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Publication Date 1989-07-01

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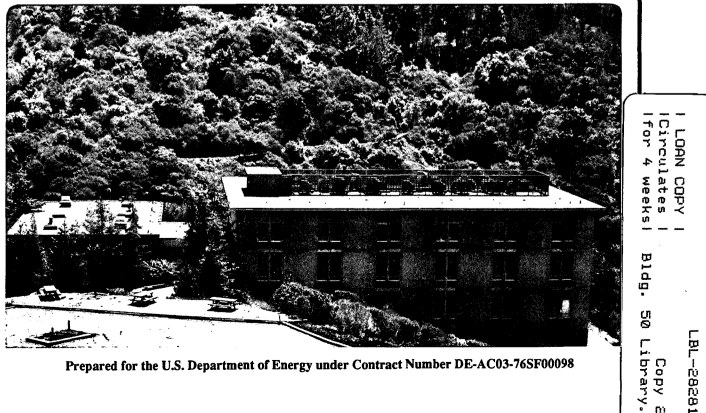
Submitted to the Journal of Materials Science

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July 1989

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Prepared for the U.S. Department of Energy under Contract Number DE-AC03-76SF00098

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Metal Coated Colloidal Particles

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July 1989

This work was supported by the Director, Office of Energy Research, Office of Basic Energy Sciences, Materials Sciences Division, of the U.S. Department of Energy under Contract No. DE-AC03-76SF00098.

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ABSTRACT

Procedures are described for coating of submicron ceramic powders with copper and nickel. The process required precoating of the core particles with a palladium catalyst. A precoating procedure was developed in which palladium chloride is reduced by sodium hypophosphite or stannous chloride on the powder surface in aqueous suspensions. Commercial electroless solutions were used to deposit copper and nickel on the catalyzed powders. The effect of various experimental parameters on the effectiveness of the surface conditioning of the particles has been investigated.

KeyWords: Electroless plating, microencapsulation, colloid, palladium, catalyst.

I. INTRODUCTION

The need for metal-coated fine powders has recently been recognized in the areas of composite materials. In the processing of metal-matrix composites, poor wetting of the ceramic particles with the molten metal often results as a consequence of the incompatibility between covalent or ionic nature of the ceramic surface and the metallic nature of the matrix. Improved wetting characteristics were obtained when the despersoids were initially coated with metals [1-8]. When ceramic powders are incorporated in metal matrices by powder metallurgy, by melt infiltration, or by mixing with molten/semi-molten metal, nonhomogeneous distribution of the ceramic component often result. Uniform distribution becomes increasingly difficult with higher volume fraction of the ceramic particulates. For the fabrication of metalmatrix composites with high ceramic particle content the dispersed phase can be coated with appropriate amounts of the desired matrix metal, and compacted to produce dense body with a relatively homogeneous distribution of the ceramic component.

The electroless depositon of copper or nickel on non-metallic objects usually requires catalysis of the surfaces to be coated. The activation of macroscopic surfaces is often achieved using Pd or Pd-Sn catalysts available as suspension of colloidal particles in the submicron-micron size range [9].

The encapsulation of fine, non-metallic powders with copper or nickel also requires heterogeneous catalysis. Unfortunately, the commercial Pd or

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Pd-Sn cataysts, owing to their particulate nature, are unsuitable to treat submicron core particles: The two kinds of particles can at best heterocoagulate and the effectiveness of the catalyst layer deposited would be greatly influenced by the relative sizes of the core and the catalyst particles.

Current techniques for electroless metal deposition on non-metalic powders usually involve relatively coarse core particles [2-5]. Moreover, the catalysis treatment is multi-step, requiring transfer of the particles between several baths. In addition, little attention has been paid to enhance stability of the suspension during micro-encapsulation. Maintaining dispersion stability becomes particularly difficult with extremely fine particles, especially when even a small weight fraction of the solids, corresponding to a very large particle number concentration, renders the average interparticle distance extremely small.

The typical process for Pd-treatment of particles is carried out in two steps [10]. First, the core particles are sensitized by immersion in an aqueous solution of stannous chloride containing hydrochloric acid. The particles are then isolated and activated by dispersing in an aqueous solution of palladium chloride and hydrochloric acid. The solids are again recovered from the activation bath, and contacted with electroless coating solutions of copper or nickel. The complexities of particle recovery from the various baths between steps dramatically increase with decreasing particle size. Thus, for the catalysis of submicron particles, reduction in the number of steps can mean saving in the corresponding equipment and operating costs.

In this work, a one-step process for the precoating of the core particles with palladium catalyst has been developed in which powder of silicon nitride and silicon carbide of various morphologies, alumina, titania, and a vinylidene copolymer were catalyzed successfully. A process for catalyst pre-coating using stannous chloride as a reducing agent was also developed. Commercial electroless solutions were used to deposit copper on the catalyzed powders. Copper-coated silicon carbide powders were consolidated to fabricate a composite material.

II. EXPERIMENTAL

A. Materials:

Aqueous solutions of palladium chloride [a], stannous chloride [b] and sodiumhypophosphite [c] were used for catalysis of colloidal silicon nitride powder [d, e] and whisker [f]; silicon carbide powder [g], platelets [h,l] and whisker [i]; titania [j], alumina [k] and a vinylidene copolymer [l]. Commercial electroless solutions were used to deposit copper [m] and nickel [n] on the catalyzed core particles. The suspensions were stabilized during microencapsulated by the addition of polymethacrylates of polymethacrylic acid [o].

B. Catalysis of the Non-Metallic Core Powders:

(i) Treatment with Pd Catalyst by Hypophosphite-Reduction:

In this method the silicon nitride particles [e] are catalyzed with palladium by the reduction of palladium chloride by sodium hypophosphite. The untreated solids were suspended in an aqueous solution of palladium chloride and ammonium hydroxide at a pH of 2.5 -3. A solution of sodium hyposphosphite was then added, and the aging was carried out with mild stirring for 1-15 minutes at 25-60°C. The effect of experimental parameters such as pH, concentration of palladium chloride, and concentration of the core particles on the catalytic activity of the treated powders was investigated. The ranges of these parameters were pH 2.5-10; palladium chloride; 40-400 mg/L and core particles 20-140 g/L. The concentration of sodium hypophosphite was typically 0.1-0.4 g/L. The resulting suspension was concentrated by sedimentation of the solids and decanting of the supernatant. This concentrated suspension can be used directly in the subsequent encapsulation step.

The pH of the palladium pre-coating bath appeared to be a critical factor influencing the success of the catalysis process. To determine the effect of this parameter, a two-part investigation was carried out. In the first part, "blank" solutions, free of the core particles, were studied. In the second, the core particles were incorporated in the systems.

The blank systems were prepared at room temperature with 50 cm³ solutions of ≈ 0.1 g/L palladium chloride, to which 0.1 cm³ of 30% hydroxide was added. The pH was adjusted to 2.5-3 by the addition of ammonium hydroxide or hydrochloric acid. To the resulting solution, 5×10^{-2} cm³ of a solution containing 500 g/L NaH₂PO₄.2H₂O was added during rapid magnetic stirring.

a.b.	Aldrich	Chemical	Company.	USA.	
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- c. Mallinckrodt Chemical Company, USA.
- d. Superior Graphite, USA.
- e. Toshiba Ceramics, Japan.
- f. UBE Industries (America), Inc.
- g. Superior Graphite, USA.
- h,i. American Matrix, Inc., USA.
- j. MCB Reagents, USA.
- k. Alcoa, USA.

1. Darvan 820, W. R. Grace & Co., USA.

- m. LC-A and LC-B Solutions, M & T Chemicals, Inc., USA.
- n. Nikland 795, Witco, Allied-Kellite Div., USA.
- o. Daxad Dispersants, W. R. Grace & Co., USA.

The colloidal Pd-Sn catalyst preparation procedure described by Graham [11] was extended to the treatment of submicron silicon carbide powders. The original formulation consisted of:

Palladium chloride $(PdCl_2)$ 1 gHydrochloric acid (HCl, 37%)60 cm³Stannous chloride22 g $(SnCl_2.2H_20)$ 22 gWater to make $\approx 1000 \text{ cm}^3$

In one liter of water containing the hydrochloric acid, the palladium chloride and the stannous chloride were dissolved in order. This solution, when aged at room temperature for about one week, yielded a Pd-Sn catalyst hydrosol.

For the surface treatment of the silicon carbide powders, an aqueous

suspension containing 100 g/L solids was prepared. The pH was adjusted to a value corresponding to the dilute acid prepared by adding 60 cm of 37% HCl to 1000 cm of deionized water. The palladium chloride and stannous chloride, in order, were then dissolved in the suspension. Finally, the dispersion was aged with mild stirring at 50-60°C overnight. In the next step, the catalyzed particles, recovered from the resulting suspension by sedimentation, were washed with 10^{-2} M HCl. In some cases, the solids were also washed with deionized water. Finally, the surface-treated powders were dried at 60°C overnight.

Since this is a two-step process, in contrast to the two-step hypophosphite reduction process, only limited experiments were conducted to investigate the activation of ceramic particles with Pd-Sn catalysts.

C. Microencapsulation with Copper:

To a concentrated suspension containing $\approx 200 \text{ g/L}$ of the catalyzed powders, ammonium polymethacrylate (Daxad-32), and aliquots of deionized water and electroless copper solutions LC-B and LC-A were added in order. The final suspensions contained $\approx 5 \text{ g/L}$ of the Daxad dispersant. The dispersions were stirred magnetically, the "extinction time", defined as the time taken for the disappearance of the blue color from the suspension medium, was measured. This semi-quantitative extinction time was taken as an index of the rate of copper coating---the shorter the duration, the faster the deposition of copper.

D. Microencapsulation with Nickel:

The silicon nitride powders, catalyzed as described in Figure 3, were tested for coating with nickel. To 50 cm of the electroless nickel coating solution, prepared by dilution as recommended by the supplier and pre-heated to 75°C, 0.8 g of the activated core particles were added during magnetic stirring. The aging was carried out until the suspension turned from grey to black in color.

E. Consolidation of the Powders:

The coated silicon carbide particles containing 72 wt% copper, after drying, appeared black due to oxidation of the metal layers. The solids, reduced in Ar-5%H₂ at 250°C for one hour, had the characteristic color of copper. The powders were uniaxially pressed into 9.53 mm (3/8 in) diameter cylindrical pellets at 51 MPa (7500 psi), and cold isostatically pressed at 680 Mpa (100,000 psi) for 3 minutes. The final pellets, annealed in Ar at 500°C, were characterized for density, hardness and microstructure.

III. Results:

A. Catalysis of the Core Powders:

(i) Treatment with Pd Catalyst by Hypophosphite-Reduction:

The blanks with $pH \ge 4$ abruptly became dark within 45 seconds of hypophosphite addition. In about 90 seconds, the system flocculated to yield black particles that settled rapidly. The observed instability of the baths indicated that $pH \ge 4$ may not be suitable for the catalysis of ultrafine ceramic particles. At pH 3, the appearance of dark color was less sudden than at pH 4, and the palladium hydrosol produced was stable towards agglomeration for a few hours. When the initial pH was reduced to 2.5, the color transition was very gradual. Dark color appeared in 2 - 4 minutes, and a relatively stable dispersion of palladium was produced. The observations indicated the choice of pH 2.5 - 3 for the palladium treatment of submicron powders using hypophosphite reduction. At pH 2.5, the reduction process was accelerated by heating the blanks to 45°C without adversely affecting the stability of the resulting hydrosol.

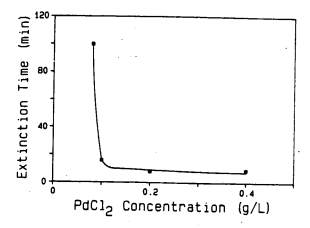


Figure 1. Plot of extinction time for silicon nitride particles catalyzed in 50 cm aqueous suspensions of (initial) pH 3, containing various amounts of palladium chloride, 0.1 cm of 30% ammonium hydroxide, 40 g/L core particles and 0.2 - 0.4 g/L sodium hypophosphite, stirred at 45-50°C for 15 min. The copper coating bath contained 27 g/L of the core particles.

The procedure for the catalysis of silicon nitride core particles at pH \geq 4 was not effective - dispersion of the treated particles in electroless copper solution were chemically stable for 12 hours. When the catalyst precoating was carried out at pH 2.5-3, the core particles could be successfully coated with copper. For a given set of experimental conditions, the extinction time was found to be relatively constant when the particles were catalyzed at pH 2.5-3.

The ammonium hydroxide in the dispersant Daxad 32 neutralized the acid in the low pH core particle catalysis suspensions. No washing of the catalyzed powders was required prior to microencapsulation with copper. Although the dispersant did not significantly affect the stability of the suspensions containing water and electroless solution LC-B, the Daxadcontaining dispersions deflocculated when electroless solution LC-A was added. Substantial agglomeration was observed in systems where the copper deposition was carried out in the absence of Daxad.

Figure 2 is a plot of extinction time as a function of palladium chloride concentration, other experimental parameters being the same. It was observed that the palladium chloride concentration has a marked influence on the success of the catalysis process. The extinction time decreased rapidly, and then became relatively constant as the palladium chloride concentration was gradually increased.

A strong dependence of extinction time on the concentration of core particles in the catalyzing suspension was also observed, as is presented in Figure 3. The extinction time, relatively constant at low particle concentrations, increased sharply and then became relatively constant as the core particle concentration was systematically increased while keeping other factors the same. The extinction time was found to decrease with increasing core particle concentration in the copper coating solutions, as is shown in Figure 4.

Table I includes the data on the extinction times for various combinations of silicon nitride and palladium chloride concentrations, for a fixed $PdCl_2/Si_3N_4$ ratio. It is observed that the extinction is significantly influenced by the actual concentrations of the core particles and palladium chloride, although the relative amounts of the two substances may be the same.

The effectiveness of the catalysis procedure was tested for several types of core particles. The powder treatment was carried out by suspending 2 g of the particles in 50 cm suspensions under the conditions described in Figure 3. The length of the extinction time for the various types of systems, reported in Table II, varies significantly with the nature and geometry of the core particles.

(ii) Treatment with Pd-Sn Catalyst:

The powder treated with the Pd-Sn catalysis process were successfully coated with electroless copper. The catalytic activity of the powders, inferred from the longer extinction time, was very poor when only 60 cm of 37% hydrochloric acid was added per liter of the suspension. A sol of the untreated particles in deionized water was found to be alkaline in nature. The alkaline substance in the powders neutralized a part of the added hydrochloric acid. The catalytic activity was found to be greatly increased when more acid was added to reach a pH value corresponding to acid solution of 60 cm of 37% hydrochloric acid per liter in deionized water. Although systematic investigation about the effect of various experimental parameters such as palladium chloride and stannous chloride concentrations, temperature, and concentration of the core particles, was not carried out. The dramatic effect of pH on the efficiency of the catalysis process was evident.

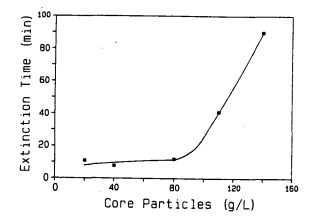


Figure 2. Plot of extinction time for silicon nitride particles catalyzed in 50 cm aqueous suspensions of (initial) pH 3, containing various amounts of the core particles, 0.2 g/L palladium chloride, 0.1 cm 3 of 30% ammonium hydroxide, and 0.4 g/L sodium hypophosphite, stirred at 45-50°C for 15 min. The copper coating bath contained 27 g/L of the core particles.

It was shown earlier [12] that macroscopic surfaces, when catalyzed by immersing in a Pd-Sn hydrosol, should be first washed with dilute hydrochloric acid. If the first rinse was done in water, the surfaces got passivated by a very thin layer of stannous hydroxide formed by the hydrolysis of stannous chloride present in the catalyst bath as a stabilizer. Treatment of the surfaces with acidic fluoride solution removes the passivating layer. Similar observations were made on the catalysis of submicron silicon carbide by the *in-situ* reduction of palladium chloride with stannous chloride. The catalytic activity of the surface treated powders was very poor when the first washing was done using deionized water. The powders were successfully coated with

copper when the solids were first washed using 0.1M hydrochloric acid.

B. Microencapsulation with Nickel:

After addition of the catalyzed silicon nitride core particles to the green electroless nickel solution at 75°C, the suspension turned black in \approx 45 seconds. After one minute, the dispersion was quenched by placing the reaction vessel in cold water and the solids were allowed to settle. The supernatant showed a weaker green color than the fresh electroless nickel solution indicating that some nickel had been removed from the solution.

C. Consolidation of the Powders:

The density of the 72% Cu-28% SiC was 4.98 g/cc. The Vickers hardness of the composite was 120, which is nearly twice the measured hardness of fully dense, annealed 99.9% copper. Scanning electron microscopy revealed a uniform distribution of the silicon carbide particles in the copper matrix (Figure 5).

IV. Discussion:

The extinction time, defined as the time required for the disappearance of blue color from the electroless copper coating solutions, was selected as the experimental index of the overall rate of dispostion of copper. Visual blue color dectection, though somewhat subjective, was chosen because of obvious ease and simplicity of the method. The uncertainty in the measurements is estimated to be ≈ 30 seconds for the extinction time of 5-10 minutes, and 2-5 minutes for the extinction times of 60-90 minutes. Thus, the error in the measurement of extinction time was typically around 10 percent.

Considering the dependence on the concentrations of the palladium chloride (Figure 2) and the core particles (Figure 3), the extinction time might be expected to be relatively invariant at fixed $PdCl_2/Si_3N_4$ ratio, theoretically corresponding to constant surface coverage of the core particles with palladium atoms. However, such a behavior was not observed (Table I), indicating that the nature of the catalyst layer is affected by the actual concentrations of palladium chloride and core particles.

The suspension of the non-catalyzed core particles in electroless copper solutions remained chemically stable for a few days. With the surface treated powders, the extinction time was found to decrease continuously with increasing core particle concentration in the copper coating baths (Figure 4). These observations provided the experimental evidence of the catalytic nature of the surface treated core particles. The curve on Figure 4 is almost linear at low core particle concentrations. This indicates that the kinetics of copper disposition are surface-reaction controlled, since only a small extent of surface area is available for coating. When more surface area is made available by higher concentration of this core particles, the rate of copper deposition should become limited by diffusion of the reactants from the bulk of the solution. This can, indeed, be inferred from the decreasing slope of the curve in Figure 4 at higher core particle concentrations.

A suspension of the untreated silicon nitride solids in electroless nickel solution was stable for two hours at 75°C, while the suspension containing particles treated with palladium catalyst turned black in <1 minute under identical conditions. Thus, the suitability of the catalysis process, using sodium hypophosphite as a reducing agent, was demonstrated for electroless microencapsulation of the fine particles with nickel.

The hypophosphite reduction process is of practical importance for several reasons. This catalytic treatment is essentially single step. The

resulting suspension, after pH adjustment, can be directly used for electroless copper deposition. Thus, the steps of particle washing and transfer between several baths are obviated. For industrial applications, the process may result in savings in the corresponding equipment and operating costs compared with the traditional multi-step processes. However, the catalytic activity of the surface-treated core particles depends on the amount of palladium on their surface. (Figure 2). Thus, a meaningful cost analysis should address optimization of the amount of palladium chloride added per unit weight of the powder catalyzed.

The catalysis process involving stannous chloride was successful only when the reduction was carried out at sufficiently low pH. It was observed that additional acid was required for powders which consumed part of the acid owing to their alkaline nature. Therefore, the measure pH value of the system was concluded to be a more important factor than just the amount of acid to be added as prescribed in the original recipe for the synthesis of Pd-Sn hydrosol [11].

It was established earlier [13] that excess of stannous chloride leads to loss of catalytic activity of the Pd-Sn catalyst hydrosol. Investigations on the effect of stannous chloride concentration on the catalytic activity of the particles treated with the Pd-Sn process may be expected to follow similar trends.

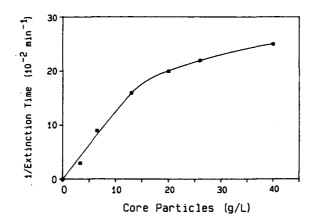


Figure 3. Dependence of extinction time on the silicon nitride concentrattion in electroless solution during micro-encapsulation with copper. The catalysis was carried out by stirring at 45°C for 15 min., 1000 cm³ aqueous suspension at pH 3, containing 40 g of the core particles, 0.2 g of Pd Cl₂, 2 cm³ of 30% NH₄OH and 0.4 sodium hypophosphite.

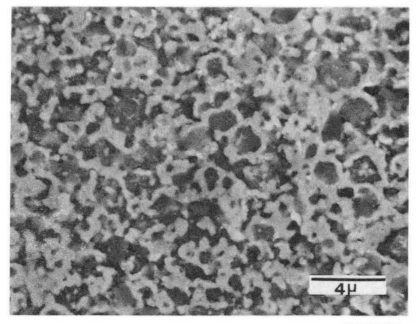
The copper coated colloidal powders were extremely reactive towards atmospheric oxygen. The powders turned black in a few minutes after exposure to air. This affinity of the copper coatings towards oxygen is attributed to the high specific surface area of the ultrafine coated particles.

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The catalysis procedure using sodium hypophosphite as a reducing agent was successfully extended to the catalysis of several shapes and sizes of ceramic powders, as well as for polymeric particles. The results indicate the versatility of the process for the treatment of wide range of core particles of varying morphology. In addition, the catalysis procedure is effective for successful electroless microencapsulation of non-metallic powders with both copper and nickel.

The catalysis experiments also indicate that universal optimum conditions do not exist for all kinds and morphologies of the core particles. In fact, under otherwise identical conditions, the effectiveness of the catalysis process varies significantly with the nature and geometry of the core particles (Table II). Therefore, for the surface treatment of a particular type of core particles, the appropriate content of the core particles, and concentrations of palladium chloride and sodium hypophosphite in the catalysis suspension should be individually determined.

Although the investigation on powders catalysis was limited to the use of sodium hypophosphite and of stannous chloride as reducing agents, substances such as hydrazine [14], dimethylamine borane [14], triethyl amine borane [14], sodium stannate [15], glucose [16, 17], aminosilanes and methanol [18, 19], aminoacids and methanol [20], ascorbic acid [21], and mixtures of amino-sugars, methanol and butyl amine [22, 23], are potential candidates for the reduction of palladium chloride for the catalysis of the non-metallic powders. The use of these reducing agents in suspensions of core particles containing palladium chloride should be investigated to prepare powders of successful electroless micro-encapsulation with copper or nickel.



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Figure 4. Scanning electron micrograph of the 72% Cu-28%SiC composite. The copper-coated SiC powders were uniaxially cold pressed at 51 MPa (7,500 psi) and cold isostatically cold pressed at 680 MPa (1000,000 psi).

V. CONCLUSION:

A one-step process, constituting the reduction of palladium chloride by sodium hypophosphite, was developed for the preparation of aqueous suspensions of activated non-metallic particles which could be directly used for electroless microencapsulation with copper or nickel. For this process, the catalytic activity of the surface treated powders was found to be strongly influenced by several experimental parameters such as the pH, concentrations of the reactants, and the nature and amount of the core particles in suspension. Several kinds of non-metallic core particles of varying morphologies were successfully catalyzed and coated with electroless copper. Consolidation of the copper coated silicon nitride powders yielded a composite having a uniform distribution of the ceramic particles.

VI. Acknowledgements:

This work was supported by Division of Materials Sciences, Office of Basic Energy Sciences, United States Department of Energy, under contract number DE-AC03-76SF00098.

VII. References:

- A. Banerji, P. K. Rohatgi and W. Reif, "The Role of Wetting in the Preparation of Metal-Matrix Composites (a Review)", Metall. 38(7), July 1984, 656-661.
- A. C. D. Chaklader and K. R. Linger, "Ceramic Metal Composites using Metal-Coated Alumina Powder", Composites, Oct 1976, 239-43.
 Hiroshi Tokisue and G. J. Abbaschian, "Friction and Wear Properites of Aluminum-Particulate Graphite Composites", Mater. Sci. Eng. 34, 1978 75-78.
- 4. F. A. Badia and P. K. Rohatgi, "Dispersion of Graphite Particles in Aluminum Castings Through Injection of the Melt", AFS Transactions, 77, 1969, 402-6. 5. Deonath and P. K. Rohatgi, "Cast Aluminum Alloy Composites Containing Copper-Coated Ground Mica Particles", J. Mater. Sci. 16, 1981, 1599-1606.
- K. Gopakumar, T. P. Murali and P. K. Rohatgi, "Metal-Shell Char Particulate Composite using Copper-Coated Particles", J. Mater. Sci. 17, 1982, 1041-48.
- B. C. Pai and P. K. Rohatgi, "Copper Coating on Graphite Particles", Mater. Sci. Eng. 21, 1975, 161-167.
- 8. T. W. Chou, A. Kelly and A. Okura, "Fibre-Reinforced Metal-Matrix Composites", Composites, 16(3), July 1985, 187-206.
- E. Matijevic', A. M. Poskanzer, and P. Zuman, "The Characterization of the Stannous Chloride/Palladium Chloride Catalysts for Electroless Plating", *Plating and Surface Finishing*, Oct 1975, 958-65.
- Hiroshi Kawakami and Shojo Takatsu, Jpn. Kokai Tokkyo Koho JP <u>60</u>, 177, 182, 11 Sept. 1985.
- 11. Peter Graham, Japanese Patent 410824.
- R. L. Cohen, "Mossbauer Studies of Colloidal Catalyst Solutions Used in Metallization of Plastics", Adv. Chem. Ser. <u>194</u>, 1981, 539-51.
- B. J. Page, "Evaluation of Mixed Tin-Palladium Activators for ABS", Plat. Surf. Finish. 74(1), 1987, 56-80.
- S. K. Jalota and K. Mitera, "pH A Necessity, an Evil or a Necessary Evil", Met. Finish. 84(7), 1986, 65-9.
- 15. Michael Gulla, "Dry Catalytic Composition", Ger. Offen. 2,644,111.
- Hitachi Chemical Co., Ltd., "Palladium-Glucose Activation for Copper Electroless Coating", Jpn. Kokai Tokkyo Koho JP 60 36, 670.
- Hitachi Chemical Co., Ltd., "Palladium-Glucose Activation for Copper Electroless Coating", Jpn. Kokai Tokkyo Koho JP 60 36, 674.
- Hitachi Chemical Co., Ltd., "Palladium Aminosilane Activation for Copper Electroless Coating", Jpn. Kokai Tokkyo Koho JP 60 36, 673.
- Hitachi Chemical Co., Ltd., "Palladium Aminosilane Activation for Copper Electroless Coating", Jpn. Kokai Tokkyo Koho JP 60 36, 669.
- Hitachi Chemical Co., Ltd., "Palladium Aminosilane Activation for Copper Electroless Coating", Jpn. Kokai Tokkyo Koho JP 60 36, 672.
- Michael Gulla, "Catalytic Metal of Reduced Particle Size", Eur. Pat. Appl. EP 163, 831.
- Hitachi Chemical Co., Ltd., "Palladium-Amino Sugar Activation for Copper Electroless Coating", Jpn. Kokai Tokkyo JP 60 36, 671.
- 23. Hitachi Chemical Co., Ltd., "Palladium-Amino Sugar Activation for Copper Electroless Coating", Jpn. Kokai Tokkyo Koho JP 60 36, 668.

<u>Table I.</u> Extinction time for $PdCl_2/Si_3N_4$ ratio = $5X10^{-3}$ for particles catalyzed in 50 cm aqueous suspensions of initial pH 3, containing various concentrations of the core particles and palladium chloride, 0.1 cm of 30% ammonium hydroxide, and 0.4 g/L sodium hypophosphite, stirred at 450-50°C for 15 min. The copper coating bath contained 27 g/L of the core particles.

No.	Palladium Chloride	Silicon Nitride	Extinction	
	Concentration	Concentration	Time	
1	0.1 g/L	20 g/L	23 min	
2	0.2 g/L	40 g/L	8 min	

Table II Extinction time for various types of core particles

Source*

Part. Con. No.	Part. Type	In copper bath	Extinction Time
1. Silicon Carbide	d	13 g/L	4 min
2. SiC Whisker	i	13 g/L	90 min
3. Si ₃ N ₄	f	13 g/L	8 min
4. SiC Platelets	h	26 g/L	70 min
5. Titania	j	13 g/L	22 min
6. Alumina	k	13 g/L	3 min
7. Polymer Particles	: 1	6 g/L	7 min

* Materials in Experimental Section

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