



Cite this: *Chem. Commun.*, 2023, 59, 13639

Received 1st August 2023,
Accepted 18th October 2023

DOI: 10.1039/d3cc03735e

rsc.li/chemcomm

Covalent organic frameworks (COFs) are a prominent class of organic materials constructed from versatile building blocks *via* reversible reactions. The quality of imine-linked COFs can be improved by using amine monomers protected with benzophenone forming benzophenone imines. Here, we present a study on substituted benzophenones in COF synthesis *via* formal transimination. 12 *para*-substituted *N*-aryl benzophenone imines, with a range of electron-rich to electron-poor substituents, were prepared and their hydrolysis kinetics were studied spectroscopically. All substituted benzophenone imines can be employed in COF synthesis and lead to COFs with high crystallinity and high porosity. The substituents act innocent to COF formation as the substituted benzophenones are cleaved off. Imines can be tailored to their synthetic demands and utilized in COF formation. This concept can make access to previously unattainable, synthetically complex COF monomers feasible.

Covalent organic frameworks (COFs) are an emerging class of organic materials.^{1,2} Linked by dynamic covalent bonds, they are constructed from organic building blocks yielding permanently porous, crystalline, and stable frameworks. Their modular design allows for precise tailoring of the framework structure and properties.³ Owing to these properties, COFs are promising organic materials for various applications including energy storage,^{4,5} catalysis,⁶ gas storage and separation,⁷ and sensing.⁸ In imine-linked COFs, multivalent amines and aldehydes reversibly form imine bonds, ultimately leading to a crystalline, stable COF. However, the mechanisms of COF formation are complex^{9–11} and not well understood. Tedious optimizations of reaction conditions are often required to obtain materials of high

Substituted benzophenone imines for COF synthesis *via* formal transimination†‡

Josefine Sprachmann, ^a Niklas Grabicki, ^a Anna Möckel, ^a Jeremy Maltitz, ^a José Refugio Monroy, ^a Glen J. Smales ^b and Oliver Dumele ^a

crystallinity and porosity. Employing specific rigid or geometrically constrained monomers,¹⁰ metal triflate catalysis,¹² using modulators,¹³ or reconstruction of COFs¹⁴ are strategies that have been developed to improve the quality of COFs.^{9,10} Dichtel and coworkers proposed a strategy generally applicable to imine COFs, where multivalent free amines are replaced by *N*-aryl benzophenone imines, which react with aldehydes in a formal transimination reaction yielding high-quality COFs (Fig. 1a).¹⁵ This strategy of using benzophenone-protected amines has two key advantages: (1) higher porosity is achieved in comparison to a conventional reaction between a free amine and aldehyde; (2) benzophenone imines are more easily obtained and purified,¹⁶ with higher stability than their free amine counterparts.

So far, this strategy has only been utilized on a few examples while most COFs published until now, were constructed from free amines (Fig. 1b). Dichtel and coworkers have used a benzophenone imine substituted phenazine to construct a redox-active COF.¹⁷ Feng and coworkers have utilized a benzophenone

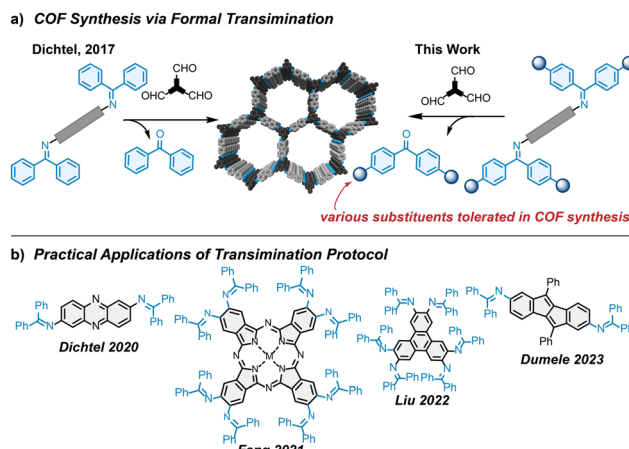


Fig. 1 (a) Schematic depiction of formal transimination approach as developed by Dichtel and coworkers and formal transimination with substituents on benzophenone. (b) Recent literature examples of benzophenone imines utilized in COF synthesis.

^a Department of Chemistry & IRIS Adlershof, Humboldt University of Berlin, Berlin 12489, Germany. E-mail: oliver.dumele@hu-berlin.de

^b Bundesanstalt für Materialforschung und -prüfung (BAM), Berlin 12205, Germany

† The PXRD data generated or analysed during this study are provided as a Source Data file at <https://doi.org/10.5281/zenodo.8337800>.

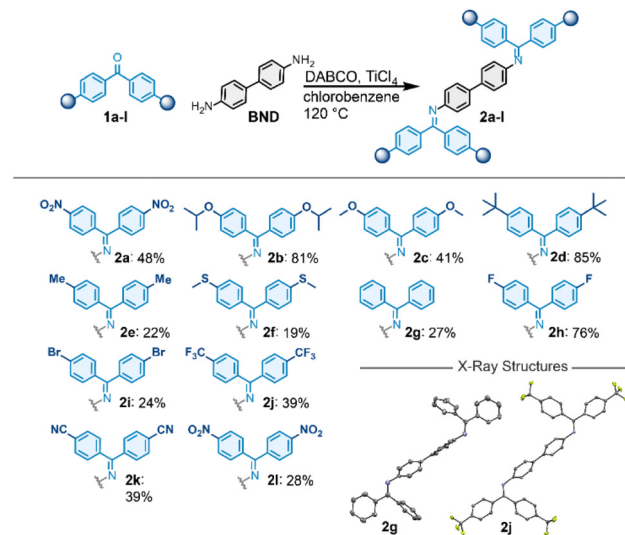
‡ Electronic supplementary information (ESI) available. CCDC 2240743, 1942397 and 2240744. For ESI and crystallographic data in CIF or other electronic format see DOI: <https://doi.org/10.1039/d3cc03735e>.



imine-decorated phthalocyanine in the formation of a conjugated COF to achieve suitable crystallinity.¹⁸ For Liu's tribenzimidazole-linked COF, the use of benzophenone imines was hypothesized to ensure a slow release of the COF node before an irreversible reaction locks the COF-linkage.¹⁹ In our lab, for the synthesis of antiaromatic dibenzopentalene (DBP) COFs, the use of benzophenone imine protection was essential as a diamino-DBP is not stable and degrades immediately at ambient conditions.²⁰ Hence, the use of benzophenone imines makes the synthesis of a robust, stable COF from an unstable building block feasible.

Until now, the formal transimination strategy has been limited to pristine benzophenone imines bearing no further substituents, though expanding this concept beyond benzophenone opens up new opportunities. One example has been shown recently where a COF building block equipped with benzophenone imines with -O*i*Pr groups for solubility was successfully used.²⁰ This is the first demonstration of how difficulties in the synthesis of complex COF building blocks can be tackled by functionalization of benzophenone imines. Another example could be conceived where electron-withdrawing groups could help further stabilize oxidatively unstable electron-rich COF building blocks. At the same time, quality optimization of the resulting COF is also in the scope of functionalized imines in COF synthesis. The above reflections have encouraged us to systematically explore the use of substituted imines in the synthesis of COFs *via* formal transimination. We synthesized a series of 12 *N*-aryl benzophenone imines with electron donating or -withdrawing groups in *para*-position. COFs with excellent crystallinity and porosity were constructed from these imines, showing that benzophenone substituents are compatible and innocent in COF formation. While electron donating and -withdrawing groups on the benzophenone moiety influence the rate constant of imine hydrolysis, the quality of COF is not clearly affected. The work shows that a variety of imines is compatible with COF synthesis, offering potential solutions for synthetically challenging COFs.

Motivated by the well-studied nonlinear Hammett plot of imine hydrolysis,^{21–23} we aimed at a substituent analysis with benzophenone imines to investigate the hypothesis of a slow-release mechanism.¹⁵ For a systematic substituent analysis we selected the *N*-benzophenone imines of benzidine (BND) and synthesized a series of benzophenones **1a–l** containing electron-withdrawing and electron-donating functional groups in *para*-position (Scheme 1). The benzophenones **1a–l** were coupled to BND, using DABCO and TiCl₄.^{15,24} The reactions gave the imines in moderate to good yields, and the crude products were conveniently purified by recrystallization, avoiding the use of column chromatography. We obtained 12 previously unreported BND-benzophenone imines spanning the Hammett scale from the electron-rich, -NMe₂ substituted **2a** to the electron-poor, -NO₂ substituted **2l**. All imines show excellent stability in ambient conditions and exhibit an improved solubility compared to the free-amine BND. Especially O*i*Pr-containing **2b** and OMe-containing **2c** show significantly enhanced solubility. Single-crystals suitable for single-crystal X-ray crystallography of CF₃-substituted imine **2j** and



Scheme 1 Synthesis of *N*-benzidine benzophenone imines **2a–l** including crystal structures of imines **2g** and **2j**.

unsubstituted imine **2g** were grown giving unambiguous proof for the structure of the imines.

The hydrolysis rates of imine bonds under reaction conditions similar to COF reaction conditions were investigated by following the intensity change at one wavelength using UV-vis spectroscopy (50 °C, 1,4-dioxane, 5000 equiv. AcOH, Fig. 2a and b). These conditions were chosen as the exact COF reaction conditions are not compatible with a spectroscopic setup (biphasic system, 120 °C, above boiling point in a pressurized tube). The hydrolysis of imines **2a–l** occurred with different rate constants

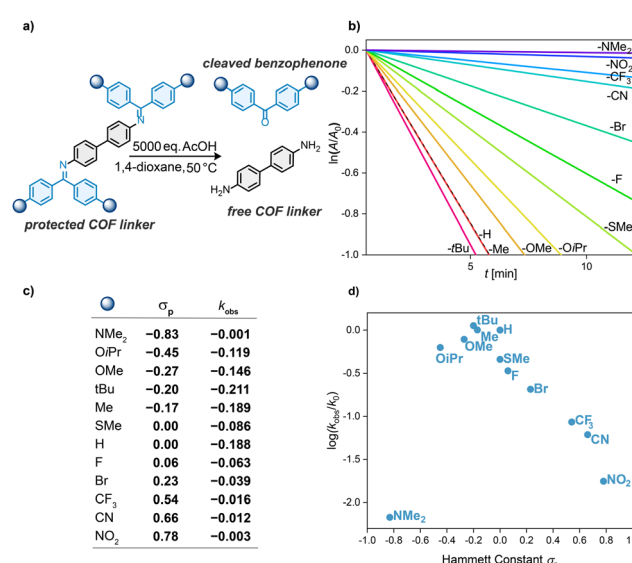


Fig. 2 (a) Reaction scheme of acid catalyzed imine hydrolysis, (b) kinetic plot $\ln(A/A_0)$ vs. t (linear regressions) following the hydrolysis of benzophenone imines at the absorption maximum in 1,4-dioxane, 50 °C, (c) reaction rate constants and *para*-Hammett parameters σ_p and (d) reaction rate constant k_{obs} vs. Hammett parameter σ_p .



k_{obs} dependent on the electronic nature of their substituent (Fig. 2c). Neutral to slightly electron-withdrawing substituents, with Hammett constants from -0.2 to 0 , resulted in the highest imine hydrolysis rates whereas the moderately electron-donating substituents, $-\text{Me}$ and $-\text{tBu}$, did not impact the reaction rate. Higher Hammett parameters led to slower hydrolysis in an approximately linear trend. With increasingly electron-rich substituents $-\text{OMe}$, $-\text{OiPr}$, and $-\text{NMe}_2$, the rate of hydrolysis was decreased (Fig. 2d). The results indicate that the conclusions drawn from earlier Hammett studies on imine formation and hydrolysis^{21–23} are also viable in COF reaction conditions. This inspired us to study the electronic modulation of the benzophenone moieties for improving COF quality. To investigate the implications on COF formation, we reacted the 12 substituted *N*-benzidine benzophenone imines **2a–l** with 1,3,5-triformylbenzene (TFB). The reaction affords BND-TFB COF, while the substituted benzophenone is being cleaved (Fig. 3). COF synthesis was carried out under solvothermal conditions (mesitylene/1,4-dioxane 1:1, 120 °C, 3 d). All substituents are tolerated in COF formation, leading to BND-TFB COF (Fig. 3a). Characterization of the material matches with previous reports (ESI,‡ Sections S5 and S6).^{15,25–27} BND-imines with $-\text{OMe}$, $-\text{Me}$, or $-\text{SMe}$ substituent lead to COFs with moderately increased Brunauer–Emmett–Teller surface area (SA_{BET}) compared to the COF from unsubstituted benzophenone imine ($-\text{H}$), while other substituents lead to COFs with slightly or markedly decreased SA_{BET} . Plotting the Hammett constant of the benzophenone substituents against the SA_{BET} of the resulting COF reveals a widely scattered plot (Fig. 3b). The most electron donating substituent $-\text{NMe}_2$ and the most electron withdrawing substituents $-\text{CN}$ and $-\text{NO}_2$ give a comparatively lower SA_{BET} . All other entries gave high SA_{BET} ranging from $1823 \text{ m}^2 \text{ g}^{-1}$ to $2388 \text{ m}^2 \text{ g}^{-1}$. While surface areas vary considerably, all COFs show high-crystallinity in their powder X-ray diffractograms (Fig. 4). To quantitatively compare crystallinities we used estimated domain sizes as a measure of crystallinity. Scherrer analysis using the full-width half maximum of the (100) reflex, revealed estimated domain sizes between 28 nm and 40 nm, typical values for COFs. The domain sizes are uniformly scattered showing that all substituents across the Hammett scale act innocent in COF formation, leading to high-quality materials (Fig. 3c). Contrary to our initial consideration, the quality of a COF is not modulated by the electronic nature of the protecting imine

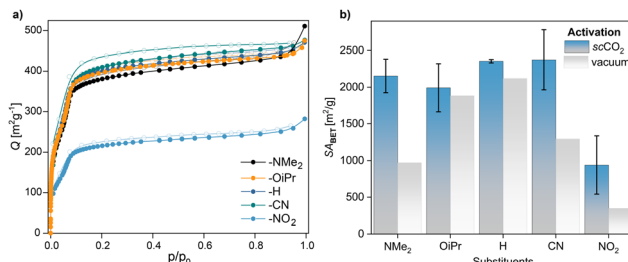


Fig. 4 (a) N_2 adsorption–desorption isotherms at 77 K and (b) improved SA_{BET} of COFs after scCO_2 activation compare to SA_{BET} after vacuum activation.

but gives reliable high-quality materials in the substituent range from $-\text{OiPr}$ to $-\text{CF}_3$ ($-0.45 < \sigma_p < 0.54$) throughout. This suggests that the mechanism by which formal transimination improves the COF quality is more complex than the previously proposed slow-release mechanism.¹⁵ The practical implication of this result lies in the variety of substituents that are tolerated in COF formation. Imine groups can be tailored to any synthetic demands, enabling the incorporation of building blocks that are not attainable with conventional methods. For instance, the $-\text{OiPr}$ group can solubilize otherwise poorly soluble monomers. Substitution with electron-withdrawing functional groups ($-\text{CN}$, $-\text{NO}_2$) can improve stability of particularly electron-rich monomers.

In addition to the COF synthesis conditions, the so-called activation process (removal of solvents from the microporous material) plays a crucial role in obtaining high-quality COFs.^{28,29} Among several methods, activation by supercritical CO_2 (scCO_2) has proven to be especially superior.³⁰ To shed further light to our study and to further investigate a possible correlation between Hammett parameter and COF quality, we replicated the COF syntheses, this time followed by scCO_2 activation. We picked five benzophenone imines across the Hammett scale and repeated the COF synthesis in duplicates. For all examples we achieved significantly higher SA_{BET} with this method compared to using vacuum activation (Fig. 4). scCO_2 might be more efficient than solvent washes in removing residual benzophenone from the COF pores. With scCO_2 activation, using benzophenone imines substituted with $-\text{NMe}_2$, $-\text{OiPr}$, $-\text{H}$ or $-\text{CN}$ all gave very high, similar surface areas, independent of the imine-substituent. Only *para*-nitro benzophenone imine consistently

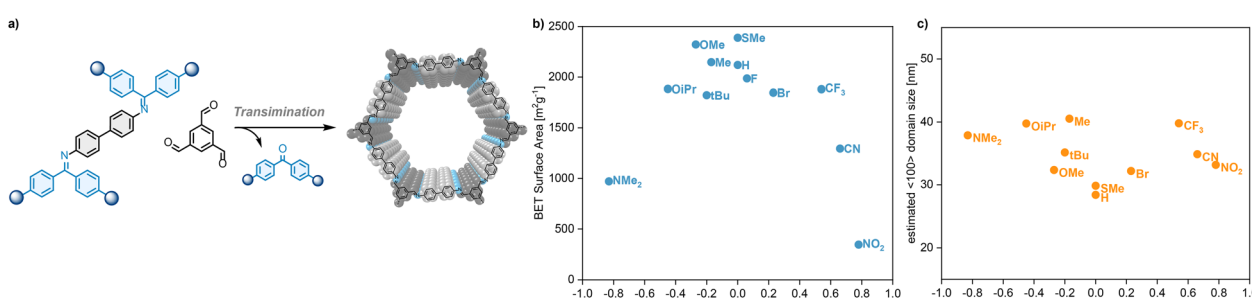


Fig. 3 (a) Reaction scheme of COF formation from substituted *N*-benzidine benzophenone imine and TFB. (b) Correlation of BET surface area and Hammett constant σ_p . (c) Correlation of estimated domain size and Hammett constant σ_p . The domain size was estimated using the Scherrer equation and is uncorrected for peak asymmetry and instrument line broadening.



gave lower porosity COFs of $SA_{BET} = 939 \text{ m}^2 \text{ g}^{-1}$. The highest surface area for BND-TFB COF was achieved using $-\text{CN}$ substituted benzophenone imine **2d**. With $2783 \text{ m}^2 \text{ g}^{-1}$, the BET surface area is only $60 \text{ m}^2 \text{ g}^{-1}$ below the theoretical surface area and the highest value for BND-TFB COF reported so far in the literature. It is interesting to note that the reaction of unsubstituted BND-benzophenone imine ($R = H$) with TFB very consistently yields COFs with a high surface area. In comparison, the functionalized benzophenone imines ($R = \text{NMe}_2, \text{O}i\text{Pr}, \text{CN}, \text{NO}_2$) also lead to high surface area COFs though with BET surface areas varying in a wider range for repeated experiments.

In summary, we have prepared a series of 12 *para*-substituted *N*-benzidine benzophenone imines and studied their hydrolysis under COF formation conditions. The imines were employed in COF reactions with TFB, all yielding high-quality BND-TFB COF. Unsubstituted benzophenone-imines give the most reliable and reproducible results while substitution with electron-donating or electron-withdrawing groups increases the variance in surface area. Although the Hammett parameter of the substituents clearly influences the hydrolysis rate of the imines, no clear trend can be observed for COF surface areas. The results indicate that electronic effects of the imines are not the decisive factor for COF quality (based on SA_{BET}). It is likely that imine hydrolysis occurs too fast for the imines to efficiently act as slow-release agent but they rather act as initial solubilizing group and modulators in the dynamic COF formation reaction. Therefore the hydrolysis rate of the imine monomer does not influence the quality of the resulting COF (*i.e.* SA_{BET}), although the imine hydrolysis rates are in a similar time range as the initial COF formation.³¹ Further research is necessary to elucidate the complex mechanism of COF formation involving benzophenone imines. We show that *para*-substitution of benzophenone imines is well tolerated in the synthesis of COFs, emphasizing the generality of the formal transimination approach. This can facilitate the synthesis of previously unattainable, synthetically complex COF monomers as benzophenone imines can be tailored to the synthetic demands, *e.g.* with solubilizing or electronically stabilizing substituents. Utilizing such substituted benzophenone imines or even imines beyond benzophenone will be a vital tool in the preparation of synthetically demanding high-performance imine COFs in the future.

We thank Edon Vitaku for helpful discussions.

Conflicts of interest

There are no conflicts to declare.

Notes and references

1 A. P. Côté, A. I. Benin, N. W. Ockwig, M. O'Keeffe, A. J. Matzger and O. M. Yaghi, *Science*, 2005, **310**, 1166–1170.

- 2 C. S. Diercks and O. M. Yaghi, *Science*, 2017, **355**, 923.
- 3 N. Huang, P. Wang and D. Jiang, *Nat. Rev. Mater.*, 2016, **1**, 16068.
- 4 S. Kandambeth, V. S. Kale, O. Shekhar, H. N. Alshareef and M. Eddaoudi, *Adv. Energy Mater.*, 2022, **12**, 2100177.
- 5 H. Gao, A. R. Neale, Q. Zhu, M. Bahri, X. Wang, H. Yang, Y. Xu, R. Clowes, N. D. Browning, M. A. Little, L. J. Hardwick and A. I. Cooper, *J. Am. Chem. Soc.*, 2022, **144**, 9434–9442.
- 6 M. Traxler, S. Gisbertz, P. Pachfule, J. Schmidt, J. Roeser, S. Reischauer, J. Rabeah, B. Pieber and A. Thomas, *Angew. Chem., Int. Ed.*, 2022, **61**, e202117738.
- 7 H. Lyu, H. Li, N. Hanikel, K. Wang and O. M. Yaghi, *J. Am. Chem. Soc.*, 2022, **144**, 12989–12995.
- 8 L. Wei, T. Sun, Z. Shi, Z. Xu, W. Wen, S. Jiang, Y. Zhao, Y. Ma and Y. B. Zhang, *Nat. Commun.*, 2022, **13**, 7936.
- 9 F. Haase and B. V. Lotsch, *Chem. Soc. Rev.*, 2020, **49**, 8469–8500.
- 10 L. Bourda, C. Krishnaraj, P. Van Der Voort and K. Van Hecke, *Mater. Adv.*, 2021, **2**, 2811–2845.
- 11 H. Li, A. M. Evans, W. R. Dichtel and J. L. Bredas, *ACS Mater. Lett.*, 2021, **3**, 398–405.
- 12 M. Matsumoto, R. R. Dasari, W. Ji, C. H. Feriante, T. C. Parker, S. R. Marder and W. R. Dichtel, *J. Am. Chem. Soc.*, 2017, **139**, 4999–5002.
- 13 T. Ma, E. A. Kapustin, S. X. Yin, L. Liang, Z. Zhou, J. Niu, L.-H. Li, Y. Wang, J. Su, J. Li, X. Wang, W. D. Wang, W. Wang, J. Sun and O. M. Yaghi, *Science*, 2018, **361**, 48–52.
- 14 W. Zhang, L. Chen, S. Dai, C. Zhao, C. Ma, L. Wei, M. Zhu, S. Y. Chong, H. Yang, L. Liu, Y. Bai, M. Yu, Y. Xu, X. W. Zhu, Q. Zhu, S. An, R. S. Sprick, M. A. Little, X. Wu, S. Jiang, Y. Wu, Y. B. Zhang, H. Tian, W. H. Zhu and A. I. Cooper, *Nature*, 2022, **604**, 72–79.
- 15 E. Vitaku and W. R. Dichtel, *J. Am. Chem. Soc.*, 2017, **139**, 12911–12914.
- 16 J. P. Wolfe, J. Åhman, J. P. Sadighi, R. A. Singer and S. L. Buchwald, *Tetrahedron Lett.*, 1997, **38**, 6367–6370.
- 17 E. Vitaku, C. N. Gannett, K. L. Carpenter, L. Shen, H. D. Abruna and W. R. Dichtel, *J. Am. Chem. Soc.*, 2020, **142**, 16–20.
- 18 H. Zhong, M. Wang, M. Ghorbani-Asl, J. Zhang, K. H. Ly, Z. Liao, G. Chen, Y. Wei, B. P. Biswal, E. Zschech, I. M. Weidinger, A. V. Krasheninnikov, R. Dong and X. Feng, *J. Am. Chem. Soc.*, 2021, **143**, 19992–20000.
- 19 Q. Zhang, F. Zhang, J. Dong, M. Shao, M. Zhu, D. Wang, Y. Guo, J. Zhang and Y. Liu, *Chem. Mater.*, 2022, **34**, 6977–6984.
- 20 J. Sprachmann, T. Wachsmuth, M. Bhosale, D. Burmeister, G. J. Smale, M. Schmidt, Z. Kochovski, N. Grabicki, R. Wessling, E. J. W. List-Kratohvil, B. Esser and O. Dumele, *J. Am. Chem. Soc.*, 2023, **145**, 2840–2851.
- 21 G. M. Santerre, C. J. Hansrote and T. I. Crowell, *J. Am. Chem. Soc.*, 1958, **80**, 1254–1257.
- 22 James O. Schreck, *J. Chem. Educ.*, 1971, **48**, 103–107.
- 23 E. H. Cordes and W. P. Jencks, *J. Am. Chem. Soc.*, 1962, **84**, 832–837.
- 24 M. Higuchi, M. Tsuruta, H. Chiba, S. Shiki and K. Yamamoto, *J. Am. Chem. Soc.*, 2003, **125**, 9988–9997.
- 25 Q. Gao, L. Bai, Y. Zeng, P. Wang, X. Zhang, R. Zou and Y. Zhao, *Chem. – Eur. J.*, 2015, **21**, 16818–16822.
- 26 L. Bai, S. Z. F. Phua, W. Q. Lim, A. Jana, Z. Luo, H. P. Tham, L. Zhao, Q. Gao and Y. Zhao, *Chem. Commun.*, 2016, **52**, 4128–4131.
- 27 L. Bai, Q. Gao and Y. Zhao, *J. Mater. Chem. A*, 2016, **4**, 14106–14110.
- 28 C. Gropp, S. Canossa, S. Wuttke, F. Gándara, Q. Li, L. Gagliardi and O. M. Yaghi, *ACS Cent. Sci.*, 2020, **6**, 1255–1273.
- 29 D. Zhu and R. Verduzco, *ACS Appl. Mater. Interfaces*, 2020, **12**, 33121–33127.
- 30 C. H. Feriante, S. Jhulki, A. M. Evans, R. R. Dasari, K. Slicker, W. R. Dichtel and S. R. Marder, *Adv. Mater.*, 2020, **32**, 1905776.
- 31 C. Feriante, A. M. Evans, S. Jhulki, I. Castano, M. J. Strauss, S. Barlow, W. R. Dichtel and S. R. Marder, *J. Am. Chem. Soc.*, 2020, **142**, 18637–18644.

