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A new droplet-forming fluidic junction for the generation of highly compartmentalised capsules[†]

A new oscillatory microfluidic junction is described, which enables the consistent formation of highly uniform and complex double emulsions, and is demonstrated for the encapsulation of four different reagents within the inner droplets (called cores) of the double emulsion droplets. Once the double emulsion droplets had attained a spherical form, the cores assumed specific 3D arrangements, the orchestration of which

appeared to depend upon the specific emulsion morphology. Such double emulsion droplets were used as

templates to produce highly compartmentalised microcapsules and multisomes. Based on these construct models, we numerically demonstrated a model chemical reaction sequence between and within the liquid

cores. This work could provide a platform to perform space/time-dependent applications, such as

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programmed experiments, synthesis, and delivery systems.

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Introduction

Droplet microfluidics¹ is usually based on liquid droplet formation in another immiscible phase, and can provide large numbers of uniform, independent, and controllable droplets.^{2,3} Such droplets have been widely applied to diverse chemistries,⁴ for instance, in d-PCR,⁵ organic synthesis,⁶ the fabrication of microparticles,7 and various cell manipulations.^{8,9} In addition, droplet microfluidics shows outstanding performance in the generation of consistent multiple emulsions,¹⁰ when compared to conventional batch methodologies, though the volumes produced are often far lower. Complex multiple emulsion droplets¹¹ can be continuously formed and processed,^{12,13} within modular or monolithically integrated, microfluidic devices and systems, which enable the fabrication of micro-scale vesicles/particles with cellularlike, internal structures¹⁴ with certain arrangement.^{15,16} Such constructs provide a generic, hierarchical platform, which can enable programmed chemical interactions that are required for the engineering of (bio)chemistry and synthetic biology.17-20

Droplet-forming, fluidic junctions are important elements in droplet microfluidics, and include the T-shaped junction,²¹ the flow-focusing junction²² (also called the crossshaped junction), and the co-flow junction.²³ The droplet formation process in these junctions has been well studied,²⁴ and indicates that the dynamic flow profiles,²⁵ the methods

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to deliver fluids,²⁶ the channel wettability and their geometries,²⁷⁻³⁰ as well as the liquid phase properties³¹ (e.g. chemical composition,³² dynamic viscosity,³³ interfacial tensions,³⁴ surfactant composition,³⁵ suspension components,³⁶ etc.) can all influence the attained droplet morphology. Generally, the strategy to form multiple emulsions is to use a series of droplet-forming junctions, within either microcapillary-based³⁷ or planar, chip-based, microfluidic devices, by either one-step methods³⁸ or multi-step methods.³⁹ The one-step method demonstrates the fine control over the formation of multi-layer droplets,⁴⁰ whilst the multi-step method exhibits the ability to generate compartmentalised droplets, by encapsulating multiple inner droplets⁴¹ (here, we adopt the terminology "cores",⁴² to describe the encapsulated droplets, so as to differentiate them from the overall multiple emulsion droplets). However, due to the existence of hydrodynamic instabilities during droplet breakup, the sequential emulsification process is constrained by the need to synchronise droplet formation at each droplet-forming junction.⁴³ In part, this is because the relatively simple architectures of these classical fluidic junctions have limited scope to spatially confine fluidic interfacial interactions. Consequentially, by using such simple junction architectures, a feed of a first emulsion, into a new secondary immiscible phase, could result in droplet polydispersity, not only due to the absolute size of the overall droplet, but also because of the variations of the encapsulated liquid cores. Therefore, generally, to obtain more precision over compartmentalised droplet formation, superior flow rate controls are required, to constrain the control of droplet morphology within a desired narrow range.43

In the work presented here, we evaluate a new bifurcated microfluidic geometry (the bat-wing junction) that was



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designed to form complex multiple emulsions, in a stepwise emulsification mechanism, *via* its feature of precision flow sectioning. We studied the flow oscillation behaviour at fluidflow bifurcations within the geometry, and studied their influence on the control of droplet breakup and size. Also, we evaluated this geometry for the generation of double emulsion droplets, where we encapsulated multiple reagents, and evaluated the precision control over the number and relative size of the inner cores and their final 3-dimensional orchestration. Further, by numerical simulations, we studied an example of how these multicore constructs could enable sequential chemical reactions within and between such cores, which may open up new applications in (bio)chemical technology, such as smart chemicals delivery and the formation of artificial cells.

Experimental

Chemicals and materials

Trimethylolpropane triacrylate (TMPTA, 246808), photoinitiator 2-hydroxy-2-methylpropiophenone (405655, 5% v/v in TMPTA), glycerol (G5516), mineral oil (M8410), oil red O (O9755), thymol blue (32728), and potassium carbonate (P1472, 3% w/w aqueous solution) were purchased from Sigma-Aldrich Corporation. The optical polymer Norland optical adhesive 61 (NOA 61) was purchased from Tech Optics Ltd.

Fabrication and setup of the microfluidic device

Circular microfluidic chips (50 mm diameter, 3 mm thick) were fabricated from polytetrafluoroethylene (PTFE) using a surface-milling machine (LPKF, C30) fitted with dual-branch, fishtail routers rotating at 30 000 rpm. The resulting rectilinear duct surfaces typically had a roughness (Ra) of $\sim 2 \mu m$ as measured with a VEECO NT3300 white-light interferometer. A 10 mm thick ground glass disc coated with a 100 um thick perfluoroalkoxy (Goodfellow Cambridge Ltd., UK) film was placed over the PTFE disc, so as to provide both (i) a continuous duct geometry with a continuous water contact angle of $\sim 108^{\circ}$ and (ii) a robust viewing window when mechanically compressed within a stainless steel integration and fluidic interface manifold (Fig. S1†).

Experiment and measurement

In the experiments, water/polymer/oil (W/P/O) was used as a template for the formation of double emulsions and fabricated using a stepwise emulsification method on a PTFE substrate microfluidic chip, without any PTFE surface treatment. Constant flow-rate, fluid delivery from 10 mL gas-tight syringes (SGE Analytical Science), through PEEK and PFA interconnects, was enabled *via* syringe displacement pumps (KD Scientific, model 789200L). The morphology of the double emulsion was controlled by the input flow rates, importantly, without adding any surfactant. The core–shell-shaped W/P segments were photo-cured, on the fly, within a length of PTFE outlet tubing (1.5 mm inner diameter) by exposure to a UV light source (5 W 365 nm-wavelength UV LED, placed 1 cm away from the tubing). Solid samples were collected in a beaker filled with a stirred carrier phase, and washed three times with acetone before analysis. A real-time video was recorded to study the droplet formation using a high-speed (400 fps) camera (MS40KD2C1, Mega Speed Corporation) mounted on a Nikon AZ100 microscope. Dimensional images of solid capsules were recorded using a Nikon MM-800 measuring microscope, and analysed with Nikon NIS-elements D 3.2 software.

Computational fluid dynamics simulation

Three-dimensional, time-dependent, computational fluid dynamics simulations were developed to model the droplet breakup process at the bat-wing junction, by using COMSOL Multiphysics 5.0 software (COMSOL Inc., Burlington, MA). The isosurface, volume arrow, velocity magnitude and pressure distributions were plotted as geometric median surfaces to demonstrate the droplet breakup process. Numerical data were obtained for the analysis.

Results and discussion

Description of bat-wing junction geometry and operation

The structures of droplet-forming junctions play an important role in droplet formation, due to the influence of spatial constraints on the delivery, and the interactions, of the fluids employed. The bat-wing junction was designed to comprise two cross-shaped intersections, aligned end-to-end and sharing the side inlets with bifurcations as shown in Fig. 1a. The two intersections were linked by an expansion zone, and terminated with an expansion outlet. These width-expanding structures were utilised to focus and dissipate the fluid flows and create local maximum velocity magnitude points, where the highest shear force occurs. The upstream and downstream bifurcations couple the pressures at the two intersections, and form a uniform pressure gradient, within the expansion zone. The widths of the bifurcations are narrower than the central channel, so as to be easily obstructed by the middle flow. The staggered geometry serves as a fluid shunt and directs the fluid delivery, either upwards or downwards, according to the local dynamic pressure distribution.

Single emulsion formation

Similar to other droplet-forming junctions, three droplet formation regimes were observed when using the bat-wing junction. As shown in Fig. 1b, the two left images show the water droplet (blue) break-up in mineral oil (red), in both dripping and squeezing regimes. The three images to the right demonstrate the TMPTA droplet (red) break-up in mineral oil (transparent), in jetting, dripping and squeezing regimes, respectively. Three-dimensional, computational fluid dynamics simulations were conducted to study the details of the droplet formation at the bat-wing junction (Fig. 1c). It was found



Fig. 1 Geometry of the bat-wing junction. a. The dimension details of the bat-wing junction geometry. The dimensions of the geometry with 600 microns depth are applied to the bat-wing junctions in all the figures of this work. b. Images of different droplet-forming regimes of the bat-wing junction. The two left images show the water droplet (blue) formation in mineral oil (red), in dripping and squeezing regimes. The three right images show the TMPTA (red) droplet formation in mineral oil (transparent) in jetting, dripping and squeezing regimes. c. Simulated images of droplet formation in different droplet-forming regimes of the bat-wing junction. The top row images show the droplet interface (yellow). The yellow films illustrate the boundary between the water phase and the oil phase. The blue vacancies in the yellow films illustrate the water phase touching the top channel wall. The bottom row images show the pressure distribution.

that the continuous phase flow was predicted to oscillate between the upstream and downstream bifurcations. This oscillation was related to the dispersed phase position, inside the expansion zone, during droplet formation (Fig. 2a and b, Video S1[†]). Continuing with the droplet formation process, while the dispersed phase (only) blocks the upstream bifurcation, the local pressure at the blockage starts to increase, forming a vortex, and most of the continuous phase flows through the downstream bifurcation. If the incoming dispersed phase blocks both the upstream and downstream bifurcations, the continuous phase is squeezed within the expansion zone that lies between the upstream and downstream bifurcations. The continuous phase accumulates in the expansion zone, squeezing the dispersed phase into an hourglass form, which eventually leads to break-off or sectioning. Since the volumes of the continuous phase from the upstream and downstream bifurcations flowing into the expansion zone vary dramatically during a droplet formation cycle (Fig. 2c), this in turn induces different flow patterns during droplet breakup (Fig. 2d). Hence, the droplet breakup point, as well as the droplet size, could be controlled by the continuous phase inflow rate (Fig. 2e). This feature of the bat-wing junction was utilised as a precise packaging tool, to assemble uniform, dispersed phase segments of definable length, during the emulsification process within the microchannel.

It was also found that a passive satellite droplet removal mechanism resulted from the squeezing regime, within the bat-wing junction. This was due to the incoming dispersed phase blocking the upstream bifurcation, just after the droplet break-up, resulting in the continuous phase being forced through the downstream bifurcation. Any formed satellite droplets were found to remain at a stationary position inside the expansion zone, and effectively merged with the incoming dispersed phase, due to the surfactant-free emulsion templates (Video S2[†]).

Double emulsion droplet formation. The double emulsion droplet formation is more complicated than that of the single emulsion, since the encapsulated water droplets within the first emulsion offer more resistance during breakup, due to the higher interfacial tension of water, over that of the water and dispersed phases. For the stepwise emulsification method, the existence of multiple hydrodynamic instabilities will influence the monodispersity of the final multiple emulsion droplets. In the next two sections, we demonstrate the performance of the bat-wing junction in forming uniform double emulsion droplets using a two-step emulsification method.

Monodispersed core-shell-shaped droplets. Here, we investigated the formation of double emulsions, using the batwing junction as a second droplet-forming junction, to section the first emulsion, in a two-step method. As shown in Fig. 3a, the first emulsion was formed at the first crossshaped, flow-focusing junction, as the inner water phase (blue, from the middle) droplets were broken up in the TMPTA flow (transparent, from sides). The shell phase inflow rate was maintained at a constant 0.2 ml h⁻¹, and the inner water phase inflow rate was increased from 0.05 ml h⁻¹ to 0.1 ml h⁻¹ until 1.4 ml h⁻¹, in 0.1 ml h⁻¹ steps, from left to right (Fig. 3a), respectively. These combinations formed different lengths of the first emulsion segment, and it was found that, by changing the continuous phase inflow rates (left transparent, right red) at the bat-wing junction, monodispersed, core-shell-shaped double emulsion droplets could be precisely sheared off, or sectioned, in various volume ratios, between the core and shell phases.



Fig. 2 Control of droplet break-up at the bat-wing junction. The channel geometry refers to Fig. 1a. a. Time sequence images of the water droplet formation in the squeezing regime. The dispersed phase blocks all the bifurcations, and the continuous phase flows into the expansion zone from two directions. The blue/clear/red phases' flow rates are 1/0.3/0.3 ml h⁻¹. b. Time sequence images of the two-phase droplet breakup simulation in the squeezing regime. The yellow cones show the local flow direction. Eddy flow (as shown by the yellow contour) appears where the blockage occurs (the expansion zone and the bifurcations) and the size changes with time. The colour gradient shows the pressure distribution. The flow rate in the top inlet is 1 ml h⁻¹ and those in the side inlets are both 0.5 ml h⁻¹. c. Continuous phase flows oscillated between the upstream and the downstream bifurcations. The average flow rates were measured at the cross-section of the coloured lines shown in b. These images also indicate that different amounts of the continuous phase flowed into the expansion zones under different inlet combinations. d. Coupled CFD simulations and laboratory experiments illustrate the different water droplet breakup locations in oil at the bat-wing junction. All dispersed phases' input rates are 2 ml h⁻¹. The continuous phase flow rates, from left to right, are 0.28 ml h⁻¹, 0.52 ml h⁻¹, 0.64 ml h⁻¹ and 1.60 ml h⁻¹, respectively. e. The relationship between the continuous phase flow rate with the droplet breakup point and droplet size. When the continuous phase flow rate is increased, the droplet breakup location.

Next, we tested the consistency, by which the bat-wing junction formed core-shell-shaped double emulsion droplets, by measuring both their yield rates and size distributions. As shown in Fig. 3b, while the inflow rate ratios of inner/shell phases were held constant, the yield rates of double emulsion droplets at the outlet were elevated by increasing the total inflow rate. The variations in time intervals between adjacent droplets entering the outlet were ±4.84%, ±4.74% and $\pm 3.47\%$, for the three groups in Fig. 3b, from left to right, respectively. At the same time, the sizes of the double emulsion droplets were reduced, while the total inflow rate was increased. The size variations (n = 100 for each group) of the inner core droplets were ±2.47%, ±2.01% and ±1.58%, and those of the total double emulsion droplets were ±1.85%, $\pm 1.85\%$ and $\pm 1.86\%$, from low to high inflow rates. These results suggested that the bat-wing junction could form highly replicated, core-shell-shaped double emulsion droplets, by a two-step method under different inflow rate combinations.

Compartmentalized double emulsion droplets. Furthermore, the bat-wing junction could be used to precisely control the number of the encapsulated droplet cores within the double emulsion droplets. This was achieved by forming a consistent first emulsion, with repeated droplet patterns, and then precisely sectioning them, at the bat-wing junction. As shown in the rest of Fig. 3, the inflow rates of the inner and shell phases were kept constant, resulting in the formation of two strings of droplets in the shell phase flow, as the first emulsion. By changing only the continuous phase flow rates, the number of droplets (which became the inner cores of the double emulsion) could be precisely tuned to a resolution of one core, as shown in Fig. 3c. This resolution could be maintained for up to 15 inner cores, as shown in Video S3.† In addition, once the shape of the double emulsion droplets had changed from segments (Fig. 3d) to spheres (Fig. 3e), the inner droplet cores self-arranged into specific 3D geometric orchestrations, which appeared to depend upon the number of inner cores that were incorporated within the double emulsion.

Multiple types of encapsulated droplets. Several groups of control experiments were conducted to form complex double



Fig. 3 Control of double emulsion formation with uniform inner droplets at the bat-wing junction. The channel geometry refers to Fig. 1a. a. Core-shell-shaped double emulsion droplet formation by the two-step method, using the bat-wing as the second droplet-forming junction. The shell phase flow rates are all 0.2 ml per hour, and the inner phase (blue) flow rates are 0.05, 0.1, 0.2, 0.3..., 1.3, and 1.4 ml per hour from left to right, respectively. b. The size distributions and the formation rates of the double emulsion droplets under different inlet combinations. From left to right, the standard deviations of the inner droplet sizes (blue) are $\pm 2.47\%$, $\pm 2.01\%$, and $\pm 1.58\%$, and the standard deviations of the total double emulsion droplet size (blue + pink) are $\pm 1.85\%$, $\pm 1.85\%$, respectively. c. Control of the inner droplet number within double emulsion droplets by tuning the continuous phase flow rate. d. Images of double emulsion droplet formation in a 600 micron deep channel, encapsulating 2, 4, 6, and 10 inner droplets. e. The 3D arrangements of the inner droplets within the spherical-shaped double emulsion droplets. The DE droplets just flowed to the 1.2 mm deep channel, from d, to attain spherical forms with the inner droplets self-arranging, moving from left to right. The top to bottom rows illustrate 4, 6, 8, and 12 inner droplets within the double emulsion droplets, respectively.

emulsions that encapsulated multiple types of inner droplet cores, within a single, shell-phase matrix. These were achieved by using two T-junctions, to generate two independent, repeating droplet groups (to become inner cores), within the shell phase flow, as the first emulsion. The patterns of these droplet groups (reagents, sizes and order) were controlled by the inflow rate combinations. Then, the first emulsions were precisely sectioned off, to form patterned double emulsions. This sectioning encapsulated the first emulsion groups of inner droplet cores, the number of which could be determined by simply tuning the continuous phase flow rates, to change the droplet breakup points within the bat-wing junction. The groups of inner droplet cores could include either even (Fig. 4a-c) or odd (Fig. 4d) numbers of such cores. As the double emulsion segments attained a spherical form, the inner droplet cores also arranged into certain geometric orchestrations, as shown in Fig. 4e. It appears that such core arrangements within the double emulsion droplets may possibly be determined by either, or both, their contiguous order or/and their sizes, within the first emulsions. In addition, a droplet-forming array, which included four T-junctions as shown in Fig. S2,† was used to produce more complicated, repeated patterns of the inner droplet core assemblies, in the shell phase flow, as the first emulsion. As shown in Fig. 4f and g, the bat-wing junction was used to consistently section off double emulsion droplets, each encapsulating four different types of inner droplet cores. The sizes and numbers of the inner droplet cores were controlled by the inflow rate combinations. As shown in Fig. 4j, by changing the inflow rates of the inner phases, which were 0.2/0.2/0.7/0.7 (ml h⁻¹, inlet combination 1), 0.4/0.4/0.5/0.5 (ml h⁻¹, inlet combination 2), 0.6/0.6/0.3/0.3 (ml



Fig. 4 Control of double emulsion formation with multiple types of inner droplets at the bat-wing junction. The channel geometry refers to Fig. 1a. All the scale bars indicate 1.2 mm. a-c. Double emulsion droplet formation from the first emulsions with even-numbered inner droplet groups. d. Double emulsion droplet formation from the first emulsions with odd-numbered inner droplet groups. e. The 3D arrangements of inner droplets within spherical double emulsion droplets. From left to right, the yellow/blue inner phase flow rates are 0.3/0.1, 0.2/0.2, and 0.3/0.1 ml h⁻¹, respectively. The shell phase and continuous phase flow rates are 0.8 ml h⁻¹ and 3.0 ml h⁻¹ as constants. f-i. Double emulsion droplet formation encapsulating four types of inner droplets, and their 3D arrangement within the spherical droplets. j. The numbers of inner core droplets within the double emulsions under different inflow combinations.

 h^{-1} , inlet combination 3), the numbers of inner droplet cores from each colour group were controlled, whilst the total number of cores remained in the range of 15–20. This was due to the (i) shell phase and continuous phase input flow rates being kept at a combined total of 0.8/2.0 ml h^{-1} for all the three combinations, (ii) the total first emulsion flow to the batwing junction remaining at a constant flow rate, and (iii) the continuous phase, which precisely sectioned the first emulsion to achieve repeated patterns. The variations in the core number partially resulted from the size variation of the different coloured water droplets, which could be reduced by higher precision pumping mechanisms. As these double emulsion droplets attained spherical forms, the inner droplet cores also self-arranged into specific 3D geometrical orchestrations, within the shell phase matrix (Fig. 4h and i).

Microcapsules and multisomes

So far, we have demonstrated how to form highly uniform double emulsion droplets that encapsulate multiple types of inner droplet cores, with controllable morphology, using the bat-wing junction and a stepwise emulsification method. The 3D geometrical arrangement of inner droplet cores appeared to depend on both their number and size and the specific contiguous order within the first emulsion. These double emulsion droplets could represent very interesting templates, to fabricate highly compartmentalised micro-constructs with given functionalities. Fig. 5a-r show solid polymeric microcapsules, which were solidified from the double emulsion droplets and produced using the methodologies described above. Once the droplets had attained an approximately spherical form, the shell phase was photopolymerised, on the fly, using a 365 nm UV light source. The photopolymerised shells formed an effectively fixed encapsulating matrix, within which the inner droplet cores, containing chemical reagents, were held in a constant, 3-dimensional geometrical orchestration. This fixation thereby enabled subsequent metrological analysis. Such multicore microcapsules, with a liquid encapsulation matrix, may be used to enable multiplereagent release or in situ chemical synthesis in relation to external stimuli. Such encapsulation matrices could include a wide range of degradable polymers (e.g. polylactic acid) and



Fig. 5 Highly compartmentalized microcapsules and multisomes, with 3D inner core droplet arrangements. All the scale bars indicate 500 microns. a-o. Solid NOA61 capsules with 2–15 uniform inner water droplets, respectively. All the capsules were produced on the same microfluidic chip by curing double emulsion droplets, which were formed using the bat-wing junction by a two-step method on a PTFE microfluidic chip. p-r. Compartmentalized, solid TMPTA capsules with multiple types of inner water droplets, fabricated by the same approach as a-o. s-v. Encapsulated droplet interface bilayers in water/squalene/TMPTA double emulsion droplets, formed on PMMA-based, planar microfluidic chips. w-y. Multisomes formed from water/squalene/3% alginate solution double emulsions on PMMA microfluidic chips. The squalene phase contains 2% 1,2-diphytanoyl-*sn-glycero*-3-phosphocholine.

semi-permeable materials (*e.g.* NOA61, as used here) or employ bilayer lipid membranes decorated with functional intercore or inter-droplet proteins. In contrast to the aforementioned constructs fixed within a photopolymerised polymer, 'soft' analogues (*e.g.* multisomes) were formed from water/squalene/TMPTA emulsions (Fig. 5s–v) or water/squalene/3% alginate emulsions (Fig. 5w–y), using the same methods as used before, but on PMMA-based microfluidic chips. The numbers of the inner water droplets were controlled, and the lipids, which were pre-suspended in squalene, were assembled around them to form monolayers. Once the double emulsion droplets had attained spherical forms, the inner

droplet cores, again self-arranged, and high-order, 3D droplet interface bilayer networks were established. Such artificial vesicles hold great potential for biotechnology developments and medical applications.⁴⁴

Finally, we simulated model sequential reactions and molecular diffusion within a typical multicore construct. This model consisted of 3D inner droplet arrangements that were based on the practical experiments demonstrated in this paper. Fig. 6a models a gas capture process within a 13-core, gas-permeable microcapsule that encapsulated the active reagents. The chemical reaction within the central core was the last in this sequential chain reaction, due to the specific



Fig. 6 Simulated sequential chemical reactions within the compartmentalised capsules. a. Reagent A is captured by the reaction when flowing over the compartmentalised microcapsules (13 inner droplets, reproduced from Fig. 5i). The top image shows the concentration of A. The microcapsule encapsulates B within the inner droplets. The four bottom images show the increasing concentration of C within the droplets. b. Demonstration of chain chemical reactions within a multisome (reproduced from Fig. 5x), which is defined to load four water-soluble reagents b, d, f and h, represented by different colours. The top image shows the model geometry. Reagent a is assumed as the molecule that can transmit through the lipid bilayer (white lines). The bottom image shows the concentrations of the chemical compounds within the multisomes.

geometrical arrangement of cores. Fig. 6b models the sequential chemical reactions within a multicore multisome that contained four types of cores, each with a different reagent. The diffusion of the water-soluble reagents was influenced by the bilayer positions, which also depended upon the inner droplet core arrangements. The reactions took place while the molecules diffused within the multisome. These demonstrations may provide the beginnings of more precise engineering tools for programmed chemistry and synthetic biology.

Conclusions

In summary, we demonstrate a novel, droplet-forming fluidic junction, for which we coined the term, the 'bat-wing junction'. Its bifurcation structures oscillate the side flows during droplet formation, and the droplet break-up point and droplet size were regulated, by adjusting only the continuous phase inflow rate. We also found that the junction acted as a passive satellite droplet removal mechanism, due to its inherent flow confinement behaviour. This junction can be used in stepwise emulsification methods, to precisely section a patterned first emulsion and then form uniform double emulsion droplets, which can encapsulate inner cores (droplets) of various sizes and reagent composition. It was found that once the double emulsions attained a spherical form in the capillary, the inner core self-arranged in certain 3D geometric orchestrations, which appeared to depend upon their sizes, numbers and order, within the double emulsion droplets. We produced highly compartmentalised microcapsules and multisomes from these double emulsion droplets, on the fly, to inherit the formed droplet morphologies. We further demonstrated by numerical simulation how these microconstructs could possibly enable sequential chemical reactions, as a generic platform for diverse applications, ranging from (bio)chemical synthesis⁴⁵/analysis,⁴⁶ cell passage,⁴⁷ smart drug delivery,^{48,49} computation⁵⁰ and self-repair materials,⁵¹ and artificial cells.⁵²

Certain limitations exist in the work reported here that could be explored further, including critically the spatial constraints due to the 2.5D nature of the planar bat-wing junction used in this study. For this, a truly 3-dimensional batwing junction could be manufactured by the adoption of novel additive manufacturing methods, such as 3D-printing.^{53,54} This could enable the more exact sectioning of a first emulsion into even more accurately defined, internally compartmentalised capsules. Also, additional precision control of individual cores (size, order, reagent concentration, *etc.*) within the first emulsion (before sectioning as double emulsion droplets) could be found by employing pulseless, fluid propulsion mechanisms. Further, sequential droplet processing could be enabled by liquid carrier phase exchange,⁵⁵ vesicle/particle surface overcoating,⁵⁶ and noninvasive in-flow droplet characterization.⁵⁷ Such improved characterisation could be enabled through non-invasive, onthe-fly, interferometric or tomographic methodologies, and could enable the increased resolution of individual core droplet volumes and dimensions.

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