En Route to a Heterogeneous Catalytic Direct Peptide Bond Formation by Zr-based Metal-Organic Framework Catalysts

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TOC



Abstract

Peptide bond formation is a challenging, environmentally and economically demanding transformation. Catalysis is key to circumvent current bottlenecks. To date, many homogeneous catalysts able to provide synthetically useful methods have been developed, while heterogeneous catalysts remain largely restricted to the studies addressing the prebiotic formation of peptides. Here, the catalytic activity of Zr_6 -based metal-organic frameworks (Zr-MOFs) towards the peptide bond formation is investigated using the dipeptide cyclization as a model reaction. Unlike previous catalysts, Zr-MOFs largely tolerate water, and reactions are carried out under ambient conditions. Notably, the catalyst is recyclable and no additives to activate COOH group are necessary, which are common limitations of previous methods. In addition, broad reaction scope tolerates substrates with bulky and Lewis basic groups. The reaction mechanism was assessed by detailed mechanistic and computational studies, and features a Lewis acid activation of carboxylate group by Zr center towards amine addition, in which an alkoxy ligand on adjacent Zr site assists lowering the barrier of key proton transfers. The proposed concepts were also used to study the formation of

intermolecular peptide bond formation. While intrinsic challenges associated with the catalyst structure and water removal limit a more general intermolecular reaction scope under current conditions, the results suggest that further design of Zr-MOF catalysts could render these materials broadly useful as heterogeneous catalysts for this challenging transformation.

KEYWORDS: Metal organic frameworks, synthesis, zirconium, amide bond, peptide bond, diketopiperazines, peptide synthesis, peptide bond formation.

Introduction

The synthesis of amides is a frequent task in every chemical laboratory. However, common procedures suitable to small laboratory scale fail to meet demanding industrial requirements.¹⁻³ Likewise, peptide bond formation is even more challenging due to the lower reactivity of the amino acid NH₂ group compared to regular amines.⁴ In these reactions, carboxylic acids usually require additional steps and/or reagents to facilitate NH₂ attack, substantially decreasing process efficiency by increasing waste generation and energy consumption. Catalysis has been explored as an alternative to streamline amide synthesis, resulting in amide bond formations directly from non-activated carboxylic acids and amines catalyzed mainly by boron⁵ and group IV/V metal⁶⁻⁹ catalysts.¹⁰ Notably, catalytic peptide bond formation remains underdeveloped, despite advancements featuring boron-,¹¹⁻¹³ metal-¹⁴⁻¹⁵ and organocatalysts¹⁶ reported recently. Although promising, these catalytic systems still require water removal from the reaction to provide good yields, making their re-use practically and economically challenging for wider industrial applications.^{1,4} Recently, we have shown that embedding Zr(IV) or Hf(IV) cations into anionic oxo clusters imparts an unusual water tolerance to the catalytic reaction, allowing formation of amide bonds under mild conditions with great experimental simplicity, and potential re-use of the catalyst.¹⁷⁻¹⁸ Here, we extend this concept by using Zr(IV)-oxo clusters embedded in metal organic frameworks (MOFs) as convenient and tailorable heterogeneous catalysts for the peptide bond formation.

The synthesis of amides directly from non-activated carboxylic acids and amines using heterogeneous catalysts is still of limited utility to synthetic chemists, despite numerous studies that have been done in the past.¹⁰ Most reactions are limited to formylation and acylation reactions, with few examples reporting more general substrates.¹⁹⁻²² Generally, these heterogenous catalysts afford amides under similar conditions to those used by homogeneous catalysts, that is, under (azeotropic) reflux at temperatures above 100 °C in anhydrous conditions. In addition, many of the heterogeneous catalysts reported so far consisted of commercially available inorganic oxides like Nb₂O₅, Al₂O₃, Zr₂O, TiO₂, SiO₂, etc, precluding control over the molecular environment surrounding the catalytic sites.^{19, 23-25} Attempts to rationally design inorganic matrixes doped with catalytically active metals have been done,²² but despite recent advances in the areas of surface organometallic and single-atom catalysis,²⁶⁻²⁷ tuning and designing heterogeneous catalysts at the molecular level remains a very challenging task, which limits to some extent the substrate scope expansion through a more rational design of novel catalysts. On the other hand, metal-organic frameworks (MOFs) have emerged as a powerful new class of heterogeneous catalysts.²⁸⁻³² Strikingly different from other common heterogeneous materials, MOFs are intrinsically porous and highly tailorable organic-inorganic hybrid materials prepared through the controlled assembly of distinct metal clusters and specially designed

organic linkers.³³ Moreover, an increasing number of methods allow engineering defects to fine-tune desired catalytic properties.³⁴⁻³⁶ However, the use of MOFs in the amide bond formation remains elusive,^{2, 37} despite the successful reports of MOFs as catalysts in esterification reactions.³⁸⁻⁴² So far only a single MOF-catalyzed amide bond formation has been reported focusing on the utility of a newly synthesized MOF material using previously reported conditions; however, this study provided little mechanistic insights that could allow further development of such chemistry.⁴³

Recently, our work intersected with MOF catalysis due to our interest in developing heterogeneous artificial peptidases based on metal-oxo clusters.⁴⁴⁻⁴⁵ To this end, we have pioneered the peptidase activity of Zr₆O₈cluster-centered MOFs (Zr-MOFs) using MOF-808,⁴⁶⁻⁴⁷ NU-1000,⁴⁸ and UiO-66⁴⁹ as catalysts. Among other practical advantages, these Zr-MOFs provided much higher hydrolysis rates than our previous Zr(IV)/Hf(IV)-polyoxometalate catalysts, prompting us to further evaluate if Zr-MOFs could also work in the reverse direction and lead to the formation of peptide bonds instead of its cleavage, as we observed previously for Zr(IV)/Hf(IV)-polyoxometalate complexes.¹⁷ Prior works addressing the formation of peptide bonds through heterogeneous catalysis have been mostly related to the prebiotic formation of peptides.⁵⁰⁻⁵⁴ They provided extensive mechanistic understanding on the interaction between amino acids and peptides with oxide surfaces, particularly silica, and the formation of peptide bonds in these environments.⁵⁵⁻⁵⁷ The formation of peptide products has been generally observed at temperatures which exceeded 100 °C. Moreover, products were frequently generated as mixtures, and were usually not isolated and purified, thereby drastically limiting the synthetic potential of these reactions for preparative organic chemistry.⁵⁸ Therefore, in view of the great structural versatility and stability,⁵⁹⁻⁶⁰ the increasing utility of Zr-MOFs in catalysis,⁶¹ and the lack of more general heterogeneous catalysts for the catalytic direct formation of amide and peptide products,^{10, 19-22, 43} repurpose of Zr-MOFs from hydrolysis to formation of peptide bonds would greatly contribute to develop a truly sustainable and practical peptide bond formation method. In this context, we report the formation of peptide bonds catalyzed by UiO-66 MOF. Inspired by our previous work,^{17, 46-49} we used the dipeptide cyclization to develop and explore the catalytic activity of Zr-MOFs towards intramolecular peptide bond formation. Furthermore, we explored the reaction mechanism by theoretical and experimental approaches, which unveiled key advantages of the multimetallic nature of the Zr-oxo cluster to the reactivity observed. Initial results towards the intermolecular peptide bond formation reaction are also presented.

Results and discussion

Reaction optimization. To develop Zr-MOFs as catalysts for the formation of peptide bonds, we have started our work by studying the cyclization of dipeptides in the presence of Zr-MOFs, following our previous successful strategy in which the hydrolytic activity of Zr(IV)/Hf(IV)-polyoxometalate complexes was repurposed to an amide bond formation reaction upon rational adjustment of the reaction conditions.¹⁷⁻ ¹⁸ Thus, the cyclization of glycylglycine (**1a**) to 2,5-diketopiperazine (**2a**) was evaluated in various solvents upon incubation of 1a with different Zr-MOFs, which were previously shown to be catalytically active for the peptide bond hydrolysis (Table 1).^{46, 48-49} Based on our previous work,¹⁷ we used DMSO as a representative organic solvent to test three well-known and easily prepared Zr-MOFs with varying characteristics: 1) MOF-808, a 6-connected MOF with 1,3,5-benzenetricarboxylate linkers;⁶² 2) NU-1000, a 8-connected MOF with 1,3,6,8-(p-benzoate)pyrene linkers;⁶³ and 3) UiO-66, a nominally 12-connected MOF with 1,4-benzenedicarboxylate (BDC) linkers.⁶⁴ The presence of free carboxylic acid groups in our substrate could result in adsorption of substrates or the resulting products to the MOF catalyst. Therefore, we have employed a 'washing step' after the reaction to ensure full material recovery. Using MOF-808 as a representative structure, we observed that stirring the crude reaction mixture for 1 h with D₂O generally ensured a high recovery of material (> 95%). We have standardized such step through all the reaction discovery and optimization stages (Table S1).

In these initial reactions using DMSO as solvent, MOF-808 and UiO-66 provided **2a** in higher yields than NU-1000 (Figures S4-S5), likely due to the lower recovery of material observed for NU-1000 (< 20%). MOF-808 formed **2a** in ~50% yield, but a small fraction of glycine (Gly, **3a**) was also detected probably arising from peptide bond hydrolysis with residual water present in the MOF structure and/or solvent (Table 1).⁴⁶ On the other hand, despite the lower yield, UiO-66 cleanly converted **1a** into **2a** (33% yield). Considering that MOF-808 and UiO-66 MOF allowed for an easier recovery of substrate and products in our system, further investigation focused on these MOFs only.

Table 1: Cyclization of GlyX (**1a,b**) to cyclo(GlyX) (**2a,b**) is affected by the MOF structure, solvent, temperature and catalyst loading.



						Yield $(\mathscr{M})^{a}$		
Entry	R	MOF	mol%	Solvent	T (° C)	1a,b	2a,b	3a,b
1	Н	MOF-808	10	DMSO	80	50	48	2
2	Н	UiO-66	10	DMSO	80	65	33	0
3	Н	MOF-808	10	MeOH	80	29	59	4
4	Н	UiO-66	10	MeOH	80	0	100	0
5	Me	UiO-66	10	MeOH	80	0	100	0
6	Me	UiO-66	10	EtOH	80	0	100	0
7	Me	UiO-66	10	ⁿ PrOH	80	0	100	0
8	Me	UiO-66	10	ⁱ PrOH	80	n.a. ^e	58	0
9	Me	UiO-66	10	MeOH	70	8	92	0
10	Me	UiO-66	10	MeOH	60	35	65	0
11^{b}	Н	UiO-66	10	MeOH	80	0	100	0
12^{c}	Н	UiO-66	10	MeOH	80	6	94	0
13 ^d	Me	UiO-66	2	MeOH	80	0	100	0

Conditions: 0.100 mmol **1a,b**, 10.0 mol% MOF, solvent (0.10 mol L⁻¹), 24 h. ^{*a*} Based on ¹H-NMR. ^{*b*} 0.05 mol L⁻¹. ^{*c*} 0.20 mol L⁻¹. ^{*d*} 168 h. ^{*e*} ⁻¹H NMR peaks of **1b** overlapped with solvent.

Following the initial screening, other solvents were probed using MOF-808 and UiO-66, identifying alcohols as optimal reaction solvents. Generally, MOF-808 adsorbed substrate and products more strongly than UiO-66, as observed by the mass recovery < 75% for solvents other than MeOH and DMSO for MOF-808. For UiO-66, only MeCN:H₂O, toluene and dioxane reactions resulted in < 90% mass recovery for those ones carried out in organic medium (Figure S4-S5). Moreover, MOF-808 showed again a greater tendency than UiO-66 to hydrolyze **1a**, as evident from the results in solvents containing water. On the other hand,

UiO-66's enhanced selectivity towards 2a was once more observed for the reaction in MeOH, which provided only desired product 2a and no side hydrolysis reaction (entry 4, Table 1). To test other alcohols, we used GlyAla (1b) instead of GlyGly (1a) to avoid overlap between product and solvent peaks in the ¹H NMR. The minimal structural changes do not affect reactivity as both substrates smoothly provide the desired cyclic adduct 2a, b in >99% in MeOH. Using 1b, ethanol and 1-propanol also provided 2b in >99% yield, while isopropanol resulted in only ~60% yield, suggesting smaller and linear alcohols are preferred as reaction solvents (entries 5-8, Table 1), though overlapped signals in the crude ¹H NMR hampered us to establish whether the lower yield in isopropanol derives from lower conversion or simply from the lower mass recovery in this reaction.

In general, the low mass recovery observed in some cases seems to be related to the available uncoordinated Zr sites, and the ability of solvent to reverse the binding of substrates and products to these sites, since lower mass recovery was generally observed for MOF-808 and NU-1000 in comparison with UiO-66. Similarly, lower mass recovery was also observed when cyclization was carried out in solvents that poorly solubilize 1a,b/2a,b (e.g., dioxane and toluene for UiO-66). These trends strongly suggests that the interplay between available Zr sites and the ability of the solvent to reverse substrates and products presumed coordination to these sites plays a key role in the overall process efficiency. When more Zr sites are available, more substrate/product molecules bind to the MOF, and if the solvent is not able to efficiently reverse this trend, a lower mass recovery is observed. Such hypothesis is also consistent with the better, but rather intriguing performance of UiO-66 for the cyclization of 1a,b in comparison with MOF-808 and NU-1000 MOFs, which intrinsically have more uncoordinated Zr sites than UiO-66. Together, these results show that the connectivity and missing-linker defects alone do not guarantee higher reaction yield as commonly indicated in the literature, since the adsorption of substrates and products to the MOF material and the ability of the solvent to reverse it directly affects the overall mass recovery after the reaction, subsequently impacting the reaction yield. Interestingly, use of water as a reaction solvent resulted in the lowest recoveries for MOF-808 and UiO-66, suggesting solubility is likely not the only factor involved in overcoming material adsorption.

Further optimization was performed by probing the effect of temperature, concentration and catalyst loading for the reaction using UiO-66 in methanol. Lower temperature decreased reaction efficiency. At 70 °C, **2b** yield slightly decreased, and it significantly dropped for temperatures of 60 °C or lower (entries 9-10, Table 1 and Table S2). In addition, dilution or concentration of the reaction did not significantly affected efficiency (entries 11-12, Table 1 and Table S3). Finally, lower catalyst loadings also provided **2b** in quantitative yields, although slower rates were observed (Table S4). Using 2 mol% of UiO-66, **2b** was formed in 33%

yield after 24 h. However, a longer reaction time of 168 h provided quantitative yield, indicating the catalyst is not deactivated by side-processes even after considerably long reaction time (entries 13, Table 1). This robust profile is usual for Zr-MOFs, but is unparalleled to other metal catalysts for the amide bond formation, and further highlights the excellent catalytic potential of Zr-MOFs for the peptide bond formation.

Control experiments. Control experiments confirmed the superiority of UiO-66 as a heterogeneous catalyst for the intramolecular peptide bond formation. In the absence of UiO-66, and when using Zr salts or BDC ligand separately, reaction yields were <10% (Table S5). Notably, soluble $Zr(^iPrO)_4$ and a combination of ZrCl₄ and BDC resulted in good yields of **2b**, though lower than with UiO-66. The performance of $Zr(^iPrO)_4$ and $ZrCl_4/BDC$ was attributed to a putative formation of Zr-oxo clusters *in situ*, which also points to the key catalytic role of UiO-66's Zr₆ clusters.⁶⁵⁻⁶⁶ Additionally, control experiments using drying agents provided low yields of **2b**, strongly indicating that the water adsorption ability of MOF is likely not reposnsible for the observed reactivity (Table S5). Finally, absence of reaction upon removal of catalyst, along with UiO-66's stability observed in this work, strongly indicates that the catalysis is heterogeneous in nature (Figure S6).

Stability, recyclability and water-tolerance of UiO-66 catalyst. Additional experiments confirmed the catalyst stability, and prompted us to probe the UiO-66 recyclability and tolerance of catalyst to water. UiO-66's excellent structural stability was confirmed through scanning electron microscopy (SEM) (Figure 1a), powder X-ray diffraction (PXRD), infrared spectroscopy (FT-IR), and thermal gravimetric analysis (TGA) of catalyst samples that were easily recovered by centrifugation from experiments with **1a,b** (Figures S7-S9). Additionally, ICP-OES revealed very little leaching of Zr(IV) ions into solution (only 0.004%) confirming its structural stability and heterogeneous nature of the reaction. Further, MOF digestion revealed ~5% of **1b** and **2b** after the reaction, indicating that minor amounts of the substrate and product remain adsorbed to the catalyst (Figures S11-S12). This suggests non-negligible changes to the pore and surface of UiO-66 after the reaction and simple MeOH washing steps. Thus, recovered UiO-66 was washed with D₂O and MeOH to remove traces of starting material and product before drying and re-using in a new reaction. This recycling protocol effectively afforded **2b** in >97% yield over 5 cycles (Figure 1b and Table S6), while no structural changes in MOF's crystalline network were detected by PXRD analysis (Figure 1c).



Figure 1: UiO-66 is a stable and recyclable catalyst with high tolerance towards water: (a) SEM images of (1) UiO-66 as synthesized and (2) after reaction with 0.100 mmol **1b**, 80 °C, 24 h, MeOH. (b) UiO-66 affords the same yields of **2b** after five reaction cycles. (c) PXRD pattern of UiO-66 as synthesized (black), and after two (red) and five (blue) reaction cycles with 0.100 mmol **1b**, 80 °C, 24 h and solvent exchange with D₂O and MeOH. (d) UiO-66 maintains its high catalytic performance even after addition of large amounts of external water.

UiO-66 catalytic cyclization of **1b** tolerates uniquely large amounts of water, and >95% yield is observed in the presence of nearly 2800 molar excess of water with respect to the catalyst (Figure 1d and Table S7). UiO-66's stability prompted us to check its tolerance to the presence of water in the reaction, as watersensitivity sharply limits large scale catalytic amide bond formations.^{1-2, 4} By adding increasing amounts of water to the standard reaction, >97% yield of **2b** was observed up until 27.8 mmol of D₂O (33% of reaction volume in Figure 1d), which corresponds to a remarkable 2778-fold excess related to the catalyst (Table S7). Further reactions showed that this tolerance is related to the amount of MeOH, and lower yields were observed when MeOH volume was reduced proportionally to the water added in order to keep reaction concentration constant. However, even when water corresponded to 70% of the solvent, a respectable 82% yield of **2b** is still observed. Such results also strongly suggest the minor contribution of water segregating ability of MOF in driving the reaction equilibrium forward, in agreement with control experiments. A maximum water absorption for UiO-66 of 0.55 g H₂O / g MOF has been reported,⁶⁷ which would mean that to adsorb the 70% water volume used in the referred reaction (~0.7 g), a much higher catalytic loading (~1.3 g of MOF, ~800 mol%) than the one used here (17 mg, 10 mol%) would be needed.

UiO-66 advantageous robustness in the presence of water was further confirmed by its better performance compared to other water tolerant Lewis acid metal salts under the same conditions (Table S8).⁶⁸ When external water was added to the reaction catalyzed by Zr(OⁱPr)₄ (20% of reaction volume), the good yield previously observed for 2b in the aforementioned control experiments decreased by ~30%, which is consistent with general sensitivity of Zr(IV) salts to moisture, given their highly favored hydrolysis to form oligomeric catalytically inactive species.^{10, 69} However, PXRD analysis of UiO-66 after reaction conducted in the presence of 50% v/v of water proved the stability of its overall structure under these conditions (Figure S13), demonstrating an intrinsic advantage of the Zr-based MOFs robustness over conventional Zr catalysts based on metal salts or complexes thereof. In addition, other Lewis acids that have been reported to tolerate water have also been tested in the presence and in the absence of water (Table S8). Even for these simple reactions, a general drop of 20 - 30% of yield was observed, with the best performing Sc(OTf)₃ providing 2b in 80% of yield in the presence of 33% volume of water, while UiO-66 affords 2b in 97% of yield under the same conditions. Considering the higher natural abundance of zirconium compared to the other the Lewis acid metal salts tested (Ti excluded), and the recyclability exhibited by UiO-66, these results underline a promising prospect of Zr-MOFs, even when one contrasts the complexity of these materials with simple, commercially available metal salts.

Reaction Scope. Several other substrates were successfully shown to undergo cyclization and could be isolated in good to excellent yields, showcasing that UiO-66 MOF catalyst can be applied to substrates bearing diverse functional groups (Table 2), which is essential for later development of intermolecular peptide bond forming methods catalyzed by Zr-MOFs. Standard substrate **1b** and Gly-Phe (**1c**) afforded product **2b** and **2c** in high purity and excellent isolated yields after a simple centrifugation and evaporation of the supernatant. In addition, indole, imidazole, hydroxyl and thioether groups in **1d,f,g,i,j** were tolerated and the corresponding products **2d,f,g,i,j** were isolated in 68 - >99% yield. The ESI-MS and NMR analysis of reactions using Gly-Lys and Gly-Arg dipeptides suggested that these substrates could also undergo cyclization; however, we could not isolate and characterize the corresponding products with our current protocol. Nevertheless, these results point to an overall good compatibility with Lewis basic functional groups.

Some steric hindrance was also tolerated since Gly-Pro (1e), a secondary amine, and the bulky side chain of Gly-Leu (1h) afforded products 2e and 2h in 92 and >99% respectively. Similar 2b and 2e yields were obtained when glycine was moved to the C-terminal position of the dipeptide substrate, while a significant drop was observed for Ser-Gly (1f) (45%), likely due to a non-productive coordination of the hydroxyl moiety. Even Ala-Ala (1i), a dipeptide without a glycine residue, was cyclized to 2i in 18 and 33% yield after 24 and 96 h, respectively. Unnatural amino acids 11,m were also successfully cyclized, showing the proteogenic nature of the substrate is not required for observing the reactivity.

Our attempts to cyclize longer oligomers such as tri-, tetra-, and pentaglycine, or unnatural amino acids with distinct carbon chain lengths (4, 6 and 7) under the same conditions were unfortunately not successful. For most cases, neither substrates or products were observed after reactions, suggesting that the substrates remained adsorbed to the catalyst, which is consistent with the significant adsorption of longer peptides (e.g., proteins) to metal oxo cluster observed in our previous works.^{44, 48} In general, cyclized products also showed different affinities for the MOF catalyst, including extreme cases like Gly-Asp (**1n**) and Gly-Glu (**1o**) which even leads to partial loss of the crystallinity rather than cyclization (Figure S14). Thus, in some cases additional washings of the catalyst to remove the products were necessary. However, although not fully optimized, our isolation protocol afforded the cyclized products in good to excellent purity, and no liquid-liquid extraction or chromatographic purification was necessary.

Table 2: Intramolecular amide bond formation from dipeptides and other amino acid substrates.



Conditions: **1b-o** (0.100 – 0.500 mmol), 10.0 mol% UiO-66, MeOH or EtOH (0.10 mol L^{-1}), 80 °C, 24 h. Isolated yields. ^{*a*} 96h reaction. ^{*b*} ¹H NMR yields.

Reaction mechanism. To gain a better understanding of the catalyst activity, we carried out a detailed experimental and theoretical investigation of the reaction mechanism. Based on our own work,^{46-49, 70} previous reports on Zr-catalyzed formation of amides directly from non-activated carboxylic acids and amines,^{17, 71} and control experiments reported here, we suggest a general Lewis acid catalytic pathway for this intramolecular amide bond formation in which the dipeptide carboxylate group is activated through

coordination to a Zr open site of the MOF (Figure 2). However, the reasons for the superior reactivity observed in alcoholic solvents were not clear at the beginning since only the solvent influence in the mass recovery discussed above does not explain why the yield of **2a,b** in MeOH or EtOH is much higher when UiO-66 MOF is used as a catalyst in comparison to when MOF-808 is employed, given that >95% of the initial mass is recovered with both catalyst (Figures S4 and S5). Such observation prompted us to further investigate whether the alcoholic solvent plays an additional role in the mechanism. Thus, we investigated whether both a superior proton transfer ability of alcoholic solvents inside the pores of UiO-66⁷² or the formation of an intermediate ester⁷³ could explain the reactivity observed.



Figure 2: Mechanistic possibilities for the Zr-MOF catalyzed intramolecular peptide bond formation

Reaction kinetics. Different kinetic experiments were used to get a clearer picture of the catalyst activity, such as potential inhibition and involvement of intermediates. Substrate consumption directly leads to product formation judging by the mirrored aspect of the kinetic profile of **1b** cyclization, disfavoring a reaction pathway involving a potential intermediate product (Figure 3a). Using a pseudo first-order kinetics model, a rate constant of 1.54×10^{-4} s⁻¹ and a half-life of 1.2 h were obtained at 80 °C (Figure 3b). Upon increasing the temperature from 60 °C to 90 °C, a \approx 17-fold reduction of the half-life from 16 hours at 60 °C to only 0.9 hours at 90 °C was observed (Figure S15a). Fitting these data to the Eyring equation, we estimated $\Delta G^{\ddagger} = 27.2$ kcal mol⁻¹ at 80 °C ($\Delta H^{\ddagger} = 22.9$ kcal mol⁻¹, $\Delta S^{\ddagger} = -12.1$ cal mol⁻¹ K⁻¹) (Figure S15b). Finally, control kinetic experiments in which the MOF is pre-saturated with product **2b** showed no

difference in yield after 3 and 5 h in comparison with the non-inhibited reaction, suggesting no product inhibition happens (Table S10).



Figure 3: a) Cyclization of GlyAla (**1b**, circles) to cyclo(GlyAla) (**2b**, triangles) as a function of time b) First order decay fit of ln[**1b**] as a function of time. *Conditions:* 0.100 mmol **1b**, 10.0 mol% UiO-66, MeOH (0.10 mol L⁻¹), 80 °C. ¹H NMR yields.

Potential involvement of an ester intermediate. Motivated by the superiority of alcoholic solvents in our reactions, previous reports on the UiO-66 MOF catalyzed esterification reactions under similar conditions,⁴⁰ the formation of peptide bonds from ester substrates,⁷⁴⁻⁷⁵ and the facile cyclization of dipeptide esters to diketopiperazine products,⁷³ we have probed the potential involvement of an intermediate ester in the reaction mechanism as depicted in Figure 2. To check whether intermediate formation of an ester and subsequent cyclization to the final products **1a,b** could account for the superior reactivity observed in alcoholic solvents, we used GlyGly-OMe as a reaction substrate. After 3 h, GlyGly-OMe afforded **2a** in 89% yield (Table S11), showing the feasibility of a potential intermediate ester cyclization. When the cyclization was blocked by protection of **1a** N-terminal position with an acetyl group, the ester was formed in ~80% of yield after 24 h, indicating that ester formation may happen in these conditions (Table S12). However, little GlyGly-OMe (~5%) was detected through ¹H NMR and ESI-MS when the reaction of **1a**

was stopped before completion (Figure S16), suggesting that if ester is involved, its cyclization is likely not the rate determining step.

Computational study. Density Functional Theory (DFT) computations were also used to examine the mechanism of Zr-MOF catalyzed dipeptide cyclization, focusing on modelling of the reaction in ethanol solvent, given the identical results for reactions involving EtOH and MeOH. Experimental characterization of UiO-66 structure revealed our system to be similar to previous reports (see SI for details), thus we based our computational model on a previous computational work on UiO-66.⁷⁶⁻⁷⁷ In this model, a representative cluster model of the MOF catalyst was chosen, comprising a single Zr_6 core unit with one BDC linker defect and an ethoxy group as a defect-capping ligand (see SI for full computational details). Glycylglycine (**1a**) in its most stable canonical form in EtOH was used as a model substrate (Table S14).

At first, we compared the relative stability of different complexes formed from 1a and the Zr₆ cluster, suggesting that the most stable form **A** involves 1a coordinating to a Zr center through its carboxylate group in monodentate fashion (Figure 4 and Figure S17). In **A** the carboxylic acid group in 1a is deprotonated and there is a neutral ethanol ligand at an adjacent Zr site. In general, other structures involving 1a monodentate binding to the free Zr site were found to be lower in energy for the deprotonated dipeptide than for its neutral form. In these complexes, the relative energy was found to vary as a function of the nature of the anchoring group within 1a that coordinates to the metal, in the order carboxylate group (most stable) < amide oxygen atom < NH₂ group.

Using structure **A** as a reactant complex, we considered two potential reaction pathways, namely the formation of cyclic product **2a** through a direct cyclization of **1a** (*Path A*) or through the formation of an intermediate ester which cyclizes in situ (*Path B*) (Figure 2). The results suggest that an ester intermediate is not involved since the calculated barrier for its formation lies at ~30 kcal mol⁻¹ above complex **A**, which is *ca*. 6 kcal mol⁻¹ higher than the barrier for the direct cyclization. This is consistent with our experimental observations that the ester is present in small amounts, but can be formed in appreciable amounts if the cyclization is blocked through protection of the dipeptide's NH₂ group. Therefore, reaction most likely proceeds through direct cyclization of **1a** (*Path A*).

Proposed catalytic cycle. Considering the DFT results, which are in line with experiments, a reaction mechanism is proposed in Figure 4 (Figure S18). From complex **A**, sequential peptide bond *trans/cis* isomerization, NH₂ attack, and water release steps take place to give the cyclized product **2a** (Figure 4). The *trans/cis* isomerization transition state (TS) features a pyramidal sp³ geometry of the amide nitrogen atom,⁷⁸⁻⁸⁰ which was computed to lie 23.1 kcal mol⁻¹ above the reactant complex (**A**), and is consistent with

experimental ΔG^{\ddagger} of 20.9 kcal mol⁻¹ for amide isomerization under similar conditions.⁸¹ Moreover, our computed *trans/cis* $\Delta G = 2.8$ kcal mol⁻¹ matches the experimental value of 3.1 kcal mol^{-1.81} Next, *cis* isomerized intermediate **B** undergoes several low barrier conformational changes that allow the NH₂ group to move closer to the carboxylate group in preparation for the rate-determining nucleophilic addition, which generates cyclic tetrahedral intermediate **C**. The TS for nucleophilic addition is predicted to lie 24.5 kcal mol⁻¹ above **A**, in good agreement with the experimentally-derived ΔG^{\ddagger} of 27.2 kcal mol⁻¹ (Figure 2 and Table S9). The 2.6 kcal mol⁻¹ discrepancy is well within the estimated error bar of the protocol used here. In the last step, cyclic intermediate **C** releases water through a low-barrier proton shift from the NH₂ to the OH group of the tetrahedral carbon. This transfer is assisted by the adjacent ethoxy group, whose absence would otherwise require formation of a highly-strained four-membered ring or the entropically disfavored involvement of an extra solvent molecule, resulting in higher energy barriers.⁸² Next, the formed water molecule is released, ultimately resulting in separate products, which are characterized by $\Delta G = -14.2$ kcal mol⁻¹ relative to the separate reactants (see SI for details).



Figure 4: SMD-B3LYP-GD3BJ/BS2 relative free energies (353.15 K, kcal mol⁻¹) for key intermediates and transition states along the computed intramolecular peptide bond formation reaction path (BS2 is a modified def2-TZVP basis set described in SI).

Intermolecular peptide bond formation. The many benefits of Zr-MOF catalysis for peptide bond formation outlined above prompted us to investigate whether *inter*molecular formation of peptide bonds could be achieved, as the MOF structure could favor the formation of this challenging amide bond.⁸³ Accordingly, glycine (3a) and UiO-66 were incubated in MeOH at 80 °C (Table 3). After 24 h, 20% of glycine was quantitatively converted into a 1a:2a mixture (~10:1 ratio) (Table 3, entry 1). Using longer reaction times increased the yield, reaching an encouraging 75% yield of a 1a:2a (1:2.6 ratio) mixture after 168 h (Table 3, entry 3). Increasing the temperature to 100 °C improved the 24 h reaction combined yield to only 50%, and more of the cyclized product was observed (Table 3, entry 4). This sluggish reactivity prompted us to consider the removal of water from the system, as it is well known that the equilibrium can be driven to the peptide bond formation through water scavenging.⁸⁴ However, using dry MeOH did not improve the yield (Table 3, entry 5), and combining dry MeOH with molecular sieves only improved the yield by 13% after 48 h (Table 3, entry 6). The apparent limited effect of molecular sieves in these reactions is puzzling, and contradicts the generally observed trend with many other catalysts reported in the literature.³⁷ This could be related to the ability of Zr-MOFs to adsorb water,⁶⁷ thereby acting both as a desiccant and a catalyst; however, this potential dual effect of the MOF structure in the intermolecular reaction would require further studies in order to be confirmed. Notably, negligible reactivity was observed with Zr(ⁱPrO)₄ under standard conditions (80 °C, no molecular sieves), and only 46% of 1a/2a combined were produced in dry conditions (100 °C, with molecular sieves) after 48 h. Further, no products were observed without MOF in the presence or absence of molecular sieves. Together, these results evidence the beneficial effect of the MOF network in enhancing the reactivity (Table S16).⁷⁵

 Table 3: Intermolecular peptide bond formation.

Р. С.	UiO-66		
H_2N H_2N O	MeOH 80 - 100 ℃	$O \dot{R}$	R R
3a , R = H	00 100 0	1a,b	2a,b
3b. R = Me			

D

Entry	R	T (°C)	Mol. Sieves?	Time (h)	Conversion (%)	Yield 1+2 (%)	1:2
1	Η	80		24	20	20	10:1
2	Н	80		96	39	39	1.2:1
3	Н	80		168	75	75	1:2.6
4	Н	100		24	50	50	1:1.1
5^a	Η	100		24	50	50	1:1.3
6	Н	100	Yes	48	63	63	1:7

7	Me	100	Yes	48	45	0	N.D.
Conditio	ns: 3a ((0.200 mmol),	10.0 mol%	UiO-66, MeOH (0.10 mol L ⁻¹), 80 °C, 1	molecular sieves 3Å	(100 mg).

¹H NMR yields. ^a Dry MeOH.

The counterintuitive selectivity favoring **1** at shorter reaction times motivated us to investigate the mechanism of this reaction by DFT modelling of the key intermediates and transition states along the reaction path, as shown in Figure 4 (see SI for more details). Our calculations estimated the energy barrier for the intermolecular reaction to be ~4 kcal mol⁻¹ higher relative to the '**3a**-free' intramolecular peptide bond formation, which is in qualitative agreement with the longer reaction times and selectivity observed for intermolecular reaction (Table 3 vs. Tables 1 and 2), as well as the non observation of triglycine or longer oligomers. However, the observed **1a**:**2a** ratios obtained at early stages of the reaction together with similar binding energies of glycine and **1a** to UiO-66 suggest that **3a** might inhibit the cyclization (Figure 5). This inhibition was confirmed by adding **3a** to a standard GlyAla (**1b**) cyclization (17% yield vs. 100% yield in absence of **3a**, Table S18).



Figure 5: SMD-B3LYP-GD3BJ/BS2 relative free energies (353.15 K, kcal mol⁻¹) for key intermediates and transition states along the computed intermolecular peptide bond formation reaction path (BS2 is a modified def2-TZVP basis set described in SI).

Attempts to extend the intermolecular reaction to other amino acids was not successful. A dimerization of alanine under the same conditions used for glycine failed to provide desired adducts (Table 3, entry 7). Attempts to optimize reaction concentration, temperature, and the presence or absence of molecular sieves were unfortunately not fruitful. In several cases, part of the initial alanine was not recovered, which was attributed primarily to its adsorption. However, aromatic signals coherent with a para substituted aromatic ring could be detected in the NMR of crude reaction mixtures, suggesting that the MOF linker reacts preferably with the amino acid substrates bulkier than glycine. Though we were not able to identify the product formed, loss of MOF network structure due to esterification of linkers has been reported before.⁸⁵ However, PXRD analysis after the reaction showed the overall structure of MOF remains intact, suggesting that the lack of reactivity could not be due to the MOF decomposition. Further attempts to facilitate the intermolecular peptide coupling led us to react H-Gly-OMe with different amino acids. However, in these reactions only cGG was observed, and dimerization of H-Ala-OMe also did not afford the linear or cyclic product (Table S17).

This unexpected lack of reactivity with other amino acids led us to probe more conventional organic substrates such as phenylacetic acid and benzylamine under the same conditions (Table 4). The *N*-benzyl-2-phenylacetamide **6** product was observed in low yields even in the presence of molecular sieves (entries 1 and 2). Preliminary optimization could elevate the yield up to 88% by exchanging the reaction solvent to dioxane, and using molecular sieves, whose effect in this case is in line with prior reports (entries 3-6).^{6, 86-88} These results show that the extent to which water removal affects the reaction depends on the substrate and/or reaction condition being used, and that even currently available Zr-MOFs might need water removal techniques to provide good yields after all. However, given that **6** was formed, the lack of reactivity observed with amino acids other than glycine suggests that it is rather the catalyst structure, and not the reaction itself or a poor water removal, that is precluding product formation. In this sense, the overall optimization of the Zr-MOF structure would be needed in order to make the catalytic activity more general toward intermolecular reactions.

 Table 4: UiO-66 catalyzes intermolecular amide bond formation.

			-66	O U
Pn (202H [∓] H2Ν	Solvent 5 80 °C, 2	, MS 3Å 14 - 48 h	N Ph H 6
Entry	Solvent	Mol. Sieves?	Time (h)	Yield 6 (%)
1	EtOH	No	24	14
2^a	EtOH	Yes	24	18
3	1,4-dioxane	No	24	39
4	1,4-dioxane	No	71	41
5	1,4-dioxane	Yes	71	88
6 ^{<i>a</i>}	1,4-dioxane	Yes	24	84 (75 ^{<i>b</i>})
$7^{a,c}$	1,4-dioxane	Yes	24	16

Conditions: **4** (0.100 mmol), **5** (0.150 mmol) 10.0 mol% UiO-66, molecular sieves 3\AA (50 mg), solvent (0.10 mol L⁻¹), 80 °C. ¹H NMR yields. ^{*a*} 3 equiv. of **5**. ^{*b*} Isolated yield. ^{*c*} No UiO-66 MOF.

Overall, these explorative results for the direct dimerization of unprotected commercially available amino acids showcases the potential of Zr-MOFs in streamlining the intermolecular catalytic peptide bond formation, although further optimization of the MOF catalyst structure is clearly necessary to expand the reaction scope to a broader range of amino acids.

Conclusion

In conclusion, we discovered the high-potential of Zr-MOFs as a new class of heterogeneous catalysts for peptide bond formation. We have shown that UiO-66 efficiently catalyzes the intramolecular formation of peptide bonds directly from free amines and non-activated carboxylic acids without any additives, drying agents, inert atmosphere or special protocols to remove water from the reaction. Good yields were obtained for a variety of substrates containing Lewis basic functional groups and bulky side chains, including a secondary amine group. Remarkably, UiO-66 is stable under the reaction conditions and shows a water tolerance that largely superseded other Zr(IV) catalysts, resulting in a straightforward recycling of the catalyst and showcasing the potential of Zr-MOFs as viable sustainable and water-tolerant catalysts for the amide bond formation. In addition, our detailed mechanistic investigation uncovered several important features for the design of future catalysts, such as 1 the interplay between MOF structure, reaction solvent and the easy recovery of substrates and products, which might have a significant impact on the overall process efficiency, 2 the key role of an alkoxy group on the adjacent metal center in lowering the barrier

of key proton transfers and *3*) the non-involvement of an ester intermediate, indicating a more general reaction scope is feasible for Zr-MOFs. Finally, we provide proof-of-concept that a similar mechanism also enables the formation of intermolecular peptide bonds directly from unprotected commercially available glycine. Even though further catalyst structure optimization is needed to expand this reactivity to other amino acids precursors, these results underline a high catalytic potential of Zr-MOFs that can impact a broad range of scientific areas.

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Notes The authors declare no competing financial interest.

ASSOCIATED CONTENT

Supporting Information. Experimental procedures, supplementary experiments, copy of NMR spectra (PDF), computational study details and results, 'xyz' coordinates of UiO-66 starting cluster model. This information is available free of charge on the ACS Publications website.

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