

STERIC EFFECTS OF PHOSPHORUS LIGANDS: *cis/trans* DISTRIBUTIONS OF $W(CO)_4L_2$ PRODUCTS

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(Received February 5th, 1980)

Summary

The *cis/trans* product distribution in the reaction of phosphines with $W(CO)_4(Me_2N(CH_2)_3NMe_2)$ depends on ligand size; the smaller the ligand the greater the *cis/trans* ratio. The ratios range from 2.9 to 0 for PPh_2Me and $P(o-Tol)_3$, respectively.

Introduction

The importance of phosphorus ligand steric effects in transition metal complexes has received considerable attention during recent years [1–6]. In this report, the effect of ligand size on the *cis/trans* product distributions of reaction 1 is unambiguously demonstrated (tmpa = *N,N,N',N'*-tetramethyl-1,3-propanediamine and L = a phosphorus ligand).



To observe steric effects only, it is necessary to choose ligands which have very similar electronic properties since both σ -donating and π -accepting abilities of ligands can affect stereochemistries [7–9]. The ligand series PPh_2Me , PPh_2Et , PPh_2i-Pr and PPh_2t-Bu provides for increasing size (cone angles of 136° , 140° , 150° and 157° , respectively) [1], while maintaining nearly identical electronic properties. Values for the $\nu(CO)$ A^1 mode of $Ni(CO)_3L$ complexes, used by Tolman as the electronic parameter for phosphorus ligands [1], are within experimental error for PPh_2Me (2067.0 cm^{-1}) and PPh_2Et (2066.7 cm^{-1}) [10]. Although corresponding values are not available for PPh_2i-Pr and PPh_2t-Bu , the electronic similarity of these ligands to PPh_2Et can be seen from other data. Grim and coworkers have found $^1J(^{31}P-^{183}W)$ values and $\nu(CO)$ bands in $W(CO)_5L$ complexes to be indicative of electronic properties [11]. Coupling constant values for the latter three ligands are identical at 240 Hz and the $\nu(CO)$ frequencies are extremely consistent [12].

Results and discussion

For the study reported here, a fivefold excess of each of the ligands PPh_2Me , PPh_2Et , $\text{PPh}_2\text{i-Pr}$, $\text{PPh}_2\text{t-Bu}$, PPh_3 , $\text{P}(p\text{-Tol})_3$ and $\text{P}(o\text{-Tol})_3$ was allowed to react with $\text{W}(\text{CO})_4\text{tmpa}$ at 40°C . Completion of reaction was determined by the lack of intense $\nu(\text{CO})$ bands characteristic of starting material complex. As reported previously [13], no evidence for *cis/trans* interconversion was observed, even after two weeks at 40°C . Since selective loss of one isomer during purification was a concern, ^{31}P NMR spectra were obtained directly from untreated product solutions. No difficulty was encountered in identifying product peaks. In fact, the spectra of PPh_2Me and PPh_2Et displayed no discernable absorptions other than free ligand, *cis* and *trans* complexes and their ^{183}W satellite peaks. The percentage of *cis* and *trans* complexes, determined by integration of the ^{31}P spectra, are listed in Table 1. Also included are results obtained in a similar manner for PPh_3 , $\text{P}(p\text{-Tol})_3$ and $\text{P}(o\text{-Tol})_3$.

The results for PPh_2Me , PPh_2Et , $\text{PPh}_2\text{i-Pr}$, and $\text{PPh}_2\text{t-Bu}$ unambiguously indicate the general trend of increasing ligand size producing increased amounts of *trans* product. This is the first known report of its kind. The effect of ethyl replacement for methyl is particularly dramatic. The reason for experimentally identical results for PPh_2Et and $\text{PPh}_2\text{i-Pr}$ is unclear. That apparent enigma is being investigated further.

Results for PPh_3 , $\text{P}(p\text{-Tol})_3$ and $\text{P}(o\text{-Tol})_3$ provide further verification of steric effects. Although these three ligands are not electronically identical, as reflected in the slightly different *cis/trans* distributions for the sterically similar ligands PPh_3 and $\text{P}(p\text{-Tol})_3$ (the cone angle for both is 145° [1]), the extremely large $\text{P}(o\text{-Tol})_3$ ligand (cone angle 194° [1]) precluded any *cis* complex formation.

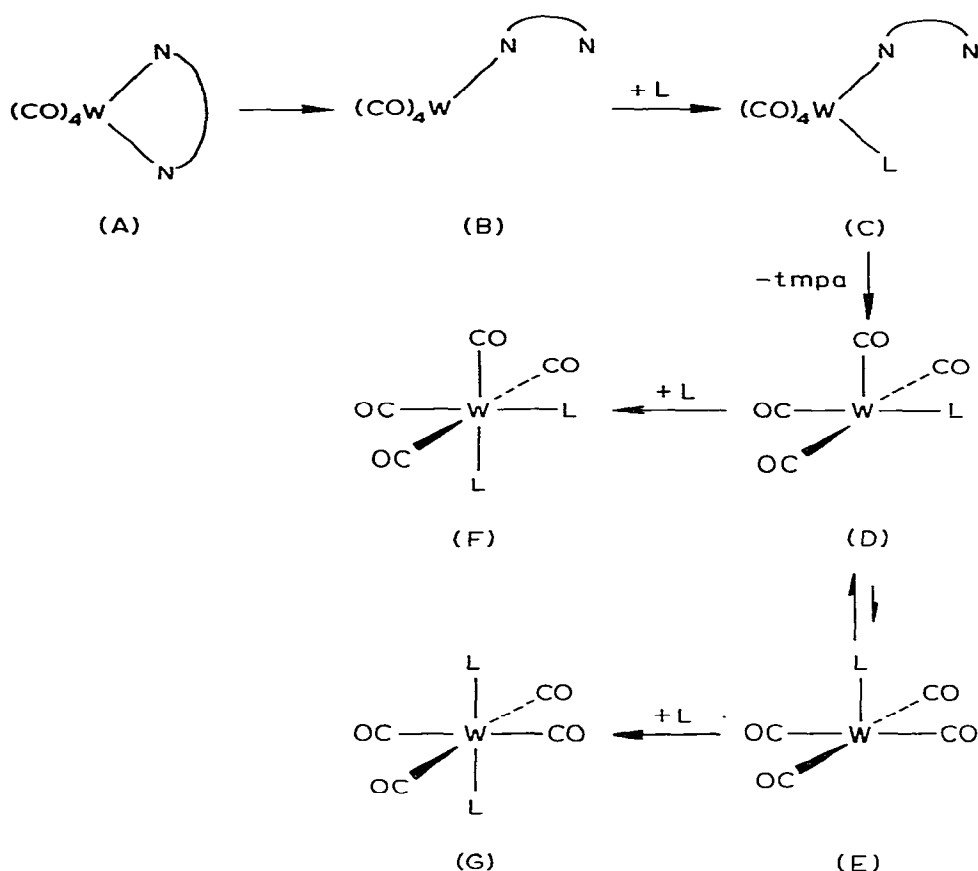
A composite reaction mechanism is depicted below. From observed rate data, the reaction has been shown to proceed through the steps $\text{A} \rightarrow \text{B} \rightarrow \text{C} \rightarrow \text{products}$ [13]. Precedence for a square pyramidal intermediate (e.g., D and/or E) comes from matrix isolation studies [14], while Darensbourg et al. [13] have suggested the equilibrium between D and E which produces *cis* (F) and *trans* (G) products, respectively.

TABLE 1

 ^{31}P CHEMICAL SHIFTS ^a AND *cis/trans* PRODUCT DISTRIBUTION OF $\text{W}(\text{CO})_4\text{L}_2$

Ligand	<i>cis</i>		<i>trans</i>	
	δ (ppm)	%	δ (ppm)	%
PPh_2Me	-6.44	74	-0.31	16
PPh_2Et	8.93	38	15.41	62
$\text{PPh}_2\text{i-Pr}$	22.01	38	29.05	62
$\text{PPh}_2\text{t-Bu}$	31.68	21	43.54	79
PPh_3	19.83	23	24.82	77
$\text{P}(p\text{-Tol})_3$	17.30	21	22.25	79
$\text{P}(o\text{-Tol})_3$	—	0	32.83	100

^a Referenced to 85% H_3PO_4 ; positive values are downfield.



Inclusion of the equilibrium step was based on two studies reported by the latter authors. The reaction of *cis*-(C₅H₁₀NH)(PPh₃)W(CO)₄ with ¹³CO resulted in stereospecific enrichment to yield *cis*-(¹³CO)(PPh₃)W(CO)₄, whereas reaction 1 with L = PPh₃ provided approximately 80% *trans*-(PPh₃)₂W(CO)₄ [13]. The results were rationalized in terms of the "site preference" model of Atwood and Brown [7]. Thus, the basal intermediate (D) is believed to be more stable thermodynamically, but steric interactions between the coordinated PPh₃ and a second, incoming PPh₃ ligand result in preferential attack of the apical intermediate (E) [13]. The results reported here are consistent with this hypothesis.

Experimental

General

All reactions, distillations and transfers were carried out in a nitrogen atmosphere. Benzene was stored over 4A molecular sieves and purged with nitrogen prior to use. *N,N,N',N'*-tetramethyl-1,3-propanediamine (tmpa) was purchased from Aldrich Chemical Co., Inc., and distilled from KOH just prior to use. Tungsten hexacarbonyl, *t*-butyldiphenylphosphine, tri-*o*-tolylphosphine, tri-*p*-tolylphosphine, and triphenylphosphine were purchased from Strem Chemicals, Inc., and used as received, Methylidiphenylphosphine, ethyldiphenylphosphine,

and *i*-propyldiphenylphosphine were synthesized by literature procedures [15]. $\text{W(CO)}_4\text{tmpa}$ was synthesized using the method reported by Dobson and Faber [16].

IR spectra were recorded using a Beckman 4250 spectrophotometer. The ^{31}P NMR spectra were recorded at 60.8 MHz using an NT-150 instrument operating with a 90° pulse angle and 12 mm probe. Trimethylphosphate was used as an external standard and deuteriobenzene (20% by volume) as the internal lock. NMR spectra were recorded directly on the reaction solutions. Isomer distributions were obtained by integration of the ^{31}P resonances. Chemical shifts, presented in Table 1, are reported relative to 85% H_3PO_4 with downfield shifts assigned positive values. The downfield signals, which also displayed the larger $^1J(\text{WP})$ values, were assigned to the *trans* complexes [17–21]. The IR spectra were also consistent with these assignments.

Reactions

Benzene was used as a solvent for the reactions involving the solid ligands, PPh_3 , $\text{P}(p\text{-Tol})_3$, and $\text{P}(o\text{-Tol})_3$. The liquid alkylidiphenylphosphine ligands were allowed to react with $\text{W(CO)}_4\text{tmpa}$ in the absence of solvent. For the solid ligands, reactions were carried out using ca. 0.045 g and 0.30 g of $\text{W(CO)}_4\text{tmpa}$ and phosphine, respectively, in 20 ml of benzene. All reactions involving the liquid ligands utilized ca. 0.16 g and 0.85 g of $\text{W(CO)}_4\text{tmpa}$ and phosphine, respectively. For example, 0.0467 g (0.110 mmol) of $\text{W(CO)}_4\text{tmpa}$ with 0.3225 g (1.059 mmol) of $\text{P}(o\text{-Tol})_3$ and 0.1661 g (0.390 mmol) of $\text{W(CO)}_4\text{tmpa}$ with 0.9036 g (4.218 mmol) of PPh_2Et were reacted. All reactions were carried out at 40°C for 48 h. After the reaction period, the IR spectra showed $\nu(\text{CO})$ bands characteristic of only *trans* and *cis* $\text{W(CO)}_4\text{L}_2$ products.

Acknowledgments

Acknowledgment is made to the donors of the Petroleum Research Fund, administered by the American Chemical Society, the Ball State University Faculty Summer Research Grant program, the Purdue University Biochemical Magnetic Resonance Laboratory, and National Institutes of Health Grant No. RR 01077 for support of this research.

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