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Preparation of the Complexes *trans*-[PtX₂(CO)L] and their Isomerisation to *cis* Isomers

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Treatment of halide-bridged dimers $[\{PtX_2L\}_2]$ (X is halide; L is tertiary phosphine or arsine) by carbon monoxide gives solutions of the *trans* isomers of $[PtX_2(CO)L]$, some of which can be isolated. These compounds readily lose CO and isomerise to *cis* derivatives. The *trans* to *cis* isomerisations are catalysed by free CO, phosphines, or halide ions, and the latter nucleophile operates even in the solid phase. Free CO exchanges with co-ordinated CO more rapidly with the *trans* isomer than the *cis*. The *trans* to *cis* isomerisations are accelerated by light or u.v. irradiation, and u.v. treatment of the thermodynamic *cis* isomers produces *trans-cis* mixtures. The mechanisms of all these processes are discussed.

MANY carbonyl complexes of platinum(II), $[PtX_2(CO)L]$ (X is halide; L is PR₃ or AsR₃), have been isolated ^{1,2} and used in spectroscopic ^{3,4} or synthetic ^{5,6} studies. They are usually prepared by CO cleavage of the halidebridged dimers *trans*-[{PtX₂L}₂]. The first report of their preparation suggested that a *trans* geometry for [PtX₂(CO)L] would be expected,¹ because of the high *trans* effect ⁷ of the tertiary phosphine or arsine ligands. In the event, the compounds isolated have always had the *cis* configuration,^{1,2} and it has become accepted practice simply to generate and use them *in situ*,⁶ confident of their geometry.

We report that the first products generated in solution by the action of CO on $[{PtX_2L}_2]$ are in fact the *trans* isomers of $[PtX_2(CO)L]$. Although these complexes readily isomerise to the *cis* derivatives, under certain conditions they can exist for considerable times and some can be isolated as solids. A preliminary account has appeared.⁸

RESULTS AND DISCUSSION

(a) Preparation and Spectroscopic Parameters.—Bubbling carbon monoxide through suspensions of $[{Pt-X_2L}_2]$ in chloroform, benzene, or toluene at room temperature produced solutions of *trans*-[PtX₂(CO)L] $(X = Cl; L = PBu_3, PEt_3, PMe_2Ph, PMePh_2, PPh_3, AsMePh_2, or AsPh_3; X = Br; L = PMe_2Ph)$ [equation (1)]. Reactions were complete in *ca.* 20 min, though if

$$trans - [\{PtX_2L\}_2] \xrightarrow{CO} trans - [PtX_2(CO)L] \longrightarrow cis - [PtX_2(CO)L] \quad (1)$$

passage of CO was continued for 1-2 h, colourless crystals of cis-[PtX₂(CO)L] precipitated.^{1,2} In the absence of an excess of CO, however, isomerisation is slow, taking 1-3 d at room temperature. Below *ca*. -30 °C, solutions of the *trans* isomers remain unchanged indefinitely in the dark, i.r. and ³¹P n.m.r. investigations revealing no trace of the *cis* isomer after several days.

Attempts to isolate the *trans* complexes by crystallisation frequently led to *cis-trans* mixtures, or even to the pure *cis* isomer. Spectroscopically pure samples of the yellow solid complexes *trans*-[PtCl₂(CO)(PMePh₂)], *trans*- $[PtCl_2(CO)(PPh_3)]$, and trans- $[PtCl_2(CO)(AsMePh_2)]$ were obtained, however, by carefully removing benzene solvent from frozen solutions at -10 °C, or CH_2Cl_2 at -50 °C. These samples have been kept at -10 °C in the dark for several days, and when redissolved showed only the original trans isomers. Attempts to analyse the compounds for C and H by the usual combustion methods failed, however, resulting in loss of CO to regenerate $[\{PtCl_2L\}_2]$ prior to combustion.

Phosphorus-31 n.m.r. spectroscopic parameters and $\nu(CO)$ from the i.r. spectra of the complexes are presented in the Table. Values of $\delta(P)$ and $\frac{1}{(Pt-P)}$ are quite similar in both isomers, and although $^{1}/(Pt-P)$ is consistently larger in the trans isomers, values are too close for this to be a reliable diagnostic test for geometry. Values of $\nu(CO)$, on the other hand, are always 20–30 cm⁻¹ higher in the *trans* isomers, and the characteristic values of ca. 2 137 cm⁻¹ are clearly diagnostic of trans compounds. Samples of trans-[PtCl₂(CO)(PEt₃)] labelled with ¹³CO [δ (¹³C), 169.9 p.p.m.; ¹J(Pt-C) 1 164 Hz; ²J(P-C) 159.6 Hz] and trans-[PtCl₂(CO)(PMePh₂)[δ (¹³C), 168.3 p.p.m.; ${}^{1}J(P-C)$ I 221 Hz; ${}^{2}J(P-C)$ 171.7 Hz] were prepared in the usual way. The most striking features of their ¹³C n.m.r. spectra are the high values of ${}^{2}J(P-C)$. These are entirely typical of the few values reported for CO trans to PR₃ at platinum,⁵ values for cis isomers being typically only 5-10 Hz.

Two interesting comparisons emerge at this stage. Firstly, a few *trans* platinum carbonyls, [PtCl₂(CO)L], have been prepared previously, generally by replacing C_2H_4 in trans-[PtCl₂(C_2H_4)L] by CO (L is substituted pyridine, pyridine N-oxide, or aniline).9 Although these compounds can be readily isolated, and appear more stable with respect to CO loss (see below) or isomerisation than the phosphine or arsine analogues described here, related amine derivatives do nevertheless isomerise on heating.¹⁰ Secondly, the generation of *trans* complexes as the first products of bridge-cleavage reactions of trans- $[M_2X_4(PR_3)_2]$ (M = Pd or Pt) by a variety of nucleophiles (now including CO) is well known and seems to be a general process.¹¹ It is a reasonable supposition that the high trans effect of PR₃ is responsible, but this kinetic phenomenon may be more subtle than that. The dinuclear complex $[Pt_2(\mu-Cl)_2Ph_2(PMePh_2)_2]$ exists in solution as similar amounts of both *cis* and *trans* isomers,^{6,12} but carbonylation produces 98% of the isomer of $[PtClPh(CO)(PMePh_2)]$ with CO *trans* to phosphorus ¹³ [which in turn isomerises to other forms,⁵ equation (2)]. Clearly some rearrangement must take place at the intermediate stages to avoid producing more of the thermodynamic isomer.

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monoxide is therefore released during the *trans* to *cis* conversion. Solutions of *cis*-[PtCl₂(CO)L] showed no tendency to lose CO when treated in this way.

The isomerisation is independent of concentration. Chloroform solutions of $trans-[PtBr_2(CO)(PMe_2Ph)]$ at four concentrations ranging from 0.025 to 0.0125 mol dm⁻³ were periodically examined by i.r. spectroscopy. After each time interval, all four solutions showed

Phosphorus-31 n.m.r.^a and i.r.^b spectroscopic characteristics for cis- and trans-[$PtX_2(CO)L$]

x	L	cis Isomers			trans Isomers		
		$\delta(P)/p.p.m.$	$^{1}J(\text{Pt-P})/\text{Hz}$	$\nu(CO)/cm^{-1}$	$\delta(P)/p.p.m.$	¹ J(Pt-P)/Hz	v(CO)/cm ⁻¹
Cl	PEt.	22.3	2 793	2 106	11.6	2885	$2\ 135$
Cl	PMe.Ph	10.4	2855	$2\ 112$	-11.0	2 933	$2\ 138$
Cl	PMePh,	-0.2	2 946	2 115 *	0.6	3 000	2 139 ª
Cl	PPh,	9.8	3 061	$2\ 115$	15.1	3 077	2 139
Cl	PBu.	14.3	2783	$2\ 106$	3.9	2865	$2\ 135$
Cl	AsMePh.			2 109			$2\ 133$
Cl	AsPh,			2 110			2 134
Br	PMe,Ph	-11.2	2822	$2\ 107$	-15.9	2825	$2\ 132$
I	PMe [•] Ph	-14.7	2 748	2 096			
Cl	$P(C_{a}H_{1})$	36.4	2850	2099			
Cl	$P(o-tolyl)_3$	5.6	3 010	2 110			

^{*a*} At 25 °C in CDCl₃: chemical shifts measured downfield of external H_3PO_4 . ^{*b*} At 20 °C in CDCl₃. ^{*c*} ν (1³CO) 2 065 cm⁻¹. ^{*d*} ν (1³CO) 2 090 cm⁻¹.

(b) Isomerisations of trans- $[PtX_2(CO)L]$.—Solutions of trans- $[PtCl_2(CO)L]$ changed entirely to the *cis* isomer after standing from 1—3 d at ambient temperatures. The bromide complex trans- $[PtBr_2(CO)(PMe_2Ph)]$ converted faster, and only *cis*- $[PtI_2(CO)(PMe_2Ph)]$ could be



detected after passing CO through chloroform suspensions of the sparingly soluble $[Pt_2I_4(PMe_2Ph)_2]$. This probably reflects an even faster conversion of a *trans* isomer to the *cis*, but the possibility cannot be ruled out that the *trans* isomer may not be formed at all with this halide.

All the *trans* complexes tended to lose carbon monoxide from solution, and their isomerisations were often accompanied by the production of 5-20% of $[{PtX_2L}_2]$. When a stream of nitrogen gas was passed through a benzene solution of *trans*- $[PtCl_2(CO)(PMePh_2)]$, total loss of CO had occurred after 5 h. An identical solution not treated by N₂ contained a 1 : 1 mixture of *cis*- and *trans*- $[PtCl_2(CO)(PMePh_2)]$ after the same time. Carbon identical isomer ratios. On the other hand, samples of trans-[PtCl₂(CO)(PMePh₂)] in benzene isomerised twice as fast as solutions of similar concentration in chloroform. This suggests a faster isomerisation in non-polar solvents, but difficulties in observing i.r. spectra in other solvents have so far prevented substantiation of this point.

The trans to cis isomerisations are dramatically catalysed by free carbon monoxide. A 0.1 mol dm⁻³ solution of trans-[PtCl₂(CO)(PMePh₂)] was completely converted to the *cis* isomer in only 35 min with passage of CO through the solution, whilst an identical, untreated solution was virtually unchanged in this time interval. A sample of trans-[PtCl₂(¹³CO)(PMePh₂)] was similarly treated with ¹²CO. After 15 min similar amounts of both trans-[PtCl₂(¹³CO)(PMePh₂)] and trans-[PtCl₂(¹²CO)(PMePh₂)] were apparent in solution, whilst after 25 min the major species was trans-[PtCl₂(¹²CO)-(PMePh₂)]. Quantities of the two *cis* isomers were by then also apparent. Again, an untreated solution of trans-[PtCl₂(¹³CO)(PMePh₂)] remained unchanged after this time interval. When ¹²CO was passed through a chloroform solution of cis-[PtCl₂(¹³CO)(PMePh₂)], exchange of CO was considerably slower than for the trans isomer, requiring ca. 1 h for 30% exchange. No trans complex was detected during this experiment.

Other nucleophiles also catalyse these *trans* to *cis* isomerisations. The addition of 0.1 molar equivalents of PMePh₂ to solutions of *trans*-[PtCl₂(CO)(PMePh₂)] caused complete conversion to the *cis* isomer in 30 min. Likewise, small amounts of $[NEt_4]Cl$, $[NBu_4]Br$, or $[NBu_4]I$ promoted complete *trans* to *cis* conversions in 1 h. Partial loss of CO was evident, presumably due to the formation of some $[PtCl_2(PMePh_2)_2]$ and $[PtCl_2X-(PMePh_2)]^-$, respectively, by CO displacement. It seemed possible that the displaced carbon monoxide

might be the true catalyst in these experiments, but this was discounted by the following experiment. To two of three identical 0.01 mol dm⁻³ chloroform solutions of trans-[PtCl₂(CO)(PPh₃)] were added 0.5 mg of PPh₃ and $[NEt_4]Cl$, respectively, and a fast stream of N_2 was passed through all three samples for 15 min. Infrared examination then revealed complete conversion to the cis isomers in the samples treated by PPh₃ or Cl⁻, but ca. 25% CO loss with no isomerisation in the untreated sample. Thus, while exchange of free and co-ordinated CO is faster than the CO-catalysed isomerisation, replacement of CO by PR₃ or X⁻ is slower than isomerisation catalysed by these nucleophiles. The halidecatalysed interconversion at least also operates in the solid phase. Grinding samples of trans-[PtCl₂(CO)L] $(L = PPh_3 \text{ or AsMePh}_2)$ with KBr caused substantial isomerisation. After the mixtures were compressed into a disc for i.r. measurements, the isomerisation continued. Grinding the complexes alone had no effect: redissolution in chloroform showed that no isomerisation had occurred.

(c) Isomerisation Mechanism.—Ligand substitution reactions at platinum usually proceed stereospecifically by an associative mechanism,¹⁴ and CO exchange probably follows this route [equation (3)]. Nucleophile-catalysed isomerisations are also associative,¹⁵ and the same five-co-ordinate intermediate is probably involved in the CO-catalysed process. The geometry change can come about either through pseudorotation, or consecutive displacement of L and/or X. These can be regarded as two extremes of one basic process.¹⁵ We note that our conditions fit exactly those predicted by Redfield and Nelson ¹⁶ as most favourable to pseudorotation, and that L and X^- (either or both of which would have to be eliminated during consecutive displacement reactions) themselves catalyse the isomerisations.

$$L \xrightarrow{X} CO \xrightarrow{*CO} L \xrightarrow{X} CO \xrightarrow{*CO} CO \xrightarrow{-CO} L \xrightarrow{X} CO (3)$$

The Scheme represents some possible routes. Trigonalbipyramidal intermediates rather than square-pyrimidal intermediates are depicted for clarity (both may be encountered during isomerisation sequences). No inference concerning the relative energies of these five-coordinate species should be drawn.^{13,15}

Intermediates A and C (Scheme) must be involved in the stereospecific exchange of CO with the trans and cis isomers, respectively, so it is tempting to suggest that the isomerisation proceeds from A through B to C. The other routes cannot, however, be ruled out. Similar schemes can be drawn for the phosphine- and halidecatalysed reactions.

The uncatalysed isomerisation is first order in Pt and involves CO loss from the trans isomer, either spontaneously or solvent assisted. In view of the ready catalysis by CO of this isomerisation, it seems very likely that the eliminated CO would catalyse the conversion

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of the remaining, undissociated, trans molecules [equations (4) and (5)]. Such a process is essentially

$$trans-[PtX_2(CO)L] \iff [PtX_2L] + CO \qquad (4)$$

$$trans-[PtX_2(CO)L] \xrightarrow{CO} cis-[PtX_2(CO)L]$$
 (5)

autocatalytic. Whilst no other systems have been claimed to conform to such a mechanism, there is good evidence that many other isomerisations of squareplanar complexes may be similar.¹⁵

A number of alternative mechanisms must be considered, all of them involving CO loss. A strong contender is the possibility of regenerating the dimeric



 $[{PtX_{2}L}_{2}]$, which might slowly but irreversibly build up the cis isomer on CO cleavage [equation (6)]. This was

$$cis-[PtX_2(CO)L] \xrightarrow{CO} [{PtX_2L}_2] \xrightarrow{CO} trans-[PtX_2(CO)L]$$
 (6)

originally suggested to explain the apparent nonappearance of the *trans* isomer.¹ Mechanisms of this type have also been shown to be involved in isomerisations of the complexes [PtXPh(CO)L].5,13 Interestingly, isomerisations of these latter complexes are inhibited, not catalysed, by CO, and treatment by L or X has no effect other than to displace CO from the molecules.

Other routes for the isomerisations include a geometry change of the three-co-ordinate (or solvated) intermediate, following CO elimination. All these mechanisms have recently been compared and discussed.¹⁵

(d) Photochemical Reactions.—Ultraviolet irradiation not only accelerated the trans to cis conversions, but promoted the conversions of cis-[PtCl₂(CO)L] into mixtures of the isomers. Prolonged irradiation at 0 °C of deuteriochloroform solutions of pure cis- or trans- $[PtCl_{2}(CO)(PMePh_{2})]$ resulted, after *ca*. 5 h, in the same equilibrium point at 68% trans. It is possible that a direct geometry change via a pseudotetrahedral intermediate is responsible for the conversions,¹⁵ but a photochemical elimination of CO is an alternative route.

Formation of some of the dimers, $[{PtX_2L}_2]$, accompanies the conversions. Some support for the latter hypothesis may be drawn from an experiment in which mixed solutions of cis-[PtCl₂(CO)(PMePh₂)] (39 mg) and $[Pt_2Cl_4(PMe_2Ph)_2]$ (32 mg) were irradiated. The experiment was interrupted before equilibrium was reached, when cis-[PtCl₂(CO)(PMePh₂)] was still the major COcontaining component. More than twice as much trans-[PtCl₂(CO)(PMe₂Ph)] was present in solution than trans- $[PtCl_2(CO)(PMePh_2)]$ (from ³¹P n.m.r. measurements). A small amount of cis-[PtCl₂(CO)(PMe₂Ph)], and some of the dimers [Pt₂Cl₄(PMePh₂)₂] and [Pt₂Cl₄(PMePh₂)-(PMe,Ph)] had also been formed. Since the transcarbonyl complexes readily lose CO, even without irradiation, this experiment cannot be taken as proof of a light-induced CO elimination as the mechanism, but the product ratios are suggestive. No reaction took place between cis-[PtCl₂(CO)(PMePh₂)] and [Pt₂Cl₄(PMe₂Ph)₂] in the dark at 0 °C.

Interestingly, the conversions of the yellow trans-[PtCl₂(CO)L] to the colourless *cis* isomers are accelerated by daylight. Though the effect is quite small in the absence of direct sunlight, it necessitates storing the trans complexes in the dark.

EXPERIMENTAL

Solution i.r. spectra were recorded on a Perkin-Elmer 577 spectrophotometer, using 0.5-mm path length NaCl solution cells. ³¹P and ¹³C n.m.r. spectra were recorded in CDCl_a on a Varian XL-100 spectrometer operating in the F.T. mode. Ultraviolet irradiations were performed in Pyrex apparatus using a Hanovia medium-pressure mercury lamp. Carbon-13 monoxide (90%) was obtained from Merke, Sharp and Dohm.

Most samples were prepared in solution (see text) and their identity and purity checked by spectroscopic methods (Table). A few were isolated as pure solids. The preparation below is typical.

trans-[PtCl₂(CO)(AsMePh₂)].--The compound [Pt₂Cl₄(As- $MePh_2$ [(0.121 g) was dissolved in CH_2Cl_2 (10 cm³) and the solution was cooled to -40 °C. Carbon monoxide gas was bubbled through the solution for 5 min. An aliquot of the solution was monitored by i.r. spectroscopy, which revealed the presence of the trans isomer only of [PtCl₂(CO)(AsMePh₂] $[v(CO) ca. 2 130 cm^{-1}]$. The remaining solution was cooled to -50 °C, and the solvent was removed by evacuating at this temperature. Yellow, solid, trans-[PtCl2-(CO)(AsMePh₂)] resulted [ν (CO) in CHCl₃ was 2 134 cm⁻¹].

After keeping the sample for 21 d. at -10 °C, no decomposition or isomerisation was detected. Several attempts at C and H analysis produced results in consistently good agreement with $[Pt_2Cl_4(AsMePh_2)_2]$.

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