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A MOF-derived Co₃O₄/nitrogen-doped carbon composite for chlorine-assisted production of ethylene oxide†

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Ethylene oxide (EO) is one of the most crucial materials in plastic industries. The traditional catalytic process requires high temperature and pressure to produce EO. A chlorine-assisted system has been reported to produce EO, but it required noble metal catalysts, which significantly increased the cost. In this work, a MOF-derived $Co_3O_4/nitrogen$ -doped carbon composite (Co_3O_4/nC) prepared through a two-step calcination method exhibited remarkable chlorine evolution reaction (ClER) activity as compared with a commercial RuO_2 catalyst, which can be attributed to the higher specific surface area and lower resistance of its porous structure and nitrogen-doped carbon. Furthermore, the Co_3O_4/nC maintained a stable potential and a high faradaic efficiency throughout the 10-hour electrolysis test.

Introduction

Ethylene oxide (EO), one of the most crucial chemicals for manufacturing consumer products and intermediates, was mass-produced with an annual production of over 34.5 Mt (2021). Taking great advantage of its strained ring structure, EO can participate in various addition reactions to open the ring. One major use of EO is its conversion to ethylene glycol, which is one of the most important precursors in the plastic industry. In addition, EO can be used to produce ethanol amines and various polymers with novel architectures. Patented by Lefort in 1935 and numerous follow-up studies, ethylene can be directly oxidized to EO on silverbased catalysts through thermal catalytic processes. However, relatively high temperature (220–280 °C) and high pressure (1–3 MPa) are required for this chemical transformation.

Alternatively, EO can be produced by a chlorohydrin process (RXN 2, 3 and 5). Powered by renewable electricity generated from solar and wind sources, chlorine (Cl_2) can be *in situ* generated through an electrochemical process under ambient conditions (RXN 1). Then, EO can be produced from a chlorine-assisted process in the formation of ethylene chlorohydrin (RXN 2 and 3), followed by its chemical reaction with

$$2Cl^- + 2e^- \rightarrow Cl_2 \tag{1}$$

$$Cl_2 + H_2O \rightarrow HCl + HClO$$
 (2)

$$HClO + C_2H_4 \rightarrow Cl-C_2H_4-OH$$
 (3)

$$2H_2O-2e^- \rightarrow H_2 + 2OH^- \tag{4}$$

$$Cl-C_2H_4-OH+OH^- \to C_2H_4O+Cl^- + H_2O$$
 (5)

In this work, a metal-organic framework (MOF)-derived porous cobalt oxide/nitrogen-doped carbon composite (Co_3O_4/NC) was studied for chlorine-assisted EO production. MOF-derived materials are well known for their unique porous structure that can offer a high specific area that is desirable for various

OH-, which gets accumulated at the cathode (RXN 4) to produce ethylene oxide via a dehydrochlorination process (RXN 5) (Fig. 1a). In the meantime, hydrogen is generated on the cathode.11 Iridium oxide (IrO2)12 and barium oxide-loaded IrO₂ 13 were reported to achieve EO production with a high faradaic efficiency (FE) of >70%. Besides, a dimensionally stable anode (DSA), made of ruthenium oxide and titanium oxide, has been proven to be an active and durable catalyst for the Cl₂ evolution reaction (CIER) and is expected to be active for the chlorine-assisted EO generation process.14 However, all these catalysts are based on noble metals, which significantly increases the overall process costs. Exploration of low-cost, robust, non-noble metal catalysts is highly desirable. Although several works showed that non-noble cobalt oxide (Co₃O₄) is an active and durable catalyst in the ClER process, 15-17 its performance is generally inferior to those of noble metal-based catalysts.

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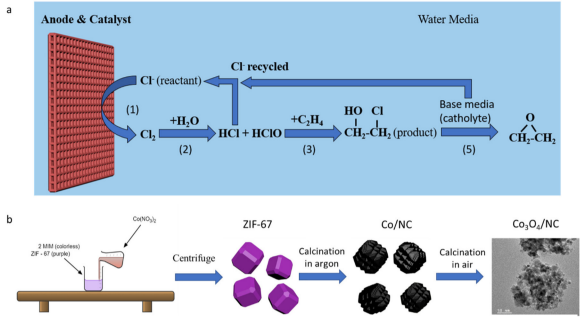


Fig. 1 (a) Scheme of the chlorine-assisted EO generation process. (b) Synthesis process.

applications, such as sensing, energy storage and catalysis. 18-20 However, they have been rarely studied for the chlorine evolution reaction (CIER). Benefiting from the highly porous structure and carbonized organic linker of MOFs as a sacrificing template, Co₃O₄ catalysts with a high specific surface area and low charge transfer resistance were prepared. Compared to the generally used direct calcination in air to turn MOFs into metal oxides, we investigated a two-step calcination method and observed that the calcination method and temperature significantly influenced the structure and ClER activity. It was seen that the optimized catalyst exhibits comparable ClER catalytic activities to the commercial ruthenium oxide (RuO2) catalyst, while Co₃O₄ has a cost of a two orders of magnitude lower than RuO₂. Finally, the Co₃O₄/NC exhibited long-term durability for chlorine-assisted EO production.

Results and discussion

Characterization of MOF-derived Co₃O₄

A cobalt containing zeolite imidazole framework (ZIF-67) was chosen as the sacrificing template due to its facile hydrothermal synthesis process and nitrogen-containing organic linker. MOF-derived Co₃O₄/NC samples were prepared from a two-step calcination treatment of ZIF-67 (Fig. 1b). The samples were named after the thermal treatment temperatures. For example, ZCO-700-300 (ZCO is short for ZIF-derived cobalt oxide) was obtained by calcinating ZIF-67 at 700 °C under argon for 1 hour, followed by calcination at 300 °C in air for 1 hour. The morphology and crystal structure of the pristine ZIF-67 after the hydrothermal synthesis were characterized by SEM (Fig. 2a) and XRD (Fig. 2d), showing a typical polyhedral

morphology with a particle size of 150-200 nm.²¹ After the first calcination step at 700 °C under an argon atmosphere, SEM (Fig. 2b) showed a slight decrease in the particle size (~100 nm). XRD (Fig. 2e) indicated the transformation of ZIF-67 into metallic Co, and the organic linker was converted to nitrogen-doped carbon (Fig. S1a†). The pore size distribution was well maintained after the first step calcination at 700 °C (Fig. S2a†). Therefore, the first step temperature was not optimized further. The metallic Co/NC can be further converted into desirable Co₃O₄/NC through the second calcination step in air, which shows a similar size to the sample after the first calcination step, as evidenced by SEM (Fig. 2c). The successful conversion of metallic Co into Co3O4 was also supported by the XRD patterns (Fig. 2f), and the XPS fitting spectra further suggested the existence of C-N bonds (Fig. S3†).

As the temperature of the second thermal treatment largely affects the content species and porous structure (Fig. S1 and S2†), it is critical to optimize its operating temperature. An extremely high temperature could collapse the small pores of the MOF and remove nitrogen-doped carbon, but a relatively low temperature would not fully oxidize Co to Co₃O₄. We kept the first step calcination temperature at 700 °C under argon and optimized the second step calcination temperature in air. As suggested by the TGA and TPO curves (Fig. S4†), ZCO-700-NA (NA indicates that the second step has not been completed yet) starts to decompose at 250 °C, while ZIF-67 starts to decompose at 350 °C. As such, the second step of calcination was designed to be performed at 250, 300 and 350 °C. To compare the two-step method developed by us with the generally used one-step method, samples were prepared via one-step calcination at 350 and 400 °C in air. If the second step temp-

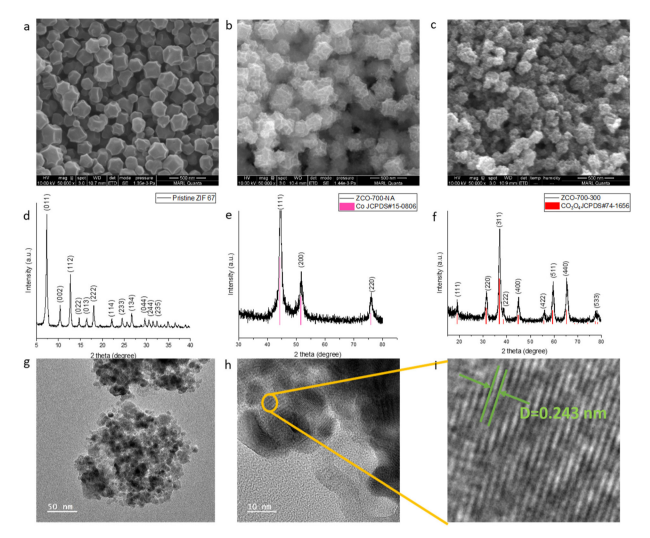


Fig. 2 Characterization of the catalysts. SEM images and XRD patterns of the (a and d) as-synthesized ZIF-67, (b and e) ZCO-700-NA, and (c and f) ZCO-700-300. (g and h) TEM images of ZCO-700-300 at different magnifications. (i) A zoom-in image of h.

erature is lower than 300 °C, Co can only be partially oxidized into Co₃O₄. For instance, as shown in the XRD patterns (Fig. S5†), ZCO-700-250 possesses both Co and Co₃O₄ phases, as compared to ZCO-700-300 with a solely Co₃O₄ phase (Fig. 2f). Further increasing the second step temperature to 350 °C would decompose all of the nitrogen-doped carbon, as shown in the TGA and TPO curves (Fig. S4†), which could undesirably increase the internal resistance of metal oxide catalysts. Energy dispersive spectroscopy (EDS) (Fig. S1†) and TGA (Table S1†) were performed to further confirm the species composition after calcination at different temperatures. EDS and TGA showed a similar trend: with an increase in the second step calcination temperature, a reduction in the amount of carbon and nitrogen was observed, along with an increase in the active content of Co₃O₄. With the second step calcination temperature over 300 °C (i.e., ZCO-700-300 and ZCO-700-350), the samples showed a similar oxygen to Co ratio, indicating that Co was fully oxidized to Co₃O₄, and 300 °C was the ideal temperature for the second calcination step.

To further understand how temperature affects the pore size distribution and surface area of ZCO samples, nitrogen physisorption was performed. The BET curves and pore size distributions are shown in Fig. S2.† All samples showed type IV hysteresis loops (Fig. S2a†), which indicates their mesoporous structures. As the second step calcination temperature increased, large-size pores appeared with an increase in their portion (Fig. S2b†). The BET surface area measured by nitrogen physisorption showed that the original pore structure in the ZIF-67 precursor collapsed during the calcination treatment, leading to a decrease in surface area (Table S1†). Therefore, the optimized second step calcination temperature was 300 °C, consistent with the EDS and TGA suggestions. The TEM image of ZCO-700-300 showed a d-spacing of 0.243 nm (Fig. 2i), which can be assigned to the Co₃O₄ (311) plane, in line with the XRD results. The above synthesis and characterization results showed the successful preparation of MOFderived Co₃O₄.

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Electrochemical measurements on MOF-derived Co₃O₄

Performance comparisons were carried out by linear scan voltammetry (LSV), as shown in Fig. 3a and S6.† ZCO-700-300 exhibited comparable performance to RuO2, and superior to commercial Co₃O₄ (C-Co₃O₄). At 100 mA cm⁻², only a 0.04 V difference in the ClER was observed between RuO2 and ZCO-700-300. Similar kinetics between ZCO-700-300 and RuO₂ was also supported by examining their Tafel slopes (Fig. 3b). The potential dependence of the ClER on ZCO-700-300 was also investigated. The FE of EO (FEEO) and current density increased by positively shifting the anodic potential, and the FE_{EO} was maintained >70% at a potential of >1.22 V vs. Ag/ AgCl (Fig. S7†). No active chlorine was detected in the absorbing vessel, and the ethylene chlorohydrin amount in the absorbing vessel was <1% in the analyte. We reasoned that this was due to the facile reaction between ethylene and hypochlorous acid (HClO). Since HClO is consumed, the reversible reaction between chlorine gas and water will always favour the production of HCl and HClO, with negligible chlorine escaping from the anolyte. Furthermore, by comparing the samples prepared at different calcination temperatures, the sample with the second step calcination temperature of 300 °C showed a superior performance to other samples (Fig. 3c), consistent with our characterization studies which showed that Co was not only fully oxidized to Co₃O₄, but also maintained a better porous structure with an optimal nitrogen doped carbon content. EIS tests were further carried out on these

samples to compare the charge transfer resistance (Fig. 3d). ZCO-700-300 showed the lowest charge transfer resistance, contributing to its good ClER activity. Although ZCO-700-250 possessed both high surface area and low charge transfer resistance, Co was not fully oxidized to Co3O4 and the remaining Co could transform to Co(OH)₂ in the aqueous electrolyte under an anodic potential. XPS confirmed the presence of Co (OH)₂ species in the ZCO-700-250 sample (Fig. S8†). It has been known that Co(OH)2 is an active oxygen evolution reaction (OER) catalyst, 22 which is the main side reaction competing for the ClER. This can be the reason for the decreased activity and selectivity of the ClER on ZCO-700-250. At the second step calcination temperature of >300 °C, a lower ClER performance could be due to the decrease in surface area (Table S1†) and increase in internal resistance. Taken together, the optimized MOF-derived Co₃O₄/NC exhibited an outstanding performance for the ClER, and careful control over the second step calcination temperature at 300 °C is critical for optimizing the ClER activity and selectivity. The performance for the ClER reached the lowest potential at 100 mA cm⁻² compared to other cobalt oxide catalysts and was even comparable to those of some noble metal catalysts, as shown in Table S2.†

The intrinsic activity was also examined by comparing their area-specific activity. It should be noted that it remains challenging to quantify the Co₃O₄ surface area in ZCO-700-300, because either the nitrogen physisorption (i.e., BET) method or the double-layer capacitance method gives the total areas of both carbon and Co₃O₄. Instead, we used the chemical surface

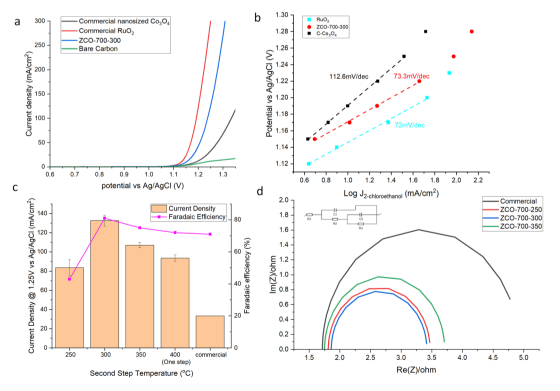


Fig. 3 CIER performance of the (a) LSV test, (b) Tafel slope, (c) current density and FE of the samples prepared at different temperatures. (d) EIS test (equivalent circuit inserted).

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area (CSA) to calculate the area-specific activity of the CIER of our reported catalysts, and the intrinsic activities of ZCO-700-300, C-Co₃O₄, and C-Co₃O₄ physically mixed with the same nitrogen doped carbon in ZCO-700-300 (C-Co₃O₄-NC) were examined. As shown in Fig. S9,† C-Co₃O₄-NC showed a similar performance compared to C-Co₃O₄. The specific activity of ZCO-700-300 was about 50% better than those of C-Co₃O₄ and C-Co₃O₄-NC. The possible reason could be (1) the heterojunction formed between carbon and Co₃O₄ during MOF calcination, which could have facilitated the electron transfer and (2) Co₃O₄ particles in ZCO-700-300 were separated by carbon, which could have increased the utilization of the particle surface area. Decoupling of these contributing factors is under way in our lab.

Furthermore, the durability of Co₃O₄/NC was evaluated at a constant current density of 100 mA cm⁻² for 10-hour electrolysis. The electrolyte was replaced by fresh electrolytes at each 2-hour interval to compensate for the consumption of the reactant Cl⁻. Electrolysis was carried out in both 0.2 M phosphate-buffered electrolyte at a pH of 6 and unbuffered 2 M KCl electrolyte. Buffered solution pH was set to 6 because it is the highest pH in which HClO is the dominant species.²³ When

the electrolyte pH is >6, the ClO ion would be the dominant species, and it could not oxidize ethylene to ethylene chlorohydrin. Whether in the buffered or unbuffered system, ZCO-700-300 maintained a remarkably stable performance in terms of both low cell potential and high FE to EO (Fig. 4a). ICP-MS confirmed a trace number of cobalt ions in the buffered anolyte (equal to <0.002% of Co₃O₄ leaching). In the unbuffered system, although the anolyte pH continuously decreased and eventually formed a strongly acidic environment, we did not observe a significant dissolution of Co₃O₄ after 10-hour electrolysis (3.6% Co₃O₄ leaching obtained from ICP-OES results). The slow cobalt dissolution rate may benefit from effective protection of Co₃O₄ by the carbon residue.^{24,25} The slightly lower FE_{EO} in the buffered system may be attributed to the use of the phosphate-buffered electrolyte, which can enhance the OER activity as phosphate has been reported to be able to facilitate the proton-coupled electron transfer process. 26,27 In addition, in terms of thermodynamic potential, the acidic environment in the unbuffered system increased the OER equilibrium potential (e.g., 0.876 V at pH 6 and 1.171 V at pH 1), while the ClER potential remained unchanged (no proton participated, 1.36 V at all pH levels). Therefore, the

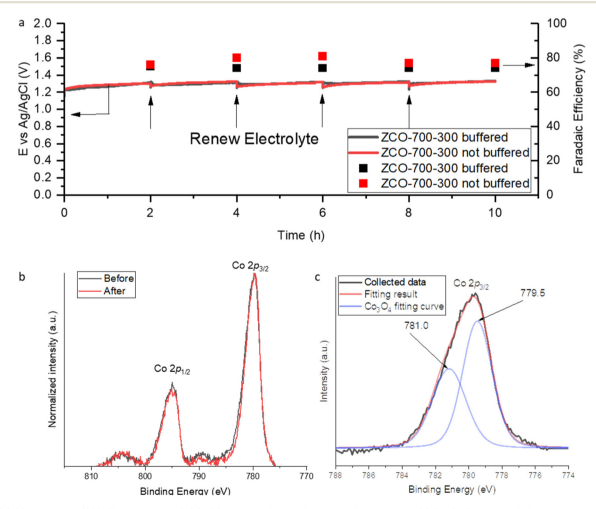


Fig. 4 (a) CP test results (90% IR compensated). (b) XPS results before and after the durability test. (c) Co₃O₄ 2p_{3/2} peak fitting curve.

standard reduction potential difference was reduced from 0.484 V to 0.136 V when the pH was changed from 6 to 1; this can also explain the slightly higher FEEO in the unbuffered electrolyte. To confirm the OER is the dominating side reaction, online gas chromatography (GC) was utilized to identify the gas product and NMR was employed to quantify soluble by-products. No signal of other C2H4 partial products was observed, indicating that O2 was the major side product (Fig. S10 and S11†). XPS was conducted to study the durability of ZCO-700-300 after 10-hour electrolysis. The peaks at 779.5 and 781.0 eV in the XPS fitting curve can be assigned to Co³⁺ and Co²⁺ in Co₃O₄, respectively. The overlapped XPS spectrum between the fresh catalyst (before the electrolysis) and spent catalyst (after 10-hour electrolysis) showed an unchanged valence state of Co (Fig. 4b and c), which indicated the robust durability of the Co₃O₄/NC catalyst.

Conclusions

In summary, we have demonstrated that a MOF-derived ${\rm Co_3O_4/NC}$ through our self-developed two-step calcination method can serve as a highly active catalyst for the chlorine-assisted production of ethylene oxide. The effects of the calcination method and temperature on the ClER activity and ${\rm FE_{EO}}$ on the ZCO samples were investigated and optimized. Benefitting from the porous structure with more exposed active sites and improved charge transfer, high ClER activity was achieved on the ${\rm Co_3O_4/NC}$ catalyst, which is comparable to a commercial ${\rm RuO_2}$ catalyst. In the 10-hour chronoamperometry test, the optimized ${\rm Co_3O_4/NC}$ catalyst exhibited remarkable stability in the buffered electrolyte, in terms of ClER activity, ${\rm FE_{EO}}$, and catalyst phase stability. The strategy for synthesizing MOF-derived ${\rm Co_3O_4/NC}$ materials could be used for the synthesis of efficient catalysts for other electrochemical reactions.

Experimental methods

Materials and chemicals

All materials and chemicals were used as received without further purification. Platinum foil (25 × 25 × 0.025 mm, 99.99%) was purchased from Strem Chemicals Inc. Cobalt(II) nitrate hexahydrate (Co(NO₃)₂·6H₂O, 98%), 2-methyl imidazole (2-MIM, 99%), ethylene oxide (EO, 50 mg mL⁻¹ in methanol) and dimethylmalonic acid (DMMA, 99%) were purchased from Sigma-Aldrich. Potassium chloride (KCl, 99%) and ethylene glycol (EG) were purchased from Fisher Chemicals. An anion exchange membrane (AEM, Fumasep FAA-3–50) was purchased from Fuel Cell Store. Commercial Co₃O₄ was purchased from US Nano Research, Inc. Commercial RuO₂ was purchased from Premetek Co.

Preparation of the precursor ZIF-67

The precursor ZIF-67 was synthesized by a hydrothermal method. 21 Typically, 0.45 g Co(NO₃)₂·6H₂O was dissolved in

3 mL of deionized (DI) water, and 5.5 g of 2-methylimidazole (2-MIM) was dissolved in 20 mL of DI water separately. Then, cobalt nitrate solution was added into 2-MIM solution slowly under stirring. The solution was stirred for 6 hours at room temperature. The purple precipitate was collected by centrifuging at 8000 rpm for 15 min and washed with water and methanol, respectively. The obtained precipitate was dried in a vacuum oven at 80 °C for 24 hours.

Preparation of ZCO samples

ZIF-derived cobalt oxide (ZCO) samples were prepared through a two-step calcination method (Fig. 1b). The first calcination step was conducted at 700 °C in a tube furnace under an argon atmosphere for 1 hour.²⁸ The second step was performed in a muffle furnace under an air atmosphere for 1 hour at different temperatures of 250, 300, and 350 °C to study the temperature effect on the activity of the samples. To compare the effect of the calcination method, the samples made by the one-step calcination method were also prepared at 350 and 400 °C under an air atmosphere.

Preparation of C-Co₃O₄-NC samples

ZIF-8 is a MOF with a similar structure to ZIF-67, which replaces Co with Zn as the metal source. ZIF-8 was chosen instead of ZIF-67, because it is easier to remove Zn, so the nitrogen-doped carbon (without metal) can be obtained. ZIF-8 was then synthesized with the same molar ratio of metal ions to the organic linker as ZIF-67 (reported in this work). The product was calcined in Ar at 700 °C for 1 hour with a temperature ramp of 5 °C min⁻¹. The product after calcination is denoted as a Zn/NC composite, which should have a similar structure to ZCO-700-NA. Zn/NC was then soaked in 0.5 M H₂SO₄ overnight to remove Zn.²⁹ The remaining Zn was <1 wt%, as confirmed by EDS. The ZIF-8 derived carbon was then mixed with commercial Co₃O₄ (C-Co₃O₄) with a 14:86 weight ratio, same as that for ZCO-700-300 according to the TGA results shown in Table S1.† The mixture was ground and dispersed in isopropanol alcohol and sprayed on 1 × 1 cm² carbon paper to reach a Co₃O₄ loading of 1 mg cm⁻². This final sample was named C-Co₃O₄-NC.

Preparation of electrodes

Commercial nanosized cobalt oxide (C–Co $_3$ O $_4$), commercial ruthenium oxide (RuO $_2$), and various self-prepared samples were loaded on hydrophilic carbon paper by spray coating. The spray ink was made of the active content and isopropanol (IPA) at a 5 mg mL $^{-1}$ concentration. The amounts of catalyst were normalized to 1 mg cm $^{-2}$ of Co $_3$ O $_4$ or RuO $_2$ according to the TGA results. Commercial samples were directly sprayed on carbon paper. Self-prepared samples were loaded by spray coating their precursor (ZIF-67 for the one-step method and ZCO-700-NA for the two-step method) on carbon paper, followed by calcination at the designated temperatures. This loading method would enhance the contact with carbon paper and prevent physical peel-off.

Characterization of the samples

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Scanning electron microscopy (SEM) images and energy dis-

persive spectroscopy (EDS) spectra were obtained using a FEI Quanta 250 FE-SEM. X-ray diffraction (XRD) data were collected from a Rigaku SmartLab diffractometer, with Cu Ko radiation ($\lambda = 0.15418$ nm) working at 40 kV and 44 mA. Transmission electron microscopy (TEM) images were captured using a JEOL 200 kV JSM-2100 scanning transmission electron microscope with a Gatan OneView 4K camera. X-ray photoelectron spectroscopy (XPS) was performed using a Kratos Amicus/ESCA 3400 instrument. Nitrogen physisorption was carried out using a Micromeritics ASAP 2020 surface area and porosity analyzer. It needs to be noted that BET measured both carbon and Co₃O₄ surface areas while carbon was inert in the ClER. Temperature-programmed oxidation was completed on the Micromeritics AutoChem II chemisorption analyzer. Leached cobalt during the durability test was determined using an inductively coupled plasma-optical emission spectrometer (ICP-OES) on a PerkinElmer Optima 8000 ICP-OES instrument and an Agilent Technologies 7700 inductively coupled plasma mass spectrometer (ICP-MS).

Electrochemical tests

Electrochemical tests were conducted on a BioLogic VSP-300 electrochemical workstation. Measurements were carried out in a typical H-type cell with an additional absorbing vessel (Fig. S12†). An anode was prepared as described in Experimental section. The cathode was platinum foil. A saturated Ag/AgCl electrode was utilized as a reference electrode. The electrolyte was 2 M KCl solution at both the anode and cathode. An anion exchange membrane was used to separate the two compartments of this cell. Ethylene gas was purged into the anolyte through a gas diffuser. An additional compartment was added to capture the chlorine gas generated on

Linear sweep voltammetry (LSV) was conducted to evaluate the activity performance of the ClER. The scanning rate of LSV was 5 mV s⁻¹. A 90% IR compensation was used to eliminate the effect of solution resistance. Electrochemical impedance spectroscopy (EIS) was performed at 1.15 V vs. Ag/AgCl electrode to measure the charge transfer resistance. Chronopotentiometry (CP) mode was utilized to test the durability of the catalyst at 100 mA cm⁻². The electrolyte was replaced with fresh electrolyte solution every 2 hours during the durability test to compensate for the consumption of chloride ions.

Product quantification

The amount of product was measured by nuclear magnetic resonance (NMR) using a Bruker NEO 400 instrument. An internal standard method was employed to quantify the product.¹⁷ Typically, the solution was made of 200 µL of D₂O, 100 μL of 15 mM DMMA solution, and 400 μL of the anolyte. The quantity was obtained by calculating the peak area ratio of the product to the internal standard and comparing it to the

calibration curve. The peak area was obtained using MestReNova software with the auto phase and baseline correction. A calibration curve generated from the known product concentration in 2 M KCl solution was used to quantify the product. An example of the NMR results and calibration curves for EG and EO are shown in the ESI (Fig. S11†). Faradaic efficiency was calculated as follows:

$$FE = \frac{2 \times F \times n}{Q}$$

where F is the Faraday constant, 96 485.33 C mol⁻¹; 2 is the number of electrons transferred when one ethylene molecule is oxidized to ethylene chlorohydrin; n is the total amount of product measured by NMR; and Q is the total amount of electrons transferred in the circuit during the whole reaction.

Active chlorine was tested using a WaterWorks free chlorine test strip. The lower detection limit was 25 ppm, corresponding to a 2 C charge. Typically, the experiment was carried out at 100 mA for 2 h, which was 720 C total charge. If the color change of the strip was not observed, we assumed that all active chlorine was consumed through reacting with ethylene.

Calculation of specific activity

Chemical surface area (CSA) is a good alternative to calculate the area-specific activity of the ClER of the reported catalysts to compare their intrinsic activity. By using the Debye-Scherrer equation, the average particle size of the metal nanoparticles could be estimated from the diffraction peak of an isolated crystal plane in XRD patterns. The average size of the Co₃O₄ nanoparticles obtained was 8 nm. The TEM image also shows that the Co₃O₄ particle size ranges from 5 to 13 nm. The specific surface area of Co₃O₄ thus can be calculated using the following equation.

$$CSA = \frac{S}{m} = \frac{S}{\rho \times V} = \frac{4 \times \pi \times r^2}{\rho \times \frac{4}{3}\pi \times r^3} = \frac{3}{\rho \times r}$$

where S is the surface area of a spherical Co particle; m is the mass; ρ is the density of cobalt oxide (6.11 g cm⁻³); V is the volume of the particle, and r is the radius of the particle. The total specific surface area of Co₃O₄ in ZCO-700-300 would be 122.75 m² g⁻¹. Then the current density based on the geometric electrode surface area could be converted to the current density based on the specific surface area. The specific activity of ZCO-700-300 was 0.108 mA cm⁻², which is 60% greater than the specific activity of commercial Co₃O₄ (0.067 mA cm⁻²). Although the surface area of the Co₃O₄ particles was not available for the ClER, the intrinsic activity of the MOF-derived Co₃O₄ with the commercial Co₃O₄ sample was calculated, based on the assumption that they exhibit same levels of catalyst utilization (= electrochemical active surface area/specific surface area).

Author contributions

T. Li synthesized and characterized MOF-derived ${\rm Co_3O_4}$ materials, and conducted most of the electrochemical tests. H. Liu helped with setting up electrochemical cells for the initial electrochemical tests and data analysis. J. Yu and W. Huang performed the ICP-MS tests and data collection and analysis. Y. Chen helped with the GC tests and some experimental design. W. Li proposed and supervised the research. T. Li, H. Liu and W. Li wrote the manuscript, and all authors discussed the results and edited the manuscript.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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