Chemically Recyclable, High Molar Mass Polyoxazolidinones via Ring Opening Metathesis Polymerization

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ABSTRACT: The development of robust methods for the synthesis of chemically recyclable polymers with tunable properties is necessary for the design of next-generation materials. Polyoxazolidinones (POxa) – polymers with five-membered urethanes in their backbones – are an attractive target because they are strongly polar and have high thermal stability, but existing step-growth syntheses limit molar masses and chemical recyclability to monomer is rare. Herein, we report the synthesis of high molar mass POxa via ring opening metathesis polymerization of oxazolidinone-fused cyclooctenes. These novel polymers show <5% mass loss up to 382–411 °C and have tunable glass transition temperatures (14–56 °C) controlled by side chain structure. We demonstrate facile chemical recycling to monomer and re-polymerization despite moderately high monomer ring strain energies, which we hypothesize is facilitated by the conformational restriction introduced by the fused oxazolidinone ring. This method represents the first chain growth synthesis of POxa and provides a versatile platform for the study and application of this emerging subclass of polyurethanes.

Synthetic polymers have revolutionized every aspect of modern life, but current mass polymer production is unsustainable as common materials generate tremendous amounts of waste¹⁻³ and pollution⁴⁻⁶ after their intended use. Therefore, next-generation materials require both competitive properties and designed end-of-life strategies to mitigate their environmental impact. Polyoxazolidinones (POxa) are an emerging class of polyurethanes with competitive properties for high temperature applications^{7,8} due to the five-membered oxazolidinone rings imparting high thermal stability^{7,9–13} and glass transition tempera-tures (T_g) .^{7,8,11–15} Despite their promise, POxa have been limited to low molar masses ($M_n \leq 32 \text{ kDa}^{8,13,15,16}$) for decades due to the step-growth nature of previous syntheses (Figure 1A) and very little work has been reported on sustainable approaches for their end-of-life.¹⁷ We sought to break this paradigm by using chain growth polymerization to access high molar mass POxa while enabling chemical recycling to monomer via depolymerization.18-21

We chose ring opening metathesis polymerization (ROMP) because (1) it is a powerful chain growth method for strained cycloalkenes^{22–24} and (2) ROMP-based polymers obtained from low ring strain monomers (~5.2 kcal/mol) are known to depolymerize at elevated temperatures via ring closing metathesis (RCM)^{25–29} (Figure 1B). This RCM strategy has recently resurfaced in the design of new polymers due to its promising contribution to chemical recycling.^{27,28,30–32} Chemical recycling to monomer is ideal for a circular polymer economy because this approach can minimize the net mass loss during recycling²¹ compared to other strategies, such as mechanical recycling,^{2.33–} ³⁵ biodegradation,^{36,37} and upcycling of commodity plastics to value added products.^{38–47} Recently, Wang and co-workers showed that the addition of *trans*-fused cyclobutene or acetonide rings to cyclooctene lowered the monomer ring strain energy (RSE), which they hypothesized was the critical factor that enabled the depolymerization of the resulting polymers.^{28,30}



Figure 1. (A) Traditional POxa from step growth polymerization (SGP) (A, B = reactive functionality). (B) Reversibility of ringopening metathesis for monomers with different ring-strain energies (RSEs). (C) Design of chemically recyclable POxa.

Herein, we utilize ROMP for the first chain growth synthesis of POxa with the oxazolidinones (Oxa) in the backbone of the polymer using *trans*-Oxa-fused cyclooctene monomers (Figure 1C). We demonstrate access to POxa with high molar mass, moderate dispersity, and control over molar mass by using a

chain transfer agent (CTA). These novel POxa show <5% mass loss up to 382 °C and have tunable T_g via side chain alteration and post-polymerization hydrogenation. Furthermore, we show clean depolymerization back to monomer, despite higher RSE than *cis*-cyclooctene (COE) calculated by density functional theory (DFT). This method provides access to POxa with higher molar masses than previous methods, which will enable future structure-property relationship studies and the addition of a broad-spectrum of new Oxa monomers for ROMP and chemical recycling to monomer.

We began our investigation by synthesizing the Oxa-fused COE monomers **M1–M4** in 4–5 steps from 1,5-cyclooctadiene (Scheme 1, see Supporting Information Section S3A for full synthetic details). Gratifyingly, **M1** readily polymerized using G2 (see Figure S1 for all catalyst structures) in dichloromethane (DCM) to yield POxa **P1** with a number-average molar mass (M_n) of 23.9 kDa; however, **P1** precipitated out during the polymerization due to poor solubility in DCM and was only soluble in *N*,*N*-dimethylformamide (DMF) and dimethyl sulfoxide (DMSO), which are incompatible with metathesis catalysts.^{48–50} Therefore, we focused on the *N*-alkylated monomers **M2–M4** which yielded polymers (**P2–P4**) that were soluble in common

metathesis solvents (e.g., DCM, CHCl₃, THF).

Scheme 1. Oxa-fused COE monomers from cyclooctadiene.



Using M2 as a model substrate, we optimized the polymerization conditions (Table 1). Except for G1 (entry 1), common metathesis catalysts (i.e., G2, G3, HG2, see Figure S1) all resulted in high conversions and molar masses ($M_n = 118-487$ kDa, entries 2-4). We selected G2 as the standard catalyst because it resulted in the lowest D and is relatively inexpensive. Increasing the initial monomer concentration from 1.0 to 2.0 M decreased conversion in 20 min and resulted in a bimodal distribution with high D (entry 5), whereas lowering the concentration to 0.5 and 0.25 M progressively decreased both conversion and molar mass (entries 6 and 7). Therefore, our standard polymerization conditions consisted of 1.0 M monomer in DCM using G2 as the ROMP catalyst at 24 °C. M3 and M4 were similarly polymerized to access high molar mass POxa $(M_n = 112.3 \text{ and } 88.1 \text{ kDa for } \mathbf{P3} \text{ and } \mathbf{P4}, \text{ respectively}) \text{ in } 1 \text{ h}$ with variable side chains (see Section S3B for synthetic details). Because these Oxa-fused COE monomers are unsymmetrical, we characterized the regioregularity of enchainment for M2-M4 using IR and NMR spectroscopies. While the side chain does affect regioregularity to some extent, all polymers had over 88% head-to-tail enchainment (see Section S2E for more details).

As expected for ROMP of COE derivatives,^{51–56} we observed evidence of slow initiation: M_n is initially very high and decreases (i.e., approaches $M_{n,theo}$) by extending the reaction time (entry 2 vs. 8–9 and Figure S5) and there was a non-linear trend between M_n and [M]:[Ru] ratio (Figure 2A). To improve control over M_n , we added *cis*-1,4-diacetoxy-2-butene as a chain transfer agent (CTA) (Figure 2B). CTAs are known to decrease the difference between $M_{n,theo}$ and M_n ,^{57,58} enable the use of catalytic Ru,^{55,59-64} and – for this CTA – result in homotelechelic polymers that can be deprotected to the corresponding diol for further functionalization.^{57,58} Despite requiring longer reaction times due to the slower rate of polymerization (78% conv. in 3 h vs. 90% without CTA), M_n had a linear relationship with [M]:([CTA]+[G2]) and was within 1–37 kDa of $M_{n,theo}$ for a wide range of molar masses, demonstrating improved control over POxa molar mass. Secondary cross-metathesis helps dispersity remain relatively constant ($D \approx 1.5$ –1.6) with and without the CTA (when t = 3 h).

Table 1 Optimization of ROMP conditions.

	M2, 100 eq Ph $[Ru] (1 eq)$ DCM $24 °C, 20 min$ $P2$ Ph Ph Ph Ph Ph Ph Ph Ph				
entry	[Ru]	conc. (M)	conv. $(\%)^a$	$M_n (kDa)^b$	D^b
1	G1	1.0	9	-	_
2	G2	1.0	82	487	1.20
3	G3	1.0	85	118	1.41
4	HG2	1.0	81	384	1.27
5^c	G2	2.0	62	203	2.35
6	G2	0.5	69	168	1.40
7	G2	0.25	48	<i>d</i>	<i>d</i>
8 ^e	G2	1.0	86	42.7	1.20
9 ^f	G2	1.0	86	34.4	1.55

^{*a*}Determined by ¹H NMR analysis of the crude reaction mixture. ^{*b*}Crude M_n and \tilde{D} were determined by size exclusion chromatography with multi-angle light scattering (SEC-MALS) in DMF. ^{*c*}[**M2**]:[Ru] = 50:1. ^{*d*}SEC peak is overlapping with small molecule peaks, but M_n of oligomers is < 1 kDa. ^{*e*}Polymerization was run for 1 h. ^{*f*}Polymerization was run for 3 h. See Figure S1 for catalyst structures.



Figure 2. Plots of M_n (black circles), $M_{n,\text{theo}}$ (open circles), and D (red triangles) as a function of (A) [**M2**]:[G2] and (B) [**M2**]:[CTA]+[G2]. Polymerization time = 3 h.

Having established the synthesis of POxa via ROMP, we next investigated the thermal properties of **P2–P4**. Thermogravimetric analysis (TGA) showed **P2–P4** had high decomposition temperatures at 5% mass loss ($T_{d,5\%}$) of 382–411 °C (Figure 3A), as expected for POxa.^{7,9–13} Differential scanning calorimetry (DSC) indicated the POxa were amorphous with glass transition temperatures (T_g) tuned by the *N*-substituent (Figure 3B). Increasing the substituent length from methyl (**P3**) to *n*-butyl (**P4**) lowered the T_g from 29 to 14 °C whereas the benzyl substituent in **P2** increased the T_g to 56 °C. These results indicate that our

modular platform for monomer synthesis enables the synthesis of POxa with tunable T_{g} 's while maintaining high $T_{d,5\%}$.



Figure 3. Thermal data – (A) TGA and (B) DSC thermograms – of **P2–P4** and (C) hydrogenated **[H]P2**.

To explore the range of properties accessible via this method, we hydrogenated the backbone olefins of P2 to form polyethylene-like polymer [H]P2 that still contains embedded polar Oxa rings (Figure 3C). Such polar-functionalized polyethylene⁶⁵ often exhibit excellent ionic conductivity,^{66–68} surface hydrophilicity,69,70 and antimicrobial properties71,72 while maintaining the mechanical strength of polyethylene. We speculated common olefin hydrogenation catalysts (e.g. Pd/C) would deprotect the benzyl group from the Oxa nitrogen⁷³ and thus used in situ-generated diimide from N-tosyl hydrazide for hydrogenation,^{56,74} resulting in 92% conversion of the olefins via ¹H NMR spectroscopy after 16 h. The $T_{d,5\%}$ increased slightly after hydrogenation (436 °C for [H]P2 vs. 408 °C for P2) whereas the $T_{\rm g}$ decreased from 56 to 42 °C due to increased backbone flexibility,75-78 indicating that post-polymerization modification provides an additional handle to tune thermal properties.

Because we are interested in the end-of-life for these materials, we explored the depolymerization of the non-hydrogenated **P2** via RCM. After a brief optimization of the depolymerization conditions (Table S4) we observed quantitative depolymerization of the polymer to **M2** using G2 in chloroform-*d* (20 mM) at 70 °C for 4 h (Figure 4). No oligomers or side reactions were observed. The recovered monomer (90% by mass) behaved identically to fresh monomer when re-polymerized using our standard conditions (see Figure S32), demonstrating the facile circular recyclability of these materials. Detrembleur and coworkers recently reported mechanical and chemical recycling of POxa-containing covalent adaptable networks,¹⁷ but this RCM approach is the first example of chemical recycling for linear POxa.



7.4 7.0 6.6 6.2 5.8 5.4 5.0 4.6 4.2 3.8 3.4 3.0 2.6 2.2 1.8 1.4 1.0 f1 (ppm)

Figure 4. ¹H NMR spectra (CDCl₃) of (A) **M2**, (B) **P2**, and (C) the crude reaction mixture of **P2** depolymerization. Depolymerization conditions: G2 (1 mol %), CDCl₃ (20 mM), 70 °C, 4 h.

Because low monomer RSE (≤ 5.3 kcal/mol) has been reported as the key factor in the depolymerization of ROMP-based polymers, 28,30 we expected M1–M4 to have lower RSE than COE, which does not readily depolymerize.^{24,79} However, the calculated RSE values were actually higher for M1-M4 (7.2-8.2 kcal/mol) than COE (6.6 kcal/mol) (Table S5), indicating that RSE is not the most important factor governing the depolymerization of these polymers. (Note that due to inconsistencies in the reported RSE of COE in the literature, ^{28,51,80,81} we calculated this value using the same method as M1-M4 at the B3LYP/6-31G (d,p) level of theory.) On the basis of the rich small-molecule literature using RCM to construct medium-sized (5-9 membered) fused rings (RSE > 7 kcal/mol), $^{82-88}$ we hypothesize the favorable depolymerization of P2 is facilitated by the conformational restriction introduced by the fused Oxa ring helping to bring the adjacent alkenes together (conceptually similar to the Thorpe-Ingold effect⁸⁹ that has previously been invoked for polymer depolymerization^{30,90}).

In summary, we successfully developed the first chain growth synthesis of POxa. We designed and synthesized novel Oxafused COE monomers with easily diversified side chains. P1 which contains free N-H groups - was insoluble in common ROMP solvents, but P2-P4 were accessed with high molar mass and moderate dispersity (M_n up to 487 kDa, $D \approx 1.5$) via ROMP. We demonstrated that the molar mass could be efficiently controlled using a commercially available CTA. This method delivers new POxa with $T_{d,5\%} > 382$ °C and tunable T_g by changing the nitrogen substituent. We showed hydrogenation of the olefin backbone of the polymer to give access to polvethylene-like POxa. We furthermore established the quantitative chemical recyclability to monomer of P2 by employing RCM to deliver pure monomer that could be readily re-polymerized. Further studies on controlled copolymerization, mechanistic insight, and mechanical properties are underway.

ASSOCIATED CONTENT

Additional polymer characterization, including SEC chromatograms, TGA and DSC thermograms, tabulated data, experimental details, calculation details, methods, reagent sources, synthetic procedures. This material is available free of charge via the Internet at http://pubs.acs.org.

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Author Contributions

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

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Notes

The authors declare no competing financial interest.

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