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Mechanisms and control of single-step microfluidic generation of multi-core double emulsion droplets

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HIGHLIGHTS

- The mechanisms of single-step formation of multi-core double emulsion were revealed.
- Phase diagrams for predicting number of encapsulated inner drops were mapped out.
- The jet pinch-off process was accompanied and affected by presence of a vortex flow.
- The formation regime of inner phase affected the number of encapsulated inner drops.
- Large single-core capsules were made from dual-core drops with unstable inner drops.

ABSTRACT

Single-step generation of monodisperse multi-core double emulsion drops in three-phase glass capillary microfluidic device was investigated using a micro-particle image velocimetry (micro-PIV) system. Phase diagrams were developed to predict the number of encapsulated inner drops as a function of the capillary numbers of inner, middle and outer fluid. The maximum stable number of inner drops cores in uniform double emulsion drops was six. Starting from core/shell drops, the formation of double emulsion drops with multiple cores was achieved by decreasing the capillary number of the outer fluid and increasing the capillary number of the middle fluid. A stable continuous jet of the middle fluid loaded with inner drops was formed at high capillary numbers of the middle fluid. Empirical correlations predicting the size and generation frequency of inner drops as a function of the capillary numbers and the device geometry were developed. Dual-core double emulsion drops were used as templates for the fabrication of polymeric capsules using “on-the-fly” photopolymerisation. The capsule morphology was controlled by manipulating the stability of the inner drops through adjusting the concentration of the lipophilic surfactant in the middle fluid. At low concentration of the lipophilic surfactant, inner drops coalesced during curing and single compartment capsules with thin shells were produced from dual-core drops. The core/shell capsules produced from multi-core drops were monodispersed and larger than those produced from core/shell drops in the same device.

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1. Introduction

Multi-core double emulsions are complex emulsions composed of several inner drops dispersed in an immiscible middle phase, which is itself dispersed in the outer phase. They can be used for encapsulation of multiple reactive components for triggered reaction and mixing [1–3], co-culture of multiple cells in cell-based therapies [4], encapsulation and sequential release of incompatible active materials and liquids while avoiding cross-contamination [5], and the production of multi-compartment capsules [6]. Conventionally, multiple emulsions are produced in two consecutive emulsification steps through high-shear mixing of immiscible liquids [7]. However, this approach is associated with broad particle size distribution, low batch-to-batch reproducibility, low encapsulation efficiency, high energy consumption, and a lack of control over the size of the drops and the number of encapsulated inner drops [8,9].

Microfluidic emulsification is a promising tool for reproducible production of monodispersed multiple emulsion drops of versatile morphology [5,10–12]. Multi-core drops are usually produced using two consecutive T-junction [8,13], co-flow [10,11,14–16], and flow-focusing [17,18] drop makers. Perro et al. [16] used two consecutive co-flow makers made of fused silica capillaries to produce structured polymersomes and map out phase diagrams predicting the number of inner drops as a function of fluid flow rates. The number of inner drops decreased as the outer and middle phase flow rates increased. Okushima et al. [13] used two consecutive T-junctions made on a Pyrex glass to produce double emulsion drops with a controlled number of inner drops. The number of inner drops was found to increase with increasing flow rate of the inner phase. Using a two-step microfluidic device composed of three coaxial glass capillaries, Kim et al. [11] observed that the number of inner drops increased with increasing middle-to-inner volumetric flow rate ratio. Chu et al. [10] derived an empirical equation to predict the number of inner drops in multi-core droplets produced in a glass capillary device consisting of two consecutive co-flow drop makers. The number of inner drops was reduced by increasing the outer phase flow rate.

In the two-step emulsification approach, the generation of inner and outer drops is spatially separated; thus, synchronising the frequency of drop generation at different steps requires delicate control of individual flow rates [5]. Single-step microfluidic emulsification methods [6,19,20], e.g., a combined co-flow and counter-current flow focusing (Fig. 1a) [21], in a three-phase glass capillary device, can overcome this disadvantage, since the flow rates of the inner and middle fluid do not need to be synchronised. Moreover, one-step processes are advantageous for very thin-shelled core/shell drops, which are difficult to create in a two-step process. In addition, the capillary device for single-step emulsification is easier to fabricate than the device composed of two sequential drop generation units [10,15], due to simpler design and smaller number of capillaries and connectors [6,22].

Three main regimes of double emulsion generation in three-phase glass capillary devices are dripping, widening jetting, and narrowing jetting. In the dripping regime, highly uniform drops are formed near the orifice of the collection tube, while widening and narrowing jetting are associated with the production of poly-dispersed large and small drops, respectively, further downstream of the orifice [23,24]. For highly uniform multi-core drops, the cores should be formed in the drooping regime and the outer drops in the dripping-to-widening jetting transition regime [6]. Lee et al. [6] used a three-phase glass capillary to produce nonspherical colloidosomes. The higher inner and middle phase flow rates and the lower outer phase flow rates favoured encapsulation of larger numbers of inner drops. Although Lee et al. [6] briefly reported the effect of flow rates on the number of encapsulated inner drops, however, the mechanism of single-step formation of multi-core double emulsion drops has not yet been fully understood and needs to be investigated.

The purpose of this work is an in-depth investigation into the mechanisms of single-step generation of multiple emulsion drops in a co-flow/flow focusing microfluidic geometry. Novel phase diagrams have been developed to depict the effect of capillary numbers of the inner, middle, and outer fluids on the number of inner drops, and determine the operating conditions for successful production of double emulsions. The flow field during jet pinch-off was visualised by seeding charged latex beads in the outer phase. The generated drops were used as templates to produce capsules with controlled number of compartments via “on-the-fly” photopolymerisation. The number of cores in the capsule was controlled by manipulating the stability of inner drops during curing by varying the concentration of lipophilic surfactant in the middle...
phase. The microfluidic method developed in this work can be used for production of relatively large silicon rubber-based capsules for encapsulation of CO₂ absorbents [25,26].

2. Experimental procedure

2.1. Materials

K₂CO₃ and glycerol were supplied by Fisher Scientific (UK). Polyvinyl alcohol (PVA, Mn = 13,000–23,000 g mol⁻¹, 87–89% hydrolysed), Pluronic® F-127 (PF-127), 2 μm carboxylate-modified polystyrene latex beads, and m-cresol purple were purchased from Sigma-Aldrich, UK. Dow Corning® 749 Fluid (DC 749) was supplied by Univar (UK). Polydimethylsiloxane oil (Dow Corning 200 fluid, 20 CST) was obtained from VWR (UK). Octadecyltrimethoxysilane (OTMS) and 2-[methoxy(polyethyleneoxy)]propyltrimethoxysilane (TMS-PEG) were purchased from Fluorochem, Ltd. (UK). Semicosil® 949 UV A/B, a UV-curing silicone rubber was donated by Wacker Chemie (UK). The reverse osmosis water was supplied from a Milli-Q plus apparatus. The fluid densities, viscosities and interfacial tensions were measured using a pycnometer, capillary viscometer, and Krüss DSA-100 pendant drop tensiometer, respectively.

2.2. Droplet production

2.2.1. Fabrication of glass capillary device

A Flaming/Brown micropipette puller (P-97, Sutter Instrument Co.) was used to pull round glass capillaries (I.D. = 0.58 mm, O.D. = 1 mm, Intracel, UK) that were used as injection and collection tubes. The tips were polished to the desired orifice size using an abrasive paper (Black Ice Waterproof T402, Alpine Abrasives, UK) and inspected using a microforge microscope (Narishige MF-830, Japan). OTMS was used to make the injection orifice hydrophobic and TMS-PEG was used to enhance hydrophilic properties of the collection tube. The injection and collection capillaries were then inserted into a square glass capillary (inner width = 1.05 ± 0.1 mm, AIT Glass, Rockaway, US) from two opposite ends, placed at the desired distance from each other and axially aligned. The capillaries were fixed into position on a microscope slide using 2-part 5–30°C part AB glue (Devcon, UK). To introduce the three phases to different radii of curvature across the interface, causing the drop to have an elongated shape, which is unstable because of the difference in capillary pressure in the thinning primary neck. At τ = 0 (Fig. 2a), the primary neck of the previous outer drop has collapsed and the previous outer drop was detached. The 11st drop was detached at τ = 95, the middle fluid thread of the previous drop disappeared after reaching a certain minimum thickness. The drop formation process can be divided into three main steps: (1) formation of inner drops by multiple break-ups of the inner fluid jet; (2) advection of inner drops through the middle fluid and their interaction with the growing perturbations on the outer interface; (3) break-up of the middle fluid jet and encapsulation of inner drops in the outer drop.

At τ = 0 (Fig. 2a), the primary neck of the previous outer drop has not yet collapsed and the 1st drop (shown in red at τ = 0), and the subsequent inner drops will be denoted as the second (2nd), third (3rd), and fourth (4th) drop, respectively. The fifth drop (shown in red at τ = 612), will become the first inner drop encapsulated in the next outer drop and will be denoted as the 1st drop. The primary neck formed downstream of the 1st drop (Fig. 1b) gets thinner and thinner until a pinch-off finally takes place. On the other hand, the secondary neck formed temporarily upstream of each inner drop disappears after reaching a certain minimum thickness.

2.3. Image analysis and particle tracking

An inverted microscope (XDS-3, GX Microscopes, UK) and Phantom V9.0 high-speed camera interfaced to a PC computer were used to observe and record the drop generation process. The time-dependent movements of inner drops were tracked by analysing successive frames captured with a resolution of 768 × 576 pixels at 2000 Hz with Imagej software. The flow field was visualised using a micro-particle image velocimetry (microPIV) system incorporating 2 μm charged latex beads seeded in the outer fluid, a visible light source and the Matlab-based PIVlab (V 1.4) software. The minimum concentration of the tracer particles of 0.167 μg/mL and 20× objective lens were used to obtain acceptable visualisation outputs.

2.4. Dimensionless numbers

Non-dimensional numbers used in this study are the capillary numbers of the inner, Ca₁, middle, Ca₂ and outer phase, Ca₃, which are the ratio of viscous to interfacial tension forces, the Weber number for the inner phase, We₁, that gives the ratio of inertial to interfacial tension force in the inner phase, and the non-dimensional time, τ:

\[
Ca_1 = \frac{\mu_1 V_{1N}}{\sigma_{12}} \quad Ca_2 = \frac{\mu_2 V_2}{\sigma_{23}} \quad Ca_3 = \frac{\mu_3 V_3}{\sigma_{3}}
\]

\[
We_1 = \frac{\rho_1 V_{1N}^2 D_1}{\sigma_{12}} \quad \tau = \frac{V_{1N} t}{R_N}
\]

where \(\rho\), \(\sigma\), \(\mu\), and \(V\) are the density, interfacial tension, dynamic viscosity and velocity, respectively (the subscripts 1, 2, and 3 stand for the inner, middle, and outer phase, respectively), \(R_N\) is the radius of the injection nozzle and \(t\) is the time. The characteristic fluid velocities are:

\[
V_{1N} = 4Q_1/\left(\pi D_1^2\right)
\]

\[
V_2 = Q_2/\left(D_2^2 - \pi D_1^2/4\right)
\]

\[
V_3 = Q_3/\left(D_3^2 - \pi D_2^2/4\right)
\]
Due to decay of the secondary neck, the maximum velocity of the 2\textsuperscript{1st} drop was higher than that of the 1\textsuperscript{1st} drop (Fig. 2b). At $\tau = 408$, the 3\textsuperscript{1st} drop was pinched off but the maximum velocity reached by that drop was lower than the maximum velocity of the 2\textsuperscript{1st} drop because the pinch-off process of the primary outer drop (marked by the red dashed rectangle) entered its final stage characterised by a high pressure in the neck. At $\tau = 612$, the 4\textsuperscript{1st} drop was pinched off and reached higher velocity than the previous inner drops, because the outer drop was detached at $\tau \approx 600$ causing the middle phase jet to accelerate afterwards (Fig. 2b). The subsequent outer drop in the blue dashed rectangle was pinched off at $\tau = 1224$.

Fig. 2b shows the temporal variations of the velocity of inner drops in dimensionless coordinates for the image sequences shown in Fig. 1. For easier comparison, the formation of each inner drop starts at $\tau = 0$. Region I ($\tau < 150$) was characterised by an increase in the velocity of inner drops and corresponds to the growth and pinch-off of each inner drop. During this step, the drop shape increasingly deviates from spherical due to increasing drag force acting from the middle fluid. Thus, the forming drop is increasingly unstable, since the difference in Laplace pressure between the upstream face of the drop and its neck increases, which causes the velocity to rise. The maximum relative velocity ($V_d/V_{1,N}$) for the 2\textsuperscript{1st}, 3\textsuperscript{1st}, and 4\textsuperscript{1st} drop of 0.15, 0.14, and 0.16, respectively, was achieved later in the cycle and was greater than the maximum relative velocity for the 1\textsuperscript{1st} drop of 0.13. The low velocity of the 1\textsuperscript{1st} drop was due to thinning of the primary neck, which prevented further advection of the 1\textsuperscript{1st} drop through the middle phase. Region II ($150 < \tau < 600$) corresponds to the decrease in the velocity of inner drops, which can be attributed to the break-up process of the primary outer drop, associated with rising pressure in the primary neck. Region III ($\tau > 600$) occurs after pinch-off of the primary outer drop and is characterised by a moderate increase in velocity followed by a slight decrease. The velocity of inner drops was increased due to dissipation of the vortex flow, while the subsequent decrease in velocity was due to the break-up of the next outer drop after encapsulation of all inner drops.

The middle phase jet has the minimum velocity just after the pinch-off of each outer drop due to recoiling of the jet interface as a result of high capillary pressure in the thread connecting the detaching drop to the remaining. This decrease in velocity of the middle phase jet in the collection tube can be observed in the video.

### Table 1

<table>
<thead>
<tr>
<th>Phase</th>
<th>Fluid Composition</th>
<th>$\rho$ (kg/m(^3))</th>
<th>$\mu$ (mPa s)</th>
<th>$\sigma$ (mN/m)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inner phase</td>
<td>85 wt% glycerol in Milli-Q water</td>
<td>1225</td>
<td>109</td>
<td>29.9</td>
</tr>
<tr>
<td>Middle phase</td>
<td>2 wt% DC 749 in Dow Corning 200 Fluid, 10 CST</td>
<td>940</td>
<td>10.4</td>
<td>31.8</td>
</tr>
<tr>
<td>Outer phase</td>
<td>70 wt% glycerol + 2 wt% PVA in Milli-Q water</td>
<td>1118</td>
<td>33</td>
<td></td>
</tr>
</tbody>
</table>

* The interfacial tension was measured in the absence of stabilisers.
recordings of double emulsion drop formation (Movies 1–3 in the Supplementary material S1). In order to identify the additional factors that can possibly affect the break-up mechanism, the flow field during the outer drop pinch-off was visualised with polystyrene tracer beads (Fig. 3).

The effect of Brownian motion can significantly affect the instantaneous velocity field [27]. In the current study, however, due to the large size of tracer particles (d = 2 μm), high characteristic velocity (V3 ~ 7.4 mm/s), and high dynamic viscosity of the outer fluid in which the tracer particles were suspended, the Brownian motion effect was negligible. The relative error due to Brownian motion, εB, can be estimated from [28]:

$$\varepsilon_B = \frac{1}{V_3} \sqrt{\frac{2D_{\text{diff}}}{\Delta t}}$$  (1)

where Δt is the time interval between two successive frames (0.5 ms), and D_{\text{diff}} (m²/s) is the diffusion coefficient of the tracer particles given by the Stokes-Einstein equation:

$$D_{\text{diff}} = \frac{\kappa T}{3\pi \mu_3 d_i}$$  (2)

where T is the absolute temperature, and κ is Boltzmann’s constant. The εB value calculated from Eq. (1) was 0.07% implying the negligible effect of Brownian motion on the normalised velocity profiles in Fig. 3. In Fig. 3a–c, it was found that the primary neck narrowing was accompanied by the formation of vortex flow tending to push backward and forward the outer interface at the upstream and downstream end of the neck, respectively, which eventually resulted in the thread break-up (Fig. 3d). Once the thread pinched off, the detached drop accelerated downstream, followed by rapid retreat of the elongated middle phase filament attached to the primary outer drop (shown in the red dashed rectangle in Fig. 3d). It is driven by the imbalance in capillary pressure across the interface and the presence of vortex flow.

On the other hand, a backflow upstream of the neck exerted a backward force that caused a deceleration of the middle phase jet and low velocity in the region shown by the black dotted box in Fig. 3d. The presence of vortex flow during necking of the middle phase jet that facilitated the jet break-up was observed numerically by Nabavi et al. [24].

### 3.2. Predicting number of inner drops

In this section, the effect of capillary numbers of inner, middle and outer fluids on the size and number of inner drops will be investigated by mapping flow patterns and identifying regions where monodisperse drops with a stable number of inner drops (N_d) can be formed.

Figs. 4a–c.1 are phase diagrams plotting N_d as a function of Ca_1, Ca_2, and Ca_3 for the fluids indicated in Table 1. The capillary numbers were varied by changing the flow rates of the inner, middle and outer fluids. The experimental data used to construct these diagrams are shown in Fig. s.2. The hatched green area represents the regions where the number of inner drops varied between 2 and 4. On the other hand, the solid black, cyan, yellow, and red areas represent the regions where N_d was stable over longer time periods, and equal to one, two, four and six, respectively. The formation of uniform drops occurred when Ca_1 was greater than Ca_2 or Ca_3 by 2–4 orders of magnitude. The maximum stable N_d value of 6 was achieved at Ca_3 = 4.6 x 10^{-3}. Monodispersed multi-core drops were created within the Ca_2 range of (1.5–4.8) x 10^{-4}. For Ca_3 < 1.5 x 10^{-4}, N_d was either 1 or varied uncontrollably between 1 and 2. For Ca_2 > 4.8 x 10^{-4}, a continuous jet of middle phase encapsulating numerous uniform inner drops was formed (Fig. 4f).

Lower Ca_3 and higher Ca_2 and Ca_1 values favoured formation of drops with larger numbers of inner drops; e.g., at Ca_1 = 0.42 and Ca_2 = 2.9 x 10^{-4}, a decrease in Ca_3 from 6.9 x 10^{-3} to 4.6 x 10^{-4} to 3.1 x 10^{-3} caused an increase in N_d from 1 to 2 to 4, respectively (Fig. s.3). A decrease in Ca_3 reduced the shear and pressure forces on the outer interface exerted by the outermost fluid, which caused the jet widening in the collection capillary. The increased jet-to-orifice diameter ratio, D_0/D_{\text{out}} is slowing down the rate of growth of perturbations on the outer interface due to its proximity to the wall [5,29], increasing the jet break-up lengths and allowing more inner drops to be encapsulated. However, Ca_3 should be large enough for the outer fluid to be able to draw the middle phase through the orifice. For Ca_3 < 3.1 x 10^{-3}, the outer fluid was not able to focus the compound jet into the collection tube, which resulted in the leakage of the inner fluid into the outer phase (Fig. 4d).

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**Fig. 3.** Flow field visualisation of the jet pinch-off using μ-PIV. The outer drop consists of two inner drops. The dimensionless time is: (a) t_0; (b) t_0 + 26; (c) t_0 + 186; (d) t_0 + 348. All scale bars are 100 μm.
When Ca₁ and Ca₃ were kept constant, N_d increased with increasing Ca₂ (Fig. 4a.1–c.1), because outer drops grew to a larger size before breaking off and thus, more inner drops could be encapsulated [23]. In addition, higher Ca₂ values at constant Ca₁ led to smaller inner drops, due to higher shear force acting on the inner fluid (Fig. 4a.2–c.2). A decrease in D₁ led to an increase in the frequency of inner drop formation, which resulted in higher N_d.

Increasing Ca₁ above a certain threshold value at constant Ca₂ and Ca₃ resulted in unsuccessful generation of multiple emulsion drops; two possible scenarios are shown in Fig. 4e and f. The flow behaviour shown in Fig. 4e was observed above the boundary line I in the phase diagram, which means at low Ca₃ and/or low Ca₂ values. In that case, due to small Q₂/Q₁ ratio, the inner fluid leaked into the outer fluid (the red dashed box in Fig. 4e). The flow behaviour shown in Fig. 4f and Movie 4 (in Supplementary material S1) occurred above the boundary line II when the volume fraction of the middle phase was higher than that of the inner phase. In that case, a high frequency of generation of inner drops suppressed perturbations of the outer interface, which led to the formation of a continuous middle phase jet loaded with numerous inner drops. This drop generation regime can be used for fabrication of hybrid microfibers [30,31].

Regarding the effect of Ca₃ on N_d, two different behaviours were observed. At Ca₃ < 1, the inertia of the inner fluid was small compared to viscous forces acting on the inner interface and N_d increased with increasing Ca₃. At 3 > Ca₃ > 1, inertial forces domi-
nated over viscous forces and N₂ decreased with increasing Ca₁ due to an increase in the size of the inner drops (Fig. 5). It should be noted that D₁ increased with increasing Ca₁ over the entire range of Ca₁ investigated, although at high Ca₂ and Ca₃ values, D₁ was nearly constant at Ca₁ < 0.4 (Fig. 4a.2–b.2).

An empirical correlation was developed to predict D₁ as a function of Ca₁, Ca₂, Ca₃, and D₉₁, based on experimental data obtained at different fluid flow rates:

\[ \frac{D₁}{D₉₁} = 0.2 \frac{Ca₁^{0.23}}{Ca₂^{0.06} Ca₃^{0.27}} \]

(3)

As shown in Fig. 6, Eq. (3) was in good agreement with the experimental data. The frequency of generation of inner drops, f₁ and the Capillary number of inner fluid, Ca₁, can be expressed as:

\[ f₁ = \frac{6 Q₁}{π D₁} \]

(4)

\[ Ca₁ = \frac{4}{π D₉₁} \frac{μ₁}{σ₁₂} Q₁ \]

(5)

Substituting Eqs. (4) and (5) into Eq. (3) one obtains:

\[ f₁ = 187.5 \frac{D₉₁^{3} \ σ₁₂^{3}}{μ₁^{3}} \frac{Ca₁^{31}}{Ca₂^{18}} \frac{Ca₃^{0.69}}{Ca₉₁^{0.69}} \]

(6)

According to Eq. (6), f₁ increases with increasing the capillary number of all fluids. In Fig. 5, an increase in Ca₁ from 0.2 to 2.6 led to an increase in f₁ by a factor of 2.2. The same trend with an increase in f₁ and D₁ as a result of increase in Q₁ was observed by numerical simulations [23]. An increase in N₉₁ from 1 to 6 at Ca₁ = 0.2–1 can be attributed to an increase in f₁ which suppressed the growth of the outer interface perturbations. On the other hand, a decrease in N₉₁ from 6 to 1 at Ca₁ > 1 can be attributed to the change in the formation regime of inner drops. For Ca₁ < 1, inner drops were formed in the dripping regime very near to the orifice. In addition, We₁ was relatively low (0.006–0.139), implying that the inertia of the inner fluid was negligible, and the drop size depended on the interplay between interfacial and viscous forces. At higher We₁ values (0.139–0.324), the inertia of the inner fluid was higher and the inner drops were formed in the transitional regime between dripping and widening jetting. As a result, the inner drops pinched off further downstream, although they were still highly uniform, with a coefficient of variation of their sizes of less than 2%. A transition from dripping to widening jetting occurred at We₁ ≈ 0.60 [23]. At We₁ = 0.96, both inner and outer drops were formed in the widening jetting mode (Fig. 5), characterised by much larger drops than in the dripping mode [23,32]. A comprehensive parametric study of the effects of fluid flow rates, fluid properties, and device geometry on the size of inner and outer drops in single-step microfluidic emulsification can be found in our previous work [23].

3.3. Production of polymeric capsules from stable and unstable inner drops

In this section, additional level of control over the size and internal structure of double emulsion drops has been achieved by controlled coalescence of inner drops. First, large monodispersed dual-core double emulsion drops were produced in dripping-to-widening jetting transitional regime by decreasing the amount of oil-soluble surfactant in the middle phase. The inner drops then coalesced within the outer drops to form monodispersed core/shell drops with thin shells. The core/shell drops formed in the dripping-to-widening jetting transitional regime are larger than the core/shell drops generated in dripping mode and can be produced at higher rates, which can be highly beneficial. The inner phase was 15 wt% aqueous solution of K₂CO₃ with trace amounts of the pH indicator m-cresol purple. The middle phase was SEMISCOSIL 949 UV A/B, a UV curable silicone adhesive, containing 0.5–2 wt% of DC 749 surfactant. The outer phase was an aqueous solution containing 70 wt% of glycerol and 1 wt% of PF-127. After generation of dual-core double emulsion drops, the middle phase was polymerised “on-the-fly” using a UV-A light source (250 W Hoene UVAHAND 250 Mobile UV lamp). The minimum energy density and UV light irradiance needed for successful photopolymerisation
were 2 J cm\(^{-2}\) and 13.8 mW cm\(^{-2}\), respectively. The morphology of the polymerised microcapsules depended on the stability of the inner drops during curing in the collection vial, which has been manipulated by changing the concentration of DC 749 surfactant in the middle phase.

When the surfactant concentration in the middle phase was 2 wt\%, the inner drops were stable and the capsules made from such drops were non-spherical with two internal cavities (Fig. 7a). When the surfactant concentration in the middle phase was 0.5 wt\%, the inner drops coalesced within the middle phase after generation of dual-core drops, but before shell curing. Therefore, destabilizing inner drops deliberately through insufficient amount of lipophilic surfactant in the middle phase is a promising approach for fabrication of relatively large single-compartment microcapsules with thin shells (Figs. 7b).

These large single-compartment silicon rubber capsules with encapsulated CO\(_2\) absorbent are highly desirable for CO\(_2\) capture in fluidised and packed bed systems due to their large gas/liquid interfacial area per unit volume (\(1.1 \times 10^{-9}\) m\(^3\)/m\(^2\)) for the capsules in Fig. 7b as compared to 220–450 m\(^2\)/m\(^3\) for typical structured packing). Also, silicon rubber has a high permeability to CO\(_2\) of 323 \(\times 10^{-9}\) cm\(^3\) cm/(s cm\(^{-2}\) cmHg) \([25,26]\). Small capsules suffer from poor fluidising properties, due to strong particle–particle adhesion, and can cause large pressure drops in packed bed systems. Therefore, microfluidic encapsulation of CO\(_2\) solvents within gas-permeable polymer shells in dripping-to-widening jetting droplet formation regime could be a promising tool to fabricate large capsules with a high capture capacity of typical liquid CO\(_2\) solvents, which are suitable for use in fluidised and packed bed reactors.

### 4. Conclusions

The single-step formation of double emulsion drops with controlled number of inner cores in three phase glass capillary devices has been investigated using high-speed imaging combined with a micro-PIV system. Phase diagrams have been constructed based on the capillary numbers of the inner, middle and outer fluid to map out the regions where monodispersed double emulsion drops with specified number of inner drops can be generated. The pinch-off process of multi-core double emulsion was caused by a combination of high capillary pressure in the neck accompanied by the presence of a vortex flow. The formation of outer drops with multiple inner drops was achieved by decreasing the capillary number of the outer phase and increasing the capillary number of the middle phase. A stable continuous jet of middle phase loaded with uniform inner drops was formed at high capillary numbers of the middle phase, which could be exploited for fabrication of hybrid microfibres. At the capillary number of the inner fluid between 0.2 and 1, the inner drops were formed in the dripping regime and any increase in \(C_b\) led to an increase in the number of inner drops. At \(C_b = 1–2\), double emulsion was formed in the dripping-to-widening jetting transitional mode and any increase in \(C_a\) within this range resulted in the smaller number of inner drops. The maximum number of stable inner drops that could be formed within monodispersed outer drops was six.

The multiple emulsion drops consisting of two inner drops were used as templates to produce polymeric microcapsules using “on-the-fly” photopolymerisation. The capsule morphology was controlled by manipulating the stability of the inner drops through adjusting the concentration of the lipophilic surfactant in the middle phase. Non-spherical two-compartment capsules were synthesised at the surfactant concentration of 2 wt\% due to high stability of the inner drops. A reduction in the surfactant concentration to 0.5 wt\% led to destabilization and coalescence of inner drops and formation of large uniform single-compartment capsules. Multi-core double emulsion drops with unstable cores can be used as an alternative to millifluidic systems for fabrication of large core/shell microcapsules. Large capsules with gas-permeable shell and a gas absorbent in the core are highly desirable for gas absorption processes in packed bed absorption columns due to high gas/liquid interfacial area per unit volume and low operating pressure drops.

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### Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.cie.2017.04.008.

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