Surface Texturing Behavior of Nano-copper Particles under Various Copper Salts System during Copper-assisted Chemical Etching

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Surface Texturing Behavior of Nano-copper Particles under Various Copper Salts System during Copper-assisted Chemical Etching

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Abstract: In this work, the effects of different copper salts on the etching behavior of n-type monocrystalline silicon wafers were detailedly studied by Cu-assisted chemical etching method. Firstly, the inverted pyramid, inverted pyramid-like and oval pit texturing structures were obtained by HF/H$_2$O$_2$/Cu(NO$_3$)$_2$, HF/H$_2$O$_2$/CuSO$_4$ and HF/H$_2$O$_2$/CuCl$_2$ etching systems. Then, the evolution of copper particles deposition behavior was studied to reveal the influencing mechanism of different anion species, the textured wafer surfaces were characterized by scanning electron microscopy (SEM) and ultraviolet-visible (UV) spectrophotometer, the etching rate, silicon wafer thinning and the deposition amount of copper particle was systematically analyzed. We conclude that the binding force between anion and cation, the oxidation of anions and the formation of complex groups [CuCl$_2$] lead to great difference in the deposition behavior of copper, resulting in different etching morphology and etching rate. The moderate size copper particles deposited from HF/H$_2$O$_2$/Cu(NO$_3$)$_2$ system make that the etching process is mild and the anisotropic etching ability can fully demonstrated, and the regular inverted pyramid structures can be formed under low thinning of silicon wafers. This work will provide guidance for controllable preparation of inverted pyramid structure and future application in high efficiency solar cells.

Key words: Cu-assisted chemical etching; etching behavior; inverted pyramid structure; copper deposition; anions ions effect
1. Introduction

Solar energy has become one of the most promising renewable energy sources to replace traditional energy sources because of its clean and pollution-free reserves [1,2], and the installed capacity of photovoltaics has increased dramatically in recent years[3]. The solar cells, which can directly convert sunlight into electrical energy, are undoubtedly the core device of photovoltaic power generation, where the single crystal silicon (sc-Si) solar cell occupied more and more market share due to the continuous technological progress and cost reduction [4,5]. Moreover, the n type silicon show huge potential in developing low-cost and high-efficiency solar cells in the future due to its natural advantages of low light attenuation, high minority carrier lifetime [6,7]. Surface texture is an important optical management strategy for high-efficient silicon solar cell devices[8]. During sc-Si solar cell processing, KOH anisotropic etching is utilized to texture silicon wafer, and the upright pyramid light trapping structure can be formed, which reduces the reflectivity of silicon wafer to about 15% [9], however, the upright pyramid is not the most effective anti-reflection structure and it is not an optimal choice in prospective HIT high efficiency cell technology which needs a efficient interfaces passivation.

Metal assisted chemical etching is an effective wet texturing method which could reduce the surface reflectivity by forming highly light-trapping nanostructures[10-12]. In previous work, we have prepared vertical, inclined and zigzag nanowire structures on multi-crystalline silicon (mc-Si) substrates by Ag-assisted chemical etching [13-15]. Besides silver, the copper also show catalytic activity, and inverted pyramids (IPs) structure with ultra-low reflectivity can be obtained by anisotropic etching of copper [16-20]. The IPs structure owns superior light-trapping performance and attracted global attention since 2001, when the research team of the University of New South Wales applied IPs structure to Passivated Emitter and Rear Locally-diffused (PERL) solar cell, which created a efficiency record of 24.7% [21]. However, this structure has not been quickly applied to photovoltaic devices on a large scale, because it needs lithography and laser processes, which were complicated and expensive. Until 2015, Wang Yan et al obtained IPs structure with low reflectivity on the silicon surface by copper-assisted chemical etching using HF/H₂O₂/Cu(NO₃)₂ solution, which brought hope for the low-cost preparation of the IPs structure [22]. We also
fabricated IPs structure with low reflectivity on mc-Si surface [23], moreover, we found that the etched morphologies in different copper salt etchants are quite different [24]. Studying the influence mechanism of anions in different copper salt etching systems will help us further understand the chemical etching mechanism of copper nanoparticle and the formation process of the inverted pyramid.

In this paper, we applied Cu-assisted chemical etching (Cu-ACE) on n-type monocrystalline silicon wafers, and the different copper salt etchant system of HF/H$_2$O$_2$/Cu(NO$_3$)$_2$, HF/H$_2$O$_2$/CuSO$_4$ and HF/H$_2$O$_2$/CuCl$_2$ was selected to explore the influence of anions species on the deposition behavior of Cu, etching rates and resulting surface morphologies. Finally, uniform IPs arrays with low reflectivity of 7.2% was prepared on silicon surface, and the etching rate was slow. Our work will contribute to a better understanding of the etching mechanism of Cu-ACE and the formation process of inverted pyramid structure.

2. Experiment process

The diamond wire sawed (100)-oriented n type crystalline silicon wafers with thickness of 156 ± 5 μm and resistivity of 1~3 Ω · cm were used. Before the experiment, the wafers were cleaned in acetone, ethanol and deionized water for 10 mins under sonication, and then immersed in 10% HF solution for 10 mins in order to remove native SiO$_2$. Subsequently, the wafers were divided into three groups different mixture. First group (3 pieces) were placed in HF/H$_2$O$_2$/Cu(NO$_3$)$_2$ (labeled NO$_3$- system), HF/H$_2$O$_2$/CuSO$_4$ (labeled SO$_4^{2-}$ system) and HF/H$_2$O$_2$/CuCl$_2$ (labeled Cl$^-$ system) solution for 7 mins, so as to analyze the difference of etching morphology, etching rate and the resulting reflectivity. Second group (12 pieces) was placed in HF/Cu(NO$_3$)$_2$, HF/CuSO$_4$ and HF/CuCl$_2$ deposition system for 5 s, 10 s, 30 s and 60 s in every system, so as to observe the evolution of copper deposition state on wafers in the absence of H$_2$O$_2$. Third group (12 pieces) was etched by the same three etching systems as the first group, and etching times were 1, 3, 5 and 7 mins in every system, in order to observe the morphology evolution of etching structures and deposited Cu nanoparticles (Cu-NPs). The detailed etching parameters were shown in tables 1 and 2. After etching, samples of group 1 were immersed into HNO$_3$ solution to eliminate residual Cu-NPs. Finally, the all samples were thoroughly rinsed with distilled water and dried in
vacuum drying chamber. Moreover, the mass change of silicon wafer before and after etching were recorded to calculate the etching rate (Ra) based on Eq. (1), the relative amounts of deposited Cu (Q) were obtained by Eq. (2). The morphologies were characterized by scanning electron microscopy (SEM), the surface reflectivity of wafers was measured by the Ocean Optic USB-4100 spectrometer.

Table 1

Etching conditions of n-type monocrystalline silicon wafers by different copper salt system

Table 2

Deposition conditions of n-type monocrystalline silicon wafers by different copper salt system

\[ R_a = \frac{\Delta m}{2 \rho S t} \] (\( \mu \text{m/min} \)) (1)

\[ Q = \frac{(m_3 - m_2)}{m_1} \times 1000 \] (2)

Where \( \Delta m \) is the mass loss of etched wafers (g), \( \rho \) is the mass density of crystalline silicon (g/cm\(^2\)), \( S \) is the silicon surface area (cm\(^2\)), and \( t \) is the etching time (min), \( m_1 \) is the mass of the as raw silicon wafer, \( m_2 \) is the mass of silicon wafer after deposition where the copper is cleaned by nitric acid, and \( m_3 \) is the mass of the wafer plus the deposited copper after deposition

3. Results and discussion

Fig. 1. (a)–(c) SEM images of n-type silicon wafer under different copper salts system; (d)–(f) the etching rate and reflectivity fitting diagram of the wafer under the three selected etching system.

Figures 1a, b, c show the top sectional SEM images of (100)-oriented crystalline silicon etched in NO\(_3^-\) system, SO\(_4^{2-}\) system and Cl\(^-\) system for 7 min at 40 °C. As we can see, the etched silicon surface presents various morphologies when the anion species are different. The Cu-NPs exhibit anisotropic etching in NO\(_3^-\) and SO\(_4^{2-}\) system, where uniform IPs with width of 0.8–1 μm were formed under NO\(_3^-\) system, and variable IP-like structures were obtained under SO\(_4^{2-}\) system. In addition, numerous oval pits with 5 μm length and 1 μm width were formed when wafers were etched in Cl\(^-\) system, this structure is similar to that of HF/HNO\(_3\) texturing method. In terms of reaction rate (see Fig. 1d), the etching rate of NO\(_3^-\) system is the lowest, followed by Cl\(^-\) system, and etching rate of the SO\(_4^{2-}\) system is the fastest. The IPs and IP-like structure owns low reflectance of 7.2% and 11.7%, the oval pits obtained by Cl\(^-\) system etching presents the highest reflectivity of
18.8% (see Figs. 1e, f), which may be due to the large size of the pit which is not conducive to the refraction of light. The results indicated that anions would greatly effect the etching behavior of Cu-NPs and leading to different morphologies and etching rate as well as reflectivity.

**Fig. 2.** SEM diagram of Cu-NPs morphology evolution of wafers treated in (a)–(d) HF/Cu(NO$_3$)$_2$ system, (e)–h HF/CuSO$_4$ system and (i)–(l) HF/CuCl$_2$ system for different time; (m)–(o) Energy Dispersive Spectroscopy (EDS) spot scan results of the deposits in HF/Cu(NO$_3$)$_2$, HF/CuSO$_4$ and HF/CuCl$_2$ systems; (p) fitting diagram of copper deposition rate variation

Figures 2a-l present the SEM images of silicon surface morphology processed in HF/Cu(NO$_3$)$_2$, HF/CuSO$_4$ and HF/CuCl$_2$ system at different time duration. Figures 2m, n, o show the EDS results of the deposited nano-particles when treated for 60 s in NO$_3^-$, SO$_4^{2-}$ and Cl$^-$ deposition systems, respectively. It can be found that numerous Cu-NPs were deposited on silicon wafers surface in HF/Cu(NO$_3$)$_2$, HF/CuSO$_4$ system. While for Cl$^-$ system, the Cu and Cl species was detected and their atomic ratio is nearly 1:1 (see Fig. 2o), indicating CuCl precipitate forms on silicon wafer surface. Figures 2a, b, c, d show the morphology evolution of Cu-NPs in HF/Cu(NO$_3$)$_2$ system, at the first 10 s stage, a large number of Cu$^{2+}$ ions capture electrons and deposit on the silicon surface, as the deposition time increases, Cu-NPs gradually grow up and form bigger round copper particles. In the HF/CuSO$_4$ system (see Figs. 2e, f, g, h), a dense copper film is formed and covered on the silicon surface at the beginning of deposition. Subsequently, some round copper particle appears when the deposition time reaches to 60 s like that of HF/Cu(NO$_3$)$_2$ system. The current results indicate that Cu$^{2+}$ ions can uniformly deposition with small size and form a dense copper film in the HF/CuSO$_4$ system. In HF/CuCl$_2$ system (see Figs. 2i, j, k, l), it is obvious that heavily agglomerated rod-shaped nano-particles appear along the saw marks regions. Figure 2p shows the relative amount of the deposited Cu-NPs under three deposition systems, it can be seen that the amount of deposited Cu-NPs in the HF/CuSO$_4$ system is slightly more than that in the HF/Cu(NO$_3$)$_2$ system at the beginning stage of 30 s, as the reaction proceeds, the amount of Cu-NPs deposited in HF/CuSO$_4$ system becomes much higher than that in HF/Cu(NO$_3$)$_2$ system, by contrast, the amount of Cu-NPs in HF/CuCl$_2$ system is the least.

**Fig. 3.** Diagram of deposition of Cu$^{2+}$ in (a) HF/CuSO$_4$ system, (b) HF/Cu(NO$_3$)$_2$ system, and (c) HF/CuCl$_2$ system

Figure 3 shows the schematic diagram of deposition process of Cu-NPs in different systems.
The deposition behavior of Cu is related to anion concentration and anion properties [24-28]. Although the concentration of Cu$^{2+}$ is the same in the three deposition systems, the anion concentration is different. Compared with NO$_3^-$ and Cl$^-$, the concentration of SO$_4^{2-}$ is the lowest, which leads to the weakest electrostatic attraction between SO$_4^{2-}$ and Cu$^{2+}$, thus the Cu$^{2+}$ ions are easier to get electrons and evenly deposit on silicon surface as shown in Fig. 3a. Moreover, the difference in the surface energy of the silicon wafer has no great influence on the Cu deposition in HF/CuSO$_4$ system, which leads to the Cu-NPs is dispersed and form a dense film. In HF/Cu(NO$_3$)$_2$ system, the higher concentration of anion concentration enhances the attraction between NO$_3^-$ and Cu$^{2+}$ and thus limits the copper ions deposition (see Fig. 3b), which causes the Cu-NPs tend to deposit at the saw marks and defect sites, slightly agglomerated and grow into bigger round particle. With the reaction proceeds, the NO$_3^-$ also oxidize Cu-NPs into Cu$^{2+}$ ions, which controls the size of the copper particles not to be too large, and the deposition rate is slower.

As for HF/CuCl$_2$ system, the concentration of Cl$^-$ is the same as that of NO$_3^-$ in HF/CuCl$_2$ system. But due to the more stable covalent bond of CuCl$_2$, the Cu$^{2+}$ tends to be more difficult to break free and deposition. Moreover, many Cu$^{2+}$ and the deposited Cu-NPs are prone to form [CuCl$_2$]$^-$ complexes in rich Cl$^-$ solution, and further form CuCl precipitate[29], as shown in Fig. 3c, the specific reaction equations are shown in Eqs. (3)-(6) [29,30].

\[
\text{Cu}^{2+} + 2\text{e}^- \rightarrow \text{Cu} \quad (3)
\]

\[
\text{Cu}^{2+} + 2\text{Cl}^- + \text{e}^- \rightarrow [\text{CuCl}_2]^-
\]

\[
\text{Cu}^{2+} + 2\text{Cu} + 4\text{Cl}^- \rightarrow 2[\text{CuCl}_2]
\]

\[
[\text{CuCl}_2]^- \rightarrow \text{CuCl} \downarrow + \text{Cl}^-
\] (6)

Besides, due to the large number of [CuCl$_2$]$^-$ complexes, the migration of ions in the solution is greatly hindered, these factors make the Cu$^{2+}$ really difficult to get electronic and deposit on silicon surface, so that the Cu particles and CuCl precipitate preferentially deposit along the saw marks and defect sites and heavily agglomerated into a rod-shaped structure. It can be seen that the chemical properties of anions greatly affect the deposition of Cu-NPs, which affects the etching behavior during the Cu-ACE.

**Fig. 4.** SEM images of nano-structures evolution of wafers etched in (a)–(d) NO$_3^-$ system, (e)–(h) SO$_4^{2-}$ system and (i)–(l) Cl$^-$ system for different time
Figure 4 presents the evolution process of IPs, IP-like and oval pit structure when silicon wafer was etched in NO$_3^-$, SO$_4^{2-}$ and Cl$^-$ system for 1 min, 3 mins, 5 mins, 7 mins. For the NO$_3^-$ system (see Figs. 4a, b, c, d). As we can see, irregular pits are formed along the saw marks at the beginning stage, then, a small amount of IPs are scattered on the silicon surface, when it’s up to 5 mins, IPs get bigger and closely gather on the surface with wide of about 800~1000 nm, moreover, the saw marks are also removed, until 7 mins, the inverted pyramids merged with each other and the IPs structure are destroyed, which is caused by excessive etching. For the SO$_4^{2-}$ system (see Figs. 4e, f, g, h), numerous pits with different sizes appear along the saw marks at the first 3 mins, when it’s up to 5 mins, the pits grows into big grooves and some IP-like structures are formed, as reaction goes on, more IP-like structures with various size can be seen, which indicates that the anisotropic etching ability of copper in SO$_4^{2-}$ system is not obvious as that in NO$_3^-$ system. As for Cl$^-$ system (see Figs. 4i, j, k, l), crescent-shaped grooves appear at the beginning of the reaction, and the crescent-shaped pits gradually widening and deepening, eventually change into elliptical pits with various lengths, as seen in Fig. 4l.

**Fig. 5.** SEM images of Cu-NPs morphology evolution of wafers etched in (a)–(d) NO$_3^-$ system, (e)–(h) SO$_4^{2-}$ system and (i)–(l) Cl$^-$ system for different time

**Fig. 6.** Diagram of average etching rate at different etching times for different system

To further explore the difference of copper etching behavior in different copper salt systems, the morphology revolution of Cu-NPs in NO$_3^-$, SO$_4^{2-}$ and Cl$^-$ system at different etching time was studied, the SEM pictures are show in Fig. 5. For NO$_3^-$ system (see Figs. 5a, b, c, d), large amount of rounded and scattered Cu-NPs was deposited on the surface at the beginning, with the extension of reaction time, some Cu-NPs were oxidized to Cu$^{2+}$ by H$_2$O$_2$, so the number of Cu-NPs is decreased,. When etching time is 7 mins, Cu-NPs are invisible but some small pits appear at the bottom of the round deep pits where Cu-NPs have been deposited. Figures 5e, f, g, h show the situation of Cu-NPs in SO$_4^{2-}$ system, it can be seen that at the beginning of the reaction, lots of Cu-NPs are deposited on the sidewalls of the pits which is similar to the situation of NO$_3^-$ system, but the size of Cu-NPs is smaller. As the reaction proceeds, the Cu particles become less and smaller. The small pits on the sidewall (see Fig. 5g) can infer that the copper-assisted chemical etching intensely happens. As for Cl$^-$ system (see Figs. 5i, j, k, l), a large amount of Cu and CuCl particles
were stacked in the crescent-shaped groove at 1 min, and the size is much bigger than that in NO₃⁻ and SO₄²⁻ system. These deposits (CuCl particles) becomes bigger as the reaction goes on, in which is contrary to the situation of NO₃⁻ and SO₄²⁻ system.

Different copper etching behaviors naturally lead to different etching rates (see Fig. 6). it is obvious that at the beginning of reaction, the etching rate of SO₄²⁻ system is much higher than the others, and rapidly decreased to 1.52 μm/min. While the etching rate of NO₃⁻ system and Cl⁻ system are very close and both stable at about 1 μm/min.

**Fig. 7. Diagram of etching behavior of Cu/CuCl in (a) SO₄²⁻ system, (b) NO₃⁻ system and (c) Cl⁻ system**

Figure. 7 elaborates the role of Cu to form different structure in the above three copper salt systems. Due to the weak electrostatic attraction, Cu evenly deposited on the silicon surface. These tiny Cu-NPs with high surface activity, lead to intense reaction, but they are quickly oxidized by H₂O₂ (see Fig. 7a), therefore, only a large number of corrosion pits and a few small Cu-NPs lied on the side wall as the reaction goes on, and the corresponding reaction rate rapidly decrease, even so the average etching rate at 7 mins is still much higher than the other two. Due to the intense reaction, the anisotropic etching ability of Cu-NPs cannot be completely demonstrated, so the regular IP structures cannot be formed but many IP-like structures instead. In NO₃⁻ system, H₂O₂ and NO₃⁻ make the Cu-NPs the proper size, which can fully demonstrate an anisotropic etching ability, and eventually form a regular IP structure as shown in Fig. 7b. Moreover, the proper size of Cu-NPs make the reaction mild and the reaction rate is always keep about 1μm/min. For the Cl⁻ system (see Fig. 7c), Cu-NPs and CuCl-NPs will both deposit on the silicon surface, the H₂O₂ can oxidize Cu-NPs, but it has no effect on the CuCl-NPs. Therefore, as the reaction proceeds, a large amount of [CuCl₂]⁻ complex ions changed into CuCl precipitate and agglomerate in the corrosion pits. Moreover, due to the uneven deposition, the resulting structure is not uniform, leading to numerous oval pits with different size, and the cutting lines are still obvious after etching, the slow etching rate of Cl⁻ system is mainly due to the slow deposition rate of Cu²⁺.

**4. Conclusion**

The effects of NO₃⁻, SO₄²⁻ and Cl⁻ on the etching morphology of n-type single crystal silicon in the copper-assisted etching process are firstly systematically analyzed, the different anions affects
the copper deposition process, result in different etching rates and behaviors.

The weak attraction between SO$_4^{2-}$ species and Cu$^{2+}$ ions leads to that Cu$^{2+}$ deposition is quick and easily form tiny Cu-NPs, which causes intense reaction process and result in lots of grooves and inverted pyramid-like structure. In the Cl$^-$ species etching system, the covalent bond of CuCl$_2$ and formation of [CuCl$_2$] complex ions lead to that the Cu$^{2+}$ ions preferentially deposit in defects sites and agglomerate into rod-shaped particles, resulting in oval corrosion pits, which is not conductive to saw marks’ removal.

In comparison, the NO$_3^-$ species show stronger oxidation and which can relieve the Cu-NPs aggregation, these Cu-NPs with proper size show better anisotropic etching ability, thus forming regular inverted pyramid structure which can almost completely disappeared saw marks and lower reflection to 7.2%. It can be seen that the type of anion does affect the etching behavior of Cu-NP to form different morphologies, through this work we can further understands the formation mechanism of inverted pyramid by HF/H$_2$O$_2$/Cu(NO$_3$)$_2$ system, moreover, it provides guidance for controllable preparation of inverted pyramid structure and future high efficiency single crystal solar cells.

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**Compliance with Ethical Standards:**

**Conflicts of interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

**Ethics approval**

The authors declare that the manuscript is not currently being considered for publication in another journal.

**Consent to participate**
I testify on behalf of all co-authors that our article submitted to Silicon.

Consent for Publication

On behalf of all the authors, I agree that our article should be published in Silicon

Availability of data and material

The authors declare this data and material has not been published in whole or in part elsewhere.

Code availability

Not applicable

Authors' contributions

All authors have been personally and actively involved in substantive work leading to the manuscript, and will hold themselves jointly and individually responsible for its content.

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Figures

Figure 1

(a)–(c) SEM images of n-type silicon wafer under different copper salts system; (d)–(f) the etching rate and reflectivity fitting diagram of the wafer under the three selected etching system.

Figure 2
SEM diagram of Cu-NPs morphology evolution of wafers treated in (a)~(d) HF/Cu(NO₃)₂ system, (e)~(h) HF/CuSO₄ system and (i)~(l) HF/CuCl₂ system for different time; (m)~(o) Energy Dispersive Spectroscopy (EDS) spot scan results of the deposits in HF/Cu(NO₃)₂ and HF/CuSO₄ and HF/CuCl₂ systems; (p) fitting diagram of copper deposition rate variation.

**Figure 3**

Diagram of deposition of Cu²⁺ in (a) HF/CuSO₄ system, (b) HF/Cu(NO₃)₂ system, and (c) HF/CuCl₂ system.

**Figure 4**

SEM images of nano-structures evolution of wafers etched in (a)~(d) NO₃- system, (e)~(h) SO₄²⁻ system and (i)~(l) Cl⁻ system for different time.
Figure 5

SEM images of Cu-NPs morphology evolution of wafers etched in (a)∼(d) NO3- system, (e)∼(h) SO42- system and (i)∼(l) Cl- system for different time.

Figure 6

Diagram of average etching rate at different etching times for different system

Figure 7
Diagram of etching behavior of Cu/CuCl in (a) SO42- system, (b) NO3- system and (c) Cl- system