



Communication

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Million-Fold Electrical Conductivity Enhancement in Fe₂(DEBDC) versus $Mn_2(DEBDC)$ (E = S, O)

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Supporting Information

ABSTRACT: Reaction of FeCl₂ and H₄DSBDC (2,5disulfhydrylbenzene-1,4-dicarboxylic acid) leads to the formation of $Fe_2(DSBDC)$, an analogue of $M_2(DOBDC)$ (MOF-74, DOBDC $^{4-}$ = 2,5-dihydroxybenzene-1,4-dicarboxylate). The bulk electrical conductivity values of both $Fe_2(DSBDC)$ and $Fe_2(DOBDC)$ are ~6 orders of magnitude higher than those of the Mn²⁺ analogues, $Mn_2(DEBDC)$ (E = O, S). Because the metals are of the same formal oxidation state, the increase in conductivity is attributed to the loosely bound $Fe^{2+}\beta$ -spin electron. These results provide important insight for the rational design of conductive metal-organic frameworks, highlighting in particular the advantages of iron for synthesizing such materials.

Metal—organic frameworks (MOFs) that display intrinsic electrical conductivity are still rare, but conductivity is emerging as an attractive complement to the inherent porosity of these materials. If high surface area is combined with electrical conductivity or high charge mobility, MOFs could find uses in fields outside traditional areas such as gas storage and separation, and make strides into batteries, supercapacitors, electrocatalysis,³ and sensing,⁴ among others. Although recent reports of conductive MOFs have crystallized several potential avenues toward improved electrical properties, including in-plane π conjugation, 4,5 through-space charge transport, 6 through-bond charge transport, 7 and doping, 7d,8 these design strategies require significant refinement.

In this context, we recently reported Mn₂(DSBDC), a MOF-74 analogue that contains $(-Mn-S-)_{\infty}$ chains, and discussed the positive effect of replacing phenoxide by thiophenoxide groups on charge mobility. 7a While this was inspired by the rich literature of organic conductors, where heavier chalcogens generally lead to better electrical properties,9 the equally compelling literature of inorganic semiconductors shows, for instance, that iron chalcogenides ¹⁰ are better intrinsic conductors than manganese chalcogenides, 11 highlighting that the metal ions may be as important as the chalcogens. Taking a cue from inorganic chalcogenides, we wanted to test the relative importance of the metal ion for MOFs with infinite chains as their secondary building units. To do so, we set out to compare the Mn²⁺ and Fe²⁺ analogues of the family of materials known as MOF-74, surmising that replacement of Mn²⁺ by Fe²⁺ would lead to superior electrical conductivity, as seen for the inorganic chalcogenides. Here, we show that within isostructural materials,

replacing Mn²⁺ with Fe²⁺ leads to a million-fold enhancement in electrical conductivity, a considerably more pronounced effect than substituting bridging O atoms with less electronegative S

 $[Fe_2(DSBDC)(DMF)_2] \cdot x(DMF)$ was isolated as dark redpurple crystals after heating a degassed and dry solution of H₄DSBDC and anhydrous FeCl₂ in N₂N-dimethylformamide (DMF) at 140 °C under an N₂ atmosphere for 24 h, and washing with additional DMF. Single-crystal X-ray diffraction analysis of $Fe_2(DSBDC)(DMF) \cdot x(DMF)$ revealed an asymmetric unit containing one Fe atom coordinated by three carboxylate groups, two thiophenoxide groups, and one DMF molecule. The sulfur atoms are coordinated in trans fashion to the Fe²⁺ atom, with Fe-S bond lengths of 2.444(2) and 2.446(2) Å. This indicates that both S atoms interact with the same d orbital of Fe²⁺, an important orbital symmetry requirement for efficient charge transport. 12 Although Fe₂(DSBDC) is isostructural with Fe₂(DOBDC)¹³ and Mn₂(DOBDC),¹⁴ its structure is only topologically related to that of Mn₂(DSBDC). As shown in Figure 1a, whereas Fe₂(DSBDC) has a single metal atom in the asymmetric unit, Mn₂(DSBDC) has two: one that is octahedrally coordinated by donors on DSBDC⁴⁻ ligands, and another that is bound by two DMF molecules. Relevantly, the two distinct metal ions in $Mn_2(DSBDC)$ reduce the symmetry of the $(-Mn-S-)_{\infty}$ chains, which may negatively affect its charge transport properties. As in other MOF-74 analogues, the $(-Fe-S-)_{\infty}$ chains in Fe₂(DSBDC) are bridged by thiophenoxide and carboxylate groups to form a three-dimensional framework with onedimensional hexagonal pores with a van der Waals diameter of \sim 16 Å (Figures 1b and S1).

When $M_2(DEBDC)(DMF)_2 \cdot x(DMF)$ are soaked in dichloromethane and then evacuated under vacuum (100 °C, 2 h), they yield $M_2(DEBDC)(DMF)_2$, a series of materials that are guestfree and where DMF completes the coordination sphere of all metal sites. Infrared spectroscopy and microelemental analysis confirmed that all guest solvent molecules were removed under these conditions (Figure S4). Surprisingly, powder X-ray diffraction (PXRD) revealed that Fe₂(DSBDC)(DMF)₂ is distorted in comparison to $Fe_2(DSBDC)(DMF)_2 \cdot x(DMF)$, a distortion that was not observed in the other three analogues (Figures S3 and S5). Mathematical simulation and DFT optimization of completely solvent-free Fe₂(DSBDC)¹⁵ gave a structure whose simulated pattern agreed well with the observed PXRD pattern of Fe₂(DSBDC)(DMF)₂ (see Figure 1c and the

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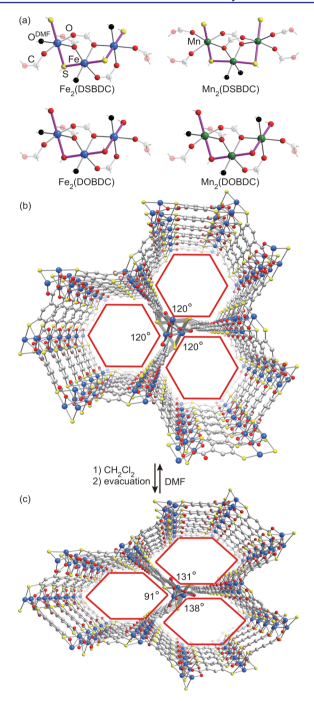


Figure 1. (a) Parts of the infinite secondary building units in $M_2(DEBDC)(DMF)_2 \cdot x(DMF)$ (M = Fe, Mn; E = S, O). The (-M-E-) $_{\infty}$ chains are represented in purple. (b,c) Partial structures of $Fe_2(DSBDC)(DMF)_2 \cdot x(DMF)$ and $Fe_2(DSBDC)(DMF)_2$ as determined by single-crystal X-ray diffraction and DFT structure optimization, respectively. H atoms and solvent molecules are omitted for clarity.

Supporting Information). This distortion is reversible: soaking $Fe_2(DSBDC)(DMF)_2$ in fresh DMF produced a crystalline phase with a PXRD pattern identical to that of $Fe_2(DSBDC)-(DMF)_2\cdot x(DMF)$ (Figure S6). To our knowledge, although breathing behavior has been observed for several classes of MOFs, ¹⁶ it has never been associated with MOF-74 analogues.

 N_2 sorption analysis revealed Brunauer–Emmet–Teller (BET) surface areas of 232, 241, and 287 m²/g for Mn₂-(DSBDC)(DMF)₂, Fe₂(DOBDC)(DMF)₂, and Mn₂-

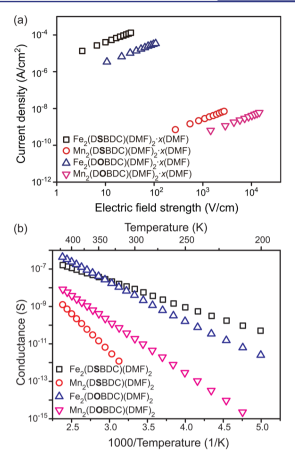


Figure 2. Electrical properties of $M_2(DEBDC)$ (M = Fe, Mn; E = S, O) pressed pellets. (a) Plots of current density versus electric field strength (J-E curves) for $M_2(DEBDC)(DMF)_2$:x(DMF) at 297 K. (b) Conductance—temperature relationship for $M_2(DEBDC)(DMF)_2$.

(DOBDC)(DMF)₂, respectively, confirming their guest-free nature and permanent porosity (Figure S8). These values are lower than those expected for completely activated MOF-74 analogues because coordinated DMF molecules occupy a significant portion of the pore volume in M₂(DEBDC)(DMF)₂. Fe₂(DSBDC)(DMF)₂ exhibited a lower BET surface area of 54 m²/g, likely because the distorted pores in this case are almost entirely occupied by coordinated DMF molecules (Figure S9).

Owing to the small crystallite size of all of these materials, single-crystal measurements of their electrical properties proved unfeasible. Accordingly, we prepared pellets of both M2-(DEBDC)(DMF)₂· κ (DMF) and M₂(DEBDC)(DMF)₂ materials, for a total of eight samples, all of which were analyzed by twoprobe current-voltage techniques. Plots of measured current density versus electric field strength, shown for the assynthesized samples in Figure 2a and for the guest-free samples in Figure S10, revealed striking differences in electrical conductivity between the Fe and Mn analogues, regardless of their solvation level. Indeed, both Fe₂(DSBDC)(DMF)₂· x(DMF) ($\sigma = 3.9 \times 10^{-6} \text{ S/cm}$) and $Fe_2(DOBDC)(DMF)_2$. x(DMF) ($\sigma = 3.2 \times 10^{-7}$ S/cm) were ~6 orders of magnitude more conductive than $Mn_2(DSBDC)(DMF)_2 \cdot x(DMF)$ and $Mn_2(DOBDC)(DMF)_2 \cdot x(DMF)$, which exhibited conductivities of 2.5×10^{-12} and 3.9×10^{-13} S/cm, respectively (Table 1).¹⁷ Although the guest-free materials showed slightly lower conductivities overall, possibly due to additional defects and grain boundaries caused by the solvent exchange and guest removal process, they reflected the same remarkable 6 orders of Journal of the American Chemical Society

Table 1. Electrical Properties of $M_2(DEBDC)$ (M = Fe, Mn; E = S, O)

	Fe ₂ (DSBDC)	Mn_2 (DSBDC)	Fe_2 (D O BDC)	Mn_2 (D O BDC)
$\sigma_{ ext{as-synthesized}} (ext{S/cm})^a$	3.9×10^{-6}	2.5×10^{-12}	3.2×10^{-7}	3.9×10^{-13}
$\sigma_{ ext{guest-free}} \ (ext{S/cm})^b$	5.8×10^{-7}	1.2×10^{-12}	4.8×10^{-8}	3.0×10^{-13}
$E_{\rm a}~({\rm eV})^c$	0.27	0.81	0.41	0.55
$E_{g} (eV)^{d}$	1.92	2.60	1.47	2.48
$\Phi (eV)^e$	3.71	3.81	2.81	3.72

^aElectrical conductivity of $M_2(DEBDC)(DMF)_2 \cdot x(DMF)$ at 297 K. ^bElectrical conductivity of $M_2(DEBDC)(DMF)_2$ at 297 K. ^cActivation energy of $M_2(DEBDC)(DMF)_2$. ^dCalculated bandgap of $M_2(DEBDC)(DMF)_2$. ^eCalculated work function of $M_2(DEBDC)(DMF)_2$.

magnitude difference in conductivity between the Fe and Mn analogues (Table 1). Although at the lower end of intrinsically conductive and porous MOFs, whose conductivity ranges between 10^{-6} and 10^2 S/cm, $^{4-7}$ the conductivity of the Fe frameworks described here is the highest in the MOF-74 family and is comparable to that of typical organic conductors (> 10^{-6} S/cm). 18

To probe the cause of the large difference between the Fe and Mn analogues, we determined the activation energy in M₂(DEBDC)(DMF)₂ by measuring their pellet conductance under variable temperature between 200 and 420 K. Working with guest-free rather than as-synthesized materials was necessary because our variable-temperature, air-free electrical microprobe setup requires that samples be passed through an evacuation chamber. However, it is reasonable to assume that because neutral guest solvent molecules are unlikely to contribute to charge transport, the activation energies of M₂(DEBDC)(DMF)₂ are not vastly different and follow the same trends as those of $M_2(DEBDC)(DMF)_2 \cdot x(DMF)$. All samples showed an increase in conductance with increasing temperature, indicative of semiconducting behavior (see Figure 2b). Fitting the respective conductance values, G, to the Arrhenius law (eq 1) revealed notable differences in activation

$$G = G_0 \exp(-E_a/kT) \tag{1}$$

energies, $E_{\rm av}$ shown in Table 1. Thus, both Fe₂(DSBDC)(DMF)₂ ($E_{\rm a}=0.27~{\rm eV}$) and Fe₂(DOBDC)(DMF)₂ ($E_{\rm a}=0.41~{\rm eV}$) had considerably lower activation energies than Mn₂(DSBDC)-(DMF)₂ ($E_{\rm a}=0.81~{\rm eV}$) and Mn₂(DOBDC)(DMF)₂ ($E_{\rm a}=0.55~{\rm eV}$), suggesting that the Fe-based MOFs have smaller band gaps and hence higher charge density than the Mn-based MOFs.

Notably, the addition of a single electron per metal ion (i.e., substitution of d⁵ Mn²⁺ for d⁶ Fe²⁺) has a much more pronounced positive effect on conductivity than changing the bridging atom from O to S, indicating that the electronic structure of the metal ions plays the most important role in charge conduction in this class of materials. To confirm the oxidation state and high-spin configuration of the Fe atoms, we measured ⁵⁷Fe Mössbauer spectra of both Fe-based MOFs. As shown in Figures S11 and S12, these spectra revealed well-resolved doublets characterized by isomer shifts of 1.172 and 1.308 mm/s, and quadrupole splittings of 3.218 and 2.739 mm/s, for Fe₂(DSBDC) and Fe₂(DOBDC), respectively. Because the isomer shifts of both MOFs fall in the expected range of high-spin

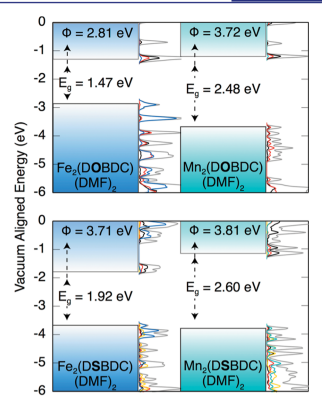


Figure 3. Calculated energy bands and projected density of states (DOS) of $M_2(DEBDC)(DMF)_2$ (M=Fe,Mn;E=S,O). The work function, Φ , and the absolute energy scale are aligned to vacuum according to ref 19. Gray curves represent total DOS. Blue, teal, yellow, red, and black curves represent projected DOS of Fe, Mn, S, O, and C, respectively.

 ${\rm Fe^{2+}}$, these experiments demonstrated that oxidation to ${\rm Fe^{3+}}$ did not occur during our experiments.

Density functional calculations were used to further probe the differences in electronic structure of M₂(DEBDC)(DMF)₂, and the significance of the additional d electron associated with the Fe²⁺ ions. The electronic density of states and ionization potentials of the guest-free system are presented in Figure 3, and detailed in the Supporting Information. Because Fe₂(DOBDC) and Mn₂(DOBDC) are structurally analogous, while Fe₂-(DSBDC) and Mn₂(DSBDC) differ in the number of metal ions in their asymmetric units, the comparison between Fe₂(DOBDC) and Mn₂(DOBDC) illustrates best the difference between Mn²⁺ and Fe²⁺. Most importantly, the valence band maximum of Mn₂(DOBDC) is composed of C-p, O-p, and Mn-d states, while in Fe₂(DOBDC) the Fe-d states dominate the valence band, with negligible contribution from ligand orbitals. This difference is attributed to the low binding energy of the filled β -spin d band of Fe²⁺, which is empty for the d⁵ high-spin Mn²⁺ ions.²⁰ Furthermore, because the lower conduction band in both MOFs is dominated by ligand-based orbitals, the band gaps are narrowed owing to a decreased work function. As a result, the calculated work functions and band gaps of Fe₂(DOBDC) are 0.91 and 1.01 eV smaller than those of Mn₂(DOBDC), respectively.

To assess the relative importance of the chalcogen atom on the charge transport, we also compared the structurally analogous $Fe_2(DSBDC)$ and $Fe_2(DOBDC)$ materials. First, the valence and conduction bands of these frameworks are flat in reciprocal space (dispersion of <100 meV in all cases). This behavior is indicative of localized orbitals rather than delocalized bands, and

is typical of many MOFs. ²¹ Thus, we anticipate that the primary mode of conduction is through charge hopping. Moreover, because the Fe²⁺ d orbitals dominate the valence band (83% of the orbital contribution), intervalence transitions between Fe atoms will proceed with little contribution from O. In contrast, due to the enhanced orbital overlap in Fe₂(DSBDC), where Fe and S orbitals contribute 53% and 14% to the valence band, transport will occur through both Fe and S in the (-Fe-S-) $_{\infty}$ chains. This mechanism lowers the charge hopping barrier and is also associated with the larger work function of Fe₂(DSBDC) compared with that of Fe₂(DOBDC). Finally, the increased contribution of C-p states to the valence band in Fe₂(DSBDC) compared to Fe₂(DOBDC) may also indicate a more efficient interchain transport, which further increases the conductivity of the former.

In summary, the synthesis of a new MOF-74 analogue based on $(\text{-Fe-S-})_{\infty}$ chains led to a material with the highest conductivity in the MOF-74 structural family. The combination of loosely bound Fe^{2+} $\beta\text{-spin}$ electrons and the low electronegativity of S atoms contributes to the higher relative conductivity of Fe₂(DSBDC). Applying similar design principles to other MOFs made from one-dimensional secondary building units should lead to further improvements in electrical properties for materials in this class.

ASSOCIATED CONTENT

S Supporting Information

Experimental details, table of X-ray refinement details, PXRD patterns, IR spectra, I-V curves, Mössbauer spectra, isotherms, computational details, and crystallographic data (CIF). The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/jacs.5b02897.

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Notes

The authors declare no competing financial interest.

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