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Voltage-induced material removal mechanism of copper for electrochemical-mechanical polishing applications

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Abstract: The current—voltage (I-V) curves, such as linear sweep voltammetry (LSV) and cyclic voltammetry (CV), were employed to evaluate the effect of electrolyte concentration on the electrochemical reaction trend. From the I-V curve, the electrochemical states of active, passive, transient and trans-passive region could be characterized. And then, the mechanism of the process of voltage-induced material removal in electrochemical mechanical polishing (ECMP) of copper was investigated. Scanning electron microscopy (SEM) and energy dispersive spectroscopy (EDS) analyses were used to observe the surface profile. Finally, the oxidation and reduction processes of the Cu surface were monitored by the repetition of anodic and cathodic potential from cyclic voltammetry (CV) method in acid- and alkali-based electrolyte.

Key words: electrochemical mechanical polishing; linear sweep voltammetry; cyclic voltammetry; HNO₃; KNO₃; electrolyte

1 Introduction

The chemical mechanical polishing (CMP) process has been widely used to obtain the global planarization of inter-metal dielectric (IMD) layers, inter-layer dielectric layers (ILD) and pre-metal dielectric (PMD) layers[1-2]. Also, the CMP is the most commonly used planarization technique in semiconductor process for ultra-large scale integrated circuit (ULSI) applications [3–4]. As its name indicates, the CMP process depends on the chemical interaction of the slurry with polishing wafer and mechanical down force applied to the wafer. Recently, the application of CMP has been especially popular in the fabrication of copper (Cu) damascene structures and multi level interconnection process[5-7]. However, low-k materials at 65 nm and below device structures, because of fragile property, require low down-force mechanical polishing for maintaining the structural integrity of under layer during their fabrication[8]. Also, the problems faced by Cu-CMP process are the lower removal rate due to the low mechanical down force required by the low-k dielectric, and to reduce dishing and erosion. To overcome these problems, the electrochemical mechanical polishing (ECMP) technology, a new planarization technology that uses electrolyte chemistry instead of abrasive slurry, was developed[8–11]. In this paper, the current—voltage (I-V) curves were employed to evaluate the effect of electrolyte concentration on the electrochemical surface reaction of Cu electrode. From this I-V curve, the electrochemical states of active, passive and transpassive region could be characterized[12–13]. It is difficult to distinguish between adsorption and desorption (or anodic and cathodic) processes. And then, we investigated how this chemical electrolytes affect the process of voltage induced material removal in Cu-ECMP. To selectively probe the electrochemical effect of ECMP, all these measurements were performed in the absence of mechanical polishing.

2 Experimental

In brief, Cu disks of 2 cm \times 2 cm size with a purity of 99.99% were used as an experimental species in a standard two-electrode glass cell containing a Pt counter electrode (CE) of mesh type and a Cu working electrode (WE) as shown in Fig.1. Electrical connection to the sample was made through a modified alligator clip at the upper part of the Cu disk, while the sample was electro-

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chemically controlled by a DC power supply system, interfaced with a personal computer as shown in Fig.1. All experiments were performed at room temperature, in the absence of the mechanical polishing. De-ionized water and reagent grade chemicals were used to prepare the electrolytes of KNO₃ and HNO₃, with different concentration levels as follows: 5%, 10% and 20% (mass fraction), respectively. Keithley 236 model I - Vmeasurement system, which consists of source measure unit (SMU) and 2361 trigger, was used to analyze the I-V curves of a Cu electrode in various electrolyte concentration and to find out the proper operating voltage at each electrochemical state. The I-V curves were plotted for each of the electrolyte concentration. From the I-V curves, the electrochemical states of active, passive, transient and trans-passive region could be characterized[12]. Appropriate operating voltages would be selected from these curves for the following quantitative tests. In quantitative analyses, the electrochemical process without the mechanical polishing process was performed in order to verify the relationship between the dissolved mass loss and applied voltages, when the selected operating voltages were applied during the constant process time such as 30, 60, 90 and 120 s, respectively. The sample masses before and after the quantitative test were measured on precision electronic scale to calculate the removal thickness of Cu disk. Finally, we fundamentally studied the chemical state and element composition of the Cu surface according to the concentration of the electrolyte and energy dispersive spectroscopy (EDS) analysis, respectively. In this way, we monitored the oxidation and reduction step of the Cu surface by the repetition of anodic and cathodic potential in KNO₃ and HNO₃ electrolyte.



Fig.1 Schematic diagram of *I*—*V* measurement system with two-electrode cell

3 Results and discussion

Fig.2 shows the I—V curves as a function of KNO₃ and HNO_3 electrolyte with concentration of 5%, 10% and 20%. As the concentrations of KNO3 electrolyte are increased, the trans-passive voltages are decreased and the current density is more increased. Thus, a plateau region, so called passive state, is not formed. This reason is because a continuous passive oxide or metallic hydroxide remains on the Cu surface. As the voltage more increases, the potential is great enough to pull the metallic ion through the passive film. Although there are changes of operating voltage and current density according to the electrolytes concentration, we point out that the ranges of proper operating voltage for subsequent ECMP experiment are between 0.5 V and 1.0 V. Also, as the concentration of HNO₃ electrolyte increases, the transient voltages are decreased and the curve slope of trans-passive state is more and more steep. This indicates that the *I*—V relationship strongly depends on the concentration of electrolyte.



Fig.2 I—V curves of KNO₃(a) and HNO₃(b) electrolyte with different concentration

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