Divergent methods for polyester and polycarbonate depolymerization with a cobalt catalyst†

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A pyridine diimine cobalt catalyst series is used for the cyclodepolymerization (CDP) of several different polyesters and polycarbonates. In the presence of isopropanol, a wide range of polymers can undergo solvolytic depolymerization. CDP of poly(propylene carbonate) revealed an unzip back-biting or random scission mechanism, depending on what conditions were used. The first example of a method capable of solvolytic and CDP on a polymer mixture in the same pot is identified.

Introduction

Polyesters and polycarbonates are important materials for a wide range of applications, such as CDs, water bottles, and packaging.‡ Additionally, emerging methods to make these materials with a range of physical properties has established a growing movement to use these polymers as degradable, recyclable or compostable replacements for non-degradable polymers made on alarming scales.§ As technology improves to increase industrial use of these materials, it is critical to establish options for their end of life.

Many polyesters and polycarbonates can be made from biofeedstocks.‖ Polyesters are often considered degradable or compostable. However, these strategies often need industrial composting, which requires an additional collection stream.§‡ As technology improves to increase industrial use of these materials, it is critical to establish options for their end of life.

Many polyesters and polycarbonates are made from biofeedstocks.‖ Polyesters are often considered degradable or compostable. However, these strategies often need industrial composting, which requires an additional collection stream.§‡ As technology improves to increase industrial use of these materials, it is critical to establish options for their end of life.

These materials can also be chemically recycled, adding another strategy to recover valuable chemicals that can be used to remake the polymer directly.‖ For polyesters and polycarbonates, there are four primary methods for chemical recycling: hydrogenation, transfer hydrogenation, solvolytic depolymerization, and cyclodepolymerization (Fig. 1 and 2).‡§§ Each of these methods have their advantages and challenges, and some methods are better for specific polymers.

Hydrogenation often breaks polyesters and polycarbonates down into diols, which are useful for numerous applications, including synthesizing new polymers.‡‡–‡‡‡ However, high pressures and temperatures are often used, posing safety and energy challenges. A safer alternative, albeit less atom economical, is transfer hydrogenation, which uses a hydrogen source like isopropanol to generate an equivalent of H2 and acetone as a by-product. Though apart from numerous examples of the transfer hydrogenation of lignin, few reports exist documenting the transfer hydrogenation of polymers.‡‡‡–‡‡‡§

Solvolytic remains one of the most studied methods for the depolymerization of step-growth polymers, particularly for poly(ethylene terephthalate) (PET).‡§‡‡ The use of water, methanol or ethylene glycol can lead to linear molecules that can be used again for polymerization.‡‡‡† Hydrolysis is relevant to many polyesters and polycarbonates, however these materials are often insoluble in water and need harsh conditions to reach efficient depolymerization.‡‡‡§‡ Methanolation and glycolysis may be more useful depolymerization methods, as is the case with PET, in which the product may regenerate new PET.‡‡‡§‡‡‡ Finally, the cyclodepolymerization (CDP) of polyesters and polycarbonates back to their cyclic precursors is an emerging area of research, as this method often applies to chain growth polymers, which are receiving much interest as replacements for non-degradable polymers. In many cases, the ceiling temperature of the polymer makes a big difference on how cyclodepolymerization can occur.‡‡‡§‡‡‡‡ Additionally, high dilution of the reaction often allows for higher conversion to
cyclic monomer, albeit with the crucial flaw of requiring large amounts of solvent.36,37 Recently, reports of manipulating ceiling temperature combined with reactive distillation methods have prevented the need for high dilution.38–40 Namely, Byers and co-workers reported exciting catalysts for the CDP of many polyesters and polycarbonates at high yields using a reactive distillation method.40

CDP of polycarbonates can take two forms: either formation of cyclic carbonates or complete depolymerization to epoxides and CO₂ (Fig. 2a).9 There are few examples of CDP of commercial polycarbonates and polystyrene to their respective cyclic monomers.31–44 In this case, CDP of the polycarbonate often leads to a cis-cyclic carbonate, a product which lacks examples of being polymerized. Many of these examples have shown the ability to perform CDP on specially designed polymers, which have been modified or synthesized with the intention that these polymers also cyclodopolymerize to polymerizable monomers.26,28,29,31,33,34,38,45–49 Similarly, scant examples of complete CDP to epoxides and CO₂ exist in the literature.50–53 To date, there is still not a clear understanding for what catalysts lead to selectivity of either form of CDP, although computational studies have helped rationalize a catalyst’s ability to lower the barriers for a specific method.49 CDP of polystyrenes has one main pathway to generate cyclic esters (Fig. 2b). Examples of CDP for common polyesters, such as poly(ε-caprolactone) (PCL), poly(δ-valerolactone) (PVL), polydioxanone (PDO), poly(caprolactam) (nylon-6), and poly(lactic acid) (PLA) are known with metal catalysts.35,40,44,45,54

Herein, we demonstrate the use of a pyridine diimine cobalt catalyst that can catalyse the cyclodepolymerization of several different polyesters and polycarbonates. In the presence of isopropanol, these catalytic methods also facilitate solvolytic depolymerization of seven different polyesters and polycarbonates, which is attributed to the base used in the reaction. To our knowledge, this is the first example of a single catalyst system demonstrating these two depolymerization strategies. We demonstrate proof of concept for achieving both methods in the same pot, which addresses considerations of mixed polymer waste.

Results and discussion

Cobalt complexes with pyridine diimine (PDI) ligands were first targeted for the depolymerization of carbonyl-containing polymers, as these complexes are easy to make and similar complexes with chiral ligand analogues have been used for stereoselective transfer hydrogenation of small molecules.55,56 Additionally, PDI cobalt complexes have been synthesized and modified regularly for application in electrocatalysis, providing a toolbox of electronic and steric modifications for these molecules.57,58 Initially, catalysts with simple substituents were targeted, with subtle modifications in steric crowding to probe the sensitivity of these catalysts for the depolymerization of polymers (Fig. 3). The PDI ligands were varied to contain mesityl (L-MES), di-isopropyl phenyl (L-DIPP), and di-ethyl
phenyl (L-DEP) substituents. The catalysts MES and DIPP were synthesized according to literature procedure while DEP was synthesized by and methods and was fully characterized (see \textit{ESI}).\textsuperscript{59,60}

Using transfer hydrogenation conditions of poly(propylene carbonate) (PPC), inspired by the recent work from Werner and coworkers, these cobalt complexes did not show any evidence for the formation of the expected transfer hydrogenation products: propane diol and methanol.\textsuperscript{15} Instead, exclusive formation of propylene carbonate (PC) was observed for all three catalysts, with reactions yielding PC, cyclodepolymerized from PPC, within 20 h at 140 °C (Table 1, entries 1–3).

Table 1  Depolymerization of PPC results after varying catalysts, base, solvent and hydrogen-donor presence\textsuperscript{a}

<table>
<thead>
<tr>
<th>Entry</th>
<th>Catalyst</th>
<th>Time (h)</th>
<th>KO\textsubscript{t}Bu (mol%)</th>
<th>Solvent(s) used</th>
<th>Solvent volume (mL)</th>
<th>% Yield\textsuperscript{b,c}</th>
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<tbody>
<tr>
<td>1</td>
<td>MES</td>
<td>20</td>
<td>5</td>
<td>iPrOH : THF</td>
<td>8 : 2</td>
<td>66</td>
</tr>
<tr>
<td>2</td>
<td>DIPP</td>
<td>20</td>
<td>5</td>
<td>iPrOH : THF</td>
<td>8 : 2</td>
<td>66</td>
</tr>
<tr>
<td>3</td>
<td>DEP</td>
<td>20</td>
<td>5</td>
<td>iPrOH : THF</td>
<td>8 : 2</td>
<td>64</td>
</tr>
<tr>
<td>4</td>
<td>MES</td>
<td>20</td>
<td>5</td>
<td>THF</td>
<td>6</td>
<td>&gt;99(0)</td>
</tr>
<tr>
<td>5</td>
<td>DIPP</td>
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<td>5</td>
<td>THF</td>
<td>6</td>
<td>78(19)</td>
</tr>
<tr>
<td>6</td>
<td>DEP</td>
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<td>5</td>
<td>THF</td>
<td>6</td>
<td>97(3)</td>
</tr>
<tr>
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<td>10</td>
<td>THF</td>
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</tr>
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<td>0</td>
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<tr>
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<td>0</td>
<td>PC</td>
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<tr>
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</tr>
<tr>
<td>21</td>
<td>DEP</td>
<td>6</td>
<td>0</td>
<td>THF</td>
<td>6</td>
<td>52(9)</td>
</tr>
<tr>
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<td>None</td>
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<td>89(1)</td>
</tr>
<tr>
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<td>None</td>
<td>10</td>
<td>5</td>
<td>THF</td>
<td>6</td>
<td>41</td>
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\textsuperscript{a} Conditions: 5 mol% catalyst with respect to the repeat unit molar mass of the polymer, 2 mmol PPC, run at 140 °C and 350 rpm in Parr reactor. Reactions with KO\textsubscript{t}Bu were performed in an air-free, N\textsubscript{2} environment. All other entries were performed exposed to air. \textsuperscript{b} Determined by comparing \textsuperscript{1}H NMR spectroscopy using mesitylene as internal standard, taken in CDCl\textsubscript{3}. \textsuperscript{c} Parentheses next to yields indicate reactions performed in duplicate; standard deviation of error in parentheses. \textsuperscript{d} NMR calculation used in place of internal standard yield calculation due to near method’s challenges in collecting full product.

Cyclodepolymerization of poly(propylene carbonate)

CDP of PPC has not been extensively studied; current reports consist of various catalysts’ ability to perform CDP on PPC to form PC.\textsuperscript{40–43} A tris(pentafluorophenyl) borane catalyst was shown to perform selective CDP with poly(cyclohexene carbonate) and PPC using CH\textsubscript{2}Cl\textsubscript{2} and 5 mol% catalyst (with respect to the repeat unit molar mass of the polymer) at 130 °C.\textsuperscript{41} The kinetics of CDP of PPC were explored with a chromium salen complex and an ammonium azide co-catalyst, which identified the importance of a nucleophilic or basic anion to initiate CDP.\textsuperscript{42} Finally, ZnEt\textsubscript{2} has commonly been used as a CDP catalyst for many designer polymers, and it can also depolymerize PPC (see \textit{ESI}).\textsuperscript{40,61} Unfortunately, the CDP of PPC in literature is difficult to compare across works, as conditions involving catalyst, solvent, reaction times, etc. vary substantially.

Notably, the PPC reaches a temperature threshold of 170 °C, at which point, the polymer is known to degrade/depolymerize into PC.\textsuperscript{61,62} For this polymer, this temperature allows for the conversion of the kinetic polymer product (PPC) to the thermodynamic cyclic carbonate (PC).\textsuperscript{7,8,42}

Once CDP selectivity was identified for these cobalt catalysts, efforts to optimize the reaction conditions for this method over transfer hydrogenation was prioritized. A tetrahydrofuran (THF)/isopropanol (iPrOH) mixture with potassium...
tert-butoxide (KO\textsubscript{t}Bu) was used to replicate Werner’s transfer hydrogenation conditions.\textsuperscript{15} THF was used to solubilize the polymer, isopropanol as the hydrogen source, and KO\textsubscript{t}Bu as a strong base.\textsuperscript{15} These conditions may not be necessary to produce PC. Without iPrOH, MES and DEP catalysts reached quantitative conversion within 20 hours at 140 °C, with DIPP displaying high but inconsistent yields of PC (Table 1, entries 4–6). These results indicate that the hydrogen source is unnecessary. CDP reactions conducted without this base yielded very similar results, if not slightly lower conversions, to those with base, save for that of DIPP, indicating base was not significantly impacting the rate of CDP (Table 1, entries 7–9).

When comparing the ligand substituents, the catalyst with mesityl (MES) and diisopropyl phenyl (DIPP) substituents showed slower CDP of PPC, while the catalyst with ethyl substituents (DEP) proved fastest and the most reproducible. It is unclear why MES is slower than DEP, however solubility could be a challenge, as these catalysts are only mildly soluble in THF at room temperature. Given DEP’s speed over DIPP and MES in the optimized conditions without iPrOH or base, this catalyst was used for further optimization. With DEP (Fig. 3), reactions shortened to 10 hours in THF—showed a distinct difference when KO\textsubscript{t}Bu was used (achieving 83% yield) versus when KO\textsubscript{t}Bu was not used (achieving just 69% yield) (Table 1, entries 10 and 11). These results indicate that presence of base does increase the rate of CDP when considering reactions that have not reached high conversions. It is anticipated that the initiation of the reaction occurs through the deprotonation of the polymer end group, which is likely an alcohol, or through nucleophilic attack of a carbonyl on the polymer chain (Fig. 4a). An O\textsubscript{Bu} alkoxide would likely be faster than a chloride for both proposed routes, supporting the higher yield of PC with KO\textsubscript{t}Bu present.

One hypothesis is that the base could exchange the Cl ligands with the –O\textsubscript{Bu} ligands, which might enhance nucleophilic attack of carboxyls in the polymer and increase the initiation rate of depolymerization. This reaction has been conducted on similar Fe complexes with PDI ligands.\textsuperscript{53} Notably, reactions with KO\textsubscript{t}Bu show a rapid color change from yellow brown to purple when solvent is added to the reaction mixture. This color change is not observed without the presence of KO\textsubscript{t}Bu. In this case, two equivalents of KO\textsubscript{t}Bu would be needed to exchange with all the chlorides on DEP. This indeed shows slightly faster conversion than DEP with one equivalent of KO\textsubscript{t}Bu (Table 1, entry 12). Additionally, stirring DEP with one or two equivalents of KO\textsubscript{t}Bu prior to the addition of PPC lead to even higher conversions, indicating the likelihood that exchanging the Cl anions for O\textsubscript{Bu} anions increases the reactivity of the DEP catalyst (Table 1, entries 13 and 14).

Unfortunately, the low solubility of DEP in most solvents at room temperature prevents the ability to characterize this exchange either by NMR spectroscopy or isolation in bulk. Future directions aim to adjust the ligands for increased solubility to confirm these hypotheses.

Once the metal catalyst has the polymer chain bound, the expected mechanism of CDP with metal-based catalysts is back-biting from alkoxide or carbonate end groups bound to the metal ion. These end groups could perform consistent back-biting reactions to directly form PC, depolymerizing the polymer through a controlled “un-zip” pathway (Fig. 4b). Furthermore, the end groups could perform transesterification reactions further down the polymer chain to form cyclic oligomers or on another polymer chain which changes the polymer molar mass (Fig. 4b). Indeed, these proposed mechanisms are well-documented across several PPC depolymerization studies.\textsuperscript{9,40,42,64} From these suspected routes, it was important to understand if reaction conditions could impact rate of CDP and control for the desired “un-zip” pathway.

Since KO\textsubscript{t}Bu is moisture-sensitive, reactivity of the catalyst without base was investigated, with the reactions pursued in the presence of air. DEP is air stable and can be left over a month without showing decomposition. Using the standard reaction conditions taken from Table 1, entry 11, batch reactions were performed to follow the progress of the reaction (Fig. 5). Yields were characterized by \textsuperscript{1}H NMR spectroscopy. The remaining PPC was also characterized by size-exclusion chromatography (SEC). Within the first 4 hours, only slow CDP is observed, while the rate rapidly increases, demonstrating a more linear trend. Once all catalysts are bound to polymer chains, this could explain the increase in rate. Before all polymer chains are deprotonated, chain transfer could occur and slow productive CDP. Following the reaction by SEC showed a consistent decrease in the dispersity of the remaining polymer chains. A duplicate reaction series shows the same general trend (see ESI†). Random nucleophilic attack on the polymer chain is observed, while the rate rapidly increases, demonstrating a more linear trend.
would likely result in broad and/or multimodal molar mass distributions, as there would be no preference for reacting with any particular part of the polymer chain. Monomodal molar mass distributions, along with the decrease in dispersity, suggest that these side reactions are not occurring, indicating selective “un-zip” backbiting reaction for CDP of PPC with PDICoCl₂ catalysts.

It was anticipated that the presence of base could lower the induction period, which would lead to productive CDP faster than without base. Other air-stable bases in addition to KO₆Bu were tested (Table 2). Two of the bases, NaOH and NaHCO₃, presented diminished yields in comparison to analogous conditions without base. NEt₃ showed a similar yield to that without base. Two bases (KOH and NaO₆Bu) resulted in comparable yields to that with KO₆Bu, without the need for inert conditions. These results indicate that a stronger base or easier ability to exchange the chlorides on DEP may be important to improve the reaction rate. Running time points with NaO₆Bu, analogous to those done without base in Fig. 5, showed inconsistent yields. Notably, reactions that did not reach full conversion with NaO₆Bu displayed a higher dispersity than the original PPC starting polymer (ESI Table S4†). Returning to KO₆Bu time points showed a more consistent increase in conversion over time. However, with this base, dispersity was variable, suggesting random chain scission. Interestingly, this similar pattern of variable dispersity with CDP is seen with use of ZnEt₂ from Byers et al.40 These results suggest that the base could be performing side reactions with or without the use of the catalyst, leading to random scission.

Table 2 Depolymerization of PPC results varying bases

<table>
<thead>
<tr>
<th>Entry</th>
<th>Base</th>
<th>Time (h)</th>
<th>% Yield⁶,c</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>KOH</td>
<td>10</td>
<td>79(8)</td>
</tr>
<tr>
<td>2</td>
<td>NaOH</td>
<td>10</td>
<td>58</td>
</tr>
<tr>
<td>3</td>
<td>NaHCO₃</td>
<td>10</td>
<td>24</td>
</tr>
<tr>
<td>4</td>
<td>NEt₃</td>
<td>10</td>
<td>61</td>
</tr>
<tr>
<td>5</td>
<td>NaO₆Bu</td>
<td>10</td>
<td>86(1)</td>
</tr>
<tr>
<td>6</td>
<td>KO₆Bu</td>
<td>10</td>
<td>83(3)</td>
</tr>
<tr>
<td>7</td>
<td>NaO₆Bu</td>
<td>2</td>
<td>40(14)</td>
</tr>
<tr>
<td>8</td>
<td>NaO₆Bu</td>
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<td>58(15)</td>
</tr>
<tr>
<td>9</td>
<td>NaO₆Bu</td>
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</tr>
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<td>NaO₆Bu</td>
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</tr>
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<td>NaO₆Bu</td>
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<td>KO₆Bu</td>
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<td>KO₆Bu</td>
<td>4</td>
<td>85(3)</td>
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<td>KO₆Bu</td>
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<td>16</td>
<td>KO₆Bu</td>
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⁶ Conditions: 5 mol% DEP and 5 mol% base with respect to the repeat unit molar mass of polymer, 2 mmol PPC, 6 mL THF, 350 rpm, run at 140 °C in Parr reactor. Reactions performed under air-free N₂ atmosphere if KO₆Bu used, otherwise performed in air.⁷ Determined by ¹H NMR spectroscopy using mesitylene as internal standard, taken in CDCl₃.⁸ Reactions performed in duplicate; standard deviation of error represented in parentheses.
and less controlled CDP. Control reactions with KO\textsuperscript{t}Bu alone did show mild conversions of PPC to PC, indicating they can facilitate CDP with and without the catalyst (Table 1, entry 23). These conditions contrast those seen with the unzipping mechanism without base. Though the chloride from DEP alone proved better at unzipping, this may be in part due to halides being weaker nucleophiles than alkoxides, making them more likely to attack end groups because of their proximity.

While THF was originally used, analogous to prior transfer hydrogenation reactions, the solvent scope was further tested to promote the fastest catalysis. Reactions performed in chlorobenzene, 2-methyl THF, and toluene did not facilitate rapid CDP, showing diminished yields to those conducted in THF (Table 1, entries 15–17). The use of PC or a PC/THF mixture both led to higher yields than those with just THF. This could be due to the higher boiling point of PC, which maintains solubility of the polymer and catalyst at elevated temperatures (Table 1, entries 18 and 19). These results suggested that the reaction could be done with polymer in the melt without added solvent, where the generated PC becomes the solvent over the course of the reaction. Gratifyingly, full conversion of PPC to PC was achieved within 10 hours when the reaction was conducted on PPC in the melt without solvent addition, in which no remaining PPC is observed by NMR spectroscopy (Table 1, entry 20). An 89% yield is achieved in just 6 hours with these neat conditions, compared to 52% yield with THF (Table 1, entries 21 and 22). This is the first example for CDP of PPC in the melt without a solvent. To ensure that presence of PC in the reaction does not lead to polymerizations, control reactions were conducted with DEP or KO\textsuperscript{t}Bu with PC as monomer and THF as solvent for 10 h at 140 °C, with no PPC formation observed (see ESI†). While reactions done with PPC in the melt could lead to greener methods for CDP, quantification of the products was more difficult if the reaction did not reach full conversion, therefore further studies were still conducted in the presence of solvent.

To identify how reaction concentration impacts CDP of PPC with DEP, the reaction concentrations were varied in two ways. In one method, the PPC and catalyst quantities were kept constant, while the THF volume was varied (Fig. 6a, blue). While this method showed no observable pressure differences on the Parr reactors as the volume of THF increased (maintaining a steady 6.0 bar), it was important to rule out any pressure contributions to the data. Therefore, the second method kept the THF volume constant, while the PPC and catalyst quantities were varied (Fig. 6a, orange). Both methods showed the same overall outcome, suggesting that pressure is not impacting the results. Four different concentrations were measured, with the most concentrated and second most dilute conditions showing the highest yields for CDP. The higher concentration reactions could show higher yields due to the catalyst initiating CDP faster, while also increasing the rate of the metal ion coming in proximity to the next carbonyl on the polymer chain. This condition also most closely matches conditions in the melt, which has shown the highest yield. The metal center could be performing chain transfer reactions between polymer chains. Dilution could prevent this side reaction. However, over-dilution could slow the metal from initiating CDP.

Next, varying the catalyst loading, while maintaining all other conditions, showed an expected increase in yield as the catalyst loading is increased from 0.5 to 5 mol% (Fig. 6b). However, when catalyst loading is increased to 7.5 mol%, a drop in yield is observed. It is unclear what causes this drop in yield, though, the added concentration of metal bound to polymer chain could lead to increased presence of chain transfer or the presence of transterification of other polymer chains, leading to lowered back-biting (Fig. 7). With smaller amounts of catalyst in fewer quantities than the polymers’ available end groups, the cobalt-alkoxide (or carbonate) may

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**Fig. 6** Graph of PC yield as a function of solvent dilution by varying THF volume (blue) or amount of catalyst and PPC relative to their dilution in THF while maintaining an overall volume of 6 mL THF (orange) (a), catalyst (DEP) loading (b), and temperature (c). Each experiment operated under conditions of 1–6 mmol PPC, 5 mol% DEP with respect to the repeat unit molar mass of the polymer, 0–18 mL THF, and 80–160 °C, run for 10 hours.

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Research Article

Cyclodepolymerization of other polymers

Once CDP of PPC was better understood, extension of studies to other polycarbonates and polyesters were warranted (Fig. 8). No observable CDP was identified for nylon 6 (7a), and small amounts of CDP was observed for PCL (2a) and poly(trimethyl-ene carbonate) (PTMC) (5a) with the DEP and THF, alone. Surprisingly, CDP of PVL was identified with DEP to form the δ-valerolactone (VL) monomer (3a). Under the standard reaction conditions provided from Table 1, PVL depolymerized with a 41% yield of VL, compared to the analogous reaction in THF alone, which yielded 7% VL. When run for 20 hours, the reaction yielded 73% product (11% by THF-only control), and when run at 180 °C for 20 h, the reaction yielded 84% product (10% by THF-only control). Prior studies have attempted depolymerization and/or thermal degradation of PVL at 250 °C, taking advantage of its relatively low ceiling temperature of 298 °C. Additionally, Byers and coworkers identify their ZnCl₂/PEG600 catalyst system to achieve 94% conversion under reactive distillation at 160 °C after 16 hours. Using DEP, we can identify high conversion without the need for reactive distillation. With reactive distillation, analogous to reactions done by Byers and coworkers, CDP of PVL at 180 °C with PEG600 for 10 hours achieved only 85% conversion. These results indicate that the DEP catalyst is more important than the presence of KO'Bu under these conditions. Similarly, poly(dioxanone) (PDO) displayed quantitative conversion under 180 °C after 20 hours (4a). Finally, PLA was converted to lactide (6a) in a moderate yield of 42%. While this is not optimized, and does not compete with the Sn(II)/alcohol catalyst system reported by Williams and coworkers, this represents one of only a few examples of CDP of PLA. Most notably, all polymers which were able to undergo any amount of CDP fared worse under the same conditions with KO'Bu added (see ESI†). These results indicate that the DEP catalyst is more important than the presence of KO'Bu for CDP.

Solvolysis

While DEP proved to be active for CDP of several selected polymers (under the conditions tried), it was still unclear whether transfer hydrogenation (the initial goal) was possible with this...
catalyst. Using the same conditions from Werner used in early PPC depolymerization trials, other polymers mentioned previously, such as PLA, PTMC, PDO, PECL, and nylon 6, in addition to poly(bisphenol A carbonate) (PBPAC) and PET, were tested. PET and PBPAC were not studied for CDP, as they are made through step-growth polymerization and are not known to have an accessible cyclic monomer. Almost all these plastics showed full depolymerization within 20 hours, though via solvolysis mechanism rather than transfer hydrogenation. The polymers and their depolymerized substrates are shown in Fig. 9. Through fairly mild conditions, DEP with KO$_t$Bu achieved complete solvolysis of a wide range of polymers, demonstrating its versatility. Control reactions with KO$_t$Bu without the catalyst showed similar conversions in most cases, suggesting the base to be the primary catalyst for this solvolysis method. In the case of PBPAC, the presence of catalyst shows greatly improved solvolysis. Additionally, reactions with PTMC with just KO$_t$Bu does not perform the solvolysis identified with the DEP. Instead, it produced a still undetermined product, with no evidence of the solvolysis product. Nonetheless, this simple base has not been previously used for solvolysis of polyesters or polycarbonates. However, the versatility for high conversions of solvolysis of many polymers under these reported conditions warrant additional studies to determine the full capability of this simple base. While these reaction conditions are not optimized, these results indicate this catalytic system is active for solvolysis of numerous polymers. Since solvolysis is not commonly done with isopropanol, additional studies will be needed to compare to other methods that prioritize hydrolysis, methanolysis, and glycolysis.

Since these two pathways seemed compatible with each other, we questioned whether they could be used in parallel in the same reaction mixture. PPC and PBPAC were selected to mix, as they are both polycarbonates, PPC shows exclusive CDP under all conditions studied, and PBPAC does not have an accessible cyclic monomer. Notably, in a mixed pot of PBPAC and PPC in a 1 : 1 molar ratio with respect to the repeat unit of the polymer, under conditions of DEP at 180 °C with KO$_t$Bu and isopropanol as solvents, CDP was maintained with PPC while solvolysis prevailed with PBPAC to bisphenol A (BPA) and diisopropyl carbonate (DIPC) (Fig. 10), with both
pathways reaching full conversion. Similarly, under conditions of just DEP with THF as solvent at 140 °C, PB PAC remained intact while PPC depolymerized to PC at 91% yield, which bodes well for isolating depolymerization products in mixed recycling streams. On the other hand, PVL under solvolysis conditions yield solvolysis products, indicating multiple depolymerization paths for one polymer with the same catalyst under different conditions.

Conclusions

A series of air-stable pyridine(diimine) cobalt dichloride catalysts were identified to be active for the selective cyclodepolymerization (CDP) of poly(propylene carbonate) (PPC) to form propylene carbonate (PC) as the exclusive product. Reaction conditions were optimized to reach high yields of PC when conducting the CDP reaction with neat polymer in the melt. Optimizations for the catalyst alone identified a controlled “un-zip” pathway to form PC, without the presence of undesirable side reactions. Alternatively, presence of a base encouraged random CDP of the polymer chain, with increases in dispersity of the polymer over the course of the reaction. Catalyst loading was found to be important, as too high or too low led to proposed inhibitive side reactions of chain transfer or transmetalation.

Finally, extensions to other carbonyl-containing polymers identified these catalysts as highly active for the selective CDP of poly(β-valerolactone) (PVL) and polydioxanone (PDO) with DEP alone. Activities of CDP for poly(trimethylene carbonate) (PTMC), poly(lactic acid) (PLA) and poly(e-caprolactone) (PCL) were observed in smaller amounts, however optimization of the conditions or the catalyst may be needed to achieve high conversion. In the presence of KO'Bu and iPrOH, DEP could promote solvolytic depolymerization of several commodity polymers. Control reactions identify the KO'Bu shows comparable activity on its own, suggesting it is the active catalyst in this reaction.

The results of these studies were used in a proof of concept mixed-recycling stream of PPC and poly(bisphenol A carbonate) (PB PAC), in which solvolysis conditions lead to the full conversion of PPC CDP and PB PAC solvolysis. CDP conditions without base or iPrOH led to selective CDP of PPC to 91% conversion, with the PB PAC being left undisturbed. These results identify promising routes for tunable deconstruction of these polymers. Future work to identify optimal changes to the metal and ligands to further improve CDP of carbonyl-containing polymers are currently underway.

Conflicts of interest

There are no conflicts to declare.

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