Mixed metal-antimony oxide nanocomposites: low pH water oxidation electrocatalysts with outstanding durability at ambient and elevated temperatures

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ABSTRACT: Electrochemical water splitting with a proton-exchange membrane electrolyte provides many advantages for the energy-efficient production of high-purity H₂ in a sustainable manner, but the current technology relies on high loadings of expensive and scarce iridium at the anodes, which are also often unstable in operation. To address this, the present work scrutinises the electrocatalytic properties of a range of mixed antimony-metal (Co, Mn, Ni, Fe, Ru) oxides synthesised as thin films by a simple solution-based method for the oxygen evolution reaction in aqueous 0.5 M H₂SO₄. Among the noble-metal free catalysts, cobalt-antimony and manganese-antimony oxides demonstrate good stability over 24 h and reasonable activity at 24 ± 2 °C, but slowly lose their initial activity at elevated temperatures. The ruthenium-antimony system is highly active, requiring an overpotential of only 0.39 \pm 0.03 and 0.34 \pm 0.01 V to achieve 10 mA cm⁻² at 24 \pm 2 and 80 °C, respectively, and most importantly, remaining remarkably stable during one-week tests at 80 °C. Detailed characterisation reveals that the enhanced stability of metal-antimony oxides water oxidation catalysts can arise from two distinct structural scenarios: either formation of a new antimonate phase, or nanoscale intermixing of metal and antimony oxide crystallites. Density functional theory analysis further indicates that the stability in operation is supported by the enhanced hybridisation of the oxygen p- and metal d-orbitals induced by the presence of Sb.

INTRODUCTION

Water electrolysers based on a proton-exchange membrane (PEM) electrolyte are currently seen as the preferred technology for the production of green hydrogen from renewables.¹⁻² Double digit megawatt PEM plants are already available and even larger installations are planned.³ Recent breakthroughs in the design of bipolar plates and cathode catalysts for the PEM electrolysers now throw a spotlight on the membrane and anode electrocatalysts as the components requiring further significant cost-efficiency improvements.⁴ Catalysts for the oxygen evolution reaction (OER) are particularly problematic in essentially all key aspects - price, availability, activity and stability. While iridium oxides provide perhaps the best combination of the activity and stability among known monometallic catalysts,¹ there is not enough of this exceptionally rare metal currently available to us to bring the PEM water electrolysis to the TW scale.⁵ Moreover, both theoretical and extensive experimental studies reveal the unavoidable degradation of iridium-based OER catalysts even under ambient conditions, ⁶⁻¹⁰ and especially in industrially relevant high-temperature tests, ¹¹ which is among the reasons for the comparatively high loadings of Ir in the anodes of PEM electrolysers.

Alternative OER catalysts based on more abundant and cheaper elements exist, in particular lead(IV)-based oxides developed through decades of research on the metal electrowinning anodes,¹² though their specific activity is significantly lower than that of iridium systems.¹³ Among other non-noble metal options, the obvious candidates are oxides of manganese, cobalt, nickel and iron, which have been widely investigated as OER catalysts for the alkaline and near neutral conditions.¹⁴⁻²¹ Monometallic oxides of Mn and Co have also been examined for applications in acidic environment, and in contrast to Ni and Fe which immediately dissolve,²² relatively stable performance for several hours was demonstrated.²³⁻²⁴ However, eventual degradation and essentially complete loss of activity is unavoidable. A noble alternative to iridium is also known – ruthenium oxides are reported to be at least as active OER catalysts.²² While the **amount of Ru in the Earth's crust is only slightly higher than that of** Ir^{25} and it continues to rise in price, ruthenium is still more than fivefold cheaper, 26 is easier to refine and is produced on an order of magnitude higher scale. 27 However, rapid degradation of RuO_2 anodes is again a major issue. $^{28\cdot30}$

Overall, it is highly unlikely that any monometallic oxide can provide an optimal combination of characteristics required to be a high-performance OER catalyst. In contrast, new materials with improved activity and/or stability emerge from the exploration of multielement oxide systems that sometimes combine the properties of individual compounds or exhibit distinct properties of their own. In the context of the design of catalysts for the low pH OER, successful approaches often emerge from combining a catalytically active oxide with the one that is highly stable under operating conditions to enable improved long-term operation with decreased losses in activity. This strategy has been broadly adopted for many years in the research on the electrowinning anode catalysts using PbO₂ as a "matrix" that stabilises oxides of cobalt, manganese, silver and other metals.^{12, 31-32} The same approach is now also applied in the design of anode catalysts for PEM water electrolysers. Improved electrochemical activity and durability during the OER in acidic solutions has been reported when catalytically active metals have been combined with the oxides of Ta^{IV 33-34} Sn^{IV 34-35} Ti^{IV 36-37} Pb^{IV 38-40} Y^{III 41} Cr^{IV 42} and also Sb^{V 43-44}

Antimony oxides (SbO_x) are of a particular interest in this context. The thermodynamic stability of SbO_x under the low pH OER conditions,⁴⁵ electrical conductivity,⁴⁶⁻⁴⁸ along with abundance and availability of Sb⁴⁹ render these compounds a suitable structural component for acid-stable water oxidation electrocatalysts. Antimony oxides have been used as stabilising interlayers and components in robust electrowinning anodes, 50-51 and have recently been also introduced to studies of OER catalysts for acidic water electrolysers. In particular, the research teams of Gregoire^{44, 52} and Lewis^{43, 53} both reported promising activity and improved stability in operation of antimonates of manganese and nickel. More recent work also described cobalt antimonate OER catalysts.⁵⁴ Thus, the stability of transition metal oxides during oxidation of acidic water can be improved via combination with the highly promising SbO_x matrix. In the case of Ru, this can also reduce the noble metal loading and thereby the cost. However, the catalysts of this type are currently underexplored and have only recently gained an increased attention.

In the search for highly active, and genuinely stable catalysts under practical operating conditions, the core aim of the present work is the investigation of the electrocatalytic activity and stability of mixed metal-antimony oxides towards the oxidation of acidic water. Specific emphasis is put on the stability, which is rarely assessed rigorously in the current literature, viz. the experiments are commonly limited to several hours and ambient temperature conditions only. Moreover, even such mild conditions cause continuous degradation of many catalysts, which sometimes remains underestimated when the stability data are recorded and presented in the galvanostatic mode. Herein, the initial tests were also undertaken under ambient conditions to identify the most promising catalysts, which were further investigated at elevated temperatures to assess and demonstrate the genuine suitability of antimony-metal oxides for operation under the conditions relevant to the PEM electrolysers. In-depth electrochemical and physical characterisation of the catalysts before and after exhaustive electrocatalytic tests coupled to the density functional theory (DFT) calculations reveal new insights into the origins of the stability of metal-antimony oxide OER catalysts on the structural and atomistic levels.

RESULTS AND DISCUSSION

All materials in this work were synthesised via annealing of precursor salts deposited from their solutions onto nominally flat glass supports covered with a thin conductive layer of F-doped SnO₂ (FTO). While sputtering and similar advanced techniques provide a better control over the morphology and composition, our choice over the much simpler drop-casting/annealing fabrication protocol was motivated by the ease of the future optimisation of this approach for the creation of high-surface area catalysts of applied significance. All electrochemical tests were undertaken in 0.5 M H₂SO₄ with measured pH of 0.3 (at ambient temperature). All galvanostatic data are presented below after manual post-correction for ohmic losses using the uncompensated (R_{u}) values measured by the electrochemical impedance spectroscopy. However, the correction was generally not applied to voltammograms, where currents cannot be always confidently attributed to the stationary catalytic process only. Extended details on the methods and procedures used in this work are provided in Supporting Information.

Electrocatalytic activity under ambient conditions

Individual metal oxides (MO_x) considered herein, *viz*. RuO_x, CoO_x, MnO_x, NiO_x and FeO_x, as well as SbO_x, were not expected to be highperformance OER catalysts of independent interest, but their properties were briefly examined to understand relevant differences in the key performance parameters that are likely to affect those of the corresponding metal-antimony oxides. X-ray diffraction (XRD) and scanning electron microscopic (SEM) analysis of the as-prepared MO_x and SbO_x indicate that the materials were dominated by single oxide phases RuO₂, Co₃O₄, Mn₂O₃, NiO, Fe₂O₃, and Sb₂O₄ (Figure S1) of variable morphology (Figure S2). Tests of MO_x and SbO_x as OER catalysts confirmed their either limited stability (RuO₂, Co₃O₄, NiO and Fe₂O₃) or activity (Mn₂O₃ and Sb₂O₄) (Figures S3-S5 and attached comments).

Further studies focused on mixed metal-antimony oxides, which are referred to as $[M+Sb]O_y$. These catalysts were synthesised with a 1:1 molar metal-antimony precursor ratio, unless stated otherwise, but the actual composition during the OER is different due to the una-voidable corrosion in acidic solutions. The degree of this corrosion for the selected key materials was quantified (Table 1) and is discussed later in the text.

Table 1. Relative loss of metals and antimony from cataly	sts
(at.%) ^a during the OER in stirred 0.5 M H ₂ SO ₄ .	

Catalyst	24 ± 2 °C ^b		° 3° 08	
	Metal	Sb	Metal	Sb
[Mn+Sb]O _y	21	17	Not analysed	
[Co+Sb]O _y	15	26		
[Ru+Sb]O _y	1	37	3	47

^a Calculation based on the amount of metal and Sb deposited onto electrodes (1 μ mol cm⁻² each) and the amounts measured in solutions after the OER tests by ICP-OES.^b 10 mA cm⁻² for 24 h and then 2.03 and 1.93 V vs. RHE for 0.5 h at each potential. ^c 10 mA cm⁻² for 192 h and then 2.03 and 1.93 V vs. RHE for 0.5 h at each potential.

First, we exclude Ni- and Fe-based systems from detailed analysis, as these materials rapidly degraded under the OER conditions (Figure S6 and S7). Attempts to improve their performance through variations in the metal : antimony ratios and annealing temperature were not successful. The effect of the latter parameter was also briefly considered for the much better performing Co-, Mn- and Ru-Sb oxide catalysts. The best results were always obtained at 600 °C (Figure S8), which is the highest value we could use due to thermal instability of FTO.⁵⁵ Hence, all results discussed here in the main text of the paper were obtained with the catalysts synthesised at 600 °C.

Voltammetric assessment of the initial OER catalytic activity of CoOx and [Co+Sb]Oy revealed lower performance of the latter (Figure S3a and 1a), which is likely associated with the reduced amount of the active cobalt oxide surface species, as also measured by cyclic voltammetry (cf. Figures S5a and S9a). Subsequently recorded chronopotentiograms reveal the significant initial degradation of the cobalt-antimony-modified electrode, though more importantly [Co+Sb]O_V catalysts do not completely lose their activity and sustain a current density of 10 mA cm⁻² at a reasonable overpotential (Figure 1b). The duration of the initial drop increased with an increase in the Co: Sb ratio used for the synthesis, while the best stabilised activity was demonstrated by the catalysts prepared with equimolar amounts of Co and Sb (Figure S10). This type of $[Co+Sb]O_{\gamma}$ demonstrated stable water oxidation at 10 mA cm⁻² at a well-reproducible IR_{u} -corrected overpotential of 0.769 ± 0.010 V (Figures 1b and S11). Subsequent potentiostatic tests at 2.03 and 1.93 V vs. RHE also did not reveal further significant losses in the catalytic activity (Figure 1c). A plausible explanation of the rapid initial loss of the performance is provided in the following section, while at this stage we conclude that $[Co+Sb]O_y$ exhibits a reasonable short-term stability during the OER at low pH and ambient temperature, yet its catalytic activity is not high.

In contrast to the cobalt-based materials, combination of Mn with Sb produced a very significant improvement in the OER catalytic activity with respect to monometallic MnO_x (Figure S3b and 1a). The [M+Sb]O_y catalysts were also able to maintain the initial performance during 24 h galvanostatic tests (Figure 1b). Variations in the Mn to Sb precursor ratio revealed that the manganese-rich materials (Mn : Sb = 2 : 1) suffer slow deterioration in performance (Figure S12), which was also recently reported for Mn antimonate with a similar starting composition and under comparable conditions.⁵² Contrasting this behaviour, the activity of [Mn+Sb]Oy synthesised herein with 1: 1 and 1: 2 ratios was highly stable. Between these two types of materials, the Sb-rich one was found to be less active, exhibiting $\eta_{\rm IR}$ for 10 mA cm⁻² of *ca* 0.70 V. The best-performing [Mn+Sb]O_y electrocatalysts with the initial equimolar Mn : Sb ratio stably maintain the OER current density of 10 mA cm⁻² for at least 24 hours under ambient conditions at a highly reproducible overpotential of 0.677 \pm 0.008 V (Figures 1b and S13).

Combination of ruthenium with the antimonate matrix produced catalysts that exhibit a stable cyclic voltammetric response typical of a robust and highly active OER catalyst up to 2.03 V *vs*. RHE, in contrast to unstable RuO_x (Figures 1a, S3e and S6e). Moreover, galvanostatic tests at 10 mA cm⁻² improved the performance of the [Ru+Sb]O_y materials by *ca* 0.04 V over the initial *ca* 10-12 h of experiments, eventuating in a reproducible stabilised *IR*_u-corrected overpotential of $\eta_{\rm IR} = 0.39 \pm 0.03$ V (Figures 1b and S14).

Overall, the analysis of the initial catalytic performance and 25 h stability at ambient temperature of the metal-antimony oxides revealed a synergistic effect of the combination of a catalytically active metal and acid-stable Sb for the $[Mn+Sb]O_y$ system, where a significant improvement in the activity was achieved. The key advantage of combining Co and Ru with the SbO_x matrix is in the substantial improvements in the stability, which is a highly favourable outcome given that the instability of the OER anode catalysts is among the most technologically pressing problems of the PEM water electrolysers. 56

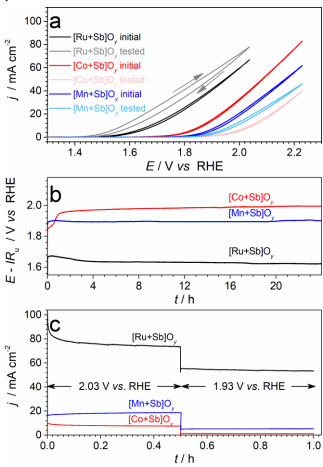


Figure 1. Electrochemical characterisation of $[Co+Sb]O_y$ (*red*), $[Mn+Sb]O_y$ (*blue*) and $[Ru+Sb]O_y$ (*black*) in stirred 0.5 M H₂SO₄ at 24 ± 2 °C: (a) cyclic voltammetry (scan rate 0.020 V s⁻¹; third scans; no $/R_u$ correction applied) recorded before (vivid traces) and after (pale traces) tests shown in panels b and c (arrows exemplify the direction of the voltammetric sweeps, which was qualitatively the same for all catalysts); (b) $/R_u$ -corrected chronopotentiograms (current density 10 mA cm²_{geom}); and (c) subsequently recorded chronoamperograms (at a non- $/R_u$ -corrected potentials of 2.03 and 1.93 V vs. RHE).

Characterisation of the mixed metal-oxide catalysts

To enable deeper understanding of the observed trends in the electrocatalytic performance of the investigated metal-antimony oxides, physical characterisation of the key materials was undertaken using the state-of-the-art techniques.

Corrosion during operation. SEM images of the $[M+Sb]O_y$ electrodes taken before and after the OER tests revealed that the initially flat, essentially featureless catalyst coatings underwent partial erosion due to the loss of material during the OER (Figure S15). The level of this loss was quantified for the key catalysts by ICP-OES (Table 1), while the changes in the catalyst surface compositions were estimated by X-ray photoelectron spectroscopy (XPS).

After 24 h galvanostatic followed by 1 h potentiostatic tests at ambient temperature, the levels of corrosion of metals and antimony from $[Co+Sb]O_y$ and $[Mn+Sb]O_y$ into the electrolyte solutions were comparable (Table 1), notwithstanding a very distinct electrocatalytic behaviour (Figure 1). XPS showed that concentrations of the metals at the surface of tested $[Co+Sb]O_y$ and $[Mn+Sb]O_y$ were also similar (Table 2). At the same time, voltammetric analysis, which provides the only reliable means of probing the catalytically active species, indicated that $[Mn+Sb]O_y$ still exhibit detectable peaks associated with Mn redox transformations after the OER (Figure S8b), while operated $[Co+Sb]O_y$ present a featureless response (Figure S8a). These observations might reflect the differences in the quasi-stabilised concentrations of the catalytically active metals in the top-most layers of the Co-Sb and Mn-Sb oxide systems. While the $[Co+Sb]O_y$ surface loses a very significant part of its cobalt, $[Mn+Sb]O_y$ is likely to maintain a higher amount of manganese available for the OER catalysis.

Table 2. Relative surface concentrations of metals for catalysts a before and after OER tests in stirred 0.5 M H_2SO_4 .

Catalyst	As prepared	Tested at		Tested at	
		24 ± 2 °C ^b	°C د		
$[Mn+Sb]O_y$	23 ± 1	12 ± 2	n.a.		
[Co+Sb]Oy	34 ± 2	9 ± 4	n.a.		
[Ru+Sb]O _y	9 ± 5	n.a.	44 ± 5		

^a at.% with respect to total metal + antimony amount quantified by XPS; data are presented as a mean \pm one standard deviation for several measurements. ^b 10 mA cm⁻² for 24 h and 2.03 and 1.93 V *vs.* RHE for 0.5 h at each potential. ^c 10 mA cm⁻² for 10 h.

The lowest level of metal corrosion was observed for $[Ru+Sb]O_y$ where the amount of lost ruthenium was not more than 1 at.% at ambient temperature, but the amount of antimony released into the electrolyte solution was significant (Table 1). This was accompanied by a slight increase in the pre-catalytic voltammetric currents (Figure S9e). Most critically, long-term tests of $[Ru+Sb]O_y$ at elevated temperature of 80 °C (*vide infra*) caused only slightly higher level of corrosion (Table 1), while XPS consistently demonstrated that the surface Ru : Sb ratio increased up to *ca* 1 : 1 (Table 2). Thus, $[Ru+Sb]O_y$ suffers a favourable corrosion of antimony that improves the electrocatalytic activity of the material towards the OER (Figures 1b and 2a) through an enrichment of the surface layer in catalytically active ruthenium.

When considered together, the observed levels of corrosion (Tables 1 and 2) and stable electrocatalytic performance of the manganese-, cobalt- and ruthenium-antimony oxides (except for the initial drop in the [Co+Sb]O_y performance) (Figure 1) suggest that a quasi-equilibrium between the solid oxides and dissolved forms of Mn/Co/Ru and Sb is established in the system. Such an equilibrium between dissolution/redeposition of metal oxides is likely to sustain the observed stable electrocatalytic operation. In other words, the examined OER catalysts most probably operate in a self-healing mode,^{38-39,57-59} with the Sb oxide matrix acting to facilitate the redeposition and suppress the dissolution of the active component.

Structural features. The structural features and the oxidation states of metals and antimony in the key $[Co+Sb]O_y$, $[Mn+Sb]O_y$ and $[Ru+Sb]O_y$ samples were probed by XAS (Figure 2), XPS (Figures S16-S18) and XRD (Figure 3). The ruthenium-antimony system was additionally investigated by TEM (Figure 4).

Fitting of the Sb 3d + O 1s spectra and comparisons to the literature⁶⁰⁻⁶¹ confirmed that antimony adopts a dominant oxidation state 5+ in all catalysts after the OER tests, as well as in most of the untreated samples (Figure S16). The Sb K-edge XANES recorded for Mn-, Co- and Ru-Sb mixed oxides reproduced well the data reported elsewhere for manganese antimonates,⁵² and are again consistent with the 5+ state (Figure 2a and S19). A slight shift to higher energy was noted in the Sb K-edge after the OER tests (Figures S19). Sb Kedge EXAFS of the investigated metal-antimony oxides are similar to each other and to Sb₂O₅ in that none of the datasets exhibit any notable second coordination sphere peaks in the Fourier Transform (FT) of EXAFS beyond the first coordination sphere apparent distances (*R*) of 2 Å (Figures 2a and S20). This would be consistent with Sb being present as a part of a highly-disordered phase or might be the consequence of how the distances in the structure cancel each other out (Figure S20 and Table S1).

The shape and position of the Mn $2p_{5/2}$ peaks for fresh [Mn+Sb]O_y are similar to those for the oxide/oxyhydroxide Mn³⁺ compounds,⁶² and show only a slight shift towards higher binding energies after tests (Figure S17a). Comparisons of the Mn K-edge XANES data for [Mn+Sb]O_y to that for the MnO, Mn₂O₃ and MnO₂ standards suggest that the oxidation state of the metal in the bulk of the as-prepared material is between 2+ and 3+, while Mn³⁺ becomes a dominant component after the OER tests (Figures 2b and S21).

The Co K-edge XANES of as-prepared $[Co+Sb]O_y$ is consistent with an oxidation state between 2+ and 3+, or a material like Co₃O₄ (Figure 2c). The Co K pre-edge intensity decreases after the OER test, which would be consistent with the loss of the Co₃O₄ phase with a tetrahedral Co²⁺ site (Figure S22a-b), which is also supported by changes in the XANES profile (Figure 2c). The major Co oxidation state on the surface can be also ascribed to 3+ based on the XPS analysis (Figure S17b) and comparisons to the literature.⁶²

Thus, both XPS and XANES indicate that the dominating oxidation states of manganese and cobalt in as-prepared metal-antimony mixed oxides are between 2+ and 3+. Although Mn and Co at the catalytic surfaces likely adopt higher oxidation states during the OER, those cannot be detected by *ex situ* analysis due to their very high oxidative reactivity and immediate conversion into states that are thermodynamically favourable under ambient environment.^{63,65}

X-ray diffractograms of $[Co+Sb]O_y$ and $[Mn+Sb]O_y$ exhibited a set of peaks typical of a tetragonal trirutile phase with major 110, 013 and 123 reflections at *ca* 27, 35 and 53°, respectively (Figure 3a-b). Qualitatively similar XRD patterns were obtained for the mixed nickel-antimony and iron-antimony samples (Figure S23). These data agree with the recent studies, ^{43,52,54} and formally suggest that the metal-antimony phase in the investigated materials is structurally similar to $CoSb_2O_6$. It is also worth noting that the mean crystallite sizes for the individual metal oxides phases detected by XRD were notably higher than those for the antimonate phase (Figure 3a-b), suggesting that the former are present as larger agglomerates as compared to the latter and therefore make significantly lower contribution to the electrocatalytically active surface area.

Well-defined FT EXAFS beyond the first coordination sphere were observed at the metal K-edges for $[Co+Sb]O_y$ and $[Mn+Sb]O_y$ (Figure 2b-c). This indicates that the metals do not simply dope the antimony oxide lattice, as was observed previously for the conceptually similar Co-Fe-Pb oxide OER catalyst,³⁸ and more complicated structural scenarios apply. To describe the metal-antimony phases, EXAFS simulations based on the crystal structure of MnSb₂O₆⁶⁶ and that of CoSb₂O₆⁶⁷ were undertaken.

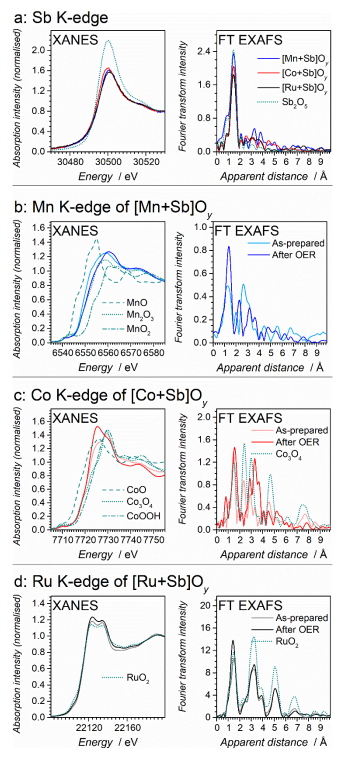


Figure 2. XAS data at (a) Sb, (b) Mn, (c) Co and (d) Ru K-edges obtained for [Mn+Sb]O_y (*blue*), [Co+Sb]O_y (*red*) and [Ru+Sb]O_y (*black*) before (*pale solid traces*) and after (*vivid solid traces*) electrocatalytic tests in stirred 0.5 M H₂SO₄ compared to the defined reference materials (*dashed* and *dotted teal* traces). (e-h) Fourier Transform of EXAFS at (e) Sb, (f) Mn, (g) Co and (h) Ru K-edges. [Mn+Sb]O_y and [Co+Sb]O_y were tested for 24 h at 10 mA cm⁻² and then for 1 h at 2.03 and 1.93 V *vs.* RHE at ambient temperature; [Ru+Sb]O_y was tested for 12 h at 10 mA cm⁻² at 80 °C.

As expected from XRD (Figure 3a), MnSb₂O₆ alone could not describe the Mn K-edge EXAFS for as-prepared [Mn+Sb]O_y (Table S2) due to the presence of a manganese oxide compound, most

likely $Mn_2O_{3,}$ which produces a peak at *ca*2.5-2.6 Å in FT EXAFS (Figures 2b and S21f). After the OER, the intensity of the Mn_2O_3 features in EXAFS are suppressed (Figures 2b and S21c, Table S2), which indicates partial loss of Mn_2O_3 and corroborates the corrosion data (Table 1). Given that this loss does not induce any notable deterioration in the catalytic activity (Figure 1b), we conclude that individual Mn oxides do not make a significant contribution to the OER performance of [Mn+Sb] $O_{y.}$

XAS data recorded at the Co K-edge for $[Co+Sb]O_y$ confirmed the XRD observations of the presence of Co₃O₄ in the as-prepared material, which vanished after the OER tests (Figure 2c and S22c,e). The XAS data of the tested catalyst are well fit with the simulations based on the CoSb₂O₆ structure⁶⁷ (Figure S22c,f and Table S3). On this basis, we interpret the initial rapid drop in the performance of $[Co+Sb]O_y$ (Figure 1b) by the dissolution of the catalytically more active Co₃O₄ phase from the material surface, while the quasi-stabilised performance achieved after *ca* 20 h is ascribed to the true catalytic activity of cobalt antimonate.

The most challenging to interpret set of physical characterisation data among examined systems was obtained for the top-performing [Ru+Sb]O_y catalyst. Analysis of the Ru 3d XP spectra is complicated by a direct overlap with C 1s signals, but it was still possible to conclude that the initial mixed Ru³⁺ + Ru⁴⁺ state of the surface undergoes oxidation during the OER to become dominated by Ru⁴⁺ (Figure S17c and S18c).⁶⁸ In turn, Ru K-edge XANES recorded for fresh and tested [Ru+Sb]O_y were in a perfect agreement with the Ru⁴⁺ state, closely resembling the spectra of RuO₂ (Figure 2d).

Interpretation of the XRD data for [Ru+Sb]O_v was not straightforward as the diffraction pattern of rutile RuO₂ is similar to that of the anticipated trirutile antimonate phase. Nevertheless, the major set of broad diffraction peaks at 28, 35, 40, 54 and 69.5° matches well the positions and relative intensities of the tabulated pattern of ruthenium (IV) oxide (*grey bars* in Figure 3c). This interpretation suggests that the Sb component(s) give rise to a set of broad low-intensity signals at ca25, 30.5 and 48° that can be attributed to highly disordered Sb₂O₅, along with two peaks at *ca* 38.5 and 44° (marked with triangles Figure 3c). One might suggest that these two reflections can be attributed to an antimonate phase which other diffraction peaks are presumably merged with the signals of RuO₂. However, analysis of the highly resolved up to R = 8 Å Ru K-edge EXAFS collected for [Ru+Sb]O_V does not support this hypothesis (Figures 2d and S24). Fitting of these data with simulations based on a RuSb₂O₆type lattice did not produce any satisfactory level of agreement (Figure S24 and Table S4). Clearly resolved FT EXAFS peaks at high R of 3.1, 3.9, 5.1, 6.8 Å present in the catalyst before and after the OER tests are associated with groups of Ru-Ru distances at 5.0, 6.8 and 7.2 Å, and cannot be explained by the RuSb₂O₆ phase. In fact, the EXAFS and XANES of [Ru+Sb]O_v are almost perfectly consistent with the RuO₂ structure, although with a slightly increased level of disorder (Figures 2d and S24, Table S4), which corroborates the major XRD signals (Figure 3c).

Taken together, XAS and XRD suggest that the ruthenium structure in $[Ru+Sb]O_y$ is very close to the RuO₂ lattice, which opens a question on the origin of the significantly improved electrocatalytic performance of the mixed oxide system as opposed to RuO₂ (Figure 1 and S4a). To shed light on this, detailed TEM investigations with elemental mapping were undertaken.

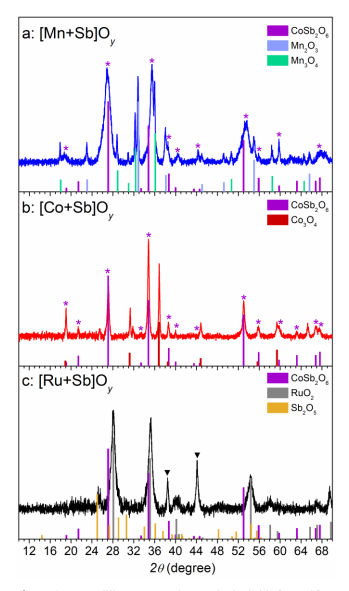


Figure 3. X-ray diffractograms of as-synthesised (a) $[Mn+Sb]O_y$, (b) $[Co+Sb]O_y$ and (c) $[Ru+Sb]O_y$. Vertical lines show tabulated positions and relative intensities for (a-c) $CoSb_2O_6$ ICSD-108964, (a) Mn_2O_3 ICDD-00-041-1442 and Mn_3O_4 ICDD-01-075-1560, (b), Co_3O_4 ICSD-36256, (c) RuO_2 ICSD-731469 and Sb_2O_5 PDF-01-071-0256. Mean crystallite sizes calculated using Scherrer equation were: (a) $d_{RD}(MnSb_2O_6) \approx 7 \text{ nm}, d_{RD}(Mn_2O_3) \approx 35 \text{ nm}, d_{RD}(Mn_3O_4) \approx 41 \text{ nm};$ (b) $d_{RD}(CoSb_2O_6) \approx 24 \text{ nm}; d_{RD}(Co_3O_4) \approx 43 \text{ nm};$ (c) $d_{RD}(RuO_2) \approx 10 \text{ nm}.$ Asterisks indicate diffraction peaks attributed to the antimonate phase. Triangles indicate peaks which assignment is not straightforward.

At low to moderate magnification, STEM-EDS analysis of $[Ru+Sb]O_y$ showed ruthenium and antimony to be intimately mixed, both before and after the OER tests (Figures 4a-b and S25). Conventional TEM imaging demonstrated that the catalyst represents a very fine assembly of nanocrystals, typically less than 10 nm in size, embedded into another material that appears significantly more structurally disordered (Figures 4c-d and S26). Higher magnification images revealed clear lattice fringes of *ca* 3 Å for the crystal-line nanoparticles, which can be attributed to RuO₂ with a relatively high degree of crystalline order (Figure 4c-d).

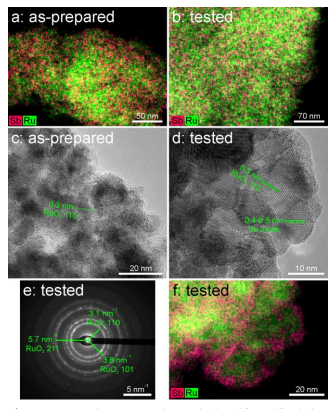


Figure 4. TEM and STEM-EDS characterisation of $[Ru+Sb]O_y$ before and after OER tests in stirred 0.5 M H₂SO₄ at 10 mA cm⁻² for 24 h at 80 °C: (a-b) medium magnification STEM-EDS mapping of the (a) asprepared and (b) tested catalyst; (c-d) TEM micrographs of the (c) asprepared and (d) tested catalyst; (e) selected area electron diffraction taken from a region containing large amounts of both Ru and Sb for the tested catalyst. (f) STEM-EDS mapping analysis of the extended area of the tested catalyst around the region shown in panel d.

This is also illustrated in the selected area electron diffraction pattern, which can be indexed as essentially pure RuO_2 (Figure 4e) notwithstanding it was recorded from a large fragment containing both ruthenium and antimony. The material closely neighbouring RuO_2 nanocrystals exhibited larger lattice fringes of *ca* 4-5 Å (Figure 4d), which cannot be ascribed to ruthenium oxide and therefore are attributed to antimony oxides. STEM-EDS mapping of this area confirmed the partial segregation of ruthenium and antimony at the nanoscale (Figure 4f), but we emphasise that the two elements were still found to be mixed very finely, even when analysed at up to 1.3M× magnification (Figures 4f and S27).

Other antimony rich areas were generally devoid of clear lattice fringes (Figure 4c-d), and although this does not definitely rule out crystallinity, it tends to support a hypothesis that Sb oxides in $[Ru+Sb]O_y$ exhibit a very high level of disorder. We also note that the Sb phase appears to coat the edges and interstices of the RuO₂ nanocrystals in the catalysts after the OER test (Figure 4f), suggesting that antimony has been lost from a previously predominating Sb-based coating of Ru oxide particles rather than a simple aggregation of two phases. This supports the corrosion data (Table 1).

Overall, the physical characterisation data reveal several fundamental differences in the investigated catalysts. As-prepared $[Mn+Sb]O_y$ and $[Co+Sb]O_y$ contain a mixture of individual metal oxides, which majorly dissolve during the OER tests, with a metal antimonate oxide phase. The latter is most likely $CoSb_2O_6$ in $[Co+Sb]O_y$, while the structure of the manganese-antimony oxide exhibits some distinctions from the published features of the antimonate phase and a significant level of disorder. At the same time, no evidence for the formation of a ruthenium antimonate was obtained. Instead, the [Ru+Sb]O_y catalyst contains highly-dispersed RuO₂ crystallites embedded into and decorated with disordered antimony oxides. One might hypothesise that an interaction and intermixing between ruthenium and antimony oxides might still occur, but at the topmost surface layers of RuO₂, especially at the abundant intergrain boundaries visualised by TEM (Figure 4c-d), rather than in the bulk phase. To assess this possibility and further assist in establishing a plausible explanation of the improved electrocatalytic performance of [Ru+Sb]O_y, theoretical analysis of the metal-antimonate systems was undertaken.

Theoretical insights into the improved stability of the Co-Sb and Ru-Sb oxides

Experimental evidence on the stabilisation against corrosion of the oxides of transition metals like Co, Mn, Ni and Ru, through their combination with the oxides that are thermodynamically stable under the OER conditions exists,^{12, 31-44} including the new results reported in the present work. However, there is a lack of the understanding behind this stabilising effect, which we aimed to address through the theoretical analysis of two systems of interest herein – Co-Sb and Ru-Sb mixed oxides. Extending the analysis to the Mn-Sb combination could not be realised due to a well-documented complex ground state magnetic structure of Mn_2O_3 , which exhibits non-collinear magnetic ordering and introduces significant uncertainties to the modelling of the electronic structure.⁶⁹⁻⁷⁰

Assessment of the electrochemical and structural stability was undertaken through the computation of cohesive energies of the materials of interest and also differences in the dissolution potentials $(\Delta E_d)^{71-73}$ for the Ru-Sb system. Both approaches have been previously validated through comparisons of the theoretical predictions and experimental electrochemical stability data for a range of systems, in particular metal oxides.⁷⁴⁻⁷⁶

For the Co-Sb combination, the simulated compositional phase diagram (Figure 5a) indicates a broad chemical potentials space that favours the formation of CoSb₂O₆ (structure shown in Figure S28) as opposed to individual oxides. Experimental data suggest that this phase dominates the [Co+Sb]O_v catalyst under the OER conditions and was therefore used for further theoretical analysis. As a point of reference in the calculations, Co_3O_4 – the dominating phase of the CoO_x control – was considered. Cohesive energies for Co₃O₄ and CoSb₂O₆ were simulated as -12.8 eV and -15.4 eV per formula unit, respectively, which indicates improved overall structural and electrochemical stability⁷⁷ of the antimonate as compared to the monometallic oxide. The improvement in cohesive energies is likely to be partially associated with the higher bond energy for Sb-O (434 kJ mol⁻¹) as compared to Co-O (397 kJ mol⁻¹),⁷⁸ although one might note that this difference does not appear sufficient to explain the calculated cohesive energies. Therefore, we hypothesised if the presence of Sb in the structure might strengthen the Co-O bond through changes in the electronic structure.

The above supposition was assessed through the analysis of the atom/orbital projected partial density of states (PDOS), calculated bond lengths and Bader charges. The PDOS for Co₃O₄ demonstrates a reasonable hybridisation of the oxygen p-orbitals with d-orbitals of cobalt in tetrahedral sites but not in octahedral ones; this is

in contrast to $CoSb_2O_6$ where only one type of Co sites is present and is strongly hybridised with O 2p (Figure S29a-b). It is also noted that the latter interact much stronger with Co 3d orbitals rather than with Sb 5p.

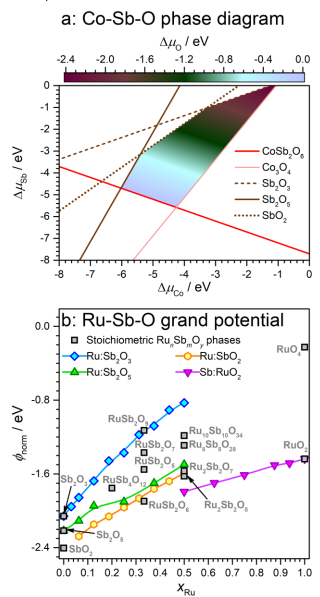


Figure 5. (a) Simulated compositional phase diagram of $CoSb_2O_6$ against individual oxides. Shaded area corresponds to the stable region of $CoSb_2O_6$, where the colour scale shows the allowed chemical potentials for oxygen; coloured lines and the corresponding spaces opposite to the shaded region present the stable regions of the identified individual oxides. (b) Normalised oxygen grand potential (ϕ_{norm}) *vs* ruthenium atomic fraction, $x_{Ru} = n/(n + m)$, calculated for $[Ru_n + Sb_m]O_y$ at T = 600 °C and 0.21 atm O₂ partial pressure. Sb:RuO₂ – Sb-doped RuO₂; Ru:SbO₂, Ru:Sb₂O₃ and Ru:Sb₂O₅ – Ru-doped antimony oxides. Calculated data are shown as symbols; lines are guides to an eye.

Since the electrocorrosion of materials is an interfacial phenomenon, the above findings were also corroborated by the analysis of the characteristic catalyst surfaces – (110) oriented facet for $CoSb_2O_6$ and (110)-A terminated surface for $Co_3O_4^{79.81}$ (Figure S28c-d). Comparisons of the PDOS of bulk and (110) surfaces suggest that the slab geometry produces surface states just above the valence band due to the undercoordinated Co^{3+} , Co^{2+} and O^{2-} ions, which renders these surfaces metallic. Similar to the bulk, 2p orbitals of the surface O in $CoSb_2O_6$ interact stronger with Co 3d as compared to Sb 5p orbitals. Enhanced O-p – Co-d interaction in $CoSb_2O_6$ is additionally verified by the calculated PDOS of sub d-orbitals of Co^{2+} in Co_3O_4 and $CoSb_2O_6$ (Figure S30).

The features of the PDOS discussed above can be expected to make the overall cohesive energy of $CoSb_2O_6$ more negative than that of Co_3O_4 and hence enhance the driving force required for the electrocorrosion of cobalt. Furthermore, the simulated Co-O and Sb-O bond lengths in $CoSb_2O_6$ were lower and higher than the corresponding expected values based on the combined ionic radii, respectively (2.08 Å calculated νs . 2.105 Å expected for high spin Co-O; 2.02 Å calculated νs . 1.98 Å expected for Sb-O). This internal compressive strain of the Co-O bonds can be associated with the higher electronegativity of Sb as compared to Co. Calculated Bader charges on Co^{2+} in Co_3O_4 and $CoSb_2O_6$ were +1.31e and +1.36e, respectively, which further supports the stronger Co-O bond in the antimonate as compared to the individual metal oxide.

Next, the ruthenium-antimony system was analysed following a similar approach. As discussed above, no robust experimental evidence for the formation of a stabile ruthenium-antimony oxide phase was obtained. In line with these experimental observations, total energies calculated by DFT simulations indicated that hypothesised RuSb₂O₆ phase is unstable with respect to the individual oxides (Table S5). This prompted us to undertake a simulation of a grand potential phase diagram of the Ru-Sb system (Figure 5b), which enables prediction of the thermodynamically stable compositions that can be formed under relevant synthesis conditions,⁸² *i.e.* at $T = 600^{\circ}$ C and 0.2 atm O₂ partial pressure in the present work.

Oxygen grand potentials were calculated for various $[Ru_n+Sb_m]O_y$ structures and compositions as a function of the metal atomic fraction, $x_{Ru} = n/(n+m)$. In addition to a broad range of stoichiometric compounds, substitutional doping of Sb into RuO₂ (Sb:RuO₂) and Ru in antimony oxides was considered. Among these different possibilities, the lowest ϕ_{norm} at $x_{Ru} = 0.5$, *i.e.* the highest stability under the conditions employed during the synthesis of $[Ru+Sb]O_{y_i}$ was calculated for the ruthenium(IV) oxide doped with antimony. In fact, Sb:RuO₂ are theoretically predicted to be more stable than the parent metal oxide within the x_{Ru} range examined (Figure 5b). At the same time, the experimental XAS data provide a compelling evidence for the dominant state of ruthenium in $[Ru+Sb]O_y$ being very similar to that in RuO₂ (*vide supra*). On this basis, further analysis focused on RuO₂ at low levels of Sb doping.

Differences in the electronic properties of the RuO₂ reference and Sb:RuO₂ were investigated based on the computations with a 2×2×2 supercell of tetragonal 6-atom RuO₂ unit cell containing 16 Ru and 32 O atoms. First, through the examination of the calculated spin polarised total density of states, we note that Sb substitution into RuO2 maintains its metallic character and even further improves the electrical conductivity as concluded from the shift of the conduction band (Figure S31), which is a positive finding from the perspective of the electrocatalytic activity. Further, and most importantly, improved stability of Ru⁴⁺ within the Sb:RuO₂ materials was confirmed by the positive difference in the dissolution potential⁷¹⁻⁷³ ΔE_{d} = 0.08 V and by the negative increase in the cohesive energies when compared to undoped RuO₂. Specifically, the cohesive energy changes from -3.28 (RuO₂) to -3.51 (Sb_{0.0625}:Ru_{0.9375}O₂) and -3.62 eV per unit formula (Ru_{0.813}Sb_{0.187}O₂). This favourable change is again explained by the enhanced strength of the Ru-O bond, which is

indicated by the stronger overlap of O 2p and Ru 3d orbitals in the PDOS (Figure S32) as well as by the higher Bader charge on ruthenium in Sb_{0.0625}: Ru_{0.9375}O₂ (+1.76e) as compared to RuO₂ (+1.70e). Enhanced charge donation from ruthenium to neighbouring oxygen atoms is also supported by the differences in the partial density of states of Ru 4d sub-orbitals (Figure S33).

In summary, the computational analysis suggests that the improvement in the stability of Ru and Co oxides upon combination with Sb oxide is majorly stemming from the electronic effects of antimony that strengthen the metal-oxygen bonds. Aggregate of the experimental and computational data for the Ru-Sb system suggests that, although the well-defined ruthenium antimonate phase is unlikely to be formed, doping of ruthenium(IV) oxide with antimony is possible. Taking into account the experimental data indicating the formation of the slightly distorted RuO₂ phase (Figures 2d, 3c and 4) and the very significant level of corrosion of antimony but not of ruthenium during the OER tests of $[Ru+Sb]O_{\gamma}$ (Tables 1 and 2), we hypothesise that Sb doping might be majorly confined to the surface of the RuO₂ crystallites. The formation of this protective layer is likely to be sufficient to provide the enhanced stability of the material against the electrocorrosion highlighted above (Figure 1) and as even more strongly emphasised through the durability tests at elevated temperature that are discussed below.

Longer-term operation at elevated temperature

A final set of tests was undertaken to assess the ability of the two most promising catalysts investigated herein, *viz.* [Mn+Sb]O_y and [Ru+Sb]O_y, to operate for extended periods of time and at elevated temperatures. Voltammetric analysis confirmed the expected positive effect of increasing the temperature on the kinetics of the OER catalysed by both Mn- and Ru-based materials (Figure 6a-b). The apparent activation energy of the OER at an *IR*-corrected overpotential of 0.6 V approximated from the backward potentiodynamic scans for [Mn+Sb]O_y is *ca* 20 ± 10 kJ mol⁻¹ (Figure S35). This is notably lower than values reported for other OER catalysts operating at low pH,⁸³⁻⁸⁴ and probably reflects the increasing instability of this material at elevated temperatures as discussed below.

While no significant deterioration of the $[Mn+Sb]O_{\nu}$ performance was observed in the voltammetric regime (Figure S36a), short-term 24 h galvanostatic tests revealed that this catalyst suffers detectable losses in activity at 60 °C (Figure S36b-c). During these experiments, the overpotential required to maintain the OER rate of 10 mA cm⁻² increased almost linearly, meaning an exponential decrease in the activity of the catalyst, at an average rate of *ca*0.001-0.002 V h⁻¹. When longer-term testing was undertaken at 80 °C, degradation at essentially the same rate was observed over the initial ca100 h of measurements (Figure 6b). Afterwards, an abrupt loss of the performance, reflected by an increase of the potential to ca2.1 V vs. RHE occurred, which was then maintained for at least another 24 h. The latter observation suggests that the change in the OER activity observed in these experiments is unlikely to be associated with the degradation of the FTO support, which would result in a complete loss of any ability of the electrode to sustain water electrooxidation due to the disruption of the electrical contact. On the basis of these new data (Figures 1b and 6b; Figure S36b), we conclude that manganese antimonate is a promising non-noble-metal-based water oxidation catalyst capable of a reasonably stable operation in acidic electrolyte solutions, but it is unlikely to be suitable for applications at 60 °C and higher temperatures.

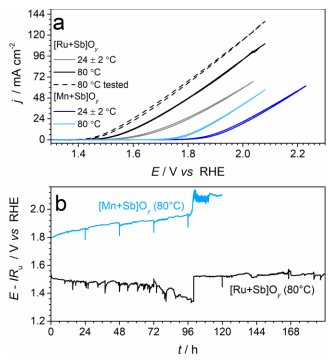


Figure 6. Performance of the [Ru+Sb]O_y (*black*) and [Mn+Sb]O_y (*blue*) catalysts during the OER in stirred 0.5 M H₂SO₄ at 80 °C: (a) cyclic voltammetry (scan rate 0.02 V s⁻¹; not / R_u -corrected; third scans) before (*vivid traces*) and after (*pale traces*) long-term tests shown in panel b (dashed curves show data collected at 24 ± 2 °C), and (b) / R_u -corrected chronopotentiograms (10 mA cm⁻²_{geom}).

The behaviour of [Ru+Sb]O_r at 80 °C was remarkably different to that of $[Mn+Sb]O_{\gamma}$ (Figures 6a and S37). First, the rutheniumbased catalyst demonstrated a robust operation for 192 h of chronopotentiometry at 10 mA cm⁻² and additional 1 h of chronoamperometry at 2.03 and 1.93 V vs. RHE (Figure 6b and S37b-c). Apart from the self-healing Co-Fe-Pb system,³⁸ we are not aware of any other recently reported OER electrocatalyst capable of similarly stable operation under such comparably harsh conditions. Second, comparisons of the cyclic voltammetric data recorded before and after long-term tests reveal a notable improvement in the performance of the catalyst. This positive change was also observed in short-term ambient temperature tests (Figure 1a) and is likely to be associated with a significant increase of the amount of ruthenium on the surface due to the loss of the catalytically inactive SbO_x (Table 2) but not Ru (Table 1). The stabilised catalytic activity for the OER of an essentially flat [Ru+SbOy] electrode at 80 °C and pH 0.3 corresponds to the reaction rate of 10 mA cm⁻²_{geom.} at an *IR*-corrected overpotential of 0.33-0.35 V (1.51-1.53 V vs. RHE).

CONCLUSIONS

The combination of transition metal and antimony oxides in a pursuit of the creation of robust and active catalysts for the electrooxidation of water under acidic conditions was found to produce very different outcomes depending on the chemical nature of the metal employed. While nickel and iron did not yield efficient catalytic systems, mixing Sb with cobalt, manganese and ruthenium resulted in materials with quite distinct structural features, but all demonstrating one very significant and critically important advantage with respect to the individual metal oxides – significantly improved operational stability. The origin of this positive effect can now be explained by the antimony-induced increase in the metal-

oxygen bond strength, which substantially suppresses the electrooxidative dissolution, as revealed through the computational studies. The theoretical treatment implemented herein might be considered for future high-throughput screening studies for the identification of robust electrocatalysts for the OER.

Another set of key findings of the present work stems from the exhaustive durability tests of the materials at different temperatures. While demonstration of apparently stable operation of the oxygen evolution reaction catalysts at low pH and ambient temperature on a short timescale does not present a very significant research challenge anymore, longer term operation at elevated temperatures of practical interest is still hard to achieve. Herein, the instability of the promising manganese-antimony oxide catalysts at 60-80 °C was revealed to indicate that this system needs to be further improved if it is intended to be designed for operation in PEM water electrolysers. Contrasting this behaviour, mixed ruthenium-antimony oxides were found to be stable in operation at 80 °C for at least one week, accompanied by negligible loss of the catalytically active metal into the solution. The lack of any recent reports on a similar performance during the electrooxidation of acidic water under comparably harsh conditions allows us to consider this result as outstanding and highly promising in the context of the development of the anode catalysts for PEM water electrolysers.

Finally, the aggregate of results presented herein suggests that there might be two different mechanisms for stabilising electrocatalytically active species like cobalt, manganese, or ruthenium oxides *via* the combination with an acid-stable "matrix" during water oxidation. The first is the formation of metal-matrix compounds either through doping or *via* transformation into new phases like antimonates. The second, less obvious and highlighted in detail herein for the ruthenium-antimony oxide system, is based on the intimate mixing of the discrete phases at the nanoscale. This mechanism might form a basis for a new promising strategy towards the design of electrocatalytic systems that exhibit outstanding stability while maintaining the high activity of the individual metal oxide.

ASSOCIATED CONTENT

Supporting Information. Experimental section and extended experimental data.

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SL designed and undertook experiments, analysed and interpreted data, and co-wrote the manuscript. MC designed and undertook experiments, analysed and interpreted data, and co-wrote the manuscript. AYadav undertook computational studies and co-wrote the manuscript. BK recorded, analysed and interpreted XAS data. JK undertook computational studies. TW collected and interpreted the TEM data. PC collected and analysed the XPS data. BJ assisted with the collection and interpretation of the XAS data. AT contributed to the data interpretation and manuscript preparation. DRM contributed to the data interpretation and manuscript preparation. RKH supervised and undertook XAS analysis, interpreted data and co-wrote the manuscript. AA designed and

supervised computational studies. AYella co-supervised the project and interpreted data. ANS devised and co-supervised the overall project, analysed and interpreted data, and co-wrote the manuscript.

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