Ultrashigh-Throughput Production of Monodisperse and Multifunctional Janus Microparticles Using in-Air Microfluidics

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Supporting Information

ABSTRACT: Compartmentalized Janus microparticles advance many applications ranging from chemical synthesis to consumer electronics. Although these particles can be accurately manufactured using microfluidic droplet generators, the per-nozzle throughputs are relatively low (≈μL/min). Here, we use “in-air microfluidics” to combine liquid microjets in midair, thereby enabling orders of magnitude faster production of Janus microparticles (≈mL/min) as compared to chip-based microfluidics. Monodisperse Janus microparticles with diameters between 50 and 500 μm, tunable compartment sizes, and functional cargo are controllably produced. Furthermore, these microparticles are designed as magnetically steerable microreactors, which represents a novel tool to perform enzymatic cascade reactions within continuous fluid flows.

KEYWORDS: liquid jets, compartmentalization, hydrogels, magnetic steering, microreactors

The anisotropic nature of Janus microparticles offers unique advantages for numerous applications. Their physical and/or chemical asymmetry has been leveraged to engineer, among others, amphiphilic particulate surfactants for Pickering emulsification, multicolored voxels for displays, compartmentalized microreactors as artificial cell mimics, and steerable microswimmers for targeted cargo delivery.

Traditionally, Janus microparticles are fabricated by immobilizing and partially modifying a particle’s surface on an interface in a 2D manner, which limits the microparticles’ morphological complexity. More advanced 3D compartmentalized Janus particles can be produced by in-line manipulating, combining, dispersing, and solidifying precursor solutions. This can be achieved using, for example, centrifugation-based dispensing, electrolydrodynamic cojetting, and microfluidic droplet generation. Although microfluidics-based production methods offer the highest control over particle morphology and complexity, they are limited by relatively low throughputs, which has been challenging their use in large-scale applications. Specifically, to yield a monodisperse product, conventional microfluidic droplet generators are restricted to operating in the dripping regime, which typically limits the per-nozzle microparticle production rates to 1 to 10 μL/min. The parallelization of these chips is the main route toward scalable fabrication and parallelization, but this strategy is highly complex and still limited to <1000 channels.

Although higher microfluidic throughputs can be achieved jetting, this approach is associated with uncontrolled jet breakup causing microparticle polydispersity. Off-chip jetting methods such as inkjet and vibrating jet technologies, have also been exploited for the rapid formation of monodisperse particles, but these are isotropic (i.e., not Janus) rather than compartmentalized Janus microparticles.

In this work, we demonstrate the ultrahigh-throughput on-the-fly fabrication of compartmentalized Janus hydrogel microreactors using “in-air microfluidics” (IAMF). IAMF is based on the controlled collision of at least two liquid microjets in midair, thereby acting as the faster and chip-free counterpart of channel-based microfluidic approaches (Figure S1). Here, we have exploited this technology for the first time to produce monodisperse Janus microparticles and microfibers by combining three liquid microjets in midair, which allowed for per-nozzle production rates that exceeded conventional chip-based droplet microfluidics by 2 orders of magnitude. To demonstrate the microparticles’ functionality, we coencapsulated paramagnetic microparticles and a combination of enzymes in the distinct Janus microcompartments. The resulting particles were leveraged as magnetically steerable microreactors, which represents a novel and effective tool to
Figure 1. Concept of 3-jet in-air microfluidics (IAMF) for Janus microparticle production. (a) A liquid Janus microjet is formed by coalescing two distinct liquid microjets (green and red) that contain a solid precursor. Piezo-electric microjet actuation generates a train of monodisperse Janus droplets that are subsequently coalesced with a third microjet (magenta) that contains a cross-linker. (b) The key stages of Janus microparticle formation during flight include: (1) Jet–jet and drop–jet coalescence, which requires $W_{\text{impact}} \leq 1$; (2) surface tension gradient-driven encapsulation of the Janus droplets, which requires $\sigma_{1} = \sigma_{2} > \sigma_{3}$ and (3) solidification, such that $\tau_{\text{encaps}} < \tau_{\text{sol}} < \tau_{\text{in-air}}$. (c) High-speed microphotograph of two microjets forming a Janus microjet (red and green) with limited initial diffusion. (d) The third microjet (magenta) can encapsulate the in-air formed train of monodisperse droplets, as revealed by high-speed (fluorescence) microscopy. (e) The collected microparticles are characterized by (f) a monodisperse size distribution, as well as (g) a clear compartmentalized morphology. The dotted blue curves indicate that mixed area $\sim 4$X smaller than the total microgel area, which corresponds to a mixed volume of $\sim 1/8$ of the microgel’s total volume.

To enable the production of compartmentalized Janus particles using IAMF, we pioneered the collision of three liquid microjets in midair (Figure 1a,b and Figure S2). First, a Janus microjet was formed by colliding two precursor solution microjets at a collisional angle $\theta$ smaller than $45^\circ$. This resulted in an impact Weber number $W_{\text{impact}} = \rho D (V \sin \theta)^2 / \sigma \lesssim 10$, with $\rho$, $D$, $V$, and $\sigma$ denoting the liquid microjet’s density, diameter, speed, and surface tension, respectively. The modest $W_{\text{impact}}$ effectively enabled liquid coalescence, while preventing “separation”, “splashing”, and “bouncing” effects. High-speed microphotography of colliding colored precursor solutions confirmed the formation of a compound microjet that was composed of two coflowing liquids, herein referred to as the “Janus microjet” (Figure 1c).

By superimposing a periodic wave on a microjet using a piezo-actuated nozzle, a stream of monodisperse droplets can be generated. Here, we demonstrate that controlled breakup of the Janus microjet is also possible by vibrating only one of the source nozzles, as surface waves continuously travel along the microjet after coalescence with another liquid microjet. Rapid-in-air stabilization of the Janus droplets was essential to prevent in-flight and postflight droplet merging. As alginate can be rapidly solidified within milliseconds by cross-linking it with divalent cations, it was selected as the prepolymer solution for the droplets. To induce on-the-fly solidification, the alginate containing Janus droplets were coalesced with a third liquid microjet (collisional angle $\theta < 45^\circ$) that contained Ca$^{2+}$ as the cross-linker. To prevent any wasting of liquids during the in-air formation process due to asynchronous droplet streams (i.e., “drop–drop” mode), in-air coalescence events always included at least one intact microjet, herein referred to as “drop–jet” and “jet–jet” mode (Figure S3).

To produce spherical particles, the Janus precursor droplet must be encapsulated (i.e., completely covered) by the cross-linker liquid. Therefore, the surface tension of the cross-linker microjet was reduced, which induced a Marangoni flow that drove rapid spreading of the Ca$^{2+}$ solution over the Janus precursor droplet surface. In contrast, the surface tensions of the Janus microjets’ polymer precursor solutions were matched to prevent the formation of bowl-shaped or core–shell microparticles. Drop-jet coalescence and encapsulation occur on a time scale $\tau_{\text{e}} \sim (\rho D^2 / \Delta \sigma)^{1/3}$, which typically is a few milliseconds. This is faster than the in-air time of droplets (i.e., $\sim 10$ to $100$ ms), which grants the on-the-fly encapsulation of the polymer precursor droplets. Indeed, Janus microparticles were successfully produced by sequentially coalescing two 0.5% alginate containing microjets and a third microjet comprising 0.2 M CaCl$_2$ and 10% ethanol (Figure 1e). Particle size analysis revealed a monodisperse size distribution, characterized by a coefficient of variation (CV) of 6% (Figure 1f).

To maintain the microparticles’ compartmentalization, it is important that the Janus droplets are solidified before significant mixing occurs. As the alginate precursor solutions are miscible, the two compartments start mixing by diffusion upon first contact over a depth of $\delta_{m} = \sqrt{D_{m} t}$, where $D_{m} \approx 1 \times 10^{-9}$ m$^2$ s$^{-1}$ denotes the diffusion of alginate. Solidification effectively stops the mixing of the alginate compartments and starts upon coalescence with the cross-linker microjet, which is typically $\sim 1$ ms after Janus microjet formation. The solidification rate is limited by the diffusion of Ca$^{2+}$ into the alginate droplet over a depth of $\delta_{c} = \sqrt{D_{c} t}$, where $D_{c} \approx 1 \times 10^{-9}$ m$^2$ s$^{-1}$ denotes the diffusion constant of Ca$^{2+}$. The outer shell of the particle has minimal mass diffusion, because it immediately solidifies. As both diffusion fronts advance at similar rate ($D_{m} \approx D_{c}$), the particle contains a partially mixed core, which is described by a diameter $D_{\text{particle}} / 2$. Therefore, the mixed volume $V_{m} \approx (D / 2)^3 \approx 1 / 8$, leaving $\sim 7 / 8$ of the particle prismatically compartmentalized. The majority of the alginate Janus particle’s compartmentalization is thus contained, as confirmed using cross-sectional analysis of Janus particles with coencapsulated red and green fluorescent nanoparticles (Figure 1g).
By operating the IAMF setup in “drop–jet” mode (Figure 2a), Janus microparticles with controlled sizes of 74 ± 6 μm (CV = 7.7%), 162 ± 10 μm (CV = 6.3%), and 488 ± 34 μm (CV = 7.0%), were readily produced by setting the microjets’ diameters $D_1 = D_2 = 50, 100,$ and 150 μm, respectively (Figure 2b, c and Figure S4). The relative compartment size of Janus microparticles could also be altered on-the-fly, simply by tuning the flow ratio of the first and second (i.e., precursor) microjet ($Q_{1,2}$), which was confirmed by measuring the relative intensities of the distinctly (red and green) fluorescently labeled Janus compartments (Figure 2d).

Plotting the data points from this study together with a representative selection of literature studies, confirmed that IAMF-based production of Janus microparticles is two to 3 orders of magnitude faster as compared to chip-based droplet microfluidics (Figure 2e). IAMF could typically produce tens of milliliters suspension containing more than 50% (v/v) Janus microparticles using a single setup (i.e., not parallelized) in a few minutes, (Figure S5). This throughput is inherent to jetting into air, for which a liquid Weber number $We = \rho D V^2/\sigma > 4$ is required. In contrast, conventional chip-based droplet microfluidics is typically operated in the dripping regime, for which $We < 4$ and Capillary numbers $Ca = \mu V/\sigma \lesssim 0.1$ are required (but usually $Ca \ll 0.1$ and $We \ll 1$ are used), to yield monodisperse microdroplets and microparticles.\(^{13,28}\)

Importantly, parallelization of microfluidic Janus droplet generators has remained scarce and did not yet reach industrial-scale throughputs.\(^{12}\) Notably, a single IAMF setup exceeded the throughput of a previously reported 128 times parallelized chip-based microfluidic Janus droplet generator. The maximum throughput of IAMF-based Janus microparticle production is bound by a gas Weber number $We_g = (\rho_g/\rho_l) We \sim 0.2$, where $\rho_g$ and $\rho_l$ are the densities of the liquid jet and the surrounding gas, respectively, above which uncontrolled wind-induced breakup is expected to occur.\(^{29}\)

In addition to Janus microparticles, various types of microfibers with tunable Janus morphologies could be readily produced by straightforwardly changing the position of the microjets. Positioning the cross-linker microjet upstream of the Janus polymer precursor microjet’s breakup point (i.e., “jet–jet” mode), a straight Janus microfiber was formed instead of microfibers (Figure 2f). Furthermore, positioning the third microjet only a few droplet diameters before the precursor microjets’ breakup point enabled the production of periodically beaded Janus microfibers (Figure 2g). Janus microfibers composed of continuous microfibers containing discrete periodic compartments (i.e., patches) could be produced by positioning the second microjet such that it broke up into a...
droplet train before colliding with the second and third microjet (Figure 2h). Thus, only changing a single parameter (i.e., the interjet position) enabled on-the-fly switching between producing fibrous and particulate matter (Movie S1).

To demonstrate the functionality of the compartmentalized Janus microparticles, we set out to produce steerable microreactors. Particularly, we created microparticles with a physicochemical anisotropy by adding paramagnetic micro-particles to one of the solid precursor microjets. The resulting Janus microparticles exhibited paramagnetic properties that allowed for their spatial manipulation and orientation within a static carrier solution in a noninvasive manner via exposure to a magnetic field (Figure 3a, b). We then investigated whether the paramagnetic Janus particles could also be positioned within flowing liquids, which is essential for their application as steerable microreactors to facilitate a multienzymatic cascade reaction within a laminar coflow of β-D-glucose, H₂O₂, and DAB. (g) The enzymes glucose oxidase (GOX) and horseradish peroxidase (HRP) can together drive a cascade reaction via the sequential enzymatic production and reaction of H₂O₂. In the presence of β-D-glucose and of 3,3′-diaminobenzidine (DAB), a successful cascade reaction is indicated by the formation of poly(DAB), which appears as a brown precipitate. (f) Janus microparticles with paramagnetic and GOX/HRP compartments could be leveraged as steerable microreactors to facilitate a multienzymatic cascade reaction within a laminar coflow of β-D-glucose, H₂O₂, and DAB. (g) Compartmentalization of the brown paramagnetic particles and the enzymes was key to enable accurate quantification of the enzymatic cascade reaction product using absorption spectrophotometry. Furthermore, spectrophotometric analysis of various reagent combinations confirmed that both β-D-glucose and DAB were essential to complete the cascade reaction. "ns" indicates no significance. * indicates significance with p < 0.01.

Figure 3. Multifunctional Janus microparticles as magnetically steerable enzymatic microreactors. Using a magnetic field, Janus microparticles with a paramagnetic compartment could be (a) moved and (b) oriented within a static carrier liquid. (c, d) The same particles could also be trapped and moved with micrometer precision within a laminar coflowing liquid. (e) The enzymes glucose oxidase (GOX) and horseradish peroxidase (HRP) can together drive a cascade reaction via the sequential enzymatic production and reaction of H₂O₂. In the presence of β-D-glucose and of 3,3′-diaminobenzidine (DAB), a successful cascade reaction is indicated by the formation of poly(DAB), which appears as a brown precipitate. (f) Janus microparticles with paramagnetic and GOX/HRP compartments could be leveraged as steerable microreactors to facilitate a multienzymatic cascade reaction within a laminar coflow of β-D-glucose, H₂O₂, and DAB. (g) Compartmentalization of the brown paramagnetic particles and the enzymes was key to enable accurate quantification of the enzymatic cascade reaction product using absorption spectrophotometry. Furthermore, spectrophotometric analysis of various reagent combinations confirmed that both β-D-glucose and DAB were essential to complete the cascade reaction. "ns" indicates no significance. * indicates significance with p < 0.01.
cascade microreactors as powerful tools for, for example, complex sequential enzymatic reactions.\textsuperscript{30}

The steerable Janus microparticles were endowed with a chemically functional domain to yield multifunctional particles that could act as steerable multienzymatic microreactors. To achieve this, we discretely coencapsulated paramagnetic microparticles and a combination of two enzymes to enable a multienzymatic cascade reaction, which was accomplished by moving the steerable enzyme-based microreactors across multiple reagent fluid flows. To this end, the enzymes glucose oxidase (GOX) and horseradish peroxidase (HRP) were selected. GOX catalyzes the oxidation of $\beta$-D-glucose to form $\beta$-D-glucono-1,5-lactone and hydrogen peroxide ($\text{H}_2\text{O}_2$), whereas HRP catalyzes the $\text{H}_2\text{O}_2$-induced oxidation of the chromogenic substrate 3,3'-diaminobenzidine (DAB) to form poly-DAB). Poly(DAB) precipitates and was used to visualize the successful enzymatic cascade reaction based on the colorimetric colorless-to-brown shift (Figure 3e).

The paramagnetic Janus microparticles were positioned inside the channel of a microfluidic device through which a $\beta$-D-glucose solution, water, and a DAB solution were laminarily coflowed. Using a magnetic field, the Janus particles were sequentially trapped within and moved across the reagent flow lines, thereby effectively acting as steerable enzymatic microreactors. A successful enzymatic cascade reaction occurred in the microparticles by sequentially incubating them within the glucose and DAB containing streams, as revealed by the formation of brown precipitate (Figure 3f). Importantly, the compartmentalized Janus design was essential for the accurate quantification of the multienzymatic reaction product. Homogeneous encapsulation of the brown paramagnetic particles hindered the accurate visual readout of the multienzymatic reaction product. In fact, spectrophotometric analysis of the negative control (i.e., without glucose and DAB) revealed a high absorption (i.e., false positive) in the presence of paramagnetic microparticles (Figure 3g). Therefore, spatially distinct encapsulation of the paramagnetic particles and enzymes in the respective Janus compartments was key to enable an accurate visual readout of the colorimetric assay’s reaction product. The multienzyme dependency of the reaction was confirmed by incubating the paramagnetic Janus microparticles with $\beta$-D-glucose or DAB alone (Figure 3g). As magnetic manipulation is easily automated and integrated in large-scale applications, our IAMF-produced steerable microreactors represent a promising tool to upscale multienzyme microreactor chemistry.

In conclusion, we pioneered a feasible method for the ultrahigh-throughput production of various Janus microparticles and microfibers, of which the function, size, and compartmentalization can be tailored and spatially organized. By using IAMF-based droplet generation, the per-nozzle Janus particle production rate was increased by two to 3 orders of magnitude as compared to conventional chip-based droplet microfluidic platforms, which is expected to facilitate IAMF’s integration into large-scale industrial applications. To demonstrate the functionality of our approach, we produced, for the first time, magnetically steerable multienzymatic microreactors by coencapsulating magnetic microparticles and multiple enzymes into distinct compartments of Janus microparticles. This approach yielded physicochemical anisotropic microparticles that acted as magnetically steerable microreactors for enzymatic cascade reactions. The facile, ultrahigh-throughput, and multifunctional nature of IAMF-based Janus production platform primes this novel technology for its widespread integration in industrial and clinical applications, for example, to facilitate multienzymatic reactions in continuous flow systems.

\section*{ASSOCIATED CONTENT}
\section*{Supporting Information}

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsami.8b05227. Experimental Section, Figures S1–S5, Table S1 (PDF)

Movie S1 (AVI)

Movie S2 (AVI)

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\subsection*{Notes}

The authors declare the following competing financial interest(s): T.K., M.K., and C.W.V. are co-inventors on a patent based on this work filed by the University of Twente (WO2017167798). T.K. and C.W.V. are co-founders and directors of the company IamFluidics BV, which is commercializing the IAMF technology. The authors declare no other conflicting interests.

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