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1. Introduction

Above the 32 nm technology node of integrated circuits, the conventional bilayer of Ta/TaN acts as a barrier layer for copper (Cu) interconnects to prevent copper diffusion, and enhance the adhesion between copper and dielectrics.¹⁻³ With the continuous reduction of technology nodes, the proportion of Ta/TaN in the whole interconnect structure is increasing, resulting in a sharp increase in interconnect resistance and RC delay due to its high resistivity.^{3,4} On the other hand, the ultrathin tantalum cannot realize conformal deposition in the high aspect ratio trenches, which brings great challenges to the subsequent deposition of the copper seed layer.⁴ To surmount these problems, ruthenium (Ru) material has attracted widespread attention and has been proposed as the most promising barrier liner. Ruthenium is a chemically stable transition metal with a high melting point (2250 °C), a resistivity (7.1 $\mu\Omega$ cm) lower than that of Ta (13.1 $\mu\Omega$ cm), and excellent adhesion with copper.⁵⁻¹¹ More importantly, the continuous copper film can be conformally

Effect of ethylenediamine on CMP performance of ruthenium in H₂O₂-based slurries

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With the aggressive scaling of integrated circuits, ruthenium has been proposed as the next generation barrier material to replace the conventional bilayer of tantalum and tantalum nitride due to its properties such as allowing direct copper electrodeposition. In this work, the effect of ethylenediamine (EDA) on the chemical mechanical polishing (CMP) properties of ruthenium in H₂O₂-based slurries was investigated. The results show that EDA or H₂O₂ alone has little effect, but the combined use of EDA and H₂O₂ significantly enhances the removal rate of ruthenium. Subsequently, the mechanism of action of ruthenium removal promoted by EDA was studied by combining CMP experiments, electrochemical experiments and surface chemical characterization methods. It is indicated that EDA molecules react with ruthenium oxide (not ruthenium metal) to generate a large number of complexes, which promotes the dissolution of ruthenium oxides and the corrosion of ruthenium. More importantly, the oxide layers on the ruthenium surface become rough and porous, and can be easily removed by mechanical action during the ruthenium CMP process. Meanwhile, the use of EDA can reduce the electrostatic repulsive force between the SiO₂ particles and ruthenium surface in the CMP process, thus further accelerating the ruthenium removal. In order to obtain an adequate removal rate selectivity of ruthenium versus copper, the corrosion inhibitors for copper were added. As a consequence, the removal rate selectivity of 1.13: 1 was obtained, while also reducing the corrosion potential difference between ruthenium and copper to 17 mV.

and directly deposited on the ruthenium surface, while no intermetallic compound is formed between copper and ruthenium.^{7,9,12,13}

Ruthenium, being a chemically stable and hard metal, is difficult to achieve the high removal rate in CMP process. To meet the requirements of efficient ruthenium removal, the researchers screened a number of oxidizers, such as KMnO4 (potassium permanganate), KClO₄ (potassium perchlorate), NaIO₄ (sodium periodate), K₃[Fe(CN)₆] (potassium ferricyanide), potassium peroxymonosulfate (oxone), $K_2S_2O_8$ (potassium persulfate), KIO₄ (potassium periodate), (NH₄)₂S₂O₈ (ammonium persulphate) and H₂O₂ (hydrogen peroxide), among which KIO4 and H2O2 receive widespread attention.^{7-9,12,14-21} H. Cui et al.²²studied the effect of various oxidizers on the removal rate of ruthenium from the perspective of surface corrosion and oxidation. The experimental results suggested that the existence of IO₄⁻ in the slurries would boost the formation of mechanically soft and porous oxide layers on the ruthenium surface, accelerate the corrosion of ruthenium surface, and greatly improve the removal rate of ruthenium. However, B. C. Peethala et al.7 and J. Cheng23 demonstrated that when KIO₄ served as an oxidizer for ruthenium, the remarkable potential gap between ruthenium and copper may bring about serious galvanic corrosion of copper. In contrast, the use of

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H₂O₂ could obtain the negligible corrosion potential difference between ruthenium and copper while forming a smooth and dense ruthenium surface.²²⁻²⁴ Turning to the slurries with H₂O₂ as oxidant, L. Jiang et al.19 reported that the added potassium ion could achieve the higher removal rate of ruthenium. This was mainly because the introduction of potassium ion increased the intensity of the reactions between H2O2 and ruthenium and the mechanical action in CMP process. H. P. Amanapu et al.8 investigated the effect of guanidine carbonate (GC) on ruthenium CMP using H₂O₂-based slurries, indicating that the increasing removal rate of ruthenium was because GC interacted with ruthenium oxides via the lone pair electrons on its nitrogen atoms to generate complexes. Subsequently, Y. C. Du et al.24 confirmed that guanidine ion (Gnd⁺) was the main factor for promoting ruthenium removal by comparing several common guanidine salts.

Most research on ruthenium CMP is mainly centered on the ruthenium oxidizers in the last decades of industrial research, but few studies on effective complexing agents for ruthenium. As the most critical chemical additive in polishing slurries, complexing agents can accelerate material removal by chelating metal ions to generate soluble species, especially in the field of barrier CMP.^{25,26} Ethylenediamine (EDA) containing two amino functional groups has been shown to be used as efficient complexing agents for enhancing the material removal rate in metals CMP.²⁷⁻²⁹ Liu et al.²⁸ achieved a high copper removal rate of 1899 nm min⁻¹ using the slurries with 100 mM EDA and 0.6% H₂O₂. Furthermore, H. P. Amanapu et al.⁸ simply mentioned that when 1 wt% EDA was applied to the pH 9 solution containing 50 mM H₂O₂ and 5 wt% SiO₂, the removal rate of ruthenium on titanium nitride increased from 4 nm min⁻¹ to 30 nm min⁻¹. Unfortunately, the authors did not elaborate on the action mechanism of EDA to boost the ruthenium removal rate. In this study, the eco-friendly H₂O₂ and EDA were used as oxidizers and complexing agents for ruthenium, respectively. It was confirmed that EDA in the presence of H₂O₂ could significantly promote the removal rate of ruthenium using CMP experiments. Then, the action mechanism of EDA promoting ruthenium removal was studied by using CMP experiments, electrochemical experiments and surface chemical characterization methods. Summarizing all the above results, the action mechanism of EDA on ruthenium was expressed. Finally, 5-methyl-benzotriazole (MBTA) and sodium dodecyl benzene sulfonate (SDBS) were added to the solutions with H₂O₂ and EDA to solve the problem about the removal rate selective of ruthenium to copper.

2. Experimental

2.1 Materials

The solutions in this work were prepared with deionized water, silica abrasives (40% purity; JinWei Group Co. Ltd., China) and analytically pure grade chemicals, in which silica abrasives were only used for CMP experiments and Zeta potential measurements. Chemicals including hydrogen peroxide (H_2O_2 ; 30% purity), ethylenediamine ($C_2H_8N_2$; 99.5% purity), ethylenediamine tetraacetic acid ($C_{10}H_{16}N_2O_8$; 98% purity), glycine

($C_2H_5NO_2$; 99% purity), potassium citrate ($C_6H_5K_3O_7$; 99.3% purity), 5-methyl-benzotriazole ($C_7H_7N_3$, 99.9% purity), sodium dodecyl benzene sulfonate ($C_{12}H_{25}SO_3Na$, 99.97% purity) and nitric acid (as pH adjuster) were obtained from Shanghai Macklin Biochemical Technology Co. Ltd while H_2O_2 purchased from Jiangtian Chemical Co. Ltd., Tianjin, China.

2.2 CMP experiments

3 inch wafers cut from 12 inch ruthenium blanket wafers (SKW Associates, Inc., USA) were polished on the E460 polisher (Alpsitec Inc., France) with a Politex Reg pad (Dow Chemical Company, USA). The process parameters of CMP were as follows: down pressure of 1.5 psi, slurry flow rate of 300 ml min⁻¹, carrier/platen speeds of 87/93 rpm, and the polishing time of 30 s. The removal rate of ruthenium was calculated by measuring the film thickness before and after CMP using a fourpoint probe (333 A; Four Dimensions Inc., USA). Each experiment was executed three times, and then averaged it. Also, the zeta potential of slurries was measured with a potential analyzer (NICOMP 380ZLS; PSS Inc.,USA).

2.3 Electrochemical experiments

The corrosion characteristics of ruthenium in the designated solutions were investigated using CHI660E electrochemical workstation obtained from Shanghai Chenhua Instrument Co. Ltd. A three-electrode cell was used to obtain the open circuit potential (E_{oc}) , potentiodynamic polarization and impedance spectroscopy (EIS) data, where ruthenium was employed as working electrode, standard calomel electrode (SCE) was served as reference electrode and platinum was used as counter electrode. Before each measurement, the ruthenium electrode was thoroughly sealed with insulating tape to expose an effective area of 1 cm², followed by polishing with 2000# sandpaper until its surface was mirror-like, and finally cleaned with deionized water and dried with pressure-air. The open circuit potential $(E_{\rm oc})$ was stabilized for approximately 1800 s to get a reliable and credible potential value. Potentiodynamic polarization was measured by employing a scan rate of 5 mV s⁻¹ over the voltage range of $E_{\rm oc} \pm 0.3$ V. EIS measurements were performed at an open circuit potential by employing the frequency range of 0.01 Hz to 1000 kHz, while an alternating current signal with the amplitude of 5 mV was applied. EIS data was analyzed and modeled by ZsimpWin software. In particular, prior to performing potentiodynamic polarization and EIS measurements, a stable E_{oc} was required.

2.4 Surface characterization

Before immersion in the solution to be tested, the sample processing method was as follows: 1×1 cm samples were cut from a 12 inch ruthenium blanket wafer and soaked in a 25 mM citric acid solution for 10 minutes, followed by rinsing with deionized water and finally dried with pressure-air. The morphological characteristics of the ruthenium soaked in the test solutions for 15 minutes was characterized by atomic force microscopy (AFM; 5600LS, Agilent) and scanning electron microscopy (SEM; Zeiss Sigma 500/VP, Carl Zeiss). The X-ray

photoelectron spectroscopy (XPS) device used herein was manufactured by Thermo Scientific and was model number ESCALAB250Xi. After testing, the XPS spectra was analyzed by CasaXPS software and calibrated according to the C1s peak (284.6 eV).

3. Results and discussion

3.1 Dependency of ruthenium RR on H₂O₂ concentration

The removal rate of ruthenium as a function of H_2O_2 concentration is present in Fig. 1, where 5 wt% SiO₂ is added to the slurries at pH 9. It is important to point out that all solutions without or with SiO₂ herein have a pH of 9 to avoid the generation of toxic RuO₄ in an acidic environment. It is obvious from Fig. 1 that the removal rate of ruthenium increases as the H_2O_2 concentration increases from 0 to 0.15 wt%, and reaches the maximum value at 0.15 wt% H_2O_2 . However, the removal rate of ruthenium decreases with the addition of H_2O_2 of 0.225 wt% or more, revealing that high concentration of H_2O_2 is not conducive to the ruthenium removal. To investigate the causes of these phenomena, the ruthenium removal mechanism in CMP process must be discussed first, as follows:

In alkaline environment, it is generally assumed that ruthenium can react with OH^- to generate a series of oxides through eqn (1)–(4).^{7,12,22,30} In the absence of H_2O_2 , the electrons



Fig. 1 (a) The ruthenium RR as a function of H_2O_2 concentration; (b) 3D AFM micrographs of ruthenium immersed in the following solutions: (b1) 0.075 wt% H_2O_2 at pH 9, (b2) 0.15 wt% H_2O_2 at pH 9 and (b3) 0.225 wt% H_2O_2 at pH 9, (b4) 0.3 wt% H_2O_2 at pH 9.

generated by the oxidation reactions are consumed by the oxygen reduction reaction, as expressed in eqn (5).^{7,22,30} After adding H_2O_2 to the alkaline solution, the H_2O_2 reduction reaction (as shown in eqn (6)) becomes the main cathode reaction, which can expedite the increase of OH⁻ concentration in the solution. Increase of OH⁻ concentration is beneficial to acceleration of the reaction presented in eqn (1)–(4), and thus promoting the generation of assorted oxides. Furthermore, H_2O_2 can react with ruthenium to generate $RuO_2 \cdot 2H_2O$, as shown in eqn (7).³⁰

$$Ru + 3OH^{-} \rightarrow Ru(OH)_{3} + 3e^{-}$$
(1)

$$Ru + 4OH^{-} \rightarrow RuO_{2} \cdot 2H_{2}O + 4e^{-}$$
⁽²⁾

$$RuO_2 \cdot 2H_2O + 4OH^- \rightarrow RuO_4^{2-} + 4H_2O + 2e^-$$
 (3)

$$RuO_2 \cdot 2H_2O + 4OH^- \rightarrow RuO_4^- + 4H_2O + 3e^-$$
 (4)

$$O_2 + 2H_2O + 2e^- \rightarrow 4OH^-$$
(5)

$$H_2O_2 + 2e^- \rightarrow 2OH^- \tag{6}$$

$$Ru + 2H_2O_2 \rightarrow RuO_2 \cdot 2H_2O \tag{7}$$

In the CMP process, the oxides generated on the ruthenium surface are rapidly abraded by mechanical action, and the exposed fresh surface is continuously oxidized or complexed, thus realizing ruthenium removal. The reason for the higher removal rate of ruthenium at rather low H₂O₂ concentration may be the formation of a discontinuous oxide layer on the ruthenium surface. However, when the concentration of H_2O_2 exceeds a certain value, the dense and thick oxide films are generated, resulting in the lower removal rate.24 It can be seen from Fig. 1b that the surface roughness of ruthenium becomes significantly smaller as the concentration of the added H₂O₂ is more than 0.15 wt%, indicating that the oxide films on the ruthenium surface is denser and smoother in this case. This is the reason why the removal rate of ruthenium shows a decreasing trend when H₂O₂ concentration is greater than 0.15 wt%.

3.2 Dependency of ruthenium RR on EDA concentration

Regarding the effect of complexing agents, the removal rate of ruthenium from slurries containing ethylenediamine tetraacetic acid (EDTA), glycine (Gly), potassium citrate (CAK) or EDA was tested, as shown in Fig. 2a. Obviously, ethylenediamine exhibited the largest removal rate of ruthenium, which indicated the best performance of ethylenediamine as the complexing agent for ruthenium in the slurry. The removal rate of ruthenium as a function of EDA concentration is present in Fig. 2, where 5 wt% SiO₂ is added to the slurries at pH 9. In the presence of 0.15 wt% H₂O₂, the removal rate of ruthenium increases from 116 Å min⁻¹ to 375 Å min⁻¹ as the EDA concentration increases from 0 to 40 mM. However, it is interesting that, without adding H₂O₂ into the slurries, the removal rate of Å min⁻¹ over the entire EDA concentration range. It can be said



Fig. 2 (a) The removal rate of ruthenium in the slurries with 5 wt% SiO₂, 0.15 wt% H_2O_2 and 40 mM complexing agent at pH 9; (b) the ruthenium RR as a function of EDA concentration.

that the introduction of EDA to 5 wt% SiO_2 did not expedite the ruthenium removal compared to the SiO_2 alone. These results suggest that EDA can effectively improve the removal rate of ruthenium, but it must be combined with H_2O_2 . The mechanism by which EDA facilitates ruthenium removal during CMP process will be described in detail below.

3.3 Mechanism of ruthenium removal promoted by EDA

Ethylenediamine is a diacidic base with dissociation constants of 6.848 and 9.928.^{29,31,32} The distribution of H_2EDA^{2+} , $HEDA^{+}$ and EDA, in solutions of different pH values can be calculated by the distribution fraction formula, as presented in Fig. 3, where EDA stands for $H_2N-CH_2-CH_2-NH_2$.

3.3.1 Interaction of EDA with SiO₂ **particles.** From Fig. 3, it is observed that, in the solution with pH 9, the main existing form of ethylenediamine is HEDA⁺. The positively charged HEDA⁺ will spontaneously adsorb around the negatively charged SiO₂ particles through electrostatic action, thereby neutralizing the negative charge on the SiO₂ particles surface. For investing in the interaction, zeta potential tests were performed on slurries with 5 wt% SiO₂, 0.15 wt% H₂O₂, various concentrations of EDA and pH 9, as shown in Fig. 4a. With the



Fig. 3 Speciation diagram of ethylenediamine in aqueous solution (EDA stands for $H_2N-CH_2-CH_2-NH_2$).

increase of EDA concentration, the absolute value of zeta potential of slurries decreases, which indicates that the negative charge on the SiO₂ particles surface decreases gradually. The neutralization of the negative charge of the SiO₂ particles reduces the electrostatic repulsive force between SiO₂ abrasives and the negatively charged ruthenium surface (isoelectric point of 4-6), thus strengthening the mechanical action and further accelerating the removal rate of ruthenium.^{33,34} To validate that the enhancement of mechanical action is not the only factor of promotion of the ruthenium removal, the effect of potassium ion with the same amount of positive charge as HEDA⁺ on removal rate of ruthenium is illustrated in Fig. 4b. Obviously, the introduction of EDA shows a greater ruthenium removal rate at the same molarity. Complexation reaction of EDA with ruthenium and its oxides will be confirmed in the following experiments.

3.3.2 Open circuit potential analysis. Open circuit potential represents the potential difference between the working electrode and the reference electrode in the absence of applied current, and can be used to reflect the corrosion or passivation state of working electrode surface. Fig. 5 depicts the $E_{\rm oc}$ curves of ruthenium in the presence of 0.15 wt% H₂O₂ and varying concentrations of EDA at pH 9. It is obvious that the $E_{\rm oc}$ of ruthenium is approximately 0.135 V to 0.205 V (*versus* SCE) over the entire EDA concentration range. Within the range of $E_{\rm oc}$ obtained here, Ru, Ru(OH)₃, RuO₂·2H₂O, RuO₄⁻⁻ and RuO₄²⁻⁻ are mainly formed on the ruthenium surface according to Pourbaix diagram of Ru-H₂O system.

Without adding EDA into the solution, the formation of passivating layer of Ru(OH)₃, and RuO₂·2H₂O (eqn (1), (2) and (7)) brings about the slow and continuous growth of E_{oc} over time. Moreover, with adding EDA and H₂O₂, E_{oc} reaches the maximum value at a certain time and then decreases continuously, which can be explained as follows: the generation of passivating layer of Ru(OH)₃ and RuO₂·2H₂O makes the E_{oc} increase firstly. However, this layer is likely to be discontinuous because the introduction of EDA promotes the dissolution of partial oxides on the ruthenium surface, resulting in the observed decline of E_{oc} over time.



Fig. 4 (a) Dependency of zeta potential of slurries on EDA concentration; (b) the ruthenium RR as a function of "X" concentration (X stands for EDA or KNO₃).



Fig. 5 $E_{\rm oc}$ -time curves of ruthenium in the designated EDA concentration.

3.3.3 Potentiodynamic polarization curves measurements. Potentiodynamic polarization curves performed on ruthenium in a solution are presented in Fig. 6, where the solution consists



Fig. 6 Potentiodynamic polarization curves for ruthenium in the designated EDA concentration.

of 0.15 wt% H₂O₂, various concentrations of EDA and pH 9. The corresponding parameters of corrosion potential (E_{corr}) and corrosion current density (J_{corr}) are obtained from Fig. 6 by Tafel method and listed in Table 1.³⁵ It is clear from Table 1 that the E_{corr} decreases with increasing EDA concentration, while the J_{corr} increases. The decrease of corrosion potential and the increase of corrosion current density indicate that the chemical action of the solution on ruthenium is enhanced and the corrosion rate of ruthenium in the solution is accelerated. Moreover, the anodic curves of ruthenium treated with EDA emerge a weak passivation region, which is probably due to the accumulation of oxides caused by the increase of the anodic current, and the adsorption of HEDA⁺.

3.3.4 EIS measurements. EIS tests were introduced to analyze the surface properties and the kinetics of electrode processes for ruthenium. Before analyzing the experimental data of EIS and fitting any theoretical model on them, it is necessary to verify whether these data fulfill the following conditions: causality, linearity and stability. It is generally believed that if the electrochemical system satisfies these conditions, the real part (Z_r) and imaginary part (Z_i) of the complex impedance (Z) will conform to the Kramers–Kronig Transformation (KKT). The KKT is simply a mathematical relationship, and is specifically expressed as follows:³⁶

$$Z_{\rm i}(w) = \frac{2\omega}{\pi} \int_0^\infty \frac{Z_{\rm r}(x) - Z_{\rm r}(w)}{x^2 - w^2} dx$$
(8)

Table 1 Corresponding electrochemical parameters of ruthenium obtained with Fig. ${\rm 6}$

EDA (mM)	$E_{\rm corr}$ (V)	$J_{ m corr}$ (A cm ⁻²)			
0	0.173	$5.843 imes10^{-6}$			
10	0.107	3.451×10^{-5}			
20	0.102	4.536×10^{-5}			
30	0.094	$6.313 imes10^{-5}$			
40	0.082	1.111×10^{-4}			

$$Z_{\rm r}(w) = Z_{\rm r}(\infty) + \frac{2\omega}{\pi} \int_0^\infty \frac{xZ_{\rm i}(x) - \omega Z_{\rm i}(w)}{x^2 - w^2} \mathrm{d}x \tag{9}$$

where $Z_{\rm r}(\omega)$ and $Z_{\rm i}(\omega)$ are functions of the angular frequency $\omega = 2\pi f; f$ refers to frequency; and $Z_{\rm r}(\infty)$ represents real part of the impedance extrapolated to infinite frequency. It should be noted that since an analytical expression for Z is not available, the values of $Z_{\rm r}$ and $Z_{\rm i}$ can only be obtained from experimental data over a limited frequency range. Obviously, it can be seen from Fig. 7 that the experimental data are highly coincident with the KKT calculated data, suggesting that the data obtained by this electrochemical system are reliable.³⁶

Nyquist and Bode plots for ruthenium electrode dipped to the solutions containing 0.15 wt% H₂O₂, varying concentrations of EDA and pH 9 are given in Fig. 8. Seeing from Fig. 8a, the Nyquist plots appear as a capacitive arc at both the high and intermediate frequency area, followed by a straight line at low frequency area. Additionally, all Nyquist plots have the same shape, revealing that the introduction of EDA only changes the value of impedance, without affecting other electrochemical properties in the system. Generally, the sum of the diameter of two capacitive arcs is called polarization resistance (R_p) .³⁷ The $R_{\rm p}$ continually decreases with the consecutive addition of EDA content, indicating that added EDA is beneficial to the corrosion of ruthenium. The similar trends can be seen from the phase angle and frequency variations in the Bode plots of Fig. 8b. The impedance modulus (|Z|) at the lowest frequency and the highest phase angle decreases with the increasing concentration of EDA, which further proves that EDA corrodes the ruthenium surface. It is particularly noted that two peaks could be seen from the phase angle curves, which reveals that the impedance spectrum has two time constants and that micropores and cracks may be formed on the surface of the ruthenium electrode.38

In order to obtain information about the relevant model for dealing with the EIS data, the Nyquist plots are deeply analyzed, as follows: the first capacitive arc at the high frequency area is caused by the parallel arrangement of film capacitance (C_f) and film resistance (R_f); the second capacitive arc appearing at the intermediate frequency area is characterized by the parallel linkage of the charge transfer resistance (R_{cf}) and double layer

capacitance (C_{dl}); Warburg impedance (W) associated with the diffusion of reactants and products at the interface between electrode and solution is used to characterize the straight line at the low frequency area. Therefore, it can be determined that the electrode process for ruthenium contains two time constants and is influenced by mass transfer process. Moreover, two capacitive arcs that are slightly depressed and the straight line with a slope that deviates by $\pi/4$ may be credited with the unevenness or surface roughness of ruthenium electrode, resulting in the replacement of C_{f} and C_{dl} by constant phase elements (CPE) to obtain an appropriate equivalent circuit.³⁹ The constant phase elements of C_{dl} and C_{f} are denoted by CPE_{dl} and CPE_f, respectively. The impedance of a CPE is expressed in following equation:³⁷⁻³⁹

$$Z_{\rm CPE} = \left[Q(j\omega)^n\right]^{-1} \tag{10}$$

where *Q* means the modulus of CPE, *j* is imaginary root, and *n* stands for CPE exponent with values between -1 and 1. The specific meaning of CPE varies with *n*, which is described as follows: when n = -1, CPE represents inductance; when n = 0, it is resistance; when n = 0.5, it stands for Warburg impedance; when n = 1, it is ideal capacitance. The impedance of a *W* can be presented as:⁴⁰⁻⁴²

$$Z_W = \sigma \omega^{-\frac{1}{2}} (1-j) \tag{11}$$

where, σ is taken as Warburg impedance coefficient. Finally, Fig. 9 shows the resulting equivalent circuit, and the fitted values are shown in Table 2. The corresponding values of $C_{\rm f}$ and $C_{\rm dl}$ in the table can be calculated by the following eqn (10), where $f_{z_{\rm im}-{\rm max}}$ is the frequency value at which the imaginary part of the impedance in the Nyquist plots is maximum.³⁷⁻³⁹

$$C = Q(2\pi f_{z_{\rm im}-\rm max})^{n-1}$$
(12)

As can be seen from Table 2, R_f , R_{ct} and the sum of both (R_p) decrease with the augmenting content of EDA. This observation suggests that the addition of EDA significantly weakens the resistance of the solution to corrosion of ruthenium and increases the active sites of ruthenium dissolution reaction,



Fig. 7 Comparison of experimental and KKT calculated data of (a) Z_r and (b) Z_i .



Fig. 8 Bode plots (a) and Nyquist curves (b) for ruthenium after treatment with different solutions.



Fig. 9 The electrical equivalent circuit used to fit the EIS data.

thereby accelerating the chemical dissolution of ruthenium and its oxides. In addition, the values of $C_{\rm f}$ and $C_{\rm dl}$ show an increasing trend as the EDA concentration increases, and the specific reason can be explained by the following expression:³⁷⁻³⁹

$$C_{\rm f} = \frac{F^2 S}{4RT} \tag{13}$$

$$C_{\rm dl} = \frac{\varepsilon^0 \varepsilon}{d} s \tag{14}$$

where *S* means the effective contact area between electrode and solution, *F* represents the Faraday constant, *d* represents the thickness of electric double layer, and ε^0 and ε represent the permittivity constants of air and the electric double layer, respectively. Therefore, the increase of $C_{\rm f}$ and $C_{\rm dl}$ could be

attributed to the expansion of effective contact area (*S*) attributed by the destruction of the dense oxide layers on the ruthenium surface by EDA molecules. In addition, the thinning of the oxide layers on the ruthenium surface caused by EDA complexation may also provoke the growth of $C_{\rm f}$ and $C_{\rm dl}$.

The increase of W indicates that the presence of EDA accelerates the diffusion of corrosion ions to the ruthenium surface, and also enhances the diffusion of soluble ruthenium complexes from the ruthenium surface to the solution. For further detailed qualitative analysis, the Warburg impedance coefficient σ that is associated with the electrolyte diffusion behavior can be obtained the Randles curves (Z_r or Z_i against $\omega^{-1/2}$) in Fig. 10a and Fig. 10b.⁴² In general, the lower the value of σ is, the easier it is for the electrolyte to diffuse into the solution through the pores in the oxide films. The slope of $Z_{\rm r}$ against $\omega^{-1/2}$ in Fig. 10a is regarded as σ because of the higher degree of linear fitting. Seeing from Fig. 10c, σ decreases with the augmenting concentrations of EDA. Increase in σ means a growth in the number of the ions permeated the oxide layers, which can be interpreted as a more porous, coarser and thinner oxide films on the ruthenium surface resulting from an increased corrosion rate using EDA. The rough and porous nature of oxide films will then be demonstrated by surface morphology measurement.

3.3.5 Surface morphology measurements. SEM observations were utilized to assess the surface morphology of ruthenium soaked in aggressive solutions. Fig. 11 depicts the SEM micrographs of ruthenium in certain solutions: (a) deionized water (blank sample); (b) 0.15 wt% H_2O_2 at pH 9; (c) 0.15 wt% H_2O_2 and 30 mM EDA at pH 9. It is clear that from Fig. 11a,

Table 2 Fitted impedance parameters of ruthenium obtained with EEC											
EDA (mM)	$R_{\rm s} \left(\Omega \ {\rm cm}^2\right)$	$R_{\rm f} \left(\Omega \ {\rm cm}^2\right)$	$R_{\rm ct} \left(\Omega \ {\rm cm}^2\right)$	$R_{\rm p} \left({\rm k}\Omega ~{\rm cm}^2 \right)$	$C_{\rm f} \left(\rm nF \ cm^{-2} \right)$	$n_{\rm f}$	$C_{\rm dl} \left(\mu { m F~cm^{-2}} \right)$	n _{dl}	$W(\mu\Omega^{-1} \ { m cm}^{-2} \ { m s}^{0.5})$		
0	85.30	8451	1316	9767	0.34	0.98	35.89	0.94	1.82		
10	68.41	1454	972	2426	0.73	0.98	38.02	0.88	3.70		
20	30.30	598	832	1430	0.96	0.98	51.68	0.87	8.75		
30	13.77	382	560	942	1.10	0.99	62.35	0.87	13.60		
40	9.90	288	513	801	1.24	1	63.01	0.88	17.70		



Fig. 10 The Randles curves ((a) Z_r against $\omega^{-1/2}$, (b) Z_i against $\omega^{-1/2}$) were obtained using the low-frequency region data of the Nyquist diagram in Fig. 6c. (c) Relationship between σ and the concentration of EDA.

when only immersed in deionized water without any corrosive ions, the ruthenium surface is remarkably smooth and not attacked, and clearly shows crystal grains with a diameter of about 30 nm. However, an increase in surface roughness is evident when ruthenium is immersed in a solution with H_2O_2 (Fig. 11b), which is usually accompanied by the formation of porous and rough oxides. A considerably deteriorated and roughed surface results from further addition of EDA to the solution containing H_2O_2 (Fig. 11c). This is because of the strong complexation of EDA, the oxide films on ruthenium surface cannot effectively frustrate attack with aggressive ions in the corrosive environments, resulting in the acceleration of the corrosion of ruthenium.

Due to the lack of quantitative information of surface roughness in SEM images, AFM, which can distinguish the changes of surface morphology at the atomic level and calculate the surface roughness of the samples, was employed to further investigate the surface topography of ruthenium dipped in different corrosive solutions. 3D AFM micrographs and corresponding height profiles of ruthenium surface immersed in solutions without and with H_2O_2 or EDA are illustrated as Fig. 12. From the 3D micrograph in Fig. 12c, it should be observed that when H_2O_2 and EDA are present in the solution simultaneously, the ruthenium surface is heavily corroded, which provokes the generation of bumpy and rough surface with deep and large pits and presents the maximum surface roughness ($S_q = 1.24$ nm), which is the result of the corrosive attack of EDA with strong complexation. The height profile in Fig. 12f also exhibits the same phenomenon, with a peak–valley difference (PV) of 7.52 nm, which is also the largest among the three samples. It is remarkable that when H₂O₂ is added alone (Fig. 12b and e), both the value of the surface roughness and peak–valley difference are slightly increased compared with the blank sample (Fig. 12a and d), which may be correlated with the formation of the rough and porous oxide layers by H₂O₂ oxidation. This observation is consistent with the electrochemical and SEM results, indicating that the synergistic effect of H₂O₂ and EDA can accelerate the corrosion of ruthenium, thereby improving the removal rate.

3.3.6 XPS measurements. XPS tests of ruthenium samples with different treatments were carried out to investigate the chemical composition of oxide layer on ruthenium surface in the absence and presence of EDA. Fig. 13 displays the fitted results of XPS spectra on the ruthenium surface after immersion in a solution with and without EDA, and the detailed analysis results of Ru(3d) are shown in Table 3. The Ru(3d) spectra of the ruthenium surface presents five peaks, including Ru peak at about 280.2 eV (3d_{5/2}) and 284.4 eV (3d_{3/2}), RuO₂··2H₂O peak at about 281.0 eV (3d_{5/2}) and 285.3 eV (3d_{3/2}) and RuO₃ peak at about 282.5 eV (3d_{5/2}).^{12,43,44} It should be noted that



Fig. 11 SEM micrographs of ruthenium in certain solutions: (a) deionized water (blank sample); (b) 0.15 wt% H₂O₂ at pH 9; (c) 0.15 wt% H₂O₂ and 30 mM EDA at pH 9.

the absence of Ru(OH)₃ on the ruthenium surface may be credited with the oxidation of Ru(OH)₃ into RuO₂·2H₂O under the action of oxidizer.¹² Furthermore, it is considered that RuO₃ exists on the bulk-phase of RuO2 · 2H2O.12 Therefore, the oxides covering the ruthenium surface are mainly RuO2 · 2H2O and RuO₃. From the results shown in Table 3, it can be seen that the content of the oxides on the ruthenium surface is reduced after immersion in the solution containing H₂O₂ and EDA. Combined with the results of electrochemical and surface morphology, it can be reasonably inferred that EDA may mainly interact with ruthenium oxides and can effectively facilitate the dissolution of ruthenium oxides. This may also explain why the addition of EDA to the solution without H₂O₂ in Fig. 2b cannot effectively improve the removal rate of ruthenium, which is mainly due to the extremely low content of oxides on the ruthenium surface in the absence of H₂O₂. In addition, a peak located at 399.2 eV in the N(1s) spectra of Fig. 13b may be ascribed to the nitrogen atoms of EDA adsorbed on the ruthenium surface.29

3.4 Action mechanism of EDA on ruthenium

Summarizing all the above results, the action mechanism of EDA on ruthenium is expressed, as presented in Fig. 14. Ruthenium can react with OH^- to generate a series of oxides, such as $RuO_2 \cdot 2H_2O$ and RuO_3 , when a solution has a pH of 9. It should be mentioned here that the oxidation of ruthenium is a gradual process, including $Ru(OH)_3$, *etc.* transitional oxidation

states. Moreover, these oxides can be slowly dissolved and converted to RuO4⁻ and RuO4²⁻ species under the action of OH-. The increase of OH- concentration in solution after adding H₂O₂ promotes the formation of ruthenium oxides, resulting in an oxide layer consisting of RuO₂·2H₂O and RuO₃ on the ruthenium surface, as presented in Fig. 14a. When EDA is present in solution with H₂O₂, EDA interacts with ruthenium oxides via the lone pair electrons on its nitrogen atoms to generate a number of complexes, thus facilitating the dissolution of the oxides on the ruthenium surface. More importantly, the ruthenium surface is severely corroded and illustrates rough and porous, as depicted in Fig. 14b. Finally, the oxides generated on the ruthenium surface are rapidly abraded by mechanical action during the ruthenium CMP process of Fig. 14c. Additionally, the addition of EDA can reduce the zeta potential for slurries, thereby reducing the electrostatic repulsive force between the SiO₂ particles and ruthenium surface, enhancing the mechanical action and further accelerating the ruthenium removal.

3.5 Removal rate selectivity

In order to achieve planarization during barrier CMP, copper dishing, which is the height difference of a copper with respect to the adjacent oxides, must be controlled.⁴⁵⁻⁴⁹ This would require that the removal rate of barrier material is greater than that of copper. Only in this way can the dishing defects caused by the copper overburden CMP step be corrected, the residual



Fig. 12 3D AFM micrographs and corresponding height profiles of ruthenium immersed in (a and d) deionized water, (b and e) 0.15 wt% H_2O_2 at pH 9 and (c and f) 0.15 wt% H_2O_2 and 30 mM EDA at pH 9.

copper be removed, and the surface non-uniformity of the patterned wafer be controlled.⁴⁹ Fig. 15 presents the removal rate for ruthenium and copper obtained using the slurries with various components (the specific information of slurries is embodied in the title of Fig. 15) and shows the removal rate selectivity for ruthenium to copper in the corresponding

slurries. It should be pointed out that in this experiment, the EDA concentration is selected as 30 mM for ensuring the stability of the slurries with SiO_2 particles and the fastest removal rate of ruthenium. Compared with slurry A, the addition of EDA in slurry B improves the removal rate of ruthenium, but the removal rate of copper is remarkably higher than that of



Fig. 13 (a) Ru(3d) and (b) N(1s) spectra of ruthenium after dipping into Ref solutions without or with 30 mM EDA (Ref: 0.15 wt% H₂O₂ at pH 9).

Table 3 The XPS results of Ru(3d) obtained with Fig. 13



Fig. 14 Schematic diagram of action mechanism of EDA on ruthenium. (a) H_2O_2 ; (b) H_2O_2 and EDA; (c) ruthenium CMP.



Fig. 15 The removal rate of ruthenium and copper treated with slurry A (Ref :5 wt% SiO₂ + 0.15 wt% H₂O₂ at pH 9), slurry B (Ref + 30 mM EDA) and slurry C (Ref + 30 mM EDA + 100 ppm MBTA + 300 ppm SDBS).

ruthenium. On this basis, the corrosion inhibitor combination for copper in another study⁵⁰ is added (as shown in slurry C), and successfully adjust the removal rate selectivity to 1.13:1.

3.6 Evaluation of galvanic corrosion

Galvanic corrosion refers to the phenomenon that when two metals contact each other or communicate through other electrolyte solutions, the metal with a lower potential is corroded preferentially.^{7,30,51,52} Since the standard electrode potentials of copper and ruthenium are 0.337 V and 0.45 V (*versus* SHE), respectively, the Cu/Ru interface is prone to galvanic corrosion, which leads to the accelerated corrosion of copper during the ruthenium barrier CMP.³⁰ It is generally considered that the galvanic corrosion can be effectively controlled only when the corrosion potential difference (ΔE_{corr})



Fig. 16 The potentiodynamic polarization plots of copper and ruthenium for different solutions, where the inhibitor consists of 100 ppm MBTA and 300 ppm SDBS (Ref: 0.15 wt% H_2O_2 + 30 mM EDA at pH 9).

is less than 20 mV. Fig. 16 illustrates the potentiodynamic polarization plots of ruthenium and copper for different solutions. Obviously, in the absence of a corrosion inhibitor of copper, the ΔE_{corr} between ruthenium and copper is negligible compared to potassium periodate and potassium periodate as oxidizers for ruthenium. MBTA and SDBS were also introduced to acquire a satisfactory removal rate selectivity, at which the ΔE_{corr} between copper and ruthenium is 17 mV.

Conclusion

The removal rate of ruthenium in SiO₂-based slurries comprised of EDA, H₂O₂ or both at pH 9 was investigated. It was demonstrated that the single use of EDA or H₂O₂ had limited effect on improving the removal rate of ruthenium, but the combination of EDA and H₂O₂ evidently facilitated the ruthenium removal, revealing that the formation of oxides was a key step, and EDA mainly reacted with oxides on ruthenium surface. Next, by exploring the effect of potassium ion with the same positive charge as HEDA⁺ (the main forms of EDA in pH 9) on removal rate of ruthenium, it was verified that the enhancement of mechanical action was not the main factor. Then, the results of electrochemical experiments, surface morphology measurements and XPS measurements showed that EDA could facilitate the dissolution of ruthenium oxides, thus facilitating the corrosion of ruthenium and forming a rough and porous oxide layer on the ruthenium surface. It was likely to be a large number of complexes formed by the interaction of the lone pair electrons on nitrogen atoms of EDA with ruthenium oxides. The rough and porous oxide layers were rapidly removed by mechanical action of SiO2 particles during the ruthenium CMP process. Finally, the removal rate selectivity of 1.13:1 and the corrosion potential difference of 17 mV were attained by adding MBTA and SDBS to the solution with EDA and H_2O_2 .

Author contributions

Yi Xu: conducting research, conceptualization of data, analysis and interpretation of data, writing-original draft preparation; Tengda Ma: conducting research, acquisition of data; Yuling Liu: review & editing; Baimei Tan: conceptualization, supervision, resources, project administration, writing-reviewing and editing; Shihao Zhang: investigation; Yazhen Wang: investigation; Guoqiang Song: experimental instrument test.

Conflicts of interest

There are no conflicts to declare.

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