

## Chlorine Photoelimination from a Diplatinum Core: Circumventing the Back Reaction

Timothy R. Cook, Yogesh Surendranath, and Daniel G. Nocera\*

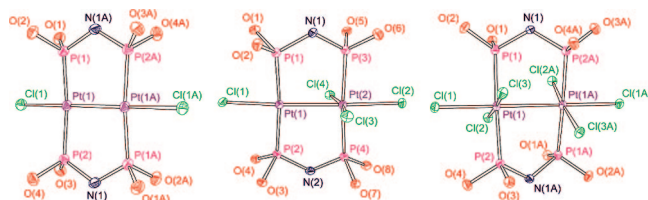
Department of Chemistry, 6-355, Massachusetts Institute of Technology, 77 Massachusetts Avenue, Cambridge, Massachusetts 02139-4307

Received September 11, 2008; E-mail: nocera@mit.edu

Solar energy may be stored by using it to rearrange the bonds of  $\text{H}_2\text{O}$  and  $\text{HX}$  ( $\text{X} = \text{Cl}, \text{Br}$ ) to  $\text{H}_2/\text{O}_2$  and  $\text{H}_2/\text{X}_2$ , respectively.<sup>1,2</sup> From a mechanistic standpoint, the oxidative half-reaction of  $\text{HX}$  is more attractive than  $\text{H}_2\text{O}$  because it only involves the coupling of a two-electron oxidation to  $\text{H}_2$  generation. The water oxidation half-reaction requires four electrons, and necessarily involves the intimate coupling of electron to proton transfer.<sup>3,4</sup> Unlike the  $1\text{e}^-$  chemistry of redox symmetric bimetallic complexes,<sup>2,5</sup> we have developed a method for hydrogen production from  $\text{HX}$  solutions utilizing a bimetallic photocatalyst that operates in two discrete two-electron steps for hydrogen evolution and halogen elimination.<sup>6,7</sup> A hydrido-halide  $\text{Rh}_2^{\text{II,II}}(\text{H})(\text{X})_2$  species is generated from the addition of two  $\text{HX}$  molecules to a parent  $\text{Rh}_2^{0,0}$  complex.  $\text{H}_2$  photoelimination is facile to furnish a two-electron mixed valence complex,  $\text{Rh}_2^{0,\text{II}}(\text{X})_2$ ; photoelimination of halogen, however, from  $\text{Rh}_2^{0,\text{II}}(\text{X})_2$  is not. Hence the overall efficiency for the  $\text{H}_2$  photocycle is limited by the efficiency of halogen photoelimination. More generally, overcoming the strong metal–halide bond is an important overall determinant of  $\text{H}_2$  production in most  $\text{HX}$  photocycles. If no pathway for bond activation is accessible, productive chemistry ceases upon hydrogen evolution.<sup>8</sup> Even if halogen photoelimination is achieved, halogen must be trapped since the back-reaction of the primary photoelimination products is rapid and favorable.<sup>9</sup> For this reason, the overall photoefficiency typically scales with the efficiency of the chemical trap. Traps are problematic in that the trap– $\text{X}$  bond provides the thermodynamic driving force,<sup>10,11</sup> thus obviating energy storage implications. In addition, the detailed process by which photoelimination proceeds is obscured as both  $\text{X}^\bullet$  and  $\text{X}_2$  can react with most traps. Improved efficiencies for  $\text{H}_2$  production therefore require increased quantum yields for  $\text{M}-\text{X}$  bond activation and new strategies to prevent the back reaction of primary photoproducts.

Our most recent investigations of  $\text{HX}$  photocycles have focused on improving the efficiency of the halogen elimination step of the photocycle by incorporating oxidizing metals, such as Au and Pt into bimetallic cores.<sup>12,13</sup> Whereas a  $\text{PtAu}$  bimetallic trihalogen complex shows markedly improved photoefficiencies for photoelimination,<sup>13</sup> it nonetheless requires a halogen trap. We now report a system that addresses both challenges of a high photoefficiency for  $\text{X}_2$  elimination in the absence of a trap. Photolysis of a  $\text{Pt}_2^{\text{III,III}}$  complex with UV or visible light (355–510 nm) eliminates halogen to deliver a two-electron mixed-valent  $\text{Pt}_2^{\text{I,III}}$  with a high quantum yield in solution. Photolysis of the complex in the solid state yields  $\text{Cl}_2$  with no need for a halogen trap.

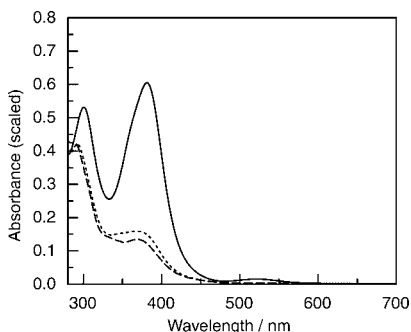
Though bimetallic complexes of group 10 metals bridged by diphosphine ligands typically promote insertion reactions,<sup>14–18</sup> they can drive oxidative chemistry.<sup>19</sup> With regard to the latter, we were intrigued by the oxidative chemistry of  $\text{Pd}_2(\text{P}-\text{P})_2\text{X}_2$  ( $\text{P}-\text{P}$  = dimethylphosphinomethane, diethylphosphinomethane) by halogens to yield valence symmetric face-to-face  $\text{Pd}_2^{\text{II,II}}(\text{P}-\text{P})_2\text{X}_4$  ( $\text{X} = \text{Cl},$



**Figure 1.** Thermal ellipsoid plots of **1** (left) **2** (center) and **3** (right) drawn at the 50% probability level; **1** and **3** crystallize with half a molecule per asymmetric unit. (O)– $\text{CH}_2\text{CF}_3$  and (N)– $\text{CH}_3$  groups of the phosphazanes are omitted for clarity.

Br) and  $\mu\text{-X}_2$  complexes ( $\text{X} = \text{I}$ ).<sup>20</sup> We find that the  $\text{d}^8 \cdots \text{d}^8$  cores of these complexes can disproportionate to deliver  $\text{d}^7\text{--d}^9$  mixed-valence complexes in the presence of phosphazane ligands. Suitable  $\text{Pt}^0$  and  $\text{Pt}^{\text{II}}$  sources (e.g.,  $(^t\text{Bu}\text{dba})_3\text{Pt}_2$  and  $\text{Pt}(\text{cod})\text{Cl}_2$ ,  $^t\text{Bu}\text{dba} = 4\text{'-tert-butyl dibenzylideneacetone}$ ,  $\text{cod} = 1,5\text{-cyclooctadiene}$ ) react with the phosphazane tefpma ( $\text{tefpma} = ((\text{CF}_3\text{CH}_2\text{O})_2\text{P})_2\text{NCH}_3$ ) to yield  $\text{Pt}_2^{\text{I,II}}(\text{tefpma})_2\text{Cl}_2$  (**1**) in analogy to the synthesis of  $\text{Pt}_2^{\text{I,II}}$  phosphine complexes.<sup>21</sup> The solid state structure of **1**, shown in Figure 1, reveals a short Pt–Pt contact of 2.6187(6) Å, similar to that of other  $\text{M}_2(\text{P}-\text{P})_2\text{X}_2$  bimetallics and consistent with a metal–metal bond.<sup>17,22</sup> A  $^3\text{1P}\{\text{1H}\}$  resonance at 113.57 ppm disappears upon the addition of 1 equiv of  $\text{PhI}\cdot\text{Cl}_2$  to a solution of **1** and two resonances appear at 100.52 and 60.62 ppm, corresponding to phosphorus atoms coordinated to the  $\text{Pt}^{\text{I}}$  and  $\text{Pt}^{\text{III}}$ , respectively, of  $\text{Pt}_2^{\text{I,III}}(\text{tefpma})_2\text{Cl}_4$  (**2**). X-ray diffraction studies confirm the two-electron mixed-valent nature of **2** (Figure 1). The  $\text{Pt}(1)\text{--Pt}(2)$  distance of 2.6187(7) Å establishes that a metal–metal bond is maintained upon oxidation. The coordination of  $\text{Cl}(3)$  and  $\text{Cl}(4)$  to  $\text{Pt}(2)$ , ( $d_{\text{Pt}(2)\text{--Cl}(3)} = 2.323(2)$  Å and  $d_{\text{Pt}(2)\text{--Cl}(4)} = 2.338(2)$  Å) completes the pseudo-octahedral geometry about the  $\text{Pt}^{\text{III}}$  center. The structure of **2** is unique among the oxidation products of  $\text{M}_2(\text{P}-\text{P})_2\text{X}_2$  bimetallics, as it represents the first example of a group 10 two-electron mixed-valent compound. Oxidation of **2** with 1 equiv of  $\text{PhI}\cdot\text{Cl}_2$ , or treatment of **1** with 2 equiv of  $\text{PhI}\cdot\text{Cl}_2$ , affords a red solution of  $\text{Pt}_2^{\text{III,III}}(\text{tefpma})_2\text{Cl}_6$  (**3**). The  $^3\text{1P}\{\text{1H}\}$  NMR spectrum of **3** shows a single resonance at 52.22 ppm. As with **1** and **2**, X-ray diffraction analysis of the complex indicates the presence of a direct Pt–Pt interaction ( $d_{\text{Pt--Pt}} = 2.7037(3)$  Å) for two metals possessing an octahedral coordination environment.

The diplatinum-based suite of  $\text{d}^7\text{--d}^7$ ,  $\text{d}^9\text{--d}^7$ , and  $\text{d}^9\text{--d}^9$  complexes defined by **3**, **2**, and **1**, respectively, complement the previously described dirhodium suite of complexes of the same d-electron count.<sup>9</sup> As shown in Supporting Information, Figure S7, the electronic profiles of **1–3** are dominated by excited states of  $\text{d}\sigma^*$  parentage as observed for the spectroscopy of  $\text{d}^7\text{--d}^7$ ,  $\text{d}^9\text{--d}^7$ , and  $\text{d}^9\text{--d}^9$  dirhodium complexes.<sup>23</sup> As the  $\text{d}\sigma^*$  absorption profile possesses both  $\text{M--M}$  and  $\text{M--X}$  antibonding character, electronic excitation into the empty  $\text{d}\sigma^*$  orbital is expected to promote halogen



**Figure 2.** Electronic absorption spectra of **3** (solid), **2** (dashed), and the solid state photolysis product (dotted) in  $\text{CH}_2\text{Cl}_2$ ,  $\lambda_{\text{irr}} > 350$  nm.

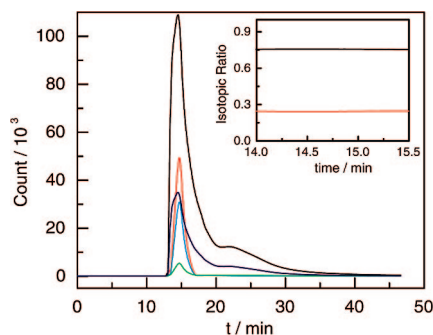
elimination as we have observed for the M–X bond photoactivation of  $\text{Rh}_2^9$  and  $\text{PtAu}^{13}$  systems.

Irradiation of 43  $\mu\text{M}$  solutions of **3** in benzene with either 405 or 510 nm light in the presence of 2,3-dimethyl-1,3-butadiene (DMBD) leads to rapid conversion to **2** as monitored by both UV–vis spectroscopy (Figure S8) and  $^{31}\text{P}\{^1\text{H}\}$  NMR (Figure S13). The quantum yield for Pt–Cl bond activation is high. Figure S9 shows the measured photoreaction quantum yields ( $\Phi_p$ ) as related to the DMBD concentration. The observed quantum yield of 38% for halogen photoelimination far exceeds all previously reported values.<sup>6,13</sup> Since both 405 and 510 nm light give the same efficiency for halogen elimination, a common excited-state must be reached by the 385 and 520 nm transition bands. Photolysis of **3** in benzene in the absence of DMBD results in reductive fragmentation of the bimetallic core to yield  $\text{Pt}(\text{tfepma})\text{Cl}_2$ . Conversely, photolysis of **3** in THF in the absence of DMBD results in its quantitative conversion to **2**. Based on dirhodium photochemistry,<sup>6,7</sup> the photoeliminated halogen is efficiently trapped by  $\alpha$ -H abstraction from THF.

The resulting suite of compounds preserves a metal–metal bond across a four-electron transformation with a  $\text{Pt}_2^{\text{III}}$  species mixed-valence species as the linchpin. The presence of metal–metal bonding across the series provides an accessible excited-state with  $\sigma^*$  character from which photochemistry can originate, in much the same way as has been previously reported for  $\text{Rh}_2$  dimers.<sup>9,23</sup>

A halogen trap is circumvented if the photoreaction is performed in the solid state. Irradiation of solid powder samples of **3** in vacuo and in ambient conditions results in elimination of  $\text{Cl}_2$  to produce (based on UV–vis,  $^{31}\text{P}\{^1\text{H}\}$  and  $^1\text{H}$  analysis, Figures 2, S12, and S14) **2** in 70% synthetic yield along with its isomer, the *cis,trans*- $\text{Pt}_2^{\text{II,II}}(\text{tfepma})_2\text{Cl}_4$  compound (see Supporting Information for structural and NMR data) in 15% yield. Generation of chlorine gas in these solid state experiments was confirmed by a mass spectrometric (MS) analysis of the gas in the headspace of a sealed, evacuated high-pressure glass vessel in which **3** was irradiated. Mass fragments corresponding to  $^{35,35}\text{Cl}_2$ ,  $^{37,37}\text{Cl}_2$ ,  $^{35,37}\text{Cl}_2$ ,  $^{35}\text{Cl}^\bullet$ , and  $^{37}\text{Cl}^\bullet$  are all observed (Figure 3); the  $\text{Cl}^\bullet$  results from  $\text{Cl}_2$  fragmentation upon electron impact ionization. The experimental abundance of  $^{35}\text{Cl}$  and  $^{37}\text{Cl}$  of 0.756(1) and 0.244(1), respectively, is in good agreement with the natural abundance of these isotopes in chlorine gas (75.78(4) and 24.22(4), respectively<sup>24</sup>). Since the possibility of a bimolecular reaction is removed in the solid state and no trap is present to collect ejected  $\text{Cl}^\bullet$  radicals, an intramolecular elimination of  $\text{Cl}_2$  from the excited-state of **3** is invoked.

The development of HX splitting photocatalysis is expanded by the results reported herein: (1) high quantum yields for halogen photoelimination have been achieved and (2) the evolution of  $\text{Cl}_2$  upon solid-state photolysis of **3** represents the first example of the thermodynamically unfavorable halogen elimination without the use



**Figure 3.** Mass spectrometry analysis of the gas evolved from solid state photolysis of **3**. Traces correspond to  $^{35}\text{Cl}$  (black),  $^{35}\text{Cl}^{35}\text{Cl}$  (red),  $^{37}\text{Cl}$  (purple),  $^{35}\text{Cl}^{37}\text{Cl}$  (blue),  $^{37}\text{Cl}^{37}\text{Cl}$  (green) mass fragments. Inset shows the fractional amount of  $^{35}\text{Cl}$  (red) and  $^{37}\text{Cl}$  (black) present.

of a chemical trap. Since  $\text{Cl}_2$  addition proceeds facily under the conditions of photolysis, we have shown authentic energy storage reactivity in the absence of a halogen trap. Further work is underway to unify the  $\text{H}_2$  and  $\text{X}_2$  photochemistry and to elucidate mechanistic details of M–X bond activation for the compounds reported here.

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**Note Added after ASAP Publication.** Typographical errors in the compound descriptions presented in the online abstract (ASAP 12/18/08) have been corrected. The revised abstract was reposted on January 7, 2009.

**Supporting Information Available:** Tables of bond lengths and angles, full experimental details and crystallographic information files (CIF). This material is available free of charge via the Internet at <http://pubs.acs.org>.

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