Subnanometer high-entropy alloy nanowires enable remarkable hydrogen oxidation catalysis

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High-entropy alloys (HEAs) with unique physicochemical properties have attracted tremendous attention in many fields, yet the precise control on dimension and morphology at atomic level remains formidable challenges. Herein, we synthesize unique PtRuNiCoFeMo HEA subnanometer nanowires (SNWs) for alkaline hydrogen oxidation reaction (HOR). The mass and specific activities of HEA SNWs/C reach 6.75 A mgPt−1 and 8.96 mA cm−2, respectively, which are 2.8/2.6, 4.1/2.4, and 19.8/18.7 times higher than those of HEA NPs/C, commercial PtRu/C and Pt/C, respectively. It can even display enhanced resistance to CO poisoning during HOR in the presence of 1000 ppm CO. Density functional theory calculations reveal that the strong interactions between different metal sites in HEA SNWs can greatly regulate the binding strength of proton and hydroxyl, and therefore enhances the HOR activity. This work not only provides a viable synthetic route for the fabrication of Pt-based HEA subnano/nano materials, but also promotes the fundamental researches on catalysis and beyond.

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Hydrogen fuel cell is considered as one of the most promising energy conversion devices due to its high energy efficiency and pollution-free feature. Over the past decades, great efforts have been devoted to developing catalysts for hydrogen oxidation reaction (HOR) for enhancing the energy conversion efficiency of the hydrogen fuel cell. The ideal catalysts for HOR should simultaneously fulfill the features of high activity, high durability as well as low cost. As the state-of-the-art catalyst, PtRu-based alloys have been widely used for HOR in recent years. Despite great progress has been achieved, PtRu-based catalysts still suffer from the drawbacks, including high cost, poor stability, and low resistance to CO-poisoning. It is thus highly desired to develop highly efficient catalysts for HOR under operating conditions.

Recently, high-entropy alloys (HEAs) have attracted increasing attention in diverse fields due to their inherent properties, including high thermal stability, strong mechanical strength, excellent corrosion resistance, and fine trade-off effect on performance. As promising catalysts in heterogeneous catalysis, HEAs generally consist of five or more metals, which display the structure-dependent synergistic effects and enhanced catalytic performance in terms of activity and durability. Moreover, since the HEAs can largely decrease the usage of noble metals and thus reduce the cost of catalyst, they have been attracting increasing research interests. For instance, Chen et al. reported a small HEAs PtNi2Fe2Co3.3Mn0.7 nanoparticles (NPs) with a mean size of ~3.4 nm as an efficient catalyst for alkaline methanol oxidation reaction, on which the synergistic effects facilitate the site-to-site electron transfer during both reduction and oxidation processes. Cui et al. demonstrated that the synergistic effects between different metals in CrMnFeCoNi HEA sul fate can regulate the electronic states and thus enhance the oxygen evolution reaction activity. Despite the great potentials of HEAs, the controlled synthesis of PtRu-based HEAs with tailored morphologies and sizes still remains great challenges, which severely impedes their applications. To this end, it is highly desirable to develop facile strategies for fabricating PtRu-based HEAs.

Herein, we fabricated PtRuNiCoFeMo HEA subnanometer nanowires (SNWs) as highly active and durable catalyst for HOR. In particular, the mass activity of HEA SNWs for HOR is 4.1 and 19.8 times higher than that of commercial PtRu/C and Pt/C, respectively. Moreover, no obvious decay of HOR performance was observed after 2000 cycles in the accelerated durability test (ADT), suggesting the promising stability of HEA SNWs for HOR. Additionally, HEA SNWs can even display superior resistance to CO poisoning compared to commercial PtRu/C and Pt/C in the presence of 1000 ppm CO. Density functional theory (DFT) calculations confirm that the strong interactions between different metals in HEA SNWs can regulate the electronic structures of different metals and thus enhance the HOR activity. In particular, Co and Ni sites maintain the highly stable valence states due to the pinning effect by nearby Fe and Mo sites, and the Pt and Ru sites modulate the overall electroactivity for superior HOR performance. This work may not only provide a facile protocol for the controllable synthesis of HEA SNWs, but also promote the fundamental researches on HEAs for catalysis and beyond.

Results

Material synthesis and characterization. For the synthesis of HEA SNWs, we adopted platinum (II) acetylacetonate (Pt(acac)2), ruthenium (III) acetylacetonate (Ru(acac)3), nickel (II) acetylacetonate (Ni(acac)2), cobalt (III) acetylacetonate (Co(acac)3), iron (III) acetylacetonate (Fe(acac)3), and molybdenum hexacarbonyl (Mo(CO)6) as the metal precursors, oleylamine (OAm) as the solvent, stearyl trimethyl ammonium bromide (STAB) as the structure-directing agent, and glucose as the reducing agent, respectively. Transmission electron microscopy (TEM) and high-angle annular dark-field scanning TEM (HAADF-STEM) images show that the obtained HEA SNWs has a mean diameter of 1.8 ± 0.3 nm (Fig. 1a and Supplementary Fig. 1). The characteristic peaks in the X-ray diffraction (XRD) pattern are ascribed to face-centered cubic (fcc) structure of Pt (JCPDS No. 04-8002) and the positive shifts of peaks in XRD pattern imply the formation of alloy (Fig. 1b). On the other hand, the weakening and broadening of peaks in the XRD pattern could be attributed to the lattice distortion in HEA SNWs. A scheme was thus provided to better understand the structure of HEA SNWs (Fig. 1c). Scanning electron microscopy energy-dispersive X-ray spectroscopy (SEM-EDS) shows that the composition of HEA SNWs is Pt/Ru/Co/Fe/Mo = 29.6/9.3/15.2/19.8/19.5 (Supplementary Fig. 2). Moreover, both HAADF-STEM and line scan analysis show that all the metals are uniformly distributed in the HEA SNWs. Furthermore, the aberration-corrected high-resolution STEM (HRSTEM) images show that the obtained HEA SNW has a distorted structure with abundant atomic steps and defect-rich lattice mismatch (Fig. 1f–h). In addition, a 3D atomic model was provided to depict the structure of HEA SNWs (Fig. 1i), which has vividly revealed the unique structure of HEA SNWs with plentiful surface atomic steps and numerous facet boundaries.

HOR performance of PtRuNiCoFeMo HEA SNWs. To evaluate the potential applications of the as-prepared HEAs, PtRuNiCoFeMo HEA SNWs were used as catalyst for HOR, a significant process for hydrogen fuel cell. Prior to HOR test, HEA SNWs were loaded on Vulcan XC-72R carbon (HEA SNWs/C, Supplementary Fig. 10a), and home-made HEA NPs (Supplementary Fig. 9) loaded on Vulcan XC-72R C (HEA NPs/C), commercial PtRu/C and Pt/C were selected as the references (Supplementary Fig. 10b–d and Fig. 11). Fourier transform-infrared (FT-IR) spectrum of HEA SNWs shows that most of the ligands have been removed from the surface of HEA SNWs/C (Supplementary Fig. 12). From the cyclic voltammograms (CVs) curves of various catalysts obtained in 0.1 M HClO4 solution at a scan rate of 50 mV s⁻¹ (Supplementary Fig. 13), we can see that the presence of Ru significantly broadens the electrical double-layer capacitor. It is found that the anode current of HEA SNWs/C increases sharply with the potential (Fig. 2a). In particular, HEA SNWs/C exhibits a high and steady current density with the increased potential, whereas a gradual decay in the current density was observed for HEA NPs/C, suggesting that HEA SNWs/C can be serve as a highly efficient catalyst for HOR. Moreover, the HEA SNWs/C for HOR was evaluated in N₂-saturated 0.1 M KOH solution to measure the anodic current. As shown in Fig. 2a (yellow line), the current density is ~0 when the potential is over 0.1 V, and the weak current density at 0–0.1 V is ascribed to Ru oxidation, which confirms the occurrence of HOR in H₂-saturated 0.1 M KOH solution. The Koutecky–Levich plot was then collected to verify the electron transport process (Supplementary Fig. 14). It is found that the slopes for HEA NPs/C, PtRu/C, and Pt/C are 10.78, 11.36, and 16.95 mA cm⁻² ppm⁻¹/², respectively. In contrast, the slope for HEA SNWs/C is 14.06 mA cm⁻² ppm⁻¹/², which is close to the theoretical value (15.06 mA cm⁻² ppm⁻¹/²), further confirming the H₂ mass-transport controlled process (Supplementary
Fig. 1 Characterizations of HEA SNWs. a HAADF-STEM image. b XRD pattern. c Crystal structure. d SEM-EDS and atomic ratio of different elements. e HAADF-STEM-EDS elemental mappings. f–h Aberration-corrected HRSTEM images. i 3D models and enlarged atomic model of HEA SNWs.

Fig. 2 HOR performance evaluation of HEA SNWs/C and other catalysts. a Polarization curves in H₂-saturated 0.1 M KOH. b Tafel plots. c Normalized mass activity and specific activity at an overpotential of 50 mV vs. RHE. d Normalized mass activity and specific activity before and after 2000 cycles of ADTs. e Relative current-time chronoamperometry response of different catalysts in 1000 ppm CO/H₂-saturated 0.1 M KOH at 100 mV vs. RHE.
Mechanism investigation. To reveal the mechanism for the enhanced HOR performance over HEA SNWs/C, the surface properties of catalyst were studied by X-ray photoelectron spectroscopy (XPS) measurement. It is noted that all the metal elements in HEA SNWs are mixed with metal state and oxidation state. Compared to HEA NPs/C, the peak in the Pt$^0$ 4f XPS spectrum of HEA SNWs/C positively shifts by 0.13 eV (Fig. 3a), while the peak in the Ru$^3$ 3p XPS spectrum negatively shifts by 0.36 eV (Fig. 3b), indicating that electrons might transfer from Pt to Ru in HEA SNWs. On the other hand, the Ni$^0$ 2p, Co$^0$ 2p, Fe$^0$ 2p and Mo$^{4+}$ 3d XPS spectra of HEA SNWs positively shift by 0.26, 0.17, 0.93 and 0.25 eV, respectively, in comparison to HEA NPs (Supplementary Fig. 22). Considering that only Ru$^3$ 3p XPS spectrum negatively shifts by 0.36 eV, we might conclude that the positive shifts of Ni$^0$ 2p, Co$^0$ 2p, Fe$^0$ 2p, and Mo$^{4+}$ 3d XPS spectra were attributed to the electron transfer from Ni, Co, Fe and Mo to Ru in HEA SNWs. To further study the influence of electron transfer on the binding strength of reactants and intermediates during HOR, we compared the d-band center of various catalysts. Compared to values of HEA NPs/C (−3.732 eV), PtRu/C (−3.631 eV) and Pt/C (−3.548 eV), the d-band center of HEA SNWs/C shifts upwards to −3.835 eV (Fig. 3c, d), which thereby weakens the adsorption of H but facilitates the adsorption of OH intermediate$^{28,39}$. Considering the significant effect of Ru on HOR that can facilitate the adsorption of OH$^\cdot$ species$^{30,31}$, we evaluated the HOR performance over HEA SNWs and PtNiCoFeMo HEA SNWs/C. The mass activity of HEA SNWs/C is 11.8 times higher than that of PtNiCoFeMo HEA SNWs/C (Supplementary Fig. 23a, b). Moreover, compared to the PtNiCoFeMo HEA SNWs/C, HEA SNWs/C displays significantly improved resistance to CO poisoning (Supplementary Fig. 23c). In particular, the HOR activity in the presence of 1000 ppm CO decreases by 26.4% after 2000 s when HEA SNWs/C was used as catalyst, which is much smaller than that of PtNiCoFeMo HEA SNWs/C (85.2%), confirming the significance of Ru on HOR. Additionally, the position of the CO stripping peak of HEA SNWs/C shifts positively compared with Pt/C, which further suggests that the presence of Ru significantly improves the resistance to CO poisoning (Fig. 3d and Supplementary Fig. 16). The aforementioned results imply that the strong electronic interaction between different elements in HEA SNWs can regulate the adsorption abilities of H and OH species, and then enhances the alkaline HOR activity (Fig. 3e).

DFT calculations were conducted to further investigate the mechanism for the enhanced HOR performance. It is noted that both Ni and Ru sites show subtle distortions while the overall HEA SNWs remain stable, which confirms the stability of HEA SNWs/C (Fig. 4a, b). With respect to the electronic structures, the strong coupling between the bonding and anti-bonding orbitals on the HEA surfaces delivers a high HOR activity with efficient site-to-site electron transfer between different metal sites. Moreover, the detailed electronic structures of HEA SNWs are also demonstrated by the projected partial density of states (PDOSs) (Fig. 4c$^{32}$). Notably, the evident overlaps between d-orbitals are observed, indicating the strong bonding between different metals. In particular, we notice the evident sharp Ni 3d orbitals locate near −1.30 eV, which is a key indicator for the strong adsorptions of H$^\cdot$ and OH$^\cdot$. Meanwhile, Co 3d orbitals show a similar position with the Ni 3d orbitals, which may further facilitate the electron transfer. Similar phenomena were observed for Fe 3d and Mo 4d orbitals, and the overlap of d orbital might not only enhance the electron transfer, but also impose the pinning effect on the 3d orbitals of Ni and Co sites for a robust electroactivity. On the other hand, Pt 5d orbital locates at the furthest position to Fermi level, suggesting that Ru plays as the electron reservoir to balance the valence state of HEA NWs during electrocatalysis. Moreover, the Ru 4d ranges from −6.0 to +4.0 eV, which enables the flexible electronic modulations and accelerates the site-to-site electron transfer on the HEA surfaces. Furthermore, the site-dependent electronic structure of each element in HEA SNWs has been demonstrated. As shown in Fig. 4d, the Pt−5d center gradually approaches to the Fermi level from the bulk (cyan color) to the surface of HEA SNWs (green color), indicating that the H$_2$ adsorption becomes much stronger on the surface than on the bulk of HEA SNWs. It is noted that the overall Pt−5d orbital is further away from the Fermi level compared to the pure Pt metal.
and the oxhydryl binding energy (OHBE) of HEA SNWs/C improved electroactivity of HEA surface, which guarantees the possibility in charge transfer and bonding chances with OH. Besides, from the bulk to the surface, the gradual absence of the eg-t2g band splitting of Co 3d orbitals leads to the efficient intra-orbital eg-t2g electron transfer rather than transfer between d-d orbitals, which further promotes the site-to-site electron transfer from catalysts to adsorbates (Fig. 4f). It is also noted that the electronic structure of Co in HEA SNWs are similar from the bulk to the surface. Such a robust electronic structure is attributed to the double pinning effect by the shielding effect from neighboring Fe/Mo and Ru orbitals. For Fe sites, the eg-t2g band splitting for 3d orbitals decreases from 3.33 to 2.79 eV from the bulk site to the surface sites, supporting a more efficient HOR. Compared with the state-of-the-art PtRu/C and Pt/C catalysts, HEA SNWs/C exhibits advantages, including high activity, enhanced stability, and promising resistance to CO poisoning, being a promising catalyst for alkaline HOR. In particular, the mass activity and specific activity of HEA SNWs reach 6.75 A mgPt−1 and 8.96 mA cm−2, which are 2.8/2.6, 4.1/2.4, and 19.8/18.7 times higher than those of HEA NP/C, commercial PtRu/C and Pt/C, respectively. DFT calculations reveal that the strong synergy significantly varies the electronic properties of different elements in HEA SNWs, which further regulates the HBE and OHBE. This work not only provides a viable synthetic route for the fabrication of Pt-based HEA subnano/nano materials, but also promotes the fundamental researches on catalysis and beyond.

Discussion

In summary, we have demonstrated a successful synthesis of PtRuNiCoFeMo HEA SNWs with abundant surface atomic steps and facet boundaries for efficient HOR. Compared with the state-of-the-art PtRu/C and Pt/C catalysts, HEA SNWs/C exhibits advantages, including high activity, enhanced stability, and promising resistance to CO poisoning, being a promising catalyst for alkaline HOR. In particular, the mass activity and specific activity of HEA SNWs reach 6.75 A mgPt−1 and 8.96 mA cm−2, which are 2.8/2.6, 4.1/2.4, and 19.8/18.7 times higher than those of HEA NP/C, commercial PtRu/C and Pt/C, respectively. DFT calculations reveal that the strong synergy significantly varies the electronic properties of different elements in HEA SNWs, which further regulates the HBE and OHBE. This work not only provides a viable synthetic route for the fabrication of Pt-based HEA subnano/nano materials, but also promotes the fundamental researches on catalysis and beyond.
Methods

Chemicals. Pt(acac)₂ (97%), Ni(acac)₂ (95%), Co(acac)₃ (98%), Fe(acac)₃ (98%), and STAB (98%) were purchased from Sigma-Aldrich. Ru(acac)₃ (97%) was supplied by Alfa Aesar. Mo(CO)₆ (98%) was purchased from Strem Chemicals Inc. Glucose (C₆H₁₂O₆, analytical reagent), potassium hydrate (KOH, analytical reagent), and STAB (98%) were purchased from Sigma-Aldrich. Ru(acac)₃ (97%) was supplied by Aladdin. HClO₄ (analytical reagent, 70–99.7%) and isopropanol (C₃H₈O, analytical reagent, ≥ 99.7%) were purchased from Tianjin Chemical Products Co. Ltd. All chemical reagents were used as received without further purification. All aqueous solutions were prepared using deionized water with a resistivity of 18.2 MΩ cm⁻¹.

Materials synthesis. In a typical synthesis of PtRuNiCoFeMo HEA SNWs, 9.8 mg Pt(acac)₂, 5.0 mg Ru(acac)₃, 3.8 mg Ni(acac)₂, 7.0 mg Co(acac)₃, 7.0 mg Fe(acac)₃, 10 mg Mo(CO)₆, 60 mg glucose, and 39.3 mg STAB were dissolved in 5 mL OAm, followed by ultrasonication for 2 h. The mixture was then heated in an oil bath at 210 °C for 5 h. The cooled product was collected by centrifugation and washed three times with a cyclohexane/ethanol (v/v = 1:9) mixture. For the synthesis of PtRuNiCoFeMo HEA NPs, all of the conditions are similar to those of HEA SNWs except for the additions of STAB and glucose.

Characterizations. XRD measurement was conducted on a SmartLab-SE powder diffractometer equipped with a Cu radiation source (λ = 0.15406 nm). SEM-EDS was observed through ZEISS Sigma 300 field emission scanning electron microscope. TEM was operated on JEM-1400 TEM at an accelerating voltage of 100 kV. HRTEM was conducted on a FEI Tecnai G3000. The X-ray photoelectron spectroscopy spectra were collected by XPS (Thermo Scientific, ESCALAB 250Xi). The carbon peak at 284.6 eV was used as the reference to correct for charging effects. The concentrations of the catalysts were determined by ICP-OES (ICAP 7000, ThermoFisher, USA). FT-IR was performed with KBr in the range of 4000–400 cm⁻¹ (Nicolet i550).

Electrochemical measurements. The electrocatalytic properties toward HOR were evaluated using a RDE (Pine, diameter of 5 mm) with CHI 7600E Electrochemical Workstation. A SCE and a graphite rod were used as the reference electrode and counter electrode, respectively. HOR measurements were performed in 0.1 M KOH solutions purged with high-purity H₂. The sweep rate was 5 mV s⁻¹ to collect the HOR polarization curves. The accelerated durability tests were carried out in 0.1 M KOH between −0.1 V and 0.4 V vs. RHE at a scan rate of 500 mV s⁻¹ for 2000 cycles. The CO antipoisoning tests were carried out in 0.1 M KOH purged with saturated 1000 ppm CO/H₂, and the long-range stability test voltage was at 100 mV vs. RHE.

The kinetic current (I_k) was calculated using the Koutecky–Levich equation:

\[
I_k = \frac{1}{f} = \frac{1}{I_a} + \frac{1}{I_d}
\]

where \(I\) is measured current and \(I_k\) is diffusion-limited current, which can be fitted to the Butler–Volmer equation and Levich equation:

\[
I_d = 0.62nFvD^{1/2}n^{1/2}C_s^{1/2}ω^{1/2} = BC_s^{1/2}ω^{1/2}
\]

in which \(n\) and \(α_a\) are the anodic and cathodic transfer coefficients (\(α_a + α_c = 1\)), \(F\) is the Faraday constant, \(η\) is the overpotential, \(R\) is the universal gas constant, \(T\) is the Kelvin temperature, \(n\) is the electrons number, \(D\) is the diffusion coefficient, \(v\) is the viscosity coefficient, \(C_s\) is the solubility, \(ω\) is the rotating speed, and \(B\) is the Levich constant, respectively.

For the CO stripping measurement, CO gas was bubbled into 0.1 M HClO₄ solution for 30 min. The electrode was quickly moved to a fresh 0.1 M HClO₄ solution for 30 min. The electrode was quickly moved to a fresh 0.1 M HClO₄ solution for 30 min.
solution and recorded the two cycles to calculate CO stripping peak. The electrochemical surface area of the noble metals was obtained using Eqs. (4) and (5).

\[
Q_{\text{adsorption}}(C) = \int \frac{1}{\text{d}E} \text{d}(\text{mA}) \quad \text{volts} ^{-1}
\]

\[
\text{ECSA} \quad \frac{m^2}{g} = \left[ \frac{Q_{\text{adsorption}}(C)}{4200 \mu F_{\text{Pt}}(\text{PtRu}) \text{mg}} \right]^{0.5}
\]

**Calculation setup.** To investigate the electronic structures, we have utilized DFT calculations within CASTEP packages in this work. To achieve accurate descriptions of the exchange-correlation energy, we applied the generalized gradient approximation (GGA) and Perdew–Burke–Ernzerhof (PBE) in all the calculations. Meanwhile, we have set the cutoff energy of the plane-wave basis to be 380 eV based on the ultra-high quality. The data that support the findings of this study are available from the corresponding author upon request. Source data are provided with this paper.

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Author contributions
X.H. conceived and supervised the research. X.H. and Y.X. designed the experiments. X.H., C.Z., Y.X., L.B., T.Y., and Q.S. performed data analysis and experiments discussions. H.Z., Z.Y., Y.F., and Y.Z. performed and analyzed the HAADF-STEM and corresponding EDS characterizations. B.H. performed the DFT simulations. X.H., L.B., C.Z., and Y.X. wrote the paper. All authors discussed the results and commented on the manuscript.

Competing interests
The authors declare no competing interests.

Additional information
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