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## Microwave-Assisted Synthesis of Co/CoO<sub>x</sub> Supported on Earth-Abundant Coal-Derived Carbon for Electrocatalysis of Oxygen Evolution

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The evident demand for hydrogen as the ultimate energy fuel for posterity calls for the development of low-cost, efficient and stable electrocatalysts for water splitting. Herein, we report the synthesis of  $Co/CoO_x$  supported on coal-derived N-doped carbon via a simple microwave-assisted method and demonstrate its application as an efficient catalyst for the oxygen evolution reaction (OER). With the optimal amount of cobalt introduced into the N-doped coal-derived, the developed catalyst achieved overpotentials of 0.370 and 0.429 V during water oxidation at current densities of 1 mA cm<sup>-2</sup> and 10 mA cm<sup>-2</sup>, respectively. There was no noticeable loss in the activity of the catalyst during continuous galvanostatic polarization at a current density of 10 mA cm<sup>-2</sup> for a test period of 66 h. The synergistic interaction of the  $Co/CoO_x$  moieties with the pyridinic and pyrollic nitrogen functional groups in the N-doped carbon, as well with the other heteroatoms species in the pristine coal favored enhancement of the OER electrocatalytic performance. © The Author(s) 2019. Published by ECS. This is an open access article distributed under the terms of the Creative Commons Attribution 4.0 License (CC BY, http://creativecommons.org/licenses/by/4.0/), which permits unrestricted reuse of the work in any medium, provided the original work is properly cited. [DOI: 10.1149/2.0281908jes]

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Renewable energy-driven electrolysis of water to produce hydrogen, the ultimate energy carrier for a green and sustainable energy future, is constrained by inefficient electrocatalysis of the underlying half-cell reactions that operate at rather high reaction overpotentials, thereby rendering the process costly and commercially unattractive compared to other hydrogen production technologies.<sup>1</sup> Commercial electrolyzers typically operate at a cell voltage of 1.8-2.0 V, which is much higher than the theoretical minimum of 1.23 V. In particular, the overpotential of the oxygen evolution reaction (OER) is considerably very high compared to that of the hydrogen evolution counter reaction and is thus the major obstacle for further improvement of the efficiency of water electrolyzers.<sup>2</sup> Ir- and Ru-oxides are the present stateof-the art catalysts for the OER, their scarcity and high-prices notwithstanding, which limits any prospects of their application for industrial scale-up.3-5 In recent years, non-noble free first-row transition metalsbased materials (such as Co, Mn, Ni, Fe, Cu), have gained noticeable popularity in developing promising low-cost OER catalysts owing to innovations in material design beneficial for enhancement of kinetic phenomena and stabilization of active sites.<sup>6-9</sup> However, the intrinsically low electronic conductivity of metal oxides and the tendency of their nanoparticles to aggregate into larger clusters thereby negating the beneficial effects of high surface area, hamper optimal utilization of the full potential of these materials as OER electrocatalysts.<sup>10</sup> On the other hand, carbon materials such as graphene, carbon nanotubes, carbon black and porous carbon doped with foreign atoms, for example, sulfur, nitrogen and boron have been demonstrated to play an important role as catalysts and catalyst supports for OER.<sup>11–14</sup> The carbon structures, and their heteroatom-modified derivatives, not only lead to enhanced electron transfer but also minimize agglomeration of the catalyst particles by their embedment within the heteroatom modified carbon matrix thus leading to OER activity enhancement.<sup>15-18</sup>

Coal with an abundant global supply is one of the three most important primary energy resources in the world. However, the direct combustion of coal to generate energy is one of the leading causes of  $CO_2$  emissions in the atmosphere, the main precursor for global

warming. It is therefore vitally important to innovate new possibilities for coal beneficiation to develop value-added products.<sup>19-22</sup> With the increasing demand and urgency of developing non-noble metal catalysts for OER, pristine coal, which essentially consists of graphene stacking structures and impurity elements such as H, O, N, S, Si, Fe, Mn, among others,<sup>23–25</sup> is seen as a potential material to prepare OER electrocatalyst. Following the reports on the application of different coals as raw materials to produce N-doped porous carbon, which showed good electrocatalytic performance toward oxygen reduction reaction (ORR) in acidic media,<sup>26-29</sup> our group has also studied the influence of acid pretreatment on the electrocatalytic performance of coal-derived carbon catalysts toward ORR in alkaline media.<sup>30</sup> Recently, we reported the possibility to directly convert earth-abundant coal into high-performance bifunctional electrocatalysts for both the oxygen evolution and oxygen reduction reactions without any acid pretreatment and additional introduction of metals.<sup>31</sup> However, further improvement of the electrocatalytic performance of the oxygen reactions, is still a challenge that we have to overcome. To the best of our knowledge, no work has been reported on the preparation of a composite electrocatalyst using coal-derived carbon as support for transitional metals for improved electrocatalysis of the OER.

In this work, we report a facile and simple microwave-assisted method to synthesize  $Co/CoO_x$  supported on N-doped coal-derived carbon, which efficiently catalyzes the OER under alkaline conditions. The synergistic interaction of  $Co/CoO_x$  with the pyridinic and pyrollic nitrogen functional groups in the N-doped coal-derived carbon matrix, as well as with the other heteroatom species present in the coal-derived carbon is believed to be the origin for the good OER performance of the catalyst.

#### Experimental

*Chemicals and solutions.*—Brown coal was got from Hailar Coal Mine, Inner Mongolia as a present and its chemical compositions obtained by industrial analysis and elemental analysis have been reported previously.<sup>30,31</sup> Cobalt acetate\*4H<sub>2</sub>O ( $\geq$ 99.5%) was bought from Sinopharm Chemical Reagent Co. Ltd. (China). NaOH ( $\geq$ 96%) and ethanol ( $\geq$ 99.8%) were got from Tianjin Rgent Chemicals Co.

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Ltd. (China). Nafion (5%) was bought from Sigma-Aldrich (China). All aqueous solutions were prepared using tri-distilled deionized water.

*Sample preparation.*—The original brown coal was first crushed by ball milling, followed by pretreatment with HNO<sub>3</sub> vapor at 200°C for 12 h in order to improve the hydrophilicity of the coal sample. The sample was then thermally treated in a horizontal quartz chemical vapor deposition (CVD) reactor where it was first flushed with nitrogen at a flow rate of 50 mL min<sup>-1</sup> for 15 min prior to raising the temperature up to 800, 950, 1050 or 1100°C at a rate of 10 K min<sup>-1</sup> in a 100 mL min<sup>-1</sup> stream of NH<sub>3</sub> and Ar (1:9 volume ratio) and maintained at this temperature for 2 h. After the reaction, the reactor was cooled down in the same atmosphere with the same flow rate for 5 h and finally switched to nitrogen with a flow rate of 50 mL min<sup>-1</sup> to room temperature. The prepared coal-derived carbon catalysts are denoted as MC800, MC950, MC1050 and MC1100, respectively.

To prepare Co/CoO<sub>x</sub> decorated MC catalyst, cobalt acetate was added to a 20 mL MC aqueous suspension (3 mg mL<sup>-1</sup>) with an initial Co to MC weight ratio of 1, 5, 10, 20, 30, 40 and 50%. The suspension was subsequently ultrasonicated for 90 minutes at room temperature with an ultrasonic cell grinder (Scientz, China) and transferred to a microwave reactor (Yuhua, China) to form Co/CoO<sub>x</sub>-MC composites under a constant power of microwave (720 W) for 3 cycles with 30 s "on" and 60 s "off". During this process, the solution was continuously stirred. Then the solution was washed with distilled water to remove unreacted salts until the filtrate solution was clear. The sample was freeze dried at -55°C for 12 h in a freeze dryer (Boyikang, China) to finally obtain the resultant catalyst powder. The prepared catalysts are denoted as Co/CoOx1-MC1050, Co/CoOx5-MC1050, Co/CoO<sub>x</sub>10-MC1050, Co/CoO<sub>x</sub>20-MC1050, Co/CoO<sub>x</sub>30-MC1050, Co/CoOx40-MC1050, and Co/CoOx50-MC1050, respectively. The reference Co/CoO<sub>x</sub> sample was prepared in a similar way by adding cobalt acetate to deionized distilled water, followed by the same above-mentioned treatment process. During microwave treatment, the metal cobalt acetate and solvent water absorb microwave radiation leading to rapid internal heating and decomposition of the metal salt to form the corresponding oxide. The agitation forces caused by the microwave heating process induce homogenous deposition of the metal oxide on the carbon matrix.

Electrochemical measurements.—The pretreatment procedure of the rotating disk electrode (RDE) ( $\emptyset = 5 \text{ mm}$ , glassy carbon) used as the catalyst support for the electrochemical measurements followed our previous work.<sup>30,31</sup> Briefly it was first polished on a polishing cloth (LECO, USA) using Al<sub>2</sub>O<sub>3</sub>-pastes of different grain sizes (1, and 0.3  $\mu$ m), followed by rinsing with water. 5 mg of the catalyst powder was dispersed in a mixture of water, ethanol and Nafion (5%) with a volume ratio of 490  $\mu$ l:490  $\mu$ l:20  $\mu$ l with the aid of ultrasonication for 30 min to prepare catalyst inks. 8.3 µl catalyst suspension corresponding to a loading of 0.210 mg cm<sup>-2</sup> was carefully dropcoated on the RDE. The electrode was left to dry at room temperature in air for at least 30 min before being investigated for the oxygen evolution reaction (OER). The electrochemical performances of the composites were investigated by a CHI760E potentiostat/galvanostat (CH Instruments, China) in combination with a speed control unit (PINE Research Instrumentation, USA) in NaOH solution saturated with O2. A Pt-stick and a Ag/AgCl/3 M KCl electrode were used as counter electrode and reference electrode, respectively. The potential was scanned between 0 to 1.0 V at a scan rate of 5 mV  $s^{-1}$ , with the electrolyte saturated with O2. The rotation speed of the RDE was maintained at 1600 rpm. The long-term stability test was first performed at a constant current density of 10 mA cm<sup>-2</sup> for 66 h in O<sub>2</sub>-saturated 1.0 M NaOH solution using a graphite electrode ( $\emptyset = 6 \text{ mm}$ ) with the same catalyst loading (0.210 mg cm<sup>-2</sup>). Another durability test was also carried out by running 1000 cyclic voltammograms in the range of 0 to 0.6 V vs. Ag/AgCl/3M KCl reference electrode at a scan rate of 100 mV s<sup>-1</sup> in O<sub>2</sub>-saturated 1.0 M NaOH. For both cases, the linear sweep voltammograms before and after the test has been recorded for comparison. The potentials in the study are reported with respect to the reversible hydrogen electrode (RHE), according to the conversion  $E = E_0 + E_{Ag/AgCl} + 0.059^*$ pH - iR<sub>s</sub>, where  $E_0$  is the measured potential vs. Ag/AgCl/3M KCl,  $E_{Ag/AgCl}$  is 0.207 V for the standard potential of the Ag/AgCl/3M KCl reference electrode with respect to the standard hydrogen electrode at zero pH, R<sub>s</sub> is the electrolyte resistance read out directly from the CHI software at the open circuit potential and i is the electrocatalytic OER current.

**Characterization of samples.**—Transmission electron microscopy (TEM) (HITACHI H-600) and scanning electron microscopy (SEM) (HITACHI SU8010) were applied to characterize the morphologies of MC1050 and Co/CoO<sub>x</sub>-MC1050. A D8 (Bruker) X-ray diffractometer with Cu K<sub>a</sub> radiation ( $\lambda = 1.5418$  Å) and a Bruker spectrometer with a 532 nm laser source were used for recording X-ray diffraction (XRD) patterns and Raman spectra, respectively. An ESCALab250Xi electron spectrometer with a 300 W Al K<sub>a</sub> radiation source was applied for X-ray photoelectron spectroscopy (XPS) measurements.

#### **Results and Discussion**

The electrochemical performance of the MC and Co/CoO<sub>x</sub>-MC catalysts toward the OER was characterized by linear sweep voltammetry (LSV) in O<sub>2</sub>-saturated 0.5 M NaOH solution (pH 13.7) at a scan rate of 5 mV s<sup>-1</sup>. During the measurements, the electrode was continuously rotated at 1600 rpm to prevent gas bubble accumulation on the electrode surface. Figure 1A presents the polarization curves of MC1050 and after its modification with different loadings of  $Co/CoO_x$ , as well as the  $Co/CoO_x$  sample as reference. MC1050 displayed insignificantly low anodic currents compared to its counterparts modified with Co/CoO<sub>x</sub> thus indicating poor activity in electro catalyzing the OER. In contrast, very promising OER activity was observed in the case of the Co/CoOx-MC1050 composites clearly highlighted by two important features, that is, a significant negative shift of the OER onset potential and a positive increase of the OER current density. The onset potential for oxygen evolution, defined as the potential corresponding to a current density of 1 mA cm<sup>-2</sup>, on Co/CoO<sub>x</sub>40-MC1050 was 1.610 V, which was the smallest among the investigated samples. Co/CoOx40-MC1050 also required the least potential (1.664 V), compare to the other samples, Co/CoO<sub>x</sub>50-MC1050 (1.711 V), Co/CoO<sub>x</sub>30-MC1050 (1.709 V), Co/CoO<sub>x</sub>20-MC1050 (1.738 V), Co/CoO<sub>x</sub>10-MC1050 (1.789 V), Co/CoO<sub>x</sub>5-MC1050 (1.773 V), Co/CoO<sub>x</sub>1-MC1050 (1.809 V), as well as pure Co/CoO<sub>x</sub> (1.731 V) to reach 10 mA cm<sup>-2</sup>. A current density of 10 mA cm<sup>-2</sup> has been proposed as the figure of merit for estimating the efficiency of a given OER catalyst for viable solar driven fuel synthesis.<sup>32</sup> The inclusion of cobalt thus evidently promoted the OER activity. However, the decrease of the electrochemical performance with increase of the amount of the cobalt precursor above 50% might be due to agglomeration of Co/CoOx particles, which decrease the active surface area exposed to the solution. In addition, a high coverage with Co/CoO<sub>x</sub> in relation to carbon seems to mask the potential contribution of carbon as a conductive matrix thereby diminishing the charge transfer rate. The influence of the thermal treatment temperature of MC in NH<sub>3</sub>/Ar on the electrocatalytic performance of the microwave-synthesized Co/CoO<sub>x</sub> toward OER was subsequently studied as shown Figure 1B. Co/CoO<sub>x</sub>40-MC1050 displayed the best OER performance with respect to the onset potential for the OER at 1 mÅ cm<sup>-2</sup> ( $E_{onset} = 1.610$  V), and the potential necessary to attain a current density of  $10 \text{ mA cm}^{-2}$  (E<sub>10</sub> = 1.664 V), which were both much better than Co/CoO<sub>x</sub>40-MC950 ( $E_{onset} = 1.627$  V and  $E_{10} = 1.685$  V) and Co/CoO<sub>x</sub>40-MC800 ( $E_{onset} = 1.644$  V and  $E_{10} = 1.718$  V), except Co/CoO<sub>x</sub>40-MC1100 that attained nearly similar values ( $E_{onset} =$ 1.614 V and  $E_{10} = 1.667$  V). High-temperature treatment would favor the incorporation of nitrogen into the lattice positions of carbon forming graphitic nitrogen thus leading to the increase of structure disorder with more surface and edge defects, which would favor the subsequent introduction of  $Co/CoO_x$ .<sup>31,33</sup> Further to this, high temperature



**Figure 1.** Linear sweep voltammograms (iR drop corrected) showing the electrocatalytic activity toward OER of (A) pure Co/CoO<sub>x</sub> and MC1050, and MC1050 modified with different loadings of Co/CoO<sub>x</sub>; (B) MC modified with Co/CoO<sub>x</sub> and thermally treated at different temperatures, namely: 800°C, 950°C, 1050°C and 1100°C; (C) OER activity of Co/CoO<sub>x</sub>40-MC1050 in O<sub>2</sub>-saturated NaOH at different concentrations that is: 0.1, 0.5 and 1 M NaOH. The electrode was maintained at a rotation speed of 1600 rpm and scan rate of 5 mV s<sup>-1</sup> during the measurements; (D) Tafel slopes derived from (C); (E) long-term chronoamperometric stability test of Co/CoO<sub>x</sub>40-MC1050 supported on a graphite electrode at a constant current density of 10 mA cm<sup>-2</sup> in O<sub>2</sub>-saturated 1 M NaOH; (F) Linear sweep voltammograms before and after the experiment in (E).

annealing of carbon, especially in the presence of transition metal species is known to increase the graphitic degree of the carbon as well as its conductivity. The electrochemical impedance spectra (EIS) of MC samples thermally treated at different temperatures were investigated at 1.65 V vs. RHE over the frequency range from 100 kHz to 0.01 Hz in 1.0 M NaOH or 5 mM K<sub>3</sub>[Fe(CN)<sub>6</sub>] in 1.0 M NaOH solution. When recording the impedance in 1.0 M NaOH, it was not straightforward to determine the charge resistance from the resulting Nyquist plots due to the complexity and heterogeneous nature of the system. EIS was thus also measured in a solution of 5 mM K<sub>3</sub>[Fe(CN)<sub>6</sub>] in 1.0 M NaOH. In this case,

during ac perturbation at the open circuit potential of the established Fe<sup>3+/</sup>Fe<sup>2+</sup> redox couple, the measured charge transfer resistance from the Nyquist plots corresponds to resistance to electron transfer across the catalyst films owing to the high reversibility of the Fe<sup>3+/</sup>Fe<sup>2+</sup> redox couple (Figure S3). The electrochemically active surface area of the prepared samples i.e. Co/CoO<sub>x</sub>-MC800, Co/CoO<sub>x</sub>-MC950, Co/CoO<sub>x</sub>-MC1050, and Co/CoO<sub>x</sub>-MC1100, loaded on the glassy carbon electrode are calculated to be 14.65, 25.25, 24.70, and 27.85 cm<sup>2</sup>, by running cyclic voltammetry in the range of 1 to 1.15 V vs. RHE at different at different scan rates (5, 10, 20, 50, 100 and 200 mV s<sup>-1</sup>) (Figure S4). The OER performance of

Sample	Catalyst loading (mg cm <sup>-2</sup> )	Electrolyte	E <sub>OER</sub> (V)@ 10 mA cm <sup>-2</sup>	η <sub>OER</sub> (V)	Tafel slope (mV dec <sup>-1</sup> )	Ref.
Co/CoOx40-MC1050	0.21	O <sub>2</sub> -sat. 1 M NaOH	1.659	0.429	58	This work
Co@CoO@CO <sub>3</sub> O <sub>4</sub> - N/C	0.25	O <sub>2</sub> -sat. 1 M NaOH	1.680	0.45	39	34
Co(OH) <sub>2</sub> NPs	0.1	O <sub>2</sub> -sat. 1 M NaOH	1.559	0.339	40	35
$Co_3O_4$	0.22	1 M KOH	1.655	0.425	44	36
Co/Co <sub>3</sub> O <sub>4</sub> 40-NG	0.21	O <sub>2</sub> -sat. 1 M NaOH	1.667	0.437	99	37
CoP/CDs-18.97%	0.35	1 M KOH	1.61	0.4	93	38
Co <sub>3</sub> O <sub>4</sub> @Ni	0.07	1 M KOH	1.650 <sup>a</sup>	0.42	65	39
De-LCO NPs (LiCoO <sub>2</sub> )	0.1	0.1 M KOH	1.62	0.39	57	40
Co <sub>3</sub> O <sub>4</sub> -C	1.76	1 M KOH	1.609	0.38	78	41
$RuO_2$	0.21	O <sub>2</sub> -sat. 1 M KOH	1.64	0.41	-	42
IrO <sub>2</sub>	0.21	O <sub>2</sub> -sat. 1 M KOH	1.70	0.47	-	42

### Table I. A comparison of the performance of cobalt-based materials as OER catalysts based on recent literature.

<sup>a</sup>The potential was initially reported vs. Ag/AgCl/3M KCl, and it was corrected vs. RHE in this table.

Co/CoOx40-MC1050 was further characterized in O2-saturated 0.1 M NaOH (pH 13) and 1 M NaOH (pH 14), respectively in order to study the influence of the electrolyte pH. Better electrocatalytic performance of the catalyst was achieved with increase of the NaOH concentration. This is not surprising because the OH<sup>-</sup> ions are reactants in the OER in alkaline solution, according to the equation:  $4OH^- \rightarrow$  $O_2 + 2H_2O + 4e^-$ . Therefore, increase of the NaOH concentration would accelerate the kinetics of the OER by increasing the exchange current density, which is directly correlated with the OH- concentration. The potential required to obtain a current density of 1 mA  $\rm cm^{-2}$ and 10 mA cm<sup>-2</sup> was 1.600 and 1.659 V, respectively, with corresponding overpotentials ( $\eta$ ) of 0.370 and 0.429 V, when Co/CoO<sub>x</sub>40-MC1050 was characterized in O2-saturated 1 M NaOH. This performance is comparable to that reported for the state-of-the-art cobaltbased OER catalysts and commercial RuO2, and surpasses some of advanced carbon materials (e.g. graphene or carbon nanotubes) supported cobalt-based catalysts and commercial IrO2 (Table I).34-42 Figure 1D shows the Tafel plots of the Co/CoOx40-MC1050 characterized in different NaOH solutions derived from Figure 1C. The Tafel slopes were 54 and 58 mV dec<sup>-1</sup> for the Co/CoO<sub>x</sub>40-MC1050 characterized in 0.5 M and 1 M NaOH, respectively, much lower than in 0.1 M NaOH (94 mV dec<sup>-1</sup>). From the general expression for the Tafel slope of an electrode reaction:  $b = \frac{2.303RT}{\alpha F}$ , where R is the universal gas constant (8.314 J K<sup>-1</sup> mol<sup>-1</sup>), T is the temperature (298 K),  $\alpha$  is the transfer coefficient, and F is the Faraday constant (96485 C mol<sup>-1</sup>), a slope of ~118.3 mV dec<sup>-1</sup>, assuming  $\alpha = 0.5$ , would be expected if the first electron transfer step (i) in the OER mechanism outlined below is rate limiting. M + OH<sup>-</sup>  $\rightarrow$  MOH + e<sup>-</sup> (i); M-OH<sub>ads</sub> (A)  $\rightarrow$  M-OH<sub>ads</sub> (B) (i'), A and B denoting surface sites characterized by different energy levels which is typical of heterogeneous catalysts, as in the present case of Co/CoO<sub>x</sub>-MC, due to surface diffusion during the OER.<sup>43,44</sup>  $MOH + OH^{-} \rightarrow MO + H_2O + e^{-}$  (ii);  $MO + OH^{-} \rightarrow MOOH + e^{-}$ (iii); MOOH + OH<sup>-</sup>  $\rightarrow$  MO<sub>2</sub> + H<sub>2</sub>O + e<sup>-</sup> (iv); and MO<sub>2</sub>  $\rightarrow$  M + O<sub>2</sub> (v). In the case of an anodic reaction, the anodic transfer coefficient  $\alpha_a$ is defined as:  $\alpha_a = \mathbf{r} \cdot (\mathbf{1} - \mathbf{\beta}) + \mathbf{n}' / \mathbf{v}$ , where  $\beta$  is the symmetry factor, v is the stoichiometric number, n' is the total number of electrons exchanged in the elementary steps preceding the rate determining step (RDS) in the mechanism, and r is a factor that can have only two values: 0 or 1, when the RDS is a chemical or electrochemical step, respectively.<sup>45–47</sup> If the chemical step after the first electron transfer is rate limiting, A Tafel slope of  $\sim$ 59.2 mV dec<sup>-1</sup> is obtained. If the second electron transfer step of the OER (step (ii)) is rate limiting, a Tafel slope of  $\sim 40 \text{ mV} \text{ dec}^{-1}$  would be obtained. The existence of a Tafel slope of about 118 mV dec<sup>-1</sup> implies kinetically slow adsorption of the OH<sub>ads</sub> intermediate (step (i)), so that a low surface coverage of the electrode with adsorbed reaction intermediates is expected in the entire OER potential region. The observed Tafel slopes of 54 mV dec $^{-1}$ 58 mV dec<sup>-1</sup> and 94 mV dec<sup>-1</sup> of the OER on Co/CoO<sub>x</sub>40-MC1050

in 0.5, 1.0, and 0.1 M NaOH, respectively, thus indicate that the OER was considerably kinetically faster in 0.5 M and 1.0 M NaOH compared to 0.1 M NaOH. At lower NaOH concentration (0.1 M NaOH), the formation of adsorbed OH<sup>-</sup> species in the primary discharge step  $(M + OH^{-} \rightarrow MOH + e^{-})$  and subsequent surface diffusion seem to be the limiting factors for the overall OER rate. On the other hand, the smaller values of the Tafel slope at higher NaOH concentrations (0.5 or 1 M NaOH) are certainly due to a different rate controlling elementary step, either M-OH<sub>ads</sub> (A)  $\rightarrow$  M-OH<sub>ads</sub> (B) or M-OH<sub>ads</sub> + OH-  $\rightarrow$  M-O<sub>ads</sub> + H<sub>2</sub>O + e<sup>-.48-50</sup> The Tafel plots for different Co/CoOx-MC1050 samples with different Co/CoOx loadings derived from Figure 1A and the samples of Co/CoO<sub>x</sub>40 loaded on MC thermally treated at different temperatures, derived from Figure 1B, are also presented in Figure S1. It is anticipated that an optimal morphology should favor rapid OH<sup>-</sup> diffusion and adsorption to the catalyst surfaces, such that, the rate-control step is exclusively dependent on the intrinsic catalyst properties. One of the most important concerns on applying carbon-based material as OER catalysts is whether they can remain stable under the strongly oxidative conditions during the OER. Moreover, a stable catalytic performance in high-concentration alkaline solutions is critical for realistic applications.<sup>51</sup> Therefore, a long-term stability chronopotentiometric test was performed by applying a constant current density of 10 mA cm<sup>-2</sup> in 1 M NaOH with  $Co/CoO_x$ 40-MC1050 deposited on a graphite electrode as the support. The potential response was stable for the investigated period of at least 66 hours, moreover, a slight decrease of the overpotential was observed (Figure 1E). Additionally, linear sweep voltammograms were also recorded before and after the chronopotentiometric experiment (Figure 1F). The LSVs further demonstrated that the OER activity of the catalyst was well preserved in the concentrated electrolyte. Slight but noticeable improvement of the OER performance was observed after the long-term chronopotentiometric experiment, which might be due to morphological changes of the catalyst film, particularly, removal of unstable amorphous carbon exposed to the electrolyte due to the forces induced by detachment of the oxygen gas bubbles and concomitant increased accessibility of the Co/CoO<sub>x</sub> based active sites. On the other hand, all metallic elements when used as catalysts for the OER undergo surface oxidation to form surface oxides, or higher oxides. Metallic cobalt for example is oxidized to cobalt oxide, cobalt oxyhydroxide (CoOOH). Depending on the size of the metallic particles, the oxidation might occur gradually. We previously observed continuous activity enhancement of cobalt and nickel boride and phosphide catalysts due to progressive oxidation of metals with potential cycling, and concomitant increase of the OER activity.<sup>52,53</sup> Therefore, the fact that the OER activity increases marginally after stability testing is certainly due to oxidation of metallic cobalt and cobalt oxide (CoO) to form cobalt oxyhydroxide (CoOOH), which is believed to be the active state for the OER. We also performed the stability test of



Figure 2. SEM images of (A) MC1050 and (B) Co/CoO<sub>x</sub>40-MC1050, and the corresponding elemental mapping.

Co/CoO<sub>x</sub>40-MC1050 by running 1000 cyclic voltammograms (CVs) between 0 and 0.6 V vs. Ag/AgCl/3M KCl at a scan rate of 100 mV s<sup>-1</sup> in 1 M NaOH saturated with oxygen (Figure S2A). Only a marginal change in the CVs was observed. The LSVs recorded before and after CV measurement also confirmed a negligible decay during the stability test. The potential difference at the current density of 10 mA cm<sup>-2</sup> was only 10 mV (Figure S2B). Because a moderate potential sweep was applied, the strong surface oxidation process would not take place in this case.

To gain further insight into the properties of the prepared catalysts, SEM and EDX elemental mapping were used to study the morphology and chemical composition of MC1050 and the optimized catalyst, Co/CoO<sub>x</sub>40-MC1050 (Figure 2). It is believed that the surface defects in MC1050, including oxygen-functional groups were active sites for the initial nucleation of Co owing to the strong electrostatic interaction between oxygen-functional groups on the surfaces and the Co cations, followed by further growth into 3-dimensional nanoparticles.<sup>37</sup> As shown in Figure 2B, Co/CoO<sub>x</sub> particles were homogeneously distributed on MC1050. The presence of a higher amount of the cobalt precursor inevitably led to the agglomeration of formed Co/CoO<sub>x</sub> nanoparticles under high-energy microwave treatment. The corresponding elemental maps of the distribution of C, N, O and Co are also displayed, clearly confirming the homogeneous distribution

of the different elements. Moreover, Al, Ca, and Si were also detected, which obviously arise from the ash in the coal.

Co/CoO<sub>x</sub>40-MC1050 was additionally characterized by highresolution TEM in order to probe the nanostructure of the prepared sample in-depth (Figure 3). Figure 3A confirms that a large proportion of the nanoparticles were embedded in the carbon framework. Both metallic Co and its oxide,  $Co_3O_4$ , were found to be present, as indicated by the observed lattice fridges with interspacing of 0.205 and 0.244 nm, corresponding to (111) and (311) crystal planes, respectively (Figure 3B).<sup>54,55</sup> Meanwhile, Ca<sub>2</sub>Al<sub>2</sub>SiO<sub>7</sub> and SiO<sub>2</sub> also coexisted in the sample, confirmed by the presence of the crystal lattice spacings of 0.200 and 0.429 nm, assigned to the (212) and (100) crystal planes, respectively. This is consistent with the findings in our previous study where similar species were observed and obviously arise from the impurities in the original coal.<sup>31</sup> Moreover, the TEM images reveal that the catalyst particles adopt a core-shell like structure with the Co/CoO<sub>x</sub> nanocrystallites being firmly embedded inside the MC matrix and surrounded with a few layers of graphene (d = 0.334 nm) (Figure 3C).

The catalyst was further characterized by Raman spectroscopy to obtain insight into the graphitization degree of the carbon materials. Figure 4 shows the Raman spectra of MC1050, and MC1050 modified with the Co/CoO<sub>x</sub> nanoparticles. The Raman spectra of both MC1050



Figure 3. (A) TEM image of  $Co/CoO_x 40$ -MC1050, and high-resolution TEM images of (B) metallic Co,  $Co_3O_4$ ,  $Ca_2Al_2SiO_7$  and  $SiO_2$  nanoparticles; and (C) metallic Co surrounded with graphene layers.



Figure 4. Raman spectra of MC1050 and Co/CoO<sub>x</sub>40-MC1050 (The spectra were normalized with respect to the corresponding D-bands).

and Co/CoOx40-MC1050 exhibit two characteristic main peaks, a Dband at ca. 1350 cm<sup>-1</sup> and a G-band at ca. 1588 cm<sup>-1</sup>, originating from vibration of sp<sup>3</sup> carbon atoms due to structural defects and the  $E_{2g}$  vibration mode of sp<sup>2</sup> carbon domains, respectively.<sup>56,57</sup> The intensity ratio of the D- and G-band  $(I_D/I_G)$ , were calculated to be 1.00 and 0.96 for MC1050 and Co/CoO<sub>x</sub>40-MC1050, respectively. The  $I_D/I_G$ ratio was similar with and without the introduction of Co/CoOx indicating that the synthesis of Co/CoOx-MC1050 composites did not introduce significant surface defects to MC1050.37 The existence of a 2D-band at 2686 cm<sup>-1</sup> and a (D+G)-band at 2921 cm<sup>-1</sup> provides additional proof for distortion of the aromatic graphitic domains.<sup>58,59</sup> The  $I_{2D}/I_{D+G}$  intensity ratio decreased after the introduction of cobalt, which is ascribed to the coverage of the MC-1050 surfaces by Co/CoO<sub>x</sub> nanoparticles and increase of the graphitic degree of the carbon. This also gives a cue that the Co/CoOx nanoparticles prefer to anchor on the surface defects. The peaks at 464 and 663 cm<sup>-1</sup> confirm the presence of  $Co_3O_4$ .<sup>37</sup>

The XRD spectrum of Co/CoO<sub>x</sub>40-MC1050 was compared with that of MC1050, as shown in Figure 5. Two peaks at  $2\theta = 26.6^{\circ}$  and at 44.7° attributed to (002) and (100) planes of graphitic carbon predominated the diffraction patterns of both MC1050 and Co/CoO<sub>x</sub>40-MC1050. In addition, both samples contained rather weakly resolved diffraction peaks, similar to our previous observations for related materials.<sup>30</sup> Particularly, no clear diffraction peaks ascribable to either metallic Co or its oxides were observed in the XRD diffraction patterns, which is partly due a low concentration of Co and CoO<sub>x</sub> species in relation to carbon in these catalysts, and also because the



Figure 5. XRD spectra of MC1050 and Co/CoO<sub>x</sub>40-MC1050.

particles are mainly embedded inside graphene layers as revealed by the high-resolution TEM images.

XPS was used to study the chemical state and interaction(s) of the various components (Figure 6). Co/CoO<sub>x</sub>40-MC1050 has clear Co  $2p_{3/2}$  and Co  $2p_{1/2}$  doublet signals at 782.5 eV and 798.4 eV, respectively, with a spin-orbit splitting of 15.9 eV indicating that Co<sup>2+</sup> rather than Co<sup>3+</sup> species were the dominant oxide form of cobalt (Figure 6A).<sup>60</sup> The corresponding shoulder peaks at a slightly high binding energy of both peaks are due to a shake-up process, which is generally observed for Co<sup>2+</sup> species in the high spin state.<sup>61</sup> The XPS results therefore further confirm that CoO was the dominant surface species. In consideration of the TEM results, and the fact that XPS is a surface sensitive technique, the detected CoO oxide is believed to be a native oxide on the surface of metallic cobalt formed due to atmospheric exposure. The deconvoluted spectrum of the Co 2p region is dominated by the  $Co^{2+}$  due to  $CoO.^{61,62}$  The thermal treatment of cobalt and carbon precursors at high temperature in an ammonia atmosphere in principle leads to the formation of nitrogen functional groups as well as cobalt coordinated to nitrogen within the carbon matrix. The Co-N coordination is weak and the Co-N group has only been observed in very few studies by XPS, where it manifests as a weak shoulder at about 778.8 eV, and by X-ray absorption fine structure studies.<sup>42</sup> In the present XPS studies, the Co-N group was not discernible, nonetheless, its presence cannot be excluded. It should however be noted that the binding energies for both Co 2p<sub>3/2</sub> and Co 2p<sub>1/2</sub> peaks are slightly higher than those reported for the standard spectra of pure CoO or Co<sub>3</sub>O<sub>4</sub>.<sup>62</sup> A similar chemical shift of binding energy of Co  $2p_{3/2}$  was also observed upon the pyrolysis of cobalt and tetramethoxy-phenyl porphyrin (CoTMPP) at 600°C in a nitrogen atmosphere.<sup>63</sup> The co-existence and interaction of the Co, N, O and C active moieties with the other elements and complex oxide species arising from the impurity elements such as Al, Mg, Ca, Fe, Mn, Si originally present in the pristine coal, is not to be neglected.<sup>31</sup> Although not definitively clear at this moment, interaction of the impurity species with the active moieties may lead to modification of the surface electronic structure of the active moieties and ultimately the OER activity of the composite catalyst. In line with this, the slight positive shift in the binding energy of the Co 2p<sub>3/2</sub> and Co 2p<sub>1/2</sub> peaks compared to that of pure cobalt oxide species is most likely due to the interaction of cobalt with the heteroatom elements in the coal. The O 1s peak can be deconvoluted into three peaks (Figure 6B). The two peaks at 531.9 and 533.3 eV are due to the presence of C=O and C-O functional groups, respectively. The peak centered at 530 eV would relate to anionic oxygen associated with oxide species. The introduction of oxygen functional groups on the surface of carbon materials during nitric acid treatment is commonly observed.<sup>64</sup> However, in this case, the introduction of cobalt led to a decrease of the total oxygen content, as manifested by the drastic decrease of the intensity of the peak at 531.9 eV. This suggests that the C=O related groups underwent decomposition during the microwave process.<sup>65</sup> Moreover, the C=O sites could act as potential anchoring sites for the nucleation and growth of cobalt during microwave treatment. Increase in the intensity of the peak at 530 eV upon the introduction of cobalt was not evident. This implies that the amount of oxygen introduced by the formation of cobalt oxide was negligible. On the other hand, XPS is a surface-sensitive technique, a drastic decrease of the amount of the detected oxygen species might be due to the MC1050 covered by Co/CoO<sub>x</sub> nanoparticles. The N 1s peak was deconvoluted into three peaks corresponding to pyridinic (398.4 eV), pyrrolic (400.1 eV) and graphitic (401.3 eV) nitrogen (Figure 6C). The influence of surface functionalization by HNO3 vapor and the subsequent thermal treatment in ammonia was previously investigated,<sup>30</sup> where a decrease of the relative O/C ratio was observed accompanied by an increase of the relative N/C ratio. Oxygen functional groups created during HNO3 vapor treatment are known to favor the functionalization of carbon with nitrogen atoms during ammonia treatment. The relative concentration of the various nitrogen species varied upon the additional introduction of cobalt as indicated by the specific intensities of the respective species in the deconvoluted N 1s spectra. The relative ratio of pyridinic



Figure 6. (A) XP Co 2p, (B) O 1s, and (C) N 1s spectra of MC1050 and Co/CoO<sub>x</sub>40-MC1050.

nitrogen decreased while that of pyrrolic nitrogen increased after the microwave treatment process. This indicates that the introduction of transition metals may partially catalyze the decomposition of nitrogen groups upon heating. In our case, it is believed that the cobalt preferentially catalyzes the decomposition of pyridinic nitrogen-contained groups. A similar observation was reported in our previous study for a catalyst containing iron.<sup>30</sup>

#### Conclusions

A simple microwave-assisted method for synthesis of a coalderived N-doped carbon matrix (MC) embedding cobalt oxide species  $(Co/CoO_x-MC)$  proved to be versatile and reliable to prepare a very efficient non-precious catalyst for the OER under alkaline conditions. Promising OER activity with overpotentials of 0.370 V and 0.429 V at a current density of 1 mA cm<sup>-2</sup> and 10 mA cm<sup>-2</sup>, respectively, was realized when an optimal amount of cobalt (40 wt%) was introduced into coal derived N-doped carbon (Co/CoOx40-MC1050). This sample also shows good stability without noticeable loss in activity during chronopotentiometric polarization at a constant current density of 10 mA cm<sup>-2</sup> for a test period of 66 h. Metallic Co and its oxides, CoO and  $Co_3O_4$ , were identified in the composite based on TEM, Raman spectroscopy and XPS analysis. The synergistic effect arising from the presence of the heteroatoms present in the pristine coal and the interaction of Co/CoOx with N-doped coal-derived carbon favored enhancement of the OER electrocatalytic performance.

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