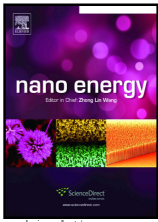
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www.elsevier.com/locate/nanoenergy

PII: S2211-2855(18)30252-0

DOI: https://doi.org/10.1016/j.nanoen.2018.04.021

NANOEN2651 Reference:

To appear in: Nano Energy

Received date: 6 March 2018 Revised date: 9 April 2018 Accepted date: 9 April 2018

Cite this article as: Jianbing Zhu, Meiling Xiao, Ping Song, Jing Fu, Zhao Jin, Liang Ma, Junjie Ge, Changpeng Liu, Zhongwei Chen and Wei Xing, Highly polarized carbon nano-architecture as robust metal-free catalyst for oxygen reduction in polymer electrolyte membrane fuel cells, Nano Energy, https://doi.org/10.1016/j.nanoen.2018.04.021

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Highly polarized carbon nano-architecture as robust metal-free catalyst for oxygen reduction in polymer electrolyte membrane fuel cells

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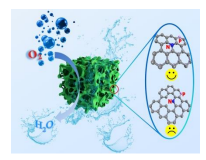
#### **ABSTRACT**

Metal-free electrocatalysts have eluded widespread adoption in polymer electrolyte membrane fuel cells due to their far inferior catalytic activity than most non-precious metal-N-C counterparts (M- $N_x$ -C) for oxygen reduction reaction (ORR), despite their distinct advantages over the M- $N_x$ -C catalysts, including lower cost and higher durability. Herein, we develop a rational bottom-up engineering strategy to improve the ORR performance of a metal-free catalyst by constructing a three-dimensional ultrathin  $N_x$ -P dual-doped carbon nanosheet. The resultant

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catalyst represents unprecedented ORR performance with an onset potential of 0.91 V, half-wave potential of 0.79 V. Impressively, a maximum power output at 579 mW cm<sup>-2</sup> is generated in the fuel cell test, the best among reported metal-free catalysts and outperforms most of the M-N<sub>x</sub>-C catalysts. The outstanding catalytic performance results from the highly active polarized carbon sites which are induced by selective graphitic nitrogen and phosphorous dual doping. Our findings provide new directions for the exploration of alternatives to Pt and bring a renew interests in the metal-free catalysts.

#### Graphical abstract



The ORR performance of metal-free electrocatalysts in acid media was successfully boosted by using electronic modulation and nanostructure engineering strategy simultaneously. The resultant NPC catalyst featured with 3D porous nanosheet structure represents unprecedented ORR performance with an onset potential of 0.91 V and half-wave potential of 0.79 V in acid medium, the best among reported metal-free catalysts.

Keywords: metal-free catalyst, active site, oxygen reduction reaction, polymer electrolyte membrane fuel cell

#### 1. Introduction

Polymer electrolyte membrane fuel cells (PEMFCs) are regarded as the most promising energy technology to replace the internal combustion engines as the vehicular power source [1, 2]. One major obstacle impedes their wide spread application is the high cost and limited reserve

of the state-of-the-art Pt-based catalysts. In this regard, great endeavors have been paid to search efficient non-precious metal catalysts (NPMCs) towards oxygen reduction reaction (ORR) [3-5]. Among the varied type of catalysts, the metal-free carbon-based materials exhibit distinct advantages over other NPMCs due to their lower cost, higher durability and simpler active site structure[6-8]. Since Gong et al firstly reported nitrogen-doped carbon nanotube arrays as ORR catalysts in Science [5], optimizing carbon nanostructure (1D nanotube [5, 9, 10], 2D nanosheet [11], 3D nanostructure[12, 13]) and modulating the doping elements (N, S, P, B, F or multiple atom doping [14-25]) have been majorly adopted to boost the catalytic performance. The regulation in carbon nanostructure addresses the accessible active site density, electronic conductivity, as well as the mass transfer properties of the catalysts [26-30]. Meanwhile, the doping elements modulate the intrinsic activity by altering the charge and spin density of carbon atoms [31-35].

Despite the enormous progress achieved, two obstacles remain for further development of the metal free catalysts: First, their catalytic performance in acid media for PEMFCs is still far away from satisfactory although a variety of heteroatoms-doped carbon materials were reported due to the low active site density and poor leached intrinsic activity. This has led to a judgement that transition metals (the M-N<sub>x</sub>-C type catalysts) are indispensable for the nonprecious catalysts in acidic solution, which greatly discourages the search for efficient acidic metal free ORR catalysts. Second, the understanding in the real active sites is also obscured due to the non-uniform structure, which hinders further rational design of high-performance metal-free electrocatalysts. Obviously, only when the intrinsic activity (doping functionalities), the activity unleashing issues (the tune in carbon structure) and the identification of active site are

simultaneously addressed can metal free catalysts become truly promising towards PEMFCs applications.

Herein, we aimed at improving the metal-free catalysts' performance through combining the electronic modulation with nanostructure engineering strategy, which is achieved by construing a series of N, P dual-doped carbon (NPC) materials featured with adjustable 3D porous nanosheet structure. Phosphorus is intentionally introduced to boost the intrinsic activity via electronic modulation effect, where its promotional effect is validated by a combination of ultraviolet photoelectron spectroscopy (UPS) analysis and density functional theory (DFT) calculation. The energy barrier of the  $O_{2(ads)}$  hydrogenation to form  $OOH_{(ads)}$  is reduced due to the significantly lowered local work function of active carbon atoms neighborhood the N, P dopants. Meanwhile, nanostructure engineering ensures the successful activity manifestation of these novel active sites, confirmed by enlarged current density. As a result, the N, P dual-doped carbon nanosheet catalyst presents unprecedented catalytic activity in acid media among the metal-free catalysts, i.e., an onset potential of 0.91 V, and half-wave potential of 0.79 V, outperforming most of the reported M-N<sub>x</sub>-C catalysts. The superb ORR performance of the derived catalyst is also confirmed by H<sub>2</sub>/O<sub>2</sub> single cell test. Our study not only paves new ways for the design of NPMCs, but also verifies the comparable contribution of metal-free active site structure to commonly recognized M-N<sub>x</sub> site, pushing the field one step ahead.

#### 2. Materials and methods

#### 2.1 Reagents

Melamine, diphenylphosphinic acid and zinc chloride were purchased from Aladdin Company. 5 wt% Nafion ionomer was obtained from Aldrich. Perchloric acid was purchased

from Alfa Aesar. Commercial state-of-the-art 20 wt% Pt/C (Johnson Matthey Company, HiSPEC<sup>TM</sup> 3000) was used as the benchmark for comparison and was denoted as Pt/C. Ultrapure water (Millipore,  $18.2 \text{ M}\Omega$  cm) was used throughout all experiments.

## 2.2 Preparation of materials

The melamine–diphenylphosphinic acid complex crystals (MDPCC) with adjustable N/P ratios were synthesized by a wet chemistry method. For a typical synthesis of MDPCC with N/P ratio of 4, melamine (2.52 g) and ultrapure water (100 mL) were added into a 250 mL single-necked, round-bottomed flask. Then flask was transferred into an oil bath and heated at 100 °C under stirring. After the solution became clear diphenylphosphinic acid (1.09 g) was added into the solution and the mixture was allowed to react at 100 °C for 3 h. Finally the solution was evaporated to dryness and obtained the precursor.

The N, P co-doped catalyst was synthesized by pyrolyzing melamine–diphenylphosphinic acid complex crystals at the target temperature. Typically, 2.0 g MDPCC was placed in a quartz boat and then pryolyze at 1100 °C under N<sub>2</sub> atmosphere for 1 h, with a heating rate of 2 °C/min. The obtained material was denotes as NPC-X-1100. For the NPC-4-1100-Zn sample, 2.0 g MDPCC precursor mixed with 500 mg ZnCl<sub>2</sub>, then pyrolyzing with the same procedure described above.

#### 2.3 Physical Characterzation

Scanning electron microscopy (SEM) measurements were performed with an XL 30 ESEM FEG field emission scanning electron microscope. Transmission electron microscopy (TEM), high resolution transmission electron microscopy (HRTEM), high-annular dark-field scanning transmission electron microscopy (STEM) and element mapping analysis were conducted on

Philips TECNAI G2 electron microscope operating at 200 kV. X-ray diffraction (XRD) measurements were performed with a PW-1700 diffractometer using a Cu  $K_{\alpha}$  ( $\lambda$ =1.5405 Å) radiation source (Philips Co.).Raman spectra were collected on a J-Y T64000 Raman spectrometer with 514.5 nm wavelength incident laser light. The textural and morphological features of the various carbon supports and catalysts prepared were determined by nitrogen physisorption at 77 K in a Micromeritics ASAP 2020. Textural properties such as the specific surface area pore volume and pore size distribution were calculated from each corresponding nitrogen adsorption–desorption isotherm, applying the Brunauer–Emmett–Teller (BET) equation and the Barrett–Joyner–Halenda (BJH). X-ray photoelectron spectroscopy (XPS) measurements were carried out on Mg  $K_{\alpha}$  radiation source (Kratos XSAM-800 spectrometer). The bulk compositions were evaluated by inductively coupled plasma optical emission spectrometer (X Series 2, Thermo Scientific USA).

### 2.4 Electrochemical measurement

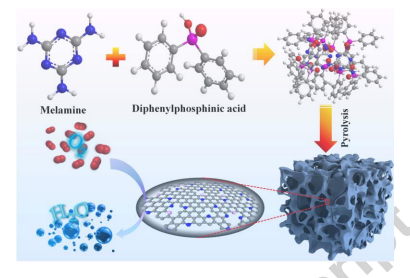
All electrochemical measurements were conducted in a conventional three-electrode cell at room temperature (~25 °C) using the 750E Bipotentiostat (CH Instruments). Non-noble metal catalyst ink was prepared by ultrasonically dispersing 5 mg catalyst in a suspension containing 50 μL Nafion (5 wt %) solution and 950 μL ethanol, while Pt/C catalyst (20 wt% Pt on Vulcan XC-72 carbon, Johnson Matthey) ink was prepared by dispersing 5 mg catalyst ultrasonically in a mixture solution containing 50 μl Nafion (5 wt %) solution, 550 μL isopropanol and 400 μL Milli-Q water. The catalyst film coated electrode was obtained by dispersing the catalyst ink on a glassy carbon rotating ring-disk electrode followed by drying in air. The catalyst loadings on RRDE were 0.4 mg cm<sup>-2</sup> for NPC-X-*T* catalysts and 40 μg<sub>Pt</sub> cm<sup>-2</sup> for Pt/C catalyst. The ORR stability was investigated by continuous potential cycling in oxygen-saturated 0.1 M HClO<sub>4</sub>

solution between 0.6 V and 1.05 V with the scan rate at 0.1 V s<sup>-1</sup>. And after 10, 000 cycles, the ORR steady-state polarization measurements were conducted in  $O_2$ -saturated 0.1 M HClO<sub>4</sub> solution with scanning rates of 5 mV s<sup>-1</sup> and rotation rate at 1600 rpm.

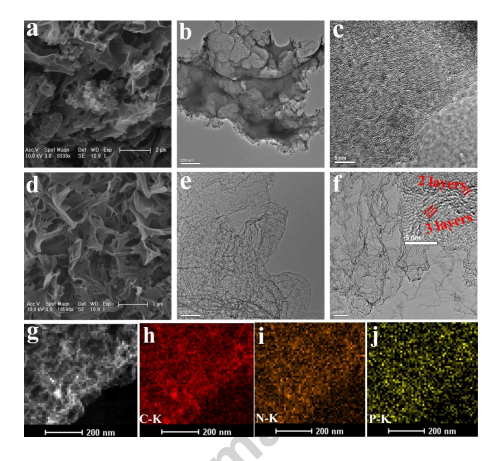
#### 3. Results and Discussion

The porous metal-free NPC catalysts were synthesized via a template-free process as illustrated in **Scheme 1**. The melamine and diphenylphosphinicacid were firstly allowed to cooperatively assemble into melamine–diphenylphosphinicacid complex crystals (MDPCC), confirmed by the Fourier Transform Infrared Spectroscopy (FTIR) together with <sup>31</sup>PNMR spectra (Fig. S1-3). Subsequently, the MDPCC underwent a one-step pyrolysis to yield the final NPC catalysts. The precursor was pyrolyzed under various temperatures (800-1100 °C) and catalysts with different melamine/diphenylphosphinicacid feeding ratios (N/P ratios) were synthesized, with resultant samples labeled as NPC-X-*T* (X represents the N/P ratios in the precursor, *T* represents pyrolytic temperature). Furthermore, the microstructures of these NPC-X-*T* catalysts were further optimized by co-pyrolyzing the MDPCC precursor in the presence of ZnCl<sub>2</sub> (NPC-4-1100-Zn).

**Scheme 1** Schematic illustration of the synthesis of NPC-X-*T* electrocatalysts.



The morphology and structure of the as-synthesized materials were examined by scanning electron microscopy (SEM) and transmission electron microscopy (TEM) images. First, increasing the pyrolysis temperature results in structure evolution from solid block morphology (NPC-4-800) into fluffy three-dimensional nanosheet structure (**Fig. 1** and Fig. S4), with NPC-4-1100 achieving the most uniform and porous structure. N and P were both successfully doped into carbon skeletons throughout the whole carbon matrix, as shown in Fig. S5. Besides, higher pyrolysis temperature leads to higher graphitization degree, as evidenced by clearer lattice fringes observed from the HRTEM and decreased  $I_d/I_g$  value in Raman spectra (Fig. S6). Second, we investigated the influence of N/P ratio in the precursor on the microstructure of the pyrolytic materials. As shown in Fig. S7, increasing the melamine ratio results in structure transformation from solid block aggregates (NPC-1-1100 and NPC-2-1100) into 3D porous nanosheet structure (NPC-4-1100 and NPC-6-1100), caused by the massive gas generated from melamine decomposition into carbon nitride gases (e.g.  $C_2N_2^+$ ,  $C_3N_2^+$ ,  $C_3N_3^+$ ). Third, the structure of the NPC-4-1100 catalysts (the one with most uniform pore structure) can be further

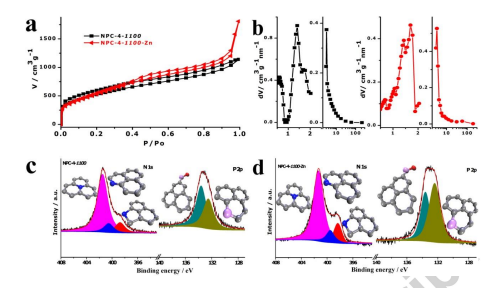


**Figure 1.**(a) SEM, (b) TEM, (c) HRTEM images of NPC-4-*1100*; (d) SEM, (e) TEM and (f) HRTEM images and (g-j)corresponding the elements maps of NPC-4-*1100*-Zn, (h) C-K, (i) N-K, (j) P-K).

tuned by adding ZnCl<sub>2</sub> into the precursor before pyrolysis. The resultant NPC-4-1100-Zn catalyst represents a three-dimensional macroporous structure, where the walls (Fig. 1d-j) exhibit well defined graphitic structure consisting of 2~4 graphene layers, much thinner than the NPC-4-1100 catalyst (inset Fig. 1f). The highly graphitized feature and the porous structure guarantee the final NPC-4-1100-Zn with good electronic conductivity, active site accessibility, and fluent mass transfer.

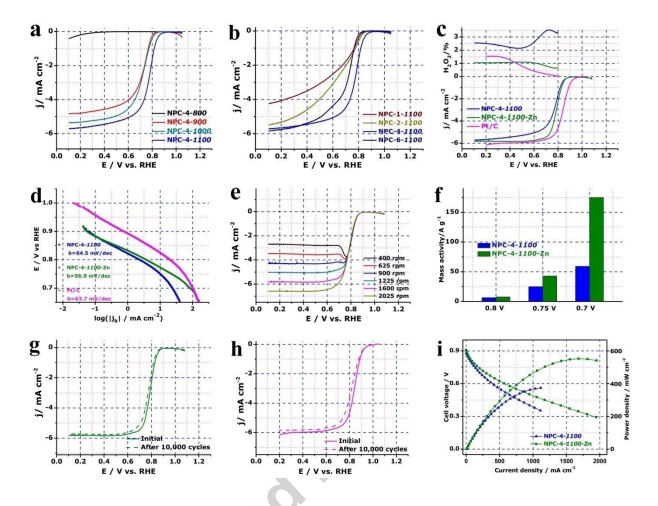
We then investigated the surface area and pore structure of the synthesized catalysts through Brunauer-Emmett-Teller (BET) technique. The  $N_2$  adsorption-desorption results (**Fig. 2a** and

Fig. S8, S9) demonstrate type IV isotherm for all the catalysts. Combining with the pore size distribution analysis, the catalysts are confirmed to exhibit hierarchical porous structure; where micropores (< 2 nm) and mesopores are simultaneously present (Fig. 2b and Fig. S8, S9). For all the NPC-X-T catalysts, higher pyrolytic temperature and larger N/P ratio leads to enlargement in surface areas, with the NPC-4-1100 catalyst exhibits the highest surface area of 1698 m<sup>2</sup> g<sup>-1</sup> and largest pore volume of 1.92 cm<sup>3</sup> g<sup>-1</sup> (Fig. S10 and Table S1). The introduction of Zn (NPC-4-1100-Zn) leads to further enhanced surface area (2048 m<sup>2</sup> g<sup>-1</sup>) and pore volume (2.79 cm<sup>3</sup> g<sup>-1</sup>, Fig. S10), which exceeds that of NPC-4-1100 and most of the reported heteroatom-doped carbon materials [36-39]. The X-ray photoelectron spectroscopy (XPS) was adopted to characterize the surface structure of the catalysts. The deconvoluted high resolution N 1s spectra (Fig. 2c, d and Fig. S11, S12) show the concurrent presence of pyridinic N (398.5 eV), graphitic N (401.3 eV), and pyrrolic N (399.5 eV). The P 2p spectra were deconvoluted into two peaks at 132.9 eV (P-C) and 133.8 eV (P-O), confirming the successful doping of P and the formation of P conjugated aromatic C rings (Fig. 2c, d and Fig. S11, S12). No signals at around 402.2 eV for N-P bonding can be found, indicating the isolated N and P species in the carbon network. The surface N and P doping concentrations are found to decrease with increase in pyrolysis temperature (Fig. S13 and Table S2), and the respective concentrations of N and P are also well tuned by varying N/P feeding ratios (Fig. S13 and Table S2). Of all the NPC-X-T catalysts, the NPC-4-1100 represents the most balanced N (5.90 at%) and P doping levels (2.14 at%), which is considered to be beneficial for ORR, provided N and P both promote the catalytic process. It is noted that while the final NPC-4-1100-Zn represents comparable P and N doping levels with NPC-4-1100, the P 2p spectra shows a higher proportion of P-C species than P-O, suggesting the promotional effect of Zn in the formation of P conjugated aromatic C rings.



**Figure 2** (a) N<sub>2</sub> adsorption–desorption isotherms and (b) the corresponding pore size distribution curves for NPC-4-*1100* and NPC-4-*1100*-Zn catalysts; High-resolution N 1s and P 2p XPS spectra for (c) NPC-4-*1100*, (d) NPC-4-*1100*-Zn catalysts.

The electrocatalytic performance of the catalysts was evaluated using rotating ring-disk electrode (RRDE) technique in O<sub>2</sub>-saturated 0.1 M HClO<sub>4</sub> solution and compared with the commercial Pt/C catalysts (Johnson Matthey, 20% Pt). A clear impact of synthetic condition on both onset potential and limiting current is observed, as illustrated in **Fig. 3**. Specifically, the NPC-4-900, NPC-4-1000, and NPC-4-1100 represent onset potentials of 0.82, 0.85, and 0.91 V, in sequence, while the NPC-4-800 shows almost no ORR activity with only a small current at ultra-high over potential (~0.5 mA cm<sup>-2</sup> at 0.1 V vs. RHE, Fig. 3a). The main reason for the ultra-low ORR activity on NPC-4-800 might be due to the insufficient electronic conductivities of NPC-4-800 (because of the low graphitization degree), as revealed by the quasi-reversible [Fe(CN)<sub>6</sub>]<sup>3-/4</sup>-redox test (Fig. S14). Meanwhile, the ORR 4-electron pathway selectivity enhanced with increasing pyrolytic temperature, as evidenced by the decreased H<sub>2</sub>O<sub>2</sub> yield from Figure S15, which corresponds to the electron transfer number of 3.76, 3.81 and 3.95 for NPC-4-900, NPC-4-1000, and NPC-4-1100 (Fig. S15b, S16), respectively. As for the catalysts with different



**Figure 3** Steady - state ORR polarization curves for NPC-4-*T* (a) and NPC-X-1100 (b) catalysts in O<sub>2</sub>-saturated 0.1 M HClO<sub>4</sub> with a scan rate of 5 mV/s and a rotation speed of 1600 rpm; (c) RDE polarization curves and peroxide yield of NPC-4-1100, NPC-4-1100-Zn and commercial Pt/C in O<sub>2</sub>-saturated 0.1 M HClO<sub>4</sub> with a scan rate of 5 mV/s and a rotation speed of 1600 rpm; (d) Tafel plots of NPC-4-1100, NPC-4-1100-Zn and Pt/C catalysts; (e) Koutecky-Levich (K-L) plots for NPC-4-1100-Zn at various potentials; (f) mass activity of ORR at different potentials on NPC-4-1100 and NPC-4-1100-Zn samples; (g, h) steady - state ORR polarization curves of NPC-4-1100-Zn (g) and Pt/C (h) before and after 10,000 potential cycles in O<sub>2</sub>-saturated 0.1 M HClO<sub>4</sub>; (i) polarization curves and corresponding power densities of membrane electrode assemblies fabricated with NPC-4-1100 and NPC-4-1100-Zn cathode catalysts.

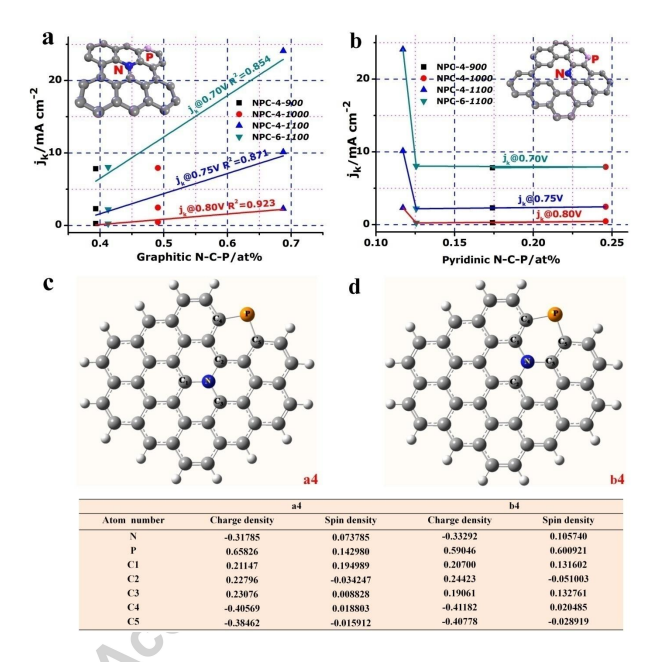
N/P ratios, the NPC-4-1100 catalyst represents the best performance with half-wave potential locates at 0.775 V as well as excellent stability (Fig S17). In comparison, while NPC-1-1100 and NPC-2-1100 catalysts are unsuccessful in reaching the diffusion limited current density plateau, the NPC-6-1100 catalyst exhibits a 75 mV negative shift in half-wave potential (E<sub>1/2</sub>=0.70 V, Fig. 3b). The RRDE measurements further confirm the lowered charge transfer number of NPC-1-1100 and NPC-2-1100 catalysts and the four-electron ORR process(n>3.85) on the NPC-6-1100 catalyst (Figure S15c, d and Fig. S18). It is noted that although quite different behaviors were observed in the mixed kinetic-diffusion control region and diffusion limited current density plateau, almost all the NPC-X-1100 catalysts showed similar onset potential, indicating the NPC-X-1100 catalysts shared the same active site at different active site densities. Additionally, NPC-4-1100 catalyst exhibited significantly enhanced ORR activity compared with the N-doped C and P-doped C catalysts (Fig. S19), further verified the synergetic effect between N and P dopants.

Of particular interest is the NPC-4-1100-Zn catalyst, which performs even better than NPC-4-1100, with E<sub>1/2</sub> at 0.79 V (15 mV positive shift) and charge transfer number at 3.95 (~1% H<sub>2</sub>O<sub>2</sub> in all potential region) obtained (Fig. 3c-e and Fig. S20). This result not only corresponds to the highest ORR performance ever reported for metal free catalysts, but also exceeds most of the M-N<sub>x</sub>-C catalysts (Table S3) and only shows a 50 mv negative shift in E<sub>1/2</sub> compared with the state-of-the-art Pt/C catalysts (E<sub>1/2</sub>=0.845 V). The analysis of Tafel slopes for NPC-4-1100 (64.5 mV/dec), NPC-4-1100-Zn (56.9 mV/dec) and Pt/C (63.7 mV/dec) catalysts (Figure 3d) revealed the similar ORR kinetic process on these electrodes, where the transfer of the first electron from the electrode to O<sub>2</sub> is the rate limiting step. The ORR stability of NPC-4-1100-Zn catalyst was also evaluated using an accelerated durability test protocol by cycling the catalysts between 0.6 and 1.0 V at 100 mV s<sup>-1</sup> in O<sub>2</sub>-saturated electrolyte. As shown in Figure 3g, after 10,000 cycles,

the NPC-4-1100-Zn catalyst showed 16 mV negative shift in half-wave potential, comparable to that of Pt/C catalyst (15 mV, Figure 3h) and nearly unchanged Tafel slop (Fig. S21). To further validate the excellent ORR performance of NPC-4-1100-Zn catalyst, single cell test was conducted. As shown in Figure 3i, the open circuit voltage of the single cell is 0.905 V, which is consistent with results from the RRDE measurements, and the maximum power output was 579 mW cm<sup>-2</sup> for the NPC-4-1100-Zn much better than that of NPC-4-1100 catalyst (393 mW cm<sup>-2</sup>). We further conducted durability test at a constant cell voltage of 0.5 V (Fig. S22). As can be seen, it delivered a decent stability performance by showing only ca. 21.5% loss of the initial current over 72 h of operation, demonstrating excellent durability of NPC-4-1100-Zn cathode electrocatalyst. This result suggests the highly promising future of the metal free catalyst to be utilized in real PEMFC system.

Based on the fact that metal-free N doped carbon-based ORR catalysts exhibiting onset potential over 0.8 V in acidic solution has never been reported[40-42], we have reason to believe that the NPC catalysts with onset potential exceeding 0.9 V must originated from the creation of intrinsically more active sites, due to the N and P co-doping. To reveal the synergistic functions of N and P in ORR, we further studied the relationship between the current densities and the different dopants' concentration, where the graphitic N-C-P (G<sub>N</sub>-C-P) and pyridinic N-C-P (P<sub>N</sub>-C-P) C-P) species (configuration shown inset Fig. 4a, b) were majorly investigated. As shown in Fig. 4a, we found an obvious linear correlation between G<sub>N</sub>-C-P concentration and ORR activity at high potential (0.8 V), where the correlation parameter R<sup>2</sup> is calculated to be 0.923, clearly evidencing the synergistic effect between graphitic carbon and C-P bond. At lower potentials, as other active sites (such as pyridinic N-C species) became active and contribute partially to the  $(R^2=0.871@0.75V)$ kinetic decreased dependences observed current, linear are

R<sup>2</sup>=0.854@0.70V), which are however still sufficient to demonstrate the synergy effect. The critical role of G<sub>N</sub>-C-P was further confirmed by excluding current contribution of pyridinic N-C active sites according to Nakamura's work (details are shown in supporting information), which results in increased correlation coefficient to 0.886 at 0.75 V and 0.875 at 0.70 V, respectively. The specific activity per G<sub>N</sub>-C-P specie is estimated to be 0.18 e<sup>-</sup>s<sup>-1</sup> at 0.8 V, much higher than that of the pyridinic N-C active site at 0.5 V (0.13 e<sup>-</sup>s<sup>-1</sup>) [29], verifying the dominated role of G<sub>N</sub>-C-P



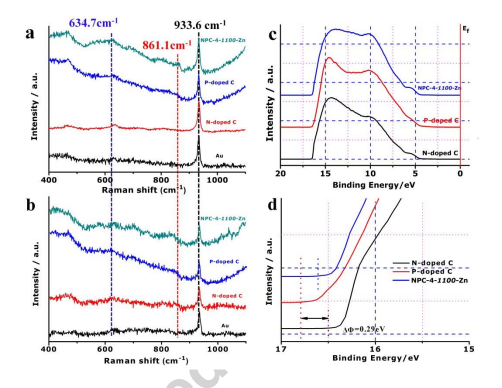
**Figure 4** (a) correlation between kinetic current densities of ORR at 0.8, 0.75, and 0.7 V versus RHE and the graphitic N-C-P species concentrations; (b) Correlation between kinetic current densities and the pyridinic N-C-P concentrations; (c, d) spin and charge density of carbon network (gray) dual-doped by N (blue) and P (orange). C1 has very high spin density, C2 and C3 have high positive charge density, and C4 and C5 have moderately high positive spin densities.

active sites. In contrast, no distinguishable correlation between the kinetic current density and  $P_N$ -C-P contents is noticed (Fig. 4b), indicating that the ORR activity is determined solely by the  $G_N$ -C-P concentration at high potential.

Density functional theory (DFT) calculations were performed to further reveal the real active site structure. It is known that the ORR activity of N doped carbon is rooted from the charge transfer from C to N due to the higher electron negativity of the latter, where the C atoms with positive charges results in increased ORR activity. For the NPC catalysts in the G<sub>N</sub>-C-P model, we found that the charge and/or spin density of neighboring C atoms further increases when P atoms are bonded to carbon atoms adjacent to graphitic N. On the contrary, for the P<sub>N</sub>-C-P type, the charge and spin density of the carbon atoms decrease with the introduction of neighboring P dopants, compared with the pristine N-doped carbon (Fig. 4c, d and Fig. S19-22 and Table S4-S7). G<sub>N</sub>-C-P species are believed as the origination for the much higher intrinsic activity, consistent with the results in Fig. 4a. The possible configurations of active site structures were then confirmed by comparing the charge/spin density of various structure models. As shown in Fig. 4c, d, the optimized models were a4 and b4 due to the highest charge/spin density achieved. Specifically, the spin density was up shifted to 0.195 for C1 in Model a4 and 0.133 for C3 in Model b4 (Fig. 4c, d and Table S4, S5), which is higher than that of N-doped C sample (0.178 for C1 in a4 and -0.008 for C3 in b4).

To confirm the more positive charge characteristics of the active C atoms, we further employed the surface-enhanced resonance Raman spectroscopy (SERRS) and used SCN<sup>-</sup> to probe the alternation in surface resonance signals after adsorption. The peaks corresponding to P–C/P-C--N (620-700 cm<sup>-1</sup>) bond stretching modes and characteristic peaks of N-C/N-C—P species (770 cm<sup>-1</sup> to 880 cm<sup>-1</sup>) show significantly attenuation in peak density after adsorption,

evidencing the positively charged nature of C atoms (Fig. 5a). Notably, the NPC-4-1100-Zn sample exhibits a more attenuation compared to N-doped C, suggesting more positive charge of C atoms due to N, P co-doping (Fig. 5b), being consistent with the DFT calculation results.



**Figure 5** (a, b) SERRS spectra of the N-doped C, P-doped C and NPC-4-1100 catalysts before (a) and after (b) injecting KSCN in 0.1 M HClO<sub>4</sub>; (c) UPS spectra collected using an He I (21.2 eV) radiation; (d) the enlarged view of the secondary electron tail threshold.

Last, ultraviolet photoelectron spectroscopy (UPS) measurements were carried out to investigate the doping effect of N and P on the work function of the catalysts. In Fig. 5c, two bands could be clearly distinguished, assignable to  $\sigma$  and  $\pi$  valence-band of graphite. The work functions determined from He I spectra is 4.70, 4.41, and 4.59 eV for the N-doped C, P-doped C, and NPC-4-1100-Zn, respectively (Fig. 5d). Compared to the N-doped C, the lowered local function of the NPC-4-1100-Zn catalyst suggests a lower energetic barrier (higher driving energy) for

donating electrons from the surface of the N, P co-doped catalyst to the adsorbed molecular oxygen, thereby facilitating the formation of the OOH species, which is known to be the rate-determining step in the ORR process.

#### 4. Conclusion

In summary, we addressed the intrinsic activity and the active sites unleashing issues of metal-free ORR catalysts using electronic modulation and nanostructure engineering strategy. Specifically, an onset potential of 0.91 V and a half-wave potential of 0.79 V were achieved in 0.1 M HClO<sub>4</sub>, ranking the first among the metal-free carbon-based electrocatalysts and is comparable to the M-N<sub>x</sub>-C catalysts. Furthermore, in-situ fuel cell test demonstrates maximum power output at 579 mW cm<sup>-2</sup>, the best for metal-free catalysts and even better than most M-N<sub>x</sub>-C materials. This result provides a sound refute to the judgement that only metal containing materials are viable nonprecious catalysts in acidic solution. In-depth study on the active site structure revealed that the positive charged carbon atoms adjacent to both graphitic N and P species are the main active sites. The enhanced charge/spin density and lower local work function due to co-doping were further revealed by DFT calculations and UPS measurements, respectively. Our findings provide new directions for the design of earth-abundant materials as highly active ORR electrocatalysts.

#### Acknowledgments

The work is supported by the National Natural Science Foundation of China (21433003, 21633008), the National Key Research and Development Program of China (2016YFB0101202), Jilin Province Science and Technology Development Program (20150101066JC,

20160622037JC, 20170203003SF), Hundred Talents Program of Chinese Academy of Sciences and the Recruitment Program of Foreign Experts (WQ20122200077).

# Appendix A. Supporting information

SEM, TEM, XRD, XPS, Raman and other electrochemical measurements are available in Supporting Information.

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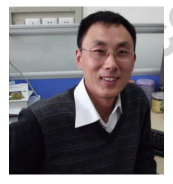


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### Highlights:

- The synthesized catalyst exhibits the best ORR performance among metal-free catalysts in acidic medium.
- Graphitic N and P polarize the adjacent carbon atoms, making them catalytically active to ORR.
- Optimized 3D porous structure guarantees activity manifestation of these active sites.