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Redox Processes of Manganese Oxide in Catalyzing Oxygen Evolution and Reduction: An in Situ Soft X-ray Absorption Spectroscopy Study

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Supporting Information

ABSTRACT: Manganese oxides with rich redox chemistry have been widely used in (electro)catalysis in applications of energy and environmental consequence. While they are ubiquitous in catalyzing the oxygen evolution reaction (OER) and oxygen reduction reaction (ORR), redox processes occurring on the surface of manganese oxides are poorly understood. We report valence changes at OER- and ORR-relevant voltages of a layered manganese oxide film prepared by electrodeposition. X-ray absorption spectra were collected in situ in O2-saturated 0.1 M KOH using inverse partial fluorescence yield (IPFY) at the Mn L_{32} -edges and partial fluorescence yield (PFY) at the O K-edge. Overall, we found reversible yet hysteretic Mn redox and



qualitatively reproducible spectral changes by Mn $L_{3,2}$ IPFY XAS. Oxidation to a mixed Mn^{3+/4+} valence preceded the oxygen evolution at 1.65 V vs RHE, while manganese reduced below Mn³⁺ and contained tetrahedral Mn²⁺ during oxygen reduction at 0.5 V vs RHE. Analysis of the pre-edge in O K-edge XAS provided the Mn–O hybridization, which was highest for Mn³⁺ (e_{σ}^{-1}). Our study demonstrates that combined in situ experiments at the metal L- and oxygen K-edges are indispensable to identify both the active valence during catalysis and the hybridization with oxygen adsorbates, critical to the rational design of active catalysts for oxygen electrocatalysis.

INTRODUCTION

Manganese oxides have rich redox chemistry and are thus a prime choice for energy storage and conversion both in nature and industrial applications. In nature, oxygen evolution is performed by a CaMn₄O₅ cluster in photosystem II (PSII),^{1,2} which has inspired much research in homogeneous and heterogeneous (photo)electrocatalysis of the oxygen evolution reaction (OER). $^{\overline{2}-9}$ The CaMn₄O₅ motif can also be found in several layered oxides,^{10–14} among which are layered birnessites $(\delta - A_x MnO_{2-y} \cdot zH_2O, A = Na, K, Ca, and Sr).^{10-13,15-18}$ In addition, Mn oxides with perovskite,^{19,20} bixbyite,²¹ and spinel²² structures can exhibit reasonable activity in catalyzing OER. Moreover, Mn oxides are among the most active for the oxygen reduction reaction (ORR),^{23–26} where the ones with perovskite,²⁷ bixbyite,²⁸ and spinel^{22,23,29,30} structures show high ORR activity. Due to suitable kinetics for both OER and ORR, Mn oxides are studied as bifunctional electrocatalysts in metal-air batteries.³¹⁻³³ Furthermore, their redox chemistry

makes Mn oxides promising candidates for charge storage in electrochemical capacitors^{34–36} and batteries^{37–39}

Mn oxides form layered, spinel, perovskite, and bixbyite structures, where Mn ions can have a wide range of valence states usually from 2+ to 4+,^{10,12,16,18,20-22,31,40-52} exhibiting different catalytic activities for OER and ORR kinetics.^{21,23,45,5} A number of studies have employed ex situ methods to correlate Mn valence with OER and ORR activity, using either X-ray absorption spectroscopy (XAS) at the Mn K-edge (hard XAS)^{10,15,22,40-42,54,55} to probe unoccupied Mn-4p orbitals or the Mn $L_{3,2}$ -edges (soft XAS)⁴³⁻⁴⁶ to probe unoccupied Mn-3d orbitals or by using X-ray photoelectron spectroscopy (XPS)^{16,20,21,31,41,43} to probe binding energies of occupied states (e.g., of Mn 2p electrons). However, ambiguities exist in

Received: June 7, 2017 Revised: July 23, 2017 Published: July 23, 2017 what Mn valence is responsible for OER or ORR activity among the *ex situ* studies in literature.

Electrodeposited and subsequently crystallized films of Mn³⁺₂O₃ (with minor Mn²⁺ impurities) show OER activity in 1.0 M KOH higher than those containing more Mn²⁺ or Mn⁴⁺ from ex situ XPS work.²¹ This study detects little Mn⁴⁺ after the oxidizing conditions of the OER.²¹ Moreover, a recent study by Wei et al.²² shows increasing activity of MnCo₂O₄ in 0.1 M KOH with reduction from Mn^{3.7+} to Mn^{3.2+} based on ex situ Kedge XAS. The highest OER activity (1.52 V vs RHE at 25 μ A/ cm²_{ox}) is observed for Mn₃O₄,²² where it is argued that octahedral Mn³⁺ is the active site. However, an optimal Mn valence in the range of Mn^{3.6+} to Mn^{3.8+} for the OER on various Mn oxides in 0.1 M KPi by ex situ XPS¹⁶ and ex situ K-edge XAS^{10,15} studies, where the catalyst is composed exclusively of Mn octahedra. Higher valences approaching Mn⁴⁺ have also been detected for these materials after the OER in both 0.1 M KOH and KPi by ex situ K-edge XAS.¹⁴ On the basis of ex situ L-edge XAS, Gorlin et al.43 find that the extent of valence changes after OER in 0.1 M KOH depends on the crystallinity that can be greatly influenced by the heat treatment in the preparation of the catalyst.

With regard to ORR kinetics, Mn perovskites with mixed valence of Mn^{3+} and Mn^{4+} have intrinsic activities in 0.1 M KOH of up to 1.3 mA/cm²_{ox} at 0.8 V vs RHE,^{23,27} which is higher than those containing exclusively Mn^{4+} as measured by *ex situ* L-edge⁴⁶ and K-edge^{22,41} XAS, which suggests that having Mn^{3+} is critical to obtain high activity.⁴⁶ In addition, spinels containing Mn^{3+} and Mn^{2+} also exhibit high activities of up to 0.7 mA/cm²_{ox} at 0.8 V vs RHE for the ORR in 0.1 M KOH,^{23,29,30} corroborating that Mn^{3+} is essential to catalyze the ORR. It has recently been shown that the e_g occupation of the octahedral site in these spinels is the activity descriptor for ORR,²² which shows the maximum activity (0.88 V vs RHE at 25 μ A/cm²_{ox}) for Mn^{3.4+}. This is the same Mn valence as reported for the most active perovskite oxide in the La_{1-x}Sr_xMnO₃ series.²³ Thus, these observations are in agreement with the hypothesis that having a mixed Mn valence between Mn³⁺ and Mn⁴⁺ can improve the ORR activity.²³

The dynamic range of accessible Mn valence states near the potentials of oxygen evolution and reduction (O_2/OH^-) suggests the valence may change during these electrocatalytic processes, which calls for detection and correlation of potentialdependent Mn valence by in situ characterization. Such studies have been performed using hard X-rays at the Mn K-edge.^{12,18,42,47–50,55,56} Mn valence changes can also be tracked using UV-vis spectroscopy after suitable calibration with another method.^{12,15,55,57,58} Zaharieva et al.¹² report an oxidation of up to 0.5 Mn valence states before the onset of OER in an *in situ* experiment in 0.1 M KPi, where Mn³⁺ at catalytic voltages is proposed as a prerequisite for OER activity in electrodeposited layered Mn oxides. Similar observations are noted by Gorlin et al.,¹⁸ where oxidation of electrodeposited Mn oxide films during the OER at 1.8 V vs RHE in 0.1 M KOH result in a mixed $Mn^{3+/4+}$ oxide with layered structure. Oxidation of electrodeposited^{51,59,60} and photodeposited⁵⁰ Mn oxide prior to the OER is also reported in near-neutral solutions of 0.1 M KPi by Mn $L_{3,2}$ -edge XAS, ^{51,60} 0.1 M Na₂SO₄ by Mn K-edge XAS,⁵⁰ and 2 M KCl by Mn K-edge XAS.⁵⁹ Addition of Au to the electrode increases the Mn valence of reactively sputtered Mn₃O₄ films as compared to identical potentials on Au-free electrodes.45

With regard to ORR, *in situ* Mn K-edge work of Lima et al.⁴⁷ reveals the reduction of β -MnO₂ and the formation of Mn₃O₄ (with tetrahedral Mn²⁺ and octahedral Mn³⁺) at 0.70 V vs RHE in 1.0 M KOH, which is in agreement with the work of Gorlin et al.¹⁸ finding the formation of Mn₃O₄ from electrodeposited Mn oxide during ORR at 0.7 V vs RHE in 0.1 M KOH. At higher ORR overpotentials (~0.3 V vs RHE), significant fractions of octahedral Mn²⁺ are observed (e.g., in the Mn(OH)₂ phase).^{47,48} Since only the valence of the octahedral sites was proposed to trend with activity,²² it is crucial to track both the oxygen coordination and valence of the active Mn cations during *in situ* experiments.

Extracting coordination and valence from soft XAS has several advantages over hard XAS. While hard XAS at the Mn K-edge probes Mn 4p orbitals,⁶¹ soft XAS at the Mn L_{3,2}-edges is ideally suited to elucidate the Mn valence and coordination symmetry because Mn 3d orbitals are probed directly.⁶² The core hole broadening at the Mn K-edge is 1.2 eV, while it is only 0.3 eV at the Mn L₃-edge.⁶³ This allows resolution of more fine structure at the Mn L3-edge so that Mn valence and coordination in mixed valence oxides can be better distinguished. It also makes in situ studies at the Mn L_{3,2}edges an attractive alternative to time-consuming analysis required for extended X-ray absorption fine structure (EXAFS) studies. Furthermore, this information is contained in an energy range of 10 eV at the Mn L3-edge, while an energy range of about 550–975 eV ($k_{\text{max}} = 12-16 \text{ Å}^{-1}$) is commonly recorded for EXAFS.⁶⁴ Thus, higher time resolution is achievable by in situ experiments at the Mn L₃-edge due to the shorter energy range and therefore acquisition time. In addition, the approximate attenuation lengths of the X-rays are a factor 100 shorter at the Mn L_3 -edge (~100 nm) as compared to the Mn K-edge (~10 μ m).⁶⁵ The attenuation length at the Mn L₃edge is thus better matched with typical thicknesses of electrodeposited films for in situ XAS of 100 nm¹⁸ or less.⁶⁶ Unfortunately, there are very few demonstrations of Mn valence changes at the Mn L_{3,2}-edges^{51,60} due to the enormous challenges associated with in situ XAS of Mn at these low energies. These X-ray transmission studies demonstrated oxidation and subsequent reduction of an electrodeposited Mn oxide during a single voltage cycle⁵¹ and Mn reduction during potentiostatic activation.⁶⁰ The soft XAS regime also gives access to analysis of the O K-edge, which provides valuable complementary insight into the electronic structure but has never before been performed in situ for Mn oxides and is rarely performed for other transition metal oxides.^{42,67–69}

In this study, we report valence changes of an electrodeposited Mn oxide film in O₂-saturated 0.1 M KOH *in situ* as a function of voltage, extending from ORR to OER regions for multiple cycles. Mn $L_{3,2}$ spectra of the electrodeposited film provide high sensitivity to valence changes of Mn ions when detected using the inverse partial fluorescence yield (IPFY).⁷⁰ Using Mn $L_{3,2}$ IPFY XAS, we found overall reversible yet hysteretic Mn redox in a single cycle and qualitatively reproducible spectral changes during repeated voltage cycling. Lastly, the Mn–O hybridization was studied *in situ* by O Kpartial fluorescence yield (PFY) XAS. Combining Mn $L_{3,2}$ and O K-edge analysis yielded a more comprehensive picture of dynamic changes of both Mn valence and Mn–O hybridization as a function of potential, which is critical to gain insights into the active sites catalyzing ORR and OER.

EXPERIMENTAL SECTION

Materials. The following Mn oxides were obtained commercially as references: MnO (99.99%, Alfa Aesar), Mn₃O₄ (97%, Sigma-Aldrich), β -MnO₂ (99.999%, Alfa Aesar), and KMnO₄ (99%, Sigma-Aldrich). Additionally, δ -K_xMnO_{2-y}·zH₂O was synthesized by reacting MnSO₄ with KMnO₄ in aqueous solution.⁷¹ Transmission electron microscopy electron diffraction of reference δ -K_xMnO_{2-y}·zH₂O powder revealed a hexagonal layered structure (space group $P6_3/mmc$; Figure S1). Further details about the references may be found in Table S1.

The electrodeposited films were prepared following the work of Nakayama et al.,⁷² where a potential of 1.33 V vs RHE was applied for a controlled charge of 10.3 mC/cm² (inset of Figure S2). All areas in this work refer to the geometric surface area of the electrode that was accessible to the electrolyte. Electrolytes for electrodeposition were prepared by adding 50 mM KCl (99% Sigma-Aldrich) and 2 mM KMnO₄ (99% Sigma-Aldrich) to deionized water (>18 M Ω cm), which had a pH of 6.5 at room temperature. Complete removal of KMnO₄ from electrodeposition was confirmed by recording a Mn L_{3,2}-edge spectrum in an XAS cell filled with water (Figure S3), which exhibited no characteristic peaks of KMnO₄. While the crystal structures of these electrodeposited thin films were difficult to resolve by X-ray diffraction, thicker films made by the same protocol can be assigned to the birnessite structure.⁷²

XAS Cell. The three-electrode XAS cell used was developed based on a previously used flow cell.⁷³ It was made from Durus Black⁷³ with a leak-free silver chloride reference electrode (Harvard Apparatus) and a Pt counter electrode (Figure S4A). The XAS window was a commercial 20 nm Au on a 100 nm silicon nitride membrane with a 5 nm Cr binding layer (Norcada Inc. NX5100C).

Electrochemical Measurements. Electrolytes of 0.1 M KOH were prepared using deionized water (>18 M Ω cm) and KOH pellets with 99.98% purity (Alfa Aesar; at MIT) or 88.15% purity (Fisher Scientific; at the beamline). For activity measurements, oxygen (99.994% purity, Airgas) was bubbled to ensure O_2/HO^- equilibrium at 1.23 V vs RHE, while argon (99.999% purity, Airgas) was bubbled prior to background measurements for at least 30 min. Electrochemical characterization at the beamline was performed using a Voltalab PGZ-301 potentiostat. The reference electrode was calibrated in a laboratory glass cell using Pt working and counter electrodes in 0.1 M H₂-saturated KOH where the average voltage at zero current was taken as the reference voltage (1.006 V). The electrolyte was drawn through the flow cell using a Palmer syringe pump at a rate of 50 μ L/min (500 μ L/min for last 1.65 V step in cycle 3 due to O_2 buildup).

The electrodeposited δ -K_xMn^{3,4+}O_{2-y}·zH₂O on AulCrO_xI Si₃N₄ was electrochemically characterized in the three-electrode XAS cell, where we found the expected features in a cyclic voltammogram in O₂-saturated 0.1 M KOH (Figure S4B, step 6 in Table S2). The redox peaks centered at ~0.8 V vs RHE were previously observed for electrodeposited α -Mn₂O₃,⁷⁴ mixed MnO/MnO₂,¹⁸ and β -Mn^{IV}O₂⁴⁷ particles in both N₂- and O₂saturated 0.1 M KOH. During potentiostatic measurements, the currents at 0.50 V vs RHE were consistently negative owing to ORR while the currents at 1.65 V vs RHE were consistently positive owing to OER (Figure S4C). Currents collected at 1.20 and 1.50 V vs RHE had negative currents, which were not expected since the potentials were higher than the OCV of 1.07 V, indicating potential parasitic currents in the XAS cell. We can exclude pH changes in 0.1 M KOH for the observed currents (Figure S4C) due to high buffering capacity of water under these conditions (Supporting calculations).

X-ray Absorption Spectroscopy. X-ray absorption measurements at both the Mn L_{3,2}-edges and O K-edges were performed at the spherical grating monochromator (SGM) beamline 11ID-1 at the Canadian Light Source.75 The window of the sample cells was mounted at an angle of roughly 45° with respect to both the incident beam and the detectors. The irradiated area on the cell window was about 0.05 mm², as estimated previously.⁷³ All measurements were conducted at room temperature in the fluorescence mode using Amptek silicon drift detectors (SDDs) with 1024 emission channels (energy resolution \approx 120 eV). Four SDDs were employed simultaneously with vanadium (200 nm) filter foils mounted to suppress oxygen fluorescence. Gaussian curves were fitted to the oxygen and Mn emission lines of all SDDs at each incident energy where only the amplitude was varied freely. The incident intensity was obtained by measuring the current on a gold grid placed before the sample chamber.

PFY was then obtained from the Mn amplitude for Mn PFY in the range 632-676 eV and from the oxygen amplitude for oxygen PFY in the range 518-586 eV. Regardless of the range, the Mn L₃-edge intensity was reduced for Mn oxide powders due to self-absorption (Figure S5A). This makes it extremely challenging to distinguish fine details of the spectra around 640 eV. The IPFY was obtained from the inverse of the oxygen amplitude in the range 632-676 eV. In this range, the IPFY was free of self-absorption (Figure S5B) because the inverse of the nonresonant oxygen absorption is effectively a measure of the X-ray attenuation length.⁷⁰ Noting that there is no suitable Mn edge at energies lower than the O K-edge, we report the conventional PFY for the O K-edge. Oxygen PFY spectra were normalized for background absorption by fitting a straight line between 522 and 528 eV and subtracting it over the whole range of the data. Finally, the average intensity between 557 and 590 eV (after the O K-edge) was normalized to unity. The background of the in situ IPFY spectra was more complex due to the variation of the oxygen content (e.g., by electrochemical oxygen evolution). It required fitting an exponential function to the points of the local minimum between the L_3 - and L_2 -edge (around 650 eV) and after the L_2 -edge (660–676 eV), which was then subtracted over the whole range of data. This was followed by the conventional subtraction of a constant before the L₃-edge and division of the average intensity around 650 eV (after Mn L₃-edge). For consistency, the same normalization was applied to powder references.

The energy axes of both O K- and Mn $L_{3,2}$ -spectra were calibrated with respect to the pre-edge in the spectrum of molecular oxygen at 530.8 eV,⁷⁶ which was acquired by subtracting the spectrum of a pristine window in an oxygen atmosphere from that of a pristine window in an argon atmosphere (Figure S6). This differential approach was necessary because the pristine window contained a thin layer of chromium oxide to bind gold and silicon nitride. We use molecular oxygen as a reference for energy calibration because its purity is easier to ensure than that of metals which spontaneously form surface oxides. Moreover, evaluation is straightforward due to the very sharp prepeak of a molecular transition that can be calculated from first principles.⁷⁶ The energy positions of key spectral features of common oxides (suppliers in Table S1) with this energy calibration are

presented in Table S3 for the Mn L_3 -edge and in Table S4 for the O K-pre-edge. These common oxides could be used for energy calibration and thus allow comparison with the work presented herein. The peak positions on our scale differ from those reported previously (Table S5), which does not affect the calculation of the Mn valence that relies on relative energies.

All experiments were performed at a flux of $\sim 7 \times 10^9$ photons/s, for which there were no indications of radiation damage in potassium permanganate (or all other Mn oxides; Figure S5), which is known for facile photoreduction.⁷⁷ Moreover, we found no indications of radiation damage at this dose in subsequent scans during the *in situ* experiment where the electrolyte was flown at 50 μ L min⁻¹ (Figure S7). At this rate, the sample volume in the cell of $\sim 10 \ \mu$ L is completely replaced every 12 s. The sequence of all XAS measurements may be found in Table S2.

Linear Combination Analysis of the Mn L_{3,2}-edges. XANES fitting results depend critically on intensity normalization. To remove the effect of sample density variations and oxygen concentration, the Mn L-edge spectra were normalized to the average between 640 and 644 eV (Figure S8). Then, the rising part of the Mn L₃-edges was fit between 633.7 and 644.7 eV using Athena,⁷⁸ where a nonlinear least-squares procedure based on the Levenberg-Marquardt algorithm is employed. The XAS samples were fit using the spectra of $Mn^{2.7+}{}_{3}O_{4}$ and δ - $K_x Mn^{3.4+}O_{2-\nu}$; zH_2O powders as standards with the constraints that the weights of each standard is between 0 and 1 and the two weights sum to 1. Other standards, particularly MnO₂ polytypes, were tested but discarded due to worse fits. While δ - $K_x Mn^{3.4+}O_{2-v} \cdot zH_2O$ matched the oxidized state of the electrodeposited film very well (e.g., Figure S8A), Mn₃O₄ differed slightly from the reduced state of the electrodeposited film (e.g., Figure S8G). Thus, the necessity of additional standards cannot be rigorously excluded. However, the spectrum of Mn₃O₄ was the best match among Mn oxides with low oxidation state. State-dependent decay effects⁷⁹⁻⁸¹ might cause fluorescence yield measurements to deviate from true absorption cross sections of the Mn_3O_4 and δ -K_xMn^{3.4+}O_{2-v}· zH_2O phases. Fit uncertainties were obtained from the diagonal elements of the covariance matrix scaled by the square root of reduced chi-square (1σ - uncertainty).

Fitting and Integration of the O K-Pre-Edge Region. O K-edge spectra were fit by Gaussian curves using the "multiple peak fit" tool in Origin 9.1. For the analysis of the Mn–O hybridization of the reference samples (Figure S9), we simultaneously fit Gaussian curves to a shoulder and the main O K-edge between 536.2 and 547.4 eV as a background, except for MnO, where an additionally fitted peak at ~530 eV was included in the background. The Gaussian curves were not offset ($y_0 = 0$). We also attempted to fit the pre-edge region, but the fits did not converge for some of the reference materials. We thus used a more robust integration approach. Experimental uncertainties were obtained by performing the analysis on individual scans and calculation of the standard deviation of the values.

For the *in situ* measurements of the electrodeposited film, we simultaneously fit three Gaussians between 520.0 and 535.5 eV (Figures S10 and S11). Two of these peaks constitute the pre-edge, while the shoulder in the O K-edge was also included as a background to the pre-edge. The range from 530.5 to 531.5 eV was excluded because an additional peak may emerge in this range. The widths of the peak centered at ~530 and ~523 eV were fixed to 1.8 and 1.4 eV. The position of the peak in the O

K-edge was fixed to 533.0 eV. The Gaussian curves were not offset ($y_0 = 0$). All other parameters were allowed to vary freely. Fit uncertainties were obtained from the diagonal elements of the covariance matrix scaled by the square root of reduced chi-square (1σ - uncertainty).

RESULTS AND DISCUSSION

Calibration and Estimation of Mn Valence of Manganese Oxides Using IPFY XAS Spectra. Soft IPFY XAS spectra at the Mn L_{3,2}-edge of powder Mn oxides were used to estimate the Mn valence as they were less susceptible to saturation effects than the PFY (Figure S5) and more reliably calibrated with common reference materials as compared to the total electron yield (TEY; Figure S5C), both of which were used in previous *ex situ* work.^{43–46,61,82} These advantages of the IPFY resulted in the greatest sensitivity for Mn valence among the investigated methods. Figure 1A compares the IPFY spectra at the Mn L_{3,2}-edges of select phase-pure Mn oxide powders (Figure S12) with rocksalt, spinel, layered birnessite, and rutile structures (Figure 1B). These oxides have Mn valence in the range from 2+ to 4+ (Figure 1C). With increasing Mn valence, the centers of gravity of the Mn L₃-edges were found to linearly shift to higher energies. Of significance is that spinels of different Mn valence (i.e., $Mn^{2.7+}{}_{3}O_{4}$ and $LiMn^{3.5+}{}_{2}O_{4}$) have considerably different Mn L3-edges, while spinel LiMn^{3.5+}2O4 and layered δ -K_xMn^{3.5+}O_{2-y}·zH₂O with comparable valence also exhibit comparable spectra. A Mn oxide film was electrodeposited by reduction of KMnO₄ at 1.33 V vs RHE. Its Mn L₃-spectrum of was found to have a sharp feature at ~642 eV and a broad peak at ~645 eV, which resembles those of spinel LiMn^{3.5+} $_{2}O_{4}$ and layered δ -K, Mn^{3.5+} $O_{2-\nu}$, $zH_{2}O$ (Figure 1A) but differs from those of rocksalt MnO (octahedral Mn^{2+} with peaks at ~640 and ~641 eV) and spinel Mn_3O_4 (tetrahedral Mn^{2+} and octahedral Mn^{3+} with a peak at ~641 eV). On the basis of XRD performed on thicker films,⁷² we assigned the electrodeposited film to δ -K_xMn^{3.5+}O_{2-v}·zH₂O (Figure S2). Using the linear correlation between the nominal Mn valence and the center of gravity of the Mn L₃-edges (Figure 1C), the electrodeposited film was found to have a valence of Mn^{3.4+} after flushing the XAS cell and exchange of the KMnO₄ solution for O_2 -saturated 0.1 M KOH (Figure S3). The calculated Mn valence of our film $(Mn^{3.4+})$ is lower than that of Mn oxides electrodeposited potentiostatically from solutions of 0.5 mM Mn chloride and 0.1 M methylphosphate at 0.58 V vs NHE $(Mn^{3.7+})^{83}$ and at 1.05 V vs NHE $(Mn^{4.0+})^{60}$ but higher than those electrodeposited by the same method at 0.85 V vs NHE (Mn^{2.8+}).⁵¹ The latter's Mn valence is unusually low, and the authors discuss that their sample was reduced by X-ray irradiation. Our films were intermediate to those made by cycling (i) between 0.0 and 0.6 V vs Ag/AgCl at 20 mV/s and 900 rpm in Mn acetate and Na sulfate of 0.1 M each (Mn^{3.1+})⁴³ and (ii) between +2.15 and 0.75 V vs NHE at 100 mV/s without rotation or stirring in 0.5 mM Mn acetate (Mn^{3.8+}).¹⁵

The O K-pre-edge accesses the density of Mn states through hybridization with oxygen states. In a completely ionic description of the Mn–O bond, all p-orbitals would be filled; thus, no X-ray absorption signal could be expected. Yet in a covalent description of the Mn–O bond, electrons are shared between Mn and O by hybridization of molecular orbitals. Thus, the background-subtracted area under the O K-pre-edge is a measure of the density of empty states.^{84,85} Further in-depth discussion may be found in ref 84. The Mn valence is a measure



Figure 1. X-ray absorption spectroscopy at the (A) Mn L_{3,2}-edges of MnO, Mn^{2.7+} $_{3}O_{4}$, LiMn^{3.5+} $_{2}O_{4}$, δ -K_xMn^{3.5+} $O_{2-\nu}$ $zH_{2}O$, and β -Mn⁴⁺ O_{2} reference powders (Table S1) compared to those of Mn oxide electrodeposited on AulCrO_x|Si₃N₄ and recorded in O₂-saturated 0.1 M KOH under flow. (B) Crystal structures of the measured oxides. (C) Mn valence obtained by calibration to the center of gravity of the Mn L₃-edge of reference Mn oxides (squares) and test references not used in the calibration (circles). Circles additionally show the fit uncertainty propagated from the calibration curve of the Mn valence (gray line) (D) O K-edges of all samples. (E) Estimation of the Mn-O hybridization according to Suntivich et al.⁸⁴ using the normalized O K-pre-edge area integrated between 529 and 532 eV. The pre-edge of the electrodeposited film is drastically reduced by 0.1 M KOH and can thus not be compared directly to the dry reference samples. O Kspectra were recorded in partial fluorescence yield and Mn L_{3.2}-spectra in inverse partial fluorescence yield. The maximum of each spectrum in panels A and D was normalized to unity and then offset. An asterisk indicates a Mn³⁺ impurity in the MnO spectrum (Figure S5D). Errors indicate the standard deviation of 3-10 spots on the samples (Tables S3 and S4).

of the empty states and increasing Mn valence is associated with a pronounced increase in the PFY O K-pre-edge features below 532 eV (Figure 1D,E). The shape and onset of the O Kpre-edges were found to drastically differ when going from MnO and Mn₃O₄ to LiMn₂O₄, while changes are subtler among LiMn₂O₄, K_xMnO_{2-y}·zH₂O, and β -MnO₂. This is consistent with the trends observed at the Mn L_{3,2}-edges (Figure 1A).

Trends in hybridization of the references were obtained from the O K-pre-edge. The angular overlap of empty Mn 3d states with O 2p states depends on the orbital symmetry, where t_{2g} states contribute about 4 times less to the spectral intensity as compared to those with e_g symmetry.⁸⁴ Therefore, the contribution of the Mn states to the area under the pre-edge is removed by division of $(h_{e_g} + 1/4 h_{t_{2g}})$, where h_{eg} are the number of holes with e_g symmetry and $h_{t_{y_e}}$ is the number of Article

those with $t_{\rm 2g}$ symmetry. As the large exchange energy mandates high spin, there are 3 holes with t_{2g} symmetry for all investigated oxides, and the number of e, holes equals the Mn valence that was determined experimentally from the calibration curve in Figure 1C. The remaining area under the pre-edge is thus proportional to the hybridization of empty states, which we assume to be equal to the hybridization of filled states (i.e., covalence).⁸⁴ O K-pre-edges normalized by this established method⁸⁴ show increasing Mn–O hybridization with greater Mn valence (Figure 1E) for integration of the preedge intensities between 529 and 532 eV (arbitrarily defined). Mn-O hybridization in LiMn₂O₄, K_xMnO_{2-v}·zH₂O, and β - MnO_2 is mostly due to Mn 3d orbitals because of the orbital energies of $Mn^{3+}~(e_g{}^1t_{2g}{}^3)$ and $Mn^{4+}~(e_g{}^0t_{2g}{}^3).^{61,86}$ It should be noted that integration over the entire pre-edge of the O K-edge in Figure S9F includes the hybridization changes from unoccupied Mn 4s and Mn 4p orbitals $(a_{1g} \text{ and } t_{1u} \text{ of } Mn^{2+})$ in addition to those from unoccupied Mn 3d orbitals (e_g and t_{2g} of Mn³⁺ and Mn⁴⁺). Thus, the entire pre-edge is not a suitable measurement of Mn-O hybridization because the orbital contributions change among the Mn oxides. The O K-edge spectrum of the electrodeposited film in O2-saturated 0.1 M KOH was dominated by the signal of the aqueous electrolyte and the contribution of the Mn oxide to the total O K-signal was small (Figure 1D).

In Situ IPFY XAS at the Mn L_{3.2}-Edge and PFY XAS at the O K-Edge of Electrodeposited δ -K_xMn^{3.5+}O_{2-v}·zH₂O films. Noticeable changes were found at different applied potentials in the IPFY XAS spectra of the Mn L_{3,2}-edges collected in situ on the electrodeposited film in a threeelectrode cell, as shown in Figure 2A. At 1.65 V (steps 10a and 18a) and 1.50 V vs RHE (step 17a), the spectra strongly resembled that of the δ -K_xMn^{3.5+}O_{2-v}·zH₂O reference, while at 0.80 V (steps 13a and 15a) and 0.50 V vs RHE (step 14a), the spectra compared well with that of the $Mn^{2.7+}_{3}O_4$ reference that contains tetrahedral Mn²⁺. The changes in the Mn valence of the electrodeposited films at different applied potentials were quantified using the calibration curve in Figure 1C. Using the spectra in Figure 2A (steps 10a-18a in Table S2), the valence was initially $Mn^{3.1+}$ at 1.65 V vs RHE, then gradually decreased with reducing potential to Mn^{2.5+} at 0.50 V vs RHE (step 15a), and finally gradually increased with greater potential to Mn^{3.3+} at 1.65 V vs RHE (step 18a). The film was reduced below an average valence of 3+ below the onset potential of ORR (~0.9 V vs RHE) and oxidized above an average valence of 3+ above the onset potential of OER (~1.6 V vs RHE). Therefore, our data support the investigated Mn oxide containing tetrahedral Mn²⁺ during ORR conditions and Mn⁴⁺ during OER conditions in addition to Mn³⁺.

A hysteresis in the Mn valence associated with stepping the voltages down and up was noted in Figure 2B, which suggests that oxidation kinetics of the film are slower than the reduction kinetics. This hypothesis is supported by the observation that only the features of the δ -K_xMn^{3.5+}O_{2-y}·zH₂O reference were noted at OCV (step 7a), while all measurements after reduction at 0.80 V vs RHE (step 8a) retained a shoulder at 641.1 eV characteristic to Mn₃O₄. A similar hysteresis in the Mn valence changes is reported for electrodeposited Mn oxide between 0.7 and 1.9 V vs RHE in 0.1 M KPi^{S1} and for nanocrystalline hydrous Mn oxides between 0.65 and 1.65 V vs RHE in 2 M KCl,⁵⁹ where it has been attributed to poor electronic and ionic conductivity of Mn₃O₄ formed at low potentials. The hysteresis of the Mn valence affects the measured current densities



Figure 2. X-ray absorption spectroscopy of the electrodeposited film at the Mn L_{3,2}-edges recorded in inverse partial fluorescence mode using the three-electrode XAS cell with an AulCrO_x|Si₃N₄ window while flowing O₂-saturated 0.1 M KOH. (A) Waterfall plot of the spectra during cycling from 1.65 V via 0.50 to 1.65 V vs RHE. (B) Trends of the Mn valence during cycling. (C) Waterfall plot of the spectra during alternating reduction (0.50 V vs RHE) and oxidation (1.65 V vs RHE) steps. (D) Trends of the Mn valence during alternating oxidation/ reduction steps. Thick lines were smoothed by a second-order spline filter⁸⁷ using an 8 pt window. Powderous references of Mn₃O₄ (short dashed line) and δ-K_xMnO_{2-y}·zH₂O (solid line) are also included. All Mn valences were obtained using the calibration curve in Figure 1C, and error bars were propagated from the fit uncertainty of the calibration curve. The step number in Table S2 is also shown in each panel.

(Figure S13C), which suggests correlation with electrocatalysis. However, further *in situ/operando* experiments are required to clearly prove a catalytic role of the observed changes in Mn valence.

Partial reversibility of spectral changes was further demonstrated by repeating experiments between 0.50 V (ORR) and 1.65 V vs RHE (OER) three times (Figure 2C), where spectra at 0.5 V vs RHE resembled $Mn^{2.7+} {}_{3}O_{4}$ and spectra at 1.65 V vs RHE resembled that of δ -K_xMn^{3.5+}O_{2-y}·zH₂O. Comparable Mn valence was obtained at the same applied potentials in these three repeats, as shown in Figure 2D, where the average valence during ORR was Mn^{2.5+} and the average valence during OER was Mn^{3.2+}.

We further fit all spectra using those from powders of $Mn^{2.7+}{}_{3}O_{4}$ and δ -K_xMn^{3.5+}O_{2-y}·zH₂O to quantify the similarity to these standards and support the assignments above (Figure

S8; further details are given in Experimental Section). In support of our previous discussion, the initial measurement at OCV (step 7a) showed (100 ± 2)% δ -K_xMn^{3.5+}O_{2-y}·zH₂O character, while reduction at 0.8 V vs RHE resulted in up to (99 ± 4)% Mn^{2.7+}₃O₄ character (step 14a) with an average of (87 ± 2)%, and oxidation at 1.65 V vs RHE was limited to (76 ± 2)% δ -K_xMn^{3.5+}O_{2-y}·zH₂O character (step 20) with an average of (71 ± 1)% (Figure S8). The excellent agreement with the Mn₃O₄ standard also corroborates Mn²⁺ having tetrahedral coordination in the investigated voltage range. The two most common analyses of soft XAS, calibration of the Mn valence and fitting standards; thus, both support partial reversibility of

and Mn⁴⁺ at OER conditions. In situ O K-edge PFY XAS spectra revealed partially reversible changes in the pre-edge with applied voltage (Figure 3). The O K-pre-edges were found to include a peak labeled (i) near 529.7 eV, which can be attributed mainly to δ -

the Mn redox as well as tetrahedral Mn²⁺ at ORR conditions



Figure 3. X-ray absorption spectroscopy of the electrodeposited film at the O K-edge recorded in partial fluorescence mode using the threeelectrode XAS cell with an AulCrO₄|Si₃N₄ window while flowing O₂-saturated 0.1 M KOH. (A) Waterfall plots of the O K-edge and (B) areas under peaks (i) and (ii) during cycling from 1.65 V via 0.50– 1.65 V vs RHE. (C) Waterfall plots of the O K-edge and (D) areas under peaks (i) and (ii) during alternating reduction (0.50 V vs RHE) and oxidation (1.65 V vs RHE) steps. The voltages and step numbers in Table S2 are shown near each spectrum. Dotted lines show exemplary fits of peak (i) and peak (ii), further fits are in Figures S10 and S11. Reference spectra of Mn₃O₄ (short dashed line) and δ -K_xMnO_{2-y}·zH₂O (solid line) powders and a clean AulCrO_x|Si₃N₄ window (long dashed line) are also included. Error bars indicate the fit uncertainty.

 $K_r Mn^{3.5+}O_{2-\nu} \cdot zH_2O$ with a minor contribution from $Mn^{2.7+}{}_{3}O_4$ at the high-energy side of the peak (Figure 3A) and a peak labeled (ii) at 532.3 eV already present in the AulCrO_x|Si₃N₄ window with a strong background from both δ -K_xMn^{3.5+}O_{2-v}. zH_2O and $Mn^{2.7+}_{3}O_4$ (Figure 3A). The spectra were further analyzed by fitting Gaussians to these two peaks and a shoulder in the O K-edge as the background (Figures S10; further details are given in Experimental Section). The area under peak (i) decreased with applied voltage to a minimum at 0.80 V vs RHE (step 15b), after which it increased again (Figure 3B). In contrast, the area under peak (ii) remained relatively constant and then dropped steeply after step 16b. An identical analysis was performed for alternating oxidation at 1.65 V vs RHE and reduction at 0.50 V vs RHE (Figures 3C and S11), where again only the area under peak (i) followed the voltage but on a background of overall reducing peak area (Figure 3D). This suggests that peak (i) correlates with electronic properties of the electrodeposited film, while the diverging trend of peak (ii) suggests that it correlates with the degradation of the AulCrO_xl Si₃N₄ window.

The ratio of the areas of peaks (i) to (ii) was employed to remove the influence of film loss due to degradation of the window material (Figure 4A). The resulting trend with voltage



Figure 4. Trends of the area ratio (solid circles) of peak (i) to peak (ii) in Figure 3A,B (A) for cycling from 1.65 V vs RHE to 0.50 V vs RHE with intermediate voltages and back; as well as (B) for alternating reduction at 0.50 V vs RHE and oxidation at 1.65 V vs RHE. For comparison, the Mn valence (open circles) obtained from the calibration data in Figure 1C of the main text is also shown. (C) Linear correlation between the area ratios of peak (i) and peak (ii) in panels A and B with the Mn valence obtained from the calibration data in Figure 1C of the main text. Error bars were obtained by propagating the uncertainties of the fitted amplitudes of peaks (i) and (ii) as well as the uncertainties of the Mn valence calibration curve.

strongly resembles that of the Mn valence (Figure 4A) obtained from analysis of the Mn L₃-edge that is based on the experimental calibration in Figure 1C. Similarity among the trends in the O K-and Mn-L edge is expected as the metal dband and oxygen p-band increase their overlap with higher valence in perovskites ABO₃ ($B = Mn^{45}$ and Ni⁸⁴), which can serve as references for the electrodeposited film as the relevant metal 3d states are well separated from hybridization with other elements (e.g., A = Ca or Pr).^{45,88} Highly similar trends between the peak area ratio and Mn valence were also observed for alternating reduction at 0.50 V vs RHE and oxidation at 1.65 V vs RHE (Figure 4B). Analysis of all recorded O K-preedges demonstrates clearly that the peak ratio scales linearly with the experimental Mn valence based on the L₃-edge (Figure 4C). Therefore, the presented analysis could be used to determine the metal valence state by in situ analysis of the O Kpre-edge if a suitable calibration was performed.

The pre-edge of the oxygen K-edge data is sensitive to both valence and hybridization as we discussed above for powder reference compounds in Figure 1. Having established the sensitivity of the in situ O K-data on Mn valence, we now analyze the hybridization, for which the peak ratios of the areas of peaks (i) and (ii) were further normalized by the weighted number of holes on the Mn cation $(e_g \text{ holes } + 1/4t_{2g} \text{ holes})^{84}$ obtained from the experimentally determined Mn valence. To emphasize relative changes, we divided by scan 10b (1.65 V) for decreasing voltage (Figure S14A), by scan 14b (0.50 V) for increasing voltage (Figure S14A), and by scan 7b (OCV) for alternating between 0.50 and 1.65 V vs RHE (Figure S14B). This normalized peak ratio decreased for reduction from Mn^{3.1+} to Mn^{2.5+} when decreasing the voltage from 1.65 to 0.50 V vs RHE and increased for oxidation from Mn^{2.5+} to Mn^{3.0+} (Figure S14A), but the obtained changes are not significant within the uncertainty of the ratios.

The in situ trends are qualitatively consistent with the ex situ trends of the reference samples in Figure 1E, where the hybridization decreased from the $Mn^{2.7+}_{3}O_{4}$ reference to the δ - $K_xMn^{3.5+}O_{2-y}$ zH_2O and $LiMn^{3.5+}O_4$ references. The spectrum of β -Mn⁴⁺O₂ (e_g⁰) showed a lower hybridization as compared to that of the Mn^{3.5+} (e_g^{0.5}) references, which might explain the lower hybridization of the electrodeposited film with Mn^{3.4+} $(e_{\sigma}^{0.6})$ valence as compared to the $\hat{Mn}^{3.0+}$ (e_{σ}^{-1}) valence when the eg orbital is emptied. The reduction of the hybridization above Mn^{3+} (e_g^{1}) can also be seen qualitatively for alternating reduction at 0.50 V vs RHE and oxidation at 1.65 V vs RHE (Figure S14B), where the hybridization decreases for reduction and increases for oxidation, albeit not significantly within the uncertainty of the ratios. Additionally, the difference in hybridization qualitatively decreases with the difference in Mn valence (Figure S14B). In summary, analysis of the O K-preedge area shows qualitative trends of changes in Mn-O hybridization during in situ experiments, where an eg occupancy near one likely corresponds to the highest hybridization.

In this report, we have analyzed changes of the Mn valence with applied voltage during *in situ* Mn $L_{3,2}$ -and O K-edge XAS. Furthermore, we analyzed changes in the Mn–O hybridization of an electrodeposited Mn oxide for the first time using *in situ* O K-edge XAS. We found a mixed Mn^{3+/4+} valence above applied potentials of 1.20 V vs RHE (i.e., clearly before the onset of the OER). This is in agreement with previous *in situ* studies on electrodeposited Mn oxides at the Mn K-^{12,18,50,59} and Mn $L_{3,2}$ -edges^{51,60} in various electrolytes (0.1 M KOH,¹⁸ 0.1 M KPi,^{12,51,60} 0.1 M NaSO₄,⁵⁰ and 2 M KCl⁵⁹). In our *in*

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situ data, there is also clear evidence for 20% Mn^{4+} ions on average during oxygen evolution at 1.65 V vs RHE (Figure 4D), which disagrees with *ex situ* XPS studies of bixbyite Mn_2O_3 .²¹ The difference might be due to the drying procedure before XPS measurements or the higher crystallinity⁴³ of the material in the *ex situ* XPS study, which would hinder accumulation of holes and remove any excess charge during the XPS measurement where the sample is grounded. *Ex situ* XPS¹⁶ and XAS^{10,14,15,43} studies also clearly show a mixed $Mn^{3+/4+}$ valence after OER in 0.1 M KOH^{14,43} and 0.1 M KPi^{10,14–16} for less crystalline electrodeposited Mn oxides in agreement with our *in situ* XAS study.

The observation of tetrahedral Mn^{2+} during ORR in our *in* situ L-edge study is likewise in agreement with previous Mn Kedge reports where the spinel structure was identified by fitting the Mn K-edge spectra to references⁴⁷ and by EXAFS analysis.¹⁸ In our data, the presence of tetrahedral Mn^{2+} can be directly seen as the emergence of a new peak, which greatly simplifies analysis. It is crucial to separate tetrahedral Mn^{2+} from the Mn valence at octahedral sites, which trend with ORR activity.²² Our study demonstrates that this is possible by *in situ* experiments at the Mn L_{3,2}-edges, which allow faster acquisition as compared to EXAFS analysis at the Mn K-edge.

In situ analysis of the hybridization qualitatively supports the special role of single eg occupancy for oxygen electrocatalysis. Understanding Mn-O hybridization is crucial for oxygen electrocatalysis because it relates to the adsorption of oxygen on the catalytic surface.³ Similar in situ O K-edge studies have previously only been available for hematite $(\alpha - Fe_2O_3)^{67,68}$ and electrodeposited Ni oxide.⁶⁹ We obtained this hybridization by analysis of the O K-pre-edge at various applied voltages, which was consistent with the trends identified based on valence from select ex situ references. The maximum of Mn–O hybridization is estimated for a valence close to Mn³⁺ with an e_o occupancy near unity, where the maximum activity for both OER and ORR was reported for perovskites^{19,23,27} and spinels²² based on ex situ analysis. Thus, the presented analyses at the Mn-L₃ and particularly O K-edges set the stage for correlations between Mn-O hybridization and catalytic activity toward testing known activity descriptors at catalytic conditions.

CONCLUSIONS

We probe changes in the Mn valence and Mn-O hybridization using IPFY XAS at the Mn L_{3,2}-edges and PFY XAS at the O Kedges of electrodeposited δ -K_xMnO_{2-v}·zH₂O. Experiments are performed in situ in 0.1 M KOH solution as a function of voltages relevant to ORR and OER. IPFY Mn L_{3.2}-edges provide greater chemical sensitivity to Mn valence as compared to the conventional PFY detection, where the center of gravity of the Mn L₃-edge is established to correlate with the Mn valence using a number of reference Mn oxides. Using this calibration, the average Mn valence of electrodeposited δ - $K_rMnO_{2-\nu}$ zH_2O has been determined as a function of the applied voltage. The valence relevant to ORR is below Mn³⁺ and contains tetrahedral Mn²⁺, while the valence relevant to OER is above Mn^{3+} and contains Mn^{4+} ions. Both are in accordance with previous ex situ and in situ studies on electrodeposited Mn oxides. Stepping back and forward between ORR and OER voltages is shown to be reversible, but there is hysteresis in the Mn valence, which can be attributed to slow oxidation kinetics of a Mn₃O₄-like phase with poor electronic conductivity as proposed by Chang et al.⁵⁹ In situ analysis of the O K-edge suggests that the Mn-O

hybridization decreases for valences above and below Mn^{3+} . Combining this *in situ* insight with previous *ex situ* studies on perovskites^{19,23,27} and spinels²² fortifies the notion that having significant Mn^{3+} (i.e., about one e_g electron) is critical in catalyzing OER and ORR. Therefore, *in situ* experiments at both the metal L- and oxygen K-edges are a powerful approach for (1) identifying the active valences during electrocatalytic reactions and (2) the metal–oxygen hybridization of active catalysts.

ASSOCIATED CONTENT

S Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpcc.7b05592.

TEM and SAED of δ -MnO₂; XAS of Mn²⁺ references; additional XAS spectra; XRD of Mn powders; electrochemical currents during XAS; Mn–-L₃ fit results; O Kfit results; physical data of Mn oxide powders; sequence of XAS measurements; analysis results of the Mn–L₃ and O K-edges; energy calibration of soft XAS in literature; calculation of absence of pH changes (PDF)

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