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Cite this: Lab Chip, 2023, 23, 72

Received 24th July 2022, Accepted 12th November 2022

DOI: 10.1039/d2lc00688j

rsc.li/loc

Integrated silicon microfluidic chip for picoliterscale analyte segmentation and microscale printing for mass spectrometry imaging

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A silicon single-chip microfluidics system that integrates microscale fluidic channels, an analyte segmentation device, and a nozzle for electrohydrodynamic-assisted printing is designed for hyphenation with MALDI mass spectrometry (MS) imaging. A miniaturized T-junction segments analytes into monodisperse picoliter oil-isolated compartments. The printing nozzle deposits generated droplets one-by-one into an array on a conductive substrate without splitting or coalescing. Virtually single-shot MS analysis is enabled due to the ultrasmall droplet volumes and highly localized printing. The signal-to-noise ratio indicates that detection limits at the attomole level are achieved for γ-aminobutyric acid. COMMUNICATION
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Introduction

Automated liquid handling and dispensing through droplet arrays is attracting increasing interest in the fields of single cell analysis, 1 drug discovery² and clinical diagnosis.³ Compared with conventional multiwell plate approaches, droplet arrays minimize sample consumption 4 and enable high throughput screening.^{5,6} A reduced sample volume lowers the detection limit for analysis techniques such as matrix-assisted laser desorption–ionization mass spectrometry (MALDI-MS), which is attributed to the decreased spreading of the dried sample spot.⁷ Owing to the isolated detection on each droplet, the droplet array also enables analyte concentration monitoring⁸ if the droplets are printed following the chronological order in which they were sampled, preserving the temporal resolution.⁹

† WS and SB contributed equally to this paper.

detection stages, temporal resolution could be compromised by Taylor dispersion, 12 where the concentration gradient of the analyte is smeared out due to longitudinal diffusion. Therefore, collecting samples as a segmented flow $9,10$ has been proposed to improve temporal resolution by confining Taylor dispersion within each segmented oil-isolated analyte droplet. If the analyte is segmented near the sampling port and then printed on a MALDI substrate, the resulting array represents a series of chemical events in a chronological order, with the temporal resolution defined by the segmentation frequency. While temporal resolution could be preserved by analyte segmentation, the volume of droplets is typically at the microliter scale $9-11,13$ resulting in increased surface spreading during deposition that lowers MS sensitivity.¹³ Smaller picoliter droplets are therefore beneficial for reduction of

For applications such as neurochemistry, $9-11$ being able to monitor the dynamics of neurotransmitter release is vital for investigating brain function. To achieve a high recovery rate in microdialysis, sampling at ultra-low perfusion flow rates is beneficial.¹⁰ However, during sampling, transportation, and

analyte spreading upon deposition, 14 increased ionization efficiency, better tolerance to salt contamination and oil interference, leading to lower limits of detection $(LODs)$ ¹⁵ However, typical microfluidic devices based on a polydimethylsiloxane (PDMS) platform are utilizing relatively large $100 \times 100 \mu m^2$ channel cross-sections with flow rates in the μ L min⁻¹ range, that, together with the small Young's modulus of PDMS and unwanted swelling, make it difficult to control the ultra-small droplet formation in the picoliter regime. Complex multistage designs¹⁶ or utilization of surfactants 17 are required to produce pL-scale monodisperse droplets. Owing to a much larger Young's modulus and precise control of microfluidic channels down to a few μm radius, silicon microfluidics provides superior control of the ultralow flow rates in the nL min−¹ range critical for maintaining droplet volumes in the picoliter regime¹⁸ and maintaining their monodispersity.¹⁹

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To transfer droplets onto a MALDI substrate while preserving their chronological order, various deposition techniques have been reported including contact printing, 2^{0} as well as contactless approaches including electrospray deposition²¹ and electrohydrodynamic (EHD) assisted printing.²² Contact printing enables deposition¹³ or formation²³ of picoliter to femtoliter droplets by passing a waterfront through an array of prepatterned hydrophilic sites. Among contactless methods, electrospray ionization has been used to introduce picoliter-scale segmented analytes for MS analysis; $10,19$ however, when used for deposition on a MALDI substrate, analytes are aerosolized and ionized in the electrosprayed jet 24 which significantly increases the spread of printed patterns and lowers MS sensitivity.¹⁴ The EHDassisted printing with electric field strength below the formation of the electrospray jet has been demonstrated to produce pL-scale droplets, $25,26$ limited, however, to a continuous non-segmented analyte flow. To achieve contactless printing of oil-segmented picoliter droplets into a MALDI array while simultaneously preserving their chronological order, high temporal resolution, and minimal spreading area, the integration of flow segmentation and printing functionalities in a single device is needed.

In this paper, a single-chip silicon integrated device is developed to segment continuous analytes into pL droplets, and then prints droplets individually into a MALDI array. Attomole-level detection sensitivity is demonstrated for a deposit site localized within the area of a single laser shot.

Fabrication

To segment analytes into a series of individual oil-isolated compartments and then print them one-by-one with the assistance of an electric field, a microfabrication platform¹⁹ based on silicon on insulator (SOI) wafers is used. It consists of three levels of aligned photolithography, several steps of dielectric deposition and etching. Here, four devices (Fig. 1A) are integrated on-chip: connection stubs for fluidic interfacing, sealed surface channels that provide microfluidic flow, a T-junction droplet generator for monodisperse analyte segmentation, and a printing nozzle for individual droplet transfer.

The fabrication is divided into two stages (Fig. 1B): channel definition (Fig. 1B.1–3) and formation of a printing nozzle (Fig. 1B.4). The chip is built on an SOI wafer (UltraSil LLC) with a doped 15 μm device layer, a 0.5 μm buried oxide layer and a 450 μm handle layer. Silicon, as a rigid substrate, eliminates channel deformation and resulting pressure fluctuations to provide stable ultralow flows down to a few nL min−¹ . To define microfluidic channels, a 0.3 μm hard mask of $\sin x$ is first deposited on top of the device layer via plasma enhanced chemical vapor deposition (PECVD) (Fig. 1B.1). Direct laser writing (Heidelberg μPG 101) is used to pattern the $\sin x$ mask, creating a line of 1 μ m diameter holes separated by 2 μm gaps along the center of the designed channel length. Inductively coupled plasma reactive ion etching (ICP-RIE) is then used to open the patterned holes on the $\sin x$ mask, and isotropic XeF₂ etching is performed to form hemispherical microfluidic channels (Fig. 1B.2). The channel radius can be tuned from 7 to 12 μm by controlling the XeF_2 etching time. A 7 μ m radius ensures that etched portions under adjacent 1 μm holes merge uniformly to form a continuous microfluidic channel. A maximum radius of 12 μm is limited by the 15 μm device layer thickness. To seal the 1 μ m holes in the SiN_x mask, a 4 μm thick layer of $\sin x$ is deposited by PECVD (Fig. 1B.3). Leab on a Chip

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To define the perimeter of the printing nozzle (Fig. 1B.4), contact photolithography is performed on the device layer side. ICP-RIE is used to etch through the SiN_x mask, then deep reactive ion etching (DRIE) is used to etch through the

Fig. 1 Fabrication of a silicon microfluidic droplet printing device. A) Schematic of an integrated chip with droplet generation and contactless printing. B) Cross-section schematic of fabrication steps including (1) SiN_x hard mask deposition on SOI and hole opening on SiN_x, (2) isotropic $XeF₂$ etching of silicon, (3) channel sealing with 4 µm SiN_x deposition, and (4) nozzle perimeter and undercut are defined by DRIE. C) SEM side view of a released nozzle. D) Photograph of an integrated chip with a T-junction droplet generator and printing nozzle. E) Photograph of a chip with electric and fluidic interfaces packaged in a 3D-printed plastic holder.

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Si device layer until the buried oxide layer is reached. Finally, aligned contact photolithography and DRIE are performed from the handle layer side (not shown) to undercut the nozzle and to release the chip from the wafer. The scanning electron microscopy (SEM) side view image of the released nozzle cantilever (Fig. 1C) shows a rectangular shape suspended silicon needle with a 15×75 µm cross-section. To efficiently electrify droplets at the nozzle outlet to ease printing, the device layer is heavily doped yielding 0.005 Ω cm resistance providing electrical contact without charging of the analyte flow. 27

The fabricated chip that integrates an upstream T-junction droplet generator, downstream printing nozzle and sealed microfluidic channels (Fig. 1D) is released from the wafer and is packaged to provide fluidic and electrical interfaces (Fig. 1E). To interface with external microfluidic pumps, the inlet/outlet of each microfluidic channel is shaped into a 170 \times 450 μm silicon 1400 μm long stub to fit with silica capillaries (Polymicro TSP530700, 650 μm ID). UV curable resin (Norland Products, NOA 68T) is applied at the stub–capillary junction to ensure airtight sealing. For electrical connection, copper tape is bound to the doped device layer via conductive adhesive (MG Chemicals 8331D). The final device (Fig. 1E) with fluidic and electrical interfaces is packaged into a 3D-printed plastic holder. Communication

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Analyte segmentation

Since the inner surface of the as-fabricated microfluidic channels is covered by a thin silicon dioxide layer, the channels' surface is hydrophilic. Therefore, the oil phase is dispersed, while the aqueous phase is continuous. To segment the aqueous analyte (10 μM fluorescein, Sigma-Aldrich in DI water) into monodisperse picoliter-scale plugs separated by oil droplets (Fluorinert FC40, Sigma-Aldrich), a cross-flow shear in a T-junction structure with a 12 μm channel radius is utilized (Fig. 2). The aqueous phase flow was maintained at a constant flow rate of 15 nL min−¹ by a syringe pump (Harvard Apparatus Pump11 Elite). The oil phase was introduced at a pressure varied between 100 and 500 mbar by a pressure pump (FlowEZ LU-FEZ-7000) to tune the flow rate ratio.

As fabricated, the microfluidic channels are hydrophilic, hence oil is the segmented phase, while the aqueous phase is continuous. Previous MS measurements²⁸ of the potential carry-over of analytes between consecutive aqueous plugs have shown that carry-over of analytes in the oil-segmented aqueous flow between aqueous plugs of similar volume and frequency is suppressed by at least 3 orders of magnitude.

Since the silicon nitride sealing on top of a surface microfluidic channel is transparent for visible light, the droplet flow can be analysed with standard fluorescence microscopy (Olympus IX73 fitted with a Canon Rebel T7i video camera). Fig. 2A displays a single frame in a video recorded during droplet formation at an oil pressure of 300 mbar $(1:2)$ oil to water flow rate ratio), which shows a

Fig. 2 Analyte segmentation into monodisperse pL droplets. A) Fluorescence microscopy image of the oil phase (black) protruding into the water phase (green) at the T-junction with a channel radius of 12 mm. B) Analyte segmentation into a series of oil-isolated droplets (green) with a constant volume travelling down the channel. C) Statistics of the droplet volume for a 12 μm radius T-junction produced at an aqueous flow rate 15 nL min−¹ and oil pressure of 300 mbar (1 : 2 oil to water flow rate ratio). The solid line is the normal distribution. D) Tuning droplet volume by adjusting the flow rate ratio at the T-junction with a channel radius of 7 μ m. Boxes cover the 25-75% data range and whiskers represent the standard deviation. The red line is a linear fit.

characteristic "necking" of the oil protruding into the aqueous phase flow and blocking the entire channel diameter. The oil droplet generates a pressure build-up in the channel that eventually overcomes the interfacial tension between phases followed by the droplet's release.¹⁷ When operating in such a "squeezing" regime, 29 the process results in a repeatable generation of almost identical aqueous plugs (Fig. 2B). Analysis of recorded videos reveals the mean aqueous phase volume of 43 pL with a monodispersity of 9% relative standard deviation (SD) (Fig. 2C). Both, a plug frequency of 2.1 Hz with an SD of 0.2 and volume, are stable for tens of minutes of continuous operation. By adjusting the oil to water flow rate ratio from 0.5 to 3 by tuning the oil phase pressure, the aqueous plug volume can be continuously tuned down to 25 pL.

Even smaller aqueous plug volumes are attainable (Fig. 2D) by decreasing the microfluidic channel radius down to just 7 μ m by adjusting the XeF₂ etching time. In this chip, with the oil phase pressure tuned from 300 to 500 mbar and the aqueous phase flow rate kept constant at 10 nL min⁻¹, the flow rate ratio is tuned from 0.2 to 1.1. The resulting mean aqueous plug volume is tuned from 4 to 9 pL (Fig. 2D). Within the whole tuning range, the monodispersity is preserved below 5% RSD and the dependence of the aqueous plug volume on the flow rate ratio shows high linearity $(R^2 =$ 0.962), confirming operation in the squeezing regime. 30

Nozzle tip design for EHD-assisted printing

Once the analyte is segmented into a train of individual pLscale aqueous plugs at a few Hz frequency, they need to be

deposited one-by-one onto a MALDI substrate without splitting or coalescing to preserve their chronological order and ensure temporal resolution of the MS analysis. The goal for the nozzle tip design, therefore, is to guarantee that the upstream pulsatile dripping at the tip^{31} is in resonance with the volume and frequency of droplets generated downstream at the T-junction. That is challenging since the pulsatile flow segmentation at the nozzle tip is defined by a balance of many forces (Fig. 3A). A viscous drag of the flow pulling on the protruding water meniscus at the nozzle tip,³² σ_K = 4/ $3\rho QU$ (where ρ is the liquid density, Q is the flow rate, and U is the average velocity of the liquid), may not be enough for ultra-slow nL min−¹ flow rates to overcome a counteracting interfacial tension, $\sigma_y = 2\pi L_y$ (where γ is the surface energy and L is the wetting perimeter), that keeps protruding water from being detached from the silicon interface (Fig. 3A). Gravitational forces, important for microliter and nanoliter volume droplets, 13 can be neglected for picoliter volumes since the Bond number (the ratio of gravity to interfacial tension) is only 10−⁵ . Therefore, under gravity alone, picoliter segmented analytes will coalesce into much larger droplets at a slower frequency, hence degrading the temporal resolution and mixing their chronological sequence for the MALDI-MS analysis. Lab on a Chip

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EHD-assisted printing²² enables fine tuning of the droplet detachment since it relies on additional electrostatic force,³³ $\sigma_E = 2\pi r^2 \epsilon_0 E_c^2$ (where r is the droplet radius, ϵ_0 is the air permittivity, and E_c is the electric field strength), that changes the force balance at the nozzle tip (Fig. 3A). Various EHD spraying regimes 31 are typically observed with the increase of electric field strength starting with dripping, followed by micro-dripping, spindle, and cone-jet. 26 Among these regimes, droplet formation in the pulsatile micro-

Fig. 3 A) Schematic of a force balance during droplet detachment from the nozzle tip. B) SEM image of the sharp nozzle design. C) SEM image of the flat nozzle design. D) Series of background-subtracted fluorescence microscopy images from a video recorded during droplet formation for sharp nozzle design with a 7 μ m radius channel at 0.64 kV and 20 nL min⁻¹. The rightmost frame is the last captured before droplet detachment. E) Series of background-subtracted fluorescence microscopy images for flat nozzle design with a 7 μ m radius channel at 0.8 kV and 20 nL min⁻¹. The rightmost frame is the last captured before droplet detachment.

dripping regime is preferred when a periodic series of monodisperse (no siblings) droplets³⁴ with a diameter about the nozzle size D are formed at an axially stable meniscus.²⁶ However, the micro-dripping regime occurs only within a narrow range of electric field strengths and flow rates.^{26,34}

To explore this parametric space to produce pL-scale aqueous droplets at a few Hz frequency, two complementary tip designs are fabricated (Fig. 3B and C) with the same width of the nozzle base of 75 μm that is either tapered down to a sharp tip (Fig. 3B) or terminated in a flat tip (Fig. 3C). This additional capability, provided by silicon microfabrication to lithographically define the micron-scale geometry of the nozzle, enables the influence of the balance of forces (Fig. 3A) by modulation of the wetting perimeter L , and concentrating the electric field at the sharpened nozzle $tip¹⁹$ as opposed to a more uniform field in the flat tip design.

To map the parametric space of a micro-dripping regime in both designs, a series of experiments were performed to electrospray a continuous aqueous phase (10 μM fluorescein in DI water) at different flow rates (5 nL min−¹ and 20 nL min⁻¹) and applied voltages. A voltage is applied between the chip and the ITO coated glass slide (Delta Tech CB-40IN-S111) located at a 2 mm distance from the nozzle tip. To quantify droplet formation, a fluorescence microscopy video was recorded, and a single background image was subtracted from all individual frames (Fig. 3D and E) to increase the droplet visibility. The resulting high-contrast images captured during EHD-assisted deposition enable the measurements of the droplet volume and frequency at different voltages and flow rates for both sharp-tip (Fig. 3D) and flat-tip (Fig. 3E) designs.

For the sharp-tip design, at voltages below 0.62 kV, large and irregular nL-scale droplets are accumulated at the tip driven mostly by gravity (Fig. 4B, left). At voltages higher than 0.72 kV (Fig. 4B, right), the EHD spraying occurs in the form of unstable spindles and microjets.26,31 The range of voltages between 0.64 kV and 0.7 kV at a constant 20 nL min⁻¹ flow rate results in pulsatile generation of almost identical droplets (Fig. 4B, center) indicative of a micro-dripping regime. 26,31 At 0.64 kV, the sharp-tip design produced droplets with a mean volume of 786 \pm 15 pL at 1.1 \pm 0.2 Hz (Fig. 3D). A gradual increase of the voltage in 0.2 kV increments results in the decrease of the droplet volume accompanied by a reduction of pulsation frequency that is summarized in a log–log plot in Fig. 4A (circles). At 20 nL min⁻¹ flow rate (closed circles), the log-log dependence shows high linearity (R^2 = 0.998, black line) with a slope of -0.895 . At the slower 5 nL min⁻¹ flow rate (open circles), the droplet frequency decreased 2× with a similar slope.

In contrast, for the flat-tip design (Fig. 4C), the voltage range for a stable micro-dripping regime is shifted to higher voltages between 0.7 kV and 0.82 kV. At 0.80 kV and 20 nL min−¹ , much larger 2100 ± 100 pL droplets are generated at a $10\times$ slower frequency of 0.12 \pm 0.01 Hz (Fig. 3E). The corresponding log–log plot (Fig. 4A, triangles) exhibits highly

Fig. 4 A) Scaling of droplet frequency vs. droplet volume for various flow rates and applied voltages for two alternative nozzle tip designs. Circles correspond to the sharp tip design, while triangles to the flat tip design. Open and closed symbols correspond to 5 nL min−¹ and 20 nL min⁻¹ flow rates, correspondingly. Solid lines are linear fits to the data. The dashed line indicates extrapolation towards the target parameters to be met for electrospraying the analytes that are presegmented at the T-junction into 10 pL droplets at 1 Hz. B) Background-subtracted fluorescence microscopy images for sharp nozzle design with a 7 μm radius channel with 20 nL min⁻¹ flow at 0.62 kV (dripping), 0.66 kV (micro-dripping), and 0.74 kV (spindle). C) Background-subtracted fluorescence microscopy images for flat nozzle design with a 7 μm radius channel with 20 nL min⁻¹ flow at 0.72 kV (dripping), 0.8 kV (micro-dripping), and 0.82 kV (spindle).

linear dependence (slope -0.384, R^2 = 0.928, black line), almost identical to both 5 nL min⁻¹ (open triangles) and 20 nL min−¹ (closed triangles) flow rates.

Scaling dependencies of EHD-assisted spraying are defined by the quasi-static force balance $\sigma_E + \sigma_K = \sigma_v$ at the time of the droplet detachment 35 (Fig. 3A). If the dimensionless electric Bond number $B_E = \varepsilon_0 E_c^2 D / \gamma$ is introduced, the force balance for the micro-dripping regime, when droplet diameter $2r$ is of the order of outlet nozzle size D and of the wetting perimeter L , can be rewritten as:

$$
f^2 V = \frac{3}{2\rho} (1 - B_E) \gamma \tag{1}
$$

Considering the narrow range of voltages for the microdripping regime, the Bond number B_E can be considered constant. Then, indeed, the slope for the flat tip design (Fig. 4A, triangles) is close to the expected −0.5 confirming f^2V invariance expected from eqn (1), invariant with respect to the flow rate as observed.

The results for the sharp tip design (Fig. 4A, circles) differ significantly from eqn (1), that is most likely defined by two separate contributions to the force balance. First, the flow velocity U is measured to be $2\times$ faster than that in the flat tip design that is likely to be induced by larger electro-osmotic drag due to higher field concentration at the sharp tip discontinuity.¹⁹ This results in 4× larger kinetic force σ_K indicating faster droplet detachment and hence higher frequency. Second, the interfacial tension σ_{ν} is 3× smaller than that in the flat tip design due to smaller wetting perimeter L (Fig. 3D). Considering the electro-osmotic acceleration, the force balance requires³⁵ $\rho r^4 f^2 \sim \varepsilon_0 E_c^2 r^2$, and eqn (1) can be rewritten as:

$$
fV = B_E^{\ 0.5} \tag{2}
$$

Indeed, the resulting slope of −1 in the coordinates of a log– log plot in Fig. 4A (circles) is close to the experimentally determined value.

Our estimate of the balancing forces at the tip end (Fig. 3A) indicates that the scaling dependence between the frequency and volume outlined in eqn (1) and (2) (Fig. 4A) is, to a first approximation, invariant to the concentration of analytes and to the concentration of various salts which are abundant in biological samples. The electrostatic force σ_E acting on a hanging droplet at the tip of an electrified nozzle does not depend³⁶ on liquid conductivity K if the electric relaxation time $t_e = \varepsilon_0 \varepsilon / K$, required for the equipotential to reach the droplet surface, is much shorter than the droplet detachment time. For example, the t_e is about 10⁻¹⁰ s for the cerebrospinal fluid ($\varepsilon = 10^9$ and $K = 2$ S m⁻¹ at low frequency³⁷) that is at least 9 orders of magnitude faster than the printing period used here. For analytes with a concentration of a few μM dissolved in DI water, the conductivity K is still large enough³⁸ (K = 2 × 10⁻⁴ S m⁻¹) to yield a t_e of 10⁻⁶ s, still 5 orders smaller than the printing period. The t_e is becoming long enough to potentially affect droplet detachment only for the DI water of grade 3 ($K = 10^{-7}$) $S \; m^{-1}$).

Resonant one-to-one droplet deposition

Neither of the nozzle tip designs could operate at resonance with the droplet train generated at the T-junction (a few pL at frequencies around 1 Hz). Therefore, under conditions explored in Fig. 4, the aqueous plugs generated at the T-junction during flow segmentation will either split into smaller droplets or coalesce into larger ones thus disrupting their chronological sequence and degrading temporal resolution. Intuition built upon the results in Fig. 4 implies that if the balance of forces can be shifted towards smaller volumes and higher frequencies, the resonance condition can potentially be within reach for the flat-tip design as indicated by extrapolation of experimental results to smaller volumes (dashed line in Fig. 4A). It has been shown³⁹ that the transition from dripping to spindle and jetting modes occurs at the critical Weber number, We = $\rho v^2 2r/\gamma$, that depends on

the flow velocity v. Widening the channel radius from 7 μm (design in Fig. 2D and 4) to 12 μm (design in Fig. 2A–C) will decrease the flow velocity (hence the critical Weber number) as much as 5-fold.³⁹ Therefore, at the same flow rate, by increasing the channel radius, the boundary from microdripping to jetting regimes will be pushed to higher voltages, so that smaller droplets could be printed.

To verify this intuition, the Si integrated chip with 12 μm radius channels equipped with a T-junction (design in Fig. 2A–C) and a flat-tip nozzle (design in Fig. 3C) was fabricated. The chip was mounted in front of an ITO coated glass slide (Delta Tech CB-40IN-S111) at a 2 mm distance. For chemical sensitivity measurements, 200 μM γ-aminobutyric acid (GABA) is added to the aqueous phase (10 μM fluorescein, Sigma-Aldrich in DI water). Under the same conditions as in Fig. 2C (aqueous flow rate = 15 nL min^{-1} and oil pressure $= 300$ mbar with a 1:2 oil to water flow rate ratio) and with application of 1.2 kV between the chip and the slide, the flow is segmented into a series of monodisperse oil-separated aqueous plugs. Lab on a Chip

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Analysis of fluorescence microscopy images taken at the outlet of a printing nozzle right before the aqueous droplet detachment (Fig. 5A) shows stable pulsatile (2 Hz frequency) generation of monodisperse droplets with a mean volume of 42 pL and 10% SD (Fig. 5B). The distribution in Fig. 5B was determined to not differ significantly (*t*-test, $p = 0.054$) from that in Fig. 2C. This confirms that the device operates in resonance with aqueous droplets leaving the nozzle individually one-by-one without splitting or coalescing.

Fig. 5 Characterization of individual droplet printing. A) Single video frame taken at detachment of a 40 pL droplet from the flat nozzle. B) Statistical analysis of the droplet volume measured at the nozzle tip. C) Fluorescent residue of an array of printed droplets after drying. D) Fluorescent residue of a single droplet residue. E) Statistical analysis of the area of the fluorescent residue.

To print an array of the droplets, the ITO glass slide was fixed on top of a motorized stage translated at a constant 500 μ m s⁻¹ speed. Once deposited, droplets evaporate leaving a periodic array of deposits on the slide surface. Bright-field and fluorescence microscopy techniques using an Axio Imager M2 (Carl Zeiss, Jena, Germany) were used to visualize residue locations with respect to fiducial marks (Fig. 5C). Images were stitched and exported using Zen 2 (Carl Zeiss, Blue edition) software.

The observed fluorescently labelled residues are separated by a constant 250 μm gap (Fig. 5C) which is expected for a stage traveling at 500 μ m s⁻¹ at 2 Hz generation and transfer frequency. Further analysis of the fluorescent residues (Fig. 5D) shows an area of 1250 μ m² (Fig. 5D) consistent with the expected 40 μm diameter of a 40 pL droplet. The distribution of fluorescent residue areas for 38 printed droplets (Fig. 5E) has a mean value of $1410 \mu m^2$ with 28% SD that confirms that each fluorescence residue corresponds to an individual aqueous droplet. No sign of oil was observed throughout the substrate that is likely due to fast evaporation of oil droplets with a volume as small as just 20 pL $(2:1)$ water to oil ratio in Fig. 2B).

Characterization of droplets during all stages of droplet generation, transfer, detachment, and landing confirms that individual, intact droplets are transferred.

Mass spectrometry of deposited droplets

Matrix-assisted laser desorption/ionization-time of flight (MALDI-TOF) mass spectrometry imaging (MSI) was used to map the chemical contents of the deposited droplet arrays. MSI can provide untargeted, multiplexed detection in biochemical systems,⁴⁰ making it an ideal detection method to couple to the Si platform for future biomedical applications. To perform MALDI-MSI, after fluorescence microscopy, the ITO glass slide was covered with a thin layer of a common MALDI matrix, α-cyano-4-hydroxycinnamic acid (CHCA). The CHCA solution was prepared at 10 mg mL^{-1} in 70% acetonitrile. An HTX-M5 Sprayer (HTX Technologies, Chapel Hill, NC) was used to apply the matrix to the glass surface at 100 μ L min⁻¹ and 60 °C, with nebulizing gas pressure at 10 psi. The fluorescence image acquired in Fig. 5C was used within FlexImaging (Bruker Corp., Billerica, MA) to identify sites for targeted MALDI-MSI of the printed droplets. Mass spectra were acquired on an UltrafleXtreme MALDI TOF/TOF mass spectrometer with a frequency tripled Nd:YAG solid-state laser (Bruker Corp.) using a spatial resolution of $50 \times 50 \mu m^2$, the "Ultra" laser setting, and 100 laser shots.

When the MALDI laser spot is focused on the target area (Fig. 6A), a clear ion signal at $m/z = 104.07$ is detected (Fig. 6B, red) corresponding to $[M + H]^{+}$ of GABA. The GABA signal taken at an adjacent blank spot representing the background (Fig. 6B, black) is 2–3 orders of magnitude smaller. The appearance of this signature peak only at the

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Fig. 6 MS-MALDI imaging analysis of a printed droplet array. A) Schematic of MALDI-MS. B) Spectra of the printed residue site (red) and blank background (black). Spectra are shifted vertically for clarity. C) Intensity profile of GABA ion ($m/z = 104.07$) at the residue site. D) Fluorescence microscopy image of the same residue site. The white crosses in C) and D) mark the same location on the slide.

residue site confirms that the signal at $m/z = 104.07$ cannot be attributed to matrix or background interference. The nearby peak at around $m/z = 102.5$ is attributable to the matrix or contamination but does not interfere or overlap with GABA signal detection. These mass spectra confirm that the analyte signal is detected at the printing site, and the signal peak is isolated well enough to avoid interference.

The unwanted spatial delocalization of the analyte can occur due to analyte diffusion beyond the initial printed droplet residue during solution-based matrix application that can significantly degrade chemical sensitivity. It has been shown¹⁴ that the analyte diffusion with the CHCA matrix application procedure is negligible for 60 pL deposited droplets. To evaluate potential analyte spreading, a spatial resolution study of the MS signal was performed to compare MSI signal mapping (Fig. 6C) with the fluorescent droplet areas (Fig. 6D). The MSI heat map shows the spatial distribution of the GABA signal at $m/z = 104.07$ (Fig. 6C). Total ion count (TIC) normalization is used to reduce the effect of differences in ionization efficiency and/or matrix– analyte co-crystallization across pixels representing adjacent spots on the ITO glass slide. Since the laser step size (pixel width) is set at 50 μm while the laser beam diameter is 100 μm, the GABA signal from a 40 μm diameter fluorescent droplet residue is expected to be confined to a 100 μm radius. The heat map shows that the GABA signal is mostly concentrated in a single 50 μm pixel with a few, lower intensity adjacent pixels within a 100 μm radius.

To identify the localization of the MS signal at $m/z =$ 104.07 relative to the position of the deposited residue, the fluorescence image (Fig. 6D) of the substrate was acquired from the same area as the MS image (Fig. 6C). The fluorescent droplet residue at the same coordinates (relative to the white crosses in Fig. 6C and D that mark the same location) is strongly co-localized with the pixel of the highest intensity GABA signal. Even with the current step-wise scanning approach for MSI, the number of pixels scanned per printed droplet is relatively small due to the ultrasmall pL droplet volume indicating virtually single-pixel sample ionization.

MALDI-MS imaging of printed droplet array

To demonstrate the capability of the system for highthroughput droplet array printing and high sensitivity MALDI-MSI detection, 41 droplets were printed in a row and then analysed by MALDI-MSI. A 1 mm long area was scanned using the same MALDI-MSI parameters above to create the TIC normalized GABA signal heat map for all 41 droplets (Fig. 7A). All printed droplets are detected as well-isolated 100 μm radius individual GABA deposits. Since the frequency for droplet transfer is 2 Hz, it takes only 20 seconds to print this line automatically, enabling further expansion into 2D arrays by adding another axis of movement to the mechanized stage. These results confirm the high throughput printing and detection capabilities of the system.

To assess the chemical sensitivity of the approach, the signal to noise ratio (SNR) and limits of detection (LODs) of MALDI-MSI for GABA detection are estimated. The LOD for MSI is expected to be improved 14 by concentrating the analyte to a small area. Owing to analyte segmentation into pL droplets, the MS signal is strongly localized with a single

Fig. 7 MSI analysis for the printed droplet array. A) Heat maps of the MSI GABA signal intensity at $m/z = 104.07$ for all 41 droplet deposits. A single line printed array is folded into three separate sections to ease figure layout. B) Statistics of MSI signal intensity at the printed residue and blank sites.

printed site not extending beyond 100 μm diameter thus promising high sensitivity (Fig. 6C). To provide a noise level reference, a blank background signal is collected during MSI scanning in the immediate vicinity of the GABA residue. The highest intensity, TIC normalized intensity at $m/z = 104.07$ for blank pixels, is 1.98, which is attributable to chemical noise and possible isobaric ions formed from the MALDI matrix and/or the ITO-coated glass surface. This highest intensity signal from blank pixels is used as a threshold to distinguish between the true GABA signal and background signal. For all pixels at each printing site, pixels with TIC normalized intensity $(m/z = 104.07)$ higher than the threshold of 1.98 are considered pixels with the GABA signal. All pixels with signal intensity above the threshold for a given printed site are summed to calculate the total GABA intensity per printed droplet area. The mean summed intensity at printed sites is found to be 206.6 \pm 44.6 (mean \pm standard deviation), while blank pixels only have 1.27 ± 0.32 (Fig. 7B). Given that each droplet contains 40 pL of 200 μM solution of GABA, the measured summed signal from each printed site corresponds to 8 femtomoles of GABA molecules. To calculate the LOD, a single point calculation was done since only one GABA concentration was printed. The mean blank intensity + 3× blank standard deviation was calculated as 2.2 to represent the signal intensity for GABA at the LOD. To determine the concentration of GABA corresponding to the LOD, the LOD signal is multiplied by the ratio of 8 fmol per mean summed droplet intensity to yield an LOD of 86 amol. Therefore, an attomole level LOD for the MSI detection of GABA was achieved using the Si integrated chip with a flat nozzle for automated, high-throughput droplet array printing. Lab on a Chip Common Access Article Common Common Common Common Common Common Call in the common Call in the

Conclusions

A silicon microfluidic chip that integrates multiple devices including microfluidic channels, analyte segmentation, a nozzle for EHD-assisted droplet printing, and packaging stubs is designed and fabricated. The T-junction analyte segmentation device operating at an ultralow nL min⁻¹ flow rate produced monodisperse droplets with tuneable volumes at the picoliter scale. The EHD-assisted printing nozzle design operates in resonance with the pre-segmented analytes to print droplets individually one-by-one without splitting or coalescing. Printed droplets on ITO glass are highly localized and well-isolated forming a periodic array that is ideal for MALDI-MSI analysis. Virtually single-pixel acquisition of mass spectra is enabled due to the ultrasmall droplet volumes and highly localized printed droplet deposits. A fewer pixels per droplet concentrate the MS signal and increase the sensitivity leading to an 86 attomole LOD for GABA. This integrated Si chip has strong potential for single cell analysis, by virtue of the reduction of the droplet volume down to the single cell level. Application of this approach to microdialysis-based neurochemistry,⁴⁰ where the sample volume is usually limited and temporal information needs to be preserved, is promising.

Author contributions

The manuscript was written through contributions of all the authors. All the authors have given approval to the final version of the manuscript.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

The research reported in this publication was supported in part by the NINDS of the NIH grants UF1NS107677 and RF1NS126061 and the NSF grant 1735252. The content is solely the responsibility of the authors and does not necessarily represent the official views of the National Institutes of Health.

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