

UNITED STATES ENVIRONMENTAL PROTECTION AGENON WASHINGTON DIG 20455

007155

4/25/1989

YENCRANDUM

CFFICE DF PESTICIDES AND TOXIC SUBSTANCES

SUBJECT: Review of the Mouse Oncogenicity Study on Endosulfan

IO: George LaPocca, PM-15 Registration Division, H75050

EFOM: Murcia van Gemert, Fh.D. /// Lune Commut. [25/89 Acting Chief, HEAS Branch, H75090

THRU: William Burnam. Actino Director, HED, H75090

Chemical: Endoscifar

Caswell No: 420

Frodest No: 8-1094

Fedistration Division had requested that EFAS review the mouse oncocenicity and rangefinding studies on the compound Endosulfan.

Conclusions:

Under the conditions of the study, endosulfan was not oncogenic when fed to male and female Hoe:NMPRE rice for 24 months at levels of 2, 6, or 18 ppm. There were no overt signs of toxicity or dose-related effects on clinical observations, food consumption, hematology, clinical chemistry, urinalysis, organ weights, macroscopic pathology, or microscopic pathology. Decreased survival in high-dose females and body weight reduction in highdose males throughout the study were considered to be compoundrelated effects. Histologically, the incidence of lymphosarcoma was high in dosed and control males and females. Based on the effects of Endosulfan on mortality at 18 ppm, the LOEL is 19 ppm and the NOEL is 6 ppm.

The classification of this study is Core Minimum.

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EFA: 68D90056 DYNAMAC No. 134-B March 22, 1989

DATA EVALUATION RECORD

ENDOSULFAN

Chronic Toxicity/Oncogenicity Feeding Study in Mice

APPROVED BY:

Robert J. Weir, Ph.D. Acting Department Manager Dynamac Corporation

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EPA: 68280056 DYNAMAC No. 134-E March 22, 1989

DATA EVALUATION RECORD

ENDOSULFAN

Chronic Toxicity/Oncogenicity Feeding Study in Mice

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DATA EVALUATION RECORD

GUIDELINE § 83-5

STUDY TYPE: Chronic toxicity/oncogenicity feeding study in mice.

ACCESSION/MAID NUMBER: 407924-01.

TEST MATERIAL: Endosulfan technical.

<u>SYNONYM(S)</u>: Thiodan; Benzolpen; 6,7,8,9,10,10-hexachlorc-1,5,5a,6,9,9a-hexahydro-6,9-methano-2,4,3-benzodicxathiepin-3cxide.

STUDY NUMBER(S): 745; TOXN No. 83.01113; Hoechst Report No. A38008.

SPONSOR: Hoechst Celanese Corporation, North Somerville, NJ.

<u>TESTING FACILITY</u>: Pharma Research Toxicology and Pathology, Heechst Aktiengesellschaft, Frankfurt, Federal Republic of Germany.

TITLE OF REPORT: Endosulfan-Substance Technical (Code: HOE 002671 OI ZD97 0003) Carcinogenicity Study in Mice, 24 Month Feeding Study.

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<u>AUTHOR(S)</u>: Donaubauer, H.H.

REPORT ISSUED: April 6, 1988.

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<u>CONCLUSIONS</u>: Under the conditions of the study, endosulfan was not encogenic when fed to male and female Hoe:NMRKf mice for 24 months at levels of 2, 6, or 18 ppm. There were no overt signs of toxicity or dese-related effects on clinical observations, food consumption, hematology, clinical chemistry, urinalysis, organ weights, macroscopic pathology, or microscopic pathology. Decreased survival in high-dose females and body weight reduction in high-dose males throughout the study were considered to be compound-related effects. Histologically, the incidence of lymphosarcoma was high in desed and control males and females. Based on the effects of endosulfan on nortality at 18 ppm, the LOFL is 18 ppm and the NOEL is 6 ppm.

Classification: Core minimum.

A. MATERIALS:

- <u>Test Compound</u>: Endosulfan technical; description: solid brown flakes; code No.:Hoe 002671 OI 2D97 0003; purity: 97.9%.
- Test Animals: Species: mice; strain: Hoe:NMRKf (SPF71); age: approximately 4 weeks; weight: males--23 g, females--22.5g at study initiation; source: Hoechst Breeding Colony, Frankfurt, West Germany.

B. STUDY DESIGN:

 <u>Animal Assignment</u>: Following 7 days of acclimation, animals were assigned to the following test groups with a computerized randomization procedure:

Test	Dose in Diet		Study	Şac	terim rifices 18 months)
Group	(ppn)	Males	Females	Males	Fenales
1 Control	0	60	60	20	20
2 LOW (LDT)	2	60	60	20	20
3 Mid (MDT)	e	60	60	20	20
4 High (HDT)	18 .	60	60	20	20

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Mice were individually housed in an environmentally controlled room with a 12-hour light/dark cycle.

The dose levels selected for this study were based on a 3month subchronic oral toxicity study (Huntingdon Research Center U.K., Report HST 229/831052) and a supplementary 42day feeding study (HOECHST AG, study No. 744, Report No. 85.0024) in mice. In the subchronic study, mice were fed the test material at doses of 2, 6, 18, or 54 ppm in the diet. Mice receiving 54 ppm, equivalent to a mean daily intake of 7.4 mg/kg body weight, exhibited convulsions and salivation and 12/20 males and 10/20 females died withir 6 weeks of study initiation. No mortality or other toxic effects were exhibited at the lower levels. The reported NOEL was 18 ppm, equivalent to a mean daily intake of 2.2 mg/kg body weight.

The supplementary 42-day feeding study, in which mice were fed the test material at doses of 18 ppm in the diet, was used to test the sensitivity of the mouse strain. Two of 10 female mice died on days 28 and 38; no other effects were reported. The maximum tolerated dose was considered to be 18 ppm (equivalent to a mean daily intake of 4.6 mg/kg body weight).

2. <u>Diet Preparation</u>: Premixes were prepared by dissolving appropriate amounts of the test compound in 20 g of vehicle (sesame oil DAB 7) and were then mixed with the appropriate amount of test diet at 14-day intervals. A premix with the same amount of sesame oil was prepared for the control animals. One kilogram of the endosulfan/sesame oil premix was added to additional feed weekly at the appropriate test diet concentrations. Storage conditions were not reported. During the study period, samples of the test compound were reanalyzed for content of active ingredient and samples of the treated food were collected and analyzed for stability, concentration, and homogeneity.

<u>Pesults</u>: Samples of the test compound analyzed four timesduring the study indicated that endosulfan was stable during the study duration (24 months) with content of active ingredient ranging from 96.7 to 97.9%. The mean recovery values of the compound from the test diet were 90.0, 91.7, and 93.3% of the nominal values for the 2-, 6and 18-ppm diets, respectively (Table 1). Concentration analyses for the control diet mixtures were not provided. The test diet was found to be stable up to 14 days of storage (Table 2).



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	Mean Concentration (mg/kg) at							
Monte	C ppm	2 ppn	6 ppr.	18 рот				
1 :	<u></u> >	2.0 ± 0.1	6.0 ± 0.2	17.1 ± 0.9				
	••	2.1 ± C.1	5.8 ± 0.7	18.1 ± 1.9				
12	• •	1.7 ± 0.1	5.4 ± 0.3	16.2 ± 2.1				
15		1.6 ± 0.1	4.8 ± 0.4	17.3 ± 0.5				
24	• •	1.8 ± 0.1	5.5 ± 0.1	15.5 ± 1.0				
Overall		1.8 ± C.2	5.5 ± 0.5	15.8 ± 1.0				
% Recovery	••	90	91.7	93.3				

TABLE 1. Analysis of Concentration of Endosulfan in Test Diet^a

*individual mean concentrations, overall mean diet concentration, and percent recovery were calculated by the reviewers.

The data for control (C ppm) diet were provided.

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	·		¥e.	an Conter	tration (75 (21 Car	:	· · · · · · · · · · · · · · · · · · ·	
	2 20	r on Day		. 6 0	on on tay		18	por or Da	<u> </u>
Month	C	7	1-	C	7	۹.	C	7	14
Initiai	1.93	1.89	2.05	é.13	6.1E	6.71	1E.2	17.6	18.5
3	1.82	22	1.55	5.54	e.33	6.44	17.3	17.7	16.2
6	1.69	1.56	1.8-	5.41	6.27	£.75	16.1	15.3	15.4
12	1.29	1.82	1.52	4.92	ć.32	6.04	18.4	15.2	16.2
18	1.71	1.59	1.81	4.70	4,75	5.85	17.0	15.2	17.1
24	1.81	7.54	1.59	5.49	4.81	5.34	14.3	12.3	15.3
Overal!	1.8:0.1	1.8:0.3	1.9:0.1	5.4±0.5	5.6±C.8	6.2:0.£	16.9:1.5	16.1±1.2	16.5:1.
Recovery	90	90	\$5	91	97	103	94	89	92

TABLE 2. Analysis of Stability of Endosulfan Test Diet^{*}

*individual mean concentrations, overall mean diet concentration and percent recovery were calculated by the reviewers.

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- Food and Water Consumption: Animals received food (Altromin-1321, standard pulverized diet) and water <u>ad</u> <u>libitum</u>.
- 4. <u>Statistics</u>: The following procedures were utilized in analyzing the numerical data. Body weights, hematology, clinical chemistry, and relative organ weights were analyzed using the parametric methods of Dunnett and Sidak and the nonparametric methods of Nimenyi/Dunnett and Nimenyi/Sidak. Reticulocyte counts were analyzed using the T-test and the Wilcoxon test. Kaplan Meier estimates and the Log-rank test were used for survival analyses.
- 5. <u>Quality Assurance</u>: A quality assurance statement was signed and dated April 22, 1985.

C. METHODS AND RESULTS:

1. <u>Observations</u>: Animals were inspected twice daily for signs of morbidity and mortality. Mice were examined weekly for neurological disturbances, opacity of the refracting medium of the eyes, impairment of dental growth, and changes in the oral muccsa. All mice were individually examined for palpable masses monthly for 6 months and twice monthly thereafter until study termination.

<u>Results</u>: The cumulative mortality and percent survival data are summarized in Table 3. At study termination (106 weeks), survival among high-dose females (28%) was significantly (p<0.05) lower than survival among the control females (45%). Survival in other dosed females and all dosed males were comparable to controls ranging from 37 to 55%. Survival in dosed animals of the satellite groups sacrificed at 12 months was comparable to controls; no deaths were reported in dosed or control animals scheduled for interim sacrifice at 18 months.

It was reported that there were no abnormal clinical sighs suggestive of a compound-related effect; individual data were not presented. It was reported that no neurological disturbances, impairments of dental growth, or changes in the oral muccsa were found in dosed animals.

Palpation of the skin was reported by the study author to reveal an equal number of masses in dosed and control mice; individual data were not reported.

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Cose Group	Mortality (Percent Survival) at Week 26 52 78 92 100								
(26	52			10c				
			<u>Ya es</u>						
C	3 (95)	5 (92)	12 (80)	16 (73)	27 (55)				
ź	1 (98)	(73) 3	14 (77)	22 (63)	33 (45)				
e	1 (98)	10 (53)	16 (73)	25 (58)	37 (38)				
18	2 (97)	15 (75)	17 (72)	23 (62)	35 (42)				
			Fenal es						
5	3 (95)	4 (93)	11 (82)	20 (67)	33 (45)				
2	2 (97)	2 (97)	10 (83)	21 (65)	36 (40)				
£	0 (100)	8 (87)	16 (73)	27 (55)	38 (37)				
18	1 (98)	12 (80)	23 (62)	30 (50)	43 (28)*				

TABLE 3. Cumulative Montelity and Percent Survival in Mice Fed Endosulfan for 24 Months[®]

"Percent survival was based on 60 mice/sex/dose of the main group.

"Significantly different from control values at p <0.05.

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2. Body Weight: Mice were weighed weekly.

<u>Results</u>: Table 4 presents mean body weight data at selected intervals. Body weights of males receiving 18 ppm were slightly but significantly (p<0.05) decreased from study weeks 2 to 17 and 26 to 31 when compared to concurrent controls; body weights of these animals remained slightly but nonsignificantly decreased thereafter to study termination. The body weights of other dosed males and all dosed females were similar to controls throughout the study.

3. Food Consumption and Compound Intake: Consumption was determined and mean daily diet consumption was calculated at the same intervals as body weight. Efficiency and compound intake were calculated from the consumption and body weight gain data.

<u>Pesults</u>: Food and mean compound consumption data at selected intervals are summarized in Table 5. Food consumption of dosed males and females were found to be similar to concurrent controls. Mean test compound intake was reported to be 0.28, 0.84, or 2.51 mg/kg/day for males receiving 2-, 6-, or 18-ppm endosulfan, respectively, and 0.32, 0.97 or 2.86 mg/kg/day for females receiving 2-, 6-, or 18-ppm endosulfan, respectively.

4. <u>Ophthalmological Examinations</u>: Ophthalmoscopic examinations were not performed but the opacity of the refracting medium of the eyes was observed weekly.

<u>Results</u>: These findings were similar in dosed and control mice.

5. <u>Hematology and Clinical Chemistry</u>: Blood was collected from the retro-orbital venus plexus of nonfasted mice at €, 12, 18, and 24 months for hematology and at 12, 18, and 24 months for clinical analysis from 10 mice/sex/group. The CHECKED (X) parameters were examined:

a. <u>Hematology</u>:

	Hematocrit (HCT)
Σ	Hemoglobin (HGB)
Σ	Leukocyte count (WBC)
X	Erythrocyte count (RBC)
Σ	Platelet count
Σ	Reticulocyte count (RETIC)
	Red cell morphology

X Leukocyte differential count^a
X Mean corpuscular HGB (MCH)
X Mean corpuscular HGB concentration (MCHC)
X Mean corpuscular volume (MCV)
Coagulation:thromboplastin time (PT)

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Recommended by Subdivision F (October 1982) Guidelines. Peticulocyte counts and differential leukocyte counts were determined from control and high-dose mice only.

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Dose Group	Kear Body Weight (g : S.D.) at Week								
(ppr)	1	13	25	52	75	92	104		
				Maies					
,C	27 : 2	35 ± 3	37 ± 3	38 : 4	39 ± 3	38 ± 3	,37 ± ,3		
2	26 ± 2	35 : 3	37 = 3	35 ± 4	35 ± 3	38 ± 3	37 = 3		
ć	27 ± 2	35 ± 3	3 ⁻ ± 4	39 ± 4	35 ± 3	37 : 3	36 z 3		
18	25 ± 2	34 ± 3*	35 : 4	37 : 3	37 z 4	36 ± 4	35 ± 4		
				Fenales					
c	23 ± 1	28 ± 2	30 ± 2	32 : 2	34 = 4	34 ± 3	3- = 3		
2	23 = 1	28 ± 2	30 ± Z	32 ± 2	33 : 3	33 : 3	33 z 3		
6	23 = 1	29 ± 2	30 ± 2	32 ± 3	34 ± 3	34 ± 3	34 ± 4		
12	23 ± 1	29 = 2	3* = 2	32 ± 4	34 2 2	3= ± 4	34 = 4		

TABLE 4. Pepresentative Results of Nean Eody Weights of Mice Fed Endosulfan for 24 Months^a

^aSased on mice of the main group.

"Significantly different from control values at p <0.05.

Dose		Mean Food Consumption (<u>g/day ± 5.D.) at Week</u>						
6-оцр (рот)	•	:3	20 20	52	75	9 2	154	(mg/kg/cay)
				Veles				
C.	5.5 ± C.5	5.1 ± 2.4	4.8 ± C.4	4.8 ± 0.6	5.3 = 0.4	5.4 ± 0.4	5,4 ± 1.0	5
2	5.5 ± C.c	5.C ± C.5	4.7 ± C.5	4.8 ± 0.4	5.3 ± C.4	5.4 ± G.4	5.5 ± 0.6	0.28
6	5.6 ± C.5	5.0 ± 0.5	4.7 ± 0.9	5.0 ± 0.5	5.3 ± 0.5	5.5 ± 0.6	5.ć 2 C.5	C.8-
15	5.4 ± 0.5	4.0 ± 0.4	4.4 ± 1.0	4.8 ± 0.5	4.9 ± 0.5	5.2 ± 0.4	5.2 ± 1.2	2.51
				Fenales				
C	5.4 ± 0.8	5.C ± C.8	4.9 = 0.6	4.8 ± 0.7	5.5 ± C.7	5.8 ± 0.8	5.9 ± C.8	e 0
2	5.3 ± 0.6	4.7 = 0.6	4.5 ± 0.5	4.5 ± 0.5	5.4 ± 0.5	5.4 ± 0.6	5.3 ± 1.1	6.32
6	5.3 ± 0.7	4.7 = 0.6	4.7 ± 0.6	4.6 ± 0.6	5.4 ± 0.6	5.4 ± 0.6	5.7 ± 0.6	5.97
18	5.3 : 0.6	6.9 ± 0.7	4.6 ± 0.8	4.8 ± 0.7	5.2 ± 0.6	5.5 ± 0.5	5.4 ± 0.8	5 2.86

TABLE 5. Representative Food and Compound Consumption of Nice Fed Endosulfan for 24 Months^e

Based on mice of the main group.

<u>Results</u>: There were no effects of dosing on hematology parameters. The study author reported that hematological changes due to aging of the mice (leukemia) were found in dosed and control animals.

b. Clinical Chemistry:

<u>Flectrolytes</u> Calcium Chloride Magnesium Phosphorus Potassium Sodium Other Albumin' Albumin'globulin ratio Blood creatinine Blood urea nitrogen Cholestercl' Globulins Glucose' Total bilirubin Direct bilirubin Total protein Triglycerides

- Enzymes X Alkaline phosphatase (ALP) Cholinesterase Creatinine phosphokinase
- X Serum alanine aminotransferase (SGPT)

Lactic acid dehydrogenase

X Serum aspartate aminotransferase (SGOT)[†] Gamma glutamyltransferase (GGT)

<u>Results</u>: There were no effects of dosing on the enzyme parameters measured.

- 6. Urinalysis: Urinalyses were not performed.
- 7. <u>Tissue Residue Determinations</u>: Sections of liver and kidney tissues from 10 mice/sex/group were collected at terminal sacrifice for residue determinations of a- and Bendosulfan. Only organs without macroscopic findings were used.

Fesults: The tissue residue results were not reported.

E. <u>Sacrifice and Pathology</u>: All animals that died and that were sacrificed on schedule were subject to gross pathclogical examination of integument, orifices, eyes, and internal organs. The CHECKED (X) tissues were collected for histological examination. In addition, the (XX) organs were weighed:

Recommended by Subdivision F (October 1982) Guidelines.

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	<u>Digestive System</u>
	Tongue
Х	Salivary glands
Σ.	Esophagus
Х	Stomach [*]
X	Duodenum
Х	Jejunum
Х	Ileum
Σ	Cecum
χ	Color
X	Rectum
	Liver
	Gailbladder'
	Fancreas
	the and water and
	Respiratory
	Trachea
XX	Lung

Cardiovasc./Hemat. X Aorta XX Heart X Bone marrow X Lymph nodes XX Spleen X Thymus

<u>Urocenital</u> XX Kidneys X Urinary bladder XX Testes X Epididymides X Prostate X Semin 1 vesicle XX Ovaries X Uterus

Neurologic XX Brain X Feripheral nerve (sciatic nerve) X Spinal cord (2 levels) X Fituitary X Eyes (certic perve)

(optic nerve)

<u>Glandular</u> XX Adrenals Lacrimal glanĝ

X Manmary gland X Thyroids Parathyroids

Harderian glands

<u>Other</u>

- Bone (sternum)
- X Skeletal muscle
- X Skin
- X All gross lesions

and masses X Nasal septum

Results:

Absolute organ weights were not <u>Organ Weights</u>: a. statistically analyzed by the study author. The relative lung and ovary weights of high-dose females were reported to be slightly but significantly (p<0.05) decreased at 12 months; the relative liver weights of high-dose males and relative ovary weights of high-dose females were found to be slightly but significantly (p<0.05) decreased at 18 months. These decreases were reported to be within the range of values for strainmatched historical controls and therefore were considered to be incidental changes. At 24 months, organ weights were similar in dosed and control males and females. Earlier changes in organ weights were not consistent over time or between sexes.

Recommended by Subdivision F (October 1982) Guidelines.

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- b. <u>Gross Patholcov</u>: The study author considered the incidence of gross lesions to be similar in dosed and control males and females.
- c. Microscopic Pathelogy:
 - 1) <u>Nonneoplastic</u>: Table 6 summarizes nonneoplastic findings in mice sacrificed at 24 months. Nonneoplastic findings at 12 and 18 months were reported by the study author to be unrelated to endosulfan administration. The nonneoplastic findings of dosed mice at 24 months were considered to be similar to concurrent controls or were considered to be normal age- and strain-related changes. The incidence of epithelial thickening in the urinary bladder was increased in dosed males and females; however, in the absence of a progression to clear proliferative change, this was considered to be of no toxicological importance.
 - 2) <u>Neoplastic</u>: Table 7 summarizes neoplastic findings in mice sacrificed at 24 months. Neoplastic findings at 12 and 18 months were considered spontaneous and of no toxicological importance. Lymphosarcoma was the nest commonly observed type of neoplasm among male and female mice. The incidence of these findings were similar in dosed and control mice, were considered to be normal ageand strain-related changes, and were not considered to be compound related.

D. STUDY AUTHOR'S CONCLUSIONS:

Dietary administration of endosulfan to male and female HOE:NMERF mice for 24 months at concentrations of 2, 6, or 18 ppm produced no overt compound-related signs of carcinogenicity. Body weight reduction was exhibited in high-dose males and increased mortality was exhibited in high-dose females. Minor changes in organ weights (lung, liver, and ovary) were observed in some high-dose males and females at 12 and 18 months only. Based on these results, the NOEL for endosulfan is 6 ppm.

	Dose Level (ppm)								
		Mal	es						
Organ Finding	0	2	6	18	0	<u> </u>	£	18	
<u>Lunes</u>	(58) ^t	(59)	(55)	(57)	(59)	(58)	(57)	(59)	
Congested	13	24	17	16	16	5	18	20	
<u>Nidneys</u> Cortical cysts(s)	(58) 2	(59) 13	(55) 8	(57) 2	(59) 0	(58) 0	(57) 1	(59) 0	
· ·	-								
Cortical foci of mononuclear cells	3	5	4	12	3	3	o	0	
<u>Urinary bladder</u> Minimal focal epi-	(58)	(59)	(55)	(57)	(59)	(58)	(57)	(59)	
thelial thickening	0	5	8	12	0	6	ç	10	
<u>Adrenals</u> Severe subcapsular	(58)	(59)	(55)	(57)	(59)	(58)	(57)	(59)	
foci of fibroblast like cells	- 0	0	2	4	. 8	10	3	8	
Areas of fatty degeneration	O	0	1	2	6 .	5	3	11	
<u>Seminal Vesicles</u> Distended	(58) 13	(59) 15	(55) 21	(57) 20					
Distended with peri pheral fibrosis	• 4	0	0	13					
<u>Uterus</u>					(59)	(58)	(57)	(59)	
Prominent fibrous tissue					0	1	1	3	
Dilated gland(s)					3	6	1	7	

TABLE 6. Representative Nonneoplastic Findings in Mice Fed Endosulfan for 24 Months^a

*Eased on mice of the main group.

¹Number in parentheses equals number of tissues examined.

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	Dose Leve: (ppm)								
	Males					-e-a	. #5		
Ongat:Finding	C	2	ć	3.	C	Z	¢	18	
<u>ntari</u>	(58)*	(59)	(55)	(57)	(52)	(52)	(57) 10	(5E) 5	
, ymphosarcoffa	4	7	19	ç	3	,6	10	2	
	(5E)	(59)	(55)	(57)	(59)	(5E)	(57)	(59)	
	9	7	12	10	2	14	11	ç	
Aprile	(53)	(54)	(49)	(50)	(55)	(51)	(50)	(49)	
Lymphosarcoma	7	5	7	5	7	12		B	
Sz een	(58)	(58)	(54)	(57)	(59)	(57)	(57)	(58)	
	10	11	12	9	9	17	16	11	
_ynphosa*cona									
<u>kioneys</u>	(5E)	(57)	(55)	(57)	(59)	(5E)	(57)	(59)	
Lynchosarcoma	9	8	Ş	8	7	10	10	9	
Pancreas	(56)	(55)	(53)	(55)	(58)	(57)	(53)	(56)	
Umphosarcome	5	4	5	Z	5	11	10	.£	
	(56)	(56)	(54)	(56)	(59)	(54)	(55)	(5E)	
	11	13	15	13	76	20	22	11	
urinary bladder	(56)	(58)	(54)	(52)	(58)	(51)	(53)	(54)	
Lymphesarcoma	1	3	3	2	3	6	6	T	
Uterus Lynprosarcome					(59) 3	(58) 5	(57) 5	(5°) 5	
<u>Gvaries</u> Cranulosa cell tumor					(59) 5	(56) 10	(56) 13	(59) 13	
Esopradus	(55)	(58)	(54)	(57)	(57)	(51)	(52)	(58)	
Lymphosanooma	1	5	3	2	2	- 4		5	
Lasal cavity	(54)	(56)	(50)	(53)	(56)	(56)	(53)	(55)	
	0	3	4	3	0	2	3	3	
Kulticentnic tumors	(17)	(16)	(20)	(17)	(26)	(29)	(27)	(18)	
	12	13	18	16	22	25	25	15	
smondsandona as cause of death	11	3	14	11	10	16	15	11	

TABLE 7. Representative Neoplastic Findings in Mice Fed Endosulfar for 24 Months

*Number in parentheses equals number of tissues examined; missing tissues were subtracted from total tissues examined.

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E. REVIEWERS' DISCUSSION AND INTERPRETATION OF RESULTS:

The study design was adequate for an oncogenicity study but not for a combined chronic toxicity/oncogenicity study, which were the guidelines referred to by the study author. There were some deficiencies in the conduct of the study and in data reporting. The control diet was not analyzed for concentration of test compound. Means, standard deviations, and percent recovery of the diet concentration and stability analyses were calculated by the reviewers. Homogeneity data were not reported. No statistical analysis was performed on absolute organ weights.

EPA Pesticide Assessment Guidelines, 1982, for chronic toxicity/oncogenicity studies suggest 15 clinical biochemistry determinations that should be performed on animals for chronic studies. Since sufficient blood was not available, only SGOT, SGPT, and alkaline phosphatase clinical biochemistry parameters were measured in this study. Urinalyses and complete ophthalmological examinations, which were not performed in this study, were also suggested in EPA Pesticide Assessment Guidelines.

Several discrepancies were reported in the histopathological data at 12 (1 discrepancy), 18 (1 discrepancy), and 24 months (5 discrepancies) involving numbers of missing tissues, numbers of tumor-bearing mice, and numbers of tumors in various tissues.

No intercurrent or sporadic deaths were indicated for 18-monthsatellite animals (10 mice/dose/sex); our reviewers consider this finding to be unusual. Histologically, the incidence of missing tissues appeared to be high in animals sacrificed at 12, 18, and 24 months; however, missing tissues were primarily from nonmajor organs.

It appears that an MTD was not achieved; the NOEL of 6 ppm was based on the effects of endosulfan on mortality and body weight at the high dose. However, survival was decreased in females only and body weights were only slightly decreased (1 to 2g) in males; these changes in body weight, although statistically significant, were not considered to be biologically significant by the reviewers. Dose levels for the study were based on results of a 3-month subchronic cral toxicity study in which convulsions and death were exhibited at 54 ppm, the highest dose tested. No toxic effects were exhibited at the lower doses tested, 2, 6, or 18 ppm. The reviewers suggest that a high-dose level of 36 ppm, for example, may have produced a greater incidence of compound-related effects.