Fabrication of palladium/platinum core-shell nanoparticles by electroless metal plating

ABSTRACT

An electroless metal plating method was used to form metallic platinum (Pt) shells on metallic palladium (Pd) nanoparticles. The electroless metal plating method comprised two steps: (1) reduction of Pd ions to fabricate Pd nanoparticles and (2) deposition of Pt nuclei and the following formation of Pt shells by immersing the Pd nanoparticles in a Pt-plating solution. TEM observation revealed that the nanoparticles had a size of 5.7±1.9 nm, were composed of Pd nanoparticles as core and Pt shells. The Pd/Pt core-shell nanoparticles were immobilized electrostatically on carbon support surface-modified with poly(diallyldimethylammonium chloride) (Pd/Pt/C). Cyclic voltammetry revealed that the Pd/Pt/C exerted hydrogen adsorption/desorption, expecting the Pd/Pt/C to function as catalyst for fuel cell.

Keywords: palladium, platinum, core-shell, particles, electroless metal plating, catalyst.

1. INTRODUCTION

Composites of plural materials are of special interest because they exhibit multiple functions derived from their individual components. Among the numerous structures that have been proposed for functional materials to date, core-shell structures, in which a metallic core particle is coated with a metallic shell, have attracted significant attention owing to their unique properties and potential applications in fields such as catalysis, optics, and biotechnology [1-3]. Core-shell structures are also interesting economically because a small amount of expensive material can be used to cover a core of inexpensive material [4].

Pt is an interesting noble metal because Pt nanomaterials and nanomaterials containing Pt exhibit extraordinary chemical and physical properties that can be harnessed for sensors, magnetic materials and catalysts [5-10]. Pt is also used as a catalyst for fuel cells [11-13], and its oxygen reduction reaction (ORR) activity is desired to be improved. According to Zhang et al.'s work, the ORR activity of Pt is dependent on support species, and a metallic palladium support promotes the activity the most efficiently among various metals examined [14]. Their results indicate that composite particles composed of Pd as core and Pt as shell will function as the catalyst with high ORR activity. From this viewpoint, several researchers studied on Pd/Pt core-shell nanoparticles for fuel cell catalysts [15-17]. Many studies produced the Pd/Pt core-shell nanoparticles with reduction of Pt ions to form Pt with reductant in the presence of Pd nanoparticles, in which the formation of Pt may take place not only on Pd nanoparticle surface but also in the solution. This may result in production of not only Pd/Pt core-shell nanoparticles but also Pt nanoparticles that are not in contact with Pd nanoparticles. This unexpected production reduces efficiency of fabrication of Pd/Pt core-shell nanoparticles.

This study aimed to develop a method for fabricating Pd/Pt core-shell nanoparticles. Our previous work performed fabrication of TiO₂/Pt core-shell nanoparticles by electroless metal plating [18]. Because reduction of metal ions to metal takes place at surface of catalyst nuclei deposited on support in the electroless metal plating process, all the generated metal nuclei should be consumed to produce metal layer on the support. The electroless metal plating process was extended to the fabrication of Pd/Pt core-shell nanoparticles. The method is composed of two steps. The first step is fabrication of Pd nanoparticles with reduction of palladium ions in aqueous solution. The Pd...
nanoparticles were used as catalyst nuclei for the following Pt-plating. The second step is Pt-plating, or formation of Pt shells on the Pd nanoparticles surfaces with reduction of platinum ions in the presence of the Pd nanoparticles. The present work also demonstrated immobilization of the Pd/Pt nanoparticles on carbon support (Pd/Pt/C), and electro-chemical measurement using the Pd/Pt/C to confirm that it has a catalytic activity.

2. EXPERIMENTAL

2.1 Chemicals

The starting reagents for the Pd nanoparticles and the Pt shells were PdCl₂ (97.0%) and hydrogen hexachloroplatinate (IV) hexahydrate (H₂PtCl₆·6H₂O) (98.5%), respectively. Ethanol (99.5%) was used as both of a solvent and a reducing reagent for Pd ions in preparation of the Pd nanoparticle colloid solution. A reducing reagent for Pt ions was hydrazine monohydrate (98.0%). Polyvinylpyrrolidone (PVP) (K-30 (Mw=40000)) was used as a dispersing agent in preparation of nanoparticles. Hydrochloric acid (HCl) (35.0-37.0%) and 2-propanol (super special grade) were used for preparing suspension of Pd/Pt/C for fabrication of Pd/Pt/C-immobilized electrode. Poly(diallyldimethylammonium chloride) solution (PDADMAC) (20 wt% in H₂O, Mw=1000,000-2000,000) was used for surface-modification of the carbon support. An electrolyte for measurement of electro-chemical property was perchloric acid (60.0-62.0%, Ultrapur). The Nafion™ dispersion solution (5%, DE520 CS type) and 2-propanol (super special grade) were added to a PDADMAC aqueous solution under sonication, and the mixture was aged at 35°C for 6 h to produce the Pd/Pt/C nanoparticles. The washing process was repeated three times.

The Pd/Pt nanoparticles were immobilized on the carbon powder for the following electro-chemical measurement with the following process. The Pd/Pt nanoparticles were expected to be colloidally stable in water, because their surface was made ionic due to oxide layer produced with passivation. The carbon surface is hydrophobic. Accordingly, the carbon surface should be hydrophilic to increase affinity between the Pd/Pt particle surface and the carbon surface. Thus, prior to the immobilization, the carbon powder was surface-modified with anionic polymer to make the carbon surface hydrophilic with the following process. The carbon particles were added to a PDADMAC aqueous solution under sonication, and the suspension of carbon particles was aged at 35°C for 24-48 h (PDADMAC-C). The initial concentrations of carbon and PDADMAC were 0.27 and 1 g/L, respectively. After the aging, the PDADMAC-C particles were washed with the same process as for the nanoparticles, in which water, however, was used for the re-dispersion. Subsequently, the colloid solution of washed Pd/Pt nanoparticles and water were added to the suspension of washed PDADMAC-C particles, and the mixture was aged at 35°C for 6 h to produce Pd/Pt-immobilized carbon particles (Pd/Pt/C). The initial concentrations of Pd, Pt, and carbon were 0.665 mM, 0.2 mM, and 0.135 g/L, respectively. After the immobilization, the Pd/Pt/C particles were washed with the same process as for the nanoparticles, in which water, however, was used for the re-dispersion. In the final re-dispersion, the concentrations of Pd, Pt, and carbon were adjusted to 1.33 mM, 0.4 mM, and 0.27 g/L, respectively, by decreasing the amount of water used for the re-dispersion.

2.2 Preparation

To obtain catalyst nuclei in electroless metal plating, Pd nanoparticles were fabricated by the reduction of Pd ions with the following process. PdCl₂ and PVP were added to a co-solvent of 60/40 (v/v) H₂O/ethanol under sonication, and the mixture was refluxed at 80°C for 3 h. The initial concentrations of Pd and PVP were 5.3 mM and 7.6 g/L, respectively, in the co-solvent.

The formation of Pd/Pt core-shell particles was achieved using an electroless metal plating method with the following process. Following previous studies on electroless Pt plating [18,19], a Pt plating solution was prepared by dissolving H₂PtCl₆·6H₂O in a 4% HCl aqueous solution to give a Pt concentration of 19 mM. The Pt plating solution and water was added to the Pd nanoparticle colloid solution at 5°C. After 1 h, hydrazine was added to the mixture to initiate Pt deposition (Pd/Pt). The initial concentrations of Pd, PVP, Pt and hydrazine were 2.9 mM, 3.8 g/L, 0.80 mM and 20 mM, respectively. After 24 h, the solvent of the Pd/Pt nanoparticle colloid solution was removed with evaporation under low pressure, and ethanol was added to the residue to re-disperse the nanoparticles in ethanol. The nanoparticles re-dispersed in ethanol were washed with a process composed of centrifugation, removal of the solvent, addition of ethanol, and shaking on a vortex mixer. The washing process was repeated three times.

2.3 Characterization

Transmission electron microscopy (TEM) was used to investigate the morphology of the nanoparticles and performed with a JEOL JEM-2100 microscope operating at 200 kV. To prepare samples for TEM, the colloid solution was dropped on a collodion-coated copper grid, and the
dispersion medium was evaporated in air, and then in a vacuum. The volume-average particle sizes were determined by measuring dozens of particle diameters in the TEM images. The particle powder was characterized by X-ray diffractometry (XRD). For preparing a powder sample for the XRD measurement, supernatant of the particle colloid was removed by decantation, and then the residue of the colloid was dried at room temperature for 24 h in a vacuum. The XRD measurements were performed with a Rigaku Ultima IV X-ray diffractometer at 40 kV and 30 mA using CuKα radiation. The study of the surface composition of particles was performed using X-ray photoelectron spectroscopy (XPS) measurements. The XPS samples were produced by the same process as that for the XRD samples. The XPS spectra were obtained using a JEOL JPS-9010 equipped with a monochromatic Mg Kα radiation source (200 W, 10 kV, 1253.6 eV). To study the composition below the surface, the particles were etched using 500 kV Ar+ sputtering for 1 min. The introduction of PDADMAC on the particle surfaces was qualitatively confirmed using ζ-potential measurements. To measure the ζ-potential of the particles, electrophoretic light scattering (ELS) was performed with a Brookhaven ZetaPlus zeta potential analyzer. Either aqueous HCl or aqueous NaOH was added to the colloid solution to vary the pH for the ELS measurement.

Electro-chemical property of the Pd/Pt/C was measured with a cyclic voltammetry. To fabricate a working electrode, the suspension of Pd/Pt/C nanoparticles, which was prepared by mixing the Pd/Pt/C powder, the 5% Nafion™ dispersion solution, and the 2-propanol, was put onto surface of a glassy carbon rotating electrode (Hokuto Denko, HR2-D1-GC5), and then was dried by applying a hot air. A reference electrode and a counter electrode were a reversible hydrogen electrode kit (ALS, 013373) and a platinum electrode (Hokuto Denko, HX-C5), respectively. The CV was performed with a Hokuto Denko HZ-5000 electrochemical measurement system operating at a scan speed of 50 mV/s in a range of 0.05-1.2 V in 0.1 M perchloric acid in nitrogen gas with no rotating.

3. RESULTS AND DISCUSSION

3.1. Pd nanoparticles

An inset of Fig. 1 (a) shows a photograph of the as-prepared Pd nanoparticle colloid solution. The solution, which was a brownish suspension prior to the reflux, became a black solution during the reflux, implying the reduction of Pd²⁺ to produce metallic Pd nanoparticles. No particle sedimentation was observed in the solution, suggesting that the Pd nanoparticles were colloidally stable. Fig. 1 (a) shows a TEM image of the as-prepared Pd nanoparticles. The particles were quasi-spherical with an average size of 3.4±1.3 nm. Fig. 2 (a) shows the XRD pattern of the nanoparticles. Several diffractions were detected at ca. 40, 46 and 68 degrees. It was hard to determine detailed peak angles for the diffractions, since they were not strong. However, these diffractions could be attributed to the (111), (200) and (220) planes of cubic metallic Pd (ICSD No.: 01-071-3757), respectively, which indicated the production of metallic Pd nanoparticles. This indication supported the implication for production of metallic Pd nanoparticles.

3.2 Pd/Pt nanoparticles

An inset of Fig. 1 (b) shows a photograph of the as-prepared Pd/Pt nanoparticle colloid solution prior to the evaporation of solvent. The solution had a color of black. Since Pt nanoparticle colloid solution is blackish as well as Pd nanoparticles, it was hard to confirm production of metallic Pt by naked eyes. The nanoparticles were also colloidally stable because of no observation of particle sedimentation in the solution. Fig. 1 (b) shows a TEM image of the nanoparticles. The particles were quasi-spherical with an average size of 3.6±1.3 nm. The particle size appeared to slightly increase with the electroless metal plating process, though there was no large increase. The increase in particle size implied formation of Pt shell on Pd nanoparticles.
An inset of Fig. 1 (c) shows a photograph of the as-prepared Pd/Pt nanoparticle colloid solution after the evaporation of solvent. The solution was still black. The nanoparticles were also colloidally stable because of no observation of particle sedimentation in the solution, so that the evaporation process did not influence the colloidal stability of the particles. Fig. 1 (c) shows a TEM image of the nanoparticles. Nanoparticles appeared to contain a core or multiple cores, and the particles had an average size of 5.7±1.9 nm. Pt shells became thick with the evaporation of solvent, which increased the particle size. The evaporation of solvent decreased the volume of solution, which increased various concentrations of species such as Pd particles, generated Pt nuclei and unreacted chemicals.

![Image](image-url)

Figure 2. XRD patterns of (a) Pd nanoparticles and (b) Pd/Pt nanoparticles after the evaporation.
Slika 2. XRD spektri (a) nanočestica Pd i (b) nanočestica Pd / Pt nakon isparavanja.

The increase in the concentrations of unreacted chemicals increased ionic strength of the solution. Increased ionic strength has been reported to compress the double layer of colloidal particles [20-22]. The evaporation of solvent therefore decreased the double-layer repulsion between the Pd nanoparticles and the Pt nuclei. The increased ionic strength was thus primarily responsible for the adsorption of the Pt nuclei on the Pd nanoparticles in the present system. The adsorption was then followed by formation of the Pt shells, or formation of core-shell structure. It should be noted that several nanoparticles appeared to contain multiple cores. The evaporation of solvent also decreased the double-layer repulsion between the Pd nanoparticles. The increased ionic strength was also responsible for the aggregation of Pd nanoparticles in the present system. The aggregates were then coated with Pt shells, which resulted in the production of nanoparticles containing the multiple Pd nanoparticles as the core. Fig. 2 (b) shows the XRD pattern of the nanoparticles. Several diffractions were detected at ca. 40, 46 and 68 degrees as well as the Pd nanoparticles. There were two possibilities in attribution of these diffractions. One was the (111), (200) and (220) planes of cubic metallic Pd, and the other was the (111), (200) and (220) planes of cubic metallic Pt (ICSD No.: 01-071-3756). It was hard to conclude which attribution was reasonable, because there is no large difference in d-spacing between cubic metallic Pd (d-spacing: 0.389 nm) and cubic metallic Pt (d-spacing: 0.392 nm), and the diffractions for the Pd/Pt nanoparticles were also not strong. According to our previous work on TiO2/Pt core-shell particles fabricated with the electroless metal plating method, the crystalline Pt shells formed on the TiO2 particles [18]. Since the electroless metal plating process in the present system was quite similar to that in our previous work, the crystalline Pt shells were considered to be also obtained in the present system, though the XRD did not remarkably provide conclusion on formation of crystalline Pt shells.

![Image](image-url)

Figure 3. XPS spectra of Pd/Pt nanoparticles (a) prior to and (b) after Ar+ etching.
Slika 3. XPS spektri nanočestica Pd / Pt (a) pre i (b) nakon Ar+ nagrizanja.
Fig. 3 shows the XPS patterns of the nanoparticles. The XPS measurement revealed existence of elements of Pd and Pt. Atomic ratios of Pd + Pt for the surfaces of the Pd/Pt nanoparticles were estimated from the XPS peak area intensity. Prior to the Ar⁺ etching, the atomic ratios of Pd and Pt were 73.2 and 26.8 atom%, respectively. These ratios increased to 74.3 atom% and decreased to 25.7 atom%, respectively, after the Ar⁺ etching. This result indicated that the etching of the particle surface revealed the surface of the Pd particles, which further supported the formation of Pt shells on the Pd nanoparticles.

3.3. Surface-modified carbon particles

Figs. 4 (a) and (b) show TEM images of the as-received and the surface-modified carbon particles, respectively. The as-received carbon particles were spherical particles with a size of 44.9±13.9 nm. The surface-modified carbon particles were also spherical particles, and had a size of 41.6±11.0 nm that was almost the same as that of the as-received carbon particles. Accordingly, it was hard to confirm existence of PDADMAC on the carbon particles. Fig. 5 shows the ζ-potentials of various carbon powders as a function of the pH. In the as-received carbon and the surface-modified carbon fabricated for a surface-modification time of 24 h, the ζ-potentials decreased and passed through isoelectric points (IEP) as the pH increased. The IEPs of the as-received carbon and the surface-modified carbon were 4.0 and 9.0, respectively. The IEP for the surface-modified carbon was higher than that for the as-received carbon. Because the PDADMAC is a cationic polymer, this shift to higher pH indicated that the PDADMAC molecules were present on the carbon powder surface, or the carbon powder surface was successfully modified with the PDADMAC. The surface-modified carbon fabricated for a surface-modification time of 48 h did not have an IEP, and had positive ζ-potentials in the range of pH examined. This indicated that an increase in surface-modification time increased the amount of PDADMAC on particles. As a result, the carbon surface became cationic.

3.4. Pd/Pt/C powder

Fig. 4 (c) shows a TEM image of Pd/Pt/C powder. Darker and lighter parts of particles were determined to be Pd/Pt particles and carbon powder, respectively, due to the electron density differences between them. The Pd/Pt nanoparticles were immobilized on the carbon, though several Pd/Pt nanoparticles were not. The ELS measurement revealed that the surface-modified carbon was cationic in the range of pH ca. 3-12, as shown in Fig. 5 (c). According to studies on zeta-potentials of Pt nanoparticles [23,24], the Pt nanoparticles have an isoelectric point of ca. 3. The Pt nanoparticle surface should be anionic at neutral pH. Since the Pd/Pt-immobilization was performed at neutral pH, the Pt surface of Pd/Pt nanoparticles and the surface of surface-modified carbon powder were probably anionic and cationic, respectively. This indicated that electrostatic attractive power acted on them. Consequently, the Pd/Pt nanoparticles were immobilized on the surface-modified carbon powder.

Fig. 6 shows a cyclic voltammetry curve for Pd/Pt/C. A negative peak detected at ca. 0.7 V was ascribed to reduction of Pt oxide. Peaks based on hydrogen adsorption/desorption were detected in a voltage range as low as ca. 0-0.3 V. The result on the detection expected the Pd/Pt/C to function as a catalyst for fuel cell. A charge required to oxidize a monolayer of hydrogen adsorbed on Pt is 210 μC/cm², and an integrated charge amount for hydrogen adsorption in a range of 0.05-0.4 V was 47.5 μC in Fig. 5, which corresponded to a total electrochemical active area of 0.226 cm². An electrochemical active area for Pt loading in catalyst (EAA) (cm²/g-Pt) is required to evaluate a Pt catalyst precisely. Accordingly, further electrochemical characterizations are required to be performed in a...
future work for evaluating catalytic properties of the Pd/Pt/C.

Figure 5. ζ-Potentials of (a) as-received carbon particles, (b) surface-modified carbon particles fabricated with a surface-modification time of 24 h, and (c) surface-modified carbon particles fabricated with a surface-modification time of 48 h

Slika 5. ζ-Potentijali od (a) ugljeničnih čestica kako su primiti, (b) površinski modifikovane čestice ugljenika izrađene sa vremenskom modifikacijom od 24 h, i (c) površinski modifikovane čestice ugljenika proizvedene sa vremenskom modifikacijom od 48 h

Figure 6. Cyclic voltammogram of Pd/Pt/C particles. The Pd/Pt/C particles used for the measurement was the sample (c) in Fig. 4

Slika 6. Ciklični voltamogram čestica Pd / Pt / C. Čestice Pd / Pt / C koje su korištene za merenje bile su uzorak (c) na slici 4

4. CONCLUSIONS

The Pd nanoparticles with the size of 3.4±1.3 nm were produced by the reduction of Pd ions, and then the Pt shells were fabricated on the surfaces of the Pd nanoparticles by using the electroless metal plating technique. The size of Pd/Pt core-shell nanoparticles after the evaporation treatment was 5.7±1.9 nm. The surfaces of carbon support powder were modified with the PDADMAC with the cationic group, and then the Pd/Pt core-shell nanoparticles with anionic surfaces were immobilized electrostatically on the surface-modified carbon support. The hydrogen adsorption/desorption took place for the Pd/Pt/C powder. A study on application of the Pd/Pt/C particles to catalyst for fuel cells is being prepared for their practical use.

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5. REFERENCES


IZVOD

IZRADA PALADIJUM / PLATINSKOG JEZGRA NANOČESTICA ELEKTROSTATSKIM METALNIM PREMAZOM

Metoda pločica bez elektrolita je korišćena da se formiraju metalne čestice školjke platine (Pt) na nanočesticama metalnog paladijuma (Pd). Metoda neplastičnog metala je obuhvatila dva koraka: (1) smanjenje jona Pd za proizvodnju Pd nanočestica i (2) deponovanje jezgra Pt i sledeće formiranje Pt školjki potapanjem Pd nanočestica u Pt- rastvor. TEM posmatranje je otkrilo da je nanočestice Pd kao jezgra i Pt školjki. Nanočestice Pt / jezgra su bili elektrostatici imobilizovani na površini ugljenika, modifikovanim sa poli (diallildimethilammonium chloride) (Pd / Pt / C). Ciklička voltametrija pokazala je da je Pt / C imala adsorpciju vodonika / desorpciju, očekujući da Pd / Pt / C funkcionira kao katalizator za gorivne čelije.

Ključne reči: paladijum, platina, jezgro -čestice, elektrostatska metalna galvanizacija, katalizator.

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