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Continuous Synthesis of Elastomeric Macroporous Microbeads

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Macroporous microbeads are synthesized by microfluidic production of silica-loaded polymeric microdroplets followed by porogen removal via selective etching. Microdroplets are produced in a flow-focusing microreactor to ensure monodispersity with uniform porogen loading. Effects of porogen size and polymer network density on the porosity and effective modulus of the microbeads are studied.

Macroporous microparticles have important applications in areas where enhanced surface area and rough surface topology are preferred over dense spherical microparticles.¹ The macroporous microparticles are utilized in applications ranging from catalysis² and selective adsorbents³, to enzyme immobilization⁴, drug delivery⁵, and cell growth scaffolding^{6,7}. Porous monoliths have been considered as reactor packing⁸ and catalyst scaffolds, but suffer from flow channelling, which is mitigated in particulate beds. Porous microparticles have been fabricated using a wide variety of materials (e.g., ceramics, oxides, polymers, and hydrogels)^{6,9-12} and methods (e.g., emulsification¹³, templating¹⁴, and self-assembly) in both bulk¹⁰ and flow reactors^{1,15,16}. Crosslinked polysiloxane networks have been demonstrated to have tunable viscoelastic and surface properties^{17,18} which can be varied to promote cell growth, as well as favourable biocompatibility and oxygen transport¹⁸. Porous particles of linear polysiloxanes have been synthesized using salt-induced osmosis of water microdroplets into the droplets of linear polysiloxane that are formed during the bulk emulsification process¹³. Despite the high degree of porosity, bulk emulsification methods have the downside of a very wide particle size distribution due to the uncontrolled breakup in the mechanical emulsification process. Over the past two decades, droplet-based microfluidic strategies -owing to precise control

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over the flow and reaction parameters– have been successfully utilized for continuous synthesis of a wide range of micro/nano structures. $^{19\mathchar{-}24}$

In this work, we demonstrate a microfluidic strategy for continuous synthesis of elastomeric macroporous microbeads with tuneable size and porosity.

Depending on the intrinsic properties of the substrate material, different microfluidic approaches have been utilized to create macroporous beads. Such methods include fabrication of intrinsically porous beads through solvent removal and spontaneous phase separation²⁵, as well as structural templating with immiscible fluids (basic and high-internal-phase emulsions (HIPE))^{14,26} and solids (e.g., salt, sugar, polystyrene, and silica)³. Unique physicochemical properties of polysiloxanes limit the feasibility of the more conventional microfluidic approaches for forming macroporous beads. For example, (Poly)hydromethylsiloxane (PHMS) is not compatible with spontaneous phase separation or solvent removal, as the network must be chemically crosslinked through hydrosilylation. Additionally, the lack of UV or thermal initiated crosslinking means that the PHMS microbeads must be formed with porogen present as the crosslinking reaction is occurring. During preliminary screening, it was found that the fluid-based templating of PHMS droplets had major drawbacks for continuous synthesis of macroporous microbeads, namely the stability and droplet size of the formed water-in-oil or gas-in-oil emulsions. The surfactant required to generate stable water-inoil emulsions would cause immiscibility issues with the PHMS and crosslinker mixture, thus preventing the hydrosilylation catalyst from mixing with the polymer phase to begin crosslinking. Thus, solid porogens were considered as the templating material. The conventional sacrificial polystyrene (PS) microspheres were rejected due to their solubility in the PHMS crosslinking solvent (toluene), which would dissolve the PS microspheres before the PHMS could solidify into a porous network. Crystalline solids (e.g., sodium chloride and sucrose) were rejected for similar reasons; the microscale particles would migrate to the fluid interface between the nonpolar

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PHMS and the polar continuous phase (water) and promptly dissolve before the PHMS network could crosslink. However, mesoporous amorphous silica microspheres are insoluble in all the fluids used in the flow synthesis of elastomeric microbeads and can be removed selectively during post-processing. Potassium hydroxide (KOH) selectively etches the amorphous silica microspheres, leaving behind the porous PHMS scaffold. An overview of this process is shown in Fig. 1.



Fig. 1 (I) Major components of the microparticles: PHMS (polymer), 1,5 – hexadiene (crosslinker), and mesoporous silica microparticles (porogen). (II) Liquid PHMS droplet loaded with silica microparticles produced by the flow-focusing microreactor. (III) Porogen-loaded PHMS microbead after crosslinking by hydrosilylation. (IV) Schematic of the formed macroporous microbead after selective removal of the solid porogen with chemical etching.

Experimental

In this work, building on our recently developed capillary-based microfluidic reactor for dense PHMS particles²⁷, we developed a flow synthesis strategy (Fig. 2) for continuous production of porous silicone elastomer microbeads with tunable size and porosity.

Chemicals

(Poly)hydromethylsiloxane (PHMS) (average Mn 1,700-3200), 1,5-hexadiene, platinum(0)-1,3-divinyl-1,1,3,3-tetramethlydisiloxane (Karstedt's Catalyst) (in Xylene, Pt-2wt%), toluene (anhydrous, 99.8%), , and sodium dodecyl sulfate (SDS), were purchased from Sigma Aldrich. A 500 molecular weight vinyl-terminated polydimethylsiloxane crosslinker (DMS-V03, hereafter V03) was purchased from Gelest Inc. and used as received. Acetone and 2-propanol (IPA) were purchased from VWR Analytical. Silica gels (HPLC grade, spherical 5, 7.5, 10 and 20 μ m APS, 120 angstroms) were purchased from Alfa Aesar. All chemicals were used as received. Deionized water was obtained in-house using a PURELAB Flex purification unit.

Experimental Setup

The microfluidic platform, shown in Fig.2, consists of a 3D coaxial flow-focusing microreactor (producing uniform microdroplets - the silicone networks), three syringe pumps

The microreactor consists of two inner coaxial borosilicate glass capillaries (0.75 mm inner diameter, 1.0 mm outer diameter) with their tips separated by ~1mm. Both inner capillaries are sheathed in an outer glass capillary (1.12 mm inner diameter, 1.5 mm outer diameter) resulting in two coaxial inner microchannels and two annular microchannels on either side of the gap. A Full diagram of the necessary fluidic connections is shown in the supplementary information, Figure S1. The capillary-based microreactor constructed using off-the-shelf components offers advantages in cost and rapid prototyping compared to expensive custom microfabricated reactors. The inlet streams including two dispersed and one continuous phase are delivered into the microreactor using syringe pumps and gas-tight glass (dispersed phase) and stainless-steel (continuous phase) syringes. The inlet side inner microchannel contains a mixture of the polymer (PHMS), crosslinker (1,5 - hexadiene or V03), solvent (toluene), and silica microspheres of selected sizes (from 1.6-10 μ m), while the inlet side annulus contains a mixture of toluene and crosslinking catalyst to begin the hydrosilylation crosslinking on chip. Water and sodium dodecyl sulfate (SDS) are fed through the outlet-side annulus as the continuous phase (with surfactant) for facile microdroplet breakup and stabilization until crosslinking is finished in the collection bath. The center microchannel of the outlet side is a flamed-tip constricted glass capillary (produced using a Narishige PC-10 capillary puller) to promote breakup of the polymer mixture jet into small droplets and carry them to the collection bath (filled with DI water) where the crosslinking is



Fig. 2 (A) Schematic of the developed microfluidic platform for the continuous synthesis of the macroporous microbeads. The inset shows the microbeads collection bath. (B) An exemplary image of the capillary-based flow-focusing microreactor utilized for the synthesis of elastomeric microbeads. $Q_1=10 \mu$ L/min, $Q_2=10 \mu$ L/min, $Q_3=200 \mu$ L/min.

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completed. Upon crosslinking of the in-flow generated droplets, spherical silicone elastomer microbeads filled with close-packed silica microspheres are formed. In the next step, the porogen (*i.e.*, mesoporous amorphous silica microspheres) is selectively removed using potassium hydroxide solution. When the silica loading inside the microbeads is close-packed, the etching process creates interconnected pores throughout the elastomeric microbead.

Reagent Preparation

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The precursor solutions were prepared as detailed in Table 1. The silica-loaded polymer precursor solution was prepared by mixing the polymer and crosslinker in the desired ratio (3:1 and 5:1 PHMS:crosslinker by mole basis). The viscosity of the polymer precursor solution was reduced by adding an organic solvent (toluene) prior to the addition of the silica microspheres. Silica microspheres of the desired size (1.6 - 10 μ m) were then added to the polymer mixture until the mixture set into a colloidal gel (*i.e.*, vial could be inverted without flow). Next, an additional 25% v/v of the polymer was added to ensure the mixture could flow in the syringe, microreactor, and tubing and connections. The crosslinking catalyst concentration was precisely tuned to ensure prompt crosslinking of the microbeads in the collection bath to prevent possible coalescence during collection while ensuring the mixture remained a liquid during the in-flow mixing and breakup inside the flow-focusing microreactor. The crosslinking catalyst concentration (i.e., crosslinking time) was tuned off-line using small (100 μ L) batch gel tests. The maximum concentration of Pt in the polymer prior to the washing steps is ~ 40 μ M which is significantly lower than EPA reported values of Pt toxicity (0.1mmol/kg)²⁸ or Pt catalyzed reactions.

DI water with 3 wt% SDS was used as the continuous phase to lower the surface energy of the polymer jet / water interface and promote breakup while preventing droplet coalescence before crosslinking in the collection vial.

Microdroplet Formation

| Stream | Components | Purpose |
|----------------|-------------------------|-----------------------|
| Q1 | Toluene | Solvent and Diluent |
| | Karstedt's Catalyst | Crosslinking Catalyst |
| Q ₂ | PHMS | Polymer |
| | 1,5-hexadiene or V03 | Crosslinker |
| | Silica gel | Porogen |
| | Toluene | Solvent and Diluent |
| Q ₃ | Water | Continuous Phase |
| | SDS | Surfactant |

Table 1. Different stream components of the flow-focusingmicroreactor shown in Fig. 2.

Before collection of the elastomeric microbeads, the flowrates of the various streams were adjusted to find a stable dripping flow regime, as droplet formation from a polymer-based liquid /solid colloid mixture is non-Newtonian. The main parameters adjusted were (a) the ratio between the two inlet streams (*i.e.*, $Q_1:Q_2$) to vary the loading of silica within the microdroplet and the viscosity of the microdroplet mixture, and (b) the total dispersed phase and continuous phase flow rates ($Q_1+Q_2 \& Q_3$) which control the velocity of the two fluids at the flow-focusing orifice and the resulting flow regime within the microreactor.

Microparticle Collection

The silica-loaded PHMS microdroplets were collected in a large water bath to prevent contact between droplets before they finish crosslinking. The microreactor outlet was inserted into the water in the collection vial to eliminate dripping of the microreactor effluent into the collection vial and its potential effects on the droplet shape and porogen loading. Following collection, the microdroplets were allowed to crosslink in the water bath for 24 h to ensure complete crosslinking step, the elastomeric microbeads were filtered (Whatman 1004-100 20 μ m pores) and washed to remove the surfactant (SDS). The microbeads washing protocol involved two rinses with DI water followed by two rinses with isopropanol and two rinses with acetone. After washing, the microbeads were dried at room temperature.

Selective Removal of Solid Porogen

Following washing and drying of the silica-loaded elastomeric microbeads, the silica microspheres were selectively removed from the elastomer network by etching with 1M KOH. A small amount of acetone (~100 μ L per mL) was added to the 1M KOH solution to reduce the surface tension and achieve complete porogen removal. When pure KOH is used, surface tension can prevent penetration of the etchant into small pores within the silica-loaded and semi-etched elastomeric microbeads. The PHMS microbeads were left in the etchant solution for 2 h to achieve complete removal of the solid porogen. Following the solid porogen removal, the washing and drying protocol was repeated, and the samples were prepared for further analysis using electron microscopy and X-ray microcomputed tomography.

Porosity Characterization

SEM Imaging

Following the synthesis, drying, and porogen removal steps, the elastomeric microbeads were characterized using scanning electron microscopy (FEI Verios 460L Field Emission SEM and Hitachi 3200N Variable Pressure SEM) to evaluate the degree of silica loading, particle size, and size dispersity as well as the pore network inside the porous microbeads.

MicroCT Imaging

Samples of the synthesized elastomeric microbeads (both etched and unetched) were subjected to a micro-computed tomography (μ CT) scan (Nikon XTH 225 ST) with a voxel size of ~6 μ m to verify that the mesoporous silica microspheres were completely etched without causing structural damage to the elastomeric microbeads in the etch bath.

Mechanical Tests: Effective Modulus

Effective moduli of the different PHMS microbeads (control, silica-

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Fig. 3 Schematic of the modulus test setup using a parallel plate configuration. Normal force and plate separation distance were measured over a compression and relaxation of an individual microbead.

loaded, and porous beads) were determined using a rheometer (TA

Instruments DHR3) in parallel plate configuration to measure force *vs.* axial compression of individual microbeads. The data was then fit to Hertz's equation, 14,29

$$F = \frac{E * D^{\frac{1}{2}}}{3(1 - v^2)} (D - H)^{\frac{3}{2}}$$
(1)

and normalized by the microbead diameter,¹⁴ resulting in the following equation

$$\frac{F}{D^2} = \frac{E^*}{3} \left(\frac{\delta}{D}\right)^{\frac{3}{2}}$$
(2)

where F is the measured axial force, D is the uncompressed microbead diameter, δ is the compression distance, E is the bulk

modulus, ν is the Poisson's ratio, and E^* is the effective modulus defined as $E/(1-\nu^2)$. The schematic of the moduli measurement setup is shown in Fig. 3.

Results

In the first step, we studied the degree of porogen packing and efficacy of the selective porogen removal using SEM imaging. Fig. 4 shows monolayer, single particle, surface morphology, and cross-sectional images of the synthesized elastomeric microbeads. The mesoporous silica microspheres used in the microreactor are on the order of 50-100X smaller than the flamed-tip opening to mitigate any possible clogging of the outlet capillary. It was observed that the silica-loaded polymer stream to the catalyst stream flow rate ratios greater than 1:1, resulted in stable polymer jet formation and breakup with uniform porogen loading inside each droplet (see supplementary video M1). Using SEM imaging, it was determined that the produced monodispersed PHMS microbeads were uniformly loaded with close-packed silica microspheres, which were removed entirely upon etching with KOH (Fig. 4 Biii and Ciii). In the next step, we investigated the effect of varying the crosslinker length on the degree of porosity within the macroporous microbeads. As shown in Fig. 4C-D, switching from the hexadiene crosslinker to the divinyl PDMS crosslinker enhanced the pore interconnectivity and surface porosity of the resulting particles. Compared to the hexadiene crosslinker, the vinyl-terminated PDMS makes the polymer network stronger and less brittle due to the additional length



Fig. 4 SEM images of PHMS microbeads (A) without silica gels $(Q_1/Q_2/Q_3 = 5/10/200 \ \mu\text{L/min})$, (B) with 5 \mathbb{Z} m silica gels $(Q_1/Q_2/Q_3 = 10/10/300 \ \mu\text{L/min})$, (C) after selective etching of the silica particles $(Q_1/Q_2/Q_3 = 10/10/300 \ \mu\text{L/min})$, (D) and etched particles using a divinyl-terminated PDMS crosslinker $(Q_1/Q_2/Q_3 = 8/12/50 \ \mu\text{L/min})$. (I) Monolayer microbeads; (II) Single intact microbead; (III) Inside the elastomeric microbeads; (IV) Surface morphology of the microbeads.

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and flexibility offered by the PDMS chains. The molar density of V03 is much lower than the hexadiene crosslinker due to the high molecular weight (500 vs. 82 g/mol) and bulky PDMS chain. The stronger network allows improved survival of the thinwalled pores during the filtering and SEM plating process. The PHMS microscaffolds produced by our microfluidic method showed an excellent monodispersity (CV< 5%, see Fig. S2 and S3 in the supplementary information) compared to the polysiloxane microbeads synthesized using bulk emulsification techniques (20%<CV< 100%). The CV of the produced microbeads begins to dramatically increase (~12% with 20 µm silica) with the size of silica particles as they begin having a significant influence on the dynamics of droplet formation at the flamed-tip opening. As the silica-loaded droplet breaks off from the bulk fluid it necks down which will be more affected with larger silica particles present in the necking region.

Next, we studied the effect of porogen size on the degree of porosity of the resulting elastomeric macroporous microbeads. Varying the size of the mesoporous silica microspheres loaded into the droplets allows for size tuning of the resulting macropores within each PHMS microbead (Fig. 5). As the size of the silica microspheres increases, the size of the resulting macropores increases as well, leading to large voids and interconnected spaces. However, increased porogen size negatively affects the stability of droplet breakup in the microreactor and can lead to rapid separation of the colloidal silica suspension in the precursor syringe, resulting in heterogeneous flow of silica microspheres. Small perturbations in the jet breakup dynamics introduced by varying porogen size



Fig. 5. PHMS microbeads synthesized with (A) 5, (B) 7.5, (C) 10, and (D) 20 μ m sized mesoporous silica microspheres using 5:1 PHMS:V03. SEM images of (I) silica-loaded microbeads, (II) single macroporous microbead, and (III) surface morphology of the macroporous PHMS microbeads. Flowrates Q₁, Q₂, and Q₃ in μ L/min: (A) 4/12/250, (B) 8/14/150, (C) 5/10/100, and (D) 8/18/200.

can have a significant impact on the range of flow rates that



Fig. 6 Volume rendering of μ CT stacked images for (A) porogen-loaded and (B) macroporous microbeads. Insets show single microbead cross-sections with identical colormaps. Microbeads produced using $Q_1 = 10 \mu$ L/min, $Q_2 = 10$

result in monodisperse dripping. Minimizing the porogen size introduces smaller breakup disturbances, but the desired pore size for a given application must be considered.

 μ L/min, and Q_3 = 300 μ L/min with 5:1 PHMS:hexadiene ratio and 5 μ m silica.

In order to study the penetration depth of the etchant solution into the porogen-loaded elastomeric microbeads, highresolution µCT imaging was performed on dried porogenloaded (control) and macroporous microbeads. High-resolution µCT imaging provides information about the internal structure of PHMS microbeads without the need for destructive analysis by cutting into the microbeads. Fig. 6 and supplementary video M-2 show reconstructed µCT images of the tested PHMS microbeads. As seen in Fig. 6, there is a dramatic density difference between the porogen-loaded (*i.e.*, unetched, Fig. 6A) and macroporous (*i.e.*, etched, Fig. 6B) elastomeric microbeads. The lack of high density regions within the porous microbeads suggest that the etching process is successful in selective removal of the utilized solid porogen (silica), even from the core (center) of the PHMS microbeads. As shown in Fig. 6B, the etched microbeads are primarily comprised of low-density void spaces while the silica-loaded PHMS microbeads possess highdensity patches throughout the microbeads (Fig. 6B).

After confirmation of the complete selective removal of the solid porogen from the elastomeric microbeads using µCT imaging, we studied the effect of macropores on the effective modulus of the in-flow synthesized microbeads. The effective moduli of the synthesized microbeads were measured using the parallel plate test described above. A representative compression and relaxation data set is provided in the supplementary information (Fig. S4). Fig. 7A shows the measured force vs microbead displacement tests for porogenfree (control), porogen-loaded, and macroporous microbeads. Fig. 7B shows the calculated effective modulus of the microbeads from linear fits of force vs displacement graphs (Eq. 2) similar to Fig. 7A. As seen in Fig. 7B, the porogen-free (control) microbeads had the highest effective modulus, followed by the porogen-loaded and macroporous microbeads. In addition, increasing the crosslinker concentration resulted in

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a more tightly bound network and thus a higher effective modulus. Furthermore, increasing the crosslinker length and elasticity (*i.e.*, V03) resulted in a stronger and more durable network, while the shorter and stiffer crosslinker (*i.e.*, hexadiene) resulted in more brittle microbeads. Particle composition (crosslinker type and ratio to the polymer) was kept constant across each group (control, silica-loaded, and etched particles) with flow rate variations only maintaining the overall dripping regime.



Fig. 7 (A) Low compression normalized stress vs strain data for PHMS microbeads crosslinked using V03 crosslinker. (B) Effective elastic modulus for different crosslinker concentrations. A and B - Red: Non-porous microbeads (control); Green: silica-loaded PHMS microbeads; Blue: PHMS macroporous microbeads. 3:1 hexadiene microbeads produced using $Q_1/Q_2/Q_3 = 10/6/300$ (5 µm silica-loaded) and 9/18/120 (control). 5:1 hexadiene microbeads produced using $Q_1/Q_2/Q_3 = 10/10/300$ (5 µm silica-loaded) and 5/10/200 (control). 5:1 V03 microbeads produced using $Q_1/Q_2/Q_3 = 8/12/200$ (5 µm silica-loaded) and 12/36/300 (control). All flowrates are in µL/min.

Conclusions

In conclusion, we demonstrated the ability to continuously synthesize monodisperse elastomeric microbeads with tunable pore size and elastic modulus. The developed microfluidic platform enabled facile synthesis of porogen-loaded elastomeric microbeads with uniform loading and excellent size distribution. Increasing the size of mesoporous silica microspheres resulted in larger interconnected macropores within the elastomeric microbeads. In addition, increasing the length of the crosslinker caused a significant increase in the degree of porosity. We expect that the method presented herein will be readily extended to other linear siloxanes and functional crosslinkers with applications in areas such as adsorbents^{3,24} and enzyme immobilization⁴, catalysis², and tissue engineering^{6,7}. The developed microbead synthesis strategy could also be applied towards the synthesis of macroporous microbeads using other substrates that can only be crosslinked chemically rather than the conventional thermal or optical triggering techniques.

Conflicts of interest

The authors declare no conflicts of interest.

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