

**Showcasing research from Professor Ming-Hua Zeng's laboratory, College of Chemistry & Chemical Engineering, Hubei University, Wuhan, China.**

Tracking the pyrolysis process of a 3-MeOsalophen-ligand based Co<sub>2</sub> complex for promoted oxygen evolution reaction

Efficient oxygen evolution reaction catalysts can be prepared *via* controlled pyrolysis of molecular platforms, and there is still minimal mechanistic understanding of such pyrolysis processes. Here, a 3-MeOsalophen-ligand based Co<sub>2</sub> complex is introduced as a precursor *via* controlled pyrolysis and systematically analysed by TG-MS, and can be used as an efficient OER electrocatalyst with a distinct core-shell Co/CoOx@NC nanostructure. This work offers quiding insight into controlled pyrolysis *via* TG-MS analysis, using a novel complex precursor for precise regulation of heteroatom-doped (3d) transition metal-based electrocatalysts.

## **As featured in:**



See Xu Peng, Ming-Hua Zeng *et al.*, *Chem. Sci.*, 2019, **10**, 4560.

rsc.li/chemical-science



# **Chemical Science**

## EDGE ARTICLE



Cite this: Chem. Sci., 2019, 10, 4560

All publication charges for this article have been paid for by the Royal Society of Chemistry

Received 29th January 2019 Accepted 11th March 2019

DOI: 10.1039/c9sc00505f

rsc.li/chemical-science

### Introduction

Finding an efficient, stable and environmentally friendly material system for electrocatalytic reactions has long been a research target pursued by scientists.<sup>1,2</sup> The electrochemical splitting of water offers a promising process for green energy utilization. In this process, the oxygen evolution reaction (OER) half reaction directly determines the overall efficiency of water splitting.<sup>3</sup> Although traditional OER catalysts such as precious metal oxides (e.g.,  $RuO<sub>2</sub>$  and IrO<sub>2</sub>) are used with good catalytic performance, new low-cost catalysts are needed for large-scale implementation.<sup>4</sup>–<sup>7</sup> Of note, cobalt-based inorganic materials have become promising catalysts because of their lower cost, easier synthesis methods, and higher catalytic activity.<sup>8-10</sup> Most

## Tracking the pyrolysis process of a 3-MeOsalophen-ligand based  $Co<sub>2</sub>$  complex for promoted oxygen evolution reaction†

Bingxin Pan,<sup>a</sup> Xu [Pe](http://orcid.org/0000-0002-7471-8467)ng[,](http://orcid.org/0000-0002-8510-5033) D<sup>\*a</sup> Yifan Wang,<sup>a</sup> Qi An,<sup>a</sup> Xu Zhang,<sup>a</sup> Yuexing Zhang, D<sup>a</sup> Thomas S. Teets  $\mathbf{D}^c$  and Min[g](http://orcid.org/0000-0002-7227-7688)-Hua Zeng  $\mathbf{D}^{*ab}$ 

Efficient oxygen evolution reaction catalysts can be prepared via controlled pyrolysis of molecular platforms, and there is still minimal mechanistic understanding of such pyrolysis processes. Here, we introduce a 3-MeOsalophen-ligated cobalt complex as a precursor to obtain a Co-based OER electrocatalyst via controlled pyrolysis under an inert atmosphere. In our case, the unique N, O chelation mode of the 3-MeOsalophen ligand (bis[3-methoxysalicylydene]-1,2 iminophenylenediamine) was used to synthesis a Co<sub>2</sub> complex  $[Co<sub>2</sub>(3-MeOsalophen)(Cl)<sub>3</sub>(CH<sub>3</sub>OH)<sub>2</sub>].$  By regulating the pyrolysis conditions, we successfully obtained a N-doped carbon Co/CoO<sub>x</sub> core–shell nanostructure. More importantly, TG-MS was first adopted for tracking the decomposition products of the complex in the pyrolysis process, further finding out the evolution mechanism from  $Co<sub>2</sub>$  to the core–shell nanostructure. As an electrocatalyst for the oxygen evolution reaction, the core-shell Co/CoO<sub>x</sub>@NC-800 nanostructure achieves an ultralow overpotential of 288 mV at 10 mA  $cm^{-2}$  in 1 M KOH solution. This work offers guiding insight into controlled pyrolysis via TG-MS analysis, using a novel complex precursor for precise regulation of heteroatom-doped (3d) transition metal-based electrocatalysts. EDGE ARTICLE<br>
Tracking the pyrolysis process of<br>  $\bullet$  Cheek forundates<br>
Consist, 2019 in 4660<br>
Class the Cost of B 3-MeOsalophen-ligand based Co<sub>2</sub> complex for<br>
Class the complex scale production reaction in the complex s

precursors for Co-based electrocatalysts are cobalt oxides, cobalt nitrides, cobalt chalcogenides, or Co-based MOFs.<sup>11-17</sup> Despite the fact that this variety of Co-based precursors bring synergic advantages for OER electrocatalysts, there is still impetus to design new precursors to prepare OER catalysts with superior electrochemical performance.

Metal–ligand coordination complexes are attractive as precursors for OER catalysts, with compositions and reactivity that can be precisely controlled.<sup>18,19</sup> The concept is similar to the use of MOF precursors, which via simple pyrolysis can convert into the  $M/MO_x@NC$  core–shell structure with catalytic properties.<sup>8,20,21</sup> The chemical state of nitrogen in the core-shell structure can be adjusted by selecting a suitable nitrogencontaining precursor and regulating the pyrolysis temperature, and there is some benefit of precise regulation of heteroatom-doped (3d) transition metal-based catalysts with a core-shell structure.<sup>22</sup> With these precedents in mind, we turned our attention to Schiff base metal complexes, an important class of compounds in many applications. Among Schiff base derivatives, salen complexes are widely used in catalytic reactions due to the advantages of the N, O coordination mode and ability to stabilize 3d metal ions. These ligands support well-defined coordination complexes and offer the potential to form materials with precise compositions under reductive pyrolysis.<sup>23</sup> However, reports of pyrolysis reactions of salen to prepare electrocatalytic materials are rare, inspiring us to investigate the pyrolysis of a complex supported by a 3-



a Hubei Collaborative Innovation Center for Advanced Organic Chemical Materials, Ministry-of-Education Key Laboratory for the Synthesis and Application of Organic Functional Molecules, College of Chemistry & Chemical Engineering, Hubei University, Wuhan, 430062, P. R. China. E-mail: pengxu@hubu.edu.cn; zmh@ mailbox.gxnu.edu.cn

b Department of Chemistry and Pharmaceutical Sciences, Guangxi Normal University, Key Laboratory for the Chemistry and Molecular Engineering of Medicinal Resources, Guilin, 541004, P. R. China. E-mail: zmh@mailbox.gxnu.edu.cn

c Department of Chemistry, University of Houston, 3585 Cullen Boulevard, Room 112, Houston, Texas 77204-5003, USA

<sup>†</sup> Electronic supplementary information (ESI) available. CCDC 1882936. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c9sc00505f

MeOsalophen ligand to access 3d metal-based nanostructures embedded in N-doped carbon networks.

Here, we introduce a simple 3-MeOsalophen-ligand based Co complex as a precursor to obtain a Co-based electrocatalyst via controlled pyrolysis under an inert atmosphere. In our case, we use the unique N, O coordination chelation mode of the 3-MeOsalophen ligand to synthesis  $Co_2$  [Co<sub>2</sub>(3-MeOsalophen)(Cl)<sub>3</sub>(CH<sub>3</sub>- $[OH]_2$ ]. By regulating the pyrolysis conditions, we successfully obtained an N-doped carbon  $Co/CoO<sub>x</sub>$  core–shell nanostructure. Our work transforms the 3-MeOsalophen ligand outside the Co metal core into N-doped carbon with controlled pyrolysis, and the TG-MS technique is introduced to study the progress of pyrolysis from the  $Co_2$  molecular precursor to N-doped carbon  $Co/CoO_x$ core–shell nanostructures.

### Results and discussion

The typical synthesis procedure of the resulting catalysts from the 3-MeOsalophen ligand to N-doped carbon  $Co/CoO<sub>x</sub>$  core– shell nanostructures is illustrated in Scheme 1. Single-crystal X-ray diffraction analysis reveals that  $[Co<sub>2</sub>(3-MeOsalophen)(Cl)<sub>3</sub>$ - $(CH<sub>3</sub>OH<sub>2</sub>]$  crystallizes in the monoclinic space group  $P<sub>2</sub>/c$  (Table S1<sup>†</sup>). The binuclear complex is bridged by two  $\mu$ 2-phenolate moieties from the salen ligand. The  $Co(m)$  ion in the complex is in a six-coordination mode, in a nearly octahedral geometry. Meanwhile, the  $Co(n)$  ion is in a five-coordination mode with a nearly trigonal bipyramidal geometry. It should be noted that the observation that the ligand  $OCH<sub>3</sub>$  substituents do not participate in coordination has been rarely reported before.  $Co(m)$  is coordinated with the deprotonated phenol oxygens O1 and O2  $(Co(m)$ –  $O1 = 1.905 \text{ Å}, \text{Co(m)}-O2 = 1.890 \text{ Å}, \text{the ligand nitrogen atoms N1}$ and N2 (Co(m)–N1 = 1.871 Å, Co(m)–N2 = 1.883 Å) and anions Cl1 and Cl2 (Co(m)–Cl1 = 2.276 Å, Co(m)–Cl2 = 2.270 Å), where N1, N2, O1, and O2 are coplanar.  $Co(n)$  is coordinated with deprotonated phenol oxygen atoms O1 and O2 (Co( $\text{II}$ )–O1 = 2.231 Å,  $Co(\pi)-O2 = 2.119$  Å), two CH<sub>3</sub>OH molecules and the anion Cl3  $(Co(\Pi)-Cl3 = 2.330 \text{ Å})$  (Fig. S3 and Table S2<sup>†</sup>). In addition, two N and two O atoms from the 3-MeOsalophen ligand complete the peripheral ligation around the metallic Co core. In the context of pyrolysis (see below), we propose that the (N, N, O, O)

coordination mode of the salen prevents cobalt leaching at low temperature and provides control over the pyrolysis process. In addition, we anticipate that the aromatic core of the ligand and the ligand  $C=N$  bonds provide the template for the N-doped carbon that surrounds the cobalt following pyrolysis. Based on these features, we initiated TG-MS studies to understand the decomposition and recombination mechanism of the 3-MeOsalophen ligand-based complexes during pyrolysis, with the goal of preparing highly active OER electrocatalysts.

The as-synthesized  $Co<sub>2</sub>$  was pyrolysed under Ar-annealing conditions to prepare the hybrid porous  $Co/CoO_x@NC-T$  (T represents the pyrolysis temperature). Of note, pyrolysis temperature plays a key role in determining the structures of the resulting samples, thus leading to different electrocatalytic activities, so we analyzed the sample composition obtained at different pyrolysis temperatures by powder X-ray diffraction (PXRD) and high-resolution transmission electron microscopy (HRTEM) (Fig. 1). The X-ray diffraction (XRD) patterns of the Co/  $CoO<sub>x</sub>(a)NCs$  are shown in Fig. 1a. It can be found that  $Co/$  $CoO<sub>x</sub>(QNC-500 exists only in the form of elemental cobalt, and$ the peaks at 44.2 $^{\circ}$ , 51.6 $^{\circ}$  and 75.9 $^{\circ}$  can be assigned to the (111), (200) and (220) facets of structured metallic cubic Co (JCPDS 15- 0806). Meanwhile, the peaks at  $41.5^{\circ}$  and  $47.4^{\circ}$  belong to the (100) and (101) facets of hexagonal Co (JCPDS 05-0727). When the pyrolysis temperature was increased from 500  $^{\circ}$ C to 600  $^{\circ}$ C, the diffraction peak of CoO first appeared, corresponding to the diffraction peak at 36.5 $^{\circ}$ , 42.4 $^{\circ}$  and 61.5 $^{\circ}$ . It should be noted that the peak of hexagonal Co disappears, and this fact suggests the transformation of small particles of hexagonal cobalt into cubic cobalt, and the agglomeration grows into cubic cobalt due to the increase in temperature. Furthermore, the peaks at  $31.3^\circ$ , 59.4 $^{\circ}$  and 65.2 $^{\circ}$  can be assigned to Co<sub>3</sub>O<sub>4</sub> when the temperature increases further to 700  $^{\circ}$ C.<sup>24,25</sup> Interestingly, when the pyrolysis temperature increases to 800 $^{\circ}$ C, we hardly find the diffraction peaks of  $Co<sub>3</sub>O<sub>4</sub>$ , which may be due to the reduction gases produced during the pyrolysis process, which would be systematically analyzed as described in Fig. 3. Of note, the stronger diffraction peak of  $Co<sub>3</sub>O<sub>4</sub>$  appears again in Co/  $CoO<sub>x</sub>@NC-900$ . The specific phase changes and facet information of  $Co/CoO<sub>x</sub>@NCs$  are summarized in Table S3.<sup>†</sup> In Edge Article<br>
McGolalophan ligand to access 3d meal-based nanostractures coordination mode of the sales proside the recupies are a present<br>
emplete in Super Scheme and Details (a Napole discrepand based Commercial 3.0 Napo



Scheme 1 The schematic illustration of the synthesis strategy from the 3-MeOsalophen ligand to Co-based complex and the finally obtained controlled pyrolysis Co/CoO<sub>x</sub>@NC nanostructures



Fig. 1 Structural information of the  $Co/CoO<sub>x</sub>$  and  $Co}$  nanostructures. (a) Powder XRD patterns of  $Co/CoO_x@NCs$  synthesized at 500-900 °C; (b) HRTEM and (c) HAADF images of Co/CoO<sub>x</sub>@NC-800; (d) elemental mapping images of Co/CoO<sub>x</sub>@NC-800.

addition, carbon peaks were negligible in all the XRD patterns of  $Co/CoO<sub>x</sub>(aNCs, presumably because of their relatively weak$ intensities. To further confirm its phase composition, the HRTEM images of  $Co/CoO<sub>x</sub>(QNC-800)$  were obtained and are shown in Fig. 1b, demonstrating that the nanoparticle is encapsulated in a carbon layer, where the Co (111) crystal plane and CoO (111) are clearly seen with a lattice spacing of 0.204 nm and 0.241 nm, respectively, consistent with the XRD results of  $Co/CoO<sub>x</sub>(QNC-800)$ . Meanwhile, the 0.345 nm spacing on the carbon layer can be attributed to the (002) plane of the graphitic carbon. The carbon layer provides good protection for the nanoparticles and improves the catalytic stability.<sup>26-28</sup> Furthermore, high-angle annular dark field scanning transmission electron microscopy (HAADF-STEM) imaging and elemental mapping were also conducted, in which we can find that metallic cobalt is located in the center of the core–shell structure, while C and N elements are uniformly distributed around, as shown in Fig. 1c and  $d^{29,30}$  In addition, the distribution of Co and O indicates that Co atoms may be bonded with O to form amorphous  $CoO<sub>r</sub>$  parts which are favorable for electrocatalytic activity.<sup>31</sup> Therefore, the aforementioned results clearly demonstrate that the  $Co/CoO<sub>x</sub>@NC-800$  sample was successfully prepared. The TEM images of other  $Co/CoO<sub>x</sub>(a)NCs$  are shown in Fig. S4,† and the particle size increases with increasing the pyrolysis temperature (600–900  $^{\circ}$ C).

To further explore the chemical bonding environment of Co/ CoOx@NC nanostructures, X-ray photoelectron spectroscopy (XPS) is effective in determining the chemical bonding properties of N, C, O and Co species in catalysts, as shown in Fig. 2a–d. The specific element details are listed in Table S4,<sup>†</sup> and XPS data for other  $Co/CoO<sub>x</sub>@NCs$  are shown in Fig. S5, S6, S8 and S9.† The XPS survey spectrum in Fig. 2a shows main peaks at 284.6, 398.4, 531.1, and 777.9 eV, which could be assigned to C 1s, N 1s, O 1s, and Co 2p, respectively.<sup>30</sup> The deconvoluted highresolution C 1s spectrum exhibited three bands, and the bands were related to the  $sp^2 C$  (284.6 eV), C–N or C–O (285.6 eV), and  $C=N$  or  $C=O(287.5 \text{ eV})$ , respectively. We observed asymmetric signals at 284.3 eV and peaks related to C–N and C–O, which



Fig. 2 The chemical bonding environment of Co/CoO<sub>x</sub>@NC nanostructures. (a) XPS survey spectra. (b) XPS survey spectrum of Co/CoO<sub>x</sub>@NC-800 for Co 2p. (c) XPS analysis-based Co<sup>2+</sup>/Co<sup>3+</sup> atomic ratios of Co/CoO<sub>x</sub>@NCs. (d) XPS survey spectrum of Co/CoO<sub>x</sub>@NC-800 for N 1s. (e) Raman spectroscopy of Co/CoO<sub>x</sub>@NCs. (f) Brunauer–Emmett–Teller (BET) values of Co/CoO<sub>x</sub>@NCs.

confirmed the successful bonding of N and O to the carbon structure. When the pyrolysis temperature was increased, there was no significant change in the percentages of the three bands of the material. This suggests that the composition of the surrounding carbon is basically stable (Fig. S5, S7 and Table S5<sup>†</sup>).<sup>32</sup> Fig. 2b shows the Co 2p spectra of Co@CoO<sub>x</sub>/NC-800, which indicate the simultaneous presence of  $Co<sup>0</sup>$  (778.6 eV),  $Co<sup>2+</sup>$  (782.0 eV) and  $Co<sup>3+</sup>$  (780.1 eV), which has a synergistic effect on the electrocatalytic processes.<sup>33,34</sup> Furthermore, we find that  $Co^{2+}$  is dominant compared with  $Co^{3+}$  in  $Co/CoO<sub>x</sub>(a)NC-800$ (Fig. 2c), which may be due to the reductive gases produced during the pyrolysis process, and this is consistent with the data in Fig. 3 and  $S8.$ <sup>†</sup> In addition, the O 1s spectra also confirmed the coexistence of oxygen defects (531.6 eV), C–O (533.5 eV) and surface oxidation of the metallic cobalt at 529.8 eV, respectively (Fig. S6†), indicating that there are abundant O vacancies in the  $Co/CoO<sub>x</sub>@NC-800$  nanostructures.<sup>35,36</sup> When the pyrolysis temperature reaches 700  $^{\circ}$ C and 800  $^{\circ}$ C, the percentage of Co–O in the materials significantly decreased (Fig. S7 and Table S6<sup>†</sup>). Meanwhile, the N 1s spectrum of  $Co/CoO<sub>r</sub>(QNC-800$  was also collected (Fig. 2d). Four main peaks associated with pyridinic-N (398.5 eV), pyrrolic-N (400.0 eV), Co–N<sub>x</sub> (399.1 eV) and graphitic-N (401.3 eV) species were observed (Fig. S9 and Table S7†).<sup>32</sup> The Co content in  $Co/CoO_x@NC-600$ ,  $Co/CoO_x@NC-700$ ,  $Co/CoO<sub>x</sub>@NC-800$  and  $Co/CoO<sub>x</sub>@NC-900$  was determined by Edge Article<br>
confirmed the successful bonding of N and O to the carbon inductively coupled plasma optical constants are<br>
of the martial. This suggests that the composition of the Corologist core shall ansostrate<br>the "int

inductively coupled plasma optical emission spectroscopy and found to be 35.75, 50.42, 40.49 and 68.42 wt%, respectively (Table S8†). These results demonstrate the formation of  $Co/CoO<sub>x</sub>(a)NC$  core–shell nanostructures.<sup>25</sup>

Raman spectroscopy was further used to characterize the degree of graphitization of the  $Co/CoO<sub>x</sub>(aNCs)$  (Fig. 2e). In the spectra, two sharp Raman peaks were observed at 1354 and  $1584\ \mathrm{cm}^{-1}$  which are characteristic of carbon materials and can be ascribed to the disordered structures/structural defects (or D band) and the graphitic G band, respectively. The intensity ratio of the D band to G band  $(I_D/I_G)$  increases from 0.75 to 1.11 for  $Co/CoO<sub>r</sub>(QNC-600-900,$  indicating that the resulting defects increase at higher pyrolysis temperature. In addition, peaks at 470.6, 515.6 and 680.9  $cm^{-1}$  are classical vibration modes of  $Co-O$ <sup>37,38</sup> Furthermore, the surface area along with pore size distribution of the  $Co/CoO<sub>x</sub>@NCs$  was evaluated. As shown in Fig. 2f, it was found that the BET surface area was 321.5  $m^2$  g<sup>-1</sup> for Co/CoO<sub>x</sub>@NC-600, 223.9 m<sup>2</sup> g<sup>-1</sup> for Co/CoO<sub>x</sub>@NC-700, 312.7 m<sup>2</sup> g<sup>-1</sup> for Co/CoO<sub>x</sub>@NC-800 and 102.319 m<sup>2</sup> g<sup>-1</sup> for  $Co/CoO<sub>x</sub>(QNC-900$ . Usually, the BET surface areas of complexes that undergo the pyrolysis process would gradually decrease as pyrolysis temperature increases,<sup>39</sup> yet the  $Co/CoO<sub>x</sub>@NC-800$ sample is nearly similar to  $Co/CoO<sub>x</sub>@NC-600$ , which may be due to the reduction gases produced during the pyrolysis process, as described further in Fig. 3. Meanwhile, the presence



Fig. 3 Controlled pyrolysis tracking analysis. (a) TG and DTG profiles of Co<sub>2</sub>. (b) TG-MS curves of the corresponding Co<sub>2</sub> pyrolysis tracking process. (c) Possible decomposition and polymerization reaction occurring in the pyrolysis process from Co<sub>2</sub> to Co/CoO<sub>x</sub>@NCs.

#### Chemical Science Edge Article

of mesopores and the pore size distribution determined by the QSDFT method are displayed in Fig. S10 and S11,† respectively. Therefore, the chemical bonding environment of Co, O and C was systematically analyzed by XPS, Raman spectroscopy and BET surface area measurement, showing that the  $Co/CoO<sub>x</sub>(a)$ -NC-800 sample has higher  $Co^{2+}/Co^{3+}$  and a reasonable  $I_D/I_G$ ratio as well as a relatively higher surface area compared with other  $Co/CoO<sub>x</sub>(aNCs)$ .

It is essential to trace the decomposition products in the pyrolysis process, and further to find out the mechanism of the pyrolysis process and the evolution of the core–shell structure. Hence, thermogravimetric analysis combined with mass spectrometry (TG-MS) was used to track the pyrolysis process. As shown in Fig. 3a, the TG and DTG curves of  $Co<sub>2</sub>$  can be divided into three areas from  $30-800$  °C. Generally, the mass decrease between 30 °C and 100 °C is attributed to the loss of CH<sub>3</sub>OH or H2O guest molecules (namely region A); moreover, the decrease between 150–350  $\degree$ C is largely due to the partial decomposition of the ligand (namely region B); and finally between 480  $\degree$ C and 800 °C we observe complete decomposition of  $Co<sub>2</sub>$  to form the core-shell structure of Co/CoO<sub>x</sub>@NC (namely region C).<sup>40-42</sup>

According to the structure of  $Co<sub>2</sub>$  and the fragments detected by TG-MS as shown in Fig. 3b, the whole pyrolysis process can be divided into several processes: (i) the process from room temperature to 100  $^{\circ}$ C only detects fragments of CH<sub>3</sub>OH, where the escape of guest molecules occurs. As the temperature increases from 100 °C to 200 °C, CH<sub>3</sub>OH is detected again, assigned to the  $CH<sub>3</sub>OH$  molecules coordinated to Co which escape the coordination sphere of the  $Co<sub>2</sub>$  complex. (ii) When the temperature is raised from 150  $\degree$ C to 300  $\degree$ C, two peaks appear in the temperature region as seen from the DTG curve. In this temperature range, several fragments are detected by TG-MS. Characteristic fragments include CH<sub>4</sub> ( $m/z = 15$ ), Cl' ( $m/z =$ 38), and CH<sub>3</sub>Cl ( $m/z = 52$ ). Of note, CH<sub>4</sub> can be attributed to the ligand centered ether bond cleavage. In addition, the chlorine atom on Co can be combined with a methyl group on a similar ligand to form  $CH<sub>3</sub>Cl$ . And the generation of  $CH<sub>3</sub>Cl$  is further increased when the temperature is increased. It may be related to the increase of  $CH_4$  and the escape of more chlorine atoms, and the extra Cl atoms escape as free radicals  $(m/z = 38)$ . The loss of coordinated methanol and chloride ligands will lead to Co becoming an unusually active center, which may facilitate the aggregation of cobalt atoms and the association with the partially decomposed fragments to form a stable core–shell nanostructure. (iii) Further increase in temperature leads to complete rupture of the whole  $Co_2$  complex, resulting in  $C_6H_6$  $(m/z = 78)$  fragment formation and ether C–O, aromatic C–C, and C–N bond cleavage processes, producing an active site that promotes ligand polymerization and release of  $H_2$ .<sup>43</sup> On the other hand,  $HC = N'$  formed by the simultaneous cleavage of the Co–N coordination bond and  $C_6H_6$  in this process participates in the formation of a carbon network, and finally forms N-doped carbon. Prominently, while the ligand undergoes decomposition and polymerization, the metal oxide formed by Co and phenol oxygen is also reduced to some extent by adjacent small molecule fragments such as  $H_2$  and CH<sub>4</sub> to form metallic Co with  $H_2O$  and CO escaping. Since  $H_2$  is generated in this

process, no  $CO_2$  ( $m/z = 44$ ) is detected. Similarly, we tracked the pyrolysis process of the 3-MeOsalophen ligand (Fig. S12 and S13†). It is found that the decomposition of the ligand produces more fragments than that of  $Co<sub>2</sub>$ , and a little sample remained after pyrolysis  $(2.32\%)$ , suggesting that the Co active site produced by the decomposition of  $Co<sub>2</sub>$  may accelerate the recombination of organic fragments, and then rapidly forms a stable carbon network, which is consistent with the aforementioned phenomena. In addition,  $H_2$  was not detected in the process of ligand decomposition, suggesting that the Co active site is crucial for the production of  $H<sub>2</sub>$ . To summarize, the whole pyrolysis process from  $Co_2$  to  $Co/CoO_x@NC$  nanostructures is depicted in Fig. 3c, indicating that the pyrolysis process of  $Co<sub>2</sub>$ occurs in a reductive atmosphere between  $520-700$  °C, accounting for the distinct nanostructure of  $Co/CoO<sub>x</sub>(QNC-800)$ formed via this process.

The catalytic ability of  $Co/CoO<sub>x</sub>(QNCs)$  for the oxygen evolution reaction was also assessed in 1 M KOH solution using a typical three-electrode configuration. The loading of catalyst on carbon paper is 1.68 mg  $cm^{-2}$ . For comparison, commercial  $RuO<sub>2</sub>$  was also examined under the same conditions. The Co/  $CoO<sub>x</sub>@NCs$  and commercial  $RuO<sub>2</sub>$  electrodes were first oxidized in alkaline medium saturated with  $O<sub>2</sub>$ , and then measured at 5 mV  $s^{-1}$  to obtain the polarization curves shown in Fig. 4a. The polarization curves obtained for  $Co/CoO_x@NC-800$  show that the material has signicant electrocatalytic properties for the OER. The  $Co/CoO_x@NC-800$  electrode displayed the smallest overpotential of 288 mV at 10 mA  $cm^{-2}$ . In contrast, Co/  $CoO_x@NC-600$ ,  $Co/CoO_x@NC-700$ ,  $Co/CoO_x@NC-900$  and commercial RuO<sub>2</sub> possessed larger overpotentials of 433 mV, 356 mV, 413 mV and 339 mV, respectively. Meanwhile, the Tafel slope for Co/CoO<sub>x</sub>@NC-800 is 98 mV dec<sup>-1</sup> (Fig. 4b). The results indicated that  $Co/CoO<sub>x</sub>(QNC-800)$  exhibits excellent electrocatalytic performance, which is comparable to that of the stateof-the-art Co-based electrocatalysts (Table S9†). To gain further insight into the electrocatalytic activity of the above electrocatalysts, the electrochemical impedance spectra (EIS) and the  $C_{d1}$  (double layer capacitance) of the electrocatalysts were obtained.<sup>36,44</sup> As shown in Fig. 4d, the  $C_{dl}$  value of the Co/ CoO<sub>x</sub>@NC-800 electrode is calculated to be 35.04 mF cm<sup>-2</sup>, which is larger than the 31.72, 26.55, 17.2 mF  $cm^{-2}$  of Co/  $CoO_x@NC-700$ ,  $Co/CoO_x@NC-900$  and  $Co/CoO_x@NC-600$ , respectively (Fig. S14†). This suggests that  $Co/CoO<sub>x</sub>@NC-800$ has more catalytically active centers in the electrolysis process. In addition, electrochemical impedance spectroscopy (EIS) was conducted to explore the electrode kinetics under OER conditions. In the Nyquist plots (Fig. 4e) of  $Co/CoO_x@NCs$ ,  $Co/$  $CoO<sub>x</sub>(a)NC-800$  could be clearly seen to have the smallest semicircle (with the smallest radius) or the smallest charge transfer resistance ( $R_{\text{ct}}$ , 7.40  $\Omega$ ) compared with the other electrocatalysts ( $R_{\text{ct}} = 13.61 \Omega$  for Co/CoO<sub>x</sub>@NC-600,  $R_{\text{ct}} = 11.09 \Omega$ for Co/CoO<sub>x</sub>@NC-700 and  $R_{ct} = 12.79 \Omega$  for Co/CoO<sub>x</sub>@NC-900). The measured impedance spectra data were fitted and evaluated using Zview software. As shown in Fig. 4e, the symbols refer to the experimental data and the lines denote the corresponding fitting results. The inset is the equivalent circuit in the form of  $R_s$  (CPE1,  $R_o$ ) (CPE2,  $R_{ct}$ ), where  $R_s$  represents the Openical Selence<br>
of mesopotes and the ports are distribution decembered by the concerns of the NoCologite on 12<br>
Operating articles. This are the system and the component than the of Co-band all interactions<br>
was systema



Fig. 4 OER activity of Co/CoO<sub>x</sub>@NC nanostructures and commercial RuO2. (a) IR-corrected polarization LSV curves. (b) Corresponding Tafel plots. (c) Comparison of overpotentials at  $j = 10$  mA cm<sup>-2</sup> corresponding to Tafel slopes. (d) Current density as a function of the scan rates for Co/CoO<sub>x</sub>@NCs. (e) Nyquist plots of catalysts at the biased potential of 1.6 V vs. RHE over the frequency range from 10 000 to 0.01 Hz. (f) Chronopotentiometric measurements of long-term stability of  $Co/CoO<sub>x</sub>@NC-800$ . The inset shows the TEM image after the OER chronopotentiometric test, scale bar is 100 nm.

solution resistance, CPE1 and CPE2 are the constant phase elements which describe electron transport at the GCE/catalyst interface and the catalyst/electrolyte interface, respectively,  $R_0$  is the oxide film resistance and  $R_{ct}$  denotes the charge transfer resistance at the catalyst/electrolyte interface.<sup>45</sup> This suggested that the electron transfer rate during the OER was faster in Co/  $CoO<sub>x</sub>@NC-800$ . The results of electrochemical impedance spectroscopy (EIS) and electrochemical double-layer capacitance  $(C_{\text{dl}})$  measurements of electrocatalysts confirm the excellent OER performance of  $Co/CoO<sub>x</sub>(QNC-800$  among Co/  $CoO<sub>x</sub>@NCs$ . More importantly, high durability is of great significance for electrocatalysts in practical applications. Chronopotentiometry (CP) was conducted to study the electrocatalytic durability of  $Co/CoO_x@NC-800$  for the OER in  $O_2$ saturated 1.0 M KOH solutions. As shown in Fig. 4f, Co/  $CoO<sub>x</sub>@NC-800$  shows negligible potential fluctuation at 50 mA  $\text{cm}^{-2}$  for 15 h. Furthermore, the phase and the corresponding peaks of XRD patterns of the  $Co/CoO<sub>x</sub>@NC-800$  sample before and after the catalysis were compared (Fig.  $S15\dagger$ ), showing that the main phase peak of the sample is well preserved. At the same time, as confirmed by the TEM image, the core-shell nanostructure morphology is retained after long-term OER operation (inset of Fig. 4f). The above indicates that the Co/  $CoO_x@NC-800$  structure from  $Co_2$  has excellent stability for the OER in alkaline media. Besides, the influence of pyrolysis

temperature on final catalytic performance is discussed in Fig. S4.† In other words, the excellent electrochemical performance of  $Co/CoO<sub>x</sub>@NC-800$  is attributed to the synergic effects of the designable 3-MeOsalophen-based Co complex as the precursor and controlled pyrolysis conditions, forming such a distinct nanostructure.

### Conclusions

In summary, we have shown that the 3-MeOsalophen-ligand based  $Co<sub>2</sub>$  complex undergoes controlled pyrolysis to form distinct core–shell  $Co/CoO_x@NC$  nanostructures which are efficient OER electrocatalysts. TG-MS analysis reveals the pyrolysis process and indicates that the decomposition of the 3- MeOsalophen-ligated  $Co<sub>2</sub>$  provides a reductive atmosphere, and the possible decomposition and polymerization mechanism of Co2 was elucidated, which facilitates the formation of core–shell nanostructures. Notably, the core–shell  $Co/CoO<sub>x</sub>(QNC-800)$ nanostructure achieves an ultralow overpotential of 288 mV at 10 mA  $cm^{-2}$  in 1 M KOH solution, which is comparable to that of the state-of-the-art Co-based electrocatalysts. This work offers a general route for designing synthetically derived core–shell nanostructures by designing selected complexes and controlling pyrolysis conditions.

### Conflicts of interest

There are no conflicts to declare.

### Acknowledgements

This work was financially supported by the National Science Fund for Distinguished Young Scholars (Grant 21525101), the BAGUI Talent and Scholar Program (2014A001), NSFC (Grants 81773775, 21805074 and 21661088), NSFC of Hubei (Grants 2017CFA006, 2018CFB151), and NSFGX (Grants 2014GXNSFFA118003, 2017GXNSFDA198040).

### Notes and references

- 1 B. Zhang, X. Zheng, O. Voznyy, R. Comin, M. Bajdich, M. García-Melchor, L. Han, J. Xu, M. Liu, L. Zheng, E. Yassitepe, N. Chen, T. Regier, P. Liu, Y. Li, P. D. Luna, A. Janmohamed, H. L. Xin, H. Yang, A. Vojvodic and E. H. Sargent, Science, 2016, 352, 333–337.
- 2 S. L. Zhao, Y. Wang, J. C. Dong, C. T. He, H. J. Yin, P. F. An, K. Zhao, X. F. Zhang, C. Gao, L. J. Zhang, J. W. Lv, J. X. Wang, J. Q. Zhang, A. M. Khattak, N. A. Khan, Z. X. Wei, J. Zhang, S. Q. Liu, H. J. Zhao and Z. Y. Tang, Nat. Energy, 2016, 1, 16184.
- 3 J. Suntivich, K. J. May, H. A. Gasteiger, J. B. Goodenough and Y. Shao-Horn, Science, 2011, 334, 1383–1385.
- 4 X. Kong, K. Xu, C. Zhang, J. Dai, S. Norooz Oliaee, L. Li, X. Zeng, C. Wu and Z. Peng, ACS Catal., 2016, 6, 1487–1492.
- 5 C. Hu and L. Dai, Angew. Chem., Int. Ed., 2016, 55, 11736– 11739.
- 6 C. McCrory, S. Jung, J. Peters and T. Jaramillo, J. Am. Chem. Soc., 2013, 135, 16977–16987.
- 7 E. Hu, Y. Feng, J. Nai, D. Zhao, Y. Hu and X. W. Lou, Energy Environ. Sci., 2018, 11, 872–880.
- 8 S. Dou, C. L. Dong, Z. Hu, Y. C. Huang, J. l. Chen, L. Tao, D. Yan, D. Chen, S. Shen, S. Chou and S. Wang, Adv. Funct. Mater., 2017, 27, 1702546.
- 9 K. Yan, J. Chi, J. Xie, B. Dong, Z. Liu, W. Gao, J. Lin, Y. Chai and C. Liu, Renewable Energy, 2018, 119, 54–61.
- 10 K. Yan, X. Shang, Z. Li, B. Dong, X. Li, W. Gao, J. Chi, Y. Chai and C. Liu, Appl. Surf. Sci., 2017, 416, 371–378.
- 11 Y. Tong, P. Chen, T. Zhou, K. Xu, W. Chu, C. Wu and Y. Xie, Angew. Chem., Int. Ed., 2017, 56, 7121–7125.
- 12 P. Chen, K. Xu, Z. Fang, Y. Tong, J. Wu, X. Lu, X. Peng, H. Ding, C. Wu and Y. Xie, Angew. Chem., Int. Ed., 2015, 54, 14710–14714.
- 13 M. Zhang, Q. Dai, H. Zheng, M. Chen and L. Dai, Adv. Mater., 2018, 30, 1705431.
- 14 Y. Lu, L. Yu, M. Wu, Y. Wang and X. W. Lou, Adv. Mater., 2018, 30, 1702875.
- 15 X. Wang, L. Dong, M. Qiao, Y. Tang, J. Liu, Y. Li, S. Li, J. Su and Y. Lan, Angew. Chem., Int. Ed., 2018, 57, 9660–9664.
- 16 K. Yan, J. Chi, Z. Liu, B. Dong, S. Lu, X. Shang, W. Gao, J. Chi, Y. Chai and C. Liu, Inorg. Chem. Front., 2017, 4, 1783–1790.
- 17 K. Yan, J. Qin, J. Lin, B. Dong, J. Chi, Z. Li, F. Dai, Y. Chai and C. Liu, J. Mater. Chem. A, 2018, 6, 5678–5686.
- 18 M. H. Zeng, Z. Yin, Z. H. Liu, H. B. Xu, Y. C. Feng, Y. Q. Hu, L. X. Chang, Y. X. Zhang, J. Huang and M. Kurmoo, Angew. Chem., Int. Ed., 2016, 55, 11407–11411.
- 19 J. Wang, L. Gan, W. Zhang, Y. Peng, H. Yu, Q. Yan, X. Xia and X. Wang, Sci. Adv., 2018, 4, eaap7970.
- 20 L. Jiao and H. L. Jiang, Chem, 2019, DOI: 10.1016/ j.chempr.2018.12.011.
- 21 H. Zhang, J. Nai, L. Yu and X. W. Lou, Joule, 2017, 1, 77–107.
- 22 P. Yin, T. Yao, Y. Wu, L. Zheng, Y. Lin, W. Liu, H. Ju, J. Zhu, X. Hong, Z. Deng, G. Zhou, S. Wei and Y. Li, Angew. Chem., Int. Ed., 2016, 55, 10800–10805.
- 23 C. Baleizao and H. Garcia, Chem. Rev., 2006, 106, 3987–4043.
- 24 H. Jin, J. Wang, D. Su, Z. Wei, Z. Pang and Y. Wang, J. Am. Chem. Soc., 2015, 137, 2688–2694.
- 25 A. Aijaz, J. Masa, C. Rçsler, W. Xia, P. Weide, A. J. R. Botz, R. A. Fischer, W. Schuhmann and M. Muhler, Angew. Chem., Int. Ed., 2016, 55, 4087–4091.
- 26 G. Xu, G. C. Xu, J. J. Ban, L. Zhang, H. Lin, C. L. Qi, Z. P. Sun and D. Z. Jia, J. Colloid Interface Sci., 2018, 521, 141–149.
- 27 Y. Z. Chen, C. Wang, Z. Yu. Wu, Y. Xiong, Q. Xu, S. H. Yu and H. L. Jiang, Adv. Mater., 2015, 27, 5010–5016.
- 28 X. Zhang, R. Liu, Y. Zang, G. Liu, G. Wang, Y. Zhang, H. Zhang and H. Zhao, Chem. Commun., 2016, 52, 5946– 5949.
- 29 S. Li, C. Cheng, X. Zhao, J. Schmidt and A. Thomas, Angew. Chem., Int. Ed., 2018, 57, 1856–1862.
- 30 K. Xu, H. Cheng, L. Liu, H. Lv, X. Wu, C. Wu and Y. Xie, Nano Lett., 2016, 17, 578–583.
- 31 S. Huang, Y. Meng, S. He, A. Goswami, Q. Wu, J. Li, S. Tong, T. Asefa and M. Wu, Adv. Funct. Mater., 2017, 27, 1606585.
- 32 J. Zhang, L. Qu, G. Shi, J. Liu, J. Chen and L. Dai, Angew. Chem., Int. Ed., 2016, 128, 2270–2274.
- 33 X. Yan, Y. Jia, J. Chen, Z. Zhu and X. Yao, Adv. Mater., 2016, 28, 8771–8778.
- 34 B. Chen, X. He, F. Yin, H. Wang, D. J. Liu, R. Shi, J. Chen and H. Yin, Adv. Funct. Mater., 2017, 27, 1700795.
- 35 J. Bao, X. Zhang, B. Fan, J. Zhang, M. Zhou, W. Yang, X. Hu, H. Wang, B. Pan and Y. Xie, Angew. Chem., Int. Ed., 2015, 54, 7399–7404. Openical Science<br>  $\frac{1}{2}$  Creative Articles. Published on 12 March 2019. The Access Article is licensed under a Creative Common Access Articles. Published and the Creative Common Access Articles. Download Unported Unpor
	- 36 H. Xu, Z. X. Shi, Y. X. Tong and G. R. Li, Adv. Mater., 2018, 30, 1705442.
	- 37 S. H. Johnson, C. L. Johnson, S. J. May, S. Hirsch, M. W. Cole and J. E. Spanier, J. Mater. Chem., 2010, 20, 439–443.
	- 38 B. S. Yeo and A. T. Bell, J. Am. Chem. Soc., 2011, 133, 5587– 5593.
	- 39 Y. Ding, Y. Niu, J. Yang, L. Ma, J. Liu, Y. Xiong and H. Xu, Small, 2016, 12, 5414–5421.
	- 40 R. A. Schäfer, D. Dasler, U. Mundloch, F. Hauke and A. Hirsch, J. Am. Chem. Soc., 2016, 138, 1647–1652.
	- 41 R. Risoluti, M. A. Fabiano, G. Gullifa, L. W. Wo and S. Materazzi, Russ. J. Gen. Chem., 2017, 87, 564–568.
	- 42 T. K. Kim, K. J. Lee, J. Y. Cheon, J. H. Lee, S. H. Joo and H. R. Moon, J. Am. Chem. Soc., 2013, 135, 8940–8946.
	- 43 L. Zhang and Y. H. Hu, J. Phys. Chem. C, 2010, 114, 2566– 2572.
	- 44 T. Tang, W. J. Jiang, S. Niu, N. Liu, H. Luo, Y. Y. Chen, S. F. Jin, F. Gao, L. J. Wan and J. S. Hu, J. Am. Chem. Soc., 2017, 139, 8320–8328.
	- 45 J. Zhang, F. Li, W. Chen, C. Wang and D. Cai, Electrochim. Acta, 2019, 300, 123–130.