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1 **Exploring Chemistry in Micro-Compartments using Guided Droplet**
2 **Collisions in a Branched Quadrupole Trap Coupled to a Single Droplet, Paper**
3 **Spray Mass Spectrometer**

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13
14 **Abstract:**

15 Recent studies suggest that reactions in aqueous micro-compartments can occur at significantly
16 different rates than those in the bulk. Most studies have used electrospray to generate a
17 polydisperse source of highly charged microdroplets, leading to multiple confounding factors
18 potentially influencing reaction rates (e.g., evaporation, charge, size). Thus, the underlying
19 mechanism for the observed enhancement remains unclear. We present a new type of
20 electrodynamic balance—the branched quadrupole trap (BQT)—which can be used to study
21 reactions in microdroplets in a controlled environment. The BQT allows for condensed phase
22 chemical reactions to be initiated by colliding droplets with different reactants and levitating the
23 merged droplet indefinitely. The performance of the BQT is characterized in several ways. Sub-
24 millisecond mixing times as fast as ~400 μs are measured for low velocity (~0.1 m/s) collisions of
25 droplets with <40 μm diameters. The reaction of ortho-phthalaldehyde (OPA) with alanine in the
26 presence of dithiothreitol is measured using both fluorescence spectroscopy and single droplet
27 paper spray (PS) mass spectrometry. The bimolecular rate constant for reaction of alanine with
28 OPA is found to be 84 ± 10 and $67 \pm 6 \text{ M}^{-1} \text{ s}^{-1}$ in a 30 μm radius droplet and bulk solution,
29 respectively, which demonstrates that bimolecular reaction rate coefficients can be quantified
30 using merged microdroplets and that merged droplets can be used to study rate enhancements due
31 to compartmentalization. Products of the reaction of OPA with alanine are detected in single
32 droplets using paper spray mass spectrometry. We demonstrate single droplets with <100 pg
33 analyte can easily be studied using single droplet mass spectrometry.

34 I. Introduction

35 Chemical reactions in micron-sized compartments are ubiquitous in nature, occurring in
36 cells, mineral pores, and atmospheric aerosols. Several recent studies suggest that reactions in
37 confined spaces occur at enhanced rates that can be several orders of magnitude faster than those
38 in the bulk.¹⁻⁷ Although the underlying mechanism remains unclear, three main factors are thought
39 to contribute—increased concentrations due to solvent evaporation, surface acidity (i.e. charge),
40 and interfacial adsorption.⁸ Many studies reporting enhanced rates of reaction have used an
41 electrospray source to generate an aerosol plume.¹⁻⁵ Electrospray sources produce highly charged
42 and rapidly evaporating droplets, which could lead to increasing reactant concentrations,
43 significant pH variability compared to the bulk (due to solvent oxidation by the charges in the
44 droplets), and complex ion-ion and ion-solvent interactions. Further, the droplet plume exhibits a
45 high surface area relative to the bulk volume, thus enhancing the role of interfacial adsorption in
46 perturbing chemical kinetics.⁸ An increase in the surface to volume ratio of the droplets has been
47 found to increase the reaction rate, suggesting the droplet surface could play a major role in the
48 observed rate acceleration.⁷ Most measurements studying rate enhancements in microdroplets use
49 a polydisperse droplet source, which limits the control of both size distribution and time evolution
50 of the reactant concentration.¹⁻⁶ Microfluidic devices capable of generating a monodisperse droplet
51 distribution have also been used to study rate enhancements in microdroplet emulsions.⁷ However,
52 it remains unclear if the rate enhancing properties attributed to the oil-water interface of
53 microfluidic devices are general to the air-water interface.

54 Another means of ensuring a monodisperse size is by limiting a study to a single droplet.
55 The contactless confinement of single, micron-sized droplets has been established as a powerful
56 method for probing the physical and chemical properties of liquids and heterogeneous interfaces.⁹

57 For example, aerosol optical tweezers (AOT) have been used to measure the surface tension,
58 viscosity and hygroscopicity of a wide variety of aqueous samples, while electrodynamic balance
59 (EDB) methods have allowed rapid mass transport and transformation processes to be
60 interrogated.⁹ AOT provide powerful, real-time characterization of droplets via the morphology
61 dependent resonances that appear in the Raman spectrum of the droplet induced by the trapping
62 laser. Also, a holographic optical trap (HOT) can manipulate arrays of droplets, and bring selected
63 droplets into contact when desired. All droplets confined in a HOT can be fully characterized using
64 cavity enhanced Raman spectroscopy to determine their size to nanometer accuracy before they
65 are mixed together.¹⁰ EDB's are extremely versatile techniques and offer advantages over AOT
66 methods due to the facile introduction of single droplets into the confinement region and the ability
67 to trap particles that strongly absorb laser light. EDB's can trap a larger range of droplet sizes than
68 AOT (from 100s of nanometers¹¹ to 100 μm diameter for EDB's⁹ compared with $<10 \mu\text{m}$ for
69 AOT¹⁰). A wide range of electrode configurations have been developed (e.g. double ring,
70 cylindrical, quadrupole) to facilitate a broad range of measurements, including the study of Mie
71 scattering,¹² hygroscopicity and phase transitions,^{13,14} heterogeneous chemistry,^{15,16} ice
72 nucleation,¹⁷⁻¹⁹ and mass transport from droplets.^{20,21} In these measurements, compositional
73 changes in confined particles are typically the result of interfacial processes, such as mass transport
74 (evaporation and/or condensation of semi-volatile and volatile material) or chemical changes due
75 to reactive gas uptake. This makes bulk initiated chemistry difficult to study in conventional EDB
76 configurations because it is hard to initiate a reaction cleanly at a well-defined time in a single
77 droplet with no external reactant source.

78 The optical probes, such as Raman spectroscopy, that are typically used to characterize
79 droplets in an EDB or AOT are limited to functional group information and cannot provide

80 information about the exact chemical composition of the droplet as it undergoes reactive
81 transformations.⁹ As evident from previous rate enhancement measurements,¹⁻⁶ mass spectrometry
82 of droplets can provide quantitative chemical information, but studies characterizing single
83 droplets with mass spectrometry are limited. Previously, free-falling droplets have been ionized
84 directly by impaction on a highly charged needle, forming a spray that is directed toward a mass
85 spectrometer.²² Several studies have performed mass spectrometry on single, levitated droplets in
86 an acoustic trap (which uses ultrasonic waves to confine particles) coupled to a laser desorption^{23,24}
87 or a direct analysis in real time (DART) ionization source.²⁵ However, because of the strong
88 electric fields in EDB's and the small size of the trapped droplet, non-destructive mass
89 spectrometry is difficult (i.e. particles need to be removed from the trap and ionized completely).
90 Previous work has shown droplets can be ejected from a double ring electrode, deposited onto a
91 matrix and ionized using matrix assisted laser desorption ionization.²⁶ However, the delay between
92 deposition and detection is not ideal for real time reaction monitoring, and direct ionization of
93 droplets as they exit the EDB is preferred. Very recently, work has quantified evaporation from
94 microdroplets by ejecting single droplets from a double ring electrode, vaporizing them on a heated
95 platform and ionizing the components using a corona discharge.²⁷

96 In order to further explore chemical transformations in micron-sized compartments, we
97 present here a new EDB electrode arrangement—the branched quadrupole trap (BQT)—that is
98 capable of merging confined droplets. Merged droplets have previously been trapped using a
99 tandem electrodynamic trap and observed spectroscopically.²⁸ The BQT allows for rapid changes
100 in composition via droplet coalescence (enabling the study of bulk initiated processes) and allows
101 for the ejection of single droplets into an ionization source for mass spectral analysis (enabling
102 chemical characterization of droplets). By coupling droplet confinement techniques with high

103 resolution mass spectrometry, it is possible to measure condensed phase chemical kinetics in
104 micro-compartments under well-defined conditions. Due to the droplet size range over which EDB
105 methods operate (1 – 10's μm) and their ability to control droplet composition by changing
106 environmental conditions, they represent a unique way to study chemical reactions in a potentially
107 interesting droplet size regime. In this work, we characterize the mixing times of merged droplets
108 and demonstrate the application of the technique to probing chemical processes in droplets using
109 both fluorescence spectroscopy and single droplet paper spray (PS) mass spectrometry.

110 **II. Experimental Section**

111 **a. Branched Quadrupole Trap Design**

112 A linear quadrupole EDB allows arrays of droplets to be confined along the axis of four
113 rods, as has been discussed previously.^{11,29} In order to facilitate merging of different droplet
114 populations, the branched quadrupole trap (BQT) design has been developed (shown in Figure 1).
115 Aqueous droplets are generated using a piezoelectric dispenser with a 50 μm orifice (Microfab,
116 Inc.) and are introduced into the top of the BQT along the linear axis. During actuation of the
117 dispenser, high voltage (typically <1 kV) is applied to an induction electrode to induce a net charge
118 on the droplet allowing it to become confined within the electric field of the BQT. Droplets
119 generally have a net charge on the order of 10^{-13} C ($\sim 10^6$ elementary charges),³⁰ which leads to
120 surface charge densities that are ~ 100 times smaller than those found in electrospray droplets.³¹
121 The BQT consists of four stainless steel trapping electrodes arranged in a quadrupole
122 configuration. An alternating voltage (V_{ac}) with an amplitude of 300-500 V and frequency of 200-
123 400 Hz is applied to the trapping electrodes to confine the charged droplets axially along the rods.

124 A set of branching electrodes that extend off the linear trap at a 60° angle allow two
125 separate droplet dispensers with to be used concurrently or simultaneously. By using two different
126 solutions in each of the dispensers, chemistry can be initiated when droplets from each of the
127 dispensers collide. Droplets dispensed from both the vertical dispenser and branch dispenser travel
128 ~ 12 cm before they enter the common, lower portion of the trap. Droplets are held in the lower
129 trap using a set of balancing electrodes that consist of stainless steel blades extending
130 symmetrically between the rods toward the center of the quadrupole trap. By applying a static
131 voltage (up to ± 500 V) to these electrodes, the gravitational force acting on the droplet can be
132 overcome, leading to contactless levitation of the droplet.

133 Once confined within the trap, droplets are illuminated with a 532-nm laser (Changchun
134 New Industries Optoelectronic Tech.) introduced axially. Scattered light from the laser is collected
135 (using an optical scheme that has been previously reported)³² and imaged in the far field with a
136 CMOS camera (Thorlabs, Inc.) using a 532-nm line pass filter. The far field image serves two
137 purposes: droplet positioning and sizing. A feedback loop controlling the voltage applied to the
138 balancing electrodes is used to keep the droplet centered in the far field image and stationary in
139 the trap. For example, as a droplet evaporates and loses mass, the magnitude of the voltage applied
140 to hold the droplet decreases to keep the droplet fixed in space. Additionally, the far field image
141 contains interference fringes from Mie scattering with distinct maxima and minima. The angular
142 spacing of these fringes, the wavelength of scattered light (532 nm) and the refractive index of the
143 droplet (1.36 for the ~ 3 M LiCl droplets)³³ are used with the geometrical optics approximation to
144 determine the radius of the droplet.³⁴ An additional 355-nm laser (JDS Uniphase) is introduced co-
145 axially with the 532 nm laser and is used to excite fluorescence in the confined droplet.

146 The BQT is housed within an environmentally controlled chamber and most experiments
147 were performed in a high relative humidity (RH ~90 %) atmosphere generated by passing 200-500
148 sccm of nitrogen gas through a water bubbler. The RH of the gas is measured with two separate
149 RH sensors (Honeywell International, Inc.) located at the inlet and outlet of the chamber. The
150 nitrogen flow exerts a downward force on the trapped droplets and facilitates droplet ejection into
151 the mass spectrometer when the DC trapping voltage is removed.

152 **b. Merging Droplets in the BQT:**

153 In a typical merging experiment, a single droplet is dispensed from the side arm dispenser
154 and held in the lower trap. A second droplet from the vertical dispenser is generated and merges
155 with the confined droplet. The voltages applied to the induction electrodes are set to produce
156 droplets of opposite polarity (droplets with the same charge polarity will not merge). The initial
157 droplet (from the side arm dispenser) has a larger net charge than that from the vertical dispenser.
158 When droplets merge, there is a decrease in the overall charge and an increase in the mass of the
159 droplet, which both cause the merged droplet to fall lower in the trap. The initial position of the
160 droplet is restored by increasing the voltage applied to the balancing electrodes using the voltage
161 feedback loop. Droplet size is determined from the fringe separation from Mie scattering in the far
162 field image both before and after merging the event. The change in radius with the merging event
163 is used to infer the size of the merged droplet. As shown in Figure 2, the radii of the triggered and
164 merged droplets are very repeatable (± 200 nm, 0.7 % relative standard deviation). The size of the
165 held droplet is controlled by changing the water activity of the initial solution with a soluble salt
166 (~0.4-6 M LiCl) and allowing the droplet to equilibrate with the trap conditions. The size of the
167 merging droplet is controlled by changing the shape (i.e., magnitude and duration) of the square
168 wave electrical pulse used to generate the droplet with the piezoelectric microdispenser.³⁵ As the

169 merging droplet diameter increases, its terminal velocity changes from 0.07 to 0.15 m/s for 39 and
170 55 μm diameter droplets, respectively. Compared to previously reported droplet merging
171 approaches in an EDB,²⁸ the BQT approach does not separate droplet coalescence from
172 spectroscopic study. Thus, fast reactions (such as mixing dynamics in droplets) can easily be
173 studied using the BQT.

174 We have applied two different methods to study the merging process and the evolving
175 chemical composition of droplets. Fluorescence emitted by the droplet is collected and focused
176 onto a photomultiplier tube (PMT, Hamamatsu Photonics) using a plano-convex lens ($f/30$ mm).
177 The fluorescence emitted by the droplet is used to quantify mixing times and reaction kinetics. A
178 paper spray (PS) ionization source is used for single droplet mass spectrometry measurements to
179 detect the products of a reaction.

180 **c. Mixing Times from Droplet Fluorescence:**

181 The mixing time in a merged droplet is measured to determine the fastest reactions that can
182 be studied using the BQT. This time is measured by quantifying the acid-induced quenching of
183 Rhodamine-B (RhB), a fluorescent dye. Fluorescence from a droplet containing 500 μM RhB and
184 LiCl is excited using the 532 nm laser and detected with the PMT using a 550 nm long pass filter
185 to remove any elastically scattered light. Fluorescence from the RhB droplet is quenched by
186 merging it with a sulfuric acid droplet (2-20% v/v). The decay of fluorescence intensity upon
187 droplet coalescence is measured with the PMT (data acquisition rate = 500 kHz). Mixing is studied
188 by: 1) changing the size of the trapped droplet and keeping the size of the merging droplet constant;
189 2) keeping the size of the trapped droplet constant and changing the size of the merging droplet;
190 and 3) changing the concentration of sulfuric acid in the merging droplet.

191 **d. Reaction Kinetics from Droplet Fluorescence:**

192 Fluorescence is used to measure chemical kinetics in a single droplet which can be directly
193 compared to those in the bulk solution. The chemical reaction between ortho-phthalaldehyde
194 (OPA) and alanine in the presence of dithiothreitol (DTT) yields an isoindole product that
195 fluoresces at ~400-500 nm (Scheme 1).^{36,37} Solutions of OPA and alanine are prepared in a 3 M
196 LiCl solution that is buffered with a 50 mM borate buffer (pH = 9). Alanine solutions have
197 concentrations of 5.3, 10.3, 15.3, 20.8 and 33.3 mM. OPA and DTT are mixed together (to form a
198 stable adduct)³⁷ and have concentrations of 5.3 mM and 7.8 mM, respectively. Fluorescence from
199 merged alanine and OPA droplets is excited by the 355 nm laser and measured with the PMT using
200 a 450±20 nm bandpass filter to remove elastically scattered laser light. At least 10 trials are used
201 for each reaction condition in the droplets. Reaction kinetics measured in droplets are compared
202 to those measured in the bulk as described in the Supporting Information.

203 **e. Single Droplet Paper Spray Mass Spectrometry:**

204 By coupling a paper spray (PS) ionization source to the exit of the BQT, we have developed
205 a new approach to determine how the chemical composition of a single droplet changes over the
206 course of a reaction. A schematic of the experimental setup is shown in Figure 3a. The details of
207 PS mass spectrometry and its applications have been previously described.^{38,39} Here, a PS
208 ionization source is generated by cutting a small triangle (~6 mm base, ~10 mm height) from
209 chromatography paper (Whatman 3MM), passing a solvent through it, and applying a 4-5 kV
210 potential. The tip of the PS is placed ~2.5 cm from the inlet of the mass spectrometer (Q-Exactive
211 Orbitrap, Thermo Fisher Scientific, Inc.). The large electric field at the tip of the paper causes a
212 spray to form that is directed toward the mass spectrometer. A 0.8-1.0 mL/hour flow of 1% formic
213 acid solution in methanol through the filter paper maintains continuous operation of the PS source.

214 The mass spectrometer was operated with a resolution of 17,500 and a maximum ion injection
215 time of 50 ms.

216 A 2-cm long piece of 1/4" stainless steel tubing is affixed to the exit of the BQT coaxial with
217 the trap. The BQT is positioned such that the exit of this tubing is ~1 cm above the tip of the PS
218 source. After merging, droplets are held in the trap for a fixed period of time. Following a delay,
219 the voltage applied to the balancing electrodes is removed and the droplet falls from the trap
220 (typically aided by the flow of humidified nitrogen), impinging on the tip of the PS source. The
221 flow of solvent dilutes the droplets and pushes the components in the droplet toward the tip of the
222 filter paper where they are ionized and sprayed into the mass spectrometer. Compared to
223 electrospray techniques with droplets,⁴⁰ the PS source slows down the ionization event and ensures
224 the entire droplet is sampled. Figure 3b shows a mass spectrum that is collected from a single, 50
225 μm diameter droplet of 0.2% citric acid (~0.6 ng of citric acid total). Due to its low vapor pressure
226 and previous use in our lab,⁴¹ citric acid is used to benchmark the sensitivity and reproducibility
227 of the single droplet PS mass spectrometry technique. The mass spectrometer is operated in
228 negative mode and the only peak observed is from deprotonated citric acid ($[\text{M-H}]^-$) at m/z 191.
229 Figure 3c shows the time profile of the peak at m/z 191. Each individual spike arises from the
230 impact of one droplet on the PS source. The average peak area of these droplets has a relative
231 standard deviation of ~15% (likely due to variation in where the droplet impacts the PS source).
232 The precision of this method could be improved with the use of an internal standard in the droplet.

233 The products of the reaction between OPA and alanine in the absence of LiCl are studied
234 using single droplet PS mass spectrometry. Because the large amount of LiCl that is used in the
235 fluorescence experiments significantly diminishes the ionization efficiency of the organic species
236 by the PS source, LiCl is not added for the single droplet mass spectrometry experiments. Solutions

237 of 10.7 mM OPA with 21.4 mM DTT and 30.0 mM alanine are prepared in a 50 mM borate buffer
238 (pH ~ 9). The mass spectrometer is operated in positive ion mode to study both the components of
239 pure, unreacted OPA and alanine droplets as well as merged, reacted droplets.

240 **III. Results and Discussion:**

241 In order to facilitate an understanding of the observed reaction kinetics, we first describe
242 the mixing times that are observed in merged droplets and the experimental parameters that control
243 them. Then, both kinetic and product analyses of the reaction between OPA and alanine in droplets
244 are presented. Reaction kinetics measured with fluorescence imaging are reported in Section III.b.
245 The OPA/alanine reaction products that were observed with single droplet PS mass spectrometry
246 are presented in Section III.c.

247 **a. Mixing Times in Merged Droplets:**

248 The fastest reaction kinetics that can be measured in a well-mixed droplet following
249 coalescence is dependent on the timescale for mixing in the merged droplets. Here, the time it
250 takes for a merged droplet to mix completely is measured by quantifying the quenching rate of a
251 fluorescent dye. RhB fluorescence is quenched by a change in pH of solution; at low pH
252 fluorescence is quenched almost entirely (Figure S-1, Supporting Information). Figure 4a shows
253 examples of how the fluorescence from differently sized RhB droplets (500 μ M) is quenched when
254 they merge with 20% (v/v) sulfuric acid droplets of a constant diameter (38 ± 3 μ m). The initial rise
255 in fluorescence intensity is due to the increase in the cross sectional area of the droplet illuminated
256 by the laser. As the coalesced droplet relaxes from initially dumbbell-shaped to spherical, its cross
257 sectional area changes, causing the observed fluorescence intensity to oscillate. A similar effect is
258 observed (and has been reported previously⁴²) in elastically scattered light (Figure S-2, Supporting

259 Information). The angular frequency and damping of this oscillation is related to the surface
260 tension and viscosity of the merged droplet, respectively.⁴² Droplets typically relax to a spherical
261 shape after $\sim 200 \mu\text{s}$. The fluorescence transients are fit to the following piecewise exponential
262 decay function to extract mixing times:

$$263 \quad I = \begin{cases} a + b, & t < c \\ a \cdot e^{-(t-c)/d} + b, & t \geq c \end{cases}, \quad (1)$$

264 where a is the quenched fluorescence intensity, b is the fluorescence intensity post-quenching, c
265 is the time delay between data collection and droplet merging, and d is the mixing time. The red
266 lines shown in Figure 4a are the fits of Eq. 1 to the fluorescence transients. Because the lifetimes
267 of surface oscillations are typically shorter than the observed mixing times and their magnitude is
268 less than the change due to quenching, surface oscillations are not explicitly considered in Eq. 1.

269 To better understand the observed mixing times, they are compared to the timescales for
270 molecular diffusion, viscous diffusion, and bulk convection. The characteristic time for molecular
271 diffusion (τ_{Diff}) is:

$$272 \quad \tau_{Diff} \sim \frac{L_{ch}^2}{D}. \quad (2)$$

273 This is the time needed for a molecule to diffuse one characteristic length (L_{ch}) in a fluid with mass
274 diffusivity D . For RhB in water, D is $4.2 \pm 0.3 \times 10^{-10} \text{ m}^2 \text{ s}^{-1}$.⁴³ L_{ch} for the merged droplet is
275 calculated from the held (D_{held}) and merging (D_{merge}) droplet diameters as follows:⁴⁴

$$276 \quad L_{ch} = \frac{2D_{held}D_{merge}}{D_{held} + D_{merge}}. \quad (3)$$

277 For the droplet conditions here, molecular diffusion time is typically $\sim 1\text{-}10 \text{ s}$.

278 The viscous diffusion timescale (τ_{visc})—which represents the time required for momentum
279 to diffuse one characteristic length scale in a fluid with kinematic viscosity ν —is:

$$280 \quad \tau_{visc} \sim \frac{L_{ch}^2}{\nu}. \quad (4)$$

281 For a ~ 3 M LiCl solution, ν is $1.4 \times 10^{-6} \text{ m}^2 \text{ s}^{-1}$.⁴⁵ For the droplet conditions here, viscous diffusion
282 is typically 1-10 ms.

283 Finally, the bulk convection timescale (τ_{conv}) is the time required for material to traverse
284 one characteristic length at a rate equal to the relative droplet velocity (U_{rel}):

$$285 \quad \tau_{conv} \sim \frac{L_{ch}}{U_{rel}}. \quad (5)$$

286 The relative velocity of the collision is calculated from the terminal velocity of the merging droplet
287 (0.07-0.15 m/s). As the merging droplet approaches the balancing electrode, its velocity increases
288 slightly due to Coulombic attraction. However, based on the voltage applied to the balancing
289 electrode, this acceleration is measured to be small (Supporting Information, Figure S-3). For
290 droplets here, the bulk convection time is typically ~ 200 - $600 \mu\text{s}$. Carroll and Hidrovo previously
291 demonstrated that as the inertia of the collision event increases, the observed mixing times is
292 shortened toward the bulk convection timescale.⁴⁴

293 Figure 4b shows the measured mixing times when the size of the held droplet is changed
294 and the diameter of the merging droplet is held constant ($D_{merge} = 38 \pm 3 \mu\text{m}$). As the size of the
295 held droplet decreases, the observed mixing time decreases toward the convection mixing time. A
296 log-log plot of initial droplet diameter vs. mixing time has a slope of 3.2 ± 0.2 (Figure S-4,
297 Supporting Information), implying that the mixing rate in this experiment scales with the volume
298 of the held droplet.

299 Figure 4c shows the mixing times when the size of the held droplet is constant ($D_{held} =$
300 $56 \pm 2 \mu\text{m}$) and the diameter of the merging droplet is changed. As the diameter of the merging
301 droplet increases, the diameter of the merged droplet increases (which causes the viscous diffusion
302 time to increase), and the terminal velocity of the merging droplet increases (which causes the bulk
303 convection time to decrease). The experimental data show that mixing times decrease with
304 increasing merging diameter. This is likely due to the increasing energy and inertia of the
305 collision.⁴⁴ If the inertia of the collision were to continue to increase, the mixing time is predicted
306 to follow the bulk convection time.⁴⁴ Finally, when the concentration of sulfuric acid is changed
307 (and the held/merging droplet diameters are kept constant), the mixing times do not change (Figure
308 S-5, Supporting Information). This suggests that the observed mixing times are controlled by the
309 size of droplets and velocity of collisions, and are not due to the concentration of reagents in the
310 in the droplets.

311 The observed mixing times reported here (reliably down to $\sim 400 \mu\text{s}$) are very similar to
312 those reported in free-droplet collision experiments (i.e. colliding droplets are not confined within
313 an electrodynamic balance).⁴⁶ They are also similar to those observed in conventional stopped flow
314 kinetics measurements ($\sim 2 \text{ ms}$),⁴⁷ but slower than those achieved in miniaturized continuous flow
315 methods such as theta capillary electrospray (mixing time $\sim 1 \mu\text{s}$)⁴⁸ and microfluidic channels
316 (mixing time $\sim 15 \mu\text{s}$).⁴⁷ Work by Lee *et al.* colliding a plume of high speed (80 m/s) 13- μm
317 droplets report mixing times of a couple of microseconds.⁶ The mixing times in the BQT allow for
318 the study of reactions with a bimolecular rate constant of up to $\sim 10^4$ - $10^5 \text{ M}^{-1} \text{ s}^{-1}$. As shown, faster
319 mixing times could be achieved by either using smaller droplets or increasing the velocity of the
320 collision. The latter could be accomplished either electrostatically (e.g. apply a higher potential to
321 the balancing electrodes, Figure S-3) or with a faster flow of gas through the trap.

322 **b. Chemical Reactions in Merged Droplets**

323 The reaction between OPA and alanine in the presence of DTT (Scheme 1) is studied in
324 droplets with the BQT and bulk solution using fluorescence spectroscopy. LiCl is added to the
325 solutions to decrease the water activity such that evaporation from the droplet is minimized and
326 the conditions in the droplet can be reproduced in the bulk. Because the water activity of a 3.0 M
327 LiCl solution is 0.87,⁴⁹ experiments are performed with a RH close to 87%, and the size of the
328 droplet does not change considerably over time of the experiment (Figure S-6, Supporting
329 information). In bulk measurements, the volumes of the reactant solutions are mixed in a 1:1 ratio,
330 and the initial reactant concentrations are half the value of the prepared solutions. In droplets, the
331 initial reactant concentrations are determined using the size of the merging droplets ($24.3 \pm 0.2 \mu\text{m}$
332 and $23.0 \pm 0.6 \mu\text{m}$ radius for alanine and OPA, respectively). The merged droplet has a radius of
333 $29.9 \pm 0.4 \mu\text{m}$. While keeping the initial OPA concentration constant ($\sim 2.6 \text{ mM}$), the rate of
334 fluorescence appearance is measured at various alanine concentrations ($\sim 2.6, 5.2, 7.7, 10.4$ and
335 16.7 mM). Figure 5 shows an example of the bulk and droplet fluorescence data that are collected
336 with ~ 2.6 and $\sim 7.7 \text{ mM}$ initial OPA and alanine concentrations, respectively. Fluorescent intensity
337 in the droplet appears at a slightly faster rate than in the bulk solution. To quantify the reaction
338 rate constants in the merged droplet and bulk, the OPA and alanine reaction is simulated and fit to
339 experimental data.

340 When the alanine concentration exceeds the OPA concentration, the final fluorescence
341 intensity does not change with increasing amounts of alanine. At these conditions, it is assumed
342 that the final concentration of the fluorescent product is equal to that of the initial OPA
343 concentration. Using this scaling, the measured fluorescence intensity is converted to
344 concentration of fluorescent product. The reaction between OPA and alanine has previously been

345 shown to follow bimolecular kinetics.³⁶ Thus, to quantify the rate of reaction in bulk and droplets,
346 an ordinary differential equations solver is used to simulate a bimolecular reaction with the initial
347 reactant concentrations set to experimental values. The bimolecular rate constant in the simulation
348 is varied to best match the simulated and experimental product concentrations. The same kinetic
349 analysis is used for both bulk and droplet experiments. The solid lines in Figure 5 represent the
350 simulated product concentrations with the optimized rate constant. The average bimolecular
351 reaction rate constants in the bulk and droplet are 67 ± 6 and 84 ± 10 $\text{M}^{-1} \text{s}^{-1}$, respectively.
352 Uncertainty corresponds to the standard deviation of the rate constants extracted at each reaction
353 condition. The simulated fits and individual rate constants extracted at each reactant condition are
354 tabulated in the Supporting Information (Figure S-7 and Table S-1). The bimolecular rate constant
355 for the reaction of alanine with OPA in the presence of DTT has previously been measured to be
356 60 ± 4 $\text{M}^{-1} \text{s}^{-1}$,³⁶ which is in good agreement with the bulk rate constant reported here.

357 The average rate constant in the 30 μm radius droplet is roughly 25% larger than the rate
358 constant in the bulk. When the polarity of the charge on the droplet is reversed (i.e. the merged
359 droplet has a net positive charge instead of net negative charge) the kinetics of the reaction are
360 unchanged (Figure S-8 in Supporting Information). This suggests that the small amount of charge
361 on the droplet surface does not affect the overall rate of reaction. Because evaporation from the
362 particle is minimized with the addition of LiCl, the small observed rate enhancement could
363 originate from enhanced surface to volume ratio in the droplet compared to the bulk. Fallah-Araghi
364 *et al.* previously measured the kinetics of a bimolecular reaction in aqueous droplets in an oil-
365 water emulsion. They observed a maximum rate enhancement by factor of ~ 40 that decreased with
366 increasing droplet radius. The observed enhancement was attributed to the increasing surface to
367 volume ratio at smaller droplet sizes. A weak adsorption of molecular species to the oil-water

368 interface was predicted to change the energetics of the reaction to favor product formation.
369 However, a rate enhancement was only observed in emulsions that had a radius smaller than ~20
370 μm .⁷ The merged droplets in this study have a radius of $29.4 \pm 0.4 \mu\text{m}$, which could indicate that
371 the droplets used here are still too large to observe a significant rate enhancement from interfacial
372 effects.

373 **c. Single Droplet Mass Spectrometry**

374 The products of the reaction between OPA and alanine are studied using the PS ionization
375 source. Figure 6a shows mass spectra from a single alanine droplet (black line) and OPA solution
376 droplet (red line). Compared to the concurrently developed single droplet mass spectrometry
377 method (which uses a corona discharge to ionize the vaporized droplet components),²⁷ the use of
378 the PS ionization source leads to less fragmentation and easier identification of reaction products.
379 The alanine spectrum has only one peak at m/z 90, which represents the protonated molecular ion.
380 The OPA spectrum has peaks at m/z 135 and 157, which are from the protonated OPA molecule
381 and the OPA/ Na^+ complex, respectively. The peak at m/z 311 corresponds to the OPA and DTT
382 adduct complexed with a Na^+ ion. OPA reacts with thiol containing compounds to create a stable
383 1,3-dihydroisobenzofuran compound that has been previously observed.³⁷ Because DTT has two
384 reactive thiol groups, a single DTT molecule can react with two OPA molecules. The peak at m/z
385 445 corresponds to this reaction product complexed with a Na^+ ion. Figure 6a also shows the mass
386 spectrum of a single merged droplet after it has reacted for 6 s (blue line). Because alanine is in
387 excess in the merged droplet, only the intensities of the peaks from OPA-containing species have
388 decreased significantly (the alanine peak remains a dominant peak). The peaks present at m/z 342
389 and 476 correspond to the protonated isoindole reaction products.^{36,37} The chemical structures for
390 each of the ions are shown in the Supporting Information (Table S-2).

391 Figure 6b shows a selected ion chromatogram from the ejected merged droplets for each
392 of the peaks of interest in the experiment (m/z 90, 311, 342, 445, and 476). Each of the ions show
393 a similar time response, and are only present when a droplet is ejected onto the PS source. A ~ 5
394 mM aqueous droplet with a radius of $30\ \mu\text{m}$ (similar to the OPA conditions in this experiment) has
395 ~ 0.5 pmol of material (< 1 ng). As shown in Figure 6b, this results in single droplet pulses with a
396 signal to noise (S/N) ratio > 100 . Assuming a S/N ratio of ~ 10 is necessary to quantify peak
397 intensities, a single, ~ 5 -mM droplet with a radius of $\sim 15\ \mu\text{m}$ could easily be detected in the current
398 configuration.

399 **IV. Conclusion:**

400 A branched quadrupole trap has been designed and constructed to merge confined droplets.
401 This trap allows for new measurements of homogeneous chemical reactions in droplets. Through
402 the quenching of fluorescence of RhB droplets by sulfuric acid droplets, consistent mixing times
403 as short as $\sim 400\ \mu\text{s}$ are obtainable using droplets moving at relative velocities of ~ 0.1 m/s. As
404 predicted by Carroll and Hidrovo,⁴⁴ mixing experiments in the BQT show that faster mixing times
405 are achievable by either decreasing the size of the merging droplets or increasing the speed of the
406 collision. With these mixing times, chemical reactions with a bimolecular rate constant up to $\sim 10^4$ -
407 $10^5\ \text{M}^{-1}\ \text{s}^{-1}$ can be studied in the BQT in its current form.

408 The ability to measure homogeneous chemical reactions in the BQT has been demonstrated
409 using both fluorescence spectroscopy and single droplet PS mass spectrometry. The reaction of
410 OPA with alanine (in the presence of DTT) is found to occur slightly faster ($\sim 25\%$) in a droplet
411 with a radius of $\sim 30\ \mu\text{m}$ than in bulk solution. Charge on the droplet and changes in the
412 concentration of reactants due to evaporation do not play a significant role in any potential rate
413 enhancement in the fluorescence experiments reported here. Thus, the small rate enhancement is

414 attributed to the larger surface to volume ratio of the droplet compared to the bulk. Single droplet
415 PS mass spectra of reacted droplets following merging show the expected reaction products. Based
416 on the observed signal levels, it is estimated that single droplets with a radius of ~15 μm with <100
417 pg of analyte could be easily detected using PS mass spectrometry.

418 We have developed a new technique for the contactless manipulation and merging of
419 micron-sized droplets to initiate chemistry. We demonstrate the applicability of fluorescence
420 imaging for measuring reaction kinetics and demonstrate the use of mass spectrometry coupled
421 with a single particle trap. Going forward, these developments will allow for rigorous probing of
422 reaction kinetics in a variety of samples spanning a wide range of sizes and concentrations.

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431 **References:**

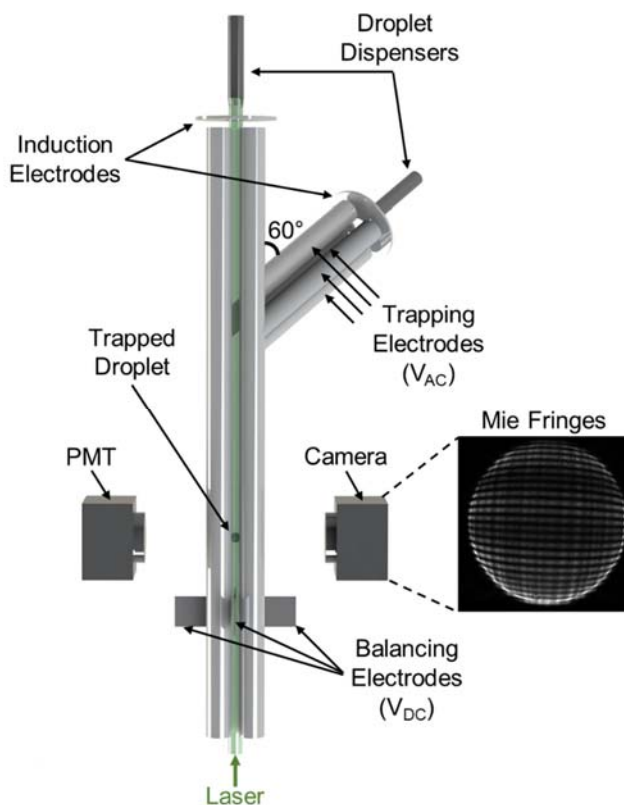
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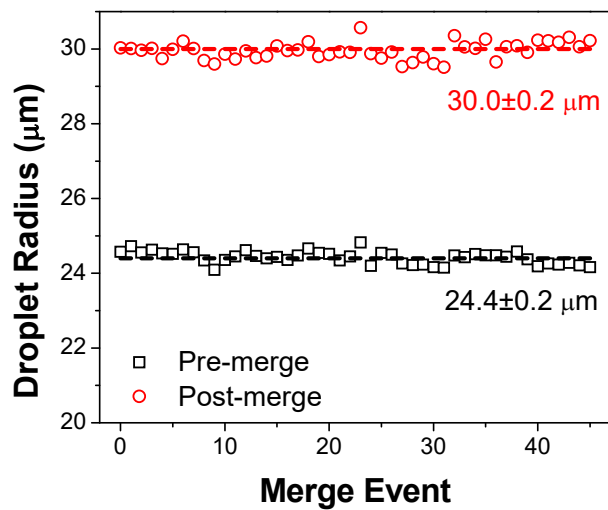
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503
 504 **Figure 1.** Schematic of the BQT. Different solutions necessary for a chemical reaction are
 505 dispensed from the two droplet dispensers. Thus, when the droplets from each of the dispensers
 506 coalesce, a chemical reaction in the merged, trapped droplet can be initiated. The chamber that
 507 houses the BQT setup and the optics used to collect and collimate the light for the camera and
 508 PMT are not shown. An example of the far field image of elastically scattered light used to
 509 determine the size of the trapped droplet is shown.

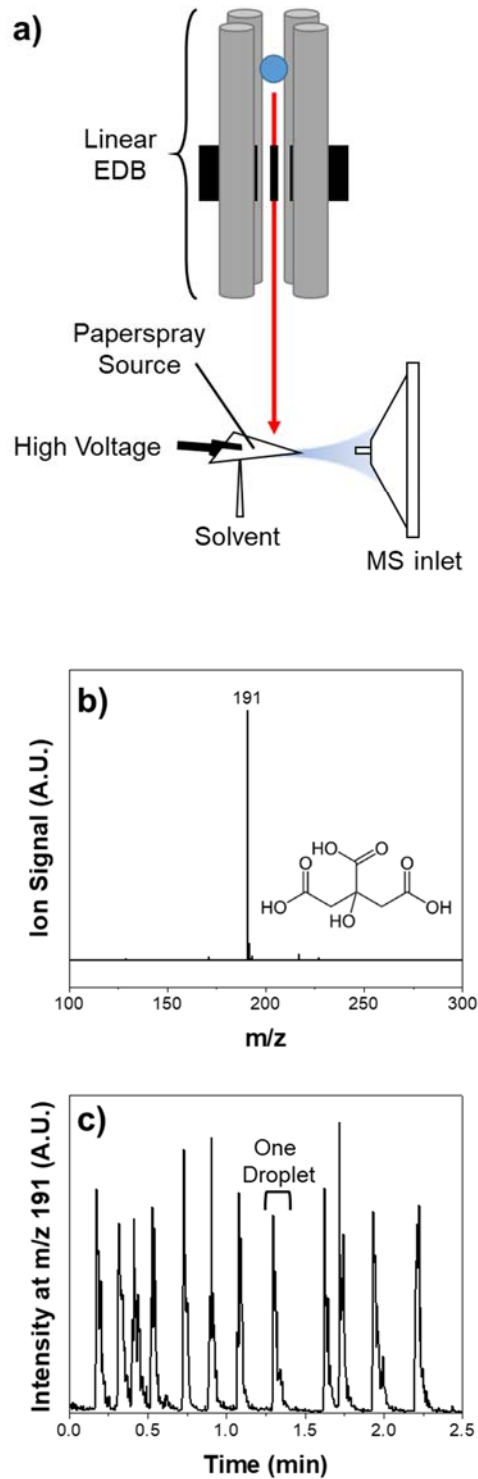
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512 **Figure 2.** Measured droplet radius before and after merging for 45 separate coalescence events
513 (~2 hours of operation). Droplets contained ~3 M LiCl. The dashed lines show the average droplet
514 radius.

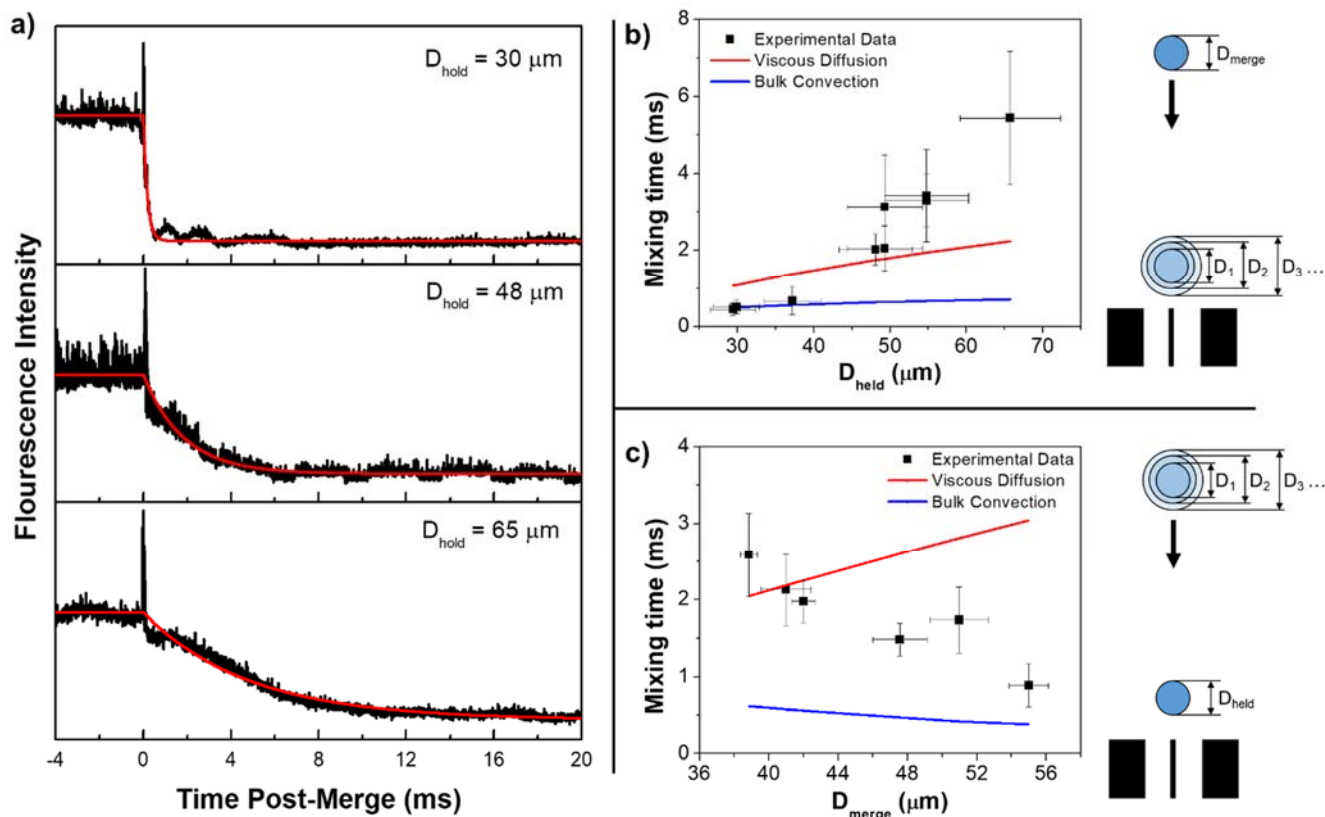
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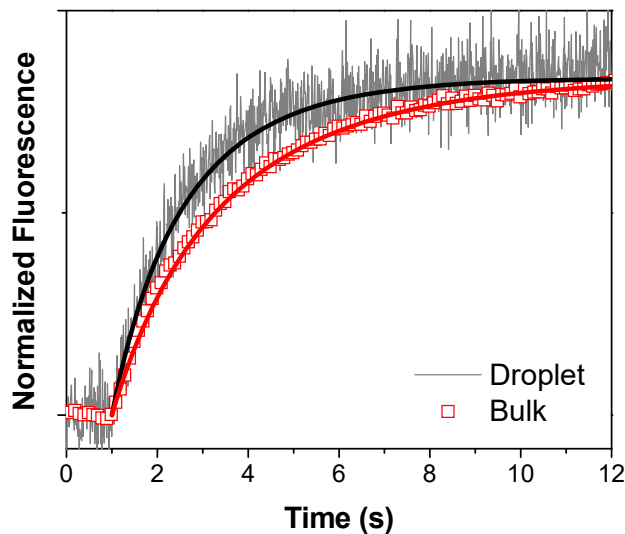
518 **Figure 3.** a) Schematic of the BQT/PS mass spectrometry interface. b) Mass spectrum from a
 519 single 50- μm diameter, 0.2% (w/v) citric acid droplet. c) Selected ion chromatogram of citric acid
 520 droplets. Each spike in the chromatogram represents one 50- μm , 0.2% citric acid droplet impacting
 521 the PS source.



522

523 **Figure 4.** a) Fluorescent quenching of differently sized RhB droplets with the coalescence of a
 524 $38 \pm 3 \mu\text{m}$, 20% (v/v) sulfuric acid droplet. The fluorescence intensity measured with the PMT
 525 (black line) is fit to Eq. 1 to extract a mixing time. The fit to the data is shown by the red line. The
 526 data shown have mixing times of 0.191 ± 0.006 , 1.82 ± 0.03 and 4.93 ± 0.06 ms for the 20, 48, and 65
 527 μm RhB droplets, respectively. b) Mixing times for experiments where the merging droplet
 528 diameter (sulfuric acid) is kept constant ($38 \pm 3 \mu\text{m}$) and the held droplet diameter (RhB) is varied.
 529 The viscous diffusion and bulk convection times are shown in red and blue, respectively. The
 530 molecular diffusion times are too long to show on this scale. The experimental mixing times scale
 531 with the volume of the held droplet. c) Mixing times for experiments where the merging droplet
 532 diameter (sulfuric acid) is increased and the held droplet diameter (RhB) is kept constant (56 ± 2
 533 μm). As the diameter of the merging droplet increases, the mixing time decreases.

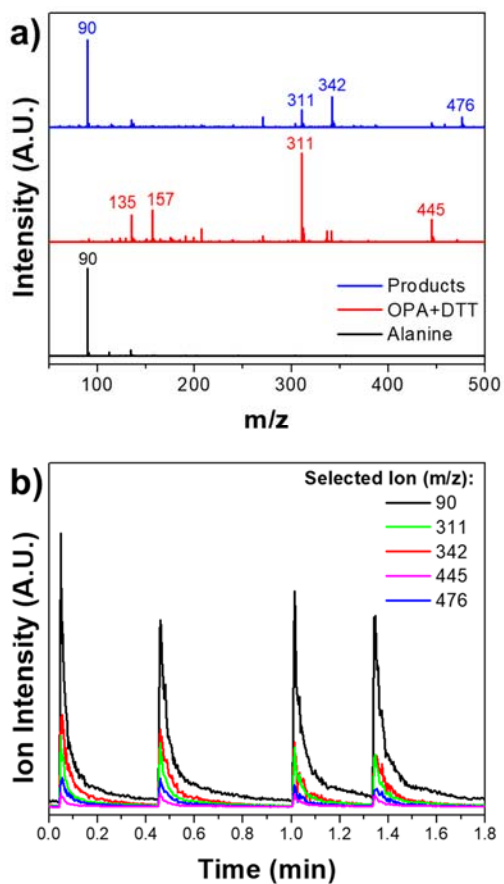
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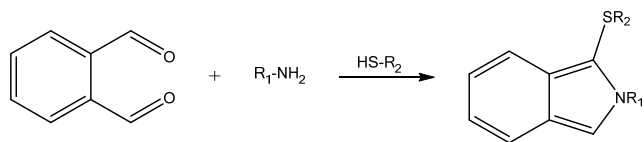
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536 **Figure 5.** Fluorescence (450 ± 20 nm) generated from the reaction of OPA with alanine in a droplet
 537 with a radius of 29.9 ± 0.4 μm (gray line) and in bulk solution (red squares). In the droplet, the
 538 initial OPA concentration is 2.4 ± 0.1 mM and the initial alanine concentration is 8.1 ± 0.4 mM. In
 539 bulk solution, the initial concentration of OPA is 2.6 mM and the initial concentration of alanine
 540 is 7.7 mM. The solid black and red lines are the best-fit product concentrations from the
 541 bimolecular reaction simulation. The average bimolecular rate constant for the reaction of alanine
 542 with OPA is found to be 84 ± 10 and 67 ± 6 $\text{M}^{-1} \text{s}^{-1}$ in the droplet and bulk, respectively. The
 543 individual rate constants for the different reaction conditions are tabulated in Table S-1.

544



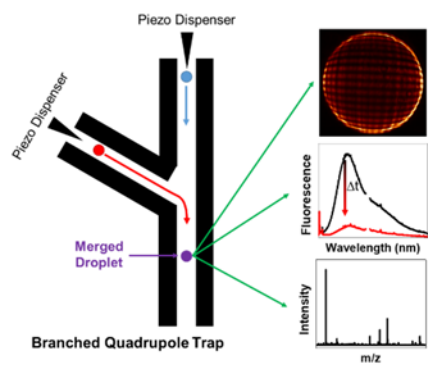
545
 546 **Figure 6.** a) Mass spectra of single droplets from alanine solution (black line), OPA/DTT solution
 547 (red line) and merged alanine/OPA droplets after 6 s of reaction (blue line). The new peaks in the
 548 products spectrum (m/z 342 and 476) correspond to the expected fluorescent products. The
 549 chemical structures of labeled peaks are given in the Supporting Information (Table S1). b)
 550 Selected ion chromatograms showing the time response of each peak of interest in the merged
 551 droplets. The signal to noise ratio for each of these peaks is >100.



552

553 **Scheme 1.** The reaction of ortho-phthalaldehyde (OPA) with a primary amine (alanine) in the
554 presence of a thiol group (dithiothreitol, DTT) yields a fluorescent isoindole compound.

555



556

557 TOC Graphic