Toxicology Research Laboratory

.



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Title Page

Study Report for Task Order No. UIC-7I

FOUR WEEK ORAL DOSE RANGE-FINDING STUDY OF WR242511 IN DOGS

Sponsor: US Army Medical Materiel Development Activity

Test Article: WR242511

Contract No.: DAMD17-92-C-2001

Study Director

Barry S. Levine, D.Sc., D.A.B.T.

In-Life Phase Completed On

March 9, 1994

Performing Laboratory

TOXICOLOGY RESEARCH LABORATORY (TRL) University of Illinois at Chicago (UIC) Department of Pharmacology 1940 W. Taylor St. Chicago, IL 60612-7353

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FOUR WEEK ORAL DOSE RANGE-FINDING STUDY OF WR242511 IN DOGS

Test Article: WR242511

Sponsor:

US Army Medical Materiel Development Activity Fort Detrick Frederick, MD 21702-5014

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SUMMARY

1.

This study evaluated the toxicity of WR242511 tartrate in dogs following four weeks of daily administration by gelatin capsule. The results are summarized in Table 1. The primary toxic effects of WR242511 tartrate were seen in the RBCs, liver and the lung. Anemia and methemoglobinemia were observed in all the dose levels tested. Compensatory changes to the anemic state included macrocytosis, reticulocytosis and increased nucleated RBCs. Decreases in body weight and food consumption were seen at the high dose level and possibly the mid dose level. Microscopic lesions were seen in the liver and the lung at all dose levels tested. Hepatocellular swelling was supported by decreases in the A/G ratio and increases in haptoglobin levels in mid and/or high dose animals. Pulmonary lesions (alveolar proteinic exudates, macrophage infiltration and acute alveolar inflammation) and mild to moderate thrombocytopenia were seen at all dose levels. Leukocytosis consisting of increased mature neutrophils, possibly secondary to stress, was seen in high animals and possibly in the mid dose female. On the basis of the findings from this study and after consultation with the Sponsor, the following three dose levels have been selected for use in the four week oral toxicity study: 0.1, 0.3 and 1.0 mg base/kg/day.

2. INTRODUCTION

This study was conducted to determine the appropriate dose levels for a four week oral toxicity study of WR242511 tartrate by gelatin capsule. The study was conducted in accordance with the specifications of the Sponsor, as indicated in Task Order UIC-7I. The FDA requires the use of two animal species, one of which is a non-rodent, in preclinical toxicology studies. The dog is a standard and accepted non-rodent species for regulatory toxicology studies, and was specified by the Sponsor. Oral administration is the intended clinical route and was also specified by the Sponsor. All methods and procedures were conducted within the spirit of the Quality Assurance Programs of the Toxicology Research Laboratory, University of Illinois at Chicago and Pathology Associates, Inc. designed to conform with FDA Good Laboratory Practices Regulations. No unforeseen circumstances affected the integrity of the study. Dosing was initiated on February 9, 1994 and the in-life portion was terminated on March 9, 1994.

3. MATERIALS AND METHODS

3.1 Test Article

WR242511 Tartrate (Lot No. DJD-08-235, Batch No. BM05816) a yellow powder, was received on June 16, 1993 from Herner & Co. and was assigned an in-house chemical number (1720614). The chemical name of the test article is 8-[(4-Amino-1-methylbutyl)amino]5-(1-hexyloxy)-6-methoxy-4-methylquinoline DL tartrate and the mole fraction of the base is 0.71. It was stored at -20 to -15°C, ambient humidity and protected from light in an amber bottle.

The test article purity had previously been determined (1/24/94) at the completion of the in-life portion of a study entitled "Thirteen Week Oral Toxicity Study of WR242511 in Rats" (UIC/TRL Study No. 107). At that time, the purity was 99.59% \pm 0.02%.

3.2 Animals

A shipment of male and female Beagle dogs were obtained from Marshall Farms, North Rose, NY on November 18, 1993. The animals were approximately 6 - 7 months old (birth dates between April 16, 1993 and May 15, 1993) upon arrival at the UIC AAALAC-accredited animal facility. Each animal was given a facility-unique animal number upon arrival. This number immediately appeared as a tag on a chain collar, and was additionally tattooed on the inner aspect of the ear on the same day. Animals were singly housed, except as noted, in runs in a temperature ($72 \pm 6^{\circ}$ F) and humidity (50 \pm 20%) controlled room with a 12 hour light/12 hour dark cycle. During the quarantine/pretest period, the animals were occasionally housed two/run within sex. The run size, typically at least 15 square feet, was adequate to house dogs at the upper weight range as described in the *Guide for the Care and Use of Laboratory Animals*, DHHS (NIH) No. 86.23. All runs were cleaned and fresh bedding was replaced daily. The runs were sanitized once every two weeks.

Certified Canine Diet No. 5007 (PMI Feeds Inc., St. Louis, MO), approximately 400 g, was provided daily from arrival until termination. Exactly 400 g were provided when food consumption was measured. The food was removed for an overnight fast ($\approx 16 - 20$ hours) prior to blood collection and scheduled sacrifice. Tap water was provided *ad libitum* from an automatic watering system in which the room distribution lines were flushed daily from arrival until termination. The water was untreated with additional chlorine or HCl. There were no known contaminants in the feed or water which were expected to influence the study. The results of the most current comprehensive chemical analyses of Chicago water are documented in files maintained by Quality Assurance.

Animals were quarantined for approximately two months, as they were selected from a shipment used for several studies. Body weights and physical examinations were done upon the dogs' arrival at the animal facility. Additionally, each dog was lightly sprayed upon arrival with Para Pyrethrin Mist for fleas, lice, and ticks. At least one week prior to dosing initiation, hematology and clinical chemistry tests, and fecal examination for internal parasites were performed. All dogs had been previously vaccinated against canine distemper, infectious canine hepatitis, leptospirosis, parainfluenza, parvo, oral papilloma, and rabies by the animal supplier. For approximately three weeks prior to dosing initiation, the animals were observed daily for signs of illness and all unusual observations were reported to the Study Director, Toxicologist, or Clinical Veterinarian. Animals were examined during quarantine and approved for use by the Clinical Veterinarian prior to being placed on test. Quarantine release was documented on the Clinical Veterinarian Log by the veterinarian prior to study initiation.

3.3 Experimental Design

Three animals of each sex were chosen from the shipment on the basis of quarantine data including body weight, food consumption and clinical pathology. These animals were randomized by sex using a table of random letters into the groups shown in the following table.

Treatment <u>Group</u>	Dose Level (mg base/kg/day)	Number of <u>Males</u>	Number of <u>Females</u>
1	0.5	1	1
2	0.9	1	1
3	1.5	1	1

WR242511 dose levels were selected on the basis of a previously conducted two week oral dose range-finding study in rats (UIC/TRL Study No. 106), following consultation with the Sponsor. The number of animals, 1 sex/dose, is the minimum number of dogs and is the number typically used in preliminary dose range-finding studies.

Following treatment group allocation, the animal's number appeared on a card visible on the front of each run. The run card additionally contained the study number, test article identification, treatment group number, sex and dose level. Run cards were color-coded as a function of treatment group.

The test article was administered once daily by gelatin capsule starting with Day 0 (February 9, 1994) for four weeks. All animals received empty gelatin capsules for the last 3 days during Week -1 to acclimate them to the procedure. The quantity of the test article (mg base/kg/day) was adjusted based on the animal's most recent body weight. The animals were dosed up to and including the day prior to scheduled necropsy on Day 28. The dogs weighed 9.7 - 10.7 kg (males) and 8.4 - 9.1 kg (females) on Day -2, and were approximately 9 - 10 months old at initiation of treatment.

Body weights of all animals were recorded at randomization in Week -1, weekly thereafter, and at termination on Day 28. Clinical signs were recorded once daily, approximately 1 - 2 hours after dosing. The general behavior, posture, locomotion, breathing pattern and coat were observed for all animals. The animals were also observed immediately prior to dosing and in the afternoon for moribundity/mortality. Physical examinations (clinical observations) which included examination of eyes and all orifices were conducted in Week -1, on Day 0 prior to dosing, and once weekly thereafter. Food consumption was measured for all animals over an approximate 24 hour period once weekly commencing with Week -1.

Hematology and clinical chemistry parameters were measured following an overnight fast approximately one week prior to dosing initiation, on Day 14 and on Day 28 at termination. In addition, overnight fasted methemoglobin levels were measured weekly commencing on Day 0, just prior to dosing. On Day 21, in addition to the measurement of methemoglobin levels prior to dosing, non-fasted levels were determined approximately 2 and 8 hours after treatment. The animals were unanesthetized and sufficient blood was collected from the jugular vein to measure the following parameters in random order. Water was available *ad libitum* during all fasting periods. Clinical pathology methodology is contained in Appendix 1.

Hematology

Activated partial thromboplastin time	Mean corpuscular hemoglobin (MCH)
Erythrocyte count	Mean corpuscular hemoglobin
Erythrocyte morphology	concentration (MCHC)
Heinz bodies	Mean corpuscular volume (MCV)
Hematocrit	*Methemoglobin
Hemoglobin	Platelet count
Leukocyte count, total and	Prothrombin time
differential	Reticulocyte count

^aMeasured with a Co-oximeter (Instrumentation Laboratory, Model No. 282). The assay was performed within one-hour of sample collection. The specimens were kept on wet ice prior to analysis.

Clinical Chemistry

Alanine aminotransferase (ALT/SGPT)	Globulin (calculated)
Albumin	Glucose
Albumin/globulin ratio (calculated)	Haptoglobin
Alkaline phosphatase	Lactate dehydrogenase (LDH)
Aspartate aminotransferase (AST/SGOT)	Phosphorus (inorganic)
Calcium	Potassium
Chloride	Sodium
Cholesterol	Total bilirubin
Creatinine	Total protein
Creatine kinase (CK)	Triglycerides
Gamma glutamyl transferase	Urea nitrogen (BUN)

Additionally, a minimum of 2.5 ml of blood was collected from the jugular vein weekly, just prior to dosing, beginning on Day 0 for the separation and isolation of plasma and cellular blood components according to the Sponsor's directives. The plasma and cell fractions resulting from separation by centrifugation were sent to COL Thomas Brewer, WRAIR, as specified by the Sponsor. The results obtained from these samples are not included in the study report.

All animals survived the four week treatment period and were sacrificed and necropsied on Day 28. This was accomplished by sodium pentobarbital anesthesia (i.v.; 20-30 mg/kg) and exsanguination. An extensive necropsy was performed under the direction and supervision of the pathologist. Terminal body weights were collected prior to routine sacrifice.

The necropsy procedure was a thorough and systematic examination and dissection of the animal viscera and carcass to include the external surface, all orifices, the cranial cavity, external surface of the brain, cross section of the spinal cord, the nasal cavity and nasal turbinates, thoracic, abdominal and pelvic cavities and their viscera, and cervical tissues and organs. The following tissues and organs were collected and fixed in 10% neutral buffered formalin (NBF).

*Adrenal glands	Nerve (sciatic)
Aorta	*Ovaries
*Brain (fore-, mid-, and hind-)	Pancreas
Cecum	Pituitary
Colon	Prostate
Diaphragm	Rectum
Duodenum	Rib with marrow
Epididymides	Salivary gland (submandibular)
Esophagus	Skin
Eyes and optic nerve	Spinal cord (thoracic, cervical)
Gall bladder	*Spleen
Gross lesions	Stomach
*Heart	*Testes
Ileum	Thymus
Jejunum	*Thyroid gland with parathyroids
*Kidneys	Tongue
*Liver (with gall bladder drained)	Tonsil
*Lungs/Bronchi	Trachea
Lymph node (submandibular)	Ureter
Lymph node (mesenteric)	Urinary bladder
Mammary gland	Uterus
Muscle (skeletal)	

Those tissues marked with an asterisk (*) were embedded in paraffin, sectioned, stained with hematoxylin and eosin, and examined microscopically in all animals necropsied on Day 28.

In addition to the collection of the aforementioned tissues and organs, five tubes of heparinized blood (≈ 250 ml) and bile samples aspirated by syringe from the gall bladder were collected at necropsy according to the Sponsor's directives. These samples were sent to COL Thomas G. Brewer, WRAIR, as specified by the Sponsor, and the results obtained from these samples are not included in the study report.

3.4 Statistical Analysis

Statistical analyses were not conducted due to the small sample size. The dose levels for all the summary and individual data are expressed on the basis of mg base/kg/day.

Quantitative data were tabulated and presented in the report. In addition to the written report, summary data tables of parameters and variability were transmitted to the Sponsor on magnetic media (computer diskette) in "ASCII" form.

RESULTS

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4.1 Mortality/Clinical Signs

Summary of clinical signs are presented in Table 2. Individual clinical signs and daily incidence of clinical signs are contained in Appendix 2.

No animals died during the study. Treatment-related daily clinical signs of cyanosis (1 - 2 hours post-dosing) were observed in all treatment groups. Signs of cyanosis were first observed beginning on Days 1 and 2 in the two high dose animals and beginning on Days 6 and 7 in the low dose animals. The severity increased from slight (barely perceptible, slight blue tinged color) blue gums and sclera and moderate (easily seen blue color) blue tongue in low dose animals to moderate blue gums and severe (marked, deep blue-purple color) tongue in mid and high dose females. The severity of these cyanotic signs appeared to reach a plateau in the second week of treatment and were thereafter observed for the remainder of the study. A few exceptions existed including an increase in severity of signs of cyanosis in the mid and high dose female in Week 4 and the disappearance of severe blue tongue in these same animals after Day 21. Pale gums and tongue (lacking a pink appearance) were also occasionally observed in all dose levels, but was frequently noted for the high dose male. Diarrhea was seen once in the low dose female and emesis was seen several times in the low dose male.

4.2 Body Weight

Individual body weights and individual weight gains are presented in Tables 3 and 4, respectively.

During the treatment period, both high dose animals lost 0.5 kg. Body weight changes in the low and mid dose animals during this period were marginal, ranging from 0.1 to -0.2 kg.

4.3 Food Consumption

Individual food consumption data are shown in Table 5.

Food consumption appeared to be occasionally decreased in mid and/or high dose animals. Food intake did not appear to be altered in low dose animals.

4.4 Clinical Pathology

Individual animal clinical chemistry data are shown in Tables 6.1 - 6.6. Individual animal hematology data are shown in Tables 7.1 - 7.6. Individual animal methemoglobin data are presented in Tables 8.1 and 8.2.

Apparent decreases in serum albumin levels and/or increases in serum globulin levels resulted in a decreased A/G ratio in mid and high dose animals. These changes suggest that WR242511 possibly produced marginal hepatotoxic changes.

Serum haptoglobin levels were below the detection limit (< 13 mg/dl) in all females on Day -7/-8 and the low dose female on Day 14. On Day 14, increases in serum haptoglobin levels were seen in mid and high dose animals, and the low dose male. On Day 28, all animals had increased serum haptoglobin levels. The occurrence of increased levels of this protein, which is synthesized by hepatocytes, is indicative of an inflammatory response, i.e. an acute phase reaction.

Dose-dependent anemia, as indicated by decreased RBC count, hemoglobin and hematocrit, were seen in mid and high dose animals and in the low dose male. The maximal effect was generally seen after two weeks of treatment (Day 14), and some resolution of the anemia was observed by Day 28 in the two higher dose levels. In contrast, the apparent anemia in the low dose male was not observed until Day 28. Compensatory increases in MCV, reticulocyte counts and/or nucleated RBCs were seen in mid and high dose animals and in the low dose male.

Mild to moderate thrombocytopenia was seen in all animals, except in the low dose female. The greatest decrease in platelet count was seen on Day 14. By Day 28, the thrombocytopenia had started to resolve. Slight increases in WBC counts were observed in high dose animals. This leukocytosis was apparently a result of increased mature neutrophil numbers.

Biologically significant elevations of methemoglobin levels were seen in all dose levels. These levels appeared to peak by Day 14 and slightly decrease thereafter. By comparing methemoglobin levels at 2 and 8 hours post dosing (Day21-2h and Day21-8h, respectively) with those measured prior to dosing on Day 21 (Day21-0h), it appeared that methemoglobin levels were at a steady-state level.

4.5 Pathology

The Pathology Report is contained in Appendix 3. A summary of microscopic lesions is shown in Table 9.

The oral administration of WR242511 was associated with changes in the lung and the liver. The lung lesions (alveolar proteinic exudates, macrophage infiltration and acute alveolar inflammation) were observed in the high dose animals, in the mid dose female, and in the low dose male. Proteinic exudate which consisted of amphorous to fibrillar gray-pink acellular material was observed in the alveolar lumen. Macrophage infiltrates were observed mainly in the interstitial tissues near terminal bronchioles, but also were present in the interstitium and in alveoli. These infiltrating macrophages were large cells with abundant, pale, vacuolated cytoplasm. Acute alveolar inflammation was characterized by neutrophil infiltrations with necrosis.

Swollen hepatocytes, a common morphologic manifestation of degenerative changes, was seen in all animals, except in the low dose male. This change was identified as large cells whose cytoplasm had a ground-glass appearance.

No other microscopic changes were considered to be related to WR242511 treatment.

DISCUSSION

This study evaluated the toxicity of WR242511 tartrate in dogs following four weeks of daily administration by gelatin capsule. The results are summarized in Table 1. No animals died during the study. Generalized cyanosis manifested clinically by blue gums, sclera and tongue was observed in all dose levels and was supported by methemoglobinemia. In addition, pale gums and tongue was observed in most animals. An increase in severity and duration was observed in the higher dose levels especially in the females. Decreases in body weights accompanied by sporadic decreases in food consumption were seen in the high dose animals and possibly in the mid dose female.

Treatment-related anemia, as indicated by decreased RBC, hemoglobin and hematocrit, was apparent in all dose levels, but not in the low dose female. Macrocytosis, reticulocytosis and an increase in nucleated RBCs were seen as compensatory responses to the anemic state in mid and high dose animals. Methemoglobinemia was seen in all dose levels throughout the study. Methemoglobin appeared to be maintained at steady state levels by Day 21.

WR242511-induced hepatocyte swelling was noted in all dose levels tested. This lesion was of minimal severity in the affected animals. The hepatocellular swelling may have been associated with the apparent decreases in the A/G ratio, and increases in serum haptoglobin levels, indicative of an acute phase reaction.

In the lung, WR242511 resulted in alveolar proteinic exudates, macrophage infiltration and acute alveolar inflammation. These lesions of minimal to mild severity were observed in the high dose animals, in the mid dose female, and in the low dose male.

Mild to moderate thrombocytopenia was seen at all dose levels. Leukocytosis consisting of increased mature neutrophils was seen in high dose animals. The neutrophilia was possibly an indirect effect of the stress produced by the anemia and methemoglobinemia.

The purpose of this study was to select dose levels for a four week oral toxicity study of WR242511 in dogs. It is anticipated that frank toxicity would occur at the high dose level, marginal or no toxicity accompanied by potentially therapeutic methemoglobin levels would be seen at the mid dose level, and no toxicity accompanied by minimal elevation in methemoglobin levels would be observed at the low dose level. On this basis and after consultation with the Sponsor, the following three dose levels have been selected for use in the four week oral toxicity study: 0.1, 0.3 and 1.0 mg base/kg/day.

6. PERSONNEL

Study Director	Barry S. Levine, D.Sc., D.A.B.T.
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Tox. Lab Supervisor	Soudabeh Soura, B.S.
Lead Technician	Teresa O'Neill, B.S.
Chemistry Specialist	Thomas Tolhurst, B.S.
Clinical Pathology	Maria Lang, A.H.T., C.V.T.

Report preparation was assisted by Rae-Jean T. Ballentine, B.S.

7. ARCHIVES

The raw data, specimens, test article reserves, and final report are archived at the Toxicology Research Laboratory (TRL), University of Illinois at Chicago (UIC), Department of Pharmacology, 1940 W. Taylor St., Chicago, IL 60612-7353.

Table 1

FOUR WEEK ORAL DOSE RANGE-FINDING STUDY OF WR242511 IN DOGS

Summary of Toxic Responses

Dose (mg base/kg/day)	0.5	0.9	1.5
Dogs/Sex	0	•	0
Deaths	0	0	0
Body Weight Gain	NE	↓ F?	Ļ
Food Consumption	NE	↓ F?	0
Clinical Signs	Blue gums Blue sclera Blue tongue Pale gums (F) Pale tongue (F)	Blue gums Blue sclera Blue tongue Pale gums Pale tongue (M)	Blue gums Blue sclera Blue tongue Pale gums Pale tongue (M)
Clinical Chemistry*	↑ нрт	ALB GLOB A/G HPT	ALB GLOB A/G HPT
Hematology ^b	$\begin{array}{c} \downarrow \text{ RBC (M)} & \uparrow \text{ MCV (M)} \\ \downarrow \text{ HGB (M)} & \uparrow \text{ METHB} \\ \downarrow \text{ HCT (M)} & \downarrow \text{ PLT (M)} \end{array}$	$\downarrow RBC \uparrow RETICS \downarrow HGB \uparrow NRBCs \downarrow HCT \uparrow METHB \uparrow MCV \downarrow PLT$	$\begin{array}{c} \downarrow RBC & \uparrow NRBCs \\ \downarrow HGB & \uparrow METHB \\ \downarrow HCT & \downarrow PLT \\ \uparrow MCV & \uparrow LEUK \\ \uparrow RETICS & \uparrow MNEUT \end{array}$
Histopathology	LUNG - Alveolar proteinic exudate (M) Macrophage infiltration (M) Acute alveolar inflammation (M) LIVER - Hepatocellular swelling (F)	LUNG - Alveolar proteinic exudate (F) Macrophage infiltration (F) Acute alveolar inflammation (F) LIVER - Hepatocellular swelling	LUNG - Alveolar proteinic exudate Macrophage infiltration Acute alveolar inflammation LIVER - Hepatocellular swelling
Conclusions	methemoglobinemia were observed macrocytosis, reticulocytosis and inc seen at the high dose level and possi all dose levels tested. Hepatocellu haptoglobin levels in mid and/or h infiltration and acute alveolar inflar Leukocytosis consisting of increased and possibly in the mid dose femal	in all dose levels tested. Compensato creased nucleated RBCs. Decreases in bly the mid dose level. Microscopic les ular swelling was supported by decrea- igh dose animals. Pulmonary lesions numation) and mild to moderate thromb mature neutrophils, possibly secondary e. On the basis of the findings from the	BCs, liver and the lung. Anemia and ry changes to the anemic state included body weight and food consumption were ions were seen in the liver and the lung at ases in the A/G ratio, and increases in (alveolar proteinic exudates, macrophage ocytopenia were seen in all dose levels. to stress, was seen in high dose animals this study and after consultation with the r week oral toxicity study: 0.1, 0.3 and 1.0

^aALB = albumin, GLOB = globulin, A/G = albumin/globulin ratio, HPT = haptoglobin

^bRBC = red blood cell count, HGB = hemoglobin, HCT = hematocrit, MCV = mean corpuscular volume, RETICS = reticulocyte count, NRBCs = nucleated red blood cells, METHB = methemoglobin, PLT = platelets, LEUK = leukocytes, MNEUT = mature neutrophils

? = Possible or marginal effect

NE = No effect

Table 2

FOUR WEEK ORAL DOSE RANGE-FINDING STUDY OF WR24511 IN DOGS

	SUMMARY OF CI	LINICAL S	IGNS		
STUDY: 133		SEX: MA	LE		
	DOSE:(mg/kg) GROUP:	0.5 1-M	0.9 2-M	1.5 3-M	(mg base/kg/day)
	Scheduled Sacrifice Blue Gums Blue Sclera Blue Tongue Pale Gums Pale Tongue	1 1 1 0 0	1 1 1 1 1	1 1 1 1 1	
	Total Number of Animals	1	1	1	
STUDY: 133	DOSE:(mg/kg) GROUP:	SEX: FEMA 0.5 1-F	0.9 2-F	1.5 3-F	(mg base/kg/day)
	Scheduled Sacrifice Blue Gums Blue Sclera Blue Tongue Pale Gums Pale Tongue	1 1 1 1 1 1	1 1 1 1 1 0	1 1 1 1 1 0	

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Table 3.1

FOUR WEEK ORAL DOSE RANGE-FINDING STUDY OF WR24511 IN DOGS

	SUMMARY	OF	BODY WEIGH	TS (Kilogr	ams)
 STUDY: 133			SEX	: MALE	
PERIOD	DOSE: (mg/kg) GROUP:	0.5 1-M	0.9 2-M	1.5 3-M	(mg base/kg/day)
DAY -7	MEAN S.D. N	10.7 0.00 1	10.6 0.00 1	10.1 0.00 1	
DAY -2	MEAN S.D. N	10.4 0.00 1	10.7 0.00 1	9.7 0.00 1	
DAY 5	MEAN S.D. N	10.3 0.00 1	10.5 0.00 1	9.4 0.00 1	
DAY 12	MEAN S.D. N	10.6 0.00 1	10.8 0.00 1	8.9 0.00 1	
DAY 19	MEAN S.D. N	10.3 0.00 1	10.5 0.00 1	9.0 0.00 1	
DAY 26	MEAN S.D. N	10.5 0.00 1	10.6 0.00 1	9.2 0.00 1	

Table 3.2

FOUR WEEK ORAL DOSE RANGE-FINDING STUDY OF WR24511 IN DOGS

	SUMMARY	OF	BODY WEIGH	TS (Kilog	rams)
STUDY: 133			SEX	: FEMA	LE
PERIOD	DOSE: (mg/kg) GROUP:	0.5 1-F	0.9 2-F	1.5 3-F	(mg base/kg/day)
DAY -7	MEAN S_D_	9.4 0.00	8.5 0.00	9.2	
DAY -2	N MEAN	1 9.1	1 8.4	1 9.0	
	S.D. N	0.00	0.00	0.00	
DAY 5	MEAN S.D.	8.6	8.1	8.6 0.00	
DAY 12	N MEAN	1 8.9	8.1	1 8.5	
	S.D. N	0.00	0.00 1	0.00	
DAY 19	MEAN S.D.	9.2	8.2	8.2	
DAY 26	N MEAN	1 9.0	1 8.2	1 8.5	
	S.D.	0.00	0.00	0.00	

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Table 4.1

FOUR WEEK ORAL DOSE RANGE-FINDING STUDY OF WR24511 IN DOGS

	SUMMARY	OF	WEIGHT GAIN	S (Kilog	rams)
STUDY: 13	3		SEX:	MALE	
PERIOD	DOSE: (mg/kg) GROUP:	0.5 1-M		1.5 3-M	(mg base/kg/day)
_ b					
DAY 5		-0.1	-0.2	-0.3	
		0.00	0.00	0.00	
	N	1	1	1	
DAY 12	MEAN	0.3	0.3	-0.5	
	S.D.	0.00	0.00	0.00	
	N	1	1	1	
DAY 19	MEAN	-0.3	-0.3	0.1	
	S.D.	0.00	0.00	0.00	
	N	1	1	1	
DAY 26	MEAN	0.2	0.1	0.2	
		0.00		0.00	
	N	1	1	1	
TOTAL GAI	N MEAN	0.1	-0.1	-0.5	
		0.00		0.00	
	N	1	1	1	

^aSuccessive periods ^bBaseline is Day -2

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Table 4.2

FOUR WEEK ORAL DOSE RANGE-FINDING STUDY OF WR24511 IN DOGS

	SUMMARY	OF	WEIGHT GAINS	5 (Kilog	(rams)	
STUDY: 133			SEX:	FEMA	LE	
PERIOD	DOSE: (mg/kg) GROUP:	0.5 1-F	0.9 2-F	1.5 3-F	(mg base/k	g/day)
_ b						
DAY 5	MEAN	-0.5	-0.3	-0.4		
	S.D.	0.00	0.00	0.00		
	N	1	1	1		
DAY 12	MEAN	0.3	0.0	-0.1		
	S.D.	0.00	0.00	0.00		
	N	1	1	1		
DAY 19	MEAN	0.3	0.1	-0.3		
	S.D.	0.00	0.00	0.00		
	N	1	1	1		
DAY 26	MEAN	-0.2	0.0	0.3		
	S.D.	0.00	0.00	0.00		
	N	1	1	1		
TOTAL GAIN	MEAN	-0.1	-0.2	-0.5		
	S.D.	0.00	0.00	0.00		
	N	1	1	1		

^aSuccessive periods

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^bBaseline is Day -2

Table 5.1

FOUR WEEK ORAL DOSE RANGE-FINDING STUDY OF WR24511 IN DOGS

		SUMMARY OF	DAILY MEAN	FOOD	CONSUMP	TION (Grams)
	STUDY:	133		SI	EX: MALE	
	PERIOD	DOSE:(mg/kg) GROUP:	0.5 1-M	0.9 2-M	1.5 З-м	(mg base/kg/day)
	DAY -6	INTAKE (g) S.D. N	400 0.0 1	257 0.0 1	400 0.0 1	
I	DAY -2	INTAKE (g) S.D. N	347 0.0 1	253 0.0 1	400 0.0 1	
	DAY 6	INTAKE (g) S.D. N	400 0.0 1	319 0.0 1	287 0.0 1	
	DAY 13	INTAKE (g) S.D. N	349 0.0 1	286 0.0 1	201 0.0 1	
	DAY 20	INTAKE (g) S.D. N	350 0.0 1	313 0.0 1	200 0.0 1	
	DAY 27	INTAKE (g) S.D.	321 0.0	256	400	

Table 5.2

FOUR WEEK ORAL DOSE RANGE-FINDING STUDY OF WR24511 IN DOGS

		SUMMARY OF	DAILY MEAN	FOOD	CONSUMPTI	ON (Grams)
	STUDY:	133		SI	EX: FEMALE	
	PERIOD	DOSE:(mg/kg) GROUP:	0.5 1-F			(mg base/kg/day)
i -	DAY -6	INTAKE (g) S.D. N	400 0.0 1	231 0.0 1	325 0.0 1	
1	DAY -2	INTAKE (g) S.D. N	355 0.0 1	266 0.0 1	261 0.0 1	
]	DAY 6	INTAKE (g) S.D. N	400 0.0 1	336 0.0 1	400 0.0 1	
	DAY 13	INTAKE (g) S.D. N	400 0.0 1	206 0.0 1	63 0.0 1	
	DAY 20	INTAKE (g) S.D. N	400 0.0 1	180 0.0 1	400 0.0 1	
	DAY 27	INTAKE (g) S.D. N	331 0.0 1	346 0.0 1	400 0.0 1	

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FOUR WEEK ORAL DOSE RANGE-FINDING STUDY OF WR242511 IN DOGS

INDIVIDUAL ANIMAL CLINICAL CHEMISTRY DATA

STUDY ID: 133 STUDY NO: 133		GROUP: 1-	F : 0.5 mg base/	kg/day		SEX: FEMALE
	ANIMAL ID TES	UNITS	DAY-7/-8	DAY 14	DAY 28	
	8017 ALT	U/L	37	33	36	
	AST	U/L	37 35	48	42	
	TP	g/dL	6.7	6.8	6.5	
	ALB			3.2	3.4	
	GLO			3.6	3.1	
	A/G	· ·	1.23	0.89		
	TBI		0.17	0.18	0.20	
	ALK		129	146	139	
	GGT	U/L	5	3	3	
	CHO	. mg/dL	214	216	195	
	TRY	mg/dL	26	46	32	
	LDH	U/L	101	125	42	
	CK	U/L	208	183	141	
	BUN	mg/dL	17.8	22.0	19.3	
	CREA	mg/dL	0.87	0.93	0.84	
	NA	mmol/L	147	149	148	
	К	mmol/L	3.94	4.22	4.02	
	CL	mEq/L	131	133	128	
	CA	mg/dL	10.9	10.3	9.8	
	IP			4.2	5.0	
	GLU				115	
	HAPT		X	X	85.2	

(--)-Data Unavailable (-)-No Units for Test X -Below limit of Detection.

FOUR WEEK ORAL DOSE RANGE-FINDING STUDY OF WR242511 IN DOGS

INDIVIDUAL ANIMAL CLINICAL CHEMISTRY DATA

STUDY ID: 133							SEX: MALE
STUDY NO: 133			GROUP: 2-M	1 : 0.9 mg base/	kg/day		
	ANIMAL II	D TEST	UNITS	DAY-7/-8	DAY 14	DAY 28	
	7951	ALT	U/L	46	32	30	
		AST		36	48	39	
		TP	g/dL	6.8	6.9	6.7	
		ALB	g/dL		3.1	3.1	
		GLOB	g/dL	3.1	3.8	3.6	
		A/G	-	1.19	0.82	0.86	
		TBILI	mg/dL	0.17	0.28	0.17	
		ALKP	U/L	122	64	65	
		GGT	U/L	5	4	6	
		CHOL	mg/dL	127	180	160	
		TRY	mg/dL	28	57	40	
		LDH	U/L	67	324	39	
		CK	U/L	118	208	98	
		BUN	mg/dL	13.7	15.0	17.1	
		CREA	mg/dL	0.67	0.74	0.72	
		NA	mmol/L	144	145	145	
		K	mmol/L	4.81	4.74	4.54	
		CL	mEq/L	126	130	QNS	
		CA	mg/dL	11.1	10.4	10.0	
		IP	mg/dL			4.9	
		GLU	mg/dL	111	108	117	
		HAPT	mg/dL	101.5	255.2	274.8	

(-)-No Units for Test

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QNS-Quantity Not Sufficient

FOUR WEEK ORAL DOSE RANGE-FINDING

		S	TUDY OF	WR242511	IN DOGS	21110	
	INDI	VIDUA	L ANIMAL	CLINICAL	CHEMIST	TRY DATA	
STUDY ID: 133 STUDY NO: 133			GROUP: 2-F	: 0.9 mg base/	kg/day		SEX: FEMALE
				DAY-7/-8			
	8000	ALT	U/L	41	34	31	
		AST	U/L	42	49	59	
		TP	g/dL	6.7	7.0	7.1	
		ALB	g/dL	3.4	3.2	2.8	
		GLOB	g/dL	3.3	3.8	4.3	
		A/G	-	1.03	0.84	0.65	
		TBILI	mg/dL	0.27	0.27	0.18	
		ALKP	U/L	74	48	73	
		GGT	U/L		2	5	
		CHOL	mg/dL	218	252	269	
		TRY	mg/dL	52 65	74	78	
		LDH	U/L	65	201	68	
		CK	U/L		133	114	
		BUN	mg/dL	15.2	18.0	15.7	
		CREA	mg/dL	0.71	0.75	0.77	
		NA	mmol/L	145	144	144	
		K	mmol/L	4.33	4.57	4.24	
		CL	mEq/L	127	132	110	
		-	ma (all	10 4	10 /	10 1	

10.6

5.5

112

--X

10.4

5.2

113

97.8

10.1

104

42.0

(--)-Data Unavailable (-)-No Units for Test

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CA

IP

GLU

HAPT

mg/dL

mg/dL

mg/dL

mg/dL

X -Below limit of Detection.

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FOUR WEEK ORAL DOSE RANGE-FINDING STUDY OF WR242511 IN DOGS

INDIVIDUAL ANIMAL CLINICAL CHEMISTRY DATA

STUDY ID: 133							SEX: MAL
STUDY NO: 133			GROUP: 3-M	: 1.5 mg base/	kg/day		
	ANIMAL I	D TEST	UNITS	DAY-7/-8	DAY 14	DAY 28	
	7950	ALT	U/L	36	30	21	
		AST			47	55	
		TP	g/dL	7.3	7.7	7.2	
		ALB			2.9	2.7	
		GLOB	g/dL		4.8	4.5	
		A/G	-	0.87	0.60	0.60	
		TBILI	mg/dL	0.15	0.18	0.12	
		ALKP	U/L	99	121	113	
		GGT	U/L	4	2	3	
		CHOL	mg/dL	216	275	178	
		TRY	mg/dL	31	69	56	
		LDH	U/L		151	82	
		CK	U/L	195	169	216	
		BUN	mg/dL	21.7	13.1	16.7	
		CREA	mg/dL	0.71	0.79	0.73	
		NA		146	146	146	
		к	mmol/L	4.43	4.32	4.08	
		CL	mEq/L	124	129	107	
		CA	mg/dL	11.3	10.7	10.2	
		IP	mg/dL	7.1	5.1	5.6	
		GLU	mg/dL	111	107	122	
		HAPT	mg/dL	33.1	378.0	228.6	

(-)-No Units for Test

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FOUR WEEK ORAL DOSE RANGE-FINDING STUDY OF WR242511 IN DOGS

INDIVIDUAL ANIMAL CLINICAL CHEMISTRY DATA -----

STUDY ID: 133 STUDY NO: 133		GROUP: 3	F : 1.5 mg base/	kg/day		SEX: FEMALE
	ANIMAL ID TH	EST UNITS	DAY-7/-8	DAY 14	DAY 28	••••••
	7999 AI	LT U/L	51	42	35	
	A	ST U/L	50	46	47	
	T	g/dL	7.0	7.4	6.7	
	AI	B g/dL	3.7	3.1	3.0	
	GI	OB g/dL	3.3	4.3	3.7	
	A	/G -	1.12	0.72	0.81	
		BILI mg/dL	0.31	0.19	0.17	
	AL	KP U/L	96	201	145	
	G	GT U/L	4	4	5	
	CI	IOL mg/dL	217	336	210	
	TE	RY mg/dL	58	93	59	
	L	DH U/L	136	173	54	
	CI	K U/L	337	139	153	
	BL	JN mg/dL	23.2	12.0	18.6	
	CF	REA mg/dL	0.87	0.87	0.79	
	NA	a mmol/L	144	146	145	
	κ	mmol/L	4.26	4.59	4.59	
	CL	mEq/L	127	129	120	
	CI		10.5	10.2	10.0	
	IF		4.6	5.7	4.9	
		U mg/dL	107	117	114	
		APT mg/dL	X	566.4	244.4	

(--)-Data Unavailable (-)-No Units for Test

X -Below limit of Detection.

FOUR WEEK ORAL DOSE RANGE-FINDING STUDY OF WR242511 IN DOGS

INDIVIDUAL ANIMAL HEMATOLOGY DATA

STUDY ID: 133 STUDY NO: 133			GROUP: 1-M : 0.5 mg bas	se/ka/dav		SEX: MALE
31001 NO. 155						
ANIMAL I	D TEST	UNITS	Day-7/-8	Day 14	Day 28	
7952	RBC	10^6/cmm	6.90	7.49	5.62	
	HGB	g/dL	17.2	18.9	14.0	
	HCT	%	48.2	53.5	41.3	
	MCV	fL	69.9	71.4	73.5	
	MCH	pg	24.9	25.2	24.9	
	MCHC	g/dl	35.7	35.3	33.9	
	RETICS	%RBCs	0.4	1.6	1.1	
	NRBC	COUNT	0	1	2	
	HB	%	0.0	0.3	0.0	
	PLT	10^3/ccm	288	99	131	
	PT	sec	7.2	7.0	7.1	
	APTT	sec	11.5	10.8	11.1	
	WBC	10^3/cmm	8.4	9.3	9.2	
	M. Neutrop		5.1	6.0	7.5	
	I. Neutrop		0.0	0.3	0.0	
	Lymphocyte		2.7	1.9	1.1	
	Monocytes		0.5	1.0	0.6	
	Eosinophil		0.1	0.1	0.1	
	Basophils		0.0	0.0	0.0	
	Atypical L		0.0	0.0	0.0	
MORPHOL	OGY OBSERVAT	IONS:	Anisocytosis Moderate Polychromasia,Slight	Polychromasia,Slight Anisocytosis,Slight		t

WBC corrected for NRBC = or > 10

FOUR WEEK ORAL DOSE RANGE-FINDING STUDY OF WR242511 IN DOGS

INDIVIDUAL ANIMAL HEMATOLOGY DATA

STUDY ID: 133 STUDY NO: 133			GROUP: 1-F : 0.5 mg bas	se/kg/day	SEX:	SEX: FEMALE
ANIMAL I	D TEST	UNITS	Day-7/-8	Day 14	Day 28	
8017	RBC	10^6/cmm	6.34	6.08	6.19	
		g/dL	15.9	15.2	15.3	
		%	44.0	42.5	43.8	
		fL	69.4	69.9	70.8	
	MCH	pg	25.1	25.0	24.7	
		g/dl	36.1	35.8	34.9	
		%RBCs	0.7	1.2	0.9	
	NRBC	COUNT	0	0	0	
	HB	%	0.2	0.2	0.2	
	PLT	10^3/ccm	352	246	258	
	PT	sec	8.0	7.7	7.9	
	APTT	sec	12.3	11.8	11.9	
	WBC	10^3/cmm	7.6	7.4	7.4	
	M. Neutrop	10^3/cmm	4.3	4.1	3.5	
	I. Neutrop	10^3/cmm	0.0	0.1	0.0	
	Lymphocyte	10^3/cmm	2.5	2.4	3.1	
	Monocytes		0.3	0.7	0.6	
	Eosinophil	10^3/cmm	0.5	0.1	0.2	
	Basophils	10^3/cmm	0.0	0.0	0.0	
	Atypical L	10^3/cmm	0.0	0.0	0.0	
MORPHOL	OGY OBSERVATI	ONS:	Anisocytosis,Slight	Polychromasia,Slight	Anisocytosis,Slight	
				Antonio anto Oltoba		

Polychromasia,Slight Anisocytosis,Slight Polychromasia,Slight Poikilocytes,Slight

WBC corrected for NRBC = or > 10

FOUR WEEK ORAL DOSE RANGE-FINDING STUDY OF WR242511 IN DOGS

INDIVIDUAL ANIMAL HEMATOLOGY DATA

TUDY ID: 133 TUDY NO: 133			GROUP: 2-M : 0.9 mg ba	se/kg/day		SEX: MALE
ANIMAL	ID TEST	UNITS	Day-7/-8	Day 14	Day 28	
7951	RBC	10^6/cmm	7.47	5.95	6.71	
	HGB	g/dL	18.4	15.2	16.6	
	HCT	%	52.2	42.8	48.9	
	MCV	fL	69.9	71.9	72.9	
	MCH	pg	24.6	25.5	24.7	
	MCHC	g/dl	35.2	35.5	33.9	
	RETICS	%RBCs	1.3	0.9	2.5	
	NRBC	COUNT	2	5	0	
	HB	%	0.2	0.2	0.0	
	PLT	10^3/ccm	379	103	146	
	PT	sec	7.1	6.9	7.1	
	APTT	sec	11.5	11.8	11.4	
	WBC	10^3/cmm	9.0	7.2	6.6	
	M. Neutrop	10^3/cmm	5.1	3.0	3.6	
	I. Neutrop	10^3/cmm	0.0	0.1	0.1	
	Lymphocyte	10^3/cmm	3.1	3.1	2.3	
	Monocytes	10^3/cmm	0.5	0.6	0.5	
	Eosinophil	10^3/cmm	0.3	0.3	0.1	
	Basophils	10^3/cmm	0.0	0.0	0.0	
	Atypical L	10^3/cmm	0.0	0.0	0.0	
MORPHO	DLOGY OBSERVAT	IONS:	Anisocytosis,Slight	Polychromasia,Slight Anisocytosis,Slight		

Slight

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WBC corrected for NRBC = or > 10

FOUR WEEK ORAL DOSE RANGE-FINDING STUDY OF WR242511 IN DOGS

INDIVIDUAL ANIMAL HEMATOLOGY DATA

STUDY ID: STUDY NO:				GROUP: 2-F : 0.9 m	a base/ka/day	S	EX: FEMALE
A	NIMAL ID	TEST	UNITS	Day-7/-8	Day 14	Day 28	
8	000	RBC	10^6/cmm	6.66	5.99	6.28	
_		HGB	g/dL	16.5	15.1	15.5	
		HCT	%	45.8	41.6	44.7	
		MCV	fL	68.8	69.4	71.2	
		MCH	pg	24.8	25.2	24.7	
		MCHC	g/dl	36.0	36.3	34.7	
		RETICS	%RBCs	0.6	1.2	2.2	
		NRBC	COUNT	0	4	6	
		HB	%	0.0	0.0	0.0	
		PLT	10^3/ccm	417	66	145	
		PT	sec	7.4	7.2	7.5	
		APTT	sec	13.0	12.6	13.1	
		WBC	10^3/cmm	9.3	10.2	10.3	
		M. Neutrop		5.8	5.8	7.4	
		I. Neutrop	10^3/cmm	0.0	0.2	0.0	
		Lymphocyte	10^3/cmm	3.2	2.7	2.4	
		Monocytes	10^3/cmm	0.3	1.2	0.5	
		Eosinophil	10^3/cmm	0.1	0.3	0.0	
		Basophils		0.0	0.0	0.0	
		Atypical L	10^3/cmm	0.0	0.0	0.0	
	MORPHOLO	GY OBSERVAT	IONS:	Anisocytosis Moderate	Polychromasia,Slight Anisocytosis,Slight		

Polychromasia, Slight

WBC corrected for NRBC = or > 10

FOUR WEEK ORAL DOSE RANGE-FINDING STUDY OF WR242511 IN DOGS

INDIVIDUAL ANIMAL HEMATOLOGY DATA

TUDY ID: 133 TUDY NO: 133		GROUP: 3-M : 1.5 mg base/kg/day					
ANIMAL I	D TEST	UNITS	Day-7/-8	Day 14	Day 28		
7950	RBC	10^6/cmm	6.19	5.11	5.84		
	HGB	g/dL	15.6	13.2	14.8		
	HCT	%	43.7	37.5	43.7		
	MCV	fL	70.6	73.4	74.8		
	MCH	pg	25.2	25.8	25.3		
	MCHC	g/dl	35.7	35.2	33.9		
	RETICS	%RBCs	0.7	1.8	1.3		
	NRBC	COUNT	0	1	2		
	HB	%	0.0	0.3	0.1		
	PLT	10^3/ccm	380	75	143		
	PT	sec	7.6	7.1	7.5		
	APTT	sec	12.1	12.6	9.9		
	WBC	10^3/cmm	10.5	15.5	20.1		
	M. Neutrop	10^3/cmm	5.6	11.3	12.9		
	I. Neutrop	10^3/cmm	0.0	0.0	0.2		
	Lymphocyte		3.9	2.6	4.6		
	Monocytes		0.4	1.4	1_4		
	Eosinophil	10^3/cmm	0.6	0.2	0.2		
	Basophils		0.0	0.0	0.0		
	Atypical L	10^3/cmm	0.0	0.0	0.8		
MORPHOLOGY OBSERVATIONS:		Anisocytosis	Polychromasia,Slight	Anisocytosis,Slight			
			Moderate	Anisocytosis,Slight			
			Polychromasia,Slight				

WBC corrected for NRBC = or > 10

FOUR WEEK ORAL DOSE RANGE-FINDING STUDY OF WR242511 IN DOGS

INDIVIDUAL ANIMAL HEMATOLOGY DATA

STUDY ID: 133 STUDY NO: 133			GROUP: 3-F : 1.5 mg ba	se/kg/day	S	EX: FEMALE
ANIMAL	ID TEST	UNITS	Day-7/-8	Day 14	Day 28	
7999	RBC	10^6/cmm	7.43	5.79	6.35	
	HGB	g/dL	18.3	14.7	15.1	
	HCT	%	50.2	40.6	45.0	
	MCV	fL	67.6	70.1	70.9	
	MCH	pg	24.6	25.4	23.8	
	MCHC	g/dl	36.5	36.2	33.6	
	RETICS	%RBCs	0.5	1.9	1.7	
	NRBC	COUNT	0	3	9	
	HB	%	0.0	0.0	0.0	
	PLT	10 ³ /ccm	295	28	169	
	PT	sec	7.3	6.7	7.1	
	APTT	sec	10.9	11.2	11.1	
	WBC	10^3/cmm	7.9	9.0	12.7	
	M. Neutrop	10^3/cmm	5.2	6.3	8.3	
	I. Neutrop	10^3/cmm	0.0	0.0	0.1	
	Lymphocyte	10^3/cmm	2.2	1.4	2.9	
	Monocytes		0.2	1.3	0.8	
	Eosinophil	10^3/cmm	0.2	0.0	0.6	
	Basophils	10^3/cmm	0.0	0.0	0.0	
	Atypical L	10^3/cmm	0.0	0.0	0.0	
MORPHO	LOGY OBSERVATI	ONS:	Anisocytosis,Slight	Anisocytosis,Slight	Anisocytosis,Slight	

Anisocytosis,Slight Polychromasia,Slight Poikilocytes,Slight Target Cells,Slight

WBC corrected for NRBC = or > 10

FOUR WEEK ORAL DOSE RANGE-FINDING STUDY OF WR242511 IN DOGS										
INDIVIDUAL ANIMAL METHEMOGLOBIN DATA TEST: Methemoglobin										
STUDY ID: STUDY NO: ABBR: MET	133							SEX: MALE UNITS: %		
ANIMAL ID	Day-7/-8	Day O	Day 7	Day 14	Day21-0h	Day21-2h	Day21-8h	Day 28		
GROUP: 1- 7952	M:0.5 mg base/k 1.0	kg/day 0.9	16.1	20.2	16.4	15.7	16.8	12.8		
MEAN SD N	1.0 NA 1	0.9 NA 1	16.1 NA 1	20.2 NA 1	16.4 NA 1	15.7 NA 1	16.8 NA 1	12.8 NA 1		
GROUP: 2-1 7951	M:0.9 mg base/k 1.0	kg/day 0.7	23.9	28.0	24.6	23.8	24.8	20.8		
MEAN SD N	1.0 NA 1	0.7 NA 1	23.9 NA 1	28.0 NA 1	24.6 NA 1	23.8 NA 1	24.8 NA 1	20.8 NA 1		
GROUP: 3-1 7950	M:1.5 mg base/k 0.8	kg/day 0.9	37.1	36.8	33.6	32.4	33.2	29.4		
MEAN SD N	0.8 NA 1	0.9 NA 1	37.1 NA 1	36.8 NA 1	33.6 NA 1	32.4 NA 1	33.2 NA 1	29.4 NA 1		

NA-Not Applicable

FOUR WEEK ORAL DOSE RANGE-FINDING STUDY OF WR242511 IN DOGS										
INDIVIDUAL ANIMAL METHEMOGLOBIN DATA TEST: Methemoglobin										
STUDY ID: STUDY NO: ABBR: METH	133							SEX: FEMALE UNITS: %		
ANIMAL ID	Day-7/-8	Day O	Day 7	Day 14	Day21-Oh	Day21-2h	Day21-8h	Day 28		
GROUP: 1-F 8017	:0.5 mg base/1 0.7	kg/day 0.6	12.8	15.4	13.2	13.1	13.7	11.4		
MEAN SD N	0.7 NA 1	0.6 NA 1	12.8 NA 1	15.4 NA 1	13.2 NA 1	13.1 NA 1	13.7 NA 1	11.4 NA 1		
GROUP: 2-F 8000	:0.9 mg base/1 0.5	kg/day 0.7	22.8	30.3	28.1	25.9	26.1	23.9		
MEAN SD N	0.5 NA 1	0.7 NA 1	22.8 NA 1	30.3 NA 1	28.1 NA 1	25.9 NA 1	26.1 NA 1	23.9 NA 1		
GROUP: 3-F 7999	:1.5 mg base/) 0.8	kg/day 0.6	28.0	37.9	32.8	32.4	32.9	29.4		
MEAN SD N	0.8 NA 1	0.6 NA 1	28.0 NA 1	37.9 NA 1	32.8 NA 1	32.4 NA 1	32.9 NA 1	29.4 NA 1		

Table 8.2

NA-Not Applicable

Contract No.: DAMD17-92-C-2001 Task Order No.: UIC-7I UIC/TRL Study No.: 133

Table 9

FOUR WEEK ORAL DOSE RANGE-FINDING STUDY OF WR242511 IN DOGS

Summary of Microscopic Lesions^a

MICROSCOPIC LESIONS		Dose (mg base/kg/day)					
ORGAN - lesion	Sex	0.5	0.9	1.5			
LUNGS - Proteinic alveolar exudate	М	1	0	1			
	F	0	1	1			
- Acute alveolar inflammation	М	1	0	2			
	F	0	1	2			
- Macrophage infiltrates	М	1	0	2			
	F	0	2	2			
LIVER - Hepatocellular swelling	М	0	1	1			
Swelling	F	1	1	1			

^aLesion severity was scored as follows:

1 = Minimal 3 = Moderate 2 = Mild 4 = Marked

For additional information see Pathology Report in Appendix 3.

Contract No.: DAMD17-92-C-2001 Task Order No.: UIC-71 UIC/TRL Study No.: 133

APPENDIX 1

Clinical Pathology Methodology

Clinical Chemistry Test Directory

STUDY:	133								
NO.	ABBR. UNITS	DESCRIPTION PRECISION CA	LCULATED	OPERAND A	OPERAND B	LOWER MALE	LIMIT FEMALE	UPPER MALE	LIMIT FEMALE
1.	ALT	Alanine Aminotransf							
f	U/L	Integer	NO			20	20	60	60
2.	AST U/L	Aspartate Aminotrar Integer	NO			20	20	60	60
3.	TP g/dL	Total Protein 0.0	NO			6.0	6.0	8.0	8.0
4.	ALB g/dL	Albumin 0.0	NO			2.7	2.7	3.7	3.7
5.	TBILI mg/dL	Total Bilirubin 0.00	NO			0.00	0.00	0.30	0.30
6.	ALKP U/L	Alkaline Phosphatas Integer	e NO			50	50	200	200
7.	GGT U/L	Gamma Glutamyl Trar Integer	sferase NO			0	0	10	10
8.	CHOL mg/dL	Cholesterol Integer	NO			150	150	250	250
9.	TRY mg/dL	Triglycerides Integer	NO			20	20	60	60
10.	LDH V/L	Lactate Dehydrogena Integer	se NO			25	25	200	200

11. CK Creatine Kinase U/L NO 50 50 300 300 Integer Blood Urea Nitrogen 12. BUN 0.0 NO 8.0 8.0 18.0 18.0 mg/dL 13. CREA Creatinine mg/dL 0.00 NO 0.50 0.50 1.00 1.00 14. NA Sodium mmol/L NO 140 150 Integer 140 150 15. K Potassium 5.25 mmol/L 0.00 NO 4.00 4.00 5.25

(REPORT CONTINUED)

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Clinical Chemistry Test Directory

..... STUDY: 133 ----

NO.	ABBR. UNITS	DESCRIPTION PRECISION	CALCULATED	OPERAND A	OPERAND B	LOWER MALE	LIMIT FEMALE	UPPER MALE	LIMIT FEMALE
16.	CL mEq/L	Chloride Integer	NO			110.0	110.0	130.0	130.0
17.	CA mg/dL	Calcium 0.0	NO			9.0	9.0	12.0	12.0
18.	1P mg/dL	Inorganic Phosp 0.0	horus NO			4.0	4.0	8.0	8.0
19.	GLU mg/dL	Glucose Integer	NO			90	90	140	140
20.	HAPT mg/dL	Haptoglobin 0.0	NO			0	0	250	250
21.	GLOB g/dL	Globulin 0.0 Ope	rand A - Operand B	TP	ALB	3.0	3.0	6.0	6.0
22.	A/G	A/G Ratio 0.00 Ope	rand A / Operand B	ALB	GLOB	0.50	0.50	1.50	1.50

(END OF REPORT)

02-JUN-1994

CLINICAL CHEMISTRY

Alanine Aminotransferase (ALT/GPT)

Modified Wroblewski & La Due procedure Ciba-Corning 550 Express Clinical Chemistry System Henry, R.J., Chiamori, N., Golub, O.J. and Berkman, S. Am. J. Clin. Path., <u>34</u>, 381, 1960.

Aspartate Aminotransferase (AST/GOT)

Modified Karmen procedure Ciba-Corning 550 Express Clinical Chemistry System Bergmeyer, H.V., Scheibe, P., and Wahlefeld, A.W. Clin. Chem., <u>24</u>, 58, 1978.

Total Protein

Biuret technique Ciba-Corning 550 Express Clinical Chemistry System Kingsley, G.R. J. Biol. Chem. <u>131</u>, 197, 1939.

Albumin

Bromocresol green method Ciba-Corning 550 Express Clinical Chemistry System Doumas, B.T. and Biggs, H.G. Standard Methods of Clinical Chemistry, 7, 175, 1972.

Total Bilirubin

Modified Walters and Gerard method Ciba-Corning 550 Express Clinical Chemistry System Ertinghausen G., Fabiny-Byrd, D.L., Tiffany, T.O., and Carey, S.J. Clinical Chem., <u>19</u>, 1366, 1973.

Alkaline Phosphatase

Modified Bessey-Lowry procedure Ciba-Corning 550 Express Clinical Chemistry System Neumann, H. and Von Vreedendaal M. Clin. Chem. Acta., <u>17</u>, 183, 1967.

Gamma Glutamyl Transferase (GGT)

JFCC Methods for Gamms Glutamyl Transferase Shaw, L.M., Stromme, J.H., London, J.L., Theodorsen, L. J. Clin. Chem. C;in, Biochem. <u>21</u> (1983) 633-646

Cholesterol

Cholesterol esterase-oxidase method Ciba-Corning 550 Express Clinical Chemistry System Rosechlow, P., et. al Z.F. Klin. Chem. V. Klin. Biochem. <u>12</u>, 226, 1974.

CLINICAL CHEMISTRY (Contd.)

Triglycerides

Tetrazolium salt reduction method Ciba-Corning 550 Express Clinical Chemistry System Klotzsch, S., et. al. Advances Automated Analysis, Vol. 1, Mediad Inc., Tarrytown, N.Y., p. 111, 1973.

Lactate Dehydrogenase

 $L \rightarrow P$ technique Ciba-Corning 550 Express Clinical Chemistry System Wacker, W.E.C., Ulmer, D.D., Valle, B.L., New England J Med. <u>225</u>, 449, 1956

Creatine Kinase (CK)

Modification of Szasz et al. procedure Ciba-Corning 550 Express Clinical Chemistry System Clin. Chem. 22 650-656, 1976.

Urea Nitrogen (BUN)

Modified urease technique Ciba-Corning 550 Express Clinical Chemistry System Talke, H. and Schubert, G.E. Klin. Wchnschr. <u>43</u>, 174, 1965.

Creatinine

Jaffe method Ciba-Corning 550 Express Clinical Chemistry System Larsen. K. Clin. Chem. Acta, <u>41</u>, 209, 1972

<u>Na+, K+</u>

Ion specific electrodes Model 614 ISE Na+/K+ Analyzer (Ciba Corning)

Chloride

Mecuric thiocyanate procedure Ciba-Corning 550 Express Clinical Chemistry System Zall, O.M., Fisher, D. and Garner, M.Q. Anal. Chem, <u>28</u>, 1065, 1956.

Calcium

Modified alizarin procedure Ciba-Corning 550 Express Clinical Chemistry System Frings, C.S., et. al. Clin. Chem., <u>16</u>, 816, 1970.

Phosphorus, Inorganic

Ammonium molybdate method Ciba-Corning 550 Express Clinical Chemistry System Fiske, C.H. and Subbarow, Y. J. Biol. Chem. <u>66</u>, 325, 1925.

CLINICAL CHEMISTRY (Contd.)

Glucose

Hexokinase method Ciba-Corning 550 Express Clinical Chemistry System Bondar, J.L. and Mead, D.C. Clin. Chem. <u>20</u>, 586, 1974.

Haptoglobin

Antigen-antibody method Ciba-Corning 550 Express Clinical Chemistry System Atlantic Antibodies Test Kit

Hematology Test Directory

STUDY: 133

NO.	ABBR. UNITS	DESCRIPTION PRECISION C	ALCULATED	OPERAND A	OPERAND B	LOWE MALE	R LIMIT FEMALE	UPPER MALE	R LIM FE
1.	RBC 10^6/cmm	Erythrocytes 0.00	NO			6.00	6.00	8.00	8.
2.	HGB g/dL	Hemoglobin 0.0	NO			12.0	12.0	19.0	19
3.	НСТ %	Hematocrit 0.0	NO			35.0	35.0	55.0	55
4.	MCV fl	Mean Corpuscular V 0.0	olume NO			57.0	57.0	70.0	70
5.	MCH Pg	Mean Corpuscular H 0.0				20.	20	25	25
6.	MCHC g/dl	Mean Corpus. Hemo. 0.0	Conc. NO			32.0	32.0	38.0	38
7.	RETICS %RBCs	Reticulocytes 0.0	NO			0.0	0.0	1.0	1.0
8.	НВ %	Heinz Bodies 0.0	NO			0.0	0.0	2.0	2.0
9.	HJ %	Howell-Jolly Bodie 0.0	s NO			0.0	0.0	2.0	2.0
10.	PLT 10^3/ccm	Platelets Integer	NO			200	200	500	50
11.	PT sec	Prothrombin Time 0.0	NO			6.0	6.0	9.0	9.0
12.	APTT sec	Act. Partial Throm 0.0	bo. Time NO			7.0	7.0	12.0	12.
13.	FIBR mg/dL	Fibrinogen Integer	NO						
14.	WBC 10^3/cmm	Leukocytes 0.0	NO			7.0	7.0	15.0	15
15.	METH %	Methemoglobin 0.0	NO			0	0	3	3

HEMATOLOGY

Erythrocyte Count Electronic counting procedure Sysmex 180A Hematology Analyzer

Hemoglobin

Cyanomethemoglobin method Sysmex 180A Hematology Analyzer

Hematocrit

Indirect method; calculated value based on volume of red cells and volume of blood

Mean Corpuscular Volume (MCV) Indirect method; calculated value based on hematocrit and red blood cell count

Mean Corpuscular Hemoglobin (MCH) Indirect method; calculated value based on erythrocyte count and hemoglobin

Mean Corpuscular Hemoglobin Concentration (MCHC) Indirect method; calculated value based on hematocrit and hemoglobin

Reticulocyte Count

New methylene blue staining procedure Brecher, G., Am. J. Clin. Path., <u>19</u>, 895, 1949.

Heinz Bodies

Methyl Violet staining technique

Platelet Count

Electronic counting procedure Sysmex 180A Hematology Analyzer

Prothrombin Time (PT)

Electra 700 coagulation machine

Activated Partial Thromboplastin Time (APTT) Electra 700 coagulation machine

Fibrinogen

Electra 700 coagulation machine

Leukocyte Count

Electronic counting procedure Sysmex 180A Hematology Analyzer

Methemoglobin

Measured with a Co-oximeter (Instrumentation Laboratory Model 282)

HEMATOLOGY (Contd.)

Leukocyte Differential Count Neutrophils - Immature (bands) Neutrophils - Mature (segs) Monocytes Basophils Lymphocytes Eosinophils Wright stain procedure Schalm, O.W., Jain, N.C. and Carroll, E.J. Veterinary Hematology, Color Plates Chapter, 3rd Edition, Lee and Febiger, 1975.

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Contract No.: DAMD17-92-C-2001 Task Order No.: UIC-7I UIC/TRL Study No.: 133

APPENDIX 2

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Individual Observations (Clinical Signs)

		INDIVI	DUAL CLINICA	L S	IGNS			
STUDY: 133 DAY 0-DAY 2	3	GROUP: DOSE:	1-M 0.5(mg/kg)		SEX:	MALE		
ANIMAL	# OBSERVATIONS		SEVERITY	LOC	ONSET	DURATION	FREQUENCY	
7952	Blue Gums Blue Sclera Blue Tongue Blue Tongue Normal Scheduled Sacr Vomit Seen In		1 1 2		DAY 7 DAY 9 DAY 8 DAY 7 DAY 0 DAY 28 DAY 3	DAY 25 DAY 27 DAY 26 DAY 11 DAY 15 DAY 28 DAY 25	7 16 10 2 9 1 7	
••••••		GROUP: DOSE:	2-M 0.9(mg/kg)		SEX:	MALE		
ANIMAL	# OBSERVATIONS		SEVERITY	LOC	ONSET	DURATION	FREQUENCY	
7951	Blue Gums Blue Sclera Blue Tongue Blue Tongue Normal Pale Gums Pale Tongue Scheduled Sacr	rifice	1 1 2		DAY 8 DAY 8 DAY 6 DAY 8 DAY 0 DAY 12 DAY 9 DAY 28	DAY 26 DAY 27 DAY 27 DAY 24 DAY 25 DAY 25 DAY 25 DAY 28	6 20 12 3 6 7 7 1	
		GROUP: DOSE:	3-M 1.5(mg/kg)		SEX:	MALE		
ANIMAL	# OBSERVATIONS		SEVERITY	LOC	ONSET	DURATION	FREQUENCY	
7950	Blue Gums Blue Sciera Blue Tongue Blue Tongue Normal Pale Gums Pale Tongue Scheduled Sacr	ifice	1 1 2		DAY 6 DAY 11 DAY 2 DAY 7 DAY 0 DAY 3 DAY 8 DAY 28	DAY 7 DAY 27 DAY 22 DAY 20 DAY 1 DAY 27 DAY 27 DAY 28	2 14 8 2 2 22 16 1	

Severity Codes

Observations Severity No. Description Blue Gums/ 1 Slight (barely perceptible, slight blue tinged color) Blue Tongue/ 2 Moderate (easily seen, blue color) Blue Sclera 3 Severe (marked, deep blue-purple color)

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		INDIVI	DUAL	CLINICA	LS	IGNS			
STUDY: 133 DAY 0-DAY 28		GROUP: DOSE:	1-F 0.5	(mg/kg)		SEX:	FEMALE		
ANIMAL #	OBSERVATIONS			SEVERITY	LOC	ONSET	DURATION	FREQUENCY	
8017	Blue Gums Blue Sclera Blue Tongue Diarrhea Normal Pale Gums Pale Tongue Scheduled Sacr	ifice		1 1 1 2		DAY 24 DAY 6 DAY 8 DAY 10 DAY 0 DAY 15 DAY 13 DAY 28	DAY 26 DAY 27 DAY 27 DAY 10 DAY 5 DAY 23 DAY 23 DAY 28	2 22 10 1 6 4 5 1	
		GROUP: DOSE:	2-F 0.9	(mg/kg)		SEX:	FEMALE		
ANIMAL #	OBSERVATIONS			SEVERITY	LOC	ONSET	DURATION	FREQUENCY	
8000	Blue Gums Blue Gums Blue Sclera Blue Sclera Blue Tongue Blue Tongue Blue Tongue Normal Pale Gums Scheduled Sacr	ifice		1 2 1 2 1 2 3		DAY 6 DAY 27 DAY 3	DAY 22 DAY 27 DAY 26 DAY 27 DAY 23 DAY 27 DAY 20 DAY 20 DAY 5 DAY 26 DAY 28	14 1 20 1 6 16 2 4 6 1	
1		GROUP: DOSE:	3-F 1.5	(mg/kg)		SEX:	FEMALE		
ANIMAL #	OBSERVATIONS			SEVERITY	LOC	ONSET	DURATION	FREQUENCY	
7999 1 1 1	Blue Gums Blue Gums Blue Sclera Blue Tongue Blue Tongue Blue Tongue Normal Pale Gums Scheduled Sacr	ifice		1 2 1 1 2 3		DAY 5 DAY 25 DAY 5 DAY 1 DAY 6 DAY 8 DAY 0 DAY 15 DAY 28	DAY 27 DAY 25 DAY 27 DAY 19 DAY 27 DAY 21 DAY 21 DAY 0 DAY 19 DAY 28	15 1 22 13 11 3 1 4 1	

Severity Codes

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Observations	Severity No.	Description
Blue Gums/	1	Slight (barely perceptible, slight blue tinged color)
Blue Tongue/	2	Moderate (easily seen, blue color)
Blue Sclera	3	Severe (marked, deep blue-purple color)

	SUMMAI	RY OF OBSER	VATION IN	CIDENC	E	
STUDY: 133		P.	SEX: MA	LE		
	PERIOD	DOSE:(mg/kg) GROUP:	0.5 1-M	0.9 2-M	1.5 3-M	
	DAY 0 No. Observed Normal		1 1 100%	1 1 100%	1 1 100%	
	DAY 1 No. Observed Normal		1 1 100%	1 1 100%	1 1 100%	
	DAY 2 No. Observed Normal Blue Tongue		1 1 100%	1 1 100%	1 0	
	SEV 1		0	0	1 100%	
	DAY 3 No. Observed Normal Blue Tongue		1 0	1 1 100%	1 0	
	SEV 1 Vomit Seen In Pale Gums	Run	0 1 100% 0	0 0 0	1 100% 0 1 100%	
	DAY 4 No. Observed Normal Blue Tongue		1 1 100%	1 1 100%	1 0	
	SEV 1 Pale Gums		0 0	0 0	1 100% 1 100%	
	DAY 5 No. Observed Normal Blue Tongue		1 1 100%	1 1 100%	1 0	
	SEV 1		0	0	1 100%	

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Severity Codes

Observations	Severity No.	Description
Blue Gums/	1	Slight (barely preceptible, slight blue tinged color
Blue Tongue/	2	Moderate (easily seen, blue color)
Blue Sclera	3	Severe (marked, deep blue-purple color)

		RY OF OBSER		THE	TDENC		
STUDY: 133			SEX:	MAL	E		
	PERIOD	DOSE:(mg/kg) GROUP:		.5 M	0.9 2-M		1.5 З-М
	DAY 6						
	No. Observed		1		1	1	
	Normal		1 100	0%	0	0	
	Blue Gums						
	SEV		•		•		400%
	1 Blue Tongue		0		0	1	100%
	SEV						
	1		0		1 100%	1	100%
	DAY 7						
	No. Observed Blue Gums		1		1	1	
	SEV						
	1		1 100	7%	0	1	100%
	Blue Tongue						
	SEV						
	1		0		1 100%	0	
	2		1 100	1%	0	1	100%
	DAY 8						
	No. Observed		1		1	1	
	Blue Gums						
	SEV						
	1		0		1 100%	0	
	Blue Sclera SEV						
	1		0		1 100%	0	
	Blue Tongue		•				
	SEV		DE DALLA				
	1		1 100		0	0	
	2 Pale Gums		0		1 100% 0	0	100%

Severity Codes

Observations	Severity No.	Description
Blue Gums/	1	Slight (barely preceptible, slight blue tinged color
Blue Tongue/	2	Moderate (easily seen, blue color)
Blue Sclera	3	Severe (marked, deep blue-purple color)

STUDY:	133			SE	X: M	ALE	5		
							0.9		1.5
		PERIOD	DOSE:(mg/kg) GROUP:		0.5 1-M		2-M		3-M
		DAY 9							
		No. Observe	ed	1		1		1	
		Blue Gums							
		SEV							
		1		1	100%	0		0	
		Blue Sclera	8						
		SEV							
		1			100%		100%	0	
		Pale Gums		0		0			100%
		Pale Tongue	8	0		1	100%	1	100%
		DAY 10							
		No. Observe	ed	1		1		1	
		Blue Gums							
		SEV							
		1		1	100%	0		0	
		Blue Sclera	3						
		SEV							
		1		1	100%	1	100%	0	
		Blue Tongue	2						
		SEV							
		1			100%		100%	0	
		Vomit Seen	In Run		100%	0		0	
		Pale Gums		0		0			100%
		Pale Tongu	2	0		0		1	100%
		DAY 11							
		No. Observe	ed	1		1		1	
		Blue Gums							
		SEV							
		1		1	100%	0		0	
		Blue Sclera	3						
		SEV							
		1		1	100%	1	100%	1	100%
		Blue Tongue	9						
		SEV							
		2			100%	0		0	
		Vomit Seen	In Run	1	100%	0		0	
		Pale Gums		0		0			100%

Severity Codes

Observations	Severity No.	Description
Blue Gums/	1	Slight (barely preceptible, slight blue tinged color
Blue Tongue/	2	Moderate (easily seen, blue color)
Blue Sclera	3	Severe (marked, deep blue-purple color)

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	SUMMA	RY OF OBSER	VATION I	NCIDENC	E				
STUDY: 133		SEX: MALE							
		DOSE:(mg/kg)	0.5	0.9	1.5				
	PERIOD	GROUP:	1-M	2-M	3-M				
	DAY 12								
	No. Observed		1	1	1				
	Normal		1 100%	0	0				
	Blue Sclera								
	SEV								
	1		0	1 100%	1 100%				
	Pale Gums		0	1 100%	1 100%				
	Pale Tongue		0	1 100%	1 100%				
	DAY 13								
	No. Observed		1	1	1				
	Blue Gums								
	SEV								
	1		0	1 100%	0				
	Blue Sclera								
	SEV		1 100%	1 100%	1 100%				
	1 Dive Tensile		1 100%	1 100%	1 100%				
	Blue Tongue SEV								
	1		0	1 100%	0				
	Pale Gums		0	0	1 100%				
	Pale Tongue		0	0	1 100%				
	Fate Tongue		0	0	1 100%				
	DAY 14				2.5				
	No. Observed		1	1	1				
	Normal		1 100%	0	0				
	Blue Gums								
	SEV		0	1 100%	0				
	1		0	1 100%	0				
	Blue Sclera SEV								
	1		0	1 100%	1 100%				
	Blue Tongue		0	1 100%	1 100%				
	SEV								
	1		0	1 100%	0				
	Pale Gums		0	0	1 100%				
	Pale Tongue		0	õ	1 100%				
	rate tongue		0	0	1 100%				

Severity Codes

Observations Severity No. Description Blue Gums/ Blue Tongue/ 1 Slight (barely preceptible, slight blue tinged color 2 Moderate (easily seen, blue color) Blue Sciera 3 Severe (marked, deep blue-purple color)

SUMMARY OF OBSERVATION INCIDENCE							
STUDY: 133	SEX: MALE						
		DOSE:(mg/kg)	0.5	0.9	1.5		
	PERIOD	GROUP:	1-M	2-M	3-M		
	DAY 15						
	No. Observed		1	1	1		
	Normal		1 100%	0	0		
	Blue Sclera SEV						
	1		0	1 100%	0		
	Pale Gums		0	1 100%	1 100%		
	Pale Tongue		0	1 100%	1 100%		
	DAY 16		12		-		
	No. Observed		1	1	1		
	Blue Gums						
	SEV 1		1 100%	0	0		
	Blue Sclera		1 100%	0	0		
	SEV						
	1		1 100%	1 100%	0		
	Blue Tongue						
	SEV						
	1		1 100%	1 100%	1 100%		
	Pale Gums		0	0	1 100%		
	DAY 17						
	No. Observed		1	1	1		
	Blue Sclera						
	SEV						
	1		1 100%	1 100%	1 100%		
	Blue Tongue						
	SEV		0	1 100%	0		
	1 Pale Gums		0	1 100% 0	0 1 100%		
	Pale Gums		0	0	1 100%		

Î

Severity Codes

Observations Severity No. Description Blue Gums/ Slight (barely preceptible, slight blue tinged color 1 Blue Tongue/ 2 Moderate (easily seen, blue color) Blue Sclera 3 Severe (marked, deep blue-purple color)

	SUMMAR								
STUDY: 133		SEX: MALE							
		DOSE:(mg/kg)		0.5		0.9		1.5	
	PERIOD	GROUP:		1-H		2-M		3-м	
	D.W. 40								
	DAY 18		1		1		1		
	No. Observed Blue Sclera						1		
	SEV		4	100%	4	100%	1	100%	
	1		1	100%		100%	1	100%	
	Blue Tongue								
	SEV		4	10.0%	•		0		
	1			100%	0		0		
	Vomit Seen In I	Kun		100%		100%	-		
	Pale Gums		0					100%	
	Pale Tongue		0		1	100%	1	100%	
	D.1.4 40								
	DAY 19		4				1		
	No. Observed		1		1		1		
	Blue Sclera								
	SEV		4	100%		400%	4	100%	
	1			100%		100%		100%	
	Pale Gums		0			100%		100%	
	Pale Tongue		U			100%		100%	
	DAY 20								
	No. Observed		1		1		1		
	Blue Sclera								
	SEV								
	1		1	100%	1	100%	1	100%	
	Blue Tongue								
	SEV								
	2		0			100%		100%	
	Pale Gums		0		0		1	100%	
	DAY DA								
	DAY 21		4		4				
	No. Observed		1		1		1		
	Blue Sclera								
	SEV			100%		100%	4	100%	
	1		1	100%	1	100%	1	100%	
	Blue Tongue								
	SEV			100%		100*		100%	
	1	D		100%		100%		100%	
	Vomit Seen In I	KUN		100%	0		0		
	Pale Gums		0		1	100%	1	100%	

Severity Codes

Observations	Severity No.	Description
Blue Gums/	1	Slight (barely preceptible, slight blue tinged color
Blue Tongue/	2	Moderate (easily seen, blue color)
Blue Sclera	3	Severe (marked, deep blue-purple color)

STUDY: 1	33		SE	X: M	ALE			
	PERIOD	DOSE:(mg/kg) GROUP:		0.5 1-M		0.9 2-M		1.5 3-M
	DAY 22							
	No. Observ	/ed	1		1		1	
	Blue Scler	а						
	SEV							
	1		1	100%	1	100%	1	100%
	Blue Tongu	Je						
	SEV							
	1		1	100%	1	100%	1	100%
	Vomit Seer	n In Run	1	100%	0		0	
	Pale Gums		0		1	100%	1	100%
	DAY 23							
	No. Observ	/ed	1		1		1	
	Blue Gums							
	SEV							
	1		0		1	100%	0	
	Blue Scler	а						
	SEV							
	1		1	100%	1	100%	0	
	Blue Tongu	Je						
	SEV							
	1		1	100%	1	100%	0	
	Pale Gums		0		0		1	100%
	Pale Tongu	Je	0		0		1	100%
	DAY 24							
	No. Observ	ved	1		1		1	
	Blue Gums							
	SEV							
	1		1	100%	1	100%	0	
	Blue Scler	a						
	SEV							
	1		1	100%	1	100%	1	100%
	Blue Tongu	Je						
	SEV							
	1		1	100%	0		0	
	2		0		1	100%	0	
	Pale Gums		0		0		1	100%
	Pale Tongu	Je	0		0			100%

Severity Codes

	Severity Codes	
Observations	Severity No.	Description
Blue Gums/	1	Slight (barely preceptible, slight blue tinged color
Blue Tongue/	2	Moderate (easily seen, blue color)
Blue Sclera	3	Severe (marked, deep blue-purple color)

50	MMARY OF OBSER	RVATION I	NCIDENC	12					
STUDY: 133	SEX: MALE								
PERIOD	DOSE:(mg/kg) GROUP:	0.5 1-M	0.9 2-M	1.5 3-M					
DAY 25									
No. Obse	rved	1	1	1					
Blue Gum									
SEV	-								
1		1 100%	0	0					
Blue Scl	era								
SEV									
1		1 100%	1 100%	1 100%					
Blue Ton	gue								
SEV		1 100%	0	0					
1 Verit Se	en In Run	1 100% 1 100%	0	0					
Pale Gum		0	1 100%	1 100%					
Pale Ton		õ	1 100%	1 100%					
rate for	300		1 1004	1 100%					
DAY 26									
No. Obse	rved	1	1	1					
Blue Gum	S								
SEV									
1		0	1 100%	0					
Blue Scl	era								
SEV		4 400%	4 4000	4 4000					
1		1 100%	1 100%	1 100%					
Blue Ton SEV	jue								
5EV 1		1 100%	1 100%	0					
Pale Gum	8	0	0	1 100%					
Pale Ton		0	Ō	1 100%					
DAY 27									
No. Obse		1	1	1					
Blue Scl	era								
SEV									
1		1 100%	1 100%	1 100%					
Blue Ton	gue								
SEV 1		0	1 100%	0					
Pale Gum		0	0	1 100%					
Pale Ton		0	0	1 100%					
Fate Tur	ave	0	0	100%					

Severity Codes

Observations	Seventy No.	Description
Blue Gums/	1	Slight (barely preceptible, slight blue tinged color
Blue Tongue/	2	Moderate (easily seen, blue color)
Blue Sclera	3	Severe (marked, deep blue-purple color)

1			SUMMAI	RY OF OBSE	RVATIO	N	INCIDENCE		
S	rudy:	133	PERIOD	DOSE:(mg/kg) GROUP:	C	-5 -M	MALE 0.9 2-M	1.5 3-M	
			DAY 28 No. Observed Scheduled Sacr	ifice	1 1 10	0%	1 1 100%	1 1002	· · · · · · · · · · · · · · · · · · ·

Severity	Codes

Observations	Severity No.	Description
Blue Gums/	1	Slight (barely preceptible, slight blue tinged color
Blue Tongue/	2	Moderate (easily seen, blue color)
Blue Sclera	3	Severe (marked, deep blue-purple color)

.

•••••	SUMMA	RY OF OBSER	VATION T	NCTDENC	Е	
STUDY: 133			SEX: FEMA			
		DOSE:(mg/kg)	0.5	0.9	1.5	
	PERIOD	GROUP:	1-F	2-F	3-F	
	DAY 5 No. Observed		1	1	1	
	Normal		1 100%	1 100%	0	
	Blue Gums					
	SEV 1		0	0	1 100%	
	Blue Sclera		°	ů.	1 1004	
	SEV		•	•	4 400%	
	1 Blue Tongue		0	0	1 100%	
	SEV					
	1		0	0	1 100%	
	DAY 6					
	No. Observed		1	1	1	
	Blue Gums SEV					
	1		0	1 100%	1 100%	
	Blue Sclera					
	SEV 1		1 100%	1 100%	1 100%	
	Blue Tongue					
	SEV 2		0	1 100%	1 100%	
			0	1 100%	1 100%	
	DAY 7					
	No. Observed Blue Gums		1	1	1	
	SEV					
	1 Dive Colore		0	1 100%	1 100%	
	Blue Sclera SEV					
	1		1 100%	1 100%	1 100%	
	Blue Tongue					
	SEV 1		0	1 100%	0	
	2		0	0	1 100%	

Severity Codes

Observations	Severity No.	Description
Blue Gums/	1	Slight (barely preceptible, slight blue tinged color
Blue Tongue/	2	Moderate (easily seen, blue color)
Blue Sclera	3	Severe (marked, deep blue-purple color)

	SUMMA	RY OF OBSER	VATI	ON II	NCI	DENC	E	
STUDY: 133			SEX:	FEM	ALE	2		
		DOSE:(mg/kg)		0.5		0.9		1.5
	PERIOD	GROUP:		1-F		2-F		3-F
	DAY 8							
	No. Observed		1		1		1	
	Blue Gums							
	SEV							
	1		0		1	100%	1	100%
	Blue Sclera							
	SEV							
	1		1	100%	1	100%	1	100%
	Blue Tongue							
	SEV			100%	•		~	
	1			100%	0		0	
	2 3		0		0	100%	0	100%
	2		0		0		1	100%
	DAY 9							
	No. Observed		1		1		1	
	Blue Gums							
	SEV							
	1		0		1	100%	1	100%
	Blue Sclera							
	SEV							
	1		1	100%	1	100%	0	
	Blue Tongue							
	SEV					0.012-0-0		
	2		0		1	100%	1	100%
	DAY 10							
	No. Observed		1		1		1	
	Blue Gums							
	SEV							
	1		0		1	100%	1	100%
	Blue Sclera							
	SEV							
	1		1	100%	1	100%	1	100%
	Blue Tongue							
	SEV							
	2		0		1	100%	1	100%
	Diarrhea							
	SEV							
	2			100%	0		0	

Severity Codes

Observations	Severity No.	Description
Blue Gums/	1	Slight (barely preceptible, slight blue tinged color
Blue Tongue/	2	Moderate (easily seen, blue color)
Blue Sclera	3	Severe (marked, deep blue-purple color)

	SUMMA	RY OF OBSER	VATION I	ACIDENC	£	
STUDY: 133			SEX: FEM	ALE		
		DOSE:(mg/kg)	0.5	0.9	1.5	
	PERIOD	GROUP:	1-F	2-F	3-F	
	DAY 11					
	No. Observed		1	1	1	
	Blue Gums					
	SEV		120			
	1		0	1 100%	1 100%	
	Blue Sclera SEV					
	1		1 100%	1 100%	1 100%	
	Blue Tongue					
	SEV					
	1		1 100%	0	0	
	2		0	1 100%	1 100%	
	DAY 12					
	No. Observed		1	1	1	
	Blue Gums					
	SEV 1		0	1 100%	1 100%	
	Blue Sclera		0	1 100%	1 100%	
	SEV					
	1		1 100%	1 100%	1 100%	
	Blue Tongue					
	SEV 1		0	0	1 100%	
	3		0	1 100%	0	
			-			
	DAY 13					
	No. Observed		1	1	1	
	Blue Gums SEV					
	1		0	1 100%	0	
	Blue Sclera					
	SEV					
	1		1 100%	1 100%	1 100%	
	Blue Tongue SEV					
	1		0	0	1 100%	
	2		õ	1 100%	0	
	Pale Tongue		1 100%	0	0	

Severity Codes

7

Observations	Severity No.	Description
Blue Gums/	1	Slight (barely preceptible, slight blue tinged color
Blue Tongue/	2	Moderate (easily seen, blue color)
Blue Sclera	3	Severe (marked, deep blue-purple color)

1

STUDY: 133		SEX: FEMA	NCIDENC ALE		
PERICO	DOSE:(mg/kg) GROUP:	0.5 1-F	0.9 2-F	1.5 3-F	
DAY 14 No. Observe	_	1	1	1	
NO. UDSErver Blue Sclera SEV		1	1	1	
1 Blue Tongue		1 100%	0	1 100%	
SEV 1		0	1 100%	1 100%	
Pale Tongue		1 100%	0	0	
DAY 15 No. Observe		1	1	1	
Blue Sclera SEV 1		1 100%	1 100%	1 100%	
Blue Tongue SEV		1 100%	1 100%	1 100%	
1 Pale Gums		0 1 100%	1 100% 1 100%	1 100%	
Pale Tongue		1 100%	0	0	
DAY 16 No. Observed		1	1	1	
Blue Sclera SEV			4 400%	4 400%	
1 Blue Tongue SEV		1 100%	1 100%	1 100%	
1 2		0	0 1 100%	1 100% 0	
Pale Gums Pale Tongue		1 100% 1 100%	1 100% 0	0	

Severity Codes

Severity No.	Description
1	Slight (barely preceptible, slight blue tinged color
2	Moderate (easily seen, blue color)
3	Severe (marked, deep blue-purple color)
	Severity No. 1 2 3

				SUMMAR	RY OF OBSE	RVATI	ON IN	CI	DENC	3			
ST	UDY:	133				SEX:	FEMA	LF	2			 	-
1			PERIOD		DOSE:(mg/kg) GROUP:		0.5 1-F		0.9 2-F		1.5 3-F		
				Observed		1		1		1			
-			Blue SEV 1	Gums Sclera		0		1	100%	0			
			SEV 1 Blue	Tongue		1	100%	1	100%	1	100%		
			SEV 1 2 Pale	Gums		0 0		010	100%	0	100%		
l l			DAY 18	bserved		1		1		1			
			SEV 1 Blue	Sclera		0		1	100%	0			
			SEV 1 Blue SEV	Tongue		1	100%	1	100%	1	100%		
			1 2 Pale	Gums		1 0 0	100%	0 1 0	100%	0	100%		

Severity Codes

Blue Gums/ I Slight (barely preceptible, slight	t blue tinged color
Blue Tongue/ 2 Moderate (easily seen, blue colo	or)
Blue Sclera 3 Severe (marked, deep blue-purpl	le color)

STUDY: 133			SEX: F	EMALI	Ξ		
	PERIOD	DOSE:(mg/kg) GROUP:	0.		0.9 2-F		1.5 3-F
	DAY 19						
	No. Observed		1	1		1	
	Blue Gums						
	SEV						
	1		0	1	100%	0	
	Blue Sclera		0		1004	0	
	SEV						
	1		1 100	Y 1	100%	1	100%
	Blue Tongue		1 100	A 1	100%		100%
	SEV						
	1		0	0		1	100%
	2		0		100%	0	
	Pale Gums		0	0			100%
	Pate duis		0	0			100%
	DAY 20						
	No. Observed		1	1		1	
	Blue Gums						
	SEV						
	1		0	1	100%	1	100%
	Blue Sclera		-				
	SEV						
	1		1 100	x 1	100%	1	100%
	Blue Tongue						
	SEV						
	1		1 100	x 0		0	
	3		0		100%		100%
	÷						
	DAY 21						
	No. Observed		1	1		1	
	Blue Gums						
	SEV						
	1		0	1	100%	1	100%
	Blue Sclera						
	SEV						
	1		1 100	% 1	100%	1	100%
	Blue Tongue						
	SEV						
	1		1 100	% 0		0	
	2		0	1	100%	0	
	3		0	0		1	100%

Severity Codes

Observations	Severity No.	Description
Blue Gums/	1	Slight (barely preceptible, slight blue tinged color
Blue Tongue/	2	Moderate (easily seen, blue color)
Blue Sclera	3	Severe (marked, deep blue-purple color)

	SUMMARY	OF OBSERVAT	IOI	N INC	IDENCE		
STUDY: 133		SEX	:]	FEMAL	E		
PER		DSE:(mg/kg) ROUP:		0.5 1-F	0.9 2-F		1.5 3-F
В	o. Observed Lue Gums		1	1	I	1	
В	SEV 1 lue Sclera SEV		0	1	100%	1 1	00%
В	1 lue Tongue SEV		1 10	00% 1	100%	1 1	00%
	1 2 ale Gums		1 10 0 1 10	1	100%	0 1 1 0	00%
В	o. Observed Lue Gums		1	1	i	1	
В	SEV 1 lue Sclera SEV		0	C)	1 1	100%
В	1 Lue Tongue SEV		1 10	00% 1	100%	1 1	100%
	1 2 ale Gums		0 0 1 10	00% 1) 100%	0	100%
P	ale Tongue		1 10	00% 0)	0	

Severity Codes

Observations	Severity No.	Description
Blue Gums/	1	Slight (barely preceptible, slight blue tinged color
Blue Tongue/	2	Moderate (easily seen, blue color)
Blue Sclera	3	Severe (marked, deep blue-purple color)

SUMMARY OF OBSERVATION INCIDENCE STUDY: 133 DOSE:(mg/kg) 0.5 0.9 1.5 PERIOD GROUP: 1-F 2-F 3-F	
DOSE:(mg/kg) 0.5 0.9 1.5 PERIOD GROUP: 1-F 2-F 3-F	
DAY 24 No. Observed 1 1 1 Blue Gums SEV	
1 1 100% 0 1 100% Blue Sclera SEV	
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	
SEV 1 100% 0 0 2 0 1100% 1100% Pale Gums 0 1100% 0	
DAY 25 No. Observed 1 1 1 Blue Gums SEV	
2 0 0 1 100% Blue Sclera SEV	
SEV 1 1 100% 1 100% Blue Tongue SEV 1 1 100% 0 <td></td>	
1 1 100% 0 0 2 0 1 100% 1 100% Pale Gums 0 1 100% 0	

Sev	erity	Codes

ObservationsSeverity No.DescriptionBlue Gums/1Slight (barely preceptible, slight blue tinged colorBlue Tongue/2Moderate (easily seen, blue color)Blue Sclera3Severe (marked, deep blue-purple color)

SUMMARY OF OBSERVATION INCIDENCE STUDY: 133 DOSE:(mg/kg) 0.5 0.9 1.5 PERIOD GROUP: 1-F 2-F 3-F	
DOSE:(mg/kg) 0.5 0.9 1.5	
PERIOD GROUP: 1-F 2-F 3-F	
DAY 26	
No. Observed 1 1 1 Blue Gums SEV	
1 1 100% 0 1 100%	
Blue Sclera SEV 1 1 100% 1 100% 1 100%	
Blue Tongue SEV 1 1 100% 0 0 2 0 1 100% 1 100%	
2 0 1 100% 1 100% Pale Gums 0 1 100% 0	
DAY 27 No. Observed 1 1 1	
Blue Gums SEV	
1 0 0 1 100% 2 0 1 100% 0 Blue Sclera	
Blue Sciera SEV 1 1 100% 0 1 100%	
2 0 1 100% 0 Blue Tongue	
SEV 1 100% 0 0	
2 · 0 1 100% 1 100%	
No. Observed 1 1 1 1 Scheduled Sacrifice 1 100% 1 100% 1 100%	

Severity Codes

Observations

Observations	Severity No.	Description
Blue Gums/	1	Slight (barely preceptible, slight blue tinged color
Blue Tongue/	2	Moderate (easily seen, blue color)
Blue Sclera	3	Severe (marked, deep blue-purple color)
Blue Tongue/	2 3	Moderate (easily seen, blue color)

Contract No.: DAMD17-92-C-2001 Task Order No.: UIC-7I UIC/TRL Study No.: 133

APPENDIX 3

Pathology Report

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FINAL PATHOLOGY REPORT FOR FOUR WEEK ORAL DOSE RANGE-FINDING STUDY OF WR242511 IN DOGS TRL STUDY NUMBER 133

PREPARED BY PATHOLOGY ASSOCIATES, INC. 10 WEST 35TH STREET CHICAGO, IL 60616

FOR

TOXICOLOGY RESEARCH LABORATORY (M/C 868) DEPARTMENT OF PHARMACOLOGY UNIVERSITY OF ILLINOIS AT CHICAGO COLLEGE OF MEDICINE 1940 WEST TAYLOR STREET CHICAGO, IL 60612-7353

SEPTEMBER 16, 1994

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Final Pathology Report Toxicology Research Laboratory Study Number 133

SECTION I

PATHOLOGY NARRATIVE

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Final Pathology Report Toxicology Research Laboratory Study Number 133

FINAL PATHOLOGY REPORT

FOUR WEEK ORAL DOSE RANGE-FINDING STUDY OF WR242511 IN DOGS

INTRODUCTION

This pathology report, submitted by Pathology Associates, Inc. (PAI) to Toxicology Research Laboratory (TRL), University of Illinois at Chicago, represents the pathology findings for the study designated as "Four Week Oral Dose Range-Finding Study of WR242511 in Dogs", Toxicology Research Laboratory Study Number 133.

EXPERIMENTAL DESIGN AND METHODS

Three groups, each composed of two (one male and one female) Beagle dogs, were given the test article (WR242511) at doses of 0.5, 0.9, and 1.5 mg base/kg/day (mbkd) as outlined in the Summary of Experimental Design (Table I) for four weeks. No control group was included. All animals in the study were sacrificed and necropsied at the termination of the study in accordance with TFL Standard Operating Procedures. Tissues required by the protocol for collection at necropsy were preserved in 10% neutral buffered formalin (Table II). No gross lesions were observed at the necropsy.

Tissues required by the protocol to be examined histologically (Table II) were processed and slides were prepared in accordance with PAI Standard Operating Procedures. These tissues were then evaluated by light microscopy, and the results were tabulated.

The pathology portion of the study was conducted in accordance with the protocol, TRL and PAI Standard Operating Procedures, and in the spirit of Good Laboratory Practices (GLP). However, as this was a non-GLP study, no Quality Assurance Statement was issued.

Microscopic findings for all groups are summarized in the Project Summary Tables (Section II). The mean group severity scores are found in the Severity Summary Tables (Section III). The mean group severity scores were determined by dividing the sum of all severity scores for a finding by the number of tissues examined. Microscopic findings in the protocol-required tissues for individual animals are presented in the Tabulated Animal Data Tables (Section IV). The codes used as entries in these tables are explained in the Report Codes Table.

RESULTS AND DISCUSSION

The Results and Discussion section is divided into two parts: Diagnostic Terms and Histopathology Findings. The Diagnostic Terms portion lists and clarifies diagnostic terminology that may be unclear. Terms listed in the Diagnostic Terms portion of this section were not necessarily considered to be test article-related. The Histopathology Findings portion of this section reports the results and provides discussion of the histopathologic evaluation of the tissues.

Diagnostic Terms

The morphologic characteristics of observations and lesions which require comment are presented in subsequent paragraphs to aid in the interpretation of the data.

Lung

Proteinic exudate in the lung consisted of amorphous to fibrillar gray-pink acellular material in the lumen of alveoli. Macrophage infiltrates were found mainly in the interstitial tissues near terminal bronchioles but occurred in other areas of the interstitium and in alveoli. Infiltrating macrophages were large cells with abundant, pale, vacuolated cytoplasm. Some of the larger interstitial macrophage infiltrates also had acute inflammation which consisted of neutrophil infiltration with necrosis. Neutrophils were also found free in some alveoli. Focal chronic inflammation consisted of a single subpleural focus of granulomatous inflammation. Macrophages within this lesion had abundant dense cytoplasm rather than abundant, pale, vacuolated cytoplasm. Lymphocytes were prevalent in this lesion and there was focal, early fibroplasia. No foreign material was seen, but the lesion was typical of a chronic foreign body response. It was considered not to be related to the other lesions described in the lung.

Liver

Swollen hepatocytes were large cells with a ground-glass appearance to the cytoplasm. These cells compressed and obliterated bile canaliculi. Hemosiderin pigment was a golden-brown granular material within Kupffer cell cytoplasm. Cellular infiltrates were focal aggregates of lymphocytes and plasma cells, usually found adjacent to central veins.

Adrenal Gland

Vacuoles in the cytoplasm of cells in the adrenal cortex were variable in size, lacked a delineating membrane, and were clear.

The remainder of the diagnoses used in this study were considered to be self-explanatory and were not discussed in this section.

Histopathology Findings

As there was only one animal per sex in each treatment group and as there was no control group, it was not possible to identify a dose response for any of the changes observed in this study. Changes interpreted as potentially related to the test article were found in the lung, liver, and possibly in the adrenal gland.

Alveolar proteinic exudate, acute alveolar inflammation, and macrophage infiltration occurred in 4 out of the 6 animals in this study (Table III, Summary of Incidence of Potentially Test Article-Related Changes). They did not occur in the low dose female or in the middle dose male. When present, these changes occurred throughout the sections evaluated. However, as sections from only the left apical lobe were evaluated, they may or may not have occurred throughout the entire lung. The absence of these changes from sections evaluated from two animals does not rule out the possibility that they may have occurred in other areas of the lung in these two animals. These changes are consistent with damage to the endothelial or epithelial barriers in the capillary bed of the lung. Similar changes have been associated with oxidative injury following administration of other chemicals. For these reasons and as they occurred in 4 out of 6 animals in this study, alveolar proteinic exudate, acute alveolar inflammation, and macrophage infiltration were considered to be test article-related changes.

Swollen hepatocytes are a common morphologic manifestation of degenerative change in hepatocytes. Swollen hepatocytes occurred in all animals in this study except the low dose male. When present, swollen hepatocytes tended to occur diffusely throughout the sections of liver evaluated. For these reasons, swollen hepatocytes were considered to be a test article-related change. Hemosiderin deposition occurs in normal animals, but can occur as a response to increased erythrocyte turn-over. Hemosiderin deposits occurred in both low dose animals and in one high dose animal in this study. The relationship of the test article to this change is uncertain, though, as hemosiderin deposits were minimal in these dogs and did not occur in one high dose or in either middle dose animal. For these reasons, hemosiderin deposits in the liver were considered to most likely not be a test article-related change. Cellular infiltrates are a common finding in untreated dogs. This change did not occur in either high dose animal and was considered an incidental change.

Vacuolation of cells in the adrenal cortex can be an incidental observation, can be a non-specific finding related to stress, or can be a direct test article-related effect. The occurrence of this change in the middle and high dose females suggests this to be a test article-related finding in this study. This interpretation is uncertain, though, as the vacuolation present in the adrenal cortex was of minimal severity in both affected dogs and could have occurred for several other reasons.

The other lesions observed were considered to be incidental findings and not to warrant further discussion.

CONCLUSIONS

Under the conditions of this study, changes in the lung (proteinic exudates, acute inflammation, and macrophage infiltrates) and liver (swollen hepatocytes) were interpreted as test article-related. Hemosiderin deposits in the liver and vacuolation of cells in the adrenal cortex may have resulted from exposure to the test article or from other causes. Based on the animals affected with these changes and on their severity scores, these two changes were considered to be not related to the test article. A no-effect level was not determined.

Michael Zmlinsa

Michael J. Tomlinson, DVM, Ph.D. Diplomate, ACVP

Date 16 1994

TABLE I

SUMMARY OF EXPERIMENTAL DESIGN

Treatment Group	Dose Level (mg base/kg/day)	Number of Males	Number of Females
1	0.5	1	1
2	0.9	1	1
3	1.5	1	1

TABLE II

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PROTOCOL-REQUIRED TISSUES

 * Adrenal glands Aorta * Brain (fore-, mid-, and hind-) Cecum Colon Diaphragm Duodenum Epididymides Esophagus Eyes and optic nerves Gallbladder Gross lesions * Heart Ileum Jejunum * Kidneys * Liver (with gallbladder drained) * Lungs/bronchi Lymph node (submandibular) Lymph node (mesenteric) Mammary gland Muscle (skeletal) 	Nerve (sciatic) * Ovaries Pancreas Pituitary Prostate Rectum Rib with marrow Salivary gland (submandibular) Skin Spinal cord (thoracic, cervical) * Spleen Stomach * Testes Thymus * Thyroid gland with parathyroids Tongue Tonsil Trachea Ureter Urinary bladder Uterus
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* These tissues from all animals were designated for processing and histopathologic evaluation.

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TABLE III

SUMMARY OF INCIDENCE OF POTENTIALLY TEST ARTICLE-RELATED CHANGES

	DOSE LEVEL mg base/kg/day	0	.5	0	.9	1	.5
	SEX	Male	Female	Male	Female	Male	Female
HISTOPATHO	DLOGIC CHANGE						
	eolus, Proteinic ammation, Acute crophage	+ + +	-	- -	+ + +	+ + +	+ + +
LIVER: Hepatocyte, S	welling	-	+	+	+	+	+

"+" = present in this animal

"-" = not present in this animal

Report Codes Table

A. Codes applying to organs

- N Tissues within normal histological limits
- A Autolysis precluding adequate evaluation
- P Paired organ missing
- U Tissues unsuitable for complete evaluation
- S Tissues not applicable to animal
- * Tissues not required by protocol

B. Codes applying to microscopic diagnoses

- 1 minimal
- 2 mild
- 3 moderate
- 4 marked
-) focal
-] locally extensive
- > multifocal
- P Present

B Neoplasm, benign

M Neoplasm, malignant without metastasis

- C Neoplasm, malignant with metastasis
- X Metastatic site (+)
- No data entered

SECTION II

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PROJECT SUMMARY TABLE

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PROJECT ID. NO: TRL133 WEEKS: 5		FATES: SEX: MA	Terminal Sac LE	rifice	PAGE 1
GROUP:		0.5 mbkd	0.9 mbkd	1.5 mbkd	
 NUMBER OF ANIMALS:		1	1	1	
		# %	# %	# %	
ADRENAL GLAND	# Ex	1	1	1	
Cortex, hemorrhage		1 (100)	0 (0)	0 (0)	
BRAIN (FORE)	# Ex	1	1	1	
BRAIN (MID)	# Ex	1	1	1	
BRAIN (HIND)	# Ex	1	1	1	
HEART	# Ex	1	1	1	
KIDNEY, LEFT Nephrocalcinosis	# Ex	1 (100)	1 1 (100)	1 1 (100)	
		1 (100)	1 (100)	1 (100)	
KIDNEY, RIGHT	# Ex	1	1	1	
Nephrocalcinosis		1 (100)	1 (100)	1 (100)	
LIVER	# Ex	1	1	1	
Hepatocyte, swelling Infiltrate, cellular		0 (0) 0 (0)	1 (100)	1 (100)	
Pigment, hemosiderin		1 (100)	1 (100)	0 (0) 0 (0)	
Fighent, neitositee in		1 (100)	0 (0)	0 (0)	
SPLEEN	# Ex	1	1	1	
Capsule, siderofibrotic p	aque	1 (100)	0 (0)	0 (0)	
THYROID GLAND	∦ Ex	1	1	1	
PARATHYROID GLAND	# Ex	1	1	1	
CANALLINGTO OLAND	WEX		1		

Project Summary Table

Project Summary Table

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SUMMARY: Incidence of NEOPLASTIC and NON-NEOPLASTIC Microscopic Findings

PROJECT ID. NO: TRL133 WEEKS: 5		FATES: Terminal Sacrifice SEX: MALE						PAGE 12
GROUP: NUMBER OF ANIMALS:		0.5	mbkd 1	0.9 m	nbkd 1	1.5	mbkd 1	
TESTIS	# Ex	#	*	#	*	#	*	
LUNG Alveolus, exudate, proteini	# Ex	1	(100)	1 0	(0)	1	(100)	
Alveolus, inflammation, acu Infiltrate, macrophage	Ite		(100)	0	(0) (0)		(100)	

PROJECT ID. NO: TRL133 WEEKS: 5	PAGE 13				
GROUP: NUMBER OF ANIMALS:		0.5 mbkd 1	0.9 mbkd 1	1.5 mbkd 1	
		# %	# %	# %	
ADRENAL GLAND Cortex, vacuolation, cytop	# Ex lasm		1 1 (100)	1 1 (100)	
BRAIN (FORE)	# Ex	1	1	1	
BRAIN (MID)	# Ex	1	1	1	
BRAIN (HIND)	# Ex	1	1	1	
HEART	# Ex	1	1	1	
KIDNEY, LEFT Nephrocalcinosis	# Ex	1 1 (100)	1 1 (100)	1 0 (0)	
KIDNEY, RIGHT Nephrocalcinosis	∦ Ex		1 1 (100)	1 0 (0)	
LIVER Hepatocyte, swelling	# Ex		1 1 (100)	1 1 (100)	
Infiltrate, cellular Pigment, hemosiderin		1 (100) 1 (100) 1 (100)	1 (100) 0 (0)	0 (0) 1 (100)	
SPLEEN	# Ex	1	1	1	
THYROID GLAND	# Ex	1	1	1	
PARATHYROID GLAND	# Ex	1	1	1	

SUMMARY:	Incidence of NEOPL	ASTI	C and	NON-N	EOPLAS	TIC Mi	crosc	opic Findir	ngs	
PROJECT ID. WEEKS: 5	NO: TRL133			ATES: TE		al Sac	rific	e		PAGE 14
GROUP:			0.5 г	nbkd	0.9	mbkd	1.5	mbkd		
NUMBER OF A	NIMALS:			1		1		1		
			*	*	#	*	#	*		
LUNG		# Ex	1		1		1			
Alveolus,	exudate, proteinic	:	0	(0)	1	(100)	1	(100)		
Alveolus,	inflammation, acut	e	0	(0)	1	(100)	1	(100)		
Infiltrat	e, macrophage		0	(0)	1	(100)	1	(100)		

Project Summary Table

MARY: Incidence of NEOPLASTIC and NON-NEOPLASTIC Microscopic Findings

SECTION III

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SEVERITY SUMMARY TABLE

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 Sever						1.5				
PROJECT ID. NO: TRL133 WEEKS: 5		F	ATES: SEX:		al Sac	rifice		PA	GE	16
GROUP: NUMBER OF ANIMALS:	0	.5 m 1		0.9 m 1		1.5 m 1				
			SEV		SEV		SEV			
ADRENAL GLAND	# Ex	1	021	1		1	521			
Cortex, hemorrhage			1.00	0		0				
BRAIN (FORE)	# Ex	1		1		1				
BRAIN (MID)	# Ex	1		1		1				
BRAIN (HIND)	# Ex	1		1		1				
HEART	# Ex	1		1		1				
KIDNEY, LEFT	# Ex	1		1		1				
Nephrocalcinosis		1	1.00	1	1.00	1	1.00			
KIDNEY, RIGHT	# Ex	1		1		1				
Nephrocalcinosis		1	1.00	1	1.00	1	1.00			
LIVER	# Ex	1		1		1				
Hepatocyte, swelling		0			1.00		1.00			
Infiltrate, cellular Pigment, hemosiderin		0	1.00	1	1.00	0				
Fighent, neitositerin			1.00	U		0				
SPLEEN	# Ex	1		1		1				
Capsule, siderofibrotic pla				0		0				
THYROID GLAND	# Ex	1		1		1				
PARATHYROID GLAND				1		1				

Severity Summary Table

	Severi	ty	Su	mmar	ут	able	2				
Í										PAGE	17
	PROJECT ID. NO: TRL133		F	ATES:	Termin	hal Sad	rifice				
	WEEKS: 5			SEX:	MALE						
	GROUP:	0	.5 m	bkd	0.9 m	nbkd	1.5 m	bkd			
	NUMBER OF ANIMALS:		1			1	1				
		-	_						_		
			#	SEV	#	SEV	#	SEV			
	TESTIS	Ex	1		1		1				
			1		1		1				
	Alveolus, exudate, proteinic		1	1.00	0		1	1.00			
	Alveolus, inflammation, acute		1	1.00	0		1	2.00			
	Infiltrate, macrophage		1	1.00	0		1	2.00			
-	Inflammation, chronic, focal		0		0		1	1.00			

* Severity calculated by the number of tissues examined.

PROJECT ID. NO: TRL133 WEEKS: 5		PAGE	18						
GROUP:	c).5 m	ibkd	0.9 п	ibkd	1.5 m	bkd		
NUMBER OF ANIMALS:		1		1		1			
		#	SEV	*	SEV	*	SEV		
ADRENAL GLAND	# Ex	1		1		1			
Contex, vacuolation, cytopl	asm	0		1	1.00	1	1.00		
BRAIN (FORE)	# Ex	1		1		1			
BRAIN (MID)	# Ex	1		1		1			
BRAIN (HIND)	# Ex	1		1		1			
HEART	# Ex	1		1		1			
KIDNEY, LEFT	# Ex	1		1		1			
Nephrocalcinosis		1	1.00	1	1.00	0			
KIDNEY, RIGHT	# Ex	1		1		1			
Nephrocalcinosis		1	1.00	1	1.00	0			
LIVER	# Ex	1		1		1			
Hepatocyte, swelling			1.00	1	1.00		1.00		
Infiltrate, cellular Pigment, hemosiderin			1.00 1.00	1 0	1.00	0 1	1.00		
SPLEEN	# Ex	1		1		1			
THYROID GLAND	# Ex	1		1		1			
PARATHYROID GLAND	# Ex	1		1		1			

Severity Summary Table

Sever	ity	Sui	nmaı	су Т	able			4	
								PAGE	19
PROJECT ID. NO: TRL133		F.	ATES:	Termin	al Sacr	ifice			
WEEKS: 5			SEX:	FEMALE					
GROUP:	0	.5 mi	bkd	0.9 m	bkd	1.5 m	bkd		
NUMBER OF ANIMALS:		1		1		1			
		#	SEV	#	SEV	#	SEV		
LUNG	# Ex	1		1		1			
Alveolus, exudate, protein	ic	0		1	1.00	1	1.00		
Alveolus, inflammation, ac	ute	0		1	1.00	1	2.00		
Infiltrate, macrophage		0		1	2.00	1	2.00		

* Severity calculated by the number of tissues examined.

SECTION IV

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TABULATED ANIMAL DATA

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	Tabulate	l Animal Data	
	PROJECT ID: TRL133 WEEKS: 5	GROUP: 0.5 mbkd SEX: MALE FATES: Terminal Sacrifice	PAGE 2
ANIMAL ID:		952	
ADRENAL GLAND			
Cortex, hemorrhage	à	1	
BRAIN (FORE)		N	
BRAIN (MID)		N	
BRAIN (HIND)		N	
HEART		N	
KIDNEY, LEFT Nephrocalcinosis		1	
KIDNEY, RIGHT			
Nephrocalcinosis		1	
LIVER			
Pigment, hemosider	·in	1	
SPLEEN		а	
Capsule, siderofib	rotic plaque	1	
THYROID GLAND		N	

	Tabula	Tabulated Animal Data		
	PROJECT ID: TRL1 WEEKS: 5	133 GROUP: 0.5 mbkd SEX: MALE FATES: Terminal Sacrifice	PAGE 22	
	ANIMAL ID:	7952		
	PARATHYROID GLAND Cyst	Ρ		
ļ	TESTIS	N		
	LUNG			
	Alveolus, exudate, proteinic			
	Alveolus, inflammation, acute Infiltrate, macrophage	1		

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Tabulated Animal Data			
	DJECT ID: TRL133 GROUP: 0.9 EKS: 5 FATES: Term	mbkd SEX: MALE minal Sacrifice	PAGE 23
ANIMAL ID:	7951		
ADRENAL GLAND	N		
BRAIN (FORE)	N		
BRAIN (MID)	Ν		
BRAIN (HIND)	Ν		
HEART	N		
KIDNEY, LEFT Nephrocalcinosis	1		
KIDNEY, RIGHT Nephrocalcinosis	1		
LIVER Hepatocyte, swelling	1		
Infiltrate, cellular	1		
SPLEEN	Ν		
THYROID GLAND	N		

Tabulated Animal Data

	Tabulated Animal Data			
	PROJECT ID: TRL133 WEEKS: 5	GROUP: 0.5 mbkd SEX: FEMALE FATES: Terminal Sacrifice	PAGE	27
ANIMAL II	D: 4	8017		
ADRENAL GLAND		N .		
BRAIN (FORE)		Ν		
BRAIN (MID)		N		
BRAIN (HIND)		Ν		
HEART		Ν		
KIDNEY, LEFT Nephrocalcinosis		1		
KIDNEY, RIGHT Nephrocalcinosis		1		
LIVER Hepatocyte, swell Infiltrate, cellu Pigment, hemoside	ular	1 1 1		
SPLEEN		Ν		
THYROID GLAND		N		

1	Tabulat	lated Animal Data		
1	PROJECT ID: TRL133 WEEKS: 5	GROUP: 0.5 mbkd SEX: FEMALE FATES: Terminal Sacrifice	PAGE 28	
	ANIMAL ID:	8017		
1	PARATHYROID GLAND	Ν		
1	OVARY	N		
1	LUNG	N		

PROJECT ID: TRL133 WEEKS: 5	GROUP: 0.9 mbkd SEX: FEMALE FATES: Terminal Sacrifice	PAGE 2
ANIMAL ID:	8000	
ADRENAL GLAND		
Cortex, vacuolation, cytoplasm	1	
BRAIN (FORE)	N	
BRAIN (MID)	N	
BRAIN (HIND)	N	
HEART	Ν	
KIDNEY, LEFT		
Nephrocalcinosis	1	
KIDNEY, RIGHT		
Nephrocalcinosis	1	
LIVER		
Hepatocyte, swelling	1	
Infiltrate, cellular	1	
SPLEEN	N	
THYROID GLAND	N	

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PAG PROJECT ID: TRL133 GROUP: 0.9 mbkd SEX: FEMALE WEEKS: 5 FATES: Terminal Sacrifice ANIMAL ID: 8000 PARATHYROID GLAND N OVARY N LUNG Alveolus, exudate, proteinic 1	
PARATHYROID GLAND N OVARY N LUNG	Æ 3
OVARY N LUNG	
LUNG	
Alveolus, exudate, proteinic 1	
Alveolus, inflammation, acute 1 Infiltrate, macrophage 2	

Tabulated Animal Data				
	PROJECT ID: TRL133 WEEKS: 5	GROUP: 1.5 mbkd SEX: FEMALE FATES: Terminal Sacrifice	PAGE	3
ANIMAL I	D:	7999		
ADRENAL GLAND Cortex, vacuolat	ion, cytoplasm	1		
BRAIN (FORE)		N		
BRAIN (MID)		N		
BRAIN (HIND)		N		
HEART		N		
KIDNEY, LEFT		N		
KIDNEY, RIGHT		N		
LIVER Hepatocyte, swel Pigment, hemosic		1 · · · · · · · · · · · · · · · · · · ·		
SPLEEN		N		
THYROID GLAND		Ν		
PARATHYROID GLAND		N		

Tabulated Animal Data

	Tabulat	ed Animal Data	•
	PROJECT ID: TRL133 WEEKS: 5	GROUP: 1.5 mbkd SEX: FEMALE FATES: Terminal Sacrifice	PAGE 32
ANIMAL	ID:	7999	
OVARY		N	
LUNG			
Atveolus, exuda	te, proteinic	1	
Alveolus, infla	mmation, acute	2	
	rophage	2	

APPENDIX 4

Protocol and Protocol Amendments

FOUR WEEK ORAL DOSE RANGE-FINDING STUDY OF WR242511 IN DOGS

1.0 PURPOSE OF THE STUDY:

The purpose of this study is to determine the toxicity of WR242511 tartrate in dogs following four weeks of daily administration by gelatin capsule. Results derived from this study will be used to determine dose levels for the "Four Week Oral Toxicity Study of WR242511 in Dogs". The protocol for this study was approved by the UIC Animal Care Committee (Appendix 1).

2.0 SPONSOR:

	2.1	<u>Name:</u>	U.S. Army Medical Materiel Development Activity
	2.2	Address:	Fort Detrick Frederick, MD 21702-5009
	2.3	Representative:	George J. Schieferstein, Ph.D.
3.0	TESTI	NG FACILITY:	
	3.1	Name:	Toxicology Research Laboratory (TRL)
	3.2	Address:	University of Illinois at Chicago (UIC) Department of Pharmacology 1940 W. Taylor St. Chicago, Illinois 60612-7353
	3.3	Study Director:	Barry S. Levine, D.Sc., D.A.B.T.
4.0	DATE	<u>S:</u>	
	4.1	Proposed Initiation of	Dosing: 02/09/94
	4.2	Proposed End of In-Li	fe Phase: 03/09/94

4.3 <u>Proposed Study Completion Date</u> (Draft Study Report): 05/06/94

STUDY NO:	REVISED P	
	ATE: 3/1	101

5.0 TEST ARTICLE

- 5.1 <u>Name or Code No:</u> WR242511 Tartrate Bottle Number will be indicated in the raw data.
- 5.2 <u>TRL Chemical No:</u> 1720614
- 5.3 <u>Physical Description:</u> Yellow powder
- 5.4 Stability and Handling of Test Article:
 - 5.4.1 <u>Temperature:</u> -20 to -15°C.
 - 5.4.2 <u>Humidity:</u> Ambient conditions at -20 to -15°C.
 - 5.4.3 Light: Protect from light.
 - 5.4.4 Special Requirements: None.
- 5.5 <u>Special Handling Procedures:</u> Standard safety precautions will be followed including gloves, eye protection, mask, and lab coats.
- 5.6 <u>Log of Test Article:</u> The amount, date, identity of person(s) removing aliquots and the purpose for which each aliquot of the test article was removed from the batch will be documented. At termination of the study, all unused test article will be returned to the Sponsor.

6.0 PERSONNEL:

Study Director	Barry S. Levine, D.Sc., D.A.B.T.
Toxicologist	Clyde W. Wheeler, Ph.D.
Pathologist	Michael J. Tomlinson, D.V.M., Ph.D., D.A.C.V.P.
Pathology Support	Ralph M. Bunte, D.V.M., D.A.C.V.P.
Analytical Chemist	Adam Negrusz, Ph.D.
Clinical Veterinarian	Terry Hewett, D.V.M., D.A.V.C.P.
Veterinarian Support	Documented in raw data
Tox. Lab Supervisor	Soudabeh Soura, B.S.
Lead Technician	Documented in raw data
Chemistry Specialist	Thomas Tolhurst, B.S.
Clinical Pathology	Maria Lang, A.H.T., C.V.T.

7.0 TEST SYSTEM:

7.1	Species:	Dog
7.2	Strain:	Beagle
7.3	Number and Sex:	3 Males and 3 Females
7.4	Age of Animals:	Approximately 7 - 9 months old at dosing initiation.
7.5	Weight of Animals:	Approximately 10 - 13 kg (males) and approximately 8 - 11 kg (females) at dosing initiation.
7.6	Source of Animals:	Marshall Farms, North Rose, NY.

- 7.7 <u>Justification for Selection of Test System:</u> The FDA requires the use of two animal species, one being a non-rodent, in preclinical toxicology studies. The dog is a standard and accepted non-rodent species for regulatory toxicology studies, and is specified by the Sponsor.
- 7.8 <u>Procedure for Unique Identification of Test System:</u> Upon arrival each animal will be given a facility unique number. This number will appear as an ear tattoo and will also appear on a cage card visible on the front of each run. The cage card will additionally contain the study number, test article identification, treatment group number and dose level. Cage cards will be color-coded as a function of treatment group. Raw data records and specimens will also be identified by the unique test animal number.
- 7.9 <u>Housing:</u> The animals will be housed in an AAALAC- accredited facility. Animals will be housed singly per run in a temperature $(65 84^{\circ}F)$ and humidity $(50 \pm 20\%)$ controlled room with a 12 hour light/12 hour dark cycle. The run size, at least 15 feet², is adequate to house dogs at the upper weight range as described in the *Guide for the Care and Use of Laboratory Animals*, DHHS (NIH) No. 86.23. All runs will be cleaned and fresh bedding replaced daily. The runs will be sanitized once every two weeks.
- 7.10 <u>Quarantine Procedure:</u> Animals will be quarantined for at least two weeks. During that time, the animals will be observed daily for signs of illness and all unusual observations will be reported to the Study Director, Toxicologist, or Clinical Veterinarian. Body weights and physical examinations will be done upon the dogs' arrival at the animal facility. Additionally, each dog will be lightly sprayed upon arrival with PARA PYRETHRIN MIST for fleas, lice, and ticks. Within one week of arrival, hematology and clinical chemistry tests, and fecal examination for internal parasites will be performed. If parasites are found, the affected animal will be treated with a vermifuge approved by the Sponsor, and at least 10 days and a negative fecal examination will elapse before the animal is used on a study. All dogs will have been vaccinated against canine distemper, infectious canine hepatitis, leptospirosis, parainfluenza, parvo,

oral papilloma, and rabies by the animal supplier. Animals will be examined during quarantine and approved for use by the Clinical Veterinarian prior to being placed on test. Any sickly animal will be eliminated from the animal selection process. If a selected animal appears sickly prior to initiation of treatment, it will be replaced by a healthy animal prior to treatment under the direction of the Study Director or Toxicologist. Quarantine release will be documented on the Clinical Veterinarian Log by the veterinarian prior to study initiation.

- 7.11 <u>Food:</u> Purina Certified Canine Diet No. 5007 (Ralston Purina Company, St. Louis, MO), approximately 400 g, will be provided daily from arrival until termination. Exactly 400g will be provided when food consumption is measured. The food will be removed for an overnight fast ($\approx 16 - 20$ hours) prior to blood collection or scheduled sacrifice.
- 7.12 <u>Water:</u> Tap water from an automatic watering system in which the room distribution lines are flushed daily will be provided *ad libitum* from arrival until termination. The water is untreated with additional chlorine or HCl.
- 7.13 There are no known contaminants in the feed or water which are expected to influence the study. The results of bi-monthly comprehensive chemical analyses of Chicago water are documented in files maintained by Quality Assurance.
- 7.14 It is not known if the animals will experience pain or distress during the study. Analgesic or anesthetic agents will confound the ability to determine the toxic potential of the test article, and therefore will not be used. If an animal is in severe pain or distress, following consultation with the veterinary staff, it will be euthanized in accordance with standard operating procedures.

8.0 EXPERIMENTAL DESIGN:

8.1 Treatment Groups:

Treatment <u>Group</u>	Dose Level (mg base/kg/dav)	Number of <u>Males</u>	Number of <u>Females</u>
1	0.5	1	1
2	0.9	1	1
3	1.5	1	1

WR242511 dose levels will be selected on the basis of a previously conducted two week oral dose range-finding study in rats (UIC/TRL Study No. 106), following consultation with the Sponsor. The number of animals, 1 sex/dose, is the minimum number of dogs and is the number typically used in preliminary dose range-finding studies.

REVISED PAGE STUDY NO: / 33 INITIAL:	B12
DATE: 3/11/94	

If toxicity is not observed after two weeks of treatment, the mid dose may be escalated above the high dose for the remainder of the treatment period.

- 8.2 <u>Frequency and Route of Administration of the Test Article:</u> The test article will be administered once daily by gelatin capsule starting with Day 0 for at least four weeks. All animals will receive empty gelatin capsules for the last 3 days during Week -1 to acclimate them to the procedure. The quantity of the test article (mg base/kg) will be adjusted based on the animals most recent body weight. The animals will be dosed up to and including the day prior to scheduled necropsy on Day 28.
- 8.3 <u>Justification of Route:</u> The oral route is the intended clinical route and is specified by the Sponsor.
- 8.4 <u>Procedure to Control Bias during the Assignment of Animals to Treatment Groups:</u>During the quarantine/pretest period, the animals will be randomized by sex using a table of random letters or numbers. The method will be documented in the raw data.
- 8.5 Test Article Vehicle: Gelatin capsules (size 000; capacity 1.37 ml).
- 8.6 <u>Test Article Dosage Form Preparation and Analyses:</u> Not applicable.
- 8.7 Type and Frequency of Observations, Tests, Analyses and Measurements:
 - 8.7.1 <u>Clinical Signs:</u> All animals will be observed once daily for clinical signs of toxicity approximately 1 2 hours after dosing. Additionally, all animals will be observed for morbidity/mortality in the afternoon and immediately prior to dosing in the morning.
 - 8.7.2 <u>Clinical Observations:</u> All animals will be subjected to a physical examination including examination of eyes and all orifices at randomization (Week -1), on Day 0 (initiation of dosing), weekly thereafter, and at termination on Day 28.
 - 8.7.3 <u>Body Weight:</u> Body weights of all animals will be recorded at randomization in Week -1, weekly thereafter, and at termination on Day 28.
 - 8.7.4 <u>Food Consumption:</u> Food consumption for all animals will be measured over an approximate 24 hour period weekly commencing with Week -1.
 - 8.7.5 <u>Clinical Pathology:</u> Hematology and clinical chemistry parameters will be measured following an overnight fast approximately one week prior to dosing initiation, on Day 14 and on Day 28 at termination. In addition, methemoglobin levels will be measured weekly commencing on Day 0, just prior to dosing. On Day 21, in addition to the measurement of methemoglobin levels prior to dosing, non-fasted levels will be determined approximately 2 and 8 hours after treatment. The animals will be unanesthetized and sufficient blood will be collected from the jugular vein to measure the following parameters in random order. Water will be available *ad libitum* during all fasting periods.

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Hematology

Activated partial thromboplastin time Erythrocyte count Erythrocyte morphology Heinz bodies Hematocrit Hemoglobin Leukocyte count, total and differential

Mean corpuscular hemoglobin (MCH) Mean corpuscular hemoglobin concentration (MCHC) Mean corpuscular volume (MCV) *Methemoglobin Platelet count Prothrombin time Reticulocyte count

^a To be measured with a Co-oximeter (Instrumentation Laboratory, Model No. 282). The assay will be performed within one-hour of sample collection. The specimens will be kept on wet ice prior to analysis.

Clinical Chemistry

- Alanine aminotransferase (ALT/SGPT) Albumin Albumin/globulin ratio (calculated) Alkaline phosphatase Aspartate aminotransferase (AST/SGOT) Calcium Chloride Cholesterol Creatinine Creatine kinase (CK) Gamma glutamyl transferase
- Globulin (calculated) Glucose Haptoglobin Lactate dehydrogenase (LDH) Phosphorus (inorganic) Potassium Sodium Total bilirubin Total protein Triglycerides Urea nitrogen (BUN)
- 8.7.6 <u>Plasma and Blood Cell Isolation:</u> Just prior to dosing, a minimum of 2.5 ml of blood will be collected from the jugular vein weekly commencing on Day 0 for the separation and isolation of plasma and cellular blood components according to the Sponsor's directives. The plasma and cell fractions resulting from separation by centrifugation will be sent to Col. Thomas Brewer, MD as specified by the Sponsor. The results obtained from these samples will not be included in the study report.
- 8.7.7 <u>Pathology:</u> All animals which die on test or are killed if moribund will be necropsied. All remaining animals which survive the four week test period will be sacrificed and necropsied on Day 28. This will be accomplished by sodium pentobarbital anesthesia (i.v.; 20-30 mg/kg) and exsanguination. An extensive necropsy will be performed under the direction and supervision of the pathologist. Terminal body weights will be collected prior to routine sacrifice.

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The necropsy procedure will be a thorough and systematic examination and dissection of the animal viscera and carcass to include the external surface, all orifices, the cranial cavity, external surface of the brain, cross section of the spinal cord, the nasal cavity and nasal turbinates, thoracic, abdominal and pelvic cavities and their viscera, and cervical tissues and organs. The following tissues and organs will be collected and fixed in 10% neutral buffered formalin (NBF).

Those tissues marked with an asterisk (*) in all treatment groups found dead, sacrificed either *in extremis* or at scheduled necropsy on Day 28 will be embedded in paraffin, sectioned, stained with hematoxylin and eosin, and examined microscopically.

In addition to the collection of the aforementioned tissues and organs, five tubes of heparinized blood (≈ 250 ml) will be collected at euthanasia and bile will be aspirated by syringe from the gall bladder at necropsy according to the Sponsor's directives. The samples will be sent to Col. Thomas G. Brewer, MD as specified by the Sponsor, and the results obtained from these samples will not be included in the study report.

8.7.8 <u>Statistical Analyses:</u> Statistical analyses will not be conducted due to the small sample size. Quantitative data will be tabulated and presented in the report. In addition to the written report, summary data tables of parameters and variability will

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be transmitted to the Sponsor on magnetic media (computer diskette) in "ASCII" form. The transcribed data on disk will no longer be considered to be GLP compliant.

9.0 RECORDS TO BE MAINTAINED:

All data generated during the conduct of the study, except those that are generated as direct computer input, shall be recorded directly, promptly, and accurately in ink in bound books with prenumbered pages or on worksheets that shall be bound during or at the conclusion of the nonclinical laboratory study. All appropriate computer and machine output shall be bound during or at the conclusion of the study. All data entries shall be dated on the day of entry and signed or initialed by the person entering the data.

Any changes in entries for whatever reason (e.g., to correct an error or transposition) shall be made so as not to obscure the original entry, shall indicate the reason for such change, and shall be dated and signed or identified at the time of data input. In computer driven collection systems, the operator responsible for direct data input shall be identified at the time of data input. Any changes in computer entries for whatever reason (e.g., to correct an error or transposition) shall be made in such a manner so as not to obscure the original entry, if possible, shall indicate the reason for such change, and shall be dated and the responsible individual shall be identified.

All recorded data shall be reviewed, signed, and dated by a knowledgeable person, other than the person making the entry, to assure adherence to procedures and to verify observations.

Upon completion of the study and submission of the final report, all raw data, documentation, specimens, test article reserves and other materials necessary to reconstruct the study will be stored in the TRL archives maintained by Quality Assurance.

All changes or revisions, and reasons therefore, to this protocol once it is approved shall be documented, signed by the Study Director and Sponsor, dated and maintained with the protocol.

10.0 REGULATORY REQUIREMENTS:

This study will be performed within the spirit of the UIC/TRL Quality Assurance Program designed to conform with FDA Good Laboratory Practice Regulations and EPA Good Laboratory Practice Standards.

Will this study be submitted to a regulatory agency? Yes If so, to which agency(ies)? Food and Drug Administration

Does the Sponsor Request that test article samples be returned? <u>Possibly: direction to be provided</u> by Sponsor.

Does the Sponsor request that samples of the test article/carrier mixture(s) be returned to the Sponsor? Not applicable

PRTL133

Office of the Vice Chancellor for Research (M/C 672) 310 Administrative Office Building 1737 West Polk Street Chicago, Illinois 60612-7227 (312) 996-4995

Appendix 1

November 22,1993

IIC The University of Illinois at Chicago

Barry S. Levine Med-Pharmacology 312 BGRC, M/C 868

Dear Dr. Levine:

The protocol indicated below has been reviewed in accordance with the Animal Care Policies of the University of Illinois at Chicago and approved on May 18, 1993.

Title of Application:

Four Week Oral Dose Range-Finding Study of WR242511 In Dogs

ACC Number: 93-033-12

This institution has Animal Welfare Assurance Number A3460.01 on file with the Office for Protection from Research Risks, NIH. Please transmit this letter of acceptable verification of your research protocol to your sponsor.

Thank you for complying with the Animal Care Policies and Procedures of UIC.

Sincerely yours,

uephine E. Thiller

Josephine B. Miller, Ph.D. Chair, Animal Care Committee

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11.0 PROTOCOL APPROVAL:

STUDY DIRECTOR:

11/19/93

Barry S. Levine, D.Sc., D.A.B.T.

Date

SPONSOR APPROVAL:

George J. Schieferstein, Ph.D. Contracting Officer's Representative (COR)

12/13/93. Date

COMMENTS FROM THE COR:

the test article; "BM05816".

Range-Finding Study of WR242511 in Dogs

specific bottle number be included in the protocol.

Study No .:	133
Title:	Four Week Oral Dose
1. Page 2	Section 5.1
Indica	te the Bottle Number of
Reason:	Sponsor requested that
2. Page 4	Section 7

Add the following section:

"7.14 It is not known if the animals will experience pain or distress during the study. Analgesic or anesthetic agents will confound the ability to determine the toxic potential of the test article, and therefore will not be used. If an animal is in severe pain or distress, following consultation with the veterinary staff, it will be euthanized in accordance with standard operating procedures."

Reason: Sponsor requested addition to the protocol.

3. Page 4 Section 8.1

Add the following sentence to the end of the first paragraph "The number of animals, 1 sex/dose, is the minimum number of dogs and is the number typically used in preliminary dose range-finding studies."

Reason: Sponsor requested addition to the protocol.

4. Page 5 Section 8.5

Change the test article vehicle section to the following "Gelatin capsules (size 000; capacity 1.37 ml)."

Reason: Clarification of the size and capacity of the gelatin capsules to be used.

5. Page 6 Section 8.7.6

Change first sentence to indicate that all blood collection will be done "just prior to dosing" and that the plasma and cellular components will be separated according to the Sponsor's directives.

Study No.: 133

Title: Four Week Oral Dose Range-Finding Study of WR242511 in Dogs

5. contd.

Reason: Clarification of the time when blood collection will be performed and that the separation will be performed according to the SOP provided by the Sponsor.

6. Page 6 Section 8.7.7

Add "(i.v.; 20-30 mg/kg)" after "sodium pentobarbital anesthesia".

Reason: Clarification of the protocol to indicate the dose and route of phenobarbital.

- 7. Page 7 Section 8.7.7
 - A) Change scheduled necropsy date from "Day 14" to "Day 28" in the second paragraph.
 - B) Change third paragraph to indicate that " ≈ 250 ml" of heparinized blood will be collected at euthanasia and bile at necropsy according to the Sponsor's directives in an SOP to be provided by the Sponsor.

Reason: Mistake in protocol (A) and Sponsor requested change in the protocol (B).

Approvals:

Barry S. Levine, D.Sc. D.A.B.T.

. .

SPONSOR APPROVAL:

STUDY DIRECTOR:

George J. Schieferstein, Ph.D. Contracting Officer's Representative (COR)

Date

Study No.: 133

Title: Four Week Oral Dose Range-Finding Study of WR242511 in Dogs

8. Page 1 Section 4.0

Add the study dates as follows:

4.1	Proposed Initiation of Dosing:	02/09/94

4.2 Proposed End of In-Life Phase: 03/09/94

4.3 <u>Proposed Study Completion Date</u> (Draft Study Report): 05/06/94

Reason: The study dates have been finalized.

9. Page 4 Section 8.1

Change the dose levels to read as follows:

"Low" = "0.5" mg base/kg/day

"Mid" = "0.9" mg base/kg/day

"High" = "1.5" mg base/kg/day

Reason: Dose levels have been selected following consultation with the Sponsor.

10. Page 5 Section 8.7.5

Change the first sentence to indicate clinical pathology parameters will be measured "approximately one week prior to dosing initiation" in place of "within one week of arrival".

Reason: Clarification of the protocol.

Approvals:

STUDY DIRECTOR:

Barry S. Levine, D.Sc. D.A.B.T.

SPONSOR APPROVAL:

George J. Schieferstein, Ph.D. Contracting Officer's Representative (COR)

3/18/74

Date

133 Study No .:

Four Week Oral Dose Range-Finding Study of WR242511 in Dogs Title:

Section 8.7.5 11. Page 5

> Add the following sentence "On Day 21, in addition to the measurement of methemoglobin levels prior to dosing, non-fasted levels will be determined approximately 2 and 8 hours after treatment."

Sponsor requested addition to the study. Reason:

Section 8.7.7 12. Page 7

> Add "Lungs/Bronchi" to the list of tissues which will be embedded in paraffin, sectioned, stained with hematoxylin and eosin, and examined microscopically.

The lungs were a potential target organ in a previously conducted rat toxicity study. Reason:

Approvals:

STUDY DIRECTOR:

Barry S. Levine, D.Sc. D.A.B.T.

George J. Schieferstein, Ph.D.

Contracting Officer's Representative (COR)

alala X

SPONSOR APPROVAL:

APPENDIX 5

Study Deviations

FOUR WEEK ORAL DOSE RANGE-FINDING STUDY OF WR242511 IN DOGS

Study Deviation

Deviation Type

Specific Deviation

Effect on Study

Protocol

On several occasions, the relative humidity in the animal room deviated outside the specified range by \leq -7%.

None. These sporadic occurrences were not considered to have had an impact on the outcome of the study.

"The detailed "Deviation Report" is contained in the raw data which is archived at the University of Illinois at Chicago, Department of Pharmacology, Chicago, Illinois.

The above deviation did not affect the integrity of the study.

Barry S. Levine, D.Sc., D.A.B.T.

a

a Date