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Published in:
Nano Letters

DOI:
10.1021/acs.nanolett.8b02062

Published: 12/09/2018

Document Version
Publisher's PDF, also known as Version of record

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Please cite the original version:
Kumar, A., Banerjee, K., Foster, A. S., & Liljeroth, P. (2018). Two-Dimensional Band Structure in Honeycomb Metal-Organic Frameworks. Nano Letters, 18(9), 5596-5602. https://doi.org/10.1021/acs.nanolett.8b02062
Two-Dimensional Band Structure in Honeycomb Metal–Organic Frameworks

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ABSTRACT: Two-dimensional (2D) metal–organic frameworks (MOFs) have been recently proposed as a flexible material platform for realizing exotic quantum phases including topological and anomalous quantum Hall insulators. Experimentally, direct synthesis of 2D MOFs has been essentially confined to metal substrates, where the strong interaction with the substrate masks the intrinsic electronic properties of the MOF. In addition to electronic decoupling from the underlying metal support, synthesis on weakly interacting substrates (e.g., graphene) would enable direct realization of heterostructures of 2D MOFs with inorganic 2D materials. Here, we demonstrate synthesis of 2D honeycomb MOFs on epitaxial graphene substrate. Using low-temperature scanning tunneling microscopy (STM) and atomic force microscopy (AFM) complemented by density-functional theory (DFT) calculations, we show the formation of a 2D band structure in the MOF decoupled from the substrate. These results open the experimental path toward MOF-based designer electronic materials with complex, engineered electronic structures.

KEYWORDS: Scanning tunneling microscopy (STM), metal–organic framework (MOF), cobalt, 4,4′-dicyanobiphenyl (DCBP), 9,10-dicyanoanthracene (DCA), epitaxial graphene

Metal–organic frameworks (MOFs) are an important class of materials that present intriguing opportunities in the fields of sensing, gas storage, catalysis, and optoelectronics.1–4 While there are a tremendous number of examples of three-dimensional, bulk MOFs, synthesis strategies for two-dimensional (2D), monolayer thick MOFs (also referred to by various names5 such as metal–organic coordination networks (MOCNs),6 surface-confined metal–organic networks (SMONs),7 metal–organic materials (MOMs), and metal–organic graphene analogues (MOGs)8) are more limited. These systems are drawing growing interest due to their very exciting properties, either in conjunction with other 2D materials or as a stand-alone platform for novel electronic materials with tunable properties.9–11

The synthetic flexibility and tunable electronic properties of MOFs stem from the choice of metal atoms, organic molecules, the linker chemistry and electronic and magnetic interactions among the building blocks.7,12–16 For example, it is possible to realize honeycomb and Kagome lattices that are expected to give rise to peculiar electronic properties. Compared to 2D covalent organic framework (COFs), 2D MOFs can incorporate metal centers with high spin–orbit coupling and magnetism, which are important building blocks for realizing exotic materials such as topological and quantum anomalous Hall insulators. It has been theoretically predicted that 2D MOFs can be turned into topological insulators by adding sufficiently strong spin–orbit interactions through the choice of the metal atom.15,16–19 This suggests MOFs as a tunable platform for realizing organic quantum materials.9,13–22 Since 2D MOFs go through reversible bond-forming reactions, their on-surface synthesis on weakly interacting substrates is easier compared to 2D COFs.21 However, experimental study of these materials requires synthesis methods that yield monolayer MOFs on weakly interacting substrates such that their intrinsic electronic properties can be probed.

Procedures for direct growth of 2D MOFs exist, e.g., through synthesis on the air–liquid interface or by chemical vapor deposition (CVD) in ultrahigh vacuum (UHV) conditions.7,22–27 CVD growth is typically carried out on metallic substrates where various types of frameworks have been studied in detail.7,12 However, the strong hybridization with the underlying substrate masks the intrinsic properties of the frameworks. This problem has been overcome in the case of single molecules by the use of ultrathin insulating films28–31 and inert 2D materials such as graphene10,32–35 that electronically decouple the molecule from the metallic substrate. Unfortunately, self-assembly and, in particular, on-surface

Received: May 21, 2018
Revised: August 8, 2018
Published: August 22, 2018
chemical reactions are a virtual terra incognita on weakly interacting, noncatalytic substrates\(^{10,36-40}\) and therefore the experimental observation of the intrinsic electronic properties of 2D MOFs has been elusive. Here, we demonstrate the controlled synthesis of high quality honeycomb MOFs on epitaxial graphene using different organic linkers (dicyanobiphenyl, DCBP, and dicyanoanthracene, DCA) with cobalt metal atoms. We characterize the structures using low-temperature scanning tunneling microscopy (STM) and atomic force microscopy (AFM). We are able to access the intrinsic electronic properties of 2D MOFs and demonstrate the formation of a strongly coupled 2D electronic system in the DCA-Co MOF by scanning tunneling spectroscopy (STS) measurements complemented by density-functional theory (DFT) calculations.

Figure 1 shows the structure of the honeycomb MOFs—DCBP\(_2\)Co\(_2\) and DCA\(_2\)Co\(_2\)—synthesized on epitaxial graphene grown on Ir(111) (G/Ir(111)), experimental details are given in the Supporting Information. Briefly, after synthesis of epitaxial graphene,\(^{31-43}\) we sequentially deposit the molecules and cobalt atoms at various temperatures. After deposition of a submonolayer coverage of the molecules, adding cobalt atoms at slightly elevated temperatures (details in the Supporting Information) results in formation of a strongly coupled 2D electronic system in the DCA-Co MOF by scanning tunneling spectroscopy (STS) measurements complemented by density-functional theory (DFT) calculations.

**Figure 1.** Overview of two MOFs. (a) An STM overview image of a honeycomb DCBP\(_2\)Co\(_2\) MOF on G/Ir(111) surface. Scale bar is 10 nm. Imaging parameters: 1.23 V and 3.3 pA. (b) Constant height frequency shift, \(\Delta f\), nc-AFM image of DCBP\(_2\)Co\(_2\) MOF acquired with a CO-terminated tip. Scale bar is 1 nm. (c) DFT-simulated structure of DCBP\(_2\)Co\(_2\) MOF on graphene. (d) STM topography image of DCA\(_2\)Co\(_2\) MOF. The scale bar is 1 nm. Imaging parameters: \(-1\) V, 15 pA. (e) DFT simulated structure of DCA\(_2\)Co\(_2\) MOF on graphene. Red parallelograms indicate the unit cells.
stoichiometry, DCA forms only mononuclear 3-fold (DCA$_3$Co) complexes which is unambiguously confirmed by nc-AFM imaging (see Supporting Information, Figures S4 and S5). We attribute the absence of 4-fold DCA$_4$Co to a larger steric hindrance compared to that of a 4-fold structure of DCBP$_4$Co.

Figure 2 compares $dI/dV$ spectra recorded on single molecules and the corresponding single metal−organic complexes. As shown in Figure 2a, $dI/dV$ spectrum recorded on a DCBP molecule shows a shoulder at 2.7 V corresponding to the lowest unoccupied molecular orbital (LUMO) (see Supporting Information, Figure S5). The peak due to the highest occupied molecular orbital (HOMO) of the molecule is not accessible within the recorded bias range of the spectrum and it lies at a bias lower than $-1.5$ V. On a single DCBP$_4$Co complex, $dI/dV$ reveals two peaks at 2.3 V and $-1$ V with

Figure 3. Electronic properties of honeycomb DCBP$_3$Co$_2$ MOF. (a) STS recorded on honeycomb DCBP$_3$Co$_2$ at the positions shown in the inset. (b) Calculated band structure and total PDOS of DCBP$_3$Co$_2$ MOF. (c, d) Experimental (panel c) and simulated STM images (panel d) at the energies indicated in the figure. Scan size is $6.2 \times 4$ nm$^2$.

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corresponding electronic states located on DCBP and Co center, respectively. On the basis of the bias-dependent STM imaging of the 4-fold phases and $dI/dV$ spectroscopy of DCBP molecule as a function of distance from Co center (see Supporting Information, Figures S5 and S6), it is clear that the peak 2.3 V originates from the LUMO of the DCBP molecule. The shift of the molecular LUMO toward the Fermi level by 0.4 V indicates that there is an electrostatic shift of the orbital energy due to the Co atom of the complex and other complexes present in the vicinity.

d$I$/d$V$ spectra on single DCA molecule on G/Ir(111) also reveal a peak at 1.8 V as shown in Figure 2b. The gating effect due to Co atoms is also observed in the DCA$_n$Co$_3$ single complexes. The $dI/dV$ spectra recorded on DCA of the complex shows that the LUMO shifts down to 750 mV and three satellite vibronic peaks also become visible. The vibronic mode energy of $\sim$200 mV fits well with the expected energy of the C−C vibration.49,50 The assignment of the peak to the molecular LUMO is also evident from the STM images (see Supporting Information Figures S4 and S7). If there was (integer) charge transfer to DCBP or DCA molecules in the respective complexes, the LUMO peak would split to singly occupied/unoccupied molecular orbital (SOMO/SUMO) peaks at negative and positive bias.51,52 $dI/dV$ spectra recorded on Co center of the complex shows an additional shoulder at the onset of the peak. We attribute this shoulder to the metal-state as the metal center becomes brighter in the STM images at sample bias beyond 0.7 V (see Supporting Information Figure S4).

Figure 3a shows $dI/dV$ spectra recorded on DCBP molecule in the DCBP$_3$Co$_2$ MOF has a peak at 1.44 V which we ascribe to the elastic LUMO peak with corresponding vibronic replica at 1.62 V. The small line-width of DCBP LUMO and the observation of satellite vibronic peaks indicate that the intermolecular electronic coupling in the framework is weak such that we have isolated molecular electronic states. The spectrum recorded on the Co center reveals a faint peak at $-1150$ mV, which is visible in the background corrected spectrum (see Supporting Information Figure S8). The state is localized only at the metal-center.

We have used DFT to calculate the band structure of the DCBP$_3$Co$_2$ MOF as shown in Figure 3b for the antiferromagnetic ground state. While DFT underestimates the band gap, it correctly captures the nature of the lowest lying bands: the occupied states have a stronger metal character compared to the unoccupied states, which are mostly composed of the ligand states (Figure 3c,d). The enhanced contrast on the metal atoms and ligands can be seen at negative and positive bias, respectively, compared to the STM topography in the gap (Figure 3c, middle panel). However, DFT seems to overestimate the bandwidth of the unoccupied ligand-derived states compared to the experiment. This could be related to how well the torsional angle between the phenyl rings of DCBP molecule is estimated by DFT as this is known to control...
the $\pi-\pi$ conjugation within the backbone of the molecule.\textsuperscript{53} The coupling is enhanced for the planar, smaller DCA linker as demonstrated below.

A substantial in-plane electronic hybridization and formation of energy bands with significant width in DCA$_3$Co$_2$ MOF is evidenced by $dI/dV$ spectroscopy and spatially resolved $dI/dV$ maps. $dI/dV$ spectrum (Figure 4, blue) recorded at the center of the ring (constituting six DCA lobes) has three peaks at 260, 480, and 860 mV. Considering the separation between the first and the second peak, $\sim 220$ mV, the second peak could still be interpreted as a vibronic satellite. However, the separation between the second and the third peak rules out vibronic origin. The spectrum at the center of DCA molecule (red curve) shows that the first peak shifts to 360 mV, while that on cobalt (cyan curve) has the first peak at 320 mV. The systematic evolution of the spectra across the framework is shown in the Supporting Information Figure S9. Comparison of the spectra at the lobe and center of DCA and cobalt in the DCA$_3$Co$_2$ MOF to that of DCA$_3$Co single complex indicates that there is additional intensity in the MOF at energies higher than 700 mV (see Supporting Information Figure S10). As the $dI/dV$ signal is directly proportional to the local density of states (LDOS), this is direct evidence of additional electronic states. Further, we have recorded $dI/dV$ maps of the same area at different energies as shown in Figure 4c. The LUMO lobes on the DCA molecules are the brightest feature at lowest energies (200 mV), while the states on the Co sites are more prominent at $\sim 300$ mV. At intermediate energies, 410 and 450 mV, there exist extra bright features in the $dI/dV$ maps superimposed on the existing framework. These features are likely to result from the formation of standing wave patterns due to the scattering of 2D electron waves from the boundaries of the finite sized DCA$_3$Co$_2$ MOF domain. This is supported by the Fourier transforms (FFT) of a large area $dI/dV$ maps (Figure 4d). Apart from the 12 outer spots (red circles) corresponding to the honeycomb structure of DCA$_3$Co$_2$ MOF, there exists internal structure which evolves continuously with the bias. The spots in this quasi-particle interference pattern correspond to scattering vectors connecting the initial and final states of the scattering process at the given energy. In addition to this joint density-of-states, they contain information on the nature of the allowed scattering processes.\textsuperscript{54–56} While quantitative analysis of the experimental patterns is difficult due to the other overlapping peaks stemming from the geometry as well as the limited sample size (number of repetitive unit cells), they indicate the formation of an extended electron system with considerable dispersion (bandwidth).

The calculated electronic band structure using DFT for the symmetric, ferromagnetic DCA$_3$Co$_2$ framework without graphene is shown in Figure 4b. The band structure on graphene has additional bands arising from the graphene and the slight shifts and splittings of the MOF states are due to the residual interaction with the graphene (see Supporting Information, Figure S11). In line with the calculations done for DCA$_3$Cu$_4$ and DCA$_3$Mn$_2$ MOFs,\textsuperscript{18,20} the band structure of DCA$_3$Co$_2$ MOF has a number of flat-bands and Dirac cones. While the antiferromagnetic structure is slightly lower in energy (by 0.05 eV), the ferromagnetic state better reproduces the experimental results (see also discussion on this in the computational methods section of the Supporting Information). LDOS maps of DCA$_3$Co$_2$ MOF for both spin configurations look very similar except for the map at $-0.15$ V corresponding to antiferromagnetic configuration (see Supporting Information, Figure S12b), where prominent metal states stand out. These are not observed experimentally, suggesting a ferromagnetic spin configuration. The presence of a large gap between $-0.7$ and $-1.2$ eV in the calculated band structure and lack of states below the Fermi energy in the $dI/dV$ spectra until $-1.5$ V suggest that the energy corresponding to the experimental Fermi level lies below the flat band at energy $-0.7$ V. The DFT calculation suggests that the bottom of the conduction band consists of a flat band and a Dirac cone stemming from the DCA states and the Kagome symmetry of the lattice. Subsequently, at higher energies, there are also relatively flat bands originating mostly from the metal atom orbitals and a band with more mixed character. This overall picture is consistent with the experiments where we first see intensity on the molecules with metal states emerging at higher energies and an overall bandwidth of $\sim 1$ eV.

In summary, we have demonstrated direct synthesis of high-quality honeycomb MOFs on epitaxial graphene surface. While DCBP$_3$Co$_2$ MOF only has weak coupling between the building blocks, DCA$_3$Co$_2$ MOF shows significant in-plane hybridization resulting in the formation of 2D electronic states with significant bandwidth. These results open the experimental path toward MOF-based designer electronic materials with complex, engineered electronic structures. The direct growth of 2D MOFs on graphene outlines possibilities of heterostructures with inorganic 2D materials with potential applications in electronics, sensors, and catalysis.

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.nanolett.8b02062.

Experimental and computational methods and additional results (PDF)

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Notes

The authors declare no competing financial interest.

**ACKNOWLEDGMENTS**

This research made use of the Aalto Nanomicroscopy Center (Aalto NMC) facilities and was supported by the European Research Council (ERC-2017-AdG No. 788185 “Artificial Designer Materials”), and the Academy of Finland (Projects no. 305635 and 311012, and Centres of Excellence Program projects no. 284594 and 284621). A.S.F. has been supported by the World Premier International Research Center Initiative (WPI), MEXT, Japan and acknowledges use of the CSC, Helsinki for computational resources.

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