

Heterobimetallic Cooperation Mediates the Transformation of White Phosphorus into Zwitterionic *catena*-Phosphonium(+)diphosphenide(−) Ligands**

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Dedicated to Professor Otto J. Scherer on the occasion of his 75th birthday

White phosphorus, P₄, the most reactive allotrope of the element, is readily available from rock phosphates and is widely used for the chemical manufacturing of a great variety of organophosphorus derivatives.^[1] In a typical industrial procedure, P₄ is treated with Cl₂ at high temperature to yield phosphorus chlorides, which are subsequently treated with organic substrates to afford the organophosphorus derivatives. The growing worldwide market for phosphorus derivatives and increasingly rigorous environmental regulations, which ban dangerous precursors such as chlorine from large-scale industrial processes, are stimulating the quest for methods that avoid chlorine and functionalize white phosphorus under milder conditions.^[2] The most efficient approach in this regard would be an as-yet unknown metal-catalyzed functionalization of P₄,^[3] although several contributions from our group^[4,5] and others^[6] have shown the importance of stoichiometric activations mediated by bimetallic transition-metal complexes. Recently, Bertrand et al. have shown that metal-free activation of P₄ can also be achieved by using two equivalents of cyclic alkyl aminocarbenes to synergically activate white phosphorus.^[7]

Herein we report our findings that white phosphorus can be activated in a stepwise fashion by two different transition-metal units, which initially activate the P₄ tetrahedron and then break the activated polyphosphorus unit into two

zwitterionic diphenyl(alkyl)phosphonium(+)diphosphenide(−) molecules.

The reaction of white phosphorus with Co(BF₄)₂·6H₂O and bis(diphenylphosphino)methane (dppm) in refluxing THF/butanol gives [Co(Ph₂PCH₂PPh₂PPPPPh₂PCH₂PPh₂)]·BF₄ (**1**·BF₄), whose synthesis^[8] represents a turning point in the coordination chemistry of white phosphorus.^[9] Complex **1**⁺ contains a surprising P₆ ligand which originates from the nucleophilic attack of two PPh₂ ends from two different dppm ligands on the η⁴-tetraphosphabutadiene moiety formed by opening of the P₄ molecule. A perusal of the bonding distances—the external P–P distances are shorter than the internal ones (2.172 vs. 2.197 Å)—has allowed Midollini and co-workers to formally attribute the oxidation state −I to Co and to describe the charge distribution of the new hexaphosphorus ligand as a P⁽⁺⁾–P=P–P=P–P⁽⁺⁾ dication where two phosphonium moieties are formally located at each P₆ end.^[8] Attempts to react this unique fragment with several electrophiles and nucleophiles were generally unsuccessful, thus indicating the high stability of the η⁴-tetraphosphabutadiene ligand.^[10]

We have found that the reaction of **1**·BF₄ with [Pt(C₂H₄)(PPh₃)₂] in dichloromethane at room temperature affords the heterobimetallic complex [Co(μ,η^{1:2:1}-P=P-PPh₂CH₂PPh₂)₂{Pt(PPh₃)₂}]BF₄ (**2**·BF₄) in almost quantitative yield by elimination of ethene and insertion of the carbene-like PtL₂ moiety into the central P–P bond of the P₆ ligand of **1**⁺ (Scheme 1).^[11] Work-up as detailed in the Experimental Section gave **2**·BF₄ as a dark brown microcrystalline material in good yield. Complex **2**⁺ was characterized by elemental analysis, ³¹P{¹H} and ¹H NMR spectroscopy, ESI mass spectrometry, and single-crystal X-ray diffraction (Figure 1).^[12] The crystal structure of **2**·BF₄ contains tetrafluoroborate anions, [Co(μ,η^{1:2:1}-P=P-PPh₂CH₂PPh₂)₂{Pt-

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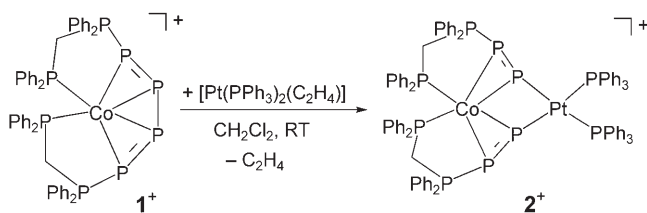
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[**] We are grateful to Thermphos International BV, Vlissingen (Netherlands), for a generous loan of white phosphorus. Our thanks also go to the NMR service of “Firenze Hydrolab”, a project sponsored by Ente Cassa di Risparmio di Firenze. Dr. Andrea Raffaelli (ICCOM CNR, Pisa, Italy) is acknowledged for running the ESI-MS experiments.



Supporting information for this article is available on the WWW under <http://www.angewandte.org> or from the author.



Scheme 1. Synthesis of complex **2**⁺.

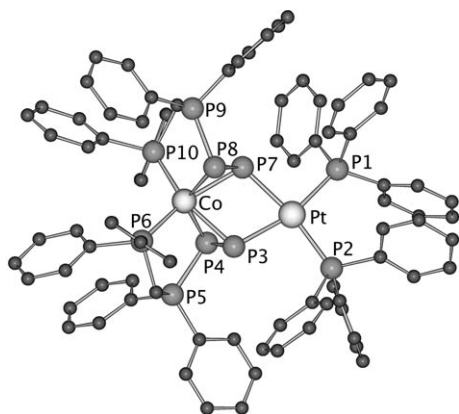


Figure 1. Molecular structure of the heterobimetallic cation 2^+ . Hydrogen atoms have been omitted for clarity. Selected bond lengths [Å] and angles [°]: Co–P10 2.226(3), Co–P6 2.216(3), P3–P4 2.179(4), P7–P8 2.185(4), Pt–P7 2.334(3), Pt–P1 2.317(3); P6–Co–P10 110.77(12), P3–Co–P4 56.67(11), P3–Co–P7 74.43(11), P7–Co–P8 56.57(11), P3–P4–P5 89.19(14), P7–P8–P9 90.06(15).

$(\text{PPh}_3)_2]^+$ cations, and interspersed molecules of ethanol and dichloromethane in 1:1:2:1 ratio.

The original P_6 “zigzag” chain in 1^+ has been cleaved upon reaction with the $\text{Pt}(\text{PPh}_3)_2$ moiety to form two separate P_3 moieties, namely RP5-P4-P3 and RP9-P8-P7 , which are nearly identical in light of their bond lengths and angles. This reaction occurs with complete regioselectivity and may be viewed as the formal oxidative cleavage of the internal $\text{P}^+-\text{P}=\text{P}$ single bond of the η^4 -tetraphosphabutadiene unit $\text{P}^{(+)}-\text{P}=\text{P}-\text{P}^{(+)}$. The P3-P7 bond length changes from 2.197(3) Å in 1^+ to 2.823(4) Å in 2^+ following the insertion of PtL_2 , while the P3-Co-P7 angle opens from $57.2(1)^\circ$ to $74.43(11)^\circ$. The distance between P3 and P7 excludes the existence of any residual weak interaction between these two atoms in 2^+ . The other bond distances (P-P and Co-P) and the P-Co-P bond angles remain virtually unchanged with respect to those in 1^+ . Thus, with the noticeable exception of atoms P3 and P7 , the coordination environment around cobalt in complexes 1^+ and 2^+ is almost superimposable (see the Supporting Information). The formal oxidative charge at the cobalt center is also maintained in 2^+ as for the parent compound 1^+ , that is, Co^{-1} . The geometry around the cobalt center is a distorted tetrahedron and the core of the bimetallic complex contains a C_2 axis of symmetry, as previously found for complex 1^+ . The coordination geometry around platinum is square planar. The torsion angles P5-P4-P3-Pt and P9-P8-P7-Pt are $154.03(12)^\circ$ and $158.23(13)^\circ$, respectively.

The most outstanding feature of 2^+ concerns the electronic nature of the two P_3 ligands which hold the two metal atoms together. Based on the electron charge distribution, the nature of this ligand can be described as a zwitterionic diphenyl(alkyl)phosphonium(+)diphosphenide(−) molecule, $\text{RPh}_2\text{P}^{(+)}-\text{P}=\text{P}^{(-)}$ ($\text{R} = \text{Ph}_2\text{PCH}_2$), where the negative charge is placed on the $\mu\text{-Co,Pt}$ -bridging P atom and the positive one on the opposite PPh_2R phosphonium P atom. The average $\text{P}=\text{P}$ bond length in the diphosphenide moiety is 2.182(4) which is longer than the value found for the only known phosphatetriylphosphonium cation (2.025(1) Å)^[13] and in agreement

with the $\text{P}=\text{P}$ bond length in diphosphenes coordinated to transition metals.^[14] In the case at hand, the diphosphenide section of the zwitterion acts as a two-electron π -donor towards Co and, additionally, as a two-electron σ -donor towards Pt^{II} . There is also some back-donation from the cobalt center into the π^* -orbital of the $\text{P}=\text{P}$ group.^[15] A similar bonding situation has been documented in diphosphene systems by Power and co-workers, who have reported a bimetallic iron complex containing a diphosphene ligand σ,π -bonded to two $\text{Fe}(\text{CO})_4$ units with a $\text{P}=\text{P}$ bond length of 2.184(2) Å, as is the case herein.^[16]

In keeping with the solid-state structure and the analysis of the electronic properties of 2^+ , the $^{31}\text{P}\{^1\text{H}\}$ NMR spectrum exhibits a temperature-invariant AA'BB'CC'DD'EE'X splitting pattern ($\text{X} = ^{195}\text{Pt}$; Figure 2). The relatively high value of the $^1J_{\text{P7,P8}}$ and $^1J_{\text{P3,P4}}$ coupling constants (289.6 Hz) matches

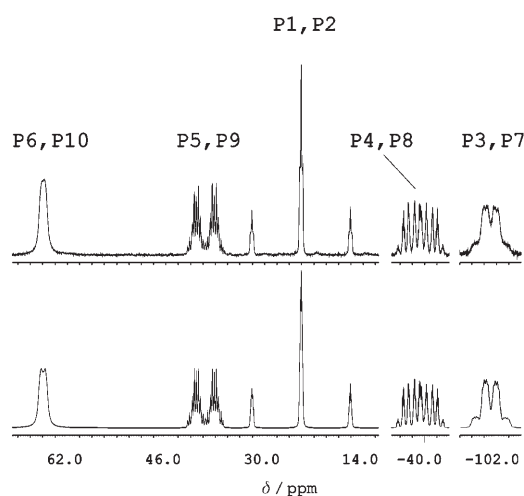


Figure 2. Computed (bottom) and experimental (top) $^{31}\text{P}\{^1\text{H}\}$ NMR spectra of 2^+ (CD_2Cl_2 , 161.89 MHz, 294 K). The labeling is the same as in Figure 1.

that found in other diphosphene complexes and supports the partial double-bond character of these bonds. The coupling constants of the two naked diphosphene P atoms with platinum ($^1J_{\text{Pt,P3}} = ^1J_{\text{Pt,P7}} = 525.0$ Hz), which are much smaller than those with the triphenylphosphine P atoms ($^1J_{\text{Pt,P1}} = ^1J_{\text{Pt,P2}} = 2585.0$ Hz), are also worth noticing. A similar behavior has been observed for other compounds containing naked P atoms or units coordinated to the $\text{Pt}(\text{PPh}_3)_2$ fragment^[11] and has been ascribed to the nature of the M-P bond.^[17]

The $\mu,\eta^{1:2:1}\text{-P}_3$ ligands found in 2^+ belong to a growing and important family of *catena*-phosphorus compounds whose systematization has been proposed recently.^[18] Whereas neutral^[19] and anionic *catena*-phosphorus species^[20] are relatively well known, *catena*-phosphorus cations are still quite rare^[18] and no examples of polyphosphorus zwitterions have been reported to date. Additionally, none of the known linear *catena*-phosphorus cations has been so far obtained from white phosphorus.^[21]

This work demonstrates that the stepwise cooperation of two different transition metal units may bring about the transformation of P_4 into intriguing organophosphorus compounds. Thus, the known reaction of white phosphorus with

Co²⁺ ions and dppm may be viewed as the first step of a modular process where the carbene-like PtL₂ unit selectively cuts the P₆ chain to yield two doubly metalated *catena*-zwitterionic diphenyl(alkyl)phosphonium(+)diphosphonide(−) ligands. The resulting RPh₂P⁽⁺⁾–P=P^(−) unit is the first example of an unsaturated phosphonium diphosphonide ligand tethering two different metal units and may be a potentially versatile functional organophosphorus group to be used with organic and organometallic species.

Experimental Section

2-BF₄: Addition of solid [Pt(PPh₃)₂(η²-C₂H₄)]^[22] (90 mg, 0.12 mmol) to a stirred solution of **1**-BF₄ (125 mg, 0.12 mmol) in 10 mL of CH₂Cl₂ under nitrogen at room temperature caused an immediate color change from dark red to dark brown. ³¹P{¹H} NMR analysis of this solution showed the complete formation of **2**-BF₄. Addition of ethanol (4 mL) and slow concentration under nitrogen gave dark brown crystals suitable for X-ray crystallography. These crystals were filtered off, washed with *n*-pentane, and dried in a stream of nitrogen. Yield: 168.7 mg (80 %). C, H analysis (%) calcd for C₈₆H₇₄BCoF₄P₁₀Pt (1758.1): C 58.75, H 4.24; found: C 58.94, H 4.29. ESI-MS: *m/z* (%) 1670.21 (51) [Co(μ₂η^{1,2,1}-P=P-PPh₂CH₂PPh₂)₂{Pt(PPh₃)₂}]⁺, 835.10 (100) [M²⁺]; ¹H NMR (400.13 MHz, CD₂Cl₂, 294 K): δ = 7.4–6.3 (m, 70H; Ph), 3.2 (m, 2H; CH₂), 2.2 ppm (m, 2H; CH₂); ³¹P{¹H} NMR: See Figure 2 and the Supporting Information.

Details of the crystal-structure determinations of **2**-BF₄·2EtOH·CH₂Cl₂ are provided in the the Supporting Information, which also includes complete ORTEP drawings, ³¹P NMR spectroscopic data, and copies of the ³¹P{¹H} NMR spectra.

Received: January 29, 2008

Published online: April 15, 2008

Keywords: cobalt · heterometallic complexes · phosphorus · platinum · zwitterions

- [1] a) D. E. C. Corbridge, *Phosphorus—An Outline of its Chemistry, Biochemistry and Technology*, 5th ed., Elsevier, Amsterdam, NL, **1995**; b) J. Emsley, *The 13th Element: The Sordid Tale of Murder, Fire, and Phosphorus*, Wiley, New York, **2002**.
- [2] J. M. Lynam, *Angew. Chem.* **2008**, *120*, 843–845; *Angew. Chem. Int. Ed.* **2008**, *47*, 831–833.
- [3] M. Peruzzini, L. Gonsalvi, A. Romerosa, *Chem. Soc. Rev.* **2005**, *34*, 1038–1047.
- [4] P. Barbaro, M. Di Vaira, S. Seniori Costantini, M. Peruzzini, P. Stoppioni, *Chem. Eur. J.* **2007**, *13*, 6682–6690.
- [5] D. Yakhvarov, P. Barbaro, L. Gonsalvi, S. M. Carpio, S. Midollini, A. Orlandini, M. Peruzzini, O. Sinyashin, F. Zanobini, *Angew. Chem.* **2006**, *118*, 4288–4291; *Angew. Chem. Int. Ed.* **2006**, *45*, 4182–4185.
- [6] F. H. Stephens, M. J. A. Johnson, C. C. Cummins, O. P. Kryatova, S. V. Kryatov, E. V. Rybak-Akimova, J. E. McDonough, C. D. Hoff, *J. Am. Chem. Soc.* **2005**, *127*, 15191–15200.
- [7] a) J. D. Masuda, W. W. Schoeller, B. Donnadiou, G. Bertrand, *Angew. Chem.* **2007**, *119*, 7182–7185; *Angew. Chem. Int. Ed.* **2007**, *46*, 7052–7055; b) J. D. Masuda, W. W. Schoeller, B. Donnadiou, G. Bertrand, *J. Am. Chem. Soc.* **2007**, *129*, 14180–14181.
- [8] a) F. Cecconi, C. A. Ghilardi, S. Midollini, A. Orlandini, *J. Am. Chem. Soc.* **1984**, *106*, 3667–3668; b) F. Cecconi, C. A. Ghilardi, S. Midollini, A. Orlandini, *Inorg. Chem.* **1986**, *25*, 1766–1770.
- [9] For leading references on the coordination chemistry of white phosphorus, see: a) M. Peruzzini, R. R. Abdreimova, Y. Budnikova, A. Romerosa, O. J. Scherer, H. Sitzmann, *J. Organomet. Chem.* **2004**, *689*, 4319–4331; b) M. Ehse, A. Romerosa, M. Peruzzini, *Top. Curr. Chem.* **2002**, *220*, 107–140; c) K. H. Whitmire, *Adv. Organomet. Chem.* **1998**, *42*, 1–145; d) O. J. Scherer, *Acc. Chem. Res.* **1999**, *32*, 751–762; e) M. Peruzzini, I. de Los Rios, A. Romerosa, F. Vizza, *Eur. J. Inorg. Chem.* **2001**, 593–608; f) C. C. Cummins, *Angew. Chem.* **2006**, *118*, 876–884; *Angew. Chem. Int. Ed.* **2006**, *45*, 862–870; g) B. P. Johnson, G. Balazs, M. Scheer, *Top. Curr. Chem.* **2004**, *232*, 1–23.
- [10] The only reaction of **1** to give a defined product was the photochemical reaction with [M(CO)₆], which led to the adduct [Co(Ph₂PCH₂PPh₂PPPPPh₂PCH₂PPh₂){M(CO)₅}]Y (M = Cr, Mo, W.; Y = BF₄, BPh₄) by coordination of one M(CO)₅ fragment to one of the central P atoms of the phosphabutadiene unit^[8b] (S. Midollini, personal communication).
- [11] A few examples of PtL₂ insertion into a P–P bond have been reported. See, for example: a) M. Di Vaira, P. Stoppioni, *Polyhedron* **1994**, *13*, 3045–3051; b) M. Di Vaira, M. Peruzzini, P. Stoppioni, *J. Chem. Soc. Dalton Trans.* **1985**, 291–295; c) M. Scheer, M. Dargatz, A. Rufinska, *J. Organomet. Chem.* **1992**, *440*, 327–333.
- [12] CCDC 676028 contains the supplementary crystallographic data for this paper. These data can be obtained free of charge from The Cambridge Crystallographic Data Centre via www.ccdc.cam.ac.uk/data_request/cif.
- [13] V. D. Romanenko, V. L. Rudzevich, E. B. Rusanov, A. N. Chernega, A. Senio, J.-M. Sotiropoulos, G. Pfister-Guillouzo, M. Sanchez, *J. Chem. Soc. Chem. Commun.* **1995**, 1383–1385.
- [14] a) L. Weber, *Chem. Rev.* **1992**, *92*, 1839–1906; b) M. Yoshifuji, *Pure Appl. Chem.* **1999**, *71*, 503–512; c) A. H. Cowley, N. C. Norman, *Prog. Inorg. Chem.* **1986**, *34*, 1–63.
- [15] The back-donation from the Co^{−1} center to the π*-orbital of the tetraphosphabutadiene ligand in **1** increases the electron density of the central P3–P7 bond, which is therefore more prone to react with the Pt⁰ moiety.
- [16] a) K. M. Flynn, H. Hope, B. D. Murray, M. M. Olmstead, P. P. Power, *J. Am. Chem. Soc.* **1983**, *105*, 7750–7751. For other examples of side-on and end-on bonded diphosphene complexes see reference [13a] and b) R. Mathieu, A.-M. Caminade, J.-P. Majoral, S. Attali, M. Sanchez, *Organometallics* **1986**, *5*, 1914–1916; c) A. G. del Pozo, A. M. Caminade, F. Dahan, J. P. Majoral, R. Mathieu, *J. Chem. Soc. Chem. Commun.* **1988**, 574–575.
- [17] a) M. Di Vaira, L. Sacconi, P. Stoppioni, *J. Organomet. Chem.* **1983**, *250*, 183–195; b) M. Di Vaira, L. Sacconi, *Angew. Chem.* **1982**, *94*, 338; *Angew. Chem. Int. Ed. Engl.* **1982**, *21*, 330.
- [18] C. A. Dyker, N. Burford, *Chem. Asian J.* **2008**, *3*, 28–36, and references therein.
- [19] The chemistry of neutral *catena*-phosphines has been developed by Baudler and coworkers: M. Baudler, K. Glinka, *Chem. Rev.* **1994**, *94*, 1273–1297, and references therein.
- [20] For *catena*-phosphorus anions, see: a) H. G. von Schnering, W. Hönle, *Chem. Rev.* **1988**, *88*, 243–273; b) S. Gómez-Ruiz, E. Hey-Hawkins, *Dalton Trans.* **2007**, 5678–5683; c) S. Gómez-Ruiz, A. Schisler, P. Loennecke, E. Hey-Hawkins, *Chem. Eur. J.* **2007**, *13*, 7974–7982; d) D. Stein, A. Dransfeld, M. Flock, H. Rügger, H. Grützmacher, *Eur. J. Inorg. Chem.* **2006**, 4157–4167.
- [21] The pentaphosphorus cluster cation [P₅X₂]⁺ has been prepared by reaction of PX₃ (X = Br, I) with Ag[Al(OC(CF₃)₃)₄] and P₄: I. Krossing, I. Raabe, *Angew. Chem.* **2001**, *113*, 4544–4547; *Angew. Chem. Int. Ed.* **2001**, *40*, 4406–4409.
- [22] R. Head, *Inorg. Synth.* **1984**, *24*, 214–215.