Scrutinizing the Pore Chemistry and the Importance of Cu(I) Defects in TCNQ-Loaded Cu₃(BTC)₂ by a Multitechnique Spectroscopic Approach

Christian Schneider,† Matthias Mendt,‡ Andreas Pöppl,† Valentina Crocella,*§ and Roland A. Fischer*†

†Department of Chemistry, Technical University of Munich, Lichtenbergstrasse 4, D-85748 Garching, Germany
‡Felix Bloch Institute for Solid State Physics, University Leipzig, Linnestrasse 5, D-04103 Leipzig, Germany
§Department of Chemistry, NIS and INSTM Reference Centre, Università di Torino, Via G. Quarello 15, I-10135 and Via P. Giuria 7, I-10125 Torino, Italy

ABSTRACT: Host–guest interactions control the fundamental processes in porous materials for many applications such as gas storage and catalysis. The study of these processes, however, is not trivial, even if the material is crystalline. In particular, metal–organic frameworks (MOFs) represent a complex situation since guest molecules can interact with different parts of the organic linkers and the metal clusters and may alter the details of the pore structure and system properties. A prominent example is the so-called retrofitted MOF material TCNQ@Cu₃(BTC)₂ that has attracted a lot of attention due to its electronic properties induced by the host–guest interactions. Only recently, structural evidence has been presented for a bridging binding mode of TCNQ to two Cu paddlewheel units; however, many issues regarding the redox chemistry of Cu₃(BTC)₂ and TCNQ are currently unsolved. Herein, we report a powerful spectroscopic approach to study the host–guest chemistry of this material. Combining IR spectroscopy in the presence of CO and EPR spectroscopy, we found that the intrinsic Cu(I) defects of the host react with the guest, forming TCNQ radical anions. This chemistry has profound implications, in particular, with respect to the performance of TCNQ@Cu₃(BTC)₂ as an electronic conductor. A decreasing availability of open Cu(II) sites with increasing TCNQ loading proves the coordinative binding of TCNQ to the paddlewheel nodes, and a heterogeneous structure is formed with different TCNQ arrangements and pore environments at low TCNQ loadings. Finally, the combined use of spectroscopic characterization techniques has proven to be, in general, a powerful approach for studying the complex chemistry of host–guest materials.

KEYWORDS: metal–organic frameworks, redox-active guests, defect chemistry, CO probe IR spectroscopy, EPR spectroscopy

INTRODUCTION

Porous materials play an important role in fundamental research and large-scale industry applications. From activated carbon in exhaust hoods to zeolites for catalytic cracking of hydrocarbons, all materials share high surface areas and pore volumes that allow guest molecules to diffuse and adsorb.¹ A newer class of porous materials named metal–organic frameworks (MOFs) has attracted tremendous attention for use in applications like gas storage, separation, and catalysis.²,³ Particularly, MOFs with coordinately unsaturated metal sites, the so-called open metal sites (OMSs), bear great potential compared to their closed coordination-shell analogues as they typically feature high heats of adsorption and intrinsic reactive centers for catalysis.⁴,⁵ Moreover, the OMSs can serve as an anchoring point for other species that are introduced into the MOF postsynthetically, for example, for medical applications.⁷ Recently, retrofitting was introduced as a concept, in which metal nodes with OMSs or labile monotopic ligands are bridged by additional linkers to fine-tune physicochemical properties.⁸ In all cases, the OMS of the MOF is involved in an interaction with the guest molecule, whose nature is typically difficult to assess. Crystallographic characterization, for example, requires a uniform distribution of the guest molecule with long-range order throughout the crystal lattice. In many cases, however, the guest occupies only a portion of the available coordination sites and thus characterizes rather as
a defect to the parent structure that cannot be determined by standard single-crystal X-ray diffraction experiments. Even at high occupancies, structure determination can be challenging due to the small sizes of the crystallites. In some cases, it is possible to track the presence of guests by powder X-ray diffraction (PXRD). For example, Cu$_2$(BTC)$_2$ (HKUST-1, BTC = 1,3,5-benzenetricarboxylate), which is constructed from dimeric Cu(II) paddlewheel units and tritopic BTC linkers, exposes OMSs at the Cu centers after removal of BTC = 1,3,5-benzenetricarboxylate), which is constructed from dimeric Cu(II) paddlewheel units and tritopic BTC linkers, exposes OMSs at the Cu centers after removal of solvents through thermal treatment. As the Cu centers are linkers, exposes OMSs at the Cu centers after removal of BTC = 1,3,5-benzenetricarboxylate), which is constructed from dimeric Cu(II) paddlewheel units and tritopic BTC linkers, exposes OMSs at the Cu centers after removal of solvents through thermal treatment. As the Cu centers are linkers, exposes OMSs at the Cu centers after removal of solvents through thermal treatment.

Figure 1. Pore chemistry in TCNQ-loaded Cu$_3$(BTC)$_2$. (a) In the idealized picture, all Cu paddlewheels are bridged by TCNQ to form one-dimensional arrays along the (111) crystal lattice. (b) For a realistic description of the arrangement of TCNQ, as well as noncoordinated TCNQ (left) and TCNQ in interaction with intrinsic framework, defects such as missing linker defects (right) must be considered. Color code: C, gray; O, red; N, blue; Cu, dark blue or green. In particular, Cu(II)—Cu(II) and Cu(II)—Cu(I) paddlewheels are depicted in blue and green, respectively. H atoms are omitted for clarity.

EXPERIMENTAL SECTION

Synthesis. Pristine Cu$_3$(BTC)$_2$ was synthesized following the procedure reported in the literature. TCNQ-loaded Cu$_3$(BTC)$_2$ samples were synthesized following our reported vapor-phase infiltration protocol. Briefly, after thermal activation to remove the solvent (a mixture of N,N'-dimethylformamide, ethanol, and H$_2$O) used for solvothermal synthesis and creating the characteristic OMS at the Cu paddlewheel nodes, Cu$_3$(BTC)$_2$ was physically mixed in a mortar with TCNQ, and then, the mixture was annealed under vacuum in a sealed glass ampoule and heated for 3 days at 180 °C to...
yield the host–guest complex. Distinct amounts of TCNQ were employed to prepare a set of samples with different molar fractions of
TCNQ hereafter referred to as xTCNQ@Cu3(BTC)2 with x = n(TCNQ)/n(Cu3(BTC)2) = 0.25, 0.5, and 1.0. During and after the synthesis, the materials were handled under inert conditions (dry Ar or vacuum) and stored in the glovebox to avoid contamination from the atmosphere.

For comparison, a sample was synthesized via solution impregnation following the published synthesis protocol of Talin et al. Therefore, the parent Cu3(BTC)2 was immersed in a dichloromethane (DCM) solution of TCNQ for 3 days (see the Supporting Information for details). The TCNQ–loaded sample, named soTCNQ@Cu3(BTC)2, was collected by filtration and dried under vacuum at room temperature.

In Situ IR Spectroscopy. IR spectra were collected in transmission mode on a Bruker Vertex 70 Fourier transform spectrophotometer, equipped with a KBr beamsplitter and a MCT cryodetector, accumulating 32 scans at a 2 cm$^{-1}$ resolution. All samples were examined in the form of thin self-supporting pellets that were prepared inside the glovebox to avoid exposure to the atmosphere. The pellets, mechanically protected with a gold envelope, were placed in a special homemade quartz cell used for operations down to liquid N2 temperature (i.e., a nominal temperature of 77 K). Transferred out of the glovebox, the cell was connected to a conventional high-vacuum glass line, equipped with a mechanical and a turbo molecular pump (capable of a residual pressure of P < 10$^{-3}$ mbar), which allows activation and in situ adsorption/desorption measurements of the employed molecular probe in a fully controlled atmosphere. Before CO adsorption, each sample was heated at 150 °C under a dynamic vacuum (10$^{-5}$ mbar) for 2 h to remove any potential residual fraction of undesired adsorbates and/or moisture. After the thermal treatment, the material was exposed to CO and cooled down to 77 K. IR spectra were recorded at this temperature, while the CO pressure was gradually decreased to 10$^{-4}$ mbar to obtain an isothermal set of spectra.

EPR Spectroscopy. All cw EPR experiments were performed with a Bruker EMX micro (X-band, 9.4 GHz) spectrometer fitted with a Bruker ER 4119HS cylindrical cavity. Low-temperature experiments were enabled by the Oxford Instruments He cryostat ESR 900. Modulation amplitudes were always set smaller than the smallest peak-to-peak linewidth of transitions in the experimental EPR spectra. Measurements at different microwave (mw) powers ensured that no line shape distortions through saturation occurred for the signals of interest, and that, in case of quantitative experiments, EPR spectra were recorded in the non-saturating mw power regime. The MATLAB toolbox EasySpin v. 5.2.25,6 employing the exact diagonalization of the spin Hamiltonian matrix, was used for spectral simulations of the EPR signals. EPR signal intensities of various EPR active species were determined by the double integration of their simulated EPR signals, if their quantification was desired.

All TCNQ–loaded samples for EPR measurements were prepared inside the glovebox by filling the material into an X-band EPR quartz glass tube connected with a high vacuum PTFE valve. The valve was closed and transferred to a vacuum station, where the EPR tube was evacuated and then flame-sealed, keeping the material under inert conditions during each step.

Results and Discussion

After the synthesis, all materials were carefully characterized to quantitatively determine the TCNQ loading in the MOF structure. Particularly, PXRD data confirmed the successful infiltration (see Figure S2). Standard characterization results of xTCNQ@Cu3(BTC)2, with x = 0.1, 1.0 in 0.1 increments were thoroughly reported in a previous publication. In short, Cu3(BTC)2 exhibits one small pore of ~11 Å and two types of large pores of ~16 Å in diameter. The Cu paddlewheel nodes are oriented in a way that the OMSs at the axial position point to the center of one large pore while being tangential relative to the other large pore (see the Supporting Information for more information). Notably, the synthesis conditions involving noninert solvents at elevated temperatures and a thermal activation of the material promote the formation of intrinsic Cu(I) sites, the so-called modified node defects.30 During the vapor-phase infiltration reaction, TCNQ diffuses into the large pores of Cu3(BTC)2 and binds preferentially to the available OMSs of two neighboring Cu paddlewheels, as evidenced by PXRD.31 At the highest possible loading (x = 1.0), the large pores can accommodate two molecules of TCNQ while still exhibiting a residual BET-specific surface area of 574 m$^2$ g$^{-1}$, that is, about one-third of the pristine Cu3(BTC)2, still allowing for the adsorption of smaller molecules such as N2 or CO. During the reaction, a small amount of Cu(TCNQ) forms as a byproduct on the external surface of the crystallites, suggesting an electron transfer from the intrinsic Cu(I) defects of the MOF to TCNQ. The electrical conductivity of the sample correlates quasi-exponentially with the amount of TCNQ employed during the reaction and amounts to 1.5 × 10$^{-4}$ S cm$^{-1}$ for 1.0TCNQ@Cu3(BTC)2.32

In Situ IR Study of CO Adsorption at 77 K. Cu3(BTC)2 by itself represents a complex system involving OMSs and a sophisticated defect chemistry. The introduction of TCNQ into the pores adds an additional layer of complexity as it can coordinate to the OMS and accept electrons from the framework. Therefore, the use of standard characterization techniques does not reveal the full picture of this interaction. In situ IR spectroscopy with probe molecules was used in the past and proved to be a powerful technique to shed light on complicated systems such as MOFs containing open metal sites. In particular, the study of CO adsorption at 77 K is one of the more common tools employed to probe the Lewis acidic sites generated by the presence of exposed metal cations.33,34 For instance, the positively charged Cu(II) species in Cu3(BTC)2 allow the formation of carbon end Cu(II)–CO adducts, while, in parallel, CO is able to interact with framework Cu(II) defect sites generated by missing carbonate units (decarboxylation during thermal activation) or with extra-framework Cu(I) species deriving from Cu3O impurities (e.g., formed during solvothermal synthesis).20,24,38 Finally, the framework Cu(II) cations in Cu3(BTC)2 are linked to partially negatively charged carbonate units, which could constitute a possible interaction site for CO through the formation of oxygen-end adducts as well (organic linker–OC).39,40 By exploiting the subtle capacity of CO to disclose the existence of sites of such different nature, it is clear how this probe molecule could be useful to clarify the host–guest structure and interactions of xTCNQ@Cu3(BTC)2. As a general comment, it is worth highlighting that only the species available for interaction with CO are effectively probed, whereas fully coordinated and nonaccessible species cannot be detected. Therefore, this method is not able to probe the Cu(TCNQ) byphase. In addition, it is important that this technique probes the bulk volume of the porous material, and contributions from the external crystallite surface have negligible contributions to the observed IR signals due to the low external surface-to-volume ratio.

Cu3(BTC)2. Before analyzing the host–guest materials, CO adsorption was carried out on the parent MOF to get a clear picture of the different Cu families. For this purpose, a thin pellet of Cu3(BTC)2 was prepared inside the glovebox and placed in a homemade IR cell for low-temperature measure-

DOI: 10.1021/acsami.9b16663
ACS Appl. Mater. Interfaces 2020, 12, 1024–1035
ments, as reported in the Experimental Section. After a thermal activation at 150 °C for 2 h, the material was exposed to ~40 mbar CO, cooled down to 77 K, and then gradually evacuated at the same temperature. The isothermal set of spectra is reported in Figure 2 (from dark blue to dark red) in the CO vibrational mode spectral range. The spectra of the materials, showing the full mid-IR spectral region before CO contact, are provided in Figure S3.

The whole set of spectra is characterized by an extremely intense and complex envelope of bands due to the CO interaction with the different surface sites. A detailed explanation of the various components and of their spectral behavior is reported in the Infrared Spectroscopy section of the Supporting Information. However, to allow a full comprehension of the IR spectra collected after CO adsorption on the xTCNQ@Cu3(BTC)2 host–guest systems, the main spectral features of CO in interaction with pristine Cu3(BTC)2 are summarized in the following. (i) The out-of-scale band located at around 2172 cm−1 is ascribed to the formation of reversible Cu(II)–CO adducts. The peculiar behavior of this band during the outgassing process is due to the gradual conversion of polycarbonyl adsorbate species into monocarbonyls.24,38 (ii) Considering the interpretation of Drenchev and co-workers, the band at 2192 cm−1 could be generated by a different Cu(II)–CO species, whereas the signal at 2148 cm−1 (probably stabilized by the presence of the 2192 cm−1 component) could be tentatively assigned to CO in interaction with the organic part of the framework.38 (iii) The bands at around 2120 and 2096 cm−1 can be ascribed to the formation of monocarbonyl Cu(I)–CO and Cu(I)–OC (in which the adsorbed CO molecule interacts through the oxygen atom) adducts due to the presence of defective Cu(I) sites.38,40,41 The component at 2120 cm−1 is very stable, persisting until very low CO coverages. (iv) The two signals at 2132 and 2106 cm−1 appear in the set of spectra just at a low CO coverage when most of the Cu(II)–CO complexes (band at around 2172 cm−1) is removed and then gradually decrease in intensity upon outgassing. These signals can be tentatively assigned to vibrational modes of bridged Cu(I)–CO–Cu(II) carboxylates in Cu(II)–Cu(I) dimers generated by the presence of missing carboxylate units.41–43 (v) The two sharp and intense signals at 2142 and 2128 cm−1 simultaneously decrease in intensity without any noticeable change in position and are characterized by a very labile nature (they quickly disappear at the early stages of the desorption process). Following the results proposed by Drenchev and co-workers,38 these bands can be ascribed to CO in interaction with the organic linker via the formation of oxygen-end adducts, presumably with the benzene ring. This assignment is confirmed by observing the modification of some specific spectral features of the organic linker at 1912 and 1896 cm−1 during the CO adsorption/desorption experiments with respect to the spectrum of the material before the contact with the probe molecule (see Figure S5).38 (vi) Finally, at a very low CO coverage, a broad signal appears at 2125 cm−1. The frequency of the component is typical of CO adsorbed on Cu(I) sites of amorphous Cu2O impurities.33

The main IR bands resulting from the interaction of CO with pristine Cu3(BTC)2 are summarized in Table 1. xTCNQ@Cu3(BTC)2. After the preliminary analysis of the pristine host framework, we turned our attention to the TCNQ-loaded samples aiming to answer three questions: “What happens to the OMSs upon adsorption of TCNQ?” and “Do we observe redox processes between Cu(I) defects and TCNQ?”, and “How does TCNQ affect the local pore environment of Cu3(BTC)2?” To this purpose, the CO adsorption at 77 K has been carried out on a series of xTCNQ@Cu3(BTC)2 samples (x = 0.25, 0.5, 1.0) on thin pellets prepared inside the glovebox, as reported in the

Table 1. Assignments of the Main IR Bands Formed upon CO Adsorption on Pristine Cu3(BTC)2 (See Figure 2)

<table>
<thead>
<tr>
<th>wavenumber (cm−1)</th>
<th>assignment</th>
<th>CO coverage</th>
<th>ref</th>
</tr>
</thead>
<tbody>
<tr>
<td>2192−2200</td>
<td>polycarbonyl/monocarbonyl Cu(II)–CO</td>
<td>high−low</td>
<td>38</td>
</tr>
<tr>
<td>2178</td>
<td>monocarbonyl Cu(II)–CO</td>
<td>medium low</td>
<td>24, 38</td>
</tr>
<tr>
<td>2172</td>
<td>polycarbonyl Cu(II)–CO</td>
<td>medium high</td>
<td>24, 38</td>
</tr>
<tr>
<td>2148</td>
<td>organic linker–OC</td>
<td>high−low</td>
<td>38</td>
</tr>
<tr>
<td>2142 and 2128</td>
<td>organic linker–OC</td>
<td>high−medium high</td>
<td>38–40</td>
</tr>
<tr>
<td>2123 and 2106</td>
<td>bridged Cu(I)–CO–Cu(II)</td>
<td>low</td>
<td>41–43</td>
</tr>
<tr>
<td>2120</td>
<td>monocarbonyl Cu(I)–CO</td>
<td>high−medium low</td>
<td>38, 40, 41</td>
</tr>
<tr>
<td>2096</td>
<td>monocarbonyl Cu(I)–OC</td>
<td>high−medium low</td>
<td>40, 41</td>
</tr>
<tr>
<td>2125</td>
<td>Cu(I)–CO (of Cu2O impurities)</td>
<td>low</td>
<td>23</td>
</tr>
</tbody>
</table>

“Wavenumber (in cm−1) from this work. "CO coverage at which the band is visible. "Cu(II)–CO adducts different from the standard ones at 2172–2178 cm−1. "Band stabilized by the 2192 cm−1 component.”
Experimental Section. The isothermal sets of spectra of the three materials with various TCNQ loadings are reported in Figure 3, showing the CO vibrational mode spectral range. The spectra have been collected by gradually decreasing the CO pressure of CO was gradually decreased from 16 (dark blue) to 10 mbar (dark red). The spectra before contact with CO are shown in Figure 3 as black curves. For a better comparison, the spectra of the three materials should be normalized to take into account the different pellet thickness. Three pellets have similar thicknesses and densities and provide comparable results. After the gas admission in the IR cell, an envelope of signals related to the CO interaction with the different surface sites appears. Considering the assignments made for the host framework, the band at around 2170 cm$^{-1}$ is generated by Cu(II)–CO adducts (polycarbonyls or monocarbonyls, depending on the CO coverage). This band decreases considerably as a function of TCNQ loading, indicating a reduced number of available Cu(II) OMSs able to interact with the probe molecule. This trend clearly supports earlier experimental and computational results, suggesting the coordination of TCNQ to the Cu(II) OMSs of Cu$_3$(BTC)$_2$. Therefore, it can be reasonably assumed that the three pellets have similar thicknesses and densities and provide comparable results. After the gas admission in the IR cell, an envelope of signals related to the CO interaction with the different surface sites appears. Considering the assignments made for the host framework, the band at around 2170 cm$^{-1}$ is generated by Cu(II)–CO adducts (polycarbonyls or monocarbonyls, depending on the CO coverage). This band decreases considerably as a function of TCNQ loading, indicating a reduced number of available Cu(II) OMSs able to interact with the probe molecule. This trend clearly supports earlier experimental and computational results, suggesting the coordination of TCNQ to the Cu(II) OMSs of Cu$_3$(BTC)$_2$.

Concerning the formation of Cu(I)–CO and Cu(I)–OC adducts (with intrinsic Cu(I) defect sites), the sharp and well-structured bands at 2120 or 2096 cm$^{-1}$, which were clearly evident in pristine Cu$_3$(BTC)$_2$, are totally absent, at any CO coverage, in all three samples. The absence of these spectral components is particularly evident in Figure 4b, which compares the spectra of all the materials at a low CO pressure (1 mbar), which is probably the best situation to observe these differences due to the lower reversibility of the signals generated by Cu(I) species. It should be noted that the Cu(I) defect signals are fully absent in the sample with the lowest TCNQ loading ($x = 0.25$, compare light green and blue curves in Figure 4), suggesting a chemical reaction of these species already induced by the presence of very small amounts of TCNQ. In contrast, in all TCNQ-loaded samples, the extra-framework Cu$_2$O impurities are always present, as evidenced by the broad and persistent (resistant against prolonged outgassing) component at 2125 cm$^{-1}$, which seems completely unaffected by the presence of TCNQ. In our previous work, we observed the formation of a byphase of Cu(TCNQ), which requires transfer of an electron to TCNQ and tentatively hypothesized Cu(I) defects being the reducing agent. This previous hypothesis is now strongly supported by the spectroscopic investigation, revealing the absence of the Cu(I) species and the presence of TCNQ radical anions (see the EPR results in the following section; Figure 5).

Looking at the envelope of bands in the region between 2145 and 2125 cm$^{-1}$, assigned to CO in interaction with the organic backbone (presumably with the benzene ring) via the formation of oxygen-end adducts, the vibrational modes are clearly perturbed by the presence of the large guest molecule inside the MOF cavities. Indeed, at high TCNQ loadings ($x = 1.0$), the sharp components at 2142 and 2128 cm$^{-1}$ in pristine Cu$_3$(BTC)$_2$ shift to 2138 and 2131 cm$^{-1}$, respectively. In contrast, for the materials with lower TCNQ loadings ($x = 0.25$ and 0.5), these bands are distinctly broader, exhibiting an intermediate situation due to the unambiguous presence of two distinct components for each signal (upon desorption up to four bands are visible in this region). These modifications become evident in Figure 4a, where the spectra of the materials at different TCNQ loadings (from 0 to 1.0), at a medium-high CO coverage, are compared in the spectral range of CO stretching vibrations. The above described spectral changes disclose a different pore environment throughout the TCNQ loading series. While all the bands are absent in pristine Cu$_3$(BTC)$_2$ (bands at 2142 and 2128 cm$^{-1}$), in the sample...
with $x = 1.0$ (bands at 2138 and 2121 cm$^{-1}$), all large pores are occupied with a maximum of two TCNQ molecules. In these two well-defined “extreme” cases, the local pore environment is therefore homogeneous. In between these two extremes, samples with $x = 0.25$ and 0.5 exhibit empty, partially, or completely filled cavities (i.e., containing 0, 1, or 2 TCNQ molecules), resulting in a more complex spectroscopic situation compared to samples with $x = 0$ and 1.0.

The unequivocal perturbation induced in the organic backbone by the guest TCNQ molecule has been also proven by directly observing the bands at 1912 and 1896 cm$^{-1}$ (assigned to combination modes of the organic framework) in the spectra of the TCNQ loading series before the CO contact, as reported in Figure S5. These vibrations have the maximum intensity in the pristine Cu$_3$(BTC)$_2$, whereas they gradually disappear upon increasing the TCNQ amount.

Finally, we have to spend a few words to evaluate the spectral behavior of the two IR bands at around 2192 and 2148 cm$^{-1}$, whose assignment has been reported to be quite controversial in the literature. The 2192 cm$^{-1}$ component is always present in all TCNQ-loaded samples, whereas the 2148 cm$^{-1}$ signal gradually disappears by increasing the guest molecule loading (see Figure 4c). This behavior demonstrates the total absence of correlation between these two spectral features, confirming the assignments proposed by Drenchev and co-workers. The band at 2192 cm$^{-1}$ is ascribed to the CO adsorption on a minor fraction of “different” Cu(II) sites, not involved in the interaction with the TCNQ guest molecule, as testified by the high intensity of this band compared to the signal due to standard Cu(II) species (2172 cm$^{-1}$), especially in the material with the highest TCNQ loading ($x = 1.0$). In contrast, the component at 2148 cm$^{-1}$ assigned to CO attached to the organic part of the framework is no longer present in 1.0TCNQ@Cu$_3$(BTC)$_2$ due to the perturbation induced by the TCNQ molecule at its maximum loading.
Table 2. Assignments of the Main IR Bands Formed upon CO Adsorption on xTCNQ@Cu$_3$(BTC)$_2$ (See Figure 4)

<table>
<thead>
<tr>
<th>wavenumber$^a$</th>
<th>assignment</th>
<th>xTCNQ@Cu$_3$(BTC)$_2$$^b$</th>
</tr>
</thead>
<tbody>
<tr>
<td>2192$^c$</td>
<td>polycarboxyl/monocarboxyl Cu(II)–CO</td>
<td>0.25, 0.5, 1.0</td>
</tr>
<tr>
<td>2178</td>
<td>monocarboxyl Cu(II)–CO</td>
<td>0.25 (strong), 0.5 (medium), 1.0 (weak)</td>
</tr>
<tr>
<td>2172</td>
<td>polycarboxyl Cu(II)–CO</td>
<td>0.25 (strong), 0.5 (medium), 1.0 (weak)</td>
</tr>
<tr>
<td>2148</td>
<td>organic linker–OC</td>
<td>0.25, 0.5</td>
</tr>
<tr>
<td>2142 and 2128</td>
<td>organic linker–OC</td>
<td>0.25 (strong), 0.5 (medium)</td>
</tr>
<tr>
<td>2138 and 2131</td>
<td>organic linker–OC</td>
<td>0.25 (weak), 0.5 (medium), 0.1 (strong)</td>
</tr>
<tr>
<td>2132 and 2106</td>
<td>bridged Cu(I)–CO–Cu(II)</td>
<td>absent</td>
</tr>
<tr>
<td>2120</td>
<td>monocarboxyl Cu(I)–CO</td>
<td>absent</td>
</tr>
<tr>
<td>2096</td>
<td>monocarboxyl Cu(I)–OC</td>
<td>absent</td>
</tr>
<tr>
<td>2125</td>
<td>Cu(I)–CO (of Cu$_3$O impurities)</td>
<td>0.25, 0.5, 1.0</td>
</tr>
</tbody>
</table>

$^a$Wavenumber (in cm$^{-1}$) from this work. $^b$Presence of the band in the xTCNQ@Cu$_3$(BTC)$_2$ samples and corresponding intensity. $^c$Cu(II)–CO addsucts different from the standard ones at 2172–2178 cm$^{-1}$.

EPR Spectroscopy. In situ IR spectroscopy results clearly evidence the existence of Cu(I) OMSs in pristine Cu$_3$(BTC)$_2$ that vanish upon TCNQ loading, indicating a possible electron transfer from Cu(I) ions to TCNQ molecules. To distinctly elucidate the formation of paramagnetic TCNQ$^-$ radicals, we decided to perform cw EPR experiments on some xTCNQ@Cu$_3$(BTC)$_2$ samples, which were absent in pristine Cu$_3$(BTC)$_2$ (see Figure 5), contrasting with the results by Talin et al. who did not observe any signal of TCNQ$^-$ radicals when they loaded TCNQ into Cu$_3$(BTC)$_2$, via solution impregnation (TCNQ loading, x ∼ 0.25).44 To scrutinize the differences between the two infiltration methods, we synthesized a reference sample, soTCNQ@Cu$_3$(BTC)$_2$, following its published synthesis protocol, which was then compared with our xTCNQ@Cu$_3$(BTC)$_2$ samples obtained by vapor-phase infiltration.

The room temperature spectra of pristine Cu$_3$(BTC)$_2$, xTCNQ@Cu$_3$(BTC)$_2$ (x = 0.25, 0.5), and soTCNQ@Cu$_3$(BTC)$_2$ exhibit an isotropic signal at g = 2.15 with a Lorentzian peak-to-peak linewidth of ΔB$_{pp}$ = 92(3) mT (Figure S6). According to previous works,25,56 we assign these signals to the electron spin S = 1 state of the antiferromagnetically coupled Cu(II)–Cu(II) paddlewheel units, subjected to interdimeric exchange processes. The disappearance of this signal at T = 14 K in all the materials confirms its antiferromagnetic nature (Figure S7). Notably, all TCNQ-loaded samples show an isotropic signal at g = 2.003(2) at T = 14 K (around 335 mT in Figure 5a) with an experimental linewidth of ΔB$_{pp}$ ≈ 1.6 mT, typical for radical species due to the presence of TCNQ$^-$44. According to the literature, the expected $^{14}$N hyperfine (hf) splitting of TCNQ$^-$ anions is of the order of 3 MHz (~0.1 mT)40,47, which is too small to be resolved in the present case. In contrast to the xTCNQ@Cu$_3$(BTC)$_2$ samples, the TCNQ$^-$ signal of soTCNQ@Cu$_3$(BTC)$_2$ exhibits shoulders at the low and high field end, which we tentatively assign to TCNQ$^-$ anions with a larger EPR linewidth. This larger linewidth might indicate a high local concentration of the contributing fraction of TCNQ$^-$ molecules, which leads to dipolar interactions between these species, resulting in a line broadening.

TCNQ Radical Anions. Since we did not observe any signals of TCNQ$^-$ radicals at room temperature (Figure S6), we performed additional EPR experiments at T = 95 K on all TCNQ-loaded samples. Like in the spectra obtained at T = 14 K, the almost isotropic signals of the TCNQ$^-$ radicals at g-values of g = 2.003(2) are visible (Figure Sb). However, their peak-to-peak linewidths increased from ΔB$_{pp}$ = 1.56(23) mT at T = 14 K to ΔB$_{pp}$ = 2.52(40) mT at T = 95 K. By increasing the temperature, the TCNQ$^-$ signals further broaden (e.g., Figure S8) until they totally vanish at room temperature (Figure S6). This observation can be explained by spin–spin interactions between the TCNQ$^-$ radicals with the S = 1 state of nearby Cu(II)–Cu(II) paddlewheel units, which becomes increasingly populated at higher temperature.45 Indeed, pristine Cu$_3$(BTC)$_2$ as well as the two TCNQ-loaded xTCNQ@Cu$_3$(BTC)$_2$ (x = 0.25, 0.5) samples and the reference sample soTCNQ@Cu$_3$(BTC)$_2$ shows the typical S = 1 signal of magnetically diluted Cu(II)–Cu(II) paddlewheel units at T = 95 K (see Figures S9 to S12),25,44 verifying that the excited S = 1 state of the paddlewheels had become significantly thermally populated at T = 95 K. This result strongly suggests that the TCNQ$^-$ radicals are located inside the pores of the Cu$_3$( BTC)$_2$ framework, and the signal is not produced by the Cu(TCNQ) byproduct that was observed to form at the outer surface of the xTCNQ@Cu$_3$(BTC)$_2$ crystallites.13

To roughly estimate the amount of TCNQ$^-$ radicals per paddlewheel unit, the formula for the magnetic susceptibility $\chi_M$ of monomeric electron spin S = 1/2 species and $\chi_{PW}$ of dimeric electron spin S = 1 species of the paddlewheels were used, as given in the Supporting Information.25 Here, the room temperature S = 1 signals of the copper paddlewheels and of the TCNQ$^-$ radicals at T = 75 K were compared with experimental spectra of an ultramarine reference sample that was measured under the same experimental conditions. This accounts for possible differences in the EPR intensities due to different coupling conditions at different temperatures. For each sample, the amount ε of TCNQ$^-$ radicals per paddlewheel unit was calculated via

$$\varepsilon = \frac{I_M(75 \text{ K})}{I_{PW}(295 \text{ K})} \frac{\chi_{PW}(295 \text{ K})}{\chi_M(75 \text{ K})}$$

(1)

where $I_M(75 \text{ K})$ and $I_{PW}(295 \text{ K})$ are the EPR intensities of the TCNQ$^-$ species at T = 75 K and of the S = 1 paddlewheel species at T = 295 K, respectively, as they have been determined by double integrating the corresponding simulated
EPR signals, normalized by means of the ultramarine reference signals. In accordance with the literature, an exchange coupling constant of $J = -370 \text{ cm}^{-1}$ was used in the Bleaney–Bowers equation for the magnetic susceptibility of the $S = 1$ spin, which had been determined for the antiferromagnetically coupled Cu paddlewheel units of Cu$_3$(BTC)$_2$·(H$_2$O)$_3$. Following this procedure, we estimate a concentration of 0.025(13), 0.016(8), and 0.010(5) TCNQ$^-$ radicals per paddlewheel units for the samples 0.25TCNQ@Cu$_3$(BTC)$_2$, 0.5TCNQ@Cu$_3$(BTC)$_2$, and solTCNQ@Cu$_3$(BTC)$_2$, respectively (Table 3). These quantitative results prove that a doubling of the TCNQ molecules per paddlewheel unit does not increase the concentration of TCNQ$^-$ radicals, suggesting a scenario where all accessible framework Cu(I) (intrinsic) defects are already oxidized at low TCNQ loadings ($x = 0.25$), and additional TCNQ is adsorbed without undergoing a redox reaction. Consequently, the concentration of TCNQ$^-$ radicals remains at the same order of magnitude in all TCNQ-loaded samples. In contrast to the results reported by Talin et al., we observe a significant amount of TCNQ$^-$ anions in the sample prepared via solution impregnation. This discrepancy could be derived by the slightly different reaction conditions, and more detailed and elaborate experiments to study the materials obtained by TCNQ solution impregnation are suggested. In this study, we show that Cu(I) defects in the crystalline solid act as the reducing agent for the TCNQ guest molecules in both samples obtained via solution impregnation or vapor-phase infiltration. Thereby, the presence of TCNQ radical anions is directly dependent on the intrinsic Cu(I) defect concentration in the pristine MOF, which, in turn, is sensible to the synthesis and activation procedure. Thus, besides the choice of the solvent, which can promote the reduction of TCNQ$^-$, the defect concentration of the parent MOF adds another layer of complexity to the synthesis and properties of TCNQ@Cu$_3$(BTC)$_2$.

Cu(II) Species. At low temperatures, pristine Cu$_3$(BTC)$_2$ as well as the samples xTCNQ@Cu$_3$(BTC)$_2$ and solTCNQ@Cu$_3$(BTC)$_2$ exhibits signals of monomeric Cu(II) species, which have a 3d$^9$ electron configuration and, therefore, an electron spin of $S = 1/2$ (Figure 6 and Figures S13 and S14). The four measured samples display broad Cu(II) background signals with no resolved $^{63,65}$Cu (nuclear spin $I = 3/2$) hf splitting, indicative for monomeric Cu(II) species in high local concentration or not well-defined coordination as they might occur at grain or outer boundaries of the MOF crystals or in extra-framework impurity species. One might expect such impurity signals since extra-framework hexaaqua Cu(II) complexes were identified in samples of Cu$_3$(BTC)$_2$ in a previous work by Pöppl et al. The present samples were activated at higher temperatures and washed for a longer time than in the work of Pöppl et al. However, extra-framework Cu(II) species might still have remained in the present samples, even if they do not occur as hexaaqua complexes due to the potential removal of all water under the harsher conditions.

Table 3. EPR-Derived Concentration c (Number per Paddlewheel Unit) of Different Paramagnetic Species Determined via Eq 1 and Tentative Assignments

<table>
<thead>
<tr>
<th>Sample</th>
<th>TCNQ$^-$</th>
<th>species A - Cu(II)–Cu(I) paddlewheel</th>
<th>species B - defective Cu(II) monomer</th>
<th>species C - defective Cu(II) monomer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cu$_3$(BTC)$_2$</td>
<td>$&lt;0.0002$</td>
<td>0.015(8)</td>
<td>0.0013(6)</td>
<td>$&lt;0.01$</td>
</tr>
<tr>
<td>0.25TCNQ@Cu$_3$(BTC)$_2$</td>
<td>0.025(13)</td>
<td>$&lt;0.0004$</td>
<td>0.0025(13)</td>
<td>0.0050(13)</td>
</tr>
<tr>
<td>0.5TCNQ@Cu$_3$(BTC)$_2$</td>
<td>0.016(8)</td>
<td>$&lt;0.0012$</td>
<td>0.0012(6)</td>
<td>0.012(6)</td>
</tr>
<tr>
<td>solTCNQ@Cu$_3$(BTC)$_2$</td>
<td>0.010(5)</td>
<td>$&lt;0.0025$</td>
<td>$&lt;0.0003$</td>
<td>$&lt;0.006$</td>
</tr>
</tbody>
</table>

*Possibly assigned to a broken paddlewheel. Same as species B$_a$ but in higher local concentration.

![Figure 6](https://example.com/f6.png)

Experimental EPR spectra of (a) pristine Cu$_3$(BTC)$_2$ (blue) and (b) sample 0.5TCNQ@Cu$_3$(BTC)$_2$ (green) measured at $T = 14$ K. In panel (a), the sum simulation (brown) is the superposition of the simulated signals of the monomeric Cu(II) species A (dark red) and B$_a$ (red). In panel (b), the sum simulation (brown) is a superposition of the signals of the Cu(II) species B$_a$ (red) and B$_b$ (light red). The asterisk labels the saturated signal of TCNQ$^-$ radicals that extends outside the frame and that overlaps with the Cu(II) signals. See Table S1 for simulation parameters.
activation conditions. In addition, the EPR spectrum of pristine Cu
\(\text{(BTC)}_2\) at \(T = 14\) K resolves the \(g_{63,65}\)Cu hf transitions of two Cu(II) species \(A\) and \(B\), between 260 and 320 m\(T\) as well as the \(g_{63,65}\)Cu hf transitions of species \(A\) and \(B\), both 320 and 340 m\(T\) (Figure 6a; see Table S1 for the spin Hamiltonian parameters). Further, minor \(g_{63,65}\)Cu hf transitions of other defective Cu(II) species are resolved for pristine \(\text{Cu}_3(\text{BTC})_2\) at \(T = 14\) K, but their low signal intensities and their overlapping with other hf transitions of different Cu(II) species prevented their clear assignment and determination of their spin Hamiltonian parameters. However, their presence indicates a varied presence of different kinds of well-defined Cu(II) defects in pristine \(\text{Cu}_3(\text{BTC})_2\).

As explained above for the TCNQ\(^{-}\) radicals, we have determined the concentration of species \(A\) in pristine Cu
\(\text{(BTC)}_2\) at \(T = 75\) K to 0.015(8) Cu(II) ions per paddlewheel unit (Table 3). Within the experimental error, this concentration is very similar to those determined for the TCNQ\(^{-}\) radicals in all three TCNQ-loaded samples. Importantly, the signal of species \(A\) was not observed for the TCNQ-loaded samples anymore (Figure 6b and Figures S13 and S14). The above reported observations suggest that species \(A\) represents Cu(II) ions of mixed valence Cu(II)–Cu(I) paddlewheel units in pristine Cu
\(\text{(BTC)}_2\). In accordance to the IR results, all diamagnetic Cu(I) ions are oxidized to Cu(II) upon TCNQ loading, and, in parallel, a corresponding fraction of the guest molecule is reduced to TCNQ\(^{-}\).

Consequently, the same concentration of TCNQ\(^{-}\) radicals is observed in the loaded samples as Cu(II)–Cu(I) paddlewheel units occur in pristine Cu
\(\text{(BTC)}_2\) (Table 3). The magnetic coupling of a potentially formed Cu(II)–Cu(II)–TCNQ\(^{-}\) spin triad is discussed in detail in the Supporting Information. If such spin triads are present in the TCNQ-loaded samples, then the present results indicate a weak exchange interaction between the Cu(II) ion and the axially coordinating TCNQ\(^{-}\) radical without significant distortion of the antiferromagnetic coupling between the two Cu(II) ions and the isotropic TCNQ\(^{-}\) signal. The \(g\)- and \(\delta\)Cu hf-tensor principal values \(g_x = 2.290(3)\) and \(A_x = 0.0170(7)\) cm\(^{-1}\) of species \(A\) are typical for Cu(II) ions in square planar to square pyramidal coordination. For a better understanding of the different Cu(II) species, we suggest pulsed EPR experiments like electron spin echo envelope modulation (ESEEM) and electron nuclear double resonance (ENDOR) experiments, which might resolve the hf coupling between the Cu(II) electron spin \(S = 1/2\) and the Cu nuclear spin \(I = 3/2\) of the adjacent \(\text{Cu}(I)\) ion. Such experiments are currently in progress. Nevertheless, a reduction of TCNQ molecules by Cu(I) ions is strongly indicated by the presented IR and EPR results.

### CONCLUSIONS

An advanced spectroscopic characterization study has been carried out by means of IR spectroscopy of adsorbed probe molecules (here, CO) and EPR spectroscopy to disclose the host–guest chemistry of a series of Cu
\(\text{(BTC)}_2\) samples containing increasing amounts of TCNQ.

The combined spectroscopic approach together with a comparison of solvent-free vapor-phase infiltration and
solution impregnation provides substantial answers to the three questions raised at the beginning of this study. By means of in situ IR spectroscopy, we clearly show that the amount of Cu(II) OMSs in Cu₃(BTC)₂ decreases monotonically by increasing the guest loading because of the occupation by TCNQ. In combination with our previous powder diffraction data, the presented results dispelled any doubt about the coordination of TCNQ to the Cu(II) dimers of the paddlewheel units. Moreover, the IR data revealed a homogeneous pore environment for the pristine and fully loaded MOF (x = 1.0; i.e., two TCNQs per large pore), whereas intermediate samples exhibited a more heterogeneous, variegated TCNQ distribution inside the microporous cavities. In addition, both IR and EPR data evidenced the presence of intrinsic Cu(I) framework defects in the volume of pristine Cu₃(BTC)₂ that disappear immediately upon exposure to even small TCNQ amounts. In return, the presence of TCNQ radical anions becomes evident in all TCNQ-loaded samples. Remarkably, the radical concentration in the TCNQ-loaded samples correlates quantitatively with the Cu(II)–Cu(I) defect concentration in the parent MOF, evidencing an electron transfer from the Cu(I) defects to TCNQ. Additional pulsed EPR measurements are currently in progress to get further insights into the nature of the different Cu(II) species. The results obtained during this study have profound implications for the fabrication and the understanding of the physical origin of the electronic conductivity and performance of devices based on TCNQ@Cu₃(BTC)₂. We find that the framework Cu(I) defect concentration directly correlates with the TCNQ radical anion concentration, thereby introducing additional charge carriers into the system, which is typically associated with an increase in electrical conductivity. In fact, the maximum TCNQ⁺ concentration is already reached at low loadings, which would result in a plateau of the conductivity at higher loadings if the introduced charge carrier was the only contribution to the conductivity. This contrasts with the experimental results. An increased charge mobility due to neutral TCNQ in the pores, or contributions from a Cu(TCNQ) surface biphase that forms under certain conditions, must also be taken into account when studying the electronic properties of these materials. The use of carefully fabricated defect-free Cu₃(BTC)₂ thin films as the starting material for the infiltration reaction under very mild conditions (low temperature, inert solvent) will likely lead to a host–guest material without the formation of TCNQ radicals. In a similar way, defect-free bulk materials should be accessible by optimization of the synthetic protocol; however, mild activation procedures to generate the OMSs without Cu(I) defect formation are scarce or even absent. The present system is yet another example where the defect chemistry of MOFs has a significant impact on the material properties and underlines the importance of advancing the research field of defect-engineered MOFs.

At this point, we like to highlight our recent computational study on structurally retrofitting metal–organic frameworks with cross-linkers (CL), in particular, Cu paddlewheel-based systems such as CL@Cu₃(BTC)₂, and the expansion to CL@NOTT-100 and CL@NOT-101. A library of 20 CLs with binding properties similar to TCNQ was evaluated. We suggest that the chemical interaction of these potentially noninnocent guest molecules with Cu paddlewheel-based MOFs beyond the simple coordination to the open copper sites may be as complex as the present case of Cu₃(BTC)₂ (HKUST-1), and it may be elucidated in detail following the methodology as we described above.

As a final, and more general, remark, it is worth highlighting that a high precision inorganic solid-state synthesis together with the application of a suite of complementary techniques of spectroscopic characterization represents a powerful approach to study the subtle, and frequently unexpectedly complex, chemistry of synergetic Guest@MOF systems, in general.

■ ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsami.9b16663.

Information about the synthesis, powder X-ray diffraction, IR, and EPR spectroscopy (PDF)

■ AUTHOR INFORMATION

Corresponding Authors
*E-mail: valentina.crocella@unito.it (V.C.).
*E-mail: roland.fischer@tum.de (R.A.F.).

ORCID
Matthias Mendt: 0000-0002-1295-0228
Valentina Crocella: 0000-0002-3606-8424
Roland A. Fischer: 0000-0002-7532-5286

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

Funding
DFG SPP1928 (COORNETs).

Notes
The authors declare no competing financial interest.

■ ACKNOWLEDGMENTS

The authors would like to thank Silvia Bordiga for insightful discussions. C.S. acknowledges his scholarships from the German Academic Scholarship Foundation and from the German Chemical Industry Fund (FCI). This work was supported by the German Research Foundation Priority Program 1928 “CCORNETs” (www.coornets.tum.de).

■ REFERENCES

High Electrical Conductivity and High Porosity in a Guest@MOF

Pujari, S. P.; Zuilhof, H.; Janek, J.; Allendorf, M. D.; Fischer, R. A.

ACS Applied Materials & Interfaces

Presence of Water.

Welle, A.; Mugnaini, V.; Meded, V.; Zharnikov, M.; Wo

2018

Frameworks.

Gilles, M. K. Cu1+ in HKUST-1: Selective Gas Adsorption in the

39400

Surfaces of Heterogeneous Catalysts by in Situ IR Spectroscopy.

Morphology and Electrical Properties of Cu3BTC2 Thin Films Before

and After Reaction with TCNQ.

Films.

Compound Cu3(BTC)2.

Lamberti, C.; Zecchina, A.; Solari, P. L.; Kongshaug, K. O.; Bordiga, S.

ESR Spectroscopy of Cupric Ions in the Metal

Organic Framework Cu3(BTC)2.

Retrofitting Adapts a Metal

Surface of HKUST-1 Organic Framework to

Organic Framework

Organic Framework to

Organic Framework

Dual-Cation Sites.

Amino-Modified SBA-15 Catalysts.

Computational and Fourier

Transform Infrared Spectroscopic Studies on Carbon Monoxide

Adsorption at Cupric Ions in the Metal

Organic Framework Compound Cu3(BTC)2.

Adsorption of Gases to Cupric Ions in the Zn(II)-Doped Porous

Coordination Polymer: Spectroscopic Features and Interaction

with High Seebeck Coefficient and Low Thermal Conductivity.


DOI: 10.1021/acsami.9b16663

ACS Appl. Mater. Interfaces 2020, 12, 1024–1035


