Phenylene-Bridged OSSO-Type Titanium Complexes in the Polymerization of Ethylene and Propylene

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Supporting Information

ABSTRACT: The dichloro titanocene complexes (OSSO_Bu)_2TiCl_2 (1) and (OSSO_Cum)_2TiCl_2 (2) bearing o-phenylene-bridged OSSO-type ligands [OSSO_Bu-H = 6,6’-(((1,2-phenylenebis(sulfanediyl))bis(methylene))bis(2,4-di-tert-butylphenen)) and OSSO_Cum-H = 6,6’-(((1,2-phenylenebis(sulfanediyl))bis(methylene))bis(2,4-bis(2-phenylpropan-2-yl)phenen)] were prepared and characterized. The X-ray structure of 1 revealed that Ti atom has an octahedral coordination geometry with an fac–fac wrapping of the [OSSO] ligand. In solution at 25 °C, 1 mainly retains the C2 symmetric structure, whereas 2 shows an equilibrium between C2- and C2-symmetric stereoisomers. Activation of 2 with (Ph3C)[B(C6F5)4] led to a highly active catalytic system with an activity of 238 kgPE·molcat⁻¹·bar⁻¹·h⁻¹; linear polyethylene with a Tm of 122 °C and Mw of 107 kDa were obtained under these conditions. Catalyst 1 displayed the moderate activity of 59 kgPE·molcat⁻¹·bar⁻¹·h⁻¹. Gel permeation chromatography analysis revealed the formation of high-molecular-weight polyethylenes with very large distributions of the molecular weights, indicating a low control of the polymerization process, probably because of the presence of different active species in solution. Density functional theory investigation provides a rational for the relative high-molecular-weight polymers obtained with these complexes. The precatalyst 2 was also active in propylene polymerization producing atactic oligomers terminated with unsaturated end groups.

INTRODUCTION

The last decades have witnessed the development of new nonmetallocene early-transition metal complexes as catalyst precursors for olefin polymerization. Among these, group 4 complexes of polydentate ligands, based on a bis(phenolate) framework with additional donors, are particularly acknowledged. Indeed, Fujita and Coates reported on bis(phenoximine) group 4 complexes that are very active in the ethylene polymerization and produce syndiotactic polypropylene under living conditions. Later on, Kol et al. developed a zirconium complex coordinated by a diamine bis(phenolate) [ONNO]-type ligand that is able to promote the living polymerization of 1-hexene with high isospecificity. Since then, several group 4 metal complexes featuring [ONNO]-, [OSSO]-, [OOOO]-, and [OPPO]-type tetradentate ligands have been reported in literature. Among these examples, the family of group 4 complexes bearing the bis(phenolate) ligands with soft-donor sulfur atoms has shown some peculiar features. As a matter of fact, the titanium complexes with 1,4-dithiabutanediyl-linked [OSSO]-type ligands are highly active precatalysts not only for the isospecific polymerization of styrene but also have shown a unique behavior in the homo- and copolymerization of diene monomers. Kol et al., following these results, introduced a methylene group between the S donor and the arene ring, obtaining an improvement of the activity in 1-hexene polymerization with respect to the analogous salan-type complex but with a concomitant loss of stereocontrol. More recently, high isoselectivity in 1-hexene polymerization was achieved by Ishii et al. using a zirconium complex with an [OSSO]-type ligand featuring both methylene and trans-1,2-cyclooctanediyl spacers in the molecular backbone. These results clearly show that the activity is profoundly affected by the dimension of the ring. Notably, it resulted that both an increase or a decrease of the number of carbon atom with respect to the cyclooctanediyl ring leads to a decrement of the catalytic activity. To further explore these structure–activity relationships in the OSSO ligands with methylene spacers, we turned our attention to OSSO ligands with the phenylene bridge. Moreover, we focused on titanium complexes because this

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metal is a cost-effective alternative to the other heavier group IV elements. Herein, we report on the synthesis and characterization of new titanium complexes, supported by o-phenylene-bridged bis(phenolato) ligands and on their activity as catalyst precursors in ethylene and propylene polymerization.

## RESULTS AND DISCUSSION

### Synthesis and Crystal Structure.

The preligands OSSO_{Bu}H and OSSO_{Cum}H [OSSO_{Bu}H = 6,6’-((1,2-phenylenebis(sulfanediyl))bis(methylene))-bis(2,4-di-tert-butylphenol) and OSSO_{Cum}H = 6,6’-((1,2-phenylenebis(sulfanediyl))bis(methylene))-bis(2,4-bis(2-phenylpropan-2-yl)phenol)] were synthesized following the published procedure, that is, by reaction between the appropriate 2-arylphenol) with OSSO tetrachloroethane, TCDE, as the solvent), a unique set of the natural range observed for this kind of angles. The two chlorides are far from each other with an angle of 104.89(6). The Ti-O angle falls in the normal range observed for this kind of angles. The two chlorides are far from each other with an angle of 104.89(6).

### Solution Structures.

The coordination of a tetradeinate ligand to a metal center can produce three octahedral stereoisomers, designated as mer–mer (trans), fac–fac (cis-α), and fac–mer (cis-β) belonging to the C_{2v}, C_{2h}, and C_{1} point groups, respectively. Cause of the weak bond between the hard titanium center and the soft sulfur atom, [OSSO]−Ti complexes have a stereochemically nonrigid coordination geometry. When dissolved in solution, they may display fluxional processes and/or equilibria between the different stereoisomers. The number of species in solution, their geometry, and the exchange processes of 1 and 2 were investigated with the aid of NMR spectroscopy.

In the 1H NMR spectrum of 1 (1,2-dideutero-1,1,2,2-tetrachloroethane, TCDE, as the solvent), a unique set of the resonances was observed in which the methylene protons of the coordinated ligand resonates as an AB pattern at 4.35 and 4.93 ppm, as expected for diastereoprotic protons in a C_{1}-symmetric environment (Figure S4, Supporting Information). This indicates that, in solution, complex 1 retains the same fac–fac ligand wrapping to the metal center observed in the solid state. The heating of the solution up to 50 °C produced only a line broadening of the signals. Differently, when the solution of 1 was cooled at subambient temperature, a second pattern of resonances was observed. The methylene protons of this new isomer gave rise to four doublets at δ 4.93, 4.46, 4.32, and 4.00 ppm, suggesting a C_{1} symmetry for the metal complex. The ratio between the C_{1} and C_{2} isomers is 1:4 at

![Figure 1. Oak Ridge thermal ellipsoid plot drawing of complex 1 (30% thermal ellipsoids). Selected bond lengths [Å] and bond angles [deg]: Ti−Cl1 = 2.271(2); Ti−Cl2 = 2.278(2); Ti−O1 = 1.835(3); Ti−O2 = 1.864(3); Ti−S1 = 2.601(2); Ti−S2 = 2.603(2); S1−Ti−S2 = 77.88(5); O1−Ti−O2 = 161.09(14); and Cl1−Ti−Cl2 = 104.89(6).](image-url)
The exchange process was investigated by appropriate exchange spectroscopy experiments (1H-1H EXSY) at −40 °C (Figure S5, Supporting Information). EXSY peaks between the signals of the methylene groups in each isomer were easily observed. In addition, a second series of less intense cross-peaks correlating the signals of the methylene groups of the two isomers were also detected. These peaks testify a rapid Δ−Λ enantiomer exchange for both the C1 and C2 isomers and a slower interconversions between the two geometric C1 and C2 isomers.

For the more steric encumbered complex 2, two sets of signals attesting the presence of the C1 and C2 isomers were observed at room temperature (Figure S7, Supporting Information). It is worth noting that the ratio between the C1 and C2 isomers is 2:1 at 30 °C. The 1H−1H EXSY (at 30 °C) proved that the Δ−Λ enantiomer exchange and the C1 and C2 interconversions occur also in this case (Figure 2).

![Figure 2. Portion of the 1H−1H EXSY NMR spectrum of 2 (τm = 0.300 s, TCDE, 30 °C, and 600 MHz).](image)

This behavior was previously observed for other titanium complexes bearing OSSO ligands containing methylene spacers, that is, ethylene-bridged and cyclooctanediyl-bridged [OSSO]-type ligands. The C1 symmetric isomer was the prevailing isomer for the ethylene-bridged complex (C1/C2 = 1:4), whereas the C2 symmetric isomer was prevailing for the cyclooctanediyl-bridged complex (C1/C2 = 2:1). Variable-temperature NMR analysis indicates a reversible conversion of two isomers at 60 °C for the titanium dichloro complex with ethylene-bridged [OSSO]-type ligand. For the titanium complex with cyclooctanediyl-bridged [OSSO]-type ligand, the NMR spectrum of the mixtures of isomers was unchanged up to 100 °C.

**Polymerization Studies.** Complexes 1–2 were tested in ethylene polymerization after activation with modified methylaluminoxane (MMAO) or Al/Bu3/[Ph3C][B(C6F5)4] in toluene at room temperature. The main results are summarized in Table 1.

Upon activation with MMAO, both complexes led to active polymerization catalysts. Complex 1 showed an activity of 10 kg mol−1 h−1, whereas using complex 2, the activity increases to 42 kg mol−1 h−1 (entries 1 and 2, Table 1). The activity of 2 decreases when the [Al]/[Ti] molar ratio is changed from 1000 to 2000 or 500 (entries 3 and 4, Table 1). The obtained polyethylene (PEs) were scarcely soluble in common aliphatic solvents. For this reason, the polymer products were extracted in boiling xylene and the soluble fractions were characterized by NMR spectroscopy, differential scanning calorimetry (DSC), and gel permeation chromatography (GPC) analysis. All PE samples were linear and showed a melting point of 135 °C. The molecular weight distributions were broad, with very high dispersity indexes (D), probably because of a partial activation of the active species or to degradation of the catalyst when treated with alkyl aluminum excess.

Treatment with Al−Bu3 (Al/Ti molar ratio of 20), followed by reaction with 1 equiv of [Ph3C][B(C6F5)4], afforded ethylene polymerization catalysts with activities higher than those obtained with MMAO. Indeed, in the case of complex 1, an increment of the activity to 59 kgPE molcat−1 bar−1 h−1 was recorded (entry 5, Table 1). Using complex 2, with bulky cumyl substituents on the phenol rings, the effect on the catalytic performance is more pronounced, with a fivefold enhancement of the catalytic activity (compare entries 2 and 6, Table 1). With the activity value of 238 kgPE molcat−1 bar−1 h−1, complex 2 can be rated as a highly active catalyst, according to the scale proposed by Gibson. The comparison between complexes 1 and 2 suggests that increasing the steric hindrance around the metal center leads to a more effective catalytic active species. The DSC analysis of the PEs suggested the formation of highly linear polymers with a melting point of about 122 °C for 2 and of about 130 °C for 1. The NMR analysis confirmed the linear nature of the obtained polymers. The 1H NMR spectrum of PE by 2 shows a singlet because of the polymethylene sequence at δ 1.16 ppm; a unique signal at δ 28 ppm in the 13C NMR spectrum was parallely observed, and no end group was detected (Figure S11, Supporting Information). The molecular weight distributions of PEs are bimodal and the D values are large, probably as a result of different active species (Figure S12, Supporting Information).

Complex 2 was also investigated in propylene polymerization. It resulted moderately active at room temperature with an activity value of 3 kgPP molcat−1 h−1 at 27 °C (entry 7, Table 1) but became inactive at low temperature. The crude products appeared as low melting waxes. In the 1H NMR spectrum signals belonging to unsaturated chain ends were clearly detected (Figure S13, Supporting Information). Two broad resonances at 4.57 and 4.65 ppm and a resonance at 1.60 ppm indicate the presence of vinylidene end group because of β-hydride transfer from the last 1,2 inserted propylene unit. In addition, three complex multiplets at 5.0, 5.4, and 5.8 ppm are due to allylic and cis-2-butenyl end groups. Allylic chain end group arises from the β-methyl transfer to the metal center from the last propylene unit inserted with a 1,2-regiochemistry or from the β-hydride transfer from the methyl of the...
propylene unit inserted with a 2,1-regiochemistry. cis-2-Butenyl end group is due to the \( \beta \)-hydride transfer from the last inserted propylene unit with 2,1 regiochemistry after 1,2 insertion. A minor signal because of the 3-butenyl chain end group is also detected; this group comes from the \( \beta \)-hydride transfer from the methyl of the last inserted propylene unit with 2,1 regiochemistry after 1,2 insertion. The signals diagnostic of these terminations were also detected in the \(^{13}\)C NMR spectrum of the oligomers obtained with 2,1 regiochemistry after 1,2 insertion.

The signals of the microstructure, as revealed by the pattern of signals in the \(^{13}\)C NMR spectrum of the polypropylene oligomers obtained by 2 (Figure 3). Vinylidene (44%) and allyl (39%) end groups are the principal end groups, and cis-2-buteryl and 3-butenyl end groups amount to 14 and 3%, respectively.

The \(^{13}\)C NMR analysis of oligopropylene showed an atactic microstructure, as revealed by the pattern of signals in the methyl pentad region. The regioregularity of the monomer insertion was not elevated. The signals in the ranges of 34.1–35.6 and 14.7–17.2 ppm because of the methylene carbons of the tail-to-tail sequence and methyl carbons of the head-to-head concatenation were observed with approximately the same area giving a total amount of regioirregular sequences of 27%.

**Molecular Modeling Studies.** To shed light on the polymerization mechanism promoted by the title catalytic systems, a density functional theory (DFT) study was carried out. We investigated chain propagation and chain termination, the elementary steps involved in the reaction, using as a model of the active species the cationic \( n \)-propyl complex deriving from the \( C_2 \) isomer of 1 (\( A_{C_2} \)). The energy profile is displayed in Figure 4.

According to the coordination insertion mechanism, the ethylene coordination to the active species leads to the \( \pi \)-complex B, from which both chain propagation and chain termination can start. The coordination energy of ethylene is quite low (−0.7 kcal/mol). The propagation reaction proceeds through the insertion of ethylene into the metal—carbon bond. This step requires overcoming a quite low barrier of 1.3 kcal/mol (Figure 4) and involves the four-membered transition state \( T_{S\beta H T} \) (Figure 5).

The resulting insertion product C is more stable than B by −18.1 kcal/mol. Alternately, the \( \pi \)-complex B can lead to the termination of the growing polymer chain through the \( \beta \)-hydrogen transfer to the monomer (BHT). The process involves the migration of a hydrogen atom in the \( \beta \) position belonging to the alkyl chain to the nearest carbon atom of the monomer. The corresponding transition state \( T_{S\beta CP} \) reminds a bis(olefin) complex, characterized by a strong metal–hydrogen interaction. As a matter of fact, the distance between the metal atom and the hydrogen atom moving from the alkyl group to the monomer was found to be 1.91 Å (Figure 5). The barrier associated with this step is 7.0 kcal/mol. A second termination process is the \( \beta \)-hydrogen elimination (BHE). Lacking the coordinated monomer, one of the hydrogen atom of the alkyl chain in \( \beta \) position can move to the metal center. It is generally reported that BHE is not the preferential chain termination reaction in the ethylene polymerization promoted by OSSO-
type complex reactions because of its substantially higher activation energy compared to the BHT process. For this reason, the barrier for this step was not calculated. The balance between $\beta$-hydrogen transfer and chain propagation has a direct consequence on the catalytic behavior: more competitive the BHT process is, longer polymeric chains are. An estimation of the competition between the two process is given by the different heights of the corresponding activation barriers ($\Delta\Delta E_{BHT-CP}$). The $\Delta\Delta E_{BHT-CP}$ for title catalytic system is 5.7; this value indicates that our catalyst system is prone in producing long polymeric chains in agreement with the experimental results.

It should be noted that the prototypical titanium complex dichloro{1,4-dithiabutanediyl-2,2'-bis(4,6-di-tert-butylphenoxo)}titanium (3), activated with MAO, leads to high-molecular-weight-branched PE. Experimental evidence supported by DFT calculations suggested that the mechanism involves the in situ formation of $\alpha$-olefins by $\beta$-hydride elimination followed by the reinsertion of these $\alpha$-olefins during the propagation step. The insertion and the BHT barriers computed for 3 at the same level of theory were lower than those obtained for our model complex: the insertion barrier was 1.0 kcal/mol, whereas the BHE barrier was 5.9 kcal/mol and the corresponding $\Delta\Delta E_{BHT-CP}$ for 3 is 4.9 kcal/mol. Thus, different polymeric architecture obtained by these catalysts can be addressed to the more competitive termination process found for complex 3 with respect to the catalytic system reported in this study.

The multimodal GPC traces of polymers obtained with our catalytic systems suggest the formation of multiple catalytically active species having different polymerization performances. Even though the origin of multimodality is still not completely clear and it is necessary to use caution in deducing the real active species from the solution structures of the precatalysts, we wonder if the coordination isomerism of the precatalysts could cause the broadening of the molecular weight distribution. Thus, we turn our attention to the other possible geometric isomers of the active species, that is, the isomer with a mer–mer wrapping of the OSSO ligand around the metal center (A$_{C_i}$) and the isomer with an fac–mer wrapping of the OSSO ligand (A$_{C_e}$). A$_{C_i}$ would not lead to insertion because it could only capture an olefin trans to the alkyl group; therefore, it was not further considered. A$_{C_e}$ was successfully optimized and showed very similar energy respect to A$_{C_e}$, but all our attempts to locate the corresponding ethylene coordination adduct were unsuccessful. The potential energy curve resulting from the nearing of ethylene to the titanium atom of A$_{C_e}$ presented a minimum when the ethylene was placed at about 5 Å from the titanium atom, a distance coherent with the formation of a van der Waals adduct, not with a coordination adduct. Thus, A$_{C_e}$ and A$_{C_e}$ should only represent inactive/dormant species that do not influence the distribution of the molecular weights. In light of these observations, the formation of multiple active species would derive from partial transformation of the precatalyst structure by the aluminum cocatalyst.

## CONCLUSIONS

The dichloro titanium complexes 1–2, supported by [OSSO]-type ligands with an $\alpha$-phenylene-bridge, were prepared and completely characterized in solution by NMR spectroscopy. For complex 1, the X-ray crystal structure revealed that the titanium complex has a $C_2$-symmetric conformation with a distorted octahedral geometry about the titanium center. In solution, 1 mainly maintained the $C_2$ symmetric structure while 2 displayed an equilibrium between $C_2$- and $C_1$-symmetric stereoisomers. Upon activation with MMAO, these complexes were found to be active for ethylene polymerization. The use of [Ph$_3$C][B(C$_6$F$_5$)$_4$], in the presence of AlBu$_3$ as an alkyllating agent and a scavenger, produced much more active catalysts; in particular, complex 2 showed a high activity of 238 kgPE·mol$_{cat}$⁻¹·h⁻¹. Linear high-molecular-weight PEs were obtained with broad dispersity indexes, probably because of the presence of different active species in solution.

DFT investigation on the elementary steps of the polymerization process revealed that the $\beta$-hydrogen transfer to the monomer, the principal termination process, is not competitive with respect to propagation, explaining the production of long polymer chains. The $\Delta\Delta E_{BHT-CP}$ values for our catalytic system are higher than that computed for the prototypical titanium complex 3, in agreement with the different polymeric architecture obtained by these catalysts. Moreover, we found that the multimodal molecular weight distribution cannot be addressed to the coordination isomerism of the precatalysts as only the active species deriving from the $C_2$ isomer is catalytically active, whereas the other possible species are inactive/dormant species.

The precatalyst 2 was also active in propene polymerization at room temperature, producing atactic oligomers terminated with unsaturated end groups.

## EXPERIMENTAL SECTION

### Materials and Methods

The description of the general procedures, and instruments employed is provided in the Supporting Information. The compounds 6,6'-(1,2-phenylenebis(sulfanediyli))bis(methylene)bis(2,4-di-tert-butylphenol) (OSSO)$_{\text{no-H}}$ and 6,6'-(1,2-phenylenebis(sulfanediyli))bis(methylene)bis(2,4-bis(2-phenylpropan-2-yl)phenol) (OSSO)$_{\text{cyc-H}}$ were synthesized following the literature procedure.

### Synthesis of the Complex (OSSO)$_{\text{no-H}}$TiCl$_2$ (1)

A solution of TiCl$_4$ (0.86 mmol, 90 μL) in toluene (4 mL) was added dropwise to a toluene solution (6 mL) of OSSO$_{\text{no-H}}$H (0.50 g; 0.86 mmol) at 25 °C. The mixture was left under stirring for 2 h, and the solvent was distilled off in vacuum.
red residue was crystallized from acetonitrile at 25 °C (0.41 g, 82%). Spectroscopic data: 1H NMR (CD2Cl2, 25 °C, 300 MHz): δ 7.34 (br s, 2H, Ar), 6.60 (br s, 2H, Ar), 6.49 (br s, 2H, Ar), 6.09 (2H, Ar), 4.39 (d, JHH = 13.4 Hz, 2H, CH−H), 3.48 (br sd, 2H, CH−H), 1.89 (s, 18H, C(CH3)3), 1.03 (s, 18H, CH3). 13C NMR (CD2Cl2, 25 °C, 600 MHz): δ 209.2 (s, 2H, Ar), 7.29 (br s, 2H, Ar), 7.14 (br s, 2H, Ar), 6.25 (br s, 2H, Ar), 4.50 (d, JHH = 13.1 Hz, 2H, CH−H), 3.73 (d, JHH = 13.1 Hz, 2H, CH−H), 1.59 (s, 18H, CH3), 1.06 (s, 18H, CH3). 

**Synthesis of the Complex (OSSO\textsubscript{4}TiCl\textsubscript{4} (2).** A solution of TiCl\textsubscript{4 (1.11 mmol, 120 μl) in toluene (8 mL) was added dropwise to a toluene solution (12 mL) of OSSO\textsubscript{4}H (0.92 g; 1.11 mmol) at 25 °C. The mixture was stirred for 2 h, and the solvent was distilled off in vacuum. The red residue was crystallized from acetonitrile at 25 °C (0.81 g, 82%). Spectroscopic data: 1H NMR (CD2Cl2, 25 °C, 300 MHz): δ 7.45 (s, 2H, Ar), 7.42~7.05 (m, Ar), 6.97 (m, Ar), 6.85~6.70 (m, Ar), 6.68 (s, Ar), 6.60 (t, Ar), 6.21 (d, Ar), 6.14 (br s, Ar), 5.07 (d, JHH = 15.1 Hz, CH−H), 4.45 (d, JHH = 15.1 Hz, CH−H), 4.22 (d, JHH = 12.0 Hz, CH−H), 3.48 (d, JHH = 12.0 Hz, CH−H), 3.08 (d, JHH = 11.3 Hz, CH−H), 2.08 (s, CH3), 2.01 (s, CH3), 1.87 (s, JHH = 12.0 Hz, CH−H), 1.78 (s, CH3), 1.74 (s, CH3), 1.68 (s, CH3), 1.65 (s, CH3), 1.63 (s, CH3), 1.37 (s, CH3), 1.28 (s, CH3), 1.12 (s, CH3). 13C NMR (CD2Cl2, 25 °C, 75.5 MHz): δ 136.3, 135.2, 131.8, 126.2, 124.7, 122.9, 43.2, 35.8, 34.7, 31.5, 30.9. Anal. Calcd for C\textsubscript{69}H\textsubscript{50}Cl\textsubscript{2}O\textsubscript{15}S\textsubscript{4}Ti: C, 71.29; H, 6.02; S, 6.72. Found: C, 71.29; H, 6.02; S, 6.72.

**Procedure for Ethylene Polymerization Using MMAO.** Ethylene polymerizations were carried out in a 250 mL Buchi glass reactor equipped with a mechanical stirrer. In a typical procedure, the reactor vessel was charged with a toluene solution of MMAO (0.1 M, 90 mL), pressurized with ethylene and vented three times. After the mixture was thermostated at the required temperature and 1 atm of ethylene, a toluene solution (10 mL) of precatalyst (10 μmol) was injected into the reactor, and the monomer was rapidly charged to the prescribed pressure. After the specific reaction time, the gaseous monomer was evacuated and the polymerization mixture was poured in ethanol acidified with aqueous HCl. The solid polymer was collected by filtration, washed with fresh ethanol, and dried in a vacuum drying oven at 80 °C. In the case of the polymerization promoted by 2, oligomers were extracted from the oligomerization reaction with CHCl\textsubscript{3} and dried on MgSO\textsubscript{4}. The excess solvent was distilled off at reduced pressures.

**Computational Details.** DFT calculations were performed with the Gaussian 09 program suite. All geometrical optimizations were performed at the BP86 level, that is, employing the exchange–correlation functionals of Becke and Perdew, respectively. The electronic configuration was described with the quasi-relativistic LANL2DZ effective core potential for Ti and S, whereas we adopted the SVP basis set for O, C, and H. To save computational resources, the alkyl groups of the OSSO ligand in ortho position were modeled with the tert-butyl groups, whereas the alkyl groups in para position were replaced by hydrogen atoms. The structures of transition states were located by applying Schlegel’s synchronous transit-guided quasi-Newton (QST2) method as implemented in Gaussian 09. All geometries were characterized as minimum or transition state through frequency calculations. The differences in energy reported in Figure 4 are in the gas phase without zero-point correction. Cartesian coordinates of all DFT optimized structures are available on request. Structures were visualized by the CYLview program.

**Single-Crystal X-ray Crystallography.** The details of the data collection for (OSSO\textsubscript{4}TiCl\textsubscript{4} (1) are given in the Supporting Information. Crystal data are reported in Table 2.

| Table 2. Crystal Data and Structure refinement for (OSSO\textsubscript{4}TiCl\textsubscript{4} |  |
|---|---|
| empirical formula | C\textsubscript{69}H\textsubscript{50}Cl\textsubscript{2}O\textsubscript{15}S\textsubscript{4}Ti |
| formula weight | 695.66 |
| temperature/K | 296(2) |
| crystal system | monoclinic |
| space group | P2\textsubscript{1}/c |
| a, Å | 14.2494(6) |
| b, Å | 15.1062(7) |
| c, Å | 17.1323(7) |
| α, deg | 90 |
| β, deg | 90 |
| γ, deg | 90 |
| cell volume, Å\textsuperscript{3} | 3617.3(3) |
| Z | 4 |
| ρ\textsubscript{m}, mg m\textsuperscript{-3} | 1.277 |
| μ(Mo Kα), mm\textsuperscript{-1} | 0.530 |
| F(000) | 1472 |
| crystal size mm | 0.15 × 0.10 × 0.10 |
| θ limits, deg | 1.457 to 24.498 |
| Refl. collected, unique (R\textsubscript{int}) | 50482/6012 (0.2416) |
| goodness-of-fit on F\textsuperscript{2} | 1.109 |
| R\textsubscript{1}(F\textsuperscript{2}), wR\textsubscript{2} [F > 2σ(F\textsuperscript{2})] | 0.0581, 0.0830 |
| largest diff. peak and hole, e Å\textsuperscript{-3} | 0.316 and −0.400 |

CCDC 1578280 contains the supplementary crystallographic data for this paper. These data can be obtained free of charge from the Cambridge Crystallographic Data Centre via www.ccdc.cam.ac.uk/data_request/cif.
ASSOCIATED CONTENT
Supporting Information
The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsomega.8b01550.

Description of the general procedures; figures giving 1H NMR spectra of proligands, complexes, and polymers; GPC trace of PE sample; and crystallographic details for complex 1 (PDF)

Crystallographic data for (OSSO\textsubscript{Bu})\textsubscript{2}TiCl\textsubscript{2} (CIF)

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Notes
The authors declare no competing financial interest.

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