Revelation of the Nature of the Reducing Species in Titanocene Halide-Promoted Reductions

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Abstract: The fundamental nature of TiIII complexes generated in tetrahydrofuran by reduction of Cp2TiCl2 has been clarified by means of cyclic voltammetry and kinetic measurements. While the electrochemical reduction of Cp2TiCl2 leads to the formation of Cp2TiCl2−, the use of metals such as Zn, Al, or Mn as reductants affords Cp2TiCl and (Cp2TiCl)2 in a mixture having a dimerization equilibrium constant of 3 × 103 M−1, independent of the metal used. Thus, we find it unlikely that the trinuclear complexes or ionic clusters known from the solid phase should be present in solution as previously suggested. The standard potentials determined for the redox couples Cp2TiCl3/Cp2TiCl2−, (Cp2TiCl)2−/(Cp2TiCl)−, Cp2TiCl/Cp2TiCl2−, and Cp2TiF/Cp2TiF2− increase in the order listed. However, the reactivity of the different TiIII complexes is assessed as (Cp2TiCl)2 ≈ Cp2TiCl ≈ Cp2Ti3+ ≫ Cp2TiCl2− in their reactions with benzyl chloride and benzaldehyde. None of the reactions proceed by an outer-sphere electron transfer pathway, and clearly the inner-sphere character is much higher in the case of Cp2Ti3+ than for (Cp2TiCl)2, Cp2TiCl, and in particular Cp2TiCl2−. As to the electron acceptor, the inner-sphere character increases, going from benzyl chloride to benzaldehyde, and it is suggested that the chlorine atom in benzyl chloride and the oxygen atom in benzaldehyde may function as bridges between the reactants in the transition state.

Introduction

Low-valent metal complexes employed as single electron transferring agents are important reagents for promoting organic synthetic transformations. One of these, which has become increasingly popular over the past few years, is bis(cyclopentadienyl)titanium chloride (Cp2TiCl2), promoting highly diastereoselective pinacol coupling reactions, epoxide openings to alkyl radicals, reductions of alkyl halides such as glycosyl bromides and vic-dibromides, Reformatsky additions, and others. The generation of the trivalent titanium species in tetrahydrofuran (THF) may be achieved by direct substitution of two of the chlorides in TiCl3 with metallated cyclopentadiene or by simple reduction of Cp2TiCl3 with metals such as Zn, Al, or Mn as reductants affords Cp2TiCl and (Cp2TiCl)2 in a mixture having a dimerization equilibrium constant of 3 × 103 M−1, independent of the metal used. Thus, we find it unlikely that the trinuclear complexes or ionic clusters known from the solid phase should be present in solution as previously suggested. The standard potentials determined for the redox couples Cp2TiCl3/Cp2TiCl2−, (Cp2TiCl)2−/(Cp2TiCl)−, Cp2TiCl/Cp2TiCl2−, and Cp2TiF/Cp2TiF2− increase in the order listed. However, the reactivity of the different TiIII complexes is assessed as (Cp2TiCl)2 ≈ Cp2TiCl ≈ Cp2Ti3+ ≫ Cp2TiCl2− in their reactions with benzyl chloride and benzaldehyde. None of the reactions proceed by an outer-sphere electron transfer pathway, and clearly the inner-sphere character is much higher in the case of Cp2Ti3+ than for (Cp2TiCl)2, Cp2TiCl, and in particular Cp2TiCl2−. As to the electron acceptor, the inner-sphere character increases, going from benzyl chloride to benzaldehyde, and it is suggested that the chlorine atom in benzyl chloride and the oxygen atom in benzaldehyde may function as bridges between the reactants in the transition state.

However, in our previous study concerning the electrode reduction of Cp2TiX2 (X = Cl, Br, and I), we showed that mixtures of Cp2TiX22−, Cp2TiX, and the dimer (Cp2TiX)2 are generated, with the former being the major species only in the case of X = Cl.15

The metal-based reductions have led to development of important reaction protocols catalytic with respect to the titanium complex, employing stoichiometric amounts of a metal reductant to regenerate the catalyst after electron transfer. Such applications significantly augment the utility of this reagent for organic synthesis, while justifying the development of more elaborate versions for asymmetric synthesis. Despite its synthetic importance, little is known about the mechanism of the electron transfers that it promotes to organic substrates. Information concerning the electronic interaction between the reductant and the substrate in the transition state would undoubtedly be of importance for understanding the reactivity of TiIII reductants in addition to the development of new applications.

A complicating feature is the structure of the species produced from the above-described reduction of Cp2TiCl2 with metals. Previous reports have shown that trinuclear complexes (denoted I in Chart 1) may be isolated in the solid state, depending on the metal employed.13,16 Such complexes have therefore been proposed to be the reducing species in solution accounting at the same time for the high dl-selectivities observed in pinacol couplings of aryl aldehydes in THF.17 On the other hand, ESR and ENDOR studies of I prepared with zinc have provided evidence for the existence of a TiIII cation in a frozen THF solution.
matrix. Preliminary cyclic voltammetric studies also gave no indications of the presence of trinuclear complexes in solution. Finally, a comparative study of the diastereoselectivities obtained in the pinacol coupling of benzaldehyde with metal-reduced Cp2TiCl2 and (Cp2TiCl)2 in the presence of trimethylsilyl chloride allowed Dunlap and Nicholas to propose the dimer to be the true reducing agent in THF. Through a combination of kinetic and electrochemical investigations, we disclose in this article that trinuclear complexes I (or ionic clusters) are not the reacting species in THF despite their isolation and characterization in the solid phase and that common TiIII-based reductants are formed in all cases, independent of the metal used. Our results substantiate the view that (Cp2TiCl)2 and Cp2TiCl are the major reducing species in solution. In addition, we provide characterization of the electron-transferring abilities of TiIII reductants with the two organic substrates, benzyl chloride and benzaldehyde, allowing further insight into the mechanism of the pertinent reduction processes.

Results and Discussions

Cyclic Voltammetry. Cyclic voltammetry studies of the three metal-reduced solutions of Cp2TiCl4 denoted Met–Cp2TiCl2 (Met = Zn, Mn, and Al) display a quite interesting picture as shown in Figure 1. All three voltammograms exhibit similar oxidation peaks at –0.8 V (designated 2/3) and –0.4 V (designated 4) vs Fe3+/Fe2+ (abbreviation for ferrocenium/ferrocene), respect-

Figure 1. Cyclic voltammograms of 2 mM solutions of Zn–Cp2TiCl2 (–), Mn–Cp2TiCl2 (– – –), and Al–Cp2TiCl2 (– • – • –) recorded at a sweep rate of 0.1 V s⁻¹ in 0.2 M Bu4NPF6/THF.

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 processes. On the reverse sweep a reduction peak (designated $\text{Met}^-/\text{Fc}$) appears at ca. $-1.2 \text{ V vs FC/Fc}$, being slightly dependent on which metal considered. The fact that the peak potentials for the three different metal-reduced solutions are the same indicates that the waves originate from identical species and that trinuclear complexes are unlikely to be present in any of these solutions. Clearly, the oxidation potential of $\text{I}$ (see Chart 1) would be expected to differ for the different metals.

The most likely candidates as common Ti$^{III}$-based species are $\text{Cp}_2\text{TiCl}_2^-$, $(\text{Cp}_2\text{TiCl})_2$, $\text{Cp}_2\text{TiCl}$, and $\text{Cp}_2\text{Ti}^{+}$. The structure of the $(\text{Cp}_2\text{TiCl})_2$ dimer well-characterized from the solid state is shown as $\text{II}$ in Chart 1, while $\text{III}$ presents a tentative suggestion for its structure in THF as discussed below. Fortunately, all of these species can be generated either separately or as a mixture (see the Experimental Section). In Figure 2, we have collected cyclic voltammograms of $\text{Cp}_2\text{TiCl}_2$, $\text{Cp}_2\text{Ti}^{+}/\text{PF}_6^-$, and the mixture $\text{Cp}_2\text{TiCl}/(\text{Cp}_2\text{TiCl})_2$ (denoted “$\text{Cp}_2\text{TiCl}$” in ref 15) to be able to compare their characteristics with those of the voltammograms of $\text{Met}^--\text{Cp}_2\text{TiCl}_2$ in Figure 1. In accordance with expectation, $\text{Cp}_2\text{TiCl}_2^-$ with an oxidation peak potential of $-1.2 \text{ V vs FC/Fc}$ is found to be the easiest one to oxidize, whereas $\text{Cp}_2\text{Ti}^{+}$ with a potential of $-0.4 \text{ V vs FC/Fc}$ is the most difficult one. Both $(\text{Cp}_2\text{TiCl})_2$ and $\text{Cp}_2\text{TiCl}$ show intermediate behavior, having essentially the same oxidation peak potentials of $-0.8 \text{ V vs FC/Fc}$; the appearance of the additional peak at $-0.4 \text{ V vs FC/Fc}$ in this case is due to the generation of $\text{Cp}_2\text{Ti}^{+}$ during the cyclic voltammetric sweep. The nice correspondence of these peaks with those recorded for the metal-generated solutions strongly suggests that the origin of peaks $1-5$ can be ascribed to the electrode processes depicted in eqs $1-4$.

1/5: $\text{Cp}_2\text{TiCl}_2^- - e \rightleftharpoons \text{Cp}_2\text{TiCl}_2$  
2: $(\text{Cp}_2\text{TiCl})_2 - e \rightleftharpoons (\text{Cp}_2\text{TiCl})_2^+$  
3: $\text{Cp}_2\text{TiCl} - e \rightleftharpoons \text{Cp}_2\text{TiCl}^+$  
4: $\text{Cp}_2\text{Ti}^+ - e \rightleftharpoons \text{Cp}_2\text{Ti}^{2+}$

It should be emphasized that because cyclic voltammetry is a dynamic technique, the species detected are not necessarily present initially in the solution but may simply be generated during the sweep. This also provides the explanation why the oxidation wave pertaining to $\text{Cp}_2\text{Ti}^{+}$ appears in the voltammogram of $\text{Cp}_2\text{TiCl}/(\text{Cp}_2\text{TiCl})_2$. Thus, one cannot a priori conclude that all four Ti$^{III}$-based molecules are present in the metal-generated solutions. Indeed, we will show below on the basis of an extended analysis involving both cyclic voltammetry and kinetic measurements that the two principal Ti$^{III}$ species are the dimer, $(\text{Cp}_2\text{TiCl})_2$, in equilibrium with its monomer, $\text{Cp}_2\text{TiCl}$, and that the following reactions apply in the description of the metal-based reductions of $\text{Cp}_2\text{TiCl}_2$.

$\text{Zn} + 2\text{Cp}_2\text{TiCl}_2 \rightarrow \text{ZnCl}_2 + 2\text{Cp}_2\text{TiCl}$  
$\text{Mn} + 2\text{Cp}_2\text{TiCl}_2 \rightarrow \text{MnCl}_2 + 2\text{Cp}_2\text{TiCl}$  
$\text{Al} + 3\text{Cp}_2\text{TiCl}_2 \rightarrow \text{AlCl}_3 + 3\text{Cp}_2\text{TiCl}$  
$2\text{Cp}_2\text{TiCl} \rightleftharpoons (\text{Cp}_2\text{TiCl})_2$

One of the prominent features of the $\text{Met}^-\text{Cp}_2\text{TiCl}_2$ solutions is the presence of metal chlorides and their possible involvement in chloride transfer reactions. This explains why the $1/5$ peaks of the $\text{Cp}_2\text{TiCl}_2/\text{Cp}_2\text{TiCl}_2^-$ redox couple in the metal-generated cases are not completely coincident with those obtained for an authentic sample of $\text{Cp}_2\text{TiCl}_2$. As seen in Figure 3, the peaks recorded for $\text{Cp}_2\text{TiCl}_2$ are gradually shifted upon addition of $\text{ZnCl}_2$ to come in closer agreement with those recorded for $\text{Zn}^-\text{Cp}_2\text{TiCl}_2$. At the same time, this is also the origin of the variation in the relative currents of the oxidation waves for the different $\text{Met}^-\text{Cp}_2\text{TiCl}_2$ solutions. Without doubt, the chloride abstraction proceeds with greater ease from $\text{AlCl}_3$ than from $\text{ZnCl}_2$ and $\text{MnCl}_2$ since peak 1 attributed to the oxidation of $\text{Cp}_2\text{TiCl}_2^-$ can only be observed in the former case (see Figure 1). The relative importance of this peak increases upon lowering the sweep rate. Moreover, addition of excess $\text{Bu}_4\text{NCl}$ to any of the solutions results in the transformation of $(\text{Cp}_2\text{TiCl})_2$, $\text{Cp}_2\text{TiCl}$ and $\text{Cp}_2\text{Ti}^{+}$ to $\text{Cp}_2\text{TiCl}_2^-$ as shown in Figure 4.

To substantiate the working hypothesis that $(\text{Cp}_2\text{TiCl})_2$ and $\text{Cp}_2\text{TiCl}$ should be the principal Ti$^{III}$ species present in the

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Figure 2. Cyclic voltammograms of 1.0 mM $\text{Cp}_2\text{TiCl}_2$ (---), 0.4 mM $\text{Cp}_2\text{Ti}^+ (---)$, and 1.5 mM $\text{Cp}_2\text{TiCl}/(\text{Cp}_2\text{TiCl})_2$ (----) recorded at a sweep rate of 0.1 V s$^{-1}$ in 0.2 M Bu$_4$NPF$_6$/THF. Currents are normalized with respect to concentration to facilitate comparison.

Figure 3. Cyclic voltammograms of 2 mM $\text{Cp}_2\text{TiCl}_2$ in the presence of 0 (--), 0.5 (---), and 1 equiv ZnCl$_2$ (----) recorded at a sweep rate of 0.1 V s$^{-1}$ in 0.2 M Bu$_4$NPF$_6$/THF. The sweep could not be extended past $-1.3 \text{ V in the negative direction because of the deleterious adsorption of ZnCl}_2$.
Met–Cp$_2$TiCl$_2$ solutions, a detailed cyclic voltammetric analysis was carried out. The first important observation is that the relative current of the Cp$_2$Ti$^+$ oxidation wave decreases with increasing sweep rate as illustrated in the case of Zn–Cp$_2$TiCl$_2$ in Figure 5. At the same time, the number of electrons involved in the $2/3$ wave goes from $1/2$ toward 1. Experiments carried out at a sweep rate of 1 kV s$^{-1}$ at a glassy carbon ultramicroelectrode confirm this tendency as wave 4 essentially disappears. Thus, it is concluded that Cp$_2$Ti$^+$ is not present in any of the Met–Cp$_2$TiCl$_2$ solutions initially but that it is generated at the electrode surface as a consequence of chemical reactions occurring during the sweep. In a previous electrochemical study on metal-free Ti$^{III}$-based solutions, Laviron et al. suggested that Cp$_2$Ti$^+$ is formed in a fast father–son reaction (eq 9) involving chloride transfer from Cp$_2$TiCl to Cp$_2$TiCl$^+$, with the latter being formed in the electrode process of eq 3.

$$\text{Cp}_2\text{TiCl} + \text{Cp}_2\text{TiCl}^+ \xrightleftharpoons[k_{-9}]{k_{9}} \text{Cp}_2\text{Ti}^+ + \text{Cp}_2\text{TiCl}_2$$

Indeed, the concomitant generation of Cp$_2$TiCl$_2$ in eq 9 is detectable on the reverse sweep (see Figure 1). However, in the present case one should also take into account that the formation of Cp$_2$TiCl$_2$ may occur through competing chloride transfers to Cp$_2$TiCl$^+$ from the metal chlorides generated in eqs 5–7.

The evidence for the existence of the equilibrium reaction, eq 8, involving Cp$_2$TiCl and (Cp$_2$TiCl)$_2$ is provided by voltammograms recorded for Zn–Cp$_2$TiCl$_2$ at various concentrations. As seen in Figure 6, the broad wave appearing at $-0.8$ V vs Fc$^+/Fc$ consists of two processes, with the first one becoming more dominant as the concentration increases. Thus, the first peak 2 is attributed to the oxidation of the dimer, eq 2, and the second peak 3 to the oxidation of the monomer, eq 3, with the relative height being affected by the presence of the equilibrium reaction in eq 8. Assignment of the broad wave to the oxidation of trinuclear complexes is ruled out by the fact that exactly the same type of splitting is observed for Al–Cp$_2$TiCl$_2$ and Mn–Cp$_2$TiCl$_2$ as well as the metal-free Cp$_2$TiCl/(Cp$_2$TiCl)$_2$ solution analyzed previously (see Figure 2).

Another interesting point is that eqs 3 and 9 involving the oxidation of Cp$_2$TiCl followed by the father–son reaction cannot be the only pathway for the generation of Cp$_2$Ti$^+$ during the cyclic voltammetric sweep. Peak 4 is also apparent at high concentrations and low sweep rates, although the oxidation goes almost exclusively through (Cp$_2$TiCl)$_2$ in eq 2 under such conditions. This implies that Cp$_2$Ti$^+$ may originate from the fragmentation of (Cp$_2$TiCl)$_2$ as well. On account of the observations that the reduction of Cp$_2$TiCl$_2$ is detectable on the reverse sweep and that the peak current ratio for the oxidation waves 2 and 4 approaches unity at high concentrations and low sweep rates, the fragmentation is proposed to follow the pathway in eq 10.

$$\text{(Cp}_2\text{TiCl)}^2_+ \xrightleftharpoons[k_{-10}]{k_{10}} \text{Cp}_2\text{Ti}^+ + \text{Cp}_2\text{TiCl}_2$$

Competing chloride transfers from the metal chlorides to (Cp$_2$TiCl)$_2$ are found to be slower processes and thus not considered further (see the Supporting Information).

On the basis of the above information, we propose that a common reaction scheme, a so-called mesh scheme, can be set up for the three Met–Cp$_2$TiCl$_2$ systems (Scheme 1).
In the Supporting Information, detailed versions of this scheme are provided for each metal. The mesh scheme incorporates the four relevant electrochemical processes, eqs 1–4, along with a number of chemical reactions, including eqs 8–10. Among the latter, eq 8 is a key reaction because the equilibrium constant, \( K_8 \), determines the actual composition of the solution while the associated rate constants are essential parameters for describing the compositional changes occurring at the electrode surface during a chemical or electrochemical reaction. The oxidation of both \( \text{Cp}_2\text{TiCl}_2 \) and \( (\text{Cp}_2\text{TiCl})_2 \) leads ultimately to the formation of \( \text{Cp}_2\text{Ti}^{2+} \) and \( \text{Cp}_2\text{TiCl}_2 \). As no peak corresponding to the reduction of \( \text{Cp}_2\text{TiCl}_2 \) can be monitored in the cyclic voltammograms, even at high sweep rates, it is proposed that \( \text{Cp}_2\text{Ti}^{2+} \) reacts rapidly with \( \text{Cp}_2\text{TiCl} \) or in particular metal chlorides to yield \( \text{Cp}_2\text{TiCl}_2 \). Likewise, the absence at low sweep rates of a cyclic voltammetric reduction wave pertaining to \( \text{Cp}_2\text{Ti}^{2+} \) formed upon oxidation of \( \text{Cp}_2\text{Ti}^{+} \) in eq 4 suggests that \( \text{Cp}_2\text{Ti}^{2+} \) will abstract chloride from the metal chlorides. On the other hand, the lack of peak 1 in double sweep experiments for Met = Zn and Mn indicates that the chloride transfer reactions may go the other way around, i.e., the metal chlorides can abstract chloride effectively from \( \text{Cp}_2\text{TiCl}_2^- \) produced upon the electrochemical reduction of \( \text{Cp}_2\text{TiCl}_2^- \) in eq 1. In general, Ti\(^{IV}\)- but not Ti\(^{III}\)-based species thus are capable of abstracting chloride from the metal chlorides with the Al–\( \text{Cp}_2\text{TiCl}_2 \) system presenting a notable exception, as peak 1 in this case is observable.

From a mechanistic point of view it is crucial to determine the exact composition of the solutions. Unfortunately, voltammograms recorded at low sweep rates will be influenced strongly by the chemical reactions preceding or following the electrode processes. For instance, the overall oxidation of the relatively facile \( \text{Cp}_2\text{TiCl}(\text{Cp}_2\text{TiCl})_2 \) equilibrium system will mainly go through the dimer under such conditions, as this is the species most easily oxidized. In principle, it should be possible to outrun all chemical reactions by applying high sweep rates. However, in the present study we found that the largest applicable value for quantitative purposes was about 50 V s\(^{-1}\); at higher sweep rates the 2 and 3 waves broadened and merged, preventing us from measuring the equilibrium constant of eq 8 directly. Other techniques such as chronoamperometry and sampled current voltammetry were considered as well,\(^{21}\) but they did not provide information that was not obtainable by cyclic voltammetry.

All relevant potentials, heterogeneous and homogeneous rate constants, and equilibrium constants have been extracted by digital simulations (using the program DigiSim 3.03)\(^{22}\) of experimentally obtained voltammograms recorded at different sweep rates of 0.1–50 V s\(^{-1}\) and concentrations of 0.17–2.0 mM. For these kinds of measurements, it is important that the number of experimental observations is large to diminish the overall uncertainty. In addition, the whole collection of voltammograms recorded for each solution of Met–\( \text{Cp}_2\text{TiCl}_2 \) has been simulated simultaneously to obtain the most consistent set of results. The main emphasis in the simulation procedure has been to reproduce the relative currents and the peak potentials with less attention paid to the shape of the relatively broad curves. As described in the Supporting Information, the procedure is based partly on a subjective judgment of which values make the better fits because of the large number of variables present.

The adjustable parameters in the simulation procedure include standard potentials, equilibrium constants, homogeneous and heterogeneous rate constants, symmetry factors, and diffusion coefficients. Of these, the last two are considered to be of less importance, and thus they are kept fixed. The values of the standard potentials, \( E^\circ \), and the homogeneous rate constants, \( k_\text{r} \), are very important for the quality of the fits, as they essentially determine the position of the peaks. For \( k_\text{r} \), we find relatively small values less than 0.1 cm s\(^{-1}\); therefore, the

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(20) Usually the presence of at least three redox states as well as chemical states would be required for describing such a mechanism by a mesh scheme. Thus, in the present case one might argue that the mechanistic scheme is of the ladder type, if the \( \text{Cp}_2\text{TiCl}_2^- \) species with its combination of Ti\(^{III}\) and Ti\(^{IV}\) atoms is not considered as representing a distinct redox state (see also: Evans, D. H. Chem. Rev. 1990, 90, 739).


(22) Rudolph, M.; Feldberg, S. W. DigiSim, version 3.03; Bioanalytical Systems, Inc.: West Lafayette, IN.
electrode processes can be considered to be quasireversible for most of the sweep rates employed. Thus, the influence on the peak potentials from homogeneous follow-up or preceding reactions is not as pronounced as if the electrode processes had been reversible. On the other hand, the homogeneous kinetics in terms of the rate constants involved exerts a large influence on the peak currents and the wave pattern.

The fits are acceptable as illustrated in Figures 7 and 8 for Zn–Cp₂TiCl₂. In general, the oxidative sweep is described more successfully than the reductive one, as the simulations seem to overestimate somewhat the reduction wave of Cp₂TiCl₂. This indicates that the products formed during the oxidative sweep are not all transformed to Cp₂TiCl₂ as assumed in the simulations, but that some of them disappear through other routes.

Indeed, it is likely as discussed elsewhere¹⁵ that a side reaction between Cp₂Ti²⁺ and the solvent, THF, may take place. The latter reaction was not incorporated in the simulations as its effect on the determination of the other parameters in any case would be small. The simulations also showed that the father–son reaction involving Cp₂TiCl⁺ in eq 9 will be outrun by chloride transfer reactions, and accordingly, $k_9$ was set equal to the upper limit of $5 \times 10^3$ M⁻¹ s⁻¹ found previously for the Cp₂TiCl(Cp₂TiCl₂) case.¹⁵ A detailed description of the exact model parameters used and a compilation of relevant fits can be found in the Supporting Information.

The essential data extracted from the simulation procedure are collected in Table 1. Note that all potentials listed are referenced against Fc⁺/Fc, but they can easily be converted to

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**Figure 7.** Recorded (−) and simulated (−−−) cyclic voltammograms of 2 mM Zn–Cp₂TiCl₂ at sweep rates of 0.2, 1, 5, and 20 V s⁻¹ in 0.2 M Bu₄NPF₆/THF. The simulation parameters employed can be found in Table 1 and in the Supporting Information.

**Figure 8.** Recorded (−) and simulated (−−−) cyclic voltammograms of Zn–Cp₂TiCl₂ in concentrations of 0.17, 0.34, 0.80, and 1.80 mM at a sweep rate of 1 V s⁻¹ in 0.2 M Bu₄NPF₆/THF. The simulation parameters employed can be found in Table 1 and in the Supporting Information.
Table 1. Relevant Data Extracted from Cyclic Voltammograms Recorded on Met—Cp₂TiCl₂ Solutions

<table>
<thead>
<tr>
<th></th>
<th>Zn—Cp₂TiCl₂</th>
<th>Mn—Cp₂TiCl₂</th>
<th>Al—Cp₂TiCl₂</th>
<th>Cp₂TiCl/(Cp₂TiCl)₂</th>
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<tbody>
<tr>
<td>E°/V vs Fe⁺/Fe⁺</td>
<td>−1.20 ± 0.03</td>
<td>−1.20 ± 0.03</td>
<td>−1.24 ± 0.03</td>
<td>−1.27 ± 0.04</td>
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<tr>
<td>kₜ/cm s⁻¹</td>
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<tr>
<td>kₜ/M⁻¹ s⁻¹</td>
<td>6.7 (2–5) × 10⁻³</td>
<td>3 (2–5) × 10⁻³</td>
<td>5 (2–10) × 10⁻⁴</td>
<td>2 (2–7) × 10⁻⁴</td>
</tr>
<tr>
<td>kₗ/M⁻¹ s⁻¹</td>
<td>2 (1–10) × 10⁻²</td>
<td>1 (0.5–5) × 10⁻⁴</td>
<td>167 (50–1000)</td>
<td>0.03 (10⁻⁴–10⁻³)</td>
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<td>kₗ/M⁻¹ s⁻¹</td>
<td>33 (10–250)</td>
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<td>kₗ/M⁻¹ s⁻¹</td>
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<tr>
<td>kₜ/ M⁻¹ s⁻¹</td>
<td>1.001–10⁻¹</td>
<td>1 (0.5–1.5) × 10⁻⁴</td>
<td>1 (0.01–10⁻⁵)</td>
<td>1 (0.01–10⁻⁵)</td>
</tr>
<tr>
<td>kₜ/ M⁻¹ s⁻¹</td>
<td>3 (1–50) × 10⁻²</td>
<td>3 (1–50) × 10⁻²</td>
<td>3 (1–50) × 10⁻²</td>
<td>3 (1–50) × 10⁻²</td>
</tr>
<tr>
<td>kₜ/ M⁻¹ s⁻¹</td>
<td>0.3 (10⁻⁵–30)</td>
<td>0.3 (10⁻₅–30)</td>
<td>0.3 (10⁻₅–30)</td>
<td>0.3 (10⁻₅–30)</td>
</tr>
</tbody>
</table>

*The tabulated values are the ones providing the best fits, while those given in parentheses describe the intervals of tolerance, as determined from a manual adjustment of the parameters. To minimize the number of adjustable parameters, the uncertainties on the kₗ values were not determined, but in general they can be assumed to be on the order of 50%. The results for Cp₂TiCl/(Cp₂TiCl)₂ are taken from ref 15 (denoted “Cp₂TiCl” therein). Standard potentials are defined as reduction potentials and can be converted to E° by adding 0.52 V. Calculations were performed by programs of a thermochemical cycle. The simulations are found to be largely unaffected by the exact value of kₗ, and accordingly the latter is set equal to the upper limit determined for the Cp₂TiCl/(Cp₂TiCl)₂ system. An upper limit can be determined from simulations; see ref 15. The simulations are found to be very sensitive toward changes in these parameters because of the reversible reactions occurring between aluminum chloride and the different titanium-based species.

SCE by adding 0.52 V. On the basis of a simple comparison of the potential values that exhibit a variation of as much as 0.8 V, the electron-donating abilities of the different Ti₃⁺ species should be in the order Cp₂TiCl⁺ ≫ (Cp₂TiCl)₂ ≫ Cp₂TiCl ≫ Cp₂Ti⁺ from a thermodynamic point of view. However, such a consideration does not take into account that the reactivity of an electron donor is dependent on other factors as well, such as self-exchange reorganization energies, electrostatic work terms, and last but not least, inner-sphere character of the transition state. In fact, the determination of heterogeneous rate constants in the range of 0.008–0.04 cm s⁻¹ for all four species would suggest that the pertinent self-exchange reorganization energies are relatively large. Usually this would imply large intrinsic barriers and thus kinetically slow electron-transfer processes, unless there are favorable interactions present between the donor and acceptor in the transition state.

According to the equilibrium data, the dominant species in all Met—Cp₂TiCl₂ solutions are Cp₂TiCl and (Cp₂TiCl)₂ while there will be contribution from neither Cp₂Ti⁺ nor Cp₂TiCl⁺. It should be noted that the latter is the major species generated in the electrochemical reduction of Cp₂TiCl₂. The value of 3 × 10⁻¹ M⁻¹ found for the equilibrium constant Kₗ is independent of the metal considered. Actually, it is even equal to that obtained for a metal-free solution, showing that the metal chlorides exert no influence on the position of the dimerization equilibrium, although the associated rate constants kₗ and kₗ⁻¹ both increase by a factor of 25 going from Met = Zn to Al. As to the composition of Met—Cp₂TiCl₂ solutions, the distribution of Ti⁺⁺ species in a 0.5 mM solution, for example, will be 0.22 mM Cp₂TiCl and 0.14 mM (Cp₂TiCl)₂ while the corresponding numbers for a 100 mM solution will be 4 and 48 mM. Depending on the exact reaction conditions (catalytic or stoichiometric), either of the two species may thus be the dominant one. In our previous study we showed that Cp₂TiCl⁺ could be generated from Cp₂TiCl and free Cl⁻ with a relatively large formation equilibrium constant of 10⁻⁴ M⁻¹. Still, Cp₂TiCl⁺ will not be present in the Met—Cp₂TiCl₂ solutions, as chloride is firmly bound in the corresponding metal chlorides. Only in the case of Al—Cp₂TiCl₂ is the chloride transfer from AlCl⁺ sufficiently fast (see the Supporting Information) to ensure production of detectable amounts of Cp₂TiCl⁺ during a cyclic voltammetric sweep. On the other hand, Cp₂TiCl⁺ can easily be made the principal Ti₃⁺ species in any of the solutions, if BnNCl is added. The structure of the oxidized dimer (Cp₂TiCl)⁺ is relatively fragile, endowed by a rate constant kₗ of 10⁻³ s⁻¹ for its fragmentation. As shown in our previous study, the latter process can be further accelerated by the presence of a nucleophile such as chloride.

Kinetic Studies. With the aim of characterizing the reactivity and electron-donating abilities of the different Ti₃⁺ species, Cp₂TiCl⁺, (Cp₂TiCl)₂, Cp₂TiCl and Cp₂Ti⁺, kinetic studies were carried out on their reactions with benzyl chloride (BnCl) and benzaldehyde (PhCHO). A description of the experiments involving Cp₂TiCl⁺ and BnCl is provided elsewhere. The measurements were performed by following the overall absorption decay of the Ti₃⁺ species and/or the equivalent buildup of Ti⁺⁺ species using excess of substrate.

Benzyl Chloride. The fact that the Met—Cp₂TiCl₂ solutions are mixtures of Cp₂TiCl and (Cp₂TiCl)₂ complicates the kinetic analysis in the sense that their reactivities have to be studied through a variation of the equilibrium concentrations. According to eq 8 the larger the overall concentration of Met—Cp₂TiCl₂ becomes, the more dominant the dimer will be. In Figure 9, we have collected kinetic traces for the buildup of Ti⁺⁺ using three different concentrations of Zn—Cp₂TiCl₂ in the presence of excess BnCl. The development is distinct in the sense that the overall kinetics becomes slower as the concentration is raised because of the influence from the equilibrium reaction in eq 8. The same behavior is seen for the two other Met—Cp₂TiCl₂ solutions.

The mechanistic basis for carrying out the fitting procedure consists of eqs 8 along with eqs 11–15, involving both Cp2TiCl and (Cp2TiCl)2 as reactive species.

\[
\begin{align*}
\text{Cp}_2\text{TiCl} + \text{BnCl} & \xrightarrow{k_{11}} [\text{Cp}_2\text{TiCl}^+ + \text{Cl}^- + \text{Bn}^*] \rightarrow \text{Cp}_2\text{TiCl}_2 + \text{Bn}^* \\
\text{Cp}_2\text{TiCl} + \text{Bn}^* & \xrightarrow{k_{12}} [\text{Cp}_2\text{TiCl}^+ + \text{Bn}] \rightarrow \text{Cp}_2\text{TiClBn} \\
(\text{Cp}_2\text{TiCl})_2 + \text{BnCl} & \xrightarrow{k_{13}} (\text{Cp}_2\text{TiCl})_2^{+} + \text{Cl}^- + \text{Bn}^* \\
(\text{Cp}_2\text{TiCl})_2 + \text{Bn}^* & \xrightarrow{k_{14}} [(\text{Cp}_2\text{TiCl})_2^{+} + \text{Bn}] \rightarrow \text{Cp}_2\text{TiCl} + \text{Cp}_2\text{TiClBn} \\
\text{Cp}_2\text{TiClBn} + \text{BnCl} & \rightarrow \text{Cp}_2\text{TiCl}_2 + \text{Bn}_2
\end{align*}
\]

The hypothesis is that in the actual electron-transfer step between \(\text{Cp}_2\text{TiCl}\) and \(\text{BnCl}\) depicted in eq 11, the benzyl radical, \(\text{Bn}^*\), is formed along with chloride which will be captured by \(\text{Cp}_2\text{TiCl}^+\) to afford \(\text{Cp}_2\text{TiCl}_2\) as revealed by previous cyclic voltammetric studies.\(^{-15}\) In principle, these two processes might take place concertedly, which is the reason for showing the initial products within brackets. \(\text{Bn}^*\) is further reduced to the benzyl anion, \(\text{Bn}^-\), in a fast second electron-transfer process (eq 12), and as this anion is expected to be a strong ligand we suggest that \(\text{Cp}_2\text{TiClBn}\) is formed (possibly concertedly). The equivalent reactions of \((\text{Cp}_2\text{TiCl})_2\) are depicted in eqs 13 and 14. Formally, the electron transfer from \((\text{Cp}_2\text{TiCl})_2\) to BnCl in eq 13 would produce \((\text{Cp}_2\text{TiCl})_2^{+}\), \(\text{Cl}^-\), and \(\text{Bn}^*\) if it was not for the immediate follow-up reaction taking place between \(\text{Cl}^-\) and \((\text{Cp}_2\text{TiCl})_2^{+}\). According to our previous cyclic voltammetric studies,\(^{-15}\) this reaction will afford \(\text{Cp}_2\text{TiCl}\) and \(\text{Cp}_2\text{TiCl}_2\). The fast reduction of \(\text{Bn}^*\) by \(\text{Cp}_2\text{TiCl}\) shown in eq 14 leads to the formation of \(\text{Cp}_2\text{TiCl}\) and \(\text{Cp}_2\text{TiClBn}\). In the final step, eq 15, \(\text{Cp}_2\text{TiClBn}\) produced in eqs 12 and 14 reacts in a relatively slow process with a second molecule of BnCl to afford \(\text{Cp}_2\text{TiCl}\) and dibenzyl, \(\text{Bn}_2\), as the final products in accordance with experimental observations (see the Experimental Section).

Analysis of the traces recorded was carried out on the basis of the above equations on the assumption that the reactions of eqs 12 and 14 would be so fast that their only influence on the kinetics would be through the reaction stoichiometry. The simulation program Gepasi 3.21 was employed,\(^{-25}\) giving a consistent set of data with the best fits depicted in Figure 9. Attempts to encompass a situation in the simulations, where the reactivity of the dimer would be negligible and the monomer with its free coordination site the only reactive species, lowered the quality of the fits. Furthermore, it was found that the equilibrium constant \(K_8\) in that instance could hardly be larger than 30 M\(^{-1}\), inconsistent with the value of 3 \(\times 10^1\) M\(^{-1}\) determined from the cyclic voltammetric measurements.\(^{-26}\)

In Table 2, we have collected the data extracted from the above measurements. As seen, the reactivity of \(\text{Cp}_2\text{TiCl}\) and \((\text{Cp}_2\text{TiCl})_2\) is largely the same with \(k_{11} = 0.66\) M\(^{-1}\) s\(^{-1}\) and \(k_{13} = 0.80\) M\(^{-1}\) s\(^{-1}\), independent of the metal. A few kinetic experiments were carried out in the presence of 0.2 M Bu4NPF\(_6\) to have exactly the same experimental conditions as in the cyclic voltammetric experiments. It was found that while \(k_{11}\) was more than halved going to 0.30 M\(^{-1}\) s\(^{-1}\), there was only a slight decrease in \(k_{13}\) to give 0.60 M\(^{-1}\) s\(^{-1}\). From an electrostatic point of view, it would indeed be expected that the largest influence should be seen on the smallest species. Still, the effects are small considering that a lowering of the rate constant by a factor of 2.5 only corresponds to an increase in the activation energy of 0.5 kcal mol\(^{-1}\). This also indicates that the charge separation in the transition state should be small.

The kinetic study of the cation \(\text{Cp}_2\text{Ti}^+\) is much simpler since it can easily be generated without any interference from the other two Ti\(^{III}\) reagents by adding TlPF\(_6\) to a solution of metal-free \(\text{Cp}_2\text{TiCl}\)/(\(\text{Cp}_2\text{TiCl})_2\) (see the Experimental Section). The mechanism for the reaction between \(\text{Cp}_2\text{Ti}^+\) and \(\text{BnCl}\) is proposed to follow eqs 16–18.

\[
\begin{align*}
\text{Cp}_2\text{Ti}^+ + \text{BnCl} & \xrightarrow{k_{16}} [\text{Cp}_2\text{Ti}^{2+} + \text{Cl}^- + \text{Bn}^*] \rightarrow \text{Cp}_2\text{TiCl}^+ + \text{Bn}^* \\
\text{Cp}_2\text{Ti}^+ + \text{Bn}^* & \xrightarrow{k_{17}} [\text{Cp}_2\text{Ti}^{2+} + \text{Bn}] \rightarrow \text{Cp}_2\text{TiBn}^+ \\
\text{Cp}_2\text{TiBn}^+ + \text{BnCl} & \rightarrow \text{Cp}_2\text{TiCl}^+ + \text{Bn}_2
\end{align*}
\]

In the initial electron-transfer step (eq 16), \(\text{Bn}^*\) is formed along with chloride, which according to the cyclic voltammetric analysis will coordinate at the Ti\(^{IV}\) nucleus either concertedly or in an immediate follow-up reaction, thereby generating \(\text{Cp}_2\text{TiCl}^+\). The benzyl radical is reduced by \(\text{Cp}_2\text{Ti}^+\) in eq 17, affording \(\text{Cp}_2\text{TiBn}^+\), which upon further reaction with \(\text{BnCl}\) (eq 18) will produce the final product \(\text{Bn}_2\) along with \(\text{Cp}_2\text{TiCl}^+\). The rate constant \(k_{16}\) can easily be determined to be 0.44 M\(^{-1}\) s\(^{-1}\) under pseudo-first-order conditions as shown in Figure 10. This result is also included in Table 2.

A comparison of the reactivity of the different Ti\(^{III}\)-based species shows quite interestingly (see Table 2) that \(\text{Cp}_2\text{Ti}^+\) exhibits almost the same reactivity as \((\text{Cp}_2\text{TiCl})_2\) and \(\text{Cp}_2\text{TiCl}\),


(26) In cyclic voltammetry, Bu4NPF\(_6\) was used as the supporting electrolyte but the presence of this salt in the kinetic experiments did not improve the quality of the fits.
whereas Cp₂TiCl₂⁻ is a much less efficient reagent. Presumably this can be explained by the fact that Cp₂TiCl₂⁻ has no free coordination site so that a binding of the acceptor to the titanium nucleus cannot take place. This leads to an intrinsically slow reaction possessing very little inner-sphere electron-transfer character. On the other hand, for Cp₂Ti⁺ and Cp₂TiCl easy replaceable solvent molecules will occupy the free ligand sites, facilitating a coordination of benzyl chloride during the reaction as illustrated by the structures IV and V in Chart 2. Note that within such structures chloride can readily be transferred as a ligand to the TiIV nucleus upon electron transfer.

In this sense, it is surprising that (Cp₂TiCl₂), having no free coordination site in the assumed symmetric structure II (known from the solid phase, see Chart 1),¹¹ should exhibit the same reactivity as Cp₂TiCl. Previous stereochemical investigations have suggested that a pentavalent coordination is a possibility,²a,j but we think that a more plausible explanation could reside in the dimeric structure of (Cp₂TiCl₂) being so weak that the same sort of coordination as suggested for Cp₂TiCl might take place during the electron transfer (see VI, Chart 2). There is even the possibility that the dimeric structure in THF from the very beginning is of the half-open type depicted as III in Chart 1, thereby leaving room for an easily accessible coordination site. Following the electron-transfer process and the transfer of chloride to the TiIV nucleus, the dimeric structure will be broken completely as outlined in eq 13. It is important to note that within this picture of concerted electron and ligand transfers the species given in the brackets of eqs 11–17 will not exist. Also investigated was whether the formation of any precursor complexes would be detectable in cyclic voltammetry. However, voltammograms of Zn–Cp₂TiCl₂ solutions recorded just after the addition of BnCl gave no such indications, nor did we see changes in the equilibrium ratio of Cp₂TiCl and (Cp₂TiCl₂).

**Benzaldehyde.** A series of kinetic experiments was carried out on the reaction between Zn–Cp₂TiCl₂ (i.e., Cp₂TiCl and (Cp₂TiCl₂)) and PhCHO. This reaction is of great synthetic importance, as it presents a convenient route for accomplishing pinacol couplings.² The two other TiIII-based species, Cp₂Ti⁺ and Cp₂TiCl₂⁻, were also included in this part of the study, although the measurements pertaining to Cp₂Ti⁺ being irreproducible had to be renounced, and the reactivity of Cp₂TiCl₂⁻ was found to be so low that only an upper limit of 0.1 M⁻¹ s⁻¹ could be provided for the rate constant.

The reaction pathway in this case is assumed to consist of an initial electron transfer from Cp₂TiCl (eq 19) or (Cp₂TiCl₂) (eq 20) to benzaldehyde, followed by dimerization of the thus formed ketyl radicals (eq 21).

\[
\text{Cp}_2\text{TiCl} + \text{PhCHO} \rightarrow \text{Cp}_2\text{CITIO} + \text{Ph} \rightarrow \text{Cp}_2\text{CITIO} + \text{Cp}_2\text{TiCl} \quad (19)
\]

\[
(\text{Cp}_2\text{TiCl})_2 + \text{PhCHO} \rightarrow 2 \text{Cp}_2\text{CITIO} + \text{Cp}_2\text{TiCl} \quad (20)
\]

\[
2 \text{Cp}_2\text{CITIO} \rightarrow \text{Cp}_2\text{CITIO} + \text{OTiClCp}_2 \quad (21)
\]

Note that the TiIV species formed upon electron transfer will be strongly coordinated at the oxygen and will not be liberated unless other strong electrophiles such as trimethylsilyl chloride or protons are added to the solution.¹¹ Moreover, it is assumed that the dimeric structure of (Cp₂TiCl₂) will be broken completely as the reaction in eq 20 progresses. In previous studies addressing the stereochemical outcome of pinacol couplings,² the possibility has been put forth that a pentavalent coordination involving two ketyl radicals can take place at the titanium nucleus in either the dimer II² or the trinuclear complex I²a,j without the need to break bonds. Similar complexes have also been proposed, where two of the chlorides in I have been displaced by ketyl radicals.¹¹²c,d The prominent feature of such arrangements is that they provide straightforward explanations for the high dl/meso ratios obtained. In contrast to this view stands the above cyclic voltammetric analysis pointing toward a rather fragile structure of the oxidized dimer (k₁₀ = 300 s⁻¹).
The rate constants obtained are all included in Table 2. While the reactivity of (Cp2TiCl)2 and Cp2TiCl in the case of benzyl chloride was found to be largely the same, this is certainly not the case for benzaldehyde, where the reactivity of (Cp2TiCl)2 surpasses that of Cp2TiCl by at least a factor of 35. On this basis the arrangement of the dimer shown as VII in Chart 2, where one of the TiIII atoms coordinates to the benzyl group in benzaldehyde during the electron transfer, seems to present a plausible structure. Simulation of the reaction schemes for BnCl, eqs 8 and 11–15, and PhCHO, eqs 8 and 19–21 (with $k_{19} = 2 \text{ M}^{-1} \text{s}^{-1}$), using the Gepasi program shows that in the case of a 100 mM Zn–Cp2TiCl2 solution (containing 100 mM of the substrates) 87.5 and 99.5%, respectively, will go through (Cp2TiCl)2. For a 0.5 mM solution, the corresponding percentages are 28.5 and 85.5%. The turnover points are reached at Zn–Cp2TiCl2 concentrations of $2.3 \times 10^{-3}$ and $4.8 \times 10^{-5}$ M for BnCl and PhCHO, respectively, on the assumption that the concentrations of the substrates are 100 mM. With respect to the product formation in the pinacol coupling reaction, it may thus be concluded that (Cp2TiCl)2 will be the species responsible, unless catalytic conditions are employed. On the other hand, this should have no implications for the dl:meso diastereoselectivities as the dimerization step in eq 21 is suggested to be the same for the two pathways involving Cp2TiCl and (Cp2TiCl)2. This point will be further discussed in a forthcoming publication.

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Figure 11. Kinetic traces recorded for the decay of TiIII at $\lambda = 800$ nm in THF for the reactions of 15 mM PhCHO with 1 (−−), 2 (−−−), and 5 mM (−−−−) solutions of Zn–Cp2TiCl2. The initial absorption is set at 0.0148 in all three cases to facilitate comparison. The dotted curves are the best fits based on simulation of eqs 8 and 19–21, providing $k_{19} < 2 \text{ M}^{-1} \text{s}^{-1}$ and $k_{20} = 70 \text{ M}^{-1} \text{s}^{-1}$.

Chart 2. Proposed Structures for the Coordination of Benzyl Chloride to Cp2Ti+ (IV), Cp2TiCl (V) and (Cp2TiCl)2 (VI) and of Benzaldehyde to (Cp2TiCl)2 (VII) during the Electron Transfer Reactions

The kinetic studies on the reactions with BnCl and PhCHO show that the reactivity order of the four principal TiIII species is: (Cp2TiCl)2 ≥ Cp2TiCl ≈ Cp2Ti+ ≫ Cp2TiCl2−. From a thermodynamic point of view, it would be expected that Cp2TiCl− with a 0.8 V lower oxidation potential than that of Cp2Ti+ should be the best electron donor. This indicates that at least the electron-transfer processes involving the three former species must be of the inner-sphere type. In this context, it is interesting to compare the extracted rate constants for the reaction between the different TiIII-based species and BnCl or PhCHO, denoted generally $k_{\text{obs}}$ (i.e., depending on the reaction considered it may correspond to $k_{11}$, $k_{13}$, $k_{16}$, $k_{19}$, or $k_{20}$), with those expected for electron-transfer reactions involving well-characterized electron donors such as aromatic radical anions. In the case of BnCl, the latter rate constants denoted $k_{ET}$ may be deduced from the log $k_{ET}$ vs $E^0$ plot shown in ref 24, where $E^0$ denotes the standard potential of the aromatic compound.

Unfortunately, the same procedure for finding $k_{ET}$ cannot be employed in the case of PhCHO, since the electron transfer from the aromatic radical anions to form the radical anion of benzaldehyde involves an energetically unfavorable equilibrium reaction. We therefore chose to calculate the required $k_{ET}$ values between outer-sphere electron donors with $E^0$ values as listed in Table 1 and benzaldehyde on the basis of the equations of Marcus and Eyring.28 The $E^0$ of benzaldehyde is $-2.37$ V vs Fc+/Fc as measured by cyclic voltammetry, the reorganization response. The same type of measurements could not be carried out for PhCHO because of the fast reactions occurring in eqs 19–21. From the best fits included in Figure 11 it is found that $k_{19} < 2 \text{ M}^{-1} \text{s}^{-1}$ and $k_{20} = 70 \text{ M}^{-1} \text{s}^{-1}$, thus showing that the influence from eq 19, in general, will be rather small.

The rate constants obtained are all included in Table 2. While the reactivity of (Cp2TiCl)2 and Cp2TiCl in the case of benzyl chloride was found to be largely the same, this is certainly not the case for benzaldehyde, where the reactivity of (Cp2TiCl)2 surpasses that of Cp2TiCl by at least a factor of 35. On this basis the arrangement of the dimer shown as VII in Chart 2, where one of the TiIII atoms coordinates to the benzyl group in benzaldehyde during the electron transfer, seems to present a plausible structure. Simulation of the reaction schemes for BnCl, eqs 8 and 11–15, and PhCHO, eqs 8 and 19–21 (with $k_{19} = 2 \text{ M}^{-1} \text{s}^{-1}$), using the Gepasi program shows that in the case of a 100 mM Zn–Cp2TiCl2 solution (containing 100 mM of the substrates) 87.5 and 99.5%, respectively, will go through (Cp2TiCl)2. For a 0.5 mM solution, the corresponding percentages are 28.5 and 85.5%. The turnover points are reached at Zn–Cp2TiCl2 concentrations of $2.3 \times 10^{-3}$ and $4.8 \times 10^{-5}$ M for BnCl and PhCHO, respectively, on the assumption that the concentrations of the substrates are 100 mM. With respect to the product formation in the pinacol coupling reaction, it may thus be concluded that (Cp2TiCl)2 will be the species responsible, unless catalytic conditions are employed. On the other hand, this should have no implications for the dl:meso diastereoselectivities as the dimerization step in eq 21 is suggested to be the same for the two pathways involving Cp2TiCl and (Cp2TiCl)2. This point will be further discussed in a forthcoming publication.27

The kinetic studies on the reactions with BnCl and PhCHO show that the reactivity order of the four principal TiIII species is: (Cp2TiCl)2 ≥ Cp2TiCl ≈ Cp2Ti+ ≫ Cp2TiCl2−. From a thermodynamic point of view, it would be expected that Cp2TiCl− with a 0.8 V lower oxidation potential than that of Cp2Ti+ should be the best electron donor. This indicates that at least the electron-transfer processes involving the three former species must be of the inner-sphere type. In this context, it is interesting to compare the extracted rate constants for the reaction between the different TiIII-based species and BnCl or PhCHO, denoted generally $k_{obs}$ (i.e., depending on the reaction considered it may correspond to $k_{11}$, $k_{13}$, $k_{16}$, $k_{19}$, or $k_{20}$), with those expected for electron-transfer reactions involving well-characterized electron donors such as aromatic radical anions. In the case of BnCl, the latter rate constants denoted $k_{ET}$ may be deduced from the log $k_{ET}$ vs $E^0$ plot shown in ref 24, where $E^0$ denotes the standard potential of the aromatic compound. Unfortunately, the same procedure for finding $k_{ET}$ cannot be employed in the case of PhCHO, since the electron transfer from the aromatic radical anions to form the radical anion of benzaldehyde involves an energetically unfavorable equilibrium reaction. We therefore chose to calculate the required $k_{ET}$ values between outer-sphere electron donors with $E^0$ values as listed in Table 1 and benzaldehyde on the basis of the equations of Marcus and Eyring.28 The $E^0$ of benzaldehyde is $-2.37$ V vs Fc+/Fc as measured by cyclic voltammetry, the reorganization

energy of the reactions is assumed to be 10 kcal mol\(^{-1}\), and for the collision frequency a value of 3 \(\times 10^{11}\) M\(^{-1}\) s\(^{-1}\) is used.\(^{28}\)

In Table 2, the \(k_{obs}/k_{ET}\) ratios are collected for BnCl and PhCHO with values ranging from less than 6.8 \(\times 10^{3}\) to 4.3 \(\times 10^{6}\). Usually, such large numbers would be indicative of substantial inner-sphere character of the transition state,\(^{29}\) but in the present case one should exert great care in doing this kind of assessment, since a thorough analysis should also take into account differences in the reorganization energies, all relevant work terms, as well as the exact association constants for the formation of precursor complexes.\(^{24}\) One could easily imagine that the latter would be of particular importance for the reactions of PhCHO. In addition, the finding of relatively small heterogeneous rate constants in Table 1 indicates that the reorganization energy for the Ti\(^{III}\) species is substantially larger than that for aromatic radical anions. At the present stage, we therefore prefer to develop a relative scale of uncorrected \(k_{obs}/k_{ET}\) ratios, which are still very useful for carrying out comparisons, rather than paying further attention at the absolute values.

For both BnCl and PhCHO, the \(k_{obs}/k_{ET}\) ratios decrease in the order \(\text{Cp}_2\text{Ti}^+\), \(\text{Cp}_2\text{TiCl}\), \(\text{Cp}_2\text{TiCl}_2\), and \(\text{Cp}_2\text{TiCl}_3^-\). This would imply that there should be substantially more inner-sphere character in the reactions of Cp\(_2\)Ti\(^+\) than those of Cp\(_2\)TiCl, (Cp\(_2\)TiCl\(_2\)), and in particular Cp\(_2\)TiCl\(_3^-\). In fact, this is in line with expectations since the cation Cp\(_2\)Ti\(^+\) compared with the anion Cp\(_2\)TiCl\(_3^-\) would be expected to have a stronger coordination in the transition state to the developing Cl\(^-\) in BnCl and O\(^-\) in PhCHO. From both a kinetic and thermodynamic point of view, the features of Cp\(_2\)TiCl\(_3^-\) (\(E^0 = -1.27\) V vs Fc\(^+/Fc\); \(k_{obs} < 0.019\) M\(^{-1}\) s\(^{-1}\) and \(k_{obs}/k_{ET} < 6.8 \times 10^3\) for BnCl) are comparable with those of SnI\(_2\) (\(E^0 = -1.41\) V vs Fc\(^+/Fc\); \(k_{obs} = 0.02\) M\(^{-1}\) s\(^{-1}\) and \(k_{obs}/k_{ET} = 250\) for BnCl),\(^{24}\) although it is hard to carry out a direct comparison of the \(k_{obs}/k_{ET}\) ratios for the reasons given above. Finally, it may be concluded that the inner-sphere character of the processes involving PhCHO is stronger than for BnCl and that this feature is getting more pronounced as the thermodynamic electron-donating abilities of the Ti\(^{III}\) complex are lowered.

**Experimental Section**

**Materials.** Most chemicals were of commercial origin unless otherwise noted. Benzyl chloride and benzaldehyde were vacuum distilled before use. THF was distilled over sodium and benzophenone under an atmosphere of dry nitrogen. Argon (99.99% purity) was passed through a column of P\(_2\)O\(_5\) (Sicapent). Tetrabutylammonium hexafluorophosphate, Bu\(_4\)NPF\(_6\), was prepared from a hot aqueous solution containing tetrabutylammonium hydrogen sulfate and potassium hexafluorophosphate, Bu\(_4\)NPF\(_6\), was prepared from a hot aqueous solution of tetrabutylammonium chloride, Bu\(_4\)NCl, was recrystallized from toluene. The metal-free crystals of TlCl then precipitated. The liquid was removed, and the crystals were washed twice with THF. A large amount of THF was subsequently added, and after 30 min of stirring the solution turned pale blue. The concentration of Cp\(_2\)TiPF\(_6\) in this saturated solution was approximately 2 mM as determined from peak current measurements.\(^{31}\)

**Apparatus.** Most of the electrochemical equipment was home-built, and a description of the experimental setup is provided in ref 32. The working electrode was a glassy carbon disk of diameter 1 mm. The electrode surface was polished using 0.25 μm diamond paste (Struers A/S), followed by cleaning in an ethanol bath. The counter electrode consisted of a platinum coil melted into glass, while a silver wire in a sintered glass containing THF/0.2 M Bu\(_4\)NPF\(_6\) + 0.02 M Bu\(_4\)NI served as the reference electrode. All potentials were reported against the ferrocenium/ferroene (Fc\(^+/Fc\)) redox couple, the potential of which has been measured to be 0.52 V vs SCE in THF/0.2 M Bu\(_4\)NPF\(_6\).\(^{23}\) All handling of chemicals was performed on a vacuum line, and at no point during the different operations was the interference of oxygen allowed. The ohmic drop was compensated with a positive feedback system incorporated in the potentiostat. The kinetic traces were recorded by means of a fiber-optic spectrometer, model S1000 (dip-probe), from Ocean Optics, using a light path length of 1 cm.

**Procedure.** In the cyclic voltammetric experiments, 0.77 g of Bu\(_4\)NPF\(_6\) (2.0 mmol) and a small magnetic bar were added to the electrochemical cell. The cell was closed and flushed thoroughly with argon for 10 min. Typically, 9 mL of freshly distilled THF and 1 mL of the appropriate standard solution containing the compound of interest were added to the cell using a syringe, and the solution was stirred for 30 s. Special care must be taken when recording voltammograms of the Met–Cp\(_2\)TiCl\(_3\) solutions, since the metal ions present may be reduced at low potentials and cause deleterious adsorption of metal on the electrode surface. In general, potentials below −1.3 V vs Fc\(^+/Fc\) should be avoided. At the end of each series of experiments a small amount of ferrocene was added, and the potential of the Fc\(^+/Fc\) couple was measured.

In the kinetic experiments, the UV–vis dip-probe was mounted vertically in a two-necked cell containing a small magnetic bar.\(^{32}\) The cell was closed and flushed with argon for 10 min before 9 mL of freshly distilled THF was added. From a standard solution containing the appropriate Met–Cp\(_2\)TiCl\(_3\), typically 1 mL was transferred to the cell. Benzyl chloride or benzaldehyde was added in excess, while the solution was stirred vigorously. The decay of Ti\(^{III}\) at the wavelength \(\lambda = 800\) nm and the concomitant buildup of Ti\(^{IV}\) at \(\lambda = 535\) nm were followed. Analysis of the products formed in the reaction of Zn–Cp\(_2\)TiCl\(_3\) with benzyl chloride showed (after acidic quenching) formation of both dibenzyl and toluene, the ratio of which increased as the reaction time was prolonged. This indicates that the reaction shown in eq 15 is relatively slow occurring in the time scale of hours.

**Conclusions**

Detailed cyclic voltammetric studies of Zn\(–\), Mn\(–\), and Al-reduced Cp\(_2\)TiCl\(_3\) solutions in THF show that the principal

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Ti$^{III}$ species present are a mixture of Cp$_2$TiCl and (Cp$_2$TiCl)$_2$, independent of the metal considered; the equilibrium constant of dimerization is determined to be $3 \times 10^3$ M$^{-1}$. There are no indications of trinuclear complexes or ionic clusters involving Cp$_2$Ti$^+$ as suggested previously. Nor do we see the formation of Cp$_2$TiCl$_2^-$ as found in the electrochemical reduction of Cp$_2$TiCl$_2$, because chloride now becomes firmly bound by the corresponding metal chlorides. According to the standard potentials determined in cyclic voltammetry the electron-donating ability of the different Ti$^{III}$-based species would be expected to be in the order Cp$_2$TiCl$_2^-$ \(\approx\) (Cp$_2$TiCl)$_2 \gg$ Cp$_2$TiCl \(\gg\) Cp$_2$Ti$^+$. However, in reality the reactivity order is (Cp$_2$TiCl)$_2 \approx$ Cp$_2$TiCl \(\approx\) Cp$_2$Ti$^+ \gg$ Cp$_2$TiCl$_2^-$, as assessed in their reactions with benzyl chloride and benzaldehyde. In general, it is found that (Cp$_2$TiCl)$_2$ will be the species responsible for the product formation, unless catalytic conditions characterized by low concentrations of Ti$^{III}$ are employed. None of the reactions studied in this article proceed by an outer-sphere electron-transfer pathway, and the inner-sphere character of the reactions of Cp$_2$Ti$^+$ is clearly much higher than those of (Cp$_2$TiCl)$_2$, Cp$_2$TiCl, and in particular Cp$_2$TiCl$_2^-$. As for the electron acceptor, the inner-sphere character of the transition state increases going from benzyl chloride to benzaldehyde, and it is suggested that Cl in benzyl chloride and O in benzaldehyde could function as bridges between the reactants in the transition state. In a forthcoming publication$^{27}$ the focus will be on metal-reduced solutions of Cp$_2$TiBr$_2$ and Cp$_2$TiI$_2$ as well as the factors influencing the reactivities and diastereoselectivities of the pinacol coupling reaction involving benzaldehyde.

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Supporting Information Available: Text giving a description of the model parameters used in the simulations and figures giving a compilation of relevant fits (PDF). This material is available free of charge via the Internet at http://pubs.acs.org.

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