How oxidation state and lattice distortion influence the oxygen evolution activity in acid of iridium double perovskites†

Maria Retuerto,*a Laura Pascual,b Oriol Piqué,c Paula Kayser,d Mohamed Abdel Salam,e Mohamed Mokhtar,†d José Antonio Alonso,†d Miguel Peña,a Federico Calle-Vallejo ic,e and Sergio Rojas id,a

In view of iridium’s scarceness and high cost, Ir-containing catalysts for water electrolyzers should have low loadings and maximal utilization. Here, we studied low-Ir-content, highly active, double perovskites (Sr2MIrO6, M = Ni, Co, Sc and Fe) for the oxygen evolution reaction (OER) in acid combining electrochemical experiments, DFT, and advanced characterization techniques. The initial OER performance depends on Ir’s oxidation state and the geometric features of IrO6 frameworks, which are tuned by the choice of M. Higher oxidation states, particularly Ir6+, enhance the OER activity: Sr2NiIrO6 and Sr2CoIrO6 display potentials of ~1.53 V at 10 mA cm⁻², comparable to the best Ir-based catalysts in the literature. However, because of their less symmetric structures, perovskites with Ir6+ are less stable, prone to surface reconstruction and their cations leach under OER conditions. These results show that improved iridium-based OER electrocatalysts in acid can be designed by balancing their activity and stability.

Water electrolysis comprises two half reactions: the formation of hydrogen at the cathode (in acid: 4H⁺ + 4e⁻ → 2H₂, hydrogen evolution reaction, HER) and the formation of oxygen at the anode (in acid: 2H₂O → O₂ + 4H⁺ + 4e⁻, oxygen evolution reaction, OER). In acidic media, the overall process is limited by the sluggish kinetics of the OER and the highly corrosive environment associated with water electrolysis. As a consequence, iridium- and ruthenium-based oxides are the state-of-the-art catalysts for the OER. Ru-based oxides typically display the lowest OER overpotentials; however, their stability in acid is insufficient. Therefore, benchmark catalysts are based on IrO₂ and IrO₃/TiO₂, yielding current densities of 10 mA cm⁻² at ~1.52 V and mass activities of ~30–50 A g⁻¹ at 1.5 V.

Since Ir is one of the rarest, ill-distributed and most expensive elements on the Earth’s crust, traded in a volatile market where slight fluctuations in demand can significantly impact supply, it is necessary to make the best possible utilization of it. In this sense, mixed oxides are promising candidates to replace pure iridium oxides as electrocatalysts for the OER. For instance, the mass percentage of Ir in IrO₂ is 86 wt%, and goes down to 59 wt% in SrIrO₃ and 36–38 wt% in Sr₂MIrO₆, if M is a first-row transition metal. To the best of our knowledge, the iridium mixed oxides with OER activity in acid electrolyte reported so far are: (i) Ir-based perovskites such as SrIrO₃/IrO₂ thin films, SrCo₀.₃Ir₀.₇O₃, Ba₂RIrO₆ (R = Y, La, Ce, Pr, Nd, Tb), La₁₂Ir₁₀O₃₅, 6H-SrIrO₃, Ir₁–SrTiO₃, Sr₂FeIrO₆ and Sr₂CoIrO₆ (ii) Ir–pyrophlores such as Bi₂Ir₂O₇, Pb₂Ir₂O₆,
Pr$_2$Ir$_2$O$_7$, Y$_2$Ir$_2$O$_7$, (iii) spinels such as Ni$_{0.34}$Co$_{0.44}$Ir$_{0.2}$O$_4$, and (iv) Ruddlesden–Popper phases such as Sr$_2$IrO$_4$ and Sr$_2$IrO$_6$.

Looking at the performance of the aforementioned compounds, we hypothesize that the OER activity of Ir mixed oxides is related to the actual environment of Ir cations in such oxides. In other words, the OER activity is likely affected by the nature of the foreign metal(s) and the actual location of the cations in the oxides. However, the roles of the foreign metals, the crystal structure including conformation, the distortions, the bond strength of the IrO$_6$ octahedra (Oh), and that of Ir’s oxidation state on the OER performance of Ir mixed oxides, are not fully understood. Evidently, this knowledge void prevents the rational design of advanced Ir-based OER electrocatalysts.

A number of studies suggest a direct relationship between the environment and oxidation state of Ir in its oxides and their OER activity. The most usual Ir oxidation states in oxides are +3 and +4, as in Sr$_2$TaIrO$_6$, Sr$_2$NbtIrO$_6$, and Sr$_2$TiIrO$_6$. However, such perovskites present cationic disorder with Ta, Nb, or Ti and Ir randomly occupying the same crystallographic position, thereby preventing the study of the actual role of Ir’s environment for the OER. A recent study illustrated the relevance of upper iridium oxides for the OER. Oxides with Ir$^{5+}$ or Ir$^{6+}$ are less common; however, double perovskites with Ir cations in upper oxidation states have been reported, for instance, Ba$_2$Ir$_2$O$_6$ (R = Ce, Tb, Y, La, Pr, Nd), La$_2$IrIrO$_6$, Sr$_2$FeCoIrO$_6$ (ref. 26) and Sr$_2$(Ca, Mg)IrO$_4$ (ref. 27). These double perovskites exhibit cationic ordering between R, Li, Fe, Co, Ca or Mg and Ir; being Ir cations located in an independent crystallographic position than the other metals, which allows studying the impact of Ir’s environment for the OER.

The suitability of OER electrocatalysts should be assessed holistically by studying their catalytic performance, including activity, durability, and stability. It is well known that the activity depends on the oxygen adsorption strength of the active sites. This strength depends on several factors, such as the position of the oxygen’s p-band with respect to the Fermi level, which affects the activity, durability and stability. It is well known that the activity, durability, and stability. The bond strength of the IrO$_6$ octahedra (Oh), and that of Ir’s oxidation state on the OER performance of Ir mixed oxides, are not fully understood. Evidently, this knowledge void prevents the rational design of advanced Ir-based OER electrocatalysts.

2. Experimental section

2.1 Synthesis of Sr$_2$MIrO$_6$ perovskites

The synthesis of Sr$_2$MiRo$_6$ (M = Ni, Co, Sc, Fe) is detailed elsewhere. Briefly, the samples were prepared by sol–gel methods. We used a solution of citric acid and HNO$_3$ to dissolve the starting materials. IrO$_2$ was not dissolved but stayed in suspension under magnetic stirring. The suspension was slowly evaporated leading to organic resins containing a homogeneous distribution of the involved cations. After evaporation, the resins were dried at 140 °C and heated at 600 °C for 12 h to decompose the organic materials and nitrates. Next, Sr$_2$NiIrO$_6$ was annealed at 900 °C for 12 h in O$_2$ flow and subsequently at 900 °C under 200 bar of O$_2$ for 48 h, to obtain the Ir$^{6+}$ perovskite oxide. For Sr$_2$MIrO$_6$ (M = Co, Sc, Fe) oxides the reactive precursors were heated in air at 1100 °C for 12 h.

2.2 Physicochemical characterization

The phase identification and purity of the obtained perovskites was determined by X-ray powder diffraction (XRD) in the Bragg–Brentano reflection geometry with CuK$_\alpha$ radiation (λ = 1.5418 Å). A detailed study of the crystal structure using Powder Neutron Diffraction (PND) and the analysis of Ir oxidation states and coordination environments using X-ray Absorption Spectroscopy (XAS) is available in the literature.

Transmission Electron Microscopy (TEM), high-resolution electron microscopy (HRTEM) and X-ray energy dispersive spectra (EDX) were recorded in a 200 kV (JEOL 2100F) field emission gun transmission electron microscope equipped with an EDX X-Max 80 spectrometer (Oxford Instruments). The specimens were prepared by depositing aliquots onto a Cu grid supporting a lacy carbon film.

X-ray photoelectron spectra (XPS) were acquired with a VG ESCALAB 200 R at a pass energy of 50 eV using a Mg Kz X-ray source. The kinetic energies of the photoelectrons were measured using a hemispherical electron analyser working in the constant-pass energy mode. A background pressure below 3 × 10$^{-8}$ mbar was kept in the chamber during data acquisition. The binding energies were determined by setting the C 1s peak at 284.6 eV. 200 scans in increments of 0.1 eV with dwell times of 50 ms were collected in order to increase the signal-to-noise ratio.
2.3 Electrochemical characterization

An Autolab PGStat 302N potentiostat/galvanostat was used to test the electrochemical performance of the oxides. The measurements were performed using a standard three-compartment glass cell and a rotating disk electrode (RDE) (Pine Research Instruments). A graphite bar was used as counter electrode. An Ag/AgCl electrode was used as reference electrode. The perovskites were deposited as an ink on a 0.196 cm\(^2\) glassy carbon electrode. The ink was prepared by mixing 5 mg of oxide with 1 mg of carbon black (Vulcan-XC-72R) to improve the electrical conductivity. The mixture was dispersed in tetrahydrofuran (THF) and 5\% Nafion and sonicated using an Ultrasonic Processor UP50H (Hielscher). The composition of the ink was 5 mg\(_{\text{o}xide}\) 1 mg\(_{\text{vulcan}}\), 0.03 mL\(_{\text{THF}}\), 10 \(\mu\)L of the ink were dropped onto the electrode to obtain a catalyst loading of 0.255 mg\(_{\text{o}xide}\) cm\(^{-2}\), or ~0.97 mg\(_{\text{ir}}\) cm\(^{-2}\), depending on the oxide.

The OER catalytic performance of the perovskites was evaluated by recording linear sweep voltammograms between 1.1 and 1.7 V at 10 mV s\(^{-1}\). The measurements were performed in an O\(_2\) saturated 0.1 M HClO\(_4\) electrolyte to assure the O\(_2\)/H\(_2\)O equilibrium at 1.23 V, at a rotation rate of 1600 rpm. The OER kinetic curves were capacitance-corrected by using the average of the anodic and cathodic curves and iR-corrected by using the formula \(E - iR_{\text{corrected}} = E_{\text{applied}} - iR\), where \(i\) is the current and \(R\) is the uncompensated resistance (\(R \sim 29 \Omega\)) as obtained from Electrical Impedance Spectroscopy (EIS) at open voltage.

Durability tests were performed by recording 50 consecutive OER cycles at 10 mV s\(^{-1}\) between 1.2 and 1.7 V vs. RHE, or by chronoamperometric tests by fixing the initial potential for each catalyst to the potential needed to reach 10 mA cm\(^{-2}\) and monitoring the evolution of the current density with time. The durability of Sr\(_2\)NiIrO\(_6\) and Sr\(_2\)CoIrO\(_6\) was further tested by recording 10 000 consecutive cycles between 1.2 and 1.7 V at 50 mV s\(^{-1}\).

2.4 Computational details

The VASP code was used to perform the DFT calculations,\(^{30}\) RPBE was the chosen exchange-correlation functional,\(^{31}\) and the projector augmented-wave (PAW) method was used to describe the effect of the inner cores on the valence electron density.\(^{32}\) The free energies of reaction were approximated as \(\Delta G = \Delta E_{\text{DFT}} + \Delta ZPE - T \Delta S\), where \(\Delta E_{\text{DFT}}\) is the DFT-calculated reaction energy and \(\Delta ZPE\) is the zero-point energy change between reactants and products. The ZPEs were obtained from vibrational frequency calculations using the harmonic approximation. As usual,\(^{31,44}\) the term \(T \Delta S\) includes the total entropies of the molecules involved in the OER, and neglects those of the adsorbates. The computational hydrogen electrode (CHE) was used to model proton–electron pairs.\(^{42}\) We assumed the following catalytic pathway for the OER: \(^{43,44}\)

\[ \text{*OOH} \rightarrow \text{*O}_2 + \text{H}^+ + e^- \]  
(4)

Within the CHE framework, the free energies of reaction can be written as a function of the adsorption energies of the three adsorbed intermediates, namely *O\(_x\), *OH, and *OOH:

\[ \Delta G_1 = \Delta G_{\text{OH}} \]  
(5)

\[ \Delta G_2 = \Delta G_{\text{O}} - \Delta G_{\text{OH}} \]  
(6)

\[ \Delta G_3 = \Delta G_{\text{OOH}} - \Delta G_{\text{O}} \]  
(7)

\[ \Delta G_4 = \Delta G_{\text{O}_2} - \Delta G_{\text{OOH}} \]  
(8)

where \(\Delta G_{\text{O}_2} = 4.92\ eV\) corresponds to the standard equilibrium potential (\(E^0 = 1.23\ V\)) multiplied by the total number of electrons per catalytic cycle (1.23 V x 4e\(^{-}\) = 4.92 eV). The adsorption energies of *O\(_x\), *OH and *OOH are defined using H\(_2\)O[l] and proton–electron pairs and are provided in Table S1 for the perovskites under study. We extracted from the free energies in eqn (5)–(8) the potential-limiting step, which determines the calculated OER overpotential (\(\eta_{\text{OER}}\)): \(^{43,44}\)

\[ \eta_{\text{OER}} = \max(\Delta G_1,\Delta G_2,\Delta G_3,\Delta G_4)/e^- - 1.23 \ V \]  
(9)

As shown in previous publications,\(^{46,47}\) the concept of electrocatalytic symmetry as expressed by the electrochemical-step symmetry index (ESSI) is a good descriptor for the OER overpotential:

\[ \text{ESSI} = \frac{1}{n} \sum_{\text{i}} \left( \frac{\Delta G_i^-}{e^- - E^0} \right) \]  
(10)

where \(\Delta G_i^-\) corresponds to the reaction energies in eqn (5)–(8) larger than 1.23 eV.

A full description of \(\delta-\varepsilon\) optimization can be found elsewhere.\(^{47}\) Briefly, the method is based on the introduction of two parameters into the adsorption energies: (I) \(\delta\), which is scaling-based and affects *O\(_x\), *OH and *OOH, and (II) \(\varepsilon\), which is scaling-free and affects *OOH only. Both parameters are in units of energy. Normally, reasonable values for \(\delta\) are in the range \([-0.3, 0.3]\) eV, whereas those of \(\varepsilon\) are in the range \([-0.3, 0]\) eV. \(\delta\) is either a weakening or strengthening of the adsorption energies, while \(\varepsilon\) is only a strengthening. See Section S2 of the ESI\(^\dagger\) for more details and the specific values obtained for the double perovskites containing Ni and Co.

We note that there are known inaccuracies of DFT for the OER modelling of some oxides e.g. RuO\(_2\),\(^{48,49}\) and some solutions have been proposed.\(^{46,50,51}\) In particular, the inaccuracies of DFT observed before for SrIrO\(_3\) (ref. 19) were corrected for in this study by using thick slabs. For further information on the computational details please refer to Section S1 of the ESI\(^\dagger\).

3. Results and discussion

3.1 Double-perovskite structure and composition

The incorporation of different M metals into Sr\(_2\)MIrO\(_6\) is expected to result in materials with different structural and chemical features, namely unit-cell volume, Ir oxidation state,
Ir–O and M–O bond lengths, structural distortions, etc. In the following, we intend to disclose the effect of those features on the OER performance of iridium mixed oxides. Fig. S1 in the ESI† shows the XRD diffractograms for Sr$_2$MlIrO$_6$ (M = Ni, Co, Sc and Fe) and the evolution of their unit-cell volumes. All the oxides are pure and exhibit the perovskite crystal structure. The volume of the unit cell increases with M in the order Co$^{3+}$ < Fe$^{3+}$ < Ni$^{3+}$ < Sc$^{3+}$, as expected from their ionic radii. Powder Neutron Diffraction (PND) data reveal that all perovskites have monoclinic symmetry with unit-cell parameters $a = b 
eq c$ and $\alpha = \gamma = 90^\circ \neq \beta$, in the space groups $P2_1/n$ (a’c’’c’’ tilting, Glazer notation) for Ni and Sc oxides, and $I2/m$ (a’’b’’b’’ tilting) for Co and Fe oxides. Both $P2_1/n$ and $I2/m$ symmetries are characteristic of distorted double perovskites with alternating MO$_6$ and IrO$_6$ complexes along the three crystallographic directions in a long-range Oh M/Ir ordering.55,56

Oxygen vacancies can play a key role in the OER activity of oxides. Therefore, the presence or absence of oxygen vacancies in the double perovskites was assessed by means of PND. This technique is one of the most reliable ones to evaluate the occupancy of light atoms such as oxygen in oxides.55 The refinement of the crystal structures reveals the absence of oxygen vacancies in Sr$_2$MlIrO$_6$, irrespectively of the atom in the M site. PND data also reveal a stoichiometric composition of the catalysts in the bulk.55,56 The most relevant crystallographic features obtained from PND are detailed in Table S2,† for instance, oxygen vacancies, Ir oxidation state and (Ir–O) distances. The surface composition of Sr$_2$MlIrO$_6$ was studied by XPS, providing a M/Ir ratio close to the nominal value of 1 in all samples. Except for Sr$_2$NiIrO$_6$, the surface is enriched in Sr in all of the studied oxides, showing Sr/Ir surface atomic ratios in the range of 1.9 to 3.1. This observation is in line with previous reports on Sr-containing perovskites, where it was reported that SrCO$_3$ forms at the surface.56

The oxidation state of M and Ir was studied by XAS and bond-valence sum (BVS) calculations using Brown’s model with PND data.33,39,57,58 XAS data indicate that Sr$_2$NiIrO$_6$ perovskites contain Ni$^{2+}$ and Ir$^{4+}$ cations, whereas the oxidation states in Sr$_2$CoIrO$_6$, Sr$_2$ScIrO$_6$ and Sr$_2$FeIrO$_6$ are close to M$^{3+}$ and Ir$^{5+}$. However, it should be noted that Ir$^{5+}$ sites are slightly more oxidized in Sr$_2$CoIrO$_6$ than in Sr$_2$FeIrO$_6$. BVS gives another estimation of the oxidation states using the bond distances calculated from PND. This technique is one of the most reliable ones to evaluate the characteristic of distorted double perovskites with alternating MO$_6$ and IrO$_6$ complexes along the three crystallographic directions in a long-range Oh M/Ir ordering.55,56

3.2 Initial OER activity: effect of M in the catalytic performance

Fig. 1a shows the catalytic activity for the OER of Sr$_2$MlIrO$_6$ (M = Ni, Co, Sc and Fe) in terms of iR-corrected current densities, normalized to the geometric area of the electrodes (0.196 cm$^2$). The non iR-corrected curves and the R obtained from EIS at open-circuit voltage are shown in Fig. S3 and S4 in the ESI.†

The current density decreases with M as Ni$^{2+}$ > Co$^{3+}$ > Sc$^{3+}$ > Fe$^{3+}$. As shown in Table 1, the current densities reached at 1.5 V are 3.3, 2.6, 1.3, and 0.3 mA cm$^{-2}$ for Sr$_2$NiIrO$_6$, Sr$_2$CoIrO$_6$, Sr$_2$ScIrO$_6$, and Sr$_2$FeIrO$_6$, respectively. The trends can also be observed in the Tafel plots in Fig. 1b.

![Fig. 1](image-url)
Taking the potential to achieve 10 mA cm$^{-2}$ as an RDE-based metric to benchmark catalysts, as suggested by Jaramillo et al.,$^{31}$ we observe that Sr$_2$NiIrO$_6$ records the lowest positive potential of 1.525 V, followed by Sr$_2$CoIrO$_6$ and Sr$_2$ScIrO$_6$ with potentials of 1.535 and 1.560 V, respectively. Clearly, Sr$_2$NiIrO$_6$ and Sr$_2$CoIrO$_6$ have similar current densities; however, Sr$_2$ScIrO$_6$ and Sr$_2$FeIrO$_6$ are less active, suggesting that M plays an important role in the OER performance of Sr$_2$MIrO$_6$. Sr$_2$FeIrO$_6$ records the lowest performance in the whole polarization range, showing a potential of 1.650 V to achieve 10 mA cm$^{-2}$.

Table 1 compares the electrochemical performance of the perovskites under study with state-of-the-art Ir mixed oxides. Sr$_2$NiIrO$_6$, Sr$_2$CoIrO$_6$, Sr$_2$ScIrO$_6$, and Sr$_2$FeIrO$_6$ display comparable or higher activity than the best Ir mixed oxides in the literature for the OER in acid media, including SrCo$_0.9$Ir$_{0.1}$O$_3$, Sr$_2$FeIrO$_6$, Sr$_2$ScIrO$_6$, and Sr$_2$CoIrO$_6$ which have a potential of ~1.47 V at 10 mA cm$^{-2}$. In all cases, it is assumed that Ir sites are the main active sites for the OER, since the overpotentials achieved by Ir mixed oxides are comparable to Ir–O phases.$^{32}$

The Tafel slopes obtained at low current densities are also shown in Table 1. The Tafel slopes for Sr$_2$NiIrO$_6$ (48 mV dec$^{-1}$) and Sr$_2$CoIrO$_6$ (52 mV dec$^{-1}$) are in line with the values reported for the best Ir mixed oxides. Indeed, most Ir- and Ru-based catalysts, including IrO$_2$ and RuO$_2$, display Tafel slopes between 40 and 60 mV dec$^{-1}$.

The initial OER activities were first correlated with the oxidation state of the Ir cations in each catalyst (Fig. 1a, inset). The trends in Fig. 1a reveal that the OER activity increases alongside the oxidation state of Ir in the double perovskites. Specifically, Sr$_2$NiIrO$_6$ with Ir$^{5+}$ cations displays the highest OER activity, followed by Sr$_2$CoIrO$_6$ with Ir$^{6+}$. The perovskites with Ir$^{5+}$ cations (Sr$_2$ScIrO$_6$ and Sr$_2$FeIrO$_6$) are the least active ones in the series. The importance of Ir’s oxidation state for the OER performance was observed previously in the La$_2$LiRo$_6$ double perovskite.$^{23}$ It was claimed that Ir$^{5+}$O$_6$ are not active for the OER, so that a previous activation step oxidizing Ir$^{5+}$ to Ir$^{6+}$ is needed to make Ir$^{5+}$-mixed oxides active. This suggests that the higher OER activity of Sr$_2$NiIrO$_6$ and Sr$_2$CoIrO$_6$ phases could be related to the presence of Ir$^{6+}$ cations. We note that Ir$^{6+}$O$_6$ bonds are stronger and, therefore, more covalent than Ir$^{5+}$O$_6$ bonds. In addition, it has been observed for metals, monoxides and several families of perovskites that, as the oxidation state of the metal centers increases, the adsorption energies of O$_2^*$, *OH and *OOH become more positive, that is, weaker.$^{37,44}$ Hence, we conclude that more covalent Ir–O lattice bonds are concomitant with weaker Ir–O bonds. It is worth noting here that in a Sabatier-type analysis, the weakening of metal–adsorbate bonds is generally beneficial for the OER when the compounds are initially located on the strong-binding (left) side of a volcano activity plot and until a limit, namely, the top of the volcano. Beyond such limit, the bonds start being undesirably weak, as discussed in the next section.

The specific OER activity of the perovskites was assessed by normalizing the OER activity by the compounds’ active surface area (Fig. 1c). The surface area can be calculated by several methods such as BET or electrochemical methods (ECSA). In the case of mixed oxides, in view of their usually low surface areas, it is a common practice to normalize the activity by the mass specific surface area ($A_S$) calculated from the diameter data of TEM particles (Section S9, ESI†). The determination of $A_S$ (eqn (S5) in the ESI†) along with a comparison between $A_S$, BET and ECSA values is provided in Section S9 in the ESI (Table S3†). The $A_S$ values obtained for the perovskites with M = Ni, Co, Sc and Fe are 2.9, 1.3, 6.8 and 1.6 m$^2$ g$^{-1}$, respectively. The different $A_S$ values obtained are in line with the different particle sizes measured, which are 276 ± 29 nm for Ni catalyst.
519 ± 152 nm for Co, 111 ± 33 nm for Sc and 376 ± 110 nm for Fe catalyst.

As observed in Fig. 1c, the specific activities follow a similar trend compared to the current densities, being Sr₂NiIrO₆ and Sr₂CoIrO₆ again the most active OER catalysts. Sr₂CoIrO₆ achieves a specific activity of 7.5 mA cm⁻² at 1.55 V, which is larger than the one reported by Grimaud et al.²⁶ of 4–5 mA cm⁻² at the same potential. Another important observation is that the specific activities of Sr₂CoIrO₆ and Sr₂FeIrO₆ increase compared to the other catalysts because their surface area is lower. This means that the low current density reached by Sr₂FeIrO₆ stems in part from its low surface area.

We studied the effect that the IrO₆ Oh distortion could have on the OER performance. The Oh distortion (Δ) in perovskites is calculated with eqn (11):⁵⁷

\[
\Delta = \frac{1}{6} \sum \left( \frac{b_i - b_{av}}{b_{av}} \right)^2
\]

(11)

where \( b_i \) and \( b_{av} \) are the individual Ir–O distances (taken from ref. 33 and 39) and \( b_{av} \) are the average (Ir–O) distances (Table S2†), see Fig. S5 in the ESI.† The inset of Fig. 1c displays the evolution of the potential with the IrO₆ Oh distortion at a specific activity of 0.5 mA cm⁻². We observe that a higher distortion inside the Oh results in higher OER activities. However, Δ gives information about the distortion inside the Oh and not about the tilting of the Oh. Both types of distortions are detailed in Fig. S5 in the ESI.† Concerning the effect of the Oh tilting in the catalytic activity, Fig. S6 in the ESI† illustrates the evolution of the activity on the perovskites under study with the monoclinic β angle and with the Oh tilting itself. Smaller monoclinic β angles indicate that the perovskite is more distorted, i.e., more deviant from the cubic structure. The results indicate that larger Oh tilting also has a positive effect on the OER activity of the perovskites.

In conclusion, we observe that both distortion and Oh tilting in the catalytic activity of the Sr₂MIrO₆ double perovskites (M = Ni, Co, Sc, Fe) is ca. 38 wt%, which is similar to the content of other Ir mixed oxides and considerably lower than SrIrO₃ (59 wt%) and IrO₂ (see Table 1). Regarding the mass activities (in A g⁻¹) of the catalysts (Fig. 1d), we note that the iridium mass fraction of all Sr₂MIrO₆ is similar among the analyzed double perovskites (see Table 1), so the trend is comparable to that of the current densities in Fig. 1a. As shown in Table 1, the mass activities obtained for Sr₂NiIrO₆, Sr₂CoIrO₆, and Sr₂ScIrO₆ at 1.525 V are comparable to state-of-the-art Ir mixed oxides and largely surpass IrO₂.

### 3.3 Computational modelling of Sr₂M₀IrO₆ perovskites

To further rationalize the trends in OER catalytic activities of the Sr₂M₀IrO₆ double perovskites (M = Ni, Co, Sc, Fe), we resorted to DFT calculations. First, we assessed the bulk configuration of such perovskites. All calculations were performed on antiferromagnetic systems, in line with the experimental antiferromagnetic ordering on these perovskites.⁵³,³⁹ Among the different antiferromagnetic spin arrangements, we found that the configuration in which the spins are aligned in the (001) plane was most stable for all the analyzed systems.

Furthermore, we calculated the OER overpotentials following the procedure described in the Computational Details section. Values of 0.40, 0.44, 0.75, and 0.79 V were obtained for M = Co, Ni, Fe, and Sc, respectively. Because of the inherent accuracy of DFT (given by the gray bands in Fig. 2a) and the proximity of some of the overpotentials calculated here, we safely conclude that DFT predicts that Sr₂M₀IrO₆ with Co and Ni are more active than those with Fe and Sc, in agreement with experiments. However, it is difficult to ascertain if double perovskites with Co are more active than those with Ni. We note as well that depending on the activity metrics, the ordering between M = Co and Ni changes, see Fig. 1a and c. These results are in line with the experimental observations in Fig. 1b. We rationalize the trends in two different ways. Firstly, in Fig. 2a, the overpotentials of the studied perovskites appear as a function of the descriptor \( \Delta G_{OH} - \Delta G_{OH} \) which renders a volcano-type activity plot for the OER.⁶¹,⁴⁴ For the sake of comparison, we also

![Fig. 2](image-url)
included computational\textsuperscript{44} and experimental\textsuperscript{46} literature data for IrO\textsubscript{2}, together with Sr\textsubscript{2}CoIrO\textsubscript{6} and Sr\textsubscript{2}NiIrO\textsubscript{6} from our own experimental data (the addition of experimental data into the computational volcano plot is made following the method described elsewhere\textsuperscript{47,48}). Fig. 2a shows that the double perovskites follow the traditional adsorption-energy scaling relations, and those with M = Co, Ni are the closest to the top of the volcano, even surpassing IrO\textsubscript{2}(110). Since most of the compounds in Fig. 2a are on the left (strong-binding) side of the volcano, weakening the adsorption energy of *O likely enhances the activity.

Secondly, from a thermodynamic standpoint the ultimate goal in OER electrocatalysis is to reach thermoneutrality,\textsuperscript{46,69,70} which is achieved by a hypothetical “ideal” catalyst for a multi-electron reaction. The OER reaction steps of such catalyst are fully symmetric, that is, with ΔG\textsubscript{i} = 1.23 eV (eqn (5)-(8)), which implies that \( \eta_{\text{OER}} = 0 \) V. To quantify how much a real catalyst approaches the ideal catalyst, one can use the electrochemical-step symmetry index (ESSI, see the Computational Details section).\textsuperscript{46} This metric for electrocatalytic symmetry is defined so that the OER overpotential is reduced as the symmetry of catalysts increases, namely when ESSI approaches zero. The studied Sr\textsubscript{2}MnIrO\textsubscript{6} compounds comply with this general guideline in Fig. 2b, where we observe a good correspondence between ESSI and \( \eta_{\text{OER}} \). The OER at Ir sites seems rather asymmetric when Sc and Fe are in the M position of Sr\textsubscript{2}MnIrO\textsubscript{6}, whereas Co and Ni help increase the electrocatalytic symmetry, thereby lowering the OER overpotential.

An interesting feature of ESSI is its ability to provide guidelines for the enhancement of electrocatalysts.\textsuperscript{47} This can be made following adsorption-energy scaling relations between *O, *OH and *OOH, or breaking them, respectively referred to as δ and ε optimization. The triangles in Fig. 2a and b show that the scaling-based δ-optimization of double perovskites with Ni would sizably reduce \( \eta_{\text{OER}} \) from 0.44 to 0.28 V if *OH binding energies decrease by 0.16 eV. Double perovskites with Co would only experience a small reduction from 0.40 to 0.35 V when *OH adsorption be strengthened by 0.05 eV. On the other hand, scaling-free ε-optimization would only benefit Sr\textsubscript{2}NiIrO\textsubscript{6}, for which a strengthening of 0.27 eV in *OH adsorption with respect to *OH would lower the overpotential to 0.17 V. All this implies that Sr\textsubscript{2}NiIrO\textsubscript{6} is more prone to further optimization, which is interesting for future studies.

3.4 Studying the initial OER activity of Ir double perovskites

Both the experimental and theoretical results presented in this work indicate that Sr\textsubscript{2}CoIrO\textsubscript{6} and Sr\textsubscript{2}NiIrO\textsubscript{6} are the most active catalysts among the studied Ir double perovskites. To verify that the initial OER activities reported in Fig. 1 can be ascribed to the perovskite phase and not to IrO\textsubscript{2}/IrOOH surface segregated phases, we characterized the most active perovskites (Sr\textsubscript{2}CoIrO\textsubscript{6} and Sr\textsubscript{2}NiIrO\textsubscript{6}) after 5 cycles in OER by TEM/EDX and XPS. We specifically chose 5 cycles because no significant changes on the current densities are observed after those. The HRTEM images shown in Fig. 3a, which correspond to the catalysts recovered after 5 cycles, indicate that the materials still display the perovskite structure with only a minor fraction of Ir particles observed at the surface.

We studied the surface of Sr\textsubscript{2}CoIrO\textsubscript{6} and Sr\textsubscript{2}FeIrO\textsubscript{6} perovskites by XPS before and after 5 cycles of reaction to compare the surface stability of a highly active catalyst and one with considerably lower activity. Both samples present Sr, Ir and M cations with Ir/M surface atomic ratios close to the initial nominal value of 1. However, the content of Sr is significantly smaller after 5 cycles than that on the initial samples (Fig. 3b),

![Image](https://example.com/image.png)
reflecting the high tendency of Sr to dissolve in the electrolyte. In the case of Sr₂CoIrO₆, the surface content of Ir increased compared to the content of M (Fig. 3b), indicating the incipient formation of Ir-based nanoparticles.

We also analyzed the evolution of the XPS-Ir 4f region (Fig. 3c). According to previous reports dealing with the stability of Ir perovskites for the OER in acid electrolyte,⁵² if the used perovskites were covered by IrOₓ/IrOOH nanoparticles, the oxidation state of Ir would change to +4/+3 after the reaction. However, as observed in Fig. 3c, the Ir 4f core-level region is not shifted to smaller binding energies (as expected for a reduction in Ir’s oxidation state) after 5 OER cycles. Instead, it remains at the same binding energy for Sr₂FeIrO₆, or shifts to slightly higher binding energies for Sr₂CoIrO₆. In line with a previous report,⁵³ we suggest that the oxidation from Ir⁵+ to Ir⁶+ is responsible for the high OER activity Sr₂CoIrO₆. This partial surface oxidation of Ir during the OER has been observed and correlated with an activity enhancement, owed to the existence of Ir in a higher oxidation state or a different molecular surface environment, caused by the oxidizing conditions of the OER.⁵⁴,⁵⁵ On the other hand, the oxidation of Ir⁵+ to Ir⁶+ is not observed in Sr₂FeIrO₆, which exhibits lower OER activity but higher stability (see below). These observations clearly indicate that the oxidation state of Ir plays a crucial role in the OER activity, which seems to be enhanced by Ir cations with upper oxidation states.

3.5 Stability and evolution of Sr₂MIrO₆ during the OER

The instability of mixed oxides during the OER in acidic media is an issue yet to be resolved. It has been reported that mixed oxides lose stability after several OER cycles, sometimes immediately after immersion in the electrolyte, prior to the OER cycling experiments.⁵⁶,⁵⁷,⁵⁸ This lack of stability leads to catalysts that are more active than the fresh mixed oxide, likely due to the deposition of highly active IrOₓ/IrOOH nanoparticles at the surface of the mixed oxides. As shown in Fig. 3, we tested the catalysts after 5 cycles of reaction and, although the perovskite was still the main phase, some signs of degradation were observed. To further evaluate the stability of Sr₂MIrO₆, we measured 50 OER cycles at 10 mV s⁻¹ between 1.2 and 1.7 V (Fig. 4a), in all cases showing an increment of the OER activity with cycling. To understand the origin of the observed behavior, we analyzed the structure and composition of the used catalysts after such 50 cycles.

Fig. 4b shows representative HRTEM images of the perovskites after 50 OER cycles. The structure of Sr₂CoIrO₆ and Sr₂NiIrO₆ collapses, as observed in the HRTEM and in the FFT in the inset. The resulting material is formed by nanometric Ir-based particles with no signal indicative of the presence of Co, Ni or Sr in the samples after 50 cycles (EDX results, Fig. S8 in the ESI†). However, the morphology of both catalysts after 50 cycles is different (Fig. S9 in the ESI†). The Ni-containing catalyst collapses into a more open structure, less dense and apparently with more surface area (Fig. S9a†), while the Co-containing catalyst presents a more compact morphology (Fig. S9b†).

On the other hand, Sr₂FeIrO₆ and Sr₂ScIrO₆ maintain the perovskite structure as suggested by the presence of the [101] perovskite planes in the Sc perovskite and the oriented structure down the [111] zone axis in the case of the Fe-containing double perovskite. The presence of Sr, Fe(Sc) and Ir in the particles can be observed by EDX analyses, see the map in Fig. 4c. However, a careful analysis of the used perovskites also reveals the presence of IrOₓ nanoparticles at the surface of the latter perovskites (Fig. 4b), indicating the reconstruction of the perovskites during the OER. These results suggest that, contrary to the activity trend, the more distorted perovskites (M = Co and Ni) are less stable than the more symmetric ones (M = Sc and Fe).

Insights onto the deactivation of representative Sr₂CoIrO₆ and Sr₂FeIrO₆ samples were obtained by analysing the composition of the electrolyte by ICP-OES. We collected electrolyte before the reaction and after 5 and 50 OER cycles. The evolution of Sr, M (Co and Fe) and Ir concentrations in the electrolyte during the OER with both catalysts is shown in Fig. S10 in the ESI†. As observed, Sr and M dissolve faster in Sr₂CoIrO₆ than in Sr₂FeIrO₆ with cycling, indicating that Sr₂FeIrO₆ is more stable for the OER in acid electrolyte. In both cases the dissolution of Ir is low, in agreement with the formation of IrOₓ particles at the surface that are not dissolved during the reaction.

These observations are in good agreement with previous studies,⁵⁶,⁷¹ reporting that more symmetric perovskites are more stable under the conditions of the H₂O/O₂ interconversion reactions, since they can be subjected to more extreme potentials without being destroyed. Conversely, distorted structures are susceptible to dissolution upon OER cycling.

We also performed chronoamperometry studies during one hour at the fixed potential where the current density is 10 mA cm⁻² (Fig. 5a). The double perovskites’ activity is stable during such steady-state durability test, losing only 5–10% of the initial
activity in all cases. This result indicates the easier formation of Ir-nanoparticles on the surface during cycling experiments compared to experiments performed at a selected current density (i.e. 10 mA cm⁻²).

In view of the high OER activity of Sr₂NiIrO₆ and Sr₂CoIrO₆, we performed a more aggressive dynamic durability test consisting of 10 000 OER cycles at 50 mV s⁻¹ between 1.2 and 1.7 V (Fig. 5b) on both catalysts. This stability test is essentially an experiment in which the catalysts are continuously cycled between those two potentials in acid media for ~55 h. In the case of Sr₂NiIrO₆, the activity loss after 10 000 cycles is small, with an increment of potential of only 10 mV to achieve the reference current density of 10 mA cm⁻². For Sr₂CoIrO₆, the activity decreases from 13.3 to 9.8 mA cm⁻² at 1.6 V, indicating severe deactivation after the durability test. The activity loss on Sr₂CoIrO₆ can also be followed from the evolution of the potential needed to achieve a current density of 10 mA cm⁻², which increases from 1.57 to 1.60 V. These results may indicate that the surface reconstruction is different in both oxides. The effect of different surface reconstruction on the OER durability has been extensively reported.\textsuperscript{20,22,24-72} Analyzing the reconstruction of the surface after 50 cycles for both oxides (Fig. S9 in the ESI†), we conclude that the Ni-containing catalyst presents less dense Ir phases than the Co-containing catalyst. This different morphology can explain the different evolution of the catalyst after the long-term durability test. Probably, during cycling, the Co-containing catalyst forms larger Ir particles more rapidly than the Ni-containing catalyst, decreasing its current density after 10 000 cycles.

4. Conclusions

In this study we report the performance of Sr₂MIrO₆ (M = Ni, Co, Sc, Fe) double perovskites for the OER in acid electrolyte and rationalize the trends using a variety of experimental and computational descriptors, such as Ir oxidation state, degree of octahedral distortion, strength of Ir-adsorbate bonds, and electrocatalytic symmetry. The OER activities are comparable to those of state-of-the-art Ir-based catalysts, following the trend Sr₂FeIrO₆ < Sr₂ScIrO₆ < Sr₂CoIrO₆ < Sr₂NiIrO₆. The higher OER activity of Sr₂NiIrO₆ and Sr₂CoIrO₆ stems from their ability to stabilize Ir⁶⁺ cations during the OER. However, the stability of both oxides is lower than those of Sr₂ScIrO₆ and Sr₂FeIrO₆ in which lower Ir oxidation states are found. Thus, low Ir oxidation states, relatively strong binding energies and high structural symmetry ensure high stability.

While all of the analyzed materials exhibit the perovskite structure during the first cycles of reaction, only the more symmetric phases, which are the lesser active, maintain the perovskite structure for more than 50 OER reaction cycles.

The correlations between iridium oxidation state, adsorption energies and OER activity in acid media reported in this work can serve as a guideline for designing more active and durable OER catalysts in acid electrolyte. Our results hint toward a necessary tradeoff between activity and stability, which can be provided by a careful selection of the metal M in Sr₂MIrO₆ double perovskites.

Conflicts of interest

There are no conflicts to declare.

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