Group 4 Metal Olefin Polymerization Catalysts Stabilized by Bidentate O,P Ligands

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A series of bis(phosphanylphenoxide) group 4 metal dichloride complexes has been synthesized via treatment of MCl₄(THF)₂ (M = Ti, Zr, Hf) with sodium salts of the phosphanylphenols. The complexes bearing diphenylphosphine donors adopt C₁-symmetric, all-cis ground-state structures as determined by X-ray crystallography. In solution an O,P ligand exchange process is observed. Eyring analyses indicate a nondissociative process proposed to proceed via a 120° rotation of the P,P,Cl or the O,P,Cl trigonal faces. A zirconium complex containing diisopropylphosphine donors adopts a configurationally rigid trans-P₂ geometry. In ethylene polymerization studies (methylaluminoxane (MAO) activation), bis(6-tert-butyl-2-diphenylphosphanylphenoxide) zirconium and hafnium catalysts gave activities up to 49 000 and 2000 g/mmol·h·bar, respectively. Both of these catalysts also afforded efficient catalyzed chain growth using ZnEt₂. The zirconium derivative afforded activities up to 13 300 g/mmol • h for propylene polymerization using "dried"-MAO (DMAO) as the cocatalyst. The polymer had a slight syndiotactic bias and is formed by a chain-end control mechanism. A dibenzyl analogue of the highly active zirconium catalyst revealed an all-cis structure by crystallography, but a mixture of isomers in solution by NMR spectroscopy. Cationic monoalkyl complexes were synthesized using either $[CPh_3][B(C_6F_5)_4]$ or DMAO and suggest a highly fluxional structure for the catalytically active species.

Introduction

Ligands containing monoanionic oxygen donors, especially phenoxides, have proved to be a popular alternative to the cyclopentadienyl fragment in post-metallocene olefin polymerization catalysts. Although simple group 4 metal bisphenoxide catalysts are active, additional neutral donors are often incorporated into the ligand frame to occupy sites of coordinative unsaturation not needed for chain growth. Such additional donors may also help to "rigidify" the structure of the complex and open up the possibility for stereoselective polymerizations using prochiral olefins.

In group 4 metal complexes, the phenoxide donors tend to be matched with relatively hard first-row nitrogen-³⁻¹⁸ and oxygen-based⁷c,f-i,¹⁴c,h,19 donors. However, there is growing

evidence that softer second-row donors may offer beneficial stabilization of the highly reactive metal center. ^{14c,h,20} While there have been a growing number of reports concerning catalysts that combine phenoxides with thioether donors ^{2a,b,7e,14c,b,19g,21,22} there has been little focus on ligands containing tertiary phosphines ^{14c,d,j,23,24} Therefore bis(*o*-phosphanylphenoxide) complexes of the group 4 metals ^{25,26} appeared attractive; additional benefits include (i) a fairly straightforward synthesis (*vide infra*) with several sites around the ligand amenable to elaboration, (ii) a rigid phenylene bridge, which should help to enforce phosphine coordination, and (iii) an active species that could

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parallel the highly active group 4 metal phenoxyimine systems³ and offer a new class of early transition metal polymerization catalyst with the potential to influence polymer microstructure and stereochemistry.

Results and Discussion

Complex Synthesis and Structure. A range of *ortho*-substituted phosphanylphenols was synthesized from the corresponding phenols using the MOM protection, lithiation,

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deprotection strategy of Rauchfuss²⁷ (Scheme 1). In each case a hitherto unreported phosphonium salt was obtained that required treatment with ammonia to effect deprotection of the proligand. This product (A) was isolated and characterized during the synthesis of 6-(*tert*-butyl)-2-diphenylphosphanylphenol and revealed a hydroxymethyl substituent attached to the phosphorus center, presumably formed by an intramolecular attack during the MOM deprotection (Scheme 2; the structure of the salt was confirmed by crystallography; see Supporting Information).

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 $\begin{array}{lll} Scheme & 1. \ Synthesis & of & Bis(phosphanylphenoxide)M(IV) \\ & Dichloride \ Complexes \ (OMOM = OCH_2OCH_3) \end{array}$

Scheme 2. Proposed Mechanism for Formation of A

The phosphanylphenols were converted to sodium salts using NaH and reacted with $MCl_4(THF)_2$ (M=Ti, Zr, and Hf) to give the desired complexes 1-6 as analytically pure air- and moisture-sensitive solids (Scheme 1). The bis-O,P structures of 1 and 3-6 were confirmed by single-crystal X-ray analyses. The structures of 3 and 4 are discussed below; analyses of 1, 5, and 6 have been reported previously and revealed all-cis structures for 1 and 6 and a *trans*-P₂ structure for 5.25

The solid-state structure of 3 revealed the presence of two crystallographically independent molecules (I and II) in the asymmetric unit (molecule I is shown in Figure 1, molecule II in Figure S4 in the Supporting Information). The coordination sphere for each molecule is distorted octahedral with cis angles being in the ranges $73.81(7)-105.87(7)^{\circ}$ and $71.81(6)-101.22(7)^{\circ}$ for molecules I and II, respectively; the two most acute angles in each case are associated with the bites of the O,P ligands. The primary difference between the molecules is not the arrangement of the donor atoms, which, like in 1 and 6, are all-cis, but the conformation of the P(1)/O(1) ligand. This difference can be readily appreciated by considering the folding of the five-membered C₂OPTi chelate rings. While for both complexes the P(2)/O(2) ring is essentially planar (the five atoms being coplanar to within ca. 0.03 and 0.01 Å for molecules I and II, respectively), the P(1)/O(1) ring in each case has an envelope conformation with the metal lying out of the C₂OP plane (coplanar to within ca. 0.02 Å). For molecule I the metal lies ca. 0.57 Å out of this plane in the direction of O(2), whereas for molecule II the metal lies ca. 0.63 Å in the opposite direction toward Cl(2'). Keeping the Ti,Cl(1),Cl(2),P(2),O(1),O(2) coor-

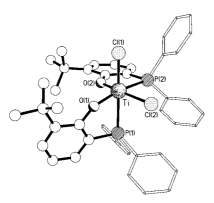


Figure 1. Molecular structure of I, one of the two independent molecules present in the crystals of 3.

Table 1. Selected Bond Lengths (Å) and Angles (deg) for Complexes 3 (molecules I and II), 4, and 6

	3 (I)	3 (II)		
	[M = Ti]	[M = Ti]	4 [M = Zr]	6 [M = Hf]
M-O(1)	1.839(2)	1.8388(19)	1.9632(15)	1.956(3)
M-O(2)	1.874(2)	1.837(2)	1.9889(14)	1.989(3)
M-P(1)	2.7055(8)	2.7408(9)	2.8872(6)	2.8628(11)
M-P(2)	2.6309(8)	2.6979(9)	2.8208(6)	2.7954(11)
M-Cl(1)	2.2679(9)	2.2794(9)	2.4239(6)	2.4058(11)
M-Cl(2)	2.3177(8)	2.3307(9)	2.4263(6)	2.4060(12)
Cl(1)-M-P(1)	178.98(4)	170.81(4)	170.66(2)	169.99(4)
Cl(2)-M-O(2)	161.34(7)	160.62(7)	154.08(5)	155.76(10)
P(2)-M-O(1)	159.34(7)	163.71(7)	152.95(5)	154.30(10)
Cl(1)-M-Cl(2)	94.95(3)	94.15(3)	96.87(2)	96.37(5)
Cl(1)-M-P(2)	92.18(3)	92.88(3)	103.30(2)	103.34(4)
Cl(1)-M-O(1)	105.87(7)	101.22(7)	101.81(5)	100.33(10)
Cl(1)-M-O(2)	93.52(7)	95.13(8)	92.55(5)	92.43(9)
Cl(2)-M-P(1)	86.07(3)	81.17(3)	85.026(19)	86.04(4)
Cl(2)-M-P(2)	88.05(3)	89.03(3)	84.69(2)	85.37(4)
Cl(2)-M-O(1)	100.04(7)	97.95(7)	102.16(5)	101.88(11)
P(1)-M-P(2)	87.96(2)	94.94(3)	85.965(16)	86.52(3)
P(1)-M-O(1)	73.81(7)	71.81(6)	68.86(5)	69.66(10)
P(1)-M-O(2)	85.54(6)	91.73(7)	89.57(4)	89.20(9)
P(2)-M-O(2)	75.03(6)	73.54(6)	69.62(4)	70.61(9)
O(1)-M-O(2)	93.57(9)	96.89(9)	99.44(6)	98.68(14)

dination sphere "stationary" (these atoms have an rms fit of ca. 0.06 Å in the two molecules), this difference can be viewed as a bending of the aryl ring of the P(1)/O(1) ligand about the P···O vector either toward or away from O(2) (see Figure S6 in the Supporting Information). This flexing is accompanied by a shift in the P(1) position between the two molecules by ca. 0.37 Å. The different conformations of the C₂OPTi rings may be caused by steric interactions between the two tert-butyl substituents as the analogous chelates in 1 are planar to within $0.04~{\rm \AA}.^{25}$ The *trans* influence of the various donors can be seen by inspection of the Ti-X bond lengths (Table 1). The Ti-P(1) bond is longer than its Ti-P(2) counterpart, reflecting the difference between a chloride and a (phen)oxide, respectively, in the trans position. Likewise, Ti-Cl(2) is longer than Ti-Cl(1), reflecting the difference between a formally monoanionic oxygen donor and a neutral phosphorus donor, and Ti-O(2) is longer than Ti-O(1) [in complex I but not, strangely, in complex II], showing the difference between chloride and phosphorus.

The crystal structure of the zirconium complex 4 (Figure 2) is very similar to that of 3-I. The geometry at the metal center is distorted octahedral with cis angles in the range 68.86(5)– $103.30(2)^\circ$. The coordination sphere is all-cis, and the two five-membered C_2OPZr chelate rings have different geometries; the P(2)/O(2)

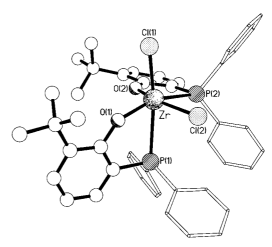


Figure 2. Molecular structure of 4.

Scheme 3. Interchange of all-cis enantiomers.

ring is flat (all five atoms coplanar to within ca. 0.03 Å), while the P(1)/O(1) ring has an envelope conformation with the metal lying ca. 0.36 Å out of the C₂OP plane (which is coplanar to within ca. 0.02 Å) in the direction of O(2).²⁸ This planarity for the P(2)/O(2) ring, and folding of the P(1)/O(1) ring, is also seen in the structure of the hafnium analogue 6,25 where the five atoms of the P(2)/O(2) ring are all coplanar to within ca. 0.03 Å, and the metal lies ca. 0.37 Å out of the $C_2O(1)P(1)$ plane (which is coplanar to within ca. 0.02 Å) in the direction of O(2).²⁹ The most noticeable difference between complexes 3 and 4 (except the obvious presence of only one independent molecule in the structure of 4) is in the pattern of bonding at the metal center (Table 1). Although the bonds to phosphorus and oxygen show the same pattern as in 3 [M-P(1) longer than M-P(2), M-O(2) longer than M-O(1)], the two Zr-Cl bond lengths are statistically identical [2.4239(6) and 2.4263(6) Å to Cl(1) and Cl(2), respectively], whereas in 3 the Ti-Cl(2) bonds were noticeably longer than those to Cl(1). This may in part be a consequence of the Zr-Cl bonds being ca. 0.1 Å longer than their Ti-Cl counterparts. This pattern of bond lengths is repeated in the hafnium analogue 6 (Table 1).

The configurations of 1-6 in solution were determined by ¹H and ³¹P NMR spectroscopy. All signals in the ³¹P spectra were shifted substantially downfield (by 16-30 ppm) from the value found for the corresponding sodium phosphanylphenoxide, consistent with coordination of the phosphines to the metal center. 23b,d If, as seen in the solid state, the PPh2-substituted complexes adopt a C_1 -symmetric, all-cis configuration, then two sets of signals are expected for the inequivalent O,P ligands. This should be conveniently observed in the ³¹P NMR spectra with two doublets anticipated for the two coupled phosphorus environments. However, only the spectrum of 3 gave this pattern $(^2J_{PP} = 6.2 \text{ Hz});$ ³¹P NMR spectra of 1, 2, 4, and 6 all returned single ³¹P resonances, apparently contradicting the crystallographic data. Therefore, variable-temperature ³¹P NMR experiments were performed. At low temperature the anticipated set of doublets was observed for each complex, thus indicating that a dynamic interchange of the all-cis enantiomers is occurring at ambient temperature (Scheme 3).

To probe the mechanism of ligand exchange, kinetic parameters for the averaging process were calculated for **2–4** and **6** using line-shape analysis³⁰ of the ³¹P NMR data measured over

the temperature range 203–363 K in toluene- d_8 ; selected ³¹P NMR spectra and calculated exchange rates are shown in Figure 3, and activation parameters are in Table 2. Significantly, for each complex a negative entropy of activation was obtained, suggesting that rearrangement of the ligands occurs via a nondissociative mechanism. ³¹ There are two possible nondissociative pathways for the interconversion of all-*cis* enantiomers in (O,P)₂MCl₂ complexes; ^{32,33} these are shown in Figure 4.

The first involves a 120° rotation about an O,P,Cl trigonal face with the two chelate rings confined to each face (C_{2v} -symmetric transition state); the second requires a 120° rotation about the P,P,Cl face with the chelate rings spanning the two trigonal faces (C_s -symmetric transition state). These mechanisms are not distinguishable by NMR spectroscopy. What is clear is that the rate of this rearrangement is determined by the steric bulk of the substituents *ortho* to the phenoxide donors, with **2**, bearing phenyl substituents, exchanging 100 times faster at 303 K than **3**, which contains *tert*-butyl groups (this difference is almost entirely entropic). Otherwise the separation of the trigonal faces, determined by the radii of the metal centers (Ti < Zr > Hf), appears to play a key role in controlling the rate of ligand exchange.

The NMR spectra of **5** afforded a single set of resonances for the O,P ligands in accord with the ground-state structure established by crystallography.²⁵ Comparable spectra were obtained at 323 and 223 K, indicating a nonfluxional structure.

Ethylene Polymerization Studies. Complexes 1–6 were investigated as catalysts for ethylene polymerization³⁴ using methylaluminoxane (MAO) as the cocatalyst; activity and molecular weight data are collected in Table 3. The productivities of the titanium catalysts, 1–3/MAO, are clearly improved when a bulky substituent is positioned *ortho* to the phenoxide donor. Increasing bulk also gives order of magnitude increases in molecular weight with concurrent decreases in the polydispersity index (PDI). Similar observations have been noted by Fujita et al. for phenoxyimine catalyst systems.³ Analyses of the polymer by ¹³C NMR spectroscopy revealed highly linear polyethylene (PE). Also visible in the polymer samples isolated using 1/MAO and 2/MAO were signals for *n*-alkyl end-groups, consistent with chain transfer to aluminum operating as the predominant termination mechanism.

The zirconium catalyst 4/MAO was found to have exceptional activity for ethylene polymerization with submicromolar quantities of catalyst required to prevent excessive exotherms and polymer yields. At reduced precatalyst loadings the concentration of cocatalyst also had to be reduced to achieve optimal activity. At a precatalyst loading of 40 nmol and employing 0.75 mmol of MAO, an optimized activity of 49 000 g/mmol • h • bar was recorded over 60 min, with a molecular weight (M_n) of 16.6 kg/mol. Altering the phosphine substituents from phenyl to isopropyl resulted in a dramatic lowering of the productivity and a very broad molecular weight distribution. Although care must be exercised when extrapolating metal-ligand geometries from precatalysts to the active species, the differing configurations of 4 (all-cis) and 5 (P-trans) may be responsible for this clear change in performance. We shall return later to scrutinize the nature of the active species derived from 4. The polyethylene from both zirconium catalysts showed evidence for chain transfer to aluminum in their ¹³C NMR spectra.

⁽²⁸⁾ The molecular structure of **5**, reported in ref 25, has folded C_2OPZr rings with the metal lying ca. 0.29 and ca. 0.36 Å out of the rest of the chelate.

⁽²⁹⁾ The chlorine, phosphorus, and oxygen atoms of the coordination sphere in **6** have been renumbered with respect to the report in ref 25 in order to match the numbering of the atoms in the structures reported here. Cl(1) and Cl(2) have swapped, as have P(1) and P(2), and O(1) and O(2).

⁽³⁰⁾ Reich, H. J. WINDNMR v7.1.6, 2002. For full kinetic data and Eyring plots see the Supporting Information.

⁽³¹⁾ Non-dissociative pathways have been proposed in other octahedral bis-bidentate Group 4 metal complexes: (a) Fay, R. C.; Lindmark, A. F. J. Am. Chem. Soc. 1983, 105, 2118. (b) Rahim, M.; Taylor, N. J.; Xin, S.; Collins, S. Organometallics 1998, 17, 1315. (c) Kakaliou, L.; Scanlon, W. J.; Qian, B.; Baek, S. W.; Smith, M. R.; Motry, D. H. Inorg. Chem. 1999, 38, 5964. (d) Otten, E.; Dijkstra, P.; Visser, C.; Meetsma, A.; Hessen, B. Organometallics 2005, 24, 4374.

^{(32) (}a) Bickley, D. G.; Serpone, N. *Inorg. Chem.* **1976**, *15*, 948. (b) Bickley, D. G.; Serpone, N. *Inorg. Chem.* **1976**, *15*, 2577.

⁽³³⁾ Bailar, J. C. J. Inorg. Nucl. Chem. 1958, 8, 165.

⁽³⁴⁾ After all polymerizations the reaction medium was examined by GC. No oligomeric products were detected in any instance.

2

18,900 s⁻¹

1.560 s

343 K

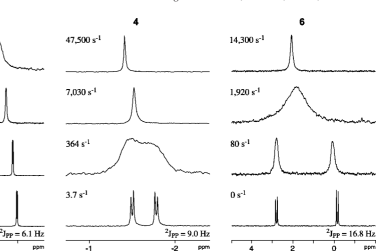
303 K

263 K

223 K

 $0 s^{-}$

12



10 Figure 3. Selected variable-temperature ^{31}P NMR spectra (toluene- d_8) for 2-4 and 6 with associated exchange rates.

3

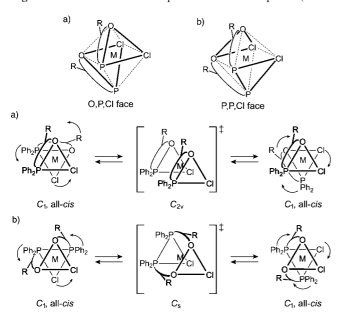
160 s⁻¹

16 s⁻¹

2.3 s⁻¹

 $0 \, s^{-1}$

12



 $^{2}J_{PP} = 6.4 \text{ Hz}$

10

Figure 4. Nondissociative pathways for ligand exchange in bis(phosphanylphenoxide)MCl₂ complexes.

Table 2. Activation Parameters for the Ligand Exchange in 2-4 and 6 (toluene-d₈ solvent)

complex (M, R)	ΔH^{\dagger} (kcal/mol)	ΔS^{\ddagger} (cal/K·mol)	$\Delta G^{\ddagger}_{298 ext{K}}$ (kcal/mol)
2 (Ti, Ph)	13.2(2)	-0.6(5)	13.4(4)
3 (Ti, ^t Bu)	12.3(2)	-12.6(5)	16.1(3)
4 (Zr, ^t Bu)	11.5(2)	-3.4(6)	12.5(3)
6 (Hf, ^t Bu)	10.8(1)	-8.3(5)	13.3(3)

The hafnium catalyst 6/MAO also showed a marked sensitivity to the quantity of cocatalyst, with an optimized productivity of 2000 g/mmol·h·bar. In all cases the polymer had a broad multimodal molecular weight distribution. It is also of note that polymerizations using 6/MAO were accompanied by significant induction times, with the vast majority of the polymer being formed in the final 30 min. Since aging a solution of 6/MAO (100 equiv) for 30 min prior to injection into the polymerization vessel did not improve the productivity, or reduce the induction time, it is proposed that insertion into the initially formed Hf—Me bonds is rate-limiting.³⁵ Furthermore, ¹³C NMR analysis of the polymer revealed that termination occurs by chain transfer to aluminum, which regenerates Hf-Me bonds and hence further limits insertion. As aluminum methyls are steadily consumed, the rate at which Hf-Me bonds are formed should decrease, resulting in an increased ethylene uptake as the run progresses.

Since chain transfer to aluminum appears to occur readily for these catalysts, we decided to investigate 4 and 6 as candidates for the catalyzed chain growth (CCG) of short chain linear alkanes using ZnEt₂ as the chain transfer agent.³⁶ Polymerizations in the presence or absence of ZnEt2 are collected in Table 4. For both catalysts introducing ZnEt2 to the reaction mixture gave a significant decrease in molecular weight and narrow molecular weight distributions (PDI 1.2–1.3) consistent with efficient CCG. Also of note was the considerable increase in the productivity of 6/MAO. This is consistent with the previous assertion that insertion into Hf-Me bonds is ratelimiting. In the presence of ZnEt₂ rapid alkyl exchange generates Hf-Et bonds into which ethylene readily inserts. Additional CCG reactions in which samples were taken at intervals through the run and the products analyzed by GC revealed Poisson distributions of PE oligomers that increased in molecular weight over time; selected distributions for 4 are shown in Figure 5 (all distributions can be found in the Supporting Information).

Propylene Polymerization Studies. Complexes 4 and 6 also showed significant productivities for the polymerization of propylene using "dried"-MAO³⁷ (DMAO) as the cocatalyst (see Table 5). At 20 °C 4/DMAO gave a viscous oil of moderate molecular weight and low PDI with a productivity of 5340 g/mmol·h·bar. Analysis of the polymer by ¹³C NMR spectroscopy (Figure 6) showed it to have a slight syndiotactic bias ([rr] = 38.5%; cf. 25.0% for atactic polymer). Full pentad

^{(35) (}a) Wester, T. S.; Johnsen, H.; Kittilsen, P.; Rytter, E. Macromol. Chem. Phys. 1998, 199, 1989. (b) Liu, Z.; Somsook, E.; White, C. B.; Rosaaen, K. A.; Landis, C. R. J. Am. Chem. Soc. 2001, 123, 11193.

^{(36) (}a) Britovsek, G. J. P.; Cohen, S. A.; Gibson, V. C.; Maddox, P. J.; van, Meurs Angew. Chem., Int. Ed. 2002, 41, 489. (b) Britovsek, G. J. P.; Cohen, S. A.; Gibson, V. C.; van Meurs, M. J. Am. Chem. Soc. 2004, 126, 10701. (c) van Meurs, M.; Britovsek, G. J. P.; Gibson, V. C.; Cohen, S. A. J. Am. Chem. Soc. 2005, 127, 9913. (d) Arriola, D. J.; Carnahan, E. M.; Hustad, P. D.; Kuhlmann, R. L.; Wenzel, T. T. Science 2006, 312, 714. (e) Kempe, R. Chem.-Eur. J. 2007, 13, 2764.

⁽³⁷⁾ DMAO is MAO with residual AlMe₃ removed under vacuum (see Supporting Information). DMAO has been shown to improve the performance of several non-metallocene catalysts: (a) Hasan, T.; Ioku, A.; Nishii, K.; Shiono, T.; Ikeda, T. Macromolecules 2001, 34, 3142. (b) Hagimoto, H.; Shiono, T.; Ikeda, T. Macromol. Rapid Commun. 2002, 23, 73. (c) Furayama, R.; Saito, J.; Ishii, S.; Mitani, M.; Matsui, S.; Tohi, Y.; Makio, H.; Matsukawa, N.; Tanaka, H.; Fujita, T. J. Mol. Catal. A: Chem. 2003, 200, 31.

Table 3. Ethylene Polymerization Results for Complexes $1-6^a$

precatalyst (loading (µmol))	MAO loading (mmol)	yield PE (g)	activity ^b	$M_{\rm n}^{\ c} \ ({\rm kg/mol})$	$M_{\rm w}^{\ c} \ ({\rm kg/mol})$	$M_{\rm w}/M_{\rm n}$
1 (5.0)	2.50	0.41	41	1.9	529	278^{d}
2 (5.0)	2.50	6.15	615	19.2	44.5	2.3
3 (5.0)	2.50	3.90	390	782	1,370	1.8
4 (0.1)	2.50	2.24	11 200	8.2	21.0	2.6
4 (0.1)	1.25	5.34	26 700	12.0	71.8	6.0
4 (0.1)	0.75	5.00^{e}	$100,000^{e}$	13.1	54.3	4.1
4 (0.04)	0.75	3.92	49 000	16.6	143	8.6
5 (5.0)	2.50	0.35	35	1.0	109	109 ^f
6 (1.0)	2.50	1.10	550	1.0	8.3	8.3^{g}
6 (1.0)	1.25	2.13	1070	2.0	228	114^{h}
6 (1.0)	0.75	4.00	2000	6.6	580	88^i

^a Conditions: 400 mL glass reactor, mechanical stirring, toluene (200 mL), 2 bar ethylene, 25 °C, 1 h (except e). ^b Activities in g/mmol·h·bar. ^c Determined by high-throughput GPC. ⁱ Trimodal: $M_{pk} = 1.7$, 51.5, and 899 kg/mol. ^h Trimodal: $M_{pk} = 1.2$, 18.6, and 562 kg/mol. ^d Multimodal: $M_{pk} = 1.2$, $M_{pk} = 1.2$, 1.5 kg/mol. ^e Run time 15 min. ^f Bimodal: $M_{\rm pk}=0.5$ and 25.9 kg/mol. ^g Bimodal: $M_{\rm pk}=0.6$ and 10.5 kg/mol.

Table 4. CCG on ZnEt₂ Using 4 and 6^a

precatalyst (loading (μmol))	ZnEt2 loading (mmol)	yield (g)	activity ^b	$M_n^c(g/\text{mol})$	$M_{\rm w}{}^c({ m g/mol})$	$M_{ m w}/M_n$
4 (0.5)	0	1.05	25 200	2600	13 900	5.3
4 (0.5)	4	1.25	30 000	250	300	1.2
6 (1.5)	0	0.17	461	800	3000	3.8
6 (1.5)	4	3.75	10 000	300	400	1.3

^a Conditions: Schlenk, magnetic stirring, MAO (1 mmol), toluene (100 mL), 1 bar ethylene, 25 °C, 5 min (4)/15 min (6). ^b Activities in g/mmol·h·bar. ^c Determined by high-throughput GPC.

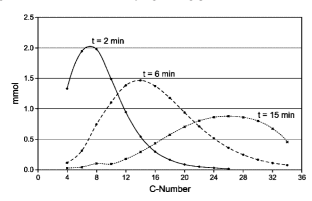


Figure 5. Oligomer distributions from 4/MAO with ZnEt₂. analysis³⁸ indicates this selectivity is induced by chain-end control, with the probability of a rac insertion (P_r) at 61.7%. No chain end-groups were visible in the NMR spectrum of the polymer. By lowering the polymerization temperature to -20°C, and hence increasing the concentration of propylene in solution, the activity of 4/DMAO was increased to 7010 g/mmol·h·bar. At -30 and -50 °C activities decreased, indicating that insertion becomes rate-limiting. At lower temperatures stereoselectivities did improve slightly; for example at -50 °C P_r = 65.9%. A polymerization in liquid propylene at room temperature³⁹ gave a productivity of 13 300 g/mmol·h, which is among the highest reported productivities for a postmetallocene propylene polymerization system. By using MAO as the cocatalyst the molecular weight of the polymer could be decreased such that isobutyl end-groups became visible in the ¹³C NMR spectrum. ⁴⁰ This not only indicates that chain transfer to aluminum operates but that insertion occurs in a 1,2-fashion (Scheme 4).

Consistent with the observations for ethylene polymerization, 6/DMAO (772 g/mmol · h · bar at 25 °C) gave a somewhat lower productivity than 4/DMAO, but this could be increased, to 949 g/mmol·h·bar, by raising the polymerization temperature to 50 °C. Again the polymer was isolated as a viscous oil, with NMR analysis revealing a reduced level of stereocontrol (P_r = 56.2% at 25 °C) and isobutyl end-groups.

Toward the Active Species. In order to rationalize the stereoselectivity of 4/DMAO for propylene polymerization, more information was required on the structure of the catalytically active species. Treating $ZrBn_4$ (Bn = CH_2Ph) with 2 equiv of 6-(tert-butyl)-2-diphenylphosphanylphenol, or reacting 4 with 2 equiv of BnMgBr, gave the dibenzyl complex 7 as an analytically pure air- and moisture-sensitive yellow solid (Scheme 5).

Recrystallization from heptane yielded crystals suitable for an X-ray structure determination. The molecular structure (Figure 7) revealed an geometry analogous to that found for its dichloride relative 4, with the oxygen, phosphorus, and carbon donor atoms all lying mutually cis. The geometry at the metal center is a little more distorted from octahedral for 7, the cis angles ranging between 65.70(8)° and 112.86(11)° (cf. $68.86(5)-103.30(2)^{\circ}$ in 4) and the two most acute angles being associated with the bites of the O,P ligands (Table 6). The P(2)/ O(2) five-membered C_2OPZr chelate ring is again flat, all five atoms being coplanar to within ca. 0.02 Å, while the P(1)/O(1)ring has an envelope conformation, the metal lying ca. 0.53 Å out of the C₂OP plane (which is coplanar to within ca. 0.01 Å) in the direction of O(2). Despite the change from chloride to benzyl ligands, the pattern of Zr-P and Zr-O bonding is similar, with Zr-P(1) longer than Zr-P(2) (carbon vs oxygen in the *trans* position) and Zr-O(2) longer than Zr-O(1) (carbon vs phosphorus in the trans position). For the two benzyl ligands, Zr-C(8) (trans to oxygen) is longer than Zr-C(1) (trans to phosphorus). There is no evidence of any multiple bonding between the benzyl ligand and the metal center.

The room-temperature solution-state NMR spectra of 7 revealed broad peaks for the benzyl and tert-butyl protons, indicative of a complex with an intermediate rate of O,P ligand exchange (Figure 8). Additionally, a sharp peak in the tert-butyl region suggested that a more rigid geometric isomer might also

⁽³⁸⁾ Pentad intensities determined by line fitting using MestReC v4.7.0.0. These were compared to best-fit chain-end and enantiomorphic site control models. (a) Busico, V.; Cipullo, R. Prog. Polym. Sci. 2001, 26, 443. (b) Bovey, F. A.; Tiers, G. V. D. J. Polym. Sci. 1960, 44, 173. (c) Shelden, R. A.; Fueno, T.; Tsunetsugu, T.; Furukawa, J. J. Polym. Sci., Part B 1965, 3, 23,

⁽³⁹⁾ Conditions: 1 L steel autoclave, mechanical stirring, 4 (1 μmol), MAO (2.5 mmol), 500 mL of liquid propylene, 25 °C, 1 h.

⁽⁴⁰⁾ Polymerization conditions: as for Table 5 except MAO (5.0 mmol) used as cocatalyst. See Supporting Information for ¹³C NMR spectrum.

Table 5. Propylene Polymerizations Using 4 and 6^a

precatalyst (loading (µmol))	temp (°C)	activity ^b	$M_{\rm n}^{\ c}({\rm kg/mol})$	$M_{\rm w}^{\ c}({ m kg/mol})$	$M_{ m w}/M_{ m n}$	$[rr]^d(\%)$
4 (2.5)	20	5,340	37.0	62.7	1.70	38.5
4 (2.5)	-20	7,010	52.9	79.1	1.49	40.5
4 (2.5)	-30	4,500	58.4	95.5	1.63	42.1
4 (2.5)	$-50^{\rm e}$	1,750 ^e	18.3	28.4	1.6	44.6
6 (10.0)	50	949	14.9	34.4	2.30	33.7
6 (10.0)	25	772	18.2	35.1	1.93	36.0
6 (10.0)	0	112	7.79	12.7	1.63	37.3

^a Conditions: Schlenk, magnetic stirring, DMAO (2.5 mmol), TIBAL (125 μ mol), heptane (100 mL) (except e), 2 bar propylene, 30 min (4)/1 h (6). ^b Activities in g/mmol·h·bar (except e). ^c Determined by GPC. ^d Determined by ¹³C NMR (101 MHz). ^e Liquid propylene (100 ml); activity in g/mmol • h.

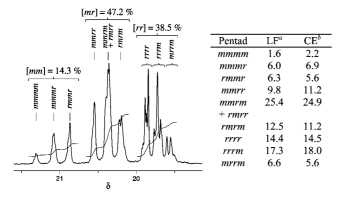


Figure 6. ¹³C NMR (101 MHz, methyl region) of polypropylene formed with 4/DMAO at 20 °C, with pentad analysis. "From linefitting analysis. ^bCalculated for chain-end control ($P_r = 61.7\%$).

Scheme 4. Isobutyl-Terminated Polymer from 1,2-Propylene Insertions and Chain Transfer to Aluminum

$$\begin{array}{c|c} M & & \\ & & \\ \hline & i) \text{ AlMe}_3 \text{ / Al}^{\text{I}} \text{Bu}_3 \\ \hline & ii) \text{ H}^{\text{+}} \\ \hline & 2 \text{ isobutyl endgroups} \end{array}$$

5. Synthesis of a Bis(phosphanylphenoxide) **Zirconium Dibenzyl Complex**

be present. The room-temperature ³¹P NMR spectrum revealed broad signals of approximately equal intensity at 2.3 and -9.3ppm, consistent with inequivalent but exchanging phosphorus environments in the all-cis structure. However, the spectrum also included a very broad peak at 5 ppm and a sharper signal at -14.6 ppm. It is proposed that these peaks are due to trans-O2 and trans-P2 geometric isomers. To probe whether this mixture could be converted to a single geometric product, a sample of 7 in C₆D₆ was heated to 50 °C for 2 h. No change in the isomer distribution was observed. It is also of note that the ratio of the isomers was dependent on the synthetic route employed, with a greater proportion of the all-cis isomer formed using BnMgBr (ca. 80% compared to ca. 53% using ZrBn₄). Therefore, the observed distribution is not an interchanging equilibrium mixture but rather represents a kinetic distribution of products.

On cooling to 223 K, the broad signal at 5 ppm in the ³¹P NMR spectrum sharpened to a singlet. Additionally, the broad signals assigned to the all-cis isomer resolved to two sharp doublets (${}^{2}J_{PP} = 21 \text{ Hz}$), consistent with a configurationally rigid C_1 -symmetric complex. The high-field peak does not signifi-

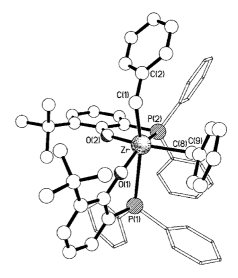


Figure 7. Molecular structure of 7.

Table 6. Selected Bond Lengths (Å) and Angles (deg) for 7

		. ,	. 0,
Zr-O(1)	1.981(3)	Zr-O(2)	2.013(3)
Zr-P(1)	3.0194(11)	Zr-P(2)	2.8583(10)
Zr-C(1)	2.277(4)	Zr-C(8)	2.310(4)
P(1) - Zr - C(1)	160.56(11)	P(2)- Zr - $C(1)$	112.86(11)
P(2) - Zr - O(1)	149.73(8)	P(2) - Zr - C(8)	79.59(11)
O(2)-Zr- $C(8)$	147.94(13)	O(1)-Zr- $C(8)$	107.34(14)
P(1) - Zr - P(2)	85.47(3)	O(1)-Zr- $C(1)$	95.09(13)
O(1) - Zr - O(2)	99.10(11)	O(2)-Zr- $C(1)$	94.55(14)
C(1)- Zr - $C(8)$	100.73(17)	P(1) - Zr - O(1)	65.70(8)
P(1) - Zr - O(2)	86.06(8)	P(2)- Zr - $O(2)$	68.49(7)
P(1) - Zr - C(8)	88.53(12)		

cantly alter upon cooling and is hence assigned to the trans-P₂ isomer by comparison to the variable-temperature behavior of 5. The low-temperature ¹H NMR spectrum of 7 includes five distinct benzyl signals in the region 1.9-3.0 ppm. Four of these are of equal intensity and are assigned to the four inequivalent benzyl protons of the all-cis isomer.41 It was anticipated that the benzyl protons of the trans-O₂ and trans-P₂ structures should give at least two signals. Since only one benzyl signal (2.68 ppm) remains unassigned, it is proposed that these two isomers have benzyl signals with coincidental chemical shifts. It should be noted that the low-temperature ¹³C NMR spectrum contains four signals for the benzylic carbons (64–74 ppm) and the quaternary carbon of the tert-butyl groups (34-35 ppm), consistent with there being one C_1 - and two C_2 -symmetric

The activation parameters for O,P ligand exchange in the all*cis* isomer of **7** ($\Delta H^{\dagger} = 10.5(3)$ kcal/mol, $\Delta S^{\dagger} = -10.4(9)$ cal/

⁽⁴¹⁾ A COSY spectrum revealed cross-peaks for the peaks at 2.93 and 1.95 ppm and at 2.33 and 2.17 ppm, indicating that each pair is bound to the same benzylic carbon.

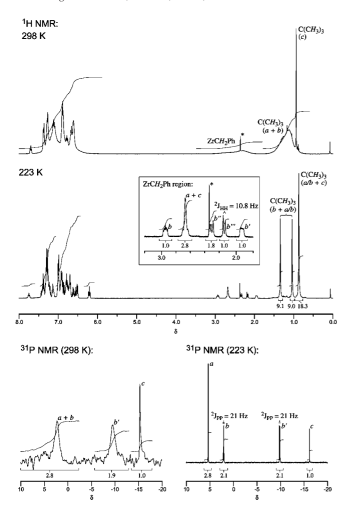


Figure 8. Variable-temperature ¹H and ³¹P NMR spectra of **7** (from ZrBn₄) in CDCl₃; * = residual toluene; a = trans-O₂, b = all-cis, c = trans-P₂ geometric isomer.

K·mol, $\Delta G^{\dagger}_{298\mathrm{K}} = 13.6(6)$ kcal/mol) were obtained using the same methodology as for the dichloro complexes.³⁰ Again, negative ΔS^{\dagger} values are indicative of a nondissociative process. Predictably, given the increased steric influence of the benzyl ligands compared to chlorides, the rate of isomerization was considerably slower than in 4 (approximately 7-fold at 298 K).

Of the reagents that can be used to abstract alkyl groups, [Ph₃C][B(C₆F₅)₄] is particularly attractive since both the borate anion and eliminated hydrocarbon are typically noncoordinating, thus facilitating study of the base-free cation. 42 Hence, 7 was treated with 1 equiv of [Ph₃C][B(C₆F₅)₄] at room temperature in an NMR-scale reaction in C₆D₅Br. Near-quantitative formation of the cationic benzyl complex 8 was observed in the ¹H NMR spectrum (Figure 9), which included two equimolar signals assignable to the CH_2 groups of the zirconium-bonded benzyl (2.84 ppm) and the eliminated 1,1,1,2-tetraphenylethane (3.82 ppm). Also of note are the relatively high-field shifts for the aromatic protons of the benzyl ligand (6.12-6.64 ppm), suggesting η^2 -coordination. Further evidence for this bonding mode was provided by a ¹H-coupled ¹³C NMR spectrum, which included a triplet at 79.3 ppm for the benzylic carbon with a $^1J_{\rm CH}$ value of 148 Hz. 43,44 Only one C(CH₃)₃ signal and one phosphorus signal are observed in the ¹H and ³¹P NMR spectra,

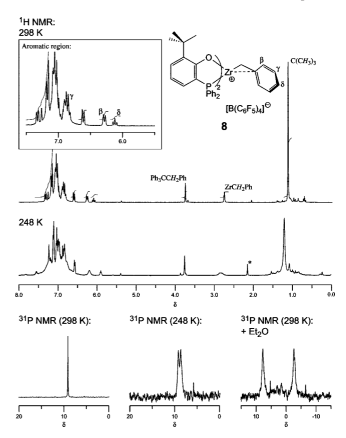


Figure 9. Variable-temperature 1H and ^{31}P NMR spectra of **8** in $C_6D_5Br.*=$ residual toluene.

respectively, indicating that the cation is either highly fluxional or adopts a square-based pyramidal structure with the benzyl ligand in the apical position. At low temperature the 1 H NMR spectrum significantly broadens and the 31 P NMR spectrum reveals two distinct, though broad, resonances indicative of a C_1 -symmetric ground state akin to the parent six-coordinate complex 7. To further probe the structure of 8, a small amount of ether was added to the NMR sample. The room-temperature 31 P NMR then contained two separate, equal intensity signals at 8.3 and -2.1 ppm, consistent with a six-coordinate ether complex in which trigonal face rotation is restricted by the additional steric constraints imposed by binding of a molecule of diethyl ether. Near-identical 19 F NMR spectra were obtained with or without ether coordination, consistent with $B(C_6F_5)_4$ acting as a noncoordinating counterion.

An analogous cationic species 8' could be obtained upon treatment of 7 with 50 equiv of DMAO in C_6D_6 (Figure 10). The 1H NMR spectrum clearly shows signals for an η^2 -bound benzyl moiety, and the ^{31}P NMR spectrum reveals a sharp singlet almost identical to that seen for 8. Analogous NMR-scale experiments were performed using the dichloride complex

⁽⁴²⁾ Coordination of solvent cannot typically be ruled out. A chlorobenzene adduct of a zirconium benzyl cation has been structurally characterized: Wu, F.; Dash, A. K.; Jordan, R. F. *J. Am. Chem. Soc.* **2004**, *126*, 15360.

⁽⁴³⁾ Values above 130 Hz indicate some distortion of the benzyl ligand. Examples of Zr complexes in which a solution state η^2 -interaction has been confirmed by X-ray analysis include: (a) Tsurugi, H.; Matsuo, Y.; Yamsgata, T.; Mashima, K. *Organometallics* **2004**, *23*, 2797. (b) Shao, P.; Gendron, R. A. L.; Berg, D. J.; Bushnell, G. W. *Organometallics* **2000**, *19*, 509. (c) Bei, X. H.; Swenson, D. C.; Jordan, R. F. *Organometallics* **1997**, *16*, 3282. (d) Crowther, D. J.; Borkowsky, S. L.; Swenson, D.; Meyer, T. Y.; Jordan, R. F. *Organometallics* **1993**, *12*, 2897. (e) Jordan, R. F.; Lapointe, R. E.; Bajgur, C. S.; Echols, S. F.; Willett, R. *J. Am. Chem. Soc.* **1987**, *109*, 4111. (f) Latesky, S. L.; McMullen, A. K.; Niccolai, G. P.; Rothwell, I. P.; Huffman, J. C. *Organometallics* **1985**, *4*, 902.

⁽⁴⁴⁾ It has recently been suggested that in the absence of coordinated solvent benzyl ligands may in fact bind in an η^3 manner: Sassmannshausen, J.; Track, A.; Stelzer, F. *Organometallics* **2006**, *25*, 4427.

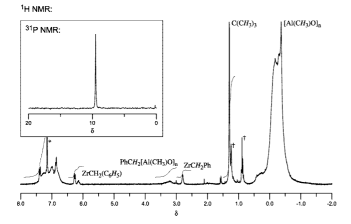


Figure 10. ¹H and ³¹P NMR spectra of **8**′ (7 + DMAO) in C_6D_6 (*). † = residual heptane in DMAO.

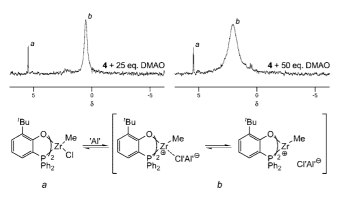


Figure 11. ^{31}P NMR spectra of 4 + DMAO in C_6D_6 .

4 with differing amounts of DMAO. The ³¹P NMR spectra reveal a low-intensity sharp signal and a larger broad peak (Figure 11). The low-intensity signal is tentatively assigned to a (O,P)₂Zr(Me)Cl complex and the broad peak to (O,P)₂Zr⁺Me. In the absence of the additional alkyl—metal interactions seen for **8/8'** it is proposed that the aluminoxane anion can reversibly access the zirconium center. Such dynamic interactions, and the nonuniform nature of aluminoxanes, can hence explain the observed broadening. ⁴⁵ Increasing the concentration of DMAO increases both the yield of the cationic species and the likelihood of cation—anion interactions.

Interestingly, when a solution of **8** (formed using [Ph₃C][B(C₆F₅)₄]) was placed under an atmosphere of ethylene, the yellow color was retained⁴⁶ and no polymer was formed, suggesting that the η^2 -benzyl interaction is too strong to be displaced by ethylene.⁴⁷ This was reproduced under polymerization conditions (Table 7) with no polymer formed at 25 °C. However, running the polymerization at 50 °C did give polymer, suggesting that an $\eta^2 - \eta^1$ rearrangement may occur at elevated temperature. Furthermore, polymer could be formed at room temperature using MAO as the cocatalyst, with a steady increase in ethylene uptake measured throughout the run. This suggests gradual exchange of the η^2 -benzyl ligands for methyls by the AlMe₃ present in the cocatalyst.

Given the fluxional nature of the catalytically active species derived from $\bf 4$ or $\bf 7$ it is perhaps less surprising that the

Table 7. Ethylene Polymerizations with 7^a

cocatalyst ^b	temp (°C)	yield PE (g)	activity ^c	$M_{\rm n}^{d}$ (kg/mol)	$M_{\rm w}^{}$ (kg/mol)	$M_{\rm w}/M_{\rm n}$
ТВ	25	0.00	0			
TB	50	1.56	3900	6.3	16.2	2.6
MAO	25	2.79	14 000	7.8	20.6	2.6

^a Conditions: glass reactor, mechanical stirring, 7 (0.2 μmol), MAO (2.5 mmol) or [Ph₃C][B(C₆F₅)₄] (0.2 μmol) and AlⁱBu₃ (0.5 mmol), toluene (200 mL), 2 bar ethylene, 1 h (TB)/30 min (MAO). ^b TB = [Ph₃C][B(C₆F₅)₄]. ^c Activities in g/mmol • h • bar. ^d Determined by GPC vs PS standards.

polypropylene formed is virtually atactic. However, it should be noted that there are fluxional zirconocenes that can form isotactic-atactic stereoblock polymers. 48 Furthermore, the highly syndiotactic polymer generated using bis(phenoxyimine)titanium catalysts is thought to be a direct consequence of an isomerization process that occurs after each monomer insertion.⁴⁹ For a fluxional catalyst to give stereoenriched PP via an enantiomorphic site control mechanism, two requirements must be met. First, the ground-state structure(s) must be able to influence the orientation of the polymer chain in order to control the mode of monomer insertion. Second, the rate of geometrical isomerization must be comparable to, but preferably less than, the rate of insertion. Assuming that the O,P ligand framework in 4/DMAO can control monomer insertion, this means the rate of ligand reorganization must be slower than the rate of monomer insertion, which is calculated as 71 s⁻¹ at 20 °C. 50 A crude comparison with the ligand exchange rates calculated for 4 and 7 (3400 and 596 s⁻¹, respectively) indicates that the fluxional processes in 4/DMAO are too rapid for any stereocontrol to be imparted.

Conclusions

A number of bis(phosphanylphenoxide) group 4 metal complexes have been synthesized in order to assess the effect of incorporating phosphine donors into early transition metal based olefin polymerization catalysts. The geometries of these complexes are dependent upon the phosphine substituents, with disopropyl- and diphenyl-phosphine zirconium derivatives possessing *trans*-P₂ and all-*cis* structures, respectively.

Polymerization studies have highlighted a *tert*-butyl-substituted zirconium catalyst that shows excellent activity in the production of polyethylene and polypropylene. In fact this catalyst is one of the most active post-metallocene catalysts reported to date, suggesting that there is considerable scope for utilizing the combination of hard donors with soft phosphine donors in early transition metal polymerization catalysis. It can be expected that linking the O,P ligands to one another might increase the configurational stability of the catalyst, which may then offer the possibility for stereoselective α-olefin polymerizations.

Experimental Methods

The synthesis of the phosphanylphenols, phosphanylphenoxides, and complexes 1-6 has been reported elsewhere. ²⁵ ZrBn₄ was synthesized according to a published procedure. ⁵¹ Air-sensitive procedures were performed under nitrogen using standard Schlenk techniques, with solvents being distilled over standard drying agents

⁽⁴⁵⁾ Hawrelak, E. J.; Deck, P. A. Organometallics 2003, 22, 3558.

⁽⁴⁶⁾ Kress et al. have observed a single ethylene insertion, indicated by a lack of polymer coupled with a loss of color: Gauvin, R. M.; Osborn, J. A.; Kress, J. *Organometallics* **2000**, *19*, 2944.

⁽⁴⁷⁾ Bochmann, M.; Lancaster, S. J. Organometallics 1993, 12, 633.

⁽⁴⁸⁾ Lin, S.; Waymouth, R. M. Acc. Chem. Res. 2002, 35, 765.

⁽⁴⁹⁾ Miliano, G.; Cavallo, L.; Guerra, G. J. Am. Chem. Soc. 2002, 124, 13368.

⁽⁵⁰⁾ Insertion rate (s^{-1}) = polymer yield (g)/[fw(propylene) (g/mol) × cat. loading (mol) × run time (s)].

⁽⁵¹⁾ Felten, J. J.; Anderson, W. P. J. Organomet. Chem. 1972, 36, 87.

The numbering scheme below is used in the assignments of the O,P precursors and complexes:

Synthesis of A. To a solution of 1-tert-butyl-2-(methoxymethoxy)benzene (14.55 g, 75.0 mmol) in ether (150 mL) was added n-butyllithium (30.6 mL, 76.5 mmol), and the mixture was stirred for 12 h. Chlorodiphenylphosphine (14.15 mL, 78.8 mmol) was added to the mixture dropwise at -78 °C. The mixture was stirred at room temperature for 3 h and then filtered. The filtrate was concentrated to dryness and the residue dissolved in THF (75 mL) and 5 M HCl (75 mL). The mixture was stirred and heated at 50 °C for 12 h. The mixture was poured into water (500 mL), filtered, and washed with water (3 \times 100 mL). The crude product was triturated in methanol, filtered, and dried under vacuum to give a colorless powder (16.69 g, 41.6 mmol, 56% yield). Anal. Calcd (%) for C₂₃H₂₆ClO₂P (400.88): C, 68.91; H, 6.54. Found: C, 68.87; H, 6.49. ¹H NMR (400 MHz, CDCl₃): δ 9.40 (br, 1H, Ar-O H), 7.74–7.70 (m, 3H, Ar- $H_{(4,D)}$), 7.58–7.53 (m, 8H, Ar- $H_{(B,C)}$), 7.29 (br, 1H, CH₂O H), 7.02 (dt, J = 7.7 Hz, J = 3.6 Hz, 1H, Ar- $H_{(5)}$), 6.59 (ddd, ${}^{3}J_{HP} = 13.9 \text{ Hz}$, ${}^{3}J_{HH} = 6.9 \text{ Hz}$, ${}^{4}J_{HH} = 1.4 \text{ Hz}$, 1H, Ar- $H_{(6)}$), 5.43 (s, 2H, C H_2 OH), 1.47 (s, 9H, C(C H_3)₃). ¹³C{¹H} NMR (101 MHz, CDCl₃): δ 159.5 (d, ${}^{2}J_{CP} = 3.0$ Hz, Ar- $C_{(2)}$), 144.7 $(d, {}^{3}J_{CP} = 6.6 \text{ Hz}, \text{Ar-}C_{(3)}), 135.3 (\text{Ar-}C_{(4)}), 134.5 (\text{Ar-}C_{(D)}), 133.8$ (d, ${}^{2}J_{CP} = 9.2 \text{ Hz}$, Ar- $C_{(6)}$), 133.2 (d, ${}^{3}J_{CP} = 9.4 \text{ Hz}$, Ar- $C_{(C)}$), 130.0 (d, ${}^{2}J_{CP} = 12.0 \text{ Hz}$, Ar- $C_{(B)}$), 123.6 (d, ${}^{3}J_{CP} = 13.9 \text{ Hz}$, Ar- $C_{(5)}$), 119.3 (d, ${}^{1}J_{CP} = 82.7 \text{ Hz}$, Ar- $C_{(A)}$), 109.6 (d, ${}^{1}J_{CP} = 89.4 \text{ Hz}$, Ar- $C_{(I)}$), 61.0 (d, ${}^{1}J_{CP} = 62.6$ Hz, P $CH_{2}OH$), 35.3 ($C(CH_{3})_{3}$), 30.4 $(C(CH_3)_3)$. ³¹P{¹H} NMR (162 MHz, CDCl₃): δ 20.9 (s). MS (negative FAB; m/z): 399 [M]⁻, 333 [M - Cl - CH₂OH]⁻, 188 $[M - {}^{t}Bu-2Ph]^{-}$. IR (KBr; cm⁻¹): 3142 (OH), 2868br (OH). Crystals suitable for X-ray analyses were grown from a chloroform solution layered with pentane.

Synthesis of 7. Method 1: A solution of ZrBn₄ (0.312 g, 0.68 mmol) in toluene (20 mL) was cooled to -10 °C and a solution of 2-*tert*-butyl-6-diphenylphosphanylphenol (0.477 g, 1.43 mmol) in toluene (20 mL) added dropwise. The mixture was stirred for 30 min at -10 °C and for 30 min at 25 °C. The solvent was removed under vacuum and the residue triturated with pentane (5 mL), filtered, and dried under vacuum to give **7** as a yellow powder (0.547 g, 0.58 mmol, 85% yield). Anal. Calcd (%) for C₅₈H₅₈O₂P₂Zr (940.25): C, 74.09; H, 6.21. Found: C, 73.82; H, 6.10. Three different isomers were detected by NMR analysis: *trans*-O₂ (isomer *a*), all-*cis* (*b*), and *trans*-P₂ (*c*). ¹H NMR (400 MHz, CDCl₃, 298

K): δ 7.70 (t, J = 8.3 Hz, Ar-H), 7.38–6.61 (br m, Ar-H), 3.00–1.80 $(br, 4H(a) + 4H(b) + 4H(c), C H_2Ph), 1.50-0.80 (br, 18H(a) +$ 18H(b), C(C H_3)₃), 0.94 (s, 18H(c), C(C H_3)₃). ¹H NMR (400 MHz, CDCl₃, 223 K): δ 7.74 (t, J = 7.8 Hz, Ar-H), 7.43–7.12 (m, Ar-H), 7.02–6.68 (m, Ar-H), 6.64–6.50 (m, Ar-H), 6.21 (t, J = 8.1Hz, Ar-H), 2.96–2.90 (m, 1H(b), C H_2 Ph), 2.71–2.64 (m, 4H(a) + 4H(c), C H_2 Ph), 2.33 (d, $^2J_{HH} = 10.8$ Hz, 1H(b), C H_2 Ph), 2.17 (d, $^{2}J_{HH} = 10.8 \text{ Hz}, 1H(b), C H_{2}Ph), 1.97-1.92 \text{ (m, } 1H(b), C H_{2}Ph),$ 1.34 (s, 9H(a/b), $C(C H_3)_3$), 1.04 (s, 9H(a/b), $C(C H_3)_3$), 0.87 (s, 18H(a/b) + 18H(c), C(C H_3)₃). $^{13}C\{^{1}H\}$ NMR (101 MHz, CDCl₃, 223 K): δ 169.1, 169.0, 168.9, 168.7, 166.4, 166.1, 147.6, 147.5, 146.0, 145.8, 137.8, 137.3, 137.2, 137.2, 137.1, 134.0, 133.6, 133.6, 133.5, 133.5, 133.3, 133.1, 133.1, 132.8, 132.7, 132.2, 132.1, 131.9, 131.8, 131.6, 131.5, 131.3, 131.2, 131.1, 131.0, 130.8, 130.5, 130.1, 129.9, 129.9, 129.8, 129.4, 129.3, 129.2, 129.0, 128.9, 128.7, 128.4, 128.2, 128.0, 127.9, 127.7, 127.2, 125.5, 125.5, 125.2, 123.7, 123.5, 121.3, 120.7, 120.5, 120.2, 119.7, 119.7, 119.6, 119.4, 119.3, 118.3 (Ar-C), 73.9, 73.7, 69.6, 63.5 (CH₂Ph), 34.8, 34.7, 34.4, 34.0 $(C(CH_3)_3)$, 29.2, 28.8, 28.3 $(C(CH_3)_3)$. $^{31}P\{^1H\}$ NMR (162 MHz, CDCl₃, 298 K): δ 5.3–2.4 (br, 2P(a) + 1P(b) [49%]), -9.5 (br, 1P(b) [33%]), -15.3 (s, 2P(c) [18%]). $^{31}P\{^{1}H\}$ NMR (162 MHz, CDCl₃, 223 K): δ 5.3 (s 2P(a) [35%]), 2.0 (d, ${}^{2}J_{PP} = 21$ Hz, 1P(b) [27%]), -9.8 (d, ${}^{2}J_{PP} = 21$ Hz, 1P(b) [27%]), -16.2 (s, 2P(c)[11%]).

Method 2: To a solution of **4** (0.301 g, 0.36 mmol) in toluene (30 mL) cooled to -78 °C was added dropwise PhCH₂MgBr (1.24 M in THF; 0.60 mL, 0.74 mmol), giving immediate formation of a yellow solution. The mixture was allowed to warm to room temperature over 12 h, over which time a colorless precipitate was formed. The suspension was filtered through Celite and the solvent then removed under vacuum. The residual yellow oil was triturated with pentane and the resultant precipitate filtered and dried under vacuum to give **7** as a yellow powder (0.181 g, 0.19 mmol, 53% yield). Anal. Calcd (%) for C₅₈H₅₈O₂P₂Zr (940.25): C, 74.09; H, 6.21. Found: C, 73.95; H, 6.10. 31 P{ 1 H} NMR (101 MHz, CDCl₃, 298 K): δ 2.3 (br, 2P(a) + 1P(b) [60%]), -9.2 (br, 1P(b) [40%]), -15.2 (s, 2P(c) [<1%]).

Synthesis of 8. In a nitrogen filled glovebox, a C₆D₅Br solution of [CPh₃][B(C₆F₅)₄] (1 equiv) was added to a C₆D₅Br solution of 7 (1 equiv) at 25 °C. The orange solution was transferred to an NMR tube fitted with a Youngs tap valve. The NMR data was acquired immediately. ¹H NMR (250 MHz, C₆D₅Br, 298 K): δ 7.34 (dd, ${}^{3}J_{HH} = 7.4 \text{ Hz}$, ${}^{2}J_{HH} = 1.6 \text{ Hz}$, 2H, Ar-H), 7.28–6.98 (m, 34 H, Ar-H), 6.93–6.82 (m, 8H + 2H, Ar-H + Ar- $H_{(\gamma)}$), 6.62 (d, J =7.8 Hz, 2H, Ar-H), 6.28 (d, ${}^{3}J_{HH} = 7.5$ Hz, Ar- $H_{(\beta)}$), 6.12 (t, ${}^{3}J_{HH}$ = 7.3 Hz, Ar- $H_{(\delta)}$), 3.81 (s, 2H, Ph₃CC H_2 Ph), 2.83 (s, 2H, ZrC H_2 Ph), 1.22 (s, 18H, C(C H_3)₃). ¹H NMR (400 MHz, C₆D₅Br, 248 K): δ 7.50–6.40 (br m, 48H, Ar-H), 6.24 (br s, Ar- $H_{(\beta)}$), 5.94 (br t, ${}^{3}J_{HH} = 7.5 \text{ Hz}$, Ar- $H_{(\delta)}$), 3.78 (s, 2H, Ph₃CC H_{2} Ph), 2.84 (br s, 2H, ZrC H₂Ph), 1.21 (s, 18H, C(C H₃)₃). ¹³C NMR (63 MHz, C_6D_5Br , 298 K): selected δ 165.3 (br, Ar- $C_{(I)}$), 150.6 (s, Ar- $C_{(\alpha)}$), 79.3 (t, ${}^{1}J_{CH} = 148 \text{ Hz}$, Zr $CH_{2}Ph$), 35.2 (s, $C(CH_{3})_{3}$), 29.7 (q, $^{1}J_{CH} = 123 \text{ Hz}, C(CH_{3})_{3}).$ $^{31}P\{^{1}H\} \text{ NMR (101 MHz, C}_{6}D_{5}Br, 298)$ K): δ 9.4 (s). ³¹P{¹H} NMR (162 MHz, C₆D₅Br, 248 K): δ 8.4 + 7.9 (2 unresolved br s). $^{19}F\{^{1}H\}$ NMR (235 MHz, C_6D_5Br , 298 K): $\delta -135.9$ (s, o-BC₆ F_5), -166.4 (t, ${}^3J_{\text{FF}} = 21.2$ Hz, p-BC₆ F_5), -170.2 (pt, $J_{FF} = 18.5$ Hz, $m\text{-BC}_6F_5$).

The reaction was repeated and two drops of dry Et₂O added to the solution. The NMR data were acquired immediately. ¹H NMR (250 MHz, C₆D₅Br, 298 K): selected δ 7.59 (d, ³ J_{HH} = 7.5 Hz, 2H, Ar-H), 7.18–6.40 (br m, 49 H, Ar-H), 2.50 (br s, 2H, ZrC H_2 Ph). ³¹P{¹H} NMR (101 MHz, C₆D₅Br, 298 K): δ 8.3 (br s), -2.1 (br s). ¹⁹F{¹H} NMR (235 MHz, C₆D₅Br, 298 K): δ -136.0 (s, o-BC₆ F_5), -166.6 (t, ³ J_{FF} = 21.0 Hz, p-BC₆ F_5), -170.4 (pt, J_{FF} = 18.4 Hz, m-BC₆ F_5).

Reaction of 7 with DMAO. In a nitrogen-filled glovebox, a C_6D_6 solution of DMAO (50 equiv) was added to a C_6D_6 solution of **7**

(1 equiv) at 25 °C. After 20 min, the pale yellow solution was transferred to an NMR tube fitted with a Youngs tap valve, and NMR data were acquired. ¹H NMR (250 MHz, C_6D_6): δ 7.41–6.70 (dd + br m, ${}^3J_{\rm HH}$ = 6.9 Hz, ${}^4J_{\rm HH}$ = 2.1 Hz, 33H, Ar-H), 6.26 (d, ${}^3J_{\rm HH}$ = 7.8 Hz, 2H, Ar- $H_{(\beta)}$), 6.14 (t, ${}^3J_{\rm HH}$ = 6.7 Hz, 1H, Ar- $H_{(\delta)}$), 3.24 (br, 2H, DMAO-C H_2 Ph), 2.80 (s, 2H, Zr-C H_2 Ph), 1.30 (s, 18H, C(C H_3)₃), 0.5 > 23>(-1.0) (br m, Al-C H_3). 31 P{ 1 H} NMR (101 MHz, C_6D_6): δ 9.4 (s).

Ethylene Polymerization Procedure. A 400 mL glass Fisher-Porter vessel was dried at 50 °C for 12 h. The Fisher-Porter head, equipped with an injection port (fitted with a rubber septum), pressure gauge, and mechanical stirrer, was flame-dried for 5 min and fitted to the glass vessel. The reactor was then purged with nitrogen for 45 min, charged with solvent, purged with 1 bar of ethylene overpressure for 5 min (to give 2 bar total ethylene pressure), and placed in a room-temperature water bath. Scavenger (MAO or AliBu₃) was introduced at least 5 min before catalyst injection. A solution of precatalyst (for polymerizations using MAO) or activated catalyst (for polymerizations using [CPh₃][B(C₆F₅)₄]) in toluene was introduced via syringe to initiate the polymerization. The reaction mixture was rapidly stirred throughout. Polymerizations were terminated by venting the overpressure followed by addition of 5 mL of 2 M HCl_(aq). A sample of the reaction mixture was taken for GC analysis. The solid product was precipitated from the mixture by addition of MeOH (400 mL). After filtration the polymer was washed with a large quantity of MeOH and dried under vacuum at 60 °C for 12 h.

Catalyzed Chain Growth Procedure. A Schlenk, containing 100 mL of toluene and a magnetic stirrer under a nitrogen atmosphere, fitted with a rubber septum was purged with ethylene for 5 min and the overpressure then vented (1 bar total ethylene pressure). MAO and, when required, ZnEt₂ were then added. If GC samples were to be taken during the run, then a known quantity of 2,2,4,4,6,8,8-heptamethylnonane was added as the GC standard. The catalyst was introduced via syringe. Reactions were terminated and worked up in the same manner as for the ethylene polymerizations above.

Propylene Polymerization Procedures. Method 1 (-30 to 50 °C): A Schlenk, containing 100 mL of heptane, DMAO, and a magnetic stirrer under a nitrogen atmosphere, fitted with a rubber septum was purged under 1 bar of propylene overpressure for 5 min (to give 2 bar total propylene pressure). AliBu₃ was added as a scavenger and the temperature of the vessel adjusted as required, allowing 20 min for the solvent to saturate with propylene. A toluene solution of precatalyst treated with 100 equiv of DMAO was introduced via syringe to initiate the polymerization. The reaction mixture was rapidly stirred throughout. Polymerizations were terminated by venting the overpressure followed by addition of 5 mL of 2 M HCl_(aq). After warming to room temperature, a further 50 mL of 2 M HCl(aq) was added, and the layers were separated. The organic fraction was washed with 2 M HCl_(aq) (50 mL) and water (50 mL), dried (MgSO₄), and filtered. The solvent was removed under vacuum and the polymer dried under high vacuum for 12 h.

Method 2 (-50 °C): The vessel was prepared as in method 1, except no solvent was added, and then placed in a cold bath at -50 °C. When 100 mL of propylene had condensed, the Schlenk was isolated from the propylene feed. A toluene solution of precatalyst treated with 100 equiv of DMAO was introduced via syringe to initiate the polymerization. The reaction mixture was rapidly stirred throughout. Polymerizations were terminated by addition of 5 mL of 2 M HCl_(aq). The vessel was **SLOWLY** vented and warmed to room temperature. The polymer was isolated as for method 1.

Acknowledgment. BP Chemicals and, subsequently, Innovene and Ineos are thanked for financial support of this work.

Supporting Information Available: Crystallographic details, VT-NMR spectra, kinetic data, polymer NMRs, and GC results. OM7008149