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1 Introduction

Although homogeneous Brønsted acids, such as $H₂SO₄$ and HCl, are widely used as cheap catalysts, they are impractical to recycle, and they cause severe equipment corrosion as well as environmental pollution.¹ As a result, heterogeneous Brønsted acid catalysts are highly desirable.² Solid supports, such as organic polymers³ and porous materials,⁴ have been commonly utilized as heterogeneous Brønsted acidic catalyst carriers.⁵ Among several porous materials reported in the literature, metal–organic frameworks (MOFs) demonstrated several advantages in their highly tailorable nature, porous structure and large surface area.⁶ MOF materials have been widely utilized as catalytic materials for a wide variety of organic transformations.⁷ In general, Brønsted acid/MOF systems have been obtained using organic ligand bearing a pre-installed acid functional group.⁸ However, this approach limits the variety of Brønsted acid MOFs that can be prepared and increases the cost of Brønsted acid MOFs.⁹ More synthetic approaches to access Brønsted acid MOFs are therefore necessary for the preparation of low pK_a catalytic materials.¹⁰

Post-synthetic modification (PSM) of porous MOFs is a rapid approach for the synthesis of functionalized MOF scaffolds

Development of a Brønsted acid Al–MIL-53 metal– organic framework catalyst and its application in [4 + 2] cycloadditions†

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Two Al–MIL-53 derived metal–organic frameworks (MOFs) were developed to serve as efficient heterogeneous Brønsted acid catalysts. Sulfonic acid functional groups were successfully incorporated into the framework by post-synthetic modification (PSM) using commercially available reagents. The synthesized Al–MIL-53-RSO₃H and Al–MIL-53-ArSO₃H Brønsted acid–MOF catalysts were fully characterized by SEM, powder XRD, FTIR, TGA and N_2 adsorption/desorption isotherms. An efficient [4 + 2] cycloaddition of 2-vinyl-substituted phenols was evaluated using the newly synthesized Al–MIL-53– RSO₃H and Al–MIL-53–ArSO₃H catalysts. The Al–MIL-53–RSO₃H and Al–MIL-53–ArSO₃H catalysts were found to be compatible with a variety of substituted substrates; finally, they can be recycled five times without compromising the yield or selectivity of the reaction. PAPER
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through stable covalent bonds.¹¹ Brønsted acidic moieties can be introduced easily and rapidly taking advantage of PSM methods.¹² Yaghi and co-workers¹³ developed a PSM procedure that employs 1,3-propanesultone to synthesize IRMOF-3 derived Brønsted acid MOF; Wang and coworkers instead¹⁴ applied a similar PSM strategy to the synthesis of Brønsted acid MOF catalysts. More recently, Cohen and co-workers have developed an anhydride approach for the modification of aminofunctionalized MOFs using the modified MOFs for organocatalytic reactions.¹⁵ For the installation of a highly acidic sulfonic group, the choice of the PSM reagent is highly important. For example, the use of chlorosulfonic acid as sulfonating agent could lead to the complete collapse of the crystalline structure.¹⁶ For this reason, a commercially available reagent for the introduction of a sulfonyl group under mild reaction condition would be highly desirable.¹⁷

Hetero-Diels–Alder (HDA) reactions are among the most useful organic transformations in heterocyclic synthesis as they can afford a broad range of structural complexity.¹⁸ ortho-Quinone methide $(o\text{-}QM)^{19}$ is composed of a cyclohexadiene motif in conjugation with a carbonyl and a methylene group. ortho-Quinone methide intermediates can undergo 1,4-conjugate addition reactions with nucleophiles and $[4 + 2]$ cycloaddition reactions, including HDA reaction, with various dienophiles.²⁰ Chromanes have been prepared by HDA reactions with an *o*-quinone methide used as dienophile.²¹ However, only very limited examples of such reactivity²² have been reported applied to the synthesis of 4H-chromene derivatives, which are useful in medicinal chemistry.²³ An unreactive alkyne is generally required to access 4H-chromenes via an HDA reaction. Alternatively, a tandem $\begin{bmatrix} 4 & + & 2 \end{bmatrix}$ /elimination reaction sequence may result in the construction of 4H-chromenes from

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the corresponding 2π partners. The elimination of a stable molecule, such as a benzylic substituent, is the common method for o-QM generation. It has been proposed that the mild protonation of 2-ethynylphenol might result in a benzylic carbocation intermediate. The generated benzylic carbocation would subsequently tautomerize to the desired o-QM intermediate.

In this work, we wish to report two Al–MIL-53 derived Brønsted acid MOFs for $[4 + 2]$ cycloadditions/elimination catalysis tandem reaction.²⁴ The newly synthesized Al-MIL-53- $RSO₃H$ and Al–MIL-53–ArSO₃H MOF catalysts bear sulfonic acid groups, and were both obtained from commercially available reagent. A variety of substituted 4H-chromenes was generated from different 2-vinyl-substituted phenols. Furthermore, the synthesized Al–MIL-53–RSO₃H and Al–MIL-53–ArSO₃H catalysts can be readily filtered and separated from the reaction solution, allowing five-time recycle and reuse of the catalyst without compromising the yield and selectivity of the reaction. BSC Advances Workerstonding 2r: partners, the elimination of a stable \overline{S} Results and discussion moted on the more and the properties are the published in the more and the published under a case of the more and the p

2 Experimental section

2.1 Preparations of MOF support

Al–MIL-53–NH2 was prepared from 2-aminoterephthalic acid, per literature procedure.²⁵

2.2 Preparations of Al–MIL-53– $RSO₃H$

1.85 g of Al–MIL-53–NH₂ (0.85 mmol based on a MW of 2181.28 ${\rm g}$ mol $^{-1}$) was suspended in 20 mL of CHCl₃; then, 2.0 equiv. of 1,3-propanesultone (207 mg, 1.7 mmol) were added. The mixture was stirred slowly at 25 \degree C for 12 h, after which the solvent was decanted. Fresh DMF (10 mL) was added once a day for three days; $CHCl₃$ was then used to rinse the crystals once a day for two days. The crystals were dried under vacuum at 40 °C before use.

2.3 Preparations of Al–MIL-53–ArSO₃H

1.85 g of Al–MIL-53–NH₂ (0.85 mmol based on a MW of 2181.28 ${\rm g}$ mol $^{-1})$ were suspended in 20 mL of CHCl $_3$; 2.0 equiv. of o-sulfobenzoic acid anhydride (313 mg, 1.7 mmol) were then added. The mixture was stirred slowly at 25 \degree C for 12 h, after which the solvent was decanted. Fresh DMF (10 mL) was added once a day for three days; $CHCl₃$ was then used to rinse the crystals once a day for two days. The crystals were dried under vacuum at 40 \degree C before use.

2.4 $[4 + 2]$ cycloaddition reaction conditions

In a general catalytic reaction, 1 mol% of the Al–MIL-53–NH₂ derived MOF acid was added to 1.00 mL of chlorobenzene in an oven-dried round-bottom reaction vessel. 2.0 mmol of organic substrate was added then. The solution was stirred at 40 $^{\circ}$ C for 3 h. The reaction mixture was passed through a silica gel plug and eluted with 5 mL of dichloromethane. Then, the filtrate was concentrated under reduced pressure and the residue was purified by flash chromatography over silica gel to afford the dimeric product. The yield was calculated based on the isolated product.

3 Results and discussion

Al–MIL-53–NH2 was synthesized using 2-amino-1,4 benzenedicarboxylate ($NH₂-H₂BDC$) as the organic linker.

Subsequently, Al–MIL-53–RSO₃H and Al–MIL-53–ArSO₃H were modified post-synthetically using the corresponding reagent in chloroform solvent (Scheme 1).

The Al–MIL-53–NH₂ crystals appeared to be uniform nanorods 35 nm wide and 210 nm long (Fig. 1a). The synthesis of the Al-MIL-53-RSO₃H catalyst was achieved by treating Al-MIL-53- $NH₂$ crystals with 1,3-propanesultone in chloroform solution (Scheme 1). As evidenced by scanning electron microscopy, the structural morphology was maintained during the postsynthetic modification of the Al-MIL-53-NH₂ crystals (Fig. 1b). Similarly, the structural morphology of Al–MIL-53-ArSO₃H was also maintained (Fig. 1c).

Powder X-ray diffraction (PXRD) studies were performed to confirm the successful synthesis of Al-MIL-53-NH₂ MOF and its sulfonated derivative.¹⁷ Post-synthetic modification with 1,3propanesultone did not cause the decomposition of the Al–MIL-53–NH₂ crystalline structure, underlining the stability of the Al– MIL-53–NH2 derived Brønsted acid (Fig. 2b). Moreover, treatment with o-sulfobenzoic acid anhydride also maintained the crystalline structure of Al–MIL-53–NH₂ (Fig. 2c).

The modification ratio of aromatic sulfonic moiety on Al-MIL-53-RSO₃H and Al-MIL-53-ArSO₃H was measured by ¹H NMR, by comparing the spectra of the modified materials with that of Al–MIL-53–NH₂ (Fig. 3). Al–MIL-53–NH₂ digested in the presence of HF clearly showed three peaks. The singlet proton represents the aromatic proton adjacent to the amino functional group. The two doublets instead belong to the protons in the para and meta positions with respect to the amino group. Proton nuclear magnetic resonance studies of the digested

Scheme 1 Schematic illustration of the post-synthetic modification of $Al-MIL-53-NH₂$

Fig. 1 SEM of (a) $Al-MIL-53-NH_2$, (b) $Al-MIL-53-RSO_3H$, (c) $Al-MIL-53-H_2$ $53-ArSO₃H$ and (d) recycled Al-MIL-53-ArSO₃H

Fig. 2 Powder XRD of (a) Al–MIL-53–NH₂, (b) Al–MIL-53–RSO₃H and (c) $Al-MIL-53-ArSO₃H$.

sample indicated that 20% of Al–MIL-53–RSO₃H amino groups were post-synthetically functionalized (Fig. 3b). This observation was similar to results previously reported in literature; similarly, partial $NH₂$ group conversion has also been reported.²⁶ In a similar fashion, Al–MIL-53–ArSO₃H showed a modification ratio of 21% (Fig. 3c). These results clearly demonstrated the successful post-synthetic modification of amino functional groups, as well as the new installation of new acidic moieties. Additional evidence for sulfonic acid modification was obtained via FT-IR spectra of Al–MIL-53–NH₂ and Al– MIL-53–RSO₃H (Fig. S2[†]). The 1254 cm⁻¹ peak corresponding to the free amine group was significantly reduced, which suggests the conversion of free amino groups to alkyl sulfonic groups (Fig. S2b and c†). However, FT-IR provided limited information in terms of detailed structure, since the peak of ${SO_3}^-$ is not significant on FT-IR spectra.

The specific surface areas of the products were analyzed by $N₂$ adsorption–desorption measurements at 77 K. As shown in Fig. 4, the trace of the Al–MIL-53–NH₂ was a type-I isotherm. Al–

Fig. $3⁻¹H NMR spectra of (a) digested Al-MIL-53-NH₂, (b) digested$ Al-MIL-53-RSO₃H and (c) digested Al-MIL-53-ArSO₃H.

Fig. 4 Nitrogen adsorption/desorption isotherms of (a) Al–MIL-53– $NH₂$, (b) Al–MIL-53–RSO₃H and (c) Al–MIL-53–ArSO₃H.

MIL-53-NH₂ is a highly porous MOF material with a BET surface area of 540 m^2 g^{-1} , calculated by collected nitrogen adsorption isotherms (Fig. 4a). The surface area was reduced to 351 m² g⁻¹ for $AI-MIL-53-RSO₃H$ upon post-synthetic modification, presumably due to the pore structure filling by aromatic sulfonic groups (Fig. 4b). Similarly, the BET surface area for Al– MIL-53-ArSO₃H was calculated to be 312 m² g⁻¹ (Fig. 4c). However, both Al-MIL-53-RSO₃H and Al-MIL-53-ArSO₃H with reduced BET surface area performed well in our catalysis study, likely because the highly porous structure and the flexibility of the newly installed organic functional group allows the free entry and exit of the reaction substrate. Such a reduction in surface area was also observed in several post-synthetic modi fications of MOFs reported in literature.²⁷

Additionally, thermal and structural stabilities of the parent Al–MIL-53–NH₂, as well as of the modified Al–MIL-53–RSO₃H and Al-MIL-53-ArSO₃H catalysts were examined by thermal gravimetric analysis (TGA). The weight loss of Al–MIL-53 began around 200 °C, proving its high thermal stability during organic catalysis. The weight loss of the modified Al–MIL-53–RSO₃H and Al-MIL-53-ArSO₃H sample began around almost identical temperatures (Fig. 5). The TGA results proved the high thermal stability of the Al-MIL-53-RSO₃H and Al-MIL-53-ArSO₃H materials, which ensures their stability in the reaction as a heterogeneous sulfonic acid catalyst. It is important to note that the stability of the material is high enough to prevent the \cos of $SO₃$ and other sulfur anhydrides, even in relatively harsh conditions.

Compound 1a was prepared following a previously reported literature procedure; the o-quinone methide intermediate was obtained through tautomerization of the benzylic carbocation.²⁸ As reported in Table 1, cycloaddition reactions were run in the absence of catalyst (Table 1, entry 1) and with the basic Al–MIL-53–NH2 (Table 1, entry 2) as control reactions, affording no yield of 2a. Homogeneous acids were evaluated for the synthesis of 4H-chromene 2a (Table 1, entries 3–5) also affording no or poor yields and selectivity. Furthermore, hydrochloric acid and acetic acid only showed minimal reactivity at 40 $^{\circ}$ C after 3 h, presumably because of the low acidity in terms of their pK_a (Table 1, entries 3–4). Only strong Brønsted acid p-TsOH functioned as an efficient catalyst in the $[4 + 2]$ cycloaddition reaction, which gave 4H-chromene 2a a yield of 56% (Table 1, entry 5). The unsatisfactory yield was due to the incompletion of the chromene formation reaction pathway, which left significant amount of reaction intermediate 3. The turnover-number (TON) was calculated based on the total conversion of 4H-chromene 2a. 2a was then subjected to the reaction in presence of 1 mol% Al-MIL-53-RSO₃H, yielding 89% of product (Table 1, entry 6). Remarkably, Al-MIL-53-ArSO₃H gave the desired $4H$ -chromene product 2a in almost quantitative yield after 3 h in toluene (Table 1, entry 7). The TON of Al–MIL-53–ArSO₃H under the optimized reaction conditions was calculated to be 98. This **BSC Advances**

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Fig. 5 TGA of (a) $Al-MIL-53-NH₂$, (b) $Al-MIL-53-RSO₃H$ and (c) $Al MIL-53-ArSO₃H.$

Table 1 HDA reaction of 2-ethenylphenol with different catalysts⁴

| Entry | Catalyst | Solvent | Yield ^{<i>a</i>} of 2a | $Yield^b$ of 3 | Conv. of 1 |
|-------|-------------------------------|---------------------------------|------------------------------------|-------------------|---------------|
| 1 | | PhCH ₃ | 0% | 0% | |
| 2 | $Al-MIL-53-NH2$ | PhCH ₃ | 0% | 0% | |
| 3 | HCl | PhCH ₃ | $< 5\%$ | $< 5\%$ | |
| 4 | ACOH | PhCH ₃ | $< 5\%$ | $< 5\%$ | |
| 5 | p -TsOH | PhCH ₃ | 56% | 42% | 98% |
| 6 | $Al-MIL-53-RSO3H$ | PhCH ₃ | 85% | 11% | 96% |
| 7 | Al-MIL-53-ArSO ₃ H | PhCH ₃ | 98% | $< 1\%$ | 99% |
| 8 | $Al-MIL-53-ArSO3H$ | CH ₂ Cl ₂ | 92% | $< 1\%$ | 93% |
| 9 | Al-MIL-53-ArSO ₃ H | THF | 47% | 9% | 56% |
| 10 | $Al-MIL-53-ArSO3H$ | CH ₃ CN | 90% | $< 1\%$ | 91% |
| 11 | Al-MIL-53-ArSO ₃ H | EtOH | 36% | 6% | 42% |
| | | | | | |

Reaction conditions: $1a$ (2.0 mmol), 1 mol% catalyst based on acid group, solvent (5 mL) at 40 $^{\circ}$ C for 3 h. b Isolated yield.

observation strongly demonstrated the unique property of the Al–MIL-53 derived Brønsted acid catalyst. Only the Al–MIL-53 derived acid catalyst achieved the desired cycloaddition product in high yield and high selectivity. Further screening indicated that aprotic solvents (e.g. PhCH₃, CH₂Cl₂, THF, was calculated based on the amount of isolated product $CH₃CN$ appeared to be more compatible (Table 1, entries 7–10), while protic solvents such as ethanol only gave very poor yields of the desired product 2a (Table 1, entry 11). The optimized reaction condition was set to use 1 mol% solid acid catalyst. Chapman reported a homodimerization of carpanone and only 46% yield was achieved using Pd^{2+} catalyst.²⁹ Wan has developed a photo initiated dimerization of phenol type compound and 88% yield was achieved after 3 h.³⁰ Schaus also reported an iron-catalyst dimerization of 2H-chromenes; 85% yield and 3 : 1 diastereoselectivity were achieved after 12 h.³¹ Our strategy effectively assemble two oQM precursor, furnished a dimeric structure in high yield and selectivity, while the catalyst loading was low. Our methodology demonstrated high industrial practical use in terms of low catalyst usage, high yield and high selectivity, when compared with literature reports.

Under the optimized conditions, a broad range of 1,1 diphenylethylenes (1a–1h) was tested. The reaction showed good tolerance toward electron-rich and electron neutral functional groups. A wide range of functional groups, such as methoxy, methyl, and dioxo, performed well (Table 2, entries 1– 6). The non-substituted ethylene (1b) and methyl substituted ethylene (1c) also afforded the desired products 2b and 2c in good yield (Table 2, entries 2–3). meta-Substitution patterns were also tested using methoxy groups, which provided the desired product in good yield (2d, 85% yield). In addition, very electron-rich aromatic rings, such 3,4-benzodioxole also

Table 2 Substrate scope of the $AI-MIL-53-ArSO₃H$ catalyzed cycloaddition reaction⁴

 a 2.0 mmol of substrate 1 and 0.02 mmol of Al–MIL-53-ArSO₃H were dissolved in 2.0 mL of toluene and stirred at 40 $^{\circ}$ C for 3 h. The yield was calculated based on the amount of isolated product.

afforded good yields (2e, 76% yield). Generally, more electronrich functional groups provide increased yields in only 3 hours. The decreasing yield of 2e is likely due to the instability of the methylenedioxy moiety under acidic condition (Table 1, entry 5). Highly electron-rich substrate 1f provided almost quantitative yield of cycloaddition product 2f (Table 1, entry 6). Furthermore, an electron-deficient aromatic group also worked well (Table 2, entry 7). A sterically more hindering substrate was also tested affording a compromised yield 2h (Table 2, entry 8).

A deuterium exchange experiment was performed to further study the quinone methide mechanism (Scheme 2). Deuterated phenol 4 was synthesized and subjected to 60 \degree C thermal condition. However, there was no deuterium migration to the vinyl position in 7. This observation allows us to exclude the possibility of a 1,5-hydride shift mechanism for the generation of the quinone methide species.

In order to examine its recyclability, the Al-MIL-53-ArSO₃H catalyst was isolated and reused in five reaction cycles. The catalyst was centrifuged from the reaction solution and washed with chloroform. The strong covalent bond between the aromatic sulfonic acid moiety and amino group on Al–MIL-53– ArSO3H ensures the chemical stability of the active sites, which remains good, affording up to 98% yield after five reuses of the same batch of catalyst (Fig. 6). Furthermore, the supernatant liquid of the toluene suspension showed no catalytic reactivity toward the vinyl phenol substrate, which is evidence for no leakage of Al-MIL-53-ArSO₃H catalyst. The SEM spectra and Xray powder diffraction pattern of the Al-MIL-53-ArSO₃H catalyst were characterized after five reuses.

These data were indistinguishable from those collected from a batch of fresh catalyst (Fig. 1d and S1†). The catalytic reaction solution was analyzed by ICP-AES and showed no detectable amounts of alumina ions. These facts clearly demonstrate that there was no catalyst decomposition or leakage over the cycloaddition process.

Fig. 6 The catalyst recycling of Al–MIL-53–ArSO₃H in cycloaddition reaction of 2a

Scheme 3 Proposed mechanism for the $AI-MIL-53-ArSO₃H$ promoted cycloaddition reaction.

On the basis of the aforementioned experimental results and mechanistic findings, a possible mechanism is proposed in Scheme 2. 1,1-Diphenylethylene 1a equilibriums to form quinone methide 4 (Scheme 3). Then quinone methide 4 reacts with another equivalent of 1a through a concerted hetero-Diels– Alder reaction, to afford dimeric compound 3. Driven by steric effects, the fully substituted C-2 eliminates to generate the desired product 2a under acidic catalytic condition. The conversion of 3 to 2a was also tested using the synthesized Al– $MIL-53-ArSO₃H$ and quantitative yield was obtained for the aromatic group elimination.

4 Conclusions

In conclusion, heterogeneous Al–MIL-53–RSO₃H and Al–MIL- $53-Arg₃H$ catalysts bearing an aromatic sulfonic acid group were synthesized. The structural morphology of both Al–MIL-53– $RSO₃H$ and Al–MIL-53–Ar $SO₃H$ were well retained after postsynthetic modification using commercial available reagents. Al-MIL-53-RSO₃H and Al-MIL-53-ArSO₃H showed good to high activity and selectivity in the $[4+2]$ cycloaddition at only 1 mol% catalyst loadings. Our methodology demonstrated higher reaction efficiency and higher selectivity than other literature reported dimerization system, while only low catalyst loading was utilized. Al-MIL-53-ArSO₃H demonstrated superior catalytic activity over its non-aromatic partner Al–MIL-53–RSO₃H. The Al-MIL-53-ArSO₃H catalyst was chemically stable, possibly due to its strong covalent bonds and it did not show any leaching during cycloaddition catalysis. Further studies involving new applications of the Al–MIL-53 derived catalysts are in progress.

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