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PAPER

View Article Online
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Cite this: Lab Chip, 2022, 22, 3025

Horseshoe lamination mixer (HLM) sets new standards in the production of monodisperse lipid nanoparticles†

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Microfluidic mixers promise unique conditions for the controlled and continuous preparation of nanoparticles by antisolvent precipitation. Nanoparticles may enable encapsulation of drug or mRNA molecules in the form of carrier nanoparticles or can provide higher bioavailability in the form of drug nanoparticles. The ultimate goal in microfluidic approaches is the production of nanoparticles with narrow size distributions while avoiding contaminations and achieving sufficiently high throughput. To achieve this, a novel microfluidic precipitation device was developed and realized by two-photon polymerization: mixing elements were designed in such a way that the liquids undergo a repeated Smale horseshoe transformation resulting in an increased interfacial area and mixing times of less than 10 ms. These elements and an additional 3D flow focusing ensure that no organic phase is exposed to the channel walls. The integration of a fluidic shield layer in the flow focusing proved to be useful to delay the precipitation process until reaching a sufficient distance to the injection nozzle. Lipid nanoparticle preparation with different concentrations of castor oil or the hard fat Softisan® 100 were performed at different flow rates and mixing ratios with and without a shield layer. Flow rates of up to $800~\mu l~min^{-1}$ and organic phase mixing ratios of up to 20% resulted in particle sizes ranging from 42 nm to 166 nm with polydispersity indices from 0.04 to 0.11, indicating very narrowly distributed, and in most cases even monodisperse, nanoparticles. The occurrence of fouling can be completely suppressed with this new type of mixing elements, as long as Dean vortices are prevented. Moreover, this parameter range in the horseshoe lamination mixer provided a stable and continuous process, which enables a scalable production.

Received 14th March 2022, Accepted 4th July 2022

DOI: 10.1039/d2lc00240j

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1 Introduction

Micromixers play an important role in biomedical diagnostics or in the microscale production of nanomedicines. Mixing can be accelerated in microscopic channels due to the large surface-to-volume ratio of the liquids involved, making the mixing time identical or even smaller than the reaction time of chemical or physical processes (*e.g.*, precipitation) and thereby allowing better control of the products. However, laminar flow conditions are maintained in most microflows. In the absence of turbulence, diffusion transport is the dominant factor for mixing. Design configurations of the

Passive micromixers have become increasingly popular in the field of nanoparticle preparation in the last decades and were used for the production of polymeric nanoparticles, ^{6,7} drug nanoparticles, ^{8,9} liposomes ^{10,11} or lipid nanoparticles (LNPs). ^{12,13} The formulation of drug nanoparticles or nanoparticulate drug carrier systems represents a common pharmaceutical approach for improving the bioavailability of poorly soluble drugs after oral administration or to process them into injectable products. ^{14–17} Moreover, drug carrier nanoparticles are under intensive investigation with regard to site-specific drug delivery (drug targeting) after intravenous injection, *e.g.* for the treatment of cancer cells. ^{14,18–20} Nucleic acids encapsulated in cationic lipid-containing nanoparticles can be applied for vaccination (a prominent example being the mRNA-based COVID-19 vaccines) and are being

microchannels can reduce diffusion paths by creating liquid lamellae through advection or distributive mixing. Such mixing processes are either actively supported by an external energy input or passively supported by static geometric features that manipulate the streamlines upstream or in the mixing channel.^{1,5}

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[†] Electronic supplementary information (ESI) available. See DOI: https://doi.org/

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intensively explored for a wide range of therapeutic applications such as the treatment of cancer, infections, or hereditary diseases.²¹ Microfluidic systems are a good option for the production of such nanoparticle formulations. 22-24 The particle size and particle size distribution width are important quality criteria of pharmaceutical nanoparticles, e.g., with regard to the dissolution rate of drug nanoparticles or the biodistribution of nanoparticulate drug carriers. While the dissolution rate of drug nanoparticles always benefits size, 16,17 smallest possible particle intravenously injected pharmacokinetic behavior of nanoparticles is often considered optimal for drug targeting in a size range between 70 nm or 100 nm to 200 nm. 25,26

Pharmaceutical nanoparticles can be prepared in micromixers using the bottom-up method of antisolvent precipitation. Nanoparticle-forming constituents (carrier materials and/or drugs) are dissolved in an appropriate solvent and then thoroughly mixed with an antisolvent, typically water. 16,27 With progressing diffusion, the solutes in the ternary mixture exceed a critical concentration and spontaneous nucleation starts. As nucleation proceeds, the concentration of the substance in the solvent falls below the threshold for nucleation, but growth of nucleated particles can continue to occur. Homogeneous and rapid mixing of the liquids involved in the precipitation process is crucial for producing small particles with a narrow size distribution.²⁸ The latter is of particular importance for nanoparticle dispersions that are prone to Ostwald ripening (such as many drug nanosuspensions) as it counteracts the ripening process.29,30

The ultrafast mixing in microfluidic systems provides optimal conditions for antisolvent precipitation.^{8,31-33} While several microfluidic systems already produced LNPs with sizes below 200 nm and allowed to control particle sizes by adjusting process and formulation parameters, the achievement of narrow size distributions appeared to be more challenging. 13,34-36 Another issue with almost all microfluidic precipitation processes is material accumulation at the channel walls, known as fouling, which becomes a serious disadvantage with increasing microchannel surfaceto-volume ratio. Material can attach to the channel wall and continue to grow into microparticles. This ongoing process can lead to changes in the streamlines or even to complete clogging of the system. Microparticles detaching from the channel wall are carried into the nanodispersion and must be filtered out afterwards.³⁷ Furthermore, microparticles being a thousand times bigger than a single nanoparticle reduce the precipitation yield.

Different strategies to minimize fouling have been reported, such as the use of integrated cleaning techniques employing ultrasound, 38 the use of high pressure mixers to generate large shear stresses, 39-41 tailored surface properties of the channel walls, ^{39,42-44} or adapted composition of the solutions involved. 40,41,45 Nozzle configurations with 3D reduce hydrodynamic flow focusing material-wall interactions, while abrupt path and cross-section changes should be minimized. 41,46 An additional protective separation layer through an additional stream 47,48 or mixing by 3D emulsification8 can create a sufficient distance between particles and channel surfaces and improve the quality of the product. Ideal mixing conditions can be achieved with the baker's transformation, in which a high number of layers adding up to a chaotic distribution of all particles - is achieved by repeated stretching, cutting, and stacking of the solutions. 49-52 However, if the wall contact of a phase is to be avoided, cutting as required for the baker's transformation is not feasible in a microfluidic mixer (Fig. 1). Recently, a concept of a microfluidic mixer based on the Smale horseshoe transformation,⁵⁰ where the cutting is replaced by folding, was introduced, but wall contacts of both phases still occur in the described geometry.⁵³

We recently reported on a coaxial lamination mixer (CLM) preventing channel geometries material-wall interactions while ensuring efficient and fast mixing.54 This unique micromixer, fabricated using two-photon polymerization (2PP), includes inlet filters, a nozzle for coaxial injection, and a series of coaxial lamination elements that increase the interfacial area and reduce the layer thickness of the organic phase. Nanodispersions with castor oil and the hard fat Softisan® 100 with particle sizes between 56 nm and 115 nm and polydispersity indices between 0.04 and 0.12 can be prepared with the CLM system. Fouling can be completely prevented by the CLM system within a wider range of flow conditions, but can still occur at high flow rates above 200 µl min⁻¹ and at flow ratios of organic phase to total flow rate above 15%. Processing hard fat at a concentration of 5 mg ml⁻¹ produced material deposits growing at the injection nozzle.

To solve the remaining limitations in microfluidic nanoparticle precipitation and to fully exploit the potential of microfluidics, the present work is based on a completely novel micromixer concept that appears to be feasible by means of 2PP. 3D mixing elements inducing Smale horseshoe transformation offer the possibility of extending the operating window for stable high throughput production of LNPs without sacrificing the major advantage of CLM, which is that wall contacts for the organic lipid solution are avoided. Moreover, a protective fluidic shield layer promises to prevent even the initial contact with the nozzle outlet that was previously identified as the critical area for fouling.

2 Materials and methods

2.1 Femtosecond laser ablation and two-photon polymerization

Borofloat® 33 borosilicate glass (Schott, Mainz, Germany) in the form of 4-inch wafers, was used as the base substrate for the 3D printed channel system. A femtosecond laser system (microSTRUCT c, 3D Micromac AG, Chemnitz, Germany) was used to subdivide the wafer into individual chips after the surface was partially roughened, through holes were created and alignment marks for the 2PP were engraved. Roughening

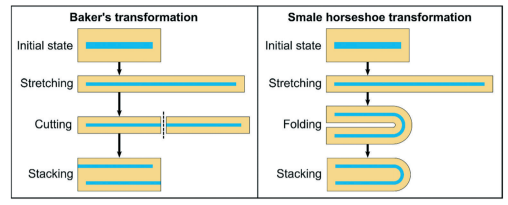


Fig. 1 Schematic representations of the baker's transformation (left) and the Smale horseshoe transformation (right). In both cases, the blue colored fluid is surrounded by the yellow colored fluid in the initial state. Only the cutting process in the baker's transformation causes the blue fluid to contact the wall.

of the glass surface was found necessary for good adhesion of 2PP printed structures. A 2PP system (Photonic Professional GT1, Nanoscribe GmbH, Eggenstein-Leopoldshafen, Germany) equipped with a 25× magnification objective lens (LCI "Plan-Neofluar" 25×/0.8 Imm Korr Ph2, Zeiss, Jena, Germany) was used to 3D print the channel system by dip-in lithography, in which the objective lens is immersed in the resist.

IP-S (Nanoscribe GmbH), a negative-tone resist developed for 2PP, was chosen as the print material, which had a high transparency and hydrophobic surface properties after polymerization. Printing time of a channel system ranged from 14 to 17 hours, depending on the channel design but can in future be reduced with the installation of the next generation of the 2PP device (Photonic Professional GT2, Nanoscribe GmbH). After 2PP printing was complete, any remaining resist on the substrate was removed in a 70 °C developer bath (mr-Dev 600, micro resist technology GmbH, Berlin, Germany), while any remaining resist in the channel system was flushed out by forcing the developer through the substrate inlets. Using a hotplate, the channel system was dried and post-cured at 190 °C for 10 min. Lastly, a UV adhesive (Blufixx PW, Blufixx GmbH, Wesseling, Germany) was cast over the channel system and cured to stabilize and protect the microchannel system from external mechanical stresses or impacts. A more detailed description is given in our previous work.54

2.2 Design of the horseshoe lamination mixers

The coaxial lamination mixer (CLM) in the previous work⁵⁴ allowed fouling-free precipitation at flow rates not exceeding 200 μl min⁻¹. This limited the throughput of the system. The stretching and folding occurs simultaneously in the CLM making the shape and position of the organic phase lamellae not ideally controllable. This resulted in non-uniform diffusion lengths and a closer approach to the channel walls. The horseshoe lamination mixer is therefore designed to separate the stretch and fold processes in time and space.

The horseshoe lamination mixers (HLM) consist of a main channel between the through glass inlet and outlet holes. The inlets are equipped with 2PP printed filter discs with a pore diameter of 20 µm (Fig. 2a) providing protection against particles that can cause clogging of the system.⁵⁴ The channel initially has a round cross-sectional shape with an inner diameter of 200 µm and changes into a rectangular shape after 1 mm. Two systems, A (ESI:† system A view) and B (ESI:† system B view), were designed with slightly different flow focusing sections.

In system A, the rectangular main channel has crosssectional dimensions in height and width of 200 µm × 250 μm and holds a wing-shaped nozzle, which is supplied through a single side channel (Fig. 2b; ESI:† Video S1). For injecting the organic phase (liquid B), the nozzle with a rectangular opening in height and width of 30 \times 200 μm contains a grid of pillars each with a size of 5 μ m \times 10 μ m arranged with a distance of 13.6 µm to each other. The pillars act both as a support structure for the thin wall of the nozzle and as a diffuser to homogenize hydrodynamic pressure and flow velocity, thus achieving a symmetric injection of the organic phase into the main channel.

System B has a main channel with a cross-sectional dimension in height and width of $240 \times 280 \mu m$ (Fig. 2c). The organic phase (liquid B) enters the system through an inlet from one side and is injected through a wing-shaped nozzle with a rectangular opening in height and width of 30 × 200 μm. An additional shield phase (fluidic shield layer, liquid C) comes from the opposite side and is injected to the main channel through a 15 µm wide opening, which completely encloses the injection region of the organic phase. The ringnozzle for this shield phase contains homogenization pillars of the same geometry and spacing as in the organic phase nozzle.

In both systems, a series of thirteen consecutive horseshoe lamination mixing (HLM) elements, each with a length of 650 μm, is located behind the flow focusing section (Fig. 2d). In the first part of these elements, the stretching of the liquid takes place by a transition of channel geometry from a

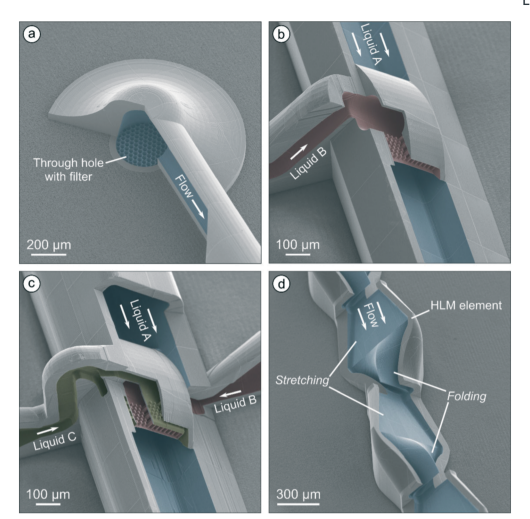


Fig. 2 SEM micrographs of the key components of the HLM systems printed on a glass substrate. The channels were printed for these SEM micrographs as partially open structures to see the interior; colorization inside the channels highlights the liquid flow. a) Channel inlet with filter structure printed on top of a through hole. b) Flow focusing section with the injection nozzle of type A system. Liquid B (red) flows from a side channel into the rectangular nozzle and enters the main channel surrounded by liquid A (blue). c) Flow focusing section of a type B system. Liquid B (red) and C (yellow) flow from two opposing inlets into a nozzle with two openings. Liquid B enters the main channel surrounded by liquid A (blue). d) View on the mixing section with horseshoe lamination mixing elements in series.

rectangular shape of 250 × 100 µm (width and height), to an hourglass shape with a width of 500 µm and a height of 47.5 μ m at the sides and 25 μ m in the center. In the second part, the channel is folded and assumes a rectangular shape with the same dimensions as at the beginning of the HLM element. The repetitive stretching and folding achieves a continuous increase in surface area the organic and aqueous phase simultaneously decreasing the layer thickness of the organic phase. This results in intensified diffusion, and thus, faster mixing. The developed HLM design provides very smooth 3D transition geometries and enables mixing in a very compact format but suitable for increased throughput. The borofloat silicate glass with through holes for the inlets and outlet secures the necessary stability of the 3D printed system (Fig. 3a). The holes of the main channel inlet and the outlet are positioned at a distance of 11.6 mm to each other.

2.3 Flow characterization

2.3.1 Flow simulations. The flow trajectories and concentration distributions were obtained assuming continuous fluidic media which are incompressible and Newtonian. The flow is laminar and in a steady state and unaffected by gravity and other body forces. Simulations based on the continuity eqn (2.1) and the Navier–Stokes momentum eqn (2.2) were performed by computational fluid dynamics using ANSYS FLUENT 20® (ANSYS Inc., Canonsburg, USA).

$$\nabla \cdot \vec{u} = 0 \tag{2.1}$$

$$\rho \vec{u} \cdot \nabla \vec{u} = -\nabla p + \mu \nabla^2 \vec{u} \tag{2.2}$$

where \vec{u} denotes the velocity vector, ρ the fluid density, μ the dynamic viscosity, p the pressure. Mixing of fluids is assumed to be an isothermal process which is analyzed by determining

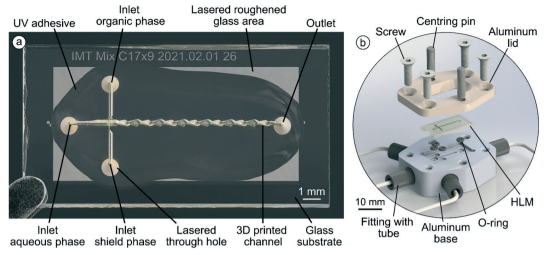


Fig. 3 a) Photograph of the microfluidic glass chip with a 3D printed channel of system B. The chip contains four through holes (3 inlets and 1 outlet). Dust particles on the chip have been digitally removed to not distract the viewer. b) Exploded view of the mounting bracket together with the HLM system.

the solute concentration c in the system based on the convection-diffusion eqn (2.3).55

$$\vec{u} \cdot \nabla c = D \nabla^2 c \tag{2.3}$$

where D is the molecular diffusion coefficient. The working fluids are water and ethanol with a diffusion coefficient of 1.2×10^{-9} m² s⁻¹ at 293.15 K. The mutually influencing properties of the fluids during the mixing are linearly interpolated. Because chemical inertness of the system is assumed and effects of fluid solubilities are neglected, the energy equation is not considered. The viscosity of waterethanol mixtures is given as the share-weighted sum of the viscosities of the two components. Minimal volume changes due to the solution of ethanol in water and any influences of the dissolved or precipitated lipid on the flows are neglected. At the outlet, zero pressure boundary condition was employed, while at both inlets, uniform velocities were employed. No-slip boundary conditions were assumed at the inner walls. 100% water was assigned to the inlet for the aqueous phase and 100% ethanol to inlet for the organic phase.

The computer-aided design of the HLM was discretized by 3D tetrahedral elements with inflation layers. The mesh size was confirmed by mesh convergence observed behind the first two HLM elements (ESI:† Mesh Benchmark and validation for the simulation of concentration distribution, Table S1). The calculation of the numerical analysis was carried out in double precision mode. For the solution methods, the algorithm SIMPLEC (semi implicit pressure linked equation-consistent) was selected coupling pressure and velocity fields. For the calculation the Green-Gauss node mode was used for the gradient discretization, the spatial discretization PRESTO! (PREssure STaggering Option) scheme for the spatial pressure discretization and the third-order MUSCL (Monotonic Upstream-centered

Conservation Laws) scheme for the momentum equation

The mixing quality was quantitatively determined at crosssectional planes normal to the flow direction between the HLM elements using the mixing index:56,57

$$MI = 1 - \sqrt{\frac{\sigma_{\rm m}^2}{\sigma_{\rm max}^2}} \tag{2.4}$$

 $\sigma_{\rm m}$ represents the standard deviation of concentration in a cross-sectional plane at any given place across the channel calculated as

$$\sigma_{\rm m} = \sqrt{\frac{1}{N} \sum_{k=1}^{n} (C_k - \bar{C})^2}$$
 (2.5)

where N is the number of uniformly distributed sampling points, C_k the concentration of the mass fraction at sampling point k, and \bar{C} the concentration averaged over the crosssection. σ_{max} is the maximal standard deviation of the concentration at the specified cross-section along the channel:57

$$\sigma_{\text{max}} = \bar{C}(1 - \bar{C}) \tag{2.6}$$

MI_i shall not only be considered as function of the position of the mixing plane x_i , but also as function of the elapsed residence time $t_{RT,i}$, which represents the average time required for the solution to reach the cross-sectional plane i. $t_{RT,i}$ is calculated as

$$t_{\text{RT},i} = \frac{V_i}{Q_{\text{T}}},\tag{2.7}$$

where V_i is the volume of the channel from the nozzle tip up to the plane i and Q_T is the total volume flow rate.

2.3.2 Experimental analysis of mixing. The mixing time $t_{\rm mix}$ was experimentally determined by means of light

microscopy (VHX 5000, Keyence GmbH, Neu-Isenburg, Germany) based on a chemical solution reaction of iodine with sodium thiosulfate, which becomes invisible as decolorization occurs. The vellow-brown color of the iodine decolorizes to colorless iodide and tetrathionate ions in a quasi-immediate reaction when mixed well with the sodium thiosulfate. This allows a point to be determined for a complete reaction of iodine and sodium thiosulfate, and thus to indicate complete mixing of the solutions. 58,59 Ethanol, absolute (HPLC grade), was purchased from Fisher Scientific (Loughborough, United Kingdom) and purified water was obtained by reverse osmosis and sterile filtration using an water purifier (membraPure, Hennigsdorf, Germany). For the first phase, representing the aqueous phase, 33.9 mg ml⁻¹ sodium thiosulfate (Sigma Aldrich, Steinheim, Germany) was dissolved in purified water and for the second phase, representing the organic phase injected through the nozzle, 50.8 mg ml⁻¹ iodine (Sigma Aldrich, Steinheim, Germany) was dissolved in ethanol. For these studies, channel systems were printed with up to three HLM elements behind the initial flow focusing section. After the last HLM element, the channel keeps its rectangular shape and dimensions of 250 μ m \times 100 μ m (section 2.2). This shape also corresponds to that of the channel without HLM elements. Syringe pumps (Nemesys Base120 + low-pressure modules, Cetoni GmbH, Korbussen, Germany) equipped with glass syringes (2.5 ml glass syringe, Innovative Labor Systeme, Stützerbach, Germany) pumped the liquids into the HLM system at variable total volume flow rates Q_T and fixed ratios φ_{OP} of the organic phase flow rate to the total volume flow rate $Q_{\rm T}$. $\varphi_{\rm OP}$ was initially set to values of 5%, 10%, 15%, and 20% and a high flow rate Q_T was activated so that decolorization of the iodine was not evident. Below a critical flow rate $Q_{T,critical}$, complete decolorization of the iodine can be observed at a certain point behind the nozzle. From the

$$t_{\text{mix}} = \frac{V_{\text{Ch}}}{Q_{\text{T critical}}} \tag{2.8}$$

2.4 Preparation of nanoparticles

 $t_{\rm mix}$ is given as:

2.4.1 Materials. Castor oil, refined (Ph. Eur., Lamotte Oils, Bremen, Germany) and the solid hard fat Softisan® 100 (melting range: 33.5–35.5 °C), a gift from IOI Oleo (Witten, Germany), were used as lipids for the preparation of nanoparticles. The non-ionic surfactant polysorbate 80 (Ph. Eur.) was purchased from Sigma Aldrich (Steinheim, Germany). Purified water (section 2.3.2) was used for the aqueous phase, while ethanol (section 2.3.2) was used for the organic phase and the shield phase. Lipid and surfactant were added to the organic phase, while the shield phase contained only ethanol. The surfactant polysorbate 80 was

CAD design, the channel volume $V_{\rm Ch}$ for the section between

the nozzle and the observation point can be determined and

present in all ethanolic lipid solutions and acted as a stabilizer for the lipid nanodispersions preventing coalescence. Lipid and surfactant concentrations in the respective ethanolic phases can be obtained from Table 1.

The ethanolic solutions and the purified water were filtered with a 0.2 μm syringe filter Puradisc 25 TF (WhatmanTM, 0.2 μm polytetrafluoroethylene, GE Healthcare UK) prior to use. The precipitation of Softisan® 100 nanoparticles leads to nanodispersions in which the particles are present as a supercooled melt. For this reason, the systems are referred to here as nanodispersions rather than solid lipid nanoparticles. The nanodispersions were stored in the dark at room temperature.

2.4.2 Flow control setup. The microfluidic chip was placed into a mounting bracket consisting of a base and a lid, each made of aluminum, with screw connections and O-rings for a tight sealing with the inlets and outlets (Fig. 3b). Each fluidic port is attached to an M6 threaded fitting and was connected via polytetrafluoroethylene tubes to glass syringes, 2.5 ml syringes for the organic phase and the shield phase and a 5 ml syringe for the aqueous phase. Syringe pumps (Nemesys Base 120 + low-pressure modules, Cetoni GmbH, Korbussen, Germany) controlled the volume flow rates of each phase and were automated by a script, which ensured identical preparation procedures for all nanodispersions with the following sequence: 1. fill the channels with the solutions with a standard flow rate and flow rate ratio, 2. drive the currently defined flow rate and flow rate ratio for sample collection, 3. drive a transition time for flushing the output tubing with the nanodispersion, 4. start a waiting time to collect the target volume of 500 µl nanodispersion and 5. run a standard flow rate and flow rate ratio for the transition to the next sample collection. The sample volumes were collected in Eppendorf Safe-Lock tubes (Eppendorf AG, Hamburg, Germany), requiring different precipitation times ranging from 10 seconds to 5 minutes, depending on the total flow rate Q_T . All precipitation experiments were performed in triplicate.

For nanoparticle preparation with system A and different lipid solutions, the total flow rates $Q_{\rm T}$ of 100 μ l min⁻¹, 200 μ l min⁻¹, 400 μ l min⁻¹, 800 μ l min⁻¹, and 1600 μ l min⁻¹ as well as the flow rate ratios $\varphi_{\rm OP}$ of 5%, 10%, and 20% were used. At higher $\varphi_{\rm OP}$, the nanodispersion may become unstable due to the resulting high content of solvent.³⁴

With system B, $Q_{\rm T}$ and $\varphi_{\rm OP}$ were fixed at 100 μ l min⁻¹ and 10%, respectively, while flow rate ratios of the shield phase

Table 1 Concentrations of castor oil/Softisan® 100 and polysorbate 80 in the organic phase

Ethanolic castor oil solution		Ethanolic Softisan® 100 solution	
Castor oil	Polysorbate 80	Softisan® 100	Polysorbate 80
5 mg ml ⁻¹	2.5 mg ml ⁻¹	2 mg ml ⁻¹	1 mg ml ⁻¹
10 mg ml^{-1}	5 mg ml^{-1}	5 mg ml^{-1}	2.5 mg ml^{-1}
20 mg ml^{-1}	10 mg ml^{-1}	-	_
40 mg ml^{-1}	20 mg ml^{-1}		

 $\varphi_{\rm SP}$ of 1%, 3%, 5%, and 7% were used. The resulting flow rates for each phase can be obtained from the ESI† (Table S2). The increase of $\varphi_{\rm SP}$ causes an increase of the overall ethanol concentration in the nanodispersions. $\varphi_{\rm OP}$ was kept fixed to keep the inflow of the organic phase and thus its layer thickness constant. In a second experiment, $\varphi_{\rm OP}$ was fixed at 10% and flow rates $Q_{\rm T}$ of 100 μ l min⁻¹, 200 μ l min⁻¹, 400 μ l min⁻¹, and 800 μ l min⁻¹ were set. $\varphi_{\rm SP}$ of 3% and 5% were used for 2 mg ml⁻¹ and 5 mg ml⁻¹ Softisan® 100, respectively.

2.5 Dynamic light scattering (DLS)

The z-average value and polydispersity index (PDI) of the nanodispersions were determined by dynamic light scattering using a Nanoseries ZS Zetasizer (Malvern Instruments, Malvern, United Kingdom). The z-average represents the intensity-weighted hydrodynamic diameter of the particles. Prior to the measurements, the nanodispersions were diluted with purified water, which had been filtered with a 0.2 µm syringe filter (Whatman® Puradisc 25). The dilution of the samples was adjusted according to the used flow rate ratios $\varphi_{\rm OP}$ and $\varphi_{\rm SP}$. This ensured that a constant concentration of solvent was present in all cuvette samples. A polystyrene cuvette (Omnilab GmbH, Bremen, Germany) contained 1 ml of sample, which was initially equilibrated in the Zetasizer for 5 minutes at 25 °C and then measured three times at a backscatter angle of 173° for 120 s. Particle size parameters were calculated assuming a dynamic viscosity of 0.8872 mPas and a refractive index of 1.330 for water at 25 °C. The obtained z-average values are expected to be slightly underestimated when the influence of the solvent on the viscosity is neglegted.⁴⁵ The samples were measured within two days of their preparation. For comparison with the results from cryo-SEM, the number-weighted diameter was calculated in the Zetasizer on the basis of the Mie theory, 61 assuming a refractive index of 1.46 representative for triglyceride particles (imaginary part 0.01).62

For samples prepared with system A at the flow rate ratios $\varphi_{\rm OP}$ of 5%, 10%, and 20%, the cuvette was loaded with 400 µl, 200 µl, and 100 µl of nanodispersion, respectively, and filled up to 1 ml with purified water. For samples from system B and with both used lipids, the cuvette was loaded with 182 µl, 154 µl, 133 µl, and 118 µl for the flow rate ratios $(\varphi_{\rm OP} + \varphi_{\rm SP})$ 11%, 13%, 15%, and 17%, respectively, and filled up to 1 ml with purified water.

2.6 Electron microscopy

The microchannel components were coated with a thin gold layer before they were imaged by scanning electron microscopy (SEM; Phenom XL, LOT-QuantumDesign GmbH, Darmstadt, Germany) at 10 kV accelerating voltage using the backscattered electron detector.

Nanodispersions were investigated with cryo-SEM. The samples were frozen using a high-pressure freezer (Leica EM ICE, Leica Microsystems GmbH, Wetzlar, Germany) with liquid nitrogen at 2×10^8 Pa. In a high-vacuum coater (Leica EM ACE600, Leica Microsystems GmbH, Wetzlar, Germany) at ~150 °C, the samples were freeze-fractured with a knife and partially sublimated to remove the water-ethanol mixture in order to expose the emulsion droplets. The surface was sputtered with a 4 nm platinum layer. The micrographs were taken in cracks of the platinum layer, as better particle imaging was observed in these areas. The freeze-fractured nanodispersions were imaged by Cryo-SEM (Helios G4 CX, Thermo Fisher Scientific, Waltham, USA) at 2 kV accelerating voltage using the immersion mode. Nanodispersions were investigated after three weeks of preparation.

3 Results and discussion

3.1 Characterization of two-phase flow (system A)

3.1.1 Stretch and fold simulation. Streamline simulations of two-phase flow in system A (which do not take diffusion into account) are displayed in Fig. 4. The injection of the organic phase provides a centered flow focusing and a good enclosure by the aqueous phase. The injected stream is compressed and accelerated by the aqueous phase. As the flow rate ratio φ_{OP} increases, the initial ratio of organic to aqueous phase velocity increases and the organic phase layer grows thicker. When passing an HLM element, the organic phase stretches and folds around the center (ESI:† Video S2). The images in Fig. 5a-d represent the cross-sections of the main channel after the injection, and after two, four and eight HLM elements when diffusion is not taken into account. After the second element, the organic phase appears in four layers (Fig. 5b). The number of layers ideally doubles with each element (Fig. 5c and d), so that after n elements 2^n layers of the organic phase with a thickness that ideally has reduced by a factor of 2^{-n} are found.

To ensure accurate results in the simulation of the concentration distribution based on the convection–diffusion eqn (2.3) a meshing convergence analysis and a validation by a simplified experiment were performed (ESI:† Mesh Benchmark and validation for the simulation of concentration distribution). Fig. 5e–h illustrate simulations of the organic phase concentration distributions in system A, taking interdiffusion into account. In Fig. 5e, the position of the cross-sectional image was set right before the first HLM element. The concentration of the organic phase in its core has reduced to below 0.4. The homogenization increases with each element, and after eight elements, a practically uniform distribution of the organic phase is reached.

However, a small area with lower concentration of the organic phase can still be observed in the channel, which remains even after the eighth HLM element. In the lower right corner of the channel in Fig. 5b, the region is not occupied by the streamlines of the organic phase, and this corner is not reached by diffusion as can be seen in Fig. 5f. The HLM elements that force the folding each have a symmetrical structure, but after the first folding the cross sectional loop is closed on one side and open at the other. As

Folding

50 µm —

Paper

Isometric view Top view Main channel Flow Nozzle Sque on some of the second Streamlines (not colored) organic phase **HLM** element Velocity Injection 0.20 [m/s] 0.15 0.10 Stretching 0.05 0.00 Flow Side view

Fig. 4 Simulated streamlines of the organic phase with color-coding of the flow velocities. The pictures show the injection and the flow through the first HLM element in different views. The aqueous phase streamlines are not shown.

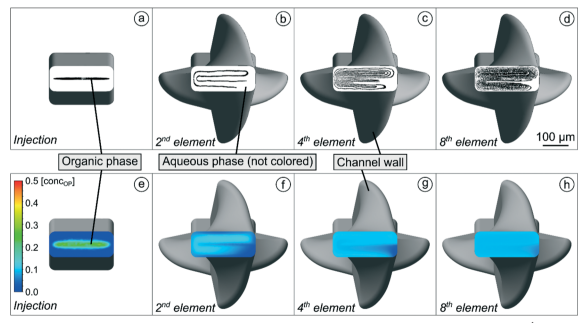


Fig. 5 Cross-sectional views of simulated streamlines of the organic phase obtained at a total flow rate Q_T of 100 μ l min⁻¹ and a flow rate ratio φ_{OP} of the organic phase of 10% in system A. The first row of images illustrates the streamlines behind the injection nozzle (a), and after the second (b), fourth (c) and eighth (d) HLM element. The second row of images shows the concentration of the organic phase behind the injection nozzle (e), and after the second (f), fourth (g) and the eighth (h) element.

a consequence, these ends of the organic phase experience a slightly different local volume flow (ESI:† Video S2). This repeats for each element and leads to this small area that the

organic phase reaches in a delayed manner and that will probably not contribute to precipitation. Nevertheless, the uniform concentration for a large part of the cross-section Lab on a Chip

manage that maining is madely determined by the stretching angle other limited maintains (

proves that mixing is mainly determined by the stretching and folding of the organic phase.

The simulation reveals that the undiluted organic phase does not touch the channel walls during the stretching or folding process. The layer of organic phase and the wall only get closer with each pass through an HLM element (ESI:† Video S2). As long as precipitation of LNPs consumes the lipid content before diffusion reaches the walls, no fouling should be expected.

3.1.2 Simulation of mixing with succesive HLM elements. Fig. 6a shows the layer length l_{OP} and layer thickness d_{OP} of the organic phase after injection and after one, two, three and four HLM elements as obtained from the simulation with system A, not considering diffusion. Streamlines obtained with more than four elements did not allow for clearly identifying the boundaries of the extremely thin layers. l_{OP} increases exponentially by an average factor of 1.93 (ideally assumed as 2) with each element both for φ_{OP} of 10% and 20%, while d_{OP} decreases exponentially by an average factor of 0.56 (ideally assumed as 0.5) with each element for $\varphi_{\rm OP}$ of 10% and 20%. This results in a calculated stretching of $l_{\rm OP}$ from the measured 221 μm after injection by a factor of around 5200 to 1.17 m behind the thirteenth element. Simultaneously, $d_{\rm OP}$ decreases from 8.1 μm before the first HLM element to 5 nm after the thirteenth element at 10% φ_{OP} and from 15.2 $\mu\mathrm{m}$ to 9 nm at 20% φ_{OP} . It is worth mentioning that according to $t_{\text{Diff}} = d_{\text{OP}}^2/2D$, with D as the mutual diffusion coefficient of 0.80 × 10⁻⁹ m² s⁻¹ of a 20 vol% ethanol in water mixture (25 °C), $^{63-65}$ a layer thickness of $d_{\rm OP}$ = 40 nm would equilibrate by diffusion in less than a microsecond. Optimization of the folding process, so that the lateral ends of the organic phase are not retracted, could slightly enhance the efficiency of the stretching to be closer to 2.

Fig. 6b shows the mixing index in dependence on the influence of the elapsed residence time t_{RT} for different Q_{T} . The data points in the curves indicate a cross-sectional plane after each HLM element. MI = 0 is obtained when the mixing has not started, while MI = 1 means the absence of any concentration gradients in the cross-sectional plane. The mixing increases steadily, but the progression of mixing slows down with each element. On one hand, the nozzle with its high aspect ratio and the HLM elements are designed to promote initially fast diffusion between the phases, which decreases with the reduction of concentration gradient in time.⁶⁶ On the other hand, the HLM elements prevent early contact of the organic phase with the channel wall. Additionally, only a small spot with a lower concentration of the organic phase remains even after the eighth element (Fig. 5d). It is located where the folded lines do not reach.

At higher $Q_{\rm T}$, a smaller mixing index is achieved after the same number of HLM elements, because less time is given for diffusion. As a result, more elements have to be passed before mixing is completed. However, this is compensated by the shorter elapsed residence time $t_{\rm RT}$ of the solutions in the channel at higher $Q_{\rm T}$, and results in a shorter mixing time

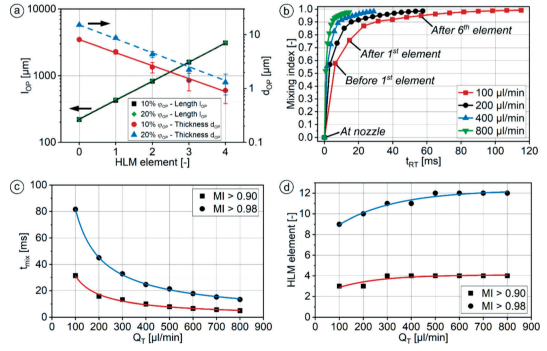


Fig. 6 a) Simulated layer length $l_{\rm OP}$ and thickness $d_{\rm OP}$ of the organic phase in system A. The total flow rate $Q_{\rm T}$ was set to 100 μ l min⁻¹ and the flow rate ratios $\varphi_{\rm OP}$ to 10% and 20%. Data sets were fitted using nonlinear regression, whose R^2 for $l_{\rm OP}$ and $d_{\rm OP}$ deviate from 1 less than 0.001. b) Influence of elapsed residence time $t_{\rm RT}$ and $Q_{\rm T}$ on the mixing index, where $\varphi_{\rm OP}$ was set to 10%. c) Influence of $Q_{\rm T}$ on $t_{\rm mix}$ (the flow residence time after which a mixing index greater than 0.9 or 0.98) is reached in the simulations. The curves are a guide for the eye only. d) The number of HLM elements after which a mixing index greater than 0.9 or 0.98 is reached in the simulations over $Q_{\rm T}$. The curves are a guide for the eye only.

nevertheless (Fig. 6c). The simulations show that good mixing (MI > 0.9) is obtained after 31 ms, 16 ms, 10 ms and 5 ms for the flow rates 100 μ l min⁻¹, 200 μ l min⁻¹, 400 μ l min⁻¹ and 800 ml min⁻¹, respectively. Lower flow rate ratios $\varphi_{\rm OP}$ would further reduce the mixing time due to the lower thickness of the organic phase layer.

The number of HLM elements required to reach mixing indices higher than 0.90 or 0.98 (saturation value for MI) in dependence on $Q_{\rm T}$ is given in Fig. 6d (detailed values in ESI:† Table S3). It can be obtained, regardless of $Q_{\rm T}$, good mixing (MI > 0.9) can already be achieved with only four elements, while nearly perfect mixing can be reached with all thirteen HLM elements. This applies with $\varphi_{\rm OP}$ of 10%, and with the flow conditions considered in the simulations, *i.e.*, without formation of secondary flows at high Reynolds numbers.

3.1.3 Evaluation of mixing experiments. Fig. 7 shows the times t_{mix} as experimentally obtained decolorization of iodine when mixed with sodium thiosulfate in dependence on the number of HLM elements for varied flow rate ratios φ_{OP} (ESI:† Table S4). In a channel with no HLM element, the iodine solution decolorized after 67 ms and after 1902 ms at φ_{OP} of 5% and 20%, respectively. An increase in φ_{OP} resulted in a larger layer thickness d_{OP} and a corresponding large increase in mixing time of several hundred milliseconds; in particular, a doubling of φ_{OP} , and thus of d_{OP} , led to an 5.6-fold average increase of the mixing time (according to $t_{\text{Diff}} = d_{\text{OP}}^2/2D$ with a quadrupling of t_{mix} would be estimated). The integration of one HLM element reduced $t_{\rm mix}$ by a factor of 4.2 on average, so that the time decreased to 16 ms and 373 ms at φ_{OP} of 5% and 20%, respectively. With a second integrated HLM element, $t_{\rm mix}$ was further reduced by a factor of 3.2 on average. As the organic phase is stretched, increased diffusion already occurs between the elements causing subsequent elements to have a progressively smaller effect on reducing t_{mix} . This decreasing effect of additional elements on $t_{
m mix}$ continues. However, in a system with a third HLM element, the further reductions in mixing time range between 1.2 and 2 indicating that for

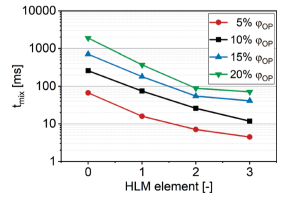


Fig. 7 Mixing time $t_{\rm mix}$ experimentally observed with iodine solution in sodium thiosulfate solution depending on the number of HLM elements inside a channel. $\varphi_{\rm OP}$ were set at 5%, 10%, 15% and 20%.

thicker layers (higher φ_{OP}) the additional element has a higher influence (Fig. 7 and ESI:† Table S4).

During the folding process, parts of the organic phase layer at the folding point remained thicker than the average. These thicker parts were more expressed at higher φ_{OP} in the simulations (data not shown) and showed in the microscope in the form of stripe-like streams of the iodine solution at higher Q_{T} , where diffusion at the observation point was not completed. The resulting slower diffusion explains why the decrease in mixing time was lower at the ratios φ_{OP} of 15% and 20% from the transition of a system of two to three HLM elements.

While $t_{\rm mix}$ obtained from simulations indicates good mixing after 10 ms (MI > 0.9) at a $\varphi_{\rm OP}$ of 10% with $Q_{\rm T}$ of 400 μ l min⁻¹ after the third HLM element (Fig. 6b–d, ESI:† Table S3), the experiment revealed complete mixing at 13 ms at a $\varphi_{\rm OP}$ of 10% with $Q_{\rm T}$ of 410 μ l min⁻¹ after the third HLM element (Fig. 7, ESI:† Table S4).

3.2 Lipid nanoparticle preparation in two-phase flow (system A)

3.2.1 Influence of flow rate ratio and total flow rate on LNP size distribution. The total flow rate $Q_{\rm T}$ and the flow rate ratio $\varphi_{\rm OP}$ can be used to adjust the mixing time (section 3.1.2) and thus the z-average value (Fig. 8). With increasing $Q_{\rm T}$ and thereby faster mixing, particle sizes decreased steadily at all $\varphi_{\rm OP}$. In the experiments, lipid nanoparticles with z-average values ranging from 45 nm to 85 nm and PDI values ranging from 0.04 to 0.23 were produced, while particles with PDI values below 0.07 can be considered monodisperse. This compares with the previously best results obtained with microfluidic antisolvent precipitation of castor oil, with z-average values between 44 nm and 90 nm and PDI values from 0.1 to 0.3, with a bimodal size distribution for the smallest z-average value, as we have recently reported. 34,45

The standard deviations obtained through n=3 experiments for the particle sizes (0.1–0.5 nm) and the PDI values (0.003–0.010) were very small. A respective doubling of $\varphi_{\rm OP}$ from 5% to 10% and to 20% led to a stepwise increase in particle size by an average factor of 1.17 for all $Q_{\rm T}$ below 800 μ l min⁻¹ (Fig. 8a). Due to the increasing layer thickness $d_{\rm OP}$ with increasing $\varphi_{\rm OP}$, the concentration of solvent and lipid in the channel increases, and longer diffusion and mixing times are required. This leads to a slower supersaturation of the lipid, a lower nucleation rate and thus a stronger growth phase causing the formation of larger particles. ⁶⁸

The PDI values increased slightly and steadily with increasing $Q_{\rm T}$ until they varied strongly between different $\varphi_{\rm OP}$ at a $Q_{\rm T}$ of 1600 μ l min⁻¹ (Fig. 8b). At and below 800 μ l min⁻¹, the PDI values did not differ significantly between the ratios $\varphi_{\rm OP}$. Different effects could be responsible for the slow increase of PDI values with increasing $Q_{\rm T}$. At higher $Q_{\rm T}$, the organic phase flows through more HLM elements before reaching complete diffusion. The simulations reveal a decreasing distance of the organic phase from the wall

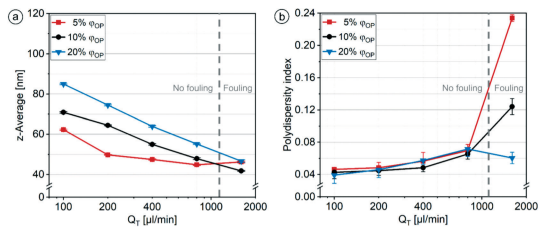


Fig. 8 Influence of total flow rate Q_T and flow rate ratio φ_{QP} on z-average (a) and PDI values (b) (n = 3 experiments, \pm standard deviation). The ethanolic phase contained 5 mg ml⁻¹ castor oil and 2.5 mg ml⁻¹ polysorbate 80. A dashed gray line separates data points of experiments associated with fouling from those where no fouling was observed. PDI graph in (b): standard deviation bars for Q_T below 1600 μ l min⁻¹ only point in one direction for better visibility.

surfaces with each additional HLM element, causing parts of the organic solution to flow more slowly through the channel because of non-slip conditions at the wall. Additionally, the variance in the layer thickness of the organic phase increases with each element due to accumulations of the organic phase at the folding points (section 3.1.1). Both effects lead to higher variance in the speed of mixing and a broader particle size distribution.69

3.2.2 Influence of castor oil concentration on LNP size distribution. An increase in castor oil concentration led to an increase in particle size, while the z-average value decreased with increasing flow rate Q_T (Fig. 9a). At the highest considered castor oil concentration of 40 mg ml⁻¹, z-average values of 142 nm to 166 nm were produced with Q_T between 100 μl min⁻¹ and 800 μl min⁻¹. PDI values in a range of 0.05 to 0.07 at a castor oil concentration of 10 mg ml⁻¹ were almost the same as obtained with 5 mg ml⁻¹ castor oil (Fig. 9b). At a castor oil concentration of 20 mg ml⁻¹ and 40 mg ml⁻¹, the PDI began to increase across all Q_T to values ranging from 0.06 to 0.09 and from 0.08 to 0.11 respectively. A beginning instability of the precipitation process at the nozzle may cause non-uniform nucleation or enhance agglomeration of the precipitated nanoparticles. Moreover, higher lipid concentrations could cause the supersaturation to start earlier but homogenization to be completed later. This could lead to longer periods of nucleus formation and particle growth.⁷⁰ At the used Q_T below 1600 μl min⁻¹, no fouling occurred on the HLM elements at any of the castor oil concentrations used.

Compared to our previous work,54 the HLM system offers the possibility of using higher lipid concentrations while achieving smaller particle sizes and more narrow size distributions. Calculating the production efficiency of the HLM system in the form of lipid processing per hour with the consideration that the PDI should be below 0.07 for monodisperse spherical particles and assuming that all lipid

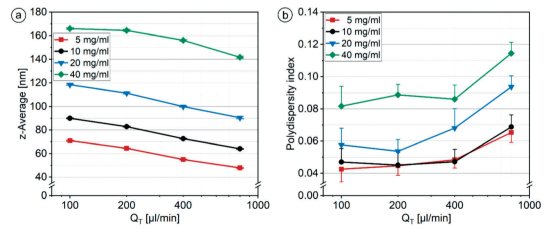


Fig. 9 Influence of total flow rate Q_T and lipid concentration of castor oil on z-average (a) and PDI values (b) $(n = 3 \text{ experiments}, \pm \text{ standard})$ deviation). The flow rate ratio φ_{OP} was set to 10%. The z-average and PDI values for the lipid concentration of 5 mg ml⁻¹ were taken from the result of section 3.2.1. PDI graph: standard deviation bars only point in one direction for better visibility.

precipitates as nanodroplets with a size below 200 nm, this new mixer can process 48 mg h^{-1} (ESI:† Table S5). Compared to our previous work with the CLM and under identical conditions, this results in a 4-fold increase in productivity.

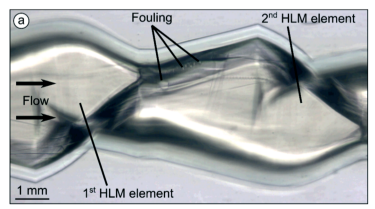
3.2.3 Fouling on nozzle and HLM elements at high flow rates. The flow behavior changed between a total flow rate $Q_{\rm T}$ of 800 μ l min⁻¹ and 1600 μ l min⁻¹, and at a $Q_{\rm T}$ of 1600 μ l min⁻¹ fouling occurred at the first and second HLM elements with all flow rate ratios $\varphi_{\rm OP}$ (Fig. 10a; ESI:† Video S4). Larger lipid droplets were observed at the ends of the flat channels and were dragged along by the flow. This led to a strong increase in the PDI (Fig. 8b). In particular, the PDI value suddenly quadrupled to 0.23 at a $\varphi_{\rm OP}$ of 5% and a $Q_{\rm T}$ of 1600 μ l min⁻¹.

The simulation shows that at a Q_T of 1600 μ l min⁻¹, the organic phase at the folded open ends appears slightly bent over and is in contact with the channel wall (Fig. 10b). The design of the HLM elements causes a helical rotation of the fluids. In the elements at the open ends, centrifugal forces may cause a Dean effect during the folding at sufficiently high flow rates, forming a secondary flow. 4,71,72 The dimensionless Dean number can be used to characterize the strength of the secondary flow in the curved channel and is defined as, De = Re $\sqrt{d_{\rm hydr}/2R}$ where Re is the Reynolds number (Re = $u \cdot d_{hvdr}/v$), u the average velocity of the liquids, d_{hydr} the hydraulic diameter of the channel, R the radius of curvature, and v the kinematic viscosity. The lateral ends of the channel lie on a helical path whose radius varies, and locally reaches a minimum value of $R = 300 \mu m$. For $Q_T =$ 1600 μl min⁻¹, this results in a Reynolds number of 102 with $v_{\text{Water}} = 1 \text{ mm}^2 \text{ s}^{-1}$, $u = 1.48 \text{ m s}^{-1}$, $d_{\text{hydr}} = 68.8 \text{ } \mu\text{m}$ and a Dean number of 35. In the literature, a formation of secondary vortices in the channel with a Reynolds number and a Dean number in this range is described. 73,74 The centrifugal forces start to affect the streamlines of the organic phase above 100 μl min⁻¹, while at 100 μl min⁻¹ and below the streamlines remain unchanged (ESI:† Fig. S5).

To avoid such secondary vortices, either the width of the nozzle can be reduced to avoid the organic phase approaching the channel ends, or the HLM element must be redesigned with a larger radius of curvature during the folding. Simulations with an adapted nozzle and an HLM element, whose folding section was extended by a factor of 4 in distance, show improved flow conditions without contact of the organic phase with the channel wall at $Q_{\rm T}$ of 3200 μ l min⁻¹ and $\varphi_{\rm OP}$ of 5% (ESI:† Video S3). By this extension of the folding section in the HLM element, the Dean number is reduced by a factor of 3.3. Nanoparticle preparations were not performed using the system with extended HLM element, but all following results were still obtained with the HLM element from section 2.2.

While fouling was very clearly pronounced at a φ_{OP} of 5%, only minor fouling with larger isolated microdroplets was observed at a φ_{OP} of 20%. The dynamic viscosity of an ethanol-water mixture with 20 vol% ethanol is around 1.6 times higher than of water. This provides a more stable laminar flow characterized by a lower Reynolds number and thus a lower Dean effect. The simulations show that the contact of the organic phase with the channel wall is slightly lower at a φ_{OP} of 20% than with 5% (data not shown). However, since in the simulations the viscosity of the mixed fluid is linearly interpolated between the values of the pure phases, it is difficult to assess how strong this trend is in reality.

When preparing nanodispersions with castor oil at a concentration of 10 mg ml⁻¹ and above, fouling accumulated slightly on the injection nozzle (Fig. 11a). First deposits formed at the lateral ends of the nozzle, which grew over time. The processing of solid lipids poses an even higher risk of fouling since solid deposits on channel walls will be more stable. Nanoprecipitation with Softisan® 100 revealed fouling directly at the nozzle but not at any of the HLM elements when $Q_{\rm T}$ was smaller than 1600 μ l min⁻¹ (Fig. 11b; ESI:† Video S5). Fouling started at the nozzle edge and grew along the interface between the aqueous and organic phases. As



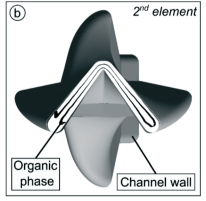
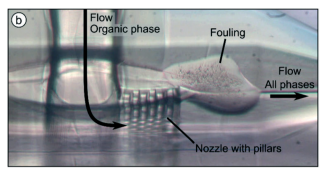


Fig. 10 a) Microscopic image of the first two HLM elements during a precipitation process with 5 mg ml⁻¹ castor oil. The flow rate Q_T and flow rate ratio φ_{QP} of the organic phase were set to 1600 μ l min⁻¹ and 5%, respectively. Positions of fouling, observed after 20 s operation, are indicated. b) Cross-sectional view of simulated streamlines of the organic phase within the second HLM element during the folding process obtained at Q_T of 1600 μ l min⁻¹ and φ_{QP} of 5%.



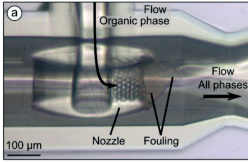


Fig. 11 Microscopic images of the nozzle of system A during a precipitation process with 40 mg ml⁻¹ castor oil (a) and with 5 mg ml⁻¹ Softisan® 100 (b). The total flow rate Q_T and flow rate ratio φ_{QP} of the organic phase were set to 100 μ l min⁻¹ and 10%, respectively. Positions of fouling, observed after 2 min (a) and 9 min (b) operation, are indicated.

growth continued, a tubular structure with decreasing diameter formed, which strongly influenced the mixing process. To prevent the accumulation of fouling on the nozzle, a nozzle with a fluidic shield layer (system B) was developed to enclose the organic phase.

3.3 Microfluidic characterization of three-phase flow (system B)

Simulation of the injection of three phases in system B confirms a complete enclosure of the organic phase by the shield phase (Fig. 12a and b). The thickness of the enclosing shield phase depends on the flow rate ratio φ_{SP} of the shield phase. The lateral ends of the shield phase reveal a greater thickness than its upper and lower sides (Fig. 12c). When the shield phase is injected into the main channel, its width is increased due to height compression by the surrounding aqueous phase. While the uniform distribution of the shield phase decreases if the pillar density in the nozzle is too low, a too high density runs the risk of not being able to clean the channel from the residual photoresist when the system is printed with 2PP. The number and distribution of pillars in the nozzle was chosen to ensure a sufficiently uniform distribution of the shield phase around the organic phase.

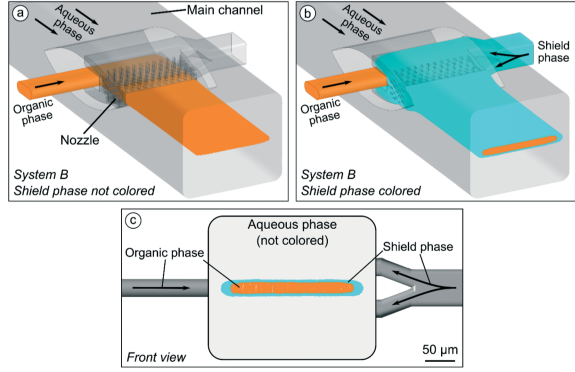


Fig. 12 Visualization of streamline simulation of organic and shield phase at the injection nozzle of system B. The total flow rate Q_T was set to 100 μ l min⁻¹ and the flow rate ratios of the organic phase φ_{OP} and the shield phase φ_{SP} were both set to 10%. In image a), only the organic phase is visualized, while in image b) and c), the shield phase is visualized as well. The simulated streamlines of the aqueous phase are not visible in all images. Vertical white lines in the organic phase are "digital" voids caused by Ansys when distributing the number of streamlines.

Simulations showed that the patterns of the subsequent stretching and folding by the HLM elements did not show any obvious differences compared to system A.

3.4 Comparison of two-phase flow (system A) with three-phase flow (system B)

3.4.1 Effect of fluidic shield layer on fouling. The most severe fouling on the nozzle of system A was observed with high castor oil concentrations and with the use of Softisan® 100 solutions. For this reason, these formulations were used to investigate the effect of the fluidic shield layer. Fouling on the nozzle could be prevented when system B was used (Fig. 13; ESI:† Video S5). The pure solvent of the shield layer initially prevents a supersaturation of the lipid at the interface between the aqueous and organic phase. The phases must first diffuse through the shield layer before precipitation can occur.

The additional shield phase will slightly slow down the mixing with the aqueous phase and will cause an increase of the total concentration of solvent in the nanodispersion. A too high solvent fraction may reduce long-term stability of the nanodispersion. Therefore, the addition of shield phase should be kept to the minimum necessary to prevent fouling in the channel system. The flow rate ratio φ_{SP} of the shield phase was varied to determine the limit of fouling suppression. The risk of fouling increases with higher lipid concentration in the organic phase. The intensity of fouling decreased rapidly with increasing φ_{SP} , until at φ_{SP} of 3%, no fouling could be detected using 20 mg ml⁻¹ and 40 mg ml⁻¹ castor oil solutions and with 2 mg ml-1 Softisan® 100 solution. φ_{SP} of 5% was necessary to prevent fouling when using the 5 mg ml⁻¹ Softisan® 100 solution. Interestingly, fouling, at too low φ_{SP} , formed only on one side of the nozzle, specifically on the opposite side of the inlet for the shield phase. This implies that the organic or shield phase did not flow into the main channel with a perfectly uniform distribution (as predicted by the simulation), but rather

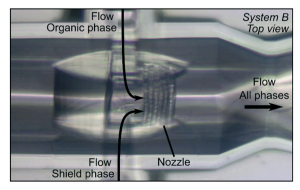


Fig. 13 Microscopic image of the nozzle of system B during the precipitation process of 5 mg ml $^{-1}$ Softisan® 100 after 9 min operation. The total flow rate Q_T was set to 100 μ l min $^{-1}$ and the flow rate ratios φ_{OP} and φ_{SP} were set to 10% and 5%, respectively. No fouling was visible in contrast to system A (Fig. 11b).

followed the path of least resistance from the inlet to the opening of the nozzle. In the simulation, the velocity distributions in the nozzle show that the organic and shield phases leave the nozzle 7% and 2% faster on the side of their respective inlets. Optimizations of the pillar arrangement in the nozzle and the nozzle ring opening could improve the distribution of the shield phase around the organic phase and, thus, fouling might be prevented even at smaller φ_{SP} .

3.4.2 Effect of fluidic shield layer on precipitation process. Fig. 14a shows the z-average values of the castor oil and Softisan® 100 nanoparticles obtained across the investigated flow rate ratios φ_{SP} at a constant total flow rate Q_T of 100 μ l \min^{-1} and a flow rate ratio φ_{OP} of 10%. When φ_{SP} increased at constant φ_{OP} , the z-average values of castor oil particles did not change, while the z-average values of Softisan® 100 particles decreased. When φ_{SP} changed from 1% to 5%, the z-average values decreased from 68 nm to 62 nm and from 105 nm to 83 nm at 2 mg ml⁻¹ and 5 mg ml⁻¹ Softisan® 100 respectively. The shield phase had a positive influence on the PDI values (Fig. 14b). The best reduction in PDI was achieved with the 5 mg ml⁻¹ Softisan® 100 solution with a change from 0.11 to 0.05 when φ_{SP} of 5% was used. For comparison: with system A, the strongest fouling on the nozzle with this lipid was observed at a concentration of 5 mg ml⁻¹.

The results show that the shield phase further improved the performance of the HLM system, especially when solid lipids were precipitated. The suppression of fouling and improved quality of the nanoparticle dispersions demonstrate that flow focusing is a crucial factor for the stable production of lipid nanoparticles with narrow size distribution.

3.4.3 Influence of flow rate ratio and total flow rate on Softisan® 100 nanoparticles. Nanodispersions with Softisan® 100 were prepared with two-phase and three-phase flow to better illustrate the effect of the shield phase on the precipitation process. The lowest identified flow rate ratio $\varphi_{\rm SP}$ with no fouling were used for particle preparation, 3% for 2 mg ml $^{-1}$ and 5% for 5 mg ml $^{-1}$ Softisan® 100. *z*-Average values of 42 nm to 83 nm and PDI values of 0.05 to 0.07 were realized with these Softisan® 100 concentrations, with flow rates $Q_{\rm T}$ of 100 μ l min $^{-1}$ to 800 μ l min $^{-1}$ and a flow rate ratio $\varphi_{\rm OP}$ of 10% used in system B (Fig. 15). In other investigations with solid glycerides, particle sizes were mostly in the range of 100 nm to 300 nm and PDI values of 0.15 to 0.34 were achieved. $^{13,35,60,76-78}$

For both systems under investigation, the z-average values continuously decreased with increasing $Q_{\rm T}$ for all Softisan® 100 concentrations (Fig. 15a), while the nanoparticles without using a shield phase were up to 19 nm larger. For both Softisan® 100 concentrations and with increasing $Q_{\rm T}$, PDI values decreased with system A and increased with system B. Fouling was observed to be lower when higher flow rates were used in system A, possibly due to higher shear stress at the nozzle and thus less change in flow conditions. The increase of the PDI with increasing $Q_{\rm T}$ for system B with Softisan® 100 showed a similar trend as the increase of the PDI for

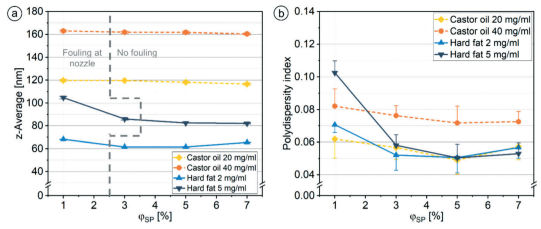


Fig. 14 Influence of the flow rate ratio φ_{SP} of the shield phase and the lipid solution (castor oil and the hard fat Softisan® 100) on z-average (a) and PDI values (b) (n = 3 experiments, \pm standard deviation). The total flow rate Q_T and the flow rate ratio φ_{OP} of the organic phase was set to 100 μl min⁻¹ and 10%, respectively. PDI graph (b): standard deviation bars only point in one direction for better visibility.

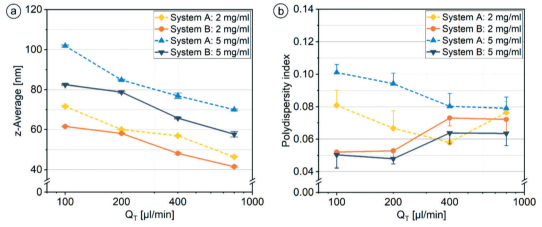


Fig. 15 Influence of total flow rate Q_T and lipid concentration of Softisan® 100 on z-average (a) and PDI values (b) (n = 3 experiments, \pm standard deviation). The flow rate ratio φ_{OP} of the organic phase was set to 10%. PDI graph: standard deviation bars only point in one direction for better visibility.

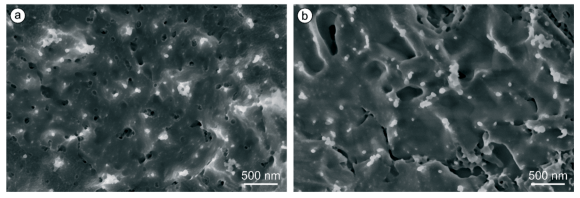


Fig. 16 Cryo-SEM images of nanodispersions prepared with 5 mg ml⁻¹ castor oil (a) and 5 mg ml⁻¹ Softisan® 100 (b). a) The nanodispersion with castor oil was prepared with system A, a total flow rate Q_T of 100 μ l min⁻¹ and a flow ratio φ_{OP} of 5%. b) The nanodispersion with Softisan® 100 was prepared with system B, Q_T of 100 μ l min⁻¹ and flow rate ratios φ_{OP} and φ_{SP} of 10% and 5%, respectively.

system A with castor oil (section 3.2.1), but was a bit stronger (Fig. 15b). The additional fluidic shield layer surrounding the organic phase acts as an additional barrier through which the organic and aqueous phase must diffuse. If the enclosure of the organic phase is not uniform, this variance can add to the variance caused by the stretching and folding process (section 3.1.2) when more HLM elements must be passed as Q_T increases, resulting in a higher PDI at higher Q_T .

3.5 Electron microscopic evaluation of cryo-stabilized nanodispersions

Fig. 16 presents the cryo-SEM images of a castor oil (Fig. 16a) and a Softisan® 100 nanodispersion (Fig. 16b). In addition to clearly (almost fully) visible emulsion droplets, some droplets protrude partially out of the surface only with their crests and appear to have a diameter of a few nanometers. During sublimation of the aqueous phase, individual emulsion droplets penetrate the surface until they are completely exposed. Additionally, larger clusters of agglomerates with lengths of up to 300 nm can be seen in the images. Intensity-weighted particle size distributions from DLS measurements that are especially appropriate to detect larger particle fractions gave no hint for the presence of agglomerates with sizes around 300 nm. Previous work proved shortterm stability of comparable nanodispersions.^{34,54} Thus, agglomerates may have formed during sample preparation. Cryo-TEM, which uses a different sample preparation technique, could be helpful to clarify whether such nanodispersions contain agglomerates or not.⁷⁹

The nanodroplets appear to be homogeneous in size with a roundish shape. The evaluation of 160 castor oil droplets and 100 Softisan droplets from several micrographs (disregarding still embedded particles and agglomerates) reveals diameters of 52 ± 7 nm for castor oil and 69 ± 9 nm for Softisan® 100 nanodroplets. Compared to the DLS results discussed above, the castor oil (Fig. 8a) and Softisan® 100 particles (Fig. 15a) are 10 nm and 14 nm smaller, respectively, according to cryo-SEM. In the routine evaluation of the DLS measurements, the intensity-weighted hydrodynamic diameter of the particles was obtained, while the evaluation of the cryo-SEM images yielded a number-weighted distribution. Thus, the measured z-average values may be slightly larger.80 When the number-weighted hydrodynamic is calculated for the respective measurements, a mean particle size of 48 nm is obtained for the castor oil and 64 nm for the Softisan® 100 nanoparticles. As described in section 2.5, a slight underestimation of the particle size measured by DLS is to be expected for the samples examined here since the presence of ethanol was neglected in the evaluation. Overall, the particle sizes from cryo-SEM and DLS can thus be considered as in good consistence.

4 Conclusions and outlook

A novel horseshoe lamination mixer (HLM) was designed and printed on a glass substrate by two-photon polymerization, which offers the necessary 3D resolution to realize such complex 3D structures of the injection and HLM elements. Typical printing times of several hours per system, depending on the resolution and scanning strategy, currently make mass production impractical. But there is currently no other manufacturing technology for such small 3D structures. The complexity of the 3D structures does not allow microfabrication by injection molding, conventional lithographic or by soft-lithography. However, improving the speed of 2PP processing is an active field in the further industrial development of this technology. The performance of the HLM was demonstrated for antisolvent precipitation to produce lipid nanodispersions containing either castor oil or Softisan® 100 as potential drug carrier material. The central injection of the organic phase, leading to a thin layer, ensures a large contact interface between the liquid phases, while avoiding contact of the organic phase with the channel wall. Subsequently, HLM elements further enlarge the contact interface and enhance diffusive mixing between organic and aqueous phase. Each HLM element consists of a stretching section and a folding section. When passing a series of thirteen HLM elements in one mixing channel, the organic solution is folded into layers, the number of which doubles with each HLM element. The HLM system results in mixing times below 10 ms after three HLM elements. Moreover, the contact of the organic phase with the channel walls is completely prevented within a range of flow conditions. By the introduction of an additional phase via a second nozzle, the organic lipid phase is completely enclosed with a fluidic shield layer before contact with the aqueous phase. A minimum flow rate ratio of this shield phase is necessary for a complete enclosure of the organic phase and successful prevention of fouling on the nozzle surface. This enables the fouling-free use of castor oil and Softisan® 100 in a wider operational window including higher lipid concentrations. In the case of Softisan® 100, use of the shield phase further reduces the particle size compared to the system without the shield phase. Castor oil or Softisan® 100 nanoparticles can be obtained with the HLM system with particle sizes ranging from 42 nm to 166 nm within the optimal flow conditions and without fouling. PDI values between 0.04 to 0.11 indicate narrow up to monodisperse particle size distributions.

Higher flow rate ratios do not widen the particle size distribution. Additionally, the repeated production of samples displays excellent reproducibility. The, in part, extraordinarily small particle sizes obtained suggest that the use of even higher lipid concentrations may also result in particle sizes below 200 nm. In consequence, the lipid content of the nanodispersions would be increased, probably leading to a reduced effort for sample concentration up to lipid contents desired for pharmaceutical applications. The HLM system provides a fouling-free operation range up to

800 µl min⁻¹ across flow rate ratios from 5% to 20% organic lipid phase. This extends the optimal range of our previous micromixer, the coaxial lamination mixer (CLM), which is mainly restricted to total flow rates up to 200 µl min⁻¹ and flow rate ratios up to 15%. The improved total flow rate also corresponds to an increased throughput of the systems, which can be extended further with suggested design optimizations. Moreover, the use of higher concentrations of castor oil and Softisan® 100 in the organic phase than in the CLM is successful with the HLM system. When using equal lipid concentrations and flow rate ratios, smaller particles with narrower size distributions are obtained in the HLM system. While lipid nanoparticles with similar sizes have already been generated in other micromixers, comparably narrow particle size distributions with simultaneous high throughput have, to our knowledge, never been achieved with microfluidic processes so far. The performance and high practical benefits of the HLM system are also demonstrated by the fact that stable long-term production is possible with complete avoidance of fouling.

Future work should include simulations that allow nonlinear interpolation of the properties of the mixture, which could help for designing optimized HLM elements for operation at higher flow rates without the occurrence of secondary vortices. To further increase throughput by an order of magnitude or more one should demonstrate the parallelization of such mixing systems. A microfluidic glass chip with integrated channels for uniform partitioning into multiple parallel mixing channels combined with several flow focusing nozzles as well as HLM elements can be realized by combining femtosecond laser structuring of glass with 2PP printing. Such hybrid systems could further increase the productivity of microfluidic nanoparticle precipitation many times over without the need to increase the number of feed pumps.

Author contributions

Peer Erfle: conceptualization, investigation, methodology, visualization, writing – original draft, writing – review & editing. Juliane Riewe: conceptualization, methodology, writing – review & editing. Songtao Cai: methodology, investigation, writing – review & editing. Heike Bunjes: conceptualization, supervision, writing – review & editing. Andreas Dietzel: conceptualization, supervision, writing – review & editing.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

P. Erfle acknowledges financial support from the 'QUANOMET-Niedersächsische Wissenschaftsallianz' provided by the Ministry of Science and Culture (MWK) of Lower Saxony (Germany) and the technical support for the

cryo-SEM evaluation by Bogdan Semenenko of the Institute for Particle Technology, Technische Universität Braunschweig. J. Riewe acknowledges financial support from Boehringer Ingelheim and the MWK of Lower Saxony within the graduate program "µ Props – Processing of Poorly Soluble Drugs at Small Scale". This work has been carried out within the framework of the SMART BIOTECS alliance between Technische Universität Braunschweig and Leibniz Universität Hannover, an initiative supported by the MWK of Lower Saxony.

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