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Selective Oxidation of C–H Bonds through a Manganese(III) Hydroperoxo in MnII-Exchanged CFA-1

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ABSTRACT: Partial substitution of ZnII by MnII in Zn₅(OAc)₄(5,5′-bibenza[d][1,2,3]triazole)₃ (CFA-1) results in a MnIII species supported by three nitrogen ligands and a charge-balancing anion, a structure reminiscent of those found in molecular “scorpionate” complexes. Unlike molecular manganese(II) scorpionates, Mn-CFA-1 is capable of catalytically activating oxygen from air to oxidize C–H bonds up to 87 kcal/mol in strength. A series of in situ spectroscopic studies, including diffuse-reflectance UV−vis, diffuse-reflectance infrared Fourier transform spectroscopy, and X-ray absorption spectroscopy, reveal that catalysis likely proceeds through a manganese-(III) hydroperoxo that is only accessed in the presence of a hydrogen-atom donor. These results demonstrate that the site isolation provided in metal–organic frameworks enables the generation and utilization of highly reactive species for catalysis that are inaccessible in molecular systems.

INTRODUCTION

Selective oxidations are key steps in the transformation of both petroleum and biofeedstocks into commodity chemicals and ultimately into consumer products. Often, oxidations are achieved with the intermediacy of reactive metal oxo species, which typically suffer either from indiscriminate reactivity and a lack of selectivity toward a range of C–H bonds or from bimolecular decomposition.¹−⁴ Manganese is an element of interest for these transformations because of its prevalence in biological transformations involving oxygen-atom transfer and the ease of access to multiple redox states.⁵−⁸ However, manganese-based molecular catalysts are particularly prone to bimolecular decomposition, while most isolated molecular examples of manganese(IV) oxo species display sluggish oxygen-atom-transfer reactivity, at least, in part, because of the strongly donating ligand fields and sterically bulk typically required for their isolation.⁹−¹¹ Metal–organic frameworks (MOFs) confer several advantages for heterogeneous small-molecule chemistry, lending such attributes as ease of product separation and site isolation, coupled with well-defined, tunable catalytic sites.¹²−¹⁵ The combination of these features enables access to unique coordination geometries around metal ions that are otherwise inaccessible in molecular analogues and sometimes lead to unusual reactivity.¹⁶,¹⁷ The carboxylate and azolate binding moieties that often serve as the node-to-linker interface in MOFs offer relatively weak ligand fields, especially compared to those found in the ligands of molecular manganese complexes. These weaker ligand fields combined with site isolation make it possible to access and utilize highly reactive metal–oxygen species.¹⁸

Here, we took inspiration from a family of molecular manganese complexes colloquially known as “scorpionates”, which feature manganese–oxygen species that are primed for reactivity.¹⁹,²⁰ Indeed, several molecular scorpionate-based manganese(III) peroxo species have been isolated, albeit through harsh oxidizing agents such as hydrogen peroxide or potassium superoxide.²¹−²⁵ No catalysis has been reported with these species, presumably either because they are sterically accessible but plagued by instability or because they are relatively stable but sterically inaccessible. We surmised that the incorporation of such species within scorpionate-like MOFs would unlock the reactivity potential that has not yet been achieved with molecular manganese scorpionates. This approach is inspired by earlier work wherein Kuratowski clusters in materials such as Zn₅Cl₄(BTDD)₃ (MFU-4) and Zn₅(OAc)₄(bibta)₃ (CFA-1) serve as excellent platforms for site-isolated scorpionate chemistry.²⁶−³³ The secondary building units (SBUs) in both of these materials offer metal sites supported by three nitrogen ligands and a charge-balancing anion, comprising a ligand field that is far weaker than those typically reported for known molecular manganese(IV) oxo species.³⁵ Altogether, these properties make CFA-1, made from a more accessible ligand than MFU-4, a particularly attractive platform for the isolation of reactive...

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high-valent manganese–oxygen species potentially capable of selective catalytic C–H activation chemistry.

## RESULTS AND DISCUSSION

Mn-CFA-1 (Figure 1) can be accessed via cation exchange at room temperature by gently stirring CFA-1, a crystalline powder, with 50 equiv of MnCl₂·4H₂O in N,N-dimethylformamide (DMF) for 7 days. Thorough washing of the solid material isolated from this reaction with DMF and methanol, followed by activation under dynamic vacuum at 180 °C, yields a crystalline material with a Brunauer–Emmett–Teller (BET) surface area of 1990 m²/g, consistent with that of the CFA-1 starting material (see the Supporting Information). Analysis by inductively coupled plasma optical emission spectrometry (ICP-OES) indicates the incorporation of two manganese atoms per SBU, and the 'H NMR of an acid-digested material revealed the presence of some residual acetate anions.

A comparison of the X-ray absorption near-edge spectroscopy (XANES) features collected on the activated material sealed in a capillary under an inert atmosphere to Mn⁴⁺ reference compounds indicated that the manganese present in Mn-CFA-1 is in the 2⁺ formal oxidation state (Figure 2). Exposure of activated Mn-CFA-1 to air for several hours led to only a slight shift in the edge energy (+0.2 eV) relative to pristine Mn-CFA-1 and is indicative of the coordination of a small amount of water to Mn⁴⁺ rather than oxidation to Mn⁶⁺ or higher. Indeed, formal oxidation would be expected to shift the edge energy by several electronvolts, as evidenced by Mn⁶⁺ standards such as MnO₂ and Mn(OAc)₃ (Figure 2).

The first signs of intriguing reactivity of the manganese scorpionate moieties in Mn-CFA-1 came from stoichiometric oxidations of substrates such as cyclohexene to products including cyclohexen-1-ol, cyclohexen-1-one, and cyclohexene oxide using a variety of oxidants including tert-butylsulfonyl-2-iodosylbenzene and tert-butyl hydroperoxide. The more remarkable reactivity was observed, however, when using air as the terminal oxidant, which led to catalytic and selective formation of alcohol and ketone products from a variety of substrates including ethylbenzene, cyclohexene, cumene, and 1-hexene but not including substrates with stronger C–H bonds such as toluene and cyclohexane. Evidencing the key role of manganese in this reactivity and the heterogeneity of the manganese catalytic site, all-zinc CFA-1 does not enable catalysis under identical conditions, while ICP-OES of a postcatalysis filtrate sample presented no detectable manganese traces in solution. Additionally, powder X-ray diffraction (PXRD) of the solid material postcatalysis indicated good crystallinity that was unchanged from that of the Mn-CFA-1 starting material (Figure S1), eliminating the possibility of framework decomposition.

The heterogeneous nature of the catalysis was further confirmed by a size-exclusion experiment. Thus, Mn-CFA-1 was found to catalyze oxidation of 1,3,5-trisopropylbenzene for a total of 3.3 turnovers per manganese, whereas cumene oxidation was 2.5 times more efficient, with 8.3 turnovers per manganese. It is expected that both of these substrates have comparable benzylic C–H bond strengths; however, cumene at its narrowest width is approximately 4.2 Å, while the narrowest width of 1,3,5-trisopropylbenzene is approximately 7.3 Å. The channels of CFA-1 are only 6.2 Å at their widest aperture. Presumably, whereas cumene is able to diffuse through the pores of the material and undergo oxidation catalyzed by the site-isolated manganese species within, 1,3,5-trisopropylbenzene is likely only able to access manganese species on the surface of the MOF crystals, resulting in more limited conversion. These results are in line with previous observations of size-selective catalysis with MOFs.

Because of the previously mentioned observation of acetate upon acid digestion of Mn-CFA-1, the impact of the charge-balancing anion identity on catalysis was investigated. Attempts at performing cation exchange on CFA-1 with Mn(OAc)₃·4H₂O, which would theoretically yield a material featuring only acetate anions, were unsuccessful. Additionally, attempts at complete anion exchange of acetate for chloride in Mn-CFA-1 resulted in a poorly crystalline material with severely lowered surface area. Fortunately, the pentanuclear zinc cluster that is the SBU of CFA-1 is a motif that can be found in other MOFs.

[Figure 2. XANES data for Mn-MOFs compared with manganese standards.]
previously reported that highly sensitive catalysis, such as ethylene dimerization to selectively yield 1-butene, proceeds identically in both the CFA-1 and MFU-4\textsuperscript{l} platforms;\textsuperscript{27} given their remarkably similar geometries, this was expected to be the case for the oxidation reactions discussed in this work as well. Thus, Mn-MFU-4\textsuperscript{l} was synthesized in a manner similar to that described for Mn-CFA-1 and slightly modified from what has been previously reported.\textsuperscript{29} MFU-4\textsuperscript{l} was gently stirred in a DMF solution containing 50 equiv of MnCl\textsubscript{2}-4H\textsubscript{2}O for 7 days. The solid material isolated from this reaction was washed thoroughly with DMF and methanol and then activated under a high vacuum at 180 °C until a pressure of 10\textsuperscript{−5} Torr was achieved. This procedure yielded a high-quality material with good crystallinity and a BET surface area of 3570 m\textsuperscript{2}/g, consistent with that of the MFU-4\textsuperscript{l} starting material and with the value previously reported for Mn-MFU-4\textsuperscript{l}. ICP-OES analysis indicated the incorporation of 1.8 manganese atoms per SBU, and the \textsuperscript{1}H NMR of the acid-digested material

| catalyst     | oxidant | time  | alcohol\textsuperscript{a} | ketone\textsuperscript{a} | sum of the alcohol and ketone products | ratio of alcohol/ketone |
|--------------|---------|-------|-----------------------------|-----------------------------|---------------------------------------|------------------------|
| Mn-CFA-1     | air     | 15 min| 2.8                         | 3.2                         | 6                                     | 1:1.1                  |
| Mn-CFA-1     | air     | 1 h   | 3.8                         | 5.0                         | 8.8                                   | 1:1.3                  |
| Mn-CFA-1     | air     | 6 h   | 3.8                         | 6.9                         | 10.7                                  | 1:1.8                  |
| Mn-CFA-1     | O\textsubscript{2} (2 bar) | 18 h  | 5.4                         | 15.8                        | 21.2                                  | 1:2.9                  |
| Mn-MFU-4\textsuperscript{l} | air     | 18 h  | 0.9                         | 0.5                         | 1.4                                   | 1:0.55                 |
| Mn(CO\textsubscript{2}CH\textsubscript{3})-MFU-4\textsuperscript{l} | air     | 18 h  | 4.7                         | 14.7                        | 19.4                                  | 1:1.3                  |
| Mn(CO\textsubscript{2}CF\textsubscript{3})-MFU-4\textsuperscript{l} | air     | 18 h  | 5.5                         | 11.1                        | 16.6                                  | 1:2.0                  |

\textsuperscript{a}Turnovers of cyclohexen-1-ol calculated with respect to manganese. 

\textsuperscript{b}Turnovers of cyclohexen-1-one calculated with respect to manganese.

**Scheme 1. Proposed Catalytic Cycle for Oxidation of Cyclohexene by Mn-CFA-1 in Air**

The solid material isolated from this reaction was washed thoroughly with DMF and methanol and then activated under a high vacuum at 180 °C until a pressure of 10\textsuperscript{−5} Torr was achieved. This procedure yielded a high-quality material with good crystallinity and a BET surface area of 3570 m\textsuperscript{2}/g, consistent with that of the MFU-4\textsuperscript{l} starting material and with the value previously reported for Mn-MFU-4\textsuperscript{l}. ICP-OES analysis indicated the incorporation of 1.8 manganese atoms per SBU, and the \textsuperscript{1}H NMR of the acid-digested material
confirmed that, indeed, no acetate anions were present. Extended X-ray absorption fine structure (EXAFS) analysis was performed on this material and yielded a fit consistent with the substitution of manganese for zinc into the MOF framework, complete with a charge-balancing chloride anion. Details of this analysis can be found in the Supporting Information.

Tellingly, for the influence of the anion on reactivity, Mn-MFU-4l that is not ever exposed to acetate, does not catalyze the oxidation of cyclohexene with air, with the reactivity observed being merely stoichiometric (Table 1). Subjecting Mn-MFU-4l to a sodium acetate solution yields Mn-(CO$_2$CF$_3$)-MFU-4l, a material with good crystallinity where 2.7 chlorides per SBU are replaced by acetates (see the Supporting Information). This acetate-exchanged Mn-MFU-4l does enable catalytic oxidation of cyclohexene in the presence of air, conclusively demonstrating the critical role of acetate for oxidation activity by manganese in both Mn-CFA-1 and Mn-MFU-4l. Adding a large excess of sodium acetate to Mn-CFA-1 did not significantly increase the number of turnovers for cyclohexene oxidation, suggesting that acetate is not consumed in the reaction, but rather facilitated it as a ligand on the manganese. Importantly, replacing only 0.9 of the chloride anions in each SBU of MFU-4l by trifluoroacetate produces a material, Mn(CO$_2$CF$_3$)-MFU-4l (see the Supporting Information), that is as active as Mn-MFU-4l with a more significant acetate content, underscoring the importance of weak ligand-field anions in promoting oxidation catalysis with manganese-substituted Kuratowski clusters.

Visual inspection of the reaction progress provides first clues to the potential mechanism of O$_2$ activation and reactivity with the manganese-exchanged MOFs. Thus, when substrates with sufficiently weak C–H bonds are added to beige Mn-CFA-1 in air, the MOF turns black upon contact and then rapidly lightens to dark brown over the course of 5 min; as the reaction continues overnight, the material returns to its original color. This series of color changes is not observed when the substrate is introduced to Mn-CFA-1 in the absence of oxygen, nor is it ever observed when the substrate has C–H bonds that are too strong to be oxidized, such as toluene. These visual observations combined with the decline in catalyst performance over time suggest that a highly reactive species forms initially only in the presence of both O$_2$ and substrate and that this species is responsible for the majority of the observed reactivity, before disappearing from the reaction mixture. A plausible scenario is one where, in the presence of a suitable hydrogen-atom donor, manganese(III) hydroperoxo is formed and kicks off the catalytic cycle shown in Scheme 1. Although rare, the few manganese(III) hydroperoxo species reported in the literature are accessed in a similar manner: a Mn$^{II}$ complex is oxidized by molecular oxygen in the presence of a hydrogen-atom donor to yield manganese(III) hydroperoxo, presumably via a very transient manganese(III) superoxo. In support of this hypothesis, the addition of only 10 equiv of benzoquinone as a scavenger, results in a more than 60% reduction in the oxidized product produced by Mn-CFA-1 in air. Although the proposed manganese(III) hydroperoxo was not isolable in our hands, we assign its relative instability to the comparitively weaker ligand field conferred by Mn-CFA-1 relative to those in reported manganese(III) hydroperoxo molecular species. Notably, the same reasoning explains the increased reactivity of the putative manganese(III) superoxo here, which is able to oxidize stronger C–H bonds than previously seen with molecular systems.

In line with our qualitative visual observations, in situ spectroscopic experiments confirmed oxidation of Mn$^{II}$ in Mn-CFA-1 as well as changes in its electronic structure upon exposure to cyclohexene in air. Although Mn$^{II}$-CFA-1 in air has no strong absorptions in diffuse-reflectance (DR) UV–vis (Figure 3a), new absorptions at 439 and 650 nm appear upon the addition of 1 drop of cyclohexene. These bands are not associated with pure cyclohexene itself and likely indicate the formation of Mn$^{III}$, wherein d–d transitions become spin-forbidden.

In situ XANES measurements provided more conclusive evidence for the formation of Mn$^{III}$ upon exposure of Mn-CFA-1 to both cyclohexene and air. Thus, whereas exposing Mn-CFA-1 to a flow of simulated air (20% O$_2$ in N$_2$) produced no visible changes in the XAS spectrum, adding cyclohexene to the in situ gas flow caused an abrupt change in the edge energy and line shape (Figure 3b). More specifically, a shift in the edge energy of +0.5 eV and a shift of the preedge feature by +0.3 eV were consistent with the oxidation of manganese. Although not as dramatic as those observed for molecular manganese scorpionates undergoing oxidation from Mn$^{II}$ to Mn$^{III}$, we note that the Mn$^{II}$ in our system is likely not a dominant species when the system reaches the steady state. Indeed, in light of the catalytic cycle proposed in Scheme 1, we

![Figure 3](image-url)

**Figure 3.** (a) DR UV–vis and (b) XANES data of Mn-CFA-1 in air, before and after exposure to cyclohexene.
do not anticipate that 100% of the manganese in the sample would be oxidized from MnII at any given point in time. Of even greater interest is the decrease in the intensity of the preedge feature at 6540.3 eV upon exposure to cyclohexene. The preedge features of Mn-XANES spectra are closely related to the symmetry at the manganese site and here suggest that the manganese site reaches a more centrosymmetric environment upon exposure to air and cyclohexene, resulting in a less intense preedge feature. This would be in line with the manganese atom gaining another ligand (such as the hydroperoxo).

Additional evidence for the formation of manganese(III) hydroperoxo came from in situ diffuse-reflectance infrared Fourier transform spectroscopy (DRIFTS; Figure 4). Exposing Mn-CFA-1 to a 10 s “pulse” of O2 flowing through a saturator filled with cyclohexene caused the appearance of a band at 903 cm⁻¹ over the course of 20 min, which then gradually disappeared. This frequency is in the region of the O–O bond stretch of the putative hydroperoxo species, as numerous manganese(III) side-on peroxy, manganese(III) hydroperoxo, and manganese(III) alkylperoxo species have been reported in the same region. One other band of interest that exhibits the same transient behavior as that at 903 cm⁻¹ (that is, it grows in and then disappears over the course of approximately 20 min) is 827 cm⁻¹. Intriguingly, this frequency is most in line with those of terminal manganese(IV) oxo species. Although such species are extremely rare, one reported instance for the Mn–O stretching frequency in a manganese(IV) oxo places it at 845 cm⁻¹.

Although the Mn-CFA-1 and Mn-MFU-4 systems allow the identification of rare, transient manganese(III) hydroperoxo and potentially of a manganese(IV) oxo species with notable reactivity toward C–H bonds, the catalytic activity of these high-valent manganese species is quickly quenched by water formed during the reaction. The appearance of water is readily discernible through in situ DRIFTS experiments (Figure S5). Even increasing the availability of the oxidant by performing the reaction under 2 bar of O2 and increasing the duration to 18 h did not result in more than a marginal increase in the catalytic performance (Table 1). This is also consistent with the only transient appearance of spectroscopic signatures of MnIII-OOH and manganese(IV) oxo that are not discernible beyond the first 30 min of the reaction. Indeed, it is highly likely that the proposed catalytic cycle is only in effect initially, and further turnovers achieved after the inaugural few rounds of catalysis are instead the result of radical processes. Indeed, the addition of 10 equiv of 2,4,6-tri-tert-butylphenol, a free-radical scavenger that is too large to diffuse into the pore of Mn-CFA-1, to the reaction inhibits conversion of cyclohexene to oxidized products and, interestingly, keeps the product distribution closer to 1:1 for reactions run for longer than 1 h (Table 2).

Figure 4. In situ DRIFTS of Mn-CFA-1 after exposure to O2 and cyclohexene. Spectra were collected over the course of 15 min; a feature at 903 cm⁻¹ grows in (a) and then diminishes (b), as does a feature at 827 cm⁻¹, which also grows in (c) and then diminishes (d).
Table 2. Effect of the Free-Radical Inhibitor 2,4,6-Tri-tert-butylphenol and Reaction Time on the Total Catalyst TON and Selectivity<sup>a</sup>

| catalyst | oxidant | addition of a free-radical inhibitor<sup>b</sup> | time | alcohol<sup>c</sup> | ketone<sup>d</sup> | sum of the alcohol and ketone products | alcohol/ketone ratio |
|----------|---------|-----------------------------------------------|------|-----------------|---------------|---------------------------------|-------------------|
| Mn-CFA-1 | air     | no                                           | 1 h  | 3.8             | 5.0           | 8.8                            | 1.1:1             |
| Mn-CFA-1 | air     | yes                                          | 1 h  | 0.4             | 0.5           | 0.9                            | 1:1.3             |
| Mn-CFA-1 | air     | no                                           | 6 h  | 3.6             | 6.9           | 10.7                           | 1:1.8             |
| Mn-CFA-1 | air     | yes                                          | 6 h  | 1.4             | 1.6           | 3                              | 1:1.1             |

<sup>a</sup>Typical reaction conditions: 10 mg of MOF, 1 mL of cyclohexene, 22 °C. <sup>b</sup>10 equiv of 2,4,6-tri-tert-butylphenol. <sup>c</sup>Turnovers of cyclohexen-1-ol calculated with respect to manganese. <sup>d</sup>Turnovers of cyclohexen-1-one calculated with respect to manganese.

The detrimental effect of water on catalysis is made obvious when water is added directly to Mn-CFA-1 prior to the addition of cyclohexene, which reduces product formation by 75%. Although clearly a limitation in batch processes, the generation of reaction-limiting water in situ could be mitigated by implementing catalysis in a flow system. In a setup where all products (desired and otherwise) are swept out of the catalyst bed before having a chance to further react, Mn-CFA-1 could exhibit increased turnover numbers (TONs) for olefin oxidation and potentially increased selectivity. Indeed, a time point study indicated that the bulk of the observed activity for Mn-CFA-1 occurs within the first 15 min of the reaction. Even during this time, the selectivity shifts from an initial alcohol/ketone ratio of approximately 1:1 to the favoring of ketone products over alcohol products as the reaction progresses. The implementation of a flow approach employing Mn-CFA-1 is ongoing.

**CONCLUSION**

The foregoing results highlight how the site isolation and relatively weak ligand fields generally exhibited by MOFs, here exemplified by CFA-1, enable the activation of O<sub>2</sub> by a Mn<sup>II</sup> species in a scorpionate environment, followed by reactivity that has not been accessible in molecular systems. Mn-CFA-1 is capable of catalytically oxidizing weak C–H bonds using air as the terminal oxidant with the proposed intermediacy of Mn<sup>III</sup>OOH. The data provide evidence that this catalysis is influenced by the identity of the charge-balancing anion, and preliminary results suggest that more weakly coordinating species, such as trifluoroacetate, promote even greater catalytic activity. The reaction is inhibited by the inherent formation of water as a side product, likely because of the highly reactive nature of the intermediates in our proposed catalytic cycle, making this catalyst an intriguing candidate for continuous-flow chemistry.

**ASSOCIATED CONTENT**

* Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.inorgchem.9b02068.

Details of experimental procedures and results, PXRD data, nitrogen isotherm data, and EXAFS analysis (PDF)

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**AUTHOR CONTRIBUTIONS**

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

**NOTES**

The authors declare no competing financial interest.

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