**Electrical Conductivity in a Porous, Cubic Rare-Earth Catecholate**

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**ABSTRACT:** Electrically conductive metal–organic frameworks (MOFs) provide a rare example of porous materials that can efficiently transport electrical current, a combination that is favorable for a variety of technological applications. The vast majority of such MOFs are highly anisotropic in both their structures and properties: Only two electrically conductive MOFs reported to date exhibit cubic structures that enable isotropic charge transport. Here we report a new family of intrinsically porous frameworks made from rare-earth nitrates and hexahydroxytriphenylene. The materials feature a novel hexanuclear secondary building unit and form cubic, porous, and intrinsically conductive structures, with electrical conductivities reaching $10^{-3}$ S/cm and surface areas of up to 780 m$^2$/g. By expanding the list of MOFs with isotropic charge transport, these results will help us to improve our understanding of design strategies for porous electronic materials.

**RECENT ADVANCES IN ELECTRICALLY CONDUCTIVE METAL–ORGANIC FRAMEWORKS (MOFS) SIGNIFICANTLY EXPANDED THE RANGE OF APPLICATIONS FOR THIS BROAD CLASS OF HYBRID ORGANIC–INORGANIC MATERIALS, ACHIEVING IMPORTANT MILESTONES IN ELECTROCHEMICAL CATALYSIS, ELECTROCHROMISM, ELECTROCHEMICAL SENSORS, AND OTHER RELATED FIELDS.**

**Graphene**10 and silicon nanowires11 is of significant fundamental interest to develop electrically conductive MOFs with isotropic charge transport.12 Remarkably, to date, only two MOFs with cubic symmetry have been reported to show intrinsic (non-guest-based) electrical conductivity.13,14 The highest conductivities to date are observed in MOFs based on ligands with catechol- and semiquinone-derived functional groups, such as hexaaminotriphenylene,9,15 and hexahydroxytriphenylene (HHTP),3,16 among others.17 A number of factors makes these ligands especially viable for making electrically conductive frameworks, including the strong $\pi$–d conjugation between the metal and the ligand, the ligands’ propensity for $\pi$–$\pi$ stacking, and, crucially, their redox activity: The oxidation of the catechol moieties forms semiquinoid radicals that serve as the charge carriers in the redox activity: The oxidation of the catechol moieties forms semiquinoid radicals that serve as the charge carriers in the materials.17–26 It is therefore unsurprising that of the two conductive MOFs with cubic symmetry, one is based on a ligand belonging to this class, dihydroxybenzoquinone (DHBQ). The cubic framework $[\text{Fe}_2(\text{DHBQ})_3][\text{Bu}_4\text{N}]_2$ shows an impressive conductivity of 0.16 S/cm.14 This value is controllable through postsynthetic redox chemistry, where the radical semiquinoid linkers can be reduced to their closed-shell catechol form to lower the free carrier concentration. This framework, however, is nonporous due to the bulky charge-compensating tetrabutylammonium cations blocking the small pores. Nevertheless, partly inspired by these results, we turned to the larger, chemically related HHTP ligand to target a potentially porous, isotropically conductive MOF.

The reaction of concentrated solutions (>0.5 M) of hydrated rare-earth nitrates $\text{M(NO}_3)_3\cdot\text{nH}_2\text{O}$ ($\text{M} = \text{Y, Eu, La}$) with HHTP in a mixture of water and $\text{N,N'}$-dimethylimidazolidinone (DMI) at temperatures above 150 °C yields large octahedral yellow crystals (Figure S11) of $[\text{M}_6(\mu_6-\text{NO}_3)(\text{HOTP})_2]^{3+}$ ($\text{M}_{\text{HOTP}}$, $\text{M} = \text{Y, Eu}$; HOTP = 2,3,6,7,10,11-hexaoxytriphenylene). The charge balance is provided by five monoanions (hydroxides or nitrates), which correspond to a fully reduced ligand, HOTP$^{−}$. This assignment is corroborated by the color of the crystals: The yellow-brown color matches the color of unoxidized, air-free solutions of HHTP. Minor oxidation of HHTP leads to the formation of intensely colored blue or purple species. X-ray diffraction analysis of single crystals revealed that all three compounds crystallize in the cubic space group $Fd\bar{3}m$. (See Figure 1 and Tables S2 for structural parameters.) The secondary building units (SBUs) consist of hexanuclear $\mu_6$-nitro catecholate clusters (Figure 2b), wherein the metal ions alternate above and below the plane of the central nitrate ion in a chairlike conformation. Six HHTP ligands surround each secondary building unit (SBU) and connect neighboring SBUs into tetrahedral cages (Figure 2c). The cages themselves are connected at the vertices to provide a net with 6- and 3-connected nodes and overall $spn$ topology (Figure 2a). The same topology has been previously observed in several MOFs.
most notably in MOF-808, \(^{28}\) and is homologous to the faujasite zeolite structure type (Figure 1).

The SBU shows significant crystallographic disorder, as may be expected given the coordination flexibility of rare-earth ions. Metal atoms in \(\text{Eu}_6\text{HOTP}_2\) and \(\text{Y}_6\text{HOTP}_2\) have similar eight-coordinate environments (Figure S5), with each metal connected to one oxygen atom of the central bridging nitrate, four catecholate oxygen atoms of three different HOTP linkers, and a disordered combination of aqua and nitrate ligands oriented toward the pore or the inner cavity of the tetrahedral cage. Notably, this SBU has not been previously observed in MOFs. The discovery of new SBUs is a driver for expanding the reticular chemistry of MOFs, \(^{29}\) yet reports of new inorganic building units whose structural features are conserved across different metals have become increasingly rare. The new SBU is particularly notable because it features catecholate ligands, which thus far have given rise exclusively to single-ion SBUs, with very few exceptions. \(^{22,30–32}\) As with the earliest SBUs made from carboxylates, the hexanuclear SBU reported here is also a close mimic of a molecular hexalanthanide cluster that has been previously reported (Figure S6). \(^{33}\) This similarity reinforces the strategy of targeting high-symmetry clusters known in molecular chemistry as potential SBUs for novel MOFs, a strategy commonly employed in reticular chemistry with more traditional functional groups such as carboxylates and azolates. \(^{34}\)

Owing to its propensity to \(\pi\)-stack, HHTP rarely forms 3D structures. \(^{12}\) Indeed, only one other report details the formation of isotropic, 3D MOFs from HHTP. \(^{27}\) Instead, this ligand preferably leads to stacked structures \(^{16}\) reinforced by close contacts between its extended aromatic cores. Notably, although the 2D sheets and the close contacts in most HHTP-based materials lead to some of the highest electrical conductivities among porous MOFs, \(^{35}\) the electrical transport in these materials is anisotropic. In fact, almost all conductive MOFs reported thus far have lower symmetry than cubic and therefore exhibit anisotropic transport. \(^{12}\)

To probe whether \(\text{M}_6\text{HOTP}_2\) represents a rare example of a cubic, electrically conductive MOF, we tested the electrical conductivity of activated crystalline powders with a two-probe apparatus that has been previously described. \(^{36}\) The activation of the crystals involved heating under a dynamic vacuum at 95 °C for 18 h. We note that thermogravimetric analysis of as-synthesized samples exchanged sequentially with water, methanol, and acetonitrile showed a first significant mass loss...
at ~90 °C (Figure S1). A representative sample of Y₆HOTP₂, activated as previously described, also showed permanent porosity, as indicated by a type I N₂ adsorption isotherm, which can be fit to the Brunauer–Emmett–Teller (BET) model¹⁵ to give an apparent surface area of 780 m²/g. This value is in line with the reported surface area of the topologically equivalent MOF-808 (1140 m²/g)²⁸ which has near-identical unit-cell parameters to those of Y₆HOTP₂.

Relevantly, although as-synthesized crystals of M₆HOTP₂ are yellow, they slowly darken (Figure S12) over time, even under rigorous air-free conditions, likely indicating at least the partial oxidation of catechol moieties in HOTP₆⁻ ligands to semiquinones. This transformation happens over the course of many weeks at room temperature, but heating or exposure to air causes significant acceleration. Although prolonged exposure to air leads to the eventual loss of crystallinity, up to complete amorphization (Figure S8), heating under inert conditions preserves the structure while still accelerating the color change.

We tracked this color change by in situ diffuse reflectance spectroscopy of a powder of Y₆HOTP₂ heated to 90 °C under a dinitrogen atmosphere, as shown in Figure 3. The spectra closely resemble those reported for chemically similar molecular clusters, such as the trinuclear Co(III) HOTP complex,³⁸ as well as the trinuclear Fe(III) complex with a hexaaminotriphenylene-based ligand.³⁹ Importantly, spectral changes observed for Y₆HOTP₂ upon heating also mimic the behavior of the Fe(III) complex upon oxidation. Specifically, the MOF spectra exhibit a decrease in relative intensity of the bands at 300–450 nm (attributed to π−π* transitions within the aromatic core) and at 800 nm (attributed to ligand-to-metal charge transfer) as well as the appearance of a new band at 1000–1200 nm, commonly associated with the π−π* transitions in radical semiquinoid moieties.³⁸,³⁹

The spectral changes previously discussed confirm the partial oxidation of the material and the potential formation of free charge carriers that could improve the electrical conductivity. Indeed, both Y₆HOTP₂ and Eu₆HOTP₂ show electrical conductivities on the order of 10⁻⁶ to 10⁻⁵ S/cm (Figure 4) upon activation, across several independently synthesized batches. Although not as high as the bulk conductivities of HHTP-derived MOFs that exhibit close HOTP stacking (10⁻⁴–10⁻² S/cm),³,¹⁶ these values are remarkable²⁄² for 3D, isotropic porous MOFs, where charges are expected to transport with equal efficiency along the three Cartesian coordinates.

It is rare for intrinsic porosity and electrical conductivity to coexist within one material. It is even rarer for such compounds to be structurally isotropic. In this work, we presented a new family of MOFs that combine significant porosity with electrical conductivity in a cubic framework. The relatively high conductivity shown by these materials demonstrates that targeting low-dimensional pathways is not mandatory for achieving efficient charge transport. We hope these results will inspire more detailed studies into the dimensionality of charge transport in this exciting class of materials and eventually lead to a better understanding of how porosity interacts with conductivity.

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**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.0c01713.

- Additional experimental details and procedures (PDF)
- X-ray crystal structure of Eu₆HOTP₂ (CIF)
- X-ray crystal structure of Y₆HOTP₂ (CIF)
- X-ray crystal structure of La₆HOTP₂ (CIF)

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Figure 3. Diffuse reflectance spectra for Y₆HOTP₂, showing the evolution of the electronic structure of the material on heating under an inert atmosphere (black, initial spectrum; yellow, final spectrum). The broad band at 1000–1200 nm is often considered a fingerprint of semiquinoid radicals in chemically related compounds.

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Figure 4. Current–voltage characteristics of Y₆HOTP₂, obtained on pressed pellets in a two-probe configuration, showing perfect linear behavior in accordance with Ohm’s law. The linear fit of the data (dashed line) reveals the conductivity of this representative device to be 2.0 × 10⁻⁵ S/cm.
Author Contributions
The manuscript was written through contributions of both authors. Both authors have given approval to the final version of the manuscript.

Notes
The authors declare no competing financial interest.

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