

SPECTROSCOPIC AND THERMAL STUDIES ON PLATINUM(II) IODIDE COMPLEXES WITH PRIMARY AMINES

G. FARAGLIA and L. SINDELLARI

*Dipartimento di Chimica Inorganica, Metallorganica ed Analitica dell' Università,
via Loredan 4, 35100-Padova (Italy)*

S. SITRAN

Istituto di Chimica e Tecnologia dei Radioelementi del C.N.R., Padova (Italy)

(Received 13 October 1986)

ABSTRACT

The complexes *cis*- and *trans*-[PtL₂I₂], [PtBua₃I]I and [PtL₄]I₂, where L = butan-1-amine (Bua) or hexan-1-amine (Hea), have been prepared and characterized by elemental analyses, IR and ¹H NMR spectra and TG, DTG, DTA measurements. Thermal degradation of the 1:3 and 1:4 species yields the corresponding *trans* complexes as intermediates. Moreover, the *cis* species isomerize to *trans* without decomposition. In the ¹H NMR spectra of the 1:2 and 1:4 derivatives, the NH₂ proton signal is characteristic of the stoichiometry and geometry, whereas [PtBua₃I]I decomposes in most solvents, releasing one of the ligand molecules to give *trans*-[PtBua₂I₂] solutions.

INTRODUCTION

In the past few years we have reported on various platinum(II) halide complexes with hexan-1-amine (Hea) and propan-1-amine (Pra) having the formulae *cis*- and *trans*-[PtL₂X₂], [PtL₃X]X and [PtL₄]X₂ (X = Cl or Br), which were characterized either in the solid state (IR and thermal analysis) or in solution (¹H NMR spectra) [1,2]. We extended the study to the iodo-derivatives [3], due to their use as intermediates in the preparation of drugs for anti-tumour tests [4,5]. When aquo-ion solutions are used in biological tests particular attention must be paid to the purity of the starting platinum iodide amino complexes. In this line, a recent paper reports the preparation of pure *cis*-[Pt(NH₃)₂I₂] through formation of solvates with formamides and its behaviour in aqueous KI and HI solutions [6].

In this paper we report the preparation of the PtI₂ complexes with hexan-1-amine and butan-1-amine (Bua) and their characterization by IR, ¹H NMR and thermal analysis data.

EXPERIMENTAL

The reagents used were PtI_2 (Johnson Matthey), $\text{K}_2[\text{PtCl}_4]$ (Fluka), hexan-1-amine (C. Erba) and butan-1-amine (Janssen). The reactions were generally carried out at room temperature.

*Preparation of the complexes**trans-[PtHea₂I₂]*

The complex was prepared by reacting PtI_2 (0.7 mmol) with Hea (1.6 mmol) in benzene (5 cm³) with stirring (6 h). The resultant yellow solid was filtered, washed with *n*-pentane and dried in vacuo. Further fractions of product were obtained by partial evaporation of the residual benzene solution and subsequent addition of *n*-pentane (yield, 80%). The compound can be recrystallized from benzene/*n*-pentane. The melting point range was 110–112°C and the experimental composition was: C, 22.2; H, 4.7; N, 4.3% (calculated (for $\text{C}_{12}\text{H}_{30}\text{I}_2\text{N}_2\text{Pt}$): C, 22.1; H, 4.7; N, 4.3%). It was also prepared in a nearly quantitative yield by stirring (4 h) a PtI_2 (0.5 mmol) suspension in a Et_2O solution of Hea (1.1 mmol in 5 cm³). The reaction went on in heterogeneous phase, yielding the slightly soluble *trans* isomer. Pure samples were also obtained by evaporation to dryness of carefully filtered acetone solutions of the crude product and by heating $[\text{PtHea}_4]\text{I}_2$ under reduced pressure (ca. 80°C).

cis-[PtHea₂I₂]

Aqueous KI (3.7 mmol in 2.0 cm³ of H_2O) was added to an aqueous solution of $\text{K}_2[\text{PtCl}_4]$ (0.9 mmol in 5 cm³). The dark red solution was carefully filtered from a yellow undissolved residue. Addition of Hea (1.8 mmol) with vigorous stirring yielded a pinkish solid, which was filtered, washed with H_2O and dried under reduced pressure. The solid, washed with Et_2O , was recrystallized from benzene/*n*-hexane and finally washed with Et_2O (yield, 60%, m.p., 102–103°C; found: C, 22.1; H, 4.7; N, 4.3%).

[PtHea₄]\text{I}_2

PtI_2 (0.5 mmol) was allowed to dissolve in a Et_2O solution of Hea (2.2 mmol in 5 cm³; 4 h). Slow evaporation of the filtered yellow solution yielded white crystals of the compound, which were washed with *n*-pentane. The compound can be purified by evaporation of solutions in *n*-hexane, in which the complex is slightly soluble (yield, 90%; m.p., 84–85°C; C, 33.7; H, 7.2; N, 6.6%; calculated (for $\text{C}_{24}\text{H}_{60}\text{I}_2\text{N}_4\text{Pt}$): C, 33.8; H, 7.1; N, 6.6%).

trans-[PtBua₂I₂]

The yellow solution obtained by reacting PtI_2 (0.4 mmol) and Bua (0.8 mmol) in benzene (3 cm³) was filtered from the small undissolved residue (a

mixture of *cis* and 1:3 species from IR spectra). The first solid fraction obtained after addition of *n*-hexane was discarded (*cis-trans* mixture). Further addition of *n*-hexane gave the yellow compound. It can be recrystallized from benzene/*n*-hexane (yield, 70%; m.p., 140–142°C; C, 16.2; H, 3.7; N, 4.7%; calculated (for $C_8H_{22}I_2N_2Pt$): C, 16.1; H, 3.7; N, 4.7%). The compound can be easily obtained by thermal decomposition of the corresponding 1:3 and 1:4 complexes under reduced pressure (ca. 130°C).

cis-[PtBua₂I₂]

Prepared by the method used for *cis*-[PtHea₂I₂]. The yellow solid was recrystallized from benzene (yield, 60%; m.p., 140–142°C; C, 16.1; H, 3.7; N, 4.7%).

[PtBua₃II]

The reaction of PtI₂ (0.4 mmol) with Bua (1.3 mmol) in Et₂O yielded (6 h, with stirring) a white solid and a yellow solution. The solid, filtered and washed with Et₂O, was [PtBua₄]I₂ impure for the 1:3 complex (by IR spectra). The residual Et₂O solution, kept at ca. –20°C, yielded greenish-white crystals of the compound, which were filtered, washed with small fractions of cold Et₂O (in order to eliminate traces of the *trans* species) and finally with *n*-pentane (yield, 50%; C, 21.7; H, 5.1; N, 6.3%; calculated (for $C_{12}H_{33}I_2N_3Pt$): C, 21.6; H, 5.0; N, 6.3%).

[PtBua₄]I₂

A suspension of PtI₂ (0.6 mmol) in a Et₂O solution of Bua (2.9 mmol in 5 cm³) yielded, overnight with stirring, a white solid, which was filtered and washed with Et₂O and *n*-pentane. It was recrystallized from benzene/*n*-pentane (yield, 80%; m.p., 127–129°C; C, 26.0; H, 6.1; N, 7.5%; calculated (for $C_{16}H_{44}I_2N_4Pt$): C, 25.9; H, 6.0; N, 7.6%).

Measurements

The IR spectra were recorded by using either a Perkin Elmer 580B spectrophotometer (4000–400 cm^{–1}) or a Bruker FT IR instrument (450–100 cm^{–1}) as Nujol mulls between KBr and polyethylene discs. The ¹H NMR spectra were obtained using a Jeol FX 90 Q spectrometer. The TG, DTG, and DTA curves in dinitrogen (flux rate, 250 cm³ min^{–1}; heating rate, 5°C min^{–1}) were recorded on a Netzsch STA 429 thermoanalytical instrument (reference material, neutral Al₂O₃).

RESULTS AND DISCUSSION

The compounds (Table 1) were generally prepared in organic media by reaction of platinum iodide and amine in the appropriate molar ratio, except

TABLE 1

Solubilities^a and infrared frequencies in the 3500–3000 cm⁻¹ and 1650–1500 cm⁻¹ regions

Compound	Solubility	Frequency (cm ⁻¹)					
		MeOH	C ₆ H ₆	Me ₂ CO	Et ₂ O	CH ₂ Cl ₂	$\delta(\text{NH}_2)$
<i>trans</i> -[PtHea ₂ I ₂]	~i		vs	vs	sls	vs	1578(s)
<i>cis</i> -[PtHea ₂ I ₂]	s		vs	vs	~i	vs	1565(s)
[PtHea ₄] ₂	sls		s	s	s	s	1608(wbr), 1590(sh)
<i>trans</i> -[PtBua ₂ I ₂]	s		vs	vs	s	vs	1578(s)
<i>cis</i> -[PtBua ₂ I ₂]	sls		sls	vs	i	s	1565(s)
[PtBua ₃] ₂	s		s	vs	sls	vs	1574(s)
[PtBua ₄] ₂	vs		s	vs	i	vs	1610(wbr), 1590(sh)

^a At room temperature: i, insoluble; s, soluble, sls, slightly soluble; vs, very soluble. The complexes are insoluble in H₂O (except for [PtBua₄]₂, sls) and in *n*-hexane (except for [PtHea₄]₂, sls). All the complexes are soluble in DMSO and in DMF.

for *cis* derivatives, which were prepared by reaction of $[\text{PtI}_4]^{2-}$ and amine in water. The *trans* isomers were also obtained by thermal decomposition of the higher stoichiometry derivatives and by thermal isomerization of the parent *cis* species. Owing to their solubility in organic solvents (Table 1) the complexes can be easily purified, except for $[\text{PtBua}_3\text{I}]\text{I}$, which generally releases one ligand molecule, yielding yellow solutions of *trans*- $[\text{PtBua}_2\text{I}_2]$. Since the process is slow in diethylether, the compound has been prepared in this solvent by reaction of platinum iodide and Bua (molar ratio 1 : 3.1) and stored at about 5°C. In analogous conditions, the 1 : 3 Hea derivative was isolated in a low yield and impure form for $[\text{PtHea}_4]\text{I}_2$. Attempts to recrystallize $[\text{PtHea}_3\text{I}]\text{I}$ were unsuccessful, owing to its easy decomposition and to its solubility, very close to that of the parent 1 : 4 complex.

The trend of the infrared absorptions in the $\nu(\text{NH})$ and $\delta(\text{NH}_2)$ regions (Table 1) is strictly related to stoichiometry and geometry and, apart for small halide-dependent shifts, it is very close to that of the chloro- and bromo-derivatives [1]. The *trans* species show, beyond 3000 cm^{-1} , three sharp absorptions, whereas the *cis* species and $[\text{PtBua}_3\text{I}]\text{I}$ present a strong, broad band at 3200 cm^{-1} and 3180 cm^{-1} , respectively. Owing to the presence of strong hydrogen bonds with iodides, the 1 : 4 complex absorptions are at lower energy (ca. 3100 cm^{-1}), the corresponding bending mode bands being at 1610 cm^{-1} . The halide-dependent $\delta(\text{NH}_2)$ absorptions shift in the *trans*- $[\text{PtHea}_2\text{X}_2]$ series from 1587 cm^{-1} ($\text{X} = \text{Cl}$) to 1582 cm^{-1} ($\text{X} = \text{Br}$) and 1578 cm^{-1} ($\text{X} = \text{I}$) and in the analogous *cis* series from 1578 cm^{-1} ($\text{X} = \text{Cl}$) to 1570 cm^{-1} ($\text{X} = \text{Br}$) and 1565 cm^{-1} ($\text{X} = \text{I}$). A similar behaviour was observed for the parent platinum halide complexes with Pra. The platinum-iodine stretching frequencies (Table 2) are as expected. The *cis* isomers present a strong band at ca. 177 cm^{-1} , shifted to higher frequencies in the *trans* isomers (191 cm^{-1}). The compound $[\text{PtBua}_3\text{I}]\text{I}$ exhibits, together with $\nu(\text{Pt}-\text{I})$ at 188 cm^{-1} , a strong absorption at 278 cm^{-1} , common to the 1 : 4 species (289 cm^{-1}), probably to be assigned to Pt-N stretching [7-10].

The thermal analysis data are summarized in Table 3. The *trans* species melt without decomposition (Hea, 115°C ; Bua, 143°C) and their degradation starts at ca. 190°C . The thermograms of *cis*- $[\text{PtBua}_2\text{I}_2]$ (Fig. 1) clearly show the isomerization exotherm at 122°C , followed by the melting endotherm of the *trans* species, as observed for *cis*- $[\text{PtPra}_2\text{I}_2]$ (isomerization exotherm 125°C ; melting exotherm, 203°C). The isomerization process starts in *cis*- $[\text{PtHea}_2\text{I}_2]$ (Fig. 2) at 90°C , then melting of the *cis-trans* mixture takes place (100°C) followed by further isomerization in the melt (107°C). At higher temperatures the thermograms coincide with those of the *trans* derivatives, as for the 1 : 4 complexes after releasing of two ligand molecules. The thermal behaviour of $[\text{PtBua}_3\text{I}]\text{I}$ differs from that of $[\text{PtPra}_3\text{I}]\text{I}$, which releases one ligand molecule in a single step (109°C) to give *trans*- $[\text{PtPra}_2\text{I}_2]$. As is shown in Fig. 3, the release of the first ligand

TABLE 2

Infrared frequencies (450–100 cm⁻¹; ν (Pt–I) are underlined) and ¹H NMR data in CDCl₃ (ppm; T, ca. 25 °C)

Compound	Infrared frequency (cm ⁻¹)	NMR chemical shift (ppm)		
		NH ^{a,b}	α CH ₂ ^b	β CH ₂
<i>trans</i> -[PtHea ₂ I ₂]	403(vw), 339(w), 267(mw), 224(vw), <u>191(s)</u> , 88(w)	3.25	2.9	1.4 ^c
<i>cis</i> -[PtHea ₂ I ₂]	401(vw), 327(vw), 254(vw), 178(m)	4.35	2.9	1.7
[PtHea ₄]I ₂	405(vw), 342(vw), 289(sbr), 225(wbr), <u>135(wbr)</u>	5.65	2.7	1.85
<i>trans</i> -[PtBua ₂ I ₂]	346(w), 265(mw), <u>191(s)</u> , 89(w)	3.25	2.9	1.4 ^c
<i>cis</i> -[PtBua ₂ I ₂]	331(vw), 253(vw), <u>176(m)</u>	4.35	2.9	1.65
[PtBua ₃]I	443(vw), 433(vw), 278(sbr), 228(vw), <u>188(m)</u>	5.6w–3.2 ^d	2.7–2.9	
[PtBua ₄]I ₂	455(wbr), 430(vw), 347(w), 289(sbr), 180(wv)	5.65	2.7	1.80

^a Coupling with ¹⁹⁵Pt (J ca. 60 Hz) is observed.^b Broad signals.^c Superimposed on the signals of the other CH₂ groups.^d The solution, initially colourless, turns to yellow in 5 min, owing to formation of the *trans* species.

TABLE 3

Thermal data of the complexes (in dinitrogen)

Compound	Decomposition interval (°C)	TG weight loss (%)		DTA peak temperature (°C) ^a
		Experimental	Calculated	
<i>trans</i> -[PtHea ₂ I ₂]	195–370	68.6	70.0 (2Hea + 2I)	115(m), 220(ex), 345(d)
<i>cis</i> -[PtHea ₂ I ₂]	185–370	68.7	70.0(2Hea + 2I)	95(iso), 100(m), 107(iso), 222(ex), 345(d)
[PtHea ₄]I ₂	80–190	23.6	23.7(2Hea)	95(m), 137(d)
	190–370	52.4	53.4(2Hea + 2I)	225(ex), 345(d)
<i>trans</i> -[PtBua ₂ I ₂]	190–370	66.8	67.2(2Bua + 2I)	143(m), 216(ex), 345(d)
<i>cis</i> -[PtBua ₂ I ₂]	190–370	66.4	67.2(2Bua + 2I)	122(iso), 143(m), 216(ex), 345(d)
[PtBua ₃]I	55–140	11.0	10.9(1Bua)	90(d), 135(shd), 143(m)
	180–370	59.2	59.9(2Bua + 2I)	216(ex), 345(d)
[PtBua ₄]I ₂	95–145	19.7	19.7(2Bua)	134(d), 143(m)
	175–370	52.8	54.0(2Bua + 2I)	216(ex), 345(d)

^a ex, exotherm; m, melting endotherm; d, decomposition endotherm; iso, isomerization exotherm.

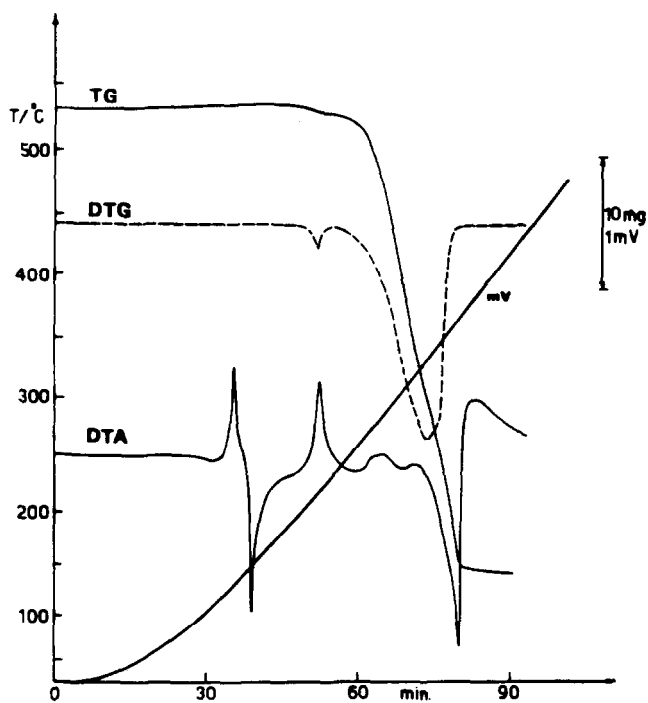


Fig. 1. Thermograms of *cis*-[PtBua₂I₂] (56.42 mg).

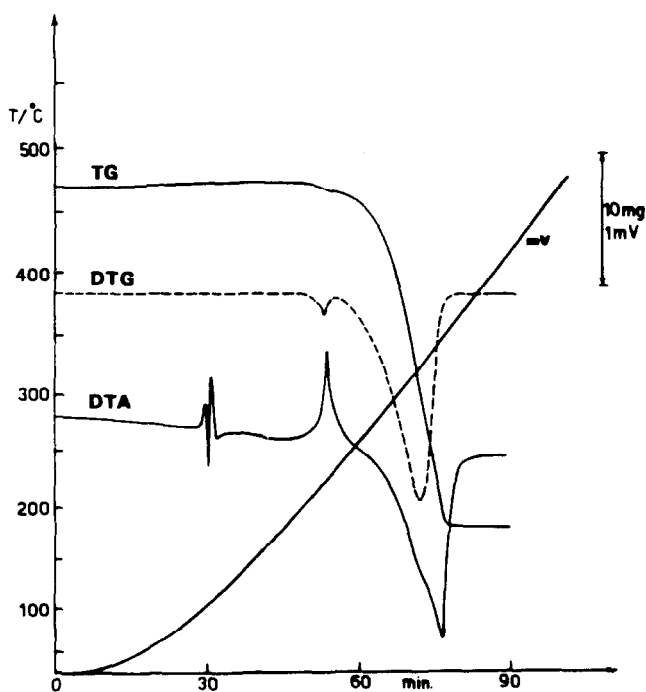


Fig. 2. Thermograms of *cis*-[PtHea₂I₂] (39.04 mg).

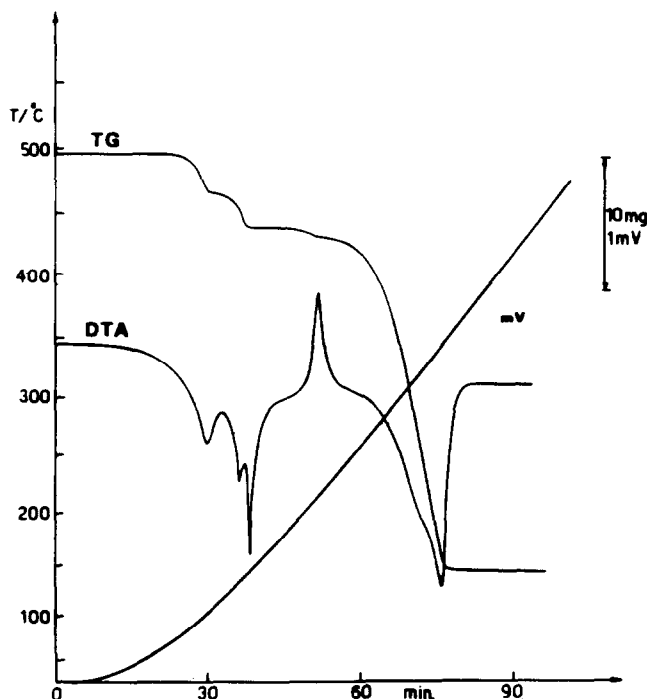


Fig. 3. Thermograms of $[\text{PtBua}_3\text{I}]\text{I}$ (45.54 mg).

molecule takes place in two steps, the related DTA peaks being at 90°C and 135°C . IR and ^1H NMR spectra of $[\text{PtBua}_3\text{I}]\text{I}$ samples heated up to 90°C are in accordance with the formation of *trans*- $[\text{PtBua}_2\text{I}_2]$ and, in part, of $[\text{PtBua}_4\text{I}_2]$. Both species are absent in the starting product, whose IR spectrum shows an absorption minimum below 3100 cm^{-1} , where the 1:4 complex absorbs strongly. The relative intensity of the steps depends on the amount of substance in the crucible. If a few milligrams are used, the second step is minimized.

The ^1H NMR data in deuterated chloroform are reported in Table 2. Whereas the αCH_2 and βCH_2 resonances are quite close for all complexes, the NH_2 proton signal is affected by stoichiometry and geometry (Fig. 4). For the *trans* species it is observed upfield (3.25 ppm) with respect to the corresponding signal in the *cis* isomer (4.35 ppm), allowing the detection of *cis* to *trans* isomerization processes in solution. The NH_2 resonances of the 1:4 complexes are downfield with respect to those of the 1:2 complexes, as observed previously for analogous halides. The ^1H NMR spectrum of a solution of $[\text{PtBua}_3\text{I}]\text{I}$ is initially similar to those of the chloro- and bromo-homologues and presents two NH_2 signals, due to the different environment of the ligand molecules in the square planar $[\text{PtBua}_3\text{I}]^+$ moiety. In a short time the solution turns from colourless to yellow and the downfield signal

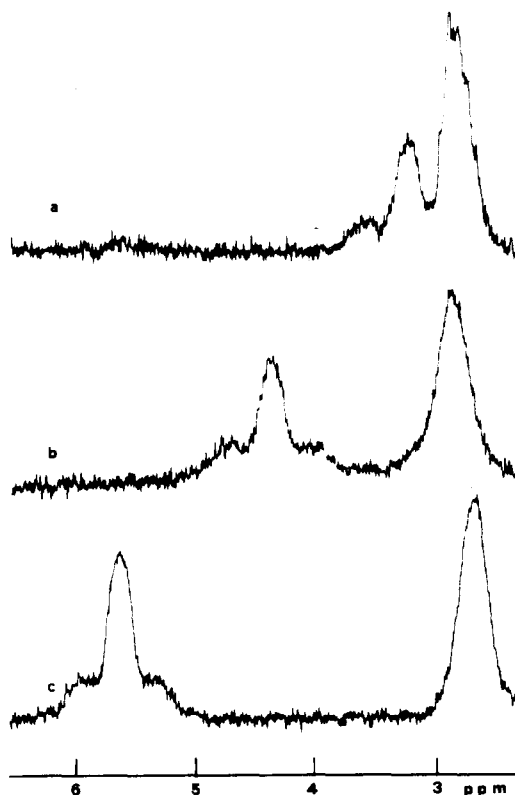


Fig. 4. ^1H NMR spectra in CDCl_3 : a, *trans*- $[\text{PtHea}_2\text{I}_2]$; b, *cis*- $[\text{PtHea}_2\text{I}_2]$; c, $[\text{PtHea}_4]\text{I}_2$.

decreases progressively, the final spectrum coinciding with that of *trans*- $[\text{PtBua}_2\text{I}_2]$. In conclusion, either the *cis* species or $[\text{PtBua}_3\text{I}]\text{I}$ present a low stability in solution. Among the complex series prepared, the bromo-derivatives seem to be the more useful intermediates in the preparation of aquo-ion solutions and compounds of biochemical interest, owing to the easy purification and the solubility and stability characteristic.

ACKNOWLEDGEMENT

The authors thank Mrs Franca Marzola for far-infrared spectra registration.

REFERENCES

- 1 G. Faraglia, L. Sindellari and S. Sitran, *Thermochim. Acta*, 78 (1984) 159.
- 2 V. Cherchi, G. Faraglia, L. Sindellari and S. Sitran, *Transition Met. Chem.*, 10 (1985) 76.

- 3 G. Faraglia, L. Sindellari, V. Cherchi and S. Sitran, *Transition Met. Chem.*, 11 (1986) 98.
- 4 P.C. Hydes and B.W. Malerbi, U.S. 4,225,529 (Cl.260-49R; C07F15/00; 1980); *Chem. Abstr.*, 94 (1981) 77033m.
- 5 M.L. Tobe, A.R. Khokhar and P.D.M. Braddock, *Ger. Offen.*, 2,715,492 (Cl.A61K31/28; 1977); *Chem. Abstr.*, 88 (1978) 99314g.
- 6 G. Raudaschl, B. Lippert, J.D. Hoeschele, H.E. Howard-Lock, C.J.L. Lock and P. Pilon, *Inorg. Chim. Acta*, 106 (1985) 141.
- 7 C. Engelter, A.T. Hutton and D.A. Thornton, *J. Mol. Struct.*, 44 (1978) 23.
- 8 M. Pfeffer, P. Braunstein and J. Dehand, *Spectrochim. Acta, Part A*, 30 (1974) 341.
- 9 M. Pfeffer, P. Braunstein and J. Dehand, *Inorg. Nucl. Chem. Lett.*, 8 (1972) 497.
- 10 G.W. Watt, L.K. Thompson and A.J. Pappas, *Inorg. Chem.*, 11 (1972) 747.