RSC Advances



PAPER

View Article Online
View Journal | View Issue



Cite this: RSC Adv., 2021, 11, 28949

Micro-area investigation on electrochemical performance improvement with Co and Mn doping in PbO₂ electrode materials†

Ze Lv, Zhen Chen, **D* Qiang Yu, Wei Zhu, Hongjun You, Bangyao Chen, Zhaoyi Zheng, Yuanyuan Liu and Qi Hu

PbO₂-Co₃O₄-MnO₂ electrodes, used in the electrowinning industry and in the degradation of organic pollutants, have demonstrated an elevated performance through macroscopic electrochemical measurements. However, few reports have investigated localized electrochemical performance, which plays an indispensable role in determining the essential reasons for the improvement of the modified material. In this study, the causes of the increase in electrochemical reactivity are unveiled from a micro perspective through scanning electrochemical microscopy (SECM), X-ray diffraction (XRD), Raman microscopy (Raman), and X-ray photoelectronic energy spectroscopy (XPS). The results show that the increase of electrochemical reactivity of the modified electrodes results from two factors: transformation of the microstructure and change in the intrinsic physicochemical properties. Constant-height scanning maps indicate that the electrochemical reactivity of the modified electrodes is higher than that of the PbO₂ electrode on the whole and high-reactivity areas are orderly distributed, coinciding with the observations from SEM and XRD. Thus, one of the reasons for the improvement of the modified electrode performance is the refinement of the microscopic morphology. The other reason is the surge of the oxygen vacancy concentration on the surface of the coating, which is supported by XRD, Raman and XPS. This finding is detected by the probe approach curve (PAC), which can quantitatively characterize the electrochemical reactivity of a substrate. Heterogeneous charge transfer rate constants of the modified electrode are 4-5 times higher than that of the traditional PbO₂ electrode. This research offers some insight into the electrochemical reactivity of modified PbO2 electrodes from a micro perspective.

Received 23rd May 2021 Accepted 14th August 2021

DOI: 10.1039/d1ra04006e

rsc.li/rsc-advances

1. Introduction

As a kind of ceramic electrode material, PbO₂ has excellent corrosion resistance and a strong oxidation ability, and it can withstand large currents. Despite the virtues mentioned above, there is huge room for improvement in terms of the electrochemical reactivity of the PbO₂ electrode. Researchers have modified and refined PbO₂ electrodes by inserting ions, 1,2 doping with active and inert particles, 3,4 and changing the structure of PbO₂. $^{5-7}$ As a result, the electrochemical reactivity of the PbO₂ electrode is enhanced and the application range of the PbO₂ electrode is expanded. In the electrocatalytic reaction, element doping can produce more oxygen defects, increase active sites, and improve the redox ability of electrode materials. In general, to verify the enhancement of new electrodes, the voltammetry charge (q_t) , charge transfer resistance (R_{ct}) and

Faculty of Science, Kunming University of Science and Technology, Kunming 650093, China. E-mail: chenzhen@kust.edu.cn

† Electronic supplementary information (ESI) available. See DOI: 10.1039/d1ra04006e

exchange current density extracted from cyclic voltammetry (CV), AC impedance (EIS) and linear sweep voltammetry (LSV), respectively, are persuasive evidence. However, these electrochemical analysis parameters obtained from macroscopic electrochemical measurements only reflect the average performance of the entire electrode area. They cannot illustrate the connection between the microscopic morphology, the difference of element composition and the macroscopic properties, which should not be ignored. This limitation has significantly hampered research on the mechanism of strengthening performance. The electrode reaction is a heterogeneous catalytic reaction. The catalytic reaction occurs at the interface between the electrode and electrolyte. The surface of the material provides a place for the liquid phase reactant molecules to participate in a reaction. Therefore, the microscopic morphology plays an important role in the reactivity and selectivity of electrocatalytic materials. Actually, some researchers have stated that the most important factor determining the catalytic performance is the surface morphology of the electrode material.8-11 It is a misconception that all tiny regions on an electrode share the same electrochemical

reactivity. To solve this problem, a research method with high spatial resolution is needed to analyze the electrochemical reactivity variation on different micro-regions and further to establish the relationship between electrode performance and microscopic morphology. SECM is a microscopic method that can better understand complicated electrochemical systems. SECM explores the reactivity changes at the micrometer-level and provides a brand new way to look into the reaction mechanism and kinetics of the electrode interface. At the same time, electrochemical imaging technology can collect the spatial analysis information of the substrate surface related to the surface topography structure and composition, creating a powerful approach to discover the relationship between the microtopography structure and the reactivity.

SECM can carry out electrochemical measurements on the micrometer scale. The ultramicro probe moves above an area close to the substrate and the Faraday current generated by the electrochemical reactions around the probe are recorded to study the electrochemical process and morphology of the substrate. SECM has evolved many valuable functions including the feedback mode, 12,13 substrate generation - probe collection mode,14 redox competitive mode,15 and surface inquiry mode.16 For one thing, SECM breaks through the limitations of traditional electrochemical methods with high-resolution, on-site, and three-dimensional observations, 17-19 which are closer to the actual situation of the substrate. For another thing, SECM can obtain the dependence of the electron transfer rate on the substrate potential and can also obtain quantitative information about the uneven distribution of the substrate. Wang²⁰ successfully characterized the relationship between electrochemical reactivity and crystallographic orientation on the surface of a polycrystalline platinum electrode with the help of SECM and electron backscatter diffraction (EBSD), they determined the structure-reactivity relationship of the electrocatalytic effect of the polycrystalline platinum electrode surface on the HOR in the sulfuric acid solution. The relationship shows that the catalytic activity of the hydrogen oxidation reaction (HOR) increases in the order of Pt(100) < Pt(110) < Pt(111) near the standard potential of hydrogen. Huai²¹ employed electron transfer rate constants to simulate and reevaluate the vulcanization reaction on malachite, combined with X-ray photoelectron spectroscopy (cryo-XPS) to characterize the corresponding surface morphology and confirmed the vulcanizing agent at two different concentrations, providing corresponding guidance for malachite flotation. Yao²² disclosed detailed information on the kinetics and reaction mechanism of Cu(II)/Cu(I) redox reactions determined SECM in systems with excess chloride. This work provides valuable insights for analyzing the electrode kinetics of unstable metal complexes in aqueous solutions. Zheng23 used SECM and EIS to study the localized electrochemical reactivity and heterogeneous charge transfer rate constant of the β-PbO₂ electrode. The results illustrated that the β-PbO₂ electrode is significantly different in terms of electrochemical reactivity. When the substrate potential is maintained at 0.5 V, the $[Fe(CN)_6]^{4-}$ oxidation reaction rate on the localized β-PbO₂ electrode is the largest, achieving the best constant-height image resolution; besides, the EIS also

demonstrated that the charge transfer process of $[Fe(CN)_6]^{4-}$ is the easiest when the substrate is 0.5 V.

In this project, SECM is employed to reveal the essential reasons for the enhancement of the performance of the PbO₂- Co_3O_4 -MnO₂ composite electrode in the $[Fe(CN)_6]^{3-}$ $[Fe(CN)_6]^{4-}$ system. At first, electrochemical information on the substrate is collected by moving the probe in a constant-height scanning method, thousands of current space coordinates are collected and a three-dimensional map is drawn, through which localized electrochemical reactivity in different micro areas is presented clearly. Then, the distribution regularity of the highreactivity area is expected to compare with the microscopic image showing morphology. So the relationship between modified morphology and electrochemical reactivity will be concluded. Secondly, the nonlinear fitting of the PAC at different substrate potentials is carried out to quantitatively study the electrochemical reaction reactivity difference of the micro-area and to explore the relationship between the characteristics of the modified electrode and its electrochemistry reactivity. Finally, XRD, Raman, XPS, and other characterization methods are combined to study the physicochemical properties of the electrode. SECM is an effective method to detect the electrochemical reactivity distribution on the surface of composite electrode, providing a new way to study modified PbO2 electrodes and the relationship between electrochemical reactivity and microscopic structure and intrinsic physicochemical properties.

2. Experiments and methods

Materials and preparation of the PbO_2 electrode and PbO_2 - Co_3O_4 - MnO_2 electrode

Lead nitrate and sodium fluoride (XILONG SCIENTIFIC CO., LTD), copper nitrate, cobalt nitrate, and 50% manganese nitrate water solution (Tianjin Fengchuan Chemical Reagent Technology Co., Ltd), and potassium ferricyanide and potassium chloride (shanghai Aladdin Biochemical Technology CO., LTD) were obtained.

The PbO $_2$ electrode was prepared by anode oxidation in a plating solution composed of 190 g L^{-1} Pb(NO $_3$) $_2$, 0.5 g L^{-1} NaF, and 15 g L^{-1} Cu(NO $_3$) $_2$. The anode was a stainless steel sheet (10 mm \times 50 mm \times 1 mm), the cathode was a stainless steel cylinder. The electrodeposition was conducted under a constant-current power source of 5 mA cm $^{-2}$ for 45 minutes, the temperature was 45 °C.

The β-PbO₂ intermediate layer was prepared *via* anodic oxidation at a current density of 5 mA cm⁻² for 15 minutes through using a constant-current power source. Stainless steel sheets were employed as the anode. A stainless steel cylinder served as the cathode. The composition of the electrodeposition solution was 190 g L⁻¹ Pb(NO₃)₂, 0.5 g L⁻¹ NaF, 0.5 g L⁻¹ AEO-7, and 15 g L⁻¹ Cu(NO₃)₂, the pH was 3, and the bath temperature was 40 $^{\circ}$ C.

The PbO_2 – Co_3O_4 – MnO_2 electrode was prepared by anode oxidation in a plating solution composed of 190 g L⁻¹ $Pb(NO_3)_2$, 15 g L⁻¹ $Cu(NO_3)_2$, 4 g L⁻¹ $Mn(NO_3)_2$, 40 g L⁻¹ $Co(NO_3)_2$, 0.5 g L⁻¹ NaF, and 10 g L⁻¹ citric acid. The anode was a β - PbO_2 intermediate layer, the cathode was a stainless steel cylinder. The electrodeposition was conducted under a constant-current power source of 5 mA cm⁻² for 45 minutes, the temperature was 45 °C.²⁴

Scanning electrochemical microscopy

A diagram of the SECM instrument is shown in Fig. S1 (ESI†). It is mainly composed of three parts: the electrochemical part (double potentiostat, probe, electrolytic cell, substrate, and various electrodes), the piezoelectric controller (to accurately control the position of the probe and base), and a computer (including an interface).

Scanning electrochemical microscopic imaging

The "feedback mode" and "substrate generation – probe collection mode" were used to characterize the substrate surface in the constant-height scanning maps. A SECM image is obtained by scanning the tip in the x–y plane and monitoring the UME tip current, i_T, as a function of the UME tip location, the so-called constant height mode. The feedback mode is based on a given probe potential and substrate potential, collecting the probe current with the distance d between the probe and the substrate to obtain a local electrochemical reactivity distribution map. In the experiment, the UME tip has a quiet time of 60 s and the step distance is 1 μ m, all experiments are in 10 mmol L⁻¹ K₃[Fe(CN)₆] and 0.1 mol L⁻¹ KCl solution.

Probe approach curves analysis

The PACs measurement adopts the feedback mode, and a PbO₂-Co₃O₄-MnO₂ electrode of about 10 mm \times 50 mm \times 1 mm was installed at the bottom of the electrolytic cell as the first working electrode, with an exposed area of 14.51 mm² (the area of the small holes in the electrolytic cell). The Pt ultramicrodisk electrode (Shanghai Chenhua company) with an effective radius $a = 5 \mu m$ and shielding coefficient RG = 1.5 (for related calculations, refer to ESI†) was used as the second working electrode. Ag/AgCl/3 mol L⁻¹ KCl serves as the reference electrode; Pt wire serves as an auxiliary electrode. 10 mmol L^{-1} $K_3[Fe(CN)_6]$ solution was used as the redox medium, 0.1 mol L⁻¹ KCl solution was used as the supporting electrolyte, and the probe voltage was 0.05 V vs. Ag/AgCl to control the probe tip reaction, as shown in reaction 1 to reduce $K_3[Fe(CN)_6]$. The probe stabilization status during the experiment is detailed in the ESI.†

$$[Fe(CN)_6]^{3-} + e^- \rightarrow [Fe(CN)_6]^{4-}$$
 (1)

As Fig. 1a illustrates, the probe was moved to the position 100 μm above the PbO₂-Co₃O₄-MnO₂ electrode keeping an open circuit potential. This location is theoretically far enough from the substrate thus the current detected in the tip is ruled by the flux of $[Fe(CN)_6]^{3-}$ from the bulk solution to the tip. This situation refers to "hemispherical diffusion" and a correlative detail mathematical formula is presented in the ESI.† First, the UME tip was parallel to the substrate until the UME tip was close to the surface of the substrate, and then the probe was returned to a certain distance and next an approximation was performed to obtain the corresponding probe approach curve. When the UME tip is close to the substrate, the PbO₂-Co₃O₄-MnO₂ electrode will interfere with the probe current in a positive feedback manner, the diffusion current will increase. As shown in Fig. 1b, this is due to the fact that the [Fe(CN)₆]⁴⁻ generated on the UME tip can be oxidized on the surface of the PbO₂-Co₃O₄-MnO₂ electrode, the current collected by the probe eventually increases. This process is called positive feedback diffusion. The corresponding oxidation reaction is represented by reaction (2). Fig. 1c shows the probe approach curves in the corresponding positive feedback mode. Under the positive feedback conditions, the electron transfer is enhanced, leading to a significant increase in the electron transfer rate constant k. This was to infer the essential reason for the improved performance of the modified electrode.

$$[Fe(CN)_6]^{4-} - e^- \rightarrow [Fe(CN)_6]^{3-}$$
 (2)

The nonlinear fitting of probe approach curves on a substrate

In this study, the nonlinear fitting method of PACs was used to quantitatively distinguish the electrochemical reactivity changes of the two electrodes by plotting the Y-axis as $I_{\rm T}$ ($I_{\rm T}=i_{\rm T}/i_{\rm T,\infty}$), the tip current normalized by the current is far from the substrate) and the X-axis as the L (L=d/a), the tip-substrate separation normalized by the tip radius) curve. Both X and Y are given in dimensionless form. Since this plot involves only dimensionless variables, it does not depend upon the concentration or diffusion coefficient of O. Through the nonlinear

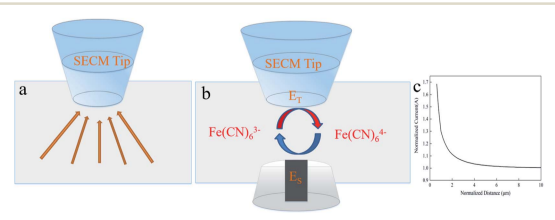


Fig. 1 (a) Hemispherical diffusion. (b) positive feedback diffusion, and (c) positive feedback curve.

fitting of the following equations, the kinetic information of the charge transfer process in the substrate can be described and the charge transfer rate constant k of the reaction can be determined. The equation details of the tip approach curves can be found in the ESI.†

Physical characterization analysis

The microscopic surface morphology of the PbO₂–Co₃O₄–MnO₂ electrode was obtained by SEM (Phenom ProX). XRD was used to analyze the phase composition and crystal structure. The chemical composition and elemental states on the PbO₂–Co₃O₄–MnO₂ electrode surface were analyzed through X-ray energy spectroscopy (EDS, EDAX-PHOENIX, EDAX, USA) and X-ray photoelectron spectroscopy (XPS, PHI5000 Versa ProbeII, Ulvac-PHI, Japan). Raman imaging on the surface of the substrate used a confocal Raman microscope (Raman, Renishaw) to analyze the Raman intensity at different positions of the substrate and the origin was used to map and analyze the Raman surface scan imaging data.

3. Results and discussion

Micromorphology and elemental composition of two electrodes

Fig. 2(a and b) shows the micromorphology of the PbO₂–Co₃O₄–MnO₂ electrode and PbO₂ electrode. Compared with the pure PbO₂ electrode, the microscopic morphology of the PbO₂–Co₃O₄–MnO₂ electrode has transformed into a spherical structure with a distinct boundary. After doping with Co₃O₄ and MnO₂, the grain size of PbO₂ is significantly refined, rendering

the grain smaller, compact and uniform. On the other hand, the PbO_2 prepared by the traditional way is pyramid-shaped, which is the typical structure and morphology of β - PbO_2 .

The EDS analysis results of the PbO2-Co3O4-MnO2 electrode and PbO₂ electrode are shown in Fig. 2c and d. The active layer of the PbO₂-Co₃O₄-MnO₂ electrode includes four elements: O, Pb, Co, and Mn. The SEM image and the corresponding element map are shown in Fig. S2 (ESI†). The content of the four elements is 72.99%. 23.53%, 1.34%, and 1.14%. The relative content of Co and Mn is low. During the electrodeposition process, the plausible explanation is that the process of incorporating Co and Mn into PbO2 involves at least two stages, the first stage is the chemical adsorption of Co2+ and Mn2+ on the growing PbO2 deposit; the second stage is the electro-oxidation of Co²⁺, Mn²⁺ to Co³⁺ and Mn⁴⁺ (Co₃O₄ and MnO₂) are incorporated within PbO2.25 With the increase of lead oxide on the electrode surface; the occurrence rate of the oxygen evolution reaction on the electrode surface is accelerated26 and the oxygen evolution reaction forms competition with oxidation reaction,27 ultimately leading to a decrease in the deposition of Mn and Co.

The relationship between electrochemical reactivity and morphology and structure of microspheres

SECM is a high spatial resolution method for the micro-area on the electrode. It can directly obtain surface electrochemical reactivity information associated with localized morphology. Under the feedback mode of SECM, the probe is maintained at a constant height. In a solution of 10 mmol $\rm L^{-1}$ $\rm K_3[Fe(CN)_6]$ and 0.1 mol $\rm L^{-1}$ KCl, the constant-height surface scanning of the two electrodes is conducted when the probe is maintained at

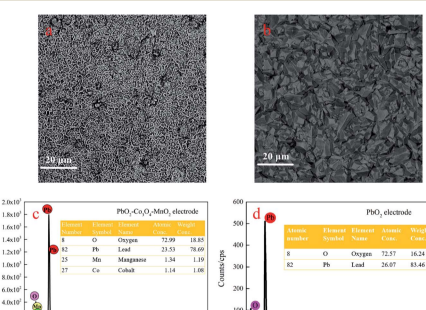


Fig. 2 SEM images of (a) the $PbO_2 - Co_3O_4 - MnO_2$ electrode and (b) the PbO_2 electrode. (c) Energy spectrogram analysis for the $PbO_2 - Co_3O_4 - MnO_2$ composite electrode. (d) Energy spectrogram analysis for the PbO_2 electrode.

6

Energy/KeV

Paper

i_/nA i /nA 70 70 1.728 1.640 60 60 1.692 1.535 50 50 40 40 30 30 1.620 1.324 20 20 10 584 1 218 30 40 50 30 40 50 20 60 10 20 60

Fig. 3 SECM map of the $80 \mu m \times 80 \mu m$ constant-height surface at the open circuit potential of (a) the PbO₂-Co₃O₄-MnO₂ electrode and (b) the traditional PbO₂ electrode, $d=3 \mu m$.

0.05 V (vs. Ag/AgCl) and the substrate potential is open-circuit potential. Fig. 3 presents constant-height surface scanning of the two electrodes at the open-circuit potential on an 80 $\mu m \times$ 80 μm scale, the UME tip is 3 µm away from the substrate. All tip currents are normalized with $i_{T,\infty}$, the correspondence relationship between the normalized currents and colors of all graphics uses the scale on the right. As shown in Fig. 3, every point on the constant-height surface scanning shows positive feedback (which enhances the oxidation of $[Fe(CN)_6]^{4-}$). The probe current detected during the scanning is proportional to the oxidation capacity of the corresponding micro-area on the substrate. Therefore, the highreactivity area with a higher current is the red area and the lowreactivity area with a lower current is blue. As the feedback current is a function of the probe-substrate distance, the higher position corresponds to the higher current (red area) and the lower position corresponds to the lower current (blue area).

X (μm)

Fig. 3a shows that the active area of the PbO₂–Co₃O₄–MnO₂ electrode is spherical, highly coinciding with the sizes, shapes, and distribution of the PbO₂–Co₃O₄–MnO₂ morphology observed by SEM. What is striking about the constant-height scanning image of the PbO₂–Co₃O₄–MnO₂ electrode is that electrochemical reactivity on the spherical center is higher than on the edge around the circle, and the active area of the electrode as a whole is evenly distributed. In contrast, the active region of the traditional PbO₂ electrode shown in Fig. 3b is not uniformly distributed. Also, judging from the normalized currents, the electrochemical reactivity of the modified electrode is higher than that of the traditional PbO₂ electrode.

Several factors are known to be responsible for this improvement. After the correction of the PbO₂–Co₃O₄–MnO₂ electrode due to the addition of two elements, the grain shape changes significantly, the grain size decreases, and the degree of tightness between the grains increases after the addition of two elements, thereby reducing the variation between the high and low active areas and distributing active points on the electrode surface in a uniform way.

Since the probe tip current is normalized, the currents in the two pictures are comparable. The probe current of the modified electrode is significantly higher than the traditional PbO₂ electrode. SECM results demonstrate that the electrocatalytic reactivity of the PbO₂–Co₃O₄–MnO₂ electrode is elevated. Evidently, the transformation of the morphology contributes to this enhancement. Due to the doping of Co and Mn elements, the preferred growth orientation of the PbO₂ changes, and the grains are refined into a spherical shape. With the specific surface area of the electrode increasing, more active area is fully exposed to the electrolyte for the electrochemical reaction, providing more positions for electron transfer. The presented evidence supports the idea that the catalytic performance of the PbO₂–Co₃O₄–MnO₂ electrode is closely related to the topography of the electrode surface.^{28,29}

Probe current curves (red lines) produced by the tip scanning a certain distance over the surface of the two electrodes are displayed on the top half of Fig. 4. What stands out in this chart is the pattern of the curve. Due to the messy topography of the traditional PbO_2 electrode, electrochemical reactivity of the traditional PbO_2 electrode is unevenly distributed. Regarding the

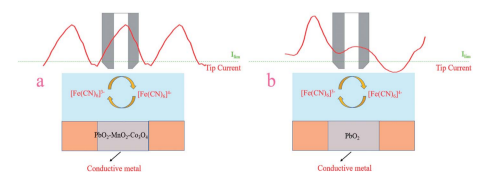


Fig. 4 The positive feedback response curve of the probe current at open circuit potential for (a) the $PbO_2-Co_3O_4-MnO_2$ electrode and (b) traditional PbO_2 electrode.

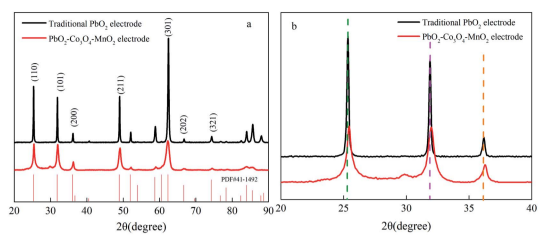


Fig. 5 (a) XRD patterns of the PbO₂ electrode (black) and PbO₂ – Co_3O_4 – MnO_2 electrode (red). (b) Local magnified XRD patterns of the 20° – 40° .

modified PbO₂ electrode, the shape is transformed into a uniform spherical shape and thus the active area is evenly distributed and the gap of electrochemical reactivity among different points narrows. These results further support the hypothesis that the electrochemical reactivity of the PbO₂–Co₃O₄–MnO₂ electrode has a remarkable correlation with the morphology, and the electrochemical reactivity of the modified electrode is enhanced by the refinement of the microscopic structure.

The grain structural information of the samples was revealed by XRD. The XRD patterns of the traditional PbO $_2$ electrode and PbO $_2$ –Co $_3$ O $_4$ –MnO $_2$ electrode are shown in Fig. 5. In Fig. 5a, the peaks of two electrodes are strong at 25.4°, 31.9°, 36.2°, 49.4°, and 62.5°, which could be ascribed to β -PbO $_2$ (PDF#41-1492). The results illustrate that these two electrodes mainly consist of β -PbO $_2$. Besides, it can be seen that the diffraction peak intensity and half-width of the peaks from the two electrodes are significantly different. The diffraction peak intensity of the modified electrode is weaker than that of the traditional PbO $_2$ electrode, indicating that the preferred crystal orientation of the β -PbO $_2$ was converted because of the ion-doped modification. In addition, the half-peak width of the modified electrode is larger than that of the traditional PbO $_2$ electrode. According to the Debye–Scherrer formula,

$$D = \frac{K\lambda}{\beta \cos \theta}$$

the average crystal size of the PbO_2 – Co_3O_4 – MnO_2 electrode is estimated to be 10.45 nm, while the traditional PbO_2 electrode is 15.68 nm. This result proves that the crystal size of the modified electrode is smaller than that of the traditional electrode.

Fig. 5b is an amplified XRD image of $20^{\circ}-40^{\circ}$. It shows that the (101), (110), and (200) crystal planes in the PbO₂–Co₃O₄–MnO₂ electrode shift into higher diffraction angles, this is because the Co and Mn element doped into the PbO₂ unit cell have smaller radii than the Pb element, causing the diffraction peaks of the doped modified electrode to shift to a larger angle. The unit cell parameters and unit cell volume are correspondingly reduced (as shown in Table 1). The smaller size of the crystals after doping Co and Mn into the PbO₂ lattice meant

more point defects formed within the PbO₂ lattice, which also shows that the concentration of surface oxygen vacancies is greater than that of traditional PbO₂.³⁰ Both Co and Mn oxides are rich in oxygen vacancies.³¹ Under the influence of the two elements, the diffraction peak intensity of the modified electrode is lower than that of the traditional PbO₂ electrode and the growth of each crystal plane is inhibited, reshaping the preferred growth orientation of the grains.

In summary, after the addition of more conductive particles into the PbO₂ electrode, the added particles will not only undergo electrochemical oxidation but will also replace the preferential growth sites, causing the surface morphology of the electrode material to change. The addition of Co and Mn changes the micro-morphology of the original PbO₂ electrode, which not only increases the specific surface area of the electrode material but also reduces the grain spacing on the surface of the electrode material. The greater the number of atoms exposed to the electrolyte, the more active sites there are on the electrode surface and the better the catalytic reactivity.³²

The relationship between the electrochemical reactivity of the electrodes and the electrodes' physicochemical properties

The current response of the SECM constant-height surface scanning mainly depends on the micro-topography, conductivity, and electrochemical reactivity of the substrate. The advantage of PAC is that it can interpret localized electrochemical reactivity without any influence from the micro-topography of the substrate. Therefore, the positive feedback mode of SECM is adopted to study the variation in the electrochemical reactivity of the substrate under different potentials. The feedback current at the probe is determined by the

Table 1 Lattice constants of two electrodes

Electrode	A (Å)	B (Å)	C (Å)	V	Crystallite-size/nm
PbO ₂ PbO ₂ -Co ₃ O ₄ -MnO ₂				83.450 82.900	

localized intrinsic physicochemical properties of the substrate and can reflect the difference in the reactivity of the substrate surface. With a tiny distance between the probe and the substrate, mass transfer can take place in a rapid way. Being immune to the influence of concentration polarization is the theoretical basis for the SECM to quickly determine the electron transfer reaction. When the electron transfer step is treated as a speed control step, the change in substrate potential directly affects the speed of the electron transfer step on the substrate and the entire electrode reaction process. The obtained PAC is subjected to nonlinear fitting to compute a heterogeneous reaction rate constant. This rate constant can adequately mirror the conductivity and electrochemical reactivity of the substrate surface.33-35

Basing on the theoretical model established by Lefrou, 36 the nonlinear fitting eqn (5) (ESI†) is used to achieve the best fit between the theoretical PAC and the experimental PAC, and the normalized abscissa is used. The calculation of the theoretical calculation of the SECM tip approach curves can be found in the ESI.† Fig. 6 shows the probe approach curves fitting diagrams of points A and B on the two substrate electrodes with different potentials. The two points A and B are active sites on the two electrodes, respectively. The black curve in the figure is the initial experimental data and the red dotted line is the probe fitting. The theoretical approximation curve, in all cases, fits the experimental data very well. The heterogeneous charge transfer rate constant k of the regeneration process of the electrochemically active medium is obtained. It can be seen from Table

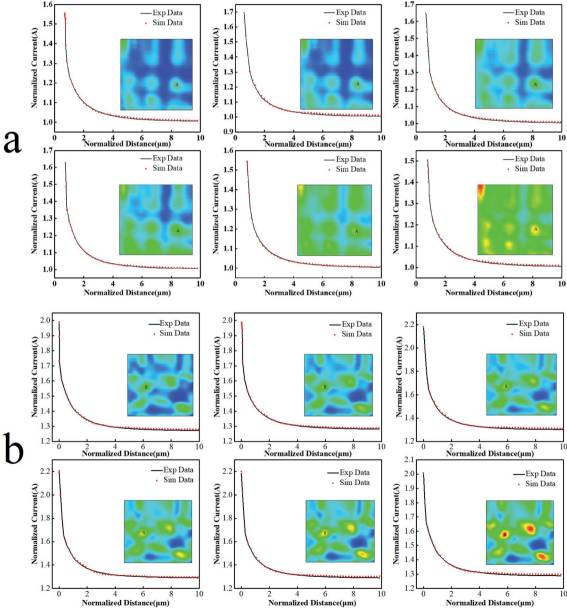


Fig. 6 The UME probe approach curve with (a) the PbO₂-Co₃O₄-MnO₂ electrode and (b) the traditional PbO₂ electrode, $E_t = 0.05 \text{ V}$, $E_s = 0.1$ -1.0 V, increment distance = 1 μ m, quiet time = 60 s.

Table 2 The values of charge transfer rate constants at points A and B at different potentials

Electrode	0.1 V	0.3 V	0.5 V	0.7 V	0.9 V	1.0 V
PbO ₂ -Co ₃ O ₄ -MnO ₂ PbO ₂	8.387 2.574		8.701 2.657			10.364 3.135

2 that with the substrate potential becoming more positive, the k value gradually increases, suggesting that the electron transfer on the surface becomes easier and the reactivity of active regions of the two electrodes increases. Accordingly, the rate of the Fe²⁺ oxidation reaction on the substrate speeds up and thus a large

amount of Fe³⁺ diffuses to the UME surface. As the concentration of the redox medium rises, the reduction current detected by the probe increases.

Whether it is a traditional PbO_2 electrode or a modified PbO_2 electrode, the charge transfer rate constant of both increases with the increase in the potential, as does the reactivity of the active site. Under the same potential, the heterogeneous charge transfer rate constant of the modified electrode is about 3 times higher than that of the traditional electrode. It is indicated that the electrochemical reactivity of the modified electrode is enhanced due to the change of its physical and chemical properties.

As discussed above, the alteration of physicochemical properties gives rise to the enhancement of electrochemical

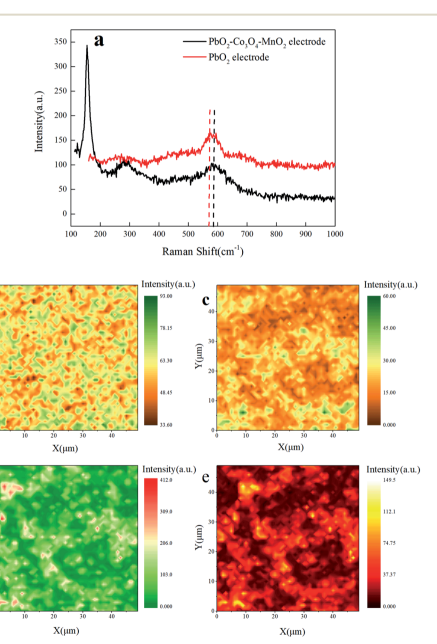


Fig. 7 (a) The Raman spectrum of the $PbO_2-Co_3O_4-MnO_2$ electrode between 0 and 1000 cm⁻¹ using 532 nm excitation. (b) The Raman microscopy image of the PbO_2 electrode is composed of 576 cm⁻¹ correlation spectral distribution. The Raman microscopy image of the $PbO_2-Co_3O_4-MnO_2$ electrode. (c) The relative spectral distribution composition of 591 cm⁻¹. (d) The relative spectral distribution composition of 143 cm⁻¹.

Y(µm)

reactivity on the PbO₂-Co₃O₄-MnO₂ electrode unveiled by the PAC experiments. To further clarify the alteration of the physicochemical properties, Raman microscopy was utilized to identify the mechanism of doped elements by capturing their Raman intensity. Fig. 7a illustrates that the peak at 576 cm⁻¹ is attributed to the Raman active vibrational mode of Pb-O in PbO₂.37 After doping with Co and Mn elements, a new band emerges at 143 cm⁻¹, which is attributed to the F_{2g} mode of Co-O;38 the weak peak at 274 cm⁻¹ is assigned to the band of Mn-O.39 It shows that Mn and Co were successfully deposited into the modified electrode. It also can be observed that the main band position at 591 cm⁻¹ of the modified electrode is blueshifted relative to the main peak position of the traditional

PbO₂ electrode and the intensity of the Raman spectrum is weakened and the width becomes wider, indicating that Co and Mn are successfully doped into the electrode and part of it enters the PbO2 lattice. Meanwhile, the number of oxygen vacancies on the surface of the modified electrode increases40 and the electrochemical reactivity is enhanced, which is consistent with the results of XRD.

Using the above spectrum as a reference, information about the distribution of the electrode component can be obtained. Fig. 7b is the Raman microscopy image of the main band position of the PbO₂ electrode. In Fig. 7b, the shape of the green region with high Raman intensity is similar to the rutile structure of PbO2 and the Raman intensity distribution is consistent

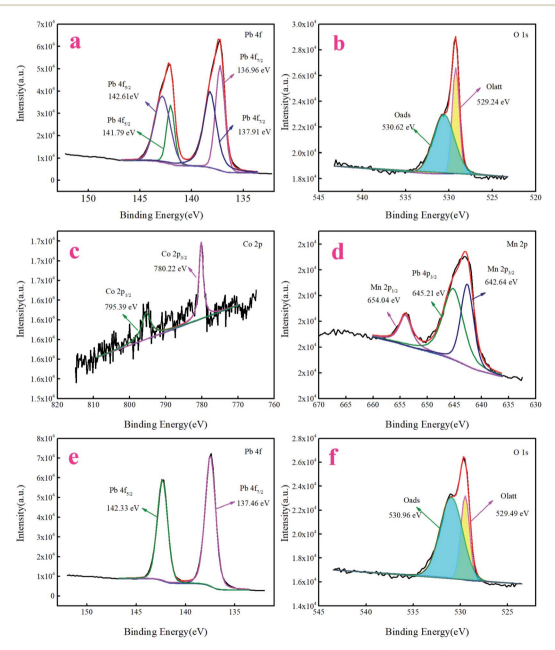


Fig. 8 XPS spectrum and fitted data of (a) Pb 4f, (b) O 1s of the as-obtained PbO₂ electrode and (c) Co 2p, (d) Mn 2p, (e) Pb 4f, and (f) O 1s of the as-obtained PbO₂-Co₃O₄-MnO₂ electrode.

RSC Advances Paper

with the electrochemical reactivity distribution (Fig. 3b). Fig. 7c-e are the Raman mapping scanning images of the modified electrode conducted at 591 cm⁻¹, 143 cm⁻¹, and 274 cm⁻¹, respectively. It can be seen from the figures that the addition of Co and Mn elements makes the Raman intensity and distribution of Pb in the modified electrode significantly different. Fig. 7d and e are the Raman mapping scanning images under the characteristic band of the Co and Mn elements. The Raman intensity is high in the middle and low on the edge, roughly coinciding with the electrochemical reactivity distribution of the PbO₂-Co₃O₄-MnO₂ electrode (Fig. 3a). These results confirm the association between the doping of Co and Mn elements and the transformation of the microscopic morphology which offers more active electrochemical reaction sites. This also accords with our earlier observations on SECM.

Coupled with the Raman spectra results, the second reason for the enhancement of electrochemical reactivity is increasing the number of surface oxygen vacancies which results from the entrance of Co and Mn elements into the PbO₂ lattice. Namely, it adequately explains how the physicochemical properties change, making conclusions from PAC more insightful and comprehensive.

XPS is an analysis technique for the active components of electrode. It can study the relationship between the composition and performance of the active phase and infer the reaction mechanism. X-ray photoelectron spectroscopy (XPS) was carried out to study the compositions and redox behavior of two electrodes. The full spectrum of the two electrodes (Fig. S3 and S4†) can be found in the ESI.† Fig. 8a and b illustrates the Pb 4f and O 1s XPS spectra of traditional PbO2. As can be seen from Fig. 8a, the binding energy of Pb 4f is 136.96 eV, 137.91 eV, 141.79 eV, and 142.61 eV, respectively, a 4.7 eV and 4.83 eV energy variation between the Pb 4f_{7/2} and Pb 4f_{5/2} peaks is observed, evidencing the existence of PbO2 in the traditional electrode.41 The O 1s species in the high-resolution XPS spectra (Fig. 8b) revealed signals at 529.24 eV and 530.62 eV, corresponding to surface lattice oxygen (Olatt) and surface adsorption oxygen (Oads).42

For the modified electrode, the Co 2p XPS spectra (Fig. 8c) show two major peaks at 780.22 eV and 795.39 eV, which are ascribed to the typical Co 2p_{3/2} and Co 2p_{1/2} orbitals, respectively. A 15 eV energy variation between the Co $2p_{3/2}$ and Co $2p_{1/2}$ 2 peaks is observed, evidencing the existence of Co₃O₄ in the modified electrode. 43,44 From Fig. 8d, one can observe the appearance of two signals at BE = 642.64 eV and 654.04 eV, assignable to Mn 2p_{3/2} and Mn 2p_{1/2}, respectively, and the interval between the two peaks is 11.4 eV, which indicates that the oxidation state of Mn in the modified electrode is MnO₂ (ref. 45) because the binding energy in the figure is 645.21 eV, which is Pb 4p. Since the binding energy of the stronger Pb 4p peak, Pb 4p_{3/2} is similar to that of the O 1s peak, the two peaks coincide and need to be fitted. The Pb 4f XPS spectra (Fig. 8e) for the modified electrode exhibited two well-defined, symmetric peaks centered at 137.46 eV and 142.33 eV, corresponding to $4f_{7/2}$ and $4f_{5/2}$, respectively, which are consistent with the spectral values for PbO₂.41,46 As can be seen from the O 1s spectrum (Fig. 8f), two peaks centered at 529.49 eV and 530.96 eV arise from lattice oxygen (O_{latt}) and adsorbed oxygen (O_{ads}). 42,47

The changes in oxygen species surface coverage and binding strength cause the change of the electrochemical reactivity of the modified electrode. To determine the existence of oxygen on the electrode surface and quantitatively calculate the relationship between oxygen species on the surface and doping elements, the peak separation fitting results of the O 1s spectrum peaks of the two electrodes are listed in Table 3. The redox performance of the electrocatalytic materials mainly depends on the activation of molecular oxygen and the type of surface or lattice oxygen.48 For an oxygen-deficient material, the adsorbed oxygen species on the surface of the PbO₂ electrode is considered to be more active and more important for the oxidation reaction. Typically, lattice oxygen species are involved in selective oxidation reactions, while adsorbent oxygen species such as O^{2-} , O_2^{2-} , and O^- are given priority in the process of complete oxidation.49 Therefore, in the catalytic process, adsorbed oxygen is the most active oxygen species. Moreover, the concentration of adsorbed oxygen is closely related to the number of oxygen vacancies and the ratio of Oads/Olatt can be used to illustrate the relative abundance of oxygen vacancies on the electrode material surface. Therefore, compared to the undoped PbO2 electrode, it was found that the adsorbed oxygen content of the modified electrode is higher than that of the traditional PbO₂ electrode and the Oads/Olatt value of the modified doped electrode is greater than that of the traditional PbO2 electrode, which means that there is more active oxygen on the surface of the doped modified electrode. The species and the number of oxygen vacancies also increased correspondingly, indicating that the doped modified electrode materials have better electrochemical reactivity. The results discussed above show that the role of the doping element is to change the surface properties of PbO₂.

In addition, previous studies on doping WC particles⁵⁰ and CNTs⁵¹ concluded that improvement of the electrodes is driven by their good electrical conductivity and fast electron transfer. These doping substances present on the electrodes in a way of

Table 3 XPS fitting parameters and crystallites of the PbO₂ electrode and PbO₂-Co₃O₄-MnO₂ electrode

		Binding energ	gy/eV			
Electrode	Crystallite-size/nm	O 1s (O _{ads})	O 1s (O _{latt})	O _{ads} /O _{latt} (atom%)	O _{ads} (content%)	O _{latt} (content%)
PbO ₂	15.68	530.62	529.25	1.06	51.46	48.54
PbO ₂ -Co ₃ O ₄ -MnO ₂	10.45	530.96	529.49	2.31	69.79	30.21

Paper **RSC Advances**

physical combination and then enhance the electrochemical reactivity. Concerning the PbO2-Co3O4-MnO2 electrode, some Co and Mn ions have intruded into the PbO₂ lattice. Hence, more oxygen vacancies are successfully introduced into the doped modified electrode. This conclusion has been verified through XRD, Raman, and XPS. In other words, the promotion of the PbO2-Co3O4-MnO2 electrode does not rely on the formation of Co and Mn oxides. The oxides do not directly participate in the electrocatalytic reaction. In a word, the transformation of the microscopic morphology and increasing of oxygen vacancies are important driving factors of enhancement of electrochemical reactivity on the PbO2-Co3O4-MnO5 electrode.

Conclusions

This paper proposes that the performance improvement of the doped modified electrode is due to the change of the microscopic morphology of the modified electrode and the increase of the oxygen vacancy concentration on the electrode surface.

The feedback mode of SECM successfully observed the electrochemical reactivity distribution of the PbO2-Co3O4-MnO2 electrode and the traditional PbO2 electrode and concluded that the electrochemical reactivity of the modified electrode was higher than that of the traditional PbO₂ electrode. Combined with SEM, the shape of the electrochemically active region is consistent with the micro-topography structure. Through XRD, the microscopic morphology and structure of the modified electrode are observed to have changed due to the doping of Co and Mn elements. In addition, comparing the SECM constant height surface scan with the Raman surface scan can conclude that the electrochemical reactivity distribution is related to the morphology of the substrate. The active area of the modified electrode is distributed in a circular shape, while the traditional PbO₂ electrode presents a pyramid shape. In summary, the performance improvement of the modified electrode is partly due to the change of the micromorphology.

For another thing, the nonlinear fitting of the PACs quantitatively analyzes the localized reactivity of the two electrode surfaces. The heterogeneous reaction rate constant of the modified electrode is about 3 times higher than that of the traditional PbO2 electrode. Finally, combining the XPS and Raman results, the increase in the k of the modified electrode stems from the increase in the concentration of surface oxygen vacancies, it is these oxygen vacancies serving as active sites that give a boost to the electrochemical reactivity of the modified electrode. In contrast to the mechanism of doping WC particles and CNTs, the promotion of the PbO₂-Co₃O₄-MnO₂ electrode does not rely on the formation of Co and Mn oxides and they do not directly participate in the electrocatalytic reaction. It is believed that this investigation provides a new micro perspective on the causality toward electrochemical reactivity improvement of PbO2-Co3O4-MnO2 electrodes and provides a new platform for the study of the structure-reactivity relationship of the electrode surface, which can be readily extended to many electrochemical processes of other electrode materials.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was financially supported by the National Natural Science Foundation of China (No. 51464021).

Notes and references

- 1 Y. Zhou, Z. Li, C. Hao, Y. Zhang, S. Chai, G. Han, H. Xu, J. Lu, Y. Dang, X. Sun and Y. Fu, Electrochim. Acta, 2020, 333, 135535-135547.
- 2 Z. Souri, A. Ansari, D. Nematollahi and M. Mazloum-Ardakani, J. Electroanal. Chem., 2020, 862, 114037-114052.
- 3 X. Wang, R. Xu, S. Feng, B. Yu and B. Chen, RSC Adv., 2020, 10, 1351-1360.
- 4 C. Chen, X. Wang, R. Xu, Y. Zhang, S. Feng, A. Ju and W. Jiang, RSC Adv., 2021, 11, 6146-6158.
- 5 L. Gui, Z. Chen, B. Chen, Y. Song, Q. Yu, W. Zhu, Q. Hu, Y. Liu, Z. Zheng, L. Ze, H. You and F. Yeasmin, J. Hazard. Mater., 2020, 399, 123018-123033.
- 6 N. Comisso, S. Cattarin, P. Guerriero, L. Mattarozzi, M. Musiani and E. Verlato, Electrochim. Acta, 2016, 200,
- 7 S. Abaci, U. Tamer, K. Pekmez and A. Yildiz, Appl. Surf. Sci., 2005, 240, 112-119.
- 8 S. Ghasemi, M. F. Mousavi, H. Karami, M. Shamsipur and S. H. Kazemi, *Electrochim. Acta*, 2006, **52**, 1596–1602.
- 9 D. Pech, T. Brousse, D. Bélanger and D. Guay, Electrochim. Acta, 2009, 54, 7382-7388.
- 10 Z. Wang, Y. Liu and V. M. Linkov, J. Power Sources, 2006, 160, 326-333.
- 11 Z. Li, Y. Yang, A. Relefors, X. Kong, G. M. Siso, B. Wickman, Y. Kiros and I. L. Soroka, J. Colloid Interface Sci., 2021, 583,
- 12 C. G. Zoski, B. Liu and A. J. Bard, Anal. Chem., 2004, 76, 3646-
- 13 D. S. A. Yin Jing, S. Mehraeen, R. J. LeSuer and B. P. Chaplin, J. Mater. Chem. A, 2018, 6, 23828-23839.
- 14 X. Zeng, D. Liu, S. Wang, S. Liu, X. Cai, L. Zhang, R. Zhao, B. Li and F. Kang, ACS Appl. Mater. Interfaces, 2020, 12, 37047-37053.
- Morkvenaite-Vilkonciene, A. Ramanaviciene 15 I. A. Ramanavicius, RSC Adv., 2014, 4, 50064-50069.
- 16 D. Zigah, J. Rodriguez-Lopez and A. J. Bard, Phys. Chem. Chem. Phys., 2012, 14, 12764-12772.
- 17 L. Danis, S. M. Gateman, C. Kuss, S. B. Schougaard and J. Mauzeroll, ChemElectroChem, 2017, 4, 6–19.
- 18 A. J. Bard, F. R. Fan, D. T. Pierce, P. R. Unwin, D. O. Wipf and F. Zhou, Science, 1991, 254, 68-74.
- 19 H. Luo, C. Dong, S. Gao, C. Du, K. Xiao and X. Li, RSC Adv., 2014, 4, 56582-56595.
- 20 Y. Wang and D. O. Wipf, J. Electrochem. Soc., 2020, 167, 146502-146511.

- 21 Y. Huai, Y. Qian and Y. Peng, *Appl. Surf. Sci.*, 2020, **531**, 147334–147342.
- 22 Y. Meng, J. Chen and K. Liang, *Anal. Chem.*, 2020, **92**, 10420–10424.
- 23 Z. Zheng, Q. Yu, Z. Chen, W. Zhu, Q. Hu, Y. Liu, L. Gui and Y. Song, J. Electroanal. Chem., 2020, 878, 114699–114707.
- 24 Y. Liu, W. Zhu, Z. Chen, Q. Yu, Q. Hu, Z. Zheng, L. Gui and Y. Song, *Int. J. Hydrogen Energy*, 2021, **46**, 6380–6394.
- 25 A. B. Velichenko, R. Amadelli, E. A. Baranova, D. V. Girenko and F. I. Danilov, *J. Electroanal. Chem.*, 2002, 527, 56–64.
- 26 Y. Yao, C. Huang, H. Dong, F. Wei and X. Chen, *Russ. J. Electrochem.*, 2019, 55, 364–369.
- 27 A. B. Velichenko, E. A. Baranova, D. V. Girenko, R. Amadelli, S. V. Kovalev and F. I. Danilov, *Russ. J. Electrochem.*, 2003, 39, 615–621
- 28 J. González-Prior, R. López-Fonseca, J. I. Gutiérrez-Ortiz and B. de Rivas, *Appl. Catal., B*, 2016, **199**, 384–393.
- 29 Z. Chen, S. Wang, W. Liu, X. Gao, D. Gao, M. Wang and S. Wang, *Appl. Catal.*, A, 2016, 525, 94–102.
- 30 Q. Ma, Z. Chen, S. Zhong, J. Meng, F. Lai, Z. Li, C. Cheng, L. Zhang and T. Liu, *Nano Energy*, 2021, 81, 105622–105633.
- 31 Z. Han, Y. Liu, J. Deng, S. Xie, X. Zhao, J. Yang, K. Zhang and H. Dai, *Catal. Today*, 2019, **327**, 246–253.
- 32 W. J. Xue, Y. F. Wang, P. Li, Z.-T. Liu, Z. P. Hao and C. Y. Ma, *Catal. Commun.*, 2011, **12**, 1265–1268.
- 33 S. E. Pust, D. Scharnweber, C. Nunes Kirchner and G. Wittstock, *Adv. Mater.*, 2007, **19**, 878–882.
- 34 S. E. Pust, D. Scharnweber, S. Baunack and G. Wittstock, *J. Electrochem. Soc.*, 2007, **154**, C508–C514.
- 35 A. Maho, F. Kanoufi, C. Combellas, J. Delhalle and Z. Mekhalif, *Electrochim. Acta*, 2014, 116, 78–88.
- 36 C. Lefrou, J. Electroanal. Chem., 2006, 592, 103-112.

- 37 H.-W. Yeh, C.-J. Chang, G. G. Huang and P.-Y. Chen, *J. Electroanal. Chem.*, 2019, **834**, 64–70.
- 38 L. Ma, S. Chen, H. Li, Z. Ruan, Z. Tang, Z. Liu, Z. Wang, Y. Huang, Z. Pei, J. A. Zapien and C. Zhi, *Energy Environ.* Sci., 2018, 11, 2521–2530.
- 39 T. Gao, M. Glerup, F. Krumeich, R. Nesper, H. Fjellvag and P. Norby, *J. Phys. Chem. C*, 2008, **112**, 13134–13140.
- 40 Y. C. Huang, B. Long, M. N. Tang, Z. B. Rui, M. S. Balogun, Y. X. Tong and H. B. Ji, *Appl. Catal.*, *B*, 2016, **181**, 779–787.
- 41 J. Morales, G. Petkova, M. Cruz and A. Caballero, *J. Power Sources*, 2006, **158**, 831–836.
- 42 Y. Xia, X. Bian, Y. Xia, W. Zhou, L. Wang, S. Fan, P. Xiong, T. Zhan, Q. Dai and J. Chen, Sep. Purif. Technol., 2020, 237, 116321–116328.
- 43 K. Xiang, Z. Xu, T. Qu, Z. Tian, Y. Zhang, Y. Wang, M. Xie, X. Guo, W. Ding and X. Guo, *Chem. Commun.*, 2017, 53, 12410–12413.
- 44 C. Mahala, M. D. Sharma and M. Basu, *New J. Chem.*, 2019, 43, 15768–15776.
- 45 E. Hastuti, A. Subhan, P. Amonpattaratkit, M. Zainuri and S. Suasmoro, RSC Adv., 2021, 11, 7808–7823.
- 46 K. Kannan, G. Muthuraman and I. S. Moon, *Mater. Lett.*, 2014, 123, 19–22.
- 47 S. H. Xie, Y. X. Liu, J. G. Deng, J. Yang, X. T. Zhao, Z. Han, K. F. Zhang and H. X. Dai, *J. Catal.*, 2017, 352, 282–292.
- 48 Y. Li and W. Shen, Chem. Soc. Rev., 2014, 43, 1543-1574.
- 49 A. Amri, X. Duan, C.-Y. Yin, Z.-T. Jiang, M. M. Rahman and T. Pryor, *Appl. Surf. Sci.*, 2013, 275, 127–135.
- 50 Y.-Y. Ma, Z.-L. Lang, L.-K. Yan, Y.-H. Wang, H.-Q. Tan, K. Feng, Y.-J. Xia, J. Zhong, Y. Liu, Z.-H. Kang and Y.-G. Li, *Energy Environ. Sci.*, 2018, 11, 2114–2123.
- 51 Y. Xia, J. Feng, S. Fan, W. Zhou and Q. Dai, *Chemosphere*, 2021, 263, 128069-128078.