Zirconocene-Catalyzed Dimerization of α-Olefins:  
DFT Modeling of the Zr-Al Binuclear  
Reaction Mechanism

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Abstract: Zirconocene-mediated selective dimerization of α-olefins usually occurs when precatalyst  
(η5-C5H5)2ZrCl2 is activated by minimal excess of methylalumoxane (MAO). In this paper, we present  
the results of density functional theory (DFT) simulation of the initiation, propagation, and termination  
stages of dimerization and oligomerization of propylene within the framework of Zr-Al binuclear  
mechanism at M-06x/DGDZVP level of theory. The results of the analysis of the reaction profiles allow  
to explain experimental facts such as oligomerization of α-olefins at high MAO/(η5-C5H5)2ZrCl2  
ratios and increase of the selectivity of dimerization in the presence of R2AlCl. The results of DFT  
simulations confirm the crucial role of the presence of chloride in the selectivity of dimerization.  
The molecular hydrogen was found in silico and proven experimentally as an effective agent that  
increases the rate and selectivity of dimerization.

Keywords: density functional theory; dimerization of α-olefins; polymerization; single-site  
catalysts; zirconocene

1. Introduction

Despite half-century of the fruitful study of zirconocene-catalyzed polymerization of α-olefins,  
some aspects of the reaction mechanism are unclear. One of the clouds obscuring the clear sky of  
Cosse-Arlman cationic coordination-insertion mechanism combined with few termination event  
pathways [1–7] (Scheme 1) is a highly selective dimerization of α-olefins with a formation of  
methylenealkanes [8–13].  

Highly selective dimerization of α-olefins was detected when (η5-C5H5)2ZrCl2 [8–10] or  
Z(η5-C5H4)2ZrCl2 (Z-bridge between Cp rings) [13] have been activated by minimal excess of  
methylalumoxane (MAO). In the first publication of Bergman [9], the 1:1 Al/Zr ratio was used. In our  
experiments, we successfully applied two-stage activation of zirconocene dichlorides by the reaction  
with triisobutylaluminium (TIBA) and 5–10 equivalents of MAO [13]. Given that the reaction of LZrCl2  
with TIBA results in the formation of Zr-Al hydride complexes [14–17], we proposed that the addition  
of minimal amounts of MAO results in the formation of cationic Zr-Al hydride species that catalyze  
dimerization of α-olefins. Relatively stable di-Zr complexes [18] could hardly be seen as prototype  
of catalytic species. Zr-Al complexes can vary in a number of coordinated Al atoms [15,19–27]. To explain  
experimental facts, we recently proposed a binuclear Zr-Al model (Scheme 2) [13,28] involving the
catalytic species that are closely related to discrete ionic pairs containing Zr-(μ-Me)_2AlMe_2 fragments formed at the initial stage of zirconocene dichloride activation by MAO [29–35]. In References [13,28] we assumed that Zr-(μ-Cl)-Al species maintain selective dimerization, whereas Zr-(μ-H)-Al complexes catalyze the formation of higher oligomers.

In our calculations, we considered [(η^5-C_5H_5)_2Zr-H]^+ as a starting stationary point I-0 for traditional cationic mechanism. The formation of π-complex I-1 with propylene molecule was found to be highly exergonic (−17.5 kcal/mol), this complex easily transformed to [(η^5-C_5H_5)_2Zr–C_3H_7]^+ via transition state TS-1 with low activation barrier (2.1 kcal/mol). n-Propyl cationic complexes can be stabilized by additional agostic Zr-H bonding, the results of our calculations demonstrated that β-agostic complex I-1_b is 10.6 kcal/mol more stable than α-agostic complex I-1_a (Scheme 3).

In the present paper, we report the results of DFT modeling of initiation, propagation, and termination stages of α-olefin dimerization and oligomerization catalyzed by cationic (η^5-C_5H_5)_2Zr-Al species derived from R_2AlX (R = Me, iBu; X = H, Cl, Me, Scheme 2) in comparison with traditional cationic mechanism.

2. Results

Propylene as an easiest α-olefin was chosen to minimize the time of calculations. For simulations of the reaction mechanisms, we used Gaussian-09 program package [36] at M-06x/DGDZVP [37,38] level of theory (for details, see Section 4.1). The results of the optimization of stationary points and transition states are provided in section S1 in the Supporting Information.

2.1. DFT Modeling of the Initiation Stage

In our calculations, we considered [(η^5-C_5H_5)_2Zr-H]^+ as a starting stationary point I-0 for traditional cationic mechanism. The formation of π-complex I-1 with propylene molecule was found to be highly exergonic (−17.5 kcal/mol), this complex easily transformed to [(η^5-C_5H_5)_2Zr–C_3H_7]^+ via transition state TS-1 with low activation barrier (2.1 kcal/mol). n-Propyl cationic complexes can be stabilized by additional agostic Zr-H bonding, the results of our calculations demonstrated that β-agostic complex I-1_b is 10.6 kcal/mol more stable than α-agostic complex I-1_a (Scheme 3).
The propensity of \([\eta^5\text{-C}_5\text{H}_5]_2\text{Zr--H}\] to form stable adducts with \(\text{R}_2\text{Al--X} (\text{R} = \text{Me}, \text{iBu}; \text{X} = \text{H}, \text{Cl}, \text{Me})\) has been confirmed by considerable (more than 30 kcal/mol) lowering of the free energies in formation of the starting complexes \(\text{I-0X}\) by the formula \([\eta^5\text{-C}_5\text{H}_5]_2\text{Zr(\mu-H)(\mu-X)AlR}_2\]^+. To compare the relative free energies and free enthalpies of the species that correspond to mononuclear and binuclear mechanisms, this negative change was accounted by the subtraction of \(\Delta G (\text{R}_2\text{AlX})\) and \(\Delta H (\text{R}_2\text{AlX})\) from the values of \(G\) and \(H\) calculated for \(\text{Zr--XAlR}_2\) species. The values of free energies and free enthalpies of stationary points and transition states for both mechanisms relative to \(G\) (I-2\text{a}) and \(H\) (I-2\text{b}) are presented in Table 1.

The coordination of propylene with a formation of \(\text{I-1X}\) is exergonic for \(\text{X} = \text{H}\) and endergonic for \(\text{X} = \text{Cl}\) and \(\text{Me}\). The difference in free energies of the insertion transition state \(\text{TS-1X}\) and \(\text{I-1X}\) is minimal for \(\text{X} = \text{Cl}\), but in general the value of the activation barrier of the formation of alkyl complexes \(\text{I-2X}\) is lower for \(\text{X} = \text{H}\) (Table 1). In the propyl cationic complexes \(\text{I-2X}\), \(\text{Zr(\mu-X)Al}\) coordination retains. Taking into consideration the possibility of the agostic \(\text{Zr-H}\) bonding, we calculated geometries and free energies of four types of \(\text{I-2X}\) and found that the most stable are agostic complexes \(\text{I-2X}\text{a}\) and \(\text{I-2X}\text{bo}\) (Scheme 3). The formation of \(\beta\)-agostic complexes by isobutyl fragment (for \(\text{R} = \text{iBu}\)) was also accounted and found to be insignificant (see section S1 the Supporting Information for details). The complexes \(\text{I-2H}\text{bo}\) (Figure 1) are remarkably stable (−28.0 and −28.1 kcal/mol for \(\text{R} = \text{Me}\) and \(\text{iBu}\), respectively) in comparison with other \(\beta\)-agostic complexes containing \(\text{Zr(\mu-X)Al}\) fragments (\(\text{X} = \text{Cl}, \text{Me}\)). Higher stability of \(\text{I-2H}\text{bo}\) with outside \(\beta\)-H coordination can be explained by the minima of steric distortions driven by \(\text{Zr(\mu-X)Al}\) fragments for \(\text{X} = \text{H}\). Extremely low free energies of \(\text{I-2H}\text{bo}\) and closely related \(\text{I-4H}\text{bo}\) (−39.7 and −42.5 kcal/mol for \(\text{R} = \text{Me}\) and \(\text{iBu}\), respectively) are important factors that allow to explain the results of oligomerization experiments (see Section 2.3).

### 2.2. DFT Modeling of the Propagation Stage and Dimer Formation

Stationary points and transition states for coordination/insertion of the second and third propylene molecules and for chain termination are presented in Scheme 4. For cationic mechanism, the coordination of the second propylene molecule was energetically favorable (−4.4 kcal/mol), the activation barrier of the second monomer insertion \(\Delta G^\# = G(\text{TS-2}) - G(\text{I-3})\) was 10.4 kcal/mol. The calculated activation barrier for chain release via transfer to monomer (TS-3, Scheme 4) was 17.1 kcal/mol. The most stable product of the second monomer insertion was \(\beta\)-agostic 2-methylpentyl zirconocene cation \(\text{I-4b}\). This complex can eliminate 2-methylpentene-1 via \(\text{TS-4}\) with a formation of \(\text{I-5}\) (\(\Delta G^\# = 8.7\) kcal/mol) or coordinate (I-6) and insert (TS-5) the third propylene molecule (\(\Delta G^\# = 8.4\) kcal/mol).
Therefore, the chain propagation in the case of $[\eta^5\text{C}_5\text{H}_5]_2\text{Zr}^{-}\text{C}_6\text{H}_{13}]^+$ was found to be preferable, with the minimal difference in free energies between transition states of chain propagation and chain termination via $\beta$-hydride elimination (0.3 kcal/mol).

![Figure 1](image-url)  
Figure 1. Calculated geometries of I-2H-bo complexes for (a) R = Me and (b) R = $^1$Bu. The distances Zr-H are specified (Å).

Scheme 4. Stationary points and transition states of the propagation stages and termination events.

Within the framework of binuclear mechanism, we failed to find stationary points containing coordinated $R_2\text{AlX}$ and propylene molecules. Hence, the coordination of propylene resulted in loss of Zr-X bonding, and the stationary points I-3 and I-6 as well as transition states TS-2, TS-3, and TS-5 had been common for mononuclear and binuclear mechanisms. The fundamental difference between mononuclear and binuclear mechanisms had arisen at the stage of the formation of 2-methylpentyl complexes I-4X followed by $\beta$-hydride elimination via TS-4X to $\pi$-complexes I-5X.

We optimized the geometries of I-4X, TS-4X, and I-5X for $R = \text{Me}$ and $^1\text{Bu}$. The results of our calculations clearly demonstrated that $\beta$-agostic complexes I-4X_bo have lower free energies in comparison with isomeric $\beta$-agostic complexes I-4X_bi and $\alpha$-agostic complexes I-4X_a (Scheme 4).
The only exception was I-4Cl_a (R = Me) that was 1.2 kcal/mol more stable in comparison with I-4X_bo.

The nature and geometries of the transition states TS-4X deserves separate consideration. In these transition states, Al atoms demonstrated cooperative effect (Figure 2). For R = Me, the values of the activation barriers \( \Delta G^\circ = G(\text{TS-4X}) - G(\text{I-4X}) \) were 17.7 (H), 13.6 (Cl), and 16.2 (Me) kcal/mol. Thus, \( \beta \)-hydride elimination had been the most affected by Me_2AlCl coordination at Zr atom. This pattern was also manifested for R = ^1^Bu (Table 1).

![Figure 2. Calculated geometries of the transition states TS-4X (R = Me): (a) X = H; (b) X = F; (c) X = Cl; (d) X = Me. The distances Zr-H, Al-H and \( \beta \)-C-H are specified (Å).](image)

The comparison of the relative free energies of chain termination transition states TS-4X and chain propagation transition state TS-5 allowed us to make the conclusion that \( \beta \)-hydride elimination is energetically preferable for X = H and Cl. For X = Me, \( \beta \)-hydride elimination and dissociation of \([\eta^5\text{C}_5\text{H}_{5}]_2\text{Zr(C}_8\text{H}_{13})-\text{AlMe}_3]^+\) with subsequent coordination-insertion of propylene molecule being equal. Apart from that, in the case of \([\eta^5\text{C}_5\text{H}_{5}]_2\text{Zr(C}_8\text{H}_{13})-\text{AlMe}^\text{Bu}_2]^+\), the chain propagation was preferable (Table 1).

### Table 1. Calculated free energies and free enthalpies (kcal/mol) of stationary points and transition states for mononuclear and binuclear reaction mechanisms. 1

| Structure | Cationic Mechanism | R = Me | R = ^1^Bu |
|-----------|-------------------|--------|-----------|
|           |                   | R = Me | R = ^1^Bu | R = Me | R = ^1^Bu |
| I-0       | G                 | 27.5   | -8.4      | -4.6    | -2.7  | -11.8 | -6.3  | -3.5   |
| I-0X      | H                 | 40.8   | -9.9      | -6.7    | -2.6  | -11.2 | -8.6  | -6.6   |
| I-1       | G                 | 10.0   | -13.5     | 0.0     | 5.1   | -14.9 | -1.8  | 5.3    |
| I-1X      | H                 | 9.8    | -26.7     | -14.0   | -1.6  | -29.7 | -17.3 | -12.2  |
| TS-1      | G                 | 12.1   | -7.0      | 4.3     | 10.6  | -9.2  | 3.7   | 11.8   |
| TS-1X     | H                 | 11.4   | -21.9     | -10.9   | -4.1  | -25.2 | -12.8 | -6.9   |
| I-2 b     | G                 | 0.0    | -13.1     | -7.0    | -1.6  | -16.3 | -9.8  | -1.4   |
| I-2X bi   | H                 | 0.0    | -26.9     | -19.5   | -11.6 | -28.9 | -23.0 | -15.2  |
| I-2X a    | G                 | 10.6   | -22.1     | -13.0   | -7.8  | -21.0 | -12.7 | -6.1   |
| I-2X a    | H                 | 10.6   | -35.9     | -28.4   | -22.6 | -36.7 | -29.3 | -24.7  |
| I-2 b     | G                 | 0.0    | -28.0     | -10.4   | -9.1  | -28.1 | -11.0 | -6.8   |
| I-2X bo   | H                 | 0.0    | -41.9     | -26.9   | -24.0 | -43.7 | -28.8 | -26.7  |

| I-3       | G                 | -4.4   | -16.3     | -3.2    | 6.0   |
| I-3 b     | H                 | -15.6  | -3.2     |
| TS-2      | H                 | -9.3   |          |
Table 1. Cont.

| Structure | Cationic Mechanism | R = Me | R = ^1^Bu |
|-----------|-------------------|--------|----------|
|           |                   | X=H    | X=Cl    | X=Me    | X=H    | X=Cl    | X=Me    |
| TS-3      | G                  | 12.7   | 12.7    | 12.7    | 12.7   | 12.7    | 12.7    |
|           | H                  | −1.9   | −1.9    | −1.9    | −1.9   | −1.9    | −1.9    |
| I-4_b     | G                  | −12.5  | −27.1   | −19.1   | −12.7  | −28.0   | −22.4   | −13.8   |
| I-4X_bi   | H                  | −26.4  | −52.0   | −45.9   | −38.0  | −55.1   | −49.8   | −43.3   |
| I-4_a     | G                  | −3.0   | −34.8   | −23.6   | −20.7  | −33.9   | −22.8   | −19.0   |
| I-4X_a    | H                  | −15.7  | −61.9   | −51.7   | −48.2  | −62.8   | −53.8   | −50.6   |
| I-4_b     | G                  | −12.5  | −39.7   | −22.4   | −20.8  | −42.5   | −23.7   | −18.7   |
| I-4X_bo   | H                  | −26.4  | −68.3   | −51.8   | −49.1  | −71.0   | −53.9   | −51.6   |
| TS-4      | G                  | −3.8   | −22.0   | −9.9    | −4.6   | −21.4   | −6.5    | 0.3     |
| TS-4X     | H                  | −17.9  | −50.5   | −38.9   | −32.1  | −51.6   | −38.3   | −33.2   |
| I-5       | G                  | −8.2   | −30.5   | −18.6   | −12.3  | −31.2   | −17.4   | −10.4   |
|           | H                  | −19.7  | −56.6   | −45.0   | −37.7  | −59.3   | −45.6   | −40.2   |

\(^1\) Relative to I-2_b.

2.3. Dimerization and Oligomerization of 1-Hexene: Experimental Study

We performed a series of experiments using \((\eta^5-C_5H_5)_2^2ZrCl_2\) as a precatalyst. In these experiments, \((\eta^5-C_5H_5)_2^2ZrCl_2\) (0.1 mmol) was suspended in 200 mmol of 1-hexene and treated with TIBA solution (2 mmol) at 60 °C. After 10 min of stirring (no reaction detected), the calculated amount of MMAO-12 solution was added. The mixtures were analyzed by GC and \(^1\)H NMR spectroscopy after 1, 2, 3, and 4 h. The final results of these experiments are presented in Table 2. We found that after activation of \((\eta^5-C_5H_5)_2^2ZrCl_2\) by 200 eq. MMAO-12 typical oligomerization proceeds (Table 2, run 1), the ratio of dimer, trimer, tetramer, and pentamer of 1-hexene can be interpreted by standard Flory distribution. The same results have been obtained by Kissin [39,40]. In the presence of 100 eq. MMAO-12 (Table 2, run 2), the relative rate of the dimer increased. In the following experiments, the rate of higher oligomers decreased with lowering of MMAO/Zr ratios (Table 2, runs 3–6, see Figure S2 in the Supporting Information). It should be noted that the rate of the reaction had a local minima at MMAO/Zr = 20 (Table 2, run 4).

The addition of R2AlCl resulted in an increase in the relative proportion of dimer. The best results were obtained when Me2AlCl was added (Table 2, run 7). The increasing of Et2AlCl/Zr ratio resulted in slowing down the reaction (Table 2, runs 8–11, Figure S3 in the Supporting Information). We also found that the addition of trimethylalumium (Table 2, run 12) resulted in deceleration of the reaction with a formation of large amounts of higher oligomers. The activity of the catalyst and the selectivity of dimerization substantially increased in the atmosphere of the molecular hydrogen (Table 2, run 13; this issue is discussed below).

\[\text{1 Relative to I-2_b.}\]
Table 2. Zirconocene-catalyzed oligomerization of 1-hexene (in bulk, 60 °C, 4 h).  

| Run | MMAO-12 eq. | R₂AlX eq. | 1-Hexene Conv., % | Product Distribution (wt.%)|
|-----|-------------|-----------|-----------------|----------------------------|
|     |             |           |                 | 2-C₆ | C₁₂ | C₁₈ | C₂₄ | C₃₀ | Higher Oligomers |
| 1   | 200         | 0         | 99              | 1.7  | 53.8| 13.2| 5.3 | 3.2 | 21.7           |
| 2   | 100         | 0         | 98              | 1.9  | 69.5| 7.5 | 3.8 | 2.4 | 14.1           |
| 3   | 50          | 0         | 96              | 2.0  | 75.1| 6.9 | 3.1 | 1.6 | 7.7            |
| 4   | 20          | 0         | 93              | 2.0  | 77.8| 5.2 | 2.3 | 1.1 | 4.8            |
| 5   | 10          | 0         | 97              | 3.4  | 82.3| 5.9 | 1.5 | 0.3 | 3.6            |
| 6   | 5           | 0         | 95              | 3.8  | 81.6| 5.4 | 1.3 | 0.3 | 2.6            |
| 7   | 10 Me₂AlCl, 1 | 98 | 3.6  | 87.4 | 3.2 | 1.1 | 0.3 | 2.5 |
| 8   | 10 Et₂AlCl, 1 | 88 | 3.6  | 78.1 | 2.9 | 1.0 | 0.3 | 2.2 |
| 9   | 10 Et₂AlCl, 2 | 97 | 3.8  | 85.7 | 3.3 | 1.2 | 0.4 | 2.7 |
| 10  | 10 Et₂AlCl, 5 | 59 | 2.0  | 53.2 | 1.1 | 0.3 | <0.1| <0.2|
| 11  | 10 Et₂AlCl, 10 | 31 | 0.6  | 29.1 | 0.3 | <0.1| <0.1| <0.2|
| 12  | 10 Me₃Al, 2  | 61 | 0.9  | 36.6 | 9.6 | 3.3 | 1.4 | 9.2 |
| 13  | 10 Me₂Al, 2  | 99 | 2.8  | 88.8 | 3.2 | 1.4 | 0.6 | 2.1 |

1 Activation by 20 eq. of TIBA for 10 min at 60 °C followed by the addition of given amount of MMAO-12.  
2 The data were obtained by GC, the rate of higher oligomers was determined by the weighing of the residue after stirring the treated reaction mixture at 250 °C and 0.1 Torr.  
3 Oligomerization at 1 bar of the molecular hydrogen.

3. Discussion

Using the data presented in Table 1, we draw free energy profiles for the oligomerization of propylene catalyzed by zirconocene cation and by (η⁵-C₅H₅)₂Zr–XAlR₂ cationic complexes for X = H, Cl, and Me. These profiles are presented in Figure 3a,b for R = Me and iBu, respectively. We calculated the free activation energies for two reaction pathways, namely, chain termination with a formation of 2-methylpentene-1 (vinylidene dimer of propylene) via TS-4 and chain propagation via TS-5. These values were determined as a difference between G (TS-4) or G (TS-5) and free energies of the most stable intermediates I-4. The values of the free activation energies for insertion of the second monomer molecule TS-2, β-hydride elimination after this insertion TS-4, and insertion of the third monomer molecule TS-5 are presented in Figure 3. It is the difference between TS-4 or TS-4X and TS-5 that determines the main direction of the reaction as a selective dimerization or oligomerization.

The first conclusion from the analysis of the reaction profiles (Figure 3) is the preference of β-hydride elimination after insertion of the second molecule of monomer in (η⁵-C₅H₅)₂Zr–XAlR₂ cationic complexes for the case of X = H and Cl. Within the framework of cationic mechanism and for X = Me, oligomerization and dimerization of propylene seem to be equal by the value of the activation barrier. This conclusion allows to explain higher oligomer content at higher MAO/Zr ratios (Table 1, runs 1 and 2) by the formation of the catalytic particles that are mononuclear cationic Zr complex or binuclear Zr complex with Me₃Al. At high MAO/Zr ratios (100 and more) the large excess of MAO plays the role of “R₂AlX sponge”, the ability of MAO to form stable complexes with Me₂AlCl and Me₃Al had been confirmed experimentally [41–44]. Note that the formation of oligomers was detected by Bergman when (η⁵-C₅H₅)₂ZrMe₂ was activated with 10 eq. of MAO instead of (η⁵-C₅H₅)₂ZrCl₂ [10].

When X = Cl, DFT modeling predicted the preferable formation of vinylidene dimers at low MAO/Zr ratios. In our experiments (Table 2, runs 7–9), we detected the highly positive effect of the addition of R₂AlCl to the reaction mixture. Bergman [9,10] demonstrated that the presence of Cl is crucial for high selectivity of dimerization; recently the chemists of Idemitsu [45,46] and our laboratory [13,47] applied the addition of R₂AlCl to reach maximum yields of α-olefin dimers.
Our calculations predicted that hydride complex I-2H_bo is ~17 kcal/mol more stable than I-2Cl_bo and can, therefore, be considered as a typical “dormant site” for the binuclear mechanism. Evidently, the formation of alkyl-hydride Zr-Al complexes is more than likely under the reaction conditions used in our experiments. Moreover, these inactive species can make up the most of Zr-containing particles. Keeping in mind that the metal-hydride agostic interaction makes the main contribution to the stabilization of I-2H_bo, we proposed that the molecular hydrogen can activate this dormant site by the cleavage of agostic Zr-βH bonding. We made optimizations of different structures formed from I-2H_bo and H2 molecule and found a novel stationary point I-2H_H2 (Figure 4). In this complex, the agostic bonding Zr-H is absent that potentially facilitates the formation of stationary points and transition states of the propagation stage which is completed by selective formation of the dimer when X = H.

![Figure 3](image-url). Reaction profiles for zirconocene-catalyzed oligomerization of propylene for mononuclear and binuclear reaction mechanisms for R = Me (a) and sBu (b). The free energies of stationary points and transition states are presented relative to I-2_b. The free activation energies for insertion of the second propylene molecule (via TS-2), β-hydride elimination with a formation of dimer (TS-4), and insertion of the third propylene molecule (TS-5) are presented in color.
In complex I-2H_H2, the molecule of H₂ plays the role of the additional activator. The interatomic distance d(H-H) in I-2H_H2 was only 0.1 Å higher than in hydrogen molecule. This complex may be stabilized with a formation of α-agostic complex I-2H_a_H2 (see S1.9 in the Supporting Information). The free energy of this complex is 12.2 kcal/mol higher than 1-2H_bo. It is evident that the similar complex can be formed from 1-4H_bo. Apparently, the molecular hydrogen can activate the “dormant” β-agostic hydride complexes with an increase in selectivity of the dimer formation. To confirm this assumption, we performed the catalytic experiment at 1 bar of H₂ (Table 2, run 13) and detected 99% conversion of the monomer with 89% dimer selectivity without the formation of saturated hydrocarbons. This experiment has clearly demonstrated the role of the molecular hydrogen as an activator, but not as a reactant.

4. Materials and Methods

4.1. DFT Calculations

The initial cartesian coordinates of the stationary points had been found by PRIRODA program (version 4.0, M.V. Lomonosov Moscow University, Moscow, Russia) [48] using the 3ζ basis. The final optimization and determination of the thermodynamic parameters for stationary points and transition states were carried out using Gaussian 09 program [36] for gas phase at 298.15 K, the root mean square (RMS) force criterion was 3 × 10⁻⁴ kcal/mol Å⁻¹. The M-06x functional [37] and DGDZVP basis set [38,49] were used in the optimizations. As was demonstrated earlier, M-06x functional is one of the most correct functionals for calculations of the free energies in DFT modeling of zirconocene-catalyzed reactions [50]. Transition states were found by energy scanning with sequential changing of key geometric parameters with a step of 0.01 Å followed by Berny optimization and confirmed by intrinsic reaction coordinate (IRC) simulations (see Figure S1 and find *_IRC.gif animation files in the Supporting Information).

4.2. General Experimental Remarks

TIBA (1 M solution in hexane, Merck, Darmstadt, Germany), MMAO-12 (1.52 M solution in toluene, Merck), (η⁵-C₅H₅)₂ZrCl₂ (Merck), and CDCľ₃ (99.8% ²H, Cambridge Isotope Laboratories, Inc., MS, USA) were used as purchased. 1-Hexene (Merck) was stored over Na wire and distilled under argon. The ¹H NMR spectra were recorded on a Bruker AVANCE 400 spectrometer (400 MHz, Bruker, MS, USA) at 20 °C. The chemical shifts are reported in ppm relative to the solvent residual peaks. The distribution of oligomers produced in zirconocene-catalyzed reactions was measured by gas chromatography (GC) method. GC analysis was carried out with a KRISTALL-2000M gas chromatograph (Meta-chrom Ltd., Yoshkar-Ola, Russian Federation) equipped with a SolGel-1ms (60 m × 0.25 mm × 0.25 μm) column and a flame ionization detector. Helium was used as a carrier gas at a rate of 1.364 cc/min and with a split ratio of 73.3:1. The injection temperature was 320 °C, and the

Figure 4. Calculated geometries of I-2H_bo (a) and dihydrogen complex I-2H_H2 (b). The distances Zr–H, Zr–C(Alkyl), and H–H are specified (Å).
4.3. Dimerization and Oligomerization Experiments

1-Hexene (25 mL, 200 mmol) and TIBA (2 mL of 1 M solution in hexane, 20 mmol) were mixed in a two-necked flask prefilled with argon, which was then placed in a thermostated bath with diethylene glycol. After maintaining the external bath at 60 °C for 2 min, zirconocene precatalyst (0.1 mmol) was added to the flask. After 20 min of stirring, given amounts of MMAO-12 (1.52 M solution in toluene) and R₂AlCl (if applied, 1 M solutions in hexane) were added. The reaction probes were analyzed by NMR and GC after 1, 2, 3, and 4 h. After 4 h of the reaction, 1 mL of methanol and 1 mL of water were added, the organic phase was separated, and the residue was extracted with hexane. The combined organic phases were evaporated under reduced pressure, the residue was stirred at 250 °C (0.1 Torr) to remove C₁₂–C₃₀ reaction products and to determine the weight of the higher oligomers.

5. Conclusions

In the present paper, we report the results of DFT modeling of the binuclear mechanism of zirconocene-catalyzed oligomerization of α-olefins (by the case of propylene) that consider the catalytic particles as a cationic species with coordinated R₂AlX fragments (X = H, F, Cl, Me). We have proposed this mechanism earlier [13,28] for specific reaction conditions, when starting zirconocene dichloride precatalyst has been activated by minimal excess of MAO (up to 10 equivalents). The comparison of the reaction profiles for traditional mononuclear [(η⁵-C₅H₅)₂Zr-Alkyl]⁺ and binuclear [(η⁵-C₅H₅)₂Zr-Alkyl(R₂AlX)]⁺ species demonstrated the qualitative difference between mononuclear and binuclear mechanisms. Without R₂AlX coordination, oligomerization is a preferable reaction pathway. If X = H, highly stable β-agostic complexes 1-2X_bo form, and the reactions slow down. If X = Cl, the formation of vinylidene dimers becomes a main direction of the reaction. The transition states of β-hydride elimination TS-4X (X = H, Cl) demonstrate explicit Zr-Al cooperative effect. For X = Me, there is no significant assistance for β-hydride elimination. These results of DFT modeling correlate with the results obtained in the experimental study of 1-hexene oligomerization.

We also found that molecular hydrogen under “low MAO” conditions demonstrates the chemical behavior that is not typical for Ziegler-Natta processes. In the presence of hydrogen, the dimerization accelerates with substantial increase of selectivity, without the formation of the products of hydrogenolysis. We used the “molecular hydrogen” probe in DFT calculations to find and visualize the mechanism of this effect and found that the complex 1-2H_bo can react with H₂ without cleavage of the H–H bond but with the complete loss of β-agostic coordination. Thus, H₂ acts as an additional activator for hydride complex that grows as an active and selective dimerization catalyst.

Obviously, the mechanistic concept proposed and presented in our paper is approximate and something speculative. However, we believe that Zr-Al binuclear mechanistic concept, which takes into account the direct impact of R₂AlX in catalytic process, will be a valuable addition to traditional mechanism of single-site polymerization of α-olefins. The simplicity of the model is attractive to compare in silico the reactivity of zirconium complexes with different ligand environment under industrially important “low MAO” conditions and for the design of prospective zirconocene oligomerization and polymerization catalysts. An extensive experimental and theoretical work in this field is underway in our laboratory.

Supplementary Materials: The following are available online. DFT calculations data: plots of the molecular geometries, energies, and cartesian coordinates for all stationary points and transition states; Figure S1: The dependence of the composition of the reaction mixtures (1-hexene oligomerization in bulk, 60 °C, 4 h) from Al₂(MMAO-12)/Zr ratio; Figure S2: The effect of R₂AlCl on the composition of the reaction mixtures (1-hexene oligomerization in bulk, 60 °C, 4 h, Al₂(MMAO-12)/Zr = 10), in pdf. format. 19 animation files for all transition states and 3 animation files demonstrating the IRC procedure (in gif. format).

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**Sample Availability:** Samples of vinylidene oligomers of 1-hexene are available from the authors.