Mechanistic Parameters of Electrocatalytic Water Oxidation on LiMn$_2$O$_4$ in Comparison to Natural Photosynthesis

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Targeted improvement of the low efficiency of water oxidation during the oxygen evolution reaction (OER) is severely hindered by insufficient knowledge of the electrocatalytic mechanism on heterogeneous surfaces. We chose LiMn$_2$O$_4$ as a model system for mechanistic investigations as it shares the cubane structure with the active site of photosystem II and the valence of Mn$^{3.5+}$ with the dark-stable S1 state in the mechanism of natural photosynthesis. The investigated LiMn$_2$O$_4$ nanoparticles are electrochemically stable in NaOH electrolytes and show respectable activity in any of the main metrics. At low overpotential, the key mechanistic parameters of Tafel slope, Nernst slope, and reaction order have constant values on the RHE scale of 62(1) mV dec$^{-1}$, 1(1) mV pH$^{-1}$, $-0.04(2)$, respectively. These values are interpreted in the context of the well-studied mechanism of natural photosynthesis. The uncovered difference in the reaction sequence is important for the design of efficient bio-inspired electrocatalysts.

Introduction

In light of the globally increasing energy consumption, efficient electrocatalytic energy storage and conversion are paramount for the transition from dwindling fossil sources to sustainable sources owing to the intermittent nature of renewable energy.[1,2] Among the available options, energy storage in hydrogen bonds is highly attractive due to the large gravimetric energy densities of the resulting chemical fuels.[3,4] Water is the most abundant sustainable proton source, which makes water splitting a cornerstone of sustainable fuels.

The scientific grand challenge of water splitting is the large overpotential of the half-reaction that oxidizes water to oxygen,[5] that is, the oxygen evolution reaction (OER). The best currently known electrocatalysts for the OER suffer from overpotentials around 0.3 V higher than the thermodynamic limit,[6] which is very large as it equals, for example, the span between the least and most active perovskite oxides.[7] Property–activity relationships have proven a promising approach in the search for more active catalysts based on earth-abundant materials. Yet, further targeted progress on earth-abundant electrocatalysts is hampered by insufficient experimental insight into the nature of intermediates, the rate-limiting step (RLS), and the mechanistic pathway.

The best understood catalyst for oxygen evolution by water oxidation is the CaMn$_2$O$_4$ cofactor of photosystem II (PSII),[2, 10–12] the paragon of all artificial photosynthesis work and blueprint for solar fuels catalysts.[6, 13–18] The structure of the active site is known with high resolution[8, 9, 19] and furthermore, the active states have been studied for decades by various experimental and theoretical methods.[2, 10–12, 20–23] The mechanism of water oxidation in natural photosynthesis is the so-called S-state cycle or Kok cycle[24] (Figure 1). In the dark-stable state S$_4$, the four Mn ions have an average valence of Mn$^{3.5+}$. Light flashes advance the catalytic cycle by oxidizing Mn ions to Mn$^{4+}$ in state S$_3$. This high-valent state drives oxygen evolution at the site marked with an asterisk after an additional light flash.[25] The state resulting after oxygen evolution, S$_0$, can then be oxidized back to state S$_1$ by a fourth flash. In contrast, the active states of heterogeneous surfaces are much less defined and extremely difficult to study experimentally. Most insight into the mechanisms on electrocatalytic surface is derived from the analysis of often ambiguous electrochemical parameters.

Electrochemical analysis of mechanistic parameters is very mature.[26–30] Recently, Minguzzi et al.[31] introduced dynamic
voltage–pH diagrams for the discussion of kinetic information akin to thermodynamic information in a Pourbaix diagram.\[32\] The approach was further extended by Haghighat and Dawlaty\[33, 34\] to display the electron-transfer coefficient (i.e., Tafel slope) and reaction order. These studies inspired us to seek a convenient graphical representation of mechanistic parameters in pH-dependent studies with high current and voltage resolution.

Here, we discuss LiMn$_2$O$_4$ in alkaline media as an electrocatalytic model for the active site of PSII, with which it shares the cubane motif. The $\lambda$-LiMn$_2$O$_4$ materials system has been previously investigated using photochemical methods and X-ray absorption spectroscopy (XAS). Mn loss is analyzed using rotating-ring disk electrodes (RRDE) to identify conditions under which potential Mn loss interferes least with oxygen evolution. Then, the currents owed to oxygen evolution are measured by RRDE at various alkaline pH values to obtain the Tafel slope, Nernst slope, and reaction order. These studies inspired us to seek a convenient graphical representation of mechanistic parameters in pH-dependent studies with high current and voltage resolution.

Results and Discussion

The expected cubic crystal structure of semiconducting LiMn$_2$O$_4$ was confirmed by XRD at room temperature (Figure 2a). Rietveld analysis of the powder XRD patterns confirmed the space group of Fd$ar{3}$m (#227) and a lattice parameter of $a=8.15\ \text{Å}$ (Table 1). LiMn$_2$O$_4$ can exhibit a charge-ordered structure with space group I4$_1$/amd at low temperatures,\[40\] with the first-order transition of ordering occurring at 280–290 $\text{K}$,\[41, 42\] but we found no indication of the charge-ordered structure in our analysis. The unit cell contained 8 formula units in a volume of 541.46(1) $\text{Å}^3$, which resulted in a bulk density of 4.436(1) $\text{g/cm}^3$ of the LiMn$_2$O$_4$ particles. Furthermore, there are 6 Mn atoms on the most stable (001) surface, which yields a surface density of 9 x $10^{14}$ $\text{Mn cm}^{-2}$. The crystal structure consisted entirely of edge-sharing octahedra with distances of 1.93 Å (Mn–O) and 2.88 Å (Mn–Mn) that exhibited Mn–O–Mn bond angles of 96.7°. The Mn–O bond length is intermediate to the average bond lengths of octahedral Mn$^{3+}$–O (2.015 Å) and Mn$^{4+}$–O (1.899 Å),\[44\] which supports mixed Mn$^{3+}$–$^{4+}$ valence. The structure can also be interpreted as consisting of Mn$_2$O$_4$ cubane units where voids are filled with Li (Figure 2a, inset). Thus, the cubane motif relates LiMn$_2$O$_4$ structurally to the CaMn$_2$O$_4$ cofactor of PSII (Figure 1).

Morphology and particle sizes of the LiMn$_2$O$_4$ powder were analyzed by scanning electron microscopy (SEM). The particle shapes were well approximated by ellipsoids (Figure 2b, inset).
and can thus be projected as ellipses with principle axes a and b. Histograms of the equivalent diameter of a circle \(d_e = \sqrt{2ab}\) were generated based on 200 randomly selected particles (Figure 2b). They were fit by a lognormal distribution with most frequent diameter (i.e., mode) of 69 nm, from which a mean particle diameter of 79(43) nm was calculated. The vendor specifies particle sizes of <500 nm based on BET analysis, which was in agreement with our microscopic analysis. Finally, the particle distribution of the equivalent diameter \(d_e\) and experimental bulk density \(\rho\) were used in the Cauchy expression \((A = 6\rho / \Sigma d_e^{2}/\Sigma d_e^{3})\) to calculate a specific area of 12.264(2) m² g⁻¹ for LiMnO₄.

Soft X-ray absorption spectroscopy (XAS) was used to elucidate the valence of the LiMnO₄ powder. The Mn L₃ edge involves a transition from Mn₂P₂O₇ core to Mn 3d valence states. However, the wave functions of the core and valence states overlap significantly so that a complex multiplet fine structure is detected.\(^\text{[46]}\) Spectra were acquired in the surface-sensitive total electron yield (TEY; escape depth \(\approx 7 \text{ nm}\))\(^\text{[47,48]}\) and the bulk-sensitive inverse partial fluorescence yield (IPFY).\(^\text{[49]}\) The IPFY is effectively a measure of the X-ray attenuation length akin to X-ray transmission experiments and as such less susceptible to saturation effects compared to the conventional partial fluorescence yield (PFY).\(^\text{[49]}\) The spectra were normalized to peak β for better comparison of the energy positions and relative intensities of the features in the Mn L₃ edges.

The peak positions and shapes of LiMnO₄ are typical for those of Mn\(^{3+}^{\text{+}}\) oxides in octahedral coordination.\(^\text{[50]}\) The spectra showed a sharp peak near 642.0 eV (denoted α in Figure 2c) and a broader peak with higher intensity near 644.5 eV (denoted β in Figure 1c). Peak β was intermediate to that of the Mn\(^{2+}\)O and β-Mn\(^{4+}\)O, which indicated a mixed Mn\(^{3+}\) valence. Peaks α and β were shifted to higher energies in the TEY spectra, which indicated surface oxidation. Mn valences of 3.5(3) and 3.9(2) were determined from the IPFY and TEY spectra of LiMnO₄ using Mn\(^{2+}\)O\(₄\), La\(₂O₃\)Mn\(^{4+}\)O\(₃\) and β-Mn\(^{4+}\)O\(₂\) as references for calibration (more detail in the Experimental Section). Thus, the determined Mn valences support slight surface oxidation, which is, however, not significant within the uncertainty of the calculation method.

The crystal field splitting and Mn–O covalence was studied by analysis of the OK pre-edge, which accesses the unoccupied manganese density of states through hybridization with unoccupied oxygen states, that is, holes, in antibonding \(\sigma^*\) and \(\pi^*\) orbitals. The conventional PFY is reported for the OK edge because there was no suitable absorption edge for IPFY analysis. The shape of the TEY and PFY spectra resembled that of the β-Mn\(^{4+}\)O\(₂\) reference, yet peaks γ and δ in Figure 2d were shifted to lower energies in LiMnO₄ compared to β-Mn\(^{4+}\)O\(₂\), in which the TEY spectra were less shifted than the PFY spectra. It should be noted that the PFY spectrum is broader, which might affect the apparent peak positions. Peak γ can be assigned to spin-up holes in the \(e_g\) orbitals of the Mn\(^{3+}\)- (one \(e_g\) hole) and Mn\(^{2+}\) ions (two \(e_g\) holes),\(^\text{[51,52]}\) which further overlap with the \(t_{2g}\) spin-down holes of the Mn\(^{3+}\) ion.\(^\text{[51]}\) Peak δ can be assigned to the energetically close spin-down \(e_g\) holes on Mn\(^{4+}\) and spin-down \(t_{2g}\) holes on Mn\(^{3+}\) ions.\(^\text{[53]}\) The hybridization between these frontier orbitals of Mn and O can be experimentally determined by integration of the spectral intensity under peaks γ and δ normalized by the contribution of the holes to spectral intensity \((e_g\) holes + \(t_{2g}\) holes).\(^\text{[54]}\) We assume that the integral under the peaks is a good estimation for the covalence of the Mn–O bond, which was highest for LiMnO₄ among β-Mn\(^{4+}\)O\(₂\) and Mn\(^{2+}\)O\(₂\).\(^\text{[50]}\) High covalence was also supported by the Mn–O–Mn angle of 96.7° obtained by XRD analysis. It could lead to oxygen radicals,\(^\text{[55]}\) which are also discussed in the mechanism of natural photosynthesis (Figure 1, S₃).\(^\text{[10]}\)

In summary, physical characterization established that LiMnO₄ shares the cubane structure with the CaMnO₄ cofactor of PSII and the Mn\(^{3.5+}\) valence with the dark-stable S₃ state in the mechanism of natural photosynthesis. High Mn–O covalence potentially fosters oxygen radicals, which are discussed to play an important role prior to oxygen evolution in natural photosynthesis.

The electrochemical current of LiMnO₄ was analyzed using a RRDE setup in NaOH with molarities between 0.1 M (pOH 1 = pH 13) and 1.0 M (pOH 0 = pH 14). The buffer capacity of the used alkaline electrolytes is at least four-fold higher than that of the commonly used 0.1 M phosphate buffer at pH 7\(^\text{[50]}\) so that electrolyte additives could be omitted to avoid possible side reactions. Voltammetry showed decreasing currents with cycling where the current at the scan apex halved within 10 cycles (Figure 3a). The first cycle exhibited broad wave with onset near 1.4 V versus RHE (reversible hydrogen electrode), which was absent in later cycles. To investigate the origin of these currents, the ring was set to reduce MnO\(₂\) or MnO\(₂\), which are thermodynamically favorable in the investigated voltage range.\(^\text{[32]}\) The uncorrected ring currents were about 3 orders of magnitude smaller than the disk currents, indicating a small effect. During the first cycle, the ring currents rose simultaneously with the disk currents at 1.4 V versus RHE, that is, well before the onset of oxygen evolution. At voltages above 1.6 V versus RHE, the ring currents saturated in the first

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**Figure 3.** Stability during voltage cycling of LiMnO₄-carbon composite electrodes. \(a\) Select voltammetry cycles of disk currents and corresponding ring currents (open diamonds). The ring was set to 1.2 V versus RHE to detect reduction of high-valent Mn species. A contrast ring measurement of carbon (solid squares) is included. Representative high-resolution transmission electron microscopy (HR-TEM) of \(b\) LiMnO₄ ink and \(c\) of an ink-casted electrode cycled 10 times between 1.25 and 1.75 V versus RHE. All electrochemical measurements were conducted in O\(₂\)-purged 0.25 M NaOH (pH 13.4) at 10 mVs⁻¹ and 1600 RPM.
cycle, whereas the disk currents rose exponentially. The deviation of the ring and disk currents at high currents indicated that the currents could be measured independently (i.e., no detectable crosstalk). The release of Mn from LiMn$_2$O$_4$ quickly abated with cycling. The ring currents of the 5th and 10th cycle (open diamonds in Figure 3a) were small (< 0.1 mA) and equal within the noise level. Moreover, the detected currents were barely above a background measurement of a glassy carbon electrode loaded only with acetylene black carbon. During the later cycles, the onset of the ring currents of LiMn$_2$O$_4$-carbon electrodes shifted to higher voltages and might coincide with oxygen evolution as reported previously for electrodeposited Mn$^{1+/4+}$ oxides. Nonetheless, these currents were within the background and did not affect the interpretation of our results.

Post-mortem transmission electron spectroscopy (TEM) established the absence of significant structural changes owing to Mn loss. Particles in the as-prepared LiMn$_2$O$_4$ ink (Figure 3b) were compared with particles on an electrode prepared from the same ink after 10 cycles between 1.25 and 1.75 V versus RHE at pH 13.4 (Figure 3c). Representative TEM images demonstrated crystallinity up to the surface of both the ink and the cycled particles. No indications of surface amorphization or phase changes were observed, which is in clear contrast to perovskite oxides cycled under comparable conditions. We conclude that the cubane structure of LiMn$_2$O$_4$ is preserved for at least 10 voltage cycles and selected the 5th cycle for further evaluation of catalytic activity.

Capacitance-corrected disk currents during the 5th cycle were independent of the purging gas. Little hysteresis was observed above 1.65 V versus RHE, whereas capacitive currents introduced hysteresis at lower voltages. We corrected for capacitive currents by averaging the anodic and cathodic voltage scans (Figure 4a) which resulted in the expected vanishing currents before an exponential current rise at approximately 1.5 V versus RHE. The capacitive-corrected disk currents were equal within experimental spread in O$_2$-purged and Ar-purged NaOH at pH 13 (Figure 4b). This indicated that the LiMn$_2$O$_4$ redox was independent of the oxygen pressure at the electrode. Although the equilibrium potential of oxygen redox is defined at standard conditions, that is, 1 bar O$_2$ pressure, the expected voltage shift due to the pressure term in the Nernst equation could not be resolved within the experimental uncertainty. We conclude that experiments performed in both O$_2$-purged and Ar-purged NaOH can be compared directly, which enables oxygen detection at the ring in Ar-saturated electrolyte.

Qualitative oxygen detection at the ring was established by oxygen reduction in Ar-saturated NaOH electrolytes (Figure 4c). Quantitative detection is hindered by trapping of bubbles at the disk and the PTFE (polytetrafluoroethylene) spacer. Instead of the conventional collection efficiency, the ring currents were scaled to match the disk currents between 20 and 100 μA, where we expect low bubble formation. The scaling factors at 1600 rpm were 0.062(22) at pH 13 and 0.061(14) averaged over all investigated pH values during the 5th cycle. The deviation to the expected value of 0.424 for the investigated RRDE setup suggested significant trapping of oxygen bubbles. Yet, the similarity of the scaling factors among all investigated electrodes indicated that the oxygen flux to the ring was reproducible for low disk currents. Although the currents of the capacitance-corrected disk and scaled ring were identical within experimental spread due to the scaling factor, the Tafel slope (dlog(i/R)/dE) differed within one standard deviation, which is analyzed further below. It should be noted that Tafel slopes are independent of multiplied factors and the scaling factor mainly facilitates easier comparison between the ring and disk currents.

Both previously discussed processes, Mn loss and oxygen evolution, were independent of pH on the RHE scale within experimental scatter. Mn loss was investigated in O$_2$-saturated NaOH at pH 13.0, 13.4, and 14.0 during the 1st cycle, where it was most pronounced (Figure 5a). The Pourbaix diagram predicts an increasing equilibrium potential for MnO$_2$ formation and possibly MnO$_4^{2-}$ formation when increasing the pH from 13 to 14 on both the RHE and NHE (normal hydrogen

![Figure 4](image1.png) Exemplary electrocatalytic measurements of LiMn$_2$O$_4$-carbon composite electrodes. (a) Cyclic voltammetry of disk currents during the 5th cycle as measured and capacitance-corrected by averaging anodic and cathodic voltage scans. Arrows indicate the scan direction. (b) Comparison of the capacitance-corrected disk currents in Ar- and O$_2$-purged NaOH at pH 13. (c) Comparison of the disk currents in Ar-purged NaOH at pH 13 (line) and corresponding scaled ring currents of the anodic scan (open circles). All electrochemical measurements were conducted in 0.10 M NaOH (pH 13.0) at 10 mV s$^{-1}$ and 1600 RPM. Error bars indicate the standard deviation (1σ) of 3 electrodes.

![Figure 5](image2.png) Dependence of Mn loss and oxygen evolution on pH of LiMn$_2$O$_4$-carbon composite electrodes. (a) Cyclic voltammetry of disk currents and corresponding ring currents of Mn reduction during the 1st cycle in O$_2$-purged NaOH. (b) Capacitance-corrected disk currents and corresponding ring currents of oxygen reduction during the 5th cycle in Ar-purged NaOH. All electrochemical measurements were conducted at 10 mV s$^{-1}$ and 1600 RPM. Error bars indicate the standard deviation (1σ) of 3 electrodes.
electrode) scales. The average ring current owed to MnO₂ reduction was indeed highest at pH 14, yet identical to that of the other pH values within experimental scatter. Thus, no significant pH dependence of Mn loss could be detected within experimental uncertainty and the cumulative Mn loss up to the 5th cycle is not expected to affect electrocatalysis.

The capacitance-corrected disk currents and scaled ring currents fall onto each other in Ar-saturated NaOH during the 5th cycle at all investigated pH values, that is, pH 13.0, 13.2, 13.4, and 14.0, up to currents of approximately 0.5 mA (Figure 5b). This demonstrated that Mn loss, albeit present, was reproducible enough to avoid large scatter in later cycles. Furthermore, no pH dependence is expected on the RHE scale when hydroxide is the reactant of the OER, which is discussed further below.

In summary, the RRDE experiments demonstrated that side reactions were negligible and the scaled ring currents matched the capacitance-corrected disk currents at any pH low currents. Therefore, we assign the scaled ring currents to the catalytic rate of oxygen evolution (\(\text{O}_2\)) depletion, which is the relevant current for mechanistic analysis.

The activity of LiMn₃O₄ was determined in the most common metrics (Figure 6 and Table 2) for comparison with other electrocatalysts. We analyzed the capacitance-corrected disk currents due to the aforementioned issues of obtaining quantitative information from ring currents and lack of ring data for comparison. Thus, the disk currents were normalized by the disk area of 0.126 cm² \(\text{disk}^2\), oxide mass loading of 50(1) µg \(\text{disk}^2\) or the experimentally determined oxide area of 6.1(2) cm² \(\text{disk}^2\). Additionally, the surface turnover frequency (TOF-s) and bulk turnover frequency (TOF-b) were calculated using the Mn surface \((9 \times 10^{14} \text{Mn cm}^{-2})\) and bulk \((4.436(1) \text{g cm}^{-3})\) densities obtained from XRD analysis. The current densities were independent of pH below 1.7 V versus RHE within the experimental scatter and diverge for higher voltages at pH 13.4 and 14.0. Furthermore, higher currents will lead to more oxygen bubble formation, which is also expected to contribute to the divergence of disk currents at high voltages. The (unscaled) ring currents were above noise for currents > 0.6 µA (Figure 6 inset), which gives an onset of oxygen detection of about 1.59 V versus RHE independent of pH. Moreover, the ring currents overlap within experimental uncertainty between the onset and approximately 1.68 V versus RHE.

Previous reports of the OER activity of LiMn₃O₄ are inconsistent. Cady et al. studied composite electrodes of LiMn₃O₄ carbon, and neutralized Nafion in 1.0 M NaOH, which showed no activity above background. It is conceivable that the high Nafion concentration of their electrodes impedes electronic conduction, making LiMn₃O₄ electrochemically silent and catalytically inactive. On the other hand, Wei et al. studied the overpotential of composite electrodes of carbon, Nafion, and spinel oxides including LiMn₃O₄ at 25 µA cm⁻² in 0.1 M KOH. They reported similar overpotentials for LiMn₃O₄ and Co₃O₄ which is in agreement with the small difference in overpotentials of LiMn₃O₄ in our study and Co₃O₄ in Jung and co-workers (see also next paragraph). Moreover, we assigned the reduction currents at the ring to oxygen evolved at the disk (Figures 4b–c, 5b, 6). Therefore, we conclude that LiMn₃O₄ is active for electrocatalytic oxygen evolution.

A common activity metric in the field of solar fuels is the voltage to obtain \(j_{\text{disk}} = 10 \text{mA cm}^{-2}\) because it matches a 10% efficient solar cell. The LiMn₃O₄–carbon composite electrode provided 10 mA cm⁻² at 0.520(2) V overpotential (1.749(2) V versus RHE) at pH 14, which is similar to or better than other first-row transition metal nanoparticles such as MnO \((0.51(4) \text{V})\), Mn₃O₅ \((0.53(4) \text{V})\), MnO₂ \((0.50(3) \text{V})\), Co₃O₄ \((0.50(1) \text{V})\), and NiFeOₓ \((0.51(1) \text{V})\), but higher than that of MnOₓ \((0.43(2) \text{V})\). The nanoparticles in the latter study of Jung and co-workers have surface areas within one order of magnitude of LiMn₃O₄ and twice the oxide loading used herein \((0.8 \text{mg cm}^{-2})\).

The catalyst mass correlates with the cost and weight of an electrolysis device and is thus another common normalization. The LiMn₃O₄–carbon composite electrode reached 10 Ag⁻¹ at 1.705(3) V versus RHE \((0.476(3) \text{V overpotential})\) at pH 13.4 and 14.0, which is similar to Ca₂MnO₄ \((1.70 \text{V versus RHE; 5th cycle;} \text{“sub-micron”})\) and β-MnO₂ \((1.71 \text{V versus RHE; 5 m² g}^{-1})\). Yet, the overpotential is higher than that of Mn oxides with higher surface area, for example, solvent-free α-MnO₂–SF \((1.64 \text{V versus RHE; 112 m² g}^{-1})\) and oxides with similar sur-

| Table 2. Electrocatalytic activity of LiMn₃O₄–carbon composite electrodes. |
| --- |
| Metric | \(j\) | \(E\) [V versus RHE] \(\text{at 1.68 V versus RHE}^a\) |
| disk area | 10 mA cm⁻² | 1.749(2) | 2.1(1) mA cm⁻² |
| mass | 10 Ag⁻¹ | 1.705(3) | 5.3(4) Ag⁻¹ |
| oxide area | 100 µA cm⁻² | 1.714(3) | 42(3) µA cm⁻² |
| TOF-s \(b\) | – | – | 0.07(1) O₂ Mn⁻¹ s⁻¹ |
| TOF-b \(b\) | – | – | 0.002(2) O₂ Mn⁻¹ s⁻¹ |

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ChemSusChem 2017, 10, 4479 – 4490 www.chemsuschem.org 4483 © 2017 The Authors. Published by Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim.
face area that exhibit higher activity per oxide surface, for example, ball-milled Ba$_{0.5}$Sr$_{0.5}$CoO$_{2-x}$Fe$_{x/2}$O$_{3-δ}$ (1.53 V versus RHE; 3.9 m$^2$g$^{-1}$).\[\text{[25]}\]

Ideally, the current would be normalized by the number of active sites for comparison of the intrinsic activity of materials without geometric effects such as surface area and roughness.\[\text{[27]}\] However, the number of active sites is difficult to determine for oxide surfaces. Instead, two proxies are commonly used: normalization by the oxide area or the total number of metal ions either on the surface or in the bulk, from which the TOF is obtained. In both normalizations, it is assumed that all surface ions are active, which likely leads to underestimation of intrinsic activity, particularly if all bulk ions are considered. The LiMn$_2$O$_4$-carbon composite electrode reached 100 $\mu$Acm$^{-2}$ at 1.714(3) V versus RHE (0.485(3) V overpotential), which is similar to LaMnO$_3$/C (1.68 V versus RHE)\[\text{[29]}\] and Ba$_2$Mn$_3$O$_{12}$/C (1.66 V versus RHE)\[\text{[62]}\] particles and La$_{0.5}$Sr$_{0.5}$MnO$_3$ thin films (1.68(2) V versus RHE)\[\text{[61,62]}\] but clearly less active as compared to, for example, Ba$_{0.5}$Sr$_{0.5}$CoO$_{2-x}$Fe$_{x/2}$O$_{3-δ}$ thin films (1.53 V versus RHE).\[\text{[28]}\] The TOF of the LiMn$_2$O$_4$-carbon composite electrode was 0.00255(2) O$_2$Mn$^{-1}$s$^{-1}$ at 1.68 V versus RHE (0.45 V overpotential), which is higher than the TOF reported for $\beta$-MnO$_2$ (0.0012 O$_2$Mn$^{-1}$s$^{-1}$)\[\text{[66]}\] intermediate to electrodeposited MnO$_{n/2}$H$_{n}$ (0.0033(2) = 0.007 O$_2$Mn$^{-1}$s$^{-1}$),\[\text{[61]}\] and less than the TOF of Mn oxides with higher surface area such as $\alpha$-MnO$_2$-SF (0.0047 O$_2$Mn$^{-1}$s$^{-1}$).\[\text{[66]}\] The TOF-s of the LiMn$_2$O$_4$-carbon electrode is more than an order of magnitude higher (0.07(1) O$_2$Mn$^{-1}$s$^{-1}$) than the TOF-b. The TOF can also be obtained for the CaMn$_2$O$_4$ cofactor of PSII, which approximately works at an overpotential of 0.3 V\[\text{[72]}\] and has a TOF-s$=TOF$-b$ \approx$ 100 O$_2$Mn$^{-1}$s$^{-1}$ at that lower overpotential.\[\text{[2]}\] It should be noted that the nature of the catalyst, environmental conditions, and driving force for catalysis differ between LiMn$_2$O$_4$ and PSII.

We conclude that composite electrodes of LiMn$_2$O$_4$-carbon exhibited respectable electrocatalytic activity in any metric. LiMn$_2$O$_4$ shares the cubane structural motif and average Mn$^{3.5+}$ valence with the dark-stable $S_1$ state of natural photosynthesis as we discussed above. These commonalities raise the question whether there are parallels between the photocatalytic mechanism of PSII and the electrocatalytic mechanism of LiMn$_2$O$_4$.

The electrocatalytic mechanism of oxygen evolution is defined by three partial derivatives that relate the key parameters of the reaction kinetics:\[\text{[27]}\]

$$\left(\frac{\partial E}{\partial \phi}\right)_i = -\left(\frac{\partial E}{\partial \log [i]}\right)_{\phi} \times \left(\frac{\partial \log [i]}{\partial \phi}\right)_E$$

The subscripts indicate constant variables for which additionally temperature and pressure should be held constant. From left to right, these partial derivatives are known as the Nernst slope ($\partial E$/\partial $\phi$), the Tafel slope ($\partial E$/\partial $\log [i]$), and the reaction order with respect to $\phi$ ($\log [i]$/\partial $\phi$). The reaction order with respect to $\phi$ is identical to that with respect to $\log [OH^-]$ and $-\log [H^+]$ ($\phi = -\log [H^+] = 14 - pOH = 14 + \log [OH^-]$). We discuss the reaction order with respect to hydroxide because hydroxide is the reactant in alkaline electrolytes (4OH$^- + O_2 + 4e^- + H_2O$). Furthermore, determination of all of the partial derivatives can lift ambiguities in mechanistic assignments that exist in assignments based on a single derivative.

The values of these partial derivatives depend on the choice of the reference potential with the exception of the Tafel slope. Here, we provide the values for voltages relative to both NHE and RHE. The NHE scale is commonly employed in the field of solar fuels, whereas the RHE scale is common in the field of technical electrocatalysis. Voltages on the NHE scale are proportional to the Gibbs free energy of the reaction but the potential of water oxidation depends on $\phi$, whereas it is 1.23 V under standard conditions on the RHE scale independent of $\phi$.

The Nernst slope ($\nu = \partial E$/\partial $\phi$) is 0 mV$^{-1}$ph on the RHE scale and $-59$ mV$^{-1}$pH on the NHE scale when electron and proton/hydroxide transfers are coupled. At standard conditions, the Nernst slope on these two scales can be converted using:\[\text{[25,73]}\]

$$(\partial E_{\text{NHE}}/\partial \phi)_i = (\partial E_{\text{RHE}}/\partial \phi)_i + 59 \text{ mV} \text{pH}^{-1}$$

where $E_{\text{NHE}}$ is the voltage on the NHE scale and $E_{\text{RHE}}$ is the voltage on the RHE scale. The Nernst slope approaches 59 mV$^{-1}$pH on the RHE scale and 0 mV$^{-1}$pH on the NHE scale when the investigated electrochemical process becomes independent of $\phi$, that is, electron and proton/hydroxide transfer become decoupled and an imbalanced number of electron and proton/hydroxide are transferred.

The Tafel slope ($b = \partial E$/\partial $\log [i]$) is perhaps the most widely investigated among the three partial derivatives. It does not require $\phi$-dependent studies and is independent of the choice of the reference potential. The value of the Tafel slope depends mainly on the ratio of electrons transferred before the RLS to electrons transferred after the RLS but also on the surface coverage of the intermediates\[\text{[27,30]}\] and the reorganization energy of the surroundings of the active site.\[\text{[74]}\] The Tafel slope can be used as a diagnostic for a given RLS in a mechanistic sequence, for example, a value of 59 mVdec$^{-1}$ indicates that there is an electron transfer prior to a limiting chemical step under the assumption of low surface coverage (no blocking) and metallic conduction (fast electron transfer). Most values of the Tafel slope cannot be uniquely assigned to a RLS in a given mechanistic sequence.

The reaction order with respect to $\phi$ ($\rho = \partial \log [i]/\partial \phi$) depends on the reference potential. The reaction orders on the NHE and RHE scale can be converted by

$$\rho_{\text{NHE}} = \rho_{\text{RHE}} + (59/b) \text{ mV dec}^{-1}$$

where $b$ is the value of the Tafel slope in the investigated $\phi$ range. The expected reaction order of an electrochemical process that does not depend on $\phi$ is $\rho = 0$. On the RHE scale, this is obtained at constant overpotential of oxygen electrocatalysis, whereas the free energy is constant on the NHE scale. Reported reaction orders of the OER spread considerably in the literature\[\text{[76]}\] and can have fractional values, which may be owed to side reactions or competing intermediates on the
was obtained on the RHE scale. We concluded above that the scaled ring currents correspond to the catalytic rate of oxygen evolution and therefore, we focused on the analysis of the mechanistic parameters of the ring currents but also provide the disk currents for comparison with the literature.

The Nernst slope ($\nu = \Delta E / \Delta \log i \Delta \log pH$) of the scaled ring was $1(1)$ mV pH$^{-1}$ on the RHE scale and $-60(1)$ mV pH$^{-1}$ on the NHE scale. The Nernst slopes at the disk were $-2(2)$ and $-63(2)$ mV pH$^{-1}$ on the RHE and NHE scales, respectively. They were thus identical within experimental uncertainty in the investigated current range between 30 and 130 mA (Figure 7c). A Nernst slope of $-71(1)$ mV pH$^{-1}$ was previously determined for electrodeposited MnO$_2$, between pH 11.4 and 13.3,$^{(25)}$ which is identical to the Nernst slope at the disk within experimental uncertainty. The Nernst slope depended on the current above 130 mA. It might be possible that the scan rate was too fast to ensure quasi-equilibrium at these higher currents and thus reaction rates. Alternatively, the transfer of electrons and hydroxide could have been decoupled owing to hydroxylation of neighboring sites (i.e., breakdown of the low coverage condition).

The disk currents show a clear transition to another region of constant Nernst slope at currents below 10 mA. The Nernst slope of $-39$ mV pH$^{-1}$ on the NHE scale suggested a process, in which 3 electrons and 2 protons/hydroxide are transferred. Possible reactions involving MnO$_2$ and MnO$_2^2$- have more negative Nernst slopes, and oxidation to HMnO$_4^-$ has an equilibrium potential much higher than the upper scan boundary (Table 3). It is conceivable that delithiation competes with deprotonation as the onset voltage of LiMnO$_4$ delithiation is near 1.5 V versus RHE in brine.$^{(76)}$ This analysis illustrated the diagnostic value of the Nernst slope at the disk to identify and exclude side reactions.

The reaction order with respect to OH$^-$ ($\nu = \Delta E / \Delta \log i \Delta \log pH$) was $-0.04(2)$ on the RHE scale between 1.59 and 1.66 V on the scaled ring and 0.23(2) on the capacitance-corrected disk between 1.52 and 1.58 V (Figure 7d). In the voltage range of the ring, the reaction order of the disk decreases toward negative values, which could be due to changes in the Tafel slope or a side reaction such as the aforementioned delithiation.

**Figure 7.** Key mechanistic parameters of LiMnO$_4$-carbon composite disks (small circle, solid lines) and corresponding scaled ring currents (large circles, dotted lines) of oxygen reduction. Tafel plot on the (a) RHE and (b) NHE scales. (c) Nernst slope ($\Delta E / \Delta \log i / \Delta \log pH$) as function of current. Reaction order with respect to OH$^-$ ($\Delta E / \Delta \log i / \Delta \log pH$) on the (d) RHE and (e) NHE scales. (f) Tafel slopes ($\Delta E / \Delta \log i / \Delta \log pH$) as function of pH. All electrochemical measurements were conducted in Ar-purged NaOH at 10 mVs$^{-1}$ and 1600 RPM. Error bars indicate the standard deviation of 3 electrodes. Thin dotted lines indicate constant derivatives and the shaded area indicates where $\Delta E / \Delta \log i / \Delta \log pH$ and $\Delta E / \Delta \log i / \Delta \log pH$ are simultaneously constant.

**Table 3.** Predicted Nernst slopes and observed Nernst slopes of LiMnO$_4$.

| Reaction[a] | $\Delta E / \Delta \log i / \Delta \log pH$ | $\Delta E / \Delta \log i / \Delta \log pH$ | $\Delta E / \Delta \log i / \Delta \log pH$ |
|-------------|---------------------------------|---------------------------------|---------------------------------|
| $4OH^- + 4e^- + 2H_2O$ | 0.401 | -59 | 0 |
| $MnO_2 + 4OH^- \rightarrow MnO_4^- + 3e^- + 2H_2O$ | 0.595 | -79 | -20 |
| $MnO_2 + 4OH^- \rightarrow MnO_4^- + 2e^- + 2H_2O$ | 0.600 | -118 | -59 |
| $MnO_2 + 2OH^- \rightarrow HMnO_4^- + 2e^- + 3H_2O$ | 2.09 | -39 | 20 |
| $LiMnO_4 - Li^+ + MnO_2$ | $\approx 0.9^{[7]}$ | 0 | 59 |
| observed for LiMnO$_4$ (0.74–0.81 V versus NHE) | n/a | -63(1) | -2(1) |
| observed for LiMnO$_4$ (< 0.72 V versus NHE) | n/a | -4(1) | 20(1) |

[a] Based on Ref. [77]. [b] Equilibrium potential on the NHE scale.$^{(77)}$ [c] Uncertainties (1σ) of the fit to a constant are indicated. [d] Nernst slope of the disk on the NHE scale. [e] Nernst slope of the disk on the RHE scale. [f] In aqueous brine against Ag counter.$^{(78)}$ The estimated experimental onset of delithiation from LiMnO$_4$ is $\approx 0.7$ V versus NHE ($\approx 1.5$ V versus RHE).
Converted reaction orders with respect to OH\(^-\) of 0.19, 0.13, 0.18, 0.18, and 0.18 on the RHE scale were previously reported for disks of La\(_{0.5}\)(Ca\(_{0.5}\))Mn\(_2\)O\(_4\), La\(_{0.5}\)Sr\(_{0.5}\)Mn\(_2\)O\(_4\), La\(_{0.5}\)K\(_{0.5}\)Mn\(_2\)O\(_4\), La\(_{0.5}\)Sr\(_{0.5}\)Mn\(_2\)O\(_4\), and BaMn\(_2\)O\(_4\) pellets,\(^{30,79}\) respectively. These values are similar to the reaction order at the LiMn\(_2\)O\(_4\) disk.

The reaction order on the NHE scale had an average value of 0.83(1) both on the scaled ring between 0.74 and 0.81 V versus NHE and on the capacitance-corrected disk between 0.70 and 0.79 V versus NHE (Figure 7e). Using Equation (3), reaction orders of 0.91(3) and 0.94(4) can be calculated for ring and disk. The Tafel slopes of disk and ring were not constant at pH 13 below 0.81 V versus NHE, which might cause the deviation between the calculated and measured values. The shape of the reaction order traces on the NHE and RHE scales differed, which is a consequence of the differing current profiles as function of pH on the NHE and RHE scales. The reader is referred to the in-depth discussion of Koper.\(^{76}\) Reaction orders with respect to OH\(^-\) reported on the NHE scale differ greatly from −1.2(1) for electrodeposited Mn\(_2\)O\(_4\),\(^{27,29}\) to −0.5 for MnO\(_2\) paste\(^{76}\) and 0.65 for LaMnO\(_3\) pellets.\(^{30}\) The spread may be partially caused by the Tafel slopes being 52(8),\(^{76}\) 60(3) mV dec\(^{-1}\), and 83(1) mV dec\(^{-1}\), respectively.

The constants of current Nernst slope and voltages of constant reaction order defined a region in the Tafel plot, in which neither current nor voltage depended on pH (shaded areas in Figures 7a, b). On the RHE scale, the data of all investigated pH values fell into this region, whereas the data at pH 13 fell outside on the NHE scale. This dependence of the coupled mechanistic parameters [Eq. (1)] could contribute to the large spread of reported reaction orders and Tafel slopes on the NHE scale. The currents at which the pH scale can be evaluated independent of pH on the RHE scale spanned less than an order of magnitude. Although it is somewhat arbitrarily recommended to evaluate Tafel slopes over more than 2 current magnitudes,\(^{27}\) this criterion appears to be too strict for a complex reaction such as the OER investigated by conventional macroscopic electrodes. Instead, we propose to evaluate the Tafel slope where the Nernst slope and reaction order are constant.

The values of the Tafel slope on the scaled ring and the capacitance-corrected disk were both independent of pH within experimental uncertainty (Figure 7f), yet had different values. The ring Tafel slope was 62(1) mV dec\(^{-1}\), whereas the disk Tafel slope was 83(1) mV dec\(^{-1}\). Disk Tafel slopes between 70 and 90 mV dec\(^{-1}\) are typical for composite electrodes containing carbon; Tafel slopes of 88, 70, and 85 mV dec\(^{-1}\) were reported for MnO, Mn\(_3\)O\(_7\), and MnO\(_2\), respectively.\(^{80}\) The significance of these values is unknown and they do not correspond to commonly discussed mechanisms of the OER.\(^{27,30,74}\) Aside from intermediate surface coverage on these oxides (i.e., not low coverage or full coverage),\(^{27,30}\) it is conceivable that the Tafel slope is influenced by Mn proportionation,\(^{81}\) which would affect the number of electrons transferred prior to the RLS. Yet, Tafel slopes of about 60 mV dec\(^{-1}\) were also previously reported for diverse manganese oxides such as composite electrodes of MnO\(_2\) and carbon (61 mV dec\(^{-1}\)), electrodeposited MnO\(_2\) (60(3) mV dec\(^{-1}\)),\(^{79}\) and epitaxial La\(_{0.5}\)Sr\(_{0.5}\)Mn\(_2\)O\(_4\) disks (65(5) mV dec\(^{-1}\)).\(^{61}\) The latter was also studied on a RRDE setup, where the ring Tafel slope was 66(6) mV dec\(^{-1}\) (the values are given for setup used at the Institute of Materials Physics (IMP) in Ref. [61], which is identical to the one used in this study). These literature values further corroborate that the ring Tafel slope of 62(1) mV dec\(^{-1}\) is representative of oxygen evolution on LiMn\(_2\)O\(_4\).

Interpretation of mechanistic parameters of the Nernst slope, reaction order, and Tafel slope calls for proposing a suitable reaction mechanism. Historically, the three mechanistic parameters are mainly discussed in the context of four mechanistic sequences\(^{27,30}\) that are based on the works of Bockris,\(^{29}\) Krasil’shikov,\(^{62}\) O’Grady,\(^{83}\) and Kobussen.\(^{84,85}\) The latter mechanistic sequence is also used in contemporary theoretical work\(^{86,87}\) and has received much attention lately.

Instead of postulating a mechanistic sequence based on the classic reaction sequences (in the field of oxygen electrolysis), we take a bio-inspired approach by interpreting the mechanistic parameters of LiMn\(_2\)O\(_4\) in the context of the well-established mechanism of natural photosynthesis (Figure 1). This approach suggests itself due to the structural and chemical similarities between the CaMn\(_2\)O\(_4\) cluster of PSII and LiMn\(_2\)O\(_4\) established herein.

In the following Gedankenexperiment, let us assume natural photosynthesis was an electrocatalytic process with a reaction sequence identical to that of the actual biocatalytic process (Table 4). We further assume that the CaMn\(_2\)O\(_4\) cluster of PSII is an electrochemical surface with low hydroxide coverage in an alkaline electrolyte and exhibits quasi-equilibrium between the reaction steps. In natural photosynthesis, it is undisputed that the transition from S\(_2\) (via the elusive S\(_3\)) to S\(_3\) is the slowest step (time constant \(\approx 1.5\) ms) and thus rate-limiting (Table 4).\(^{2,10,12,20-23}\) Starting from the dark-stable state S\(_1\), two
electrons are transferred before the limiting step \( (n_p) \), that is, during \( S_i^{2+} \rightarrow S_i^{3+} \) and \( S_i^{3+} \rightarrow S_i^{4+} \). One electron is transferred during the limiting step \( (n_o) \). Using the formula derived by Fletcher:[24]

\[
b = \frac{59}{[n_p + 0.5 n_o (1 - \Delta)]},
\]  

(4)

an electrocatalytic surface would have a Tafel slope of \( 24/(1 - \Delta) \) \( \text{mV} \text{dec}^{-1} \) or a lower Tafel slope of \( 24 \) \( \text{mV} \text{dec}^{-1} \) if the positive reorganization energy of the double layer is neglected. It is worth noting that low Tafel slopes are highly desirable for electrocatalytic applications. A Tafel slope of \( 24 \) \( \text{mV} \text{dec}^{-1} \) has recently been reported for the OER on FeNi layered double hydroxide on Ni foam.[92] However, it is significantly lower than the Tafel slope of \( 62(1) \) \( \text{mV} \text{dec}^{-1} \) observed for \( \text{LiMn}_2\text{O}_4 \).

We also calculated non-vanishing reaction orders with respect to \( \text{OH}^- \) on the NHE and RHE scales for the RLS of the photosynthetic mechanism as an electrocatalytic process. On the NHE scale, each step that includes a hydroxide transfer increases the reaction order by one because the formation of these steps depends on the hydroxide concentration and the population of preceding steps. This pedestrian derivation is supported by the more rigorous analysis of Shinagawa et al.,[93] who found that the reaction rate depends on the concentrations of the hydroxide concentration with a polynomial degree equivalent to the steps that included hydroxide transfer. Thus, the co-factor of PSII as an electrocatalytic surface would have a reaction order with respect to \( \text{OH}^- \) on the NHE scale of \( \mu_{\text{NHE}} = 3 \), which corresponds to a reaction order of \( \mu_{\text{RHE}} = 0.5 \) on the RHE scale when the reorganization energy is neglected. This also disagrees with the observed reaction orders on \( \text{LiMn}_2\text{O}_4 \).

Tafel slopes and reaction orders were also calculated under the assumption that any of the other steps in the extended S-state cycle[23] are rate-limiting (Table 4). Equation (4) indicates that the Tafel slope decreases when more electrons are transferred before the RLS or when an electron is transferred during the RLS. The step \( S_i^{2+} \rightarrow S_i^{3+} \) matches the experimentally obtained Tafel slope and reaction order of \( \text{LiMn}_2\text{O}_4 \). However, any process would produce a Tafel slope of \( 39 \) \( \text{mV} \text{dec}^{-1} \) when one electron is transferred prior to a RLS without electron transfer. This step could be breaking of the \( \text{O} \cdots \text{H} \) bond (as in \( S_i^{2+} \rightarrow S_i^{3+} \)), making of the \( \text{O} \cdots \text{O} \) bond or breaking the \( \text{Mn} \cdots \text{O} \) bond (oxygen release). However, \( \text{O} \cdots \text{H} \) bond breaking should have fast kinetics for high concentrations of \( \text{OH}^- \) (\( \text{pK}_a = 14 \) in alkaline electrolytes) and thus dioxygen formation or release are most likely limiting. It should be noted that the details of the latter two steps are still debated for natural photosynthesis.[9,12,20-23,94,95]

The reaction order of the \( S_i^{2+} \rightarrow S_i^{3+} \) transition of 0 on the RHE scale also matches the value of \(-0.04(2)\) observed for \( \text{LiMn}_2\text{O}_4 \) within 2 standard deviations. In natural photosynthesis, the electron transfer occurs first (\( S_i^{3+} \rightarrow S_i^{4+} \)), followed quickly by deprotonation (\( S_i^{2+} \rightarrow S_i^{3+} \)). The sequence of events is possibly identical during electrocatalysis on \( \text{LiMn}_2\text{O}_4 \) due to an identical Mn valence as well as cubane structure of \( \text{LiMn}_2\text{O}_4 \) compared to the \( S_i \) state and the presence of the base hydroxide in the electrolyte. Although the electrochemical reaction order can give no insight into the sequence, the obtained values clearly indicate the transfer of a hydroxide.

The combined mechanistic parameters suggest coupled transfer of electrons and hydroxide. \( \text{LiMn}_2\text{O}_4 \) showed a Nernst slope of \( 1(1) \) \( \text{mV} \text{pH}^{-1} \) on the RHE scale (\( -60(1) \) \( \text{mV} \text{pH}^{-1} \) on the NHE scale), which indicates coupling of electron and hydroxide transfer. The obtained Tafel slope mandates the transfer of one electron and the reaction order the transfer of one hydroxide. It could be transferred either before or during the RLS. As braking of \( \text{O} \cdots \text{H} \) bonds is unlikely involved in the RLS, the mechanistic parameters point toward a coupled electron and hydroxide transfer before the RLS.

All discussed data point toward an early RLS in the mechanism of \( \text{LiMn}_2\text{O}_4 \) compared to natural photosynthesis. In the S-state cycle of photosynthesis, two coupled charge transfers occur before the RLS that includes an additional coupled charge transfer (Table 4). In contrast, \( \text{LiMn}_2\text{O}_4 \) has only a single coupled charge transfer before a RLS without electron transfer. PSII contains only a single cubane unit embedded in the protein matrix. Recent diffraction experiments[96] suggest the oxygen marked by an asterisk in Figure 1 as the site of \( \text{O} \cdots \text{H} \) bond formation. In \( \text{LiMn}_2\text{O}_4 \), comparable sites are available at any corner of the cubane and the surface was slightly oxidized. This should make it more likely that a single electron transfer sufficiently oxidizes the surrounding of a potential active site in \( \text{LiMn}_2\text{O}_4 \), the bulk of which consists of equal numbers of randomly distributed \( \text{Mn}^{3+} \) and \( \text{Mn}^{4+} \) as supported by the non-charge ordered structure of our \( \text{LiMn}_2\text{O}_4 \) powder (Figure 2a; Table 1).

Oxygen evolution reduces Mn in natural photosynthesis and electrocatalytic Mn oxides.[90] Thus, an early limiting step in the mechanism of \( \text{LiMn}_2\text{O}_4 \) means that further oxidation steps are required to return the electrocatalyst to the initial state. These steps are likely fast at the highly oxidizing voltages of the OER because voltages above approximately 0.8 V versus RHE were found sufficient to oxidize \( \text{Mn}^{3+/4+} \) oxide,[50] whereas the mechanistic parameters were evaluated above 1.58 V versus RHE (i.e., > 0.78 V overpotential with respect to \( \text{Mn}^{2+/3+} \) oxidation). Changes in the RLS of electrocatalytic oxides have recently been demonstrated by using electrolyte additives[97] or surface functionalization.[98] These approaches might enable the later fast steps in the mechanistic sequence to enhance reaction kinetics. Furthermore, control over the electrons transferred prior to the RLS allows reducing the Tafel slope.

We conclude that the steps of the S-state cycle of natural photosynthesis can explain the observed mechanistic parameters on \( \text{LiMn}_2\text{O}_4 \). However, we cannot judge the validity of our assumption that the mechanistic sequence of these two catalysts is identical, which would require elaborate in situ experiments to test the proposed hypothesis.

**Conclusions**

We discussed \( \text{LiMn}_2\text{O}_4 \) as an electrocatalytic model for the active site of natural photosynthesis, that is, the \( \text{CaMn}_2\text{O}_4 \) co-factor of photosystem II, to gain comparative insight into the
mechanism of the oxygen evolution reaction (OER). By ex situ soft X-ray absorption spectroscopy (XAS), we established a valence of Mn\(^{3.5+}\) akin to the dark-stable S1 state in the mechanism of natural photosynthesis. Mn dissolution was negligible after 5 cycles, which was established using rotating-ring disk electrode (RRDE) measurements and high-resolution TEM. LiMn\(_2\)O\(_4\) nanoparticles show respectable activity in all the main metrics, namely 0.520(2) V overpotential at 10 mA cm\(^{-2}\), 1.705(3) V versus reversible hydrogen electrode (RHE) at 100 µA cm\(^{-2}\) and a surface turnover frequency of 0.07(1) O\(_2\)Mn\(^{3-}\)s\(^{-1}\) at 0.45 V overpotential. Additional RRDE measurements were used to probe the oxygen evolved at the disk. Using the catalytically relevant ring currents, we obtained a Tafel slope of 62(1) mV dec\(^{-1}\), Nernst slope of 1(1) mV pH\(^{-1}\), and reaction order with respect to OH\(^-\) of $(-0.04(2)$ at low overpotentials on the RHE scale. The Nernst slope ($\sim$60(1) mV pH\(^{-1}\)) and reaction order ($0.83(1)$) were also evaluated on the normal hydrogen electrode (NHE) scale. These mechanistic parameters indicate a rate-limiting step (RLS) without electron transfer that is preceded by a single electron transfer, which is in contrast to natural photosynthesis, for which three holes are accumulated prior to oxygen release. It should be noted that an early RLS does not exclude a mechanism in which holes (i.e., oxidation equivalents) are accumulated as the pristine material is already in a high oxidation state (Mn\(^{3.5+}\)). We calculated the Tafel slope and reaction order under the assumption that the S-state cycle of natural photosynthesis is an electrochemical process. The late RLS of the active site in natural photosynthesis would result in a desirable low Tafel slope of about 24 mV dec\(^{-1}\). This suggests that engineering materials with a photosystem-like mechanism having a late RLS (i.e., more than electron transfer before the RLS) provide low Tafel slopes at low overpotential. Our mechanistic analysis thus outlines a strategy to lower the Tafel slope of bio-inspired electrocatalysts for the production of sustainable fuels with high efficiency.

**Experimental Section**

LiMn\(_2\)O\(_4\) was bought from Sigma-Aldrich (Art. #725129, LOT #MKBF0675V) and used as received.

Powder XRD patterns were obtained with a Bruker D8 diffractometer using monochromatized CuK\(_\alpha\) radiation. SEM was performed using an FEI Nova Nano SEM 650 operated at 5 and 15 kV, where a through-lens detector was used. TEM studies were performed using a FEI Titan 80–300 environmental microscope operated at 300 kV in high vacuum mode. The analyzed samples were drop cast onto a lacey carbon film fixed on a 3 mm copper grid directly before the TEM experiments.

Soft XAS measurements at the MnL and OK edges were performed at the spherical grating monochromator (SGM) beamline 11ID-1 at the Canadian Light Source.\(^{[100]}\) Samples were prepared by covering the glassy carbon electrode (GCE) with a thin film of pellets (Applichem, >99% in milliQ water. Electrolytes with pH between 13 and 13.4 were prepared by diluting the stock solution with milliQ water. MnO\(_2\) was detected at the ring of the RRDE by reduction at 0.2 V versus SCE in NaOH purged with O\(_2\) (Air Liquid, 99.999%). Oxygen was detected at the ring of the RRDE by reduction at $-0.4$ V versus SCE in NaOH purged with Ar (Air Liquid, 99.999%).

The experimental protocol was identical for all ink measurements and pH values. The electrode was conditioned at 0.25 V versus SCE for 2 min and cyclic voltammetry between 0.25 and 0.75 V versus SCE with 2 mV per step was performed at 10 mVs\(^{-1}\) for 10 cycles with the ring either at 0.2 or $-0.4$ V versus SCE. The electrolyte
was allowed to rest for 2 min at open circuit (OC) and then electrochemical impedance spectroscopy (EIS) was performed between 100 kHz and 1 Hz with an AC amplitude of 25 mV at OC to determine the uncompensated resistance. After IR correction of the voltage, the data were interpolated either on an equidistant voltage axis with 5 mV resolution or an equidistant log current axis with 20 points per decade. Reaction orders were obtained on the equidistant voltage data, whereas the Tafel and Nernst slopes were obtained on the equidistant log current data.

Acknowledgements

Financial support by the CRC 1073 (projects C05 and 202) of the Deutsche Forschungsgemeinschaft (DFG) is gratefully acknowledged. Research described in this work was performed at the Canadian Light Source, which is supported by the Natural Sciences and Engineering Research Council of Canada, the National Research Council Canada, the Canadian Institutes of Health Research, the Province of Saskatchewan, Western Economic Diversification Canada, and the University of Saskatchewan.

Keywords: manganese oxide • reaction mechanisms • solar fuels • water oxidation • water splitting

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