Electroless deposition of Ru films on Si substrates with surface pretreatments

Jing-Yu Chen, Shiau-Lin Huang, Pu-Wei Wu *, Pang Lin

Department of Materials Science and Engineering, National Chiao Tung University, Hsinchu 300, Taiwan

ARTICLE INFO

Available online 9 June 2012

Keywords:
Electroless deposition
Ru film
Si substrate
Surface pretreatment

ABSTRACT

Electroless Ru depositions on Si substrates undergoing surface pretreatments such as hydrogen fluoride (HF) etching, HF etching and activation, as well as HF etching, sensitization, and activation are explored. The plating bath contains K2RuCl5·xH2O, NaClO, NaOH, and NaNO2 at appropriate ratios. Continuous electroless Ru film is unable to obtain on the as-received Si wafer because the native oxide inhibits the process of nucleation and growth. In contrast, after surface pretreatments, dense and continuous Ru films are observed at reasonable growth rates. Contact angle measurements indicate a relatively hydrophilic surface after sensitization and activation, which leads to faster Ru film growths and larger surface roughness as compared to the HF-etched sample. X-ray photoelectron spectra confirm the formation of Sn and Pd nuclei and their presence promotes the heterogeneous growth of Ru films as evidenced by images from scanning electron microscope. In addition, depth profiling from Auger electron spectrometer suggests a uniform composition across the film thickness despite part of the Ru exists in an oxidized form.

1. Introduction

Ruthenium (Ru) and ruthenium oxide (RuO2) have attracted considerable attention recently because of their unique physical and chemical properties for many industrial applications. For example, Ru is explored as a diffusion barrier layer and as an electrocatalyst in semiconductor and electrochemical devices, respectively [1,2]. In addition, RuO2 is widely studied as a promising pseudocapacitive material [3]. Conventional synthetic routes for Ru and RuO2 involve vacuum-based and solution-based approaches. The vacuum-based technique includes physical vapor deposition [4], chemical vapor deposition [5,6], and atomic layer deposition [7–9]. Unfortunately, these processes often require expensive equipment and they are known for unnecessary precursor waste. In contrast, the solution-based approach such as sol–gel synthesis [10], electroplating [3,11,12], and electroless deposition [13–19] are credited with simple setups, efficient use of precursors, as well as scalability in outputs. In the solution-based approach, the electroplating technique can be conducted via an externally applied driving force in potentiostatic or galvanostatic mode to direct deposits on conductive substrates. In contrast, the electroless deposition technique can be applied on both conductive and non-conductive platforms as reducing agents in the plating bath are responsible for reducing ions on the substrates instead.

In principle, the electroless deposition approach involves an autocatalytic reaction to form deposits via a heterogeneous nucleation and growth [20]. Notable advantages for the electroless deposition technique are wide substrate selection, sophisticated shape tolerance, and free of externally applied driving force. To date, electroless deposition of metals including Ni, Cu, Ag, Au, Pd, and Pt have been demonstrated in various formulations and their formation mechanisms are well-established [21–25]. On the other hand, metals such as Ru, Rh, Re, Os, and Ir have received few attention and functional electroplating baths still require further investigation [26]. In the case of Ru, preliminary experiments have been carried out on substrates including Cu, C, Pd-InGaAs, Si, and acrylonitrile butadiene styrene [13–19]. It is realized that for non-conductive substrates, selective surface pretreatments are rather necessary to ensure a uniform film formation. Earlier, Chang et al. attempted Ru electroless deposition on Si substrates but surface pretreatments were not discussed [15]. Recently, we have developed an oxidative-reductive Ru electroless plating bath and demonstrated its applicability on a Cu substrate [18,19]. Since Cu is known as a conductive surface for electroless plating, it becomes our interest to explore identical formulation on non-conductive surface such as Si. Preliminary study using as-received Si substrates concluded that a dense Ru film was unable to form. Hence, in this work, we investigate the effect of surface pretreatments on Si substrates for the formation of continuous Ru film via an electroless route.

2. Experimental details

A 6-inch boron doped p-type single crystalline Si wafer with (100) orientation was broken into small pieces (2 × 2 cm2) and they were used as the substrates for Ru electroless deposition. The resistance and thickness for the Si wafer were 1–30 Ω cm and 625 ± 25 μm, respectively. Prior to surface pretreatment, the Si samples were rinsed...
with deionized water and acetone to remove any debris and possible contaminant. Afterward, it was immersed in a hydrogen fluoride (HF) solution at 25 °C for 10 min. The HF solution contained HF (J.T. Baker; Buffered Oxide Etch 6:1) and H2O2 (SHOWA; 35 wt.%) at a 1:50 volume ratio. Subsequently, the Si samples were further subjected to two different steps including activation, as well as sensitization and activation, respectively. The activation step involved immersing the Si substrate in a mixture of 0.1 wt.% PdCl2 and 1 wt.% HCl in deionized water at 40 °C for 10 min. The sensitization step engaged the Si substrate in a solution containing 0.3 wt.% SnCl2 and 2.5 wt.% HCl in deionized water at 25 °C for 3 min. Upon completion of surface pretreatments, the samples were submerged in a Ru electroless plating bath at 40 °C for different times. The chemical ingredients in the plating bath contain K2RuCl5, H2O, NaClO, NaOH, and NaNO2, at appropriate ratios, and the relevant oxidative–reductive mechanism for Ru film formation has been reported elsewhere [18].

A field-emission scanning electron microscope (FE-SEM; JEOL-JSM-6500F) with 15 keV operating voltage was used to observe morphology and thickness for the deposited films. We took the cross-sectional SEM images and selected three different locations to estimate their respective heights. An energy dispersive spectrometer (EDS; Oxford 7557) was utilized for elements detection. The acceleration voltage was 15 keV and the acquisition time was 70 s for the EDS measurement. A contact angle set-up consisting of lens, light source, and a power supply was adopted to determine hydrophilicity of Si substrates before and after surface pretreatments. An atomic force microscope (AFM; Vecco Dimension 5000 Scanning Probe Microscopy) was involved to determine roughness for the deposited films. Tapping mode was utilized for AFM measurement for 1×1 μm2 scanning area. An Auger electron spectrometer (AES, Thermo Microlab 350) was employed to perform depth profiling for the deposited films. The depth profile was divided into 100 layers with Ar ions bombardment at 5 s per layer. An X-ray photoelectron spectrometer (XPS; Thermo Microlab 350) with Mg target was used to identify oxidation states of Sn and Pd on Si substrates after surface pretreatments. For the XPS measurements, additional Pt particles were sputtered on the Si surface serving as a standard for correction with metallic Pt binding energy at 4f7/2 at 71.2 eV. The Si surface was bombarded with Ar ions which were operated with 3 kV and 1 μA for 30 s before recording the XPS.

3. Results and discussion

Table 1 provides the contact angles for Si samples before and after surface pretreatments, as well as the roughness before and after Ru electroless depositions. As listed, the removal of native oxide from the as-received Si wafer after HF etching rendered a relatively hydrophobic surface. However, after further pretreatments of activation or sensitization and activation, the Si surfaces were transformed to hydrophilic again. The roughness values were consistent with what we expected as activation steps produced Pd nuclei that led to a larger roughness especially for the sample undergoing sensitization and activation steps. In addition, after Ru electroless deposition, more Pd nuclei on the Si surface resulted in a larger thickness variation for the deposited Ru.

Table 1

<table>
<thead>
<tr>
<th>Contact angle (°)</th>
<th>Roughness (nm)</th>
<th>Before Ru deposition</th>
<th>After Ru deposition</th>
</tr>
</thead>
<tbody>
<tr>
<td>As-received Si substrate</td>
<td>37.97</td>
<td>0.2</td>
<td>N/A</td>
</tr>
<tr>
<td>HF etching</td>
<td>79.75</td>
<td>0.3</td>
<td>3.72</td>
</tr>
<tr>
<td>HF etching + activation</td>
<td>17.08</td>
<td>3.6</td>
<td>11.61</td>
</tr>
<tr>
<td>HF etching + sensitization + activation</td>
<td>12.84</td>
<td>4.3</td>
<td>11.79</td>
</tr>
</tbody>
</table>

Fig. 1 demonstrates the XPS profiles for the Si samples after surface pretreatments. As shown in Fig. 1(a), the Pd 3d5/2 and 3d3/2 were identified at 334.4 and 339.9 eV, respectively for the Si substrate after activation pretreatment. These values agree well with what was reported earlier for metallic Pd 3d lines [27]. Hence, we realize that the activation step is effective in distributing Pd nuclei on the Si surface for subsequent Ru deposition. Fig. 1(b) and (c) exhibits the Sn 3d and Pd 3d lines after sensitization and activation pretreatments, respectively. As shown in Fig. 1(b), the profile revealed notable signals at 484.5 and 493.4 eV, which are consistent with metallic Sn from literature [27]. In addition, there were presence of Pd 3d lines at 335.4 and 341.1 eV, and these values are slightly larger than those in Fig. 1(a), indicating that moderate Pd oxidation to PdO was possibly occurring [28,29]. The shoulder at 332 eV in Fig. 1(c) is from Pt 3d3/2 which is...
attributed to the sputtered Pt prior to the XPS measurements. Our results suggest that metallic Sn and oxidized Pd are formed on the Si surface after sensitization and activation treatments.

In order to confirm the formation of Ru films on the Si sample, the EDS was employed after 15 min of plating time. The estimated Ru amounts for activation as well as sensitization and activation were 1.06 and 0.93 at.%, respectively. In contrast, the Si sample after only HF etching was merely 0.81 at.%. Earlier, XPS results on the plated films atop Cu substrates indicated a composition makeup of 92.5 at.% Ru and 7.5% RuO₂ [19]. Since we used identical plating formulation to carry out electroless deposition in this study, we believe that the resulting composition was not notably altered from previous work. The acceleration voltage for the EDS is 15 keV, which is in line with what are typically used in similar measurements. Hence, the signal depth for element detection in EDS is likely to be several micrometers. Considering that the thickness for the plated Ru films was only in sub-micron regime (<100 nm), it is not unexpected that the recorded Ru amount in terms of atomic percentage was largely subdued.

Fig. 2 presents the SEM images of Ru films after 40 min of electroless plating. As shown in Fig. 2(a), the as-received Si wafer revealed Ru deposit that was partially formed. Previously, we purchased wafers with 500 nm SiO₂ on surface and determined that with or without surface treatments (HF etching, hydrophilic silane treatment, etc.), the Ru films were unable to deposit. This is because the SiO₂ is known to be unstable in alkaline solution and the pH value for the Ru plating bath is 12–13. Thus, nucleation and growth of stable nuclei are unable to proceed as the underlying SiO₂ keeps dissolving during electroless plating. Since the Si wafer always has its native SiO₂ on its surface (~3 nm), surface pretreatment is thus necessary to render it more susceptible to Ru deposition. As shown in Fig. 2(b), Si sample after HF etching revealed a relatively flat Ru film with scattered particles atop. In contrast, for Si samples after HF etching and activation, as well as HF etching, sensitization, and activation shown in Fig. 2(c) and (d), respectively, the Ru films demonstrated granular morphologies with notable thickness variations. These results are not unexpected since the presence of nuclei promotes heterogeneous island growths that coalesce into a solid film eventually. This mechanism is substantiated by the apparent voids observed in Fig. 2(c).

Fig. 3 provides the average thickness and standard deviation for the Ru films on Si samples after surface pretreatments. Apparently, a linear relationship between the deposition time and film thickness was observed for all samples. To accurately estimate the growth rate, the first data point at 15 min was excluded. As a result, the remaining data points lead to $y = 27.44 + 1.07x$, $R^2 = 0.94$ for HF etching, $y = 25.35 + 1.53x$, $R^2 = 0.98$ for HF etching and activation, and $y = 17.03 + 1.85x$, $R^2 = 0.99$ for HF etching, sensitization, and activation. Obviously, the slower growth rate observed for the HF etching sample is attributed to the lack of nucleation sites for Ru electroless deposition. An alternative possibility is the relatively hydrophobic surface that discourages wetting of Ru plating bath. For samples undergoing additional sensitization and activation steps, there appeared a slightly faster growth rate as compared to those of activation step, which is rationalized by its excessive Sn and Pd nuclei on the Si surface. However, we must admit that the effect of sensitization seems to be rather moderate.

Fig. 4 presents the AES depth profile for the Ru film after 30 min of electroless plating on the Si sample undergoing HF etching, as well as sensitization and activation. The atomic percentage for Ru was clearly observed when the etching time was varied from 0 to 330 s. Assuming the etching rate of standard SiO₂ (2 Å s⁻¹) is identical to that of Ru, the thickness for the Ru film is estimated at 66 nm. This value is reasonably close to what was observed in SEM cross-sectional view at 71.9 ± 9.5 nm. As shown in the depth profile, from 0 to 25 s the atomic percent for O was slightly larger than that for Ru, suggesting that the surface for the Ru film was partially oxidized becoming Ru oxide or hydroxide instead. This behavior is not unusual as the Ru is...
known to form RuO$_x$ and Ru(OH)$_x$ when it is exposed to atmosphere or electrolyte. When the etching time was between 25 and 145 s, the atomic percentage for Ru and O remained relatively unchanged at 40–50%, inferring that the composition for the Ru film was rather uniform. According to our previous work [19], the electroless Ru plating bath produces a composite Ru/RuO$_2$ film and the exact Ru/RuO$_2$ makeup is contingent on the processing parameters involved. Therefore, it is likely to have this amount of oxygen throughout the deposited film. Lastly, when the etching time was between 145 and 330 s, the atomic percentage for Si became larger than those of Ru and O. This behavior was caused primarily by surface roughness at the interface between Si and Ru that renders both signals to be detected simultaneously. Another remote possibility is the formation of sili-
cides such as RuSi or Ru$_2$Si$_3$ because the binary phase diagram of Ru and Si indicates so when the Ru/Si ratio is larger than 1 [30]. However, since our samples were not subjected to any heat treatment after the electroless depositions, the formation of intermetallic ruthenium silicide was very unlikely.

4. Conclusions

We explored electroless Ru depositions on Si substrates undergoing surface pretreatments including HF etching, sensitization, and activation. Contact angle measurements indicated a relatively hydrophilic surface after sensitization and activation, which led to faster Ru film growths and larger surface roughness as compared to the HF-etched sample. XPS confirmed the formation of Sn and Pd nuclei and their presence facilitated the heterogeneous growth of Ru films. Results from AES depth profiling suggested a uniform composition across the film thickness despite part of the Ru was present in an oxidized form. We determined that without surface pretreatment, Ru film was not able to form. In contrast, after HF etching, sensitization, and activation, dense Ru films were obtained at reasonable growth rates.

Acknowledgments

Financial support from the National Science Council of Taiwan (100-ET-E-009-002-ET) is noted.

References