Unraveling the Nature of Sites Active toward Hydrogen Peroxide Reduction in Fe-N-C Catalysts

Chang Hyuck Choi,* Won Seok Choi, Olga Kasian, Anna K. Mechler, Moulay Tahar Sougrati, Sebastian Brüller, Kara Strickland, Qingying Jia, Sanjeev Mukerjee, Karl J. J. Mayrhofer, and Frédéric Jaouen*

Abstract: Fe-N-C catalysts with high O₂ reduction performance are crucial for displacing Pt in low-temperature fuel cells. However, insufficient understanding of which reaction steps are catalyzed by what sites limits their progress. The nature of sites were investigated that are active toward H₂O₂ reduction, a key intermediate during indirect O₂ reduction and a source of deactivation in fuel cells. Catalysts comprising different relative contents of FeₙCₓ moieties and Fe particles encapsulated in N-doped carbon layers (0–100%) show that both types of sites are active, although moderately, toward H₂O₂ reduction. In contrast, N-doped carbons free of Fe and Fe particles exposed to the electrolyte are inactive. When catalyzing the ORR, FeₙCₓ moieties are more selective than Fe particles encapsulated in N-doped carbon. These novel insights offer rational approaches for more selective and therefore more durable Fe-N-C catalysts.

The oxygen reduction reaction (ORR), a key reaction for acidic fuel cells, is today most efficiently catalyzed by Pt materials.[1] However, the large-scale deployment of acidic fuel cells will be confronted with the grand challenge of developing inexpensive catalysts with high activity and stability.[2] While metal–nitrogen–carbon catalysts (M-N-C, especially M=Fe or Co) are promising,[3] further advances and fundamental insights into the factors governing their catalysis and durability are needed to meet the stringent industrial requirements.

The nature of active sites in pyrolyzed Fe-N-C catalysts is a highly complex and debated topic.[4] The simultaneous presence of multiple Fe species in most catalysts investigated hitherto has obscured the interpretation of their ORR behaviors, especially since multiple sites may lead to unexpected synergies. In particular, several types of active sites may be necessary to catalyze the multielectronic ORR, which may proceed either through a direct 4e⁻ pathway on a single site (O₂ → H₂O₂), or through indirect pathways involving a desorbed H₂O₂ intermediate (O₂ → H₂O₂ → H₂O) on two sites. These may be identical in nature (consequential 2e⁻ × 2e⁻ mechanism), or different (bifunctional 2e⁻ + 2e⁻ mechanism). If the ORR proceeds via the bifunctional mechanism, one type of site (S1) reduces O₂ to H₂O₂ and another (S2) catalyzes the peroxide reduction reaction (PRR). Under this hypothesis, only Fe-N-C catalysts comprising a high density of sites S1 and S2 may catalyze the ORR with an apparent low % H₂O₂. PRR catalysis is also highly desirable for improved durability of Fe-N-C catalysts because even low H₂O₂ production can lead to significant degradation during fuel cell operation.[5]

From advanced spectroscopy (Mössbauer[6] and X-ray absorption spectroscopy (XAS),[7] and mass spectrometry[8]) or molecular-probe approaches,[9] there is a growing consensus that FeₙCₓ moieties or N-C species encapsulating Fe particles (Fe@N-C) catalyze the transfer of the first 2e⁻ during ORR (S1).[10] Then, if the ORR mostly follows the 2e⁻ + 2e⁻ pathway, highly selective Fe-N-C catalysts should be characterized by a high number of sites S2.[11] It has been recently suggested that Fe particles or pyridinic-N groups might be the site S2.[12] However, improved understanding on...
the nature of the sites active toward PRR is still pivotal for improved durability. To determine whether Fe-N-C materials catalyze the ORR via a direct, consequential, or bifunctional mechanism, we have investigated the PRR activity on a set of catalysts, including model catalysts only comprising either FeN\textsubscript{x}C\textsubscript{y} moieties or Fe particles.

In the continuation of our recent studies\cite{10b,13}, the catalysts were synthesized by pyrolysis of Fe\textsubscript{II} acetate, 1,10-phenanthroline (Phen), and a Zn\textsuperscript{II} zeolitic imidazolate framework (ZIF-8). Three catalysts were first prepared, named FeNC-wet-1, FeNC-dry-1, and FeNC-dry-0.5, the labeling referring to homogenization conditions and Fe content before pyrolysis (see Methods in the Supporting Information). A catalyst pyrolyzed without Fe\textsubscript{II} acetate was also prepared (“NC”). FeNC-wet-1, FeNC-dry-1, and FeNC-dry-0.5 showed Fe content of ca. 3.4, 3.0, and 1.5 wt% after pyrolysis, respectively, while about 100 ppm Fe was detected with inductively coupled plasma mass spectrometry (ICP-MS) for NC.\cite{14} X-ray diffraction (XRD) and Raman spectroscopy revealed carbonization of ZIF-8 and Phen after pyrolysis (Supporting Information, Figure S1). X-ray photoelectron spectroscopy (XPS) shows similar N-doping level and N-components for all catalysts (Supporting Information, Figure S2, Table S1).

To identify the Fe structures in the catalysts, we examined their morphology with transmission electron microscopy (TEM), which revealed a larger number of Fe particles in FeNC-wet-1 than in FeNC-dry-1, with most Fe particles embedded in N-doped carbon layers (Supporting Information, Figure S3). \textsuperscript{57}Fe Mössbauer spectroscopy showed that the crystalline Fe particles seen in TEM images correspond to α-Fe, Fe carbide, or γ-Fe (Figure 1a–c; Supporting Information, Table S2). The relative absorption area owing to Fe particles was 47% for FeNC-wet-1 and 8% for FeNC-dry-1. No signal assigned to Fe particles was detected for FeNC-dry-0.5, the Mössbauer spectrum of which shows only the doublets D1 and D2 (FeN\textsubscript{x}C\textsubscript{y} moieties). These observations were confirmed by extended X-ray absorption fine structure (EXAFS), with a lower intensity of the Fe-Fe backscattering signal for FeNC-dry-1 vs. FeNC-wet-1 (Figure 1d). As previously reported\cite{10b}, a strong Fe-N(O) interaction without Fe-Fe interaction for FeNC-dry-0.5 confirms the sole presence of FeN\textsubscript{x}C\textsubscript{y} moieties. Quantification of the absolute content of each Fe component was derived from the Mössbauer spectra fittings (Figure 1e).\cite{6c} Thus, Fe-N-C catalysts with high (FeNC-wet-1) and low content (FeNC-dry-1) of Fe particles are identified, along with those devoid of Fe particles (FeNC-dry-0.5) and nearly devoid of Fe altogether (NC). Furthermore, a catalyst labeled FePhen@MOF-\textsubscript{ArNH\textsubscript{3}} exclusively comprising Fe particles was investigated.\cite{10a} Its Fe content was 3.1 wt% (Figure 1e), with Fe being present as metallic, carbide, and nitride particles.

The electrochemical properties were then measured using a rotating disk electrode (RDE) in 0.1 m HClO\textsubscript{4} electrolyte.

![Figure 1](https://example.com/figure1.png)

**Figure 1.** \textsuperscript{57}Fe Mössbauer absorption spectra and their fittings with five spectral components for a) FeNC-wet-1, b) FeNC-dry-1, and c) FeNC-dry-0.5. d) Fourier transforms of the EXAFS spectra. e) Absolute Fe contents of the two main sub-groups of Fe species.

![Figure 2](https://example.com/figure2.png)

**Figure 2.** Electrochemical characterization. a) ORR polarization curves, b) \textsubscript{H}2\textsubscript{O} formation measured with RRDE, and c) PRR polarization curves: 900 rpm and 800 \textmu g cm\textsuperscript{-2} (additionally, 100 \textmu g cm\textsuperscript{-2} for RRDE studies). d) Correlations between PRR activity and absolute content of either total Fe or Fe sub-components. PRR activity is reported as the slope dJ/dE seen in (c) at high potential (see the Supporting Information).
All four of the Fe-N-C catalysts showed high ORR activity (Figure 2a), comparable to that of other high-performing Fe-N-C catalysts.\textsuperscript{[14]} The NC catalyst showed poor ORR kinetics, as expected in acidic medium.\textsuperscript{[4,10a,b]} Rotating ring disk electrode (RRDE) measurements at 800 μg cm\textsuperscript{-2} loading showed low H\textsubscript{2}O\textsubscript{2} production (< 6%) during ORR for the four Fe-N-C catalysts (Figure 2b and S4). However, at 100 μg cm\textsuperscript{-2}, up to 32% H\textsubscript{2}O\textsubscript{2} production was measured. This indicates that H\textsubscript{2}O\textsubscript{2} produced at a given catalytic site may be consecutively reduced to H\textsubscript{2}O. This suggests that, on some Fe sites at least, H\textsubscript{2}O\textsubscript{2} may desorb and then re-adsorb on the same type or on another type of Fe sites having a higher affinity for H\textsubscript{2}O\textsubscript{2}. Furthermore, the positive correlation between % H\textsubscript{2}O\textsubscript{2} and absolute content of Fe particles at low loading (Supporting Information, Figure S4, right) suggests FeN\textsubscript{C}, moieties catalyze the ORR to H\textsubscript{2}O more selectively than Fe@NC sites. Owing to 15–30% H\textsubscript{2}O\textsubscript{2} produced during ORR at low loading, investigating the PRR activity of different sites in Fe-N-C catalysts is important to rationally improve their selectivity.

The PRR activity was measured in Ar-saturated 0.1 M HClO\textsubscript{4} electrolyte containing 1.3 or 10 mM H\textsubscript{2}O\textsubscript{2} (Figure 2c). The NC catalyst showed no PRR activity, indicating that surface N-groups unpromoted by Fe cannot be a site S2. In contrast, high PRR current on FeNC-dry-0.5 demonstrates that FeN\textsubscript{C}, moieties are PRR-active. This is a first major finding of the present study, enabled by the synthesis of Fe-N-C catalysts free of Fe particles.\textsuperscript{[10b,15]}

Previous work hypothesized that Fe-N-C catalysts catalyze the ORR via a bifunctional 2e\textsuperscript{-} + 2e\textsuperscript{-} mechanism in acidic medium; the site S1 for 2e\textsuperscript{-} ORR being FeN\textsubscript{C}, moieties and site S2 for PRR being Fe particles.\textsuperscript{[12a]} However, the turnover frequency of FeN\textsubscript{C}, moieties for PRR is much lower than that for ORR (FeNC-dry-0.5: lower PRR than ORR current at 0.8 V\textsubscript{RHE} for the same reactant concentration of 1.3 mM, Figure 2a,c). This suggests that ORR on FeN\textsubscript{C}, moieties predominantly occurs via the direct 4e\textsuperscript{-} pathway. This is also supported by the low H\textsubscript{2}O\textsubscript{2} production measured for FeNC-dry-0.5 at low loading (Figure 2b).

At this stage, we however do not know yet whether Fe particles present in other Fe-N-C catalysts are PRR active. If such particles are PRR inactive, the PRR activity should be linearly correlated with the absolute FeN\textsubscript{C}, content. However, no linear correlation between FeN\textsubscript{C}, content and PRR activity is observed (Figure 2d, center). This suggests that FeN\textsubscript{C}, sites are not the sole active site for PRR but that Fe particles should also be considered. Some correlation is observed between PRR activity and content of Fe particles (Figure 2d, right), but a better correlation is observed between PRR activity and the total Fe content (Figure 2d, left). Significant PRR activity of FePhen@MOF-ArNH\textsubscript{2} also demonstrates that Fe particles are PRR active, with an activity commensurate to those of FeN\textsubscript{C}, moieties (Figure 2c,d).

While Fe particles are PRR-active, it is still unclear whether they need to be directly exposed to the electrolyte for catalyzing the PRR. Fe particles in direct contact with electrolyte have previously been suggested to be a site S2.\textsuperscript{[12a,16]} Recently, we showed that electrolyte-exposed Fe particles can survive in acidic environments at > 0.7 V\textsubscript{RHE}, which is due to stabilization of ferric hydroxide species.\textsuperscript{[13]} The presence of Fe particles exposed to acidic electrolyte during short RDE studies is thus possible.

To elucidate whether Fe particles must be directly exposed to the electrolyte to catalyze the PRR, the PRR activity of FeNC-wet-1 and FeNC-dry-1 was measured before and after dissolution (electrochemical dissolution\textsuperscript{[10b]} of electrolyte-exposed particles (Figure 3). Operando analysis using a scanning flow cell (SFC) connected with ICP-MS (Supporting Information, Figure S5) showed that a significant fraction of electrolyte-exposed Fe particles in FeNC-wet-1 were removed during 100 cyclic voltammograms (CV) between 0 and 1 V\textsubscript{RHE} (Figure 3a). The dissolved Fe reaches 9% of the total content of Fe particles present in pristine FeNC-wet-1. FeN\textsubscript{C}, moieties are highly stable during this treatment, as previously demonstrated by negligible Fe leaching from FeNC-dry-0.5.\textsuperscript{[13b]}

The PRR activity of FeNC-dry-1 and FeNC-wet-1 were however unmodified, even after extended dissolution up to 1000 CVs (Figure 3b and c). An extrapolation predicts that the vast majority of electrolyte-exposed Fe particles are removed after 1000 CVs (Supporting Information, Figure S6). As previously reported,\textsuperscript{[13]} the dissolution of exposed Fe particles did not decrease the ORR activity (Supporting Information, Figure S7). Therefore, it is concluded that electrolyte-exposed Fe particles in pristine FeNC-wet-1 are inactive toward PRR and ORR. It demonstrates that Fe@N-C sites are therefore PRR active and can be a site S2, as previously hypothesized.\textsuperscript{[10a]}

In conclusion, we elucidated the PRR and ORR reactivity of four different structures existing in Fe-N-C catalysts, that is, metal-free N-C groups, electrolyte-exposed Fe particles, FeN\textsubscript{C}, moieties and Fe@N-C. The ORR pathways on Fe-
N-C catalysts could consequently be established (Figure 4). Contrary to previous hypotheses, N-C groups and electrolyte-exposed Fe particles were shown to be inactive toward PRR in acidic medium. We also demonstrated that FeN\(_x\)C\(_y\) moieties and Fe@N-C species are moderately active toward PRR, proving their possible roles in both direct (major path) and indirect 4e\(^–\) ORR pathways. Since the synthesis of Fe-N-C catalysts with only FeN\(_x\)C\(_y\) moieties\(^{[10c,13]}\) or their combination\(^{[3b,d,17]}\) is now controllable, the understanding of the nature of active sites for PRR provided herein offers new insights for the rational design of advanced Fe-N-C catalysts with high selectivity and expectedly improved durability in polymer electrolyte fuel cells.

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Conflict of interest

The authors declare no conflict of interest.

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