Design and synthesis of coordination polymers with chelated units and their application in nanomaterials science

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The advances and problems associated with the preparation, properties and structure of coordination polymers with chelated units are presented and assessed. The metalloligand strategy, and columnar-layered strategy, as well as approaches based on supramolecular building blocks and supramolecular building layers in the design of coordination polymers with chelated units are considered. The conventional (solvent evaporation, diffusion, hydro(solvo)thermal, ionothermal, microfluidic, surfactant-thermal, and in situ spacer methods as well as synthesis in supercritical CO₂) and alternative (microwave-assisted, electrochemical, mechanochemical, and sonochemical methods) synthesis routes are described. Special attention is paid to the post-synthetic approaches to the preparation of coordination polymers with chelated units including post-synthetic modification, post-synthetic deprotection, and post-synthetic exchange. The thermal transformations of coordination polymers, resulting in the formation of nanocomposite materials, are considered. The bibliography includes the papers published within last five years.

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1. Introduction

Coordination polymers (CPs) are objects of increasing interest to researchers. These advanced structures can be considered from two different points of view: only scientific knowledge or practical orientation. To date, owing to the efforts of specialists in various fields such as polymer, coordination, material, bioinorganic and analytical chemistry, photochemistry, catalysis, etc., considerable advances in this field of chemistry have been achieved. It is important that CPs are porous materials with adjustable size of pores, have vast surface areas and intriguing structural topologies. A great interest in the development of new CPs is associated with their promising applications, especially in such areas as ion exchange, adsorption, and separation processes, biomedical applications, drug delivery, sensor technologies, heterogeneous and biomimetic catalysis, luminescence, proton conductivity, etc.

In addition, CPs have gained more and more attention as nanocomposite materials precursors, and they serve not only as a starting material, but also as stabilizers of formed nanoparticles. From the view of the design of materials, solid-state thermolysis of CPs with different structures and compositions is a simple and rational way to synthesize novel nanostructured materials. Thermal decomposition of CPs under various conditions has been widely studied with the aim to prepare a variety of nanomaterials with desired sizes and morphologies. It is important that various nano-sized materials, for example, carbon, metals, metal oxides, etc., can be obtained using CP or their precursors.

Coordination polymers with chelated units are an interesting topic in the branch of coordination chemistry due to its specific properties and high stability than ordinary complexes due chelate or macrocyclic effect. They relate to a special class of polymeric metal chelates containing metal in the main chain, whose special feature is breaking a polymer backbone after a metal is removed.

The present review aims to describe the latest advances in the field of the coordination polymers with chelated units including main strategies and approaches to their design as well as synthetic methodologies of their preparation. It contains the basic directions of using coordination polymers as precursors of nanostructured materials, a conclusion and the main challenges.

2. Main strategies and approaches

To date, a wide variety of strategies and approaches to the design of coordination polymers with chelated units have been developed. Among them, metalloligand and pillar-layered strategies as well as approaches based on supramolecular building blocks and layers are the most common.

2.1. Metalloligand strategy

Though a great amount of building units has brought to vast variety of target materials, using well-defined metal complex as a building unit (i.e. metalloligand) is unique for building projected architecture. CP assembly based on metalloligand approach proceeds mainly in two stages: (1) synthesis of a metalloligand by interaction between well-defined ligands and metal ions (mostly 3d metal ions), and (2) reaction of the metalloligand with a second type of a metal ion, which plays a role of nodal units in a framework. The metalloligand used as a building unit provides structural rigidity, which puts ancillary functional groups in predetermined conformation. These ancillary functionalities can then coordinate a secondary metal ion and take part in self-assembling through such weak intermolecular interactions as H-bonds. Two classes of metalloligands are respectively distinguished, which have competitive M–L-bond sensitive groups, and H-bond sensitive functional groups. It is this that metalloligand differs from a simple complex in which the metal ion is coordinated with a certain number of “ordinary ligands” that do not contain an attached functional group. It is important that both metal complexes with monofunctional ligands and metal chelates can act as a metalloligand. The latter constitute the main group of metalloligands used to create CP with chelated units.

Among metalloligands, the widest applied as building units are metalloporphyrins (M-Pp). By integration of different functional peripheral substituents into β-pyrrol and/or meso-positions, M-Pp can have several multifunctional properties. With M-Pp used as building units for metal-organic frameworks (MOFs), easy molecular modification of Pp widens possibilities of structural design. Since most elements of periodic table can be included in Pp, different metal centers can be integrated into porous surface of MOFs without changing a framework topology. Moreover, using M-Pp as building units for MOFs gives an opportunity to adapt photochemical, redox, catalytic, and other properties of frameworks.

Most widely used Pp-based ligands for CPs building are tetrakis(4-carboxyphenyl)-Pp (L1) and tetrakis(4-pyridyl)-Pp (L2). The various multiporphyrin frameworks with different topology and dimensionality are built on the basis of coordination of their peripheral carboxyphenyl or pyridine fragments. Therefore, L1 and L2 are almost inexhaustible sources for building either H-bonding or M–L-driven CPs. It should be noted that for design of Pp-based MOFs, clusters formed by rigid Lewis acid metals, such as Zr6, Hf6, Zr8, Hf8, and Fe3, are used as most stable secondary building unit (SBUs). Thus, systematic study of Zr- and Hf-Pp MOFs has brought to discovery of isostructural PCN-221(M) based on metalloligands and Zr6 or Hf6 clusters. In particular, cluster Zr6O18 is characterized by idealized Zr6 cube, in which each Zr atom is in one vertex and each face of the cube is crowned with one μ4-oxygen atom. On each face of the cube carbonate from Pp ligands bridges two Zr atoms thus obtaining 3D MOF with very rare (4,12)-connected fctw topology, in which two types of polyhedral cages with diameters ~1.1 and ~2.0 nm and cage opening ~0.8 nm are found. As another example, we shall notice a solvothermal reaction of Fe–L1, ZrCl4, and benzoic acid in DEF during 48 h at 120 °C, which leads to PCN-222(Fe) with a framework consisting of square-planar Pp ligands linked to Zr6 clusters. In this CP, each Fe–L1 block is coordinated with four 8-connected clusters Zr6 with twisted
angle forming 3D network with Kagome-like topology (Fig. 1). It is remarkable that the mesoporous framework contains large 1D hexagonal open channels with a larger diameter 3.7 nm ranking among the largest for MOFs. In addition, this framework is preserved even after immersion in concentrated HCl, which is rare in MOF materials. We shall also notice highly stable MOFs with 3D nanochannels PCN-224(Ni, Co, Fe) assembled with six-connected Zr$_6$ cluster and M–L$_1$ metalloligands. The PCN-224 series not only demonstrates high BET surface area (2600 m$^2$/g), but also stays unchanged at pH of aqueous solution from 0 to 11.

Metal–metalloporphyrin frameworks (MMPFs) form a class of coordination networks self-assembled using reasonable choice of metal-containing SBUs and specially developed M-Pp metalloligands. As an example of CP of MMPF series, we shall consider MMPF-6 based on FeL$_1$–Cl metalloligand and SBU Zr oxide cluster, which is obtained under solvothermal conditions (Fig. 2).

Porous MMPFs including [Zn$_2$(HCOO)$_2$(MnL$_1$)], [Cd$_2$-(HCOO)$_2$(MnL$_1$)], [Zn$_2$(HCOO)(FeL$_1$)] and [Cd$_3$(H$_2$O)$_6$(μ$_2$-O)(FeL$_1$)$_2$] units are synthesized by heating the mixture MCl–L$_1$ (M = Mn and Fe) and nitrates M’ (M’ = Zn or Cd) in mixed solvent from DMF and acetic acid. First three compounds are built from M’$_2$(COO)$_4$ paddlewheel subunits linked with ML$_1$ bridge and formate ligands for formation of their 3D

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**Fig. 1** The crystal structure and underlying network topology of PCN-222(Fe). The Fe–L$_1$ (a; blue square) is connected to four 8-connected Zr$_6$ clusters (b; light orange cuboid), generating a 3D network with Kagome-like topology (d, e) and large 1D channels (c; green pillar). Zr: black spheres, C: gray, O: red, N: blue, and Fe: orange. [Reprinted with permission from ref. 56. Copyright 2012, Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim.]

**Fig. 2** (a) FeL$_1$–Cl metalloligand. (b) Zr$_6$O$_{18}$(CO$_2$)$_6$(H$_2$O)$_8$ SBU. (c) Hexagonal and triangular 1D channels of MMPF-6. Color scheme: C, gray; O, red; Cl, green; Zr, turquoise. [Reprinted with permission from ref. 57. Copyright 2012, American Chemical Society.]
connectivity. Formate column is heterogeneously connected with M and M’ cations in two first CPs and links homogeneously M’ cations in third CP. A μ₂-O bridging FeL₁ dimer plays a role of decadentate ligand for linking of 10 Cd cations with formation of interesting 3D coordination network of fourth CP (Fig. 3).

The reaction of M-L₁ (M = Co, Ni, Cu, VO) with lead nitrate gives MOFs with [Pb₂(M-L₁)] units, which differ in framework topologies and Pp configurations changing from flat to wavy to bowl-shaped.⁶⁸ It is important that Pp cores play pivotal role in the formation of different coordination frameworks by controlling Pp stacking, L₁ coordination modes and the coordination spheres of the Pb(n) cations.

We should also notice CZJ-1 with [Zn₂(MnOH-L¹)(L’)] unit, where L’ = N,N’-di(4-pyridyl)-1,4,5,8-naphthalenetetracarboxyldimide.⁶⁴ This porous MOF has a type of twice interpenetrated cubic α-Po topology, in which Zn₂(COO)₄ paddlewheel clusters are bridged by MnOH–L¹ in 2D sheets additionally linked by bridging organic pillared spacer L’ with formation of 3D porous structure.

Among other used Pp ligand, we shall notice 5,15-bis(dicarboxyphenyl)-Pp. Thus, for example, frameworks with [Cu₂(ML)₆]₄⁻ unit have been obtained from ML metalloganids, where M = Zn, Ni, Pd, MnCl, Ru(CO),⁸⁴ using two different methods: interaction between Cu nitrate and ML in DMF at 80 °C during 24 h or by diffusion of triethylamine vapor into the abovementioned mixture. It is interesting that this framework forms 3D network consisting of two-component combination (Cu₂ paddlewheel nodes and ZnL building units) and having internal spherical cavity of 20 Å in diameter. This interior cavity is surrounded totally by 16 available metal centers of two different metals, i.e. with eight Zn atoms from ZnL units and eight Cu atoms from paddlewheel units (Fig. 4).

In another interesting example, narrowly distributed nano- and micro-meter CPs are obtained from Mn[n]–L metalligand and Co acetate, in which morphologies are diversified from amorphous spheres in crystalline cubes through different period of reaction (Fig. 5).⁸⁵

In order to develop CPs based on Pp ligands with a possibility to coordinate more metal centers, more than four carboxyl or Py groups were involved as, for example, in octatopic Pp-ligand 5,10,15,20-tetrakis(3,5-biscarboxyl-phenyl)-Pp.⁶⁹,⁷⁰ In particular, it is used for building of three porous M-Pp frameworks ZJU-18, ZJU-19 and ZJU-20,⁷¹ whose structures are 3-periodical, binodal, edge-transitive tbo networks (Fig. 6) showing intercrossed porous windows of about 11.5 Å and porous cavities of about 21.3 Å in diameter. These three isostructural MOFs were synthesized by heating Mn[n]Cl-L and MnCl₂, NiCl₂ or CdCl₂ mixture, respectively, in mixed solvent from DMF and acetic acid at 80 °C during a week.

We shall also notice MOF based on tetrazolyl-Pp with {[Mn(n)]Cl[Mn(n)Cl-L]₄} unit, called UTSA-57, which is built from Mn₄Cl₄(L)₈(H₂O)₄ clusters, linked with metalloganid Mn₄Cl₄-L (Scheme 1).⁸⁶ UTSA-57 has a rare scu MOF topology and is built of 8-connected bridging SBUs with 1D square
nanotube-like 20 Å channels along c axis. In addition, UTSA-57 presents 3D stable microporous structure.

An example of metalloligand for MOF formation through H-bonds can be Co(II)-5,10,15,20-tetra(4-(4-acetateethyl)phenoxy)phenyl-Pp. In it, M-Pp molecules are linked with each other by intermolecular H-bonds with formation of 2D layer, which is additionally linked by π···π interactions for formation of 3D supramolecular structure having one type of micropores with 3.98 × 6.47 Å. MOF has a permanent porosity with Langmuir surface area of 158.79 m² g⁻¹ and BET surface area of 97.70 m² g⁻¹. In another example, M-Pp coordination framework [Co(H₂L)]ₙ, where H₂L = meso-tetra(4-carboxyphenyl)-Pp, having 2D layered coordination network structure shows H-bonds between carboxyl groups within 2D layer as well as between neighboring layers.

The popular metalloligands are metallosalens (salen is N,N₀-bis(salicylidene)ethylenediamine), in which additional functional groups, such as carboxylate, pyridine and benzoate-groups are in para- and meta-positions with respect to OH-groups on aromatic ring. As an example, it can be noted using solvothermal synthetic strategy for CPs building based on metallosalens as spacers. In this strategy, a metalloligand based on carboxy-functionalized salen spacer 2,6-bis[(4-carboxy-anilino)carbonyl] pyridine and a metal salt are solved in DMSO/DMF mixture, and then heated, which leads to CPs formation with governed size from nano- to micro-meter level. The ligand chelates metal cations in the center of the structure and provides two terminal carboxylate groups for CP growth. The particle size is regulated by changing the reaction temperature, so that a temperature increasing leads to bigger particles. Initially formed nanowires aggregate with formation of cub-like clusters, which undergo intrastructural fusion into uniform cubic-shaped particles. In addition, the size of the formed structures is influenced by DMSO/DMF ratio, at that higher content of DSMO leads to bigger cubes.

A special attention should be given to the metalloligand strategy in synthesis of heterometallic CPs, since used metalloligand already containing ion of an initial metal fits ideally for interaction with respective secondary metal ions. In conventional synthesis using the reaction between mixture of metal salts and organic ligands, it is very difficult to control synthesis of heterometallic complexes, since there is always high probability of formation of homometallic polynuclear complexes. At the same time, when metalloligand strategy is used, side reaction is minimal, and exclusively heterometallic CP can be obtained. A great number of metalloligands is developed, which provide targeted production of ordered heterometallic architectures.

Interesting heterometallic Cu(n)–Zn(n) CPs 2A and 2B are synthesized by interaction of CuL metallaligand, where H₂L = N,N₀-bis(salicylidene)-1,3-propanediaimine, with Zn perchlorate and sodium diycyanamide (dca) using different molar ratios of reagents (Scheme 2). The obtained compounds are 2D and 3D CPs,
respectively, with a common \([\text{Cu}L_2\text{Zn(NCN)}_2\text{]}_2\) trinuclear unit, in which dca fragments have \(\mu_{1,5}\)-bridging coordination regime. It is interesting that the isomeric CPs is a rare example of \(\text{genuine supramolecular isomerism}\). It should be noted that the reaction of similar \([\text{Cu}L_2\text{]}\) metalloligand (where \(\text{H}_2\text{L}\) is \(\text{N},\text{N}'\text{-bis(2-methylsalicylidene)-1,3-propanediamine}\)) with zinc perchlorate hexahydrate and sodium dicyanamide, in a \(2 : 1 : 2\) molar ratio at room temperature, resulted in a heterometallic discrete trinuclear complex, \([\text{Cu}L_2\text{Zn(NCN)}_2\text{]}_2\) (1).

Cu(i)-salen CPs \([\text{NiL}_2[C\text{u(i)CN]}_9]_n\), \([\text{Cu(ii)L}_2[C\text{u(i)CN]}_9]_n\) and \([\text{NiL}[\text{Cu(ii)L]}_2]_n\) are obtained by direct linking metallosalen precursors with \([\text{Cu(i)CN]}_n\) chains and Cu\(_{12}\) clusters using the metalloligand strategy (Scheme 3).\(^{101}\) In these CPs, efficient integration of catalytically active transition metals of Ni(ii)/Cu(ii)salen units and photoactive Cu(i) species in one solid polymer is achieved, which makes it possible to obtain materials with double catalytic properties.

Heterometallic CPs based on metallosalens have shown their efficiency in a range of catalytic processes, in particular, Mn(ni)/Cu(n)-salen MOFs in separation of chiral and achiral small molecules,\(^{99}\) Mn(ni)/Ni(n)-salen 1D CPs in olefin epoxidation,\(^{90}\) Cu(n)/Cu(n)-salen-based 1D CP in three-component Strecker reaction.\(^{102}\)

Wide range of metalloligands is presented by acac-complexes, which are used, for example, in synthesis of heterometallic CPs with \([\text{NiAg}_3(\text{acac})_3(\text{NO}_3)_2]_n\), \([\text{NiAg}_3(\text{acac})_3(\text{ClO}_4)_2]_n\) and \([\text{NiAg}_3(\text{acac})_3(\text{CF}_3\text{SO}_3)_2]_n\) units, where acac is acetylacetonate anion, through integration of respective Ag salts into the reaction with Ni(acac)\(_2\) metalloligand (Scheme 4).\(^{103}\) It is important that anions in CPs are decisive factors for formation of different structures. For example, first CP is outstanding 3D chiral framework, second CP has 2D inorganic layers, and third CP shows 2D (4,5)-connected layered architecture.

We shall notice crystal engineering of heterometallic CPs based on FeL\(_3\) and Fe(L')\(_3\) chelates, where \(L = 1,3\text{-di}(3\text{-pyridyl})\) propane-1,3-dionato and \(L' = 1,3\text{-di}(3\text{-pyridyl})\) propane-1,3-dionato, which act as metalloligands in reactions with Ag salts.
forming CPs with \([\text{AgFeL}]_n\) and \([\text{AgFe} (L')_n]_n\) units (Scheme 4). It is important that structure of obtained CPs depends on nature of used counter-ions. Thus, reaction between \(\text{FeL}_3\) and \(\text{AgBF}_4\) leads to CP, in which Ag centers link metalloligands in discrete nanotubes, while the reactions with \(\text{AgPF}_6\) and \(\text{AgSbF}_6\) have given CPs, in which metalloligands are bound in sheets. At the same time, interaction between \(\text{FeL}_3\) and \(\text{AgNO}_3\) leads to layered CP with \([\text{Ag}_3\text{Fe} (L')_3]_n\) unit.

Metalloligand strategy is used for production of heterometallic 3D CPs containing Cu(II) and trimethyltin as nodes. The first step of this synthetic way was in situ preparation of generated 2D CP with \([\text{Cu} (\mu-LH)]_n\) unit (\(LH_2 = \text{pyridine-2,5-dicarboxylic acid}\)), whose reaction with \(\text{Me}_3\text{SnCl}\) has given heterometallic 3D CP with \([\text{Cu}(\text{Me}_3\text{Sn})_2(\mu-L)_2]_n\) unit (Scheme 6). The obtained CP is 4,4-connected polymer with \(\text{sqc}\) topology, which contains the paddlewheel-shaped core consisting of two heterometallic Sn(IV)/Cu(II) macrocycles.

In another interesting example, 3d–4d heterometallic Co(II)–Zn(n) and Co(II)–Cd(n) CPs are obtained from two Co(II)-based metalloligands containing appended groups of arylcarboxylic acids 5-[picolinamido]-isophthalic acid and 4-{[pyridine-2-carbonyl]-amino}benzoic acid in different positions. Arylcarboxylate groups coordinate secondary metal ions, Zn(n) and Cd(n), with production of different 3D networks (Fig. 7). It is important that all networks show ordered positions of secondary metal ions and unique, even unprecedented network topologies.

An example of using chelating bpy ligands is porous 3D CP with \([\text{La}_4\text{Cu}]_n\) unit based on tetranuclear \(\text{La}_4\) cluster and Cu(i)-metalloligand, including 6,6’-dimethyl-5,5’-dicarboxy-bpy, where bpy is 2,2’-bipyridine. In flexible porous heterometallic MOFs of rock-salt-type based on [CoL]_3 metallo-ligand (\(H_2L = 4,4’\text{-dicarboxy-bpy}\)) and trivalent lanthanide cations La, Ce, Pr, Nd, Sm, Eu, Gd, Tb, and Er, six carboxylates in the upper part of each coordination octahedron of Co(II)-metalloligand are bound with M cations. If a MOF contains M cation smaller than Nd(III), the MOF is crystallized in cubic spatial group \(\text{Fm}3\text{m}\), while other MOFs with larger M are crystallized in low-symmetric rhombic spatial group \(\text{Fd}3\text{m}\) due to asymmetric 10-coordinated bicapped structure of square antiprism of larger M cation.

There is an interesting using of macrocyclic metalloligand NiL (\(H_2L = 1,4\text{-dihydro-2,3-dioxo-5,6:9,10:13,14\text{-trienzeto}[1,4,8,11\text{tetraazacyclotetradeca-7,11-diene-7,12-dicarboxylic}]}) for production of CPs with \([\text{Mg} (\text{NiL})]_n\), \([\text{Zr}_2(\text{NiL})]_n\), and \([\text{Ph}(\text{NiL})]_n\) units having infinite one-chained helical structures. All NiL metalloligands in CPs are located in head-to-tail positions (Fig. 8, left column), in order to form very similar single-twisted helical CPs (Fig. 8, right column).

### 2.2. Pillar-layered strategy

Pillared MOFs are widely studied platforms with numerous applications; they consist of 2D layers, which have ligating sites...
for linking with ditopic columns, as a rule, derivatives of 4,4'-bipy, where 4,4'-bipy is 4,4'-bipyridine, leading to 3D architectures. Functionalization either with spacers within 2D layers or ditopic columnar ligands allows to easily adjusting structures and properties of pillared MOFs.67

Most widely used for building pillared-layered CPs are M-Pp, since they form rigid 2D sheets, which are easily connected by pillared ligands. Thus, for example, Pp paddlewheel frameworks (PPF family) are obtained using M-Pp and columnar molecules for connection of paddlewheel SBUs.110 The PPF assembling can be considered as two-staged process. At the first stage, carboxyl groups of M-Pp, where Pp = tetra-(4-carboxyphenyl)-Pp, bind to $\text{M}_2(\text{COO})_4$ paddlewheel SBUs (M = Zn, Co), which are insensitive to Pp metals. In the resulting 2D layer consisting of M-Pp and paddlewheel SBUs, axial paddlewheel directions are easily available for connection with ancillary ligands. At the second stage,
the class of bipyridyl molecules is used as pillars for linking 2D layers occupying axial sites. Therefore, for demonstration of targeted M-Pp pillaring, the controlled strategy of production of Pp-based pillared-paddlewheel frameworks PPF-11, including metal pairs Zn/Zn, Co/Co, Mn/Zn and Fe/Zn, is developed, where first and second metal are the metal centers in a Pp core and in a paddlewheel cluster, respectively, by changing pillaring ligands from 4,4'-bipy to 2,2'-dimethyl-4,4'-bipy. These compounds have 3D framework, in which 2D layers are pillared with a steric controlled 2,2'-dimethyl-4,4'-bipy remaining structurally unbound metal centers inside Pp. Therefore, using this approach, a desired control over pillar coordination is reached.

Another example is F-MOF and DA-MOF, which consist of two Zn(n)-Pp columns of [5,15-dipyridyl-10,20-bis[pentafluorophenyl]porphyrin]-Zn(n) and [5,15-bis[4-(pyridyl)ethynyl]-10,20-diphenylporphinato]-Zn(n), respectively (Scheme 7). Photogenerated exciton migrates at a pure distance up to ~45 Pp pillars during its lifetime in DA-MOF, and only up ~3 Pp pillars in F-MOF with high anisotropy along specific direction. This means that Pp molecular structures play important role in

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Fig. 8 Coordination environments of metal centers, and metal-binding mode, [head-to-tail] arrangement and saddle shape of NiL (left column) and left-(M) and right-handed (P) helical chains (right column) in [Mg(NiL)] (top), [Zr2(NiL)2] (middle) and [Pb(NiL)2] (bottom). Mg, Zr, and Pb, pine green; Ni, cyan; N, blue; O, red; C, grey. [Reprinted from ref. 109 with permission of Springer.]
exciton jump. Great distance and directed energy transfer in DA-MOF suggest promising applications of this compound for development of efficient light-harvesting and energy-transporting materials.

2.3. Approach based on supramolecular building blocks

Other approaches to MOFs design were inspired by success in supramolecular chemistry. Thus, supramolecular building blocks (SBB) approach is a powerful method for achievement of hierarchical assembling, which provide qualitatively far higher level of control over structural properties and porous structures. This concept is based on using preliminary chosen 0-periodical metal–organic polyhedron (MOP) as a building unit, which allows to self-assembling greater and more complicated ensembles of several metallic nodes and organic spacers at the supramolecular level. As compared with molecular building units, these supramolecular cages or clusters can sharply increment porous architecture, and very often they have well-defined coordination geometry, which results in considerable degree of predictability in synthetic design. MOFs having high symmetry based on highly-connected polyhedral cage of molecular building units, which actually are SBBs, can provide elegant control over a structure due to their high connectivity, and also can design themselves features of limited nanospace and very high surface area. In most detail, this approach was studied for the example of M-Pp CP by MOP building using $[\text{M}_2(\text{carboxylate})_4]$ paddlewheel fragments. It is important that MOPs can be functionalized either on vertices, or on faces.

Using MOPs as building units promotes scale of MOFs to higher complexity level, and can give interesting MOF platforms. Thus, for example, 24-, 18-, or 12-connected MOPs were used as SBBs for formation of highly connected rht-MOFs, gea-MOFs, and fcu-MOFs, respectively. Moreover, several isophthalate derived Pp ligands were successfully prepared during building nanometer polyhedral

![Scheme 7](image)

Scheme 7 Synthesis routes of the isostructural F-MOF and DA-MOF compounds and photographs of resulting crystals. [Reprinted with permission from ref. 19. Copyright 2013, American Chemical Society.]

![Fig. 9](image)

Fig. 9 The three types of polyhedral cages presented in MMPF-3: (a) cubohemioctahedron, (b) truncated tetrahedron, and (c) truncated octahedron. (d) 3D structure of MMPF-3 illustrating how its polyhedral cages are connected. [Reprinted with permission from ref. 61. Copyright 2012, Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim.]
cage-containing MMPFs. Thus, MMPF-3 were obtained solvothermally from Pp ligand 5,15-bis(3,5-dicarboxyphenyl)-10,20-bis(2,6-dibromophenyl)-Pp and Co$_2$(H$_2$O)$_2$(CO$_2$)$_4$ molecular building units, which afford cubohemioctahedral SBBs serving as 12-connected nodes in the resulting network of fcu topology. It is worth noticing that MMPF-3 shows three types of polyhedral cage (Fig. 9): a cubohemioctahedron, truncated tetrahedron, and truncated octahedron, which are interlinked.

SBBs are obtained from custom-designed tetra-dicarboxy-Pp self-assembled with M(II) (M = Zn and Cd) cations, which are uniform polyhedrons based on Pp molecular building units. Faces of Pp fragments link triangular M$_2$(CO$_2$)$_3$ or M(CO$_2$)$_3$ fragments with formation of small cubicuboctahedral SBBs, which, in turn, are fused with adjacent SBBs on opposite faces of each Pp fragment. The resulting highly symmetric broadened topology pecu networks, MMPF-4 (M = Zn) and MMPF-5 (M = Cd) show two different polyhedral cages and are permanent microporous (Scheme 8).

2.4. Approach based on supramolecular building layers

We shall also notice another conceptual strategy, in particular, the approach based on supramolecular building layers (SBLs), which made it possible to design and produce pillar MOFs with far higher complexity (Fig. 10). This unique and powerful assembling strategy is based on using pre-targeted 2-periodical MOF layers as SBLs for deliberate building 3-periodical functional MOFs. This is carried out by chemical cross-linking layers through available bridging sites on layers (for example, open metal site or modified positions of organic spacer). SBL structural method requires reasonable choice of organic ligands, which will be pillars for layers. Definitely, this unique strategy is not confined to complicated pillar; it is also applied to simple pillar. As a result of countless combinations of cross-linkings, numerous MOFs with certain 3D network topologies can be pre-targeted developed and synthesized using preliminary obtained SBLs. Additional advantage of this approach is the fact that the common framework and topology of the network will remain unchanged basing on pillar layers, which provides almost unlimited expansion of the limited space (for example, cavities and porosity). In addition, it should be noted that if pores or windows of the layers remain unexpanded (i.e. expansion takes place only due to pillars), interpenetration of MOFs is excluded. No less important is the fact that modularity typical of this method provides easy functionalization or introduction of additional functionalities for deliberate applications.
SBL approach is used for deliberate synthesis of MOF with open massif of amide or amine functionalities inside the porous system. Two MOF platforms, eea-MOF and rtl-MOF based on pillaring kgm-a or sql-a layers with hetero-functional 3-connected organic building units, are directed and built deliberately for desired incorporation and demonstration of amide or amine functionalities [Fig. 11 and 12]. Distinctive kgm-a and sql-a networks can be considered as ideal blueprints for targeted 2-periodical MOFs based on assembling square building units produced from well-known metal paddlewheel clusters used as molecular building units.
3. Synthetic methodologies

3.1. Conventional synthesis

By now, quite many approaches to MOF synthesis have been developed, which are usually conventionally divided into two big groups: conventional synthesis and alternative methods of synthesis.\textsuperscript{120-123} The term conventional synthesis is applied to reactions performed without heating or using traditional electric heating without parallelization of reactions. Most MOF synthesis is carried out in liquid phase, in which separate solutions of metal salt and ligand are mixed together or a solvent is added to a solid salt and a ligand mixture in a reaction vessel. In order to obtain new MOFs, different factors should be taken into account during synthesis, such as nature of a metal ion, structural features of organic ligand, counterion, and some experimental variables including temperature of reaction, ratio of reacting agents and their concentrations, pH, system of solvents, crystallization methods, etc.\textsuperscript{124} Even small variations of only one from several factors having effect on the synthesis process can change dramatically structure of the formed MOFs.

Thus, if reaction between metal(II) acetates and a [1-(2-hydroxyphenyl)propyldiene]-hydrazide ligand is performed in 1:2 (M:L) molar ratio, only Mn(II) forms 1:2 complex in ethanol, while Cu(II) and Zn(II) form 1:1 1D CPs in methanol (Scheme 9). Interaction of ligand with Mn(II) acetate in methanol instead of formation of expected Mn(II) complex leads to heterocyclic derivative of quinazoline 2,2-dimethyl-3-[1-(2-hydroxy-phenyl)-propyldieneamino]-2,2-dimethyl-2,3-dihydro-1H-quinazolin-4-one.

Using interaction between 2-substituted 8-hydroxyquinolinolate ligand (HL) including Py group and Zn or Cd salts, CPs with [ZnL\textsubscript{2}]\textsubscript{00}, and [CdL\textsubscript{2}]\textsubscript{00} units are produced (Scheme 10).\textsuperscript{125} The first CP has 2D square network containing \textit{meso}-helical chains, while the second CP is a 2D network made of binuclear Zn\textsubscript{2} SBUs. By incorporation of two dicarboxylate anions into the reaction system, four CPs are obtained with [ZnL\textsubscript{2}L\textsubscript{2}L\textsubscript{2}L\textsubscript{2}L\textsubscript{2}]\textsubscript{00}, [ZnL\textsubscript{2}L\textsubscript{2}L\textsubscript{2}L\textsubscript{2}L\textsubscript{2}]\textsubscript{00}, [CdL\textsubscript{2}L\textsubscript{2}L\textsubscript{2}L\textsubscript{2}L\textsubscript{2}]\textsubscript{00}, and [CdL\textsubscript{2}L\textsubscript{2}L\textsubscript{2}L\textsubscript{2}L\textsubscript{2}]\textsubscript{00} units (H\textsubscript{2}L\textsubscript{00} = 1,4-benzenedicarboxylic acid, H\textsubscript{2}L\textsubscript{00} = 4,4’-biphenyldicarboxylic acid).

The L\textsubscript{1}-based CPs show 2D structures built by cyclic hexamers Zn\textsubscript{4}L\textsubscript{4}, which are divided in half by coordinated H\textsubscript{2}L\textsubscript{00}. In CPs based on L\textsubscript{00}, four ligands bridge 1D ML chains into 2D layered structure or present an interesting 3D structure in which L\textsubscript{00} ligands link binuclear Cd units into multiple \textit{meso}-helical chains along \textit{a} and \textit{b} axes, respectively. Therefore, skeletons of dicarboxylate anions play important role in assembling different MOF structures.

In the case of using organic acid, a base is needed, usually, amine, which is used to deprotonate the acid (future bridging fragment) and to initiate the reaction. At that, it is important to exclude competitive coordination of the base and organic acid.

3.1.1. Solvent evaporation method. A solvent evaporation is most conventional and widely used method of crystal growth which carried out by evaporation or cooling a saturated solution. For this method, the following conditions are needed: crystals grow in saturated solutions, solubility increases as temperature rises, and crystals can appear during cooling.
which should be well controlled, since the cooling rate and final temperature are interrelated. Its main drawback is long duration of the process as compared to other known ordinary methods.

As an example of a simple water-based method of slow evaporation, we shall consider synthesis of Pb-based CP with \([\text{PbCl_2(HL)}]_n\) unit, where \(\text{H}_2\text{L} = \text{pyrazine-2,3-dicarboxylic acid},\) under optimized conditions.\(^\text{127}\) In the same way, the CP with \([\text{Pb}_2\text{L}_2]_n\) unit is obtained without any additives. Reactions between \(\text{Zr(n)}\) salt and \(\text{Pp-tetracarboxylic acid}\) lead to MOF with two types of open channels presenting MOF distinguished by \((4,8)\)-connected \(\text{sqc}\) network. It is interesting to notice that MOF remains unchanged in boiling water and in aqueous solutions with pH changing from 1 to 11, surprisingly wide range, which MOF can endure.\(^\text{128}\) 3D heterobimetallic CP with \([\text{Mn}_2\text{L}_2\text{Hg(SCN)}]_n\) asymmetric unit (\(\text{L} = 2\text{-benzoylpyridine}\)) is synthesized by slow addition of \(\text{NH}_3\text{SCN}\) and \(\text{Hg(SCN)}_2\) aqueous solution to methanol solution of Mn chloride and L followed by mixing at room temperature during 12 h.\(^\text{129}\)

The final dimensionality of crystal structure depends on such important factor as \(\text{M–L}\) ratio. It is known that 1 : 1 ratio often leads to 1D structures, and high coefficients, as a rule, advance CP of higher dimensionality.\(^\text{130}\) Not only covalence has relation to these structures: bridging action of non-covalent interactions such as H-bonds or π–π stacking can change final structures even with very similar components. This result depends strongly on reactivity of involved types and H-bonded donor/acceptor interactions.

Reactions of alkyl substituted aromatic monocarboxylic acids (HL) with transition metal ions in presence of chelating and bridging N,N-donor ligands (L’) give either discrete or polymer complexes depending on position of a substituent on benzoic acid and ability of ancillary ligand to work as chelating or bridging ligand.\(^\text{131}\) For example, when phen was used as an ancillary ligand, and for carboxylic acid was taken 4-tert-butybenzoic acid, discrete complexes were deposited. However, for the same ancillary ligand with 2,4,6-trimethylbenzoic acid, 1D helical polymers with \([\text{ML}_2(L’)]_n\) unit (\(\text{M} = \text{Mn, Co, Cu and Zn}\)) were obtained. This clearly shows ability of \(\text{o,o}-\text{disubstituted benzoic acids}\) to form helical polymers, while \(p\)-substitution on aryl ring leads to discrete complexes.

Ancillary ligands, such as bpy and phen, are often used with multicarboxylate ligands for building new functional CP architectures with chelated units.\(^\text{132}\) Phen as an ancillary ligand is most preferable since it generates 3D supramolecular interactions, such as aromatic π–π stacking and –CH–π interaction stacking, which provides stability of supramolecular ensembles.\(^\text{133–135}\) In this connection, reasonable combination of bridging carboxylates with phen-like chelating ligands and metal ions has generated many interesting coordinated architectures. As length of a linear spacer increases, a tendency to interpenetration of 3D CP also increases, and if a structure is already penetrated, a degree of interpenetration, as a rule, increases with length of a spacer ligand.

Counter-ions are present in CP structure, when neutral ligands are used. They can have effect on geometrical environment of a metal ion (more or less coordinating counter-ions), and can control generally CP structure by involving into weak interactions or acting as guest molecules in voids of a crystal architecture. As a typical example, we shall consider assembling tripyridyltriazole ligand 3,4-bis(2-pyridyl)-5-(3-pyridyl)1,2,4-triazole (L) with dca anion, where dca is dicyanamide, and different Cd(n) salts in \(\text{CH}_3\text{OH–H}_2\text{O}\) medium, which leads to CPs with \([\text{Cd}_2\text{L}_2\text{dca(H}_2\text{O})]_n\), \([\text{CdL(dca)}\text{ClO}_4]_n\) and \([\text{CdL(dca)}\text{X}_n\text{]}(\text{X}^- = \text{Cl}^–\text{and Br}^–)\) units.\(^\text{136}\) In the first two CPs, Cd(n) centers are linked by L spacers forming \([\text{Cd}_2\text{L}_2]\) bimetal units, which are bridged with dca anions, thus leading to 1D motif and 2D layer, respectively. The latter two CPs reflect 3D isostructural

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**Scheme 10** Synthesis of CPs based on 2-substituted 8-hydroxyquinolinate ligand including Py group, where \(\text{H}_2\text{BDC}\) is \(1.4\)-benzenedicarboxylic acid and \(\text{H}_2\text{BPDC}\) is \(4.4\)-biphenylidicarboxylic acid. [Reproduced from ref. 126 with permission from The Royal Society of Chemistry.]
homochiral coordination networks built by 1D [CdL]_n helical units and dca bridges. Therefore, Cd(II) salt counter-ions are responsible for different CP structures. We shall also notice that assemblies of the same ligand with Cu(II) chloride and dca anion at different M–L ratios give CPs with [Cu_2L(dca)]_n and [CuL(dca)Cl]_n units displaying 1D double-chained massif and 3D chiral dia network, respectively (Scheme 11).

Depending on nature of anion of the initial metal salt Cu(NO_3)_2 or Cu(ClO_4)_2, mixing their aqueous solutions with methanol L solution [L = 3-(2-pyridyl)-4-(4-pyridyl)1,2,4-triazole] leads either to a mononuclear complex [CuL(H_2O)][NO_3]_2, or to 1D linear CP with [CuL]_n unit, respectively. It is interesting that incorporation of NaN_3 in these assembled systems under similar conditions gives two different 2D layered networks with [CuL(N_3)]_n units, both of which contain double inorganic anions as counter-ions (NO_3^-/N_3^- and ClO_4^-/N_3^-, respectively). In addition, 1D linear massif [CuL(N_3)]_n is obtained with N_3^- compensating positive charges with subtle change in reaction conditions. It should be noted that stepwise structural transformations between these crystal phases can also be achieved by direction of azide anion (Scheme 12). In other example, Cd(II) CPs with [Cd(L)(H_2O)Cl]_n, [Cd(L)(H_2O)Br]_n and [CdLBr]_2 units are assembled from CdX_2 (X = Cl, Br, I) and the same ligand. The first two CPs are isostructural and show 1D loop-like chain, while the third complex has prominent dimeric macrocyclic motif. It is interesting that another 1D chain [CdL(SCN)]_n can be reached when NH_4SCN is incorporated in the assembled system of third complex. Thus, halogenide- and thiocyanate-anions in these coordination complexes act not only as counter-ions, but as structure-directing agents.

We shall also notice 3,3'- di(pyrazinamoyl)-bpy (H_2L) ligand consisting of different linking units, which can facilitate CP formation. Reaction of the ligand with Cu acetate in presence of Et_3N·HCl gives CP with [Cu_2L_2Cl_2(OAc)]_n unit, at that trans-configuration of bpy-unit presented in the ligand remains in the resulting CP. In other words, a bpy fragment is not chelating,
and \( {\text{L}}^{2-} \) anion coordinates two Cu ions by bis-tridentate mode through amide and pyridine nitrogen atoms (Fig. 13). Bimetallic unit is linked to equivalent unit through a pair of 1,3-acetate and \( \mu \)-chlorine bridges with resulting tetrameric \( [\text{Cu}_{4}\text{L}_{2}\text{Cl}_{2}(\text{OAc})_{2}] \) core. Coordination of one of pyrazole substituents from each ligand to the third Cu center of rigid paddle-wheel \( [\text{Cu}_{3}(\text{OAc})_{4}] \) unit leads to unique 1D polymer structure.

As the chosen solvents change, the reactions between 2,5-thiophenedicarboxylic acid (\( \text{H}_{2}\text{L} \)) and Mg nitrate give four different CPs \( \text{Mg}_{2}\text{L}_{2}(\text{SO}_{4})_{2} \cdot 0.5\text{Sol} \), \( \text{Mgl}(\text{DMSO}) \), \( [\text{Mg}_{2}\text{L}_{2}(\text{Ac})] \) and \( [\text{Mg}_{2}\text{L}_{2}(\text{DMF})_{2}(\text{EtOH})(\text{H}_{2}\text{O})_{2}] \) (\( \text{Sol} = \text{ethylene glycol} \)).\(^{140}\) Coordinated molecules of solvents take different coordination regimes, which play important role in building CP structure. Thus, in the first CP, the Sol molecules as bidentate bridging ligands advance linking MgL layers in 3D framework. DMSO molecules in the second CP and \( \text{Ac}^- \) anions in the third CP are coordinated in \( \mu_{2-}\text{way} \) and \( (k^2-k')_{13} \) regime, respectively, leading to infinite chains as SBU. And, finally, in layered 2D structure of the fourth CP, coordination of DMF, EtOH and \( \text{H}_{2}\text{O} \) molecules provides induction of the non-centrosymmetrical structure.

There is interest in fast, inexpensive, and ecological methods of production of high quality MOFs with properties rarely obtained under normal conditions.\(^{141}\) The method is based on using organic salts (instead of their homological protonated organic ligands) as sources of anion spacers, so that their solubility and necessary deprotonation stage are achieved substantially in aqueous solution. For example, high quality MOF-74 based on zinc acetate and 2,5-dihydroterephthalic acid usually obtained in organic solvents can also be synthesized in water and at room temperature.

We shall also notice (green coordination modulation method) for CP synthesis using ecological ethanol and alkali additive.\(^{142}\) In particular, most variable architectures, including rod crystals, dual-core shelled peanut-like crystals and double-head dandelion-like crystals, are obtained through changes in parameters of the reaction.

For synthesis of phase-pure MOFs, an easy seed-mediated approach is developed, which allows to skipping the MOF nucleation stage, which generates mixed cores.\(^{143}\) In addition, using this method, phase-pure MOF isomers with separate porous structures are obtained, thus showing universality of the method.

3.1.2. Diffusion methods. Diffusion method is one of classic methods for synthesis of well-defined crystalline materials. The most often used procedure is liquid–liquid diffusion, in which two or more solutions containing solved reagents are used. As compared with fast mixing solutions containing metal ions and bridging ligands, which makes it possible to obtain microcrystalline product, slow diffusion leads to formation of crystals with pronounced morphologies.

Of interest is a production of 3D plywood-like Ni\([\text{II}]\) hexaazamacrocyclic CP with [Ni\([\text{II}]\text{L}_{4}\text{H}_{4}\text{L}'\)] \( \cdot \text{Sol}_x \) unit \( (L = 1,3,6,9,11,14\)-hexaazatricycl[12.2.1.16,9]octadecane, \( \text{H}_{4}\text{L}' = 1,2,4,5\)-benzenetetracarboxylic acid) using slow diffusion.\(^{144}\) The CP structure includes [Ni\([\text{II}]\text{L}_{4}^{0+}\)] cation and [\( \text{H}_{4}\text{L}'\text{L}_{4}^{0+}\)] anion in molar ratio 1 : 1, in which Ni\([\text{II}]\) ions are connected through bridging ligand \( \text{H}_{4}\text{L}'\text{L}_{4}^{0+} \) based on weak Ni-O coordination interactions with 1D chain formation. Alternating cross-like 1D chains are stacked in the plywood structure and connected with each other through H-bonds, giving 3D network (Fig. 14).

There is such interesting synthetic method as vapor diffusion, which includes solution of a metal source and a ligand in a solvent, for example, DMF, and its positioning in an open container surrounded with a volatile base solution, such as...
thermal methods are the most efficient way for production of various CPs. Initially, these methods were developed for synthesis of zeolites, but then they were adapted to synthesis of MOFs and occur so efficient that by now the majority of CPs is obtained particularly using these methods. Hydro(solvo)thermal methods are widely applied in synthesis of new CPs, since their products have high purity, good dispersion, are easily controlled, etc.\textsuperscript{126-128} The hydro(solvo)thermal reactions are associated with using organic or inorganic solvent at high temperature (usually 80–260 °C) and autogeneous pressure in a closed system (as a rule, teflon-lined autoclaves or glass vessels).\textsuperscript{14} An obvious advantage of hydro(solvo)thermal methods is that they make it possible to alleviate the solubility problems of chelating ligands and improve the reactivity of the reagents during crystallization. Undoubtedly, during the synthesis it is necessary to take into account some parameters of the reaction, including composition of reagents, temperature and pressure, concentration, time of reaction, pH, solubility, etc.\textsuperscript{149} Drawbacks of this method are long time of reaction, low percentage yield, necessity in high temperatures and use of toxic solvents (for example, DMF).\textsuperscript{128}

Among a vast variety of polytopic ligands used in hydro(solvo)thermal MOF synthesis, the leading role, undoubtedly belongs to carboxylate ligands,\textsuperscript{14} therefore they, first of all, were used in design of CPs with chelated units. Thus, solvothermal reaction of CO\textsubscript{2} or Zn\textsubscript{2} salts with dianhydride of 1,2,4,5-tetracarboxylic acid (H\textsubscript{4}A) leads to formation of 3D CPs with [CO\textsubscript{2}L\textsubscript{2}]\textsubscript{m} and [Zn\textsubscript{2}L\textsubscript{2}]\textsubscript{m} units. In the first polymer, 1D CO\textsubscript{2} carboxylate chains are interconnected by organic fragment of L\textsubscript{2} \textsuperscript{14} ligands. In the second polymer, 1D ZnL\textsubscript{2} ribbons are interconnected by Zn\textsubscript{2}(H\textsubscript{2}O)\textsubscript{4} building units creating 3D porous framework with 1D rectangular channels.\textsuperscript{150} Often in similar synthesis, ancillary ligands such as 1,2-di(4-pyridyl)ethylene (L\textsubscript{2}) are also used apart from the main ones. It is with this ligand, CP with [Zn\textsubscript{2}L\textsubscript{2}(L\textsubscript{2})\textsubscript{3}]\textsubscript{m} unit is obtained, which has a complicated 3D framework with (3,4)-connected \textbf{fsh} topology.\textsuperscript{151} An interesting example is 3D CP with [Cd\textsubscript{2}(L\textsubscript{2})\textsubscript{3}]\textsubscript{m} unit synthesized based on a main ligand of 1,2,4-benzenetricarboxylic acid (H\textsubscript{3}A) and ancillary ligand of 1,4-bis[imidazolyl]butane (L\textsubscript{4}) under hydrothermal conditions. The obtained CP relates to extraordinary examples of trinodal (4,6)-connected network with \textbf{fsc} topology.\textsuperscript{152} It is worth paying attention to CP with [Mg\textsubscript{2}L\textsubscript{2}]\textsubscript{m} (L = perylene-3,4,9,10-tetracarboxylic acid dianhydride) with special hexagonal tube-like morphology, which is produced by self-assembly by hydrothermal method.\textsuperscript{153} It is interesting that this morphology of [Mg\textsubscript{2}L\textsubscript{2}] is mostly controlled by Ostwald ripening and self-structuring mechanism, at that using organic solvents as additives, the preparation of CPs of different morphologies, for example, hexagonal rings and snowflakes, and also CP sizes from micrometer to nanometers can be controlled.

Hydrothermal reaction of La(NO\textsubscript{3})\textsubscript{3} with 2-pyrimidinecarbonitrile in water gives 3D MOF with [LaL\textsubscript{3}(H\textsubscript{2}L\textsubscript{4})\textsubscript{2}]\textsubscript{m} unit (L = oxalate), which has channels with trapped water molecules.\textsuperscript{154} It is important that CP is built from La atoms linking with bis-chelating oxalate ligands.

La-based CPs microplates are prepared using a simple (bottom-up) method by hydrothermal reaction of La(III) and \textit{green ligand} ethylenediaminetetraacetic acid (EDTA).\textsuperscript{155} Under similar conditions, analogous Ce-EDTA CPs microplates are also prepared. It is important that the mechanism of La-EDTA CPs formation follows the Ostwald ripening mechanism, and its morphology can be easily adjusted by variation of the amount of EDTA and the total concentration of reagents. In addition, also several types of micro/nanostructures have been obtained, such as nanocluster particles, nanofibers, big or uniform microplates.

MOF with [CdL(phen)]\textsubscript{m} unit is hydrothermally synthesized using heterocyclic 2,2′-biquinoline-4,4′-dicarboxylate ligand (L) and N-containing ancillary ligand.\textsuperscript{156} The complex is infinite 2D framework networks containing aesthetic \textit{meso}-helix, which are linked with formation of 6-connected 3D supramolecular architecture through aromatic π–π stacking and C–H···O–H-bonds interactions.

CPs with [ZnL(L\textsubscript{2})\textsubscript{m}][ZnL(L\textsubscript{2})\textsubscript{m}], [Cd\textsubscript{2}L(L\textsubscript{2})\textsubscript{m}]\textsubscript{m} and [Cd(L\textsubscript{2})\textsubscript{m}][Cd(L\textsubscript{2})\textsubscript{m}]\textsubscript{m} units are obtained through interaction under solvothermal conditions between 1,2-phenylenedioxydiacetic acid (H\textsubscript{2}A) and hydroquinone-O,O′-diacetic acid (H\textsubscript{2}B) with Zn and Cd nitrates in presence of ancillary ligand 1,1′-butanediylbis[imidazole] (L\textsubscript{1}).\textsuperscript{157} It should be also noted hydro(solvo)thermal synthesized CPs with [Cd(phen)L\textsubscript{2}]\textsubscript{m} ([Cd(phen)[L\textsubscript{2}]\textsubscript{m}]\textsubscript{m} and [Cd(phen)L\textsubscript{2}]\textsubscript{m}) units based on 2-methyl-(H\textsubscript{2}A) and 2-ethyl-1H-imidazole-4,5-dicarboxylic acid (H\textsubscript{2}B)\textsubscript{158} The CPs are infinite chains, in which HL and HL′ in μ\textsubscript{2}-regime are linked with Cd(II) atoms, while phen or bpy ligands chelate the central ions (Fig. 15).

An interesting example is hydro(solvo)thermally synthesized CPs with [Mn\textsubscript{2}L\textsubscript{2}(phen)]\textsubscript{m}, [Mn\textsubscript{2}L\textsubscript{2}(phen)]\textsubscript{m} and [Mn\textsubscript{2}L\textsubscript{2}(phen)]\textsubscript{m} units based on 1,4-naphthalene dicarboxylic acid (H\textsubscript{2}A)\textsubscript{159} 1D–2D ligands take various coordination regimes in different solvents and concentrations, which provides formation of different crystal structures. Thus, first CP has Mn\textsubscript{2} dimers linked with L\textsubscript{2} spacers, and stacked in 2D structures of grid pattern. The next CP has 3D framework, which is formed by Mn\textsubscript{2} dimers and L\textsuperscript{2} spacers, and each MnO\textsubscript{2}N\textsubscript{2} node of the third CP is linked with other node through L\textsuperscript{2} ligands with formation of 2D structure (Scheme 13).
It should be also noted synthesized under hydrothermal conditions CPs with \([\text{CdL}(\text{phen})]\) (1), \([\text{NiL}(\text{phen})]\) (2), \([\text{CdL}_3(\text{phen})]\) (3), \([\text{CdL}(\text{phen})]\) (4), \([\text{NiL}(\text{phen})]\) (5), \([\text{NiL}(\text{phen})]\) (6), \([\text{CdL}(\text{phen})]\) (7), \([\text{CdL}(\text{phen})]\) (8), \([\text{CdL}_3(\text{H}_2\text{O})]\) (9), and \([\text{CdL}_3(\text{phen})]\) (10) units including the main ligand 3-carboxy-1-(4′-(2′-carboxy)biphenylmethyl)-2-oxidopyridinium (H₂L) and ancillary (L′) ligands 1,4-bis(imidazol-1-ylmethyl)benzene (compounds 1 and 2), 2-(6′-(pyridin-2′-yl)-4′-p-tolylpyridin-20-yl)pyridine (compounds 3 and 5), 2-(4′-(4′-tert-butylyphenyl)-6′-(pyridin-2′-yl)pyridin-20-yl)pyridine (compounds 4 and 6), 1,10-(1,6-hexanediyl)bis(imidazole) (compound 7), 1,10-(1,4-butanediyl)bis(imidazole) (compound 10). It is important that the crystal structures of the obtained CPs are influenced to a great extent by L-anions and N-donor ligands.

Solvothermal reactions of terphenyl-2,5,2′,5′,2″-tetracarboxylic acid (H₄L) and M(n) in presence of bpy give three CPs with \([\text{ML}_n(\text{bpy})]\) (M = Co, Ni) and \([\text{Cu(H}_2\text{L})(\text{bpy})]\) n units. Of interest are systematic arrays of CPs including bpy chelating ligand and different derivatives of isophthalic acid such as 5-(4′-methylphenyl) group (1D linear chain linked with L bridge). OH group (1D zigzag chains linked by H-bonding and π-π stacking interactions to form a 3D supramolecular structure), 5-(4-carboxybenzylamino) group (1D, 2D and 3D frameworks), 5-(imidazol-1-yl-methyl) group (double-chained and 3D supramolecular structures through H-bond/π-π interactions), 4,5-di(3′-carboxyphenyl) group (0D, 1D, 2D and 3D structures) etc. Such arrays make it possible to reveal fundamental structure–property relations to define possible directions of practical applications for CPs.

It should be also noted using 3,5-disulfobenzoate (L) in hydrothermal synthesis of Ln-based 2D MOFs with \([\text{LnL}(\text{phen})]\) n unit, where Ln = La, Pr, Nd. They belong to two 2D structural types, and their networks have different topologies: the isostructural La and Pr compounds are uninodal 5-connected CP 2-periodical (6,3) networks and the Nd compound has a binodal 3- and 6-connected kgd network.

It is interesting the using flexible fatty carboxylic acids, for example heptane and hexane diacids, as bridging ligands, and phen derivatives, for example, 2-(3-pyridyl)imidazo[4,5-f]phen, as chelating ligands. Macrocyclic ML complexes based on 2,3-dioxo-5,6,15,15-dibenzo-1,4,8,12-tetraazaacyclo-pentadeca-7,13-diene (H₂L) are applied as main building units for preparation of CPs with chelated units is linking monomer metal chelates with various bridging ligands. Thus, CP with \([\text{Co}_2(\text{H}_4\text{L})_2(\text{phen})]_{\text{n}}(\text{N}_2)_n\) unit has been obtained by hydrothermal method using a phen as ancillary chelating ligand and bridging azide-linking ligand. Macroyclic ML complexes based on 2,3-dioxo-5,6,15,15-dibenzo-1,4,8,12-tetraazaacyclo-pentadeca-7,13-diene (H₂L) are applied as main building units for preparation of CPs with \([\text{CdL}(\text{phen})]\) n, and \([\text{MnL}(\text{phen})]\) n units including bridging ligands 5-aminoisophthalic acid (H₅L) or 5-aminoisophthalic acid (H₅L'){. It is important that CPs consist of different 1D chains formed by L and L′ or L″ bridge, and they are bound through H-bonds into 2D architecture.

The 2D CPs with \([\text{ZnL}(\text{phen})]\) n, \([\text{ CdL}(\text{phen})]\) n, \([\text{ZnL'}(\text{phen})]\) n, and \([\text{ZnL''}(\text{phen})]\) n units are obtained as a result of reaction of Zn(n)- or Cd(n)-phen with biphenyl-4,4′-dicarboxylic acid (H₄L) and 4,4′-azidobenzoic acid (H₄L′). The coordination-induced effect of their morphology is shown; in
particular, grain and flowery morphology of non-coordinated ligands H$_2$L and H$_2$L', respectively, changes with coordination of metal salts. The ridged surface, cracks of 2 µm width, parallelepiped structure and feathery appearance are established for the CPs, respectively (Scheme 14).

At the same time, when using bpy-4,4'-dicarboxylate (H$_2$L) and the respective transition metal[n] salt under hydrothermal conditions, CPs with [MnL(H$_2$O)$_2$]$_n$, [NiL(H$_2$O)$_2$]-2H$_2$O]$_n$, [NiL(H$_2$O)$_3$]$_n$ and [CuL(H$_2$O)$_2$]$_n$ units are obtained, in which each metal ion is coordinated with chelating bpy fragment, two carboxylate oxygen L$^2$– and water molecules.\(^1\)

There is interest in other 3d–4f heterometallic CPs with [Ln$_2$CO$_2$L$_4$(HL)$_2$(H$_2$O)$_2$]$_n$ (Ln = Eu, Gd, Tb, Dy, Ho, Tm, Lu) unit shaped as 3D sandwich-like framework.\(^2\)

Heterocyclic polycarboxylates are widely used in synthesis of CPs with chelated units.\(^3\) Thus, hydrothermally synthesized are 2D lanthanide CPs with [LnL(HL)$_n$](Ln = Pr, Nd, Sm, Eu and Tb, H$_2$L = pyridine-2,3-dicarboxylic acid) unit, which are built from sheets consisting of carboxylate-bridging Ln dimers folded along [100] direction and held together by H-bonds between carboxylic groups.\(^4\) Using 1,4-bis(1H-imidazol-1-yl) butane (L') as an ancillary ligand, CPs with [ZnL(L')]$_n$ and [CdL(L')]$_n$ units are obtained, first CP of which shows wave-like layered structure, where layers are additionally stacked through π–π interactions to generate 3D supramolecular architecture (Fig. 16).\(^5\) For the second CP, 3D three-fold interpenetrating diamondoid architectures are typical.

It should be also noted the using pyridine-2,5-dicarboxylic,\(^6\) pyridine-2,4-dicarboxylic,\(^7\) and pyridine-2,6-dicarboxylic\(^8\) acids in the design of CPs.

Under hydrothermal conditions from Co(OH)$_2$ and chelidamic acid in the molar ratio 1 : 1, the Co(n) CP with [Co$_{13}$L]$_n$ unit is obtained, which crystallizes into 1D stair-like structure.\(^9\)

There is interest in CP with [Ca$_2$L(L')]$_n$ unit based on pyrazine-2-carboxylic (L) and benzene-1,3,5-tricarboxylate (L') synthesized by hydro(solvo)thermal method forming a double layered network, in which inorganic zigzag chains of coordination Ca polyhedrons are bound with organic ligands.\(^10\) CPs with [M$_2$L$_4$]$_n$ unit are obtained through interaction between 2,3-pyrazinedicarboxylic acid (H$_2$L) and Y and Sm nitrates under hydrothermal conditions in presence of ammonia for pH regulation. The obtained CPs are isostructural and show a 3D network structure based on [M$_2$L$_4$(H$_2$O)$_2$]$_n$ building units.\(^11\)

Different synthetic strategies are used in hydro(solvo)thermal synthesis of CPs based on the different imidazole-dicarboxylic acids: imidazole-4,5-dicarboxylic,\(^12\) 2-p-methoxyphenyl-1H-imidazole-4,5-dicarboxylic,\(^13\) 1-(4-carboxybenzyl)-1H-imidazole-4,5-dicarboxylic,\(^14\) and 2-(pyridine-3-yl)-1H-imidazole-4,5-dicarboxylic\(^15\) acids.

CPs with [Pb$_2$L]$_n$ and [MnL]$_n$ units based on imidazo[4,5-f] phen (L) are synthesized using hydrothermal method.\(^16\) In the structure of the first polymer, a Pb(n) center is six-coordinated and has octahedral geometry. In the second CP, Mn(n) ion is six-coordinated with formation of distorted octahedral configuration, and the polymer shows 1D supramolecular chain formed via H-bonds.

Hydro(solvo)thermal method proves to be very efficient for production of chiral CPs, which are of great interest not only due to intriguing variety of architectures and topologies, but

Scheme 14  CPs obtained by the reaction of metal [Zn(n), Cd(n)], phen separately with biphenyl-4,4'-dicarboxylic acid and 4,4'-azodibenzoic acid. [Reprinted with permission from ref. 170. Copyright 2012, American Chemical Society.]
because of their potential applications in many areas. It is important that chiral CPs can be developed controllably by targeted choice of ligands or metal centers. Among the developed by new methods of designing chiral materials, it is worth noticing incorporation of chiral ligands, chiral templates, or chiral ancillary agents, and also spontaneous resolution without chiral auxiliary. From this view, the most rational method of design of chiral CPs is integration of a chiral center into the structure, such as a chiral ligand; however, this process is not always convenient because of difficulty of a chiral ligand synthesis. In addition, more attention is attracted to use of achiral ligands for building various chiral frameworks, especially by different conformed rotations of a ligand skeleton or self-assembling achiral building units. Principally important strategy for structuring chiral materials is development of left-hand and right-hand helical structures, which requires efficient transfer of stereochemical information between neighboring chiral helices. Considerable contribution to transfer of chirality between helixes can be made using supramolecular interactions including M−L bonds and non-covalent interactions, such as H-bonds and π−π stacking interactions. Therefore, it is promising to use associated functionalities providing supramolecular interactions and performing efficient chiral transfer along whole supramolecular assembly. At the same time, enantioselective synthesis is straight and efficient method of synthesis of enantiopure chiral-open MOF using enantiopure building units as precursors of reagents. Therefore, many chiral-open frameworks with high thermal stability are obtained using enantioselective synthesis and are successfully used in enantioselective catalysis and separation. It should be noted that chirality is a unique property of CPs with chelated units when tris-chelate is used.

In order to integrate inherent chirality of a ligand and chirality induced by conformation rotations of asymmetric skeleton of the ligand during its coordination, CPs with [CoL(Py)$_2$]$_n$, [NiL(Py)$_2$]$_n$, [CuL(Py)$_2$]$_n$, [Zn$_2$L$_2$(Py)$_2$]$_n$ and [Cd$_2$L$_2$- (Py)$_2$]$_n$ units are obtained through the reaction between (1R,3S)-1,2,2-trimethyl-3-[[5-(2-hydroxy)phenyl-1,3,4-oxadiazol-2-yl)] cyclopentanecarboxylic acid (H$_2$L) and transition metal ions. Among them, the first two CPs had only right-hand helical chains with homochiral helical discrimination due to formation of interhelical non-classic H-bonds C–H⋯O. Other polymers contained simultaneously alternating right- and left-handed helical chains, interaction between which was due to π−π interactions (CP with Cu atoms) and interchain bonds M–O–M (CPs with Zn and Cd atoms) (Scheme 15). In these compounds, chirality is stipulated by inherent chirality of a ligand and conformation twisting the main asymmetric chain of a ligand. Therefore, the effect of metal ion nature on structure of a chiral CP is obvious.

Using solvothermal method with Cd[n] salt and enantiopure chiral ligand based on d-isosorbide (L), a homochiral luminescent porous CP with [CdL(H$_2$O)]$_n$ unit has been synthesized, which is 2D porous material and forms 1D channel along a axis with the size ~6.2 × 4.4 Å$^2$. A chiral CP with [AgSCN(phen)]$_n$ unit obtained using solvothermal reaction between AgSCN and phen shows unusual asymmetric 1D network formed from a zigzag [Ag–S–Ag]$_n$ chain with thiocyanate ions and phen ligands hanging along opposite sides of the chain, respectively (Fig. 17a). It is important that

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**Fig. 16** (a) Coordination environment of the Zn(ii) atom in CP. (b) View of the 2D undulating layer of CP. (c) View of the 3D supramolecular architecture connected by face-to-face π–π interactions. [Reprinted from ref. 174. Copyright 2013, with permission from Elsevier.]
asymmetry of the chain is clear in $b$ direction (Fig. 17b), and in one unit cell there are two [AgSCN(phen)$_n$]$_n$ chains stacked in opposite direction (Fig. 17c). It is interesting to notice that the scheme of the unit cell along $b$ axis is very similar to Chinese “Tai Ji” patterning, which means two opposite principles of nature (Fig. 17d).

Also it is worth noticing homochiral CPs with [ML$_n$$(H_2L)$]$_n$ unit, where $M = Zn$ or Co, based on $\nu(+)$ camphoric acid (L) and 4,4'-methylenebis(3,5-dimethylpyrazole) (H$_2$L) obtained through interaction between ligands and transition metal ions under hydrothermal conditions.$^{205}$

Of interest is chiral infrequent azolate-based octanuclear metallomacrocyclic 3D [Co$_4$L$_2$(H$_2$O)$_2$]$_n$ complex based on 5'- (pyridin-2-yl)-2$H$,4'$H$-3,3'-bi(1,2,4-triazole) (H$_2$L) (Fig. 18). It is important that asymmetric polymer unit consists of one-half Co1 and Co3 ions and one independent Co2 ion, two L$_2^+$ anions, two aqua-ligands, and four and half lattice water molecules, at that each Co$_{1-3}$ ion is six-coordinated and takes slightly distorted octahedral geometry.

Solvothermal synthesis is successful approach to obtaining a broad series of 2D and 3D Pp MOFs.$^{63,207-211}$ For this purpose, most often spacers based on Pp with carboxylic acid and Py-linked functional groups are used, in particular, meso-tetra(4-carboxyphenyl)-Pp and meso-tetra(4-pyridyl)-Pp.$^{22,79,82}$ During building MOFs based on Pp spacer, macrocycles can be metalized in situ with the same metal ions, which present in SBU's,$^{58,59,61}$ that is attractive approach for development of high density metallic sites in 3D nanospace, and can be especially enticing for catalysis, if metal centers inside Pp rings, and in SBU's are catalytically active.$^{45,61}$

As a typical example, it should be noted solvothermal reaction of Fe(II) chloride with the tetrakis(4-carboxyphenyl)-Pp in the presence of different bases, which results in the formation of four Fe-Pp MOFs representing three different topologies and inorganic SBU (Scheme 16).$^{212}$ In particular, depending on the synthesis conditions, isolated Fe(II) octahedra, diiron(II) paddlewheel dimers or extended [Fe(III)OH)$_3O_4$]$_n$ chains can be obtained controllably.

Of interest are solvothermally synthesized rare indium-based porous MMPF-7 and MMPF-8 with the pts topology constructed by self-assembly of In(III) and two custom-designed meso-tetra(4-carboxyphenyl)-Pp.$^{213}$

Among other Pp ligands for CP synthesis, there is interest in octatopic Pp ligand tetrakis(3,5-bis([4-carboxyphenyl]phenyl)-Pp, which forms {pillar-free} highly porous M-Pp framework [[Zn$_2$(H$_2$O)$_2$]$_2$·[ZnL]$_2$]$_n$ called UNLPF-1 in the solvothermal reaction of L with Zn nitrate in DMF and acetic acid at 80 °C during 72 h.$^{209}$ UNLPF-1 has 3D non-interpenetrating structure with widespread type of SBU, in particular, square paddlewheel [Zn$_3$(COO)$_4$(H$_2$O)$_2$], which connects four L ligands,
and each L ligand links eight in situ generated (four above and four below Pp plane) SBUs (Fig. 19). It is important that two adjacent Pp macrocycles along with four paddlewheel SBUs form a cavity of great sizes $14.5 \times 23.7 \, \text{Å}^2$. In addition, the same ligand is linked to in situ generated a distorted Co trigonal prism SBU, producing strong, (6, 8, 8)-connected MMPF-2 with msq topology.\textsuperscript{238}

During hydrothermal synthesis of Cd-Pp CPs based on meso-tetra-(pyridyl)-Pp, the Pp molecule shows three types of coordination regimes, in which each Pp molecule is linked to one, two, four, and five Cd centers.\textsuperscript{214}

Another attractive ligand 5,10,15,20-tetrakis(4,4-dipyridylamino-phenylene)-Pp can be coordinated with 4–7 metal centers using its multiple peripheral Py and a Pp core leading to rich structural variety: H-bonding 1D chains linked by $[(\text{H}_2\text{O})\text{Cl}_2]^2$ fragments, 3D structure formed by $\pi-\pi$ stacking interactions between interpenetrating 2D networks, 2D structure with big cavities consisting of 50- and 70-members metal-lomacrocycles (Fig. 20), complex 2D structure linked with zigzag chains, and stair-like 2D structure containing binuclear $[\text{Cd}_2(\text{CO}_3)_4]$ subunits. It is important that 4,4'-dipyridylaminophenylene fragments can rotate around Pp framework, leading to good conformation flexibility of a Pp ligand.\textsuperscript{215}

Coordination of Mn(II) and Zn(II) acetates with 5,15-bis(4-carboxyphenyl)-10,20-dipyridyl-Pp ($\text{H}_4\text{L}$) gives two CPs with $[\text{Mn}(\text{HL})]_n$ and $[\text{Zn}(\text{HL})]_n$ units, respectively.\textsuperscript{216} First CP has 2D coordination networks, which are then bound by H-bonds with formation of a 3D network structure. In the second polymer, 1D coordination zigzag chains are obtained, and are then linked by H-bonds into a 2D structure (Scheme 17). It is important that carboxyl residuals in the CPs are noncoordinated and the Py moieties are either coordinated or noncoordinated, at that noncoordinated carboxyl and Py fragments take part in intermolecular H-bonds, which are favorable for formation of H-bonding networks.

There is a great practical interest in using a solvothermal approach for production of uniform thin films based on Zr-Pp
In particular, the fabricated MOF-525 thin film is electrochemically targeted in aqueous solution and can be used as an amperometric nitrite sensor. It should be noted the microcrystalline structure containing up to 10 different types of bivalent metals, namely, Mg, Ca, Sr, Ba, Mn, Fe, Co, Ni, Zn, and Cd, based on MOF-74 with \([\text{Zn}_2\text{L}]_n\) unit (L = dioxidoterephthalate), which has high CO\(_2\) adsorption capacity under practical atmospheric pressure conditions. The one-pot reaction makes it possible to obtain MOF-74 with 2 (Mg and Co), 4 (Mg, Co, Ni, and Zn), 6 (Mg, Sr, Mn, Co, Ni, and Zn), 8 (Mg, Ca, Sr, Mn, Fe, Co, Ni, and Zn), and 10 (Mg, Ca, Sr, Ba, Mn, Fe, Co, Ni, Zn, and Cd) different types of bivalent metals. It turned out that all metal ions used in MOF synthesis are integrated in the same MOF-74 structure, and metal ions are heterogeneously distributed within each of crystalline particles. It should be noted that this approach is also used for incorporation of other metal ions (for example, Ca, Sr, Ba, and Cd), of which the original structure of the MOF cannot be made in the form of monometallic MOF.

Presently combinations of different methods are very popular, and hydro(solvo)thermal method is no exception. In particular, hydro(solvo)thermal hybrid methods are often applied for synthesis of CPs and functional materials, including nanomaterials. A considerable body of studies has been performed for hybridization of the hydrothermal method with microwaves, electrochemistry, ultrasound, mechanochemistry, optical radiation and hot pressing. Contrary to ordinary heating hydro(solvo)thermal method, which requires a lot of time (as a rule, from half to several days) and high electrical power (more than thousand watts), microwave-assisted heating is a greener approach to synthesis of materials for shorter time (from several minutes to several hours), and lower energy consuming (hundreds watts), as a result of direct and uniform heating a matter. As an example, we can consider UV-absorber \((\text{Ti}_5\text{O}_{12}\text{F}_{12})\) obtained using microwave-heating-assisted hydrothermal synthesis, whose 3D network is built from \(\infty(\text{Ti}_5\text{O}_{12}\text{F}_{12})\) infinite inorganic layers separated by guanidinium (L) cations. Under UV irradiation at 254 nm during 40 h, white microcrystalline powder transformed into light-purple-gray due to reduction of Ti(IV) to Ti(III), which is confirmed by magnetic measurements.

### 3.1.4. Urothermal synthesis

Recently urothermal synthesis has opened a new approach to creation of porous framework materials with promising applications. It is based on using different derivatives of urea as solvents. One of very useful features of urothermal synthesis is reversible coordination of urea derivatives to metal sites, which allows them to competitively link with metal sites of a framework, and in many cases they can be easily removed after crystallization for generation both porosity and open metal sites. Competition for coordination to metal sites among derivatives of urea and other solvents, such as DMF or DEF, is an interesting aspect of this process of synthesis, and can also be used to provide additional structural control.

### 3.1.5. Ionothermal synthesis

Using room temperature ionic liquids or deep eutectic solvents in CP production is called ionothermal synthesis. Usually in MOFs synthesis ionic liquid...
liquids serve as solvents, reagents, structure-directing agents, charge-compensating templates in anion networks or functional inclusions. Ionic liquids attract attention as a solvent for chemical synthesis due to their unique properties, such as, actually, zero vapor pressure, excellent solvating properties, easy secondary using, and high thermal stability. They received much attention as promising media for design and production of known and unprecedented new MOFs. As compared with water or traditional molecular organic liquid, different and easily tunable properties of the solvent for the ionic liquid supplement considerable attention to eco-friendly and safe solvents. Unfortunately, high cost of ionic liquids limits their usage in large scale synthesis of MOFs.

Most works concerned with ionothermal MOF synthesis is focused on ionic liquids obtained from 1-alkyl-3-methylimidazolium. Thus, 3D MOF \([\text{EMIM}][\text{In}_2(\mu_3-\text{OH})_2\text{L}_2\cdot2\text{H}_2\text{O}]_n\) based on 1,2,4,5-benzenetetracarboxylate (L) is synthesized using ionic liquid bromide 1-ethyl-3-methylimidazolium (\[\text{EMIM}\] Br) as a solvent.\(^{233}\) In this polymer, 7-coordinated pentagonal bipyramidal In(1) and octahedral In(2) atoms are linked by \(\mu_3\)-OH groups with formation of infinite \([\text{In}_2(\mu_3\text{-OH})_2]_n\) inorganic chain along \(a\) axis, which are additionally extended by L ligands for creation of 3D anion microporous framework. The \[\text{EMIM}\]^+ cations occupy 1D channels acting as a template and charge-compensating species.

### 3.1.6. Synthesis in supercritical \(\text{CO}_2\)

Eco-efficient method based on using only supercritical \(\text{CO}_2\) as a solvent without the addition of any other additive or co-solvent is applied for synthesis of 1D Cu(II) MOFs.\(^{234}\) For example, neutral Cu(acac)$_2$ chelates and two linear spacers, the bidentate 4,4'-bipy and trimethylene-4,4'-bipy, were subjected to interaction under compressed \(\text{CO}_2\) at 60 °C and 20 MPa during 4 or 24 h, respectively (Fig. 21). The reaction yield of the CPs synthesized through the supercritical route was close to 100%, since both the reagents were almost completely consumed in the performed experiments. It should be noted that success reached in synthesis of different 1D MOFs is due to high solubility of reagents in supercritical \(\text{CO}_2\).

### 3.1.7. Surfactant-thermal method

Another approach to ensure a high degree of control over size and shape of MOF crystals and over their textural properties is use of surfactants (for example, amphiphiles and block-copolymers).\(^{235,236}\) It is important that the surfactants play different roles in control over morphology of MOF crystals. In particular, under special conditions, surfactants are self-assembling to form micelles, which can act as nanoreactors.\(^{232}\) In addition, in this case, MOFs grow on external surface of a micelle, not inside, so that a surfactant works as a molecular template, not a nanoreactor leading to formation of meso- and/or macro-pores, which together with intrinsic microporosity of MOF leads to particles with hierarchical porosity.\(^{237,238}\)

As compared with organic solvents, surfactants are more thermally stable and more environmentally. At the same time, surfactants with low or no pressure vapor make reactions in the surfactant medium possible at far higher temperatures. In addition, surfactants have more multifunctional properties than ionic liquids, such as cationic, anionic, neutral, zwitterionic, acidic, basic, etc. Which is more important, their low cost and commercial availability of surfactants make them ideal reaction medium for production of functional materials. Presently, the surfactant medium is used for preparation of nano-materials, and it should be noted that use of surfactants as reaction medium for the growth of crystalline materials is unprecedented. Janus characteristics of surfactants, which contain hydrophobic and hydrophilic groups, can increase efficiently solubility of metal ions and organic ligands, making them ideal medium for MOF growth.

We note the study of length-controlled synthesis of 1D M-Pp CP using \(\text{bottom-up}\) strategy under action of anion surfactant.\(^{239}\) For example, using this method, the 1D structure of

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**Fig. 21** The preparation of Cu(II) 1D MOFs using an eco-efficient method based exclusively on supercritical \(\text{CO}_2\) as a solvent. [Reproduced from ref. 234 with permission from The Royal Society of Chemistry.]
nanorods and nanowires are obtained. The strategy of growth of crystalline materials is developed: growth of crystalline chalcogenide materials using surfactant-thermal method.\textsuperscript{322,223,224} Chiral Ni(ii) mononuclear chelate [NiL(H$_2$O)$_2$] and 3D CP [NiL(H$_2$O)$_2$]$_n$ based on thiazolidine 2,4-dicarboxylic acid (H$_2$L) are synthesized by the reaction between Ni acetate and H$_2$L in aqueous solution at 25 and 80 \degree C, respectively.\textsuperscript{246} From the same procedure with polyvinylpyrrolidone (PVP) used as a surfactant, another respective micron size Ni(ii) CP [NiL(H$_2$O)$_2$]$_n$ is obtained at 25 and at 80 \degree C (Fig. 22).

Very important is the fact that surfactants can be capping agents or inhibitors, thus providing steric stabilization, which leads to the formation of nanoparticles and has a different effect on various faces of a crystal causing growth anisotropy of MOF crystals. In this case surfactants slow down crystal growth rate, providing spatial stabilization of MOF nanoparticles\textsuperscript{223,224} and relative value of different faces of a crystal, which leads to anisotropic growth.\textsuperscript{257,258,243,244} Capping agents, such as trisodium citrate, are used for limitation of growth of nanoparticles and stabilization of their aggregation. Their effect is, as a rule, not long, though using trisodium citrate with Lee and Meisel method can bring to particles, which are stable for a month.

3.1.8. In situ spacer synthesis. The approach called \textit{(in situ} spacer synthesis), in which organic spacers are generated in reaction medium \textit{in situ} from initial materials, attracts more interest in the recent years.

As an example, we shall consider solvothermal reaction of Pb acetate with flexible 1,3-bis(4-pyridyl-3-carboxyl)-propane (H$_2$L) under different conditions of synthesis through \textit{in situ} ligand transformation reaction, which give three genuine coordination polymers [PbL$_2$]$_n$, [Pb$_2$L$_3$]$_n$ and [Pb$_2$L$_3$]·2H$_2$O$_n$ (Fig. 23).\textsuperscript{246} In these compounds, L$^2$ ligand shows different coordination conformations and regimes adjusted to different synthesis conditions, including temperature of reaction, cooling rate and dopants, and designs various architectures linking different building units. Two first polymorphs show 3D framework with 1D channels built from binuclear ring-like [Pb$_2$(L$^2$)$_2$] and binuclear semiring-like [Pb$_4$L$_2$] units, respectively. The last polymorph also has 3D architecture; however, it is built from binuclear ring-like [Pb$_2$(L$^2$)$_2$] units interconnected with L$^2$ ligand.

We shall also notice CPs with [Cu(n)(mu$_3$-L)$_2$]$_n$ [Cu(n)Cu(n)(mu$_3$-L)$_2$]$_n$ units based on chelating ligand of 2-pyrazinocarboxylate (HL), which is formed \textit{in situ} by hydrolysis of the initial N,N’-di-hydro-1,3,5,7-tetraoxobenzo[1,2-c:4,5-c’]dipyrrrole-2,6(1H,3H)-dipyrryl-3-pyrazinocarboxamide upon its reaction with Cu(n) chloride, iodide, and perchlorate, respectively.\textsuperscript{246} The first CP has 2D double-layer structure, which can be considered as a binodal (3,3)-connected network, the second polymer is (4,4) net, and the third CP has a 3D open framework built by linking 1D sawtooth chains of Cu(n) iodide clusters with [Cu(n)(mu$_3$-L)] units. It is interesting that all compounds cannot be synthesized directly using HL and the respective Cu ions as initial reagents under the same conditions of reaction. Therefore, the structural diversity of this system is adjusted \textit{in situ} by hydrolysis reactions.

3.2. Alternative synthesis routes

In hydro(solvo)thermal processes, considerable volumes of a solvent are necessary to produce great quantities of CP for commercial using, which is environmentally unfavorable, and makes products costly. It is also necessary to accelerate crystallization process and generate uniform crystals with diminished size; therefore it is important to develop easily available, non-expensive, fast, and commercially viable synthetic routes. Many efforts were taken for development of alternative methods of CP synthesis, such as electrochemical, sonochemical, microwave, and mechanochemical methods. These methods provide possibilities for MOF synthesis in reduced time and with high quality, which is favorable for commercial application of MOFs. In this view, detailed studies were performed for optimization of MOF synthesis conditions to obtain high yields of solid products for industrial applications.\textsuperscript{243,244}

Fig. 22. Schematic representation of the generation of discrete 1, 3D framework 2, and 1D left helical chain 3 triggered by temperature or polyvinylpyrrolidone. [Reprinted with permission from ref. 242. Copyright 2012, American Chemical Society.]
3.2.1. Microwave-assisted synthesis. It is important to emphasize that microwave-assisted MOF syntheses are mainly focused on the following moments: acceleration of crystallization, formation of nanometer materials, improvement of purity of a final product, and selective synthesis of polymorphs. This is reached by direct heating solvents and increase in crystal nucleation rate. To optimize the reaction conditions, systematic studies of changes in the composition and process parameters (solvent, irradiation time, temperature of reaction, power level, the molar ratio of reagents, reagent concentration, etc.) were carried out. It should be noted that microwave irradiation makes it possible to accelerate synthesis of fine crystals as compared with conventional heating. On the whole, advantages of this method are high efficiency, phase selectivity, particle size diminishing, and morphology control.

As a typical example, we consider microwave-assisted synthesis of Co-MOF-74 under different conditions, in particular, temperature of synthesis, power level of microwave irradiation, and time of synthesis. Optimized Co-MOF-74 was synthesized at 180 °C at 180 W power in a microwave oven during 1 h, which resulted in the highest textural properties (1314 m² g⁻¹ of BET surface area) with high product yield (about 76% per ligand). Co-MOF-74 (ca. 50 µm long and 8 µm wide) obtained by microwave heating was smaller in size than the solvothermally prepared Co-MOF-74 (ca. 300 µm long and 70 µm wide).

Another interesting example is production of rare earth based CP submicrospheres from pyridine-2,5-dicarboxylic acid and M(NO₃)₃ using the method of light microwave heating during 5 min with DMF as a solvent. Submicrospheres have diameters 100–400 nm, and the microspheres’ surface is smooth. Straw-sheaf-like Tb-based CP architectures are successfully synthesized using a simple and ecological microwave heating in great scales during 15 min without any template or surfactant using 1,2,4,5-benzenetetracarboxylic acid as organic building unit. The measured specific surface area of Tb-based CP was 152.51 m² g⁻¹.
Individual straw-sheaf has length in the range 70–90 μm, and average diameter in the range 5–8 μm.

3.2.2. Electrochemical synthesis. In order to exclude metal salts usage, metal ions are continuously integrated via anode dissolution into the reaction mixture, which contains dissolved spacer molecules and conducting salt.\textsuperscript{249} Other advantage of electrochemical route is a possibility to launch a continuous process and a possibility to obtain higher content of solids as compared with batch reactions. Undoubtedly, electrochemical MOF synthesis provides alternative method of synthesis, which does not require additional external source of heat, however, it is too expensive to be used at industrial scale. It is important that this approach works through formation of the reagent ions in a solution, thus it is unnecessary to use bases. It is worth noticing that electrochemical methods suits well for formation of MOF membranes due to intrinsic coating reached on scaffolds due to a MOF product deposition.

3.2.3. Mechanochemical synthesis. In mechanochemical synthesis, the mechanical break of intramolecular bonds followed by chemical transformation occurs.\textsuperscript{250} Mechanochemical synthesis is a promising alternative method for systematic and large scaled CP production. Its advantages include the following: lowering of contamination, low cost, simplicity of the process, easiness of processing, efficient reaction rate, selectivity, and solution of a problem of low solubility of reagents. Mechanoysis or milling is currently used as fast, scaled, and potentially non-toxic way of MOF synthesis. MOFs can also be obtained using mechanochemistry, i.e. grinding of two or more solids using a mechanical ball mill and thus avoiding use of solvents.\textsuperscript{250} This is very aggressive technique, which makes it possible to obtain products from micro- to nanocrystals at room temperature in a shorter time.

Using twin screw and single screw extruders has been shown for continuous synthesis of different CPs, including Ni(salen)-based.\textsuperscript{251} Quantitative conversions are reached with the formation of products at the kg h\textsuperscript{-1} rates, which after activation show surface areas and volume of the pores equivalent to materials obtained by conventional solvent-based method.

High proton conductivity material has been obtained using economical and ecologically pure mechanochemistry.\textsuperscript{252} The reaction is a new synthetic strategy with respect to materials related to technology of fuel cells.

3.2.4. Sonochemical synthesis. It is important that using ultrasound of high intensity provides easy, ecological, and universal synthetic tool for production of nanostructured materials, which are often unavailable for conventional methods.\textsuperscript{253}

Sonochemical methods through uniform and accelerated nucleation also provide decrease in crystallization time and far less sizes of particles, than those provided by conventional solvothermal synthesis. For this purpose, a solution of mixture of substrates for this MOF structure is input in a horn-type Pyrex reactor equipped with an ultrasonic bar with regulated output power without external cooling. As an example, we shall consider sonochemical synthesis of nanometer MOF with [Pb\textsubscript{2}(N\textsubscript{3})(NO\textsubscript{3})Q\textsubscript{2}] unit, where HQ is 8-hydroxyquinoline.\textsuperscript{254}

Sonochemically are also obtained nanostructures of CPs with [Pb\textsubscript{2}L\textsubscript{4}]\textsubscript{n} (HL = 2-pyrazinecarboxylic acid) and [Pb(L\textsubscript{1})\textsubscript{2}]\textsubscript{n} (HL = 2-quinolinocarboxylic acid) units.\textsuperscript{255}

There is an interest in sonochemical synthesis of nano-sized Pb(u) CP with 1H-1,2,4-triazole-3-carboxylate ligand.\textsuperscript{256}

High quality Mg-MOF-74 crystals (1.640 m\textsuperscript{2} g\textsuperscript{-1} BET surface area) with particle size about 0.6 μm are successfully synthesized during 1 h by sonochemical method after addition of triethylamine as a deprotonating agent. It is interesting that mesopores are formed, probably, due to competitive linking triethylamine to Mg(u) ions.\textsuperscript{258}

4. Post-synthetic transformations

In the recent years, post-synthetic transformations have become powerful tools for synthesis of the CPs with chelated units, which cannot be reached with de novo synthesis due to limited linker solubility, thermal stability, chemical stability, functional group compatibility, and undesired intervention between metal ions and linker of functional fragments during MOF assembling. Removal or substitution of solvents and guest molecules, action of chemically active vapors and external stimuli, such as heat, light or mechanochemical force, induce such structural transformations, and they often manifest themselves in different physical properties, such as color, magnetism, luminescence, chirality, porosity, etc. because of a change in coordination number and geometry, dimensionality, interpenetration, etc. More radical transformations associated with exchange of metal ions, pillar ligands, and insertions of additional ligands between layers were also represented.

Post-synthetic transformations include post-synthetic modification (PSM), post-synthetic deprotection (PSD) and post-synthetic exchange (PSE).

4.1. Post-synthetic modification

Post-synthetic modification (PSM) is a single-crystal to single-crystal (SC–SC) transformation and is the process including transformation of side groups on surface or in pores of MOFs in order to change physical or chemical properties. This can be reached with chemical reaction, adsorption, ligand or metal exchange, light, and pressure stimuli. Post-synthetic MOF modification opens new dimension of structural transformations, which cannot be reached by conventional synthesis.

There are metal centers and initial ligands with functional groups, which can take part in a reaction under normal conditions of MOF synthesis for the preparation of MOFs with specified crystal structures and pore sizes. Then the functional groups on ligands can react with post-synthetic reagents to obtain final desired functionalities.\textsuperscript{257} PSM MOFs can be realized using multiple interactions, among which covalent and coordinate bonds are the most important. Post-synthetic methods, which modify MOFs with chemical reagents with the lattice structure unchanged, make it possible to integrate functional groups in MOFs.\textsuperscript{258}
Chelating groups can be introduced more easily by post-than by presynthetic functionalization in MOFs since they may compete in the coordination to inorganic nodes. There are various examples in the literature that show how to produce chelating groups by PSM. It is important that potential chelating fragments can be introduced in MOFs by covalent post-synthetic modification, so that they do not cause interference to the synthesis, but are still available for further coordination modification.

In particular, universality of 2-amino-1,4-benzenedi-carboxylate as a building block has been showed: it can be smoothly subjected to PSM in three different prototypic MOF topologies: IRMOF-1, DMOF-1 (D = dabco), and UMCM-1. The important features of PSM on IRMOF-3 include (1) facile introduction of a wide range of functional groups using simple reagents (e.g., anhydrides and isocyanates), (2) the introduction of multiple (as many as four different) substituents into the MOF lattice, and (3) control over reaction conditions to preserve the crystallinity and microporosity of the resultant MOFs.

As a typical example, we note imine condensation of IRMOF-3 wetted in toluene solution and excess of salicylaldehyde during 7 days, which has given a PSM-product with ~13% conversion of amino groups. Then V(O)acac$_2$ was loaded in the framework for coordination with the salicylidene groups, and the second step was almost quantitative (Scheme 18). In this case, post-synthetic derivatization of a porous material gives functionalized material, which links a metal complex VO(acac)$_2$ in contrast to non-functionalized precursor, which is inactive for complex linking.

In conversion of roles, aldehyde can be a part of MOF, as in the case of ZIF-90 (isostructural to ZIF-8, but with a tag aldehyde group). Specifically, reduction of aldehyde to alcohol functionality was successfully achieved by reacting ZIF-90 with NaBH$_4$ in methanol at 60 °C for 24 h to give ZIF-91 with about 80% conversion (Fig. 24A). The chemical versatility of the aldehyde group was highlighted by performing another organic transformation on ZIF-90. Reaction of ZIF-90 with ethanolamine in methanol at 60 °C gave ZIF-92 (Fig. 24B). Quantitative conversion to the imine was completed within three hours.

Tandem PSM is useful, since it can be realized in one-pot synthesis. PSM MIL-101(Fe)-NH$_2$ with Py-2-carboxaldehyde and NiCl$_2$ in the one-pot reaction has given ~30% conversion of amino groups into pyridine imino groups (Scheme 19). Nickel centers are integrated as catalytic sites into pores by coordination with a nitrogen atom of imino group and a pyridine nitrogen atom. Ni complex is formed, first of all, to stop any competition with aldehyde reagent for amino groups.

Additional set of examples is observation that traditional batch hydro(solvo)thermal methods of synthesis, as a rule, exclude direct incorporation of free-base Pp as a spacer in MOF, since spacers spontaneously link metal ions present as building blocks and nodes. A promising particular solution of the abovementioned problem is post-synthetic treatment afterwards renamed and generalized in PSM, i.e. chemical substitution (atom, ion or functional group, addition, substitution or transformation) of a framework without a change in parent topology. Really, a task with participation of metal-free Pp or salen-based spacers was first met by assembling metallized spacer in variants of compounds, and then by substitution of non-structural (i.e. spacer-localized) metal ions with a pair of protons. The second stage of PSM accepts new metal ions.
It should be noted that covalent modification of bridging ligands is studied in detail. A special interest is MOF, where metal centers are subjected to modification. Also a possibility is demonstrated of PSM via coordination of additional metal ions with centers in bridging ligands and addition or removal of metal ions in metal centers.

Of particular note is the linker installation method, in which a stable Zr-MOF with inherent missing linker sites, namely, PCN-700, with coordinatively unsaturated Zr_6 clusters was employed and linkers bearing different functional groups were post-synthetically installed (Fig. 25). At first, the bpy-5,5'-dicarboxylate and terphenyl-4,4'-dicarboxylate were sequentially installed by treating PCN-700 crystals with solutions of bpy-5,5'-dicarboxylate and terphenyl-4,4'-dicarboxylate in DMF at 75 °C. The modified MOF bears open bpy sites that readily react with CuI in acetonitrile to form the metalated framework. According to the crystal structure, the Cu^+ center is chelated by a bpy group and further coordinated with an N from acetonitrile and an I^- ion as a counter-ion. The Cu^+ center is arranged around the 1D channel along the c-axis, whereas the terphenyl-4,4'-dicarboxylate controls the accessibility of the Cu^+ center by partially blocking the channels. Therefore, the size selectivity of the whole material can be tuned by changing the size of substituents on the terphenyl-4,4'-dicarboxylate linker. In particular, the terphenyl-4,4'-dicarboxylate linker was functionalized by methyl groups, phenyl groups, and hexyl groups, which is expected to result in different selectivity toward the substrates.

Unsaturated metal sites were used in an alternative method, in which MOF with [Cu_L]_3 unit (L = 1,3,5-benzenetricarboxylate) was modified with en followed desolvation under vacuum to prepare free coordinating fragments.

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4.2. Post-synthetic deprotection

PSD is useful for synthesis of functional MOFs, such as those which have groups with a trend to metal complexes formation in direct synthesis. The process includes elimination of complexity in MOF and in production of big cavities. PSD is sometimes also called protection, complexation, and deprotection.267

By now, PSD just begins to draw attention as a post-synthetic method of functionalization. The concept of this method is that a protective functional group is incorporated as an organic spacer, and then the protective group is removed by the post-synthetic method. It should be noted that post-synthetic MOF modification is realized for the preparation of single-site catalysts.268

Protection–deprotection methods open new possibilities for (unlock) of protected functionalities for production of highly active and/or coordinating fragments on MOFs. This can facilitate immobilization of metal–organic catalysts in MOF cavities.269

4.3. Post-synthetic exchange

Post-synthetic exchange (PSE) also known as a building block replacement261 includes replacement of basic structural components of preliminary assembled MOFs. PSE methods include (I) solvent-assisted linker exchange (SALE), (II) non-bridge ligand replacement, and (III) transmetalation. These one-stage or tandem PSE processes suppose exchange of key structural MOF components, which, in turn, should provide evolution of initial MOF structures for MOFs design consisting of absolutely new components, supposedly, using SC–SC transformations.

SALE is one of most widely used PSE approaches for adjustment of MOF functionality. SALE realization, as a rule, involves holding MOFs crystals in a solution of a spacer substitution candidate in a thorough chosen solvent.279 It is worth noticing that SALE proceeds through exchange of ligands in undamaged MOFs, but not by solution and re-crystallization. Therefore, topology of a parent MOF is reproduced in a daughter structure. SALE was demonstrated for a range of MOFs,276–278 including such, seemingly, inert compounds as ZIFs and, which is more surprising, UiO-66,274,275 the material, which is characterized by unusually strong M–L bonds.

Thus, modification of UiO-66 is executed with catechene using SALE in DMF/H₂O solution during 2 days at 85 °C. After metalation, Cr-metalized MOFs were synthesized (Fig. 26).276 Importantly, insulated metal-monocatechelato fragment is achieved in very strong MOF by two principally different strategies: PSD and PSE. As compared with PSD, PSE is easier and more efficient approach to functioning for MOFs access, which cannot be synthesized directly under solvothermal conditions. Metalation of catechene functionality remained in MOFs led to unprecedented Fe-monocatechelato and Cr-monocatechelato types.

PSE was also employed as a strategy to introduce dimercapto functionality into UiO-66.277 In particular, PSE was performed by incubating solid UiO-66 in an aqueous solution of 2,3-dithiacatechene-1,4-benzendicarboxylic acid (tcat-H₂bdc) for 24 h at 85 °C. The linker-exchanged thiocalcat material, UiO-66-TCAT, was isolated as a yellow microcrystalline powder using centrifugation. Immobilized thiocalcat ligands on the UiO-66-TCAT provide an excellent platform to achieve accessible and unsaturated mono(thiocalcat) metal centers (Scheme 20). The metalation of 40% thiocalcat-functionalized UiO-66-TCAT using Pd(OAc)₂ in CH₃Cl₂ afforded dark-brown solids. The crystallinity of UiO-66-PdTTCAT was maintained upon metalation. It is important to notice that Pd-metalated MOFs are efficient, heterogeneous, and recycled catalysts for regioseleetive functionalization of sp² C–H bond. This material is a rare example of chelating-assisted C–H functionalization performed by a MOF catalyst.

Consecutive process of post-synthetic exchange of ligands was used for production of a range of mono-, di-, and tri-functionalized mesoporous MOFs.278 Using this process, orthogonal functional groups are established, and after that post-synthetically modified with a dye and quencher molecules, illustrating the level of structural and functional complexity that can be achieved within this system. It is important that the degree of completion and the relative ease of ligand exchange can depend on the lability of the M–L bond within the MOF.

An extended Pp MOFs family is developed, which includes directly variety of M-Pp through M-Pp and M-bipyridyl Pp as building units, which have large available channels and active metal sites (Scheme 21).279

A simple substitution of a solvent is basic for PSM chemistry. As a typical example, we note a solvent replacement in MOF with [CdL₂(CIO₄)₂]ₙ unit, where L = (S)-2,2’-diethoxy-1,1’-binaphthyl-6,6’-bis(4-vinylpyridine), in which ethanol was reversibly exchanged with benzene in chiral 1D hexagonal channel. It is assumed that this MOF has a good potential for heterogeneous asymmetric catalysis, since the structure can be supported after replacement of a solution.279

Post-synthetic transmetalation has become a prominent way of synthesis of new crystalline materials based CPs and is especially powerful for orientation of metastable phases, which are unavailable for conventional high temperature synthetic methods.280,281 By now, PSD just begins to draw attention as a post-synthetic method of functionalization. The concept of this method is that a protective functional group is incorporated as an organic spacer, and then the protective group is removed by the post-synthetic method. It should be noted that post-synthetic MOF modification is realized for the preparation of single-site catalysts.268

Protection–deprotection methods open new possibilities for (unlock) of protected functionalities for production of highly active and/or coordinating fragments on MOFs. This can facilitate immobilization of metal–organic catalysts in MOF cavities.269

Fig. 26 Schematic view of catechene-functionalized UiO-66 after metalation for catalytic oxidation of alcohols to ketones. [Reprinted with permission from ref. 276. Copyright 2014, American Chemical Society.]
at the atomic level structural characterization. Thus, relative occupation of cation on two crystallographically different metal sites in Fe(II), Cu(II), and Zn(II)-exchanged versions of microporous MnMnL MOF \( (L = 1,3,5\text{-benzenetristetrazolate}) \), was established. It is important that using dispersion differences between Mn, Fe, Cu, and Zn, degree and localization of cation exchange are determined. In addition, the replacement degree for Mn(II) depends on the identity of the substituting metal.\(^{283}\)

Different amounts of Co and Ni were substituted in Mg-MOF-74 using one-pot solvothermal reaction.\(^{284}\) Based on elemental analysis, Co and Ni were more efficiently incorporated into MOF-74 framework from solution than Mg was. In addition, temperature of the reaction has stronger effect on the final composition of metal in these mixed metallic MOF-74 structures than a solvent composition of the reaction.

Fast room-temperature cation exchange is also observed during soaking of CP with \([\text{Pb}_2\text{Cl}_2 \text{HL}_2 \text{(H}_2\text{O})_2]_n\) unit based on pyrazine-2,3-dicarboxylic acid (\(\text{H}_2\text{L}\)) in Cu(II) aqueous solutions (Scheme 22).\(^{127}\)

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**Scheme 20** Schematic view of the synthesis of UiO-66-TCAT and UiO-66-PdTCAT. [Reprinted with permission from ref. 277. Copyright 2015, American Chemical Society.]

**Scheme 21** Schematic illustration of the introduction of 4,4’-bipy to the crystals of PPF-18 (a) and PPF-20 (b) for the transformation of PPF-27 (c) and PPF-4 (d), respectively. [Reproduced from ref. 110 with permission from The Royal Society of Chemistry.]
Reaction of MOF with anhydrous zinc acetate in DMF at 100 °C afforded a highly crystalline purple material, consistent with 90% occupancy of the Pp center sites by Zn(II). Visible light catalytic activity of this Zn-Pp based material is shown on sacrificial segregation of hydrogen from water.

Scheme 22 Cation exchange by simple soaking of Pb(n) MOF into an aqueous solution of Cu(n) cation at room temperature. [Reprinted with permission from ref. 127. Copyright 2015, American Chemical Society.]

Series of metallosalen-based MOFs is prepared using post-synthetic modification of Mn[III]SO-MOF, Mn[III](salen)-based MOF. Treatment of Mn[III]SO-MOF with H2O2 leads to removal of Mn3+ ions from salen pillar, which then can be remetalated with different metal precursors with formation of isostructural MSO-MOF materials [Scheme 23]. Presence of new metallosalen pillars in MSO-MOF is entirely confirmed. It is important that remetalated Mn[III]SO-MOF material reflects similar catalytic activity and porosity to initial MOF.

Exchange of metal cations within a MOF has also been demonstrated for M-Pp MOF. Thus, easy route to M-Pp-based nanoreactor with post-synthetic metal-ion exchange with active metal cations and catalytically inactive MMPF, which consists of nanoscopic multifaced cages is developed.40 Local coordination of the active metal center in Pp macrocycle after the metal-ion exchange was crystallographically identified.

It should be noted cationic PCN-223(Fe) formed by synthetic treatment of PCN-223, Pp Zr-MOF with sh-p-a network, which is excellent recycled heterogeneous catalyst for hetero-Diels–Alder reaction.286

Two isostructural Zr-Pp and Hf-Pp MOF (FJI-H6 and FJI-H7) are rationally synthesized and built from 2.5 nm cubic cages.287 It is worth noticing that in FJI-H6 and FJI-H7 two nitrogen atoms of the Pp ring are not deprotonated, therefore metal ion of another type is integrated in the framework. Submergence of FJI-H6 or FJI-H7 single crystals in 0.5 M solution of Cu(NO3)2 in DMF at the temperature 85 °C during 72 h leads to metalated FJI-H6(Cu) or FJI-H7(Cu). As was expected, both in FJI-H6 and FJI-H7 Cu(n) ions were inbuilt in Pp rings, and Cu(n) ions in plane-square N4 coordination with two axial sites were exposed, which is typical of bivalent metal ions in Pp MOFs.

Of interest are isostructural Pp MOFs, in particular, UNLPF-10b, -11, and -12, which consist of In[III], Sn[IV]-Cl2-, and Sn[IV]-Pp building units, respectively.288 Metallation with high valence metal cations (In[III] and Sn[IV]) considerably changes electronic structure of Pp macrocycle and provides high oxidation photoexcited state, which can be subjected to efficient reduction extinguishing processes to facilitate organic reactions.

It should be noted Pp-based MOF MMPF-5, in which each small cubicoctahedral cavity consists of faces of six Cd[II]-metalated tetrakis(3,5-dicarboxyphenyl)porphine units interlinked by eight triangular Cd(CO3)3 fragments (Fig. 27a).60 In MMPF-5 crystal structure, aCd(n) cation within Pp ligand ring is far from Pp plane (Fig. 27b), which points to weak linking Cd(n) cations in Pp macrocycle and on a possibility to replace big Cd(n) by a smaller cation of a bivalent metal. Submerging the MMPF-5 crystals into DMSO solution of Co(NO3)2 at 85 °C for two days gives Co(n)-exchanged MMPF-5(Co). Complete substitution of Cd(n) by Co(n) is confirmed exclusively within Pp macrocycles, however, Cd(n) cations in the framework have remained untouched, possibly, due to their strong chelating with six carboxylic oxygen atoms. Therefore, small cubicoctahedral cavity in MMPF-5(Co) is distinguished by faces of six Cd[II] coordination with two axial sites were exposed, which is typical of bivalent metal ions in Pp MOFs.

Post-synthetic incorporation of Zn into free-base of Al(n) MOF with formation of light-harvesting MOFs proceeds by similar way.41 Reaction of MOF with anhydrous zinc acetate in DMF at 100 °C afforded a highly crystalline purple material, consistent with 90% occupancy of the Pp center sites by Zn(n). Visible light catalytic activity of this Zn-Pp based material is shown on sacrificial segregation of hydrogen from water.

Scheme 23 Demetalation of Mn(n)SO-MOF and its subsequent remetalation. [Reprinted with permission from ref. 285. Copyright 2011, American Chemical Society.]
Of interest is post-synthetic metalation of reliable chiral and porous BINAP-based Zr-MOFs UiO topologies with Rh(nbd)$_2$BF$_4$ and [Rh(nbd)Cl]$_2$/AgSbF$_6$, where nbd is norbornadiene, to give highly active enantioselective one-site solid catalysts for reaction of asymmetric cyclization of 1,6-enynes.

Extremely flexible and strong Cd-MOF anion accessible with tetra-acid spacer, in particular, 3,3',5,5'-tetrakis(p-carboxyphenyl)-2,2',6,6'-tetramethoxy-1,1'-biphenyl, shows abundant post-synthetic metal exchange of metal sites with the metal ion set, which differs significantly by their charges, ion radius, and chemical origin. In particular, there are as many as 16 isostructural new MOFs with transition, lanthanide, and basic groups of metal ions.

Taking into account universality and practical usefulness of PSM and PSE for access to other difficult-to-synthesize MOFs as bulk materials, their application to MOFs in thin film can also be useful. Among the most universal and useful approaches to synthesis of MOF films is layer-by-layer (LbL) coordination, which is called liquid phase epitaxy. LbL approach successfully and repetitively inputs solutions of each building block of a framework in contact with functionalized substrates, and then a MOF film grows. It is important that composition of additional layer not necessarily coincides with previous layers. Reliability of LbL-assembled pillared-paddlewheel MOF films in direction of transformation into new or modified MOFs through SALE and post-synthetic metalation has been demonstrated. In particular, it is shown that LbL synthesis provides MOFs, which are unavailable through other de novo strategies (Scheme 24). MOF contains 5,10,15,20-(4-carboxyphenyl)-Pp units as tetratopic spacers, which coordinate pairs of Zn(II) ions in paddlewheel mode, and 4,4'-bipy spacer, which work as Zn(II)-ligating spacers or pillars between Pp defined layers. A dabco was used as a spacer for exchange or replacement of 4,4'-bipy pillars MOF. In addition, after 12 h reaction in excess magnesium bromide diethyl etherate, the free-base Pp containing films grown of quartz slides were transformed into M-Pp form.

Nevertheless, there are some drawbacks of post-synthetic approaches. For example, it is difficult to solve spatial distribution of functional groups post-synthetically included in MOF. As a result, most MOF structures are hypothetical. In some cases, a degree of post-synthetic incorporation of functional groups in MOFs is very limited. Moreover, further studies should investigate chemical principles of their specific phenomena, including SALE and transmetalation in order to improving the direct synthesis of MOFs with required chelating functionality.

5. Thermolysis of coordination polymers as a route to nanostructures materials

Recently CPs have become widely used as templates for a variety of nanomaterials. Unlike other templates, they produce monodisperse nanoparticles of high quality, while maintaining the porosity and morphology of the template precursor. Especially CPs widely used for the preparation of metal, metal oxide and carbon nanomaterials.

As a typical example, we note the strategy of using MOF thermolysis for the preparation of metal and metal oxides nano-
sized particles, such as Cu/CuO, Co/Co$_3$O$_4$, ZnO, Mn$_2$O$_3$, MgO and CdS/CdO. According to this strategy, during thermolysis in N$_2$, metals with a reduction potential of $-$0.27 V or higher contained in MOFs always form pure metal nano-sized particles, whereas metals with a reduction potential lower than $-$0.27 V form metal oxide nano-sized particles (Fig. 28). It is important that there is a relationship between the size of the formed nanoparticle and the distance between the SBU inside the MOF precursors. In addition, the crystallinity of the carbon matrix was also influenced by the thermolysis conditions (N$_2$ and air).

5.1. Synthesis of carbon materials

Interest in various forms and allotropic modifications of carbon is very wide from use as adsorbents, carriers for catalysts and drug delivery to the electrode materials. The highly porous carbon materials may be prepared by various methods, including physical and chemical activation of the carbon, polymer aerogels carbonization and templating methods using zeolites and mesoporous silica. Often, however, obtained mesoporous carbon materials are characterized by a broad polydispersity to micron-sized particle and their fuzzy shape.

The direct thermal transformation of CPs was used for the preparation of porous carbons with special morphologies (for example, cube and polyhedron etc.), and their pore structures may be easily varied depending on the as-formed metal or metal oxide nano-sized particles. However, using thermolysis of CPs for the preparation of nanocomposites is restrained by their high cost.

Aluminium–diethylenetriamine pentaacetic acid (Al–DTPA) microfibers were used as cost-effective precursors (Fig. 29) to prepare nitrogen-doped carbon microfibers (NCF).

In particular, thermal transformation and acid-leaching of Al–DTPA leads to the well-defined NCFs. It is important to underline that the microfiber morphology of Al–DTPA persists in the resulting carbon materials (Fig. 30B–E). The typical NCFs prepared at 900 °C (NCF-900) have the diameters of $\sim$500 nm and lengths of about 50 mm (Fig. 30B–D). It should be noted that some graphitization regions are observed in the high-resolution TEM image of NCF-900 (Fig. 30E). Increasing thermolysis temperature leads to a slight increase in the graphitization degree of carbon microfibers (Fig. 30E inset). Moreover, considerable interconnected pores are homogeneously distributed in the whole microfibers (Fig. 30D). The pore size distribution curves exhibited the mesopores with...
producing various carbon materials with catalytic and magnetic participation in di-molecules and a carbon source. Due to the high thermal in air at 800 °C, there is a simple manner, based on the MOF structures as template and length (50 mm) (Fig. 30A). The prepared -Al₂O₃ microfibers can be also used as the precursors of γ-Al₂O₃ microfibers. After thermalization of Al-DTPA in air at 800 °C, the well-defined microfiber morphology in γ-Al₂O₃ was observed in SEM images (Fig. 30F). As compared to Al-DTPA microfibers, there is decreasing the diameter (350 nm) and length (50 nm) (Fig. 30A). The prepared γ-Al₂O₃ has a smaller BET surface area (50 m² g⁻¹) and pore size distribution in the range of 2–15 nm.

Various types of porous carbon materials may be prepared in a simple manner, based on the MOF structures as template molecules and a carbon source. Due to the high thermal stability, the nanoporous structure and the possibility to penetrate into the pores of small molecules with their subsequent participation in different (ship-in-bottle) reactions, MOFs can be considered as rigid templates similar to the mesoporous silica gel and zeolites. Given the wide variety of MOFs of different compositions and morphology, the carbon materials can be produced with controlled pore texture and surface area. Moreover, the synthetic strategy based on MOFs allows to producing various carbon materials with catalytic and magnetic properties.⁶⁶

Advantages of MOFs templates can be realized very efficiently in the production of various composite carbon materials, for example, type M/MO@C (Fig. 31). The general approach of the synthesis is based on the fact that the 1st stage of polymerization is carried out in MOFs carbon precursor (Fig. 31a), then a M@C nanocomposite is formed by thermolysis (Fig. 31b), and finally post-thermal treatment is carried out to convert the metal to metal oxide nanoparticles (Fig. 31c). The porosity of the carbon matrix and M/MO nanoparticle sizes are controlled in stages.⁶⁷

It should be noted the obtaining a porous cobalt oxide-carbon hybrid by carbonizing in an inert atmosphere and subsequently air-calcing nanocrystals of ZIF-67.⁶⁸ It is important that the synthesized carbon composite material can act as an efficient electrochemical water oxidation catalyst.

5.2. The preparation of metal oxide materials

As a rule, the basis of the methods for producing metal oxide nanocomposites from CP is a single-stage solid-phase thermalization of the precursor source, as, in general, quite comfortable, well-reproducible and easily scalable process. The methodology was used in the synthesis of nanoparticles of various metal oxides, for example, PbO,⁶⁹ ZnO,⁷⁰ Fe₂O₃,⁷¹ Co₃O₄,⁷² CdO,⁷³ CuO/Cu₂O,⁷⁴⁷⁵ Tl₂O₃,⁷⁵ etc. We consider some of the most common examples.

Thus, [Pb₂(2,9-dimethyl-phen)(μ-N₃)(μ-ClO₄)]₉ (ref. 316) and [Cd(2,9-dimethyl-phen)(1-(2-thienyl)3,3,3-trifluoroacetone)₂]₉ (ref. 317) were used for PbO and CdO nanoparticles preparation, respectively. Thermal decomposition of the 3D CP [Pb(phen)(1-N₃)(1-NO₂)]₆ containing an azide-anion ligand gave PbO nanoparticles.⁷¹

ZnO nanomaterials with desired morphologies have been obtained by thermal decomposition of appropriate CPs and MOF precursors.⁵²,⁶⁹,¹¹³ For example, calcination of infinite rectangular-tubular helices [ZnCl₂₃] and [ZnBr₂₃] (L is bis(3-pyridyl)cycloctetramethylenesilane) at 500 °C gives uniform hexagonal tubular spire crystals of 1.2 × 1.2 × 4.0 µm³ dimensions and spheres, respectively.⁷³ SEM images of the products of [MX₃L] thermalization at 500 °C (Fig. 32) show that the compounds ultimately change to zinc(II) oxide crystals. For [ZnCl₂₃], calcination at 500 °C leads to the growth of each ZnO single crystal.

The Zn(n) oxide and Cd(n) oxide nanoparticles were prepared by thermolysis of Zn(n) and Cd(n) CPs, [Zn(L)₃(N₃)₃]₉, [Zn(L)₃(SCN)₃]₉, [Cd(L)₂(N₃)₃]₉, and [Cd(L)₂(SCN)₉]₉, where L is 2-aminomethylpyridine.⁷³ SEM images exhibit the average size of prepared ZnO and CdO nanoparticles of 60–70 nm.
A series of lead-containing CPs have been applied for PbO nanoparticle formation. PbO nanoparticles serve as catalyst for multicomponent organic synthesis reactions, for instance Paal-Knorr reaction and oxidative coupling of methane, due to the significant catalytic property with operational simplicity, high reactivity, environmental friendliness, reduction of reaction times, and reusability of PbO nanoparticles. In this respect, a CP of lead(ii) nitrate complexes with bridging trans-1,2-bis-(4-pyridyl) ethene (L) ligand, [Pb(µ-NO3)3(L)(MeOH)]n, was used to prepare PbO nanoparticles via two different methods.

The nanostructure of 1D copper CP, [Cu(HL)NO3]n (H2L is [2-1-(2-hydroxypropylimino)ethyl] phenol), was used as precursor to obtain single phase CuO nano-sized particles by thermolysis. According to the SEM images the diameters of the prepared products were equal to 40 nm (Fig. 33).

Calcinations of the nano-sized copper(ii) CP, [Cu(L)(H2O)2]n [L = 2,6-pyridinedicarboxylate], at 500 °C under air yielded CuO nanoparticles. After calcining the cerium-based CPs with three isomers of benzenedicarboxylic acid at high temperature, micro/nanostructures of ceria such as nanoparticles, nanorods, and microflowers with retained morphologies were achieved.

Thermolysis of the porous CPs with nanorod morphology based on a catechol-substituted Pp [meso-tetrakis(3,4-dihydroxyphenyl)-Pp] led to carboniferous materials containing metal oxide nanoparticles with large surface areas (up to 800 m² g⁻¹).

The advantages of these CP precursors are simplicity of processing and long-range ordering that allow the synthesis of metal oxides with various morphologies. Phase composition and properties of the resulting metal oxide nanocomposites depend on the crystal structure and morphology of the starting CP precursor, its particle size, and the synthesis conditions (the influence of the temperature, environment, the nature of the surfactant and etc.).

5.2.1. Influence of the original morphology and particle size of coordination polymers. Size and morphology of the produced nanomaterials are highly dependent on the type and morphology of the CP precursor.

Porous γ-Fe2O3 nanoparticles were prepared via a solid-state transformation process of a mesoporous iron(II) carboxylate crystal, MIL-100(Fe). The N2 adsorption-desorption analysis also demonstrates the mesoporous character of the derived γ-Fe2O3 material, with type IV N2 isotherm at 77 K. This material has a relatively large specific surface area 123.5 m² g⁻¹, which is presumed to be benefited from the two-step calcination of the template MIL-100(Fe). On the other hand, the pore size distribution reveals that most of the pores are focused on 10 nm, indicating the mesopores are effective formed for the γ-Fe2O3.

Of interest is using MOF thermolysis to develop the porous iron oxides. As an example, we note the fabrication of spindle-like mesoporous α-Fe2O3 using MOF MIL-88(Fe) as template. The prepared spindle-like mesoporous α-Fe2O3 was composed of clustered Fe2O3 nanoparticles with size of ~20 nm. In another example, a facile synthesis of porous Fe2O3 nanocubes by simultaneous oxidative decomposition of Prussian blue nanocubes was demonstrated. The derived porous Fe2O3 nanocubes are composed of very fine Fe2O3 nanoparticles with size of several nanometers. It is important that MOFs show distinct advantages due to their well-ordered crystalline
structure, high porosity, large surface area, and tunable pore size, in comparison with other templates.

It should be noted that during the MOF pyrolysis it can also be prepared metal oxide nanostructure consisting of the same primary particles, but with a different secondary and tertiary architecture thus modulating the macroscopic structure of the initial CP. For example, two types of Co$_3$O$_4$ nanomaterials with plate-like and rod-like morphology of the nanoparticles, based on the Co-containing plate-shaped ([Co$_8$(L)$_3$(DMF)$_2$]$_{16}$, p-MOF and rod-shaped ([Co(L)(DMSO)]$_n$, r-MOF) CPs, consisting of the same composite units (Co$^{2+}$ and 1,4-benzenedicarboxylate (L)), were prepared through pseudomorphic transformation.\textsuperscript{238}

Thermolysis of nano plates of two Cd(II)-based MOFs, [Cd$_2$(L)$_2$(L')$_2$ (DMF)$_n$ (TMU-8)] and [Cd(L)(4,4'-bipy)$_n$ (DMF)$_n$ (TMU-9)] where H$_2$L is V-shaped flexible dicarboxylate ligand 4,4'-oxybis(benzoic acid) and the L' is N-donor ligand 1,4-bis(4-pyridyl)-2,3-diaza-1,3-butadiene, leads to CdO nano-sized particles (Fig. 34).\textsuperscript{339} SEM images of the products exhibit the formation of aggregation of CdO nano-sized particle in the range of 60–160 nm. Particle sizes of CdO prepared by thermolysis of TMU-9 are smaller than those prepared from calcination of TMU-8 that is associated with the influence of the structures of these two MOFs on the final morphologies of CdO particles.\textsuperscript{23}

The 1D CP, [Zn(4,4'-bpy)(H$_2$O)$_4$]$_n$(H$_2$L = acetylene-dicarboxylic acid), and 3D MOF, Zn(L)$_2$: L'$_2$ (L’ is triethylamine) were applied as precursors for the synthesis of ZnO nanocrystals via direct thermal decomposition.\textsuperscript{34} Interestingly, changing the precursor from 1D CP to 3D MOF changes the morphology from nanorods to nanoparticles.

Thermolysis of the bulk powder and nano-sized Ni(II) MOF with cubic building units and 1D open channels, i.e. [Ni$_{16}$ (Ni$_4$L$_{12}$)(H$_2$O)$_{20}$(H$_2$O)$_4$]$_n$ (H$_2$L = 4,5-imidazoledicarboxylic acid), at 700 °C under air yielded NiO nanoparticles, which size and morphology depends on the particle size of the initial compound.\textsuperscript{341} The crystalline and nano-structures of two Pb(n) CPs, [Pb$_2$(L)$_4$(MeOH)]$_n$ (L = 2-pyrazinacarboxylic acid) and [Pb(L)$_2$]$_n$ (L’ = 2-quinolinencarboxylic acid), were thermoloyzed to prepare PbO nanoparticles with various morphology and size.\textsuperscript{234}

It should be noted that N,S-containing ligands were used not only to prepare metal sulfide nanoparticles, as expected, but also metal oxides, more thermodynamically stable in air. Thus, a ZnO nanostructure was prepared by thermolysis of nanoparticles of a Zn(II) CP, [Zn(L)(CH$_3$COO)]$_n$ (L = 2,2'-diamino-5,5'-dimethyl-1,4'-bithiazole), at 500 °C under air.\textsuperscript{342}

Saving macroscopic MOFs structure during their thermal transformation may serve as a promising strategy in the design of nanomaterials with predetermined properties and morphology.

5.2.2. Influence of temperature. Temperature is an important parameter controlling the phase composition and morphology of the resulting nanostructures. Thus, controlled thermolysis of [Mn$_2$(L)(DMA)]$_n$ (L is meso-butane-1,2,3,4-tetracarboxylate, DMA is N,N-dimethylacetamide)\textsuperscript{343} and Mn$_3$(L') (L’ is 2,5-dihydroxyterephthalate)\textsuperscript{344} by varying the temperature and the atmosphere leads to the formation of a series of nanoporous manganese oxides with different degrees of oxidation: MnO, Mn$_2$O$_4$, Mn$_3$O$_8$, and Mn$_2$O$_3$ and preserving the original morphology of the starting precursor (Fig. 35). In addition, varying the thermolysis temperature (300–700 °C) of Mn$_3$(L')\textsuperscript{344} leads to reduction of lattice defects, increasing the crystallinity degree of the thermolysis products. At the same time, the crystal surface becomes uneven and a plurality of voids appears with temperature increasing. The prepared MnO are characterized by type IV adsorption isotherms, indicating their mesoporous structure. It should be noted that in this way
rare low-valence oxides MnO and metastable phases of Mn$_2$O$_8$ with nanoporous architecture can be obtained, i.e. MOF thermal transformation can significantly enrich the composition, structure and functional properties of the resulting nanoporous metal oxide materials.

Similar patterns are characteristic of thermal transformations of MIL-100(Fe).\textsuperscript{346} The average size of nanorods Fe$_2$O$_3$, obtained at temperatures of 350, 550 and 750 °C is 50, 150 and 200 nm, respectively. Character of adsorption-desorption isotherms in this case is consistent with the general trend of the formation of the mesoporous structure of metal oxide nanocomposites obtained by MOF thermolysis. As a rule, surface area, pore volume, and pore size decreases with increasing thermolysis temperature, which also is in agreement with the fact that the thermally controlled crystal growth reduces the porosity with increasing process temperature.

Thermolysis temperatures of two silver\textsuperscript{[i]} CPs, [Ag(HQ)(Q)], and \{[Ag(HQ)$_2$]NO$_3$\}, has effect on the formation of silver nanomaterials.\textsuperscript{347} In particular, increasing temperature leads to increase of agglomeration of silver nanoparticles. Calcination of sinusoidal CP [Ag(L)](ClO$_4$), where L is a diethylbis(4-pyridyl)silane, at 400 °C gives circle morphology with evolving burned organics. At the same time, at 600 °C, network circles involving a silver(0)/silver chloride (chlorargyrite)/silicon(IV) oxide composite with a micro-sized convexo-concave surface are formed.\textsuperscript{348} However, thermolysis of [Ag(L)](BF$_4$) crystals at 600 °C gives silver(0) nanocomposites without silicon(IV) oxide.

It should be noted the zinc(II) oxide nanoparticles prepared by thermolysis of \{Zn(L)$_2$(H$_2$O)$_3$\}, \{Zn(L’)$_2$(H$_2$O)$_3$\}, and [Zn(H$_2$O)$_6$]$(L’)_2$, where L is nicotinic acid N-oxide, L’ is picolinic acid N-oxide and L’’ is isonicotinic acid N-oxide, at two temperatures (200 and 600 °C) using two methods (with surfactant and without surfactant).\textsuperscript{349} Analysis of the SEM images of the prepared ZnO nanoparticles exhibits that small and spherical ZnO nanoparticles were prepared by thermolysis of compounds at 200 °C using surfactant (oleic acid).

5.2.3. Influence of surfactants. It should be noted using surfactants in the synthesis of nanomaterials to change the morphology and particle size. For example, Zn(II) CPs were used for the preparation of nano-sized materials in the presence of surfactants.\textsuperscript{350,351}

Pure phase ZnO nanoparticles with various morphologies and sizes were synthesized by thermolysis of zinc(II) CPs, [Zn(L)(OAc)]ClO$_4$, [Zn(L)I$_2$], and [Zn(L)$_3$](ClO$_4$)$_2$(H$_2$O)$_2$, based on the same ligand L.\textsuperscript{352}

Of interest is the synthesis of PbO nanoparticles (~30 nm) by thermolysis of a Pb(II) complex, \{Pb$_2$(L)$_3$(NO$_3$)$_2$\}:MeOH, where L is $N,N',N''$-tris(pyrid-3-ylmethyl)-1,3,5-benzenetricarboxamide, at 180 °C with oleic acid as a surfactant.\textsuperscript{353}

We note also PbO nanoparticles (~25 nm) obtained by thermolysis of plate-shaped nanostructures of a 1D lead(II) CP involving the Pb$_2$(μ-I)$_2$ fragment, \{Pb(L)$_3$\}(L = neocuproine or 2,9-dimethyl-phen), with oleic acid as a surfactant.\textsuperscript{354}

5.3. Mixed-oxide nanocomposites

Heterometallic chelates are of interest for preparation of functional oxide materials (e.g. metal-oxide ceramics). The main advantages of using heterometallic chelates as precursors of
nanomaterials are low temperature of thermolysis, the exact ratio of metal ions, the formed nanoparticle size control, etc.

As a typical example, we consider using mixed lanthanide CPs for the production of rare earth oxides.\textsuperscript{335} In particular, well-defined cylindrical or spherical micro-morphologies can be prepared by varying solvents (water or THF) which were discovered in the resulting oxides after thermal treatment. Such bimetallic CPs were synthesized by the interaction of chlorides or nitrates of Ce, Nd, Gd with 2,5-dihydroxy-1,4-benzoquinone and used as precursors for mixed oxide ceramics. Thermolysis at 850 °C of the CPs leads to a mixture of BaCuO\textsubscript{3} and CuO. It is important that the thermolysis conditions have effect on the phase compositions of nanoparticles. Mesoporous nickel ferrite spheres were obtained after the spheres were calcined at 300 °C in air to remove the structure-directing agent and increase crystallinity. Mesoporous NiFe\textsubscript{2}O\textsubscript{4} spheres of high specific surface area (278 m\textsuperscript{2} g\textsuperscript{-1}) with a highly crystalline framework were prepared by adjusting the amount of structure-directing agent and the calcining condition.

6. Concluding remarks and outlook

As the contents of this review show, the current stage of development of CPs with chelated units has reached its peak in the accumulation of experimental facts and their theoretical
interpretation and generalization; all major groups of researchers are involved in this area of science.

How we see development of the chemistry of coordination polymers with chelated units?

Firstly, it is the search for more “precision” of the directed synthesis of the polytopic ligands containing chelating sites with controlled flexibility, allowing design of the unstrained metalallocycles. In this regard, structural homogeneity of CPs with chelated units and the lack of functional groups capable of forming other coordination centers are important. It is important to develop CPs with chelated units, chelating centers which include non-traditional heteroatoms. The optimization of the synthesis methods of CPs with chelated units will continue by reducing the number of synthetic steps (for example, by combining the synthesis of chelating polytopic ligands and CPs on their basis, etc.).

It should be noted that the CPs with chelated units make important contribution to developing the chemistry of polymeric metal chelates. Among the most important tasks in this direction are the continued development and optimization of low cost, high-yielding synthetic routes toward final CP with minimal impurities or contaminations; the continued introduction of new “green” chemistries and solvent-free or aqueous solution of preparation processes (in the absence of organic solvents) into synthetic methodologies of CP, which would minimize the cost and environmental impact; control over site-specific functionality (for example, core–shell–periphery); design of shape/morphology changing CP on application of an external stimulus; further investigation into CP for targeted biological applications.

The promising way will be further development of nanomaterials preparation by thermolysis of CPs. This is an intensely developed field of chemistry and materials science, which possesses its objects and investigation methods. Information in this field is accumulated continuously collecting the facts that seemed strange earlier and completing logical structuring of new ways of production of nanomaterials with high marketing potential. Nevertheless, till now, as a rule, it is impossible to recognize correlations between content, structural features, and properties of the nanomaterials prepared by this method, which in many respects restrains scientifically grounded approach to structuring of these materials and prediction of their promising properties.

Conflicts of interest

No potential conflict of interest was reported by the authors.

References

1 K. Naka, Metal Organic Framework (MOF), in Encyclopedia of Polymeric Nanomaterials, ed. S. Kobayashi and K. Müllen, Springer, 2015.

2 Metal-Organic Frameworks: Materials Modeling towards Engineering Applications, ed. J. Jiang, CRC, 2015.

3 R. Zhai, F. Jiao, H. Lin, F. Hao, J. Li, H. Yan, N. Li, H. Wang, Z. Jin, Y. Zhang and O. Qian, Chin. J. Chromatogr., 2014, 32, 107–116.

4 Metal–Organic Framework Materials, ed. L. R. MacGillivray and C. M. Lukehart, Wiley, 2014.

5 H.-C. Zhou, J. R. Long and O. M. Yaghi, Chem. Rev., 2012, 112, 673–674.

6 W. Lu, Z. Wei, Z.-Y. Gu, T.-F. Liu, J. Park, J. Park, J. Tian, M. Zhang, Q. Zhang, T. Gentle III, M. Bosch and H.-C. Zhou, Chem. Soc. Rev., 2014, 43, 5561–5593.

7 O. L. Ortiz and L. D. Ramirez, Coordination Polymers and Metal Organic Frameworks: Properties, Types, and Applications, Nova Science Publishers, 2012.

8 M. O’Keeffe and O. M. Yaghi, Chem. Rev., 2012, 112, 675–702.

9 S.-L. Wu, J.-P. Zou, M.-H. Chen, H.-B. Yang, M.-J. Li, X.-B. Luo, F. Luo, M.-F. Wu and G.-C. Guo, Polyhedron, 2012, 48, 58–67.

10 R. Haldar, N. Sikdar and T. K. Maji, Mater. Today, 2015, 18, 97–116.

11 K. E. Knope, D. T. Lill, C. E. Rowland, P. M. Cantos, A. Bettencourt-Dias and C. L. Cahill, Inorg. Chem., 2012, 51, 201–206.

12 W. Q. Kan, J. F. Ma, Y. Y. Liu and J. Yang, CrystEngComm, 2012, 14, 2316–2326.

13 J.-R. Li, J. Sculley and H.-C. Zhou, Chem. Rev., 2012, 112, 869–932.

14 H. Wu, Q. Gong, D. H. Olson and J. Li, Chem. Rev., 2012, 112, 836–868.

15 K. Sumida, D. L. Rogow, J. A. Mason, T. M. McDonald, E. D. Bloch, Z. R. Herm, T.-H. Bae and J. R. Long, Chem. Rev., 2012, 112, 724–781.

16 M. P. Suh, H. J. Park, T. K. Prasad and D.-W. Lim, Chem. Rev., 2012, 112, 782–835.

17 Y. He, W. Zhou, G. Qian and B. Chen, Chem. Soc. Rev., 2014, 43, 5657–5678.

18 S. Qiu, M. Xue and G. Zhu, Chem. Soc. Rev., 2014, 43, 6116–6140.

19 H.-J. Son, S. Jin, S. Patwardhan, S. J. Wezenberg, N. C. Jeong, M. So, C. E. Wilmer, A. A. Sarjeant, G. C. Schatz, R. Q. Snurr, O. K. Farha, G. P. Wiederrecht and J. T. Hupp, J. Am. Chem. Soc., 2013, 135, 862–869.

20 S. Jin, H.-J. Son, O. K. Farha, G. P. Wiederrecht and J. T. Hupp, J. Am. Chem. Soc., 2013, 135, 955–958.

21 P. Horcajada, R. Gref, T. Baati, P. K. Allan, G. Maurin, P. Couvreur, G. Férey, R. E. Morris and C. Serre, Chem. Rev., 2012, 112, 1232–1268.

22 W. Morris, B. Voloskiy, S. Demir, F. Gándara, P. L. McGrler, H. Furukawa, D. Cascio, J. F. Stoddart and O. M. Yaghi, Inorg. Chem., 2012, 51, 6443–6445.

23 J. Liu, P. K. Thallapally, B. P. McGrail, D. R. Brown and J. Liu, Chem. Soc. Rev., 2012, 41, 2308–2322.

24 C. He, D. Liu and W. Lin, Chem. Rev., 2015, 115, 11079–11108.

25 K. Lu, C. He and W. Lin, J. Am. Chem. Soc., 2015, 137, 7600–7603.
26 Z. Hu, B. J. Deibert and J. Li, Chem. Soc. Rev., 2014, 43, 5815–5840.
27 L. E. Kreno, K. Leong, O. K. Farha, M. Allendorf, R. P. Van Duyne and J. T. Hupp, Chem. Rev., 2012, 112, 1105–1125.
28 Y. Cui, Y. Yue, G. Qian and B. Chen, Chem. Rev., 2012, 112, 1126–1162.
29 J. Heine and K. Muller-Buschbaum, Chem. Soc. Rev., 2013, 42, 9232–9242.
30 M. Yoon, R. Srilambalaji and K. Kim, Chem. Rev., 2012, 112, 1196–1231.
31 J. Liu, L. Chen, H. Cui, J. Zhang, L. Zhang and C.-Y. Su, Chem. Soc. Rev., 2014, 43, 6011–6061.
32 P. Phuengphai, S. Youngme, P. Gamez and J. Reedijk, Dalton Trans., 2010, 39, 7936–7942.
33 T. Zhang and W. Lin, Chem. Soc. Rev., 2014, 43, 5982–5993.
34 A. H. Chughtai, N. Ahmad, H. A. Younus, A. Laypkov and F. Verpoort, Chem. Commun., 2015, 44, 6804–6849.
35 J. Canivet, S. Aгуado, Y. Schuurman and D. Farrusseng, J. Am. Chem. Soc., 2013, 135, 4195–4198.
36 A. N. Mlinar, B. K. Keitz, D. Gygi, E. D. Bloch, J. R. Long and A. T. Bell, ACS Catal., 2014, 4, 717–721.
37 Y. Zhang, Y. Zhou, Y. Zhao and C. Liu, Catal. Today, 2016, 263, 61–68.
38 A. Bhunia, S. Dey, J. M. Moreno, U. Diaz, P. Concepcion, K. Van Hecke, C. Janiak and P. Van Der Voort, Chem. Commun., 2016, 52, 1401–1404.
39 S. Ou and C.-D. Wu, Inorg. Chem. Front., 2014, 1, 721–734.
40 P. Ramaswamy, N. E. Wong and G. K. H. Shimizu, Chem. Soc. Rev., 2014, 43, 5913–5932.
41 C.-Y. Sun, X.-L. Wang, X. Zhang, C. Qin, P. Li, Z.-M. Su, D.-X. Zhu, G.-G. Shan, K.-Z. Shao, H. Wu and J. Li, Nat. Commun., 2013, 4, 2717.
42 Y. Cui, F. Zhu, B. Chen and G. Qian, Chem. Commun., 2015, 51, 7420–7431.
43 J. Pan, C.-P. Liu, F.-L. Jiang, M.-Y. Wu, L. Chen, J.-J. Qian, K.-Z. Su, X.-Y. Wan and M.-C. Hong, CrystEngComm, 2015, 17, 1541–1548.
44 X.-P. Wang, T.-P. Hu and D. Sun, CrystEngComm, 2015, 17, 3393–3417.
45 A. D. Pomogailo and G. I. Dzhhardimalieva, Nanostructured Materials Preparation via Condensation Ways, Springer, 2014.
46 A. D. Pomogailo and G. I. Dzhhardimalieva, Metallopolymetric Hybrid Nanocomposites, Nauka, 2015.
47 B. I. Kharisov, O. V. Kharissova and U. O. Méndez, J. Coord. Chem., 2013, 66, 3791–3828.
48 O. Carp, Materials Obtained by Solid-State Thermal Decomposition of Coordination Compounds and Metal-Organic Coordination Polymers, in Reactions and Mechanisms in Thermal Analysis of Advanced Materials, ed. A. Tiwari and B. Raj, Scrivener Publishing LLC, 2015.
49 Y. Song, X. Li, L. Sun and L. Wang, RSC Adv., 2015, 5, 7267–7279.
50 J.-K. Sun and Q. Xu, Energy Environ. Sci., 2014, 7, 2071–2100.
51 B. I. Kharisov, H. V. Rasika Dias, O. V. Kharissova, V. M. Jiménez-Pérez, B. O. Pérez and B. M. Flores, RSC Adv., 2012, 2, 9325–9358.
52 M. Y. Masoomi and A. Morsali, Coord. Chem. Rev., 2012, 256, 2921–2943.
53 A. D. Pomogailo and I. E. Uflyand, Adv. Polym. Sci., 1990, 97, 61–105.
54 A. D. Pomogailo and I. E. Uflyand, Makromolekulaynye Metallokholiety Macromolecular Metal Chelates, Khimiya, 1991.
55 G. Kumar and R. Gupta, Chem. Soc. Rev., 2013, 42, 9403–9453.
56 D. Feng, Z.-Y. Gu, J.-R. Li, H.-L. Jiang, Z. Wei and H.-C. Zhou, Angew. Chem., Int. Ed., 2012, 51, 10307–10310.
57 Y. Chen, T. Hoang and S. Ma, Inorg. Chem., 2012, 51, 12600–12602.
58 X.-S. Wang, M. Chrzanowski, C. Kim, W.-Y. Gao, L. Wojtas, Y.-S. Chen, X. P. Zhang and S. Ma, Chem. Commun., 2012, 48, 7173–7175.
59 X.-S. Wang, M. Chrzanowski, W.-Y. Gao, L. Wojtas, Y.-S. Chen, M. J. Zaworotko and S. Ma, Chem. Sci., 2012, 3, 2823–2827.
60 X.-S. Wang, M. Chrzanowski, L. Wojtas, Y.-S. Chen and S. Ma, Chem.–Eur. J., 2013, 19, 3297–3301.
61 L. Meng, Q. Cheng, C. Kim, W.-Y. Gao, L. Wojtas, Y.-S. Cheng, M. J. Zaworotko, X. P. Zhang and S. Ma, Angew. Chem., Int. Ed., 2012, 51, 10082–10085.
62 X.-L. Yang, M.-H. Xie, C. Zou, Y. He, B. Chen, M. O’Keeffe and C.-D. Wu, J. Am. Chem. Soc., 2012, 134, 10638–10645.
63 A. Fateeva, P. A. Chater, C. P. Ireland, A. A. Tahir, Y. Z. Khimyak, P. V. Wiper, J. R. Darwent and M. J. Rosseinsky, Angew. Chem., Int. Ed., 2012, 51, 7440–7444.
64 V. S. P. K. Neti, A. J. Metta-Magaña and L. Echegoyen, J. Coord. Chem., 2013, 66, 3193–3198.
65 D. Feng, H.-L. Jiang, Y.-P. Chen, Z.-Y. Gu, Z. Wei and H.-C. Zhou, Inorg. Chem., 2013, 52, 12661–12667.
66 J. A. Johnson, X. Zhang, T. C. Reeson, Y. S. Chen and J. Zhang, J. Am. Chem. Soc., 2014, 136, 15881–15884.
67 B. J. Burnett and W. Choe, CrystEngComm, 2012, 14, 6129–6131.
68 J. M. Falkowski, T. Sawano, T. Zhang, G. Tsun, Y. Chen, J. V. Lockard and W. Lin, J. Am. Chem. Soc., 2014, 136, 5213–5216.
69 Z. Zhang, L. Wojtas, M. Eddaoudi and M. J. Zaworotko, J. Am. Chem. Soc., 2013, 135, 5982–5983.
70 Z. Zhang and M. J. Zaworotko, Chem. Soc. Rev., 2014, 43, 5444–5455.
71 Z. Zhang, L. Zhang, L. Wojtas, P. Nugent, M. Eddaoudi and M. J. Zaworotko, J. Am. Chem. Soc., 2012, 134, 924–927.
72 Z. Zhang, L. Zhang, L. Wojtas, M. Eddaoudi and M. J. Zaworotko, J. Am. Chem. Soc., 2012, 134, 928–933.
73 A. Fidalgo-Marijuan, G. Barandika, B. Bazán, M.-K. Urtiaga and M. I. Arriortua, CrystEngComm, 2013, 15, 4181–4188.
74 L. Xu, J. Wang, Y. Xu, Z. Zhang, P. Lu, M. Fang, S. Li, P. Sun and H.-K. Liu, CrystEngComm, 2014, 16, 8656–8659.
326 R. Gupta, S. Sanotra, H. Nawaz Sheikh, B. Lal Kalsotra, V. Kumar Gupta and Rajnikant, J. Coord. Chem., 2012, 65, 3917–3931.
327 L. Wang, H. Zou, Y. Li, X. Li and S. Zhong, Mater. Manuf. Processes, 2017, 32, 484–488.
328 S. Jin, J. P. Hill, Q. Ji, L. Kumar Shrestha and K. Ariga, J. Mater. Chem. A, 2016, 4, 5737–5746.
329 L. Hashemi and A. Morsali, J. Inorg. Organomet. Polym., 2012, 22, 272–277.
330 M. A. Alavi, A. Morsali, S. W. Joo and B. K. Min, Ultrason. Sonochem., 2015, 22, 349–358.
331 A. Mehrani, A. Morsali, Y. Hanifehpour and S. W. Joo, Ultrason. Sonochem., 2014, 21, 1430–1434.
332 S. Hei, Y. Jin and F. Zhang, J. Chem., 2014, 2014, 546956.
333 F. Meng, Z. Fang, Z. Li, W. Xu, M. Wang, Y. Liu, J. Zhang, W. Wang, D. Zhao and X. Guo, J. Mater. Chem. A, 2013, 1, 7235–7241.
334 N. Nasihat Sheno, A. Morsali and S. Woo Joo, Mater. Lett., 2014, 117, 31–33.
335 M. Hu, A. A. Belik, M. Imura, K. Mibu, Y. Tsujimoto and Y. Yamauchi, Chem. Mater., 2012, 24, 2698–2707.
336 L. Zhang, H. B. Wu, S. Madhavi, H. H. Hng and X. W. Lou, J. Am. Chem. Soc., 2012, 134, 17388–17391.
337 L. Zhang, H. B. Wu, R. Xu and X. W. Lou, CrystEngComm, 2013, 15, 9322–9335.
338 K. J. Lee, T.-H. Kim, T. K. Kim, J. H. Lee, H.-K. Song and H. R. Moon, J. Mater. Chem. A, 2014, 2, 14393–14400.
339 M. Y. Masoomi and A. Morsali, Ultrason. Sonochem., 2016, 28, 240–249.
340 A. Abbasi, M. Gharib and M. Najafi, J. Sci., Islamic Repub. Iran, 2016, 27, 217–221.
341 A. Tahmasian and A. Morsali, Inorg. Chim. Acta, 2012, 387, 327–331.
342 A. Hosseinion, S. Jabbari, A. Reza Mahjoub and M. Movahedi, J. Coord. Chem., 2012, 65, 2623–2633.
343 J. H. Lee, Y. J. Sa, T. K. Kim, H. R. Moon and S. H. Joo, J. Mater. Chem. A, 2014, 2, 10435–10443.
344 H. Jiang, C. Wang, H. Wang and M. Zhang, Mater. Lett., 2016, 168, 17–19.
345 T. K. Kim, K. J. Lee, J. Y. Cheon, J. H. Lee, H. S. Joo and H. R. Moon, J. Am. Chem. Soc., 2013, 135, 8940–8946.
346 Z.-M. Liu, S.-H. Wu, S.-Y. Jia, F.-X. Qin, S.-M. Zhou, H.-T. Ren, P. Na and Y. Liu, Mater. Lett., 2014, 132, 8–10.
347 K. Akhbari, N. B. Bahman, A. Morsali and P. Retailleau, J. Iran. Chem. Soc., 2016, 13, 165–169.
348 S. Y. Moon, M. W. Park, T. H. Noh and O.-S. Jung, J. Mol. Struct., 2013, 1054–1055, 326–332.
349 F. Marandi, L. Hashemi, A. Morsali and H. Krautscheid, Ultrason. Sonochem., 2016, 32, 86–94.
350 F. Shahangi Shirazi and K. Akhbari, Inorg. Chim. Acta, 2015, 436, 1–6.
351 M. Moeinian and K. Akhbari, J. Solid State Chem., 2015, 225, 459–463.
352 A. Mehrani, A. Morsali and P. Ebrahimpour, J. Coord. Chem., 2013, 66, 856–867.
353 G. H. Shahverdizadeh, F. Hakimi, B. Mirtamizdoust, A. Soudi and P. Hojati-Talemi, J. Inorg. Organomet. Polym., 2012, 22, 903–909.
354 B. Mirtamizdoust, B. Shaabani, A. Khandar, H. Pourradi, Y. Abbasytyula, H. Goudarziabshar, D. Viterbo, G. Croce and P. Hojati-Talemi, J. Inorg. Organomet. Polym., 2012, 22, 1293–1299.
355 T. Demars, M. Boltoeva, N. Vigier, J. Maynadié, J. Ravaux, C. Genre and D. Meyer, Eur. J. Inorg. Chem., 2012, 3875–3884.
356 Z. V. Dobrokhотова, N. V. Gogoleва, E. N. Зорина-Тихонова, М. А. Кискин, V. V. Chernyshev, A. L. Эмелина, M. A. Bukov, A. S. Goloveshkin, I. S. Bushmarинов, A. A. Сидоров, A. S. Богомьяков, M. L. Ковба, V. M. Novотортсев и I. L. Еременко, Eur. J. Inorg. Chem., 2015, 2015, 3116–3127.
357 A. V. Гавриков, P. S. Коротеев, Z. V. Dobrokhотова, A. B. Ильюхин, N. N. Ефимов, D. I. Кирдянкин, M. A. Bykov, M. A. Руимин и V. M. Новотортсев, Polyhedron, 2015, 102, 48–59.
358 N. Заузолькова, Z. Dobrokhотова, A. Лермонтов, E. Зорина, A. Эмелина, M. Bukov, V. Chernyshev, A. Сидоров, M. Кискин, A. Богомьяков, A. Лытвиненко, S. Котолитов, Y. Великодньи, M. Ковба, V. Новотортсев и I. Еременко, Solid State Chem., 2013, 197, 379–391.
359 M. A. Руимин, Z. V. Dobrokhотова, A. L. Эмелина, M. A. Bukov, N. V. Гоголева, K. S. Gavrichев, E. N. Зорина, Tikhonova, L. I. Demina, M. A. Kiskin, A. A. Sidorov, I. L. Eremenko and V. M. Novotortsev, Polyhedron, 2015, 87, 28–37.
360 D. Hong, Y. Yamada, M. Sheehan, S. Shikano, C.-H. Kuo, M. Tian, C.-K. Tsung and S. Fukuzumi, ACS Sustaina. Chem. Eng., 2014, 2, 2588–2594.
361 Y. Xia, B. Wang, G. Wang, X. Liu and H. Wang, ChemElectroChem, 2016, 3, 299–308.