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Detection of high-valent iron species in alloyed oxidic cobaltates for catalysing the oxygen evolution reaction

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Iron alloying of oxidic cobaltate catalysts results in catalytic activity for oxygen evolution on par with Ni-Fe oxides in base but at much higher alloying compositions. Zero-field $^{57}$Fe Mössbauer spectroscopy and X-ray absorption spectroscopy (XAS) are able to clearly identify Fe$^{4+}$ in mixed-metal Co-Fe oxides. The highest Fe$^{4+}$ population is obtained in the 40–60% Fe alloying range, and XAS identifies the ion residing in an octahedral oxide ligand field. The oxygen evolution reaction (OER) activity, as reflected in Tafel analysis of CoFeO$x$ films in 1 M KOH, tracks the absolute concentration of Fe$^{4+}$. The results reported herein suggest an important role for the formation of the Fe$^{4+}$ redox state in activating cobaltate OER catalysts at high iron loadings.
imensional reduction of first row metal oxides gives rise to metallate oxygen evolving catalysts (M-OECs) that exhibit high activity for the oxygen evolution reaction (OER)\(^ {1-5}\). Electrosynthesis of oxidic cobaltates and nickelates in the presence of phosphate and borate (Co\(_{3-x}\)P\(_2\)B\(_2\)O\(_7\)\(^{6,7}\), Co\(_x\)B\(_{3-x}\)O\(_4\)\(^{8,9}\), NiB\(_x\)O\(_4\)\(^{10,11}\), MnP\(_x\)\(^{12,13}\)) results in clusters of 10–60 metal atoms, as determined from in situ pair distribution functional analysis\(^ {3,9,14-19}\). The self-healing property of the M-OECs\(^ {2,20-22}\) allows them to promote water-splitting under benign conditions. Under such conditions, the catalysts may be easily interfaced with materials for direct conversion of water to oxygen and hydrogen at high efficiency\(^ {23-26}\), as well as interfaced with biological organisms to perform artificial photosynthesis\(^ {27}\) at efficiencies greatly exceeding natural photosynthesis\(^ {28,29}\). The metalloclusters possess a high edge-to-area ratio that engenders high activity, as revealed by isotopic labelling studies\(^ {30}\) that show the critical O–O bond formation step to occur by proton-coupled electron transfer (PCET) at cluster edge sites\(^ {31-36}\). Moreover, the electronic charge in M-OECs can delocalize within the clusters\(^ {37,38}\) giving rise to electron/ hole transport\(^ {39}\) that can maximally couple to the ion transport needed to support the OER\(^ {40,41}\).

Iron doping of metal oxide films has long been known to increase overall OER activity of metal oxide OER catalysts\(^ {42,43}\). The behaviour of Fe in Ni-OECs has been revisited\(^ {44}\), and the role of Fe has been ascribed to various factors, including active site Fe\(^ 4+\) or higher valent species\(^ {45-47}\), near neighbour Fe effects on Ni resulting from strain on the oxide lattice\(^ {48-51}\), active oxygen intermediates at Ni–Fe sites\(^ {52-54}\), Fe induced partial-charge transfer to Ni sites\(^ {55,56}\), and Fe acting as a Lewis acid that promotes charge transfer to the ligands and favourable energetics for Ni oxyl formation\(^ {57,58}\). Quizzically, though detected by Mössbauer spectroscopy, the presence of Fe\(^ {4+}\) does not correlate with the observed catalytic activity\(^ {59}\). Iron loading has also been shown to affect the OER activity of Co-OEC\(^ {60-62}\), but at very different alloying loads. Whereas Ni-OECs show maximal activity with Fe loadings of ~5 mol% Fe\(^ {42,55}\), the maximal activity of Co-OECs is observed for Fe loadings of ~40 mol% Fe\(^ {61}\). These higher loadings suggest different roles for Fe in enhancing M-OEC activity at high versus low alloying.

We now report the zero-field\(^ {57}\)Fe Mössbauer and X-ray absorption spectroscopy (XAS) of Co-OEC alloyed with Fe from 0 to 100% and show that, unlike Fe alloyed in Ni-OECs, the presence of Fe\(^ {4+}\) tracks OER activity, suggesting that Fe\(^ {4+}\) is intimately involved as a redox activator of OER. The results suggest different roles for Fe in alloyed M-OEC catalysts. At low loadings such as in (NiFe)-OECs, OER is performed by Ni active sites and Fe promotes the PCET activation of OER. At high loadings, as is observed here for Fe-alloyed Co-OEC catalysts, the redox properties of Fe appear to play a prominent and more direct role in promoting OER.

**Results**

A series of CoFeO\(_x\) films with varying Fe content were prepared by cathodic deposition upon the reduction of nitrate to induce a high local basic pH near the electrode, resulting in the electrodeposition of a CoFe hydroxide film\(^ {31}\), which was then converted to CoFeO\(_x\) with the application of an anodic potential. Metal elemental compositions were determined by inductively coupled mass spectrometry (ICP-MS) of digested films after electrochemical measurements. Previous studies have shown that Fe and Co deposit homogeneously\(^ {31}\) and are detected by SEM/EDS analysis\(^ {63}\). As Fe content increases, both the cathodic and anodic features of the Co\(^ {2+}\)/Fe\(^ {3+}\)/Fe\(^ {4+}\) couple are shifted towards higher potentials (Supplementary Fig. 1). CoFeO\(_x\) films with 40–80 mol% Fe in 1 M KOH (pH 14) that was scrubbed of trace metal contaminants display Tafel slopes of ~30 mV/dec (Supplementary Fig. 2), similar to previously published results\(^ {61}\). We note that the lowest Tafel slopes in CoFeO\(_x\) films are obtained at much higher Fe:Co ratios than observed for NiFeO\(_x\) films.

The electronic structure of Fe centres in CoFeO\(_x\) films was probed with zero-field\(^ {57}\)Fe Mössbauer spectroscopy. A representative\(^ {57}\)Fe Mössbauer spectrum is given in Fig. 1; the spectra of all CoFeO\(_x\) film samples are given in Supplementary Fig. 3 and Supplementary Fig. 4. The spectra are reproducible and sensitive to Fe population changes between samples with 10 mol% Fe differences (Supplementary Fig. 5). Two species of Fe are detected in the\(^ {57}\)Fe Mössbauer spectra. Fits of the spectra furnish corresponding isomer shifts (Δ) and quadrupole splittings (|ΔE\(_Q\)|) for one species with Δ ~ 0.3 mm/s and |ΔE\(_Q\)| ~ 0.7 mm/s and the other species with Δ ~ 0.2 mm/s and |ΔE\(_Q\)| less than the resolved linewidth (~0.3 mm/s). These values, which are somewhat sensitive to total Fe alloying concentrations and fittings (Supplementary Fig. 7), do not correspond to either FeO\(_x\)\(_{3-}\) or metallic Fe\(^ {66,67}\). One species matches the Mössbauer parameters of the high spin (HS) Fe\(^ {3+}\) in the oxide ligand field of NiFeO\(_x\)\(^ {3,68,69}\) or metallic Fe\(^ {66,67}\). The Mössbauer parameters of the second Fe species correspond to those observed previously for Fe\(^ {4+}\) in NiFeO\(_x\)\(^ {70}\), and is consistent with theoretical calculations showing the persistence of Fe\(^ {4+}\) in NiFeO\(_x\)\(^ {71}\). The Fe\(^ {3+}\)/Fe\(^ {4+}\) ratio, which may be determined from the Mössbauer parameters, shifts towards Fe\(^ {3+}\) at low and high Fe concentrations (Supplementary Fig. 6) with a maximal absolute Fe\(^ {4+}\) concentration observed between 40 and 60% Fe loading (Fig. 2). Strikingly, as Fig. 2 illustrates, the population of Fe\(^ {4+}\) in CoFeO\(_x\) films tracks OER activity as reflected in Tafel slopes. We observe a direct correlation between absolute Fe\(^ {4+}\) content and low Tafel slopes (30 mV/dec), implicating the important role of Fe\(^ {4+}\) in enhancing OER activity in CoFeO\(_x\) films at high Fe alloying concentrations. The maximum in activity is likely a result of Fe becoming the dominant compositional metal. Unary Fe oxide films are inferior OER catalysts even as ultrathin sub-monolayer films\(^ {72}\). Thus the observed maximum in activity is consistent with the active site for OER becoming dominated by an Fe-only composition at high iron loadings in excess of 50% loading.

The assignment of Fe\(^ {3+}\) and Fe\(^ {4+}\) as deduced by Mössbauer spectroscopy is supported by Fe K-edge XAS of CoFeO\(_x\) samples with varying Fe content. The X-ray absorption near edge
and Fe\textsuperscript{4+} spectra and corresponding composition 50% Fe:50% Co (black line), and calculated Fe\textsuperscript{3+} slope (red circle) with absolute Fe\textsuperscript{4+} given in Supplementary Fig. 8. As the proportions of Fe\textsuperscript{3+} composition. Analogous data for several other compositions are shown in Fig. 3. Several important observations can be made between the Fe\textsuperscript{3+} and Fe\textsuperscript{4+} species. There is a large edge shift from \textasciitilde7124 to \textasciitilde7128 eV between the Fe\textsuperscript{3+} and Fe\textsuperscript{4+} species. Additionally, the 1s \to 3d pre-edge intensity of the Fe\textsuperscript{3+} species is significantly higher than that for Fe\textsuperscript{4+} (inset of Fig. 3a). The R-space amplitude of the Fe\textsuperscript{3+} species is significantly lower than the Fe\textsuperscript{4+} species (Fig. 3c). This decreased amplitude suggests a lower coordination number. There is also a clear contraction of the first and second shell scattering distances for Fe\textsuperscript{4+}. The pre-edge region reflects transitions to the many-electron excited states of the metal centre. The spectral intensity of the pre-edge derives from both electric quadrupole and electric dipole mechanisms. In a centrosymmetric ligand field (e.g., \(O_h\)), the electric dipole contribution is parity forbidden, and only the quadrupole intensity is present. Conversely, deviation from centrosymmetry (e.g., \(T_d\)) results in a significant increase in the pre-edge intensity. This increase in intensity derives from electric dipole allowedness, which is consistent with the amount of 3d-4p mixing in a non-centrosymmetric ligand field\textsuperscript{73}. These observations, together with a low coordination number from the low amplitude R-space data, suggest that Fe\textsuperscript{3+} is present in a distorted ligand field lacking inversion symmetry—either \(T_d\) or square pyramidal ligand field geometries are likely possibilities\textsuperscript{73}. Along similar lines for Fe\textsuperscript{4+}, the low pre-edge intensity suggests a more symmetric \(O_h\) ligand field, which will largely exhibit electric quadrupole intensity. A more symmetric ligand field is also consistent with the higher amplitude R-space data and small \(\Delta E_C\).

The combination of Mössbauer, XAS, and EXAFS data support the assignment of a high-valent Fe\textsuperscript{4+} species in a symmetric \(O_h\) ligand field. An alternative scenario to consider is a low-symmetry Fe\textsuperscript{3+} centre, though it must be in a strongly electron withdrawing environment. For instance, negative isomer shifts can be obtained in low-symmetry Fe\textsuperscript{3+} complexes in the presence of strong backbonding (e.g., \(K_2[Fe\textsuperscript{3+}(CN)\textsubscript{6}]\) or \(Na_2[Fe\textsuperscript{3+}(CN)\textsubscript{6}(NO)]\))\textsuperscript{74}. Additionally, when \([Fe\textsuperscript{3+}(CN)\textsubscript{6}]\) is coordinated in supramolecular assemblies (e.g., Prussian blue analogues) involving metal–metal interactions. Such second-sphere coordination of the Fe–CN bonds by another metal ion can shift the Fe\textsuperscript{δ} by \textasciitilde0.1 mm/s. Thus, metal–metal or charge transfer interactions in CoFeO\textsubscript{x} could effectively decrease the Fe-based s electron density and give rise to a negative δ and high Fe K-edge energy. However, this scenario would result in an isomer shift that gradually shifts more negative as this Fe species is surrounded by more Co centres. This is not observed here; the isomer shift of the Fe species remains relatively constant at δ \textasciitilde0.2 mm/s and in fact becomes slightly less negative with increasing Co concentration (Supplementary Fig. 7). Similarly, the Fe\textsuperscript{4+} XAS spectra obtained from linear combination fits using different Co:Fe ratios are very similar (Supplementary Fig. 9). These considerations, together with the weak ligand field imposed by oxide coordination and the consistency between the Mössbauer and Fe K-edge XAS data, suggest that the Fe species observed here can be assigned to a high-valent Fe\textsuperscript{4+} centre in an \(O_h\) ligand field.

**Fig. 2 Correlation of Tafel slope with Fe\textsuperscript{4+} composition.** Overlay of Tafel slope (red circle) with absolute Fe\textsuperscript{4+} (green triangle) population in CoFeO\textsubscript{x} films with increasing Fe content. Tafel measurements were run in triplicate, and the average value is shown on the graph at 95% confidence limits.

**Fig. 3 X-ray absorbance spectra of CoFeO\textsubscript{x}.** a Fe K-edge X-ray absorbance spectra and corresponding b k-space and c R-space for CoFeO\textsubscript{x} with the composition 50% Fe:50% Co (black line), and calculated Fe\textsuperscript{3+} (blue line) and Fe\textsuperscript{4+} spectra (green line). Inset of a highlights the pre-edge region.

Discussion

Iron activates M-OECs for OER but its role appears to differ with the nature of the M-OEC and the condition under which it operates. Although most OER is performed in concentrated base, the large-scale deployment of renewable energy storage has prompted interest in performing OER in neutral water sources\textsuperscript{75,76}. For this line of investigation, M-OECs excel owing to their stability arising from their self-healing properties\textsuperscript{22}. At neutral pHs, Fe\textsuperscript{3+} plays a role in OER that appears to be derived
from non-redox properties. PCET activation of water is impaired since water is a poor proton acceptor; Fe may act as a Lewis acid to increase the acidity of OH (aqua/hydroxo) moieties that are coordinated to M-OECs and thereby lower the reduction potential for the M3+/M4+ couple and lead to a greater population of M4+ in the Fe-doped catalysts. This in turn gives rise to increased oxyl character (M(IV)... O ↔ M(III)... O). This Lewis acidity behaviour is supported by the observation that Fe doping in NiPbO4 shows no enhancement in OER at solution pH values commensurate with the pKa of Fe4+. Moreover, OER enhancement may be replicated by non-redox active, Lewis acidic cations in Fe-free Ni-OECs. In concentrated base, OH− can adequately serve the role of a proton acceptor and the influence of the Fe4+ is diminished. When Fe4+ is implicated in OER, as has been proposed in numerous studies, OER appears to occur at the M (Co or Ni) metal centre with Fe4+ promoting the activation of OER at the M of the M-OEC. Such proposals are consistent with the electronic structure of first-row transition metal centres confronting the “oxo-wall”79. Moving to the right in the periodic table, the d-electron count for the M4+ formal oxidation state increases and in a tetragonal oxide ligand field, the d5x2 and dxz orbitals are populated, preventing electron donation from terminal oxygen to the metal centre. Consequently, the M–O bond strength is much weaker for Co4+ than for Fe4+, which formally accommodates a kinetically more inert metal-oxo double bond. From a kinetics perspective, we believe that Fe4+ is not the active site from which OER occurs but rather OER occurs from Co centres with the Fe4+ participating as a redox cooperative centre where Fe4+ enhances the oxidizing power of a Co:Fe active site (Supplementary Fig. 1) versus a Co4+–only active site. Thus, we believe that for both NiFeOx and CoFeOx systems, Fe3+ functions as a Lewis acid in promoting PCET reactivity for the OER. However, unlike NiFeOx OER activity in CoFeOx tracks the Fe4+ alloying concentration, suggesting that the redox properties of the Co4+ centre is further enhanced by the presence of redox active Fe4+ centres.

In conclusion, we have spectroscopically detected and characterized a high-valent Fe4+ centre in CoFeOx thin film OECs. Spectroscopic data suggest that this Fe4+ centre is located in a symmetric O3+ oxide ligand field. The correlation between Fe4+ content and OER activity in CoFeOx thin films suggests an important role of this high-valent state in the mechanism of O–O bond formation and oxygen evolution and supports the merits of exploring mixed-metal oxide oxygen evolution catalysts.

Methods

Materials. Catalysts with specific Fe:Co ratios were prepared by electrodeposition from metal nitrate salt solutions that were degassed. After deposition, the film was rinsed briefly in Type I water and then submerged in KOH buffer. Films were held at a constant potential of 0.84 V in 1 M KOH pH 14 or 1.0 V in 0.1 M KOH pH 13 for 3 h to convert the film to the oxohydroxide form before further electrochemical analysis. To obtain films of various thicknesses, the total deposition time was altered between 30 and 120 s and the current held during deposition was changed between 0.5, 1.0 and 5.0 mA/s. The exact film loading was obtained from ICP-MS analysis of the films.

Electrochemistry. All electrochemical experiments were conducted at room temperature (23 ± 1 °C). Electrode potentials were converted to the NHE scale using E(NHE) = E(Ag/AgCl) + 0.197 V. Overpotentials for the OER from water were computed using η = E(NHE) − (0.23 V − 0.059 V × pH).

Spectroscopy. CoFeOx catalyst with natural 57Fe abundance were prepared for Mössbauer spectroscopy at 77 K. The data were calibrated and fit to linear combinations of symmetric pairs of Lorentzian peaks. Fe K-edge XANES spectra were collected at beamline 12BM-B at the Advanced Photon Source at Argonne National Laboratory. Reconstructed spectra of the pure Fe4+ and Fe3+ species were obtained through linear combinations of the XAS spectra of the various CoFeOx films.

Data availability

Experimental procedures, characterization of compounds electrochemical and spectral data are available in the Supplementary Information. All data are available from the authors on reasonable request.

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Author contributions
N.L., R.G.H., and D.H. conducted the experiments. N.L., R.G.H., D.H., L.X.C., and D.G.N. designed experiments and analysed and interpreted the data. D.G.N. prepared a draft of the manuscript and finalized the manuscript.

Competing interests
The authors declare no competing interests.

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