Size-controlled large-diameter and few-walled carbon nanotube catalysts for oxygen reduction†

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We demonstrate a new strategy for tuning the size of large-diameter and few-walled nitrogen-doped carbon nanotubes (N-CNTs) from 50 to 150 nm by varying the transition metal (TM = Fe, Co, Ni or Mn) used to catalyze graphitization of dicyandiamide. Fe yielded the largest tubes, followed by Co and Ni, while Mn produced a clot-like carbon morphology. We show that morphology is correlated with electrocatalytic activity for the oxygen reduction reaction (ORR). A clear trend of Fe > Co > Ni > Mn for the ORR catalytic activity was observed, in both alkaline media and more demanding acidic media. The Fe-derived N-CNTs exhibited the highest BET (∼870 m² g⁻¹ and electrochemically accessible (∼450 m² g⁻¹) surface areas and, more importantly, the highest concentration of nitrogen incorporated into the carbon planes. Thus, in addition to the intrinsic high activity of Fe-derived catalysts, the high surface area and nitrogen doping contribute to high ORR activity. This work, for the first time, demonstrates size-controlled synthesis of large-diameter N-doped carbon tube electrocatalysts by varying the metal used in N-CNT generation. Electrocatalytic activity of the Fe-derived catalyst is already the best among studied metals, due to the high intrinsic activity of possible Fe–N coordination. This work further provides a promising route to advanced Fe–N–C nonprecious metal catalysts by generating favorable morphology with more active sites and improved mass transfer.

Introduction

Polymer electrolyte fuel cells (PEFCs) are promising clean energy conversion devices due to their high efficiency (up to 85%) and lack of undesirable emissions, including CO₂. However, the oxygen reduction reaction (ORR) at the cathode of H₂/air PEFCs is a sluggish reaction, which requires a catalyst containing a large amount of precious metals (e.g., Pt) to achieve sufficient reaction activity and stability. Due to the high cost and scarcity of Pt, new nonprecious metal catalysts (NPMCs) derived from earth-abundant elements that match the high performance and durability of Pt catalysts are desperately needed. Recently, highly graphitized nitrogen-doped nanocarbon materials derived from nitrogen–carbon precursors via a high-temperature approach in the presence of transition metals (TMs, e.g., Fe, Co, Ni) have attracted substantial attention due to their exceptionally high activity and stability compared to other NPMC formulations. According to other researchers and ourselves, in situ formation of graphitized carbon nanostructures in the M–N–C catalysts is a critical factor dictating active site generation and is linked to the measured ORR activity. The currently prepared carbon nanostructures in M–N–C catalysts include tubes, onion-like carbon, and platelets (multi-layered graphene). As a result, their surface areas are much higher than conventional multi-walled carbon nanotubes (MWNTs). As a result, their surface areas are much higher than conventional MWNTs. In addition, the large diameter provides a mesoporous structure that facilitates mass transfer into and out of the tubes. Thus, these novel carbon nanotubes provide a new opportunity to develop high-performance
Nanoporous metal–organic frameworks (MOFs) have shown promise as templates for preparing carbon-based nonprecious metal catalysts (NPMCs) for the oxygen reduction reaction (ORR). In this work, we focused on optimizing the activity by using metal–organic frameworks (MOFs) as templates. In this work, as a replacement for expensive MOFs, a carbon black, BlackPearl 2000, was used to prepare the tube-like catalysts. Importantly, we observed the size-dependence of the carbon nanotubes produced using different TMs including Ni, Co, Fe, and Mn, with a low-cost precursor (e.g., dicyandiamide, price: $1–2 kg⁻¹). The tube diameter and level of nitrogen incorporation correlate directly with the measured catalytic activity for the ORR. Although many studies have explored the role of TMs in NPMC preparation, few reports have systematically correlated the choice of metal, resulting morphologies, and catalyst properties. Herein, we discovered the morphology and chemical doping of carbon nanotubes can be fine-tuned by selection of the TM used in their preparation. This, in turn, provides control of electrochemical properties and catalytic activity. Fe-derived carbon nanotubes have the largest diameter, followed by Co- and Ni-derived ones. While the detailed mechanism through which the TM controls the tube size remains uncertain, we show that the TM choice plays a significant role in NPMC preparation, influencing key catalyst properties including morphology, surface area, elemental composition, and ORR activity. Thus, this work provides a new path toward development of high-performance carbon-based NPMCs by controlled formation of favorable nanocarbon structures.

Experimental section

Catalyst synthesis

Large-diameter and few-walled nitrogen-doped carbon nanotubes (N-CNTs) were prepared by first mixing 24 mmol dicyandiamide with 6 mmol of one of the metal acetates (MAc, M = Fe, Co, Ni or Mn) in 250 mL ethanol. The solution was heated to 78 °C in a covered 2 L Erlenmeyer flask and held at this temperature for 5 h. Next, 0.2 g BlackPearl 2000 carbon black was added and the mixture was stirred for an additional 24 h before uncovering the flask and allowing the solvent to evaporate. The resulting dry powder was heated to 1000 °C for 1 h in a tube furnace under a constant N2 flow of 50 mL min⁻¹. The heat-treated sample was then suspended in 0.5 M H2SO4 at 80 °C for 5 hours to remove unstable metal species. Finally, a second heat treatment, at 1000 °C for 3 h, was carried out to remove the functional groups and oxidative carbon layers in the catalysts to further improve the ORR activity.

Physical characterization

The size and morphology of catalysts were characterized using transmission electron microscopy (TEM) on a JEOL JEM-2010 or a JEOL 2100F microscope at a working voltage of 200 kV. Samples were prepared for imaging by drop casting a dilute suspension of the catalyst onto a carbon-coated TEM grid. Catalyst morphology was also studied using scanning electron microscopy (SEM) on a Hitachi SU 70 microscope at a working voltage of 5 kV. The crystalline phases present in each sample were identified using powder X-ray diffraction (XRD) on a Rigaku Ultima IV diffractometer with Cu K-α X-rays. X-ray photoelectron spectroscopy (XPS) was performed using a Kratos AXIS Ultra DLD XPS system equipped with a hemispherical energy analyzer and a monochromatic Al Kα source. The monochromatic Al Kα source was operated at 15 keV and 150 W; pass energy was fixed at 40 eV for the high resolution scans. All samples were prepared as pressed powders supported on a metal bar for the XPS measurements. Brunauer Emmett Teller (BET) surface area was measured using N2 adsorption/desorption at 77 K on a Micromeritics TriStar II. Samples were pre-treated at 150 °C for 3 h under vacuum prior to nitrogen physisorption measurements.

Results and discussion

Electrochemical measurements

All electrochemical measurements were performed using a CHI Electrochemical Station (Model 660b) in conventional three-electrode cell at room temperature. A glassy carbon rotating disk electrode (GC-RDE) was used as the working electrode. Each catalyst was mixed with isopropanol and a Nafion solution to produce an ink that was drop-cast onto the GC-RDE and air-dried for 30 min at 60 °C. The catalyst-coated GC-RDE was subjected to cyclic voltammetry (CV) in N2-saturated 0.5 M H2SO4 electrolyte at a scan rate of 20 mV s⁻¹ to determine the electrochemically accessible surface area. Then, to measure the electrocatalytic activity for the ORR, rotating ring disc electrode (RRDE) measurements were performed in O2-saturated 0.5 M H2SO4 and in 0.1 M NaOH at room temperature and a rotation rate of 900 rpm. A graphite rod was used as the counter electrode. An Hg/HgSO4 (K2SO4-sat., 0.705 V vs. RHE) or an Ag/AgCl electrode (KCl-sat., 0.996 V vs. RHE) was employed as the reference electrode in acidic and alkaline electrolytes, respectively. The loading of catalyst was controlled at 800 µg cm⁻² for all measurements. ORR steady-state polarization curves were recorded using discrete potential steps of 0.05 V at intervals of 30 s from 1.0 V to 0 V vs. RHE.

Controlled morphologies and structures

We developed a facile and low-cost synthesis strategy to prepare a new type of large-size and few layered tube catalysts by using various metals, dicyandiamide, and Blackpearl 2000 carbon black. The tube size can be controlled via the choice of metal (Fe, Co, or Ni). The Fe-derived tube catalyst exhibits the largest size followed by Co- and Ni-derived ones as evidenced by direct SEM image observation (Fig. 1). Synthesis using Fe, Co or Ni yielded highly graphitized tubes with bamboo-like structures. Based upon measurement of approximately 35 tubes from each sample, the Fe-derived carbon nanotubes exhibited the largest average diameter (Fig. 1a,b and S1†), 112 nm, while the tubes derived from Co and Ni had average diameters of 61 and 42 nm, respectively (Fig. 1c,d and S1†).

Prior work suggests that the growth of carbon nanotubes is triggered by dicyandiamide with 6 mmol of one of the metal acetates (MAc, M = Fe, Co, Ni or Mn), resulting in a tube-like morphology. In this work, we discovered that the morphology and chemical doping of carbon nanotubes can be fine-tuned by selection of the TM used in their preparation. This, in turn, provides control of electrochemical properties and catalytic activity. Fe-derived carbon nanotubes have the largest diameter, followed by Co- and Ni-derived ones. While the detailed mechanism through which the TM controls the tube size remains uncertain, we show that the TM choice plays a significant role in NPMC preparation, influencing key catalyst properties including morphology, surface area, elemental composition, and ORR activity. Thus, this work provides a new path toward development of high-performance carbon-based NPMCs by controlled formation of favorable nanocarbon structures.
with 3d-electrons of the TMs.\textsuperscript{29} Thus, the formation of metal (Fe, Co or Ni) nanoparticles by reduction of the metal salts is expected to initiate the formation of carbon nanotubes. We believe the growth mechanism of carbon nanotubes is similar to that of traditional carbon nanotubes. The tip-growth mechanism is clearly evidenced by SEM images like Fig. 1b, showing Fe-derived tubes with metal particles attached at the tips of the tubes prior to acid leaching.

Additional images showing the typical morphology of Fe-derived carbon nanotubes are provided in Fig. S2,\textsuperscript{†} illustrating the highly porous and 3D carbon nanotube network architecture generated by this synthesis process. Notably, the diameter of Fe-derived carbon nanotubes (up to 150 nm) was significantly larger than the size of traditional N-doped carbon nanotubes prepared by CVD methods (usually less than 50 nm).\textsuperscript{30–34} Tang et al.,\textsuperscript{34} employed a CVD method to prepare N-doped carbon nanotubes using nanowire templates. The N-CNT growth method demonstrated here is also much simpler and lower cost than traditional CVD.\textsuperscript{34,35} The unique capability to tune the tube size over a wide range by varying type of metals (e.g., Fe, Co, and Ni) was discovered, which further influences the ORR activity of the corresponding catalysts. The trend of increasing tube diameter with decreasing atomic number of the TM did not continue for Mn. As shown in Fig. 1e, irregular clot-like structures were generated, rather than carbon nanotubes. Fig. 1f shows carbon materials synthesized under identical conditions in the absence of any metal. No features other than those associated with the Black-

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Fig. 1 SEM images of carbon catalysts prepared using: (a) Fe, (b) Fe but before acid leaching treatment, (c) Co; (d) Ni; (e) Mn; (f) no TM. The mean diameters of the tubes prepared using each metal were (a,b) Fe (112 nm), (c) Co (61 nm), and (d) Ni (42 nm). Diameter distributions are provided in ESI, Fig. S1.\textsuperscript{†}
pearl carbon particles were observed, suggesting that no significant amount of highly graphitized nanocarbon forms in the absence of metal.

The morphologies of carbon nanotubes were further studied by TEM imaging (Fig. 2). In good agreement with SEM results, a clear trend of tube size was observed in the sequence Fe > Co > Ni. In addition, Mn-derived nanocarbon exhibited a hollow, irregular, quasi-spherical carbon particle morphology. This observation is consistent with prior studies of nucleation and growth of carbon nanotubes using TM catalysts. Overall, the most active catalysts for carbon nanotube formation are Fe, Co and Ni. Furthermore, HR-TEM images shown in Fig. 3 were further used to reveal the subtle structures of carbon nanotubes and nanoshells observed in Fe- and Mn-derived samples, respectively. Unlike traditional carbon nanotubes with continuous hollow structures, these ultra-large diameter carbon nanotubes present bamboo-like morphologies, in which a number of cages are connected continuously (Fig. 3a). The wall thickness of a typical carbon tube was 6–10 layers of carbon planes (Fig. 3b). The term “few-walled” is used in this work to describe graphitic structures with few layers (less than 10 layers). Thus, we refer to these large carbon tubes as few-walled carbon nanotubes. Notably, the carbon layers in the carbon nanotube exhibit substantial defects such as vacancies and dislocations, which can be beneficial for electrocatalysis. In good agreement with the observation in Fig. 1b, during the formation of carbon nanotubes, carbon shells are growing along the metallic agglomerates and ultimately encapsulate some of them. Rather than the bamboo-like carbon nanotubes derived from Fe, isolated carbon cages were observed in the Mn-derived sample as shown in Fig. 3c. Notably, well-defined and highly ordered carbon layer structures were produced in the presence of Mn (Fig. 3d).

These TEM imaging results suggest that Mn is highly effective in catalyzing the graphitization of carbon precursors (i.e. dicyandiamide) during the high temperature treatment. However, lack of defects in the ordered carbon structures can be a disadvantage for electrocatalysis and may significantly limit the electrocatalytic activity. These structural observations are in good agreement with the much lower doping level of nitrogen and low activity for the ORR measured with the Mn-derived samples discussed below. ESI Fig. S3A-D† show the dependence of tube morphology upon heating temperature. We found that 700 °C is the minimum temperature required for the formation of tubes, and further increasing the temperature from 700 °C to 1000 °C increased the yield of material with tube morphology, suggesting higher temperature is more favorable the tube growth. The dependence of ORR activity upon the heating temperatures during the synthesis was studied for the best performing Fe-based catalysts (Fig. S4†). The sample treated at 1000 °C showed the highest activity in both acidic and alkaline electrolytes. Although previous studies have reported production of nitrogen-doped carbon catalysts with tunable physical and chemical properties, here we provide the first demonstration that the diameter of carbon tubes can be manipulated by varying the TM used in the precursor mixture. Given the important role played by the TM in controlling nanocarbon morphology, we anticipate similarly strong effect on the catalytic properties of these nanocarbons.
Powder XRD (Fig. 4a) showed the presence of graphitic carbon, metal carbides, and metals in each of the carbon nanotube samples only subjected to the first treatment and before acidic leaching. The diffraction peak near $2\theta = 26.5^\circ$, assigned to the (002) plane of graphitic carbon, is present in all samples. The enlarged view of these peaks in Fig. 4b shows a slight shift to lower diffraction angles following the trend of Fe $<$ Co $\approx$ Ni $<$ Mn. The fact that the Fe-derived sample exhibits the largest shift, 0.1°, implies that it has the largest lattice constant among the samples. In the case of Fe-derived carbon nanotubes, the XRD peaks at 43.6°, 45.0° and 51.0° were assigned to iron carbide (Fe$_7$C$_3$) and the peaks at 44.5° and 64.7° are attributed to metallic Fe. In the XRD patterns for Co and Ni-derived carbon nanotubes, metallic Co and Ni were identified, along with small amounts of Co$_2$C and NiC. The XRD pattern of the Mn-derived sample showed the presence of Mn$_3$C but no metallic Mn in the final product. These observations suggest that the metallic species (e.g. Fe, Co and Ni) may be important for generating a large-diameter tube morphology during the high-temperature graphitization process.

Nitrogen doping

Nitrogen doping into nanocarbon electrocatalysts has been hypothesized by many to be important to their ORR activity. We employed XPS to systematically study the effect of the TM on the level and position of nitrogen doping in these nanocarbon samples. Results are summarized in Table 1. The N content in the carbon nanotubes derived from Fe is the highest (4.49 at%), followed by Co (4.22 at%), Ni (3.86 at%), and Mn (1.78 at%). This trend is consistent with the shifts of the (002) peaks observed in the XRD patterns. The peak position shifts to lower angle with increased N-doping, suggesting that higher nitrogen doping produces a larger $d$-spacing between graphite layers. In addition, elemental analysis showed that traces of metal remained in these carbon nanotube samples. During the leaching treatment with 0.5 M H$_2$SO$_4$ at 80 °C for 5 hours, those metallic species that are fully enclosed by carbon shells are inaccessible to the acid and are not removed. The residual content of Ni in the nanocarbon was higher than that of Co and Fe (Ni $>$ Co $>$ Fe). The amount of residual metal species in the carbon nanotubes decreased with increasing tube diameter. This suggests that leaching out the metal residues from the smaller tubes is more difficult than from larger tubes, which is reasonable. Prior studies have proven that such metal species are inactive for the ORR, especially in acidic electrolyte. Thus, removal of these inactive species can increase the activity and enhance the stability of the nanocarbon electrocatalysts.

We also analyzed high-resolution N 1s XPS spectra (Fig. 5) to determine the chemical state of the nitrogen dopant in these nanocarbons. Each sample featured two dominant nitrogen peaks at 401.2 and 398.7 eV. These can be assigned to graphitic and pyridinic nitrogen, respectively. This suggests that nitrogen atoms are doped both within the carbon planes (graphitic) and at the edges (pyridinic). Nitrogen atoms can be regarded as n-type dopants in carbon, and have been associated with high ORR activity. The ratio of graphitic nitrogen to pyridinic nitrogen (Table 1) has been considered to be an important factor in determining the ORR catalytic activity. The graphitic nitrogen content in Mn-derived carbon clots (60%) is notably higher than that in
Fe, Co and Ni-derived carbon nanotubes, which were all nearly identical (54–55%). In ESI Fig. S5† it compares the XPS spectra of C 1s for the samples derived from Fe, Co, Ni and Mn. Overall, the broad peak can be decomposed into three components. The sharp peak at 284.7 eV can be assigned to the C–C bonds in graphitic carbon structures. The full width at half-maximum (FWHM) of Mn-derived carbon clots was 1.10 eV, which was smaller than carbon nanotubes derived from Fe (1.25 eV), Co (1.25 eV), and Ni (1.20 eV). The comparison of C 1s spectra suggests a relatively high degree of graphitization in the Mn-derived nanocarbon. This result is consistent with its XRD pattern, which showed the sharpest (002) peak among the nanocarbons studied here. The peak at 286.2 eV is assigned to C–O. The largest C–O content was observed in Fe-derived catalysts, consistent with its relatively high oxygen content, as shown in Table 1. These samples all showed a very small contribution from C=O at higher binding energy, near 288.1 eV.

**BET and electrochemical surface areas**

In addition to nitrogen doping, the ORR activity is strongly influenced by the nanocarbon morphology, including its surface area and pore size distribution. Thus, BET surface areas of these nanocarbons were measured using N2 physical adsorption at 77 K. The measured surface areas as function of the metal used for the synthesis are shown in Fig. S6.† The Fe-derived carbon nanotube sample, which showed the largest tube diameters, also exhibited the highest surface area (868 m² g⁻¹), followed by the nanocarbons derived from Co (466 m² g⁻¹), Ni (391 m² g⁻¹), and Mn (167 m² g⁻¹). The nitrogen adsorption–desorption isotherms for these samples are compared in Fig. S7.† The Fe-derived sample exhibits the most dominant mesoporous feature, consistent with the largest tube size. Thus, the Fe-derived carbon nanotubes with largest diameter are expected to provide the largest number of active sites for the ORR with favorable mass transport.

While BET surface area determined by N2 adsorption is important to characterize porous catalysts, electrochemically accessible surface area (Sₐ) is more relevant for electrocatalysis, and was measured to determine the utilization of surface area available for the ORR. Fig. 6 shows CV curves for each sample in the absence of O₂. These were used to compute the Sₐ in 0.5 M H₂SO₄ solution (a) and 0.1 M NaOH solution (b). The upper branch of the voltammograms between 0.2 and 0.6 V in acidic or alkaline solution is attributed to pseudocapacitance associated with the electrolyte-accessible surface area of the nanocarbon catalyst. Thus, the effective Sₐ (m² g⁻¹) can be deduced from the gravimetric double layer capacitance C (F g⁻¹). C can be represented as: \[ C = \frac{I}{vm} \]  
where I is the current, m is the electrode mass, and v is the scan rate (here 20 mV s⁻¹). The Sₐ value can then be determined as: \[ Sₐ = \frac{C}{C_{GC}} \]

where C_{GC} is the double layer capacitance (F m⁻²) of a glassy carbon electrode surface, typically 0.2 F m⁻².⁴² The values of Sₐ calculated from eqn (1) and (2) are summarized in Table S1† for both acidic and alkaline electrolytes. Compared to other samples, the Fe-derived carbon nanotubes show the highest Sₐ, reaching 419 m² g⁻¹ and 403 m² g⁻¹ in acidic and alkaline solutions, respectively. The values of Sₐ decreased in the same order, Fe > Co > Ni > Mn, in both electrolytes.

**Electrocatalytic activity for the ORR**

Electrocatalytic activity of the nanocarbon catalysts for the ORR was evaluated in both acidic and alkaline electrolytes using a RRDE at room temperature and a rotation speed of 900 rpm. Generally, TM-derived N–M–C catalysts exhibit improved ORR activity, relative to metal-free carbon catalysts. Thus, we did not compare with the poor catalytic activity of the sample prepared without any metal, but focus on comparing the results obtained using the different TMs. Fig. 7a and b compare the steady-state ORR polarization curves measured in 0.5 M H₂SO₄ and 0.1 M NaOH, respectively. Notably, in order to minimize the possible effects of post-treatment on ORR catalytic activity, these samples are not the final catalysts, but were only subject to the first heat treatment and the acid leaching step. At this synthesis stage, the different TMs indeed lead to significantly different catalytic activity for the ORR. In both acidic and alkaline media, the onset potential of the ORR for TM-derived carbon samples follows the order of Fe > Co > Ni > Mn, indicating that the nature of the catalytically active sites varied with the metals used for synthesis. In particular, the most positive onset potential, measured with the Fe-derived sample, reached 0.89 V, followed by Co-(0.84 V), Ni-(0.60 V), and Mn-(0.58 V) derived samples. The Fe-derived sample also showed the most positive half-wave potential (E₁/₂) at 0.71 V, indicating that it provided the largest number of active sites available for the ORR. Similarly, the Fe-derived catalyst also exhibited the highest activity in the alkaline solution, again showing the most positive onset and E₁/₂ potentials.

Based on our previous effort to develop high-performance carbon-based NMPCs, a second heat treatment is able to remove the oxygen-containing functional groups and corrosive carbon species from catalysts, thereby further enhancing the
ORR activity. Thus, for the two most promising Fe- and Co-derived catalysts, we applied a second heat treatment at 1000 °C in N₂ for 3 h. A comparison of ORR activities of these samples before and after the second heat treatment at a rotation rate of 900 rpm is shown in ESI Fig. S8.† Both Co-derived and Fe-derived catalysts exhibited dramatically improved activity in both the kinetically-controlled and mass-transfer controlled potential regions, indicating a significant increase in the number of active sites for the ORR. In particular, the onset potential of the Fe-derived catalyst was shifted from 0.89 V to 0.95 V in acidic solution. In the alkaline solution measurement, the onset potential was also positively shifted by 30 mV, to 1.05 V, but the change was much smaller than in acidic solution. These results clearly show improved catalytic performance after the second heat-treatment. Notably, the difference in activity between Fe- and Co-based catalysts subjected to the second heating is much smaller in alkaline electrolyte, relative to acidic electrolyte. Once again, it is very likely that the active sites for the ORR in acid and alkaline media are different. The final Fe- and Co-derived carbon nanotube catalysts after the second heating treatment were further compared with the commercial 20 wt% Pt/C catalysts. ORR activities in acid and alkaline electrolytes are shown in Fig. 7c and d, respectively. Like other state-of-the-art nonprecious metal catalysts, in the more challenging acidic media, Co- and Fe-based catalysts are still inferior to Pt/C. However, in the alkaline media, the ORR activities are quite comparable to, or even exceeded, that of the Pt/C, demonstrating their great promise to replace precious metals in alkaline fuel cells and metal-air batteries.

The high resolution XPS spectra of the N 1s region for the catalysts synthesized using Fe and Co before and after the second heat-treatment are shown in ESI Fig. S9 and Table S2.† After the second heat-treatment, the total N content of the Fe-derived catalyst was reduced from 4.49 at% to 2.93 at%. However, the relative graphitic N content increased by around 6.5%, relative to the pyridinic N content. In contrast, the second heat-treatment did not lead to a drop of total nitrogen content for the Co-derived catalyst, but reduced the relative graphitN content. Comparing the high resolution XPS spectra of Fe and Co-derived catalysts in the C 1s region (ESI Fig. S10†), C–C and C=O peaks became narrower in both cases. This suggests an increased level of graphitization after the second heat treatment. Also, the total O content dropped significantly in both Fe and Co derived catalysts. Therefore, after the second heat treatment, both catalysts exhibited a higher degree of graphitization and lower concentration of inactive surface ligands, as indicated by the reduced O content. We conclude that those factors enhanced ORR performance in both Fe and Co derived catalysts. Notably, the reduced total N content does not result in a decrease of ORR activity for Fe-based catalysts. Once again, relative to total N content, achieving an optimal ratio of graphitic to pyridinic N is more important to ORR activity.

In addition to onset potential, the limiting current density also varied significantly between catalysts derived using different TMs. The Fe-derived catalyst showed the highest limiting current density. This can be attributed to two factors. First, its high electrochemically accessible surface area with mesoporosity provides efficient mass transfer within catalyst layers, thereby increasing limiting current. Second, the Fe-derived catalyst showed the highest selectivity toward four-electron reduction of O₂ to H₂O over two-electron reduction to H₂O₂, as indicated by results in Fig. 7e and f. These show H₂O₂ yields in 0.5 M H₂SO₄ and 0.1 M NaOH, respectively. Fe-derived catalysts only yield 5% and 2% peroxide in the acidic and alkaline electrolytes, respectively. In contrast, the Mn-derived sample showed low four-electron selectivity, with 50% and 15% peroxide yield in acid and alkaline electrolytes, respectively. The selectivity of Co and Ni-derived samples fell between those of Fe- and Mn-derived samples. This order was consistent with the order of their limiting current densities.

Based on current understanding along with the correlation of catalytic activity and physical properties, the ORR activity order of Fe > Co > Ni > Mn was determined by differences in electrochemically accessible surface area and nitrogen doping. At present, the detailed nature of active sites in the N–M–C catalysts is still of debate, centered on the function of the transition metal during the catalyst synthesis. Some researchers have asserted that metals only catalyze nitrogen doping into
tubes with di-
ad Ni are very e-
catalyst preparation. Among studied metals, Fe, Co
emerged as a new type of high-performance nonprecious metal
catalyst for the ORR. In this work, we build upon our previous
emphasized that key morphology-related factors such as surface
area, defect concentration in carbon nanostructures, and
optimal nitrogen doping play important roles in improving
overall catalyst performance. Compared to Fe, Co, and Ni, we
show that Mn was much more effective in catalyzing the for-
mation of perfect graphite-like carbon structures, as demon-
strated by the sharp carbon (002) peak in XRD, the more
negative binding energy in C 1s XPS, and the highly ordered
structures visible in HR-TEM images. However, Mn failed to
induce sufficient nitrogen doping to produce a highly active
catalyst. The perfect graphitic structures with low nitrogen
content produced with Mn resulted in the lowest ORR activity
among the catalysts considered here. In contrast, a substantial
concentration of defects and dislocations was identified in Fe-
based carbon catalysts, which were the most active electroca-
lysts. This supports the view that defects in the carbon nano-
tubes catalyzed by Fe are critical for achieving high ORR
activity. In addition, Fe-based catalysts have the most favorable
morphology, providing more active sites (high electrochemi-
cally active surface area) and efficient mass transfer (large-size
tubes) for the ORR electrocatalysis.

Notably, although Fe-derived carbon nanotube catalysts had
the highest ORR activity, they had the lowest ratio of electro-
chemically accessible surface area to BET surface area among
the four catalysts studied here. Thus, the Fe-derived carbon
nanotube catalyst still has the highest potential for further
ehancement of ORR catalytic activity by further optimizing
morphology and structure to increase the catalyst utilization,
i.e. active surface area that is electrochemically accessible and,
therefore, contributes to ORR activity.

Conclusion
Large-size and few-walled N-doped carbon nanotubes have
emerged as a new type of high-performance nonprecious metal
catalyst for the ORR. In this work, we build upon our previous
effort to demonstrate a novel approach for fine-tuning the
diameters of large-size carbon nanotubes by varying the TM
used for catalyst preparation. Among studied metals, Fe, Co
and Ni are very effective in catalyzing formation of carbon
tubes with different sizes following the order Fe (110 nm) > Co
(60 nm) > Ni (40 nm). Rather than tubes, Mn yields cage-like
nanocarbon with a relatively high degree of graphitization, low
concentration of defects, and low nitrogen content. In con-
trast, substantial numbers of dislocations and defects were
found in Fe-based bamboo-like carbon nanotubes. Meanwhile,
ORR activity measured with these nanocarbon samples in
both acid and alkaline electrolytes followed the same
sequence, Fe > Co > Ni > Mn in terms of their onset/half-wave
potentials, four-electron selectivity, and limiting current den-
sities. Notably, the critical second heat-treatment during the
synthesis reduces the ORR activity difference between Co- and
Fe-based catalysts, and this difference was much smaller in
alkaline media than that in acid. These results strongly
suggest that the active sites for the ORR in acid and alkaline
media are significantly different.

This work establishes a synthesis–morphology–property
relationship that can provide a basis for further catalyst
improvements. Compared with other nanocarbons, the unique
large-size and few-walled carbon nanotubes derived from Fe
have the largest surface area, both in terms of traditional BET
surface area and the more relevant electrochemically acces-
sible surface area. This favorable morphology provides the
highest number of active sites for the ORR. The level of nitro-
gen doping was consistent among these nanotube samples,
which allowed us to specifically study the morphology-depend-
ence of the ORR activity. Although the intrinsic activity of the
catalyst may be associated with the nature of likely M–N moi-
eties (Fe–N or Co–N), morphology clearly also plays an impor-
tant role in governing the catalyst activity. This work provides
new insights into the origins of catalytic activity, which can be
used to design and synthesize advanced nanocarbon catalysts
for electrochemical energy conversion.

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