Programmable Porous Polymers via Direct Bubble Writing with Surfactant-Free Inks

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ABSTRACT: Fabrication of macroporous polymers with functionally graded architecture or chemistry bears transformative potential in acoustic damping, energy storage materials, flexible electronics, and filtration but is hardly reachable with current processes. Here, we introduce thiol–ene chemistries in direct bubble writing, a recent technique for additive manufacturing of foams with locally controlled cell size, density, and macroscopic shape. Surfactant-free and solvent-free graded three-dimensional (3D) foams without drying-induced shrinkage were fabricated by direct bubble writing at an unparalleled ink viscosity of 410 cp (40 times higher than previous formulations). Functionalities including shape memory, high glass transition temperatures (>25 °C), and chemical gradients were demonstrated. These results extend direct bubble writing from aqueous inks to nonaqueous formulations at high liquid flow rates (3 mL min⁻¹). Altogether, direct bubble writing with thiol–ene inks promises rapid one-step fabrication of functional materials with locally controlled gradients in the chemical, mechanical, and architectural domains.

KEYWORDS: polymer foams, 3D printing, nonaqueous foam, thiol–ene, functionally graded materials

INTRODUCTION

Solid polymeric foams constitute a widely applied material class with applications in cushioning (shoes, sofa, bed),¹ oil sequestration,² energy storage,³ robotics,⁴ pressure sensing,⁵,⁶ and scaffolds for artificial tissues.⁷⁻⁹ The widespread use of foams lies in their exceptional performance as compared to bulk materials, such as excellent weight/stiffness ratios, low heat transfer, permeability, and load-bearing potential by a single construct.¹⁰ Compared to natural foams, such as bone and bamboo, synthetic foams possess multiple shortcomings including poor control of the three-dimensional (3D) pore arrangement, outer shape complexity, and, especially, gradients in pore size, density, stiffness, and other relevant properties that would enable the integration of multiple functions by a single foam construct.¹⁰⁻¹² Conventional, synthetic foams rely on the introduction of air by mechanical action or chemical addition, resulting in a broad distribution of the cell size, that is, polydisperse foams. In recent years, monodisperse foams have been typically made in a two-step process, by first depositing monodisperse solids/emulsions in a matrix, and then removing the deposited material.¹³⁻¹⁷ While these approaches provide monodisperse foams with either open- or closed-cell architectures,⁷ the requirement of surfactant, limited throughput (typically a few mL h⁻¹), use of multiple precursor liquids, and multistep processes inhibit their immediate industrial viability.¹⁵,¹⁸ Additionally, integrating complex 3D geometries¹⁹⁻²⁴ and a range of porosities within a single construct is still challenging.²⁵ A simple, robust, and scalable technique to produce surfactant-free foams with defined cellular size/structure and 3D cell arrangement with a broad access to aqueous/nonaqueous monomers and minimal shrinkage/postcuring would therefore represent a major step forward.²⁶

Direct bubble writing is a new printing technique that relies on the rapid solidification of bubbles upon impact on a substrate to ensure high fidelity printing.⁵ To print bubbles, pressurized gas is injected through the core of a core–shell nozzle with a coflowing outer shell fluid (aqueous or organic inks) to produce gas-filled bubbles with tunable size and porosities.

In our previous work, we demonstrated that acrylate-based direct bubble writing possessed gas-type dependent cellular structure (i.e., nitrogen = closed-cell or air = open-cell), locally tunable foam properties (density/modulus) via control over the bubble size/distribution, and the ability to print 3D lattices, shells, and out-of-plane pillars.⁷ Key benefits of this process include high throughput (10 mL min⁻¹ liquid flow rate), on...
the-fly tunable cell size/architecture, simple setup, and minimal clogging. However, direct bubble writing suffers from substantial shrinkage (28% in each dimension), use of surfactant for bubble stabilization, a limited monomer selection, and a prolonged drying step to remove the solvent (water, forming 59% of the initial weight). To overcome these issues, we advance direct bubble writing to a solvent-free thiol–ene platform. Thiol–ene photopolymerization is insensitive to oxygen, does not need a surfactant, has negligible shrinkage, requires minimal postcure processing, allows for rapid postpolymerization modification, and enables access to higher glass transition temperature materials ($T_g > 23^\circ C$). Such a robust platform provides tunable chemical/mechanical properties with simple “click” chemistries, further decreasing the barrier to applying the direct bubble writing platform.

**METHODS AND MATERIALS**

**Materials.** All reagents were obtained commercially and used without further purification unless otherwise specified. Triallylisocyanurate, pyrogallol, pentaerythritol tetrameracaptopropionate (PETMP), dipentaerythritol hexakis(3-mercaptopropionate) (DiPETMP), and 2,2-dimethoxy-2-phenylacetophenone (DMPA) were purchased from Sigma-Aldrich. Darocure 1173 was donated by iGM Resins.

**Printing Ink.** Pyrogallol (0.1 g, 0.79 mmol) serves as a radical inhibitor to stabilize the ink formulations and is first dissolved into triallylisocyanurate (40 g, 0.16 mol), followed by pentaerythritol tetrameracaptopropionate (PETMP, 29.4 g, 0.06 mol), dipentaerythritol hexakis(3-mercaptopropionate) (DiPETMP, 31.4 g, 0.04 mol), and 4.2 g of the photoinitiator Darocure 1173 as shown in Figure 1a. Inks containing excess thiol were prepared similarly but with a 1.2:1 thiol-to-alkene ratio by increasing amounts of DiPETMP (41.9 g, 0.54 mol) and PETMP (39.2 g, 0.08 mol).

**3D Printing Setup.** Printing resins are loaded into 30 mL of HSW soft-jet luer-lock syringes and mounted into a NE-8000 New Era Pump Systems syringe pump. The loaded syringe is connected to the shell nozzle of a custom core–shell nozzle (for details, see ref 6) by a 1 mm diameter, 48 cm long tubing. Pressurized air connects to the core of this nozzle, resulting in the direct ejection of resin bubbles into air as depicted in Figure 1b (photographs of the setup are provided in Figure S1). The bubble train is deposited onto the print surface, where the resin is photopolymerized by ultraviolet (UV) light. The alignment of the UV light and the bubble deposition location is ensured by mounting the print nozzle and two fiber-optic light guides to a custom machined aluminum bracket, which is bolted to a LuZBot TAZ 6 3D printer. Print paths for each geometry are written as G-code scripts and loaded onto the 3D printer via an SD card. Air pressure is controlled via a Matlab-controlled Alicat Scientific electronic pressure controller, and resin flow is controlled via the syringe pump feed rate. UV light intensity for each light guide is controlled independently via two OmniCure light sources. One light source (OmniCure S2000) cures a stream of air-infused resin 7.5 mm from the nozzle tip at an angle of 18° from horizontal. This precuring does not fully cure the ink and is designated as a “stream cure”. The other light source (OmniCure S1000) cures the stream 78 mm from the nozzle tip at 70° from horizontal, and it is designated as “surface cure”. Printing parameters studied include air pressure (0–40 kPa), print speed (30–100 m s$^{-1}$), resin flow rate (1.5–4.0 mL min$^{-1}$), and UV-cure intensity (55.4–621 mW cm$^{-2}$). Unless otherwise indicated, each sample was printed with a constant resin flow rate of 3 mL min$^{-1}$ and air pressure of 23 kPa with a surface cure intensity of 30.6 mW cm$^{-2}$ and a stream cure intensity of 301 mW cm$^{-2}$. The printing speed for each sample was varied using G-code commands to print 30 mm long sections for each printing speed. After printing, the samples were postcured with unfiltered UV light before being removed from the printing surface. Printed parts were exposed for 30 s by unfiltered UV light within 5 min after printing to ensure that the samples were thoroughly cured.

**UV Intensity Study.** The influence of UV-cure intensity on resin stream width and print quality was studied by independently varying stream and surface UV-cure intensity. Two experimental conditions were used: the first varying stream cure intensity with a constant surface cure intensity of 30.6 mW cm$^{-2}$ and the second varied stream cure intensity with a constant surface cure intensity of 106 mW cm$^{-2}$. The samples were then studied using a microscope to determine the print quality at each cure intensity.

**Mechanical Testing.** The influence of air pressure on foam mechanical properties was studied via uniaxial compression testing using a MTS 810 hydraulic test frame. Printing paths and speeds for each sample were controlled using G-code commands to print 50 mm long sections for each printing speed. After printing, the samples were postcured with unfiltered UV light before being removed from the printing surface. The samples were compressed to a strain of 50% at a constant cross-head speed of 50 mm min$^{-1}$ using a 100 kN load cell. Fewer than four samples were tested at each air pressure.

**Rheology.** Ink formulations were evaluated with an ARES-G2 rheometer from TA Instruments (New Castle, DE) equipped with a UV light guide accessory. Following a 14 s equilibration period, the samples were irradiated by UV light from an unfiltered OmniCure S2000 UV light source via a fiber optic cable for 30 ms. Unfiltered UV light intensity was calibrated at the sample location using an external fiber-optic light guide accessory.
radiometer. Photoreology experiments were conducted by loading samples between an aluminum (bottom) and quartz (top) 20 mm diameter parallel plates with a sample thickness of 0.5 mm. Experiments were conducted using automatic strain adjustment at 10 Hz in “fast sampling” mode to resolve rheological behavior at fast cure speeds and to prevent over torque of the instrument. The viscosity was determined as a function of the shear rate with a stainless steel 2.5 mm cone and plate geometry (0.1 rad) at 25 °C. The samples were presheared at 10 s⁻¹ prior to steady shear experiments to ensure homogenous samples before monitoring steady-state shear viscosity.

**Postpolymerization Modification.** Printed porous filaments prepared from inks containing 20 mol % excess thiol were postmodiﬁed via thiol–ene and thiol-Michael reactions. For thiol–ene modiﬁcation, the thiol-functional ﬁlaments were submerged in a tetrahydrofuran (THF) solution containing ﬂuorescein o-methacrylate (0.025 g, 0.06 mmol) and DMPA (0.01 g, 0.04 mmol) and irradiated with 4 mW cm⁻² UV light for 30 min to ligate the ﬂuorescein dye to the excess thiol surface. For thiol-Michael modiﬁcation, thiol-functional ﬁlaments were submerged in either an aqueous solution of Texas Red C2 maleimide (5 mg, 0.007 mmol) or a THF solution of 1-[4-(trifluoromethyl)phenyl]-1H-pyrrole-2,5-dione (CF₃-MA, 0.025 g, 0.1 mmol) containing triethylamine for 12 h.

**Characterization.** X-ray photoelectron spectroscopy was performed using a Thermo Fisher ESCALAB Xi+ spectrometer equipped with a monochromatic Al X-ray source (1486.6 eV). Measurements were performed using the standard magnetic lens mode and charge compensation. The base pressure in the analysis chamber during spectral acquisition was at 3 × 10⁻⁷ mbar. Spectra were collected at a takeoff angle of 90° from the plane of the surface. The pass energy of the analyzer was set at 150 eV for survey scans with an energy resolution of 1.0 eV; the total acquisition time was 220 s. Binding energies were calibrated with respect to C 1s at 284.8 eV. All spectra were recorded using the Thermo Scientiﬁc Avantage software; data ﬁles were translated to VGD format and processed using the Thermo Avantage package v5.9904. Polymerization kinetics were measured by real-time Fourier transform infrared (FTIR) spectroscopy using a Nicolet 8700 FTIR spectrometer with a KBr beam splitter and MCT/A detector. Each sample was exposed to a UV light with an intensity of 100 mW cm⁻² using an Omnicure Series 1000 light source. Series scans were recorded at approximately 2 scan s⁻¹ with a resolution of 4 cm⁻¹. Thiol conversion was monitored by integrating the peak at 2557 cm⁻¹ over time.

**RESULTS AND DISCUSSION**

To use direct bubble writing without both solvent and surfactant, a library of thiol and alkene monomers were probed for their ability to foam through a core–shell nozzle. While typical monomers have viscosities less than 10 cP, we found that higher viscosity monomers (90–2500 cP) still provide highly controlled bubble trains. The final ink formulation had a viscosity of 410 cP and is comprised of hexa- and tetra-functional thiol monomers along with a trifunctional alkene and a photoinitiator (see Figure 1a). To study the influence of formulation changes to ink curing, different photoinitiator concentrations were evaluated in 30 ms pulse-UV rheometer experiments (Figure S2a). The rate at which the ink’s complex viscosity builds after UV exposure monotonically increased as a function of the Darocur 1173 concentration, which was varied from 0 to 4%. The 4% Darocur 1173 concentration reached a complex viscosity greater than 20 000 cps in less than 0.4 s. Additionally, photoinitiators Darocur 1173 and SpeedCure TPO-L (2,4,6-trimethylbenzoyldi-phenylphosphinate) were evaluated at different cure intensities (2, 30, 60 mW, with a 30 ms pulse, at 4 wt %) (Figure S2b). Clearly, the light intensity had a greater effect than the photoinitiator type, highlighting the importance of light exposure intensity and location. Darocur 1173 was chosen for the ﬁnal formulation since this photoinitiator consistently yielded higher viscosity materials compared to the TPO-L.

To probe the bubble ejection regime from the core–shell nozzle for this highly viscous ink, we systematically varied the flow rate of the liquid resin mixture (Ql) and the air pressure (Pair). Figure 1b depicts the experimental setup of the nozzle conﬁguration to study bubble ejection and simultaneously print 3D structures. As shown in Figure 1c–h, at all ﬂow rates tested (1.5 ≤ Ql ≤ 4.0 mL min⁻¹), ﬁve distinct ejection mechanisms were observed. At low Pair, pure-liquid resin jetting/dripping occurred (Figure 1c). Here, the gas is unable to overcome the threshold pressure required to enter the liquid as previously described. As Pair increases, the length of the jet increased until a critical point where small bubbles become continuously entrained within a jet of resin to form a connected bubble stream (Figure 1d). A further increase in Pair leads to a transition away from ﬁlaments and to the production of disconnected monodisperse bubbles (droplets with a liquid shell and a gas core; Figure 1e). As Pair is again further increased, multibubble ejections (Figure 1f) and
eventually random spraying occurs (Figure 1g). To afford the highest control and minimize defects during printing, we selected a flow rate $Q_\text{f}$ of 3 mL min$^{-1}$ for all experiments. Higher flow rates are also well-controlled but result in too short print times for our syringe-fed setup.

Multiple configurations of the UV light guides were evaluated (i.e., a single stream/surface light or double stream/surface lights). Interestingly, if both fiber-optic lines are positioned in the stream cure position, the UV-curing of this connected bubble stream leads to the production of porous fibers, which at present cannot be printed in a controlled fashion. Alternatively, when just a single light source was used (stream or surface), the ink was allowed to flow too much, leading to poor retention of the porous structures. The combination of a stream light with a surface light allowed the ink to flow enough for higher-fidelity layer-by-layer printed structures. In Figure 2, we determine the optimal curing conditions of the bubbles by varying the intensity of the stream curing from 55 to 621 mW cm$^{-2}$ at two different surface cure intensities (30 and 106 mW cm$^{-2}$). At low stream UV intensities (55–174 mW cm$^{-2}$), the ink that impacts the substrate is able to flow, leading to regions of high resin density and regions with bubbles. This result is readily explained by the reduction in curing, allowing time for the resin to separate from the bubbles prior to polymerization as seen in the 55.4 mW cm$^{-2}$ printing condition. At medium cure intensities (244–408 mW cm$^{-2}$), the bubbles are distributed over the entire filament and a more uniform structure was observed (similar thickness throughout the printed line with uniform bubble placement). At 483–621 mW cm$^{-2}$ and at higher surface cure intensity (106 mW cm$^{-2}$), a tortuous printed line was observed. We assume that the stream was cured immediately upon impact with the surface, such that capillary forces were prevented from relaxing the internal surfaces (the bubble air–resin interface) as well as the external surfaces (smoothening the edges of the filament). Eventually, 30.6 mW cm$^{-2}$ surface cure with 301 mW cm$^{-2}$ stream cure was selected for all experiments moving forward.

To print objects with multiple layers, the influence of print speed on the resulting line width (Figure 3a) and the layer thickness as a function of the number of layers (Figure 3b) were systematically studied. To assess the line width, filaments were formed by laterally moving the printhead in the $x$-$y$ direction at programmed print speeds. For printing speeds of $V > 20$ mm s$^{-1}$, continuous filaments are formed with decreasing filament width as the velocity of the printhead is increased (Figure 3a). As the ejected filaments are cured in situ, they can be stacked to produce 3D architectures. As Figure 3b shows, the total thickness ($L_T$) increases linearly (1.4 mm layer$^{-1}$) with the number of applied layers ($L_n$), indicating that the local impact and solidification process are robust with respect to the height of the construct.

With the ability to print consistently, we examined the influence of the gas pressure on the resulting cell size, foam density, and Young’s modulus. As the gas pressure ($P_{\text{air}}$) increases from 21.7 to 22.6 kPa, the average bubble diameter (DB) increases from 107 ± 8 to 173 ± 15 µm, respectively (Figure 4a). The coefficient of variance ($CV = \sigma/m$ was determined, where $\sigma$ is the standard deviation and $m$ is the mean cell size as detailed in Figure S3. In the monodisperse ejection regime, a tight distribution of 4.9% $< CV < 6\%$ was achieved. This is an improvement compared even to the recent microfluidic techniques for which $CV \approx 20\%$ was reported as the state of the art for solid foams.\textsuperscript{36} Figure 4b shows that increasing the pressure also decreases the density of the foams from 845 ± 25 kg m$^{-3}$ at 21.7 kPa to 501 ± 26 kg m$^{-3}$ at 22.6 kPa. It is well-known that the density of foams controls their stiffness.\textsuperscript{37} Figure 4c shows Young’s modulus $Y$ as a function of the density, revealing tunability over more than an order of magnitude.
magnitude (from 0.32 ± 0.17 to 5.5 ± 0.7 MPa). The dashed line in Figure 4c shows the exponent $\alpha$ of the relationship $Y = \rho^\alpha$, where $\rho$ is the relative density. The observed value $\alpha = 3.2$ exceeds the literature value for both open-cell foams ($\alpha = 2$) and closed-cell foams ($\alpha = 3/2$). Although the theory was derived for the low-$\rho$ limit, this high value suggests that part of the foam mass hardly contributes to the stiffness (it may be contained in the nodes between the struts). Therefore, Young’s modulus is highly tunable over the achieved porosity fraction of ~50%, enabling one-step and single-material production constructs with locally tailored properties. As an example, we printed a basic multilayer construct in which we modulated $P_{\text{air}}$ during printing, resulting in distinct regions of small cells ($D_\text{h} = 90 \pm 12 \mu m$), nonporous material, and larger cells ($D_\text{h} = 122 \pm 23 \mu m$) (Figure 5). Here, the porous sections were readily bent, while the nonfoamed center section exhibited notably higher stiffness. This concept could be readily leveraged for 3D printing of functionally graded wearables such as helmets, where cushioning and hard protection must be integrated in a single piece.

In addition to functional gradients derived from the foam’s morphology, thiol–ene inks enable the fabrication of porous materials with independently controlled chemical functionalization. As a salient consequence of the step-growth thiol–ene polymerization process, thiol–ene materials are inherently functional with thiol and alkene groups. Off-stoichiometric inks can be employed to produce materials that are predominately thiol or alkene functional materials where the functional group can be employed to produce materials that are predominately polymeric. 38 Here, we demonstrate this ability by changing the monomer ratio of thiol to alkene within the ink to print foams comprising excess thiols, which can undergo simple postpolymerization reactions (Figure 6a). As an example, two foam filaments were printed, one from an excess thiol ink (1.2:1 SH/alkene) and another from a stoichiometric ink (1:1 SH/alkene). As indicated in Figure S4, thiol conversion plateaus at approximately 80% upon photopolymerization of inks containing 20 mol % excess thiol-functional groups, whereas stoichiometric inks achieve >95% thiol conversion. The thiol-functional and control filaments were submerged in a THF solution containing fluorescein-ortho-methacrylate and DMPA and irradiated with 4 mW cm$^{-2}$ UV light to ligate the fluorescein dye to the excess thiol surface. After multiple washes and gentle sonication, fluorescent microscopy revealed the dye attached efficiently to the filament containing excess thiol filament, while only faint fluorescence was observed for the control filament (Figure 6b). Similarly, thiol-functionalized and control filaments were submerged in an aqueous solution of Texas Red C2 Maleimide and trimethylenamine to ligate Texas Red C2 to the foam’s surface via thiol-Michael addition; again, only the filament fabricated with excess thiol in the ink showed evidence of efficient ligation (Figure 6c). Additional surface characterization of the off-stoichiometric resins via X-ray photoelectron spectroscopy (XPS) (before and after postfunctionalization with CF$_3$-MA) is provided in Figure S4. This demonstration confirms that simple postproduction thiol–ene reactions can successfully add new functionality to the foams.

As a final example, we demonstrate shape-memory applications at an easily accessible temperature range (Figure 7a) enabled by the higher $T_g$ ($40$ °C) of the cured thiol–ene inks as compared to our previously used acrylate-based inks. 6 As shown in Figure 7b, a three-dimensional flower that deploys in water was printed on a flat surface at room temperature. Covalent cross-links within the network define the permanent shape of a flower. Subsequently, the flower was heated above $T_g$ to 60 °C to make the material conformable. The flower was folded into a temporary shape and then cooled below its $T_g$ to lock the temporary shape under an elastic strain. Upon contact with warm water ($T_{\text{water}} > T_g$), the flower unfolded into its original printed shape within 10 s, as shown by time-stamped images in Figure S5, driven by the entropic recovery of the elastic strain within the polymer network. The mechanism for shape memory in cross-linked polymer networks using $T_g$ as the switch is well-documented in literature. 39–42 with several examples focusing on thiol–ene materials. 33–45 This example highlights the one-step fabrication of shape-programmable memory foams, enabling complex patterns that save space and are deployable to predefined shapes upon application of heat. 36

## CONCLUSIONS

The combination of thiol–ene-based inks and direct bubble writing yielded a rapid one-step fabrication of foams with narrow cell size distributions in the absence of both surfactant and solvent. The high viscosity of the used thiol and alkene monomers yielded a viable strategy toward a surfactant-free fabrication of porous solids. In contrast, direct bubble writing with low-viscosity bubbles ($\eta \approx 10$ mPa·s) required a surfactant to prevent their rupture prior to impacting the surface. This newfound processing window for inks with a sufficiently high viscosity to omit surfactants but with sufficiently low viscosity for bubble ejection from core–shell nozzles is highly relevant, especially since predicting and preventing the rupture of bubbles are still challenging. 37 Furthermore, the lack of solvent, coupled with gelation at high thiol–ene monomer conversion, 27–29 ensured that no significant shrinkage of the printed parts was observed. The use of high $T_g$ monomer pairs in direct bubble writing was exploited in a simple shape-memory application. Additionally, altering the ratio of thiol–ene monomers allowed for the production of excess thiol foams to undergo postpolymerization modification via two different thiol–ene reactions. These examples highlight direct bubble writing as a new route toward

![Figure 5. Printed construct with two porous regions connected by a nonporous region. The graph depicts the bubble diameter distribution for this continuously printed multilayer porous solid. Small bubbles: $P_{\text{air}} = 21.2$ KPa; no bubbles: $P_{\text{air}} = 18$ KPa; large bubbles: $P_{\text{air}} = 22.2$ KPa.](https://dx.doi.org/10.1021/acsami.0c07945)
functionally graded materials with independently tunable architectural and chemical gradients.

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsami.0c07945.

The Supporting Information includes: Figure S1. Overview of the setup; Figure 2. Time-resolved rheological properties of the resin and curing dynamics; Figure S3. Automated cell size analysis; Figure S4. Unfolding of memory foam (PDF)

Figure 6. Off-stoichiometry resin and postfunctionalization. (a) Filaments were printed with either a thiol-to-ene ratio of 1.2:1 (yellow spheres) or a 1:1 control. After printing, the excess functional group is reacted with either (i) fluorescein α-methacrylate, THF, DMPA, and 4 mW cm⁻² UV light or (ii) Texas Red C₂ Maleimide, water, and triethylamine. These reactions respectively yield materials with a green or red fluorescent functional group that is covalently bonded to the resin. (b) Optical bright-field micrograph of fluorescein labeled (left), unlabeled (middle), and Texas Red labeled (right). (c) Fluorescein labeled porous solid under 395–455 nm excitation and 480 nm LP filter. (d) Texas Red C₂ Maleimide labeled porous solid under 540–580 nm excitation with a 610 nm LP filter. Scale bar = 4 mm.

Figure 7. Thermal and shape-memory properties of a 3D printed porous solid. (a) Differential scanning calorimetry of cured thiol–ene foam. (b) Shape memory cycle for a 3D printed flower. The diameter of the flower is 10 cm.

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Notes

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