SU-8 and PDMS Based Hybrid Fabrication Technology for Combination of Permanently Bonded Flexible and Rigid Features on a Single Device

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Abstract

In this article, a novel hybrid fabrication technology is presented that uses both a flexible polymer (polydimethylsiloxane - PDMS) and a rigid polymer (SU-8). A covalent bond between the flexible and rigid polymer layers is achieved using an oxygen plasma treatment during a layer-by-layer direct spin-on process. Precise alignment of the features in each layer and a highly repeatable method are achieved by this new process.

As a proof-of-concept, we successfully fabricated PDMS-based flexible microfluidic devices with SU-8-based rigid world-to-chip/chip-to-world interconnects. The bond strength between the PDMS and SU-8 layers is measured by three methods: 1) Instron® microtester to pull apart the layers; 2) voice coil actuator to test the bond between interconnects and the substrate; and 3) microfluidic pressurization test to evaluate the bond strength along the channels. The bond strength between the flexible PDMS layer and the rigid SU-8 features is very strong; the bond between these two polymers does not fail during these evaluations although the integrity of the PDMS layer itself fails during the microtester evaluation. Additionally, the layer-by-layer direct-spin on process resulted in a repeatable process and precise alignment of the features in each layer, which are necessary in order to achieve consistent performance from the fabricated devices.

The rigid SU-8 interconnects fabricated onto a flexible PDMS device serve as a world-to-chip/chip-to-world interconnects for the direct connection with Tygon® tubing. Three different designs of hybrid (PDMS and SU-8 based) microfluidic devices are designed, fabricated and tested. Each variation differed in the microchannel design in order to demonstrate the versatility of the process to make devices on multiple scales and patterns. These hybrid microfluidic devices are capable of functioning without leakage up to pressures of 85.85 ±3.56 kPa. Although microfluidic channels with interconnects are shown as a proof-of-concept, the fabrication process demonstrated herein could be utilized to develop a number of more sophisticated microfluidic and biomedical devices.
1 Introduction & background

SU-8 and PDMS have been widely used for many years as the polymers of choice in the fabrication of a number of different polymer-based microfabrication processes. Initially, PDMS and SU-8 were mainly used to fabricate devices with microfluidic or biomedical applications [1-3] and SU-8 was additionally used to make molds for PDMS based flexible devices [1-3]. SU-8 has also been used in the fabrication of active devices, such as gears, and actuators [4]. PDMS is a desirable material in the fabrication of microfluidics because of its proven bio-compatibility [5], over-all flexibility and relatively low materials cost.

Generally, PDMS is used for fabricating flexible devices and SU-8 is used for fabricating rigid and strong micro-structures. Until now, systems that include a combination of rigid and flexible polymers have been realized using PDMS and other rigid substrate materials, such as glass, PMMA and silicon [6-8]. In such micro-systems, the PDMS based flexible features and the other rigid features were fabricated separately and combined in the end by surface treatment of the PDMS followed by bonding at the interface of these dissimilar materials. Besides, rigid layers are generally used as a base substrate upon which the devices are built, and primarily used for metal connections or structural rigidity. Hence, the over-all flexibility of the completed device is lost in such cases. Additionally, the existing approaches have limitations in the precision in the alignment of bonded features.

In many cases, it is desirable to fabricate rigid structures directly on flexible devices without sacrificing over-all flexibility of the resulting device. An example of a potential application of these hybrid structures are rigid microfluidic interconnects integrated directly onto a flexible microfluidic chip. These rigid interconnects can be used for chip-to-chip and/or chip-to-world/world-to-chip interconnects. It has been shown that combinations of rigid and flexible materials usually result in the most robust interconnects [9].

Previously, microfluidic chips with interconnect structures were fabricated either using a flexible polymer material (e.g., PDMS) or rigid polymer materials (e.g., SU-8) [9]. However, in these previous designs, the connections between the device and the “world” were fabricated using the same material as the microfluidic chips (i.e. SU-8 interconnects onto the SU-8 microfluidic chips, and PDMS interconnects onto the PDMS microfluidic chips). SU-8 based microfluidic chips with SU-8 interconnects worked well for chip-to-chip and/or world-to-chip/chip-to-world connections. However, the PDMS interconnects based chips were difficult to align and connect together because of the inherent characteristics of the PDMS, such as stiction between surfaces and structural deformation.

Building rigid interconnects on a flexible device would make these structures more functional and adaptable to a wide variety of applications. Hence, a fabrication technology that can incorporate the rigid (e.g., SU-8) interconnect structures directly onto the flexible (e.g., PDMS) microfluidic device is required for such devices. Fabrication procedures need to be
developed that can incorporate rigid interconnect structures directly onto the PDMS layer. Additionally, to the best of our knowledge there is no direct spin-on and self-registration process demonstrated in the literature for fabricating rigid polymer structures on top of any other flexible polymer support.

We present here a fabrication method to fabricate SU-8-based microstructures directly onto flexible PDMS based devices. This fabrication process features a layer-by-layer direct spin-on process to prepare an SU-8 layer on top of a PDMS support. Using this new approach, we are able to align multiple layers of PDMS and/or SU-8 that may be useful in more complex designs and applications. The proposed fabrication technology also incorporates a strong adhesion between the PDMS and the SU-8 layers. Using the hybrid fabrication technology, fully flexible microfluidic devices (PDMS-based) are fabricated with the rigid (SU-8) interconnects directly patterned onto the fluidic inlet/outlet ports. The devices presented here are merely for a proof-of-concept, but the capabilities of this technology are not limited to this simple demonstration. Many more complex biomedical and microfluidic devices could be possible using the proposed fabrication technology.

2 Fabrication principle and hypothesis

In this section, a detailed explanation is provided to describe the hybrid fabrication technology to make flexible devices using both the SU-8 and PDMS polymers. First, the hybrid fabrication principle is explained using the simplified process of the proposed technology. And finally, different designs of the hybrid microfluidic devices are described based on the fabrication technology.

2.1 Fabrication principles

In this section, a simplified set of fabrication steps are briefly described for our new fabrication process. The fabrication process for preparing the SU-8 molds for the patterning of PDMS microfluidic devices to be used as the patterned base layer of our device (PDMS) are not described in detail here. These are relatively well known technologies [10]. Details are provided for the fabrication process to make hybrid microfluidic devices (see Section 2.4). The cured, patterned PDMS layer served as the base layer for demonstrating our new approach to making hybrid devices. The exact process parameters for patterning the SU-8 layer(s) were adopted from previous experiments [11][12], and are not elaborated upon in further detail within this manuscript. We do, however, explain the important process steps and considerations necessary to achieve a permanently bonded SU-8 layer onto the PDMS layer by direct spin-on process.

The simplified fabrication process of the proposed hybrid fabrication technology is shown in Figure 1. In the first step of this fabrication process, the PDMS layer was prepared on a clean glass substrate using standard soft lithography techniques (Figure 1(a)). This PDMS layer was exposed to the oxygen (O₂) plasma (using an Axic Benchmark 800 Plasma System) to
activate the surfaces of the PDMS (Figure 1(b)). An appropriate formulation of the SU-8 layer was spun (using Headway Research Inc. spin coater) onto the PDMS layer directly after the preparation of the activated PDMS surfaces (Figure 1(c)). It is very important to coat the SU-8 layer onto the activated PDMS as soon as possible in order to form the strongest bond between these two polymer layers (PDMS and SU-8). A detailed explanation for a possible mechanism behind the strong interactions between the SU-8 and PDMS are discussed in Section 2.2. A specific formulation of the SU-8 and spin coating parameters were used to achieve the desired thickness of the SU-8 layer as described in prior literature [11][12]. The SU-8 layer was soft baked on a hot plate (Torrey Pines Scientific EchoTherm HS40) followed by exposure to UV light (using Quintel Corporation Q-2001CT) through an appropriate photolithography mask (Figure 1(d)). The UV exposed SU-8 layer was heated again for the post-exposure bake process (Figure 1(e)), cooled to room temperature and immersed in an SU-8 developer bath composed of propylene-glycol-monoether-acetate (PGMEA) (Figure 1(f)). This SU-8 developer dissolved the uncrosslinked SU-8. Finally, the PDMS layer and the permanently bonded SU-8 structures were peeled-off from the glass substrate (Figure 1(g)).

The resulting hybrid, free-standing polymer stack contained a PDMS base layer and patterned SU-8, which was permanently bonded to the PDMS to provide a combination of

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Figure 1: The simplified hybrid fabrication process: (a) On a cleaned substrate, pour and cure a mixture of the Sylgard 184 (PDMS) elastomer and its curing agent; (b) expose the PDMS layer to an O₂ plasma to activate the surface; (c) spin the appropriate SU-8 formulation for the desired thickness of SU-8 and soft-bake this layer with the parameters defined previously [11]; (d) selectively expose the SU-8 layer to a UV source using the desired mask pattern; (e) heat the SU-8 layer for the post exposure bake; (f) remove the uncrosslinked SU-8 using the SU-8 developer; and (g) peel-off the hybrid polymer stack from the base substrate.
flexibility and rigidity combination with precise alignment of the features in both layers.

### 2.2 Hypothesis behind the hybrid fabrication technology

Previously, the SU-8 structures were directly prepared on a PDMS layer to obtain easily peel-able SU-8 devices [11] or SU-8 based sacrificial layers [12]. These stand-alone SU-8 structures were easily fabricated because the adhesion of the SU-8 layer with the PDMS layer is inherently poor. Additionally, the SU-8 is a mechanically rigid material whereas the PDMS is a mechanically flexible material, which helps to detach the fully processed SU-8 layer just by mechanical bending the PDMS layer [11][12].

We introduce herein a hybrid polymer stack incorporating both the PDMS and SU-8 layers with a strong bond between these two polymer layers. To achieve a strong adhesion between these two materials, the PDMS surfaces were treated with oxygen plasma before the SU-8 layer was spun onto the PDMS. In the previous technologies [11][12], the PDMS surface was not treated by any means before processing the SU-8 layer on the surfaces of the PDMS. Hence, the additional step of oxygen plasma treatment created a strong bond between the PDMS and SU-8 layers. All the other SU-8 processing steps, such as the soft bake, the post-exposure bake, the UV exposure, the solvent development and the hard bake remained identical in both methodologies.

The cured PDMS contains repeating chains of dimethylsiloxane units (-O-Si(CH)₃₂-)ₙ. Exposing these polymer chains to oxygen plasma creates silanol groups (Si-OH) on the PDMS surfaces [13][14]. It has previously been demonstrated that the epoxy groups of SU-8 can strongly bond with the (Si-O) groups of silanol on the exterior of plasma treated PDMS surfaces (or SiO₂) [15]. The SU-8 polymer chain can, therefore, strongly bond with the functional groups generated by the oxygen plasma treatment of the PDMS surfaces.

In order to demonstrate a successful proof-of-concept, we used surface activation parameters that were optimized in prior experiments [13]. In future, this study could be further extended to determine the influence of different activation parameters on quality of the hybrid flexible structures created by processes described herein.

### 3 Design and fabrication of the hybrid microfluidic devices

After optimizing the SU-8 and PDMS based technology, we designed and made a proof-of-concept device to demonstrate the utility of this method to fabricate hybrid flexible devices. Three different designs were prepared for demonstrating a hybrid microfluidic device. The following sections describe in detail the process flow to design and make these hybrid microfluidic device.
3.1 Designs of the hybrid microfluidic devices

As a proof-of-concept for the hybrid fabrication process, we prepared hybrid microfluidic devices using the combination of flexible PDMS microfluidic channels and rigid SU-8 interconnects made directly on the PDMS layer. Three different designs were prepared for this demonstration. The first device we designed and fabricated was a five channel microfluidic device (Figure 2(a)); the second was a 'Y-channel' mixer (Figure 2(b)); and the third a '+' channel' dielectrophoresis (DEP) device (Figure 2(c)).

As shown in Figure 2, all three devices were designed using three flexible layers of PDMS and one rigid layer of SU-8 on top of all of the PDMS layers. The top two PDMS layers

![Figure 2](image-url)
and the layer of SU-8 interconnects were prepared on the same substrate using a layer-by-layer direct spin-on process to form the hybrid polymer stack. The bottom PDMS layer was prepared on a separate substrate and bonded with the hybrid polymer stack using the oxygen plasma to activate this interface [13][14].

The multilayer SU-8 mold for the PDMS channel (middle) layer and the PDMS fluidic port (top) layer was prepared on a base substrate. Subsequently, the PDMS mixture was poured and cured. Finally, the rigid SU-8 interconnect layer was directly processed onto the PDMS layer when the PDMS layer was still on the mold. By using this layer-by-layer direct spin-on approach, we were able to achieve accurate alignment of the features using a highly repeatable fabrication process.

All the hybrid microfluidic devices had a base that was 10-mm wide and 22-mm long. The first device (Figure 2(a)) consisted of different sized microfluidic channels, which were used in the microfluidic pressurization test. The five different widths of these microchannels were 50 μm, 100 μm, 200 μm, 300 μm and 500 μm. All these channels were 20 mm long. The second device (Figure 2(b)) was a 'Y channel' microfluidic device generally used as a simple mixer of two different types of fluids. Both the inlet branches were 200 μm wide and the mixer channel was 400 μm wide. The third device (Figure 2(c)) was the ‘+ channel’ device, which is often used for dielectrophoresis (DEP) experiments. All the channels were 400 μm wide in this DEP device.

The bottom PDMS layer was 500-μm thick, the microchannel layer was 100-μm thick and the layer supporting the microfluidic ports was also 100-μm thick. The microfluidic ports were 500 μm in diameter for all the devices. The SU-8 interconnect layers were 250-μm high with an inner diameter of 600 μm and an outer diameter of 1000 μm.

3.2 Detailed fabrication flow of the hybrid microfluidic device

In this section, the detailed flow is described for the fabrication process to create the dielectrophoresis device. The same fabrication process flow, although using different photolithography masks, was used to fabricate other two devices. In the end, photolithography masks were also prepared that accommodated all the designs on a same substrate. As mentioned earlier, the top two PDMS layers along with the SU-8 interconnect layer were processed on the same substrate using the layer-by-layer self-registration process to obtain a reproducible alignment between the processed layers. The procedures for the fabrication process to assemble the hybrid microfluidic device are outlined in Figure 3.
To reliably process the SU-8 layers with different thicknesses, accurate process parameters are essential. 

**Figure 3:** The Step-by-step fabrication process for creating the hybrid microfluidic devices. 

(a) On a cleaned glass substrate, a 100-μm thick layer of SU-8 was spun and soft baked. 
(b) This soft baked SU-8 layer was selectively exposed to UV irradiation through a photolithography mask designed for the mold of a microchannel. 
(c) A second layer of 100-μm thick SU-8 was directly spun onto the first layer of SU-8 and both layers were baked using the appropriate parameters for soft bake of the second SU-8 layer. 
(d) Expose the second layer of SU-8 to UV irradiation through a photolithography mask to define the inlet and outlet ports, followed by heating for the post-exposure bake. 
(e) The uncrosslinked SU-8 was removed by rinsing these stacked layers in SU-8 developer, and these multiple layers of stacked SU-8 structures were hard baked. 
(f) The PDMS mixture was prepared using Sylgard 184® elastomer and the appropriate curing agent kit. A mixture of both components was poured over the multilayer SU-8 mold and cured at an elevated temperature. 
(g) The PDMS surfaces were plasma treated and a 250-μm thick SU-8 layer was spun coat directly onto these surfaces to make the rigid interconnect structures. 
(h) After soft baking this SU-8 layer, it was exposed to UV irradiation through a photolithography mask appropriate for patterning the interconnects. This assembly was heated for the post-exposure bake, and (i) solvent developed to dissolve the unexposed (uncrosslinked) SU-8. 
(j) The hybrid polymer stack was peeled away from the supporting substrate and bonded with the bottom PDMS piece by plasma activation.
parameters for each fabrication step should be optimized. For the SU-8 fabrication process, these steps consists of spin, softbake, UV exposure, post exposure bake, development and hard bake. Similarly, for PDMS, these processing steps consist of mixing elastomer and curing agent, degassing, pouring on to mold and curing. The parameters for the fabrication of the SU-8 and PDMS layers used in these experiments were previously determined and reported in the literature [11][12].

Using a clean glass substrate, a 100-μm thick SU-8 layer was spun for the microchannels mold (Figure 3(a)). The first SU-8 layer was heated on a hot plate and exposed to UV irradiation through the photolithography mask for the appropriate microchannels (Figure 3(b)). A second layer of 100-μm thick SU-8 was spun onto the first layer of UV exposed SU-8 (Figure 3(c)). The resulting stack of SU-8 was heated on a hot plate using the soft bake parameters of the second SU-8 layer. Both of the SU-8 layers were subsequently exposed using the photolithographic mask for microfluidic ports (Figure 3(d)). Subsequently, both the SU-8 layers were heat treated on a hot plate for the post-exposure bake and rinsed extensively using the SU-8 developer (Figure 3(e)). The solvent developed stack of SU-8 layers was then rinsed with a fresh portion of the SU-8 developer and followed with a rinse of isopropyl alcohol (IPA). The resulting multilayer SU-8 molds were hard baked for the structural stability. These SU-8 molds were subsequently used to make tri-layer structure of PDMS/PDMS/SU-8.

The PDMS mixture was prepared by mixing the Sylgard 184® elastomer and the curing agent in 10:1 (w/w) ratio. The de-gassed mixture was poured onto the multilayer structure of SU-8 and any excess amount of the uncured PDMS was scraped off from these molds. The removal of the excess PDMS was an important step to achieve PDMS layers of the desired thickness and upon which were created the microfluidic inlet and outlet ports. Additionally, the scraping action also ensures that PDMS was removed from the tops of the mold for the microfluidic ports and, hence, there remained clear openings to these ports after removal of the PDMS from these SU-8 molds. The PDMS mixture cast against this mold was cured at an elevated temperature (Figure 3(f)).

Now, the multilayer SU-8 molds and the PDMS stack is exposed with the oxygen plasma. The polymer stack (the SU-8 molds and the PDMS layer) prepared on the glass substrate was treated with the oxygen plasma. Right after the glass substrate with the polymer stack was removed from the surface treatment, a 250-μm thick SU-8 layer was spun onto the PDMS layer to create the rigid interconnect structures (Figure 3(g)). The SU-8 layer was soft baked and exposed to UV irradiation through the photolithography mask for creating the pattern of the interconnect structures (Figure 3(h)). The UV exposed SU-8 layer was then heated for the post-exposure bake and solvent developed using the SU-8 developer (Figure 3(i)).

The processed hybrid polymer stack of PDMS layers and the SU-8 interconnects were gently peeled away from their molds. The bottom layer of PDMS within this hybrid stack was bonded with a flat layer of PDMS, which was prepared separately, to seal the channels of the
hybrid microfluidic device (Figure 3(j)). The resulting hybrid microfluidic device was flexible and included permanently bonded rigid SU-8 interconnects for chip-to-chip and/or world-to-chip/chip-to-world interconnections.

The completed hybrid microfluidic devices were visually inspected and mechanically tested for functionality and performance as discussed in the following section.

4 Experimental results & discussions

The fabrication process to make hybrid polymer-based flexible devices, including hybrid microfluidic devices, was tested for different aspects during and after each of the fabrication steps. An especially important component was the bond strength between the PDMS layer and the SU-8 layer (the interconnects), which we tested many times during the technology development. The completed PDMS and SU-8 based hybrid microfluidic device was also tested with the microfluidic pressurization test to determine functionality of the interconnects for chip-to-chip and/or world-to-chip/chip-to-world connections. Each experimental test along with the results obtained from these tests are discussed in the following sub-sections.

4.1 Bond strength measurement between the SU-8 and the PDMS layers

Before designing and fabricating applications using the hybrid fabrication technology, such as the hybrid microfluidic devices, it was very important to evaluate the bond strength between the PDMS and SU-8 layers produced by this hybrid fabrication process. To determine the bond strength using the Instron® microtester, a simple rectangular SU-8 slab was made on top of a flat piece of PDMS (Figure 4(a)).

![Figure 4: The design and test set-up for measuring the bond strength between the SU-8 and PDMS layers. (a) The design for the bond strength test. A 100-μm thick SU-8 layer was fabricated on top of a 1-mm thick PDMS layer. Both of the polymer layers were processed to obtain a permanently bonded section of the SU-8 to a portion of the PDMS surface and the other half of the SU-8 is peelable as this portion of the PDMS was blocked during the plasma treatment. (b) The test set-up using the Instron® Microtester to determine the bond strength. The peeled polymer layers were attached to the tester arm and mechanically pulled apart to determine the bond strength.](image-url)
The Instron® microtester is a mechanical tester capable of very accurate displacement and includes a PC-based measurement system. The test samples were attached to the top and the bottom arms of the tester and mechanically pulled apart while measuring the respective force. To be accommodated in this test set-up, the PDMS and the SU-8 based test structures were fabricated specifically to attach securely into the grippers of the top and the bottom arms (Figure 4(a)). To obtain such structures, a portion of the PDMS layer was covered with a piece of glass during the surface treatment. Hence, one-half of the subsequently processed SU-8 layer was permanently bonded to the PDMS layer and the other half of the SU-8 layer was easily peel-able from the PDMS layer (this technology is explained in [11][12]).

After fabrication of the test structures, the SU-8 layer from the untreated PDMS area was peeled-off and securely attached into the mechanical tester (Figure 4(b)). The PDMS layer was attached to the fixed (bottom) arm of the microtester and the peeled SU-8 layer was attached to the movable (top) arm of the microtester. Both the arms of the microtester were moved using the defined linear displacement and the respective force was measured during this process.

After more than 20 measurements of different samples, it was clear that the PDMS layer fails and ruptures every time without damaging the bond between the PDMS and the SU-8 layers (Figure 5). The bond between the two polymers was intact in all the samples at the end of the measurement. It is clear that the bond between the two polymers is strong and it isn't possible to separate the polymer layers.

4.2 Fabrication related results

During the fabrication of the PDMS and SU-8 based hybrid microfluidic device, optical microscopy images were captured using the optical microscope (Olympus MX40) to check acceptability of the processed features after each step. Some of the important optical images are included here (Figure 6 to Figure 9).
In Figure 6, a high magnification image is shown for the multilayer SU-8 molds. The final components within the mold were precisely aligned and clearly defined after the hard bake step because of the layer-by-layer direct spin-on process. The molds in the image also adhered very well to the glass substrate. A very uniform and consistent fabrication process was achieved because of the precisely tuned parameters for a wide variety of the substrate materials [11][12].

Figures 7, 8 and 9 show the hybrid polymer stacks after the fabrication process was completed on the substrate and while the stacked structure was still on the glass substrate. The precisely aligned and permanently bonded SU-8 interconnects are clearly visible in all three images. The accuracy of the alignment because of the layer-by-layer direct spin-on process is
clearly visible from the high magnification image of Figure 9. Additionally, consistency of the processed layers throughout the substrate was excellent even after multilayer hybrid fabrication process (Figure 7).

The optical microscope images clearly showed that the hybrid polymer fabrication process using multilayer processing helps to produce uniform, well-defined, and reproducible features. Additionally, the self-registration process for the multi-layer fabrication was able to provide high accuracy in the layer-to-layer alignment.

4.3 Interconnect adhesion test

After the polymer-based hybrid microfluidic device was fabricated, the bond strength between the SU-8 interconnects and the top PDMS layer was once again tested using the voice coil actuator (VCA) as described in [11]. The graphical presentation of the test set-up is shown in Figure 10.

In the VCA-based test set-up (Figure 10), a needle was attached to the VCA and the pointed end of the needle was aligned to the SU-8 interconnect of the hybrid microfluidic device. Force applied to the interconnects were in an attempt to use the VCA to detach these SU-8 structures from the top PDMS layer (the PDMS port layer). The force generated through the VCA is linearly proportional to the current supplied to the VCA [11].

Using the VCA-based adhesion test, it was not possible to detach the SU-8 interconnects from the PDMS layer. During all the tests, the SU-8 interconnects deformed the PDMS substrate and moved away from the needle point. The acquired analytic data was, therefore, due to the strong adhesion between the PDMS layer and the SU-8 interconnect.

![Figure 10: The graphical presentation of the test set-up for the interconnect adhesion test using the voice coil actuator (VCA). The fabricated hybrid microfluidic device was placed on a glass substrate and the needle attached to the VCA was used to push the SU-8 interconnects, trying to remove them from the PDMS layer. In all the tests, however, the SU-8 interconnect slipped from the needle because of the flexibility of the PDMS layers.](image-url)
4.4 Microfluidic test

After complete fabrication of the hybrid microfluidic devices, a microfluidic test was performed on each of the devices to test their desired functionality. It was also important to verify reliable completion of the process and absence of leakage in each of the devices. One of the microscopic images of the microfluidic functionality test is shown here (Figure 11). The 'Y-channel' device (Figure 2(c)) was injected with two separate solutions of de-ionized (DI) water each containing a different dye molecule with a unique color (Figure 11).

During the microfluidic testing the devices were directly connected to the syringe pump with Tygon® tubing placed over the rigid SU-8 interconnects (Figure 11). The functional test of these devices was performed using a flow rate of 0.05 mL/min created using a syringe pump (Harvard Apparatus '11'). As shown in Figure 11, there was no leakage in these hybrid microfluidic devices around interconnections, as well as from any other layer within these devices. The lack of leaks was partially attributed to the design of each interconnect. The outer diameter of each interconnect was slightly larger than the inner diameter of the Tygon® tubing (importance of this was explored in detail within [16]).

Besides the microfluidic functional test of the hybrid microfluidic devices, we also performed a microfluidic pressurization test of the devices. For the microfluidic pressurization test, a microfluidic device was used that contained different channel widths (Figure 2(a)). The microfluidic test set-up was prepared using the syringe pump and a pressure sensor (Omega PX26-005GV and Omega PX26-030GV) [11]. All the pressurization experiments were
performed at a flow rate from 0.01 mL/min to 0.10 mL/min on three separate devices. The determination of the leakage pressure was performed using the 50-µm wide channels with a flow rate from 0.01mL/min until leakage of the pumping fluid was observed under the optical microscope. The simulation results obtained from COMSOL are used as a base line to select the flow rates for each pressurization test. The simulation results along with the experimental measurements from three different devices are plotted in Figure 12.

As shown in Figure 12, the microfluidic pressurization results are plotted in two parts. The upper half of the plot represents the pressurization test results obtained from the simulation and from the experimental measurements for the 100-µm, 200-µm, 300-µm and 500-µm wide channels. As mentioned above, these tests were performed from flow rates of 0.01 mL/min to
0.10 mL/min. Similarly, the bottom half of the plot represents the pressurization test results and simulation associated with the 50-μm wide channels. As shown in Figure 12, the pressure in all the microfluidic channels increased linearly with the flow rate. Additionally, none of the channels leaked at flow rates up to 0.10 mL/min in all three devices. Another important aspect for avoiding leakage and repeatable pressure responses was consistency of the feature size, which is also visible from the plots in Figure 12. The repeatable results from the different devices indicated repeatable alignment of the features in all the experiments.

Similarly, for the leakage pressure measurements using the 50-μm wide channels, channels were used from three different devices. In all three devices, the leakage between the Tygon® tubing and the rigid SU-8 interconnect is observed when the flow rate is increased to 0.2 mL/min. The average pressure at which these three devices failed was 85.85 ± 3.56 kPa. Errors were calculated from the standard deviation of three independent measurements.

The hybrid fabrication technology developed using the PDMS and the SU-8 materials was very useful for fabricating hybrid microfluidic devices that can be modularly used for many biomedical and microfluidic applications. The SU-8 interconnects provided easy chip-to-chip and/or world-to-chip/chip-to-world interconnections without requiring any gaskets or special set-ups. This modularity is especially useful for the biomedical devices that require polymer-based disposable sub-systems.

5 Conclusions

A hybrid fabrication technology using the flexible polymer (PDMS) and the rigid polymer (SU-8) is successfully presented in this article. A permanent bond between the flexible and rigid polymer layers is achieved using oxygen plasma treatment of the PDMS followed by a direct spin-on process to create the SU-8 features. Additionally, very accurate alignment and a reliable process are achieved because of the layer-by-layer processing and the direct spin-on SU-8 process on top of the PDMS layer.

To prove the usability of the hybrid fabrication technology, we developed hybrid microfluidic devices with rigid interconnects. The fabrication technology is, however, not limited to only this application, it can be utilized to develop many sophisticated microfluidic and biomedical applications using these low cost polymers. Three different designs of the hybrid microfluidic device were fabricated and tested.

From the adhesion tests between the two polymer layers, we conclude that the bond strength between the two polymer layers is extremely strong. Because of such a strong bond, the integrity of the PDMS material failed while the bond between the two polymer layers remained intact during all experiments. This proof of permanent bond was also tested using different test set-ups at different stages of the technology development.
The microfluidic test performed on the hybrid microfluidic devices proved modularity of the rigid SU-8 interconnects for the world-to-chip/chip-to-world interconnections. The microfluidic pressurization test performed on all the devices demonstrated a linear response to pressure with respect to flow rates from 0.01 mL/min to 0.10 mL/min. The experimental test response obtained from three different devices also demonstrated the precise alignment achieved in this process and the repeatable process yield. The 50-μm wide channels of the hybrid microfluidic devices leaked upon reaching a flow rate of 0.20 mL/min and a pressure of 85.85 ± 3.56 kPa.

6 References


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