

Scheme 3. Synthesis of $[\text{Mes}^*\text{P}_4(\text{Cl})\text{Mes}^*][\text{GaCl}_4]$ ($3[\text{GaCl}_4]$). The isomeric ratio between 3a^+ and 3b^+ varies from 1:8 (-50°C , 3 h) to 3:4 (-80°C , 12 h) depending on the reaction conditions.

reaction (ca. 40 % by mole fraction). Full conversion of **1** to **2**⁺ could be confirmed by a trapping reaction (see below).

The resonances were assigned on the basis of computed NMR data (Supporting Information). The calculated gas-phase structure of **2**⁺ (PBE0/6-31G(d,p)) reveals *C*₁ symmetry and incorporates a P=P double bond (2.05 Å; Wiberg bond index: 1.49; cf. 2.034(2) Å in $\text{Mes}^*\text{P}=\text{PMes}^*$)^[35] that stabilizes the positive charge. In agreement with the Lewis formula, the highest partial charge is therefore found at one of the Mes^* -substituted P atoms (+0.45 *e*), which adopts a nearly planar coordination environment ($\Sigma(\angle \text{P}) = 359.9^\circ$). Even though the minimum structure displays four inequivalent P nuclei, a relatively fast exchange between two enantiomers accounts for both the observed NMR pattern and the signal broadening (Figure 1). As the dynamic equilibrium renders both Mes^* -

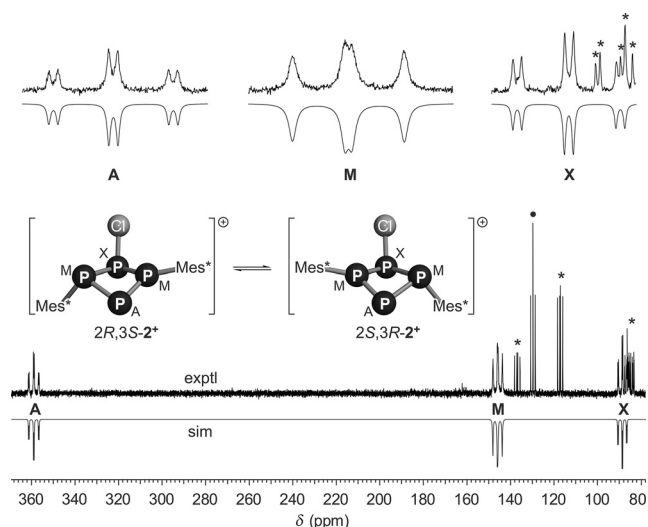


Figure 1. Experimental and simulated in situ ^{31}P NMR spectrum of the reaction of **1** and GaCl_3 (-60°C). The AM_2X pattern is caused by a dynamic equilibrium between two enantiomers of **2**⁺ (calculated structures). Labeled lines/signals are caused by starting material (●) and other intermediates (*).

substituted P atoms equivalent, a single resonance at the center of their individual shifts ($\delta_{\text{calc}} = 206.5, 48.6$ ppm; $\delta_{\text{av}} = 127.6$ ppm) is expected. Taking this into consideration, the calculated and experimental data show reasonable agreement (Supporting Information, Table S7). Two other intermediates were observed at temperatures below -20°C , which could not

be identified unambiguously (Supporting Information, Figure S1).

The color of the reaction mixture faded slowly to orange/yellow over a period of one day at low temperatures. After full conversion of the starting material, two main products were formed, which were identified as *exo-exo* and *endo-exo* isomers of $[\text{Mes}^*\text{P}_4(\text{Cl})\text{Mes}^*]^+$ (**3a**⁺, **3b**⁺, Scheme 3) on the basis of NMR data (85 % yield). Intriguingly, the chlorine atom of the reaction product is connected to one of the Mes^* -substituted P atoms, resulting in the formation of a novel, bicyclic triphosphinophosphonium framework. According to DFT calculations, these constitutional isomers of **2**⁺, which are formed after a formal 1,2-Cl shift, are energetically favored by 46.3 kJ mol⁻¹ (**3a**⁺) or 45.3 kJ mol⁻¹ (**3b**⁺) with respect to intermediate **2**⁺ (ΔG^{298}).

Addition of *n*-pentane facilitated crystallization at -80°C and afforded colorless crystals that were characterized as $\text{CH}_2\text{Cl}_2/\text{C}_6\text{H}_5\text{F}$ solvate of **3a** $[\text{GaCl}_4]$ (40 % yield of isolated product). The molecular structure was determined by single-crystal X-ray diffraction (Figure 2). All P–P bond lengths are

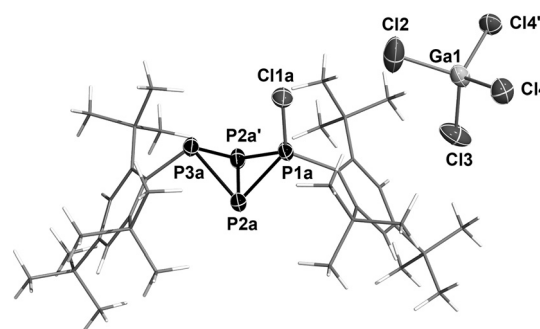


Figure 2. Molecular structure of **3a** $[\text{GaCl}_4]$. Ellipsoids are set at 50 % probability (173 K). Selected bond lengths [Å] and angles [$^\circ$]: P1a–P2a 2.150(2), P1a–Cl1a 2.009(2), P2a–P2a' 2.244(2), P2a–P3a 2.244(2); Cl1a–P1a–P2a 117.65(7), Cl1–P1a–Cl1a 118.7(2), P2a'–P1a–P2a 62.91(7), P1a–P2a–P2a' 58.55(3), P3a–P2a–P2a' 59.99(3), P1a–P2a–P3a 83.60(6), P2a'–P3a–P2a 60.02(7); P1a–P2a–P2a'–P3a $-101.68(3)^\circ$.^[56]

within the range of typical single bonds (cf. $\Sigma r_{\text{cov}} = 2.22$ Å),^[36] although the bonds to P1a (2.150(2) Å) are somewhat shortened owing to bond polarization. Similarly, the P1a–Cl1a bond (2.009(2) Å) is slightly shorter than the sum of covalent radii (2.10 Å).^[36] A distorted tetrahedral coordination environment is found for P1a, which can be compared to tetracoordinate P atoms in cyclic phosphinohalophosphonium ions or bicyclic $\text{TerP}_4\text{Me} \cdot \text{B}(\text{C}_6\text{F}_5)_3$ ($\text{Ter} = 2,6$ -dimesityl-phenyl).^[18,32] All other P atoms exhibit typical trigonal pyramidal coordination (bond angles 58° – 96°). The fold angle of the butterfly-shaped bicyclic P_4 scaffold amounts to $101.68(3)^\circ$, which compares well to known tetraphosphabicyclo[1.1.0]butane frameworks (88.6° – 108.1°).^[25,26,32,37–41] No close contacts are found between anion and cation. Several attempts to crystallize the *endo-exo* isomer **3b** $[\text{GaCl}_4]$, which is actually formed in excess according to ^{31}P NMR spectra, remained futile.

The solid-state Raman spectrum of **3a** $[\text{GaCl}_4]$ (at -60°C) shows characteristic bands at 340 cm^{-1} (GaCl_4 “breathing”

mode), 485 cm⁻¹ (P₄ deformation), 513 cm⁻¹ (P₄Cl “breathing”), 588 cm⁻¹ (P–C stretching at tricoordinate P), 615 cm⁻¹ (P–Cl stretching), and 633 cm⁻¹ (P–C stretching at tetracoordinate P). Upon warming, the sample started to decompose above 0 °C as indicated by decreasing signal intensities (Supporting Information, Figure S2).

To further investigate the electronic structure of **3a**⁺ and **3b**⁺, density functional theory (DFT) calculations were performed.^[42] For both isomers, NBO^[43] analysis revealed that the positive charge is mostly localized at the phosphonium center (P1, +0.76 *e*), but also resides at the tricoordinate Mes* substituted atom (P3, +0.37 *e*). The bridgehead atoms (P2) are only slightly positive (+0.12 *e*), whereas the Cl atom bears a slightly negative charge (−0.16 *e*). Thus, the overall charge of the P₄Cl scaffold amounts to +1.21 *e*, or +1.37 *e* if only the P₄ unit is regarded. Comparison with neutral Mes*P₄Mes* (+0.54 *e*, Supporting Information) or even dicationic [(Ph₃As)P₄(AsPh₃)]²⁺ (+0.12 *e*)^[26] confirms the unique charge distribution within both isomers of **3**⁺.

In the ³¹P NMR spectrum, **3a**⁺ and **3b**⁺ are characterized by AMX₂ patterns (Figure 3). A characteristic high

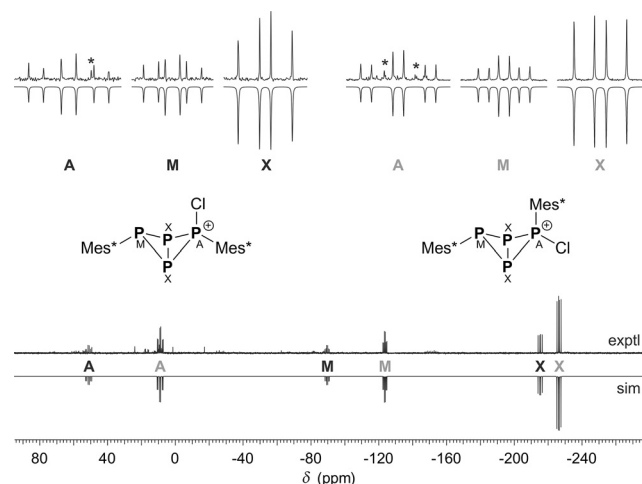


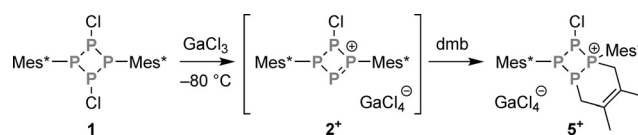
Figure 3. In situ ³¹P NMR spectrum at 0 °C displaying the formation of **3a**⁺ and **3b**⁺. Starred lines in the enlarged section are caused by impurities that are due to incipient decomposition at this temperature.

field shift is observed for the bridgehead atoms (P_X; **3a**⁺: −214.5 ppm; **3b**⁺: −226.4 ppm), which compares to known tetraphosphabicyclo[1.1.0]butanes (−220 to −370 ppm),^[25,26,32,37–41,44–46] but is still shifted slightly downfield owing to the positive charge of the P₄ scaffold. The tricoordinate bridging atom (P_M) displays a moderate high-field shift (**3a**⁺: −89.5 ppm; **3b**⁺: −123.6 ppm) within the typical range, whereas the tetracoordinate P atom (P_A) is shifted significantly downfield (**3a**⁺: +51.2 ppm; **3b**⁺: +9.0 ppm), which has not yet been observed for nuclei within bicyclic P₄ frameworks. All of the experimental values are in good agreement with theoretical data (Supporting Information, Tables S8 and S9).

At ambient temperature, solution NMR spectra indicate that **3a**⁺ and **3b**⁺ decompose within hours.^[47] One of the decomposition products was identified as

[Mes*P(H)(Cl)*t*Bu][GaCl₄] (**4**[GaCl₄]; Supporting Information). Remarkably, a *t*Bu group is transferred from a Mes* substituent to the P atom of **4**⁺ in the course of this reaction, which demonstrates the intrinsic lability of **3a**⁺ and **3b**⁺. Displacement of *t*Bu groups from Mes* substituents has previously been observed in the presence of Lewis acids or under thermal conditions.^[6,48–51]

In a next series of experiments, we aimed to further establish the identity of low-temperature intermediate **2**⁺. For that purpose, **1** was treated with GaCl₃ in the presence of dimethylbutadiene (dmb) as trapping reagent.^[52,53] This time, no red color was observed upon addition of GaCl₃. According to ³¹P NMR data, the reaction led to quantitative formation of tetraphosphabicyclo[4.2.0]octenium salt **5**[GaCl₄] (Scheme 4), which can be regarded as [4+2] cycloaddition



Scheme 4. Synthesis of [Mes*P₄(Cl)(C₆H₁₀)Mes*][GaCl₄] (**5**[GaCl₄]).

product of intermediate **2**⁺ and dmb. In this vein, the trapping reaction also proved quantitative conversion of **1** to **2**⁺ at low temperatures.

At ambient temperature, cation **5**⁺ is characterized by four broad signals in the ³¹P NMR spectrum, indicating unresolved dynamic effects. Cooling to −80 °C revealed dynamic equilibria between several conformers or rotamers (Supporting Information, Figure S3). Heating to +80 °C barely improved the resolution of the NMR spectrum and led to slow decomposition of **5**⁺. In the ¹H NMR spectrum, some of the signals are also significantly broadened, which is most likely due to hindered rotation of some *t*Bu groups.

Crystallization at 5 °C yielded colorless crystals of **5**·[GaCl₄]·CH₂Cl₂ (yield of isolated product after workup: 45 %; see the Supporting Information). The solid-state structure was determined by single-crystal X-ray diffraction (Figure 4).

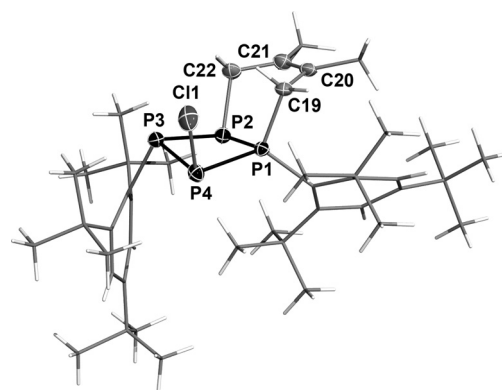


Figure 4. Molecular structure of **5**⁺.^[56] Ellipsoids are set at 50% probability (173 K). Selected bond lengths [Å] and angles [°]: P1–P2 2.195(1), P1–C19 1.846(3), C19–C20 1.514(4), C20–C21 1.334(4), C21–C22 1.500(4), P2–C22 1.871(3), P4–Cl1 2.068(1); P2–P1–P4 91.78(4), C19–P1–P2 102.3(1), C19–P1–P4 116.1(1); P1–P2–P4–P3 159.03(1).

The bicyclic system is folded along the P1–P2 and C19–C22 axes (angles between least squares planes: P3–P4–P1–P2/P1–P2–C22–C19/C22–C19–C20–C21 72.1°/63.2°). The P₄ ring system is rather flat (fold angle: 159°), but well within typical parameters (110°–173°).^[54,55] P1 sits in a distorted tetrahedral environment, and the other P atoms exhibit trigonal pyramidal coordination. The P1–P2 bond (2.195(1) Å) is slightly shorter than the other three P–P bonds (av. 2.25 Å). The structural data of the P₄ ring system correspond to other known cyclotriphosphinophosphonium ions.^[16–18] The Mes* moiety at P1 is highly distorted owing to Pauli repulsion between the *o*-tBu groups and the P₄C₆H₁₀ scaffold; the P atom is bent out of the least-squares plane of the phenyl ring by 52.1°. No close contacts are found between anion and cation. As expected, the positive charge is essentially localized at P1 (+0.92 *e*) according to NBO analysis. The partial charges of the other P atoms are only slightly positive (av. +0.32 *e*). To exclude the possibility that **5**⁺ was formed by a “reverse” reaction of **3a**⁺ or **3b**⁺ with dmb, a solution of both salts was treated with the latter. This did not lead to the formation of **5**⁺; in fact, no reaction was observed after two hours based on ³¹P NMR data.

In conclusion, the synthesis of the first tetraphosphabicyclo[1.1.0]butan-2-ium salt, **3**[GaCl₄], which incorporates a positively charged, bicyclic P₄ scaffold, is achieved by selective chlorine abstraction starting from [CIP(μ-PMes*)]₂ (**1**). **3**[GaCl₄] is temperature-sensitive and only stable at low temperatures. The formation of **3**⁺ was studied by means of low-temperature NMR experiments. This led to the discovery of tetraphosphonium intermediate **2**⁺, which could be identified on the basis of DFT calculations. Moreover, a trapping reaction with dmb could provide chemical evidence for the identity of **2**⁺. Currently, we are investigating the reactivity of **1** with other Lewis acids and under different conditions to diversify the product scope.

Keywords: bicyclic compounds · NMR spectroscopy · phosphorus cations · phosphorus compounds · ring systems

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