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Authors: Xuewei Wang; Meng Sun; Stephen Ferguson; J. Damon Hoff; Yu Qin; Ryan Bailey; Mark Meyerhoff

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Ionophore-based biphasic chemical sensing in droplet microfluidics

Xuwei Wang,^{*[a]} Meng Sun,^[a, b] Stephen A. Ferguson,^[a] J. Damon Hoff,^[b] Yu Qin,^[a] Ryan C. Bailey,^[a] and Mark E. Meyerhoff^[a]

Abstract: Droplet microfluidics is an enabling platform for high-throughput screens, single cell studies, low-volume chemical diagnostics, and microscale material syntheses. Analytical methods for real-time and *in situ* detection of chemicals in the droplets will greatly benefit these applications, but they remain limited. Herein, we report a novel heterogeneous chemical sensing strategy based on functionalization of the oil phase with rationally combined sensing reagents. Sub-nanoliter oil segments containing pH-sensitive fluorophores, ionophores, and ion-exchangers enable highly selective and rapid fluorescence detection of physiologically important electrolytes (K^+ , Na^+ , and Cl^-) and polyions (protamine) in sub-nanoliter aqueous droplets. Electrolyte analysis in whole blood is demonstrated without suffering from optical interference from the sample matrix. Moreover, an oil phase doped with a boronic acid-appended aza-BODIPY dye allows indication of H_2O_2 in the aqueous droplets, exemplifying sensing of targets beyond ionic species.

Over the past two decades, droplet microfluidics has emerged as a unique subcategory of microfluidics that has found numerous applications in drug screening, directed evolution, single-cell analysis, and medical diagnosis, as well as for the synthesis of materials and molecules.^[1] In pressure-driven droplet microfluidics, an aqueous stream and a water-immiscible oil stream intersect via a microchannel junction, such as a T-junction or a flow-focusing geometry. Discrete and monodisperse aqueous droplets or plugs are generated at femtoliter to nanoliter volumes at a frequency ranging from Hz to kHz. This technique provides new opportunities to screen drug candidates and enzyme mutants in a reagent-conservative and high-throughput fashion, to perform chemical testing in small volumes of bodily fluids for minimally invasive diagnostics, and to study individual cells/bacteria/viruses with low dilution factors and negligible liquid evaporation. The effective internal circulation within the flowing liquid segments and the favored interfacial mass transfer between microscale compartments also make droplet microfluidics an efficient platform for chemical synthesis and liquid-liquid extraction.^[1a,2]

In these applications, quantitative analysis of (bio)chemicals within droplets is essential for the determination of biomarker concentrations, the indication of enzymatic reactions/chemical interactions, examination of biochemical

activities of cells/bacteria, and the monitoring of synthesis and extraction processes. Indeed, a wide variety of detection modalities including fluorescence spectroscopy, bright-field microscopy, infrared spectroscopy, surface-enhanced Raman spectroscopy, mass spectrometry, nuclear magnetic resonance spectroscopy, and electrochemistry have been employed in droplet microfluidics.^[3] Among them, fluorescence-based techniques are perhaps most common because their fast measurement rates can match the high droplet generation frequency and their high sensitivity is suited to the very low volumes of sample confined in the droplet. Current fluorescence-based tests are primarily restricted to enzyme assays (e.g., alkaline phosphatase^[4] and β -galactosidase^[5]) using fluorogenic substrates, metabolite detection (e.g., L-lactate^[6] and ethanol^[7]) using coupled enzyme reactions, and bioaffinity assays (e.g., immunoassay^[8] and DNA hybridization assays^[9]) using fluorophore-labeled biomacromolecules. Also, these tests exclusively occur in the aqueous phase of the droplet microfluidics. Herein, we, for the first time, exploit the use of the oil segments in droplet microfluidics as a chemical sensing phase for targets in their adjacent aqueous droplets. This new strategy not only greatly extends the spectrum of analytes that can be quantified via the droplet microfluidics technology but also offers several key inherent advantages compared to chemical sensing in aqueous droplets. First, many host-guest recognition chemistries such as ionic complexation are favored in a low dielectric constant medium.^[10] Second, the low aqueous solubility is a challenge in the design of certain molecular probes, which is another reason that addition of an organic solvent into the assay solution is commonly practiced; in contrast, the design of highly hydrophobic probes is relatively easy by the introduction of hydrophobic moieties such as long alkyl chains. Third, the oil segments as sensors do not suffer from optical interference from the color and/or turbidity of the aqueous sample matrix. Lastly, fabrication and operation of the microfluidic chip are simplified because there is no need to use an additional aqueous stream to introduce sensing reagents into the sample phase.

The first targets of our methodology are electrolytes, such as K^+ , Na^+ , and Cl^- . Quantification of these ions is important for medical diagnostics, monitoring of ion-channel functions, and study of ion binding to biomacromolecules. Droplet microfluidics may greatly benefit these applications, but *in situ* sensing of these electrolytes has not been reported in droplet microfluidics. The most successful technique for electrolyte analysis is the ionophore-based ion-selective sensors (electrodes and optodes).^[11,12] Interestingly, these sensors rely on a water-immiscible organic phase such as plasticized membranes or micro/nano-particles to test analytes in an aqueous sample, which resembles the biphasic scaffold of droplet microfluidics. Therefore, we take advantage of this similarity to incorporate ion sensing functionalities into droplet microfluidics. Notably, there is one report on the use of an ion-selective electrode to detect ions (Mg^{2+}) in a continuous aqueous phase separated from droplet

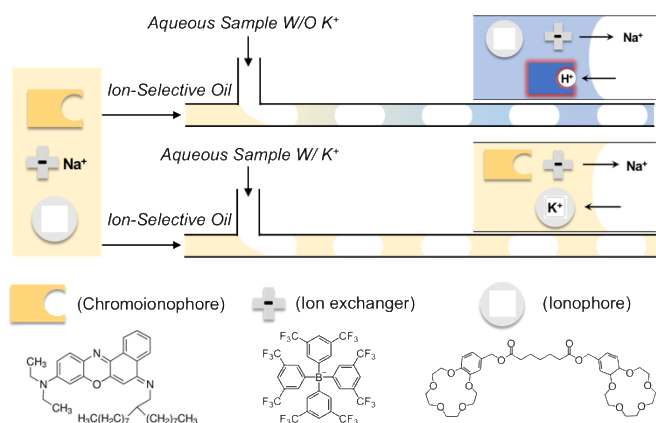
[a] Dr. X. Wang, Dr. M. Sun, Dr. S. Ferguson, Prof. R. Bailey, Prof. M. Meyerhoff
Department of Chemistry
University of Michigan
930 N University, Ann Arbor, MI 48109, USA
E-mail: wangxue@umich.edu

[b] Dr. M. Sun, Dr. J. D. Hoff
Department of Biophysics
University of Michigan
930 N University, Ann Arbor, MI 48109, USA

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microfluidics,^[13] but the requirement of phase separation sacrifices most attractive features of droplet microfluidics such as the throughput and the capability to study individual cells.

Scheme 1 shows the principle of optical sensing of cations (K^+ as an example) in droplet microfluidics. Highly hydrophobic sensing chemicals including a pH indicator dye (fluorescent chromoionophore) as an optical read-out element, an ionophore as the ion recognition element, and a cation exchanger to prevent interference from anions are dissolved into a water-immiscible oil. By using two infusion pumps, this oil phase is merged with an aqueous solution within a T-junction microchannel under conditions that generated segmented flow. For a buffered aqueous sample without the target cations, protons from this sample can transfer into the oil phase and protonate a chromoionophore having an appropriate pK_a . This process is accompanied by expulsion of the hydrophilic Na^+ from the oil segments into the aqueous droplets to maintain electroneutrality of the oil. In contrast, when the aqueous sample contains target cations, these ions will be extracted into the oil segments due to the very high binding affinity of the ionophore to the target ion. This ionic complexation process competes with the protonation of the chromoionophore, and the degree of protonation determines the absorbance and fluorescence properties of the pH-sensitive dye in the oil segments.



Scheme 1. Operation principle of biphasic K^+ sensing platform in droplet microfluidics.

The color and fluorescence of the oil phase can be monitored by high-speed bright-field video and time-resolved laser-induced fluorescence, respectively, at the distal end of the downstream channel, which is 3.5 cm from the junction. As can be seen from Figure 1A, the oil segments are blue when the K^+ concentration is low and become orange for a higher K^+ concentration, which results from the protonated and deprotonated chromoionophore, respectively (see Figure S1A for the absorbance spectra). The color difference is small because of the short optical path length ($40\ \mu\text{m}$) in the microchannel. Therefore, we employ fluorescence spectroscopy to perform quantitative ion analysis. Based on the fluorescence spectra of the protonated and deprotonated chromoionophore in bulk dioctyl sebacate (Figure S1B), an excitation laser at 630 nm and an emission wavelength band of 673 – 738 nm were chosen. As shown in Figure 1B, significant fluorescence is observed only from the oil segments phase because of the high hydrophobicity of the dye (chromoionophore III, $\log P=10.5$, from ChemDraw). A higher concentration of K^+ in the aqueous phase leads to a lower

fraction of protonated chromoionophore molecules, which yields less fluorescence in the oil segment due to the unfavored charge transfer between the donor and acceptor groups in the chromoionophore. The mean fluorescence signal (F) of all oil segments during a 0.5 s period is obtained from a Gaussian fit of the photon intensity (photons per second) emitted from the oil phase. The fluorescence of the completely protonated (F_{max}) and deprotonated (F_{min}) chromoionophore in the segmented oil is acquired when the aqueous phase consists of 0.1 M HCl and 0.1 M NaOH, respectively. The degree of protonation ($1-\alpha$) is then calculated by the following equation:

where $[CH^+]$ is the concentration of the protonated

$$1 - \alpha = \frac{[CH^+]}{[C_T]} = 1 - \left(\frac{F_{max} - F}{F_{max} - F_{min}} \right)$$

chromoionophore, and $[C_T]$ is the total concentration of the chromoionophore. As is convention in the field of ion-selective optodes,^[12] we plot the degree of protonation against the concentration of the analyte ion in the aqueous phase to create the calibration curve (Figure 1C). Like traditional ion-selective optodes,^[12] this sensing method has a wide dynamic range covering several orders of magnitude of analyte concentration. Half protonation of the chromoionophore occurs at around 10^{-3} M K^+ ($[K^+]/[H^+] = 10^{4.4}$), which represents a sensitivity similar to those of other K^+ optodes.^[14] The standard deviation shown in Figure 1C could be translated into a concentration error of $\pm 3\%$ for 10^{-3} M K^+ and $\pm 1\%$ for 10^{-2} M K^+ based on the linear regression formula shown in Figure S2. Furthermore, because of the specificity of the ionophore, this system is at least 1000-fold more sensitive toward K^+ than other cations including Na^+ , Li^+ , Ca^{2+} , and Mg^{2+} .

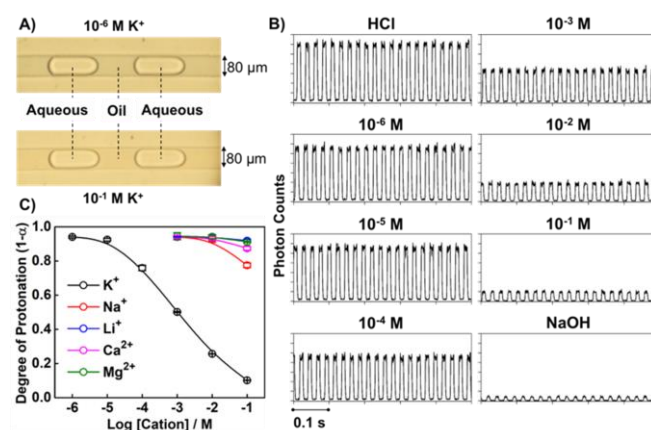


Figure 1. A) Bright-field images of the segmented fluids in the presence of 10^{-6} M and 10^{-1} M KCl in the aqueous phase. The used oil is dioctyl sebacate, and the chip is made of PDMS; B) Fluorescence trace of the segmented flow in the presence of different concentrations of KCl in Tris-HCl buffer at pH 7.4 and 0.1 M HCl or NaOH; C) Response curve of this sensing method toward different cations based on the 0.5-s test (data points are average \pm SD for $n=3$ measurements).

One difference of this sensing scheme from traditional ion-selective optodes is the equal volume of the sample phase and the sensing phase. Depletion of the analyte in the sample becomes possible, which is similar to the ion-selective nano-optodes operating in exhaustive sensing mode.^[15] According to the deprotonation of the chromoionophore, 4.5% and 0.7% of K^+ is extracted into the oil phase when the sample has 10^{-3} M and 10^{-2} M of KCl, respectively.

Based on the flow rate (2 $\mu\text{L}/\text{min}$ for each phase) and the frequency of segment generation (~ 41 segments per second for each phase), the size of each oil segment and each aqueous droplet is estimated to be ~ 0.8 nL. A test time of 0.5 s corresponds to an aqueous sample volume of ~ 17 nL and ~ 21 measurement events. Taking the 10^{-3} M KCl sample as an example, we obtained a relative standard deviation of only 1.2% for the averaged fluorescence from the 21 oil segments. The negligible variation between different oil segments and their fluorescence intensities may allow further reduction of the sample volume and the measurement time. The use of an ultra-small volume of sample to perform a test is critical for the concept of minimally invasive diagnostics such as those based on fingerstick blood sampling or a microneedle-based painless blood draw, especially when a large number of parameters are to be quantitated from a single sample.

Figure 2 shows the fluorescence change as a function of distance along the channel after droplet generation. When the aqueous phase has 10^{-6} M KCl, the fluorescence of the oil phase reaches 95% equilibrium after 2.5 cm of travel, corresponding to the process of chromoionophore protonation. This 2.5-cm distance takes ~ 1.8 s to traverse based on a moving velocity of ~ 170 μm per 12 ms (each segment). For a high concentration of K^+ , the chromoionophore in the oil phase only needs to be partially protonated or remains deprotonated, which takes a shorter time to reach equilibrium. Such response times are more than 50-fold shorter than conventional polymeric membrane-type ion-selective optodes with film thicknesses of a couple of μm ^[14a] and microsphere-type ion-selective optodes with diameters of ~ 20 μm .^[14b] This is also faster than ion sensing in parallel flow-based microfluidics.^[16] The very fast response time is likely related to the lower viscosity of the pure liquid oil phase compared to the commonly used polymer-plasticizer mixture, and the enhanced convection within both aqueous droplets and oil segments due to the friction-induced internal flow

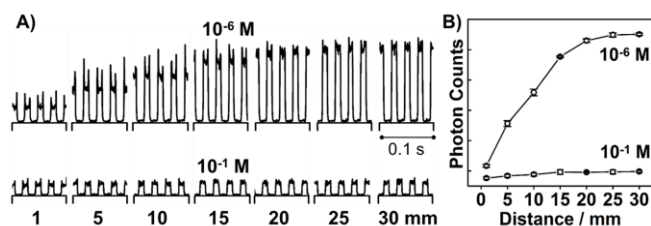


Figure 2. A) Fluorescence trace of the segmented flow at different locations along the channel after the merging junction; B) Mean photon intensity of the oil segments as a function of distance from the junction (data points are average \pm SD for $n=3$ measurements).

circulation.^[17]

Traditional fluorescent probes are usually susceptible to optical interference from colored and/or turbid samples such as whole blood. Indeed, although the formation of stable blood droplets has been reported in an image-based study of blood coagulation,^[18] optical chemical analysis in whole blood has not, to the best of our knowledge, been reported previously in droplet microfluidics. Interrogation of such complicated samples is also quite challenging for other analytical techniques such as mass spectrometry and Raman spectroscopy. In contrast, our biphasic sensing scheme uses physically separated sensors and samples. Since both the laser illumination and the fluorescence collection are perpendicular to the PDMS chip, the signal generated from the oil segment can be monitored without suffering from optical interference from the blood. As shown in

Figure 3A, the blood droplets do not exhibit any fluorescence under the employed experimental conditions, but they are able to induce fluorescence in the oil segments based on ion extraction. The K^+ concentration in the blood sample is calculated to be 2.6 ± 0.4 mM, which reasonably matches the concentration of 2.2 mM obtained by a commercial blood gas/electrolyte analyzer (Figure S2). The error may be related to spontaneous hemolysis of blood in the microchannel. Indeed, when the blood sample was sonicated for 5 s before being introduced into the microfluidics chip, significantly reduced fluorescence is observed from the oil segments due to the release of K^+ from the broken blood cells (Figure 3). Extraction of lipophilic compounds from biological samples into the oil segment might be another source of error. Such non-selective extraction could be prevented by using a fluoruous oil phase with perfluorinated sensing chemicals.

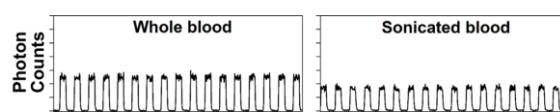


Figure 3. Fluorescence trace of the segmented flow for analysis of whole blood and sonicated whole blood. A 1:1 dilution of the whole blood with 0.1 M Tris-HCl buffer at pH 7.4 was employed to control the sample pH.

One feature of ionophore-based ion-selective detection using combined reagents is that the selectivity and the sensitivity can be adjusted by using different sensing reagents and different ratios of those reagents. For example, the use of a Na^+ ionophore and a tetraphenylborate-type cation exchanger in its potassium salt form renders the oil segments selective toward Na^+ (Figure S3). By using a mercuracarborand "anti-crown ether" ionophore for Cl^- , a quaternary ammonium-type anion exchanger, and a less basic chromoionophore (chromoionophore I), fluorescent sensing of Cl^- can also be achieved (Figure S4). In addition, highly selective ionophores for other inorganic cations such as Ca^{2+} , Mg^{2+} , Pb^{2+} , Zn^{2+} , Cd^{2+} , Hg^{2+} , Ag^+ , and other inorganic anions such as F^- , NO_3^- , NO_2^- , SO_4^{2-} , CO_3^{2-} are available.^[11] A wide range of organic ions (e.g., creatinine and choline derivatives),^[19] polyions (e.g., protamine, heparin, and DNA),^[20] and even uncharged organic molecules (e.g., phenols, boronic acids, saccharides, and H_2O_2)^[21] have also been detected by biphasic sensors. Therefore, the oil-based sensing concept promises to bring a wealth of new analytical capabilities to the droplet microfluidics and advance its applications.

Herein, one additional example of this new chemical sensing approach is highlighted by detection of protamine, an arginine-rich protein with 21 positive charges. The oil phase is 1,2-dichloroethane with chromoionophore I and dinonylnaphthalene sulfonic acid (DNNSH). Selective extraction of protamine over singly charged cations into the oil segments is driven by the strong cooperative ion-pairing interaction between one multiply-charged protamine and multiple singly-charged DNNS anions.^[20] Such extraction competes with protonation of the chromoionophore and induces decreased fluorescence in the oil segments. As shown in Figure 4, this method exhibits a fluorescent response toward protamine over a dynamic range of 1 to 100 $\mu\text{g}/\text{mL}$, which is comparable to other protamine-sensitive optodes, but is achieved here in a much shorter timescale (~ 1.2 s vs. >10 min).^[20b,20c] The strong ion association interaction also makes conventional polyion-selective sensors

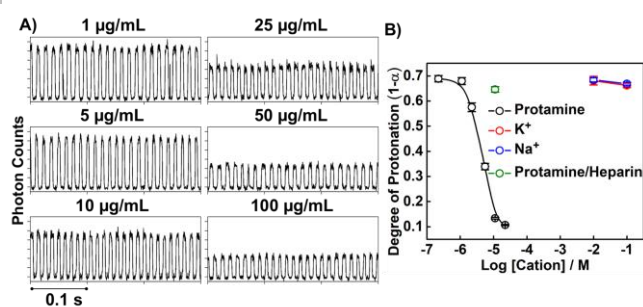


Figure 4. A) Fluorescence trace of the segmented flow in the presence of different concentrations of protamine in 50 mM Tris-HCl buffer at pH 7.4 as the aqueous phase (excitation: 630 nm; emission: 673–738 nm); B) Calibration curves toward protamine and small cations. The green circle is the response toward a 50 µg/mL protamine solution containing 12 U/mL heparin (data points are average \pm SD for $n=3$ measurements).

irreversible.^[20] However, in our sensing scheme, every oil segment is only used for one measurement, and fresh oil segments are generated continuously. Therefore, there is no requirement for reversibility of the biphasic sensing chemistry, which is another distinct advantage of this droplet microfluidic-based sensing scheme. If the protamine is neutralized by polyanions such as heparin, the fluorescence response is prevented because the complexed polycations are no longer able to form ion-pairs with DNNS anions (Figure 4B, green circle). Based on this mechanism, we are currently working on continuous monitoring of blood heparin levels using droplet microfluidics, which would be useful during extracorporeal procedures (e.g., cardiopulmonary bypass surgery) to determine the actual concentration of this anticoagulant in blood. Moreover, this same polyion sensing method has other indirect applications such as enzyme assays, enzyme inhibitor/activator tests, and aptasensing,^[20] which might now be integrated into a droplet microfluidics platform.

In another proof-of-principle example, the chemical sensing of neutral species is demonstrated using droplet microfluidics. The limited aqueous solubility is an obstacle in the application of some BODIPY and azaBODIPY dyes. However, we successfully use a boronic acid-functionalized azaBODIPY dye in the oil phase to detect H₂O₂, an important reactive oxygen species, in the aqueous droplet based on the oxidative conversion of boronic acids to phenols in the oil phase (Figure S5).^[21d]

In summary, the oil phase in droplet microfluidics provides a promising scaffold for sensing of ionic, polyionic, and non-ionic species in aqueous sample droplets. This sensing scheme is fast, reagent-economic, and compatible with complicated sample matrices. The ultimate goal of this endeavor lies in the clinical analysis of chemical species from ultra-small volumes of bodily fluids, as well as the high-throughput screening of drugs that target ion channels on individual whole cells—both of which are not readily achievable by current analytical techniques.

Acknowledgements

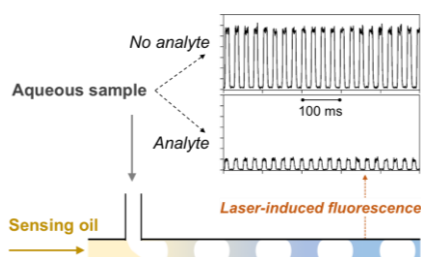
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Keywords: analytical methods • biphasic sensing • ionophores • molecular recognition • droplet microfluidics

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COMMUNICATION

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Xuewei Wang,* Meng Sun, Stephen A. Ferguson, J. Damon Hoff, Yu Qin, Ryan C. Bailey, Mark E. Meyerhoff

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