Tuning the Stability of Pd(IV) Intermediates Using a Redox Non-innocent Ligand Combined with an Organolanthanide Fragment

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Supporting Information

ABSTRACT: The unique combination of a divalent organolanthanide fragment, Cp*₂Yb, with bipyrimidine (bipym) and a palladium bis-alkyl fragment, PdMe₂, allows the rapid formation and stabilization of a Pd(IV) tris-alkyl moiety after oxidative addition with MeI. The crucial role of the organolanthanide fragment is demonstrated by the substitution of bipym by the 4,5,9,10-tetraazaphenanthrene ligand, which drastically modifies the electronic structure and tunes the stability of the Pd(IV) species.

While organometallic complexes of Pd(IV) and Pt(IV) have been known for several decades,† the expansion of their utility in organic synthesis urges the chemical community to improve or find new methods that allow the easy access of high oxidation intermediates of the late transition metals, such as Pd(III), Pd(IV), Ni(III), Ni(IV), and Cu(III). Besides, there is a growing debate in the scientific community about the electronic structure of these high-valent intermediates, and more examples of well-characterized compounds are required to address accurately their chemical nature.

The first occurrence of isolated organometallic Pd(IV) complexes appeared in 1975, with the oxidation by chlorine of a series of palladium bis-C₆F₅ complexes coordinated by different chelates. A decade later, Canty reported the oxidative addition of dimethylpalladium complexes bearing bipyridine and phenanthroline ligands, leading to complexes of Pd(IV) that were stable enough for isolation. This seminal work largely developed this area, and other tris-methyl complexes of Pd(IV) appeared. Since these developments, their formation mechanism and relative stability have been studied with high interest. Therein, the chelate ligand is crucial, because of its geometric (ligand symmetry, rigidity, dynamic coordination) or electronic (vide infra) role. As such, the fine-tuning of the ligand that allows the stabilization of various reactive intermediates is needed to develop novel reactions.

Recent detailed reports by Bergman and Tilley showed that the coordination of a Lewis acid in the platinum bipyrazine (bpyz) and platinum bipyrimidine (bipym) bis-aryl complexes leads to a dramatic acceleration of the biaryl reductive elimination, implementing an electronic remote control strategy to manipulate the stability and reactivity of group 10 organometals using bimodal ligands. Van der Vlugt et al. reported another strategy based on the tuning of non-innocent ligands, allowing intramolecular electron transfer(s), to achieve homolytic bond cleavage and radical reactivity at a Pd(II) metal center, instead of the usual two-electron reactivity.

In this context, the comprehensive electronic studies made on different N-heterocyclic adducts of divalent lanthanides demonstrated that their ground state is the result of multiple factors. Thus, Cp*₂Yb(bipy) is a multi-configurational singlet composed of an open-shell (f₁³, L⁺⁻) and a closed-shell (f₁⁴, L⁺⁺) state in a 0.83:0.17 ratio, while Cp*₂Yb(phen) is a triplet state (f₁⁴, L⁺⁺). A chemical ramification of these observations is that the bipy adduct is stable while the phen adduct dimerizes, featuring a reversible C–C bond. This dichotomy was rationalized by the symmetry of the molecular orbital of the ligand (C₂ᵥ symmetry) that is getting populated: the b₁ orbital in the bipy complex has large coefficients on the N-atoms, allowing for easy magnetic communication. In contrast, the a₁ orbital in the phen complex has small coefficients on the N-atoms, discouraging magnetic communication.

Knowing this, we thought it would be possible to tune the relative stability of group 10 organometals using organolanthanides bearing different redox non-innocent ligands. In this Communication, we wish to present the synthesis and chemical structure analysis of a new type of organometals that combine a divalent lanthanide, Cp*, Yb, and a transition metal, PdMe₂, bridged by redox non-innocent ligands, bipym and 4,5,9,10-tetraazaphenanthrene (taphen), as the analogues of bipy and phen. The formation and stabilization of a Pd(IV) tris-alkyl moiety is discussed as a proof of concept.

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The bipym and the taphen bis-methylpalladium complexes, (bipym)PdMe₂ (1) and (taphen)PdMe₂ (2), were synthesized from (tmeda)PdMe₂ in the (Scheme 1). Both complexes produced red needles suitable for X-ray diffraction (XRD) analysis (see Supporting Information (SI)). Complexes 1 and 2 were combined with Cp*₂Yb(OEt)₃ in toluene to yield the heterobi-metallic complexes Cp*₂Yb(bipym)PdMe₂ (3) and Cp*₂Yb(taphen)PdMe₂ (4), respectively. Both 3 and 4 are dark toluene solutions from which dark brown blocks of 3 and dark purple needles of 4, suitable for XRD studies, were isolated (Figure 1).

The Cp*—Yb average distances in 3 and 4 are similar (2.025(8) and 2.29(1) Å, respectively) and are indicative of the oxidation state of the Yb center. Typical values of 2.30–2.31 Å are observed for YbII complexes, whereas longer Cp*—Yb distances (2.43–2.50 Å) are indicative of YbIII complexes. They oscillate between these limiting values when the electronic ground state is multiconfigurational. Within this picture, the Cp*—Yb distances in 3 and 4 would better agree with YbIII complexes. This means that an electron has been transferred to the ligand. In 3, this can be further noticed by comparing the C—C distance that links the two pyridine heterocycles with that of the neutral bipym in 1: 1.417(2) vs 1.491(6) Å, respectively. A reduced distance is indicative of a ligand-based reduction, as noted in the recent work of Wieghardt et al. In comparison, the mono-reduced bipy ligand features an average short C—C distance of 1.42(1) Å. In 4, a similar trend is observed when comparing the diazo distance in 2 (1.295(5) Å) and that in 4 (1.38(1) Å). An increased distance is indicative of a reduction. The Pd—N(ave) and Pd—C(ave) distances in 3 (2.129(4) and 2.022(3) Å) and 4 (2.174(4) and 1.990(3) Å) are comparable to those in 1 (2.135(5) and 2.028(5) Å) and 2 (2.152(5) and 2.028(8) Å), indicating that the coordination environment of the Pd center is only slightly affected by the lanthanide coordination.

¹H NMR spectra of 3 and 4 were recorded in toluene-d₈ and are consistent with C₂ᵥ-symmetric species with three bipym (taphen) proton shifts of 256.3 (73.9), −3.49 (−135.1), and −159.4 (−295.7) ppm, one Cp* methyl shift of 7.91 (13.44) ppm, and one palladium methyl shift of 6.03 (−8.46) ppm. Since the signals are largely shifted in both cases compared to usual diamagnetic positions, the ¹H NMR data are in agreement with an oxidized Yb center and thus a reduced LPdMe₂ fragment (L = bipym, taphen). The variable-temperature NMR of 4 revealed a Curie behavior (linear δ vs 1/T plot over the −80 to 60 °C temperature range, see SI, Figure S7).

Temperature-dependent magnetic data were recorded in the 2–300 K temperature range (Figure 2). The χT values at room temperature (RT) are 2.28 and 2.62 emu-K·mol⁻¹ for 3 and 4, respectively. The related Cp*₂Yb(phen) and Cp*₂Yb(bipy) complexes have a χT value of 2.36 (triplet) and 0.72 emu-mol⁻¹·K (singlet), respectively, at 300 K. In 3 and 4, the net χT value at RT is thus in better agreement with a triplet state that is significantly populated at RT. For 4, when the temperature decreases, the χT value monotonously decreases to 2.10 emu-mol⁻¹·K at 15 K and then drops to 1.4 emu-mol⁻¹·K at 4 K. This behavior is typical for Yb complexes and is attributed to the depopulation of the crystal field states: the overall magnetic data of 4 are in agreement with a triplet ground state (S = 1). The situation found in 3 is drastically different. The χT value decreases faster with decreasing temperature until a first inflection point around 50 K and a second one at 15 K. This behavior was already reported for Cp*₂Yb(L) complexes and may be explained as follows: (i) At low temperature, the ground state is an open-shell singlet ground state (F¹,L∗) mixed with a closed-shell singlet (F¹₄,L⁰); the ground state is multiconfigurational, and the magnetic susceptibility is temperature independent. (ii) The inflection point at 15 K indicates the presence of a second open-shell singlet ground state with a different ratio of open-shell vs closed-shell, while the temperature increases, this low-lying state is getting populated (Boltzmann). The presence of the two low-lying singlet states is in good agreement with two similar b₁ LUMO and LUMO+1 orbitals in 1 (see SI, Figure S29). (iii) The inflection point at 50 K indicates the temperature at which the triplet state (S = 1) is getting populated up to RT. The relative energy of the triplet—singlet gap can be qualitatively estimated from the χT value at RT (300 cm⁻¹, SI, Figure S14).

The magnetic studies and NMR data agree well with different electronic ground states for 3 and 4. In 4, the electron is lying on the ligand with little participation of the Yb metal center, while in 3, a multi-configuration state is present and the Yb metal center has a strong influence on the electron located on the ligand. The next step is to understand how it may affect the reactivity at the Pd center: this strong contrast noted between 3 and 4 may allow tuning the stability of PdIV intermediates. A simple oxidative addition has been performed to test this conjecture.

The oxidative additions of 1 and 2 by MeI were followed by ¹H NMR in toluene-d₈ and thf-d₈ for solubility reasons. In agreement with Canty’s previous experiments, the oxidative addition is rather slow at RT (SI, Table S1). For 1 and 2, the reductive elimination is fast and starts when the PdIV intermediate is not entirely formed (SI, Table S2). When studied on the isolated PdV complexes, the reductive elimination is fast as well in both cases (see SI, Figures S16–S23 and Tables S1 and S2, for details).

The reaction of 1 equiv of MeI with 4 at RT in toluene-d₈ (Scheme 2) leads to a slight change of color from dark red to brown, accompanied by precipitation of a dark powder. The ¹H NMR analysis of the product performed several minutes after the reaction indicates the presence of several species whose ratio changes over time and the characteristic ¹H NMR singlet of
The latter complex crystallizes in toluene-crystallization of NMR at RT reveals only the formation of ethane and further ORTEP of (LUMO in SI, Figure S30).

particular bond, in agreement with the population of a b1 orbital establishing a strongly localized

The 1H NMR spectrum of 8 in toluene-d8 reveals three bipyprotons at 264.5, 4.70, and −168.4 ppm, close to the values for 3, in agreement with a similar structure and electronic structure. Two additional signals are present for the methyl groups in a 2:1 ratio, and the Cp8 has two resonances (5.99 and 6.34 ppm), in agreement with a C3 symmetry and indicative of the presence of a bonded I-atom in solution on the NMR time scale. This contrasts with Canty’s findings that the cationic form of the complex is the key for the reductive elimination to occur. The spectra were very similar in coordinating solvents such as thf-d8 and pyridine-d5. Addition of CD3I to 3 leads to a scrambling of the deuterated methyl positions, even when the reaction is performed at low temperature. This can be explained by a possible alkyl halide exchange in these complexes during the oxidative addition. The oxidative addition has also been performed with MeOTf, and it only resulted in fast reductive elimination (SI, Figure S28), which is indicative of the importance of the Pd–I coordination in the PdIV stability.

The clean and fast synthesis of 8 allows its crystallization in a pure form. We took advantage of this to study the kinetics of the reductive elimination process to understand the influence of the organolanthane fragment. These studies were performed in toluene-d8, a non-coordinating and non-dissociative solvent, and showed first-order kinetics. A satisfactory Eyring plot was obtained between 10 and 30 °C (Figure 4), revealing ΔHf = 24(1) kcal·mol−1 and ΔSf = 6(6) cal·mol−1·K−1. The ΔHf is significantly increased compared to Canty’s measurements of the (bipy)PdMe2 complex in several different solvents (from 8.4 to 18.6 kcal·mol−1). Thus, it seems that the mechanism of the reductive elimination is not modified compared to that reported earlier, but the barrier is increased because of a better stability of the octahedral tris-methylpalladium complex that is due to the presence of the lanthanide fragment. The net result of this is expressed by the half-life time of 8 being more than 1 h at RT, while those of 1, 2, and 4 were too fast to be measured in our hands (minutes).

Accordingly, the electronic structure of 8 was studied by solid-state magnetism and features a behavior similar to that of 3: complex 8 is a multi-configurational singlet ground state with a triplet that is substantially populated at RT. The only notable difference is the absence of an inflection point at 50 K, explained by the presence of a sole singlet ground state below the triplet in energy. This difference accounts for the different geometry (square planar vs octahedral) and indicates how the presence of the organolanthane fragment may influence the electronic structure of the overall molecule, playing a role in the reactivity patterns at the transition metal center.

In summary, we have successfully synthesized molecules containing both lanthanide and reactive PdMe2 fragments with two different bridging ligands in which an electron is transferred...
from the lanthanide fragment, influencing drastically the rapidity at which the oxidative addition with MeI occurs by making the ligands more strongly donating and allowing the clean isolation of PdIV species. The different symmetry of the LUMOs of the (L)PdMe₂ fragments induces large differences in the electronic structure of the overall molecules. Complex 4 has a singlet ground state and 3 a triplet ground state. Moreover, in 3, the electron is transferred on the b₁ orbital that has significant spin density on the N-atoms coordinated to the Pd center. These considerations have a large consequence on the relative stability of the PdMe₁₁ fragment. This work implements an additional strategy for the fine-tuning of the formation and relative stability of high oxidation states in organometallic reactions, by modifying soundly the redox non-innocent ligand and the divergent lanthanide fragment.

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