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### **Enantioselective Small Molecule Synthesis by Carbon Dioxide-Fixation using a Dual Brønsted Acid/Base Organocatalyst**

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#### **Abstract**

Carbon dioxide exhibits many of the qualities of an ideal reagent – it is nontoxic, plentiful, and inexpensive. Unlike other gaseous reagents, however, it has found limited use in enantioselective synthesis. Moreover, unprecedented is a tool that merges one of the simplest biological approaches to catalysis – Brønsted acid/base activation – with this abundant reagent. We describe a metal-free small molecule catalyst that achieves the three component reaction between a homoallylic alcohol, carbon dioxide, and an electrophilic source of iodine. Cyclic carbonates are formed enantioselectively.

#### **Graphical Abstract**



The current global economic and environmental landscape has accelerated research into carbon dioxide  $(CO<sub>2</sub>)$  capture and storage  $(CCS)$  technology across a broad range of chemical disciplines. The most notable advancements have been made in the areas of materials chemistry,<sup>1</sup> carbon storage engineering, and alkylamine–based "scrubbing" systems.<sup>2</sup> The threat of car-bon dioxide accumulation as a greenhouse gas has motivated these sequestration strategies, but this gaseous reagent holds immense potential value as an abundant and nontoxic C1-building block<sup>3</sup> for carbon-carbon bond formation and carbonheteroatom functionalization reactions in chemical synthesis.<sup>4,5</sup> Unfortunately, the underlying features that contribute to car-bon dioxide's low general toxicity and ease of handling render it relatively inert as a chemical reactant.<sup>6</sup> This is punctuated by the contrasting abundance of *enantioselective* chemical reactions using hydrogen (H<sub>2</sub>),<sup>7</sup> oxygen  $(O_2)$ ,<sup>8</sup> and even carbon monoxide  $(CO)$ .<sup>9</sup> Chemical technologies that preferentially form one

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Supporting Information

Experimental procedures and spectroscopic data for all new compounds, and X-ray data (cif) for **2a**. This material is available free of charge via the Internet at [http://pubs.acs.org.](http://pubs.acs.org)

handedness (enantiomer) of a chiral product have direct application to drug development and new materials. Despite the virtues of high temperature and/or pressure to address poor reactivity, transformations employing  $CO<sub>2</sub>$  as a reagent<sup>10</sup> are typically limited to either Lewis basic substrates with sufficient nucleophilicity to react with the poorly electrophilic  $CO<sub>2</sub>$ <sup>3</sup> or metal-based reagents to increase the rate of  $CO<sub>2</sub>$  incorporation, often through a metal carboxylate intermediate (Figure 1A). $^{11,12}$  We sought a reagent, ideally a catalyst, that could both over-come these barriers to reactivity and/or unfavorable equilibria while simultaneously controlling stereoselection – in essence, a catalyst that could stabilize a substrate-CO<sub>2</sub> adduct, but still activate this adduct toward subsequent carbon-oxygen bond formation. Unprecedented is the use of a metal-free catalyst to stabilize the adduct of a weak nucleophile with carbon dioxide, such as a carbonic acid-base complex, while effectively guiding it toward enantioselective carbon-oxygen bond formation. Metal-based systems include CO<sub>2</sub> insertion into activated epoxides generating almost exclusively 5-membered cyclic carbonates, including enantioselective kinetic resolutions (Figure 1A).<sup>13</sup> Additionally, Yamada has reported a silver(I)-based alcohol desymmetrization using carbon dioxide at high pressure to pre-pare five-membered cyclic carbonates.<sup>14</sup>

We posited that a properly balanced Brønsted acid-Brønsted base bifunctional catalyst might lower the barrier to  $CO<sub>2</sub>$  incorporation and/or assist in the stabilization of the resulting adduct<sup>15</sup> as a prelude to its use as an oxygen nucleophile in a subsequent enantioselective carbon-oxygen bond-forming step.<sup>16</sup> If this could be achieved using a metal-free catalyst – an organocatalyst – the virtues of minimalism (symmetrical catalyst, low temperature, atmospheric pressure, near-neutral pH) would apply, suggesting broad impact. In this report, we validate this design by the development of a carboxylation/alkene function-alization reaction of homoallylic alcohols to produce chiral cyclic carbonates.

Homoallylic alcohol **1a** became the basis for developing a tandem alcohol carboxylation– alkene iodocarbonation reaction due to the lower steric demand of a 1,1-disubstituted alkene. The standard reaction to which others are compared involved chilling (−20 °C) a toluene solution of homoallylic alcohol (**1a**, 0.4 M) prior to addition of *N*-iodosuccinimide (Table 1, entry 1) and carbon dioxide (balloon). These catalyst-free conditions returned starting material following a 48 h reaction period. Compared to an otherwise identical reaction, addition of a strong base (sodium hydride, Table 1, entry 2) delivered the desired cyclic carbonate, but in only 16% yield.<sup>17</sup> Substitution of a more polar solvent (e.g. THF) for toluene increased the yield marginally (Table 1, entry 3). Based on precedence for Brønsted basic amines to react directly with carbon dioxide, or  $CO<sub>2</sub>/H<sub>2</sub>O$  combined, several amine bases were examined (Table 1, entries 4–5), as well as hydrogen bond-donor/acceptor amines (e.g. TBD, Table 1, entry 6) in an attempt to accelerate the desired reaction.<sup>18, 19</sup> These extensive attempts generally provided three outcomes: 1) return of unreacted homoallylic alcohol, 2) formation of apparent iodoetherification products, or simply 3) low yields (<15%) of the desired carbonate. Similarly, good hydrogen bond-donors, such as TFA or thiourea **3** <sup>20</sup> (Table 1, entries 7–8) failed to deliver any significant amount of carbonate.

Our ultimate goal was to explore the ability of a Brønsted ac-id/base combination to promote the reaction. Use of pyrroli-dine-substituted bis(amidine) **7** ('PBAM') resulted in an 18% yield of **2a**, however the carbonate was formed in a promising 39% ee (Table 1,

entry 9) at −20 °C. The analogous catalyst incorporating *trans*-stilbene diamine ('StilbPBAM' (**8**)) instead of *trans*-cyclohexane diamine provided the product in 33% yield and similar ee (36% ee, Table 1, entry 10). It was noted in these early experiments that the addition of molecular sieves (MS 4A) resulted in more consistent reactions as judged by conversion and/or yield (Table 1, entry 11). In reactions with-out sieves, formation of a precipitate appeared to correlate with lower yields, and varying enantioselectivity (particularly with free base (**7** or **8**; *vide infra*). Exploration of strong Brønsted acid additives (1:1 ligand:acid) led to moderate differences in enantioselection (Table 1, entries 12–15), with catalyst complex **8**·HNTf<sub>2</sub> providing product with 91% ee (Table 1, entry 15). Some sensitivity of both yield and selectivity to concentration was noted, $21$  with lower concentrations leading to depressed yield and selectivity (Table 1, entries 16–17). We reinvestigated several achiral amine bases under these optimized conditions, with only marginal improvement in yield (Table 1, entries 18–21). Attempts were also made to simulate the Brønsted acid/base character of catalyst **8**·HNTf<sub>2</sub> using monobasic amines in combination with varying amounts of Brønsted acid (e.g. Table 1, entry 21), none resulting in significant improvement of yield. Collectively, these results suggest an underlying order to the hydrogen bonding network in the key selectivity-determining step, if not unique reactivity associated with the proper positioning of a Brønsted acid and base in the same molecule, in this carbon dioxide–fixating reaction.

Application of conditions optimized for homoallylic alcohol **1a** to a range of similar substrates is outlined in Table 2. α-Substituted styrene derivatives were scrutinized using the mild conditions developed (**2a–m**, Table 2). Nominal substitution of the aromatic ring led to equally positive outcomes, with **2a–2c** formed in 91–95% ee, and high chemical yield (82– 96% yield) (Table 2, entries 1, 3–5). Substitution near the alkene was not tolerated, as no substrate conversion was observed to produce **2d** or **2e** (Table 2, entries 6–7). Increasing the reaction temperature to 0 °C led to complex mixtures suggestive of competing intermolecular iodoetherification. However, a β-naphthyl-substituted alkene led to good enantioselection and yield (**2f**, 90% ee, 88% yield) (Table 2, entry 8). Anisole derivatives (*meta*-and *para*-substituted) provided generally good enantioselectivity (80–90% ee) and higher chemical yield (97%, **2h**), but lower yield for **2g** (Table 2, entries 9–10). Halogensubstituted arenes led to a range of results, mostly related to reactivity; while selectivity remained high, some reached only partial conversion de-spite extended reaction times. Halogen substitution *meta* and *para* to the alkene provided consistently good enantioselection (87–90% ee) (Table 2, entries 11–14). Reactivity varied greatly among **2i– 2l**, however, and suggested that the alkene nucleophilicity might be a key determinant of reactivity. Carbonate **2m** was produced in nearly quantitative yield and 91% ee (Table 2, entry 15).

Alkenes bearing aliphatic substituents are often regarded as challenging substrates in stereoselective difunctionalization reactions.<sup>22</sup> 3-Alkyl butenols were prepared and converted to carbonates **2n–2p** with promising levels of enantioselectivity (up to 74% ee) (Table 2, entries 16–18). Iodocarbonate **2o**, derived from the sterically–unencumbered 3 methyl-but-3-ene-1-ol, a widely available isoprenyl feedstock, formed in a moderate 68% ee. Although not a focus of these investigations, allylic alcohols reacted sluggishly but

exhibited good yield and lower enantioselection. In an effort to probe the adaptation of this method to larger amounts, a gram-scale experiment using 3 mol % catalyst led to the carbonate in 89% ee and 79% yield (Table 2, entry 2). Finally, spirocyclic carbonate **2q** was prepared from the corresponding trisubstituted alkene in moderate yield (63%) and encouraging enantioselection (69% ee) (Scheme 1).

Several experimental observations are worth mention in ad-dition to the trends summarized above. Foremost among these, use of a carbon dioxide balloon attached to a degassed reaction established that the steady-state carbon dioxide concentration is significant in chilled toluene and could be reached within 40 minutes – far shorter than the time to complete conversion to carbonate (monitored by in situ IR). The rate of  $CO<sub>2</sub>$  absorption was affected insignificantly by most every factor examined, including temperature, the presence of MS 4A, and stirring rate. The correlation between high chemical yield and molecular sieves can be explained by the formation of a complex between the catalyst, adventitious water, and carbon dioxide, which we hypothesize to be the carbonic acid salt (Figure  $2^{2^3}$ . This complex precipitates from the reaction mixture when using **8**·HNTf2, but its formation appears reversible, reverting to active catalyst when a dessicant (MS 4A) and dry gas (argon) are added. Although a non-covalent complex is hypothesized in our work, some nucleophilic amines can form a covalent adduct with  $CO_2$ .<sup>24</sup> When not in competition with water, the alcohol substrate can entrain carbon dioxide, forming an intermediate and transient alkyl carbonic acid salt with the bifunctional catalyst. This intermediate, in reaction with NIS, forms a complex which then collapses to the cyclic carbonate either stepwise or directly.<sup>25</sup>

While cyclic carbonates are valuable in their own right,  $2<sup>6</sup>$  and they are prepared here using carbon dioxide as a phosgene surrogate, Scheme 2 details several notable transformations. First, reduction by stannane provided carbonate **9** in 86% yield when applied to iodocarbonate **2a**. Straightforward carbonate hydrolysis with a basic resin in methanol led to the versatile epoxide **10**. This is particularly significant from the viewpoint that carbon dioxide is effectively used as an equivalent to epoxidation, which normally requires an electrophilic source of oxygen (e.g. a peracid or dioxirane). Full reduction employing a stronger reducing agent (LiAlH4) leads to tertiary alcohol **11** in 71% yield. This carbon dioxide fixation method therefore offers a simple two step equivalent to metal-free oxidations of homoallylic alcohols,  $27$  for which carbon dioxide is converted to either dialkyl carbonate or methanol.

In summary, a mild and operationally straightforward carbon dioxide fixation reaction has been developed using a dual Brønsted acid/base catalyst that presents hydrogen bond-donor and acceptor functionality to activate and orient substrates in an enantioselective reaction. This metal-free method employs relatively weak nucleophiles (homoallylic alcohols) in the CO2 fixation step, generating transient acids that add to an alkene in combination with *N*iodosuccinimide. The catalysts deployed here use the virtues of Brønsted acid/base activation *alone* to achieve highly enantioselective carbonate synthesis.<sup>28</sup> From a different viewpoint, this carbon dioxide fixation method circumvents approaches dependent on phosgene29 as a source of carbonate protecting group for 1,3-diol prepared through

stereoselective synthesis. Numerous enantioenriched small molecules might be prepared using carbon dioxide as a source for carbon-oxygen bond formation.

#### **Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

#### **Acknowledgments**

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#### **Figure 1.**

Enantioselective methods using  $CO_2$  as a reagent, contrasting state-of-the-art metalmediated reactions with this report of Brønsted acid/base catalysis.



**Figure 2.** 

Catalyst Inactivation Pathway and its Recovery Using MS 4A



**Scheme 1.**  Iodocarbonation of a Trisubstituted Alkene.

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#### **Scheme 2.**

Conversions of Carbonate Products (Conservation of Enantiomeric Excess (ee) Observed in All Cases)

**Table 1**

Development of an Enantioselective CO<sub>2</sub>-Capture Reaction using a Homoallylic Alcohol Development of an Enantioselective CO2-Capture Reaction using a Homoallylic Alcohol







 $\overline{1}$ 

 $^d\!$  Catalyst prepared as the 1:1 acid salt, except entry 21. *a*Catalyst prepared as the 1:1 acid salt, except entry 21.

 $b$ MS denotes molecular sieves 4A, employed at a concentration of 1 g/mmol relative to the alcohol. *b*MS denotes molecular sieves 4A, employed at a concentration of 1 g/mmol relative to the alcohol.

*c*Isolated yield.

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 $d$  Enantiomeric excess (ee) determined by HPLC using a chiral stationary phase. Reactions are 0.4 M toluene unless otherwise noted. *d*Enantiomeric excess (ee) determined by HPLC using a chiral stationary phase. Reactions are 0.4 M toluene unless otherwise noted.

 $R$ eaction temp = 0 °C. 3-Methyl-3-buten-1-ol was converted in 60% yield under identical conditions. *e*Reaction temp = 0 °C. 3-Methyl-3-buten-1-ol was converted in 60% yield under identical conditions.

 $f_{\rm 20\,mol}$  % catalyst employed.  $f_{\rm 20\ mol}$  % catalyst employed.

 $^8S$  mol % catalyst employed, with results analogous to those using 10 mol % catalyst. *g*5 mol % catalyst employed, with results analogous to those using 10 mol % catalyst.



# **Table 2**

Initial Scope of an Enantioselective CO2-Capture Reaction using a Homoallylic Alcohol *a*



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for **2a** assigned using X-ray analysis, remaining examples assigned by analogy.

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 $b$  6.8 mmol (1.0 g) substrate was employed under the optimized conditions (1 atm CO<sub>2</sub>) utilizing 3.0 mol % catalyst for 48 h. *b*6.8 mmol (1.0 g) substrate was employed under the optimized conditions (1 atm CO2) utilizing 3.0 mol % catalyst for 48 h.

 $\emph{c}_{10\;\rm{mol}}$  % catalyst loading. *c*10 mol % catalyst loading.

 $d$  Reaction temperature was –50  $^{\circ}{\rm C}.$ *d*<br>Reaction temperature was −50 °C.

 $^e\rm{It}$  was noted that purified  $2\rm{g}$  was prone to decomposition. *e*It was noted that purified **2g** was prone to decomposition.