Abstract: The pnictogenyl Group 13 compounds (Dipp,Nacnac)M[E(SiMe$_3$)$_3$]Cl and (Dipp,Nacnac)M(EH)$_2$ (Dipp,Nacnac = H$_2$C(Me)$_n$N(Ar)$_2$, Ar = Dipp = 2,6-iPr$_2$C$_6$H$_4$; M = Al, Ga, In; E = P, As) were successfully synthesized. The salt metathesis between (Dipp,Nacnac)MCl$_2$ and Li(E(SiMe$_3$)$_2$) only led to monosubstituted compounds (Dipp,Nacnac)M(E(SiMe$_3$)$_2$)Cl [E = P, M = Ga(1), In (2); E = As, M = Ga (3), In (4)], regardless of the stoichiometric ratios used. In contrast to the steric effect of the SiMe$_3$ groups in 1–4, the reactions of the corresponding halides with LiP$_2$DME (or KAsH$_3$) facilitated the dipnictogenide compounds (Dipp,Nacnac)M(EH)$_2$ (E = P, M = Al (5), Ga (6), In (7); E = As, M = Al (8), Ga (9)), avoiding the use of flammable and toxic PH and AsH$_3$ for their synthesis. The compounds 5–9 are the first examples of monomeric Group 13 dipnictogenides and disilanides in which the metal center is bound to two terminal PH$_2$ and AsH$_3$ groups, respectively. In contrast to the successful synthesis of the indium dipnictogenide (Dipp,Nacnac)In(PH$_3$)$_2$; the reaction of (Dipp,Nacnac)InCl$_3$ with KAsH$_2$ led to an indium mirror due to the instability of the target product.

Pnictogenyl-substituted Group 13 metal compounds have emerged as an important type of molecules, having enormous potential to be used in material science of for example, nanoparticles, optoelectronic layers, or semiconductors. Moreover, the highly reactive EH$_2$ groups (E = N, P, and As) probably enable further substitution to obtain heterobimetallic species containing M–E(H)–M’ moieties. Given the Lewis acidity of the Group 13 metal center and the Lewis basicity of the EH$_2$ groups, these types of complexes readily undergo oligomerization to give dimers (R$_2$AI$\text{NH}_3$)$_2$ [R = SiMe$_3$, N(SiMe$_3$)$_2$] and [P$\text{H}^+\text{M}^-$(NH$_3$)$_2$] (M = Al, Ga, Ph$^+$ = terphenyl ligands); trimers (R$_2$AI$\text{NH}_3$)$_3$ (R = CH$_3$, tBu), or pseudo oligomers Al[R$_2$Al$\text{NH}_3$]$_3$ (R = SiMe$_3$, N(SiMe$_3$)$_2$) (A, B, C, Scheme 1). Therefore, steric crowding is essential to isolate stable monomeric species with parent EH$_2$ substituents. In 2004, the first monomeric aluminum diamide supported by a β-diketiminato substituent was synthesized from the corresponding chloride derivative with NH$_3$ in the presence of an N-heterocyclic carbene (D, Scheme 1). Subsequently, the corresponding gallium diamide was obtained through a similar procedure. Nonetheless, no progress on its heavier analogues has been made until now. This is caused by the extreme flammability and toxicity of PH$_3$ and AsH$_3$, respectively. To date, only very few of the heavier pnictogenyl Group 13 compounds with parent EH$_2$ (E = P, As) substituents have been reported. The lithium phosphanylanilane Li(Al(Ph)$_3$)$_2$ is known as the first aluminum phosphane representing a convenient PH$_3$ transfer reagent although it is only stable in ethereal solution. In 2000, Driess et al. reported the synthesis of disobutylphosphanyl(phenyl)alane [Bu$_2$Al(Ph)$_3$], which gives in situ in THF a monomolecular Bu$_2$Al(THF)PH$_3$ species (E, Scheme 1). Power and co-workers reported phosphanyl alanes and -gallanes ring compounds stabilized by bulky terphenyl ligands (F, Scheme 1). Previoulsy, our group was able to obtain aluminum and gallium monophosphanide complexes [(CO)$_5$W]$\text{PH}_2$NMe$_2$(NMe$_3$) (M = Al, Ga), which were prepared through H$_2$ elimination between Me$_3$N-MeH$_2$ (M = Al, Ga) and PH$_2$W[CO]$_5$(G, Scheme 1). Similarly, versatile oligomers can be obtained by using different conditions and different Lewis bases during the synthesis. Moreover, lithium arsanylanilane Li(Al(AsH)$_3$)$_2$ was also synthesized by the reaction of LiAlH$_4$ with AsH$_3$ In general, far less research has been made in arsanyl (AsH$_3$) chemistry because of its toxicity, light sensitivity, and instability. In addition to the aluminum and gallium diamide compounds (D, Scheme 1), the heavier analogues of Group 13 metal dipnictogenides are still unknown. Moreover, the preparation of pnictogenyl Group 13 compounds with an EH$_2$ moiety generally requires the pnictogen hydrogen compound-eEH$_2$, which, in addition to their flammability and toxicity, is invalid for stoichiometrically controlled reactions. Therefore, the quest for the stabilization of heavier Group 13 metal dipnictogenide species of type H is still open, and an appropriate
novel synthetic route should be developed. In 1993, Cowley prepared the base-stabilized phosphanyl- and arsanylalanes monomers Me₃N·AlH₂Mes(EMes=2,4,6-Me₃-C₆H₃) by metathesis reactions.[14] More recently, we reported on the synthesis of Lewis base-stabilized pnictogenylboranes based on metathesis,[15] demonstrating the feasibility of this type of reaction in the preparation of pnictogenyl metal compounds. In addition, the Lewis base-stabilized parent phosphanyl- and arsanylboranes H₂EBH₂·Me₃N (E=P, As) also were obtained by the alcoholysis of the corresponding (SiMe₃)₂EBH₂·Me₃N with methanol,[15g] which seems to be an alternative synthetic route. Herein, we report on the synthesis and characterization of first Group 13 metal diphosphanide and diarsanide compounds through salt metathesis, in which the sterically bulky β-diketiminate ligand was employed for the stabilization of the target products.

Initially, we explored the approach of Group 13 metal dipnicogenide compounds with SiMe₃ substituents. Treatments of (Dipp₂Nacnac)MCl₂ (Dipp₂Nacnac=HC[C(Me)N(Ar)]₂, Ar=2,6-iPr₂C₆H₃; M=Ga, In) with lithium bis(trimethylsilyl)phosphane [LiP(SiMe₃)₂] or arsanide [LiAs(SiMe₃)₂] in toluene only gave the monosubstituted compounds 1–4, respectively (Scheme 2). In 1–4, one chlorine atom has been replaced by a P(SiMe₃)₂ or As(SiMe₃)₂ group. However, even by using an excess amount of the lithium pnictogenide reagents, it was not possible to substitute both chlorine atoms, which is attributed to the steric hindrance of the bulky β-diketiminate ligand as well as the size of trimethylsilyl groups. The compounds 1–4 are well soluble, for example in toluene. In the ¹H NMR spectra of 1–4, characteristic resonances of the β-diketiminate ligand are observed, for example, the singlet at δ = 4.97 ppm for 1 [2: 4.84, 3: 4.93, and 4: 4.81 ppm] corresponds to the γ-H in each compound. In addition, the ¹H NMR spectra of 1 and 2 also show a doublet at δ = 0.09 and 0.02 ppm, respectively, with a 3 JPH coupling constant of 4.6 Hz, corresponding to the trimethylsilyl groups. In comparison, the singlets assigned to the trimethylsilyl groups in the ¹H NMR spectra of 3 and 4 were observed at δ = 0.10 and 0.07 ppm, respectively. Moreover, the ³¹P{¹H} NMR spectra of 1 and 2 show a singlet at δ = −255.0 and −252.9 ppm, respectively.

Single-crystal X-ray diffraction was carried out to confirm the structures of 1–4. All four compounds crystallize in the monoclinic space group P2₁/m, the molecular structure of 1 is depicted in Figure 1. The metal center is in a tetrahedral coordination mode binding to a β-diketiminate ligand, a chlorine atom as well as a bis(trimethylsilyl)phosphane or arsanide group, respectively. The N-M-N angles (M=Ga, In) in 1–4 were determined to 96.03(13) (1) and 95.95(9)° (3) for the gallium

![Scheme 1](image1)

**Scheme 1.** Selected examples of Group 13 metal pnictogenide compounds with EH₂ (E=N, P) substituents.

![Scheme 2](image2)

**Scheme 2.** Synthesis of compounds 1–4. Yields are given in parenthesis.

![Figure 1](image3)

**Figure 1.** Molecular structure of 1 with thermal ellipsoids at 30% probability level. Carbon-bound hydrogen atoms are omitted for clarity.
phosphinanes 5–7 were isolated in yields of 32, 34.3, and 1%, respectively. The quite low yield of isolated material for 7 is imputed to the extreme photosensitivity of indium compounds because the solution of 7 immediately became turbid upon exposure to light.

Compounds 5–7 are air and moisture sensitive and soluble in almost all common organic solvents. They were fully characterized by multinuclear NMR spectroscopy and X-ray diffraction. In the $^1$H NMR spectra of 5–7, typical resonances of $\beta$-diketiminate backbones are observed. The resonances of $\gamma$-H are detected as a singlet at $\delta = 4.97$ (5), 4.83 (6) and 4.75 ppm (7), which are comparable to those of (Dipp$_2$Nacnac)Al(NH$_2$)$_2$ ($\delta = 4.88$ ppm)\(^{[25]}\) and (Dipp$_2$Nacnac)Ga(NH$_2$)$_2$ ($\delta = 4.76$ ppm).\(^{[38]}\) Moreover, the $^1$H NMR spectrum of 5 shows a doublet at $\delta = 0.74$ ppm ($J_{\text{HH}} = 172.0$ Hz), which was assigned to the PH$_2$ moiety, whereas in the spectra of 6 and 7, the signal splits into a doublet of doublets at $\delta = 1.03$ ppm ($J_{\text{HH}} = 175.7, J_{\text{HH}} = 5.1$ Hz) for 6 and 0.90 ppm ($J_{\text{HH}} = 168.0, J_{\text{HH}} = 4.6$ Hz) for 7 (Table 1).

Table 1. Resonances of the PH$_2$ groups in $^1$H NMR spectra of compounds 5–7.

<table>
<thead>
<tr>
<th>Complex</th>
<th>$\delta$ (ppm)</th>
<th>multiplicity</th>
<th>$J_{\text{HH}}$ (Hz)</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>0.74</td>
<td>d</td>
<td>172.0</td>
</tr>
<tr>
<td>6</td>
<td>1.03</td>
<td>dd</td>
<td>157.7</td>
</tr>
<tr>
<td>7</td>
<td>0.90</td>
<td>dd</td>
<td>168.0</td>
</tr>
</tbody>
</table>

The AB spin systems in 6 and 7 are a result of the nonplanarity of the C$_x$N$_y$M ring, because the gallium and indium atoms are relatively oversized. The same coupling constants also were observed in the $^{31}$P NMR spectra. In contrast, singlets at $\delta = -273.4$ (5), -262.1 (6), and -285.5 ppm (7) were detected in the $^{119}$P(H) NMR spectra, falling in the range of the typical resonances of terminal PH$_2$ groups.\(^{[14,10,12a]}\) In addition, the LIFDI mass spectra of 5 and 6 show the most intense peak at m/z 477.3 and 519.4, respectively, assigned to the ionic fragment of [M$^+$–PH$_2$]. However, mass spectroscopy could not be used for the characterization of 7 because of its light and air sensitivity.

The structures of 5–7 were further determined by single-crystal X-ray diffraction. Compounds 5 and 6 crystallize in the triclinic space group $P\bar{1}$, whereas 7 crystallizes in the monoclinic space group $P2_1/n$. The molecular structure of 5 is depicted in Figure 2 and those of 6 and 7 are included in the Supporting Information. These three compounds are isostructural because their metal centers adopt a distorted-tetrahedral geometry in the coordination with two phosphorus and two nitrogen atoms, respectively. The phosphorus atoms adopt a pyramidal geometry with different orientations. The N-M-N angles ($M =$ Al, Ga, In) decrease following the order of Group 13 as 97.47(5) (5), 95.81(6) (6), and 89.10(6) (7). This indicates the nonplanarity of the C$_x$N$_y$M rings ($M =$ Al, Ga, In), which is in agreement with the coupling in the $^{19}$P NMR spectra. In contrast, the P-M-P angles slightly increase in the sequence of 115.26(2) (5), 118.18(19) (6), and 122.55(2) (7). In compound 5, the Al–P bond lengths are 2.3474(5) and 2.3718(5) Å, thus being comparable to those containing parent PH$_2$ units in [(CO)$_2$W(H$_2$)PAH$_2$(NMe$_2$)$_2$] ($n = 1$, 2.3671(1); $n = 2$, 2.4323(2)Å),\(^{[11,21]}\) [(CO)$_2$Cr(H$_2$)PAH$_2$(NMe$_2$)] (2.3831(1) Å),\(^{[22]}\) [(CO)$_2$WPH$_2$(NMe$_2$)] (2.3868(6) Å),\(^{[12a]}\) and [(ArAl)$_2$(μ-PH$_2$)$_2$(μ-PH)PH$_2$] [Ar$'$ = C$_6$H$_5$-2,6(C$_6$H$_5$-2,4,6-Me$_3$)] (2.3782(2) Å).\(^{[34]}\) The Ga–P bond distances of 6 (2.3286(5) and 2.3532(5) Å) are close to those observed in [(CO)$_2$W(H$_2$)PAH$_2$(NMe$_2$)] (2.3490(2) Å)\(^{[11]}\) and 1 (2.3310(9) Å), whereas 7 has slightly longer Ga–P bonds (2.5255(7) and 2.5052(6) Å) compared with that of 2 (2.4808(8) Å).

Furthermore, it was of interest if this synthetic procedure could also be used for the synthesis of the more sensitive bis-AsH$_2$-substituted derivatives. The treatment of (Dipp$_2$Nacnac)MX$_2$ ($X = I$, $M =$ Al; $X = Cl$, $M =$ Ga)\(^{[16,19]}\) with two equivalents of KAsH$_2$ resulted in the new aluminum and gallium diarsanide complexes 8 and 9 (Scheme 3). However, the
reaction of (Dipp,Nacnac)InCl$_3$ with KAsH$_2$ led to an indium mirror as well as [Dipp,Nacnac]H as the only product detectable by $^1$H NMR spectroscopy [24] due to the high sensitivity, even though the compounds were handled in the absence of light. Compound 9 is also quite unstable in solution to give [Dipp,Nacnac]H, whereas the aluminum derivative 8 shows a higher stability. Similarly, the $^1$H NMR spectra of 8 and 9 display characteristic signals for the β-diketiminate ligands such as the singlets for $\gamma$-H at $\delta = 4.99$ and 4.86 ppm and the septets for CHMe$_2$ at $\delta = 3.43$ and 3.44 ppm, respectively. In addition, the resonances for AsH$_3$ groups are observed as a singlet at $\delta = 0.28$ and 0.65 ppm, respectively. In the LiFDI mass spectrum of 8, the peak of the [M$^-$−AsH$_3$] fragment is observed at m/z 521.2, whereas the EI mass spectrum of 9 shows not only the [M$^-$−AsH$_3$] but also the [M$^-$−2AsH$_3$] fragments at m/z 563.1(95) and 487.2(22), respectively.

Compound 8 was further characterized by single crystal X-ray diffraction and represents the first monomeric aluminum diarsanide that has been structurally characterized so far (Figure 3). Compound 8 crystallizes in the monoclinic space group $P2_1/n$, in which the fourfold-coordinated aluminum center is bound to a β-diketiminate ligand and two AsH$_3$ substituents. Both arsenic atoms adopt a pyramidal geometry, indicating the existence of the lone pairs of electrons. The N(1)-Al(1)-N(2) angle in 8 (97.77(4)°) is similar to those of 5 (97.47(5)°) and (Dipp,Nacnac)Al[NH$_2$]$_2$ (95.71(4)°) [25]. The average As−Al bond length of 2.474(3) Å in 8 is comparable to those observed in lithium arsanylalanate [Li(DME)]$_2$[(AlH$_3$As)$_2$Li(DME)] (DME = 1,2-dimethoxyethane) (av. 2.472(3) Å) [26] whereas they are longer than the corresponding bond in 3 (2.4196(4) Å).

Computational DFT studies (see the Supplementary Information) indicate that the reactions leading to 1–4 and 5–9 are exergonic and that the formation of a solid salt is the driving force for the reaction. The computed thermodynamic data indicate that the stability of Group 13 metal diphosphanides and diarsanides decreases in the order Al > Ga > In. The processes of the decomposition of In-containing derivatives into indium metal with the formation of [Dipp,Nacnac]H, E$_0$ and H$_3$ as by-products are exergonic at room temperature by 10 and 53 kJ mol$^{-1}$ for E = P or As, respectively, which is in agreement with the experimentally observed low stability of 7 and the absence of its diarsanide analog.

The analysis of the electronic structure reveals that the HOMOs of 5–9 are essentially lone pairs of electrons located at pnicotogenyl groups. Thus, 5–9 can potentially serve as bidentate Lewis bases. However, the large E-M-E angles (115–123°, see above), which are unfavorable for the formation of 4-membered cycles, suggest that such compounds could be rather bridging than chelating ligands.

In summary, we provided a smooth synthetic route to metal phosphonide and arsenide compounds through salt-metathesis reactions and a series of β-diketiminate ligand-stabilized pnicoteryl Group 13 complexes (Dipp,Nacnac)M[E(SiMe$_3$)$_2$]Cl (E = P, As; M = Ga, In) and (Dipp,Nacnac)M(EH)$_3$ (E = P, M = Al, Ga; In; E = As, M = Al, Ga) have been synthesized. The utilization of alkali-metal pnicnides (LiE(SiMe$_3$)$_2$, LiPH$_2$, and KAsH$_2$) avoids the use of flammable and toxic PH$_3$ and AsH$_3$ for their synthesis. Due to steric hindrance, reactions of LiE(SiMe$_3$)$_2$ (E = P and As) with the corresponding metal chlorides only led to monosubstituted products 1–4. When using LiPH$_2$-DME and KAsH$_2$ as starting materials, the monomeric Group 13 diphosphanides and diarsanides 5–9 were obtained. For the first time, these complexes contain two terminal PH$_3$ and AsH$_3$ groups, respectively. Notably, the reaction of (Dipp,Nacnac)InCl$_3$ with KAsH$_2$ led to an indium mirror and [(Dipp,Nacnac)H] due to the intrinsic instability, implying the high photolability of such species. The good solubility and stability of aluminum diphosphanide and diarsenide compounds promise an interesting subsequent reactivity pattern, which is in the focus of current investigations. Moreover, the rather low stability of the M−E bonds might result in potential PH$_3$ and AsH$_2$ transfer reagents.

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Conflict of interest

The authors declare no conflict of interest.

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