

Tandem Frustrated Lewis Pair/Tris(pentafluorophenyl)borane-Catalyzed Deoxygenative Hydrosilylation of Carbon Dioxide

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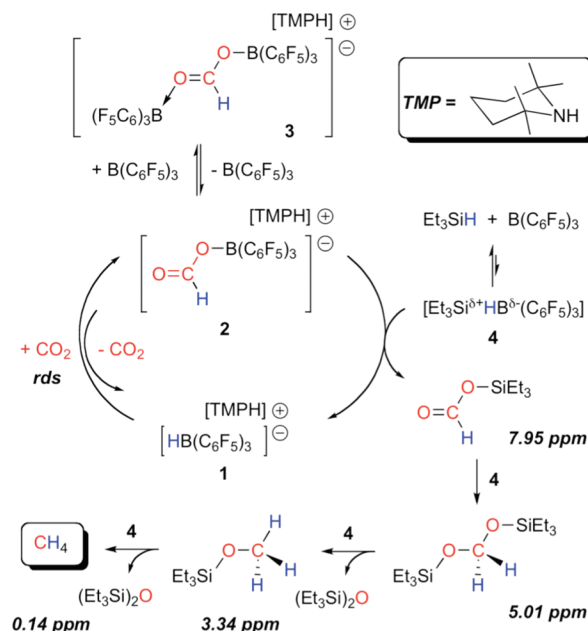
Abstract: The frustrated Lewis pair system consisting of 2 equiv of 2,2,6,6-tetramethylpiperidine (TMP) and tris(pentafluorophenyl)borane $[B(C_6F_5)_3]$ activates carbon dioxide to form a boratocarbamate–TMPH ion pair. In the presence of triethylsilane, this species is converted to a silyl carbamate and the known ion pair $[TMPH]^+[HB(C_6F_5)_3]^-$, which recently was shown to react with CO_2 via transfer of the hydride from the hydridoborate to form the formatoborate $[TMPH]^+[HC(O)OB(C_6F_5)_3]^-$. In the presence of extra $B(C_6F_5)_3$ (0.1–1.0 equiv) and excess triethylsilane, the formatoborate is rapidly hydrosilylated to form a formatosilane and regenerate $[TMPH]^+[HB(C_6F_5)_3]^-$. The formatosilane in turn is rapidly hydrosilylated by the $B(C_6F_5)_3/Et_3SiH$ system to CH_4 , with $(Et_3Si)_2O$ as the byproduct. At low $[Et_3SiH]$, intermediate CO_2 reduction products are observed; addition of more CO_2/Et_3SiH results in resumed hydrosilylation, indicating that this is a robust, living tandem catalytic system for the deoxygenative reduction of CO_2 to CH_4 .

The utilization of carbon dioxide as a sustainable and nontoxic C1 feedstock for the production of value-added chemical products such as carboxylic acids or fuels such as methanol and methane is of current interest.¹ The high thermodynamic stability of CO_2 necessitates its catalytic activation and coupling to a thermodynamic driver for efficient conversion. Transition-metal-based catalysts have played a dominant role in CO_2 conversion, but recently, an increasing number of organocatalytic CO_2 reduction schemes have emerged.² For example, N-heterocyclic carbenes (NHCs) reversibly form zwitterionic adducts $NHC \cdot CO_2$ that are considered key intermediates in the reductive deoxygenation of CO_2 using diphenylsilane as a sacrificial reducing agent, affording CH_3OH upon workup.³

In this context, activation of CO_2 by transition-metal-free “frustrated Lewis pairs” (FLPs)⁴ has led to the development of stoichiometric reductions of CO_2 to CH_3OH . Here, the FLPs form bridging carboxylate species⁵ that can accept hydrogen from ammonia borane⁶ or via a thermally driven, multistep self-reduction in which the key step is a reversible B–H bond addition of hydridotris(pentafluorophenyl)borate to one C=O double bond of CO_2 , affording the formatoborate anion $[HC(O)OB(C_6F_5)_3]^-$.⁷ Ultimately, hydrolysis of CH_3O-LA ($LA = BX_3$ or AlX_3) is required in order to obtain methanol.

Boron–hydrogen bond addition to CO_2 mediated by phosphonium or ammonium borate ion pairs formed via FLP hydrogen splitting thus offers a potential entry point into catalytic CO_2 fixation in the presence of a suitable reducing agent (oxygen acceptor). We have shown that perfluoroarylboranes are excellent catalysts for the reductive hydrosilylation of carbonyl functions⁸ and C–O bonds,⁹ a potentially useful reaction for subsequent steps in the reductive deoxygenation of CO_2 to CH_4 .

Scheme 1



The ammonium hydridoborate ion pair **1** formed by treatment of the FLP $B(C_6F_5)_3/2,2,6,6$ -tetramethylpiperidine (TMP) and hydrogen¹⁰ (32 mM, C_6D_5Br) reacted with CO_2 (2–4 atm) in the presence of Et_3SiH (18 equiv) at 56 °C to afford the previously reported⁷ formatoborate **2** exclusively (see Scheme 1).¹¹ The reaction was monitored by 1H and ^{19}F NMR spectroscopy, and integration versus an internal standard ($C_6H_5CF_3$, 9 mM) revealed that no Et_3SiH was consumed. Thus, although the formation of **2** is reversible,⁷ there does not appear to be sufficient free $B(C_6F_5)_3$ present under these conditions to activate silane for further reduction of **2**.

Accordingly, we carried out a reaction under identical conditions with an additional 1.0 equiv of $B(C_6F_5)_3$ (relative to **1**) present. This resulted in the immediate and complete conversion of **2** back into **1** at room temperature and the appearance of the products of CO_2 hydrosilylation. Further monitoring of the reaction by 1H and ^{19}F NMR spectroscopy at 56 °C showed that silane was gradually consumed and that CH_4 along with 2 equiv of $(Et_3Si)_2O$ were formed as the ultimate reaction products. Minor amounts of bis(triethylsilyl)acetal, $(Et_3SiO)_2CH_2$, (~10%) were also present. Interestingly, **1** and $B(C_6F_5)_3$ were the only boron-containing compounds detectable during the reaction but diminished in favor of a new species, **3**, upon complete silane consumption. At the same time, the characteristic signals of HCO_2SiEt_3 , $\{Et_3SiO\}_2CH_2$, and Et_3SiOCH_3 in C_6D_5Br became evident in the 1H NMR spectra. Upon addition of further silane equivalents and pressurization with fresh CO_2 , these partially reduced intermediates were depleted and methane formation resumed, indicating a “living” catalytic system.

Collectively, these observations suggest that the chemistry depicted in Scheme 1 is operative. Analysis of the ^{19}F NMR spectra indicated that the coordinated $\text{B}(\text{C}_6\text{F}_5)_3$ in compound **3** is labile in solution.¹² Thus, the equilibrium between **3** and $2/\text{B}(\text{C}_6\text{F}_5)_3$ is rapid and provides a source of free borane to activate the Et_3SiH present via **4**, as previously reported.⁸ The fact that neither **3** nor **2** was detected in the reaction mixture in the presence of silane implies that B–H bond addition of **1** to CO_2 is the rate-limiting step. Consequently, the overall rate of silane consumption showed a zeroth-order concentration dependence over four half-lives (~ 240 min) when the reaction was monitored at 56°C by ^1H NMR spectroscopy (see Figure S2 in the Supporting Information). As the silane concentration decreased, the concentration of CH_4 increased at a rate one-quarter that of silane consumption while **[1]** remained constant, as expected. Once **2** was generated, the reaction with $\text{B}(\text{C}_6\text{F}_5)_3$ -activated silane **4** was rapid and eventually produced CH_4 and $(\text{Et}_3\text{Si})_2\text{O}$ (as shown in Scheme 1) via well-documented $\text{B}(\text{C}_6\text{F}_5)_3$ -mediated transformations.¹³

In accord with this postulate, the reaction of a 2:1 mixture of **2** and $\text{B}(\text{C}_6\text{F}_5)_3$ (in equilibrium with **3**) with Et_3SiH (1.2 equiv vs **2**) at room temperature instantly afforded $(\text{Et}_3\text{SiO})_2\text{CH}_2$ (57%) along with starting material (42%) and trace amounts of $\text{Et}_3\text{SiOCH}_3$ and CH_4 , suggesting that the triethylsilylformate intermediate is highly reactive toward **4**. Therefore, its production from **2** is critical. Here, the involvement of a silylium cation to react with anionic **2** greatly enhances its conversion rate in comparison with that for the reduction of **2** with further equivalents of anionic **1**.⁷ In other words, silylium ion transfer to the formate moiety of **2** from **4** is Coulombically favored over hydride transfer from **1** and occurs under much milder conditions.⁷ The reaction of **4** with **2** gives highly reactive $\text{HCO}_2\text{SiEt}_3$ but also regenerates **1** for rate-limiting activation of CO_2 .

Compound **3** was generated separately by treatment of solutions of **2** with $\text{B}(\text{C}_6\text{F}_5)_3$ and was isolated by slow hexane diffusion into the reaction mixture at -30°C . Its solid-state structure was elucidated by single-crystal X-ray diffraction, and an ORTEP diagram is shown in Figure 1 along with selected metrical data implying that the negative charge is delocalized over the bridging formate and flanking borane fragments. This compound was proposed by Ashley et al.⁷ as an intermediate in the conversion of **2** to $[\text{H}_3\text{COB}(\text{C}_6\text{F}_5)_3]^-[\text{TMPPH}]^+$ that accepts hydride from **1**; here it serves as a reservoir of borane catalyst for hydrosilylation of **2** and subsequent intermediates.

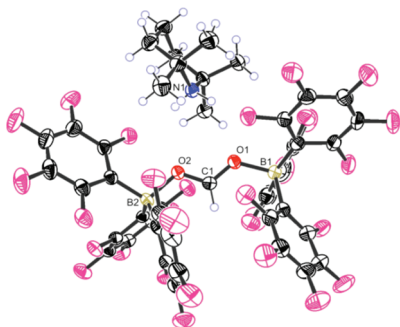
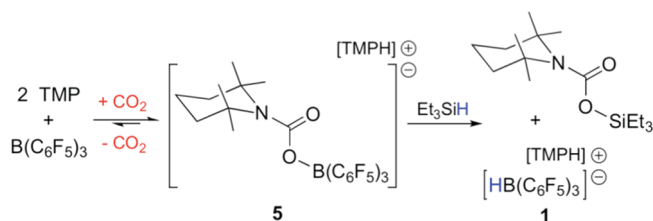


Figure 1. ORTEP plot for **3** (50% thermal displacement ellipsoids). Selected bond lengths (Å) and angles (deg): C1–O1, 1.256(3); C1–O2, 1.268(3); O1–B1, 1.587(3); O2–B2, 1.584(3); O1–C1–O2, 120.2(2); O1/C1–H, 119.9. The largest deviation from the least-squares plane through atoms B1, O1, C1, O2, and B2 is 0.015 Å for O1.

The above experiments were performed with isolated **1**, but this can be bypassed simply through the use of a 2:1 mixture of TMP/

Scheme 2



$\text{B}(\text{C}_6\text{F}_5)_3$ with CO_2 in the presence of silane (Scheme 2). In the absence of Et_3SiH , ion pair **5** was generated and could be characterized by X-ray crystallography.¹¹ As with other complexes of this type,^{5,6} CO_2 activation is reversible, and small quantities of free borane are accessible to added silane, which rapidly converts **5** into the triethylsilyl carbamate and **1**, which in turn becomes available for catalytic reduction of CO_2 to CH_4 as in Scheme 1.

In summary, ammonium borate **1** and $\text{B}(\text{C}_6\text{F}_5)_3$ act in tandem to catalytically convert CO_2 to CH_4 using triethylsilane as the reductant. The rate-limiting step involves transfer of hydride from **1** to CO_2 , suggesting that a more nucleophilic hydridoborate might improve the rate of conversion. However, use of the less Lewis acidic borane $\text{B}(4\text{-C}_6\text{F}_4\text{H})_3$ ¹⁴ resulted in a kinetic profile essentially identical to that obtained for $\text{B}(\text{C}_6\text{F}_5)_3$ (Figure S3). Ongoing work will explore a range of boranes, amines, and sacrificial reductants aimed at increasing the turnover frequencies on the basis of the mechanistic details uncovered through these detailed spectroscopic studies.

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Supporting Information Available: Crystallographic data for **3** and **5** (CIF), additional experimental and spectroscopic details, and complete ref 1a. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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- (11) See the Supporting Information for experimental details.
- (12) ^{19}F NMR spectra of mixtures of **2** and various amounts of $\text{B}(\text{C}_6\text{F}_5)_3$ in $\text{C}_6\text{D}_5\text{Br}$ exhibited broad resonances consistent with the presence of **2**, **3**, and free $\text{B}(\text{C}_6\text{F}_5)_3$ in rapid equilibrium.
- (13) This mechanistic scheme is similar in form to that proposed for a $\text{Zr}/\text{B}(\text{C}_6\text{F}_5)_3$ -based catalytic system for the slow hydrosilylation of CO_2 . See: Matsuo, T.; Kawaguchi, H. *J. Am. Chem. Soc.* **2006**, *128*, 12362–12363.
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