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Electrosynthesis of Cyclic Carbonates from Epoxides and Atmospheric Pressure Carbon Dioxide†

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The use of CO₂ for the preparation of value-added compounds has dramatically increased due to increased global warming concerns. We herein report the first example of a direct copper cathode/magnesium anode system that effectively converts epoxides and carbon dioxide to cyclic carbonates under mild electrochemical conditions at atmospheric pressure.

Carbon dioxide is arguably the main contributor to the increased concentration of gases within the atmosphere, in fact since the dawn of the industrial revolution CO₂ concentration within the atmosphere has annually increased. The main cause being identified as the combustion of fossil fuels. Methods that involve reducing the amount of CO₂ released into the atmosphere from industrial processes are therefore of significant value. Current commercial methods involve CO₂ capture via absorption using amine based materials post combustion. Pre-combustion methods also exist in which fuel is firstly converted into H₂ and CO₂ followed by a capture process. Once captured storage of CO₂ is achieved by storing it in unused oil/gas wells or under the ocean. This is a highly energy intensive process and unlike other waste streams CO₂ has a variety of possible uses such as a C₁ building block in organic synthesis.

To chemists carbon dioxide presents itself as a cheap, non-toxic and highly abundant carbon source, and if effectively activated can allow for carboxylation of organic molecules. Therefore fixation of CO₂ into organic molecules is an area of increasing interest and offers a much more environmentally sound alternative to current storage solutions. Currently, the greatest obstacle for establishing industrial processes based on CO₂ as a raw material is that in general a large energy input is required to transform CO₂.

There are several useful products which can be prepared from CO₂, for example polycarbonates which are also synthesised from epoxides and CO₂ are commercially important as they have several applications, for example as eyewear lenses and exterior automotive components. Perhaps the two most effective solutions have been reported independently by Coates and Lee. Coates zinc based catalyst 1 has been employed in the copolymerisation of cyclohexene oxide and CO₂ and Lee’s salen cobalt catalyst 2 has been used in the copolymerisation of propylene oxide and CO₂ (Figure 1). Cyclic carbonates, which are also produced from epoxides and CO₂, are widely used in the manufacture of products including solvents, paint-strippers, biodegradable packaging, as well as having other applications in the chemical industry. Again, however, these processes do require high energy input, including high temperatures, high pressures and the use of purified carbon dioxide.

Scheme 1. Current processes for CO₂ incorporation into epoxides.

There are a plethora of methods for the formation of cyclic carbonates from epoxides and CO₂, but recent efforts have been focused on the development of catalytic systems which are able to transform epoxides to cyclic carbonates at temperatures below 100 °C and at atmospheric pressure. Currently the only successful system is that reported by North and co-workers which employs a bimetallic salen complex such as 3 (Figure 1) and tetrabutylammonium bromide as a co-catalyst, although later developments have incorporated the co-catalyst into the salen substructure.

Fig. 1. Current catalysts for CO₂ incorporation into epoxides.

We were interested in developing a low energy alternative in line with our current research portfolio aimed at electrosynthesis coupled with semiconductor photoelectrodes to drive light assisted electrosynthetic reactions. We were therefore attracted to the possibility of employing an electrosynthetic system as this type of

† Electronic Supplementary Information (ESI) available: Experimental procedures and characterization of compounds 5a-k. See DOI: 10.1039/b000000x/
process could be designed to be cost neutral in terms of energy consumption if combined with a suitable solar powered energy source.  Duiñach has previously reported the electrochemical carboxylation of epoxides using a nickel(II) cyclam complex, and in a related process Yuan and co-workers have reported an electrochemical system using high CO₂ pressures. Both of these systems unfortunately have several drawbacks, including expensive potentially toxic catalysts, non-user friendly solvents and in the latter case high CO₂ pressures. We believed that it should be possible to deliver a step change in this technology by the development of an electrochemical system which is easy to set up, cheap, reliable, requires no expensive catalysts, runs under atmospheric CO₂ pressures and at ambient temperatures. In order to achieve this goal we set about screening a range of electrode materials for the electrosynthetic incorporation of CO₂ into our test substrate styrene oxide 4a (Table 1). The reactions were run in a single compartment cell, with acetonitrile as solvent, tetraethylammonium bromide as electrolyte and 60 mA current. We were delighted to find that good to excellent conversions to cyclic carbonate could be achieved under a variety of conditions (Table 1). Excellent conversion to cyclic carbonate 5a was achieved using the copper cathode/magnesium anode combination (Table 1 entry 1). It is important to note that in the absence of applied current we observed little to no cyclic carbonate (3%) and only recovered unreacted epoxide. We also did not observe the formation of any polymeric materials from the possible competing polycarbonate reaction or any other side products that have previously been observed for the corresponding reactions carried out at high temperature and/or pressure.

We next turned our attention to optimizing this mild system (Table 2), we found that high conversion to cyclic carbonate could still be achieved if we reduced the amount of tetrabutylammonium bromide to 1.0 equivalents (it is important to note here that we typically recover 90-95% of this material after the reaction by precipitation with Et₂OAc), however, below this level we observed only a 80% conversion (Table 2, entry 3). We also found that the use of tetrabutylammonium bromide is essential for the reaction to proceed, if we replaced the bromide counter-ion with tetrafluoroborate then we observed only 17% conversion to cyclic carbonate (Table 2, entry 5). The reactions can be carried out at room temperature (77% conversion after 6 h, Table 2, entry 7), however, for convenience mild heating to 50°C is employed (Table 2, entry 6); A combination of heating (50°C) and electrolysis (60 mA) affords the cyclic carbonate 2a in 99% conversion and 96% isolated yield. Reduced reaction times, for example, 3.5 h results in only 76% conversion (Table 2, entry 8).

Table 2 Optimization studies

<table>
<thead>
<tr>
<th>Entry</th>
<th>Supporting Electrolyte (equiv.)</th>
<th>T (°C)</th>
<th>t (h)</th>
<th>Conv. (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Bu₄NBr (2.0)</td>
<td>7 h rt</td>
<td>12 h 50 °C</td>
<td>19 &gt;99</td>
</tr>
<tr>
<td>2</td>
<td>Bu₄NBr (1.0)</td>
<td>7 h rt</td>
<td>12 h 50 °C</td>
<td>19 98</td>
</tr>
<tr>
<td>3</td>
<td>Bu₄NBBr (0.5)</td>
<td>7 h rt</td>
<td>12 h 50 °C</td>
<td>19 80</td>
</tr>
<tr>
<td>4</td>
<td>Bu₄NBBr (1.0)</td>
<td>25 °C</td>
<td>6</td>
<td>77</td>
</tr>
<tr>
<td>5</td>
<td>Bu₄NPF₆ (1.0)</td>
<td>50 °C</td>
<td>6</td>
<td>17</td>
</tr>
<tr>
<td>6</td>
<td>Bu₄NBBr (1.0)</td>
<td>50 ºC</td>
<td>6</td>
<td>99</td>
</tr>
<tr>
<td>7</td>
<td>Bu₄NBBr (1.0)</td>
<td>25 °C</td>
<td>6</td>
<td>77</td>
</tr>
<tr>
<td>8</td>
<td>Bu₄NBBr (1.0)</td>
<td>50 °C</td>
<td>3.5</td>
<td>76</td>
</tr>
</tbody>
</table>

General conditions: Cu cathode, Mg anode, CO₂ (1 atm, balloon), MeCN, single compartment cell, 60mA;  Conversion evaluated from the ¹H NMR spectrum by integration of epoxide vs cyclic carbonate peaks;  Conversion evaluated from the °H NMR spectrum by integration of epoxide vs cyclic carbonate peaks, isolated yield after column chromatography is shown in parenthesis; On average 90-95% of the Bu₄NBBr is recovered after each reaction by precipitation with Et₂OAc.

With our optimized conditions in hand we screened the applicability of this system over a range of substrate types (Table 3). We were delighted to find that excellent yields of cyclic carbonates were observed including electron rich and poor aromatic systems and a range of aliphatic epoxides were also effectively converted to the corresponding cyclic carbonate (these yields represent some of the highest reported to date). Sensitive functionalized substrates such as 4i-k, are also tolerated under the reaction conditions affording excellent yields of the corresponding cyclic carbonates 5i-k.

In order to explore the mild nature of these reaction conditions we were interested in the applicability of this system to enantiopure epoxides. Application of our optimized conditions to the CO₂ incorporation of (S)-styrene oxide afforded the cyclic carbonate in excellent yield (97%) with retention of configuration and only a slight loss of optical purity (99.5:0.5 e.r. to 98.6:1.4 e.r.).

In conclusion we have developed a powerful new tool in the combat against CO₂ emissions for the synthesis of value added compounds. This process is one of only a handful of approaches that allow this reaction to proceed at atmospheric pressure and at
ambient-mild temperatures. The yields obtained are comparable or better than those already reported in the area and our approach benefits from not employing an additional catalyst into the reaction, which in some cases can be toxic or expensive. The equipment required to perform this CO₂ incorporation reaction is cheap and should be readily available in any undergraduate teaching facility i.e. copper wire, magnesium ribbon and a power supply. We are currently looking at the mechanism (early indications show that it does not involve MgBr₂ but may well be related to that postulated by North and co-workers) and scalability of this process as well as applications of CO₂ incorporation towards other types of organic molecules and the development of a self-contained solar driven process.

Notes and references

1 Representative procedure for the synthesis of phenyl ethyle carbonate 5a. Styrene oxide (0.12 g, 1.0 mmol) and CO₂ (balloon) in acetonitrile (150 mL) were electrolysed (constant current: 60 mA) for 6 h in a single compartment cell (Mg anode and Cu cathode) containing BuNBr (0.32 g, 1.0 mmol) as supporting electrolyte at 50 °C. On completion the reaction mixture was washed with aqueous 0.1 M HCl (50 mL) followed by extraction with EtO₂ (3 x 35 mL). The combined organic extracts were then dried over MgSO₄ and evaporated under reduced pressure to afford a amber oil, which was suspended in EtOAc (100 mL). After 1 h the precipitated BuNBr (0.30 g, 95%) was removed by filtration and the solvent evaporated under reduced pressure to afford an amber oil. This crude material was purified by column chromatography on silica gel eluting with ethyl acetate:light petroleum. Colourless solid (0.157 g, 96%).


11 For example, isomerisation of the epoxide to ketone and aldehydes or addition of H₂O to afford the ring opened diol: see: J.-Q. Wang, X.-D. Yue, F. Cai, L.-N. He, Catal. Commun. 2007, 8, 167-172.

12 GC/MS and H1,12C NMR data revealed only the reaction products and BuNBr.

13 In a independent reaction addition of MgBr₂ to the reaction without electrolysis resulted in complete recovery of starting materials.

14 The electrode materials required for large scale applications are currently commercially available. Mg anodes are used in the protection of ship hulls. See for example: http://www.corpro.co.uk/pdf/Anodes/Magnesium/Magnesium-Anodes.pdf

15 Spectroscopic data for phenyl ethylene carbonate 5a were consistent with those reported in the literature: see: J.-L. Jiang, F. Gao, R. Hua, X. Qiu, J. Org. Chem. 2005, 70, 381-383.