One-step synthesis of Fe$_3$PtPd(OH)$_2$[Picolinic acid]$_8$(H$_2$O)$_4$ hybrid nanorods: efficient and stable electrocatalyst for oxygen reduction reaction in alkaline solution

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Design and synthesis of effective electrocatalysts for oxygen reduction reaction in alkaline environments is critical to reduce energy losses in alkaline fuel cells. We have systematically evaluated new approaches for reducing the Pt content while retaining the activity of a Pt-based catalyst with hydrolytic phases containing hydroxide moieties in addition to metal ions and ligands. We report for the first time architectured organic-inorganic hybrid nanorod catalyst, which is fabricated by solvothermal reaction of K$_2$MCl$_4$ (M = Pd, Pt) with picolinic acid (PA) (chelating agent) in the presence of FeCl$_2$. Excess base produces isostructural coordination M-PA complex to Fe-OH chains. A generic formula can be written as Fe$_3$PtPd(OH)$_2$[PA]$_8$(H$_2$O)$_4$. The electrocatalytic activities of the hybrid nanorods are explored for oxygen reduction reaction (ORR) in alkaline medium. The onset potential of ORR is significantly reduced with a positive shift of about 109 mV and twice the reduction current density is observed in comparison with Pt/C with the same mass loading. We believe that this work may lead towards the development of heterodoped organic-inorganic hybrid materials with greatly enhanced activity and durability for applications in catalysis and energy conversion.

Energy is one of the biggest challenges in the 21st century and there has been an ever increasing demand for environmental friendly high-power energy sources. Fuel cells with their high energy and power density have been widely considered as green and efficient alternative energy sources$^{1-2}$. The cathodic oxygen reduction reaction (ORR) is at the heart of fuel cell performance, and efficient ORR electrocatalysts are essential for practical applications. The common commercial catalysts used in ORR are Pt and Pt based precious metal electrocatalysts, which exhibit excellent catalytic activity, but suffer from prohibitive cost, susceptibility to methanol crossover, and poor stability in the electrochemical environments$^{3-7}$. Recent efforts in electrocatalysis have focused on decreasing the Pt content in fuel cell electrocatalysts or replacing it with less expensive materials$^{8-12}$. In the face of these considerable challenges, one of the pathways to address these problems is to increase the ORR activity as well as reduce cost by alloying a second metallic element such as Pd, Fe, Ni, and Co with pure Pt$^{12-16}$. These catalysts are more active for ORR than their Pt counterparts. For example, recent research has demonstrated that Pd doped Pt systems show excellent ORR performance in both alkaline and acidic electrolytes$^{17,18}$. The cost of Pd, however, is currently about one-third that of Pt, and it is at least 50 times more abundant than Pt. In addition, for ORR, it has been found that a small amount of Fe doping is effective for enhancing the ORR activities of Pt or Pd alloys, possibly due to the enhanced structure disorder and conductivity$^{19,20}$. For example, the unique core–shell structures of Pd/FePt NPs with 5 nm Pd core encircled with a FePt shell generate 12 times more current (5 nm/1 nm Pd/FePt

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NPs) than commercially available pure-Pt catalysts. On the other hand, transition-metal species, such as metal hydroxides or layered double hydroxides, have recently gained noticeable popularity in various energy systems owing to their low cost and high theoretical activity. For example, Lei et al. report on the highly active and durable Ni₅Co₁₋ₓ(OH)₂ catalyst for ORR. Li et al. observed an excellent ORR performance on NiCoFe-LDH. These results demonstrate that 3D metal in the hydroxide state is a promising catalyst for ORR. These studies give the idea that the use of noble and non-noble metals together in the hydroxide or complex form, instead of the reduced form, can effectively increase ORR activity. But usually, heterogeneous electrocatalysts suffer from extensive leaching of the active metal species during reactions and eventually lose their catalytic activity.

Herein, for the first time, we report an architectured organic-inorganic hybrid nanorod electrocatalyst, which was fabricated by the solvothermal reaction of K₂MCl₄ (M = Pd, Pt) with picolinic acid in the presence of FeCl₂ (in which hydroxide forms easily in the basic solution). The excess base produces isostructural coordination solids in which ‘complex ligands’, containing palladium or platinum, coordinate to metal hydroxide chains. A generic formula can be written as: Fe₃M₂(OH)₂[PA]₈(H₂O)₄, where M²⁺ = Pd and Pt. PA is known to be stable in solvothermal conditions and also to form stable complexes with soft metals via -NH coordination and hard metals –OH coordination. This synthesis method allows to form a catalyst with a uniform distribution of metal ions at structure. In these conditions, the Pt content of the catalyst decreases while the activity of a Pt-based catalyst with hydrolytic phases containing hydroxide moieties will be retained. Also, this structure decreases metal leaching in successive runs and increases reusability of the catalyst. To study the effect of the presence of Pt and Pd on ORR activity, three types of hybrid nanorods, including Fe₃Pt₂[PA]₈(OH)₂(H₂O)₄, Fe₃Pd₂[PA]₈(OH)₂(H₂O)₄ and Fe₃PtPd[PA]₈(OH)₂(H₂O)₄, were synthesized, and the effect of their compositions on ORR activity was studied. These hybrid nanorods displayed substantially enhanced ORR activity as compared with that of commercial Pt/C catalysts in 0.1 M KOH solution. It is critical to highlight that, to the best of our knowledge, there have been no previous reports of either supported or activated Fe(OH)₂ with Pt and Pd complexes. These hybrid nanorods are a promising new catalyst candidate for practical fuel cell applications.

**Results**

**Characterization of hybrid nanorods.** FT-IR spectroscopy, X-ray diffraction (XRD), scanning electron microscopy (SEM), EDX, and transmission electron microscopy (TEM) were applied for characterization. FTIR spectra for the as-prepared nanohybrids are shown in Fig. 1a. The FTIR spectra of nanohybrids were roughly attributed to the following: 3412 cm⁻¹ to the O-H and N-H stretching vibration of the Fe hydroxide layer and PA amine groups; 3056 cm⁻¹ due to the alkyl C-H stretching of PA; 1687 cm⁻¹ due to the carboxylic acid C=O stretching of PA and the OH stretching band of H₂O; 1506–1591 cm⁻¹ due to the aromatic C=C bending; 1327 cm⁻¹ due to the ν(CO₂) asymmetric; and 1276 cm⁻¹ due to the ν(CO₂) symmetric; 1154 cm⁻¹ due to the –C=N stretching; and 500–900 cm⁻¹ to the Fe-O, Fe-O-Fe, and O-Fe-O lattice vibrations. These observations confirmed the formation of Pt/Pd-PA complex and Fe-hydroxide at all three hybrid nanorods.

Figure 1b illustrates the XRD pattern of hydrothermally synthesized hybrid nanorods. From Fig. 1b, it can be seen that the observed reflections clearly indicate the formation of a single phase compound without any impurity traces. XRD results revealed that all of three formed hybrid nanorods have a monoclinic crystal structure with a C2/c space group (JCPDS card no. 00-049-2426). The structure of monoclinic nanorod hybrids is composed of two different structures including metal hydroxide (C2/c space group and JCPDS card no. 00-030-0147) and M-PA complex with monoclinic structure (P21/c space group and JCPDS card no. 00-044-1812), which are held...
together by two different types of linkage. The first type of linkage consists of infinite chains with a –Fe/PA/M/PA/Fe/PA/M– repeat unit. The second type is a series of Fe hydroxide strips. Each individual layer has metal hydroxide ‘ribs’, which are linked to their nearest neighbors by a row of PA/M/PA moieties.

The surface composition of the Fe₃PtPd(OH)₂[PA]₈(H₂O)₄ was analyzed by X-ray photoelectron spectroscopy (XPS, Figure S1). The survey spectrum (Figure S1a) shows carbon, oxygen, nitrogen, iron, palladium and platinum species. The high-resolution XPS spectra of C 1 s (Figure S1b) showed that the C 1 s region contains five components corresponding to C=O, C–OH, C=O, HO-C=O and C=N species can be further separated out. The O 1 s region (Figure S1c) contains four components corresponding to Pt/Pd/Fe-OH, C=O, C-O, and O-C=O bonds. The N 1 s, Pd 3d and Pt 4 f XPS spectra are shown in Fig. 2d–f, respectively. The increase in the binding energy of element N and decrease in the binding energies of element Pd and Pt was occurred due to the fitting the surplus pair electrons of element N in the free electron orbital of element Pd and Pt in the chelating process. The positive shifts of the N 1 s peak and negative shifts of the Pd 3d and Pt 4f peaks suggest that the intermolecular coordination of palladium/platinum with nitrogen of the picolinic acid happened. The shift of binding energy could affect the interaction strength between Pd (II)/Pt (II) ions and picolinic acid. In Figure S1g, the XPS spectrum of the Fe shows double peaks with binding energies at 706.3 eV and 719.8 eV, corresponding to Fe 2p₃/2.

Figure 2. Structural and compositional analyses of the hybrid nanorods. (a) Low- and high- magnification FE-SEM images of the hybrid nanorods, (b) TEM images of the hybrid nanorods, (c) EDS spectrum of the hybrid nanorods.
and Fe 2p1/2, respectively. Compared to the standard spectra of metal Fe(II), the binding energies of Fe decreases slightly. These results combined with obtained results from O 1 s spectra confirm the Fe(OH)2 formation. XPS was used to confirm the valence state of the Pd, Pt and Fe on the catalyst. As shown in Fig. 2e–g, the Pd 3d, Pt 4f and Fe 2p XPS spectra of catalyst are attributed to Pd(II), Pt(II) and Fe(II).

Morphological properties of all three hybrid nanorods were analyzed by FE-SEM and TEM, and the images are shown in Fig. 2. From FE-SEM and TEM images, it is apparent to notice the formation of the regular shaped hybrid nanorods with ~100 nm thickness and 2–3 µm length for all three hybrid nanorods (Fig. 2a,b). The elemental analysis of the as-prepared hybrid nanorods was obtained using EDX and ICP-AES. According to Fig. 2c, the formulas of the hybrid nanorods are derived as Fe3Pt2[PA]8(OH)2(H2O)4, Fe3Pd2[PA]8(OH)2(H2O)4, and Fe3PtPd[PA]8(OH)2(H2O)4, which is strong evidence for the formation of the hybrid nanorods. SEM-EDX mapping analysis for the hybrid nanorods (Figure S2) proved the uniform distribution of Pt, Pd, Fe and C, N, O (Picolinic acid) inside structure. The uniform structure lead to enhancement in both catalytic activity and durability toward the oxygen reduction reaction.

**Activity and performance of hybrid nanorods for the ORR.** We first compared the electrochemical behavior of the hybrid nanorods with each other and with Pt/C for ORR using cyclic voltammograms (CVs) cell in oxygen-saturated 0.1 M KOH (Fig. 3a). The detailed information can be found in Table 1. As can be seen in Fig. 3a, distinct peaks corresponding to ORRs can be observed for all the hybrid nanorods. In Fig. 3a, single cathodic reduction peaks at −0.3 V, −0.18 V, and −0.09 V can be observed in an O2-saturated solution for the Fe3Pt2[PA]8(OH)2(H2O)4, Fe3Pd2[PA]8(OH)2(H2O)4, and Fe3PtPd[PA]8(OH)2(H2O)4 electrodes, respectively. Most interesting is that in the O2-saturated solution, ORR peak potential of the Pd based hybrid nanorods is more positive than that of Pt based hybrid nanorods. Additionally, compared with Fe3Pt2[PA]8(OH)2(H2O)4, Fe3Pd2[PA]8(OH)2(H2O)4, a significant increase of the peak current density and an obvious positive shift of the peak potential can be found on Fe3PtPd[PA]8(OH)2(H2O)4. Commercial Pt/C catalysts exhibit a reduction peak at −0.27 V for ORR. Obtained results showed that the peak potential for reduction of oxygen at the hybrid nanorods, especially Fe3PtPd[PA]8(OH)2(H2O)4, is more positive than that of commercial Pt/C, shifting positively by about 0.18 mV. The higher peak current density and more positive reduction potential observed in CVs suggests that Fe3PtPd[PA]8(OH)2(H2O)4 catalysts exhibit much better catalytic activity toward ORR.

**LSVs** in O2-saturated 0.1 M KOH (Fig. 3b) were conducted to further investigate the ORR activity of Fe3Pt2[PA]8(OH)2(H2O)4, Fe3Pd2[PA]8(OH)2(H2O)4, and Fe3PtPd[PA]8(OH)2(H2O)4, and then they were compared with the commercial Pt/C catalyst. The catalytic parameters for the ORR are summarized in Table 1. The

**Table 1.** Summary of the important performance parameters of the ORR catalysts extracted from the LSV results shown in Fig. 3.

| Catalyst            | Onset potential | Current density (mA/cm2) at −0.8 V | Half-wave potential |
|---------------------|-----------------|-----------------------------------|---------------------|
| Fe3Pt2[PA]8(OH)2    | −0.014          | −0.34                             | −0.3                |
| Fe3Pd2[PA]8(OH)2    | 0.024           | −1.38                             | −0.18               |
| Fe3PtPd[PA]8(OH)2   | 0.087           | −2.36                             | −0.99               |
| Pt/C                | −0.022          | −0.99                             | −0.27               |

and Fe 2p3/2, respectively. Compared to the standard spectra of metal Fe(II), the binding energies of Fe decreases slightly. These results combined with obtained results from O 1 s spectra confirm the Fe(OH)2 formation. XPS was used to confirm the valence state of the Pd, Pt and Fe on the catalyst. As shown in Fig. 2e–g, the Pd 3d, Pt 4f and Fe 2p XPS spectra of catalyst are attributed to Pd(II), Pt(II) and Fe(II).
obtained results from Fig. 3b demonstrate that the ORR process catalyzed by the Fe₃PtPd[PA]₈(OH)₂(H₂O)₄ and Fe₃Pd₂[PA]₈(OH)₂(H₂O)₄ are a two-step two-electron pathway with the formation of intermediate HO₂⁻ ions, consistent with the reports in the literature 28,29. By contrast, Fe₃Pt₂[PA]₈(OH)₂(H₂O)₄ exhibits a one-step process for ORR. As Fig. 3b shows, the onset potentials measured for all the hybrid nanorods are positively moved from that of Pt/C. The observed onset potentials order for the hybrid nanorods and Pt/C was Fe₃PtPd[PA]₈(OH)₂(H₂O)₄ (E_{onset} = 0.087 V) > Fe₃Pd₂[PA]₈(OH)₂(H₂O)₄ (E_{onset} = 0.024 V) > Fe₃Pt₂[PA]₈(OH)₂(H₂O)₄ (E_{onset} = −0.014 V) > Pt/C (E_{onset} = −0.022 V). It is clear that the ORR onset potential measured for all the hybrid nanorods are positively moved from that of Pt/C. These results indicate that ORR activity of the Pd based hybrid nanorods is better than that of the Pt based hybrid nanorod, which is in line with previous reports 30,31. However, the Fe₃PtPd[PA]₈(OH)₂(H₂O)₄ catalyst displays the most positive onset potential and the largest limiting current density compared with Fe₃Pt₂[PA]₈(OH)₂(H₂O)₄ and Fe₃Pd₂[PA]₈(OH)₂(H₂O)₄ catalysts. The limiting current density on Fe₃PtPd[PA]₈(OH)₂(H₂O)₄ is nearly twice and seven times as large as that on Fe₃Pd₂[PA]₈(OH)₂(H₂O)₄ and Fe₃Pt₂[PA]₈(OH)₂(H₂O)₄, respectively. Furthermore, the Fe₃PtPd[PA]₈(OH)₂(H₂O)₄ catalyst exhibits considerable catalytic performance with better onset potential, half-wave potential, and reduction current compared to the commercial Pt/C. The onset potential of Fe₃PtPd[PA]₈(OH)₂(H₂O)₄ is more positive than that of Pt/C, shifting positively by about 109 mV. The enhanced specific activity for the Fe₃PtPd[PA]₈(OH)₂(H₂O)₄ catalyst should be attributed to the strain arising from the mismatch in the lattice constant among Pd, Pt, and Fe, and the ligand effect reflecting the electronic coupling among the three metals 3,32–35. These results suggest that the ORR catalytic activity of Fe₃PtPd[PA]₈(OH)₂(H₂O)₄ is better than that of commercial Pt/C catalysts, Fe₃Pt₂[PA]₈(OH)₂(H₂O)₄ and Fe₃Pd₂[PA]₈(OH)₂(H₂O)₄, indicating the advanced role of doped heteroatoms to enhance ORR activity.

To demonstrate the practical electrocatalytic activity of Fe₃PtPd[PA]₈(OH)₂(H₂O)₄ towards oxygen reduction, the CV of the electrode in 0.1 M KOH solutions saturated with N₂ or O₂ was investigated at a scan rate of 10 mV s⁻¹. As shown in Fig. 4, the CV curve of Fe₃PtPd[PA]₈(OH)₂(H₂O)₄ obtained from the N₂-saturated solutions show featureless voltammetric currents in the range of 0.2 to −0.8 V, while the CV curve obtained from the O₂ saturated solution exhibit well-defined cathodic peaks corresponding to the reduction of oxygen. This clearly demonstrates the electrocatalytic activity of Fe₃PtPd[PA]₈(OH)₂(H₂O)₄ towards oxygen reduction.

Stability studies. The durability of the catalysts and the long-term stability of the electrocatalytic activity for ORR are of prominent concern in fuel cells. The stabilities of the Fe₃PtPd[PA]₈(OH)₂(H₂O)₄ and Pt/C electrodes towards oxygen reduction are shown in Fig. 5. The dotted lines are the cycling difference from the 1st cycle to the 4000th cycle. Pt/C catalysts display a rapid decay of the signal (up to 20%) current depression after the 4000th cycle, indicating a poor stability. In contrast, the response of the Fe₃PtPd[PA]₈(OH)₂(H₂O)₄ electrode retains acceptable stability throughout the entire experiment, with only 5% current diminishations after the 4000th cycle. These results demonstrate the higher durability of Fe₃PtPd[PA]₈(OH)₂(H₂O)₄ compared with Pt/C. The promoted electrochemical stability may be due to the stronger interaction force between Pt/Pd-PA and F-OH groups than the force between Pt and C.

Scan rate effect. Figure 6 shows the representative CV curves of the Fe₃PtPd[PA]₈(OH)₂(H₂O)₄ electrode in a 0.1 M KOH aqueous electrolyte at various scan rates ranging from 5 to 100 mV s⁻¹. Clearly, an ORR peak within 0.1 to −0.4 V is visible in all the CV curves. Furthermore, a linear relation between the peak current at different scan rates and the square root of the scan rate is observed, confirming that the redox reaction is a diffusion-controlled process.
These results evidently suggest that the great promise of this hybrid Ni foam supported the Fe$_3$PtPd$_8$(OH)$_2$(H$_2$O)$_4$ nanorods electrode for high-performance ORR by long cycle life, excellent onset potential, half-wave potential, and reduction current.

**Discussions**

In summary, we have fabricated an advanced organic-inorganic hybrid nanorod catalyst with the generic formula of Fe$_3$PtPd$_8$(OH)$_2$(H$_2$O)$_4$, which boasts a greater electrocatalytic ORR activity than that of current commercial Pt/C catalysts. Specifically, the enhancement in the specific activity can be attributed to a uniform distribution of Pt and Pd complexes at Fe-OH structure, ligand effect, and strain effect arising from the lattice mismatch between Pd, Pt, and Fe. The new types of heterodoped structures may provide opportunities for further development of catalysts with high activities and a long lifetime for practical ORR applications in alkaline solutions.

**Materials**

**Synthesis of hybrid nanorods.** Hybrid nanorods with a molar ratio of Pd:Pt from 1.0 to 0.1 were prepared by solvothermal treatment. A similar synthesis method was previously reported.$^{24}$ Briefly, aqueous solution A (0.1 mol K$_2$MCl$_4$ (M = Pt, Pd with the molar ratio of 0:1, 1:1 and 1:0) in 25 mL of CO$_2$-free deionized water), solution B (picolinic acid 0.2 mol in 25 mL of CO$_2$-free deionized water), and solution C (FeCl$_2$ 0.2 mol in in 25 mL of CO$_2$-free deionized water) were prepared. Solutions (A) and (B) were first mixed, and the pH of the solution was adjusted to 10.0 by addition of 2 M of NaOH solution. Then, solution C was added dropwise with stirring. The solution pH was controlled in the range of 9.5–10.5. The mixture was placed in a 100 mL Teflon-lined stainless
auto clave and heated at 150 °C for 15 h. After being cooled to room temperature, the obtained precipitate was filtered and washed with a large amount of water until the pH value of the waste water reached 7, and then it was dried at 70 °C for 6 h.

**Oxygen reduction reaction procedure.** Electrochemical measurements for evaluation of ORR catalytic activity of the hybrid nanorods were performed using a computer-controlled potentiostat (CHI 760 C, CH Instrument, USA) with a typical three-electrode system. A nickel foam electrode (0.3 mm diameter) was used as the working electrode, a Pt foil as the counter electrode, and a saturated Ag/AgCl electrode as the reference electrode. All the experiments were conducted at room temperature (25 °C). For working electrode preparation, 1.5 mg of the catalyst was dispersed in a mixture of 0.5 mL ethanol and 20 μL of 5% Nafion under ultrasonication for 20 min. Next, 10 μL of the dispersion was uniformly dropped onto the nickel foam electrode and dried at room temperature and under ambient conditions. The commercial Pt/C (20 wt% Pt on Vulcan XC-72) electrode was prepared by the same procedure.

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Author Contributions
H. Asiabi and M. Shamsayei planned and supervised the project; Y. Yamin advised on the project; E. Saievar-Iranizad and A. Bayat designed and performed experiments; H. Asiabi and S. Kamari Kaverlavani carried out the electrochemical property tests.

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