	₹ 20 <sup>0</sup>	hr	Wks	<u>p/1 %</u>	32	<u>p/1 %</u>	8	1/0	% 2	Male	Female M+	M+F
<b>20</b>	3.15	0.27	1.2	۲.	0/5	0	2/5	40	2/10	20	15.0	10.4	12.7
468	3.09	0.18	1.0	6	2/10 20	20	3/10	30	5/20	25	12.2	11.0	11.6
368	2.62	0.08	1.0		6/0	Φ	1/9	Π	1/18	9	15.0	14.1	14.6
36A	2.22	0.00	teret Beret	12-13	6/0	0	6/0	0	0/18	0	15.0	15.0	15.0
46A	2.00	0.33	1.0		01/0	0.	01/0	0	0/20	0	15.0	15.0	15.0
33	9.88	0.02	4.0	12	U/4	0	0/4	0	0/8	0	15.0	15.0	15.0
			, <b>)</b>										
					•	ļ				-			

Mean survival time determined after recording deaths during exposure day and a 14-day post-exposure observation period (total of 15 days).

Means + SD calculated from multiple daily filter-determined aerosol mass concentration data.

20

than duration, is the determining factor.

Body weights of the survivors were monitored at various times for 14 days after the single expo unes to 2.00, 2.22, 2.62, 3.09 or 3.15 mg/l of the aerosol. Mean body weights are recorded in Table A-1 of the Appendix. Figure 1 shows the mean body weights of male and female rats which survived the exposures to 2.00, 3.09 and 3.15 mg/l as recorded over a 14-day period. There were marked decreases in body weights of both sexes immediately after the single exposure with gradual recovery of the starting weight range after about 9 days and continued increases thereafter. No control animals were included in any of these studies. ムーンション・ショー たいとうたいとう ないとうとう しょくろうかい

Thus although these preliminary mortality experiments were conducted with rats of various ages they demonstrated that  $LC_{50}$  studies with single exposures at logarithmically spaced concentrations, as required in the contract, were not feasible within our system because of the nature of the biologic responses (20 to 25 percent mortality) in the range of aerosol concentration levels physically attainable with the generators (maximum capacity at our standardized 500 l/min airflow rates approximately 3 mg/l for 1 hr).

# 2. Multiple Exposures

A series of multiple exposures were therefore conducted in which 6 - to 8-week-old male and female rats inhaled RP/BR aerosol for 1 hr per day on 5 consecutive days at concentrations ranging from 1.56 to 3.05 mg/l. In another study rats received five daily 4-hr exposures to 0.35 and 0.99 mg/l of RP/BR aerosol. The results from these experiments are shown in Table 4. With four of the experiments (Nos. 65A and B, and 57A and B) control groups exposed to filtered air were also included. Mortality rates, mean survival times, body weights and overall clinical observations were made for a total of 19 days (during the 5 exposure days and for an additional 14-day observation period. Necropsies were done on all animals that died spontaneously during the study. Survivors were necropsied on Day 19.

Mortality and mean survival data show that for the 1-hr exposures, mortality increased from 0 and 10 percent 1.56 mg/l to 80 and 100 percent at 3.05 mg/l for male and 1 male ats resepctively. The overall change in mortality of the two sexes combined was 5 to 90 percent. Mean survival times were correspondingly decreased or unchanged. In the 4-hr exposures, only one female rat died in the entire study and that was at the lower exposure concentration. Observations from the follow-up gross necropsy did not explain the unexpected death of this animal.

The LC<sub>50</sub> value calculated from the mortality data of the 5 daily 1-hr exposure dose response studies shown in Table 4 for the



MORTALITY AND MEAN SURVIVAL TIME® OF 6- TO 8-WEEK-OLD MALE AND FEMALE RATS AFTER 5 DAILY EXPOSURES TO RP/BR AEROSOL Table 4.

	RP/BR	Aerosol	RP/BR Aerosol Exposure Daily			Mortalitv	itv		۰.	e H	ftean Survival	_
•	Conc	[/6m	Period.	Mal	e	Fena	le.	J+W	}	•	ime Days	
Exp. No.	Mean	Mean ± SD <sup>D</sup>	h: H	<u>e</u> 1/0	8	<u>077 %</u>	se	0/1	82	a e	e Fenale	M+F
52A	3.05	0.29	1.0	\$15	0 <i>3</i>	5/5	100	01/6	06	6.6	2.0	4.3
658	2.49	0.67	1.0	4/10	40	5/10	50	9/20	45	12.6	11.1	11.8
528	1.99	0.13	1.0	3/10	30	4/10	40	7/20	35	13.9	12.6	13.3
. 65Å	1.56	0.10	1.0	01/0	0	01/1	10	1/20	م	19.0	17.8	18.4
65	0	•	1.0	01/0	0	01/0	C	0/20	C.	19.0	19.0	اد.0
578	0.99	0.07	4.0	01/0	0	01/0	0	02 <i>1</i> U	0	19.0	19.0	0.01
57A	0.35	0.05	4.0	01/0	0	1/10	10	1/20	เว	19.0	17.7	18.4
57	0		4.0	0/10	0	01-10	0	02/0	0	19.0	19.0	19.0
•	•		· •			,						·
<sup>a</sup> Mean sur	vival t observa	ime deter ation per	Mean survival time determined after recording d ernosure observation period (total of 19 days)	ter rec	ording 9 days	g death: :)	s duri:	ng 5 ex	posure	days an	Mean survival time determined after recording deaths during 5 exposure days and a 14-day post- ernosure observation period (total of 19 days)	post-

exposure observation period (total of 19 days).

<sup>b</sup> Overall means <u>+</u>SD calculated from 5 daily means obtained from multiple daily filter-determined aerosol mass concentration determinations.

aerosol concentrations of 0, 1.56, 1.99, 2.49 and 3.05 mg/l was 2.32 mg/l with 95 percent confidence intervals of 1.99 and 2.73 mg/l respectively.

Figures 2 and 3 show the mean body weights in the survivors these 5 daily 1-hr exposure studies. (The data are recorded from In Table A-2 of the Appendix.) It can be seen that at all four concentrations there are marked decreases in body weights of both male and female rats during the exposure period. Although no controls were included with the exposures at 3.05 and 1.99 mg/l (Figure 2) it appears that survivors of the 3.05 mg/l exposure concentration did not even recover their starting weight during the 14-day observation period. The second set of experiments (Figure 3) definite weight losses after 1.56 and 2.49 mg/l exposures show relative to air-exposed conrols with a more marked effect after the exposure to the higher concentration. Although the rate of weight gain appeared to be similar in exposed and control rats after the initial weight loss in the RP/BR exposure groups, the exposed animals never attained the weight of the controls.

The corresponding mean body weights for the 4-hr exposure studies are shown in Figure 4 and recorded in Table A-2 of the Appendix. It appears that for both concentrations there was only a slight initial weight loss in exposed female rats relative to controls and recovery was complete by the end of the observation period. In males there appeared to be a somewhat more marked effect during and after exposure to 0.99 mg/l of the aerosol. The single "dipping" point observed on Day 8 in the 0.35 mg/l exposure group was the result of limited food availability over the week-end because of a non-aligned feeder.

Clinical observations were made twice daily in all six studies during the five exposure days and in the 14-day observation period that followed. Control animals appeared normal throughout. An occasional incidence of crusted eye was observed in 5 to 10 percent of all groups including the controls. As the RP/BR concentration was increased in the four 1-hr exposure experiments from 1.56 to 3.05 mg/1 the fraction of animals shouling labored breathing at some time during the observation period increased from 20 percent (occasional incidence) to 100 percent (through the entire 19 days). The fraction of animals which were lethargic at some time increased from 20 percent to 60 percent. Reddish discharge from the nose was observed in 30 percent of the animals exposed to 3.05 mg/1 of the aerosol only.

In the experiments in which the rats were exposed five times for 4 hr to filtcred air, 0.35 mg/l RP/BR or to 0.99 mg/l RP/BR, a crusted eye was occasionally seen in 5 to 10 percent of the i mais. Ten percent of the controls and 20 percent of the 0.35 mg/l group were hyperreactive at some time. Ten percent of the

**IIT RESEARCH INSTITUTE** 









\_\_\_\_\_

animals exposed to 0.99 mg/l were lethargic at some time during observation.

All animals which died were necropsied and observations were made of lungs, trachea, esophagus, liver, stomach, intestines, cecum. No control animals were included for experiments 52A and 52B (3.05 mg/l and 1.99 mg/l aerosol for five 1-hr exposures). In general, the observations on these spontaneous deaths included overall mottled red lungs with occasional dark red foci, occasional tan foci on the liver, and gas-filled stomach. Intestines and cecum. Controls included in other studies and all surviving animals were killed by CO<sub>2</sub> inhalation and exsanguination at the conclusion of the observation period. Control animals (Experiments 57 and 65) exhibited the same mottled red lungs with occasional pinpoint or larger red foci and an occasional tan focus on the liver as observed in the exposed groups (0.35 mg/l and 0.99 mg/l RP/BR for five 4-hr exposures for Experiments 57A and 57B respectively). Observations in necropsies of animals exposed to 1.56 mg/l RP/BR showed a greater incidence of red mottling and red foci in the lungs. Necropsies of animals exposed to 2.49 mg/l RP/BR (Experiment 65B) yielded similar observations with the addition of occasional necrotic foci on the liver in spontaneous deaths.

This comparison of the 1-hr and 4-hr studies in terms of body weight changes and of clinical observations demonstrated marked toxic effects on both parameters after the 1-hr exposures but negligible effects after 4-hr exposures to lower concentrations of the RP/BR. These results again indicate the importance of the exposure concentration relative to duration since for similar or higher CxT values in the 4-hr studies there were minimal effects compared to the 1-hr exposure.

# C. Definitive Studies

D

These preliminary range finding data were used as a basis for . the final design for the Phase II definitive studies. These studies were conducted to provide information on selected biologic response parameters after four dally exposures to 0.5 mg/l of RP/BR aerosol and comparing the effects at 1.0 - and 3.5 - hr exposure durations. The biologic endpoints selected for this study were pulmonary free cell parameters (total and differential cell counts cellular protein and ATP levels in the free cells obtained by and tracheobronchial lavage), body weights and potential structural necropsy changes (gross and histopathology). Pulmonary bactericidal activity was not chosen for this study because of the. very high sensitivity of this assay. The selected parameters were tested immediately following the last exposure and after a 14-day recovery period. In an additional study rats were exposed to 0.5 mg/1 of RP/BR for 3.5 hr daily 4 days/week for 4 weeks and examined for histopathologic changes with and without recovery.

IIT RESEARCH INSTITUTE

# 1. Experimental Design

The experimental design with exposure group allocations for study Nos. 76, 77 and 78 is shown in Table 5. In Study Nos. 76 and 77 four groups of 14 male and 14 female rats were exposed for 1.0 or 3.5 hr daily, four times per week for one week to 0.5 mg/l of RP/BR aerosol. Four groups were exposed to filtered air. In Study Nos. 76A and B, two RP/BR and two air-treatment groups (1 and II: V and VI) were exposed for 1.0 hr per day on 4 consecutive days. In Study Nos. 77A and B, the remaining four treatment groups (III and IV; VII and VIII) received 4 daily 3.5 hr exposures to RP/BR or air. One RP/BR and one air group from each exposure duration (treatment groups 1 to 1V) were sacrificed subsequent to the last exposure. Pulmonary lavages with total and differential cell counts, cellular protein and ATP levels were done for 10 male and 10 female rats from each group. The remaining 4 males and 4 females were killed, necropsied and tissues from 2 rats each sex were processed for histopathologic examination of the of respiratory tract. The other RP/BR and air treatment groups from each exposure duration were maintained for a 14-day recovery period (treatment groups V to VIII) at the end of which the same numbers of animals from each group as listed above were used for pulmonary lavage, necropsy and histopathology.

Study No. 78 was conducted to determine if any gross structural changes in the respiratory tract could be discovered relative to controls from lihalation of 0.5 mg/l of RP/BR aerosol, 3.5 hr daily, four days per week over a four-week period at the completion of the exposures and after a 14-day recovery period. Pulmonary lavage parameters were not examined in this study.

Animals in all three studies were weighed on a regular schedule and clinical observations for toxic signs were made twice daily except for weekends and holidays.

# 2. Exposure to RP/BR Aerosol

The rats were exposed to RP/BR aerosol or filtered air in 1-m<sup>3</sup>-sized inhalation chambers as described in the methods section. For animals receiving the same exposure regimen but different post-exposure treatments (that is, with or without recovery; such as treatment groups 11 and VI and IV and VIII, respectively), common chambers were used and, therefore, common exposure concentrations were calculated. However, in order to facilitate the time-consuming and labor-intensive processes of pulmonary lavage and necropsy of the test animals, the exposures were staggered. Exposure of 5 male and 5 female rats from each group for lavage and 2 of each sex for necropsy were started one day and the remaining (5+2) rats per sex per group were started on the next day (Monday through Thursday and Tuesday through Friday, respectively). Therefore, there were actually five days of

IIT RESEARCH INSTITUTE

Table 5. EXPOSURE GROUP ALLOCATION FOR STUDY NOS. 76, 77 AND 78

Necropsy eħ No. of Rats per Sex Total Lavage Necro Lavage 010 010 00 2 2 77 14 1¢ 20 20 7 2 2 Recovery Period (14 Days) Total Weeks No/Week Exposure Hr/day 1.0 1.0 ŝ 3.5 0.1 . . . . . . ŝ s. N Target Conc. (mg/l) 0.5 0.5 0.5 5.0 0.5 0 Ò Treatment Group ٨ŧ T N ------۵ -Study No. 76A 768. 77A 778 7.8

- :

two rats per sex and per treatment group, were necropsied after the exposures and after the 14-day recovery period, respectively: no treatment group designation given.

. 201

multiple daily chamber readings from which the means and standard deviations (SD) for aerosol mass concentration and particle size values for the entire four exposure day study were calculated. Thus, in Study Nos. 76 and 77, for the 1.0-and 3.5-hr aerosoi exposures, 15 (3 per day on 5 days) and 20 (4 per day on 5 days) filter and integrated photosensor samples, respectively, were collected. For Study No. 78, 64 filter and photosensor samples (4 per day on 16 days) were taken. Aerosol mass concentration and particle size data for the various exposure periods were calculated as overall means from the multiple daily means averaged over the entire exposure period. Percent phosphoric acid was determined from one filter-collected sample per week. The aerosol mass concentration and particle size data for Study Nos. 76, 77 and 78 are summarized in Table 6.

و م م م م م م

The mass median aerodynamic diameters (MMAD) shown in the table for Study Nos. 76 and 77 were higher than the values we usually observed (0.45 to 0.63  $\mu$ m). In reviewing the original Quartz Crystal Microbalance impactor data for these studies it was found that for all replicate samples impactor stage No. 7 (cut diameter = 0.84  $\mu$ m) collected between 45% and 60% of the total particle mass, while stage No. 6 (cut diameter = 1.69  $\mu$ m) collected nothing. Thus, the resulting aerosol size distribution was one tailed, and, because the mass median aerodynamic diameter is a single point property of the distribution, the median particle size appears markedly different from the expected value. The mass mean aerodynamic diameter, being a property of the overall distribution, was not as severely affected with values (0.61, 0.57 and 0.54 for Study Nos. 76, 77 and 78) which were essentially equivalent to mean values previously determined under similar conditions during the aerosol homogenity studies of Phase I.

## 3. Mortality and Clinical Observations.

No deaths occurred in any of the studies. All animals appeared normal during the exposure interval and 14-day recovery period except for 2 males for which crusted eyes were recorded during the recovery period. No other treatment-related changes were seen.

#### 4. Body Weights

According to the experimental design for Study Nos. 76 and 77 only the animals designated for lavage were weighed before each of the four exposures and at predetermined intervals during the recovery period. Mean body weights are summarized for treatment groups I through VIII for the four exposure days in Table A-3 and those for recovery animals (treatment groups V through VIII) taken over an 18-day period in Table A-4 of the Appendix. Two separate analyses were conducted, one on the results of the 4 exposure days and one over the entire 18-day period including 4 exposure and 14

#### IT RESEARCH INSTITUTE

31

Table 6. AEROSOL MASS CONCENTRATION, PARTICLE SIZE AND PERCENT H<sub>3</sub>Po4 FOR STUDY NOS. 76, 77 and 78

		RP.	RP/BR Exposure	e				Aerosol <sup>'d</sup>		
Study No.	Treatment Group	Hr/day	No/week	Total Weeks	Recovery	Mass Concentration, mg/ Filter Sample Photosenso	ration, mg/l Photosensor	MMADb .	Particle Size ADb og	H2P04. 2
764	-	0.1	-	-						
		•	•	•	•	0.62 ± 0.06	$0.53 \pm 0.07$	0.53 ± 0.07 0.94 + 0.02 2.28 + 0.08	-2 28 + 0 08	6 09
768	· 1A	0 1	÷	°	+					
AL	A :	3.5	- <b>3</b> '	<b></b>	ı ,					
178	1 f f A	3.5	्र म	-	•	U.5U I U.UZ	0.46 ± 0.01	0.46 ± 0.01 0.78 ± 0.00 2.53 ± 0.03 62.7	2.53 ± 0.03	62.7
	<b>iy</b>	3.5	<b>.</b>		+	0.51 ± 0.04	0.50 ± 0.05	0.50 ± 0.05 0.61 ± 0.14 1.92 ± 0.21 67.8 ± 2.48	1.92 ± 0.21	67.8 ± 2.4

All data given as Mean  $\pm$  SD from daily means except for percent  ${
m H_3PO_4}$  determined from one filter sample per week. For methods see text.

Mass Median Aerodynamic Diameter.

No treatment group designation given.

Ę

32
recovery days. The data were analyzed for between timepoint differences as well as linear trends in weight gain over time.

A multivariate ANOVA of the 4-day exposure period revealed that exposure duration was not a significant factor (there was no treatment by duration interaction) and that the average level of body weights over the four-day period was not significantly affected. However, a significant exposure by sex interaction (F=5.46, df=4/149, p<0.0004) was found. Table 7 displays the daily body weight means averaged over both durations for control and exposed male and female rats. Inspection of the data demonstrated significant (p<0.05) differences for individual time points between exposed and control male but not female rats and that the exposure had a significant effect on the linear rate of weight increase in male but not in female rats.

Analysis of the recovery animals over the entire 18-day period revealed that the previously noted sex by exposure interaction for days 1 to 4 was absent. Again, exposure duration was not found to be a significant factor. A significant overall exposure effect was found (F=8.63, df=4/68, p<.0001). Inspection of the average daily body weight means for control and exposed animals (Table 8) revealed that the average level of body weights was significantly over the entire 18-day period in exposed animals decreased (p<0.008) and that individual timepoint comparisons were also significant (p<0.05). However, the rate of body weight gain was not significantly different between exposed and control animals. This indicates that treatment produces a decrease in body weight that does not return to within the normal control limits in the recovery period.

The mean body weights measured in Study No. 78 in which rats received 16 exposures of 3.5-hr daily duration for 4 days per week are summarized in Table A=5 of the Appendix. The rats were weighed before the first exposure and then weekly on Fridays for the four exposure weeks and two recovery weeks. Statistical analysis of the data was conducted for both between timepoint body weicht differences and trends in body weight measurements over time. Nø significant treatment by sex interaction was found. The average body weights of exposed animals over time were significantly lower those of controls (p<0.002). However, RP/BR exposure did not than affect the overall rate of body weight gains relative to those in As shown in Table 9 significantly lower body weights controls. were seen in exposed rats at all timepoints after the initial preexposure weighing on Day 1: Day 5 (p<0.0002), Day (p<0.00008), Day 19 (p<0.0002), Day 26 (p<0.008), Day Day 12 33 (p<0.002), and Day 39 (p<0.02). The attenuation of the probability values on Day 33 and Day 39 Indicates a recovery effect in which treated animals return to within the 92 percent limits of controls. These body weight measurements are, however, still significantly different (p<0.02) from controls.

#### IIT RESEARCH INSTITUTE

MEAN BODY WEIGHTS<sup>a</sup> OF MALE AND FEMALE RATS DURING FOUR DAYS OF EXPOSURE TO RP/BR AEROSOLS (Study Nos. 76A and 77A)

Table

	RP/BR Exposure	osure		Mean Bo	dv Weights	Mean Body Weights (g) Determined	termined
Treatment	(ma/1) <sup>5</sup>	Duration			on Stu	on Study Day:	
Groups	Mean ± 50	(hr/day)	Sex	ا د	2	<b>.</b>	+
	0	1.0,3.5	£	201.4	208.7	213.7	- 219.4
11, IV	0.56 ± 6.07	1.0,3.5	X	201.1	204.3*	208.2*	213.1*
÷ ; ; ; ;	0	1.0,3.5	<b>بد</b>	138.9	142.6	144.1	146.8
. At . I	0.56 ± 0.07	1.0,3.5	ia.	139.1	141.5.	142.9	146.6

<sup>a</sup> Calculated for 20 rats per point, averaging over the 1.0 and 3.5 hr exposure duration, because there was no exposure by duration interaction.

b Aerosol mass concentration calculated from daily gravimetric filter readings.

<sup>c</sup> Study Day I = first exposure day. \*

Significantly different from controls (p < 0.05).

MEAN BODY WEIGHTS OF MALE AND FEMALE RATS DURING RP/BR AEROSOL EXPOSURE AND RECOVERY DAYS **a** Table

(STUDY NOS. 76B and 77B).

conc. <sup>a</sup> Duration         Mean Body Weights (g)         Determined on Study Day           (mg/1)         hr/day)         hr/day)         Mean Body Weights (g)         Determined on Study Day           Mean ± SD         for ¼ days         Sex         1 <sup>c</sup> 2 <sup>c</sup> 3 <sup>c</sup> 4 <sup>c</sup> 5           Mean ± SD         for ¼ days         Sex         1 <sup>c</sup> 2 <sup>c</sup> 3 <sup>c</sup> 4 <sup>c</sup> 5           0         1.0,3.5 <sup>d</sup> M.F <sup>e</sup> 171.5         176.7         180.0         184.6         188.4           0.55 ± 0.09         1.0,3.5         M.F         169.5         172.1*         174.3*         178.6*         181.2*		RP/BR Exposure	xposure	,	•					
Mean ± SD         for ¼ days         Sex         1 c         2 c         3 c         4 c         5           0         1.0,3.5 <sup>d</sup> M, F <sup>e</sup> 171.5         176.7         180.0         184.6         188.4           0.55 ± 0.09         1.0,3.5         M, F         169.5         172.1*         174.3*         178.6*         181.2*	reatment	сонс. <sup>а</sup> {ma/1}	Duration hr/dav)		Mean Body	Weights	(g) Detc	rmined on	Study Da	а ``>
9 1.0.3.5 <sup>d</sup> M.F <sup>e</sup> 171.5 176.7 180.0 184.6 188.4 0.55 ± 0.09 1.0.3.5 M.F 169.5 172.1* 174.3* 178.6* 181.2*	Group	Nean 1 SD	for 4 days	Sex	- <sup>-</sup> c	2 <sup>c</sup>	3 <sup>c</sup>	4c	5	0
.09 1.0.3.5 M.F 169.5 172.1* 174.3* 178.6* 181.2*	V. VII	œ	1.0,3.5 <sup>d</sup>	<b>ດ</b> ນ	171.5	176.7		184.6	188.4	193.1
	111A.1V	0.55 ± 0.	1.0,3.5	M, F	169.5	172.1*	174.3*	178.6*	161.2*	185.5*

35

(Page 1 of 2)

bereicher ihreiten bereichten Weicher die Verschreiten bereichten bereichten bereichten bereichten bereichten b

Table 8 (Continued). MEAN BODY WEIGHTS OF MALE AND FEMALE RATS DURING RP/BR AERCSOL EXPOSURE AND

AND RECOVERY DAYS (STUDY NOS. 76B and 77B)

	RP/BR E	RP/BR Exposure	,					,
Treatment	cenc. <sup>a</sup> (mg/1)	Duration hr/day)		Mean	Body Weig	ht (q) De	Mean Body Weight (q) Determined on Dav <sup>b</sup>	on Dav: <sup>b</sup>
Group	Hean ± SD	for 4 days	Sex	Sex 8	=	14	16	-18
11A'A	•	ł.ψ.ȝ.5 <sup>d</sup> M.F <sup>e</sup> 201.6 211.9	R.Fe	201.6	211.9	223.5	223.5 226.5	230.3
111A" 1A	0.55 ± 0.09 1.0,3.5	1.0,3.5	۲. ۲.	195.0*	M.F 195.0* 205.8* 215.5*	215.5*	218.8*	222.4*

Calculated from daily gravimetric filter readings for both exposure durations combined.

Study Day 1 = first exposure day.

Exposure days.

Data combined for 20 male and 20 female rats per point because there was no exposure by sex interaction. Data averaged over the 1.0 and 3.5 hr durations because there was ro exposure by duration interaction. Significantly different from controls (p <0.05). MÉAN BODY WEIGHTS<sup>a</sup> or male and female rats during four weeks of exposure four days/week to RP/BR Aekusols and two weeks of recovery Table 9.

RP/BR Ex	posure								
Aerosol				ncew	Body Valates		-	U	
- Juon	Duration				rear out weights (9), betermined on Study Day;	191, verei	mined on	otudy Day:	
(1/6m)	(hr/day)	Sex	2-	29	12 <sup>d</sup>	p61	26 <sup>d</sup>	33 <sup>e</sup>	39 <sup>e</sup>
0	3.5	.H,F <sup>f</sup>	168.8	188.4	214.3	232.5	248.5	266.9	280.3
0.51 ± 0.04	3.5		168.1	180.1***	203.1***	203.1*** 221.8***	239.8**	257.9*	270.9*
			,						

Weights are from 20 or 18 rats of each sex per group on Days 1 through 26 and Days 33 through 39, respectively.

Data given as Mean ± SD calculated from daily gravimetric filter readings.

Study Day 1 = first exposure day.

Exposure period.

Recovery period.

Data was averaged across sex because there was no exposure x sex interaction.

\* p <0.05 \*\* p <0.01 Significant difference from control:

\*\*\* p <0.001

## 5. Pulmonary Free Cell Parameters

Results of the pulmonary lavage experiments are summarized in Table 10 showing total cell counts, percent macrophages determined from differential counts and total protein and ATP levels In the Statistical analysis of these data revealed no lavaged cells. difference between the effects of 1.0- and 3.5- hr exposures (no duration by exposure interaction) over the entire exposure and observation pariods. In the macrophages of rats examined immediately after the fourth exposures a significant exposure by sex interaction was found for ug total protein per 10° cells For females, averaging over exposure duration PP/BR (p<0.001). aerosol inhaiation exposure produced an increase in total protein (control = 25.23 and exposed = 28.09 ug protein per  $10^{2}$ cells). whereas males exhibited a decrease (control = 25.24 and exposed = 22.99 ug protein per 10<sup>2</sup> in the absence of other cells). treatment by sex interactions the data obtained from the two sexes were combined for all other parameters in Table 11. In addition, In the absence of an exposure by duration interaction the experimental observations were also averaged over the two exposure durations. When examined immediately after the last. RP/BR exposure, (treatment groups 1 through IV) significant overall decreases in ATP content per  $10^5$  cell (p<0.002) and in ATP per ug protein (p<0.008) were observed. Statistical analysis of the data obtained after the recovery period (treatment groups & through VII) revealed no significant exposure-related effects; hence recovery from these exposure effects was complete within 14 days.

#### 6. Gross Necropsy and Histopathologic Observations

Gross observations made on the rats during necropsy for the 4-day and the 4-week exposure studies are summarized in the Appendix in "Pathology Report, Part I". After the 4-day exposure study mottled red lungs were found in many of the rats in both the treated and filtered air groups and in both the exposure and recovery groups. The small number of animals per group makes it difficult to determine the significance of this lesion. Other lesions which occurred with some frequency were uninary calculi in males especially in the recovery group, mottled brown kidneys, and distended uteri. These and the remaining lesions observed were regarded as incidental findings and were present in both the control and treated animals.

No compound related gross pathologic lesions were observed in rats from the experimental group sacrificed after the 4-week exposures, nor in rats sacrificed after two weeks of recovery period. All lesions observed at necropsy were considered to be spontaneous in nature and unrelated to inhalation of RP/BR aerosols.

From the 4-day exposure studies two rats per sex per group

IIT RESEARCH INSTITUTE

EFFECTS OF FOUR DAYS OF EXPOSURE TO RP/BR AEROSOLS ON PULMONARY FREE CELLS LAVAGED Table 10.

9

FROM THE LUNGS OF MALE AND FEMALE RATS<sup>a</sup>

	ļ	. 59	SE	0.6	0.3	0.6	0.3	0.7	0.3	0.4	0.4
	phages	Females	Mean ± SE	96.8 ±	98.5 ± 0.3	∓ 0*66	98.9 ± 0.3	98.1 ±	98.9 ± 0.3	± 1.96	98.0 ± 0.4
ints	* Macrophages	Males	Mean ± SE	4.0 ± 0.6-	98.9 ± 0.4	99.1 ± 0.4	98.9 ± 0.5	97.1 ± 1.0	98.7±0.6	98.2 ± 0.4	97.8 ± 1.0
Cell Counts	106	Feinales	Mean ± SE	6.57 ± 0.41	$6.25 \pm 0.47$	7.15 ± 0.48	6.06 ± 0.75	7.88 ± 0.74	8.47 ± 0.73	10.03 ± 1.15	9.84 ± 1.21
	Total × 10 <sup>6</sup>	Males	- Mean ± SE	8.79 ± 0.61	9.20 ± 0.81	8.21 ± 0.61	7.51 ± 0.51	11.67 ± 1.70	10.52 ± 1.02	7.86 ± 1.08	12.69 ± 1.40
RP/BR Exposure		buration	(hr/day)	0.1	0.1	3.5	3.5	0.1	0.1	3.5	<b>3.</b> 2
RP/BR F	Conc.		Tean .5U	0	0.02:0.00	0	0.50.0.02	0	0.62:0.06	0	0.50.0.02
		lrealment	e roup	م_م		<u>م</u> 	<b>N</b>	<b>u u</b> >	-	211 <sub>2</sub>	
		Ann a		764		774		768		778	
		·							•		

(Page 1 of 2)

Table 10 (Continued). EFFECTS OF FOUR DAYS OF EXPOSURE TO RP/BR AEROSOLS ON PULMONARY FREE CELLS LAVAGED

C

D

FROM THE LUNGS OF MALE AND FEMALE RATS<sup>a</sup>

		RP/BR Conc	RP/BR Exposure	ng Prote	Protein/10 <sup>5</sup> Cells	ATP fq x	ATP fq x 10 <sup>8</sup> /10 <sup>5</sup> Cells	ATP for v 10	ATP for v 106/110 Brotheir
Study No.	Group	(mg/l) Mean •SD	Duration (hr/day)	Males Mean ± SE	Females Mean ± SE	Males Mean ± SE	Femaler Mean ± SE	Males Mean ± SE	Females
76A	ر <u>م</u> م	0 0.62+0.06	0.1	26.26 ± 2.10 21.51 ± 1.10	25.48 ± 0.96 26.65 ± 0.70	0.59 ± 0.09 0.48 ± 0.07	0.80 ± 0.11 0.54 ± 0.10	$2.42 \pm 0.42 \\2.27 \pm 0.37$	
	، مر =>	0.50±0.02	3.5	26.14 ± 1.18 24.47 ± 0.91	24.99 ± 1.06 29.53 ± 1.59	$1.10 \pm 0.13$ $0.86 \pm 0.07$	1.11 ± 0.08 1.01 ± 0.11	4.18 ± 0.46 3.52 ± 0.29	$4.50 \pm 0.33$ $3.51 \pm 0.38$
768		0.62±0.06	0.1	20.91 ± 0.56 23.29 ± 1.44	22.41 ± 1.45 21.46 ± 0.42	0.61 ± 0.09 0.55 ± 0.13	$0.77 \pm 0.15$ $0.70 \pm 0.09$	3.00 ± 0.44 2.50 ± 0.63	3.24 ± 0.55 3.27 ± 0.41
	۲۱۱ <sup>۲</sup> ۲۱۱۱	0.50+0.02	3.5	$21.16 \pm 2.37$ $22.62 \pm 3.30$	25.25 ± 2.91 21.21 ± 0.80	0.60 ± 0.13 0.56 ± 0.14	0.76 ± 0.14 0.59 ± 0.15	2.28 ± 0.30 3.28 ± 0.64	3.00 ± 0.56 2.76 ± 0.60
Ó	10 rats per group.	• dno							

b Tested immediately after the last exposure.

Tested after a 14-day recovery period following the last exposure.

(Page 2 of 2)

OVERALL MEAN VALUES<sup>a</sup> FOR ENDPOINT PARAMETERS MEASURED ON PULMONARY FREE CELLS LAVAGED FROM THE LUNGS OF RATS AFTER EXPOSURE TO RP/BR AEROSOLS Table II.

		RP/BR E	Exposure		•			
Study No.	Treatment Groups	Conc. (mg/l) Mean ± SD <sup>b</sup>	Duration (hr/day) for 4 days	Recovery Period (14 day)	Total cells x 10 <sup>6</sup>	lug Protein 10 <sup>5</sup> cells	$\frac{\text{ATP fg x10^8}}{10^5 \text{ cells}} = \frac{\text{A}}{10^5 \text{ cells}}$	ATP fg x 10 <sup>6</sup> ug protein
764, 774	1 1 1 1	Ö	1.0,3.5	ŀ	7.73	IJ	0.92	3.64
76A, 77A	11,1V	0.55 ± 0.09	1.0,3.5	ŀ	7.25	U	0.73**	2.85***
768, 778	V, VI	<b>0</b>	1.0,3.5	÷	9.35	22.79	0.65	2.78
768, 778	V1, V111	0.55 ± 0.09	1.0,3.5	¥	10.38	21.42	0.61	2.95
	r							

Mean values obtained averaging over replication, sex, and exposure duration since no duration by exposure or sex by exposure interactions were found.

b Calculated from daily gravimetric filter readings for both durations.

<sup>C</sup> Not reported because of a significant exposure by sex interaction (see text).

Significant difference from control: \*\* p <0.01 \*\*\* p <0.001

÷

were submitted for histologic evaluations of the respiratory tract. Because gross examination of the kidneys from several animals revealed mottling, the following tissues were trimmed and processed to paraffin blocks: trachea, pulmonary lymph nodes, each lung lobe, nasal turbinates, and kidneys. The paraffin blocks were then shipped to Experimental Pathology Laboratories, Inc. where hematoxylin and eosin stained slides were prepared and examined. The results are summarized in "Pathology Report, Part II" and included in the Appendix.

The microscopic changes and a detailed listing of all tissues evaluated are presented in the Histophatology incidence Tables. All lesions are summarized by sex and treatment group and presented in the Summary Incidence Tables. A correlation of lesions observed at necropsy with the corresponding microscopic observation, where possible, is presented in the Correlation of Gross and Microscopic Findings Tables. The gross observations in these tables were transcribed from the necropsy sheets provided with the paraffin blocks.

A summary of the results of the microscopic examinations indicate that administration of RP/BR at the concentrations and for the duration of exposure used in this study did not produce treatment-related changes in the kidney, trachea, or the nasai turbinates at the levels examined.

Several animals which recovered for 14 days following either a 1.0 - hour or 3.5-hour exposure (for four days) to RP/BR combustion products had focal creas of Interstitial reaction with alveolar macrophages which may be treatment related. However, these animals also had moderate to marked lymphold hyperplasia of pulmonary lymph nodes which suggests an infectious agent such as Sendal virus, Mycoplasma pulmonis or pneumonia virus of mice (PVM) may have lesions. Although Inguirles produced these to Harlan/Sprague-Dawley, Inc., supplier of the rats indicated that results of viral serology and bacteriology assays of this rat shipment showed Sendal and Mycoplasma sp to be negative there was a positive PVM titer.

#### LY\_ CONCLUSIONS

 $\bigcirc$ 

2

D

•

Exploratory experiments with male and female Sprague-Dawley rats exposed to RP/BR aerosols ranging from 0.5 to 3 mg/l for durations of 1 to 4 hr showed highly significant decreases in pulmonary bactericidal activity to inhaled <sup>35</sup>S-K\_pneumoniae in exposed relative to control rats after single as well as multiple exposures. The higher exposure concentrations also produced significant decreases in total cell counts in the pulmonary free cells obtained by tracheobronchial lavage from the exposed rats. The subsequent mortality studies with exposures to RP/BR aerosols ranging from 2.00 to 3.15 mg/l mass concentration demonstrated that the maximum mortality resulting from a single 1-hr exposure to approximately 3 mg/l of the aerosol (maximum generator capacity) was 20 to 25 percent, while 2.62 mg/l resulted in 6 percent deaths. Thus it became evident that  $LC_{50}$  studies with single exposures at logarithmically spaced concentrations, were not feasible in the range of aerosol concentration levels physically attainable with the generators at the air flow rates required in our inhalation exposure system.

A single 4-hr exposure to 0.88 mg/l with a CxT value similar to those in the 3.09 and 3.15 mg/l 1-hr studies caused no deaths thereby suggesting that exposure concentration is the determining factor rather than duration.

A follow-up multiple exposure study with rats inhaling RP/BR aerosols for 1-hr dally for 5 consecutive days at concentrations of 1.56, 1.99, 2.49 and 3.05 mg/l showed mortality rates ranging from 5 to 90 percent with an estimated  $LC_{50}$  value of 2.32 mg/l. In contrast, when rats received five daily 4-hr exposures to 0.35 or 0.99 mg/l of RP/BR only one died and comparison of body weights and of clinical observation for the 1- and 4-hr studies showed markedly negative effects after the 1-hr and practically negligable ones after the 4-hr exposures. Thus the importance of exposure concentration over duration in terms of producing effects at comparable CxT values became evident again.

In the definitive studies male and female rais received four daily exposures to 0.5 mg/l of RP/BR aerosol for 1.0 or 3.5 hr durations. In an additional experiment rats were exposed to 0.5 mg/l of RP/BR for 3.5 hr daily 4 days per week for 4 weeks.

No deaths occurred in any of the studies and generally no treatment-related clinical observations were seen during any of the exposures on the following 14-day, recovery periods. Statistical analysis of body weights showed that exposure duration was not a significant factor. When examined over the exposure and recovery periods combined, the 4-day as well as the 4-week-RP/BR exposures produced significant weight decreases in rats of both sexes and the weights did not return to the normal control levels within tha recovery period.

No compound-related gross pathologic lesions were observed during necropsies immediately following the last epxosure, or after a 14-day recovery period in either the 4-day or the 4-week exposure group.

Exposure duration did not have a significant effect on the free cells lavaged from the lungs. Total and differential cell counts were not affected. Cellular ATP levels were significantly

decreased when examined immediately after the fourth exposure, however recovery was complete after 14 days.

Microscopic examination of the tissues immediately following the 4-day exposure and after a 2-week recovery period did not reveal any treatment-related changes in kidneys, trachea and nasal turbinates. The lungs of several animals had focal areas of interstitial reaction with alveolar macrophages which may have been treatment-related. However, in the opinion of the pathologist, the presence of a possible reaction of marked lymphoid hyperplasia of pulmonary lymph nodes which suggests an infectious agent and the information received from the animal supplier about a positive PVM titer in the breeding colony indicates no compound-related effect.

Thus these range finding studies have demonstrated that inhalation of C.5 mg/l RP/BR aerosol for 1.0 and 3.5 hr dally for 4 days, or for 3.5 hr 4 times weekly for 4 weeks resulted in decreased body weights relative to control animals, but the exposures did not produce gross pathologic changes and the microscopic changes observed in lung tissue could not be considered compound- related. The changes found In pulmonary lavage parameters after four exposures were no longer present after the recovery period. Since the concentration level, 0.5 mg/l, used in these studies produced minimal toxicity, the Phase III studies will use concentration levels at and above this level.

## LITERATURE CITED

- Research and Development on Inhalation Toxicologic Evaluation of Red Phosphorus/Butyl Rubber Combustion Products, Phases I Report. C. Aranyl, August 1983, Contract No. DAMD17-82-C-2121.
- Effects of Subchronic Exposure to a Mixture of 03, S02, and (NH4)2S04 on Host Defenses of Mice.
   C. Aranyi, S. C. Vana, P. T. Thomas, J. M. Bradof and J. D. Fenters, 1983, J. Toxicol. and Environ. Health 12:55-71.
- Protein Measurement with the Folin Phenol Reagent.
   H. Lowery, N. J. Rosenbrough, A. L. Farr and R. J. Randall, 1951, J. Biol. Chem. 193:265-275.
- 4. The Linear Relationship Between the Quantity of Serum and the Quantity of Virus Neutralized. F. L. Horsfall, Jr., 1939, J. Exp. Med. 70:209-222.

45

 Estimation of the LD50 and its Error by Means of Logarithmic-probit Graph Paper, L. C. Miller and M. L. Tainter, 1944, Proc. for Exp. Biol. Med. 57:261-264.

AFPENDIX

TABLES

··· ,	8P/88			W	Mean Body Weights (g) Determined on Study Day:	(g) Determined on	Study Day:	
Study No.	conc. (mg/1)	Sex	Age	J.	2	3	4	
36A	2.22±0.00	× 4	12-13	384.8±15.7(10) 233.2±11.7(10)	369.6±19.5(10) 231.2± 6.7(10)	367.2± 18.0(10) 233.4± 6.5(10)		
368	2.62±0.08	.≭ti⊾.	12-13	387.5±27.5 (9) 221.9±10.6 (9)	379.0±27.4 (9) 220.0±10.9 (9)	374.9±27.7 (9) 220.2±11.8 (9)		
464	2.00±0.33	X 14.	<u>م</u> م	139.0±17.9(10) 137.1± 7.9(10)	123.6±16.5(10) <sup>d</sup> 119.9± 8.5(10) <sup>d</sup>			138.5±23.7(10) 135.6± 7.6(10)
468	3.09±0.18	24	<b>.</b>	133.9±20.7(10) 133.3± 9.3(10)	115.4±11.8(8) <sup>d</sup> 114.3± 8.5(9) <sup>d</sup>			116.6±16.0 (8) 115.3±15.4 (7)
50	3.15±0.27	Z Ia	1	174.6±15.6( 5) 154.2±14.0 (5)	143.4±14.9 (5) 126.0±10.5 (4)	132.2±15.1 (5) 114.0± 5.6 (3)	132.4±19.2 (5) 117.3± 9.1 (3)	137.4±23.0 (5) 127.3± 9.0 (3)
								(Page
÷.,	- - -				,			1 of

ľ

そのには第二人のためを見てためのという。 第二人のないです

ことが、 思える かいかい 相応的 いたものの

MEAN BODY WEIGHTS<sup>a</sup> of male and female rats exposed one time to RP/BR Aerosols AND MONITORED FOR 14 DAYS

Table A-1 (Continued),

Study No.	conc." (mg/1)	Sex	Age wks	1		6	15 15
36A	2.22±0.00	X 4.	- 12-13	370.3±15.1(10) 233.6± 7.7(10)			382.0±12.9(10) 226.3±23.9(10)
89	2.62±0.08	<b>X</b> ii.	12-13	378.8±30.4 (9) 228.1±10.9 (9)		• • •	387.2±29.4 (9) 230.9±10.7 (8)
46A	2.00±0.33	. X 4	ف		155.2±28.6(10) 147.7±12.5(10)	160.8±28.0(10) 149.7±15.4(10)	196.7±35.4(10) 170.0±11.7(10)
468	3.0910.18	X I.	, <b></b>		129.0±21.9 (8) 131.7±17.0 (7)	3 .1±25.5 (8)  37.4±15.9 (7)	161.8±37.1 (8) 163.4±14.7 (7)
20	3.15±0.27	× 14	~~	••	155.0±24.5 (5) 143.0± 6.6 (3)		202.6±34.2 (5) 168.3± 4.9 (3)

Bata given as Mean 1 SB (number of rats).

B Mean ± S0 for gravimetric filter samples.

c Exposure day = Study Day 1.

These weights were recorded after exposure on Day 1

MEAN BODY WEIGHTS<sup>a</sup> of male and female rats<sup>d</sup> exposed five times to RP/BR Aerosols and monitored for 14 Cays Table A-2.

	RP/BR		· · · · · · · · · · · · · · · · · · ·		Mara Badu Maiakta (a) Dataminad an Study Davi	.ved vbutt	
Study No.	mg/ml Mean ± 50	Sex	d.e	rean budy wery	3 <sup>6</sup>	4 011 Juny 097.	2e
52A	3.05±0.29	X 14.	206.6±11.2 (5) 166.6±18.5 (5)	174.9±11.6 (5) 148.1± 7.7 (4)	163.0±14.5 (5) 140.8± 9.7 (4)	150.5±15.6 (4) 137.0 (1)	130.0± 9.2 (2) 138.0 (1)
528	1.99±0.13	×. L.	213.2±21.1(10) 169.2±10.8(10)	198.0±17.5(10) 156.6±15.0(10)	187.1±22.3 (9) 154.8± 6.9 (8)	186.8±22.8 (8) 151.5±12.5 (8)	192.0±21.7 (7) 155.4±10.6 (7)
- 15	œ	<b>X</b> in	205.1±10.5(10) 137.4± 6.0(10)	204.3±10.7(10) - 137.8± 6.8(10)	211.8±10.4(10) 140.2± 4.4(10)	217.2±10.9(10) 146.3± 5.3(10)	221.1±11.2(10) 148.5± 6.7(10)
57A	0.35±0.05	X. in	205.1±11.1(10) 138.6± 8.2(10)	204.0±12.6(10) 136.7± 9.6(10)	210.7±12.7(10) 140.7±13.5(10)	216.0±12.8(10) 141.4±15.1(10)	219.3±12.4(10) 143.8±17.5(10)
578	0.99±0.07	ž u	205.8±12.0(10) 138.0± 7.0(10)	194.9±12.3(10) 135.1± 8.9(10)	198.1±10.4(10) 137.2± 8.6(10)	202.1±10.3(10) 139.3±10.6(10)	203.4± 9.8(10) 149.8±10.1(10)
65	0	z u	145.4±10.9(14) 127.4± 8.4(10)	149.1±11.5(10) 129.1± 7.6(10)	157.3±12.1(10) 132.9± 8.2(10)	162.8±11.7(10) 137.5± 7.3(10)	167.5±11.3(10) 140.1±7.1(10)
65A	1.56±0.10	X in	144. 3±10.7(10) 121. 9±12.0(10)	145.9± 9.3(10) 123.J± 8.4(10)	141.2±13.4(10) 121.6±10.0(10)	144.9±11.8(10) 125.3±11.1(10)	147.9±12.3(.0) 128.9± 9.0(10)
658	2.49±0.07	X W	140.8±16.0(10) 124.8±11.8(10)	131.1±16.2(10) 112.4±11.0(10)	125.5±15.8(10) 109.1±10.5 (8)	128.8±16.8 (8) 108.9±10.4 (8)	128.4±19.0 (8) 113.8± 9.7 (6)

(Page t as 3)

Table A-2 (Continued). MEAN BODY WEIGHTS<sup>a</sup> OF MALE AND FEMALE RATS<sup>b</sup> EXPOSED FIVE TIMES TO RP/BR

AEROSOLS AND MONITORED FOR 14 DAYS

Study No.	mg/ml Mean ± SD	Sex	6 7 8	7	8	10
52A	3.05±0.29	ж ч.			125.0 (1)	
528	1.5310.13	X u		•	204.9±20.9 (7) 171.1± 8.0 (7)	
27	O	X 14.		• • • •	223.7±29.8(10) 157.0± 7.6(10)	240.5±18.4(10) 161.4± 7.2(10)
57A	0.35±0.05	2 u. 1			209.9±38.3(10) 150.0±25.4(10)	232.2±22.2(10) 153.8±30.3(10)
578	0.940.07	X LL			219.4± 7.9(10) 145.4±20.4(10)	231.2± 8.1(10) 154.4±13.9(10)
65	o	30 km -	173.3±12.5(10) 142.3± 8.2(10)	181.7±12.9(10) 146.4± 8.7(10)	188.4±13.2(10) 149.1± 9.5(10)	200.5±14.5(10) 152.7± 9.6(10)
65A	1.56±0.10	II L	152.6±12.1(10) 131.2± 7.4(10)	162.4±12.7(10) 136.5± 8.4(10)	169.8±11.8(10) 137.9± 9.9(10)	-183.7±12.4(10) 146.4± 9.0 (9)
658	2.49±0.07	<b>1</b> 2 H.	135.2±19.8 (6) 116.2±10.8 (6)	144.8±21.6 (6) 121.7± 9.8 (6)	146.5±29.1 (6) 122.0±12.6(6)	149.3±32.6 (6) 135.2± 9.2 (5)

(Page 2 of 3)

Table A-2 (Continued). MEAN BODY WEIGHTS<sup>a</sup> OF MALE AND FEMALE RATS<sup>b</sup> EXPOSED FIVE TIMES TO RP/BR

DAYS
14
FOR
MON'I TORED
AND
<b>AEROSOLS</b>

No.	mg/ml Mean ± SD	Sex	12		15		17		19	
52A	3.05±0.29	2. L.	120.0	Ξ	139.0	Ξ	149.0	(1)	161.0	(Ξ)
528	1.99±0.13	X in	216.3±40.4 (7) 182.3±16.5 (6)	.4 (7) .5 (6)	238.0±43.2 (7) 177.7±32.9 (6)	2 (7) 9 (6)	250.7±40.3 (7) 190.2±20.5 (6)	() () ()	262.9±39.1 195.7±16.6	1 (7) 6 (6)
57	0	X 4.	251.3±17.1(10) 166.6± 7.1(10)	(01)1.	260.9±17.7(10) 171.7± 7.5(10)	7(10) 5(10)	266.7±17.2(10) 173.8± 9.8(10)	(01) (01)	273.0±17.4(10) 178.5±11.0(10)	4 (10) 4 (10)
57A	0.35±0.05	. X L.	244.8±19.6(10) 168.2±10.5 (9)	.6(10) .5 (9)	254.8±15.5(10) 175.8±12.5 (9)	5 (10) 5 (9)	265.4±15.5(10) 177.7±14.1 (9)	(01) (6)	271.6±15.6(10) 182.0±12.5 (9)	6(10) 5 (9)
578	0.99±0.07	X 4.	244.2± 8.8(10) 162.3±13.0(10)	.8(10) .0(10)	256.8± 8.9(10) 167.4±11.8(10)	(01) 8(10)	263.6±10.3(10) 172.9±12.3(10)	(01)	270.2±10.7(10) 175.3±13.7(10)	7(10) 7(10)
65	0	X L	211.2±13.7(10) 155.7±10.3(10)	.7(10) .3(10)	231.8±16.0(10) 163.8±11.3(10)	0(10) 3(10)	243.7±15.5(10) 168.8±13.4(10)	(01) (10)	255.6±16.8(10) 174.8±11.1(10)	8(10) 1(10)
65A	1.56±0.10	ĨL	196.2±12.6(10) 149.4± 9.4 (9)	.6(10). .4 (9).	217.0±12.6(10) 147.7±19.7 (9)	(01) 9 (6) 1	230.9±14.3(10) 157.4±12.6 (9)	(01) (6)	244.2±15.3(10) 166.3±12.9 (9)	3(10) 9 (9)
658	2.49±0.07	<b>X</b> 4	165.0±29.0 (6) 141.8± 9.2((5)	.0 (6) .2((5)	185.2±26.2 (6) 152.8± 8.6 (5)	(2)	198.5±28.8 (6) 153.6±17.4 (5)	3 (6) 4 (5)	211.5±28.9 (6) 160.8±16.9 (5)	9 (6) 9 (5)

(Page 3 of 3)

Exposure days.

Study Day I = first exposure day.

Data given as Mean ± SD (number of rats).

Rats were 6 to 8 weeks old when used. Mean ± 5D for gravimetric filter samples. Table A-3. MEAN BODY WEIGHTS<sup>a</sup> of Male and Female Rats During four daily exposures to RP/BR Aerosol

	KB/BK	KB/BK Aerosol	•			· .	
freatment	Conc.b	Duration	· · ·	Mean Bod	y Weights (g),	Mean Body Weights (g), Determined on Study Day	Study Day:
Group	(1/bu)	(Hr/day)	Sex	lc,d	2 <sup>d</sup>	9q	p†
>	0	0.1	<b>35</b> 14.	202.4±8.1 138.1±8.8	209.3±10.0 141.9± 7.9	215.0±8.9 142.4±8.7	219.9±9.3 145.1±8.3
11, 11	0.62±0.06	1.0	52 LL	201.8±6.8 137.0±6.9	205.2± 5.6 140.1± 6.7	210.2±6.3 141.7±7.3	214.4±7.5 145.0±6.0
117.11	o	3.5	x u	200.3±9.6 139.7±7.1	208.0± 8.1 143.4± 8.3	212.5±8.6 145.7±7.9	218.9±8.7 148.4±8.4
1117.71	0.50±0.02	3.5	12. Lu	200.4±9.1 141.3±6.6	203.4± 8.3 142.9± 5.9	206.3±8.5 144.3±5.3	211.7±9.7 148.2±6.9

<sup>a</sup> Mean ± SD from body weights of 20 male or female rats, respectively.

51

b Mean  $\pm$  SD calculated from the mean daily gravimetric filter determinations.

c Study Day 1 = first exposure day.

Exposure days.

Table A-4. MEAN BODY WEIGHTS<sup>a</sup> OF.MALE.AND FEMALE RATS DURING FOUR DAILY EXPOSURES TO RP/BR AEROSOLS

í

AND FOLLOWED BY 14 DAYS OF RECOVERY

	Aerosol	'е -	ı	. Mear	Mean Body Weights (g) Determined on Study Day:	i (g) Determin	ied on study Ve	:Ye
freatment Group	conc.b (mg/l)	Duration (hr/day)	Sex	lc,d	2 <sup>d</sup>	م م	r <sup>†</sup> q	5
>	0	1.0	E in	202.2± 8.5 137.2± 6.0	207.2±11.4 141.3± 6.4	213.6±10.0 140.9± 6.8	219.0± 9.6 143.9± 7.4	222.4±11.6 148.1± 7.6
1	0.62±0.06	1.0	£L	201.8± 8.9 133.5± 5.6	205.1± 7.0 136.3± 3.7	209.5± 8.3 137.1± 5.4	213.9± 9.6 141.0± 5.3	217.0±10.1 144.2± 5.2
111	0	3.5	X LL	204.2± 8.1 139.2± 7.6	211.5± 6.0 143.4±10.2	216.0± 5.0 146.2± 7.9	222.6± 6.1 149.5± 7.1	226.6± 5.3 152.7± 8.0
1117	0.50±0.02	3.5	، ۲۲ اس	199.6± 8.7 143.1± 7.5	201.9± 8.1 144.9± 5.4	204.8± 8.3 145.7± 5.0	210.4± 9.9 148.9± 7.4	212.7± 6.3 150.8± 6.7

(Page 1 of 2).

Table A-4 (Continued). MEAN BODY WEIGHTS<sup>a</sup> OF MALE AND FEMALE RATS DURING FOUR DAILY EXPOSURES TO

3

RP/BR AEROSOLS AND FOLLOWED BY 14 DAYS OF RECOVERY

Treatment	Aerosol ronc.b.	Ourarion			Mean Body We	ights (g) Det	Mean Body Weights (g) Determined on Study Day:	udy Day:	
Group	(1/6m)	(hr/day)	Sex	9.	8	=	14	16	18
	0	1.0	X 44	227.6±10.8 151.3± 8.3	238.5±13.8 155.0± 7.1	252.4±13.9 163.1± 8.8	267.8±17.1 168.5± 8.5	272.0±17.8 170.0±10.7	276.3±18.5 170.5± 8.7
17	0.62±0.06	0.	£ LL	223.3±10.7 147.8± 5.8	236.0±11.9 152.2± 6.2	252.7±15.5 160.4± 4.5	264.1±17.2 171.1± 8.7	269.9±17.7 173.1± 8.3	273.7±17.5 175.9± 8.8
A I I	o	3.5	X LL	232.9± 7.1 156.7± 6.5	245.3± 6.9 163.3± 8.5	258.7±13.4 168.8± 6.6	273.0±11.5 179.2± 7.9	278.0±11.0 180.4± 8.2	286.5± 9.3 :82.3± 7.7
<b>VIII</b>	0.50±0.02	3.5	× u	219.2±10.2 151.5± 7.4	230.2±10.7 161.6± 7.5	243.5±13.8 166.7± 9.5	255.7±12.6 170.9±17.2	259.3±12.6 172.9±14.6	265.6±14.7 174.2±16.7
A tean t Rean t	Mean ± SD from body weights of 10 male or female rats respectively. Mean ± SD for aerosol mass concentration calculated from daily mean gravimetric filter determinations.	weights of I mass conc	10 mal entrat	e or female r ion calculate	or female rats respectively. M calculated from daily mean	ely. mean gravimet	ric filter de	terminations.	(Paç 
1 ADRIC P	Study Uay 1 = Tirst exposure day.	exposure da	۲.				· ,		;e
Exposure davs.	e dave.		•						2

f 2)

MEAN BODY WEIGHTS<sup>a-</sup> of Male and Female Rats During Four Weeks of Exposure Four Days/Week to RP/BR Aerosol and two weeks of recovery Table A-5.

:

forr a	Duration		•	Mean Bc	rear body weights (g), betermined on study uay:	No nati titi la lan	areay way.		
(1/6m)	(hr/day)	Sex	1 p. c	5 <sup>c</sup> .	12 <sup>c</sup>	-19 <sup>c</sup>	26 <sup>c</sup> .	- 33 <sup>d</sup>	39 <sup>d</sup>
0	3.5	Ľ	199.2 ± 9.8	224.4 ± 10.1		281.2 ± 10.7	258.3 ± 10.3 281.2 ± 10.7 302.7 ± 12.3 328.2 ± 13.1	328.2 ± 13.1	345.3 ± 15.6
0.51 ± 0.04	35	£	199.3 ± 8.3	214.1 ± 9.1	244.9 ± 11.4	267.1 ± 13.9	14.1 ± 9.1 244.9 ± 11.4 267.1 ± 13.9 289.8 ± 16.7 315.8 ± 20.8 332.9 ± 20.1	315.8 ± 20.8	332.9 ± 20.1
	3.5	<b>اب</b>	138.3 ± 8.0	152.3 ± 8.2	170.3 ± 8.6	183.7 ± 10.7	52.3 ± 8.2 170.3 ± 8.6 183.7 ± 10.7 194.3 ± 13.3 205.6 ± 16.3 215.2 ± 18.3	205.6 ± 16.3	215.2 ± 18.3
6.51 ± 0.04	3.5	-	136.8 ± 8.2	146.1 ± 8.4	161.3 ± 8,7	176.5 ± 9.3	16.1 ± 8.4 161.3 ± 8.7 176.5 ± 9.3 189.8 ± 10.6 199.9 ± 13.0 268.8 ± 14.2	199.9 ± 13.0	208.8 ± 14.2

Date given as Mean ± SD for aerosol mass concertration calculated from daily gravimetric filter readings and for weights from 20 or 18 rats per group on Days 1 through 26 and Days 33 through 39, respectively.

Study Day i = first exposure day.

Exposure period.

Recovery period.

# PATHOLOGY REPORT PART I: GROSS OBSERVATIONS (IITRI)

÷ .

#### FOUR-DAY INHALATION TOXICITY STUDY OF RP/BR IN RATS:

Pathology and Pulmonary Lavage (Study 76/77)

#### PATHOLOGY SYNOPSIS

No compound-related gross pathologic lesions were observed in Sprague-Dawley rats from the experimental groups sacrificed after treatment which they received and sacrificed 14 days after their last exposure.

Microscopic examination of the tissues did not reveal any treatment-related changes in kidneys, trachea and nasal turbinates. However, lungs of several animals had focal areas of interstitial reaction with alveolar macrophages which may be treatment-related. It is not clear at this study period whether there was a possible reaction of marked lymphoid hyperplasia of pulmonary lymph nodes which suggests an infectious agent.

Vladiture S. Rac

Vladislava S. Rac, D.V.M., M.S. Head, Pathology Pathology Section Life Sciences Research

NECROPSY OBSERVATIONS

Introduction

In accordance with the experimental protocol, examination of organs was performed on 64 (32 male and 32 female) Sprague-Dawley rats for IITRI Project L6139, Study Number 76 A and B Phase II. The rats were divided into eight groups according to treatment received, length of exposure, and time of sacrifice.

#### IIT RESEARCH INSTITUTE

Thirty-two rats (16 males and 16 females) were exposed to 0.5 mg/l of RP/BR aerosol. Sixteen of the 32 rats (8 males and 8 females) were exposed for one (1) hour. The remaining 16 rats were exposed for 3-1/2 hours. Both groups were exposed to the aerosol once a day for four consecutive days.

The remaining 32 rats (16 males and 16 females) were exposed to filtered air. As with the aerosol, sixteen of the 32 rats (8 males and 8 females) were exposed for one (1) hour, while the remaining 16 were exposed for 3-1/2 hours. Both groups were exposed to the filtered air once a day for four consecutive days.

Eight rats (4 males and 4 females) from each exposure group (0.5 mg/l RP/BR and filtered air) and at each time exposure (1-hr and 3-1/2 hrs) were sacrificed and necropsied within four hours (0 days) after the last exposure (Exposure Group). The remaining 32 rats were sacrificed and necropsied 14 days after their last exposure (Recovery Group). The group number, treatment, number of animals per group per sex, corresponding exposure levels, time of exposure, and time elapsed before sacrifice are outlined below.

EXPOSURE GROUP	TREATMENT	NUMBER MALES	NUMBER FEMALES	EXPOSURE LEVEL (mg/l)	EXPOSURE TIME (hr)*	TEBS**
1	Filtered Alr Contro	51 4	4	0.0	1	0 days
11	RP/BR Aerosol	4	4	0.5	1	0 days
	Filtered Air Contro	51 4	4	0.0	3 1/2	0 days
1 V	RP/BR Aerosol	4	4	0.5	3 1/2	0 days
Υ V	Filtered Alr Contro	<b>b</b> 1 4	4	0.0	1	14 days
V 1	RP/BR Aerosol	4	4	0.5	1	14 days
V 1.1	Filtered Air Contro	1 4	4	0.0	3 1/2	14 days
VIII	RP/BR Aerosol	4	4	0.5	3 1/2	

\* Hours per day for 4 consecutive days \*\* TEBS + Time Elapsed Before Sacrifice

#### Materials and Methods'

The rats were sacrificed with carbon dioxide either within four hrs (0 days) or 14 days after the last exposure, bled from the dorsal aorta, and necropsied. The organs were examined and fixed in 10% neutral buffered formalin for a period no less than 48 hrs before further processing. The select organs from 32 of the 64 rats necropsied (2 males and 2 females from each of the eight treatment groups) were submitted to the histology laboratory and were processed according to the experimental design.

#### Results and Discussion

Mottled red lungs were observed in many of the rats in both the treated and filtered air control groups and in both the exposure and recovery groups. The small number of animals per group makes it difficult to determine the significance of this iesion.

Other lesions which occurred with some frequency were urinary calculi in males especially in the recovery group, mottled brown kidneys, and distended uterl. These and the remaining lesions observed were regarded as incidental findings and were present in both the control and treated animals.

60.

Caul a. Ihmp

Carol A. Thompson, D.V.M., M.S. Senior Veterinary Pathologist Pathology Section Life Sciences Research





1999年にいたいための調理をいたいた



63 .



INHALATION STUDY - KAIS Recovery Group (14 days)



2 34

≽

Ň



V

Ÿ



ŕ

Ĩ.

F

## FOUR-WEEK INHALATION TOXICITY STUDY OF RP/BR IN RATS

(Study 78)

#### NECROPSY OBSERVATIONS

No compound related gross pathologic lesions were observed in Sprague-Dawley rats from the experimental group sacrificed after 16 exposures at 0.5 mg/l for 3.5 hours, nor in rats sacrificed after two weeks recovery period. All lesions observed at necropsy were considered to be spontaneous in nature and unrelated to treatment with Red Phosphorus/Butyl Rubber Combustion Products.

Vlandighue S. Dae

Vladislava S. Rac, D.V.M., M.S. Head, Pathology Pathology Section Life Sciences Research

IIT RESEARCH INSTITUTE

Females  $\sim$  $\sim$ 3.5 HOUR EXPOSURE 0.5 mg/1 BP/BR Males I. ı 2  $\sim$ ı ŧ 1 Females FILTERED AIR CONTROL NECROPSY OBSERVATIONS Rats Sacrificed After 16 Exposures ł  $\sim$ . • Males 2 1  $\overline{\mathbf{O}}$ Blue ingesta present in the lumen Scattered red foci URINARY BLADDER Mottled red Mottled red Distended ORGANS Lesions Calculi KIDNEYS STOMACH UTERUS L6139 SN 78 LUNGS

,

ee j
Females I . 1 3.5 HOUR EXPOSURE 0.5 mg/l BP/BR Males ŧ ł  $\sim$ 1 ~ Females FILTERED AIR CONTROL 1 1 NECROPSY OBSERVATIONS Recovery Sacrifice C Males , Blue ingesta present in the lumen Scattered red foci URINARY BLADDER Mottled red Mottled red Distended ORGANS Lesions Calculi KIDNEYS **STOMACH** UTERUS LUNGS L6139 SN 78 • ۰.

## PATHOLOGY REPORT PART II: HISTOPATHOLOGY (EPL)

EXPERIMENTAL PATHOLOGY LABORATORIES, INC.

P 1921-942-6-

) }.

イント しょうじ

EPL

### IITRI PROJECT NUMBER LO6139 PHASE II STUDY 76/77

### FOUR-DAY INHALATION TOXICITY STUDY OF RP/BR IN RATS: PATHOLOGY AND PULMONARY LAVAGE

PATHOLOGY REPORT

### Submitted to:

IIT Research Institute Chicago, Illinois 60616

September 9, 1983

# EPL

### EXPERIMENTAL PATHOLOGY LABORATORIES, INC.

TABLE OF CONTENTS

	Page
PATHOLOGY SUMMARY	, 1
SUMMARY INCIDENCE TABLES	
Males	8 10
HISTOPATHOLOGY INCIDENCE TABLES	. •
Males	14 19
CORRELATION OF GROSS AND MICROSCOPIC FINDINGS TABLES	
Males	24

# Page 1 (EPL)

PATHOLOGY SUMMARY

11、「ないないという」」

C

### Page 2 (EPL)

### EXPERIMENTAL PATHOLOGY LABORATORIES, INC.

EPL

### IITRI PROJECT NUMBER LO6139 PHASE II STUDY 76/77

### FOUR-DAY INHALATION TOXICITY STUDY OF RP/BR IN RATS: PATHOLOGY AND PULMONARY LAVAGE

### PATHOLOGY SUMMARY

Microscopic examinations were performed on selected tissues from male and female Sprague-Dawley rats. The study was designed to determine the toxicity of RP/BR (Red Phosphorus/Butyl Rubber) combustion products administered by multiple inhalation exposure as evaluated by histopathology and pulmonary lavage immediately and after a 14-day recovery period. This report contains the histopathologic findings. The experimental design for this study was as follows:

Exposure Group	Group Name	Conc.	Exposure*	No. Animal Male Feral	-
I - 1 hr.	Filtered Air Control (O hour)	<b>0</b>	1 hour	14 14	
II - 1 hr.	RP/BR (O hour)	.5 mg/1	1 hour	14 14	•
111 - 3.5 hr.	Filtered Air Control (O hour)	0	3.5 hou <del>r</del>	14 14	۱.
IV - 3.5 hr.	RP/BR (0 hour)	.5 mg/1	3.5 hour	14 14	
	Filtered Air Control (14 day)		1 hour + 14 day recovery	14 14	,
¥1. → 1 hr.	RP/BR (14 day)	.5 mg/1	l hour + 14 day recovery	14 14	
	Filtered Air Control (14 day)	0	3.5 hour + 14 day recovery	14 14	
VIII - 3.5 hr.	RP/BR (14 day)	.5 mg/1	3.5 hour + 14 day recovery	14 14	•

\*All exposures are for four (4) consecutive days.

-

Page 3 (EPL)

### EXPERIMENTAL PATHOLOGY LABORATORIES, INC.

According to protocol, two rats per sex per group were necropsied and gross and histologic evaluations of the respiratory tract were conducted. Gross examination of the kidneys from several animals revealed mottling. Therefore, the following tissues were trimmed and processed to paraffin blocks: trachea, pulmonary lymph nodes, each lung lcbe, nasal turbinates, and kidneys. The paraffin blocks were then shipped to Experimental Pathology Laboratories, Inc. where hematoxylin and eosin stained slides were prepared and examined.

### RESULTS

EPL

The microscopic changes and a detailed listing of all tissues evaluated are presented in the Histopathology Incidence Tables. All lesions are summarized by sex and treatment group and presented in the Summary Incidence Tables. A correlation of lesions observed at necropsy with the corresponding microscopic observation, where possible, is presented in the Correlation of Gross and Microscopic Findings Tables. The gross observations in these tables were transcribed from the necropsy sheets provided with the paraffin blocks.

Several lesions were found in the lung and pulmonary lymph nodes of a few male and female recovery animals that may be treatment related. No lesions at all were detected in the nasal turbinates and only a few animals had focal areas of epithelial hyperplasis in the traches.

, Page 4 (EPL)

### EXPERIMENTAL PATHOLOGY LABORATORIES, INC.

EPL

Changes were found in pulmonary lymph nodes, lung, and kidney in both control and treated animals. Minimal to mild lymphocyte hyperplasia and occasionally hemorrhage were found in the pulmonary lymph nodes of most animals. Moderate to marked lymphocyte hyperplasia was usually seen in association with multifocal accumulations of macrophages, lymphocytes, and interstitial thickening in the lung. These changes were more severe in one Group VI male and one Group VI female and in both Group VIII females, but were also present in Group V and Group VII (control) animals. This pulmonary interstitial reaction could be related to the test substance, but the lesions are also compatible with a subclinical Sendai virus or Mycoplasma pulmonis infection. The marked lymphoid hyperplasia of the pulmonary lymph nodes is indicative of intense antigenic stimulation and makes the latter possibility more likely. Renal changes seen in a few animals consisted of minimal to mild tubular hyperplasia, dilatation, and/or lymphocytic infiltrate. These changes are probably the very earliest lesions of chronic progressive nephropathy, a spontaneous syndrome that commonly occurs in laboratory rats.

One interesting renal finding was a nephroblastoma, a malignant embryonal neoplasm, in rat Number 116 which received four one-hour exposures to .5 mg/l RP/BR and was then necropsied. This large tumor was certainly present before the initiation of the experiment and was probably congenital. It is in no way treatment related.

Page 5 (EPL)

### EXPERIMENTAL PATHOLOGY LABORATORIES, INC.

### CONCLUSION

EPL

The results of this microscopic examination indicate that administeration of RP/BR at the concentrations and for the duration of exposure used in this study did not produce treatment-related changes in the kidney, trachea, or the nasal turbinates at the levels examined.

Several animals which recovered for 14 days following either a 1 hour or 3.5 hour exposure (for four days) to RP/BR combustion products had focal areas of interstitial reaction with alveolar macrophages which may be treatment related. However, these animals also had moderate to marked lymphoid hyperplasia of pulmonary lymph nodes which suggests an infectious agent such as Sendai virus or Mycoplasma pulmonis may have produced these lesions.

77

W. O. IVERSON, D.V. M. Diplomate, ACVP

W01/sfh

-----

# SUMMARY INCIDENCE TABLES

Page 6 (EPL)

Page 7 (EPĹ)

### EXPERIMENTAL PATHOLOGY LABORATORIES, INC.

EPL

### QUALITY ASSURANCE REPORT CERTIFICATION

Client Name 1	IT Research Institute	Client Study No.	76/77
Study Director	Dr. William O. Iverson	Pathologist <u>Dr. Will</u>	iam O. Iverson
Study Title	Four-Day Inhalation Tox Pathology and Pulmonary		in Rats:

Test Article Red Phosphorus/Butyl Rubber Species Sprague-Dawley Rats

All parts of the pathology phase of this study, including the final report, were reviewed by Experimental Pathology Laboratories Quality Assurance Unit on <u>June 20 - September 9, 1983</u>. All findings were reported to the Study Director and Management.

Janet Milazio 9/9/83

SUMMARY INCIDENCE TABLE

L06139 Phase II Study 76/77 4 Day Sacrifice Male Rats

I amount reserves at Participation & Specie reference beie

	Group I	Group II	Group III	Group IV		
TRACHEA (NO. EXAMINED)	(2)	(2)	. (2)	(2)		
Focal Hyperplasia					ł	
PULMONARY LYMPH NODES		1				1
(NO. EXAMINED)	(2)	(2)	(2)	(2)	,	
Lymphoid Hyperplasia	2	1	2	2		
Hemorrhage	1					
Macrophage Hyperplasia		1	•	1		] <u>.</u>
Edema						
LUNG (NO. EXAMINED)	(2)	(2)	(2)	(2)		 
Hemorrhage	1	1		1		
Atelectasis	•	2	2	2		· ·
Alveolar Macrophages		1				 
Focal Lymphocyte Aggregate			1			 •
Interstitial Thickening		I				 
				······		
MASAL TURBINATE (NO. EXAMINED)	(2)	(2)	(2)	(2)		·
		· · · · · · · · · · · · · · · · · · ·				
KIDNEY (NO. EXAMINED)	(2)	(2)	(2)	(2)		
Tubular_Hyperplasia			<b>1</b>			
Lymphocytic Infiltrate			·	•		
<u>Congestion</u>					······	. <b></b>
·	· · · · · · · · · · · · · · · · · · ·				· · · · · · · · · · · · · · · · · · ·	-
		9				
		·····				
		·		· · · · · · · · · · · · · · · · · · ·		
				• •		

Page 8 (EPL)

Page 9 (EP

### SUMMARY INCIDENCE TABLE

LO6139 Phase II Study 76/77 14 Day Recovery Sacrifice Male Rats

minumal Pathology Laboratorie

	Group V	Group VI	Group VII	Group VIII		
TRACHEA (NO. EXAMINED)	(2)	(2)	(2)	(2)		1
Focal Hyperplasia		1				
PULMONARY LYP. I NODES						
(NO. EXAMINED)	(2)	(2)	(2)	(2)		
Lymphoid Hyperplasia	2	2	2	2		
Hemorrhage	·	1	1	1		
Macrophage Hyperplasia		1	2	_1		
Edema		1	2	1		
			·			
LUNG (NO. EXAMINED)	(2)	(2)	(2)	(2)		
Hemorrhage	2	· 2	1	2		
Atelectasis	1	1	, 2	1		
Alveolar Macrophages	1	1				
Focal Lymphocyte Aggregate	1	1			ı	_
Interstitial Thickening		. 1				
NASAL TURBINATE (NO. EXAMINED)	(2)	(2)	(2)	(2)		
KIDNEY (NO. EXAMINED)	(2)	(2)	(2)	. (2)		
Tubular Hyperplasia	1	2	2	2 ·	· · ·	
Tubular Dilatation		1	.1			
Lymphocytic Infiltrie		1			· .	
Congestion		· .		1		. 
-						
			· · · · · · · ·		: • • • • • • • • • • • • • • • • • • •	
					. ,	1

Page 10 (EPL)

## SUMMARY INCIDENCE TABLE

LO6139 Phase II Study 76/77 4 Day Sacrifice Female Rats

	Group I	Group II	Group III	Group IV		
TRACHEA (NO. EXAMINED)	(2)	. (2)	(2)	(2)		1.
Focal Hyperplasia						
	:					1
PULMONARY LYMPH NODES					1	
(NO. EXAMINED)	(2)	(2)	(2)	(2)		· · · · · · · · · · · · · · · · · · ·
Lymphoid Hyperplasia	1	2	2	1		
Hemorrhage	1	1	1			
Macrophage Hyperplasia	1			,		
Edema						
Yellow-Green Pigment		·				
						÷
LUNG (NO. EXAMINED)	(2)	(2)	(2)	(2)		L
Atelectasis	2	1	1	1		
Alveolar Macrophages					· · · · · · · · · · · · · · · · · · ·	
Focal Lymphocyte Aggregate				1		
Interstitial Thickening				•		
NASAL TURBINATE (ND. EXAMINED)	(2)	(2)	(2)	(2)	-	
				•	· ·	· · · · ·
KIDNEY (NO. EXAMINED)	(2)	(2)	(2)	(2)		
Nephroblastoma		1				
Tubular Hyperplasia		·				· · · · · · · · · · · · · · · · · · ·
Lymphocytic Infiltrate						
				,		
				· · · · · · · · · · · · · · · · · · ·	•	· · .
	· · · · · · · · · · · · · · · · · · ·					
	· · · · · · · · · · · · · · · · · · ·			· · · · · · · · · · · · · · · · · · ·	· · · · ·	
	د. مراجع محمد محمد الامر			• • • • • •		
					· · ··-	
	·					

Page 11 (EPL

### SUMMARY INCIDENCE TABLE

L06139 Phase II Study 76/77 14 Day Recovery Sacrifice Female Rats

	Group V	Group VI	Group VII	Group VIII		
TRACHEA (NO. EXAMINED)	(2)	(2)	(2)	(2)		
Focal Hyperplasia		· · · · · · · · · · · · · · · · · · ·		1		
PULMONARY LYMPH NODES		к н Стала стала стал		· · · · · · · · · · · · · · · · · · ·		
(NO. EXAMINED)	(2)	(2)	(2)	(2)		
Lymphoid Hyperplasia	2	2	2	2		
Hemorrhage		1		1		·
Macrophage Hyperplasia	1	2		2		
Edema	1					
Yellow-Green Pigment	,	1		1		
	(2)	(2)		(2)		
LUNG (NO. EXAMINED)	(2)	(2)	(2)	(2)		
Atelectasis	1		2		· · · · ·	
Alveolar Macrophages	·····	1		2		
Focal Lymphocyte Aggregate	·	1		2	   	
Interstitial Thickening		1		2		
			· · ·			
ASAL TURBINATE (NO. EXAMINED)	(2)	(2)	(2)	(2)		. <b>.</b> .
(IDNEY (NO. EXAMINED)	• (2)	(2)	(2)	(2)		
Nephroblastoma						
Tubular Hyperplasia	+		1		· · ·	
Lymphocytic Infiltrate	••••••••••••••••••••••••••••••••••••••	• • • • • • • • • • • • • • • • • • •	1	1		
						·
			· ·			••••
		1				
		· · · · · · · · · · · · · · · · · · ·			,	
						- • `,
Pi l		• _ •••• •	<del></del>			

HISTOPATHOLOG

HISTOPATHOLOGY INCIDENCE TABLES

Page 13 (EPL)

,

Exposure Group	Group Name	Conc.	Exposure*
I - 1 hr.	Filtered Air Control (O hour)	0	1 hour
II – 1 hr.	RP/BR (O hour)	. 5 mg/1	1 hour
III - 3.5 hr.	Filtered Air Control (. hour)	0	3.5 hour
IV - 3.5 hr.	RP/BR (O hour)	. 5 mg/1	3.5 hour
V - 1 hr.	Filtered Air Control (14 day)	0	1 hour + 14 day recovery
VI - 1 hr.	RP/BR (14 day)	. 5 mg/1	l hour + 14 day recovery
VII - 3.5 hr.	Filtered Air Control (14 day)	0	3.5 hour + 14 day recovery
VIII - 3.5 hr.	RP/BR (14 day)	. 5 mg/1	3.5 hour + 14 day recovery

\*All exposures are for four (4) consecutive days.

Lubel 39 Phase 11 Study 76/77 A Day Sacrifice Male Rats Male Male Myperplasia Macrophage Myperplasia Edema	p 1 Croup 11 Croup 11 A 4 0 B 0 A 4 0 Croup 11		Group Group 2 X X 4 4 0 1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	>	
ase ii 77 77 77 6000 78 78 89 289 289 289 280 211 396 Hyperplasia 396 Hyperplasia			Graup 2 2 X X 8 0 1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2		
rifice rifice					
a sia 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2					
x X Sia					
a 2 sia					
a 2 2 3 5 1 a					
a 2 sia					
2 1a					
e					<u></u>
Macrophage Hyperplasia Edema					
Edema	2		<b></b>		Ţ
					, 
TUNG X					Ţ
Hemorrhage 2					
Atelectasis	2 2		2 1		
Alveolar Macrophages					
Focal Lymphocyte Aggregate		<b>F-1</b>			
Laterstitial Thickening					
1.1		>			Pag
-				·	ge I.
					14
	-0-				(EP ] ]

X = Not Remarkable

HISTOPATHOLOGY WCIDENCE FABLE HISTOPATHOLOGY WCIDENCE FABLE Suedy 10/11 Suedy 10	<b>,</b>	•					-				•	,	· · · ·	•	•	• ,			•			•	•		•	•••	Pac	e e	15	 (EP	L)	
HISTOPATHOLOGY INCLOENCE TABLE HISTOPATHOLOGY INCLOENCE TABLE ULOP 76/11 ULOP 76/11 UL	÷				ļ	· · · · · ·									<u> </u>		<u> </u>	<b></b>				ļ				ļ		ļ		]		
HISTOPATHOLOGYNKIDENCE TABLE 66139 Phase II tudy 36/17 66139 Phase II tudy 36/17 tudy														 	<u> </u>												$\left  \right $				ere/Hig	
HISTOPATHOLOGYNKIDENCE TABLE 66139 Phase II tudy 36/17 66139 Phase II tudy 36/17 tudy					ļ										ļ	<u> </u>				-							<u> </u>				narkable itely Sev	
HISTOPATHOLOGYNKIDENCE TABLE 66139 Phase II tudy 36/17 66139 Phase II tudy 36/17 tudy	÷.								10	4	×			╞──					<b> </b>		Z	. 						+			Not Rei Modera	
IHSTOPATHOLOGY INCIDENCE TABLE       IHSTOPATHOLOGY INCIDENCE TABLE       G6139 Phase 11       G013 Phase 11       Colspan="2">G013 Phase 11       Colspan= 2       Colspan= 2 <td co<="" td=""><td></td><td>·</td><td></td><td></td><td>Grou</td><td></td><td></td><td></td><td>8</td><td>8</td><td>×</td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td><u> </u></td><td></td><td></td><td></td></td>	<td></td> <td>·</td> <td></td> <td></td> <td>Grou</td> <td></td> <td></td> <td></td> <td>8</td> <td>8</td> <td>×</td> <td></td> <td><u> </u></td> <td></td> <td></td> <td></td>		·			Grou				8	8	×																	<u> </u>			
HISTOPATHOLOGYINCIDENCE TABLE 66139 Phase 11 tudy 36/17 tudy 36/17 tudy 36/17 tudy 36/17 tudy 36/17 tudy 36/17 tudy 36/17 tudy 14 tudy 36/17 tudy 14 tudy 36/17 tudy 14 tudy 14 t				1.												-		' 	-						   .		┨		-		el.	
HISTOPATHOLOGY INCIDENCE TABLE HISTOPATHOLOGY INCIDENCE TABLE 66139 Phase 11 60139 Phase 11 60139 Phase 11 60139 Phase 11 60131 6013 60 6 7 6 6 7 7 7 7 7 7 7 7 7 7 7 7 7 7	Ŧ	•	•	•						-																				1	Autolys Modera	
HISTOPATHOLOGY INCIDENCE TABLE HISTOPATHOLOGY INCIDENCE TABLE 06139 Phase 11 tudy 76/17 100 Jay Sacrifice 100 KT 100 KT 100 Jay Sacrifice 100 KT 100 Jay Sacrifice 100 KT 100 KT		,																			. 	. 	ļ			Í					1 H	
IHISTOPATHOLOGY INCIDENCE TABLE         IHISTOPATHOLOGY INCIDENCE TABLE         G6139 Phase 11         G6139 Phase 11         G6139 Phase 11         Day Sacrifice         A model         A model         Day Sacrifice         A model						•		-	• 0	2	X																				, E	
Initial State     Initial State     Initial State     Initial State       06139 Phase 11     Page 11     Page 11     Page 11       1493 76/17     Page 11     Page 11     Page 11       Page 11     Page 12     Page 12     Page 14       Page 11     Page 14     Page 14     Page 14       Page 11     Page 14     Page 14     Page 14       Page 14					Grou				8	6		1			-			 		 	[						 	ļ		·	n e Sectic	
Initial Phase 11     Initial Phase 11       06139 Phase 11     06139 Phase 11       1uudy 76/17     06139 Phase 11       06139 Phase 11     06139 Phase 11       0613 Phase 11     06139 Phase 11       0613 Phase 11     06139 Phase 11       0613 Phase 11     0       0613 Phase 11     0       0614 Phase 11     0       1004 Phase 11     0       11     0       0613 Phase 11     0       11     0       0613 Phase 11     0       11     0       0613 Phase 11     0       11     0       11     0       0613 Phase 11     0       11     0       11     0       11     0       11     0       11     0       11     0       11     0       11     0       11     0       11     0       11     0       11       11		ЭLЕ						<u> </u>	• •														<b> </b>				-		<u> </u>		Sectio pht complet	
66139 Phase 11 tudy 76/17 tudy 76/17 Bay Sacrifice ale Rats Tubular Dilatation LEUM HYMUS HYMUS RIMARY BLADDER RIMARY BLADDER RIMARY BLADDER RIMARY BLADDER	Ċ	E TA				·			,												ļ					<u> </u>			 	<b>]</b> .	P P 8	
66139 Phase 11 tudy 76/17 1095 Shase 11 tudy 76/17 1095 Sacrifice ale Rats Tubular Hyperplasia Tubular Hyperplasia Tubular Dilatation LEUM LEUM LEUM LEUM LEUM RINARY BLADDER RINARY BLADDER RINARY BLADDER RINARY BLADDER RINARY BLADDER		ENC			Ī			<b></b>																 					 			
66139 Phase 11 tudy 76/17 1095 Shase 11 tudy 76/17 1095 Sacrifice ale Rats Tubular Hyperplasia Tubular Hyperplasia Tubular Dilatation LEUM LEUM LEUM LEUM LEUM RINARY BLADDER RINARY BLADDER RINARY BLADDER RINARY BLADDER RINARY BLADDER		NCID			1															 	<u> </u>								ļ	-10	arnt mat øre/Higi	
06139 Phase 11 tudy 76/17 tudy 76/17 Bay Sacrifice ale Rats Tubular Pilatation LEUM HYWUS HYWUS RIMARY BLADDER RIMARY BLADDER RIMARY BLADDER	•	GΥ ΙΙ	·		2 S		,		<b>œ</b>	4	×		 	   	ľ. 1	 			Z	 		 		 		 			ļ			
06139 Phase 11 tudy 76/17 tudy 76/17 Bay Sacrifice ale Rats Tubular Pilatation LEUM HYWUS HYWUS RIMARY BLADDER RIMARY BLADDER RIMARY BLADDER		10LC			ŀ										i	<b> </b>		   									   '	<b> </b>	<b> </b>		<b>C - 10</b>	
66139 Phase 11 tudy 76/17 1095 Shase 11 tudy 76/17 1095 Sacrifice ale Rats Tubular Hyperplasia Tubular Hyperplasia Tubular Dilatation LEUM LEUM LEUM LEUM LEUM RINARY BLADDER RINARY BLADDER RINARY BLADDER RINARY BLADDER RINARY BLADDER		PATH										 	 		 	 	1	' 			 		 			 			1		×,	
66139 Phase 11 tudy 76/17 1095 Shase 11 tudy 76/17 1095 Sacrifice ale Rats Tubular Hyperplasia Tubular Hyperplasia Tubular Dilatation LEUM LEUM LEUM LEUM LEUM RINARY BLADDER RINARY BLADDER RINARY BLADDER RINARY BLADDER RINARY BLADDER		ISTO										·			1		   	   					j			 						
06139 Phase 11 tudy 76/17 Day Sacrifice ale Rats Tubular Hyperplasia Tubular Dilatation Lymphocytic Infiltrate Congestion LEUM RINARY BLADDER RINARY BLADDER		Ţ			roup			<u> </u>						! •	 												 					
06139 Phase 11 tudy 76/17 Day Sacrifice alle Rats IDNEY IDNEY IDNEY IDNEY IDNEY IDNEY IDNEY IDNEY IDNEY IDNEY IDNEY IDNEY IDNEY IDNEY IDNE IDNEY IDNE IDNEY IDNEY IDNEY IDNE IDNEY IDNE IDNE IDNE IDNE IDNE IDNE IDNE IDNE					١	z	⇒ <b>≯</b>	. co w					.	,  !	  `		.			· ·							r 					
06139 P tudy 76 tudy 76 Day Sa Lympho Lympho LEUM LEUM RINARY RINARY						`<	Ż -	∑ <b>⊀</b>				; ; ;										•						  .		•		
06139 P tudy 76 Day Sa Day Sa Ibula Tubula LEUM LEUM RINARY RINARY		· .				1	·																									
06139 7 tudy 76 Day 5a Bay 5a ale Rat Tubula Lubula Lubula Lubula Lubula Lubula Lubula Lubula LEUM						,		· , ·						te																	v lette	
06139 P tudy 76 Day Sa Bay Sa IbhEy Tubula Leum Leum RINARY RINARY			•	• 2								asia	noi	iltra													: : :		· .			
06139 7 tudy 76 Day 5a Bay 5a ale Rat Tubula Lubula Lubula Lubula Lubula Lubula Lubula Lubula LEUM					•	ce						erple	atat	Inf		 					ER										1.1 Ja.1	
06139 7 tudy 76 Day 5a ale Rat IbhEY LEUM LEUM LEUM RINARY RINARY	<b>.</b>	•		1356	. 111		· ·					- Hyp	- Dil	:ytic	tion						JL ADD		а - 4 л								HARDEN J.	
			ι.	39 PI	y 7.61	y Sac Rats		1			£Υ	bular	bular	mphoc	ngest		z		US			м, -							- -			
			•	1001	Stud	4 Da	1 				KI0N	10	Tu	Ly.	C,		ILEU		WAHI		URIN	•					· .			· ·		

6

Ĩ

ij.

,ŧ

• 2

.

•

	3	Group V				eroup					
14 Uay Recovery Sacrifice Male Rats	z⊃\$. ≺z-										
	<u> </u>		· · · · · · · · · · · · · · · · · · ·	2 0 L 8 0		0.4	0 1 1		9 1 6 2		
TRACHEA	×	×		×		×	~		×		
Focal Hyperplasia		1		2							
PULMONARY LYMPH NODES					-						
Lymphoid Hyperplasia				1 4		2	1		2 2		
Henorrhage				-1			1		-		
Macrophage Hyperplasia				2		2	1		2		
Edema				1		2	1				
t ung						<b> </b>					
kemorrhage				1			2		-		
Atelectasis		2		2		-	2		-		
Alveolar Macrophages				2							
Focal Lymphocyte Aggregate			'	3							
laterstitial Thickening				<b>6</b>							
NASAL TURBINATE	X	×		××		×	><	·	×		_
					-						
				-11-	,						EPL
			Key	P = Prrson	4 1	Ē		X Sis	1	emerkable	
			×		2 Z 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2						z

.•\*

n n

.

• .

Page 17 (EPL) C X = Not Remarkable 4 = Moderately Severe/High Group VIII 1 2 z 9 1 2 A = Autolysis 3 = Moderate þ C Group VII 10 2 2 Z Z N = No Section 2 = 514ht 1 = Incomplete Section 0 4 ---Z , HISTOPATHOLOGY INCIDENCE TABLE L Group VI P = Present 1 = Mountal 5 = Severe/High -12-108 ----5 2 و مليه <u>-----</u> Z z . . XeY 1 Group V × **6**0 ---z Ż **نه 4** Ş ہ ک kalarisissessies ilistiiriingis fukuristeregs, tere LO6139 Phase II Study 76/77 14 Day Recovery Sacrifice Male Rats Lymphocytic Infiltrate Tubular Hyperplasia **Tubular** Dilatation URINARY BLADDER Congestion SUMYHT X IDNE X ILEUN ....

## Page 18 (EPL)

Exposure Group	Group Name	Conc.	Exposure*
I - 1 hr.	Filtered Air Control (O hour)	C	1 hour
II - 1 hr.	RP/BR (O hour)	. 5 mg/1	1 hour
III - 3.5 hr.	Filtered Air Control (O hour)	0	3.5 hour
IV 3.5 hr.	RP/BR (O hour)	. 5 mg/1	3.5 hour
V - 1 hr.	Filtered Air Control (14 day)	0	l hour + 14 day recovery
VI - 1 hr.	RP/BR (14 day)	. 5 mg/1	1 hour + 14 day recovery
VII - 3.5 hr.	Filtered Air Control (14 day)	0	3.5 hour + 14 day recovery
VIII - 3.5 hr.	RP/BR (14 day)	. 5 mg/1	3.5 hour + 14 day recovery

\*All exposures are for four (4) consecutive days.

• Page 19 (EPL) X = Not Remarkable 4 = Moderately Severe/High Group IV З × Ó × --+ - 6 × × HN0 × -A = Autolysis 3 = Moderate Group III 4 3 4 × 2 -----× N = No Section 2 = Slight I = Incomplete Section × × 8 1 8 × ---+ -HISTOPATHOLOGY INCIDENCE TABLE P = Present 1 = Muumal 5 = Severe/High -13-Group 11 2 × 10  $\sim$ × 2 Q × ----\*\*\*\* × × ----Key Group 1 ~ × ≍ 3 0 ----× × 0 ----4  $Z \supset \Sigma$ @ w @ Lynumental Particology Laboratories, inc. < < z 5 Focal Lymphocyte Aggregate Interstitial Thickening Macrophage Hyperplasia Alveolar Macrophages Lymphoid Hyperplasia **fellow-Green** Pigment PULMONARY LYMPH NODES Focal Hyperplasia 4 Day-Sacrifice L06139 Phase 11 NASAL TURBINATE Atelectasis Hemorrhage Female Rats Study 76/77 Edema TRACHEA UNG. 123

Page 20 ([FL])         4 Baje 20 ([FL])	L06139 Phase 11 Study 76/77	,	Group 1				G	Group	11			9	Group	111		ł	ł	Group	VI di	_	ł		ſ
KIJNET       X <th>4 Bay Sacrifice Female Rats</th> <th></th> <th>۰,</th> <th></th> <th></th> <th></th> <th></th> <th></th> <th></th> <th><u>.</u></th> <th></th> <th></th>	4 Bay Sacrifice Female Rats													۰,							<u>.</u>		
KUNEY     X																							
phrohlastona Bullar Hyperplasta Maphozytic Infiltrate UUS WAGH WACH MACH MACH MACH MACH MACH MACH MACH M	KIDNEY	,											n ×	× t					° ×	+			<u> </u>
Bullar typerplasia Sultar typerplasia mphocytic lafilitrate us us us us voltar Meridia Mathematical Meridia M	Nephroblastoma	· · ·			<u> </u>			d						1				ļ				 	
mphocytic Infiltrate     I     I     I     I     I     I     I       US     US     I     I     I     I     I     I     I       US     I     I     I     I     I     I     I     I       US     I     I     I     I     I     I     I     I       US     I     I     I     I     I     I     I     I       AGN     I     I     I     I     I     I     I     I       AGN     I     I     I     I     I     I     I     I       AGN     I     I     I     I     I     I     I     I       AGN     I     I     I     I     I     I     I     I       AGN     I     I     I     I     I     I     I     I       AGN     I     I     I     I     I     I     I     I       AGN     I     I     I     I     I     I     I     I       AGN     I     I     I     I     I     I     I     I       AGN     I     I     I	Tubular Hyperplasta			<u> </u>	İ																		
ULS ULS ULS ULS ULS ULS ULS ULS	Lymphocytic infiltrate		   					   															
								<u> </u>															
uls	UTERUS		-			1294 1		 						<b> </b>					z	<u> </u>			ĺ
UIS						. 												, 					
AACH AACH 	EHYMUS							ļ		<u> </u>				i 									
AACH								 										-					
	STOMACH					<u>                                      </u>	]   		]														
-Id- -Id-			  .						<u> </u>									<b> </b>					· 
				 	<b></b>			<u> </u>					<b> </b>		 		 			 			
-14- -14-			. 	 		 	 											. 					
				ļ																			
														•									
			   ·																				
				] 							 			. 				ļ			  .		
-14- Key P = Present N = No Section A = Autolysis X = Not Remarkable																							
-14- Key P = Present N = No Section A = Autolysis X = Not Remarkable																							
Key P = Present N = No Section A = Autolysis X = Not Remarkable									4-		,												(
	-0-0-0-00		,	•.			1	tuoto		1	No Co				1	Autoto	đ				ohla		L

5

Ċ

C

Page 21 (EPL) X – Nof Remarkable 4 – Moderately Severe/Hign Group VIII 4 1 4 2 ---- $\sim$ 2 З - $\times$ -8 4.1 × 4 3  $\sim$ e × A m Autolysia A m Muderati , IIV JUDA 4  $\sim$ × 2 ----× \*\*\*\* N = No Section 2 = Sector 1 = Incomplete Section 109 × **,....** × . . , -15-Croup V: Key: P = Prevent 1 = -Minumal 5 = Severe/High 4 ò × e i ,..... c <del>, .....</del> i-d <del>dire</del>t × \*\*\*\* , ----2 4 ж 2 -----344 × Group Y ~ 3 ĊΦ, ≍ ŝ × × ~ -× × N ~ 2 D S @ W E Experimental Pathology Eaboratories, Inc. \$ « J -< 2 Focal Lymphocyte Aggregate 14 Day Recovery Sacrifice Interstitial Thickening Macrophage Hyperplasia Lymphoid Hyperplasia Yellow-Green Pigment Alveolar Macrophages PULMONARY LYMPH NODES Focal Hyperplasia 06139 Phase 11 NASAL TURBINATE Atelectasis Hemorrhage Study 76/77 Fenale Rats Edema TRACHEA LUNG EPL 33 •

.₹

L06139 Phase II Study 76/77	Group V	Group VI	Group VII		Group VIII
14 Day Recovery Sacrifice Female Rats	2320	•			
	8 3 F	- 2 4	2 F		2 F F
KIDNEY	XX	XX			X
Nephrobl as toma					
Tubular Hyperplasia					
Lymphocytic Infiltrate					
UTERUS			1		
	•		· · · · · · · · · · · · · · · · · · ·		
THYMUS	Z				
STOMACH			N		
			1		
				· ·	
	-	-16-		-	
		- Present Mormal	N = No Section 2 = Shohl	A Autolysis 3 - Moderate	X = NotRemarkable 4 = Modaratalo Savara/Horb
Expressmental Patinslergy Eaboratories, inc.	Hes, HC.	5 Sovere/High		ŧ.	

## Page 23 (EPL)

# CORRELATION OF GROSS AND MICROSCOPIC FINDINGS TABLES

,

L06139 Phase 11 Study 76/77 4 Day Sacrifice

I.

CORRELATION OF GROSS AND MICROSCOPIC FINDINGS

.1

					-Dosage Level
Animal Number	Client's fissue	Client's fissue Identification	ō	Client's Gross Observations	Microscopic Observations
98	Lungs		Red		llemorrhage
-					
	· · ·				
				-	
					age
1					
		•			
· · ·		•		-17-	
	•				

ŀ

L06139 Phase II Study 76/77 4 Day Sacrifice

CORRELATION OF GROSS AND MICROSCOPIC FINDINGS

Species:	es: Rats	sex. Males	Group Number: 11 - Sacrificed	Dosage Level: •5 mg/1
~ Z .	Animat Number	Client's Fissue Identification	Client's Gross Observations	Microscopic Observations
8	, .	Lungs	Pale red	No corresponding lesion
		tungs - all lobes	Scattered 0.1 x 0.1 cm dark red foci	Hemorrhage
		lleum	reyer's patch, Vio X Vio X Vio Cill, White	No section
		Thymus	Scattered dark red foci	No section
80	0	Lungs	Mottled red	No corresponding lesion
	•			
97				
بر				
. •			-18-	

Ammain         Clearts 1 seare identification         Microscopic         Microsco	Species: Rats	Sex: Males	Group Number: 111 - Sacrificed	Dosage Level: 0 mg/1	
86     Lungs     Mottled red	Animat Number	Client's Tissue Identification	Client's Gross Observations	Microscopic Observations	
	86	Lungs	Mottled red	No corresponding lesion	
					.,
					,
					• *
					·
	•				

Microscopic Observations 1 Dosage Level: •5 mg/1. Hemorrhage No section SCOPIC FINDINGS , · Mottled Proteinaceous calculus present in lumen Client's Gross Observations Group Number: IV - Sacrificed CORRELATION OF GROSS AND A 1 Client's Tissue Identification Sex Males Urinary bladder Lungs 06139 Pha\_e II tudy 76/77 Day Sacrifice pecies: Rats Animal 104

[

Ľ

		Sacrifice
Phuse 11	76/77	Recovery
L06139	Study 7	14 Day

# CORRELATION OF C. JOSS AND MICROSCOPIC FINDINGS

pecies: Rats	Sex Males	Group Number V - Sacrificed	Dosage Level: 0 mg/1
Animat Number	Clurint's Tissue Identification	Client's Gross Observations	Microscopic Observations
90	Lungs - all lobes	Red mottled	Hemorrhage
•	Urinary bladder	Hists calculus	No section
106	Lungs	Mottled red	Hemorrhage
•			
٩.			
,			
-			

÷

÷.,;

	•	fice
	ı	Sacrifice
anna Suint	٠	S
Phase	16/11	Recovery
		Ďay R
L06139	Study	14

CORRELATION OF GHOSS AND MICROSCOPIC FINDINGS

•

Species Rats	Sex: Males	Group Number: VI Sacini ficed	Douatge Level: •5 mg/1
Animal Number	Client's Tissue Identification	Client's Gross Observations	Microscopic Observations
92	Lungs	Scattered dark red foci	Hemorrhage .
	Thymus	Mottled red	No section
	Urinary bladder	White calculus	No ŝection
108	Lungs	Mottled red	Focal lymphocyte aggregate
101			
•			
· · · · · · · · · · · · · · · · · · ·			

		Sacrifice
-		lery
Phase	16/17	Recovery
06139	Study	Day
ē	St	*

CORRELATION OF GROSS AND MICROSCOPIC FINDINGS

Species: Rats

Group Number: VII - Sacrificed

Dosage Level 0 mg/1

Microscopic Observations No corresponding lesion No corresponding lesion No section No section Hemorrhage No section Client's Gross Observations Scattered dark red foci White calculus White calculus Mottled brown Mottled red Mottled red **Client's Tissue Identification** Males. Sex: Urinary bladder **Urinary** bladder Kidneys Thymus Lungs Lungs Animat 94 110

Species: Rats	Sex: Males	Group Number: VIII - Sacrificed	Dosage Level: .5 mg/1
Animat Number	Clent's Tissue Identification	Cherit's Gross Observations	Microscop. Observations
96	Lungs	Mottled red	Hemorrhage
112	Lungs	Mottled red	Hemorrhage
	Kidneys	Mottled brown	No corresponding lesion
	Urinary bladder	White calculus	No section
		· .	
-			
			andre and Break and and a first of the same and an any state of the same state of the same state of the same s
	•		

LOG139 Phase Il Study 76/77 4 Day Sacrifice

CORRELATION OF GROSS AND MICHOSCOPIC FINDINGS

	ervations	Ľ	c										****	ne o contra na suma o contra contra na suma na sum			
. 0 mg/1	Microscopic Observations	to corresponding lesion	No corresponding lesion														
Dosage Level: 0 mg/1		to corre	No corre:														-
	·								-	-							
iced	Client's Gross Observations		-	· .													
I - Sacrificed	Client's Gro	ed	ed														
C oup Number: I		Mottled red	Mottled red	,													
	cation			•	· ·			,			•				•		
Sex: Females	Client's Tissue Identification								•							•	and the second se
	Client's	Lungs	Lungs									•				•	
Species: Rats	Animal Smber	114 L	130 L					•				 T				-	

Ē

÷

•					,	• • • • • • • • • • • • • • • • • • • •	,		 •			Page	= 33	5 (E	P
	Dosage Level: .5 mg/1	Microscopic Observations	Nephroblastoma												
CORRELATION OF GROSS AND MICROSCOPIC FINDINGS	Group Number: <b>II - Sacrificed</b> Dr	Client s Gross Observations	Tan, firm mass, 1.0 x 1.1 x 0.8 cm												
	Sex Females	Client's Tissue Identification	Kidney - Ii. anterior pole												
LO6139 Phase II Study 76/77 4 Day Sacrifice	Species: Rats	Animat Numbar	116		-			10							

:
.

÷

Advised Advisor Cuencial Tissue deminication Cuents Tissue deminication Advisor Tissue deminications Ad	Species: Rats	Species: Rats Sex: Females Group Numbe	Group Number: 111 - Sacrificed Dos	Dosage Level: 0 mg/1
Lungs     Mottled red       Nottled red     No corresponding lesion	Animat Number			Microscopic Observations
		1		corresponding
	,			
				·
	·			

LO6139 Phas Study 76/77 A Day Sacri	e ii fice	GROS		••••
Species: Rats	ts sex females	Group Mumber: IV - Sacrificed	Dosage Level: •5 mg/1	
Animal Number	Chent's Tissue Identification	Client's Gross Observations	Microscopic Observations	
120	Lungs	Mottled	No corresponding lesion	
136	Uterus - horns	Dilated, contain clear fluid	No section	•••
				• • •
				<b>'</b>
•				
•				
				•
•				••••
				•
				Page
				= 35
				_ ( E
-				2L)
	•			•

• . •

:

L06139 Phase II Study 76/77 14 Day Recovery Sacrifice

CORRELATION OF GROSS AND MICROSCOPIC FINDINGS

.

Ģ

	<u> </u>	1	1			T	<del></del>	·•	<del></del>		-,			<u> </u>	age	36	<u>(EP</u>
												ł					
			1					•									
			•														
suo																	
irvati																	
Obse										ľ							
Microscopic Observations																	
rosci	, i																
Mic	ļ ,																
	No section																
	ect																
	0																
						<u> </u>			+			·				<u> </u>	ļ 
						l							ĺ				
		1					, 										
												1					
Client's Gross Observations											1			ł			
serva							1				.				, '		
çõ																	
Gros							, ,										
ant's	,																
Cle		:															
	red																
	ed																,
	Mottled red		· ·			•											
		, (															
					J			· · ·									
Lo																	-
ilicat						·			•							•	
ldent														i ·			
SWe				,					•			•					
1.5.1																	
Chent's Tissue Identification													•				
. 4	Sn												•	7	۰		
•	Thymus										•						
	+-		-														
ber 19									-								
Animat Number	136			ļ						·. ·							•.
			-		<u> </u>		t n t	, <b>1</b>	· •		-						

Study 76/77	study 76/77			
y Re	14 Day Recovery Sacrifice		CORRELATION OF GROSS AND MICROSCOPIC FINDINGS	DINGS
Species: Hals	15	Sex fenales	Group Number: VI - Sacrificed	Dosage Level: .5 mg/1
Animal Number	5	Cient's Tissue Identification	Client's Gross Observations	Microscopic Observations
	Fungs		Mottled red	Nc corresponding lesion
•	tungs		Mottled red	No corresponding lesion
	Kidneys		Mottled brown	No curresponding lesion
,				
	,			
	,			
	1			

ļ

ļ

ļ

rage 3/ LEFL)

.

,

- '

· · ·,

LOG139 Phase II Study 76/77 14 Day Recovery Sacrifice

CORRELATION OF GROSS AND MICROSCOPIC FINDINGS

いたは豊からたとれた。 日本ののための主義にいたか。

×.

		<b>.</b>		 •				<b>*</b>		<del></del>		+	Pa	age	38	(EPL	<u>)</u>
Dbservations	sion		ston						•								
Microscopic Observations	onding le		onding le														
	No corresponding lesion	No section	No corresponding leston														
	Z	Z	Z				· · ·					   					
rvalions	   .:	b) ue															
ross Obse		11 amount of ingesta		-				,									
Client's Gross Observations		ill amo															
	ed red	Contains smal discolored	Mottled red														
	Mottle	Conta di sc	Mottle														
entification								÷				•					
•																	
Client's T		4								•						с. 	
	Fungs	Stomach	Lungs								۰.			•			
Animat Number								······································			•	•					
₹₹	126	142				1.1	а I			•							

•

Study 76/ 14 Day Re	Study 76/77 14 Day Recovery Sacrifice CORF	CORRELATION OF GROSS AND MICROSCOPIC FINDINGS	GS
Species: Rd	Rats <sub>Sex</sub> Females	Group Number: VIII - Sacrificed	Dosage Level: .5 mg/1
Animal Number	Client's Tissue Identification	Client's Gross Observations	Microscopic Observations
128	Lungs	Mottled red	Focal lymphocyte aggregate
144	Lungs	Pale	No corresponding lesion
•.			
•			
•			
• • •			
	·		

### PERSONNEL SUPPORTED BY THIS PROJECT DURING THE PHASE II STUDIES

Catherine Aranyi, Principal Investigator James D. Fenters, Co-Investigator Jeannie Bradof William O'Shea Kirit Parikh Narayanan Rajendran Viadislava Rac Josephine M. Reed Alan Snelson Carol Thompson Stanley Vana W. O. Iverson, Consultant Veterinary Pathologist, (EPL, Inc.) Robert Gibbons, Consultant Biostatistician

### DISTRIBUTION LIST

No. of Coples 25 Commander ATTN: SGRD-UBG US Army Medical Bloengineering Research and Development Laboratory Fort Detrick, Frederick, MD 21701-5010 Cummander US Army Medical Research and Development Command ATTN: SGRD-RMS Fort Detrick, Frederick, Maryland 21701-5012 12 Defense Technical Information Center (DTIC) ATTN: DTIC-DDAC Came: on Station Alexandria, VA 22304-6145 1 Dean School of Medicine Uniformed Services University of the Health Sciences 4301 Jones Bridge Road Bethesda, MD 20814-4799 Commandant Academy of Health Sciences, U.S. Army ATTN: AHS-CDM Fort Sam Houston, TX, 78234-6100 Commander U.S. Army Medical Bloengineering Research and Development Laboratory ATTN: SGRD-UBD-A/Librarian Fort Detrick, Frederick, MD 21701-5010

END

FILMED

10-85

DTIC