A microfluidic chemical/biological sensing system based on membrane dissolution and optical absorption

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A microfluidic chemical/biological sensing system based on membrane dissolution and optical absorption

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Abstract
A microfluidic system to sense chemical and biological analytes using membranes dissolvable by the analyte is demonstrated. The scheme to detect the dissolution of the membrane is based on the difference in optical absorption of the membrane and the fluidic sample being assayed. The presence of the analyte in the sample chemically cleaves the membrane and causes the sample to flow into the membrane area. This causes a change in the optical absorption of the path between the light source and detector. A device comprising the microfluidic channels and the membrane is microfabricated using liquid-phase photopolymerization. A light emitting diode (LED) and a detector with an integrated amplifier are positioned and aligned on either side of the device. The state of the membrane is continuously monitored after introducing the sample. The temporal dissolution characteristics of the membrane are extracted in terms of the output voltage of the detector as a function of time. This is used to determine the concentration of the analyte. The absorption spectra of the membrane and fluidic sample are studied to determine the optimal wavelength that provides the maximum difference in absorbance between the membrane and the sample. In this work, the dissolution of a poly(acrylamide) hydrogel membrane in the presence of a reducing agent (dithiothreitol—DTT) is used as a model system. For this system, with 1 M DTT, complete membrane dissolution occurred after 65 min.

Keywords: chemical sensing, biological sensing, optical absorption, dissolvable membrane, microfluidics

(Some figures in this article are in colour only in the electronic version)

1. Introduction
The sensing of chemical and biological events is of critical importance in various applications [1, 2]. Currently, there are many sensing schemes, for instance surface plasmon resonance (SPR) [3–5], chemiluminescence [6–8], ellipsometry [9, 10], impedance spectroscopy [11–13], micromechanosensors [14], nanowire field-effect transistors [15, 16] and capillary zone electrophoresis with UV detection (CZE-UV) [17]. Recently, enzymatic degradation of thin polymer films has been used for biosensing, for instance, using SPR detection [18], and impedance spectroscopy [13]. The SPR technique was capable of detecting analyte concentrations in the range of pM, but required expensive instrumentation. With impedance spectroscopy, at high electrolyte concentration (>2 mM), no changes in impedance were observed upon degradation due to similarity in the impedances of the degradable material and the surrounding electrolyte [18]. CZE-UV-based devices are relatively less portable and less cost effective and require specialized
Dissolvable hydrogels with cleavable crosslinkers have been designed and fabricated in microfluidic channels [19]. Upon exposure to certain chemicals, the crosslinks are cleaved and the hydrogels dissolve in the sample, making them promising as components of microfluidic sensors intended for one-time usage. In our previous work [20], we demonstrated a microfluidic sensing mechanism for bio/chemical analytes using membranes which dissolve in the presence of an analyte. Dissolution of the membrane causes the sample solution to flow into on-chip interdigitated electrodes, bringing about a change in the resistance between the electrodes. The approach gives an indication of the end-point of membrane dissolution, resembling the operation of an on–off switch. With 1 M analyte concentration, it took $\sim 1813$ s for the membrane to become porous and allow electronic detection of the analyte. However, no information regarding the concentration of the analyte can be obtained by using a single membrane.

Here, we describe a sensing system based on optical absorption which provides continuous monitoring of the state of a sensing membrane. Complete dissolution of the membrane is not required to estimate the presence and concentration of the analyte thereby reducing the detection time. Further, the concentration of the analyte can be determined by tracking the temporal dissolution characteristics of a single membrane, simplifying the system design compared to our previous work. Moreover, the time required to estimate the concentration is now shorter compared to the previous approach wherein two membranes with different dissolution times were required, the bottleneck being the membrane with longer dissolution time. A microfluidic device incorporating a bio/chemical recognition membrane is positioned in the optical path between a light emitting diode (LED) and a detector. Presence of the analyte in the sample chemically cleaves the crosslinked membrane and eventually allows the sample to flow into the sample area. The difference between the optical absorbance of the membrane and the sample causes a change in the intensity of the light received by the detector upon dissolution. In this work, we use poly(acrylamide) (PAAm) based hydrogel membranes dissolved by dithiothreitol (DTT, the analyte). The microfabrication process is based on liquid-phase photopolymerization and is compatible with integrated circuit (IC) technology [21, 22].

2. Principle of operation

The sensing of the analyte is accomplished by the dissolution of an analyte-sensitive membrane. The event is detected by tracking the change in optical absorption of the path between a light source and a detector. The schematic diagram of the principle is shown in figure 1.

2.1. Selection of the model system

It has been shown that disulfide crosslinked PAAm hydrogels can be disintegrated in the presence of DTT [19, 20]. Here, we use this as a model for the sensing system. A disulfide crosslinked hydrogel membrane forms the sensing element. The presence of the analyte (here, DTT) breaks the disulfide bonds of the hydrogel, slowly rendering it porous and ultimately dissolves it completely. The dissolution of the hydrogel is a complicated multi-step process [23]. It involves the following steps: (a) diffusion of the disulfide reducing agent into the hydrogel network, (b) reduction of the disulfide crosslinker inside the hydrogel, (c) swelling of the hydrogel during de-crosslinking as a result of decreased crosslink density, (d) disentanglement of the linear (or negligibly crosslinked) PAAm chains after de-crosslinking and, finally, (e) the dissolving of the linear PAAm into the surrounding solution. The dissolution time depends mainly on the physical dimensions of the hydrogel as well as the concentration of the analyte. Higher analyte concentrations break more bonds of the hydrogel in a given interval of time and hence show a quicker response. For example, in our sensor, a 1 M DTT solution gives a stable output in about 70 min and a 0.25 M DTT solution gives a stable output in about 125 min.

2.2. Absorption spectra of the hydrogel membrane and the sample

In order to obtain a significant change in the detector’s output voltage upon dissolution of the membrane, it is necessary to employ a wavelength at which the difference between the absorption of the membrane and that of the sample is a maximum. Hence, we studied the absorption spectra of the hydrogel membrane and the sample before choosing the light source and detector. Figure 2 shows the absorption spectra of the hydrogel membrane, 1 M DTT solution and water, as measured by a spectrometer (USB 2000 Fiber Optic Spectrometer, Ocean Optics Inc., Dunedin, FL, USA). From the spectra, the absorption of both the membrane and the sample at visible wavelengths is almost equal. The difference between the absorbance increases as the wavelength moves towards the UV region ($\leq 400$ nm). It is then found that the difference between the absorbances of the membrane and DTT is maximum at $\sim 340$ nm. Therefore, for the experiments, an ultraviolet (UV) LED and UV photodiode with an integrated amplifier having peak emission and peak sensitivity, respectively, at 340 nm are chosen. Different
concentrations of DTT do not cause appreciable difference in the absorption spectra. Apart from using the spectrometer, this is verified by monitoring the output voltage of the detector after introducing 1 M DTT and 0.25 M DTT sequentially into a device having no membrane. No significant change in the detector’s output voltage is observed. Water is used as a control signal and hence its spectrum is also measured, representing samples that do not contain the analyte.

2.3. Operation of the sensing system

The system performs the temporal measurement of the optical absorption of the membrane in the presence of the sample. Absorption may be characterized by the Beer–Lambert law, \( I_T = I_0 \exp(-\alpha L) \), where \( I_0 \) is the intensity of the incident light, \( I_T \) is the intensity of the transmitted light, \( \alpha \) is the absorption coefficient of the absorbing material (either the membrane or the sample) and \( L \) is the absorption path length. The change in \( I_T \) is attributed to two reasons which are discussed in detail later: (i) diffusion of the sample solution into the membrane and (ii) dissolution of the membrane due to the presence of the analyte in the sample. Before diffusion/dissolution, the membrane is the absorbing material and has a relatively higher absorption (at \( \sim 340 \) nm), as compared to the sample. Upon introduction of the sample, the sample diffuses into the membrane and causes a change in the absorption of the membrane and hence, \( I_T \). Presence of the analyte in the sample dissolves the membrane and allows the sample to flow into the region originally occupied by the membrane resulting in a further change in \( I_T \). The difference in \( I_T \) is used to sense the dissolution of the membrane and hence the presence of the analyte. The sample is guided towards the membrane using microfluidic channels. The LED sources light onto the membrane and the detector provides an output voltage proportional to the intensity of light transmitted through the membrane. As the membrane gets dissolved, the output voltage of the detector increases. The dissolution is complete and the output voltage saturates at a particular value. By tracking the output voltage of the detector, the concentration of the analyte can be determined.

3. Device fabrication and experimental setup

The microfluidic channels and the hydrogel membrane are microfabricated using liquid-phase photopolymerization (LP), a process described in detail elsewhere [24]. The device is positioned between a light emitting diode (LED) and a detector. The output of the detector is monitored on a multimeter and stored on a computer.

3.1. Device fabrication

Figure 3 shows the fabrication process. A pre-cleaned microscope glass slide of dimension 7.5 cm \( \times \) 2.5 cm \( \times \) 1 mm (Fisher Scientific, Pittsburgh, PA) is used as the substrate. First, a rectangular metal pattern is created on the substrate, which serves as a mask for photopatterning the hydrogel membrane, as well as to block light from the LED in regions not having the membrane. The substrate is sputtered sequentially with 0.05 \( \mu \)m titanium (Ti), 0.4 \( \mu \)m copper (Cu) and 0.05 \( \mu \)m Ti metal layers (figure 3(a)). The bottom Ti serves as an adhesion layer for Cu, while the top Ti prevents oxidation of Cu. This step is carried out in a clean room (Wisconsin Center for Applied Microelectronics, UW-Madison) with a CVC-601 dc magnetron sputterer (power: 900 W; current: 203
1.0 A). A rectangular pattern of dimension 200 µm × 3 mm is selectively etched into the metal layers (figure 3(b)). The top Ti is etched using hydrofluoric acid:water = 1:100. Cu is etched using acetic acid:hydrogen peroxide:water = 1:1:10. The Cu etchant does not etch the underlying Ti. The bottom Ti is etched to expose the underlying glass.

Next, the microfluidic channels are fabricated using LP3. The glass slide is flipped vertically, and a 375 µm deep polycarbonate gasket with an adhesive liner is adhered onto the glass. The cavity thus formed between the gasket and the substrate is filled with an iso-bornyl acrylate-based pre-polymer solution. This solution consists of a monomer: isobornyl acrylate, crosslinker: tetraethylene glycol dimethacrylate, photoinitiator: 2, 2-dimethoxy-2-phenylacetophenone in the ratio (by weight) 1.9:0.1:0.06 [21, 22]. The operation of this solution resembles that of a negative photoresist. When exposed to UV radiation, the photoinitiator absorbs the light energy and uses it to bond the monomer with the crosslinker and form a polymer. The regions exposed to ultraviolet (UV) light polymerize and harden whereas the other regions remain liquid and can be flushed away with a solvent. The device is exposed to a UV light source (Acticure 4000, Exfo Life Sciences and Industrial Division, Mississauga, Ontario, Canada) through a film photomask (3600 dpi, Silverline studio, Madison, WI, USA) having the pattern of the microfluidic channels. The photopolymerization conditions are intensity = 7.8 mW cm−2 and exposure time = 22.5 s. The liquid pre-polymer polymerizes in regions exposed to UV forming the channel walls (figure 3(c)) and the unpolymerized pre-polymer is flushed in a bath of ethanol (100%) for ∼3 min. The device is baked on a hotplate at 50 °C for ∼30 min.

The hydrogel membrane is then defined using LP3. The fluidic channels are now filled with a poly(acrylamide)-based pre-hydrogel solution which consists of a monomer—acrylamide; crosslinker—cystaminebisacrylamide; photoinitiator—(4-benzoylbenzyl)trimethyl-ammonium chloride; co-initiator—N-methyl-diethanolamine; a solvent—water in the ratio (by weight) 0.15:0.00 374:0.02:0.02:1 [19, 23]. The device is now flipped vertically so that the metal pattern is on top. The pre-polymer liquid does not drain from the microfluidic channels due to surface tension. The liquid is homogeneous, the other factor which can cause non-homogeneity is the spatial distribution of light intensity during photopolymerization. To compensate for this, the sample is kept on a rotating circular disc during photopolymerization, which spatially averages the dose given to the sample. The hydrogel membrane needs to be homogeneous. Since the pre-hydrogel liquid is homogeneous, the other factor which can cause non-homogeneity is the light intensity during photopolymerization. To compensate for this, the sample is kept on a rotating circular disc during photopolymerization, which spatially averages the dose given to the sample. The hydrogel membrane is self-aligned with the metal pattern and any changes in the detector’s output can be attributed only to changes in the membrane and not the surrounding media. The pre-hydrogel solution polymerizes to form the hydrogel membrane (figure 3(d)). The unpolymerized solution is flushed with ethanol. The device is again baked on a hotplate at 50 °C for 5 min to remove the residual solvent in the hydrogel.

3.2. Experimental setup

Figure 4 shows the experimental setup. The device is coupled to a micropositioner and aligned with a UV-LED light source (UVTOP-340 nm, TO-39 package, flat window, Sensor Electronic Technology, Inc., Columbia, SC, USA) and a UV photodetector with an integrated amplifier (JIC 159 A, Electro Optical Components, Inc., Santa Rosa, CA, USA). The LED is biased at 5.2 V and has a peak emission wavelength of 340 nm and a spectrum half-width of 20 nm. Holes are drilled in a microscope glass slide and the leads of the LED are inserted into the holes and adhered onto the glass slide using double/bubble epoxy (#04004, Elementis Specialties, Inc., Belleville, NJ 07109, USA). The glass slide having the LED is coupled to a micro-positioner (Model S-926, Signatone Corporation, Gilroy, CA, USA) with the light emitting surface of the LED facing the device. The detector’s sensitivity peaks at 340 nm and has a spectral range of 315 nm to 395 nm with an active area of 0.965 mm2. It provides an output voltage proportional to the intensity of the UV radiation incident on its detecting surface. The detector is mounted on a bread board adhered to a lab jack (Fisher Scientific, Pittsburgh, PA, USA) and is kept fixed. An external feedback resistor of 1 MΩ is used to stabilize the gain of the amplifier in the detector. The micropositioners and the lab jack are adhered onto a glass slide (coupled to a micro-positioner) having the channel walls (figure 3(c)). The unpolymerized pre-hydrogel solution polymerizes to form the hydrogel membrane (figure 3(d)). The unpolymerized solution is flushed with ethanol. The device is again baked on a hotplate at 50 °C for 5 min to remove the residual solvent in the hydrogel.
4. Experiments and results

The variation in the intensity of the transmitted light (in terms of the output voltage of the detector) is recorded as a function of time for two different concentrations of DTT—0.25 M and 1 M, and water which is used as a control to show the response of the system when the sample does not contain the analyte. During experiments, the device is aligned between the LED and detector using the micropositioners coupled to the device and LED. The positioners are adjusted so that the detector shows the maximum possible output voltage when the LED is driven with 5.2 V. The sample solution is prepared by dissolving DTT (99%, SigmaAldrich) in a 0.3 M Tris buffer of pH 7.4 (Molecular biology grade, Fisher Scientific). 1 M DTT is made by dissolving 1.54 g of DTT in 10 ml of the 0.3 M Tris buffer. For other concentrations, the weight of DTT to be dissolved in the buffer is calculated proportionately. After positioning the device, the sample solution with appropriate analyte concentration is introduced into the microfluidic channel having the pattern of the hydrogel membrane through filling ports on the polycarbonate cartridge using a pipette. Upon introduction of the sample, the intensity of the transmitted light changes and is attributed to two factors: (i) diffusion of the sample into the hydrogel membrane soon after introducing the sample into the device and (ii) dissolution of the membrane when the sample contains the analyte, as discussed below.

4.1. Diffusion of the sample/water into the hydrogel membrane

Diffusion of the sample into the membrane is one of the factors contributing to the change in the detector’s output voltage. After microfabrication of the hydrogel membrane, the hydrogel is in a dehydrated state and has a relatively high absorption of the incident light as compared to the hydrated state. Figure 5(a) shows an SEM image of the hydrogel membrane. Upon introduction of the sample (or water) into the device, water diffuses into the membrane. Since water has lesser absorption than the hydrogel membrane, there is a change in the output voltage of the detector, as the optical path between the LED and the detector is then occupied partly by the sample and partly by the membrane. Figures 5(b) and (c) shows images of the membrane soon after the introduction of the sample into the system. The membrane is 3 mm long and 200 µm wide. The curve labelled ‘water’ in figure 6 shows the response of the detector when the sample is water (has no DTT) and can be used as a control signal. The diffusion process does not indicate the presence of the analyte, and further information is required. This is done by testing for the dissolution of the membrane.
and saturates.

When the membrane is completely dissolved, the output voltage of the detector ceases to increase, since the absorption of the media between the LED and detector. Upon completion of dissolution, due to a change in the optical absorbance of the path between a light source and a detector. The detector transduces the dissolution event into an output voltage by continuously monitoring the change in the intensity of light transmitted through the membrane. A difference of ∼0.3 V is observed between the undissolved and dissolved states of the membrane. A single membrane can be used to estimate the concentration of the analyte by tracking the dissolution characteristics of the membrane over time. The fabrication process is compatible with conventional integrated circuit technology and the sensing system is relatively simple.

Membrane-dissolution-based sensing allows a range of species to be detected by defining membranes specific to the analyte. It may find applications in the detection of toxins which exhibit protease activity and can be highly sensitive and highly specific. For instance, neurotoxin-specific peptide sequences can be incorporated into the hydrogel network, similar to the approach reported in [25] where hydrogel scaffolds incorporate fibrinogen and polyethylene glycol. Corroborating the output voltage of the detector with the absorption spectra for different conditions such as diffusion into the membrane and its dissolution would be an interesting experiment and will be part of our future work. Presently, the sample is introduced manually into the microfluidic chamber. A sample acquisition system can be added that interfaces the sensor to the outside world and automates its operation. Such a system could leverage the one reported in [26] with integrated microfluidic components such as valves, mixers and pumps realized also using LP3. Improvements in the sensitivity of the membrane can be made by optimizing its chemical composition, physical dimensions as well as the reaction conditions. In our future work, we will extend this sensing system to other biological and chemical analytes.

4.2. Temporal dissolution characteristics of the hydrogel membrane

After the sample diffuses into the membrane, the analyte starts dissolving the hydrogel membrane. This causes the output voltage of the detector to further increase with time and saturate upon completion of dissolution, due to a change in the optical absorption of the media between the LED and detector.

Figures 5(d) and (e) show the images of the membrane before and after dissolution. The images are taken on an opaque substrate so that the dissolution process is visible. The DTT concentration used is 1 M. In figure 5(d), the dissolution process has just started. In figure 5(e), the membrane has been dissolved. The dissolution characteristics are shown in figure 6. Following the introduction of the sample containing the analyte, as time progresses, the disulfide bonds holding the polymer hydrogel membrane are cleaved by the analyte. The sample solution flows into the membrane area and the output of the detector starts to increase, since the absorption of the sample is less than that of the membrane at the chosen wavelength (figure 2). When the membrane is completely dissolved, the output voltage of the detector ceases to increase and saturates.

Two concentrations of the analyte are studied as examples in the experiments—0.25 M and 1 M. Higher concentrations of the analyte cleave a larger number of disulfide bonds per unit time and hence results in quicker dissolution of the membrane. The final output voltage with both concentrations is the same. This shows that the hydrogel membrane is completely dissolved in both cases. A difference of ∼0.3 V is observed between the polymerized and completely dissolved states of the hydrogel membrane. The dissolution characteristics of a membrane, thus the output voltages of the detector, depend on the concentration of the analyte, which can be estimated by tracking the output voltage over time. For example, the dissolution characteristics for a host of analyte concentrations can be obtained and stored. By monitoring the dissolution characteristic of the analyte of unknown concentration with the stored characteristics, a best-fit estimate of the concentration can be made.

Other schemes such as impedance spectroscopy have been studied, where the membrane is patterned on interdigitated electrodes. In such a system, the effective impedance would depend not only on the membrane’s impedance, but also on the analyte concentration. Hence, the impedance for different combinations of membrane thickness and analyte concentration could be the same. This makes the target concentration non-deterministic, since the composition of the sample is unknown. We found that the optical absorptions for different concentrations of the analyte were similar, making the target concentration deterministic.

5. Conclusions

A microfluidic chemical and biological sensing system using membranes dissolvable by an analyte is realized. Dissolution of the membrane by a sample solution containing the analyte causes the sample to flow into the membrane area and changes the optical absorbance of the path between a light source and a detector. The detector transduces the dissolution event into an output voltage by continuously monitoring the change in the intensity of light transmitted through the membrane. A difference of ∼0.3 V is observed between the undissolved and dissolved states of the membrane. A single membrane can be used to estimate the concentration of the analyte by tracking the dissolution characteristics of the membrane over time. The fabrication process is compatible with conventional integrated circuit technology and the sensing system is relatively simple.

Membrane-dissolution-based sensing allows a range of species to be detected by defining membranes specific to the analyte. It may find applications in the detection of toxins which exhibit protease activity and can be highly sensitive and highly specific. For instance, neurotoxin-specific peptide sequences can be incorporated into the hydrogel network, similar to the approach reported in [25] where hydrogel scaffolds incorporate fibrinogen and polyethylene glycol. Corroborating the output voltage of the detector with the absorption spectra for different conditions such as diffusion into the membrane and its dissolution would be an interesting experiment and will be part of our future work. Presently, the sample is introduced manually into the microfluidic chamber. A sample acquisition system can be added that interfaces the sensor to the outside world and automates its operation. Such a system could leverage the one reported in [26] with integrated microfluidic components such as valves, mixers and pumps realized also using LP3. Improvements in the sensitivity of the membrane can be made by optimizing its chemical composition, physical dimensions as well as the reaction conditions. In our future work, we will extend this sensing system to other biological and chemical analytes.
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