Understanding Interactions between Manganese Oxide and Gold That Lead to Enhanced Activity for Electrocatalytic Water Oxidation

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ABSTRACT: To develop active nonprecious metal-based electrocatalysts for the oxygen evolution reaction (OER), a limiting reaction in several emerging renewable energy technologies, a deeper understanding of the activity of the first row transition metal oxides is needed. Previous studies of these catalysts have reported conflicting results on the influence of noble metal supports on the OER activity of the transition metal oxides. Our study aims to clarify the interactions between a transition metal oxide catalyst and its metal support in turning over this reaction. To achieve this goal, we examine a catalytic system comprising nanoparticulate Au, a common electrocatalytic support, and nanoparticulate MnO$_{x}$, a promising OER catalyst. We conclusively demonstrate that adding Au to MnO$_{x}$ significantly enhances OER activity relative to MnO$_{x}$ in the absence of Au, producing an order of magnitude higher turnover frequency (TOF) than the TOF of the best pure MnO$_{x}$ catalysts reported to date. We also provide evidence that it is a local rather than bulk interaction between Au and MnO$_{x}$ that leads to the observed enhancement in the OER activity. Engineering improvements in nonprecious metal-based catalysts by the addition of Au or other noble metals could still represent a scalable catalyst as even trace amounts of Au are shown to lead a significant enhancement in the OER activity of MnO$_{x}$.

INTRODUCTION

Electrochemical water oxidation, also known as the oxygen evolution reaction (OER), is a key energy conversion reaction in a number of clean energy technologies, including rechargeable metal–air batteries, electrolysis cells, and solar fuel production.1,2 Widespread commercialization of these technologies is limited in part by the scarcity and high cost of the best known catalysts for the OER, ruthenium and iridium oxides.3–5 Nickel oxides (NiO$_{x}$) present a viable alternative to precious metal oxides in alkaline environments and are currently used in commercially available alkaline electrolysers.6 Other nonprecious metal oxides, including manganese oxides (MnO$_{x}$) and cobalt oxides (CoO$_{x}$), have also demonstrated promising OER activity under alkaline conditions.6–19 To facilitate the development of nonprecious metal OER catalysts with improved activities, it is necessary to identify the specific catalytic sites that participate in the reaction and accurately determine their turnover frequencies. Although isolating site-specific turnover frequencies can be challenging in oxide electrocatalysts, the likelihood of success can be improved if the catalytic activity is studied on well-defined materials deposited on inert supports.20

There have been conflicting reports in the recent literature regarding the role metal supports play in the OER activity of metal oxide catalysts. A study, which used Pt(111) and Au(111) single crystal surfaces as supports for OER catalysts, demonstrated that the OER activity of four first row transitional metal oxides did not vary with the nature of the metal support and was linearly dependent on the coverage of the support by the metal oxide.6 Other reports, however, have shown that the OER activities of nickel,14 cobalt,15 and manganese oxides22,23 were influenced by the nature of the underlying support, and that the OER turnover frequency of a bulk metal oxide was inferior to that of a submonolayer amount of the same metal oxide deposited on a metal support.17,21 Our study, which focuses on one particular metal–supported transition metal oxide system, MnO$_{x}$/Au, conclusively demonstrates that adding a noble metal to a metal oxide OER catalyst can have a significant impact on its electrocatalytic activity. We show that this impact cannot be explained simply by surface area effects and we identify interesting changes in the red-ox properties of both MnO$_{x}$ and Au when the two materials are present in one composite catalyst. Using our experimental data and previous literature results, we develop a hypothesis about potential OER active sites.

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EXPERIMENTAL SECTION

Preparation of rotating disk electrode substrates: Rotating disk electrode substrates were prepared from 200 mm long glassy carbon (GC) rods (diameter 5 mm, Sigradur G HTW Hochtemperatur-Werkstoffe GmbH). Before deposition of MnO or Au nanoparticles, the rods were processed by Stanford crystal shop to produce 4 mm long pieces with one side lapped and chamfered and the other side polished to a root mean square (RMS) surface roughness of less than 50 nm.

Preparation of substrates for in situ X-ray absorption spectroscopy (XAS) characterization: Electrode substrates for in situ XAS characterization were prepared from 200 mm long GC rods (diameter 10 mm, Sigradur G HTW Hochtemperatur-Werkstoffe GmbH). Before deposition of MnO or Au nanoparticles, the rods were processed by the Stanford crystal shop to produce ca. 100–200 μm thick wafers with one side lapped and the other side polished to a surface RMS roughness of less than 50 nm.

Synthesis of catalyst materials: MnO nanoparticles were prepared with a sputtering system (Nanossys500, Mantis Deposition Ltd.) using a glow discharge magnetron sputtering with an inert gas condensation beamline 10 Lightsource (SSRL).

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In situ X-ray absorption spectroscopy (XAS) measurements were carried out at the SSRL at beamline 6-2 E52. The beamline was operated with a double-crystal Si(311) monochromator with a Rh-coated mirror to reject the high order harmonics. The X-ray beam was focused with a parabolic mirror to a spot size of 230 μm fwhm (V) by 400 μm fwhm (H) at the sample position. The X-ray energy was calibrated to the pre-edge absorption peak (6543.4 eV) of potassium permanganate. The Mn Kα signals were resolved by 6 spherically bent analyzer crystals Ge (335) installed on the high energy resolution X-ray emission spectrometer28 and detected by a silicon drift detector in photon counting mode. Multiple scans were collected at different sample positions at the potentials of interest. All spectra were normalized to have an edge jump of unity after linear backgrounds were subtracted from the raw data. The error bars reported in the spectra and the residuals represent one standard deviation based on Poisson statistics and standard error propagation.

In situ electrochemical characterization was performed in a similar setup as reported previously.29 Completely separate electrochemical cells (custom built cells, Adams & Chittenden), reference electrodes (Ag/AgCl, BASi) and counter electrodes (100 mm length/5 mm diameter glassy carbon rod, Si gradur G HTW Hochtemperatur-Werkstoffe GmbH) were used with the MnO-GC and MnO/Au-GC samples to avoid the possibility of contaminating the MnO-GC sample with trace amounts of Au. The working electrodes (GC wafers with deposited nanoparticulate catalysts) were glued to the window of the electrochemical cell using epoxy. The schematic of the in situ electrochemical cell is shown in Supporting Information, Figure S1. Prior to performing XAS measurements, cyclic voltammograms were collected at 20 mV s\(^{-1}\) in nitrogen-saturated 0.1 M KOH electrolyte by first scanning the potential from 0.05 to 1.1 V and then from 0.05 to 1.65 V. To record in situ Mn XANES spectra at an OER relevant potential, the potential was scanned from 0.05 V to a vertex potential of 1.65 V at 20 mV s\(^{-1}\) and held at 1.65 V for 15 min. All measurements were iR-compensated to 85% and are reported vs RHE by assuming a potential shift of 0.960 V. The average measured resistance between the working and the reference electrodes was \(\sim 70 \Omega\).

Effect of Au: To further study the effect of Au on the OER activity of MnO-GC nanoparticles, additional electrochemical experiments were carried out. For these experiments, the potential was scanned on a MnO sample and bare GC support from 0.05 V to 1.8 V in a N\(_2\) saturated environment. Then 0.1 mM Hau(III)Cl\(_3\) hydrate (Sigma Aldrich, 99.9999% metals basis) was added to each electrolyte and the samples were repeatedly scanned starting in the OER region, 1.2 to 1.8 V, with the cathodic scan progressively reaching more reductive potentials with each cycle to deposit increasing amounts of Au on the surface of the catalyst.

RESULTS AND DISCUSSION

Because the influence of the underlying substrate on the OER activity of MnO\(_x\) has not yet been clearly established in literature, we prepared nanoparticulate catalysts on inert GC supports to directly evaluate the impact of Au. These nanoparticulate catalysts consisted of sputtered MnO catalysts28 and evaporated Au catalysts. The first sample consisted of only MnO deposited on GC (MnO-GC). The second and third samples consisted of both MnO and Au nanoparticles: one sample with MnO sputtered on top of Au nanoparticles evaporated onto GC (MnO/Au-GC), while in the other Au was evaporated on top of the MnO-GC catalyst (Au/MnO-GC).

Schematic representations and SEM images of the MnO-GC, MnO/Au-GC, and Au/MnO-GC catalysts are shown in Figure 4921.
1a,b. From the images, we can estimate that the MnO loading is less than 1 μg (calculations are presented in the Supporting Information). Total electron yield (TEY) Mn L-edge XAS characterization of the Mn surface oxidation state in the three catalysts is presented in Figure 2. In the figure, the XAS spectra of the composite samples consisting of both MnO and Au are compared to the spectra of MnO-GC and two powder controls: MnF2 and MnO. The surface of MnO, which is known to oxidize in air, was not sputtered prior to characterization and represents oxidized “MnO”.

of the composite samples consisting of both MnO and Au are compared to the spectra of MnO-GC and two powder controls: MnF2 and MnO. In these experiments, MnF2 was used as a MnII powder standard in addition to commercially purchased MnO powder because of the known surface oxidation of the MnO phase in air.27,30 Comparison of the spectra of the powder standards to the spectra of the as-deposited MnO/Au-GC and Au/MnO-GC catalysts using this surface-sensitive technique illustrates that after the addition of Au, MnO nanoparticles exhibit the oxidized form rather than the reduced form of the MnII phase. It is not clear whether this slight difference in the starting surface oxidation state results in a change in the crystal structure of the material to form a different bulk phase. Our attempts to characterize the crystallinity of the catalysts with and without the addition of Au using conventional and synchrotron grazing incidence X-ray diffraction (XRD) yielded only reflections corresponding to the GC support (Supporting Information, Figures S2 and S3). This indicated that either both types of samples were amorphous or that the crystallinity of the nanoparticles could not be detected using grazing incidence XRD.

Electrochemical characterization of the catalysts in 0.1 M KOH electrolyte shown in Figure 3 demonstrates that the composite catalysts, consisting of both MnO and Au nanoparticles, have significant OER activity, while the OER current density of the MnO-GC catalyst is negligible by comparison.

Figure 2. Total electron yield of Mn L-edge XAS of the three catalytic samples, MnO-GC, MnO/Au-GC, and Au/MnO-GC, and two powder standards, MnF2 and MnO. The surface of MnO, which is known to oxidize in air, was not sputtered prior to characterization and represents oxidized “MnO”.

Figure 3. Cyclic voltammetry characterization of the three catalytic samples, MnO-GC, MnO/Au-GC, and Au/MnO-GC. Characterization was performed in 0.1 M KOH electrolyte, at 20 mV/s, and 1600 rpm.
Table 1. OER Activities of MnOx Electro catalysts at $\eta = 300$ and 400 mV

| catalyst material/ type (author) | $I_{\eta=300}$ mA cm$^{-2}$ | $I_{\eta=300}$ A g$^{-1}$ (est.)<sup>a</sup> | TOF<sub>$\eta=300$</sub> s<sup>−1</sup> (est.)<sup>b</sup> | $I_{\eta=400}$ mA cm$^{-2}$ | $I_{\eta=400}$ A g$^{-1}$ (est.)<sup>a</sup> | TOF<sub>$\eta=400$</sub> s<sup>−1</sup> (est.)<sup>b</sup> |
|---------------------------------|-------------------------------|-----------------------------------|--------------------------|---------------------|-----------------------------------|--------------------------|
| MnO/Au-GC (this work)           | 0.09                          | 100                               | 0.01                     | 0.23                | 200                               | 0.03                     |
| Au/MnO-GC (this work)           | 0.04                          | 30                                | 0.006                    | 0.14                | 100                               | 0.02                     |
| MnO-GC (this work)              | 0.007                         | 6                                 | 0.001                    | 0.04                | 10                                | 0.002                    |
| $\beta$-MnO$_2$-$\alpha$-Mn$_2$O$_3$ (Morita<sup>c</sup>) | N/A                           | N/A                               | N/A                     | N/A                 | N/A                               | N/A                     |
| $\gamma$-MnOOH/Au (El-deab<sup>d</sup>) | N/A                           | N/A                               | N/A                     | N/A                 | N/A                               | N/A                     |
| $\alpha$-Mn$_2$O$_3$ (Gorlin<sup>e</sup>) | 1                             | 6                                 | 0.002                    | 2.34                | 10                                | 0.006                    |
| MnO-ALD (Pickrahm<sup>f</sup>)  | 0.4                           | 20                                | 0.003                    | 0.61                | 30                                | 0.005                    |
| MnOOH/Pt(111) (Subbaraman<sup>g</sup>) | N/A                           | N/A                               | N/A                     | N/A                 | N/A                               | N/A                     |
| Mn$_2$O$_3$ (Trottochaud<sup>h</sup>) | 0.002                         | 2                                 | 0.0004                   | N/A                 | N/A                               | N/A                     |
| $\beta$-MnO$_2$ (Fekefe<sup>i</sup>) | N/A                           | N/A                               | N/A                     | N/A                 | 6                                 | 0.4                      | 0.0001                   |

<sup>a</sup>Details of the calculations are presented in the Supporting Information. <sup>b</sup>Details of the calculations. <sup>c</sup>The authors report TOF for $\eta = 600$ mV.

The Supporting Information (Figure S4) also presents characterization of Au evaporated onto GC in the absence of MnO (Au-GC) to rule out simple additive effects of MnO-GC and Au-GC. From the inset in Figure S4 it is clear that the OER current of the composite samples is significantly greater than the sum of the individual activities of MnO-GC and Au-GC. Table 1 compares the OER activity metrics of MnO-GC, MnO/Au-GC, and Au/MnO-GC, including geometric current density, mass activity, and turnover frequency (TOF), all calculated at either 300 or 400 mV overpotential ($\eta$), to the activity metrics of some of the best MnOx electrocatalysts reported in literature. Although Table 1 focuses purely on electrocatalysts, Supporting Information, Table S1 presents a similar comparison to exceptional manganese oxide photochemical water oxidation catalysts. The tabulated activity metrics for manganese oxide catalysts reveal that the catalytic behavior of MnO-GC in the absence of Au is consistent with expectations. Although its current density normalized to the geometric surface area is low in magnitude, the mass activity and TOF values of the catalyst match the performance of the best MnOx OER catalysts. Furthermore, normalization of MnO-GC OER current by its capacitance produces similar surface-area normalized activity to that of the previously reported MnO thin film catalyst (Supporting Information, Figure S6) deposited by atomic layer deposition (ALD).31

From Tables 1 and S1, it is also apparent the two composite catalysts consisting of both MnO and Au nanoparticles exhibit unusually high OER activities. Each mass activity and TOF is about an order of magnitude higher than the corresponding metrics for the previously reported highly active thin film MnOx catalysts. The only other MnOx catalyst with similar activity was published by Subbaraman and co-workers, who electro-deposited MnOx islands on a Pt(111) support. Although the authors did not include a CV of MnOx demonstrating its OER activity, they reported an OER current (on a geometric basis) of 5 mA cm$^{-2}$ at ca. $\eta = 560$ (1.78 V). This geometric current density is similar to the geometric current densities of our composite catalysts after the background OER current of Pt(111) substrate is subtracted. From the results of Subbaraman and co-workers, it is not clear whether or not the use of a noble metal support has an impact on the measured OER activity of MnOx catalyst. Direct comparison between the activity metrics of MnO-GC and Au/MnO-GC catalysts in the present report, however, convincingly demonstrates that the addition of a noble metal to MnO nanoparticles sputtered onto a GC substrate will lead to a significant enhancement in their OER activity.

The influence of adding Au to the OER activity of MnOx could occur either through a modification of the MnO phase of the starting catalyst, thus creating a bulk MnOx phase with intrinsically higher OER activity than the original catalyst, or through the formation of active sites at the interface of MnOx and Au. To probe this further, we characterized the catalytic surface after exposure to OER conditions. In Figure 4, we present ex situ TEY Mn L-edge XAS spectra of MnO-GC, MnO/Au-GC, and Au/MnO-GC catalysts previously exposed to an OER relevant potential of 1.65 V. The corresponding chronoamperometry curves used to generate the OER catalytic surfaces are shown in Supporting Information, Figure S7. As expected, exposure to oxidative potentials leads to oxidation of MnO nanoparticles in all three cases. The presence of Au, however, favors formation of less oxidized MnOx nanoparticles. Furthermore, in the Au/MnO$_2$-GC catalyst where Au is deposited on top of the MnO nanoparticles, Mn assumes the lowest oxidation state among the three OER samples (Supporting Information, Figure S8). One potential explanation for this result is that after exposure to OER conditions, the Mn oxidation state may vary as a function of a distance from Au, with MnO$_2$ located away from Au.

To further study the possible interface effect in the composite samples, we compared Mn L-edge XAS spectra of

![Graphical representation of Mn L-edge XAS spectra](Image 328x308 to 561x497)

Figure 4. Total electron yield Mn L-edge XAS of the three catalytic samples, MnO-GC, MnO/Au-GC, and Au/MnO-GC, and two powder standards, MnO$_2$ and Mn$_2$O$_3$. 

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MnO/Au-GC and MnO-GC samples in TEY and auger electron yield (AEY) detection modes. Although both AEY and TEY detection modes are surface sensitive, the probing depth of AEY is shallower than that of TEY. If an interface effect is present, one would expect the measured oxidation state of the oxidized MnO nanoparticles in the MnO/Au-GC sample to vary as a function of probe depth as illustrated in the schematic in Supporting Information, Figure S9, resulting in different TEY and AEY spectra. Figure S9 also presents the AEY and the TEY spectra of MnO-GC and MnO/Au-GC for the as-deposited and for the OER samples. Comparison of AEY and TEY spectra of MnO, in the absence of Au (the MnO-GC sample) in Figure S9 demonstrates a lack of variation in Mn oxidation state as a function of distance from the GC support, both before and after applying an OER relevant potential. Making the same comparisons with the MnO/Au-GC sample, however, shows that while the as-deposited sample has a negligible difference in the Mn spectra between AEY and TEY mode, a more significant spectral difference is observed for the sample exposed to OER relevant potentials. In that sample, the top of the nanoparticle, which is located away from the Au-MnO interface, is likely more oxidized, resembling the spectra of the MnO-GC sample exposed to OER conditions. These results support the hypothesis that there is an interface effect between Au and MnO, after MnO is exposed to high oxidative potentials. In conventional heterogeneous catalysis, such an interface effect has been previously observed with MnO2 and other reducible metal oxides, such as TiO2 and Ce2O3, in contact with Au. The ex situ nature of the Mn L-edge XAS characterization, however, does not allow us to differentiate between an interface effect that emerges when the catalyst is under OER conditions or an interface effect that emerges after the catalyst is removed from the electrochemical cell. To answer this question, in situ methods were required.

In situ characterization was performed with a bulk-sensitive technique using Mn K-edge XAS. Our measurements focused on the X-ray absorption near edge structure (XANES) region, which probes the bulk electronic structure of the catalyst. In the experiments, MnO and Au nanoparticles were deposited on ~200 μm thick GC wafers to allow X-ray illumination of the backside of the electrodes. We characterized two samples, MnO-GC and MnO/Au-GC, with a slightly reduced loading of MnO compared to the ex situ samples (Supporting Information, Figure S10). These samples were held at 1.65 V during the Mn K-edge XAS measurement. Despite the MnO/Au-GC sample having a higher OER current throughout the entire experiment, our in situ characterization shown in Figure S5 does not reveal a measurable difference between the XANES spectra of Mn in the MnO-GC and the MnO/Au-GC samples. The observation of the same average Mn oxidation during the electrochemical evolution of oxygen for both samples suggests that a small subset of sites rather than the bulk phase is critical to the enhanced OER catalysis in the samples containing Au nanoparticles; for example, this could occur by means of some of the MnOx sites locally interacting with Au in such a way as to form OER active sites with modified properties during electrochemical characterization.

To further explore this hypothesis we performed electrochemical characterization of the MnO-GC catalyst before and after adding 0.1 mM of H[Au(III)Cl4] to the electrolyte. Figure 6 shows that as we electrodeposit more Au onto the surface by sweeping to more reductive potentials, the OER activity increases, eventually reaching a current density greater than 10 mA cm−2 at 1.8 V. When the same experiment was performed with a bare GC substrate, we also observed an increase in the OER activity with increased electrodепosition of Au, but the activity remained low, below 1 mA cm−2 at 1.8 V. These experiments demonstrate that the addition of Au salt to the characterization electrolyte will lead to a similar enhancement in OER activity of MnO-GC catalyst as evaporation of Au onto the catalytic surface prior to electrochemical characterization. Some dissolution of Au into electrolyte is expected to occur at the highly oxidative potentials necessary for the OER. We took advantage of this fact and also performed experiments with the MnO-GC sample in the setup previously used to characterize samples containing evaporated Au nanoparticles, but without cleaning the cell. Thus, trace Au was expected to be present to form a sample which we refer to as “trace-Au/MnO-GC”. Supporting Information, Figure S11 shows that although the morphology of the MnO nanoparticles in trace-Au/MnO-GC had not changed while evaluating OER activity, the OER current increased with time during a potential hold at 1.65 V for 30 min (after first cycling the catalyst from 0.05 to 1.65 V at 20 mV s−1), ultimately rising to the same level of OER activity as that for the Au/MnO-GC sample. This result further illustrates that the electrolyte conditions can have a significant effect on the activity of MnOx, with the addition of only a trace amount of Au leading to a significant increase in the OER activity.

Supporting Information, Figure S12 compares cyclic voltammograms of the Au-GC, Au/MnO-GC, and trace-Au/MnO-GC samples. Analysis of electrochemical features of the samples confirms the presence of Au in trace-Au/MnO-GC sample and reveals that the presence of MnO2 has an influence on the red-ox properties of Au. Specifically, the addition of MnO2 leads to formation of multiple oxidation/reduction peaks as opposed to the one clear oxidation peak and the one clear reduction peak observed in the Au-GC sample. The presence of multiple oxidation/reduction peaks in both Au/MnO-GC and
trace-Au/MnO-GC could indicate that the interactions between Au and MnO₂ have led to a formation of Au sites with different strengths of Au−OH adsorption. Since reducible metal oxides such as MnO₂ are known to interact with noble metals to form a more oxidized noble metal at the interface of the two materials, this result is consistent with the presence of interfacial effects between MnO₂ and Au.

After taking into account our ex situ and in situ spectroscopic characterization as well as electrochemical studies, we propose that the observed enhancement in the OER activity of MnO₂ in the presence of Au is caused by the local participation of Au in the OER catalysis, for example through dissolution of Au and its re-deposition onto a subset of MnO₂ sites, rather than a bulk change in the starting properties of MnO₂. This interpretation is consistent with all of the results in our study and is also supported by the published literature in the area of OER catalysis for bulk MnO₂ materials. As described previously, Table 1 convincingly demonstrates that the samples consisting of both MnO₂ and Au had anomalously high OER activity when compared to other active MnO₂ OER catalysts. The only MnO₂ catalyst that had been reported to have similar OER activity was a manganese oxyhydroxide prepared using a completely different method, an electrodeposition, but also in the presence of a noble metal, Pt(111). Similar to Au, Pt is known to interact with transition metal oxides to form a reduced oxide and an oxidized metal at the interface of the two materials and dissolve at oxidizing potentials relevant to the OER. Therefore, the presence of Pt during the OER could lead to an enhancement in the OER activity of MnO₂ that is similar to the enhancement observed in the presence of Au. Additionally, a recent work from Najafpour and Sedigh reports that several of MnO₂ phases, including Mn₂O₃, α-Mn₃O₄, β-MnO₂, and K-birnessite can convert to the same layered MnO₂ structure under water oxidizing conditions. As discussed by the authors, this observation deemphasizes the nature of the starting MnO₂ phase in the OER catalysis and highlights the importance of the electrolyte conditions during water oxidation. Therefore, the presence of a noble metal in the catalyst sample or the electrolyte could have a greater influence on the formation of the OER active sites than the as-deposited phase of MnO₂.

The effect of electrolyte conditions on the OER activity of electrolytically active catalysts is not limited to MnO₂. Trotochaud and co-workers have recently reported transformation of NiO thin film catalyst into a layered hydroxide/oxyhydroxide OER catalyst during electrochemical characterization. During the transformation, the catalyst scavenged Fe impurities from the solution incorporating them into the final active catalytic structure. It is possible that like MnO₂, NiO could also interact with noble metal additives to form OER active sites with enhanced TOFs. For example, in a different study, Yeo and co-workers have already demonstrated a higher TOF with a catalyst consisting of only 0.14 monolayers of NiO deposited on Au relative to the TOF of a bulk NiO catalyst, indicating that an interaction between NiO and Au exists. Although Trotochaud and co-workers have also attempted to assess the role of Au by characterizing NiO catalysts on Au/Ti and ITO supports, their study involved conformal NiO thin films with limited electrochemical accessibility of the Au support. In contrast, in the study by Yeo and co-workers, the Au oxidation/reduction features were still present in the submonolayer sample, which meant that Au had an opportunity to locally impact OER activity by either interacting with the adjacent NiO or by dissolving into the electrolyte at the high oxidizing potentials necessary for the OER and subsequently integrating into the NiO catalyst. Thus, the electrochemical accessibility of the Au likely plays a role and should be an important consideration when designing catalyst morphologies for future studies.

CONCLUSIONS

In an investigation of catalysts for the oxygen evolution reaction (OER), we have shown that a composite catalyst of MnO nanoparticles in the presence of Au exhibited an enhancement in electrocatalytic activity by 1 order of magnitude over the best...
reported pure MnO₂ catalysts. A difference in the Mn L-edge X-ray absorption spectra of as-deposited catalysts originally indicated that a change in the starting properties of MnO₂ could be responsible for the observed increase in the OER current. However, subsequent electrochemical measurements as well as ex situ and in situ Mn XAS characterization of the OER relevant samples identified more likely possibilities: namely local, interfacial effects between the Au and MnO₂. Even trace amounts of Au were sufficient to activate the catalyst for the OER, further indicating that a local interaction between Au and MnO₂ₐₚ, impacting only a subset of MnO₂ sites, is the likely cause of the observed OER enhancement. As the intrinsic OER activity of other nonprecious metal oxides such as CoO₃ and NiOₓ (Supporting Information, Table S2)⁶,¹⁷,²¹ are even higher than that of MnO₂, investigating the role of Au in the OER activities of a number of nonprecious metal-based transition metal oxide catalysts could lead to even higher-performance materials.

■ ASSOCIATED CONTENT

Supporting Information
Figures and tables as described in the text; details on X-ray absorption spectra normalization; details on the calculations of the OER activity metrics. This material is available free of charge via the Internet at http://pubs.acs.org.

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Notes
The authors declare no competing financial interest.

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