Evolution of Morphology of Nano-Scale CuO Grown on Copper Metal Sheets in 5 wt% NaCl Solution of Spray Fog Environment

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ABSTRACT
Nano-scale copper oxide with various morphologies is synthesized via the thermal oxide method and growth in a 5 wt% NaCl solution of spray fog environment. The nano-scale copper oxide is grown on copper metal sheets via the thermal oxide method at 650°C for 60 minutes. Nano-scale copper oxide grains and nanowires are induced on copper metal sheets then placed in 5 wt% NaCl solution of salt spray fog environment. Significant changes in particle size and morphology are observed with increasing salt spray fog treatment time. The morphology of nano-scale copper oxide varies from nanograins to nanowires, tetrahedron, and icositetrahedron. The morphologies and structures of the obtained nano-scale copper oxide are investigated by scanning electron microscopy and energy-dispersive spectroscopy. Possible growth mechanisms are discussed.

Keywords: Nano; CuO; Morphology; NaCl Solution; Spray Fog

1. Introduction
Nanoscale one-dimensional (1D) and two-dimensional (2D) materials such as nanowires, nanorods, nanoribbons, and nanosheets have attracted a lot of interest due to their unique physical properties and potential applications in nanodevices.

Copper oxide (CuO) is a well known p-type semiconductor and a photovoltaic material. It has a narrow band gap (1.2 eV). CuO is desirable due to its availability and abundance of the starting materials, non-toxicity, and low production cost [1]. Consequently, it is widely used in applications such as catalysis [2], sensors [3], solar cells [4,5], and field-emission emitter materials [6]. However, the chemical and physical properties of CuO strictly depend on its size and morphology [7]. Many studies have synthesized various CuO nanostructures. A number of methods have been used to synthesize and control the size and morphology of nano-scale CuO, such as thermal evaporation [1], solution synthesis [7], thermal oxidation [8], sol-gel [9], hydrothermal method [10], electrospinning [11] and microwave hydrothermal synthesis [12].

These methods have been used to synthesize CuO nanomaterials with various morphologies, such as nanoparticles, 1D nanowires, nanorods, and nanobelts; 2D nanosheets, nanoleaves, and nanowiskers; and three-dimensional peachstone-like, boat-like, and ellipsoid-like nanostructures [10]. The morphology of nano-scale CuO is related to growth environment conditions, in addition to processing methods.

In this work, nano-scale CuO with various morphologies is synthesized via the thermal oxide method and growth in a 5 wt% NaCl solution of spray fog environment. The morphology of nano-scale CuO varied from nanograins to nanosheets and nanoneedles with increasing salt spray fog treatment time. Possible growth mechanisms are discussed.

2. Experimental Procedure
CuO with various morphologies was synthesized via the thermal oxide method and growth in a 5 wt% NaCl solution of spray fog environment.

Copper (99.99 wt% Cu) substrate samples with dimensions of 10 mm (width) × 10 mm (length) × 1 mm (thickness) were prepared according to the following procedure: the substrate was polished with various grinding papers (400, 600, 800, 1200, 2000, and 4000 grade) and then ultrasonically cleaned with acetone, ethanol, and de-ionized water for 10 minutes sequentially to remove the impurities and native oxides on its surface. The samples were then dried by blowing with high-pressure nitrogen gas.

Thermal oxidation of the Cu sheets was carried out in...
Evolution of Morphology of Nano-Scale CuO Grown on Copper Metal Sheets in 5 wt% NaCl Solution of Spray Fog Environment

a resistance-heated tube furnace in a pure oxygen atmosphere at a temperature of 650°C for 60 minutes. Nano-scale CuO grains were induced on the copper metal sheets and then placed in 5 wt% NaCl solution of spray fog environment. The treatment times of the samples in the spray fog environment were 1, 7, 14, 21, and 28 days, respectively. The temperature of the spray fog environment was kept in 45°C.

The surface morphology of the samples was observed using a scanning electron microscope (SEM, Hitachi S3000N, Japan) operated at 30 kV. The chemical composition of the nanomaterials was confirmed by energy-dispersive X-ray analysis (EDX, Horiba, Japan). The phases and structure of the nanomaterials were identified by grazing-incidence X-ray diffraction (GIXRD) using a Rigaku D/MAX 2500 multipurpose X-ray thin-film diffractometer with monochromatic high-intensity CuKα radiation (λ = 0.15418 nm).

3. Results and Discussion

3.1. Phase and Structure Analysis

The X-ray diffraction patterns of polycrystalline Cu metal sheets prepared by polishing and thermal oxidation are shown in Figure 1. Figure 1(a) shows the diffraction peaks of an as-polished Cu metal sheet, which are assigned to the (111), (200) and (220), planes of face-centered cubic (fcc) Cu. This result is in agreement with the standard data from JCPDS card No. 85-1326. The XRD patterns of CuO samples on Cu metal sheets prepared via thermal oxidation are shown in Figure 1(b). The XRD peaks can be indexed as monoclinic structured CuO (JCPDS card No. 80-1917), cubic structured cuprous oxide (Cu2O) (JCPDS card No. 78-2076), and fcc Cu metal from the substrate (JCPDS card No. 85-1326). The diffraction peaks of the monoclinic structured CuO are assigned to the (110), (–111), (111), (20–2), (11–3), (31–3), (400), and (22–3) planes. The peaks of cubic structured Cu2O are assigned to the (111) and (200) planes.

Figure 2 shows the XRD patterns of nano-scale CuO prepared via the thermal oxide method and then placed in a 5 wt% NaCl solution of spray fog environment for 0 (only thermal oxidation), 1, and 28 days, respectively. The diffraction pattern of the copper(II) chloride (CuCl2) appeared for the sample prepared via thermal oxidation and then placed in a 5 wt% NaCl solution of spray fog environment for 1 day, in addition to the original production peaks of CuO, Cu2O, and Cu metal substrate (Figure 2(b)). The diffraction peak of the monoclinic structured CuCl2 is assigned to the (–405) planes (JCPDS card No. 79-1635). The peaks of CuCl2 increased with increasing salt spray fog treatment time. Figure 2(c) shows that the diffraction planes of CuCl2 were (002), (111), (112), (–313), and (–221) for the sample treated using 5 wt% NaCl solution of spray fog for 28 days.

The XRD results indicate that the main phases of the nano-scale CuO grown on pure Cu metal substrates after thermal oxidation were CuO and Cu2O. CuCl2 was induced on pure Cu metal substrates when the samples were placed in 5 wt% NaCl solution of spray fog environment.

3.2. Morphology Observation

Figure 3 shows the surface morphology of polycrystalline Cu metal sheets prepared by polishing. The surface of the Cu metal sheets did not have any surface chemical products.

SEM micrographs of nano-scale CuO and nanowires
Evolution of Morphology of Nano-Scale CuO Grown on Copper Metal Sheets in 5 wt% NaCl Solution of Spray Fog Environment

Figure 3. Surface morphology of polycrystalline Cu metal sheet prepared by polishing.

grown via thermal oxidation on Cu metal sheets for 60 minutes at 650°C are shown in Figure 4(a). The morphologies of nano-scale surface products formed after thermal oxidation followed by exposure to 5 wt% NaCl solution of spray fog environment for various durations significantly change, as shown in Figures 4(b)-(f). Figure 4(a) shows Ctahdral-shaped grains about 1 μm aggregated on the nanowire after 1 day in a 5 wt% NaCl solution of spray fog environment. The Ctahdral-shaped grain size increased with increasing spray fog treatment time. The Ctahdral-shaped grains are replaced by a six-horn-shaped icositetrahedron. Each crystal has six apexes, each of which is connected to four sides to form an icositetrahedron [13]. Icositetrahedron-shaped crystals can be observed in the Figure 4(d). When the salt spray fog treatment time was increased to 28 days, the edges and corners of the icositetrahedron-shaped crystals become more extrusive (Figure 4(f)). The crystal size increased from about 1 μm (1 day) to about 8 μm (14 days) and then decreased to about 4 μm (28 days).

The chemical composition of the nano-scale oxide grown on pure Cu metal substrates after thermal oxidation and salt spray fog treatment was confirmed by EDX. Figure 5 shows the nanowire crystal after thermal oxidation that is CuO. The nano-scale oxide grown in a salt spray fog environment, which formed Ctahdral-shaped and icositetrahedron-shaped crystals, was Cu2O. The crystal shape similar to a tetrahedron was CuCl2, as shown in Figure 6.

3.3. Growth Mechanisms of Nano-Scale Copper Oxides

There are two major growth models, namely vapor-liquid-solid (VLS) catalyst-assisted growth [8] and vapor-solid (VS) growth [14], for describing the growth of 1D nanostructures. Kaur et al. [8] synthesized CuO nanowires via the thermal oxidation of copper foils and found that the growth of the nanowires is initiated by the condensation of the vapor of the material at the tip of the droplet in the VLS process. The morphology of a 1D structure has a round droplet, which is usually found near the tip of the wire. However, the CuO nanowires prepared via thermal oxidation had no droplets at their tips, indicating that the VLS mechanism is not applicable.

The oxidation process of the Cu metal substrate includes two steps [15]:

\[
\begin{align*}
4Cu + O_2 &\rightarrow 2Cu_2O \\
2Cu_2O + O_2 &\rightarrow 4CuO
\end{align*}
\]

In this work, the nano-scale Cu grains and nanowires were induced on copper metal sheets by thermal oxidation. The Cu atoms or ions diffused to the surface and then reacted with oxygen ions that formed CuO in the initial stage. When thickness of CuO film increased with increasing oxidation time, Cu atoms or ions difficult diffusion to reactive with surface O2. Consequently, the CuO, Cu2O, and Cu phases were obtained in the samples.

The following mechanism has been proposed for the oxidation and reduction processes of copper in NaCl solution [16]:

\[
\begin{align*}
Cu + Cl^- &\rightarrow CuCl\text{(ads)} + e^- \\
CuCl\text{(ads)} &\rightarrow CuCl\text{(film)} \\
CuCl + e^- &\rightarrow Cu + Cl^- \\
CuCl\text{(ads)} + Cl^- &\rightarrow CuCl_2 \\
CuCl_2^- &\rightarrow Cu^{2+} + 2Cl^- + e^-
\end{align*}
\]

The reactions in Equations (3)-(5) represent the dissolving, chelating, and film-forming processes of copper, respectively. The reactions in Equations (6) and (7) are a competition of the growth and dissolution of the corrosion products on the surface of the specimens.

Nano-scale CuO and Cu2O were grown on the Cu metal substrate and then placed in 5 wt% NaCl solution of spray fog environment. The morphologies of nano-scale CuO variation were dominated by an electrochemical mechanism. Zhao et al. [13] synthesized Cu2O microcrystals with various shapes using an electrochemical method and found that the Cu2O crystal shapes changed from Ctahdral to icositetrahedral with increasing solution PH value. The cations in the solution have little influence on the morphology of copper, but the anions have a complex influence. Hu et al. [17] found that Cl can form in certain local regions of copper. Ctahdron, nanowire, and prism structures are obtained besides cubic copper when the concentration of Cl reaches a certain value. The Cl adsorption on special crystal faces may
Evolution of Morphology of Nano-Scale CuO Grown on Copper Metal Sheets in 5 wt% NaCl Solution of Spray Fog Environment

4. Conclusions

Nano-scale CuO grains and nanowires were induced on copper metal sheets and then placed in 5 wt% NaCl solution of spray fog environment. The following conclusions were obtained:

- Nano-scale copper oxide likely developed as CuO nanograins and CuO nanowires, and then Ctahe-dral-shaped Cu₂O and tetrahedral-shaped CuCl₂ ag-

cause some crystal faces with a faster growth rate to gradually shrink or even disappear with the growth of crystals. This reasonably explains why the crystal size increased from about 1 μm (1 day) to about 8 μm (14 days) and then decreased to about 5 μm (28 days). The diffraction plane (−405) of CuCl₂ disappeared when the samples were treated using 5 wt% NaCl solution of spray fog for 28 days.

Figure 4. Surface morphology of polycrystalline Cu metal sheet prepared by thermal oxide method followed by exposure to 5 wt% NaCl solution of spray fog environment for (a) 0 (only thermal oxidation); (b) 1; (c) 7; (d) 14; (e) 21; and (f) 28 days.
Evolution of Morphology of Nano-Scale CuO Grown on Copper Metal Sheets in 5 wt% NaCl Solution of Spray Fog Environment

CuO nanowires prepared via thermal oxidation regarding likely follow the VS mechanism.

Nano-scale CuO and Cu2O were grown on a Cu metal substrate and then placed in a 5 wt% NaCl solution of spray fog environment. The morphology of nano-scale copper oxide was dominated by an electrochemical mechanism.

The Cl− adsorption on special crystal faces may cause some crystal faces with a faster growth rate to gradually shrink or even disappear with the growth of crystals.

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