## **Synthesis, Structure, and Absolute Configuration of Chiral, Pseudooctahedral Perfluoroalkyl-Substituted Dlmethylphosphonate Complexes**

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The substitution-labile complexes  $\eta^5$ -CpCo(X)(PPh<sub>2</sub>N(R)C\*H(Me)Ph)(I), 6(a,b)( $\alpha,\beta$ ), react with trimethyl<br>phosphite to afford  $\eta^5$ -CpCo(X)(PPh<sub>2</sub>N(R)C\*H(Me)Ph)(P(O)(OMe)<sub>2</sub>), 7(a,b) $\alpha$ , and  $\eta^5$ -CpCo(X)(P(OMe)<sub>3</sub>  $R = H$  and requires intramolecular P=O- $H - N$  hydrogen bonding, which develops early in the transition state to "anchor" the aminophosphine ligand. The chiral, Co-epimeric Arbuzov products were separated and characterized. The absolute configuration of  $(+)_{436}$ -7ba  $\infty$  asset determined crystallographically.  $(+)_{436}$ -7b experiment characterized. The absolute comparation of  $(\frac{1}{36} + \frac{1}{160})$  and was determined crystallizes in the monoclinic system  $P2_1$  with  $a = 11.0941$  (20) Å,  $b = 16.9200$  (22) Å,  $c = 16.927$  (3) Å,  $\beta = 91.241$  (15 and was found to contain two independent molecules in the asymmetric unit. Absolute configurations of  $(-)_{436}$ -7ba,  $(-)_{436}$ -7aa, and  $(+)_{436}$ -7aa were assigned on the basis of chiroptical evidence. P=O- $\cdot$ -H-N hydrog  $n-C_3F_7$  in the solid state. Nuclear Overhauser difference (NOED) spectra show that the solid-state conformation persists in solution.

## **Introduction**

Recently we reported<sup>3</sup> evidence demonstrating that a chiral cobalt atom in an asymmetric pseudooctahedral<sup>4,5</sup> environment *can* function **as an** effective stereogenic center for the formation of P-chiral phosphonate  $(!P(O)R(OMe)^{1-})$ complexes with good optical yields (de  $\geq 80\%$ , cf. (Scheme I). Thus, substitution of iodide in the prochiral complex 1 with 1 equiv of  $PR(OMe)_2$  gives initially<sup>6-12</sup> the chiral phosphonite complex *(R,Sc,)-2,* which subsequently dealkylates via an Arbuzov-like rearrangement<sup>6,8,9,12-18</sup> to afford the phosphonate 3 as a diastereomeric mixture  $R, S_{\text{Co}}; R, S_{\text{P}}$ . Phosphonato-Co(III) complexes have also been prepared by direct oxidative addition of dialkyl phosphonates  $(HP(O)(OR)_2)$  to  $Co(I)$  substrates<sup>19</sup> and by halide displacement with the corresponding sodium **salts.2o** 

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Although the Co- and P-chiral phosphonate products are sufficiently configurationally stable in solution to allow reliable determinations of optical yields, slow epimerization at the cobalt center does occur. We have not examined the Co epimerization of 3 in detail. However, there is ample literature precedent establishing both a dominant dissociative activation mode for loss of stereochemistry at the metal in piano stool complexes $^{21,22}$  and the lability of cobalt-halide bonds.<sup>10-12,23,24</sup> It was of interest then, in the context of our studies of optical induction from chiral transition-metal atoms, to prepare derivatives in which this mode of metal epimerization was blocked.<sup>13</sup> This report describes the preparation, characterization, crystal structure, and conformational analysis of the analogous phosphonate derivatives **7** in which iodide is replaced by a substitution-inert perfluoroalkyl group. $^{13,25}$ 

## **Results and Discussion**

**Preparation and Characterization of the Phosphonato Compounds 7a,b.** Preferential substitution of labile CO in  $\eta^5$ -CpCo(X)(CO)(I) (X = perfluoroalkyl) by P-donor ligands<sup>13,25,26</sup> occurs even in the absence of halide  $acceptors<sup>27</sup>$  and represents a more general synthetic route to the required substrates  $6(a,b)(\alpha,\beta)$  than oxidative addition to phosphine-substituted  $\eta^5$ -CpCo(CO)(P-donor) Co(I) complexes.<sup>28</sup> The substitution procedure (cf. The substitution procedure (cf. Scheme II) reported by Brunner et al.<sup>26</sup> gave good yields of the known complexes  $6a\beta$ ,  $6b\alpha$ , and  $6b\beta$  as well as the new complex **6aa.** Resolution of the resulting ca. **1:l**  mixture of diastereomeric products in the case of  $6(a,b)\alpha$ was conveniently achieved by fractional crystallization

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from methylene chloride/ hexane. The diastereomers of  $6$ b $\alpha$  show distinctive <sup>1</sup>H NMR spectra<sup>26</sup> in CDCl<sub>3</sub> (less soluble diastereomer, Cp, **4.98** ppm, s; CMe, 0.92 ppm, d, J = **6.8** Hz; more soluble diastereomer, Cp, **4.93** ppm, s; CMe,  $1.34$  ppm,  $d, J = 6.8$  Hz.). The chiral cobalt center in  $(R, S_{\text{Co}})$ -6(a,b)( $\alpha, \beta$ ) is stereolabile<sup>26</sup> at temperatures **required** for further reaction according to Scheme **II;** hence, diastereomeric mixtures were used for all subsequent reactions.

The deep-purple perfluoroalkyl complexes  $6(a,b)(\alpha,\beta)$ exhibit two parallel reaction channels on treatment with trimethyl phosphite. The sec-aminophosphine $^{29}$  complexes  $6(a,b)\alpha$  react predominantly via iodide displacement/ Arbuzov dealkylation to give orange phosphonato products  $7(a,b)\alpha$  (cf. discussion below), while the tert-aminophosphine complexes  $6(a,b)\beta$  prefer to substitute a more labile aminophosphine ligand. The purple perfluoroalkyl phosphite complexes **8a,b** isolated in the latter case were characterized spectroscopically (cf. Tables I1 and 111) and by direct preparation from the parent CO complexes **4a,b**  (cf. Scheme 11).

Attempts to prepare dimethylphosphonato complexes containing the tert-aminophosphine via a two-step sequence involving iodide substitution followed by dealkylation failed. NMR experiments showed that direct reaction of **8a** with **58** in benzene at **60** "C or acetone at 25 °C results in slow, incomplete phosphite substitution to give  $6(a,b)\beta$ . Directed halide substitutions also failed. Treatment of  $8b$  with  $AgBF<sub>4</sub>$  in acetone gave a yellow precipitate of AgI and the expected labile solvent complex, which was characterized by its 'H *NMR* **spectrum** (Cp, **5.63**  ppm, s;  $P(OMe)_3$ , 3.93 ppm; d,  $J = 10.7$  Hz). In situ<br>treatment of the acetone complex with aminophosphine  $5\beta$  in an NMR tube afforded  $9b\beta$ , presumably the same intermediate required18 in the Arbuzov sequence leading to  $7b\beta$ . The structure of  $9b\beta$  is supported by its <sup>1</sup>H NMR

<sup>(29)</sup> By analogy with the accepted nomenclature for amines, compound  $5\alpha$ , which has one remaining unsubstituted N-H, is referred to as primary, while  $5\beta$ , which has no remaining N-H groups, is tertiary.



<sup>a</sup> <sup>1</sup>H (300.1 MHz) NMR chemical shifts in ppm relative to internal TMS; <sup>19</sup>F (282.37 MHz) NMR chemical shifts relative to internal CFCl<sub>3</sub>; m = multiplet;  $s =$  singlet;  $d =$  doublet;  $u =$  unresolved; *J* values given in Hz; solvent = CDCl<sub>3</sub>.  $^b(^2J_{HP}, ^3J_{HH})$ .  $^c(^3J_{PH}, ^3J_{NHH}, ^3J_{HH})$ .  $^d$  AB multiplet,  $^4J_{FF}$ and  $v_{\text{FF}}$  unresolved.





<sup>a 13</sup>C (75.47 MHz) NMR chemical shifts in ppm relative to CDCl<sub>3</sub> = 77.0;  $d =$  doublet; nm = not measured; *J* values in Hz.

Table IV. Preparative Conditions for  $6(a,b)(\alpha,\beta)$  and  $8a,b$ 

compd	solvent	time, h	temp, °C	elution EtOAc:hex- ane	% yield
$6a\alpha$	CH <sub>2</sub> Cl <sub>2</sub>	2.0	20	1:5	95
$6a\beta$	CH <sub>2</sub> Cl <sub>2</sub>	14.0	20	1:2	95
6ba	CH,Cl,	2.0	20	1:15	92
$6b\beta$	CH <sub>2</sub> Cl <sub>2</sub>	0.25	20	1:2	95
8а	$CH_2Cl_2$	1.0	20	1:3	80
8Ь	CH <sub>2</sub> Cl <sub>2</sub>	1.0	20	2:5	63

spectrum, which shows signals for Cp  $(5.58 \text{ ppm}, \text{d}, \frac{3}{H} = 0.4 \text{ Hz})$ , P(OMe)<sub>3</sub>  $(3.92 \text{ ppm}, \text{d}, \frac{3}{H} = 10.7 \text{ Hz})$ , and  $-0.4$  **112**),  $\Gamma$  (OMe)<sub>3</sub> (3.32 ppm, d,  $\sigma$ )<sub>HP</sub> =  $10.7$  Hz), and coordinated 5*β* (PPh 7.5 ppm, m; NMe 2.40 ppm, d,  $\sigma$ )<sub>PH</sub> = 6.5 Hz; CMe 1.63 ppm, d,  $\sigma$ <sup>3</sup> $J$  = 6.9 Hz; CH 5.15 ppm, = 6.5 Hz; CMe 1.63 ppm, d,  $^{3}J = 6.9$  Hz; CH 5.15 ppm, m). Two Co epimers of  $9b\beta$  are expected. However, the

'H NMR spectrum was consistent with the presence of a single diastereomer. Presumably the Co configuration is labile,  $^{26,20}$  and the equilibrium strongly favors<sup>13</sup> one epimer. Treatment of  $9b\beta$  with iodide (as NaI or  $(n-C_4H_9)_4$ NI) did not result in dealkylation since no methyl iodide was de**tected** via lH NMR. Traces of the simple Co-substitution product **8b** but no coordinated **58** were observed.

Unlike the tert-aminophosphine derivatives, complexes  $6(a,b)\alpha$  react smoothly with trimethyl phosphite in benzene to afford phosphonato producta via an Arbuzovlike rearrangement. The orange phosphonato products  $7(a,b)\alpha$  form as a ca.  $50/50$  mixture of diastereomers

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**Figure 1.** Circular dichroism spectra of  $(-)_{436}$ -7b $\alpha$  (solid line) and  $(+)$ <sub>436</sub>-7b $\alpha$  (dotted line) in toluene.

differing only in the configuration at the cobalt atom. Separation of the diastereomeric mixture was readily accomplished using radial thick layer chromatography followed by crystallization from methylene chloride/hexane. The Co-epimeric diastereomers are configurationally stable in solution. A sample of diaster exercically pure  $7b\alpha$  was heated at 60  $\rm{^{\circ}C}$  for several weeks in a sealed NMR tube without any measurable isomerization. The higher  $R_f$  zone for both  $7a\alpha$  and  $7b\alpha$  was determined to be levorotatory  $f(x)_{436} = -2192$  and  $-2090$ , respectively), while the lower  $R_t$  zones were dextrorotatory ( $\alpha$ <sub>1436</sub> = 2040 and 3095, respectively). Their structures are supported by  ${}^{1}H$ ,  ${}^{19}F$ , and <sup>13</sup>C NMR data. Distinctive pairs of diastereotopic methoxy doublets are observed in the 'H and 13C NMR spectra for each Co epimer (cf. Tables II and III). The  $n-C_3F_7$  derivatives also show well-resolved <sup>19</sup>F AB patterns<sup>31</sup> for the diastereotopic  $\alpha$ -CF<sub>2</sub> and  $\beta$ -CF<sub>2</sub> groups<sup>32</sup> (cf. Table II). plings of the seven-spin ABMNX<sub>3</sub> system were not resolved. Characteristically, the <sup>1</sup>H NMR chemical shift parameters correlate with the chromatographic  $R_i$  value **as** well **as** the chiroptical properties (cf. discussion Ibelow) for each pair of diastereomers examined in this study. *As*  found previously for other pairs of related Co-epimeric phoephonato complexes3 the 'H NMR chemical shift of C\*-Me in the higher  $R_f$  (-)<sub>436</sub>-7a,b $\alpha$  is in each case less shielded than its lower  $R_f$   $(+)$ <sub>436</sub> diastereomer. The expected3 **P-0** stretch at ca. **1150** cm-' was obscured by strong CF absorptions. However, the expected  ${}^4J_{FF}$  and much smaller  ${}^3J_{FF}$  cou-

Absolute Configurations of  $(\pm)_{436}$ -7(a,b) $\alpha$ . The absolute configurations of  $(\pm)_{436}$ -7(a,b) $\alpha$  were assigned on the **basii** of circular dichroism (CD) spectra and, in the *case*  of (+)436-7ba, crystallographic evidence. Figures **1** and **2**  show that the CD spectra of the high- and low- $R_i$  diastereomers of  $7(a,b)\alpha$  are quasi-mirror images as expected for piano stool transition-metal epimer^.^\*^^^ **A** single-crystal X-ray structure of low- $R_f$  (+)<sub>436</sub>-7b $\alpha$  allows unambiguous assignment<sup>34</sup> of its absolute configuration at the chiral cobalt center (cf. Figures **3** and **4) as** well **as** confirmation of the absolute configuration at carbon which was known to be S. The absolute configurations for  $(+)$ <sub>436</sub>-7b $\alpha$  and  $(+)$ <sub>436</sub>-7ba' were determined using the NRCVAX utility



**Figure 2.** Circular dichroism spectra of  $(-)$ <sub>436</sub>-7a $\alpha$  (solid line) and  $(+)$ <sub>436</sub>-7a $\alpha$  (dotted line) in toluene.



**Figure 3.** Molecular geometry and absolute configuration of  $(+)$ <sub>436</sub>-7**b** $\alpha$ .



**Figure 4.** Molecular geometry and absolute configuration of  $(+)_{436}$ -7**b** $\alpha'$ .

BIVOET<sup>35</sup> in which a solved structure is used to identify the Friedel pairs most affected by anomolous dispersion and

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**Table V. Nuclear Overhauser Difference Data"** 



<sup>a</sup> % reduction of irradiated peak in parentheses; NOE enhancement in %  $\pm$  0.5. <sup>b</sup> Assigned to o-phenyl protons of PPh<sub>2</sub>. <sup>c</sup> Low field. <sup>d</sup> High field. <sup>e</sup>Corrected for overlapping C\*H signal. <sup>*f*</sup> Averaged value. <sup>g</sup>C\*H also irradiated.





Parameters reported for both independent molecules in the unit cell.

to compare the calculated relative magnitudes of  $F(+++)$ and  $F(--)$  with those measured. The hand of the working model was confirmed. The probability of this assignment being incorrect is was calculated<sup>35</sup> to be  $2 \times 10^{-8}$ .

Consideration of the chiral complex  $(+)$ <sub>436</sub>-7(a,b) $\alpha$  as a pseudotetrahedral case with  $\eta^5$ -Cp effectively occupying one coordination site and use of the modified Cahn-Ingold-Prelog rules<sup>36-38</sup> with the ligand priority series  $Cp$ 





" Parameters reported for both independent molecules in the unit cell.

 $P(0)(OMe)<sub>2</sub>>(S)-(-)$ - $PPh<sub>2</sub>NHCH(Me)Ph > X$  fixes its absolute configuration as  $S_{\text{Co}}$ ;  $S_{\text{C}}$ . CD spectra confirm that  $(-)$ <sub>436</sub>-7(a,b) $\alpha$  is its Co epimer with absolute configuration  $R_{\text{Co}}$ ; S<sub>c</sub>. Since the CD spectra of high- $R_f$  (-)<sub>436</sub>-7a $\alpha$  and high- $R_f$  (-)<sub>436</sub>-7b $\alpha$  are isomorphous, the former can be assigned the absolute configuration  $R_{\text{Co}}$ ;  $S_{\text{C}}$ . Similarly, CD data allow assignment of the low- $R_f$  (+)<sub>436</sub>-7a $\alpha$  diastereomer as  $S_{Co}$ ;  $S_{C}$ .

**Crystal Structure and Solid-state Conformation.**  Suitable crystals of  $(+)$ <sub>436</sub>-7ba were obtained by slow cooling of methylene chloride/ hexane solutions. The solid-state structure, determined by X-ray techniques (cf. Experimental Section for details), revealed two crystallographically independent molecules in the asymmetric unit  $(Z = 4)$ . Positional parameters and selected bond lengths and angles are given in Tables VI and VII. Representational drawings, **shown** in Figure 3 and **4,** define piano stool structures with approximate **90°** bond angles about a pseudooctahedral cobalt. The coordination sphere consists of an unexceptional  $\eta^5$ -Cp ring which occupies three *facial* 

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Table VIII. Summary of **CrYstalloeraDhic** Data **for** 

$(+)_{436}$ -7b $\alpha^a$				
formula М	$C_{30}H_{31}CoNO_3P_2F_7$ 707.45			
crystal habit	orange prism			
crystal size, mm	$0.3 \times 0.1 \times 0.03$			
crystal system	monoclinic			
space group	P2.			
cell constants				
a, Å	11.0941 (20)			
b, A	16.9200 (22)			
c, Å	16.927 (3)			
$\beta$ , deg	91.241 (15)			
V, A <sup>3</sup>	3176.6 (9)			
z	4			
$D_x$ , g cm <sup>-3</sup>	1.479			
F(000)	1447.80			
scan type	<i>0120</i>			
abs correction	none			
$2\theta_{\text{max}}$ , deg	44.8			
h,k,i ranges	-11,11,0; 17,0,18			
refins measd	5010			
unique reflns	4957			
obsd reflns $(I_{\text{net}} > 2.5\sigma I_{\text{net}})$	2042			
$R_{\rm f}$ ( <i>I</i> > 2.5 $\sigma(I)$ )	0.084			
$R_w$ $(I > 2.5\sigma(I))$	0.051			
GOF	1.97			
no. of params	360			
largest peak in final diff map, $e^{\pi} A^{-3}$	0.450			
deepest hole in final diff map, $e^- A^{-3}$	$-0.420$			

 ${}^aR_f = \sum (F_o - F_c)/\sum F_o$ ;  $R_w = (\sum (w(F_o - F_w)^2)/\sum (wF_o^2))^{1/2}$ ; GOF =  $(\sum (w(F_o - F_w)^2)/(no.$  reflns - no params))<sup>1/2</sup>.



**Figure 5.** Comparison of conformations for  $(+)$ <sub>436</sub>-7b $\alpha$  and  $(+)_{436}$ -7b $\alpha'$ .

coordination sites. A P-bound dimethylphosphonato<sup>20</sup> group, an aminophosphine, and a perfluoropropyl group fill the remaining coordination positions. Except for **a**  minor change in orientation of the  $\eta^5$ -Cp ring (cf. Figure **51,** the fundamental Structure of the two independent molecules is identical. Both the  $P(N)-Co-P(O)$  and the P(N)-Co-C bond angles open slightly to ca. **95'** in order to accommodate the steric requirement of the aminophosphine ligand. No significant differences in C-P bond lengths were apparent for the aminophosphine and dimethylphosphonate ligands. As found previously,<sup>3,23,39</sup> the geometry of the dimethylphosphonate ligand is distorted from tetrahedral by a closing of the 0-P-0 angle to **102'**  and a concomitant opening of the Co-P=0 angle to 121

Strong intramolecular N-H--O=P hydrogen bonding is evident in the solid-state structure. The N-0 bond

distance of 2.843 Å for  $(+)_{436}$ -7b $\alpha$  is markedly less than the value of **3.0516 A** typically **found for** strongly hydrogenbonded O<sub>"H-N</sub> systems.<sup>40</sup> The secondary hydrogenbonding interaction controls the solid-state conformation and resulta in a distorted six-membered **(Co-P-0.-H-**N-P) boat. As found previously<sup>3,40</sup> for related complexes containing the aminophosphine ligand *Sa* and an accessible oxygen-donor site, the  $\eta^5$ -Cp ring assumes a pseudoequatorial position bisecting both the PPh<sub>2</sub> and P(OMe)<sub>2</sub> groups. This conformation places the ortho protons of both phenyl groups proximal (ca. 2.3  $\AA$  to the  $\eta^5$ -Cp ring hydrogens) and, as a further consequence, forces the  $\eta^5$ -Cp ring and  $N(R)C^*$  into a anti rather than the preferred<sup>41</sup> gauche relationship. The planes of the pseudoequatorial PPh and  $\eta^5$  groups are also orthogonal, allowing for an attractive  $\beta$ -phenyl effect.<sup>24,42</sup> Since the alternate conformation which places a perfluoroalkyl group pseudoequatorial and  $\eta^5$ -Cp pseudoaxial would also permit the preferred gauche  $\eta^5$ -Cp/N(R)C\* arrangement<sup>41</sup> as well as the stabilizing  $\beta$ -phenyl effect, we are forced to conclude that a further destabilizing steric interaction must be encountered. Recalling that the cobalt atom is nearly octahedral<sup>5</sup> with bond angles close to 90° rather than 109°, we suggest that the latter conformation would force destabilizing gauche interactions between the perfluoroalkyl group and both PPh<sub>2</sub> phenyl groups.

Solution Conformation. **'H** nuclear Overhauser effect difference (NOED) spectra<sup>43</sup> were measured in order to eatablish whether the solid-state conformation was retained in solution for  $(+)$ <sub>436</sub>-7ba. Comparative NOED data were also obtained for  $(-)$ <sub>436</sub>-7ba,  $(-)$ <sub>436</sub>-7aa, and  $(+)$ <sub>436</sub>-7aa. Table **V** displays the quantitative NOE enhancements measured. Both Co epimers of the  $n-C_3F_7$  (7b $\alpha$ ) and CF<sub>3</sub>  $(7a\alpha)$  series show parallel effects; hence, we propose that their solution conformations are *similar.* Irradiation of the Cp **signal** gives large positive enhancements *(ca.* **4-7%)** for the ortho protons of the  $PPh<sub>2</sub>$  groups consistent with a solution conformation in which a pseudoequatorial Cp is gauche to both rings **as** found in the solid state of  $(+)$ <sub>436</sub>-7b $\alpha$ . Positive OMe  $\rightarrow$  Cp NOE enhancements are **also** expected between the Cp and OMe groups since they **also** define a simiiar gauche relationship. However, Table **V** shows that the measured effects are very small *(ca.* **51%)**  and cannot be considered diagnostic. Presumably the increased flexibility associated with the  $Me-O-P$  link increases the average distance with respect to the Cp ring and results in very low enhancements. The NOED evidence strongly supports the premise that the solid-state conformation of  $(+)$ <sub>436</sub>-7b $\alpha$  is retained in solution. If the alternate conformation in which Cp is pseudoaxial were significantly populated, NOE enhancements would be expected between  $Cp$  and only one  $PPh<sub>2</sub>$  o-phenyl. The NOED results do not eliminate the possibility of free rotation about the Co-P(N) bond. However, we feel this is highly unlikely. Confirmation of this fact derives from an IR study which demonstrated that the  $\nu_{\text{NH}}$  is not affected by dilution, showing that hydrogen bonding persists in solution.

## Experimental Section

Reagents **and** Methods. *All* **manipulations were performed under a nitrogen atmosphere using standard Schlenk techniques.** 

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Nitrogen gas was purified by *passing* it through a **series** of columns containing activated molecular sieves **(3 A),** granular phosphorus pentoxide, and DEOX catalyst **(100** "C). Toluene, benzene, and ether were distilled from blue Solutions of sodium benzophenone ketyl. Methylene chloride was freshly distilled from  $P_4O_{10}$ . Acetone and ethyl acetate were distilled from activated **3-1**  molecular sieves. The progress of reactions was monitored by analytical thin-layer chromatography (precoated TLC plates, **silica**  gel **F-254,** Merck). NMR spectra were recorded on a General corded as thin films on a Mattson Polaris FTIR instrument with 4-cm<sup>-1</sup> resolution. Optical rotation measurements were determined<br>in toluene (ca. 1 mg/mL) in a 1-cm path length cell using a Perkin-Elmer Model 241 polarimeter. CD spectra were determined in toluene *(ca.* **1** *mg/mL)* on a **Jasco** J **40** A apparatus using a 0.1-cm path length cell. Melting points were determined in sealed capillaries and are uncorrected. Elemental analyses were performed by Guelph Chemical Laboratories, Inc. Chromatographic separations were carried out using a Chromatotron (Harrison Associates) with 1–2-mm-thick (silica gel<sub>eo</sub>)PF<sub>254</sub> (Merck) adsorbant. The compounds  $\eta^b$ -CpCo(CO)(I<sub>2</sub>),<sup>44</sup> (S)-(-)- $PPh_2NRC^*H(Me)Ph,$ <sup>45</sup>  $\eta^5$ -CpCo(X)(CO)(I) (4a,b<sup>46</sup>), and  $\eta^5$ - $CpCo(X)(PPh<sub>2</sub>NRCH(Me)Ph)(I)$  (6a $\beta$ , 6b $\alpha$ , and 6b $\beta^{26}$ ) were prepared using the established procedures. Commerical samples of trimethyl phosphite were distilled before **use.** Samples of CF31 and n-C3F71 (Aldrich) were used **as** received.

Proton NOED spectra were determined under steady-state conditions on a GE GN3OO-NB instrument at **25.0 0.1** "C. Data were collected using interleaved experiments of **16** or **32** transients cycled **12-16** times through the list of decoupling frequencies. In each experiment, the decoupler was gated on in continuous wave (CW) mode for **6 s** with sufficient attenuation to give an approximate **70-90%** reduction in intensity of the irradiated peak. A **60-8** delay preceded each frequency change. A set of four equilibrating scans was employed to equilibrate the spins prior to data acquisition. No relaxation delay was applied between successive scans of a given frequency. Difference spectra were obtained on **16K** or zero-filled **32K** data tables which had been digitally filtered with a **0.1-Hz** exponential line-broadening function. Quantitative data were obtained by integration.

Crystal Structure Determination. Intensity data were collected at room temperature on an Enraf-Nonius CAD4 (Mo **K,** radiation with graphite monochromater) automated diffractometer controlled by the NRCCAD program.<sup>47</sup> Data were collected with profile analysis and **8/28** scan mode.

Structure Solution and Refinement. Orange prisms of  $(+)_{436}$ -7ba were grown from dilute solutions of methylene chloride/hexane at -30 °C. Unit cell dimensions for  $(+)_{436}$ -7ba were derived from **24** reflections with a **20** range of **20.00-28.OOo.**  Systematic absences for the total data set were consistent with the space group  $P2_1$ . Structures were determined by direct methods and refined by successive cycles of least squares. All calculations were carried out using the NRCVAX suite of crystalstructure-solving programs.48 The phenyl rings were refined **as**  rigid groups to improve the reflections/parameters ratio. The absolute configuration for  $(+)_{436}$ -7b $\alpha$  was determined using the NRCVAX utility BIVOET.<sup>35</sup>

Carbonyl-Substitution Reactions of 4a,b. Preparation **of**   $6(a,b)(a,\beta)$  and  $8a,b$ . These complexes were prepared by treatment of 4a,b with the appropriate P-donor ligand at room temperature in methylene chloride solvent. In a typical prepa-<br>ration, 0.5934 g (1.954 mmol) of  $(S)$ -(-)-PPh<sub>2</sub>NHC\*H(Me)Ph was added to a solution of 0.7396 g  $(1.651 \text{ mmol})$  of  $\eta^5$ -CpCo(n- $C_3F_7$ (CO)(I), **4b**, in 20 mL of methylene chloride and stirred at ambient temperature for **2** h. A rapid evolution of CO was observed. Removal of volatiles under oil pump vacuum left a dark solid which was purified by radial thick layer chromatography (SiOz/ethyl acetakhexane **1:15)** to give **1.096 g (1.513** mmol,  $91.7\%$ ) of  $\eta^5$ -CpCo(n-C<sub>3</sub>F<sub>7</sub>)(PPh<sub>2</sub>NHC\*H(Me)Ph)(I), 6b $\alpha$ , as a ca. 1:1 mixture of  $R, S_{\text{Co}}$  epimers.  $6a\alpha$ ,  $6a\beta$ ,  $6b\beta$ ,  $8a$ , and  $8b$  were prepared in a similar fashion. Analytical and spectroscopic data are reported in Tables 1-111. Preparative details are summarized in Table IV.

Preparation of  $\eta^5$ -CpCo(X)(PNR\*)(P(O)(OMe)<sub>2</sub>),  $7(a,b)a$ . A solution of **0.8037** g **(1.1086** mmol) of 6b and **0.1478** g **(1.1918**  mmol) of  $P(OMe)_3$  in ca. 10 mL of benzene was prepared in a Carius tube and heated to **120** "C for **17** h. Removal of volatiles followed by filtration and radial thick layer chromatography (SiO,/EtOAc:hexane **15)** separated two orange zones. The lower  $R_t$  zone was identified as  $(\tilde{S}_{C_{02}}S_{C})$ -7b $\alpha$  (86.4 mg, 11.0%) on the basis of spectroscopic and crystallographic evidence. The higher R product **(95.5** mg, **12.2%)** was characterized **as** its Co epimer,  $(R_{Co}S_C)$ -7b $\alpha$  (cf. text).

Similarly, refluxing **0.1381** g **(0.2210** mmol) of 6a with **0.0316**  g **(0.2630** mmol) of P(OMe)3 in benzene for **4** h gave after chromatography  $(SiO<sub>2</sub>/EtOAc:hexane 1:1) (S<sub>Co</sub>,S<sub>C</sub>)$ -7aa  $(0.0321 g,$ **24.3%)** as a low- $R_f$  orange zone and  $(R_{Co}S_C)$ -7aa (0.0360 g, 27.2%) as a high- $R_i$  orange zone.

Reaction of  $6(a,b)\beta$  with Trimethyl Phosphite. In a typical reaction, a NMR tube was charged with equimolar amounts of  $6b\beta$  and  $P(\text{OMe})_3$  (ca. 0.01 mmol) dissolved in 0.4 mL of CD<sub>2</sub>Cl<sub>2</sub>. **'H** NMR spectra at ambient temperature showed that the major product was  $\eta^5$ -CpCo(n-C<sub>3</sub>F<sub>7</sub>)(P(OMe)<sub>3</sub>)(I), 8b. Similar results were obtained for  $6a\beta$ .

Directed Halide Substitution of  $n^5$ -CpCo(n -C<sub>3</sub>F<sub>7</sub>)(P- $(OMe)_3$  $(I), 8b.$  A solution of 0.0215 g  $(0.1101 \text{ mmol})$  of AgBF<sub>4</sub> in acetone was slowly added via syringe to a stirred solution of **0.0350 g (0.0643 mmol) of 8b in acetone. After 0.5 h, the resulting** yellow precipitate of AgI was centrifuged, and a solution of **0.0209**  g  $(0.0655 \text{ mmol})$  of  $6\beta$  in acetone was added. After stirring at 20 "C for **10** min, the solvent was removed under oil pump vacuum, leaving an orange red solid  $(0.0475 \text{ g}, 82.5\%)$  as  $[\eta^5$ -CpCo $(n$ - $C_3F_7$ )(PPh<sub>2</sub>N(Me)CH(Me)Ph)(P(OMe)<sub>3</sub>)]BF<sub>4</sub>, 9b $\beta$ .

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Registry **No.** 4b, **1212852-8;** *6aa* (isomer **1),142260-&4-2;** *6aa*  (isomer **2), 142394-44-3;** 6aj3 (isomer **l), 65073-89-4;** 6aj3 (isomer **2), 65137-29-3;** 6ba (isomer **l), 65073-91-8;** 6ba (isomer **2),**   $65137-27-1$ ;  $6b\beta$  (isomer 1),  $65073-92-9$ ;  $6b\beta$  (isomer 2),  $65137-26-0$ ;  $(S_{\text{Co}}S_{\text{C}})$ -7aa, 142260-87-5;  $(R_{\text{Co}}S_{\text{C}})$ -7aa, 142393-00-8;  $(S_{\text{Co}}S_{\text{C}})$ -7ba, **142293-83-2;** (Rc,,Rc)-7bj3, **142394-45-4;** 8a, **142260-85-3;** 8b, **142260-86-4;** 9bj3, **142260-89-7; (S)-(-)-PPh,NHC\*H(Me)Ph, 71554-93-3.** 

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Supplementary Material Available: Tables of atomic and thermal parameters, anisotropic temperature factors, and bond distances and angles **(13** pages). Ordering information is given on any current masthead page.