Harnessing liquid-in-liquid printing and micropatterned substrates to fabricate 3-dimensional all-liquid fluidic devices

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Systems comprised of immiscible liquids held in non-equilibrium shapes by the interfacial assembly and jamming of nanoparticle−polymer surfactants have significant potential to advance catalysis, chemical separations, energy storage and conversion. Spatially directing functionality within them and coupling processes in both phases remains a challenge. Here, we exploit nanoclay−polymer surfactant assemblies at an oil−water interface to produce a semi-permeable membrane between the liquids, and from them all-liquid fluidic devices with bespoke properties. Flow channels are fabricated using micropatterned 2D substrates and liquid-in-liquid 3D printing. The anionic walls of the device can be functionalized with cationic small molecules, enzymes, and colloidal nanocrystal catalysts. Multi-step chemical transformations can be conducted within the channels under flow, as can selective mass transport across the liquid−liquid interface for in-line separations. These all-liquid systems become automated using pumps, detectors, and control systems, revealing a latent ability for chemical logic and learning.
he ability to shape immiscible liquids into prescribed architectures and reconfigure them on-demand is an emerging design paradigm in soft-matter materials chemistry\textsuperscript{1–9}. To trap a liquid indefinitely in a nonequilibrium shape within another liquid, a suitable elastic film must be assembled at the liquid–liquid interface\textsuperscript{3}. Assemblies of nanoparticle–polymer surfactants (NPSs), composed of colloidal nanoparticles originating from one of the liquids and polymer surfactants from the other, offer a compelling means both to assemble this elastic film and impart nanoparticle-derived functionality upon an all-liquid system\textsuperscript{3}. NPS assembly at the interface leverages complementary chemical functionality at the surface of the nanoparticle and at the polymer surfactant chain end(s). This scheme is versatile, allowing the fabrication of reconfigurable systems of discrete liquid droplets\textsuperscript{10,11}, long liquid threads\textsuperscript{6,12}, molded objects\textsuperscript{13}, and bicontinuous interfacially jammed emulsions\textsuperscript{2}. Still nascent in their development, structured liquids presently lack clear design rules for controlling their 2D or 3D architectures, spatially directing functional components within each liquid phase, and coupling physicochemical processes across the liquid–liquid interface so as to create autonomous chemical systems capable of performing useful work, processing information, or executing logical functions.

Here, we harness the structured liquid paradigm to fabricate 3D all-liquid fluidic devices that are infinitely reconfigurable and endowed with spatially programmable functions. In turn, we gain insights into their potential for rendering chemical systems of arbitrary complexity to perform tasks, including chemical separations, multi-step chemical transformations, and chemical logic.

Results

All-liquid fluidic device fabrication. To fabricate all-liquid fluidic devices, glass supports are first coated with a superhydrophobic polymer\textsuperscript{7} before undergoing photo-patterning with a variety of superhydrophilic channel architectures (Fig. 1a). A polar phase—e.g., an aqueous dispersion of anionic 2D nanoclays (10 mg mL\textsuperscript{−1} at pH 7.0)—is then deposited onto the superhydrophilic region prior to immersion in an oil phase—e.g., a solution of homobiecholeic amine-terminated poly(dimethylsiloxane) (H\textsubscript{2}N-PDMS-NH\textsubscript{2}, M\textsubscript{w} \sim 27 kg mol\textsuperscript{−1}, 10% w/w) in either toluene, silicone oil, or dodecane (Supplementary Fig. 1). Interfacial forces are effective in pinning and confining the aqueous phase in arbitrarily complex geometries and a wide range of channel widths (635–3000 \textmu m) (Supplementary Fig. 2).

Once the liquid phases are in place on the patterned glass support, NPS film formation at the water–oil interface is extremely rapid\textsuperscript{11,14,15}. Its kinetic trajectory was monitored via time-evolution of interfacial tension (IFT) for pendant droplets of 2D nanoclay dispersions (0.5–10 mg mL\textsuperscript{−1}) within an oil phase consisting of H\textsubscript{2}N-PDMS-NH\textsubscript{2} (10% w/w) in toluene (Fig. 1b). The concentration of nanoclay influences the rate of diffusion to the interface, the ionic strength of aqueous phase and the areal density at the interface at steady-state (Supplementary Fig. 3a); the latter two ultimately dictate the IFT\textsuperscript{16,17}. In all cases, the system reached steady-state rapidly. Furthermore, NPS assembly into an elastic film upon interface compression was irreversible, as evidenced by the film’s wrinkling behavior upon droplet retraction (Fig. 1c, Supplementary Fig. 3b and 4, and Supplementary Movie 1). The topography of the NPS film at the water–oil interface was revealed using in-situ atomic force microscopy (AFM)\textsuperscript{16} (Fig. 1d). Here, we immersed the AFM tip in the aqueous phase and brought it into contact with the water–oil interface, where a tiny drop of silicon oil (viscosity \(\eta = 60,000\) cSt) with 10% w/w of H\textsubscript{2}N-PDMS-NH\textsubscript{2} had been pinned on the silicon substrate and subsequently enshrouded by a dispersion of 2D nanoclays (10 mg mL\textsuperscript{−1}). The data showed well-packed nanoclays at the liquid–liquid interface: no structural aberrations were visible.

NPS films exhibit the necessary mechanical strength and elasticity to render patterned all-liquid microchannels shape-persistent under high flow rates, even at sharp curves defined by superhydrophilic surface patterns: e.g., in U-bends of serpentine channels (Fig. 1). 2D nanoclay particles are held strongly to the interface by the surfactant. Incomplete packing of nanoclays yields pores that expose surfactant capable of binding a second nanoclay layer on top of the first (Fig. 1d). These nanoclay–nanoclay interactions ultimately determine the mechanical properties of the NPS films for small displacements both compressive and extensive. To quantify the elasticity of NPS films, we compressed them uniaxially in a planar geometry in a Langmuir–Blodgett trough (Supplementary Fig. 5). The wavelength of the wrinkles, \(\lambda\), is directly related to the bending modulus, \(B\), by \(\lambda = \frac{2t}{B} \left( \frac{1}{\pi} \right)\)\textsuperscript{18,19}, where the density difference between oil and water is \(\Delta \rho = 70\) kg m\textsuperscript{−3}, and \(g = 9.8\) m s\textsuperscript{−2}. \(\lambda\) was obtained by image analysis of 2748 wrinkles to be \(\lambda = 113 \pm 22\) \textmu m, giving a bending modulus on the order of 10\textsuperscript{4} k\textsubscript{B}T for the NPS film—at least two orders of magnitude larger than that typically measured in nanoparticle assemblies at a liquid–liquid interface\textsuperscript{20,21}. To demonstrate the value these elastic properties bring to all-liquid fluidic devices, we pumped an aqueous solution of resazurin dye (50 \textmu g mL\textsuperscript{−1}, pH 7.0) through a channel \(\sim 139\) mm in length and \(\sim 2\) mm in width at a flow rate of 10 mL h\textsuperscript{−1}, significantly faster than the maximum reported in liquid microfluidics that lack a functional interfacial membrane (Fig. 1e, f). Only patterned microchannels bearing NPS walls were capable of guiding flow along the channels at these rates without aberrations to the all-liquid architectures; absent these elastic NPS walls, the aqueous phase irreconcilably accumulates a volume at the entry point of the channel (Supplementary Movie 2). Thus, while all-liquid fluidics have, in the past, made use of surface patterning and surface adhesion to produce complex all-liquid structures\textsuperscript{22–25}, these schemes relied on surface tension and contact angle hysteresis\textsuperscript{26,27} for pumping one fluid within the other, passively or actively. This set limits for the types of fluids and the flow rates that can be used, and restricted the chemistries that can be loaded into each phase. In our scheme, rather than flow being driven by Laplace pressure, contact angle hysteresis or electric field strength\textsuperscript{28}, the internal overpressure under flow is balanced by the elastic NPS wall. As a result, all-liquid fluidic systems may now be used at high flow rates (up to 10 mL h\textsuperscript{−1}) at markedly reduced surface tensions and, notably, in the presence of surface-active species. The maximum flow rate of all-liquid fluidic devices will of course depend both on the cross-sectional area of the channel as well as the channel architecture, and can be explored endlessly as such.

Membrane permeability. The irregular shapes of nanoclays yield NPS films with residual interstitial spaces, if the assemblies are confined to 2D, which are selectively permeable to solutes in either liquid phase\textsuperscript{29}. Overlapping nanoclays may also contribute to the film’s observed permeability. We demonstrated selective mass transfer across the interface (Fig. 2a) by carrying out crossover experiments using a charge-neutral fluorescent dye (dye 1, Fig. 2b and Supplementary Fig. 6) that has an affinity for both water and toluene, and compared the crossover rate with those of anionic resazurin (2) and cationic rhodamine 6G (3). We introduced a solution of H\textsubscript{2}N-PDMS-NH\textsubscript{2} in silicone oil
(10% w/w) into a quartz cuvette containing a dispersion of nanoclays (9 mg mL$^{-1}$) and dye (100 µg mL$^{-1}$) and let the system form an NPS interfacial film; owing to the insolubility of dyes in silicone oil, these reporters remain barricaded inside the aqueous phase. We then added toluene to the oil phase and observed the rate at which dye partitioned into the toluene phase by monitoring the optical absorbance of the aqueous phase (Fig. 2b, Supplementary Fig. 7 and 8); the decrease in optical absorption indicated successful partitioning of the dye into the organic phase. In contrast, neither of the ionic dyes partitioned into the organic phase (Supplementary Fig. 7). The partitioning of non-ionic dye was further confirmed in the patterned all-liquid device using laser scanning confocal microscopy (LSCM) (Fig. 2c and Supplementary Fig. 9); our success in this format suggested that the semi-permeable NPS film could be utilized for in-line chemical separations under flow. To demonstrate this feature, a solution of dyes 1 and 2 was pumped through the channel (0.2 mL h$^{-1}$), which resulted in a color change after ~1 h under continuous flow, both within the aqueous channel as well as in the overlaid toluene (Fig. 2d and Supplementary Movie 3). More than 93% of dye 1 was separated from the aqueous dye mixture by liquid–liquid partitioning into the toluene phase; dye 2 was retained in channel (Fig. 2e and Supplementary Fig. 10).

**Membrane functionalization.** Beyond their use as structural elements, anionic nanoclays further allow cationic molecules, enzymes, and nanoparticles to be anchored to the NPS wall (Fig. 3a–c). In turn, all-liquid fluidic devices are primed for biphasic chemical synthesis. We were initially drawn to this possibility by noting changes in the optical absorption of rhodamine 6G in the presence of nanoclays during the crossover experiments: the molar extinction spectrum of cationic dyes changed over time, consistent with adsorption to the anionic nanoclays (Supplementary Fig. 7), whereas no such changes were observed for anionic or non-ionic dyes. We confirmed that
cationic small molecules were immobilized on the NPS walls by using LSCM to visualize the spatial distribution of methylene blue (Fig. 3d and Supplementary Fig. 11) and rhodamine 6G (Supplementary Fig. 12) in the device. After pumping cationic dyes into the channel, aging the system such that they diffuse to the interface, and rinsing with water to remove unbound objects, the immobilized fluorophores each tracked the curvature of the NPS wall. We were similarly able to anchor to the wall cationic enzymes (e.g., horseradish peroxidase, HRP) as well as cationic nanocrystal catalysts (e.g., cetyltrimethylammonium bromide (CTAB)-modified Pt nanocrystals, \( \zeta = +19.2 \pm 1.3 \text{ mV} \)). While the fluorescence of adsorbed dye-labeled HRP (lissamine rhodamine B sulfonyl chloride (LRSC)-labeled HRP) was readily visualized using LSCM (Fig. 3e and Supplementary Fig. 13), the immobilization of CTAB-Pt nanocrystals with respect to the underlying nanoclay tiles at the liquid–liquid interface required in-situ AFM to properly visualize nanocrystal density (Fig. 3f, and Supplementary Fig. 14 and 15). Notably, we did not observe any nanocrystal mobility at the NPS surface in the in-situ AFM experiments, even upon repeated imaging over a fixed area; the nanocrystals are well-anchored to the nanoclays.

**All-liquid microreactor.** We carried out flow-driven multistep chemical transformations using all-liquid fluidic devices configured with HRP@NPS walls (Supplementary Fig. 16 and 17). In a first demonstration, we pumped a solution of 3,3',5,5'-tetramethylbenzidine (TMB) and hydrogen peroxide (Fig. 3g) through...
the channel (1 mL h\(^{-1}\)), which led to the oxidation of TMB to yield a yellow-colored diimine product\(^{30}\) (Supplementary Fig. 18). We visually tracked the trajectory of this reaction along the channel length, as the substrate, intermediate, and product are chromogenically distinct. In a second demonstration, we pumped a solution of phenol, 4-aminoantipyrine (4-AAP) and hydrogen peroxide (Fig. 3h) through the channel (1 mL h\(^{-1}\)), which led to the oxidative coupling of phenol and 4-AAP to yield a red-colored quinone-imine conjugate\(^{31}\). While this second sequence is more complex, the kinetics were fast, likewise allowing the reaction’s trajectory to be visualized in the channel via the intense color of the final product. Given our successes with these reactions, we next implemented an expanded set of chromogenic oxidative coupling reactions to demonstrate how the spatial programmability of function should be coupled to active programming and re-programming of the flow paths of the device to create a type of chemical logic.

**Programmable, reconfigurable, and self-healing microreactors.** An advantage to using NPS films to stabilize all-liquid fluidic devices is that additional microchannels are straightforward to introduce via direct-write methods (e.g., 3D liquid-in-liquid printing), even after the device has been configured and its local chemistry defined by the user (Fig. 4). To contextualize this advance, without the aid of micropatterned substrates and NPS films, constructing channel-like aqueous threads in oil is not typically possible, as the thermodynamic driving force required to reduce interfacial area breaks up aqueous threads into droplets\(^{32}\). In that wall-forming is fast, and the jammed assembly counteracts in a timely manner the compressive force in the system arising from Plateau–Raleigh instabilities\(^{6,12,33}\) both patterned 2D channels and 3D printed inter-channel microtubes were stable for at least 2 h, i.e., while devices were in use. The stability of our printed microchannels allows them to be used as bridges between different regions on the substrate, or to couple the device to an exogeneous entity (Fig. 4a–d).

We found that despite the perturbative nature of anchoring, printing, and bridging liquid microchannels, NPS films self-heal on a time-scale of ms-to-s (Supplementary Fig. 19). This allows spatially programmed functionality to be connected in a user-defined sequence in real time. Printed channels retained their structure during flow, as demonstrated by the pumping of aqueous blue dye through three liquid regions connected by two...
Fig. 4 Reconfigurable 3D all-liquid microreactors with logical optical outputs. a Schematic of fabricating 3D microchannels stabilized by nanoclaysurfacant interfacial assembly to connect the separated microchannels by printing of water in oil. Disconnecting these microchannels is also possible by cutting these “bridges” off. b Adjacent channels on a patterned substrate connected by a 3D printed overpass. Na-fluorescein (0.1 mg mL$^{-1}$) has been added to the clay dispersion to clarify features of the constructs. Scale bar: 2 mm. c Fabrication of a 3D bridge linking separate channels (middle panel) and clipping the bridge (right panel). Scale bar: 500 µm. d Time-lapse showing a dyed solution pumped through liquid channels CH 1, CH 2, and CH 3, which are connected by printed 3D bridges. Scale bar: 5 mm. e Chemical transformations yielding chromogenically distinct products are conducted consecutively by forming and then breaking bridges between the inlet and various patterned channels containing different reagents. The bridges are built using liquid-in-liquid 3D printing to yield a “turn-on” state, and mechanically broken to yield a “turn-off.” Inset photographs on top show the solutions collected in capillaries after carrying out these chromogenic reactions in vials. DAOS: N-ethyl-N-(2-hydroxy-3-sulfopropyl)-3-methylaniline, sodium salt. TOOS: N-ethyl-N-(2-hydroxy-3-sulfolpropyl)-3-methylaniline, sodium salt. Scale bars: 1 mm (top) and 5 mm (bottom)

Discussion

The ability to spatially program and re-program on-demand the functionality and geometric layout of a fluidic device, including channel compartmentalization, flow paths between channels, and extent of chemical coupling between the immiscible fluids comprising the device opens exciting directions in responsive soft-matter chemical systems. Our studies uncover a latent learning ability in such devices, in that physiochemical sensing or detection of channel properties and contents can be used to direct the architecture of the device to achieve a specific outcome. Maturation of the design concept led to devices that can execute complex tasks in a logical manner by reversible compartmentalization of function and direction of chemo-energetic flows that operate far from equilibrium conditions$^{34–37}$. The potential for this system to exhibit autonomous learning is evident. Such devices may also be arrayed to generate deep or dark data$^{38}$ for machine learning, e.g., from all-liquid (bio)chemical transformations and screens, to build knowledge and understanding from chemical logic.

Methods

Fabrication of photopatterned glass substrates. We employed a previously published protocol$^7$ with slight modifications (Supplementary Figure 1). Briefly, 25 µm-thin strips of Teflon film (American Durafile Co.) were placed at the edges of one 3-(trimethoxysilyl)propyl methacrylate-modified glass plate (76.2 × 25.4 × 1 mm, VWR Microscope Slides) and one fluorinated glass slide was clamped on top of it. A polymerization mixture consisting of 2-hydroxyethyl methacrylate (24% w/w), ethylene dimethacrylate (16% w/w), 1-decanol (12% w/w), cyclohexanone (48% w/w) and 2,2-dimethoxy-2-phenylacetophenone (photoinitiator, 1% w/w with respect to monomers) was injected into the mold between the glass slides and irradiated for 15 min with 12.0 mW cm$^{-2}$ UV light (260 nm). After
removing the glass cover, the newly formed polymer layer was washed extensively with ethanol. The superficial surface was removed by using an adhesive film ( "Scotch tape" ). The polymer layer was then immersed into dichloromethane (50 mL containing 4- pentyenoic acid (112 mg, 1.14 mmol) and 4-(dimethylamino)pyridine (DMAP, 56 mg, 0.46 mmol) and N,N'-disopropylcarbodiimide (176.5 µL, 1.14 mmol), followed by stirring the solution at room temperature for 4 h. The plate was washed extensively with acetone and dried using a stream of nitrogen.

Surface patterning was performed using thiol-yne reactions. The polymer layer was wet with ethyl acetate solution containing 10 vol% 1 H,1 H,2 H,2H-perfluorodecanethiol, covered by a photomask, and irradiated by UV light (260 nm) for 30 s. After removing the photomask, washing with acetone and drying, the polymer layer was wet with an ethanol−water mixture (1:1 v/v) containing 2-mercaptoethanol (15% v/v) and 2,2-dimethoxy-2-ethoxyacetophenone (1% w/v). Finally, the polymer layer was irradiated by UV light (365 nm, 2 mW cm−2) for 120 s, and washed extensively with ethanol followed by drying. Static water contact angles (WCAs) were measured after each modification (Supplementary Fig. 1b).

Modification of the surface with 1 H,1 H,2 H,2H-perfluorodecanethiol endows the polymer surface with superhydrophobicity (static WCA being as high as 163.1°), as functionalization of the surface with 2-mercaptoethanol transforms the polymer into a superhydrophilic surface (static WCA = 48°). Due to the extreme difference in wettability of the superhydrophilic area compared to the superhydrophobic barriers, water is spontaneously repelled by the barriers, but fills the superhydrophilic regions.

**All-liquid device fabrication.** An appropriate volume of the nanoclay dispersion (10 mg mL−1) was applied to each superhydrophilic area to in-fill the pattern’s contour. The entire surface was then covered with a solution of H2N-PDMS-NH2 and DMAP (56 mg, 0.46 mmol) and 4-(dimethylamino)pyridine (50 mL containing 12% w/w) in the cuvette (Supplementary Fig. 7a) was monitored using the DSA Advance software after gentle adding 1.5 mL toluene solution of NH2-PDMS-NH2 (10% w/w) in the cuvette (Supplementary Fig. 7a). Adsorption of dye 1 (1 mg mL−1) in a dispersion of nanoclays (10 mg mL−1) was introduced to a microchannel bearing an NPS interferential film. The channel was interrogated by LSCM and the field of view set at the edge of the channel. Fluorescence (excitation, 405 nm; detection range, 510–680 nm) from the dye showed that dye 1 is fully three orders of magnitude larger than the value obtained by direct measurements on nanoparticle assemblies20,21. The 2D Young’s modulus, Y2D = Y2D of 5 N m−1 is fully three orders of magnitude larger than the water−oil IFT as measured in Fig. 1b. These results are remarkable for two reasons. First, our ultra-thin NPS assembly has bending moduli orders of magnitude larger than those measured previously in the literature19,40 or predicted by theory21. We attribute this to both the overlapping structure of the nanoassembly and the steric interactions between the bound polymer brushes that reinforce the film. Second, the large Young’s modulus of the film underscores the enhanced structural robustness and higher flow rates that the film allows us to achieve in comparison to methods in which pumping and structure are derived solely from surface forces.

**Synthesis of amphiobal dye 1.** The synthesis of dye 1 (Supplementary Fig. 6a) was carried out over two steps. Briefly, to a dichloromethane solution (20 mL) of cyanoacetic acid (10 mmol), triethylene glycol monomethyl ether (10 mmol), and DMAP (1 mmol), a solution of dicyclohexylcarbodiimide (10 mmol) in dichloromethane (10 mL) was added dropwise at 0 °C. The mixture was allowed to warm to room temperature and was stirred for 2 h. After filtering the solid byproduct in the mixture under gravity, the filtrate was dried using MgSO4, concentrated and purified by column chromatography (silica, 10–80% ethyl acetate in hexane) to obtain a pure 2-(2-(methoxyethoxy)ethoxy)ethyl 2-cyanoacetate (71% yield). Next, to a solution of 4-(dimethylamino)benzaldehyde (3.35 mmol) and 2-(2-(methoxyethoxy)ethoxy)ethyl 2-cyanoacetate (3.35 mmol) in THF (7.5 mL) was added 1,8-diazabicyclo[5.4.0]undec-7-ene (DBU) (3.40 mmol) in THF (0.5 mL). The solution was concentrated under vacuum after stirring for 1 h at room temperature. Dye 1 was purified via silica gel chromatography (10–100% ethyl acetate in hexanes) (68%) yield.1 H NMR (500 MHz, CDCl3) δ 8.08 (s, 1H), 7.95 (d, 2H, J = 9.0 Hz), 6.71 (d, 2H, J = 9.1 Hz), 4.43 (m, 2H), 3.83–3.81 (m, 2H), 3.73–3.75 (m, 2H), 3.70–3.67 (m, 4H), 3.58–3.56 (m, 2H), 3.32 (s, 3H), 13.12 (s, 6H) ppm. These data were consistent with a previous report41. Dye 1 dissolves in water and in toluene. This molecular rotor shows differentiated fluorescence in water and in toluene (Supplementary Fig. 6b).

**Permeability of NPS films at the liquid−liquid interface.** Mass transfer across the NPS assembly intercuvette in a cuvette (Supplementary Fig. 7a) was monitored using 5 µL of dye 1, resazurin (dye 2), or rhodamine 6G (dye 3) (each 1 mg mL−1 in water). The dyes were combined with 45 µL nanoclay dispersion (10 mg mL−1, pH 7) in the cuvette, and then covered with 50 µL solution of H2N-PDMS-NH2 nanoclay dispersion (10% w/w) in the cuvette (pH 7). After 15 min of dye 1 was added (1.4 mg mL−1) in a nanoclay dispersion (10 mg mL−1) was introduced to a microchannel bearing an NPS interferential film. The channel was interrogated by LSCM and the field of view set at the edge of the channel. Fluorescence (excitation, 405 nm; detection range, 510–680 nm) from the dye showed that dye 1 is fully three orders of magnitude larger than the value obtained by direct measurements on nanoparticle assemblies20,21. The 2D Young’s modulus, Y2D = Y2D of 5 N m−1 is fully three orders of magnitude larger than the water−oil IFT as measured in Fig. 1b. These results are remarkable for two reasons. First, our ultra-thin NPS assembly has bending moduli orders of magnitude larger than those measured previously in the literature19,40 or predicted by theory21. We attribute this to both the overlapping structure of the nanoassembly and the steric interactions between the bound polymer brushes that reinforce the film. Second, the large Young’s modulus of the film underscores the enhanced structural robustness and higher flow rates that the film allows us to achieve in comparison to methods in which pumping and structure are derived solely from surface forces.
remove the unreacted dye. LRSC-labeled HRP showed an emission maximum of 395 nm in 2-(N-morpholino)ethanesulfonic acid buffer (MES, 10 mM, pH 6.5) after excitation at 370 nm (Supplementary Fig. 13c).

To adsorb the enzyme onto the NPS wall, 100 µg mL⁻¹ of LRSC-labeled HRP in MES buffer (pH 6.5) was infused through the liquid channel at a flow rate of 0.15 mL h⁻¹ for 12 min. After aging for 60 min, the channel was washed by introducing pure MES buffer for 30 min at a flow rate of 0.5 mL h⁻¹ (Supplementary Fig. 13d). The distribution of HRP on the wall of the device was visualized using LSCM (Fig. 3e).

All-liquid fluidic devices with wall-bound cationic nanorods. CTAB-coated platinum nanocrystals (Pt NCs) were prepared according to a reported method with minor modifications. Briefly, an aqueous solution of K₂PtCl₄ (1 mM, 10 mM) was mixed with an aqueous solution of CTAB (8.4 mL, 19 mM) in a 20 mL vial at room temperature. The mixture was then heated at 50 °C until the solution became clear, after which an ice-cold aqueous solution of NaN₃H₂O (0.6 mL, 50 mM) was added, and the reaction mixture stirred for 6 h at 50 °C. Excess CTAB was removed by centrifuging the solution twice at 14,000 × g for 30 min. The precipitate was collected and re-dispersed in 50 mL water. The size and size distribution of the NCs was observed using a Zeiss Gemini Ultra-55 analytical Field Emission Scanning Electron Microscope (Supplementary Fig. 14). A Malvern Zetasizer Nano instrument was used to measure the zeta potential.

Cationic CTAB-coated Pt NCs were immobilized onto the anionic NPS assembly and visualized directly at the liquid–liquid interface using an in-situ AFM technique (Fig. 3f). A minimal drop of silicon oil (viscosity η = 60,000 cSt) with 10% w/w of H₂N-PDMS-NH₂ was placed on a silicon substrate and 30 µL of NPS dispersion and substrate were added, allowing the NCs to adsorb the anionic NPS assembly through complementary electrostatics (Supplementary Fig. 15a). While imaging, the AFM tip was immersed in the NCs dispersion and brought directly into contact with the water–oil interface (Supplementary Fig. 15b, c).

All-liquid microreactors-on-a-chip. Using an all-liquid fluidic device configured with a glass substrate (viscosity η = 10 cSt) with 10% w/w of H₂N-PDMS-NH₂ was placed on a silicon substrate and 30 µL of pure MES buffer was pumped through the channel at a flow rate of 1 mL h⁻¹ for 10 min (Supplementary Fig. 16). After 15 min aging for enzyme immobilization, pure MES buffer was pumped through the channel at a flow rate of 1 mL h⁻¹ for 10 min to remove the free HRP. TMB (10 times diluted from commercial TMB from Sigma, pH 6.5) or 4-AP/AAP periphery and substrates (5 mM 4-AP, 25 mM phenol, and 1 mM H₂O₂ in MES buffer, pH 6.5) was introduced. Two separate needles were connected to the inlet of the channel to avoid enzyme and substrate contamination. Conversion of substrates into colored products were observed along the length of the channel (Fig. 3g), which tracked color changes monitored exogenously in a cuvette (Supplementary Figs. 17, 18).

Spatially programmable all-liquid microreactors for chemical logic. Micro-channels linking otherwise isolated regions on the patterned glass substrates were created using a wet-etching technique (Fig. 3h). A minimal drop of silicon oil (viscosity η = 10 cSt) with 10% w/w of H₂N-PDMS-NH₂ was placed on a silicon substrate and 30 µL of nanoclay dispersion (10 mg mL⁻¹ in water) was added on top of the silicone oil to cover the oil completely. The sample was placed in a Cypher ES-sealed sample cell for 30 min, which allowed the nanoclay and polymer surfactants to assemble at the water–oil interface. Excess nanoclay was removed by exchanging the aqueous phase with water twice, after which the CTAB-coated Pt NC dispersion was added, allowing the NCs to adsorb the anionic NPS assembly through complementary electrostatics (Supplementary Fig. 15a). While imaging, the AFM tip was immersed in the NCs dispersion and brought directly into contact with the water–oil interface (Supplementary Fig. 15b, c).

channel to turn-on the route, the 4-AAP/H₂O₂ solution was injected into the inlets at a flow rate of 0.15 mL h⁻¹ to trigger the chromogenic reactions to generate products with different colors under flow (Supplementary Fig. 25c). Bridges were cut off to prevent chemical contamination after finishing a desired reaction in one of the 2D channels. The conversion of substrates into various colored products in CHs 1–3 was consistent with expected color changes, e.g., when reactions were conducted in a cuvette (Fig. 4e). No color change was observed in CH4, where HRP was absent.

Data availability
All data supporting the key findings of this study are available within the article and its Supplementary Information files or from the corresponding author upon reasonable request.

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References
1. Liu, A. J. & Nagel, S. R. Jamming is not just cool any more. Nature 396, 21–22 (1998).
2. Balazs, A. C., Emrick, T. & Russell, T. P. Nanoparticle polymer composites: where two small worlds meet. Science 314, 1107–1110 (2006).
3. Huang, C. et al. Bicontinuous macroporous materials from Bijel templates. Adv. Mater. 22, 4836–4841 (2010).
4. Cui, M., Emrick, T. & Russell, T. P. Stabilizing liquid drops in nonequilibrium shape by the interfacial jamming of nanoparticles. Science 342, 460–463 (2013).
5. Huang, C. et al. Bicontinuous structured liquids with sub-micrometre domains using nanoparticle surfactants. Nat. Nanotechnol. 12, 1060–1063 (2017).
6. Forth, J. et al. Reconfigurable printed liquids. Adv. Mater. 30, 1707603 (2018).
7. Feng, W. et al. Surface patterning via Thiol–Yne click chemistry: an extremely fast and versatile approach to superhydrophilic-superhydrophobic micropatterns. Adv. Mater. Interfaces 1, 1400269 (2014).
8. Yang, Z., Wei, J., Sobolev, Y. I. & Grzybowski, B. A. Systems of mechanized and reactive droplets powered by multi-responsive surfactants. Nature 553, 313–318 (2018).
9. Zarzar, L. D. et al. Dynamically reconfigurable complex emulsions via tunable interfacial tensions. Nature 518, 520–524 (2015).
10. Toor, A., Lamb, S., Helms, B. A. & Russell, T. P. Reconfigurable microfluidic droplets stabilized by nanoparticle surfactants. ACS Nano 12, 2563–2572 (2018).
11. Huang, C. et al. Structured liquids with pH-triggered reconfigurability. Adv. Mater. 28, 6612–6618 (2016).
12. Liu, X. et al. Liquid tube formation and stabilization using cellulose nanocrystal surfactants. Angew. Chem. Int. Ed. 56, 12594–12598 (2017).
13. Shi, S. et al. Liquid letters. Adv. Mater. 30, 1705800 (2018).
14. Forth, J., Skaff, H., Emrick, T., Dinsmore, A. D. & Russell, T. P. Nanoparticle assembly and transport at liquid–liquid interfaces. Science 229, 226–229 (2003).
15. Montelongo, T. et al. Electrotunable nanoplasmonic liquid mirror. Nat. Mater. 16, 1127–1133 (2017).
16. Chai, Y., Lukito, A., Jiang, Y., Ashby, P. D. & Russell, T. P. Fine-tuning nanoparticle packing at water–oil interfaces using ionic strength. Nano Lett. 17, 6453–6457 (2017).
17. Jiménez-Ángelés, F. & Firoozabadi, A. Tunable substrate wettability by thin water layer. J. Phys. Chem. C 120, 24688–24696 (2016).
18. Cerda, E., Ravi-Chandar, K. & Mahadevan, L. Thin films: wrinkling of an elastic sheet under tension. Nature 419, 579–580 (2002).
19. Cerda, E. & Mahadevan, L. Geometry and physics of wrinkling. Phys. Rev. Lett. 90, 074302 (2003).
20. Leahy, B. D. et al. Geometric stability and elastic response of a supported nanoparticle film. Phys. Rev. Lett. 105, 058301 (2010).
21. Mueggenburg, K. E., Lin, X.-M., Goldsmith, R. H. & Jaeger, H. M. Elastic membranes of close-packed nanoparticle arrays. Nat. Mater. 6, 656–660 (2007).
22. Walsh, E. J. et al. Microfluidics with fluid walls. Nat. Commun. 8, 816 (2017).
23. Zhao, B., Moore, J. S. & Beebe, D. J. Surface-directed liquid flow inside microchannels. Science 291, 1023–1026 (2001).
24. Atencio, J. & Beebe, D. J. Controlled microfluidic interfaces. Nature 437, 648–655 (2005).
25. Casavant, B. P. et al. Suspended microfluidics. Proc. Natl. Acad. Sci. USA 110, 10111–10116 (2013).
26. Walker, G. M. & Beebe, D. J. A passive pumping method for microfluidic devices. *Lab Chip* 2, 131–134 (2002).
27. Hong, L. & Pan, T. Three-dimensional surface microfluidics enabled by spatiotemporal control of elastic fluidic interface. *Lab Chip* 10, 3271–3276 (2010).
28. Tokarz, M. et al. Single-file electrophoretic transport and counting of individual DNA molecules in surfactant nanotubes. *Proc. Natl. Acad. Sci. USA* 102, 9127–9132 (2005).
29. Dinsmore, A. D. et al. Colloidosomes: selectively permeable capsules composed of colloidal particles. *Science* 298, 1006–1009 (2002).
30. Rye, D. B., Saper, C. B. & Wainer, B. H. Stabilization of the tetramethylbenzidine (TMB) reaction product: application for retrograde and anterograde tracing, and combination with immunohistochemistry. *J. Histochem. Cytochem.* 32, 1145–1153 (1984).
31. Allain, C. C., Poon, L. S., Chan, C. S. G., Richmond, W. & Fu, P. C. Enzymatic determination of total serum cholesterol. *Clin. Chem.* 20, 470–475 (1974).
32. Powers, T. R., Zhang, D., Goldstein, R. E. & Stone, H. A. Propagation of a topological transition: the Rayleigh instability. *Phys. Fluids* 10, 1052–1057 (1998).
33. Toor, A., Helms, B. A. & Russell, T. P. Effect of nanoparticle surfactants on the breakup of free-falling water jets during continuous processing of reconfigurable structured liquid droplets. *Nano Lett.* 17, 3119–3125 (2017).
34. Henson, A. B., Gromski, P. S. & Cronin, L. Designing algorithms to aid discovery by chemical robots. *ACS Cent. Sci.* 4, 793–804 (2018).
35. Greenaway, R. L. et al. High-throughput discovery of organic cages and catenanes using computational screening fused with robotic synthesis. *Nat. Commun.* 9, 2849 (2018).
36. Aspuru-Guzik, A., Lindh, R. & Reiher, M. The matter simulation (re)evolution. *ACS Cent. Sci.* 4, 144–152 (2018).
37. Steiner, S. et al. Organic synthesis in a modular robotic system driven by a chemical programming language. *Science* 363, eaax2211 (2019).
38. Raccuglia, P. et al. Machine-learning-assisted materials discovery using failed experiments. *Nature* 533, 73–76 (2016).
39. Knoche, S. et al. Elastometry of deflated capsules: elastic moduli from shape and wrinkle analysis. *Langmuir* 29, 12463–12471 (2013).
40. Vella, D., Aussillous, P. & Mahadevan, L. Elasticity of an interfacial particle raft. *Europhys. Lett.* 68, 212–218 (2004).
41. Sutharsan, J. et al. Rational design of amyloid binding agents based on the molecular rotor motif. *ChemMedChem* 5, 56–60 (2010).
42. Zhou, Y., Hu, N., Zeng, Y. & Rusling, J. F. Heme protein—clay films: direct electrochemistry and electrochemical catalysis. *Langmuir* 18, 211–219 (2002). 
43. Yamazaki, Y. et al. Nanocrystal bilayer for tandem catalysis. *Nat. Chem.* 3, 372–376 (2011).

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**Author contributions**

B.A.H. and W.F. designed and B.A.H. directed the experiments. W.F. carried out pendant drop tensiometry, mass transfer experiments, LSCM, and all-liquid device fabrication and testing. Y.C. and P.D.A. carried out in-situ AFM experiments. J.F. and T.P.R. carried out mechanical characterization of NPS films and liquid-in-liquid 3D printing. W.F., J.F., T.P.R. and B.A.H. wrote the manuscript with contributions from all co-authors.

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