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The evolution pathway from iron compounds to Fe$_1$(II)-N$_4$ sites through gas-phase iron during pyrolysis

Jingkun Li$^{1,4,8}$, Li Jiao$^2$, Evan Wegener$^3$, Lynne Larochelle Richard$^4$, Ershuai Liu$^4$, Andrea Zitolo$^5$, Moulay Tahar Sougrati$^1$, Sanjeev Mukerjee$^4$, Zipeng Zhao$^6$, Yu Huang$^{5,7}$, Fan Yang$^6$, Sichen Zhong$^8$, Hui Xu$^8$, A. Jeremy Kropf$^3$, Frédéric Jaouen$^1$, Deborah J. Myers$^{*3}$, and Qingying Jia$^{*4}$

$^1$Institut Charles Gerhardt Montpellier, UMR 5253, CNRS, Université Montpellier, ENSCM, Place Eugène Bataillon, 34095 Montpellier cedex 5, France
$^2$Department of Chemical Engineering, Northeastern University, Boston, Massachusetts, 02115, United States
$^3$Chemical Sciences and Engineering Division, Argonne National Laboratory, Lemont, Illinois, 60439, United States
$^4$Department of Chemistry and Chemical Biology, Northeastern University, Boston, Massachusetts, 02115, United States
$^5$Synchrotron SOLEIL, L’orme des Merisiers, BP 48 Saint Aubin, 91192 Gif-sur-Yvette, France
$^6$Department of Materials Science and Engineering, University of California, Los Angeles, California, 90095, United States
$^7$California NanoSystems Institute (CNSI), University of California, Los Angeles, California, 90095
$^8$Giner, Inc, Newton, Massachusetts, 02466, United States.

*1 is the present address of J.L.; 4 is the previous address.

ABSTRACT: Pyrolysis is indispensable for synthesizing highly active Fe-N-C catalysts for the oxygen reduction reaction (ORR) in acid, but how Fe, N, and C precursors transform to ORR-active sites during pyrolysis remains unclear. This knowledge gap obscures the connections between the input precursors and output products, clouding the pathway toward Fe-N-C catalyst improvement. Herein, we unravel the evolution pathway of precursors to ORR-active catalyst comprised exclusively of single atom Fe$_1$(II)-N$_4$ sites via in-temperature X-ray absorption spectroscopy. The Fe precursor transforms to Fe oxides below 300 °C, and then to tetrahedral Fe$_1$(II)-O$_4$ via a crystal-to-melt-like transformation below 600 °C. The Fe$_1$(II)-O$_4$ releases a single Fe atom that diffuses into the N-doped carbon defect forming Fe$_1$(II)-N$_4$ above 600 °C. This vapor phase single Fe atom transport mechanism is verified by synthesizing Fe$_1$(II)-N$_4$ sites via “non-contact pyrolysis” wherein the Fe precursor is not in physical contact with the N and C precursors during pyrolysis.

INTRODUCTION

Hydrogen fuel cell electric vehicles (FCEVs), based on proton exchange membrane fuel cells (PEMFCs), were first sold commercially in Japan in 2014, with the launch of the Toyota Mirai,$^1$ followed by limited commercialization in Europe, Korea, Canada, and the United States in 2015 by both Toyota and Hyundai. The catalysts used for the oxygen reduction reaction (ORR) in PEMFCs are Pt-alloys.$^2$ While advances have been made in reducing platinum usage in PEMFCs, the cost of automotive fuel cell systems is still prohibitively high, due primarily to the high loading of Pt in the PEMFC cathode.$^{1,2}$ The key to widespread and sustainable commercialization of FCEVs from a cost perspective is the significant reduction of Pt loading$^2$ or the replacement of Pt-alloys with earth-abundant, inexpensive materials (i.e., platinum group metal (PGM)-free). The leading PGM-free catalysts for the ORR in PEMFCs are single transition metal atoms (M=Mn, Fe or Co) embedded in a nitrogen-doped carbon matrix (M-N-C).$^{3-9}$ Recently, Ye et al.$^9$ reported an Fe-N-C catalyst that delivers 570 mW·cm$^{-2}$ in H$_2$/air PEMFCs. Nevertheless, this state-of-the-art performance is insufficient for automotive PEMFCs.$^2$ Further progress in performance necessitates improvements in the turnover frequency and/or in the density of active sites in M-N-C materials, both of which are impeded by the lack of understanding of the formation of the active site(s) in the synthesis of M-N-C catalysts. The current synthesis strategies for M-N-C catalysts consist of combining sources of M, N, and C either in a single compound or in separate compounds and pyrolyzing the compound(s) typically in the 900 to 1100 °C temperature range.$^6-8, 10, 11$ These synthesis strategies primarily rely on an empirical approach to choice of precursor, M content, and pyrolysis settings to increase catalytic activity. The underlying mechanism governing the competitive formation of metal-based active sites and byproducts during pyrolysis has remained elusive. Rational synthesis of improved M-N-C catalysts requires unraveling the current "black box" synthesis approach of input precursors and output products.
The complexity of M-N-C catalyst structures arises largely from the pyrolysis process, which was demonstrated in the 1980s to be essential to producing highly active M-N-C catalysts for the ORR in acidic media. The material resulting from the high temperature pyrolysis, however, is often a mixture of species including N-doped carbon defects or edges, M-N moieties, and inorganic particles of metal, metal oxides, carbides, nitrides, etc. All these species have been proposed to be ORR active in alkaline and even in acidic electrolyte. It thus remains inconclusive which site(s) are responsible for the superior ORR activity of M-N-C catalysts. This issue has been addressed by efforts to gain some control over the final products. By lowering the Fe content in the precursors (<1 wt%), some research groups managed to produce Fe-N-C catalysts with predominantly Fe(II)-N4 (Fe: denotes the single atom configuration without direct Fe-Fe bonds) moieties, as evidenced by Mössbauer and X-ray absorption spectroscopy (XAS). These catalysts exhibit high ORR performance despite their low gravimetric and volumetric density of Fe(II)-N4 sites. Recently, Fe(II)-N4 moieties were visualized in the Fe-N-C catalyst by scanning transmission electron microscopy coupled with electron energy loss spectroscopy. These studies indicate that the Fe(II)-N4 moiety has a high turnover frequency towards the ORR and likely is the major contributor to the ORR activity of pyrolyzed Fe-N-C catalysts. This finding identifies an obvious pathway towards Fe-N-C catalyst advancement via increasing the Fe(II)-N4 site density. However, the electrochemically active Fe(II)-N4 site density saturates at a very low Fe content (<3 wt%), despite substantial efforts to optimize the type and composition of precursors and pyrolysis conditions. The marginal improvement in Fe-N-C catalysts reported in recent years using the trial-and-error synthesis approach calls for a return to the original question: how is the Fe(II)-N4 site formed during pyrolysis? Herein, we directly monitor the stepwise transformation from Fe precursors to Fe(II)-N4 sites during pyrolysis and validate our site formation mechanistic model via “non-contact pyrolysis” where the Fe and N-C precursors are not in physical contact during pyrolysis.

RESULTS AND DISCUSSION

In-temperature XAS at the Fe K-edge was measured on the mixture of Fe, N, and C precursors throughout the pyrolysis and cooling processes and after exposure of the material to air at room temperature after the pyrolysis. We gradually increased the temperature of the samples to 1000 °C and then decreased the temperature to room temperature in a flowing Helium atmosphere, holding at targeted temperatures during XAS measurements. Two different routes were chosen for in-temperature XAS measurements. The first route used a mixture of iron(II) acetate (FeAc2), 1,10-phenanthroline monohydrate, and a Zn-based metal organic framework (MOF), zeolitic imidazolate framework eight (ZIF-8), mixed via ball milling (see Supplementary). These precursors are typical for synthesis of MOF-derived Fe-N-C catalysts and have been shown to generate active Fe-N-C catalysts with predominantly Fe(II)-N4 sites. However, with this route, the thermal evolution of the Fe compound is convoluted with the thermal evolution of the N and C precursors. To deconvolute the effect of pyrolysis on the Fe and C-N constituents, the second route used a mixture of FeCl2·4H2O, as the iron precursor not containing N and C, and a heat-treated N-doped carbon matrix (N-C) as the N and C precursor. Following the procedure developed for pyrolysis of MOF-derived Fe-N-C catalysts, the N-C was synthesized by homogenizing ZIF-8 and 1,10-phenanthroline dry powders via low-energy planetary ball-milling, followed by heat treatment in Ar at 1050 °C for one hour and a second heat treatment in NH3 at 900 °C for 15 minutes. Multiple characterizations (Figure S1) showed that the N-C has a porous structure with a Brunauer-Emmett-Teller (BET) surface area of 950 m2·g−1, of which 770 m2·g−1 is contributed from micropores according to the t-plot analysis (Figure S1E). X-ray photoelectron spectroscopy (XPS) indicated 0.46 at% of residual Zn, 2.86 at% of O, 6.51 at% of N, and, notably, an ultra-high content of pyridinic N, as manifested by the prominent peak at 398.3 eV (Figure S1D). The simultaneous presence of abundant micropores and pyridinic N preferentially located at the defects and edges of the carbon matrix signifies the presence of abundant nitrogen-doped microporous defects.

XAS of the FeCl2·4H2O and N-C mixture. As compared to the as-received FeCl2·4H2O, the FeCl2·4H2O mixed with N-C via ball-milling exhibits a much lower peak intensity of the X-ray absorption near edge structure (XANES) spectrum at room temperature (Figure 1A). Meanwhile, the intensity of the Fe-O (O from H2O) scattering, with a peak at approximately 1.5 Å in the Fourier transform of the extended X-ray absorption fine structure (FT-EXAFS) scattering, note that the distances in the FT-EXAFS are not corrected for phase shift, drops dramatically; whereas the Fe-Cl scattering peak at approximately 2.1 Å remains largely unchanged (Figure 1B). These results indicate that ligating water molecules are partially detached upon ball-milling. As the temperature increases to 300 °C (denoted as FeCl2·NC-300), the Fe-Cl scattering peak disappears, and new peaks emerge. The Fe-O peak at 1.4 Å and the newly-emerging Fe-Fe peaks at 2.7 Å and 4.6 Å overlap those of the α-Fe2O3 standard (Figure 1B). Meanwhile, the edge energy increases and the XANES spectrum approaches that of α-Fe2O3 (Figure 1A). These results indicate that ferric oxides are formed at 300 °C. As the temperature gradually increases to 600 °C, the XANES shifts to lower energy approaching that of the Fe(II)O standard (Figure 1A), indicating the reduction of ferric oxides to ferrous oxides. The intensities of both the Fe-O and Fe-Fe peaks drop, with the Fe-Fe peaks eventually disappearing at 600 °C, leaving behind one prominent peak at 1.5 Å (Figure 1B). The peak can be fit well with an Fe-O path with a bond distance of 1.99 ± 0.01 Å and a coordination number of 3.6±0.4 (Table S2 and Figure S2). This Fe-O configuration was previously reported, in molten Fe salts, to be a tetrahedral Fe(II)-O4 moiety with an Fe-O bond distance of 1.98±0.02 Å. The tetrahedral structure of the Fe(II)-O4 species observed at 600 °C is supported by the high intensity of the 1s → 3d transition feature of XANES at 7112 eV, as this feature is minimal for centrosymmetric Fe species, such as the plane Fe(II)-N4 in iron phthalocyanine (FePc), and is intense in non-centrosymmetric Fe species, especially for the tetrahedral geometry (Figure S3). The observed transition from ferric oxides, with a local structure of octahedral Fe-O6 with Fe-Fe bonds, to tetrahedral Fe(II)-O4 without Fe-Fe bonds, is akin to the crystal-to-melt transition of Fe.
salts to Fe(II)-O₄ upon melting above 1300 °C.23 Here the Fe(II)-O₄ is not in a molten state since it is observed at much lower temperature. Moreover, Fe(II)-O₄ is unstable by itself owing to the unbalanced cationic and anionic charge. It is rather stabilized by N-C in a solid state via electron delocalization that restores the charge balance, such as is observed for Fe(II)-N₄ sites imbedded in a graphene plane.16

The in-plane Fe(II)-N₄ moieties emerge at approximately 600 °C. This is signified by the appearance of a peak at 7117 eV in the first derivative of the XANES spectra, overlapping that of the Fe(II)-Pc standard (Figure 1C). This peak arises from the 1s → 4p transition with simultaneous ligand to metal charge transfer, and is the fingerprint of the in-plane Fe(II)-N₄ structure.12, 13 As the temperature increases to 1000 °C, the XANES spectrum at energies above 7117 eV gradually shifts to higher energies, approaching that of Fe(II)Pc (Figure 1A), indicating the transformation from the tetrahedral Fe(II)-O₄ to in-plane Fe(II)-N₄. This shift continues rather than reversed during cooling to room temperature, which indicates that the Fe(II)-O₄ → Fe(II)-N₄ transformation is an irreversible thermal process. After cooling to room temperature, but before exposure to air, the Fe speciation (FeCl₂-NC-End) is dominated by in-plane Fe(II)-N₄ moieties, as verified by both the XANES and EXAFS spectra. The in-plane Fe(II)-N₄ fingerprint shoulder at 7117 eV becomes prominent in the first derivative of the XANES (Figure 1C) and even discernible in the XANES spectrum of FeCl₂-NC-End (Figure 1A). The EXAFS fitting (Figure S6) gives a first shell Fe-N coordination number of 4.3±0.4 and bond length of 1.91±0.01 Å, close to that of Fe(II)Pc (1.93±0.01 Å).20 This Fe-N bond distance is much shorter than that of Fe-O (1.99±0.01 Å) as expected from the in-plane structure versus the tetrahedral structure.

Figure 1. (A) XANES, (B) Fourier Transform (FT)-EXAFS and (C) the first derivative of the XANES spectra of FeCl₂-NC-T collected with temperature increasing from room temperature to 1000 °C and cooling down to room temperature. (D) ⁵⁷Fe Mössbauer spectrum (5 K) and fitting of FeCl₂-NC-Air. (E) ORR performance of FeCl₂-NC-T catalysts. RDE polarization plots were obtained in room temperature O₂-saturated 0.1 M HClO₄ from 0.05 to 1.05 V with a scan rate of 1 mV s⁻¹ at a rotation rate of 1600 rpm. The corresponding cyclic voltammograms are displayed in Figure S4.

Upon exposure to air (FeCl₂-NC-Air), the XANES spectrum shifts positively and the fingerprint shoulder at 7117 eV becomes barely discernible (Figure 1A), indicating the oxidation of the surface Fe(II)-N₄ via adsorption of an O₂ or OH ligand, forming Fe₃(III)-N₄O₂/2OH sites. Low temperature (5 K) ex situ Mössbauer identifies two doublets D1 and D2 in FeCl₂-NC-Air (Figure 1D and Table S3). D1 and D2 are the most common doublets identified in Fe-N-C materials and have recently been assigned to Fe₃(III)-N₄O₂ and Fe₃(II)-N₄, respectively.10, 24 These combined results conclusively confirm the formation of gas-phase accessible in-plane Fe(II)-N₄ upon pyrolysis at temperatures >600 °C.

In parallel, X-ray diffraction (XRD) measurements were conducted on the same mixture after it was subjected to pyrolysis at various temperatures. No Fe-related signals are observed throughout the temperature range up to 1000 °C (Figure S7). This result indicates the lack of long-range order in all the Fe species, including the oxides observed by XAS, consistent with the lack of prominent FT-EXAFS peaks at high R values (Figure 1B). The ORR activities of these mixtures were assessed using a rotating disk electrode (RDE) in an O₂-saturated 0.1 M HClO₄ electrolyte. The ORR activities of the catalysts pyrolyzed at 200 °C and 400 °C are similar to that of N-C (Figure 1E). The ORR activity improves dramatically as the pyrolysis temperature increases to 600 °C, coinciding with the emergence of Fe(II)-N₄ sites. The activity continues to improve with increasing temperature up to 1000 °C, reaching a half-wave potential of 0.8 V (all the potentials reported here are versus the reversible hydrogen electrode and are not IR-corrected) (Figure 1E), which coincides with the increasing transformation from Fe₃(II)-O₄ to Fe₃(II)-N₄. Not only is the RDE-determined activity comparable to that of state-of-the-art Fe-N-C,3, 6, 25 but the intimate correlation between the increasing ORR activity and the increasing relative content of Fe₃(II)-N₄ in the sample provides startling evidence that the Fe(II)-N₄ moiety is responsible for the high ORR activity of Fe-N-C in acidic electrolyte.

Nearly the same thermal evolution process was observed in the in-temperature XAS experiment on the MOF-based mixture: FeAc₂ → ferric oxides with a local octahedral Fe-O₆ structure → ferrous oxides → tetrahedral Fe(II)-O₄ → in-plane Fe(II)-N₄ (Supplementary Section 3). The two different but representative groups of Fe, N, and C precursors undergoing the same thermal evolution pathway suggests that this is likely a common pathway towards the formation of Fe(II)-N₄ during the pyrolysis of mixtures of Fe, N, and C precursors. This pathway is, however, missing details regarding the mechanism for transformation of tetrahedral Fe(II)-O₄ to in-plane Fe(II)-N₄. The question remains as to how the Fe(II)-O₄ transforms to a moiety with a very
different local geometry and ligand environment. An alternative transformation from tetrahedral Fe-N₄ to active FeN₄ was recently proposed to occur at temperatures ≥ 800 °C during the pyrolysis of Fe-doped ZIFs to form Fe-N-C catalysts. However, EXAFS cannot distinguish the Fe-O₃ structure from Fe-N₄ since it cannot distinguish O from N as the scattering neighbor.

To address this uncertainty, we conducted the in-temperature XAS on the mixture of FeCl₂·4H₂O and nanoscale SiO₂ powders in an environment free of N and C. The SiO₂ was deliberately chosen to dilute FeCl₂·4H₂O for XAS measurements in the transmission mode, as it can also serve as a support with high thermal stability as an alternative to N-C. The thermal evolution involving the decomposition of FeCl₂·4H₂O to form a species with one prominent FT peak at approximately 1.4 Å at 600 °C was again observed (Figure 2A). The fitting of the EXAFS spectrum at 600 °C gives an Fe-O coordination number of 3.9 ± 0.6 and bond length of 2.00 ± 0.02 Å (Table S5), thus conclusively confirming the formation of tetrahedral Fe(II)-O₄ rather than Fe(II)-N₄ or Fe(II)-O₃. More importantly, this XAS spectrum at 600 °C (FeCl₂-SiO₂-600) nearly overlaps that of the FeCl₂-NC-600, with trivial differences arising from a small fraction of Fe(II)-O₄ being already converted to Fe(II)-N₄ in FeCl₂-NC-600 (Figure 2A and Figure S12). This result verifies the formation of Fe(II)-O₄ upon the pyrolysis of FeCl₂·4H₂O mixed with either N-C or SiO₂ and, by inference, pyrolysis of the FeA₄-ZIF-B₈ mixture.

As the temperature of the FeCl₂-SiO₂ mixture increases from 600 °C to 1000 °C, in the absence of N-C the Fe(II)-O₄ partly transforms to ferrous oxides and then Fe nanoparticles, rather than Fe(II)-N₄ (Figure 2A) (Supplementary Section 4). The Fe(II)-O₄ and Fe nanoparticles are present throughout the process of cooling to room temperature, as seen in Figure 2B, wherein the growth of the FT-EXAFS peaks is mainly attributed to the decreasing Debye-Waller factor with decreasing temperature. The XANES spectra remain largely unchanged during cooling (Figure S11B). The presence of Fe(II)-O₄ at room temperature verifies that it is a stable species when anchored onto a substrate. The observed thermal evolution pathway of the FeCl₂-SiO₂ mixture may be representative of the typical pathway for the formation of single atom catalysts supported on oxides.

Collectively, the thermal evolution of the Fe precursors in the three mixtures converges to the formation of tetrahedral Fe(II)-O₄ at 600 °C, and then diverges at higher temperatures with/without N-C (Figure 2C). This thermal decomposition generally follows the Fe-O phase diagram that depicts the transformation of ferric oxides to ferrous oxides around 300 °C in an oxygen-deficient gaseous environment and then to Fe nanoparticles around 600 °C with all the oxygen neighbors removed. A natural hypothesis deduced from the Fe-O phase diagram is that complete removal of the oxygen neighbors from Fe(II)-O₄ leaves behind a single free Fe atom (Fe₁), which may form Fe₁(II)-N₄ if captured by the N-C defect nearby: Fe₁(II)-O₄ → Fe₁ → Fe₁(II)-N₄, or sinter into aggregated Fe clusters in the absence of N-C defects (Figure 2C).

A unique aspect of this so-called impregnation mechanism lies in the formation of Fe₁. The Fe₄ is essentially a gas phase iron that expands to fill the volume available, from which we deduce that it is not necessary to mix the Fe precursors with N and C during pyrolysis to produce active Fe-N-C catalysts, as previously asserted, since the Fe₁ released from the Fe precursor can diffuse or otherwise transported into N-C defects to form Fe₁(II)-N₄. To test this hypothesis, we developed a so-called non-contact pyrolysis procedure wherein the Fe precursor and N-C are separately placed in two boats in the furnace without direct physical contact.
between the two materials during pyrolysis (Figure 3A). In addition to FeAc₂ and FeCl₂·4H₂O, that were shown above to transform to Fe₃(II)·Nₓ sites when mixed with N and C precursors, α-Fe₂O₃ was also investigated as the Fe precursor since ferric oxides were observed at intermediate temperatures during both synthesis routes via in-temperature XAS.

![Diagram](image)

**Figure 3.** (A) Schematic illustration of the non-contact pyrolysis wherein the Fe precursor and N-C were separately placed in two boats in the furnace during pyrolysis with Fe placed upstream of the gas flow. (B) ORR performance of indicated catalysts. RDE polarization plots were obtained in room-temperature O₂-saturated 0.1 M HClO₄ from 0.05 to 1.0 V with a scan rate of 1 mV·s⁻¹ at a rotation rate of 1600 rpm. Solid lines represent catalysts obtained by non-contact pyrolysis; dashed line catalysts obtained by regular pyrolysis for comparison. (C) H₂-O₂ and (D) H₂-air fuel cell polarization curves. Cathode: ~4.0 mg·cm⁻² of FeCl₃/NC-1000; Anode: 0.3 mg·cm⁻² Pt/C; Membrane: NR212 membrane (Ion Power); 200 and 1000 mL·min⁻¹ gas fed with 100% RH at anode (H₂) and cathode (O₂/air), respectively; electrode area 5 cm². The back pressures during the fuel cell tests are 1 bar reactant gas. (E) XANES and (F) FT-EXAFS spectra of the three catalysts obtained by non-contact pyrolysis, and of the Fe foil and FePC-O₂. (G) TGA of the three Fe precursors including FeCl₃·4H₂O, FeAc₂, and α-Fe₂O₃.

The ORR performance of the N-C subjected to the non-contact pyrolysis with FeCl₂·4H₂O at 1000 °C (denoted as FeCl₂/NC-1000) is essentially the same as that of the mixture of FeCl₂·4H₂O and N-C pyrolyzed at 1000 °C (FeCl₂-NC-1000) (Figure 3B). This catalyst also exhibits reasonably high performance in a PEMFC membrane-electrode assembly, delivering an iR-corrected current densities of 105 mA·cm⁻² and 400 mA·cm⁻² at 0.8 V and 0.7 V, respectively, in H₂-O₂ (Figure 3C), and a maximum power density of 0.28 W·cm⁻² in H₂-air (Figure 3D). This relatively high performance is likely related to an enrichment of surface active sites, as expected from the vapor deposition feature of the non-contact pyrolysis. This is the first demonstration of a highly active Fe-N-C PEMFC cathode catalyst that is synthesized without mixing the Fe precursor with N and C precursors. Driven by the proof-of-concept, optimizations of the precursors, non-contact pyrolysis, and the corresponding PEMFC testing are underway in our groups.

The formation of Fe₃(II)·Nₓ in the non-contact pyrolysis material, as reflected by the RDE and PEMFC results, is confirmed by the post-pyrolysis XAS and low temperature ⁵⁷Fe Mössbauer measurements. The amount of Fe deposited onto N-C is ~1.5 wt%, as estimated from the edge step of the XANES, and ~1.8 wt% by X-ray fluorescence (XRF). The XANES aligns well with the oxygen adduct of Iron(III) phthalocyanine-tetrasulfonic acid (FePc-O₂⁻) (Figure 3E). Correspondingly, the EXAFS shows a prominent Fe-N/O peak around 1.5 Å that nearly overlaps the Fe-N/O scattering peak of FePc-O₂⁻ (Figure 3F). The small Fe-Fe peaks indicate the co-presence of Fe nanoparticles and oxides, which are evident in the XRD pattern (Figure S14). In agreement with the XAS, the ⁵⁷Fe Mössbauer spectrum collected at 5 K identifies four components: D1 (assigned to Fe₃(III)-Nₓ-O₂), another doublet with higher isomer shift assigned to high spin Fe²⁺, γ-Fe, and α-Fe (Figure S15). Excellent fits were obtained without inclusion of a D2 doublet. These results demonstrate that the non-contact pyrolysis produces highly active Fe-N-C catalysts with predominately Fe₃(II)-Nₓ sites and provides support for the proposed gas-phase transport-impregnation mechanism. However, these results do not necessarily indicate that the gas phase Fe species is Fe₂. The anhydrous FeCl₂ has a relatively low boiling point of ~1000 °C and possesses a vapor pressure of ~1 atm at 1000 °C.²⁸ Gas phase FeCl₂ may exist and be transported from the Fe-precursor boat to the N-C boat at high temperatures, forming Fe₃(II)-Nₓ sites once contacting the N-C. When mixed with N-C, the FeCl₂·4H₂O, however, undergoes a completely different evolution pathway via the formation of iron oxides and Fe₂. As illustrated in Figure 2C. Therefore, to verify the existence of gas phase Fe₂, non-contact pyrolysis was also conducted on α-Fe₂O₃ and FeAc₂ to exclude the formation of gas phase iron chlorides.

The ORR activity of the N-C subjected to the non-contact pyrolysis with α-Fe₂O₃ at 1000 °C (Fe₃O₄/N-C-1000) is much higher than that of N-C, lower than that of FeCl₂-NC-600, but has the same ORR onset potential (Figure 3B). These results suggest the presence of Fe₃(II)-Nₓ sites as in FeCl₂-NC-600, but with lower site density. Indeed, the XANES and EXAFS of Fe₃O₄/N-C-1000 nicely overlap with those of FeCl₂/N-C-1000 and FePc-O₂⁻ (Figure 3E and 3F), confirming the formation of Fe₃(II)-Nₓ. The very small XANES edge step for this material indicates that only small amounts of Fe were deposited on N-C, estimated at approximately 0.03 wt% by XRF. This is expected from the high thermal stability of α-Fe₂O₃, with a decomposition temperature higher than 1000 °C, as seen by the thermogravimetric analysis (TGA) results (Figure 3G). This validation of the non-contact pyrolysis synthesis using α-Fe₂O₃ further support the gas phase transport-impregnation mechanism.
They also indicate that ionic compounds with stable crystalline structures are not ideal Fe precursors for Fe-N-C synthesis due to the high energies needed to break the stable Fe-Fe bonds to release Fe. It was shown that metal (Pd or Ni) nanoparticles can release single free metal atoms that form metal-Ns sites upon impregnation into N-C defects, but this only occurs at >1000 °C as the competitive sintering process dominates at lower temperatures.29-31

The thermal stability of the ferric oxides generated in situ from FeCl3·4H2O or FeAc2·4H2O mixed with N-C is much lower than that of bulk α-Fe2O3, as it is readily reduced to Fe(II)-O4 with its Fe-Fe bonds cleaved in the 300 °C to 700 °C temperature range (Figure 1B and Figure S8B). Note that the ferric oxides generated in situ are invisible to XRD (Figure S7), which indicates that the clusters are ultra-small and/or amorphous. We thus deduce that the N-C disturbs the long-range crystalline structure of Fe oxides thereby weakening the Fe-Fe bonds, and/or traps small clusters of Fe oxides, which have a reduced number of Fe-Fe bonds as compared to α-Fe2O3. This promotes the transformation from Fe2O3 with fewer and weakened Fe-Fe bonds, to Fe(II)-O4. In addition, the stabilization of Fe(II)-O4 in the solid state by the N-C or SiO2 substrates further promotes the crystal-to-melting transformation of Fe-O4 → Fe(II)-O4 at much lower temperatures. Without N-C, the α-Fe2O3 with stable crystalline structures releases few Fe atoms and thus forms fewer Fe(II)-N4 sites at 1000 °C.

Surprisingly, the ORR performance of the N-C subjected to the non-contact pyrolysis with FeAc2 at 1000 °C is only slightly better than that of N-C, and much worse than that of Fe2O3/N-NC-1000 (Figure 3B), indicating the absence of Fe(II)-N4. Both XAS (Figure 3E and 3F) and XRD (Figure S14) of the N-C after the non-contact pyrolysis show Fe nanoparticles and oxides as the predominant Fe species. The Fe content of this material is approximately 0.5 wt%, as estimated from both the XANES edge step and XRF. These results indicate that Fe atoms were released from FeAc2 and transported onto the N-C, forming Fe clusters rather than Fe(II)-N4. One key difference between FeAc2 and α-Fe2O3 is that the former decomposes at ~300 °C, whereas the latter decomposes above 1000 °C (Figure 3G). At 300 °C, sintering of Fe into aggregated clusters dominates,29 due to the short mean free path of Fe at this relatively low temperature, leading to the formation of Fe nanoparticles. In addition, 300 °C is below the temperature (~660 °C) of the transform from Fe(II)-O4 to Fe(II)-N4, and thus even if Fe atoms reach the N-C substrate, they tend to form iron oxides rather than Fe(II)-N4. These issues can be addressed by thorough mixing of FeAc2 with N and C precursors prior to pyrolysis. In the mixture, the N-doped defects are in close proximity to the Fe sources. It can be hypothesized that N-C stabilizes Fe(II)-O4 as the reservoir of Fe through a wide temperature range of 400-1000 °C (Figure S8). This promotes the formation of Fe(II)-N4, despite the short mean free path of Fe. In comparison with FeAc2, α-Fe2O3 releases much fewer Fe atoms at much higher temperature (~1000 °C), and thus the mean free path of Fe is longer. Consequently, these Fe atoms can reach the N-C substrate and form Fe(II)-N4. Similar to this proposed mechanism, Li’s group recently observed the transformation of bulk Pd to Pd-N4 at elevated temperature and accordingly proposed that this conversion was driven by the capture of mobile Pd atoms on the defects of N-C.29

An important implication of the non-contact pyrolysis results is that the threshold temperature for the formation of Fe2(II)-N4 of ~600 °C is determined by the threshold temperature for the impregnation of Fe into the N-C defect (Fe → Fe(II)-N4), rather than the release temperature of Fe. That is, it is determined by the inherent thermal stability of Fe(II)-N4. The Fe(II)-O4 has a lower thermal stability as it forms and decomposes at lower temperatures. This difference in thermal stability between Fe2(II)-N4 and Fe(II)-O4 accounts for the competition between the two species during pyrolysis above 600 °C. The Fe2(II)-N4 gradually wins the competition as the temperature reaches 1000 °C. This fundamental limitation determines the necessity for multiple pyrolyses with an optimized temperature of ~1000 °C to drive the Fe2(II)-O4 → Fe2(II)-N4 transformation.

CONCLUSION

We unraveled the thermal evolution pathway during the pyrolysis of Fe-N-C catalysts: Fe precursor → Fe oxides (octahedral Fe-O4) → tetrahedral Fe2(II)-O4 → Fe → Fe2(II)-N4. The demonstration of formation of Fe2(II)-N4 via non-contact pyrolysis (i.e., with the Fe precursor and nitrogen-doped carbon in separate boats) reveals the presence of gas-phase iron at mild temperatures and opens up an avenue for the synthesis of single-atom catalysts via vapor deposition approaches.

ASSOCIATED CONTENT

Supporting Information

Supporting information (PDF) includes: Experimental Section; Results and Discussion: Section 1. N-C Characterizations (Figure S1); Section 2. In-temperature XAS on the mixture of FeCl3·4H2O and N-C (Figure S2-S7 and Table S2-S3); Section 3. In-temperature XAS on the mixture of MOF-based mixture (Figure S8-S10 and Table S4); Section 4. In-temperature XAS on the mixture of FeCl3·4H2O and SiO2 (Figure S11-S13 and Table S5); Section 5. Non-contact pyrolysis of three different Fe precursors (Figure S14). The Mössbauer spectrum of FeCl3/N-C-1000 measured at 5 K (Figure S15). This material is available free of charge via the Internet at http://pubs.acs.org.

AUTHOR INFORMATION

Corresponding Author

* Deborah J. Myers: dmyers@anl.gov
* Qingying Jia: q.jia@northeastern.edu

Author Contributions

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Notes

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