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One-Electron Oxidation of $[M(P^tBu_3)_2]$ ($M = Pd, Pt$): Isolation of Monomeric $[Pd(P^tBu_3)_2]^+$ and Redox-Promoted C–H Bond Cyclometalation

Thibault Troadec, Sze-yin Tan, Christopher J. Wedge, Jonathan P. Rourke, Patrick R. Unwin, and Adrian B. Chaplin*

Abstract: Oxidation of zero-valent phosphine complexes $[M(P^tBu_3)_2]$ ($M = Pd, Pt$) has been investigated in 1,2-difluorobenzene solution using cyclic voltammetry and subsequently using the ferrocenium cation as a chemical redox agent. In the case of palladium, a mononuclear paramagnetic Pd^I derivative was readily isolated from solution and fully characterized (EPR, X-ray crystallography). While in situ electrochemical measurements are consistent with initial one-electron oxidation, the heavier congener undergoes C–H bond cyclometalation and ultimately affords the 14 valence-electron Pt^II complex $[Pt(\kappa^2_{PC}P^tBu_2CMe_2CH_2)(P^tBu_3)]^+$ with concomitant formation of $[Pt(P^tBu_3)_2H]^+$.

Over the past few decades a rich variety of chemistry has emerged based on the reactions of palladium and platinum complexes in the 0 and +II formal oxidation states, epitomized by the omnipresence of palladium catalyzed cross-coupling reactions in contemporary organic chemistry.^[1,2] In contrast, the organometallic chemistry of well-defined complexes of these elements bearing formal +I oxidation states is much less established and examples are largely limited to unstable or dinuclear species with distinct metal–metal bonds.^[3,4] Halogen bridged palladium complexes of the type $[Pd(\mu-X)(P^tBu_3)_2]$ ($X = Br, I$) are notable examples and are believed to act as reservoirs for reactive $\{Pd^0(P^tBu_3)\}$ fragments in catalytic transformations.^[5] In other systems, Pd^I and Pt^I species have been postulated as intermediates, but with little supporting evidence.^[6] With a view to isolating well-defined mononuclear complexes in the +I oxidation state relevant to catalysis, we report herein our work involving one-

electron oxidation of widely used and commercially available palladium(0) and platinum(0) complexes of tri-*tert*-butylphosphine $[M^0(P^tBu_3)_2]$ ($M = Pd, \mathbf{1a}$; $Pt, \mathbf{1b}$).

As a starting point we determined the redox potentials of $\mathbf{1a}$ and $\mathbf{1b}$ by cyclic voltammetry (CV) in the weakly coordinating solvent 1,2-difluorobenzene (0.2 M $[nBu_4N][PF_6]$ electrolyte, Figure 1).^[7] Reversible one-electron oxidation was observed at $E_{1/2} = -0.44$ V ($\mathbf{1a}$) and $E_{1/2} = -0.10$ V ($\mathbf{1b}$) relative to $Fc/[Fc]^+$ ($Fc =$ ferrocene). The electrochemical characteristics of closely related cyclic alkyl(amino) carbene (CAAC) analogues have recently been studied by CV and the redox potentials of $\mathbf{1a}$ and $\mathbf{1b}$ are similar in magnitude to those found for $[M^0(CAAC)_2]$ ($M = Pd, -0.60$ V; $Pt, -0.07$ V) in THF (0.1 M $[nBu_4N][ClO_4]$).^[8] Consistent with the generation of a stable Pd^I species ($\mathbf{2a}$), the peak current ratios (i_p^{red}/i_p^{ox}) in the palladium voltammograms are essentially unity (ca. 0.99). Conspicuously lower ratios were observed for the platinum complex (ca. 0.90).

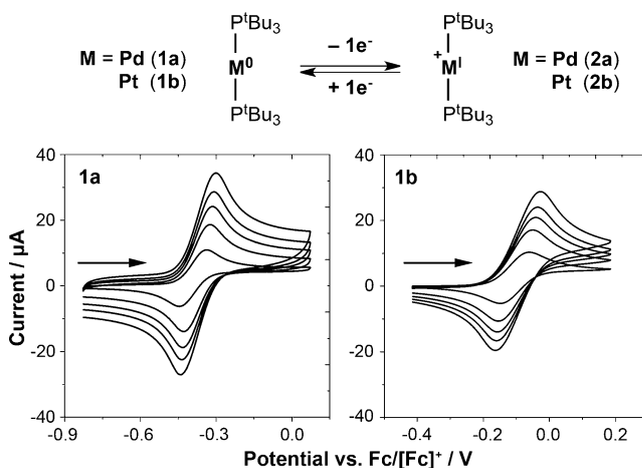


Figure 1. Cyclic voltammograms for the oxidation of $\mathbf{1a}$ and $\mathbf{1b}$ in 1,2- $C_6H_4F_2$ (2 mM $\mathbf{1}$; 0.2 M $[nBu_4N][PF_6]$ electrolyte; glassy carbon working electrode, Pt counter electrode and Ag wire reference electrode; scan rates = 10, 30, 50, 70, and 100 $mV s^{-1}$).

Encouraged by these data, $\mathbf{1a}$ was reacted with one equiv of $[Fc][PF_6]$ in 1,2-difluorobenzene at 293 K and dark blue $[Pd^I(P^tBu_3)_2][PF_6]$ $\mathbf{2a}$ was subsequently isolated in 92% yield following addition of *n*-pentane. The electrochemical characteristics of isolated $\mathbf{2a}$ are equivalent to those measured in situ starting from $\mathbf{1a}$ ($E_{1/2} = -0.42$ V; see Supporting

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Information). This new paramagnetic species was additionally characterized in solution using UV/Vis spectroscopy ($\lambda_{\text{max}} = 667 \text{ nm}$), ESI-HRMS (positive ion mode, $510.2736 \text{ m/z } [M]^+$; calculated 510.2740 m/z), and EPR spectroscopy. The EPR spectrum (1,2- $\text{C}_6\text{H}_4\text{F}_2$ glass at 200 K, Figure 2), shows a superposition of a single resonance at $g = 2.316(5)$ with a lower intensity sextet arising from hyperfine coupling to ^{105}Pd

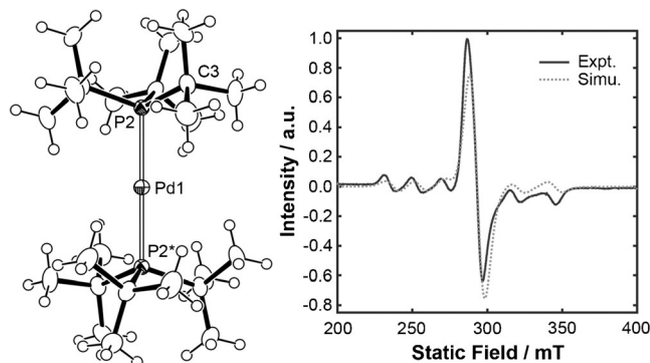
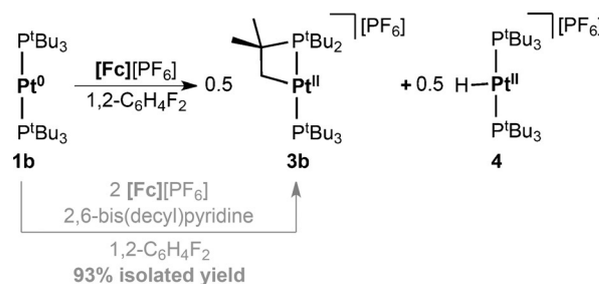


Figure 2. The solid-state structure^[22] and EPR spectrum of **2a** (1,2- $\text{C}_6\text{H}_4\text{F}_2$ glass, 200 K, a.u. = arbitrary units).^[11] Ellipsoids are set at 50% probability; anion omitted for clarity. The starred atom is generated by the symmetry operation $1-x, 1-y, 1-z$. Selected data: Pd1–P2 2.3470(6) Å; P2–Pd1–P2* 180°, Pd1–P2–C3 108.81(5)°.

($I = 5/2$, 22% abundance), corroborating formation of an $S = 1/2$ Pd^{I} species. The unusually large ^{105}Pd hyperfine coupling of approximately 25 mT, and lack of resolved coupling to ^{31}P ($I = 1/2$, 100% abundance) is consistent with strong localization of the unpaired electron spin on the Pd center. Complex **2a** crystallizes in the high-symmetry cubic space group $Pa\bar{3}$ with the palladium atom on a center of inversion (Figure 2). In comparison to **1a**, the Pd–P bond length is significantly elongated, from 2.285(3) to 2.3469(6) Å ($\Delta(\text{Pd–P}) = +0.062(4)$ Å); the P–Pd–P angles in both cases are symmetry enforced at 180°.^[9] To the best of our knowledge, this is the first example of an unsupported two-coordinate Pd^{I} complex. A similar bond length elongation has been noted in closely related NHC complexes of $\text{Ni}^{\text{0}}/\text{Ni}^{\text{I}}$ ($\Delta(\text{Ni–C}) = +0.08(2)$ Å).^[10] Isolated **2a** is air-sensitive in solution, but shows good stability under an argon atmosphere. For instance, under argon the EPR spectrum intensity was essentially unchanged after 24 h at 293 K (15 mm). However, slow degradation of **2a** was observed by UV/Vis spectroscopy under high dilution conditions ($t_{1/2} \approx 30 \text{ h}$; 0.15 mM), which we attribute to the presence of adventitious water as the rate of degradation increased significantly when water was added deliberately. Moreover, **2a** can be stored in the solid-state in air (72 h) with no evident change by UV/Vis spectroscopy.

When preparation of the analogous Pt^{I} complex **2b** was attempted by reaction of **1b** with one equiv of $[\text{Fc}][\text{PF}_6]$, a 1:1 mixture of the new diamagnetic cyclometalated complex $[\text{Pt}^{\text{II}}(\kappa^2_{\text{PC}}\text{-P}^{\text{t}}\text{Bu}_2\text{CMe}_2\text{CH}_2)(\text{P}^{\text{t}}\text{Bu}_3)][\text{PF}_6]$ **3b** and known Pt^{II} hydride $[\text{Pt}^{\text{II}}(\text{P}^{\text{t}}\text{Bu}_3)_2\text{H}][\text{PF}_6]$ **4** ($\delta(^1\text{H}) -36.30 \text{ ppm}$; $^2J_{\text{PtH}} = 8.6$, $^1J_{\text{PtH}} = 2590 \text{ Hz}$; $\delta(^{31}\text{P}) 86.3 \text{ ppm}$; $^1J_{\text{PtP}} = 2621 \text{ Hz}$) was formed within 15 min instead, as indicated by ^1H and



Scheme 1. Chemical oxidation of **1b**.

^{31}P NMR spectroscopy (Fc observed; Scheme 1).^[12] This outcome suggests only transient stability of **2b** in solution, with subsequent C–H bond homolysis accounting for the divergence from fully reversible one-electron oxidation of **1b** observed by CV.^[13] Reaction of **1b** with two equiv of $[\text{Fc}][\text{PF}_6]$ in the presence of excess hindered base 2,6-bis(decyl)pyridine (5 equiv), which is able to deprotonate **4**, resulted in selective formation of **3b** within 15 min. In this manner, **3b** was isolated in 93% yield following successive crystallizations from 1,2- $\text{C}_6\text{H}_4\text{F}_2$ to remove ferrocene, excess base, and pyridinium salt.^[13] For comparison, no significant reaction was detected by ^1H or ^{31}P NMR spectroscopy on mixing of **1b** and 2,6-bis(decyl)pyridine in 1,2-difluorobenzene at 293 K (24 h) or heating **1b** alone in 1,2-difluorobenzene at 353 K (24 h).

Two independent but structurally similar cations are observed in the solid-state structure of **3b** (one is shown in Figure 3), both illustrating adoption of a T-shaped coordination geometry^[14] and cyclometalation of one of the *tert*-butyl substituents; these are identified by distinctly acute Pt1–P2–C3 angles [90.0(3)/89.5(3)°] and Pt1–C4 bond lengths of 2.063(17)/2.065(17) Å. The **3b** cation is formally a 14 valence-electron (VE) complex, but is stabilized by adoption

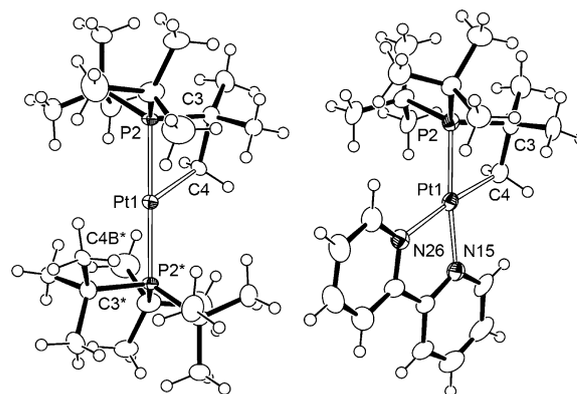


Figure 3. Solid-state structures of **3b** and **6**.^[22] Ellipsoids are set at 50% and 30% probability, respectively; minor disordered components and anions omitted for clarity; only one of the two independent molecules is shown for **3b**. Starred atoms in **3b** are generated by the symmetry operation $1-x, 2-y, 1-z$. Selected data **3b**: Pt1–P2 2.297(2) Å, Pt1–C4 2.063(17) Å, Pt1...C4B* 2.83(2) Å; P2–Pt1–P2* 180°, Pt1–P2–C3/C3* 90.0(3)°. **6**: Pt1–P2 2.235(2) Å, Pt1–C4 2.077(10) Å, Pt1–N15 2.080(7) Å, Pt1–N26 2.156(7) Å; P2–Pt1–N15 166.3(2)°, C4–Pt1–N26 175.4(3)°, Pt1–P2–C3 88.5(3)°.

of an agostic interaction between the non-cyclometalated phosphine ligand and Pt center (Pt1...C4B 2.83(2)/2.84(2) Å). In solution, the structure of **3b** was fully corroborated by NMR spectroscopy (CD₂Cl₂, 298 K). Formation of the metallacycle is apparent by distinctive ¹H and ¹³C methylene resonances at δ(¹H) 2.75 ppm (²J_{PtH} = 110 Hz) and δ(¹³C) 10.3 ppm (¹J_{PtC} = 670 Hz) with platinum satellites, two doublet ³¹P resonances with a large (*trans*) ²J_{PP} coupling constant and platinum satellites (δ(³¹P) 59.1 ppm (¹J_{PtP} = 2896 Hz, ²J_{PP} = 317 Hz, P¹Bu₃), δ(³¹P) 25.2 ppm (¹J_{PtP} = 1916 Hz, ²J_{PP} = 317 Hz, P¹Bu₂CMe₂CH₂)), and a platinum chemical shift of δ(¹⁹⁵Pt) –3816 ppm (225 K). Although the signals associated with the non-cyclometalated phosphine ligand broadened on cooling to 185 K, the agostic interaction could not be definitively resolved by ¹H NMR spectroscopy.

Cyclometalation reactions of Pt^{II} complexes have extensive precedent.^[14a,15] For instance, T-shaped complexes [Pt^{II}(κ²_{PC}-PR₂C₆H₃MeCH₂)(PR₂Xyl)]⁺ (R = Cy, Ph; Xyl = 2,6-dimethylphenyl) with similar structural and spectroscopic metrics compared to **3b**, were prepared by cyclometalation reactions involving halide abstraction from [Pt^{II}(PR₂Xyl)₂(Me)Cl] and subsequent elimination of methane.^[16] Intramolecular C–H bond activation of P¹Bu₃ in [Pt^{II}(P¹Bu₃)₂HX] (X = Cl, Br, I, OTf, NO₂) has also been described and results in coordinatively saturated products [Pt^{II}(κ²_{PC}-P¹Bu₂CMe₂CH₂)(P¹Bu₃)X].^[17] In the case of **3b**, the presence of a Pt^{II} intermediate preceding cyclometalation can be discounted on the basis of the electrochemical characteristics of **1b**. Instead the formation of **3b** and **4** presumably occurs via concerted bimetallic (radical) oxidative addition,^[18] or proceeds through a common Pt^{III} alkyl hydride intermediate [Pt^{III}(κ²_{PC}-P¹Bu₂CMe₂CH₂)(P¹Bu₃)H]⁺ (**5**). In the latter case, subsequent comproportionation (**5** + **2b**), disproportionation (via a Pt^{IV} alkyl dihydride), or Pt–H bond homolysis (i.e. 2 × **5** → 2 × **3b** + H₂; **3b** + H₂ → **4**) would afford the observed 1:1 mixture of **3b** and **4**.^[19]

Seeking to gain more insight into this mechanism, trapping of the postulated intermediate **5** was attempted by coordination of 2,2'-bipyridine (bipy). However, oxidation of **1b** with either one or two equiv of [Fc][PF₆] in the presence of one equiv of bipy resulted in formation of a new cyclometalated complex [Pt^{II}(κ²_{PC}-P¹Bu₂CMe₂CH₂)(bipy)][PF₆] **6** instead, alongside protonated phosphine (δ(³¹P) 54.2 ppm). The identity of this new complex was verified by independent synthesis from **3b** and bipy in 1,2-C₆H₄F₂ (97% yield of isolated product). As with **3b**, the cyclometalated phosphine in **6** is characterized by an acute Pt1–P2–C3 angle (88.5(3)°) and bears a similar Pt1–C4 bond length of 2.077(10) Å (Figure 3). Moreover, both solution and solid-state data are fully consistent with a coordinatively saturated metal complex. Notably, the substantially higher *trans*-influence of the methylene ligand is reflected in different Pt–N bond lengths (Pt1–N15, 2.156(7) versus Pt1–N26, 2.080(7) Å); the associated ¹³C resonance shows a reduced ¹J_{PtC} coupling in comparison to **3b** (580 versus 670 Hz). Stronger Pt–P bonding is apparent in **6** relative to **3b**, on the basis of a shorter Pt–P bond (2.235(2) versus 2.297(2)/2.299(3) Å), and a larger ¹J_{PtP} coupling constant determined by ³¹P NMR spectroscopy (3105 versus 1916 Hz). A platinum chemical shift of δ(¹⁹⁵Pt)

–3788 ppm (225 K) was also measured for **6** and is very similar to that of **3b** (δ(¹⁹⁵Pt) –3816 ppm).

Reaction of isolated **3b** with H₂ (1 atm) results directly in the formation of **4**, which is reconcilable with Pt–H bond homolysis or disproportionation (via an unstable Pt^{IV} alkyl dihydride intermediate) during the formation of **3b/4**. However, the underlying mechanism is still not completely clear at this time. For instance, we cannot discount the formation of **3b** through a pathway involving deprotonation of **5** (mediated by **1b**^[20] or 2,6-bis(decyl)pyridine) and a second one-electron oxidation. The redox potential of the associated Pt^I/Pt^{II} couple, assessed by CV experiments using both isolated **3b** (*E*_{1/2} = –1.90 V, irreversible) and **6** (*E*_{1/2} = –1.68 V, *i*_p^{ox}/*i*_p^{red} ≈ 0.96), indicates that such a one-electron oxidation is at least conceptually feasible using [Fc][PF₆] (see Supporting Information for CVs).

Motivated by the cyclometalation observed on oxidation of **1b**, we have also preliminarily investigated whether similar reactivity can be induced in the palladium analogue. Our studies are on-going, but we do note that reaction of **1a** with two equiv of [Fc][PF₆] in the presence of excess 2,6-bis(decyl)pyridine (5 equiv) resulted in the gradual appearance of a diamagnetic complex with spectroscopic characteristics consistent with cyclometalation (**3a**; δ(³¹P) 57.0, –1.3 ppm; ²J_{PP} = 316 Hz).^[21] However, this species was only formed in situ in about 30% yield after 72 h at 293 K, as measured by NMR spectroscopy (using an internal standard), and the resulting reaction mixture has proved intractable so far to further characterization.

In summary, we have described a simple method for accessing the reaction chemistry of mononuclear palladium and platinum complexes bearing a +I formal oxidation state, as demonstrated by one-electron oxidation of [M⁰(P¹Bu₃)₂] (M = Pd, Pt) using [Fc][PF₆]. While the Pd^I derivative was readily isolated from solution and fully characterized, the heavier congener undergoes C–H bond cyclometalation to afford the 14 VE Pt^{II} complex [Pt^{II}(κ²_{PC}-P¹Bu₂CMe₂CH₂)(P¹Bu₃)]⁺ with concomitant formation of [Pt^{II}(P¹Bu₃)₂H]⁺. Future work is focused on charting the reactivity and catalytic activity of these novel Group 10 species, and will be published in due course.

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