Effects of the PdO nanoflake support on electrocatalytic activity of Pt nanoparticles toward methanol oxidation in acidic solutions

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ABSTRACT

We prepared PdO nanoflake thin films on carbon cloths by reactive sputtering deposition, and studied electrocatalytic performance of Pt nanoparticles electrodeposited on the PdO thin film in methanol oxidation reaction (MOR) in acidic media. The PdO nanoflake thin film exhibits a cyclic voltamperometric (CV) behavior similar to metallic Pd electrodes because a metallic Pd surface layer is formed on the PdO thin film electrode under the CV measurement condition. The nanoflake morphology of the PdO electrode provides a large surface area for Pt nanoparticle loading, resulting in a large electrochemical active surface area (ESA). The Pt/PdO electrode has a high electrocatalytic activity toward MOR, and the good electrocatalytic performance of the Pt/PdO electrode is ascribed to a high CO tolerance and the large ESA. We suggest that the high CO tolerance of the catalyst on the Pt/PdO electrode is a result of the synergism of the bifunctional mechanism and the electronic effect. XPS analyses indicate that negative charge transfer occurs from the PdO support to the Pt nanoparticles, indicating the presence of the electronic effect. Pt nanoparticles on the PdO support can greatly alleviate the nanoflake damage during the CV measurement, which results from anodic dissolution of metal Pd from the PdO support in acidic media.

1. Introduction

Platinum has a superior electrocatalytic activity toward methanol oxidation reaction (MOR) and, therefore, Pt or Pt-based alloys is presently the most preferred catalyst used in direct methanol fuel cells (DMFCs). However, practical commercialization of DMFCs using Pt as the catalyst is impeded by many materials difficulties, such as degraded electrocatalytic activity of MOR due to CO poisoning, sluggish oxygen reduction reaction, methanol crossover and the high cost of the Pt catalyst. As far as MOR is concerned, the most widely studied subject in DMFC research is how to enhance the electrocatalytic activity of the anode with a minimized Pt loading. Researchers try to reduce the size and optimize the distribution of Pt nanoparticles to minimize the use of the precious Pt catalyst and concurrently increase the electroactivity surface area (ESA) for methanol oxidation. In addition, Pt-based alloys, such as Pt–Ru, Pt–Ni and Pt–Pd, have been synthesized to improve the electrocatalytic performance of the anode for MOR [1–8]. Metal oxides, such as TiO2, RuO2 and CeO2, were also used as the Pt support to enhance the electrocatalytic efficiency of the Pt catalyst [9–12]. In previous works, we used nanostructured TiO2 and NiO thin films as the support of Pt nanoparticles and studied the effect of the oxide supports on the enhancement of the electrocatalytic activity of the Pt catalyst toward MOR [11,13]. We ascribed the observed electroactivity enhancement to the synergism of the electronic effect and the bifunctional mechanism, which is a combination result of electronic and chemical interactions between the hydrous oxides and Pt nanoparticles. This synergism widely occurs to electrodes using transition metal oxides as the Pt nanoparticle support. We are therefore interested in the enhancement effect of different metal oxide supports on electrocatalytic activity of Pt nanoparticles toward MOR. In this study, we prepared PdO nanostructured thin films, which have a large surface area, by reactive sputter deposition (RSD), and investigated their influence on the electrocatalytic performance of Pt nanoparticles in MOR in acidic media.

Electrochemical properties of PdO is rarely reported in the literature although electrochemical behavior of metallic Pd has been extensively documented [14–25]. However, because the Pd/PdO redox reaction is the primary electrochemical process in the cyclic voltamperometry (CV) test of Pd electrodes, electrochemical properties of PdO may be well interpreted in terms of the CV of metallic Pd. Pd exhibits a high electrocatalytic activity toward MOR and, therefore, has been considered a possible catalyst replacement for Pt in MOR. Since Pd has a much higher dissolution rate than Pt in acidic electrolytes [26,27], MOR electrocatalyzed by the Pd electrode was mostly studied in alkaline solutions. Anodic dissolution of the Pd electrode occurs in acidic media when repetitive Pd oxide formation and reduction take place on the anode during a CV test [28–30]. As a consequence, only a very few reports on MOR catalyzed by Pd-based anodes in acidic media are available in the
In this study, we pulse-electrodeposited Pt nanoparticles on the PdO thin film electrode (denoted by Pt/PdO), and found that the electrodeposited Pt nanoparticles could greatly reduce the anodic dissolution of the PdO electrode in the acidic electrolyte. Moreover, the PdO support can enhance the electrocatalytic activity of Pt nanoparticles toward MOR in the acidic solution; the enhancement is ascribed to the synergism of the bifunctional mechanism and the electronic effect.

2. Experimental

PdO nanoflakes were deposited on carbon cloths at room temperature in a radio frequency magnetron sputter deposition system. Before the PdO deposition, the carbon cloth was immersed in the aqueous solution of 5 M HNO₃ at 25 °C for 5 h, followed by a rinse with deionized (DI) water. The RSD condition for the PdO nanoflake thin film has been described in detail previously [33]. The palladium target was 2 in. in diameter with a purity of 99.99%. The RSD was performed with a gas mixture of Ar (20 sccm) and O₂ (20 sccm) at the working pressure of 9 × 10⁻³ torr and an RF power of 50 W. The as-deposited PdO nanoflake thin film was directly used for electrochemical study without further film treatment.

Pt nanoparticles were electrodeposited on the PdO nanoflake thin film in a mixed aqueous solution of 0.001 M H₂PtCl₆ at room temperature by galvanostatic pulse plating in a two-electrode cell system. The PdO nanoflake support was used as the working electrode and a Pt disk (1.5 cm in diameter) as the counter electrode. The Pt electrodeposition condition, the time durations for the positive current pulse (5 mA) and the negative current pulse (−10 mA) were 1 and 2 ms, respectively. A total of 400 pulse cycles were performed to deposit Pt nanoparticles on the PdO nanoflake support. For comparison, we also pulse-electrodeposited Pt particles on the carbon cloth support (denoted by Pt/C), and prepared a 10 nm thick Pt thin film on the carbon cloth by e-beam deposition (denoted by blanket-Pt). For the preparation of the Pt/C electrode, following the pulse deposition conditions were used: the DC pulse current: −1.0 mA, the pulse duration: 100 ms, the cycle period: 200 ms, and the total number of pulse cycles: 200. The Pt loading of the electrodes was determined by inductively coupled plasma mass spectrometry. Under the sample preparation conditions described above, the Pt/PdO, the Pt/C and the blanket-Pt electrodes had a Pt loading of 0.076, 0.031 and 0.081 mg cm⁻², respectively.

Electrochemical measurements were performed at room temperature in a three electrode cell system with a saturated calomel reference electrode (SCE). The Pt/PdO electrode was used as the working electrode and a Pt disk as the counter electrode. All aqueous solutions were prepared with DI water. The cyclic voltammetry measurement of methanol oxidation was conducted in a deaerated solution of 0.5 M H₂SO₄ and 1 M CH₃OH within the potential range from −0.2 to 0.5 V with a scan rate of 20 mV s⁻¹. The CO stripping measurement was conducted in a 0.5 M H₂SO₄ solution with a scan rate of 20 mV s⁻¹. The CO adsorption on the Pt catalyst was performed by flowing a 1.0% CO/N₂ gas mixture in the 0.5 M H₂SO₄ aqueous solution at 0.1 V for 30 min. Before the CO stripping measurement, the solution was purged with pure N₂ gas for 10 min to remove CO remaining in the solution. The chronoamperometry measurement for methanol oxidation was carried out in the solution of 0.5 M H₂SO₄ + 1 M CH₃OH at 0.5 V for 3600 s.

Surface morphology was studied by scanning electron microscopy (SEM, JEOL JSM-6500F). The chemical composition of the carbon cloth samples was examined by X-ray photoelectron spectroscopy (XPS, Thermo VG 350). The energy calibration for XPS spectra was performed using the Au 4f and the Pt 4f binding energies as the references. The microstructure of the PdO thin film and Pt nanoparticles was studied by transmission electron microscopy (TEM, JEOL JEM-3000F) and X-ray diffractometry (XRD, PANalytical X’Pert Pro) with the Cu Kα source. TEM specimens were prepared by mechanical separating nanoflakes from the carbon cloth using ultrasonic agitation in ethanol.

3. Results and discussion

3.1. Electrochemical properties of the PdO thin film electrode

Fig. 1(a) shows the SEM image of as-deposited PdO nanoflakes on a carbon fiber selected from the carbon cloth shown in the inset. The uniform PdO thin film sputter-deposited on the carbon cloth is composed of continuously interconnected flake-like nanostructures as shown by the magnified SEM image in Fig. 1(b). The PdO nanoflakes are very thin with a flake thickness of ~15–20 nm according to the SEM image, and XRD analysis (not shown) indicates that the room-temperature deposited PdO thin film is amorphous. Pd is generally regarded as a precious metal and, therefore, the use of the PdO support may be considered costly for practical DMFC applications. However, since the height of PdO nanoflakes on the sample is less than 150 nm according to the cross-sectional SEM image (not shown), the consumption of Pd is reasonably low. The nanoflake-like morphology of the as-deposited PdO thin film results in a large surface area, providing enormous electrochemical active sites. Because PdO, a p-type semiconductor, has a low electrical resistivity, we had little difficulty in the
electrodeposition of Pt nanoparticles and electrochemical measurements on the PdO nanoflake thin film.

3.1.1. Electrochemical measurements of the PdO electrode in aqueous H₂SO₄ solutions

To study the electrocatalytic activity of the PdO nanoflake thin film toward MOR in acidic media, we first performed CV measurements of the thin film in an aqueous solution of 0.5 M H₂SO₄, and the CV result is shown in Fig. 2. The CV measurement has a scan potential range between −0.2 and 0.9 V vs. SCE with the scan rate at 20 mV s⁻¹. The cyclic voltammogram shows that, after the first cycle scan, the PdO thin film exhibits a CV behavior characteristic for metallic Pd. To examine if metallic Pd was electrochemically formed on the PdO nanoflake thin film in the acidic solution, we studied chemical states present on the PdO thin film by XPS. Fig. 3 shows Pd 3d XPS spectra of the PdO thin film before and after the CV measurement. For comparison, the Pd 3d spectrum of a metallic Pd thin film, which was sputter-deposited on the carbon cloth, is also presented in the figure. Curve fitting was performed assuming that the Pd⁰ peak has an asymmetric shape and the Pd²⁺ and Pd⁴⁺ peaks have a Gaussian profile. The dashed lines in the figure mark the binding energies of the Pd(0), Pd(II) and Pd(IV) states, which are referred to the literature [34–36]. The as-deposited nanoflake thin film has the Pd 3d₃/₂ peak maximum at 336.5 eV, suggesting that the PdO is the primary chemical state on the film surface. The curve fitting of the broad Pd 3d doublet indicates that PdO₂ is also present in the as-deposited film. After the first cycle of the CV scan, the film surface is greatly reduced to metal Pd as shown by the dramatic change in the shape of the Pd 3d peak profile, which has a large signal component situated at 335.2 eV. Because the Pd 3d signals of the PdO and/or Pd(OH)₂ states are more intense than that of the metal Pd, the metallic Pd surface layer electrochemically formed on the PdO thin film has a thickness much less than the probe depth of the Pd 3d photoelectron, which is about 5 nm when the Mg Kα line is used for the XPS analysis. Although thin, the metallic Pd surface layer must decisively govern the CV behavior of the PdO nanoflake thin film after the first CV cycle.

For CV curves with the scan cycle number larger than one, the anodic and cathodic peaks below 0 V vs. SCE are due to hydrogen desorption and adsorption reactions, respectively, on the metallic Pd surface layer. Palladium has CV responses in acidic media similar to platinum [16,30,37]. In a cyclic voltammogram of metallic Pd in the H₂SO₄ solution, three pairs of CV peaks at potentials below 0.9 V can be observed; the three peak pairs are associated with three redox reactions: hydrogen adsorption/desorption, hydrogen absorption/evolution and Pd oxidation/PdO reduction. Metallic Pd has a great hydrogen absorption capability; the cathodic and anodic peaks due to the hydrogen absorption/evolution reactions may span a large potential range (−0.1 to 0.3 V vs. SCE) depending on the thickness of the metallic Pd layer, and overlap the peaks due to hydrogen adsorption/desorption reactions [38–40]. The anodic wave beginning around 0.45 V results from oxidation of the metallic Pd surface layer during the positive sweep. Since the CV scan limit (0.9 V vs. SCE) is far below the potential required for the formation of PdO₂ (1.4 V) [29,30,41], the Pd(II) state must be the primary oxidation state on the PdO thin film in the potential range between 0.45 and 0.9 V. Electro-oxidation of metal Pd in acidic media has been proposed to proceed via the following reactions [29,30,36]:

\[
Pd + H₂O → PdO + 2H⁺ + 2e⁻, \quad (1)
\]

\[
Pd + 2H₂O → Pd(OH)₂ + 2H⁺ + 2e⁻. \quad (2)
\]

As to be discussed below, the presence of the surface hydroxide is important to the enhancement in the electrocatalytic activity of Pt nanoparticles, which were electrodeposited on the PdO thin film, toward MOR. The cathodic peak between 0.15 and 0.45 V in the reverse sweep is due to the reduction of the surface oxide and hydroxide which are formed during the preceding positive sweep.

3.1.2. Methanol electro-oxidation on the PdO electrode

Fig. 4 shows a cyclic voltammogram of the PdO thin film in an aqueous solution of 1 M CH₃OH + 0.5 M H₂SO₄. After the first scan cycle, the CV profile of the PdO thin film is very similar to that of a Pt electrode. In addition to the hydrogen peaks below 0 V, two anodic peaks are situated in the potential range between 0.2 and 0.8 V. We have performed a CV measurement for the as-deposited PdO thin film in the potential range between 0.4 and 0.9 V, where PdO reduction is unimportant, and the result shown in the inset of Fig. 4 indicates that PdO has little electrocatalytic activity toward MOR. Therefore the anodic peak situated at 0.58 V must be due to MOR occurring on metallic Pd sites, which are produced as a result of the PdO reduction during the potential sweep from −0.2 to 0.45 V. Previous study shows that Pd has an electrocatalytic behavior similar...
MOR between 0.4 and 0.9 V, the large anodic current density above, the PdO electrode has little electrocatalytic activity toward potentials above 0.45 V in the acidic solution and, as discussed [24,47,48]. Because metal Pd sites are absent on the PdO electrode during the reverse sweep from the upper limit potential of the CV scan methanol oxidation on metal Pt sites, which were reactivated during the forward sweep [11,12,45,46]. Note that the MOR anodic peak in the positive sweep has a shoulder feature as well, which is especially obvious for the tenth CV cycle; the shoulder peak is around the onset potential (0.45 V) of the Pd oxidation on the nanoflake thin film. We thus believe that the observation of the shoulder features for both the forward and the reverse anodic peaks is associated with the Pd/PdO redox reaction on the PdO electrode in the acidic solution.

3.2. Electrocatalytic activity of the Pt/PdO electrode toward MOR

Fig. 5(a) shows the SEM image of the PdO thin film after the pulse-electrodeposition of Pt nanoparticles. The Pt nanoparticle-loaded PdO thin film has a flake shape much less sharp than the as-deposited PdO thin film shown in Fig. 1 because of the accumulation of Pt nanoparticles on the ridge of the nanoflake. The nanoparticle accumulation is likely due to that the electric field is concentrated around the ridge area during the electrodeposition, resulting in a higher deposition rate of Pt nanoparticles on the ridge than on the sidewall. The XRD spectrum of the Pt/PdO thin film is shown in Fig. 5(b). The two diffraction peaks situated at 40.2° and 68.3° correspond to the Pt (1 1 1) and the (2 2 0) lattice planes, respectively. The other diffraction peaks shown in the XRD spectrum are due to the hexagonal lattice structure of graphite [49]. The average particle size of Pt nanoparticles on the PdO nanoflakes thin film is 5.5 nm according to the Debye-Scherrer calculation based on the peak width of the Pt (1 1 1) diffraction peak [2,43]; the derived particle size is very close to the size observed in high resolution TEM (HRTEM) images discussed below.

Fig. 6(a) shows the bright field TEM image of electrodeposited Pt nanoparticles on PdO nanoflakes, which were separated from the Pt/PdO electrode by ultrasonic agitation in the ethanol solution. Spots of dark contrast are densely distributed on the PdO nanoflakes. The HRTEM images of the edge of a nanoflake (Fig. 6(b) and (c)) indicate that the dark spots are due to Pt nanocrystals; the two lattice spacings labelled by A and B correspond to the Pt (1 1 1) and (2 0 0) lattice planes, respectively. The size of most Pt nanoparticles is in the range between 3.0 and 5.5 nm, which is reasonably in agreement with the average size derived from the XRD analysis. The Pt nanoparticles uniformly spread over the PdO nanoflake although particle aggregation occurs in some areas. Because of the nanoscale size and the good dispersion of Pt nanoparticles on the PdO nanoflake, a large portion of Pt surface atoms on the nanoparticles measured in the potential range of 0.4–0.65 V (in the reverse sweep) is unlikely due to methanol oxidation on the PdO electrode. Therefore, we believe that the anodic peak in the reverse CV sweep is a result of oxidation of carbonaceous adspecies on the PdO electrode. Accompanied with the anodic peak is a shoulder peak with the peak maximum around 0.33 V, which is close to the peak potential of the PdO reduction (Fig. 2). Note that the MOR anodic peak in the positive sweep has a shoulder feature as well, which is especially obvious for the tenth CV cycle; the shoulder peak is around the onset potential (0.45 V) of the Pd oxidation on the nanoflake thin film. We thus believe that the observation of the shoulder features for both the forward and the reverse anodic peaks is associated with the Pd/PdO redox reaction on the PdO electrode in the acidic solution.
must locate in the peripheral area that adjoins the PdO substrate. As a consequence, numerous Pt electrochemical active sites are subject to strong local chemical and electronic modifications induced by the PdO substrate. These modifications can greatly influence the electrocatalytic activity of the Pt nanoparticles toward MOR.

3.2.1. Methanol electro-oxidation on the Pt/PdO electrode

The cyclic voltammogram of the Pt/PdO electrode in the aqueous solution of 1 M CH₃OH and 0.5 M H₂SO₄ is shown in Fig. 7. For comparison, the figure also presents CV curves of an electron-beam deposited Pt thin film and electrodeposited Pt particles; both types of the Pt electrocatalyst were deposited directly on the carbon cloth, and we refer to the Pt thin film and the Pt particle electrodes thereafter as blanket-Pt and Pt/C, respectively. The MOR peak of the Pt/PdO electrode has the peak maximum around 0.62 V with the onset potential at 0.45 V. The Pt/PdO electrode has both the MOR peak potential and the onset potential less positive than the Pt/C and the blanket-Pt electrodes, indicating that the Pt/PdO electrode has a lower overpotential for methanol electro-oxidation. Moreover, the Pt/PdO electrode has a much larger intensity ratio of the MOR peak (denoted by $I_f$) to the anodic peak in the reverse scan (denoted by $I_b$) than the other two electrodes. The $I_f/I_b$ ratio of the Pt/PdO electrode is 1.87, and the ratio for the Pt/C and the blanket-Pt electrodes is 1.28 and 1.22, respectively. We may use the $I_f/I_b$ ratio as an index to describe the resistance of the electrocatalyst against CO poisoning in MOR. The larger $I_f/I_b$ ratio of the Pt/PdO electrode indicates that Pt nanoparticles on the PdO support have a higher CO tolerance. The efficient removal of CO adspecies is crucial for improving the electrochemical performance in DMFCs. To examine the CO tolerance of the Pt/PdO electrode, we performed the CO stripping measurement for the electrode and the result is presented in Fig. 8. Also shown in the figure are the CO stripping CVs of the Pt/C, the blanket Pt and the as-deposited PdO thin film electrodes. The onset and the peak potentials of the CO stripping peak are listed in Table 1. The Pt/PdO electrode has the CO oxidation peak potential smaller than the PdO thin film electrode by 0.12 V. According to the Pd 3d₃/₂ spectrum (Fig. 3(d)), the PdO thin film is extensively reduced after being polarized at 0.1 V in the 0.5 M H₂SO₄ solution (i.e., the polarization condition for the CO adsorption treatment). Therefore, the CO stripping CV feature of the
PdO thin film electrode actually stems from metal Pd surface sites. The large difference in the stripping potential between the Pt/PdO and the PdO thin film electrodes suggests that the CO stripping peak of the Pt/PdO electrode has a trivial contribution from metal Pd sites. Compared with the Pt/C and the blanket-Pt electrodes, the Pt/PdO electrode has smaller CO stripping peak potential and onset potential, indicating that CO adspecies are easier to be oxidized on the Pt/PdO electrode.

The electrochemical surface area (ESA) of a Pt based catalyst is often determined by the total charge in the hydrogen adsorption (or desorption) region in a CV voltammogram. However, metal Pd has a great capacity of hydrogen absorption, the ESA of the Pt catalyst on the PdO support might be overestimated if the hydrogen region is used to calculate the ESA. We thus used the CO stripping peak to estimate the ESA of the studied electrodes, which is calculated by the following equation:

$$\text{ESA} = \frac{S_R}{M_{Pt}} = \frac{Q_{CO}}{0.484 \times M_{Pt}}$$

where $S_R$ is the real surface area of the catalyst, $M_{Pt}$ represents the Pt loading (mg) on the electrode, $Q_{CO}$ (mC) is the total charge calculated from the CO stripping peak, and the value 0.484 (mC cm$^{-2}$) corresponds to the charge density required for the electro-oxidation of one monolayer of CO adspecies on the Pt surface [50]. The calculated ESA and the real surface area of the three Pt-based electrodes are listed in Table 1. The Pt/PdO electrode has a much larger $S_R$ (14.409 cm$^2$) than the other two electrodes and thus exhibits a much higher electrocatalytic activity toward MOR as shown by the CV curves in Fig. 7. The large $S_R$ of the Pt/PdO electrode is ascribed to the uniform distribution of dense Pt nanoparticles on the PdO nanoflake substrate, which provides a large surface area (12.50 cm$^2$ as listed in Table 1) for the Pt loading. Note that the $S_R$ of the PdO substrate, which was also determined by CO stripping CV measurement, may not truly represent the Pt loading area because the surface PdO was reduced to metallic Pd during the CV measurement and the nanoflake geometry could be significantly modified. However, the large loading area of Pt nanoparticles can still be realized by the measured $S_R$ of the PdO nanoflake support.

### 3.2.2. Synergism of the bifunctional mechanism and the electronic effect

Compared with the two electrodes without the PdO support, the Pt/PdO electrode has a lower overpotential for MOR and a high CO tolerance, suggesting that the PdO nanoflake must play an important role in the enhancement of the electrocatalytic performance of the Pt nanoparticles. The enhancement in the electrocatalytic activity of Pt catalysts toward MOR is generally explained by the bifunctional mechanism and the electronic effect models [1,5,7,11,13,51]. The bifunctional mechanism is widely used to describe how hydroxyl surface groups oxidize CO adspecies on neighboring Pt adsorption sites, thus avoiding the CO poisoning. For the Pt/PdO electrode, Pt nanoparticles are surrounded by abundant Pd-OH surface groups, which are produced when metal Pd sites are oxidized in the acidic electrolyte via Eq. (2). CO adspecies on the periphery of a Pt nanoparticle are readily oxidized by neighboring Pd-OH surface groups because of immediate interactions of the CO adspecies with Pd-OH groups surrounding the Pt nanoparticle [11,13]. This CO oxidation is via a reaction mechanism just like the widely accepted bifunctional mechanism occurring to Pt based binary alloys, such as Pt-Pd and Pt-Ru. Further adsorption of OH species on free Pt sites on the nanoflake may occur by means of OH adspecies migration from the PdO support and/or dissociative adsorption of H$_2$O molecules from the electrolyte. These OH adspecies can progressively oxidize CO adspecies in the inner surface area of the nanoparticle via the Langmuir–Hinshelwood (L–H) reaction mechanism, which describes that two diffusing surface reactants collide with each other to form the product in a surface reaction. The CO oxidation reaction can free more Pt electrocatalytic sites on the nanoparticle, thereby allowing further methanol adsorption and oxidation. A smaller Pt nanoparticle should be less susceptible to the CO poisoning because the bifunctional mechanism and the L–H mechanism can be more kinetically active on the nanoparticle that has a large peripheral region and short diffusion distance.

Besides the bifunctional mechanism, electronic interactions between Pt nanoparticles and the PdO support can also improve the electrocatalytic activity of the Pt catalyst toward MOR. The electronic interaction may modify the electronic structure of the Pt nanoparticle and thus alter the chemisorption behavior of CO adspecies on the nanoparticle. If the adsorption strength of CO adspecies becomes weaker as a result of the electronic structure modification of the Pt nanoparticle, the oxidation of CO adspecies by surrounding OH adspecies may be kinetically favored, leading to a better catalytic activity of the Pt nanoparticle toward MOR.

Fig. 9 shows Pt 4f spectra of the Pt/PdO, the Pt/C and the blanket-Pt electrodes before the electrochemical measurement. The Pt 4f$_{7/2}$ peak of the Pt/PdO electrode is situated at 70.8 eV, which negatively shifts from that of the blanket Pt thin film (71.4 eV) by ~0.6 eV. Previous studies have shown that the binding energy of core level electrons of metal nanoparticles shifts positively from that of the bulk counterpart as a function of the particle size [52–55]. The binding energy shift due to the nanosize effect generally increases with decreasing the size of metal nanoparticles, and the size effect on the energy shift becomes unimportant when the particle size is larger than 3 nm [54]. Therefore, the large negative shift in the Pt 4f$_{7/2}$ binding energy is unlikely to result from the nanosize

### Table 1

<table>
<thead>
<tr>
<th>Pt/PdO</th>
<th>Pt/C</th>
<th>Blanket-Pt</th>
<th>PdO</th>
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<td>Onset potential (V)</td>
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<td>0.52</td>
<td>0.52</td>
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<tr>
<td>Peak potential (V)</td>
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<td>0.58</td>
<td>0.63</td>
</tr>
<tr>
<td>$i_0/i_b$</td>
<td>1.87</td>
<td>1.28</td>
<td>1.22</td>
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<td>Real area ($S_R$) (cm$^2$)</td>
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<td>2.548</td>
<td>4.228</td>
</tr>
<tr>
<td>ESA (cm$^2$ mg$^{-1}$)</td>
<td>189.6</td>
<td>82.2</td>
<td>52.2</td>
</tr>
</tbody>
</table>

* The PdO electrode was used as the Pt support, and hence only its $S_R$ is given here.

Fig. 8. CO stripping CV curves for the Pt/PdO, the PdO thin film, the Pt/C and the blanket Pt electrodes. The scan rate was 20 mV s$^{-1}$. 
effect but is caused by an electronic interaction between the Pt nanoparticle and the PdO support. The negative binding energy shift indicates that negative charge transfer from the PdO support to Pt nanoparticles must occur. The Pt/C electrode has the Pt 4f peak maximum at 71.2 eV, the less negative Pt 4f energy shift compared with the Pt/PdO electrode indicates that Pt particles on the Pt/C electrode are subject to a smaller electronic modification.

In combination with the CV measurements, the XPS analyses clearly show the importance of the PdO substrate to the electronic effect, which can enhance the electrocatalytic activity of Pt nanoparticles toward MOR. The electronic effect must bePrevailing in the periphery of a nanoparticle. Surface atoms in the peripheral region of a smaller Pt nanoparticle should experience a larger electronic modification than those in the inner area as a result of their close proximity to the interface between the Pt nanoparticle and the PdO support, at where the charge transfer takes place [11,13]. A smaller Pt particle size will result in a stronger electronic effect because the peripheral Pt surface atoms occupy a large portion of the surface area of the particle. Most Pt particles on the Pt/C electrode have a size in the order of a few tens of nanometers. The poorer electrochemical performance of the Pt/C electrode compared with the Pt/PdO electrode may be partly due to the larger particle size.

3.2.3. Electrochemical stability of the Pt/PdO electrode in acidic media

To study the electrocatalytic stability of the Pt/PdO electrode in the solution of 1 M CH₃OH + 0.5 M H₂SO₄, we carried out chronoamperometric test at 0.5 V. The chronoamperograms shown in Fig. 10 clearly demonstrates that the Pt/PdO electrode has a much better chronoamperometric performance than the Pt/C and the blanket-Pt electrodes. The Pt/PdO electrode has a gradual decrease in the current density within the first 30 min, and then its current density keeps steady at ~0.8 mA cm⁻². On the other hand, the Pt/C and the blanket-Pt electrodes show a low current density of 0.27 mA cm⁻² and 0.02 mA cm⁻², respectively, after one hour of the chronoamperometric test. The much better chronoamperometric performance of the Pt/PdO electrode can be attributed to the better CO tolerance of Pt nanoparticles. Metal Pd is liable to anodic dissolution in acidic electrolytes and, therefore, is considered an unsuitable electrocatalyst in DMFCs [27,29,30]. The occurrence of anodic dissolution of a Pd electrode during a CV measurement is due to that the surface PdO and Pd(OH)₂, which are formed on the Pd electrode in the forward sweep, is reduced in the reverse sweep, producing Pd²⁺ ions in the solution [29,30]. Since the PdO nanoflake has a CV behavior similar to metal Pd, the anodic dissolution is also expected to occur to the PdO nanoflake electrode. However, the observation that the Pt/PdO electrode has a steady current density after 30 min of the chronoamperometric test suggests that the electrode has a better chemical stability in the acidic solu-

![Fig. 9. Pt 4f spectra of the Pt/PdO, the Pt/C and the blanket-Pt electrodes before the electrochemical measurement.](image)

![Fig. 10. Chronoamperograms of the Pt/PdO electrode, the Pt/C, the blanket-Pt and the PdO electrodes in the aqueous solution of 1 M CH₃OH + 0.5 M H₂SO₄ at room temperature for 1 h. The oxidation potential was kept at 0.5 V vs. SCE.](image)

![Fig. 11. SEM images of (a) the PdO nanoflake electrode and (b) the Pt/PdO electrode after 10 CV cycles in the solution of 1 M CH₃OH + 0.5 M H₂SO₄.](image)
Fig. 11(a) shows the SEM image of a PdO nanoflake electrode after 10 CV cycles in the solution of 1 M CH₃OH + 0.5 M H₂SO₄. The nanoflake morphology of the PdO/C electrode was severely damaged after the CV test, indicating vigorous anodic dissolution of Pt occurred. In contrast, the electrodeposited Pt nanoparticles seem to be able to suppress the anodic dissolution of the PtPdO. Fig. 11(b) shows that the nanoflake feature of the PtPdO electrode after 10 cycles of the CV exhibits a trivial change from that of the as-prepared electrode shown in the SEM image in Fig. 5. It has been reported that the change in the surface chemical composition of Pd-containing alloys can affect the electrode dissolution behavior of the alloys [30, 56, 57]. The electrodeposition of Pt nanoparticles may electrochemically stabilize the PdO nanoflake surface, thereby alleviating the anodic Pd dissolution during the CV measurement. Besides, the chronoamperometric test was performed at 0.5 V, which is above the potential range where the PdO reduction takes place. Therefore, active anodic Pd dissolution as a result of the PdO reduction was unlikely to occur in the test, and a stable chronoamperometric behavior was thus obtained. This argument is supported by the chronoamperometric result of the as-deposited PdO thin film electrode shown in Fig. 10. The PdO electrode is completely inactive under the experimental condition, indicating the absence of metal Pd sites on the electrode. It is likely that the electrochemical stability of the Pt/PdO electrode depends on the amount of Pt loading, and hence an optimal ratio of Pt to PdO (or the surface coverage of Pt nanoparticles on PdO) may exist, which can lead to desirable electrochemical performance and stability of the electrode in the acidic electrolyte. This dependence requires further study, and we are undertaking the study using atomic layer deposition, instead of electrodeposition, to better control the Pt loading on the substrate.

4. Conclusion

We prepared PdO nanoflake thin films on carbon cloths by reactive sputtering deposition, and pulse-electrodeposited Pt nanoparticles on the PdO thin films. The nanoflake morphology of the PdO support provides a large surface area for Pt nanoparticle loading, resulting in a large ESA. The electrocatalytic activity of the PdO thin film and the Pt/PdO electrode toward MOR was studied in acidic media. The PdO nanoflake thin film has a CV behavior similar to a metallic Pd electrode. According to XPS analysis, a thin metallic Pd surface layer is produced on the PdO nanoflake thin film after the first cycle of the CV test in the aqueous H₂SO₄ solution. Because of the repetitive PdO reduction and Pd oxidation in the acidic solution, methanol electro-oxidation on the PdO nanoflake thin film exhibits a CV feature that is closely related to the Pd/PdO redox reaction. The Pt/PdO electrode has a higher electrocatalytic activity toward MOR than the Pd/C and the blanket-Pt electrodes. The much better electrocatalytic performance of the Pt/PdO electrode is ascribed to a high CO tolerance and the large ESA. The high CO tolerance of the Pt-PdO electrode is a result of the synergism of the bifunctional mechanism and the electronic effect. The XPS study shows that negative charge transfer occurs from the PdO support to the Pt nanoparticles. The electronic interaction may modify adsorption properties of adspecies on the Pt nanoparticles, and thus affect electrochemical properties of the Pt/PdO electrode. The nanoflake morphology of the as-deposited PdO thin film is seriously damaged after the CV test because of the anodic dissolution of metal Pd in acidic media. However, the anodic dissolution is greatly alleviated when Pt nanoparticles are electrodeposited on the PdO nanoflake thin film.

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