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Activating Janus charge distribution on the P-doped Ni₃S₂/Co₉S₈ interface for enhancing charge-matched urea adsorption: boosting high current hydrogen production *via* coupled urine degradation†

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The urea oxidation reaction (UOR) is emerging as a thermodynamically favorable alternative to the oxygen evolution reaction, offering significant potential for energy-efficient H₂ production and simultaneous treatment of urea-rich wastewater. However, the 6 e⁻ transfer process of the UOR results in sluggish kinetics, necessitating the development of highly efficient electrocatalysts. Herein, a Janus charge distribution surface is constructed by incorporating phosphorus (P) into the Ni₃S₂/Co₉S₈ heterojunction to enhance the UOR performance and accelerate urea-assisted H₂ production. The P incorporation facilitates electron transfer from Ni₃S₂ to Co₉S₈, creating a local electrophilic/nucleophilic interface that enhances the adsorption of urea molecules with the electron-withdrawing C=O group and electron-donating amino groups. As a result, the modified P-Ni₃S₂/Co₉S₈ exhibits ultralow potentials of 1.22, 1.30 and 1.39 V (*versus* the reversible hydrogen electrode) to reach 10, 100 and 1000 mA cm⁻² for the UOR, respectively. Remarkably, when alkaline urine is used as the electrolyte, the P-Ni₃S₂/Co₉S₈ catalyst, functioning as a bifunctional electrocatalyst in an anion-exchange membrane electrolyzer, can stably deliver a high current density of 1000 mA cm⁻² for H₂ production over 180 h. This work highlights the importance of designing electrocatalysts by activating interfacial charge distribution to enhance reactant adsorption and trigger chemical bond cleavage.

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1 Introduction

Urea is one of the simplest organic molecules and serves as a crucial component for various life forms. For animals and humans, urea is generated as a nitrogenous end product from the decomposition of proteins and is excreted in urine.¹ Approximately 265 million tons of urine containing 2–2.5 wt% urea are produced daily, which is more than 500 times the

market demand for urea.² After the urine-rich wastewater is released directly into the environment, the untreated urea decomposes naturally into toxic ammonia/nitrate, or nitrogen oxide, causing irreversible environmental damage. Thus, the treatment of urine-rich wastewater has become a major environmental concern.^{3,4} Conventional methods for urea removal, such as adsorption, thermal hydrolysis and chemical oxidation, typically require high energy consumption and are ineffective in treating high loads of urea nitrogen. The electrocatalytic urea oxidation reaction (UOR) offers a promising approach for treating urea/urine wastewater, operating efficiently under ambient conditions with renewable electricity input. Moreover, the UOR has a favorable thermodynamic equilibrium potential of 0.37 V (all potentials are reported *versus* the reversible hydrogen electrode, RHE), compared to the oxygen evolution reaction (OER) of 1.23 V. Theoretically, replacing the OER with the UOR for the coupled hydrogen evolution reaction (HER) can reduce energy consumption for H₂ production by up to 70%.⁵ In addition to its energy-saving potential, the UOR not only has the potential to save energy but also offers advantages in material accessibility and environmental friendliness. Unlike water electrolysis, which depends on abundant water resources, the

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UOR utilizes urea from wastewater, converting it into valuable products like hydrogen. This approach not only addresses wastewater management challenges but also promotes resource recycling.⁶ These dual benefits make the UOR a more environmentally friendly and sustainable hydrogen production method compared to traditional water electrolysis.

Despite the thermodynamic advantages, the UOR involves a $6e^-$ transfer step ($\text{Co}(\text{NH}_2)_2 + 6\text{OH}^- \rightarrow \text{N}_2 + 5\text{H}_2\text{O} + \text{CO}_2 + 6e^-$), which results in sluggish reaction kinetics,⁵ making the development of efficient electrocatalysts critical and urgent.^{7,8} In nature, urea decomposition is accelerated by the urease-catalyzed process, where the amine group is bonded to one nickel site of the enzyme and the oxygen atom is attached to another. For the UOR occurring in a heterogeneous electrocatalytic manner, the adsorption behaviors of electrocatalysts to urea are crucial in determining the performance. Accordingly, significant efforts have been made to regulate the adsorption energy or configuration of urea through heteroatom doping,^{9–11} vacancy incorporation^{12,13} and interface engineering.^{14,15} Among these strategies, interface engineering is notable for constructing heterojunctions with adjustable surface states.^{16,17} Strong coupling between different components facilitates electron transfer and redistribution at the interface, forming a Janus charge distribution surface.^{18,19} Given that urea molecules consist of one electron-withdrawing group (C=O) and two electro-donating groups (amino), the electrophilic domain at the interface benefits the $-\text{NH}_2$ group adsorption, while the nucleophilic domain favors C=O group adsorption. Therefore, constructing heterojunctions and optimizing the charge redistribution at the interface are believed to enhance urea adsorption and improve the UOR performance. Additionally, understanding the adsorption behaviors at the interface for reactants is crucial for guiding the design of active heterojunction electrocatalysts.

Transition metal sulfides, characterized by their unfilled outer d-orbitals and 3d valence electron structure, have emerged as promising catalysts due to their high catalytic activity, low cost, and excellent conductivity.²⁰ The recent advances of pentlandites for wastewater treatment further exemplify the versatility and applicability of transition metal sulfides in addressing environmental challenges.^{21,22} These materials exhibit unique advantages in applications such as the organic electro-oxidation reaction,²³ OER,²⁴ and HER.²⁵ Recent studies have highlighted their significant potential in bifunctional catalysis, making them ideal alternatives to precious metal catalysts. For example, Mo^{4+} -doped NiS has demonstrated remarkable performance in the UOR and HER, attributed to synergistic effects that enhance electron transfer and provide abundant active sites.⁹ Similarly, Fe-modified Ni hydroxysulfide (Fe-NiSOH) nanosheet arrays exhibit rapid self-reconstruction through *in situ* sulfur leaching, promoting the formation of active species.²⁶ This design achieves low overpotentials and excellent stability in both water and seawater oxidation. These advancements underscore the potential of transition metal sulfides in developing efficient and durable electrocatalysts for sustainable energy applications.

Herein, we construct a P-modified $\text{Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$ heterostructure (P- $\text{Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$) as a bifunctional electrocatalyst for the UOR and HER. The incorporation of P aims to regulate the charge distribution at the interface and further modulate the adsorption behaviors of urea molecules. Experimental results and theoretical calculations indicate that the introduced P promotes electron transfer from Ni_3S_2 to Co_9S_8 and enhances urea adsorption at the electrophilic (Ni_3S_2) and nucleophilic (Co_9S_8) interfaces. Benefiting from the modified heterointerface, P- $\text{Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$ exhibits ultralow potentials of 1.22, 1.30 and 1.39 V at 10, 100 and 1000 mA cm^{-2} , respectively, for the UOR. Furthermore, as a bifunctional electrocatalyst, P- $\text{Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$ requires only 1.70 V to drive 500 mA cm^{-2} in the UOR coupled HER test, which is approximately 500 mV lower than the OER//HER system. Importantly, a flow electrolyzer equipped with the P- $\text{Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$ electrode and supplied by urine can stably operate at 1000 mA cm^{-2} for over 180 h, further indicating its potential for energy-saving H_2 production and urine wastewater treatment.

2 Results and discussion

2.1 Synthesis and characterization

The P-doped $\text{Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$ heterojunction (P- $\text{Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$) was synthesized through a two-step process involving a hydrothermal reaction followed by phosphorization treatment, as illustrated in Fig. 1a. Initially, the $\text{Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$ precursor was prepared on nickel foam (NF) *via* a facile hydrothermal reaction, using NF as the Ni source and substrate and $\text{CoCl}_2 \cdot 6\text{H}_2\text{O}$ and thiourea as the Co and S sources, respectively. Fig. 1b shows the XRD pattern of the $\text{Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$ precursor. The diffraction peaks at 44.7° , 51.9° and 76.5° are attributed to the NF substrate (PDF no. 04-0850). Additionally, the remaining peaks can be indexed to hexagonal Ni_3S_2 (PDF no. 44-1418) and Co_9S_8 (PDF no. 02-1459), indicating the coexistence of Ni_3S_2 and Co_9S_8 phases after hydrothermal synthesis. The $\text{Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$ precursor exhibits a structure composed of interconnected nanosheets with a smooth surface, grown vertically on the NF substrate, as shown in the scanning electron microscopy (SEM) images (Fig. S1†). After phosphorization treatment, the diffraction peaks of the Ni_3S_2 and Co_9S_8 phases become more pronounced due to the annealing process at 400 °C for 2 h, while no diffraction peaks from P-related species are detectable (Fig. 1b). Fig. 1c displays that the framework of the nanosheets is basically maintained in P- $\text{Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$, but the surface becomes rough with the formation of many tiny nanosheets. This hierarchical structure significantly enhances the exposure of the effective surface, thereby promoting contact between reactants and active sites and improving the electrochemical performance.²⁷ During the phosphorization process, the doping level of P was controlled by using different amounts of NaH_2PO_2 (P content ranging from 5 to 30 mg). Fig. S2, S3,† and 1c present the XRD and SEM results of $\text{P}_{(5,15,20,30)}\text{-Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$. As $\text{P}_{(15)}\text{-Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$ exhibits the highest electrocatalytic activity, this sample is denoted as P- $\text{Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$ and used for the following detailed characterization. High-resolution transmission electron microscopy (HRTEM) images of P- $\text{Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$ (Fig. 1d and S4†)



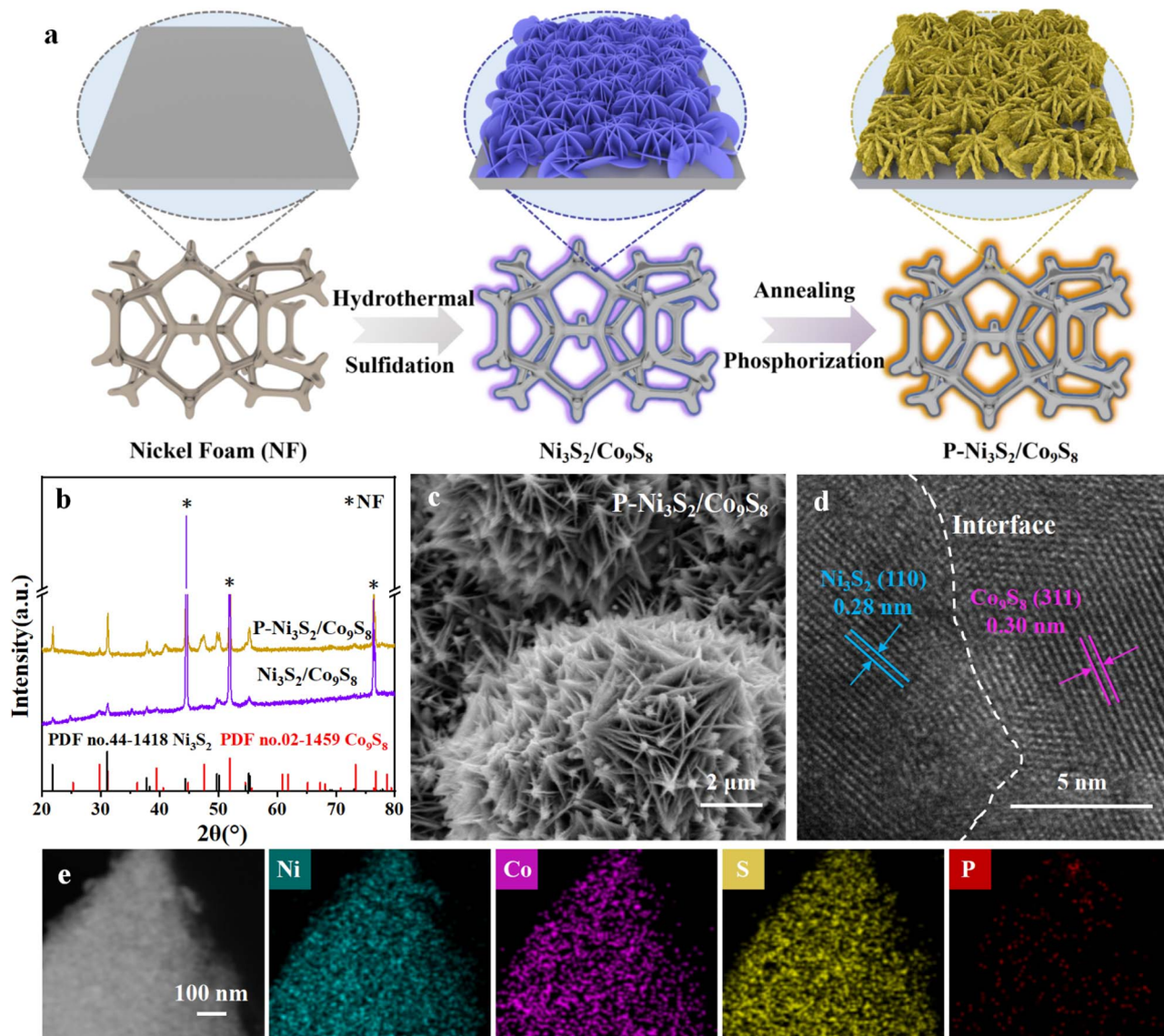


Fig. 1 (a) Schematic illustration of the synthetic procedure of P-Ni₃S₂/Co₉S₈. (b) XRD patterns of Ni₃S₂/Co₉S₈ and P-Ni₃S₂/Co₉S₈. (c) SEM image, (d) HRTEM image, and (e) HAADF-STEM image with the corresponding energy-dispersive spectrometry (EDS) mapping images of P-Ni₃S₂/Co₉S₈.

clearly resolve the lattice fringes with spacings of 0.28 and 0.30 nm, corresponding to the (110) plane of Ni₃S₂ and (311) plane of Co₉S₈, respectively, consistent with the XRD results. Additionally, a clear interface is observed between the adjacent Ni₃S₂ and Co₉S₈ regions. The high-angle annular dark-field scanning TEM (HAADF-STEM) and the corresponding energy-dispersive spectrometry (EDS) mapping images (Fig. 1e) display the homogeneous distribution of Co, Ni, S, and P elements throughout the nanosheet, with a P-doping amount of approximately 1.01 wt%. These results confirm the successful fabrication of the P-doped Ni₃S₂/Co₉S₈ heterojunction structure through phosphorization treatment of the Ni₃S₂/Co₉S₈ precursor. The surface chemical states and valence electron states of P-Ni₃S₂/Co₉S₈ and Ni₃S₂/Co₉S₈ were investigated by X-ray photoelectron spectroscopy (XPS). The XPS survey spectra (Fig. S5[†]) show the presence of P along with Ni, Co and S

elements in P-Ni₃S₂/Co₉S₈, further confirming the incorporation of P through phosphorization treatment. The P 2p spectrum of P-Ni₃S₂/Co₉S₈ presents two typical species corresponding to P-Co/Ni (130.3 and 129.4 eV) and P-O (133.3 eV)²⁸ (Fig. 2a). In the Ni 2p_{3/2} spectrum of P-Ni₃S₂/Co₉S₈, the peaks at 855.9 and 857.1 eV are assigned to Ni²⁺ and Ni³⁺, while the peak at 852.8 eV is attributed to the metallic Ni⁰ state.¹⁰ Similarly, the Co 2p_{3/2} spectrum exhibits three peaks at 778.0, 780.3, and 781.5 eV, corresponding to Co⁰, Co³⁺, and Co²⁺, respectively.²⁹ The peaks at 162.0 and 163.2 eV observed in the S 2p spectrum of P-Ni₃S₂/Co₉S₈ can be attributed to S 2p_{3/2} and S 2p_{1/2} of S²⁻ species in Co/Ni sulfide.³⁰ Compared with Ni₃S₂/Co₉S₈, the Ni 2p, Co 2p, and S 2p XPS spectra undergo shifts in binding energy after P-doping (Fig. 2b–d).³¹ Specifically, the Ni 2p peak shifts toward higher binding energy by about 0.3 eV, and the Co 2p peak shifts toward lower binding energy by about



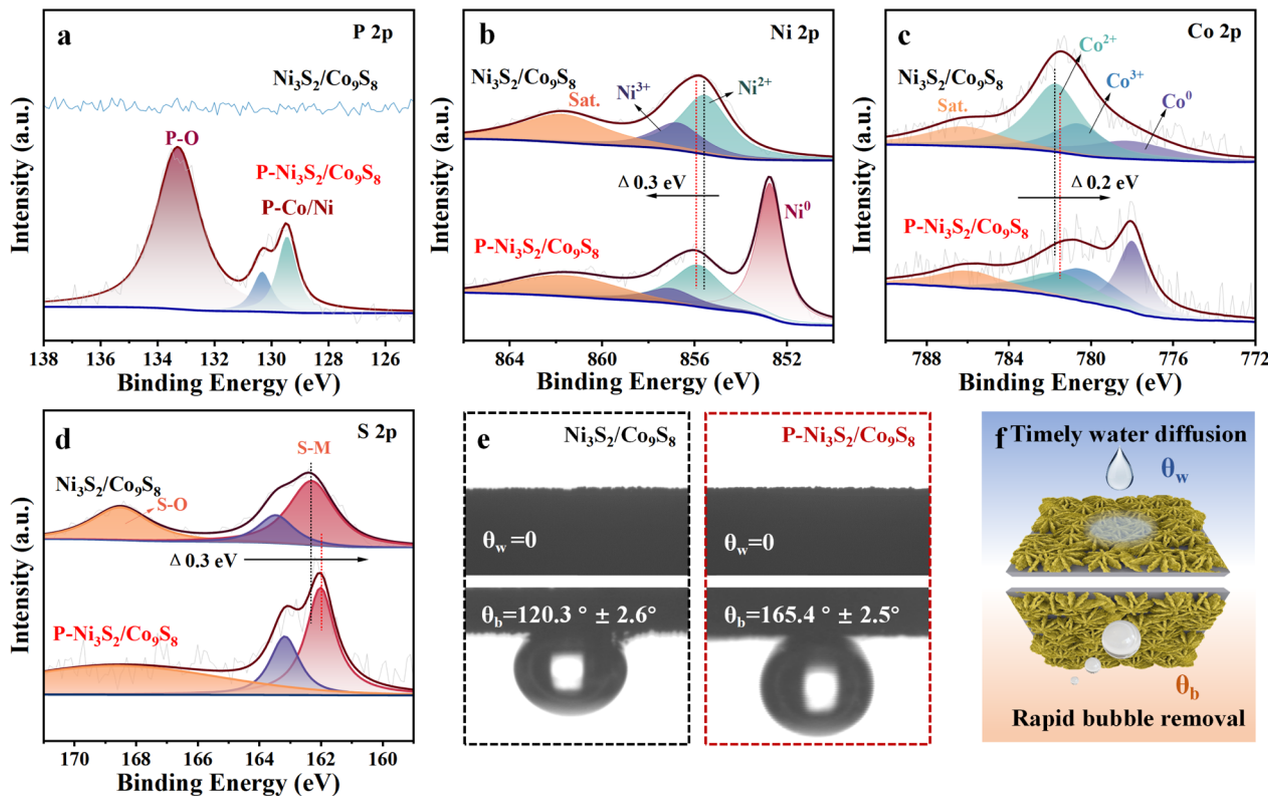


Fig. 2 XPS spectra of (a) P 2p, (b) Ni 2p, (c) Co 2p and (d) S 2p for $\text{Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$ and $\text{P-Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$. (e) Static water/bubble contact angle images of $\text{Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$ and $\text{P-Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$. (f) Schematic illustration of the superhydrophilic and superaerophobic properties of $\text{P-Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$.

0.2 eV. The binding energy in the S 2p spectrum is reduced by 0.3 eV. These shifts clearly indicate electron transfer from Ni_3S_2 to Co_9S_8 after P decoration, forming a Janus charge distribution at the $\text{P-Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$ heterointerface. These results further confirm the successful incorporation of P into $\text{Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$, which regulates the electronic environment at the heterojunction.

For electrocatalytic reactions proceeding in aqueous electrolyte with gaseous products, the hydrophilic/aerophobic properties of the electrode are essential for practical applications at industrial current densities.^{32,33} As shown in Fig. 2e, both $\text{Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$ and $\text{P-Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$ are superhydrophilic, with the static water contact angle (θ_w) close to 0° , enhancing the accessibility of the electrolyte to the catalyst surface during high-current electrochemical reactions. Specifically, the adhesion of gas bubbles to the $\text{P-Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$ surface is effectively reduced compared to $\text{Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$. The underwater gas bubble contact angle (θ_b) of $\text{P-Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$ ($165.4 \pm 2.5^\circ$) is significantly larger than that of $\text{Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$ ($120.3 \pm 2.6^\circ$) due to the rougher surface of $\text{P-Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$ after P doping. Moreover, the gas bubbles do not stably adhere to its surface during the test and are dispersed immediately upon contact, showing superaerophobicity.³⁴ The superhydrophilic/superaerophobic properties of $\text{P-Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$ not only ensure intimate contact between the reactants and the electrode surface but also facilitate the quick removal of gaseous products (Fig. 2f), which is critical for stable operation at high current densities.³⁵

2.2 Electrocatalytic performance for the UOR

The electrocatalytic performance of $\text{P-Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$ for the UOR was investigated in alkaline media with urea as the electrolyte using a standard three-electrode system. All potentials in the linear sweep voltammetry (LSV) polarization measurements were 85% *iR*-corrected and referenced to the reversible hydrogen electrode (RHE), unless otherwise stated. First, the effects of urea and KOH concentrations on the UOR performance were examined. The LSV curves of $\text{P-Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$ in Fig. S6a† indicate that the UOR performance is relatively insensitive to urea concentration. In contrast, the KOH concentration significantly influences the UOR performance of $\text{P-Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$, with the polarization current deteriorating as the KOH concentration decreases (Fig. S6b†). The performance remains high between 0.5 and 1.0 M KOH, which is advantageous for treating waste urea resources with varying urea concentrations. In this study, the electrolyte selected for UOR performance measurement was 1.0 M KOH with 0.33 M urea (an average concentration in urine). Subsequently, the influence of P doping level on the UOR performance was investigated, as shown in Fig. S7.† Clearly, the introduction of different amounts of P positively affects the catalyst kinetics of the UOR. Upon comparison, $\text{P-Ni}_3\text{S}_2/\text{Co}_9\text{S}_8$ with a 15 mg P-doped precursor exhibits the most outstanding UOR performance, characterized by the most significant increase in current and the lowest Tafel slope. Given that the OER is the primary competing reaction for the UOR, the LSV curves of the $\text{P-Ni}_3\text{S}_2/$



Co₉S₈ electrocatalyst for the UOR and OER are compared in Fig. 3a. For the OER measurement, the polarization curve was reversely swept to avoid the influence of Ni and Co oxidation.³⁶ The current density of the OER starts to increase at 1.45 V and reaches 500 mA cm⁻² at 1.65 V. In contrast, the LSV curve for the UOR shifts to a much lower potential range. The onset potential is as low as 0.97 V, and P-Ni₃S₂/Co₉S₈ displays a lower potential of 1.34 V to reach 500 mA cm⁻². Notably, P-Ni₃S₂/Co₉S₈ can deliver a high current density of 1000 mA cm⁻² at a potential of 1.39 V, while the anodic current of the OER is almost negligible at the same potential. These results indicate that the highly active P-Ni₃S₂/Co₉S₈ can maintain high UOR selectivity at large current densities, effectively mitigating the impact of OER competition. To demonstrate the advantage of P doping in

promoting UOR performance, the LSV curves of P-Ni₃S₂/Co₉S₈ and Ni₃S₂/Co₉S₈ for UOR are depicted in Fig. 3b. The UOR potentials required for P-Ni₃S₂/Co₉S₈ to reach 100 and 500 mA cm⁻² are merely 1.22 and 1.28 V, respectively, which are significantly lower than the 1.33 and 1.37 V required for Ni₃S₂/Co₉S₈. The NF substrate exhibits negligible activity toward the UOR. In addition, the Tafel slope of P-Ni₃S₂/Co₉S₈ is determined to be 57 mV dec⁻¹, lower than that of the control electrocatalyst (Fig. 3c). The low Tafel slope is also well supported by the observed rapid increase in current density, indicating fast reaction kinetics toward the UOR. The electrochemical surface areas (ECSAs) of P-Ni₃S₂/Co₉S₈ and Ni₃S₂/Co₉S₈ were evaluated by calculating their electrochemical double-layer capacitance (*C*_{dl}) from the non-faradaic regions of cyclic voltammetry (CV)

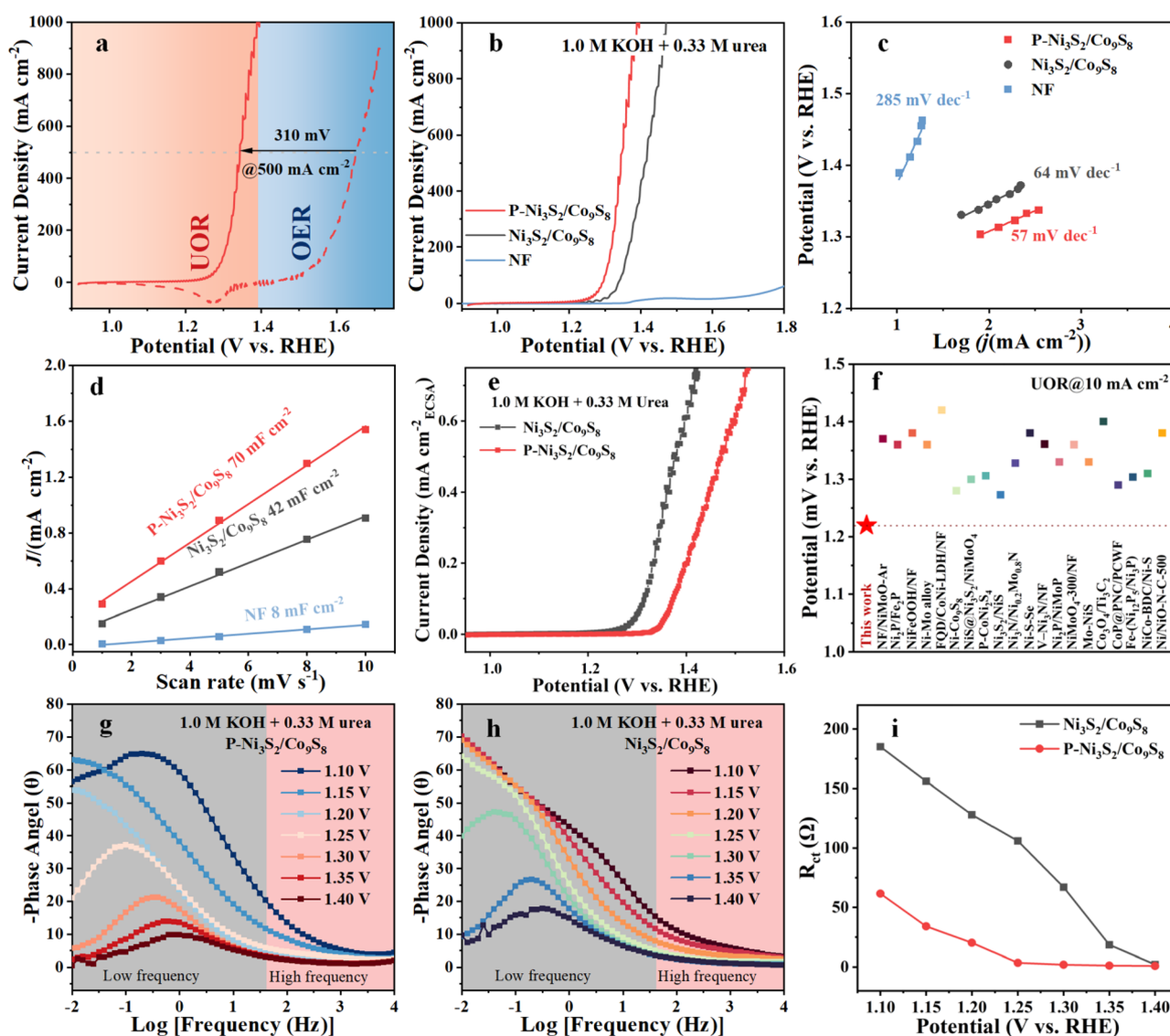


Fig. 3 Anodic oxidation performance. (a) LSV curves of P-Ni₃S₂/Co₉S₈ for the UOR (1.0 M KOH with 0.33 M urea) and the OER (1.0 M KOH). (b) LSV curves, (c) Tafel slopes, and (d) *C*_{dl} values of P-Ni₃S₂/Co₉S₈, Ni₃S₂/Co₉S₈, and NF during the UOR. (e) ECSA-normalized LSV curves of P-Ni₃S₂/Co₉S₈, Ni₃S₂/Co₉S₈, and NF during the UOR. (f) Comparison of the overpotential for the UOR of our catalyst and other reported Ni/Co-based electrocatalysts. Bode phase plots of (g) P-Ni₃S₂/Co₉S₈ and (h) Ni₃S₂/Co₉S₈ at different potentials during the UOR. (i) Fitted *R*_{ct} values of P-Ni₃S₂/Co₉S₈ and Ni₃S₂/Co₉S₈.



curves (Fig. S8† and 3d). The C_{dl} value of P-Ni₃S₂/Co₉S₈ is 1.7 and 8.8 times higher than those of Ni₃S₂/Co₉S₈ and NF, respectively, implying the positive effect of P doping on increasing the ECSA. Moreover, the higher ECSA-normalized current density of P-Ni₃S₂/Co₉S₈ in Fig. 3e further demonstrates its superior intrinsic activity. The turnover frequency (TOF) was employed to evaluate the intrinsic catalytic activity. The number of active sites was calculated based on the charge capacity obtained by integrating the CV curves (Fig. S9†). Fig. S10† shows that the TOF values of P-Ni₃S₂/Co₉S₈ consistently exceed those of Ni₃S₂/Co₉S₈. The high TOF of 8.47 s⁻¹, is gained for P-Ni₃S₂/Co₉S₈ at 1.4 V, which greatly surpasses that of the Ni₃S₂/Co₉S₈ heterojunction (3.69 s⁻¹), further confirming the enhanced UOR intrinsic activity after P doping. The excellent UOR performance of P-Ni₃S₂/Co₉S₈ surpasses those of the state-of-the-art Co/Ni-based electrocatalysts (Fig. 3f and Table S1†).

To investigate the dynamic evolution of the electrocatalyst interface during the UOR, *in situ* electrochemical impedance spectroscopy (EIS) measurements were performed with the applied potential gradually increased from 1.10 to 1.40 V.³⁷ Fig. 3g and h show the Bode plots of P-Ni₃S₂/Co₉S₈ and Ni₃S₂/Co₉S₈. An apparent phase peak at 1.25 V in the low-frequency region (0.1 Hz) for P-Ni₃S₂/Co₉S₈ implies the occurrence of a charge transfer process associated with the UOR.³⁸ As the applied potential increases, the intensity of the phase angle continuously decreases, and the position of the peak gradually shifts towards higher frequencies. The position of the peak reflects the time domain of the charge transfer process, which is related to the kinetics of the electrochemical reaction.³⁹ The phase angle of P-Ni₃S₂/Co₉S₈ shifts more rapidly towards the high-frequency region compared to Ni₃S₂/Co₉S₈, indicating the faster catalytic UOR kinetics after P introduction. As displayed in Fig. S11,† the Nyquist plots of P-Ni₃S₂/Co₉S₈ are steep lines at low applied potentials (<1.25 V), indicating extremely high charge transfer resistances and that the UOR has not yet occurred. The Nyquist plot begins to deviate from the imaginary part after 1.25 V, which is associated with the charge transfer of the UOR process, consistent with the corresponding Bode plots. As the applied potential increases to 1.30 V, significant mutation occurs in the Nyquist plot of P-Ni₃S₂/Co₉S₈, and a complete small semicircle appears, indicating the start of the UOR.³⁹ In contrast, the Nyquist plot of Ni₃S₂/Co₉S₈ changes at 1.30 V, and a complete semicircle is observed at 1.35 V. Consequently, according to the equivalent circuit (Fig. S12†), Ni₃S₂/Co₉S₈ shows a later decrease in R_{ct} (charge transfer resistance) compared to P-Ni₃S₂/Co₉S₈ (Fig. 3i), implying faster polarization and oxidation behavior of the UOR on the surface of P-Ni₃S₂/Co₉S₈.

In light of the UOR electrocatalytic mechanism, the CV curves of P-Ni₃S₂/Co₉S₈ for the UOR and OER were compared (Fig. 4a). In the CV curve for the OER (1.0 M KOH), a broad oxidation peak is observed at 1.3–1.4 V, corresponding to the oxidation reaction of M²⁺ (M = Co/Ni). Subsequently, a reduction reaction peak of M³⁺ appears during the backward sweep. These observations suggest that P-Ni₃S₂/Co₉S₈ undergoes pre-oxidation to produce catalytic high-valence species during the OER process, which has been reported for other Co/Ni-based

electrocatalysts.⁴⁰ Upon the addition of urea, the oxidation peak disappears, and the current density increases rapidly at the location where the M²⁺ oxidation begins. Following this, a reduction peak appears during the reverse sweep, indicating that the high-valence M³⁺ probably also act as the active species in the UOR process.⁴¹ Given the similar active sites in the OER and UOR, the adsorption and activation of OH⁻, which is the first step in the OER (OH⁻ + * → OH_{ads} + e⁻),⁴² also plays a crucial role in the UOR in alkaline solutions. Therefore, the adsorption and activation behaviors were detected by evaluating their OER performance. The LSV curves in Fig. 4b show that P-Ni₃S₂/Co₉S₈ exhibits superior OER performance with a higher polarization current than that of Ni₃S₂/Co₉S₈. Moreover, in a low OH⁻ concentration environment (Fig. S13†), P-Ni₃S₂/Co₉S₈ also displays better OER performance. These results imply that the introduction of P could facilitate the effective capture of OH⁻ on the P-Ni₃S₂/Co₉S₈ surface. Furthermore, changes in the active sites at the electrode–solvent interface were monitored by open-circuit potential (OCP) measurement to evaluate the urea adsorption behavior on the catalyst surface. When 0.33 M urea was injected, the potential equilibrium was disrupted due to the exchange of urea and ions on the catalyst surface, resulting in a significant change in OCP.⁴³ In Fig. 4c, the observed change in OCP for P-Ni₃S₂/Co₉S₈ (0.38 V) is more pronounced compared to Ni₃S₂/Co₉S₈ (0.33 V), confirming that more urea molecules are adsorbed to the Helmholtz layer of the P-Ni₃S₂/Co₉S₈ electrode. The faster decrease in OCP for P-Ni₃S₂/Co₉S₈ upon the addition of urea (inset in Fig. 4c) further provides the evidence of the promoting effect of P on dynamic urea adsorption.⁴⁴

To further investigate the adsorption behavior of Ni₃S₂/Co₉S₈ and P-Ni₃S₂/Co₉S₈ towards the two functional groups (–NH₂ and C=O) of the urea molecule, urea analogues (acetamide, thiourea and ethylenediamine) with similar structural groups were used for comparison. As shown in Fig. 4d–f, the LSV curves for Ni₃S₂/Co₉S₈ and P-Ni₃S₂/Co₉S₈ in the acetamide electrocatalytic oxidation reaction (AER), thiourea electrocatalytic oxidation reaction (TER), and ethylenediamine electrocatalytic oxidation reaction (EER) were shifted to higher potentials compared to the UOR, indicating a decrease in performance. In particular, the degradation caused by the –NH₂ alteration (AER) was much more pronounced than that caused by the C=O alteration (TER). P-Ni₃S₂/Co₉S₈ showed the least degradation performance in the EER (Fig. 4f), which has two –NH₂ groups like the urea molecule, but Ni₃S₂/Co₉S₈ showed significant degradation. As displayed in Fig. S14,† the results of the OCP tests also showed that P-Ni₃S₂/Co₉S₈ has a significantly lower adsorption strength in the absence of the –NH₂ group (AER). Compared to Ni₃S₂/Co₉S₈, P-Ni₃S₂/Co₉S₈ exhibits higher –NH₂ adsorption (TER and EER). These results suggest that the improved performance of P-Ni₃S₂/Co₉S₈ is likely due to its enhanced adsorption strength of –NH₂ during the oxidation reaction. In parallel, programmed temperature rise desorption (TPD) tests were conducted on P-Ni₃S₂/Co₉S₈ and Ni₃S₂/Co₉S₈ to investigate the adsorption behavior towards different groups of urea molecules. In a butylamine/He atmosphere, the desorption temperature for P-Ni₃S₂/Co₉S₈ is higher than that for Ni₃S₂/Co₉S₈, demonstrating stronger adsorption of P-Ni₃S₂/Co₉S₈ for the –NH₂ group



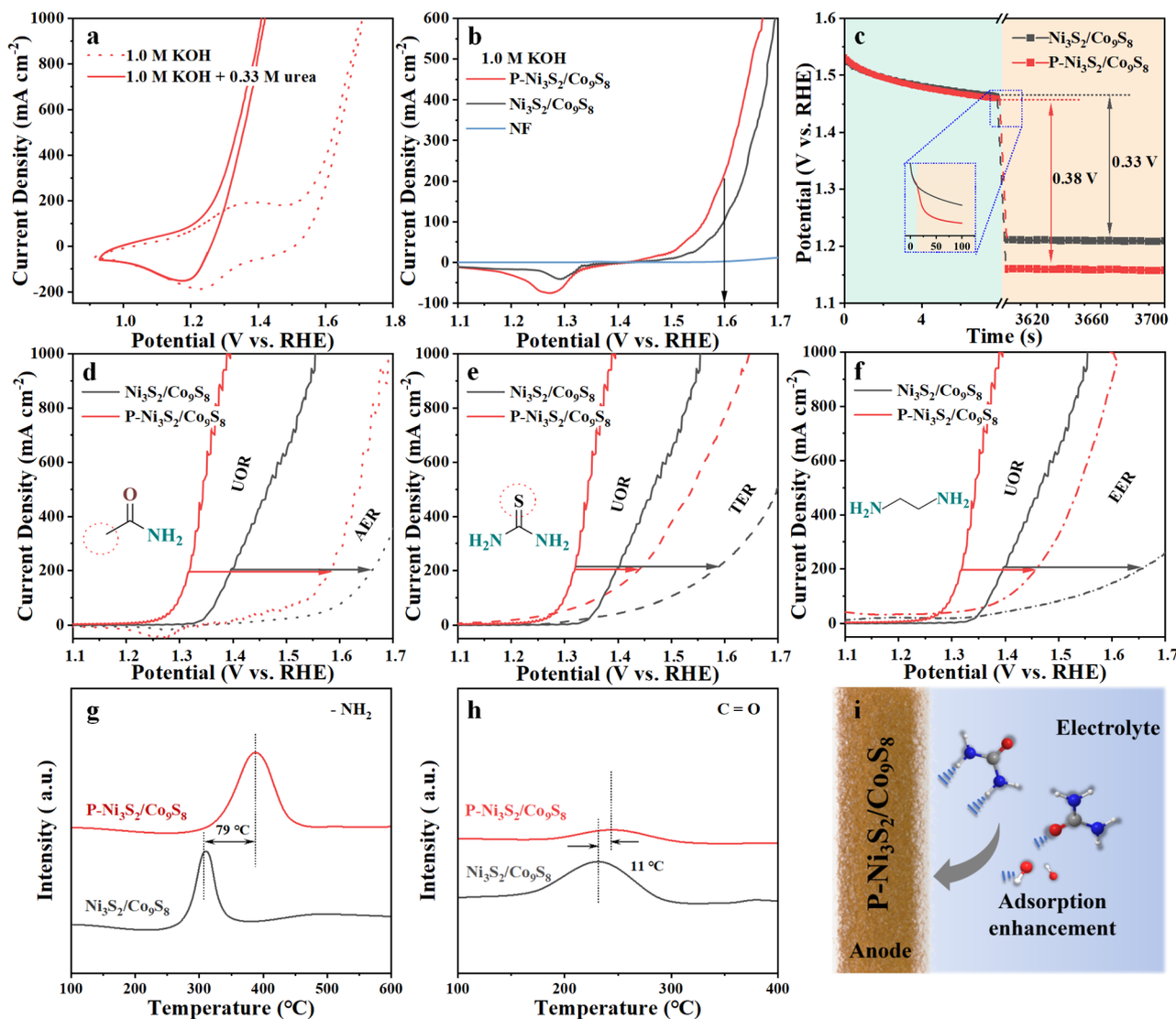


Fig. 4 (a) CV curves of P-Ni₃S₂/Co₉S₈ during the UOR and OER. (b) Backswept LSV curves of P-Ni₃S₂/Co₉S₈, Ni₃S₂/Co₉S₈, and NF in 1.0 M KOH. (c) OCP curves of P-Ni₃S₂/Co₉S₈ and Ni₃S₂/Co₉S₈ in 1.0 M KOH before and after 0.33 M urea was injected. LSV curves of P-Ni₃S₂/Co₉S₈ and Ni₃S₂/Co₉S₈ in 1.0 M KOH with 0.33 M urea or with (d) acetamide, (e) thiourea and (f) ethylenediamine. TPD spectra of P-Ni₃S₂/Co₉S₈ and Ni₃S₂/Co₉S₈ in (g) butylamine/He and (h) CO atmospheres. (i) Schematic diagram of the initial reaction in the inner Helmholtz layer.

(Fig. 4g). The desorption temperature of P-Ni₃S₂/Co₉S₈ for urea molecules in the CO-TPD test was similar to that of Ni₃S₂/Co₉S₈ (Fig. 4h), but the desorption temperature of P-Ni₃S₂/Co₉S₈ was slightly higher than that of Ni₃S₂/Co₉S₈. These results suggest that the enhanced urea adsorption should be mainly attributed to the -NH₂ group, which is consistent with the LSV test results for urea analogue substrates (Fig. 4i).

The surface electronic structures and chemical states of P-Ni₃S₂/Co₉S₈ were investigated using XPS after the OER and UOR tests (Fig. S15[†]). After the OER test, the peak intensity of the P/S element is considerably decreased, and the signals of P/S-Co/Ni disappear. Compared to the initial P-Ni₃S₂/Co₉S₈, the surface concentrations of Ni^{0/2+} and Co^{0/2+} species are significantly lower after the OER test, while the ratios of Ni³⁺ and Co³⁺ increase. This suggests that the Ni^{0/2+} and Co^{0/2+} species were oxidized to Ni³⁺ and Co³⁺, respectively.⁴⁵ Furthermore, the O 1s

spectrum reveals that the characteristic peaks corresponding to O-MO_x species become dominant after the OER test, confirming the reconstruction of the catalyst surface into metal oxyhydroxide species. In contrast, the XPS results after the UOR test are significantly different from those after the OER test. The proportion of P/S-O characteristic peaks increase after the UOR test, although a certain proportion of P/S-Co/Ni remains detectable, implying partial surface oxidation on the catalyst. The peaks corresponding to Ni^{0/3+} species disappear, and only Ni²⁺ species are present. For the Co element, the Co^{3+/2+} ratio does not change significantly. Similarly, the O 1s spectrum exhibits no significant changes. Based on these results, it is proposed that the proton in the urea molecule may fill the hydrogen defect in the intermediate metal hydroxyl species, thereby avoiding a phase transition to metal oxyhydroxide. This finding is



consistent with the previously reported nucleophilic oxidation mechanism mediated by metal redox pairs (M^{2+}/M^{3+}).⁴⁶

Density functional theory (DFT) calculations were performed to elucidate the effectiveness of P decoration in enhancing the UOR performance of the P-Ni₃S₂/Co₉S₈ heterostructure. Based on experimental results and theoretical calculations, the optimized model for P-Ni₃S₂/Co₉S₈, where surface S atoms are partially replaced by P atoms, was employed for the following DFT study (Fig. S16†).^{47,48} The Ni₃S₂/Co₉S₈ model is similar to that of P-Ni₃S₂/Co₉S₈, except for the absence of P atom substitution. The effect of P incorporation on the charge distribution at the Ni₃S₂/Co₉S₈ heterointerface was first investigated *via* analyzing the charge density difference. Fig. 5a and b indicate that electrons transfer from Ni₃S₂ to Co₉S₈. Upon P incorporation, the charge distribution significantly increases from 0.08 e⁻ (Ni₃S₂/Co₉S₈) to 0.28 e⁻ (P-Ni₃S₂/Co₉S₈), forming a Janus charge distribution interface with electrophilic P-Ni₃S₂ and nucleophilic P-Co₉S₈. This Janus interface facilitates the selective adsorption of urea's electron-withdrawing group (C=O) and electron-donating group (amino). For heterogeneous electrocatalysis, the adsorption behaviors of reactants on electrocatalysts typically play a crucial role in affecting the corresponding reaction processes. To determine the adsorption behaviors of urea, all possible NN-terminal and NO-terminal adsorption configurations at the interfaces of Ni₃S₂/Co₉S₈ and P-Ni₃S₂/Co₉S₈ were considered (Fig. S17†). As compared in

Fig. 5c, the stable adsorptions for each sample were screened according to the calculated adsorption energies. Accordingly, urea tends to adsorb as a NN-terminal configuration with an adsorption energy of -2.07 eV at the interface of Ni₃S₂/Co₉S₈. For P-Ni₃S₂/Co₉S₈, a NO-terminal configuration (N adsorbed on the Ni site and O adsorbed on the Co site) with a more negative adsorption energy of -2.23 eV is favorable, confirming enhanced urea adsorption upon P incorporation. Additionally, the local charge density difference for Ni₃S₂/Co₉S₈ and P-Ni₃S₂/Co₉S₈ adsorbed urea was calculated, and the results are shown in Fig. 5d. Upon Bader charge analysis, the P-decorated hetero-junction displays a larger charge density difference of 0.13 e⁻ compared to 0.11 e⁻ for Ni₃S₂/Co₉S₈. Thus, the incorporation of P not only increases the charge density difference but also promotes electron transfer from the adsorbed urea molecule to the P-Ni₃S₂/Co₉S₈ interface, which is crucial for enhancing urea adsorption. Thus, the incorporation of P favors electron transfer from the adsorbed urea molecule to P-Ni₃S₂/Co₉S₈, thereby enhancing urea adsorption. To further quantitatively investigate the bonding strengths between active sites and the adsorbed urea, crystal orbital Hamiltonian population (COHP) analysis was carried out.⁴⁹ Fig. 5e-h compare the integrated COHP (ICOHP) calculated by integrating COHP over all levels up to the Fermi level. The ICOHP for P-Ni₃S₂/Co₉S₈ (-1.64 and -0.94 eV) is more negative compared to Ni₃S₂/Co₉S₈ (-1.30 and -0.04 eV), further confirming a stronger coupling between the

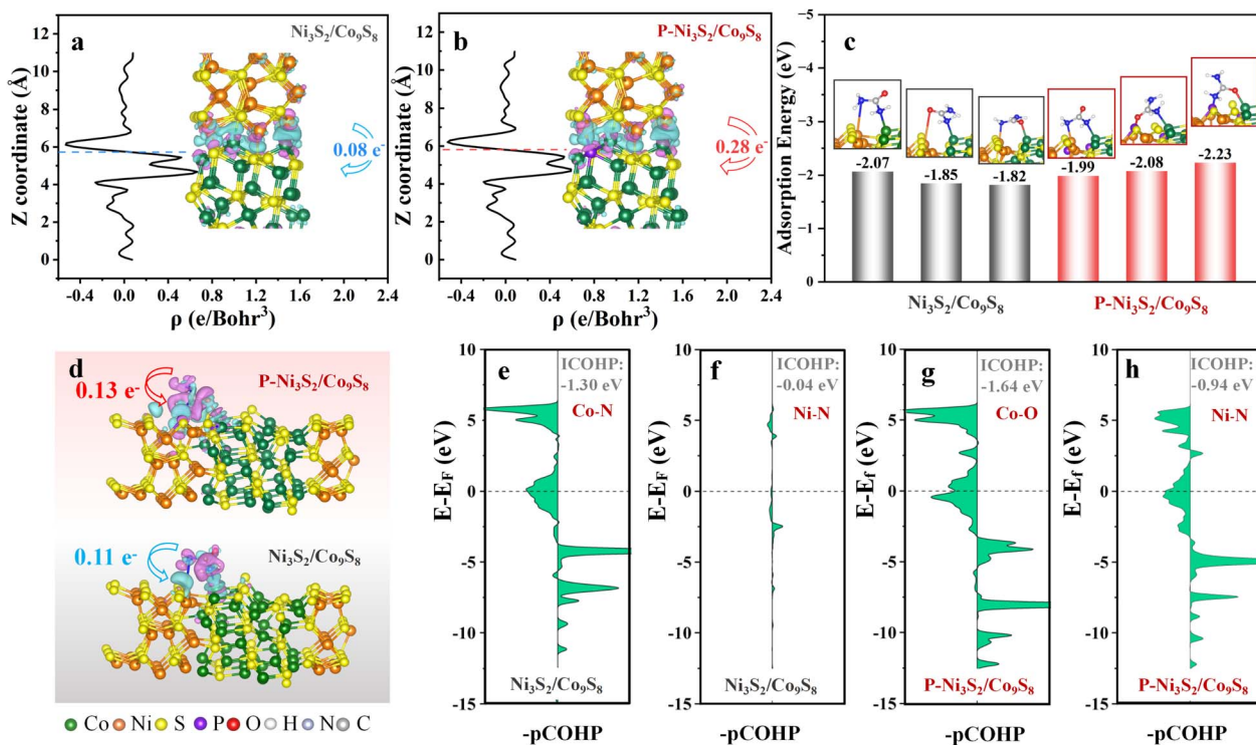


Fig. 5 Average charge density difference of (a) Ni₃S₂/Co₉S₈ and (b) P-Ni₃S₂/Co₉S₈. (c) Three optimized adsorption structures and adsorption energies for CO(NH₂)₂ on Ni₃S₂/Co₉S₈ and P-Ni₃S₂/Co₉S₈. (d) The difference charge density and Bader charge analysis for CO(NH₂)₂ on Ni₃S₂/Co₉S₈ and P-Ni₃S₂/Co₉S₈, where the isosurface value is set to 0.0015 e Å⁻³ and the positive and negative charges are shown in cyan and magenta, respectively. The green, orange, yellow, purple, red, white, blue, and gray balls represent Co, Ni, S, P, O, H, N and C atoms, respectively. Projected crystal orbital Hamiltonian population (pCOHP) analysis between CO(NH₂)₂ and (e) and (f) Ni₃S₂/Co₉S₈ and (g) and (h) P-Ni₃S₂/Co₉S₈.



transition metal (Ni/Co) and urea (N/O) orbitals upon P incorporation. This stronger orbital coupling, induced by P incorporation, directly contributes to the enhanced adsorption and activation of urea molecules on the heterojunction surface. All these results indicate that the incorporation of P leads to modification in the electronic structure of the heterojunction, thereby enhancing adsorption and promoting UOR activity.

2.3 Electrocatalytic performance for the HER

In addition to its remarkable UOR activity, the electrocatalytic HER performance of the P-Ni₃S₂/Co₉S₈ was also evaluated in an alkaline electrolyte (1.0 M KOH) using a three-electrode system. The LSV curves in Fig. 6a show that the potential of P-Ni₃S₂/Co₉S₈ is significantly lower than that of Ni₃S₂/Co₉S₈ in the current density range of 0–1000 mA cm⁻². Specifically, to achieve current densities of 10, 100, and 500 mA cm⁻², P-Ni₃S₂/Co₉S₈ requires overpotentials of 63, 162, and 192 mV, respectively, which are much lower than those of Ni₃S₂/Co₉S₈ (88, 232, and 313 mV, Fig. 6b). Meanwhile, P-Ni₃S₂/Co₉S₈ shows a significantly lower Tafel slope of 85 mV dec⁻¹ compared to 116 mV dec⁻¹ for Ni₃S₂/Co₉S₈ (Fig. 6c), indicating faster HER kinetics on the surface of P-Ni₃S₂/Co₉S₈. Meanwhile, a high TOF value of 7.08 s⁻¹ can be achieved at -0.2 V for P-Ni₃S₂/Co₉S₈ (Fig. S18†), which is 4.9 times higher than that of Ni₃S₂/Co₉S₈, demonstrating its exceptionally high intrinsic catalytic activity. The

HER performance of P-Ni₃S₂/Co₉S₈ exhibited a satisfactory level. Compared with reported Ni-based catalysts, although it did not reach the optimum, it was on a par with the upper-middle range (Table S1†). Furthermore, *in situ* EIS was performed to further probe the alkaline HER kinetics. As shown in Fig. 6d and e, the Bode plots for P-Ni₃S₂/Co₉S₈ and Ni₃S₂/Co₉S₈ show similar trends in phase angles, while P-Ni₃S₂/Co₉S₈ exhibits a greater downward trend in the phase angle response due to more efficient electron transfer (Fig. 6f).⁵⁰ The Nyquist plots of P-Ni₃S₂/Co₉S₈ and Ni₃S₂/Co₉S₈ at potentials of 0.2–0.45 V show a gradual shift from a steep line to a semicircle (-0.2 V) as the applied potential increases (Fig. S19†). Subsequently, the semicircle diameter becomes progressively smaller as the potential increases, suggesting that the reaction rate increases. P-Ni₃S₂/Co₉S₈ consistently has a smaller semicircular diameter than Ni₃S₂/Co₉S₈ at the same potential, further confirming the faster electron transfer at the P-Ni₃S₂/Co₉S₈ interface. Consequently, the introduction of P in the heterojunction structure enhances electron transfer and accelerates the kinetics of the HER. The HER activity of P-Ni₃S₂/Co₉S₈ was also evaluated in a 1.0 M KOH solution containing urea to verify the feasibility of UOR as an alternative to the OER for coupling with the HER. As shown in Fig. S20,† the nearly overlapping LSV curves indicate that urea has a negligible effect on the HER performance of P-Ni₃S₂/Co₉S₈, ensuring its potential application in the energy-saving UOR coupled H₂ production.

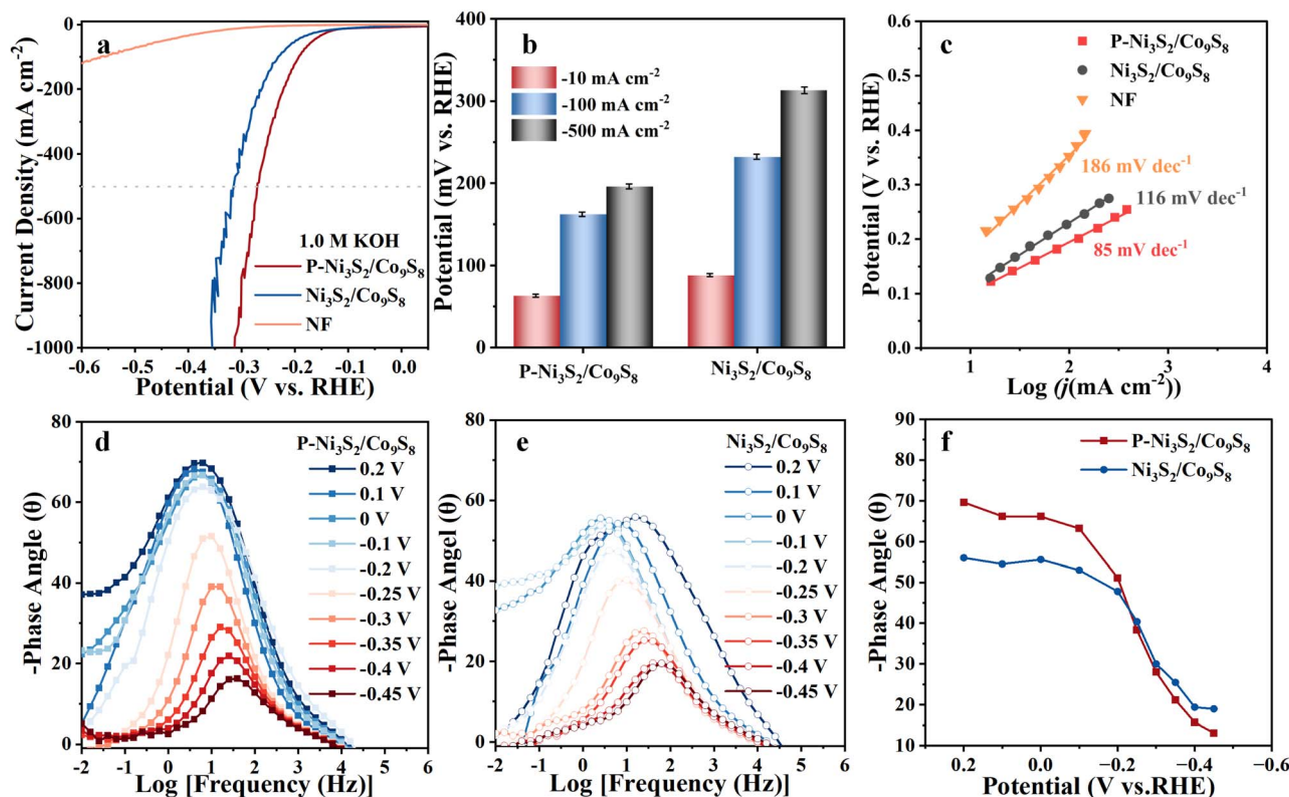


Fig. 6 Electrocatalytic HER performance. (a) LSV curves of P-Ni₃S₂/Co₉S₈, Ni₃S₂/Co₉S₈, and NF in 1.0 M KOH. (b) Comparison of potentials at different current densities between P-Ni₃S₂/Co₉S₈ and Ni₃S₂/Co₉S₈. (c) Tafel slopes of P-Ni₃S₂/Co₉S₈, Ni₃S₂/Co₉S₈, and NF for the HER. *In situ* EIS characterization. Bode phase plots of (d) P-Ni₃S₂/Co₉S₈ and (e) Ni₃S₂/Co₉S₈ at different potentials during the HER. (f) Corresponding response of the phase angle to the applied potential.



2.4 Electrocatalytic activity toward overall flow-mode urea electrolysis

Considering the outstanding bifunctional electrocatalytic activities of P-Ni₃S₂/Co₉S₈ towards the UOR and HER, we constructed a paired UOR//HER two-electrode cell, employing P-Ni₃S₂/Co₉S₈ as both the anode and cathode (Fig. S21†). As depicted in Fig. S21b,† the UOR//HER system in 1.0 M KOH containing 0.33 M urea exhibits a significantly lower onset potential and a more rapid current increase compared to the OER//HER system in 1.0 M KOH. Specifically, to achieve a high current density of 500 mA cm⁻², the UOR//HER system requires a potential of only 1.70 V, approximately 500 mV lower than that of the OER//HER system. These results underscore the potential of the bifunctional P-Ni₃S₂/Co₉S₈ for urea-assisted electrolysis as an energy-saving H₂ production pathway, potentially replacing conventional water electrolysis. Inspiringly, to further evaluate the practicality of P-Ni₃S₂/Co₉S₈ for urea-assisted H₂ production, an anion exchange membrane (AEM) flow electrolyzer was assembled, effectively enhancing mass transfer and reducing concentration polarization (Fig. 7a and S22†). When using artificial urine (with pH adjusted to 14) as the electrolyte, the LSV curve may fluctuate due to the presence of impurities such as Cl⁻, Mg²⁺, and Ca²⁺. The flow-mode UOR//HER device operates efficiently, demonstrating a significantly reduced cell voltage compared to the OER//HER (Fig. 7b). At 25 °C and a high current density of 1000 mA cm⁻², the flow-mode UOR//HER

device achieves a 34.5 ± 0.98% reduction in electricity consumption relative to the OER//HER system, demonstrating the effectiveness of the UOR//HER system in minimizing energy expenditure.⁵¹ The stability of an electrocatalyst under high current density is a crucial metric for evaluating in practical applications. As illustrated in Fig. 7c, the P-Ni₃S₂/Co₉S₈ catalyst exhibits excellent stability across different temperatures. At 25 °C, it maintains 92.1% of its initial voltage after 180 hours of continuous operation at 1000 mA cm⁻². Initially, a slight voltage increase is observed, which can be attributed to pre-oxidation effects.⁵² Subsequently, the voltage increases gradually, likely due to the gradual depletion of urea concentration and minor influences from urine components on the catalyst. However, these factors have a minimal impact on the overall performance. Notably, the degradation rate of 0.044% corresponding to the 92.1% voltage retention is significantly better than those reported in several literature references (Table S2†), where higher degradation rates are typically observed under similar conditions. Furthermore, the XRD pattern, SEM image, and EDS spectrum of P-Ni₃S₂/Co₉S₈ following the durability test exhibit negligible variation in morphology, phase, and composition (Fig. S23 and S24†). The potential leaching of Ni and Co from the P-Ni₃S₂/Co₉S₈ catalyst into the electrolyte was detected with ICP-AES. Following a 180 h durability test, the amount of dissolved Ni and Co was found to be less than 1.0 μg cm⁻² (corresponding to the geometric area of the electrode, Table S3†).

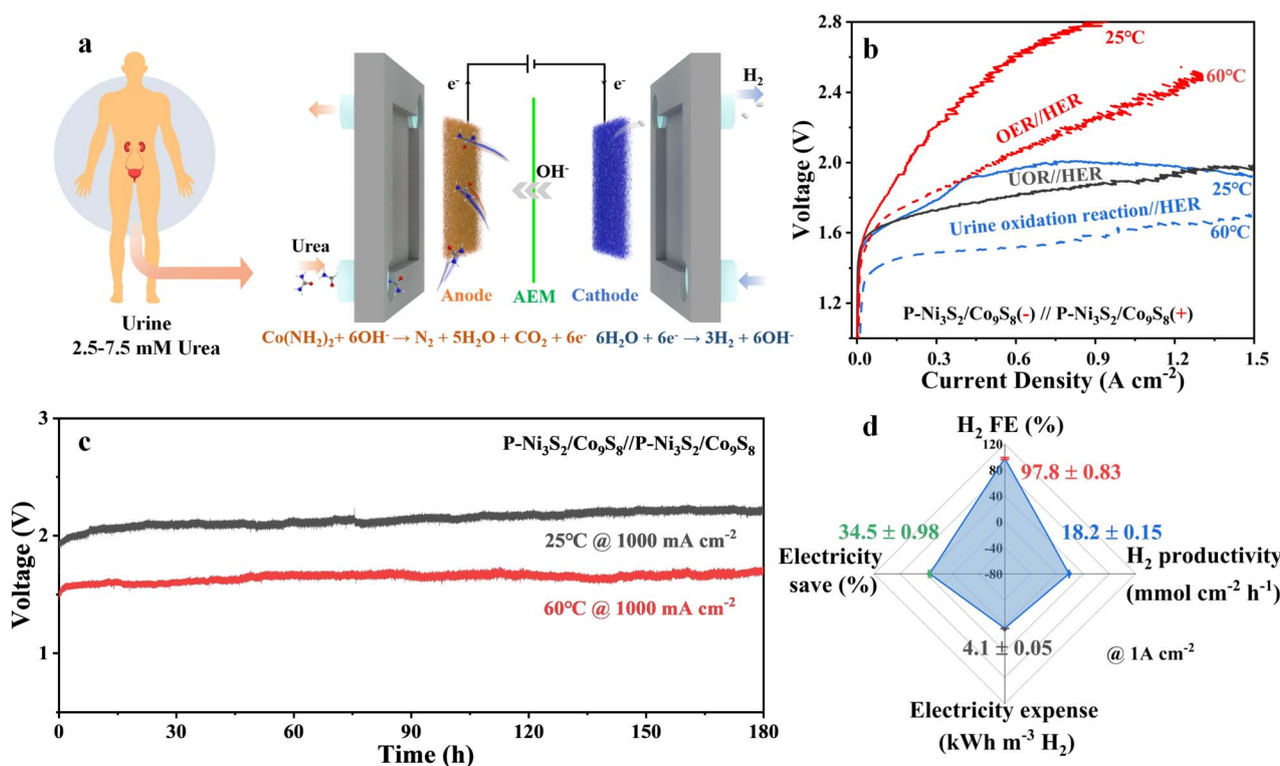


Fig. 7 Urea-assisted electrolysis performance in an AEM electrolyzer. (a) Schematic illustration of the urea electrolyzer using P-Ni₃S₂/Co₉S₈ as both the anode and cathode. (b) Polarization curves of P-Ni₃S₂/Co₉S₈(-)//P-Ni₃S₂/Co₉S₈(+). (c) Durability cell voltage–time plots for P-Ni₃S₂/Co₉S₈(-)//P-Ni₃S₂/Co₉S₈(+) at a constant current density of 1000 mA cm⁻² (the inset shows the photographs of the AEM electrolyzer). (d) The comprehensive performance of P-Ni₃S₂/Co₉S₈.



This value is significantly lower than the Ni and Co amounts in the pristine P-Ni₃S₂/Co₉S₈, thereby effectively ruling out the possibility of substantial dissolution of these metals from the catalyst. These findings reliably demonstrate that P-Ni₃S₂/Co₉S₈ possesses outstanding stability for the UOR. The faradaic efficiency (FE) of H₂ production was monitored during the stability test by periodically comparing the amount of theoretical and measured H₂ (Fig. S25†). The FE for H₂ is maintained at a high level (97.8 ± 0.83%), with a productivity of 18.2 ± 0.15 mmol cm⁻² h⁻¹, accompanied by a low energy consumption of 4.1 ± 0.05 kWh m⁻³ H₂ (Fig. 7d). More impressively, when the temperature is increased to 60 °C, the catalyst not only exhibits remarkable enhancement in performance but also maintains outstanding stability, achieving a voltage retention of 93.5%. Gas chromatography (GC) analysis was conducted on the anode gaseous products at different time intervals. The results reveal the generation of N₂ and O₂ during the UOR process (Fig. S26 and Table S4†). Here, O₂ is produced through the competitive OER. Notably, N₂ is identified as the predominant product, exhibiting a high FE of 71.29 ± 1.67%, while the FE of O₂ is only 0.51 ± 0.02%. This indicates that the majority of electrons are utilized for catalyzing urea oxidation rather than the OER. Analysis of the anolyte by ion chromatography (IC) reveals that the primary liquid product is NO₂⁻ (Fig. S27†), with a FE of 37.52 ± 1.90%. Therefore, taking all factors into account, after the long-term stability test, the main products from the UOR were N₂ and NO₂⁻.⁵³ Overall, given its remarkable activity and stability, the P-Ni₃S₂/Co₉S₈ electrocatalyst holds great promise for industrial application in energy-saving H₂ production *via* urea-assisted electrolysis.

3 Conclusions

In summary, a P-modified Ni₃S₂/Co₉S₈ heterostructure with a Janus charge distribution interface is constructed as an efficient electrocatalyst for urea-assisted H₂ production. The incorporation of P effectively enhances the electron transfer from Ni₃S₂ to Co₉S₈ at the interface. The resultant electrophilic P-Ni₃S₂ and nucleophilic P-Co₉S₈ domains facilitate the adsorption of electron-donating amino and electron-withdrawing carbonyl groups of urea, thereby modulating the adsorption energy and interaction. Consequently, the prepared P-Ni₃S₂/Co₉S₈ exhibits enhanced UOR performance with low potentials of 1.22, 1.30, and 1.39 V (*vs.* RHE) at current densities of 10, 100, and 1000 mA cm⁻², respectively. As a bifunctional electrocatalyst, P-Ni₃S₂/Co₉S₈ also displays excellent HER activity, requiring a minimal potential of 65 mV at 10 mA cm⁻². Accordingly, an AEM flow electrolyzer equipped with this bifunctional P-Ni₃S₂/Co₉S₈ and supplied with alkaline urine enables stable H₂ production at 1000 mA cm⁻² for 180 h. Compared to conventional water splitting, the UOR//HER system achieves a satisfactory H₂ productivity of 18.2 mmol cm⁻² h⁻¹ with 34.5% electricity saving. This work presents a powerful strategy for activating the Janus charge distribution at heterojunction interfaces, thereby optimizing the adsorption behaviors of reactants and improving the performance.

Data availability

All data can be found in the main article or the ESI.†

Author contributions

C. W. (funding acquisition: lead; resources: lead; writing – review & editing: equal); H. L. (funding acquisition: equal; resources: equal; validation: equal; writing – review & editing: equal); Y. S. (conceptualization: lead; data curation: lead; investigation: lead; methodology: lead; project administration: supporting; validation: lead; visualization: lead; writing – original draft: lead); X. Z. (data curation: supporting; investigation: supporting; validation: supporting); H. G. (data curation: supporting; formal analysis: supporting; validation: supporting); W. L. (funding acquisition: supporting; validation: supporting; writing – review & editing: supporting).

Conflicts of interest

There are no conflicts to declare.

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