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3D printing *via* polymerization-induced microphase separation using acrylate macromonomers instead of macroRAFT agents†

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Polymerization-induced microphase separation (PIMS) is a versatile technique for manufacturing nanostructured materials. Combining PIMS with 3D printing enables the fabrication of complex objects with nanoscale features, opening possibilities in diverse applications, including nanostructured ceramics, solid polymer electrolytes, and ion-exchange materials. Traditionally, PIMS utilizes polymers synthesized by reversible addition–fragmentation chain transfer (RAFT) polymerization (macromolecular Chain Transfer Agents, macroCTAs). However, RAFT-based PIMS can introduce undesirable color and odor into the final materials. To address these limitations, this study explores the use of macromonomers, polymers terminated with acrylate or methacrylate groups, as alternatives to macroCTAs. We synthesized a series of polycaprolactone (PCL) variants with identical molecular weights but differing terminal functionalities: acrylate, methacrylate, trithiocarbonate, and dithiobenzoate. This library enabled a direct comparison of macromonomer and macroCTA approaches for nanostructured material fabrication *via* PIMS. Small-angle X-ray scattering (SAXS) was employed to determine nanodomain sizes. Notably, both acrylate and trithiocarbonate-terminated PCLs yielded comparable nanodomain sizes. Exploiting PCL degradation, we fabricated nanoporous, 3D-printed objects by selectively etching the PCL from materials formed with both trithiocarbonate and acrylate-terminated PCL. Critically, the acrylate-terminated macromonomer-based PIMS system produced transparent, colorless materials with well-defined microstructures. This work demonstrates the potential of macromonomers to overcome the inherent limitations of RAFT–PIMS, providing a cleaner and more versatile pathway to synthesize advanced nanostructured materials.

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Introduction

Additive manufacturing, commonly known as 3D printing, has significantly simplified the fabrication of a wide range of materials with complex shapes and customizable properties.^{1,2} More recently, advancements in controlling structures at multiple scales, particularly at both the nano- and macroscale, have further expanded its potential.^{3–9} One notable approach involves polymerization-induced phase separation (PIPS), which generates distinct regions or phases within the material in the microscopic range (from 50 nm to micrometres). This process occurs during polymerization, where phase separation takes place as the polymerization reaction progresses. The key concept behind PIPS is that the growing polymer chains

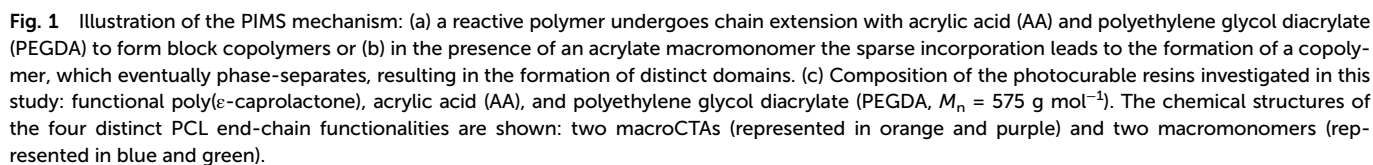
become increasingly incompatible with the surrounding medium (often the monomer or solvent), leading to phase separation.^{10,11} Levkin and coworkers pioneered the combination of digital light processing (DLP) and PIPS to form 3D-printed hierarchical macro- and mesoporous materials, which can be employed as super-hydrophobic materials, in gas separation, and water repellents for microfluidic applications, as well as superoleophilic materials for the development of oil-absorbent systems.¹²

Inspired by the early work from Seo and Hillmyer on polymerization induced microphase separation (PIMS),¹³ we introduced a novel 3D printing technology by integrating photoinduced reversible addition–fragmentation chain-transfer (RAFT) with PIMS in photocuring 3D printing, enabling a nanoscopic scale below 100 nm.¹⁴ The PIMS methodology relies on an *in situ* chain-extension reaction of a macromolecular chain transfer agent (macroCTA) with a monomer and a crosslinker, resulting in the formation of a “block” copolymer. This reaction induces microphase separation between the incompatible segments, while simultaneous crosslinking arrests the separation and forms disordered nanostructured materials (Fig. 1a). The resulting nanostructured materials

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†Electronic supplementary information (ESI) available: Materials and methods, NMR spectra of purified poly(ϵ -caprolactone), molecular weight distributions obtained by SEC of modified poly(ϵ -caprolactone)s, FTIR spectra of modified poly(ϵ -caprolactone)s, UV-vis spectra, and SAXS profiles (Fig. S1–S34 and Tables S1–S4). See DOI: <https://doi.org/10.1039/d5py00226e>





To achieve this, poly(ϵ -caprolactone) (PCL-OH) was selected as a model polymer and synthesized *via* ring-opening polymerization (ROP). The terminal hydroxyl group of PCL was then functionalized with either RAFT agents or acrylate/methacrylate end-groups. The RAFT agents, 4-cyano-4-[(dodecylsulfate

Results and discussion

To study the effect of the end group on the PIMS process and minimize the influence of macroCTA chain length on both the

Table 1 Properties of the synthesized end-group modified poly(ϵ -caprolactone) polymers

Polymer	SEC (RI, DMAC) ^a		¹ H NMR		
	M_n^{Corr} (g mol ⁻¹)	$\bar{D}$	X_n^b	$M_n^{\text{NMR } c}$ (g mol ⁻¹)	End group fidelity, f^d (%)
PCL _{3k} -OH	3350	1.15	25	3182	—
PCL _{5.5k} -OH	3800	1.17	47	5643	—
PCL _{11k} -OH	7380	1.06	99	11 805	—
PCL _{3k} -CPADB	3890	1.17	28	3788	70
PCL _{3k} -BTPA	4100	1.17	30	3866	100
PCL _{5.5k} -BTPA	4400	1.35	53	6438	90
PCL _{11k} -BTPA	7400	1.07	97	11 538	100
PCL _{3k} -acrylate	3600	1.13	27	3323	100
PCL _{5.5k} -acrylate	4500	1.27	46	5600	65
PCL _{11k} -acrylate	8100	1.13	102	11 850	54
PCL _{3k} -methacrylate	4400	1.36	27	3389	71

^a Determined by DMAC-SEC analysis using PS standards and applying a correction factor of 0.56.⁴⁰ ^b X_n calculated from the ¹H NMR spectrum: $X_n = I_{4.05 \text{ ppm}/2}$. ^c $M_n^{\text{NMR } c} = X_n \times M_w(\text{CL}) + M_w(\text{benzyl alcohol})$. ^d End group fidelity calculated using NMR:

$$f(\%) = \frac{I_{\text{end-chain}}}{I_{\text{initiator}}(5.11 \text{ ppm})/2} \times 100.$$

morphology and nanoscale size, a two-step synthesis approach was employed to ensure consistent molecular weight across all the macroCTAs and macromonomers. In the first step, poly(ϵ -caprolactone) (PCL-OH) polymers with molecular weights ranging from 3000 to 11 000 g mol⁻¹ were synthesized *via* thermal ring opening polymerization (ROP) following previous procedures.³⁸ Benzyl alcohol (BnOH) was selected as the initiator to facilitate the determination of molecular weight by NMR, and tin(II) 2-ethylhexanoate (Sn(oct)₂) acted as the catalyst (Fig. 2a). The ROP method was preferred over polycondensation to achieve a narrow molecular weight distribution ($\bar{D} < 1.2$).³⁹ Polymerization proceeded to near-complete conversion, as determined by ¹H NMR by monitoring the shift of the H_e signal from 4.28 ppm to 4.14 ppm (Fig. S1†). Upon reaching 99% monomer conversion, the resulting polymers were purified by precipitation (Table 1, Fig. S2–S7†). ¹H NMR was also employed to calculate the degree of polymerization (X_n) of purified polymers by comparing the polycaprolactone signals at 4.14 ppm (H_e) with that of the CH₂ of the benzyl group (H_b) from the initiator. Size exclusion chromatography (SEC) confirmed monodisperse molecular weight distributions for the different PCLs, although a slight discrepancy between the molecular weights determined by NMR and SEC was observed, likely due to the use of polystyrene standards in SEC (Fig. S2–S7†). In the second step, the terminal hydroxyl group of the PCL was functionalized. For RAFT-terminated PCLs, coupling reactions with BTPA and CPADB in the presence of 1-ethyl-3-(3-dimethylaminopropyl)carbodiimide (EDC) and 4-dimethylaminopyridine (DMAP) yielded PCL-BTPA and PCL-CPADB macroCTAs (Table 1, Fig. 2, Fig. S8–S15†). For vinyl-terminated PCL, reactions with 2-isocyanatoethyl methacrylate (IEM) or 2-isocyanatoethyl acrylate (IEA) produced PCL-acrylate and

PCL-methacrylate macromonomers, respectively (Table 1, Fig. 2, Fig. S16–S23†).

The resulting PCLs were purified and characterized by ¹H NMR and DMAC-SEC (Fig. 2b, Fig. S8–S23†). For PCL-BTPA and PCL-CPADB macroCTAs, ¹H NMR spectra (shown in Fig. 2b) confirmed the presence of characteristic signals corresponding to BTPA and CPADB. Specific signals at 1.93 ppm (H_j) and 2.67 ppm (H_i) confirm the presence of CPADB (purple line, Fig. 2b and Fig. S8†), while signals at 0.98 ppm (H_p) and 4.81 ppm (H_n) indicated the presence of BTPA in PCL-BTPA (orange line, Fig. 2b and Fig. S14†). For PCL_{3k}-acrylate, the ¹H NMR spectrum exhibited vinyl bond signals (H_u, H_t and H_{u'}) at 6.44, 6.15 and 5.86 ppm, respectively (blue line, Fig. 2b and Fig. S16†). Similarly, the two characteristic vinyl signals of PCL_{3k}-methacrylate were observed clearly at 6.12 ppm (H_v) and 5.58 ppm (H_{v'}) (green line, Fig. 2b and Fig. S22†). End-chain fidelity, detailed in Table 1, was determined by comparing these end-group signal integrals to the -CH₂ (H_b) at 5.11 ppm from the benzyl initiator fragment. Despite general success, ¹H NMR also revealed incomplete end-capping in certain samples, indicated by a residual terminal -CH₂OH signal (H_g at 3.64 ppm). This was apparent for PCL-methacrylate and PCL-CPADB (Fig. 2b), and notably more pronounced for higher molecular weight PCL-acrylates (5.5 and 11 kg mol⁻¹; Fig. S18 and S20†). This likely arises from either lower reactivity of the functionalizing agent or sterically hindered access to the hydroxyl terminus on longer chains. Complementary techniques provided further verification. DMAC-SEC equipped with a UV detector (310 nm) was employed to verify the presence of the trithiocarbonate or dithiobenzoate groups on the PCL (Fig. S8–S15†). Fourier-Transform Infrared (FTIR) spectroscopy further confirmed the presence of a double bond at $\nu = 1529 \text{ cm}^{-1}$ (Fig. S24†).

Photocurable resin formulations containing functionalized PCLs ($M_n = 3000 \text{ g mol}^{-1}$), acrylic acid (AA) as the monomer, poly(ethylene glycol) diacrylate (PEGDA, average $M_n = 575 \text{ g mol}^{-1}$) as the cross-linker, and diphenyl(2,4,6-trimethylbenzoyl) phosphine oxide (TPO) as the photo-initiator were prepared. AA was selected as the resulting PAA block polymer can phase separate, due to the segregation strength. The segregation strength χN is used as a measure of the thermodynamic driving force, where χ is the measure of the incompatibility between the two blocks and N is the total degree of polymerization.^{41,42} For the PCL-*b*-P(AA-*stat*-PEGDA) block copolymer, χ is estimated to be 0.92 at 25 °C indicating that the PCL and P(AA-*stat*-PEGDA) are thermodynamically incompatible (ESI, Note 1†).⁴³

Components were mixed and sonicated to achieve transparent, homogeneous solutions. Polymerization kinetics were determined *via* assessment using total reflectance-Fourier transform infrared (ATR-FTIR) spectroscopy under violet light irradiation ($\lambda_{\text{max}} = 405 \text{ nm}$ and $I_0 = 2.08 \text{ mW cm}^{-2}$) under open-to-air conditions. The molar ratio of [AA]/[PEGDA] and the weight percentages (wt%) of PCL_{3k}-R and TPO were fixed at 9/1, 21 wt%, and 1 wt%, respectively. Vinyl bond conversion was monitored by tracking the decrease in the absorption peak



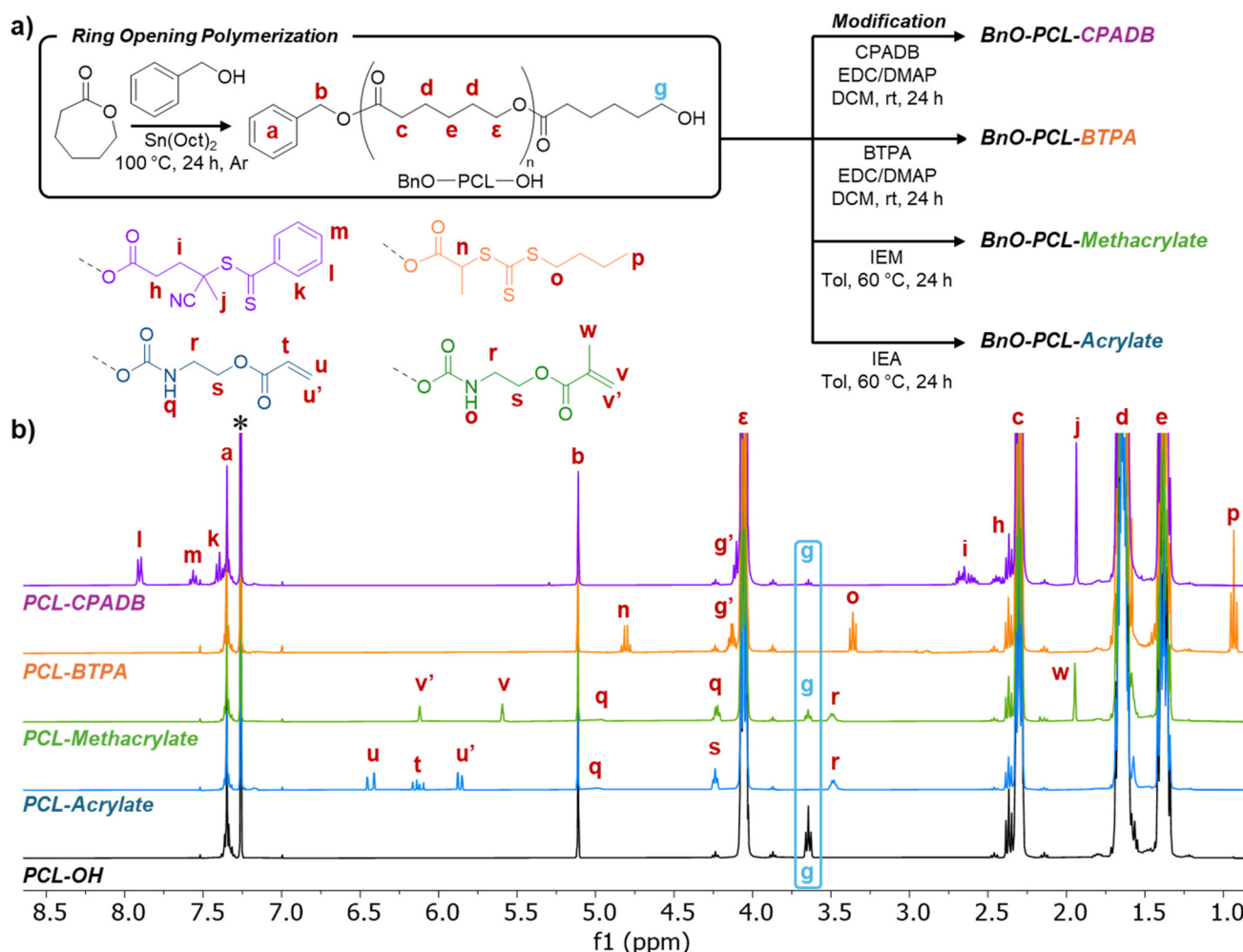


Fig. 2 Synthesis and characterization of functionalized PCL polymers. (a) Synthetic route for RAFT-terminated-PCLs (PCL-BTPA and PCL-CPADB) and acrylate/methacrylate-terminated-PCL (PCL-acrylate and PCL-methacrylate) via a two-step process. (b) ^1H NMR spectra of the starting PCL-OH and the four modified PCL polymers: PCL-acrylate, PCL-methacrylate, PCL-BTPA and PCL-CPADB (CDCl_3 (*), 400 MHz, 298 K).

at $\approx 1620\text{ cm}^{-1}$, corresponding to the vinylic $\text{C}=\text{C}$ stretching mode. As shown in Fig. 3a, resins formulated with acrylate- or methacrylate-functionalized PCL cured rapidly, reaching high vinyl bond conversion ($\geq 90\%$) within 20 seconds. In contrast, the PCL_{3k} -BTPA containing resin required over 45 seconds to achieve similar conversion (Fig. 3a). This kinetic difference is attributed to the trithiocarbonate group in PCL_{3k} -BTPA, which absorbs light at 405 nm , thereby reducing TPO photoinitiator activation. Furthermore, the PCL_{3k} -CPADB resin showed minimal vinyl bond conversion even after one minute of violet light exposure, consistent with previous reports of CPADB's ineffectiveness under these conditions. This ineffectiveness is due to its low fragmentation efficiency with acrylates and its susceptibility to rapid photodegradation (Fig. S25†).⁴⁴

Next, the formulated resins were 3D printed using an AnyCubic Photon on Mono SE ($\lambda_{\text{max}} = 405\text{ nm}$ and $I_0 = 2.3\text{ mW cm}^{-2}$) to fabricate simple objects, such as prism or discs (Fig. 3, Fig. S25, Table S1†). For consistency, the $[\text{AA}]/[\text{PEGDA}]$ molar ratio was fixed at 9/1, and the TPO concentration was

1 wt% for all the formulations. The PCL_{3k} -R content was varied from 15 to 35 wt% (Table S1†). However, the PCL_{3k} -CPADB resin required a higher irradiation intensity ($I_0 = 20\text{ mW cm}^{-2}$) to cure the resin, producing a yellow and very brittle object (Fig. 3c). These printed samples were then characterized by small-angle X-ray scattering (SAXS) to investigate their internal nanostructure and phase separation. Initial visual inspection revealed that PCL-BTPA and PCL-acrylate resins yielded transparent samples, while PCL-CPADB and PCL-methacrylate resins produced translucent samples, suggesting the formation of large domains ($>200\text{ nm}$).

Fig. 4 presents the SAXS profiles of 3D-printed objects as a function of macroCTA or macromonomer content. Resins formulated with PCL_{3k} -BTPA and PCL_{3k} -acrylate exhibited broad single scattering peaks across all loadings, indicating the formation of disordered microphase-separated materials (Fig. 4a and c). The scattering peak position shifted to higher q values with an increase in PCL content, consistent with the observations in other similar systems.²⁴ For instance, with 21 wt%



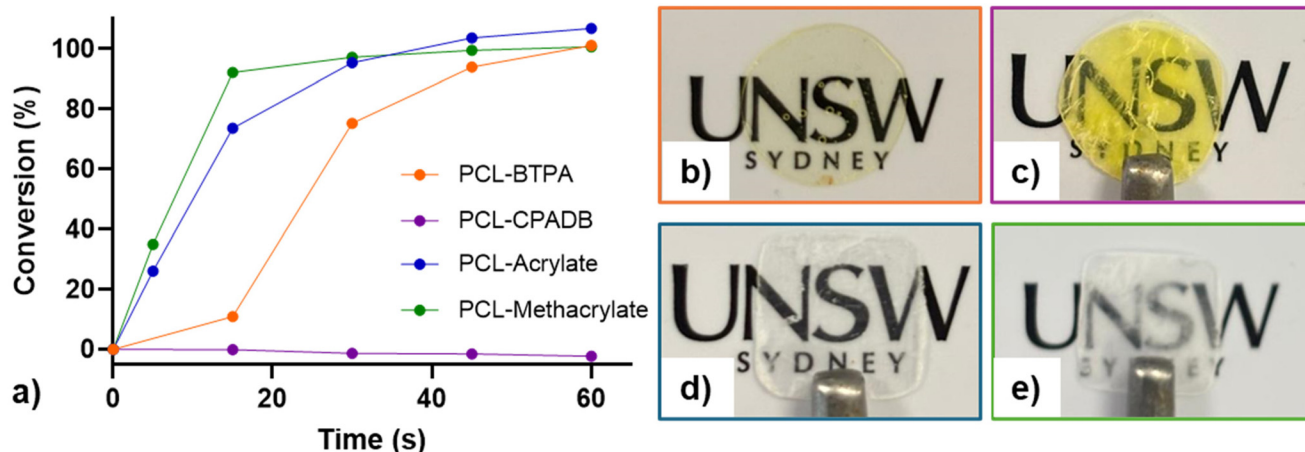


Fig. 3 Effect of the PCL end-group on photopolymerization kinetics and 3D printing. (a) Vinyl bond conversion as a function of time for resins containing different PCL polymers. Photographs of the resulting films prepared from the corresponding resins: (b) PCL_{3k}-BTPA, (c) PCL_{3k}-CPADB, (d) PCL_{3k}-acrylate, and (e) PCL_{3k}-methacrylate. Resin formulations: [AA]/[PEGDA] = 9/1 (molar ratio), PCL-R = 21 wt%, and TPO = 1 wt%. Kinetics experiments were performed in triplicate. Double bond conversion was monitored by ATR-FTIR under 2.08 mW cm⁻² violet light irradiation (λ_{max} = 405 nm).

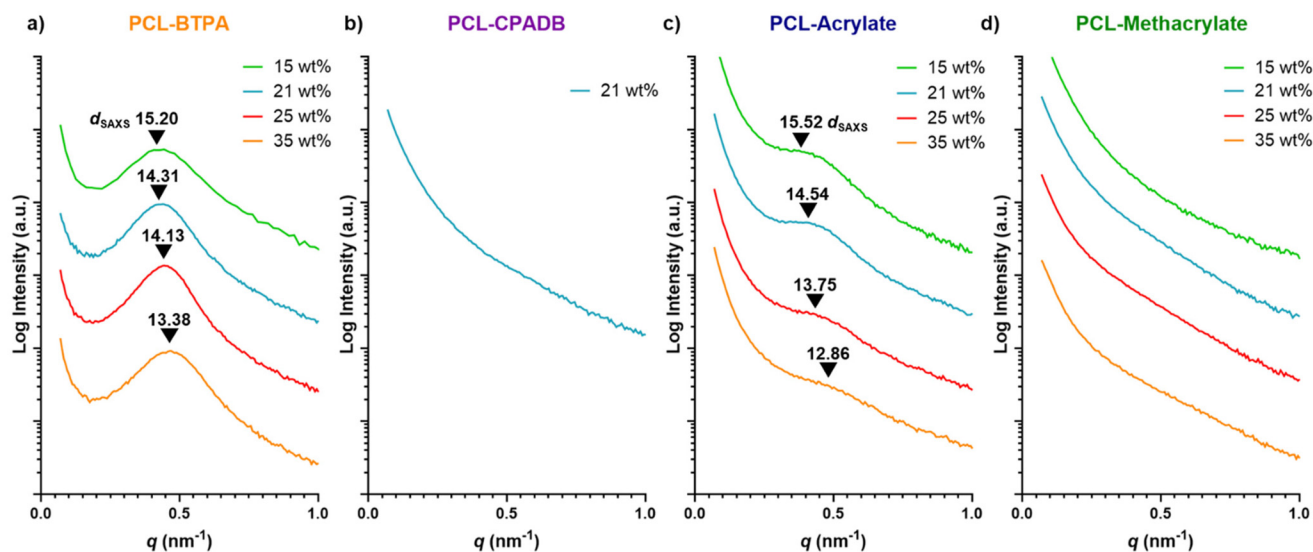


Fig. 4 SAXS profiles and the corresponding domain spacing (d_{SAXS}) values for materials 3D printed with varying loadings of (a) PCL_{3k}-BTPA, (b) PCL_{3k}-CPADB, (c) PCL_{3k}-acrylate, and (d) PCL_{3k}-methacrylate. Loading: 15 wt% (green), 21 wt% (blue), 25 wt% (red) and 35 wt% (orange). [AA]/[PEGDA] = 9/1 (molar ratio); TPO = 1 wt%.

loading for PCL_{3k}-BTPA, the peak appeared at 0.439 nm⁻¹, corresponding to a d_{SAXS} value of 14.31 nm, shifting to 0.469 nm⁻¹ (d_{SAXS} of 13.38 nm) at 35 wt% (Fig. 4a). A similar trend was observed for PCL-acrylate, with the peak shifting from 0.432 nm⁻¹ (d_{SAXS} = 14.54 nm) at 20 wt% to 0.488 nm⁻¹ (d_{SAXS} = 12.86 nm) at 35 wt% (Fig. 4c). Furthermore, comparing the two, the PCL_{3k}-BTPA signals were sharper and more intense, suggesting differences in phase separation efficiency; this may reflect more dispersed nanodomains resulting from the free radical polymerization of PCL-acrylate compared to the controlled radical polymerization mechanism of PCL-BTPA. In contrast, the SAXS profiles of objects made with

PCL_{3k}-CPADB and PCL_{3k}-methacrylate showed no discernible scattering peaks (Fig. 4b and d). This lack of features strongly indicates macrophase separation, with domain sizes exceeding the typical SAXS detection limit (>~100 nm). These results are attributed to the inefficiency of CPADB in promoting chain extension for the PCL-CPADB system due to its degradation, and the low reactivity of methacrylate groups during copolymerization with acrylates (AA and PEDGA) for the PCL-methacrylate system.

The observed variations in morphologies can be explained by the reactivity of the PCL end-chains. The trithiocarbonate group in PCL_{3k}-BTPA is a highly efficient chain transfer agent



in acrylate polymerization, which promotes effective chain extension, leading to the formation of diblock copolymers that undergo nanoscale phase separation. Conversely, the dithiobenzoate group in PCL_{3k}-CPADB is a less efficient chain transfer agent with acrylates and acrylamides,^{45,46} resulting in macrophase separation.^{25,28,29,47} Additionally, this group is more susceptible to photolysis, as evidenced by the color change from reddish to yellowish observed after printing and by the decrease in the characteristic 360 nm absorbance peak of the dithiobenzoate group (Fig. S25 and S26†). Building on these encouraging results, we also investigated resins containing PCL_{xk}-BTPA with molecular weights $M_n = 5.5$ and 11 kg mol⁻¹. The printed samples from these resins exhibited a single scattering peak that shifted to a lower q value with an increase in X_n . This shift corresponds to an increase in domain spacing, consistent with the formation of higher molecular weight block copolymers (Fig. S27 and S28†). However, the resins formulated with PCL-acrylate (5.5k and 11k) did not produce high-quality printed objects suitable for SAXS analysis.

While these initial results were encouraging, we encountered gelation of the resins within the vat after only a few minutes of 3D printing, preventing the preparation of larger and more complex objects. This effect was particularly prominent with the higher molecular weight PCL (PCL_{5.5k}-BTPA and PCL_{11k}-BTPA) (Fig. S29†). We hypothesized that this issue was due to the low miscibility of PCL in the AA/PEGDA mixture. To mitigate this and improve resin stability for longer print times, we investigated two strategies: replacing AA with alternative monomers or adding a co-solvent to enhance PCL solubility. Initially, we tested several alternative monomers, including methyl acrylate (MA), *N,N*-dimethylacrylamide (DMA) and isobornyl acrylate (IBoA) (Fig. 5a and Table S2†). SAXS analysis of the resulting materials revealed microstructures in all cases, as evidenced by the broad, single scattering peaks characteristic of disordered microphase-separated systems (Fig. 5a). Interestingly, normalization of the SAXS scattering plots by intensity (I^*) revealed that the peak breadth increased with DMA, MA, and IBoA compared to the AA formulation. This suggests that these alternative monomers led to the formation of more polydisperse nanodomains compared to AA, which yielded sharper interfaces. However, despite these promising microstructures, the gelation issue during 3D printing persisted with all tested monomers, hindering their practical application in 3D printing.

As an alternative approach, we investigated the use of a solvent to enhance PCL solubility in our resin formulations. Toluene was selected due to the high solubility of PCL. It was added at 10 wt% and 40 wt% concentrations. Remarkably, 10 wt% toluene alone successfully eliminated gelation, resulting in a stable resin suitable for extended printing (Fig. S29†). Having successfully resolved the gelation problem, we then examined the photopolymerization kinetics of the 10 wt% toluene-containing resins before fabricating complex objects. Subsequent photopolymerization kinetic studies on the 10 wt% toluene-containing resins, maintaining a constant

[AA]/[PEGDA] molar ratio (9/1), PCL_{3k}-R weight percentage (19 wt%), and TPO concentration (1 wt%), demonstrated similar kinetic profiles to those of the toluene-free formulations (Fig. S30†). Specifically, resins formulated with PCL-acrylate and PCL-methacrylate achieved near-complete double-bond conversion within 10 seconds, while PCL-BTPA-based resins required approximately 40 seconds to reach a comparable conversion level.

SAXS profiles of the 3D-printed samples revealed that while toluene addition did not significantly alter the broadness of the SAXS signals, it induced a shift in the maximum scattering peak to higher q values, indicating a decrease in d_{SAXS} from 14.11 nm to 13.36 nm as the toluene concentration increased from 10% to 40% (Fig. 5b and e, Table S3†). Moreover, at higher X_n , the SAXS signal became sharper, suggesting the formation of more well-defined nanodomains. A similar trend was noted with PCL_{xk}-acrylate with 10 wt% toluene, with d_{SAXS} values increasing from 12.09 nm (PCL_{3k}-acrylate) to 14.82 nm (PCL_{11k}-acrylate) with a significant change in the SAXS intensity. Interestingly, the SAXS signal intensity of samples prepared with 40 wt% toluene appeared slightly reduced in comparison with the one with 10 wt% toluene (Fig. 5c and Fig. S31†). Conversely, PCL-methacrylate and PCL-CPADB formulations remained opaque/translucent and exhibited no discernible SAXS signals upon toluene addition.

Moreover, the mechanical properties of these formulations were investigated, which revealed that samples prepared with PCL_{3k}-BTPA exhibited a slightly lower storage modulus and a $\tan \delta$ peak which shifted to a lower temperature compared to those prepared with PCL_{3k}-acrylate, when both used AA and PEGDA (Fig. S32†). Interestingly, the $\tan \delta$ peak temperature for the PCL_{3k}-BTPA samples shifted significantly higher, from approximately 0 °C to approximately 30 °C, when IBoA and PEGDA were used instead of AA and PEGDA (Fig. S33†).

Subsequently, we assessed the influence of end-chain functionality on printing resolution and macroscopic characteristics by fabricating objects with complex geometries. Resins formulated with PCL_{3k}-BTPA, PCL_{3k}-CPADB, PCL_{3k}-acrylate, and PCL_{3k}-methacrylate (19 wt%) in the presence of 10 wt% toluene and [AA]/[PEGDA] equal to 9/1 were tested alongside a control resin containing unmodified PCL (PCL-OH). Complex snowflake structures (28.9 × 25.7 × 2.0 mm) were successfully printed using a layer thickness of 50 μm, except for the resin based on PCL_{3k}-CPADB which failed to produce acceptable prints. PCL-BTPA and PCL-OH formulations required 30 second exposure time, while the acrylate- and methacrylate-functionalized PCL resins produced high-resolution prints with a shorter 20 second exposure time (Fig. 6).

The objects shown in Fig. 6 exhibit notable differences in transparency and color, which can be attributed to variations in phase separation behavior and the presence or absence of the RAFT agent. Objects fabricated with PCL_{3k}-BTPA and PCL_{3k}-acrylate were transparent (Fig. 6b and c). However, the RAFT-based formulation (PCL_{3k}-BTPA) exhibited a yellowish tint, while the PCL_{3k}-acrylate object was colorless. In contrast, objects printed with PCL_{3k}-methacrylate appeared translucent,



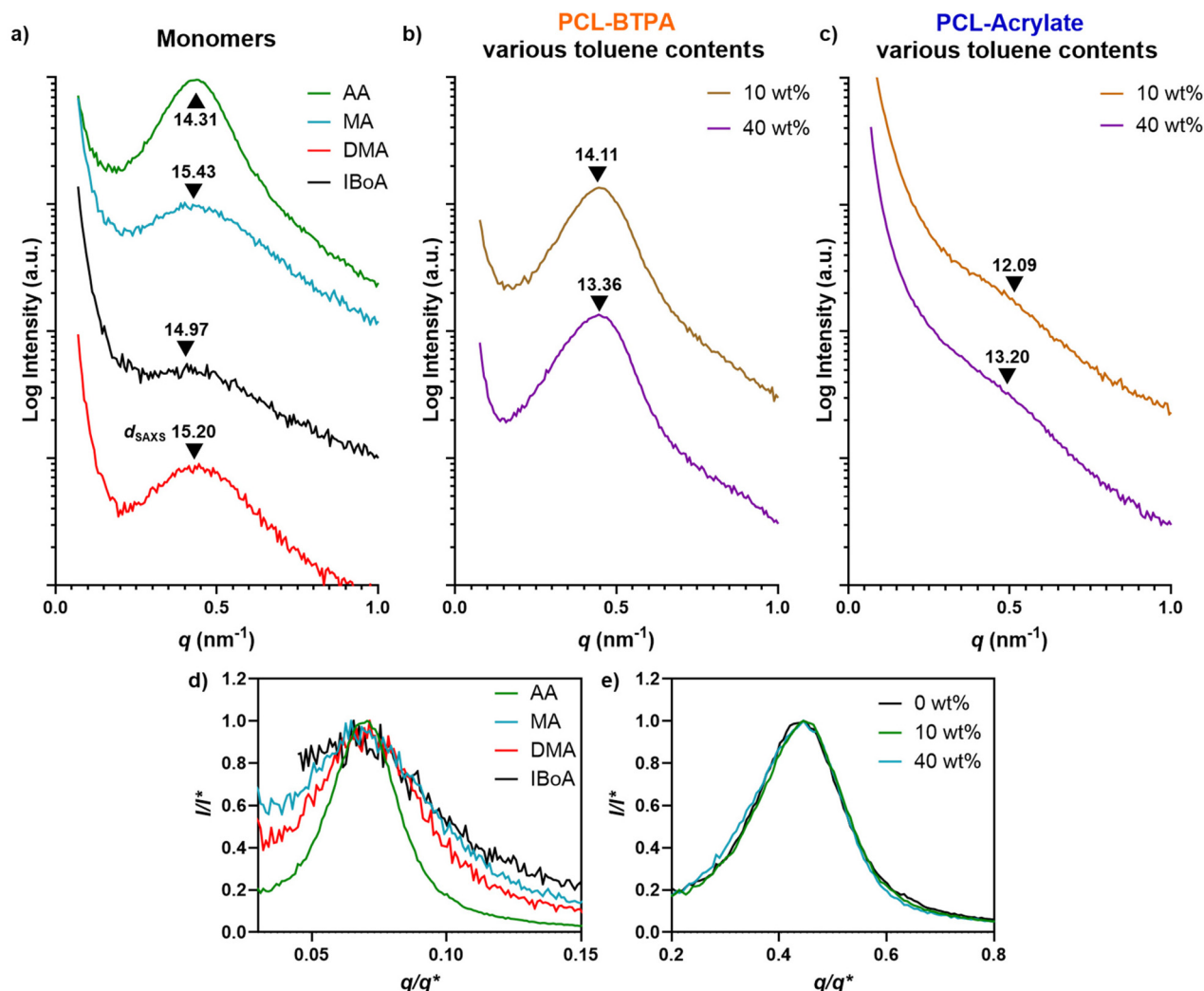


Fig. 5 Effect of the monomer and co-solvent on microphase separation. SAXS profiles and the corresponding domain spacing (d_{SAXS}) values of 3D printed materials using (a) PCL_{3k}-BTPA with different co-monomers (AA, MA, DMA, and IBoA), (b) PCL_{3k}-BTPA with AA and toluene, and (c) PCL_{3k}-acrylate with AA and toluene. (d and e) Scaled SAXS data (normalized intensity, I^*) comparing peak broadness and position (q) for PCL_{3k}-BTPA with different co-monomers and toluene contents, respectively. Note: [M]/[PEGDA] = 9/1 (molar ratio); TPO = 1 wt%.

suggesting the formation of larger domains (>100 nm), consistent with the lack of SAXS signals (Fig. 4d). This effect was even more pronounced in the control formulation (PCL_{3k}-OH, Fig. 6e), where the objects were fully opaque, indicating significant macrophase separation. These results demonstrate that formulations based on PCL-BTPAs enable high-resolution 3D printing of complex objects, with tunable nanodomain sizes (Fig. 6b-d).

Due to the importance of porous materials for a wide range of applications,^{48,49} the final part of our study focused on creating nanoporous materials by leveraging the degradability of PCL through controlled solvent-based etching, allowing for tunable pore size.^{23,34-37} Initial etching experiments were performed on 3D-printed objects fabricated from the PCL_{3k}-BTPA formulation. After printing, the objects were immersed in a 0.5 M NaOH aqueous solution at 60 °C for one and two days. SEM images were acquired before etching, after one day, and after

two days of etching (Fig. 7a). Using AA as the comonomer resulted in no observable pores after etching, likely due to swelling and collapse of the poly(acrylic acid) (PAA) domains upon freeze-drying. To address this limitation, AA was replaced with a hydrophobic monomer (IBoA) resulting in the formation of water-insoluble domains, preventing pore collapse. Additionally, the PCL M_n was increased from 3000 to 11 000 g mol⁻¹ to promote the formation of larger pores. SEM images revealed that samples before etching exhibited no visible pores, but pores emerged after one day of etching (Fig. 7a). After two days, pore coalescence was observed, indicating structural evolution over time. An identical procedure was followed for 3D-printed objects made from PCL_{11k}-acrylate (Fig. 7b), revealing pore formation with broader dispersity, as expected due to the uncontrolled nature of the polymerization mechanism.

To verify the successful etching of PCL, weight loss measurements were conducted (Table S4[†]), complemented by



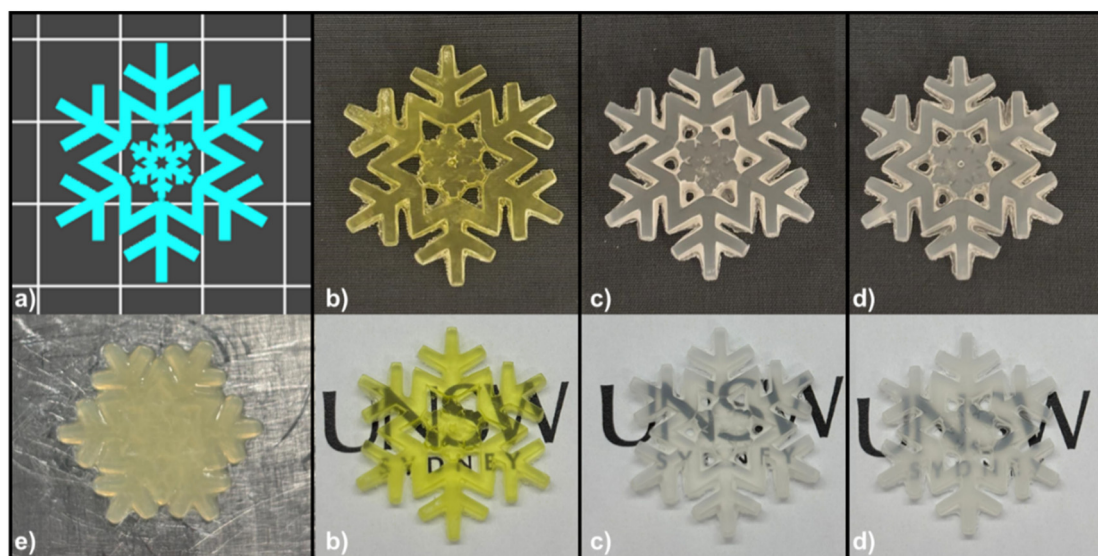


Fig. 6 3D Printing of complex snowflake structures. (a) Digital model. (b–e) Photographs of 3D-printed snowflakes using resins formulated with (b) PCL_{3k}-BTPA, (c) PCL_{3k}-acrylate, (d) PCL_{3k}-methacrylate, and (e) PCL_{3k}-OH (control). [AA]/[PEGDA] = 9/1 (molar ratio); PCL-R = 19 wt%; toluene = 10 wt%; and TPO = 1 wt%. Layer thickness: 50 μ m. Cure times: 30 s (PCL_{3k}-BTPA and PCL_{3k}-OH) and 20 s (PCL_{3k}-acrylate and PCL_{3k}-methacrylate).

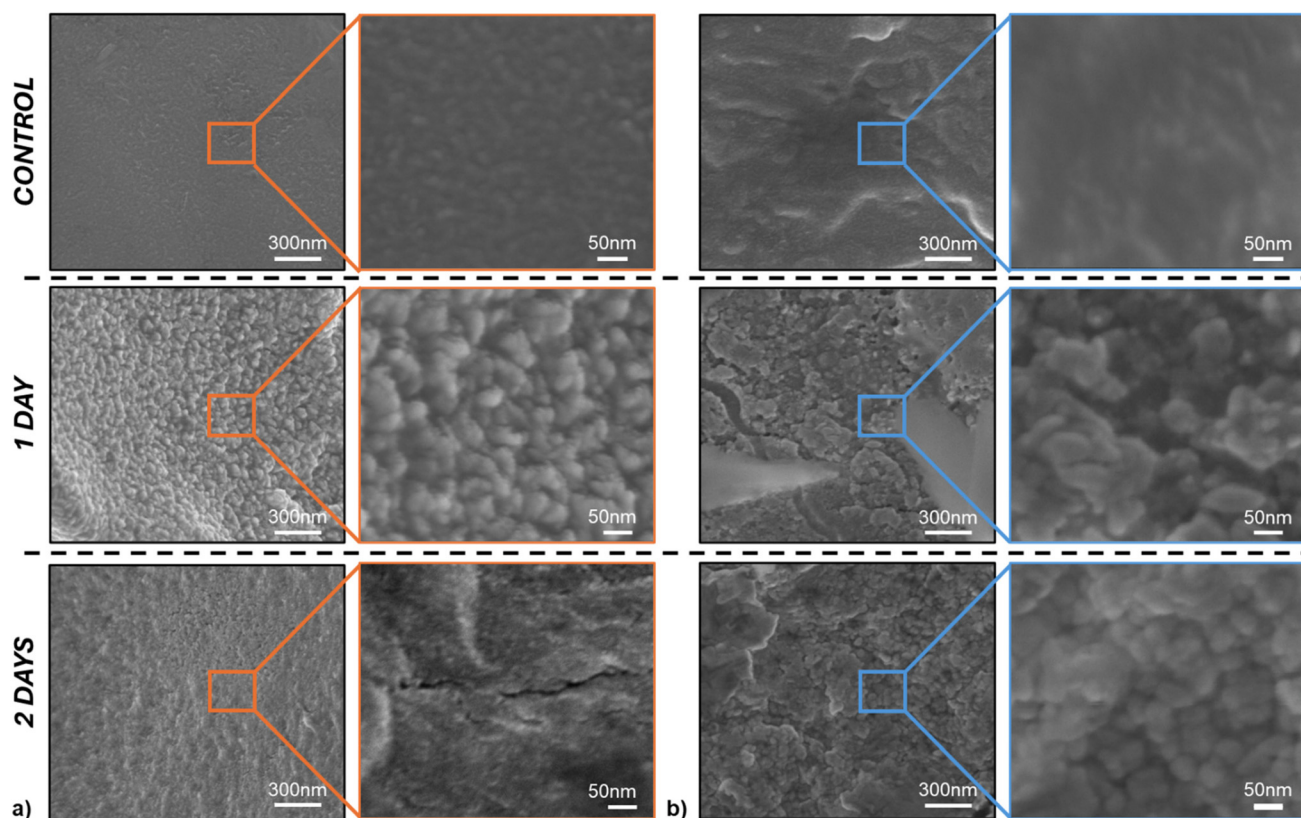


Fig. 7 SEM images of freeze-dried 3D-printed object prepared with (a) PCL_{11k}-BTPA and (b) PCL_{11k}-acrylate in the presence of IBoA, PEGDA and 10 wt% toluene, following selective etching of the PCL domains. Images show the control sample before etching (top), after 1 day of treatment (middle), and after 2 days of treatment (bottom) in a 0.5 M NaOH aqueous solution at 60 $^{\circ}$ C.



ATR-FTIR analysis. The weight loss data showed different kinetics depending on the formulation. PCL_{11k}-BTPA samples exhibited a 21.7 wt% loss after one day of etching, which corresponds well with the theoretical PCL content targeted for removal (e.g., ~29 wt% in that formulation). In contrast, PCL_{11k}-acrylate samples required two days to achieve a comparable weight loss (25 wt%), suggesting slower degradation, potentially due to network differences influencing the etching process. Additionally, ATR-FTIR analysis (Fig. S34†) supported successful PCL degradation. Spectra acquired after etching showed a marked decrease in the ester C=O peak (~1730 cm⁻¹) and the appearance of features indicative of hydrolysis products: specifically, the broad O-H stretching of carboxylic acids (~3300–2500 cm⁻¹) and the carboxylate anion (COO⁻) absorption near 1570 cm⁻¹.

Conclusion

In conclusion, this study successfully developed 3D-printable resins containing both RAFT-terminated and vinyl-terminated PCLs for the fabrication of nanostructured materials *via* PIMS. A systematic comparison of these resins revealed distinct nanodomain structures, significantly influencing the materials' optical properties. Notably, the use of PCL_{3k}-acrylate enabled the fabrication of high-resolution, colorless nanostructured materials, overcoming the limitations of traditional RAFT systems, which often yield yellowish products. Furthermore, the addition of toluene (up to 40 wt%) improved resin stability, facilitating extended print durations and the creation of complex 3D-printed objects. Finally, selective etching of the PCL component under basic conditions successfully produced nanoporous materials, demonstrating the versatility of this approach. These findings open new avenues for advanced applications in diverse fields such as filtration, catalysis, and tissue engineering.

Data availability

The data supporting this article have been included as part of the ESI.†

Conflicts of interest

The authors declare no conflict of interest.

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