

Functionalization of Pentaphosphorus Cations by Complexation**

Anup K. Adhikari, Christoph G. P. Ziegler, Kai Schwedtmann, Clemens Taube, Jan. J. Weigand* und Robert Wolf*

Abstract: The chemistry of polyphosphorus cations has rapidly developed in recent years, but their coordination behavior has remained mostly unexplored. Herein, we describe the reactivity of $[P_5R_2]^+$ cations with cyclopentadienyl metal complexes. The reaction of $[Cp^{Ar}Fe(\mu-Br)]_2$ ($Cp^{Ar} = C_5(C_6H_4-4-Et)_5$) with $[P_5R_2][GaCl_4]$ ($R = iPr$ and 2,4,6-Me₃C₆H₂ (Mes)) afforded bicyclo[1.1.0]pentaphosphanes (**1-R**, $R = iPr$ and Mes), showing an unsymmetric „butterfly“ structure. The same products **1-R** were formed from $K[Cp^{Ar}]$ and $[P_5R_2][GaCl_4]$. The cationic complexes $[Cp^{Ar}Co(\eta^4-P_5R_2)][GaCl_4]$ (**2-R** [$GaCl_4$], $R = iPr$ and Cy) and $[(Cp^{Ar}Ni)_2(\eta^{3,3}-P_5R_2)][GaCl_4]$ (**3-R** [$GaCl_4$]) were obtained from $[P_5R_2][GaCl_4]$ and $[Cp^{Ar}M(\mu-Br)]_2$ ($M = Co$ and Ni) as well as by using low-valent „ $Cp^{Ar}M^1$ “ sources. Anion metathesis of **2-R** [$GaCl_4$] and **3-R** [$GaCl_4$] was achieved with $Na[BArF_{24}]$. The P_5 framework of the resulting salts **2-R** [$BArF_{24}$] can be further functionalized with nucleophiles. Thus reactions with $[Et_4N]X$ ($X = CN$ and Cl) give unprecedented cyano- and chloro-functionalized complexes, while organo-functionalization was achieved with $CyMgCl$.

Because of the large P–P single bond energy (ca. 200 kJ mol⁻¹ in typical compounds), the structural diversity of polyphosphorus derivatives is unrivalled in main group chemistry.^[1–6] Seminal contributions of Baudler, von Schnering, and many others have unraveled a plethora of neutral polyphosphanes and polyphosphanide anions,^[2–5] and the coordination behavior of such species has been investigated extensively in the past.^[5,6] Transition-metal polyphosphides such as pentamethylpentaphosphaferrocene are used as building blocks for 1D and 2D polymers and spherical supramolecules.^[7,8] Alkylation, protonation, halogenation,

reduction, and oxidation reactions of polyphosphorus units in neutral transition-metal complexes have been reported,^[9] while the reaction of main-group-element nucleophiles such as amides, phosphanides, hydrocarbyl anions, and N-heterocyclic carbenes (NHCs) is a versatile route to anionic complexes with functionalized polyphosphido ligands.^[10] In addition, Cummins and co-workers have very impressively demonstrated the synthetic potential of polyphosphido transition metalate anions for the synthesis of useful and novel P compounds.^[6c,11]

In comparison to the well-established neutral and anionic polyphosphorus compounds, the chemistry of polyphosphorus cations has only just begun to flourish a decade ago.^[12] Recent studies clearly show the substantial influence of cationic substituents, for example, NHCs^[13] and phosphanes,^[14] on their stability and reactivity.^[15] However, the coordination chemistry of cationic polyphosphanes has scarcely been studied thus far. In contrast to a variety of neutral compounds **I–IV**^[16–19] (Figure 1) and anionic compounds **V–VII**,^[10,19] cationic polyphosphorus complexes with late transition metals (such as **VIII**) are extremely rare.^[20] In particular, the direct use of cationic polyphosphanes as ligands has not been investigated. This prompted us to study whether cationic polyphosphane complexes are accessible by reaction of

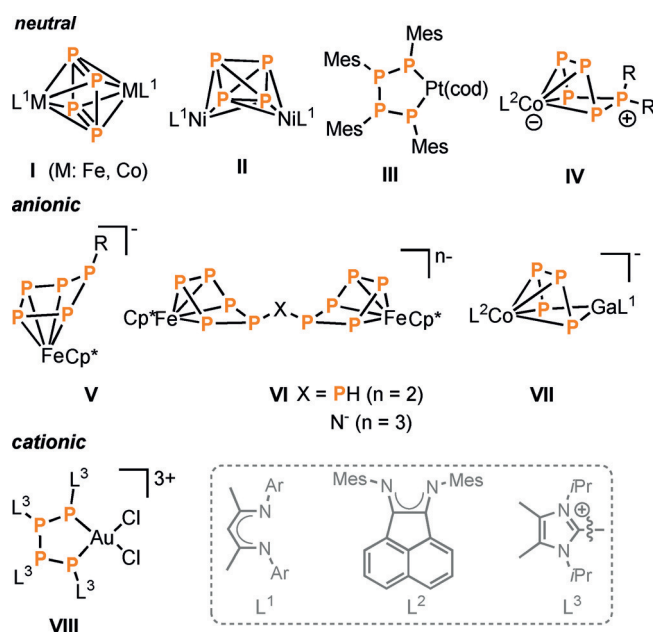


Figure 1. Selected late-transition-metal polyphosphorus complexes with P_4 and P_5 units. Mes = 2,4,6-trimethylphenyl, Cp^* = pentamethylcyclopentadienyl.

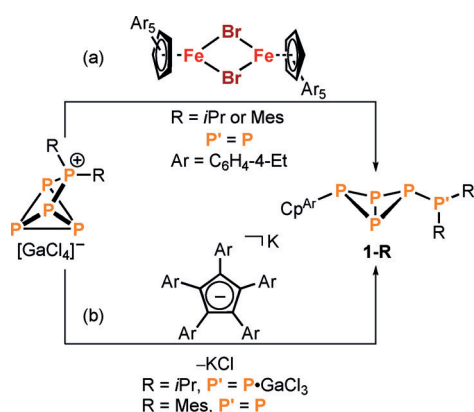
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Scheme 1. Reaction of $[P_5R_2][GaCl_4]$ with a) $[Cp^{Ar}Fe(\mu-Br)]_2$ and b) $K[Cp^{Ar}]$. $Cp^{Ar} = C_5(C_6H_4-4-Et)_5$.

suitable low-valent metal complexes with selected phosphorus cations.

$[P_5R_2]^+$ cations (Scheme 1) seemed to be ideal candidates for such an investigation as they are readily accessible with a wide range of substituents by insertion of carbene-analogous phosphonium cations $[PR_2]^+$ into P–P bonds of white phosphorus.^[21,22] Although their reactivity with elemental sulfur, selenium, and N-heterocyclic carbenes has been studied,^[23,24] their coordination behavior is completely unknown. Herein, we describe reactions of cyclopentadienyl iron, cobalt, and nickel complexes with such cations, which result in the formation of neutral bicyclo[1.1.0]pentaphosphanes **1-R** ($R = iPr$ and Mes) and cationic P_5R_2 complexes $[(Cp^{Ar}Co)(\eta^4-P_5R_2)]^+$ (**[2-R]**⁺; $Cp^{Ar} = C_5(C_6H_4-4-Et)_5$, $R = iPr$ and Cy) and $[(Cp^{Ar}Ni)_2(\eta^{3:3}-P_5R_2)]$ (**[3-R]**⁺). Unusual (pseudo)halogen and organo-substituted P_3R_2X ($X = Cl, CN,$ and Cy) frameworks are the result of post-functionalization reactions of **[2-R]**⁺ with $[Et_4N][X]$ ($X = CN$ and Cl) and the Grignard reagent $CyMgCl$.

As the cyclopentadienyl iron halide complex $[Cp^{Ar}Fe(\mu-Br)]_2$ ^[25] had been successfully used for white phosphorus activation,^[26] it was selected as the starting point for our investigation. To our surprise, the reaction of this Fe complex with $[P_5R_2][GaCl_4]$ salts ($R = iPr$ and Mes ; $Mes = 2,4,6-Me_3C_6H_2$; Scheme 1a)^[23] afforded remarkable, unsymmetrically substituted bicyclo[1.1.0]pentaphosphanes **1-R** as the major product. The isopropyl-substituted pentaphosphane **1-iPr** is formed in the reaction of $[Cp^{Ar}Fe(\mu-Br)]_2$ and $[P_5iPr_2][GaCl_4]$ and can be identified by an apparent AMXY₂ spin system with multiplets at $\delta = -38.9, -87.0, -251.9,$ and -325.0 ppm in the $^{31}P\{^1H\}$ NMR spectrum.

Unfortunately, **1-iPr** rapidly decomposes in solution (even in apolar solvents such as *n*-hexane) into diphosphane P_2iPr_4 , white phosphorus, and further unknown products. Despite its low stability, yellow crystals of the gallium trichloride adduct **1-iPr-GaCl₃** suitable for X-ray crystallography were obtained from the reaction of $K[Cp^{Ar}]$ with $[P_5iPr_2][GaCl_4]$, and the molecular structure is depicted in Figure 2a.^[27] The core of the structure is an unsymmetrical, *exo,exo*-substituted P_4 „butterfly“ unit, showing a similar folding ($P2\cdots P4$ 2.8047(10) Å) as other *exo,exo*-substituted bicyclo[1.1.0]tetraphosphanes.^[28] As a consequence, the bridgehead

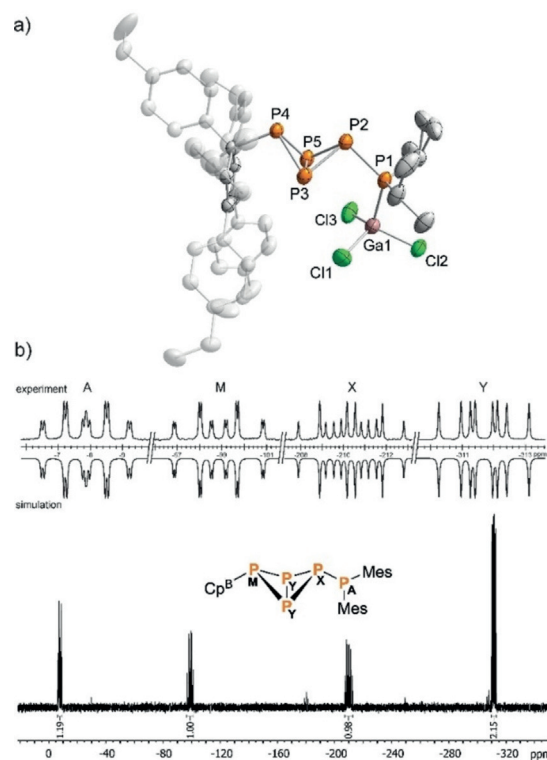
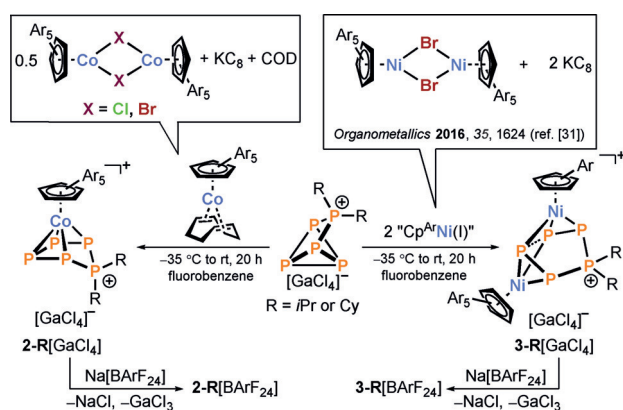


Figure 2. a) Solid-state molecular structure of compound **1-iPr-GaCl₃**. Ellipsoids set at 50% probability; H atoms are omitted for clarity. Selected bond lengths [Å] and angles [°]: P1–P2 2.2104(11), P2–P3 2.2299(12), P2–P5 2.2219(11), P3–P4 2.2124(11), P4–P5 2.2218(11) P3–P5 2.1764(11), P2…P4 2.8047(10), P1–Ga1 2.3857(10); P1–P2–P3 97.57(4), P1–P2–P5 102.19(5), P3–P2–P5 58.53(4), P2–P3–P4 78.31(4), P3–P4–P5 58.79(3), P2–P5–P4 78.28(4). b) $^{31}P\{^1H\}$ NMR spectrum of compound **1-Mes** in C_6D_6 at room temperature. $\delta(P_A) = -7.9$ ppm, $\delta(P_M) = -98.9$ ppm, $\delta(P_X) = -210.4$ ppm, and $\delta(P_Y) = -311.7$ ppm; $^1J_{AX} = -207.3$ Hz, $^1J_{XY} = -162.1$ Hz, $^1J_{MY} = -186.4$ Hz, $^2J_{AY} = 114.2$ Hz, $^2J_{MX} = 268.9$ Hz, and $^3J_{AM} = 15.4$ Hz.

$P3-P5$ bond (2.1764(11) Å) is shorter than the remaining P–P bonds (2.2104(11)–2.2299(12) Å). The $^{31}P\{^1H\}$ NMR spectrum of **1-iPr-GaCl₃** recorded in C_6D_6 (see the Supporting Information, Figure S8) is in agreement with the molecular structure (AMXY₂ spin system; $\delta = 16.4, -107.3, -267.6,$ and -324.2 ppm). Note that the chemical shifts differ from those of **1-iPr** obtained using the Fe complex because of the $GaCl_3$ coordination (see above). As for **1-iPr**, a detailed spectroscopic characterization of **1-iPr-GaCl₃** was hampered by its rapid decomposition in solution. In contrast, however, the mesityl-substituted derivative **1-Mes** can be synthesized by an analogous procedure using $K[Cp^{Ar}]$ and $[P_5Mes_2][GaCl_4]$ (Scheme 1b). **1-Mes** is significantly more stable than **1-iPr** (see Figure S87 for a $^{31}P\{^1H\}$ NMR monitoring study of its decomposition). **1-Mes** can be isolated as a pure, greenish-yellow solid in a moderate yield of 37%. The $^{31}P\{^1H\}$ NMR spectrum of **1-Mes** in C_6D_6 (Figure 2b) shows an AMXY₂ spin system similar to those of related tetraphosphacyclo[1.1.0]butanes.^[27–29] An iterative fit of this spectrum revealed typical $^1J_{PP}$ coupling constants of -207.3 Hz, -162.1 Hz, and -186.4 Hz and a $^2J_{PP}$ coupling constant $^2J_{MX}$ of 268.9 Hz. The latter, comparably very large $^2J_{PP}$ coupling constant indicates

that the lone pairs of P_M and P_X are directed towards one another (see the Supporting Information for the full analysis of this spectrum).^[28]

Reactions of $[P_5R_2][GaCl_4]$ ($R = iPr$ and Cy) with $[Cp^{Ar}M(\mu-Br)_2]$ ($M = Co$, 1 equiv; $M = Ni$, 2 equiv)^[25] afford $[(Cp^{Ar}Co)(\eta^4-P_5R_2)][GaCl_4]$ (**2-R** $[GaCl_4]$) and $[(Cp^{Ar}Ni)_2(\eta^{3,3}-P_5R_2)][GaCl_4]$ (**3-R** $[GaCl_4]$) as dark-red or pink crystalline solids in low yields of 15–25%. These reactions presumably involve a disproportionation of the metal(II) precursors to metal(I) and metal(III) compounds, which is induced by phosphorus ligand coordination, explaining the lower than expected yield. However, **2-R** $[GaCl_4]$ and **3-R** $[GaCl_4]$ can be formed in good to high yield (55–85%) by reacting $[P_5R_2][GaCl_4]$ salts directly with the cobalt(I) and nickel(I) compounds, that is, $[Cp^{Ar}Co(cod)]$ (1 equiv, $cod = 1,5$ -cyclooctadiene)^[30] and the transient „ $Cp^{Ar}Ni^{I+}$ “ (2 equiv, generated from $[Cp^{Ar}Ni(\mu-Br)_2]$ and KC_8 ; Scheme 2).^[31]



Scheme 2. Synthesis of **2-R** $[GaCl_4]$, **3-R** $[GaCl_4]$, **2-R** $[BArF_{24}]$, and **3-R** $[BArF_{24}]$. $R = iPr$ and Cy , $BArF_{24} =$ tetrakis[3,5-bis(trifluoromethyl)phenyl]borate, $Ar = C_6H_4-4-Et$.

Single crystals of **2-R** $[GaCl_4]$ ($R = iPr$ and Cy ; dark red blocks) and **3-*iPr*** $[GaCl_4]$ (pink-violet blocks) were obtained from THF/*n*-hexane at $-35^\circ C$ and analyzed by X-ray crystallography (Figure 3). The pentaphosphorus ligand is η^4 -coordinated to the metal, resulting in an envelope conformation of the P_5 fragment in **2-R** $[GaCl_4]$. The arrangement of the P2, P3, P4, and P5 atoms is almost planar with P–P bonds that are slightly shorter than typical single bonds. For **2-R** $[GaCl_4]$, the P–P bonds of the tetracoordinate P atom (P1) are approximately 0.025 Å longer than those between the coordinated P atoms. **3-*iPr*** $[GaCl_4]$ has an unprecedented molecular structure where two „ $Cp^{Ar}Ni^{I+}$ “ fragments are η^3 -coordinated to a *cyclo*- P_5iPr_2 ring. The P_5 unit is in a twist conformation as evidenced by the dihedral angle between the P2–P3–P4 and P3–P4–P5 planes of -67.9° . This ligand features one unusually long P–P bond (P3–P4 2.5002(16) Å) due to the coordination of each P atom to two Ni centers, while the remaining P–P distances (P1–P2 2.1835(17) Å, P1–P5 2.1859(15) Å, P2–P3 2.1672(16) Å, P4–P5 2.1542(16) Å) are similar to those of **2-R** $[GaCl_4]$.

$^{31}P\{^1H\}$ NMR spectra of **2-R** $[GaCl_4]$ in C_6D_6 show three multiplets of an AMM'XX' spin system in agreement with

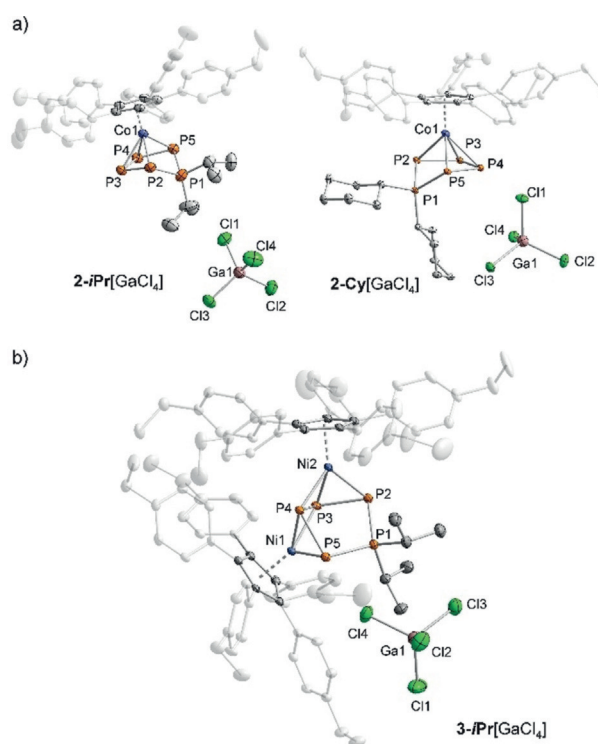


Figure 3. Solid-state molecular structures of complexes **2-R** $[GaCl_4]$ ($R = iPr$ and Cy) (a) and **3-*iPr*** $[GaCl_4]$ (b). Ellipsoids set at 50% probability; H atoms, solvent molecules in **2-Cy** $[GaCl_4]$, and disorder in the ethyl groups and the tetrachlorogallate anion of **2-*iPr*** $[GaCl_4]$ are omitted for clarity. The crystal of **2-*iPr*** $[GaCl_4]$ contained a second crystallographically independent molecule with very similar structural parameters; only one of these molecules is shown. Selected bond lengths [Å] and angles [°] for **2-*iPr*** $[GaCl_4]$: P1–P2 2.1543(12), 2.1592(12); P2–P3 2.1382(12), 2.1362(12); P3–P4 2.1371(12), 2.1370(12); P4–P5 2.1267(12), 2.1377(12); P1–P5 2.1449(12), 2.1558(11); Co1–P2 2.3227(9), 2.3316(9); Co1–P3 2.3219(10), 2.3249(9); Co1–P4 2.3396(9), 2.3368(9); Co1–P5 2.3269(9), 2.3225(10); P2–P1–P5 97.09(5), 97.15(4), P1–P2–P3 99.87(5), 101.08(5); P2–P3–P4 105.21(5), 105.36(5); P3–P4–P5 104.26(5), 104.43(5); P4–P5–P1 100.91(5), 101.50(4); **2-Cy** $[GaCl_4]$: P1–P2 2.1633(6), P2–P3 2.1361(7), P3–P4 2.1414(7), P4–P5 2.1327(7), P1–P5 2.1658(6), Co1–P2 2.3294(5), Co1–P3 2.3213(6), Co1–P4 2.3443(6), Co1–P5 2.3376(5); P2–P1–P5 96.58(2), P1–P2–P3 101.01(3), P2–P3–P4 105.12(3), P3–P4–P5 104.48(3), P4–P5–P1 101.27(3); **3-*iPr*** $[GaCl_4]$: P1–P2 2.1835(17), P2–P3 2.1672(16), P3–P4 2.5002(16), P4–P5 2.1542(16), P1–P5 2.1859(15), Ni1–P3 2.2388(12), Ni1–P4 2.2519(13), Ni1–P5 2.2547(15), Ni2–P2 2.2473(13), Ni2–P3 2.2671(13), Ni2–P4 2.2202(12); P2–P1–P5 108.73(6), P1–P2–P3 97.38(6), P2–P3–P4 96.11(6), P3–P4–P5 92.73(6), P4–P5–P1 100.25(6).

a persistent, symmetrical P_5R_2 ligand. The spectral parameters were determined by an iterative fitting procedure in each case (see the Supporting Information for details). For **2-R** $[GaCl_4]$, the data are similar to those of related α -diimine complexes (**IV** in Figure 1). $^{31}P\{^1H\}$ NMR spectra of **3-R** $[GaCl_4]$ display three multiplets of an AA'MXX' spin system. The very small magnitude of the $^1J_{AA'}$ coupling constants in **3-R** $[GaCl_4]$ ($^1J_{AA'} = -7.5$ Hz for **3-*iPr*** $[GaCl_4]$) and $^1J_{AA'} = -18.7$ Hz for **3-Cy** $[GaCl_4]$) is noteworthy. This observation agrees well with a weak P–P bond in the β -position with respect to the phosphonium center (see the discussion of the X-ray structure above), and the remaining $^1J_{PP}$ couplings

are unexceptional; for **3-*i*Pr**[GaCl₄]: $^1J_{MX} = ^1J_{MX'} = -420.4$ Hz and $^1J_{AX} = ^1J_{AX'} = -346.2$ Hz. The fluxional behavior of the Ni complexes is another notable aspect, which was observed by variable temperature (VT) $^{31}\text{P}\{^1\text{H}\}$ NMR studies on **3-*R***-[GaCl₄] in [D₈]THF (Figure 4). A symmetrical AA'MXX'

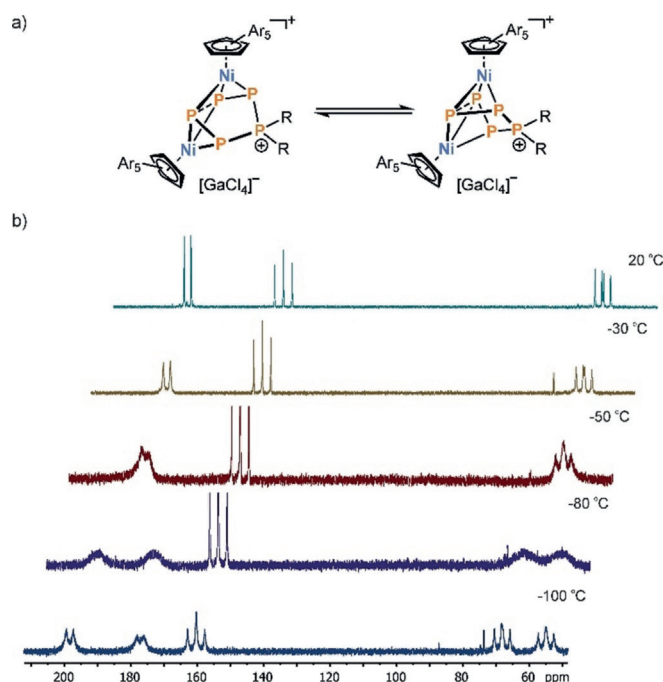
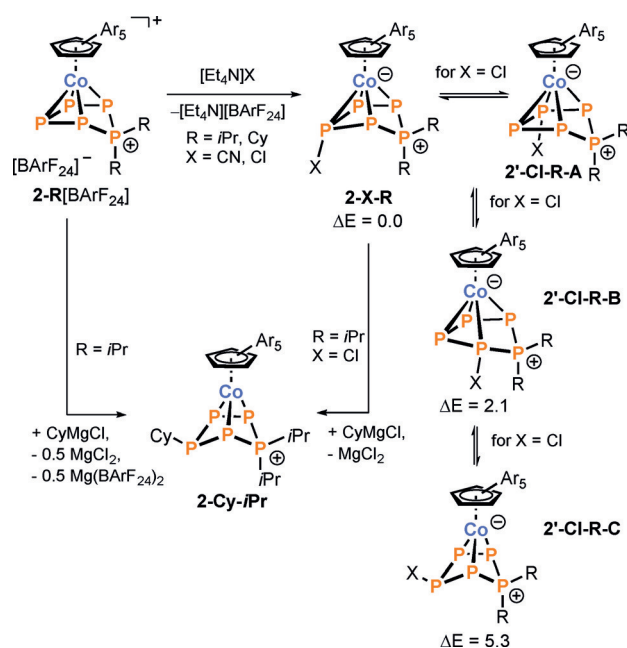


Figure 4. a) Dynamic behavior of **3-*R***[GaCl₄] at room temperature; Ar = C₆H₄-4-Et. b) VT- $^{31}\text{P}\{^1\text{H}\}$ NMR spectra of complex **3-*i*Pr**[GaCl₄] in [D₈]THF.

spin system is observed at room temperature, which decoalesces into an AEMXY spin system below -80°C (see Figure 4b and Figure S88). The prevalence of an intramolecular exchange process as shown in Figure 4a was additionally confirmed by a ^{31}P - ^{31}P EXSY experiment on **3-*Cy***[GaCl₄] in [D₈]THF at -80°C , which shows correlation peaks at $\delta = 204$ and 179 ppm (Figure S89). The NMR spectroscopic data of **2-*R***[GaCl₄] and **3-*R***[GaCl₄] thus clearly indicate that the coordination mode of the *cyclo*-P₅R₂ ligand observed in the solid-state structures is retained in solution.

Aiming at the further functionalization of the P₅R₂ ligands, reactions of [**2-*R***]⁺ and [**3-*R***]⁺ with nucleophilic anions were examined. In order to facilitate such studies, the [GaCl₄]⁻ counteranion was first replaced by reacting **2-*R***[GaCl₄] and **3-*R***[GaCl₄] with Na(BArF₂₄) (Scheme 2). The desired salts **2-*R***[BArF₂₄] and **3-*R***[BArF₂₄] can be isolated as pure compounds in moderate to high yields (47–79%); their cations show nearly identical structural and spectroscopic data as the tetrachlorogallate salts (see the Supporting Information for details). Initial reactivity tests revealed that chloro and cyano substituents are readily incorporated into the cobalt complexes, while the dinickel species **3-*R***[BArF₂₄] are unreactive. Treatment of **2-*R***[BArF₂₄] with [Et₄N]X (1 equiv, X = CN and Cl) as shown in Scheme 3 selectively gives [Cp^{Ar}Co{η⁴-P₅R₂(X)}] (**2-*X-*R****,



Scheme 3. Functionalization of the *cyclo*-P₅R₂ backbone of **2-*R***[BArF₂₄]; Ar = C₆H₄-4-Et. Relative electronic energies (ΔE in kcal mol⁻¹) are given for the calculated isomers **B** and **C** of the truncated model complex [Cp^{Ph}Co{η⁴-P₅iPr₂(X)}] **2'-*X-*i*Pr-A*** (see Table S17 for thermal enthalpies and free enthalpies).

X = CN and Cl, R = *i*Pr and Cy; Figure 5a) according to ^{31}P NMR spectroscopy. The products can be isolated as pure, dark-red, crystalline solids in 65–75% yield. Single-crystal X-ray diffraction analysis on red plates obtained from Et₂O/*n*-pentane show that the (pseudo)halogen substituent is connected to the β-position with respect to the phosphonium center of the pentaphosphorus ligand in all cases. The molecular structures of **2-CN-*i*Pr** and **2-Cl-*i*Pr** are shown in Figure 5a. The Co–P bond of the cyano- or chloro-substituted phosphorus atom (**2-CN-*i*Pr**: Co1–P3 2.8558(13) Å; **2-Cl-*i*Pr**: Co1–P3 2.7301(17) Å) is elongated by 0.4 Å to 0.5 Å compared to the remaining Co–P bonds. As a result, the substituted P atoms P1 and P3 are below the P2–P4–P5 plane. The dihedral angle P2–P3–P4–P5 is 23.20(8)° in **2-CN-*i*Pr** and 16.2(1)° in **2-Cl-*i*Pr**. For **2-CN-*i*Pr**, the P3–C46 and C46–N1 bond lengths of 1.831(4) Å and 1.141(5) Å indicate the P–C single and C≡N triple bonds.^[32] The P4–Cl1 bond (2.205(2) Å) in **2-Cl-*i*Pr** is 0.16 Å longer than in typical alkyl or aryl chlorophosphanes.^[22c] The P–P bond lengths in the *cyclo*-P₅R₂ ring are in a close range (2.1502(15)–2.1804(17) Å for **2-CN-*i*Pr** and 2.142(2)–2.1691(18) Å for **2-Cl-*i*Pr**), showing that the P₅ ring bonds are intact.

The $^{31}\text{P}\{^1\text{H}\}$ NMR spectrum of **2-CN-*i*Pr** in C₆D₆ shows five multiplets of an AEMXY spin system at room temperature, which is in agreement with the unsymmetrical solid-state structure (see Figures S62 and S63). The $^1J_{PP}$ coupling constants obtained by iterative fitting (-324.7 Hz to -400.2 Hz) are typical for P–P single bonds, while the $^2J_{PP}$ coupling constants ($^2J_{AE} = -3.9$ Hz, $^2J_{AM} = -7.7$ Hz, $^2J_{MY} = 80.3$ Hz, $^2J_{EX} = 50.0$ Hz, $^2J_{XY} = -6.4$ Hz) span a large range. Small $^2J_{PP}$ couplings such as $^2J_{AE}$ and $^2J_{AM}$ are typical for tetracoordinate P atoms.^[19–23]

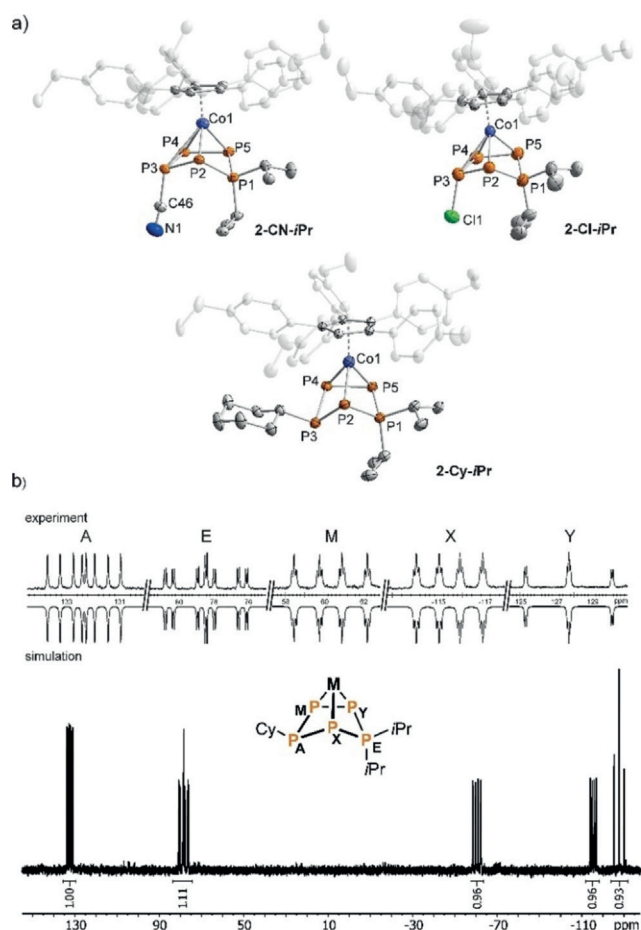


Figure 5. a) Solid-state molecular structures of **2-CN-iPr** (top left), **2-Cl-iPr** (top right), and **2-Cy-iPr** (bottom). Ellipsoids set at 50% (**2-CN-iPr** and **2-Cy-iPr**) and 30% (**2-Cl-iPr**) probability; H atoms, solvent molecules (Et₂O and *n*-pentane in **2-CN-iPr**), and disorder of the ethyl groups of Cp^{Ar} (in **2-CN-iPr** and **2-Cy-iPr**) and *cyclo*-P₃CNiPr₂ (in **2-CN-iPr**) are omitted for clarity. Selected bond lengths [Å] and angles [°] for **2-CN-iPr**: P1–P2 2.1615(9), P2–P3 2.1730(13), P3–P4 2.1804(17), P4–P5 2.1502(15), P1–P5 2.1541(10), Co1–P2 2.3173(7), Co1–P3 2.8558(13), Co1–P4 2.3072(13), Co1–P5 2.3031(7), P3–C46 1.831(4), C46–N1 1.141(5); P2–P1–P5 97.31(4), P1–P2–P3 99.68(4), P2–P3–P4 96.15(6), P3–P4–P5 109.36(6), P4–P5–P1 100.91(5), P2–P3–C46 110.87(14), P4–P3–C46 109.29(13), P3–C46–N1 173.3(4); for **2-Cl-iPr**: P1–P2 2.1691(18), P2–P3 2.142(2), P3–P4 2.1510(19), P4–P5 2.150(2), P1–P5 2.1458(18), Co1–P2 2.3227(14), Co1–P3 2.7301(17), Co1–P4 2.3214(16), Co1–P5 2.2987(15), P4–Cl1 2.205(2); P2–P1–P5 97.11(7), P1–P2–P3 100.62(8), P2–P3–P4 98.77(8), P3–P4–P5 109.08(8), P4–P5–P1 100.51(7), P2–P3–Cl1 110.07(9), P4–P3–Cl1 110.93(9); for **2-Cy-iPr**: P1–P2 2.1506(14), P2–P3 2.2327(16), P3–P4 2.2185(15), P4–P5 2.1493(15), P1–P5 2.1697(15), Co1–P2 2.2831(12), Co1–P4 2.2648(11), Co1–P5 2.3152(11); P2–P1–P5 95.55(5), P1–P2–P3 87.95(6), P2–P3–P4 86.30(5), P3–P4–P5 102.87(6), P4–P5–P1 99.77(6). b) ³¹P{¹H} NMR spectrum of complex **2-Cy-iPr** in C₆D₆ at room temperature; δ(P_A) = 132.3 ppm, δ(P_E) = 78.4 ppm, δ(P_M) = –60.4 ppm, δ(P_X) = –115.5, and δ(P_Y) = –127.7 ppm; ¹J_{AM} = –211.2 Hz, ¹J_{MY} = –396.4 Hz, ¹J_{EY} = –382.8 Hz, ¹J_{EX} = –302.7 Hz, ¹J_{AX} = –158.5 Hz, ²J_{AY} = –2.6 Hz, ²J_{AE} = 76.7 Hz, ²J_{EM} = 19.4 Hz, ²J_{MX} = 19.5 Hz, and ²J_{XY} = 16.3 Hz.

Unlike **2-CN-R**, the ³¹P{¹H} NMR spectrum of **2-Cl-R** shows an AMM'XX' spin system with broadened MM' and XX' multiplets at room temperature. Variable-temperature ³¹P{¹H} NMR measurements show that the signals decoalesce

upon cooling, and an AEMXY spin system appears below –50 °C in [D₈]THF (Figures S92 and S93). This observation is postulated to be due to an exchange of the chlorine atoms between the P3 and P4 atoms of the *cyclo*-P₅R₂ ring (Scheme 3). Indeed, DFT calculations on the truncated model complex [Cp^{Ph}Co{η⁴-P₅iPr₂(X)}] **2'-X-iPr** (X = Cl and Cy; Cp^{Ph} = C₅(C₆H₅)₅, PBE/def2-TZVP level, see the Supporting Information for further details) gave several low-energy isomers, which likely form an equilibrium in solution (see Scheme 3). Isomer **2'-Cl-iPr-A**, which corresponds to the crystallographically observed species, has the lowest energy compared to the *exo*-isomer **2'-Cl-iPr-C** (+5.3 kcal mol⁻¹) and the P2-substituted derivative **2'-Cl-iPr-B** (+2.1 kcal mol⁻¹; see Scheme 3 and the Supporting Information for details). The experimental ³¹P{¹H} NMR spectrum of **2-Cl-iPr** (Figure S92) shows a second isomer (AEMXY spin system) below –70 °C, which could correspond to **2-Cl-iPr-B** or **2-Cl-iPr-C**.^[33]

Initial attempts toward the organofunctionalization of **2-R**[BARF₂₄] with CyMgCl led to a product mixture (Figure S83) with [Cp^{Ar}Co{η³-P₅iPr₂(Cy)}] (**2-Cy-iPr**) as the major species, showing an AEMXY spin system. **2-Cy-iPr** is formed cleanly in the reaction of **2-Cl-iPr** with CyMgCl (Scheme 3), from which it can be isolated as a dark-red, crystalline solid in 68% yield. The molecular structure determined by single-crystal X-ray diffraction is shown in Figure 5a. Different from **2-X-iPr** (X = CN and Cl, also shown in Figure 5a), an *exo* configuration of the Cy substituent is observed, that is, the cyclohexyl substituent points towards the cyclopentadienyl ring. The pentaphosphorus ligand is clearly η³-coordinated (Co1–P2 2.2831(12) Å, Co1–P4 2.2648(11) Å, and Co1–P5 2.3152(11) Å) and shows a boat conformation as a result. The P–P bonds of the cyclohexyl-substituted P atom P3 (P2–P3 2.2327(16) Å, P3–P4 2.2185(15) Å) are elongated by 0.04 Å to 0.09 Å compared to the chloro- and cyano-substituted P atoms in **2-X-iPr** (X = CN and Cl). The AEMXY spin system of **2-Cy-iPr** was successfully simulated by iterative fitting (Figure 5b) and compared to **2-CN-R**, revealing significant differences concerning the ¹J_{PP} and ²J_{PP} coupling constants. For example, the magnitudes of ¹J_{AM} = –211.2 Hz and ¹J_{AX} = –158.5 Hz are much smaller than in the cyano-substituted compounds (see above). Such changes likely arise from the different conformations of the P₅ ring. DFT calculations on the model complex **2'-Cy-iPr**, where the ethyl groups on the cyclopentadienyl ligand were replaced by hydrogen atoms, suggest that the *exo* isomer is 8.6 kcal mol⁻¹ more stable than the *endo*-configured product. DFT calculations of NMR chemical shielding (Table S21) for the *exo* isomer suggest the assignment shown in Figure 5b; the calculated chemical shifts are in fairly good agreement with the experimentally observed values.

It seems important to note that reactions of [P₅Mes₂]-[BARF₂₄] with [Et₄N]X (X = Cl and CN) and CyMgCl failed to give functionalization products (see the Supporting Information for details). This observation illustrates that the transition-metal coordination is of key importance here to achieve the functionalization of the P₅R₂ scaffold.

In summary, we have comprehensively studied the hitherto unknown coordination behavior of cationic poly-

phosphorus compounds of type $P_5R_2^+$. These studies have resulted in unusual cationic complexes **2-R**[GaCl₄] and **3-R**[GaCl₄] (R = *i*Pr and Cy), while the first neutral bicyclo-[1.1.0]pentaphosphanes (**1-R**) were obtained by treatment of [P₅R₂][GaCl₄] (R = *i*Pr and Mes) with [Cp^{Ar}Fe(μ-Br)]₂, or, more conveniently, with K[Cp^{Ar}]. The functionalization of the polyphosphorus ligands in the [**2-R**]⁺ cations is straightforward after anion exchange of [GaCl₄]⁻ for [BArF₂₄]⁻, and the resulting mixed substituted complexes **2-X-R** (X = CN and Cl) can be further modified as demonstrated by the synthesis of **2-Cy-iPr** from **2-Cl-iPr**. The „functionalization by complexation“ approach thus potentially enables the preparation of hitherto inaccessible and multiply substituted polyphosphorus scaffolds in the coordination sphere of transition-metal cations. This strategy is complementary to previously reported methods for the synthesis of functionalized polyphosphorus complexes,^[9,10c,11,19] and further extensions should give access to a broad range of novel P_n frameworks.

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Conflict of interest

The authors declare no conflict of interest.

Stichwörter: Cobalt · Koordinationsverbindungen · Nickel · P-Liganden · Phosphor

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