

Metal-organic frameworks (MOFs) bring new life to hydrogen-bonding organocatalysts in confined spaces

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Hydrogen-bonding organocatalysis has emerged as promising biomimetic alternative to Lewis acid catalysis. Urea, thiourea and squaramide moieties represent the most common hydrogen-bond donors used for the preparation of these catalysts. However, their significant tendency to undergo self-quenching (self-aggregation) often decrease their solubility and reactivity. Recently, scientists have found a promising way around this problem by immobilizing the hydrogen-bonding organocatalysts on metal-organic frameworks (MOFs). Along with advantageous modular synthesis and recycling properties, the tunable porosity and topology of MOFs also allows fast mass transport and/or interactions with substrates. Herein, we highlight the existing examples dealing with the fabrication and testing of hydrogen-bonding organocatalyst-containing MOFs, providing also our vision for further advances in this area. The results derived from these works will likely serve as inspiration for the future development of superior hydrogen-bonding organocatalysts to accomplish in confined spaces chemical transformations that are either slow or unaffordable under standard homogeneous conditions.

Introduction

Most of the chemical reactions that occur in living systems are high-energy reactions that would be unaffordable or would occur very slowly without the catalysis provided by enzymes. This has inspired scientists during many decades to develop artificial catalysts^{1,2} within a general quest for economic profit. As a matter of fact, catalysts are nowadays the workhorses of the chemical transformations in industry (ca. 85-90% of the products of chemical industry are fabricated through catalytic processes).³ However, the increasing on-demand production of bulk chemicals, fine chemicals and fuels is also threatening the stability of the ecosystems in our planet due to environmentally harmful wastes. Thus, the stricter environmental regulations by the governments have motivated the replacement of numerous homogeneous reactions by greener and economical heterogeneous catalytic processes.⁴

Within this context, supramolecular coordination chemistry offers a promising biomimetic platform with well-defined functional groups suitable for catalysis.⁵⁻⁹ Features such as numerous metal geometries, tunable bond directionality and self-correcting kinetic reversibility have allowed the development of variety of metal-organic materials (MOMs) with major relevance in the field of catalysis.^{10,11} Among the vast field of MOMs,¹² and supported by the gradual maturation of crystal engineering,¹³ nanoscale metal-organic frameworks (MOFs)^{14,15} are taken a pivotal role in the rational design of eco-friendly heterogeneous catalysts¹⁶ beyond their more traditional applications in gas

storage and gas separation,¹⁷⁻²² sensing,^{23,24} ion exchange,²⁵ drug delivery^{26,27} and light harvesting.²⁸ MOFs are composed of inorganic nodes (metal ions or clusters) judiciously linked by polytopic functional organic *struts* to form uniform three-dimensional (3D) arrays of high surface area (up to 10400 m² g⁻¹),²⁹ large pores (98 Å)³⁰ and low density (0.13 g cm⁻³).³¹ In general, the active sites of MOF-based catalysts can be either metal centers with unsaturated coordination environments, catalysts encapsulated in the pores and the sites inherent in the organic framework.³² Along with the advantages of their modular synthesis^{33,34} and recycling properties, the tunable porosity and topology of MOFs also allows fast mass transport and/or interactions with substrates. These features, often challenging in homogeneous states, differentiate MOFs from other nanoporous materials such as zeolites and activated carbons. Overall, MOFs constitute ideal candidates for heterogeneous catalysis if the framework crystallinity and regularity are retained under the reaction conditions or during post-synthetic modifications strategies that can be used to introduce functional groups on the MOFs in order to extend their catalytic activity.^{35,36}

Despite of the extensive use of MOFs in heterogeneous catalysis during the last decade,^{32,37} some drawbacks are usually associated to unsatisfactory yields, poor selectivities or limited substrate scope.³⁷ Moreover, among the plethora of MOF-based catalysts, hydrogen-bonding organocatalysts³⁸ have met MOFs only recently. Hydrogen-bonding organocatalysts have emerged as biomimetic alternative to Lewis acid activation³⁸⁻⁴⁰ by means of mainly urea, thiourea or squaramide moieties. This is relevant because although a significant number of bond-forming reactions can be catalyzed by homogeneous hydrogen-bonding organocatalysts,⁴¹ their tendency to undergo self-quenching (self-aggregation) often decrease their solubility and reactivity.

Herein, we highlight the recent contributions involving the fabrication and use of hydrogen-bonding organocatalysts immobilized on MOFs. We believe that these unique examples are paving the way for the development of superior hydrogen-

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bonding organocatalysts for complicated chemical transformations.

Urea-containing MOF catalysts

Farha, Hupp, Scheidt and co-workers reported the first urea-containing metal-organic framework (MOF) with the idea of preventing the self-association that homogeneous catalysts, such as ureas, undergo in their reactions. This self-quenching is responsible of the consequent decrease of the solubility and reactivity of the catalysts (Figure 1, top).^{42,43} For this purpose, the authors envisioned that this self-quenching may be avoided if the urea catalyst was incorporated into coordination polymers using an appropriate metal ion, which possess defined reaction environments and high porosity (Figure 1, bottom).⁴³

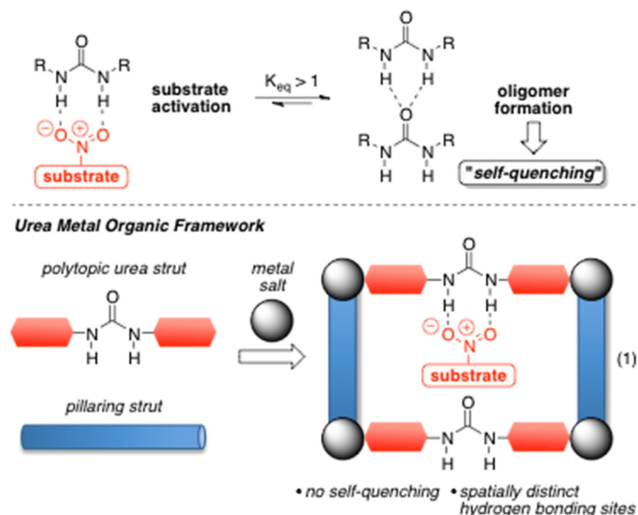
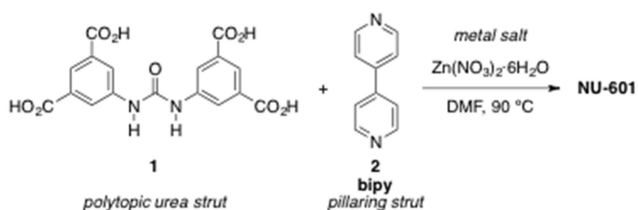


Fig. 1 Synthetic strategy of Urea MOF derivatives. Adapted with permission from ref. 43. Copyright © American Chemical Society.

Based on this pioneering idea, **NU-601** MOF was synthesised under solvothermal conditions starting from urea **1** and 4,4'-bipyridine (**bipy**, **2**) as *pillaring strut* (Scheme 1).^{43,44}



Scheme 1 Synthesis of MOF **NU-601**.⁴³

After 2 days of heating, light-yellow needle-shaped crystals of **NU-601** MOF were obtained. The structure of $Zn_2(\text{bipy})_2(\text{1-4H})$ contains 2D sheets of *strut 1* pillared with **2** and Zn_2 nodes, featuring large channels in all directions as disclosed in Figure 2.

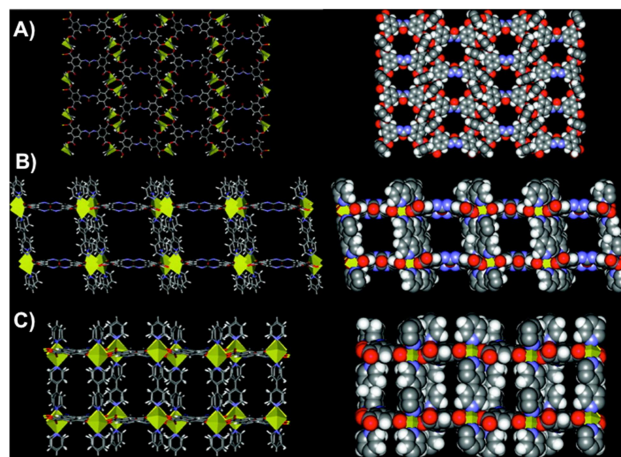
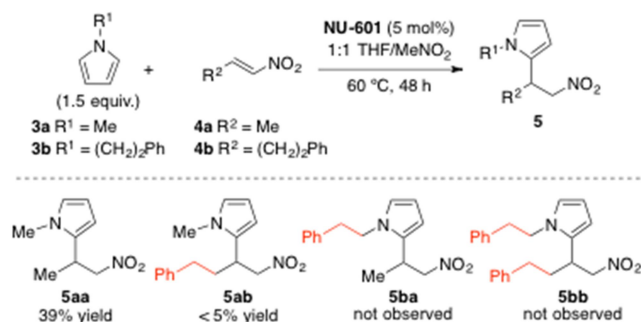


Fig. 2 Stick and space-filling representations of **NU-601** ($P2_1/c$ space group): A) view down the *a* axis (pore size 12.05 Å × 13.95 Å). B) view down the *b* axis (pore size 11.38 Å × 13.69 Å). C) view down the *c* axis (pore size 11.38 Å × 4.87 Å). Adapted with permission from ref. 43. Copyright © American Chemical Society.

Since DMF molecules remained tightly hydrogen-bonding to the N-H groups of the urea in the solid, in order to activate the structure, the exchange of the DMF with MeNO_2 during 4 days was carried out. After this exchange/activation process, the effectiveness of **NU-601** as hydrogen-bond donor catalyst was explored in the Friedel-Crafts alkylation between nitroalkenes and pyrroles.⁴⁵

In the screening of the reaction, a higher reactivity was observed when using **NU-601** as the catalyst compared with the reactivity of diphenylurea (90% conv. at 18 h vs 65% conv.) and the control reaction (21% conv. at 48 h). Then, in order to demonstrate if the catalysis took place on the surface of the MOF or within the pores, the authors employed larger substrates as steric probes (Scheme 2).



Scheme 2 Size-exclusion catalysis experiments.⁴³

For this aim, pyrrole **3b** and nitroalkene **4b** were also employed, and a rapid decrease of the reaction rate was detected. These results provided strong evidence that **NU-601** exhibited reagent size selectivity since large substrates shown significantly diminished yields because they cannot get into the porous network. **NU-601** also proved to be reusable, maintaining its reactivity after one cycle of use (33% vs 39% yield after 48 h of reaction) and suffering only a slight degradation of reactivity after five cycles (29% vs 39% yield after 48 h of reaction).

The same research group reported in 2013, as extension of

their research work,⁴⁶ a new class of urea-based **UiO-67** derivatives as plausible catalysts.⁴⁷ **UiO-67** is formed by a cubic metal-organic framework of cationic $Zr_6O_4(OH)_4$ nodes and biphenyl-4,4'-dicarboxylate (**bpdc**) 7 linkers, providing two pore sizes with diameters of 23 and 12 Å (Figure 3).⁴⁸

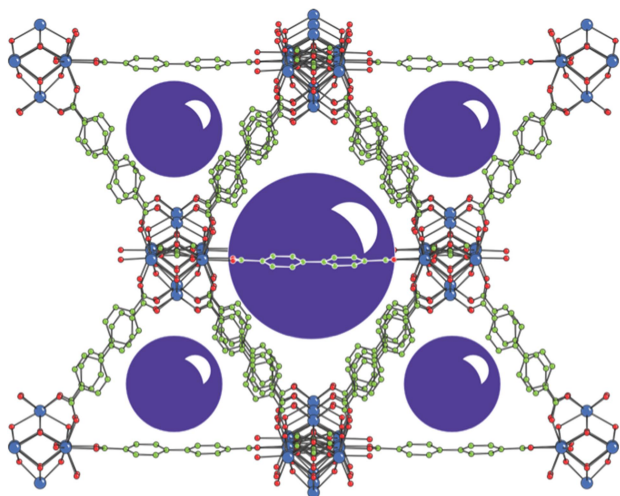
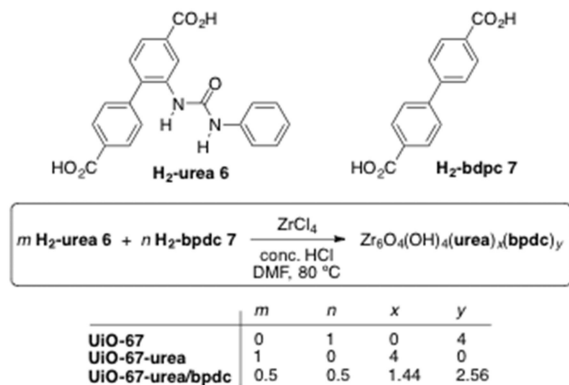


Fig. 3 **UiO-67** structure represented by a single octahedral cage (large sphere). The face of each octahedral is shared with eight smaller tetrahedral cages (small spheres). Reprinted with permission from ref. 46. Copyright © Royal Society of Chemistry.

In this study, the authors incorporated for the first time the urea scaffold into the **UiO-67** with the main aim of designing plausible hydrogen-bond donating catalysts. In order to evaluate the effect of the porosity on the catalytic activity of the materials, different species were prepared by means of various **H₂-urea/H₂-bpdc** ratios during the solvothermal synthesis (Scheme 3).



Scheme 3 Synthesis of derivatives of MOF **UiO-67**.⁴⁷

All prepared species were explored in the nitroaldol (Henry)⁴⁹ reaction between nitromethane **9** and benzaldehyde **8a** as the model reaction. The results are disclosed in Scheme 4.



Scheme 4 Screening of the nitroaldol (Henry) reaction.⁴⁷

The mixed *strut* **UiO-67-urea/bpdc** framework was found to be the most active catalyst (67% yield after 24 h). The results were very similar in the control experiment and when using $Zr_6O_4(OH)_4$ by itself (**UiO-67**) (14% and 11%, respectively). Interestingly, the use of **UiO-67-urea** afforded the same poor results (19% yield). The best catalytic activity showed by **UiO-67-urea/bpdc**, was attributed to its high Brunauer–Emmett–Teller (BET) surface area (1550 m² g⁻¹) compared with **UiO-67-urea** (390 m² g⁻¹) measured by the authors. It is remarkable that the pore size distribution on **UiO-67-urea/bpdc** did not suffer a significant decrease in the pore diameters (21.5 and 12 Å) in comparison to the precursor **UiO-67** (23 and 11.5 Å). In contrast, both structures are significantly larger than that of the pure *strut* **UiO-67-urea** (12 and 9 Å).

More recently, Hu, Liu and co-workers have developed novel functionally diverse urea-derived MOF hydrogen-bond-donating heterogeneous catalysts for their application in the Friedel–Crafts alkylation reactions.⁵⁰ The heterogeneous catalysts were synthesised following a post-synthetic modification (PSM) approach, which has been proven to be an effective tool to introduce functional groups in MOFs.^{36,51,52} Thus, three different urea-containing **Cr-MIL-101** (chromium(III) terephthalate MOF) heterogeneous catalysts were prepared. **Cr-MIL-101-NH₂** was first prepared starting from **Cr-MIL-101**, and then the former was submitted to react with the appropriate isocyanate in CH₃CN at 120 °C in a sealed tube for 12 h. It is noteworthy that both aliphatic and aromatic isocyanates were successfully anchored onto the **MIL-101** walls *via* covalent incorporation leading to **Cr-MIL-101-UR1–3** in nearly quantitative conversion and catalyst loading of 2.61, 2.48 and 1.85 mmol g⁻¹, respectively (Figure 4).⁵⁰ Gas sorption behavior of these MOFs demonstrated that they were still highly porous after post-synthetic modification (*i.e.*, Brunauer–Emmett–Teller (BET) surface area were 1409.9 m² g⁻¹, 1178.6 m² g⁻¹ and 829.3 m² g⁻¹, respectively).

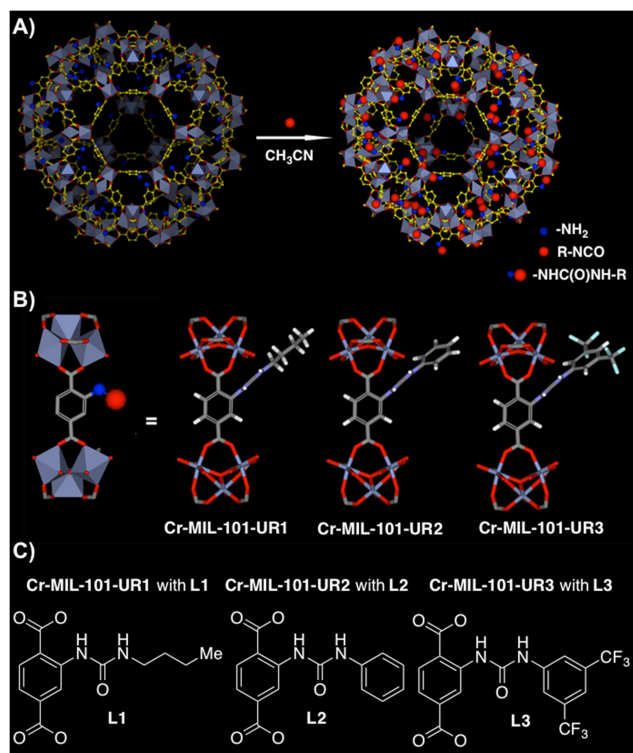
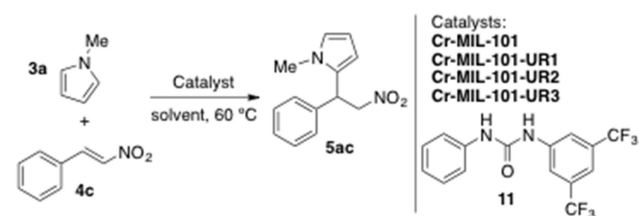


Fig. 4 A) PSM approach of Cr-MIL-101-NH₂ with isocyanates to generate Cr-MIL-101-UR1–3. B) Schematic representation of the ligand structure and [Cr₃O(CO₂)₆] cluster nodes. C) Differently substituted ureas present in the final synthesised MOFs. Adapted with permission from ref. 50. Copyright © Royal Society of Chemistry.

The catalytic activity of Cr-MIL-101-UR1–3 in the Friedel–Crafts alkylation reaction was tested in the model reaction between *N*-methylpyrrole (**3a**) and *trans*-β-nitrostyrene (**4c**) (Table 1).^{45,50}

Table 1 Screening of the Friedel–Crafts alkylation reaction.^{a,50}

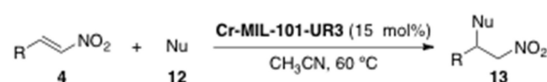


Entry	Catalyst	Solvent	Time (h)	Yield (%) ^b
1	Cr-MIL-101-UR3	DMF	36	Trace
2	Cr-MIL-101-UR3	THF	36	26
3	Cr-MIL-101-UR3	Acetone	36	27
4	Cr-MIL-101-UR3	CH ₃ CN	60	96
5 ^c	Cr-MIL-101-UR3	CH ₃ CN	24	95
6 ^d	Cr-MIL-101-UR3	CH ₃ CN	24	90
7 ^e	Cr-MIL-101-UR3	CH ₃ CN	24	73
8 ^c	Cr-MIL-101-UR1	CH ₃ CN	24	72
9 ^c	Cr-MIL-101-UR2	CH ₃ CN	24	87
10	–	CH ₃ CN	24	Trace
11	Cr-MIL-101	CH ₃ CN	48	22
12 ^c	11	CH ₃ CN	24	79

¹⁵ ^a Reactions performed with **3a** (0.2 mmol), **4c** (0.1 mmol), and catalyst (15 mol%) in the solvent (0.15 mL). ^b Determined using ¹H-NMR. ^c Ratio 5:1 for **3a**:**4c** used. ^d 10 mol% catalyst loading. ^e 5 mol% catalyst loading.

After exploring different reaction conditions and catalysts, the best results were achieved with Cr-MIL-101-UR3 (entries 4 and 20 5), which bears appropriate steric hindered and electron-withdrawing substituents. The catalyst loading could be reduced to 10 mol%, without strong influence in the yield (entry 6) and better reactivity was found in comparison with urea **11**, a homogeneous catalyst (95% vs 79% yield, entries 5 and 12, 25 respectively). The other two catalysts (Cr-MIL-101-UR1 and Cr-MIL-101-UR2) gave poorer results than Cr-MIL-101-UR3 (entries 8 and 9 compared with entry 5). The better reactivity found in the case of the later is attributable to the spatial isolation achievable in the porous MOF environment. With the optimized 30 conditions in hand, other electron-rich nucleophiles (**12a–c**) and a broad scope of different nitroalkenes (**4c–m**) were explored, achieving the same good reactivity (Table 2).

Table 2 Screening of the Friedel–Crafts alkylation reaction.^{a,50}



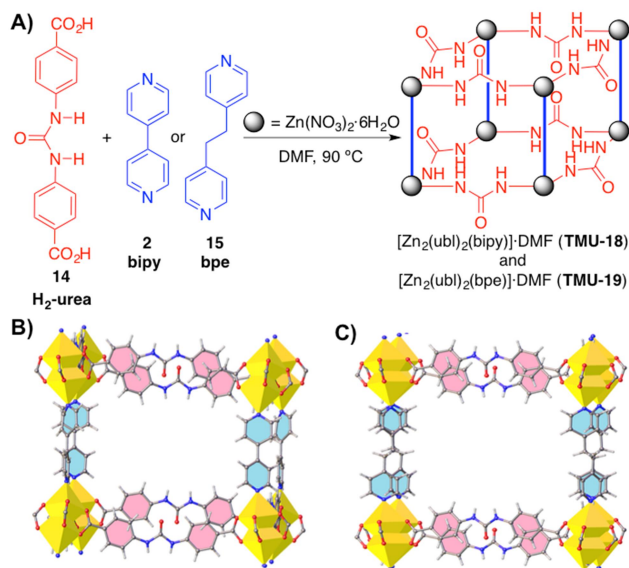
Entry	4 (R)	Nu (12)	13	Yield (%) ^b
1	4c (Ph)			95
2	4c (Ph)			93
3	4c (Ph)			94
4	4d (4-MeO-Ph)	12c	13dc	90
5	4e (2-MeO-Ph)	12c	13ec	81
6	4f (4-CF ₃ -Ph)	12c	13fc	90
7	4g (4-Cl-Ph)	12c	13gc	88
8	4h (2-naphthyl)	12c	13hc	91
9	4i (2-furyl)	12c	13ic	92
10	4j (2-thiophenyl)	12c	13jc	93
11	4k (Pr)	12c	13kc	65
12		12c	13lc	40 (80) ^c
13		12c	13mc	17 (76) ^c

³⁵ ^a Reactions performed with **4** (0.1 mmol), **12** (0.12 mmol), and catalyst (15 mol%) in CH₃CN (0.15 mL). ^b Isolated yield. ^c Isolated yield using catalyst **11**.

Interestingly, with long-chained or large nitroalkenes as 40 substrates (**4l–m**), better results were obtained using urea **11** (entries 12 and 13). This fact supports that the heterogeneous catalysis mainly occurred within the pores of the MOF, thus exhibiting reagent size selectivity. Moreover, the study of the recyclability of Cr-MIL-101-UR3 provided that the catalyst was 45 easily isolated from the reaction suspension by centrifugation and

reused at least four times with little loss of reactivity, while retaining its crystallinity as confirmed by PXRD analysis.

Recently, Morsali and co-workers have developed another example of MOFs containing urea units inside the *struts* of the MOF.⁵³ Two different MOFs (**TMU-18** and **TMU-19**) were synthesized by reacting $Zn(NO_3)_2 \cdot 6H_2O$, the urea-based ligand (**H₂-urea** **14**) and either **bipy** (**2**) or **bpe** (**15**) as *pillaring struts* under solvothermal conditions (Scheme 5). These MOFs are modified versions of the urea-containing MOF synthesized by Farha, Hupp, Scheidt and co-workers in 2012,⁴⁵ and exhibited different characteristics. Both MOFs showed large channels along the *bc*-plane with aperture size of $13.5 \times 9.9 \text{ \AA}$, for **TMU-18**, and along the *b*-axis with aperture size of $12.1 \times 10.6 \text{ \AA}$ for **TMU-19**.



Scheme 5 A) Synthesis of catalytic urea-containing MOFs **TMU-18** (triclinic *P1*) and **TMU-19** (orthorhombic *Pnna*). B-C) View along the pore direction of **TMU-18** and **TMU-19**, respectively. In contrast to **TMU-18**, TGA analysis indicated a lower thermal stability for **TMU-19**. Adapted with permission from ref. 53. Copyright © Royal Society of Chemistry.

These urea-based MOFs, especially **TMU-18**, showed good catalytic activity in regioselective methanolysis of diverse epoxides (Figure 5 and Table 3). In the case of styrene oxide (**16a**) both catalysts provided very high conversions after 140 h (100% and 95%, entry 3). However, for other three less reactive epoxides (**16b-d**) poorer conversions were obtained in general, even in the presence of double amount of catalyst (50 mol% vs 25 mol%). In all cases, very low conversions were obtained after the same reaction time when no catalyst was used (entries 1, 5, 7 and 9 vs 2, 6, 8 and 10, respectively). A fifth bulky epoxide (**16e**) was tested in order to support the happening of the reaction within the pores of the frameworks, and the low conversion reached after 140 h indicates it so (entry 11).

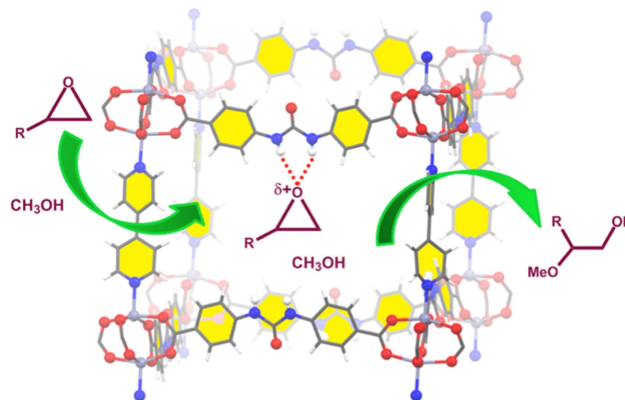


Fig. 5 Activation of epoxides by **TMU-18** for their regioselective methanolysis. Reprinted with permission from ref. 53. Copyright © Royal Society of Chemistry.

Table 3 Catalytic activities of **TMU-18** and **TMU-19** in the methanolysis of epoxides **16a-e**.^{a,53}

Entry	Substrate	Major product	Catalyst loading (mol%)	Conv. (%) ^{b,c} TMU-18	Conv. (%) ^{b,c} TMU-19
1			–	19	
2	16a	17a	25	78 (96)	72 (97)
3 ^d	16a	17a	25	100 (98)	95 (98)
4	16a	17a	50	100 (98)	95 (98)
5			–	22	
6	16b	17b	50	78 (96)	67 (91)
7			–	26	
8	16c	17c	50	64 (81)	53 (77)
9			–	16	
10	16d	17d	50	51 (78)	48 (72)
11 ^d			25	9	6

^a Reactions performed with **16a-e** (0.2 mmol) and the appropriate amount of urea catalyst in methanol (3 mL) at 60 °C. ^b GC yield using internal-standard method. ^c The data in parentheses correspond to the selectivity for the major product. ^d 140 h reaction time.

It is noteworthy to mention that the reaction conversion was lower when 40 mol% of urea powder was used as catalyst (15% vs 35% and 31%, with **TMU-18** and **TMU-19**, respectively, after 24 h), which proves the importance of including the urea moieties

within the MOF structures. In addition, both catalysts were recovered, washed and used one more time, but they showed a decrease in their catalytic activity (39% vs 45% and 29% vs 41% for **TMU-18** and **TMU-19**, respectively, after 40 h) in agreement with a gradual loss of crystallinity through the cycles as verified by PXRD measurements. However, these results open new possibilities for designing reusable urea-containing MOF catalysts for reactions carried out in aggressive media for the metal centers of the MOFs such as MeOH.

Contemporaneously, Wang, Zhao and co-workers, along with their research line,⁵⁴ designed a novel (3,24)-connected *rht*-type metal organic framework (MOF **Cu-UBTA**). It was prepared by a solvothermal reaction between an unsymmetrical hexacarboxylate ligand **H₆-L (18)** containing a urea group and **Cu(NO₃)₂** in DMF at 75 °C for 3 days (Figure 6).⁵⁵

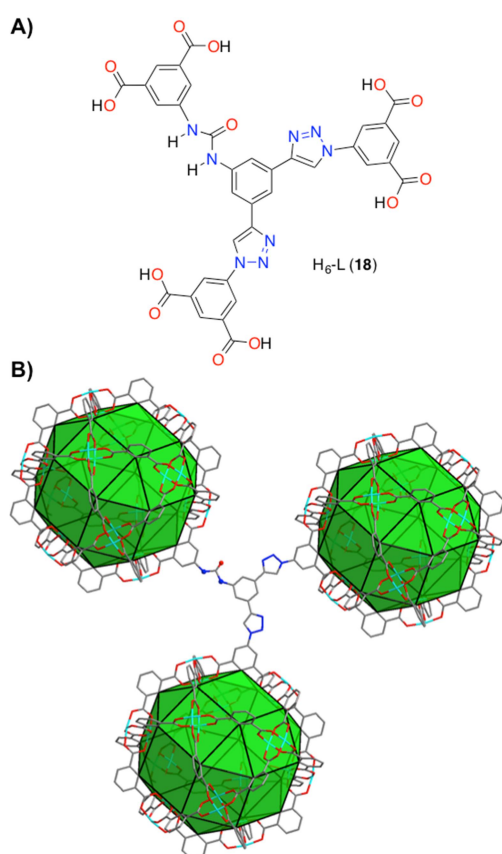


Fig. 6 A) Structure of the hexacarboxylate acid **H₆-L (18)**. B) Single crystal structure of the (3,24)-connected *rht*-topological framework in **Cu-UBTA**. Tetragonal space group *I4/m*, unit cell dimensions $a = b = 31.4206 \text{ \AA}$, $c = 44.8239 \text{ \AA}$. BET surface area = $3134 \text{ m}^2 \text{ g}^{-1}$. Adapted with permission from ref. 55. Copyright © Royal Society of Chemistry.

As a proof of its high porosity, the application of **Cu-UBTA** in gas storage and separation was further evaluated towards **CO₂**, **H₂** and **N₂**. The results disclosed in Figure 7 suggested a high selectivity with **CO₂** and **H₂** against **N₂**.

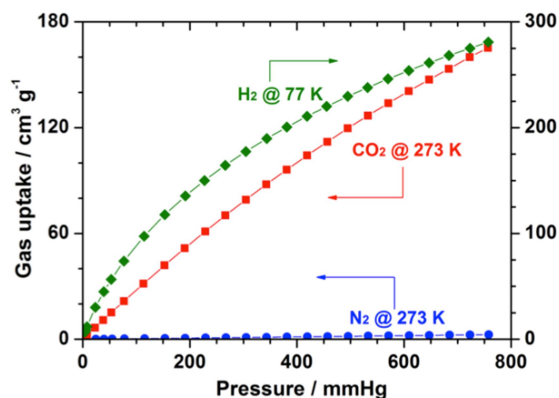
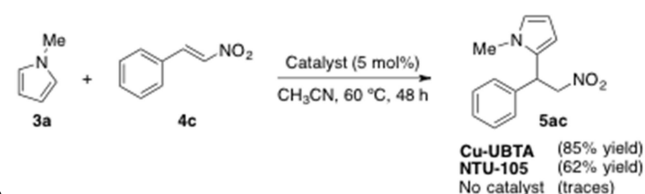


Fig. 7 Gas sorption isotherms of **Cu-UBTA** for **CO₂** and **N₂** (at 273 K), and **H₂** (at 77 K). Reprinted with permission from ref. 55. Copyright © Royal Society of Chemistry.

The catalytic activity of **Cu-UBTA** was also evaluated using the Friedel–Crafts alkylation reaction between *N*-methylpyrrole (**3a**) and *trans*- β -nitrostyrene (**4c**) as a model reaction (Scheme 6).



Scheme 6 Model Friedel–Crafts alkylation reaction catalysed by **Cu-UBTA**.⁵⁵

With **Cu-UBTA** the desired product was obtained in 85% yield after 48 h, whereas only traces of the product were observed in the control experiment without catalyst. In order to confirm the utility of the urea moiety in the catalytic system, the authors used their previous MOF **NTU-105**⁵⁴ without urea as the catalyst under the same reaction conditions. Interestingly, **NTU-105** exhibited a lower catalytic activity in relation to **Cu-UBTA**. This fact confirmed that the urea group in *rht*-MOF **Cu-UBTA** could also function as a hydrogen-bond-donating site to catalyse the Friedel–Crafts reaction in a synergic effect with the open metal sites, since the moderate catalytic activity shown by **NTU-105** was attributed to the Lewis acidic open copper sites in the structure. The scope of the application of this catalyst was extended to other nucleophiles (**12c**) and nitroalkenes **4**, leading to the desired products with good to excellent yields (Table 4). The recyclability of the MOF was also explored and the catalyst was reused after a simple centrifugation, showing similar catalytic activity in successive reaction cycles (81% for the 2nd run, 75% for the 3rd run). Moreover, PXRD measurements of the recycled catalyst revealed that the MOF retained its crystallinity after the catalytic cycles.

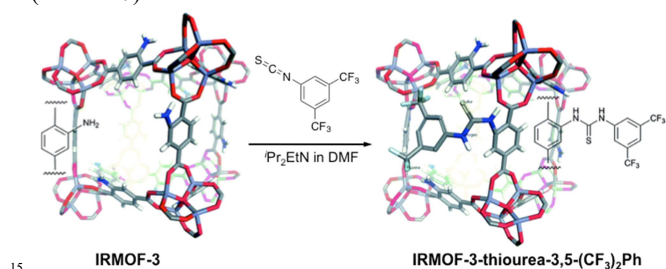
Table 4 Scope of the Friedel-Crafts reaction.^{a,55}

Entry	4 (R)	Yield (%) ^b 3a + 4	Yield (%) ^b 12c + 4
1	4d (4-MeO-Ph)	92	76
2	4f (4-CF ₃ -Ph)	92	91
3	4g (4-Cl-Ph)	91	79
4	4i (2-furyl)	86	47
5	4j (2-thiophenyl)	87	66
6	4l (4-Me-Ph)	88	76
7	4m (4-t-Bu-Ph)	93	63
8	4n (4-F-Ph)	91	83

^a Reactions were carried out with **3a** or **12c** (0.2 mmol), **4** (0.1 mmol) and catalyst **Cu-UBTA** (5 mol%) in CH₃CN. ^b Determined by ¹H-NMR using internal-standard method.

Thiourea-containing MOF catalysts

With the idea in mind to prevent also the self-quenching and to make possible the recycling of thiourea catalysts, Wang and co-workers have recently immobilised thiourea functionalities into **IRMOF-3** following a post-synthetic modification strategy (PSM).⁵⁶ The synthesis was addressed with **IRMOF-3**, a porous MOF with primary amine groups in its structure, which was further modified adding an isothiocyanate under basic conditions (Scheme 7).



Scheme 7 PSM approach between **IRMOF-3** with 3,5-bis(trifluoromethyl)phenyl isothiocyanate. Adapted with permission from ref. 56. Copyright © Royal Society of Chemistry.

The introduction of the thiourea functionalities did not affect the nanomorphology of **IRMOF-3** as shown in Figure 8, since similar cubic crystals were produced in both cases. This indicates the high similarity of both crystalline structures. In addition, the obtained nano MOF catalyst showed good thermal stability (up to ~344 °C) similar to **IRMOF-3**.

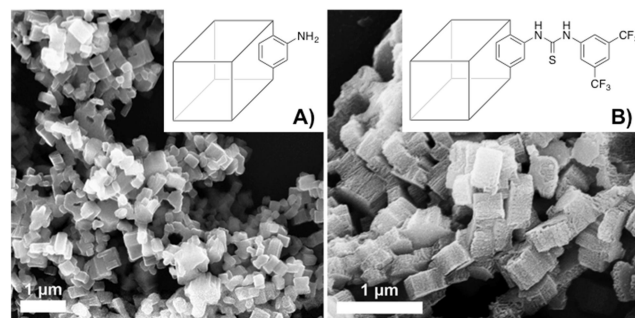
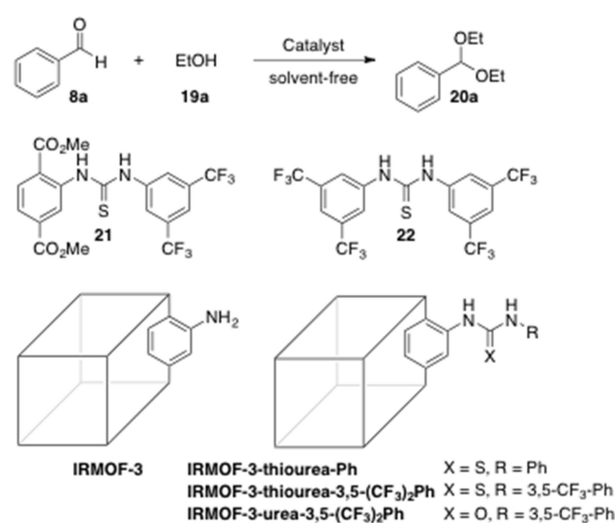


Fig. 8 SEM images of nanoscale A) **IRMOF-3** and B) **IRMOF-3-thiourea-3,5-(CF₃)₂Ph** derivative. Reprinted with permission from ref. 56. Copyright © Royal Society of Chemistry.

An initial screening of the acetalisation reaction of benzaldehyde (**8a**) at room temperature was performed testing all the catalysts synthesised (Table 5).

Table 5 Screening of the acetalisation reaction.^{a,56}



Entry	Catalyst ^b	Loading (mol%)	Yield (%) ^c	TON
1	–	–	0	0
2	<i>p</i> -TsOH	10	99	10
3	thiourea 21	1	81	81
4	thiourea 22	1	89	89
5	IRMOF-3	10	6	0.6
6	IRMOF-3-thiourea-Ph	0.2	15	75
7	IRMOF-3-urea-3,5-(CF₃)₂Ph	0.2	64	320
8	IRMOF-3-thiourea-3,5-(CF₃)₂Ph^d	0.2	96	480
9	Bulk (IRMOF-3-thiourea-3,5-(CF₃)₂Ph^d)	0.2	82	410
10	IRMOF-3-thiourea-3,5-(CF₃)₂Ph^e	0.2	98	490

^a The reaction was performed using benzaldehyde (**8a**) (1.0 mmol), dry ethanol **19a** (4.0 mmol) and organocatalyst (0.2 mol%) at room temperature for 12 h. ^b MOF catalysts are used in nanoscale unless otherwise noted. ^c Determined by GC-MS. ^d 17% thiourea incorporation. ^e 26% thiourea incorporation.

As shown in Table 5, **IRMOF-3** did not work after 12 h (entry 5) and this is in agreement with the fact that the thiourea moiety seems to be responsible of the acetalisation reactivity.⁵⁷ The phenyl substituted **IRMOF-3-thiourea-Ph** catalyst exhibited low reactivity (entry 6) in comparison to the 3,5-bis(trifluoromethyl)phenylthiourea derivative **IRMOF-3-thiourea-3,5-(CF₃)₂Ph**

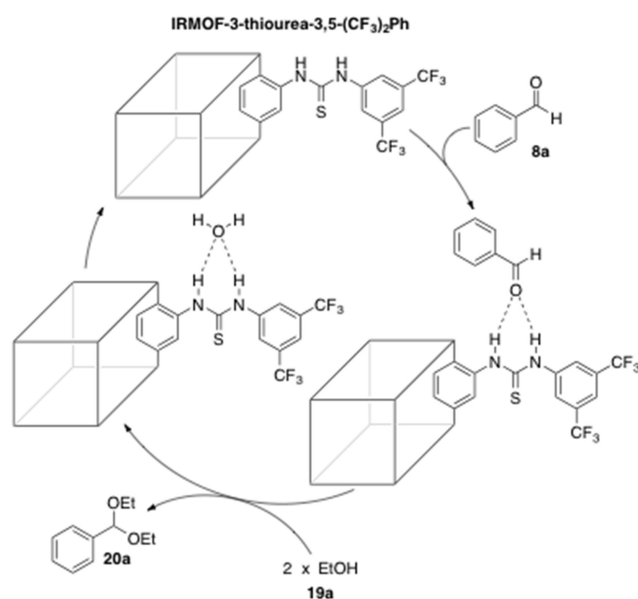
(entry 8). The latter afforded higher reactivity than its urea counterpart (entry 7), and also showed better activity than its bulk partner (regular MOF instead of nano MOF) (entry 9), which suggested a higher utilization rate of the MOF surface and pore. A higher thiourea loading (26% vs 17% thiourea incorporation) on the catalyst led to a comparable yield (98% vs 96%, respectively), with the turnover numbers (TON) up to 490 (vs 480) (entry 10). With the best reaction conditions in hand, **IRMOF-3-thiourea-3,5-(CF₃)₂Ph** (with 26% thiourea incorporation) was applied successfully to a range of substrates for acetalisation and Morita–Baylis–Hillman reactions (Table 6).

Table 6 Scope for the acetalisation and Morita–Baylis–Hillman reactions.⁵⁷

Entry	Substrate	Nu	Product	Yield (%)
1 ^a				92
2 ^a				91
3 ^a				96
4 ^a				< 5
5 ^b				73
6 ^b				81

^a Reaction conditions: For acetalisation (entries 1-4): aldehyde (**8a-d**) (1 mmol), dry alcohol (**19a,b**) (4 mmol) and catalyst (0.2 mol%) at room temperature for 12 h. Yields were determined by GC-MS. ^b For Morita–Baylis–Hillman reaction (entries 5 and 6): aldehyde (**8a,e**) (1 mmol), 2-cyclopenten-1-one (**23**) (2 mmol), 1,4-diazabicyclo[2.2.2] octane (0.5 mmol) and catalyst (2 mol%) were stirred at 4 °C for 24 h. The product was purified by column chromatography on silica gel and the yield was based on the isolated product.

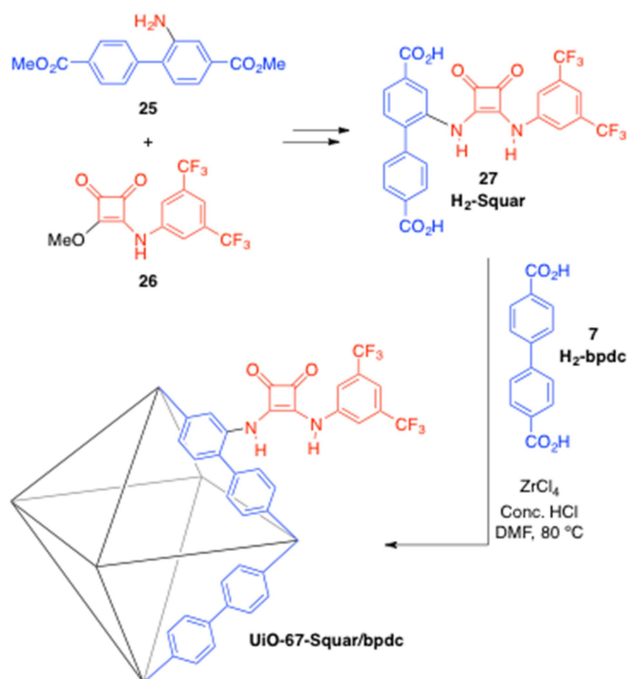
The strong covalent bond between the thiourea moiety and the amino group on **IRMOF-3** ensures the stability of the catalytic functional groups, which maintains over 95% yield after 5 cycles. PXRD and FTIR analyses showed no differences before and after the recycling experiments. The mechanism depicted in Scheme 8 was proposed for the catalytic acetalisation of aldehydes. It starts with the coordination of the catalyst with de carbonyl group. Then, the nucleophilic attack of two alcohols through an ion-pair intermediate, which leads to bezaldehyde acetal and H₂O binding, takes place. And finally, the dehydration of H₂O through ligand exchange regenerates the catalyst. In the case of the M–B–H reaction the nucleophile is compound **23**, in the presence of a base.



Scheme 8 Mechanistic proposal for the acetalisation reaction catalysed by **IRMOF-3-thiourea-3,5-(CF₃)₂Ph**.⁵⁷

Squaramide-containing MOF catalysts

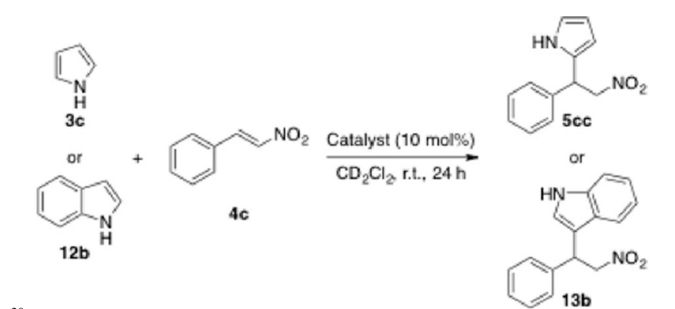
Hupp, Farha, Mirkin and co-workers have recently developed the first example of a MOF bearing acidic squaramide moiety: **UiO-67-Squar/bpdc**.⁵⁸ In the synthesis of this material, the preparation of squaramide-functionalized *strut* (H₂-Squar, **27**) was first achieved from dimethyl 2-aminobiphenyl-4,4'-dicarboxylate (**25**) and 3-((3,5-bis-(trifluoromethyl)phenyl)amino)-4-methoxycyclobut-3-ene-1,2-dione (**26**) (Scheme 9). Then it was incorporated into a zirconium-cluster based **UiO-67** MOF by reaction with the biphenyl chain (H₂-bpdc, **7**) in presence of ZrCl₄, as source of metal. This 1:1 molar ratio mixed-*strut* approach was used to accommodate the relatively sterically demanding squaramide motif and to maintain open pores.



Scheme 9 Synthesis of squaramide-based MOF **UiO-67-Squar/bpdc** and its structure. BET surface areas: $2500 \text{ m}^2 \text{ g}^{-1}$ (**UiO-67**), $1700 \text{ m}^2 \text{ g}^{-1}$ (**UiO-67-Squar/bpdc**).⁵⁸

MOF **UiO-67-Squar/bpdc** was employed successfully as the catalyst for the Friedel–Craft reaction between indole (**12b**) and nitrostyrene (**4c**) (Table 7, entry 6). The introduction of different amounts of squaramide groups into this MOF affected in a great extent the reaction rate. When all the biphenyl groups were substituted with squaramide units (**UiO-67-Squar**), the reactions showed similar outcomes compared to those obtained with the control experiment using only the non-substituted MOF (**UiO-67**) (entries 2 and 5). This lack of catalytic activity may be due to the formation of intermolecular bonds that block reactive squaramide centers and/or the shrinking of the pores of the MOF. However, when they employed **UiO-67-Squar/bpdc** with only 50% of the biphenyl groups linked to squaramide moieties, the results observed were much higher than the control experiment (Table 7, entry 6). The authors attributed this improvement in the reactivity to a decrease in the self-aggregation of squaramides, since any cooperative effect was observed between **UiO-67** and free squaramide (**Me₂-Squar** (**27'**)), the methyl esters analogues of squaramide **27**) (Table 7, entry 4). Interestingly, using low-polarity solvents at higher temperature (50 °C), the corresponding product **5cc** was obtained in nearly complete conversion (entries 7 and 8).

Table 7 Selected screening of the Friedel–Crafts reaction using different catalysts.^{a,58}



Entry	Substrate	Product	Catalyst	Yield (%) ^b
1	12b	13b	–	0
2	12b	13b	UiO-67	22
3	12b	13b	Me₂-Squar (27')	0
4 ^c	12b	13b	UiO-67 + Me₂-Squar (27')	21
5	12b	13b	UiO-67-Squar	22
6	12b	13b	UiO-67-Squar/bpdc	78
7 ^d	12b	13b	UiO-67-Squar/bpdc	93
8 ^e	12b	13b	UiO-67-Squar/bpdc	95
9	3c	5cc	–	0
10	3c	5cc	UiO-67	8
11	3c	5cc	UiO-67-Squar/bpdc	51

^a Reactions performed with **3c** or **12b** (0.03 M), **4c** (0.02 M) and the catalyst (10 mol%) in CD_2Cl_2 at r.t. ^b Determined by ¹H-NMR spectroscopy. ^c 10 mol% MOF and 10 mol% methyl-ester analogous of squaramide **27** (**Me₂-Squar**, **27'**). ^d 1,2-Dichlorobenzene-*d*₄ was used as solvent, at 50 °C. ^e Toluene-*d*₈ was used as solvent, at 50 °C.

In the case of using pyrrole (**3c**) as nucleophile, a lower conversion was achieved, under the same conditions describe for indole (compare entries 2 and 6 with 10 and 11, respectively). Remarkably, this catalyst could be reused through multiples cycles, yet a little decrease in the yield is observed over successive catalytic cycles.

Summary and outlook

In conclusion, the modular synthesis, tunable topology, large surface area and intrinsic porosity of MOFs make them very promising materials for heterogeneous catalysis. The well-defined confined space imposed by MOFs provides a unique functional microenvironment with catalytic sites uniformly arranged along the 3D supramolecular structure. The versatile field of MOF-based heterogeneous catalysis is now offering a new world of possibilities for hydrogen-bonding organocatalysts by overcoming self-aggregation and solvation issues that are usually found in homogeneous conditions. The first examples of urea-, thiourea- and squaramide-containing MOFs, have demonstrated their effectiveness in relevant chemical transformations such as Friedel–Crafts reactions, nitroaldol (Henry) reactions, acetalisation, alcoholysis of epoxides and Morita–Baylis–Hillman reactions. In general, the results derived from these studies support the conclusion that catalysis occurs mainly within the pores of the functionalised MOFs, showing improved catalyst stability and substrate size selectivity in comparison to the analogous homogeneous catalysts. Moreover, the organocatalyst-containing MOFs could be easily separated

and recycled several times without loss of activity and/or major leaching problems.

These materials can be synthesized using either organocatalyst-containing organic *struts* or by post-synthetic modification of suitable functionalised MOFs. Obviously, the enhancement of the thermochemical stability of the hybrids as well as the achievement of larger size cavities, constitute major aspects to consider when engineering new MOF-based organocatalytic systems that are expected to compete with other porous materials such as zeolites and mesoporous silica. However, the development of hydrogen-bonding organocatalyst-containing MOFs is still in its infancy with only the few examples summarized in this contribution. Many other examples are still necessary to demonstrate the reliability and broad use of these materials. In this sense, we anticipate intense research activities focused on asymmetric transformations *via* the introduction of chiral hydrogen-bonding organocatalysts, cascade or tandem reactions through the engineering of multiple catalytic sites and the activation of new reactive pathways that are otherwise unfeasible. The development of new generation of MOF materials with the ability to respond to external stimuli that are common during a given reaction would facilitate the control of the catalytic function as an autonomous on-off unit. Moreover, proper comparative studies in terms of overall costs, reaction times, catalyst loading and TON/TOF numbers with the current homogeneous catalysts used in industry will be necessary to establish the real impact of these new heterogeneous organocatalysts. The report of isolated yields in future works could also provide additional credibility to the potential advantages of these heterogeneous catalysts.

We are convinced that the examples and results highlighted in this contribution will serve as inspiration for the development of superior hydrogen-bonding organocatalysts to accomplish, in confined spaces, chemical transformations that are either slow or unaffordable under homogeneous conditions.^{59,60}

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