

## Original Research Article

# Nickel/CNTs Composite Electroplating

### ABSTRACT

In this paper we present some results on making Ni/CNTs composite by electroplating technique. The work **aimed** creating an advanced nano-composite coating surfaces by taking **advantaged** mechanical properties of the CNTs. The Watts solution was used for nickel plating experiments. We utilized two different methods in **dispersion** CNTs in the Watts solution. In the first method, CNTs were mixed in the Watts solution by ultrasonication. In the second one, the CNTs were chemically functionalized by using diazo coupling reaction in sulfanilic acid ( $\text{NH}_2\text{C}_6\text{H}_4\text{SO}_3\text{H}$ ) and nitrous acid ( $\text{HNO}_2$ ) before mixing in the Watts solution. In order to investigate the role of CNTs additives in coating, nickel was plated onto copper substrates with and without CNTs additives. The Vickers hardness of coatings was measured by Shimadzu Micro Hardness Tester.

*Keywords: Nickel, Carbon Nanotubes, Nanocomposite, Electroplating, Coating*

### 1. INTRODUCTION

Electroplating involves the coating of an electrically conductive object with a layer of metal by using electrical current through electrolyte solution [1]. In order to satisfy wear resistance and intensify hardness of coating, the electroplating technology with micro/nano size materials such as SiC,  $\text{Al}_2\text{O}_3$  and carbon powders were utilized. Many researches showed that the coatings have better mechanical properties by adding the smaller and harder nanoparticles. [2-3]

Carbon nanotubes (CNTs) were known as one of the strongest materials, both in terms of tensile strength and elastic modulus [4-5]. These special properties of the CNTs **are open** a promising way when we use CNTs to make Ni/CNTs composite plating [6-7]. In this paper, we present some results on **fabrication** Nickel/CNTs (Ni/CNTs) nanocomposite by electroplating technique.

### 2. MATERIAL AND METHODS

The CNTs **was** used in this research **production** by CVD method at the Institute of Materials Science. In this method, we used iron nanoparticles formed on  $\text{CaCO}_3$  support as catalyst and substrate that was placed in a thermal CVD system in a gas mixture of acetylene, hydrogen and nitrogen [8].

The electroplating system was built as **the outline** in figure 1. It consisted of a computer connected to an Aligent E3640A power supply by RS-232 connection, and the Watts bath was put on a heating magnetic stirrer. The anode was nickel bar and cathode was copper substrate. In order to control and investigate the influence of plating parameters **in** the electroplating process, we programmed **software for control the** Aligent E3640A by Visual Basic language. The system **can be** worked at some modes such as constant current, constant voltage, pulsed current and pulsed voltage.

41 An important requirement is carbon nanotubes have to be settled before plating to increase  
 42 dispersion ability in the solution and against gathering. We used both physical and chemical  
 43 methods for well dispersion of CNTs in Watts bath. In the physical method we put CNTs into  
 44 the Watt bath, then used ultrasonic vibration for 6 hours in order to get well dispersion of  
 45 CNTs in the Watt bath. In the chemical method, we used diazo coupling reaction by the  
 46 following steps:

- 47 1) Put CNTs into  $\text{HNO}_3$  solution.
- 48 2) Use Ultrasonic vibration for 30 minutes to make up defect on the CNTs.
- 49 3) Use filter paper to filter CNTs from  $\text{HNO}_3$  solution.
- 50 4) Mix the CNTs with aqueous solution of sulfanilic acid ( $\text{NH}_2\text{C}_6\text{H}_4\text{SO}_3\text{H}$ ) at  $70^\circ\text{C}$ .
- 51 5) Drop the solution of sodium nitrite ( $\text{HNO}_2$ ) into the solution in step 4 and keep stirring.
- 52 6) Filter and dry CNTs.

53 We found that after functionalizing by above steps, the CNTs have a better ability to disperse  
 54 in the plating solution. This can be explained as following: after diazo coupling reaction, the  
 55 CNTs remained function groups of  $-\text{N}=\text{N}-\text{C}_6\text{H}_4-\text{SO}_3\text{Na}$ . These function groups will be  
 56 dissociated in the plating solution. The CNTs will be charged the same electrical charge, and  
 57 they will generate Coulomb propulsive force between each other.

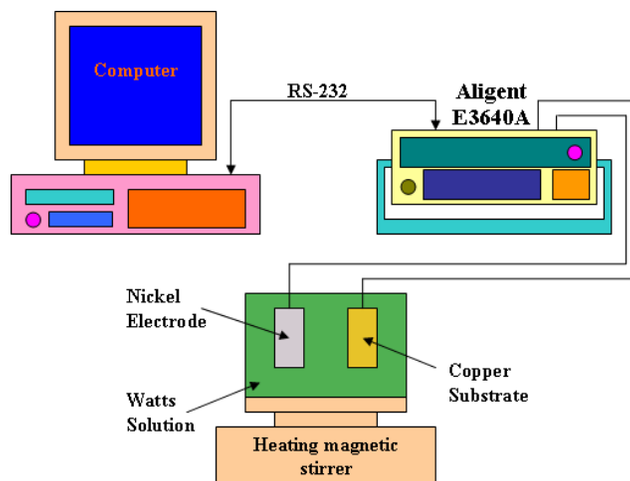
58 We employed Watts solution for nickel electroplating experiments. A standard the solution  
 59 contained: [9]

60 300 g/l Nickel Sulfate,  $\text{NiSO}_4 \cdot 6\text{H}_2\text{O}$

61 50 g/l Nickel Chloride,  $\text{NiCl}_2 \cdot 6\text{H}_2\text{O}$

62 40 g/l Boric Acid,  $\text{H}_3\text{BO}_3$

63 The volume of Watts bath was 1 liter with a pH of 4.5. The temperature of Watts bath was  
 64 kept at  $50^\circ\text{C}$ . We used 10 Hz pulsed current for all plating process in the experiment, therein  
 65 duty cycle for pulsed current is 50%, high current density was  $15\text{A}/\text{dm}^2$  and low current  
 66 density was zero. The Watts bath was stirred at 6.5 rps of rotation speed. The plating  
 67 process with CNTs additives was done using the same conditions above with 5g/l of CNTs-  
 68 concentration in Watt bath.  
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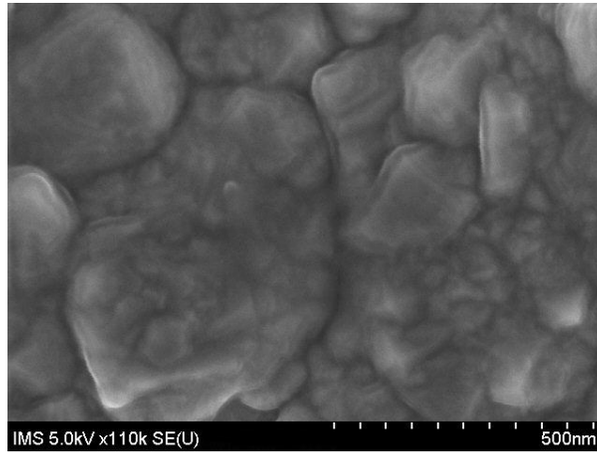


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 71  
 72 **Fig. 1. Schematics of the Ni/CNTs electroplating system**

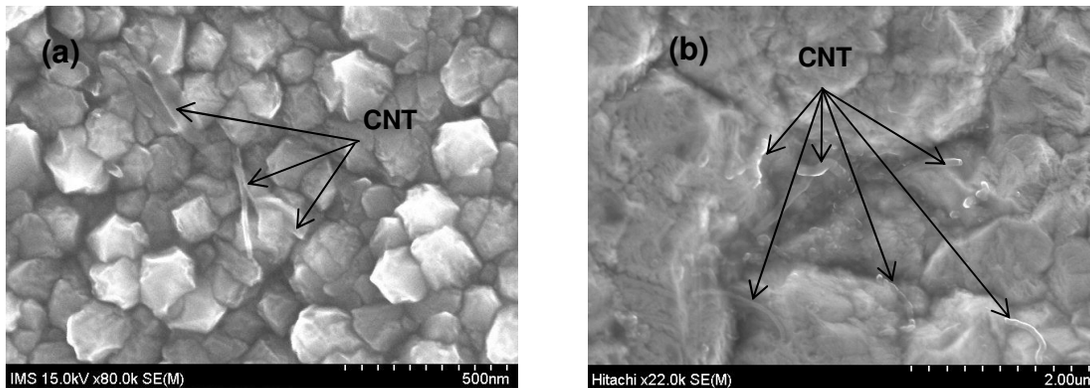
### 73 3. RESULTS AND DISCUSSION

74  
 75 Figure 2 is the SEM image of the normal Ni electroplating surface. It is seen that the size of  
 76 crystal particles of the electroplating layer is relatively large and un-uniform, ranging from  
 77 200 nm to 500 nm. Figure 3 is the SEM image of the Ni/CNTs electroplating surface. The  
 78 SEM image in figure 3a showed that in the case of using non-functionalized CNTs additives,  
 79 the crystal particles of coating is very uniform with size in the range from 100nm to 150nm.

80 Besides that, we observed the presence of the CNTs merging into the Ni coating. The SEM  
 81 image in figure 3b showed that in the case of using functionalized CNTs additive, the crystal  
 82 particles of coating appear unclearly, but there are a lot of CNTs on the plated surface. It  
 83 also showed that the density of functionalized CNTs on coating (figure 3b) was higher CNTs  
 84 density of non-functionalized CNTs on coating (figure 3a).  
 85

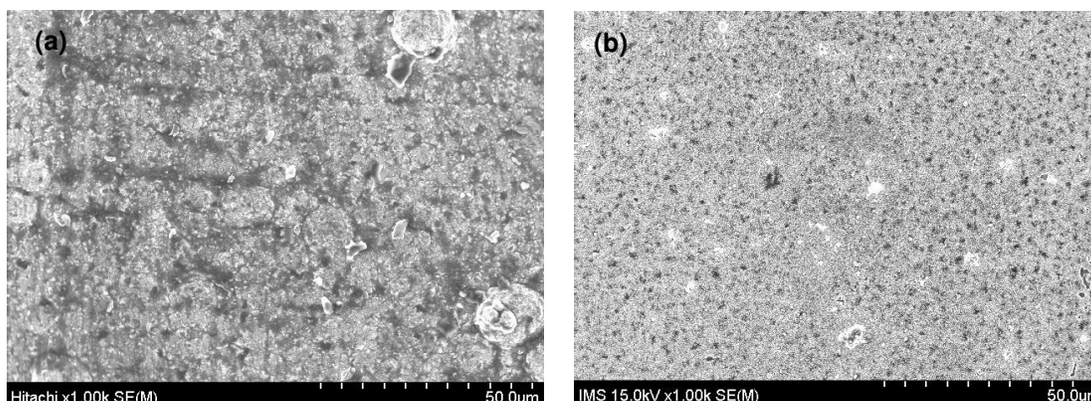


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87  
88 **Fig. 2. The SEM image of normal Nickel coating surface**  
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90  
91  
92 **Fig. 3. The SEM image of non-functionalized Ni/CNTs coating surface (a) and**  
 93 **functionalize Ni/CNTs coating surface (b)**  
94

95 The SEM image of non-functionalized CNTs nickel coating at low magnification was shown  
 96 in figure 4a. It is clear that the surface of coating is not very smooth and there are some  
 97 micro CNTs gathering clusters. This means the dispersion of CNTs in the solution was not  
 98 good enough and the CNTs tend to gather each other to form micro cluster on the Ni plated  
 99 surface. The SEM image in figure 4b showed that the functionalized Ni/CNTs coating is  
 100 much smoother compare to non-functionalized Ni/CNTs coating. This is attributed to the  
 101 better dispersion of the functionalized CNTs in the Watts solution.  
 102



103  
104 **Fig. 4. The SEM image at low magnification of non-functionalize Ni/CNTs coating**  
105 **surface (a) and functionalize Ni/CNTs coating surface (b)**  
106

107 **Table 1. Concentration of elements on copper substrate by EDS analysis**  
108

Element	Weight%
O K	3.12
Cu K	64.32
Zn K	32.56
<b>Totals</b>	100.00

109  
110 **Table 2. Concentration of elements on non-functionalized Ni/CNTs coating by EDS**  
111 **analysis**  
112

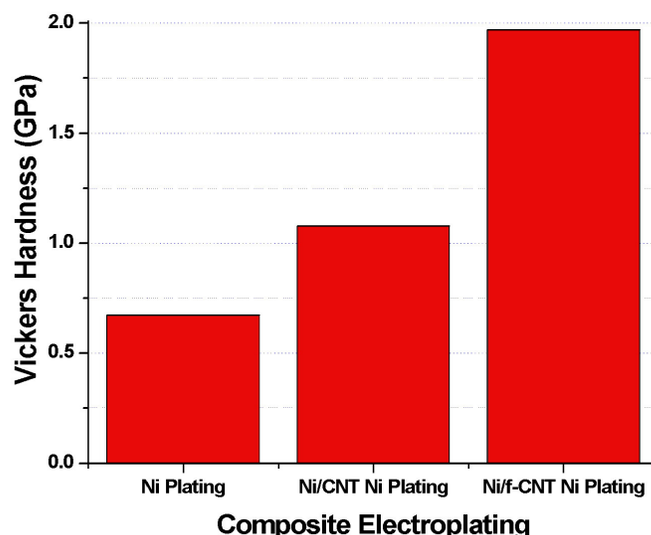
Element	Weight%
C K	11.59
O K	1.71
Ni K	86.70
<b>Totals</b>	100.00

113  
114  
115 **Table 3. Concentration of elements on functionalized Ni/CNTs coating by EDS**  
116 **analysis**  
117

Element	Weight%
C K	12.51
N K	5.77
O K	15.63
Ni K	14.61
Cu K	35.58
Zn K	15.91
<b>Totals</b>	100.00

118  
119 Table 1 is the result of EDS analysis on copper substrate, the result showed that the  
120 substrate was copper alloy with 64.32 %wt Cu, 32.56%wt Zn and 3.12%wt O of composition.  
121 Table 2 is the result of EDS analysis on non-functionalized Ni/CNTs coating. We can see  
122 presence of Ni, C, O, N elements corresponding to the 11.59% wt C, 86.7% wt Ni, and  
123 1.71% wt O of composition. This confirmed the presence of the CNTs on the non-  
124 functionalized Ni/CNTs coating. Table 3 is the result of EDS analysis on functionalized

125 Ni/CNTs coating. Apart from Zn and Cu from copper substrate, we see presence of C, O, N  
 126 elements corresponding to the 12.51% wt C, 5.77%wt N, 15.63%wt O of composition. This  
 127 also confirmed the presence of CNTs on the coating and also confirmed the presence of N  
 128 from -N=N- dimidiate functional groups. From the EDS analysis result in table 2 and table 3,  
 129 the ratio of C weight over Ni weight in the functionalized Ni/CNTs coating and the non-  
 130 non-functionalized Ni/CNTs coating are 0.86, 0.13 respectively. So, it is concluded that the  
 131 functionalized Ni/CNTs coating have more CNTs than that of the non-functionalized Ni/CNTs  
 132 coating. This is **in consistence** with the result **observing** by SEM.  
 133



134  
 135  
 136 **Fig. 5. The Vickers hardness of the electroplating coating**  
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138 **We used Shimadzu Micro Hardness Tester** to measure Vickers hardness of the coatings.  
 139 The results in figure 5 showed that the hardness of the normal Ni coating, non-functionalized  
 140 Ni/CNTs coating and functionalized Ni/CNTs coating were 0.672GPa, 1.08 GPa, 1.97 Gpa,  
 141 respectively. **It concluded** that the non-functionalized Ni/CNTs coating is 1.8 times harder  
 142 than the normal Ni coating, and the functionalized Ni/CNTs coating is nearly 3 times harder  
 143 than the normal Ni coating. This is explained by the **content** of CNTs in functionalized  
 144 Ni/CNTs coating **was higher** than that of normal Ni coating and non-functionalized Ni/CNTs  
 145 coating. The results showed that CNTs **is as great promising of** additive components for  
 146 Ni/CNTs nanocomposite electroplating.  
 147

#### 148 **4. CONCLUSION**

149  
 150 The SEM images and EDS results affirmed the presence of **the** CNTs on the Ni/CNTs  
 151 coating. The functionalized Ni/CNTs coating has higher CNTs density than that of the non-  
 152 non-functionalized Ni/CNTs coating. The functionalized Ni/CNTs coating is much smoother  
 153 compare to the non-functionalized Ni/CNTs coating. The hardness **also enhance** in Ni/CNTs  
 154 coating. The hardness of the non-functionalized Ni/CNTs coating and of the functionalized  
 155 Ni/CNTs coating is 1.8 and 3 times **harder than** the normal Ni coating, respectively. The  
 156 results showed that CNTs **is as great promising of** additive components for Ni/CNTs  
 157 nanocomposite electroplating.

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