Switchable Catalysis Improves the Properties of CO₂-Derived Polymers: Poly(cyclohexene carbonate-8-ε-decalactone-8-cyclohexene carbonate) Adhesives, Elastomers, and Toughened Plastics

Gregory S. Sulley, Georgina L. Gregory, Thomas T. D. Chen, Leticia Peña Carrodeguas, Gemma Trott, Alba Santmartí, Koon-Yang Lee, Nicholas J. Terrill, and Charlotte K. Williams*

ABSTRACT: Carbon dioxide/epoxide copolymerization is an efficient way to add value to waste CO₂ and to reduce pollution in polymer manufacturing. Using this process to make low molar mass polycarbonate polyols is a commercially relevant route to new thermosets and polyurethanes. In contrast, high molar mass polycarbonates, produced from CO₂ generally under-deliver in terms of properties, and one of the most widely investigated, poly(cyclohexene carbonate), is limited by its low elongation at break and high brittleness. Here, a new catalytic polymerization process is reported that selectively and efficiently yields degradable ABA-block polymers, incorporating 6–23 wt % CO₂. The polymers are synthesized using a new, highly active organometallic heterodinuclear Zn(II)/Mg(II) catalyst applied in a one-pot procedure together with biobased ε-decalactone, cyclohexene oxide, and carbon dioxide to make a series of poly(cyclohexene carbonate-8-decalactone-8-cyclohexene carbonate) [PCHC-PDL-PCHC]. The process is highly selective (CO₂ selectivity >99% of theoretical value), allows for high monomer conversions (>90%), and yields polymers with predictable compositions, molar mass (from 38–71 kg mol⁻¹), and forms dihydroxyl telechelic chains. These new materials improve upon the properties of poly(cyclohexene carbonate) and, specifically, they show good thermal stability (T₅₀ ~ 280 °C), high toughness (112 MJ m⁻³), and very high elongation at break (>900%). Materials properties are improved by precisely controlling both the quantity and location of carbon dioxide in the polymer chain. Preliminary studies show that polymers are stable in aqueous environments at room temperature over months, but they are rapidly degraded upon gentle heating in an acidic environment (60 °C, toluene, p-toluene sulfonic acid). The process is likely generally applicable to many other lactones, lactides, anhydrides, epoxides, and heterocumulenes and sets the scene for a host of new applications for CO₂-derived polymers.

INTRODUCTION

Devising methods to efficiently use carbon dioxide in chemical synthesis is a fundamentally important challenge.¹⁻⁴ The ring opening copolymerization (ROCoP) of carbon dioxide and epoxides is a truly catalytic process that enables high CO₂ uptake and allows for the partial substitution of polluting and costly petrochemicals (epoxides) with cheaper carbon dioxide.⁵,⁶ Current academic and industrial research focuses on applying ROCoP to make low molar mass (Mₙ) poly(carbonate/ether carbonate) polyols, i.e., chains with Mₙ = 1–10 kg mol⁻¹ and hydroxyl chain-end groups.⁶,⁷ These CO₂-polylols deliver high-performance thermosets or polyurethanes that are used to make flexible foams for furniture, rigid foams for home insulation, scratch-resistant coatings, and thermoplastic polyurethanes.⁸–¹⁴ Life cycle analysis comparing CO₂-polylols with industrially used polyether polylols, made using 100% epoxide, shows a triple win in terms of greenhouse gas emissions: for every molecule of CO₂ polymerized, two more molecules are saved in avoiding epoxide usage.¹⁵ Other reports describe how to integrate CO₂/epoxide ROCoP with industrial CO₂ capture technologies.³,¹⁶ Carbon dioxide/epoxide ROCoP can also deliver high molar mass polycarbonates (Mₙ = 10–100 kg/mol⁻¹) and among the most widely synthesized is poly(cyclohexene carbonate) (PCHC).¹⁷,¹⁸ There are many reports of catalysts delivering PCHC, from CO₂/cyclohexene oxide (CHO) ROCoP; the highest performing are dinuclear metal complexes or bicomponent metal complex and cocatalyst
Our research group has developed macrocyclic dinuclear catalysts that operate at 1 bar carbon dioxide pressure, and the most active are heterodinuclear Zn(II)/Mg(II) complexes. Despite the many advances in catalyst development, there is a paucity of reports addressing the properties and applications for PCHC. Nearly 20 years ago, Koning et al. reported that PCHC has a high glass transition temperature (105−115 °C), high stress at break (11−42 MPa), and excellent tensile modulus (2400−3600 MPa), but is limited by a very low elongation at break (0.5−2%) and by its brittleness. Since this time, material development has been limited and any current application of PCHC necessitates plasticizers and additives to overcome its brittleness. One strategy to improve the properties of PCHC, and more generally CO2-polycarbonates, is to incorporate them into block polymers. Although there are reports of the syntheses of copolymers of PCHC with petroleum derived blocks, e.g., poly(acrylates), -ethers, -styrenes, -vinyl ethers, syntheses of copolymers of PCHC with petroleum derived blocks are rare. Although there are reports of the synthesis of copolymers of PCHC with petroleum derived blocks, e.g., poly(acrylates), -ethers, -styrenes, -vinyl ethers, and -siloxanes, none investigate the resulting materials properties. Very recently, Rieger and co-workers applied switchable catalysis to make block and statistical copolymers of poly(β-butyrolactone) and PCHC. The block copolymers show phase separation, and at 50 wt % PCHC content the materials’ tensile elongation at break values reach 5% (PCHC = 0.5−2%). By applying a better choice of soft block material, switchable catalysis could significantly improve upon PCHC properties, but the method is still hampered by several catalyst and process limitations. First, the switchable catalytic rates are still too low; for example, under monomer mixture conditions CO2/epoxide ROCOP TOFs are 1−50 h−1 and lactone ROP values are 1−100 h−1. Second, when producing higher molar mass block polymers, there are difficulties controlling the chain end groups, which leads to mixtures of structures (vide infra). Third, the overall block polymer molar masses are too low for block phase separation, which limits the resulting thermal-mechanical properties.

To address these barriers, a new catalyst selective for the production of high molar mass ABA block polymers, featuring poly(cyclohexene carbonate) (PCHC) as the A block and poly(ε-decalactone) (PDL) as the B block, is proposed. The block combination is designed so that the rigid structure of PCHC, together with its high glass transition temperature (Tg ∼ 110−120 °C), serves as a “hard” block and benefits the resulting material with a high tensile modulus. To accompany it, PDL is a useful “soft” block due to its amorphous structure, low Tg (∼60 °C) and relatively low entanglement molar mass. The polymers are also designed to increase the bio-based content, as ε-decalactone is sourced from plants and CO2 is a common byproduct of bioprocessing, as well as being a common industrial waste. CHO is currently sourced from petroleum, but routes from fatty acid coproducts could be implemented to increase bio-based content. Finally, by combining carbonate and ester blocks, obviating plasticizer usage, it is envisaged that (chemical) recycling will be facilitated, while the rigid and substituted main chain structures should provide the balance of sufficient material stability.

### RESULTS

To prepare the new ABA triblock polymers, it is essential to overcome catalyst activity limitations, to control chain end-groups, and to deliver high molar mass polymers (Mn > 50 kg mol−1). To apply a one-pot switchable procedure, a single catalyst that can be directed between the CHO/CO2 ROCOP and ε-DL ROP cycles is needed (Scheme 1). To increase the catalytic rates, a Zn(II)/Mg(II) catalyst, featuring para-di(methyl)amino-benzoate coligands, is selected as it shows some of the highest absolute rates in CHO/CO2 ROCOP, especially at 1 bar pressure (Figure 1). First it is necessary to evaluate the catalytic performance in lactone ROP. It is essential to solve a well-known problem affecting epoxide/CO2 ROCOP: most catalysts deliver a mixture of α,ω-dihydroxyl telechelic and α-hydroxyl-ω-carboxylate end-capped polymer chains, observed through bimodal molar mass distributions and by end-group analysis.

The mixed chain end-groups arise because the copolymerization is initiated both from the catalyst (e.g., from the benzoate coligand) and from diols (e.g., 1,2-cyclohexane diol, CHD); the diols either contaminate the epoxide or are rapidly formed upon reaction of the epoxide with even ppm quantities.
of water (see Scheme S1 for illustrations of the ROCOP initiation and chain transfer reactions). The problem is exacerbated when using catalyst + cocatalyst systems because additives like PPNCl may also initiate chains; fortunately this is obviated using the Zn(II)/Mg(II) catalyst because it functions without cocatalyst. Unsolved, the mixed end-group phenomenon would be problematic because the desired ABA polymers would be contaminated by AB type PCHC-PDL chains.

One method to suppress the catalyst initiated polymer chains is to carry out polymerizations in the presence of excess diol, e.g., 1,2-cyclohexane diol (CHD), or with additional water present. The diol is an efficient chain transfer agent and increases the quantity of $\omega$-$\omega$-hydroxyl telechelic chains (Scheme S1). It was already observed that monomodal molar mass distributions were observed in CHO/CO$_2$ ROCOP when the catalyst:CHD ratio exceeded 1:8.\textsuperscript{31} To assess the applicability of the strategy to lactone ROP, $\varepsilon$-DL polymerization was conducted in excess CHO as solvent, to mimic subsequent switch catalysis conditions, and using a ratio of Zn/Mg:CHD of 1:8. Under these conditions, poly($\varepsilon$-decalactone) (PDL) was efficiently prepared, but it showed a bimodal molar mass distribution, by SEC analysis, and mixed chain end-groups, by MALDI-TOF mass spectrometry (Figure 1, top). Polymerizations conducted using a greater quantity of CHD (Zn/Mg: CHD = 1:20) resulted in PDL showing a monomodal molar mass distribution (Figure 1, middle). The majority of chains were dihydroxyl telechelic but the overall molar mass was very much reduced ($M_n = 5.3$ kg mol$^{-1}$), consistent with the controlled polymerization mechanism. It is clear that the strategy of adding progressively greater quantities of diol is not at all suitable for the delivery of high molar mass polymers.

An alternative approach is to prepare a catalyst featuring a labile, organometallic ligand, such as an alkyl or aryl group, that reacts with diol in situ to form the metal-alkoxide initiator.\textsuperscript{32,49,57} One benefit could be that fewer equivalents of diol (vs catalyst) are needed to deliver effective catalysis and higher molar mass polymers are feasible. To test the hypothesis, a new Zn(II)/Mg(II) catalyst featuring two pentafluorophenyl coligands was prepared via a two-step...
The macrocyclic ligand was first reacted with 1 equiv of \([\text{Mg(N(SiMe}_3)_2]\) to form, in situ, the monomagnesium complex, which was then reacted with \([\text{Zn(C}_6\text{F}_5)_2]\) to produce the desired \([\text{LZnMg(C}_6\text{F}_5)_2]\) catalyst in good isolated yield (70%). The complex shows a sharp 1H NMR spectrum, at room temperature, with eight clearly separated signals for the benzylic and methylene protons (Figure S1). This NMR feature is highly diagnostic of pure heterodinuclear complex formation, as is the observation of two mutually coupled aromatic proton signals (Figure S1). The 13C{1H} and 19F{1H} NMR spectra are also fully consistent with the formation of a low-symmetry heterodinuclear complex, and the separate signals for each of the two phenyl and benzyl carbons are particularly diagnostic (Figures S2–S6).

The new catalyst was tested in CHO/CO2 ROCOP, at 1 bar CO2 pressure and in the presence of 4 equiv of CHD, and showed excellent activity and selectivity (TOF = 101 h⁻¹, 99% selective for carbonate linkages, Table S1, entry 1). It produced ABA-20

$\text{ABA-20}$  
$\text{80:20}$  
$\text{232:68}$  
$55.9$  
$[1.16]$  
$\text{48}$  
$n.d.$  
$286$

$\text{ABA-26}$  
$\text{74:26}$  
$\text{307:131}$  
$71.9$  
$[1.16]$  
$\text{50}$  
$69$  
$290$

$\text{ABA-28}$  
$\text{72:28}$  
$\text{200:93}$  
$46.4$  
$[1.07]$  
$\text{48}$  
$n.d.$  
$272$

$\text{ABA-40}$  
$\text{60:40}$  
$\text{168:135}$  
$49.4$  
$[1.08]$  
$\text{49}$  
$110$  
$278$

$\text{ABA-50}$  
$\text{50:50}$  
$\text{147:173}$  
$59.8$  
$[1.10]$  
$\text{50}$  
$110$  
$118$  
$283$

$\text{ABA-76}$  
$\text{24:76}$  
$\text{59:225}$  
$38.0$  
$[1.12]$  
$\text{44}$  
$n.d.$  
$254$

$\text{ABA refers to PCHC-}b\text{-PDL-}b\text{-PCHC. See Table S1 for experimental data on monomer loading and conversion.}$

$\text{ABA-#} : \# \text{ denotes weight fraction of PCHC (wt %).}$

$\text{Determined by } ^1\text{H NMR spectroscopy using the integrals for PCHC (4.66 ppm) and PDL (4.85 ppm) (Figure S8).}$

$\text{Determined on the basis of overall monomer conversion calculated from the } ^1\text{H NMR spectra (See Table S2).}$

$\text{Determined by SEC, in THF, using narrow dispersity polystyrene standards.}$

$\text{\(M_n\)/\(M_w\) (Figure S12).}$

$\text{Determined by DSC analysis from the second heating cycles.}$

$\text{\(T_g\) values reported as the midpoint of each transition (Figures S13–S18).}$

$\text{Acquired by DMTA. Upper \(T_g\) values reported as the maxima in tan(\(\delta\)) (Figures S19–S22); n.d. = not determined.}$

$\text{Determined by TGA. Reported as temperature at 5\% mass loss (Figures S23–S28).}$

**Table 1. PCHC-b-PDL-b-PCHC (ABA Triblock) Characterization Data**

| polymer $^{a,b}$ | wt % PDL:PCHC$^c$ | DP PDL:PCHC$^d$ | $M_n$ (kg mol⁻¹) $^{[D]}/$ $[D]$ | $T_g$SM$^b$ (°C) | $T_g$DMTA$^b$ (°C) | $T_{5\%}$ $^{i}$ (°C) |
|------------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| ABA-20           | 80:20           | 232:68          | 55.9 $/[1.16]$  | $\text{48}$     | n.d.            | 286             |
| ABA-26           | 74:26           | 307:131         | 71.9 $/[1.16]$  | $\text{50}$     | 69              | 290             |
| ABA-28           | 72:28           | 200:93          | 46.4 $/[1.07]$  | $\text{48}$     | n.d.            | 272             |
| ABA-40           | 60:40           | 168:135         | 49.4 $/[1.08]$  | $\text{49}$     | 110             | 278             |
| ABA-50           | 50:50           | 147:173         | 59.8 $/[1.10]$  | $\text{50}$     | 110             | 118             | 283             |
| ABA-76           | 24:76           | 59:225          | 38.0 $/[1.12]$  | $\text{44}$     | n.d.            | 254             |

$^{a}$ABA refers to PCHC-b-PDL-b-PCHC. See Table S1 for experimental data on monomer loading and conversion.

$^{b}$ABA-#: # denotes weight fraction of PCHC (wt %).

$^{c}$Determined by 1H NMR spectroscopy using the integrals for PCHC (4.66 ppm) and PDL (4.85 ppm) (Figure S8).

$^{d}$Determined on the basis of overall monomer conversion calculated from the 1H NMR spectra (See Table S2).

$^{e}$Determined by SEC, in THF, using narrow dispersity polystyrene standards.

$^{f}M_n/M_w$ (Figure S12).

$^{g}$Determined by DSC analysis from the second heating cycles. $T_g$ values reported as the midpoint of each transition (Figures S13–S18).

$^{h}$Acquired by DMTA. Upper $T_g$ values reported as the maxima in tan(δ) (Figures S19–S22); n.d. = not determined.

$^{i}$Acquired by TGA. Reported as temperature at 5% mass loss (Figures S23–S28).
only dihydroxyl telechelic PCHC with a monomodal molar mass distribution and an overall $M_n$ of 12.8 kg mol$^{-1}$ (Table S1, Figure S7). It was also tested for $\varepsilon$-DL ROP, using identical conditions to previous runs and with only 4 equiv of CHD. It showed good activity, with a TOF of over 4500 h$^{-1}$, and yielded high molar mass PDL with a monomodal, narrow dispersity distribution ($M_n = 33.1$ kg mol$^{-1}$, $Đ = 1.10$) (Table S1, Entry 2). The polymer chains were all $\alpha,\omega$-dihydroxyl telechelic as determined by MALDI-TOF mass spectrometry (Figure 1, bottom).

The new Zn(II)/Mg(II) organometallic catalyst was used to prepare a series of ABA triblock polymers using a one-pot procedure (Figure 1, Scheme 2). As a representative example, the preparation of ABA-50 is described. The reaction was conducted in two stages: first the catalyst was mixed with 600 equiv of DL and 700 of CHO resulting in efficient $\varepsilon$-DL ROP and the selective formation of dihydroxyl telechelic PDL. Near complete PDL formation (>90%) occurred within 20 min.

Next, CO$_2$ was added at a pressure of 20 bar, and the polymerization stirred for a further 20 h until it reached >90% PCHC conversion. A sample of the crude material showed the formation of a high molar mass polymer (59.8 kg mol$^{-1}$, $Đ = 1.10$) (Table 1, entry 5).

To better understand the monomer selectivity, a representative polymerization with the same wt % composition in the product (ABA-50), was performed in a Schlenk tube fitted with an in situ ATR-IR spectroscopic probe and using only 1 bar CO$_2$ pressure. Under these conditions there is less efficient reaction stirring, and so the overall reaction times are longer (rates are lower), but the catalytic selectivity is the same (Scheme 2).$^{21}$ The reaction shows an exponential growth of PDL, consistent with a first order dependence of rate on DL concentration. The polymerization kinetics are in line with previous investigations of lactone ROP.$^{59}$ Upon changing the gas atmosphere from nitrogen to carbon dioxide, the DL ROP stops and PCHC slowly begins to form (Scheme 2). The in situ analysis clearly shows the very high reaction selectivity: during the first reaction phase there is only formation of PDL, and once the CO$_2$ is added into the vessel, there is no further conversion of DL.

In terms of ABA-50 characterization data, the $^1$H NMR spectrum shows quantitative selectivity for carbonate linkages (>99%) and for polymer formation (>99%) in the PCHC blocks. There are no signals corresponding to ether linkages or to cyclic carbonate (Figure S8). Moreover, since each monomer is >90% converted, the composition of the block polymer closely matches the starting monomer stoichiometry. The polymer composition remains the same after purification to remove excess epoxide (Figures S8–S9). The $^{13}$C($^1$H) NMR spectrum shows two different types of carbonyl signal, assigned to -ester and -carbonate blocks, respectively, and shows there is a lack of transesterification (Figure S10). Chain end-group analysis, by $^{31}$P($^1$H) NMR spectroscopy,$^{42}$ shows only PCHC hydroxyl end-groups and no remaining PDL end-groups (Figure S11). Together the data provide strong support for the ABA triblock structures.

The same switchable process was applied using a range of monomer compositions and producing a systematic series of different block polymers (Table 1 and Table S2). In all cases, the reactions were very well controlled with polymer...
composition being predictable from monomer stoichiometry. In all cases, monomer conversions were >90% and the overall molar mass was sufficiently high (45–60 kg mol⁻¹, DP = 300) to drive phase separation. The series comprises DL-CHO monomer ratios from 1:3.3 to 3:1, and delivers ABA polymers to drive phase separation. The series comprises DL:CHO (0.1 mol % vs monomers) and proceeded effectively at low catalyst loadings (0.1–0.05 mol % vs monomers) and proceeded efficiently showing monomer conversions >90% within 20 h. A significant advantage of switchable catalysis is the removal of any chain-end scission process as the principal degradation. Overall, the chain-end scission process as the principal degradation.

The reactions were all effective at low catalyst loadings (0.1–0.05 mol % vs monomers) and proceeded efficiently showing monomer conversions >90% within 20 h. A significant advantage of switchable catalysis is the removal of any chain-end scission process as the principal degradation. The series comprises DL-CHO monomer ratios from 1:3.3 to 3:1, and delivers ABA polymers to drive phase separation. The series comprises DL:CHO (0.1 mol % vs monomers) and proceeded effectively at low catalyst loadings (0.1–0.05 mol % vs monomers) and proceeded efficiently showing monomer conversions >90% within 20 h. A significant advantage of switchable catalysis is the removal of any chain-end scission process as the principal degradation. Overall, the chain-end scission process as the principal degradation.

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The series of polymers were all characterized using a range of spectroscopies and SEC to confirm their compositions and molar masses (Table 1). SEC analyses show minor amounts of lower molar mass PCHC (Figure S12). This shows for Co addition when ppm quantities of water are also present as a contaminant in the gas stream. The residual water remains even after drying (using drying columns) and has already been shown to form diol (1,2-cyclohexane diol, CHD), which is a chain transfer agent. In terms of the block polymer properties, minor residual PCHC is not considered deleterious particularly as the new triblocks are designed to optimize PCHC properties. The series of samples are labeled ABA-#, where the number refers to the weight percentage of PCHC (hard), which is determined from the composition data.

The series of polymers, from ABA-20 to ABA-76, were analyzed by thermogravimetric analysis (TGA) to assess thermal stability relevant for any future processing and application (Figures S23–S28). Onset degradation temperatures (T_{d,95}) were generally around 280 °C, which is consistent with purified PCHC. The decomposition profiles differ according to the block polymer compositions and two step-profiles are observed (Figure 2c). For example, for ABA-50, the first degradation corresponds to 50% weight loss, consistent with the degradation of the PCHC. The second degradation, occurring at 322 °C, corresponds to 50% weight loss, consistent with PDL degradation. In all samples, the outer PCHC blocks are thermally degraded first, consistent with a chain-end scission process as the principal degradation. Overall, the T_{d,95} value is at least 170 °C higher than the upper glass transition temperature for the PCHC blocks (T_g ∼110 °C), and this temperature range facilitates thermoplastic processing. Differential scanning calorimetry (DSC) shows the two blocks are amorphous and only glass transitions are observed (Figures S13–S18). For example, ABA-50 shows two T_g values at similar temperatures to the homopolymers (T_g,PDL ≅ −58 °C, T_g,PCHC ≅ 110–120 °C) and the temperature range facilitates thermoplastic processing. Differential scanning calorimetry (DSC) shows the two blocks are amorphous and only glass transitions are observed (Figures S13–S18). For example, ABA-50 shows two T_g values at similar temperatures to the homopolymers (T_g,PDL ≅ −58 °C, T_g,PCHC ≅ 110–120 °C). Some of the other samples only show the lower T_g by DSC analysis, although in all cases its value is close to −60 °C. According to the Fox–Flory relationship, block miscibility would give rise to T_g,blend values from −23 to 7.5 °C, and thus, the blocks are likely to be phase separated. The apparent absence of the upper T_g in the DSC thermograms of low PCHC content samples is likely caused by its reduced concentration and the low DSC signal intensity (Figures S13–S18).

Some of the samples were further investigated by dynamic mechanical thermal analysis (DMTA) to identify their upper T_g and showed typical behaviors of viscoelastic materials (Figures 2b, S19–S22). The DMTA data for ABA-50 show that as the temperature increases from 30–100 °C, there is a gradual decrease in storage modulus E' (from 200 MPa) and an increase in loss modulus (E''). Correspondingly, the tan(δ), and glass transition temperature, is maximized at 118 °C. The T_g value for ABA-50 of 118 °C is consistent with the value obtained by DSC. DMTA also allows determination of the upper T_g (PCHC) for other samples and helps substantiate the phase separation (Figures S19–S22). Small-angle X-ray scattering (SAXS), applied to ABA-50, shows a strong principal scattering peak (with intensity q^4) corresponding to a domain spacing (d) of around 21 nm (d = 2π/q^2) (Figure 2d). The data are consistent with phase separation but suggest that there is only limited long-range order in the block polymer and, because higher order reflections are not observed, the exact block morphology could not be determined. Similar SAXS results were obtained for samples ABA-40 and ABA-26 (Figures S29–S30).

Mechanical Properties. ABA-50, -40, -28, -26 were successfully processed into films suitable for tensile mechanical experiments. Films of ~150 μm were prepared via solvent casting from a methylene chloride solution (30 wt %) and were dried by solvent evaporation under ambient conditions for 72 h, and then in vacuo at 40 °C for a further 48 h. Dumbbell shaped samples were prepared, according to ISO 527–2 type SB, and uniaxial extension experiments, conducted according to ISO 527, were used to derive stress vs strain relationships (10 mm min^−1 cross-head speed). The mechanical properties of the triblock polymers vary as a function of the relative PCHC (hard-block) weight fraction (Table 2). The tensile stress–strain curves show that samples ABA-50 and ABA-40 have behaviors typical of ductile plastics with yield points at ~9% and ~17% strain, respectively (Figure 3 red and blue curves). Beyond the yield points, both materials exhibit plastic deformation and undergo significant strain-hardening (Figure S31a)). ABA-50 shows the highest tensile strength (20 MPa) and a very high elongation at break (900%). It shows a tensile toughness (U_t), estimated from the stress–strain data, of 112 MJ m^−3.

ABA-28 exhibits elastomeric behavior and does not show any obvious yield point. Accordingly, the samples of ABA-28 quickly recover their original shapes once stress is released and show no permanent plastic deformation even beyond the point of fracture (Figures 3, orange curve, and S31(b)). ABA-28 shows high elasticity, with high values of elongation at break (>1000%) and a low ultimate tensile strength. To explore further, ABA-26 was synthesized with a similar composition to ABA-28 but with higher molar mass (M_n = 70 kg mol⁻¹).

### Table 2. Mechanical Properties of ABA Triblock Polymers as a Function of Block Composition and Molar Mass

| polymer | wt % PCHC | M_n (kg mol⁻¹) | E_'* (MPa) | σ_'* (MPa) | ε_'* (%) | ε_d (%) |
|---------|-----------|----------------|-------------|-------------|----------|---------|
| ABA-50  | 50        | 59.8 [1.10]    | 238 ± 35    | 20 ± 2      | 9.0 ± 1.7 | 900 ± 104 |
| ABA-40  | 40        | 49.4 [1.08]    | 81 ± 13     | 5.1 ± 0.2   | 17 ± 2    | 1920 ± 235 |
| ABA-28  | 28        | 46.4 [1.07]    | 3.0 ± 0.8   | 0.85 ± 0.04 | N.A.     | 1060 ± 54  |
| ABA-26  | 26        | 71.9 [1.16]    | 3.9 ± 0.7   | 1.4 ± 0.1   | N.A.     | 1400 ± 69  |

“Young’s modulus. bTensile strength. cStrain at yield point. dStrain at break. Mean values ± std. dev. from measurements conducted independently on at least 5 specimens.”
Increasing the overall polymer molar mass significantly increases the elongation at break but only slightly changes the ultimate tensile strength. Such high elasticity, low tensile strength materials are of future interest as mimics of elastin and other soft natural materials. The two elastomers, ABA-26 and ABA-28, were subjected to ten repeated stress–strain cycles, where samples were stretched to 200% elongation, the stress removed and the sample relaxed at 10 mm min⁻¹, and then the stress repeated (Figures S32–S33). Both materials showed a slight reduction in Young’s modulus and stress between the first and second cycles, which indicates slight plastic deformation in the first cycle. Thereafter both materials show high elastic recovery values, specifically these were 75% for ABA-26 and 95% for ABA-28.

The two plastics, ABA-40 and ABA-50, showed strain-induced plastic-to-rubber transitions (Figures S34–S35). In hysteresis experiments (0–200% strain, 10 cycles), a clear yield point was observed in the first cycles, yet subsequent cycles showed elastic profiles. It is hypothesized that the plastic-to-rubber transition is due to a change in the polymer microstructure so that there is isolation and alignment of PCHC domains within the PDL matrix. Similar behavior has previously been observed for poly(styrene-butadiene-styrene) (SBS) materials and for block polyesters.

The remaining samples containing high quantities of PCHC or PDL, respectively, and pure PDL could not be processed into freestanding films. For example, ABA-20 is highly viscoelastic and cannot retain its shape on removal from the mold (Figure S36(a)). In contrast, ABA-76 is very brittle, and shatters under minimal stress (Figure S36(b)). Attempts to analyze a 50:50 wt % mixture of PCHC and PDL showed macrophase separation with the brittle PCHC layer being suspended above the softer PDL layer. The polymer blends could not be processed or analyzed further and would clearly be very difficult to practically apply (Figure S36(c)). One future application for these block polymers could be as blend compatibilization.

Samples of ABA-20 were assessed as pressure sensitive adhesives. Oscillatory rheology was used to probe its viscoelastic properties. A frequency sweep reveals the regions over which the viscous (⁢) and elastic (⁢) components dominate (Figure 4a). The shear storage modulus, ⁢, and shear loss modulus, ⁢, at frequencies consistent with adhesive bonding (10⁻¹ rad s⁻¹) and debonding (10² rad s⁻¹), were used to construct a viscoelastic window. The data suggest ABA-20 could be used as a high shear, permanent adhesive (Figure 4b). The bonding aspect of the window (lower boundary) lies below the Dahlquist criterion (⁢ ≤ 3 × 10⁵ Pa), indicating ABA-20 should be a contact efficient or pressure-sensitive adhesive (PSA) without needing further additives. A fixed oscillation vs time experiment showed a stable storage modulus (⁢), at 2.6 × 10⁵ Pa, and at a value which is consistently higher than the loss modulus (⁢); i.e., ABA-20 is a soft, viscoelastic solid (Figure S37). The adhesive properties of ABA-20 were assessed using 180° peel, and loop-tack tests, using pressure-sensitive tape council (PSTC) grade polished stainless steel as the substrate and 5 μm thick PETE film (Mylar) as the adhesive backing. The samples were coated onto the PETE sheet, from a 30 wt % solution in DCM using a wound wire rod, and the film thickness was ~60 μm. ABA-20 displays a maximum peel force ~10 N cm⁻¹ (Figure S38). The loop tack test shows an average maximum tack force ~8.7 N cm⁻¹ (Figure S39). A cross-hatch test, with ABA-20 coated onto PSTC-grade stainless steel, showed only minimal coating removal on application and removal of scotch tape (ISO classification: 1). These data are indicative of strong adhesive bonding to steel (Figure S40).
The ABA triblock polymers, PCHC-b-PDL-b-PCHC, are prepared with straightforward and efficient catalysis. Lactone ring-opening polymerization delivers the central PDL soft-block, and by simply adding carbon dioxide into the reaction vessel, two outer PCHC hard-blocks are attached. The new catalyst increases reaction rates, maximizes polymer molar masses, and controls its structure. It is applied at low loadings, from 1:1000–1:2000 (catalyst: CHO + DL) and in toluene solutions (3 M); it is highly active with turnover frequency (TOF) values for DL ROP exceeding 4000 h$^{-1}$ and for CHO/CO$_2$ ROCOP of 100 h$^{-1}$. The high performance allows reactions to be conducted using minimized catalyst residues (0.1–0.05 mol %), at acceptable reactor residence times (<24 h) and with monomer conversions >90% obviating monomer removal strategies at the end of reaction. The catalytic activity is up to 20 times higher than previously reported CO$_2$/CHO ROCOP (TOF = 5 h$^{-1}$ under comparable conditions) and 90 times higher than previously reported lactone ROP (TOF = 50 h$^{-1}$ for ε-caprolactone ROP under comparable conditions).$^{7,50}$ The catalysis is also well-controlled and delivers predictable block compositions and structures. By applying the new process, ABA block polymers with variable PCHC weight ratios and carbon dioxide uptakes are easily produced.

By controlling the CO$_2$ utilization and position in the polymer chain, the properties vary from viscoelastic pressure sensitive adhesives (ABA-20), to high elongation thermoplastic elastomers (ABA-26 and ABA-28) to, at highest CO$_2$ utilization, toughened plastics (ABA-40 and ABA-50). There is clearly a much broader scope of applications for PCHC than was previously envisaged. Considering specifically adhesives applications ABA-20, with a ~6 wt % CO$_2$ content, shows good performance as a single-component pressure-sensitive adhesive. It has a low dynamic elastic modulus, below the Dahlquist criterion for adhesives, and, in contact with steel surfaces, shows both high peel (10 N cm$^{-1}$) and tack strengths (8.7 N cm$^{-1}$). It has higher peel strength than commercial adhesives such as duct, masking, electrical, and Scotch tape, and these commercial adhesives are formulated products, comprising mixtures of polymers, plasticizers, fillers, and tackifiers. In contrast, the data for ABA-20 are stand-alone, but future investigations should also apply these CO$_2$-derived polymers in formulations.$^{71}$

The promising characteristics of ABA-20 suggest it could be used instead of styrene, polyacrylate, or natural rubber, currently applied as the polymeric component in adhesives. ABA-20 also shows equivalent, or better, performance than leading triblock polymer pressure-sensitive adhesives, applied with tackifiers, which were reported in the literature. $^{72–76}$ It is compared to bio-based block polyester and polyacrylate based adhesives, which were applied with various tackifiers, some biobased (see Figure S41 for illustrations of their structures).$^{72–79}$ In particular, it is bench-marked against ABA block polymers, comprising polyolactide A-blocks and soft B-blocks derived from menthoxide or ε-decalactone (Figures S, S41)$^{72,74,75,77–79}$ ABA-20 shows higher peel force strength than the corresponding bioderived ABA polyester or acrylate polymer/tackifier blends and illustrates the benefits of PCHC rigid blocks. $^{74,75,78}$ Further, higher operating temperatures are anticipated since PCHC ($T_g = 110–120$ °C) has a considerably higher glass transition temperature than PLA ($T_g = 50–60$ °C). Compared to the highest performing polyacrylate pressure-sensitive adhesives, particularly those mimicking muscle adhesion, the peel force and tack results are modest,$^{80,81}$ demonstrating the significant scope to optimize structures in the future.$^{82}$

Increasing the CO$_2$ content to ~15 wt % in sample ABA-50 yields a toughened plastic. ABA-50 shows significantly higher toughness, through the incorporation of the soft PDL midblock, than PCHC alone and overcomes the well-known brittleness of PCHC (Figure 6). Compared with bioderived poly(limonene carbonate) (PLLmC), prepared by ROCOP of limonene oxide/CO$_2$, ABA-50 shows significantly higher tensile toughness ($U_T$): ~3.15 MJ m$^{-3}$ for PLLmC, $^{83}$ compared to 112 MJ m$^{-3}$ for ABA-50. One future application signaled by these results is as a compatibilizer and impact modifier for well-known PCHC or PLLmC. A very recent report of PBL-b-PCHC showed at 50 wt % PCHC incorporation a similar tensile strength of 17 MPa but with just 1.3% elongation at break.$^{81}$ In terms of benchmarking the performance of ABA-50 against other plastics, it shows higher toughness, but considerably lower tensile strength, than the well-known commercial polycarbonate, prepared from bisphenol A ($U_T$ ~ 19 MJ m$^{-3}$) (Figure S42). It also significantly improves upon the toughness of popular biobased plastic polylactide (PLA), with toughened PLA showing $U_T$ ~ 18 MJ m$^{-3}$, and maintains comparable ultimate tensile strength to toughened PLA (25.8 MPa).$^{84,85}$

Given the encouraging properties of ABA-50 and the straightforward method to prepare it, future research could focus on how to exploit its high toughness and increase tensile strength. For example, using isotactic PCHC blocks to enhance rigidity or incorporation of vinyl-cyclohexene oxide as a cross-linkable group in the carbonate block could both offer means to increase tensile strength.$^{36,46}$ An additional advantage of using commercially available vinyl-substituted epoxides is the facility to “pattern” the resulting materials and install functionalities that control hydrophilicity, filler binding, and/or pH responsiveness.$^{87}$ Another feature of these triblock
polymers is that they contain a high proportion of bioderived raw material and are easily prepared from commercially available monomers. Considering that ε-decalactone is biosourced from castor oil via ricinoleic acid, and that the CO₂ content is straightforward to control, the polymers off theoretical renewable contents from 47% (ABA-76) to 86% (ABA-20). Currently, cyclohexene oxide is produced from petrochemicals, but routes from triglyceride coproducts could be adopted in the future if fully biobased content were required.

In terms of end-of-life scenarios, the ester and carbonate linkages comprising the polymer backbone are expected to ultimately render the materials degradable. Moreover, degradation products, in the form of diols, hydroxy-acids, and CO₂, are not anticipated to be toxic. Preliminary stability studies showed that films of ABA-50 are stable to mass loss in acidic and basic media at ambient temperature for at least a period of 9 months. Nonetheless, the samples could be fully degraded by dissolving them in toluene, warming to 60 °C (to accelerate the rate of degradation), and treatment with a weak acid ([p-toluene sulfonic acid] = 5 mM). Under these conditions rapid loss of polymer molar mass occurred over a period of 96 h (Figure S43).

**CONCLUSIONS**

A switchable polymer synthesis protocol, using a new heterodinuclear Zn(II)/Mg(II) organometallic catalyst, allows for the fast, efficient, and controllable preparation of fully degradable ABA block polymers. The protocol is applied to prepare a logical series of ABA type poly(cyclohexene carbonate)-b-poly(ε-decalactone)-b-poly(cyclohexene carbonate), with 6–31 wt % of the material derived from CO₂. The polymers show promising thermal and mechanical properties, and progressively greater CO₂ utilization allows control over applications spanning viscoelastic adhesives, thermoplastic elastomers, and tough plastics. These block polymers overcome the well-known brittleness of PCHC, one of the most widely studied CO₂-derived polymers. Further, a range of new properties for CO₂-containing polymers are demonstrated, which expands their application scope well beyond polyols for polyurethane. Viewed from the perspective of aliphatic polyesters, the cyclohexylene carbonate units confer reinforcement, mechanical strength, and increased service temperature to the resulting copolymers and provide a viable and, in some cases, better performing alternative “hard” block to PLA. The range of future polymers and applications enabled by the polymerization process could indeed be very large, since it is expected to be generally applicable to a wide range of lactones, lactide, epoxides, anhydrides, and heterocumulenes.

**ASSOCIATED CONTENT**

- Supporting Information
  The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.9b13106.

  Detailed experimental procedures for catalyst synthesis and polymerizations, and catalyst and polymer characterization data (1D and 2D NMR, SEC, DSC, DMTA, TGA, SAXS) (PDF)

**AUTHOR INFORMATION**

**Corresponding Author**
Charlotte K. Williams — Department of Chemistry, Chemistry Research Laboratory, University of Oxford, Oxford OX1 3TA, U.K.; orcid.org/0000-0002-0734-1575; Email: charlotte.williams@chem.ox.ac.uk

**Authors**
Gregory S. Sulley — Department of Chemistry, Chemistry Research Laboratory, University of Oxford, Oxford OX1 3TA, U.K.
Georgina L. Gregory — Department of Chemistry, Chemistry Research Laboratory, University of Oxford, Oxford OX1 3TA, U.K.
Thomas T. D. Chen — Department of Chemistry, Chemistry Research Laboratory, University of Oxford, Oxford OX1 3TA, U.K.
Leticia Peña Carrodeguas — Department of Chemistry, Chemistry Research Laboratory, University of Oxford, Oxford OX1 3TA, U.K.
Gemma Trott — Department of Chemistry, Chemistry Research Laboratory, University of Oxford, Oxford OX1 3TA, U.K.

**Figure 6.** Ashby plots of tensile toughness (Uₜ) vs (a) tensile strength (σ) and (b) elongation at break (εₜ) for comparison of ABA-50 with selected commercial and literature materials. Materials classified as polycarbonate (▲), polyester (●), or poly(carbonate-b-ester-b-carbonate) (■).
Alba Santmartí — Department of Aeronautics, Imperial College London, London SW7 2AZ, U.K.
Koon-Yang Lee — Department of Aeronautics, Imperial College London, London SW7 2AZ, U.K.; orcid.org/0000-0003-0777-2292
Nicholas J. Terrill — Beamline I22, Diamond Light Source, Harwell Science and Innovation Campus, Didcot OX11 0DE, U.K.

Complete contact information is available at: https://pubs.acs.org/10.1021/jacs.9b13106

Notes
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