Report Number: M-601

Date: March 12, 1980

Oral Rangefinder Study of T-2998CoC in Pregnant Rats

Experiment 1	No		:
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0680RR0018

Conducted At:

St. Paul, Minnesota

Dosing Period:

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January 20, 1980 to January 29, 1980

Study Director:

 2/24/81
Date

2/24/8_[

2/25/8/ Date





Exhibit 1231

State of Minnesota v. 3M Co., Court File No. 27-CV-10-28862

3M_MN02327266

Introduction

This oral rangefinder study was conducted to determine the upper dose level of T-2998CoC for a subsequent oral teratology study in rats. The study was sponsored by 3M Commercial Chemical Division, St. Paul, Minnnesota and was conducted by the Safety Evaluation Laboratory,

St. Paul, Minnesota. The study was conducted in accordance with the Safety Evaluation Laboratory's Standard Operating Procedures for such studies. The storage location for the raw data and a copy of the final report is maintained in the Safety Evaluation Laboratory's record archives.

Methods

Thirty-six time-mated Sprague-Dawley derived female rats from Charles River Breeding Laboratory were used in the study. The animals were indiscriminately removed from the shipping boxes by Animal Care personnel and placed in the rack of cages from the left to right starting at the top and working down. Later the Study Director assigned dose groups by vertical rows. The rats were housed individually in hanging stainless steel cages with wire mesh floors and fronts in a temperature and humidity controlled room. Purina Laboratory Chow and water were available ad libitum. The lights were on a 12 hour light/dark cycle.

The animals were observed daily from day 3 through day 20 of gestation for abnormal clinical signs. Body weights were recorded on days 3, 6, 9, 12, 15 and 20 of gestation and the rats dosed accordingly using a constant dose volume of 5 ml/kg of body weight. T-2998CoC was suspended in corn oil and administered daily by oral intubation at doses of 150, 100, 75, 50 or 25 mg/kg/day to groups of 6 rats on days 6 through 15 of gestation. A control group of 6 rats received only corn oil by oral intubation on the same days. On day 20 of gestation the rats were killed by cervical dislocation and each uterus, including its contents, was examined immediately to determine if the animal was pregnant. Because two previous teratology studies (Experiment Nos: 0680TR0008 and 0680TR0010) with chemically related compounds resulted in fetuses with teratogenic changes in the lens of the eye, a few fetuses were also taken at day 20 of gestation and examined for eye abnormalities.

Blood samples from three rats in each dose group were taken before the first dose and at day 20 of gestation. Liver specimens were also taken from the same rats on day 20 of gestation. The plasma samples and liver specimens were frozen and submitted to the sponsor.

Results and Discussion

The oral administration of T-2998CoC at 150, 100, 75, 50 or 25 mg/kg/day to rats during the period of organogenesis (days 6 through 15 of gestation) did not result in any deaths. A toxic effect of reduced body weight gain occurred between days 6 and 9 of gestation in the 150 mg/kg/day dose group (Table 1).

The two nonpregnant 150 mg/kg/day rats had a more severe effect on body

 $\frac{a}{b}$ Experiment No. 0680RR0018

3M_MN02327267

weight on day 9 of the study than the pregnant high dose dams (Appendix I). They lost a considerable amount of weight and one was observed to have urinary incontinence on days 11, 12 and 13. The pregnant dams of the 100, 75, 50 and 25 mg/kg/day dose groups did not have abnormal clinical signs and gained weight at comparable levels to the 0 mg/kg/day group.

Four fetuses were examined from each of four dams in the 150 and 25 mg/kg/day dose groups for eye changes. All of the readable fetuses sectioned had eye changes consisting of one or more of the following: large lens clefts, dark streak running one-half to three-quarters of the way through the lens or disorganized lens fibers (Table 2). The lens abnormalities occurred in the same location as those observed in the two Experiment Nos: 0680TR0008 and previous teratology studies (0680TR0010) on chemically related compounds. The abnormalities in this study appeared more pronounced than in the previous studies. In the previous studies, the teratogenic effect was a developmental eye abnormality which appeared to be an arrest in development of the primary lens fibers forming the embryonal lens nucleus, followed by secondary aberrations of the secondary lens fiber of the fetal nucleus. The same general morphological changes occurred in this rangefinder study with T-2998CoC.

Conclusion

The objective of determining an upper dose level for an oral rat teratology study was met in this study. The above results suggest that the 150 mg/kg/day dose level would be an appropriate high dose in a rat teratology study because of the toxic effect of reduced body weight gain. In addition to the toxic effect of reduced body weight gain, the teratogenic effect of lens abnormality was observed and is likely to be reproduced in a teratology study.

Table 1

Oral Rangefinder Study of T-2998CoC in Pregnant Rats

Mean Body Weight Gains of Pregnant Rats

With Standard Deviations (g)

	Day
	6 9 12 15 20
Control	30 18 21 29 76 4.2 7.4 7.5 1.6 10.7
150 mg/kg/day	21 5 35 2 1 2 84 5,5 17,8 8,8 13,8 12,1
100 mg/kg/day	29 15 17 19 84 4.1 5.1 4 4 12.6 13.5
75 mg/kg/day	27 11 21 19 74 6,6 10,0 2,7 10,5 12,6
50 mg/kg/day	10 16 37 77 6.5 3.7 5.6 7.3 16.6
25 mg/kg/day	24 16 24, 29 82 2.6 6.6 6.9 9.3 5.8

 $[\]frac{a}{c}$ Significantly higher than the control (Dunnett's t test p < 0.05)

Oral Rangefinder Study of T-2998CoC in Prequant Rats Ratios of Fetuses with Eye Changes to Fetuses Examined

High Dose Group (150 mg/kg/day) Low Dose Group
(25 mg/kg/day)

16/16

15/15<u>b</u>

4.

 $[\]frac{a}{b}$ Four fetuses examined from each of four dams $\frac{b}{b}$ One fetus not examined because eye architecture destroyed in sectioning

Appendix I

Oral Rangefinder Study of T-2998CoC in Pregnant Rats

Individual Body Weights (g) and Mean Body Weights
with Standard Deviation for Pregnant Rats

				Da	У		
		3	6 	9 	12	15	20
er MG.	'K(ኤ'ኒክተ	h.					
NIR	316	194	223	244	269	297	382
N1R	B17	186	214	228	262	25.	376
N1E	318	192	217	227	253	ک اتا کے	265
N1R	319	207	239	250	258	285	360
N1R	346	190	231	257	280	311	369
Mi	EAN	195			264	293	369
STAN.	DEM	7. 7	1 0. 3	11.5	10.5	11 . 5	8. 2
NON 1	PREGMA	NT AN	IIMALS	•			
		A COS	1991 19	224	215	223	222
N1R	فائيك	704	tii, -b'	t i.a. * 1	E L - C		
N1R		104	414 ak. ak.		ay	ban ten	
N1R	<u>320</u>					2.5	ناك -
	326 MG./KG/			D	ay		ريان -
		z Z		D 9 216	ay 12 257	15 287	367
150 (MG./KG/	z Z	15	D	ay 12	25	367 344
150 (018	MG/KG/ 321		222 218 191	216 217 222	257 257 257 244	287 261 243	367 344 314
150 (018 018	MG/KG/ 321 324	202 193	0 222 218	216 217	ay 12 257 257	257 261	367 344
150 (01R 01R 01R 01R	MG/KG/ 321 324 325 347 EAN	202 193 177 206	222 218 191 232 216	216 217 222 226	257 257 257 244 262 255	287 261 243 278 267	367 344 314 378 351
150 (01R 01R 01R 01R	MG/KG/ 321 324 325 347 EAN	202 193 177 206	222 218 191 232 216	216 217 222 226	257 257 257 244 262 255	287 261 243 278	367 344 314 378 351
150 01R 01R 01R 01R	MG/KG/ 321 324 325 347 EAN	202 193 197 206 195 12. 9	222 218 191 232 216 17, 5	216 217 222 226 228 4.6	257 257 257 244 262 255	287 261 243 278 267	367 344 314 378 351
150 01R 01R 01R 01R	MGZKGZ 321 324 325 347 EAN DEV PREGNA	202 193 197 206 195 12. 9	222 218 191 232 216 17.5	216 217 222 226 228 4.6	257 257 257 244 262 255 7. 7	287 261 243 278 267, 19, 4	367 344 314 378 351 28. 3

Appendix I (Continued)

Oral Rangefinder Study of T-2998CoC in Pregnant Rats Individual Body Weights (g) and Mean Body Weights with Standard Deviation for Pregnant Rats

				Da	y		
			to	9	12	15	26
(Photo)	Mar Fra	441					
Pak	326	164	193	218	229	253	327
HIR	327	214	240	248	268	265	331
H1H	328	at a	266	تعادات	317	349	452
Pik	329	200	235	245	256	265	353
F1H	330	185	218	234	248	268	363
PIR	348	189	218	240	263	296	371
M	EAN	202	232	247	264	282	366
STAN	DEV	33. 6	31. 3	30. 4	29. 6	34. 9	45. 5

		Day	7			
3	6	9	12	15	20	

75	MG.	4816	10	f4't'

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01R	331	192	221	243	265	268	346
01R	332	198	213	228	249	271	346
01R	333	172	203	215	235	263	346
01R	334	211	243	236	261	276	326
01R	335	193	216	225	244	268	321
01R	349	200	231	248	265	293	383
MEI STI FIN.	AN DEV		221 14. 1				346 20. 6

Appendix I (Concluded)

Oral Rangefinder Study of T-2998CoC in Pregnant Rats

Individual Body Weights (g) and Mean Body Weights with Standard Deviation for Pregnant Rats

Day							
		3	6	9	12	15	20
50 MC	i/KG/I	vff5f					
R1R	336	193	219	236	253	276	350
R1R	337	177	201	213	235	259	338
R1R	338	226	251	262	283	314	397
R1R	339	170	198	218	237	254	308
R1R	340	187	226	245	267	304	378
R1R	350	192	229	243	276	308	382
Mi	EAM	191	221	236	259	286	359
STAN	DEV	19 4	19.6	18. 2	20 - 1	26. 2	33. 0

		Day				_
3	6	9	12	1.5	20	

25 MG/KG/DHY

S1R	342	216	239	266	283	304	388
S1R	343	207	234	249	279	304	383
S1R	344	185	208	227	253	292	369
S1R	345	200	219	233	249	270	348
S1R	351	205	233	238	268	307	398
ME: STAN.	AN DEV		227 12. 8	243 15. 4	266 15. 2		377 19. 4

NON PREGNANT ANIMALS

S1R 341 187 203 219 220 228 238

DISTRIBUTION LIST

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