

Cite this: *J. Mater. Chem. A*, 2025, 13, 39831

## Balancing Ru–O bond covalency and strength via atomic Ta doping for robust acidic oxygen evolution

Caini Ma,<sup>†ab</sup> Ning Zhang,<sup>†c</sup> Depeng Wang,<sup>ab</sup> Jian Ren,<sup>ab</sup> Tao Jiang,<sup>ab</sup> Tong Liu,<sup>abd</sup> Alexandra A. Zverko,<sup>e</sup> Haixia Zhong,<sup>†\*abd</sup> and Xinbo Zhang,<sup>†\*abd</sup>

Ruthenium dioxide electrocatalysts hold promise for achieving high oxygen evolution reaction activity in proton exchange membrane water electrolyzers (PEMWEs). However, the control of their stability remains extremely challenging due to their easy oxidation and dissolution during electrochemical reactions. Herein, we address these challenges by constructing atomically dispersed Ta–O–Ru asymmetric local motifs for an efficient and stable OER process via a simple molten salt-assisted method. The optimized candidate (RuO<sub>x</sub>–10Ta) exhibits a low overpotential of 189 mV and high durability exceeding 700 hours at 10 mA cm<sup>-2</sup>, boasting a six-fold longer lifespan than commercial RuO<sub>2</sub>. *In situ* characterizations and DFT calculations reveal that in this asymmetric configuration, high-valent Ta dopants downshift the O 2p and Ru 4d band centres, thereby moderating the Ru–O covalency to facilitate water dissociation and intermediate transformation. Meanwhile, this Ta–O–Ru motif with a robust Ta–O bond acts as an electron and structural buffer that suppresses lattice oxygen loss and Ru overoxidation, enhancing the overall catalyst stability. When integrated into a PEMWE device as an anode catalyst, it achieved a low cell voltage of 1.639 V at 1 A cm<sup>-2</sup> and sustained stable operation for over 200 hours at 200 mA cm<sup>-2</sup>. These findings highlight the potential of asymmetric coordination engineering as a generalizable strategy for developing robust and efficient OER catalysts under harsh acidic conditions.

Received 6th September 2025  
Accepted 20th October 2025

DOI: 10.1039/d5ta07271a

rsc.li/materials-a

<sup>a</sup>State Key Laboratory of Rare Earth Resource Utilization, Changchun Institute of Applied Chemistry, Chinese Academy of Sciences, No. 5625, Renmin Street, Chaoyang District, Changchun, Jilin, 130022, P. R. China. E-mail: hxzhong@ciac.ac.cn; xbzhang@ciac.ac.cn

<sup>b</sup>School of Applied Chemistry and Engineering, University of Science and Technology of China, No. 96, Jinzhai Road, Baohe District, Hefei City, Anhui, 230026, P. R. China

<sup>c</sup>State Key Laboratory of Critical Metals Beneficiation, Metallurgy and Purification, Zhengzhou University, No. 100, Kexuedadao Road, Zhongyuan District, Zhengzhou City, Henan, 450001, P. R. China

<sup>d</sup>China-Belarus Belt and Road Joint Laboratory on Advanced Materials and Manufacturing, Changchun Institute of Applied Chemistry, Chinese Academy of Sciences, No. 5625, Renmin Street, Chaoyang District, Changchun City, Jilin, 130022, P. R. China

<sup>e</sup>Faculty of Mechanical Engineering, Belarusian National Technical University, 65, Nezavisimosti Ave., 220072, Minsk, Republic of Belarus

† These authors contributed equally to this work.



Haixia Zhong

Dr. Haixia Zhong has been a Professor in the State Key Laboratory of Rare Earth Resource Utilization, Changchun Institute of Applied Chemistry, Chinese Academy of Sciences since 2022. She received her PhD from the University of Chinese Academy of Sciences in 2017 under the guidance of Professor Xinbo Zhang. Then, she worked as an Alexander von Humboldt Postdoctoral Fellow at the Dresden University of Technology (host: Professor Xinliang Feng). Her research interest is focused on the development of hybrid electrocatalysts and the application of hydrogen and ammonia-related energy conversion and storage.



# 1 Introduction

A proton exchange membrane water electrolyser (PEMWE), featuring a rapid response to intermittent renewable electricity (e.g. solar and wind), compact design, and high-purity hydrogen production,<sup>1–4</sup> is pivotal in decarbonized energy systems. Nevertheless, its efficiency is fundamentally constrained by the sluggish anodic oxygen evolution reaction (OER)<sup>5,6</sup> and the fast deactivation of catalysts.<sup>7,8</sup> Although IrO<sub>2</sub> represents a state-of-the-art acidic OER catalyst, the scarcity and prohibitive cost of iridium severely impede the large-scale PEMWE deployment.<sup>9–11</sup> In contrast, Ru-based catalysts, offering better economic feasibility and superior activity, are emerging as a promising alternative for the acidic OER.<sup>12–14</sup> Despite their promising attributes, bulk RuO<sub>2</sub> catalysts undergo structural degradation that stems from their inherent electronic structure. In essence, strong Ru–O covalency in ruthenium oxides enables the activation of lattice oxygen and participation in the OER process, thereby significantly improving the catalytic activity.<sup>15</sup> Unfortunately, such involvement of lattice oxygen accelerates the generation and outward diffusion of oxygen vacancies, progressively eroding the structural integrity. Concomitantly, the removal of lattice oxygen raises the local Ru valence and promotes the formation of high-valent and volatile RuO<sub>4</sub>, leading to the irreversible loss of active sites.<sup>13–17</sup> Consequently, it is crucial to balance the activity and stability in RuO<sub>2</sub> with fine-tuning the electronic structure toward efficient and stable OER.

To break the activity–stability trade-off, several strategies including strain engineering,<sup>16,17</sup> defect engineering,<sup>18,19</sup> interface engineering,<sup>20–22</sup> high-entropy alloy design<sup>23</sup> and heteroatom incorporation<sup>24–30</sup> have been explored. Among these, foreign atom doping offers a powerful approach to directly modify the local catalytic microenvironment for activity enhancement. Concurrently, effective doping can optimize the electronic configuration of active sites and enhance the atomic cohesion, thereby suppressing the lattice oxygen loss and ruthenium dissolution. For example, Guo *et al.* demonstrated that Er doping results in the 4d–2p–4f hybridization in the Er-doped RuO<sub>2</sub> catalyst, which continuously tunes the Ru–O covalency, increases the Ru–O–vacancy formation energies and strengthens the oxygen intermediate adsorption, thereby resulting in enhanced activity and stability.<sup>29</sup> In this context, developing an asymmetric M–O–Ru active motif with an optimized local structure through metal atom doping engineering is attractive to boost an effective and stable OER process. To avoid the non-uniform doping sites or even phase separation that will sacrifice the mass activity,<sup>31</sup> key criteria are necessary for rational doping engineering toward building the asymmetric Ru active motif, including: (i) lattice compatibility with the matched ionic radii between M and the host cation Ru<sup>4+</sup>, which ensures the stable solid-solution formation with rich atomic interfaces and no inert phase segregation;<sup>32,33</sup> (ii) lattice-oxygen buffering through strong M–O interactions for stabilizing lattice oxygen, suppressing V<sub>O</sub> aggregation and the collapse of the Ru oxide lattice;<sup>34</sup> (iii) electron buffering capability for the dynamic

charge redistribution in M–O–Ru linkages, which is favourable for the fine-tuning of the Ru–O covalency and the adjustable valence state of M against the oxidation condition.<sup>35</sup>

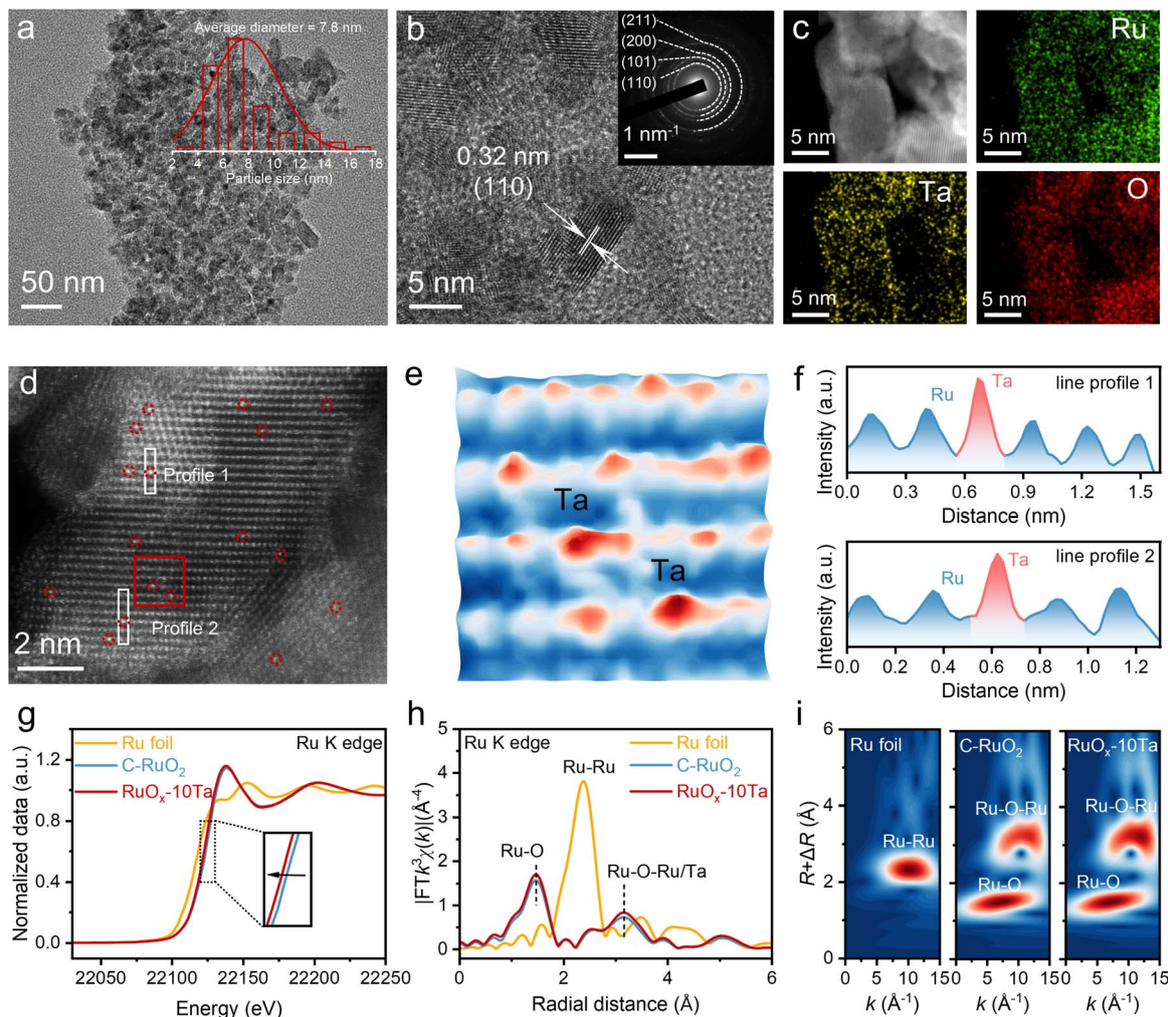
Herein, we present the example of atomically dispersed Ta–O–Ru asymmetric centres for robust OER through Ta doping regarding the strong Ta–O bond (839 kJ mol<sup>−1</sup>),<sup>36,37</sup> higher and adaptable valence (Ta<sup>5+</sup> vs. Ru<sup>4+</sup>)<sup>38</sup> and similar ionic radius (Ta<sup>5+</sup> 64 pm ≈ Ru<sup>4+</sup> 62 pm).<sup>39</sup> The resulting atomic dispersion of Ta within the RuO<sub>x</sub> matrix enables finely modulated local environments of Ru and O sites. Both experimental results and theoretical calculations reveal that the asymmetric Ta–O–Ru unit, characterized by moderate Ru–O covalency and strong Ta–O–Ru linkage, effectively promotes the lattice oxygen-involved OER process, wherein excessive lattice oxygen loss and Ru overoxidation are simultaneously suppressed. As expected, the optimized RuO<sub>x</sub>–10Ta exhibits high OER performance with an outstanding mass activity of 477.17 A g<sub>Ru</sub><sup>−1</sup> at 1.5 V vs. RHE, which is 62% higher than the pristine RuO<sub>x</sub> and 13-fold superior to commercial RuO<sub>2</sub>. This is achieved alongside a low overpotential of 189 mV at 10 mA cm<sup>−2</sup> and remarkable stability exceeding 700 h at 10 mA cm<sup>−2</sup> in 0.5 M H<sub>2</sub>SO<sub>4</sub> (6-fold superior to commercial RuO<sub>2</sub>), ranking among the most advanced Ru-based OER catalysts. Furthermore, a PEMWE device equipped with a RuO<sub>x</sub>–10Ta anode demonstrated a low cell voltage of 1.639 V at 1 A cm<sup>−2</sup> and maintained stable operation for over 200 hours at 200 mA cm<sup>−2</sup>, highlighting its good feasibility for the practical application. Our work highlights the potential of rationally developing atomically dispersed M–O–Ru asymmetric centres for acidic OER.

## 2 Results and discussion

### 2.1 Preparation and structural characterization

A series of tantalum-doped Ru oxide nanoparticles (RuO<sub>x</sub>–αTa, α = 5, 10, and 20, corresponding to the nominal Ta loadings of 5%, 10%, and 20%, respectively) and pristine RuO<sub>x</sub> were synthesized *via* a modified molten salt-assisted method (see details in the SI, Fig. S1).<sup>40,41</sup> The molten NaNO<sub>3</sub> was used to facilitate the metal-ion exchange and oxide formation. X-ray diffraction (XRD) patterns showed that all RuO<sub>x</sub>–αTa (α = 5, 10, and 20) and RuO<sub>x</sub> presented a similar single rutile phase compared to C–RuO<sub>2</sub> (commercial RuO<sub>2</sub>), with no detectable impurities from Ta<sub>2</sub>O<sub>5</sub> (Fig. S2). A systematic downshift of the diffraction peaks is observed as the Ta content increases, which is likely due to lattice expansion caused by the substitution of Ru with the larger Ta atoms. SEM imaging combined with elemental mapping (Fig. S3) indicated that the RuO<sub>x</sub>–10Ta sample consists of interconnected nanoparticles, wherein the Ru, Ta, and O elements are uniformly distributed without detectable Na signal, suggesting the effective removal of the NaNO<sub>3</sub> salt after synthesis. Inductively coupled plasma mass spectrometry (ICP-MS, Table S1) confirmed that the elemental composition of RuO<sub>x</sub>–10Ta closely matches the nominal loading values. In addition, transmission electron microscopy (TEM) images (Fig. 1a, S4 and S5) show well-dispersed nanoparticles for RuO<sub>x</sub>–5Ta/10Ta/20Ta with an average size of ~7.8 nm, slightly smaller than the homemade RuO<sub>x</sub> (9.3 nm,





**Fig. 1** Structural characterization of  $\text{RuO}_x-\alpha\text{Ta}$ . (a) TEM image with the diameter distribution (inset) of  $\text{RuO}_x-10\text{Ta}$ . (b) HR-TEM image of  $\text{RuO}_x-10\text{Ta}$  with the SAED image (inset). (c) Elemental mappings of  $\text{RuO}_x-10\text{Ta}$ . (d) Aberration-corrected HAADF-STEM image. (e) 3D atom-overlapping Gaussian function fitting mapping of the selected red square area in (d). (f) Intensity profiles along the white rectangle regions in (d). (g) Ru K-edge XANES spectra of  $\text{RuO}_x-10\text{Ta}$  and the corresponding reference samples. (h) Fourier transforms of the  $k^3$ -weighted spectra at the Ru K edge of  $\text{RuO}_x-10\text{Ta}$  and the corresponding reference samples. (i) Wavelet transform for the  $k^3$ -weighted EXAFS signals at the Ru K-edge of  $\text{RuO}_x-10\text{Ta}$  and the corresponding reference samples.

Fig. S6). The slight reduction in particle size upon Ta incorporation can be attributed to the effect of dopants on nucleation dynamics during molten synthesis.<sup>42</sup> In Fig. 1b, well-resolved lattice fringes with  $d$ -spacings of 0.32 nm are attributed to the (110) planes of the rutile  $\text{RuO}_2$  phase, similar to those of C- $\text{RuO}_2$  (Fig. S7). The selected area electron diffraction (SAED) pattern analysis (inset of Fig. 1b) confirms the polycrystalline rutile structure with clear (110), (101), (211) and (200) diffraction rings for  $\text{RuO}_x-10\text{Ta}$  and no detectable  $\text{Ta}_2\text{O}_5$  signatures, which is consistent with the XRD results. In addition, the energy-dispersive X-ray spectra (EDS, Fig. S8) and elemental mapping together with EDS line-scans (Fig. S9) demonstrate the homogeneous distribution of Ta throughout  $\text{RuO}_x$  nanoparticles. As shown in Fig. 1d, the aberration-corrected high-angle annular dark field scanning transmission electron microscopy (HAADF-STEM) image reveals that Ta atoms were atomically dispersed throughout the lattice of the  $\text{RuO}_x$  matrix.

It was found that Ta ( $Z = 73$ ) dopants appeared as isolated and significantly brighter dots within the atom columns (encircled in red), owing to their stronger electron scattering compared to the surrounding Ru ( $Z = 44$ ) and O ( $Z = 8$ ) atoms. The distinct atomic-scale distribution was further corroborated by 3D Gaussian fitting mapping (Fig. 1e) and intensity profile analysis (Fig. 1f). These results evidence the high dispersivity of the Ta dopants, which indicates abundant Ta-O-Ru motifs. Raman spectroscopy was used to further investigate the crystal structure of  $\text{RuO}_x-10\text{Ta}$  (Fig. S10). The absence of discernible peaks assignable to crystalline  $\text{Ta}_2\text{O}_5$  excludes the phase segregation. Three peaks which can be assigned to the  $E_g$ ,  $A_{1g}$  and  $B_{2g}$  modes of rutile  $\text{RuO}_2$  are broadened compared to the pristine  $\text{RuO}_x$ ,<sup>28</sup> which is associated with the fact that the substitutional doping of Ta impaired the crystalline symmetry.<sup>43,44</sup>

X-ray photoelectron spectroscopy (XPS) was performed to characterize the chemical states and the electronic



configurations of the catalysts (Fig. S11). In the Ta 4f region (Fig. S11b), two peaks centred at 24.95 eV and 26.85 eV are assignable to Ta 4f<sub>7/2</sub> and 4f<sub>5/2</sub>, respectively, consistent with successful Ta incorporation. These binding energies fall between those of Ta<sup>5+</sup> (25.90/27.90 eV) and Ta<sup>4+</sup> (23.30/25.20 eV),<sup>45,46</sup> suggesting an intermediate Ta oxidation state (+4 to +5). The slight decrease in average valence is likely caused by oxygen vacancies and other structural defects formed during molten-salt synthesis.<sup>40,47</sup> Such valence state is beneficial for the electron donation from Ta to Ru *via* Ta–O–Ru linkage.<sup>28</sup> The overlapping region with the O 2s signal was not analysed due to the peak interference. Compared to the commercial RuO<sub>2</sub> (Ru<sup>4+</sup>, Ru 3d<sub>5/2</sub> located at 280.50 eV and 3d<sub>3/2</sub> at 284.70 eV),<sup>41,47,48</sup> the Ru 3d<sub>5/2</sub> peak of the homemade RuO<sub>x</sub> shifts by ~0.10 eV toward lower binding energy, indicating a slight electron enrichment around Ru. This modest negative shift is possibly related to the molten salt synthesis as well. Crucially, upon Ta doping (RuO<sub>x</sub>–10Ta), the shift further increases to ~0.20 eV, demonstrating further electron donation to Ru from Ta *via* the Ta–O–Ru linkage. O 1s XPS deconvolution (Fig. S11d and e) uncovered four oxygen species: lattice oxygen (529.02 eV), oxygen vacancy (530.00 eV), adsorbed –OH (531.30 eV) and adsorbed H<sub>2</sub>O (532.86 eV).<sup>47,49</sup> Compared with C–RuO<sub>2</sub>, the home-made RuO<sub>x</sub> possessed an elevated oxygen-vacancy concentration, attributed to the etching effect of the molten salt synthesis.<sup>40,47,49</sup> Significantly, the introduction of Ta increased the amount of lattice oxygen, which was attributable to the presence of more surrounding O<sup>2–</sup> ions for achieving a higher valence of Ta and strong binding between Ta and O. The concomitant increase in Ru electron density and lattice oxygen content suggested the effective electronic modulation by the Ta–O–Ru configuration. Moreover, no Na-related signal was detected from RuO<sub>x</sub>–10Ta (Fig. S11a), further ruling out residual Na and its possible effects on the OER performance.<sup>38</sup> H<sub>2</sub> temperature-programmed reduction (H<sub>2</sub>-TPR) (Fig. S12) corroborated the tuned bond strength and oxygen lability *via* Ta doping. RuO<sub>x</sub> exhibited a significant shift of the reduction peak to a lower temperature (~50 °C) compared to C–RuO<sub>2</sub>, indicating enhanced lattice oxygen activity. Upon Ta incorporation, the peak slightly shifted to higher temperature with decreased area and minor splitting, suggesting fewer labile oxygen species for RuO<sub>x</sub>–10Ta, which was probably attributable to stronger Ta–O–Ru interactions.<sup>51,52</sup> The signal for RuO<sub>x</sub>–10Ta remained lower than that of C–RuO<sub>2</sub>, indicating that the lattice oxygen was maintained in a more activated state relative to C–RuO<sub>2</sub>, which indicates the appropriate oxygen activity and strong stability of the Ta–O–Ru motifs. Furthermore, X-ray absorption spectroscopy (XAS) was employed to probe the environment of Ru sites for RuO<sub>x</sub>–10Ta. Fig. 1g depicts the Ru K-edge X-ray absorption near-edge structure (XANES) spectrum of RuO<sub>x</sub>–10Ta, alongside Ru foil and C–RuO<sub>2</sub> as references. A shift in the absorption edge of RuO<sub>x</sub>–10Ta toward lower energy compared to that of C–RuO<sub>2</sub> reference (Fig. 1g) demonstrated a moderately reduced Ru oxidation state (<+4) in RuO<sub>x</sub>–10Ta.<sup>25,53</sup> This result is in line with the XPS data, further verifying the electron transfer in Ta–O–Ru linkages, which increases electron density around the Ru sites and thereby prevents overoxidation of Ru during the OER

processes. The Fourier-transformed extended X-ray absorption fine structure (FT-EXAFS) spectrum (phase-uncorrected) of RuO<sub>x</sub>–10Ta at the Ru K-edge shows a prominent peak at ~1.55 Å affiliated with the first Ru–O coordination shell, which is identical to rutile RuO<sub>2</sub> (Fig. 1h).<sup>25</sup> Quantitative FT-EXAFS fitting reveals an average Ru–O coordination number of 5.68 (Fig. S13 and Table S2), which is significantly lower than the ideal value of 6.00 for octahedrally coordinated Ru in C–RuO<sub>2</sub>, indicating the generation of oxygen vacancies (O<sub>v</sub>) around the Ru sites upon Ta incorporation. In addition, the Ru–O bond length shows only a slight increase (1.98 Å for C–RuO<sub>2</sub> and 1.99 Å for RuO<sub>x</sub>–10Ta), in good agreement with the XRD results. It is thus clear that the overall rutile RuO<sub>2</sub> lattice remains intact despite the introduction of O<sub>v</sub> for RuO<sub>x</sub>–10Ta. Wavelet-transform (WT) analysis, providing simultaneous resolution in *k* and *R*-space, allows for the discernment of atoms sharing similar coordination environments and proximities. In Fig. 1i, the WT contour plot of RuO<sub>x</sub>–10Ta exhibits enhanced scattering intensity around *k* = 12 Å<sup>–1</sup> compared to C–RuO<sub>2</sub>, which can be attributed to the heavier element Ta that is incorporated into the RuO<sub>x</sub> matrix, further indicating the formation of the Ta–O–Ru linkage.<sup>54</sup> Overall, these comprehensive structural and electronic investigations confirm that Ta atoms are atomically dispersed in the RuO<sub>x</sub> lattice and contributed to the fine electronic modulation, which is anticipated to play a pivotal role in the electrocatalytic performance.

## 2.2 Electrocatalytic performance

The acid OER performance of RuO<sub>x</sub>– $\alpha$ Ta was systematically evaluated in a three-electrode system. Crucially, RuO<sub>x</sub>–10Ta exhibited high mass activity (477.17 A g<sub>Ru</sub><sup>–1</sup> at 1.5 V vs. RHE), representing a 62% improvement over pure RuO<sub>x</sub> (294.68 A g<sub>Ru</sub><sup>–1</sup>) and 13-fold higher than C–RuO<sub>2</sub> (35.35 A g<sub>Ru</sub><sup>–1</sup>) (Fig. 2a and S14), consistent with the TOF (turnover frequency) results (Fig. S15). It is worth noting that suitable Ta doping is crucial in improving the OER catalytic activity of Ru sites. Less or more Ta doping would cause catalytic activity compromise, which is due to the smaller amounts of Ta–O–Ru active motifs or the excessive site dilution and even phase separation, respectively.<sup>28,55</sup> In addition, commercial Ta<sub>2</sub>O<sub>5</sub> and homemade TaO<sub>x</sub> (Fig. S16) exhibited negligible current densities (<0.2 mA cm<sup>–2</sup>) even at a high potential of 1.70 V vs. RHE, underscoring the intrinsically inert nature of Ta oxides toward OER. In the linear sweep voltammetry (LSV) tests (Fig. 2b), RuO<sub>x</sub>–10Ta required a low overpotential of only 189 mV to achieve a current density of 10 mA cm<sup>–2</sup>, surpassing RuO<sub>x</sub>–5Ta (202 mV), RuO<sub>x</sub>–20Ta (210 mV) and C–RuO<sub>2</sub> (250 mV) (Fig. S17). It should be noted that RuO<sub>x</sub>–10Ta with lower Ru loading still exhibited high performance comparable to RuO<sub>x</sub>, further indicating the crucial role of Ta doping in enhancing the intrinsic activity of Ru. The high activity is further reflected by the smallest Tafel slope (55.9 mV dec<sup>–1</sup>, Fig. 2c), compared to RuO<sub>x</sub> (59.5 mV dec<sup>–1</sup>), RuO<sub>x</sub>–5Ta (65.5 mV dec<sup>–1</sup>), RuO<sub>x</sub>–20Ta (66.2 mV dec<sup>–1</sup>) and C–RuO<sub>2</sub> (147.4 mV dec<sup>–1</sup>), indicative of favourable OER kinetics.<sup>56,57</sup> The double-layer capacitances (C<sub>dl</sub>) were extracted from the cyclic voltammogram (CV) curves in the non-faradaic region (Fig. 2d



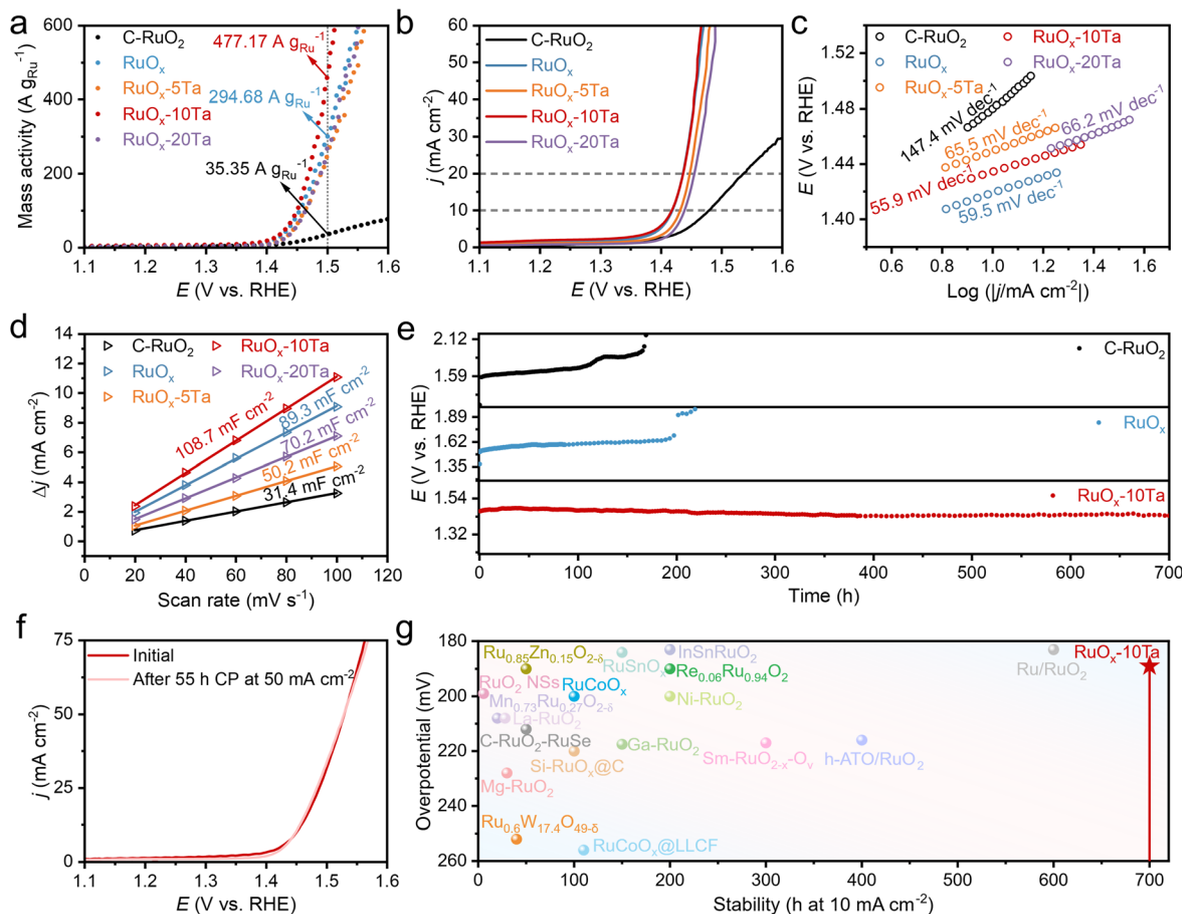


Fig. 2 Electrocatalytic performance. (a) LSV curves normalized by the Ru mass of  $\text{RuO}_x\text{-}\alpha\text{Ta}$  ( $\alpha = 5, 10$  and  $20$ ), homemade  $\text{RuO}_x$  and  $\text{C-RuO}_2$ . (b) LSV curves of  $\text{RuO}_x\text{-}\alpha\text{Ta}$ , homemade  $\text{RuO}_x$  and  $\text{C-RuO}_2$ . (c) Tafel plots derived from the polarization curves in (b). (d) Electrochemical double-layer capacitance ( $C_{dl}$ ) of the electrocatalysts. (e) Chronopotentiometric plots of  $\text{C-RuO}_2$ ,  $\text{RuO}_x$  and  $\text{RuO}_x\text{-}10\text{Ta}$  at  $10 \text{ mA cm}^{-2}$ . (f) LSV curves of  $\text{RuO}_x\text{-}10\text{Ta}$  before and after 55 h CP at  $50 \text{ mA cm}^{-2}$ . (g) Comparison of the overpotential and durability of  $\text{RuO}_x\text{-}10\text{Ta}$  at  $j = 10 \text{ mA cm}^{-2}$  with those of the previously reported OER catalysts in acidic media.

and S18) for the estimation of the electrochemical surface area (ECSA).  $\text{RuO}_x\text{-}10\text{Ta}$  shows the largest  $C_{dl}$  ( $108.7 \text{ mF cm}^{-2}$ ) compared with  $\text{RuO}_x\text{-}5\text{Ta}$  ( $50.2 \text{ mF cm}^{-2}$ ),  $\text{RuO}_x\text{-}20\text{Ta}$  ( $70.2 \text{ mF cm}^{-2}$ ),  $\text{RuO}_x$  ( $89.3 \text{ mF cm}^{-2}$ ) and  $\text{C-RuO}_2$  ( $31.4 \text{ mF cm}^{-2}$ ). This highest  $C_{dl}$  value for  $\text{RuO}_x\text{-}10\text{Ta}$  indicates its largest ECSA, suggesting the most abundant accessible active sites among the series of catalysts. Electrochemical impedance spectroscopy (EIS) further demonstrated the lowest charge-transfer resistance ( $R_{ct}$ ) for  $\text{RuO}_x\text{-}10\text{Ta}$ , highlighting the improved charge transfer efficiency introduced by Ta doping (Fig. S19).<sup>20</sup>

Moreover,  $\text{RuO}_x\text{-}10\text{Ta}$  delivered long-term durability under acidic OER conditions. At the current density of  $10 \text{ mA cm}^{-2}$  in  $0.5 \text{ M H}_2\text{SO}_4$ ,  $\text{RuO}_x\text{-}10\text{Ta}$  maintains stable performance over 700 hours without apparent decay, far exceeding the pristine  $\text{RuO}_x$  (less than 200 h) and  $\text{C-RuO}_2$  (120 h), and most of the reported Ru-based electrocatalysts (Fig. 2g and Table S4). Even at a higher current density of  $50 \text{ mA cm}^{-2}$  (Fig. S20),  $\text{RuO}_x\text{-}10\text{Ta}$  also demonstrated impressive durability for more than 300 h.

Post-OER characterizations corroborated the structural integrity of the catalyst after long-term electrolysis.  $\text{RuO}_x\text{-}10\text{Ta}$  exhibits negligible variation of the LSV curve after 55 hours running at  $50$

$\text{mA cm}^{-2}$  (Fig. 2f), demonstrating good activity retention. Additionally, the recovered  $\text{RuO}_x\text{-}10\text{Ta}$  remains structurally and compositionally stable. SEM shows a continuous network of connected nanoparticles, and compositional mapping indicates uniform Ru and Ta distributions (Fig. S21). TEM further shows well-dispersed particles with only slight growth (average size: 9.3 nm) and minimal agglomeration (Fig. S22a). The well-resolved lattice fringes and SAED pattern confirmed the retention of the rutile phase (inset of Fig. S22b). Consistently, EDS maps (Fig. S22c and d) show homogeneous Ru and Ta without evidence of phase segregation. High-resolution XPS spectra after OER (Fig. S23) exhibit high-valent  $\text{Ru}^{6+}$  species in  $\text{RuO}_x$  and  $\text{C-RuO}_2$ . However, such Ru species were not observed for  $\text{RuO}_x\text{-}10\text{Ta}$ , suggesting the suppressed Ru overoxidation as a result of Ta doping. This is consistent with the high  $S$ -number (stability number,  $2.70 \times 10^5$ ) of  $\text{RuO}_x\text{-}10\text{Ta}$ , calculated from ICP-MS analysis of the electrolyte after 400 h CP at  $10 \text{ mA cm}^{-2}$ . Furthermore, the O 1s spectra revealed a higher lattice oxygen content, as evidenced by a larger peak area in  $\text{RuO}_x\text{-}10\text{Ta}$  relative to both undoped  $\text{RuO}_x$  and  $\text{C-RuO}_2$ , demonstrating better preservation of the oxygen framework under harsh electrochemical conditions.



Additionally, the generality of the asymmetric active unit was further extended by introducing Zr, Hf, and Nb dopants, which also feature strong bonding to oxygen, and adjustable high valence states similar to Ta. All variants exhibited good OER activity with overpotentials below 220 mV at 10 mA cm<sup>-2</sup> (Fig. S24) and significantly prolonged durability at 50 mA cm<sup>-2</sup> when compared to pristine RuO<sub>x</sub> (Fig. S25). Structural analyses (XRD, TEM and XPS) confirmed the analogous modulation of the crystallinity, Ru oxidation states and oxygen species (Fig. S26–S28), which further affirms that the asymmetric M–O–Ru active unit *via* high-valent doping engineering can effectively enhance the catalyst durability, while preserving the high activity.

### 2.3 Mechanism investigation

To unravel the oxygen evolution process on RuO<sub>x</sub>-10Ta, electrochemical and *in situ* spectral characterizations were performed. First, pH-dependent linear sweep voltammetry (LSV) revealed a clear activity decline with increasing pH on RuO<sub>x</sub> and RuO<sub>x</sub>-10Ta (Fig. 3a, b and S30), which is a hallmark of the lattice oxygen participation process (LOM) as the LOM is inherently sensitive to proton availability.<sup>58,59</sup> Furthermore, the

tetramethylammonium cations (TMA<sup>+</sup>) were used as mechanistic indicators. It was found that peroxy-like (\*O<sub>2</sub><sup>2-</sup>) intermediates generated *via* the LOM pathway feature longer lifetimes than the superoxo-like (\*O<sub>2</sub><sup>-</sup>) species produced through adsorbate evolution mechanisms (AEM), thus enabling the direct probing of \*O<sub>2</sub><sup>2-</sup> during the OER process.<sup>60,61</sup> As the electrophilic site, TMA<sup>+</sup> selectively stabilizes the negative \*O<sub>2</sub><sup>2-</sup> species through coulombic trapping, thereby inhibiting the O–O coupling kinetics and inhibiting the LOM pathway. A marked suppression in OER performance was observed for both RuO<sub>x</sub> and RuO<sub>x</sub>-10Ta along with the addition of 0.1 M TMA<sup>+</sup>, corroborating the involvement of the LOM pathway on RuO<sub>x</sub> and RuO<sub>x</sub>-10Ta (Fig. S31 and 3c). The hypothesis is also strongly supported by *in situ* differential electrochemical mass spectrometry (DEMS) using <sup>18</sup>O isotopic labelling,<sup>62</sup> wherein distinct <sup>34</sup>O<sub>2</sub> signals were detected (Fig. 3d and S32) following electrochemical pre-conditioning in H<sub>2</sub><sup>18</sup>O (see Experimental section in SI). In addition, *in situ* attenuated total reflection infrared absorption spectroscopy (ATR-SEIRAS) measurements were carried out to detect the reaction intermediates. As the potential increases, the absorption bands at 1170 cm<sup>-1</sup> (RuO<sub>x</sub>) and 1190 cm<sup>-1</sup> (RuO<sub>x</sub>-10Ta) exhibit

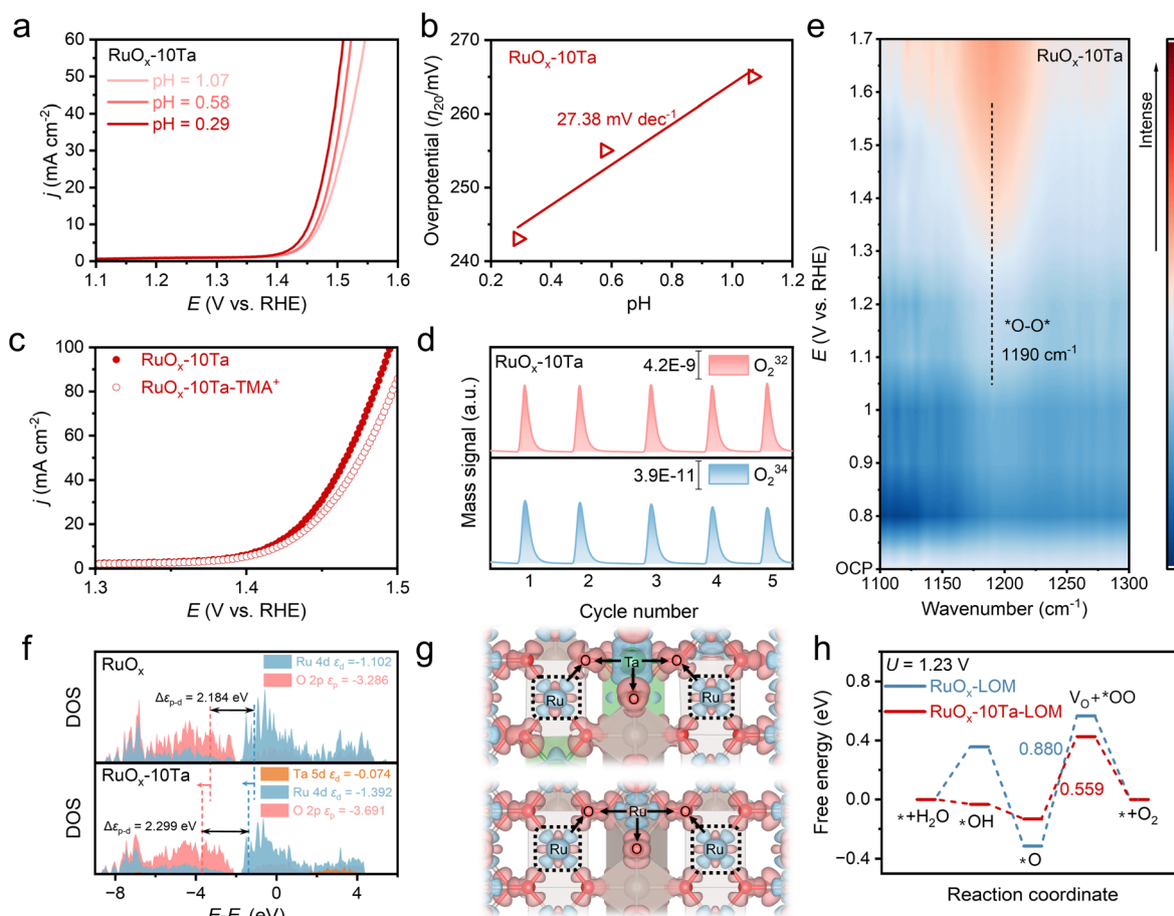


Fig. 3 OER mechanism studies. (a) pH-dependent LSV curves of RuO<sub>x</sub>-10Ta. (b) Overpotentials of RuO<sub>x</sub>-10Ta at 20 mA cm<sup>-2</sup> as a function of pH values. (c) LSV curves of RuO<sub>x</sub>-10Ta in 0.5 M H<sub>2</sub>SO<sub>4</sub> electrolyte with or without TMA<sup>+</sup>. (d) *In situ* DEMS with <sup>18</sup>O-labelled measurements for Ta–RuO<sub>x</sub>. (e) *In situ* ATR-SEIRAS measurements of RuO<sub>x</sub>-10Ta. (f) DFT-calculated DOS curves for RuO<sub>x</sub> and RuO<sub>x</sub>-10Ta. (g) DFT-calculated CDD maps for RuO<sub>x</sub> and RuO<sub>x</sub>-10Ta. (h) Free energy diagram of LOM reaction pathways for RuO<sub>x</sub> and RuO<sub>x</sub>-10Ta at U = 1.23 V.



progressive intensification (Fig. S33 and 3e), which is assignable to the  $*\text{O}-\text{O}*$  intermediate.<sup>20,63–65</sup> This observation provides direct molecular spectroscopic evidence that the oxygen evolution reaction on both  $\text{RuO}_x$  and  $\text{RuO}_x-10\text{Ta}$  catalysts proceeds through the lattice oxygen-mediated pathway. Additionally, *in situ* EIS provided further insights into the electrochemical processes on the catalysts (Fig. S34). In the Bode plots, the high-frequency region represents the oxidation of the catalysts and  $\text{RuO}_x-10\text{Ta}$  exhibited a near-zero phase angle, indicating minimal surface oxidation.<sup>66</sup> The low-frequency region corresponds to the interfacial charge transfer during OER.  $\text{RuO}_x-10\text{Ta}$  exhibited a smaller phase angle, demonstrating the facilitated charge transfer and enhanced reaction kinetics of OER.<sup>67,68</sup>

To better understand the OER mechanism with the atomically dispersed Ta–O–Ru asymmetric motifs, density functional theory (DFT) calculations were conducted. Based on the aforementioned analyses, computational models of pristine  $\text{RuO}_x$  and Ta-doped  $\text{RuO}_x$  were constructed (Fig. S35–S38). The densities of states (DOS) (Fig. 3f) for  $\text{RuO}_x$  and  $\text{RuO}_x-10\text{Ta}$  show a downward shift of the O 2p band centre from  $-3.286$  eV ( $\text{RuO}_x$ ) to  $-3.691$  eV ( $\text{RuO}_x-10\text{Ta}$ ) and of the Ru 4d centre from  $-1.102$  eV ( $\text{RuO}_x$ ) to  $-1.392$  eV ( $\text{RuO}_x-10\text{Ta}$ ), suggesting a moderately reduced Ru–O covalency and the stabilization of both Ru and O sites.<sup>29,50,58</sup> The gap between the O 2p band and Ru 4d band centres for  $\text{RuO}_x-10\text{Ta}$  increases from 2.184 to 2.299 eV ( $\Delta = 0.115$  eV), further evidencing a moderate attenuation of the Ru–O covalency, which is favourable for

suppressing over-oxidation while stabilizing the lattice oxygen. In addition, the charge density difference (CDD) maps (Fig. 3g) clearly reveal the characteristic charge redistribution induced by Ta doping. Pronounced electron depletion on Ta atoms, along with enhanced electron accumulation on neighbouring O atoms, suggested the strong interaction between Ta and O and the strong Ta–O bond,<sup>69</sup> which is beneficial for stabilizing lattice oxygen. Moreover, this charge redistribution further increases the electron density at adjacent Ru sites with more electron accumulation (Fig. S39) compared to  $\text{RuO}_x$ , consistent with the XAS and XPS results. This would be beneficial for suppressing the overoxidation of Ru sites and thereby improving stability.<sup>70</sup> Furthermore, OER pathways were investigated for analysis of their intrinsic activity. Accordingly, the rate-determining step (RDS) of AEM and LOM pathways can be assigned to the formation of  $*\text{OOH}$  and the generation of  $\text{V}_\text{O} + *\text{O}_2$ , respectively. As shown in Fig. 3h, S35 and S36, both  $\text{RuO}_x$  and  $\text{RuO}_x-10\text{Ta}$  deliver lower energy barriers (0.880 eV and 0.559 eV) for the RDS of LOM pathways than the AEM ( $\text{RuO}_x$ : 1.122 eV,  $\text{RuO}_x-10\text{Ta}$ : 0.756 eV). These results indicate that the subtly attenuated Ru–O covalency of  $\text{RuO}_x-10\text{Ta}$  preserves the high intrinsic activity through lattice oxygen participation. It should be noted that the kinetic barrier for the initial water dissociation step is significantly reduced on  $\text{RuO}_x-10\text{Ta}$  (Fig. 3h, S40 and S41), which is in line with the *in situ* Raman observations of enriched free- $\text{H}_2\text{O}$  populations at the Ta-doped interface (Fig. S42).<sup>61,63,71,72</sup> The facilitated water dissociation contributes to the refilling of oxygen vacancies ( $\text{V}_\text{O}$ ) by the  $*\text{OH}$  species, limiting excessive

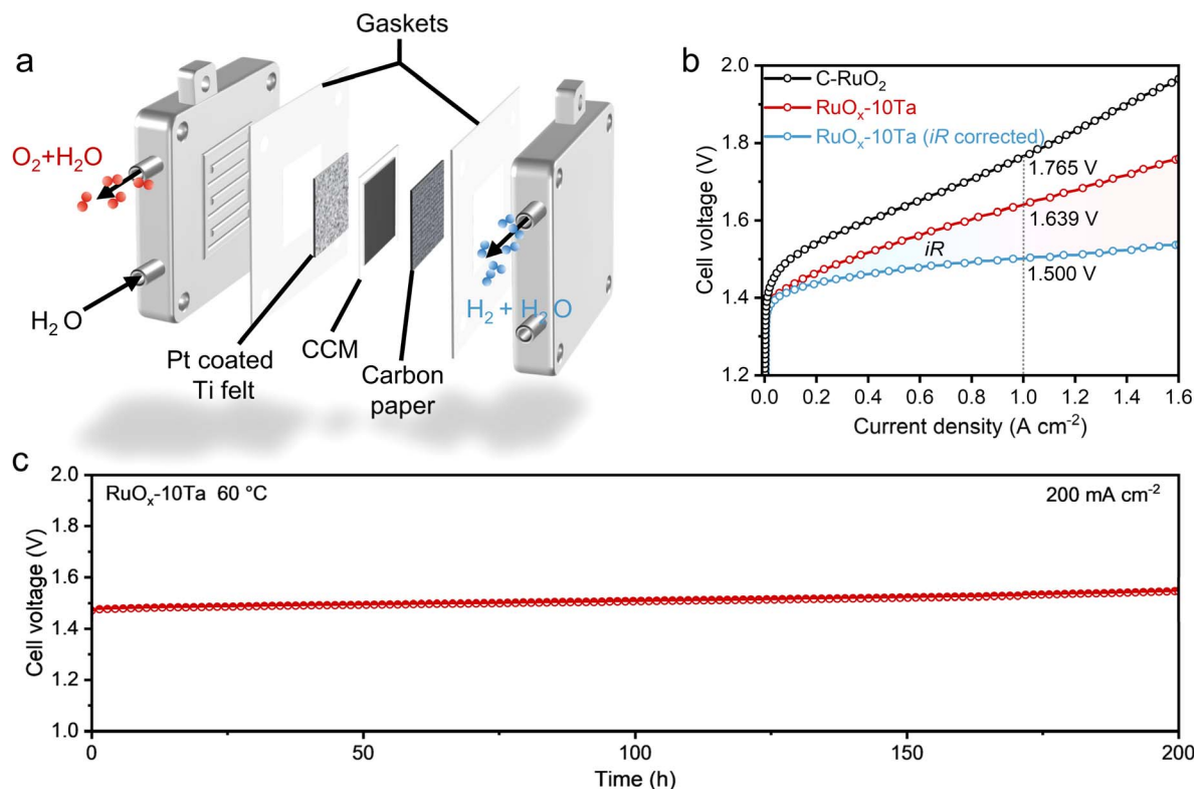


Fig. 4 Performance of the PEMWE device using the  $\text{RuO}_x-10\text{Ta}$  OER catalyst. (a) Schematic of a PEMWE. (b) Current–voltage polarization curves of the PEMWE cell with  $\text{RuO}_x-10\text{Ta}$  and C– $\text{RuO}_2$ . (c) Chronopotentiometry curve using  $\text{RuO}_x-10\text{Ta}$  at 60 °C and 200 mA cm<sup>-2</sup>.



lattice oxygen loss and structural degradation.<sup>29,73</sup> Overall, the Ta–O–Ru asymmetric motif effectively balances the Ru–O bond covalency and strength to favour LOM activity, while inhibiting structural degradation.

## 2.4 Performance of PEMWE devices

Motivated by the high OER performance of RuO<sub>x</sub>–10Ta, we constructed a PEM electrolyser using RuO<sub>x</sub>–10Ta and 40 wt% Pt/C as anode and cathode catalysts to evaluate the potential of our catalysts in practical application (Fig. 4a). The performance evaluation of all devices was conducted at 60 °C using deionized water. As depicted in Fig. 4b, the PEMWE cell employing RuO<sub>x</sub>–10Ta as an anode catalyst achieves a current density of 1 A cm<sup>−2</sup> at a cell voltage of only 1.639 V (1.500 V after *iR* compensation), which is markedly lower than that required by the commercial RuO<sub>2</sub>-based PEMWE system (Fig. S43). Impressively, RuO<sub>x</sub>–10Ta exhibits excellent long-term durability, maintaining stable operation for over 200 hours at a high current density of 200 mA cm<sup>−2</sup> (Fig. 4c). The corresponding degradation rate is as low as 3.75 μV h<sup>−1</sup>, indicating a negligible performance decay. Overall, these merits indicate the active and stable feature of RuO<sub>x</sub>–10Ta, with promising practical application prospects.

## 3 Conclusion

In this work, we overcome the intrinsic activity–stability trade-off in Ru-based acidic OER catalysts by engineering the asymmetric Ta–O–Ru active motifs *via* atomically doping Ta into the catalyst. Fine-tuning of the Ru–O covalency to balance the lattice oxygen activity and strong Ta–O–Ru bonds for structural robustness were simultaneously achieved. Combined experimental and theoretical analyses identified the dominant LOM pathway, which preserves high activity while ensuring enhanced stability. The resulting RuO<sub>x</sub>–10Ta delivers a high mass activity of 477.17 A g<sub>Ru</sub><sup>−1</sup> and an exceptional structural robustness under accelerated degradation conditions (700 h @ 10 mA cm<sup>−2</sup>), realizing about 6-fold lifespan extension over commercial RuO<sub>2</sub>. This robustness provides a foundation for high-current-density operation, as demonstrated by the stable operation of a PEMWE device at 200 mA cm<sup>−2</sup> for over 200 hours. This study demonstrates that asymmetric coordination engineering represents a powerful and broadly applicable strategy for enhancing the activity and stability in acidic OER catalysis. It opens avenues for the design of high-performance electrocatalysts based on controlled asymmetric local electronic structures.

## Author contributions

Conceptualization: X. Zhang and H. Zhong; methodology: D. Wang, C. Ma, and H. Zhong; validation: T. Liu, C. Ma, and H. Zhong; investigation: C. Ma, N. Zhang, J. Ren and T. Jiang; data analysis and assembly, C. Ma, D. Wang, J. Ren, T. Jiang and Alexandra A. Zverko; writing-original draft preparation: C. Ma and N. Zhang; writing-review and editing: C. Ma, N. Zhang, and H. Zhong; funding acquisition: X. Zhang and H. Zhong. All

authors have given approval for the final version of the manuscript.

## Conflicts of interest

The authors declare no conflicts of interest.

## Data availability

All data supporting the findings of this study are included in the manuscript and its supplementary information (SI). Additionally, the raw datasets are available from the corresponding authors upon reasonable request. Supplementary information: extended methods, datasets, and figures that support the main text. See DOI: <https://doi.org/10.1039/d5ta07271a>.

## Acknowledgements

This work was supported by the National Key R&D Program of China (2021YFB4000603), the National Natural Science Foundation of China (52273277 and U24A2062), Jilin Province Science and Technology Development Plan Funding Project (SKL202302039) and the Youth Innovation Promotion Association CAS (2021223). These authors thank the staff of beamline BL13SSW at the Shanghai Synchrotron Radiation Facility for the experimental support. H. Zhong acknowledges funding from the National Natural Science Foundation of China Outstanding Youth Science Foundation of China (Overseas).

## Notes and references

- 1 Y. Wang, M. Zhang, Z. Kang, L. Shi, Y. Shen, B. Tian, Y. Zou, H. Chen and X. Zou, *Nat. Commun.*, 2023, **14**, 5119.
- 2 H. Gao, Z. Xiao, S. Du, T. Liu, Y. C. Huang, J. Shi, Y. Zhu, G. Huang, B. Zhou, Y. He, C. L. Dong, Y. Li, R. Chen and S. Wang, *Angew. Chem., Int. Ed.*, 2023, **62**, e202313954.
- 3 Y. Wang, R. Ma, Z. Shi, H. Wu, S. Hou, Y. Wang, C. Liu, J. Ge and W. Xing, *Chem*, 2023, **9**, 2931–2942.
- 4 Y. Zhou, H. Zhong, S. Chen, G. Wen, L. Shen, Y. Wang, R. Chen, L. Tao and S. Wang, *Carbon Energy*, 2024, **7**, 23146–23157.
- 5 L. Han and Z. Xiang, *Fundam. Res.*, 2025, **5**, 624–639.
- 6 C. Rong, K. Dastafkan, Y. Wang and C. Zhao, *Adv. Mater.*, 2023, **35**, 2211884.
- 7 C. Xia, F.-M. Li, C. He, S. Zaman, W. Guo and B. Y. Xia, *Fundam. Res.*, 2024, DOI: [10.1016/j.fmre.2024.04.017](https://doi.org/10.1016/j.fmre.2024.04.017).
- 8 Y. Jia, Y. Li, Q. Zhang, S. Yasin, X. Zheng, K. Ma, Z. Hua, J. Shi, C. Gu, Y. Dou and S. Dou, *Carbon Energy*, 2024, **6**, e528.
- 9 J. Zhao, Y. Guo, Z. Zhang, X. Zhang, Q. Ji, H. Zhang, Z. Song, D. Liu, J. Zeng, C. Chuang, E. Zhang, Y. Wang, G. Hu, M. A. Mushtaq, W. Raza, X. Cai and F. Ciucci, *Nat. Nanotechnol.*, 2025, **20**, 57–66.
- 10 M. Bernt, A. Siebel and H. A. Gasteiger, *J. Electrochem. Soc.*, 2018, **165**, F305–F314.
- 11 B. Huang, Y. Ma, Z. Xiong, Z. Xiao, P. Wu, P. Jiang and M. Liang, *Energy Environ. Mater.*, 2020, **4**, 681–686.



- 12 C. Wei, Z. Wang, K. Otani, D. Hochfilzer, K. Zhang, R. Nielsen, I. Chorkendorff and J. Kibsgaard, *ACS Catal.*, 2023, **13**, 14058–14069.
- 13 R. Jiang, D. T. Tran, J. Li and D. Chu, *Energy Environ. Mater.*, 2019, **2**, 201–208.
- 14 Y. Wang, H. Yan and H. Fu, *eScience*, 2024, **5**, 100323.
- 15 M. Qi, X. Du, X. Shi, S. Wang, B. Lu, J. Chen, S. Mao, H. Zhang and Y. Wang, *J. Am. Chem. Soc.*, 2025, **147**, 18295–18306.
- 16 X. Cao, L. Miao, W. Jia, H. Qin, G. Lin, R. Ma, T. Jin and L. Jiao, *Nat. Commun.*, 2025, **16**, 6217.
- 17 Y. Xu, Z. Mao, J. Zhang, J. Ji, Y. Zou, M. Dong, B. Fu, M. Hu, K. Zhang, Z. Chen, S. Chen, H. Yin, P. Liu and H. Zhao, *Angew. Chem., Int. Ed.*, 2024, **63**, e202316029.
- 18 L. Zhou, Y. Shao, F. Yin, J. Li, F. Kang and R. Lv, *Nat. Commun.*, 2023, **14**, 7644.
- 19 W. He, X. Tan, Y. Guo, Y. Xiao, H. Cui and C. Wang, *Angew. Chem., Int. Ed.*, 2024, **63**, e202405798.
- 20 L. Deng, S. F. Hung, Z. Y. Lin, Y. Zhang, C. Zhang, Y. Hao, S. Liu, C. H. Kuo, H. Y. Chen, J. Peng, J. Wang and S. Peng, *Adv. Mater.*, 2023, **35**, e2305939.
- 21 J. Chen, Y. Ma, T. Huang, T. Jiang, S. Park, J. Xu, X. Wang, Q. Peng, S. Liu, G. Wang and W. Chen, *Adv. Mater.*, 2024, **36**, 2312369.
- 22 J. Chen, Y. Ma, C. Cheng, T. Huang, R. Luo, J. Xu, X. Wang, T. Jiang, H. Liu, S. Liu, T. Huang, L. Zhang and W. Chen, *J. Am. Chem. Soc.*, 2025, **147**, 8720–8731.
- 23 J. Chen, J. Ma, T. Huang, Q. Liu, X. Liu, R. Luo, J. Xu, X. Wang, T. Jiang, H. Liu, Z. Lv, T. Yao, G. Wang, X. Zheng, Z. Li and W. Chen, *Angew. Chem., Int. Ed.*, 2025, **64**, e202503330.
- 24 Y. Yu, F. Liu, X. Liu, L. Li, B. Zhou, H. Liao, P. Tan and J. Pan, *Chem. Eng. J.*, 2025, **514**, 163295.
- 25 Y. Shen, X. L. Zhang, M. R. Qu, J. Ma, S. Zhu, Y. L. Min, M. R. Gao and S. H. Yu, *Nat. Commun.*, 2024, **15**, 7861.
- 26 X. Wang, W. Pi, Z. Li, S. Hu, H. Bao, W. Xu and N. Yao, *Nat. Commun.*, 2025, **16**, 4845.
- 27 C. Zhou, L. Li, Z. Dong, F. Lv, H. Guo, K. Wang, M. Li, Z. Qian, N. Ye, Z. Lin, M. Luo and S. Guo, *Nat. Commun.*, 2024, **15**, 9774.
- 28 H. Liu, Z. Zhang, J. Fang, M. Li, M. G. Sendeku, X. Wang, H. Wu, Y. Li, J. Ge, Z. Zhuang, D. Zhou, Y. Kuang and X. Sun, *Joule*, 2023, **7**, 558–573.
- 29 L. Li, G. Zhang, C. Zhou, F. Lv, Y. Tan, Y. Han, H. Luo, D. Wang, Y. Liu, C. Shang, L. Zeng, Q. Huang, R. Zeng, N. Ye, M. Luo and S. Guo, *Nat. Commun.*, 2024, **15**, 4974.
- 30 W. Zheng, Y. Zhao, K. Jiang, F. Xie, L. Meng, S. Gao, J. Li, J. Lan, M. Luo, L. Liu and Y. Tan, *Nat. Commun.*, 2025, **16**, 6716.
- 31 Z. Zhang, S. Lu, M. Zhu, F. Wang, K. Yang, B. Dong, Q. Yao and W. Hu, *J. Am. Chem. Soc.*, 2025, **147**, 22806–22817.
- 32 B. Yuan, Q. Dang, H. Liu, M. G. Sendeku, J. Peng, Y. Fan, L. Cai, A. Cao, S. Chen, H. Li, Y. Kuang, F. Wang and X. Sun, *Nat. Commun.*, 2025, **16**, 4583.
- 33 D. Zhang, M. Li, X. Yong, H. Song, G. I. N. Waterhouse, Y. Yi, B. Xue, D. Zhang, B. Liu and S. Lu, *Nat. Commun.*, 2023, **14**, 2517.
- 34 X. Ping, Y. Liu, L. Zheng, Y. Song, L. Guo, S. Chen and Z. Wei, *Nat. Commun.*, 2024, **15**, 2501.
- 35 W. Hu, B. Huang, M. Sun, J. Du, Y. Hai, W. Yin, X. Wang, W. Gao, C. Zhao, Y. Yue, Z. Li and C. Li, *Adv. Mater.*, 2024, **37**, 2411709.
- 36 M. Imran, Z. Dai, F. Hussain, W. Xia, R. Chen, F. Wu and L. Li, *Energy Storage Mater.*, 2025, **74**, 103950.
- 37 Y. Wang, Y. Zhu, L. Zhang, H. Huang, C. Liu, B. Gao, W. Liu, H. Yan, J. Jian, Z. Huang, J. Feng, T. Yu, Z. Zou and Z. Li, *Angew. Chem., Int. Ed.*, 2025, **64**, e202506326.
- 38 N. Zhang, J. Du, N. Zhou, D. Wang, D. Bao, H. Zhong and X. Zhang, *Chin. J. Catal.*, 2023, **53**, 134–142.
- 39 Z. Yu, F. Ning, B. Li, Z. Sun, W. Chu and D. Xia, *J. Phys. Chem. C*, 2019, **123**, 18870–18876.
- 40 J. Xu, H. Jin, T. Lu, J. Li, Y. Liu, K. Davey, Y. Zheng and S.-Z. Qiao, *Sci. Adv.*, 2023, **9**, eadh1718.
- 41 C. Hu, K. Yue, J. Han, X. Liu, L. Liu, Q. Liu, Q. Kong, C. Pao, Z. Hu, K. Suenaga, D. Su, Q. Zhang, X. Wang, Y. Tan and X. Huang, *Sci. Adv.*, 2023, **9**, eadf9144.
- 42 X. Li, L. Zhou, H. Wang, D. Meng, G. Qian, Y. Wang, Y. He, Y. Wu, Z. Hong, Z.-F. Ma and L. Li, *J. Mater. Chem. A*, 2021, **9**, 19675–19680.
- 43 Y. Wang, R. Yang, Y. Ding, B. Zhang, H. Li, B. Bai, M. Li, Y. Cui, J. Xiao and Z. Wu, *Nat. Commun.*, 2023, **14**, 1412.
- 44 J. Yang, K. An, Z. Yu, L. Qiao, Y. Cao, Y. Zhuang, C. Liu, L. Li, L. Peng and H. Pan, *ACS Catal.*, 2024, **14**, 17739–17747.
- 45 S. Hao, H. Sheng, M. Liu, J. Huang, G. Zheng, F. Zhang, X. Liu, Z. Su, J. Hu, Y. Qian, L. Zhou, Y. He, B. Song, L. Lei, X. Zhang and S. Jin, *Nat. Nanotechnol.*, 2021, **16**, 1371–1377.
- 46 Q. Yu, Z. Zhang, S. Qiu, Y. Luo, Z. Liu, F. Yang, H. Liu, S. Ge, X. Zou, B. Ding, W. Ren, H. M. Cheng, C. Sun and B. Liu, *Nat. Commun.*, 2021, **12**, 6051.
- 47 X. Li, G. Chen, Y. Liu, R. Lu, C. Ma, Z. Wang, Y. Han and D. Wang, *Energy Environ. Sci.*, 2025, **18**, 4200–4209.
- 48 X. Chen, Y. Cheng, B. Zhang, J. Zhou and S. He, *Nat. Commun.*, 2024, **15**, 6278.
- 49 D. Chen, R. Yu, K. Yu, R. Lu, H. Zhao, J. Jiao, Y. Yao, J. Zhu, J. Wu and S. Mu, *Nat. Commun.*, 2024, **15**, 3928.
- 50 Y. Lin, Z. Tian, L. Zhang, J. Ma, Z. Jiang, B. J. Deibert, R. Ge and L. Chen, *Nat. Commun.*, 2019, **10**, 162.
- 51 N. Richards, L. A. Parker, J. H. Carter, S. Patisson, D. J. Morgan, N. F. Dummer, S. E. Golunski and G. J. Hutchings, *Catal. Lett.*, 2021, **152**, 213–226.
- 52 J. Lee, P. Tieu, J. Finzel, W. Zang, X. Yan, G. Graham, X. Pan and P. Christopher, *JACS Au*, 2023, **3**, 2299–2313.
- 53 S. Zuo, Z.-P. Wu, D. Xu, R. Ahmad, L. Zheng, J. Zhang, L. Zhao, W. Huang, H. Al Qahtani, Y. Han, L. Cavallo and H. Zhang, *Nat. Commun.*, 2024, **15**, 9514.
- 54 Y. Zhang, N. Cao, K. Wang, M. Yan, X. Zhang and P. Xie, *ACS Catal.*, 2025, **15**, 8833–8845.
- 55 F. Ospina-Acevedo, L. A. Albitar, K. O. Bailey, J. F. Godínez-Salomón, C. P. Rhodes and P. B. Balbuena, *ACS Appl. Mater. Interfaces*, 2024, **16**, 16373–16398.
- 56 J. Zhang, L. Xu, X. Yang, S. Guo, Y. Zhang, Y. Zhao, G. Wu and G. Li, *Angew. Chem., Int. Ed.*, 2024, **63**, e202405641.



- 57 W. Zhu, F. Yao, K. Cheng, M. Zhao, C. J. Yang, C. L. Dong, Q. Hong, Q. Jiang, Z. Wang and H. Liang, *J. Am. Chem. Soc.*, 2023, **145**, 17995–18006.
- 58 Z. Shi, Y. Wang, J. Li, X. Wang, Y. Wang, Y. Li, W. Xu, Z. Jiang, C. Liu, W. Xing and J. Ge, *Joule*, 2021, **5**, 2164–2176.
- 59 Y. Pan, X. Xu, Y. Zhong, L. Ge, Y. Chen, J. P. M. Veder, D. Guan, R. O'Hayre, M. Li, G. Wang, H. Wang, W. Zhou and Z. Shao, *Nat. Commun.*, 2020, **11**, 2002.
- 60 N. Yao, H. Jia, J. Zhu, Z. Shi, H. Cong, J. Ge and W. Luo, *Chem*, 2023, **9**, 1882–1896.
- 61 L. Wu, W. Huang, D. Li, H. Jia, B. Zhao, J. Zhu, H. Zhou and W. Luo, *Angew. Chem., Int. Ed.*, 2024, **64**, e202413334.
- 62 C. Lin, J. L. Li, X. Li, S. Yang, W. Luo, Y. Zhang, S. H. Kim, D.-H. Kim, S. S. Shinde, Y. F. Li, Z. P. Liu, Z. Jiang and J. H. Lee, *Nat. Catal.*, 2021, **4**, 1012–1023.
- 63 L. Deng, S. F. Hung, S. Liu, S. Zhao, Z. Y. Lin, C. Zhang, Y. Zhang, A. Y. Wang, H. Y. Chen, J. Peng, R. Ma, L. Jiao, F. Hu, L. Li and S. Peng, *J. Am. Chem. Soc.*, 2024, **146**, 23146–23157.
- 64 L. Deng, H. Chen, S. F. Hung, Y. Zhang, H. Yu, H. Y. Chen, L. Li and S. Peng, *J. Am. Chem. Soc.*, 2024, **146**, 35438–35448.
- 65 S. Zhao, S. F. Hung, L. Deng, W. J. Zeng, T. Xiao, S. Li, C. H. Kuo, H. Y. Chen, F. Hu and S. Peng, *Nat. Commun.*, 2024, **15**, 2728.
- 66 S. Zi, J. Zhu, Y. Zhai, Y. Hu, N. Zhang, S. Li, L. Liu, L. An, P. Xi and C. H. Yan, *Angew. Chem., Int. Ed.*, 2025, **64**, e202413348.
- 67 X. Y. Zhang, H. Yin, C. C. Dang, H. Nie, Z. X. Huang, S. H. Zheng, M. Du, Z. Y. Gu, J. M. Cao and X. L. Wu, *Angew. Chem., Int. Ed.*, 2025, **64**, e202425569.
- 68 Y. Hao, S. F. Hung, W. J. Zeng, Y. Wang, C. Zhang, C. H. Kuo, L. Wang, S. Zhao, Y. Zhang, H. Y. Chen and S. Peng, *J. Am. Chem. Soc.*, 2023, **145**, 23659–23669.
- 69 X. Zhang, Y. Zhang, B. O. Protsenko, M. A. Soldatov, J. Zhang, C. Yang, S. Bo, H. Wang, X. Chen, C. Wang, W. Cheng and Q. Liu, *Nat. Commun.*, 2025, **16**, 6921.
- 70 C. Li, C. Yuan, X. Huang, H. Zhao, F. Wu, L. Xin, X. Zhang, S. Ye and Y. Chen, *eScience*, 2025, **5**, 100307.
- 71 R. Ram, L. Xia, H. Benzidi, A. Guha, V. Golovanova, A. Garzón Manjón, D. Llorens Rauret, P. S. Berman, M. Dimitropoulos, B. Mundet, E. Pastor, V. Celorrio, C. A. Mesa, A. M. Das, A. Pinilla-Sánchez, S. Giménez, J. Arbiol, N. López and F. P. García de Arquer, *Science*, 2024, **384**, 1373–1380.
- 72 L. Wu, W. Huang, D. Li, B. Zhao, H. Zhou and W. Luo, *Angew. Chem., Int. Ed.*, 2025, **64**, e202420848.
- 73 N. Zhang, X. Liu, H. Zhong, W. Liu, D. Bao, J. Zeng, D. Wang, C. Ma and X. B. Zhang, *Angew. Chem., Int. Ed.*, 2025, **64**, e202503246.

