Tunable Electrical Conductivity of Flexible Metal–Organic Frameworks

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ABSTRACT: We present the computational design of naphthalene diimide (NDI)-containing metal pyrazolate MOFs (M(NDIDP), M = Zn, Co, Fe) judiciously bestowed with both framework flexibility and electrical conductivity. M(NDIDP) MOFs exhibit "wine rack" type flexibility, allowing them to partake in pressure-induced structural transitions at low pressures and room temperature. The MOFs are also equipped with closely packed redox-active NDI moieties, which results in dispersive conduction band minima for n-type charge transport and conductivity. Remarkably, structural transitions of the designed MOFs are found to facilitate the rearrangement of NDI moieties. As a result, conductivity of M(NDIDP) MOFs can be finely tuned, which is evidenced by the gradual shift from 1D to 2D charge transport and up to 7.5-fold reduction in carrier effective mass (2.76 m_0 to 0.37 m_0) when transitioning between different structural configurations. The unprecedented discovery of flexible MOFs with tunable electrical conductivity presented in this work firmly establishes MOFs as versatile candidate materials for multifunctional electronics.

KEYWORDS: metal–organic framework, electrical conductivity, flexibility, stimuliresponsive behavior, density functional theory

Introduction

The high modularity of metal-organic frameworks (MOFs) with multiple design parameters (e.g., metals ions/clusters,¹ organic linkers,² topologies,³ and defects^{4,5}) has led to the discovery of a large number of MOFs with a wide range of properties and functionalities. Judicious choice of the framework components makes it possible to tailor these materials for a variety of applications, ranging from gas storage and separation⁶ to catalysis,⁷ drug delivery,⁸ water harvesting,^{9,10} etc. More recently, research has shown that electrical conductivity can be induced in MOFs.¹¹⁻¹⁵ Due to their porosity and conventional use of electrically inactive components (i.e., closed-shell metals, σ-bridged organic linkers), MOFs have long been considered as electrical insulators. However, by synthesizing MOFs from redox-active metals and linkers that allow significant delocalization of charge carriers, it becomes possible to secure long-range charge transport (CT) pathways in MOFs to induce conductivity. Experimental conductivities of MOFs have reached up to 2500 S/cm,¹⁶⁻¹⁸ and as such, they have been successfully integrated in field-effect transistors,¹⁹ supercapacitors,²⁰ chemiresistive sensors,²¹⁻ ²³ and electrocatalysts.²⁴ While substantial progress has been made, research thus far has mainly focused on inducing the conductivity in MOFs. To make further breakthroughs, however, focus should be shifted towards synergistically combining conductivity with other properties unique to MOFs, allowing novel application schemes unfeasible in other materials.

One of the most intriguing properties of MOFs that differentiates them from other materials is framework flexibility.²⁵ Generally, MOFs are considered to be more flexible than other solid crystalline materials as they inherently possess several structural degrees of freedom (e.g., linker rotation, lattice deformation). Here, however, we specifically refer to flexibility as those cases where MOF constituents participate in large amplitude structural transitions whilst conserving the original framework topology. Such flexible MOFs are capable of undergoing significant expansion or contraction of the framework, which can drastically alter their pore geometry and internal arrangement of framework components. This can be viewed akin to a switch or a dial, as the framework is able to effectively transition between clearly distinct states or structural configurations via external stimuli (e.g., guest adsorption, temperature, light, pressure).^{26,27} As such, framework flexibility can effectively instill new external parameters (e.g., stimuli and time) for the control of MOF properties, opening up many new applications.²⁸ Previous studies have utilized the flexibility of MOFs for selective chemical sensing,²⁹ switchable catalysis,³⁰ or band gap modulation.³¹ However, to the best of our knowledge, no

one has linked framework flexibility directly with the electrical conductivity of MOFs, which could potentially lead to tunable electrical conduction properties.

In this computational study, we have performed the modular design of MOFs that simultaneously harness both framework flexibility and electrical conductivity. To guarantee the framework flexibility of our designed MOFs, a well-known flexible MOF (i.e., M(BDP),³²⁻³⁴ $H_2BDP = 1,4-di(1H-pyrazol-4-yl)$ benzene) was adopted as the template framework. Then, central benzene groups of the linkers were replaced with redox-active 1,4,5,8-naphthalene diimide (NDI) moieties to impart electrical conductivity, resulting in the newly proposed MOFs referred to as M(NDIDP). Density functional theory (DFT) modeling of M(NDIDP) MOFs was performed to discover the presence of multiple metastable phases with vastly different internal arrangement of the NDI moieties. Next, molecular dynamics (MD) simulations with DFTderived force fields were carried out to show that the MOFs can undergo pressure-induced phase transitions at room temperature, firmly establishing their framework flexibility. DFT electronic band structure analysis revealed that M(NDIDP) MOFs exhibit notable dispersions in the conduction band minima for efficient n-type charge transport and conductivity. Strikingly, conductivity of M(NDIDP) was also found to be highly configuration-dependent, as evidenced by the shift from 1D to 2D charge transport and up to 5.7-fold increase in band dispersion (7.5fold reduction in carrier effective mass) between different structural configurations within a single material. This effectively demonstrates that electrical conductivity of M(NDIDP) MOFs can be finely tuned via application of external stimuli, making them excellent candidate materials for multifunctional electronics.

Results and Discussion

Modular Design of M(NDIDP)

Many flexible MOFs have been synthesized to date, and diverse structural motifs for framework flexibility have been implemented. One of the most common structural motifs for flexibility is the "wine rack" motif, where one-dimensional metal clusters are present as pillars and bidentate linkers serve as spokes that bind the pillars together (**Figure 1a**). In such MOFs, coordination bonds between the metals and linkers function as hinges, and they are collectively arranged in a manner that allows for a significant expansion and contraction of the framework. Representative MOFs of this motif are MIL-47,³⁵ MIL-53,³⁶ and M(BDP),^{32–34} which are

widely recognized for their stepwise gas adsorption behavior arising from the large variability in the pore volume. Here, it is important to note that flexibility in these MOFs is conferred by their unique metal-linker coordination environment and topologies. Hence, by performing further structural modifications without altering the coordination environment, new properties could be instilled in the wine rack MOFs whilst retaining their flexibility.



Figure 1. (a) Flexible MOFs with wine rack structural motif built from 1D metal clusters and bidentate organic linkers. (b) Bidentate pyrazolate linkers considered for MOF construction. (c) Structure of the newly proposed M(NDIDP) MOFs shown along with the metals that have been used for the construction of the three separate MOFs.

Recently, Wentz et al. reported switchable electrical conductivity in a zinc pyrazolate MOF via reversible n-doping of the framework.³⁷ Interestingly, this MOF is isoreticular to M(BDP) but displays two key structural differences: redox-active NDI moieties are present as central moieties in the linkers, and the pyrazolate binding moieties are methylated at the 3 and 5 positions (**Figure 1b**). Electrical conductivity is effectively conferred to the MOF by the

presence of NDI moieties, as they are stacked together in a regular fashion to facilitate CT. However, the methyl groups of pyrazolate binding moieties are found to significantly hinder the MOF from partaking in large amplitude structural transitions despite the wine rack motif (see **Supporting Information**). As such, we conjectured that modular substitution of benzene with NDI in M(BDP) could induce electrical conductivity in the resulting MOF while retaining the framework flexibility, thereby allowing for stimuli-controlled conductivity.

Resultantly, MOFs isoreticular to M(BDP) yet containing NDI moieties were created *in silico* (**Figure 1c**). First, a four-linker unit cell of M(BDP) was procured from the Cambridge Structural Database (refcode: QUPZAE and QUPZIM). The central benzene moieties of the BDP linkers were then computationally substituted with NDI, resulting in the proposed MOF structures. In the process, the original unit cell of M(BDP) was expanded along the linker directions to easily accommodate the redox-active NDI moieties. Three different MOFs were built from Zn, Co, and Fe metal pillars, which allows further investigation of how different metal ions affect the resulting properties. The new MOFs are herein referred to as M(NDIDP) (M = Zn, Co, Fe, NDIDP = *N*,*N*-di(1H-pyrazol-4-yl)-1,4,5,8-naphthalenediimide). It must be stated that the NDI-containing linkers incorporated into M(NDIDP) have yet to be synthesized. Nonetheless, judging from the abundance of similar NDI-containing organic linkers and their successful incorporation into MOFs,^{38–42} we expect the linker synthesis to become possible in the near future.

Flexibility of M(NDIDP): Energy vs. Volume Profile

Next, calculations were performed to determine whether framework flexibility was successfully manifested in our M(NDIDP) MOFs. As previously discussed, flexibility targeted in this work is characterized by large amplitude expansion and contraction of the framework. In such a case, structural degrees of freedom exhibited by the MOF would allow a wide range of energetically accessible configurations, often with multiple metastable phases present as minima in the (free) energy profile. Conversely, a rigid MOF would be characterized by a narrow, parabolic energy profile with a single global minimum. Note that the expansion and contraction of flexible MOFs often incur vast differences in their unit cell volume. Hence, energy-versus-volume, or E(V) profiles for each of the M(NDIDP) MOFs were obtained at the DFT level of theory (see **Figure 2**). Here, each point corresponds to an optimized structural

configuration at the given unit cell volume, plotted according to its energy difference from the global energy minimum. The E(V) profiles of all three M(NDIDP) MOFs exhibit a significantly wide range of energetically accessible configurations (i.e., $\Delta E < 2$ eV) bounded by the lower and higher volume limits. At these volume extrema, energy rises steeply upon further structural compression or expansion that surpasses the range of motion allowed by the flexible MOFs. Provided that a multitude of energetically accessible configurations are observed over a wide volume range as opposed to the case of rigid MOFs,^{43–45} one can conclude that M(NDIDP) MOFs have the potential to show framework flexibility.



Figure 2. E(V) profiles of (a) Zn(NDIDP), (b) Co(NDIDP), and (c) Fe(NDIDP). Dark green and magenta points correspond to the (meta)stable *np* phase and *lp* phase of the MOFs, for which the corresponding MOF configurations are shown as insets. Lighter green and lighter

magenta points indicate the squeezed (sq) and expanded (exp) configurations discussed later. Light blue points for Co(NDIDP) indicate intermediate (int1 and int2) configurations also discussed later. Further structural information of the highlighted phases is shared in **Table S1**.

The global energy minima of the three M(NDIDP) MOFs are found in the lower volume regime, at V = 1867.6, 1875.2 and 1908.3 Å³ for Zn, Co, and Fe(NDIDP) respectively. MOF configurations at these volumes (see Figure 2 insets and Figures S1-S3) correspond to the narrow pore (np) phase, and hence all three M(NDIDP) MOFs are expected to energetically favor existing in the np phase over other structural configurations at 0 K. Interestingly, another energy minimum is found at V = 3735.4 Å³ for Zn(NDIDP), and also for Co(NDIDP) at V =3688.9 Å³, which correspond to a metastable large pore (*lp*) phase. In the case of Fe(NDIDP), however, such an additional *lp* energy minimum is not observed, and the *np* phase is the only thermodynamically stable structural configuration. Nonetheless, the E(V) profile of Fe(NDIDP) is far from being strictly parabolic and displays a large, shoulder-like feature in the higher volume range. This indicates that the higher volume configurations of Fe(NDIDP) are still energetically accessible, yet these configurations cannot be stably sustained by the framework at 0 K. Such differences in the E(V) profiles can be explained by the softer binding tendency of Fe, which would be more lenient towards configurational changes around the coordination environment. As such, one can conclude that the binding tendency of the metal determines the stability of the additional, metastable lp phase that could be exhibited by the MOF. For the sake of consistency in the analyses that follow, the $V = 3667.4 \text{ Å}^3$ configuration of Fe(NDIDP) was chosen to serve as the lp phase, as it is the closest in unit cell volume to Zn(NDIDP)-lp and Co(NDIDP)-lp.



Figure 3. Internal arrangement of the NDI moieties in Co(NDIDP)–lp and Co(NDIDP)–np. The parallel stacking distance (top left of each panel) and planar alignment offset (bottom left of each panel) in the *z*-direction, as well as the inter-NDI distances in the *x*- and *y*-direction

(right of each panel), are presented. As for the *np* phase, herringbone packing configuration of NDI moieties is separately shown with their edge-to-face distance.

Having installed the redox-active NDI moieties to induce CT, their relative arrangements within M(NDIDP) under the different, energetically accessible configurations were studied. Figure 3 shows the internal NDI arrangements of Co(NDIDP) in the (meta)stable *lp* and *np* phases as a case study. In Co(NDIDP)–*lp*, NDI moieties are arranged in a paralleldisplaced manner along the z-axis. The stacking distance between adjacent NDI moieties is 3.26 Å, and the planar alignment offset is 5.44 Å. These distances constitute reasonable contact between NDI moieties for π orbital overlap, which is essential for the anticipated CT. Along the x-axis (resp. y-axis), the inter-NDI distance surpasses 7.00 Å (resp. 5.52 Å) at the closest contact (i.e., N–O distance between imides of adjacent NDI), effectively prohibiting π orbital overlap in these directions. In Co(NDIDP)-np, NDI moieties retain their parallel-displaced configuration along the z-axis, but with notable differences from the lp phase. The stacking distance between NDI moieties is decreased to 3.12 Å, and planar alignment is increased to 5.85 Å. Interestingly, contraction of the framework into the *np* phase also leads to the onset of edge-to-face interactions between adjacent NDIs along the x-axis, as evidenced by the short NDI–H distances of 2.47 Å. The inter-NDI distances along the *y*-axis surpass 8.43 Å, further prohibiting orbital overlap in this direction. Hence, Co(NDIDP)-np exhibits two-dimensional, herringbone packing of NDI in the xz-plane, where the orbital overlap between NDI moieties could extend along the two packing dimensions. As shown in Table S2, similar internal NDI arrangements in the lp and np phases are also found in Zn(NDIDP) and Fe(NDIDP). All in all, vastly distinct internal NDI arrangements are observed in the different structural configurations of M(NDIDP), which could give rise to notably different CT properties.

Flexibility of M(NDIDP): Pressure vs. Volume Equation of State

For a MOF to be truly flexible, transitions between different metastable phases and structural configurations must be possible. Hence, the thermodynamic feasibility of structural transitions in M(NDIDP) MOFs was subsequently investigated. In doing so, we specifically focused on the pressure-induced structural transitions in M(NDIDP) materials, where an external pressure or mechanical stress is used as the stimulus. To this end, DFT-based force fields were derived using the QuickFF software for each of the three M(NDIDP) MOFs and their pressure-versus-volume or P(V) equations of state at T = 300 K were derived following

the established protocol,^{46,47} resulting in the profiles shown in **Figure 4**.



Figure 4. P(V) equations of state at T = 300 K for (a) Zn(NDIDP), (b) Co(NDIDP), and (c) Fe(NDIDP). Gray dotted line pertaining to the ambient pressure is drawn at 0.1 MPa. Regions shaded in red indicate the mechanically unstable volume range as determined by positive $\partial P/\partial V$ values. In the case of Fe(NDIDP), a secondary partial P(V) curve is shown in light green. This corresponds to the sheared configurations that have deviated away from the main mode of flexibility along the *x*-axis, as discussed in more detail in the **Supporting Information**.

Owing to their structural motif, the three M(NDIDP) structures exhibit P(V) equations of state similar to other wine rack MOFs.^{27,46,48} Specifically, an intermediate, mechanically unstable region with positive $\partial P/\partial V$ values is found to separate two mechanically stable *np* and *lp* branches, which extend toward low volumes (below 2250 Å³) and high volumes (above 3340 Å³ for Fe(NDIDP), 3740 Å³ for Co(NDIDP), and 3790 Å³ for Zn(NDIDP)), respectively. Each point on these two mechanically stable branches can, in principle, be thermodynamically stabilized by applying a suitable isotropic stress, either positive or negative in sign, given by the pressure value at the specific volume point. Provided the presence of two separate, mechanically stable branches in the P(V) equation of state, one can conclude that framework flexibility has been successfully manifested in M(NDIDP) MOFs at room temperature. At atmospheric pressure, i.e. P = 0.1 MPa, two (meta)stable intersections with negative $\partial P/\partial V$ values are found for Zn(NDIDP) and Co(NDIDP), whereas only one is found in the lower volume range for Fe(NDIDP). These results are in accordance with the DFT-derived E(V)profiles, as they confirm the thermodynamic (meta)stability of both lp and np phases under ambient conditions for Zn(NDIDP) and Co(NDIDP). Correspondingly, in Fe(NDIDP), the P(V)equation of state shows thermodynamic stability only for the np phase at atmospheric pressure.

Next, the critical pressures needed to induce structural transitions between the lp and np phases were extracted. From the P(V) equation of state, they can be determined by identifying the local minimum and maximum in the pressure profile. In Zn(NDIDP), the equation of state shows a pressure minimum at -234 MPa, and a maximum at 3 MPa, which can be identified with the np-to-lp and lp-to-np transition pressures, respectively. As for Co(NDIDP), these transition pressures are -224 and 13 MPa. For the softer Fe(NDIDP) MOF, these pressures amount to -217 and -42 MPa. Hence, while the lp-to-np transition in either Zn(NDIDP) and Co(NDIDP) can be easily induced by applying an external pressure of 3 and 13 MPa, the same transition in Fe(NDIDP), as well as the np-to-lp transitions in all three materials, would require negative pressures. Note that such negative pressures are not yet practically achieved in MOFs. Nonetheless, one can envision that finite pulling forces of corresponding magnitude (applied via embedding the MOF into a flexible membrane, etc.) would induce the np-to-lp transition.

An interesting feature found in the P(V) equations of state is an anomaly in the curve around $V = \sim 1650$ Å³, at elevated pressures of ca. 1.5 GPa (Figure S4). Structural configurations to the right of the anomaly (higher volume, Figure S5) closely resemble the *np* phase and retain the 4-fold rotational symmetry around the 1D metal clusters. On the other hand, configurations to the left of the anomaly (lower volume, Figure S6) are all rotationally distorted around the metal clusters, and the 4-fold symmetry is reduced to 2-fold symmetry. This means that at higher pressures, the framework becomes rotationally distorted to alleviate the stress being exerted. Note that the reduction of rotational symmetry interestingly improves the parallel alignment between adjacent NDI moieties, which could be more favorable for π orbital overlap and CT. As such, it can be conjectured that by applying pressures of ~1.5 GPa or higher, rotational distortions could be induced in M(NDIDP), which could potentially lead to further enhancements in their electrical conductivity. Here, we additionally note that in the case of Fe(NDIDP), a separate P(V) curve was observed for the volume range between 3250 and 3800 Å³ (see **Supporting Information**). Corresponding structural configurations are shown in **Figure S7**, and they are all found to be sheared in the *z*-axial direction compared to the main configurations. As these configurations deviate away from the main mode of flexibility under interest here (i.e., expansion/contraction based on the wine rack motif), we did not further investigate them. Nonetheless, this hints that other modes of flexibility could also exist in M(NDIDP), giving rise to more structural configurations with their own, unique NDI arrangements and resulting CT properties.

Electrical Conductivity of M(NDIDP) and Configuration-Dependent CT

Following the thorough affirmation of their framework flexibility, M(NDIDP) MOFs were analyzed to determine whether or not electrical conductivity has been successfully induced. In order for a MOF to harness conductivity, it must be engineered with long-range CT pathways for electron or hole transport.^{11,12} As for M(NDIDP), the installation of redox-active NDI as the central linker moiety and their closely packed arrangement within the framework are expected to provide the necessary CT pathways. The presence of valid CT pathways in a MOF can be deduced from its electronic band structure, where dispersive frontier bands would be present from significant orbital overlap between the proposed pathway components. Such band dispersions give rise to small effective carrier mass during band-like charge transport, which then guarantees high carrier mobility for observable conductivity. As such, DFT electronic band structures of each of the M(NDIDP) MOFs were calculated and analyzed to determine whether our modular design approach has successfully secured such pathways in these materials.

Electronic band structures and projected density of states (PDOS) of Co(NDIDP) are presented in **Figure 5**, and those of Zn(NDIDP) and Fe(NDIDP) are shown in **Figures S8** and **S9**. Band structures are drawn along the Γ -X and Γ -Z high symmetry paths, corresponding to

the directions in which close-packing of the NDI units was previously observed. Given their dissimilar NDI arrangements, calculations were performed for both lp and np phases. Remarkably, band structures of all three M(NDIDP)s in both phases exhibit notable dispersion in their conduction band minimum (CBM), which is ideal for n-type CT. From the PDOS and band-decomposed charge density plots (Figure S10), sole contributors to the CBM are revealed to be the C and O atoms of the NDI moieties. This shows that the CBM of M(NDIDP) MOFs is exclusively composed of the orbitals provided by NDI. Dispersion of the CBM hence arises from the orbital overlap between closely packed NDI units, which effectively completes the CT pathways in M(NDIDP) MOFs as intended by our modular design approach. Band gaps (Eg) of the MOFs under the two phases are found well within the semiconducting range, with values ranging between 0.16 and 2.39 eV (see Figures 5, S8, and S9). As such, all three MOFs in both *lp* and *np* phases are predicted to show electrical conductivity, which can be triggered by populating the CBM with electrons via excitation or n-doping of the system. Note that large dispersions are also present in the valence band maximum (VBM) in some cases, especially for Fe(NDIDP). Such VBM dispersions are suspected to be artefacts of the DFT methods adopted in our study, and should not be directly interpreted as indications of efficient hole transport (see Supporting Information).



Figure 5. Electronic band structures and projected density of states (PDOS) of (a) Co(NDIDP)–lp and (b) Co(NDIDP)–np. Conduction band minima corresponding to NDI are shown separately in the bottom right. Green bands are filled valence bands, and blue bands are empty conduction bands. Solid and dotted bands (with lighter color) correspond to up and down spins.

In M(NDIDP)–lp, dispersion of the CBM is limited to the Γ –Z path, and the band remains strictly flat along the Γ –X path. This is in accordance with the internal NDI arrangement observed in the lp phase, where parallel-displaced packing along the *z*-axis was found. As such, CT in M(NDIDP)–lp is expected to propagate one-dimensionally along this axis. Dispersion is measured to be 83, 130, and 212 meV for Zn, Co, and Fe(NDIDP)–lp, respectively. Here, we note the increasing trend of CBM dispersion in the presented order of MOFs. This is also attributed to the different binding tendencies of metal ions, which becomes softer from Zn to Co, then Fe. Softer metal ions would allow the MOF to further distort its structure, better optimizing the orbital overlap between stacked NDI moieties. This is indeed corroborated by their internal NDI arrangements (see **Table S2**), where the adjacent NDI moieties become more parallelly aligned in the same order. Electron effective mass at the CBM edges (m_e^*) were also calculated and are presented in **Table 1**. Although m_e^* values along the Γ –Z path in M(NDIDP)–lp are relatively high, they still constitute significant improvements over the non-conducting case (i.e. $m_e^* \rightarrow \infty$), and hence effectively validating the presence of CT pathways.

M(NDIDP)	large pore (<i>lp</i>)		narrow pore (<i>np</i>)	
	Γ–Χ	Γ–Ζ	Γ–Χ	Γ–Ζ
Zn		83 meV	73 meV	200 meV
		$m_{\rm e}^{*} = 4.37m_0$	$m_{\rm e}^{*}=3.02m_{0}$	$m_{\rm e}^{*} = 1.51 m_0$
Со	$\sim 0 \text{ meV}$	130 meV	121 meV	211 meV
	$m_{\rm e}^* \rightarrow \infty$	$m_{\rm e}^* = 2.76 m_0$	$m_{\rm e}^{*}=2.36m_{0}$	$m_{\rm e}^{*} = 1.64 m_0$
Fe		212 meV	403 meV	633 meV
		$m_{\rm e}^* = 1.71 m_0$	$m_{\rm e}^* = 1.17 m_0$	$m_{\rm e}^* = 0.73 m_0$

Table 1. CBM dispersion values and corresponding effective masses of electron (m_e^*) obtained from the band structures of M(NDIDP)–*lp* and M(NDIDP)–*np*. (*m*₀: electron rest mass)

In discussing the case of M(NDIDP)–np, we first note the apparent splitting of the CBM at the Γ point. Such phenomenon arises from the presence of two NDI moieties along the *x*-axis within the chosen M(NDIDP) unit cell, which come in contact with one another in the np phase (see **Figure 3**). Consequently, two pairs of NDI moieties are formed, and their orbital interactions within the cell cause the CBM to undergo band splitting at the Γ point. Note that a single NDI moiety serves as one electronic unit, yet the unit cell of M(NDIDP)–np contains

two electronic units along both x and z-axis (see **Supporting Information**). Therefore, the CBM should rather be understood as having been folded due to the presence of multiple electronic units within the unit cell.⁴⁹ This interpretation is further substantiated by the convergence of CBM and CBM+1 band energies observed at the X and Z symmetry points. As such, set of bands all corresponding to the lowest unoccupied orbitals of NDI were collectively regarded as the CBM, and band dispersion and m_e^* were calculated accordingly.

The CBM of M(NDIDP)–*np* along the Γ –Z path shows significant dispersions of 200, 211, and 633 meV for Zn, Co, and Fe(NDIDP)-np, respectively, which constitutes an up to 3fold increase compared to the *lp* phase. Such enhancements are attributed to the tighter packing of NDI along the z-axis (see Table S2), which drastically improves the orbital overlap between adjacent NDI moieties. Interestingly, band dispersion is newly observed along the Γ -X path as well in all three MOFs. This confirms that 2D herringbone packing of NDI exhibited by M(NDIDP)-np can provide sufficient orbital overlap along both dimensions, resulting in twodimensional CT. Dispersion along the Γ -X path are 73, 121 and 403 meV for Zn, Co, and Fe(NDIDP)–*np*, which are comparable to the Γ –Z path in M(NDIDP)–*lp*. Along both high symmetry paths, the previously observed correlation between binding tendencies of metal ion and the resulting band dispersion are found to persist. Trends in the band dispersion are correspondingly carried over to m_e^* (see **Table 1**), where the m_e^* is notably reduced along the Γ -Z path compared to M(NDIDP)–*lp*, and finite m_e^* values are newly observed along the Γ -X path. The smallest m_e^* of 0.73 m_0 obtained for Fe(NDIDP)–np along the Γ –Z path is noteworthy, as it is smaller than the electron rest mass and comparable to the values reported for other known conductive MOFs.^{13,50,51} Note that Eg of Co(NDIDP) decreases from 1.78 to 1.58 eV as the MOF transitions from lp to np phase. Similar reduction of Eg is observed for the other two MOFs (see Figures S8 and S9), where Eg decreases from 2.39 to 2.20 eV, and from 0.54 to 0.16 eV in Zn and Fe(NDIDP), respectively. These changes in Eg are consistent with what was previously reported for MIL-47 and MIL-53.³¹

Results thus far demonstrate that while significant conductivity is predicted for all three M(NDIDP)s, their precise CT properties vary drastically depending on their structural configurations. Between the lp and np phases, differences in the CT properties are characterized by an up to 3-fold increase in band dispersion (2.9-fold reduction in m_e^*) and a transition from 1D to 2D transport. Such differences are explained by changes in the internal NDI arrangement,

which progresses from 1D parallel-displaced packing to 2D herringbone packing as the MOF transitions from the *lp* to the *np* phase.

Further Exploration of Configuration-Dependent CT

Based on the findings thus far, we conclude that two material properties, namely flexibility and conductivity, have been successfully instilled and synergistically combined in M(NDIDP), resulting in tunable electrical conductivity. Next, one could envisage that conductivity of M(NDIDP) could be further tuned by taking advantage of their flexibility and exploring more of their allowed structural configurations. Hence, additional structural configurations of M(NDIDP) were selected and analyzed for their unique CT properties. Two intermediate configurations located between the lp and np phases, herein referred to as int1 and int2, were first extracted (light blue points in Figure 2). Additionally, "squeezed" (sq) and "expanded" (exp) configurations were obtained as structures smaller than the np phase or larger than the *lp* phase, respectively, whilst exhibiting an energy difference of less than 1.5 eV from the corresponding phases at 0 K (light green and light magenta points in Figure 2). While these configurations are not thermodynamically accessible under ambient conditions, they are still of considerable interest since external stimuli (e.g., guest loading, mechanical stress, isotropic pulling) could effectively lead to their stabilization. Band structures were subsequently calculated for the chosen configurations in Co(NDIDP) (Figure S11 and S12), and the results are presented in Figure 6 and S13.

In Co(NDIDP)–*exp* (Figure 6a), where the MOF is further expanded past the *lp* phase (Figure 6b), significant enhancement in CT is interestingly observed. While the CBM remains flat along the Γ –X path, band dispersion along the Γ –Z path increases from 130 to 280 meV, and m_e^* decreases from 2.76 m_0 to $0.91m_0$. E_g is found to be 1.73 eV (see Figure S13), which is slightly lower than Co(NDIDP)–*lp*. It is also noted that energy of the CBM at the Z point is higher than the energy at the Γ point, whereas the inverse trend was observed in the *lp* phase. These phenomena are collectively attributed to the reconfiguration of NDI moieties accompanying further framework expansion (see Table S2). The internal NDI arrangement of Co(NDIDP)–*exp* exhibits a notable decrease in the *z*-axial stacking distance from 3.26 (*lp*) to 3.12 Å (*exp*), which improves the orbital overlap between NDI moieties and therefore explains the increased band dispersion. Concurrently, the planar alignment offset surges from 5.44 (*lp*)

to 6.63 Å (*exp*). Such a large difference in NDI alignment is significant enough to induce a change of sign in the transfer integral between adjacent moieties, hence inverting the energy trend along the Γ -Z path. As presented in **Figure S14**, notably enhanced CT properties are consistently observed in Zn(NDIDP)–*exp* and Fe(NDIDP)–*exp* as well (structures shown in **Figure S15**) compared to their *lp* counterparts. This effectively demonstrates that if M(NDIDP) can be further expanded past the *lp* phase, for instance by an isotropic pulling of the material, their electrical conductivity would be notably enhanced.



Figure 6. Evolution of the conduction band minimum (CBM) of Co(NDIDP) upon decreasing volume under all of the structural configurations considered. Full electronic band structures of the additional structural configurations (*exp*, *int1*, *int2*, *sq*) are shared in **Figure S13**. Solid (blue) and dotted bands (lighter blue) correspond to up and down spins.

Next, two intermediate *int1* and *int2* configurations (**Figures 6c** and **6d**) between the metastable *lp* and *np* phases were analyzed, which allow one to study the evolution of the CBM as Co(NDIDP) undergoes the *lp*-to-*np* transition. Along the Γ -X path, dispersion is found to monotonically increase as follows: 0 meV (*lp*), 5 meV (*int1*), 85 meV (*int2*), and 121 meV (*np*). This trend is straightforwardly explained by the effect of framework contraction on the internal NDI arrangement of Co(NDIDP) (see **Tables S2**). Pairs of NDI moieties along the *x*-axis become closer to each other from one configuration to the next, leading to the gradual onset of their orbital overlap and CT in the *x*-axial direction. Along the Γ -Z path, band dispersions are 130 meV (*lp*), 181 meV (*int1*), 86 meV (*int2*), and 211 meV (*np*). Curiously, band dispersion decreases in Co(NDIDP)–*int2* while the other three configurations show an increasing trend. This can be attributed to the large *z*-axial stacking distance of 3.6 Å observed in Co(NDIDP)–*int2* as opposed to 3.26 Å (*lp*), 3.40 Å (*int1*), and 3.12 Å (*np*), which has significantly increased to better accommodate the pair interactions newly taking place in the *x*-axial direction. These

results demonstrate that correlation between the structural transition of M(NDIDP) and the resulting CT properties is nonlinear and rather complex. Nonetheless, a gradual transition from 1D to 2D transport is undoubtedly observed, establishing that the conductivity of M(NDIDP) MOFs can be finely tuned by accessing the intermediate configurations between the *lp* and *np* phases.

Remarkably, contraction of Co(NDIDP) past the np phase (Figure 6e) to Co(NDIDP)sq (Figure 6f) also results in vastly improved CT properties. In Co(NDIDP)-sq, CBM dispersion is measured to be 330 meV and 738 meV along the Γ -X and Γ -Z paths. These values constitute substantial CT enhancement over the case of Co(NDIDP)-np, especially along the Γ -Z path where the dispersion has increased by a factor of 3.5. Large dispersion in the CBM leads to significantly smaller m_e^* , where the effective mass is reduced from 2.36 m_0 to 1.84 m_0 along the Γ -X path, and from 1.64 m_0 to 0.37 m_0 along the Γ -Z path. 0.37 m_0 is one of the lowest carrier effective masses that has been observed in a MOF, to the best of our knowledge. Eg is significantly decreased from 1.58 eV (np) to 1.00 eV (sq) (see Figure S13). Again, such markedly improved CT properties are associated with the internal NDI arrangement of Co(NDIDP)-sq (see Table S2). In this structural configuration, NDI moieties are even more tightly packed together: the z-axial stacking distance is reduced from 3.12 to 2.14 Å, and the NDI-H distance is also reduced from 2.47 to 2.16 Å. This effectively maximizes the orbital overlap between adjacent NDI moieties, which further boosts CT and the resulting conductivity. Note that the extreme contraction of Co(NDIDP) into the sq configuration leads to an uneven spacing between NDI moieties in the z-axial direction (see Figure S16), which creates an opening between the folded bands of CBM at the Z point. Based on these results, we conclude that the highest conductivity of Co(NDIDP) can be attained by further contracting or compressing the MOF past the *np* phase.

Similar CT enhancements were not observed in the readily procured sq configurations of Zn(NDIDP) and Fe(NDIDP) (**Figures S17 and S18**). In the case of Zn(NDIDP)–sq, only a 1.33-fold increase in the band dispersion is observed compared to the np phase. As for Fe(NDIDP)–sq, band dispersion has rather diminished compared to Fe(NDIDP)–np along both symmetry paths. We attribute these differences in CT properties to the visibly different structural configuration of Co(NDIDP)–sq (see Figure **S12**) as opposed to Zn or Fe(NDIDP)– sq (see Figure **S17**), where a counter-clockwise rotational distortion around the 1D metal cluster is observed. Such rotational distortion around the 1D metal cluster has also been computationally predicted for the low volume configurations of other wine rack MOFs in previous studies.^{52–55} As conjectured previously, this distortion induces a much smaller angle between the edge-to-face NDI pairs compared to Co(NDIDP)–*np*, Zn(NDIDP)–*sq* or Fe(NDIDP)–*sq* (see **Table S2**), which is highly beneficial for π -orbital overlap. As a result, notable CT enhancement of Co(NDIDP)–*sq* is attributed to the observed rotational distortion. Note that similarly distorted structural configurations of Zn(NDIDP) and Fe(NDIDP) were nonetheless observed from the MD simulations (see **Figure S6**) for lower unit cell volumes (< 1650 Å³) at room temperature. Hence, it is predicted that framework contraction past the *np* phase could lead to the highest conductivity in all three M(NDIDP) MOFs, as long as the rotational distortion around the metal cluster simultaneously takes place. Altogether, our analysis effectively demonstrates that electrical conductivity of M(NDIDP) is highly tunable, and many other structural configurations with unique CT properties exist for these flexible MOFs.

Conclusion

In this study, we have demonstrated that framework flexibility and electrical conductivity can be synergistically combined into a single MOF via a modular design approach. The designed MOFs, M(NDIDP) with M = Zn, Co, Fe, show significant framework flexibility and are capable of pressure-induced structural transitions at low pressures and room temperature. M(NDIDP) MOFs also exhibit configuration-dependent CT properties, which are characterized by transition from 1D to 2D transport and up to 5.7-fold increase in band dispersion (7.5-fold reduction in effective carrier mass) between different structural configurations. As a result, tunable electrical conductivity is predicted for the M(NDIDP) MOFs, where the degree of CT in the MOF can be controlled via application of external stimuli. The design of flexible and electrically conductive MOFs performed in this work demonstrates the practicability of simultaneously inducing multiple material properties in the same MOF via the modular design approach. We anticipate that our design approach can be extended to discover more flexible and conductive MOFs for adsorptive sensing, conformal electronics, molecular switches, and many other relevant applications that may benefit from stimuli-induced changes in electrical conductivity.

Computational Methods

DFT Calculations

Kohn-Sham density functional theory (DFT) calculations were performed to obtain the optimized configurations of each of the M(NDIDP) MOFs and analyze the resulting electronic structures. All DFT calculations in this work were carried out with the Vienna *Ab initio* Simulation Package (VASP).^{56–58} The projector augmented wave (PAW) method^{59,60} was used to describe the interactions between the ions and electrons. The exchange-correlation term was described within the generalized gradient approximation (GGA) using the Perdew-Burke-Ernzerhof (PBE) functional.⁶¹ In the newly designed MOFs, it is important to accurately account for the van der Waals interactions between the NDI moieties given their high degree of aromaticity. Hence, DFT-D3 dispersion corrections with Becke-Johnson damping were employed.⁶² Γ -centered k-point grids with spacings less than 0.3 Å⁻¹ were used in the calculations. The kinetic energy cutoff was set at 600 eV, and self-consistent field (SCF) convergence criteria was fixed at 1×10⁻⁶ eV. Gaussian smearing was performed with a sigma value of 0.01 Å. Given the presence of unpaired d-orbital electrons, spin polarization was considered for Co(NDIDP) and Fe(NDIDP). High-spin ferromagnetic ordering along each 1D metal cluster was observed to be the most stable, and hence used to report the system energies.

Geometry optimizations were performed to construct the E(V) profiles of each material. In each optimization, a volume constraint was applied. That is, the total volume of the unit cell was fixed, but the internal atomic coordinates and the lattice parameters were relaxed until the energy minimum was found. Relaxation was continued until the Hellmann-Feynman forces became lower than 0.01 eV/Å for all the atoms. After successfully optimizing the MOF at a given volume, length of the lattice parameter *a* was incrementally increased or decreased to provide the starting configuration for the adjacent volume point. This procedure was repeated until the entire volume range of interest was covered. Electronic band structures were subsequently calculated for the configurations of interest. While the PBE functional can provide an accurate description of the structure of these materials, it is known to underestimate the band gaps of semiconducting materials. Hence, the HSE06 hybrid functional⁶³ was used to more accurately describe the band structures. In doing so, revised D3 damping parameters for HSE06 reported by Moellmann and Grimme were used.⁶⁴ Ten equidistant k-points within the first Brillouin zone were sampled along the high symmetry paths of interest, i.e. the Γ -X and Γ –Z paths. Carrier effective masses were calculated with the sumo package,⁶⁵ using a least-squares fit with five sample points and assuming non-parabolicity of the bands.

Molecular Dynamics Simulations

Force field-based molecular dynamics (MD) simulations were performed to study the pressure-induced structural transitions of M(NDIDP). To this end, for each of the DFT-optimized structures, first the Hessian matrix was calculated with VASP using 0.015 Å displacements and an SCF convergence criterium of 1×10^{-8} eV for improved accuracy. The *ab initio* periodic Hessian and optimized structure were then taken as inputs to the QuickFF software^{66,67} to derive the covalent force field parameters for interactions including cross terms and using the force field atom types provided in **Figure S19**. Gaussian-like atomic charges were used to describe the electrostatic interactions, and were determined using the Minimal Basis Iterative Stockholder (MBIS) partitioning scheme as implemented in HORTON.⁶⁸ To this end, GPAW calculations^{69,70} were performed on the optimized geometries to obtain the all-electron density necessary for the MBIS scheme. The radii of the Gaussian charges were obtained by following the fitting approach suggested by Chen and Martínez.⁷¹ Finally, the MM3 Buckingham potential⁷² was used to account for the van der Waals interactions.

Using these force fields, MD simulations were performed in Yaff for a variety of volumes to obtain the P(V) equations of state.⁷³ Simulations were conducted for $1 \times 1 \times 2$ supercells of the M(NDIDP) MOFs in the $(N, V, \sigma_a = 0, T)$ ensemble, where σ_a refers to the deviatoric stress component of the stress tensor.⁴⁶ In this ensemble, the unit cell *V* is constrained, but the simulation cell shape can fluctuate freely. From these simulations, the average pressure the material exerts on its environment is extracted. This average pressure equals the external pressure exerted on the material in mechanical equilibrium and can thus be used to construct the macroscopic pressure-versus-volume equations of state. A Nosé-Hoover chain thermostat^{74,75} with a chain length of 3 and a 0.1 ps relaxation time was used to control the system temperature. σ_a was controlled using the Martyna-Tobias-Tuckerman-Klein barostat⁷⁶ with a 1 ps relaxation time. The velocity Verlet integration algorithm was used for the van der Waals interactions with a smooth cutoff. For the electrostatic interactions, an Ewald summation was performed with a real-space cutoff of 15 Å, a splitting parameter of 0.213 Å⁻¹, and

reciprocal space cutoff of 0.32 Å⁻¹. Each simulation consisted of a 0.5 ns equilibration run followed by a 0.5 ns production run. The obtained P(V) equations of state were locally smoothened using the Savitzky-Golay filter.⁷⁷

Supporting Information

The Supporting Information is available free of charge.

Structures of M(NDIDP) in *lp* and *np* phases; P(V) equation of state in the low V range and corresponding configurations; Secondary phase structures of Fe(NDIDP); Band structures of Zn(NDIDP) and Fe(NDIDP) in *lp* and *np* phases; Band-decomposed charge density plots of the CBM of M(NDIDP); Additional structural configurations of Co(NDIDP) and their band structures; Structures and band structures of the *exp* configuration of Zn and Fe(NDIDP); NDI arrangement in the z-axial direction for Co(NDIDP)-*sq*; Structures and band structures of the *sq* configuration of Zn and Fe(NDIDP); Atom type assignment in M(NDIDP) for force field generation; Lattice parameters of M(NDIDP); Internal NDI arrangements of M(NDIDP); Flexibility and conductivity of Zn(NDIDP)-CH₃; Secondary phase configurations of Fe(NDIDP); VBM dispersion in M(NDIDP); CBM splitting/folding in smaller volume configurations; Data availability (PDF)

Notes

The authors declare no competing financial interest.

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TOC Graphic

