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Constructing four-in-one catalysts to realize ultralow voltage hydrogen production at ampere-level current densities†

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An anion exchange membrane water electrolyzer (AEMWE) is emerging as key technology for hydrogen production. However, its widespread application requires further reduction of cost and improvement of efficiencies. Here, we synthesize a four-in-one catalyst ($V_{SA}\text{-CoN}_x$) to achieve high-efficiency coupling hydrogen production by combining with the hydrazine oxidation reaction (HzOR) and the urea oxidation reaction (UOR). The as-synthesized $V_{SA}\text{-CoN}_x$ exhibits excellent performance in all the four reactions of HzOR, UOR and hydrogen/oxygen evolution reactions (HER/OER). The HER–HzOR coupling system only requires an ultra-low voltage of 0.21 V to deliver an ampere-level current density (1 A cm^{-2}), while the conventional HER–OER AEMWE needs nearly an input of 1.88 V. Remarkably, this HER–HzOR coupling system largely reduces the energy expenditure of the AEMWE by approximately 90%, which hits a record in the low energy cost for all water electrolysis systems known to date. Given the energy consumption of the traditional AEMWE of approximately 4.56 kW h Nm^{-3} of H_2 at a current density of 1 A cm^{-2} , the HER–HzOR AEM electrolyzer only requires 0.51 kW h Nm^{-3} of H_2 . This HER–HzOR coupling system not only significantly lowers the energy expenditure of large-scale H_2 production but also addresses the hydrazine-associated environmental pollution.

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Broader context

To construct a sustainably developing society, hydrogen, as a promising clean energy carrier, stands out due to its high energy density and zero-carbon emissions. An anion exchange membrane water electrolyzer (AEMWE) to produce hydrogen has been considered as one of the most promising hydrogen production technologies. However, its high energy costs have severely limited the extensive applications. Using small molecule oxidation as an alternative to the oxygen evolution reaction (OER) may be a useful solution to reduce the energy expenditure of large-scale H_2 production. In this work, we develop a four-in-one catalyst, which can catalyze four reactions, to achieve high-efficiency coupling hydrogen production by combining with the hydrazine oxidation reaction (HzOR) and the hydrogen evolution reaction (HER). The HER–HzOR coupling system only requires an ultralow voltage of 0.21 V to deliver an ampere-level current density (1 A cm^{-2}), while the conventional HER–OER AEMWE needs nearly an input of 1.88 V, which largely reduces the energy expenditure of the AEMWE by approximately 90%, hitting a record in the low energy cost for all water electrolysis systems known to date. Simultaneously, this HER–HzOR coupling system also alleviates the hydrazine-associated environmental pollution. It is expected that this work provides a useful coupling strategy to reduce the energy expenditure of large-scale H_2 production.

1. Introduction

Building a sustainably developing society heavily relies on effective utilization of clean energy sources such as solar, wind, and hydrogen. As a promising energy carrier, hydrogen energy stands out due to its high energy density and zero carbon emissions.¹ Harnessing renewable energies for water electrolyzers (WEs) has been increasingly considered as a highly promising approach to H_2 production.² The cathodic hydrogen evolution reaction (HER) and the anodic oxygen evolution reaction (OER) are usually coupled in a WE cell.³ Compared with

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the HER, the four-electron transfer OER is a relatively slow process with sluggish kinetics.⁴ The high theoretical overpotential (1.23 V vs. reversible hydrogen electrode (RHE)) of the OER severely hinders its practical applications,⁵ which requires high energy cost in the overall water splitting (OWS) reaction.⁶ Previous studies mainly focused on optimizing the catalysts for the HER and/or OER to achieve high-performance OWS.⁷ To reduce the energy cost of OWS, the electrochemical hydrogen-production coupled with an alternative oxidation (EHCO) reaction may be a brand-new route for green H₂ production.

In the EHCO route, coupling the HER and thermodynamically favorable small molecule oxidation with relatively low theoretical potentials to replace the OER can not only reduce the energy consumption of the overall reaction, but also simultaneously yield high-value-added products or facilitate wastewater treatment in the anode.⁸ In fact, a series of organic molecular oxidation reactions, such as those involving methanol,⁹ ethanol,¹⁰ urea,¹¹ 5-hydroxymethylfurfural,¹² and hydrazine,¹³ have been used to replace the OER. Among them, urea and hydrazine, as the common substances in wastewater, are harmful to the environment and human beings. Compared with the OER (1.23 V vs. RHE), the urea and hydrazine oxidation reactions (UOR/HzOR) can be performed at much lower standard potentials (0.37 V vs. RHE for UOR, and -0.33 V vs. RHE for HzOR). Therefore, the anodic UOR or HzOR coupled with the cathodic HER can not only effectively reduce the energy consumption of H₂ production, but also provide an opportunity to purify wastewater rich in urea or hydrazine, demonstrating significant potential for practical applications.^{11d,14} In this scenario, developing high-performance multi-functional catalysts is extremely important, which not only targets desirable reactions with high selectivity, but also significantly streamlines the design and fabrication process of catalytic electrodes.¹⁵

Recently, various catalysts, including noble metals^{14c,15a,16} and non-noble metals,¹⁷ have been developed to improve efficiencies of H₂ production. Heteroatom doping engineering has been widely used as an effective modification strategy of catalyst development, due to its synergistic effects in optimizing the electronic structure, regulating the adsorption of intermediates, and promoting electron and mass transfers.¹⁸ Previous studies have shown that the incorporation of W-O groups with strong adsorption capacity into CoP can serve as H₂O dissociation and N₂H₄ dehydration sites, effectively promoting the coupling of the HER with the HzOR.^{13a} Similarly, Mo-doping was reported to optimize the electronic structure and achieve thermoneutral ΔG_{H^+} for the HER, which therefore raises dehydrogenation kinetics from *N₂H₄ to *NHNH₂ for the HzOR.¹⁹ However, few studies have been reported at ampere-level current density for large-scale H₂ production. Actually, developing multifunctional catalysts with excellent HER, OER, UOR and HzOR activities at ampere-level current density is crucial for H₂ production *via* an EHCO route.

Herein, we synthesize a single-atom V-doped CoN_x four-in-one-functional catalyst (V_{SA}-CoN_x), where V doping can not only optimize the surface structure of catalysts, but also promote the generation of active substances and inhibit the accumulation of

high-valent metal species in the high-voltage reaction process. The HER-UOR coupling system based on V_{SA}-CoN_x as bifunctional catalysts with anion exchange membrane (AEM) only requires an input of 1.64 V to deliver a current density of 1 A cm⁻², while the HER-HzOR AEM coupling system even lowers the potential to 0.21 V under the same conditions, hitting a record in the low energy consumption for all water electrolysis systems known to date. This work offers a new route of devising multifunctional catalysts to achieve green H₂ production at ampere-level current densities by the EHCO approach, paving the way for a sustainable future in green hydrogen development.

2. Results and discussion

2.1. Synthesis and structure characterization

The catalyst V_{SA}-CoN_x was synthesized through a hydrothermal and nitridation procedure (Fig. 1a). First, the treated nickel foam (NF) was put into a homogeneous solution containing vanadium (V) and cobalt (Co) metal salts, and the precursor V-Co(OH)₂ was obtained by a hydrothermal method. Then, the final single-atom V-doped CoN_x sample (V_{SA}-CoN_x) can be obtained by calcination in a nitrogen atmosphere. For comparison, the CoN_x catalyst was also prepared *via* the same method, but without V salt in the aqueous solution.

Compared to the two samples of Co(OH)₂ and CoN_x without V-doping, V-doping can effectively reduce the aggregation of the V_{SA}-CoN_x and achieve better dispersion as shown in Fig. 1b and Fig. S1-S4 (ESI[†]). Moreover, the surface hydrophilicity (aerophobicity) of the V_{SA}-CoN_x was changed by V-doping (Fig. S5, ESI[†]), where V_{SA}-CoN_x has a smaller liquid contact angle and a relatively larger bubble contact angle than CoN_x. The change can not only maximize the active site of the V_{SA}-CoN_x catalyst in the reaction process, but also effectively prevent the formation of large bubbles resulting in bubble blockage. Fig. 1c and Fig. S6 (ESI[†]) confirm the existence of the CoN_x phase in V_{SA}-CoN_x, and the lattice fringe with 0.213 nm matches with the (200) plane of CoN_x. HAADF-STEM and electron energy loss spectroscopy (EELS) also proved the existence of elements N, V and Co, where V is atomically dispersed in the catalyst (Fig. 1d). Furthermore, the TEM-EDS mapping indicates a homogeneous distribution of Co, V, and N throughout the nanorod (Fig. 1e). In addition, the XPS results of V_{SA}-CoN_x and CoN_x (Fig. S7, ESI[†]) indicate that V-doping does not produce a strong electron interaction on the main catalyst phase, which suggests that the electron interaction between V and CoN_x is not responsible for the enhancement of catalyst activity.

The X-ray diffraction (XRD) pattern (Fig. 2a) shows that both samples match well with the standard card JCPDS#15-0806, which indicates that no additional crystalline phases were formed in the V-doping process. However, upon closer examination of the enlarged areas in Fig. 2b, it is observed that the diffraction peaks of V_{SA}-CoN_x have shifted to lower angles relative to CoN_x. This shift becomes more obvious with an increase in the X-ray diffraction angle, suggesting a lattice expansion,



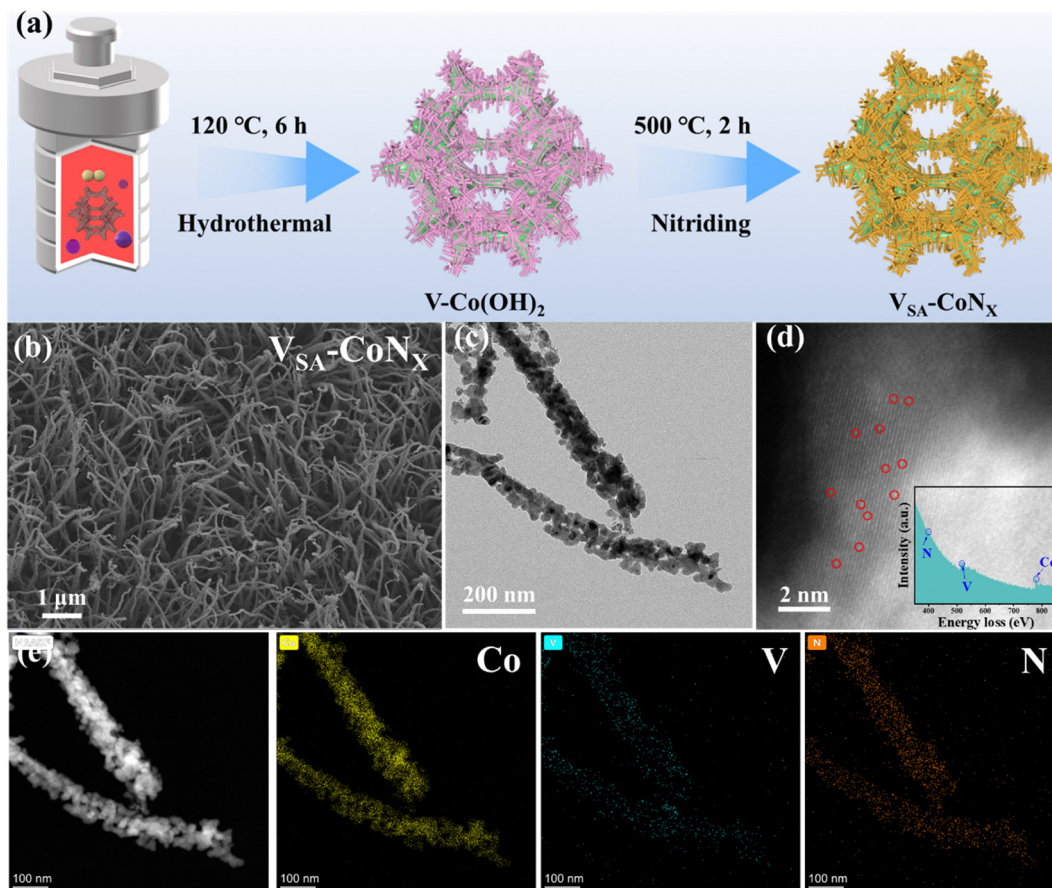


Fig. 1 Synthesis illustration and morphological characterization of $V_{SA}\text{-CoN}_x$. (a) Synthesis route. (b) SEM, (c) TEM, and (d) HAADF-STEM images (the inset in (d) is the EELS analysis). (e) STEM-EDS mappings.

as dictated by the Bragg diffraction formula. This observation indicates that the incorporation of V into the CoN_x structure has led to an increase in the interplanar spacing within the lattice.^{6b,20} The electronic structure and coordination environment of the catalyst were also analyzed by X-ray absorption spectroscopy (XAS). Fig. 2c depicts the Co K-edge XANES spectra of the Co foil, CoO, CoN_x and $V_{SA}\text{-CoN}_x$. The absorption edge of the Co K-edge of CoN_x and $V_{SA}\text{-CoN}_x$ is between Co foil and CoO, suggesting that the state of Co is situated between 0 and +2. The over-lapped Co K-edge XANES spectra of CoN_x and $V_{SA}\text{-CoN}_x$ also verify that the electronic environments of Co sites were not significantly influenced by the incorporation of V, which is in accordance with the results of Co $2p_{3/2}$ XPS.²¹ The XANES spectra at the V K-edge (Fig. 2d) further provide comparable insights, revealing that the valence state of V in $V_{SA}\text{-CoN}_x$ lies between +3 and +4 when compared with the standard samples (V foil, V_2O_3 , VO_2 , and V_2O_5). Furthermore, the Fourier transform (FT) of the k^3 -weighted EXAFS oscillations in the R space for both Co and V in $V_{SA}\text{-CoN}_x$, as shown in Fig. 2e and f, elucidates the local coordination environment around these elements. The predominant peaks at approximately 1.46 and 1.47 Å are attributed to the first shell coordination of Co–N and V–N/O, respectively. Notably, a significant peak at around 2.15 Å is observed in the $V_{SA}\text{-CoN}_x$ sample, which supports

the presence of a Co–V bond. This peak aligns with the one at approximately 2.16 Å in the R -space of the V K-edge. Moreover, the Co–N bond length in $V_{SA}\text{-CoN}_x$ increased significantly from 1.40 to 1.46 Å, compared to the CoN_x sample without V-doping, which further proves that due to the incorporation of metal V, more unsaturated metal sites were formed, resulting in a lattice expansion of the CoN_x phase.²² Strain engineering can often be used as a very effective method to improve the performance of catalytic materials.²³ The EXAFS fitting was also carried out to identify the coordination structure of the Co atom in $V_{SA}\text{-CoN}_x$ (Fig. S8, ESI[†]), and the fitting parameters are shown in Table S6. The EXAFS fitting curves of the other samples (Co foil, CoO and CoN_x) are shown in Fig. S9 and S10 (ESI[†]). Compared with CoN_x , the atomic distance of the Co–N bond in $V_{SA}\text{-CoN}_x$ increases from 1.97 to 2.02 Å due to the V-doping. The average coordination number of Co–Co in CoN_x samples was 4.5. The Co–Co coordination number in $V_{SA}\text{-CoN}_x$ decreased to 4.1, while Co–V bonds appeared with a coordination number of 0.8. We conducted ICP tests on the catalytic materials. As shown in Table S7 (ESI[†]), the ratio of metals Co to V is approximately 4 : 1, which is consistent with the parameters fitted by using EXAFS. All these results suggest that V atoms may partially replace Co atoms in the CoN_x and exists in the single-atom form (Fig. S11, ESI[†]).



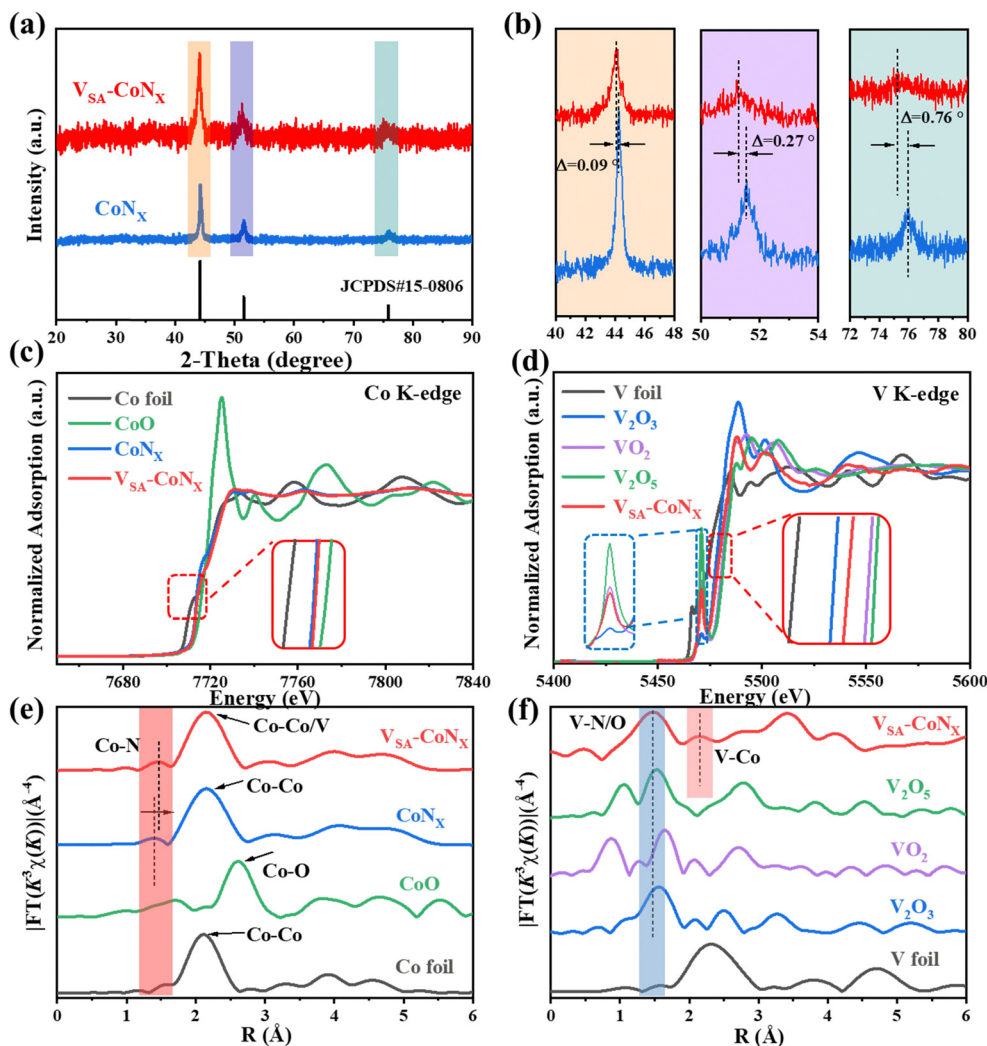


Fig. 2 XRD characterization and coordination environment of samples. (a) XRD patterns of $V_{SA}\text{-CoN}_x$ and CoN_x . (b) The XRD of magnified regions with different colors. (c) Co K-edge and (d) V K-edge XANES spectra, and (e) and (f) the corresponding Fourier transforms of EXAFS spectra of different samples.

2.2. HER-OER and HER-UOR performance

The HER and OER performances of all the samples were tested in a 1 M KOH solution, where the applied potentials relative to the reversible hydrogen electrode (RHE) were converted using the equation: $E_{\text{RHE}} = E_{\text{Hg/HgO}} + 0.925 \text{ V}$ (Fig. S12, ESI[†]). Fig. 3a and Fig. S13 (ESI[†]) show that $V_{SA}\text{-CoN}_x$ only needed overpotentials of $\sim 135 \text{ mV}$ at a current density of 100 mA cm^{-2} for the HER and 1.565 V to yield a current density of 50 mA cm^{-2} for the OER (Fig. 3c), which exceeded the commercial Pt/C catalyst ($\eta_{100} = 148 \text{ mV}$) for the HER and commercial RuO_2 ($\eta_{50} = 1.580 \text{ V}$) for the OER. The Tafel slope also proves its superior chemical kinetics (Fig. S14 and S15, ESI[†]). Fig. S16 and S17 (ESI[†]) show the negligible change in LSV before and after the stability test, which confirmed its high stability. Moreover, characterization of $V_{SA}\text{-CoN}_x$ after the HER stability test indicates that it well maintained the morphology (Fig. S18, ESI[†]) and electronic structure (Fig. S19, ESI[†]). It is noted that, unlike in the HER, after the OER long-term stability test, the original

nanowires of the catalyst surface exhibit an obvious nanosheet region (Fig. S20, ESI[†]). According to previous reports, the region of the nanosheet should be the *in situ* formed high-valence Co species (CoOOH) during the high-voltage stability test. The lattice constants corresponding to the CoOOH and the XPS tests also confirmed this observation (Fig. S21, ESI[†]).²⁴

Electrochemical impedance spectroscopy (EIS) further demonstrates that $V_{SA}\text{-CoN}_x$ has the faster catalytic process (Fig. S22, S23 and Table S1, ESI[†]). Furthermore, EIS measurements were conducted at various voltages to probe the charge transfer resistance (R_{ct}) and the electrocatalytic kinetics involved in the HER. When the applied voltages were ranging from -0.015 V to -0.085 V (Fig. S24a and b, ESI[†]), $V_{SA}\text{-CoN}_x$ presented significantly lower mass transfer impedance. The Bode phase plots in Fig. S24c and d (ESI[†]) illustrate the phase angle as a function of frequency, which provides insights into the reaction dynamics at the electrolyte-catalyst interface and within the catalyst. The peaks in the phase angle at low and



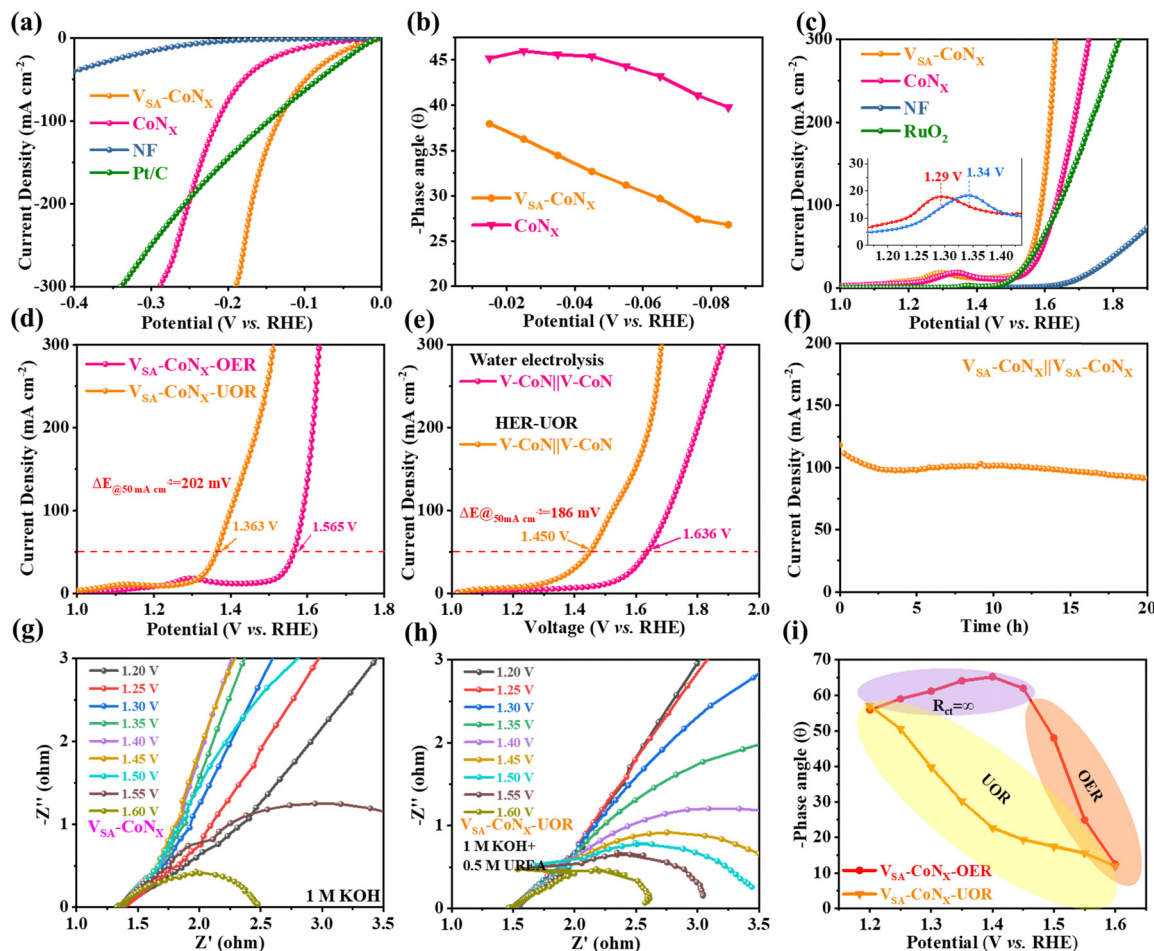


Fig. 3 Electrochemical performance of the electrodes for water electrolysis and urea electrolysis. (a) The HER polarization curves and (b) phase peak angles of $V_{SA}\text{-CoN}_x$ and CoN_x . (c) The OER polarization curves. (d) Polarization curves of $V_{SA}\text{-CoN}_x$ in the 1 M KOH electrolyte in the presence and absence of urea. (e) The LSV curves of $V_{SA}\text{-CoN}_x||V_{SA}\text{-CoN}_x$ for the water (HER–OER) and urea electrolysis (HER–UOR). (f) The stability test of $V_{SA}\text{-CoN}_x||V_{SA}\text{-CoN}_x$ for the HER–UOR. Nyquist plots of at different voltages of $V_{SA}\text{-CoN}_x$ for the (g) OER and (h) UOR. (i) Phase peak angles of $V_{SA}\text{-CoN}_x$ for the OER and UOR.

high frequencies are associated with the charge transfer at the electrolyte–catalyst interface and the electron conduction through the catalyst’s inner layer, respectively.²⁵ In the low-frequency region, the noticeably smaller phase peak angle observed for $V_{SA}\text{-CoN}_x$ (Fig. 3b) indicates that the incorporation of V into the CoN_x structure enhanced the charge transfer at the electrolyte–catalyst interface. This enhancement contributes to faster HER kinetics by facilitating the transfer of electrons between the electrolyte and the catalyst, which is crucial for the efficiency of the HER process.

Considering the excellent performance of the HER and OER of $V_{SA}\text{-CoN}_x$, we assembled a two-electrode electrolyzer by using $V_{SA}\text{-CoN}_x$ as a bifunctional catalyst for the anode and cathode. As shown in Fig. S25 (ESI[†]), it needs 1.636 V to provide a current density of 50 mA cm^{-2} . Moreover, the current density (50 mA cm^{-2}) negligibly attenuates during the stability test over 200 h (Fig. S26, ESI[†]). Compared with CoN_x without V-doping, $V_{SA}\text{-CoN}_x$ shows much better catalytic performance. Notably, the HER/OER and OWS performance based on the $V_{SA}\text{-CoN}_x$ catalyst are standing out among the state-of-the-art catalysts (Table S2, ESI[†]).

Different from the OER process (1.23 V vs. RHE), the electrochemical oxidation process of urea requires only 0.37 V.²⁶ Inspired by the above excellent HER/OER performance of $V_{SA}\text{-CoN}_x$, we further explored its UOR performance. Fig. 3d and Fig. S27 and S28 (ESI[†]) show that the UOR performance of the $V_{SA}\text{-CoN}_x$ in the 1 M KOH electrolyte containing 0.5 M urea. Impressively, the $V_{SA}\text{-CoN}_x$ catalyst only needs a potential of 1.363 V to deliver a current density of 50 mA cm^{-2} in the UOR. Compared to the OER in the KOH electrolyte without urea, the overpotential of $V_{SA}\text{-CoN}_x$ presents a 202 mV decrease at 50 mA cm^{-2} after the addition of urea. In order to verify the feasibility of using the UOR as a substitute of the OER, a two-electrode HER–UOR coupling system was constructed with KOH solution containing 0.5 M urea as the anode. The cell voltage of the HER–UOR system at 50 mA cm^{-2} is only 1.450 V (Fig. 3e), which is much lower than that of 1.636 V for OWS. Moreover, the potential of HER–UOR coupling system is apparently lower than OWS in all current densities (Fig. S29, ESI[†]), and also lower than many other electrocatalysts in HER–UOR systems (Table S3, ESI[†]). The time–current curve in Fig. 3f at



about 100 mA cm^{-2} also demonstrates its excellent stability. Similar to OER, high-valence Co species appeared on the catalyst surface after UOR stability testing (Fig. S30, ESI[†]), and XPS results (Fig. S31, ESI[†]) also demonstrated the conversion of Co and the sacrifice of surface V.

To unveil the mechanism of activity enhancement for $V_{SA}\text{-CoN}_x$ in different electrolytes, the EIS at different potentials in different electrolytes was also measured to understand the relationship between interfacial dynamics, electron transfer and mass transfer in the processes of OER and UOR.²⁷ It is clear from Nyquist that the R_{ct} is infinite when the voltage is below 1.4 V without the addition of urea (Fig. 3g), which indicates that no reaction occurred in the lower voltage range. With the increase of voltage, the corresponding phase angle in KOH even tended to increase (Fig. 3i and Fig. S32a, ESI[†]). In contrast, the phase angle of Bode phase plots in the urea-containing electrolyte decreased continuously as the potential increases (Fig. 3h–f and Fig. S32, ESI[†]), indicating that the catalyst can react in all the range of applied voltages. This intrinsically fast kinetics also explains why the UOR performed better than the OER.²⁸

In order to further investigate the structural changes of the prepared catalysts at high voltages, a series of *in situ* Raman tests were performed. Fig. 4a shows the time-dependent *in situ* Raman diffraction peaks of the catalyst at a fixed voltage, and the original catalyst ($t = 0 \text{ min}$) exhibits the characteristic peaks of the initial CoN_x and CoO_x .²⁹ When a constant voltage was applied, $V_{SA}\text{-CoN}_x$ yields new Raman peaks corresponding to Co–O (H) at approximately 499 and 573 cm^{-1} . The contour map

was gradually formed (Fig. 4a₁–d₁) with the intensity increase of corresponding peaks, and the Raman peak corresponding to VO_x also appeared in 867 cm^{-1} and was stabilized over time.³⁰ Unlike $V_{SA}\text{-CoN}_x$, although CoN_x had a similar diffraction peak (Co–O (H)) around 499 cm^{-1} (Fig. 4b), its peak intensity did not become stronger over time, even if the time was prolonged to 15 min. The above analysis shows that the two as-prepared catalysts (CoN_x and $V_{SA}\text{-CoN}_x$) will undergo a pre-oxidation process in the alkaline electrolyte, and the V-doping can effectively promote the pre-oxidation of the catalyst surface, which is beneficial for the following reaction.

After the pre-oxidation process, different voltages were applied to investigate the phase transition of catalysts in the real process. Fig. 4c shows that the Raman peaks of $V_{SA}\text{-CoN}_x$ at 479 and 573 cm^{-1} shifted to 461 and 568 cm^{-1} , respectively, when the voltage was increased to 1.275 V, indicating the formation of a higher valence Co species (CoOOH) on the surface during the reaction. It is consistent with the previous study where the *in situ* generated high-valence Co species at high voltages were regarded as the real active substances.³¹ Concurrently, the Raman peak at 867 cm^{-1} gradually disappeared with the increase of the voltage, which also explains the disappearance of the surface V after the stability test under high voltages (Fig. S21 and S31, ESI[†]). Fig. 4d shows the Raman spectra of CoN_x at different voltages. Clearly, a diffraction peak corresponding to the high-valence Co species appeared at 1.325 V. The peak intensity (CoOOH) of CoN_x increased while the peak at 499 cm^{-1} disappeared with increasing voltage, indicating that CoN_x underwent a continuous oxidation under

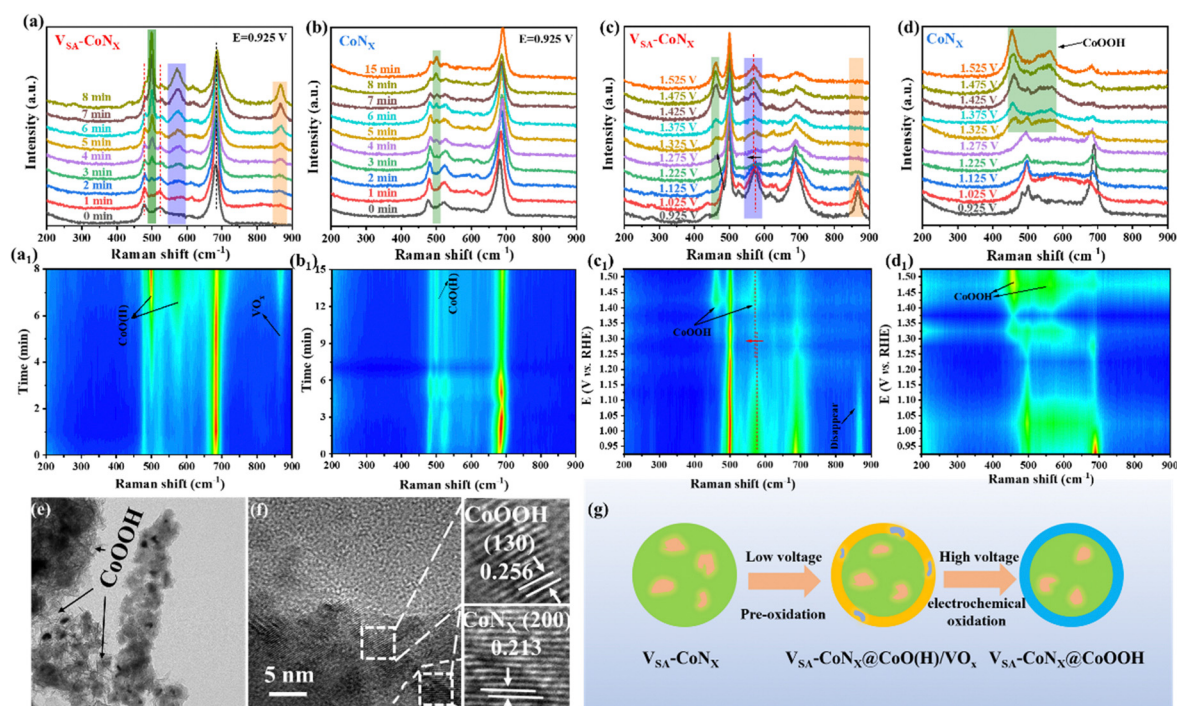


Fig. 4 Time-dependent *in situ* Raman spectra of (a) $V_{SA}\text{-CoN}_x$ and (b) CoN_x at a constant voltage. Potential-dependent *in situ* Raman spectra of (c) $V_{SA}\text{-CoN}_x$ and (d) CoN_x . (a₁)–(d₁) The corresponding contour maps. (e) TEM and (f) HRTEM images of $V_{SA}\text{-CoN}_x$ after the UOR stability test. (g) Schematic representation of active species in $V_{SA}\text{-CoN}_x$ under different voltages.



high voltage, and the surface was all converted into high-valence Co species. However, for $V_{SA}\text{-CoN}_x$ (Fig. 4c), with the increase of voltage, the intensity of the corresponding peak of CoOOH kept constant, meaning that V-doping can effectively inhibit the over-oxidation of Co at the high voltage. The TME and HRTEM images of $V_{SA}\text{-CoN}_x$ after the UOR (Fig. 4e and f) show the apparent coexistence of CoOOH and CoN_x . Therefore, V-doping promotes the production of active substances and inhibits the accumulation of high-valency species (Fig. 4g), which makes it promising to achieve excellent catalytic activity and stability at the high voltage.

2.4. HER-HzOR performance

In addition to urea, hydrazine has also been widely used as industrial raw materials. As a common type of pollutants, it is significant to develop an effective strategy to degrade hydrazine sewage for environmental protection.³² Compared with the urea oxidation, the hydrazine oxidation (HzOR) can be performed at an even lower standard voltage (-0.33 V vs. RHE), which means that the HER-HzOR system may further lower

energy consumption of H_2 production. Since the products of the HzOR are nitrogen and water, it is a greenhouse gas emission-free reaction. Therefore, the HzOR is an ideal alternative for the anodic reaction to couple with the HER. Impressively, $V_{SA}\text{-CoN}_x$ showed a surprisingly low potential ($\eta_{50} = -0.033$ V), and also presented the lower mass transfer resistance to prove the origin of its excellent performance (Fig. S33, ESI†). Compared with the traditional anodic OER, the voltage-drop reached 1.598 V at 50 mA cm^{-2} (Fig. 5a), which was larger than 1.5 V in the wide current ranges (Fig. 5b), indicating its extraordinary promise to replace the OER. Therefore, a two-electrode HER-HzOR system with $V_{SA}\text{-CoN}_x$ as bifunctional catalysts was constructed, and it only required 0.064 V to reach 50 mA cm^{-2} (Fig. 5c), which was significantly lower than 1.636 V at OWS and 1.450 V at the HER-UOR system. In addition, only 0.302 V was required at 300 mA cm^{-2} , which shows an extremely low energy cost for practical green H_2 production (Fig. S34 and Table S4, ESI†). The stability of the two-electrode electrolyzer was also tested and is shown in Fig. 5d. During the stability test, the current density continued

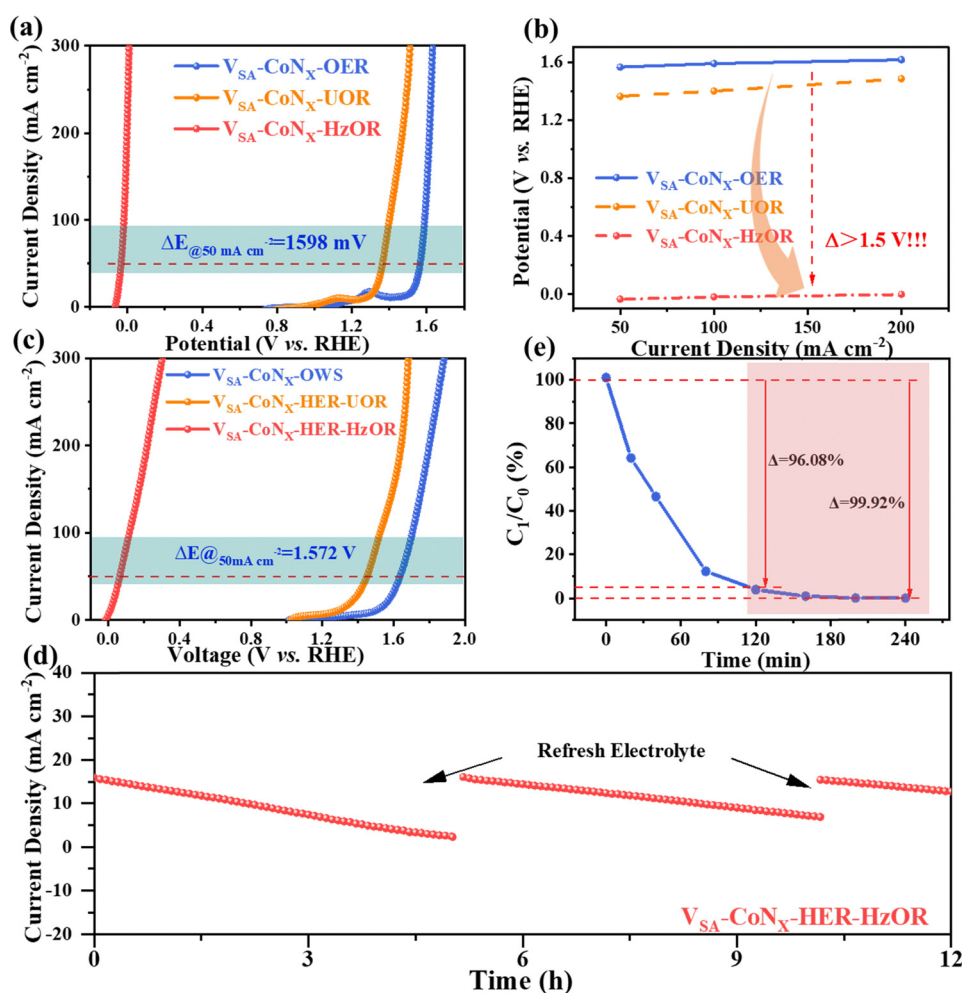


Fig. 5 Electrocatalytic performance of the electrodes for hydrazine electrolysis. (a) Polarization curves and (b) driving potentials at different current densities of $V_{SA}\text{-CoN}_x$ for the OER, UOR and HzOR. (c) Polarization curves of $V_{SA}\text{-CoN}_x$ for the OWS, HER-UOR and HER-HzOR. (d) The stability test of $V_{SA}\text{-CoN}_x\text{-HER-HzOR}$. (e) Degradation of the hydrazine electrolyte.



to decrease, but it is noted that once the electrolyte was refreshed, and the current density was restored immediately. This indicates that the decay of current density was due to the decrease of the hydrazine concentration in the electrolyte. Similar to the HER, the morphology (Fig. S35, ESI[†]) and electronic structure (Fig. S36, ESI[†]) of the catalyst did not change obviously after the stability test. Since the waste water containing hydrazine is very harmful and carcinogenic to the environment, the degradation rate of hydrazine by the HER-HzOR system was also studied by UV-vis spectroscopy. As expected, in the HER-HzOR system, the degradation rate of hydrazine in the anode electrode can reach over 95% at 2 h by two cycles, and the hydrazine concentration can be reduced to 6 ppb in 3 h, lower than the 10 ppb required by the U. S. Environmental Protection Agency (EPA) (Fig. 5e and Fig. S37, S38, ESI[†]).^{15a} This HER-HzOR coupling system is promising to the largely lower energy expenditure of H₂ production while simultaneously treating wastewater containing hydrazine.

2.5. HER-OER (UOR/HzOR)-AEMWE performance

As mentioned above, V_{SA}-CoN_x has shown excellent catalytic performance in the HER, OER, UOR and HzOR as multifunctional

catalysts. However, the H-type electrolyzer at the laboratory stage is still far from the industrial level. Therefore, to further test the commercial prospect of the catalyst material, AEMWEs using V_{SA}-CoN_x as bifunctional catalysts were also constructed (Fig. 6a). A current density of 1 A cm⁻² was reached with an input voltage of 1.88 V on the AEMWE for OWS, which is superior to many reported catalysts (Fig. 6b and Table S3, ESI[†]). To test the adaptability of the AEMWE for intermittent wind and solar powers, we also simulated this type of electricity fluctuation by changing the input voltage and the input current in a short time to study its response. Clearly, the AEMWE with V_{SA}-CoN_x as bifunctional catalysts exhibited excellent response to the change of voltage or current (Fig. S39, ESI[†]), which offers the possibility for the use of renewable electricity.

Encouraged by above excellent performance, we further constructed the HER-UOR AEMWE cell, and it needs a lower input voltage of 1.64 V to deliver a current density of 1 A cm⁻², which exhibits apparently lowered energy consumption for H₂ production, compared with the AEMWE for OWS. To further reduce the energy cost, the AEMWE system with the HzOR replacing the OER was constructed, which only required a voltage of 0.21 V to reach a current density of 1 A cm⁻², resulting in a

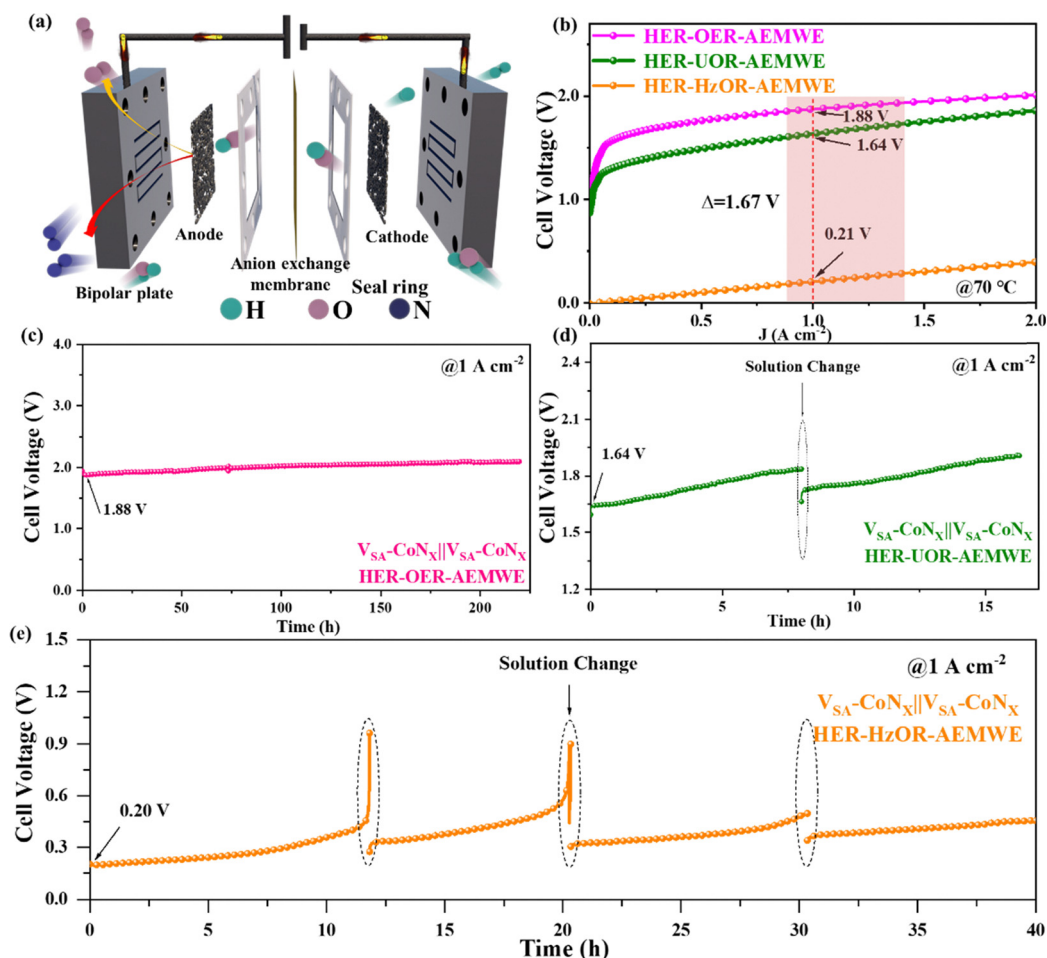


Fig. 6 (a) Schematic illustration of the two-electrode MEA flow electrolyzer based on samples. (b) LSV curves for the HER-OER-AEMWE, HER-UOR-AEMWE and HER-HzOR-AEMWE. The stability test of V_{SA}-CoN_x||V_{SA}-CoN_x for the (c) HER-OER-AEMWE, (d) HER-UOR-AEMWE and (e) HER-HzOR-AEMWE.



voltage-drop of 1.67 V from the conventional AEMWE. This HER–HzOR AEM electrolyzer shows an unprecedented ultralow voltage to deliver a current density of 1 A cm⁻², setting a new record among all AEM water electrolysis systems reported to date (Table S5, ESI†). Compared with conventional AEMWE devices (Fig. S40, ESI†), the HER–UOR and HER–HzOR systems only consume about 87.23% and 11.17% energy for the same amount of H₂ generation. An evaluation of energy consumption revealed that at 1 A cm⁻², if the energy consumption of the traditional AEMWE is about 4.56 kW h Nm⁻³ of H₂, it is 3.95 kW h Nm⁻³ of H₂ (equivalent to 43.954 kW h kg⁻¹ of H₂) for the HER–UOR AEM system and only 0.51 kW h Nm⁻³ of H₂ (equivalent to 5.628 kW h kg⁻¹ of H₂) for the HER–HzOR AEM system,³³ which is apparent lower than that of commercial PEM electrolyzers, about 4.5 to 5.0 kW h Nm⁻³ of H₂ (50.00–55.56 kW h kg⁻¹ of H₂).³⁴ The ultralow energy cost of the HER–HzOR AEM system for producing hydrogen provides a broad prospect for practical applications.³⁵

The stability of the traditional AEMWE cell was further tested (Fig. 6c), which maintained over 200 h at 1 A cm⁻², providing a promising potential for traditional water electrolysis. The HER–UOR system also shows excellent stability (Fig. 6d), where the voltage increase was caused by the decrease of the urea concentration. When the urea electrolyte was refreshed, the initial voltage was also recovered, accompanying with a slight increase. Similarly, the HER–HzOR system also exhibited the same behavior (Fig. 6e), which was due to the decrease of the hydrazine concentration. Table S7 (ESI†) also shows that the weight of the catalyst remained almost the same before and after the reaction, and the ratio of the two metals, Co and V, did not change significantly. This further confirms that the degradation of performance observed during the stability test was ascribed to the change in the hydrazine concentration, rather than the dissolution of the catalyst itself. Owing to the excellent multifunctional properties of V_{SA}-CoN_x for the HER, OER, UOR and HzOR, the AEMWE systems based on our catalysts under multi-operating conditions (including HER–OER, HER–UOR and HER–HzOR) presented outstanding performance for H₂ production. In particular, the HER–HzOR system only consumed about 11% energy of the traditional AEMWE for producing the same amount of hydrogen, which hits a record of ultralow energy cost at a current density of 1 A cm⁻² among all AEM water electrolysis systems reported so far.

3. Conclusions

In summary, a four-in-one catalyst of V_{SA}-CoN_x has been developed, and the V_{SA}-CoN_x can efficiently catalyze four reactions HER, OER, UOR and HzOR. By using the V_{SA}-CoN_x catalyst as both an anode and a cathode, the assembled AEMWE systems can achieve coupling H₂ production under multi-operating conditions, including HER–OER, HER–UOR and HER–HzOR conditions. In particular, the HER–UOR and HER–HzOR AEM coupling systems only require 1.64 and 0.21 V to achieve an ampere-level current density (1 A cm⁻²), while the conventional HER–OER AEMWE system needs 1.88 V. The energy consumption

of the HER–HzOR ECHO system is just about 11% of the conventional AEMWE system, which hits a record in the low energy expenditure for all water electrolysis systems known to date. Simultaneously, the HER–HzOR ECHO system can efficiently degrade hydrazine over 95% in 2 h and achieve the treatment of wastewater containing hydrazine. At a current density of 1 A cm⁻², if the energy consumption of the conventional AEMWE is about 4.56 kW h Nm⁻³ of H₂, it is 3.95 kW h Nm⁻³ of H₂ (equivalent to 43.954 kW h kg⁻¹ of H₂) for the HER–UOR AEM electrolyzer and 0.51 kW h Nm⁻³ of H₂ (equivalent to 5.628 kW h kg⁻¹ of H₂) for the HER–HzOR AEM electrolyzer. In short, this study offers a new perspective of achieving ultralow-cost H₂ production by coupling alternative oxidation reactions under multiple operating conditions.

Author contributions

D. C. conceived and designed the project. M. X. performed all the experiments and analyzed the results. M. H., G. X. performed a part of characterization. Q. Z. and M. S. analyzed the electrochemical data. Z. L., J. Y. and P. W. gave the supervision. M. X. and D. C. wrote the manuscript. All authors have reviewed, discussed and approved the results and conclusions of this article.

Data availability

The data supporting this article have been included as a part of the ESI.†

Conflicts of interest

The authors declare no competing interest.

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