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Introduction

Metal–organic frameworks (MOFs) are porous coordination polymers containing potential voids.¹ They are composed of inorganic nodes, referred to as secondary building units (SBUs), linked together by multitopic ligands serving as linkers. Due to the numerous possible combinations of metals or SBUs,² including homometallic or heterometallic clusters,³ with organic linkers of variable geometries and coordinating groups^{4,5} it is not surprising that over 100 000 MOF structures have been published so far.⁶ MOFs provide a broad spectrum of structures and topologies,⁷ leading to tuneable and sometimes predictable properties and functionalities. These advantages, together with their porosity, high specific surface area,^{8,9} and tuneability of their properties,¹⁰ predetermine

Zirconium-based metal-organic frameworks: the relation between linker connectivity, structure stability, and catalytic activity towards organophosphates[†]

HINESE

HEMICAL

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Metal–organic frameworks (MOFs) are studied for many applications, however, there are only a few examples of commercialization. One of the reasons behind this is that the stability of MOFs is still unknown. Much attention has been devoted to the rational synthesis of novel MOFs, yet the predictability of MOF stability is so far limited. The present study compares the stability in a water environment with pH ranging from 3.0 to 11.0 of four zirconium-based MOFs constructed from ditopic, tritopic, and tetratopic linkers, namely UiO-66 (benzene-1,4-dicarboxylic acid), MOF-808 (benzene-1,3,5-tricarboxylic acid), MIP-200 (5,5'-methylenediisophthalic acid), and PCN-222 (5,10,15,20-tetrakis(4-carboxyphenyl)porphyrin). Finally, to delineate the nature of the defects induced by the linker release, we tested the MOFs treated at a certain pH for the degradation of model organic pollutant methyl paraoxon. It is clear that both MOFs based on tetratopic linkers are much more stable than UiO-66 and MOF-808 composed of di- and tritopic linkers, respectively. It should be noted that the kinetics of the linker release were also significantly slower for tetratopic linkers. At the same time, the connectivity of the Zr₆ cluster did not play such an important role. MIP-200 proved to be the most stable MOF from the series in an aqueous environment; however, the loss of a small amount of monocarboxylic acid from the structure allowed thermal recrystallization of MIP-200 to an unknown phase so far.

MOFs for a broad spectrum of potential applications, *e.g.* gas storage and separation,¹¹⁻¹³ separation of volatile organic compounds,¹⁴ adsorption of pollutants from liquids,^{15,16} drug delivery and cancer therapy,^{17,18} energy applications,¹⁹ water harvesting and storage,^{20,21} sensing,^{22,23} and catalysis.^{24,25}

In spite of their great application potential, the number of industrial processes utilizing MOFs is so far limited. One of the reasons is the majority of MOFs' insufficient stability, i.e., low resistance to degradation during operating conditions.²⁶ There are several aspects of stability involving thermal, mechanical, or chemical stability, or their combinations. The chemical stability or more specifically, stability in a water environment, is crucial for many applications including adsorption and catalytic degradation of pollutants, sensing, antibacterial and antiviral use,^{22,27-31} drug delivery,³² or special applications such as using MOFs as pesticide carriers.³³ All those applications require aqueous media; therefore, MOFs' hydrolytic stability is of crucial importance. Laboratory testing is often limited to neat water; however, in real-life applications (the use of real contaminated water, body fluids, etc.) the aqueous medium contains many other dissolved compounds such as buffers or salts, and has different

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pH values, making the system much more complex and challenging for MOFs.

While the predictability of MOF synthesis has made significant progress in recent years, the predictability of MOF stabilities is so far unclear. It is understood that the MOF stability in aqueous media is strongly influenced by the strength of the coordination bonds between metals and linkers. In accordance with the hard and soft acid and base (HSAB) theory, the utilization of carboxylate linkers in combination with hard metals (Al(III), Zr(IV), Hf(IV), Ti(IV), etc.) leads to the construction of MOFs with increased stability in comparison with, e.g., the original MOF-5.34-39 Alternatively, the utilization of phosphonate linkers^{40,41} or oxalate⁴² can result in MOF structures of exceptional stability. The strength of coordination bonds, however, is not the only factor playing a critical role in the stability of MOFs. It is also affected by the length of connecting agents, the pK_a of a linker and its solubility in a given environment, or the SBUs' hydrophobicity and connectivity (the number of linker molecules bound to an SBU). It is assumed that, in general, higher connectivity, shorter connecting agents, and lower solubility of the linker lead to MOFs with a high stability in the water environment.^{36,37,43-47}

Many of the stability studies in aqueous media that were presented in the literature are based on a post-exposure analysis of the corresponding solids using powder X-ray diffraction (XRPD), gas adsorption, scanning electron microscopy (SEM), infrared spectroscopy (FTIR), or their combination with computational methods.^{26,32,45,46,48–53} For example, when MOFs' stability is inferred from XRPD patterns before and after the treatment, while ignoring the mass balance, presence of an amorphous phase, and changes in the chemical composition, erroneous conclusions can be drawn. In our recent studies, we have demonstrated that the structure of UiO-66, a Zr-based MOF considered one of the most stable MOFs, is compromised already at neutral pH or in the presence of commonly used buffers, such as TRIS or phosphate-buffered saline.^{54,55}

Here, we investigate the effect of the SBU connectivity and linker topicity on MOF stabilities in aqueous environments of different pH and correlate the changes made to MOFs in those conditions with catalytic activity in organophosphate degradation. We selected four Zr-MOFs constructed using di-, triand tetratopic carboxylate linkers - UiO-66, MOF-808, MIP-200, and PCN-222 (see Fig. 1). UiO-66 is based on terephthalate (benzene-1,4-dicarboxylate, BDC²⁻) linkers and the 12-connected Zr₆O₈ SBU,⁵⁶ forming an fcu-topology (in the ideal defect-free crystal).^{27,57,58} Next, MOF-808 is based on trimesate (benzene-1,3,5-tricarboxylate, BTC³⁻) linkers and 6-connected SBUs.⁶² MIP-200, which is a relatively young member of the Zr-MOF family and rated as exceptionally stable, is composed of tetratopic 5,5'-methylenediisophthalate (MDIP⁴⁻) linkers and 8-connected SBUs.⁵⁹ The last of the series, PCN-222/MOF-545, prepared simultaneously by Yaghi et al.60 and Zhou et al.63 (hereafter abbreviated as PCN-222), features 8-connected SBUs and tetratopic linkers (5,10,15,20-tetrakis(4-carboxyphenyl)porphyrin, abbreviated here as TCPP4-), forming a hexagonal structure.

The stability was studied by the combination of HPLC, analysing released linkers from the MOF, and the post-exposure analysis of the remaining solids (XRPD, N₂ adsorption, FTIR, differential thermal analysis (TGA/DTA), and chemical composition). Additionally, to put Zr-MOF stability into the perspective of an application, the parent and post-exposure MOFs were used for the catalytic degradation of methyl paraoxon (DMNP), an organophosphate (phosphate ester) commonly used as a simulant of organophosphate-based pesticides⁶⁴ and nerve chemical warfare agents.65 It is well described that missing-linker or missing-cluster defects in Zr-MOFs play a crucial role in catalytic reactions^{66,67} because open-metal sites as catalytic centres for the degradation of act organophosphates.^{68–71} In our previous work,⁵⁴ we have shown that the removal of BDC²⁻ from UiO-66 increases its catalytic activity considerably due to the increased number of openmetal sites. Here, we present a comparison of the catalytic activities of UiO-66, MOF-808, MIP-200, and PCN-222 after treatment at various pHs on the degradation of DNMP. The catalytic activity in the degradation of DMNP serves as an indication of whether the removal of linker in tested MOFs is connected with the formation of catalytically active defects or rather a structural collapse.

Experimental section

Instruments and materials

The list of chemicals is summarized in Table S1, ESI.[†] Similarly, instrumentation and measurement conditions (*i.e.*, XRPD, amorphous content, HPLC analyses of the linkers, monocarboxylates, DMNP and 4-NP, ICP-MS analyses of Zr, FTIR, gas adsorption and TGA/DTA, catalytic experiments) are described in ESI.[†]

Syntheses of MOFs

Synthesis of UiO-66,^{55,56} MOF-808,⁷² MIP-200,⁵⁹ and PCN-222 ^{60,73} followed the modified procedures published in referenced papers. The procedures are described in detail in ESI.[†]

Acid-base titration with released linker monitoring

The pH dependence of the amount of released linkers was determined using a procedure published by Klet *et al.*⁷⁴ and further elaborated by our group.⁵⁴ A beaker was charged with 50 mg of activated MOF (heated at 120 °C, 4 h, dynamic vacuum) and 50 mL of water, followed by 3 min of sonication to obtain a suspension that did not sediment within several minutes. The suspension was then stirred, and the pH was adjusted by the addition of 0.1 M NaOH using an automatic titrator Metrohm 906 Titrando. In each step, 0.005 mL of the NaOH solution was added with the rate of addition set to 0.04 mL min⁻¹. During the titration, the pH value gradually increased, and a 0.2 mL sample was taken immediately after reaching a preset pH value. The samples were immediately filtered through a PTFE microfilter (Whatman, 0.2 µm) and ana-

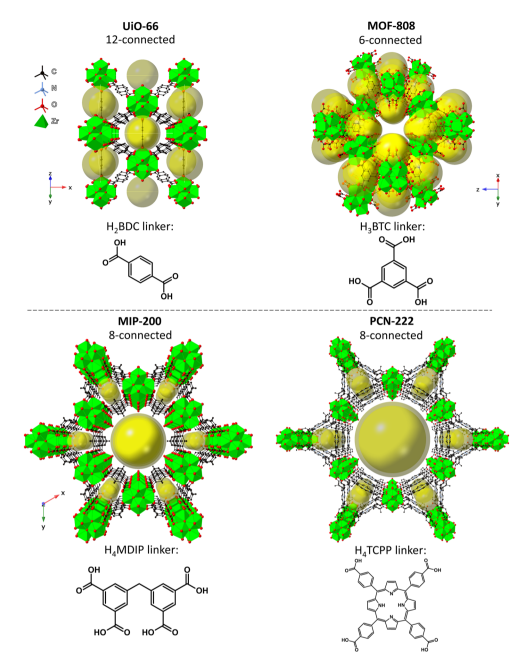


Fig. 1 Idealized structures of the tested MOFs: UiO-66⁶¹ (top left), MOF-808⁶² (top right), MIP-200⁵⁹ (bottom left), and PCN-222⁶³ (bottom right). The references refer to the corresponding CIF sources.

lysed using HPLC. These experiments provided data on the immediate release of the linkers at a certain pH; however, the kinetics of the linker release are not reflected in the results.

The same procedure was utilized for the determination of the immediate linker solubility at different pHs. 50 mg of the linker was mixed with 50 mL of water, followed by 3 min of sonication, and after that, the suspension of the linker in water was stirred for 2 h to achieve the equilibrium concentration at 25 \pm 1 °C. The immediate solubility of the linker during titration experiments indicates how the linker release from the MOF structures is limited by the linker solubility.

Kinetics of the linker release at different pHs

The kinetic stability of selected MOFs was studied at the natural pH of the MOF suspensions and at pH 6.0, 7.0, 8.0, 9.0, 10.0, and 11.0. In the case of MIP-200, a pH value of 12.0 was also used. In each kinetic experiment, 250 mg of MOF was suspended in water and sonicated for 3 min. The exact amount of added water was calculated for each pH to provide the total volume of 250 mL after the addition NaOH. After that, 0.1 M NaOH solution was added to reach the target pH value. The pH was kept constant during the kinetic experi-

ments using an automatic titrator (Metrohm 906 Titrando) which continuously adjusted pH with 0.1 M NaOH. The samples (0.2 mL) were taken out within 0 to 240 min (the periods were shorter during the first 60 min to achieve better rendering of the kinetic curves), filtered through PTFE microfilters (Whatman, 0.2 µm), and analysed by HPLC. Each kinetic experiment was repeated at least three times. After the kinetic experiments, the residual solids were collected by centrifugation (4000 rpm, Hermle ZK 496), washed three times with water and five times with acetone, air-dried, and reactivated by vacuum-drying at 120 °C. Only in the case of MIP-200, the samples were air-dried and reactivated under a vacuum only at laboratory temperature because at higher temperatures, MIP-200 recrystallizes, see below. All these samples were analysed by XRPD, N₂ adsorption, FTIR, and TGA/DTA, and the composition was determined by dissolving 10 mg of solid in 10 mL of 0.1 M NaOH and subsequent analysis of the solution. All released carboxylic acids were determined by HPLC-DAD. The metal content was determined after the acidic dissolution of 10 mg of the sample by ICP-MS analysis. Analogous analyses were also performed with the parent MOFs.

Results and discussion

Synthesis and characterization of MOFs

Zirconium-based UiO-66 was obtained by the solvothermal reaction of ZrCl₄, terephthalic acid (H₂BDC), and acetic acid as a modulator in the 1:1:4.8 molar ratio resulting in a microcrystalline powder. The XRPD patterns of the parent UiO-66 confirmed the reported structure with a broad diffraction peak between 2 and $6^{\circ} 2\theta$, indicating numerous structural defects caused by the formation of reo-phase (Fig. S1, ESI[†]).^{58,75} The N₂ adsorption isotherm has the typical shape for UiO-66, with a calculated BET specific surface area (S_{BET}) of 1263 m² g⁻¹ (Fig. 2D, Table 1). This value is higher than the theoretical surface area due to the presence of structural defects.⁷⁶ The TGA/DTA curves show a weight loss of approximately 10 wt% below 420 °C that includes release of moisture, residual solvents, and a part of monocarboxylic acids,⁵⁸ whereas the sharp weight loss of 45 wt% between 420 and 550 °C is, according to the MS analysis of evolved gases, related to the complete combustion of the BDC^{2-} linkers (Fig. S2, ESI^{\dagger}). The result is in good agreement with the content of 48.5% BDC²⁻ determined

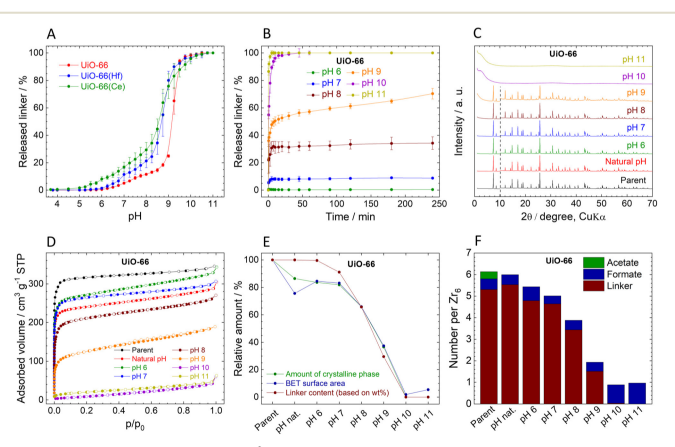


Fig. 2 Stability of UiO-66. (A). Immediate release of BDC^{2-} from Zr-based UiO-66 compared with UiO-66(Hf) and UiO-66(Ce). (B). Time-dependent BDC^{2-} release from UiO-66 at different pH values. (C). Comparison of XRPD pattern of the parent and post-exposure UiO-66; the diffractograms are normalized and shifted vertically to avoid overlaps. The dashed line at 10° 2 θ separates the zoomed area to visualize the diffractions at higher angles. (D). Adsorption (filled dots) and desorption (empty dots) isotherms of N₂ measured at 77 K. (E). Decrease of the crystalline phase percentage, S_{BET} , and linker content (all data are normalized, taking the parent UiO-66 as a default state with 100% values). (F). Molar ratios of the ligands in the parent and post-exposure UiO-66 to the Zr₆ metal clusters.

Table 1 Conversion of DMNP in relationship to structural changes in UiO-
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Sample	Crystalline phase/%	Linker removal after 4 h/%	$S_{\rm BET}/{ m m}^2~{ m g}^{-1}$	Conversion DMNP at 30 min/%	Conversion DMNI at 120 min/%
UiO-66 parent	100	0.0	1238	66	96
UiO-66 nat. pH ^a	87	<lod< td=""><td>937</td><td>73</td><td>94</td></lod<>	937	73	94
UiO-66 pH 6	84	0.4	1047	79	97
UiO-66 pH 7	82	8.8	1030	90	98
UiO-66 pH 8	66	34.3	816	96	100
UiO-66 pH 9	37	70.4	464	70	92
UiO-66 pH 10	0	100	23	4	6
UiO-66 pH 11	0	100	67	5	6

by HPLC analysis of the MOF dissolved in a basic solution. The mass content of all components is summarized in Tables S3 and S4 in ESI.†

MOF-808 was prepared according to the optimized procedure published previously.⁷² ZrCl₄ and H₃BTC were mixed in a 3:1 molar ratio in a 1:1 DMF/acetic acid mixture, followed by crystallization at 120 °C for 72 h. The XRPD pattern corresponds to that of a well-crystalline MOF-808 (Fig. S3, ESI†).⁷² The N₂ adsorption isotherm exhibits the shape typical for

MOF-808 containing multiple types of pores (Fig. 3D).⁷⁷ The thermogravimetric curve did not show any sharp features as in the case of UiO-66, which is typical for MOF-808 (Fig. S4, ESI†).^{72,77} The mass plateaued at 580 °C with a final weight loss of approximately 49 wt%. The MS analysis of evolved gases indicated that residual DMF, acetic acid, acetone, and moisture are released below 400 °C, representing approximately 20% of the total weight. The weight loss assigned to the linker combustion at higher temperatures is approximately

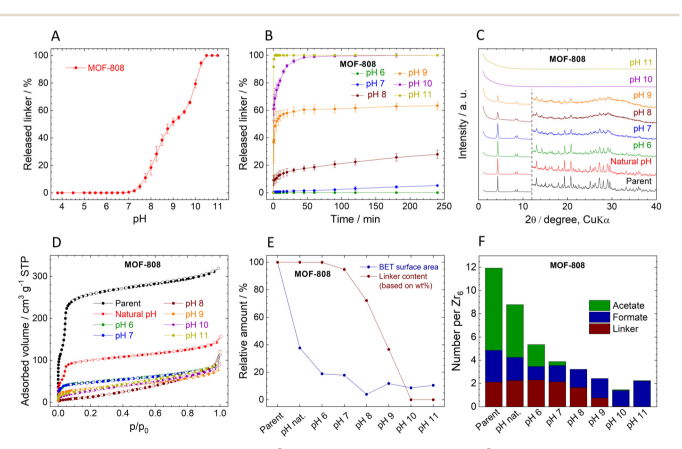


Fig. 3 Stability of MOF-808. (A). Immediate release of BTC^{3-} from MOF-808. (B). Time-dependent BTC^{3-} release from MOF-808 at different pH values. (C). Comparison of XRPD patterns of the parent and post-exposure MOF-808; the diffractograms are normalized and shifted vertically to avoid overlaps. The dashed line at 12° 2 θ separates the zoomed area to visualize the diffractions at higher angles. (D). Adsorption (filled dots) and desorption (empty dots) isotherms of N₂ measured at 77 K. (E). Decrease of S_{BET}, and linker content (all data are normalized, taking the parent MOF-808 as a default state with 100% values). (F). Molar ratios of the ligands in the parent and post-exposure MOF-808 to the Zr₆ metal clusters.

29%, which is in good agreement with 28.2 wt% of the BTC^{3-} determined by HPLC after the solid dissolution. The mass content of all components is summarized in Tables S5 and S6 in ESI.[†]

MIP-200 was prepared by a slight modification of the published procedure.⁵⁹ When following the original synthetic protocol,⁵⁹ we obtained a mixture of two crystalline phases. Several additional diffraction lines (for example 7.15, 8.73, and 12.14° 2θ) that appeared in the XRPD pattern correspond to MIP-201,⁷⁸ another MOF with 6-connected Zr₆ clusters (Fig. S5, ESI[†]). To obtain pure MIP-200 phase, the amounts of reactants and solvents were reduced to half with respect to the original procedure while the ZrCl₄ to H₄MDIP ratio was kept at 3:1. The crystallization was performed in DURAN glass bottles instead of a Teflon-lined autoclave. Possibly the surface of the Teflon insert promoted preferential crystallization of the MIP-201 phase. The S_{BET} area of 733 m² g⁻¹ calculated from N₂ adsorption isotherm (Fig. 4D) is below the reported value of ~1000 m² g⁻¹, 59 which is probably caused by the modified washing procedure. In the original report, MIP-200 was treated with boiling water before the adsorption measurements. We decided to forego this treatment to avoid dissolution of the linker and/or monocarboxylic acids. This hypothesis was confirmed by treating MIP-200 in neat water which increased S_{BET} to 938 m² g⁻¹ (see stability study below). The thermogravimetric analysis (Fig. S6, ESI†) displayed a total weight loss of 51% at 570 °C. The initial 12% being released between 50 and 210 °C can be attributed to the residual solvents, mainly acetone and moisture. The further 39 wt% released above 210 °C can be attributed to the linker combustion, which is in line with 42.1 wt% of the MDIP^{4–} determined by HPLC after the solid dissolution. For the total mass content of all components, see Tables S7 and S8 in ESI.†

PCN-222 was synthesized by a modified and scaled-up procedure^{60,73} from zirconyl chloride octahydrate and H₄TCPP in a molar ratio of 3 : 1 under modulated solvothermal conditions. The XRPD pattern (Fig. S7, ESI†) of the parent PCN-222 confirms its phase purity. The adsorption isotherm of N₂ showed a typical shape for PCN-222, which contains two types of pores in the structure with a calculated S_{BET} of 1869 m² g⁻¹ (Fig. 6D). Thermogravimetric analysis is rather inconclusive (Fig. S8, ESI†); there is not a sharp drop attributable to the combustion of the linker; moreover, mass loss is not finished at 800 °C, in which is consistent with earlier

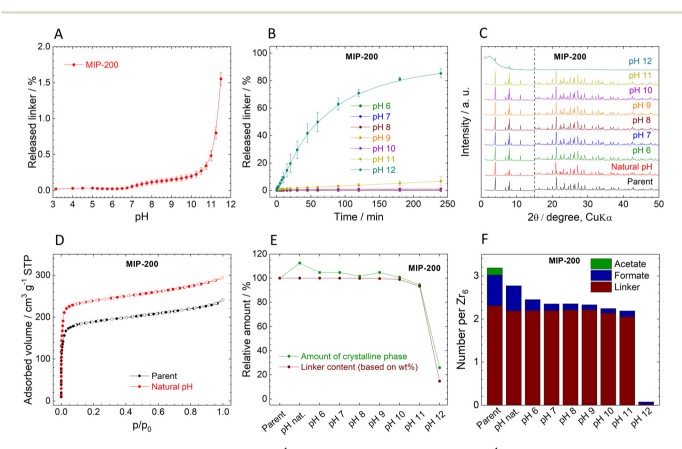


Fig. 4 Stability of MIP-200. (A). Immediate release of $MDIP^{4-}$ from MIP-200. (B). Time-dependent $MDIP^{4-}$ release from MIP-200 at different pH values. (C). Comparison of XRPD pattern of the parent and post-exposure MIP-200, the diffractograms are normalized and shifted vertically to avoid overlaps. The dashed line at 15° 2 θ separates the zoomed area to visualize the diffractions at higher angles. (D). Adsorption (filled dots) and desorption (empty dots) isotherms of N₂ measured at 77 K. (E). Changes of the crystalline phase percentage and linker content (all data are normalized, taking the parent MIP-200 as a default state with 100% values). (F). Molar ratios of the ligands in the parent and post-exposure MIP-200 to the Zr_6 metal clusters.

reports.^{79,80} The content of TCPP^{4–} determined by HPLC after the solid dissolution was 57.2 wt%. For total mass content of all components, see Tables S9 and S10 in ESI.†

Chemical stability of the MOFs in water

The chemical stability of UiO-66, MOF-808, MIP-200, and PCN-222 predetermines their applicability in many catalytic reactions, drug delivery systems, adsorption and water treatments, *i.e.*, in all applications that require an aquatic environment. The stability, evaluated by the release of the linkers from the MOF structure, was studied in pHs ranging from the natural value (pH of unadjusted water–MOF suspensions) to 11.0. In our stability studies, we used two approaches:

(i) *Immediate release of the linkers* – the titration of the MOF suspensions from the natural pH by the consecutive addition of 0.1 M NaOH solution. After reaching a selected pH, the immediate concentration of the released linker in the water phase was determined by HPLC. Clearly, the obtained titration curves describe the immediate linker release and bear no information about the long-term behaviour, which is relevant to reaction conditions in many applications.

(ii) *Kinetic stability* assessed by the time-dependent release of the linkers under constant pH. The target pH was adjusted by the addition of 0.1 M NaOH solution and then the pH value was kept constant during the experiment by an automatic titrator. The samples were taken out at predefined times and filtered, and concentration of the released linker was determined using HPLC. After 4 h, the remaining solids were collected and analysed and catalytic tests were done using these samples. The pH values for the kinetic experiments were selected based on the results from the titration curves, *i.e.*, at the MOFs' natural pH, at pH 6.0, which appears to be the pH at which some of the MOFs start to decompose; and at pH values from 7.0 to 11.0.

Stability of UiO-66

The observed immediate linker release from UiO-66 is in good agreement with our previous work.⁵⁴ The titration curve can be divided into four characteristic regions (Fig. 2A): (i) at pH 3.8 (natural pH) to 5.0, the amount of the released BDC²⁻ linker is below the detection limit (<0.01 mg L^{-1}), indicating the stability of UiO-66 in this pH region. (ii) At pH 5.25-8.75, moderate linker release rising from 0.05% to 18.4% of the total amount indicates relatively low pH sensitivity of UiO-66 in this region. (iii) pH 9.00-9.75 is characterized by high sensitivity towards pH with a sharp increase of the released linker amount - up to 97.3%; (iv) above pH 9.75, the linker release is complete. In general, UiO-66 can be considered stable only at pHs below 5, since higher pH values lead to gradual linker release and finally collapse of the structure at 10.0. Having the complete data on the Zr-based UiO-66 (hereafter labelled as UiO-66), we performed the same experiments with the Hf- and Ce-analogues of UiO-66, hereafter labelled as UiO-66(Hf) and UiO-66 (Ce) (syntheses in ESI,[†] XRPD patterns in Fig. S1, ESI[†]). The curves of immediate linker release (Fig. 2A) from UiO-66(Hf) and UiO-66(Ce) have a similar shape to that of UiO-66;

however, they indicate that the linker release is more pronounced. In the case of UiO-66(Ce), a small amount of BDC^{2–} linker (0.4%) was already released at the natural pH of 3.7. The lower pH stability of UiO-66(Hf) in comparison with UiO-66 is surprising because it is contrary to the generally accepted predictions based on the higher dissociation enthalpy of the Hf– O bond, according to which carboxylate-based Hf-MOFs should be more stable than their Zr analogues.^{81,82} It should be noted that the crystallite sizes, determined by the integral breadth of the first two diffraction lines, follow the trend UiO-66 ~ UiO-66(Hf) > UiO-66(Ce), see Fig. S1, ESI† and detailed discussion therein. Therefore, the smaller crystallite size of UiO-66(Ce) when compared to UiO-66 and UiO-66(Hf) can be partly responsible for the lower chemical stability. Further studies were performed only with Zr-based UiO-66.

The measured kinetics of the linker release from UiO-66 suggest that it is stable at its natural pH (3.8) for 4 h, as the H_2BDC concentration was below the detection limit during the entire experiment. Therefore, the corresponding curve is omitted in Fig. 2B. It is noteworthy, that at pH 6.0, the BDC²⁻ release is still below 0.5%. However, increasing pH led to much higher amounts of BDC²⁻ released. The kinetic curves of BDC²⁻ release are very fast, with BDC²⁻ release reaching a plateau within 30 min. The only exception is pH 9.0, where the BDC²⁻ is released throughout the 4 h of the experiment, probably due to the high sensitivity of UiO-66 to pH change in this region (see above).

The UiO-66 samples collected after the 4 h treatment at different pHs display XRPD patterns with the preserved original structure up to pH 8.0, where 34% of the BDC²⁻ linker was already released (Fig. 2C). This observation is in good agreement with our previous works.^{54,55} The diffraction line at $12^{\circ} 2\theta$, which is related to the degree of hydroxylation of the Zr_6 clusters⁸³ increases with the increasing pH. We highlight that the diffraction pattern of the sample treated at pH 9.0 still corresponds to the structure of UiO-66 even though there is 70% linker release and a significant reduction of the sample crystallinity (see below). When UiO-66 was treated at pH 10.0 and 11.0, the diffraction lines disappeared completely accompanied by the detected 100% linker release, indicating total destruction of the UiO-66 framework. Since no crystalline phase was detected in these samples, the products of the hydrolysis are probably amorphous zirconium oxide-hydroxide species with partially coordinated formate anions (Fig. 2F and Table S3, ESI[†]).

The content of the amorphous phase estimated using XRPD (parameters and results in Table S11, ESI†) follows the same trend as the linker release (Fig. 2E; the decrease in the crystalline phase is presented relative to the parent UiO-66-100%). Surprisingly, the formation of an amorphous phase occurred also in UiO-66 treated only with the natural pH, where no linker release was detected. Because this change cannot be caused by the BDC^{2–} release, we attribute the process to the removal of the acetate ligands, which are commonly bound to the defects on the zirconium coordination sites. As can be seen from HPLC analyses, acetates were

immediately and completely washed out already at the natural pH, whereas the formate ligands are strongly bound even above pH 10.0 (Fig. 2F, Tables S3 and S4, ESI†). The content of formate ligands is even increased at pH 10.0 and 11.0, probably due to the alkaline hydrolysis of residual DMF in UiO-66.⁸⁴ The increase of the amorphous phase content at higher pH correlates well with the extent of the linker release (Fig. 2E). We assume that linker and ligands are partially removed from the framework and the crystal structure is still preserved the vacated coordination sites are then occupied by water and OH⁻ ligands, to satisfy the charge and coordination number of the Zr₆ SBU as it is generally suggested for all Zr-MOFs with lower cluster connectivity than 12.⁸⁵ However, as soon as the structure starts to collapse this assumption is not valid.

The adsorption isotherms of N₂ are shown in Fig. 2D. Calculated S_{BET} after pH treatment is in good agreement with the structural changes described above (Fig. 2D and E, Table 1). At pH 6.0 and 7.0, S_{BET} slightly increased in comparison with the natural pH sample, which can be attributed to the increasing number of defects caused by the slight BDC²⁻ release.⁵⁸ At higher pH, S_{BET} gradually decreased and finally at pH 10.0 and 11.0 displayed no measurable porosity. After the normalization, the decrease in the S_{BET} corresponds to the decrease in the crystalline phase content, as shown in Fig. 2E. Also, FTIR spectra of samples treated at pH 10 and 11 (Fig. S9, ESI⁺) show only broad peaks in the region of 1400–1600 cm⁻¹, corresponding to adsorbed water molecules and coordinated OH⁻ ligands, and weak unresolved peaks in the region below 1000 cm⁻¹ attributed to the Zr-O stretching vibrations. The complete absence of vibrations of the linker molecules confirms the observed 100% BDC^{2-} release (Tables S3 and S4, ESI[†]).

Stability of MOF-808

The immediate linker release from MOF-808 is similar to that of UiO-66 (Fig. 3A). The titration curve can be divided into three regions: (i) at pH 3.8 (natural pH) to pH 6.5, the MOF is stable, and the concentration of the released linker is below the detection limit (*i.e.*, 0.05 mg L⁻¹). (ii) At pH 6.5–10.25, there is a steep increase in released BTC^{3–}, from approximately 0.03% for pH 6.5 to 94% at pH 10.25. The curve has stair-like characteristics in this range, with a region of lower sensitivity from pH 8.75 to 9.5. (iii) Above pH 10.25, the structure collapses, as documented by the complete release of the linker.

The kinetic curves of BTC^{3-} release (Fig. 3B) confirm the stability of MOF-808 at natural pH 3.8, *i.e.*, no released linker was detected, and therefore, the curve is not included in Fig. 3B. Only traces of BTC^{3-} were detected at pH 6.0 after 4 h (0.03%, close to the LOQ), indicating that MOF-808 is more stable in acidic environment than UiO-66. At pH 7.0 and 8.0, 5% and 33% of BTC^{3-} were released, respectively, which is comparable to UiO-66. At pH 10.0, MOF-808 was totally destroyed as evidenced by 100% BTC^{3} release. Importantly, in contrast to UiO-66, the kinetics of the release are much slower. The fast release at the beginning is followed by a slow continuous release over the following 240 min period.

The MOF-808 samples collected after the 4 h treatment at different pHs display XRPD patterns with the preserved original structure up to pH 9.0 even though the material already lost 63% of BTC^{3-} (Fig. 3C). However, the XRPD patterns of the samples treated at pH 7.0 and higher indicate a decrease in crystallinity. Probably due to the lower Zr_6 cluster connectivity of MOF-808 even small releases of BTC^{3-} have detrimental effects on the XRPD patterns. At pH 10.0 and 11.0, when all the BTC^{3-} was released, the material became amorphous. The FTIR spectra also confirm the loss of organic matter and formation of zirconium oxide-hydroxide (Fig. S10, ESI†). Unfortunately, MOF-808 cannot withstand homogenization with internal standard (ZnO), see Table S12, ESI† for additional data, and therefore, the determination of the crystalline phase content was omitted in Fig. 3E.

The composition of MOF-808 changes after the treatment at different pHs (Fig. 3F, Tables S5 and S6, ESI†). Parent MOF-808 contains approximately 7 molecules of acetates and 2.7 of formates per Zr_6 oxometallic cluster, which is more than the 6-connected Zr_6 cluster can theoretically accommodate together with BTC^{3-} . Thus, we assume that the excess of monocarboxylic acids is adsorbed in the pores even though the MOF was thoroughly washed with acetone. Interestingly, simple washing with water removed 33% of acetates and 29% of formates; the rest of the acetates were gradually washed at higher pHs together with a part of formates that are bound strongly even at highly basic pHs. The presence of adsorbed acetates and formates was confirmed by TGA/DTA with MS detection of evolved gases. At temperatures over 120 °C evolution of acetate was observed, (see Fig. S4, ESI†).

The adsorption isotherms of N_2 are shown in Fig. 3D. The adsorbed amount of N_2 and the calculated S_{BET} significantly decreased even after treatment at natural pH, with further decreases at increasing pH leading to complete loss of porosity at pH 8.0. The decrease in S_{BET} is in stark contrast with the loss of BTC³⁻ (see Fig. 3E and Table 2). However, it is in line with the evident decrease of crystallinity seen by XRPD. Moreover, the low amount of released BTC³⁻ might be due to re-adsorption to the remaining MOF-808 where it is blocking pores. In other words, based on linker release, the stability of

 Table 2
 Conversion of DMNP in relationship to structural changes in MOF-808

Sample	Linker removal after 4 h/%	$\frac{S_{ m BET}}{ m m^2 g^{-1}}$	Conversion DMNP at 30 min/%	Conversion DMNP at 120 min/%
MOF-808 parent	0.0	962	89	98
MOF-808 nat. pH ^a	<lod< td=""><td>362</td><td>74</td><td>97</td></lod<>	362	74	97
MOF-808 pH 6	0.04	181	32	63
MOF-808 pH 7	5.2	171	14	29
MOF-808 pH 8	27.9	37	14	27
MOF-808 pH 9	63.4	113	15	29
MOF-808 pH 10	100	82	6	11
MOF-808 pH 11	100	100	6	11

^a Natural pH is 3.8.

Stability of MIP-200

In contrast to UiO-66 and MOF-808, MIP-200 based on 8-connected Zr_6 clusters was significantly more stable during the titration experiment. Up to pH 11.0, the amount of the released MDIP⁴⁻ was close to the detection limit, and at pH 11.5 only 1.6% of the linker was released into the solution (Fig. 4A).

The exceptional hydrolytic stability of MIP-200 is further confirmed by the kinetic curves (Fig. 4B). At pH 6.0, MDIP⁴⁻ release was 0.013%, which was only slightly above the limit of detection (0.05 mg L^{-1}). A more evident MDIP⁴⁻ release occurred at pH 9.0 and 11.0; however, it did not exceed 0.5% and 7%, respectively. Since MIP-200 can withstand harsh conditions up to pH 11.0, we extended the testing pH range up to pH 12.0 where the amount of released MDIP⁴⁻ rapidly increased to 85%. In contrast to UiO-66 and MOF-808, the kinetic curves for MIP-200 showed continuous progress. For this reason, in the case of pH 11.0 and 12.0, we prolonged the time range to 72 h and 120 h, respectively. Interestingly, at pH 11.0, MDIP⁴⁻ was released during the whole timeframe of the experiment, while at pH 12.0 the release was completed within 8 h (Fig. 5A). The XRPD patterns of the resulting solids confirmed that the structure of MIP-200 is partially preserved with a high degree of amorphization at pH 11.0, while at pH 12.0, the structure is completely destroyed (Fig. 5B).

The XRPD patterns of the parent and post-exposure MIP-200 (Fig. 4C) indicate good preservation of the crystallinity up to pH 11.0. However, when samples treated at pH 6.0 and higher were reactivated at 100 °C overnight in a vacuum, the XRPD patterns revealed rather broad peaks of a new crystalline phase (compare Fig. 4C with Fig. S12A†). Meanwhile, the sample treated in neat water (natural pH) preserved the original structure even after the activation. As can be seen from Fig. 4F (see also Tables S7 and S8, ESI†), treatment with pH 6.0 and higher is related to the release of monocarboxylate

ligands. To confirm our hypothesis that the loss of monocarboxylates causes thermal instability, we have taken MIP-200 treated at pH 7.0 for 4 h and treated it overnight in 1 M acetic acid, 1 M formic acid or the same medium that is used for MIP-200 synthesis (mixture of acetic anhydride with formic acid). In all three cases, when the sample was then activated at 100 °C, the XRPD pattern did not change from the parent MIP-200 (see Fig. S12C†). Moreover, we found that the temperature is the responsible parameter for these changes; a vacuum alone did not induce a phase change (see Fig. S12B†). This was also confirmed by temperature-resolved XRPD (Fig. S13, ESI†). For this reason, the reactivation of MIP-200 was done only in a vacuum at laboratory temperature, and we omitted the measurement of adsorption isotherms.

The behaviour of MIP-200, which can be rearranged by simple heating when monocarboxylic acids are removed, while their reintroduction leads to restoring temperature stability, is quite unusual. A similar phenomenon was recently observed by Zhou *et al.*⁸⁶ where the rearrangement to a new phase was observed upon treatment with strongly coordinating solvents followed by activation under mild conditions. The process led to higher surface area, and moreover, in the published study the process was reversible.

The changes in the amorphous phase content are in line with the release of $MDIP^{4-}$ (see Fig. 4E). It is evident that the amorphous content did not increase too much when MIP-200 was treated with pH from natural to 11.0. Treatment with pH 12.0 is connected with high linker release, which led to partial destruction of the structure, as confirmed by an increase in the amorphous phase content. Adsorption isotherms of N₂ were measured only for the parent MIP-200 and the sample treated at natural pH (Fig. 4D). As discussed above, the increase in the adsorption amount is probably related to washing off the monocarboxylic acids. The FTIR spectra of MIP-200 (Fig. S11, ESI†) showed no significant changes even when the new crystalline phase was formed, indicating that the formation is not connected with visible changes in the building block constitution. After treating MIP-200 at pH 12.0,

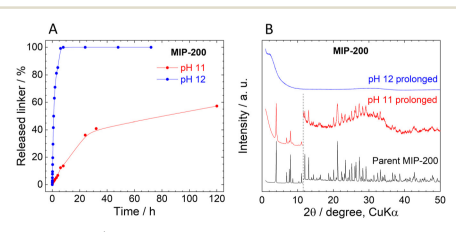


Fig. 5 (A). Prolonged time-dependent $MDIP^{4-}$ release from MIP-200 at pH 11.0 and 12.0. (B). XRPD patterns of the parent and post-exposure MIP-200 measured after kinetic experiments presented in panel A; the diffractograms are normalized and shifted vertically to avoid overlaps. The dashed line at 11.5° 2 θ separates the zoomed area to visualize the diffractions at higher angles.

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the FTIR spectrum showed a partial disappearance of vibrations belonging to organic matter which is in good agreement with the intensive loss of linker.

Stability of PCN-222

The immediate release of $TCPP^{4-}$ from PCN-222 shows a similar trend to MIP-200 (Fig. 6A). High linker release was observed only at high pHs when the amount of released $TCPP^{4-}$ reached 1%, 4%, and 43% at pH 10.5, 11.25, and 11.5, respectively.

The kinetic curves of TCPP⁴⁻ release confirm the low amounts of released linker (Fig. 6B). At the natural pH of 3.2, the release was below the detection limit of 0.005 mg L⁻¹, while at pH 6.0 and 7.0, the initially released TCPP⁴⁻ was later reincorporated back into the MOF structure. More significant linker release was observed at pH 8.0 and 9.0, but still, it was only 0.5% and 1.7% of the total amount, respectively. PCN-222 released 12.3% of the linker at pH 10.0 while partially keeping the crystal structure. The complete destruction of the network at pH 11.0 was accompanied by the 100% release of the linker. Similarly to MIP-200, the kinetic curves of PCN-222 showed continuous progress. The comparison of post-exposure XRPD patterns with that of parent PCN-222 showed that the PCN-222 structure was preserved (although slightly modified) even at pH 10.0, when 12.3% of TCPP⁴⁻ was released (Fig. 6C). pH 11.0 led to amorphization, which is in line with complete TCPP⁴⁻ release. Unfortunately, because of low mechanical stability of PCN-222, the amorphous content was not determined; for details, see Table S12, ESI.[†]

The composition of PCN-222 changed after the treatment at different pH (Fig. 6F, Tables S9 and S10, ESI[†]). Similarly to MOF-808, the number of formate ligands exceeds the number of coordination sites of the Zr_6 clusters, indicating that formates are partially adsorbed in the pores. This is supported by the fact that approximately 50% of formates were quickly removed at natural pH. In this case, no acetate ligands were present, since the synthesis was performed without the addition of acetic acid.

 N_2 adsorption isotherms confirm the stability of PCN-222 (Fig. 6D, Table 4). Very small changes up to pH 7.0 can be associated with the release of formates. Up to pH 10.0, the characteristic shape of the isotherms is preserved, indicating that two types of pores with different sizes are still present in the structure; however, the adsorbed gas volume drops signifi-

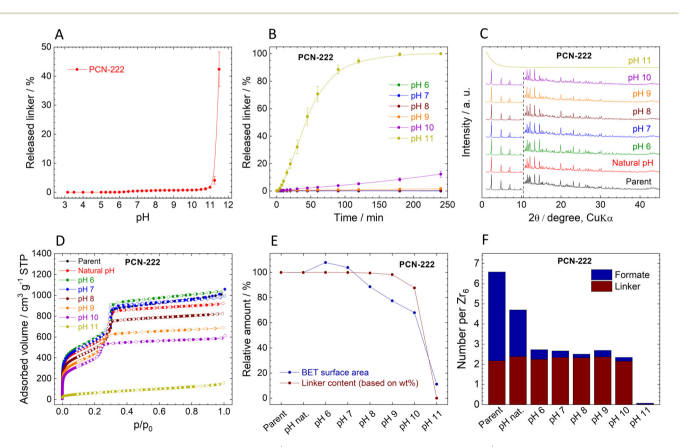


Fig. 6 Stability of PCN-222. (A). Immediate release of TCPP⁴⁻ from PCN-222. (B). Time-dependent TCPP⁴⁻ release from PCN-222 at different pH values. (C). Comparison of XRPD pattern of the parent and post-exposure PCN-222, the diffractograms are normalized and shifted vertically to avoid overlaps. The dashed line at 10.5° 2θ separates the zoomed area to visualize the diffractions at higher angles. (D). Adsorption (filled dots) and desorption (empty dots) isotherms of N₂ measured at 77 K. (E). Decrease in S_{BET} and linker content (all data are normalized, taking the parent PCN-222 as a default state with 100% values). (F). Molar ratios of the ligands in the parent and post-exposure PCN-222 to the Zr₆ metal clusters.

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cantly. The collapse of the structure at pH 11.0 is associated with loss of porosity. The correlation of the linker contents and S_{BET} show that both parameters are applicable for analysing the stability of PCN-222 and pH-induced changes to its structure (Fig. 6E). However, the N₂ adsorption seems to be more sensitive because it recorded decrease in S_{BET} before the significant amount of linker was released (Fig. 6E). The FTIR spectra also confirm the loss of organic matter and formation of zirconium oxide–hydroxide at pH 11.0 (Fig. S14, ESI[†]).

Summary of the MOFs stability

To summarize the above results, it is clear that the linker topicity and cluster connectivity have significant effects on the hydrolytic stability of Zr-MOFs. All MOFs are stable in neat water; however, around neutral pH, UiO-66 and MOF-808 (based on ditopic and tritopic linkers, respectively) start to degrade (compare Fig. S15 and S16, ESI[†]). Meanwhile, the MOFs based on tetratopic linkers, MIP-200 and PCN-222, survived up to pH 10.0 and 9.0, respectively. At the same time, it should be noted that the kinetics of the linker release get slower with increasing topicity. In the presented study, we investigated stability only in the time range of up to 4 h, which might be too short for a true assessment of MOFs based on tetratopic linkers.

Based on the literature, 36,37,45,46 it is expected that, in general, higher connectivity leads to more stable MOFs. Here, we report that the relationship between stability and Zr₆ cluster connectivity is not as straightforward. The structure of UiO-66, due to the 12-connected clusters, can survive a higher degree of linker removal accompanied by the formation of structural defects, whereas in the case of MOF-808 with 6-connected clusters, even trace amounts of the released linker lead to the collapse of the structure instead of the formation of defects. PCN-222 and MIP-200 with 8-connected clusters can also resist partial removal of linkers. It seems that a cluster connectivity of 6 is the critical point.

To prove that the linker release from MOFs is not governed by the limited solubility of the linkers, we determined the solubility of all four objective linkers as a function of pH (Fig. S17, ESI†). In all cases, the solubility was much higher than the concentrations detected in the tested solutions.

Finally, it is evident that a complex analysis based on linker release together with detailed post-exposure analysis is necessary to accurately describe the stability of MOFs and an approach limited to only some of those aspects may not be sufficient.

Catalysis using MOFs pre-treated at different pH values

For the catalytic tests, the parent and post-exposure MOFs with a different number of removed linkers and/or monocarboxylate ligands were used. Wide range of Zr-MOFs including UiO-66,^{68,71} MOF-808,^{70,87} and PCN-222 ^{68,86} has been tested for catalytic degradation of organophosphates so far. MIP-200 was tested as a catalyst for other reactions,⁸⁸ although it has not been tested for organophosphate degradation to date. In our previous work,⁵⁴ we described that the removal of BDC^{2–} from UiO-66 increased the activity because the creation of missing linker defects is directly related to new open-metal sites acting as catalytic centres.⁶⁹ Here, we took the recovered MOFs treated for 4 h at a given pH to test if the formed defects are connected with the formation of catalytically active centres or rather a structural collapse. It should be noted that we assume that the formed missing linker defects are compensated by coordinated water and OH⁻ ligands.

As a substrate for the model degradation reaction, we chose methyl paraoxon (DMNP), an organophosphate used as a simulant of pesticides or nerve warfare agents. DMNP is a suitable model compound because of its good solubility in water and ease of differentiating between substrate adsorption and catalytic degradation leading to 4-nitrophenol (4-NP) as the main product (see Fig. 7). Commonly, the catalytic degradation of DMNP is done in a buffer environment. However, buffers very often induce new defects or even MOF collapse.⁵⁴ For this reason, we tested all MOFs in neat water without pH adjustment (at natural pH values ranging from 3.9 to 4.6). After all catalytic experiments, we analysed the linker content in the solution. In all cases, the concentration of the released linker during the catalytic experiments was below the detection limit. Fig. 8 depicts the conversion of DMNP to 4-NP using different parent and post-exposure MOFs. The curves represent the formation of 4-NP. For the comparison of DMNP decrease with 4-NP formation, see Fig. S18, ESI.†

Conversion of DMNP to 4-NP using the UiO-66 catalyst followed the trends described earlier (Fig. 8 top left and Table 1). The parent UiO-66 is active; however, upon treatment with neat water (natural pH) or at pH 6.0, the reaction rate accelerated slightly. At this pH, the changes in the structure involve mainly the removal of acetate ligands from metal sites (see Table S3, ESI[†]). Treatment at pH 7.0 and 8.0 led to a significant increase in the catalytic activity, which is in good agreement with the higher amount of released BDC²⁻, while the MOF structure was still preserved. Obviously, the removal of the linker leads to the formation of catalytically active centres. However, at pH 9.0, the high linker release (70%) is already connected with a considerable degradation of the structure which results in a slower rate of reaction. Complete linker release at pH 10.0 and 11.0 is associated with a significant loss of catalytic activity. The remaining catalytic activity can be attributed to the resulting zirconium oxide-hydroxides.^{89–91}

MOF-808 does not follow the same trend as UiO-66 (Fig. 8 top right). The highest rate of degradation was recorded for

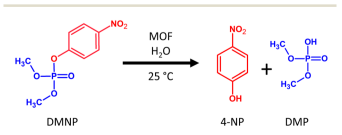


Fig. 7 Hydrolysis of DMNP to 4-NP and dimethylphosphate (DMP) catalysed by a Zr-MOF.

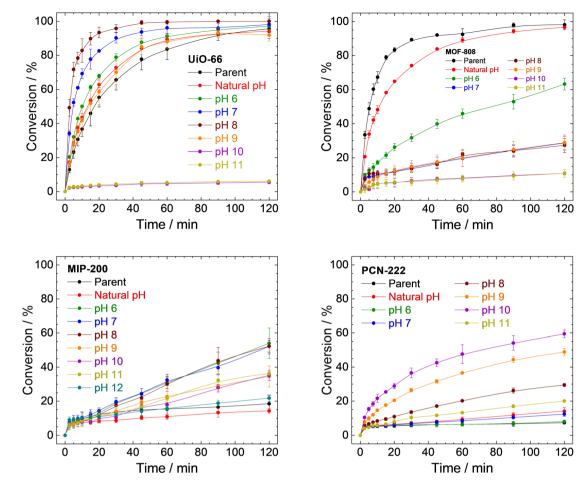


Fig. 8 Catalytic conversions of DMNP to 4-NP using parent UiO-66, MOF-808, MIP-200, PCN-222, and samples prepared by treating MOFs with a different pH for 4 h. The conversions are represented as an increase of 4-NP concentration related to the total amount of DMNP. The corresponding decrease in DMNP concentrations is presented in Fig. S18 in ESI.†

the parent MOF-808, and any treatment led to a decrease in activity. MOF-808 treated with neat water at natural pH is still comparable to the parent MOF-808; however, already treatment with pH 6.0, where only 0.04% of BTC^{3-} was released along with acetate and formate anions, displays a significant decrease in activity. A plausible explanation is that it is related to the damage to the structure (as evidenced by adsorption isotherms of N₂ and XRPD patterns; see Table 2 and Fig. 3C) instead of the formation of catalytically active defects even when a small portion of linker is removed.

MIP-200 does catalyse DMNP degradation; however, the decrease in DMNP concentration is much faster than the creation of 4-NP (compare Fig. S18, ESI†) in this case. This is probably due to the fast adsorption of DMNP coupled with a very slow degradation. As shown in Fig. 8 (bottom left) and Table 3, the parent MIP-200 and MIP-200 treated at natural pH provided only negligible activity. Acceleration of DMNP conversion was observed in the case of MIP-200 treated at pH between 6.0 and 11.0. In these cases, only a negligible amount of MDIP^{4–} was released; therefore, the removal of acetate and formate anions is mainly responsible for the activity enhance-

ment. The catalytic activity of MIP-200 treated at pH 12.0 decreased again to levels of the parent MIP-200, probably due to the large number of removed MDIP^{4-} accompanied by a loss of crystallinity.

Similarly to MIP-200, simultaneous adsorption and catalytic degradation were observed for PCN-222 (see Fig. 8 and S18, ESI†). The parent PCN-222 and PCN-222 treated at natural pH and pH 6.0 and 7.0 are only slightly active due to the negligible removal of the linker. The removal of formate ligands did not play any significant role in this case. The increase in the reaction rate was observed mainly when PCN-222 treated at pH 8.0, 9.0, or 10.0 was used. This matches well with the higher linker removal (compare conversion with linker removal in Table 4) accompanied by the formation of new catalytically active defects, as was observed for UiO-66. When PCN-222 was treated at pH 11.0, we observed total release of the linker and structural collapse, as well as a steep decline in the catalytic activity.

In general, the removal of linker molecules leads to an increase in catalytic activity, with the exception of MOF-808. This can be explained by the lower Zr_6 cluster connectivity in

 Table 3
 Conversion of DMNP in relationship to the detected changes in MIP-200

Sample	Crystalline phase/%	Linker removal after 4 h/%	$S_{\rm BET}/{ m m}^2~{ m g}^{-1}$	Conversion DMNP at 30 min/%	Conversion DMNI at 120 min/%
MIP-200 parent	100	0	733	14	19
MIP-200 nat. pH ^a	112	<lod< td=""><td>938</td><td>9</td><td>14</td></lod<>	938	9	14
MIP-200 pH 6	104	0.01	n.a.	19	54
MIP-200 pH 7	104	0.04	n.a.	20	52
MIP-200 pH 8	102	0.17	n.a.	18	52
MIP-200 pH 9	104	0.40	n.a.	14	35
MIP-200 pH 10	101	1.11	n.a.	11	35
MIP-200 pH11	95	6.77	n.a.	12	36
MIP-200 pH12	26	85.34	n.a.	12	22

 Table 4
 Conversion of DMNP in relationship to the detected changes in PCN-222

Sample	Linker removal after 4 h/%	$\frac{S_{\rm BET}}{{ m m}^2 { m g}^{-1}}$	Conversion DMNP at 30 min/%	Conversion DMNP at 120 min/%
PCN-222 parent	0	1870	6	7
PCN-222 nat. pH ^a	<lod< td=""><td>1869</td><td>7</td><td>14</td></lod<>	1869	7	14
PCN-222 pH 6	0.008	2015	6	8
PCN-222 pH 7	0.04	1942	7	12
PCN-222 pH 8	0.48	1657	14	29
PCN-222 pH 9	1.73	1448	26	49
PCN-222 pH 10	12.34	1270	37	59
PCN-222 pH 11	100	209	10	20
^{<i>a</i>} Natural pH is 3.2.				

MOF-808 when compared to the other tested MOFs. While the structure of UiO-66 with 12-connected clusters can be preserved even when a relatively high percentage of BDC^{2-} is removed and open metal sites are formed, the structure of MOF-808 collapses, and no catalytic centres are formed. For this reason, we assume that defect engineering based on the removal of linkers is not suitable for MOF-808. MOFs based on tetratopic linkers have in their parent form very low activity, which can be somewhat enhanced, however, even fine tuning of defects in the structure does not lead to catalytic activity comparable to that of UiO-66 or parent MOF-808. To summarize the catalytic part, defect engineering through linker removal can be used as a tool to increase the catalytic activity only for MOFs with higher connectivity.

Conclusions

This work examined the stability of UiO-66, MOF-808, MIP-200, and PCN-222 in pHs between 3.0 and 11.0 to delineate the relationship between linker topicity, Zr_6 cluster connectivity, and stability in aqueous media of various pHs. It should be noted that each MOF has its specifics and therefore we were changing more parameters at once.

We found that MOFs constructed from tetratopic linkers have significantly enhanced stability. Even though PCN-222 is

a mesoporous MOF with S_{BET} close to 2000 m² g⁻¹, it could tolerate a neutral or slightly basic environment much better than UiO-66 formed from BDC²⁻. At the same time, tetratopic linkers have much slower kinetics of release and we cannot rule out damage to their frameworks if MIP-200 or PCN-222 would be treated at neutral pH for days or weeks. Nevertheless, MIP-200 was the most stable MOF in aqueous media from the tested series.

The relationship between Zr_6 connectivity and stability is much more complicated. 12-connected UiO-66 can tolerate the loss of more than 30% of linker without detectable collapse of the structure. At the same time, when MOF-808 with 6-connected clusters loses even a small amount of the linker or monocarboxylic acid the structure starts to collapse. 8-connected MIP-200 and PCN-222 can tolerate a small linker release without structural damage. Studies with more MOFs would be needed to have more definite conclusion; however, from these results, it seems that as long as the actual connectivity in the Zr_6 cluster is higher than 6, defects can be created in the structure of a MOF without a structural collapse. It is, however, necessary to evaluate the effect of monocarboxylic acids as demonstrated on MIP-200 where their absence resulted in loss of thermal stability.

Additionally, we demonstrated that the determination of linker release is a sensitive tool for stability assessment only in the case of UiO-66. This is probably because, in all other studied cases, the MOFs were able to adsorb the released linker. In those cases, measurement of the adsorption isotherm or determination of the catalytic activity in organophosphate degradation gave a better picture of the structure. Analysis of the amorphous content is a powerful tool; however, it cannot be widely applied because of low mechanical stability of many MOFs.

Finally, the obtained results show that MOFs based on tetratopic linkers are not very active in organophosphate degradation. This might be related to the slow linker release, which makes the Zr_6 cluster less accessible to the organophosphate substrate. As there is a strong relationship between the preservation of the structure and catalytic properties, testing catalytic activity can also disclose whether defect engineering leads to the formation of defects or directly to the collapse of the structure.

Data availability

The data supporting this article have been included as part of the ESI.†

Conflicts of interest

There are no conflicts to declare.

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