Surface enrichment of Ir on the IrRu alloy for efficient and stable water oxidation catalysis in acid†

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Inducing the surface enrichment of active noble metal can not only help to stabilize the catalyst but also modify the catalytic performance of the catalyst through electronic and geometric effects. Herein, we report the in situ surface enrichment of Ir on IrRu alloy during the oxygen evolution reaction (OER). The surface enrichment of Ir was probed by ex situ high-resolution transmission electron microscopy (HRTEM), in situ X-ray absorption spectroscopy (XAS), and electrochemical Cu stripping, leading to complementary characterizations of the dynamic reconstruction of the IrRu alloy during OER. Guided by the density functional theory (DFT), an IrRu alloy with low Ir content (20 wt%) was constructed, which displayed a low overpotential of only 230 mV to deliver an OER current density of 10 mA cm⁻² in 0.1 M HClO₄ solution and maintained stable performance for over 20 h. To investigate the practical application potential, a proton exchange membrane (PEM) water electrolyzer using the IrRu alloy as the anode catalyst was assembled, which required a low cell voltage of only 1.48 V to generate a current density of 1 A cm⁻².

Introduction

Hydrogen is considered as the ultimate energy carrier as obtaining energy by oxidizing hydrogen produces only water, which greatly pushes the goal towards carbon neutrality. To promote the environment-friendly production of hydrogen, water electrolysis powered by renewable electricity stands out as a promising route, including proton exchange membrane (PEM) water electrolyzer, alkaline exchange membrane (AEM) water electrolyzer, and solid oxide electrolyzer (SOE). Among them, PEM water electrolyzer has the advantages of much higher current and power density, lower resistance, thinner membrane, lower gas crossover, shorter start-up times, and wider operating temperatures with smaller stack areas, which contribute to the development of water electrolyzer stacks with lower costs. However, the performance of PEM water electrolyzer is still severely limited by its anode, where the catalyst sustains high oxidation potential as well as a strong acidic environment. Currently, noble metals, especially Ru and Ir, are the only choices that give acceptable performance for the acidic oxygen evolution reaction (OER). Ru is the most active element for acidic OER; however, it is very unstable due to the dissolution of high valence ruthenium oxides under oxidative potentials. Zhao et al. reported defect-rich RuO₂ nanosheets, which could reduce the acidic OER overpotential at 10 mA cm⁻² to 210 mV but only sustained no more than 6 h. By co-alloying Ru with other transition metals (e.g., Cr and Co), its stability could be improved but still it is far from being acceptable. In comparison, Ir is much more stable in acidic OER. However, its activity is inferior to Ru and it is also more expensive than Ru.

To maintain an optimal balance between performance and cost, minimizing the noble metal usage and at the same time making full use of every noble metal site is an effective...
strategy. Loading active metal on the support can increase the active metal dispersity; however, it is only applicable to a few acid-resistant supports, considering the harsh reaction environment. Besides, tuning the intrinsic performance of OER catalysts is crucial. This can be achieved by alloying Ir or Ru with noble and/or non-noble metals, by which the electronic structure can be tuned with high flexibility. Selected examples include IrAu, IrW, IrSr, IrLi, IrNi, RuCr, and RuY. Among these, the IrRu alloy exhibits the highest acidic OER performance by maximizing the merits of both metals. Although extensive efforts have been devoted to developing IrRu alloy OER electrocatalysts, few can achieve both high catalytic activity and extensive electronic interaction.

To further balance cost and stability, structures with Ir oxide surface and heterogeneous core were designed and constructed by simple deposition, chemical leaching, electrochemical leaching and sequential reduction. The core could tune the surface electronic structure for better reactivity, while the Ir-only surface could maintain structural stability due to its much higher acid endurance. However, the poor tunability of the shell could cause troubles. If the Ir enriched shell is too thick, the core could hardly influence the electronic structure of the shell, and the intrinsic activity will not be optimized, not to mention the increased high cost induced by the thick Ir shell. If the shell is too thin or porous, the shell will not be able to protect the core from dissolution. Hence, a promising structure can be expected as the Ir-rich shell with proper thickness can balance the protection to the core from the reactive reaction and at the same time maintain an effective electronic interaction triggered by the core.

Herein, we reported an IrRu alloy catalyst loaded on TiC that could form an Ir-enriched surface during OER. This Ir-enriched surface could protect Ru from dissolution in the core, and the adsorption energetics of the Ir shell to OER intermediates could be tuned by varying the Ir/Ru ratio. The precise surface structure was probed by both microscopic and electrochemical characterizations, and in situ X-ray absorption spectroscopy (XAS) detected the valence state and coordination environment change during the reaction, verifying the in situ surface reconstruction to form the Ir-enriched shell, which greatly improved the catalytic stability in acidic OER. The IrRu alloy (Ir/Ru = 1/4) on TiC exhibited an overpotential as low as 230 mV at 10 mA cm⁻², with the cost normalized performance greatly surpassing the state-of-the-art catalysts. After a 20 hour stability test at 10 mA cm⁻², only 50 mV overpotential increase was noticed. To investigate the practical application potential, a PEM water electrolyzer using the IrRu alloy on TiC as the anode catalyst was assembled, which required a cell voltage of only 1.48 V to reach a current density of 1 A cm⁻².

Results and discussion

Structure characterization

The IrRu alloy nanoparticles (NPs) with different Ir/Ru ratios (denoted as IrₓRuᵧ) were synthesized through reduction in ethylene glycol. The physical structure of IrₓRuᵧ/TiC was characterized by high-resolution transmission electron microscopy (HRTEM), as shown in Fig. 1. An intense signal of TiC was found in the HRTEM image with a lattice spacing of about 0.22 nm corresponding to the (111) facet. The existence of the TiO₂ signal indicates the native oxidation of TiC, which was also reported in previous works. The native oxide could help NPs to be bonded more strongly to the surface of TiC. IrₓRuᵧ NPs with diameters of about 3–5 nm can be found uniformly decorated over the oxidized TiC support. Lattice spacings of 0.207 nm and 0.213 nm can be assigned to the (101) and (002) facets of the Ru(hcp) structure, respectively, while the lattice spacing of 0.22 nm corresponds to the (111) facet of the IrO₂ structure. The hcp structure was found to dominate in IrₓRuᵧ/TiC, while the fcc structure dominates in IrₓRuᵧ/TiC. A similar structure was also observed in the high-angle annular dark-field (HADDF) imaging, as displayed in Fig. S2 and S3. The X-ray diffraction (XRD) pattern, as displayed in Fig. S1, shows a weakening of the Ru signal with decreasing Ru content, which corresponds well with other reports on the IrRu alloy.

To examine the geometric and electronic structure of the as-prepared IrRu alloy, X-ray absorption spectroscopy (XAS) was performed. As shown in the X-ray near-edge fine structure (XANES), the white line intensity of Ir in IrₓRuᵧ/TiC gradually increases with increasing Ru content (Fig. 1d), corresponding to increasing electron transition from the occupied 2p³/₂ orbital to the partially occupied Ir 5d orbitals. The threshold position for Ru in IrₓRuᵧ/TiC shifts positively with increasing Ru content, suggesting an increase in the Ru valence state (Fig. 1g). These results are reasonable as Ir and Ru can be both oxidized in air, and Ru is more easily oxidized as compared to Ir. The Fourier-transformed extended X-ray absorption fine structure (EXAFS) spectra are shown in Fig. 1e and h, with detailed fitting results displayed in Fig. S5 and S6.

Tables S2 and S3 show that the Ir–Ir intensity decreases while Ru–M intensity increases with decreasing Ir/Ru ratio. The corresponding wavelet transform plots are also presented in Fig. 1f and i, from which the intensity shifts were observed between neighboring Ir- and Ru-centered coordination. Based on IrO₂ and RuO₂ standard sample profiles, the peak with the lowest coordination radius was assigned to M–O (metal–oxide) coordination. With increase k and R, M–Ir and M–Ru peaks were gradually seen. This again confirms the compositional analysis from the EXAFS and TEM results.

In situ surface enrichment characterization

To investigate how the surface structure of IrₓRuᵧ/TiC evolves during OER, in situ XAS was conducted. IrₓRuᵧ/TiC was examined because of its high measured stability. With increasing applied anodic potential, the white line intensity of Ir for IrₓRuᵧ/TiC gradually increases, with an average Ir valence state higher than +4 when the potential exceeds 1.5 V vs. saturated calomel electrode (SCE) (Fig. 2a and S7a†). The high-valence Ir is recognized as the active site for OER. The Fourier transformed EXAFS shows the same trend of iridium oxidation along

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with applied anodic potential (Fig. 2c). However, the in situ XANES spectra of Ru show that the Ru in Ir$_x$Ru$_y$/TiC maintains a low valence state (less than +2) with increasing applied anodic potential, even at 1.6 V vs. RHE (Fig. 2b and S7b†). This is further confirmed by the in situ Fourier transformed EXAFS spectra, which displays constant intensity of Ru–O coordination along with increasing anodic potential (Fig. 2d).

Ir oxidizes drastically, while Ru maintains a low valence state under OER condition, which may seem to be against the conventional understanding that Ru can oxidize more easily than Ir. To explain this unusual observation, we characterized the surface structure change of Ir$_x$Ru$_y$/TiC using the electrochemical Cu stripping method (Fig. 2e). After OER, the Ir$_x$Ru$_y$/TiC catalyst was first reduced at 0.02 V vs. SCE for 300 s and then transferred into another electrolyte with the Cu precursor for the subsequent deposition of Cu. Afterward, the catalyst was transferred into a Cu-free electrolyte to perform cyclic voltammetry measurement. As shown in Fig. 2e, the peak appearing at about 0.3 V vs. RHE is associated with bulk Cu deposition, and the peaks at about 0.4 V and 0.55 V vs. RHE can be assigned to Cu underpotential deposition (UPD) over Ru and Ir, respectively. It can be found that the surface Ru content rapidly decreased after a few OER cycles. However, the Ir signal can still be found even after 75 CV cycles. This provides a strong proof that the catalyst’s surface was in situ transformed into an Ir-enriched surface after OER, possibly due to ruthenium oxide dissolution. By quantifying the coordination number from Fourier transformed EXAFS, the trend of change in Ir–O coordination confirms that the surface was gradually oxidized with increasing anodic potential. Interestingly, the ratio ofIr–Ru coordination to Ir–Ir coordination also decreases with increasing anodic potential, indicating that the surface was gradually covered with Ir oxide, while the surface Ru was etched into the solution.
Theoretical investigation

The in situ surface enrichment of Ir on the IrRu alloy during OER was confirmed by advanced spectroscopic and electrochemical characterization. To investigate how this unique Ir-enriched surface structure influences the OER performance of the IrRu alloy, density functional theory (DFT) calculations were performed on Ir1Ru3, Ir1Ru1, Ir3Ru1, and pure Ir, representing the Ir$_x$Ru$_y$ alloy with different Ir/Ru ratios (Fig. 3a). All the surfaces were covered with a thin layer of Ir oxide to simulate the Ir-enriched surface. Due to the higher electronegativity of Ir as compared to Ru, electrons move from Ru to Ir in the bulk IrRu alloy and more negative charges accumulate on Ir along with the decreasing Ir/Ru ratio (Fig. S9†). The charge of the Ir atoms in the Ir oxide layer follows the same trend as that in the bulk (Fig. 3b). The d-band center of the active site near the Fermi level is usually linked with the adsorption energetics of reactive intermediates. As shown in Fig. 3b and S10†, the d-band center of the Ir atoms in the Ir oxide layer decreases from Ir$_3$Ru$_1$ to Ir$_1$Ru$_3$, indicating the weaker adsorption of oxygenated species on the Ir$_x$Ru$_y$ alloy with a lower Ir/Ru ratio.

Then, we calculated the energetics of OER at $T = 298.15$ K and $P = 1$ atm (Fig. 3c). It was found that the rate-determining step (RDS) of OER is at the 3rd electron-transfer step, namely, the oxidation of O* to OOH*, which is limited by the weak adsorption of OOH* on Ir$_1$Ru$_3$, Ir$_1$Ru$_1$, Ir$_3$Ru$_1$, and Ir with a free energy barrier of 0.45 eV, 0.49 eV, 0.51 eV, and 0.52 eV, respectively (Fig. 3c). Thus, we can predict that the OER activity on the Ir$_x$Ru$_y$ alloy will follow an increasing order along with decreasing Ir/Ru ratio. However, with increasing Ru content in the Ir$_x$Ru$_y$ alloy, the surface Ru atoms can be more easily

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Fig. 2  In situ XANES for (a) Ir and (b) Ru. In situ Fourier transformed EXAFS for (c) Ir and (d) Ru in Ir$_1$Ru$_4$/TiC. (e) Cu stripping of Ir$_1$Ru$_4$/TiC.

Fig. 3  (a) Structural models of Ir$_3$Ru$_5$, Ir$_1$Ru$_4$, and Ir$_3$Ru$_1$ used in the DFT calculation. The balls in green, brown, and red represent Ir, Ru, and O atoms, respectively. (b) The calculated Bader charge of Ir atom in the Ir oxide layer, overpotential of OER, and d-band center of Ir atom in the Ir oxide layer. (c) Calculated OER Gibbs free energy diagram at 1.23 V vs. RHE, $T = 298.15$ K, and $P = 1$ atm.
leached, causing an instability issue. Therefore, tuning the ratio of Ir/Ru in the Ir\textsubscript{x}Ru\textsubscript{y} alloy to balance the OER activity and stability is crucial to deliver optimal catalytic performance.

**Electrocatalytic performance**

The electrocatalytic performance of the Ir\textsubscript{x}Ru\textsubscript{y} alloy NPs with different Ir/Ru ratios was examined in 0.1 M HClO\textsubscript{4} solution. As shown in Fig. 4a, the linear sweep voltammetry (LSV) test indicates the highest OER activity for Ir\textsubscript{1}Ru\textsubscript{4}/TiC, which exhibits the lowest overpotential of only 230 mV at 10 mA cm\textsuperscript{-2}, surpassing the commercial IrO\textsubscript{2} catalyst and most state-of-the-art OER catalysts reported in the literature. The OER performance of other Ir\textsubscript{x}Ru\textsubscript{y}/TiC is displayed in Fig. 4a with the activity trend matching well with the DFT prediction. Besides the activity, the stability of the catalyst was examined by chronopotentiometry at 10 mA cm\textsuperscript{-2}, as displayed in Fig. 4b. The Ir\textsubscript{1}Ru\textsubscript{4}/TiC can be sustained for more than 20 hours with less than 50 mV potential increase, which also surpasses most of the noble-metal-based OER catalysts in acid reported in the literature (Table S7\textsuperscript{†}). From the inductively coupled plasma-optical emission spectrometry (ICP-OES) results of the samples before and after the 5 hour OER test (Table S1\textsuperscript{†}), the Ir/Ru ratio could be quantified to change from 1 : 3.9 to 1 : 2.5, corresponding to the fast dissolution of Ru and surface enrichment of Ir, as shown before. Apart from the activity and stability, economic consideration is also crucial for the large-scale deployment of the PEM water electrolyzer. We compared the activity normalized by the noble metal cost, as shown in Fig. 4c, together with the stability (detailed comparison can be found in Table S7\textsuperscript{†}). It is clear that Ir\textsubscript{1}Ru\textsubscript{4}/TiC outperforms the other acidic OER catalysts, reaching a current of 1 A at the cost of 3.3 USD. To further study the practical application potential at an industrialized scale, a PEM water electrolyzer was assembled using the as-prepared Ir\textsubscript{1}Ru\textsubscript{4}/TiC as the anode catalyst (Fig. 4d), which is able to deliver a current density of 1 A cm\textsuperscript{-2} at a cell voltage as low as 1.49 V.

**Conclusion**

OER-induced surface enrichment of Ir on the IrRu alloy targeted for excellent OER performance was for the first time probed by ex \textit{situ} high-resolution transmission electron microscopy, \textit{in situ} X-ray absorption spectroscopy, and electrochemical Cu stripping. The characterization results indicated the fast dissolution of Ru and oxidation to form an active and protective Ir oxide shell on the surface of the IrRu alloy during OER. The Ir-enriched surface not only protected Ru in the core against oxidation but also offered catalytically active sites that drive the water oxidation reaction. Furthermore, the core with a high content of Ru could not only optimize the electronic structure of the Ir shell, but also significantly reduce the Ir usage, which allowed the Ir\textsubscript{1}Ru\textsubscript{4}/TiC catalyst to deliver a current of 1 A at a cost of only 3.3 USD, significantly outperforming the other acidic OER catalysts reported in the literature.

**Data availability**

All data is available in the main text or the ESI.\textsuperscript{†}

**Author contributions**

J. Z., X. L. and L. B. designed the idea. J. Z. and X. L. conducted the material synthesis and characterizations. Y.-F. J. and C. Q. X., and J. L. conducted the density functional theory calculation. S.-F. H. conducted the in-situ XAS characterization. W. L.
conducted the high resolution TEM characterization. J. Z., X. C., H. B. Y., D-S. L., T. Z., Y. L., J. L., and B. L. write the manuscript.

Conflicts of interest

Authors declare no competing interests.

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