Oxygen reduction on bimodal nanoporous palladium-copper catalyst synthesized using sacrificial nanoporous copper

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# Abstract

Nanoporous copper (NP-Cu), as a sacrificial support, was used for the synthesis of bimodal nanoporous palladium-copper (BNP-PdCu) for oxygen reduction reaction (ORR) electrodes in fuel cells. The catalytic performance of BNP-PdCu in ORR per electrochemical surface area was enhanced by the dissolution and removal of supporting NP-Cu, which indicates that the intrinsic catalytic properties of palladium are improved by the proposed synthesis strategy including galvanic replacement of copper with palladium, following copper dissolution. Cu remained on Pd surfaces even after dissolution of Cu. Additionally, significant local lattice contraction was observed at the ligament surface. First-principles calculations on the adsorbing oxygen species on Pd show that both lattice contraction and alloying with copper increase the binding energies of oxygen species to the Pd surface. The high ORR activity of the present BNP-PdCu is suggested to be mainly due to the Cu-ligand effect.

Keywords: Pd; nanostructure; surface reaction

# Introduction

Catalytic electrodes in fuel cells are typically made from platinum, which is rare in the earth’s crust. The scarcity is the main reason for the high and fluctuating price of platinum and prevents the wide application of fuel cells. The performance of fuel cells is often governed by the oxygen-reduction reaction (ORR) at an air electrode.[1] Thus, various studies on the optimization of catalytic platinum electrodes have been conducted.[1–3]

Another effort is the substitution of platinum by other elements. Palladium is one possible element for the substitution because the electronic structure of palladium is similar to that of platinum, which is clearly understood from the periodic table. From the viewpoint of surface area, nanostructuring of palladium is a key strategy for the efficient production of the air electrode. However, it has been found that the catalytic properties of palladium for ORR are much lower than those of platinum.[4] For this reason, many studies have been carried out to achieve a better catalytic performance of palladium than platinum by efforts such as nanoparticle formation[5] and alloying.[6]

Nanoporous metal synthesis, by dealloying or selective dissolution of one element from a binary alloy, has been developed after a report in 2001[7] and is now another technology for nanostructuring. Dealloying can create three-dimensional nanoporous structures in various metals, which generate pores and ligaments in the nanometer-size range.[8,9] It is well known that nanoporous Raney copper can be synthesized by dealloying Cu–Al alloys in alkaline solutions.[10] Nanoporous copper (NP-Cu) can also be produced by dealloying ductile Cu–Mn alloys.[11–13] Manganese is easy to dissolve even in neutral solution;[14] thus the precursor Cu–Mn alloys may be favorable for efficient production of NP-Cu.

Noble metals, including palladium, can be deposited in the form of a dense film or fine particles on copper by electroless plating or galvanic replacement.[15–17] Thus, nanoporous palladium with a skeletal structure replicated from NP-Cu can be produced by electroless plating of palladium on NP-Cu. The supporting NP-Cu can be removed readily by HNO3 which dissolves copper selectively, and leaves bimodal nanoporous platinum with residual copper (BNP-PdCu) (Fig. 1). In addition, the distribution of lattice spacing from surface to core is significantly varied in nanostructured materials including nanoporous metals,[18,19] causing peculiar surface chemistry.[18,20–22] In this study, BNP-PdCu for ORR catalysts was fabricated by electroless plating of palladium on the sacrificial NP-Cu and subsequent dissolution of NP-Cu. We focus on high ORR activity per surface area of fabricated BNP-PdCu catalysts, using first-principles calculations on the adsorption of oxygen species on palladium surfaces.

# Results

STEM images (Fig. 2) indicated the bimodal structure of the BNP-PdCu after dissolution of copper. The nanoporous hollow structure with an average pore diameter of 23 nm (Fig. 2(a)), which is replicated from NP-Cu (Fig. 1(b)), was composed of finer Pd aggregates with 5–10-nm diameters (Fig. 2(b)), forming the bimodal nanoporous structure. STEM-EDXS revealed that Cu remained even after dissolution by HNO3, although Cu and Pd exhibited a homogeneous distribution without surface segregation (Fig. 2(c)). HR-TEM observation (Fig. 2(d)) showed significant local lattice contraction at ligament surfaces. The remaining Cu in the sample after dissolution by HNO3 was confirmed by EDXS (Fig. 3(a)), where the atomic ratio of Cu/Pd was 0.25 before dissolution and 0.15 after dissolution. On the other hand, manganese was not detected by EDXS and ICP-AES in BNP-PdCu before and after the dissolution of copper. Thus, the contribution of manganese was eliminated in the following discussion on the catalytic properties of the BNP-PdCu samples.

XRD patterns of BNP-PdCu before and after copper dissolution are shown in Fig. 3(b). Before copper dissolution, the broad peaks for palladium were detected with minor peaks for metallic copper. The absence of copper peaks in the XRD pattern even before dissolution of copper suggested that a considerable amount of copper was removed at the stage of the displacement deposition of palladium. Comparison of the present Cu/Pd composition ratio with the case of Cu/Pt[23] also suggested the removal of copper during the displacement deposition of palladium. The broad peaks for palladium were detected whereas no copper was detected by XRD after dissolution by HNO3. Pd remained undissolved after immersion in HNO3 for removal of skeletal Cu. An inspection of the XRD patterns showed no significant peak shift of Pd (Table I), compared with the case of Cu/Pt.[23]

CV curves for BNP-PdCu before and after copper dissolution (Fig. 4(a)) were typical for palladium[24–26] with peaks for oxygen desorption. Copper dissolution by HNO3 caused no significant change in the redox process during CV scanning. On the other hand, no redox peaks from copper were detected. Considering that elemental mapping in TEM for BNP-PdCu showed no clear segregation of palladium and copper (Fig. 2(c)), one can suggest that the change in the chemical state of surface Cu by alloying with Pd and/or that the surface coverage by Cu is not so high.

The ORR activity of the BNP-PdCu samples was examined by RDE (Fig. 4(b)). In the polarization curves, the current density at a high potential of 1.1 V was almost independent of the electrode rotation rate, which corresponded to the kinetic region of the ORR. As the potential was decreased from 1.1 V, transitional regions from a kinetic- to a diffusion-controlled reaction rate were observed, with a lower current density at a higher electrode rotation rate. Below 0.4 V, the current density was almost constant and a diffusion-controlled reaction occurred.

The specific activities of BNP-PdCu before and after dissolution of copper were 0.32 and 0.61 mA cm−2, respectively, according to the RDE results. These values were significantly higher than those of the conventional Pd/C catalyst and comparable with the values reported recently for other nanostructured Pd-based alloys.[27–31] On the other hand, the mass activity of BNP-PdCu before and after copper dissolution was 0.023 and 0.034 A mgPd−1, respectively. The copper dissolution increased the specific activity of the BNP-PdCu. A higher activity per ECSA (not per mass or geometric area) for BNP-PdCu after copper dissolution meant that the copper dissolution enhanced the intrinsic ORR catalytic nature of the palladium.

# Discussion

In the case of platinum, which has been examined intensively in previous studies, lattice straining affects the overall catalytic performance for ORR in platinum;[32–34] lattice contraction is favorable for efficient ORR catalysis on platinum.[32] However, it has been reported that both lattice contraction[35–37] and expansion[27] in palladium catalysts enhances the overall catalytic performance for ORR. XRD analyses revealed that the apparent lattice constants of palladium in BNP-PdCu are close to those for bulk palladium (Table I), which is in contrast with the case of BNP-Pt[23]. However, HR-TEM of BNP-PdCu showed significant local lattice contraction at ligament surfaces, as exemplified in Fig. 2(d). That is, the lattice spacing at the surface of ligaments is smaller than that in the bulk part of ligaments. This lattice disorder at the surface of nanoligaments is often observed in nanoporous metals and is attributed to the complex three-dimensional nanostructure with positive and negative curvatures.[18,19,22] The present BNP-PdCu is replicated from NP-Cu; therefore, the lattice disorder is likely to be prominent, considering the increased complexity in the replicated BNP structure.

It is known that the binding of oxygen species to Pd surfaces is too strong and it is difficult to cause further reduction of the oxygen species to form water on Pd surfaces.[46,47] Therefore, a weak oxygen binding is desirable to balance the rates of electron transfer/O−O bond breaking and removal of O species from Pd surfaces.[27] As shown above, Cu remained on the BNP-PdCu surface even after dissolution by HNO3, in addition, a lattice disorder was generated at its surface. Therefore, it is suggested that the Cu-ligand effect and/or the strain effect led to the high ORR activity of BNP-PdCu. To obtain insights in the origins of the high ORR activity of BNP-PdCu, the binding energies of O and O2 to Pd and PdCu were calculated. The results are shown for Pd in Fig. 5 and for PdCu in Fig. 6. The most stable binding sites for Pd were the fcc for O and the t-f-b for O2, regardless of the strain. However, the most stable binding sites for PdCu were different from those for Pd. The most stable binding sites and binding energies are listed in Table II. Table II shows that the binding energies of the oxygen species are increased (negatively decreased) by the contraction strain for Pd, while the binding energies are decreased (negatively increased) by the expansion strain, which indicates that the contraction strain enhances the desorption of oxygen species on Pd although the expansion strain disturbs the desorption. Thus, the contraction strain leads to the enhanced ORR activity of Pd. In addition, Table II shows that the binding energies are increased by the addition of Cu, regardless of the strain. This suggests that the ORR activity of Pd is enhanced owing to the Cu-ligand effect. The Cu-ligand effect can significantly enhance the ORR activity of Pd because the binding energies are largely increased by the addition of Cu.

The densities of states (DOS) for a palladium atom of the Pd (PdCu) surface binding to an oxygen atom are shown in Fig. S1 in Supplementary Material. The electronic states for Pd were hardly changed by the presence of strains, which corresponds to the result that the strain had little effect on the d-band center for Pd (Table II). However, the electronic states were changed by the presence of Cu (see Fig. S1(a) and (d)). Both the downshift and upshift of the d-band center were caused by the addition of Cu (Table II). Thus, the variation in binding energy by the addition of Cu cannot be explained only from the viewpoint of the d-band center. The charge density distribution between the palladium and oxygen atoms is shown in Fig. 7. The charge density between the palladium and oxygen atoms was lower for PdCu than for Pd. Therefore, it is proposed that the binding energy increased by the addition of Cu is related to the reduced charge density.

As shown above, the contraction strain and the addition of Cu enhance the ORR activity of Pd. Hence, the high ORR activity of BNP-PdCu may be related to both the strain effect and the Cu-ligand effect. The present experimental results showed that the ORR activity was enhanced by the dissolution of Cu, although the contraction strain might be reduced and the amount of Cu on Pd was reduced by the dissolution. The Pd surface that is covered by Cu before the dissolution cannot provide the sites for ORR. In such a case, only the strain effect enhances the ORR activity because the Pd surface that is not covered by Cu provides the sites for ORR. After the dissolution, however, not only the strain effect but also the Cu-ligand effect enhances the ORR activity because Cu remains on the Pd surface. It is therefore suggested that the high ORR activity of BNP-Pd mainly arises from the Cu-ligand effect.

# Conclusions

Efficient ORR BNP-PdCu catalysts were fabricated by using NP-Cu as a sacrificial metallic support. Cu remained on Pd surfaces even after dissolution of Cu. The high ORR activity of BNP-PdCu was suggested to mainly arise from the Cu-ligand effect.

# Experimental methods

Copper-manganese alloy ingots with the composition of Cu0.3Mn0.7 were fabricated by the arc melting of pure copper and manganese (Nilaco, Tokyo, Japan) in an Ar atmosphere. After heating at 1123 K for 24 h in an Ar atmosphere, the button-shaped alloy ingot was water-quenched,[40] followed by cold rolling to 0.2-mm thickness.

Dealloying to fabricate NP-Cu was conducted by free corrosion of the rolled Cu0.3Mn0.7 sheet in 1 mol L−1 HCl for 24 h, where manganese was dissolved selectively and copper formed a nanoporous structure spontaneously (Fig. 1(b)).[11–13] After recovery by percolation and washing in pure water, the NP-Cu was pulverized manually in an agate bowl.

NP-Cu (0.05 g) was mixed with 5 wt% Nafion ethanolic solution (1 g) and of water (5 g). After homogenization in an ultrasonic bath, the suspension was fixed on a glassy carbon (GC) rotation disk electrode (RDE) with 5-mm diameter. The displacement deposition of palladium on the fixed NP-Cu was conducted by immersing the GC RDE in diluted HCl containing 0.1 mol L−1 PdCl2 at 273 K for 1 h at 600 rpm.[41] Thus, the GC electrode included NP-Cu with a fine palladium deposition; the bimodal nanoporous structure of palladium supported on NP-Cu was expected to form already at this step (Fig. 1(a)). The GC electrode that included BNP-PdCu was immersed in 0.1 mol L−1 HNO3 for 18 h to dissolve the supporting copper.

We conducted cyclic voltammetry (CV) at room temperature by using the GC electrode loaded with BNP-PdCu as a working electrode. To evaluate the electrochemical surface area (ECSA) of the BNP-PdCu samples, CV was conducted in degassed 0.1 mol L−1 H2SO4 with a scan rate of 10 mV s−1. A reversible hydrogen electrode (RHE) and Pt wire were used as a reference and counter electrode, respectively. For electrochemical cleaning, CV was repeated until a steady current-potential cyclic curve was attained. The charge () associated with the reduction of oxidized monolayer was calculated by using the cathodic peak between 0.6 and 0.8 V (vs. RHE), which was then converted to the ECSA () of the sample:[42]  
where the subscripts “sample” and “bulk” represent the BNP-PdCu and bulk palladium, respectively.

The working GC-RDE-loaded BNP-PdCu was rotated for ORR activity measurements during CV in an electrolyte of O2-saturated 0.1 mol L−1 H2SO4 with a scan rate of 10 mV s−1. The kinetically-controlled current , which is an index of the ORR activity of the BNP-PdCu, was calculated according to:

where is the measured current; is a constant and is the RDE rotation speed. was derived from the intersection at by the Koutechy–Levich plot of (at 0.70 V vs. RHE) against .[43]

Substances on the GC RDE were collected for SEM, scanning transmission electron microscopy (STEM) and high-resolution transmission electron microscopy (HR-TEM). Energy-dispersive X-ray spectroscopy (EDXS) was used for elemental analyses in the microscopic observations. X-ray diffraction (XRD) patterns of the samples were collected under a parallel-beam configuration. The concentrations of palladium and other elements in the samples were determined by inductively-coupled plasma atomic emission spectroscopy (ICP-AES) after dissolving the samples in 20 mL of nitrohydrochloric acid.

# Calculation methods

First-principles calculations on the adsorption of O and O2 on (111) Pd and PdCu surfaces were carried out to understand the origins for the catalytic activity of BNP-PdCu. The first-principles calculations were performed within the framework of density functional theory[44,45] as implemented in the CASTEP code.[46] The plane-wave basis set was used to calculate the electronic properties, and the generalized gradient approximation of the Perdew-Burke-Ernzerhof[47] functional was used for the exchange-correlation potential for the structure optimizations. Ultrasoft pseudopotentials[48] were used for all the elements in the calculations. We applied cutoff energies of 400 eV. The Brillouin zone was sampled using 5×5×1 Monkhorst-Pack *k*-point meshes.[49] Periodic boundary conditions were applied in the x, y and z directions.

Atomic models are shown in Fig. S2 in Supplementary Material. The rectangular cell consisted of 4 layers of the (111) plane of Pd with a vacuum gap of 15 Å between repeated slabs (Fig. S2(a)). The PdCu model was constructed by substituting one Pd atom at each layer except the top lay with one Cu atom to avoid direct reaction of a Cu atom with an oxygen atom (Fig. S2(b)). A 2% tensile and compressive strain was applied parallel to the exposed surface to identify the strain effects. All cells were constructed from 2 × 2 surface unit cells. The top three layers were relaxed while the bottom layer was frozen during geometry optimization calculations. The binding energy of an oxygen species to a substrate () can be given by:

where is the internal energy of an oxygen species-binding substrate, is the internal energy of a substrate and is the internal energy of an oxygen species. The most stable binding site of an oxygen species was defined as the site where the binding energy was the lowest among the various binding sites shown in Fig. S2, based on a previous study.[50]

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# Supplementary Material

To view supplementary material for this article, please visit [URL to be provided].

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# Tables

Table I. Lattice constants of palladium in bimodal nanoporous palladium-copper.

|  |  |  |
| --- | --- | --- |
| Plane index | Lattice constant (nm) and change from that in bulk palladium ( = 3.890 nm) | |
| Before Cu dissolution | After Cu dissolution |
| 111 | 3.880 (−0.27%) | 3.879 (−0.29%) |
| 200 | 3.888 (−0.05%) | 3.885 (−0.13%) |
| 220 | 3.874 (−0.42%) | 3.879 (−0.29%) |
| 311 | 3.882 (−0.21%) | 3.889 (−0.04%) |
| Average | 3.881 (−0.24%) | 3.883 (−0.19%) |

Table II. Most stable binding sites, their binding energies and d-band centers by first-principles calculations on binding of O and O2 to (111) Pd and PdCu surfaces.

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| System | | Strain | Stable binding site | Binding energy (eV) | d-band center (eV) |
| Pd | O | 0% | fcc | −5.05 | −1.01 |
| +2% | fcc | −5.10 | −1.01 |
| −2% | fcc | −4.99 | −1.01 |
| O2 | 0% | t-f-b | −1.44 | −0.67 |
| +2% | t-f-b | −1.47 | −0.59 |
| −2% | t-f-b | −1.39 | −0.68 |
| PdCu | O | 0% | top1 | −4.74 | −1.10 |
| +2% | hcp3 | −4.93 | −0.99 |
| −2% | top3 | −4.79 | −0.97 |
| O2 | 0% | t-f-bc | −1.27 | −0.64 |
| +2% | t-f-bc | −1.43 | −0.61 |
| −2% | t-b-tb | −1.29 | −0.62 |

# Figure captions

**Figure 1.** (Color online) (a) Schematic illustration of synthesis of bimodal nanoporous PdCu using nanoporous copper as a sacrificial support. (b) Scanning electron microscopy image of nanoporous copper fabricated by dealloying of Cu–Mn alloy.

**Figure 2.** (Color online) Scanning transmission electron microscopy images of bimodal nanoporous PdCu. (a), (b) and left panel in (c) are high-angle annular dark-field images, and the center and right panels in (c) show the elemental mapping by energy-dispersive X-ray spectroscopy. Arrows in (a) shows diameters of replicated pores from nanoporous copper. (d) High-resolution transmission electron microscopy image of bimodal nanoporous PdCu, showing surface lattice contraction.

**Figure 3.** (Color online) (a) Energy-dispersive X-ray spectroscopy results and (b) X-ray diffraction patterns of bimodal nanoporous PdCu before and after copper dissolution.

**Figure 4.** (Color online) (a) Cyclic voltammetry curves for bimodal nanoporous PdCu before and after copper dissolution. The scan rate was 10 mV s−1 and the electrolyte was 0.1 mol L−1 H2SO4. (b) Polarization curves for rotation disk electrodes that load bimodal nanoporous PdCu before and after copper dissolution. The scan rate was 10 mV s−1 and the electrolyte was 0.1 mol L−1 H2SO4.

**Figure 5.** (Color online) Binding energies of O and O2 to Pd surface, (a) O binding energy at 0% strain for fcc, hcp and top sites, (b) O2 binding energy at 0% strain for t-f-b, t-h-b and t-b-t sites, (c) O binding energy at 2% contraction strain for fcc, hcp and top sites, (d) O2 binding energy at 2% contraction strain for t-f-b, t-h-b and t-b-t sites, (e) O binding energy at 2% expansion strain for fcc, hcp and top sites, (f) O2 binding energy at 2% expansion strain for t-f-b, t-h-b and t-b-t sites.

**Figure 6.** (Color online) Binding energies of O and O2 to PdCu surface, (a) O binding energy at 0% strain for fcc, hcp and top sites, (b) O2 binding energy at 0% strain for t-f-b, t-h-b and t-b-t sites, (c) O binding energy at 2% contraction strain for fcc, hcp and top sites, (d) O2 binding energy at 2% contraction strain for t-f-b, t-h-b and t-b-t sites, (e) O binding energy at 2% expansion strain for fcc, hcp and top sites, (f) O2 binding energy at 2% expansion strain for t-f-b, t-h-b and t-b-t sites.

**Figure 7.** (Color online) Charge density distribution between palladium and oxygen atoms, (a) Pd surface with 0% strain, (b) Pd surface with 2% contraction strain and (c) PdCu surface with 0% strain.