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# Formation of the spirocyclic, Si-centered cage cations $[ClP(NSiMe_3)_2Si(NSiMe_3)_2P_5]^+$ and $[P_5(NSiMe_3)_2Si(NSiMe_3)_2P_5]^{2+}$ ;

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On account of our interest in  $P_4$  activation by phosphenium ion insertion into P-P bonds we have developed synthetic routes to bicyclic N-P-Si-heterocycle **7** and probed its reactivity towards  $GaCl_3$  and  $P_4$ . A  $GaCl_3$ -induced rearrangement of **7** leads to the *in situ* formation of spirocyclic, Si-centered phosphenium ions. Their insertion into P-P bonds of one or two  $P_4$  tetrahedra yields polyphosphorus cages  $[CIP(NSiMe_3)_2Si(NSiMe_3)_2P_5]^+$  (**19**<sup>+</sup>) and  $[P_5(NSiMe_3)_2Si(NSiMe_3)_2P_5]^{2+}$  (**13**<sup>2+</sup>).

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Introduction

White phosphorus  $(P_4)$  is an archetypal building block for the syntheses of polyphosphorus cations featuring a high P to substituent ratio. In contrast to highly substituted cations  $R_n P_m$ (n > m), which are obtained via synthetic methods based on catena or cyclic polyphosphanes,2 the targeted preparation of P-rich cations  $R_n P_m$  (n < m) is achieved by taking advantage of the tetrahedral P<sub>4</sub> framework. 1,3 In a seminal paper, Krossing et al. reported that dicoordinated phosphenium ions, like other predominantly electrophilic ambiphiles,4 are able to insert into a P-P bond of the P4 tetrahedron.5 This was exploited for the preparation of a series of  $P_5X_2^+$ -cages (X = Cl, Br, I). 5,6 We expanded this methodology and prepared a series of symmetrically and unsymmetrically-substituted R<sub>2</sub>P<sub>5</sub><sup>+</sup>- and  $RP_5Cl^+$ -cations (R = aryl, alkyl,  $R_2N$ ).<sup>7</sup> The additional stability added by organo-substituents allowed for the stepwise insertion of up to three [Ph<sub>2</sub>P]<sup>+</sup> phosphenium ions into P-P bonds of one P<sub>4</sub> molecule yielding mono- to tri-cationic [Ph<sub>2</sub>P<sub>5</sub>]+-, [Ph<sub>4</sub>P<sub>6</sub>]<sup>2+</sup>- and [Ph<sub>6</sub>P<sub>7</sub>]<sup>3+</sup>-cages.<sup>8</sup> A complementary study exploited 1,3-dichloro-cyclo-1,3-diphosphadiazane ClP(NDipp)2-PCl (Dipp = 2,6-diisopropylphenyl), as a twofold phosphenium ion source for the stepwise activation of two P4 tetrahedra which yielded the mono- and dicationic species [CIP- $(NDipp)_2P_5|^+$  (1<sup>+</sup>) and  $[P_5(NDipp)_2P_5]^{2+}$  (2<sup>2+</sup>, Fig. 1).<sup>9</sup> The related P<sub>5</sub><sup>+</sup>-species 3[GaCl<sub>4</sub>] was obtained by the reaction of P<sub>4</sub> with the four-membered heterocycle Cl2Si(NSiMe3)2PCl and

 $(R = SiMe_3 E = AI, n = 2)$ 

Fig. 1  $P_5^+$ -cages 1<sup>+</sup>, 2<sup>2+</sup>, 3<sup>+</sup> and 4 featuring a fused four-membered heterocycle.

GaCl<sub>3</sub> while the isolobale Al-species 4 is the result of the reaction of phosphenium zwitterion Cl<sub>2</sub>Al(NSiMe<sub>3</sub>)<sub>2</sub>P with P<sub>4</sub>. <sup>10</sup>

Only recently, cationic polyphosphorus cages have emerged as valuable synthetic building blocks for three purposes. First, they can be selectively fragmented in reactions with carbenes which results in the formation of  $P_n$ -species (e.g. a  $P_5$ <sup>+</sup>-cage cation yields a  $P_2$ - and cationic  $P_3$ <sup>+</sup>-species). Second, they can be oxidized with selenium or sulphur which allows for the targeted preparation of cationic phosphorus-chalcogen cages. Third, they can be used for the controlled release of  $P_4$  due to the reversibility of the phosphenium ion insertion.

With the intention to further expand the range of methods for the *in situ* generation of phosphenium ions for  $P_4$  activation, we investigated the reactivity of bicyclic phosphorus-nitrogen–silicon heterocycle 7 with  $GaCl_3$  and  $P_4$ . A  $GaCl_3$ -induced rearrangement reaction was observed which formally gives access to the spirocyclic, Si-centered compound  $ClP-(NSiMe_3)_2Si(NSiMe_3)_2PCl$ . Chloride abstraction by  $GaCl_3$  and insertion of the respective phosphenium ion in P-P bonds of one or two  $P_4$  molecules yields the Si-centered polyphosphorus cages  $[ClP(NSiMe_3)_2Si(NSiMe_3)_2P_5]^+$   $(19^+)$  and  $[P_5(NSiMe_3)_2Si(NSiMe_3)_2P_5]^+$   $(13^{2+})$ .

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<sup>†</sup> Electronic supplementary information (ESI) available: Crystallographic data, <sup>31</sup>P NMR spectra of reaction mixtures and general experimental information. CCDC 1060461–1060464. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c5dt01512j

## Results and discussion

**Paper** 

### Synthetic routes towards and characterization of 7

As part of our ongoing interest in four-membered phosphorus–nitrogen-element heterocycles as precursors for phosphenium ions <sup>12</sup> that can insert into a P–P bond of P<sub>4</sub> <sup>10a</sup> we revisited the synthesis of heterocycle **6**. This compound is synthesized by the reaction of iminophosphane 5 with SiCl<sub>4</sub> according to a procedure reported by Niecke and co-workers (Scheme 1). <sup>13</sup> Compound **6** was obtained from the reaction mixture by distillation (40 °C,  $8 \times 10^{-2}$  mbar) and isolated in 41% yield in accordance with the literature report. Surprisingly, however, a second fraction was obtained at higher temperatures (105 °C,  $2 \times 10^{-3}$  mbar) and identified as bicyclic compound 7 (18% yield). <sup>14</sup> The <sup>31</sup>P{<sup>1</sup>H} NMR spectrum of 7 dissolved in C<sub>6</sub>D<sub>6</sub> shows a singlet resonance at  $\delta$ (P) = 211.8 ppm which is in the typical chemical shift region of diphosphadiazanes featuring amino-groups on P. <sup>15</sup>

The molecular structure of 7 is  $C_{2V}$ -symmetric and features an distorted planar four-membered [NP]2-ring (largest deviation from the plane 0.195 Å, Fig. 2). All four N atoms exhibit a trigonal planar arrangement (angular sums range from 358.8 (3)° to 360.0(3)°) whereas the P atoms are involved in pyramidal bonding environments (angular sums P1: 286(3)° and P2: 286.2(2)°). The P1-N1 and P2-N2 bonds are almost orthogonal to the [NP]<sub>2</sub>-plane (N1-P1···P2: 95.84(4)°, N2-P2···P1: 95.84 (4)°). The  $C_{2V}$ -symmetric arrangement of the N<sub>2</sub>SiCl<sub>2</sub>-moiety in 7 is rare<sup>16</sup> and contrasts other known diamido-cyclodiphosphazane compounds compounds coordinating main group fragments. 15 Compounds of type (PNR)2NR2EXn typically exhibit  $C_{S}$ -symmetric, seco-heterocube type structures in which the main group element fragment (e.g.  $EX_n = Mg$ , BPh, AlCl, GaCl, Ge, Sn, SnCl<sub>2</sub>, SiCl<sub>2</sub>, AsCl, SbCl, BiCl, R = t-Bu, Ph) is coordinated by three N atoms and occupies the edge of a distorted cube.15

The molecular arrangement observed for 7 in the solid state also persists in solution. The observation of two sets of resonances for its chemically inequivalent Me<sub>3</sub>Si-groups in the  $^1$ H NMR spectrum ( $\delta(H) = 0.21$  and 0.40 ppm) confirm the  $C_{2V}$ -symmetry. The  $^{13}$ C{ $^1$ H} NMR spectrum shows a triplet resonance which is assigned to the carbon atoms of the Me<sub>3</sub>Si-groups attached to the [PN]<sub>2</sub>-ring ( $\delta(C) = -0.1$  ppm,  $^3J(CP) = 3.6$  Hz). A *pseudo*-triplet is observed for the Me<sub>3</sub>Si-groups adjacent to the N<sub>2</sub>SiCl<sub>2</sub>-moiety ( $\delta(C) = 2.3$  ppm,  $|^3J(CP) + ^5J(CP)| = 5.0$  Hz) and is a result of an AA'X<sub>3</sub>X'<sub>3</sub> spin system (A =  $^{31}$ P,

Scheme 1 Preparation of compounds 6 and 7; (a)  $+SiCl_4$ ,  $-Me_3SiCl$ , neat, 80 °C, 3 d.

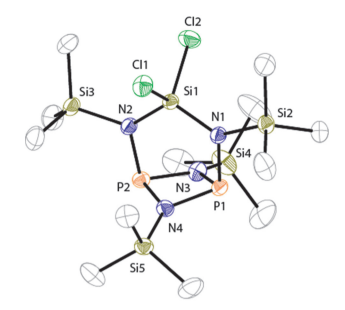


Fig. 2 Molecular structure of 7 (hydrogen atoms are omitted for clarity and thermal ellipsoids are displayed at 50% probability); selected bond lengths [Å] and angles [°]: N1–P1 1.734(1), N3–P1 1.721(1), N4–P1 1.719 (4), N2–P2 1.738(1), N3–P2 1.724(1), N4–P2 1.720(1), P1···P2 2.5235(6), N1–Si1 1.708(1), N1–Si2 1.773(1), N2–Si1 1.710(1), N2–Si3 1.774(1), N3–Si4 1.728(1), N4–Si5 1.740(1), Si1–Cl2 2.0554(6), Si2–Cl1 2.5235(6); P1–N4–P2 94.39(6), P1–N3–P2 94.19(7), N3–P1–N4 84.06(7), N3–P2–N4 83.96(7), N1–P1–N3 100.48(6), N1–P1–N4 102.10(6), N2–P2–N3 100.29(7), N2–P2–N4 101.91(6), N1–P1···P2 95.84(4), N2–P2···P1 95.58(4), Si1–N1–P1 117.14(7), Si1–N2–P2 117.18(8), N1–Si1–N2 114.19(6), Cl1–Si1–Cl2 104.35(2).

 $X={}^{13}\mathrm{C})$  with a comparatively large  ${}^2J(\mathrm{P_AP_{A'}})$  coupling constant. The same arguments account for the *pseudo*-triplet resonance in the  ${}^{29}\mathrm{Si}\{^1\mathrm{H}\}$  NMR spectrum of 7 ( $\delta(\mathrm{Si})=7.7$  ppm) which is assigned to the Me<sub>3</sub>Si-groups adjacent to the N<sub>2</sub>SiCl<sub>2</sub>-moiety ( $|{}^2J(\mathrm{SiP})| + {}^4J(\mathrm{SiP})| = 12.5$  Hz). The triplet resonance corresponding to the Me<sub>3</sub>Si-groups attached to the [PN]<sub>2</sub>-ring ( $\delta(\mathrm{Si})=1.8$  ppm,  ${}^3J(\mathrm{SiP})=11.7$  Hz) and the resonance of the SiCl<sub>2</sub> moiety ( $\delta(\mathrm{Si})=-47.4$  ppm) are observed in the expected regions. The same arguments account for the pseudo-triplet resonance of the sicl<sub>2</sub> moiety ( $\delta(\mathrm{Si})=-47.4$  ppm) are observed in the expected regions.

Due to the low isolated yield of 7, it was of interest to develop an alternative synthetic approach. Compound 8 is the head to tail dimer of iminophosphane 5 and was obtained according to a literature known procedure. The solvent free reaction of 8 with an excess of SiCl4 gave selectively and quantitatively compound 9 via Me<sub>3</sub>SiCl elimination (Scheme 2).<sup>19</sup> Compound 9 is the trans-conformer of an unsymmetricallysubstituted diphosphadiazane derivative and, thus, shows an AX spin system in its  ${}^{31}P\{{}^{1}H\}$  NMR spectrum ( $\delta(P_A)$  = 219.5 ppm,  $\delta(P_X) = 232.3$  ppm,  ${}^2J(P_AP_X) = 12.0$  Hz). The  ${}^1H$ NMR spectrum of 9 shows singlet resonances for the SiMe<sub>3</sub>groups on the [NP]<sub>2</sub>-ring and for both pointing towards the plane of the four-membered ring. The SiMe<sub>3</sub> group pointing away from the ring, however, shows a relatively large <sup>4</sup>*I*(HP) coupling constant ( $\delta(H) = 0.27$  ppm,  ${}^4J(P_AH) = 3.7$  Hz). For structurally related compounds, this was rationalized by the

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Scheme 2 Synthesis of 9 via reaction of 8 and SiCl<sub>4</sub>; (a) +SiCl<sub>4</sub>, -Me<sub>3</sub>SiCl, rt, 12 h, 96%; and subsequent transformation to 7; (b) neat, 185 °C, 10 min, 25%.

close proximity of the CH3-groups to the lone pair of electrons on the adjacent P atom.20 Due to the same reason, two relatively large 4J(PH) coupling constants (3.4 and 3.7 Hz) were observed for the isomer of 9 in which the SiCl<sub>3</sub>-group points towards the face of the four-membered ring (9'). 19 The chlorosubstituted Si atom in 9 appears as doublet resonance in the typical range in the <sup>29</sup>Si NMR spectrum  $(\delta(Si) = -27.3 \text{ ppm})^{21}$ with a remarkably large coupling constant of  ${}^{2}I(SiP_{x}) =$ 26.0 Hz. A coupling constant of similar magnitude is observed for the Me<sub>3</sub>Si-group pointing away from the four-membered ring  $(\delta(Si) = 7.8 \text{ ppm}, {}^2I(SiP_A) = 32.4 \text{ Hz})$ , whereas those pointing to the faces of the [NP]2-ring exhibit rather small coupling constants ( $\delta(Si) = 4.9$  and 0.2 ppm;  ${}^{2}J(SiP) = 4.2$  and 3.0 Hz). Similarly the <sup>13</sup>C{<sup>1</sup>H} NMR spectrum reveals rather small coupling constants for the latter Me<sub>3</sub>Si-groups ( ${}^{3}J(CP) = 8.0$  and 10.5 Hz) and a large coupling constant for the Me<sub>3</sub>Si-group pointing away from the  $[NP]_2$ -ring plane  $(^3J(CP_A) = 19.8 \text{ Hz})$ .

It is assumed, that 7 is obtained from 9 via an intermediate of type 10 which forms upon rotation<sup>22</sup> of the P-N bond involving the SiCl<sub>3</sub>-substituted N atom and inversion<sup>23</sup> of one of the P centers in 9 (Scheme 2). The arrangement of the SiCl<sub>3</sub>-group and one of the SiMe<sub>3</sub>-groups in intermediate 10 allows for Me<sub>3</sub>SiCl elimination yielding 7. Indeed, heating 9 for ten minutes to 185 °C results in conversion to 7 in 25% yield.<sup>24</sup>

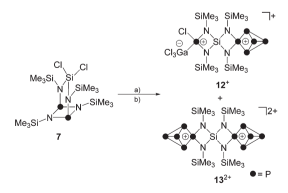
## Reactivity of 7 towards GaCl<sub>3</sub> and P<sub>4</sub>

Diphosphadiazanes and related compounds bearing chloroand Me<sub>3</sub>Si-substituents are known to undergo a variety of distinct reactions with Lewis acids. Next to halide abstraction, 10 also elimination of Me<sub>3</sub>SiCl, <sup>25</sup> migration of Me<sub>3</sub>Si-, <sup>26</sup> chloro- or methyl-substituents<sup>27</sup> and P-N bond cleavage reactions are reported.<sup>28</sup> Especially the latter reaction is of interest, since it is assumed to proceed via phosphenium ion intermediates. Thus, bicyclic compound 7 is a promising substrate for the generation of phosphenium ions and, subsequently, insertion of the latter into P-P bonds of P4.

Scheme 3 Formation of Lewis-acid/base adduct 11 via the reaction of 7 and GaCl<sub>3</sub>; (a) +GaCl<sub>3</sub>, C<sub>6</sub>H<sub>5</sub>F, r.t., 1 h.

Thus, the reaction of 7 with the Lewis acid GaCl<sub>3</sub> was probed (Scheme 3). The addition of one equivalent of GaCl<sub>3</sub> to a solution of 7 in C<sub>6</sub>H<sub>5</sub>F gave a reddish colored reaction mixture which was investigated by means of <sup>31</sup>P{<sup>1</sup>H} NMR spectroscopy. The formation of the Lewis-acid/base adduct 11 is proposed on the basis of the observation of two broad resonances in an approximate ratio of 1:1 ( $\delta(P_A)$  = 115 ppm,  $\Delta \nu_{1/2}$  = 1200 Hz;  $\delta(P_X)$  = 179.6 ppm,  $\Delta \nu_{1/2}$  = 600 Hz). The coordination of GaCl3 to a P atom is very likely, since they have been identified as the most basic sites in related compounds.29 The resonance at low field is tentatively assigned to the tri-coordinated P atom. Accordingly, the resonance at high field corresponds to the tetra-coordinated P atom and exhibits a stronger line broadening due to the coordination of GaCl<sub>3</sub>.

Interestingly, P4 does not react with 11 which indicates that the latter is not a suitable phosphenium ion source. Adding two equivalents of GaCl<sub>3</sub> to a solution of 7 in C<sub>6</sub>H<sub>5</sub>F, however, results in the rapid consumption of in situ generated 11. This reaction yields a complex mixture of not identified products and bodes well for the generation of reactive intermediates that might be able to activate P4. Indeed, mixtures of 7, P4 and GaCl<sub>3</sub> in 1:1:2 and 1:2:4 stoichiometries were bright red and the consumption of P4 accompanied by a color change to brown was observed. Subsequent investigation by means of <sup>31</sup>P{<sup>1</sup>H} NMR spectroscopy revealed the formation of P<sub>5</sub><sup>+</sup> cage cation 12<sup>+</sup> and bridged bis(P<sub>5</sub><sup>+</sup>)-cage dication 13<sup>2+</sup> by their characteristic A<sub>2</sub>MOX<sub>2</sub> and A<sub>2</sub>MX<sub>2</sub> spin systems (Scheme 4).



Scheme 4 Reaction of 7 with P<sub>4</sub> and GaCl<sub>3</sub> in various stoichiometries; (a)  $+P_4$ , +2,  $GaCl_3$ ,  $C_6H_5F$ , r.t., 24 h; (b) +2,  $P_4$ , +4,  $GaCl_3$ , r.t.,  $C_6H_5F$ , 24 h; the equation depicts the major products and is not balanced; anions of the products are not depicted.

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In addition, both spectra reveal significant amounts of P<sub>4</sub> but neither remaining 7 nor the corresponding adduct 11 were observed which is attributed to unidentified side reactions. The products 12<sup>+</sup> and 13<sup>2+</sup> are formally obtained by the insertion of phosphenium ions based on ClP(NSiMe<sub>3</sub>)<sub>2</sub>Si(NSi-Me<sub>3</sub>)<sub>2</sub>PCl which features a Si-centered spiro[3.3]heptane-motif, into P-P bonds of P<sub>4</sub>. Interestingly, both reactions yield 12<sup>+</sup> and  $13^{2+}$  in comparable ratios (1:1.8 for (a) and 1:1.5 for (b), Scheme 4). This contrasts the anticipated increase of the amount of dication 132+ in the presence of excess P4 and GaCl<sub>3</sub>. In addition, the formation of large quantities of 13<sup>2+</sup> in the reaction of 1:1:2 stoichiometry indicates that the reaction rate of a GaCl3-induced rearrangement of 7 yielding the spiromotif is slow compared to that of the subsequent phosphenium ion insertion.

A tentative mechanism for the formation of such a Si-centered spiro[3.3]heptane-type species from 7 is illustrated in Scheme 5. While the reaction of 7 with one equivalent GaCl<sub>3</sub> gives the Lewis-acid/base adduct 11 (vide supra), an excess of GaCl<sub>3</sub> is assumed to initiate a chloride abstraction from the SiCl<sub>2</sub>-moiety. This is facilitated by the tendency of bicyclic compounds of type 7 to form a seco-heterocube-type structure. 15 Thus, compound 7 might be in equilibrium with the related derivative 7<sup>I</sup>. Compound 7<sup>I</sup> features a hypervalent penta-coordinated Si-moiety which might favor the sequestering of a chloride anion. Thus, it is assumed that the reaction with GaCl<sub>3</sub> proceeds via chloride abstraction and formation of the cationic seco-heterocube 14<sup>+</sup> featuring a tetra-coordinated

7 
$$Cl \odot N R R + GaCl_3 - GaCl_4$$
 $R \oplus N R R R$ 
 $R \oplus N R R R$ 
 $R \oplus N R$ 

Scheme 5 Suggested GaCl<sub>3</sub>-induced rearrangement mechanism of 7 to phosphenium ion 18+ featuring a Si-centered spiro[3.3]heptane-type motif; R denotes Me<sub>3</sub>Si-substituents; for reasons of simplification the reactions of intermediates or products possessing a di-coordinated P atom with P4 as well as retransfer of a chloride ion to a di-coordinated P atom are not considered.

N atom. The next two steps constitute P-N bond cleavage and formation reactions yielding intermediates 15<sup>+</sup> and 16<sup>+</sup> via formal retention of the seco-heterocube-type structure. Similar P-N bond ruptures were reported as decomposition pathways of diamino-cyclo-diphosphadiazanes.<sup>26</sup> Intermediate 16<sup>+</sup> features a diphosphadiazane [NP]<sub>2</sub>-ring which is assumed to react via a cyclo-reversion reaction to intermediate 17<sup>+</sup>. Cation 17<sup>+</sup> features an aminoiminophosphane moiety similar to 5 which is tethered to a four-membered SiN2P-ring. Intramolecular nucleophilic attack of the imino-N atom on the chloro-substituted Si atom initiates a GaCl3-mediated transfer of a chloride anion from the Si atom to a di-coordinated P atom to give formally the phosphenium ion 18<sup>+</sup>. Cation 18<sup>+</sup> features the Sicentered spiro[3,3]heptane-motif and, thus, is assumed to be accountable for the formation of cages cations 12<sup>+</sup> and 13<sup>2+</sup> via insertion into a P-P bond of P4, chloride abstraction by GaCl3 and subsequent insertion into a P-P bond of a second P4 tetrahedron.

Attempts to isolate a gallate salt of 12<sup>+</sup> from both reaction mixtures were unsuccessful, possibly due to the fluxional coordination behavior of the GaCl3-molecule to the PCl-function. However, slow diffusion of n-hexane into the reaction mixture of 1:1:2 stoichiometry yielded compound 19[GaCl<sub>4</sub>] in low yields (10%, Scheme 6). The <sup>31</sup>P{<sup>1</sup>H} NMR spectrum of 19[GaCl<sub>4</sub>] dissolved in CD<sub>2</sub>Cl<sub>2</sub> shows an ABMX<sub>2</sub>Y spin system which is in accordance with the  $C_s$  symmetry of the molecule (Fig. 3). The mirror plane is defined by the tetra-coordinated P atom and the two adjacent P atoms. The P-Cl unit is included in the plane and exhibits a spatial proximity to one of the two bridge-head P atoms. The chemical shifts and coupling constants involving the P5+cage motif are similar to those observed for the cage cation 3<sup>+</sup>. The chloro-substituted P atom in  $\mathbf{19}^+$  exhibits a singlet resonance ( $\delta(P_Y)$  = 166.8 ppm) in the typical range of silyl-substituted diamino-chlorophosphanes. 13,30 A 4/(PP) coupling between this P atom and the tetra-coordinated P atom of the P5+-cage is not resolved. The <sup>1</sup>H NMR spectrum of 19<sup>+</sup> reveals three singlet resonances assigned to the chemically different Me<sub>3</sub>Si-groups which integrate in a 2:1:1 ratio. The high-field resonance ( $\delta(H)$  = 0.31 ppm) exhibits the highest intensity and is assigned to the two Me<sub>3</sub>Si-groups bonded to the four-membered ring which incorporates the P-Cl moiety. The chemical shift is comparable to related four-membered ring compounds (6:  $\delta(H)$  = 0.18 ppm, <sup>13</sup> 7:  $\delta(H) = 0.21$  ppm). Both resonances assigned to the Me<sub>3</sub>Si-groups bonded to the N atoms adjacent to the

Reaction of 19[GaCl<sub>4</sub>] with GaCl<sub>3</sub>; (a) +GaCl<sub>3</sub>, CD<sub>2</sub>Cl<sub>2</sub>, r.t., Scheme 6

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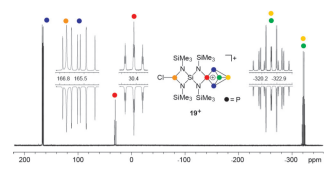


Fig. 3 <sup>31</sup>P{<sup>1</sup>H} NMR spectrum of **19**[GaCl<sub>4</sub>] (CD<sub>2</sub>Cl<sub>2</sub>, r.t.); insets show experimental (upwards) and fitted spectra (downwards); ABMX2Y spin system of 19<sup>+</sup>:  $\delta(P_A) = -322.9$  ppm,  $\delta(P_B) = -320.2$  ppm,  $\delta(P_M) =$ 30.4 ppm,  $\delta(P_X) = 165.5$  ppm,  $\delta(P_Y) = 166.8$  ppm,  ${}^1J(P_AP_B) = -189.8$  Hz,  ${}^{1}J(P_{A}P_{X}) = -143.4 \text{ Hz}, {}^{1}J(P_{B}P_{X}) = -147.7 \text{ Hz}, {}^{1}J(P_{M}P_{X}) = -245.1 \text{ Hz},$  $^{2}J(P_{A}P_{M}) = 18.8 \text{ Hz}, ^{2}J(P_{B}P_{M}) = 18.0 \text{ Hz}.$ 

 $P_5^+$ -moiety exhibit a low field shift ( $\delta(H) = 0.67$  ppm and 0.76 ppm) and are comparable to the corresponding resonance of  $3^+$  ( $\delta(H) = 0.68$  ppm). <sup>10a</sup> The <sup>29</sup>Si $\{^1H\}$  NMR spectrum exhibits a resonance at  $\delta(Si) = -58.0$  ppm which is assigned to the Si spiro-atom. This resonance reveals a doublet of doublet splitting caused by  ${}^{2}J(SiP)$ -couplings to the tetra-coordinated P atom ( ${}^{2}J(SiP_{M}) = 8.5 Hz$ ) and the chloro-substituted P atom ( ${}^{2}J$  $(SiP_{y}) = 18.5 \text{ Hz}$ ).

The molecular structure of 19<sup>+</sup> is depicted in Fig. 4 and the P-P bond lengths and angles in the P<sub>5</sub><sup>+</sup>-moiety are comparable to those of 3<sup>+</sup>. 10a Both four-membered rings are almost planar (largest deviation from the planes N1: 0.026 Å and N3: 0.022 Å) and exhibit a perpendicular arrangement (angle between both planes: 89.79(9)°). Due to the steric limitations of the fourmembered heterocycles the spiro-Si atom exhibits a distorted tetrahedral arrangement with two rather small (N1-Si3-N2: 85.4(1)°, N3-Si3-N4: 87.7(1)°) and two widened (N1-Si3-N3: 122.9(1)°, N2-Si3-N4: 118.8(1)°) N-Si-N angles. Alternating bond lengths are observed within the two four-membered rings. The P-N bonds involving the tetra-coordinated P atom are shorter (N2-P1: 1.656(3) Å, N1-P1: 1.661(3) Å) than those involving the tri-coordinated P atom (N3-P6: 1.716(3) Å, N4-P6: 1.711(3) Å) and both magnitudes of bond lengths are also observed in the related cages 3<sup>+</sup> and 2<sup>+</sup>. 9,10a The Si-N bonds in the [SiN<sub>2</sub>P]-ring fused to the P<sub>5</sub><sup>+</sup>-cage (N1-Si3: 1.755(3) Å, N2-Si3: 1.751(3) Å) are of similar lengths as observed for  $3^{+}$ . <sup>10a</sup> In contrast, the Si-N bonds in the second [SiN2P]-ring are shorter (N3-Si3: 1.711(3) Å, N4-Si3: 1.704(3) Å).

The addition of GaCl<sub>3</sub> to a solution of 19[GaCl<sub>4</sub>] in CD<sub>2</sub>Cl<sub>2</sub> yields the previously mentioned Lewis-acid base adduct 12<sup>+</sup> (Scheme 6). The  $A_2MOX_2$  spin system observed in the  $^{31}P\{^1H\}$ NMR spectrum of  $12^+$  suggests a time averaged  $C_{2v}$ -symmetry of the molecule in solution which can be explained by a fluxional behavior of the coordinated GaCl<sub>3</sub> molecule (Fig. 5). The A<sub>2</sub>MX<sub>2</sub> part of the A<sub>2</sub>MOX<sub>2</sub> spin system corresponds to the P<sub>5</sub><sup>+</sup>cage moiety and reveals comparable chemical shifts and coupling constants as observed for 3+.10a The resonance corresponding to the O part of the spin system is very broad ( $\delta(P_O)$  =

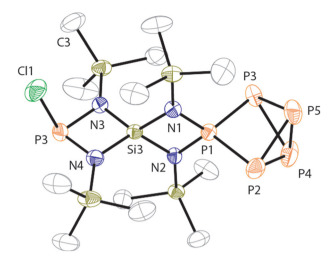


Fig. 4 Molecular structure of 19<sup>+</sup> in compound 19[GaCl<sub>4</sub>]·C<sub>6</sub>H<sub>5</sub>F (hydrogen atoms are omitted for clarity and thermal ellipsoids are displayed at 50% probability); selected bond lengths [A] and angles [°]: N2-P1 1.656(3), N1-P1 1.661(3), N1-Si3 1.755(3), N2-Si3 1.751(3), N3-Si3 1.711(3), N4-Si3 1.704(3), N3-P6 1.716(3), N4-P6 1.711(3), Cl1-P6 2.238(2), P1...Si3 2.445(1), P6...Si3 2.472(1), P1-P2 2.167(1), P1-P3 2.165(1), P2-P4 2.249(2), P2-P5 2.238(2), P3-P4 2.251(2), P3-P5 2.229(2), P4-P5 2.169(2); N1-P1-N2 91.5(1), P1-N1-Si3 91.4(1), P1-N2-Si3 91.7(1), N1-Si3-N2 85.4(1), N3-Si3-N4 87.7(1), N1-Si3-N3 122.9(1), N2-Si3-N4 118.8(1), N3-P6-N4 87.2(1), N3-P6-Cl1 102.2(1), N4-P6-Cl1 101.7(1), P3-P1-P2 91.15(5), P1-P2-P5 83.95(5), P1-P2-P4 82.53(5), P5-P2-P4 57.82(5), P5-P4-P2 60.83(5), P2-P5-P3 87.67(5).

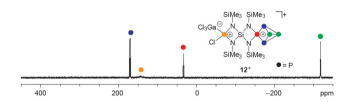


Fig. 5 <sup>31</sup>P{<sup>1</sup>H} NMR spectrum of 12[GaCl<sub>4</sub>] (CD<sub>2</sub>Cl<sub>2</sub>, r.t.); A<sub>2</sub>MOX<sub>2</sub> spin system of 12<sup>+</sup>:  $\delta(P_A) = -318.3$  ppm,  $\delta(P_M) = 33.4$  ppm,  $\delta(P_O) = 144$  ppm  $(\Delta \nu_{1/2} = \sim 1200 \text{ Hz}), \ \delta(P_X) = 170.8 \text{ ppm}, \ ^1J(P_AP_X) = -144.8 \text{ Hz}, \ ^1J(P_MP_X) =$ -248.6 Hz,  ${}^{2}J(P_{A}P_{M}) = 19.0 \text{ Hz}$ .

144 ppm,  $\Delta \nu_{1/2} = \sim 1200$  Hz) which is caused by the fluxional behavior of the coordinated GaCl3 molecule and its quadrupole moment.

Dication 13<sup>2+</sup> was isolated as a [Ga<sub>2</sub>Cl<sub>7</sub>]<sup>-</sup> salt from the reaction of 7, P<sub>4</sub> and GaCl<sub>3</sub> in a 1:2:4 stoichiometry (Scheme 4) by the addition of n-hexane. This gave a brown oil which was isolated by decanting the supernatant and upon addition of small amounts of 1,2-C<sub>6</sub>H<sub>4</sub>F<sub>2</sub> yielded a suspension containing a yellow microcrystalline material of 13[Ga<sub>2</sub>Cl<sub>7</sub>]<sub>2</sub>. This compound was isolated by filtration in low yields (20%) in an approximate purity of 75%. Further purification by recrystallization from CH2Cl2/n-hexane leads to significantly decreased yields (7% yield in >90% purity). The <sup>31</sup>P{<sup>1</sup>H} NMR spectrum of 13[Ga<sub>2</sub>Cl<sub>7</sub>]<sub>2</sub> dissolved in CD<sub>2</sub>Cl<sub>2</sub> shows an A<sub>2</sub>MX<sub>2</sub>-spin system in accordance with the  $C_{2v}$ -symmetry of the molecule

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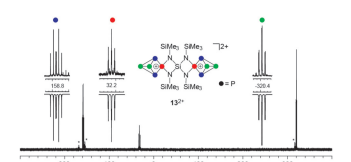


Fig. 6  $^{31}$ P{ $^{1}$ H} NMR spectrum of 13[Ga<sub>2</sub>Cl<sub>7</sub>]<sub>2</sub> (CD<sub>2</sub>Cl<sub>2</sub>, r.t.); unidentified side products are marked with asterisks; A<sub>2</sub>MX<sub>2</sub> spin system of 13<sup>2+</sup>:  $\delta$ (P<sub>A</sub>) = -320.4 ppm,  $\delta$ (P<sub>M</sub>) = 32.2 ppm,  $\delta$ (P<sub>X</sub>) = 158.8 ppm,  $^{1}$ J(P<sub>A</sub>P<sub>X</sub>) = -142.7 Hz,  $^{1}$ J(P<sub>M</sub>P<sub>X</sub>) = -259.1 Hz,  $^{2}$ J(P<sub>A</sub>P<sub>M</sub>) = 20.0 Hz.

(Fig. 6). The  $C_2$ -axis includes both tetra-coordinated P atoms and the spiro-Si atom and the mirror planes are defined by the four-membered [SiN<sub>2</sub>P]-rings. The resonances and coupling constants of  $13^{2+}$  are similar to those observed for the related compounds  $19^+$  and  $3^+$ .<sup>10 $\alpha$ </sup>

Single crystals of compound 13[Ga<sub>2</sub>Cl<sub>7</sub>]<sub>2</sub> were obtained by diffusion of *n*-hexane in a  $CH_2Cl_2$  solution of  $13[Ga_2Cl_7]_2$  at −35 °C. The compound crystallizes with two independent formula units in the asymmetric unit. Two of the four Ga<sub>2</sub>Cl<sub>7</sub><sup>-</sup> anions are highly disordered exhibiting unusually high thermal displacement parameters (see ESI† for details). Single crystals of 13[Ga<sub>2</sub>Cl<sub>7</sub>][GaCl<sub>4</sub>] were obtained by layering the supernatant solution of the reaction mixture of the synthesis of  $13[Ga_2Cl_7]_2$  with *n*-hexane at -35 °C. The data obtained by X-ray single crystal structure determination was of higher quality and, thus, the molecular structure of 13[Ga<sub>2</sub>Cl<sub>7</sub>]-[GaCl<sub>4</sub>] is discussed (Fig. 7). The P-P bond lengths and angles in the P<sub>5</sub><sup>+</sup>-moieties are comparable to those of related P<sub>5</sub><sup>+</sup>-cage compounds. 9,10a The P-N bonds in 132+ are rather short (av. P-N: 1.666(8) Å) which is a typical feature of P-N bonds involving phosphonium moieties. The Si-N bond lengths in the fourmembered rings are nearly identical (av. Si-N: 1.732(8) Å) and are between the two types of bond lengths observed for 19<sup>+</sup> (av. Si-N: 1.753(6) Å and Si-N: 1.7007(6) Å).

## Conclusions

The bicyclic P–N–Si heterocycle 7 was targeted as source for the *in situ* generation of phosphenium cations for  $P_4$  activation. In this context, two distinct synthetic protocols for its preparation were thoroughly investigated and gave insights into the formation of the bicycle. The reaction of 7 with  $GaCl_3$  initially yields adduct 11. This adduct is not stable and subsequently rearranges to give *in situ* spirocyclic, Si-centered compound  $ClP(NSiMe_3)_2Si(NSiMe_3)_2PCl$ . The latter species gives access to polyphosphorus cage cations  $[ClP(NSiMe_3)_2Si(NSiMe_3)_2P_5]^+$  (19<sup>+</sup>) and  $[P_5(NSiMe_3)_2Si(NSiMe_3)_2P_5]^{2+}$  (13<sup>2+</sup>) in the presence of  $GaCl_3$  and  $P_4$ . We are continuing to investigate the Lewis-acid mediated generation of phosphenium ions

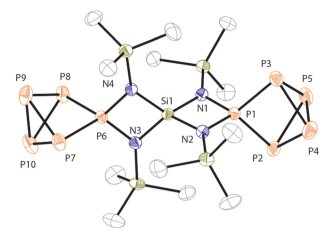


Fig. 7 Molecular structure of  $13^{2+}$  in compound  $13[Ga_2Cl_7][GaCl_4]$  (hydrogen atoms are omitted for clarity and thermal ellipsoids are displayed at 50% probability); selected bond lengths [Å] and angles [°]: N1–P1 1.667(2), N2–P1 1.671(2), N3–P6 1.663(2), N4–P6 1.665(2), Si1–N1 1.731(2), Si1–N2 1.732(2), S1–N3 1.736(2), Si1–N4 1.728(2), P1···Si1 2.4452(8), P6···Si1 2.4277(8), P1–P2 2.1540(8), P1–P3 2.1499(8), P2–P4 2.245(1), P2–P5 2.245(1), P3–P4 2.246(1), P3–P5 2.2464(9), P4–P5 2.168(1), P6–P7 2.1511(8), P6–P8 2.1572(2), P7–P9 2.240(1), P7–P10 2.252(1), P8–P9 2.2415(9), P8–P10 2.242(1), P9–P10 2.172(1); N1–P1–N2 90.81(9), N3–P6–N4 90.99(9), N1–Si1–N2 86.73(9), N3–Si1–N4 86.49(9), P1–N1–Si1 91.26(9), P1–N2–Si1 91.12(9), P6–N3–Si1 91.14(9), P6–N4–Si1 91.34(9), P3–P1–P2 92.59(3), P1–P2–P5 81.73(3), P1–P2–P4 83.19(3), P5–P2–P4 57.69(3), P5–P4–P2 61.25(3), P2–P5–P3 87.60(3), P7–P6–P8 92.30(3), P6–P7–P10 82.39(3), P6–P7–P9 82.89(3), P10–P7–P9 57.83(3), P10–P9–P7 61.36(3), P7–P10–P8 87.48(3).

for  $P_4$  activation from related phosphorus–nitrogen-element bicycles. Furthermore, studies directed towards the utilization of  $\mathbf{12}^+$ ,  $\mathbf{19}^+$  and  $\mathbf{13}^{2+}$  as synthetic building blocks will be the target of future efforts.

## Experimental

## General

General information on materials and methods as well as  $^{31}P\{^{1}H\}$  NMR spectra of reaction mixtures are given in the ESI.†

#### Synthesis of Cl<sub>2</sub>Si(NSiMe<sub>3</sub>)<sub>2</sub>(PNSiMe<sub>3</sub>)<sub>2</sub> (7)

*Method A*: The literature reported synthesis of 5 was performed on a 20 mmol scale. <sup>13</sup> Compound 5 was removed by distillation from the reaction mixture (40 °C,  $8 \times 10^{-2}$  mbar). The remaining colorless, slushy residue was dissolved in  $C_6H_5F$  (5 mL) yielding a turbid suspension. The solvent was removed *in vacuo* yielding a sludgy residue which was redistilled employing a short Vigreux column (5 cm). Compound 7 was obtained as colorless oil (1.789 g, 3.51 mmol, 18%, 105 °C,  $2 \times 10^{-3}$  mbar) which solidified shortly after distillation. *Method B*: 9 (305 mg, 0.50 mmol, 1.0 eq.) was heated to 185 °C for 10 min. In the course of the reaction a colorless liquid is formed accompanied by the condensation of Me<sub>3</sub>SiCl on

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colder parts of the reaction vessel. After cooling to ambient temperature the reaction mixture remains a liquid. Isolation of 7 from this mixture proceeds as described in method A.

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m.p.: 55.6–57.8 °C; Raman (300 mW, [cm<sup>-1</sup>]):  $\nu$  = 2959 (390), 2899 (100), 1410 (11), 690 (10), 645 (22), 613 (51), 562 (4), 349 (34), 185 (30), 141 (6), 75 (10); IR (ATR, [cm<sup>-1</sup>]):  $\nu$  = 2956 (w), 1408 (vw), 1249 (s), 1098 (vw), 973 (vw), 942 (w), 883 (w), 830 (vs), 776 (vw), 754 (vw), 713 (w), 682 (w), 643 (vw), 556 (s), 450 (m); <sup>1</sup>H NMR (C<sub>6</sub>D<sub>6</sub>, [ppm]):  $\delta$  = 0.21 (18H, s, H1), 0.39 (18H, s, H2); <sup>13</sup>C{<sup>1</sup>H} NMR (C<sub>6</sub>D<sub>6</sub>, [ppm]):  $\delta$  = -0.08 (6C, t, C1, <sup>3</sup>J(CP) = 3.6 Hz), 2.7 (6C, pseudo-t, C2, <sup>3</sup>J(CP) = 5.0 Hz); <sup>29</sup>Si {<sup>1</sup>H} NMR (C<sub>6</sub>D<sub>6</sub>, [ppm]):  $\delta$  = -47.4 (1Si, s, Si3), 1.8 (2Si, t, Si1, <sup>2</sup>J(SiP) = 11.7 Hz), 7.7 (2Si, pseudo-t, Si2, <sup>2</sup>J(SiP) = 11.8 Hz); <sup>15</sup>N NMR (C<sub>6</sub>D<sub>6</sub>, [ppm]):  $\delta$  = -397 (t, N1, <sup>1</sup>J(NP) = 55 Hz), -374 (d, N2, <sup>1</sup>J(NP) = 75 Hz); <sup>31</sup>P{<sup>1</sup>H} NMR (C<sub>6</sub>D<sub>6</sub>, [ppm]):  $\delta$  = 211.8 (s); elemental analysis for C<sub>12</sub>H<sub>36</sub>Cl<sub>2</sub>N<sub>4</sub>P<sub>2</sub>Si<sub>5</sub>: calcd: C 28.3, H 7.1, N 11.0; found: C 28.5, H 7.3, N 10.6; MS-ESI-EM: 473.0943 [M-Cl<sup>-</sup>], calcd: for C<sub>12</sub>H<sub>36</sub>Cl<sub>1</sub>N<sub>4</sub>P<sub>2</sub>Si<sub>5</sub>: 473.0945.

## Synthesis of (SiMe<sub>3</sub>)<sub>2</sub>N(PNSiMe<sub>3</sub>)<sub>2</sub>N(SiMe<sub>3</sub>)(SiCl<sub>3</sub>) (9)

Compound **8** (1.114 g, 2.00 mmol, 1.0 eq.) was suspended in  $SiCl_4$  (6.796 g, 40.0 mmol, 40.0 eq.) and stirred for 12 h at ambient temperature. After removal of all volatiles *in vacuo* **9** was isolated in quantitative yields as colorless solid (1.187 g, 1.92 mmol, 96%).

m.p.: 180.2–182.5 °C; Raman (300 mW, [cm<sup>-1</sup>]):  $\nu = 2957$ (37), 2906 (100), 1410 (15), 686 (13), 651 (4), 641 (64), 586 (21), 488 (4), 436 (38), 351 (21), 206 (37), 107 (27), 78 (18); IR (ATR, [cm<sup>-1</sup>]):  $\nu = 3139$  (vw), 3047 (m), 2958 (vw), 1406 (m), 1251 (s), 1060 (m), 923 (m), 834 (vs), 778 (vw), 755 (w), 712 (w), 677 (w), 572 (vw), 559 (m), 506 (w), 432 (vw); <sup>1</sup>H NMR (C<sub>6</sub>D<sub>6</sub>, [ppm]):  $\delta$  = 0.17 (18H, s, H5), 0.27 (9H, d, H3,  ${}^{4}J(HP) = 3.7 \text{ Hz}$ ), 0.52 (9H, s, H4), 0.64 (9H, s, H1);  ${}^{13}C\{{}^{1}H\}$  NMR ( $C_6D_6$ , [ppm]):  $\delta = 1.4$  (6C, t, C5,  ${}^{3}J(CP) = 2.5 \text{ Hz}$ , 4.7 (3C, d, C1,  ${}^{3}J(CP) = 10.5 \text{ Hz}$ ), 4.8 (3C, d, C3,  ${}^{3}J(CP) = 19.8 \text{ Hz}$ ), 5.0 (3C, d, C4,  ${}^{3}J(CP) = 8.0 \text{ Hz}$ ); <sup>29</sup>Si{<sup>1</sup>H} NMR (C<sub>6</sub>D<sub>6</sub>, [ppm]):  $\delta = -27.3$  (1Si, d, Si2,  ${}^{2}J$ (SiP) = 26.0 Hz), 0.2 (1Si, d, Si4,  ${}^{2}J(SiP) = 3.0$  Hz), 1.5 (2Si, t, Si5,  $^{2}J(SiP) = 7.4 Hz$ , 4.5 (1Si, d, Si2,  $^{2}J(SiP) = 4.2 Hz$ ), 7.6 (1Si, d, Si3,  ${}^{2}J(SiP) = 32.4 \text{ Hz});$  <sup>15</sup>N NMR (C<sub>6</sub>D<sub>6</sub>, [ppm]):  $\delta = 67.1 \text{ (t, N2, }$  ${}^{1}J(NP) = 50 \text{ Hz}$ , 93.8 (d, N1,  ${}^{1}J(NP_{X}) = 90 \text{ Hz}$ ), 113.9 (d, N3,  ${}^{1}J(NP_{A}) = 90 \text{ Hz}; {}^{31}P\{{}^{1}H\} \text{ NMR } (C_{6}D_{6}, [ppm]): AX \text{ spin system:}$  $\delta(P_A) = 218.6$  (d, P1,  $\Delta \nu_{1/2} = 42$  Hz,  ${}^2J(P_AP_X) = 12$  Hz),  $\delta(P_X) =$ 

232.2 (d, P2,  $\Delta\nu_{1/2}$  = 37 Hz,  $^2J(P_AP_X)$  = 12 Hz); elemental analysis for  $C_{15}H_{45}Cl_3N_4P_2Si_6$ : calcd: C 29.1, H 7.3, N 9.1; found: C 28.8, H 7.3, N 8.6.

# Reaction of 7, $P_4$ and $GaCl_3$ in 1:1:2 and 1:2:4 stoichiometries

1:1:2: Compound 7 (256 mg, 0.50 mmol, 1.0 eq.) was added to a suspension of P<sub>4</sub> (62 mg, 0.50 mmol, 1.0 eq.) in C<sub>6</sub>H<sub>5</sub>F (5 mL). A solution of GaCl<sub>3</sub> (176 mg, 1.00 mmol, 2.0 eq.) in C<sub>6</sub>H<sub>5</sub>F (2 mL) was added dropwise to the suspension giving a red colored reaction mixture which was stirred for 12 h at ambient temperature. In the course of the reaction the color of the reaction mixture changed to yellow and the dissolving of  $P_4$  was observed. The reaction mixture was investigated by means of <sup>31</sup>P{<sup>1</sup>H} NMR spectroscopy (see ESI†). 1:2:4: Compound 7 (256 mg, 0.50 mmol, 1.0 eq.) was added to a suspension of P<sub>4</sub> (124 mg, 1.00 mmol, 2.0 eq.) in C<sub>6</sub>H<sub>5</sub>F (5 mL). A solution of GaCl<sub>3</sub> (352 mg, 2.00 mmol, 4 eq.) in C<sub>6</sub>H<sub>5</sub>F (4 mL) was added dropwise to the suspension giving a red colored reaction mixture which was stirred for 12 h at ambient temperature. In the course of the reaction the color of the reaction mixture changed to brown and the dissolving of P4 was observed. The reaction mixture was investigated by means of <sup>31</sup>P{<sup>1</sup>H} NMR spectroscopy (see ESI†). n-Hexane (2 mL) was added leading to the formation of a brown oil. The supernatant was removed, diluted with C6H5F (6 mL) and layered with *n*-hexane (3 mL) at -35 °C. Small amounts of crystalline material of 19[GaCl<sub>4</sub>] (41 mg, 10%), suitable for X-ray single crystal structure determination, were obtained within a few days. The remaining oil was washed with *n*-hexane  $(3 \times 3 \text{ mL})$ transforming it into a brown sludge. All volatiles were removed in vacuo and the sludge was suspended in 1,2-C<sub>6</sub>H<sub>4</sub>F<sub>2</sub> (2 mL) leading to the formation of a yellow, microcrystalline solid. The supernatant was removed and the yellow powder was washed with 1,2-C<sub>6</sub>H<sub>4</sub>F<sub>2</sub>/n-hexane (1:1 mixture, 2 × 2 mL). The obtained yellow powder consisting of 13[Ga2Cl7]2 in an approximate purity of 75% (determined by 31P{1H} NMR spectroscopy, 20% yield, 145 mg) was isolated by filtration and dried in vacuo. Recrystallization from a CH2Cl2 solution by slow diffusion of *n*-hexane yielded crystalline material of 19  $[Ga_2Cl_7]_2$  (purity > 90%) which was suitable for single crystal structure determination. Isolation was conducted via filtration and removal of all volatiles in vacuo (7% yield, 55 mg). Single crystals of 13[Ga2Cl7]-[GaCl4] were obtained by layering the diluted supernatant of the reaction mixture with n-hexane at −35 °C.

**m.p.:** 115.0–116.9 °C; **Raman (300 mW, [cm<sup>-1</sup>]):**  $\nu = 2961$  (37), 2998 (100), 1412 (24), 636 (37), 552 (63), 442 (22), 398 (24), 376 (14), 344 (43), 152 (44), 121 (13); **IR (ATR, [cm<sup>-1</sup>])**:

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 $\nu = 2956$  (vw), 2897 (vw), 1411 (vw), 1254 (m), 1123 (vw), 993 (s), 905 (s), 822 (vs), 756 (vw), 724 (s), 690 (w), 639 (w), 548 (vw), 509 (w), 462 (m); <sup>1</sup>H NMR (CD<sub>2</sub>Cl<sub>2</sub>, [ppm]):  $\delta = 0.31$  (18H, s, H1), 0.67 (9H, s, H2), 0.76 (9H, s, H3); <sup>13</sup>C{<sup>1</sup>H} NMR  $(CD_2Cl_2, \lceil ppm \rceil)$ :  $\delta = 1.4$  (6C, d, C1,  ${}^3J(CP) = 3.1$  Hz), 2.2 (3C, m, C2), 3.1 (3C, m, C3);  $^{29}Si\{^{1}H\}$  NMR (CD<sub>2</sub>Cl<sub>2</sub>, [ppm]):  $\delta =$ -58.0 (dd, Si4,  ${}^{2}J(SiP_{Y}) = 18.5$  Hz,  ${}^{2}J(SiP_{M}) = 8.5$  Hz), 7.4 (d, Si1,  ${}^{2}J(SiP) = 8.4 \text{ Hz}$ , 10.9 (s, Si2), 11.7 (s, Si3);  ${}^{15}N$  NMR  $(CD_2Cl_2, [ppm]): \delta = 96 \text{ (s, N1), } 115 \text{ (s, N2), } 125 \text{ (s(br), N3);}$ <sup>71</sup>Ga{<sup>1</sup>H} NMR (CD<sub>2</sub>Cl<sub>2</sub>, [ppm]):  $\delta = 249.6$  (s); <sup>31</sup>P{<sup>1</sup>H} NMR  $(CD_2Cl_2, [ppm])$ : ABMX<sub>2</sub>Y spin system:  $\delta(P_A) = -322.9$ ,  $\delta(P_B) =$ -320.2,  $\delta(P_M) = 30.4$ ,  $\delta(P_X) = 165.5$ ,  $\delta(P_Y) = 166.8$ ,  ${}^{1}J(P_AP_B) =$  $-189.8 \text{ Hz}, {}^{1}J(P_{A}P_{X}) = -143.4 \text{ Hz}, {}^{1}J(P_{B}P_{X}) = -147.7 \text{ Hz}, {}^{1}J$  $(P_M P_X) = -245.1 \text{ Hz}, {}^2 J(P_A P_M) = 18.8 \text{ Hz}, {}^2 J(P_B P_M) = 18.0 \text{ Hz};$ elemental analysis for C<sub>12</sub>H<sub>36</sub>GaCl<sub>5</sub>P<sub>6</sub>N<sub>4</sub>Si<sub>5</sub>: calcd: C 17.8, H 4.5, N 6.9; found: C 17.1, H 4.4, N 5.5.

<sup>1</sup>H NMR (CD<sub>2</sub>Cl<sub>2</sub>, [ppm]):  $\delta$  = 0.37 (18H, s, H2), 0.75 (18H, s, H1); <sup>31</sup>P{<sup>1</sup>H} NMR (CD<sub>2</sub>Cl<sub>2</sub>, [ppm]): ABMOX<sub>2</sub> spin system:  $\delta$ (P<sub>A</sub>) = -318.3,  $\delta$ (P<sub>M</sub>) = 33.4,  $\delta$ (P<sub>O</sub>) = 144 ( $\Delta$  $\nu$ <sub>1/2</sub> = ~1200 Hz),  $\delta$ (P<sub>X</sub>) = 170.8, <sup>1</sup>J(P<sub>A</sub>P<sub>X</sub>) = -144.8 Hz, <sup>1</sup>J(P<sub>M</sub>P<sub>X</sub>) = -248.6 Hz, <sup>2</sup>J(P<sub>A</sub>P<sub>M</sub>) = 19.0 Hz. Compound 12[GaCl<sub>4</sub>] was independently synthesized by addition of GaCl<sub>3</sub> (18 mg, 0.10 mmol, 1.0 eq.) to a solution of 19[GaCl<sub>4</sub>] (64 mg, 0.10 mmol, 1.0 eq.) in CD<sub>2</sub>Cl<sub>2</sub> (1 mL). The obtained colorless solution was stirred for 30 min at ambient temperature and subsequently investigated by <sup>1</sup>H and <sup>31</sup>P{<sup>1</sup>H} NMR spectroscopy.

SiMe<sub>3</sub> SiMe<sub>3</sub> 
$$\square$$
 [Ga<sub>2</sub>Cl<sub>7</sub>]<sub>2</sub>

$$SiMe_3$$
 SiMe<sub>3</sub>  $\bullet$  = P
$$13[Ga2Cl7]2$$

m.p.: 164.5–167.5 °C (decomposition); Raman (250 mW, [cm<sup>-1</sup>]):  $\nu = 2962$  (10), 2898 (17), 1095 (10), 633 (13), 549 (100), 441 (11), 397 (17), 384 (10), 354 (10), 138 (81) the Raman measurement was hampered by strong fluorescence effects; IR (ATR, [cm<sup>-1</sup>]):  $\nu = 2956$  (vw), 2898 (vw), 1409 (w), 1257 (s), 994 (vs), 899 (m), 814 (vs), 759 (vw), 729 (w), 692 (vw), 637 (w), 544 (w), 490 (w), 441 (vw), 409 (w); <sup>1</sup>H NMR (CD<sub>2</sub>Cl<sub>2</sub>, [ppm]):  $\delta = 0.72$  (36H, s, CH<sub>3</sub>); <sup>13</sup>C{<sup>1</sup>H} NMR (CD<sub>2</sub>Cl<sub>2</sub>, [ppm]):  $\delta = 13.7$  (s, Si(CH<sub>3</sub>)<sub>3</sub>), the Si atom of the SiN<sub>4</sub>-moiety was not detected; <sup>15</sup>N NMR (CD<sub>2</sub>Cl<sub>2</sub>, [ppm]):  $\delta = 115$  (s); <sup>31</sup>P{<sup>1</sup>H} NMR (CD<sub>2</sub>Cl<sub>2</sub>, [ppm]):  $\delta = 13.7$  (s, Figure 1);  $\delta = 115$  (s); <sup>31</sup>P{<sup>1</sup>H} NMR (CD<sub>2</sub>Cl<sub>2</sub>, [ppm]):  $\delta = 13.7$  (s) NMR (CD<sub>2</sub>Cl<sub>2</sub>, [ppm]):  $\delta = 115$  (s); <sup>31</sup>P{<sup>1</sup>H} NMR (CD<sub>2</sub>Cl<sub>2</sub>, [ppm]):  $\delta = 13.7$  (s)  $\delta = 115$  (s); <sup>31</sup>P{<sup>1</sup>H} NMR (CD<sub>2</sub>Cl<sub>2</sub>, [ppm]):  $\delta = 13.7$  (s)  $\delta = 115$  (s); <sup>31</sup>P{<sup>1</sup>H} NMR (CD<sub>2</sub>Cl<sub>2</sub>, [ppm]):  $\delta = 13.7$  (s)  $\delta = 115$  (s); <sup>31</sup>P{<sup>1</sup>H} NMR (CD<sub>2</sub>Cl<sub>2</sub>, [ppm]):  $\delta = 13.7$  (s)  $\delta = 115$  (s); <sup>31</sup>P{<sup>1</sup>H} NMR (CD<sub>2</sub>Cl<sub>2</sub>, [ppm]):  $\delta = 13.7$  (s)  $\delta = 115$  (s); <sup>31</sup>P{<sup>1</sup>H} NMR (CD<sub>2</sub>Cl<sub>2</sub>, [ppm]):  $\delta = 13.7$  (s)  $\delta = 115$  (s); <sup>31</sup>P{<sup>1</sup>H} NMR (CD<sub>2</sub>Cl<sub>2</sub>, [ppm]):  $\delta = 13.7$  (s)  $\delta = 1$ 

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