Effect of Immersion Time on the Defect Formations and Properties of the Electroplating Chrome

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Abstract - The objective of this work was to investigate the effect of immersion time on the mechanical properties and the defect formation in the electroplating chromium. A polymer was used as substrate, whereas cooper and nickel were used as underlayer materials prior to chrome plating. Three electrolyte solutions bath were prepared to plate copper, nickel, and chrome layers. The electroplating process was conducted using a current of 5 Ampere and a voltage of 12 Volt. The processing variable was carried out by varying the immersion time with 20 minutes, 30 minutes, and 40 minutes. The morphology of the cross-sectional surface of the product was observed using an optical microscope, whereas the mechanical properties were evaluated using a Vickers microhardness tester. The result demonstrates that the microhardness increases as the increase of the immersion time. However, by increasing the immersion time, there was a trend in the formation of fibrous defect found either in the body or interfacial adhesion between Cu/Ni/Cr layers. This defect was attributed to the thickness inhomogeneity of each layer.

Keywords – Plating, Cu/Ni/Cr, time, defects, hardness

I. INTRODUCTION

The rapidly developing technology in automotive industries has resulted in demand to reduce weight and fuel consumption. Polymer is a kind of plastic that has great potential to replace automotive components fabricated by conventional metallic materials [1]. The advantages of plastics are ease of fabrication, flexibility in design, and weight saving. Although a bright surface finish can be obtained by painting or moulding in colour parts, a reliable process is required to produce good decorative surface [2]. Recently, electroplating nickel/chrome was introduced to produce a high-quality and bright reflective surfaces [1]. Electroplating chrome were available in different industries such as aircraft, vehicles, ships, and military sectors. Chrome has some distinct characteristics that are advantages in resisting wear. A thick chrome deposit provides additional wear surface, whereas thinner chrome is widely used for decorative coating. Electroplating chrome on a plastic therefore produces both bright finishes for decorative matter and wear resistance. Since polymer is a non-conductive material, metallization of non-conductive surface is therefore required a method to convert non-conductive plastic surface into conductive by coating with metal seeded surfaces for a subsequent copper deposition [3].

The industry of electroplating plastic, however, experienced many difficulties to produce high-quality products [4]. The major problem in the plating on plastic firstly came from the preparation stage in producing the initial adherent of metal which is electrically conductive material on the plastic surface which is non-conductive material. This step is the key factor in obtaining sufficient durability for the practical application of the chromed plastic. Some studies have reported that the adhesion of metal on a plastic surface has been overcome by pasting carbon prior to copper in the upper layer. Nickel was added in the next upper layer before chromium in the topcoat since it can be responsible for enhancing good corrosion resistance and supporting a lustrous metallic appearance. In a multilayer coating, however, adhesion between layers is of great concern. During plating, the thermal expansion mismatches between the layers may lead to the generation of a tiny crack. When the stresses become greater than the strength of the coating, it may develop a pattern of tiny cracks and the porosities. The crack often form an interlacing pattern to the base metal allowing corrosive liquid or gas to penetrate causing oxidation reaction. The objective of this work was to study the electroplating Cu/Ni/Cr on a polymer substrate. The effect of immersion time on the microstructure and the mechanical properties of the product were observed.

II. EXPERIMENTAL PROCEDURES

The material used as substrate in this work was a polymer that cut into a rectangular box with dimension of 50 mm long, 50 mm wide, and 2 mm thickness. The electroplating was prepared using three different electrolyte solutions that includes the solution for electroplating copper, nickel, and chrome. For electrolyte bath used to plate copper, the solution was composed of copper cyanide 53 g/l, potassium cyanide 103 g/l, and brightener 1.25 cc/l. The bath temperature for electroplating copper was kept in the range of 60-70 °C. The electrolyte solution used for plating nickel included nickel sulphate 300 g/l, nickel chloride 60 g/l, boric acid 50 g/l, brightener magnesium SS 2.5 cc/l, brightener magnesium AM 2.5 cc/l. The electrolyte bath for plating nickel was maintained at temperature of 50-60 °C, whereas the pH of the solution was 4-4.5. The electrolyte solution for plating chrome contained
chromic acid 250 g/l, sulphuric acid 2.5 g/l, and potassium cyanide KC-15 10 g/l. The chemical compositions of the solution were determined by weighing each proportion in the weighing balance and subsequently dissolved in a specific volume of distilled water. The apparatus for the plating includes a plastic box used as the bath to contain the plating solution, and copper rod as the cathode. The anode was attached to the rectifier as the power supply, while copper wire was used to hang the steel sample. The schematic arrangement of the experiment is given in Fig. 1. Prior to electroplating process, the surface of the polymer was pasted using graphite manually. The process of electroplating was conducted varying the immersion time with 20, 30, and 40 minutes. After electroplating, the samples were rinsed in distilled water and dried. The surface of the product was then visually observed. The physical appearance of the surface and thickness of each layer were observed using an optical microscope. The thickness of the layer was also measured experimentally by subtracting the weight of the samples before and after the electroplating. The different weight of the samples before and after electroplating was used to evaluate the average thickness of the layer by dividing the weight by the density of the layer material, and subsequently divided by the area of the samples. The mechanical properties was evaluated using a Vickers microhardness tester with diamond pyramid (136°) using a load of 300 gf and penetration time of 15 second.

Fig. 1 Electrolyte bath for chrome plating

III. RESULTS AND DISCUSSION

A. Surface Morphology

After the samples were covered by graphite, the samples were plated using cooper and subsequently plated by nickel. Three immersion times for the electroplating cooper and nickel were varied by 20, 30, and 40 minutes. Fig. 2 and Fig. 3 show the surface of the samples after covered by cooper layer and nickel with three different immersion time, respectively. The thickness of the cooper layer and nickel layer were given in Fig. 5. The topcoat which is chrome layer was described in Fig. 4. It can be seen in Fig. 5 that an increase of the electroplating time increases the thickness of the layer. Comparing the deposition of cooper and nickel, the deposition rate of cooper is higher than that of the nickel.

B. Microstructure

Fig. 6 shows the microstructure of the multilayer product observed in their cross sectional areas after plated using different immersion times. It was observed in all samples that the thickness of each layer is non-uniform. This is due the different factors affecting the deposition of the metal such as the non-uniformity of the graphite surface which was pasted manually, the current instability, and the condition of electrolite solution. The effect of immersion time on the electroplating cooper, nickel, and chrome as given in Fig. 6 (a-c) revealed that the plating thickness increased with increasing immersion time. This is inline with Faraday’s second law of electrolysis. It was also observed in all samples in Fig. 6 that there was fibrous defect shaped like long cracks or scratches found either in the body of the coating or in the interfacial surface between the layers. An increase in the immersion time seems to reduce the quantity of the defect. Defective parts is not acceptable since it can produce losses and spoil the products image. The defect found in the body and interfacial of the coating has led to an insufficient adhesion and porosities. This is a danger for the material used as corrosion and oxidation resistances since air from outdoor can infiltrate into the metal substrate and may cause the oxidation reaction. When this defect appeared in the surface of the toplayer, it may reduce the attractiveness of the product.

C. Mechanical Properties

Fig. 7 shows the microhardness of copper and chrome layer after plated using electroplating time with 20, 30, and 40 minutes. The measurement was taken three times for each area in order to improve the accuracy of the results. It can be seen that an increase in the immersion time increased the microhardness of the product either in cooper or chrome layer. The microhardness of chrome is higher than that of the cooper. The increase of the chrome hardness as the increase of the immersion time is higher than that of the cooper. The hardnesses of copper for immersion time of 20, 30, and 40 were 110.77 ± 6.57 HV, 130.60 ± 6.03 HV, and 159.03 ± 1.41 HV, respectively. The hardnesses of the chrome for immersion time of 20, 30, and 40 were 184.73 ± 21.18 HV, 236.63 ± 24.17 HV, and 363.70 ± 39.18 HV, respectively.

IV. CONCLUSIONS

This work has reported a study of the electroplating Cu/Ni/Cr on a polymer substrate. Graphite was pasted manually on the polymer surface to convert the non-conductive material into conductive material. Three
electrolyte baths were used to conduct the electroplating copper, nickel, and chrome. The effect of immersion time on the microstructure and mechanical properties were reported. It was observed that the thickness of each layer was non uniform. The fibrous defect was observed to appear either in the body of layer or in the interfacial between the layer.

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REFERENCES


Fig. 2 Copper layer deposited by electroplating time of: (a) 20 minutes, (b) 30 minutes, and (c) 40 minutes

Fig. 3 Nickel layer deposited by electroplating time of: (a) 20 minutes, (b) 30 minutes, and (c) 40 minutes

Fig. 4 Chrome layer deposited by electroplating time of: (a) 20 minutes, (b) 30 minutes, and (c) 40 minutes
Fig. 5 Thickness of copper and nickel underlaying chrome plated using different immersion times
Fig. 6 The optical microstructure of the Cr/Ni/Co layers electroplated for (a) 20 minutes, (b) 30 minutes, and (c) 40 minutes.

Fig. 7 Hardness of copper and chrome layers electroplated for three different immersion times.