Nanoporous platinum electrode grown on anodic aluminum oxide membrane: Fabrication, characterization, electrocatalytic activity toward reactive oxygen and nitrogen species

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HIGHLIGHTS

- A new type of nanoporous platinum (NPt) electrode was fabricated on porous aluminum oxide membrane at the first time.
- Electrocatalytic mechanisms of NPt electrode toward NO and nitroxyl were proposed, which depended on pH value of buffer.
- NPt electrode exhibited three times larger catalytic current to O2, NO and nitroxyl than those on nanofoil-Pt electrode.

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ABSTRACT

A new type of nanoelectrode, nanoporous platinum (NPt) electrode was prepared on aluminum oxide membrane by thermal evaporation deposition. The morphology, conductivity and electrocatalytic activity of NPt electrode were characterized and compared with those of nanofoil-Pt electrode through scanning electron microscopy (SEM), electrochemical impedance spectroscopy (EIS) and cyclic voltammetry (CV) techniques, respectively. SEM images showed that “nanocavities” observed in NPt electrode were actually 2-dimensional enclosures by linked nanoparticles. It was different from the conventional arrays of “nanocavities” formed on homogeneous metal films. EIS data indicated that NPt electrode possesses higher conductivity. Compared with that on nanofoil-Pt electrode (14.05 Ω·cm²), the impedance spectrum on NPt electrode exhibits a semicircle portion with much smaller diameters (1.24 Ω·cm² for NPt-100, 1.48 Ω·cm² for NPt-200). Meanwhile, the response sensitivity of NPt electrode to O₂ is 0.85 mA cm⁻², which is larger than that of nanofoil-Pt electrode (0.54 mA cm⁻²). The largest catalytic current for nitric oxide (NO) was obtained in buffer with pH value of 9.4 while for Angeli’s salt (AS) was obtained in buffer with pH value of 5.4. Additionally, electrocatalytic mechanisms of NPt electrode toward NO and AS were proposed, which indicating it depended on pH value of buffer solution.

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1. Introduction

Platinum (Pt) nanoparticles have been received wide interests in various fields, such as biosensor [1], catalysis [2], fuel cell [3], optoelectronic device [4] and so on. Template synthesis is an effective method for preparing nanomaterial with defined dimension [5]. Present techniques to form nanomaterials on templates have been developed including chemical vapor deposition (CVD), sol-gel deposition, polymerization, and electrochemical deposition [6]. However, Pt nanomaterials for electro-catalysis application were depended on synergic effect of other matrix, such as Nafion and carbon, which involves complicated fabrication process. On the other hand, porous anodized aluminum oxide (AAO) substrates have become feasible, low-cost, versatile inorganic materials for wide applications in filtration [7], catalysis [8], biosensing [9] and nanofilm-fabrication [10]. AAO substrates consist of highly-ordered and uniform nanopores with large depth/width ratio (>1000) [11]. The pore diameter and depth can be adjusted by the anodization conditions — acid types, pH value, temperature and anodizing time [12]. AAO is the typical template employed for nano-structural film fabrication. AAO provides a rough surface with large surface area for supporting metal nanoparticles and formed a conductive nano-structural network which facilitates measurement of electrochemical property. Until now, there is few work based on AAO template for Pt nanoparticle film-fabrication.

Reactive nitrogen species including oxide species—nitric oxide (NO) and reduced species—nitroxyl (HNO) in biological system have been known for many years. NO plays an important role in various physiological processes, serving as an endothelial-derived relaxing factor [13], a neurotransmitter [14], and an immune system mediator [15]. Electrocatalytic detection of NO was mainly operated in acidic medium, which limited the application pH range [16]. Therefore, it is significant to study electrocatalytic detection of NO in whole pH range cover from acidic to alkaline solution. Nitroxyl (HNO), a protonated one-electron reduction product of nitric oxide, is a transient species that has been implicated in the biological activity of nitric oxide. It is donated from Angeli’s salt (AS):

\[ \text{HN}_2\text{O}_2^+ \rightarrow \text{HNO} + \text{NO}_2 \]  

In aqueous solution, it spontaneously dimerizes to give nitrous oxide, which results in its role in biology greatly neglected or dismissed:

\[ 2\text{HNO} \rightarrow \text{N}_2\text{O} + \text{H}_2\text{O} \]  

However, many reports have been published to study of HNO biology in recent years. Cetraz [17] et al. investigated a newly synthesized platinum-nitroxyl complex for their potential to circumvent cisplatin resistance in RT112 bladder cancer cells. Recent progress has been made on the elucidation of the mechanisms associated with HNO biological signaling. Importantly, the biochemical mechanisms described for HNO bioactivity are consistent with its unique and novel chemical properties/reactivity [18]. Doctorovich et al. reported the reactions of HNO with metal porphrins. Kinetic and mechanistic studies of these reactions show two alternative operating mechanisms: reaction of the metal center with HNO or with the donor molecules such as trioxodinitrate (Angeli’s salt) [19].

In this work, nanoporous Pt electrode fabricated on AAO substrate was used as a novel electrode for electro-catalysis of reactive nitrogen species-NO and HNO. Meanwhile, Pt nanoparticle was prepared on N-type silicon wafer as a control system, named as nanofilm-Pt electrode. Their morphology and electrochemical performances were studied by scanning electron microscopy (SEM), cyclic voltammetry (CV) and electrochemical impedance spectroscopy (EIS). The electro-catalysis of reactive oxygen (O2) and nitrogen species (NO and HNO) on nanoporous Pt electrode are compared with those on nanofilm-Pt electrode.

2. Experimental section

2.1. Reagent

Porous anodic aluminum oxide (AAO) membrane with average pore-size of 100 and 200 nm was purchased from Whatman (Whatman International Ltd.). The circle-shaped AAO membranes are in a diameter of 13 mm with a thickness of 60 μm. Highly purity NO (99.9%) and O2 (99.7%) were obtained from Zhonghong industry gas company (Shenzhen). Angeli’s salt (AS) was purchased from Cayman chemical co. Stock solutions of AS was prepared in 10 mM NaOH and stored at ~20 °C for further use. All other reagents were of analytical grade, and were used without further purification.

Standard saturated NO solution were prepared in the following procedure: firstly, oxygen-free deionized water was prepared by bubbling N2 gas for 30 min. Then the deionized water bubbled with NO gas for 1 h. Standard saturated O2 solution was also prepared in the similar way. The standard solutions were freshly prepared for each experiment and kept in a glass flask tightly with a rubber septum. The saturated concentration of NO and O2 in water at 25 °C are 1.9 mM and 0.263 mM, respectively. 50 mM phosphate-buffered saline (PBS) containing 0.1 M KCl with pH value of 7.0 was used in this work.

2.2. Preparation of nanofilm-Pt and NPt electrode

The fabrication processes of nanofilm-Pt and NPt electrode are as follows: firstly, N-type silicon wafer was rinsed with a standard “piranha” solution, a mixture of 3:1 (v/v) 98% H2SO4 and 30% H2O2. Meanwhile, AAO substrate was sonicated in anhydrous ethanol for 30 min, rinsed with water, dried in air, and then fixed on a silicon plate. After the pretreatment, the cleaned silicon wafer, pore-size of 100 and 200 nm AAO substrates were used as templates for nanofilm-Pt, NPt-100 and NPt-200 electrodes, respectively. The Pt film were then fabricated by Nexdep Thin film 400 deposition system in base vacuum of approximately 1 × 10⁻7 Torr. The deposition rate was set at 0.1 Å/s. The thicknesses of Pt nanofilm on the three types of electrodes were around 40 nm.

2.3. Measurements

Electrochemical measurements were performed at a CHI 660 E electrochemical workstation (CHI Instruments, Inc., USA). A three-electrode system was employed with Ag/AgCl electrode as reference electrode, a platinum plate as counter electrode, and homemade nanofilm-Pt or NPt electrode (diameter, 3 mm) as working electrode. All the potentials in this work were with respect to Ag/AgCl electrode and all measurements were carried out at room temperature (25 ± 2 °C). EIS experimental was recorded using VersaSTAT4 potentiostat/Galvanostat (Princeton Applied Research, AMETEK Inc, USA). The amplitude of modulation potential is 5 mV, and frequency range is from 10³ to 10⁻³ Hz. All measurements were carried out at room temperature (25 ± 2 °C). Data analysis was performed with ZSimpWin 3.21 using the proposed equivalent circuit. Additionally, scanning electron microscopy (SEM) was obtained on a SU-70 (Hitachi) field emission scanning electron microscopy at an operation potential of 5 kV.
3. Results and discussion

3.1. Morphological characterization

As demonstrated in Fig. 1A and B, small nanoparticles could be observed in SEM images of nanofilm-Pt electrodes, which is due to many platinum nanoparticles having been deposited on silicon surface. In SEM images of nanoporous platinum electrodes (seen in Fig. 1C–F), platinum nanoparticles and holes are linked with each other regularly. Several structural features emerged from the micrographs. First, platinum film was deposited on AAO substrates mainly by surrounding the nanopores. Second, platinum film mainly consists of nanoparticles in good homogeneity. So, platinum nanoparticles link with each other along the AAO substrate frames to form a compact structure. The peculiar structure suggests that electric properties of as-fabricated nanoporous platinum electrode would be affected not only by that of individual particles but also by their linear and ring shaped aggregates. The individual particles and their linear and ring shaped aggregates make the electrode possess larger active electrochemical area and higher conductivity, which facilitate the electron transfer and result in higher electrocatalytic activity of nanoporous platinum electrode. At the same time, the “nanocavities” observed in the nanoporous platinum electrode are actually the 2-dimensional enclosures by linked nanoparticles and are different from the conventional arrays of “nanocavities” formed on a homogeneous metal film [20,21].

3.2. CV characterization

3.2.1. Ru(NH)₆³⁺/²⁺

Fig. 2 demonstrated CVs of Ru(NH)₆³⁺/²⁺ on nanofilm-Pt and NPt electrodes at different scan rates. At scan rate of 50 mV/s, three pair of reversible redox peaks at −0.177 V, −0.188 V, and −0.210 V could be observed for nanofilm-Pt, NPt-100 and NPt-200 electrodes, respectively. Meanwhile, peak currents are linearly related to scan rates in the range from 5 to 50 mV/s on both nanofilm-Pt and NPt electrodes, which indicating a surface electron transfer process occurred [22]. Interestingly, after NPt electrodes undergone CV scanning in Ru(NH)₆³⁺/²⁺ solution, two pairs of broad redox peaks at −0.221 V and −0.225 V in blank buffer solution emerged for NPt-100 and NPt-200, respectively. It may due to Ru(NH)₆³⁺/²⁺ embedded in the nanopore of NPt electrodes after CV scanning processes. For control, only weak reduction peak around −0.2 V was observed for nanofilm-Pt electrode in blank buffer solution (Fig. 2D), which is due to the different structure between NPt and nanofilm-Pt electrodes.

![Fig. 1. Field-emission SEM images of nanofilm-Pt (A, B), NPt-100 (C, D) and NPt-200 (E, F) electrode surfaces. The magnitudes are × 20000 and × 80000, respectively.](image-url)
and NPt-200 (F) electrodes in blank PBS buffer. The scan rate; after CV scanning processes, CV responses of nano-electrodes at different scan rates. Compared with those on nanofilm-Pt electrode, the peak current increased near double and the formal potential shifted from 0.221 V to 0.234 V at scan rate of 50 mV on NPt electrode. It suggests that NPt electrodes possess larger electro-active area. Meanwhile, as shown in the inset figures, peak currents are linear with the scan rate in the range from 5 to 50 mV/s, which indicating a surface electron transfer process occurred.

3.2.2. Fe(CN)₆³⁻/⁴⁻. 
Fig. 3 demonstrated CVs of Fe(CN)₆³⁻/⁴⁻ on nanofilm-Pt and NPt electrodes at different scan rates. Compared with those on nanofilm-Pt electrode, the peak current increased near double and the formal potential shifted from 0.221 V to 0.234 V at scan rate of 50 mV on NPt electrode. It suggests that NPt electrodes possess larger electro-active area. Meanwhile, as shown in the inset figures, peak currents are linear with the scan rate in the range from 5 to 50 mV/s, indicating a surface electron transfer process occurred.

3.2.3. Calculation of electrochemical area 
Electrochemical areas of nanofilm-Pt and NPt electrodes can be determined by Randles-Sevcik equation for a reversible redox couple [23]:

$$I_p = \left( 2.69 \times 10^5 \right) n^{3/2} A D^{1/2} v^{1/2} C_0$$

(3)

where $I_p$ is peak current (A), $n$ is electron transfer number, $A$ is electrode area (cm²), $D$ is diffusion coefficient of electro-active species (cm²/s), $C_0$ is bulk concentration of the same species (mol/cm³), and $v$ is scan rate (V/s). If a reversible electro-active species is used with a known diffusion coefficient, electrode area can be calculated from peak current of a cyclic voltammogram. The diffusion coefficient of 5 mM K₃Fe(CN)₆ in a PBS solution is $1 \times 10^{-5}$ cm²/s at 25 °C [24]. Therefore, electrochemical areas of nanofilm-Pt and NPt electrodes can be obtained from CV data (shown in Fig. 3). Results are listed in Table 1. Obviously, geometric area of nanofilm-Pt electrode is nearly-double larger than that of NPt electrode. It means that electrochemical probe produces larger CV response on NPt electrode than that on nanofilm electrode. Possible reason is as follows: owing to the presence of one pore between the two patch of nanoparticle, the diffusion mode of electrochemical probe is global diffusion on the NPt electrode. While on the nanofilm electrode, the diffusion mode is planar diffusion.

3.3. EIS characterization
Concerning the special structure of nanofilm and nanoporous platinum electrode, present Randles model was not applicable [25,26]. Two new equivalent circuit models with paralleled (RQ) or (RT) component was proposed for nanofilm-Pt and NPt electrodes in this work, shown in Fig. 4. Electronic elements in new equivalent circuits are described below. Solution resistance ($R_s$), is related to the resistance between reference electrode and working electrode. $R_s$ is the electron-transfer resistance; $R_0$ accounts for double layer capacitance; $Q$ acts as a nonlinear capacitor related to film in nanopore and electrode surface; $W$ is the Warburg diffusion resistance; $T$ is the impedance caused by global diffusion. Fig. 5 illustrates the Nyquist plots of nanofilm-Pt, NPt-100 and NPt-200 electrodes in 5 mM Fe(CN)₆³⁻/⁴⁻ solution. Fitted data was also shown as solid points. The impedance spectra include a semicircle and linear portion. The semicircle portion at higher frequency range corresponds to electron-transfer-limited process, and linear portion at lower frequency range represents the.

![](image)

**Fig. 2.** CV responses of nanofilm-Pt (A), NPt-100 (B) and NPt-200 (C) electrodes in PBS buffer containing 5 mM Ru(NH₃)₆Cl₃²⁻ at scan rate of (a) 5, (b) 10, (c) 20, (d) 30, (e) 40, (f) 50 mV/s, respectively; the inset shows relationship between the peak current and scan rate.

**Fig. 3.** CV responses of nanofilm-Pt (A), NPt-100 (B) and NPt-200 (C) electrodes in PBS buffer containing 5 mM Fe(CN)₆³⁻/⁴⁻ at scan rate of (a) 5, (b) 10, (c) 20, (d) 30, (e) 40, (f) 50 mV/s, respectively; the inset shows relationship between the peak current and scan rate.

**Table 1** Comparison of the geometric and electrochemical areas on the nanofilm-Pt electrode with those on the NPt electrode.

| Electrodes | Geometric area a (mm²) | Electrochemical area b (mm²) |
|------------|------------------------|-----------------------------|
| Nanofilm-Pt | 7.05 ± 0.06 | 5.96 ± 0.08 |
| NPt-100 | 3.82 ± 0.04 | 8.95 ± 0.09 |
| NPt-200 | 2.16 ± 0.03 | 9.38 ± 0.11 |

a The geometric area are calculated from the following equation: $A = \pi r^2$ ($r = 1.5$ mm), for NPt-100 and NPt-200 electrodes, the filling factor are 0.5433 and 0.3084, respectively.

b Calculated from equation (3).
diffusion-limited process. Obviously, a semicircle portion with a straight line for the spectrum of Fe(CN)$_6^{3-}$ can be observed on nanofilm-Pt electrode (Fig. 5A). While the spectrum of Fe(CN)$_6^{3-}$ on NPt electrodes shows two quasi-straight lines in high frequency region and low frequency region, respectively (Fig. 5B and C). Compared with that on nanofilm-Pt electrode, there are lines with different slopes on NPt electrodes. It indicates different diffusion-limited processes of Fe(CN)$_6^{3-}$ on NPt electrode surface. Meanwhile, compared with that on nanofilm-Pt electrode (14.05 $\Omega$ cm$^2$), the impedance spectrum on NPt electrodes exhibited a semicircle portion with much smaller diameters (1.24 $\Omega$ cm$^2$ for NPt-100, 1.48 $\Omega$ cm$^2$ for NPt-200). It means that the electron-transfer rate increases rapidly on NPt electrodes.

As demonstrated in Fig. 5, the EIS data was fitted by proposed equivalent circuit effectively. Results are listed in Table 2. $C_{dl}$ accounts for double layer capacitance, which depends on many variables including electrode potential, temperature, ionic concentrations, oxide layers, and electrode roughness etc. These variables are the same except electrode roughness between the nanofilm Pt and NPt electrodes. The larger the electrode roughness, the thicker the diffusion layer. Therefore, the larger the double layer capacitance. Owing to the nanopore structure, the electrode roughness of NPt electrode is larger than that of nanofilm Pt electrode. Therefore, $C_{dl}$ on NPt electrode (155 and 340 $\mu$F/cm$^2$) is larger than that of nanofilm Pt electrode (24.3 $\mu$F/cm$^2$). The Warburg diffusion resistance ($W$) was incorporated in equivalent circuit model due to planar diffusion occurred on nanofilm electrode. Meanwhile, $\Gamma$ impedance was incorporated in equivalent circuit model due to global diffusion occurred on NPt electrode. Since there are two different diffusion types on the two electrodes, no comparability of the $W$ and $\Gamma$ value for nanofilm and NPt electrodes.

### 3.4. Electrocatalytic activity of NPt electrode toward O$_2$, NO, AS

Since NPt-200 electrode possesses a little larger electrochemical area than that of NPt-100 electrode, the former was adopted for electrocatalytic activity study in the following experimental. Under normal condition (101.325 kPa, 25 $^\circ$C), the concentration of saturated O$_2$ in water is 0.263 mM. Fig. 6 shows CV responses of nanofilm-Pt and NPt-200 electrode in the presence of saturated O$_2$. The electrocatalytic reduction potential occurred at $-0.012$ V for nanofilm-Pt electrode and $0.094$ V for NPt-200 electrode, respectively. The positive shift in potential is 0.1058 V, which indicating NPt electrode possesses higher electrocatalytic activity for. Meanwhile, the catalytic sensitivity of NPt-200 electrode to O$_2$ is 0.85 mA cm$^{-2}$, which is larger than that of nanofilm-Pt electrode (0.54 mA cm$^{-2}$).

Fig. 7 demonstrated the catalytic responses of nanofilm-Pt and NPt-200 electrode to NO in buffers with different pH values. For

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**Table 2**

| Parameters       | Nanofilm-Pt | NPt-100 | NPt-200 |
|------------------|-------------|---------|---------|
| $R_s$ ( $\Omega$ cm$^2$) | 10.24       | 15.06   | 16.20   |
| $C_{dl}$ ( $\mu$F/cm$^2$) | 24.3        | 155     | 340     |
| $R_{ct}$ ( $\Omega$ cm$^2$) | 14.05       | 1.24    | 1.48    |
| $Q_w$ | 0.0033 | 0.0016 | 0.0016 |
| $Q_n$ | 0.70      | 0.53    | 0.58    |
| $R_1$ ( $\Omega$ cm$^2$) | 3.45        | 44.13   | 160.17  |
| $Q_{w0}$ | NA        | 0.00015 | 0.0021  |
| $Q_{n0}$ | NA        | 0.75     | 0.93     |
| $R_2$ ( $\Omega$ cm$^2$) | NA         | 490     | 529     |
| $\Gamma$ ( $\Omega$ cm$^2$) | $1.37 \times 10^6$ | $1 \times 10^6$ | $7.25 \times 10^6$ |
| $\Delta$ ( $\Omega$ cm$^2$) | NA         | 4.06    | 59.29   |

NA: not applicable.
nanofoil-Pt electrode, reduction peaks were observed at $-0.40$ V in pH 5.4 solution, $-0.75$ V in pH 7.4 and pH 9.4 solutions, respectively. Meanwhile, oxidation current was observed in pH 9.4 solution although no peak emerged. While no oxidation current was observed in pH 5.4 and 7.4 solution. For NPt-200 electrode, reduction peaks were observed at $-0.52$ V in pH 5.4 solution, $-0.68$ V in pH 7.4 and $-0.60$ V in pH 9.4 solutions. Meanwhile, oxidation peak current was observed at $0.90$ V in pH 5.4 solution, $0.84$ V in pH 7.4 and $0.95$ V in pH 9.4 solutions. At all the three pH values (5.4, 7.4 and 9.4) of buffer solutions, NPt-200 electrode exhibited almost ten times larger catalytic current both in the oxidation and reduction potential range than those for nanofoil-Pt electrode. It indicates that the NPt electrode exhibits better catalytic activity to NO than that of nanofoil-Pt electrode. Meanwhile, the catalytic current increases with the increase of pH value, which indicating that alkaline condition is beneficial to NO electrocatalysis.

Fig. 8 demonstrates CV responses of AS in different pH solution on nanofoil-Pt and NPt-200 electrodes. Generally, oxidation and reduction peaks emerged on both nanofoil-Pt and NPt-200 electrodes in all the three pH value (5.4, 7.4 and 9.4) of buffer solutions except that on nanofoil-Pt electrode in pH value of 9.4. It may be due to more stability of AS in pH 9.4 solution and weaker catalytic activity of nanofoil-Pt electrode. Comparatively, larger catalytic current was observed on NPt-200 electrode in the same pH value solution, and larger catalytic current was obtained in the pH value of 5.4 at the same electrode. The reason is as follows: first, due to its frame and porous structure, NPt-200 electrode possesses larger electro-active area than that of nanofoil-Pt electrode; second, among the three pH value of buffer solutions, the decomposing rate of AS to HNO is fastest in pH 5.4.

Fig. 7 shows CV responses of nanofoil-Pt and NPt-200 electrodes to different concentration of AS in pH 7.4 buffer. Obviously, an increase in anodic peak current and a decrease in cathodic peak current were observed after the addition of AS. Meanwhile, the response currents are linear with the AS concentrations. The increase for NPt-200 electrode was larger than that for nanofoil-Pt electrode at the same concentration of AS. Obviously, with the increase of AS concentration, the oxidation potential shifts positively and the reduction potential shifts negatively.

3.5. Electrocatalytic mechanism of NPt electrode toward NO and HNO

The electrocatalytic reduction mechanisms of NPt electrode toward NO and HNO are as follows, taking NO as an example:

When solution pH $> 7$,

$$\text{Pt}^* + \text{NO}_{(sol)} \rightarrow \text{Pt} - \text{NO}_{(ads)} \quad (4)$$

$$\text{Pt} - \text{NO}_{(ads)} + 2\text{H}^+ + 2e^- \rightarrow \frac{1}{2}\text{N}_2 + \text{H}_2\text{O} + \text{Pt}^* \quad (5)$$

When solution pH $< 7$,

$$\text{Pt}^* + \text{H}^+_{(sol)} + e^- \rightarrow \text{Pt} - \text{H}_{(ads)} \quad (6)$$

$$2\text{Pt} - \text{H}_{(ads)} + \text{NO} + e^- \rightarrow 2\text{Pt}^* + \text{H}_2\text{O} + \frac{1}{2}\text{N}_2 \quad (7)$$
The electrocatalytic oxidation mechanisms of NPt electrode toward NO and HNO are as follows, also taking NO as an example:

\[ \text{Pt}^+ + 2\text{e}^- \rightarrow \text{Pt}^{2+} \quad \text{(fast)} \quad (8) \]

\[ \text{Pt}^{2+} + \text{NO}_{\text{ads}} \rightarrow \text{Pt}^{2+} - \text{NO}_{\text{ads}} (9) \]

\[ \text{Pt}^{2+} - \text{NO}_{\text{ads}} + 2\text{H}_{\text{2}}\text{O} - 3\text{e}^- \rightarrow \text{NO}_2^+ + 4\text{H}^+ + \text{Pt}^{2+} (10) \]

Then toward cathodic potential scan, the reduction of Pt oxide layer occurred:

\[ \text{Pt}^{2+} + 2\text{e}^- \rightarrow \text{Pt}^0 \quad (11) \]

4. Conclusions

In this work, a new type of nanoporous platinum (NPt) electrode was successfully prepared on a porous aluminum oxide membrane by thermal evaporation deposition technology. The morphology study showed that the stereo-structure of “nanocavities” on NPt electrode make it possesses larger specific surface area. EIS data indicated that NPt electrode possesses higher conductivity. The electrocatalytic studies indicated that NPt electrode exhibited higher electrocatalytic activity to reactive oxygen (O_2) and nitrogen species (NO and Angeli’s salt) than those of nanofilm electrode. In electrocatalytic reduction of O_2, the positive shift in reduction potential was 105.8 mV, which indicating NPt electrode possesses higher electrocatalytic activity. In both the oxidation and reduction potential range, NPt-200 electrode exhibited almost ten times and three times larger catalytic current than those for nanofilm–Pt electrode toward electrocatalysis of NO and AS, respectively. Electrocatalytic oxidation and reduction mechanisms toward NO and AS depend on pH value of buffer solution.

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