Fabrication of nanoporous junctions using off-the-shelf Nafion membrane

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Abstract

This paper reports a method to fabricate nanoporous junctions in microfluidic devices using off-the-shelf Nafion membrane. These nanoporous junctions are needed for various applications based on the phenomenon of ion concentration polarization (ICP). The advantages of this unique method compared to other fabrication methods of cation selective nanofluidic junctions are the simpler fabrication process, the better durability, and the increase in ICP repulsion force. Thus, microfluidic devices fabricated with that method have the potential to enhance the separation efficiency of the ICP process.

Keywords: microfluidics, nanofluidics, nanoporous

(Some figures may appear in colour only in the online journal)

1. Introduction

Ion concentration polarization (ICP) is a fundamental electrochemical transport phenomenon that occurs when ion current passes through an ion-selective membrane. Often called ion depletion or enrichment process [1], this phenomenon happens due to the mismatch of charged carriers at the membrane interface. The membrane (either nanochannel or nanoporous membrane) preferentially or only conducts one sign of ion such as cations in cation exchange membrane. As a result, ion concentration gradients are generated on both sides of the membrane. Once ICP is triggered near the cation exchange membrane interface, the concentrations of both cations and anions decrease on the anodic side of the junction (i.e. ion depletion) and increase on the cathodic side (i.e. ion enrichment). Furthermore, any charged particles, cells and other small colloids will be similarly depleted or enriched in the same analogy.

Recently, ICP has attracted a great deal of attention from the lab-on-a-chip research community because of its potential applications such as sample concentration [2–5], desalination [6, 7], separation and mixing [8, 9]. Further applications of ICP have been reported recently. Jeon et al [10] presented a novel separation method, which can continuously separate the particles based on the ICP phenomenon and the electrophoretic mobility of micro- and nano-sized particles. MacDonald et al [7] reported a method of out-of-plane ion concentration polarization for scalable water desalination, which has the advantages of increasing the throughput and the ability of integrating multiple devices in a single system. Most of the previously reported ICP-based devices used Nafion resin to deposit a thin nanoporous membrane with a thickness varying from 191 nm using microflow patterning method [2] to 200 μm by filling a laser-cut micro channel [7].

The advantage of the fabrication method of depositing a cation-selective junction inside microfluidic devices using Nafion resin as reported by Han’s group [2–6] is the quick formation of a thin Nafion layer on a clean glass slide. However, this method requires an extra polydimethylsiloxane (PDMS) microchannel serving as a master mould and needs several fabrication steps. We observed that this deposition process was not straightforward as expected. The limitations are as follow. The negative pressure required for sucking out the Nafion solution needs to be controlled accurately. The Nafion film does not adhere firmly on the glass slide and can peel off easily. The edges and surface of the fabricated Nafion trip are not smooth. Moreover, the thickness and width of the deposited Nafion trip could not be controlled precisely since the evaporation unpredictably of solution under heating process.

To simplify the fabrication process, we used an off-the-shelf Nafion membrane (NR-212, DuPont Co., USA) with a
2. Methods

We fabricated the microfluidic device for demonstration of a separation process with an integrated cation selective membrane using two different methods, to compare the two fabrication concepts. To create a nanoporous junction, the previously reported desalination device used a Nafion resin to fill a microchannel [6] (thereafter called as resin-made-device). Our proposed method uses an off-the-shelf Nafion strip integrated onto a thin PDMS layer (thereafter called as membrane-made-device).

2.1. Nanojunction based on Nafion resin

Figures 1(a) and (b) show the configuration of the separation device and the layers of the device fabricated by the proposed method, respectively. With this configuration, we first fabricated the devices using the Nafion resin. The fabrication process is summarized as follows. A square glass slide served as the substrate for the device. Gold electrodes with a thickness of 300 nm were sputtered on the glass substrate. The thickness of deposited electrode is a compromise between reducing the electrical resistance and minimizing the gap between the PDMS channel layer and glass substrate for leakage-free bonding. The sputtered electrodes consist of two layers. A 50 nm thick titanium layer was used to enhance the adhesion of the gold layer to the glass substrate. A 250 nm thick gold layer was deposited on top of this adhesion layer. Gold was selected due to its good conductivity and its resistance to oxidation. The Nafion junction was formed by the microflow patterning technique [2]. This technique used a PDMS mould having a microchannel with a width of 100 μm and a height of 10 μm. The mould was temporarily bonded on the glass substrate. The microchannel was then filled with 10 wt% solution of Nafion perfluorinated ion-exchange resin (Sigma Aldrich, St Louis, USA). Negative pressure was applied on the microchannel outlet first to fill it and then to suck out the resin to control the deposited thickness. Subsequently, the sample was baked in an oven at 110 °C for 10 min to cure the deposited Nafion strip. The top layer of the device was made of PDMS and contains the microfluidic channel network fabricated by the standard soft lithography process. Negative photoresist SU-8 (MicroChem Corp., USA) with a thickness of 100 μm was first coated on a silicon wafer. The channel features were defined by exposing the SU-8 layer through a mask to UV light. The PDMS channel was casted on, cured and then peeled off from the SU-8 mould. Finally, the surfaces of the PDMS part and the glass substrate were treated with oxygen plasma (Harrick PDC-32G) for 1 min before bonding together with the help of the precise alignment system described in the later section.

2.2. Nanojunction based on off-the-shelf Nafion membrane

To compare the efficiency of separation process in the devices fabricated with the two different methods, we used the same channel dimension to fabricate another device with

thickness of 50 μm instead of the Nafion resin to deposit a thin cation-selective membrane on a glass substrate. This off-the-shelf membrane provides better mechanical durability and a longer working time. Moreover, a thicker membrane generates a stronger electrical repulsive force around the membrane [4]. In this paper, we report a unique and simple fabrication method to integrate a thin off-the-shelf Nafion strip into a microfluidic device for applications with ICP. We chose the fabrication of a separation device to demonstrate and compare the two fabrication concepts. To create a nanoporous junction, the previously reported desalination device used a Nafion resin to fill a microchannel [6] (thereafter called as resin-made-device). Our proposed method uses an off-the-shelf Nafion strip integrated onto a thin PDMS layer (thereafter called as membrane-made-device).

Figure 1. The separation device: (a) working concept of the device. (b) Integration of a 50 μm thick Nafion membrane on a PDMS/glass substrate. (i) Preparing and cleaning a glass slide. (ii) Coating the PDMS mixture with a spin speed of 1450 rpm. (iii) Cutting the Nafion membrane and placing it carefully on top of the PDMS surface. Placing the sample at room temperature overnight to let PDMS reflow to form a flat surface. Next, placing the sample into the oven at 80 °C for 2 h. (iv) Bonding the top PDMS with microchannel network to the PDMS/glass substrate with the Nafion membrane using a precision alignment system.
our proposed method, as described in details below. Sylgard 184 and a curing agent (Dow Corning Inc.) were mixed in a 10 : 1 ratio by weight. The mixture was degassed for 1 h in a vacuum chamber. Subsequently, 5 ml of the PDMS mixture was poured onto a square glass slide and spin coated in two steps: 500 rpm for 10 s to spread the PDMS over the surface, and then increase to 1450 rpm for 30 s.

Figure 2 shows the measured height of the Nafion membrane relative to the PDMS surface. The results show that the mismatch between the PDMS surface and the Nafion surface is proportional to the spin speed. Tuning the spin speed can match the surface of the Nafion and the surrounding PDMS.

Next, a thin Nafion trip was placed on top of this spin-coated surface and kept in a clean place overnight to allow PDMS to reflow and form a flat surface. To make sure that the thin PDMS film is evenly coated, the substrate can be spin coated again before curing in an oven. The microchannels of the device were fabricated following a general PDMS soft-lithography process. The master mould was made with SU-8 photoresist (MicroChem Corp., USA) and standard photolithography process. After peeling off from the mould, the PDMS part was bonded to the PDMS-glass substrate with the integrated a Nafion membrane using oxygen plasma treatment and a precise alignment system, figures 2(c) and (d).

An aligner was designed and assembled from the Thorlab, Inc. (USA) standard components, to position of the Nafion strip accurately against the concentrated channel, figure 1(a). The alignment process was performed as follows. First, both PDMS top and bottom layers were cleaned by oxygen plasma for 45 s. Then, the PDMS-glass substrate integrated with Nafion strip was quickly placed on the sample holder of the microscope. Concurrently, the top PDMS microchannel layer was attached on the sample holder of the alignment head, figure 2(d). Next, adjusting the four degrees of freedom (three translations and one rotation) brought the Nafion trip to the right position parallel to the concentrated channel. During the alignment process, we could observe the structures to be aligned either via microscope eyepieces or a camera attached to the microscope. From the experimental results, the alignment gap could be achieved in the range from 5 to 10 μm, figure 3(c). Since the active PDMS surface would disappear over time, all steps need to be performed quickly. With our alignment system, the whole process only took 10–15 s.

Figures 3(a) and (b) depict the device design and the fabricated device, respectively. Figure 3(c) shows the position of the Nafion strip relative to other microchannels, which are visualized by food dye in figure 3(d). The Nafion strip (yellow) was integrated with a thin PDMS layer and aligned parallel to the PDMS surface. The alignment gap could be achieved in the range from 5 to 10 μm, figure 3(c). Since the active PDMS surface would disappear over time, all steps need to be performed quickly. With our alignment system, the whole process only took 10–15 s.
to the concentrated channel. Red channels were inlet (A) connected to V\(^+\), concentrated outlet (B) and diluted outlet (C). The blue channel was the buffer solution PBS connected to GND.

### 2.3. Experimental set up

The device was subsequently tested for its functionality. In our experiments, all liquids were kept in 5 ml Hamilton glass syringes, which were driven by syringe pumps (neMESYS, Cetoni, Germany) to provide the pre-set flow rates to the device. A high-speed CMOS camera (Phantom Miro eX4) attached to an inverted epi-fluorescent microscope (Nikon Eclipse TE2000-S) was used to capture the fluorescent images of the channels. The colour images were taken by a DSLR camera (Nikon D700) mounted on the right camera port of the microscope via a specialized adapter combining with an extension tube. A solution of 10µM fluorescein sodium salt C\(_20\)H\(_{10}\)Na\(_2\)O\(_5\) (Sigma Aldrich, St. Louis, USA) in DI water (MiliQ system, Milipore, USA) was used as the inlet sample. We also prepared another sample with a concentration of 500 µM to measure the reference fluorescence intensity value. For the experiments on devices used the off-the-shelf Nafion membrane, the phosphate buffer solution (PBS) with the concentration of 100µM was used to fill the buffer channel for the electric ground. The prepared solution had a pH value of approximately 8 and a conductivity of 0.75 mS cm\(^{-1}\) measured by a portable pH/conductivity meter kit (Thermo Scientific Orion 4-Star Plus, USA). Since most ions are rejected from the diluted outlet and collected at the concentrated outlet, the pH value of the concentrated stream was much higher than that of the diluted stream and the sample flow. For calibration purpose, we also measured the conductivities at various concentrations. The fluorescein sodium salt solutions with concentrations of 100µM and 500µM have corresponding conductivities of 1.9 mS cm\(^{-1}\) and 7.5 mS cm\(^{-1}\), respectively. The electric fields were provided by a high voltage dc power supply (Model PS350, Stanford Research System, Inc.). The fluorescent dye has an excitation wavelength of 540 nm and an emission wavelength of 610 nm. The separation process was observed with the help of a filters set consisted an excitation filter for 540 nm, a dichroic mirror for 565 nm and an emission filter for 605 nm. After recording the images with the high-speed camera, the fluorescence intensity and distance were analysed by using a customized MATLAB program.
3. Results and discussions

3.1. Separation process

We characterized the fluorescence intensity at the area located in front of the Nafion junction to demonstrate the continuous separation. Once the ICP is triggered, all charged ions are repelled away from the Nafion membrane and formed a boundary to prevent any charged species to enter this region. In order to compare the separation performance of the two fabricated devices, the fluorescence intensities have been normalized to the intensity of the 10μM input sample used in the device with off-the-shelf membrane. Figure 4 shows the normalized fluorescent signals at representative locations close to the junction in both devices. Three lines A, B, C were defined at the input stream, preconcentrated plug, and concentrated stream, respectively. The device made of the off-the-shelf membrane demonstrated a significant enhancement in separation efficiency. Figure 4(b) indicates that the device with off-the-shelf membrane had a 50-fold concentration enhancement, which was much higher than the resin-made-device, figure 4(a). Comparing the preconcentrated curves (B) of these two devices shows that the membrane-based device provides a sharper localized boundary at the ICP region, whereas the surrounding charges adjacent to the ICP boundary in resin-based device was not localized. The charged species were dispersed thus the starting part of the fluorescence intensity curve B in resin-based device was brighter than that of the membrane-based device. This might be a result of the stronger repulsion force caused by a higher conductance in the nano-junction fabricated from the Nafion membrane. The material purity and the uniform pore size of the fabricated junction are believed to be the main contribution factors. Moreover, we noticed that the lower intensity in the later part of the preconcentrated curve (B) proved a higher improvement in charged species rejection of the device based on the Nafion membrane. This behaviour may be useful for the separation processes utilizing ICP. Importantly, the separation efficiency would be improved if optimizing the channel geometries, especially at the bifurcated branches. For instance, the concentrated stream (i.e. slanted channel) can be narrowed down to enhance the concentration factor contributed by the geometrical factor, as reported recently on the concentrators utilizing ICP [5, 11].

3.2. ICP boundary displacement

Figure 5 shows the measured distance between the ICP boundary and the edge of the Nafion membrane in the Nafion-resin device (circles) and the off-the-shelf Nafion membrane device (squares), respectively. At first, the distance \( \Delta x \) is almost linear to the applied voltage. Increase the applied voltage did not shift the ICP boundary further as a result of the limiting current behaviour. Obviously, under the same input conditions, the new fabrication technique provided stronger ICP force and thus repels charges and ions stronger, figure 5(b).

Experimentally, we also observed that the ICP boundary fluctuated significantly and may disappear if the applied voltage is too high. These effects could presumably be explained by the breakdown of the PDMS substrate under the threshold voltage. In this situation, short circuit occurred leading to a significant increase in the current, which was monitored continuously by an ammeter (Keithley). In our experiments, the current was limited to 1μA by the dc power supply. Therefore, the applied voltage can only be varied in a limited range. Importantly, with our proposed method, a clear ICP boundary and nice alignment position of Nafion trip can be observed as shown in the left image in figure 5(a).

3.3. Operation time

We also experimentally tested the two types of devices for continuous operation under the same input parameters. We...
fixed the flow rate at 100 μl min⁻¹ and an applied electric field at 100 V cm⁻¹. In this experiment, we intentionally test the devices at high input values to check the lifetime of device. We observed that devices made with Teflon solution could work properly only for 30 min, after that the Nafion junction became ineffective. The edges of the Nafion junction were dissolved slowly. ICP boundary became significantly unstable and disappeared after 1 – 2 min. The malfunction of the junction is probably caused by the lack of structural integrity of the Nafion junction deposited by solution. In contrast, devices made with off-the-shelf Nafion membrane run continuously for more than 45 min without any leakages or removal of the Nafion strip. It demonstrated the integrity of the self-sealing Nafion strip embedded in PDMS layer. Hypothetically, the operation time is considered as depending strongly on the input flow rate and applied electric field. In our test, the device was operated at higher input values of applied electric field and flow rate (100 V cm⁻¹ and 100 μl min⁻¹) than the previous work (75 V cm⁻¹ and 0.1 – 20 μl min⁻¹) [6].

To increase the throughput and lengthen the operation time, especially for the real applications requiring the scale-up or multiplexed configuration such as desalination, the wider channels combined with a wider Nafion strip can be applicable. By optimizing the balance between the channel height, Nafion dimensions, sample flow rate, and applied electric field, the ICP boundary could be positioned properly leading to a significant increase in the operation time of the device.

4. Conclusions

We have demonstrated a robust method to fabricate the microfluidic devices for the applications of ICP phenomenon using an off-the-shelf Nafion membrane. Two types of separation devices were fabricated using Nafion resin and off-the-shelf Nafion membrane to compare their performance. Ease of fabrication, low cost, and reliability are the advantages of the
new fabrication method. The devices can be fabricated thanks to the help of a customized precise alignment system for the bonding process. This system allowed positioning the Nafion trip accurately in place against the microchannel in the top layer. Under the same channel dimension and input parameters, the device based on off-the-shelf Nafion membrane provided a stronger ICP repulsion force than that fabricated with Nafion resin. Thus, the separation efficiency could be enhanced. Moreover, the use of off-the-shelf Nafion membrane leads to better mechanical integrity and a longer lifetime of the device compared to the existing method. The failure of the junction made with Nafion resin may be caused by the poor adhesion to the glass slide. The off-the-shelf Nafion strip provides a better mechanical durability because it is strongly embedded into the PDMS-glass substrate by the self-sealing reflow process of PDMS. Since our device worked under the combination of a hydrodynamic force and an electric field, a Nafion membrane would be more suitable for high flow rates and high applied voltages than the thin Nafion junction formed by resin. Our fabrication method is especially useful for continuous concentration [11] or separation process, whereas larger charges and ions need to be filtered out to one outlet. The fabrication method can also be applied to mixing applications required strong electroconvection under ICP zone. The current limitation of this proposed method is the manual placing of Nafion strip on the spin coated PDMS-glass substrate. This issue can be improved by using a customized and automated aligner to scale up the current approach.

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