Synthesis of Thallium Platinate at High Pressure

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The double oxide $Tl_2Pt_2O_7$ has been prepared by reaction of Tl_2O_3 with platinum metal or PtO_2 at 1000° and 40 kbars pressure. The brown solid is insoluble in aqua regia and is thermally stable to 750° at atmospheric pressure. The cell is face-centered cubic with $a=10.132\pm0.004$ A and the structure is assumed to be of the cubic pyrochlore type based on space group Pt_3 . The measured density is 11.12 g/cm³ and the computed value is 11.63 g/cm³, with Z=8. Fifteen diffraction lines were observed and used in a least-squares refinement to determine the one unknown coordinate. The R factor is 3.6% based on observed maxima only. Each platinum is bonded octahedrally to six oxygens at 2.08 ± 0.04 A, each thallium to six oxygens at 2.32 ± 0.04 A and two oxygens at 2.19 ± 0.002 A. Some general aspects of the pyrochlore structure are discussed. The infrared spectrum of $Tl_2Pt_2O_7$ (to 200 cm $^{-1}$) has four sharp maxima at 684, 562, 449, and 363 cm $^{-1}$.

Introduction

Information concerning the preparation and properties of anhydrous platinum oxides is scanty and often contradictory. Three oxides (PtO, Pt₃O₄, and PtO₂) and two double oxide compositions (M^IPt₃O₄ and M₂^I-PtO₃, where M is lithium or sodium) have been reported in relatively recent publications.

Moore and Pauling² treated PtCl₂ with KNO₃ and on the basis of very meager crystal data, concluded that the product was PtO, isostructural with the corresponding palladium oxide. The existence of PtO has been questioned by later workers.³ Galloni and Roffo⁴

determined the structure of several crystals of a cubic oxide which had been obtained from a platinum wire electrode. The assigned formula was Pt₃O₄, but confirmatory analysis was not possible on the small sample, and its true composition remains in doubt. Ariya, et al.,⁵ reported that the oxidation of platinum sponge at oxygen pressures to 310 bars produced only one thermodynamically stable oxide, Pt₃O₄. Its structure, however, was not that of the Galloni compound. Platinum dioxide has been prepared by fusing H₂PtCl₆ with an alkali metal nitrate.^{3,6}

Waser and McClanahan⁷ report that cubic NaPt₃O₄

⁽¹⁾ Based on work performed under the auspices of the U. S. Atomic Energy Commission. Paper presented at 153rd National Meeting of American Chemical Society, Miami Beach, Fla., April 1967.

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is the product obtained when chloroplatinic acid is fused with sodium carbonate at 850°, but Galloni and Busch³ contend that sodium is not an essential constituent of the oxide and that its true composition is Pt₃O₄. This structure may consist of a solid solution having the formula Na_xPt₃O₄. Scheer, et al.,⁵ report that the fusion of mixtures of platinum metal with lithium or sodium carbonates in a magnesia boat leads first to the formation of the double oxide composition M_xPt₃O₄ (as described above) and then to M₂PtO₃. The latter composition is obtained after the reactants have been heated gradually over a 20-hr period to 1000°.

The present paper describes the preparation and properties of a new platinum oxide, Tl₂Pt₂O₇. This oxide was first synthesized during an investigation into the effects of high pressure on the stability relationships between cubic type C and monoclinic type B rare earth sesquioxides.⁹ When the isostructural (type C) Tl₂O₃ was studied at high pressure, a reaction occurred which led to the formation of the thallium-platinum double oxide.

Experimental Section

Materials.—The 2-ml platinum foil cups used to contain powder samples in the high-pressure syntheses analyzed 99.9+% Pt. In experiments requiring platinum in a powdered form the metal was prepared by thermal decomposition of K₂PtCl₆. Potassium chloride was removed from this product by washing it thoroughly with water. Thallium sesquioxide was prepared by the method of Rabel¹⁰ in which an alkaline thallous solution is oxidized and precipitated as virtually anhydrous Tl₂O₃ by the addition of dilute hydrogen peroxide. The sesquioxide was separated by filtration and dried at 500°. Analysis of the product gave 89.4% thallium (89.49% theoretical).

Platinum dioxide was prepared by heating K₂PtCl₄ or K₂PtCl₆ in molten KNO₃ at 350° for 24 hr. Excess KNO₃ was dissolved in water and the insoluble product treated with aqua regia to dissolve any metallic platinum and leave a residue of PtO₂. Analysis gave 86.1% platinum (85.92% theoretical).

Methods.—The reactant powders for high-pressure experiments were tamped firmly to a depth of about 5 mm in platinum foil cups 6.3 mm in height and in diameter. The sample was covered with a platinum disk, the projecting portion of the cup wall crimped to close the container, and the resulting pellet compressed in a 3-ton hydraulic press to 0.5 kbar. Several of these platinum-encased samples were then loaded into the sample cavity of a pyrophyllite tetrahedron. Details of the high-pressure sample assembly have been described previously. The tetrahedron was then exposed to the desired pressure in the high-pressure apparatus, heated to induce reaction, cooled rapidly to room temperature, and finally returned slowly to atmospheric pressure.

The reaction products were recovered from the tetrahedron and investigated by chemical analysis, X-ray diffraction, infrared, differential thermal, and thermogravimetric methods. A 114.59-mm diameter X-ray camera was used with nickel-filtered copper radiation. Calculations were based on λ 1.5417 A for Cu K α . Infrared spectra were obtained with a Beckman IR-12 spectrophotometer on thallium bromide disks containing 1% of the oxide sample, or on Nujol mulls. Simultaneous dtatga analyses were carried out on a Mettler recording thermoanalyzer.

Initial high-pressure experiments on pure Tl₂O₃ in platinum foil envelopes gave evidence of a phase transition in the sesquioxide after 1 hr at 40 kbars pressure at 550°. This phase has tentatively been indexed as a hexagonal cell isostructural with a recently reported phase of In₂O₃.¹¹ Additional information on this compound will be given in a subsequent publication. Experiments at 40 kbars and 1000° indicated that a further reaction had occurred at the higher temperature. The platinum foil envelope became brittle and partially disintegrated, while the outer surface of the sesquioxide disk was transformed from black to brown. A qualitative analysis of the brown powder revealed that it contained both thallium and platinum.

In subsequent runs powdered platinum was mixed with thallium sesquioxide to promote formation of the double oxide. The product in these experiments consisted of a brown oxide pellet mixed with unreacted excess platinum metal. Solubility tests indicated that the new oxide was insoluble in concentrated mineral acids including aqua regia. This observation permitted purification of the double oxide and suggested the presence of trivalent thallium and tetravalent platinum. Analysis confirmed this suggestion and indicated a 1:1 molar ratio for thallium and platinum. The equation for synthesis of the new oxide then becomes

$$3\text{Tl}_2\text{O}_3 + 2\text{Pt} \longrightarrow \text{Tl}_2\text{Pt}_2\text{O}_7 + 2\text{Tl}_2\text{O}$$

The thallous oxide is dissolved during the purification procedure.

The synthesis was then further simplified by mixing platinum dioxide and thallium sesquioxide in the required proportions

$$Tl_2O_3 + 2PtO_2 \longrightarrow Tl_2Pt_2O_7$$

The aqua regia purification step was retained in order to remove any traces of platinum metal from the product.

Analysis.—The double oxide has been analyzed in several ways. A sample which had been heated slowly in air to 1000° left a metallic residue comprising 42.84% of the original weight. X-Ray diffraction and spectrographic analysis identified the product as platinum. Other samples were analyzed chemically for thallium and platinum after solution in an HCl-HClO4 mixture at 275° in sealed heavy-wall glass tubes. The dissolved thallium(III) and platinum(IV) were converted to the bromides with HBr and bromine water. Thallium was then extracted with ether, reduced to Tl(I) with SO2, and precipitated as TlI. Platinum was precipitated from the ether-extracted aqueous solution with magnesium powder and weighed as the metal. In two samples the thallium content averaged 45.0% and platinum 42.4%. Another portion of the oxide was analyzed for oxygen by the KBrF4 method;12 12.2% oxygen was found. Theoretical values for Tl₂Pt₂O₇ are 44.86% thallium, 42.85% platinum, and 12.29% oxygen.

Synthesis of $Tl_2Pt_2O_7$ at Atmospheric Pressure.—Several ambient pressure methods for the synthesis of $Tl_2Pt_2O_7$ were investigated. In one experiment the direct combination of the individual oxides, PtO_2 and Tl_2O_3 , was attempted. The compacted mixed oxides gave no evidence of reaction after several days at 400° but after 4 days at 520° X-ray and infrared analyses indicated that a partial reaction had occurred. Attempts to further the reaction at somewhat higher temperatures resulted in decomposition of the PtO_2 .

Another preparative procedure involved the oxidation of the alloy, TIPt,¹³ which had been prepared by heating an equimolar mixture of the elements for 18 hr at 750° in an evacuated and sealed silica tube. Formation of the alloy was confirmed by its X-ray diffraction pattern. Oxidation experiments on powdered TIPt were run at constant temperature. After 30 days at 350° the empirical composition reached TIPtO₂, but powder X-ray diffraction data on the product showed the presence of Tl₂O₃ and a small amount of a second phase which may have been a

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platinum oxide. No Tl₂Pt₂O₇ was detectable. When the temperature was increased to 500°, the formation of the double oxide was discernible by both X-ray and infrared methods, but a pure product could not be prepared.

Results and Discussion

Thermal Stability.—Both of the constituent oxides of Tl₂Pt₂O₇ have limited thermal stability. A tga curve of PtO2 run at 6°/min indicated that trace amounts of oxygen are lost at temperatures as low as 350° but that the main decomposition reaction occurs in the temperature interval between 600 and 680°. A similar curve for type-C Tl₂O₃ showed the onset of thermal decomposition at 625°. However, no appreciable weight change was noted in a sample of Tl₂Pt₂O₇ until the furnace temperature reached 750°. Rapid decomposition began at 800° and was complete at 925° Thus the double oxide structure is appreciably more stable than either of its constituent oxides.

Structure.—We have been unable to prepare single crystals of Tl₂Pt₂O₇; hence the structural work has been confined to powder methods. Our observed X-ray pattern is indexible in terms of a small facecentered-cubic cell measuring 5.066 A. In order to account for this pattern, one could describe the structure as derived from the fluorite cell and the composition as MO_{1.75} (where M includes both metals distributed statistically throughout the cation lattice sites and one-eighth of the anion sites are left vacant). However, in view of the oxidation states and the radii of the metals involved, this possibility seems highly unlikely. The high-pressure synthesis also argues against a structure containing vacant anion sites. In addition, regardless of the reactant T1:Pt ratio, each preparation gave a 1:1 metal ratio in the final product and an identical unit cell dimension. All of these facts indicate that the constituent atoms must be located on ordered lattice sites.

The composition of this oxide, Tl₂Pt₂O₇, is proper for a cubic pyrochlore-type structure based on the space group Fd3m-Oh7. Known pyrochlore structures show cell dimensions on the order of 10 A. The observed Xray intensities are in agreement with this structure and the bond distances derived are consistent with known distances for the atoms involved. We accordingly feel justified in doubling the apparent cell to give a true cell of 10.132 ± 0.004 A and a calculated density of 11.63 g/cm³ with Z = 8. The pycnometrically measured density is 11.12 g/cm³.

Equivalent positions in the pyrochlore-type cell, as given in the "International Tables of Crystallography,"14 are $(0, 0, 0; 0, \frac{1}{2}, \frac{1}{2}; \frac{1}{2}, 0, \frac{1}{2}; \frac{1}{2}, \frac{1}{2}, 0) + 16$ Tl in (c) 1/8, 1/8, 1/8; 1/8, 3/8, 3/8; 3/8, 1/8, 3/8; 3/8, 3/8, 3/8, 1/8; 16 Pt in (d) ⁵/₈, ⁵/₈, ⁵/₈; ⁵/₈, ⁷/₈, ⁷/₈; ⁷/₈, ⁵/₈, ⁷/₈; ⁷/₈, ⁷/₈, $\frac{5}{8}$; 8 O_I in (a) 0, 0, 0; $\frac{1}{4}$, $\frac{1}{4}$, $\frac{1}{4}$; 48 O_{II} in (f) x, 0, 0; \bar{x} , 0, 0; $\frac{1}{4} + x$, $\frac{1}{4}$, $\frac{1}{4}$; $\frac{1}{4} - x$, $\frac{1}{4}$, $\frac{1}{4}$; 0, x, 0; 0, \bar{x} , 0; $\frac{1}{4}$, $\frac{1}{4}$ + x, $\frac{1}{4}$; $\frac{1}{4}$, $\frac{1}{4}$ - x, $\frac{1}{4}$; 0, 0, x; 0, 0, \bar{x} ; $\frac{1}{4}$, $\frac{1}{4}$, $\frac{1}{4} + x$; $\frac{1}{4}$, $\frac{1}{4}$, $\frac{1}{4} - x$.

Films of varying exposure times were produced in

(14) "International Tables for X-Ray Crystallography," Vol. 1, The Kynoch Press, Birmingham, England, 1952.

order to allow measurements of line intensities within reasonable density ranges. Intensities were obtained from densitometer tracings of the films and the data were then reduced in the usual manner. However, it was necessary to include an absorption correction because of the very large μ value of 2245 cm⁻¹ for this compound. For the particular capillary used, and with an estimate of the packing in the capillary based on a measurement of the grain sizes, a value of $\mu r = 13.5$ was obtained. This value served as the basis for the absorption correction. The oxygen contribution to the scattering is dominated by the heavy cation scattering, and this, with the high absorption, places a severe limitation on the accuracy of the coordinate determination for the oxygen atoms.

An approximate value of x was found by trial to satisfy intensity requirements. A least-squares refinement, based on "A Fortran Crystallographic Least Squares Program," by Busing, Martin, and Levy, led to a final value of $x = 0.270 \pm 0.006$ for the 48fold O_{II} position. The structure factors were computed with atomic scattering factors for the metals given in Vol. III of the "International Tables for Crystallography" and the O2- scattering curve reported by Tokonami. 15 The calculated structure factors were also corrected for anomalous dispersion using values given by Cromer. 16

The R factor, omitting zeros, is 3.6% based on R = $\Sigma\{|F_{\rm o}| - |F_{\rm c}|\}/\Sigma|F_{\rm o}|$. The low R factor does not measure the accuracy of the x-coordinate determination because of the dominance of the cation scattering. This is indicated by the high error in x.

The observed and computed intensities are presented in Table I. A reflection hkl indicated by an asterisk calculates zero intensity because of the point positions for this structure. A dashed line in the $\sin^2 \theta_0$ column indicates that the line was not observed on the film. Numerical values of I_0 given in the table for these unobserved reflections are limiting intensities on the densitometer tracings corresponding roughly to the noise level of the tracing background. Clearly, one may observe lines visually on the film of intensities substantially lower than those detectable by the densitometer. We have indicated that for our case no additional lines were observed visually on the film after prolonged exposures. The final calculated intensities have been corrected for the effects of thermal motion using a value of 1.80 A² for B. Absolute values for I_0 are obtained by multiplying the calculated quantities by 107.

In the proposed structure, each platinum atom is bonded to six oxygen atoms, with Pt-6(O_{II}) = $2.08 \pm$ 0.04 A. This value is in good agreement with the Pauling¹⁷ and Ahrens¹⁸ radii sum of 2.05 A (0.65 + 1.40 A for ionic bonding or 1.31 + 0.74 A for covalent bonding). Each thallium atom is bonded to eight

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Table I

X-Ray Powder Diffraction Data for Tl₂Pt₂O₇

	hkr.	sin200	sin20c	10	1 _c	hks	sin ² 0	sin ² ec	I _o	I e
	111		0.01761	< 5.0	0.24	∫ 951a ₁		.6192	< 0.5	.02
	200*		.02375	< 4.0	0	[773α ₁	**	.6192	< 0.13	.02
	220		.04691	< 2.0	0.11	(10,2,2a1	.6254	.6255		
	311		.06446	< 2.0	< 0.01	10,2,242	.6287	.6287	100	
	222	0.07034	.07031	87.6	84.4	66601	.6254	.6255	6.9	6.9
	400	.09365	.09365	28.4	29.8	666a2	.6287	.6287		
	331		,1112	< 1.0	.50	953α1	**	.6657	€ 0.2	e .01
	420*		,1170	< 1.0	0	(10,4,0%,	**	.6715		
	422		.1403	< 1.0	.05	8642		.6715	< 0.4	0
1	511		.1577			10,4,20,		.6946	< 0.2	< .01
	333		.1577	< 2.0	.12	(11,1,10)		.7119		
	440	.1862	,1867	31.9	32,6	77501		.7119	< 0.2	< .01
	531	.1002	,2041	< 0.7	.07	∫ 880a ₁	.7406	.7406		
-	600*		,2099		,	88002	.7442	.7444	3,5	2,6
1	442*		,2099	< 1.1	0	f11,3,1a,	.,,	.7579		
			.2330	0.4		97101		.7579	< 0.5	.04
	620			€ 0.6	< .01	95501		.7579	< 0.5	.04
	533	-	.2504	< 0.6	< .01					
	622	.2563	.2562	21.8	28.2	\$10,4,4å ₁		.7637	< 0.4	0
	444	.2796	.2793	7.1	6.5	88201	55	.7637		
	711		.2964	< 1.0	.05	\$10,6,0a1	**	.7868	< 0.3	€ .01
L i	551	**	.2964	-		86601		.7868		
	640#		.3024	< 0.5	0	$\int 11, 3, 3a_1$.8040	< 0.3	.01
	642		.3255	< 0.4	< .01	973a ₁		.8040		
1	731		.3428	< 0.9	.02	∫10,6,2a ₁	.8099	.8098	12.4	9.4
1	553	- 48	.3428	2 0.7	.02	10,6,242	.8145	.8139	-	7.15
	800	.3718	.3717	3.8	3.8	[12,0,0a]	.8333	.8328		
	733		.3880	< 0.4	0.6	12,0,000	.8373	.8371		-
1	820≘	**	.3947			88441	.8333	.8328	7.9	5.4
	644#		.3947	< 0.7	0	884#2	.8373	.8371		
٢	822	-	.4179			(11,5,1a,		.8500		
-	660		.4179	< 0.7	< .01	77747		.8500	< 0.3	.02
-	751		.4351			12,2,00,		.8558	< 0.2	0
	555		.4351	< 0.7	.01	[12,2,2a,		.8789		
	662	.4410	.4409	8.9	9.1	10,6,40,		.8789	< 0.3	.01
	840	.4640	.4640	7.3		(11,5,3a)	**	.8961		
				7.3	7.1	97501		.8961	< 0.3	.01
	911	77	.4813	€ 0.6	.04	(12,4,0a,	.9251	,9250		
	753	500	.4813			1			7.1	7.2
	842*		,4870	< 0.3	0	[12,4,0a ₂	.9296	.9296		
	664		,5101	< 0.3	< .01	991#1	**	.9422	< 0.1	01
	931		.5274	< 0.3	.01	[12,4,2a1	**	.9480		
	844	.5564	.5562	6.2	6.8	10,8,0%		.9480	< 0.4	0
	$933a_{1}$.5736			88621		.9480		
	$771\alpha_1$.5736	- 0.8	.03	10,8,201		.9706	< 0.1	< .01
	755a1		.5736			[13,1,1a ₁	**	,9882		
10	,0,0*		.5793	-0.5	0	11,7,101		.9882	€ 0.6	.06
1	860≎		.5793	< 0.5	U	11,5,501		.9882	50.0	.06
1	0,2,0		.6024	- 0.5		99341	**	.9882		
	362		.6024	< 0.5	< .01	10,6,64	.9938	.9940	8.4	13.5

oxygens with Tl–2(O_I) = 2.19 ± 0.002 A and Tl–6-(O_{II}) = 2.32 ± 0.04 A. The weighted average bond length is 2.29 A for the eight bonds, as compared with the Pauling radii sum (0.95 + 1.40 A) of 2.35 A. Three oxygen–oxygen distances are found: O_I–O_{II} = 2.74 ± 0.06 A, O_{II}–O_{II} = 2.55 ± 0.01 A, and O_{II}–O_{II} = 3.30 ± 0.09 A.

Figure 1 illustrates a comparison of the Tl₂Pt₂O₇ cell dimension with other known pyrochlore-type compounds, including the rare earth titanates, ¹⁹⁻²¹ ruthenates, ²² stannates, ²³ and zirconates, ¹⁹ together with Cd₂Ta₂O₇ and Cd₂Nb₂O₇, ^{24,25} The Tl₂Pt₂O₇ parameter is seen to fall very close to the line connecting the ruthenate(IV) pyrochlores. The agreement in cell dimension is excellent, since Ru(IV) has been assigned a radius of 0.63 A by Pauling ¹⁷ and 0.67 A by Ahrens. ¹⁸

Pyrochlore Structure.—Figure 2 illustrates a portion of a pyrochlore-type unit cell—one of the eight units which constitute the true cell of this space group.

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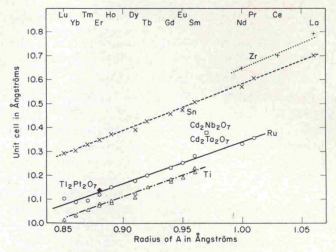


Figure 1.—Cell dimensious of A₂B₂O₇ compounds.

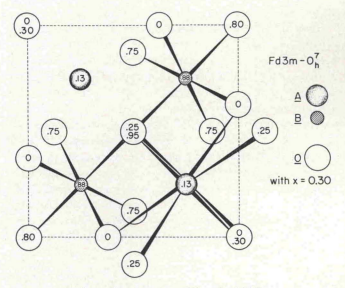


Figure 2.—Atomic positions in A₂B₂O₇.

The distorted octahedral arrangement of oxygen atoms about each B atom and the eightfold configuration about each A atom to form a distorted cube can be seen. Since each oxygen atom in the three-dimensional array of BO₆ octahedra is shared between two B metal atoms, the over-all composition becomes $(B_2O_6)_n$ for the octahedral lattice. The space group requires that all B–O distances are equivalent, but not that the octahedra are symmetrical. It also requires that six of the eight A–O bonds be equivalent; the other two can be, but usually are not, equal in length to the six.

Since the A, B, and O_I atom sites are invariant in the unit cell and the O_{II} sites contain only a single variable, one can plot metal-oxygen and oxygen-oxygen distances (in terms of fraction of the unit cell parameter) as a function of x. As shown by Figure 3, the $A-2(O_I)$ distances are independent of x, but the $A-6(O_{II})$ distances do vary with x; the eight bonds become equal at x = 0.25. The A-8(O) line represents the weighted average bond length between these atoms. Distances between atoms in the octahedra are given by the $B-6(O_{II})$ line and the two $O_{II}-O_{II}$ lines. The oxygenoxygen distances are equivalent and the octahedra are



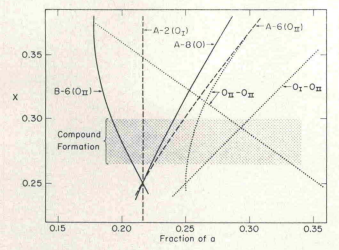


Figure 3.—Atomic distances in pyrochlore-type structures Fd3m-Oh7.

regular at x = 0.3125. The O_I-O_{II} line represents the distance between an O_I atom and the nearest O_{II} atom in an adjacent octahedron.

The shaded area indicates the observed range of x in those pyrochlores whose oxygen positions have been located. The region of compound formation appears to lie approximately midway between x = 0.3125, where regular octahedra obtain, and x = 0.2500, where perfect cubes are formed.

The minimum cell size which has been reported among the pyrochlore-forming compounds, 9.801 A in Sc₂Ti₂O₇, 20 must represent a virtual lower limit for the structure because of the volume requirements imposed by the 56 oxygen atoms in each unit cell.

Infrared Data.—The infrared spectrum of Tl₂Pt₂O₇ is somewhat unusual when compared with other pyrochlore spectra. Figure 4 illustrates the spectra of two typical examples, Yb₂Ru₂O₇ and Y₂Sn₂O₇, which were prepared for comparison purposes. A representative titanate pyrochlore spectrum has been given by Knop, et al.21 Figure 5 shows the absorption spectrum of Tl₂-Pt₂O₇, characterized by four sharp absorption bands at 684, 562, 449, and 363 cm⁻¹. It is apparent that the general features of the spectra are similar, but that the absorption maxima in the platinum compound are much sharper. This effect may be due to increased covalent character of the bonding in Tl₂Pt₂O₇. Although the ruthenium electronegativity¹⁷ is comparable with platinum (2.2), thallium (1.8) is appreciably more electronegative than the rare earths (1.1–1.2).

The strong broad absorption bands of the erbium titanate spectrum have tentatively been assigned²¹ to Ti-O modes; the higher frequency maximum has been assigned to a Ti-O stretching vibration and the lower one, to a bending mode. It has also been suggested that

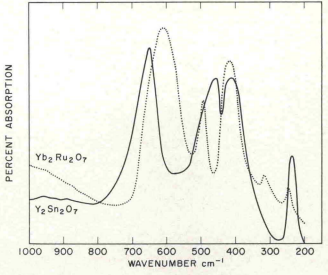


Figure 4.—Infrared spectra of pyrochlores.

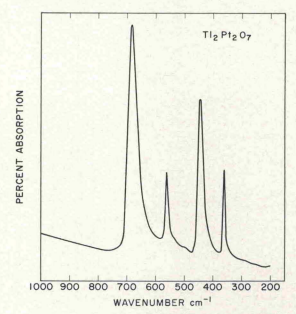


Figure 5.—Infrared spectrum of thallium platinate.

the minor features (shoulders and subsidiary maxima) are associated with the oxygen environment of the erbium atoms. A similar interpretation for the Tl₂Pt₂O₇ spectrum would assign the 684-cm⁻¹ band to a Pt-O stretching mode, the 449-cm⁻¹ band to an O-Pt-O bending mode, and the two weaker bands at 562 and 363 cm⁻¹ to Tl-O_I stretching and bending modes. Further work is required to establish the validity of these tentative assignments.

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