Water Stability of Metal-Organic Framework HKUST-1

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Abstract  Metal-organic frameworks (MOFs) are a new class of crystalline porous materials, having wide pores which provide extremely high specific surface areas, larger than those typically observed in more common porous materials like zeolites or activated carbons. This peculiar property combined with the frequent presence of open-metal sites, makes them very promising for a broad range of applications, spacing from gas storage or separation, drug delivery, toxic air removal, chemical detectors, etc. However, the industrialization of MOFs is still facing difficulties because of their low stability to water or even air moisture, a substance difficult to avoid in applications like those mentioned. Another issue concerning MOF industrialization consists in the low bulk density of the micrometric powder grains, which typically constitute such materials. Unfortunately, their low mechanical stability seems to make difficult even to enhance the packaging by simple mechanical compaction. In this review, we will particularly focus on the current state of art involving the MOF HKUST-1, especially on the degradation process involved when this MOF interacts with water molecules. Furthermore, we will show the connection between water and mechanical stability, bringing to attention of a study where solid tablets of HKUST-1 powder have been realized without any loss of crystallinity or porosity because of an accurate study on the effects of different degree of hydration during the tableting phases. In addition, in order to highlight the causes of damages induced in the framework upon interaction with water, a comparison with other two copper carboxylate MOFs will be shown, namely STAM-1 and STAM-17-OEt, which differ from HKUST-1 uniquely for the organic ligands.

Keywords  water stability, HKUST-1, mechanical compaction, MOF hydrolysis pattern, degradation process

Introduction

Metal-organic frameworks (MOFs) are a promising class of porous crystalline materials with great potentialities in a wide range of industrial applications. Composed of combinations of metallic knots and organic linkers, more than 70,000 different structures are currently recognized as MOFs.\textsuperscript{[1-9]} Their specific surface area typically exceeds 1,000 m$^2$/g, reaching sometimes even 10,000 m$^2$/g.\textsuperscript{[10]} Such a remarkable porosity is due to the possibility to combine up to 50 Å long extended organic molecules via different possible configurations, arising from the coordination geometry of the metallic junctions and it opens the doors for applications in gas storage (H$_2$, CH$_4$, ...) and then in the field of renewable energies.\textsuperscript{[1-3,9]} If the metallic junction is composed of a polynuclear cluster arising from a chelation of two or more metal ions, it is called secondary building unit (SBU).\textsuperscript{[1-3,5]} The metal sites may also have a particular affinity with some gaseous substances, thus allowing to selectively adsorb pollutants like CO$_2$, NH$_3$, cyclohexane and many others, leading to potential applications in the removal of toxic industrial chemicals.\textsuperscript{[2,3,7-12]} However, only very few applications have been already introduced in commercial use,\textsuperscript{[13,14]} mainly because of two big challenges: the extremely sensitivity to water (even to air moisture)\textsuperscript{[8,10]} and the typical poor packaging of MOFs, generally synthesized in the form of powders with low bulk density.\textsuperscript{[15,16]} Both the issues have been constantly investigated by researchers, but several significant aspects are still unclear. Examples of these are the deep knowledge of the hydrolysis/dissociation mechanisms induced by the interaction with water molecules or even finding cheap methods, which might allow to produce water-resistant or well-packaged MOFs without loss in porosity, crystallinity, or performance.\textsuperscript{[8,15,16]} In this review, both these topics are covered, specially focusing on the water stability of the MOF HKUST-1, after a brief overview based on MOFs in general. Even other two MOFs will be treated, STAM-1 and STAM-17-OEt, composed of the same SBU commonly called paddlewheel but different organic linkers which lead to markedly diverse properties.

Since its first appearing in a scientific journal in 1999, HKUST-1 or Cu-BTC, has been one of the most investigated MOFs.\textsuperscript{[18,19]} In fact, it is widely used as model for MOF research because of the ease of getting a high quality industrial-scale HKUST-1 material and for its peculiar properties as selective adsorbent.\textsuperscript{[51,12]} The other two MOFs shown in this review, STAM-1 and STAM-17-OEt, are much more recent but very promising, especially in the research field.\textsuperscript{[20,21]} In fact, despite the similarities with HKUST-1, they have a high water stability and a flexible crystalline structure, which means that the crystal undergoes spontaneous lattice transitions when subjected to determined external stimuli.\textsuperscript{[20-23]} A deeper understanding of this aspect can contribute to giving a significant boost to the search of stable MOF structures.

Water Stability in Metal-Organic Frameworks

"Water is everywhere!" is the overtone of the review about the water adsorption in MOFs written by Canivet and co-authors.\textsuperscript{[15]} In fact, even though MOFs have great potentialities for methane or hydrogen storage and for the capture of carbon dioxide from flue gases,\textsuperscript{[4,7,11,12,15,24]} outside of the laboratories natural or industrial gas streams or even the simple air have generally significant percentages of water...
vapor/humidity, and therefore the MOF behavior in the presence of water is a critical aspect to investigate.\[8\] Even worse, for example, water vapor is actually one of the main components in industrial flue gas, then it cannot be ignored when studying performances in the capture of gases like CO\(_2\). Unfortunately, even the first discovered MOFs, MOF-5 and HKUST-1, turned out to be strongly water-sensitive with critical consequent effects, and also afterwards it has been remarked that, generally, MOF performances in humid environments are typically not enough to fit the industrial requirements.\[8,18,19\] A possible dehydration step of bio-fuel streams is downright unthinkable, because it would require an energy exceeding that released by the combustion process itself.\[8\] However, it is also true that, for determined applications, certain soluble MOFs called Bio-MOFs are interesting for in vivo medical applications, for example, as drug carriers.\[7,10,15\] Introducing a definition as general as possible, a water stable MOF should have the property that, after handling it in laboratory air and/or exposing it to a reasonable amount of air moisture, the porous structure is retained.\[8\] In addition, from an industrial point of view, it might also be crucially important that the water-stable material could return to its original form at the end of each cycle of adsorption/gas separation or purification; in this cases, a reversible hydration process would be required.\[8\]

### Structural factors linked to water stability

Since the beginning of the first MOF synthesis, the strength of the metal-ligand bond has constituted a problem. The first MOFs, indeed, frequently lacked of architectural stability and that typically collapsed upon removal of solvents or exchange of ions in the pores, and then of a total absence of chemical stability because of such relatively weak bonds.\[1\] After the introduction of charged chelating linkers (whom the first one was actually the BTC linker), the metal-ligand bond has been increased and, additionally, such change of the linkers could balance that of the metal centers and avoid ion accumulation and even the interpenetration. Because of the combination of such linkers with SBUs, the removal of the solvent did not lead to the collapse of the structure for many new MOFs.\[1\] However, such bonds are frequently strong enough to avoid water decomposition.

After several efforts and compared studies, some structural factors governing the stability of MOFs in presence of water have been individuated.\[1,8\] Generally, MOF water resistance may depend on both the electronic and the steric effects of the ligand on the metal node.\[8,15\] In fact, the main structural property that makes a MOF water stable is the inertness of the metal cluster: in this case, the degradation mechanism is not favored.\[8,15\] Another point of view for the study of the (in)stability of a MOF is to evaluate the strength of metal-ligand coordination bond, because it typically represents the weak point of the structure.\[8\] Even if a specific MOF has not an inert metal, there are cases in which we can talk of kinetic stability: this latter property is governed by the activation energy barrier, which depends on the specific pathway of the potential dissociation reaction.\[8\] Two examples of kinetic factors, which can play an important role in the increase in hydrolysis activation energy, are hydrophobicity and ligand steric.\[8\] The reasoning at the basis is that a dissociation mechanism takes place only if the oxygen of a water molecule can sufficiently approach the metal to allow the establishment of an interaction between the electron orbitals.\[8,15\] In addition, it is also necessary that the energy associated to such interaction is large enough to exceed the activation energy barrier of the reaction.\[8,15\] It is expected that MOFs having this characteristic could decompose when submerged in liquid water but they would be stable after exposure to high humidity conditions.\[8\] Despite this, there are few notable exceptions which does not have inert metals, but are thermodynamically stable: probably in some cases so strong kinetic factors are involved that they make the barrier of the irreversible degradation fatal high.\[8,25,26\]

Two main degradation mechanisms acting in MOFs exposed to water have been outlined:

1. The ligand displacement reaction, which involves the inclusion of a water molecule into the metal-ligand bond;
2. The hydrolysis reaction, which implicates the breakage of the metal-ligand bond and the dissociation of the water molecule.\[15,27\]

The more diffused dissociation mechanism in MOF field is the latter and, as shown in the following section, it is also the reaction that occurs in HKUST-1.

Pore hydrophobicity is another property that can avoid water adsorption into the pores of a MOF or it can simply prevent the formation of water clusters around the metal centers.\[8\] In fact, water clustering within MOF pores seems to have a significant contribution in promoting degradation reactions.\[8,25,26\] For example, some works have reported on degradation mechanism phenomena in IRMOFs, showing that the breakage of the metal-oxygen bonds had happened only upon the formation of water clusters next to the metal centers.\[28,29\]

As it is probably clear at this point, a deep understanding about the breakdown mechanism of MOFs interacting with water is a crucial step for their development in commercial or industrial applications and technologies.\[8,18,19\] In the following section, the state of the art about the stability of HKUST-1 will be developed, explaining HKUST-1 decomposition process in great detail.

### HKUST-1

HKUST-1 (Hong Kong University Science and Technology) or Cu\(_2\)(BTC)\(_2\) was synthesized in 1999 by Chui and coworkers, and it is formed by copper paddlewheels and tritopic benzen-1,3,5-tricarboxylate (BTC) linkers.\[18\] It has a face-centered-cubic crystal lattice, with three distinct internal pores, displaying openings of about 9, 11 and 14 Å, respectively.\[18,19,30\] By standard synthesis protocols, HKUST-1 crystalizes with a water molecule coordinated to each Cu\(^{2+}\) ion.\[18,19,31\]

The dehydrated structure is typically obtained by a simple heating process at temperature ranging from 370 to 420 K under low pressure.\[19,18,31\] The desorption of the water molecules from the copper ions leads to the changing of the color of the MOF from turquoise to dark-blue.\[18,31,32\] Its bulk density is about 0.4 g/cm\(^3\), whereas its calculated crystalline density (for a dehydrated HKUST-1) is ~0.88 g/cm\(^3\).\[33,34\]

It is well-known that massive hydration conditions or water cycles seriously compromise the structure and the performances of HKUST-1.\[8,15,21,35-46\] However, only few works face the water degradation issue at atomic scale level.

Among the various points of view and different hydrolysis models, the main cornerstone is that HKUST-1 is a very hydrophilic MOF: in fact, it is well-known that polar molecules like water have a large affinity for the Cu\(^{2+}\) of the paddlewheel and such copper ions are defined open-metal sites for the MOF.\[8,15,31,35-46\] In contrast to some other MOFs which may show the coordination sites blocked by ligands, in HKUST-1 the metal sites are on the walls of the pores, making the Cu sites completely accessible.\[8\] For this reason, the dehydration process for this kind of MOFs is also frequently named activation, because it makes the open metal sites active. If the color of the dehydrated MOF is dark violet/blue navy, it only takes few minutes of exposure to air moisture to drastically change it.
again into light turquoise.\textsuperscript{[31,35,41,51-53]}

However, in general, the copper ions are not the only possible HKUST-1 adsorption sites for small molecules. In this regard, several systematic searches of all the potential and preferential adsorption sites have been deeply carried out via molecular simulations.\textsuperscript{[38,47,53-55]}

In Figure 1, the main potential adsorption sites of HKUST-1 are represented with the spheres marked by the numbers I, I’, II and III. It has been observed that a water molecule preferentially adsorbs close to Cu\textsuperscript{2+} and then in sites I and I’, whereas apolar molecules like methane adsorb preferably at the site III.\textsuperscript{[47]}

![Figure 1](image1.png)

**Figure 1** Representation of possible sorption sites I, I’, II, and III in the structure of HKUST-1 viewed along the [100] direction. Copper atoms in pink, oxygens in red, carbons in grey, hydrogens in white. Reprinted with permission from J. Am. Chem. Soc. 2012, 134, 1466–1469. Copyright 2012 American Chemical Society.\textsuperscript{[34]}

More in detail, site I corresponds to that occupied by the water molecules directly coordinated with Cu, whereas in site I’ there can be found water molecules bonded through hydrogen bonds to water molecules directly adsorbed at site I.\textsuperscript{[34,47]}

Gul-E-Noor and co-authors\textsuperscript{[46]} confirmed and deepened this result experimentally by a \textsuperscript{1}H MAS NMR study. They studied the water-MOF interaction for HKUST-1 during a continuous hydration under air moisture for 80 h. Their interpretation of the data is based on the idea of a hydration process which can be divided in 2 main steps:

(1) During the first phase of air exposure, they have observed the progressive adsorption of water on the unoccupied metal sites (i.e., sites I), detecting a co-existence of paddlewheels with either one or two water molecules per Cu pair (namely species H\textsubscript{2}O-I and H\textsubscript{2}O-II, respectively). With the increasing of the exposure time up to 20 h, the number of paddlewheels with two water molecules has grown. The authors have estimated an instant value of 1.2 H\textsubscript{2}O/Cu at exposure time of 9 h 30 min.\textsuperscript{[46]}

(2) After 20 h, they have observed the appearing and then the increasing of a new signal, which have arisen from paddlewheels with more than two water molecules per Cu pair (namely species H\textsubscript{2}O-III), due to the addition of a second water molecule per Cu site (which corresponds to the occupation of site I’ of Figure 1). Furthermore, from the 56\textsuperscript{th} to the 80\textsuperscript{th} hour, only this signal is detected on the NMR spectra with no further changes in intensity.\textsuperscript{[46]}

![Figure 2](image2.png)

**Figure 2** Water adsorption/desorption isotherms of HKUST-1 measured at 25 °C. Adsorption – full points, desorption – empty points. Reprinted and adapted with permission from J. Phys. Chem. C 2018, 123, 1730–1741. Copyright 2018 American Chemical Society.\textsuperscript{[56]}

In the lower pressure range (\(p/p_0^\ast = 0.1–0.3\)), the curve consists of a two-step trend, indicating two distinguishable processes. Presumably, the first one indicates water coordination to the copper sites and, subsequently, the second process represents either the filling of the rest of the large hydrophilic pores or the filling of the smaller ones, which have no metal site and then have a hydrophobic character, enhanced by the BTC linkers.\textsuperscript{[38]} Concerning higher relative pressures, at about \(p/p_0^\ast = 0.4\) there is a saturation region and after \(p/p_0^\ast = 0.9\) another slight increases, probably due to the water condensation inside the pores.\textsuperscript{[38]} The small hysteresis of the desorption branch can be attributed to the strong hydrogen bonds established among the water molecules adsorbed.\textsuperscript{[39]} A residual of not-desorbed water was observed in the lower pressure range: the cause is probably the chemisorption-favorable nature of the bonds established by the water molecules on the copper sites, meaning that high temperatures are always necessary to fully desorb such molecules and then activate the metal sites.\textsuperscript{[8,39]}

### Models Concerning HKUST-1 Water Degradation

Many works have drowned the possible decomposition processes, which might take place in HKUST-1 framework. The scope of this section is to offer an overview of the main common cornerstones of such models, although it is quite difficult to summarize them in a single all-inclusive picture, since each work is based on experimental setups which had involved different environments with different relative humidity, different exposure time, hydration conditions and temperatures as well as consequent effects studied from completely different points of view. For example, although all of these works generally show PXRD patterns with drastic reduction in the intensity of the peaks or a dramatic decrease of the BET area values after
a certain degree of hydration, it is hard to definitively establish after which hydration level the first traces of hydrolysis occur within the lattice and, even more important, the details on what happens at atomic scale level.[8,15,21,35,38,41] Despite this, with some precautions, it is possible to define a general description of the water degradation process. For example, a common conclusion is that the decomposition mechanism starts with the breakdown of one or more Cu—O bonds of the paddlewheel unit. Ab initio molecular dynamics (AIMD) simulations have shown that the introduction of water molecules in the framework leads to a “violent oscillation” of the Cu—O bond length, whereas the first occupation of a metal site causes a distancing between the copper and its adjacent bonded oxygens.[55] In the view of Xue and co-authors,[55] the paddlewheel unit with the elongated Cu—O bond lengths may be considered as a precursor state for the MOF hydrolytic breakdown.[55] However, it has been estimated that high water concentrations occur for the trigger of the hydrolysis.[55] Another common result reached in many other works is the identification of the water condensation (or clustering) phenomenon inside the pores as a necessary condition for the begin of the hydrolysis in HKUST-1.[8,15,21,38,41,42,51,57] As already mentioned in the previous section, this idea has been widely applied to MOFs. In particular, the presence of condensed water as the hydrolysis begins was experimentally determined by Tan and co-authors,[42] via pressure and temperature dependence studies. They have also observed that the HKUST-1 structure had remained intact when the experimental conditions did not favor the water condensation. Accordingly, Gioine and co-authors[67] showed that HKUST-1 samples exposed to water vapor significantly above the boiling temperature (about 420 K) were surprisingly stable whereas at lower temperatures like 350 K, they were readily hydrolyzed at lower temperatures like 350—370 K, at which water can condense.

Details about the HKUST-1 degradation process

Depending on the water content inside HKUST-1 network, different decomposition products may be observed.[20,40,57,58] One of the most significant identifications of such products has been done by Todaro and co-authors, by the use of EPR spectroscopy.[58] In the following the structural processes taking in HKUST-1 upon interaction with air moisture for a long time will be explained in detail.

The hydrolysis effects have been extrapolated by analyzing the changes registered in the EPR spectra acquired at both 77 and 300 K, which are reported in Figures 3 and 4, respectively.[55]

![Figure 3](image3.png)

**Figure 3** EPR spectra obtained at 77 K for a sample of HKUST-1 powder at different times of exposure to air moisture. Reprinted with permission from J. Phys. Chem. C 2016, 120, 12879–12889. Copyright 2016 American Chemical Society.[55]

![Figure 4](image4.png)

**Figure 4** EPR spectra obtained at 300 K for a sample of HKUST-1 powder at different times of exposure to air moisture. Reprinted with permission from J. Phys. Chem. C 2016, 120, 12879–12889. Copyright 2016 American Chemical Society.[55]

Furthermore, enabling the hydration of the HKUST-1 sample exclusively through the small opening of the long and narrow EPR glass tube (about 0.3 cm diameter and about 16 cm in length), they have been able to slow down the hydration so that they could identify and analyze each step of the process in great detail.[55] Combining their results with other similar studies, the main stages of the decomposition pathway identified until now are resumed in the following paragraphs.[35,37,43,44,55]

**First stage: the water adsorption on the Cu sites.** As above mentioned, the preferential adsorption site in HKUST-1 is on the copper, and then the first stage is the simple adsorption of the water molecule on the Cu²⁺ ions. In fact, upon adsorption, the oxygen of the adsorbed H₂O shares a free electron pair with the Cu²⁺ ion, establishing an out-of-plane bond.[5,55] This would lead to an elongation of the Cu—O bonds.[55] Xue and co-authors[55] observed by DFT calculations that the first adsorbed water molecule might already tilt down and break the Cu—O bond, leading to a ligand displacement (Figures 5 and 6), but water dissociation is both thermodynamically and kinetically unlikely at this stage, and therefore hydrolysis is not yet favored at this low water loading.[55]

The elongation of the Cu—O bonds also causes an expansion of the dimensions of the unit cell of the framework, detectable by PXRD.[55]
Todaro and co-authors observed during the first 20 days of exposure to air moisture only reversible changes to the EPR spectra of their samples. As already mentioned, the paramagnetic centers present in an activated HKUST-1 sample are the solely E'1(Cu) triplet centers (namely the S = 1 spin centers of the coupled paddlewheel of the activated HKUST-1) and [Cu(H2O)6]2+ centers, which are synthesis defects also present in the pristine material.53

The main effect observed during this period of 20 days is a strong reduction in intensity (about 65%) of the EPR signal obtained at room temperature: a possible interpretation given by the authors was a drastic reduction of the concentration of E'(Cu) centers53 such effect would be due to the formation of a C = bond between the Cu2+ ions within the paddlewheel.53 The formation of this bond would not damage irreversibly the network, but it would make the paddlewheels involved EPR silent.53 Also, the BET area values obtained by nitrogen iso- thermal did not show any reduction, supporting the hypothesis of a preserved framework. According to the authors, the hypothesis that the authors have obtained a complete recovery of the EPR signal.53 In Figure 7, the stage just described is represented by the transition from (a) to (b). However, subsequent measurements carried out by EXAFS fits showed that there is a distance of the two Cu2+ ions, making the establishment of an O bond between them hardly probable. For this reason, the question has remained open but, anyway, it is certain that the first stage consists of a softly hydrated material affected by a fully reversible change.

Second stage: the beginning of hydrolysis. From the 20th to the 50th day of hydration, Todaro and co-authors observed drastic changes in the EPR spectra of their samples. Those obtained at room temperature showed a narrowing of the resonance, whereas those acquired at 77 K exhibited an intense resonance centered at about 320 mT, similarly to the peak ascribed to the S = 1/2 spin centers of the [Cu(H2O)6]2+ but with a lineshape well distinguishable from that.53 Similar changes in the EPR signal have been also found elsewhere.14 According to their interpretation, such changes are due to the appearing of a new paramagnetic center within the material, which has been named E'(Cu).53 This new EPR signal would arise from the progressive formation of Cu2+/Cu3+ paddlewheels, which contribute to the EPR signal as S = 1/2 spin centers.

More in detail, the oxidation state of one of the copper of the paddlewheels (the non-silent ones) changes from 2+ to 1+, but the resulting Cu+ ion is EPR silent: therefore, the EPR signal of E'(Cu) centers arise only from one Cu2+ of the pair in such new paddlewheel structures (Figure 7).55 The hydrolysis of the Cu—O bonds, leading to the partial detaching of a carboxylate bridge and the reduction of the coordination number from 5 to 4, allows the change in both the oxidation state and in the geometry, which becomes tetrahedral.55 The adsorbed water substituting the ligand is dissociated and it has transferred one of the two hydrogens to the oxygen of the carboxylate bridge, after overcoming a very low barrier.55 However, this configuration is only metastable because it suffers to a relevant local strain, and then it quickly relaxes to another structure. The presence of strain is also suggested by the absence of the hyperfine structure on the peak centered at ~320 mT.55 This stage of the process is already irreversible: it has been demonstrated that, once at this phase, the attempt of an activation process is not sufficient to recover the initial EPR signal. The peaks of the corresponding diffraction pattern also show a significant intensity reduction. This stage can be therefore taken as the beginning of the HKUST-1 hydrolysis.55 These results are supported also by many other works that have found a lowering of the coordination of the copper units, leading exactly to the formation of Cu2+/Cu1+ unites, affecting the catalytic performances and the chemical reactivity.55

Xue and co-authors have found that at least three water molecules per paddlewheel in its local environment occur to totally detach one ligand.55 However, their model of “decomposed paddlewheel” obtained via DFT calculation was slightly different from that reported by Todaro and co-authors: indeed in their structure the Cu—O bond involving the second Cu2+ has been broken by ligand displacement and, even though the whole ligand has been consequently detached, each copper ion maintains the same number of bonds, because the water oxgens take the place of the carboxylate oxgens (Figure 6).

Third stage: the relaxation into a more stable configuration. From the 50th to the 180th day, Todaro and co-authors observed a further progressive narrowing of the resonance detectable at 300 K and also the presence of a characteristic hyperfine quadruplet on the main peak visible at 77 K and arising from the Cu2+/Cu3+ paddlewheels (Figures 3 and 4). This would be the effects of the last stage of the decomposition process. Todaro and co-authors have described how the
relaxation leads to a new $S = 1/2$ spin center named $E'2(Cu)$ (Figure 7e), arisen after the further detaching of a second carboxylate bridge from the $\text{Cu}^2+/\text{Cu}$ paddlewheel of the $E'1(\text{Cu})$ center.\[35\] The new structure has then two detached carboxylate bridges facing one another, so that the $\text{Cu}^+$ has a $T$-shaped configuration and the $\text{Cu}^2+$ ion, coordinated to a water molecule, maintains the square-pyramidal geometry.\[35\] The characteristic hyperfine quadruplet, again easily resolvable in this last stage, would indicate the loss of the strain found in the $E'1(\text{Cu})$ center.\[35\]

Forth stage: the conversion to monomeric complexes. Todaro and co-authors suggested that the $E'2(\text{Cu})$ center is stable enough to be considered the final stage of the decomposition process.\[35\] In contrast, Pöppl and co-authors\[35\] have also studied the degradation process occurred in HKUST-1 upon exposure to air moisture, and they individuated a strong similarity between the spectroscopic parameters of the final product of the decomposed HKUST-1 and those of the $[\text{Cu}(\text{H}_2\text{O})_6]^{2+}$ monomeric complexes.\[43,44\] For this reason, they suggest that in the final products of the decomposition process of HKUST-1 exposed to air there are coppers completely cut off from the framework and possibly arranged in mononuclear complexes.\[43\] Similar results involving the presence of a subsequent stage involving only paramagnetic monomeric $\text{Cu}^{2+}$ centers have also been suggested by Terracina and co-authors.\[60\]

The hydrated HKUST-1 samples have been investigated also by SEM images (Figure 8) acquired after different exposure time by Todaro and co-authors.\[35\] Even though no morphological change is observed after the first 20 days (that is, when only $E'(\text{Cu})$ silent center and $E'(\text{Cu})$ centers co-exist in the material) compared to the pristine material, after 36 and 188 days of hydration, small holes appear on the surface of the grains. Thus, these results are in perfect agreement with the EPR results above mentioned.\[35\]

Figure 8 SEM images of a sample of HKUST-1 exposed to air (300 K, 70% RH) for different times: (a) pristine material (0 days of hydration), (b) 20, (c) 36, and (d) 188 days. Reprinted with permission from J. Phys. Chem. C 2016, 120, 12879–12889. Copyright 2016 American Chemical Society.\[35\]

Other considerations

In the last years, several efforts have been spent on the research of methods to preserve HKUST-1 (or more in general water sensitive MOFs) from destructive interaction with water.\[58,61,64,65\] For example, in some works many researchers have tried to introduce hydrophobic component which would fill the pores limiting the possibility of water condensation or to build a kind of surface barrier, like an external hydrophobic layer.\[58,61,64,65\] Unfortunately, these methods typically require a not negligible price in terms of pore accessibility.\[58,64\] Other studies have focused on the fact that hydrolysis induces superficial defects already creating a sort of surface barrier, which prevents the interaction of the internal pores with potential external gases or substances. In principle, healing such surface barrier might recover the properties of the pristine MOF.\[66–70\] For example, Müller and co-authors\[66\] found an easy method to dissolve such surface defects and then recover the pristine performances; however, such dissolving process led to a loss of about 20% in mass of the material. Finally, McHugh and co-authors\[51\] suggested that hydrolytic stability can be induced in a MOF like HKUST-1 introducing sacrificial bonds between the Cu centers and the carboxylate ligands. Such idea led to the design of the MOF STAM-17-OEt, which we will discuss in the section of “Comparison with Other Copper Carboxylate MOFs: STAM-1 and STAM-17-OEt”.

Role of humidity on mechanical compaction processes

Recently, Terracina and co-authors\[56\] deeply characterized the role and the effects of the water molecules during the phases of the production of HKUST-1 tablets. This detailed study has been made possible by the development of a new method involving EPR spectroscopy, which allows the estimation of the amount of damages induced in the MOF structure upon the application of a mechanical pressure. In other words, they have been able to recognize for the first time that, as occurs upon hydrolysis, the tabletting process always induces the formation of $S = 1/2$ paramagnetic centers in the materials, arising from the distortion and/or breaking of a variable fraction of the paddlewheels of the material. The authors have observed that the number of damaged paddlewheels increases on increasing the applied pressure. From this method, one can obtain quantitative estimates how many paddlewheels have been affected by mechanical compaction in correspondence of the metal group (Figure 9).\[56\]

Figure 9 EPR spectra of some representative tablets obtained just after tabletting and of the powder sample (in black), acquired at 77 K. The pressure applied ranges from 0 (powder sample) to 40 MPa. The insets in both the graphs (a) and (b) show the zooms of the spectrum regions enclosed by the dashed panels. Reprinted with permission from J. Phys. Chem. C 2018, 123, 1730–1741. Copyright 2018 American Chemical Society.\[56\]
Furthermore, by applying this method to the tablets produced from powder differently hydrated, they have unveiled that the number of defects (individuated as \( S = 1/2 \) spin centers emerged upon the packaging treatments) is strictly related to the level of hydration of the starting powders. In the tablets produced from HKUST-1 activated powders, a negligible degradation of the paddlewheels has been observed: the percentage of paddlewheels damaged by tableting with a pressure of about 40 MPa has been estimated to be as low as 4%. But there is more: the authors have found that an additional thermal activation just after the compaction can further reduce this defective EPR signal, the so-called “cramplike zones”, or else weak interactions between near dehydrated paddlewheels that can be broken upon interaction with water. The presence of such “sacrificial bonds” makes it “hemimble”. Unfortunately, compared to HKUST-1, they have a significantly lower inner porosity and surface area.\(^{[20,21]}\)

The BET surface areas of the tablets have confirmed that the tablets produced have an optimal inner porosity, comparable to that of the starting powder (Table 1). On the contrary, by using powder samples with different degrees of hydration, the EPR spectra acquired after their compaction have highlighted a strong presence of defects, which increase on increasing the hydration.\(^{[56]}\)

**Table 1** BET surface area of the HKUST-1 powder sample compared with that of some of the tablets produced from activated powders.

| Sample                  | BET area \((\text{m}^2/\text{g})\) |
|-------------------------|------------------------------------|
| Powder                  | 1620                               |
| 40 MPa                  | 1685                               |
| 40 MPa reactivated\(^b\) | 1935                               |
| 40 MPa, after 5 weeks   | 1848                               |
| 40 MPa reactivated,\(^a\) after 5 weeks | 1622 |

\(^a\)The analysis has been performed in the relative pressure range of the increasing trend of the Rouquerol plot. \(^b\) Reactivated just after the compaction.

Furthermore, prolonging the EPR study up to five weeks from the compaction (of activated, slightly hydrated or highly hydrated powders), it has been found that the tablets produced by activated powders and reactivated just after the mechanical tableting were the only which did not show any sign of delayed structural instability, even after five weeks.\(^{[56]}\)

Summarizing, following this specific albeit simple compaction protocol, the authors proved that it is possible to obtain a stable HKUST-1 tablet having a structure with the pristine porosity and crystallinity but with a high bulk density, simply taking care to activate the material just before and just after tableting.

**Comparison with Other Copper Carboxylate MOFs: STAM-1 and STAM-17-OEt**

In order to better understand the causes of the trigger of hydrolysis in HKUST-1 (but also, more in general, in MOFs), it can be useful to involve in the study also other similar structure. In the following, we will talk about two other copper carboxylate MOFs, which are water stable despite the similarities with HKUST-1 material. Although these two MOFs have the same copper paddle-wheel building units of HKUST-1 (Figure 10a), instead of the BTC linker (Figure 10b), the organic linkers which compose STAM-1 and STAM-17-OEt are monomethyl BTC linker (Figure 10c) and 5-ethoxy isophthalate linker (Figure 10d), respectively.

The resulting frameworks have different topologies and properties.\(^{[20,21]}\) More in detail, both STAM-1 and STAM-17-OEt exhibit flexibility triggered by adsorption/removal of water and, furthermore, STAM-17-OEt also shows the presence of significative porosity and surface area. In contrast to HKUST-1, they have a significantly lower inner porosity and surface area.\(^{[20,21]}\)

**Figure 10** Components of the MOFs HKUST-1, STAM-1, and STAM-17-OEt: (a) copper paddle-wheel unit, (b) BTC linker (for HKUST-1), (c) monomethyl BTC linker (for STAM-1), and (d) 5-ethoxy isophthalate linker (for STAM-17-OEt).

The authors, accordingly with previous literature, have supposed that when on average more than one water molecule approaches each copper ion of the paddle-wheel, the differences in evolution rise up.\(^{[60]}\)

These considerations can provide an answer to the question: why three similar MOFs have such a different water stability? In fact, such difference may be reasonably linked to the amount of free space available in the pores of the materials: indeed, the pore size is significantly larger in HKUST-1 than in HKUST-17.

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STAM-1 and STAM-17-OEt. It must be said also that other theories have been suggested, for example, involving that a significant role is determined by the hemi-lability property of the MOF STAM-17-OEt. In this view, HKUST-1 would undergo hydrolysis immediately after the adsorption of the first water molecules, whereas in STAM-17-OEt this Cu—O breakage would be prevented because of the breakage of the sacrificial bonds or “crumple zones”, which require a minor energy quantity. However, this is in sharp contrast with the common evolution between HKUST-1 and STAM-17-OEt observed during the first few hours and, furthermore, with the reversibility of such earlier phases.

Conclusions and Perspectives

Since the beginning of the research on MOFs, their susceptibility to hydrolysis has been highlighted and a huge number of works have been published in not more than two decades. A certain quantity of highly stable MOF structures have been developed, but they are only a tiny percentage compared to the totality of MOFs. Furthermore, despite being relatively easy to obtain a higher confidence about the instability of a MOF, the opposite is not simple. For this reason, more systematic experimental studies are still necessary in order to deeply characterize the water stability properties of lots of MOFs and, even more, the causes of their behavior, involving as many techniques as possible. After a long and focused work carried out on HKUST-1, a deep understanding about the degradation mechanism induced by hydrolysis has been reached, with a complete comprehension about the stability limits of the material, as well as the development of a simple protocol which allows the production of compacted HKUST-1 powders, as displayed also in this review. Even some strategies to prevent hydrolysis in HKUST-1 have been developed, although they cannot fully solve the issue, because the performances of such materials result likewise compromised. Similarly, the development of copper carboxylate MOFs which are highly water stable seems also promising even though not yet complete, since the performances of such resistant MOFs known until today are not equally promising as those of HKUST-1.

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Author Contributions

Both the authors collected relevant literatures, wrote, revised and organized this manuscript.

Conflict of Interest

The authors declare no conflict of interest.

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