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PLATINUM GROUP METAL SCHIFF BASE COMPLEXES— I. PLATINUM COMPLEXES

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Abstract—Schiff base ligands (L), prepared from benzaldehyde, salicylaldehyde and 2- or 3-aminopyridines or 3-aminoquinoline, have been isolated and reacted with K_2PtCl_4 , *cis*-[Pt(DMSO)₂Cl₂], [Pt(COD)Cl₂] and [PtL'Cl₂]₂ (L' = C₂H₄, PEt₃, PPh₃ PMe₂Ph) in different molar ratios, to yield complexes of general formula [PtLCl₂]_n (n = 1 or 2) and *cis*- or *trans*-[PtLL'Cl₂]. The Schiff bases and their platinum complexes have been characterized physicochemically.

Platinum complexes containing nitrogenous ligands have received much attention, especially since Rosenberg's discovery¹ that cis-[Pt(NH₃)₂Cl₂] (cisplatin) is an effective antineoplastic agent (first generation) for treating tumours. Thus, cis- $[Pt(NH_3), \{(OCO), C--CH_2CH_2CH_3\}]$ (carboplatin) and cis-[Pt(PrⁱNH₂)₂(OH)₂Cl₂] (iproplatin) were prepared (second generation).^{2,3} In view of the importance of such complexes and our intrinsic interest in studying the interaction between platinum and various nitrogen ligands, and the biological activity of such complexes,⁴⁻⁷ we now describe the reaction of a wide range of Schiff bases (L) (rarely used as ligands to platinum^{8, 9}) derived from benzaldehyde or salicylaldehyde and 2- or 3-aminopyridines or 3-aminoquinoline (Scheme 1) and K_2PtCl_4 , cis-[Pt(DMSO)₂Cl₂], [Pt(COD)Cl₂] (COD = 1.5-cyclopentadiene) and $[PtL'Cl_2]_{\gamma}$ $(L' = C_2H_4, PEt_3, PPh_3, PMe_2Ph)$. To the best of our knowledge this work is novel.

EXPERIMENTAL

General

¹H NMR spectra were recorded in $(CD_3)_2CO$ at 25°C on a Jeol Datum no. Ex-1 90 MHz instrument.

IR spectra were recorded on a Unicam SP 2000 spectrophotometer in the 200-4000 cm⁻¹ range using Nujol mulls and CsI discs.

UV-vis spectra were recorded on a Shimadzu UV-vis spectrophotometer UV-160 for 10^{-3} M solutions of the compounds in DMF at 25°C, using a 1 cm quartz cell.

Elemental analyses were carried out on a CHN Analyser, Type 1106 (Carlo Erba) at the Department of Chemistry, College of Science, University of Mosul, Mosul, Iraq.

Conductivity measurements were made on 10^{-3} M solutions of the complexes at 25°C using a conductivity meter model 4070 (Jenway).

Starting materials

$$\begin{split} &K_2 PtCl_4 \quad and \quad PtCl_2 \quad were \quad used \quad as \quad supplied \\ by \quad Fluka. \quad cis-[Pt(DMSO)_2Cl_2], \quad [Pt(COD)Cl_2] \\ (COD = 1,5-cyclooctadiene), \qquad [Pt(C_2H_4)Cl_2]_2, \\ [Pt(PEt_3)Cl_2]_2, \quad [Pt(PPh_3)Cl_2]_2 \quad and \quad [Pt(PMe_2Ph) \\ Cl_2]_2 \quad were \quad prepared \quad according \quad to \quad literature \\ methods.^{10,11} \end{split}$$

The Schiff bases (Scheme 1) were prepared by using the following standard method.¹²

The aldehyde derivative (benzaldehyde or salicylaldehyde) (0.01 mol) in MeOH (15 cm³) was mixed with 2- or 3-aminopyridines or 3-aminoquinoline (0.01 mol) in MeOH (15 cm³) in 100 cm³ round bottomed flask, and the mixture was boiled

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Scheme 1. The Schiff bases used in the complexation with platinum.

under reflux for ca 1 h. It was allowed to cool to room temperature and on further cooling (ice bath), the resulting precipitate was removed by filtration, washed separately with H₂O then recrystallized from MeOH. In some cases, the Schiff base was obtained as an oil. It was then treated with petroleum spirit (40–60°C) in an ice bath with continuous stirring until the oil solidified.

Preparation of complexes

Direct reactions giving $[PtLCl_2]_n$. A solution of K_2PtCl_4 (0.5 mmol) in distilled H_2O (5 cm³) was added to a solution of the Schiff base (L) (0.5 mmol) in EtOH (5 cm³), and an immediate precipitate formed. The mixture was stirred for several hours to ensure complete reaction. In some cases, precipitation occurs slowly and stirring overnight is necessary. The solid was then filtered off, washed several times with small portions of H_2O , cold EtOH and then with n-hexane and dried *in vacuo* for several hours.

Ligand displacement reactions giving $[PtLL'Cl_{2}]$: from cis- $[Pt(DMSO)_{2}Cl_{2}]$ (1:1 molar ratio). Cis- $[Pt(DMSO)_{2}Cl_{2}]$ (0.5 mmol) was suspended in CHCl₃ (15 cm³) and to this was added a solution of the Schiff base (L) (0.5 mmol) in CHCl₃ (15 cm³). The reaction mixture was heated to the boiling point of CHCl₃ under stirring for *ca* 30 min. The solution became clear, it was filtered through celite while it was hot and CHCl₃ was evaporated to a minimal volume, then ether was added to the point of turbidity. The solid thus formed was filtered off, washed several times with ether and dried *in vacuo* for several hours, yielding *cis*-[PtL(DMSO)Cl₂].

From cis-[Pt(DMSO)₂Cl₂] (1: 2 molar ratio). This type of reaction was carried out in a similar manner to those for 1: 1 molar ratio but using a 1: 2 molar ratio of *cis*-[Pt(DMSO)₂Cl₂] to Schiff base respectively, leading to *cis*-[PtLL'Cl₂] (L = L').

From $[Pt(COD)Cl_2]$. $[Pt(COD)Cl_2]$ (0.5 mmol) was suspended in CHCl₃ (10 cm³) and to this was added a solution of the Schiff base (L) (0.5 mmol) in CHCl₃ (10 cm³). The reaction mixture was heated to the boiling point of CHCl₃ under stirring for *ca* 30 min; no odour of COD was detected. The mixture was filtered through celite while hot and CHCl₃ was evaporated to leave a minimal volume. Ether was then added until turbidity. The solid thus formed was filtered off, washed several times with ether and dried *in vacuo* for several hours, giving *cis*-[PtL(COD)Cl₂].

Bridge-cleavage reactions giving trans-[PtLL'Cl₂]. The [PtCl₂L']₂ bridged complex (0.16 mmol) was dissolved in acetone (10 cm³) and to this was added a solution of the Schiff base (L) (0.32 mmol) in acetone (10 cm³). The reaction mixture was heated under reflux for *ca* 15 min, then filtered through celite. The filtrate was reduced in volume to *ca* 5 cm³ and petroleum spirit (40–60°C) was added to the point of turbidity and the sample was left in a cold place overnight. The crystals formed were filtered off, washed several times with petroleum spirit and dried *in vacuo* for several hours, yielding *trans*-[PtLL'Cl₂] (L = C₂H₄, PEt₃, PPh₃, PMe₂Ph).

RESULTS AND DISCUSSION

Physical data for the Schiff bases (II-16, III-7, III, IV) (Scheme 1) are listed in Table 1. The ¹H NMR spectra data for selected bases (I8, 13, 14, III, 5, III, IV) are given in Table 2. In one case (I8), the spectrum revealed the presence of signals related to *trans* (76%) and *cis* (24%) isomers. Thus, the *trans* isomer predominates in solution.

increases.¹³⁻¹⁶ The v(C=N) values together with v(Pt-Cl) values are good tools to distinguish between structure types A or B (B_{im} and B_{py}), since in types A, the *cis*-Cl₂ species gave two clear v(Pt-Cl) bands at *ca* 310 and 340 cm⁻¹,^{5,6,17} whereas in type B, the bridged chlorine species also gave two v(Pt-Cl) values, but the terminal one is at *ca* 330 cm⁻¹ and the bridged one is at *ca* 280 cm⁻¹ (much less intense than the terminal one).^{9,10}



The N-imine (central azomethine) and N-pyridine sites in the Schiff bases are coordinated with platinum metal ion either in bidentate fashion via both sites or monodentate via one site.

The physical properties of the complexes prepared from K_2PtCl_4 , *cis*-[Pt(DMSO)₂Cl₂], [Pt(COD)Cl₂] and [PtL'Cl₂]₂ (L' = C₂H₄, PEt₃, PPh₃, PMe₂Ph) with Schiff bases (L) are listed in Table 3; ¹H NMR data for selected complexes (most of them are poorly soluble) are listed in Table 2.

Direct reactions

The reaction between K_2PtCl_4 with Schiff bases (Scheme 1) in 1:1 molar ratio tentatively gave two types of complexes, type A and type B (Scheme 2). The Schiff base in type A coordinated as a bidentate ligand, giving four-membered ring complexes, while in type B, the Schiff base coordinated as a monodentate ligand via the most active site, either the Nimine (type B_{im}) or the N-pyridine (type B_{py}). These results were confirmed by CHN elemental analyses and by IR spectroscopy (Table 3). The IR spectra of the complexes obtained were compared with those of the free Schiff bases. A new band which appeared in the 492–568 cm^{-1} or 247–313 cm^{-1} regions is tentatively attributed to v(Pt-N) mode for N-imine or N-pyridine, respectively.⁶ The $v(C=N)_{im or py}$ of the free Schiff base is perturbed upon coordination; it either decreases or

Ligand displacement reactions

The reaction of cis-[Pt(DMSO)₂Cl₂] and [Pt(η^4 -COD)Cl₂] with Schiff bases (L) in 1:1 molar ratio lead to complexes of the general formulae cis- $[PtL(DMSO)Cl_2]$ and $cis-[PtL(\eta^2-COD)Cl_2]$, in which the Schiff base again seems likely to coordinate via the N-imine or N-pyridine site, type C $(C_{im} \text{ or } C_{py})$ complexes (Scheme 2). This fact was confirmed by the C=N, Pt-N and Pt-Cl stretching frequencies. It seems that complete displacement of 2DMSO or COD by one mole of the ligand did not occur. The presence of the remaining DMSO molecule was confirmed by the v(Pt-S)and v(S=0) modes which appeared in the 418–453 and 1155–1177 cm⁻¹ regions respectively (sulphurbonded sulphoxide).⁶ Further confirmation was provided by the ¹H NMR spectrum of cis-[PtL(DMSO)Cl₂] (Table 2), which exhibits a methyl resonance at $\delta \simeq 3.3$ ppm with ${}^{3}J({}^{195}\text{Pt-S-}$ CH) = 21 Hz related to coordinated DMSO. The resonance of the organic residue (L) remains almost constant upon coordination, apart from the CH=N resonance, which is shifted downfield by ca 1.5 ppm and is assigned to N-imine coordination (cf e.g. ligands III and III with their cis-[PtL(DMSO)Cl₂] complexes) (Table 2).

Similar behaviour was observed with the complexes *cis*-[PtL(η^2 -COD)Cl₂], in which COD coordinated more likely as a monodentate ligand with

	Vield	c M		Fo	und (Calc.)	%	Se	lected IR ^a bands	(cm ⁻¹)		VII
r	(%)	(°C)	Colour	С	Н	z	v(C==N) _{py}	ν(C==N) _{im}	v(C==C)	۷(HO)	λ_{\max} (nm)
II	89	61–69	yellow	72.0	4.9	13.8	1569 m	1623 s	1600 m	3425 b	267,385
				(72.7)	(5.1)	(14.1)					
12	80	136-140	white	72.9	5.1	13.1	1578 s	1636 s	1614 m,sh		275
13	85	86– 98	vellow	(73.6) 68.3	(5.7) 5.1	(13.2)	1575 m	1619 m	1595 m	3401	775570
1	2	2		(68.4)	(5.3)	(12.3)				10+0	010.0.117
I4	90	87-89	yellow	73.2	5.1	12.9	1578 s, sh	1620 s	1600 m	3066	295,329
;	i I			(13.6)	(5.7)	(13.2)					
IS	79	156-160	orange	59.0	3.9	17.1	1587 s	1659 s,sh	1600 s	3410	282,380
ì	ç			(59.3)	(3.7)	(17.3)					
9	83	y0-y4	white	66.1 (67-7)	4.9	12.0	1590 s	1632 s	obscured		290,331
17	06	158-160	brown	54.8	3.1	(177)	1 591 m	1650 m sh	1605 m		776 385 5
				(55.1)	(3.1)	(16.1)					0.00.00.14
I8	80	130-132	off-white	68.0	5.4	11.2	1585 s	1625 s	1608 s		313,355
				(68.7)	(5.3)	(11.5)					
6I	75	92–96	off-white	64.5	4.4	17.2	1582 m	1627 s	1602 m		282
				(64.7)	(4.6)	(17.4)					
I10	81	8284	yellow	66.0	5.2	16.2	1596 s	1640 s	1596 s		292
				(62.9)	(5.1)	(16.5)					
111	76	118-120	white	55.9	4.6	9.8	1596 m	1660 s	1614 s	3200	238,291
				(56.5)	(4.4)	(10.1)					
112	80	148-150	pale yellow	58.1	5.0	9.4	1600 w	1650 m	1623 s	3205	282
			;	(57.9)	(4.8)	(9.7)					
113	95	166–168	yellow	60.0	4.4	15.9	1578 s	1632 s,sh	1590 s		309,389
				(60.7)	(4.3)	(16.3)					
I14	93	165–168	yellow	58.7	4.7	14.2	1591 s,sh	1654 s,sh	1623 s,sh		312,391
				(58.5)	(4.5)	(14.6)					
I15	71	116-120	grey	54.9	3.7	15.3	1591 m	1659 s	1614 s	3289	314,375
,				(55.6)	(3.5)	(16.2)					
116	83	89102	yellow	59.0	4.0	16.8	1591 m	1650 m	1610 m	3207	314,385
				(59.3)	(3.7)	(17.3)					
III	85	68-70	orange	72.1	4.9	14.1	1583 m, sh	1641 s	1596 s	3333	305,333
				(72.7)	(5.1)	(14.1)					

Table 1. Physical properties of the Schiff bases L II-16, III-7, III, IV (Scheme 1)

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Table	2.	۱H	NMR	data ^a	for	selected	Schiff	bases	and
			S	elected	l coi	nplexes			

												selected complexes
				_			~).5	Compo	ound	δ (ppm), J values and assignments ^b
378	337		865	237,311	273	335	777 380	· · · · · · · · · · · · · · · · · · ·	278,37(18		<i>trans</i> (76% proportion) : 2.2 (d, 2CH ₃), 6.3 (s, Ph), 7.2–7.4 (m, py), 9.7 (s, CH=N) <i>cis</i> (24% proportion) :
3403			3156		3167		2464		3495	I13		2.4 (d, 2CH ₃), 7.0 (s, Ph), 7.5–7.9 (m, py), 11.0 (s, CH=N) 4.0 (s, OCH ₃), 6.5–7.2 (m, Ph), 7.9–8.1 (m,
s	В		s	s	s	Ш	49	116,111	E	I14		py), 8.7 (s, CH==N) 3.85 (s, CH ₃ CO), 4.0 (s, OCH ₃), 6.5–7.6 (m, Ph), 7.7–8.9 (m, py), 10.25 (s, CH==N)
1592	1600		1592	1605	1609	1587	1619	0101	1587	111		6.8–7.9 (m, Ph), 8.4–8.6 (m, py), 8.9 (s, CH=N), 12.75 (s, OH)
s	В		s,sh	s	s	s	£	3	E			7.55 (d), 7.9 (d, $J = 10$ Hz, Ph), 8.4–8.6 (m, py), 8.7 (s, CH=N) 4.85 (s, CH ₂), 6.6–8.7 (m, Ph and py), 8.75
1629	1628		1620	1632	1636	1620	1637	7001	1636	IV		(s, CH==N) 6.8-9.05 (m, Ph and quin), 9.2 (s, CH==N), 12.9 (s, OH)
569 m	591 m		578 s,sh	581 s	587 s,sh	269 m	L03	111 / 0/	587 m	cis-[Pt] L =	L(DM II	$1SO)Cl_2]3.45 (s, {}^{3}J({}^{195}Pt-S-CH) = 21.1 Hz,DMSO), 6.5-7.9 (m, Ph and py), 9.05 (s,CH=N)$
1	15	(1	11	11		-		L =	= I10	$2.75 (s, CH_3), 2.9 (s, CH_3), 3.4 (s, {}^{3}J({}^{195}Pt-S-CH) = 20.7 Hz, DMSO), 6.2-8.2 (m, Ph and py), 10.4 (s, CH=N)$
11.8	(12.0 11.2	(11.4	11.9 (12.0	13.1 12.9	10.9	(11.3 10.0	(10.1	(13.2	11.0	L =	= II1	3.3 (s, ${}^{3}J({}^{195}\text{Pt-S-CH}) = 21.0 \text{ Hz},$ DMSO), 6.7–9.4 (m, Ph and py), 10.7 (s, CH==N), 12.0 (s, OH)
4.0	(3.9) 4.7	(4.5)	4.0 (3.9)	4.3 (4.2)	4.0	(3.6) 5.2	(4.7) 5.4	5.7)	4.4 (4.8)	L =	= II5	3.5 (s, ${}^{3}J({}^{195}Pt-S-CH) = 25.3$ Hz, DMSO), 7.0–9.1 (m, Ph and py), 10.1 (s, CH=N)
0.	(6. 8.	(£.)	r. 6.	. 8 .	<u>)</u> 0:	(6; -2	(8)	. (9)	4)	L =	: III	3.3 (s, ${}^{3}J({}^{195}\text{Pt-S-CH}) = 22.3 \text{ Hz}$, DMSO), 4.93 (s, CH ₂), 6.5–9.2 (m, Ph and py), 10.2 (s, CH=N)
61	(61 62	(63	(61 (61	65 166	57	(57 61	(6 (6	c (13	<u>(</u> (((((((()))))	trans-[L =	[PtL(0 = I4	$C_{2}H_{4})Cl_{2}$ 2.4 (s, CH ₃), 4.25 (s, ² J(¹⁹⁵ Pt-CH) = 65
brown	off-white		grey	off-white	brown	off-white	elle	yenow	yellow	L =	= I13	Hz, C ₂ H ₄), 0.0–8.5 (iii, Fit and py), 9.4 (s, CH==N) 4.3 (s, OCH ₃), 4.3 (s, ${}^{2}J({}^{195}\text{Pt-CH}) = 66$ Hz, C ₂ H ₄), 7.2–8.85 (m, Ph and py), 9.2 (s
0-62	2–56		6–188	274	2194	0-112	0 E 4	+	8-199	L =	= II5	CH=N) 4.3 (s, ${}^{2}J({}^{195}Pt-CH) = 65.5 Hz, C_{2}H_{4}),$ 7.6-8.6 (m, Ph and py), 9.3 (s, CH=N)
Ŷ	ζ.		18	7.	19.	11	ŭ	n	19	trans-[L =	[PtL(I = II1	$PMe_2Ph)Cl_2$ 1.6 (d, CH ₃), 1.8 (d, CH ₃ , $J = 5.6$ Hz), 6.7–8.4 (m, Ph and py), 9.0 (s, CH=N),
73	79		83	76	86	81	0 <i>6</i>	с к	80	sllu «Do	ownfie	10.25 (s, OH)
112	II3		114	IIS	116	117	H	III	IV	loin a solve Null	ent. ultipli	cities in parentheses.

			Ţ	able 3. T	he prop	erties of p	datinum comj	olexes of the	Schiff bases					
			ž	Eour	of Calo	70		Selecte	d IR ⁶ bands	(cm ⁻¹)				
Complex	Structural assignment ^a	Colour	(°C) (Dec.)	C		v X	$\nu(C=N)_{py}$ (Δ)	ν(C==N) _{im} (Δ)	v(Pt-Cl)	v(PtN)	v(others)	$\lambda_{\rm max}$ $\lambda_{\rm max}$ (nm)	$\Lambda(\Omega^1 cn)$ DMF	lctivity 1 ² mol ⁻¹) MeNO ₂
$[PtLCl_2]_n = 13 n = 2$	ď	mollen	748 750	30.0	ч с	2 Y	1 573	12120	00	10		000	u c	
	Ē	brown	007-047	(31.6)	(2.4)	(2.7)	(-2)	(-3)	330w,	W040		086	C7	71
$\mathbf{L}=\mathbf{I6}, n=2$	B	deep	216-218	31.3	2.4	5.4	1585m	1632m	288m,	506m		237.	16	12
		yellow		(31.4)	(2.2)	(5.6)	(-5)	(0)	310m			300		
L = I8, n = 1	Α	deep	180-182	33.1	2.7	5.3	1578s	1641s	288m,	526m,		325,	39	35
		yellow		(32.9)	(2.5)	(2.6)	(– –)	(+16)	308m	261w		420		
L = I10, n = 2	\mathbf{B}_{py}	light	284-286	32.0	2.7	7.9	1578m	1632s	288w,	254w		306,	10	5
		brown		(32.2)	(2.5)	(8.1)	(-18)	(-8)	312m, sh			428		
L = III, n = 1	A	yellow	276-278	29.1	2.3	5.4	1564m	1690s	285m,	492w,		237,	48	10
				(28.8)	(2.2)	(5.2)	(-32)	(+30)	305m	268w		290		
L = I13, n = 2	Bim	yellow	200-204	30.0	2.2	7.9	1580m	1650s,sh	291w,	568w		308,	22	19
				(29.8)	(2.1)	(8.0)	(+2)	(+18)	319m			388,		
	I											444		
L = II4, n = 2	Bin	yellow-	120-122	28.9	2.4	7.3	1591s	1650s	295w,	553w		313,	16	7
		brown		(30.4)	(2.4)	(1.6)	(0)	(-4.5)	323m			389, 433		
L = III, n = 2	\mathbf{B}_{in}	yellow	262-264	30.8	2.3	5.8	1587w	1620s	288w,	547w		236,	40	22
				(31.0)	(2.2)	(0.9)	(+4)	(-21)	322w			293		
L = II4, n = 1	A	yellow-	264–268	29.0	2.0	5.5	1590s	1600s	308m,	528w,		208	10	9
		brown		(28.9)	(1.8)	(2.6)	(+12)	(-20)	342s	252w				
L = II5, n = 2	$\mathbf{B}_{\mathbf{p}y}$	off-white	136-140	30.0	2.0	5.7	1596s	1632m	288w,	265w		234,	40	
				(29.8)	(1.9)	(5.8)	(+15)	(0)	322m			301	22	
L = II6, n = 2	B.	brown	288–290	28.1	2.0	5.2	1590s,sh	1632s	302w,	496w		327,	11	9
				(28.0)	(1.8)	(5.4)	(+3)	(-4)	326m			412		
cis-[PtLL'Cl ₂]														
$\mathbf{L} = \mathbf{I}\mathbf{I},$	C _{in}	yellow	170-174	30.8	2.7	4.9	1573s	1618s	293w,	524w	449s,	297,	21	£
L' = DMSO				(31.0)	(3.0)	(5.2)	(+4)	(-5)	310w,		1155m	381		
L = I10 ,	C _{py}	yellow-	140–144	32.0	3.0	7.0	1578m	1641s	308m,	274m	453s,	274	14	5
L' = DMSO		brown		(32.1)	(3.2)	(1.0)	(-18)	(+1)	331m,sh		1168s			

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ę	4	5	6	S		7		D		51		59		45		50		54		ę		×		-			1		5		4	
13	15	17	16	11		26	-	10		52	i	72		48		53		60		7		11		4			4		21		20	
250,	277 254,	302 302	255,	301 295.	360	253,	301	230, 384		247,	289	234,	345	251.5,	385	245,	294	246,	295	261,	381	261,	316	244,	289,	393	286,	382	224,	396	300	
453s,	1170s 427s,	418s,	1177s 451w	442w		441w		429W		444w		431m		431w		442w		431m		455w		458w		455m			454m		482m		462w	
516m	502m	547w	553w	248m		248m		24/M		520m,b		567m,b		543w		537w		547w		251w		248w		250w			290w		535w		517m,b	
301s,	326s 302w, 220	330m 303w,	336w 308s,	320m 303s.	316m	305s,	312m 202	308S, 317s		309s		319s		315m		333s		313m		319s		302w		319m			325w		314w		306w	
1652s	(+20) 1677m	(+4) 1641s	(+9) 1661s	(+29) 1624m	(+4)	1636m	(+4)	10355 (-1.5)		1663s	(+43)	1641s	(+)	1632w	(-18)	1641s	(6+)	1632m	0	1628s	(+8)	1641s	(+16)	1641s	(6+)		1641s	(+)	1674s	(+19.5)	1655m	(+19)
1585s	(+7) 1578m	(-3) 1591s	(+4) 1578m	(-12) 1583m	(+5)	1592w	(+5)	(-3)		1578s	(0)	1578m	0	1587w	(-4.5)	1565w	(-16)	1587m	0)	1582s	(+4)	1587m	(+2)	1587w	(6+)		1586s	(+8)	1592s	(+0.5)	1593m	(9+)
7.1	(7.0) 5.2	(U.C) 8.4 8.5	(5.0) 4.9	(4.6) 4.1	(4.6)	4.1	(4.8)	4.8 (4.5)		5.2	(5.5)	7.4	(1.6)	7.9	(7.8)	5.2	(5.5)	5.7	(5.5)	4.6	(4.7)	4.4	(4.5)	6.2	(6.6)		5.2	(5.4)	5.2	(5.2)	3.5	(3.6)
3.0	(2.8) 2.7	(7,7)	(3.2) 3.1	(3.8) 3.6	(3.5)	3.9	(4 .1)	3.9) (3.9)		3.2	(3.2)	2.9	(2.7)	2.8	(2.4)	2.6	(2.5)	3.0	(3.2)	4.9	(4.5)	4.8	(4.5)	3.9	(4.1)		3.4	(3.3)	3.5	(3.4)	3.2	(3.1)
29.0	(30.0) 30.1	(30.0) 31.1	(32.4) 41.2	(41.7) 39.3	(39.6)	42.9	(43.0)	4.2.9 (46.3)		35.4	(35.6)	32.7	(32.7)	30.9	(31.3)	33.0	(32.9)	35.2	(35.6)	38.0	(38.3)	37.9	(38.2)	35.5	(35.6)		47.0	(47.4)	47.0	(47.1)	46.3	(46.4)
200-204	67–80	100-104	128-130	180-184		142–144		180-182		80-82		170-172		72–74		120-124		140–144		128-130		144–148		108 - 110			138-140		82–86		90–92	
pale	yellow yellow	yellow	white	off-white		yellow	:	yellow		yellow		yellow		cacao		yellow		yellow		off-white		off-white		yellow			yellow		yellow		brown	
C.	C _{im}	C _{im}	C	U U	ki -	C_{py}	C	C py		D		D		D _{im}		\mathbf{D}_{im}		D		D _{by}		D		D_{py}			D		D		D.	
$\mathbf{L}=\mathbf{I13},$	L' = DMSO L = II5 ,	L = UMSU $L = III,$	L' = DMSO L = I 6,	L' = COD L = II4 .	L' = COD	$\mathbf{L} = \mathbf{III},$	L' = COD	L = IV, L' = COD	<i>trans</i> -[Pt1.]_(C1,1	$\mathbf{L} = \mathbf{I4},$	$L' = C_2 H_4$	L = II3,	$L' = C_2 H_4$	L = 116,	$L' = C_2 H_4$	L = II5,	$L' = C_2 H_4$	$\mathbf{L} = \mathbf{III},$	$L' = C_2 H_4$	L = I4 ,	$\mathbf{L}' = \mathbf{E}\mathbf{t}_3\mathbf{P}$	L = I8 ,	$\mathbf{L}' = \mathbf{E} \mathbf{t}_3 \mathbf{P}$	L = II 3,	$\mathbf{L}' = \mathbf{E} \mathbf{t}_3 \mathbf{P}$		L = I13,	$L' = PPh_3$	$\mathbf{L}=\mathbf{I14},$	$\mathbf{L}' = \mathbf{P}\mathbf{P}\mathbf{h}_3$	L = II6,	$L' = PPh_3$

Platinum group Schiff base complexes-l

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			ň	Four	d (Calo	707		Selected	i IR ⁶ bands ((cm^{-1})			ζ	
	Structural		.d.o	IIDO.I	חורמור	0/ (U V-VIS		cuvity
Complex	assignment ^a	Colour	(Dec.)	C	Н	Z	(Δ)	(Δ)	v(Pt-Cl)	v(Pt-N)	v(others)	A _{max} (nm)		MeNO ₂
L = IS, I ′ = Me.PhP	D _{py}	yellow	130–134	36.9	3.2	6.4 (6.5)	1582s	1642s	302m,sh	288m	472m	261, 307	17	10
L = III, L' = Me ₂ PhP	\mathbf{D}_{py}	yellow- brown	110-112	(1.17) 39.0 (39.9)	(3.5) (3.5)	(5.0) (4.7) (7.7)	() 1587s (+4)	(-9)	327m	303w	472m	256, 271,	13	e
L = II 3, L' = Me ₂ PhP	$\mathbf{D}_{\mathbf{py}}$	light brown	8890	39.0 (38.7)	3.9 (3.4)	(4.3)	1579m (-12.5)	1629s (+1)	323w	313w	439m	366 304	4	7
<i>cis</i> -[PtLL'Cl ₁] L = L' = III	Ë	pale	92–94	, 44.0 0	3.5) oo	1582s	1632s	288w.	549m		301.	14	6
L = L' = 115	$\mathbf{E}_{\mathbf{py}}$	brown pale	230–234	(43.5) 41.1	(3.0) 2.9	(8.5) 8.1	(-1) 1596s	(-9) 1632s	310w 295w,	247m,		377 344	17	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
		yellow		(41.2)	(2.6)	(8.0)	(+14)	(0)	302w	275m				
^a The symbol i	m represents cool	rdination v	ia the imine	site and	the sym	bol py re	presents coord	lination via t	he pyridine s	ite.				

Table 3-continued

^{*b*} In Nujol mulls, v(others); L' = DMSO, 418–453 cm⁻¹ for v(Pt—S), 1155–1177 cm⁻¹ for v(S=O); L' = COD or C₂H₄, 431–451 cm⁻¹ for v(HC=CH); $L' = PEt_3$, PPh₃ or PMe₂Ph, 439–482 cm⁻¹ for v(Pt—P).



Scheme 2. The suggested structures for the Schiff base/platinum complexes.

platinum and was not totally removed from the coordination sphere. The characteristic odour of COD was not detected from the reaction mixture. Additional confirmation is that a band appeared in the 441–451 cm⁻¹ region characteristic of the v(HC=CH) mode.

In the 1:2 molar ratio reaction between cis-[Pt(DMSO)₂Cl₂] and Schiff bases (L), the complexes obtained were assigned to the formula cis-[PtL₂Cl₂], complex type E (Scheme 2), and no sign of DMSO could be detected by IR spectroscopy.

Bridge-cleavage reactions

These were carried out using the bridge starting complex $[PtL'Cl_2]_2$ $(L' = C_2H_4, PEt_3, PPh_3,$ PMe₂Ph) with Schiff bases (L) in 1:2 molar ratio, respectively. The complexes obtained were assigned the general formula trans-[PtLL'Cl₂] (type D complexes) and were identified by their CHN elemental analyses, IR and ¹H NMR spectra. The v(Pt-Cl) mode for *trans*-Cl₂ species is well identified by the single band^{4,18} in the 310-330 cm⁻¹ region for all the complexes obtained (Table 3). The Schiff bases again coordinated via either the N-imine or the Npyridine and this was confirmed by the significant change in the v(C=N) values (see below). The ¹H NMR of trans-[PtL(C_2H_4)Cl₂] (L = I4, II3, II5) showed a single resonance at ca 4.3 ppm with $^{2}J(^{195}\text{Pt-CH}) \cong 65$ Hz assignable to coordinated C_2H_4 .^{4,18} The resonance of the residual organic species (L) was also observed and its significance is that the δ (CH=N) value was shifted downfield by ca 0.5 ppm (Table 2) upon coordination. This gave

further support to what the IR suggested; namely that the coordination of the N-imine site of the Schiff base had certainly taken place. Additional support for our argument is that the δ (CH==N) value for the free base (III) (8.9 ppm) remains almost constant upon coordination in the complex *trans*-[PtL(PMe₂PH)Cl₂] (9.0 ppm) assigning Npyridine coordination and this is in good agreement with that found by IR spectroscopy, Δ [ν (C==N)_{py}] = 12.5 cm⁻¹ and Δ [ν (C==N)_{im}] = 1 cm⁻¹.

The UV-vis spectra of 10^{-3} M solutions of the complexes showed bands at ~250 and ~370 nm (Table 3) due to the Schiff base, plus a band which appeared, in some cases, at ~400 nm attributed to the *d*-*d* transitions of platinum.

With few exceptions, the molar conductivities of 10^{-3} M solutions of almost all the complexes [PtLCl₂]_n (n = 1 or 2), cis-[PtLL'Cl₂] (L' = DMSO, η^2 -COD, L) and trans-[PtLL'Cl₂] (L' = PEt₃, PPh₃, PMe₂Ph) indicate that they are non-electrolytes in both MeNO₂ and DMF (Table 3). On the contrary, all the complexes trans-[PtL(C₂H₄)Cl₂] (L = **14**, **II3**, **II6**, **II5**, **III**) gave results consistent with some 1:1 electrolytes in both solvents. This may be described on the basis of ligand replacement; C₂H₄ is a labile ligand which can easily be replaced by donor solvents and may give the ionic species [PtL(solv.)₂Cl]⁺Cl⁻.

In one experiment, the complex $[PtLCl_2]_n$ (L = II5), with molar conductivity in DMF of 40 Ω^{-1} mol⁻¹ cm², was left in solution for 24 h; the conductivity increased to 84 Ω^{-1} mol⁻¹ cm², which is clearly assigned to 1:1 conductivity¹⁹ and this may similarly be described as solvent interaction to give $[PtL(solv.)_2Cl]^+Cl^-$ species.

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