Fabrication of sub-60-nm contact holes in silicon dioxide layers

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Abstract

We have developed a fabrication technique, comprising electron-beam writing, chemical shrinking, and silicon dioxide etching, for the fabrication of sub-60-nm contact holes. The dimensions of the contact holes after the electron-beam writing, chemical shrinkage, and plasma etching steps were 140, 93, and 53 nm, respectively. We carefully evaluated the critical process parameters, such as mixing-bake temperature, mixing-bake time, plasma etch selectivity, and the profile shapes. A higher mixing-bake temperature led to an overhang of the contact hole in the resist; the optimal mixing-bake conditions occur when heating at 110 °C for 70 s. A fluorinated mixture of gases (CHF₃/CF₄ = 1:1) was used to etch the nano-scale contact holes in the silicon dioxide layer. The formation of side-wall polymers during the plasma etch phase contributed further to the contact hole shrinkage in addition to resist chemical shrinkage. The ratio of the hole’s perimeter to its area affects the shrinkage dimensions of the etch process, especially for smaller contact holes. The pattern reduction due to side-wall polymer deposition in etch process has the inverse relationship with the diameter of contact hole.

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1. Introduction

The progress of research into integrated circuits (ICs) is focused mainly on two goals: (1) increasing circuit performance (mainly by increasing circuit speed) and (2) increasing the functional complexity of the circuits. Scaling down device sizes is considered a very effective means of achieving these goals, but discovering technologies for achieving sub-60-nm contact holes is still a challenge [1,2].

The ITRS 2002 Update[2] predicts that, for ASIC fabrication, the critical dimensions (CDs) for contact holes after etching will be required to be 65 and 45 nm by 2007 and 2010, respectively. Technologies that use chemical shrinkage for patterning sub-100-nm contact holes in resists have been reported previously [3]. In that report, the authors proposed using a polymeric shrinking agent that cross-links on
the pattern’s sidewalls at a suitable mixing-bake temperature, but such a shrinkage technique faces the problem of defect formation in the resist pattern at higher mixing-baking temperatures. Therefore, such chemical shrinkage technology has not been used previously for the generation of sub-60-nm features. Solving this problem, and to achieve the goal of preparing sub-60-nm contact holes in a silicon dioxide layer, requires controlling the shrinkage temperature for the ca. 90-nm contact holes in the resist and improving the etch process that allows further shrinkage to occur.

In this paper, we investigate chemical shrinkage process parameters that are used in conjunction with electron-beam lithography for preparing contact holes during ASIC fabrication. We have evaluated the various bake processes and process stabilities, and we discuss the cross-sectional images of holes formed by the chemical shrinkage process at various mixing temperatures. Upon completing the chemical shrinkage procedure, we also evaluated a series of dry etch recipes for the silicon dioxide layer.

2. Experimental

Table 1 lists the experimental flows and parameters used for preparing the sub-60-nm contact holes in the silicon dioxide layer. Electron-beam exposure was performed on a Leica Weprint 200 stepper. The silicon dioxide layer was grown by wet oxidation using a mixture of hydrogen (8000 cm$^3$/min) and oxygen (4999 cm$^3$/min) gases at 978 °C in a low-pressure furnace. The electron-beam energy was 40 keV, the beam size was 20 nm, and the exposure dose was 14 μC/cm$^2$. The developer for the JSR positive electron-beam resist (MES-1EG) was an aqueous 2.38% tetramethylammonium hydroxide (TMAH) solution. A positive-tone electron-beam resist was spin-coated on a silicon wafer (150 mm diameter) and baked at 110 °C for 120 s. The thickness of the resist film was ca. 650 nm. After exposure and a post-exposure bake (110 °C for 120 s), the wafer was developed using the TMAH solution. Again, a hard-bake was applied to the wafer (110 °C for 120 s). A chemical shrinkage procedure was then applied. The chemical shrinking agent (AZ R-200, Clariant) was coated over the resist pattern, and the wafer was again soft-baked. The film thickness was ca. 400 nm. A mixing-bake step at 110 °C for 70 s was then undertaken. The chemical shrinking agent reacted with the resist acid that diffused from the resist pattern. A cleaning step, using AZ R-2, which can wash away unreacted materials, was applied to the process. A hard-bake step was performed at 110 °C for 2 min. Finally, we applied a plasma process using a mixture of CHF$_3$ and CF$_4$ gases to etch the underlying silicon dioxide layer. Critical dimensions were evaluated using either an in-line scanning electron microscope (SEM, Hitachi S-6280) or a cross-sectional SEM (Hitachi S-4000).

3. Results and discussion

3.1. Chemical shrinkage processes for the electron-beam resist

As has been reported previously [3], the mixing-bake temperature and time are the critical factors
for fabricating sub-100-nm contact holes in a resist when using the chemical shrinkage technique. Basically, the residual acid in the resist pattern diffuses into the side-wall regions of the shrinkage agent, which leads to cross-linkage reactions of the shrinkage agent. The extent of hole shrinkage in the positive resist depends upon the intrinsic acid diffusion behavior, such as its diffusion coefficient, and the time. Fig. 1 illustrates the effect of mixing-bake temperature on the CDs of various contact holes for the positive electron-beam resist. Characterization using top-down SEM clearly indicates that the CDs of contact holes at any size gradually narrow as the mixing-bake temperature is increased from 100 to 120 °C. An initial hole size of 140 nm formed by electron-beam patterning is suitable for shrinking down to the sub-100-nm level. We evaluated the profile of the contact holes at various mixing-bake temperatures by using cross-sectional SEM. The SEM pictures in Fig. 2 clearly illustrate the profile of the contact holes. Initially (prior to applying the chemical shrinkage agent), the contact holes have vertical sidewalls and a smooth surface. After a 105 °C mixing-bake (subsequent to applying the chemical shrinkage agent), the sidewalls and surface covered with shrinkage agent appear to be slightly distorted. When the mixing-bake temperature was increased to 110 °C, the contact holes remain open, but we observe an overhang effect in the contact holes formed after heating at either 115 or 120 °C. We attribute this finding to the effect of thermal flow of the shrinkage agent on the resist’s sidewalls at higher bake temperatures. This type of defect is not observed when viewing from by SEM from above the surface, but is very apparent from the cross-sectional SEM image. Clearly, the upper limit of the mixing-bake temperature for the chemical shrinkage technique is 110 °C.

Fig. 3 demonstrates the effect of mixing-bake times on the CDs of various contact holes for the positive resist at 110 °C. We observe that the hole dimensions decrease rapidly during the first 60 s, but then gradually reach a plateau. Therefore, we believe that the optimal time for the mixing-bake process in this study is 70 s. The reason for this trend with respect to mixing-bake time is that the process is dependent on the abundance of diffusion acid in the resist pattern, which is limited. If the amount of acid in the resist is increased, the CD will become narrower. The initial 140-nm hole formed by electron-beam patterning is the only one of the holes that we have studied that is suitable for being shrunk down to a sub-100-nm hole. Next, we compare the effects of the soft-bake, mixing-bake, and hard-bake processes after applying the chemical shrinkage agent. The control conditions for soft-baking, mixing-baking, and hard-baking are 85 °C for 70 s, 110 °C for 70 s, and 110 °C for 2 min, respectively. Fig. 4 clearly indicates that the hard-bake temperature has no significant effect on the hole shrinkage ratio. The

Fig. 1. Dependence of various mixing-bake temperatures for 70 s on the critical dimensions (CDs) of contact holes formed by the chemical shrinkage technique. The initial hole sizes were 140, 180, 240, and 360 nm, respectively; top-down SEM was used to measure CD.

Fig. 2. Cross-sectional SEM image of contact holes in the resist after various mixing-bake temperatures.

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soft-bake temperature does have a significant effect above \(110^\circ C\), which we ascribe to the thermal flow effect discussed earlier. Among these baking processes, the mixing-bake temperature exhibits the most significant effect on the hole shrinkage ratio.

The hole shrinkage mechanism is closely dependent on the abundance of residual acid in the resist pattern. We exposed the wafer in the cleanroom environment (class 10) after initial contact hole definition for various delay hours. Fig. 5 indicates that the shrinkage ratio fluctuates between 33\% and 34\% after various delay times. Although not illustrated here, the SEM images for these holes taken from above are very similar. It has been reported in the literature that a chemically amplified resist is sensitive to the molecular base, and leads to T-top and footing problems for the positive resist when not immediately developed (post-exposure delay) [4]. The molecular base in the cleanroom might have reacted to some extent with the surface acid. Most acids under the resist film, however, are not influenced by the molecular base from the air in the cleanroom. Therefore, the diffusion of the acids out of the resist still occurs during the mixing-bake process and the delay time has no effect.

### 3.2. Fabrication of 53-nm contact holes

The fabrication of sub-60-nm contact holes in a silicon dioxide layer by the chemical shrinkage technique has not been reported previously. To ensure the applicability to nano-fabrication techniques, the resist should tolerate the etching process. Fig. 6 depicts the etch selectivity that we estimate from the ratio of the plasma etch rates of silicon dioxide and the positive resist under various mixtures of gases. The selectivity gradually increases upon increasing the ratio of CHF\(_3\) from 0 to 0.75, and then increases abruptly upon a further increase in the gas ratio to 1. It has been suggested in the literature [5] that oxygen byproducts formed during silicon dioxide etching can react with carbon residues, especially at fluorine/carbon ratio <2. Hence, the polymer formation blocks any further etching process.

The positive resist (650 nm thick) was coated onto a wafer upon which a silicon dioxide film had been grown. A variably shaped electron-beam was
used to pattern a 140-nm hole in the positive resist (Fig. 7(a)). The chemical shrinkage process was then undertaken by spin-coating the shrinkage agent onto the resist pattern, followed by a soft-bake at 85 °C for 70 s. The wafer was then subjected to a mixing-bake at 110 °C for 70 s. The residual acid diffuses out from the resist pattern into the shrinkage agent, which leads to acid-induced cross-linkage reactions taking place. After washing off the unreacted shrinkage agent and baking again at 110 °C for 2 min, the contact hole in the resist layer now has a 93-nm diameter (Fig. 7(b)). At the stage, the shrinkage ratio is ca. 33.6%. After the resist pattern had shrunk, the wafer was sent for plasma etching to fabricate a contact hole. Interestingly, the dimension of the contact hole in the silicon dioxide is not 93 nm: Fig. 8 indicates that the hole size is 53 nm. The total shrinkage ratio of the hole diameter after the chemical shrinkage and plasma etch processes is 62.1%. What happens to the contact holes in the silicon dioxide layer during the plasma etch process? The side-wall deposition of residual polymers during the plasma etch process plays a significant role in narrowing the contact hole dimensions.

The etch mechanism for contact hole fabrication in a silicon dioxide layer is very complicated, with the dimensions of the pattern formed during the etch process being controlled by a balance...
between the amount of polymer deposited and the etch conditions. Explanations have been proposed in the literature [6–9] regarding micro-loading and aspect ratio-dependent etching (ARDE) to explain the observations made during etch processes. Micro-loading describes the variations of the etch rate between areas having different pattern densities, with features in low-pattern-density areas etching faster than features in high-pattern-density areas. Effects that are due to the pattern dimensions, which includes effects related to transport of etchant species into the pattern, or transport of etch products out of the pattern, are generally referred to as ARDE. Table 2 lists a series of data for the contact holes obtained after the chemical shrinkage and dry etch processes. The smaller holes exhibit a higher shrinkage percentage for the etch process than the larger holes. We attribute this observation to the pattern dimension effect. As the pattern size is reduced, the probability for the flux of incoming polymer species to interact with the sidewall of the contact hole increases. Fig. 9 depicts the relationship between the ratio of the hole dimension before and after etching and the ratio of the hole perimeter to the hole area of these nano-scale contact holes. We find that a linear dependence exists for the series of holes studied. This finding suggests the pattern reduction arising from side-wall polymer deposition during the etch process has an inverse relationship to the pattern diameter. The probability of transportation of a polymer species onto the sidewall of a contact hole is related linearly to the inverse of the contact hole size (after chemical shrinking). As a consequence, the contact hole in the silicon dioxide layer becomes smaller than expected after the etching process. We estimate that the uniformities (1 sigma) of the proposed method for contact hole formation after the processes of lithography, chemical shrinkage, and plasma etching are 3.21, 3.16, and 2.76 nm, respectively.

### Table 2

The diameters of contact holes formed after the processes of chemical shrinkage and plasma etching in SiO₂, and their ratios ($R = \text{hole radius}$)

<table>
<thead>
<tr>
<th>Hole diameter after chemical shrinkage ($x$) (nm)</th>
<th>Hole diameter after dry etching in SiO₂ ($y$) (nm)</th>
<th>$y/x$ (%)</th>
<th>Perimeter/area ($2\pi R / \pi R^2$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>246</td>
<td>233</td>
<td>94.7</td>
<td>0.0163</td>
</tr>
<tr>
<td>149</td>
<td>128</td>
<td>85.9</td>
<td>0.0268</td>
</tr>
<tr>
<td>133</td>
<td>98</td>
<td>73.6</td>
<td>0.0301</td>
</tr>
<tr>
<td>119</td>
<td>79</td>
<td>66.6</td>
<td>0.0336</td>
</tr>
<tr>
<td>93</td>
<td>53</td>
<td>53.3</td>
<td>0.0430</td>
</tr>
</tbody>
</table>

Fig. 9. Linear dependence between the ratio of the hole dimension after and before etching and the ratio of the hole perimeter to hole area during the etching of the nano-scale contact holes.

4. Conclusions

We have established a successful fabrication technique for preparing sub-60-nm contact holes in a silicon dioxide layer by electron-beam lithography. We have discussed in detail the many factors that influence the performance of the
shrinkage process, such as the mixing-bake temperature, mixing-bake time, and hole dimensions before and after chemical shrinkage. Using this chemical shrinkage technique (mixing-bake of 110 °C for 70 s) and an etch gas of CHF$_3$/CF$_4$ (1:1), we obtained a minimum hole dimension of 53 nm. This technology meets the requirements [2] for contact hole fabrication in the year 2009. We propose that a nano-hole effect occurs during the etch-assisted shrinkage reaction because smaller holes have a higher percentage of polymer deposition in the resist sidewall than do larger holes.

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References