

Exploring the Reactivity of Carbon(0)/Borane-Based Frustrated Lewis Pairs**

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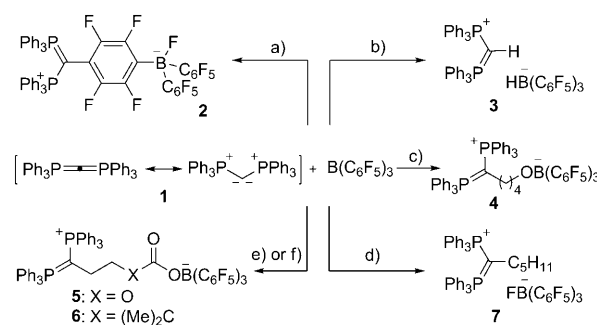
Dedicated to Professor Rosario Fernández

Since the unveiling of the concept of frustrated Lewis pairs (FLPs) by Stephan et al.,^[1] the chemistry of these systems has flourished. Arguably their most attractive application has been the heterolytic activation of H₂,^[2] and the subsequent development of metal-free hydrogenation catalysis directly employing dihydrogen^[3] rather than a surrogate.^[4] Although several other bonds such as C–O,^[5] C–H,^[6] B–H,^[7] S–S,^[8] and C–C,^[9] have also been cleaved by using FLPs, these systems largely rely on P- or N-based Lewis bases combined with a polyfluorinated borane.^[10] The sole exceptions are the sterically crowded carbene 1,3-di-*tert*-butyl-1,3-imidazol-2-ylidene (IrBu) in combination with B(C₆F₅)₃, a pair that contains a C-derived base,^[11] and the use of Al(C₆F₅)₃ instead of a borane as Lewis acid.^[12] Clearly, extension of the FLP concept to include currently unexplored partners is desirable, as it may lead to the discovery of a range of interesting new applications.

In our research to broaden the range of bases that can be used in FLP chemistry, we were inspired by the computational investigations of Tonner and Frenking on the nature of carbodiphosphoranes.^[13,14] They proposed that these compounds should be considered to comprise two phosphine ligands coordinated to a central zero-valent carbon atom that retains its four valence electrons. This view has been subsequently confirmed experimentally by the work of Bertrand et al., Fürstner et al., and others.^[15]

The available information suggests that C⁰ compounds must be exceptionally good nucleophiles. In fact, the calculated proton affinity for carbodiphosphoranes surpasses the values reported for amines, phosphines, and even N-heterocyclic carbenes. It can be envisaged therefore that, if sufficiently sterically hindered, C⁰ compounds should be qualified to function as bases in the framework of FLP chemistry. Herein, in an attempt to address this issue, the pair hexaphenylcarbodiphosphorane (**1**)/B(C₆F₅)₃ is studied and its reactivity towards several small molecules evaluated.^[17]

Initially, we carried out the reaction of **1** with B(C₆F₅)₃ in toluene at room temperature. The NMR spectroscopic data of the obtained product suggested formation of **2** by nucleophilic attack at the *para* position of the pentafluorophenyl ring and trapping of the generated fluoride anion by the boron atom. X-ray crystallographic analysis later confirmed the structure of **2** (see Supporting Information).^[18] However, when the same reagents were mixed at –78 °C, NMR spectroscopy indicated no interaction between the partners, that is, “frustration”. Purging this stoichiometric mixture with H₂ resulted in formation of a white precipitate which could be isolated in 91 % yield. The NMR data support the formulation for this product as [(Ph₃P)₂CH][HB(C₆F₅)₃] (**3**; Scheme 1). The cation exhibits a ¹H signal at δ = 1.73 ppm with ²J(¹H, ³¹P) of 5.4 Hz and a ³¹P{¹H} resonance at δ = 21.3 ppm, while the



Scheme 1. Some reactions of FLP **1**/B(C₆F₅)₃. Reagents and conditions (yields): a) toluene, –78 °C → RT (74 %); b) H₂ 1 atm, toluene, –78 °C → RT (91 %); c) THF, –78 °C → RT (76 %); d) *n*C₅H₁₁F, toluene, –78 °C, quant.; e) ethylene carbonate, toluene, –78 °C → RT (84 %); f) 2,2-dimethyl-γ-butyrolactone, toluene, 78 °C → RT (71 %).

anion gives the expected ¹¹B signal of a borohydride at δ = –25.5 ppm with a ¹J(¹H, ¹¹B) of 100 Hz. Single crystals were obtained by slow diffusion of pentane into a solution of **3** in CH₂Cl₂, and X-ray structure analysis confirmed not only the proposed structure (Figure 1), but also the ability of **1**/B(C₆F₅)₃ to function as an FLP.

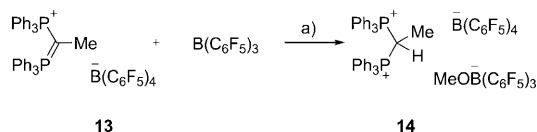
As expected for an FLP, **1**/B(C₆F₅)₃ in solution in THF resulted in ring opening of the ether to produce phosphonium borate **4**, also confirmed by X-ray analysis (see Supporting Information). This reactivity was extended to ethylene carbonate and the non-enolizable ester 3,3-dimethyl-γ-butyrolactone to produce zwitterionic species **5** and **6**, respectively. Moreover, addition of one equivalent of 1-fluoropentane to a suspension of **1**/B(C₆F₅)₃ at –78 °C generates [(Ph₃P)₂C-

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[**] We thank Prof. A. Fürstner for generous support and constant encouragement. The NMR spectroscopy and X-ray crystallography departments of our institute are also gratefully acknowledged for excellent support, as well as H. Bruns and R. Schinzel for the preparation of starting materials.

Supporting information for this article is available on the WWW under <http://dx.doi.org/10.1002/anie.201002119>.

O–H bond of methanol, something that **13** alone is not able to do (Scheme 3). In fact, **13** can be crystallized from MeOH without generating any protonated product.



Scheme 3. Cleavage of the O–H bond of methanol with the pair **13**/B(C₆F₅)₃. Reagents and conditions (yields): a) MeOH (1 equiv), toluene, RT, **14** (86%).

This reveals a degree of frustration that, albeit clearly diminished in comparison to the **1**/B(C₆F₅)₃ pair, is still remarkable considering the cationic nature of the employed base. Cationic Lewis acids like [CPh₃]⁺ have been studied in the context of FLPs;^[12] however, this is the first example of the counterintuitive use of a cationic base in this field. The formulation of compound **14** is supported by a ¹H NMR signal at δ = 1.95 ppm for the Me group with ³J(¹H, ³¹P) of 16.0 Hz and a ³J(¹H, ¹H) of 7.6 Hz, a ³¹P{¹H} resonance at δ = 28.6 ppm and two ¹¹B{¹H} signals at δ = –2.3 and –15.2 ppm. In addition, during attempts to crystallize **14**, crystals of **14**[B(C₆F₅)₄]₂ were obtained and its structure confirmed by X-ray analysis (Figure 3).

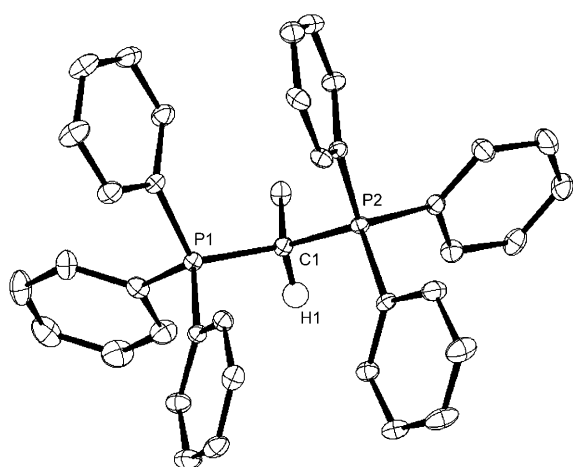


Figure 3. Molecular structure of **14**[B(C₆F₅)₄]₂ in the solid state. (B(C₆F₅)₄)₂ anions and hydrogen atoms, except that bonded to C1, were removed for clarity; ellipsoids set at 50% probability).^[16]

In conclusion, this study demonstrates that the combination of hexaphenylcarbodiphosphorane as carbon-based Lewis base and B(C₆F₅)₃ forms a novel frustrated Lewis pair. The system is not only able to achieve the classical H–H, C–O, and C–H bond cleavages, but also unprecedented activation of Si–H bonds and alkyl fluorides. The peculiar electronic situation of the C⁰ base makes the system unique: after the first protonation or alkylation, some degree of

frustration persists. The application of these FLPs in homogeneous catalysis is currently under investigation.

Received: April 9, 2010

Published online: July 2, 2010

Keywords: boranes · carbodiphosphoranes · donor–acceptor systems · frustrated Lewis pairs · hydrogen activation

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