TECHNICAL NOTE

Deep SiO\textsubscript{2} etching with Al and AlN masks for MEMS devices

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Abstract. Silicon oxide-based materials such as quartz and silica are widely used in Microelectromechanical Systems (MEMS). One way to enhance capability of their deep plasma etching is to increase selectivity by the use of hard masks. Though this approach was studied previously, information on the use of hard masks for etching of silicon-oxide based materials on 200 mm substrates is scarce. We present results of etching process development for amorphous silicon oxide using Al and AlN masks with a view of application of results for etching of silica and quartz. Three gas chemistries (C\textsubscript{4}F\textsubscript{6}/O\textsubscript{2}, CF\textsubscript{4} and SF\textsubscript{6}) and their mixtures were compared in an industrial reactive ion etch (RIE) chamber with two plasma sources. It was established that pure SF\textsubscript{6} is the best etchant and AlN is better mask than Al for providing higher selectivity and sidewall angle close to vertical. A range of etching parameters for micromasking-free etching was established and etched structures of up to 4:1 aspect ratio were created in 21\textmu m-thick oxide using the process with etch rate of 0.32-0.36 \textmu m/min and selectivity to AlN mask of (38-49):1.

1. Introduction
In the last decade a few MEMS applications have required deep anisotropic etching of doped and undoped silicon oxide films as well as silicon oxide-based glasses and quartz. Among these groups are quartz resonators, optical MEMS and micro total analysis systems (\muTAS) or bio-MEMS. Previously, deep etching of quartz, silica and other glasses was evaluated for substrates with diameter \leq 150 mm and found application in the prototyping of mentioned devices. Therefore, engineers continued to put efforts into improvement of deep etch of these materials [1-9]. It is known that deep and high aspect ratio etching of glasses is not feasible with standard photoresist masks due to low selectivity and mask profile degradation during prolonged process, therefore, the use of metal or hard masks is preferable.

A few metal masks with different gas mixtures have been investigated so far. In [2] electroplated Cr hard mask was created by wet etching and used for etching of borosilicate glass in RIE chamber with two RF plasma sources (inductively coupled plasma (ICP) and capacitively coupled plasma (CCP)) using SF\textsubscript{6}/Ar chemistry: 32 \textmu m-deep trenches with aspect ratio 2:1 have been achieved. However, limited selectivity SiO\textsubscript{2}/Cr of only 35:1 required very thick (3\textmu m) Cr mask, which is an obstacle for using this approach for etching structures with higher aspect ratios. In [3], even thicker, 5 \textmu m Ni mask was created by electroplating with subsequent lift-off and used for etching different glasses with selectivity (23-35):1. Aspect ratio of 2.5 was reported for 100 \textmu m deep structures with sidewall angle 78\textdegree. Very thick (300 \textmu m) Ni shadow mask was investigated recently for deep etching of quartz in RIE system, however, the process
displayed micromasking effect which was only partially reduced by adding Ar to SF₆, despite low aspect ratio (<1) [4].

Further increase of aspect ratio requires higher selectivity to mask. In [5, 6] this issue was solved by converting Al mask into Al₂O₃ by using alternating etch–oxidize steps. During oxidation steps, the surface layer of Al was converted into Al₂O₃ that yielded selectivity SiO₂/Al₂O₃ about 100:1. Due to high selectivity to mask, 5μm-thick SiO₂ waveguides were etched with profile close to vertical using initial Al mask of only 100 nm. However, ratio of oxidation time to etch time was about 1:1, hence, the process duration increased twice and throughput suffered considerably.

Other researchers also reported on high selectivity possible with Al₂O₃ and AlN masks. In [7] it was reported that AlN mask provided very high selectivity to SiO₂ when the gas composition of C₄F₆/SF₆ mixture, pressure and bias power are properly tuned. Similarly, in [8] it was reported that in ICP-RIE system with SF₆/O₂ plasma, the etch rate of AlN can be brought down to 1 nm/min. Though both paper results were obtained only with blanket wafers without any mask pattern, they are a good hint for researches to continue process development in this direction. Also it should be noted that in all cited papers the experiments were performed with wafers of diameter ≤ 150 mm. Now mainstream MEMS industry widely adopts 200 mm substrates but information on deep glass etch on these substrates is scarce so far. Due to complexity of plasma etching, the transfer of a process to wafers of larger diameter rarely reduces simply to scaling of basic process parameters in accordance with scaling of wafer diameter. Besides, sometimes full scaling is not possible due to limitations of RF generators and pumps. In most cases an extensive design of experiment is required. Thus, the purpose of this paper is to investigate possibilities for deep etching of undoped silicon oxide glass (USG) with hard masks on 200 mm substrates in industrial equipment. Since chemical composition of USG, silica and quartz is the same it is feasible to use USG for etching study with a view of potential application of results for etching of silica and quartz.

2. Experimental

We deposited amorphous USG films with thickness up to 21 μm by plasma enhanced chemical vapor deposition (PECVD) on 200 mm silicon wafers in a Sequel Express Concept 2 chamber from Novellus Systems, Inc. using SiH₄/N₂O/N₂ gas chemistry. USG etch process development was performed in Omega® advanced planar source (APS) process module from SPTS Technologies which has two high density 13.56 MHz plasma sources (ICP and CCP), designed to etch materials which are difficult to etch using conventional RIE or ICP sources. The process chamber is of metallic construction and its walls are heated to 130°C to reduce the level of deposition on it, thereby increasing process stability and the mean time between cleans. Through the use of permanent magnets, the plasma is confined, contributing to an increase in plasma density necessary for the successful etching of strongly bonded materials [9]. Wafer is sitting on the cooled chuck with electrostatic clamping and helium backside cooling. The distance from the wafer chuck to the top chamber plate is 225 mm. If otherwise is not stated, the temperature of the chuck was maintained by the chiller at 10 °C.

Two types of hard masks were evaluated in this study. First mask was Al deposited by conventional sputtering. Second mask was AlN deposited by reactive sputtering with pulsed direct current magnetron and RF biased chuck in a Sigma® Deposition System from SPTS Technologies. It should be mentioned that before deposition of AlN the surface of USG was subjected to chemical mechanical polishing to remove USG layer of about 200 nm. This step reduced the surface roughness of USG from 6.2 to 0.2 nm (rms) and enabled high quality of deposited AlN. Patterning of Al mask was performed by standard
lithography and etching with photoresist mask while patterning of AlN was performed with dual mask of photoresist and SiN. The latter was deposited in Sequel Express Concept 2 chamber. Before lithography step, the stress on wafers was minimized by depositing compensating SiN layer on the backside. It allowed to maintain maximal wafer warpage within 50 μm. SiN etching was performed in APS chamber with CF3/H2/He chemistry followed by photoresist stripping and wet clean. Etching of Al and AlN was performed with photoresist and SiN masks, respectively, in the same Omega® etch system using ICP-CCP metal etch process module incorporating a radial coil design. A basic gas mixture of Cl2/BCl3/Ar was used for Al etching while for AlN a buffer gas was added as described in [11]. For both masks the resulting sidewall profile was close to vertical. Tilted scanning electron microscope (SEM) image after AlN hard mask etching is presented in figure 1.

![SEM image of AlN mask](image)

**Figure 1.** Tilted SEM- image of 2.0 μm-thick AlN mask with top remaining SiN layer after etching in Omega® ICP-CCP process module.

Remaining SiN on top of AlN was not removed and served as an additional mask during short initial period of USG etching. Remaining photoresist mask on Al was removed before USG etch. For both Al and AlN masks, the wafers were subjected to wet clean for metal etch by-products removal before USG etch.

### 3. Results and discussion

#### 3.1 Aluminum mask

Initial recipe for 150 mm wafers reported in [10] had the following set of parameters: coil (ICP) power 1800 W, platen (bias) power 150 W, pressure 6 mT, flow rates C4F8/O2= 80/10 sccm, chuck temperature 55°C. To scale this recipe for etching of 200 mm wafers, we would need to ramp up RF powers proportionally to the increase of wafer area i.e. by 1.8 times. Calculation yields required coil power of 3200 W and platen power of 270 W. However, requirement of the increased coil power could not be fully satisfied in our 200 mm APS process module due to limitation of maximal power of the RF generator available at our facility. Thus, we tested the recipe with coil power 2000 W and platen power 300 W (other parameters were taken from the recipe for 150 mm wafers). The outcome of etch process was smooth etched bottom, however, sidewall was tapered with the angle of 83° (figure 2), comparing to angle ≥ 88° reported for 150 mm wafers.
Definitely, the lack of coil power for 200 mm wafers resulted in the non-sufficient ion density. As explained in [13], the result was an increase in passivating neutral-to-ion flux ratio leading to more tapered profile due to increasing re-deposition of C-rich byproducts on sidewalls. Therefore, in order to achieve vertical profile we decided to study effect of reduction of C/F ratio in the gas chemistry by diluting C$_4$F$_8$ with CF$_4$. Figure 3 shows the dependence of USG etch rate and sidewall angle in 5-μm wide trench versus C$_4$F$_8$ percentage in the mixture with CF$_4$. It is seen that the reduction of C$_4$F$_8$ percentage resulted in improvement of the sidewall angle even at a lower chuck temperature of 10°C used in these experiments. With full exclusion of C$_4$F$_8$, sidewall angle reached the value of 88°. Figure 4 presents cross-sectional image of 4μm- thick USG etched in pure CF$_4$ plasma.
**Figure 3.** USG etch rate and sidewall angle versus percentage of \(\text{CF}_4\) in the mixture with 80 sccm of \(\text{CF}_4\) (coil/platen power 1600/200 W, pressure 4 mT).

**Figure 4.** Cross-sectional SEM image of 4 \(\mu\)m-thick USG etched in pure \(\text{CF}_4\) with resulting sidewall angle 88° (coil/platen power 1600/200 W, flow rate 80 sccm, pressure 4 mT). Cross-section was done by focused ion beam with preliminary deposition of protective Pt layer.

However, it is also obvious from figure 4 that the facet on Al mask generated by sputtering during USG etch will affect pattern transfer fidelity during deeper etching. Therefore, it is desirable to reduce energy of bombarding ions in order to increase selectivity to mask. Obvious approach in this case is to reduce platen power. However, we found that with the reduction of platen power down to 100 W a phenomenon of micromasking happened in the vicinity of Al mask even in features as large as 18 \(\mu\)m (figure 5).

**Figure 5.** Tilted SEM image of USG etched with Al mask using \(\text{CF}_4\) chemistry at 100 W platen power (coil power 1600 W, flow rate 80 sccm, pressure 4 mT).

The cause of micromasking is sputtering and re-deposition of Al mask as confirmed by the presence of strong Al peak in EDX spectrum of micromasked area. Further experiments proved that with pure
CF<sub>4</sub> chemistry, the minimal platen power allowing reliably to avoid micromasking in features of ≥ 5.0µm was 150 W. Therefore, this value of platen power was chosen for evaluation of high aspect ratio etching. Figure 6 presents results of etching of 10 µm-thick USG using Al mask and CF<sub>4</sub> chemistry. An average etch rate in 5 µm-wide trench was 0.39 µm/min, selectivity USG/Al was 17:1.

Figure 6. Cross-sectional SEM image of 5.0 µm-wide trench etched with Al mask in 10 µm-thick USG using CF<sub>4</sub> chemistry (coil/platen power: 1800/150 W; flow rate 100 sccm; pressure 5mT). Inset: EDX spectrum of by-products re-deposited on sidewall.

It is seen that sidewall profile at the top part of the trench became tapered due to formation of a facet on Al mask and its subsequent transfer into USG layer. Besides, there is visible necking below the top facet due to by-product re-deposition on the sidewall. Though initial Al mask of 2.0 µm was not fully consumed, deeper etching using the same mask thickness and etch recipe was not feasible due to expected bigger tapering of profile and critical dimensions (CD) loss at a trench top. From the inset in figure 6, we see that Al is a substantial part of the sidewall by-products. Thus, for feasibility of deep etching it is imperative to increase selectivity of USG etch to Al mask on the top corner of the pattern i.e. to reduce sputtering effect. However, as mentioned earlier, the possibility of reduction of platen power was already exhausted by reducing it to 150 W. Therefore, we explored another option for increasing selectivity of etching based on the idea of papers [5, 6] where authors converted Al mask into Al<sub>2</sub>O<sub>3</sub> by introducing oxidation steps. However, instead of using alternating etch–oxidize steps we presumed that adding O<sub>2</sub> into gas mixture will create competing process of oxidation on Al surface and, thus, decrease the sputtering of Al mask. Hence, selectivity to Al mask was expected to increase. Indeed, selectivity to Al increased with the same bias power of 150 W from 17:1 in pure CF<sub>4</sub> to 23:1 with added 20 sccm O<sub>2</sub>. However, this improvement was not sufficient. To understand why approach with added oxygen was not so effective is possible based on the information in [10] where it was reported that adding O<sub>2</sub> to C<sub>4</sub>F<sub>8</sub> increased the angular etching yield of SiO<sub>2</sub> by shifting...
conditions from net deposition to net etching. Similarly, we explained an increased faceting of Al mask with added O2 in our case: adding O2 resulted in the depletion of adsorbed carbon, and Al surface became more exposed to the attack by sputtering ions. Thus, increased sputtering almost canceled the effect of oxidation in our etching chamber. Moreover, [10] reported that for Pyrex etch with Al mask, an increase in O2 concentration even resulted in the selectivity drop. Therefore, we concluded that adding O2 into the gas mixture cannot replace approach with cycles of Al oxidation and etching. However, the latter approach also was rejected because of very low throughput. Obviously, instead of trying to oxidize Al mask in the course of etching, it is better to deposit directly the mask of Al2O3 or other material providing higher selectivity.

3.2 Aluminum nitride mask

Paper [8] reported, based on blanket wafer study, that both Al2O3 and AlN had very high etch selectivity to USG. This is expected because bond dissociation energies for Al-O (502 kJ/mol) and for Al-N (368 kJ/mol) are noticeably higher than for Al-Al (264 kJ/mol) [14]. Therefore, we continued our development using AlN hard mask deposited by reactive sputtering. Initial testing on blanket AlN wafers revealed that with the same CF4 etching recipe as in figure 6, the selectivity of USG/AlN reached the value of 28:1 comparing to 17:1 for USG/Al. Etching of patterned wafers using AlN mask gave smooth etched bottom. Figure 7 shows that 16.5 μm-deep USG etch was accomplished with 2.0 μm-thick AlN mask, however, the sidewall shows the same two distortions as in the case of Al mask: one is a facet at the top portion of the sidewall and another is a sidewall necking, right below the facet.

Figure 7. Cross-sectional SEM image of 16.5 μm-thick USG etched with AlN mask using CF4 plasma (coil/platen power: 1800/150 W; flow rate 80 sccm; pressure 5mT). Pt layer was deposited for protection of the structure during specimen cutting by focused ion beam.
Thus, comparing to Al mask, AlN mask provided fabrication of higher aspect ratio structures in SiO₂, due to higher selectivity, however, additional process tuning was required to overcome issues of corner sputtering and sidewall necking. We presumed that an increase of selectivity would be useful to suppress both mentioned distortions of sidewall profile. Indeed, with an increase of overall selectivity we could expect an increase of selectivity at a top corner as well. At the same time, an increase of selectivity would result in less mask material sputtered and accordingly less material re-deposited on sidewall, thus, decreasing necking. To achieve a decreased necking it seemed reasonable to reduce carbon content in the gas chemistry because we saw high carbon content of 23.4% in the composition of by-products on the sidewall (figure 6). A study of effect of carbon content reduction was implemented by adding SF₆ into gas chemistry and reducing accordingly the fraction of CF₄. Results for the etch rate and selectivity versus percentage of SF₆ in the mixture with CF₄ for blanket wafers are presented in figure 8.

![Figure 8](image-url)

**Figure 8.** Etch rate and selectivity for blanket wafers versus SF₆ percentage in the mixture SF₆/CF₄ (coil/platen power: 1850/150 W; total flow rate 100 sccm; pressure 5 mT).

It is seen that with the change from 100% CF₄ to 100% SF₆ the selectivity increased almost two-fold though the etch rate of USG somewhat dropped. To compare effects of CF₄ and SF₆ plasma on the sidewall re-deposition we used simplified approach of etching 4 μm-thick USG layer with AlN mask and comparing etch CD bias in 2 μm-wide trenches. It was found that for CF₄ plasma, the trench width after etching shrank by 0.63 μm while for SF₆ plasma only by 0.17 μm. Thus, comparing to CF₄, the SF₆ process provided both better selectivity to AlN mask and less by-product re-deposition on sidewalls. Therefore, we continued SF₆ plasma process optimization. Figure 9 presents USG etch rate and selectivity versus platen power using SF₆ plasma.
It is seen that selectivity drastically improved with the decrease of platen power, however, the etch rate dropped. Similarly to CF$_4$ etching, there was minimal threshold chuck power of 160 W required to provide micromasking-free etching. To strike a balance between selectivity, etch rate and prevention of micromasking the platen power of 180 W was chosen for etching of patterned wafers. First, we performed partial etching of wafer having trenches of different width and, using cross-sectional SEM images, gathered information on the pattern width dependence of the etch rate (figure 10).

From figure 10 it follows that with the increase of trench width from 2 to 25 μm the etch rate increased by 20% and then saturated until trench width reached 75μm. With subsequent increase of the trench width, the etch rate slightly dropped as a result of microrloading effect. Also we observed a drop of the average etch rates in trenches with an increased etch depth. For 5μm-wide trenches this drop comprised 16% with the increase of etch depth from 3 to 20μm. This information was used for deciding etch times for full opening of targeted trenches in 21 μm-thick USG on silicon. Figure 11 presents cross-section of 5 μm-wide trench etched by optimized SF$_6$ recipe.
Comparing to image in figure 6, where etching was done with Al mask using CF₄ plasma, by-products layer on the sidewall in figure 11 is much thinner and re-deposited Al comprises only 7 atomic % comparing to 23% in case of Al mask. Still some faceting and necking effects is visible in this trench with 4:1 aspect ratio. Figure 12 shows tilted view of arrays of 15 µm-wide cavities after etching with optimized recipe in SF₆ plasma.

**Figure 11.** Cross-sectional SEM image of 21 µm-thick USG etched in SF₆ plasma with AlN mask (flow rate 100 sccm, coil/platen power 1850/180 W, pressure 5 mT). Inset: EDX spectrum of by-products re-deposited on sidewall.

<table>
<thead>
<tr>
<th>Element</th>
<th>Atomic %</th>
</tr>
</thead>
<tbody>
<tr>
<td>C</td>
<td>9</td>
</tr>
<tr>
<td>O</td>
<td>51</td>
</tr>
<tr>
<td>F</td>
<td>4</td>
</tr>
<tr>
<td>Al</td>
<td>7</td>
</tr>
<tr>
<td>Si</td>
<td>26</td>
</tr>
<tr>
<td>S</td>
<td>2</td>
</tr>
<tr>
<td>Pt</td>
<td>1</td>
</tr>
</tbody>
</table>

**Figure 12.** Tilted SEM image of 21 µm-thick USG etched in SF₆ plasma with AlN mask (gas flow rate 100 sccm, coil/platen power 1850/180 W, pressure 5 mT).

It is seen that the sidewall profile of 15-µm wide cavities is close to vertical and etched bottom is smooth. It should be noted that the etch selectivity for patterned wafers is lower than for blanket wafers. We assume that oxygen released during USG etch on patterned wafers helps to remove atoms of sulfur.
adsorbed on the surface of AlN mask, thus, making AlN surface exposed to ion bombardment and etching by sputtering. Higher selectivity for patterned wafers was achieved with reduction of platen power from 180 W to 160 W; however, this value of platen power can be used only for trenches with the width of more than 5 µm due to appearance of micromasking on the bottom of narrow trenches at high aspect ratios.

Overall, our study confirmed good prospects of AlN hard mask for deep USG etching because high values of selectivity and possibility to achieve smooth etched bottom have been demonstrated. The results are also applicable to etching of silica and quartz, bearing in mind that, due to higher density of quartz, its etching rate will be slower by 15-20%.

Table 1 presents summary of the etching processes used in this study.

<table>
<thead>
<tr>
<th>Mask</th>
<th>Gas chemistry</th>
<th>Platen power, W</th>
<th>Coil power, W</th>
<th>USG etch rate, µm/min</th>
<th>Selectivity to mask</th>
<th>Profile,°</th>
<th>Aspect ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al</td>
<td>C₄F₈/O₂</td>
<td>300</td>
<td>2000</td>
<td>0.52</td>
<td>20:1</td>
<td>83</td>
<td>2:1</td>
</tr>
<tr>
<td>Al</td>
<td>CF₄</td>
<td>150</td>
<td>1850</td>
<td>0.39</td>
<td>17:1</td>
<td>88</td>
<td>2:1</td>
</tr>
<tr>
<td>AlN</td>
<td>CF₄</td>
<td>150</td>
<td>1850</td>
<td>0.39</td>
<td>24:1</td>
<td>88</td>
<td>3:1</td>
</tr>
<tr>
<td>AlN</td>
<td>SF₆</td>
<td>180</td>
<td>1850</td>
<td>0.36</td>
<td>38:1</td>
<td>88</td>
<td>4:1</td>
</tr>
<tr>
<td>AlN</td>
<td>SF₆</td>
<td>160</td>
<td>1850</td>
<td>0.32</td>
<td>49:1</td>
<td>88</td>
<td>2:1</td>
</tr>
</tbody>
</table>

Depending on the targeted application, further process optimization may be required for attaining higher values of etch depth or aspect ratio. Another area of improvement is enhancement of etch rate which can be achieved by more detailed process window study. Judging from the mentioned above values of bond dissociation energies of Al-N and Al-O, further improvement of selectivity and other parameters is expected with replacement of AlN mask by Al₂O₃ mask.

4. Summary
We performed optimization of deep etching with hard masks for undoped silicon oxide on 200 mm wafers. Comparing to Al mask, AlN mask enabled higher selectivity and etching profile closer to vertical. Out of three studied gas chemistries - C₄F₈/O₂, CF₄ and SF₆, the latter provided the least amount of by-products re-deposited on sidewalls and, hence, is most suitable for high aspect ratio etching. Optimized etching process with SF₆ plasma yielded structures of 4:1 aspect ratio in 21µm-thick USG. AlN mask is easier to manufacture comparing to such masks as Ni and Cr, which require complicated processes of electroplating and lift-off. Thick AlN mask (up to 3 µm) can be prepared by standard processes of reactive sputtering, lithography and plasma etching. Based on confirmed selectivity values we expect that masking and etching capability developed in this work is extendable for etching of silica and quartz to the depth around 100 µm.

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References: