



Cite this: *Soft Matter*, 2017,
13, 921

Received 28th September 2016,
Accepted 23rd December 2016

DOI: 10.1039/c6sm02208a

www.rsc.org/softmatter

3D printing of self-assembling thermoresponsive nanoemulsions into hierarchical mesostructured hydrogels

Lilian C. Hsiao, † Abu Zayed Md Badruddoza, Li-Chiun Cheng and Patrick S. Doyle*

Spinodal decomposition and phase transitions have emerged as viable methods to generate a variety of bicontinuous materials. Here, we show that when arrested phase separation is coupled to the time scales involved in three-dimensional (3D) printing processes, hydrogels with multiple length scales spanning nanometers to millimeters can be printed with high fidelity. We use an oil-in-water nanoemulsion-based ink with rheological and photo-reactive properties that satisfy the requirements of stereolithographic 3D printing. This ink is thermoresponsive and consists of poly(dimethyl siloxane) droplets suspended in an aqueous phase containing the surfactant sodium dodecyl sulfate and the cross-linker poly(ethylene glycol) dimethacrylate. Control of the hydrogel microstructure can be achieved in the printing process due to the rapid structural recovery of the nanoemulsions after large strain-rate yielding, as well as the shear thinning behavior that allows the ink to conform to the build platform of the printer. Wiper operations are used to ensure even spreading of the yield stress ink on the optical window between successive print steps. Post-processing of the printed samples is used to generate mesoporous hydrogels that serve as size-selective membranes. Our work demonstrates that nanoemulsions, which belong to a class of solution-based materials with flexible functionalities, can be printed into prototypes with complex shapes using a commercially available 3D printer with a few modifications.

1. Introduction

Self-assembly is a popular method to engineer hierarchical microstructure in many types of soft materials including polymeric hydrogels.¹ Hydrogels can be formed through chemical or physical crosslinking methods, and are particularly attractive because they provide a hydrated environment with tunable diffusion profiles

and mechanical properties.^{2–7} Existing strategies to prepare structured hydrogels include the use of freeze drying^{8,9} and sacrificial templating.^{10–12} These methods only provide a coarse level of control over the resultant porous microstructure. Spinodal decomposition and phase transitions *in situ* have recently emerged as viable methods to generate a variety of bicontinuous materials including bijels,¹³ demixing polymers,¹⁴ and nematic liquid crystals.¹⁵ These multiphase systems display a rich variety of microstructure due to the non-equilibrium thermodynamic instability of the constituent building blocks. Temperature responsiveness is inherent to many of these systems and represents a simple and effective tuning parameter to access microstructure in the micron length scale. Responsive colloids that form gel networks represent a particularly effective method of self-assembling such types of structure. Their characteristic domain size depends on the thermodynamics and kinetics of the highly non-equilibrium process.^{16,17} Large, heterogeneous voids with strands that range from hundreds of nanometers to tens of microns are commonly observed in colloidal gels.¹⁸

Here, we show that when the dynamics of gelation are coupled to the time scales involved in three-dimensional (3D) printing processes, hydrogel materials with multiple length scales spanning nanometers to millimeters can be printed rapidly. 3D printing has gained popularity in recent years as an additive manufacturing platform that offers unparalleled flexibility in creating structures with arbitrary shapes and material properties.^{19,20} Compared to ink jetting and extrusion-based methods, stereolithographic (SLA) printing is a layer-by-layer prototyping technique that shows excellent compatibility with liquid polymeric materials. A traditional SLA 3D printer operates by photopolymerization at the surface of a resin reservoir, in which the constructed model is progressively built by moving downwards into the ink bath.²¹ Modern SLA 3D printers utilize an inverse setup and a layer-by-layer process to improve the printing resolution of each printed layer. First, a build platform is immersed into the resin tank containing the precursor ink. The microstructure is then photopolymerized using a 405 nm laser and actuated micromirrors, which patterns the desired motif

Department of Chemical Engineering, Massachusetts Institute of Technology, Cambridge, MA 02139, USA. E-mail: pdoyle@mit.edu

† Present address: Department of Chemical and Biomolecular Engineering, North Carolina State University, Raleigh, NC 27695, USA.



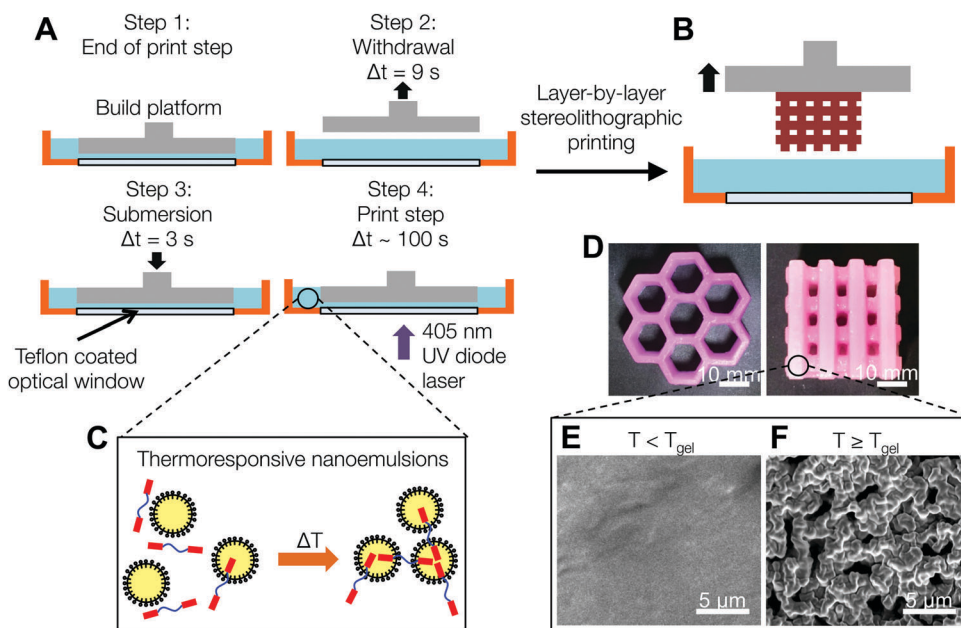


Fig. 1 3D printing of self-assembled thermoresponsive nanoemulsion inks. (A and B) Schematic of commercially available stereolithographic (SLA) printer with a custom-modified Teflon window to enhance oxygen permeability. Layer-by-layer SLA printing consists of four steps in which the desired motif is photocrosslinked, followed by high strain-rate withdrawal and submersion steps. The process repeats at a z-step size of 100 μm . (C) Nanoemulsions (yellow) stabilized by surfactants in the resin tank are heated to induce self-assembly through interdroplet bridging of PEGDMA gelators (red are dimethacrylate groups). (D) 3D honeycomb and woodpile structured hydrogels formed by using the nanoemulsion inks. (E and F) The internal morphology of the 3D printed scaffold is either homogeneous at $T < T_{\text{gel}}$ or interconnected when $T \geq T_{\text{gel}}$.

through an optically transparent window. Finally, the platform is withdrawn and re-immersed again to print the next layer, with automated wiper actions between each print step to remove remnant debris on the optical window (Fig. 1). Oxygen inhibition of the free radical polymerization is an important parameter to monitor as it controls the overall printing speed.²² An oxygen permeable poly(dimethyl siloxane) (PDMS) layer is typically placed above the optical window to ensure a thin lubrication layer between the printed model and the resin tank, which prevents the model from adhering permanently to the window.^{19,23,24}

In order to leverage SLA printing for the fabrication of self-assembled hydrogels, the rheological properties of the ink should be compatible with the movement of the build platform. High strain rate deformations are applied to the ink as the platform is repeatedly immersed and retracted over the duration of the printing process. Many viscoelastic self-assembled materials can exhibit fluid instabilities²⁵ and yield irreversibly under these types of large stresses.²⁶ Thus, in order to print 3D mesostructured hydrogels with high speed and fidelity, the ink must be able to recover its original microstructure rapidly after yielding. In addition, the material should exhibit liquid-like response²⁷ during platform movement such that it conforms to the base of the printed motif. Here, we demonstrate that thermoresponsive nanoemulsions, which belong to a class of multiphase materials, can be printed into hierarchical structures using a modern 3D SLA printer when the rheology of the complex fluid during the printing process is accounted for.

2. Experimental section

Materials

We present a oil-in-water nanoemulsion-based ink with rheological and photoreactive properties that satisfy the requirements of SLA 3D printing. Chemicals are purchased from Sigma-Aldrich and used without further purification unless noted. The nanoemulsions consist of a continuous phase with 33 vol% poly(ethylene glycol) dimethacrylate (PEGDMA, molecular weight $M_n = 750 \text{ g mol}^{-1}$) and 200 mM sodium dodecyl sulfate (SDS) dissolved in deionized (DI) water, as well as an oil phase consisting of poly(dimethyl siloxane) (PDMS) droplets (viscosity, $\eta = 5 \text{ cP}$) suspended at a volume fraction of $\phi = 0.25$. The hydrophobic fluorescent dye Nile red (excitation wavelength = 550 nm, emission wavelength = 626 nm) is dissolved in the PDMS at a concentration of 0.05 mg mL⁻¹ to enable confocal microscopy imaging of the nanoemulsions.

To synthesize the nanoemulsions, a crude emulsion is first prepared by adding the oil phase into the continuous phase under stirring. The crude emulsions are then passed through a high pressure homogenizer (Emulsiflex-C3, AVESTIN) at a pressure of 15 000 psi for 20 passes. A heat exchanger is used to chill the sample to 5 °C between each pass. After homogenization, the nanoemulsions are placed in a centrifuge at 5000 rpm to remove any large impurities and the supernatant is stored at 4 °C until further use. The final diameter of the droplets is measured using dynamic light scattering (90Plus PALS, Brookhaven Instruments) after dilution to $\phi = 0.002$ with a solvent of 33 vol% PEGDA in DI water ($\eta = 5.8 \text{ cP}$). The diameter of the nanoemulsion droplets synthesized using this homogenizing method is $2a = (40 \pm 8) \text{ nm}$.²⁸

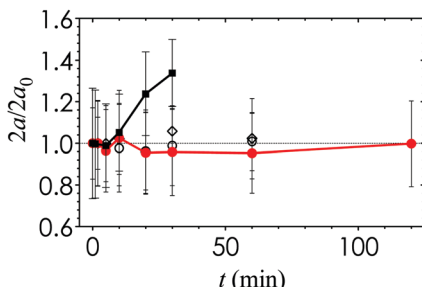


Fig. 2 Change in PDMS droplet size as a function of time for various photoinitiator additives. The additives are: 2 vol% butyl acetate (black open circles), 2 vol% butyl acetate with 0.067 g mL^{-1} TPO (black open diamonds), 2 vol% butyl acetate with 0.1 g mL^{-1} TPO (red filled circles), and 2.9 vol% hexanediol diacrylate with 0.08 g mL^{-1} TPO and 0.08 g mL^{-1} Darocur. Error bars represent the polydispersity of the droplets computed from fitting the raw autocorrelation data from dynamic light scattering with a log-normal distribution.

In order to synthesize the SLA printer ink, we require a photoinitiator additive that is compatible with the nanoemulsions and that is tailored to the wavelength of the laser source. The photoinitiator 2-hydroxy-2-methylpropiophenone (Darocur) is typically used in photocuring applications. However, we find that the nanoemulsion droplet size becomes unstable and increases significantly when Darocur is incorporated (Fig. 2). An alternative photoinitiator is phenylbis(2,4,6-trimethylbenzoyl) phosphine oxide (TPO), which exhibits significant absorbance in the $\lambda = 385$ to 420 nm wavelength range and high molar extinction coefficients.²⁹ TPO is commonly used in photocurable resins because of the overlap between its absorbance spectrum and the emission wavelength of the near-UV light sources in 3D printing. We find that adding 0.1 g mL^{-1} of TPO in a 2 vol% co-solvent of butyl acetate provides the best balance between solubility, nanoemulsion stability and photocurability. The droplet size remains stable over a period of 2 h, which is comparable to the total print time. The photoinitiator additive is added to the nanoemulsion suspension at 2 vol% to generate fresh samples prior to each 3D printing experiment.

SLA printer modifications and setup

During the polymerization step in the chemical crosslinking of PEGDMA, the presence of oxygen slows the reaction by scavenging reactive photoinitiator radicals.²² While this is normally undesirable, having a sufficiently thick oxygen inhibition layer below the cure zone is necessary to avoid scaffold adhesion to the optical window.^{19,23,24} We modify the resin tank of a commercially available desktop SLA printer (Form 2, FormLabs) by replacing the PDMS layer above the optical window with a thin, optically transparent Teflon coating to ensure a sufficient lubrication layer resulting from the unpolymerized ink.¹⁹ Teflon is effective because it has one of the highest oxygen permeabilities in polymeric materials ($1600 \text{ barrers} = 1.2 \times 10^{-14} \text{ m}^2 \text{ s}^{-1} \text{ Pa}^{-1}$)³⁰ that is double that of PDMS ($800 \text{ barrers} = 0.6 \times 10^{-14} \text{ m}^2 \text{ s}^{-1} \text{ Pa}^{-1}$).³¹ The procedure to modify the resin tank is as follows. After removal of the PDMS layer, the surface of the window is treated with a high frequency electrode gun (BD-10AS, Electro-Technic Products).

The plasma-treated window is evenly coated with 1H,1H,2H,2H-perfluoroctyltrimethoxsilane (Matrix Scientific) and baked in an oven ($T = 80 \text{ }^\circ\text{C}$) for at least 8 h. The window is then cleaned with pure ethanol, dried, and coated with a layer of 1 wt% poly(4,5-difluoro-2,2-bis-trifluoromethyl-1,3-dioxole-*co*-tetrafluoroethylene) (Teflon AF 2400) in tetradeacafluorohexane (FC-72). After the Teflon coating is completely dried, the edges of the optical window are reinforced with a UV adhesive (NOA 65, Norland Optics) to prevent nanoemulsion ink leakage into the optical chamber of the 3D printer.

To print hierarchical materials using the thermoresponsive nanoemulsions, the 3D printer is placed in an environmental incubator (Model 3110, Thermo Forma) set to six different temperatures ($T = 22.0 \text{ }^\circ\text{C}$, $28.5 \text{ }^\circ\text{C}$, $31.5 \text{ }^\circ\text{C}$, $34.0 \text{ }^\circ\text{C}$, $38.0 \text{ }^\circ\text{C}$, $39.5 \text{ }^\circ\text{C}$). The setup is allowed to equilibrate for at least 2 h prior to starting the printing experiment. Approximately 250 mL of the freshly prepared nanoemulsion ink is loaded into the resin tank and allowed to rest quiescently for 10 min to induce thermogelation. Samples used for confocal microscopy and scanning electron microscopy imaging are printed as slabs ($20 \text{ mm} \times 80 \text{ mm} \times 1.5 \text{ mm}$) with 15 layers (thickness = $100 \mu\text{m}$ per layer). The total print time for each slab is 32 min.

Rheological characterization

Rheological measurements on the nanoemulsions are carried out using a stress-controlled rheometer (AR-G2, TA Instruments) equipped with an aluminum 2° cone-and-plate geometry (diameter = 60 mm, truncation gap = $58 \mu\text{m}$). Small amplitude oscillatory measurements (strain $\gamma = 5 \times 10^{-4}$, angular frequency $\omega = 20 \text{ rad s}^{-1}$) are used in the temperature ramp (ramp rate = $0.5 \text{ }^\circ\text{C min}^{-1}$) and temperature jump studies ($T = 22.0 \text{ }^\circ\text{C}$, $30.0 \text{ }^\circ\text{C}$, $35.0 \text{ }^\circ\text{C}$, $40.0 \text{ }^\circ\text{C}$, $45.0 \text{ }^\circ\text{C}$). The linear viscoelastic moduli, G' and G'' , are monitored over the duration of the oscillatory measurements. In order to avoid slip, ultra fine sandpaper (320 grit) is adhered to the top geometry and the bottom Peltier plate for stress sweep experiments ($\gamma = 1 \times 10^{-4}$ to 15, $\omega = 20 \text{ rad s}^{-1}$), repeated yielding/recovery experiments ($\gamma = 10$, $\omega = 20 \text{ rad s}^{-1}$ and $\Delta t = 20 \text{ s}$ for the yield step; $\gamma = 5 \times 10^{-4}$, $\omega = 20 \text{ rad s}^{-1}$ and $\Delta t = 80 \text{ s}$ for the recovery step), as well as steady shear experiments (shear stress $\sigma = 0.01$ to 100 Pa) that measure the change in nanoemulsion viscosity as a function of shear rate. We use pure nanoemulsions that has been passed through a $1.5 \mu\text{m}$ nylon filter in the rheological studies. A temperature ramp oscillatory experiment on nanoemulsions containing 2 vol% of the photoinitiator mixture is conducted to check for consistency in the gelation temperature T_{gel} .

Sacrificial templating and bead infusion studies

The nanoemulsions are removed from the printed hydrogels using a gradual solvent transfer process. Briefly, samples are rinsed with pure ethanol, transferred into a 1 : 1 v/v isopropanol/ethanol mixture for 30 min, followed by a first soak in pure toluene for 2 h. The toluene bath is replaced with fresh solvent and samples are allowed to sit for at least 12 h to ensure complete removal of the PDMS, which is verified by a loss in fluorescence. To re-hydrate the samples, a reverse solvent transfer



into 1:1 v/v isopropanol/ethanol, 1:1 v/v ethanol/DI water, and finally pure DI water is used. Samples are kept in DI water for at least 24 h to swell to their equilibrium state.

Fluorescent carboxylated polystyrene beads (5 wt%, diameters = 50 nm, 200 nm, and 1.0 μ m, excitation/emission wavelengths = 505/515 nm) are purchased from Invitrogen and used without further purification. We place the porous hydrogel samples in 200 μ L of the bead suspension for 4 h. After rinsing with DI water, the samples are ready for confocal microscopy imaging.

Microscopy imaging

We use confocal laser scanning microscopy (CLSM) to visualize the internal structure of the printed samples containing nanoemulsions and those containing fluorescent beads after sacrificial templating. Imaging is performed on a Nikon A1R CLSM equipped with a resonant scanner head, diode lasers emitting at 488 and 561 nm, and a 60 \times oil immersion objective (NA = 1.4, working distance = 13 mm). Samples with dimensions of 5 mm \times 5 mm \times 1 mm are directly cut from the printed hydrogels and placed on a #1.5 coverslip for imaging in the *xy*- and *xz*-planes. The *xy*-plane images used for structural analysis have dimensions of 60.9 μ m \times 60.9 μ m with a pixel size of 119 nm. A fast Fourier transform (FFT) is applied to the raw images using the image processing software ImageJ (NIH). The radially averaged scattered light intensity of the FFT images, $I(q)$, is obtained using a radial profile plugin to ImageJ.

Porous hydrogel samples ($T = 22$ °C, 34.0 °C) are also imaged with a high resolution scanning electron microscope (HRSEM, Zeiss) at 5 kV accelerating voltage. Samples are cut into thin (~ 1 mm) sections, dried for at least 48 h at room temperature, and then adhered with carbon tape onto SEM stubs. All samples are sputter coated with approximately 10 nm of a Au-Pd alloy prior to imaging.

Mechanical characterization of filled and porous hydrogels

Nanoemulsion inks are photopolymerized into a traditional dogbone shape (cross sectional area, $A_0 = 1.00 \times 10^{-5}$ m 2 , testing area length, $L_0 = 15$ mm, thickness = 2 mm) for stress-strain mechanical testing. Briefly, 800 μ L of nanoemulsions is added into a dogbone-shaped mold made out of a transparent polycarbonate sheet. A microscope coverslip is used to cover the top of the mold. The filled mold is placed in a vacuum oven (heated to the various temperatures used in the 3D printing) purged with nitrogen gas in order to remove oxygen from the samples. This permits even curing during the photopolymerization and reduces the formation of air bubbles. After an hour, the mold is placed under a UV lamp for 5 minutes to fully photopolymerize the hydrogels. To generate porous hydrogels for mechanical testing, we further treat the dogbones with toluene and rehydrate using the procedure that is described in the experimental section under sacrificial templating.

Both ends of the hydrogel dogbones are attached to cardboard scaffold using UV glue (NOA 65, Norland Optics). The cardboard pieces are clamped directly onto a set of 10N load cells on an Instron 8840 MicroTester. The testing procedure involves the application of an uniaxial extensional strain at a rate of 0.01 mm s $^{-1}$ until rupture

occurs. Young's modulus is obtained using the relation, $E = \sigma_e/\epsilon = (FL_0)/(A_0\Delta L)$, where σ_e is the tensile stress, ϵ is the applied strain, F is the applied load, L_0 and A_0 are the original dimensions of the hydrogel dogbone, and ΔL is the deformation length measured by the Instron tester. The engineering rupture stress is obtained using the relation $\sigma_{rup} = F_{rup}/A_0$ where F_{rup} is the measured force at which the sample fractures. A minimum of six independently prepared samples are used to generate each data point.

3. Results and discussion

Printing of self-assembled nanoemulsion hydrogels

The schematic in Fig. 1A and B shows the dynamical process used to print nanoemulsion-filled hydrogels with macroscopic features (honeycomb and multi-tier woodpile hydrogels are shown in Fig. 1D). Using temperature as a tuning parameter, we are able to print hydrogels that have internal microstructure that range from nanometer-sized droplets to mesoscopic structures. At $T < T_{gel}$, the nanoemulsions are photo-crosslinked into a homogeneous matrix. When the temperature is raised beyond a critical gelation point T_{gel} , the hydrophobic end groups on the PEGDMA molecules partition into the oil phase and serve as interdroplet bridges³² (Fig. 1C), leading to the formation of a viscoelastic colloidal gel with different microstructures (Fig. 1E and F). Depending on application needs, these domains can either serve as pathways for the transport of hydrophobic molecules or be extracted to leave behind interconnected pores.

Rheological properties of the nanoemulsion inks

The viscoelasticity of the gelled nanoemulsions is an inherent material property that is directly correlated with the stress-bearing microstructure and dynamics of the gel network.^{33,34} As such, the internal length scale of the printed hydrogel scaffold can be estimated *a priori* by measuring the elastic (G') and viscous moduli (G'') of the nanoemulsions as a function of temperature using small amplitude oscillatory rheology.

The temperature ramp experiments shown in Fig. 2 are performed at a heating rate of 0.5 °C min $^{-1}$ to generate high resolution viscoelastic data at small temperature intervals. As temperature increases, G' and G'' increase rapidly and reach a plateau at high temperatures. The point at which $G' = G''$ sets the value of $T_{gel} = 29.2$ °C for this particular nanoemulsion system, although the gel temperature can be easily adjusted by replacing PEGDMA with other crosslinking molecules with two or more hydrophobic end groups.³² To account for the effect of the heating rate, we perform temperature jump measurements that are representative of print conditions. Fig. 3A indicates that G' and G'' follow the same trend as the temperature ramp experiments, and that T_{gel} is found within a similar range. The rheology data shown in Fig. 3A and 4B-D are generated using pure nanoemulsions. Addition of the photoinitiator additive results in a minor reduction in T_{gel} to 27.6 °C (Fig. 3B), although the overall rheological behavior remains similar.

In order to characterize the rheology of the nanoemulsion inks during the printing process, we estimate the compressive



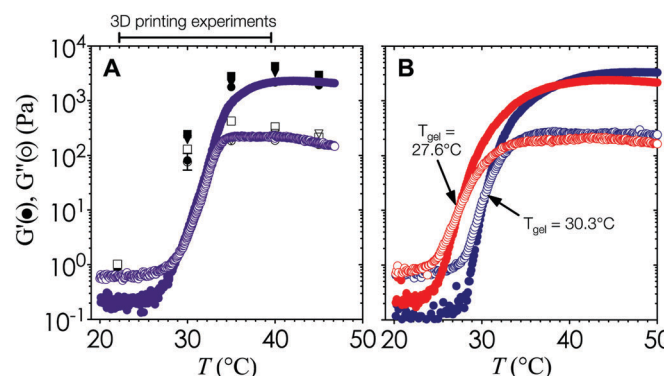


Fig. 3 Temperature responsiveness of the nanoemulsion inks. (A) Temperature ramp experiments captured at a heating rate of $0.5\text{ }^{\circ}\text{C min}^{-1}$ (purple) are overlaid with viscoelasticity data from the linear regime in the stress sweep measurements (squares), fully recovered samples in the yielding/recovery measurements (circles), and discrete temperature jump experiments (down triangles). (B) Comparison of the values of T_{gel} for samples with (red) and without the photoinitiator additives (blue). Solid symbols represent G' and open symbols represent G'' .

strain, γ , as the relative vertical displacement of the fluid induced by the platform movement, and the strain rate, $\dot{\gamma}$, using the linear velocity of the motorized stage through the ink (Fig. 4A). The motorized platform moves at an average speed of 1.27 cm s^{-1} in the withdrawal and re-submersion steps (steps 2 and 3 in Fig. 1A). There is a constant gap of $h_{\text{gap}} = 100\text{ }\mu\text{m}$, set by the z -resolution of the printer, between the optical window and the bottom of the printed model. The total height of the fluid stays relatively constant at $h_{\text{total}} \sim 7.5\text{ mm}$. Yielding of the gels occurs at strain units ($\gamma < 0.01$) lower than the deformation imposed during the print process by the build platform in the printer ($\gamma \sim 0.98$).

Fig. 4B shows that there is a transition from the linear G_{lin}' at low γ to the nonlinear regime beyond the yield strain γ_y , where G' decreases by more than three orders of magnitude. We note here that the angular frequency (ω) of the oscillatory rheology is set to 20 rad s^{-1} to generate high resolution data on the rheometer. This value of ω provides the correct values G' and G'' particularly when they become insensitive to differences in ω above the gel point of a viscoelastic suspension.³⁵ We use repeated cycles of large amplitude oscillatory shear (LAOS) and small amplitude oscillatory shear (SAOS) to investigate the structural recovery of the inks under the deformation imposed during the printing process. In the LAOS cycle, a strain amplitude much larger than the yield strain of the nanoemulsion is applied to simulate the withdrawal and submersion steps of printing. In the SAOS cycle, a small strain is used to probe the viscoelastic recovery of the ink during the print step. The timescales and parameters of the LAOS and SAOS cycle are matched to the printing step timescales as shown in Fig. 1A. Fig. 4C shows that the gelled nanoemulsions recover to its original state rapidly and reversibly once the deformation ceases. This structural recovery is key to printing hydrogels with continuous and well-controlled internal microstructure in all three dimensions.

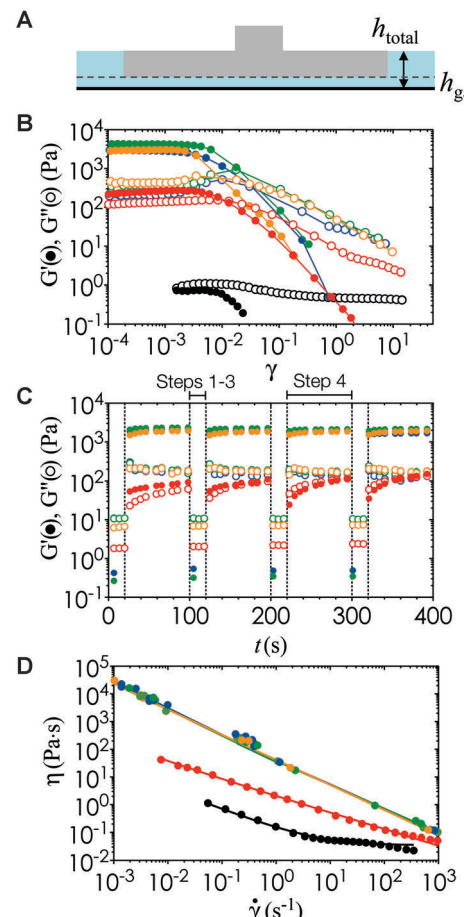


Fig. 4 Rheology of thermoresponsive nanoemulsions. (A) Schematic of the motorized build platform imposing a compressive strain at a known shear rate on the nanoemulsion ink. For our printing setup, $\gamma = 0.98$ and $\dot{\gamma} = 1.7\text{ s}^{-1}$. (B) Stress sweep experiments determines the linear regime and the yield strain at $T = 22\text{ }^{\circ}\text{C}$ (black), $30\text{ }^{\circ}\text{C}$ (red), $35\text{ }^{\circ}\text{C}$ (orange), $40\text{ }^{\circ}\text{C}$ (green) and $45\text{ }^{\circ}\text{C}$ (blue). Solid lines guide the eye. (C) Repeated large amplitude and small amplitude oscillatory experiments show that nanoemulsions recover rapidly and reversibly. Dashed lines mark the boundaries between yielding and recovery steps. The time scales and applied deformations are chosen to match the print and retraction steps in the 3D printer. Solid symbols represent G' and open symbols represent G'' . (D) Steady shear rheology show yield stress behavior and shear thinning over the range of shear rates probed. Solid lines are Herschel–Bulkley fits.

Within the operation range of the moving build platform ($\dot{\gamma} = 1.7\text{ s}^{-1}$), the nanoemulsions undergo shear thinning with a viscosity upper range of $\sim 100\text{ Pa s}$ (Fig. 4D). The yield stress, σ_y , and the power law exponent, n , can be obtained from fitting the Herschel–Bulkley model $\sigma = \sigma_y + k\dot{\gamma}^n$ to the viscosity data, where k is a consistency parameter and the steady shear viscosity is $\eta = \sigma/\dot{\gamma}$.³⁶ A purely Newtonian liquid has constant viscosity at all shear rates ($n = 1$), whereas $0 \leq n < 1$ for a shear thinning fluid. These material properties are shown in Table 1 for nanoemulsions at different temperatures. In addition, we perform steady state characterization of the thermoresponsive nanoemulsions at $T = 22\text{ }^{\circ}\text{C}$ and $40\text{ }^{\circ}\text{C}$, which encompasses the full range of conditions tested in our study. We observe some viscosity hysteresis of the nanoemulsions at $T = 22\text{ }^{\circ}\text{C}$,

Table 1 Rheological properties of nanoemulsion inks

T (°C)	σ_y (Pa)	n	G_{lin}' (Pa)
22.0	0.06	0.75	0.17
30.0	0.08	0.41	3.24
35.0	22	0.25	899.4
40.0	25	0.30	2224
45.0	28	0.28	2210

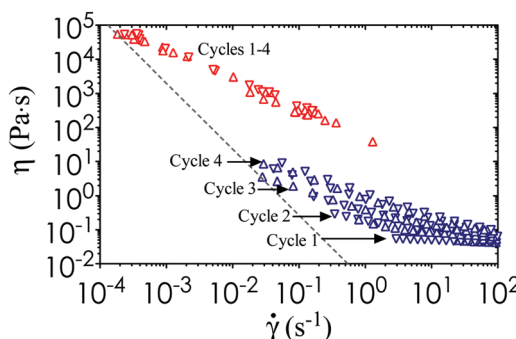


Fig. 5 Viscosity hysteresis in thermoresponsive nanoemulsions. Steady state viscosity plotted as a function of shear rate for four upwards (up triangles) and downwards (down triangles) stress sweep cycles at $T = 22\text{ }^{\circ}\text{C}$ (red) and $40\text{ }^{\circ}\text{C}$ (blue). Dashed line represents instrument sensitivity limits.

but none at $T = 40\text{ }^{\circ}\text{C}$ (Fig. 5). This gradual increase in viscosity over multiple cycles at low temperatures could be explained by the formation of flow-induced clusters from the Brownian

relaxation of the nanoemulsions and structural relaxation of the PEGDMA matrix with respect to viscous forces.³⁷ Nevertheless, the printed microstructure is unlikely to be affected by the shear imposed by the build platform given the long equilibration time during the print step. The yield stress of the material does present issues if the print area is not properly refilled after the withdrawal of the build platform. Classical Saffman-Taylor instabilities can also lead to viscous fingering that are comparable to the length scale of the self-assembled gel network within the nanoemulsions.³⁸ Thus, wiper operations are necessary to spread the ink out evenly between each layer-by-layer cycle in the printing process.

Hydrogel microstructure and mechanical properties

Fig. 6A is a panel of representative confocal laser scanning microscopy (CLSM) images in the xy -plane that show the range of microstructures of the hydrogels printed with the self-assembling nanoemulsions. Visual inspection shows that the scaffold is mostly homogeneous below the gel temperature at $T = 22.0\text{ }^{\circ}\text{C}$. Finely percolated networks form near the gel point at $T = 28.5\text{ }^{\circ}\text{C}$, which develop into spinodal-like structures at $T = 34.0\text{ }^{\circ}\text{C}$. As the temperature is further increased to $T = 39.5\text{ }^{\circ}\text{C}$, arrested phase separation results in a decrease in the length scale of the colloid-rich regions. We compute the scattered intensity $I(q)$ from a fast Fourier transform (FFT) applied to the raw 2D images (Fig. 6B), where q represents the wave vector or inverse length in real space. The characteristic gel

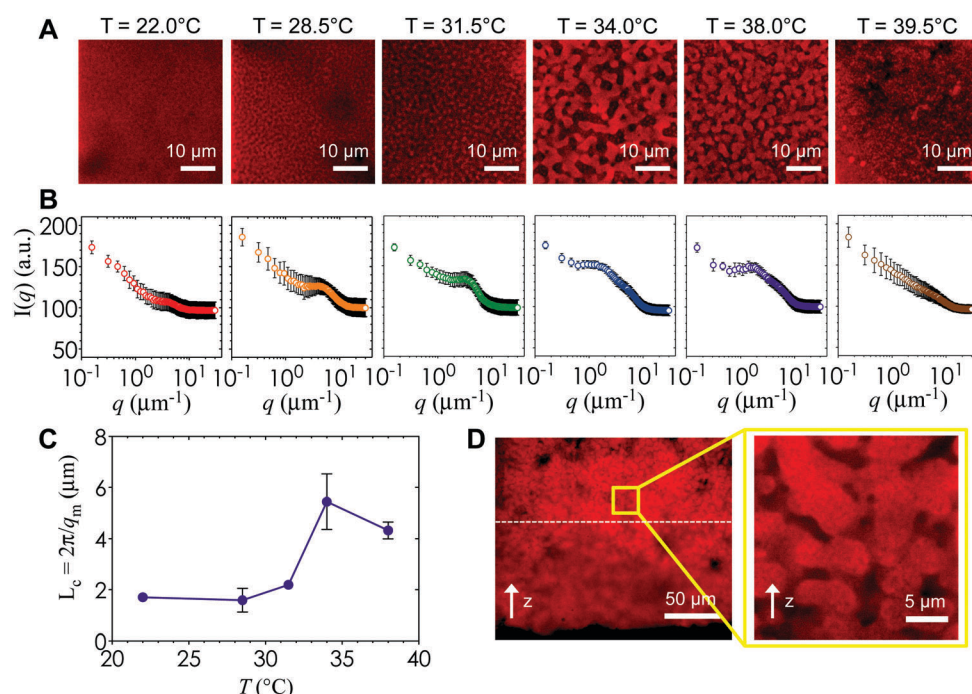


Fig. 6 Microstructure of 3D printed hydrogels using thermoresponsive nanoemulsions. (A) Representative 2D confocal laser scanning microscopy (CLSM) images of the internal microstructure of printed hydrogels, where fluorescent regions belong to poly(dimethyl siloxane) (PDMS) droplets. (B) Intensity $I(q)$ as a function of the wave vector q , for samples printed at $T = 22.0\text{ }^{\circ}\text{C}$ (red), $28.5\text{ }^{\circ}\text{C}$ (orange), $31.5\text{ }^{\circ}\text{C}$ (green), $34.0\text{ }^{\circ}\text{C}$ (blue), $38.0\text{ }^{\circ}\text{C}$ (purple) and $39.5\text{ }^{\circ}\text{C}$ (brown). (C) Characteristic length scales of the self-assembled PDMS droplets from $L_c = 2\pi/q_m$, where q_m is obtained from peaks in $I(q)$. Error bars in (B) and (C) are standard deviations from 3 independent measurements within a sample. (D) Zoomed out image in the xz -plane of a hydrogel scaffold printed at $T = 34.0\text{ }^{\circ}\text{C}$. Dashed line indicates boundary between two printed layers. Inset: Zoomed in image of the same sample.



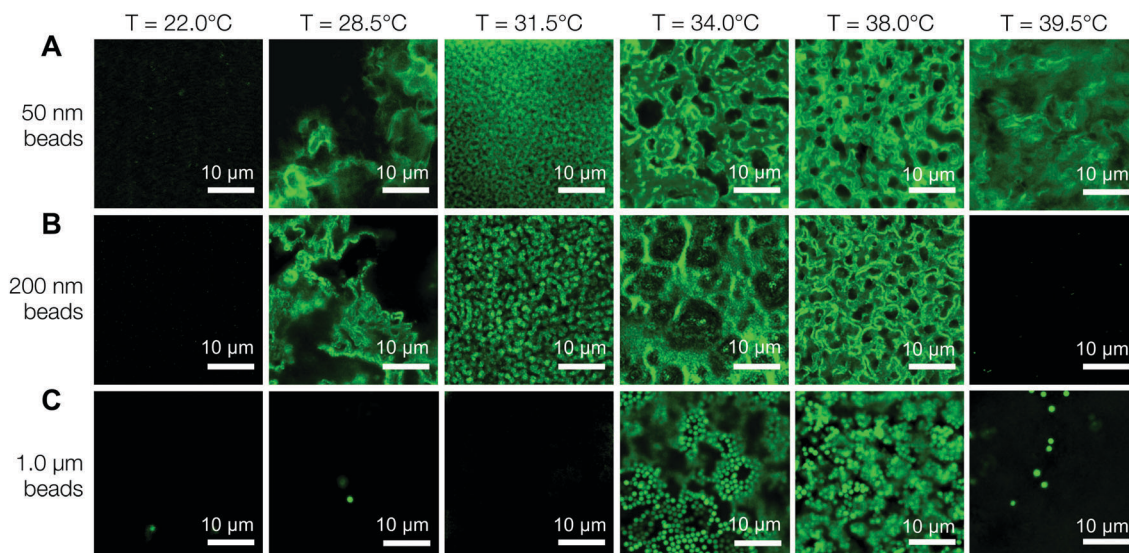


Fig. 7 Printed porous hydrogels as size-selective membranes. Representative CLSM images of mesoporous hydrogel samples in which fluorescent polystyrene beads have been introduced. Beads of diameters (A) 50 nm, (B) 200 nm and (C) 1.0 μ m are allowed to diffuse into the interconnected porous network. Fluorescence here indicates the passage of beads into the sample. All images are captured at $z = 10$ μ m above the bottom of the sample.

length scale, $L_c = 2\pi/q_m$, is defined by the local peak q_m in the $I(q)$ plots where possible and is plotted in Fig. 6C. The value of L_c increases gradually to a maximum at $T = 34.0$ °C, followed by a decrease that is brought on by kinetic arrest at higher temperatures.¹⁶ These results are in agreement with our earlier work on a similar system of thermoresponsive nanoemulsions in which samples were cast into molds.³⁹ In addition, we verify that the connective mesoscale structure is preserved between each printed layer. A zoomed out CLSM image of the hydrogel scaffold in the direction perpendicular to the printing (xz-plane) is shown in Fig. 6D, where a dashed line indicates the boundary of the two layers as observed in HRSEM images in Fig. 8A.

The interconnectivity of the gel networks formed by the self-assembled thermoresponsive nanoemulsions allows for the transport of molecules and mesoscopic objects. We demonstrate this capability by using the nanoemulsions as a sacrificial template that is extracted from the 3D printed hydrogels through solvent transfer with pure toluene. After rehydration, we immerse the samples in aqueous solutions containing fluorescent, carboxylated polystyrene beads with diameters of 50 nm, 200 nm, and 1.0 μ m. Fig. 7 illustrates the capability of the printed porous hydrogels to serve as size-selective membranes. Beads of all sizes are able to diffuse into hydrogels having comparatively larger pore size ($T = 34.0$ °C and 38.0 °C) but not those with smaller pores ($T = 28.5$ °C, 31.5 °C, 39.5 °C). The control scaffold sample printed at $T = 22.0$ °C ($T < T_{gel}$) does not permit the passage of the smallest bead size (50 nm) because of the isolated pore structure within it. A representative 3D CLSM image of a porous sample printed at $T = 34.0$ °C and treated with 200 nm beads shows that the fluorescent beads are able to diffuse into the channels in x , y , and z -directions (Fig. 8B). The xz-image of a porous sample ($T = 34.0$ °C) in Fig. 8C is captured from an xyz-stack; here, the imaging height is limited to 24 μ m due to significant scattering from the remainder of the sample.

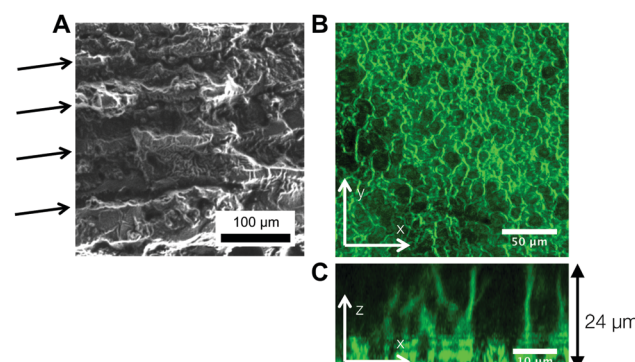


Fig. 8 Images of printed hydrogels in three dimensions. (A) HRSEM of porous hydrogel scaffold ($T = 34.0$ °C) in the xz-plane, showing layered structure from SLA printing. Arrows indicate approximate location of boundaries spaced 100 μ m apart. (B and C) CLSM images of the same scaffold treated with fluorescent polystyrene beads ($2a = 200$ nm) in the (B) xy-plane and (C) xz-plane. The images in (B and C) are captured as an xyz-stack, where the z-dimension is limited to 24 μ m because of significant backscatter from the rest of the sample.

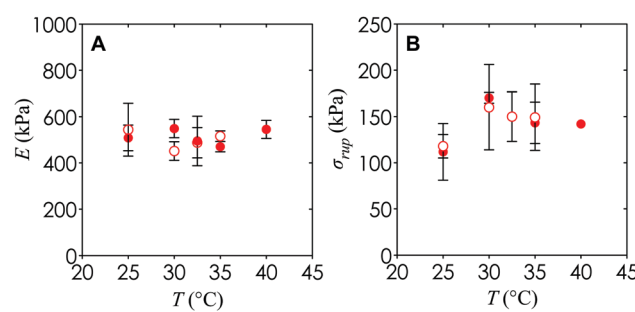


Fig. 9 Mechanical properties of filled and porous hydrogel scaffolds. (A) Young's modulus and (B) engineering rupture stress of the filled (solid) and porous (open) as a function of T . Error bars are standard deviations from 6 independent samples.

An additional observation is that the Young's modulus and engineering rupture stress for hydrogels with and without nanoemulsions remain similar to within the experimental uncertainty (Fig. 9). This suggests that the mechanical properties of the hydrogel can be tuned completely independently from the internal architecture necessary for diffusion of nutrients or cell passage, which could be important in future applications of this material towards synthetic microvasculature and 3D cellular scaffolds.

4. Conclusions

Thermoresponsive nanoemulsions belong to a class of stimuli-responsive materials that are compatible with 3D SLA printing to generate hierarchical materials with tunable mesoscale structures. The PEGDMA molecules in the nanoemulsion ink serve two important functions: (1) they induce colloidal gelation in which the characteristic length scale can be tuned based on the temperature, and (2) they provide photochemically cross-linked hydrogel materials on demand. Using a commercially available 3D printer with modifications to the resin tray to enhance oxygen permeability, we demonstrate that these nanoemulsions can serve as self-assembling precursor inks to be rapidly photopolymerized into macroscale shapes with the desired internal features. Correct selection of the photoinitiator is critical to maintain the stability of the nanoemulsions over the course of the printing process and to provide photoreactive response with the near-UV ($\lambda = 405$ nm) light sources that are commonly used in commercial 3D SLA printers. In addition, the rapid structural recovery and shear thinning rheological properties of the ink is necessary for generating hierarchical materials with an interconnected morphology that allows the mass transport of mesoscopic objects into the hydrogels. However, due to the yield stress behavior of these inks, wiper operations between successive print steps are needed to spread the ink out evenly.

Future possibilities for these mesoporous materials include the crystallization of hydrophobic molecules in nanoemulsion-carrying channels for controlled or triggered drug delivery,⁴⁰ synthetic vasculature,⁴¹ or in understanding the deformation mechanics of tissue-like models that can be printed in multiple shapes and sizes.⁴² Other applications involve retaining the nanoemulsions in the printed structure to serve as reservoirs for hydrophobic drugs,⁴³ modulating the mass transfer of gases,⁴⁴ or acting as sinks for metals/organics in water treatment processes.⁴⁵

Competing financial interest

The authors declare no competing financial interest.

Acknowledgements

This work was supported primarily by the MRSEC Program of the National Science Foundation under DMR – 1419807. The authors thank Prof. G. McKinley for the use of the rheometer and J. F. Hamel for loan of the environmental incubator.

References

- 1 S. C. Glotzer and M. J. Solomon, *Nat. Mater.*, 2007, **6**, 557.
- 2 J. Malda, J. Visser, F. P. Melchels, T. Jüngst, W. E. Hennink, W. J. Dhert, J. Groll and D. W. Hutmacher, *Adv. Mater.*, 2013, **25**, 5011.
- 3 C.-C. Lin and K. S. Anseth, *Pharm. Res.*, 2009, **26**, 631.
- 4 B. V. Slaughter, S. S. Khurshid, O. Z. Fisher, A. Khademhosseini and N. A. Peppas, *Adv. Mater.*, 2009, **21**, 3307.
- 5 L. Hockaday, *et al.*, *Biofabrication*, 2012, **4**, 035005.
- 6 A. Khademhosseini and R. Langer, *Biomaterials*, 2007, **28**, 5087.
- 7 M. P. Cuchiara, D. J. Gould, M. K. McHale, M. E. Dickinson and J. L. West, *Adv. Funct. Mater.*, 2012, **22**, 4511.
- 8 P. B. Welzel, J. Friedrichs, M. Grimmer, S. Vogler, U. Freudenberg and C. Werner, *Adv. Healthcare Mater.*, 2014, **3**, 1849.
- 9 L.-W. Xia, R. Xie, X.-J. Ju, W. Wang, Q. Chen and L.-Y. Chu, *Nat. Commun.*, 2013, **4**, 2226.
- 10 A. N. Stachowiak, A. Bershteyn, E. Tzatzalos and D. J. Irvine, *Adv. Mater.*, 2005, **17**, 399.
- 11 Y. J. Lee and P. V. Braun, *Adv. Mater.*, 2003, **15**, 563.
- 12 P. X. Ma and J.-W. Choi, *Tissue Eng.*, 2001, **7**, 23.
- 13 M. N. Lee and A. Mohraz, *Adv. Mater.*, 2010, **22**, 4836.
- 14 Y. S. Nam and T. G. Park, *J. Biomed. Mater. Res.*, 1999, **47**, 8.
- 15 H. Diestra-Cruz, E. Bokusoglu, N. L. Abbott and A. Acevedo, *ACS Appl. Mater. Interfaces*, 2015, **7**, 7153.
- 16 J. Conrad, H. Wyss, V. Trappe, S. Manley, K. Miyazaki, L. Kaufman, A. Schofield, D. Reichman and D. Weitz, *J. Rheol.*, 2010, **54**, 421.
- 17 V. J. Anderson and H. N. Lekkerkerker, *Nature*, 2002, **416**, 811.
- 18 P. J. Lu, E. Zaccarelli, F. Ciulla, A. B. Schofield, F. Sciortino and D. A. Weitz, *Nature*, 2008, **453**, 499.
- 19 J. R. Tumbleston, *et al.*, *Science*, 2015, **347**, 1349.
- 20 A. S. Gladman, E. A. Matsumoto, R. G. Nuzzo, L. Mahadevan and J. A. Lewis, *Nat. Mater.*, 2016, **15**, 413.
- 21 M. N. Cooke, J. P. Fisher, D. Dean, C. Rimnac and A. G. Mikos, *J. Biomed. Mater. Res., Part B*, 2003, **64**, 65.
- 22 S. C. Ligon, B. Husár, H. Wutzler, R. Holman and R. Liska, *Chem. Rev.*, 2013, **114**, 557.
- 23 D. Dendukuri, P. Panda, R. Haghgoie, J. M. Kim, T. A. Hatton and P. S. Doyle, *Macromolecules*, 2008, **41**, 8547.
- 24 D. Dendukuri, D. C. Pregibon, J. Collins, T. A. Hatton and P. S. Doyle, *Nat. Mater.*, 2006, **5**, 365.
- 25 K. J. LeBlanc, S. R. Niemi, A. I. Bennett, K. L. Harris, K. D. Schulze, W. G. Sawyer, C. Taylor and T. E. Angelini, *ACS Biomater. Sci. Eng.*, 2016, **2**, 1796.
- 26 R. G. Larson, *The Structure and Rheology of Complex Fluids*, Oxford University Press, New York, NY, 1999.
- 27 B. G. Compton and J. A. Lewis, *Adv. Mater.*, 2014, **26**, 5930.
- 28 A. Gupta, H. B. Eral, T. A. Hatton and P. S. Doyle, *Soft Matter*, 2016, **12**, 1452.
- 29 A. A. Pawar, G. Saada, I. Cooperstein, L. Larush, J. A. Jackman, S. R. Tabaei, N.-J. Cho and S. Magdassi, *Sci. Adv.*, 2016, **2**, e1501381.
- 30 I. Pinna and L. G. Toy, *J. Membr. Sci.*, 1996, **109**, 125.
- 31 T. Merkel, V. Bondar, K. Nagai, B. Freeman and I. Pinna, *J. Polym. Sci., Part B: Polym. Phys.*, 2000, **38**, 415.



32 M. E. Helgeson, S. E. Moran, H. Z. An and P. S. Doyle, *Nat. Mater.*, 2012, **11**, 344.

33 L. C. Hsiao, R. S. Newman, S. C. Glotzer and M. J. Solomon, *Proc. Natl. Acad. Sci. U. S. A.*, 2012, **109**, 16029.

34 L. C. Hsiao, H. Kang, K. H. Ahn and M. J. Solomon, *Soft Matter*, 2014, **10**, 9254.

35 H. H. Winter and F. Chambon, *J. Rheol.*, 1986, **30**, 367.

36 H. Zhu, Y. Kim and D. De Kee, *J. Non-Newtonian Fluid Mech.*, 2005, **129**, 177.

37 J. Kim and M. E. Helgeson, *Phys. Rev. Fluids*, 2016, **1**, 043302.

38 J. Nase, A. Lindner and C. Creton, *Phys. Rev. Lett.*, 2008, **101**, 074503.

39 L. C. Hsiao and P. S. Doyle, *Soft Matter*, 2015, **11**, 8426.

40 A. Z. M. Badruddoza, P. D. Godfrin, A. S. Myerson, B. L. Trout and P. S. Doyle, *Adv. Healthcare Mater.*, 2016, **5**, 1960.

41 N. Doshi, B. Prabhakarpandian, A. Rea-Ramsey, K. Pant, S. Sundaram and S. Mitragotri, *J. Controlled Release*, 2010, **146**, 196.

42 T. W. Sirk, K. S. Khare, M. Karim, J. L. Lenhart, J. W. Andzelm, G. B. McKenna and R. Khare, *Polymer*, 2013, **54**, 7048.

43 H. Z. An, M. E. Helgeson and P. S. Doyle, *Adv. Mater.*, 2012, **24**, 3838.

44 H. Z. An, E. R. Safai, H. B. Eral and P. S. Doyle, *Lab Chip*, 2013, **13**, 4765.

45 A. Sayari, S. Hamoudi and Y. Yang, *Chem. Mater.*, 2005, **17**, 212.

