4,6-Dimethyl-2-mercaptopyrimidine as a potential leveler for microvia filling with electroplating copper

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The impact of 4,6-dimethyl-2-mercaptopyrimidine (DMP) as a potential leveler on microvia electroplating copper filling from an acid cupric sulfate electrolyte, including polyethylene glycol (PEG), bis(3-sulfopropyl) disulfide (SPS), and chloride ion, was investigated. The results of electrochemical measurement, atomic force microscopy (AFM), and X-ray photoelectron spectra (XPS) revealed that DMP adsorption on the copper surface inhibited copper deposition. Different surface morphology and crystalline orientation was observed after plating using electroplating solutions with different concentrations of DMP. Analytical data were obtained by field emission scanning electron microscope (FE-SEM) and X-ray diffractometer (XRD), respectively. In addition, quantum chemical calculations and molecular dynamics (MD) simulations were used to investigate the interaction mechanism between DMP and copper.

1. Introduction

Currently, multifunctional and portable electronic products, such as smart phones, tablets, and wearable devices, are rapidly developing in the direction of miniaturization and integration. Not only IC chips substrates but also motherboards of electronic products need high-density interconnections by method using stacked microvias. 1,2 In order to meet the demands of electronic products with multifunctions, high frequency, and high-speed performance, 3 the microvias should be entirely filled by electroplating copper without seams or voids; i.e., so-called “superfilling” or “bottom-up filling.” 4 However, according to reported literature, 5 superfilling cannot be achieved due to uneven local current density contributions unless some particular additives are used which have significant influence on performance and orientation of the copper deposition. 6,7

To meet industrial requirements, several additives, such as an accelerator, a suppressor, and a leveler, must be simultaneously added to the electroplating copper solution. 7 In this additive system, the accelerator not only combined with chloride ion enhanced copper deposition at the bottom but also improved grain structure of the electroplating copper; the suppressors quickly adsorbed on the copper surface in the presence of chloride ion, forming a saturated film that seriously hindered diffusion of Cu 2+ to the copper surface and inhibited the copper deposition rate. 8,9 For the sake of improving microvia electroplating copper filling performance and promoting copper surface uniformity, the leveler also was added to the electroplating solution. 10 This was demonstrated in a large number of studies which showed that interactions among the additives in this system played a very significant role in the bottom-up filling. 3 Dow et al. 11 reported that ABPV can spontaneously adjust interaction between the accelerator and the suppressor from a bad filling pattern to a perfect pattern at low chloride ion concentrations. Moffat et al. 9 proposed that dodecytrimethylammonium chloride (DTAC) can deactivate the adsorbed SPS accelerator due to ion-pair interaction between the cationic head group of the DTAC and the anionic tail group of the adsorbed SPS accelerator.

As is well known, a leveler is an indispensable part of an electroplating additive system, which improves distribution of the current density and promotes microvia filling yields. However, the commonly used levelers in copper electrodeposition are quaternary ammonium cations, such as Jauns Green B (JGB), Diazine Black (DB), Methylene Blue (MB), N-butyl-methyl piperidinium bromide (PP 14Br), etc. 12-17 It is a very time-consuming and expensive job to develop additives which can be used for microvia filling. 18 Until now, only few additives used for microvia filling clearly have been reported.

According to previous research, organic compounds with several heteroatoms (e.g., sulfur, nitrogen), conjugated double bonds, and/or polar functional groups, which have strong coordination and adsorption capability to the copper surface, can usually be used as inhibitors for corrosion of copper. 19
Interestingly, the behavior of these organic compounds showed that they may be used as levelers for copper superfilling. At present, only a few compounds, such as 2-mercaptopyridine (2-MP), 6-aminobenzothiazole, and 4-amino-2,1,3-benzothiadiazole, have been studied as levelers for electroplating copper in microvias. Based on these structures, 4,6-dimethyl-2-mercaptopyrimidine (DMP) (Fig. 1) also appeared promising to improve filling performance, and its price is cheaper than JGB.

Another paper has reported DMP as an effective leveler, but it has no detailed research. Therefore, the purpose of the present work is to study filling performance in detail and electrochemical behaviors of different formulas with or without DMP, plus surface morphology, crystalline structure, thermal shock, and thermal cycle tests after copper plating. It has been proven that plating formulas containing DMP can completely meet requirements for PCB application reliability. To further study the interaction between DMP and the surface of copper, AFM and XPS patterns research was carried out. Also, both molecular dynamics (MD) simulation and quantum chemical calculations were used to analyze the active sites and adsorption behavior of DMP on copper surface. Based on the results of electrochemical measurements, AFM, XPS, and theoretical calculations, we can further explain the filling effect through combining with a microvia electroplating copper filling mechanism.

2. Experimental

2.1 Electroplating

The acidic electroplating Cu conditions, pretreatment techniques, and electroplating procedures have been presented in detail elsewhere. Test boards consisted of PCB fragments with a large number of microvias fabricated by CO₂ laser ablation. The dimensions of these test boards were 6 × 13 cm² with an effective size of 6 × 9 cm². Diameters and depths of the microvias were 100 μm and 80 μm, respectively. Prior to microvia electroplating copper filling, the sidewalls of these microvias were metalized by electroless copper. Following that, flash plating copper (electroplating copper) was carried out to increase the thickness of copper layers up to about 5 μm to prevent electroless copper oxidation.

Test boards were plated at a current density of 1.94 A dm⁻² for 70 min at 25 °C. Two phosphorus-containing copper plates with a size of 6 × 13 cm² were used as electroplating anodes and were directly placed in a Haring cell with a 1.5 L working volume. Constant agitation was produced by a wide stretch of air bubbles flowing at medium speed during electroplating of copper to guarantee efficient mass transfer. The composition of the base electrolyte was 0.88 mol L⁻¹ CuSO₄·5H₂O, 0.54 mol L⁻¹ H₂SO₄. Additives, such as HCl, SPS, PEG with a molecular weight of 8000, and DMP, were separately added to the Haring cell through dilution from stock solutions. The filling performance of various plating formula was assessed according to the cross-sections of microvia, which were examined using an optical microscope (OM, JX23, Guangdong Zheng ye Technology Co., LTD., China). Before examining the cross-sections of microvia, which were polished continuously with a series emery papers (320, 2000, and 5000 grit), they were etched to observe microstructures of the copper deposits.

2.2 Electrochemical measurements

All electrochemical measurements were conducted in a 250 mL glass beaker including 100 mL electroplating solution by a CHI660E electrochemical workstation with a conventional three-electrode system. The electroplating solution temperature was kept at 25 °C during electrochemical tests. A copper billet with a work area of 1 cm² by epoxy resin sealing was used as the working electrode (WE). The counter electrode (CE) was a platinum plate and the reference electrode was a saturated calomel electrode (SCE) with a Luggin capillary.

Prior to each measurement, the WE was abraded by a series of emery papers (320, 800, 2000, and 5000 grit) and was soaked in electroplating solution for 40 min until an almost steady-state open circuit potential (OCP) was obtained. Subsequently, electrochemical impedance spectroscopy (EIS) measurements were carried out at the OCP in a frequency range from 10⁵ Hz to 10⁻⁴ Hz with an amplitude of 5 mV s⁻¹. The EIS data were analyzed and fitted by Zsimpwin 3.10 software. Lastly, potentiodynamic polarization measurements were implemented in the potential range from OCP to −1.0 V with a scan rate of 5 mV s⁻¹. All potential values obtained during electrochemical experiments were referred to SCE. The same electrochemical measurements were implemented 3 times to ensure good repeatability.

2.3 Reliability tests

A copper plating sample was placed in a constant temperature oven at 150 °C for 6 h and then immersed fully in a lead-free solder furnace at 288 °C for 10 s. Then it was taken out and cooled to room temperature as one test cycle. The whole test requires six repeated cycles.

Then the copper plating sample was placed in a cycle test chamber, at temperatures of 125 °C and −55 °C for 15 min and working at room temperature for 20 min, respectively. This process is a test cycle, and the entire test requires 100 cycles.

2.4 FE-SEM and XRD measurements

Surface morphologies of the copper deposits obtained from different plating solutions were observed by FE-SEM, (JEOL-
Structure orientation of the Cu electrodeposition was researched and analyzed by XRD (Tongda TD-3500, China) using Cu Kα radiation (1.54056 Å) at room temperature. The diffractograms were recorded in 2θ range from 30° to 100° with a step size of 0.07°.

2.5 AFM and XPS measurements

The interaction between DMP and copper surface was investigated by AFM and XPS patterns. Copper sheets with surface area size of 1 cm², which were intercepted from PCB, were used for AFM and XPS measurements. The copper sheets had to go through the following steps for preparation: initially, they were abraded and polished using all kinds of emery papers (320, 800, 2000, and 5000 grit) and polishing cloths with alumina polishing powder, respectively. Then, the polished copper sheet surfaces were washed ultrasonically with ethanol and ultrapure water, and then dried at room temperature to obtain a shining mirror-like copper surface. Lastly, they were immersed in 1 ppm DMP aqueous solution for 30 min. After soaking, they were rinsed with ultrapure water and dried at room temperature.

AFM analysis was implemented with a MFP-3D-BIO (Uxford), working in the AC mode with Q-control. The XPS spectra tests were carried out using a Thermo Fisher ESCALAB 250Xi System at room temperature and employed Al Kα (1486.6 eV) as the incident radiation source.

2.6 Calculation methods

Quantum chemical calculations were performed by Gaussian 03W software. The DMP molecular structure was absolutely optimized by density functional theory (DFT) using B3LYP functional with the 6-311++G(d,p) basis set. The most popular quantum chemical parameters, such as the energy of the frontier molecule orbital (E_HOMO, E_LUMO), energy gap (ΔE = E_LUMO − E_HOMO), and dipole moment (μ), were calculated and considered. These parameters would be used to further explain the interaction mechanism between DMP and the copper surface.

Furthermore, interaction between DMP and the Cu (111) surface was studied by employing the forcite module in Material Studio software. The molecular dynamics (MD) simulation was carried out at 298 K with a time step of 1.0 fs and simulation time of 500 ps. The binding energy E_{Cu-DMP} between the DMP and Cu (111) surface was calculated as:

\[ E_{Cu-DMP} = E_{complex} - E_{Cu} - E_{DMP} \]

where \( E_{complex} \) was the total energy of the copper crystal together with the adsorbed DMP, \( E_{Cu} \) and \( E_{DMP} \) were the total energy of the copper crystal and free DMP molecular, respectively.

3. Results and discussion

3.1 Electroplating test

Filling results of electroplating copper solutions with different formulas are shown in Fig. 2 and 3, and their filling performance is defined as the following equation:
where $\eta$ is the filling efficiency of microvia, $A$ is the copper plating thickness inside the microvia, and $B$ is the dielectric layer thickness added to the copper plating thickness on the top surface of PCB. According to a previous research report, a standard used to determine whether the filling effect is good depends on whether the filling performance of the microvia is more than 85% and the void in the middle of microvia is visible (Table 1).

As can be seen from part a of Fig. 2, a large void was observed in the middle of the microvia; it could be attributed to non-uniform distribution of current density when no organic additives were present in the electroplating solution. This phenomenon disappeared when 200 ppm PEG (basic 2) or 1 ppm SPS and 200 ppm PEG (basic 3) was added to a bath solution, but did not turn into bottom-up filling and instead to conformal deposition.

The bottom-up filling appeared, as shown in Fig. 2d–i, by adding a small amount of DMP to the electroplating solution. Even 1 ppm DMP could cause perfect filling performance (86%) of the bath solution. Similar filling results were also obtained in other electroplating solutions using DB or JGB as a leveler. This phenomenon indicated that the synergistic effect between PEG-8000 and SPS was recovered by adding a small amount of DMP to the electroplating solution. However, when the DMP concentration was higher than 7 ppm, then filling performance of the electroplating solution declined. According to previous research, a reasonable interpretation explains this phenomenon is that excess DMP spread to the microvia, and seriously influenced the adsorption of SPS in them.

When Cl$^-$ concentration decreased from 30 ppm to 10 ppm, the filling results of microvia and surface topography of a test board surface are shown in Fig. 3 and 4, respectively. As can be seen from Fig. 4, a uniform, light and smooth copper surface appeared in a large number of copper nodules, which form a texture that traces the air bubble flow. Similar electroplating results were obtained with other plating formula having low Cl$^-$ concentrations, which may be caused by local Cl$^-$ concentrations lower than the critical concentration for steady PEG adsorption. As shown in Fig. 3, the microvia filling turns into explosive growth regardless of the concentration of DMP. This phenomenon may be attributed to the effect of accelerating SPS–Cl$^-$, which is greater than the inhibition of PEG–DMP–Cl$^-$ due to low chloride ion concentration easily leading to locally insufficient chloride ion coverage. In addition, some scholars consider that PEG–DMP–Cl$^-$ is a strong inhibitor of copper deposition. Nevertheless, because the Cl$^-$ concentration is too low, PEG–DMP–Cl$^-$ is not completely covering the copper surface, thus leading to the occurrence of this phenomenon.

### 3.2 Filling mechanism

According to the established model for simulating superfilling, SPS spread to the microvia in the presence of chloride ions and accumulated at the metal/electrolyte interface located at the bottom, where the complex copper deposition process containing a large number of reactions is displayed in the following equations:

\[
R\text{-S-S-R} + 2H^+ + 2e^- \rightarrow 2R\text{-S-H} \quad (1)
\]

\[
4R\text{-SH} + 2Cu^{2+} \rightarrow 2R\text{-S-Cu}^{2+} + 2R\text{-S-S-R} + 4H^+ \quad (2)
\]

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

\[
CuCl + e^- \rightarrow Cu + Cl^- \quad (4)
\]

### Table 1: Formulas of the microvia electroplating baths

| CuSO$_4$.5H$_2$O (g) | H$_2$SO$_4$ (M) | Cl$^-$ (ppm) | SPS (ppm) | PEG-8000 (ppm) | DMP (ppm) |
|----------------------|----------------|--------------|-----------|----------------|-----------|
| Basic 1              | 220            | 0.54         | 30        | 0              | 0         | 0         |
| Basic 2              | 220            | 0.54         | 30        | 1              | 200       | 0         |
| Basic 3              | 220            | 0.54         | 30        | 1              | 200       | 0         |
| Basic 4              | 220            | 0.54         | 30        | 1              | 200       | 0         |
| Basic 5              | 220            | 0.54         | 30        | 1              | 200       | 2         |
| Basic 6              | 220            | 0.54         | 30        | 1              | 200       | 3         |
| Basic 7              | 220            | 0.54         | 30        | 1              | 200       | 5         |
| Basic 8              | 220            | 0.54         | 30        | 1              | 200       | 7         |
| Basic 9              | 220            | 0.54         | 30        | 1              | 200       | 9         |
| Basic 10             | 220            | 0.54         | 10        | 1              | 200       | 1         |
| Basic 11             | 220            | 0.54         | 10        | 1              | 200       | 2         |
| Basic 12             | 220            | 0.54         | 10        | 1              | 200       | 3         |
| Basic 13             | 220            | 0.54         | 10        | 1              | 200       | 5         |
where R is the $(\text{CH}_2)_n\text{SO}_3^-$. The thiol group strongly adsorbs on a copper surface, forming insoluble thiol–copper complex, and the $\text{SO}_3^-\text{H}$ group can uninterruptedly surround Cu$^{2+}$ ions due to an electrostatic force, inducing these Cu$^{2+}$ ions to be dewatered. At this moment, the evaporated Cu$^{2+}$ ions can be transferred to the chloride ions and accept electrons, thus accelerating copper deposition with the assistance of chloride ion.\textsuperscript{32,34} Inversely, PEG is a nonionic surfactant which tends to interact with transition-metal ions and form a complex (Cu$^+$PEG) in the bath solution; this ability of PEG is dependent on the lone-pair electrons of oxygen. Previous research has shown\textsuperscript{35} that this complex can only adsorb on the outer Helmholtz layer of the electrode. Thus, the adsorption behavior of PEG in a bath solution without Cl$^-$ is weak. However, once Cl$^-$ exists in electroplating solution, Cl$^-$ acts as a dynamic anchor to immobilize the Cl$^-\text{Cu}^+\text{PEG}$ complex (PEG–Cu$^+$Cl$^-$), which strongly adsorbs on the inner Helmholtz layer of the cathode to inhibit copper deposition.\textsuperscript{36} The specific reaction is shown in the following equation:

\[
n\text{Cu}^+ + \text{HO}((\text{CH}_2)_n\text{O})\text{H} \rightarrow \text{HO}((\text{CH}_2)_n\text{OCu})_n((\text{CH}_2)_n\text{O})y-n\text{H}
\]

\[
n\text{Cu}^+ + n\text{Cl}^- + \text{HO}((\text{CH}_2)_n\text{O})\text{H} \rightarrow \text{HO}((\text{CH}_2)_n\text{OCuCl})_n((\text{CH}_2)_n\text{O})y-n\text{H}
\]

Therefore, this partial accelerating effect at the microvia bottom and the inhibiting effect on the copper surface led to superfilling of microvia.

### 3.3 Electrochemical analysis

In order to get insight into the interaction between additives and the influence of different DMP concentrations on microvia filling performance, potentiodynamic polarization curves and electrochemical impedance spectroscopic were implemented.

The effect of different formulas on potentiodynamic polarization curves at 298 K are shown in Fig. 5. It can be seen that deposition potential appeared at about −0.02 V using bath 1. In comparison, when PEG was added to bath 1, the deposition potential appeared as an obviously negative shift −0.13 V. This phenomenon mainly comes from PEG–Cl$^-$, which strongly adsorbs on the surface of copper to inhibit copper deposition. According to relevant research,\textsuperscript{37} if the electroplating solution contains only PEG and Cl$^-$ without an accelerator, and the deposition rate of copper at the bottom of the microvia is not constrained by mass-transfer since Cu$^{2+}$ concentration is high, then this leads to conformal deposition. This result is consistent with our filling experiment (see Fig. 2b). When 1 ppm SPS was added to electroplating baths containing PEG and Cl$^-$, the deposition potential appeared as a slightly positive shift (−0.12 V). This acceleration mainly derives from SPS–Cl$^-$. However, the plating formulas also lead to conformal deposition (see Fig. 2c) due to the coverage increment of SPS on the copper surface. Therefore, excessive SPS correspondingly reduced the inhibiting effect induced by PEG–Cl$^-$ on copper deposition, resulting in a relatively fast copper deposition on the copper surface and a reversely slow copper deposition in the microvia.

With addition of DMP, the deposition potential of copper encountered an obviously negative shift (−0.21 V), and along with the cathode current density, progressively decreased. The result showed that the addition of DMP also could increase cathodic polarization even when SPS, PEG, and Cl$^-$ simultaneously exist in a bath solution, and this phenomenon became more serious with increasing DMP concentration. According to results of the AFM, XPS, and theoretical calculation (in Sections 3.6, 3.7 and 3.8), the parallel adsorbing of DMP to copper surface is through the pyrimidine ring. That increased the contact area between DMP and copper surface, and further increased the diffusion of copper ions to the cathode surface space steric hindrance. This phenomenon showed that DMP further increased an inhibiting effect caused by PEG–Cl$^-$ and co-inhibition of copper deposition. At the same time, a lot of SPS accumulated via bottom and accelerated copper deposition with the assistance of chloride ion, and thus caused different deposition rates between the copper surface and the bottom of the microvia. Hence, perfect bottom-up filling was obtained (see Fig. 2d–h). However, as the DMP concentration was increased to 9 ppm, the inhibition of the bath solution became very strong and more DMP molecules spread to the microvia, quenched catalyst activity of SPS, and affected copper deposition.\textsuperscript{38} Therefore, the microvia filling performance declined (see Fig. 2i).

As shown in Fig. 6, Nyquist plots with different formulas can be divided into two parts according to frequency. High frequency impedance corresponding a depressed capacitive
loop, which is related to double-layer capacitance ($C_{dl}$), and charge transfer resistance ($R_{ct}$), indicating\textsuperscript{40} the inhomogeneities and roughness of the copper working electrodes. The low frequency impedance is straightforward, namely the Warburg impedance ($W$), which can be explained by dissolved cuprous chloride species transferring from the copper surface to the electroplating solution or to soluble oxygen invading the copper surface.\textsuperscript{40} As can be seen from the Nyquist plots, the shape, type, and number of the capacitive loops almost stayed the same with addition of SPS, PEG, and DMP, but the diameters were different.\textsuperscript{8} Results showed that addition of the additives slightly changed the electrodeposition mechanism.\textsuperscript{41,42}

Usually, corresponding electrochemical impedance data was obtained by employing the equivalent circuits of Zsimpwin software and fitted data are listed in Table 2. This type of equivalent circuit is shown in Fig. 7, including solution resistance ($R_s$), charge transfer resistance ($R_t$), Warburg impedance ($W$), resistance of protective film formed on copper surface ($R_f$), and constant phase angle elements (CPE).

As can be seen from Table 2, the values of both $R_s$ and $R_t$ remain nearly unchanged with the addition additives. However, the $R_{ct}$ values changed a lot. As is well known, the copper deposition contained two steps:\textsuperscript{43}

$$\text{Cu}^{2+} + e^- \rightarrow \text{Cu}^+ \quad (7)$$

$$\text{Cu}^+ + e^- \rightarrow \text{Cu} \quad (8)$$

Among them, reaction (7) is deemed to be the rate-determining step. The size of $R_{ct}$ usually is used to assess the suppression and acceleration ability of additives on copper deposition. From Table 2, the corresponding $R_{ct}$ value is 62.79 $\Omega$ only when Cl\textsuperscript{-} is added to a bath solution. As is well known, the slight extent of depolarization caused by adding Cl\textsuperscript{-} alone will accelerate copper surface deposition. The reaction mechanism was expressed as the above eqn (3) and (4). In addition, the copper deposition rate at the opening of the microvia is more than that of the bottom due to the uneven distribution of local current density. Therefore, a void occurred in the microvia interior (see Fig. 2a). In contrast, the $R_{ct}$ value increased to 144 $\Omega$ when the electroplating baths included Cl\textsuperscript{-} and PEG. Many investigations\textsuperscript{44} have demonstrated that Cl\textsuperscript{-} acts as an active site of PEG adsorption on the electrode surface to form a complex (PEG–Cu\textsuperscript{2+}–Cl\textsuperscript{-}), which inhibited copper surface deposition. However, copper deposition rate of the microvia bottoms is too slow due to the low current density. Therefore, the conformal deposition appeared. After adding 1 ppm SPS to plating solution with PEG and Cl\textsuperscript{-}, the $R_{ct}$ value decreased to 109.3 $\Omega$. This phenomenon showed that SPS–Cl\textsuperscript{-} destroyed the suppression ability of PEG–Cl\textsuperscript{-} on the copper surface, and accelerated copper surface deposition. At this moment, the copper deposition rate also relatively increased due to accumulated SPS–Cl\textsuperscript{-} at the microvia bottom. Therefore, the deposition rate difference between the bottom of microvia and copper surface are nearly the same, and the conformal deposition was also obtained (see Fig. 2c).

As expected, the $R_{ct}$ values increased with DMP added to an electroplating solution with SPS, Cl\textsuperscript{-}, and PEG. This phenomenon had shown a suppression effect of the DMP on the copper surface and this effect was more remarkable with DMP concentrations up to a 7 ppm saturated value. This result can be considered to perfect DMP coverage on the copper surface during electroplating, and almost completely prevented further copper deposition with the assistance of PEG. Hence, bottom-up filling was obtained. However, as the DMP concentration was further increased, the ability of inhibiting the electroplating solution was too strong to obtain good filling performance (see Fig. 2i). All experimental results are consistent with the potentiodynamic polarization curve measurements.

### 3.4 Reliability measurements

Fig. 8 shows the cross-sections of microvias after a thermal shock test and thermal cycle test, respectively. As can be seen from

\begin{table}[h]
\centering
\begin{tabular}{|c|c|c|c|c|c|}
\hline
\textbf{Basic} & \textbf{$R_s$ (\text{\Omega cm}^2)} & \textbf{$R_t$ (\text{\Omega cm}^2)} & \textbf{$R_{ct}$ (\text{\Omega cm}^2)} & \textbf{CPE} & \textbf{$W$} \\
\hline
1 & 0.4722 & 1.722 & 62.79 & 4.796 & 0.496 & 0.280 \\
2 & 0.5327 & 2.067 & 144.0 & 2.437 & 0.588 & 0.076 \\
3 & 0.5850 & 3.451 & 109.3 & 3.459 & 0.544 & 0.087 \\
4 & 0.5765 & 5.456 & 236.7 & 1.060 & 0.630 & 0.078 \\
5 & 0.5771 & 4.409 & 335.3 & 1.024 & 0.677 & 0.041 \\
6 & 0.6112 & 6.924 & 346.8 & 0.862 & 0.688 & 0.033 \\
7 & 0.5511 & 5.374 & 363.0 & 1.004 & 0.696 & 0.037 \\
8 & 0.5730 & 6.084 & 394.9 & 0.892 & 0.696 & 0.031 \\
9 & 0.6942 & 7.148 & 397.2 & 0.874 & 0.678 & 0.031 \\
\hline
\end{tabular}
\caption{Impedance parameters of copper in electroplating solution with different formulas}
\end{table}
Fig. 8, copper film peeling off, foaming phenomenon, and plate blasting after reliability tests did not appear. This means that the reliability of interconnections of PCBs is good and in accord with the quality standard in IPC-A-600H-2010.

3.5 Effect of different concentrations of DMP on copper deposits morphology and crystalline orientation

The surface morphology of copper deposition obtained by adding different concentrations of DMP to the electroplating solution is shown in Fig. 9. Only two concentrations, namely, 1 and 2 ppm, can obtain a uniform, compact, and smooth surface, as shown in Fig. 9a and b. When DMP concentration increased to 3 ppm, a small quantity of acicular crystals appeared on the copper surface and became denser with increasing DMP concentration. This means that higher DMP concentrations would destroy the synergistic effect on a copper surface. A similar plating result was obtained with the electroplating additives system using DB as a leveler.

The influence of different concentrations of DMP on the crystalline structure of the copper deposits was characterized by XRD. As shown in Fig. 10, the diffraction pattern for the electroplating film exhibits four important peaks at 43.2°, 50.3°, 73.9°, and 90.1°, which correspond to (111), (200), (220) and (311) of copper, respectively. Previous research showed that the value of intensity ratios (111)/(220) decreased when leveler was added to an electroplating solution, so we do not discuss that here. From Fig. 10, the intensity of (111) increased and the intensity of (220) decreased with increasing DMP concentrations. When the concentration of DMP increased to 5 ppm, the value of ratios I_{(111)}/I_{(220)} reached a saturated value and the (111) orientation became absolutely dominant. A similar phenomenon was also obtained in another electroplating additive formula. Jiang et al. confirmed that higher EDTA concentrations added to an electroplating solution caused the (111) orientation to be dominant.

3.6 Interaction between DMP and the copper surface of PCB

Two-dimensional (2D) and three-dimensional (3D) AFM graphs of the copper surface washed and polished by alumina polishing powder and treated without DMP are shown in Fig. 11a and b. Note that there were a few clear tiny scratches which derived from the polishing process on the copper surface. Furthermore, that part of the position of the copper surface was very angular. Fig. 11c and d presented AFM graphs of copper surfaces which have been soaked in 1 ppm DMP aqueous solution. Compared with Fig. 11a and b, the tiny scratches on the copper surface were not so obvious, and the angular degree of the entire copper
surface was decreased. Mean surface roughness of the copper surface decreased from 8.941 nm (polished copper surface) to 6.403 nm. This phenomenon could be attributed to DMP adsorbed on the surface of the copper and it changed the copper surface morphology.49,50

The high-resolution XPS survey scan spectra of copper surface is shown in Fig. 12a, which taken over a large binding energy region manifested the presence of carbon, nitrogen, oxygen, sulfur, and copper species. The binding energy of C1s peak at 284.8 eV was used as a standard value. The specific XPS peaks of N1s and S2p are shown in Fig. 12b and c, respectively. From Fig. 12b, we can observe a distinct peak at the binding energy of 399.7 eV, which could be attributed to the C–N or C≡N bonds in DMP.51,52 As shown in Fig. 12c, we can also observe a peak at the binding energy of 162.7 eV, corresponding to the thiolate bonds in DMP.53,54 The N1s peak and S2p peak show the authentic existence of DMP molecules on the copper surface. Combined with the results of AFM and XPS patterns, it indicated that an ordered and compacted adsorption layer of DMP was obtained after the copper surface was soaked in the DMP aqueous solution. Therefore, DMP exhibited an effective levelling ability and is beneficial to microvia filling.

3.7 Quantum chemical calculations

Quantum chemical calculations play a significant role to research interactions between additive molecules and the metal surface. Therefore, to gain insight into the adsorption behaviors of DMP on a copper surface, density functional theory (DFT) was implemented. Optimized geometrical configuration and density distributions of the frontier molecular orbital of DMP are shown in Fig. 13. Corresponding quantum chemical parameters, which include frontier molecular orbital energies ($E_{\text{HOMO}}$, $E_{\text{LUMO}}$), energy gap ($\Delta E = E_{\text{LUMO}} - E_{\text{HOMO}}$), and dipole moment ($\mu$), are collected in Table 3.

![Molecular Structure](image)

Table 3 | Quantum chemical parameters for DMP
| $E_{\text{HOMO}}$ (eV) | $E_{\text{LUMO}}$ (eV) | $\Delta E$ (eV) | $\mu$ (D) |
|-----------------|-----------------|------------|------------|
| DMP             | −0.243          | −0.046     | 0.197      | 3.41       |

According to previous reports55,56 the transition state was formed because of an interaction between the frontier molecular orbital of the reaction reagent. It is generally accepted that the $E_{\text{HOMO}}$ is usually relevant to electron donating ability, and the greater value of $E_{\text{HOMO}}$ indicated the molecular donating electrons ability became stronger. Conversely, the $E_{\text{LUMO}}$ is often associated with the ability of molecules to accept electrons and the lower value of $E_{\text{LUMO}}$ indicated the ability of molecules to accept electrons became stronger.53,54 Adsorption ability of the organic molecules on the metal surface was increased with the decreasing of $E_{\text{LUMO}}$ or increasing of $E_{\text{HOMO}}$.56-59 Therefore, $\Delta E$ is generally associated with chemical stability of the leveler and could be used to judge adsorption ability.60 In addition, the dipole moment also plays a significant role to research the adsorption mechanism of a leveler.61 According to a previous investigation62 a high $\mu$ would lead to strong adsorption of the leveler on a copper surface due to electronic force. From Table 3, the molecule of DMP has a low value of $\Delta E$ and a high value of $\mu$, indicating that DMP could strongly adsorb on the copper surface and result in increased cathodic potential and charge transfer resistance; thus, it inhibited copper surface deposition. Besides, as can be seen from Fig. 13, the HOMO of DMP is distributed on the −SH group, which means that the −SH was an active group donating electrons to low-energy empty molecular orbitals of copper to form a coordinate/covalent bond. The LUMO of DMP is located on N atoms, indicating N atoms were an active site for electrons from Cu to attack. These results are in accordance with the AFM and XPS measurements.

3.8 Molecular dynamic simulation

For a long time, MD simulation, which has been widely used, was a useful tool to study the interaction between the additives and copper.63 Therefore, MD simulations have been established to pattern the adsorption mechanism of DMP on the surface of Cu (111)44 in this article. The top and side views of the optimized equilibrium configuration of DMP are shown in Fig. 14. It can be seen that the DMP molecule can be parallelly adsorbed on the copper surface through the pyrimidine ring, increasing the
degree of surface coverage and inhibiting copper deposition. This parallel mode fully demonstrated our conjecture for the above experimental results. Moreover, the value of adsorption energy between Cu (111) surface and the DMP molecule obtained from MD simulation is 250.5 kJ mol⁻¹. This high absolute value indicated the strong adsorption ability between DMP and copper surface.⁶⁵,⁶⁶ These results further proved that the adsorption of DMP on copper surface increased cathodic potential and charge transfer resistance, inhibited the copper surface electrodeposition, and increased filling performance of microvia. This result is similar to Janus Green B, which was an effective leveler proved by several scholars.⁶⁷

4. Conclusions

DMP was used as a leveler in microvia electroplating copper filling experiments. The main results of the study are summarized as follows:

After plating copper filling experiments, a void can be clearly seen in the middle of the microvia because no organic additives were present in the bath solution. In contrast, when the electroplating solution contained only PEG and Cl⁻ without an accelerator, the deposition rate of copper at the bottom of the microvia was not constrained by mass-transfer, which lead to conformal deposition. After 1 ppm SPS was added to the electroplating solution, which correspondingly reduced the inhibiting effect caused by PEG–Cl⁻ on copper deposition, it resulted in a relatively fast copper deposition on the copper surface and a reversely slow copper deposition in the microvia. Therefore, conformal deposition was also obtained. That means only added DMP to an electroplating solution can obtain a perfect filling effect. However, excess DMP added to an electroplating solution caused too strong of an inhibiting ability, quenched catalyst activity of SPS, and affected copper deposition rate, which resulted in the filling performance of electroplating solution to decline. In addition, if only 10 ppm Cl⁻ is added to an electroplating solution with SPS, PEG, and DMP, then a stable adsorption ability of PEG and DMP can’t be obtained. Inversely, SPS can facilitate copper deposition. Therefore, the test board surface appeared with a large number of copper nodules, and the microvia filling had explosive growth.

With addition of SPS, the cathodic polarization shift to a more positive value and the value of $R_{ct}$ decreased. However, if DMP or PEG are added to the bath solution, the cathodic polarization and charge transfer resistance are increased with the assistance of Cl⁻, indicating that they could inhibit copper electrodeposition.

The FE-SEM images and XRD measurements indicated that a uniform, compact, and smooth surface can be obtained; when lower concentrations of DMP were added to electroplating solution, the (111) orientations became dominate. However, when higher DMP concentrations were added to the electroplating solution, acicular crystals appeared on the copper surface, and the preferential orientations of electroplating copper transformed from (220) to (111).

AFM and XPS measurements demonstrated DMP adsorbed on the cathodic surface and inhibited copper surface deposition. Hence, it exhibited an effective levelling ability and is beneficial to microvia filling.

The results obtained from quantum chemical calculations and MD simulation indicated N atoms and –SH of the DMP molecule were active groups that can be parallely adsorbed on the surface of copper through the pyrimidine ring due to electronic forces and increased the degree of surface coverage and inhibited copper deposition.

Conflicts of interest

There are no conflicts to declare.

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