Improved Electrocatalytic Activity and Durability of Pt Nanoparticles Supported on Boron-Doped Carbon Black

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Abstract: A facile strategy is proposed to synthesize boron-doped ECP600 carbon black (B-ECP600), and the catalyst of Pt supported on boron-doped ECP600 (Pt/B-ECP600) shows smaller particle sizes and a higher electrochemical surface area (95.62 m²·g⁻¹) and oxygen reduction reaction activity (0.286 A·mg⁻¹ for mass activity; 0.299 mA·cm⁻² for area specific activity) compared to the catalyst of Pt supported on ECP600 (Pt/ECP600). The results show that the boron doping of the carbon supports plays an important role in controlling the size and dispersion of Pt nanoparticles and the O₂ adsorption/dissociation of the oxygen reduction reaction. A further accelerated durability test proves that boron doping can greatly enhance the stability of carbon support and thus improves the electrochemical performance of the catalyst during the long-time running. All these results suggest boron-doped carbon has great potential for application in fuel cells.

Keywords: ORR; boron doping; electrocatalyst; Pt; carbon black

1. Introduction

Proton exchange membrane fuel cells (PEMFCs) have wide application prospects for their high energy density and low pollution [1,2]. Pt loaded on carbon black is the most commonly used electrocatalyst for cathode oxygen reduction reaction (ORR) of PEMFCs. However, in addition to the high cost, several drawbacks of the catalyst, such as low activity and poor durability in practical applications, have limited the further commercialization of the Pt/C catalyst [3,4]. Carbon hybrid materials, metal-free carbon-based materials, and alloy catalysts are considered as effective approaches to solve the above problems [5–9]. For example, doping heteroatoms such as N, P, and B into the frameworks of carbon materials can improve their electrocatalytic activity, as doped heteroatoms can break the electrical neutrality of adjacent carbon atoms and create active sites, thus facilitating the adsorption of oxygen molecules and the oxygen reduction reaction [10–12]. The advantages of doped carbon materials as the catalyst supports include promoting the dispersion of the catalyst, improving Pt nanoparticle adhesion, and better adsorption strength for oxygen-containing species [12–16].

Recently, a variety of studies on doped carbon materials have made some achievements. Cao et al. prepared the nitrogen and boron co-doped hollow carbon tube (NB-HCT) catalyst from the fruit fiber of platanus by a facile three-step method [17]. Tam et al. synthesized B-doped graphene quantum dots (BGQD) as a metal-free electrocatalyst for oxygen reduction reaction by carbonizing the glucose with different concentrations of boric acid through hydrothermal treatment [18]. Du et al. synthesized N-doped carbon-stabilized PtCo nanoparticles (PtCo@NC) via thermal
decomposition of a Pt-incorporated Co-based zeolitic imidazolate framework (Pt@ZIF-67) [19]. Nsabimana et al. synthesized boron-doped ordered mesoporous carbons (BOMC) by a nano-casting method, using SBA-15 mesoporous silica as template, and 4-hydroxyphenylboronic acid and sucrose as boron and carbon sources, respectively [16]. Despite the progress that has been made in the research of carbon hybrid materials, the following problems still exist in the actual production or application process: there are only a few studies demonstrating the synthesis of B-doped carbon materials and their applications due to the difficulty of synthesis, the raw materials are environmentally unfriendly and unhealthy, the process is complex and not suitable for mass production, and some doped carbon materials only show good ORR activity in an alkaline environment, which is not suitable for the acidic environment of PEMFCs, and so on [15,20–22].

In this study, we doped boron into ECP600 carbon black by a simple method: Nickel was firstly loaded into carbon black, and then boron was substituted into carbon black with sodium borohydride (NaBH₄) as a structure-directing agent. Furthermore, the catalysts of Pt nanoparticles supported on ECP600 with and without boron-doping were prepared. The result of the electrochemical test shows that the platinum catalyst using B-doped ECP600 (B-ECP600) as support exhibits much higher electrocatalytic activity (95.62 m²·g⁻¹ of the electrochemically-active surface area (ECSA) and 0.286 A·mg⁻¹ of mass activity (MA)) and better stability than the same type of catalysts supported on ECP600 without boron-doping. In addition, X-ray powder diffraction (XRD), transmission electron microscopy (TEM), X-ray photoelectron spectroscopy (XPS), Fourier-transform infrared spectroscopy (FTIR), and the nitrogen adsorption–desorption isotherm have also been used to explain the effects of B-doped carbon blacks. Meaningfully, the above doping process is simple and easy to repeat and thus has great potential for large-scale production.

2. Results and Discussion

2.1. Characterization of B-ECP600

The XRD patterns of ECP600 and B-ECP600 are shown in Figure 1a. The XRD patterns of ECP600 and B-ECP600 both exhibit two diffraction peaks, which correspond to the (002) diffraction peak of graphite at 2θ value of ca. 25° and the (100) diffraction peak at ca. 43.5°. Compared to the ECP600, a slight shift of the (002) diffraction peak toward the negative 2θ value can be seen from the XRD pattern of B-ECP600, proving that the incorporation of B leads to the increase of interlayer spacing.

The existence of B in B-ECP600 was proved by XPS. The XPS spectra of B1s narrow-scans of the B-ECP600 are shown in Figure 1b. By analyzing the detailed B1s XPS spectra, we could find a peak at 191.9 eV, corresponding to the oxidized state of B. The atomic concentrations of B and Ni in the B-ECP600 were 0.5% and 0.09%, respectively, as estimated from the intensities of the corresponding XPS peaks. Figure 1c shows the C1s XPS spectrum of the B-ECP600. We could accordingly deconvolve C1s peak into 3 peaks at 283.8, 285.4, and 289.2 eV, corresponding to C=C/C–C (carbon backbone), C–O (epoxy and/or alkoxy), and COOR (carboxylic and/or lactones), respectively [23]. The results of the FTIR (Figure 1d) also show that B-ECP600 has varied oxygen-containing groups.

The nitrogen adsorption–desorption isotherms and the corresponding pore size distribution curves of B-ECP600 and ECP600 are presented in Figure 1e,f, respectively. It can be seen from Figure 1a that the nitrogen adsorption–desorption isotherms of B-ECP600 and ECP600 exhibit type IV curves with the hysteresis loops at lower (0.4–0.8) and higher (0.8–1.0) relative pressures, which indicate the existence of mesopores and macropores [24]. As shown in Figure 1f, the primary pore size of B-ECP600 is 3.66 nm, while ECP600 displays a primary pore size of 3.80 nm. The Brunauer–Emmett–Teller (BET) surface area of B-ECP600 and ECP600 were calculated to be 957.5 and 1416.5 m²·g⁻¹, respectively. A lower BET surface area indicates that the mesopores and micropores in B-ECP600 have been collapsed, and thus, the conductivity was improved [25]. The total pore volumes are 1.75 and 2.07 cm³·g⁻¹ for B-ECP600 and ECP600, respectively.
Figure 1. (a) XRD results of ECP600 and B-ECP600; (b) B1s; (c) C1s narrow-scan of B-ECP600; (d) FTIR spectrum of B-ECP600; (e) nitrogen adsorption–desorption isotherms; and (f) pore size distributions of ECP600 and B-ECP600.

The morphology and structure of ECP600 and B-ECP600 were characterized by TEM. The TEM image of ECP600 (Figure 2a) shows the branched chain structure composed of carbon nanospheres. Figure 2b shows the TEM image of B-ECP600 without obvious carbon nanospheres. It can be seen that the structure of B-ECP600 has become much thinner and exhibits a partial sheet-like feature. This structural change is due to the NaBH₄ as a structure-directing agent [26]. This unique structure of B-ECP600 can provide more defect sites for platinum particles in the process of catalyst preparation and thus improve the electrochemical performance of the catalysts.
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Figure 2. TEM images of (a) ECP600 and (b) B-ECP600.

The proposed mechanism for the formation of B-ECP600 is schematically presented in Scheme 1. First, Ni atoms were loaded on ECP600 under a temperature of 170 °C. Especially at high temperature, ethylene glycol, as a relatively gentle reducing agent, could keep the reaction rate in a suitable range, so that Ni atoms could be more evenly dispersed on carbon black, which was beneficial to the subsequent boron doping. At the same time, the whole reduction process was carried out in nitrogen atmosphere, which could prevent the oxidation of the reduced Ni under high temperature. On the one hand, the Ni (0) supported on ECP600 could catalyze the decomposition of the NaBH4 into solid B(s) [27]. The generated B(s) underwent an alloying process immediately with the active Ni (0) to form amorphous Ni–B alloys and then were further oxidized by oxygen in the air [28]. On the other hand, NaBH4 can be hydrated into BO(OH)2, which exhibits triangle plane symmetry. The hydroxyl groups of BO(OH)2 could condense with hydroxyl groups from carbon nanospheres. Thus, the carbon nanospheres were interlinked by boron–oxygen–carbon bonds and finally formed the boron-doped carbon [23].
2.2. Characterization of Pt/B-ECP600

The structural characterization and the Pt particle size distributions of the Pt/ECP600 and Pt/B-ECP600 catalysts are shown in Figure 3. By comparing the TEM images in Figure 3a,b, we can find that Pt nanoparticles tend to aggregate to form large particle clusters on the ECP600 surface, while the Pt nanoparticles on B-ECP600 are uniformly distributed and exhibit good dispersion. It can be seen from the histograms in Figure 3c,d that the average diameter of 100 Pt particles in Pt/ECP600 was 2.90 nm and that of 100 particles in Pt/B-ECP600 was 2.19 nm, which is consistent with the results from the analysis of the TEM image. B-ECP600 has more defect sites, which provides a strong interaction and anchoring location for nonuniform nucleation of Pt, which is one of the reasons for the high dispersion of Pt/B-ECP600 [16]. The smaller particle size and better dispersion of Pt/B-ECP600 may contribute to higher Pt utilization and better electrochemical performance compared to Pt/ECP600 [29].

![Scheme 1. Schematic representation of the formation mechanism of B-ECP600.](image-url)
The detailed Pt4f XPS spectra of the Pt/B-ECP600 sample are shown in Figure 4a. It is clearly shown that there are two peaks corresponding to Pt4f7/2 and Pt4f5/2, which are evident for the presence of Pt nanoparticles on B-ECP600. The XRD patterns of the prepared Pt/ECP600 and Pt/B-ECP600 are shown in Figure 4b. Both samples show the (002) plane of carbon at 2θ value of ca. 25°. The characteristic diffraction peaks located at 2θ values of around 39.8°, 46.3°, and 67.6° correspond to the (111), (200), and (220) planes of Pt, suggesting both catalysts display the characteristics of the Pt face-centered cubic (fcc) structure. The XRD pattern of Pt/B-ECP600 also shows that no PtNi alloy was formed, despite the addition of nickel in the preparation process. By analyzing XRD patterns with the Bragg formula, we found that the lattice spacing of Pt/ECP600 and Pt/B-ECP600 was calculated to be 0.225 and 0.226 nm, which agrees well with the result of TEM.

Figure 4. (a) XPS patterns of Pt 4f for Pt/B-ECP600; and (b) XRD results of ECP600 and B-ECP600.
2.3. Electrochemical Properties of Pt/B-ECP600

Pt/ECP600, Pt/B-ECP600, and B-ECP600 were characterized electrochemically in N2-saturated 0.1 M HClO₄ solution at a sweep rate of 50 mV/s (Figure 5a). The ECSAs of Pt/B-ECP600 and Pt/ECP600 were 95.62 and 76.57 m²·g⁻¹Pt⁻¹, respectively, calculated by integrating the underpotentially deposited hydrogen adsorption charge and normalizing with scan rate, Pt loading, and the charge value of 210 µC·cm⁻² for the Pt surface. The higher ECSA of Pt/B-ECP600 is due to the relatively smaller particle size and uniform deposition of Pt nanoparticles on B-ECP600. As seen in Figure 5a, the maximum of the oxide reduction peak in the cathodic sweep of Pt/B-ECP600 is slightly more negative than that of the Pt/ECP600, which is consistent with the general belief that smaller Pt particles lead to a negative shift in peak potential for the reduction of the oxygenated species [30,31]. The negative shift of reduction peak of Pt/B-ECP600 also suggests an advanced electrocatalytic activity of the Pt/B-ECP600 toward ORR compared with Pt/ECP600. To better understand the characteristics of catalysts supported on B-doped ECP600, the electrocatalytic behavior of Pt/B-ECP600 and Pt/ECP600 was studied by investigating their electrocatalytic activity in the oxygen reduction reaction. Figure 5b shows the linear sweep voltammetry (LSV) polarization curves of two prepared catalysts in O₂-saturated 0.1 M HClO₄ solutions at the temperature of 30 °C. The ORR half-wave potential of Pt/B-ECP600 was about 0.891 V compared to the 0.878 V of Pt/ECP600, indicating the improved activity of Pt/B-ECP600. The mass activity (MA) and area specific activity (SA) of Pt/B-ECP600 were calculated to be 0.286 A·mg⁻¹Pt⁻¹ and 0.299 mA·cm⁻², respectively, by normalizing the kinetic current at 0.9 V with the mass and the ECSA of the Pt on the GC electrode. The MA and SA of Pt/ECP600 were 0.217 A·mg⁻¹Pt⁻¹ and 0.283 mA·cm⁻², respectively, as shown in Table 1. For comparison, He et al. have prepared boron-doped Pt/C catalyst by adding doped boron into commercial Pt/C (20 wt%, JM) catalyst and improved activities (ECSA went from 71 to 73 m²·g⁻¹Pt⁻¹; MA (@0.95V) went from 0.050 to 0.150 A·mg⁻¹Pt⁻¹) [28]. The electrochemical performances of the catalysts were greatly improved by boron doping.

![Figure 5](image_url)

**Figure 5.** Electrocatalytic properties of the prepared catalysts: (a) Cyclic voltammogram (CV) curves and (b) linear sweep voltammetry (LSV) polarization curves of Pt/B-ECP600 and Pt/ECP600.

**Table 1.** The electrochemically-active surface area (ECSA), mass activity (MA), and area specific activity (SA) of Pt/ECP600 and Pt/B-ECP600.

| RDE TESTING | ECSA (m²·g⁻¹Pt⁻¹) | MA (A·mg⁻¹Pt⁻¹) | SA (mA·cm⁻²) |
|-------------|-------------------|-----------------|--------------|
| Pt/ECP600   | 76.57             | 0.217           | 0.283        |
| Pt/B-ECP600 | 95.62             | 0.286           | 0.299        |

The result of XRD suggests that the enhancement of ORR activities of Pt/B-ECP600 is not due to the PtNi alloy effect. Owing to the smaller electronegativity of B, boron doping is favorable to capture the O₂ molecules [12]. However, it can be seen from Figure 6a that the B-ECP600 exhibits poor activity in acidic media, indicating that B-ECP600 might not be oxophilic enough to initiate chemisorption of O₂. Previous studies suggest that boron oxide could facilitate O₂ adsorption and
weaken O and OH binding energy on Pt (111) to a respective optimal range of better electrocatalysis for ORR [8,28,32,33]. Therefore, the enhancement of ORR activities of Pt/B-ECP600 is considered to be due to the electronic effect of B-ECP600 on the Pt-O and Pt-OH chemical bonding, which is implemented by the electron-deficient boron oxide dominating electron transfer [28]. On the other hand, the spillover effect [34] may also have played an important role in improving ORR performance of Pt/B-ECP600: Due to the oxyphilicity of boron dopant on B-ECP600 supports [12], a part of adsorbed oxygen atoms dissociated from O₂ molecules on Pt may be transferred to the adjacent B atoms and further reduced to H₂O. The analyses above suggest that the B-ECP600 has a synergetic coupling effect in the ORR electrocatalysis, and in some way, boron on B-ECP600 has taken part in the ORR and thus increased the overall ORR performance, as shown in Scheme 2.

Furthermore, in order to compare the stability of carbon supports, we assessed the long-term durability of the Pt/B-ECP600 and the Pt/ECP600 by an accelerated durability test (ADT) under a scanning rate of 100 mV s⁻¹ from 1.0 to 1.6 V (vs. reversible hydrogen electrode (RHE)) in 0.1 M HClO₄. We compared the ECSAs of Pt/B-ECP600 and Pt/ECP600, respectively, after 5 K cycles of ADTs, because the ECSA of catalysts is greatly affected by the carbon supports. It can be seen from Figure 6a–c that both as-prepared catalysts exhibit the decrease of ECSA during the ADT, while the ECSA after the ADT of Pt/B-ECP600 is still much higher than that of Pt/ECP600. As shown in Figure 6d, the ECSA of Pt/B-ECP600 after the ADT was calculated to be 62.68 m²·g⁻¹Pt⁻¹, exhibiting a loss of 34.4%. In contrast, the ECSA of Pt/ECP600 after the ADT was 44.28 m²·g⁻¹Pt⁻¹ with a drop of 42.2%. After a relatively high potential range ADT, the main causes of ECSA loss are the aggregation and falling off of Pt due to corrosion of carbon support [35,36]. The result of ADTs indicates that the B-ECP600 has better corrosion resistance compared with the ECP600. It can be seen from Figure 6a,b that the hydrogen adsorption/desorption peaks near 0.27 V of Pt/B-ECP600 and Pt/ECP600, and both become more pronounced after the ADT, suggesting that the particle size of the two samples has been increased after ADTs [31]. Figure 7 shows the TEM images of the two catalysts after the ADT. Partial Pt nanoparticles of Pt/ECP600 were aggregated after the ADT, while the Pt nanoparticles of Pt/B-ECP600 still maintained good dispersion and small size. As calculated from the TEM images, the average particle size of Pt/B-ECP600 changed from 2.19 to 2.48 nm, while the average particle size of Pt/ECP600 changed from 2.90 to 3.25 nm. In addition, the MA of Pt/B-ECP600 after the ADT was calculated to be 0.230 A·mg⁻¹Pt⁻¹, still much higher than that of Pt/ECP600 (0.180 A·mg⁻¹Pt⁻¹). These results demonstrate that, as the support of catalysts, B-ECP600 is remarkably durable compared to ECP600, which suggests using B-ECP600 as support for Pt catalysts is a promising way to improve the durability of the catalysts.
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being washed with ultrapure water and dried at 60 °C for 10 h. Then, the nickel–carbon was dissolved
were added when the mixture cooled to room temperature. The nickel–carbon was obtained after
prepared sodium borohydride aqueous solution (10 wt %). After being ultrasonically stirred for 1 h, the mixture was heat-
(1 wt %), 130 mL ethylene glycol, 85 mL ultrapure water, and 1g sodium
hydroxide aqueous solution (10 wt %). After being ultrasonically stirred for 1 h, the mixture was heat-
NaOH), nickel chloride hexahydrate (NiCl2•6H2O, 99.0%), sodium borohydride (NaBH4, 99.0%), chloroplatinic acid hexahydrate (H2PtCl6•6H2O, 99.0%), perchloric acid (HClO4, 70.0~72.0%),
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ECP600 and Pt/ECP600 after 5 K cycles of ADTs; and (d) ECSAs of Pt/B-ECP600 and Pt/ECP600 before and after 5 K cycles of ADTs.

Figure 6. Electrocatalytic durability of carbon supports. CV curves of (a) Pt/B-ECP600 and (b) Pt/ECP600
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before and after 5 K cycles of accelerated durability tests (ADTs); (c) CV curves of Pt/B-ECP600 and
Pt/ECP600 after 5 K cycles of ADTs; and (d) ECSAs of Pt/B-ECP600 and Pt/ECP600 before and after 5 K cycles of ADTs.

Figure 7. TEM images of (a) Pt/B-ECP600 and (b) Pt/ECP600 after 5K cycles of ADTs.
3. Materials and Methods

3.1. Chemicals

Carbon black (Ketjenblack ECP600JD) was obtained from Lion Corporation (Japan). Sodium hydroxide (NaOH), nickel chloride hexahydrate (NiCl$_2$•6H$_2$O, 99.0%), sodium borohydride (NaBH$_4$, 99.0%), chloroplatinic acid hexahydrate (H$_2$PtCl$_6$•6H$_2$O, 99.0%), perchloric acid (HClO$_4$, 70.0~72.0%), ethyl alcohol (C$_2$H$_6$O, 99.7%), ethylene glycol (C$_2$H$_6$O$_2$, 99.5%), and hydrochloric acid (HCl, 38.0%) were all obtained from Sinopharm Chemical Reagent Co., Ltd. (Beijing, China). All aqueous solutions were prepared using ultrapure water (18.25 M$\Omega$ cm$^{-1}$) purged by an ultrapure purification system (EPED-10TS, Chinese EPED Corporation).

3.2. Preparation of Support and Catalysts

B-ECP600 carbon black was prepared via a simple method using nickel as the medium and sodium borohydride as the boron source. A total of 430 mg carbon black was mixed with 13.62 g NiCl$_2$•6H$_2$O/EG solution (1 wt %), 130 mL ethylene glycol, 85 mL ultrapure water, and 1 g sodium hydroxide aqueous solution (10 wt %). After being ultrasonically stirred for 1 h, the mixture was heat-treated at 170$^\circ$C for 1 h in nitrogen atmosphere. A total of 3 mL hydrochloric acid and 20 mL EA were added when the mixture cooled to room temperature. The nickel–carbon was obtained after being washed with ultrapure water and dried at 60$^\circ$C for 10 h. Then, the nickel–carbon was dissolved into 250 mL ultrapure water and mixed with 5 mL prepared sodium borohydride aqueous solution (16.7 wt %) under ultrasonic stirring until there were no bubbles in the solution. The B-ECP600 was finally obtained after being washed with ultrapure water and dried at 60$^\circ$C for 10 h.

Carbon-supported Pt nanoparticle catalysts (Pt/B-ECP600 and Pt/ECP600) were prepared via the conventional microwave method. First, 4.08 g H$_2$PtCl$_6$•6H$_2$O/EG solution (1.96 wt %) was mixed with 70 mg corresponding carbon support materials (B-ECP600 and ECP600), 21 mL EG, and 14 mL ultrapure water that was ultrasonically stirred for 1 h, and then 1 g sodium hydroxide aqueous solution (10 wt %) was added into the mixture and ultrasonically stirred for 10 min. Next, the mixture was microwave-heated to 130$^\circ$C from the initial temperature of 13$^\circ$C, and then mixed with 0.3 mL hydrochloric acid and 5 mL EA. Pt nanoparticle catalysts supported by carbon were synthesized after being washed with ultrapure water and dried at 60$^\circ$C for 10 h. The total Pt loading was maintained at 30 wt % for catalysts.

3.3. Physical Characterization

The crystalline phase of the support materials and the catalysts was analyzed by recording their Power X-ray diffraction (PXRD) patterns on a Bruker D8 Advance diffractometer equipped with a Cu-Kα radiation source. The images of transmission electron microscopy (TEM) were taken on a Tecnai F20 at an accelerating voltage of 200 KV. The X-ray photoelectron spectroscopy (XPS, ULVAC-PHI, Chigasaki, Japan) data were recorded by a PHI 5000 VersaProbe with an Al Kα radiator at 5 kV. The nitrogen sorption isotherms of the B-ECP600 were collected at 77K by an ASAP 2020. Brunauer–Emmett–Teller (BET, Micromeritics, Norcross, GA, USA) was used for the specific surface area and porosity evaluations. A Fourier-transform infrared spectrometer (FTIR, NEXUS870, Nicolet, Madison, WI, USA) was used to identify the chemical structure functional groups contained in B-ECP600.

3.4. Electrochemical Characterization

All the electrochemical measurements were performed at a certain temperature (30$^\circ$C) with a three-electrode cell system provided by Gamry instruments. A glassy-carbon rotating disk electrode (RDE, Pine Research Instrumentation, diameter: 5 mm, area: 0.196 cm$^2$) was used as the working electrode, and a platinum wire was used as the counter electrode. A reversible hydrogen electrode (RHE) was used as the reference electrode in a 0.1 M HClO$_4$ solution. Catalyst ink was made by mixing 2 mg of catalyst with 1600 µL ethanol, 400 µL ultrapure, and 14.2 µL 5 wt % Nafion. A total of
10 µL of catalyst ink was dropped on the working electrode and dried under an incandescent lamp for 30 s. The RHE zero potential was estimated with the potential at which the current crossed zero by measuring the steady-state polarization curves of the hydrogen electrode reactions in 0.1 M HClO₄ saturated with H₂. A cyclic voltammetry (CV) measurement was performed in an N₂-saturated 0.1 M HClO₄ solution at a scan rate of 50 mV·s⁻¹ ranging from 0.05 to 1.05 V. The electrochemically active surface area (ECA) was calculated from the charge of hydrogen desorption between 0.05 and 0.4 V and corrected with the Pt capacity of 210 µC·cm⁻² Pt [19]. The ORR polarization was performed in an O₂-saturated 0.1 M HClO₄ solution at a sweep rate of 10 mV·s⁻¹ and a rotation speed of 1600 rpm at the temperature of 30 °C. The accelerated durability tests (ADTs) were conducted in a 0.1 M HClO₄ solution at a scan rate of 200 mV·s⁻¹, ranging from 0.6 to 1.0 V for 5000 cycles.

4. Conclusions

In this study, we successfully synthesized boron-doped ECP600 in a facile way and prepared catalysts with boron-doped ECP600 as the support. The structure of boron-doped ECP600 was investigated by TEM, XRD, BET specific surface area, FTIR, and XPS analyses. In addition, the structures and the electrochemical activities of Pt Pt/B-ECP600 and Pt/ECP600 were compared. Due to the boron-doping, Pt/B-ECP600 exhibits the better dispersion and the smaller size of Pt nanoparticles, and thus leads to much higher ECSA than Pt/ECP600. Additionally, the ORR activities of Pt/B-ECP600 were found to be remarkably higher than those of Pt-ECP600. The excellent ORR performance of Pt/B-ECP600 comes from the synergistic effect of the Pt nanoparticles and B-ECP600: on the one hand, boron oxides can promote the adsorption and dissociation of O₂ on Pt. On the other hand, part of oxygen atoms on Pt may transport to the nearby B atoms for further reduction and thus enhance the efficiency of ORR. Moreover, B-ECP600 shows outstanding long-term stability in acidic solution compared with ECP600. To sum up, B-ECP600 is highly promising as a supporting material for PEMFCs, owing to its simple preparation process, enhanced-electrocatalytic activity, and the stability of the acidic environment when using B-ECP600 as the support.

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