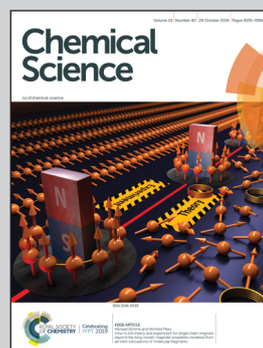


Showcasing research from Professor Kornienko's laboratory, Department of Chemistry, University of Montreal, Quebec, Canada and from Professor Amandine Guiet's laboratory, Institut des Molécules et Matériaux du Mans, France.

Investigation of mixed-metal (oxy)fluorides as a new class of water oxidation electrocatalysts

Energy research is a central focus in the field of chemistry. Within this context, electrocatalyst development and understanding is critical towards realizing practical electricity to fuel technologies. As the oxygen evolution reaction (OER) is a critical bottleneck in many systems, this research focused on developing and understanding a new class of OER-active materials. To this end, we report on the synthesis and characterization of a series of mixed-metal (Ni, Co, Fe) fluorides and oxyfluorides and find them to be exceptionally active OER catalysts, expanding the community's toolset of catalytic materials.

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# Investigation of mixed-metal (oxy)fluorides as a new class of water oxidation electrocatalysts†

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The development of electrocatalysts for the oxygen evolution reaction (OER) is one of the principal challenges in the area of renewable energy research. Within this context, mixed-metal oxides have recently emerged as the highest performing OER catalysts. Their structural and compositional modification to further boost their activity is crucial to the wide-spread use of electrolysis technologies. In this work, we investigated a series of mixed-metal F-containing materials as OER catalysts to probe possible benefits of the high electronegativity of fluoride ions. We found that crystalline hydrated fluorides,  $\text{CoFe}_2\text{F}_8(\text{H}_2\text{O})_2$  and  $\text{NiFe}_2\text{F}_8(\text{H}_2\text{O})_2$ , and amorphous oxyfluorides,  $\text{NiFe}_2\text{F}_{4.4}\text{O}_{1.8}$  and  $\text{CoFe}_2\text{F}_{6.6}\text{O}_{0.7}$ , feature excellent activity (overpotential for 10  $\text{mA cm}^{-2}$  as low as 270 mV) and stability (extended performance for >250 hours with ~40 mV activity loss) for the OER in alkaline electrolyte. Subsequent electroanalytical and spectroscopic characterization hinted that the electronic structure modulation conferred by the fluoride ions aided their reactivity. Finally, the best catalyst of the set,  $\text{NiFe}_2\text{F}_{4.4}\text{O}_{1.8}$ , was applied as anode in an electrolyzer comprised solely of earth-abundant materials, which carried out overall water splitting at 1.65 V at 10  $\text{mA cm}^{-2}$ .

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## 1 Introduction

The rapidly growing consumption of fossil fuels to meet the expanding energy demands of today's society is leading to negative consequences to the environment.<sup>1,2</sup> Global warming, ocean acidification, extreme weather events and low air quality are emerging problems. To mitigate further environmental changes, fossil fuels may be replaced by alternative energy sources. Such sources include wind, hydro, and solar power, and their conversion to electrical power is being developed. However, renewable sources are typically intermittent, presenting an obstacle for their widespread use. As such, the conversion of renewable electricity to energy-dense fuels and value-added chemicals is important to increase the penetration of renewables in the market.

A key technology within this context is the electrolysis of  $\text{H}_2\text{O}$  and/or  $\text{CO}_2$  into  $\text{H}_2$  and C-containing fuels. These reduction reactions are balanced by the oxidation of  $\text{H}_2\text{O}$  into  $\text{O}_2$  and, as such, the continual development of highly active, cost efficient and stable oxygen evolution reaction (OER) catalysts is important to render this technology economically viable. The majority

of efforts in recent years have focused on metal-oxide OER catalysts as alternatives to Ir and Ru oxides that were traditionally used.<sup>3</sup> The highest performing class of these OER catalysts is often iron based oxides/oxyhydroxides containing Ni or Co, and in select cases, these 3d metal oxides outperform the precious metal standards.<sup>4,5</sup> The presence of strain,<sup>6</sup> defects,<sup>7,8</sup> and dopants<sup>9</sup> as well as exfoliation<sup>10</sup> and an amorphous structure<sup>11</sup> have been shown to further boost the performance of metal oxides. We point the interested reader to several recent reviews and perspectives on metal-oxide OER catalysts.<sup>12–17</sup>

The utilization of inductive effects in electrocatalysis is an effective method to modulate materials' performance.<sup>18–20</sup> Substitution with metals having different electronegativity will induce the tendency to donate or withdraw electron density. In the context of OER, a large number of coordinately unsaturated sites on the catalyst with an electron-deficient configuration would boost water oxidation performance.<sup>21–23</sup> Thus, metal fluorides should be promising candidates for high-performance catalysts, given that fluorine is the highest electronegative element and therefore abstract electrons from the neighboring metals. As a consequence, the electronic structure of the transition-metal active sites is modified. However, the poor electronic conductivity of pure metal fluorides  $\text{M}_x\text{F}_y$  hinders their use as highly efficient electrocatalyst. In this context, the use of oxyfluorides  $\text{M}_x\text{O}_y\text{F}_z$  can be a good alternative as they offer a good chemical and thermal stability as well as an enhanced electronic conductivity while preserving key characteristics due to the strong electronegativity of fluorine (3.98 for F

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vs. 3.44 for O).<sup>24</sup> Indeed, oxyfluorides are present in a large range of applications such as ceramic glasses, laser cooling systems, optical amplifiers, and lithium-ion batteries.<sup>25,26</sup> Even if the advantages of the introduction of fluorine element in changing the chemical properties and electronic structures have been demonstrated,<sup>27</sup> only rare reports were found in the literature on fluorides or oxyfluorides as efficient catalyst for water oxidation.<sup>28–33</sup> The lack of studies could be explained by the challenging task to prepare oxyfluorides due to the difficulty to stabilize both fluorine and oxygen anions despite their similar ionic radii (F 1.31 Å, O 1.38 Å). Indeed, the number of iron-based oxyfluoride synthetic methods remains modest in the literature compared to pure oxides and fluorides.<sup>27</sup> As fluoride precursors are frequently sensitive to air humidity and can be easily hydrolyzed, especially at high temperatures, iron oxyfluoride FeOF was first synthesized by solid state reaction in a sealed platinum tube at 950 °C for 24 h from a mixture of Fe<sub>2</sub>O<sub>3</sub> and FeF<sub>3</sub>.<sup>34</sup> FeOF was also tempted to be prepared by solid–gas reaction with the F<sub>2</sub> through the fluorination of Fe<sub>3</sub>O<sub>4</sub> magnetite at 120 °C but only the formation of an oxyfluoride layer was observed at the surface of nanoparticles.<sup>35</sup> In order to avoid the use of pure and sensitive fluoride precursors and toxic F<sub>2</sub>, the synthesis of iron-based oxyfluorides using hydrated fluoride precursors was developed by Zhu *et al.*<sup>36</sup> The authors succeed to obtain FeOF nanorods using FeF<sub>3</sub>·3H<sub>2</sub>O in 1-propanol at 200 °C for 24 h. Other hydrated fluorides precursors such as FeSiF<sub>6</sub>·6H<sub>2</sub>O could also be used to obtain FeO<sub>x</sub>F<sub>2–x</sub> oxyfluorides through their thermal decomposition between 150 to 300 °C.<sup>37</sup> More recently, successive dehydration at 240 °C followed by dehydroxylation at 350 °C of the hydrated iron hydroxyfluoride FeF<sub>2.2</sub>(OH)<sub>0.8</sub>·0.33H<sub>2</sub>O (ref. 38) lead to the successful preparation of a lacunar oxyfluoride with the formulation FeF<sub>2.2</sub>O<sub>0.4</sub>□<sub>0.4</sub>.<sup>39</sup>

Following this strategy, the preparation of new iron-based hydrated fluorides M<sup>2+</sup>Fe<sub>2</sub><sup>3+</sup>F<sub>8</sub>(H<sub>2</sub>O)<sub>2</sub> (M = Mn, Co, Ni, Cu) by microwave heating assisted solvothermal synthesis from metal salts, aqueous hydrofluoric acid and methanol as solvent was previously reported.<sup>40</sup> These hydrated crystallized phases were further calcinated under ambient air to obtain the corresponding amorphous oxyfluorides. Though those resulting amorphous oxyfluorides were tested as cathode active material in Li-ion batteries, neither a thoroughly study of the morphology and electronic structure of those new amorphous iron-based mixed-metal oxyfluorides nor their viability as OER catalysts have been conducted.

In this paper, we set out to synthesize a series of Ni–Fe and Co–Fe (oxy)fluorides in both their crystalline and amorphous structure. Crystalline hydrated fluorides, NiFe<sub>2</sub>F<sub>8</sub>(H<sub>2</sub>O)<sub>2</sub> and CoFe<sub>2</sub>F<sub>8</sub>(H<sub>2</sub>O)<sub>2</sub> and amorphous oxyfluorides NiFe<sub>2</sub>F<sub>4.4</sub>O<sub>1.8</sub> and CoFe<sub>2</sub>F<sub>6.6</sub>O<sub>0.7</sub> materials were synthesized and characterized by XRD diffraction, thermal analysis and electronic microscopies. Their subsequent electrochemical characterization revealed each of these materials to be exceptionally active OER catalysts while Raman and X-ray photoelectron spectroscopies offered mechanistic clues to their superior activity. Finally, the highest performing NiFe<sub>2</sub>F<sub>4.4</sub>O<sub>1.8</sub> was combined with another earth abundant catalyst, cobalt sulfide, in

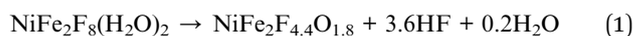
a proof-of-concept overall water electrolysis system based on earth-abundant materials.

## 2 Results and discussion

### 2.1 Synthesis and characterization

The hydrated fluorides, NiFe<sub>2</sub>F<sub>8</sub>(H<sub>2</sub>O)<sub>2</sub> and CoFe<sub>2</sub>F<sub>8</sub>(H<sub>2</sub>O)<sub>2</sub>, were synthesized through a facile microwave heating assisted solvothermal synthesis. In brief, metal chlorides are mixed together with aqueous hydrofluoric acid (HF<sub>40%</sub>) and methanol. The mixture is heated by microwave irradiation at 160 °C for 30 min and leads to green and pink crystalline powders for NiFe<sub>2</sub>F<sub>8</sub>(H<sub>2</sub>O)<sub>2</sub> and CoFe<sub>2</sub>F<sub>8</sub>(H<sub>2</sub>O)<sub>2</sub> respectively. The X-ray powder diffraction patterns, indexed in the monoclinic system with the C2/m space group, show that these metal hydrates are isostructural with Fe<sub>3</sub>F<sub>8</sub>(H<sub>2</sub>O)<sub>2</sub> (Fig. 1a and b and S1†).<sup>41,42</sup> The structures were determined using the Rietveld method (ESI†).

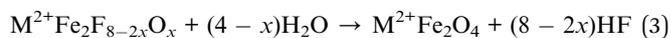
Those resulting crystalline hydrated fluorides were used as precursors to obtain the corresponding amorphous oxyfluorides by an appropriate treatment under ambient air. Their structural and compositional evolutions with the temperature were monitored by thermogravimetric and thermogravimetric analyses (Fig. 1c and d and S2†). The diffraction peaks positions of the hydrated crystallized phases (blue domain) shift above 180 °C to lower 2θ values and their intensities decrease. The last phenomenon is related to the elimination of water and hydrogen fluoride molecules leading to the amorphous phase (green domain) as confirmed by mass spectroscopy coupled thermogravimetric (MS-TGA) analysis under N<sub>2</sub> (Fig. 2). Further calcination and hydrolysis at higher temperature allows the formation of the corresponding crystallized spinel M<sup>2+</sup>Fe<sub>2</sub><sup>3+</sup>O<sub>4</sub> structures. The formulations of the intermediate stabilized amorphous oxyfluorides were determined through the following chemical reactions. For NiFe<sub>2</sub>F<sub>8</sub>(H<sub>2</sub>O)<sub>2</sub>, the experimental weight loss (21.0 wt%) corresponds to NiFe<sub>2</sub>F<sub>4.4</sub>O<sub>1.8</sub> following the reaction (1):



And in the case of CoFe<sub>2</sub>F<sub>8</sub>(H<sub>2</sub>O)<sub>2</sub>, to the reaction (2). It must be noted that CoFeF<sub>5</sub>·7H<sub>2</sub>O as impurity was detected by XRD (Fig. S1†) and quantified (8% molar) by Mössbauer spectrometry.<sup>40</sup> This amount has to be taken into account to obtain the 16.2% experimental weight loss.



Compared to TGA under N<sub>2</sub>, TGA under ambient air shows identical first weight losses (Fig. S2†) but for temperatures above 400 °C, slow weight losses related to a hydrolysis occur that leads to spinel oxides (Table S1†) according to reaction (3):



Consequently, the stabilized amorphous oxyfluorides phases were prepared by thermal decomposition of M<sup>2+</sup>Fe<sub>2</sub><sup>3+</sup>F<sub>8</sub>(H<sub>2</sub>O)<sub>2</sub>



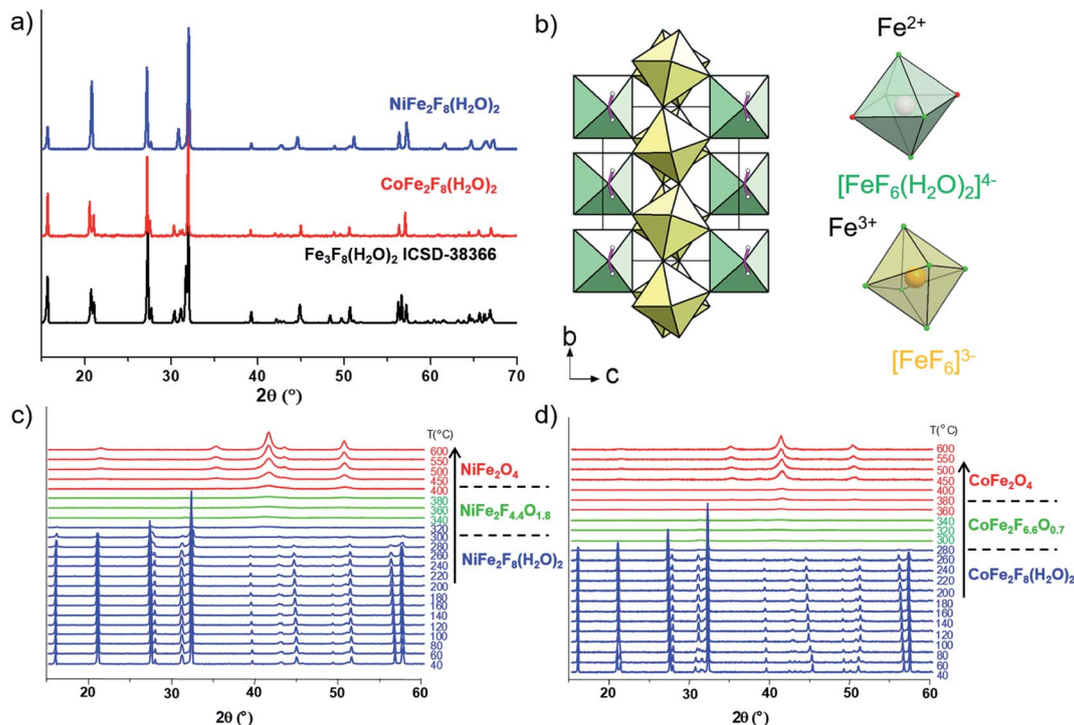


Fig. 1 (a) XRD patterns of  $\text{NiFe}_2\text{F}_8(\text{H}_2\text{O})_2$  and  $\text{CoFe}_2\text{F}_8(\text{H}_2\text{O})_2$  hydrated phases collected at room temperature compared to that of  $\text{Fe}_3\text{F}_8(\text{H}_2\text{O})_2$  (ICSD-38366), (b) projection along a axis of  $\text{MFe}_2\text{F}_8(\text{H}_2\text{O})_2$  structure. Thermal evolution of the X-ray diffractograms under dry air of (c)  $\text{NiFe}_2\text{F}_8(\text{H}_2\text{O})_2$  and (d)  $\text{CoFe}_2\text{F}_8(\text{H}_2\text{O})_2$ .

under air for 1 h at 340 °C and 320 °C for Ni and Co, respectively. Electronic microscopies (SEM and TEM) together with nitrogen sorption have been performed to determine the size and the morphology of the Ni–Fe and Co–Fe based compounds before and after calcination. As revealed by SEM (Fig. S3†), microsized particles are obtained for the hydrate fluorides which is in good accordance with the sharpness of the peaks in the diffractograms (Fig. 1a). The decomposition of those crystalline fluorides leads in both cases to a significant decrease of the particle size (Fig. S3†). In order to probe the atomic distribution in those resulting amorphous oxyfluorides, energy dispersive spectroscopy (EDS) elemental mapping, carried out by SEM, shows that

both metals (Ni/Fe and Co/Fe) were homogeneously dispersed without phase segregation within the resolution capacity of the instrument (Fig. S3†). The final Fe to metal ratio of 2 present in the initial hydrated fluoride precursors was also confirmed. This nanostructuration though the thermal treatment was further investigated by transmission electron microscopy (TEM) and  $\text{N}_2$  sorption. As shown in Fig. 3 for high magnification, fine structures are observed for both materials and in the case of Ni–Fe amorphous oxyfluorides, pores of less than 10 nm could be detected. This emerging porosity is probably related to the precursor's decomposition. Indeed, the HF and  $\text{H}_2\text{O}$  gas molecules liberated during the thermal decomposition could act as

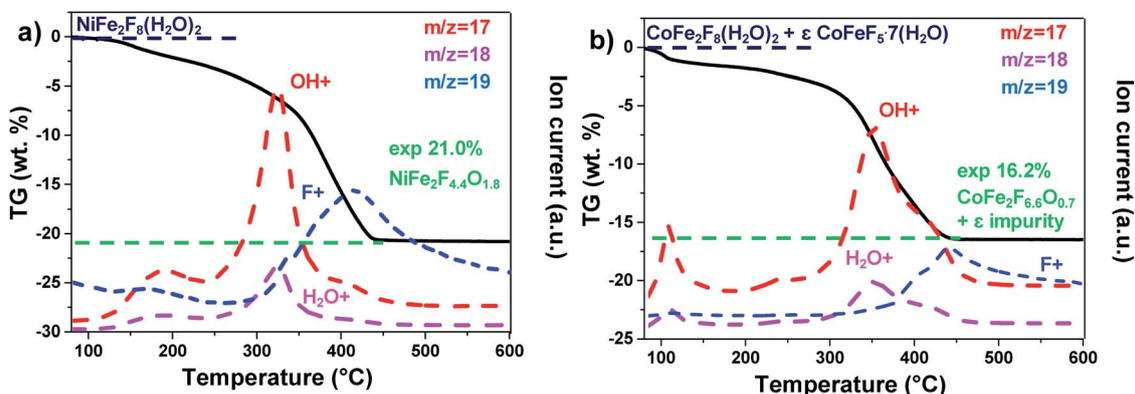


Fig. 2 MS coupled TGA analysis under  $\text{N}_2$  of (a)  $\text{NiFe}_2\text{F}_8(\text{H}_2\text{O})_2$  and (b)  $\text{CoFe}_2\text{F}_8(\text{H}_2\text{O})_2$ .



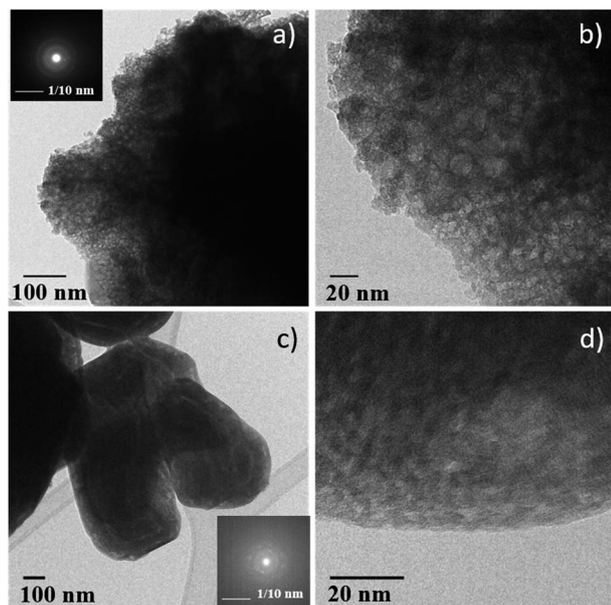


Fig. 3 TEM micrographs of (a) and (b)  $\text{NiFe}_2\text{F}_{4.4}\text{O}_{1.8}$  and (c) and (d)  $\text{CoFe}_2\text{F}_{6.6}\text{O}_{0.7}$ . Insets: corresponding SAED.

a self-generated porogen.<sup>43</sup> The porosity enhancement between  $\text{NiFe}_2\text{F}_{4.4}\text{O}_{1.8}$  and  $\text{CoFe}_2\text{F}_{6.6}\text{O}_{0.7}$  seems to be related to the increase of the number of lost HF molecules. Indeed, as shown in reaction (1) and (2), a release of 3.6HF molecules is determined for  $\text{NiFe}_2\text{F}_8(\text{H}_2\text{O})_2$  whereas only 1.4 for  $\text{CoFe}_2\text{F}_8(\text{H}_2\text{O})_2$ . The amorphous character of those oxyfluorides, evidenced by thermodiffraction, was further confirmed by TEM as diffuse electron diffraction patterns were obtained by selected area electron diffraction (SEAD) on several grains and no distinct diffraction fringes at higher resolution could be observed (Fig. 3). TEM analyses could not be performed on the hydrated fluorides as they were not stable under the electron beam.

$\text{N}_2$  sorption measurements were carried out to determine the specific surface area ( $\text{SA}_{\text{BET}}$ ) of the fluorinated materials before and after thermal treatment. As expected for microsized hydrated fluorides, the measured surface areas are less than  $10 \text{ m}^2 \text{ g}^{-1}$ . However, for the corresponding oxyfluorides obtained after calcination, the  $\text{SA}_{\text{BET}}$  is drastically increased up to  $76 \text{ m}^2 \text{ g}^{-1}$  and  $30 \text{ m}^2 \text{ g}^{-1}$  for  $\text{NiFe}_2\text{F}_{4.4}\text{O}_{1.8}$  and  $\text{CoFe}_2\text{F}_{6.6}\text{O}_{0.7}$  respectively confirming the porogen effect of the  $\text{H}_2\text{O}$  and HF release during the thermal treatment.  $\text{N}_2$  adsorption/desorption isotherms shows type IV hysteresis corresponding to mesoporous structure according to the IUPAC classification.<sup>44</sup> In the case of  $\text{NiFe}_2\text{F}_{4.4}\text{O}_{1.8}$ , the BJH pore-size distribution analysis (Fig. 4a inset) shows an average pore diameter inferior to 10 nm for Ni-Fe phase, value in good agreement with the TEM observation.

X-ray photoelectron spectroscopy (XPS) was subsequently utilized to probe the electronic structure of the transition metal species within the hydrated fluoride and oxyfluoride catalysts (together labelled (oxy)fluorides). In these measurements, both the binding peak position and peak shape provide element-specific information on oxidation state and chemical

environment. The binding energies, and specifically the  $2p_{3/2}$  peaks (denoted with a \* in Fig. 5), of our materials were compared to those found in the national institute of science and technology (NIST) database.<sup>45</sup> The Ni  $2p_{3/2}$  binding energies were determined to be at 858.1 and 855.6 eV for  $\text{NiFe}_2\text{F}_8(\text{H}_2\text{O})_2$  and  $\text{NiFe}_2\text{F}_{4.4}\text{O}_{1.8}$ , respectively (Fig. 5a). The values for  $\text{NiF}_2$  have been measured at 857.4–858.2 eV. In comparison, the Ni  $2p_{3/2}$  peak is typically found at 855–856 eV for  $\text{Ni}(\text{OH})_2$ , and at 854–855 eV for NiO. This indicates that the F withdraws electron density from the Ni in our materials as their binding energies, especially that of  $\text{NiFe}_2\text{F}_8(\text{H}_2\text{O})_2$ , are positively shifted in comparison to Ni(II) oxides/hydroxides. The Co  $2p_{3/2}$  peaks of  $\text{CoFe}_2\text{F}_8(\text{H}_2\text{O})_2$  and  $\text{CoFe}_2\text{F}_{6.6}\text{O}_{0.7}$  were centered at 782.9 eV and 781.2, respectively (Fig. 5b). Likewise, these peaks are shifted slightly higher in binding energy as compared to CoO (~780.4 eV),  $\text{Co}(\text{OH})_2$  (781–782 eV),  $\text{Co}(\text{OH})\text{O}$  (~780 eV) and  $\text{Co}_3\text{O}_4$  (779–780 eV) and closer to those of  $\text{CoF}_2$  (783.0 eV) and  $\text{CoF}_3$  (782.4 eV). The same can be said for the Fe  $2p_{3/2}$  peaks, which were found at 712.9 eV ( $\text{NiFe}_2\text{F}_{4.4}\text{O}_{1.8}$  and  $\text{CoFe}_2\text{F}_{6.6}\text{O}_{0.7}$ ) and 714.4 eV ( $\text{NiFe}_2\text{F}_8(\text{H}_2\text{O})_2$  and  $\text{CoFe}_2\text{F}_8(\text{H}_2\text{O})_2$ ) (Fig. 5c). These shifted the most in comparison to (709–710 eV),  $\text{FeOOH}$  (711–712 eV),  $\text{Fe}_3\text{O}_4$  (709–710 eV) and  $\text{Fe}_2\text{O}_3$  (710–711 eV) and are closer to  $\text{FeF}_3$  (~714 eV). The O 1s spectra for  $\text{NiFe}_2\text{F}_8(\text{H}_2\text{O})_2$  and  $\text{CoFe}_2\text{F}_8(\text{H}_2\text{O})_2$  are similar to that of pure water (~533 eV) while the oxyfluoride O 1s spectra featured only a red-shifted peak at ~530 eV as the O was incorporated in the lattice (Fig. S4†). Similarly, the F 1s spectra displayed a peak at 685.2 eV for  $\text{NiFe}_2\text{F}_8(\text{H}_2\text{O})_2$  and  $\text{CoFe}_2\text{F}_8(\text{H}_2\text{O})_2$  that is slightly red-shifted to 684.8 eV upon their conversion to oxyfluorides.

In summation, the XPS investigation points to all of the transition metals in the new synthesized (oxy)fluorides being electron-poor relative to their oxide analogues, induced by the presence of the highly electronegative F anions. The Fe metal cation likely experiences the largest magnitude of these effects and this is especially pronounced in the hydrated fluorides. Because the exact position of the peaks is not linearly proportional to oxidation state and electronic structure, we do not yet draw quantitative conclusions regarding the magnitude of inductive effects conferred by the F anion.

## 2.2 Electrocatalysis

To evaluate the electrochemical performance of the (oxy)fluoride materials, the as-prepared powders were sonicated together with a Nafion binder and carbon nanotube conductive adhesive to generate a catalyst ink. The catalyst ink was then drop-cast onto a carbon paper electrode and dried prior testing in 1 M KOH. Several (~3–6) electrodes were prepared for each measurement and error bars represent standard deviations from multiple electrodes. Cyclic voltammetry (CV) tests of the catalysts in 1 M KOH electrolyte illustrated that each material undergoes a series of redox-changes prior to OER catalysis, indicating the transformation to a catalytically active state (Fig. 6a). This is common in mixed-metal oxides of Ni, Co, and Fe, in which typically the surface evolves into an oxyhydroxide phase at a potential in the range of where our redox peaks are located.<sup>4</sup> This may indicate that the surface of our (oxy)fluorides



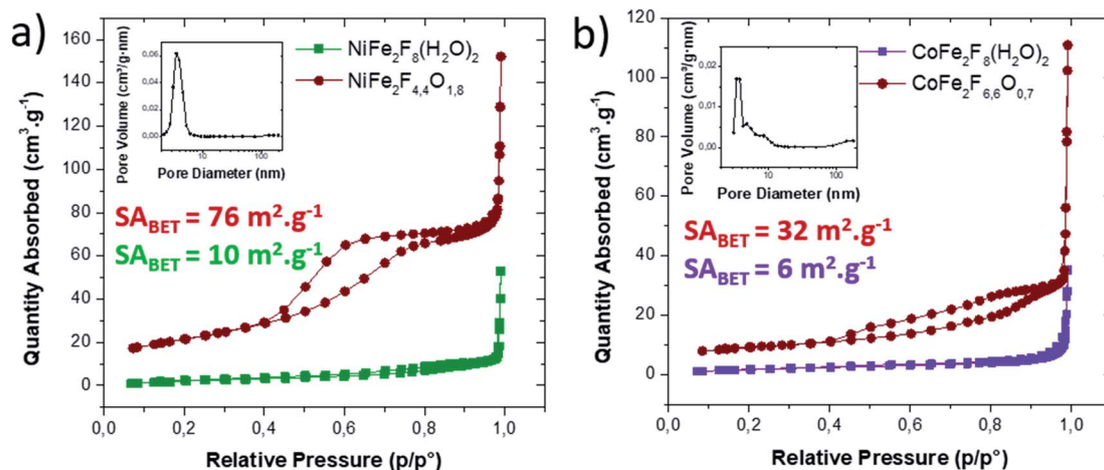


Fig. 4  $N_2$  adsorption/desorption isotherm of (a)  $NiFe_2F_{4.4}O_{1.8}$  and (b)  $CoFe_2F_{6.6}O_{0.7}$  before (green/purple) and after calcination (brown). Inset: corresponding BJH pore size distribution analyzed from the desorption branch.

also evolve into similar phases. The differences in peak shapes here reflect variance in the physical and electronic structure of the transition metals in each material. We also did not witness drastic changes in the CV shape or increases in OER activity as is sometime observed in Ni-Fe and Co-Fe oxides as a result of structural or compositional changes during this conditioning phase.<sup>46,47</sup> Following this redox transformation, the OER kinetics are exceptionally high, evidenced by the low Tafel slopes (40–60  $mV\ dec^{-1}$  range) (Fig. 6b) and low overpotential to attain a geometric current density of 10  $mA\ cm^{-2}$  (250–350  $mV$ ) (Fig. 6c). The performance of the (oxy)fluorides is comparable to state-of-the-art mixed-metal oxide OER catalysts.<sup>48,49</sup> In this series, the  $NiFe_2F_{4.4}O_{1.8}$  was consistently the highest performing material. The series of materials' performance is likely also influenced by their surface area and consequently the quantity of active sites exposed to the solution, with  $NiFe_2F_{4.4}O_{1.8}$  exhibiting both the highest surface area ( $76\ m^2\ g^{-1}$ ) and electrocatalytic

activity. An interesting point is that each material exhibits a current crossover in the CV, around 1.5 V (*i.e.* the current in the reverse scan is slightly higher). This may indicate an *in situ* surface reconstruction during the CV that serves to activate the material during the CV cycle and could be an interesting aspect to explore with *operando* techniques in the future.

The precise benefit to OER catalysis conferred by the F anions within  $NiFe_2F_{4.4}O_{1.8}$  is illustrated by comparing its electrochemical response to the crystalline  $NiFe_2O_4$  (Fig. 6f). The electron-withdrawing nature of F is evident through a shift in the redox potential for the Ni-oxidation peak, which is shifted 50  $mV$  more positive. This indicates that the Ni is harder to oxidize in  $NiFe_2F_{4.4}O_{1.8}$ . However, once oxidized the  $NiFe_2F_{4.4}O_{1.8}$  catalyst oxidized water much more rapidly, with an earlier onset potential and quickly increasing OER current. In contrast,  $NiFe_2O_4$  requires <100  $mV$  overpotential to attain similar currents.

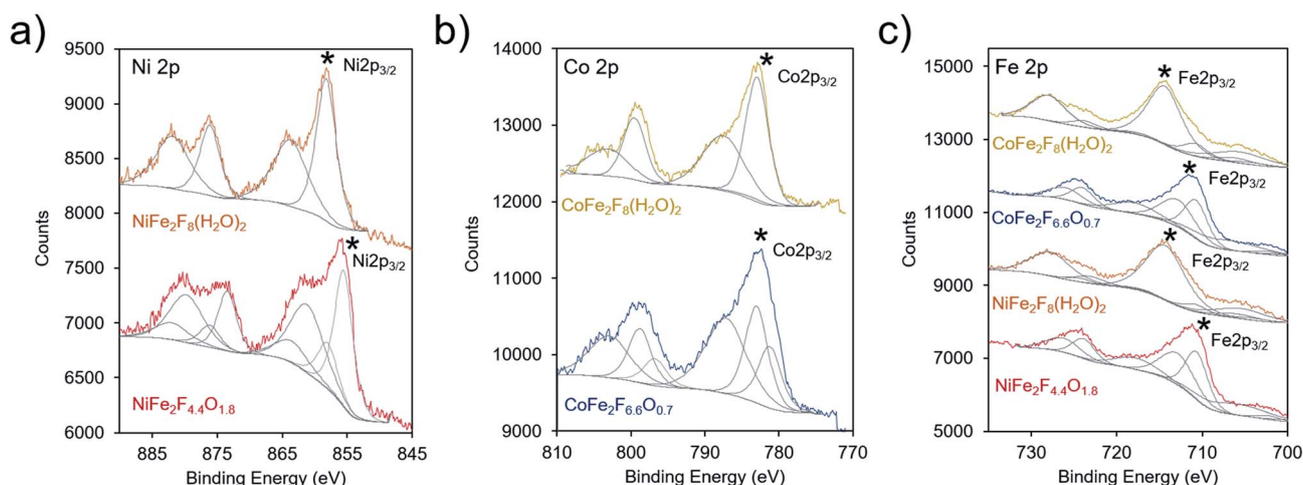
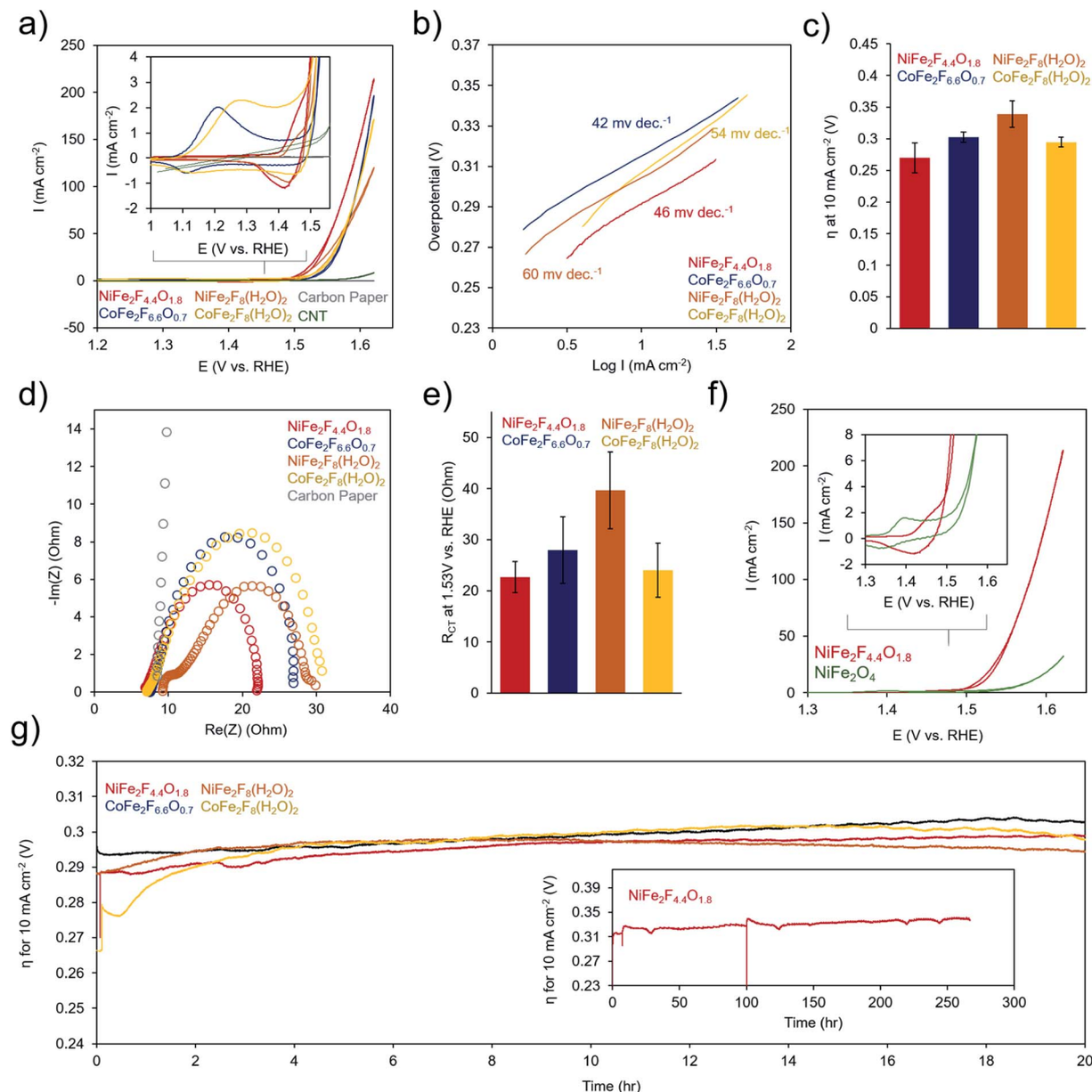


Fig. 5 XPS spectra of the (oxy)fluorides and oxyfluorides. The (a)  $Ni\ 2p_{3/2}$ , (b)  $Co\ 2p_{3/2}$  and (c)  $Fe\ 2p_{3/2}$  were acquired and spectra subsequently fit to (grey traces) compare with database data of F-free oxides. The  $2p_{3/2}$  peaks discussed are denoted with a \*.





**Fig. 6** Electrochemical characterization of oxyfluoride catalysts. (a) Cyclic voltammetry of each material's currents and redox prior to catalysis in the inset. (b) The Tafel slopes for each material lie in the 40–60  $\text{mV dec}^{-1}$  range. (c) Average and standard deviation of overpotential necessary to attain  $10 \text{ mA cm}^{-2}$ . (d and e) This EIS measurements, acquired at 300 mV overpotential, point to  $\text{NiFe}_2\text{F}_{4.4}\text{O}_{1.8}$  as the catalyst with the lowest charge transfer resistance (f) the beneficial effect of the fluorine species is evident when comparing the Ni oxyfluoride to the Ni spinel, in which the redox wave is shifted in the positive direction for the spinel and the OER catalysis is slower. (g) The performance was tested over a prolonged 20 h chronopotentiometric test at  $10 \text{ mA cm}^{-2}$  with  $\text{NiFe}_2\text{F}_{4.4}\text{O}_{1.8}$  measured over 270 h.

Electrochemical Impedance Spectroscopy (EIS) was performed at 300 mV overpotential for all samples and the data, which was in the form of a semi-circle, was fit using a Randles equivalent circuit model (Fig. 6d and e). However,  $\text{NiFe}_2\text{F}_8(\text{H}_2\text{O})_2$  featured two semi-circles, indicative of both a charge-transfer resistance and a significant resistance from the material's limited conductivity had to be fit with a separate model. The models and fitting are presented in Fig. S5.† In these spectra, the high-frequency intercept at  $\sim 7$  ohms reflects the solution-resistance and the low-frequency intercept of the

semicircle at around 20–40 ohms resistance corresponds to the charge transfer kinetics at the catalyst surface. The lowest charge-transfer resistance of  $\text{NiFe}_2\text{F}_{4.4}\text{O}_{1.8}$  ( $23 \pm 3$  ohms) corresponds to its rapid OER catalysis. Finally, the stability of (oxy)fluorides was evaluated through chronopotentiometric measurements at  $10 \text{ mA cm}^{-2}$  (Fig. 6g). Over a period of 20 h, each sample experiences only minimal ( $\sim 30$  mV) performance losses. The stability of  $\text{NiFe}_2\text{F}_{4.4}\text{O}_{1.8}$  is demonstrated with a duration of  $\sim 270$  h (Fig. 6g inset). Prior to efforts at optimization of material structure and morphology, these initial



electrochemical results already indicate that these (oxy)fluorides are excellent candidates for OER-enabled technologies such as electrolyzer or air batteries.

To elucidate the molecular dynamics (oxy)fluorides throughout the catalytic process, Raman spectroscopy was used to probe them before and after catalysis (4 h chronoamperometry at 300 mV overpotential in 1 M KOH). Spectra of crystalline  $\text{NiFe}_2\text{F}_8(\text{H}_2\text{O})_2$  and  $\text{CoFe}_2\text{F}_8(\text{H}_2\text{O})_2$  shows several strong bands, as common to crystalline metal-oxides (Fig. 7a and b). However, the bands of hydrated fluorides significantly widen and decrease in intensity, indicating a loss of crystallinity during catalysis. In contrast, the spectra of the amorphous oxyfluorides,  $\text{NiFe}_2\text{F}_{4.4}\text{O}_{1.8}$  and  $\text{CoFe}_2\text{F}_{6.6}\text{O}_{0.7}$ , show a small evolution with weak and wide bands before and after OER testing. Likely, the OER-active state of each material is an amorphous final state on the material's surface and is at a higher-valence oxidation state than the as-made material. The spectra of these materials after catalysis do not match those of  $\text{NiOOH}$ ,<sup>50</sup>  $\text{CoOOH}$ ,<sup>50</sup> amorphous  $\text{CoO}_x$ ,<sup>51,52</sup> or various iron oxide phases,<sup>53</sup> indicating that the surface of the (oxy)fluorides tested here are not transformed to  $\text{NiOOH}$  or  $\text{CoOOH}$ .

The TEM analysis was also conducted on the amorphous  $\text{NiFe}_2\text{F}_{4.4}\text{O}_{1.8}$  and  $\text{CoFe}_2\text{F}_{6.6}\text{O}_{0.7}$  after the OER catalysis (Fig. S6†). In both cases, these oxyfluorides remain amorphous

confirming the Raman analysis. Their morphology slightly changes to thin sheet-like appearance and no additional amorphous layer is observed. Metal leaching was also observed during catalysis. Indeed, the  $\text{Fe}^{3+}$  to  $\text{M}^{2+}$  ratios measured by EDS-TEM vary from 2 to 1.4 for  $\text{NiFe}_2\text{F}_{4.4}\text{O}_{1.8}$  corresponding to a Fe leaching and from 2.1 to 2.4 for  $\text{CoFe}_2\text{F}_{6.6}\text{O}_{0.7}$  corresponding to a Co leaching. Those differences in the composition through leaching are in good agreement with the slight differences in the Raman spectra.

In order to probe surface intermediates and rate-limiting steps of the OER cycle of the (oxy)fluoride materials, we utilized methanol oxidation as facile method to detect surface-bound \*OH. As \*OH is a very electrophilic intermediate, it will react with methanol, and thus give rise to methanol oxidation currents when present in substantial quantities.<sup>54</sup> Upon the addition of 10 mM methanol, we noted enhanced currents beginning at 1.0 V vs. RHE for all examples except for  $\text{NiFe}_2\text{F}_8(\text{H}_2\text{O})_2$  (Fig. S7†). This result points to (though does not prove) \*OH coverage on the (oxy)fluoride surfaces prior to OER initiation and that the \*OH deprotonation step as possibly being rate limiting. On the other hand, the limiting step for  $\text{NiFe}_2\text{F}_8(\text{H}_2\text{O})_2$  may be the adsorption of \*OH. This is especially interesting as even changes in a material's stoichiometry induce notable changes in mechanism.

While a complete mechanistic picture of these oxyfluoride materials is not yet available, there exist a number of promising approaches to obtain complementary pieces to this puzzle. X-ray absorption spectroscopy (XAS), especially when operated *in situ*, has been previously used to elucidate how changes in oxidation states and chemical environments of the transition metal species influence catalytic activity in metal-oxide materials and would be similarly useful to this system.<sup>49,55–58</sup> XPS, performed in specialized instrumental setups would also provide complementary information regarding electronic structure as a function of applied bias and reaction time.<sup>59</sup> Furthermore, techniques such as electrochemical quartz crystal microbalance measurements may impart information on voltage-dependent surface or bulk reconstruction by cross-comparing currents and *in situ* changes in mass.<sup>60</sup>

As a proof of concept, the best performing material,  $\text{NiFe}_2\text{F}_{4.4}\text{O}_{1.8}$ , was combined with another earth-abundant hydrogen evolution reaction (HER) catalyst, cobalt sulfide ( $\text{CoS}_x$ ),<sup>61,62</sup> and utilized in an overall water electrolysis cell (Fig. 8a and S8†). In a two-electrode configuration, overall water electrolysis initiated at  $\sim 1.60$  V and reached  $100 \text{ mA cm}^{-2}$  at 1.80 V. Chronopotentiometric testing at  $10 \text{ mA cm}^{-2}$  pointed to a stable performance of this composite system at  $\sim 1.65$  V for 24 h. In this configuration, the OER overpotential was  $\sim 270$  mV, in line with some of the highest-performing Ni-Fe oxides<sup>48</sup> and the HER overpotential was  $\sim 100$  mV. This metric is comparable to the performance achieved with benchmark precious-metal (*e.g.* Pt, Ir, Ru) containing systems<sup>63</sup> that also need typically 1.55–1.60 V to reach  $10 \text{ mA cm}^{-2}$  and points to the promise of oxyfluorides as cost-effective OER components of next-generation electrochemical technologies.

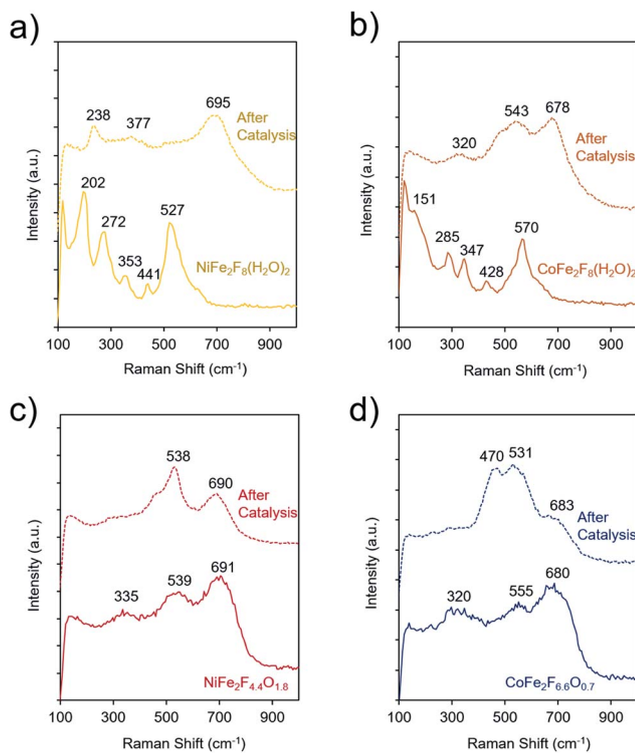


Fig. 7 Raman spectroscopy of the (oxy)fluorides before and after catalytic testing (4 h at 1.53 V in 1 M KOH). The spectra of crystalline  $\text{NiFe}_2\text{F}_8(\text{H}_2\text{O})_2$  (a) and  $\text{CoFe}_2\text{F}_8(\text{H}_2\text{O})_2$  (b) exhibit a number of strong bands which are lost and give way to broader bands after catalysis. In contrast, the amorphous  $\text{NiFe}_2\text{F}_{4.4}\text{O}_{1.8}$  (c) and  $\text{CoFe}_2\text{F}_{6.6}\text{O}_{0.7}$  (d) feature broader bands which undergo considerably less changes during catalytic testing.



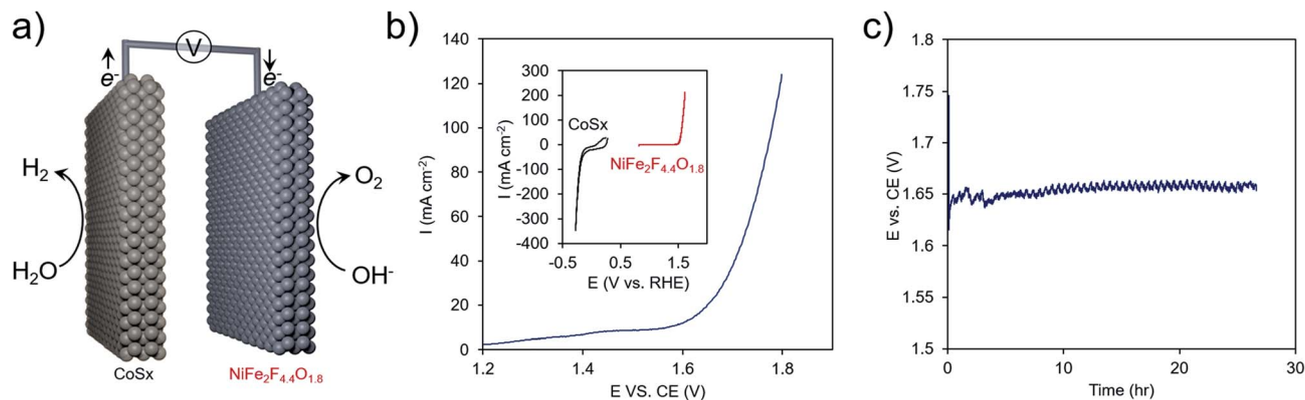


Fig. 8 (a) Overall water electrolysis with NiFe<sub>2</sub>F<sub>4.4</sub>O<sub>1.8</sub> as the anode. (b) A NiFe<sub>2</sub>F<sub>4.4</sub>O<sub>1.8</sub> was integrated with a CoS<sub>x</sub> cathode to put together an overall water electrolysis system comprised of earth-abundant materials. (c) This system featured an onset of ~1.60 V and required ~1.65 V to generate a stable current of 10 mA cm<sup>-2</sup>.

### 3 Concluding remarks

In summary, we present a study on the synthesis and electrocatalytic applications of mixed-metal (oxy)fluorides as OER catalysts. Crystalline hydrated fluorides, CoFe<sub>2</sub>F<sub>8</sub>(H<sub>2</sub>O)<sub>2</sub> and NiFe<sub>2</sub>F<sub>8</sub>(H<sub>2</sub>O)<sub>2</sub>, were prepared by microwave heating assisted solvothermal synthesis. Subsequent calcination of the hydrated fluorides leads to the formation of amorphous oxyfluorides NiFe<sub>2</sub>F<sub>4.4</sub>O<sub>1.8</sub> and CoFe<sub>2</sub>F<sub>6.6</sub>O<sub>0.7</sub>. The (oxy)fluorides are speculated to benefit from the fluorine anions withdrawing electron density away from the Co, Ni, and Fe species, which are likely responsible for the exceptional electrocatalytic properties of each material. Finally, the best catalyst, NiFe<sub>2</sub>F<sub>4.4</sub>O<sub>1.8</sub>, associated with a CoS<sub>x</sub> HER catalyst leads to a highly performing water electrolyzer comprised of only earth-abundant element catalysts. This study may open up avenues towards the utility of (oxy)fluoride materials for energy-related applications and rational routes for harnessing inductive effects conferred by fluorine species.

## 4 Materials and methods

### 4.1 Synthesis

The hydrated fluorides M<sup>2+</sup>Fe<sub>2</sub><sup>3+</sup>F<sub>8</sub>(H<sub>2</sub>O)<sub>2</sub> (M<sup>2+</sup> = Co, Ni) was obtained by solvothermal reaction using a MARS-5 Microwave Digestion System (CEM Corp.) from starting reactants of chloride precursors (Alfa Aesar), 9.45 mL of absolute methanol 'MeOH' (233 mmol, 24.7 mol L<sup>-1</sup>, 99.8%, Sigma Aldrich) and 0.55 mL hydrofluoric acid solution (15 mmol, 27.6 mol L<sup>-1</sup>, Riedel De Haen). A constant concentration [M<sup>II</sup>] + [Fe<sup>III</sup>] = 0.1 mol L<sup>-1</sup>, a ratio [Fe<sup>III</sup>]/[M<sup>II</sup>] = 2 and a constant volume of liquid (HF and MeOH) were fixed. The M<sup>II</sup>/Fe<sup>III</sup>/HF/MeOH ratio is 1/2/44/699. The mixtures are placed in Teflon autoclaves and heated at 160 °C for 30 min with stirring. After cooling, the solid products are filtered, washed with 2 mL of ethanol and dried in a furnace under air.

CoFe<sub>2</sub>F<sub>8</sub>(H<sub>2</sub>O)<sub>2</sub> and NiFe<sub>2</sub>F<sub>8</sub>(H<sub>2</sub>O)<sub>2</sub> were put in a furnace at 320 °C and 340 °C, respectively, during 1 h (heating/cooling rate of 2 °C min<sup>-1</sup>) giving the amorphous oxyfluorides with M<sup>2+</sup>M<sub>2</sub><sup>3+</sup>F<sub>8-2x</sub>O<sub>x</sub> formulations.

### 4.2 Characterization methods

**X-ray powder diffraction (XRPD).** X-ray diffraction patterns were collected in the range 10° ≤ 2θ ≤ 150° on a Panalytical MPD-PRO diffractometer equipped with a linear X'Celerator detector with a CoK<sub>α</sub> radiation (1.789 Å) used to avoid the X-ray fluorescence. Rietveld refinements were performed by using the Fullprof profile refinement program. This diffractometer belongs to the "X-ray Diffusion and Diffraction" technical platform of IMMM (Le Mans University). Data were collected in the [10–100°] 2θ scattering angle range with a 0.0131° step.

**Scanning electronic microscopy (SEM).** SEM images of the powders were obtained using a JEOL microscope (JSM 6510LV) at an acceleration voltage of 20 kV. Elementary quantitative microanalyses were performed using an Energy Dispersive Spectrometry (EDS) OXFORD detector (AZtec software). The microscope belongs to the "Electron Microscopy" technical platform of IMMM (Le Mans University).

**Transmission electron microscopy (TEM).** The TEM study (SAED and HREM) was performed on a JEOL JEM 2100 HR electron microscope operating at 200 kV and equipped with a side entry ±35° double-tilt specimen holder. The microscope belongs to the "Electron Microscopy" technical platform of IMMM (Le Mans University). The samples for transmission electron microscopy investigation were prepared by ultrasonically dispersing the raw powder in ethanol, depositing a drop of the resulting suspension onto a holey carbon-coated copper grid and finally drying the grid in air.

**Nitrogen sorption.** N<sub>2</sub> sorption isotherms were measured at 77 K using a TriStar II 3020 (Micrometrics). The film samples were degassed under vacuum at 100 °C for 12 h prior measurement. The surface areas were calculated using the Brunauer–Emmett–Teller (BET) method.

**Thermal analysis.** Mass Spectroscopy coupled Thermo Gravitimetric Analysis (TGA-MS) was performed using a Netzch STA 449 F3 coupled with a QMS 403 C mass spectrometer. The thermoanalytical curves were recorded together with the ion current curves in the multiple ions detection probe. A constant purge nitrogen gas flow of 80 mL min<sup>-1</sup> and a constant heating rate of 5 °C min<sup>-1</sup>



were applied. The thermogravimetric (TGA) experiments were carried out with a thermoanalyzer SETARAM TGA 92 with a heating rate of  $5\text{ }^{\circ}\text{C min}^{-1}$  from room temperature up to  $900\text{ }^{\circ}\text{C}$  under dry air (Alphagaz, mixture of oxygen (20%) with nitrogen (80%),  $\text{H}_2\text{O} < 3\text{ ppm}$ ). X-ray thermodiffraction (HT-XRD) was performed under dry air in an Anton Parr XRK 900 high temperature furnace with the diffractometer already described. The samples were heated from  $40$  to  $600\text{ }^{\circ}\text{C}$  at a heating rate of  $10\text{ }^{\circ}\text{C min}^{-1}$ . X-ray diffraction patterns were recorded in the  $[5\text{--}60^{\circ}] 2\theta$  range with a scan time of  $10\text{ min}$  at  $20\text{ }^{\circ}\text{C}$  intervals from room temperature to  $400$  and at  $100\text{ }^{\circ}\text{C}$  intervals from  $400$  to  $600\text{ }^{\circ}\text{C}$ .

**Electrochemical measurements.** To fabricate electrodes, sample powders were sonicated in ethanol together with 1% Nafion and 1% multi-walled carbon nanotubes (40 nm diameter, purchased from Sigma Aldrich) to make a catalyst ink. The ink was pipetted onto Toray carbon paper to load  $1\text{ mg cm}^{-2}$  of catalyst and dried at  $80\text{ }^{\circ}\text{C}$  for 20 minutes. Typically, a geometric surface area of  $0.2\text{ cm}^2$  was employed. Electrochemical characterization was performed using a Biologic VMP 150 potentiostat equipped with impedance capability. Prior to voltammetry or amperometry, ohmic drop compensation (85%) was performed with EC-Lab software, using the ZIR function, which the solution resistance through recording the impedance value at a single, high frequency value (100 kHz) at open circuit. The measured solution resistance was usually around 5–8 ohms with a distance of approximately 2 cm between the working and reference electrodes. A carbon rod and Ag/AgCl electrode were used as counter and reference electrodes. To ensure that the potential of the reference electrode did not drift over time, the reference was periodically checked against a “master” electrode which did not undergo testing. CVs were later performed by making an equivalent ink but with the Nafion omitted, as its acidic nature may have had an obstructive effect on OER catalysis (Fig. S9†). Methanol oxidation was performed by measuring two consecutive CVs, one in  $1.0\text{ mol L}^{-1}$  KOH and another immediately after the addition of a small quantity of methanol to reach 10 mM concentration. The solution was stirred during electrolysis with a Teflon stir bar at 1000 rpm. For two-electrode measurements of overall water electrolysis,  $\text{CoS}_x$  was prepared through electrodeposition onto carbon paper from a  $\text{CoCl}_2$  and thiourea containing aqueous solution, using a well-established recipe.<sup>62</sup> Briefly, a carbon paper electrode was cycled in a pH 7, 100 mM  $\text{CoCl}_2$  and 0.5 M thiourea between  $-0.2$  and  $-1$  for 4 cycles, then  $-1.2$  and  $0.3\text{ V vs. Ag/AgCl}$  for 26 cycles at  $5\text{ mV s}^{-1}$ , rinsed with water, dried and kept in ambient conditions prior to use. A smaller voltage was used for the first 4 cycles to promote homogeneous coverage of the electrode with the  $\text{CoS}_x$ .

**Raman spectroscopy.** Raman spectra were acquired using a Renishaw InVia spectrometer and a 514 nm 30 W laser. Spectra were first acquired at low power (99.5% laser attenuation) to ensure that the laser irradiation of the samples did not alter them. Typical acquisition times were 180 seconds. Several spectra were acquired to verify sample homogeneity and representative spectra were incorporated into the manuscript.

**X-ray photoelectron spectroscopy.** XPS spectra were acquired with a VG ESCALAB 3 Mark II spectrometer with a Mg K $\alpha$  source operating at 300 W (15 kV, 20 mA). Prior to characterization,

samples were deposited onto conductive copper tape and no other modifications made. The C 1s peak (285.0 eV) was used for energy calibration and the background for all spectra was subtracted using a Shirley method. High-resolution scans were taken at 0.1 eV steps following a survey scan at 1.0 eV step size.

## Conflicts of interest

There are no conflicts to declare.

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