Journal of Visualized Experiments

Accumulation and Analysis Methods of Cuprous Ion in Copper Sulfate Plating Solution --Manuscript Draft--

Article Type:	Invited Methods Article - JoVE Produced Video
Manuscript Number:	JoVE59376R1
Full Title:	Accumulation and Analysis Methods of Cuprous Ion in Copper Sulfate Plating Solution
Keywords:	Copper plate; solution; cuprous ion; monovalent copper ion; BCS; absorbance; concentration; plating film; surface structure; quality; evaluation; on-site
Corresponding Author:	Toshiaki Koga
	JAPAN
Corresponding Author's Institution:	
Corresponding Author E-Mail:	t-koga@aist.go.jp
Order of Authors:	Toshiaki Koga
	Yoshitaro Sakata
	Nao Terasaki
Additional Information:	
Question	Response
Please indicate whether this article will be Standard Access or Open Access.	Open Access (US\$4,200)
Please indicate the city, state/province, and country where this article will be filmed . Please do not use abbreviations.	Tosu-shi Saga 841-0052, JAPAN

TITLE:

Accumulation and Analysis of Cuprous Ions in a Copper Sulfate Plating Solution

234

1

AUTHORS:

5 Toshiaki Koga, Yoshitaro Sakata, and Nao Terasaki

6 7

Advanced Manufacturing Research Institute, Department of Electronics and Manufacturing, National Institute of Advanced Industrial Science and Technology (AIST), Tosu-shi Saga, Japan

8 9 10

Corresponding Author:

11 Toshiaki Koga (t-koga@aist.go.jp)

12 13

Email Addresses of Co-authors:

14 Yoshitaro Sakata (yoshitaro-sakata@aist.go.jp) 15 Nao Terasaki (nao-terasaki@aist.go.jp)

16 17

KEYWORDS:

copper plate, solution, cuprous ion, monovalent copper ion, BCS, absorbance, concentration, plating film, surface structure, quality, evaluation, on-site

20 21

22

23

SUMMARY:

Here, accumulation of cuprous ions in a copper sulfate plating solution in a model experiment and an analysis based on quantitative measurements are described. This experiment reproduces the accumulation process of cuprous ions in the plating bath.

242526

27

28

29

30

31

32

33

34

35

36

37

38

39

40

41

42

ABSTRACT:

Knowledge of the behavior of cuprous ions (monovalent copper ion: Cu(I)) in a copper sulfate plating bath is important for improving the plating process. We successfully developed a method to quantitatively and easily measure Cu(I) in a plating solution and used it for evaluation of the solution. In this paper, a quantitative absorption spectrum measurement and a time-resolved injection measurement of Cu(I) concentrations by a color reaction are described. This procedure is effective as a method to reproduce and elucidate the phenomenon occurring in the plating bath in the laboratory. First, the formation and accumulation process of Cu(I) in solution by electrolysis of a plating solution is shown. The amount of Cu(I) in the solution is increased by electrolysis at higher current values than the usual plating process. For the determination of Cu(I), BCS (bathocuproinedisulfonic acid, disodium salt), a reagent that selectively reacts with Cu(I), is used. The concentration of Cu(I) can be calculated from the absorbance of the Cu(I)-BCS complex. Next, the time measurement of the color reaction is described. The color reaction curve of Cu(I) and BCS measured by the injection method can be decomposed into an instantaneous component and a delay component. By analysis of these components, the holding structure of Cu(I) can be clarified, and this information is important when predicting the quality of the plating film to be produced. This method is used to facilitate the evaluation of the plating bath in the production line.

43 44

INTRODUCTION:

As printed circuit boards become denser and multilayered, management of plating solutions during the manufacturing process becomes more important to maintain product quality. In copper sulfate electroplating, the monovalent copper ion (cuprous ion: Cu(I)) has been determined to be one of the main causes of the large roughness and dull finish of the copper plating surface. The behavior and role of Cu(I) in the plating process¹⁻⁵, the effect of each additive, and the holding structure⁶⁻⁸ have been investigated. It is necessary to analyze Cu(I) in the plating solution, but it was difficult to quantify its concentration because of the instability of Cu(I) in an aqueous solution. Therefore, the on-site analysis of Cu(I) in the plating bath is an effective tool for controlling the plating solution.

545556

57

58

59

60

61

62

63

64

65 66

67

45

46

47

48

49

50

5152

53

We performed colorimetric analysis using an aqueous chelating reagent, BCS (bathocuproinedisulfonic acid, disodium salt), to establish on-site quantitative analysis of Cu(I) in a copper sulfate plating solution. The BCS can be used to quantify the Cu(I) concentration in the aqueous solutions⁹⁻¹¹. The cuproine type color reaction reagent, which has been conventionally used for the determination of Cu(I), is hydrophobic and extraction with alcohol is necessary. It was shown that BCS is hydrophilic and can directly measure Cu(I) in an aqueous solution. Two molecules of BCS coordinate to one Cu(I) to form 1:2 complexes that absorb visible light at wavelengths between 400 and 550 nm (See Figure 1). We established a method to determine the concentration of Cu(I) in the plating solution from the measurement of the absorbance of the Cu(I)-BCS complex^{12,13}. In the first part of this protocol, a method of accelerating the Cu(I) formation in a copper sulfate plating solution in a model experimental system and the quantitative measurement of the Cu(I) concentration in a plating solution are described. This is fundamental to clarify the process of formation and accumulation of Cu(I) in the plating bath.

68 69 70

71

72

73 74

75

Further, it was shown that the color reaction of Cu(I) and BCS can be divided into rapid reaction components and relatively slow reaction components. This increases the uncertainty in the absorbance measurement. To overcome this problem, we developed a method of measuring reaction curves by an injection method^{14,15}. The second part shows the measurement of Cu(I) based on the injection method. By analyzing the components obtained by the injection method, it is possible to approximate the understanding of the Cu(I) formation mechanism and holding structure in solution.

76 77 78

79

80

81

82

83

Conventionally, it has been claimed that Cu(I) in a plating solution is instantly oxidized to cupric ions (Cu(II)). We have confirmed that there are several millimoles (mmol/L) of Cu(I) in the plating bath of the production line¹². According to this experiment method, the accumulation of Cu(I) similar to the plating bath can be reproduced even in the beaker of the laboratory. This is a fundamental technology to elucidate the Cu(I) production and accumulation process in a copper sulfate electroplating solution, which was unknown¹⁴. Furthermore, by controlling Cu(I) in the plating solution, it is also possible to predict the effect of Cu(I) on the quality of the plating film¹⁵.

84 85

PROTOCOL:

868788

NOTE: Please check all related material safety data sheets (MSDS). Please wear protective

equipment when experimenting with the copper sulfate plating.

899091

1. Preparation of the copper sulfate plating solution

92

NOTE: The copper sulfate plating aqueous solution is prepared by combining sulfuric acid (0.5 mol/L), copper sulfate (0.4 mol/L), chlorine (Cl, 1.41 mmol/L), polyethylene glycol (PEG; MW 4000: 0.025 mmol/L), bis(3-sulfopropyl) disulfide (SPS, 0.003 mmol/L), and Janus Green B (JGB, 0.004 mmol/L) in pure water.

97

98 1.1. Place a stir bar in a 1 L beaker and pour in 600 mL of pure water. Add sulfuric acid (95.0%: 49.04 g) in small portions while stirring. Leave it until the solution cools down.

100

1.2. Add copper sulfate (99.5%: 99.876 g) to the solution little by little. Stir for 30 min.

102

1.3. Add 23.7 mL of hydrochloric acid (0.02 mol/L), 0.1 g of polyethylene glycol, 1 mL of 1 mg/L
 SPS solution, and 1 mL of 2 mg/L JGB solution.

105

1.4. Transfer the solution to a volumetric flask (1 L). Add pure water and adjust to 1 L. Transfer the copper sulfate plating solution to a polyethylene container and store it at room temperature in the dark.

109

2. Formation of the Cu(I) in the plating solution

110111

2.1. Pour 150 mL of the copper sulfate plating solution into a 200 mL beaker. Put the stir bar in the beaker and stir at 500 rpm. Leave the plating solution in advance at room temperature (23 °C ± 1 °C) for 1 hour.

115

2.2. Insert a tube into a beaker and let nitrogen flow (about 85 mL/min). Deoxygenate the plating solution with nitrogen gas for over 30 min.

118

2.3. Cut the 0.3 mm thick copper plate with metal shears to 9.5 cm x 2 cm dimensions. Cut the platinum plate with a thickness of 0.1 mm in the same way.

121

2.4. Wash the copper plate and the platinum plate with ethanol and rinse with pure water. Dry
 with nitrogen gas.

124

2.5. Attach the copper plate and platinum plate to the fixing jig, insert it inside the beaker and fix
 it. The immersed area of each plate to the plating solution is 4 x 2 cm² (See Figure 2).

127

NOTE: The jig consists of an acrylic beaker fixing part (**Figure 3** (1)) and metal electrode parts (**Figure 3** (2)). The electrode part consists of the parts to fix the plate, and the part connects to the cord from the power supply.

131

2.6. Connect the electrode (anode) of the copper plate to the positive end of the power supply

133 (Figure 3 (3)), and the electrode of the platinum plate (cathode) to the negative end of the power supply (Figure 3 (4)).

135

2.7. Turn on the power supply at a constant current of 1.0 A (current density: 62.5 mA/cm²). Cu(I) is formed in the plating solution according to the electrolysis time, and Cu(I) concentration (accumulated amount) is maximized in about 10 min.

139

NOTE: If the plate is inserted while the stirrer rotates, the plating solution may scatter and the beaker may light up. Please install the jig before turning on the power to avoid danger.

142

2.8. Turn off the power after 10 min and stop the stirrer. Leave it for about 10 min until the particles settle.

145

3. Quantitative measurement of the Cu(I)

146147

3.1. Prepare the BCS solution (10⁻² mol/L) by dissolving 0.36 g of the molecule in 100 mL of pure water. Stir the solution and dissolve the BCS in an excess amount relative to the monovalent copper. Store the BCS solution in a light-proof container and store the container in the dark.

151

NOTE: In the measurement, the BCS concentration in the sample solution is adjusted to 1,000 times or more the Cu(I) concentration.

154

155 3.2. Add 60 mL of acetic acid (1 mol/L) and 25.2 mL of NaOH solution (1 mol/L) to 120 mL of pure water to prepare a neutralizing solution (buffer solution).

157158

3.3. Put a stir bar in the absorption measurement cell (optical path length: 1 cm) and pour in 2.5 mL of neutralization solution and 219 µL of BCS solution.

159160161

3.4. Mix in 22 µL of plating solution sample (step 2.9). Stir for 20 min.

162

NOTE: In order to ensure that the function of BCS is normal, the pH of the sample solution to be measured should not fall below 4. BCS selectively forms a complex with Cu(I). The Cu(I)-BCS complex absorbs in the visible region (400 to 550 nm), and the neutralizing solution develops an orange color (Figure 4).

167

3.5. Measure the absorption spectra of the sample solution (3.4) with an UV/vis spectrophotometer (wavelength range: 400–600 nm) (Figure 5e).

170

NOTE: There are no constrained measurement apparatus and conditions, and it is desirable to make them identical in one experiment series.

173

174 3.6. Calculate the concentration of Cu(I) using the Lambert-Beer law:

175

176 $A = \varepsilon I c$

where A is absorbance, L is the optical path length, ε is the molar absorption coefficient (BCS: 1.2 \times 10⁴ at 485 nm), and c is the molar concentration (mol/L) of the solute.

NOTE: Because the optical path length is 1 cm, the Cu(I) concentration in the cell is simply the absorbance divided by the molar extinction coefficient. The value obtained by multiplying the ratio 125 (the fold dilution with the neutralizing solution) is the Cu(I) concentration of the plating solution.

4. Injection measurement of Cu(I) and BCS color reaction curves

4.1. Use a UV/vis spectrophotometer with time measurement function of more than 20 min for injection measurement. The spectrometer should have a sample chamber cover with a syringe port (**Figure 6** left) and a thermostat cell holder with a stirrer.

4.2. Use a square cell of 1 cm x 1 cm for the absorbance measurement. Put s stir bar in the absorption cell.

4.3. Pour 2.5 mL of the neutralized solution prepared in 3.2 and 219 μ L of the BCS solution prepared in 3.1 into the cell. Maximize stirrer rotation speed.

4.4. Set the measurement time to 1270 s in the time measurement mode at 485 nm and start. One min after starting, inject 22 μ L of the plating solution sample (2.9) with a pipette from the syringe port of the chamber cover. Reaction curves of Cu(I) and BCS will be acquired (**Figure 6** right).

REPRESENTATIVE RESULTS:

The concentration of Cu(I) in the plating solution can be determined from the absorbance at 485 nm of Cu(I)-2BCS chelate. **Figure 5** shows the absorption spectra of the plating solutions that were electrolyzed for 0, 4, 6, 8 and 10 min. The Cu(I) concentration tends to increase from 0 to 10 min depending on the electrolysis time. However, as a result of the time-resolved measurement, a delay component appeared in addition to the instantaneous component in the reaction between BCS and Cu(I). This reduces the signal-to-noise ratio (S/N ratio) of the absorbance value and prevents accurate determination of Cu(I) concentration. It is preferable to use the injection method to determine the Cu(I) concentration, because the change in absorbance caused by the injection of plating solution is measured by time decomposition (**Figure 6**).

Information on the Cu(I) holding structure in the plating solution is obtained by numerical analysis of the reaction curve. In general, Cu(I) is quickly oxidized to Cu(II) in an aqueous solution; but in the plating solution it is considered to be stabilized by forming a complex with an additive (especially PEG)¹⁴. The reaction curve reflects the chelation process of Cu(I) and BCS. The reaction curve is composed of a component that increases immediately after the plating solution injection and a component that slowly increases over several tens of min. These components suggest that

there are multiple holding structures of Cu(I) in the plating solution. Characteristics of the plating solution involved in Cu(I) can be evaluated by analyzing the reaction curve. Assuming that the reaction of Cu(I) with BCS is a first order reaction with respect to the Cu(I) concentration, we obtained the following reaction kinetics of the absorbance, At:

226 At = A0 + AL $[1 - \exp(-t/TL)]$

t is the time from the start of measurement, A0 corresponds to a component that reacts instantaneously (absorbance at t=0) and AL corresponds to a component that reacts slowly (At - A0). TL is the time constant of the AL component. To simulate the color reaction curve, we applied the formula to the original analysis software (software may be commercially available)^{13,15}. A curve simulating the change in the absorbance of the color reaction of the electroplating solution is shown in **Figure 7**. From the simulation, the parameters (A0, AL, TL) related to Cu(I) accumulation are quantified. The simulation results in this figure were A0 = 0.053, AL = 0.098, TL = 13.6 min, and $r^2 = 0.998$. **Figure 8** (graph) plots the simulation value A0 in the plating solution that was electrolyzed for different times. Although the value of A0 did not change greatly until 4 min of electrolysis, an increase corresponding to electrolysis time was seen from 6 min to 10 min.

Plating was carried out on a copper substrate for 10 min with the electrolysis solutions to investigate the effect of Cu(I) on the quality of the copper plating such as roughness and morphology. **Figure 8** shows the SEM (Scanning Electron Microscope) images of the film surface structure deposited with electrolysis solutions. The film structure at 0 min and at 4 min of electrolysis plating are nearly indistinguishable. There are fine particles adsorbed densely with a size of several tens of nanometers and a smooth surface morphology. After 6 min of electrolysis plating, there is some swelling on the surface. After 10 min of electrolysis plating, there is a large chunky roughness.

FIGURE LEGENDS:

- **Figure 1: Structure and absorption spectrum of Cu (I)-BCS complex.** Fresh copper sulfate plating solution and electrolysis solution. Since Cu(I) is accumulated in the plating solution by electrolysis, the absorption spectrum of Cu(I)-BCS complex is observed in the electrolysis plating solution sample.
- Figure 2: Schematic diagram of the equipment for electrification experiment (left) and representative conditions of the electrolysis experiment (right).
- Figure 3: Picture of a combinations of parts to be energized in the experiment. Attach the jig with the electrode plate to the glass beaker and connect it to the power supply. (1) Acrylic beaker fixing part, (2) metal electrode parts, (3) copper plate electrode (anode), and (4) platinum plate electrode (cathode).
- Figure 4: Absorption measurement of Cu(I). Absorption measurement procedure (left) and

photos of sample solution (right). Fresh copper sulfate plating solution (blue) and electrolysis solution (orange). Since Cu(I) is accumulated in the plating solution by electrolysis, it is colored orange in the electrolysis plating solution sample.

Figure 5: Absorption spectra of Cu(I)-BCS in electrolysis solutions. Electrolysis time: (a) 0, (b) 4, (c) 6, (d) 8, and (e) 10 min. Since the absorbance of Cu(I)-BCS generally increases as the electrolysis time becomes longer, it is considered that the amount of Cu(I) accumulated in the plating solution is increased. This figure is a modification of Figure 2 of Koga et al. 2018¹⁵.

Figure 6: Injection measurement. Left: Picture of chamber cover. There is a syringe port at the top of the cell; insert a pipette there and inject sample solution. Right: Reaction curve of plating solution which was electrolyzed at 1.0 A for 10 min. A sharp increase in absorbance immediately after injection and a gentle increase are clearly observed.

Figure 7: Simulation of absorbance of plating solution (1.0 A, 10 min). ♦: measured point, solid line: fitting curve. This figure is a modification of Figure 4 of Koga et al. 2018¹⁵.

Figure 8: Deposition versus electrolysis time. (Graph) Normalized absorbance fitting parameters are plotted against electrolysis time, AO. (Pictures) SEM images of the plating film surface that were deposited in each electrolysis solution (times above pictures are electrolysis times).

DISCUSSION:

Figure 2 schematically shows a system for electrolysis experiment. The jig is an ordered item, which consists of an acrylic part to be fixed to beakers and metal parts for attaching plates and for connecting with the power supply. By this mechanism, the immersion area of the plates becomes constant, and the relationship between the current value and the current density is kept constant. In our conditions, immersion is 4 cm x 2 cm, and the current density will be 62.5 mA/cm² with a current of 1 A. In the accumulation procedure of Cu(I), a copper plate is attached to the anode and a platinum plate is attached to the cathode. In order to increase the accumulation efficiency of Cu(I), it is preferable to deoxidize the plating solution with nitrogen gas beforehand.

Quantitative measurement of Cu(I) consists of a simple procedure. Pour the neutralization solution and BCS solution into the cell and mix the plating solution (**Figure 4**). It is necessary to stir for more than 20 min until Cu(I) and BCS react sufficiently. This is to ensure the accuracy of the measurement by sufficiently advancing the reaction. If Cu(I) is contained in the plating solution, the sample solution appears orange and an absorption spectrum having a peak at 485 nm is obtained. Changes in solution color due to the complex formation were dramatic and surprised many copper plating technicians.

It is confirmed that Cu(I) accumulates in the solution when a current is passed through the copper sulfate plating solution (**Figure 5**). The absorption spectrum shows the shape of the Cu(I)-BCS complex, which is suitable for calculating the Cu(I) concentration from the absorbance at 485 nm. Although the current value is arbitrary, Cu(I) is hardly accumulated at a current value of 0.2 A,

and a higher current value is required. Although the accumulation amount of Cu(I) tends to increase with electrolysis time, it is saturated by excessive current (for example, electrolysis for more than 10 min at 1.0 A). The accumulation amount of Cu(I) increased by electrolysis for 10 min when the current value was 0.5 to 1.0 A¹⁴. When an excessive current flowed (for example, at 1.0 A for 20 min), the Cu(I) concentration decreased. This is thought to be related to the formation of copper particles due to the progress of the disproportionate reaction.

The reaction of Cu(I) and BCS in the plating solution has multiple time components, which often make the accurate determination of the concentration difficult. In order to solve this problem, an injection measurement is desirable (**Figure 6**). In this measurement, the absorption intensity of the Cu(I)-BCS complex is acquired as a changed amount from the baseline before injection of the plating solution, so it can be determined more accurately. In addition, since the reaction curve can be simply numerically analyzed, the concentration can be known with high accuracy even if the reaction is not completed. The components of the reaction curve are thought to reflect the retention structure of Cu(I) in the plating solution¹⁴.

It is important to model the holding structure of Cu(I) in the plating solution against the assertion that Cu(I) in the plating bath instantaneously oxidizes Cu(II). We propose the following model from analysis of characteristics of the current amount, formation, and accumulation of Cu(I). A portion of the Cu(I) eluted from the copper plate is retained in solution in the form of a Cu(I)-PEG complex. In early stages of the complex formation, chloride ions are thought to play a role as a temporary stabilizer for Cu(I)^{6,8}. Cu(I) coordinated to PEG is incorporated inside the three-dimensional structure, and it is in a hydrophobic environment. When the formation of Cu(I) is promoted, excess Cu(I) is coordinated to the surface of the PEG and may be in the vicinity of the liquid. Since Cu(I) on the surface reacts promptly with BCS, it will reflect the A0 component of the reaction curve. Since the Cu(I) inside the PEG is protected from BCS attack, it has a slow AL component. It has been pointed out that the A0 component mainly influences the quality of the plating film¹⁵. This information is important for management of the plating solution.

By accelerating the denaturation of the plating solution and verifying the accumulated Cu(I) concentration and the holding structure, it is possible to clearly characterize the plating solution. This is important not only for understanding the plating process but also for predicting the quality of the plating film to be produced. From the verification of the SEM image, it was shown that the Cu(I) concentration, especially the AO component, is strongly involved in the generation of the roughness of the plating film (**Figure 7**). On-site measurement of Cu(I) gives new indications for the management of plating baths.

This research can contribute to the management of the plating bath based on optical measurement. We aim to develop a system that can evaluate the state of the plating bath on the production line on-time and in situ.

DISCLOSURES:

We have nothing to disclose.

ACKNOWLEDGEMENTS:

We thank Miss. Hirakawa for her great contribution to this research.

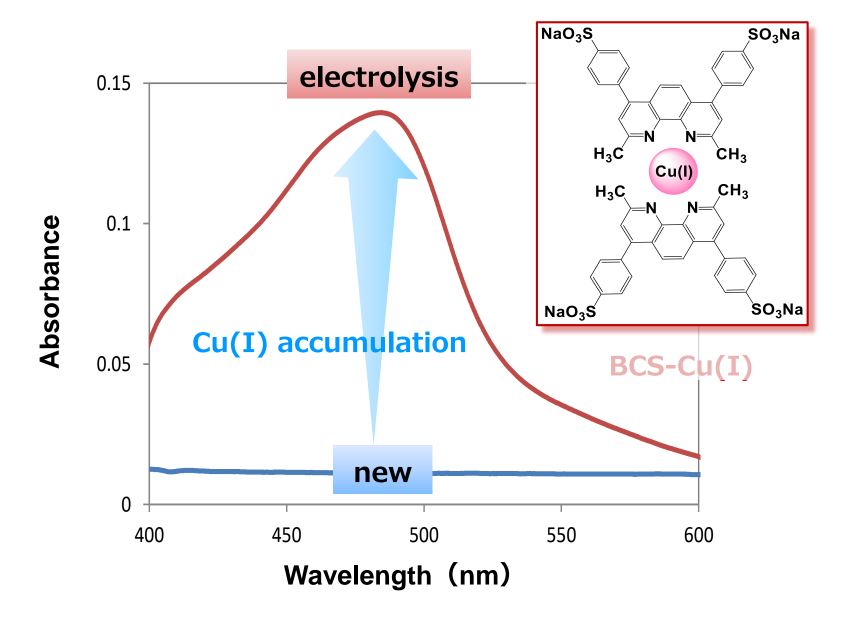
355 356

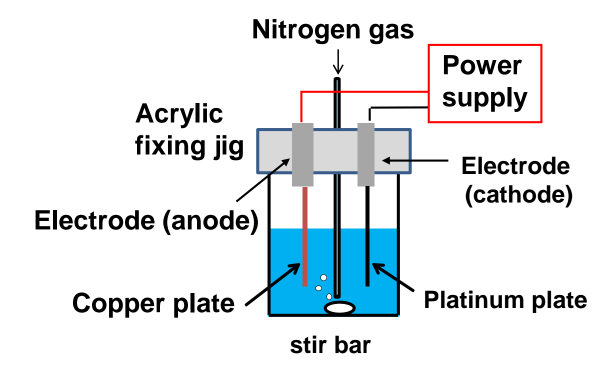
353

REFERENCES:

- 1. Kondo, K., Akolkar, R. N., Barkey, D., Yokoi, M. Copper *Electrodeposition for Nanofabrication* of *Electronics Devices* Chap. 1, Springer, New York (2014).
- 2. Kondo, K., Nakamura, T., Okamoto, N. Correlation between Cu (I)-complexes and filling of via cross section by copper electrodeposition. *Journal of Applied Electrochemistry*. **39**, 1789-1795 (2009).
- 362 3. Healy, J. P., Pletcher, D., Goodenough, M. The Chemistry of the additives in an acid copper electroplating bath, Part II. The instability of 4,5-dithiaoctance-1,8-disulphonic acid in the bath on open circuit. *Journal of Electroanalytical Chemistry*. **338**, 167-177 (1992).
- 4. Frandon, E. E., Walsh, F. C., Campbell, S. A. Effect of thiourea, benzotriazole and 4,5-dithiaoctane-1,8-disulphonic acid on the Kinetics of Copper Deposition from Dilute Acid Sulphate Solution. *Journal of Applied Electrochemistry*. **25**, 574-583 (1995).
- 5. Gabrielli, C., Mocoteguy, P., Perrot, H., Zdunek, A., Sanz, D. N. A Model for Copper Deposition in the Damascene Process Application to the Aging of the Deposition Bath. *Journal of The Electrochemical Society*. **154** (1), D13-D20 (2007).
- 6. Yokoi, M., Konishi, S., Hayashi, T. Adsorption Behavior of Polyoxyethyleneglycole on the Copper Surface in an Acid Copper Sulfate Bath. *Denki Kagaku* **52**, 218-223 (1984).
- 7. Pan, S. Z., Song, L. X., Chen, J., Du, F. Y., Yang, J., Xia J. Noncovalent Interaction of Polyethylene Glycol with Copper Complex of Ethylenediaminetetraacetic Acid and Its Application in Constructing Inorganic Nanomaterials. *Dalton Transactions*. **40**, 10117-10124 (2011).
- 8. Feng, Z. V., Li, X., Gewirth, A. A. Inhibition Due to the Interaction of Polyethylene Glycol, and Copper in Plating Bath: A Surfce-Enhanced Raman Study. *The Journal of Physical Chemistry* B. **107**, 9415-9423 (2003).
- 9. Palmer, J. Determination of Copper Species in Atmospheric Waters. *The Plymouth Student Scientist* **7** (2), 151-184 (2014).
- 381 10. Faizullah, A., Townshend, A. Spectrophotometric Determination of Copper by Flow Injection 382 Analysis with an On-Line Reduction Column. *Analytica Chimica Acta* **172**, 291-296 (1985).
- 383 11. Koga, T., Hirakawa, C., Takeshita, M., Terasaki, N. Quenching Characteristics of 384 Bathocuproinedisulfonic Acid, Disodium Salt in Aqueous Solution and Copper sulfate plating 385 solution. *Japanese Journal of Applied Physics*. **57**, 04FL04-1-5 (2018).
- 12. Noma, H., et al. Analysis of Cu(I) in Copper Sulfate Electroplating Solution. *Journal of The Surface Finishing Society of Japan.* **63**, 124-128 (2012).
- 13. Noma, H., et al. Analysis of Cu(I) Complexes in Copper Sulfate Electroplating Solution by Using Reaction Kinetics with a Chelate Regent. *ECS Transactions*. **58** (17), 77-88 (2014).
- 390 14. Koga, T., Nonaka, K., Sakata, Y., Terasaki, N., Electrochemical Formation and Accumulation of
- 391 Cu(I) in Copper Sulfate Electroplating *Solution. Journal of The Electrochemical Society.* **165** (10), 392 D423-D426 (2018).
- 393 15. Koga, T., Nonaka, K., Sakata, Y., Terasaki, N. Spectroscopic and Electrochemical Analysis of
- 394 Cu(I) in Electroplating Solution and Evaluation of Plated Films. *Journal of The Electrochemical* 395 *Society.* **165** (10), D-467-D471 (2018).

396





Electrolysis parameter

Anode: copper plate

Cathode: platinum plate

Immersion area: $4 \times 2 \text{ cm}^2$

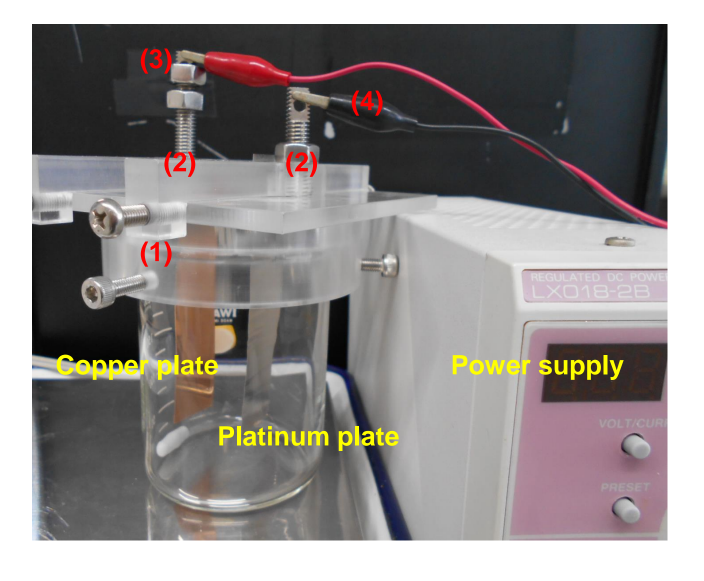
Bubbling gas: nitrogen

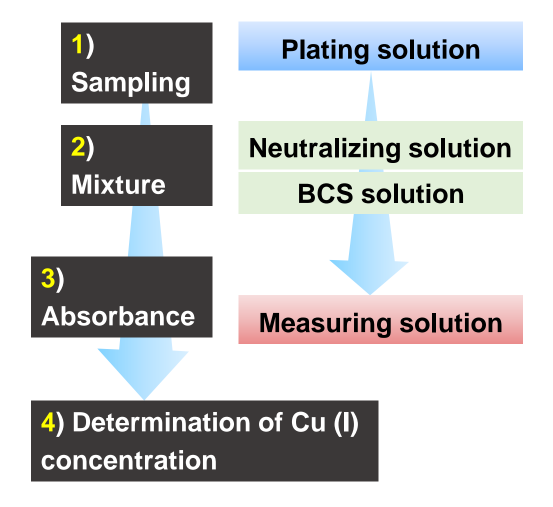
Current value: 0.1 to 1.0 A

Electrolysis time: 0 to 20 min

Experimental bath

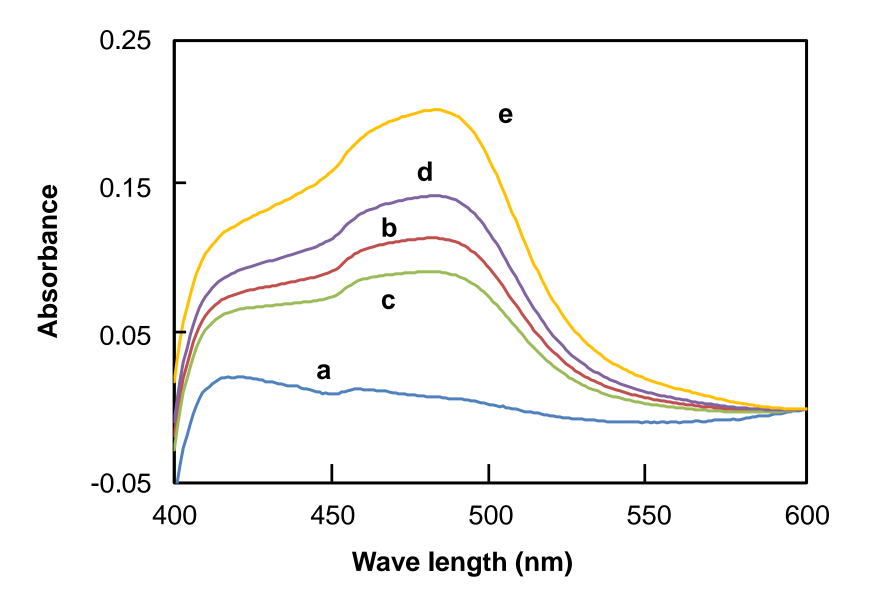
Current density: 1 A = 62.5 mA/cm²

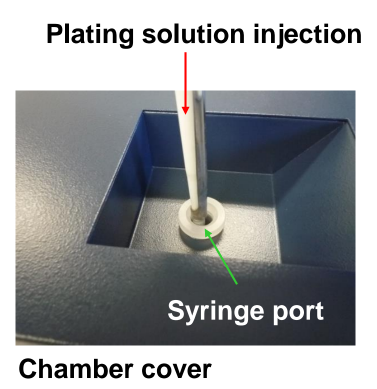


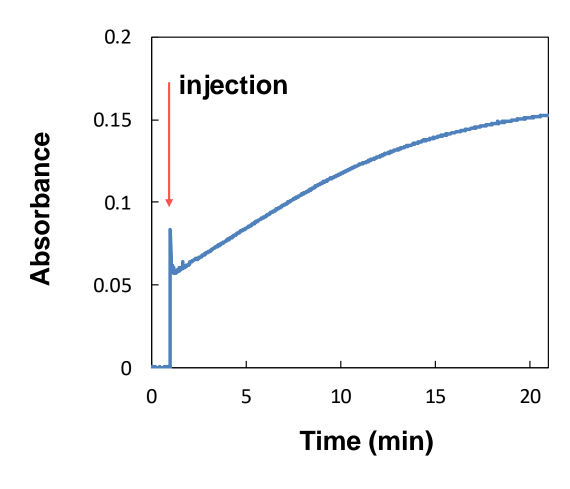


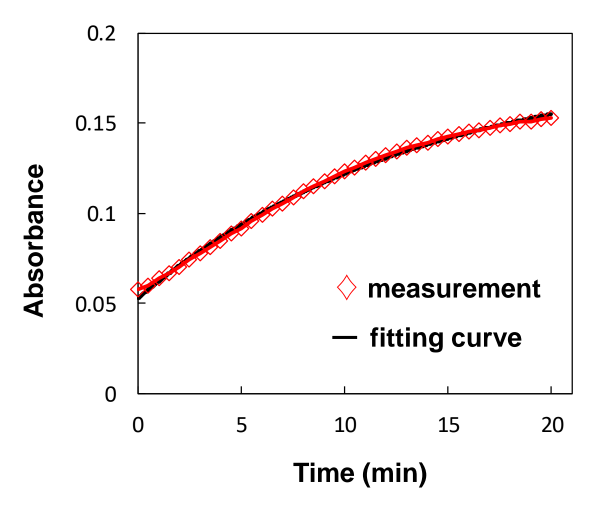


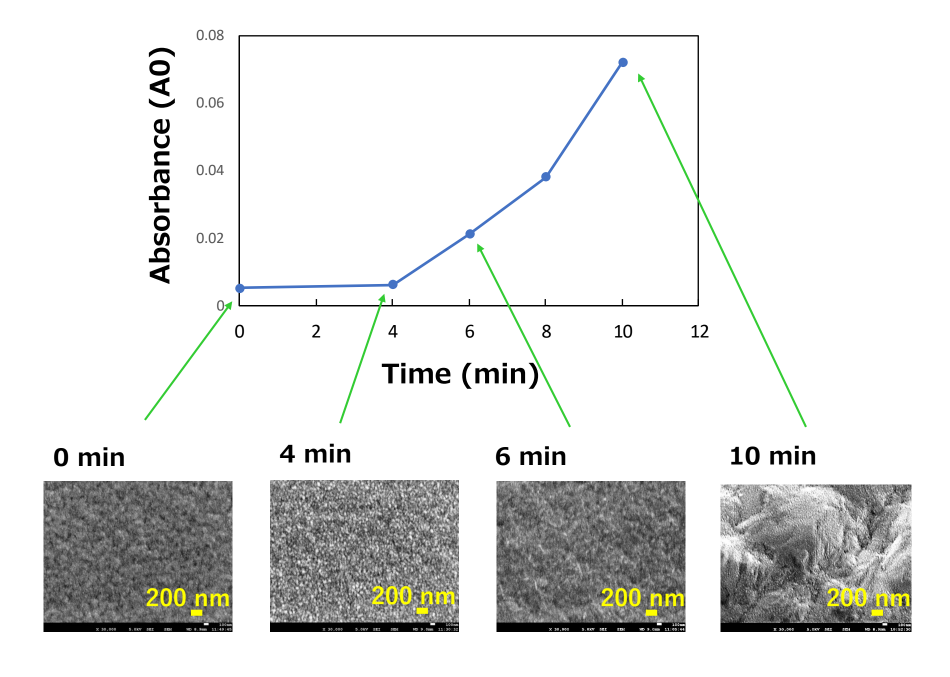
new electrolysis











Name of Material/ Equipment	Company	Catalog Number	Comments/Description
Acetic acid	Wako	016-18835	
BCS	Dojindo	B002	
Copper plate	YAMAMO TO-MS	B-60-P05	
Copper sulfate	Wako	033-04415	
Hydrochorinic acid	SIGMA- ALDRICH	13-1750-5	
JGB	Wako	106-00011	
Magnetic stirrer	Iuchi	HS-30D	
NaOH	NACALAI TESQUTE	31511-05	
PEG4000	Wako	162-09115	
Platinum plate	NILACO	PT-353326	
Power supply	TAKASAG O	LX018-28	
SPS	Wako	327-87481	
Stir bar	AS ONE	1-5409-01	
Sulfuric acid	Wako	192-04696	
Syringe port	JASCO	CSP-749	
Thermostat cell holder with a stirrer	JASCO	STR-773	
UV/vis Spectrophotometer	JASCO	V-630	



ARTICLE AND VIDEO LICENSE AGREEMENT

	Accumulation and Analysis Methods of Cuprous Ion in Copper Sulfate Plating Solution		
Author(s):	Todniaki koga, Yoshitaro Sakata. Nao Terasaki		
	Author elects to have the Materials be made available (as descr com/publish) via: Access Open Access	ibed at	
r -/- 1	ect one of the following items: or is NOT a United States government employee.		
	or is a United States government employee and the Materials were prepare his or her duties as a United States government employee.	ed in the	
· ·	or is a United States government employee but the Materials were NOT prepara his or her duties as a United States government employee.	ed in the	

ARTICLE AND VIDEO LICENSE AGREEMENT

Defined Terms. As used in this Article and Video License Agreement, the following terms shall have the following meanings: "Agreement" means this Article and Video License Agreement; "Article" means the article specified on the last page of this Agreement, including any associated materials such as texts; figures, tables, artwork, abstracts, or summaries contained therein; "Author" means the author who is a signatory to this Agreement; "Collective Work" means a work, such as a periodical issue, anthology or encyclopedia, in which the Materials in their entirety in unmodified form, along with a number of other contributions, constituting separate and independent works in themselves, are assembled into a collective whole; "CRC License" means the Creative Commons Attribution-Non Commercial-No Derivs 3.0 Unported Agreement, the terms and conditions of which can be found at: http://creativecommons.org/licenses/by-nc-

nd/3.0/legalcode; "Derivative Work" means a work based upon the Materials or upon the Materials and other preexisting works, such as a translation, musical arrangement, dramatization, fictionalization, motion picture version, sound recording, art reproduction, abridgment, condensation, or any other form in which the Materials may be recast, transformed, or adapted; "Institution" means the institution, listed on the last page of this Agreement, by which the Author was employed at the time of the creation of the Materials; "JoVE" means MyJove Corporation, a Massachusetts corporation and the publisher of The Journal of Visualized Experiments; "Materials" means the Article and / or the Video; "Parties" means the Author and JoVE; "Video" means any video(s) made by the Author, alone or in conjunction with any other parties, or by JoVE or its affiliates or agents, individually or in collaboration with the Author or any other parties, incorporating all or any portion of the Article, and in which the Author may or may not appear.

- 2. Background. The Author, who is the author of the Article, in order to ensure the dissemination and protection of the Article, desires to have the JoVE publish the Article and create and transmit videos based on the Article. In furtherance of such goals, the Parties desire to memorialize in this Agreement the respective rights of each Party in and to the Article and the Video.
- Grant of Rights in Article. In consideration of JoVE agreeing to publish the Article, the Author hereby grants to JoVE, subject to Sections 4 and 7 below, the exclusive, royalty-free, perpetual (for the full term of copyright in the Article, including any extensions thereto) license (a) to publish, reproduce, distribute, display and store the Article in all forms, formats and media whether now known or hereafter developed (including without limitation in print, digital and electronic form) throughout the world, (b) to translate the Article into other languages, create adaptations, summaries or extracts of the Article or other Derivative Works (including, without limitation, the Video) or Collective Works based on all or any portion of the Article and exercise all of the rights set forth in (a) above in such translations, adaptations, summaries, extracts, Derivative Works or Collective Works and (c) to license others to do any or all of the above. The foregoing rights may be exercised in all media and formats, whether now known or hereafter devised, and include the right to make such modifications as are technically necessary to exercise the rights in other media and formats. If the "Open Access" box has been checked in Item 1 above, JoVE and the Author hereby grant to the public all such rights in the Article as provided in, but subject to all limitations and requirements set forth in, the CRC License.



ARTICLE AND VIDEO LICENSE AGREEMENT

- 4. Retention of Rights in Article. Notwithstanding the exclusive license granted to JoVE in Section 3 above, the Author shall, with respect to the Article, retain the non-exclusive right to use all or part of the Article for the non-commercial purpose of giving lectures, presentations or teaching classes, and to post a copy of the Article on the institution's website or the Author's personal website, in each case provided that a link to the Article on the JoVE website is provided and notice of JoVE's copyright in the Article is included. All non-copyright intellectual property rights in and to the Article, such as patent rights, shall remain with the Author.
- 5. Grant of Rights in Video Standard Access. This Section 5 applies if the "Standard Access" box has been checked in Item 1 above or if no box has been checked in Item 1 above. In consideration of JoVE agreeing to produce, display or otherwise assist with the Video, the Author hereby acknowledges and agrees that, Subject to Section 7 below, JoVE is and shall be the sole and exclusive owner of all rights of any nature, including, without limitation, all copyrights, in and to the Video. To the extent that, by law, the Author is deemed, now or at any time in the future, to have any rights of any nature in or to the Video, the Author hereby disclaims all such rights and transfers all such rights to JoVE.
- Grant of Rights in Video Open Access. This Section 6 applies only if the "Open Access" box has been checked in Item 1 above. In consideration of JoVE agreeing to produce, display or otherwise assist with the Video, the Author hereby grants to JoVE, subject to Section 7 below, the exclusive, royalty-free, perpetual (for the full term of copyright in the Article, including any extensions thereto) license (a) to publish, reproduce, distribute, display and store the Video in all forms, formats and media whether now known or hereafter developed (including without limitation in print, digital and electronic form) throughout the world, (b) to translate the Video into other languages, create adaptations, summaries or extracts of the Video or other Derivative Works or Collective Works based on all or any portion of the Video and exercise all of the rights set forth in (a) above in such translations, adaptations, summaries, extracts, Derivative Works or Collective Works and (c) to license others to do any or all of the above. The foregoing rights may be exercised in all media and formats, whether now known or hereafter devised, and include the right to make such modifications as are technically necessary to exercise the rights in other media and formats. For any Video to which this Section 6 is applicable, JoVE and the Author hereby grant to the public all such rights in the Video as provided in, but subject to all limitations and requirements set forth in, the CRC License.
- 7. Government Employees. If the Author is a United States government employee and the Article was prepared in the course of his or her duties as a United States government employee, as indicated in Item 2 above, and any of the licenses or grants granted by the Author hereunder exceed the scope of the 17 U.S.C. 403, then the rights granted hereunder shall be limited to the maximum

- rights permitted under such statute. In such case, all provisions contained herein that are not in conflict with such statute shall remain in full force and effect, and all provisions contained herein that do so conflict shall be deemed to be amended so as to provide to JoVE the maximum rights permissible within such statute.
- 8. **Protection of the Work.** The Author(s) authorize JoVE to take steps in the Author(s) name and on their behalf if JoVE believes some third party could be infringing or might infringe the copyright of either the Author's Article and/cr Video.
- 9. Likeness, Privacy, Personality. The Author hereby grants JoVE the right to use the Author's name, voice, likeness, picture, photograph, image, biography and performance in any way, commercial or otherwise, in connection with the Materials and the sale, promotion and distribution thereof. The Author hereby waives any and all rights he or she may have, relating to his or her appearance in the Video or otherwise relating to the Materials, under all applicable privacy, likeness, personality or similar laws.
- Author Warranties. The Author represents and warrants that the Article is original, that it has not been published, that the copyright interest is owned by the Author (or, if more than one author is listed at the beginning of this Agreement, by such authors collectively) and has not been assigned, licensed, or otherwise transferred to any other party. The Author represents and warrants that the author(s) listed at the top of this Agreement are the only authors of the Materials. If more than one author is listed at the top of this Agreement and if any such author has not entered into a separate Article and Video License Agreement with JoVE relating to the Materials, the Author represents and warrants that the Author has been authorized by each of the other such authors to execute this Agreement on his or her behalf and to bind him or her with respect to the terms of this Agreement as if each of them had been a party hereto as an Author. The Author warrants that the use, reproduction, distribution, public or private performance or display, and/or modification of all or any portion of the Materials does not and will not violate, infringe and/or misappropriate the patent, trademark, intellectual property or other rights of any third party. The Author represents and warrants that it has and will continue to comply with all government, institutional and other regulations, including, without limitation all institutional, laboratory, hospital, ethical, human and animal treatment, privacy, and all other rules, regulations, laws, procedures or guidelines, applicable to the Materials, and that all research involving human and animal subjects has been approved by the Author's relevant institutional review board.
- 11. JoVE Discretion. If the Author requests the assistance of JoVE in producing the Video in the Author's facility, the Author shall ensure that the presence of JoVE employees, agents or independent contractors is in accordance with the relevant regulations of the Author's institution. If more than one author is listed at the beginning of this Agreement, JoVE may, in its sole



ARTICLE AND VIDEO LICENSE AGREEMENT

discretion, elect not take any action with respect to the Article until such time as it has received complete, executed Article and Video License Agreements from each such author. JoVE reserves the right, in its absolute and sole discretion and without giving any reason therefore, to accept or decline any work submitted to JoVE. JoVE and its employees, agents and independent contractors shall have full, unfettered access to the facilities of the Author or of the Author's institution as necessary to make the Video, whether actually published or not. JoVE has sole discretion as to the method of making and publishing the Materials, including, without limitation, to all decisions regarding editing, lighting, filming, timing of publication, if any, length, quality, content and the like.

Indemnification. The Author agrees to indemnify JoVE and/or its successors and assigns from and against any and all claims, costs, and expenses, including attorney's fees, arising out of any breach of any warranty or other representations contained herein. The Author further agrees to indemnify and hold harmless JoVE from and against any and all claims, costs, and expenses, including attorney's fees, resulting from the breach by the Author of any representation or warranty contained herein or from allegations or instances of violation of intellectual property rights, damage to the Author's or the Author's institution's facilities, fraud, libel, defamation, research, equipment, experiments, property damage, personal injury, violations of institutional, laboratory, hospital, ethical, human and animal treatment, privacy or other rules, regulations, laws, procedures or guidelines, liabilities and other losses or damages related in any way to the submission of work to JoVE, making of videos by JoVE, or publicat on in JoVE or elsewhere by JoVE. The Author shall be responsible for, and shall hold JoVE harmless from, damages caused by lack of sterilization, lack of cleanliness or by contamination due to

CORRESPONDING AUTHOR

the making of a video by JoVE its employees, agents or independent contractors. All sterilization, cleanliness or decontamination procedures shall be solely the responsibility of the Author and shall be undertaken at the Author's expense. All indemnifications provided herein shall include JoVE's attorney's fees and costs related to said losses or damages. Such indemnification and holding harmless shall include such losses or damages incurred by, or in connection with, acts or omissions of JoVE, its employees, agents or independent contractors.

- 13. Fees. To cover the cost incurred for publication, JoVE must receive payment before production and publication the Materials. Payment is due in 21 days of invoice. Should the Materials not be published due to an editorial or production decision, these funds will be returned to the Author. Withdrawal by the Author of any submitted Materials after final peer review approval will result in a US\$1,200 fee to cover pre-production expenses incurred by JoVE. If payment is not received by the completion of filming, production and publication of the Materials will be suspended until payment is received.
- 14. Transfer, Governing Law. This Agreement may be assigned by JoVE and shall inure to the benefits of any of JoVE's successors and assignees. This Agreement shall be governed and construed by the internal laws of the Commonwealth of Massachusetts without giving effect to any conflict of law provision thereunder. This Agreement may be executed in counterparts, each of which shall be deemed an original, but all of which together shall be deemed to me one and the same agreement. A signed copy of this Agreement delivered by facsimile, e-mail or other means of electronic transmission shall be deemed to have the same legal effect as delivery of an original signed copy of this Agreement.

A signed copy of this document must be sent with all new submissions. Only one Agreement is required per submission.

Name:	Tushiaki koga
Department:	Advanced Manufacturing Reserch Institute
Institution:	AICT
Title:	Accumulation and Analysis Methods of Cuprous Ion in Copper Sultate Plating Solution
Signature:	Toshraki koya Date: November 8, 2018

Please submit a signed and clated copy of this license by one of the following three methods:

- 1. Upload an electronic version on the JoVE submission site
- 2. Fax the document to +1.866.381.2236
- 3. Mail the document to JoVE / Attn: JoVE Editorial / 1 Alewife Center #200 / Cambridge, MA 02140

Dear Editor

JoVE59376

"Accumulation and Analysis Methods of Copper Ion in Copper Sulfate Plating Solution" Toshiaki Koga, Yoshitaro Sakata, and Nao Terasaki

We thank Editor for careful reading our manuscript and for giving useful comments. We appreciate that you are interested in our report. We have revised the manuscript JoVE59376 on the base of the editor comments.

We look forward to a publication of our manuscript in Journal of Visualized Experiments.

Sincerely

Toshiaki Koga

Our responses to the reviewew comments are as follows. The correction is described in red in the text.

1. Please take this opportunity to thoroughly proofread the manuscript to ensure that there are no spelling or grammar issues.

Reply:

I reviewed the spelling and grammar.

2. Figure 1: Please include a space between numerical values and their corresponding units (e.g., 1 A, 62.5 mA/cm2).

Reply:

We modified the new Figure 2.

3. Please define all abbreviations before use.

Reply:

We have defined SEM.

4. Please expand the Short Abstract to briefly describe the applications of this protocol.

Reply:

Short abstract was expanded.

Accumulation of cuprous ion in copper sulfate plating solution in model experiment and analysis based on quantitative measurement are described. This experiment reproduces the accumulation process of cuprous ion in the plating bath, which contributes to the elucidation of its mechanism and prediction of the quality of the plating film.

5. Please expand the Short Abstract to briefly describe the applications of this protocol.

Reply:

Introduction was expanded.

Conventionally, it has been claimed that Cu(I) in a plating solution is instantly oxidized to cupric ions (Cu(II)). We have confirmed that there are several millimoles (mmol/L) of Cu(I) in the plating bath of the production line¹². According to this experiment method, the accumulation of Cu(I) similar to the plating bath can be reproduced even in the beaker of the laboratory. This is a fundamental technology to elucidate Cu(I) production and accumulation process in copper sulfate electroplating solution which was unknown¹⁴. Furthermore, by controlling Cu(I) in the plating solution, it is also possible to predict the effect of Cu(I) on the quality of the plating film¹⁵.

6. Please remove commercial language from the manuscript text: JASCO.

Reply:

I excluded JASCO.

7. Please remove commercial language from the manuscript text: JASCO.

Reply:

	ded JASCO.
	ease remove commercial language from the manuscript text: JASCO.
	viewed the protocol.
8. Pl	ease revise the protocol text to avoid the use of any personal pronouns (e.g., "we",
9. Pl	"our" etc.). ease move the discussion about the protocol to the Discussion. Please add more details to your protocol steps.
	viewed the protocol.
	Please provide the concentration of sulfuric acid.
	ed in the concentration of sulfuric acid and copper sulfate.
12 . F	Please specify the temperature that is appropriate for storing the plating solution.
Reply: I enter	red the storage conditions.
13. F	Please specify the current value and predetermined time used in this step.
•	fied the current value and time.
	Please list an approximate volume of BCS solution to prepare.
	CS solution (10-2 mol / L) is prepared by dissolving 0.36 g of the molecule in 100 pure water.

16.	Please describe how to perform curve fitting.
	fitting was removed from the PROTOCOL and inserted into RESULTS.
17.	Please expand to explain the Representative Results in the context of the technique you described.
Reply I expa	: inded REPRESENTATIVE RESULTS and made FIGURE LEGENDS.
18.	Please revise the Discussion to explicitly cover the following.
	ISSION was revised and added in red.
	Please revise the Discussion to explicitly cover the following.
	ISSION was revised and added in red.
	Please include an Acknowledgements section.
Reply Added	: d acknowledgment.
	References: Please do not abbreviate journal titles. Please do not apply superscript atting to reference numbers in this section.
Reply The it	: ems pointed out was revised.
	Table of Materials: Please sort the items in alphabetical order according to the name of ial/equipment.

15. Please describe how to obtain the reaction curve and how to perform a numerical

analysis.

Reply:

They are arranged in alphabetical order.

Dear Reviewer 1

JoVE59376

"Accumulation and Analysis Methods of Copper Ion in Copper Sulfate Plating Solution" Toshiaki Koga, Yoshitaro Sakata, and Nao Terasaki

We thank reviewer for careful reading our manuscript and for giving useful comments. We appreciate that you are interested in our report. We have revised the manuscript JoVE59376 on the base of the reviewer comments.

We look forward to a publication of our manuscript in Journal of Visualized Experiments.

Sincerely

Toshiaki Koga

Our responses to the reviewew comments are as follows. The correction is described in red in the text.

1. In lines 62-63 it is preferred to add a figure or sketch and referred to show the Skeltonstructure of BCS-Cu(I).

Reply:

I changed Figure 3 to Figure 1 and added a guide to Figure 1.

(See Figure 1).

2. in line 88, is it chlorine or chloride and is the source for this material?

Reply:

Yes, add hydrochloric acid so that the chlorine concentration becomes 1.41 mmol/L (1.4).

3. line 136 instead of (minus) it is o preferred to use (negative).

Reply:

Fixed plus to positive, minus to negative.

4. in line 118 and under it is prefer to point out to a figure of that jig for good presentation.

Reply:

A picture of the jig was added as Fig. 3 and an explanation was added in the sentence.

- 2.6. The jig consists of an acrylic beaker fixing part (Fig. 3 (1)) and metal electrode parts (Fig. 3 (2)). The electrode part consists of the parts to fix the plate and the part connect to the cord from the power supply.
- 2.7. Connect the electrode (anode) of the copper plate to the positive of the power supply (Fig. 3 (3)), and the electrode of the platinum plate (cathode) to the negative of the power supply (Fig. 3 (4)).

5. is it true to insert the electrode in the solution before turning on the current? Why?

Reply:

If the plate is inserted with the stirrer rotating, there is a possibility that the plating solution may splash, and the beaker may light up. In order to avoid danger, load the jig beforehand and let the current flow.

6. in line 208, definition of symbols is required.

Reply:

I added the definition in the text.

t is the time from the start of measurement, A0 increases the value of a component that reacts instantaneously (absorbance at t = 0) and AL increases the value of a component that reacts slowly (At - A0). TL is the time constants of AL component.

7. in line 266 what is meant by: a copper plate is attached to the anode and platinum is attached to the cathode?

Reply:

We think that the copper plate of the anode dissolves by electrolysis and promotes the accumulation of copper ions in the plating solution. The same result could be obtained by using a copper plate for the cathode. However, we chose platinum plates, as it is presumed that the plate does not change and the influence on the plating solution is small, and we can use the same plate repeatedly in different experiments.

8. It is also needed to confirm the effect of Cu(I) on the shape of the deposit in accordance with the suggested its formed generation with time> I think it is preferred to add SEM s figure

Reply:

Thank you for your advice. I will use Figure 8.

9. Conclusions are needed

Reply:

I added it to the discussion and described it in the red.

Dear Reviewer 2

JoVE59376

"Accumulation and Analysis Methods of Copper Ion in Copper Sulfate Plating Solution" Toshiaki Koga, Yoshitaro Sakata, and Nao Terasaki

We thank reviewer for careful reading our manuscript and for giving useful comments. We appreciate that you are interested in our report. We have revised the manuscript JoVE59376 on the base of the reviewer comments.

We look forward to a publication of our manuscript in Journal of Visualized Experiments.

Sincerely

Toshiaki Koga

Our responses to the reviewew comments are as follows. The correction is described in red in the text.

1. It must be clearly stated why only BCS was chosen, is it economic?

Reply:

I added it in the red during the introduction.

The cuproine type color reaction reagent, which has been conventionally used for the determination of Cu(I), is hydrophobic and extraction with alcohol is necessary. It was shown that BCS is hydrophilic and can directly measure Cu(I) in aqueous solution.

2. The effect of temperature, pH,operating current/voltage and Cu sulfate concentrations on this solutions must be shown, as these are the industrial plating constrains.

Reply:

The relationship between plating solution conditions and Cu(I) accumulation is interesting. However, we did experiments under the same conditions, so we do not have knowledge of the effects of temperature and sulfuric acid concentration. The relationship with the current value is a major problem and it is described in reference 14. I added a sentence in the discussion.

Although the accumulation amount of Cu (I) tends to increase with the electrolysis time, it is saturated by excessive current (for example, electrolysis for more than 10 minutes at 1 A). The Cu (I) accumulation amount increased with the current value by electrolysis for 10 minutes when the current value was 0.5 to $1.0 \, A^{14}$.

3. The section 2.2. is not clear.

Reply:

The explanation of 2.2 has moved to RESULTS.

4. line no. 109, error on section no.

Reply:

I fixed the mistake.

5. line no. 138, do not agree with 'thoroughly melt BCS'

Reply:

As you pointed out, 10⁻² mol/L BCS can not be completely melted. Correct the sentences and add them.

3.1. The BCS solution (10⁻² mol/L) is prepared by dissolving 0.36 g of the molecule in 100 mL of pure water. Stir the solution and dissolve the BCS in an excess amount relative to the monovalent copper. Store the BCS solution in a dark container and store it in the dark.

NOTE: In the measurement, the BCS concentration in the sample solution is adjusted to 1000 times or more the Cu(I) concentration.

6. line no. 284, please give the value of excessive current.

Reply:

It depends on experimental conditions, but we added one example.

Although the accumulation amount of Cu (I) tends to increase with the electrolysis time, it is saturated by excessive current (for example, electrolysis for more than 10 minutes at 1 A). The Cu (I) accumulation amount increased with the current value by electrolysis for 10 minutes when the current value was 0.5 to $1.0 \, A^{14}$.

7. Why the study is limited to 10 minutes.

Reply:

When an excessive current flows, the correlation between electrolysis time and Cu(I) accumulation amount collapses. For 1.0 A electrolysis, 10 minutes is the limit.

8. It is better to have results and discussion together.

Reply:

Revised REPRESENTIVE RESULTS (insert 2.2), and put the FIGURE LEGENDS term.

Dear Reviewer 3

JoVE59376

"Accumulation and Analysis Methods of Copper Ion in Copper Sulfate Plating Solution" Toshiaki Koga, Yoshitaro Sakata, and Nao Terasaki

We thank reviewer for careful reading our manuscript. We appreciate that you are interested in our report.

We look forward to a publication of our manuscript in Journal of Visualized Experiments.

Sincerely

Toshiaki Koga