Ring-Opening Copolymerizations of a CO2-derived δ -Valerolactone with ϵ -Caprolactone and L-Lactide

Ryan J. Anderson, Rachel L. Fine, Rachel M. Rapagnani, Ian A. Tonks*

Department of Chemistry, University of Minnesota – Twin Cities, 207 Pleasant St SE, Minneapolis MN 55455

TOC Graphic:



Abstract

Ring-opening random, gradient, and block copolymerizations of the CO₂-derived δ -valerolactone 3-ethyl-6-vinyltetrahydro-2H-pyran-2-one (EtVP) with ε -caprolactone (ε -CL) and L-lactide (LLA) are reported. By employing both concurrent and sequential addition strategies, a variety of thermal and physical properties could be accessed. Concurrent copolymerization of EtVP with ε -CL yielded gradient copolymers with low glass transition temperatures, while block copolymerizations via sequential addition led to semi-crystalline materials regardless of monomer feed ratios. For LLA copolymerizations, glass transition temperatures increased with LLA incorporation regardless of addition method, but higher T_g values were observed in block copolymerizations from sequential addition. Tensile testing of a *poly*(EtVP-*b*-LLA) with molar ratio of 40:60 EtVP:LLA resulted in $\sigma = 0.8$ MPa, E = 5.6 MPa and 83% elongation at break. The chemical recyclability of EtVP-based copolymers was explored as an end-of-life option. Both ε -CL and LLA copolymers could be recycled, with block copolymers giving higher yields of recycled monomers than random copolymers.

Introduction

The disubstituted lactone, 3-ethylidene-6-vinyl-tetrahydro-2H-pyran-2-one (**EVP**) offers substantial promise as an inexpensive entry point into CO_2 -derived materials because it can be selectively synthesized through efficient Pd-catalyzed telomerization of butadiene and CO_2 .^{1,2} Biodegradable and chemically recyclable **EVP**-derived polyesters can be accessed through the ring-opening polymerization (ROP) of either the semi- or fully-hydrogenated δ -valerolactone **EVP** derivatives, 3-ethyl-6-vinyltetrahydro-2H-pyran-2-one (**EtVP**) or 3,6-diethyltetrahydro-2H-pyran-2-one (**DEP**).^{3,4} However, *poly*(**EtVP**) and *poly*(**DEP**) are both amorphous polymers: although they may find use in flexible/ductile materials⁵ or as adhesives,⁴ there is significant interest in tuning the materials properties of these polymers in an effort to expand their applications.

Ring-opening copolymerization (ROCOP) of lactones has been well-demonstrated as a method for accessing polyesters with a wide array of thermal and mechanical properties.⁶ ε -caprolactone (ε -CL) and L-lactide (LLA) are two of the most used and well-studied lactones for copolymer synthesis. $Poly(\varepsilon$ caprolactone) (PCL), which can be bioderived^{7,8}, has many biomedical applications due to its strong permeability, elasticity, and thermal properties.⁹ More specifically, PCL is semi-crystalline with a melting point (T_m) of 59-64°C and a glass transition temperature (T_g) of -60 °C, which results in high toughness at human body temperatures.^{10,11} PCL is also biodegradable, though it typically requires 2-3 years for complete degradation to occur in biological media.¹² Bulk PCL also exhibits low tensile strength (16-27 MPa) and elastic modulus (250-430 MPa).^{13,14} On the other hand, *poly*(L-lactic acid) (PLLA) is an attractive bio-based¹⁵ polymer with high tensile strength (28-50 MPa) and elastic modulus (1200-3250 MPa), though it suffers from poor toughness (2.0-6.0% elongation at break).^{13,16–19} Due to these factors, copolymers of PCL and PLLA have been developed to address the weaknesses of the individual homopolymers.^{19–22} While copolymer properties can be tuned through monomer feed ratios, molecular weights, and lactone choice, different copolymer microstructures (random, gradient, block copolymers) can also affect the overall thermal and mechanical properties. For example, PLLA-PCL random copolymers lack industrially relevant mechanical properties, as the randomization¹⁹ of the structure results in a decrease in overall polymer crystallinity compared to the their block counterparts.²³ Comonomer reactivity ratios can be used to predict the type of copolymer (alternating, blocky, gradient or random) that will form under ROP conditions.^{24,25}

Given the successes in the synthesis of **PCL/PLLA** copolymers *via* ROCOP,²⁶ we hypothesized that ε-CL and LLA would be ideal candidates for copolymerization with EtVP. We envisioned a copolymer comprised of EtVP and ε-CL could be easily processable and potentially chemically recylable.²⁷ Alternately, in a LLA/EtVP copolymer the softness of *poly*(EtVP) could balance out the brittleness of PLLA, potentially yielding good mechanical strength and high toughness for biomedical applications.²⁸ In both cases, EtVP incorporation would promote enhanced biodegradation potential, recyclability, and provide the opportunity for post-polymerization modification through functionalization of the side chain vinyl groups.

No **EtVP**-based copolymers have been reported to date. However, there have been a few recent copolymer syntheses with **EVP** (Figure 1), which were developed because **EVP** is reticent to homopolymerize.^{29–32} The Ni group has reported the use of β -butyrolactone (Figure 1, reaction A),²⁹ **ε-CL** (Figure 1, reaction B),³⁰ and cyclohexene oxide (Figure 1, reaction C)³¹ as comonomers for the ROP of **EVP** to synthesize random copolymers; similarly, the Tang group has used propylene oxide (Figure 1, reaction D).³² In all cases, low molar mass polymers with high dispersities were obtained, especially at high feed ratios of **EVP**. These limitations could be due to the α , β -unsaturated ester moiety of **EVP**, which generally impedes productive ring-opening and gives rise to deleterious Michael self-addition.³³ **EtVP** lacks this α , β -unsaturation and thus undergoes well-controlled ROP^{3,5}, so it is a good candidate for accessing more controlled copolymer microstructures *via* ROCOP. Herein, we report the copolymerization of **EtVP** with **ε**-**CL** and **LLA**. By altering when monomer addition occurs, copolymer microstructure can be controlled, leading to a broader range of thermal and physical properties (Figure 1, right).



Figure 1. Left: previous reports of copolymerizations with **EVP**. Right: controlled copolymerizations of ϵ -**CL** and **LLA** with **EtVP** can lead to various types of copolymer microstructure (this work).

Results and Discussion

Initial copolymerization attempts with ε -CL were conducted with 1,5,7-triazabicyclo[4.4.0]dec-5ene (TBD)³⁴ and 3-phenyl-1-propanol (3-PPA) as an initiator (Table 1) under neat conditions, building on previous studies of TBD-catalyzed ring-opening polymerization of EtVP.³ Across all feed ratios, high molar mass polymers (up to 54.0 kg/mol) could be achieved by employing 2.5 mol% TBD and 0.5% 3-PPA. At higher feed ratios of EtVP (Table 1, entries 4 and 5), lower molar masses are observed, which may be a result of increased undesirable TBD initiation.³ Copolymer composition demonstrated good agreement with monomer feed ratios, indicating good control of the polymerization. T_g increased with the molar fraction of EtVP, from -60.5 at 19% EtVP in the polymer (Table 1, entry 2) to -33.1 °C at 100% EtVP (Table 1, entry 5). Melting points were observed when a high ratio of ε -CL (74%) was incorporated (Table 1, entry 2). Higher feed ratios of ε -CL also impacted the dispersity (D), increasing from 1.2 in pure *poly*(EtVP) (entry 5) to 1.7 in pure PCL (entry 1). This increase is most likely due to the propensity for PCL chain ends to undergo transesterification,³⁵ especially in the presence of TBD.³⁶ Transesterification can be minimized by employing shorter reaction times (Figure S18).

Table 1. Synthesis and properties of gradient copolymers from EtVP and E-CL.



Entry	EtVP/ε-CL (mol/mol) ^ь	Polymer EtVP (%) ^c	M _{n,theo} (kDa)	M _{n,SEC} (kDa) ^d	Ð	Τ _g , Τ _m (°C)
1	0/100	0	22.9	54.0	1.7	n.d. ^e , 57.0
2	20/80	19	22.2	44.8	1.6	-60.5, 29.4
3	50/50	48	24.7	27.1	1.5	-50.8, none
4	80/20	74	26.4	23.7	1.2	-44.9 <i>,</i> none
5	100/0	100	25.1	17.8	1.2	-33.1, none

^aReaction conditions: 2.5 mol% TBD, 0.5 mol% 3-PPA, neat, 22 °C, 20 h. ^binitial molar ratios. ^ccalculated from ¹H NMR spectroscopy. ^ddetermined by THF SEC using polystyrene standards. ^enot determined.

Kinetic studies were undertaken to further understand the reactivity of **EtVP** with ϵ -CL (Figure S17). The initial rate of polymerization was found to be 3.3 M h⁻¹ at an **EtVP**: ϵ -CL molar ratio of 50:50, 5% TBD and 1% 3-PPA (Figure S17A). This is substantially faster than the homopolymerization of **EtVP** (1.44 M h⁻¹).³ Reactivity ratios were determined by utilizing a simple non-terminal model (Figure S19).²⁴ It was found that $r_{ttVP} = 1.37$ and $r_{\epsilon-CL} = 0.91$ when using a feed ratio of 40:60 **EtVP**: ϵ -CL, indicating the formation of a somewhat random copolymer with some gradient character. For reference, the reactivity ratios of monomers in the syntheses of *poly*(δ -valerolactone(VL)-*co*- ϵ -CL) and *poly*(LLA-*co*- ϵ -CL) gradient copolymers were found to be $r_{VL} = 12.92$, $r_{LLA} = 17.48$ and $r_{\epsilon-CL} = 0.10.^{24}$ A terminal model²⁵ was also explored, but no reasonable fit could be found for the obtained data set.

Next, block copolymerizations of **EtVP** and **\epsilon-CL** were attempted, envisioning that the resultant diblock copolymers would be more crystalline. Employing a sequential monomer addition strategy, block copolymers [*poly*(**CL**-*b*-**EtVP**)] with varying amounts of **EtVP** incorporation were synthesized (Table 2). In the sequential addition reactions, the order of monomer addition as well as the presence of solvent had a significant impact on the copolymer synthesized (Table S3). It was found that initial polymerization of ϵ -**CL** to full conversion in concentrated toluene (8-9 M) followed by subsequent chain extension through **EtVP** addition without pre-polymer purification resulted in the highest molar mass copolymers, and block copolymerizations under these conditions are reported in Table 2. Chain extension of purified *poly*(**EtVP**) was also explored and resulted in similar copolymers, but required more intensive purification steps and was less convenient (Figure S20). Molar masses and dispersities of *poly*(**CL**-*b*-**EtVP**) were similar to the gradient copolymers (Table 1) synthesized under identical feed ratios and catalyst/initiator loadings.

Table 2. Synthesis and properties of block copolymers from EtVP and ε-CL.



Entry	EtVP/ε-CL (mol/mol) ^b	EtVP Conversion (%) ^c	Polymer EtVP (%) ^c	M _{n,SEC} (kDa) ^d	Ð	Τ _{g,} Τ _m (°C)
1	0/100	-	0	54.0	1.7	n.d. ^e , 57.0
2	20/80	37	7	-	-	-
3	50/50	64	42	30.6	1.7	n.d. ^e , 54.5

4	80/20	67	74	17.3	1.5	n.d. ^e , 54.8
5	100/0	81	100	17.8	1.2	-33.1, none
^a Reaction	conditions: 2	2.5 mol% TBD,	0.5 mol% 3-PPA,	8-9 M toluene,	22 °C, 24 h.	^b initial molar

Reaction conditions. 2.5 mol% TBD, 0.5 mol% 5-PPA, 8-9 M toldene, 22 C, 24 H. mittal mola
ratios. ^c calculated from ¹ H NMR spectroscopy. ^d determined by THF SEC using polystyrene
standards. ^e not determined.

The microstructure of the gradient and block copolymers can be differentiated by ¹³C NMR spectroscopy (Figure 2). For the 50:50 poly(CL-b-EtVP) block copolymer (Figure 2, middle; from Table 2 Entry 3), two sets of resonances are present in the carbonyl region of the ¹³C NMR spectrum at 174.9 and 173.5 ppm. These signals are consistent with long blocks of each monomer in the block copolymer, as a mixture of poly(EtVP) and PCL homopolymers (Figure 2, top) has identical chemical shift values for each polymer. In contrast, the random/gradient copolymer poly(EtVP-q-CL) (Figure 2, bottom; from Table 1 entry 3) exhibits two additional sets of resonances at 175.8 and 172.7 ppm corresponding to the heterosequences EtVP-E-CL and E-CL-EtVP, respectively. These heterosequence resonances were assigned by ¹H-¹³C HMBC spectroscopy (Figure S11). The presence of EtVP-EtVP and E-CL-E-CL homosequences, as well as heterosequences and their ratios, further indicate that poly(EtVP-g-CL) is a gradient copolymer. Based on ¹³C NMR carbonyl integrations, 21.4% correspond to EtVP-EtVP, 25.4% to EtVP-ε-CL, 26.4% to ε-CL-EtVP and 26.5% to ε-CL-ε-CL in the 50:50 gradient copolymer. The average length of monomer block (I) and degree of randomness (R) were also determined for the gradient copolymer using ¹³C NMR spectroscopy.³⁷ It was found that $I_{cL} = 3.0$ and $I_{EtVP} = 2.9$ with $R_{cL} = 0.62$ and $R_{EtVP} = 0.76$; where R = 1 for a random copolymer and < 0.5 for a blocky copolymer. The thermal properties of the block copolymers were drastically impacted compared to the gradient copolymers. For example, in the case of 50:50 EtVP:ε-CL, a melting point of 54.5 °C was observed for the block copolymer (Table 2, entry 3), but no melting point was present for the gradient copolymer (Table 1, entry 3).



Figure 2. ¹³C NMR of 50:50 *poly*(**EtVP**-*g*-**CL**) (bottom), 50:50 *poly*(**CL**-*b*-**EtVP**) (middle) and a mixture of *poly*(**EtVP**) and **PCL** (top).

Encouraged by the ability to synthesize gradient and block copolymers with ϵ -CL, copolymerizations of EtVP with LLA were next explored (Table 3). LLA is not very soluble in EtVP, and as a result reactions were carried out with toluene as the solvent. Reactions were run as concentrated as possible owing to the low T_c of EtVP. Here, both concurrent polymerizations as well as sequential additions for block copolymer synthesis were attempted. Regardless of the feed strategy, molar masses of the EtVP/LLA copolymers were lower than the EtVP/ ϵ -CL copolymers. For concurrent addition (random) copolymers, *poly*(EtVP-*co*-LLA), a feed ratio of 50:50 EtVP:LLA resulted in the highest molar mass copolymer (Table 3, entry 2), and no clear molar mass trend was observed with molar feed ratios. A broader range of molecular weights and dispersities were found for the block copolymers, *poly*(EtVP-*b*-LLA) (Table 3, entries 4-6), compared to the random copolymers. For example, a maximum M_n of 27.3 kg/mol and a D = 1.5 was obtained with 21% EtVP incorporation in *poly*(EtVP-*b*-LLA) (Table 3, entry 4).

Table 3. Synthesis and properties of **EtVP** and **LLA** copolymers from concurrent and sequential addition. Inset graph: tensile testing of 40:60 **EtVP:LLA** block copolymer (5 trials).



Entry	Addition		LLVF	rorymen	IVI n,SEC	Ð	/ g
		(mol/mol) [♭]	Conversion (%) ^c	EtVP (%) ^c	(kDa) ^d		(°C) ^e
1	concurrent	20/80	74	23	13.7	1.6	30.0
2	concurrent	50/50	75	49	18.6	1.5	5.0
3	concurrent	80/20	69	77	15.6	1.9	-16.8
4	sequential	20/80	99	21	27.3	1.5	40.8
5	sequential	50/50	89	55	16.7	1.4	-25.6, 30.5
6	sequential	80/20	88	84	23.9	2.5	-22.9

^aReaction conditions: 2.5 mol% TBD, 0.5 mol% 3-PPA, 3.24 M toluene (respective to **LLA**), 22 °C, 25 h. ^binitial molar ratios. ^ccalculated from ¹H NMR spectroscopy. ^ddetermined by THF SEC using polystyrene standards.

All copolymers synthesized were amorphous, even when LLA incorporation was 79% (Table 3, entry 4). However, the T_g values were significantly impacted by LLA incorporation in both the random and block copolymers. For example, the T_g values of poly(EtVP-co-LLA) spanned from -16.8 °C with 77% EtVP incorporation to 30.0 °C with 23% EtVP incorporation (Table 3, entries 1-3). A similar trend can be seen poly(ϵ -CL-co-LLA), where T_g increased from -23 to 19 °C when the molar fraction of ϵ -CL decreased from 46% to 18%.²⁰ The T_g of the poly(EtVP-b-LLA) block copolymers increased from -22.9 to 40.8 °C when increasing from 16% to 79% LLA incorporation (Table 3, entries 4-6). While it is hard to directly compare $T_{\rm g}$ values of the two copolymer types due to a difference in the degrees of **EtVP** incorporation, it is expected that blocky poly (EtVP-b-LLA) would have higher T_g values than the random of poly (EtVP-co-LLA) copolymers based on comparison to PCL-PLLA copolymers³⁸. The microstructure of 50:50 LLA copolymers was investigated via ¹³C NMR spectroscopy (Figure S38 and S51). Resonances consistent with EtVP-LLA and LLA-EtVP heterosequences were the most predominant in poly(EtVP-co-LLA), although trace homosequences could be observed. This could provide evidence for a random copolymer when employing concurrent addition with EtVP-LLA, although reactivity ratios could not be established because of signal overlap between EtVP and LLA. For poly(EtVP-b-LLA), trace heterosequences were observed which can be explained by the increase in EtVP conversion after the addition of LLA.

Next, mechanical testing of the **LLA** copolymers was undertaken. Initially *poly*(**EtVP**-*co*-**LLA**) with a molar ratio of 50:50 was used for tensile testing. While this polymer exhibited excellent elasticity, its

tensile strength was very low. *Poly*(**EtVP**-*b*-**LLA**) with 40% **EtVP** was then investigated. After testing five samples, the average tensile strength was found to be 0.8 MPa with an elastic modulus of 5.6 MPa and an 83% elongation at break (Table 3 inset graph, right). While the elongation has significantly improved, tensile strength remains low compared to **PLLA** and **PLLA** copolymers.¹⁷

Finally, the chemical recyclability of **EtVP**-based copolymers was explored (Figure 3) employing the commonly used catalyst, tin(II) 2-ethylhexanoate (SnOct)₂. *Poly*(**EtVP**-*g*-**CL**) with 48% **EtVP** could be recycled at 185 °C under vacuum with 35% combined yield of the two monomers, whereas *poly*(**CL**-*b*-**EtVP**) was recycled with 61% combined yield of the two monomers. More **EtVP** was recovered relative to **ε**-**CL** in both cases (64% and 51% for gradient and block, respectively). The residual material for both copolymers was primarily unreacted **PCL** with some *poly*(**EtVP**) still present (Table S6). **LLA** copolymers resulted in lower recyclability, with only 27% yield for *poly*(**EtVP**-*co*-**LLA**) and 35% yield for *poly*(**EtVP**-*b*-**LLA**). In this case, residual material was a mixture of decomposition or cross-linked products (Figure S62). A mixture of **LLA** and meso-lactide was recovered with **EtVP**. Compared to the homopolymer *poly*(EtVP) (>85% recovery), monomer recovery of the copolymer systems is significantly lower; nonetheless, these results show the potential for chemical recycling of **EtVP**-based copolymers although a different catalyst system^{27,39,40} may be required for higher monomer recovery/selectivity.



Figure 3. Chemical recycling of EtVP based copolymers employing Sn(Oct)₂ as a depolymerization catalyst.

Conclusion

This work has expanded the synthetic polymer chemistry of the CO₂-derived lactone **EtVP** through ring-opening copolymerizations with ϵ -CL and LLA. Polymer properties and microstructures could be tuned through concurrent and sequential copolymerization strategies, which led to the formation of either block, gradient, or random copolymers. ϵ -CL block copolymers resulted in semi-crystalline polymers regardless of the molar ratio employed. For LLA, copolymers remained amorphous, and mechanical testing showed improved elasticity relative to PLLA. Furthermore, ϵ -CL and LLA copolymers could be chemically recycled back to monomer utilizing Sn(Oct)₂. While this work lays the foundation for EtVP-based copolymers, investigation into triblocks and other end-of-life options may further improve the potential applications of these CO₂-based (co)polymers.

Supporting Information

Full experimental details and data (.pdf) The Supporting Information is available free of charge on the ACS Publications website.

Acknowledgements

Funding for this work was provided by the NSF Center for Sustainable Polymers (CHE-1901635) and the University of Minnesota (Doctoral Dissertation Fellowship to R.M.R.). Instrumentation for the University of Minnesota Chemistry NMR facility was supported from a grant through the National Institutes of Health (S100D011952).

Corresponding Author

*itonks@umn.edu

References

- (1) Rapagnani, R. M.; Tonks, I. A. 3-Ethyl-6-Vinyltetrahydro-2H-Pyran-2-One (EVP): A Versatile CO2-Derived Lactone Platform for Polymer Synthesis. *Chem. Commun.* **2022**, *58* (69), 9586–9593. https://doi.org/10.1039/d2cc03516b.
- (2) Tang, S.; Lin, B. L.; Tonks, I.; Eagan, J. M.; Ni, X.; Nozaki, K. Sustainable Copolymer Synthesis from Carbon Dioxide and Butadiene. *Chem. Rev.* **2023**. https://doi.org/10.1021/acs.chemrev.3c00847.
- (3) Rapagnani, R. M.; Dunscomb, R. J.; Fresh, A. A.; Tonks, I. A. Tunable and Recyclable Polyesters from CO2 and Butadiene. *Nat. Chem.* **2022**, *14* (8), 877–883. https://doi.org/10.1038/s41557-022-00969-2.
- (4) Lou, Y.; Xu, L.; Gan, N.; Sun, Y.; Lin, B. L. Chemically Recyclable Polyesters from CO2, H2, and 1,3-Butadiene. *Innovation* **2022**, *3* (2), 100216. https://doi.org/10.1016/j.xinn.2022.100216.
- (5) Lou, Y.; Xu, J.; Xu, L.; Chen, Z.; Lin, B. L. Chemically Recyclable CO2-Based Solid Polyesters with Facile Property Tunability. *Macromol. Rapid Commun.* 2022, 43 (20), 1–8. https://doi.org/10.1002/marc.202200341.
- Bińczak, J.; Dziuba, K.; Chrobok, A. Recent Developments in Lactone Monomers and Polymer Synthesis and Application. *Materials (Basel)*. 2021, 14 (11), 1–20. https://doi.org/10.3390/ma14112881.
- Acik, G. Bio-Based Poly(ε-Caprolactone) from Soybean-Oil Derived Polyol via Ring-Opening Polymerization. J. Polym. Environ. 2020, 28 (2), 668–675. https://doi.org/10.1007/s10924-019-01597-7.
- (8) Pyo, S. H.; Park, J. H.; Srebny, V.; Hatti-Kaul, R. A Sustainable Synthetic Route for Biobased 6-Hydroxyhexanoic Acid, Adipic Acid and ε-Caprolactone by Integrating Bio- And Chemical Catalysis. *Green Chem.* **2020**, *22* (14), 4450–4455. https://doi.org/10.1039/d0gc01454k.
- Malikmammadov, E.; Tanir, T. E.; Kiziltay, A.; Hasirci, V.; Hasirci, N. PCL and PCL-Based Materials in Biomedical Applications. J. Biomater. Sci. Polym. Ed. 2018, 29 (7–9), 863–893. https://doi.org/10.1080/09205063.2017.1394711.
- Mishra, N.; Goyal, A.; Khatri, K.; Vaidya, B.; Paliwal, R.; Rai, S.; Mehta, A.; Tiwari, S.; Vyas, S.; Vyas, S. Biodegradable Polymer Based Particulate Carrier(s) for the Delivery of Proteins and Peptides. *Antiinflamm. Antiallergy. Agents Med. Chem.* **2008**, 7 (4), 240–251.

https://doi.org/10.2174/187152308786847816.

- (11) Pitt, C. G. *Biodegradable Polymers as Drug Delivery Systems*; 1991; Vol. 17. https://doi.org/10.3109/03639049109043830.
- (12) Abedalwafa, M.; Wang, F.; Wang, L.; Li, C. Biodegradable Poly-Epsilon-Caprolactone (PCL) for Tissue Engineering Applications: A Review. *Rev. Adv. Mater. Sci.* **2013**, *34* (2), 123–140.
- (13) Engelberg, I.; Kohn, J. Physico-Mechanical Properties of Degradable Polymers Used in Medical Applications: A Comparative Study. *Biomaterials* 1991, *12* (3), 292–304. https://doi.org/10.1016/0142-9612(91)90037-B.
- (14) Eshraghi, S.; Das, S. Mechanical and Microstructural Properties of Polycaprolactone Scaffolds with One-Dimensional, Two-Dimensional, and Three-Dimensional Orthogonally Oriented Porous Architectures Produced by Selective Laser Sintering. *Acta Biomater.* **2010**, *6* (7), 2467–2476. https://doi.org/10.1016/j.actbio.2010.02.002.
- (15) Pretula, J.; Slomkowski, S.; Penczek, S. Polylactides—Methods of Synthesis and Characterization. *Adv. Drug Deliv. Rev.* **2016**, *107*, 3–16. https://doi.org/10.1016/j.addr.2016.05.002.
- Kaihara, S.; Matsumura, S.; Mikos, A. G.; Fisher, J. P. Synthesis of Poly(L-Lactide) and Polyglycolide by Ring-Opening Polymerization. *Nat. Protoc.* 2007, 2 (11), 2667–2671. https://doi.org/10.1038/nprot.2007.391.
- (17) Farah, S.; Anderson, D. G.; Langer, R. Physical and Mechanical Properties of PLA, and Their Functions in Widespread Applications — A Comprehensive Review. *Adv. Drug Deliv. Rev.* 2016, 107, 367–392. https://doi.org/10.1016/j.addr.2016.06.012.
- (18) Rasal, R. M.; Hirt, D. E. Toughness Decrease of PLA-PHBHHx Blend Films upon Surface-Confined Photopolymerization. J. Biomed. Mater. Res. - Part A 2009, 88 (4), 1079–1086. https://doi.org/10.1002/jbm.a.32009.
- (19) Jikei, M.; Suga, T.; Yamadoi, Y.; Matsumoto, K. Synthesis and Properties of Poly(L-Lactide-Co-Glycolide)-b-Poly(ε-Caprolactone) Multiblock Copolymers Formed by Self-Polycondensation of Diblock Macromonomers. *Polym. J.* **2017**, *49* (4), 369–375. https://doi.org/10.1038/pj.2016.126.
- (20) Dakshinamoorthy, D.; Peruch, F. Block and Random Copolymerization of μ-Caprolactone, L-, and Rac-Lactide Using Titanium Complex Derived from Aminodiol Ligand. J. Polym. Sci. Part A Polym. Chem. 2012, 50 (11), 2161–2171. https://doi.org/10.1002/pola.25983.
- Qian, H.; Bei, J.; Wang, S. Synthesis, Characterization and Degradation of ABA Block Copolymer of L-Lactide and ε-Caprolactone. *Polym. Degrad. Stab.* 2000, 68 (3), 423–429. https://doi.org/10.1016/S0141-3910(00)00031-8.
- (22) Zhang, J.; Xu, L.; Xiao, W.; Chen, Y.; Dong, Z.; Xu, J.; Lei, C. Ring-Opening Polymerization of ε-Caprolactone with Recyclable and Reusable Squaric Acid Organocatalyst. *Eur. Polym. J.* **2021**, *157* (June), 110643. https://doi.org/10.1016/j.eurpolymj.2021.110643.
- (23) Naddeo, M.; Sorrentino, A.; Pappalardo, D. Thermo-Rheological and Shape Memory Properties of Block and Random Copolymers of Lactide and ε-Caprolactone. *Polymers (Basel).* 2021, 13 (4), 1– 19. https://doi.org/10.3390/polym13040627.
- (24) Beckingham, B. S.; Sanoja, G. E.; Lynd, N. A. Simple and Accurate Determination of Reactivity

Ratios Using a Nonterminal Model of Chain Copolymerization. *Macromolecules* **2015**, *48* (19), 6922–6930. https://doi.org/10.1021/acs.macromol.5b01631.

- (25) Lynd, N. A.; Ferrier, R. C.; Beckingham, B. S. Correction: Recommendation for Accurate Experimental Determination of Reactivity Ratios in Chain Copolymerization (Macromolecules (2019) 52: 6 (2277–2285) DOI: 10.1021/Acs.Macromol.8b01752). *Macromolecules* 2019, *52* (20), 7961. https://doi.org/10.1021/acs.macromol.9b01984.
- Jin, M.; Hoye, T. R. Lactone Ring-Opening Equilibria in Methanol by 1H NMR Analysis: An Assessment of the Ring-Opening Polymerizability of Lactone Monomers. *Macromolecules* 2023, 56 (3), 1122–1129. https://doi.org/10.1021/acs.macromol.2c01141.
- (27) Su, J.; Xu, G.; Dong, B.; Yang, R.; Sun, H.; Wang, Q. Closed-Loop Chemical Recycling of Poly(∈-Caprolactone) by Tuning Reaction Parameters. *Polym. Chem.* 2022, 13 (41), 5897–5904. https://doi.org/10.1039/d2py00953f.
- (28) Li, Y.; Thouas, G. A.; Chen, Q. Z. Biodegradable Soft Elastomers: Synthesis/Properties of Materials and Fabrication of Scaffolds. *RSC Adv.* 2012, 2 (22), 8229–8242. https://doi.org/10.1039/c2ra20736b.
- Yue, S.; Bai, T.; Xu, S.; Shen, T.; Ling, J.; Ni, X. Ring-Opening Polymerization of CO2-Based
 Disubstituted δ-Valerolactone toward Sustainable Functional Polyesters. ACS Macro Lett. 2021, 10 (8), 1055–1060. https://doi.org/10.1021/acsmacrolett.1c00341.
- (30) Chen, K.; Zhu, Z.; Bai, T.; Mei, Y.; Shen, T.; Ling, J.; Ni, X. A Topology-Defined Polyester Elastomer from CO2 and 1,3-Butadiene: A One-Pot-One-Step "Scrambling Polymerizations" Strategy. *Angew. Chemie - Int. Ed.* 2022, *61* (46). https://doi.org/10.1002/anie.202213028.
- (31) Wang, Z.; Zheng, W.; Yue, S.; Chen, K.; Ling, J.; Ni, X. Random Terpolymer of Carbon Dioxide, Butadiene and Epoxides: Synthesis, Functionalization and Degradability. *Chinese J. Chem.* **2024**. https://doi.org/10.1002/cjoc.202400050.
- (32) Zhao, Y.; Zhang, X.; Li, Z.; Li, Z.; Tang, S. Functional and Degradable Polyester-Co-Polyethers from CO2, Butadiene, and Epoxides. ACS Macro Lett. 2024, 315–321. https://doi.org/10.1021/acsmacrolett.4c00071.
- (33) Garcia Espinosa, L. D.; Williams-Pavlantos, K.; Turney, K. M.; Wesdemiotis, C.; Eagan, J. M.
 Degradable Polymer Structures from Carbon Dioxide and Butadiene. ACS Macro Lett. 2021, 10 (10), 1254–1259. https://doi.org/10.1021/acsmacrolett.1c00523.
- (34) Dove, A. P. Organic Catalysis for Ring-Opening Polymerization. *ACS Macro Lett.* **2012**, *1* (12), 1409–1412. https://doi.org/10.1021/mz3005956.
- (35) Pratt, R. C.; Lohmeijer, B. G. G.; Long, D. A.; Waymouth, R. M.; Hedrick, J. L. Triazabicyclodecene: A Simple Bifunctional Organocatalyst for Acyl Transfer and Ring-Opening Polymerization of Cyclic Esters. J. Am. Chem. Soc. 2006, 128 (14), 4556–4557. https://doi.org/10.1021/ja060662+.
- (36) Martello, M. T.; Burns, A.; Hillmyer, M. Bulk Ring-Opening Transesterification Polymerization of the Renewable δ-Decalactone Using an Organocatalyst. ACS Macro Lett. 2012, 1 (1), 131–135. https://doi.org/10.1021/mz200006s.
- (37) Fernández, J.; Meaurio, E.; Chaos, A.; Etxeberria, A.; Alonso-Varona, A.; Sarasua, J. R. Synthesis and Characterization of Poly (I-Lactide/ε-Caprolactone) Statistical Copolymers with Well Resolved

Chain Microstructures. *Polymer (Guildf)*. **2013**, *54* (11), 2621–2631. https://doi.org/10.1016/j.polymer.2013.03.009.

- (38) Nalampang, K.; Molloy, R.; Punyodom, W. Synthesis and Characterization of Poly(L-Lactide-Co-ε-Caprolactone) Copolymers: Influence of Sequential Monomer Addition on Chain Microstructure. *Polym. Adv. Technol.* **2007**, *18* (3), 240–248. https://doi.org/10.1002/pat.880.
- (39) Alberti, C.; Enthaler, S. Depolymerization of End-of-Life Poly(Lactide) to Lactide via Zinc-Catalysis. *ChemistrySelect* **2020**, *5* (46), 14759–14763. https://doi.org/10.1002/slct.202003979.
- McGuire, T. M.; Buchard, A.; Williams, C. Chemical Recycling of Commercial Poly(I-Lactic Acid) to I-Lactide Using a High-Performance Sn(II)/Alcohol Catalyst System. J. Am. Chem. Soc. 2023, 145 (36), 19840–19848. https://doi.org/10.1021/jacs.3c05863.