Element–Element Bond Formation upon Oxidation and Reduction

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Dedicated to Professor Herbert W. Roesky on the occasion of his 85th birthday

Abstract: The redox chemistry of \([\text{Cp}^\text{II} \text{Co}]_2\) is investigated. Both compounds can be oxidized and reduced twice. That way, the monocations \([\text{Cp}^\text{II} \text{Co}]_2(\eta^5 \text{P}^\text{III} \text{E}_4)] X = E = P, X = BF_4 (\text{A} 2), [\text{FAI}] (\text{B} 3); E = As, X = BF_4 (\text{A} 4), [\text{FAI}] (\text{B} 4))

Introduction

Oxidation and reduction reactions have been widely used for element–element bond formation or cleavage reactions. For example, the elemental modifications of sulfur (S8), phosphorus (P4 or red phosphorus), or gray arsenic can be easily degraded in reduction processes by bond cleavages to the corresponding S8\(^{2-}\), P8\(^{3-}\), or As8\(^{3-}\) units, respectively.[1] On the other hand, oxidation reactions lead, in general, to bond formation. In the case of S8, one new S–S bond is formed upon oxidation to S8\(^{2+}\), while for the oxidation of white phosphorus, an additional aggregation takes place to form a P4\(^{3+}\) moiety.[2]

Analogous reactivity can be observed for cyclic phosphines[3,4] or arsines such as Bu3P (E = P, As). Both compounds can be reduced and degraded to E2 fragments by using elemental potassium.[5,6] While this behavior is particularly true for saturated main group compounds containing lone pairs, the situation is different for unsaturated species. In this case, reduction leads in general to the population of \(\pi^*\) orbitals and the formation of radical anions, which can result in bond formations.[4] For example, phosphines[6] diboryl compounds,[6b] and (TPB)Cu+ (TPB = tris[2-(diisopropylphosphino)phenyl]borane)[6c] form new formal one-electron bonds (P–P, B–B, and Cu–B) upon reduction. Polynictogen ligands in the coordination sphere of transition metals also reveal interesting redox chemistry. The redox chemistry of pentaphosphaferrocene \([\text{CP}^\text{III} \text{Fe}]_2(\eta^1 \text{P}_3)]\)[3] \((\text{CP}^\text{III} = \text{pentamethylcyclopentadienyl})\) was studied spectroelectrochemically by Winter and Geiger,[6] and experimentally by our group, generating one oxidized and two reduced products.[7] Depending on the reducing agent, one or two electrons can be transferred to \([\text{CP}^\text{III} \text{Fe}]_2(\eta^1 \text{P}_3)]\). When potassium hydride was used, a single-electron transfer occurred leading to a proposed intermediate \([\text{CP}^\text{III} \text{Fe}]_2(\eta^1 \text{P}_3)]\), which dimerized to the dinuclear complex \([\text{CP}^\text{III} \text{Fe}]_2(\eta^1 \text{P}_3)_2\). When elemental potassium was used in excess, the reaction gave the dianionic species \([\text{CP}^\text{III} \text{Fe}]_2(\eta^1 \text{P}_3)_2^2-\) (A; Scheme 1). On the other hand, when thianthrenium hexafluoroantimonate was used as a strong oxidant, the dinuclear compound \([\text{CP}^\text{III} \text{Fe}]_2(\eta^1 \text{P}_3)_2^2-\) (B) was isolated. In these reactions, the polyphosphorus ligand in \([\text{CP}^\text{III} \text{Fe}]_2(\eta^1 \text{P}_3)_2\) responds to the addition or withdrawal of one electron either by folding the P5 ring or by forming a new external P–P bond.[7] The redox chemistry of the analogous arsenic compound \([\text{CP}^\text{III} \text{Fe}]_2(\eta^1 \text{As}_3)_2]\) leads, upon reduction, to a mixture of anionic species containing As8, As10, As10+, and As11 ligands, while all attempts

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Scheme 1. Selected examples of reduced and oxidized polynictogen ligand complexes.
towards an oxidation have failed thus far.\[6\] Similar behavior, including the formation of external E–E bonds upon oxidation, was reported for the complexes \([\text{CpMo(CO)}_3_{(\mu_1,\mu_1,\eta^1-\eta^1-E,\eta^1-E)}] (E = \text{P–Bi}), which results for \(E = \text{P}. As in the formation of the complexes C and D (Scheme 1).\[9\]

Not only sandwich complexes with an \(E_n\) ligand as a lower deck show an instructive redox chemistry, but also homometallic triple-decker complexes with the corresponding \(E_n\) ligand as a middle deck. \([\text{Cp}^5\text{Mo}](\mu_1,\mu_1,\eta^1-\eta^1-P)_2]\), for example, can be easily oxidized to \([\text{Cp}^5\text{Mo}](\mu_1,\mu_1,\eta^1-\eta^1-P)_2]^+ (E), retaining its initial triple-decker geometry in the solid state, whereas the cyclo-P$_n$ ligand in E tends to slightly distort in a bis-allylic manner.\[10\] Therefore, in contrast to the usual polypnictogen complexes, here, oxidations have the opposite effect, namely of elongating P–P bonds, but of also strengthening Mo–Mo bonds.

Intrigued by the diversity of the structural changes observed upon oxidation and/or reduction of the polypnictogen (E$_n$) ligand complexes, we were interested in using E$_n$ ligand complexes that combine the features of triple-decker complexes and separated E$_n$ units, and we were keen to explore whether the redox behavior follows the traditional pathway (oxidation: forming a bond; reduction: cleaving a bond) or whether new avenues to novel and structurally unprecedented products are opened. Therefore, we investigated the redox chemistry of the cobalt complexes \([\text{Cp}'''\text{Co}](\mu_1,\mu_1,\eta^1-\eta^1-E,\eta^1-E)_2]\) (E = \(1\)), \(2\)): \(\text{Cp}''''' = \text{1,2,4-tris-}(\text{tert-butyl})\text{cyclopentadienyl})\), which are easily accessible by the reaction of the toluen complex \([\text{Cp}'''\text{Co}](\mu_1,\mu_1,\eta^1-\eta^1-\text{C}_2\text{H}_4)]\) with white phosphorus or yellow arsenic on gram scale.\[11\]

**Results and Discussion**

To gain first insight into their redox properties, the frontier molecular orbitals of 1 and 2 were computed (Figure 1). The HOMO shows bonding character within the E$_2$ units and antibonding character between them. Therefore, the abstraction of electrons by oxidation should induce a cyclization while the E–E distance of the former E$_2$ unit will be elongated. The situation for the LUMO, which is a linear combination of the \(\pi^*\) orbitals of the E$_2$ units, is the other way around. The LUMO shows bonding character between the two separated E$_2$ units and antibonding character within the E$_2$ units. Again, the population of this orbital by adding electrons through reduction should induce a cyclization to form an E$_3$ ligand while the bond within each E$_2$ unit will be elongated.

To obtain an overview of how redox processes proceed, cyclic voltammetry measurements were performed in different solvents. We found that the nature of the redox process is strongly dependent on the solvent used. In the case of 1, there are two reversible oxidation processes in CH$_2$Cl$_2$ at –367 and +351 mV and at most, irreversible reduction around –2350 mV vs. \([\text{Cp}_2\text{Fe}]^+/\text{Cp}_2\text{Fe}^+\). When the solvent is changed to THF, only one reversible oxidation at –336 mV and one reversible reduction at –2304 mV were observed (Figure 2). The cyclic voltammogram of 2 in DME shows two reversible oxidations at –463 and +48 mV and two reversible reductions at –2144 and –2644 mV vs. \([\text{Cp}_2\text{Fe}]^+/\text{Cp}_2\text{Fe}^+\) (see Figure 3).

For the chemical oxidation, we chose Ag$^+$ as a suitable oxidant, because of the rather low oxidation potentials of 1 and 2.\[12\]

Both compounds can be oxidized by using one equivalent of an Ag$^+$ salt containing a weakly coordinating anion [X] (X = BF$_3$ or [FAI]: [FAI] = [FAI(OC$_3$F$_7$)(C$_5$F$_3$)]) leading to the isostroical compounds \([\text{Cp}'''''\text{Co}](\mu_1,\mu_1,\eta^1-\eta^1-E,\eta^1-E)_2][\text{X}] [E = \text{P}, X = \text{BF}_3 (3a; 66 %), [FAI] (3b; 56 %); E = \text{As}, X = \text{BF}_3 (4a; 65 %), [FAI] (4b; 42 %); Eq. (1)], in which two new E–E bonds have been formed to generate a new triple-decker sandwich complex with a cyclobutadiene-like formal E$_3^-$ middle deck.
might be viewed as cyclobutadiene-like units. In 3b, the E₄ ligand has a trapezoid shape with three shorter P–P bonds (P₁–P₂ 2.2205(6), P₂–P₃ 2.2358(6), P₃–P₄ 2.2069(7) Å) and one longer one (P₁–P₄ 2.3139(6) Å). In 4b, the As₄ ligand is disordered over three positions with site occupancies of 5, 25, and 70%, preventing the accurate description of the As₄ unit.

DFT optimization of the geometries of 3a and 3b (BP86/def2-TZVP level of theory; 3a', 3b') revealed a rectangular E₄ ligand (similar to that observed in the crystal structure) for 3a and a trapezoid-shaped ligand for 3b, which is more widened than in 3b (2.5133 Å (optimized geometry of 3b')) and 2.3139(6) Å (experimental structure of 3b)). The geometry optimizations were both started from the atomic coordinates obtained from the X-ray structures. However, starting from a symmetric cyclo-P₄ ligand, this geometry is retained in the optimized structure of 3a', the same as for the final trapezoid-shaped ligand in 3b'. The energy difference between the two isomers is only 3 kJ mol⁻¹, with the trapezoid isomer being favored. Both geometries represent minima on the energy hypersurface (see the Supporting Information). The structural differences between 3a and 3b can be attributed to packing effects as the counterions differ significantly in size. DFT calculations show a similar behavior for the arsenic compounds 4a and 4b. The formation of one P–P bond and the generation of a trapezoid-shaped ligand are reminiscent of the coordination of 1 to [W(CO)₅] fragments leading to ([(Cp"Co)₂(η⁴−η³−η²−η¹−1)P₄][W(CO)₅]) (13). However, in this complex, the P₁–P₂ and P₃–P₄ distances are significantly shorter (2.070(8) and 2.093(9) Å), the P₂–P₃ distance is quite the same (2.276(8) Å), and the P₁–P₄ distance is longer (2.962(8) Å) than the corresponding distances in 3b. An analogous As compound is unknown. The compounds 3a/3b and 4a/4b are paramagnetic. The ¹H NMR spectra in solution at room temperature reveal strongly shifted broad signals for the Cp⁻ ligands. The EPR spectra (solid and in frozen solution at 77 K) show an isotropic resonance with the gₘ values indicating one unpaired electron each (77 K, solid: 3a: gₘ = 2.037; 3b: gₘ = 2.024; 4a: gₘ = 2.121; 4b: gₘ = 2.094; see the Supporting Information). The Evans NMR spectra reveal effective magnetic moments of 1.23 μₑ and of 2.01 μₑ for 3b and of 2.01 μₑ for 4b, corresponding to about one unpaired electron each.

The pnictogen ligands in the starting materials 1 and 2 can be described as separated E₄⁻⁻⁻⁻ units (if Co₃ moieties are assumed). Upon oxidation, one electron is removed, and formally, a cyclo-E₄⁻⁻⁻⁻ ligand is formed. The SOMO (3a: Ω = -224, see the Supporting Information) reveals that the unpaired electron is located mainly on the E₄⁻⁻⁻⁻ ligand (electron density within the short E–E bonds and between them with a minor contribution of the Co atoms, see the Supporting Information). The spin density is homogeneously distributed over all E and Co atoms, and the absence of a hyperfine coupling to Co in the EPR spectra also underlines the presence of a formal cyclo-E₄⁻⁻⁻⁻ ligand. Likewise, the decrease in the formal charges (Mulliken charges, see the Supporting Information) of the E₄ ligand as compared to the starting complexes emphasizes this description. It has to be noted that this description of 3a and 4a is only a formalism as the electron density in the SOMOs is also distributed over

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Figure 3. Cyclic voltammogram of 2 in DME vs. [Cp₃Fe]/[Cp₃Fe]⁺ (electrolyte N′Bu₄PF₆, scan rate: 100 mVs⁻¹, room temperature).

Figure 4. Structure of the cations in 3a (left) and 4a (right) in the solid state. Thermal ellipsoids set at 50% probability. Hydrogen atoms, anions and solvent molecules are omitted for clarity.

Crystals suitable for single-crystal X-ray structure analysis were obtained from concentrated solutions in CH₂Cl₂, layered with pentane (3b, 4a, 4b) or hexane (3a) at −30°C. As the compounds 3a/3b and 4a/4b are isostructural and differ solely in the used anion, only the structures of 3a and 4a in the solid state are depicted in Figure 4.

The structures reveal triple-decker complexes with cyclo-E₄ ligands coordinating in an η⁴−η³ fashion to two [Cp"Co] fragments. The geometry of the cyclo-E₄ ligand is slightly distorted and differs a little in all compounds. For 3a and 4a, there are rectangular cyclo-E₄ ligands with two shorter E–E bonds (P₁–P₂ 2.1860(9), P₂–P₃ 2.1837(8), As₁–As₂/As₃–As₄ 2.3882(2) Å) and two longer E–E bonds (P₁–P₄ 2.3022(8), P₂–P₃ 2.2960(8), As₁–As₄/As₂–As₃ 2.5198(2) Å), which
both Co atoms and the Cp" ligands, and the covalent character of the bonding between the E¢ ligand and the metal has to be considered.

Upon reacting 1 and 2 with Ag[TEF] ([TEF] = [Al(OC-(CF$_3$)$_3$)$_2$]), the dications [(Co"$_2$(μ-η"'-η"-Et)"_2][TEF] (E = P (5), As (6)) were obtained as crystalline compounds in yields of 68 and 69%, respectively [Eq. (2) in Figure 5], which now represent cyclo-E¢$_2$-middle decks with four equivalent bonds.

The structures of 5 and 6 in the solid state (Figure 5) show triple-decker complexes with a cyclo-E¢ ligand as a middle deck. In 5, the P$_2$ unit is planar with very similar P–P distances (2.236(2) and 2.239(2) Å) representing single bonds, which was confirmed by the Wiberg bond indices (WBI) of 0.92.[14] The P–P bonds in 5 are longer than those in the complexes with a cyclo-P$_2$ ligand as an end-deck as in [Cp"Co$_2$(μ-η'-η-E)$_2$] (2.1557(18)-2.1699(14) Å).[15] In contrast to 5, the cyclo-As$_4$ ligand in 6 is slightly folded (fold angle 3°). The As–As bond lengths are between 2.4355(9) and 2.4759(9) Å and lie in the range of single bonds, as confirmed by WBIs between 0.80 and 0.93 (the DFT-optimized geometry reveals a trapezoid-shaped As$_4$ ligand with a fold angle of 6.4°).[14] The As–As distances are also longer than in complexes with a cyclo-As$_4$ ligand as an end-deck such as [Cp"Nb(CO)$_2$(η'-η-As)$_4$] (2.345(4)-2.409(4) Å).[14] The structural motif of the dication is reminiscent of the Cp$_2$Mo (C$_4$Me$_3$Bu)-substituted compound [(Cp$_2$Mo)$_2$(μ-η'-η-As)$_4$] [CoCl$_2$(thf)$_2$], obtained, however, by starting from As$_4$(SiMe$_3$)$_4$ as an As source.[17] This complex shows As–As distances of 2.4552(10) and 2.4680(11) Å. In the 1H NMR spectrum of 5, two broad singlets centered at δ = 1.82 and 1.71 ppm for the 'Bu groups of the Cp" ligand can be detected, but no signals for the H atoms bonded to the Cp ring, indicating a dynamic process in solution. The 30P[1H] NMR spectrum at room temperature shows one broad singlet at δ = 494.0 ppm (ν$_{os}$ = 4000 Hz). Upon cooling to −80°C, the signals in the 1H NMR spectra broaden further, and still no resonance for the Cp-bonded H atoms can be found. The signal in the 30P[1H] NMR spectra broadens too and eventually disappears completely. Upon warming an NMR sample of 5 in ortho-difluorobenzene containing a C$_6$D$_6$ capillary to 80°C, the signals in the 1H NMR spectra remained unchanged. In the 30P[1H] NMR spectrum, a singlet can be observed whose intensity decreases at 80°C. In the 1H NMR spectrum of 6, three singlets centered at δ = 6.50, 1.66, and 1.50 ppm can be detected, with the signal for the Cp-bonded H atoms (6.50 ppm) being downfield-shifted by approximately 2 ppm in comparison to the starting material. Both 5 and 6 are EPR-silent.

The abstraction of another electron from the E¢$_2$ ligand in 3 and 4 leads to a further change in the geometry, and the ligand can now be described as a cyclo-E¢$_2$ ligand. Identical P–P bonds in the completely planar middle deck of 5 indicate the presence of an aromatic P$_2$¢ ligand. The HOMO of 5 shows the electron density to be homogeneously distributed over all P atoms (indicating four equivalent bonds) and both Co atoms (in the shape of a d orbital, see the Supporting Information). In the case of 6, the As¢$_2$ ligand is not completely planar, and the As–As distances differ slightly from each other. The optimized geometry shows a trapezoid-shaped ligand with one side being more open than the other. The structure in the solid state suggests that, therefore, the HOMO shows electron density within the three shorter As–As bonds and at the Co atoms (d orbitals). As the formal charges ( Mulliken charges, see the Supporting Information) of the E¢ ligand in 5 and 6 decrease further compared to the monocations and relative to 1 and 2, the formal description of the ligand as E¢$_2$ seems appropriate.

All mentioned oxidations of 1 and 2 are fully reversible. The addition of stoichiometric amounts of KC$_6$ selectively yields back the starting materials 1 and 2. In the related cyclic voltammograms, both compounds 1 and 2 show reversible reduction processes at rather negative redox potentials.

Potassium graphite was chosen as a suitable reducing agent. By using a small excess of > 1 equiv of KC$_6$, the monoanions [K(18-c-6)(dme)]$_2$[(Cp"Co)$_2$(μ-η'-η-As)$_2$] (E = P (7), As (8)) can be obtained in crystalline yields of 56 and 40%, respectively, which now feature cyclobutadiene-like E¢$_2$-middle decks [Eq. (3)]. Because of their anionic character, 7 and 8 are extremely sensitive towards air and moisture.

The structures in the solid state (Figure 6) reveal a rectangular cyclo-E¢ middle deck for both compounds. Two shorter E–E bonds (P1–P2/P3–P4 2.1288(9) Å, As1–As2/As3–As4 2.3074(16) Å) and two longer E–E bonds (P1–P4/P2–P3 2.3606(8) Å, As1–As4/As2–As3 2.5852(15) Å) are present. All bonds are in the range of shortened and elongated single bonds, respectively, as also confirmed by the WBIs for the short (7: 1.18, 8: 1.15) and long bonds (7: 0.68, 8: 0.63).[14,18]

Figure 5. Structure of the dications in 5 (left) and 6 (right) in the solid state. Thermal ellipsoids set at 50% probability. Hydrogen atoms, anions, and solvent molecules are omitted for clarity.
Compounds 7 and 8 are both paramagnetic as indicated by the paramagnetic shift of the signals of the Cp” ligand in the 1H NMR spectra in solution and by the absence of signals in the 31P{1H} NMR spectrum (for 7). Both compounds are EPR-active and show resonances in solution (frozen solution) and in the solid state (at room temperature and at 77 K). The spectra of both compounds show a rhombically structured signal at 77 K in the solid state (7: \( g_x = 2.0840, g_y = 2.0638, g_z = 1.9897 \); 8: \( g_x = 2.2765, g_y = 2.0724, g_z = 2.0752 \)). The effective magnetic moment was determined by the Evans method to be 1.93 \( \mu_B \) (7) and 2.35 \( \mu_B \) (8), respectively, corresponding to about one unpaired electron each.

The DFT calculations reveal that the SOMO (7: \( \alpha = -189 \); 8: \( \alpha = -225 \)) is mainly located between \( E_1 @ E_4 \) and \( E_2 @ E_3 \) in an bonding fashion with some participation of a \( d \) orbital of the Co atoms and an antibonding \( \pi \) orbital of the Cp” ligands. The spin density is spread over the \( E_1 \) ligand and the Co atoms while being mainly located on Co, explaining the rhombic signals in the EPR spectra of 7 and 8. The summed up Mulliken charges (see the Supporting Information) also indicate that the additional electron was transferred to the \( E_4 \) ligand and the formal description as \( E_4 @ \) moieties seems to be appropriate, although the spin density distribution indicates a partial charge delocalization also on the Co atoms and the Cp” ligands.

According to the cyclic voltammograms, 1 shows only one reversible reduction while 2 reveals two of them. Based on this observation, we attempted to access the doubly reduced product of 1. Indeed, the reduction of 1 leads to the dianionic compound \([K(18-c-6)(dme)\_2][K(18-c-6)][(Cp”Co)(\mu_3,\mu_3\eta^1:\eta^1-P_4)] \) (9), which was obtained in crystalline yields of 75\% [Eq. (4)]. Probably, during the reaction, first the monoanion 7 is formed followed by a further reduction to the dianion 9. The second reduction step occurs by cleavage of a P–P bond in 7 followed by rearrangement of the P unit. Compound 9 is extremely sensitive towards air and moisture.

The structure of 9 in the solid state (Figure 7) reveals a prismatic structural motif consisting of two \([Cp”Co]\) fragments and four P atoms. There are two shorter P–P bonds (P1–P2 2.1582(11), P3–P4 2.1615(11) Å) and one longer one (P2–P3 2.2646(10) Å). The P1–P4 distance is with 2.8396(12) Å too long for a P–P bond, but below the sum of the van der Waals radii (\( S_{vdW} = 3.80 \) Å).\(^{19} \) Therefore, an interaction between the two nuclei can be expected. This is indeed confirmed by the WBI of 0.24. The \( P_1 \) ligand can be best described as being butadiene-like. The Co2P4 scaffold is also isoelectronic to the Ni2P4 unit in the complex \((Cp”Ni)(\mu_3,\mu_3\eta^1:\eta^1-P_4) \)\(^{22} \).
In the $^{31}$P($^1$H) NMR spectrum of 9 in THF, two doublets centered at $\delta = 20.6$ and $-7.5$ ppm with a $J_{PP}$ coupling constant of 263 Hz can be assigned to the two inequivalent P atoms in 9. The fact that the cyclic P$_{8}$ ligand in 7 rearranges upon addition of another electron is not obvious. The first step of the reaction would be formally a reduction to 

$$[(\text{Cp}^*\text{Co})_2(\mu_1\eta^1\eta^2\eta^2\text{P}_3)]^- \quad (1\text{-}9\text{a})$$

However, 1-9a is 41.46 kJ mol$^{-1}$ higher in energy than the anion of 9. The coordination of a [K(18-c-6)]$^+$ unit to the edge of the P$_8$ ligand in 9 contributes with 193.73 kJ mol$^{-1}$ to the stabilization of the product. Interestingly, when the potassium counternions were fully separated from the P$_8$ ligand complex by using 2,2,2-coupling to the stabilization of the 8-19 it cannot be obtained by the reduction of 9. Hence, by reacting 1 with an excess of potassium graphite in the presence of the 2,2,2-coupling, only the monoanion 7 is obtained.

Both reductions of 1 and the reduction of 2 are fully reversible. The addition of stoichiometric amounts of AgBF$_4$ to 7, 8, and 9, respectively, yields selectively the related starting materials. While the Co$_2$As$_3$ prism is known as a structural motif of the samarium compound $[(\text{Cp}^*\text{Co})_2(\mu_1\eta^1\eta^2\eta^2\text{Sm})]$ it cannot be obtained by the reduction of 2 with two equivalents of potassium graphite. Probably, an additional coordination to a Lewis acid such as Cp*Sm is needed to obtain a stable compound. From the stoichiometric reaction, only the monoanion 8 can be isolated. By using an excess of potassium (fourfold or higher), one arsenic atom is abstracted, and [K(dme)$_2$]$][(\text{Cp}^*\text{Co})_2(\mu_1\eta^1\eta^2\eta^2\text{As}_3)]$ (10) can be isolated in crystalline yields of 26% [Eq. (5)]. In this reaction, the formation of a black precipitate is observed. Compound 10 is extremely sensitive towards air and moisture.

The structure in the solid state (Figure 8) reveals a bent triple-decker complex with an allylic As$_2$ ligand. The As$_1$–As$_2$ (2.4216(4) Å) and As$_2$–As$_3$ (2.4055(4) Å) bonds are in the range of single bonds, which was also confirmed by WBI of 0.99.[24] The As$_1$–As$_3$ distance is 3.0224(5) Å, and too long to be considered as a bond. As the distance is still below the sum of the van der Waals radii ($\Sigma_{\text{vdW}} = 3.76$ Å), an interaction between the two atoms can be expected, which was confirmed by a WBI of 0.20.[19] Alternatively, 10 can be described as a nido cluster with a square-pyramidal Co$_3$As$_2$ core according to the Wade–Mings rules. There are barely any triple-decker sandwich complexes known that include Cp ligands and a P$_3$ ligand as a middle deck, and none with an As$_3$ ligand. The Ni compounds $[(\text{Cp}^*\text{Ni})_2(\mu_1\eta^1\eta^2\eta^2\text{P}_3)]^+$, $[(\text{Cp}^*\text{Ni})_2(\mu_1\eta^1\eta^2\eta^2\text{P}_3)]^-$, and the heterobimetallic $[(\text{Cp}^*\text{Co})(\text{Cp}^*\text{Ni})(\mu_1\eta^1\eta^2\eta^2\text{P}_3)]$ have been reported.[23] While the nickel complexes formally possess one or two electrons more than 10, the latter is isoelectronic to 10. Therefore, 10 represents the first example of a triple-decker complex with an allylic As$_2$ ligand.

**Conclusion**

We have shown that the cobalt complexes $[(\text{Cp}^*\text{Co})_2(\mu_1\eta^1\eta^2\eta^1\text{E}_3)]$ (E = P (1), As (2)) exhibit a unique redox chemistry far from the usual behavior of a triple-decker complex or polynictogen rings and cages. Instead, they display a distinctive platform where E-E bond formations can be performed using both oxidation and reduction reactions, with these processes being completely reversible. By reduction and oxidation, the two separate E$_2$P$_2$ units in 1 and 2, respectively, can be transformed into cyclobutadiene-like E$_2$P$_2$ moieties (3a, 3b, 4a, 4b) and E$_4$P$_6$ moieties (7, 8), respectively, or finally into cyclo-E$_4$P$_2$ ligands in 5 and 6. Further reduction of the monoanions 7 and 8 leads either to the cleavage of a P–P bond followed by rearrangement into a Co$_2$P$_3$ scaffold or, in the case of an (As$_2$)$_2$ unit, to the abstraction of one arsenic atom to yield a bent triple-decker complex with an allylic As$_2$ ligand as a middle deck (10), which represents the first example of a triple-decker complex with such a ligand. All redox processes (except for the formation of 10) are fully reversible and can be selectively reversed when stoichiometric amounts of KC$_6$ or AgBF$_4$, respectively, are added. Even the formation of two P–P bonds as well as the cleavage of one P–P bond in the formation of 9 can be reverted stepwise upon addition of one or two equivalents of AgBF$_4$. Moreover, these results also clearly show the similarities and the different behavior of the (As$_2$)$_2$ entity in 2 and the (P$_3$)$_2$ moiety in 1 within these redox processes.

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Conflict of interest

The authors declare no conflict of interest.

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