**Research Articles** 

Angewandte International Edition www.angewandte.org

High Entropy Oxides Hot Paper

How to cite: Angew. Chem. Int. Ed. **2022**, 61, e202200086 International Edition: doi.org/10.1002/anie.202200086 German Edition: doi.org/10.1002/ange.202200086

# **Entropy Enhanced Perovskite Oxide Ceramic for Efficient Electrochemical Reduction of Oxygen to Hydrogen Peroxide**

Ziliang Chen<sup>+</sup>, Jie Wu<sup>+</sup>, Zhengran Chen, Hongyuan Yang, Kai Zou, Xiangyong Zhao, Ruihong Liang,\* Xianlin Dong, Prashanth W. Menezes,\* and Zhenhui Kang\*

Abstract: The electrochemical oxygen reduction reaction (ORR) offers a most promising and efficient route to produce hydrogen peroxide (H<sub>2</sub>O<sub>2</sub>), yet the lack of costeffective and high-performance electrocatalysts have restricted its practical application. Herein, an entropyenhancement strategy has been employed to enable the low-cost perovskite oxide to effectively catalyze the electrosynthesis of  $H_2O_2$ . The optimized Pb-(NiWMnNbZrTi)<sub>1/6</sub>O3 ceramic is available on a kilogram-scale and displays commendable ORR activity in alkaline media with high selectivity over 91 % across the wide potential range for H<sub>2</sub>O<sub>2</sub> including an outstanding degradation property for organic dyes through the Fenton process. The exceptional performance of this perovskite oxide is attributed to the entropy stabilization-induced polymorphic transformation assuring the robust structural stability, decreased charge mobility as well as synergistic catalytic effects which we confirm using advanced in situ Raman, transient photovoltage, Rietveld refinement as well as finite elemental analysis.

## Introduction

Hydrogen peroxide  $(H_2O_2)$  is one of the most significant and fundamental industrial chemicals for several frontier fields

[*]	Dr. Z. Chen, <sup>+</sup> J. Wu, <sup>+</sup> Prof. Z. Kang Institute of Eurotional Nano and Soft Materials (ELINSOM)	
	Jiangsu Key Laboratory for Carbon-based Functional Materials and Davies	
	Joint International Research Laboratory of Carbon-Based Functional	
	Soochow University Suzhou 215123 (China)	
	E-mail: zhkang@suda.edu.cn	
	Dr. Z. Chen, <sup>+</sup> H. Yang, Dr. P. W. Menezes Department of Chemistry: Metalorganics and Inorganic Materials Technische Universität Berlin Straße des 17 Juni 135, Sekr. C2, 10623 Berlin (Germany) E-mail: prashanth.menezes@mailbox.tu-berlin.de	[+
	Z. Chen, K. Zou, Prof. R. Liang, Prof. X. Dong Key Laboratory of Inorganic Functional Materials and Devices Shanghai Institute of Ceramics Chinese Academy of Sciences 588 Heshuo Road, Jiading District, Shanghai 201800 (China)	
	E-mail: liangruihong@mail.sic.ac.cn	

including environment, agriculture, and energy.<sup>[1]</sup> Currently, the World Health Organization (WHO) has listed  $H_2O_2$  as the crucial disinfectant against SARS-CoV-2 that causes the COVID-19 pandemic.<sup>[2]</sup> The generally adopted industrial method for large-scale  $H_2O_2$  production is the anthraquinone cycling process.<sup>[3]</sup> However, this process demands straightforward participation of hydrogen and oxygen gases coupled with catalysis using expensive palladium, followed by a tedious separation procedure to limit the substantial amount of organic wastes.<sup>[4]</sup> In this regard, it is crucial to develop an alternative method to synthesize  $H_2O_2$  by decreasing energy consumption, production cost, and waste generation.

In recent years, the electrochemical synthesis of  $H_2O_2$ from the oxygen reduction reaction (ORR) through the 2e<sup>-</sup> transfer process has aroused great interest as it can theoretically overcome all the drawbacks which are parasitic in the traditional anthraquinone method.<sup>[5]</sup> Nevertheless, ORR is a competitive reaction and favors a 4e<sup>-</sup> pathway to form water molecules.<sup>[6]</sup> Therefore, developing an electrocatalyst featuring high activity and selectivity towards a 2e<sup>-</sup> pathway is the prerequisite for the sustainable electrosynthesis of  $H_2O_2$  via ORR.<sup>[7]</sup> In this context, several carbonaceous materials, transition metal-based compounds as well as their hybridization have been investigated as advanced electrocatalysts for the electrosynthesis of  $H_2O_2$ .<sup>[8]</sup> Nevertheless, most of these electrocatalysts can still only be produced at the (milli)gram level, which is likely to restrict

Material Chemistry Group for Thin Film Catalysis—CatLab Helmholtz-Zentrum Berlin für Materialien und Energie Albert-Einstein-Str. 15, 12489 Berlin (Germany) E-mail: prashanth.menezes@helmholtz-berlin.de Prof. X. Zhao Key Laboratory of Optoelectronic Material and Device Department of Physics Shanghai Normal University Shanghai 200234 (China)

◎ © 2022 The Authors. Angewandte Chemie International Edition published by Wiley-VCH GmbH. This is an open access article under the terms of the Creative Commons Attribution License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited.

Dr. P. W. Menezes

<sup>[&</sup>lt;sup>+</sup>] These authors contributed equally to this work.

the large-scale application. Besides, the preparation procedures for such electrocatalysts are usually demanding as they require a rigorous coordination chemistry environment, control of the morphology as well as inhibition of (nano)particles agglomeration.

Moreover, most of the reported electrocatalysts in the catalytic process undergo severe oxidation at high operation voltage, and the overpotential window to attain a sustained high peroxide selectivity is narrow ( $\approx 0.4$  V), thus inevitably leading to poor durability and efficiency.<sup>[5a,9]</sup> Although continuous progress has been achieved over the years to develop catalysts with high efficiency for the electrosynthesis of H<sub>2</sub>O<sub>2</sub>, there are still several formidable challenges to enabling an electrocatalyst that can be readily prepared in large quantities to economically convert O<sub>2</sub> to H<sub>2</sub>O<sub>2</sub> over a broad voltage range whilst concurrently maintaining robust stability and yield.

Perovskite oxides are a category of complex oxides with the chemical formula  $ABO_3$ , in which A atoms have larger cation radii than those of B atoms that are surrounded by the network of corner-sharing  $BO_6$  octahedra.<sup>[10]</sup> The unique crystal structure combined with the advantages of tunable composition and low cost render them highly promising for efficient electrocatalysts.<sup>[11]</sup> Previous studies have demonstrated that perovskite oxides are capable of effectively catalyzing the 4e<sup>-</sup> ORR, showing their potential for fuel cells.<sup>[12]</sup> Cho et al. reported that the catalyst based on  $La_{0.3}(Ba_{0.5}Sr_{0.5})_{0.7}Co_{0.8}Fe_{0.2}O_{3-\delta}$  delivered a four-electron transfer pathway, however, in the absence of La, the electron transfer number of  $Ba_{0.5}Sr_{0.5}Co_{0.8}Fe_{0.2}O_{3-\delta}$  was decreased to 3.2, which meant that the 2e<sup>-</sup> ORR pathway began to compete during ORR reaction.<sup>[13]</sup> From this report, it can be concluded that the 2e<sup>-</sup> pathway for ORR is feasible with perovskite oxide electrocatalysts albeit the selectivity is unsatisfactory. This shows that it is indeed possible to drive a 2e<sup>-</sup> pathway-dominated ORR by tuning the composition and structure of perovskite oxides, yet the related investigation is still lacking. On the other hand, highentropy materials (alloys, oxides, and sulfides) have recently received extensive interest as novel electrocatalysts for a variety of catalytic applications including hydrogen evolution reaction, oxygen evolution reaction, and 4e<sup>-</sup> pathwaydominated ORR.<sup>[14]</sup> These high-entropy materials can be several synthesized by appealing entropy-driven strategies.<sup>[15]</sup> Owing to the intrinsic traits of tunable composition, cocktail effect, as well as high corrosion and oxidation resistance, all of these high-entropy electrocatalysts showed impressive catalytic activity. Therefore, based on the premises, combining perovskite oxide with the concept of high-entropy to form high-entropy perovskite oxides might be an effective approach to realize highefficiency 2e<sup>-</sup> ORR.

Bearing the above points in mind, herein, a high-entropy perovskite oxide ceramic Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> derived from the Pb(ZrTi)<sub>1/2</sub>O<sub>3</sub> prototype was deliberately synthesized *via* the thermal-induced solid solution method for the efficient H<sub>2</sub>O<sub>2</sub> production by ORR. Here, the metal species in the *B* site are expected to be able to drive  $2e^-$  ORR, while the inert Pb in *A* site is anticipated to serve as a

barrier to isolate the surrounding B atom and thereby promotion of 2e<sup>-</sup> ORR.<sup>[16]</sup> Strikingly, by combining in situ Raman, ex situ transient photovoltage measurement (TPV) with Rietveld refinement and finite element analysis (FEA), we disclose that upon the construction of high-entropy, the crystal structure transforms from the initial tetragonal-type  $Pb(ZrTi)_{1/2}O_3$  to the cubic-type  $Pb(NiWMnNbZrTi)_{1/6}O_3$ , showing a decreased internal lattice strain. Meanwhile, the presence of high-entropy decreases the charge transfer ability and optimizes the surface charge density, contributing to the selectivity towards 2e<sup>-</sup> ORR. Moreover, all B-site metal species could serve as the catalytic active center and synergistically perform catalysis. Benefiting from these merits, the as-synthesized Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> exhibited a selectivity as high as 96 % at the potential of 0.4 V vs. RHE. In addition, it also showed negligible performance decay even after 12 h of duration at 0.4 V vs. RHE, together with an outstanding dye degradation property for methylene blue (MB), methyl orange (MO), and rhodamine B (RB). The performances attained here are not only significantly better than those of Pb(ZrTi)<sub>1/2</sub>O<sub>3</sub> but also one of the best among previously reported electrocatalysts. Most importantly, the synthetic strategy adopted here for the preparation of catalysts can easily be scaled-up (to the kilogramlevel) and has huge potential for future practical applications.

Angewandte

Chemie

#### **Results and Discussion**

#### Synthesis, Phase, and Microstructural Features

Figure 1a schematically illustrates the synthesis procedure for the high-entropy perovskite oxide. First, target metal oxides and carbonate powders were mixed homogeneously by high-energy ball-milling. Then, the mechanical-treated powders were pressed as pellets for high-temperature sintering under the air to obtain the Pb(NiWMnNbZrTi)<sub>1/</sub> <sub>6</sub>O<sub>3</sub> ingots. Finally, the as-prepared ingots were crushed into powders for further use. Remarkably, as visually presented in Figure 1b, the catalyst could be prepared up to a kilogram-scale only by a one-batch operation. The inductively coupled plasma mass spectrometry (ICP-MS) indicated that the atomic ratio of Pb : Ni:W:Mn : Nb:Zr:Ti was 1:0.164:0.165:0.160:0.163:0.174:0.174 (Table S1), which was quite close to the designed composition, demonthe successful preparation of the strating Pb-(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> by thermal-solid diffusion. Rietveld refinement of the X-ray powder diffraction pattern (XRD) showed a pure perovskite oxide phase crystallizing in cubic structure ( $Pm\bar{3}m$ ) with refined lattice parameters a=b=c=4.0245(4) Å (Figure 1c, Table S2, and S3). For comparison, the pure Pb(ZrTi)<sub>1/2</sub>O<sub>3</sub> was also synthesized under the same condition (Figure 1d, Table S1). Interestingly, the  $Pb(ZrTi)_{1/2}$  $_{2}O_{3}$  phase was crystallized in the tetragonal structure with lower symmetry (P4mm) (Table S2 and S4). Moreover, based on the Williamson-Hall method, the lattice strain for  $Pb(NiWMnNbZrTi)_{1/6}O_3$  (+1.05(5)%) was found to be much smaller as compared to that (+1.60(4)%) of Pb-

## **Research Articles**



**Figure 1.** a) Schematic synthesis of high-entropy ceramic Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> compound. b) Digital image of the kilogram-level production of Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub>. Rietveld refinement of XRD patterns for c) Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> and d) Pb(ZrTi)<sub>1/2</sub>O<sub>3</sub>. e) Comparison of lattice strain of Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> and d) Pb(ZrTi)<sub>1/2</sub>O<sub>3</sub>. e) Comparison of lattice strain of Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> and d) Pb(ZrTi)<sub>1/2</sub>O<sub>3</sub>. Green line, red " + " symbol, blue line, and green vertical bars showed in Figures (c) and (d) represent the observed data, fitted data, differentiation, and the positions of the reflection peaks of the phase in the XRD patterns. High-resolution XPS spectra of f) Ni 2p, g) W 4f, h) Mn 2p, i) Nb 3d, j) Ti 2p, and k) Zr 3d in Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub>.

 $(ZrTi)_{1/2}O_3$  (Figure 1d). Such results clearly indicated that an entropy enhancement could induce the release of lattice strain and led to the polymorphic phase transformation of  $Pb(ZrTi)_{1/2}O_3$ , which was expected to improve the structural stability. Figure 1f-k reveals the high-resolution X-ray photoelectron spectroscopy (XPS) spectra of Ni 2p, W 2p, Mn 2p, Nb 2p, Zr 2p, and Ti 2p, respectively, where their respective oxidation states have been shown. Specifically, it could be seen from Figure 1f that Ni  $2p_{1/2}$  and Ni  $2p_{3/2}$  were distributed at 873.7 and 855.3 eV, respectively, both of which could be deconvoluted into Ni<sup>3+</sup> and Ni<sup>2+</sup>, suggesting the presence of Ni–O bond.<sup>[17]</sup> The peaks at 37.4 and 35.4 eV in Figure 1g could be assigned to W  $4f_{5/2}$  and W  $4f_{7/2}$ , respectively, which were widely identified as  $W^{6+}$  in  $WO_3$ .<sup>[18]</sup> The spectrum displayed in Figure 1h could be divided into two peaks at 645.4 and 642.2 eV, which were attributed to the Mn<sup>4+</sup> and Mn<sup>3+</sup>, respectively.<sup>[19]</sup> As presented in Figure 1i, two characteristic peaks at 209.3 and 206.6 eV could be ascribed to Nb  $3d_{3/2}$  and Nb  $3d_{5/2}$  in the niobium oxide.<sup>[20]</sup> Moreover, the Ti–O,<sup>[21]</sup> Zr–O,<sup>[22]</sup> and Pb–O<sup>[23]</sup> bonds could also be observed in the high-resolution Ti 2p (Figure 1j), Zr 3d (Figure 1k), and Pb 4f (Figure S1a) XPS spectra, respectively. The high-resolution O 1s XPS spectra further confirmed the presence of metal-O bonds (Figure S1b).<sup>[24]</sup> For comparison, the XPS spectra for all elements in Pb(ZrTi)<sub>1/2</sub>O<sub>3</sub> are shown in Figure S2. The comparative results demonstrated that the entropy enhancement could effectively modify the electronic structure of the compound (For details refer to Figure S2).

To further acquire the microstructure information, the field emission scanning electron microscopic (FESEM) image for Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> was conducted (Figure 2a) that exhibited uniform dispersion of pseudosphere particles with sizes ranging from nanoscale to microscale.

**Research Articles** 



*Figure 2.* a) FESEM image, b) TEM image; c) corresponding SAED pattern, d) high-magnified TEM image, e, f) high-resolution TEM images, g) HAADF image of representative Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> particle and the corresponding elemental mapping of h) Ni, i) W, j) Mn, k) Ti, l) Zr, m) Nb, n) Pb, and o) O.

Subsequently, the transmission electron microscopic (TEM) image of the representative Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> particle was recorded (Figure 2b). The selected area electron diffraction (SAED) pattern of the chosen region of the particle (Figure 2b) showed diffraction rings/dots that can be precisely indexed to (110), (111), (200), (210), (211) and (220) facets of Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> phase (Figure 2c), respectively. Moreover, the result obtained from SAED is in good agreement with the XRD pattern. The magnified TEM image displayed in Figure 2d shows a number of nanocrystallites (as marked by the dotted circle). High-resolution TEM (HRTEM) images for the region A and B of Figure 2d are presented in Figure 2e and 2f, respectively, in which (110) and (202) facets with an intersection angle of about  $60^{\circ}$  and (110) and (-210) facets with an intersection angle of about 108° belonging to Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> phase were confirmed. In addition, the high angle annular darkfield scanning TEM (HAADF-STEM) pattern exhibited the bright region indicating the homogeneous distribution of metal species. The corresponding EDS mapping further substantiated the high compositional uniformity of the Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> phase within the entire particle (Figure 2g–o). Besides, the phase purity, structure, and composition of Pb(ZrTi)<sub>1/2</sub>O<sub>3</sub> were also verified by the HRTEM and EDS mapping results (Figure S3).

#### Effect of Entropy on ORR Activity

Motivated by the unique structure and composition, the electrochemical ORR activity of Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> to produce  $H_2O_2$  was systematically examined by employing the rotating ring disk electrode (RRDE) technique in 0.1 M KOH, together with PbZrO<sub>3</sub>, PbTiO<sub>3</sub>, Pb(ZrTi)<sub>1/2</sub>O<sub>3</sub>, Pb-(MnNbZrTi)<sub>1/4</sub>O<sub>3</sub>, Pb(WNbZrTi)<sub>1/4</sub>O<sub>3</sub>, and Pb(NiNbZrTi)<sub>1/4</sub>O<sub>3</sub> for comparison (Figure S4–S11). The collection efficiency of the RRDE was determined as 0.37 (Figure S12). Figure 3a provided the linear sweep voltammetry (LSV) curves of the electrocatalysts at the scan rate of 10 mV s<sup>-1</sup>, in which both the disk and ring current density are presented.



*Figure 3.* a) LSV curves of Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub>, Pb(ZrTi)<sub>1/2</sub>O<sub>3</sub>, Pb(MnNbZrTi)<sub>1/4</sub>O<sub>3</sub>, Pb(WNbZrTi)<sub>1/4</sub>O<sub>3</sub>, and Pb(NiNbZrTi)<sub>1/4</sub>O<sub>3</sub>, recorded at 1600 rpm and 10 mV s<sup>-1</sup> (bottom part), together with the corresponding H<sub>2</sub>O<sub>2</sub> current on the ring electrode (upper part). b) Calculated electron transfer number and c) selectivity of H<sub>2</sub>O<sub>2</sub> within the potential sweep. d) Comparison of stability of Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> and Pb(ZrTi)<sub>1/2</sub>O<sub>3</sub> at a fixed disk potential of 0.1 V. e) Comparison of ORR activity of Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> with previously reported electrocatalysts; High-resolution XPS spectra of f) Ni 2p, g) W 4f, h) Mn 2p, i) Ti 2p, j) Zr 3d, and k) Nb 3d in Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> after the chronoamperometry test.

Notably, the onset potentials at the disk and ring electrodes around  $\approx 0.76 \text{ V}$  vs. RHE were both for Pb-(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub>, and the potential to deliver a high current density of 1.0 mA cm<sup>-2</sup> was 0.66 V vs. RHE, showing the outstanding ORR catalytic activity. Moreover, according to the disk and ring current density, the average electron transfer number (n) towards ORR for all the electrocatalysts was calculated and compared in Figure 3b and Figure S13. As expected, the *n* for Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> was calculated to be  $\approx 2.08$  in the potential range of 0.3–0.7 V vs. RHE, which was the lowest value among the tested electrocatalysts, substantiating a 2e<sup>-</sup>-dominated ORR pathway. Notably, the selectivity of H2O2 production for Pb-(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> was estimated above 91 % within the potential range from 0.1 to 0.7 V vs. RHE (Figure 3c). To strengthen our claims, we also synthesized other highentropy perovskites with quinary and novenary B-site metals based on the  $Pb(ZrTi)_{1/2}O_3$  prototype, such as Pb-(NiWMnNbTi)<sub>1/5</sub>O<sub>3</sub> (Figure S4, Figure S14), Pb-Pb- $(NiWMnNbZr)_{1/5}O_3$ (Figure S4, Figure S15) and

(NiWMnNbTiZrFeVMo)<sub>1/9</sub>O<sub>3</sub> (Figure S4, Figure S16). By comparing to the  $2e^-$  ORR activity of Pb(NiWMnNbZrTi)<sub>1/</sub>  $_6O_3$ , it could be further confirmed that the  $2e^-$  ORR activity was gradually increased with the enhancement of entropy in our studied system within a certain entropy range (Figure S17). Moreover, the partial incorporation of La into Asite metal of Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> was also carried out and the achieved Pb<sub>0.9</sub>La<sub>0.1</sub>(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> compound still exhibited the comparable  $2e^-$  ORR activity to Pb-(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> (Figure S18 and S19).

On the other hand, since the high durability is the prerequisite for the practical production of  $H_2O_2$  by ORR, the long-time stability measurement of Pb-(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> was conducted *via* chronoamperometry at a potential of 0.1 V (Figure 3d). Remarkably, the activity remained unchanged for 12 h, demonstrating the robust electrochemical production of  $H_2O_2$ . However, compared to Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> could be obtained after 12 h, implying the advantage of high entropy in improving

Angew. Chem. Int. Ed. 2022, 61, e202200086 (5 of 10)

the structural stability. Such exceptional ORR performance of Pb(NiWMnNbZrTi)1/6O3, in terms of selectivity and durability, surpasses most of the recently reported electrocatalysts thus ranking it among one of the best nonprecious-metal electrocatalysts (Figure 3e, Table S5). In order to reveal the advantages of Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> in the production of  $H_2O_2$ , we evaluated the decomposition degree of H<sub>2</sub>O<sub>2</sub> on different catalysts (Figure S20). A certain amount of electrocatalysts were added into 0.1 M KOH containing 10 mM H<sub>2</sub>O<sub>2</sub>, and the Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> possessed the lowest H2O2 decomposition among all the measured electrocatalysts. To further verify the highly selective  $H_2O_2$  production, the determination of  $H_2O_2$  was performed by the cerium sulfate (Ce<sup>4+</sup>) method. Upon gradually undergoing the process of Ce<sup>4+</sup> titration, the color of the solution turned orange-yellow. The theoretical concentration of  $H_2O_2$  produced at 34 mA cm<sup>-2</sup> for 1 h was calculated to be 864 mg L<sup>-1</sup> whereas, in the real measurement, the  $H_2O_2$  concentration achieved by the Ce<sup>4+</sup> method was about  $807 \text{ mg L}^{-1}$ , which was quite close to the theoretical value. The above results signified the high selectivity of Pb(NiWMnNbZrTi)<sub>1/6</sub>O3 towards H2O2 by ORR.

#### Insights into the Entropy-Improved ORR Activity

To acquire deeper insights into the impressive activity, selectivity, and durability, XPS was performed, and the composition of the Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> after the above chronoamperometry test was examined. As seen in Figure 3f-k and Figure S21, the high-resolution XPS spectra suggested that all chemical elements were preserved after the durability test. Moreover, after the chronoamperometry test, the binding energy for B-site metal species of Ni 2p, Mn 2p, Ti 2p, Zr 3d, and Nb 3d was shifted to the higher energy by around 0.3 eV, while W 4f shifted to lower energy by 0.3 eV. This difference could arise from the surface change during catalysis, suggesting that B-sites were the main active sites for ORR.<sup>[24]</sup> Note that the binding energy of high-resolution XPS spectra for A-site Pb was basically unchanged before and chronoamperometry test, implying its weak contribution to catalytic activity. These results were similar to those of Pb(ZrTi)<sub>1/2</sub>O<sub>3</sub> after the chronoamperometry test (Figure S22). To further evaluate whether and how  $Pb(NiWMnNbZrTi)_{1/6}O_3$  phase changed during the ORR process, the HRTEM, SAED, HAADF as well as the corresponding EDS mapping was recorded. As displayed in Figure S23, no new phase was identified after the chronoamperometry test and the attained results matched quite well with the initial Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub>. However, some defects such as lattice distortion were more apparent on the surface of the particles, which could be ascribed to the catalysis-induced surface reconstruction (Figure S23). Similar to the Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub>, the HRTEM, SAED as well as HAADF of  $Pb(ZrTi)_{1/2}O_3$  after chronoamperometry test also indicated the implicit surface reconstruction (Figure S24). To further compare the compositional stability, the element content in the 0.1 M KOH electrolyte after the chronoamperometry test was measured with ICP. The results obtained in Table S6 clearly revealed a higher element dissolution for  $Pb(ZrTi)_{1/2}O_3$  than that of Pb-(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub>.

To gain an understanding of the active structure of Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub>, in situ Raman spectra were performed in 0.1 M KOH at various ORR potentials (-0.2--0.6 V vs. SCE). As seen in Figure 4a, the *B*-site metal-O bond for all metal species in Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> gradually increased with the increase of the applied operation potential. This indicated that all B-site metal species possibly synergistically catalyzed the  $O_2$  to produce  $H_2O_2$ . This could be due to the synergistic catalysis effect of high entropy materials although the related mechanism can be complex. Such observations have also been made for many high entropy materials in typical catalytic reactions including OER and 4e<sup>-</sup>-dominated ORR.<sup>[25]</sup> It is important to note that, a change of Pb-O signal was also found in the in situ Raman spectra, which could be ascribed to the strong intrinsic adsorption ability of Pb to hydroxyl groups.<sup>[26]</sup> Interestingly, it is worth mentioning that  $Pb(ZrTi)_{1/2}O_3$  was first thought of as 4e<sup>-</sup> ORR electrocatalyst, however, our present investigation reveals that it can also show an intrinsic catalytic activity towards the production of H2O2. This could be verified in Figure 4b where the signals for Zr-O and Ti-O bonds increased gradually with increasing the operating voltage.

On the other hand, the TPV was applied to investigate and compare the electron transfer behavior of the Pb-(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> and Pb(ZrTi)<sub>1/2</sub>O<sub>3</sub> catalysts. TPV curves of Pb(NiWMnNbZrTi)<sub>1/6</sub>O3 and Pb(ZrTi)<sub>1/2</sub>O3 are displayed in Figure 4c. Especially, the TPV curve for Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> was more oscillating as compared to that for Pb(ZrTi)<sub>1/2</sub>O<sub>3</sub>, implying the distinct electron transfer process between these two oxides. Moreover, a slower TPV signals decay tendency was observed for Pb(ZrTiMnWNiNb)<sub>1/6</sub>O<sub>3</sub> (Figure S25) indicating the electron accumulation effect and sluggish kinetics of electron transfer.<sup>[27]</sup> Thus, the formation of high entropy perovskite oxide weakens the charge transfer ability, which may block the binding of oxygen with electrons. As the  $4e^-$  ORR process requires a fast electron transfer ability and sufficient reaction of oxygen with electrons, the diminished charge depletion ability and charge transfer kinetics caused by the high entropy possibly suppressed the 4e<sup>-</sup> ORR process and facilitated the 2e<sup>-</sup> ORR process.<sup>[9,28]</sup> To further analyze the TPV data, continuous wavelet transformation (CWT) was performed on TPV data. The three-dimensional continuous wavelet transformation results of Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> and Pb(ZrTi)<sub>1/2</sub>O<sub>3</sub> were displayed in Figures 4d and e, in which three parameters including time, frequency, and intensity were involved. Low-frequency denotes slow electron transportation, and high frequency represents fast electron transportation. Taking the low-frequency 5 Hz as an example, the Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> exhibits a lagging peak than that of Pb(ZrTi)<sub>1/2</sub>O<sub>3</sub> manifesting the accumulation of electrons on the Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> surface (reaction sites) rather than the transfer (inset in Figure 4c).

**Research Articles** 



**Figure 4.** In situ Raman spectra of a) Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> and b) Pb(ZrTi)<sub>1/2</sub>O<sub>3</sub> in O<sub>2</sub>-saturated 0.1 M KOH from -0.2 to -0.6 V  $\nu$ s. SCE with an interval voltage of 0.05 V. c) Comparison of TPV curves between Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> and Pb(ZrTi)<sub>1/2</sub>O<sub>3</sub> The inset in (c) shows the comparison of CWT curves between Pb(NiWMnNbZrTi)<sub>1/2</sub>O<sub>3</sub>. 3D CWT patterns of d) Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> and e) Pb(ZrTi)<sub>1/2</sub>O<sub>3</sub>. f) Comparison of EMD curves between Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> and Pb(ZrTi)<sub>1/2</sub>O<sub>3</sub>. g) Stress distribution diagram of Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> system under the fluid field (the left and right ruler represents the stress distribution of fluid and particle, respectively). Surface charge map of the h) Pb(NiWMnNbZrTi)<sub>1/2</sub>O<sub>3</sub> and i) Pb(ZrTi)<sub>1/2</sub>O<sub>3</sub> particle at steady state.

Similarly, a comparison at higher frequencies also supported the same conclusion (Figure S26).

In order to better understand the electron transfer process on the catalyst surface, empirical mode decomposition (EMD) was also utilized to refine the meaningful TPV data. EMD is an adaptive time-space analysis method and performs operations that partition a series into intrinsic mode functions (IMFs) without leaving the time domain. These modes contribute to various signals contained within the data. By the EMD method, the white noise is averaged out, and then the persistent part that survived from the averaging process provides real and meaningful signals (Figure 4f).<sup>[29]</sup> It was found that the Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub>

delivered a higher intensity distribution than  $Pb(ZrTi)_{1/2}O_3$ , i.e, electrons were easier to accumulate and be trapped on the surface of the  $Pb(NiWMnNbZrTi)_{1/6}O_3$ , which was consistent with the TPV and CWT results. Therefore, based on the above results, it was clear that the electron transfer on the Pb(NiWMnNbZrTi)\_{1/6}O\_3 surface was slower, which probably affected the binding reaction of active sites with sufficient oxygen leading to the preferrable  $2e^-$  pathway. In the case of Pb(ZrTi)\_{1/2}O\_3, the electrons stay on the catalytic sites for a shorter time causing more electrons to react with oxygen and allowing the whole reaction pathway to deviate from the  $2e^-$  pathway.

© 2022 The Authors. Angewandte Chemie International Edition published by Wiley-VCH GmbH

Furthermore, the stress, electrical, and charge distribution on the surface of Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> and Pb- $(ZrTi)_{1/2}O_3$  particle was modeled by the FEA (Figure 4g and Figure S27, S28). The results showed that both the charge (8.15×10<sup>-8</sup> C) and charge density (2.03×10<sup>-2</sup> Cm<sup>-2</sup>) for Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> were lower than those for Pb- $(ZrTi)_{1/2}O_3$  (1.54×10<sup>-7</sup> C, 3.69×10<sup>-2</sup> Cm<sup>-2</sup>) (Figure 4h,i). The less positive charge density on the surface of Pb-(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> not only suggested the possible trap of more electrons on the surface as compared to Pb(ZrTi)<sub>1/2</sub>O<sub>3</sub> towards catalysis but also implied its weaker binding ability with –OOH, and thus being able to easily dissociate the bond between the substrate and –OOH to produce the H<sub>2</sub>O<sub>2</sub>.

#### **Extended Dye Degradation Property**

Inspired by the above exciting findings, a tandem reaction was conducted by coupling ORR with dye degradation. As illustrated in Figure 5a, Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> was loaded on the carbon paper, and then assembled in an H-cell electrolyzer containing O<sub>2</sub>-saturated 0.1 M KOH solution. Figure 5b shows the LSV curves obtained under the Ar and O<sub>2</sub>. It could be seen that the attained current density was negligible in the Ar-saturated 0.1 M KOH, while a large current density was observed in the O<sub>2</sub>-saturated 0.1 M KOH, which demonstrated a strong oxygen reduction ability of Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub>. Subsequently, the electrolytic cell was discharged at 34 mA cm<sup>-2</sup> for 30 minutes, and then an appropriate amount of electrolyte in the cathode area was taken out, acidified, and then reacted with Ce<sup>4+</sup> to determine H<sub>2</sub>O<sub>2</sub> productivity. Remarkably, in addition to the excellent activity and selectivity, the Pb-(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> led to a H<sub>2</sub>O<sub>2</sub> production rate as high as 1.1 mol g<sub>cat</sub><sup>-1</sup> h<sup>-1</sup>. Besides, the long-term stability test result delivered only a very minor current decrease ( $\approx 2\%$ ) even after 12 h (Figure 5c).

It is well known that the Fenton reaction between  $H_2O_2$ and Fe<sup>2+</sup> will generate hydroxyl radicals, and therefore, the Fenton reaction is often used in treating pollutants and wastewater. Inspired by this, we further employed Pb-(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> as an electrocatalyst to generate H<sub>2</sub>O<sub>2</sub> for the degradation of organic dyes through the Fenton reaction (details in the Experimental Section). Three organic dyes, MB, MO, and RB, were used to evaluate the degradation ability. Figure 5d-f shows the optical image of MB, MO, and RB where fading of color to decolorization was observed within the first 15 mins. Figure 5g-i displays the UV/Visible absorption spectra of MB, MO, and RB. At a fixed dye concentration (30 ppm), all three dyes exhibited a large decrease in their absorption peak within the first two minutes, and no obvious absorption was visible at 15 min. Moreover, the rate of dye degradation for the investigated dyes was much faster than the previous literature reported ORR catalysts (Table S7). The presented results demon-



*Figure 5.* a) Digital image of H-type electrolyzer assembled by working electrode (Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> loaded onto carbon paper), counter electrode (IrO<sub>2</sub> loaded onto carbon paper), and a SCE reference electrode in 0.1 M KOH. Inset in (a) shows the schematic diagram of H-type electrolyzer. b) LSV curves of the catalyst in O<sub>2</sub> and Ar-saturated 0.1 M KOH. c) *i–t* curve of the H<sub>2</sub>O<sub>2</sub> production in the H-type electrolyzer. Digital images for the decolorization of d) MB, e) MO, and f) RB against the degradation time; UV/Visible spectra of the g) MB h) MO and i) RB solution at different time intervals.

Angew. Chem. Int. Ed. 2022, 61, e202200086 (8 of 10)

© 2022 The Authors. Angewandte Chemie International Edition published by Wiley-VCH GmbH

strate that Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> is not only an excellent catalyst for the large-scale production of  $H_2O_2$  but also can spontaneously be applied to effectively degrade a variety of pollutants in wastewater by the Fenton reaction.

### Conclusion

In summary, a novel high-entropy perovskite oxide ceramic (Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub>) was synthesized on a large scale by a thermal-triggered solid solution method and was demonstrated as an efficient electrocatalyst for the electrochemical reduction of O2 to H2O2. The as-synthesized Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> was found to be a selective catalyst for the electrosynthesis of H<sub>2</sub>O<sub>2</sub> in alkaline solution producing over 91% of selectivity across a wide potential range from 0.1 to 0.7 V and was remarkably stable without any loss of current even after 12 h duration at 0.4 V. In addition, the entropy-enhanced Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> was directly compared with the low-entropy Pb(ZrTi)<sub>1/2</sub>O<sub>3</sub> that showed a significant difference in the performance of ORR. Further in situ and ex situ investigations combined with computational analysis unveiled that the entropy increase not only induced a polymorphic phase transition accompanied with a release of lattice strain but also reduced electron migration capability on the surface, thus improving structural stability as well as enhancing the catalytic activity for the selective production of H<sub>2</sub>O<sub>2</sub>. Furthermore, the robust H<sub>2</sub>O<sub>2</sub> production enabled by  $Pb(NiWMnNbZrTi)_{1/6}O_3$  electrode ensured the effective electro-Fenton process and efficient degradation of several organic pollutants including MB, MO, and RB, showing enormous potential for this electrode for onsite water treatment applications. Finally, the integration of perovskite with high entropy as well as the related findings for the application of ORR to produce H<sub>2</sub>O<sub>2</sub> not only establishes the representative Pb(NiWMnNbZrTi)<sub>1/6</sub>O<sub>3</sub> as the new benchmark 2e<sup>-</sup> ORR electrocatalysts in alkaline solution but also provides new mechanistic insights and opens a new design avenue for cost-effective, stable and efficient earth-abundant transition metal compound electrocatalysts.

#### Acknowledgements

This work was supported by the National Key R&D Program of China (2020YFA0406104, 2018YFC0308603), National MCF Energy R&D Program of China (2018YFE0306105), the Natural Science Foundation of Jiangsu Province (BK20210735), and the *CIC* and the *111* Project. The authors gratefully acknowledge the funding from Alexander von Humboldt (AvH) Foundation and Deutsche Forschungsgemeinschaft (DFG, German Research Foundation) under Germany's Excellence Strategy-EXC 2008/1–390540038-UniSysCat. H. Yang also thanks China Scholarship Council (CSC) for the Ph.D. fellowship. P.W.M. greatly acknowledges support from the German Federal Ministry of Education and Research in the framework of the project Catlab (03EW0015A/B). Open Access funding enabled and organized by Projekt DEAL.

## **Conflict of Interest**

The authors declare no conflict of interest.

## **Data Availability Statement**

The data that support the findings of this study are available from the corresponding author upon reasonable request.

**Keywords:** Electrocatalysis • High Entropy • Hydrogen Peroxide • Oxygen Reduction Reaction • Perovskite Oxide Ceramic

- a) E. Jung, H. Shin, B. H. Lee, V. Efremov, S. Lee, H. S. Lee, J. Kim, W. Hooch Antink, S. Park, K. S. Lee, S. P. Cho, J. S. Yoo, Y. E. Sung, T. Hyeon, *Nat. Mater.* **2020**, *19*, 436–442; b) C. Xia, S. Back, S. Ringe, K. Jiang, F. Chen, X. Sun, S. Siahrostami, K. Chan, H. Wang, *Nat. Catal.* **2020**, *3*, 125–134.
- [2] Z. Tang, N. Kong, X. Zhang, Y. Liu, P. Hu, S. Mou, P. Liljestrom, J. Shi, W. Tan, J. S. Kim, Y. Cao, R. Langer, K. W. Leong, O. C. Farokhzad, W. Tao, *Nat. Rev. Mater.* **2020**, *5*, 847–860.
- [3] C. D. Gheewala, B. E. Collins, T. H. Lambert, Science 2016, 351, 961–965.
- [4] a) Y. J. Lin, I. Khan, S. Saha, C. C. Wu, S. R. Barman, F. C. Kao, Z. H. Lin, *Nat. Commun.* **2021**, *12*, 180; b) G. F. Han, F. Li, W. Zou, M. Karamad, J. P. Jeon, S. W. Kim, S. J. Kim, Y. Bu, Z. Fu, Y. Lu, S. Siahrostami, J. B. Baek, *Nat. Commun.* **2020**, *11*, 2209.
- [5] a) S. Siahrostami, S. J. Villegas, A. H. Bagherzadeh Mostaghimi, S. Back, A. B. Farimani, H. Wang, K. A. Persson, J. Montoya, *ACS Catal.* **2020**, *10*, 7495–7511; b) Q. Yang, W. Xu, S. Gong, G. Zheng, Z. Tian, Y. Wen, L. Peng, L. Zhang, Z. Lu, L. Chen, *Nat. Commun.* **2020**, *11*, 5478; c) B. Q. Li, C. X. Zhao, J. N. Liu, Q. Zhang, *Adv. Mater.* **2019**, *31*, 1808173.
- [6] C. Tang, L. Chen, H. Li, L. Li, Y. Jiao, Y. Zheng, H. Xu, K. Davey, S. Z. Qiao, J. Am. Chem. Soc. 2021, 143, 7819–7827.
- [7] E. Jung, H. Shin, W. Hooch Antink, Y.-E. Sung, T. Hyeon, ACS Energy Lett. 2020, 5, 1881–1892.
- [8] a) D. A. Kuznetsov, Z. Chen, P. M. Abdala, O. V. Safonova, A. Fedorov, C. R. Muller, J. Am. Chem. Soc. 2021, 143, 5771– 5778; b) Z. Pan, K. Wang, Y. Wang, P. Tsiakaras, S. Song, Appl. Catal. B 2018, 237, 392–400; c) N. Thiyagarajan, D. Janmanchi, Y. F. Tsai, W. H. Wanna, R. Ramu, S. I. Chan, J. M. Zen, S. S. Yu, Angew. Chem. Int. Ed. 2018, 57, 3612– 3616; Angew. Chem. 2018, 130, 3674–3678; d) S. Thundiyil, S. Kurungot, R. N. Devi, ACS Appl. Mater. Interfaces 2021, 13, 382–390.
- [9] J. Gao, B. Liu, ACS Mater. Lett. 2020, 2, 1008–1024.
- [10] G. Chen, Y. Zhu, H. M. Chen, Z. Hu, S. F. Hung, N. Ma, J. Dai, H. J. Lin, C. T. Chen, W. Zhou, Z. Shao, *Adv. Mater.* 2019, *31*, 1900883.
- [11] a) C. Baeumer, J. Li, Q. Lu, A. Y. Liang, L. Jin, H. P. Martins, T. Duchon, M. Gloss, S. M. Gericke, M. A. Wohlgemuth, M. Giesen, E. E. Penn, R. Dittmann, F. Gunkel, R. Waser, M. Bajdich, S. Nemsak, J. T. Mefford, W. C. Chueh, *Nat. Mater.* 2021, 20, 674–682; b) C. Sun, J. A. Alonso, J. Bian, *Adv. Energy Mater.* 2021, 11, 2000459; c) J. Dai, Y. Zhu, H. A. Tahini, Q.

Angew. Chem. Int. Ed. 2022, 61, e202200086 (9 of 10)

<sup>© 2022</sup> The Authors. Angewandte Chemie International Edition published by Wiley-VCH GmbH

Lin, Y. Chen, D. Guan, C. Zhou, Z. Hu, H. J. Lin, T. S. Chan, C. T. Chen, S. C. Smith, H. Wang, W. Zhou, Z. Shao, *Nat. Commun.* **2020**, *11*, 5657; d) S. Chen, Y. Su, P. Deng, R. Qi, J. Zhu, J. Chen, Z. Wang, L. Zhou, X. Guo, B. Y. Xia, *ACS Catal.* **2020**, *10*, 4640–4646.

- [12] a) S. Samira, J. C. A. Camayang, K. Patel, X.-K. Gu, E. Nikolla, ACS Energy Lett. 2021, 6, 1065–1072; b) Q. Ji, L. Bi, J. Zhang, H. Cao, X. S. Zhao, Energy Environ. Sci. 2020, 13, 1408–1428; c) A. Safakas, G. Bampos, S. Bebelis, Appl. Catal. B 2019, 244, 225–232; d) Y. Zhou, Q. Gu, Y. Li, L. Tao, H. Tan, K. Yin, J. Zhou, S. Guo, Nano Lett. 2021, 21, 4861–4867; e) Y. Q. Lyu, F. Ciucci, ACS Appl. Mater. Interfaces 2017, 9, 35829–35836; f) T. D. Thanh, N. D. Chuong, J. Balamurugan, H. Van Hien, N. H. Kim, J. H. Lee, Small 2017, 13, 1701884.
- [13] J. I. Jung, H. Y. Jeong, J. S. Lee, M. G. Kim, J. Cho, Angew. Chem. Int. Ed. 2014, 53, 4582–4586; Angew. Chem. 2014, 126, 4670–4674.
- [14] a) T. A. A. Batchelor, J. K. Pedersen, S. H. Winther, I. E. Castelli, K. W. Jacobsen, J. Rossmeisl, *Joule* 2019, *3*, 834–845; b) T. Li, Y. Yao, B. H. Ko, Z. Huang, Q. Dong, J. Gao, W. Chen, J. Li, S. Li, X. Wang, R. Shahbazian-Yassar, F. Jiao, L. Hu, *Adv. Funct. Mater.* 2021, *31*, 2010561; c) N. Kumar Katiyar, K. Biswas, J.-W. Yeh, S. Sharma, C. Sekhar Tiwary, *Nano Energy* 2021, *88*, 106261.
- [15] a) S. Nie, L. Wu, L. Zhao, X. Zheng, S. Yang, P. Zhang, *Chem. Catal.* **2021**, *1*, 648–662; b) S. Hou, X. Ma, Y. Shu, J. Bao, Q. Zhang, M. Chen, P. Zhang, S. Dai, *Nat. Commun.* **2021**, *12*, 5917; c) S. Nie, L. Wu, L. Zhao, P. Zhang, *Nano Res.* **2021**, https://doi.org/10.1007/s12274-021-3803-3; d) Y. Yao, Z. Liu, P. Xie, Z. Huang, T. Li, D. Morris, Z. Finfrock, J. Zhou, M. Jiao, J. Gao, Y. Mao, J. J. Miao, P. Zhang, R. Shahbazian-Yassar, C. Wang, G. Wang, L. Hu, *Sci. Adv.* **2020**, *6*, eaaz0510.
- [16] a) F. Ye, T. Wang, X. Quan, H. Yu, S. Chen, *Chem. Eng. J.* **2020**, 389, 123427; b) J. F. Carneiro, M. J. Paulo, M. Siaj, A. C. Tavares, M. R. V. Lanza, *ChemElectroChem* **2017**, 4, 508–513; c) Z. Xu, J. Liang, Y. Wang, K. Dong, X. Shi, Q. Liu, Y. Luo, T. Li, Y. Jia, A. M. Asiri, Z. Feng, Y. Wang, D. Ma, X. Sun, *ACS Appl. Mater. Interfaces* **2021**, *13*, 33182–33187; d) M. Wang, N. Zhang, Y. Feng, Z. Hu, Q. Shao, X. Huang, *Angew. Chem. Int. Ed.* **2020**, *59*, 14373–14377; *Angew. Chem.* **2020**, *132*, 14479–14483; e) C. Liu, H. Li, J. Chen, Z. Yu, Q. Ru, S. Li, G. Henkelman, L. Wei, Y. Chen, *Small* **2021**, *17*, 2007249; f) L. R. Aveiro, A. G. M. da Silva, V. S. Antonin, E. G. Candido, L. S. Parreira, R. S. Geonmonond, I. C. de Freitas, M. R. V. Lanza,

P. H. C. Camargo, M. C. Santos, *Electrochim. Acta* **2018**, *268*, 101–110.

Angewandte

Chemie

- [17] J. Yin, Y. Li, F. Lv, Q. Fan, Y. Q. Zhao, Q. Zhang, W. Wang, F. Cheng, P. Xi, S. Guo, ACS Nano 2017, 11, 2275–2283.
- [18] T. Paik, M. Cargnello, T. R. Gordon, S. Zhang, H. Yun, J. D. Lee, H. Y. Woo, S. J. Oh, C. R. Kagan, P. Fornasiero, C. B. Murray, ACS Energy Lett. 2018, 3, 1904–1910.
- [19] S. Rehman, T. Tang, Z. Ali, X. Huang, Y. Hou, Small 2017, 13, 1700087.
- [20] K. A. Resende, A. H. Braga, F. B. Noronha, C. E. Hori, *Appl. Catal. B* 2019, 245, 100–113.
- [21] J. Liu, P. Wang, J. Fan, H. Yu, J. Energy Chem. 2020, 51, 253– 261.
- [22] J. Xia, H. Guo, M. Cheng, C. Chen, M. Wang, Y. Xiang, T. Li, E. Traversa, J. Mater. Chem. A 2021, 9, 2145–2151.
- [23] J. Varghese, T. Ghoshal, N. Deepak, C. O'Regan, R. W. Whatmore, M. A. Morris, J. D. Holmes, *Chem. Mater.* 2013, 25, 1458–1463.
- [24] N. Cao, Z. Chen, K. Zang, J. Xu, J. Zhong, J. Luo, X. Xu, G. Zheng, *Nat. Commun.* 2019, 10, 2877.
- [25] a) L. Zhang, W. Cai, N. Bao, Adv. Mater. 2021, 33, 2100745; b) Y. Ma, Y. Ma, Q. Wang, S. Schweidler, M. Botros, T. Fu, H. Hahn, T. Brezesinski, B. Breitung, Energy Environ. Sci. 2021, 14, 2883–2905; c) K. Mori, N. Hashimoto, N. Kamiuchi, H. Yoshida, H. Kobayashi, H. Yamashita, Nat. Commun. 2021, 12, 3884.
- [26] W. T. Mook, M. K. Aroua, M. H. Chakrabarti, C. T. J. Low, P. V. Aravind, N. P. Brandon, *Electrochim. Acta* **2013**, *94*, 327– 335.
- [27] J. Wu, Y. Zhou, H. Nie, K. Wei, H. Huang, F. Liao, Y. Liu, M. Shao, Z. Kang, J. Energy Chem. 2022, 66, 61–67.
- [28] a) J.-C. Dong, X.-G. Zhang, V. Briega-Martos, X. Jin, J. Yang, S. Chen, Z.-L. Yang, D.-Y. Wu, J. M. Feliu, C. T. Williams, Z.-Q. Tian, J.-F. Li, *Nat. Energy* **2019**, *4*, 60–67; b) J. Lai, B. Huang, Y. Tang, F. Lin, P. Zhou, X. Chen, Y. Sun, F. Lv, S. Guo, *Chem* **2018**, *4*, 1153–1166.
- [29] X. Mi, H. Liu, Y. Li, Energy Convers. Manage. 2019, 180, 196– 205.

Manuscript received: January 3, 2022 Accepted manuscript online: March 2, 2022 Version of record online: March 23, 2022