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P(v)-bis(amidophenolate) ligand cooperation: stoichiometric C=O-bond cleavage in aldehydes and ketones<sup>†</sup>

Simon B. H. Karnbrock, Christopher Golz and Manuel Alcarazo 🝺 \*

The cooperation between a geometrically constrained, highly electrophilic phosphorus(v) center, and an electronically rich tetradentate bis(amidophenolate) ligand enables the cleavage of the C=O bond from typical aldehydes and ketones delivering iminio phosphoramidate species. The amphiphilic nature of these products, which is demonstrated through their reaction with typical Lewis acids and bases, enables their use as a mild source of silylium cations from silanes, allowing the selective reductive coupling of aldehydes to ethers under catalytic conditions.

The design of catalysts that strategically make use of metal-ligand cooperativity as a tool to facilitate the bond-breaking/forming steps at the substrate during a catalytic cycle has become increasingly popular. Among other advantages, these catalysts often serve as environmentally benign alternatives to previously developed systems in which a possible active role of the ligand was not considered.<sup>1</sup> However, and despite that undeniable impact, the exploitation of cooperativity in homogeneous catalysis has been mostly limited to transition metal-mediated processes; the transfer of such designing principles to p-block element-based catalysis is, in comparison, underdeveloped.<sup>2</sup> Remarkable exceptions are Radosevich catalytic hydrogenations and hydroborations using geometrically distorted phosphoramidites and phosphorus triamides,<sup>3</sup> the hydroboration of aldehydes and ketones mediated by N-heterocyclic germylenes,<sup>4</sup> and the digallane-catalysed hydroamination of alkynes introduced by Fedushkin,<sup>5</sup> among others.<sup>6</sup>

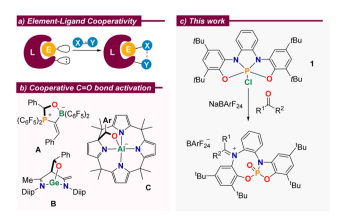
Cooperative behaviour between p-block elements and their surrounding ligands is more commonly found in stoichiometric transformations, which certainly have reduced synthetic utility, but serve to explore non-obvious modes of substrate activation and provide models for the development of new catalytic processes.<sup>2</sup> In this regard, and due to their ubiquity,

Institute of Organic and Biomolecular Chemistry, University of Göttingen,

E-mail: manuel.alcarazo@chemie.uni-goettingen.de

the activation of carbonyl moieties is very appealing. Numerous reports evidence that aldehydes or ketones readily bind to amphiphilic p-block complexes with concomitant cleavage of the C–O  $\pi$ -bond. For example, Erker described the reaction of a geminal alkylidene-bridged phosphine/borane Lewis pair with benzaldehyde to deliver compound **A** (Scheme 1b);<sup>7</sup> Wu, Liu and Zhao demonstrated that benzaldehyde cooperatively binds to Driess' N-heterocyclic germylene<sup>4</sup> to yield the corresponding adduct **B**,<sup>8</sup> and Greb and co-workers assessed the reactivity of a square planar aluminate towards aldehydes. Remarkably, the formation of adduct **C** is reversible and controllable by external stimuli.<sup>6c</sup>

We recently showed that the embedment of a highly electrophilic P(v) centre into an electron-rich, redox non-innocent, tetradentate bis(amidophenolato) scaffold results in phosphorane **1**. Once activated by chloride abstraction, this species is able to catalyse the disproportionation of 1,2-diphenylhydrazine through a mechanism that makes use of the electron reservoir character of the ligand.<sup>9</sup> These structural features, namely strong Lewis acidity at the central P(v)-centre and a high-lying occupied molecular orbital in spatial proximity, qualifies the



Scheme 1 (a) Concept of element–ligand cooperativity; (b) selected literature examples; (c) C=O bond cleavage in aldehydes and ketones.

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system for further modes of element–ligand cooperativity.<sup>2</sup> Herein, we report our studies on the reaction of **1** towards aldehydes and ketones of different structure, the evaluation of the reactivity of the iminiophosphoramidate products obtained, and their subsequent use as promotors for the reductive ether-ification of aldehydes.

At the outset of our study, chlorophosphorane  $1^{9a}$  was treated with acetone in the presence of Na[B(C<sub>6</sub>H<sub>3</sub>(CF<sub>3</sub>)<sub>2</sub>)<sub>4</sub>] (NaBArF<sub>24</sub>) as a chloride anion abstractor. The <sup>1</sup>H NMR spectrum of the crude reaction mixture clearly indicated loss of the original  $C_{\rm S}$ -symmetry of **1** and the splitting of the acetone methyl groups into two singlets at 2.67 and 2.54 ppm, which suggested the involvement of the bis(amidophenolate) side-arms in the transformation. Moreover, the <sup>31</sup>P NMR spectrum of the same reaction depicted a well-defined signal at -6.1 ppm, indicating the formation of a P(v)=O moiety.<sup>10</sup> Finally, compound **2** was isolated in a good yield by simple extraction from the reaction mixture with CH<sub>2</sub>Cl<sub>2</sub> and subsequent washing with pentane; its connectivity as an iminiophosphoramidate was unambiguously determined by X-ray crystallography (Scheme 2).

This reaction is not limited to acetone. Other aliphatic and aromatic ketones such as cyclopentanone, cyclohexanone and fluorenone were also found to be suitable substrates 3-5; but unfortunately, non-symmetric ketones invariably delivered non-separable E/Z-product mixtures. 1-Adamantyl carbaldehyde also

a) BArF<sub>24</sub> ∕<sup>t</sup>Bu NaBArF<sub>24</sub> <sup>t</sup>Bu <sup>t</sup>Bu <sup>t</sup>Bu **2** (76%) 1 BArF<sub>24</sub>  $BArF_{24}^{-}$ <sup>t</sup>Bu <sup>t</sup>Bu Bu **3** (62%) Bu 4 (73%) BArF<sub>24</sub> BArF<sub>24</sub> <sup>t</sup>Bu <sup>t</sup>Bı. <sup>t</sup>Bu t<sub>R</sub> <sup>t</sup>Bu **5** (92%) <sup>t</sup>Bu **6** (77%) c b)

**Scheme 2** (a) Reaction of ketones and aldehydes with chlorophosphorane **1**; (b) and (c) solid-state molecular structures of **2** (left) and **6** (right). Hydrogen atoms, BArF anions and solvent molecules omitted for clarity. Ellipsoids are set at the 50% probability level.<sup>11</sup>

delivers a stable adduct 6, which was isolated as a pure (*Z*)-isomer after recrystallization from hot toluene, but the iminiophosphoramidates derived from less sterically demanding aldehydes were all prone to decompose slowly.

A closer look into the X-ray structures of the carbonyl adducts 2-6 is quite informative (Scheme 2b and c for compounds 2 and 6, and Fig. S2-S4 for 3-5, ESI<sup>+</sup>). The C-N bond lengths are in all adducts within the typical range for iminium moieties,<sup>12</sup> and only vary slightly depending on the substituents at the carbon atom. Thus, the shortest C-N distance is observed for the aldiminium-derivative 6 (N1-C35; 1.297(2) Å) while the longest is found for the  $\pi$ -conjugated fluorenone derivative 5 (N1– C35; 1.3206(13) Å). A recurring feature observed throughout all the molecular structures is the close spatial proximity between the iminium carbon C35 and O1, with values ranging from 2.7028(14) Å in 4 to 3.0263(13) Å in 5, all shorter than the sum of their van der Waals radii.<sup>13</sup> This is probably a consequence of the sum of two factors; namely, the structural constraints imposed by the rigid bis(amidophenolato) scaffold, and the polarity match between the electrophilic iminium carbon and the Lewis basic oxygen atom from the P(v) = O moiety.

In order to gain a deeper understanding of the mechanism leading to the formation of 2-6, electronic structure calculations were carried out at the B3LYP-D3(BJ)/def2-TZVP//PBE-D3(BJ)/def-2SVP level of theory to evaluate the feasibility of the critical steps.<sup>14</sup> Fig. 1a shows the computed Gibbs free energies for intermediates and transition states using acetone as a model substrate. Once the chloride anion has been replaced by acetone Int I, an amino moiety of the ligand attacks the electrophilic carbonyl carbon of the coordinated acetone delivering oxazaphosphetane Int II via transition state TSI. This cooperative pathway requires a moderate Gibbs activation energy ( $\Delta G^{\ddagger}$ ) of 11.4 kcal mol<sup>-1</sup>. From Int I, cleavage of the C-O σ-bond proceeds with an even lower barrier (4.8 kcal mol<sup>-1</sup>) through **TS2**, which resembles the [2+2]cycloreversion step of an aza-Wittig reaction.<sup>15</sup> The formation of a strong P–O bond in Int III offers a qualitative explanation for the exergonic nature of this transformation.

The analysis of electron flow for this reaction pathway using intrinsic bond orbitals (IBOs) beautifully pinpoints the phosphorusligand cooperative action.<sup>16</sup> On the way to **Int II** the nitrogen lone pair attacks the carbonyl group to form the  $\sigma$ (N–C) bond of the oxazaphosphetane moiety, while simultaneously, the initial  $\pi$ (CO) bond is transformed into an oxygen-centred lone pair, thus breaking the  $\pi$ (CO) interaction (Fig. 1b). Subsequently, the dative  $\sigma$ (N–P) bond in **Int II** slowly breaks, giving rise to the  $\pi$ (N–C) bond in 2, while at the same time the  $\sigma$ (C–O) bond in **Int II** dissipates giving rise to the formation of the  $\pi$ (P=O) interaction (Fig. 1c).<sup>16</sup>

Based on the spatial proximity of non-quenched Lewis acid and base sites in 2–6, we speculated that these species might exhibit amphiphilic reactivity. Hence, using 6 as a model substrate, we initially treated this compound with an archetypical Lewis base, 4-dimethylaminopyridine (DMAP). The expected attack at the iminium functionality took place and adduct 7 could be isolated by crystallization (Scheme 3a). Contrarily, the reaction of 6 with GaCl<sub>3</sub> proved to be more complex. Instead of mere coordination of the Lewis acidic Ga-centre to O1,

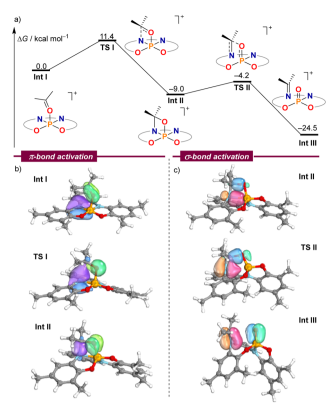
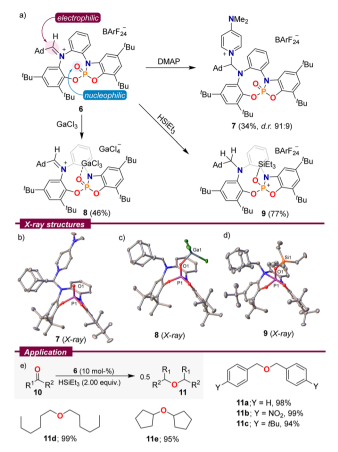


Fig. 1 (a) DFT calculations at the B3LYP-D3(BJ)/def2-TZVP//PBE-D3(BJ)/ def2-SVP level on the mechanism of the C=O bond cleavage. (b) and (c) Selected IBO plots through the reaction coordinate.

degradation of the BArF<sub>24</sub> counterion was observed, and 8 crystallized as a [GaCl<sub>4</sub>]<sup>-</sup> salt. By employing three equivalents of GaCl<sub>3</sub>, we were able to obtain 8 in 46% yield as an orange crystalline solid. As expected, by formation of 7 the C35–N1 bond distance elongates from 1.297(2) Å in 6 to 1.487(2) Å, while in 8 it is the P1– O1 distance that gets elongated as a result of coordination to the Lewis acid, from 1.4545(14) Å in 6 to 1.4878(10) Å in 8. It is worth noting that the weak pyramidalization observed at the Ga centre ( $\sum_{\angle Ga} = 345.8^{\circ}$ ) is indicative of a limited donor ability of 6.<sup>17</sup>

Interestingly, both reactivity modes can be observed upon the addition of triethyl silane to a CDCl<sub>3</sub> solution of **6**. The aldiminium resonance at 8.38 ppm in the <sup>1</sup>H NMR spectrum slowly disappears while two new signals at 3.66 and 3.25 ppm with a geminal  ${}^{2}J_{HH} = 13.4$  Hz arise, in agreement with the reduction of the iminium functionality. Simultaneously, a doublet in the  ${}^{29}$ Si-INEPT spectrum resulting from  ${}^{2}J_{PSi}$  coupling verified binding of the silylium fragment to the P==O moiety. On a preparative scale, **9** was isolated in a good yield of 77% and single crystals suitable for X-ray diffraction were grown by vapor diffusion of pentane into a toluene solution of **9**. The solid-state structure confirms the proposed connectivity. Compound **9** depicts an exceptionally long O1–Si1 bond (1.7301(11) Å),<sup>15,18</sup> which is indicative of the low basicity of the P==O fragment in **6**.

As a result of that weak O–Si interaction, it was rationalized that **9** might serve as a mild source of triethylsilylium cations, and therefore, it could be used as a pre-catalyst for transformations in which that cation plays an active role. Hence, we



Scheme 3 (a) Reactivity of 6; (b)–(d) solid-state molecular structures of 7-9. Hydrogen atoms, anions and solvent molecules in the solid-state structures were omitted. Ellipsoids are set at 50% probability;<sup>11</sup> (e) reductive etherification of aldehydes and ketones. Yields were determined by quantitative NMR spectroscopy using hexamethylbenzene as an internal standard.

evaluated the ability of **6** to catalyse the reductive etherification of aldehydes and ketones to ethers using  $Et_3SiH$  as a reductant.<sup>19</sup> The reaction proved general, and benzaldehyde derivatives of different electronic properties **10a–c**, aliphatic aldehydes **10d**, and even cyclopentanone were selectively transformed into the corresponding symmetrical ethers in excellent yields (Scheme 3e).

In summary, we report the cleavage of the C=O bond in aldehydes and ketones through the cooperation between a central P(v)-atom and its surrounding tetradentate ligand. This behaviour can be rationalized considering the high electrophilicity at the P-centre, which is a consequence of the structural constraints imposed by the multidentate bis(amidophenolate) scaffold, and the electron richness of that ligand. A preliminary application of the iminiophosphoramidate species such obtained as promotors for the reductive etherification of aldehydes and ketones is also described.

S. B. H. K. and M. A. conceived the project and designed the experiments. S. B. H. K. performed the experiments, analysed the results and carried out the computational studies. C. G. did the X-ray crystallographic studies. All authors discussed the results and S. B. H. K. and M. A. prepared the manuscript.

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## Conflicts of interest

There are no conflicts to declare.

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