Cationic Alkylzirconium Complexes Based on a Tridentate Diamide Ligand: New Alkene Polymerization Catalysts

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Summary: Alkyl abstraction from {*Me3SiN(CH2CH2-* $NSiMe_3\langle R_2(R_2/R_2; R = CH_2Ph, Me)$ *using* $B(C_6F_5)$ ₃ *affords cationic alkyl complexes stabilized by a diamide ligand. The ionic η2-benzyl adduct decomposes slowly to give a cationic cyclometalation product, which coordinates the [PhCH2B(C6F5)3]*- *anion; the methyl cation coordinates the anion [MeB(C6F5)3]*- *via a Zr*'''*Me*-*B interaction. The complexes exhibit moderate ethene polymerization activity.*

The search for alternatives to group 4 metallocenes as electrophilic "uniform site" catalysts for alkene polymerization is currently of great interest.^{1,2} Much attention has focused on nitrogen-containing ligands $([L]^-$ or $[L_2]^{2-}$), such as porphyrins,³ tetraaza[14]annulenes,⁴ tetradentate Schiff base ligands,⁵ (hydroxy $phenyl)oxazolines⁶$ and benzamidinates.⁷ Unfortunately ethene polymerization activities (for $L_2MCl_2/$ methylaluminoxane or $[L_2MR]^+$; M = Ti, Zr, Hf) have been generally disappointing. $4-7$ In contrast, and despite the application of dicyclopentadienyl and cyclopentadienylamide catalysts in kiloton-scale industrial processes, diamide complexes (the next family in this series) have received little attention in the patent⁸ or scientific literature. $9-14$ This is surprising considering the facile synthesis of diamide complexes as catalyst

(8) (a) Isotactic polypropene formation using {(Me3Si)2N}2MCl2/MAO has been claimed: Canich, A. M.; Turner, H. W. Patent Application WO 92/12162 (Exxon, 1992). (b) Sasaki, T.; Shiraishi, H.; Johoji, H.; Katayama, H. European Patent Application 571 945 (Sumitomo, 1993).

(9) A zirconium diamide/methylaluminoxane polymerization catalyst has very recently been reported: Cloke, F. G. N.; Geldbach, T. J.; Hitchcock, P. B.; Love, J. B. *J. Organomet. Chem.* **1996**, *506*, 343.

(10) For very recent examples of group 4 complexes with chelating diamide ligands, see refs 9, 13, and 14 and the following: (a) Aoyagi, K.; Gantzel, P. K.; Kalai, K.; Tilley, T. D. *Organometallics* **1996**, *15*, 923. (b) Warren, T. H.; Schrock, R. R.; Davis, W. M. *Organometallics* **1996**, *15*, 562. (c) Scollard, J. D.; McConville, D. H.; Vittal, J. J. *Organometallics* **1995**, *14*, 5478.

 $precursors⁹⁻¹⁴$ and the expected increased electrophilicity of diamide cations compared to analogues with polydentate nitrogen ligands. A program to probe the potential of diamide ligands in polymerization has now lead to the first alkene polymerization catalysts based on cationic diamide complexes. Recent reports $13,14$ of *neutral* group 4 adducts of the new tridentate ligand $[Me₃SiN(CH₂CH₂NSiMe₃)₂]$ ²⁻ lead us to report here our work involving *cationic* group 4 alkyl adducts of this ligand.

New crystalline dialkyl precursors of putative cationic complexes have been prepared from the readily available dichlorozirconium complex **1**15,16 using standard alkylation methodology (Scheme 1).17,18 The observation of ¹H NMR resonances (C_6D_5Br) for inequivalent benzyl $(2, -20 \degree C)$ or methyl groups $(3, 25 \degree C)$, as well as four different backbone dimethylene hydrogens,¹⁹ is consistent with coordination of the amino nitrogen to zirconium in a trigonal bipyramidal structure. Fluxional exchange of the environments of the axial and equato-

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(14) Clark, H. C. S.; Cloke, F. G. N.; Hitchcock, P. B.; Love, J. B.; Wainwright, A. P. *J. Organomet. Chem.* **1995**, 503, 333.
(15) Preparation of **1** proceeds via reaction of Me₃SiN(CH₂CH₂-
NLiSiMe₃)₂ (N₃Li Me $_3$ SiCl with HN(CH $_2$ CH $_2$ NH $_2$) $_2$ in hexane, followed by removal of the

amine hydrochloride byproduct by filtration and reduction to dryness.13 (16) Horton, A. D.; de With, J.; van der Linden, A. J.; van de Weg, H. Unpublished results.

(17) Synthesis of 2: Toluene (20 mL) at -78 °C was added to a mixture of **1** (3.1 mmol) and Mg(CH₂Ph)₂(dioxane)_{0.5} (3.9 mmol). After
the solution was warmed to 25 °C, the solvent was removed and **2** crystallized from pentane at -40 °C (0.84 g, 45%). Synthesis of **3**: To a suspension of **1** (0.41 mmol) in Et2O (10 mL) at -78 °C was added an $Et₂O$ solution of LiMe (0.83 mmol). After the solution was warmed to 25 °C, the solvent was removed and **3** crystallized from hexane at -40 °C (0.104 g, 58%). Synthesis of 5: Toluene (0.5 mL) and bromobenzene (0.1 mL) were added to a mixture of **2** (0.19 mmol) and B(C₆F₅)₃ (0.19 °C for 16 h, hexane (5 mL) addition afforded a precipitate, which was washed with hexane and dried, giving pure 5. Typical NMR experi-
ment: C₆D₃Br (0.7 mL) at –30 °C was added to a mixture of **2** or **3** and B(C_6F_5)₃ (each 0.01 mmol) in a 10 mL bottle at -30 °C, followed by rapid warming to 0 °C and transfer to an NMR tube. Although **4**, **6**, and **9** are formed quantitatively (>95% NMR purity), attempts to isolate the complexes in analytically pure crystalline form have not been successful.

(18) Reaction of 1 with LiMe in $Et₂O$ was recently reported to give an intractable mixture.13

(19) The complexes were characterized by 1H and 13C NMR spectroscopy and (for **2**) elemental analysis. The NMR data supports *η*1 benzyl coordination in **2**: *ipso* C, *δ* 146.8 ppm; ZrCH₂, *δ* 72.1, 64.0
ppm (¹J_{CH} = 120, 124 Hz); *ortho* H, *δ* 6.86, 6.78 ppm.

^X Abstract published in *Advance ACS Abstracts,* May 1, 1996. (1) (a) Sinclair, K. B.; Wilson, R. B. *Chem. Ind. (London)* **1994**, 857.

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R. M. *Angew. Chem., Int. Ed. Engl.* **1995,** 34, 1143.
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N.; Rosen, R. K.; Knight, 416 815 (Dow, 1990). (b) Cannich, J. A. M.; European Patent Application 420 436 (Exxon, 1990). (c) Devore, D. D.; Timmers, F. J.; Hasha, D. L.; Rosen, R. K.; Marks, T. J.; Deck, P. A.; Stern, C. L. *Organome-tallics* **1995**, *14*, 3132.

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rial benzyls and of the dimethylene hydrogens in **2**, observed at 25 °C, presumably involves dissociation of the amino nitrogen, inversion at nitrogen, and recoordination.14 The greater lability of the amino nitrogen in **2** than in dimethyl **3** (two sharp Zr*Me* 1H NMR resonances at 25 °C: *δ* 0.55 and 0.46 ppm) might reflect stabilization of the 4-coordinate species by η^2 -benzyl coordination.19

Abstraction of a benzyl group from 2 with $B(C_6F_5)_3$ in C_6D_5Br or $C_2D_2Cl_4$ solution cleanly affords a soluble product, **4**; ¹⁷ in toluene **4** is deposited as a yellow precipitate. The complex has been shown by ${}^{1}\text{H}$, ${}^{11}\text{B}$, $13C$, and $19F$ NMR spectroscopy²⁰ to consist of a benzylzirconium cation²¹ and a noncoordinated $[PhCH₂B (C_6F_5)_3$ ⁻ anion (Scheme 2).²² The pseudotetrahedral complex is stabilized by η^2 -coordination of the benzyl ligand (giving a maximum 14-electron count at Zr), as shown by the large $^1J_{CH}$ value of 142 Hz for the $ZrCH_2$ group (*δ* 68.7 ppm) and the upfield *ipso* carbon (*δ* 137.5 ppm) and *ortho* hydrogen resonances (δ 6.38 ppm).²³ An identical cation (in addition to free $NMe₂Ph$ or $Ph₃CCH₂$ -Ph) is obtained using $[PhMe₂NH][B(C₆F₅)₄]$ or $[Ph₃C]$ - $[B(C_6F_5)_4].$

Solutions of **4** (with $[PhCH_2B(C_6F_5)_3]$ ⁻ as anion) decompose slowly (16 h, 25 °C), with elimination of 1

equiv of toluene, to give a single organometallic product, 17 which may be precipitated with hexane. $1H$, $13C$, and 19 F NMR spectroscopy²⁰ and elemental analysis have confirmed that orange solid **5** is the product of C-H activation of one of the amido SiMe_3 groups in **4** (Scheme 2). Ligand activation is reflected in the observation of four different Si*Me* resonances (1:1:3:3) and a diastereotopic SiCH₂Zr group (δ 0.70, 0.37, ²J_{HH} $=$ 13.2 Hz). The zirconium cation is stabilized by coordination of the benzylborate anion,²² as shown, in particular, by the downfield *ipso* carbon resonance (*δ* 161.0 ppm; free anion, δ 148.6 ppm) and the large value of $\Delta\delta(m,p\text{-}\mathrm{F})$ of 3.9 ppm (free anion, 2.7 ppm) in the ¹⁹F NMR spectrum.24

In contrast to ionic **4**, the methyl complex **6**, obtained from the reaction of dimethyl **3** with $B(C_6F_5)_3$ in C_6D_5 -Br,¹⁷ exhibits a covalent bonding interaction between the cation, $[N_3ZrMe]^+$, and the $[MeB(C_6F_5)_3]^-$ anion (Scheme 3).25 Anion cordination is reflected in the downfield location of the B*Me* resonance (*δ* 1.38 ppm; free anion *δ* 1.13 ppm) and the large value of ∆*δ*(*m,p*-F) of 4.4 ppm.²⁴ The ZrMe group resonates at δ 0.65 ppm (¹H NMR) and at δ 55.1 ppm (¹³C NMR), slightly upfield of dimethyl **3**.

Analogous methyl cations may be obtained by protonolysis (**7**) or reaction with the trityl reagent (**8**) in C_6D_5 -Br solution (Scheme 3).17,20 Cation **7** is stabilized by NMe2Ph coordination (downfield *ortho* and *para* hydrogen resonances).17,26 For complex **8**, it is not possible to distinguish between possible weak solvent and anion coordination to the zirconium cation. The location of the Zr*Me* resonance in the complexes (**7**, *δ* 0.77 ppm; **8**, *δ* 0.75 ppm), slightly downfield from **6**, may reflect increased metal electrophilicity (compared to the THF adduct of **6**: *δ* 0.48 ppm).

Formation of methyl cations **7** and **8** proceeds via an intermediate species, **9**, cleanly obtained using 0.5 equiv of $[Ph_3C][B(C_6F_5)_4]$ (Scheme 3).^{17,20} Further reaction of **9** with another 0.5 equiv of the reagent affords **8** (20 min, 25 °C). Complex **9** exhibits two Zr*Me* resonances (*δ* 0.51, 2 Me; *δ* 0.48 ppm, 1 Me), as well as resonances for two equivalent and symmetric ${N_3Zr}$ fragments (2:1 ratio of SiMe₃ groups, four backbone methylene hydrogens). These data are consistent with a structure in which a methyl group bridges two $[N_3ZrMe]^+$ fragments, similar to recently reported metallocene complexes, $[{Cp'}_2 ZrMe}_{2}(\mu$ -Me)]⁺²⁷ Again, it is clear that a methyl group of a group 4 dimethyl complex can act as an effective donor for an electrophilic metal center.

Preliminary reactivity studies of the cationic complexes have revealed moderate ethene polymerization activity but very low propene activity. A 20-fold excess of ethene is rapidly polymerized (5 min, 25 °C) by C_6D_5Br solutions of **4** or **6–8** in an NMR tube; the conversion of 20 equiv of propene (to propene oligomers) is incomplete after 15 min. In all cases only a small

⁽²⁰⁾ Selected NMR data (see also text) (¹H NMR, C_6D_5Br , 25 °C, unless otherwise stated) are as follows. **4**: *δ* 7.24 (*p*-BzZr), 7.19 (*m*-BzZr), 3.36 (BC*H*2), 3.17, 2.99, 2.81 (4H, 2H, 2H, NC*H*2), 2.69 (ZrC*H*2), 0.22, -0.21 (18H, 9H, SiMe₃). ¹³C NMR (C₂D₂Cl₄, -35 °C): δ 136.1,
130.5 (2C, *o,m*-BzZr), 129.3 (p-BzZr), 52.7, 49.6 (2C, NCH₂), 32 (1C,
BCH₂), 0.7, -1.2 (6C, 3C, SiMe₃). ¹⁹F NMR: δ -131.63 (d), -1 (t), -167.85 (m). **5**: *δ* 7.2 (*o*,*m*-Bz), 6.68 (*p*-Bz), 3.6-2.2 (NC*H*2, BC*H*2), 0.16, 0.12, -0.02, -0.03 (3H, 9H, 3H, 9H, Si*Me*3). 13C NMR: *δ* 161.0 (1C, *ipso*-Bz), 50.9, 50.1, 50.0, 49.9 (1C, N*C*H₂), 34.5 (Zr*C*H₂), 0.9 (1C,
Si*Me*), -0.6, -0.7, -1.1 (3C, 3C, 1C, Si*Me*). ¹⁹F NMR: δ -131.96 (d),
-162.09 (t), -165.98 (m). **6**: δ 3.65 (2H, NC*H*₂), 3.1-2.8 (4 2.47 (2H, C*H*₂), 0.00, −0.01 (18H, 9H, SiMe₃). ¹³C NMR (C₆D₅Br, −25
°C): *δ* 15.7 (BMe). ¹⁹F NMR: *δ* −133.68 (d), −161.26 (t), −165.64 (m). **7**: *δ* 7.24 (*m*-Ph), 7.12 (*p*-Ph), 6.88 (*o*-Ph), 2.69 (s, 6H, N*Me*2). 13C NMR (C2D2Cl4, -25 °C): *δ* 61.5 (Zr*Me*). **9**: *δ* 3.55, 3.16, 2.84, 2.48 (4H, NC*H*2), 0.15, -0.07 (36H, 18H, Si*Me*3).

⁽²¹⁾ Schematic representation of one of two possible trigonal bipyramidal isomers (axial/equatorial groups exchanged) for **4** and **6**-**9** is arbitrary. A single species is observed spectroscopically.

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⁽²⁴⁾ The value of ∆(*m,p*-F) (19F NMR) is a good probe of coordination of $[RB(C_6F_5)_3]$ ⁻ $(R = Me, CH_2Ph)$ to cationic d⁰ metals (values 3-6 ppm indicate coordination; <3 ppm indicates noncoordination): Horton, A. D. Unpublished results.

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Scheme 3

amount of the cation reacts (giving unidentified products), reflecting slower initiation than propagation in polymerization. In an autoclave experiment, benzyl cation **4** (0.2 mmol, 7.1 bar ethene, 200 mL toluene, 25 °C, 1 L autoclave) afforded 11.0 g of polyethene in 10 min (rate 3600 g/g of Zr·h; GPC analysis, $M_w = 230000$, $M_n = 52200$. The deviation of the polydispersivity (M_w / $M_n = 4.4$) from the ideal uniform site value of 2 is partly ascribed to the nonisothermal nature of the polymerization (exotherm of $+55$ °C within 90 s). A similar reaction with propene (6.6 bar, 60 min) gave 0.45 g of

atactic propene oligomers (${}^{1}H$ NMR: $CH_{2}=C(R)$ Me end groups, $M_n = 5600$, corresponding to a rate of 20 g/g Zr'h.

The new tridentate diamide ligand affords a relatively robust framework for cationic group 4 alkyl complexes. Although the ligand is subject to degradation by $C-H$ activation of a SiMe₃ group adjacent to zirconium, the reaction is much slower than that found by us for other amide ligands.28 Similar to the situation for cationic alkylmetallocenes, the cationic *η*2-benzyl adduct of the tridentate ligand is a rather weak electrophile,^{23b} whereas the methyl cation coordinates the $[MeB(C_6F_5)_3]$ ⁻ anion, $NMe₂Ph$, or even $N₃ZrMe₂$. This work has shown that cationic diamide systems can function as effective ethene polymerization catalysts. The low propene reactivity may reflect a general inertness of fourcoordinate diamide cations toward coordination and insertion of alkenes larger than ethene. In other studies, we have found that three-coordinate analogues exhibit higher activity toward propene.²⁸

Supporting Information Available: Text and a table providing full details of the preparation, characterization, and alkene reactivity of the compounds (11 pages). Ordering information is given on any current masthead page.

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