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Modulating paired Ir–O–Ir via electronic perturbations of correlated Ir single atoms to overcome catalytic selectivity

The active center of a paired Ir–O–Ir structure was generated by introducing electronic perturbations to correlated single Ir atoms on Co₃O₄ nanosheets, with the oxygen atoms of Ir-O-Ir serving as the primary active site for the selective electrocatalysis of As(III).

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Modulating paired Ir–O–Ir via electronic perturbations of correlated Ir single atoms to overcome catalytic selectivity†

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Single-atom catalysts have been extensively utilized for electrocatalysis, in which electronic metal–support interactions are typically employed to stabilize single atoms. However, this neglects the metal–metal interactions of adjacent atoms, which are essential for the fine-tuning of selective sites. Herein, the high-loading of Ir single atoms (Ir SAs) (8.9 wt%) were adjacent to oxygen vacancy-rich Co3O4 nanosheets (Ir SAs/Co3O4). Electronic perturbations for both Ir single atoms and Co3O4 supports were observed under electronic metal–support and metal–metal interactions, thus generating Ir–O–Co/Ir units. Electrons were transferred from Co and Ir to O atoms, inducing the depletion of 3d/5d states in Co/Ir and the occupation of 2p states in O atoms to stabilize the Ir SAs. Moreover, the O atoms of Ir–O–Ir functioned as the main active sites for the electrocatalysis of As(III), which reduced the energy barrier for the rate-determining step. This was due to the stronger electronic affinities for intermediates from reduction of As(III), which were completely distinct from other coordinated O atoms of Co3O4 or IrO2. Consequently, the resultant Ir SAs/Co3O4 exhibited far more robust electrocatalytic activities than IrO2/Co3O4 and Co/Ir in the electrocatalysis of As(III). Moreover, there was a strong orbital coupling effect between the coordinated O atoms of Ir SAs and the –OH of H2AsO3, thus exhibiting superior selectivity for As(III) in contrast to other common heavy metal cations. This work offers useful insights into the rational design of intriguing SACs with high selectivity and stability for the electrocatalysis and electrochemical analysis of pollutants on an electronic level.

Introduction

Single-atom catalysts (SACs) have been broadly employed in the synthesis of interfaces for electrochemical catalysis, where various support matrices with unsaturated sites and high surface areas were used to maximize the utilization of atoms to obtain highly active catalysts.1–5 However, SACs still suffer from inadequate long-term stability and selectivity for interfacial catalysis-reactions, which are important for their practical applications.6,7 Numerous attempts have recently been made toward the design and optimization of the electrocatalytic performance of active SACs, whose stability can also be enhanced through the introduction of suitable supports including metal compounds (e.g., oxides,18 chalcogenides,19 nitrides,11,12 carbides,13,14 and phosphides25,26) and carbon-based supports.17,18 Researchers have focused on the selectivity of SACs for multistep catalytic reactions (e.g., chlor-alkali process,19 the dehydrogenation of ethanol reaction,20 and the preferential oxidation of CO in hydrogen-rich fuel gas streams;21 etc.), while the development of practical catalysts for the eradication of complex environmental contaminants (e.g., forever chemicals) remains elusive.22 For instance, many types of heavy metal ions (HMIs) coexist in wastewater generated by industrial or mining activities, including Cu(n), Cd(n), Pb(n), Hg(n), and As(m).23,24 Although SACs with high activity participate well in the catalysis and reduction of HMIs, it remains a challenge to accurately analyze HMIs in ambient waterways due to the poor selectivity of SACs. Thus, the design of selectively stabilized catalysts for the modification of electrodes is extremely significant for the practical application of electrochemical strategies.25
The electronic metal–support interactions (EMSIs) provide an efficacious approach to tune the electronic properties of catalysts, thus achieving exceptional catalytic effects.\textsuperscript{26,27} Inspired by the electronic perturbations of Pt nanoparticles interfaced with CeO\textsubscript{2}, Campbell advanced research into strong metal–support interactions (SMSIs) to the scale of electrons.\textsuperscript{28–30} Specifically, there are interfacial charge transfers driven by different chemical potentials between metals and supports, which lead to significantly perturbed electronic structures, thus creating specifically active sites to boost catalytical reactions.\textsuperscript{31} Furthermore, charge transfers at the interface are dependent on the distribution of metals on the supports, as well as the properties of the supports themselves, which strongly affect the intrinsic activities and selectivity of the catalysts.\textsuperscript{32} Thus, the selection of metals and supports is critical.\textsuperscript{33} Single atoms with uniform and explicit active sites provide an ideal platform for the study of EMSIs, which greatly simplifies the exploration of charge transfer within electronic structures.\textsuperscript{28} In terms of supports, metal oxides are more suitable candidates than other nanomaterials (e.g., carbon-based materials), since the uncoupled d states of single metal atoms tend to interact with O atoms.\textsuperscript{27} As one of the typical transition metal oxides, Co\textsubscript{3}O\textsubscript{4} nanomaterials contain highly oxidized redox couples, which have large specific surface areas and are good for adsorption.\textsuperscript{24} It has been reported in the literature that the incorporation of metal atoms into a Co\textsubscript{3}O\textsubscript{4} lattice is an effective means by which the catalytic performance of Co\textsubscript{3}O\textsubscript{4} can be significantly improved. Lou \textit{et al.} incorporated many transition metal atoms (e.g., Mn, Fe, Co, Ni, Cu, and Zn) into a Co\textsubscript{3}O\textsubscript{4} host lattice to obtain a series of metal doped Co\textsubscript{3}O\textsubscript{4} nanomaterials (M–Co\textsubscript{3}O\textsubscript{4}), which showed excellent catalytic performance for electrocatalytic oxygen evolution.\textsuperscript{28} Moreover, isolated noble metal atoms (e.g., Au, Ru, Pt, and Ir) were commendable selections for various catalytic reactions due to their advantages of unique electronic structures and irreplaceable activity. For example, a single Au atom on a Co\textsubscript{3}O\textsubscript{4} (Au/Co\textsubscript{3}O\textsubscript{4}) catalyst exhibited excellent catalytic activity and stability for the oxidation of CO, the active center of which was found to be a dynamic domain centered on the single Au atom that also encompassed the surrounding Co and O atoms (and/or O vacancies).\textsuperscript{34} Wang \textit{et al.} showed that Ru was incorporated into Co\textsubscript{3}O\textsubscript{4} to occupy the surface octahedral sites of Co\textsubscript{3}O\textsubscript{4} to form Ru–O–Co structures, which significantly increased the surface O and promoted the reducibility of Co\textsubscript{3}O\textsubscript{4} catalysts for the oxidation of chlorinated aromatics.\textsuperscript{24} Apparently, these studies still focused on the high activity and stability of catalysts. However, these strong metal–oxygen interactions may reduce the activities of single metals due to few binding sites for reactants.\textsuperscript{1} As for the premise of ensuring the stability and activity of SACs, the regulation of selective active sites still presents immense challenges. Recently, researchers discovered that there were metal–metal interactions between adjacent metal single atoms at a shortened distance, which led to the modified electronic structures of SACs.\textsuperscript{35–38} Unfortunately, since there have been few studies that focused on metal–metal interactions, due to the difficulty in stably anchoring correlated SACs on metal oxides under high loadings, it has always been difficult to distinguish the electronic perturbations of EMSIs and metal–metal interactions. We believe that the stability, activity, and selectivity of SACs can be improved through synergies between EMSIs and metal–metal interactions. Additional research should also be conducted to elucidate the charge transfer kinetics between metal single-atom sites and their supports. Moreover, for certain complex catalytic reactions, links between the selective adsorption of reaction intermediates and active sites should be established. For example, the catalytic reduction of As(III) requires the removal of three –OH groups, as it exists as a stable H\textsubscript{3}AsO\textsubscript{3} molecule in aqueous media, which requires higher catalytic activities than other HMIs.\textsuperscript{37,38}

The design of SACs based on EMSIs and metal–metal interactions to achieve the selective catalysis of As(III) provides typical modelling that is suitable for the study of the kinetics of selective catalysis. This is expected to establish a connection between the electronic perturbations of selective active sites. Therefore, we elaborately designed highly loaded Ir single atoms (Ir SAs) on oxygen vacancy-rich Co\textsubscript{3}O\textsubscript{4} nanosheets (Ir SAs/Co\textsubscript{3}O\textsubscript{4}), where the content of Ir SAs is up to 8.9 wt%. Experimental characterization, involving X-ray absorption fine structure (XAFS) spectroscopy and X-ray photoelectron spectroscopy (XPS), combined with density functional theory (DFT) calculations, revealed that there was an interfacial charge transfer due to the EMSI and metal–metal interactions. This generated electronic perturbations for Ir SAs, Ir–O–Co, and Ir–O–Ir units, where electrons were transferred from Co and Ir sites to O atoms, particularly for Ir–O–Ir structures. This resulted in greatly stabilized Ir SAs with high valence states and active Ir–O–Ir structures. These electron-enriched O atoms were verified to be the essential active sites for electrocatalysis and were completely distinct from the other coordinated O atoms of Co\textsubscript{3}O\textsubscript{4} or IrO\textsubscript{2}. Consequently, the resultant Ir SAs/Co\textsubscript{3}O\textsubscript{4} nanosheets exhibited dozens of times better electrocatalytic activity than IrO\textsubscript{2}/Co\textsubscript{3}O\textsubscript{4} and Co\textsubscript{3}O\textsubscript{4} in the electrochemical analysis of As(III). Moreover, there was a strong orbital coupling effect between the coordinated O atoms of Ir SAs and H\textsubscript{3}AsO\textsubscript{3}; thus, Ir SAs/Co\textsubscript{3}O\textsubscript{4} exhibited superior selectivity to As(III) compared to other common divalent heavy metal cations.

Results and discussion

Morphological and structural characterization of catalysts

Co\textsubscript{3}O\textsubscript{4}, IrO\textsubscript{2}/Co\textsubscript{3}O\textsubscript{4}, and Ir SAs/Co\textsubscript{3}O\textsubscript{4} were prepared and characterized, respectively. The HAADF-STEM image in Fig. 1a excludes the possibility of the aggregation of Ir single atoms with brighter contrast than Co\textsubscript{3}O\textsubscript{4}, and the lattice fringes of 0.285 and 0.243 nm belonged to the (220) and (311) planes of Co\textsubscript{3}O\textsubscript{4}, respectively.\textsuperscript{24}–39 Moreover, in the high-resolution HAADF-STEM (HADDF-HRSTEM) image (Fig. 1b) bright dots (Ir atoms, marked by red circles) occupied the Co\textsubscript{3}O\textsubscript{4} lattice, where the majority of Ir atoms existed adjacent within the cationic Co\textsubscript{3}O\textsubscript{4} lattice. The line intensity profile along the middle of the green dashed rectangles in the inset further indicated the replacement of Co sites with adjacent Ir SAs. From the scanning electron microscopy (SEM) images and transmission electron microscopy (TEM) images (Fig. S1†), similar
Co₃O₄ and Ir SAs/Co₃O₄ nanosheets were confirmed, while the surface of IrO₂/Co₃O₄ was rough due to the generation of numerous nanoparticles, which were assigned to IrO₂ according to a lattice spacing of 0.201 nm. The Ir SAs/Co₃O₄ nanosheets were devoid of nanoparticles and the corresponding elemental mapping images (Fig. S2a†) revealed the uniform distribution of Ir, O, and Co across the nanosheets. Additionally, the energy-dispersive spectrometry (EDS) results in Fig. S2b† confirmed the coexistence of Ir, O, and Co, which was consistent with the elemental mapping images. The mass ratio of Ir was calculated
to be 8.9 wt%, which aligned well with the results obtained by inductively coupled plasma mass spectrometry (ICP-MS). Moreover, in X-ray diffraction (XRD) patterns (Fig. S2†), the Ir SAs/Co3O4 peaks corresponded well with those of Co3O4 (JCPDS#276-1802),41 which further verified the independent dispersion of Ir atoms. The diffraction peaks were sharper and more substantial than those of Co3O4, which implied that the introduction of Ir atoms improved the crystallinity of Co3O4. Another peak near 40° in IrO2/Co3O4 spectra was assigned to IrO2 (JCPDS#15-0870).42 Electron paramagnetic resonance (EPR) was applied to characterize vacancies in these nanomaterials, and isotropic EPR signals with a g value of 2.033 are presented in Fig. S2d,† owing to the presence of positively charged V0.37 Moreover, for IrO2/Co3O4, the signal intensity of V0 slowly decreased with the addition of an Ir source, which meant that the concentration of V0 decreased due to the introduction of IrO2 nanoparticles, while the V0 content of Co3O4 and Ir SAs/Co3O4 was similar. The change in the Raman spectra (Fig. S2e†) further evidenced the speculation that Ir atoms occupied the Co sites. Notably, the A1g band of the Ir SAs/Co3O4 spectra shifted in contrast to that of the Co3O4 spectra, which indicated that the substitution induced lattice expansion.43 Moreover, as the A1g band depends on octahedral cations, it was demonstrated that Ir dopants occupied the Co(m) sites of octahedral units (Co6oct), which was consistent with the tendency that cations with larger radii are preferentially located at octahedral sites.44-46 XPS was employed to examine the surface chemical states of the prepared samples. Fig. S2f-†i present the XPS survey and HR-XPS spectra of the corresponding elements. In the high-resolution XPS spectra of O 1s, the characteristic peaks near 529.8 and 531.6 eV were assigned to Vo and lattice oxygen (O0) for Ir SAs/Co3O4, and the peak at 532.6 eV was due to hydroxyl groups (–OH) or surface-adsorbed water (Oads).47 Compared with Co3O4 and IrO2/Co3O4, the binding energy of O2 in Ir SAs/Co3O4 shifted negatively to 0.4 eV, while the Vo shifted to a high binding energy of 0.3 eV, which elucidated the strong interaction between Ir and O. In addition, the Vo content was reduced in Co3O4 and IrO2/Co3O4, which was consistent with the results revealed by the EPR tests. It could be found that the two fitted peaks of Co 2p3/2 correspond to Co2+ and Co3+, and the Co2+/Co3+ ratio of Ir SAs/Co3O4, Co3O4 and IrO2/Co3O4 was 2.0, 2.0 and 1.8, indicating that the valence of Co in IrO2/Co3O4 was reduced. Moreover, due to the overlap of Co 3p and Ir 4f, we used Ir 4d for comparison, and there was almost no change in the valence state of Ir. BET measurements were used to characterize the surface areas and the porous structures of the prepared samples. The N2 adsorption–desorption isotherms and the suitable aperture distribution of Ir SAs/Co3O4, Co3O4, and IrO2/Co3O4 are presented in Fig. S3.† Their surface areas were calculated to be 122.2, 109.0 and 42.5 m2 g−1, and most of the mesoporous structures were approximately 10.2, 11.6 and 8.8 nm, respectively.

X-ray absorption fine structure (XAFS) spectroscopy was further employed to characterize the electronic structures and coordination attributes of Co and Ir in the prepared samples. First, the valence states of Co and Ir were determined using X-ray absorption near edge structure (XANES) spectra. As presented in the cyan regions of Fig. 1d, the absorption edge of the Co K-edge in Co3O4 was located in the middle of Ir SAs/Co3O4 and IrO2/Co3O4 spectra, revealing that the order of the valence state of Co was Ir SAs/Co3O4 > Co3O4 > IrO2/Co3O4.37 Therefore, compared with Co3O4, the Co of Ir SAs/Co3O4 lost electrons, while the Co of IrO2/Co3O4 gained electrons. As shown in Fig. 1e, the height of the white line peak of the Ir L3-edge in Ir SAs/Co3O4 was higher than that of IrO2, which indicated that the valence state of Ir SAs exceeded +4. This implied a larger population of unoccupied 5d states of Ir atoms than that of supported single atoms.47-48 However, the height of the white line peak in IrO2/Co3O4 spectra was situated in the middle of the Ir foil and IrO2 peaks, suggesting that the valence state of Ir in IrO2/Co3O4 was between 0 and +4. To be noticed, the higher surface valence state obtained in XPS data compared with the XAFS data may be caused by the surface oxidation. The extended X-ray absorption fine structure (EXAFS) spectra were also fitted for further analysis. Fig. 1f depicts the EXAFS spectra of the Ir L3-edge (uncorrected for phase shift), with the detailed fitting results presented in Fig. S4 and Table S1.† For Ir SAs/Co3O4, the peak near 1.58 Å corresponded to the Ir–O scattering path, while the peak near 2.49 Å was different from the Ir–Ir scattering path of the Ir foil peak located at ~2.58 Å, which suggested the absence of Ir NPs/clusters, in alignment with the STEM observation.27 In detail, the first shell was fitted as Ir–O scattering (RIr–O = 2.00 Å), surrounded by 6.0 O atoms, indicating that the Ir SAs attained saturated coordination. The scattering of the second shell was fitted as Ir–O–Ir (RIr–O–Ir = 2.98 Å) and Ir–O–Co (RIr–O–Co = 3.00 Å), with coordination numbers of 4.0 and 3.3, which reflected the strong interaction between single atoms and the Co3O4 substrate. For IrO2/Co3O4, the scattering of the first shell at 1.99 Å corresponded to the Ir–O path with unsaturated coordination with a coordination number of 5.1. The second shell as Ir–Co scattering (RIRr–Co = 2.57 Å) with a coordination number of 1.0, which implied a strong IrO2 interaction with the Co3O4 substrate. The above analysis was conducive to the construction configurations for analysis of changes in electronic structures. In addition, the Co coordination environment was analyzed. The specific EXAFS spectra of the Co K-edge are shown in Fig. S5 and Table S2,† where peaks at ~1.53 Å corresponded to the Co–O scattering paths, whereas the peaks at ~2.46 and 3.02 Å were the Co–Co scattering paths.32 Compared with Co3O4, the coordination number of the Co–Co scattering at RCo–Co = 3.38 Å increased following the incorporation of Ir SAs, with a bond length of 3.35 Å. After compounding IrO2, the Co–Co bond length was 3.36 Å, which indicated that the electronic structure of Co3O4 was adjusted.

Combined with the coordination environment analyzed by XAFS, DFT calculations were used to further explore the differences in the electronic structures of Ir SAs/Co3O4, IrO2/Co3O4, and Co3O4. Fig. S6† shows the front, side, and top views of the optimized configurations. To better distinguish the surface-active sites of IrO2/Co3O4, two configurations were optimized with IrO2 and Co3O4 as the surfaces, respectively, denoted as IrO2/Co3O4(Co) and IrO2/Co3O4(Ir). First, we specifically analyzed the electron transfer process in Ir SAs/Co3O4 (Fig. 1c). As shown in Fig. 1g, the Co and Ir in Ir SAs/Co3O4 were
surrounded by blue regions, which suggested that the Co and Ir atoms lost electrons. In contrast, the O atoms were surrounded by red regions, indicating that they gained electrons. The electron transfer process aligned well with the increasing valence states of Ir and Co revealed by the XANES data. It was worth noting that the surrounding O atoms in Ir–O–Ir was darker in red than the O atoms in the Co₃O₄ bulk structure, meaning that the coordinated O atoms of Ir–O–Ir acquired more electrons and may have been more active. Furthermore, Bader charge analysis was utilized to quantify the Co and Ir atoms, with the results listed in Table S3.† In general, the charge of the Co atoms in Ir SAs/Co₃O₄ shifted from 1.34e in CoO₂ to 1.35e, indicating that the Co atoms acquired 0.01e. Conversely, the charge of Co in IrO₂/CoO₂(Co) was 1.30e, which lost 0.04e compared with Co₃O₄. Furthermore, the charge of Ir atoms in Ir SAs/Co₃O₄ increased by 0.37e compared with that of IrO₂, while the charge of Ir atoms in IrO₂/CoO₂[Ir] decreased by 0.02e. These data matched well with the changes in valence states and electron transfer processes revealed by XANES and differential charge density images. In conclusion, with the integration of Ir SAs, electrons were transferred from Co and Ir to the surrounding O atoms, which induced the depletion of the 3d/5d states of Co/Ir and occupation of the 2p states of O atoms, thus stabilizing the Ir SAs, with O atoms being extremely electron-enriched. In contrast, the Ir and Co atoms gained electrons following IrO₂ compounding, resulting in reduced valence states. Subsequently, the density of states (DOS) of Ir SAs/CoO₂, IrO₂/CoO₂, and Co₃O₄ are compared in Fig. 1h. There were stable bonding orbitals of Ir and coordinated O atoms in Ir SAs/Co₃O₄, which ranged from −10 to −2.5 eV. In the dashed box, the coordinated O atoms of the Ir SAs also had an obvious coupling effect with H₂AsO₃, implying that Ir SAs/CoO₂ had the capacity to interact with H₂AsO₃. Furthermore, the occupied state of the coordinated O of Ir SAs passed through the Fermi level, which suggested excellent electrical conductivity that is an essential prerequisite for catalytic reactions.¶ Similarly, as shown in Fig. S7,† the O of Co₃O₄ on the surface of IrO₂/CoO₂(Co) also passed through the Fermi level, which participated well in the adsorption process. However, the Ir and O bonds in IrO₂/CoO₂[Ir] were well coupled between −10 and −2.5 eV, without an occupied state near the Fermi level to accept electrons for adsorption in the O-p orbital, which meant that the Ir–O bond was stable. By integrating the electrons of the Co-d orbitals, it was found that the d-band centers of CoO₂ and IrO₂/CoO₂(Co) were located at −3.74 eV and −3.36 eV, respectively, while the d-band center of Ir SAs/CoO₂ obviously shifted to −2.73 eV, which signified that the incorporation of Ir SAs modulated the electronic structure of CoO₂ and the position of its d-band center. Undoubtedly, the distinct electronic structures of these materials can give rise to potential differences in the active sites and interaction modes of catalytic reactions.

Effective electrocatalysis and selective detection of As(III)

Ir SAs/Co₃O₄/GCE, Co₃O₄/GCE, and IrO₂/Co₃O₄/GCE were synthesized and used for electrochemical studies. The conductivity of these electrodes was examined via cyclic voltammetry (CV) and electrochemical impedance spectroscopy (EIS), with the results in Fig. S8† showing that the incoming Ir SAs successfully reduced the resistance of metal oxides, which was of significant benefit for electrochemical detection. Simultaneously, linear sweep voltammetry (LSV) data were used to calculate the turnover frequencies (TOFs) to further reveal the differences in the catalytic activities of these modified electrodes. Their LSV signals toward As(III) of 10 ppb are presented in the inset of Fig. 2a, and the spikes that emerged at about 0 V were the stripping peaks of As(0), which shows that the current on Ir SAs/Co₃O₄/GCE was far higher than that for Co₃O₄/GCE and IrO₂/Co₃O₄/GCE. Furthermore, through the shape and area of the LSV peaks, the reaction time (t) and the number of As(III) that contributed to the electrocatalytic reduction reaction (N) were obtained. The TOF was calculated using the equation TOF = N/([A × t]), where A is the active area of electrode interfaces (Fig. S9†).¶ Details regarding the TOF calculation are shown in the ESI.† In short, Ir SAs/Co₃O₄/GCE exhibited the highest activity with an efficiency of 4.47 nm⁻² s⁻¹, which was almost 20-fold and 10-fold better than Co₃O₄/GCE (0.45 nm⁻² s⁻¹) and IrO₂/Co₃O₄/GCE (0.27 nm⁻² s⁻¹), respectively, which demonstrated the prominent catalytic activity of Ir SAs/Co₃O₄/GCE toward 10 ppb As(III) (inset) on Ir SAs/Co₃O₄, IrO₂/Co₃O₄, and Co₃O₄ modified electrodes under the same conditions.
for the electroanalysis of As(III). Additionally, the electroanalytical performance of Co₃O₄/GCE and IrO₂/Co₃O₄/GCE was also confirmed under the same detection conditions using the square wave anodic stripping voltammetry (SWASV) technique with optimized conditions in Fig. S10. Moreover, the sensitivity and LOD comparisons of different modified electrodes are summarized in Fig. S11, with detailed SWASV responses and linear equations. Specifically, the sensitivities toward the As(III) of Co₃O₄/GCE and IrO₂/Co₃O₄/GCE were 0.08 and 0.04 μA ppb⁻¹, respectively, and for the LOD were 4.07 and 4.37 ppb, respectively. Moreover, the electrochemical performance obtained by Ir SAs/Co₃O₄ for the detection of As(III) was superior to that obtained by other modified cobalt-based nanomaterials (Table S4†). Considering the coexistence of other heavy metal ions, Ir SAs/Co₃O₄/GCE was implemented for the electroanalysis of Zn(II), Cd(II), Cu(II), Hg(II), and Pb(II). Fig. S12† presents the detailed SWASV responses and corresponding linear equations, and comparisons of the sensitivity and LOD for the HMs are summarized in Fig. 2b. Explicitly, the sensitivities toward Zn(II), Cd(II), Cu(II), Hg(II), and Pb(II) were 0.001, 0.22, 0.22, 0.07, and 0.23 μA ppb⁻¹, respectively, while the LOD were 360.00, 4.82, 4.32, 3.66, and 4.83 ppb, respectively. The electroanalytical performance of Ir SAs/Co₃O₄/GCE toward these heavy metal cations was far worse than that toward As(III), which verified that Ir SAs/Co₃O₄/GCE was highly selective for As(III). Further in-depth analysis was required to uncover the fundamental mechanism of why the Ir SAs/Co₃O₄/GCE exhibited excellent sensitivity and selectivity for As(III). The stability and reproducibility of the modified electrodes were significant indicators to evaluate their practicability which were further investigated by SWASV toward As(III) at 4 ppb, with the results shown in Fig. S13.† During 10 cycles of SWASV tests of the same electrodes, the relative standard deviation (RSD) was just 1.9% (Fig. S13a†). In addition, the RSD of 10 different Ir SAs/Co₃O₄ modified electrodes was only 4.0% (Fig. S13b†). Except that these RSDs of the currents were less than 5%, there was almost no change in the shape of the peak currents obtained in the above experiments, indicating the excellent stability and reproducibility of Ir SAs/Co₃O₄/GCE. Moreover, the morphologies and structures of Ir SAs/Co₃O₄ after electrochemical tests of 20 times were characterized and analyzed to further certify their durability. First, XRD patterns in Fig. S14† presented unchanged peaks of used Ir SAs/Co₃O₄ compared with fresh catalysts, and there is no peak match with Ir nanoparticles. HADDF-STEM images presented in Fig. S15a and b† show that the Ir atoms used in Ir SAs/Co₃O₄ were evenly distributed as single atoms. Besides, k²-weighted and fitting spectra for Ir L₃-edge and Co K-edge in Fig. S15c-ff further indicated an unvaried coordination environment of Ir SAs/Co₃O₄. All in all, Ir SAs/Co₃O₄ exhibited satisfactory stability, reproducibility and durability.

Charge transfer process in electrocatalysis of As(III)

To reveal the strongly reactive catalytic sites and the analysis of As(III), XPS and XAFS were used to characterize the interactions between Ir SAs/Co₃O₄ and IrO₂/Co₃O₄ and As(III). After adsorbing As(III) at 10 ppm in 0.1 M HAc-NaAc (pH = 5.0), Ir SAs/Co₃O₄ and IrO₂/Co₃O₄ were denoted as Ir SAs/Co₃O₄/As and IrO₂/Co₃O₄/As, respectively. In the XANES spectra of the Co-K edge (Fig. 3a), there was almost no change in the position of the absorption edge of Ir SAs/Co₃O₄ following the interaction with As(III). Furthermore, the unchanged valence of Co indicated that the Co sites might not be related to the active As(III) site. In contrast, the Co K-edge absorption edge of IrO₂/Co₃O₄ shifted toward higher energy levels after interacting with As(III), which meant that the Co lost electrons. As shown in the Ir-L₃ spectra (Fig. 3b), the height of the white line peaks in the Ir SAs/Co₃O₄/As and IrO₂/Co₃O₄/As spectra were higher and lower, respectively, than those prior to adsorption, which implied that the valence state of the Ir SAs increased. In contrast, the valence state of IrO₂ decreased. In the XANES spectra of the As K-edge (Fig. 3c), the position of the absorption edge revealed that the valence state of the adsorbed As was equivalent to that of NaAsO₂, indicating that no noticeable oxidation occurred. From the analysis of the XANES spectra, it was found that the active sites of Ir SAs/Co₃O₄ and IrO₂/Co₃O₄ that interacted with As(III) were quite different. In general, the Ir atoms in Ir SAs/Co₃O₄ lost electrons when interacting with As(III), while the Co and Ir atoms in IrO₂/Co₃O₄ lost and gained electrons, respectively. EXAFS spectra were fitted to analyze the coordination environments of Co and Ir after adsorbing As(III). According to the specifically fitted results of the Co-K edge (Fig. S16 and Table S1†), there was no significant change in the coordination environment of the Co of Ir SAs/Co₃O₄ and IrO₂/Co₃O₄ prior to and following adsorption. Furthermore, the coordination environment of the Ir was fitted and analyzed, with the fitted data shown in Fig. S17 and Table S2.† There was no significant change in the coordination environment of Ir in IrO₂/Co₃O₄, while the coordination environment of Ir in Ir SAs/Co₃O₄ was altered. In addition, changes in the O atoms on the surface were further analyzed via using XPS. As displayed in Fig. S18a†, the peak at 531.6 eV of Ir SAs/Co₃O₄ belonged to that of VO, which shifted to a lower energy level by 0.8 eV after interacting with As(III). Similarly, in Fig. S18b†, the VO peak in the IrO₂/Co₃O₄ spectra also shifted to a lower energy level by 0.5 eV following the interaction with As(III), indicating that the VO in both Ir SAs/Co₃O₄ and IrO₂/Co₃O₄ interacted with As(III). Although the surface area of IrO₂/Co₃O₄ was reduced by the formation of IrO₂ nanoparticles, we found that the physical adsorption toward As(III) was higher than that of Ir SAs/Co₃O₄ and Co₃O₄ in the HR-XPS spectra of As 3d (Fig. S19†), which would be further explained by the DFT adsorption configurations. However, the electrochemical sensitivity of IrO₂/Co₃O₄ is lower than that of Ir SAs/Co₃O₄ due to that the complex catalytic reaction of H₃AsO₃ not only depended on the adsorption process.

As shown in Fig. 3d, the Ir-O bond in Ir SAs/Co₃O₄ was longer by 0.3 Å after the adsorption of As(III), which further suggested that the active sites in Ir SAs/Co₃O₄ were related to Ir. Combined with the highly active O atoms of Ir-OIr and the saturated coordination environment of Ir revealed above, it was speculated that the actual reactive sites were the highly active O atoms. For IrO₂/Co₃O₄, although there was no change in the coordination environment of Ir, considering the change of its
valence state, the active sites could be IrO2 or Co3O4. Furthermore, all of the aforementioned materials were involved in catalyzing As(III), and the specific catalytic interactions could be further analyzed with the help of theoretical calculations.

DFT calculations were employed to analyze changes in the electronic structures of catalysts to reveal the actual active sites. First, the adsorption configurations were optimized, including Ir SAs/Co3O4/As, IrO2/Co3O4(Co)/As, and IrO2/Co3O4(Ir)/As, with the specific optimized configurations shown in Fig. S20.† For Ir SAs/Co3O4, the H3AsO3 molecules were spontaneously adsorbed to the concave surface formed by the VO on Ir SAs/Co3O4, and a hydrogen bond was formed between the coordinated O atoms of the Ir–O–Ir structures and the –OH in H3AsO3. For IrO2/Co3O4, two hydrogen bonds interacted with H3AsO3; specifically, the coordinating O atoms of unsaturated Co in Co3O4 interacted with –OH in H3AsO3, which was denoted as IrO2/Co3O4(Co)/As. Moreover, the H3AsO3 molecules could interact with unsaturated Ir sites, which was denoted as the IrO2/Co3O4(Ir)/As configuration.

More intuitively, the active sites that interacted with H3AsO3 and the electron transfer process were explored through differential charge density images and corresponding slice

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Fig. 3  Normalized XANES spectra of Ir SAs/Co3O4 and IrO2/Co3O4 prior to and following interactions with As(III) of (a) Co K-edge, (b) Ir L3-edge and (c) As K-edge. (d) FT-EXAFS k3-weighted χ(k) function spectra (uncorrected for phase shift) for the Ir L3-edge of Ir SAs/Co3O4 and Ir SAs/Co3O4/As. (e) Charge density difference image of Ir SAs/Co3O4/As, the isovalue was 0.005 eÅ⁻³ (green and yellow regions represent the depletion and accumulation of electrons, respectively). (f) Charge density difference image on the slice through Ir and O in Ir SAs/Co3O4 and O in H3AsO3 (blue and red regions represent the depletion and accumulation of electrons, respectively). (g) PDOS of Ir, O, and H3AsO3 in adsorption configuration.
differential charge density images. As shown in Fig. 3e and f, the Ir atoms of Ir–O–Ir in Ir SAs/Co3O4 obviously lost electrons, while O gained electrons and further transferred them to the –OH of H3AsO3 to reduce H2AsO3.

As shown in Fig. S21a and c† in IrO2/Co3O4(Co), the Co lost electrons to its coordinating O atom and further transferred them to the –OH of H3AsO3. As shown in Fig. S21b and d† in IrO2/Co3O4(Ir), As lost electrons and Ir gained electrons. The electron transfer process revealed by the differential charge density images was consistent with the results revealed by XANES. Bader charge analysis further quantitatively revealed the electron transfer process following the adsorption of H2AsO3 (Table S3†). The Ir atoms of Ir–O–Ir in Ir SAs/Co3O4 lost 0.05e, and the Ir in IrO2/Co3O4(Ir) gained 0.05e, while the Co in IrO2/Co3O4(Co) lost 0.11e.

Variations in the DOS of the Ir and O atoms in these configurations are shown in Fig. 3g. Significantly, the H2AsO3 energy levels adsorbed on the nanomaterials were altered compared with those of free H2AsO3, which also had a strong coupling with the orbital energy levels of specific atoms in the catalysts. Specifically, the O-p orbital of Ir SAs/Co3O4 moved to a lower energy level following the adsorption of H2AsO3. In the dashed box, the O-p orbitals at 0 to −1.67 eV, −5.9 to −6.3 eV, and −9.5 eV matched well with the H2AsO3 energy levels, implying that the O atom of Ir–O–Ir was the active site for interacting with H2AsO3. Similarly, the O atom of the Co3O4 in IrO2/Co3O4(Co)/As also generated an energy level that matched the orbital energy level of H2AsO3. However, in IrO2/Co3O4/Ir/As, Ir in the orbital energy level of Ir–d that matched with H2AsO3 and not that of O-p, indicating that the interactive H2AsO3 sites in IrO2 were unsaturated Ir sites. In general, the actual reactive sites for catalyzing H2AsO3 in Ir SAs/Co3O4 were the active O atoms in Ir–O–Ir, where the VO on its surface played a role in promoting the adsorption of H2AsO3. For IrO2/Co3O4, both the Ir atoms of IrO2 and the coordinated O atoms of unsaturated Co in Co3O4 were the reactive sites that catalyzed H2AsO3.

Selective reduction reaction of As(III) and the proposed catalytic mechanism

To further explore the electrocatalytic reduction reaction of As(III) on different sensing interfaces, adsorption configurations of the H2AsO3 intermediate during the electrocatalytic reduction were optimized. The adsorption and stepwise reduction configurations of H2AsO3 on Ir SAs/Co3O4, IrO2/Co3O4(Co), and IrO2/Co3O4(Ir) are shown in Fig. 4a (side view) and Fig. S20† (front and top views), and corresponding relative energies were compared in Fig. 4b. The adsorption energies (E_ads) of H2AsO3 on Ir SAs/Co3O4, IrO2/Co3O4(Co), and IrO2/Co3O4(Ir) were −1.01, −0.35, and −1.23 eV, respectively. The adsorption capacity of Ir SAs/Co3O4 for H2AsO3 was weaker than that of IrO2/Co3O4(Ir), due to the formation of Ir–As bonds in IrO2/Co3O4(Ir)/As and only hydrogen bonds in Ir SAs/Co3O4. Among the configurations of IrO2/Co3O4(Ir)/As and Ir SAs/Co3O4/As, the largest potential barriers in the stepwise reduction reaction appeared in the *H2AsO2 to *HASO process, and the energy required by Ir SAs/Co3O4 was 0.23 eV, which was lower than that of IrO2/Co3O4(Ir) (0.42 eV). Furthermore, in IrO2/Co3O4(Co)/As, although the *H2AsO2 was spontaneously converted to *HASO, the reduction of *HASO to *As had a potential barrier as high as 1.19 eV, which directly diminished the reduction efficiency of As(III) during...
electrochemical enrichment. This may have also been the explanation for the high adsorption energy, albeit poor electrochemical performance of IrO$_2$/Co$_3$O$_4$(Co). It was observed that Ir–O–Ir bonded with $^*$HASO to spontaneously reduce $^*$HASO to $^*$As. Overall, the external energy required for the rate-determining step for the catalysis of As(III) by Ir SAs/Co$_3$O$_4$ was the lowest, and the Ir–O–Ir active center had a stronger electronic affinity to intermediates, which was beneficial for the more rapid deposition of As(0) on its surface, thereby boosting the electrochemical signal. Therefore, we proposed the mechanism of the electrochemical detection process in Fig. 4c. When adding a certain amount of As(III), As(III) diffused rapidly onto the surface of Ir SAs/Co$_3$O$_4$, and was confined in the concave surface formed by Ve on the surface. Subsequently, the coordinated O atoms of Ir–O–Ir interacted with –OH of H$_2$AsO$_3$, and –OH captured the electrons of the Ir-coordinated O, further accelerating the process by which electrons transferred from the Ir to O atoms in Ir–O–Ir, thereby resulting in the reduced valence state of Ir. Additionally, the Ir–O bonds were elongated by the interaction of H$_2$AsO$_3$ and Ir–O–Ir. –OH was gradually removed from H$_2$AsO$_3$ under the reduction of the active Ir–O–Ir structure. Most of the steps in reducing As(0) were thermodynamically spontaneous, and the external energy required for the rate-determining step was low. In addition, different from other common heavy metal cations, the specific interaction with –OH made Ir SAs/V$_2$Co$_3$O$_4$ super selective toward As(III). At the applied potential, As(III) was first reduced to As(0) and then oxidized to As(III), and the SWASV signal increased linearly with the increased concentration of As(III), thus realizing the reduction and quantitative detection of As(III) with high-sensitivity.

Conclusions

In summary, high-density Ir SAs were stably anchored on oxygen vacancy-rich Co$_3$O$_4$ that was characterized using HADDF, XAFS, and other techniques to elucidate the uniform and adjacent distribution of Ir SAs and the generation of Ir–O–Ir/Co structures. Combined with DFT calculations, there was an interfacial charge transfer due to electronic metal–support and metal–metal interactions, which resulted in evident electronic perturbations for Ir single atoms and Co$_3$O$_4$. Specifically, electrons transferred from Co and Ir sites to O atoms, particularly in Ir–O–Ir structures. These electron-enriched O atoms were totally different from other coordinated O atoms, which served as essential active sites for electrocatalysis with more robust electronic affinities for the intermediates derived from the reduction of As(III). Consequently, they exhibited dozens of times better electrocatalytic activities than IrO$_2$/Co$_3$O$_4$ and Co$_3$O$_4$ in the electrochemical analysis of As(III). Furthermore, new orbital energy levels matching H$_2$AsO$_3$ appeared in the O atoms of Ir–O–Ir; thus, Ir SAs/Co$_3$O$_4$ exhibited superior selectivity for As(III) compared to other common divalent heavy metal cations. This work revealed the electronic perturbations induced by synergetic interactions between the metal–support and metal–metal from electronic structures and charge transfer processes to provide an innovative strategy for the development of highly active, selective, and stable catalysts.

Experimental section

Synthesis of Co$_3$O$_4$, IrO$_2$/Co$_3$O$_4$, and Ir SAs/Co$_3$O$_4$

First, 4.0 mmol Co(NO$_3$)$_2$·6H$_2$O and 5.0 mmol hexadecyl trimethyl ammonium bromide, CTAB were dissolved into 100 mL of water in turn, and 6 mmol sodium borohydride (SB) was quickly added and reacted for 30 min under stirring. Subsequently, 4 mL H$_2$IrCl$_6$·6H$_2$O (40 mg mL$^{-1}$) was injected into the blue-green solution at an even and slow rate of 12 mL h$^{-1}$ under continuous stirring. After reacting for another 4 h, 1.0 mmol SB was quickly added and then reacted for another 20 min to obtain a black solution. The resulting products were collected, rinsed with distilled water and acetone, and dried at 60 °C overnight. Finally, Ir SAs/Co$_3$O$_4$ nanosheets were obtained by calcining the precursor at 450 °C for 4 h under air, at a heating rate of 5 °C min$^{-1}$. For comparison, Co$_3$O$_4$ was synthesized according to the above experimental procedure without the addition of H$_2$IrCl$_6$·6H$_2$O and extra SB. Furthermore, IrO$_2$/Co$_3$O$_4$ was prepared by increasing the H$_2$IrCl$_6$·6H$_2$O content. The specific experimental conditions can be found in the ESI.$\dagger$

Electrochemical adsorption and measurements

All of the adsorption experiments and electrochemical detection were carried out in 0.1 M HAc–NaAc of pH = 5. For SWASV tests, the deposition potential and running time were set at $-1.1$ V and 150 s and the desorption potential and running time were set at $+0.8$ V and 150 s.

Determination of the turnover frequencies

The TOF value for reduction of As(III) on different electrodes was calculated using the equation $\text{TOF} = \frac{N}{A \times t}$, where $N$ is the number of As(III) participating in the electrocatalytic reduction process, which could be obtained by LSV tests. A is the active area of the electrode interface, which could be calculated according to the Randles–Sevcik equation. More specific experiments and calculation processes could be found in the ESI.$\dagger$

Data availability

The data supporting this article and other findings are available from the corresponding authors upon request.

Author contributions

X. H., W. L. and S. C. designed this project. S. C. and Y. Y. implemented the experiments. Z. S. implemented the DFT calculations. X. X., P. L. and M. Y. carried out the data analysis. S. C. and A. C. drafted the manuscript. S. C. and A. C. reviewed and edited the manuscript. All authors approved the final version.

Conflicts of interest

There are no conflicts to declare.
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References