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OF CURIUM TRICHLORIDE

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### CRYSTAL STRUCTURE AND LATTICE PARAMETERS OF CURIUM TRICHLORIDE

J. C. Wallmann, J. Fuger, J. R. Peterson, and J. L. Green

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# CRYSTAL STRUCTURE AND LATTICE PARAMETERS OF CURIUM TRICHLORIDE

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October 1965

#### ABSTRACT

CmCl $_3$  has been found to exhibit the hexagonal UCl $_3$  structure. Powder data from two samples give average lattice parameters  $a=7.380\pm0.006A$  and  $c=4.186\pm0.010A$ , where the error limits are the 95% confidence interval calculated using the standard statistical method for the average of two independent determinations.

#### INTRODUCTION

The crystal structure and lattice parameters of curium trichloride have been reported by Asprey, Keenan, and Kruse. (1)

The lattice parameters for curium trichloride calculated from data obtained by us are in serious disagreement with the work referred to above. Since tabulated values for the ionic radii of the tri-positive actinide ions have been derived largely from crystallographic data on the trichlorides, and since these radii are of value in predicting structures of a variety of compounds, we feel that it is of importance that the discrepancy in the curium trichloride data be resolved. We describe in some detail below the derivation of our experimental data, and the methods of calculation used to arrive at the lattice parameters.

#### EXPERIMENTAL

#### A. Materials

X-ray diffraction data were obtained on two samples of curium trichloride, derived from separate curium stock solutions which had been subjected to different purification procedures.

The starting material for the first sample consisted of a mixture of about 12 mg of  $Am^{243}$  and 12 mg of  $Cm^{244}$ , together with small amounts of common impurities such as Ca, Mg, Al, and Fe.

An americium-curium separation was achieved by two successive ion exchange elutions, using alphahydroxyisobutyric acid as the eluting agent. (2)

The separated curium fraction was loaded onto Dowex 50 ion exchange

resin contained in a quartz tube and moved to the bottom of the resin bed column with 2 M HCl. Curium was then stripped from the resin using 6 M HCl, the eluate being collected in three fractions. A sixty-microgram sample of the 7.5 mg middle fraction was analyzed for impurities by spectrographic emission analysis using copper spark excitation. Limits of detection for various elements by this method have been given in a previous publication. (3) The only impurities detected in the 7.5 mg curium fraction were 0.04 atom percent americium, 0.16 atom percent silicon and 0.1 atom percent calcium.

About 40 micrograms of curium were taken from this stock and transferred to a quartz microcone. The hydrochloric acid solution was evaporated to dryness, treated with a few microliters of freshly distilled nitric acid, re-evaporated to dryness and heated in air to about 600°C.

A portion of the curium oxide obtained in this way was scraped free, using a platinum wire scraper, and transferred to a quartz x-ray capillary. The capillary was connected to a vacuum line and the oxide treated for about ten minutes with one-half atmosphere of HCl<sub>(g)</sub> at  $400^{\circ}$  C. Excess HCl and water vapor formed by the reaction were removed by pumping, after which fresh HCl was again added to the system. This process was repeated several times. In the final treatment the sample was allowed to cool to room temperature in the presence of HCl, which was then pumped off, and the capillary sealed for examination of the chloride by x-ray diffraction.

Diffraction lines from this preparation were recorded on Film 1500-A.

Starting material for the second trichloride sample was about 18~mg of  $\text{Cm}^{244}$  containing a small amount of americium together with significant amounts of Fe, Al, Ca, etc.

An americium-curium separation was effected by ion exchange, using Dowex 1  $\times$  8 resin and 4.2 M LiNO at pH 2.15 as eluting agent. (4)

The curium fraction from this column operation was treated with excess  $\mathrm{NH_4OH}$  to precipitate  $\mathrm{Cm(OH)_3}$ , which was washed several times and then dissolved in a minimum volume of 0.1 M HCl. The 0.1 M HCl solution was transferred to the top of a Dowex  $50 \times 4$  resin bed contained in a quartz tube. The curium was then moved to the bottom of the resin column with 2 M HCl and stripped with 6 M HCl. This method is highly effective in separating curium from such common impurities as Al, Ca, and Fe.

A 24  $\mu gm$  sample of the purified curium sample was analyzed by copper spark emission analysis. The only impurity detected was 0.42 atom percent of americium.

A few microliters of this curium stock solution were transferred to a clean platinum plate, evaporated to dryness, treated with 10  $\mu$ l of freshly distilled nitric acid, re-evaporated and heated in air to  $600^{\circ}$  C. Subsequently the sample was re-heated in air in an open tube furnace for 10 minutes at  $675^{\circ}$  C.

After cooling, a portion of the curium oxide was transferred from the platinum plate to a quartz capillary and heated in a stream of anhydrous HCl(g) for 35 minutes at 400-500°C. The tip of the capillary was then sealed, the sample cooled to room temperature, excess HCl pumped off, and the capillary sealed.

The diffraction lines were recorded on Film 2069-A, using diffraction equipment described below.

# B. Diffraction equipment

The diffraction equipment consisted of a Model 80-000 Jarrel-Ash Microfocus x-ray source and a ll4 mm diameter Norelco Precision

Powder Camera, manufactured by the Phillips Electronics Instrument Company.

#### RESULTS AND DISCUSSION

Line positions on Film 1500-A were read twice and averaged; those on 2069-A were read once.

Following indexing, the data were transferred to cards for  $709^4$  computor determination of the most probable lattice parameter values, according to a least-squares fit of the differences between experimental  $\sin^2\theta$  values and those calculated from the assigned indices. Two computational programs were used: the LCR-2 program developed by Williams (5) and the MET-124 program of Mueller and Heaton. (6)

Lattice parameters calculated by the two programs were the same to less than 0.001 A.

Line intensities were calculated theoretically, on the basis of the assumed UCl3-type hexagonal structure, by using the POWD program developed by Smith. (7)

In Table 1 below we present a comparison of observed and calculated  $\sin^2\!\theta$  values, as well as observed and calculated line intensities for both CmCl<sub>z</sub> preparations.

TABLE 1. OBSERVED AND CALCULATED INTENSITIES AND  $\sin^2 \theta$  VALUES FOR CmCl3

		2069A		1500A	
	$_{ ext{Sin}}$ 2 $_{ heta}$ (b) $_{ ext{I}}$ (c)	$\sin^2\theta$	<pre>[d)</pre>	$\sin^2\!\theta$	I(d)
hkl	Calc. Calc.	Obs.	Obs.	Obs.	Obs.
100 в	0.0119	0.0118	7	0.0118	. 7
100 α	0.0145 98	0.0146	10	0.0145	10
110 β	0.0356	0.0355	. 7	0.0357	. 4
101 β	0.0395	0.0397	8	0.0397	. 4
110 $\overline{\alpha}$	0.0436 66	0.0437	10	0.0436	8
101 $\overline{\alpha}$	0.0485 100	0.0486	10	0.0486	10
200 <del>α</del>	0.0582 23	0.0582	8.5	0.0582	4
111 β	0.0632	0.0635	2	0.0634	2
201 β	0.0751	0.0750	7	0.0748	14
111 $\overline{\alpha}$	0.0776 11	0.0777	1,	0.0777	4
120 в	0.0830	0.0832	2	0.0831	2
201 <del>a</del>	0.0921 91	0.0920	10	0.0921	10
120 α	0.1018 16	0.1018	5.5	0.1017	14
300 в	0.1068	0.1066	14	0.1068	2.5
002 в	0.1106	0.1108	7	0.1107	. 4
121 β	0.1107	) 0.1100		0.000	
102 β	0.1225	0.1228	1	•	<b>-</b>
300 <del>a</del>	0.1309 35	0.1307	8.5	0.1310	7
002 α	0.1357 16	0.1356	10	0.1359	10
121 $\overline{\alpha}$	0.1358 76	}	<b>√</b>	· · · · · · · · · · · · · · · · · · ·	
220 в	0.1423	0.1423	1	0.1422	1
112 β	0.1462	0.1463	2	0.1463	2

TABLE 1. (Cont'd)

				2069A		1500A	
	$\sin^2_{\theta}(b)$	<sub>I</sub> (c)		$\sin^2\theta$	I(q)	$\sin^2\!\theta$	I(q)
hkl	Calc.	Calc.	٠	Obs.	Obs.	Obs.	Obs.
102 α	0.1502	10		0.1503	4	0.1504	4
130 в	0.1542			0.1544	1		<u>.</u> .
202 β	0.1581			0.1582	1.		
220 α	0.1746	14		0.1748	7	0.1746	14
112 $\bar{\alpha}$	0.1793	21		0.1794	8	0.1796	7
131 β	0.1819			0.1825	2		
130 α	0.1891	8		0.1893	7	0.1893	, 4
122 β	0.1936		1	0.1941	77	0.3003	4
202 <del>a</del>	0.1939	8	5	0.1941	7	0.1941	
302 β	0.2174	÷	1	0.0177		0.2177	0.5
401 в	0.2174	ι.	5	0.2177	2	0.2111	2.5
131 $\alpha_1$	0.2227	15		0.2232	8		cont
131 $\overline{\alpha}$	0.2230	22		-		0.2233	. 7
131 $\alpha_2$	0.2238	6		0.2239	14		•
400 α <sub>1</sub>	0.2324	3		0.2330	14		<b>-</b> .
400 <del>a</del>	0.2328	4		<u>.</u>		0.2328	2.5
212 α <sub>1</sub>	0.2371	6		0.2374	5		<b>-</b> ·
212 $\bar{\alpha}$	0.2375	9				0.2379	4 `
140 в	0.2491		•	0.2491	2	0.2492	2
222 β	0.2529		1	0.2532	4	0.2532	4
231 β	0.2530		5	0.2772	<del>''</del>	0.2772	- <del></del>
302 α_	0.2662	15		0.2667	7	· · · · · · · · · · · · · · · · · · ·	-

TABLE 1. (Cont'd)

			·	2069A		1500A	
	•	Sin <sup>2</sup> θ(b	) <sub>I</sub> (c)	$\sin^2\!\theta$	(d)	$\sin^2\!\!\dot{ heta}$	I(q)
hkl		Calc.	Calc.	Obs.	Obs.	Obs.	Obs.
401 $\alpha_1$	,	0.2662	5	0.2674	14	· <b>-</b>	•
302 <del>α</del>	·	0.2666	23	e e	-	0.0070	0
401 α		0.2667	7		· · · }	0.2670	8
230 α <sub>1</sub>		0.2760	3	0.2763	4		
230 <del>α</del>		0.2764	5		-	0.2761	14
203 в		0.2963		0.2968	2	0.2971	.2
500 в		0.2965		0.2900	2	0.2911	2
140 α1		0.3050	9 .	0.3053	4		
140 $\bar{\alpha}$		0.3055	13	, .	eed .	0.3058	5
140 α2		0.3065	14	0.3062	2	· · · · · · · · · · · · · · · · · · ·	
222 $\alpha_1$		0.3097	7	0.3096	7 .		
231 $\alpha_1$		0.3098	18		1	• • • • •	
222 ā		0.3102	11		-	0.3108	9
$231 \overline{\alpha}$		0.3103	27		}	0.7100	
222 α <sub>2</sub>		0.3113	4	0.3117	<u>L</u>	·	
231 α2		0.3114	9	0.711			
103 $\alpha_1$		0.3193	5 .	0.3198	4		
103 α		0.3198	7		<b>-</b>	0.3195	4
132 α <sub>1</sub>		0.3243	14	0.3242	14	· · ·	
132 🕏		0.3248	7	•	-	0.3246	4
123 β		0.3319		0.3324	2	0.3319	5

TABLE 1 (Cont'd)

			2069A		1500A	
	$\sin^2\!\theta$	[(c)	$\sin^2\!\theta$	(d)	$\sin^2\!\theta$	(d)
hkl	Calc.	Calc.	Obs.	Obs.	Obs.	Obs.
142 β	0.3597		2 7500	0	0.75.05	
241 в	0.3598		0.3598	2	0.3597	2
203 α1	0.3628	7	0.7677	14		
500 α <sub>1</sub>	0.3631	1	0.3637	4		
203 α	0.3634	11		7	0.3637	. 4 .
500 α	0.3637	2		ا	7 0.3637	· <del>+</del> ·
402 α <sub>1</sub>	0.3678	3	0.3682	2	-	•
402 α	0.3684	14			0.3687	2
501 α <sub>1</sub>	0.3970	6	0.3976	4	<u></u>	
501 ā	0.3976	10	<del>-</del>		0.3972	14
123 α <sub>1</sub>	0.4064	9	0.4065	4	-	
123 α	0.4071	14	-		0.4072	7
123 α2	0.4084	5	0.4082	2	••	
232 α <sub>1</sub>	0.4114	4	0.4122	2	<b>-</b>	
142 $\alpha_1$	0.4404	10	0.4406	7	• •	
241 $\alpha_1$	0.4405	7	} 0.4400	ľ	- ·	
$142 \overline{\alpha}$	0.4412	15	-		0.4413	10
241 ā	0.4413	. 10			( )	10
142 α <sub>2</sub>	0.4426	5	0.4432	4		
241 α <sub>2</sub>	0.4427	3	1	T	<del>-</del>	
233 В	0.4742		0.4749	2	<b>-</b>	•
151 α <sub>1</sub>	0.4841	8	0.4841	7.	-	

TABLE 1 (Cont'd)

•			2069A	~~~~~	1500A	
	$\sin^2_{\theta}(b)$	_ ;	Sin <sup>2</sup> θ	I(q)	$\sin^2\!\theta$	. I(q)
hkl	Calc.	Calc.	Obs.	Obs.	Obs.	Obs.
151 α	0.4849	11		•	0.4846	5
151 α2	0.4865	4	0.4867	4	<del></del>	•
133 $\alpha_1$	0.4935	5	0.4937	4	<del>-</del>	
133 α	0.4944	7	••		0.4946	4
600 α <sub>1</sub>	0.5229	2	0.5237	0.5	-	
332 α <sub>1</sub>	0.5276	4	0.5277	2		
332 ā	0.5285	6	<b>-</b>		0.5272	3
403 α <sub>1</sub>	0.5371	2	0.5373	<u>1</u>	· 	•
340 α <sub>1</sub>	0.5374	2	(0.))()	<b>T</b>	_	
403 ₹	0.5380	3	-	}	0.5371	4
$340 \overline{\alpha}$	0.5383	2		}	0.0012	
?	?	?	a · ·		0.5498	
?	?	?	а		0.5566	
250 α <sub>1</sub>	0.5664	5	0.5663	2	-	
341 α <sub>1</sub>	0.5713	5	0.5706	2		
$341\overline{\alpha}$	0.5722	8	<b>-</b>		0.5710	14
$341 \alpha_2$	0.5741	3	0.5741	2		
$233 \alpha_1$	0.5807	8	0.5806	7	. <u>.</u> .	
233 <del>a</del>	0.5816	12	. <del>.</del>		0.5808	5
233 α <sub>2</sub>	0.5836	4	0.5841	14	<del>-</del>	
114 $\overline{\alpha}$	0.5863	14	-	7	0.5851	2
512 <del>α</del>	0.5867	2	•	j	,	~ <del>-</del>

TABLE 1 (Cont'd)

	•		2069A		1500A		
	$\sin^2_{\theta}(b)$	I(c)	$\sin^2 \theta$	(d)	$\sin^2\theta$	(d)	
hkl	Calc. C	alc.	Obs.	Obs.	Obs.	Obs.	
204 <del>a</del>	0.6008	2			0.6014	2	
153 β	0.6166		<b>-</b> .		0.6169	2	
$124 \overline{\alpha}$	0.6445	3	-		0.6447	2	
602 α <sub>1</sub>	0.6583	3	0.6585	7	0.6577	4	
161 α <sub>1</sub>	0.6584	5	J. 0.000				
602 α <sub>2</sub>	0.6616	1	0.6618	4	0.6620	2	
161 α2	0.6617	3	) 3.332				
503 α <sub>1</sub>	0.6678	4	0.6679	4.	0.6681	2 .	
$304 \alpha_1$	0.6725	5	0.6725	<b>7</b>	0.6727	14	
342 α <sub>1</sub>	0.6728	3	) 0.0129	• •			
304 α <sub>2</sub>	0.6758	3	0.6766	4	0.6759	2	
342 α <sub>2</sub>	0.6762	l	}	•			
440 a <sub>1</sub>	0.6972	2	0.6976	1	<b>-</b>		
252 α <sub>1</sub>	0.7019	9	0.7019	7	0.7023	14	
252 α2	0.7054	14	0.7056	. 4	<b>-</b>		
243 α <sub>1</sub>	0.7114	5	0.7115	7	0.7115	14	
243 α <sub>2</sub>	0.7149	3	0.7151	5	. <del>-</del>		
224 α <sub>1</sub>	0.7160	3	<b>-</b>		0.7162	4.	
$13^{4} \alpha_{1}$	0.7306	5	<b>-</b> '.		0.7312	2	
351 α <sub>1</sub>	0.7455	7	0.7456	14	0.7458	4	
351 α <sub>2</sub>	0.7493	3	0.7491	2	_	•	

TABLE 1 (Cont'd)

			2069A		1500A	
•	$\sin^2\!\theta(b)$	(c)	$\sin^2\theta$	I(q)	$\sin^2\!\theta$	·I(d)
hkl	Calc.	Calc.	Obs.	Obs.	Obs.	Obs.
153 α	0.7550	.7	0.7551	5	0.7553	5
260 α1	0.7553	2	0.1771	· ) .	0.1777	
153 $\alpha_2$	0.7587	4	0.7592	14	0.7593	4
260 α <sub>2</sub>	0.7590	1	0.1792	<b>.</b>	· · · · · · · · · · · · · · · · · · ·	. 4
404 a <sub>1</sub>	0.7741	2	0.7747	2	0.7752	2
261 α <sub>1</sub>	0.7891	6	0.7891	7	0.7888	4
261 α <sub>2</sub>	0.7930	3	0.7933	4 .	· <b>-</b>	
$234 \alpha_1$	0.8177	3	-		0.8180	2
170 α	0.8279	6	0.8279	2	0.8282	2
170 α2	0.8320	3'	0.8328	4	0.8328	4
442 a <sub>1</sub>	0.8326	. 6.	0.0720		0.0)20	7
442 α <sub>2</sub>	0.8367	3	0.8366	2	<b>-</b>	
$343 \alpha_1$	0.8421	7	0.8422	2	0.8423	2
343 α <sub>2</sub>	0.8463	3	·			
144 α1	0.8467	.10	0.8466	7	0.8475	7
352 $\alpha_1$	0.8471	<u>)</u>				
144 α2	0.8510	5	0.8512	. ),	0.8516	4
352 α <sub>2</sub>	0.8513	2	0.0912	*	0.0010	4
262 α <sub>1</sub>	0.8907	4	0.8907	2	0.8911	2
205 α <sub>1</sub>	0.9046	7	0.9044	7	0.9050	5
504 α <sub>1</sub>	0.9048	3	/ O. 70++		0.3070	)

TABLE 1 (Cont'd)

				2069A		1500A	
	$\sin^2_{\theta}(b)$	I <sup>(c)</sup>		Sin <sup>2</sup> θ *	I(q)	$\sin^2\theta$	I(q)
hkl	Calc.	Calc.		Obs.	Obs.	Obs.	Obs.
205 α <sub>2</sub>	0.9091	4	}	0.9090	2 .	_	
504 a2	0.9093	1	1	0.9090	<u>-</u>		
360 α <sub>l</sub>	0.9150	5		0.9151	2	<del>-</del> ·	
360 α <sub>2</sub>	0.9196	3	1	0.9194	7	<b>-</b>	
451 α <sub>1</sub>	0.9198	11	}	0.9194		0.9204	4
163 α <sub>1</sub>	0.9293	11		0.9290	7	0.9297	5
163 α2	0.9339	4	1	0.9334	7	0.9343	. 5
$334 \alpha_1$	0.9339	7	1	○• <i>ラ</i> ノノ <sup>+</sup>			. ,
125 α <sub>1</sub>	0.9481	14		0.9478	7	0.9486	7
125 α <sub>2</sub>	0.9528	8		0.9529	14	0.9535	4
172 α1	0.9633	27	1.	0.9630	7	0.9635	7
801 α <sub>1</sub>	0.9634	8	}	0.9000			
172 α2	0.9681	14	1	0.9680	4	0.9686	14
801 α <sub>2</sub>	0.9682	4 .	5	0.3000	T .	,000	

# TABLE 1 (Cont'd)

<sup>a</sup>The two unindexable trace lines at  $\sin^2\theta$  equal to 0.5498 and 0.5566 were independently observed only on film 1500A. On re-examination of film 2069A, these features were definitely located; however, the intensities of the reflections were very low.

<sup>b</sup>Calculated using a = 7.380A and c = 4.186A with  $\lambda_{\overline{\alpha}}$  = 1.54178A,  $\lambda_{\alpha_1}$  = 1.54051A,  $\lambda_{\alpha_2}$  = 1.54433A and  $\lambda_{\beta}$  = 1.39217A.

<sup>c</sup>Calculated using the POWD intensity program assuming the atomic coordinates of UCl<sub>3</sub> and scaled such that the strongest line has an intensity of 100.

d Estimated visually relative to a value of 10 for the strongest line.

The following features of the data recorded in Table 1 should be noted:

- 1) All lines except two appearing on each film have been indexed and included in the computation of the lattice parameters.
  - 2) The two unassigned lines appear at barely detectable intensity.
- 3) The quality of film 2069A is superior to that of 1500A in the sense that the pattern is clearer ( $\alpha_1$   $\alpha_2$  separations are noted at  $\sin^2\theta$  = 0.223 in 2069A as compared with  $\sin^2\theta$  = 0.660 in 1500A).
- 4) For either film the greatest difference between an observed and a calculated  $\sin^2\!\theta$  is 0.0013.
- 5) There are no significant discrepancies between observed and calculated intensities for either film.

- 6) Lattice parameters calculated for the two preparations are the same within 0.001A.
- 7) From film 2069A the observed lattice parameters are calculated to be a =  $7.3803 \pm 0.0002A$  and c =  $4.1862 \pm 0.0002A$ , while those from film 1500A are: a =  $7.3793 \pm 0.0003A$ , c =  $4.1847 \pm 0.0002A$ . The above error limits are standard deviations for the individual patterns computed using the LCR-2 program. Drs. Asprey and Keenan have been kind enough to inform us that the films on which their computations were based were of relatively poor quality and exhibited no lines having diffracting angles above  $45^{\circ}$ . These facts probably account for the differences between their calculated parameters and ours.

The interval agreement of the powder data is consistent with error limits of ±0.001A for both lattice parameters. The use of these limits is in accordance with customary practice. From a chemical standpoint, however, it is suggested that error limits based entirely on the agreement of independent determinations would be more meaningful. Treating the results presented here as two independent determinations of the lattice parameters, the application of standard statistical methods to the average, accounting for nonstatistical sampling, gives for the 95% confidence interval:

 $a = 7.380 \pm 0.006A$   $c = 4.186 \pm 0.010A$ 

This provides a statistically meaningful basis for comparison with other groups of independent determinations. It is felt that information of this sort would be of great assistance in the recognition of anomalies due to the effect of purity, nonstoichiometry, radiation damage, etc.

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# FOOTNOTES AND REFERENCES

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