Ultradeep fused silica glass etching with an HF-resistant photosensitive resist for optical imaging applications

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Abstract

Microfluidic and optical sensing platforms are commonly fabricated in glass and fused silica (quartz) because of their optical transparency and chemical inertness. Hydrofluoric acid (HF) solutions are the etching media of choice for deep etching into silicon dioxide substrates, but processing schemes become complicated and expensive for etching times greater than 1 h due to the aggressiveness of HF migration through most masking materials. We present here etching into fused silica more than 600 μm deep while keeping the substrate free of pits and maintaining a polished etched surface suitable for biological imaging. We utilize an HF-resistant photosensitive resist (HFPR) which is not attacked in 49% HF solution. Etching characteristics are compared for substrates masked with the HFPR alone and the HFPR patterned on top of Cr/Au and polysilicon masks. We used this etching process to fabricate suspended fused silica membranes, 8–16 μm thick, and show that imaging through the membranes does not negatively affect image quality of fluorescence microscopy of biological tissue. Finally, we realize small through-pore arrays in the suspended membranes. Such devices will have applications in planar electrophysiology platforms, especially where optical imaging is required.

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

Introduction

Glass and fused silica are appealing materials for constructing microelectromechanical systems (MEMS), lab-on-a-chip and microfluidic platforms due to their chemical inertness, biocompatibility, optical transparency, mechanical rigidity, high melting point, electrical insulation, gas impermeability and ability to bond to silicon, glass and polydimethylsiloxane (PDMS) [1–3]. However, many of the wafer-scale processing methods developed for silicon are not readily transferable to glass; hence, serial fabrication techniques have been employed, such as ion track etching through glass [4] and laser machining fused silica [5] and Foturan photostructurable glass ceramics [6]. These methods have been used to realize high aspect ratio microfluidic devices [7] and planar patch-clamp electrodes [4] in glass materials. To avoid the use of specialized equipment, there has recently been an effort to adapt wafer-scale processing methods to glass, namely reactive ion etching and lithographically defined ‘wet’ etching. These methods have enabled the realization of a variety of devices including free-standing air cavities [2], micropumps [8], capillary electrophoresis microchambers [1, 9], high Q-factor resonators [10], microfluidic channels [11, 12], waveguides [13], bioanalytical devices [14] and single cell trapping wells [3], planar patch-clamp electrodes [15], and optical sensing platforms [1, 3].

Reactive ion etching is a major component of integrated circuit (IC) technology owing to its anisotropy and selectivity over masking and underlying layers [16]. Glass, however, shows an etch rate roughly an order of magnitude lower than silicon. As a result, relatively high biases are required when etching glass which compromises the masking material choice, smoothness of the etched surface and attainable etch depth [17–20]. ‘Wet’ etching borosilicate and aluminosilicate
glasses in hydrofluoric acid solutions show etch rates up to 8 μm min⁻¹ [8, 11, 21], but they exhibit isotropic etch profiles and frosted etched surfaces due to the presence of insoluble impurities [22]. Etching pure amorphous SiO₂ (fused silica/quartz), in contrast, results in optically transparent surfaces but the etch rate is on the order of 1 μm min⁻¹ [3]. While methods have been developed to reduce the loss of optical clarity during borosilicate HF etching [21, 23], fused silica still has the advantage of chemical purity which makes it compatible with CMOS processing techniques [3] and eliminates substrate autofluorescence [24]. Previous masking film materials for fused silica included chromium (Cr) [25], photoresists [13], polysilicon (polySi), amorphous silicon, aluminum, silicon nitride and chromium/gold (Cr/Au) [3]. For example, fused silica was wet etched 60 μm deep with a Cr/Au mask in 49% HF for 1 h [3] and 104 μm deep with a stress-reduced Cr mask in a heated buffered ammonium fluoride solution for 7 h [25]. If fused silica etch depths substantially greater than 100 μm are desired, very long etching times and/or concentrated HF solutions (49% by mass) are required, which causes HF to eventually migrate through most masking materials resulting in surface pitting and eventually mask deterioration and/or liftoff.

We report here a method to etch fused silica greater than 600 μm deep while keeping the substrate free of pits and maintaining a polished etched surface suitable for biological imaging, using an HF-resistant photosensitive resist (HFPR), ProTEK PSA (Brewer Science, Inc., Rolla, MO). ProTEK PSA is a negative tone resist system that consists of a thermoplastic cycloolefin copolymer which is a highly nonpolar and hydrophobic polymer. A photosensitive agent induces crosslinking which renders it insoluble in hydrocarbon-based developing solvents. The minimal free volume of the cross-linked material, in addition to the above-mentioned properties, minimizes the diffusion of polar HF molecules through the HFPR and thus renders it resistant to 49% HF [26, 27]. Undercut and mask deterioration are compared for substrates masked with the HFPR, Cr/Au and polySi alone and Cr/Au and polySi films protected by the HFPR. Additionally, surface smoothness is compared for different etch depths. We demonstrate deep trenches in fused silica wafers using just the HFPR, obviating the need of furnaces or metal evaporators to deposit masking films for many etching applications. Even though etching 640 μm deep into a wafer approximately 650 μm thick, the surface of the resulting suspended membrane was smooth enough to allow fluorescence imaging without loss of image quality. Finally, we plasma etch pore arrays in the fused silica-suspended membranes which may be used as a planar patch-clamp electrode and/or suction electrode arrays.

**Experimental details**

*Materials and substrate preparation*

Square GE 124 fused silica wafers, 25.4 mm wide, 0.2 mm thick (Structure Probe Inc., West Chester, PA), 50 mm wide, 0.2 mm thick (Marc Optics Inc., Santa Ana, CA) and 50 mm wide, 0.55–0.65 mm thick (Quartz Scientific, Inc., Vancouver, WA) were used in this study. Photolithography and etching experiments were performed in class 100 and 1000 clean rooms, respectively. All wafers were first cleaned in a piranha solution (3:1, 95% sulfuric acid: 30% hydrogen peroxide, General Chemical, Parsippany, NJ). The general process flow is shown in figure 1. For the gold masks, a 10 nm adhesion film of chromium and 200 or 240 nm of gold were evaporated with a Mark 40 Electron Beam Evaporator (CHI Industries, Fremont, CA). For polySi masks, approximately 1550 nm of polySi was deposited on both sides of the wafer in a low pressure chemical vapor deposition (LPCVD) tube furnace (Tyster Corporation, Torrance, CA) with the following parameters: 40 Pa (300 mT) pressure, 600 °C deposition temperature, and 3 h deposition time. The HFPR, ProTEK PSA (Brewer Science, Inc., Rolla, MO), was photolithographically patterned according to the manufacturer’s guidelines with a MA6 mask aligner (SUSS Microtec Ag, Garching, Germany). All wafers were patterned with pores 850–900 μm in diameter. Following development, wafers with the HFPR were further baked at 200 °C for at least 10 min to ensure a highly cross-linked film due to the long HF exposure times used here. The opposite side of all wafers used in this study was blanket protected with the HFPR to protect from HF vapor. AZ5214E photoresist (AZ Electronic Materials USA Corporation, Branchburg, NJ) and the HFPR were used to pattern the poly Si and Cr/Au films, respectively. Au and Cr were etched with Gold Etch Type TFA (Transene Company, Inc, Danvers, MA) and CR-7S Chromium Etchant (Cyantek Corp., Fremont, CA) respectively. PolySi was etched in a SLR-770 deep reactive ion etcher (DRIE) (Oerlikon, Switzerland) with the Bosch process [28]. Table 1 summarizes the major equipment required for the processing of the different masking materials.
Fused silica wet etching

Fused silica was etched in 49% (by mass) hydrofluoric acid (General Chemical) in a custom-built polyethylene o-ring wafer holder (figure 1(D)). Following etching, the wafers were immediately immersed in water and then in a piranha solution to remove the HFPR. Au/Cr were removed in their respective etchants, while polySi was removed in a custom-built xenon difluoride (XeF₂) vapor etcher [29]. Wafer thicknesses before etching were measured with a micrometer or a Wyko NT3300 optical profilometer (Veeco Instruments, Inc., Plainview, NY). Etch depths were measured with a Dektak 6 stylus profilometer (Veeco Instruments) or Wyko NT3300 optical profilometer. Surface roughness was measured with the NT3300 profilometer in phase shift scanning interferometry mode.

Pore array device

For the devices with pore arrays, pores 20–50 μm in diameter were etched through suspended fused silica membranes (0.85 mm diameter, 6–13 μm thick) on 0.2 mm thick wafers as previously described [15]. Briefly, a 10 nm titanium adhesion film and 450 nm nickel masking film were evaporated onto the wafer and electrochemically patterned in class 10 85% phosphoric acid (general chemical). The exposed fused silica was then etched in an Ulvac Neutral Loop Density 570 DRIE etching system with the following parameters: 52.0 A of current to the top and bottom electromagnetic coils, 30.6 A to the middle coil, 1000 W to the RF antenna, 90 W to the substrate electrode, 30 sccm C₃F₈, 0.4 Pa (3 mT) pressure, 50 °C chamber temperature, 150 °C shield temperature and 20 °C substrate electrode temperature. Twenty-one devices, 3.5 mm × 3.5 mm (length × width), were diced from a 25.4 mm × 25.4 mm wafer (American Precision Dicing, San Jose, CA).

Optical imaging of fluorescent neural tissue

Neural tissue from the medicinal leech Hirudo verbana was used for biological imaging experiments. A single mid-body ganglion was isolated from the leech as previously described [30] and pinned to a polymethylsiloxane (PDMS) slab. Autofluorescence from the ventral side of the ganglion was imaged with an upright microscope objective. For imaging through suspended fused silica membranes, the ganglion still attached to the PDMS slab was positioned ventral side down over the suspended membrane with a micromanipulator. Images were acquired with an inverted objective through fused silica surfaces that were HF etched 195 and 640 μm deep. Throughout the imaging experiments, tissue remained immersed in normal leech saline solution [31]. A custom dual upright and inverted microscope was used for imaging experiments. This system was constructed by attaching a modified Olympus upright BX WI microscope to an Olympus inverted IX51 microscope (Olympus America, Inc., Center Valley, PA) with a custom-built stage in between the two. A 5 W LED light (LedEngin, Inc., San Jose, CA) provided illumination at 460 nm. The filter set comprised a 460/50 nm bandpass excitation filter, 505 nm dichroic mirror, and 510 nm long pass emission filter (Chroma Technology Corporation, Bellows Falls, VT). Images were recorded with an Olympus E420 digital camera.

Results and discussion

Integrity of masking materials in 49% HF

Chromium/gold + HFPR. Gold–chromium films have widely been used as a masking material for HF etching due to the chemical inertness of Au and the strong adhesion of Cr to glass and fused silica. With just Cr/Au as a mask for etching approximately 200 μm deep into fused silica, we were able to obtain an optically transparent etched surface, although the silica surface under the mask was severely pitted (figure 2(A)). Previous reports using a Cr/Au mask also showed similar pitting features [3]. However, when we further protected the Cr/Au film with the HFPR, the Cr/Au mask at the feature’s edge was much better preserved even after almost 3 h of

| Table 1. Major processing equipment/steps required for processing masking materials. |
|---------------------------------|-----------------|-----------------|
|                                | HFPR            | Cr/Au           | PolySi          |
| --------------------------------|-----------------|-----------------|
| Deposition                      | Spin-coater     | Metal evaporator| LPCVD furnace   |
| Patterning                      | UV lithography  | UV lithography  | UV lithography  |
| Pattern transfer                | –               | Cr/Au etchant   | DRIE            |
| Film removal                    | Piranha solution| Cr/Au etchant   | XeF₂ (g)        |

Figure 2. Fused silica wafers etched in HF with different masks: (A) Wafers masked with Cr/Au alone, (B) Cr/Au combined with the HFPR and (C) the HFPR alone were etched approximately 200 μm deep. Substrates masked with (D) polySi alone, (E) polySi combined with the HFPR and (F) the HFPR alone were etched approximately 600 μm deep. Scale bars are 1 mm.
etching time (166 μm deep). Pitting was also significantly reduced compared to the wafers masked with just Cr/Au mask (figure 2(B)).

**Polysilicon + HFPR.** To etch fused silica 600 μm deep we chose a polySi mask because it has been previously demonstrated to show substantially less pitting than Cr/Au [3]. However, our attempts to use a 1.5 μm thick polySi film were unsuccessful. As in previous reports, the polySi film remained defect free at 40 min of etching in 49% HF [3], but after 2 h of etching defects in the polySi film developed, and after 5 h we had to stop the etch because the film was so pitted that HF solution leaked under the o-ring in the wafer holder (figure 2(D)). To resolve this problem, we patterned a single layer of the HFPR on the polySi film to protect it. This strategy allowed us to etch over 600 μm in fused silica, although after 9 h of etching time, the polySi and fused silica underneath were significantly pitted. Additionally, the polySi at the edge of the patterned features had deteriorated (figure 2(E)), which sometimes caused polySi debris to fall onto the flat etching surface resulting in micromasking.

**HFPR alone.** We wanted to test the feasibility of using just a photoresist for deep HF etching. Fused silica wafers masked with the HFPR alone were etched approximately 200 and 600 μm deep (figures 2(C), (F)). The HFPR never showed signs of being attacked by 49% HF. While there was significant undercut using the HFPR, the fused silica under the HFPR was not pitted even after 15 h of etching (three consecutive etching steps on the same wafer). Furthermore, the etched surface remained polished and thus suitable for optical imaging.

**Fused silica undercut and etch rates with different masking films**

Masking with Cr/Au + HFPR resulted in etch rates of $1.12 \pm 0.06 \, \mu m \, min^{-1}$ (mean ± standard deviation, $N = 7$ wafers etched for 162–184 min; figure 3(A)). This rate corresponds to a final etch depth of $202 \pm 11 \, \mu m$ in 3 h. In contrast to the substantial variability of etch rates between wafers, etch rates were spatially very uniform within individual wafers: When two wafers were processed with a mask that defined multiple trenches, the depths of these trenches were constant to within 0.33 and 0.48 μm (root-mean-square variability of etch depth of 15 measured trenches in each of two wafers) after 166 min of etching. Etch rates were very similar when the HFPR was used alone, even though that resulted in much larger undercuts compared to the Cr/Au + HFPR when etching approximately 200 μm (figure 3(A)).

Since we were interested in fabricating thin membranes (see below), this variability in the etch rate necessitated interrupting the etch process for intermediate depth measurements when we etched 600 μm deep features. PolySi + HFPR and the HFPR alone were used as masks to etch more than 600 μm deep. Compared to wafers masked with polySi + HFPR, undercut for wafers masked with the
HFPR alone was much more extensive, extending all the way to the o-ring for these long etching times (figure 3(B)). The wafer for which a profilometry trace is shown in figure 3(B) was first masked with a 900 \( \mu \text{m} \) diameter feature and etched 390 \( \mu \text{m} \) deep. The wafer was then cleaned (to measure the etch depth), and the HFPR was patterned again, but with a 3.0 mm diameter feature aligned over the original 900 \( \mu \text{m} \) feature so as not to coat over the original undercut slope, and etched again in HF. The reduced etch rate with the HFPR-masked wafers compared to polySi + HFPR-masked wafers for 600 \( \mu \text{m} \) deep etching (figure 3(B)) is likely caused by the extensive undercut with the HFPR alone combined with the increased etch depth: More fused silica is etched laterally, creating additional reaction products which must diffuse out before HF can diffuse into the mask opening.

**Smoothness of etched surfaces**

Etched surface roughness profiles were obtained from fused silica wafers using phase shift scanning interferometry \([32]\). Average roughness of etched surfaces versus etch depth is plotted in figure 4. There is no significant difference of surface roughness for different etch depths and/or different masking schemes. Because the average roughness cannot adequately capture all spatial aspects, representative three-dimensional roughness profiles are shown for surfaces etched to depths of 32, 166, and 645 \( \mu \text{m} \) in figure 5 to visualize defect patterns. To avoid any variability caused by different masking schemes, the comparison is only between the HFPR-masked wafers. The insets of figure 5 demonstrate that the spatial characteristics of these surfaces are similar regardless of etch depth. This suggests defect patterns may already be seeded on the surface in the first few minutes of etching. Because the surface roughness is very similar for a clean fused silica wafer and one patterned with the HFPR but not etched (figure 4), further investigations would be needed to determine if the origin of these defects are the result of mask patterning, surface defects in the native fused silica, and/or diffusion gradients in the liquid etching medium. In summary, we obtained surfaces with an average roughness under 10 nm at the bottom of 600 \( \mu \text{m} \) deep etched features, the deepest HF etching in fused silica reported to date.

**Fluorescence imaging of biological tissue through the etched surfaces**

To demonstrate that deep etching results in devices compatible with optical microscopy, we fluorescently imaged biological tissue through the fused silica surfaces. Neural tissue was imaged from above with an upright microscope objective (figure 6(A)). The same neural tissue was turned over and positioned over fused silica surfaces that were etched 195 and 640 \( \mu \text{m} \) deep (figures 6(B), (C)). As seen in the figure, imaging through either of the etched surfaces with an inverted microscope objective did not negatively affect the image quality.

**Pore arrays batch fabricated in suspended fused silica membranes**

A fused silica wafer (188–195 \( \mu \text{m} \) thick) was etched in HF to a depth of 182 \( \mu \text{m} \) resulting in arrays of suspended...
Figure 6. HF etching fused wafers etched does not negatively affect image quality in fluorescence microscopy. (A) Fluorescence image from neural tissue obtained with an upright microscope objective. (B–C) Fluorescence image of the same neural tissue imaged through a fused silica membrane that was HF etched (B) 195 μm deep (an 8.2 μm thick suspended fused silica membrane) and (C) 640 μm deep (a 10–16.5 μm thick suspended fused silica membrane).

Figure 7. Batch fabrication of through pore arrays in suspended fused silica membranes. (A) Deep trenches etched with HF into a fused silica wafer (21 devices). The scale bar is 3.5 mm. The inset shows a single device (3.5 mm × 3.5 mm in length and width) diced from a wafer with a pore array that was plasma etched into the suspended fused silica membrane. The next inset shows a close up of the pore array fabricated into a thin suspended fused silica membrane. The scale bar is 100 μm. (B) Schematic of a single pore array device interfaced to a PDMS fluidic chamber allowing negative pressure to be applied to immobilize biological tissue. The inset shows a cross section of a single device with a through pore array in the thin-suspended membrane.

Conclusion

We have presented a processing scheme that can wet etch at least 600 μm deep into fused silica. We achieved this with a HF-resistant photosensitive resist, ProTEK PSA. This masking process, which does not require more than standard photolithography equipment (table 1), yielded substrates free of pits and etched surfaces with an average roughness on the order of 10 nm (figure 5) albeit with significant feature undercut. When the HFPR was combined with Cr/Au or polySi films, undercut was more limited with the tradeoff of substantial surface pitting for long etch times. In the future, it may be possible to minimize the surface pitting and undercut by combining the HFPR with stress-controlled ‘hard’ masks [21, 25].

We demonstrated that etched fused silica surfaces allow for optical imaging through the device. This processing scheme will contribute to the fabrication of transparent biological devices. One application is planar patch-clamp electrodes [4, 15, 33]. Another is multielectrode arrays (MEAs) [34]: Perforated MEAs have received increased attention recently because they simultaneously enable tissue immobilization, oxygen perfusion and recording from multiple electrodes in parallel, but currently available devices are not transparent [35, 36]. We are now able to fabricate similar devices in an optically transparent substrate that will allow for optical imaging [37]. Finally, due to the simplicity of patterning a photosensitive resist, researchers will be able to test the performance of a variety of fused silica and/or glass device prototypes in a minimal amount of processing time.

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