

Selective Reduction of CO₂ to CH₄ by Tandem Hydrosilylation with Mixed Al/B Catalysts

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Supporting Information

ABSTRACT: This contribution reports the first example of highly selective reduction of CO_2 into CH_4 via tandem hydrosilylation with mixed main-group organo-Lewis acid (LA) catalysts $[Al(C_6F_5)_3 + B(C_6F_5)_3]$ {[Al] + [B]}. As shown by this comprehensive experimental and computational study, in this unique tandem catalytic process, [Al] effectively mediates the first step of the overall reduction cycle, namely the fixation of CO_2 into HCOOSiEt₃ (1) via the LA-mediated



C=O activation, while [B] is incapable of promoting the same transformation. On the other hand, [B] is shown to be an excellent catalyst for the subsequent reduction steps 2–4, namely the hydrosilylation of the more basic intermediates [1 to $H_2C(OSiEt_3)_2$ (2) to $H_3COSiEt_3$ (3) and finally to CH_4] through the frustrated Lewis pair (FLP)-type Si–H activation. Hence, with the required combination of [Al] and [B], a highly selective hydrosilylative reduction of CO₂ system has been developed, achieving high CH_4 production yield up to 94%. The remarkably different catalytic behaviors between [Al] and [B] are attributed to the higher overall Lewis acidity of [Al] derived from two conflicting factors (electronic and steric effects), which renders the higher tendency of [Al] to form stable [Al]–substrate (intermediate) adducts with CO_2 as well as subsequent intermediates 1, 2, and 3. Overall, the roles of [Al] and [B] are not only complementary but also synergistic in the total reduction of CO_2 , which render both [Al]-mediated first reduction step and [B]-mediated subsequent steps catalytic.

INTRODUCTION

Achieving efficient direct and catalytic reduction of CO₂ into CH₄ will have significant impact on addressing two currently biggest issues facing humanity: global warming largely due to the rising level of the greenhouse gas CO₂ and increasing demand on clean energy such as solar energy and natural gas CH_4 . Although CO_2 is considered to be an ideal renewable C_1 feedstock for chemicals, materials, and fuels as it is renewable, abundant, nonflammable, and inexpensive,¹ it is a highly stable, inert molecule, so it has been a challenge to develop technologically feasible and economically competitive methods to convert CO₂ into fuels, especially high-energy density, deoxygenated fuels such as methane.² Catalytic hydrosilylation of CO₂, although with silanes as an expensive hydrogen source, is a thermodynamically favored process with formation of stronger Si–O bonds,³ when compared to the catalytic hydrogenation of CO₂.⁴ Currently, transition-metal (TM) catalysts based on metals such as Ru,⁵ Co,⁶ Rh,⁷ Ir,⁸ Ni,⁹ Cu,¹⁰ and main-group Zn^{11} have been utilized but achieved limited success in the catalytic hydrosilylation of CO₂ to lower oxidation state species with low selectivity for methane formation.²

An emerging approach to enhancing catalytic performance of the CO_2 hydrosilylation systems is to couple the strong organo-Lewis acid $B(C_6F_5)_{3}$,¹² which, when used alone, is incapable of reducing CO₂,¹³ with TM complexes of Zr,¹³ Pt, Pd,¹⁴ Re,¹⁵ and Sc^{16} for fixation of CO_2 (by TM complexes) and subsequent reduction of the more basic formate, aldehvde, and methanol equivalents to CH_4 (by the borane). More recently, there is emerging interest in the development of TMfree catalysis spotlighted by frustrated Lewis pairs (FLPs).¹⁷ O'Hare et al. reported CO₂ can be hydrogenated to CH₃OH with the 2,2,6,6-tetramethylpiperidine $(TMP)/B(C_6F_5)_3$ FLP system.¹⁸ Alternatively, Stephan¹⁹ et al. demonstrated the combination of $PMes_3/AlX_3$ (Mes = mesityl, X = Cl, Br) effects the reduction of CO₂ to CH₃OH with H₃NBH₃ as a hydrogen source. However, both of these transformations require a stoichiometric amount of the FLP reagents. In this regard, Fontaine et al. discovered an ambiphilic Lewis pair system with a less Lewis acidic catecholboryl unit which can promote the release of the reduced products from the catalyst and thus render the hydroboration of CO₂ catalytic.²⁰ Ever since, major experimental and computational efforts have been directed to the study of the CO_2 reduction utilizing the FLP chemistry.^{21,22} On the other hand, only a few examples of catalytic hydrosilylation of CO2 have surfaced in the literature by utilizing either the separate reactivity of Lewis acid (LA)/Lewis

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base (LB) or the combination of them. For instance, Ying et al. reported NHC-catalyzed CO₂ reduction with H₂SiPh₂,²³ while Piers utilized $B(C_6F_5)_3$ (in excess) and TMP for deoxygenative hydrosilylation of CO₂.²⁴ Müller et al. demonstrated that silyl cations are effective in promoting conversion of CO₂ into benzoic acid, formic acid, and methanol, albeit not in a catalytic fashion.²⁵ More recently, Wehmschulte et al. reported the LAcatalyzed hydrosilylation of CO₂ by cationic aluminum species AlR_2^+ (R = Et or OAr).²⁶ However, the detailed mechanism of such LA-catalyzed CO₂ reduction remains unclear, and thus further enhancing catalytic performance through modification of catalyst structure seems challenging. In this context, it is of great interest and importance to survey suitable main-group LA candidates for effective CO2 reduction. Inspired by the dual activation of CO2 and silane R3SiH {[SiH]} by the above overviewed TM or $TM/B(C_6F_5)_3$ systems, we envisioned that a combination of the more oxophilic and higher Lewis acidic $Al(C_6F_5)_3$ {[Al]},²⁷ which could be sufficiently potent to render CO_2 fixation into silvl formate, with the less oxophilic $B(C_6F_5)_3$ {[B]}, which favors FLP-type [SiH] activation for subsequent steps of reduction,²⁸ could serve as a main-group tandem FLP system that effectively converts CO₂ to CH₄. The tunability of [Al]/[B] catalysts should allow us to develop more efficient, economical, and recyclable tandem LA catalysts based on interor intramolecular and heterogeneous platforms. Accordingly, this contribution reports the first such main-group tandem LA catalytic system for highly effective and selective CO₂ reduction into CH₄ through hydrosilylation using a pair of [Al] and [B] LAs (Chart 1) which, when each used only, is ineffective (in

Chart 1. Proposed Fundamental Steps Involved in the Mixed Main-Group [AI]/[B] LA System for Tandem Catalytic Hydrosilylation of CO₂ into CH₄



case of [B]) or only marginally effective (in case of [Al]) for catalyzing this transformation. In this context, we present herein a full account of our combined experimental and theoretical/computational investigations into this novel maingroup [Al]/[B] tandem catalyst system.

RESULTS AND DISCUSSION

Fixation of CO₂ with {Et₃SiH + [Al]} and Selective Reduction of CO₂ to CH₄ by {Et₃SiH + [AI]/[B]}. As [B] was shown to be ineffective in catalyzing reduction of CO₂ via hydrosilylation,²⁹ we turned our attention to [Al], a stronger LA based on several lines of experimental and theoretical evidence. 27,30 Most recently, we have revealed that the Al center of the even unsolvated $Al(C_6F_5)_3$ is not truly tricoordinated, but it adopts a dimeric structure with double Al...F(ortho) interactions in solid state.³¹ Such weak interactions are readily destroyed by addition of more electron-donating substrates than [Al] itself, such as toluene, ferrocene, or even fluorosilane and hydrosilane.^{30a,31,32} For instance, the unsolvated [Al] forms a stable and isolable adduct with Et_3SiH in hexanes, while the interaction between [B] and Et₃SiH could only be detected by indirect spectroscopic evidence.^{28h} These observations suggest that the activation of the Si-H bond by [Al] is stronger than by [B]. However, the hydrosilylation of ketones by Et₃SiH proceeds more efficiently with [B] than with [Al], due to the high Lewis acidity and oxophilicity of [Al] that favor its complexation with the ketone substrates, greatly suppressing the Si-H bond activation. On the other hand, the feasible dissociation between the carbonyl substrates and [B] (even at a small ratio) enables the activation of the Si-H bond by the free [B] and the subsequent hydrosilylation.^{28j} Notably, Stephan et al. showed that [Al] reacts with CO2 at 90 °C to form a dimeric Al compound $[(C_6F_5)C(O)OAl(C_6F_5)_2]_2$ by insertion of CO₂ into the Al- C_6F_5 bond,³³ while Müller reported that a [Si-H...Al] system with a coordinatively saturated, less Lewis acidic aluminum center is inactive for CO₂ fixation.³⁴

Excitingly, the stoichiometric reaction between [Et₃SiH- $Al(C_6F_5)_3$ and CO_2 at room temperature (RT) (entry 3, Table 1) revealed the rapid disappearance of the silane Si-H signal at 3.75 ppm and concomitant appearance of new signals at 8.17 ppm in ¹H NMR, at 172.8 ppm in ¹³C NMR, and at 46.0 ppm (downfield shifted from 0.0 ppm from the silane) in ²⁹Si NMR (Figure 1). These spectroscopic features are consistent with the generation of a complexed silyl formate species, which was identified as the complex HCOOSiEt₃-[Al] (1-[Al]), the structure of which was confirmed by single-crystal X-ray diffraction (SC-XRD) analysis (Chart 2). No further reduction to lower oxidation state species was detected. With 10 mol % of [Al], the generation of 1-[Al] (10% based on Et₃SiH) was observed, but no further silane conversion was achieved. Other silanes such as ⁱPr₃SiH and PhMe₂SiH also led to the formation of the corresponding silvl formate-[Al] complexes (see the Supporting Information). These observations suggest that the activation of CO_2 by [Al] likely occurs through the typical LAcarbonyl activation and the LA is not released from the

Table 1. Selected Results of CO₂ Reduction via Hydrosilylation by [Al] and [Al]/[B]

| entry | silane | cat. | conditions | [SiH] conv. $(\%)^a$ | CH_4 yield $(\%)^b$ |
|-------|---------------------|----------------------|-------------|----------------------|-----------------------|
| 1 | Et ₃ SiH | _ | 12 h, 80 °C | 0 | 0 |
| 2 | Et ₃ SiH | 5% [B] | 12 h, 80 °C | 2 | 0 |
| 3 | Et ₃ SiH | 100% [Al] | 1 h, RT | 98 | 0 |
| 4a | Et ₃ SiH | 10% [Al] | 10 h, 80 °C | 39 | 14 |
| 4b | Et ₃ SiH | 10% [Al] | 24 h, 80 °C | 54 | 16 |
| 5 | Et ₃ SiH | 5.0% [Al] + 5.0% [B] | 1 h, RT | 15 | 7 |
| 6 | Et ₃ SiH | 5.0% [Al] + 5.0% [B] | 5 h, 80 °C | 100 | 82 |

^aBased on silane consumption. ^bSee the SI for yield calculation details.

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Figure 1. Comparisons of ¹H, ¹³C, and ²⁹Si NMR spectra before (bottom) and after (top) CO₂ fixation by [Et₃SiH + Al(C₆F₅)₃] in C₆D₅Br at 25 °C.

Chart 2. CO_2 Insertion into Si-H Bond with the $[Et_3SiH + Al(C_6F_5)_3]$ System To Form the [Al]-Coordinated Silyl Formate and the Corresponding Solid-State Structure Determined by SC-XRD



resulting product under these conditions, thus requiring a stoichiometric amount of LA. To render this reaction catalytic and possibly promote further hydrosilylation of the formed intermediates, we heated the above mixture to 80 °C for 24 h, achieving a moderate conversion of 54% (calculated from the silane consumption) with a 16% CH₄ yield (entry 4b, Table 1). These results imply that the catalytic performance of [AI] decreases as the basicity of the subsequent reaction intermediates [HCOOSiEt₃, H₂C(OSiEt₃)₂, and H₃COSiEt₃, vide infra] increases. Intriguingly, when we employed a mixed LA system containing 5 mol % [AI] + 5 mol % [B], quantitative conversion of the silane with 82% CH₄ yield was achieved at 80 °C in 5 h (entry 6, Table 1; Figure 2).

Isolation of Reaction Intermediates. To elucidate the possible mechanism for this unique [Al]/[B] tandem catalyst



Figure 2. ¹H NMR (25 °C, C_6D_5Br) spectra of hydrosilylation of CO_2 catalyzed by the mixed Al/B system (entries 5 and 6, Table 1) at different time intervals (•: HCOOSiEt₃-[Al], ‡: H₂C(OSiEt₃)₂-[Al], †: free H₂C(OSiEt₃)₂ and *: toluene from [Al] and NMR solvent residue).

system, we synthesized each of the intermediates or byproducts involved in the above reduction scheme, including HCOO-SiEt₃, $H_2C(OSiEt_3)_2$, $H_3COSiEt_3$, and $Et_3SiOSiEt_3$ (Scheme 1).

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Scheme 1. Preparation of the Reaction Intermediates Involved in the Overall CO₂ Reduction Cycle

HCOOH + Et₃SiH
$$\xrightarrow{Pd/C}$$
 HCOOSiEt₃ (1)
1 + Et₃SiH $\xrightarrow{1)[B]; 2)}$ DABCO
H₂C(OSiEt₃)₂ (2)
CH₃OH + Et₃SiH $\xrightarrow{[B]}$ H₃COSiEt₃ (3)
2 $\xrightarrow{[B]}$ [H₂CO] $\left\{ \begin{array}{c} 0\\ H\\ -C\\ H \end{array} \right\}$

Compounds HCOOSiEt₃ and H₃COSiEt₃ were readily prepared from the dehydrogenative coupling of the corresponding precursors by Pd/C or [B].^{26b} On the other hand, successful examples or selective formation and isolation of $H_2C(OSiEt_3)_2$ remain scarce.^{5a,15,16} Piers et al. noted that the sequestration of the [B] catalyst to prevent further hydrosilvlation is necessary.^{16a} Interestingly, in our attempts to reduce HCOOSiEt₃ with 1 equiv of Et₃SiH, we observed selective formation of $H_2C(OSiEt_3)_2$ in 5 min, after which the product gradually underwent further rearrangement to form 1,3,5-trioxane (a formaldehyde equivalent) and hexaethyldisiloxane Et₃SiOSiEt₃. This process is believed to proceed through coordination of the catalytic amount of [B] to the substrate, as sequestration of [B] with DABCO (1,4-diazabicyclo[2.2.2]octane) prevented the aforementioned rearrangement reaction and enabled the isolation of $H_2C(OSiEt_3)_2$. In a separate NMR scale experiment, $H_2C(OSiEt_3)_2$ was left for 1 day in C_6D_5Br and remained intact, but addition of 5 mol % of [B] converted $H_2C(OSiEt_3)_2$ into 1,3,5-trioxane and $Et_3SiOSiEt_3$ in 1 h

Scheme 2. Binding Interaction of Each Intermediate with [Al] and [B] Catalysts



Figure 3. X-ray crystal structures of 1-[AI] and 1-[B] with thermal ellipsoids drawn at the 50% probability. Hydrogen atoms except H1 were omitted for clarity. Selected bond lengths (Å) and angles (deg) for 1-[AI]: C1–O1 1.277(2), C1–O2 1.242(2), Al1–O2 1.8532(16), Si1–O1 1.7717(15), C1…F8' 2.935(3), and C1…F13' 2.908(3); for 1-[B]: C1–O1 1.276(2), C1–O2 1.245(2), B1–O2 1.621(2), Si1–O1 1.7499(14), C1…F5 2.838(3), and C1…F11 2.748(2).

(Figures S21 and S22). Indeed, the formation of the formaldehyde intermediate even without a catalyst is energetically feasible for bis(phenylsilyl)acetal $H_2C(OSiH_2Ph)_2$, via intramolecular elimination of PhH₂SiOSiH₂Ph, as predicted by calculations reported by Wang et al.³⁵ Our proposed mechanism through coordination of $H_2C(OSiEt_3)_2$ to [B] will be discussed in the following sections. The final disiloxane product of CO₂ reduction, Et₃SiOSiEt₃, can also be isolated as the byproduct by vacuum distillation.

Binding Interaction between the Intermediates and [AI] or [B]. With all three intermediates in hand, we studied their interaction with [AI] and [B] (Scheme 2). As expected, the less sterically hindered HCOOSiEt₃ forms a stable adduct with both [AI] and [B]. Crystalline 1-[AI] and 1-[B] adducts can be isolated in pure form by crystallization of equimolar 1 and [AI] or [B] in hexanes at -30 °C. Further supporting evidence came from multinuclear spectroscopic data (Figures S1–9) chiefly: (1) the aldehyde proton appears at 8.17 and 7.95 ppm in ¹H NMR for 1-[AI] and 1-[B], respectively; (2) the ¹⁹F NMR spectra also exhibit typical patterns for tetracoordinate aluminum and boron species; (3) both the formate and the LA signals are present in the ¹³C NMR and assigned; and (4) the ²⁹Si NMR signals are significantly

downfield shifted for 1-[Al] (δ 46.0 ppm) and 1-[B] (δ 46.4 ppm) when compared to the free 1 (δ 26.5 ppm). Worth noting here is that the spectroscopic data of 1-[Al] is consistent with those generated by the stoichiometric reaction between CO₂ and [Et₃SiH·Al(C₆F₅)₃].

The direct structural evidence for the 1:1 complexation in 1-[AI] and 1-[B] was derived from the SC-XRD analysis (Figure 3). In the structure of 1-[Al], the Al–O2 bond is rather strong, as indicated from the short bond length of 1.8532(16) Å. Notably, the C1-O2 and C1-O1 distances are 1.242(2) and 1.277(2) Å. This specific alternation suggests that C1-O2 features a more double-bond character, while C1-O1 features a more single-bond character. In another word, if 1-[Al] is viewed as a model in which [Al] and [Si]⁺ are competing for the middle formate anion, then [Si]+ exhibits stronger interaction with the anion. Nonetheless, both C1-O2 and C1–O1 distances are shorter than the typical C–O single bond (cf. 1.43 Å) and longer than the C=O double bond (cf. 1.23 Å), indicative of a certain degree of conjugated resonance structure along the -O1-C1-O2- moiety. The middle C1 carries a partial positive charge, accounting for the further secondary interactions with two para-F atoms [C1…F contact: 2.908(3) and 2.935(3) Å] from two neighboring molecules. Scheme 3. Catalytic Hydrosilylation Reaction of Each Fundamental Step



The structural parameters for 1-[**B**] are overall similar to those of 1-[**A**l]: C1–O1 1.276(2), C1–O2 1.245(2), B1–O2 1.621(2), except that secondary C…F contacts [2.748(2) and 2.838(3) Å] were derived from an intramolecular fashion with the adjacent $-C_6F_5$ ring instead, presumably due to the smaller radius of boron and shorter bond lengths around it. Further evidence for activation of the carbonyl moieties by LAs was deduced from FT-IR analysis of the solid samples of 1-[**A**I] and 1-[**B**] (Figure S32). When compared with the uncoordinated 1 (cf. 1708 cm⁻¹), both 1-[**A**I] (1604 cm⁻¹) and 1-[**B**] (1597 cm⁻¹) exhibit significant shifts of the C==O stretching mode to lower frequencies by 104 and 111 cm⁻¹, respectively.

In contrast, the interaction of [Al] and [B] with the bulkier and less basic H₂C(OSiEt₃)₂ and H₃COSiEt₃ is different (see Scheme 2 and SI). In the case of the more sterically hindered and less acidic [B] (relative to [Al]), there was no spectral change from the ¹H NMR signals for its 1:1 mixture with $H_2C(OSiEt_3)_2$ or $H_3COSiEt_3$. However, based on the fact that $H_2C(OSiEt_3)_2$ readily undergoes the rearrangement reaction in the presence of [B] in a prolonged period, it is likely that [B] can bind to $H_2C(OSiEt_3)_2$, albeit in a very weak fashion that favors the dissociated form. Given that the bulkiness of $H_3COSiEt_3$ is less than that of $H_2C(OSiEt_3)_2$, it is plausible that [B] can also bind to $H_3COSiEt_3$ reversibly, even though the ¹H data again indicated that this equilibrium strongly favors the dissociated form. To the contrary, the NMR scale reaction between [Al] and H₂C(OSiEt₃)₂ or H₃COSiEt₃ results in spectral shifts in the ¹H signals, corresponding to complexation between [Al] and H₂C(OSiEt₃)₂ or H₃COSiEt₃. In particular, [AI] forms isolable crystalline complex with H₃COSiEt₃. We tried to perform SC-XRD analysis of the complex but failed to obtain satisfactory results after several attempts due to poor crystal quality. Nonetheless, the overall molecular skeleton and atom connectivity of this adduct could be recognized from the

crude data (Figure S34). Collectively, these observations further support that in the subsequent reduction of loweroxidation-state species $HCOOSiEt_3$, $H_2C(OSiEt_3)_2$, and $H_3COSiEt_3$, [B] should serve as a better catalyst than [Al] due to the weaker substrate-catalyst interaction, which facilitates the FLP-type Si-H activation. In addition, the byproduct $Et_3SiOSiEt_3$ does not form a detectable adduct with either [B] or [Al], which enables [Al] and/or [B] to re-enter the catalytic cycle after the last step (CH₄ generation).

Kinetic and Mechanistic Studies of Hydrosilylation of CO_2 . To further address our hypothesis on the catalytic roles of each LA in the current mixed tandem [Al]/[B] system, we carried out a kinetic study on each step of the hydrosilylation (Scheme 3), coupled with a computational investigation (vide infra). As described previously, in step 1, the fixation of CO_2 is mediated by [Al] to form 1-[Al]. The following hydrosilylation steps 2-4 proceed more efficiently with [B] than with [Al]. Among them, the reactions of HCOOSiEt₃ or H₃COSiEt₃ with 1.0 equiv of Et_3SiH in the presence of 5 mol % [B] were completed within 5 min at RT, while the slowest step, the hydrosilylation of $H_2C(OSiEt_3)_2$ under similar conditions generated a mixture of unreacted H₂C(OSiEt₃)₂, CH₄, and a trace amount of H₃COSiEt₃. Nonetheless, treatment of HCOOSiEt₃ with 3.0 equiv of Et₃SiH and 5 mol % [B] resulted in almost quantitative conversion to CH₄ in 4 h, during which $H_2C(OSiEt_3)_2$ accumulates as the detectable intermediate. In addition, the reduction of 1,3,5-trioxane with 1.0 equiv of Et₃SiH and 5 mol % [B] led to exclusive formation of H₃COSiEt₃, indicating that the hydrosilylation of trioxane is even faster than that of H₃COSiEt₃. In comparison, the hydrosilylation of the related substrates with [Al] at RT is not effective at all and only provided a trace amount of the reduction products, but at 80 °C after longer times some hydrosilylation products were observed (Scheme 3)



Figure 4. Kinetic plots under current standard conditions (left, 80 °C, 5 mol % [Al], and 5 mol % [B]) and varied loading of [Al] and [B] (right).

| Table 2. | Selected | Results o | of CO ₂ | Hydrosil | vlation a | at 80 ' | °C under | Different | Tandem | Catalyst | Loadings |
|----------|----------|-----------|--------------------|----------|-----------|---------|----------|-----------|--------|----------|----------|
| | | | | | | | | | | , | |

| catalyst composition | reaction time $(h)^a$ | max TOF $(h^{-1})^{b}$ | $r_{\rm SiH} ({\rm M \ h^{-1}})^c$ | conv. (%) ^d | CH ₄ yield (%) ^e |
|----------------------|-----------------------|------------------------|------------------------------------|------------------------|--|
| 10% [Al] + 5.0% [B] | 3 | 5.0 | 0.20 | 100 | 77 |
| 5.0% [Al] + 5.0% [B] | 5 | 6.6 | 0.13 | 100 | 82 |
| 2.5% [Al] + 5.0% [B] | 10 | 7.9 | 0.079 | 100 | 91 |
| 1.0% [Al] + 5.0% [B] | 30 | 6.03 | 0.024 | 99 | 94 |
| 5.0% [Al] + 2.5% [B] | 9 | 7.85 | 0.0785 | 100 | 80 |

^aBased on [SiH] consumption from ¹H NMR measurement. ^bBased on the steepest slope from the linear fit of conversion against time at the early stage of hydrosilylation, see Figures S27–S31. ^cRate of [SiH] consumption, calculated based on max TOF. ^dBased on [SiH] consumption. ^eSee the SI for yield calculation details.

To gain further insights of such tandem hydrosilylation of CO₂, we carried out a kinetic study of the overall reduction process. Under our standard conditions (entry 6, Table 1), the reaction was complete in 5 h. Et₃SiH consumption and CH₄ yield at the early stage of the reaction showed a linear relationship with time (Figure 4). The concentration of detectable intermediate $H_2C(OSiEt_3)_2$ (2) reached a plateau at 1.5 h and then decreased gradually. The maximum turnover frequency (TOF) of the consumption of [SiH] using the first 5 data points from the initial 2 h was calculated to be 6.6 h^{-1} , and the rate for the [SiH] conversion was $r_{SiH} = 0.13$ M h⁻¹. Variation of the [Al] catalyst concentration (from 1.0 mol % to 10 mol %, Table 2) greatly impacted the [SiH] consumption rate. Overall, the reduction rate of CO₂, as monitored by the consumption of [SiH] signal, was proportional to the [Al] loading. The only observed species at δ 4.57 ppm after complete [SiH] conversion was the 2-[Al] adduct, which is consistent with the stoichiometric reaction between 2 and [Al]. As expected, lowering the [Al] loading reduced the amount of the final 2-[Al] residue and hence improved the yield of CH_4 . In addition, decreasing the [B] concentration also resulted in a decrease of CO₂ reduction rate. Remarkably, when only 1.0 mol % of [Al] was employed, the highest CH₄ yield of 94% was achieved (Table 2).

Collectively, the above results showed that CO_2 fixation is promoted by [Al], while the subsequent hydrosilylation reactions are catalyzed by [B]. To further clarify the preferred pathway of each step, namely conventional carbonyl activation versus FLP-type silane activation, and to explain the observed activity for the CO_2 reduction into CH_4 by [Al] alone but requiring 80 °C for 24 h (entries 4a-b, Table 1), albeit being much less effective than the mixed [Al]/[B] tandem catalyst system, we performed the following additional experiments. We first mixed equimolar 1-[B] and Et₃SiH-[Al], which resulted in the instantaneous replacement of Et₃SiH by HCOOSiEt₃ to form 1-[Al]. However, the subsequent hydrosilylation was hindered, as only a trace amount of 2-[Al] was detected up to 1 h, in contrast to the rapid reduction of 1-[B] by Et₃SiH in the absence of [Al]. This observation is in line with the Si-H activation mechanism, in which the free carbonyl has to be formed to initiate the attack at the Si center of a transient Si- $H \cdots [B]$ intermediate. If the carbonyl is coordinated to an additional LA, [Al], such nucleophilic attack becomes less plausible. With this premise, we gratifyingly found that heating the mixture to 80 °C led to the formation of 2-[Al], due to thermally induced dissociation of HCOOSiEt₃ from [Al], enabling its reduction to $H_2C(OSiEt_3)_2$ that recombines with [Al] to form 2-[Al]. Addition of an excess amount of Et₃SiH into such a mixture further converted $H_2C(OSiEt_3)_2$ to CH_4 at 80 °C. In a similar fashion, addition of [Al] to a mixture of H₃COSiEt₃, [B] and Et₃SiH reduced the hydrosilylation rate of H₃COSiEt₃ to CH₄. These experiments further confirm that the reduction steps 2-4 occur through the FLP Si-H activation mechanism, and a higher temperature is necessary to facilitate the release of substrates (1, 2, and 3) from [Al], which also renders [Al] catalytic by reentering the catalytic cycle for the step 1 reduction.

In order to investigate the catalyst recyclability, the catalytic system consisting of 10 mol % [Al] and 5.0 mol % [B] was tested for 3 cycles. In the first cycle, the hydrosilylation was complete in 3 h (Figure 5). After careful removal of excess CO_2 and CH_4 under vacuum, the system was recharged with the same amount of Et_3SiH and CO_2 as in the first cycle. The hydrosilylation of the second and third cycles was complete in 6 and 14 h, respectively. This recycling experiments indicated that the [Al]/[B] catalysts survived during catalytic cycle and are recyclable. Although we detected the decrease in catalyst efficiency from cycle to cycle (presumably due to partial



Figure 5. Recyclability of CO_2 reduction with 10 mol % [Al] and 5.0 mol % [B]. Reaction time for complete SiH consumption in each cycle: 3, 6, and 14 h.

hydrolysis of the catalyst during the reloading of CO_2), quantitative conversion can still be achieved on the third load of Et_3SiH and CO_2 .

Computational Studies of Fundamental Steps 1–4.³⁶ In this section we discuss each reduction step separately starting from the hydrosilylation of CO_2 to formate and ending with the hydrosilylation of methyl silyl ether to methane. In Schemes 4, 5, and 6 numbers in green refer to [Al], while numbers in blue refer to [B]. Further, to strengthen connections with experiments, we labeled DFT structures using letters, such as A, B, C, etc., while for structures involving intermediates 1-3 we kept the same labeling scheme as that used in the description of the experimental results.

Step 1: Hydrosilylation of CO_2 to Silylformate 1-[X] (X = Al, B). The two possible reaction pathways we investigated to generate 1 from CO_2 are reported in Scheme 4.

Along pathway 1 the individual LA directly activates the CO_2 molecule toward the H-transfer from Et_3Si-H . Pathway 2, instead, starts with a FLP-type Si-H activation with

coordination of Et₃SiH to the LA, followed by the H-transfer to the CO_2 molecule through an insertion mechanism or a SN_2 transition state and by rearrangement of the initially formed $[OCOSiEt_3]^+/[H-LA]^-$ ion pair. Focusing on pathway 1, in the case of [B] we were not able to locate a CO_2 -B(C_6F_5)₂ adduct, B. Nevertheless, we found two possible four-membered transition states, B/C and B/1-[X] in Scheme 4, where the silicon attacks one oxygen of CO₂ favoring the simultaneous Htransfer from Et₃Si-H to the CO₂ carbon atom. Transition state B/C collapses into intermediate C before reaching 1-[X], favored by 9.4 kcal/mol, while transition state B/1-[X]collapses directly into product 1-[X]. The energy cost to reach transition states B/1-[X] and B/C, around 35-39 kcal/ mol, and the fact that these transition states should be formed by the encounter of three molecules due to the instability of the preformed B intermediate explain the experimentally observed inability of [B] to activate CO₂. Conversely, the CO₂-alane adduct **B** was located 6.5 kcal/mol below separated $Al(C_6F_5)_3$ and CO_2 . This interaction weakens the C=O bond of the CO_2 . moiety, as shown by the slightly elongated C=O bond distance (1.19 Å in **B** vs 1.17 Å in the free CO_2). [Al] coordination results in an increased positive charge on the C atom of the CO_2 moiety (1.07e in **B** vs 0.99e in the free CO_2), promoting the H-transfer from Et₃SiH to the electrophilic C center. This H-transfer occurs via the four-membered transition state B/C, with the [Al] coordinated O atom attacking the Si atom and an energy barrier of 22.0 kcal/mol. Transition state B/C collapses into intermediate C, which can further precipitate in the final product 1-[X] ($\Delta G = -28.8$ kcal/mol) through the almost barrier-less transfer of the [Al] to the carbonyl oxygen, via transition state C/1-[X]. The alternative transition state with the uncoordinated O atom of **B** attacking the Si atom, B/1-[X], and leading to 1-[X] directly, was located 3.5 kcal/mol above B/C. The stability of the CO_2 -[Al] adduct B and the lowenergy barrier of 22.0 kcal/mol explain the experimentally observed capability of [Al] to activate CO₂.

As for the competitive mechanism via Si-H activation, pathway 2 in Scheme 4, it involved the formation of a silane-LA adduct. Formation of such adduct was found to be favored by 7.2 kcal/mol with [Al], whereas in the case of [B] it is





^aFree energies (kcal/mol) in *n*-Hexane are reported for X = Al (numbers in green) and B (numbers in blue) (n.l. = not located).

Scheme 5. Hydrosilylation of Silylformate 1-[X] to H₂C(OSiEt₃)₂/Formaldehyde^a



^{*a*}Free energies (kcal/mol) in *n*-hexane are reported for X = Al (numbers in green) and B (numbers in blue).

disfavored by 2.7 kcal/mol. The transition state for the following insertion of CO₂ into the Si–H bond, D1/1-[X] in Scheme 4, lies at 32.7 kcal/mol for [Al] and 33.1 kcal/mol for [B]. Finally, we also tried to promote Si–H activation via the SN₂-type transition state D2/1-[X], corresponding to the concerted attack of the CO₂ oxygen to the silicon atom and transfer of the H atom to the [X] moiety. However, we were not able to locate this kind of transition state for both [Al] and [B]. Overall, calculations confirmed the experimental inefficiency of [B] in the first step of CO₂ hydrosilylation, since [B] is not able to effectively activate CO₂ through either direct coordination to CO₂ or FLP-type silane activation. As far as [Al] is concerned, calculations suggest that the reduction of CO₂ to silylformate occurs via CO₂ activation was ruled out.

Steps 2 and 3a: Hydrosilylation of Silylformate 1-[X] to $H_2C(OSiEt_3)_2$ (Step 2) and Formaldehyde (Step 3a). Starting from the LA-silylformate adduct 1-[X], we studied the mechanisms most likely operative in the second hydrosilylation step, namely from the silvlformate adduct 1-[X] to fomaldehyde. The pathways considered with the corresponding energetics are reported in Scheme 5. We considered the Htransfer step from a second Et₃SiH molecule to the starting 1-[X] via the same pathways considered above for hydrosilylation of CO₂. Specifically, we considered two transition states along the C=O activation pathway, i.e., 1-[X]/E and 1-[X]/2-[X], where the Et₃SiH molecule attacks the oxygen of the LA-O bond or the oxygen of the Si-O bond of 1-[X]. Transition state 1-[X]/E, leading directly to Et₃SiOSiEt₃ and the LAaldehyde adduct, E in Scheme 5, is favored by roughly 6-7 kcal/mol relative to transition state 1-[X]/2-[X], leading to adduct 2-[X], with the experimentally characterized

 $CH_2(OSiEt_3)_2$ intermediate 2 bound to the LA. In this conversion, the two LAs behave similarly, with an overall energy barrier through the favored transition state 1-[X]/E of roughly 29 kcal/mol. This relatively high-energy barrier can be ascribed both to the high steric pressure in this second hydrosilylation transition state (two molecules of silane plus one LA) and to the reduced electrophilicity of the formate carbon center with respect to that of CO₂, as revealed by the charges of these two C centers, i.e., 1.07e in CO₂ vs 0.74e in 1-[X]. Finally, formation of E from $1-[X] + Et_3SiH$ is exergonic by almost 4.0 kcal/mol for both LAs, which drives aldehyde formation. As for conversion of the experimentally characterized intermediate 2-[X] to the formaldehyde adduct E, it can proceed either in a single step through transition state 2-[X]/Eor through a two-step pathway via dissociation of [X] from 2-[X], followed by the direct reductive elimination of Et₃SiOSiEt₃ from 2 to release the aldehyde via transition state 2/E, following a similar mechanism reported in literature.³ According to our calculations, the one-step mechanism via transition state 2-[X]/E, with an overall energy barrier around 20 kcal/mol, is clearly favored over the direct reductive elimination from 2. Nevertheless, dissociation of the LA from **2-**[**X**] releasing intermediate **2** is energetically favored with [B], actually representing the thermodynamic product of the reaction.

As for the Si-H activation pathway, it starts with the conversion of 1-[X] into 2-[X], with the steps from 2-[X] to E already being discussed above. Thus, we focus here on the pathway from 1-[X] to 2-[X]. We calculated first the energetics involved in the release of the LA from the silylformate adduct 1-[X]. As expected, dissociation of [Al], with a ΔG of 25.2 kcal/mol, is more expensive than dissociation of [B], with a ΔG

Scheme 6. Hydrosilylation of Formaldehyde to Silyl Methanol and Then to Methane^a



^{*a*}Free energy (kcal/mol) in *n*-hexane are reported for X = Al (numbers in green) and B (numbers in blue).

of only 5.8 kcal/mol. Starting from the LA free species 1 and a preformed $Et_3SiH-[X]$ adduct, we located both the transition states for the direct formate insertion into the Si-H bond, F1/2-[X], and the direct H-transfer between Et_3SiH and the LA via SN₂-type reactivity, F2/2-[X]. Based on the energy of the H-transfer transition states, both the Si-H activation pathways we examined can be ruled out in the case of [Al], since transition states F1/2-[X] and F2/2-[X] are more than 35 kcal/mol above intermediate 1-[X]. Conversely, the insertion Si-H activation pathway is isoenergetic with the C=O activation pathway in the case of [B] and, more relevantly, the SN₂ type H-transfer mechanism via transition state F2/2-[X] shows a barrier of only 13.2 kcal/mol.

Focusing on the thermodynamic scenario, the stability of the intermediates shown in Scheme 5 seems to correlate with the different catalytic behavior in the presence of [Al] or [B]. In fact, for [Al] the thermodynamic product is the $H_2C(OSiEt_3)_2$ -[Al] adduct 2-[X], which is favored by 1.6, 5.3, and 17.1 kcal/ mol relative to E, 1-[X], and 2, respectively. As consequence, liberation of [Al] to promote further reactivity is compromised. On the contrary, intermediate 2 is the most stable species in the presence of [B], followed by E, with 1-[X] and 2-[X] being

clearly much less stable. The different thermodynamic stability of 2 and 2-[X] with [Al] and [B], together with the much lower energy barrier via the Si–H activation pathway, renders the [B] catalyst active in the reduction of silylformate to formaldehyde, in agreement with the experimental results.

Steps 3b and 4: Hydrosilylation of Formaldehyde to Silyl Methanol (Step 3b) and Then to Methane (Step 4). The reaction pathways we considered for the last two steps to form methane from the aldehyde-LA adduct E are reported in Scheme 6 with the related energies. Since no favored hydrosilylation pathway was located starting from 2 of Scheme 5, we focused on the aldehyde-LA adduct E that turns out to be the crucial species for the course of the reaction. Intermediate E can react with the third equivalent of Et₃SiH along the C=O activation pathway via the low-energy H-transfer transition state E/3-[X] with both the LAs considered, with an energy barrier around 5 kcal/mol. Alternatively, the key intermediate 3-[X] can also be reached from E along the Si-H activation pathway after dissociation of the LA to reach intermediate G. As calculated in the previous reaction sequences, dissociation of [Al] from E is clearly endergonic, whereas it is favored with [B]. The liberated formaldehyde can react with a Et₃SiH-LA adduct, intermediate H, via the already considered insertion or SN_2 -type pathways, through transition states H1/3-[X] and H2/3-[X], respectively. The insertion mechanism can be excluded since the H1/3-[X] transition state is more than 20 kcal/mol higher in energy than transition state $E/3\ensuremath{\cdot} [X]$ for both [Al] and [B]. As for the SN₂-type pathway, it is competitive for [B], since transition state H2/3-[X] is almost isoenergetic with transition state E/3-[X] along the C=O activation pathway. Differently, it can be excluded for [Al] since the transition state H2/3-[Al] is almost 24 kcal/mol higher in energy than transition state E/3-[Al].

Two different pathways were considered to generate methane from 3-[X]. The first pathway is the one-step Htransfer from the fourth equivalent of Et₃SiH molecule to the carbon of the activated adduct 3-[X] (C-O activation pathway) via transition state 3-[X]/K. This pathway can be excluded, since it requires the overcome of an energy barrier >60 kcal/mol (Scheme 6). The second pathway is a multistep process and starts with release of the LA from 3-[X], which is again strongly endergonic for [Al], whereas it is slightly exergonic for [B] (3-[X] vs 3 for [AI] and [B] in Scheme 6). After releasing LA from 3-[X], silyl methanol 3 proceeds to the last hydrosilylation step, via SN₂-type transition state I/J involving 3 and a Et₃SiH-LA adduct. This H-transfer step is rate determining, with a barrier of 16.0 kcal/mol for [B] (calculated from the most stable $3 + Et_3SiH + LA$ species) and of almost 40 kcal/mol for [Al] (calculated from the most stable 3-[X] +Et₃SiH species). As a consequence and in agreement with the experimental results, the [Al] catalyst is inactive also in this last hydrosilylation step, which proceeds smoothly with the [B] catalyst. The last H-transfer within the formal [CH₃O- $(Et_3Si)_2$ ⁺/[H-LA]⁻ adduct J to release methane via transition state J/K is almost barrier-less. As expected, formation of methane is thermodynamically favored by roughly 50 kcal/mol with respect to the silyl methanol intermediate 3. For the sake of simplicity, the insertion pathway is not reported for the conversion of 3 to methane, since it involves a clearly unfeasible barrier, as discussed in the previous sections.

Alane vs Borane: Electronic and Steric Analysis. The results reported in the previous sections showed that the experimentally observed catalytic difference between [Al] and



Figure 6. Calculated energy profiles of intermediates and transition states for [B] and [Al] pathways along the reaction coordinate.

[B] seems to be due to the much higher stability of the LAadducts with [Al] relative to [B], in intermediates 1-[X], 2-[X], E, and 3-[X] (see Schemes 5 and 6), which reflects the difficult release of the LA from the substrates and the high barrier for the Si-H bond activation. Focusing on the named intermediates, the relative stability of 1-[X], E and 3-[X] is more than 10 kcal/mol higher for [Al] than [B], respectively. This result reflects in the key transition states for Si-H activation (i.e., F2/2-[X] and H2/3-[X]) lying much higher in energy for [Al]. Moreover, 2-[X] is favored by -17.1 kcal/mol with [Al] and disfavored by 8.4 kcal/mol with [B] relative to 2, and 3-[X] is favored by -17.6 kcal/mol with [Al] and disfavored by 3.5 kcal/mol with [B] with respect to 3 (the total difference between [Al] and [B] is of 25.5 and 21.1 kcal/mol, respectively).

To better understand the large difference in the relative stability of these intermediates, we analyzed both electronic and steric effects. From an electronic point of view, the calculated electrophilicity of the two LAs (0.24 for [Al] and 0.38 for [B]) appears to indicate that the formation of [B]-adducts should be favored relative to the formation of [Al]-adducts. Indeed, the relative energies of the SN₂ H-transfer transition states relative to the preceding intermediates are in line with the electrophilicity results. Moving to sterics, we tested the relative

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stability of 2 and 3 with respect to 2-[X] and 3-[X] with XH₃ (X = Al and B) as LA. The difference in the binding energy of 2 and 3 to AlH₃ and BH₃ is only 2.7 and 4.5 kcal/mol, relative to a difference >20 kcal/mol with [Al] and [B]. Incidentally, the same analysis performed on the LA-HSiEt₃ adducts shows that the 10 kcal/mol of difference in the relative stability of the [Al]...H-SiEt₃ adduct with respect to the [B]...H-SiEt₃ one decreased to -2 kcal/mol in favor of the borane when AlH₃ and BH₃ are used as the LA. This suggests that steric effects strongly impact the binding ability of [B].

As a further test, we decomposed the gas-phase binding energy of [Al] and [B] to formaldehyde into a preparation and an interaction energy term. The first contribution is the energy paid to deform the LA and the aldehyde from their ideal conformations in the unbound state to the geometries they assume in the LA-adducts E. The interaction energy term, instead, corresponds to the energy gain due to the rigid interaction between the LA and the aldehyde frozen in the same conformation as in E. We found that the deformation of the free [B] to the geometry it has in E is 9.5 kcal/mol more expensive than the deformation of [Al], while the interaction energy between the deformed LA and formaldehyde is only 6.5 kcal/mol stronger with [Al]. The two terms cumulate in an [Al]-aldehyde bond being 16.0 kcal/mol stronger than the [B]aldehyde bond in the gas-phase.

Overall, these results indicated that the different behavior of [Al] and [B] can be ascribed to the significantly smaller size of boron that causes unfavorable steric repulsions between the C_6F_5 rings, as well as between the C_6F_5 rings and the substituents on the silane atoms, in the adducts.

In Scheme 7 we summarized the whole catalytic cycle in the presence of both [Al] and [B] LAs, while the calculated energy profiles of intermediates and transition states for [Al] and [B] pathways along the reaction coordinate were plotted in Figure 6. As the first step, hydrosilylation of CO_2 to silvlformate takes place via the [Al] catalyzed C=O activation pathway. Next, the reduction of HCOOSiEt₃ to formaldehyde proceeds through the [B] catalyzed SN₂-type Si-H activation mechanism. In the following reduction step to achieve intermediate Et₃SiOCH₃, the C=O and the Si-H activation pathways turn out as competitive with [B] since the corresponding determining barriers are almost the same in energy. Finally, in the last reduction step to methane, the SN₂ Si-H activation mechanism is again favored. The sterics of the substrate seems to play a key role, with the C=O activation pathway suffering the steric hindrance more than the SN₂ Si-H one due to a more crowded geometry of the corresponding transition state.³⁷ The thermodynamics of the adducts favors [Al]; however, in agreement with the experimental results, only [Al] is active in the first step, while [B] is active in the subsequent steps. As a consequence an exchange between the two Lewis acids is likely to take place during the four reduction steps.

CONCLUSIONS

In summary, we have developed the first example of highly selective reduction of CO_2 into CH_4 via tandem catalytic hydrosilylation by a mixed main-group $B(C_6F_5)_3/Al(C_6F_5)_3$ catalyst system. The results, obtained from our comprehensive study involving the detection, characterization, and independent synthesis of each reaction intermediate, reactions under catalytic conditions, and computational calculations as well as kinetic and mechanistic investigations, have demonstrated that [Al] is responsible for the first step of the reduction that

converts CO_2 into $HCOOSiEt_3$, while the subsequent reduction steps of $HCOOSiEt_3$ to $H_2C(OSiEt_3)_2$ to $H_3COSiEt_3$ and finally to CH_4 are catalyzed by [B]. The hydrosilylation of $H_2C(OSiEt_3)_2$, the rate-limiting step in the [B]-catalyzed FLP reduction sequences, is proposed to proceed through a twostep pathway involving the formation and reduction of formaldehyde. Our computational results further address the fixation of CO_2 into $HCOOSiEt_3$ by [Al] via the classical LAmediated C==O activation, the subsequent transformations into CH_4 by [B] through the FLP-type Si-H activation, as well as the $H_2C(OSiEt_3)_2$ reduction via the formaldehyde cycle, all of which are consistent with the experimental results.

We attribute this remarkably different catalytic behavior between $Al(C_6F_5)_3$ and $B(C_6F_5)_3$ to the higher overall Lewis acidity of [Al] derived from two conflicting factors, electronic and steric effects. While the study of the electronic term indicates that [B] has a higher electronic affinity, the steric term suggests that [B] pays much higher reorganization energy penalty due to both a smaller radius of boron and repulsion between the ortho-fluorine atoms. This stronger overall Lewis acidity of [Al], when compared to [B], renders its higher tendency to form stable [Al]-substrate (intermediate) adducts with CO_2 as well as intermediates 1, 2, and 3, hence accounting for its distinct yet complementary catalytic behaviors in the CO₂-to-CH₄ hydrosilylative reduction cycle. Overall, the roles of [Al] and [B] are not only complementary but also synergistic in the total reduction of CO_{2} , which render both [Al]-mediated first reduction step (which, when carried out alone, is a stoichiometric reaction) and [B]-mediated subsequent steps catalytic. With an optimized loading and [Al]/[B] ratio of 1.0%:5.0%, a high CH₄ production yield of 94% has been achieved. Such a catalytic system is also shown to be recyclable, based on three cycling experiments. The tunability of [Al]/[B] catalysts should allow one to develop more efficient, economical, and recyclable tandem LA catalysts based on inter- or intramolecular and/or heterogeneous catalysts.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/jacs.6b01497.

Full experimental details, additional figures, and complete ref 36 (PDF)

Crystallographic data (CIF) Crystallographic data (CIF)

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Notes

The authors declare no competing financial interest.

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(37) This result is in agreement with the observations reported in ref 28c for the hydrosilylation of acetone with Me₃SiH and B(C_6F_5)₃. In fact, comparing the barrier for the reduction of acetone with that for the reduction of the less hindered formaldehyde, the C=O pathway turns out to be much unfavored for the ketone substrate.