Cite this: DOI: 10.1039/c0xx00000x

www.rsc.org/xxxxxx

ARTICLE TYPE

Electrochemical Water Splitting by Layered and 3D Cross-linked Manganese Oxides: Correlating Structural Motifs and Catalytic Activity

Arno Bergmann,* Ivelina Zaharieva,* Holger Dau and Peter Strasser

Received (in XXX, XXX) Xth XXXXXXXXX 20XX, Accepted Xth XXXXXXXXX 20XX 5 DOI: 10.1039/b000000x

Manganese based precious metal-free electrocatalysts for the oxygen evolution reaction (OER) are promising materials for energy storage systems based on dark or photo-coupled water electrolysis, because they are active, inexpensive and of low toxicity. In this work, atomic scale structure-activity relations of two different nano-structured manganese oxides, MnOx, are established using a combination 10 of X-ray absorption, diffraction and electrochemistry. Prepared by chemical symproportionation (s-MnO_x) and impregnation (i-MnO_x), the s-MnO_x catalyst consisted of a layered structure similar to δ-MnO₂ while the i-MnO₃ displayed a mixture of tunnelled, 3D cross-linked β- and a defective γ-MnO₂. During electrocatalytic oxygen evolution the structural motifs of both MnO_x remain largely unchanged, but the oxidation state of Mn increases from 3.5 to 3.9-4. Kinetic parameters of the electrocatalytic 15 oxygen evolution reaction were extracted using Tafel slope analysis and pH titration experiment and the role of the protons abstracted was analyzed. The study uncovers fundamental differences of general importance in the catalytic activity between layered and cross-linked structures. Exclusive presence of diμ-oxo bridged Mn ions in the layered structure is coupled to a pronounced redox and charge capacity behaviour. This ensured efficient use of surface and bulk active sites, and resulted in a relatively large ₂₀ Tafel slope. Consequently, the intrinsic OER activity is especially high in the s-MnO_x. In contrast, 3D cross-linked structures with both mono- and di-µ-oxo bridged Mn ions resulted in lower intrinsic activity but smaller Tafel slope, and thus favourable activity at technological water-splitting rates. The insights of this comparative study will provide guidance in the structural design and optimization of other non

25 a Department of Chemistry, Chemical and Materials Engineering Division, The Electrochemical Energy, Catalysis and Materials Science Laboratory, Technische Universität Berlin, Straße des 17. Juni 124, 10623 Berlin, Germany. Fax: +49 (0)30-314 22261; Tel: +49 (0)30-314 29542; E-Mail: bergmann@chem.tu-berlin.de; pstrasser@tu-berlin.de 30 ^b Fachbereich Physik, Freie Universität Berlin, Arnimallee 14, 14195 Berlin, Germany. E-Mail: ivelina.zaharieva@fu-berlin.de; holger.dau@fu-berlin.de; Fax: +49 30 838 56299; Tel: +49 30 838 3581 † Electronic supplementary information (ESI) available: Details on the syntheses of the supported manganese oxides as well as further 35 experimental results on the characterization of the support, the pH dependence of the MnO_x and additional XAS data and simulations. See DOI: 10.1039/b000000x/

precious metal oxide OER catalysts.

Introduction

development active, stable 40 electrocatalysts for water splitting is a key step in the commercialization of a so-called hydrogen economy which is based on the use of molecular hydrogen as a energy storage instead of fossil carbohydrates. The attractive concept of direct chemical storage of solar energy may be denoted as artificial 45 photosynthesis and is predicted to become a frontier technology in future energy sustainability.² To realize chemical energy storage devices for individual use or application in developing countries, the (electro)catalysts need to be based on low toxic and

- earth-abundant elements such as Mn and Fe; and they have to be 50 active under benign reaction conditions. However, the most active and stable electrocatalysts are based on precious metals like ruthenium and iridium in dimensionally-stable anodes working in harsh, acidic environments which limits their application to industrial processes.³⁻⁸
- Precious-metal free electrocatalysts receive strong attention since Nocera and co-workers reported efficient water oxidation by Co oxide (CoCat) prepared via anodic electrodeposition from a cobalt nitrate/phosphate buffer solution.9 This CoCat shows promising water oxidation activity in phosphate buffer at pH 7 60 and a self-healing mechanism in Co2+-containing electrolyte was proposed. Several researchers studied the CoCat with respect to the synthetic route, catalyst structure, and the mechanism of electrochemical oxygen evolution. 10-15 The CoCat can be described as a disordered molecular-like structure consisting of 65 edge-sharing Co octahedra (or di-µ-oxo/hydroxo-bridged Co ions) which can form extended planes of cobaltates. For the CoCat the reported atomic structure of low ordered Co oxide layers appears to be largely independent of the synthesis protocol used. 12, 14, 15 Analogously, a nickel oxide catalysts active for water 70 oxidation synthesized by anodic electrodeposition from borate or phosphate buffer showed water oxidation activity and a similar

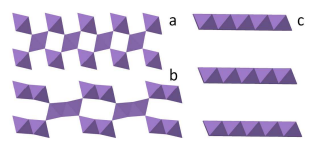


Fig. 1 Structures of selected MnO₂ modifications: β-MnO₂ (a), γ-MnO₂ (b), and δ-MnO₂ (c). The MnO₆ coordination octahedra are schematically shown. The structures are extended in the direction perpendicular to the 5 paper plane by rows of edge-sharing octahedra (in a, b, and c). This results in (1x1) tunnels of coordination octahedra (a), in (2x1) tunnels (b), and in a layered manganese oxide (c). In the layered oxides, the interlayer space is filled with water molecules and ions.

structure containing mainly di-µ-oxo/hydroxo bridged metal 10 ions. 16, 17

For the CoCat in neutral electrolyte, a chemical proton transfer step preceded by an electron transfer step in quasi-equilibrium was suggested to determine the rate of oxygen evolution which agrees well with former findings for Co-based electrocatalysts. 11,

In comparison to Co, the high abundance and low toxicity of manganese render Mn oxides as advantageous materials and make Mn-based catalysts highly favourable. 19 Biologically active Mn-containing water splitting complexes offer a possibly 20 coincidental, yet elegant analogy. In earlier works several Mn oxides varying in structure and oxidation state has been reported to be active for water oxidation, but the catalytic rates often were low and the catalyst remained without real practical application. 20-28 Structure-activity relations of water-oxidizing 25 Mn oxides have been discussed but are clearly insufficiently $understood.^{21,\,22,\,29}$

Like other binary transition metal oxides, manganese dioxides exhibit an exceptional variety of polymorphs relating, inter alia, to the small ionic radius for Mn⁴⁺. The schematic presentation of 30 Fig. 1 shows MnO₂ in layered structures with mainly edgesharing Mn octahedra such as δ-MnO₂ (Fig. 1c) or in tunneled structures (Figs. 1a & b) with a combination of corner- and edgesharing Mn octahedra such as β - and γ -MnO₂ with (1x1), (2x1) tunnels and a combination of both, respectively. 30-32

A common structural imperfection in tunneled MnO₂ materials is the de Wolff defect – an intergrowth of β -MnO $_2$ chains in γ -MnO₂ and vice versa. 33, 34 Furthermore, MnO₂ can exhibit Mn vacancies which lead to a replacement of Mn⁴⁺ ions by Mn³⁺ and results in a mean oxidation state of Mn lower than 4.35-37 The 40 presence of Mn³⁺ ions in the lattice can additionally cause microtwinning – a growth defect of the γ -MnO₂ lattice.³⁰

Synthetic δ-MnO₂ consists of MnO₆ layers where the interlayer space is filled with Na+ or K+ ions surrounded with water molecules. In case of an inhomogeneous distribution of the water 45 molecules and the foreign cations, the MnO₆ layers show a "turbostratic" arrangement and thus, are stacked without

Manganese oxides have long played a crucial role in battery applications and were structurally well characterized under 50 electrochemical battery environments³⁸. For catalytically active Mn oxides both layered and tunnelled structures have been

reported. 14, 15, 17, 20, 21, 36 In nature, water oxidation in PSII proceeds at a metal-oxido manganese-calcium complex.³⁹ Recently, a synthetic Mn-Ca water oxidation catalyst has been 55 prepared via a chemical synthesis route. 40 EXAFS studies revealed that in this amorphous material Mn ions are mainly interconnected via di-μ_{2/3}-oxo-bridges and similarly to δ-MnO₂ form layers, whereas the Ca ions are positioned in the interlayer space forming Mn₃CaO₄ cubanes or compensating the charge 60 deficiency on top of Mn-vacancies. The resulting structural motifs show striking similarities to the natural Mn-Ca complex in PSII.²² In a subsequent study, it has been shown that other redoxinert cations can be incorporated into the layered MnOx structure, but the obtained catalytic activity is highest when Ca ions are 65 present.²⁹

Recently, an electrolytically-prepared Mn oxide (MnCat) has been presented which does not contain Ca²⁺ ions but which shows electrocatalytic activity towards oxygen evolution in neutral electrolyte. In the structure of this MnCat, together with the 70 extensive di-μ-oxo bridging, a significant fraction of mono-μ-oxo bridged Mn ions was found whereas another MnOx, electrodeposited from the same media but using different electrochemistry protocol, showed almost exclusively di-u-oxo bridged Mn ions and had no catalytic activity.²¹ It was suggested 75 that Mn ions in lower oxidation state can take the structural (and perhaps also the functional) role of Ca in the catalytically active layered Mn oxides.

A synthetic tetranuclear Mn complex was incorporated into a Nafion® matrix where it decomposes and forms a layered Mn^{3+/4+} 80 oxide. 41, 42 Dissociation upon illumination results in disordered Mn²⁺ ions, which could be electrochemically re-oxidized to the layered, water-oxidizing Mn3+/4+ oxide. These finding suggest that cycling through various Mn oxidation states may be crucial in water-oxidation by amorphous Mn oxides.

As opposed to the CoCat, the mechanism of water oxidation catalysed by Mn-based catalysts is much less investigated. In 80's the mechanism of electrocatalytic oxygen evolution with Mn oxides in highly acidic and alkaline electrolyte was investigated using Tafel slope analysis, pH titration and a one electron transfer 90 step was argued to be rate-determining. 18, 26, 28 Recently, a mechanistic study of oxygen evolution at δ-MnO₂ in neutral electrolyte suggested that the disproportionation of Mn³⁺ to soluble Mn²⁺ may be a limiting factor of oxygen evolution activity at neutral pH.43 Increasing pH above 9 was suggested to 95 stabilize Mn³⁺ ions and to lead to a non-Nernstian potential dependence on pH during transition from neutral to alkaline reaction conditions and furthermore, to a higher oxygen evolution activity.43

Recently, two nano-structured manganese oxides supported on 100 multi-walled carbon nanotubes have been developed via two novel synthetic routes - symproportionation-deposition reaction and incipient wetness impregnation. These materials showed promising activity and stability for electrochemical water splitting in neutral electrolyte; the initial studies indicated that the 105 two oxides may differ in their atomic structure. 24

In this work, we studied the structure of these two catalytically active Mn oxides after deposition and after electrochemical oxygen evolution. Our goal was to investigate the structural transformations which the two catalysts undergo during oxygen

evolution reaction. The structure of as-synthesized Mn oxides and these used for electrocatalytic water oxidation were studied on different length-scales using X-ray powder diffraction (XRD), Xray absorption near-edge structure (XANES), and extended X-ray 5 absorption fine structure (EXAFS) spectroscopy. Furthermore, we investigated the electrocatalytic behaviour of these manganese oxides by cyclic voltammetry, Tafel slope analysis and pH titration experiments. The combination of structural information at the atomic level and comprehensive electrokinetic analysis 10 facilitates conclusions on crucial structure-activity relations in the synthetic Mn-oxide water-oxidation catalysts.

Experimental

Syntheses

Carbon-supported manganese oxide was synthesized via two 15 different preparation methods - syn-proportionation depositionprecipitation (s-MnO_x) and incipient wetness impregnation (i-MnO_x).²⁴ The Mn concentration was determined using inductively coupled plasma - optical emission spectroscopy (ICP-OES) to 1.69 mmol g⁻¹ and 1.75 mmol g⁻¹ (9.30 wt.% and 20 9.61 wt.%) for s- and i-MnO_x, respectively. Further details are given in the ESI†.

X-ray Diffraction

X-ray powder diffraction was conducted on the as-prepared powders with Cu K_{α} radiation in reflection mode using a Bruker 25 D8 Advance equipped with 2.3° soller slits and a positionsensitive detector (PSD)⁴⁴. Powder diffraction pattern were collected between 10° and 75° using a step size of 0.02° and counting time of 7 s. Divergence and antiscattering slit were adjusted for a constant illuminated area.

30 Electrochemical Characterization

The catalytically active MnO_x powders were deposited as thin films on glassy carbon (GC) rotating disk electrodes (Pine Instruments, Ø 5 mm). Prior to the deposition GC electrodes were polished and stepwise cleaned in ultrasonic bath using ultrapure 35 water and acetone. 5 mg of catalysts powder was suspended in a mixture of 1.99 ml of ultrapure water, 500 µl of 2-propanol and 10 µl of Nafion® solution (5wt.% of stock solution, Sigma-Aldrich) followed by an homogenization using a horn sonicator. Finally, this catalyst ink (10 µl of the catalysts suspension) was 40 dispersed on the electrode and dried in air at 60 °C for 10 min. Thus, a Mn loading of the GC electrodes of 172.8 nmol cm⁻² and 178.5 nmol cm⁻²was achieved for s- and i-MnO_x, respectively.

Electrochemical measurements were conducted with a threeelectrode rotating disk electrode (RDE) setup in a custom-made 45 glass cell using a PINE® rotator and a Biologic SP-200 potentiostat. Platinum gauze acted as counter electrode and a commercial reversible hydrogen electrode (RHE, Gaskatel) or Ag/AgCl (3M KCl, World Precision Instruments) connected via a Haber-Luggin capillary was used as a reference electrode 45, 46.

Electrochemical experiments were conducted in 0.1 M phosphate buffer (KPi) at pH7 prepared by mixing 0.1 M K₂HPO₄ (>98%, Sigma-Aldrich) and 0.1 M KH₂PO₄ (99%, Merck) aqueous solutions until the desired pH was obtained. Prior to electrochemical measurements N₂ was bubbled through 55 the electrolyte for at least 15 min and was switched to blanketing

flow during measurements. Working electrode was rotated at 1600 rpm during experiments to ensure a hydrodynamic equilibrium. Unless differently states, all electrode potentials had been corrected for ohmic losses using electrochemical impedance 60 spectroscopy and are referred to RHE.

For Tafel slope analysis current density was recorded stepwise between 1.5 and 1.9 V vs. RHE in a quasi-stationary mode. The electrode potential was increased and decreased in two separate experiments with steps of 20 mV hold for 5 min and both curves 65 were averaged. To determine the ohmic resistance, impedance spectra were recorded at every potential step between 10 kHz and 1 Hz with a modulation amplitude of 10 mV.

To investigate the effect of pH of the electrolyte at constant current and electrode potential, different aliquots of 40 wt.% 70 KOH were added every 3 min to 0.1 M phosphate buffer and pH was continuously measured. The initial pH of the phosphate buffer was ~4.7.

X-ray Spectroscopy

XANES/EXAFS experiments were carried out at the BESSY 75 synchrotron radiation source operated by the Helmholtz-Zentrum Berlin (HZB). The measurements at the manganese K-edge were conducted at the KMC-1 bending-magnet beamline at 20 K in a cryostat (Oxford-Danfysik) with a liquid-helium flow system. Further details are given ref.²¹

Samples for XAS experiments were prepared on glassy carbon cylinders, in analogy to the electrochemical RDE experiments. Electrochemical treatment in the OER range was conducted at 1.763 V vs. RHE for 3 min. After removal from the electrolyte under potential control, the electrode was dried immediately 85 using air and nitrogen and immersed in liquid nitrogen where it was until XAS measurements.

Results and Discussion

Structural Investigations

Long-range order and morphology

90 Fig. 2 shows the powder x-ray diffraction pattern of s-MnO_x and i-MnO_x recorded between 10 and 75° using Cu K_{α} radiation. The graphite structure of the MWNT_{ox} causes three strong diffraction peaks at $\sim 25^{\circ}$, $\sim 43^{\circ}$ and $\sim 53^{\circ}$ (Fig. S1).

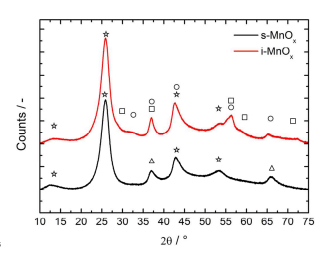


Fig. 2 Powder x-ray diffraction pattern of s-MnO_x (black) and i-MnO_x (red) between 10° and 75°, at Cu K_α radiation. Reflections are labeled by squares for β -MnO₂, by circles for γ -MnO₂, by triangles for the layered MnO₂ phase, and by stars for the reflexes caused by the support material. 5 The latter are shown in Fig. S1.

For s-MnO_x, two asymmetric reflections at ~37° and ~66° are caused by the defective crystal structure of a δ -MnO₂ phase (space group: C-1). These two reflexes are in good agreement with the reflexes arising from regular distances within the layers consisting of edge-sharing Mn octahedra. The reflexes can be assigned to the (200)/(110) and (310)/(020) doublets at ~37° and ~66°, respectively. The absence of reflexes (h k l \neq 0), especially the strong (001) reveals the absence of long-range order of the layer stacking. This can be induced by MnO₆ monolayers or variations in the interlayer distance due to "turbostratic" disorder.³¹

In the case of i-MnO_x, the diffraction pattern exhibits more than 10 reflexes arising from the γ -MnO₂ (space group: Pnma; PDF#00-044-0142) and the rutile-type β -MnO₂ (space group: 30 P4₂/mnm, PDF#01-072-1984) as a major and minor phase, respectively. Crystallite size analysis from reflex broadening is generally hindered for β - and γ -MnO₂ because size-induced broadening is superimposed by defect-induced broadening of the reflexes. Furthermore, the limited intensity of the reflexes made a 35 crystallite size analysis for the γ -MnO₂ phase in i-MnO_x impossible. In the case of β -MnO₂, the (220) reflex at ~59° is not affected by defects in the rutile lattice and could be used to estimate the size of the β -MnO₂ crystallites to be below 5 nm. 30

According to the diffraction pattern, the γ -MnO $_2$ phase of i-MnO $_x$ exhibits a significant amount of single chains of Mn octahedra in the γ -MnO $_2$ lattice (de Wolff defects) and a minor fraction of twinning faults. The presence of de Wolff defects are visible in the significant shift of the (130) reflex at ~32° with more than 3° from its ideal position. De Wolff defects can reduce the angular separation of the γ -MnO $_2$ (110) and (130) from 12 to γ° . However, the γ -MnO $_2$ (110) reflex is hidden under the C reflex at ~25°. The presence of any sizeable contribution from an α -Mn $_2$ O $_3$ phase in i-MnO $_x$ (as a possible origin of the diffraction reflex at ~32°) was excluded from XAFS characterization discussed further below.

Microtwinning in the γ-MnO₂ lattice could be extracted from the merged (221)/(240) doublet of γ-MnO₂ at ~56° but the real fraction of microtwinning is significantly lower because higher microtwinning would lead to a shift of the (130) reflex towards bigher diffraction angles, which was not observed in our data.

These findings from X-ray diffraction analysis are in agreement with earlier TEM analysis showing that s- and i-MnO_x consists of Mn oxide particles with a size of 5-10 nm attached to the MWNT_{ox}. 24

Manganese Oxidation States from XANES Spectra

To get further insights into the structure of the two catalytically active oxides at the atomic scale, we employed X-ray absorption spectroscopy at the Mn K-edge (Fig. 3).

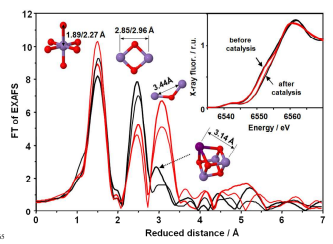


Fig. 3 X-ray absorption spectra of s-MnO_x (black) and i-MnO_x (red) deposited on electrodes before and after catalysis. The edge region of the spectra (XANES) is shown in the inset. The average oxidation state of Mn in the two samples directly after deposition on the electrode (thick lines) is around 3.5, while after 3 min of operation at 1.763 V in 0.1 M KPi at pH 7, it increases to 3.9-4.0 (thin lines). The structural motifs considered in the EXAFS simulations are schematically depicted next to the corresponding FT peak and the obtained interatomic distances are indicated. (Mn⁴⁺ ions are presented as spheres in magenta, Mn³⁺ ions in dark purple, and oxygen as red spheres.)

The XANES (X-ray absorption near-edge structure) region is particularly sensitive to the average oxidation state and local coordination geometry. 48 In general, the energy of the X-ray absorption edge increases with increasing oxidation states of the 80 absorbing atom due to the rise in the electron binding energy. 49 On the basis of calibration with Mn compounds with known structure, the average oxidation state of manganese in i-MnO_x powder was around 3, while in the s-MnO_x it was around 3.5. After deposition on the GC electrode the oxidation state in both s-85 MnO_x and i-MnO_x was estimated to be around 3.5. (Fig. S2 and S3). This indicates that both oxides contain a large fraction of Mn⁴⁺ already before applying an oxidizing potential, but a significant amount of Mn ions are in lower oxidation state (+2 or +3). The result for the s-MnO_x is in a good agreement with the 90 previously determined Mn oxidation state by Mn L-edge measurements and temperature-programmed reduction (TPR).²⁴ The oxidation state for i-MnO_x previously obtained from Mn Ledge XAS, however, was +2, explainable by Mn reduction occurring either during the thermal treatment at low oxygen 95 partial pressure and or caused by x-ray irradiation during data collection. In the present study the thermal treatment was performed at 110 °C instead of 80 °C and at ambient air. X-ray photoreduction was avoided in the present study by collecting (i) the XAS spectra at temperature of 20 K and (ii) using strictly 100 limited periods of X-ray irradiation on a single sample spot.

For transition metal based catalyst, the oxidation state of the metal ion was suggested to depend on the applied potential during electrocatalytic OER. ^{15, 21, 50, 51} To determine the extent of these changes for the Mn ions in the studied materials, we measured

XAS spectra of s- and i-MnO_x frozen in liquid nitrogen immediately after applying 1.763 V in 0.1 M KPi at pH 7 for 3 min. In both oxides, a clear shift of the edge position in the XANES spectra was observed (Fig. S2) corresponding to an 5 increase of the mean Mn oxidation state to almost +4.0. The same

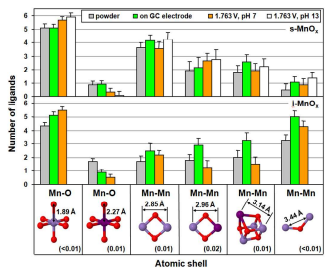


Fig. 4 Coordination numbers and interatomic distances obtained by joint simulation of the k³-weighted EXAFS spectra from s-MnO_x (upper panel) and i-MnOx (lower panel) before and after operation as water oxidation 10 catalysts in 0.1 M KPi (pH 7) or 0.1 M KOH (pH 13). The corresponding absorber-backscatter distances were kept equal for all seven spectra and are displayed together with the corresponding structural motif (colour code as in Fig. 2). The error bars represents the 68% confidence interval; the error for the interatomic distances is given in parentheses below the 15 corresponding structural motif. For further details, see Fig. S6.

shift in the edge position was observed for s-MnO_x frozen after 3 min operation at 1.763 V in 0.1 M KOH at pH 13.

Despite the similar Mn oxidation state of the two oxides after operation as catalysts, the quasi-in-situ experiment revealed 20 differences in XAS edge shape which imply that the coordination geometry of the Mn ions is modified. To address these structural differences, extended X-ray absorption fine-structure (EXAFS) spectra were analysed.

25 Structural information from EXAFS

The Fourier transforms (FTs) of the EXAFS spectra of s-MnO_x and i-MnO_x are presented in Fig. 3. In the FT, typically each peak corresponds to a specific absorber-backscatterer vector. In our case the absorber is a Mn ion, and the backscattering atoms can 30 be either oxygen atoms, or Mn ions surrounding the absorber. The distance between the absorber and backscatterer corresponds to the position of the FT peak plus about 0.4 Å. 48

Both s-MnO_x and i-MnO_x show a first peak at reduced distance of around 1.5 Å, which is typical for 6-fold oxygen coordinated 35 Mn^{4+} ions and a second peak at around 2.5 Å related to di- μ -oxo bridged Mn^{4+} ions. $^{52,~53}$ For i-MnO $_x$ there is also a well pronounced third peak, assignable to mono-µ-oxo connected Mn ions (longer Mn-Mn distance). 54 This assignment of the first three peaks can be made by comparison to the FT spectra of Mn oxides 40 with known structure (Fig. S4-B). Visual comparison to the FTs of EXAFS spectra from reference Mn oxides suggests that while the s-MnO_x has layered δ-MnO₂-like structure composed predominantly from edge-sharing (di-µ-oxo connected) Mn

octahedra (Fig. 5), the i-MnO_x contains a significant fraction of 45 corner-sharing (mono-u-oxo connected) Mn octahedra resulting in formation of tunnels.

To confirm this interpretation and to analyse the structural changes during the electrochemical water oxidation, we simulated seven k³-weighted EXAFS spectra measured from s-MnO_x and i-50 MnO_x (i) as powders, (ii) after deposition on GC electrode, (iii) and after catalytic operation at 1.763 V vs. RHE at pH 7. The GC electrode covered with s-MnOx was measured also after electrocatalytical operation at pH 13.

In the EXAFS simulations, we assumed two Mn-O distances 55 contributing to the first Mn coordination shell: the shorter Mn-O distance is assigned to Mn4+-O vectors and the longer is assignable either to Mn3+-O vectors along the elongated Jahn-Teller axis or to Mn²⁺-O vectors. The sum of the coordination numbers for these two shells was kept equal to 6 because the 60 small pre-edge feature of the XANES data suggested that in all samples the majority of the Mn ions are 6-fold coordinated. The other FT peaks were simulated with three shells of Mn backscatterer: a short Mn-Mn vector corresponding to edgesharing MnO₆ octahedra (second peak), a longer Mn-Mn distance 65 corresponding to corner-sharing MnO₆ octahedra (the third peak visible clearly in i-MnO_x samples) and an elongated di-μ-oxo bridge, which could be hidden due to the large third peak in i-MnO_x but is visible in the s-MnO_x. Such an intermediate distance was found in layered Mn oxides before and likely corresponds to 70 Mn₄O₄ cubane motifs formed by attachment of a Mn ion in an oxidation state lower than +4. Also in other water-oxidizing oxides, formation of a metal-oxo cubane has been reported.^{29, 53} The simulation results shown in Fig. S5 confirm the qualitative interpretation given above and specify internuclear distances at 75 high precision.

We approached identification of structural changes resulting from application of oxidising potential during operation of the electrocatalyst. In tendency, Mn-O distances were decreased after operation at 1.763 V, confirming the increase of the average Mn 80 oxidation state implied by the XANES data. For the Mn-Mn vectors in the s-MnO_x sample, there appears no significant change of the coordination numbers, neither due to attachment to

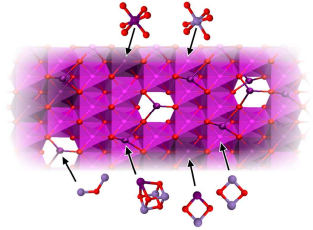


Fig. 5 Schematic model for a distorted MnO_x layer fragment formed from 85 edge-sharing (di-μ-oxo bridged) Mn⁴⁺O₆ and Mn³⁺O₆ octahedra. Some Mn ions from the layer are missing (defects), and other Mn ions are bound on top (or below) of the layer resulting in mono-μ-oxo bridges

(corner-sharing octahedra). At these defect sites, a row of edge-sharing octahedra could grow perpendicularly to the first one; this eventually may lead to formation of tunnel-like structures. Binding of a Mn³⁺ ions on top of the layer at sites without layer defects results in formation of a 5 (distorted) Mn₄O₄ cubanes. (Mn⁴⁺ ions in magenta, Mn³⁺ ions in dark purple, O in red.)

the GC electrode nor due to electrochemical treatment oxidation. For i-MnO_v, there is an increase in the EXAFS coordination of the Mn-Mn vectors caused by attachment to the electrode. 10 Interestingly, application of an oxidizing potential decreases the number of Mn-Mn vectors. Although these changes appear to be significant, due to the large error bars an unambiguous statement about extent and character of the structural changes cannot be made based on these fit results.

The EXAFS fit result in a mean Mn-Mn distance of (2.89 ± 0.01) Å between the Mn nuclei of two edge-sharing MnO₆ octahedra. The XANES data show prevalence of Mn⁴⁺ but also Mn³⁺ ions may be present. For di-µ-oxo bridged Mn⁴⁺ ions, the obtained Mn-Mn is often shorter, namely 2.86 Å.21, 53, 55, 56

20 Moreover, we find that the Debye-Waller (DW) parameter corresponding to this shell are unusually high (Fig. S5-C) indicating either a broad Gaussian-shaped distance distribution function or a bimodal distribution, both centred around a mean Mn-Mn distance of 2.89 Å. The latter option is clearly more 25 likely and implies that there are two specific Mn-Mn distances

which contribute to the second FT peak. The shorter (< 2.89 Å) and the longer one (> 2.89 Å) may be assignable to Mn^{4+} -(μ -O)₂-Mn⁴⁺ and Mn⁴⁺-(μ-O)₂-Mn³⁺ motifs, respectively.^{29, 56} For more refined analysis, we approached a 'joint-fit' of all seven EXAFS

30 spectra keeping the interatomic distances for the corresponding shells equal in all spectra. The increased significance of the jointfit approach facilitated, *inter alia*, resolution of two short Mn-Mn distances of 2.85 and 2.96 Å, respectively. The joint-fit results are shown in Fig. 4 and discussed in the following.

In the first Mn coordination sphere, short 1.89 Å Mn-O vectors typical for 6-fold coordinated Mn4+ are prevalent. A small fraction of up to 15% longer Mn-O distances of 2.29 Å, is consistent with a minor contribution of Mn2+ or Mn3+ ions (for Mn3+ ions such long Mn-O distances are found along the Jahn-40 Teller axis). 57, 58

The results of the joint-fit approach (Fig. 4) confirm the absence of major structural change for the s-MnO_x. After catalyst operation at 1.763 V, there may be a slight decrease in the number of the out-of-layer Mn³⁺ ions (cubane motifs, 3.14 Å) and 45 a slight increase of the in-layer Mn³⁺ ions (2.96 Å), resulting in larger heterogeneity of the Mn-Mn distances contributing to the second FT peak and thus a decreased peak height. These changes are relatively small and thus close to the error limit. The sum of the two short Mn-Mn vectors is around 6 as expected for 50 perfectly ordered layers of a δ-MnO₂. 53 Interestingly, this value appears to increase slightly upon catalyst operation at 1.763 V, possibly explainable by Mn-Mn vectors from out-of-layer Mn⁴⁺ ions involved in $Mn^{4+}_{4}(\mu O)_{4}$ cubane formation.

When an oxidizing potential is applied to i-MnO_x, larger 55 structural rearrangements can be observed (Fig. 4). There is a clear decrease of the mono-µ-oxo and di-µ-oxo vectors with participation of Mn³⁺ ions (2.96 Å and 3.14 Å distances). This decrease in the number of in-layer Mn3+ ions (2.96 Å) contrasts the slight increase in s-MnO_x. Also the Debye-Waller parameter 60 (distance spread) for the Mn-O distances increase at higher potential for s-MnO_x and decrease for i-MnO_x (Fig. S6), which could reflect different protonation patterns of the oxygen atoms in bridging and terminal position.

The presented results indicate clear differences between the

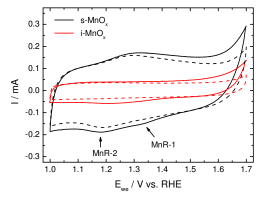


Fig. 6 Cyclic voltammograms (CV) of first (solid) and 11th (dashed) scan of s-MnOx (black) and i-MnOx (red). The CVs were recorded in 0.1 M KPi at pH 7 with a scan rate of 100 mV s⁻¹. Two minima resolved in the s-MnO_x CV are labelled by MnR-1 and MnR-2, respectively.

70 structure of the two oxides in their resting state (XRD, EXAFS) and regarding the structural changes induced by application of an oxidizing potential (EXAFS). The question arises of whether and how these structural differences relate to the elementary mechanism of the water oxidation and oxygen evolution. To 75 investigate this further, s-MnO_x and i-MnO_x were characterized electrochemically and indeed, confirmation for different processes involved in water oxidation was found in the different kinetic characteristics of the two materials, as described in the following.

80 Electrochemical and Electrokinetic Characterization

Electrochemical Characterization

Cyclic voltammetry was used to investigate the redox behaviour of the Mn ions in s- and i-MnO_x in 0.1 M KPi at pH 7. Figure 6 shows the first and the 11th potential scan recorded between 1.0 85 and 1.7 V at a scan rate of 100 mV s⁻¹ with a starting potential of 1.5 V. Both materials exhibited a mainly capacitive voltammogram between 1.0 and 1.6 V caused by the charge storage capability of both the high-surface carbon support and MnO_x. The capacitance of s-MnO_x is clearly larger than that of i-90 MnO_x, which does exceed the capacitance of the carbon support only marginally (Fig. S8). This suggests a higher extent of oxidative charging (by Mn oxidation) in s-MnO_x. At potentials above 1.6 V, the current increases inter alia due to oxygen evolution.

Both voltammograms show the differences in the redox behaviour of the Mn ions in the MnO_x. The first cathodic potential scan of both MnO_x exhibits a double reduction feature (MnR-1 and MnR-2) between ~1.45 and 1.10 V. During the subsequent anodic scan, s-MnO_x exhibits the corresponding 100 oxidation peak whereas no oxidation peak is resolved for i-MnO_x. The transferred charge during electrochemical reduction of the Mn ions in i-MnO_x is significantly lower. Comparing the potential of the reduction feature of s- and i-MnO_x with literature values might point toward the redox couple of Mn²⁺/MnO₂.⁵⁹ 105 This assignment is in line with the XANES results which show

that at an OER potential mainly Mn⁴⁺ ions are present in the catalyst. However, the presence of a double reduction feature in s- and i-MnO_x suggests a two-step electrochemical reduction of Mn⁴⁺ to Mn³⁺ and further on to Mn²⁺. In electrodeposited MnO_x

5 films, similar redox peaks were successfully connected to

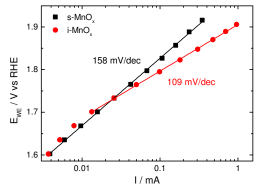


Fig. 7 Tafel plot of s-MnO_x (black) and i-MnO_x (red) extracted from quasi-stationary potential-step RDE experiments in 0.1 M KPi at pH7. All electrode potentials have been iR-corrected. The respective Tafel 10 slopes as determined by linear regression are indicated.

changes in the visible light absorption of the oxide and interpreted as Mn oxidation state changes.²¹

During continuous potential cycling between 1.0 and 1.7 V, the Mn⁴⁺ reduction features disappear completely in case of i-15 MnO_x whereas s-MnO_x continuous to exhibit both redox features even after 10 potential cycles with only slightly reduced charge. Thus, the Mn ions in i-MnO_x passivate irreversibly (but notably not completely) during initial reduction and the voltammogram becomes almost purely capacitive, whereas we observe only little 20 redox passivation in s-MnO_x.

Tafel slope analysis

Fig. 7 shows Tafel plots for s- and i-MnO_x extracted from quasi-25 stationary experiments in 0.1 M phosphate buffer at pH 7. (In Fig. 7 and elsewhere, the detected electrical current is given; the current density per cm² is by a factor of about 5 higher than the indicated current.) Both s- and i-MnO_x show a linear E vs. log(I) dependency in the potential range of 1.6-1.9 V and 1.7-1.9 V, 30 respectively. The onset potential for the electrocatalytic oxygen evolution is ~100 mV lower for s- than i-MnO_x. Linear regression of the experimental data within the mentioned potential ranges led to Tafel slopes of 158 and 109 mV/dec and thus, experimental anodic transfer coefficients, β , of 0.39 and 0.54 for s- and i-35 MnO_x, respectively. The exchange current density i₀ of s-MnO_x $(\sim 10^{-3} \text{ mA})$ is significantly higher than for i-MnO_x $(\sim 10^{-7} \text{ mA})$. Thus, s-MnO_x shows a significantly higher intrinsic electrocatalytic activity than i-MnO_x, whereas at higher electrode potentials the OER activity of i-MnO_x is superior. Thus, we 40 conclude conservatively that there are pronounced functional differences between s-MnO_x and i-MnO_x in the rate-determining process and possibly regarding number and location of active sites.

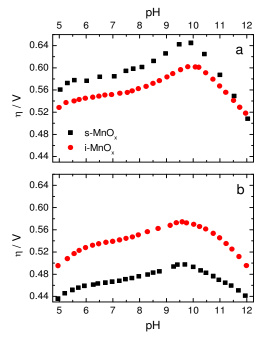


Fig. 8 pH dependency of the overpotential (η) of s- and i-MnO_x determined in 0.1 M KPi at constant currents of (a) 50 μA and (b) 20 μA . The overpotential has been calculated according to Eqn. S1.

50 pH dependence of overpotential and current

Fig. 8 shows the dependence of the overpotential, η , on the H⁺/OH⁻ concentration in the electrolyte between pH 5 and ~12 at a constant current of 20 μA and 50 μA. At both currents, the η-55 pH profile for both MnO_x can be divided into two regimes. For example at a constant current of 20 µA (Fig. 8b), the overpotential increases between pH 5 and 9.6 for s-MnO_x (i-MnO_x) from 435 mV (495 mV) to 497 mV (574 mV). Above pH 10, the overpotential decreases to 441 mV (495 mV). Thus at 60 20 µA, s-MnO_x exhibits a lower overpotential over the investigated pH range than i-MnO_x, due to its higher intrinsic electrocatalytic activity (higher exchange current density), but the principal pH dependence of the catalytic current of two oxides is the same. At 50 µA (Fig. 8a), i-MnO_x exhibits a lower 65 overpotential, due to its lower Tafel slope. Yet again, the principal pH dependence of the two oxides is the same. Thus we conclude that an intricate coupling of electrocatalysis to (de)protonation reactions is functionally decisive in both oxides. Comparison of the two panels of Fig. 8 reveals that the crossover 70 point of the Tafel plot is observable at all investigated pH values implying that at all pH, the exchange current in s-MnO_x is higher than in i-MnO_x whereas the Tafel slope always is lower in i-MnO_x. This implies that the functional differences between the two oxides indicated by different Tafel slopes and exchange 75 currents do not relate to differences in the coupling of the ratedetermining step to protonation reactions.

We also investigated the pH-dependence of the current at a constant catalytic potential of 1.35 V vs. NHE (Fig. 9).

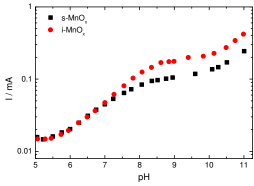


Fig. 9 pH dependence of the catalytic current (I) determined in 0.1 M KPi at a constant electrode potential of 1.35 V vs. NHE.

With increasing pH, a biphasic increase in the catalytic current is s observed. The first increase is detected for pH-values ranging from 5.7 to 8.7, that is, around the second pK of the phosphate ion (p $K_{2nd} = 7.2$, $H_2PO_4^{1-} \leftrightarrow HPO_4^{2-} + H^+$,); at pH values below 5.7 and above 8.2, plateau levels are reached. Thus the pH dependence of the catalytic current at a constant potential of 10 1.35 V vs. NHE resembles the pH titration curve of the phosphate ion. This finding suggests that stoichiometry or concentrations of [HPO₄²⁻] and [H₂PO₄¹⁻] determine catalytic activity. At even higher pH (>10), a further increase is observed, explainable by the increasing availability of OH ions at higher pH values.

15 Discussion

The oxygen evolution reaction is a complex multi-electron transfer process controlled by macroscopic, microscopic and atomistic parameters, such as the number, accessibility and character of catalytically active sites, the chemical reaction 20 mechanism, the concentration of reactant, electrical and protonic conductivity and removal of products.

The number of catalytically active sites depends on the number of accessible Mn ions for water molecules, which, in turn, is strongly connected to the structure and the morphology of the 25 electrocatalyst. If no Mn ions of the bulk material were involved in the water splitting, the number of active sites would be (most probably) directly proportional to the surface area of MnO_x and thus, particle size. However in the case of a layered MnO₂ phase, bulk Mn ions are accessible for water molecules and cations alike 30 and thus, the number of accessible active sites is more likely approaching the total number of Mn ions. Furthermore, the local coordination of the Mn ions can critically determine how accessible they are and to what extent they can contribute to the overall catalytic activity.

For the two nano-structured Mn oxide catalysts considered herein, there is a clear and important structural difference in the Mn coordination, e.g. there exist nearly exclusively di-u-oxo bridged Mn ions in the s-MnOx, whereas more mono- than di-µoxo-bridged Mn ions exist in the i-MnO_x catalyst. The 40 electrochemical analysis suggests that higher intrinsic onset activity in the low-current regime is associated with the prevalent presence of di-u-oxo bridged Mn ions in form of a layered oxide (Fig. 1c) coupled with a significantly higher electrochemical capacitance. Thus, we hypothesize that the electrochemical 45 accessibility of catalytically active Mn ions, likely one important

controlling factor for OER activity, is higher in the layered s-MnO_x structures than in the cross-linked i-MnO_x structures. This is consistent with the fact that in layered δ-like MnO₂ structure bulk Mn ions are much better accessible for the reactant 50 molecules and cations through the interlayer space. This is in clear contrast to tunnelled MnO2 structures which generally show a lower charge storage capacity compared to layered Mn structures and the charge storage capacity is proportional to the void space in the lattice. The (1x1) and (2x1) tunnels in i-MnO_x 55 exhibit certain proton conductivity but are not accessible for water molecules and cations. 35, 60 In conclusion, the participation of bulk Mn ions could lead to a larger number of active sites and thus, a higher exchange current density for s-MnO_x than i-MnO_x.

The structural motif of di-u-oxo bridged transition metal ions 60 resulting in layers (or layer fragments) of edge-sharing MO₆ octahedra have proven to be a unifying feature of many oxygenevolving Co and Ni oxide electrocatalysts prepared by electrodeposition. 12, 14, 15, 17 Looking at electrodeposited OER catalyst materials, the beneficial effect of layered oxide with 65 prevalence of di-µ-oxo bridging between metal ions and their large interlayer space filled by water and further ions becomes obvious: bulk metal ions are able to participate as catalyticallyactive sites. 15

Beyond its favourable intrinsic activity (high exchange current 70 density), Tafel slope analysis showed that layered s-MnO_x displayed inferior electrocatalytic behaviour at larger current densities reflected by a larger Tafel slope of 158 mV/dec compared to 109 mV/dec for the i-MnO_x. The unfavourably large Tafel slope of the s-MnO_x likely relates to the high level of order 75 within the layers of edge-sharing MnO₆ octahedra, which is clearly higher than found in the electrodeposited MnOx of ref.²¹ (Tafel slope of 90 mV/dec) and in water-oxidizing MnO_x particles. 21, 29 Less order implies a higher number of defect sites, e.g. Mn vacancies; the structures of putative defect sites are 80 shown in Fig. 5. These defect sites prevent prevalence of di-μ₃oxo thereby resulting in more di-µ2-O(H) bridges between Mn ions as well as in a higher number of terminal water coordination sites; both features likely are crucial for OER activity.³ And indeed, an inverse relation between di-µ3-O bridging (low 85 number of di-µ2-O(H) bridges and defect sites in general) and OER activity has been demonstrated for Co- and Mn-based oxides before 15, 21, 22, but a comparison of Tafel slopes had not been approached. We now propose that an unfavourably high Tafel slope of the OER reaction can result from a strong 90 prevalence of unreactive di-μ₃-oxo bridges between the first-row transition metal ions of the respective oxide, which is caused by the octahedral cross-linking and/or a level of structural order that is too high. This suggestion is supported by the results of the present investigation and immediately provides a guideline for 95 designing catalytic oxides with a favourably low Tafel slope. We note in passing that di-µ2-oxo bridges are present also at the ideal β -MnO₂(110) and γ-MnO₂(100) surfaces and so, similar structural motifs could participate in the electrocatalysis of s-MnO_x ('internal' surface between oxide fragment and intercalated water) as well as of i-MnO_x ('external' oxide-electrolyte surface).

Tafel slopes of ~120 mV/dec have been determined for Mn oxide electrocatalysts and from this, a one-electron transfer step has been concluded to be the rate-determining step (rds) in

MnO_x.²⁶ Herein, a lower Tafel slope of 109 mV/dec as determined for i-MnO_x does not support a simple one-electron transfer step to be rate determining during OER, neither does the value of 158 mV/dec in s-MnO_x. In line with this conclusion, the 5 detected pH-dependence of overpotential and catalytic current rule out rate-limitation by a simple one-electron transfer step, but points towards a role of proton-coupled electron transfer in the rds.

A critical factor determining the OER activity is ion transport, 10 specifically proton removal from the catalytic site and proton transfer into the electrolyte. Relevant factors may be the proton conductivity and uptake capability of the electrolyte. The log(I)pH profile of Fig. 9 reflects the 2^{nd} phosphate pK, thereby illustrating a crucial role of the phosphate protonation state. In the 15 η-pH profile of Fig. 8, we find minimal catalytic activity at maximal concentration of HPO₄² equivalent to the minimal buffer strength in the investigated pH range (Fig. S12). The buffer strength is the ability to compensate for increasing H⁺ concentration by chemical binding of the proton to a base. At pH 20 values above the 2nd phosphate pK (p $K_{2nd} \approx 7.2$), the buffer strength of the HPO₄²⁻/H₂PO₄¹⁻ ion decreases. At higher pH, the hydroxyl ions can significantly contribute to the proton uptake and the buffer strength increases. Both effects are reflected in the η-pH curves of both oxides. The beneficial influence of the 25 phosphate buffer is not related to an increase of the proton conductivity of the bulk solution. Though there are changes in the conductivity of the electrolyte (Fig. S11a), the iR-correction does not change the η -pH profile significantly (Fig. S11b). The chemical proton abstraction from the active site by phosphate 30 ions either adsorbed to the MnO_x or in solution within the Helmholtz layer appears to be decisive. Thus, Mn-based catalyst materials containing sizeable amounts of a proton-accepting base either as an intrinsic constituent - as reported for the 'cobaltphosphate catalyst⁹ - or formed during operation may be a 35 promising strategy.

Conclusions

We have uncovered the dominant atomic-scale structural motifs of two distinctly different, either layered or 3D cross-linked, water-oxidizing MnOx catalysts, and succeeded in correlating 40 these structure motifs with the respective catalytic activity. Such relations are of general importance, since they enable us to identify critical oxide characteristics controlling electrocatalytic activity.

- Our investigation suggest that the deliberate formation of activity controlling structural motifs can be controlled by simple conventional liquid metal ion precursor chemistry and is not restricted to electrodeposition methods.
- The electrokinetic analysis showed that the apparent processes during water oxidation depend on the different octahedral crosslinking in the MnO_x (with essentially the same mean oxidation state of the Mn ions), as clearly reflected in the two pronouncedly different Tafel slopes. The comparison to the structural data 55 suggests that the atomic structure is a crucial determinant of the apparent processes during OER catalysis. Comparison with electrodeposited electrocatalysts guides towards a general

character of MnO_x OER electrocatalysts.

The active catalyst state of both MnO_x is closely related to the catalyst "precursor" although the mean Mn oxidation state increases from 3.5 to ~4 upon operation at catalytic potential. Specifically, clear indications for Mn-Mn bridging mode changes in i-MnO_x upon oxidation of the catalyst are detectable. This 65 important discovery requires further in-depth investigation before its discussion in terms of the mechanism of water oxidation in the Mn oxides can be approached. We also note the interesting structural analogy between out-of-layer Mn ions linked to inlayer Mn ions via mono- u-oxo bridges resulting in corner-70 sharing MnOx octahedra and co-ordinately under-saturated (cus) metal ion sites in rutile surfaces. It is feasible that coordinative undersaturation of such out-of-layer Mn ions may confer favorable adsorption and catalytic reaction characteristics as found for rutile cus sites^{61, 62}.

Furthermore, our studies unravelled the beneficial effect of the proton uptake capability of the phosphate buffer during water splitting due to buffer strength and OH concentration. These findings – and the complex pH dependence in general – illustrate 80 the importance of proton abstraction in OER catalysis by the two herein investigated oxides. However, further studies especially on the interaction between phosphate and the MnO_x with respect to OER are necessary.

In summary, the structure-activity insights of this study provide guidelines to design and control the electrochemical OER behaviour of the complex structural chemistry of non-precious transition metal oxides. The water accessibility of the bulk oxide and the number of defect sites were tentatively identified as 90 determinants of the experimental exchange current density and Tafel slope, respectively. The presented work may pave the road for future investigations which eventually could result in a comprehensive understanding of structure-activity relations in water-oxidizing Mn oxides at the atomistic level.

95 Acknowledgements

We thank K. Mette (FHI) for collaboration regarding the synthesis, Dr. P. Chernev, Dr. M. Risch and J. Heidkamp (Freie Universität Berlin) for contributing to data collection at the synchrotron radiation source as well as M. Mertin and Dr F. 100 Schäfers for their excellent technical support at the beamline KMC-1 of the BESSY, a synchrotron radiation source in Berlin operated by the Helmholtz-Zentrum Berlin. Financial support by the Berlin cluster of excellence on Unifying Concepts in Catalysis (UniCat) is gratefully acknowledged. Arno Bergmann 105 acknowledges financial support by the Berlin Graduate school of Natural Sciences and Engineering (BIG-NSE).

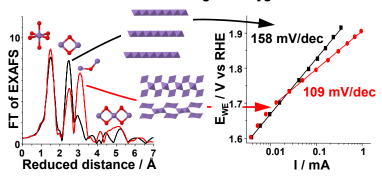
		30.	Y. Chabre and J. Pannetier, <i>Progress in Solid State Chemistry</i> ,
References		31.	1995, 23 , 1-130. J. H. Albering, in <i>Handbook of Battery Materials</i> , ed. J. O.
1.	J. O. M. Bockris, International Journal of Hydrogen Energy,		Besenhard, Wiley-VCH Verlag GmbH, 1999, pp. 85-112.
2.	1999, 27 , 731-740. T. A. Faunce, S. Styring, M. Wasielewski, G. Brudvig, B.	75 32.	J. E. Post, <i>Proceedings of the National Academy of Sciences</i> , 1999, 96 , 3447-3454.
5	Rutherford, J. Messinger, A. F. Lee, C. L. Hill, H. de Groot,	33.	P. de Wolff, Acta Crystallographica, 1959, 12, 341-345.
	M. Fontecave, D. MacFarlane, B. Hankamer, D. G. Nocera, D.	34.	S. Turner and P. R. Buseck, <i>Nature</i> , 1983, 304 , 143-146.
	M. Tiede, H. Dau, W. Hillier and L. Wang, Energy &	35.	P. Rüetschi, Journal of The Electrochemical Society, 1984, 131, 2737-2744.
2	Environmental Science, 2013.	⁸⁰ 36.	P. Rüetschi, Journal of The Electrochemical Society, 1988,
3.	H. Dau, C. Limberg, T. Reier, M. Risch, S. Roggan and P. Strasser, <i>ChemCatChem</i> , 2010, 2 , 724-761.	50.	135, 2657-2663.
¹⁰ 4.	T. Reier, M. Oezaslan and P. Strasser, <i>Acs Catal</i> , 2012, 2 ,	37.	P. Rüetschi and R. Giovanoli, Journal of The Electrochemical
••	1765-1772.		Society, 1988, 135 , 2663-2669.
5.	R. Forgie, G. Bugosh, K. C. Neyerlin, Z. Liu and P. Strasser, Electrochemical and Solid-State Letters, 2010, 13, B36-B39.	85 38.	Y. Mo, Y. Hu, I. T. Bae, B. Miller, M. R. Antonio and D. A. Scherson, <i>Journal of the Electrochemical Society</i> , 1997, 144 ,
15 6.	K. C. Neyerlin, G. Bugosh, R. Forgie, Z. Liu and P. Strasser,	20	1598.
	Journal of The Electrochemical Society, 2009, 156 , B363-	39.	H. Dau, I. Zaharieva and M. Haumann, <i>Current Opinion in Chemical Biology</i> , 2012, 16 , 3-10.
7.	B369. N. Menzel, E. Ortel, R. Kraehnert and P. Strasser,	90 40.	M. M. Najafpour, T. Ehrenberg, M. Wiechen and P. Kurz,
7.	ChemPhysChem, 2012, 13 , 1385-1394.		Angewandte Chemie, 2010, 122 , 2281-2285.
20 8.	E. Ortel, T. Reier, P. Strasser and R. Kraehnert, <i>Chemistry of</i>	41.	R. K. Hocking, R. Brimblecombe, LY. Chang, A. Singh, M.
	Materials, 2011, 23 , 3201-3209.		H. Cheah, C. Glover, W. H. Casey and L. Spiccia, Nature
9.	M. W. Kanan and D. G. Nocera, Science, 2008, 321, 1072-	40	Chemistry, 2011, 3 , 461-466.
	1075.	95 42.	R. Brimblecombe, A. Koo, G. C. Dismukes, G. F. Swiegers
10.	A. J. Esswein, Y. Surendranath, S. Y. Reece and D. G. Nocera,		and L. Spiccia, Journal of the American Chemical Society, 2010, 132, 2892-2894.
25	Energy & Environmental Science, 2010, 4, 499-504.	43.	T. Takashima, K. Hashimoto and R. Nakamura, <i>Journal of the</i>
11.	J. B. Gerken, J. G. McAlpin, J. Y. C. Chen, M. L. Rigsby, W. H. Casey, R. D. Britt and S. S. Stahl, <i>Journal of the American</i>	15.	American Chemical Society, 2012, 134 , 1519-1527.
	Chemical Society, 2011, 133 , 14431-14442.	100 44.	R. Alessio, D. B. Dell'Amico, F. Calderazzo, U. Englert, A.
12.	M. Risch, V. Khare, I. Zaharieva, L. Gerencser, P. Chernev		Guarini, L. Labella and P. Strasser, Helv Chim Acta, 1998, 81,
30	and H. Dau, Journal of the American Chemical Society, 2009,		219-230.
	131 , 6936-6937.	45.	M. Eiswirth, J. Bürger, P. Strasser and G. Ertl, <i>J. Phys. Chem.</i> ,
13.	Y. Surendranath, M. W. Kanan and D. G. Nocera, <i>Journal of</i>	105 46.	1996, 100 , 19118. Z. Liu, C. Yu, I. Rusakova, D. Huang and P. Strasser, <i>Top</i> .
1.4	the American Chemical Society, 2010, 132 , 16501-16509.	103 40.	Catal., 2008, 49 , 241-250.
14.	M. W. Kanan, J. Yano, Y. Surendranath, M. Dincâ, V. K. Yachandra and D. G. Nocera, <i>Journal of the American</i>	47.	X. Xie and L. Gao, <i>Carbon</i> , 2007, 45 , 2365-2373.
35	Chemical Society, 2010, 132 , 13692-13701.	48.	J. E. Penner-Hahn, Coordination Chemistry Reviews, 1999,
15.	M. Risch, K. Klingan, F. Ringleb, P. Chernev, I. Zaharieva, A.		190-192 , 1101-1123.
	Fischer and H. Dau, <i>ChemSusChem</i> , 2012, 5 , 542-549.	110 49.	H. Dau, P. Liebisch and M. Haumann, <i>Analytical and</i>
16.	M. Dincâ, Y. Surendranath and D. G. Nocera, <i>Proceedings of</i>	50	Bioanalytical Chemistry, 2003, 376, 562-583.
40	the National Academy of Sciences, 2010, 107 , 10337-10341.	50.	J. G. McAlpin, Y. Surendranath, M. Dincă, T. A. Stich, S. A. Stoian, W. H. Casey, D. G. Nocera and R. D. Britt, <i>Journal of</i>
17.	M. Risch, K. Klingan, J. Heidkamp, D. Ehrenberg, P. Chernev, I. Zaharieva and H. Dau, <i>Chemical Communications</i> , 2011,		the American Chemical Society, 2010, 132, 6882-6883.
	47, 11912-11914.	115 51.	D. K. Bediako, B. Lassalle-Kaiser, Y. Surendranath, J. Yano,
18.	J. O. Bockris and T. Otagawa, <i>The Journal of Physical</i>		V. K. Yachandra and D. G. Nocera, Journal of the American
45	Chemistry, 1983, 87 , 2960-2971.		Chemical Society, 2012, 134 , 6801-6809.
19.	P. C. K. Vesborg and T. F. Jaramillo, RSC Advances, 2012, 2,	52.	AC. Gaillot, D. Flot, V. A. Drits, A. Manceau, M.
20	7933-7947.	120	Burghammer and B. Lanson, <i>Chem. Mater.</i> , 2003, 15 , 4666-4678.
20.	Y. Gorlin and T. F. Jaramillo, <i>Journal of the American Chemical Society</i> , 2010, 132 , 13612-13614.	¹²⁰ 53.	T. G. Spiro, J. R. Bargar, G. Sposito and B. M. Tebo, <i>Accounts</i>
50 21.	I. Zaharieva, P. Chernev, M. Risch, K. Klingan, M. Kohlhoff,	00.	of Chemical Research, 2010, 43, 2-9.
J. 21.	A. Fischer and H. Dau, <i>Energy & Environmental Science</i> ,	54.	S. M. Webb, B. M. Tebo and J. R. Bargar, Am. Mineral., 2005,
	2012, 5 , 7081-7089.		90, 1342–1357.
22.	I. Zaharieva, M. M. Najafpour, M. Wiechen, M. Haumann, P.	125 55.	I. Saratovsky, P. G. Wightman, P. A. Pasten, JF. Gaillard and
	Kurz and H. Dau, Energy & Environmental Science, 2011, 4,		K. R. Poeppelmeier, <i>J. Am. Chem. Soc.</i> , 2006, 128 , 11188-11198.
55	2400-2408.	56.	S. M. Webb, B. M. Tebo and J. R. Bargar, <i>American</i>
23.	V. B. R. Boppana and F. Jiao, <i>Chemical Communications</i> , 2011, 47 , 8973-8975.	00.	Mineralogist, 2005, 90 , 1342-1357.
24.	K. Mette, A. Bergmann, JP. Tessonnier, M. Hävecker, L.	130 57.	S. Geller, Acta Crystallographica Section B, 1971, 27, 821-
	Yao, T. Ressler, R. Schlögl, P. Strasser and M. Behrens,		828.
60	ChemCatChem, 2012, 4, 851-862.	58.	R. E. Pacalo and E. K. Graham, Physics and Chemistry of
25.	F. Jiao and H. Frei, Chemical Communications, 2010, 46,	50	Minerals, 1991, 18 , 69-80.
2.0	2920-2922.	59.	D. R. Lide, ed., <i>Handbook of Chemistry and Physics</i> , CRC-
26.	M. Morita, C. Iwakura and H. Tamura, <i>Electrochim Acta</i> ,	60.	Press, 2001. O. Ghodbane, JL. Pascal and F. Favier, ACS Applied
65 27.	1977, 22 , 325-328. M. Morita, C. Iwakura and H. Tamura, <i>Electrochim Acta</i> ,	00.	Materials & Interfaces, 2009, 1, 1130-1139.
0.5 41.	1979, 24 , 639-643.	61.	H. Over, <i>Chem Rev</i> , 2012, 112 , 3356-3426.
28.	M. Morita, C. Iwakura and H. Tamura, <i>Electrochim Acta</i> ,	62.	H. Over, Electrochim Acta, 2013, 93, 314-333.
	1979, 24 , 357-362.	140	
29.	M. Wiechen, I. Zaharieva, H. Dau and P. Kurz, <i>Chemical</i>		
70	Science, 2012, 3 , 2330-2339.		

Broader Context

The intermittency of the electrical output power of renewable solar and wind energy devices makes efficient, cost-effective, grid-scale energy storage mandatory. Electrical power storage using molecular bonds (synthesis of fuel molecules) offers a number of advantages over storage in solid state material lattices. As electrolytic synthesis of chemical fuels is a reductive chemical process, a corresponding electrooxidation process at grid-scale level is essential. Electrochemical water oxidation to molecular oxygen is perhaps the only feasible oxidation process that can provide electrons and protons at the required grid-scale. However, inexpensive, efficient and stable water oxidation catalysts are rare, if non-existent. This is why the development of non precious water oxidation catalysts has become a scientific priority. Manganese oxides enjoy special attention as water splitters, as they form the basis for Nature's oxygen evolving complex in the Photosystem II complex. Detailed insight into structure-activity relations of inorganic Mn containing water splitting materials are still scarce. This article address this unmet need.

TOC figure:

Mn octahedral cross-linking and oxygen evolution



Electronic Supplementary Information

Electrochemical Water Splitting by Layered and 3D Cross-linked Manganese Oxides: Correlating Structural Motifs and Catalytic Activity

Arno Bergmann,*^a Ivelina Zaharieva,*^b Holger Dau^b and Peter Strasser^a

^a Department of Chemistry, Chemical and Materials Engineering Division, The Electrochemical Energy, Catalysis and Materials Science Laboratory, Technische Universität Berlin, Straße des 17. Juni 124, 10623 Berlin, Germany.

E-Mail: bergmann@chem.tu-berlin.de; pstrasser@tu-berlin.de

Fax: +49 30-314 22261; Tel: +49 30-314 29542;

^b Fachbereich Physik, Freie Universität Berlin, Arnimallee 14, 14195 Berlin, Germany.

E-Mail: ivelina.zaharieva@fu-berlin.de; holger.dau@fu-berlin.de;

Fax: +49 30 838 56299; Tel: +49 30 838 3581

Experimental

S-1. Details of experimental procedures

Syntheses

Carbon-supported manganese oxide has been synthesized via two different preparation methods – symproportionation deposition-precipitation (s-MnO $_x$) and incipient wetness impregnation (i-MnO $_x$). In both cases multi-walled carbon nanotubes (MWNT, Bayer MaterialScience AG, 95 % C) has been pre-treated with concentrated HNO $_3$ at 100°C for 16 h to remove the residual growth catalyst and amorphous carbon impurities and to functionalize their inner and outer surfaces by oxidation.

s-MnO_x has been prepared via impregnation of the carbon support with Mn²⁺ from 0.015 M manganese(II) nitrate tetrahydrate/ 0.09 M ammonium chloride solution $(Mn(NO_3)_2 \cdot 4H_2O, Merck)$ and drop wise addition of an 0.01 M aqueous potassium permanganate solution $(KMnO_4, Roth)$ under vigorous stirring. A constant pH of 8 was ensured via titration with ammonia. Afterwards, the compound was thoroughly washed, filtered and dried in air at 110 °C for 16 h.

i-MnO_x has been prepared via incipient wetness impregnation of the carbon support with a nominal loading of 10 wt.% of Mn^{2+} ions in an aqueous $Mn(NO_3)_2$ solution. The compound was consecutively dried in air for 6 h and 24 h at 60 °C and 110 °C, respectively.

Determination of the manganese loading of the carbon nanotubes was performed using a 715-ES-inductively coupled plasma (ICP) analysis system (Varian). Therefore, the powder samples were dissolved in aqua regia over night, the remaining carbon filtrated and the solution diluted with MilliQ water to the desired Mn concentration.

Calculation of overpotential n

The overpotential η in Fig. 8 has been calculated via

$$\eta = E_{NHE} + \frac{59mV}{pH \text{ unit}} \cdot pH - 1.23 \text{ V}$$

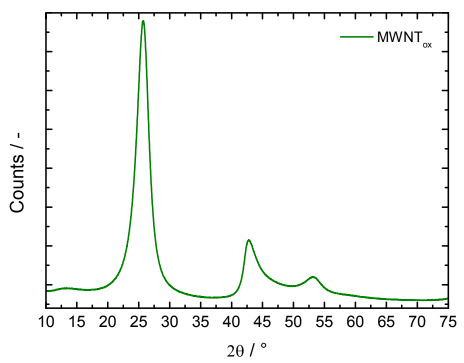


Fig. S1. Powder x-ray diffraction pattern of MWNT_{ox} between 10° and 75° at Cu K_{α} radiation.

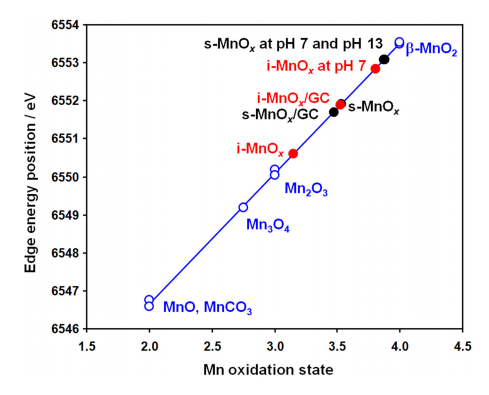


Fig. S2. Calibration line based on Mn compounds with known structure and oxidation state (in blue). Black symbols and labels indicate the edge position and estimated mean Mn oxidation state for s-MnO_x powder before (s-MnO_x) and after (s-MnO_x/GC) deposition on the electrode, and after electrocatalytic operation at 1.763 V vs. RHE for 3 min at pH 7 (0.1 M KPi) or pH 13 (0.1 M KOH). Red symbols and label represent the corresponding values for the i-MnOx.

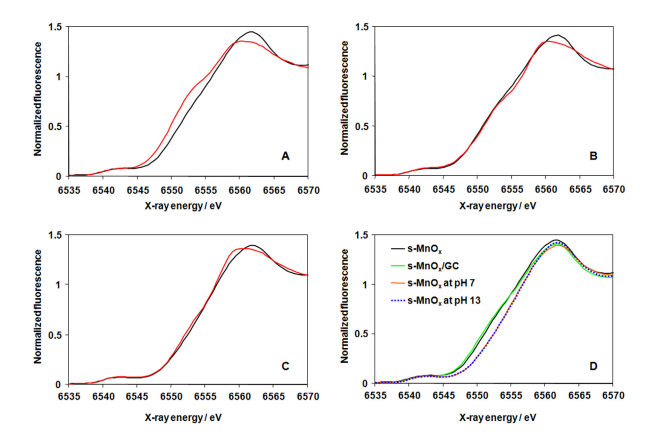


Fig. S3. XANES spectra of s-MnO_x (in black) and i-MnO_x (in red) after synthesis (A), after deposition on GC electrode (B), and after operating for 3 min at 1.763 V vs. RHE in 0.1 M KPi, pH 7 (C). In (D) is shown a comparison of the XANES spectra recorded from s-MnO_x at different stages of catalyst preparation.

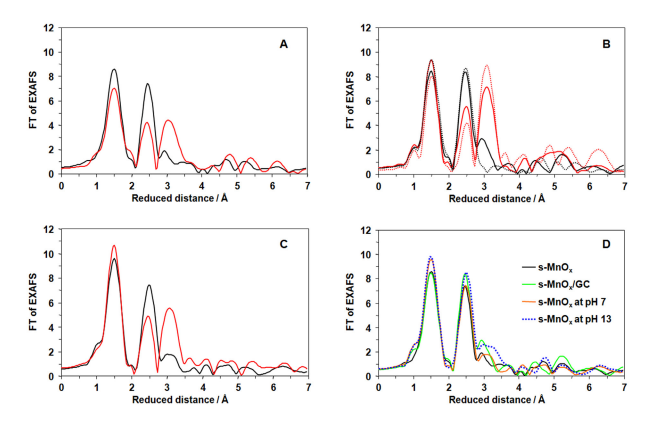


Fig. S4. Fourier-transformed EXAFS spectra of s-MnO_x (black solid lines) and i-MnO_x (red solid lines) after synthesis (A), after deposition on GC electrode (B), and after operating for 3 min at 1.763 V vs. RHE in 0.1 M phosphate buffer, pH 7 (C). In (B) the spectra from δ-MnO₂ and β-MnO₂ reference compounds are shown using black and red dotted lines, respectively. In (D) a comparison of the Fourier-transformed EXAFS spectra recorded from s-MnO_x at different stages of catalyst preparation is shown.

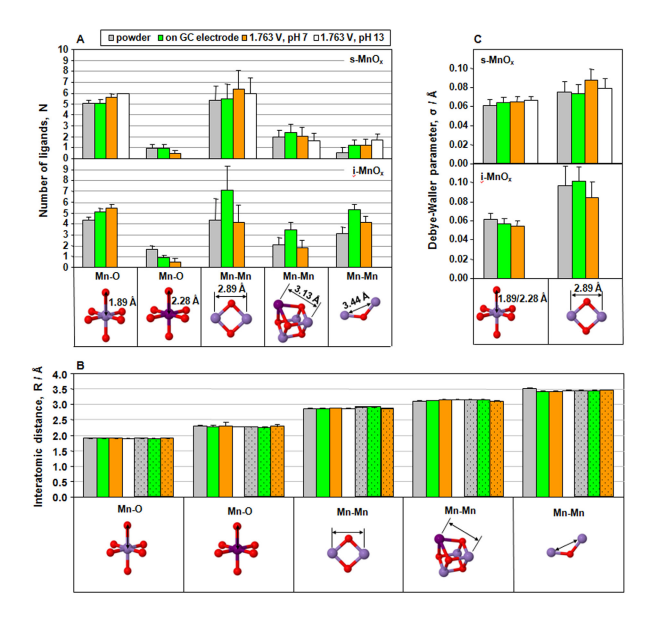


Fig. S5. Simulation results for the independent fits of the k^3 -weighted EXAFS spectra from s-MnO_x and i-MnO_x before and after operation as water oxidation catalysts in 0.1 M KPi (pH 7) or 0.1 M KOH (pH 13). (A) – EXAFS coordination numbers, (B) – interatomic distances, (C) – Debye-Waller parameters. During the simulation, the sum of the coordination numbers for the two oxygen shells was kept equal to 6 and the same Debye-Waller parameter was used for each of the two oxygen shells. The Debye-Waller parameters for the last two Mn-Mn shells were fixed to 0.063 Å. The errors represent a 68% confidence interval.



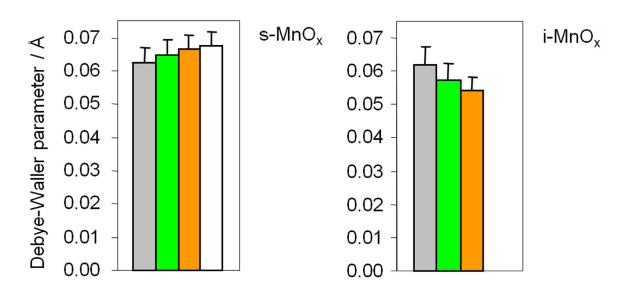


Fig. S6. Debye-Waller parameters for the oxygen shell of backscattering atoms. The other fit parameters are presented in Fig. 3 in the main text.

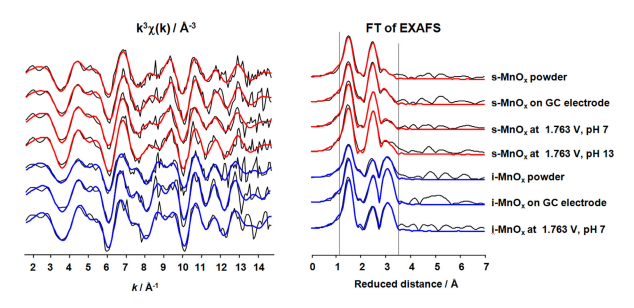


Fig. S7. k^3 -weighted experimental EXAFS spectra of the studied Mn oxides (thin black lines) and the joint-fit simulation result (red lines for s-MnO_x, blue for the i-MnO_x). The fit parameters are given in Fig. 3 of the main manuscript. The grey vertical lines indicate the range used to calculate the RF error factor. The averaged RF-values equalled 15.1% for the joint-fit simulation approach and 13.6% for simulations with independently variable interatomic distances.

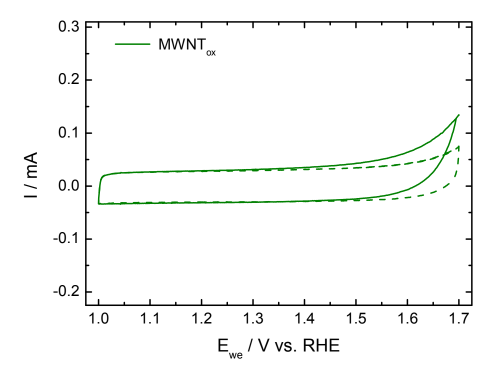
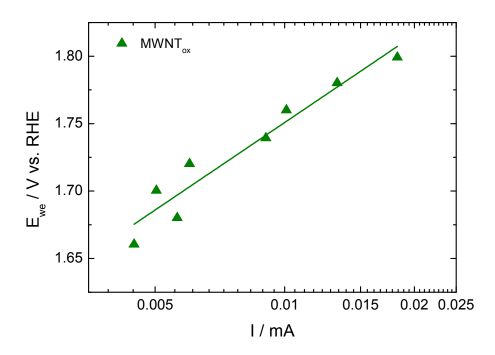


Fig. S8. First (solid) and 11th (dashed) voltammogram of MWNT_{ox} (without catalyst) recorded in 0.1 M KPi at pH 7 with a scan rate of 100 mV s⁻¹.



Background current of MWNT_{ox} without deposited catalysts. The Tafel plot of MWNT_{ox} in 0.1 M KPi at pH 7 was extracted from quasi-stationary potential-step RDE experiments at a rotation speed of 1600 rpm. All electrode potentials have been *iR*-corrected. We note that the current values are significantly lower than the corresponding figures in the presence of the investigated Mn oxide catalysts. The Tafel slope was determined to 216 mV dec⁻¹ from linear regression between 1.675 and 1.8 V.

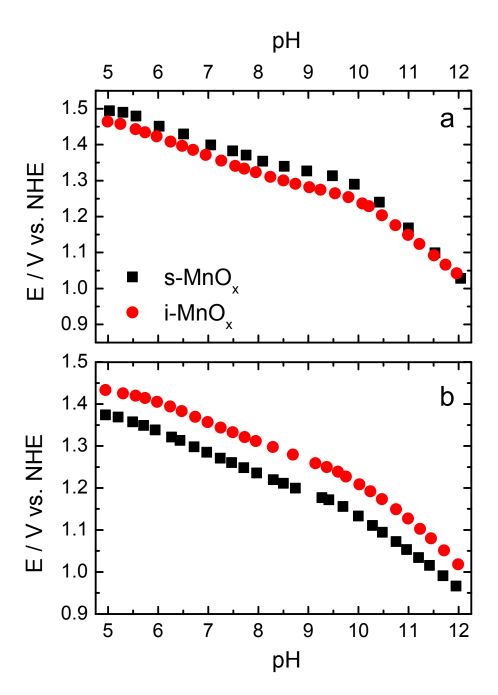


Fig. S10 pH dependence of the electrode potential vs. NHE at a constant current of (a) $50 \,\mu\text{A}$ and (b) $20 \,\mu\text{A}$ recorded during titration of aliquots of 40 wt.% KOH into 0.1 M KPi increasing the pH between 5 and 12. At each pH values the corresponding potential was detected after equilibration for 3 min.

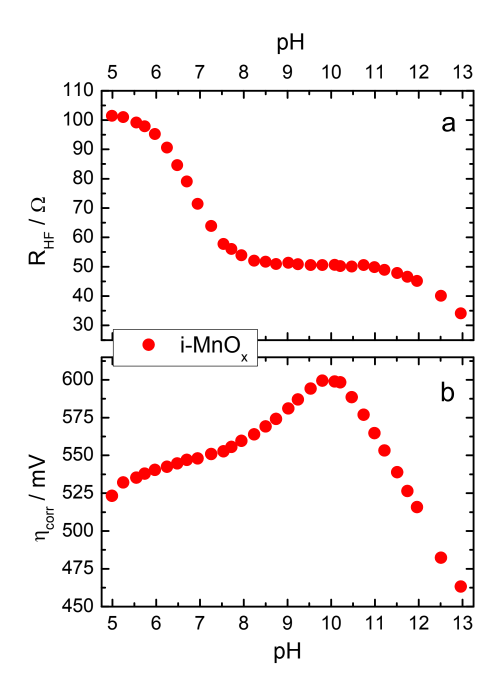


Fig. S11. pH dependence of electrolyte resistance (a) and η -pH profile after correction for ohmic losses (*iR* correction) (b). In (a), the high frequency resistance R_{HF} of the bulk electrolyte has been determined via electrochemical impedance spectroscopy at 1.35 V vs. NHE. The pH was adjusted via titration of aliquots of 40 wt.% KOH.

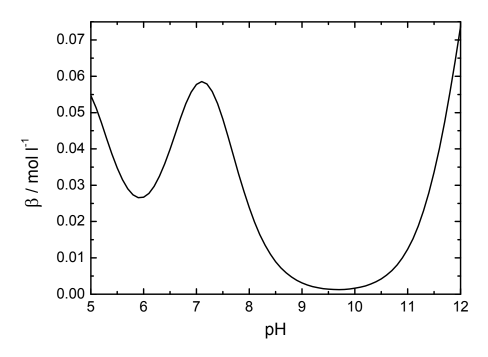


Fig. S12. pH dependence of the calculated buffer strength (β). For a concentration of the potassium phosphate buffer of 0.1 M, the buffer strength was calculated as described in ref.¹

1. A. Hulanicki, Reactions of acids and bases in analytical chemistry, Ellis Horwood Ltd, 1987.