Volcano-type relationship between oxidation states and catalytic activity of single-atom catalysts towards hydrogen evolution

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To date, the effect of oxidation state on activity remains controversial in whether higher or lower oxidation states benefit the enhancement of catalytic activity. Herein, we discover a volcanic relationship between oxidation state and hydrogen evolution reaction activity based on Os single-atom catalysts. Firstly, a series of Os SACs with oxidation states ranging from +0.9 to +2.9 are synthesized via modifying the coordination environments, including Os-N3S1, Os-N4, Os-S6, Os-C3, and Os-C4S2. A volcano-type relation between oxidation states and hydrogen evolution activity emerge with a summit at a moderate experimental oxidation state of +1.3 (Os-N3S1). Mechanism studies illustrate that with increasing oxidation states, the adsorption of H atoms on Os is strengthened due to increased energy level and decreased occupancy of antibonding states of Os-H bond until the anti-bonding states become empty. Further increasing the oxidation states weakens hydrogen adsorption because of the decreased occupancy of Os-H bonding states. In this work, we emphasize the essential role of oxidation state in manipulating activity, which offers insightful guidance for the rational design of single-atom catalysts.

Single-atom catalysts (SACs) have received growing interests in various catalytic fields due to the utmost utilization efficiency of metal atoms1–11. In addition, distinct local structure of SACs also provides an in-depth comprehension to the intrinsic structure-performance relationship during catalytic process12–20. Notably, when metal single atoms are anchored on the supports, the metal-support interaction and the heterogeneity of the supports endow single atoms with tunable electronic properties, such as oxidation state, highest occupied molecular orbital (HOMO), lowest unoccupied molecular orbital (LUMO), spin states, etc., resulting in controllable catalytic properties21–25. For instance, Rh atom doped rutile VO2 exhibited higher NH3BH3 hydrolysis activity than Rh doped monoclinic VO2 due to the higher occupied state of single Rh atoms supported on rutile VO221. Mn-C3N4 showed high solar-driven water splitting activity, which was mainly dependent on its spin state with an optimized εc occupancy (=0.95)26.

Among these electronic properties, oxidation state is considered as a typical parameter to correlate with the catalytic activity. However, specific relationship between oxidation state and catalytic activity remains controversial in whether higher or lower oxidation states
benefit the enhancement of catalytic activity for SACs. For example, reported metallic Pt SACs presented higher activities than highly oxidized Pt SACs during CO oxidation because the adsorption strengths of CO increased as the Pt atom became more metallic27. Similarly, Pt/Fe2O3 SACs with lower oxidation state exhibited more excellent 3-nitrostyrene hydrogenation activity than other Pt/Fe2O3 SACs with higher oxidation states because lower oxidation state was more beneficial for H2 adsorption and dissociation40. On contrary, Fe-N4 SACs with the highest oxidation state displayed higher benzene oxidation reaction activity compared to Fe-N4 and Fe-N2 Ca SACs because Fe-N4 facilitated the generation and activation of the vital intermediate O=Fe=O species, enhancing the ability to activate C-H bond29. Considering the controversial role played by oxidation state, it is necessary to clarify the specific role and demonstrate the underlying catalytic mechanism.

In this work, we select osmium (Os) as research object to fully discuss the relationship between oxidation state and catalytic activity due to its rich valence states (−2 to +8). Firstly, we successfully synthesized atomically dispersed Os sites with different oxidation states (from +0.9 to +2.9) by changing coordination environments, including Os-N5S0, Os-N4S1, Os-N3S2, and Os-C2S2. It was found that a volcano curve was obtained between oxidation state and hydrogen evolution reaction (HER) activity based on various Os SACs. Specifically, Os-N5S0 SACs with a moderate experimental oxidation state (+1.3) displayed the best acidic HER performance compared to other SACs, which only needed 22 mV at 10 mA cm−2. Further mechanism studies disclosed that with increasing oxidation states (decreasing valence electron numbers), the adsorption of H atoms on Os was strengthened due to the increased energy level and decreased occupancy of anti-bonding states of Os-H bond until the anti-bonding states become empty. Further increasing the oxidation states weakened hydrogen adsorption because of the decreased occupancy of Os-H bonding states near the Fermi level. This work provided Os SACs for HER and revealed the essential role of oxidation state in manipulating activity, which offers essential guidance for the rational design of SACs.

Results

Synthesis and characterization of Os SACs

To fully discuss the relationship between oxidation state and catalytic activity, we synthesized atomically dispersed Os catalysts with controllable coordination environments and verified the proposed regulation mechanism of coordination configuration to optimize HER catalytic activity. The synthetic strategy of atomically dispersed Os sites supported on N and S co-doped carbon materials (Os/CNS) is shown in Fig. 1a. First, N and S atoms were introduced into carbon black by adopting urea and 2,2'-bithienyl as the nitrogen and sulfur sources, respectively. After that, Pluronic F127 was fully mixed with CN sample in 30 mL of deionized water. Then, osmium trichloride (OsCl3) solution was slowly added into above suspension. After stirring for 6 h, the mixture was centrifuged and dried in a vacuum drying oven. Finally, the dried products were calcined in a tubular furnace at 400 °C and atomically dispersed Os/CNS sample was successfully fabricated after washing with acid solution for 12 h. Moreover, other atomically dispersed Os single atoms supported on different carriers, such as pure carbon black (C), N doped graphite carbon (CN), and S doped graphite carbon (CS), were also prepared by changing the doping heteroatom in carbon support while keeping other conditions unchanged.

To clearly obtain the atomic information of these catalysts, aberration-corrected high-angle annular dark-field scanning transmission electron microscopy (AC-HAADF-STEM) was employed. Figure 1b, c exhibited the AC-HAADF-STEM images of Os/CNS in different scale bars. It is clear that atomically dispersed Os sites were uniformly anchored on the CNS support. Subsequently, aberration-corrected TEM image of Os/CNS was shown in Fig. 1d. It is obvious that no large nanoparticles were observed. The evident lattice stripe of CNS support revealed its good crystallinity, which indicated a superior electronic conductivity. The inset of Fig. 1d corresponded to the selected area electron diffraction (SAED) of Os/CNS sample. No diffraction spots further disclosed the absence of Os particles. Then, Fig. 1e directly displayed the homogeneous distribution of Os, S, and Os elements in Os/CNS sample. In addition, to regulate the coordination structure of Os active sites, we changed the CNS into CN support while kept other synthesis conditions remained. Similarly, Os/CN displayed a uniform distribution of isolated Os atoms on the CN carrier (Fig. 1f, g). The aberration-corrected TEM image and SAED pattern verified the polycrystalline characteristics of the CN carrier (Fig. 1h). Meanwhile, the C, N, and Os elements were also uniformly dispersed across the elemental mapping (Fig. 1i). Moreover, as a control sample, Os single atom supported on CS carrier was also successfully synthesized. Obviously, Fig. 1j, k clearly revealed the even dispersion of Os atoms in Os/CS. The aberration-corrected TEM image of Os/CS also confirmed the uniform distribution of Os single atoms on CS carrier (Fig. 1j). The SAED pattern of Os/CS sample was the same as those of Os/CNS and Os/CN (inset in Fig. 1j). Afterwards, Fig. 1m exhibited the uniform distribution of Os, C, and S atoms for Os/CS. To fully discuss the influence of coordination environments, we also fabricated atomically dispersed Os/COS/2 and Os/C catalysts (Supplementary Figs. 1a–c and 2a–e). Furthermore, elemental mapping tests of Os/CS-2 and Os/C suggested the homogeneous distribution of corresponding elements on entire structure (Supplementary Figs. 3d, 4e). Additionally, the low-resolution STEM images of Os/CNS, Os/CN, Os/Cs, and Os/COS-2 were characterized (Supplementary Fig. 3a–d). The basic characteristics of pure carbon-based carriers were also completed (Supplementary Figs. 4–11). Besides, Os nanoparticles (NPs) were also synthesized for comparison (Supplementary Fig. 12a, b). As shown in Fig. 2a, it could be found from Raman spectra that the values of /I changed from 1.20 for C carrier to 1.31, 1.27, 1.29, and 1.42 for CN, CS, CS-2, and CNS carriers, respectively. Thus, the surface defects obviously increased after doping N and S atoms into carbon black due to the enlarged values of /I. The increased level of defects could contribute to the anchoring of metal atoms. According to the inductively coupled plasma-optical emission spectrometer (ICP-OES) test, the mass loading of Os for Os/C, Os/CN, Os/CNS, Os/CS, and Os/CS-2 was 3.17%, 4.56%, 4.61%, 3.80%, and 4.03%, respectively. Then, to investigate more information of Os single atom supported on different carbon materials, X-ray diffraction (XRD) and N2 adsorption-desorption isotherms tests were applied. The XRD results of different Os SACs were almost the same as that of pure carbon black, disclosing no corresponding diffraction peak of Os nanoparticles (Fig. 2b and Supplementary Fig. 13). The XRD results of different SACs were consistent with SAED, suggesting the successful preparation of various Os SACs. In addition, the nitrogen adsorption-desorption isotherm of different samples demonstrated that the specific surface areas of Os/C, Os/CN, Os/CNS, Os/CS, and Os/CS-2 were 1418.5, 1389.3, 1370.3, 1382.7, and 1325.9 m2/g, respectively (Supplementary Fig. 14a–e). The larger surface area exposed more active centers for water splitting and contributed to the adsorption and desorption of reactants and products.

X-ray photoelectron spectroscopy (XPS) was adopted to unveil the chemical composition and structure of each element in single atom samples31. In the Os 4f XPS spectra, the Os 4f7/2 and Os 4f5/2 peaks of Os/C were at 51.2 and 54.1 eV, respectively, which were higher than those (50.3 and 53.1 eV, respectively) of Os nanoparticles (Fig. 2c and Supplementary Fig. 15). The oxidation state of Os nanoparticles is nearly 0. Then, the binding energies of Os in Os/C (51.4 and 54.3 eV) were moved to larger values compared to Os/C sample (Fig. 2c). For Os/CNS, XPS survey spectrum firstly revealed the co-existence of Os, N, S, and C elements (Supplementary Fig. 16).
Then, the two characteristic peaks of Os $4f_{7/2}$ and Os $4f_{5/2}$ were obtained at 51.6 and 54.2 eV, respectively (Fig. 2c). Additionally, the binding energies of Os in Os/CN were located at 52.2 and 54.8 eV. The locations of Os $4f_{7/2}$ and Os $4f_{5/2}$ for Os/CS-2 were centered at 52.6 and 55.5 eV, respectively. Thus, the oxidation state of different Os SACs increased in the order of Os/C < Os/CS < Os/CNS < Os/CN < Os/CS-2 because of the positive movement of corresponding binding energies, suggesting that doped heteroatoms effectively regulated the oxidation state of Os sites. Moreover, Fig. 2d displayed the S 2p spectra of Os/CS, Os/CNS, and Os/CS-2. Importantly, the S 2p curve of Os/CNS can be divided into three peaks at 161.6 eV (Metal-S bond), 163.0 eV (C-S-C), and 164.2 eV (C-S-C). In addition, S 2p XPS spectra of Os/CS and Os/CS-2 also exhibited the same three peaks. The XPS spectrum of S 2p for pure CNS revealed no Metal-S bonds (Supplementary Fig. 17). Thus, it can be inferred that Os in Os/CNS, Os/CS, and Os/CS-2 directly bonded with S atom, suggesting different
coordination environments of these three samples from the traditional metal-N structures. In addition, the N 1s spectra of Os/CNS and Os/CN displayed four peaks at 398.5, 399.5, 400.8, and 401.6 eV, corresponding to pyridinic N, Metal coordinated N, pyrrolic N, and graphitic N, respectively (Supplementary Fig. 18). The successful doping of different heteroatoms precisely regulated the oxidation states of active centers, which should change the catalytic performance of atomically dispersed catalysts.

X-ray absorption fine structure (XAFS) analyses, including X-ray absorption near-edge structure (XANES) and extended X-ray absorption fine structure (EXAFS), were conducted to detect the specific coordination environments of Os SACs. The Os L-edge XANES spectra of Os SACs, OsO2, and Os nanoparticles were shown in Fig. 2e. The oxidation states of catalysts estimated from the integrated area under the white-line intensity of Os L-edge was displayed in Supplementary Fig. 19. The oxidation state of Os single atoms increased in the order of Os/C ( + 0.9) < Os/CS ( + 1.1) < Os/CNS ( + 1.3) < Os/CN ( + 1.9) < Os/CS-2 ( + 2.9), revealing that doped heteroatoms effectively regulated the oxidation state of Os sites (Supplementary Fig. 19). It was consistent with the analyses of XPS results. Furthermore, EXAFS results were collected to further acquire precise coordination configuration, including coordination numbers and bond lengths (Fig. 2f). The Fourier-transformed (FT) k^3-weighted EXAFS spectrum of Os/CN SAC exhibited a prominent peak at 1.68 Å, which was attributed to the Os-N bonds (Fig. 2f). Moreover, the first peak for Os/CNS was wider and moved positively compared to Os/CN, implying changes in coordination structure due to the doping of S atoms. Meanwhile, as displayed in Fig. 2f, the coordination environments of Os/CS-2, Os/CS, and Os/C were different from Os/CN with different shaped FT k^3-weighted EXAFS spectra. Besides, Fig. 2f further demonstrated no obvious Os-Os bonds in Os/CNS, Os/CS, Os/CN, Os/CNS-2, and Os/C compared to Os NPs. Afterwards, quantitative EXAFS fitting analyses of different Os SACs and Os NPs were used to confirm the detailed local coordination environments (Fig. 2g–i and Supplementary Fig. 20a–c). The corresponding fitting parameters were listed in Supplementary Table 1. Notably, Fig. 2g manifested that the main peak at 1.68 Å for Os/CN could be satisfactorily interpreted as Os-N first shell coordination and the corresponding coordination number was 4 with a bonding distance of 2.04 Å (Supplementary Table 1). Moreover, Os/CNS sample included Os-S and Os-N bonds in the first shell coordination (Fig. 2h) and the specific coordination numbers of Os-N and Os-S bonds were 3 and 1, respectively, implying that Os-N6S1 coordination structure was successfully constructed (Supplementary Table 1). The introduction of S atoms could effectively change the dispersion of electrons around active sites, resulting in precise regulation of oxidation state. Obviously, the best-fitting analysis of Os/CS further revealed the first shell coordination of Os-C at 1.65 Å and Os-S at 2.05 Å with a coordination number of 4 and 2, respectively (Fig. 2i and Supplementary Table 1). In addition, Os-S6 and Os-C3 coordination configurations can also be synthesized when Os is supported on CS-2 and pure C, respectively (Supplementary Fig. 20d–f). To further distinguish the formation of Os-C/N coordination in Os/CN and Os/CNS, the soft X-ray absorption spectroscopy at N K-edge and C K-edge of catalysts was applied. Based on C K-edge spectra for Os/CN and Os/CNS, the peak C1 (285.7 eV) was derived from C-C τ* (ring) excitations, the peak C2 (288.8 eV) was originated from C-N-C, and peak C3 (293.0 eV)
corresponded to C-C δ’ (ring), respectively (Supplementary Fig. 21). Importantly, C1, C2, and C3 peaks of Os/CN and Os/CNS were almost similar as that of pure carrier, indicating Os atoms did not coordinate with C atoms in carrier. In addition, N K-edge spectra for Os/CN and Os/CNS disclosed that four typical peaks (N1, N2, N3, and N4) are directly observed, which corresponded to pyridinic N, Metal coordinated N, pyrrolic N, and graphitic N species, respectively (Supplementary Fig. 22). Therefore, it further confirmed the Os atoms bonded with N atoms in Os/CN and Os/CNS, which was consistent with the analyses of XPS results. Thus, Os/CS-2, Os/CN, Os/CNS, Os/CS, and Os/C can also be regarded as Os-N4, Os-N1, Os-N2, Os-C-S2, and Os-C3 SACs in this study, respectively. Five different coordinated structure catalysts were precisely constructed in this work, which could be used to disclose the relationship between coordination environments and catalytic activity. Additionally, wavelet transform (WT) was further conducted to disclose the Os L-edge EXAFS oscillations of different Os SACs (insets in Fig. 2g–i and Supplementary Fig. 20). It is obvious that WT analyses of Os-N1, Os-N2, and Os-C-S2 exhibited the maximum intensity at approximately 4.0, 5.8, and 5.0 Å−1, respectively, which were quite different from the Os nanoparticles (13 Å−1, Supplementary Fig. 20F). Meanwhile, Os-C3 and Os-S2 also displayed the maximum intensity at nearly 4.4 and 5.9 Å−1, respectively, which were different from Os-N1 and Os-C2S2. Therefore, Os sites with different coordination environments were confirmed as atomically distributed in this work.

**Electrochemical performance of Os SACs for HER**

To disclose the influence of coordination structure on catalytic performance, we measured the electrochemical HER activities of Os catalysts in 0.5 M H2SO4 \( \text{Os/CNS} \), \( \text{Os/C} \), \( \text{Os/CS} \), \( \text{Os/CS-2} \), and 20% Pt/C required overpotentials of 22, 132, 45, 63, 233, and 13 mV to get a current density of 10 mA cm\(^{-2}\) (Fig. 3a). The onset potentials (at current density of 1 mA cm\(^{-2}\)) for Os/CNS, Os/C, Os/CS, Os/CS-2, and 20% Pt/C corresponded to 5, 47, 11, 18, 129, and 3 mV, respectively. Therefore, the HER activities of different Os SACs in this work were as follows: Os/CNS > Os/CS > Os/CS-2 > Os/C > Os/CS-2. Moreover, the LSV curves in acidic media identified that the doping of heteroatoms (N and S) effectively regulated the HER performance of SACs. Meanwhile, Os sites served as the main active centers for HER while carbon supports stabilized the metal atoms because pure carbon materials showed poor performance for HER in acidic media (Supplementary Fig. 23). In addition, the fitting Tafel slopes in acidic medium were illustrated in Fig. 3b. The Tafel slopes of Os/CNS, Os/C, Os/CS, Os/CS-2, and 20% Pt/C were 41, 133, 56, 67, 137, and 30 mV dec\(^{-1}\), respectively, interpreting the fastest HER kinetics of the Os-N1S1 among single atom samples with different coordination structures. Importantly, Os/CNS, Os/C, Os/CS, Os/CS-2 involved in Volmer-Heyrovsky mechanism. Meanwhile, at overpotential of 50 mV, the turnover frequency (TOF) values of Os/CNS, Pt/C, Os/C, Os/CS, Os/CS, and Os/CS-2 were 10.55, 6.49, 0.62, 3.60, 2.66, and 0.34 s\(^{-1}\).
respectively (Fig. 3c). TOF plots further revealed that Os/CNS illustrated highest intrinsic catalytic activity for HER in acidic medium because it displayed the largest TOF values under the same potentials in comparison with other SACs in this work (Fig. 3c). Moreover, Os/CNS also showed better HER activity compared to some reported catalysts (Supplementary Table 2). Additionally, electrochemical impedance spectroscopy (EIS) measurement in acidic medium was employed to provide further insight into the electrical conductivity of SACs in acidic media. As illustrated in Fig. 3d, Os/CNS displayed the smallest values of charge transfer resistance ($R_{ct}$) in acidic medium compared to other SACs, indicating its fastest HER kinetics. Then, the electrochemical double-layer capacitance ($C_{dl}$) was also used to estimate the electrochemically active surface area of electrocatalysts (Supplementary Figs. 24a-f, 25). Specially, the $C_{dl}$ value (29.2 mF cm$^{-2}$) of Os/CNS was larger than that of other SACs, suggesting that it exhibited larger electrochemically-active surface area (ECSA) because ECSA is positively correlated with $C_{dl}$. Moreover, stability is also an important parameter to access the HER performance of electrocatalysts. The long-term durability measurements of Os/CNS sample, including chronoamperometric curves and accelerated linear potential sweeps, were performed in 0.5 M H$_2$SO$_4$ solution. It is obvious that the current density of Os/CNS sample displayed negligible decrease after more than 50 h test (Fig. 3e). However, ECSA of Os/CNS also declined negligibly in acidic condition after 3000 cycles (inset of Fig. 3e). Os/C, Os/CN, Os/CS, and Os/CS-2 also performed excellent stability, as shown in Supplementary Fig. 26a-d. Furthermore, Supplementary Fig. 27a-d manifested that Os/CNS showed excellent structural stability due to the uniform distribution of Os atoms on carrier after HER. In Fig. 3f, oxidation states analyzed from experiments were used to correlate with HER overpotential in acidic medium. It was found that different coordination environments induced various oxidation states of Os single-atom, which directly affected catalytic performance in a volcanic curve relation. Obviously, Os/CNS showed highly active catalytic activity owing to its moderate oxidation state (+1.3). Experimental discovery verified that the oxidation state of isolated metal atom illustrated a volcanic curve with HER activity. The fundamental understanding on the role of oxidation state in coordination-dependent activity offers valuable guidance for a rational design of SACs through coordination environment engineering.

Theoretical studies between oxidation state and HER activity for Os SACs

According to the local coordination environments of graphene-supported metal SACs reported by previous works, density functional theory (DFT) calculations were employed to explore the origin of coordination-dependent catalytic activity among Os SACs. To further precisely identify the local coordination environment of each Os SACs, we constructed all possible configuration according to quantitative EXAFS fitting analyses, as shown in Supplementary Fig. 28a-k. Moreover, we also performed DFT calculations to explore thermodynamics preference of bonds formation between Os-C and Os-N in Os/CN and Os/CNS, according to energy change of the process (Supplementary Fig. 29a, b). As shown in Supplementary Fig. 29a, for Os/CNS with N source during synthesis, the endothermic process (positive value of energy change) for Os single atom, from forming Os-N$_2$S to Os-C$_2$N$_2$S$_2$ or Os-C$_2$N$_2$S local coordination, indicating that Os single atoms tended to form Os-N bond without Os-C coordination. Likewise, as for Os/CN with N source during synthesis, Os single atoms tended to form Os-N$_2$ rather than Os-C$_2$N$_2$S or Os-C$_2$ local coordination (Supplementary Fig. 29b). Thus, we did not consider candidate configuration with Os-C bonds in Os/CN and Os/CNS. The bond lengths between Os and nearest neighbor were summarized in Supplementary Table 3, which can be compared with experimental EXAFS results (Supplementary Table 1). The oxidation states of Os single-atom were estimated for all possible models by normalizing valence electron number (by Bader charges analysis) to those of bulk compounds (Os, Os$_2$O$_3$, and OsO$_2$) with known oxidation states (Supplementary Fig. 30). The results were shown in Supplementary Table 3, which could confirm whether it was in good agreement with the relative order derived from XPS and XANES analyses. We also checked the stability of the proposed possible configurations of Os SACs against metal atom aggregation, in order to confirm that the proposed possible catalysts are experimentally feasible (Supplementary Fig. 32a, b and supplementary method). All configurations meeting oxidation state, bond length and stability against sintering were labeled in red color in Supplementary Table 3, which were selected for the further investigations on HER theoretical activity.

The adsorption free energy of hydrogen, $\Delta G_{H^*}$, has been widely considered as a useful descriptor for acidic HER catalytic activity. In order to explore whether oxidation state acts as a reliable electronic-structure-involved descriptor, the relationship between oxidation state of Os single-atoms and $\Delta G_{H^*}$ for the corresponding Os SACs was built. It was found that diverse coordination environments of Os SACs brought about distinguishable oxidation states (Fig. 4a). It suggested that the electronic hybridization between center single-atom and proximal atoms from the support modified the valence electron occupation through electronic metal-support interaction (EMSI). Among all considered Os SACs, their $\Delta G_{H^*}$ were zero. For Os SACs with stronger binding strength of H, $H^*$ can be more easily captured from the electrolyte and combined with adsorbed $H^*$ to form gaseous $H_2$. What’s more, a volcanic correlation existed between oxidation state and $\Delta G_{H^*}$. Since Os-N$_2$S$_2$ possessed the moderate Os oxidation state of +1.9, it displayed the strongest binding strength of H and might accordingly show better acidic HER performance than other Os SACs with distinct Os oxidation states. Figure 4a suggested that the activity of Os SACs for acidic HER might be controlled by oxidation state as it adjusted $\Delta G_{H^*}$. Hence, oxidation state has potential to be a reliable descriptor to predict acidic HER activity of Os SACs.

DFT calculations were further employed to explore the role of oxidation state in coordination-dependent catalytic kinetics among Os SACs. According to the characteristic of Tafel slope in Fig. 3b, we inferred that Os SACs may prefer Volmer-Heyrovsky mechanism. As for the Volmer-Heyrovsky mechanism in acidic HER, the reaction pathway could be described as a three-state diagram consisting of an initial free $H_2O^-$, an adsorbed intermediate $H_*$, and gaseous $H_2$ molecule as the final product. Figure 4b showed the free energy diagram towards acidic HER of Os SACs with various coordination environments, of which structure images for hydrogen transfer process were shown in Supplementary Fig. 32a-f. The adsorption free energies of adsorbed intermediates were displayed in Supplementary Tables 4, 5. From the kinetic viewpoint, the highest reaction barrier among hydrogen transfer process is known as the kinetics of rate-determining step to describe the HER catalytic activity. $\Delta G_{H^*}$ was applied to correlate with energy barrier of atomic hydrogen capture/desorption process (Fig. 4c). When $\Delta G_{H^*} > 0$, $H^*$ can be more easily captured from the electrolyte and combined with adsorbed $H$ to form gaseous $H_2$ with higher $\Delta G_{H^*}$. Therefore, the acidic HER kinetics of Os SACs was controlled by oxidation state which adjusts $\Delta G_{H^*}$. We also applied microkinetics model and converted results in free energy diagram into polarization curve simulation (Supplementary Fig. 33). Supplementary Fig. 34 and Fig. 4d directly displayed an obvious volcanic diagram between the energy barrier (or theoretical overpotential in 10 mA/cm$^2$) and oxidation state of Os SACs. Therefore, Os-N$_2$S$_2$ with a moderate oxidation state demonstrates the lowest reaction barrier of rate-determining step and accordingly lowest overpotential among the five Os SACs in acidic HER. Besides, we also considered the possibility of Volmer-Tafel mechanism. According to the free energy diagram in Supplementary Fig. 35a, the free energy change of potential-determining step through Volmer-Tafer mechanism still showed volcano relationship with oxidation state among diverse Os SACs in...
was attributed to the adjusted energy-level of coordination-dependent activity of isolate Os sites towards acidic HER atoms. Thus, the oxidation state of isolated sites directly affected induced by oxidation state (or valence electron number) of Os single atoms. The integral of orbital which is the only unsaturated orbital of Os element. The d-band centers of Os SACs with different coordinations were derived from their projected density of state (PDOS) for Os d-orbitals (Fig. 5d). The change of hydrogen adsorption strength was determined by the anti-bonding states of Os-H bond at unsaturated occupancy (Fig. 5c). Theoretical overpotential (at 10 mA/cm²) as a function of oxidation state for different Os SACs.

Supplementary Fig. 3b, where Os-N₃S₁ displayed highest HER activity due to the moderate oxidation state.

To explore the intrinsic regulation mechanism of oxidation state for ΔG_H from a point of electron structures, we turned to focus on d-orbital which is the only unsaturated orbital of Os element. The d-band centers of Os SACs with different coordinations were derived from their projected density of state (PDOS) for Os d-orbitals (Fig. 5a). The relative order of integration of PDOS in d-orbital under Fermi energy, representing the degree of valence electron occupation in d-orbital, among Os SACs with different coordination (Supplementary Table 6), was consistent with that derived from Bader charge analysis. As presented in Fig. 5b, the less occupied d-orbital with fewer valence electrons (higher oxidation state) resulted in a higher d-band center, which yielded an increasing energy level of the molecular orbital of Os-H bond from partially occupied anti-bonding states to bonding state with unsaturated occupancy (Fig. 5c). The change of hydrogen adsorption strength was determined by the anti-bonding states of Os-H bond at first. The decreasing number of valence electrons induced anti-bonding states with higher level and lower occupancy, leading to a stronger Os-H interaction. When valence electron number kept decreasing, the hydrogen adsorption strength converted to be affected by bonding state mainly near the Fermi level. Consequently, the upshift of bonding orbital resulted in a decreasing occupancy and receded H⁺ adsorption strength. The value of integral crystal orbital Hamilton populations (ICOHP), reflecting Os-H bond strength, showed a volcanic curve relationship with d-band center (Fig. 5d). It was consistent with the analysis on Os-H bond strength (ΔG_H). Finally, coordination-dependent activity of isolate Os sites towards acidic HER was attributed to the adjusted energy-level of d-orbital, which was induced by oxidation state (or valence electron number) of Os single atoms. Thus, the oxidation state of isolated sites directly affected atomic hydrogen adsorption and molecular hydrogen formation.

Discussion

In summary, this work discloses that the HER performance of Os SACs with different coordination environments was closely correlated to the oxidation state of metal atoms from both experimental results and DFT calculations. Firstly, we prepared various Os SACs with different oxidation states by regulating coordination environments, including Os-N₃S₁, Os-N₄, Os-C₄S₂, Os_S₆, and Os-C₃. Notably, experimental results confirmed that a volcanic relation was obtained between the oxidation state of different Os SACs and HER catalytic activity. Moreover, Os-N₃S₁ with a moderate experimental oxidation state of +1.3 showed the most accelerated reaction kinetics and promising catalytic activity for acidic HER. Further intrinsic analyses revealed that the valence electron number of isolated Os atom, macroscopically reflected by oxidation state, was responsible for adjusting the energy-level of d-orbital to affect the atomic hydrogen adsorption and molecular hydrogen formation in acidic HER. Finally, this study not only offers a series of Os SACs with various coordination structures for hydrogen production, but also provides a fundamental understanding on the role of oxidation state in coordination-dependent activity. It may be broadened to other SACs and offers valuable guidance for rational design of SACs through engineering oxidation states.

Methods

Synthesis of different carbon materials

1 g activated carbon and 5 g urea were mixed in 80 mL deionized water. Then, the mixture was heated to 180 °C for 12 h in a Teflon-lined stainless steel autoclave (100 mL). After that, the mixture was dried under 50 °C and fully ground. Besides, the mixture was calcined in a tube furnace at 800 °C for 2 h under N₂ atmosphere. Finally, N doped carbon material was successfully synthesized (defined as CN). 1 g activated carbon and 3 g 2, 2-bithiophene were fully ground. Then, the mixture was calcined in a tube furnace at 800 °C for 2 h under N₂ atmosphere. Finally, S-doped carbon material was successfully synthesized (defined as CS). Activated carbon (1 g) and urea (5 g) were mixed in deionized water (80 mL). Then, the mixture was heated to 180 °C for 12 h in a Teflon-lined stainless steel autoclave (100 mL). After that, the mixture was dried at 50 °C and fully ground with 3 g of 2, 2-bithiophene. Next, the mixture was calcined in a tube furnace at
800 °C for 2 h under N$_2$ atmosphere. Finally, N, S co-doped carbon material was successfully synthesized (defined as CNS).

**Synthesis of atomically dispersed Osmium catalysts**

C/CS/CN/CNS (30 mg) and Pluronic F127 (30 mg) were fully mixed in deionized water (30 mL). Then, 0.015 mmol OsCl$_3$ was dissolved in 5 mL deionized water. After that, the OsCl$_3$ solution was doped into above suspension. Next, the mixture was fully mixed for 6 h and was centrifuged at 1300 g (acceleration of gravity) for 7 min. Then, the obtained products were washed with deionized water three times and were dried in a vacuum drying oven. Moreover, these samples were calcined at 400 °C and the prepared catalysts should be washed including sample were calcined at 400 °C under N$_2$ atmosphere for 2 h, respectively. Then, the prepared catalysts were dealt with hydrochloric acid for 12 h. Finally, Os/C, Os/CN, Os/CS, and Os/CNS were obtained, respectively.

To obtain Os/CS-2 sample, 30 mg CS and 30 mg F127 were fully stirred in 30 mL water. Then, 0.015 mmol OsCl$_3$ was dissolved in 5 mL deionized water. After that, the OsCl$_3$ solution was doped into the above mixture. Next, the mixture was fully mixed for 6 h. The obtained products were washed with deionized water for three times and were dried in a vacuum drying oven. Moreover, a porcelain plate containing sulfur powder (upstream) and a porcelain plate including sample were calcined at 400 °C under N$_2$ atmosphere for 2 h, simultaneously. Then, the prepared catalysts were dealt with hydrochloric acid for 12 h. Finally, atomically dispersed Os/CS-2 was synthesized.

**Synthesis of Osmium nanoparticles supported on CNS**

30 mg CNS was uniformly dispersed in water (20 mL) under magnetic stirring. Then, 0.06 mmol OsCl$_3$ was dissolved in 5 mL deionized water. After that, the OsCl$_3$ solution was doped into the beaker which contained CNS. Next, the mixture was fully mixed for 6 h and was centrifuged at 1300 g for 7 min. Then, the obtained product was dried in a vacuum drying oven. Finally, the obtained products were calcined in tube furnace under N$_2$ atmosphere at 600 °C for 2 h.

**Characterizations**

The morphology and structure of nanomaterials were observed on a H800 transmission electron microscope (TEM, JEM-2100F) which is coupled with energy dispersive X-ray spectroscopy (EDS) with an acceleration voltage of 200 kV. Besides, the HAADF-STEM images and EDS elemental mapping of electrocatalyst were carried out in a JEM ARM-200 microscope at 200 kV, equipped with a probe spherical aberration corrector. The X-ray diffraction (XRD) was recorded on XPert PRO MPD (DS) instrument to detect the phase composition of samples. Besides, the surface characteristics of materials were tested by X-ray photoelectron spectroscopy (XPS) carried out on ESCALAB 250 Xi system with an Mg-Ka source. The X-ray absorption fine structure spectra (Os L-edge) were collected at 1W1B station in Beijing Synchrotron Radiation Facility (BSRF). The storage rings of BSRF were operated at 2.5 GeV with a maximum current of 250 mA. Using Si (111) double-crystal monochromator, the data collection was carried out in transmission mode using ionization chamber. All spectra were collected in ambient conditions. The soft XANES spectra (Os K-edge) were measured at beamline BL12B of National Synchrotron Radiation Laboratory (NSRL).

**Electrochemical measurements**

The electrochemical measurements were carried out in a three-electrode system controlled by CHI 760E (CHEN HUA, China) electrochemistry workstation. The obtained electrocatalyst (5 mg) were mixed with water (0.2 mL), ethanol (0.8 mL) and Na$_2$SO$_4$ (15 µL). A homogeneous ink was gained after sonication for 30 min. Besides, a graphite rod was served as counter electrode. Saturated calomel electrode was used as the reference electrode. Then, 15 µL of catalyst ink was deposited on glassy-carbon electrode to obtain working electrode after the solvent was dried naturally. In this study, linear sweep voltammetry (LSV) curves for HER were carried out in 0.5 M H$_2$SO$_4$ media with a scan rate of 5 mV s$^{-1}$. The electrochemical impedance spectroscopy (EIS) tests were recorded with frequency from 0.1 to 100000 Hz.
Calculation of turnover frequency (TOF)

TOF per metal site of the catalysts was shown according to the following equation:

\[
\text{TOF} = \frac{\text{J}}{\text{2nF}} 
\]

where J is the current density (A cm\(^{-2}\)) from the LSV measurement, n is the number of active sites per geometric area (mol cm\(^{-2}\)), and F is the Faraday constant (96485.33 C mol\(^{-1}\)).

\[
\text{XAFS analysis}
\]

The acquired EXAFS data were processed according to the standard procedures using the ATHENA module implemented in the IFEFFIT software packages. The k\(^1\)-weighted EXAFS spectra were obtained by subtracting the post-edge background from the overall absorption and then normalizing with respect to the edge-jump step. Subsequently, k\(^1\)-weighted χ(k) data of Os L\(_\text{II}\) edge were Fourier transformed to real (R) space using a hanning windows (dk = 1.0 Å) to separate the EXAFS contributions from different coordination shells. To obtain the quantitive structural parameters around central atoms, least-squares curve parameter fitting was performed using the ARTEMIS module of IFEFFIT software packages.

\[\text{DFT computational details}\]

All DFT calculations with consideration of spin-polarization were conducted by using first-principle calculations, as implemented in the Vienna ab initio Simulation Package (VASP) code\(^{47,48}\). Meanwhile, we used the Perdew-Burke-Ernzerhof (PBE) functional within the generalized gradient approximation (GGA) to describe the exchange correlation energy\(^{49}\). We used the projector augmented wave (PAW) pseudo-potentials to depict the interaction between valence electron and ionic cores\(^{50}\). The Kohn-Sham equations was expanded in a plane wave basis set with cutoff energy of 500 eV after a series of test. The electronic orbital overlaps were determined in light of gaussian smearing of 0.05 eV during the geometry optimization and for the total energy computations, whilst a tetrahedron method with Blöchl corrections was applied in the accurate electronic structure calculations. The convergence threshold for the iteration in self-consistent-field (SCF) was set to be 10\(^{-6}\) eV, and that for geometry optimizations by BFGS algorithm was set to be 0.01 eV/Å on maximum force component. All of the Os SACs supported on the graphene models are constructed based on the graphene basal plane model with the supercell of 5 × 5 (12.30 Å × 12.78 Å), with a large vacuum slab of 15 Å inserted in z-direction for surface isolation to prevent interaction between two neighboring surfaces. The Monkhorst-Pack meshes of 3 × 3 × 1 k-point samplings in the Brillouin zones were used for these models. Denser k-points (12 × 12 × 1) were used for the electronic structure calculations. Considering that the solvent may influence the adsorption free energy, we employed VASPsol to include the implicit solvation effect in the calculations\(^{51}\). The vibrational frequencies of free molecules and adsorbates, which are needed to determine zero-point energies and vibrational entropies, were calculated by using the phonon modules in VASP 5.3 code. The reaction free energy calculation of HER on Os single atom catalysts supported on graphene were calculated according to electrochemical framework of calculation hydrogen electrode developed by Nørskov, which can be referred to our previous works\(^{52}\). The climbing-image nudged elastic band (CI-NEB) method was used to determine transition states and reaction paths. Through vibrational frequency analyses, all transition states have been verified by ensuring that they have only one non-negligible imaginary vibrational frequency.

\[\text{Microkinetics model and polarization curve simulation}\]

The Volmer-Heyrovsky pathway are listed by the following equations:

\[
\begin{align*}
* + H^+ + e^- & \xrightleftharpoons{\text{Ki}} H^* + H_2 \\
H^* + e^- & \xrightleftharpoons{\text{K2}} \text{H}_2 + * \\
\end{align*}
\]

where the adsorption site of hydrogen is denoted by Asterisks. Note that Eqs. (2) and (3) are both electrochemical step. Based on the above reduction step, we can gain the rate equations along with the site conservation condition:

\[
\frac{\partial \theta}{\partial t} = k_1 \theta - k_2 \theta_{\text{H*}} - k_3 \theta_{\text{H2}} + k_{-2} \theta_{\text{H2*}} 
\]

where θ is the coverage of the species. The coverage of free sites may be approximated by an effective Langmuir isotherm: θ\(_\text{H*}\) = \(\theta_0 - k_i \theta_{\text{H*}}\). The rate equation can be solved numerically at steady state, and further infer the turn over frequency (TOF). For electrochemical step, the equilibrium constant (K\(_i\)), which is associated with the reaction potential (U vs. RHE), can be expressed by

\[
K_i = \exp \left( \frac{eU - U_i}{k_BT} \right) 
\]

where \(U_i\) is the reversible potential of step i deduced by \(U_i = -\Delta G_i/e\). And the \(k_i\) is written as:

\[
k_i = A_i \exp \left( - \frac{E_{a,i}}{k_BT} \right) \exp \left( - \frac{e\beta_i(U - U_i)}{k_BT} \right) 
\]

where \(A_i\) is an effective pre-exponential factor taken as 10\(^7\) s\(^{-1}\), and the \(\beta_i\) is the symmetric factor taken to be 0.5. \(E_{a,i}\) is energy barrier of elementary step. Moreover, the rate constants for all the reverse reaction (\(k_{-i}\)) can be expressed by:

\[
k_{-i} = \frac{k_i}{K_i} 
\]

Finally, the current density (J) can be gained by:

\[
J = -2e\rho\text{TOF}_{\text{H2}} 
\]

where e is the elementary charge and ρ is the surface density of active sites.

\[\text{Data availability}\]

All data is available in the main text or the supplementary information. Source data are provided with this paper.

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D.C. designed and performed the experiments, carried out the analysis; 
H.X. performed the DFT calculations; D.C. and H.X. wrote the original 
manuscript; H.L. and C.F. revised the manuscript; J.Z. and D.C. super-
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Competing interests
The authors declare no competing interests.

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