

## Efficient Homogeneous Catalysis in the Reduction of CO<sub>2</sub> to CO

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Nature uses carbon dioxide, on a massive scale, as a one-carbon building block for the synthesis of organic molecules.<sup>1</sup> An important pathway for the consumption of CO<sub>2</sub> is its reduction to CO by the enzyme acetyl-CoA synthase/CO dehydrogenase (ACS-CODH).<sup>2</sup> Due to the large energy input required to generate it from CO<sub>2</sub>, CO is produced industrially from fossil fuels.<sup>3</sup> Even with strong reducing agents, however, overcoming the O=CO bond enthalpy of 532 kJ/mol<sup>4</sup> often presents kinetic difficulties.<sup>5,6</sup>

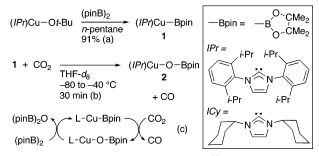
Certain metal complexes abstract oxygen readily from  $\text{CO}_2$ ,<sup>7</sup> but the resulting metal—oxygen bonds are necessarily strong, and catalytic turnover is rare.<sup>8</sup> Photolytic<sup>9</sup> and photocatalytic<sup>10</sup> approaches show promise, and synthetic electrocatalysts have achieved impressive yields and selectivities in the reduction of  $\text{CO}_2$  to CO.<sup>11</sup> However, the chemical processes involved are obscure, making it difficult to improve these systems by design, and CODH remains notably the most efficient catalyst for this reduction.<sup>12</sup> We report herein that a new carbene-supported copper(I) boryl complex abstracts oxygen from  $\text{CO}_2$  and undergoes subsequent turnover readily. Using an easily handled diboron reagent as the net oxygen acceptor,<sup>13</sup> these key steps permit unprecedented turnover numbers and frequencies for the chemical reduction of  $\text{CO}_2$  to CO in a homogeneous system.

While exploring the chemistry of organocopper(I) complexes supported by N-heterocyclic carbene (NHC) ligands,<sup>14</sup> we sought to synthesize a copper(I) boryl complex and explore its reactivity toward CO<sub>2</sub>. Metal boryls often display distinctive reactivity,<sup>15</sup> catalyzing a number of remarkable transformations.<sup>16</sup> Although C–B bond-forming reactions have been achieved using diboron compounds with catalytic<sup>17a</sup> or stoichiometric<sup>17b</sup> copper(I), welldefined copper boryl complexes have not been described.

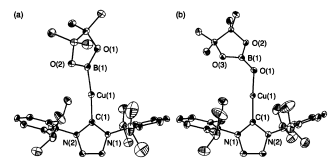
The known (*IPr*)Cu(Ot-Bu) reacts rapidly with bis(pinacolato)diboron (pinB–Bpin), forming a product identified as (*IPr*)Cu(Bpin) (**1**, Scheme 1) by <sup>1</sup>H and <sup>11</sup>B NMR spectroscopy. Diffusion of hexane vapor into a concentrated solution of **1** in toluene, carried out at -40 °C to avoid thermal decomposition,<sup>18</sup> produces single crystals suitable for analysis by X-ray diffraction. The resulting structure (Figure 1a) shows a monomeric, nearly linear coordination geometry with a Cu–B distance of 2.002(3) Å.

Complex **1** reacts with CO<sub>2</sub> under atmospheric pressure in C<sub>6</sub>D<sub>6</sub> solution, quantitatively forming a new complex within minutes. The resonance for **1** in the <sup>11</sup>B NMR spectrum, a broad singlet at 41.7 ppm, is replaced by a singlet at 21.8 ppm, indicative of boron bound to three oxygen atoms.<sup>19</sup> Single crystals of this new copper complex are grown by diffusion of hexane vapor into a concentrated toluene solution. The X-ray crystal structure (Figure 1b) reveals the product to be (*IPr*)Cu(OBpin) (**2**): The copper boryl complex abstracts oxygen from CO<sub>2</sub>, implying the release of CO as the byproduct.

To confirm the formation of CO, <sup>13</sup>C-labeled CO<sub>2</sub> is introduced to a resealable NMR tube containing a solution of **1** in THF- $d_8$  at -80 °C. After 30 min of gradual warming, analysis by <sup>11</sup>B (Figure 2a) and <sup>1</sup>H NMR spectroscopy at -40 °C indicates complete Scheme 1<sup>a</sup>



<sup>*a*</sup> Isolated yield, contains some **2** (5 mol % by <sup>11</sup>B NMR); (b) reaction is complete in <10 min at ambient temp; (c) L = IPr or ICy.

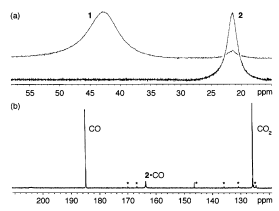


*Figure 1.* X-ray crystal structures, shown as 50% thermal ellipsoids, of boryl complex  $1 \cdot C_6 H_{14}$  (a), and borate  $2 \cdot C_7 H_8$  (b). Hydrogen atoms (calcd) and solvent are omitted for clarity. Selected bond lengths (Å) and angles (deg), (a): Cu(1)-B(1) 2.002(3), Cu(1)-C(1) 1.937(2), C(1)-N(1) 1.363(3), C(1)-N(2) 1.363(3), C(1)-Cu(1)-B(1) 168.07(16), N(1)-C(1)-N(2) 102.97(18); (b): Cu(1)-O(1) 1.8096(16), O(1)-B(1) 1.306(3), Cu(1)-C(1) 1.857(2), C(1)-N(1) 1.355(3), C(1)-N(2) 1.364(3), C(1)-Cu(1)-O(1) 174.85(10), B(1)-O(1)-Cu(1) 13.61(16), N(1)-C(1)-N(2) 103.09(18).

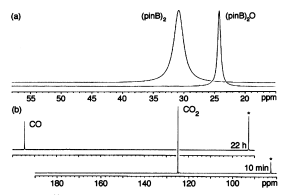
conversion of **1** to **2**. The sole labeled products visible in the <sup>13</sup>C NMR spectrum (Figure 2b) are <sup>13</sup>CO ( $\delta$  184 ppm) and an adduct ( $\delta$  164 ppm) formed reversibly from CO and borate **2**.<sup>20</sup>

Treatment of **2** in C<sub>6</sub>D<sub>6</sub> solution with pinB–Bpin smoothly regenerates **1**, forming the stable byproduct pinB–O–Bpin,<sup>21</sup> over a reaction time of about 20 min. The success of this turnover step closes a catalytic cycle for the deoxygenation of CO<sub>2</sub>. Addition of a THF solution of (*IPr*)Cu(Ot-Bu) to a 100-fold excess of pinB– Bpin under an atmosphere of CO<sub>2</sub> results in the complete conversion of pinB–Bpin to pinB–O–Bpin within 20 h at ambient temperature, as judged by <sup>11</sup>B NMR analysis of an aliquot from the reaction mixture (Figure 3a). When labeled CO<sub>2</sub> is used as the limiting reagent, in the presence of ca. 2 mol % of precatalyst **1**, the <sup>13</sup>C NMR spectrum indicates complete consumption of CO<sub>2</sub>, with CO representing the sole significant product (Figure 3b).

In the absence of copper catalyst, under otherwise identical conditions, no pinB-O-Bpin is detected, demonstrating that the diboron compound by itself is kinetically unable to reduce CO<sub>2</sub> to any observable extent. Control reactions run using copper precatalyst and pinB-Bpin in the absence of CO<sub>2</sub> (under an atmosphere



**Figure 2.** (a) <sup>11</sup>B NMR spectra before and after reaction with CO<sub>2</sub>: **1** (containing 5% **2**) and **2**; (b) <sup>13</sup>C NMR spectrum after reaction of **1** with excess <sup>13</sup>CO<sub>2</sub> (THF- $d_8$ , -80 to -40 °C, 30 min); \* denotes ligand-derived resonances; solvent and aliphatic resonances omitted for clarity.



*Figure 3.* (a) <sup>11</sup>B NMR spectra showing conversion of pinB–Bpin to pinB–O–Bpin by catalytic reduction of CO<sub>2</sub> (excess CO<sub>2</sub>, 1 mol % 1, ambient temp, THF- $d_8$ , 20 h); (b) <sup>13</sup>C NMR spectra, offset for clarity, before and after catalytic reduction of <sup>13</sup>CO<sub>2</sub> to <sup>13</sup>CO (excess pinB–Bpin, ca.. 2 mol % 1, ambient temp, THF- $d_8$ ); \* denotes pin (Me<sub>2</sub>CO)<sub>2</sub> resonance.

of argon or even dry air) show, at most, stoichiometric formation of pinB-O-Bpin with respect to copper. Thus, oxidation of pinB-Bpin by adventitious air does not contribute significantly to the formation of pinB-O-Bpin under these conditions, and essentially all conversion observed in the catalytic reactions results from the deoxygenation of CO<sub>2</sub>.

Considerably higher turnover numbers are achieved at higher reaction temperatures. Turnover of pinacolborate 2 presumably occurs much more rapidly; the boryl complex 1, generated in situ, is sufficiently stable toward decomposition to react productively with CO<sub>2</sub>. The reduction of CO<sub>2</sub> at 100 °C, using 0.1 mol % (*IPr*)-Cu(Ot-Bu) precatalyst, results in complete conversion of pinB– Bpin to pinB–O–Bpin after 20 h, corresponding to 1000 catalytic turnovers per copper.

To achieve more rapid catalytic turnover under mild conditions, the bulky *IPr* was replaced by the less sterically demanding *ICy* (1,3-dicyclohexylimidazol-2-ylidene) as a supporting ligand for copper. The complex (*ICy*)Cu(Bpin), generated in situ, is more prone to thermal decomposition than **1**, and catalytic reactions of CO<sub>2</sub> with pinB–Bpin using 1 mol % (*ICy*)Cu(Ot-Bu) display only 81% conversion at ambient temperature, with visible precipitation of copper metal after less than 1 h. However, when the reaction is run at 0 °C for 30 min and then at ambient temperature for 30 min, complete conversion of pinB–Bpin to pinB–O–Bpin is observed. This turnover frequency, corresponding to 100 turnovers within 1 h, is dramatically higher than that achieved using the *IPr* supporting ligand.

In summary, we have achieved the catalytic reduction of CO<sub>2</sub>

to CO in homogeneous solution, with high turnover numbers and frequencies, depending on the reaction conditions and supporting ligand. Both the oxygen abstraction and the catalyst turnover involve well-defined reactants and products, facilitating further study and pointing the way to future advances in catalytic reactions of CO<sub>2</sub>.

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**Supporting Information Available:** All experimental procedures; complete ref 3; structural parameters for **1** and **2**. This material is available free of charge via the Internet at http://pubs.acs.org.

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