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Homochiral layered indium phosphonates: solvent modulation of morphology and chiral discrimination adsorption†

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Assembling chiral coordination polymers into nano/microflower structures may improve their performance in applications such as chiral recognition and separation. In this study, we chose a chiral metal phosphonate system, *i.e.*, $In(NO_3)_3/R$ -, S-pempH₂ [pempH₂ = (1-phenylethylamino)methylphosphonic acid], and carried out systematic work on the self-assembly of this system in different alcohol/H₂O mixed solvents under solvethermal conditions. Enantiomeric compounds R-, S-[In_2 (pempH)₂(μ_2 -OH)₂(H_2 O)₂](In_2 (In_2) were obtained showing dense layered structures, but their morphologies varied with alcohol solvent. We obtained crystals of In_2 0 mixed solvent and spiral microflowers of In_2 1 mixed solvent and spiral microflowers of In_2 2 mixed solvent and spiral microflowers of In_2 3 mixed solvent and spiral microflowers of In_2 4 mixed solvent and spiral microflowers of In_2 5 mixed solvent and spiral microflowers of In_2 6 mixed solvent and spiral microflowers of In_2 6 mixed solvent and spiral microflowers of In_2 6 mixed solvent and In_2 6 mixed solvent and In_2 7 mixed solvent and In_2 8 mixed solvent and In_2 9 mixed solven

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Introduction

Chiral materials have attracted considerable attention owing to their potential applications in fields such as enantioselective separation, chiral sensing and asymmetric catalysis. 1-5 Chiral porous metal-organic frameworks (MOFs) or coordination polymers (CPs) are ideal candidates for chiral recognition and separation because their structures can be designed and modified to provide a pore environment for specific chiral guests.⁶⁻⁸ However, achieving precise and uniformly distributed pore structure characteristics in three-dimensional (3D) chiral MOFs is not only challenging, but also highly selective towards the size and shape of chiral guest molecules, thus restricting their adaptability to diverse chiral molecular systems. A feasible solution is to prepare two-dimensional (2D) chiral MOFs or CPs and further exfoliate them into nanosheets.9-11 The unique structural characteristics of nanosheets enable them to exhibit large surface areas, which greatly increases the adsorption sites of chiral MOF materials and allows them to be used for the recognition and separation of chiral molecules of different sizes. A notable example is given by Cui and coworkers, who fabricated nanosheets of Eu-MOF based on a chiral 1,1'-biphenyl phosphoric acid that can be used as chiral

ticular, the re-stacking of chiral nanosheets reduces the pore size, decreases the surface area, deteriorates the solubility, and ultimately results in the formation of bulk 3D materials. We envision that assembling nanosheets into nano-/micro-structures can effectively avoid the re-stacking of chiral nanosheets. In addition, whether the chirality can be transferred and amplified from the nanosheets to the macroscopic assemblies is also an extremely intriguing question.

Previous studies have shown that chiral materials with flower-like structures can exhibit macroscopic chirality, high specific surface area, and certain spatially restricted domains.

optical sensors for the detection of chiral terpenes and terpe-

noids.12 Nevertheless, despite significant progress in this

direction, challenges still remain in synthesizing 2D materials

with permanent chirality, good solubility and stability. In par-

Previous studies have shown that chiral materials with flower-like structures can exhibit macroscopic chirality, high specific surface area, and certain spatially restricted domains. As a result, both the chiroptical and selective adsorption properties of the material may be improved. ^{13–15} For instance, chiral CuO@CNC nanoflowers exhibit prominent circularly polarized luminescence (CPL) properties and can effectively induce enantioselective crystallization of L-threonine. ¹⁶ We conceived that assembling chiral CP nanosheets into nano/microflowers might improve the chiral recognition performance of the materials, but similar studies have never been reported.

In this work, we carried out a systematic investigation on the $In(NO_3)_3/R$ -, S-pempH $_2$ system, where R-, S-pempH $_2$ represent R-, S-(1-phenylethylamino)methylphosphonic acids. The In^{3+} ion and monophosphonate ligands were chosen because they can easily form dense layered structures and exhibit high thermal and water stability, akin to other metal phosphonates. $^{17-19}$ We performed the same reaction in

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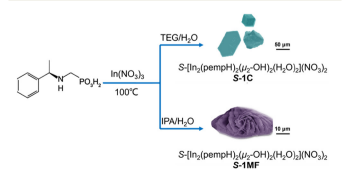
†Electronic supplementary information (ESI) available. CCDC 2402002. For ESI and crystallographic data in CIF or other electronic format see DOI: https://doi.org/10.1039/d4dt03227f

different alcohol/H2O solvents, where the alcohol can be methanol (MeOH), ethanol (EtOH), n-propanol (NPA), isopropanol (IPA), n-butanol (NBA), and triethylene glycol (TEG). In all cases, compounds R-, S- $[In_2(pempH)_2(\mu_2-OH)_2(H_2O)_2](NO_3)_2$ (R-, S-1) were obtained but with different morphologies. Specifically, we obtained crystals of R-, S-1 in TEG/H₂O mixed solvent (named R-, S-1C, where C refers to crystal) and spiral microflowers in IPA/H₂O mixed solvent (named R-, S-1MF, where MF refers to microflower) (Scheme 1). The acquisition of **R-, S-1MF** spiral microflowers is impressive because it implies that the chirality has been successfully transferred and amplified from the chiral ligand to the chiral layer and eventually to the macroscopic morphology. We further investigated the formation mechanism of R-, S-1MF and their chiral adsorption properties. It is noteworthy that although many chiral metal phosphonates are known in the literature, 20 chiral indium phosphonates have never been reported. Therefore, this work not only reports the first example of chiral indium phosphonates, but also the first example of chiral CPs with a nano/ microflower morphology.

Experimental

Materials and methods

All starting materials were of analytical reagent grade and used received without further purification. Phenylethylamino)methylphosphonic acids (R-, S-pempH₂) were prepared according to methods reported in the literature.^{21,22} Elemental analyses of C, N and H were performed with an Elementar vario Macro cube. Fourier transform infrared (IR) spectra were recorded using KBr pellets on a Bruker Tensor 27 cm^{-1} . spectrometer in the range of 400-4000 Thermogravimetric analysis (TGA) was performed on a Mettler-Toledo TGA/DSC instrument in the range of 30-600 °C under a nitrogen flow at a heating rate of 5 °C min⁻¹. Powder X-ray diffraction (PXRD) data were recorded on a Bruker D8 ADVANCE X-ray powder diffractometer (Cu-Kα) at room temperature. The scanning electron microscope (SEM) images were recorded on a Hitachi S-4800. Transmission electron microscopy (TEM) measurements were performed on a JEM-2100. UV-vis spectra



Scheme 1 Synthetic routes to S-1C and S-1MF in mixtures of TEG/ H_2O and IPA/ H_2O .

were recorded on a Shimadzu UV-3600 spectrometer. Electronic circular dichroism (ECD) spectra were recorded on a JASCO J-810 spectrophotometer at room temperature. The gas absorption and desorption isotherms were recorded on a BELSORP-Max adsorption analyzer in easy mode.

Single crystal X-ray crystallography

A suitable single crystal of *S*-1C was mounted on a loop ring, and diffraction data were collected using a Bruker D8 Venture diffractometer (Ga-K α radiation, λ = 1.34138 Å). The collected data were integrated using the Siemens SAINT program, ²³ and adsorption corrections were applied. The structures were solved using the intrinsic phasing method and refined on F^2 using full-matrix least-squares with the SHELXTL software package. ²⁴ All the non-hydrogen atoms were refined anisotropically, while hydrogen atoms bound to carbon atoms and oxygen atoms of hydroxide were refined isotropically in the riding mode. CCDC for *S*-1C (no. 2402002†) contains the supplementary crystallographic data for this paper. Details of the crystal data and refinements of *S*-1C are summarized in Table S1,† and selected bond lengths and angles of *S*-1C in Table S2.†

Synthesis of R-, S-[In₂(pempH)₂(μ_2 -OH)₂(H₂O)₂](NO₃)₂ crystals (R-, S-1C)

Compounds R-1C and S-1C were synthesized in a similar way except for the different enantiomers of the ligands, so the synthesis of S-1C is described as a representative. S-pempH₂ (21.5 mg, 0.1 mmol) was added to 5 mL deionized water in a 20 mL glass vial. After heating at 100 °C for 3 hours, a clear solution was obtained because of the dissolution of the ligand. Then, 100 µL triethylene diamine solution (4 M) was added to adjust the pH to 3.2, followed by the addition of 5 mL TEG. After the solution was mixed uniformly, In(NO₃)₃·5H₂O (60 mg, 0.2 mmol) was added and the glass vial was sealed. The mixture was placed at 100 °C for 24 hours, and transparent block-like crystals of S-1C were obtained. Yield: 16 mg, 74.4% (based on S-pempH₂). Elemental analysis calcd (%) for $C_{18}H_{32}In_2N_4O_{16}P_2$ (S-1C): C, 25.37; H, 3.79; N, 6.58%; found: C, 25.18; H, 3.88; N, 6.50%. IR (KBr, cm⁻¹): 3575(m), 3535(m), 3041(w), 2991(w), 2950(w), 2786(w), 2540(w), 2424(w), 2362(w), 1760(w), 1604(m), 1499(w), 1383(m), 1340(m), 1279(w), 1209(w), 1151(m), 1124(m), 1082(m), 1034(m), 995(m), 820(w), 788(w), 762(w), 702(w), 576(w), 492(w). For **R-1C**, yield: 14 mg, 65.1% (based on R-pempH₂). Elemental analysis calcd (%) for $C_{18}H_{32}In_2N_4O_{16}P_2$ (**R-1C**): C, 25.37; H, 3.79; N, 6.58%; found: C, 25.09; H, 3.90; N, 6.49%. IR (KBr, cm⁻¹): 3572(m), 3537(m), 3039(w), 2988(w), 2942(w), 2785(w), 2537(w), 2424(w), 2364(w), 1760(w), 1605(m), 1500(w), 1386(m), 1342(m), 1277(w), 1207(w), 1154(m), 1124(m), 1081(m), 1033(m), 996(m), 823(w), 785(w), 763(w), 703(w), 572(w), 490(w).

Synthesis of R-,S-[In₂(pempH)₂(μ_2 -OH)₂(H₂O)₂](NO₃)₂ microflowers (R-, S-1MF)

Compounds *R*-1MF and *S*-1MF were synthesized in a similar way except for the different enantiomers of the ligands, so the synthesis of *S*-1MF is described as a representative. *S*-1MF was

synthesized in the same manner as S-1C, except that the reaction solution was 90 vol% IPA/H₂O. Yield: 19 mg, 88.4% (based on S-pempH2). Elemental analysis calcd (%) for $C_{18}H_{32}In_2N_4O_{16}P_2$ (S-1MF): C, 25.37; H, 3.79; N, 6.58%; found: C, 25.37; H, 3.89; N, 6.39%. IR (KBr, cm⁻¹): 3572(m), 3536(m), 3037(w), 2990(w), 2950(w), 2788(w), 2533(w), 2424(w), 2362(w), 1760(w), 1608(m), 1497(w), 1385(m), 1337(m), 1277(w), 1207(w), 1156(m), 1124(m), 1081(m), 1032(m), 997(m), 826(w), 788(w), 756(w), 701(w), 574(w), 490(w). For *R*-1MF, yield: 17 mg, 79.1% (based on R-pempH₂). Elemental analysis calcd (%) for C₁₈H₃₂In₂N₄O₁₆P₂ (*R*-1MF): C, 25.37; H, 3.79; N, 6.58%; found: C, 25.70; H, 3.90; N, 6.41%. IR (KBr, cm⁻¹): 3574(m), 3538(m), 3036(w), 2988(w), 2948(w), 2785(w), 2536(w), 2426(w), 2358(w), 1758(w), 1605(m), 1498(w), 1386(m), 1337(m), 1278(w), 1210(w), 1154(m), 1126(m), 1085(m), 1032(m), 995(m), 818(w), 788(w), 758(w), 705(w), 577(w), 493(w).

Results and discussion

Synthesis and characterization of R-1C and S-1C crystals

We initially synthesized block-like crystals of R-, S-1C through a solvothermal reaction of $In(NO_3)_3 \cdot 5H_2O$ and R- or S-pempH $_2$ (molar ratio 2:1) in 50 vol% TEG/H $_2O$ at 100 °C. The phase purity of the bulk sample was confirmed by their powder X-ray diffraction (PXRD) patterns and elemental analyses (Fig. 1a). To examine the effect of the metal: ligand molar ratio and TEG/H $_2O$ solvent ratio on the reaction product, we conducted

similar experiments using the S-pempH $_2$ ligand. When the molar ratio was 1:1, we obtained the same material of S-1C as verified by the identical PXRD patterns (Fig. S1 \dagger), but the crystal quality was relatively poor. The reaction products obtained in 20–90 vol% TEG/H $_2$ O mixed solvents were also the same (Fig. S2 and S3 \dagger).

Compounds R-1C and S-1C exhibit identical PXRD patterns and IR and UV-vis spectra (Fig. 1a, b and S4-S6†). The UV-Vis spectra show absorption peaks at 225 nm and 252 nm, attributed to the $\pi \to \pi^*$ transitions of the benzene rings in the ligands, characteristic of the B band. 25 The solid-state circular dichroism (CD) spectra of R-1C and S-1C show mirror dichroic signals at ca. 230 nm, demonstrating their enantiomeric nature (Fig. 1c). We further conducted thermogravimetric analysis and found that the two compounds were stable up to ca. 150 °C under an N₂ atmosphere (Fig. S7†). The weight losses of R-1C and S-1C between 170 and 210 °C were 6.5% and 6.4%, respectively, which are close to the calculated value of 6.3% after removal of three water molecules. From the single crystal structural analysis given below, we noted that there are only two coordination water molecules in the structure, so the third water molecule should come from the coordinated µ-OH, which may reduce the stability of the layered structure. Indeed, the compounds continued to lose weight above 210 °C, with acceleration above 230 °C, accompanied by decomposition of the organic components and collapse of the structure.

Since R-1C and S-1C are enantiomers, we only measured the single crystal structure of S-1C. It crystallizes in the mono-

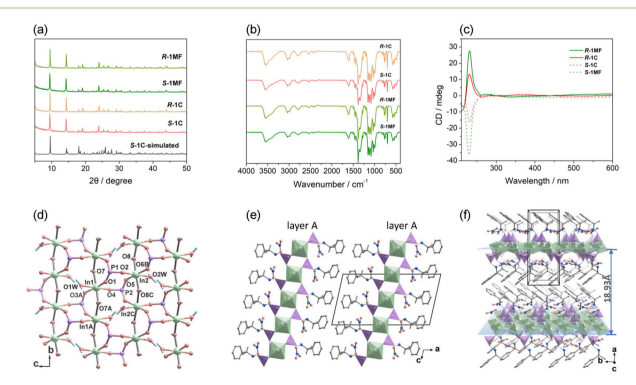


Fig. 1 (a) PXRD patterns of the as-synthesized compounds *R*-, *S*-1C and *R*-, *S*-1MF. The pattern simulated from the single-crystal data of *S*-1C is given for comparison. (b) The IR spectra of *R*, *S*-1C and *R*, *S*-1MF. (c) Solid-state CD spectra of *R*-, *S*-1C and *R*-, *S*-1MF. (d) The inorganic layer of *S*-1C. (e) Packing diagram of *S*-1C viewed along the *b*-axis. (f) Packing diagram of *S*-1C viewed along the *c*-axis.

clinic space group P2₁. The asymmetric unit contains two In³⁺ ions, two S-pempH⁻, two OH⁻, two NO₃⁻, and two coordinated water molecules (Fig. S8†). Each In3+ ion adopts six-coordination with an octahedral configuration. In 1 is coordinated by three phosphonate O atoms (O1, O3A, and O4) from three S-pempH⁻, one O atom (O1W) from one H₂O, and two μ_2 -OH⁻ (O7 and O7A). In2 is coordinated by three phosphonate O atoms (O2, O5, and O6B) from three S-pempH⁻, one O atom (O2W) from one H_2O , and two μ_2 -OH⁻ (O8 and O8C). The In1-O and In2-O bond lengths are 2.081-2.227 Å and 2.090-2.223 Å, respectively (Table S2†). The S-pempH ligand is protonated at its amino group and acts as a tri-dentate ligand using its three phosphonate oxygen atoms. Each S-pempH bridges three In atoms and vice versa, generating an inorganic layer composed of corner-sharing {In(μ₂-OH)₂O₄} octahedra and {PO₃C} tetrahedra in the bc plane (Fig. 1d). Interestingly, the equivalent In1 or In2 atoms are each bridged by μ_2 -OH⁻ units, forming infinite In-OH⁻ chains of -In1-O7-In1- and -In2-O8-In2-, respectively, which are alternatively arranged along the c-direction. Therefore, the inorganic layer of S-1C can also be viewed as consisting of two kinds of In-OH chains interconnected by PO₃ groups. The organic groups of the S-pempH⁻ ligands are suspended on either side of the inorganic layer (Fig. 1e). The hybrid layer is chiral and polar. The equivalent layers are stacked in an AAA manner and unidirectionally along the a-axis (Fig. 1e), thus leading to the formation of a polar supramolecular network structure (Fig. 1f). The distance between layers is 18.93 Å. Notably, the NO₃ counter-anions reside close to the inorganic layer and are H-bonded to the μ_2 -OH⁻ groups (O7 and O8), amino groups

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It is worth mentioning that only ten indium phosphonates have been described in the literature, $^{26-31}$ of which three are coordination polymers including two with 2D layered structures 26 and one with a 3D framework structure. 28 Compounds $In(O_3PR)(O_2P(OH)R)\cdot H_2O$ (R=Ph and $PhCH_2$) were supposed to have layered structures based on their PXRD patterns. 26 Notably, chiral indium phosphonates have never been reported. Therefore, compounds R-1C and S-1C represent the first examples of chiral indium phosphonates with a new layer topology.

(N1 and N2), and coordination water molecules (O1W and

O2W) from the same layer (Table S3 and Fig. S9†). The inter-

layer interactions are dominated by van der Waals contacts.

Solvent effect on the morphologies of In(NO₃)₃/pempH₂ assemblies: formation of *R*-1MF and *S*-1MF microflowers

Previous studies have demonstrated that solvent may have a significant effect on the morphology and even the helicity of chiral coordination polymers. Thus, we carried out a systematic study on the assembly of the $In(NO_3)_3/S$ -pempH₂ system using different polar alcohol solvents (MeOH, EtOH, IPA, NPA, and NBA) and alcohol/H₂O mixed solvents. We first conducted the reaction of $In(NO_3)_3 \cdot 5H_2O$ and S-pempH₂ (molar ratio, 1:1) in 20, 40, 60, 70, 80, 90, and 100 vol% MeOH/H₂O mixtures (total volume, 10 mL, pH = 3.2, adjusted using triethylene diamine) under solvothermal conditions

(100 °C, 1 day) and examined the solid products by scanning electron microscopy (SEM), PXRD, and IR spectroscopy (Fig. S10–S12†). PXRD and IR spectra clearly showed that the product obtained in pure methanol was the unreacted ligand, possibly due to the low solubility of *S*-pempH₂ in methanol. Although the products obtained in 20–90 vol% MeOH/H₂O had the same structure as *S*-1C, their morphologies were slightly different. The reaction in 20 vol% MeOH/H₂O produced nanoparticles, while that in 40–90 vol% MeOH/H₂O resulted in nanoplates. The nanoplates flocked together but no clear shape was observed.

We next performed the same reaction in $20{\text -}100$ vol% EtOH/H₂O mixtures (Fig. S13–S15†). Similarly, the product obtained in pure ethanol was dominated by the ligand, while those obtained in $20{\text -}90$ vol% EtOH/H₂O had the same structure as *S*-1C. From their SEM images we can see that the $20{\text -}60$ vol% products were aggregates of nanoplates/nanosheets with irregular shapes. Interestingly, we observed microparticles composed of aggregated nanosheets in the $80{\text -}90$ vol% products together with nanoparticles.

When the reaction of In(NO₃)₃·5H₂O and S-pempH₂ was conducted in 20–90 vol% NPA/H₂O or IPA/H₂O, we obtained the same product S-1C (Fig. S16–S21†). Although the products in 20–80 vol% NPA/H₂O were composed of stacked nanosheets with different sizes, the nanosheets in the 90 vol% product aggregated into circular flower-like structures (Fig. S16†). For products obtained in IPA/H₂O, the nanosheets were stacked randomly in the 20–70 vol% products, but formed microparticles with flower-like shapes in the 80–90 vol% products (Fig. S19†). Impressively, in the 90 vol% IPA/H₂O product, we observed uniformly distributed microparticles with an elliptical flower-like structure formed by the stacking of chiral nanosheets.

Finally, we performed the same reaction in 20–100 vol% NBA/H₂O mixed solvents. As shown in Fig. S22,† the 20–60 vol% products showed randomly distributed nanosheets with larger lateral dimensions. In the 70–90 vol% products, the nanosheets aggregated but no clear shape was observed. All products obtained in 20–90 vol% NBA/H₂O are the same as *S*-1C (Fig. S23 and S24†).

To sum up, the reaction of In(NO₃)₃·5H₂O and *S*-pempH₂ in 20–90 vol% alcohol/H₂O (alcohol = MeOH, EtOH, NPA, IPA, and NBA) resulted in the same compound as *S*-1C with different morphologies. A uniformly distributed product with a flower-like structure can be obtained in 90 vol% IPA/H₂O and was named *S*-1MF (MF refers to microflower). If *R*-pempH₂ instead of *S*-pempH₂ was used in a similar synthesis, we obtained the product *R*-1MF. As shown in Fig. 2, the *S*-1MF structures exhibit a highly branched and aggregated morphology with distinct left-handed spiral arrangements of the petal layers. In contrast, the *R*-1MF structures show a branched and aggregated morphology but feature right-handed spiral configurations of the petal layers.

Both types of microflowers have petals that are layered and relatively thin, with notable chirality in their spiral structures. *R*-1MF and *S*-1MF are enantiomers, verified by their identical

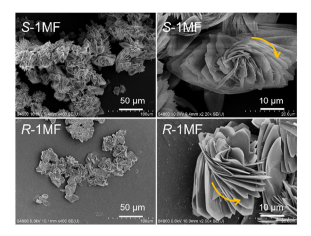


Fig. 2 SEM images of the self-assembled products of In(NO₃)₃/S-, R-pempH₂ in 90 vol% IPA/H₂O at 100 °C (S-1MF and R-1MF).

PXRD patterns and IR and UV-vis spectra (Fig. S4-S6†), as well as CD spectra, which show a dichroic signal at ca. 230 nm with an opposite sign (Fig. 1c).

Interestingly, the thickness of the nanosheets obtained in 90 vol% alcohol/H₂O varied with the solvent (Fig. 3). In 90 vol% MeOH/H2O, the nanosheets have a thickness of approximately 100-200 nm. In a 90 vol% EtOH/H2O system, the nanosheets exhibit a thickness of around 200-300 nm. The sample synthesized in 90 vol% NPA/H2O shows more compact and densely packed petals with an estimated thickness of 200-300 nm. The petals observed in the sample in 90 vol% IPA/H₂O are the thinnest, ranging from 50-100 nm. Adjusting the volume fraction of IPA has a significant impact on the size and aggregation of the microflowers. As the IPA ratio increases, the nanosheets become thinner, with reduced aggregation, ultimately leading to a uniformly distributed flower-like structure. In contrast, the nanosheets obtained in 90 vol% NBA/ H₂O are the thickest, with an estimated thickness of 300-400 nm. Despite these variations, all samples exhibit identical IR spectra and PXRD patterns (Fig. S25†).

Formation mechanism of S-1MF

In order to understand the formation mechanism of microflowers, we further investigated the changes of the In(NO₃)₃/ S-pempH₂ assemblies over time in 90 vol% IPA/H₂O at 100 °C. The reaction products of 15 to 60 min were clear solutions without any precipitates. However, TEM images revealed that dispersed sphere-like nanoparticles with sizes ranging from 10 to 30 nm appeared in the 15 min product (Fig. S26†), indicating the onset of the nucleation stage. By 30 min, the nanoparticles grew significantly to 50-100 nm and started to aggregate. At 60 min, they adopted non-spherical shapes with sizes of 300-500 nm. After 60 min of reaction, we observed precipitates, which were used for SEM measurements. As shown in Fig. 4a, the 90 min product contained irregularly shaped nanoparticles with sizes of approximately 500 nm to 1 µm. At 3 h, the particles appeared as irregularly shaped aggregates with noticeable surface textures and protrusions, reaching sizes of 1–5 μm. The nanosheet-like surface texture of the particles (size 1-8 µm) became more significant at 4 h. After 5 h of reaction, the morphology changes significantly, and microscopic structures made of stacked nanosheets can be clearly seen with sizes increasing to 5-10 µm. Loose flower-like aggregates formed after 6 h, with overall sizes of 10-20 µm. By 9 h, the aggregates became larger, forming distinct flower-like structures with overall sizes of 20-25 µm. By 18 h, the flower-like structures became more complex and distinct, with sizes of 25-30 µm and layered petal morphologies. Finally, at 24 h, the self-assembly process results in intricately layered flower-like structures (S-1MF), with overall sizes of 25-35 µm and petal thicknesses of approximately 100 nm, indicating the successful completion of the assembly process at this stage. Throughout the process, as time progresses, there is a clear trend toward increasing size, complexity, and order, with the

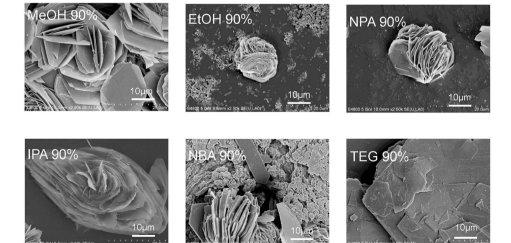


Fig. 3 SEM images of the self-assembled products of In(NO₃)₃/S-pempH₂ in 90 vol% alcohol/H₂O at 100 °C.

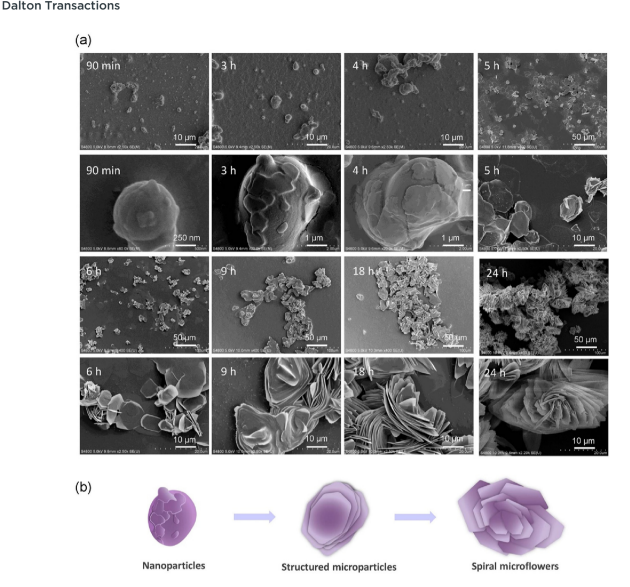


Fig. 4 (a) SEM images of the self-assembled products of $In(NO_3)_3/S$ -pempH₂ in 90 vol% IPA/H_2O at 100 °C for different reaction times. (b) Proposed mechanism for the formation of spiral microflowers.

final structures being highly organized and indicative of the meticulous self-assembly capabilities of the In(NO₃)₃/S-pempH₂ system under the given conditions. We conducted the PXRD and IR measurements for the solid products from reactions lasting between 90 min and 24 h. The results indicate that all products have the same structure as S-1C (Fig. S27 and S28†).

Based on the above observations, we hypothesize that the formation of microflowers of S-1MF occurred in three stages (Fig. 4b). First, In(NO₃)₃ and S-pempH₂ reacted in 90 vol% IPA/ H₂O to form sphere-like nanoparticles immediately; second, the nanoparticles grew into irregularly shaped micro-particles with the surfaces covered by stacked nanosheets; and third, further reaction and growth of the microparticles led to the formation of spiral microflowers. This process may be driven by surface energy minimization, anisotropic crystal growth, and specific reaction conditions, as suggested by others for the formation of flower-like structures. 35-39 In the present case, the indium phosphonate has a layered structure, in which the interlayer interactions are much weaker than the intralayer interactions. It is therefore to be expected that the In(NO₃)₃/ S-pempH₂ assemblies tend to grow anisotropically and form nanosheets, as found for other metal phosphonates. 40 The formation of spiral microflowers of S-1MF is unusual, which must be attributed to the solvent effect. Compared to other alcohols, the reaction of In(NO₃)₃ and S-pempH₂ in 90 vol% IPA/H₂O produced the thinnest nanosheets or petals. In addition, the interaction between the IPA solvent and the ligand may affect the orientation of the phenyl groups hanging outside the layers.³² All these factors may contribute to the chiral transcription and amplification during the assembly process of the In(NO₃)₃/S-pempH₂ system and hence the formation of spiral microflowers of S-1MF.

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Water stability of S-1C and S-1MF

Crystalline metal phosphonates and the corresponding nanosheets, especially those with high-valent metal ions, often exhibit high water stability. 41-43 To evaluate the water stability of S-1C and S-1MF, we conducted comprehensive stability assessments. As illustrated in Fig. 5, both compounds exhibited remarkable stability when exposed to boiling water for 24 h, indicating their resistance to hydrolysis and structural breakdown under high temperatures. The two samples were also immersed in aqueous solution at different pH values (adjusted using 0.01 M HCl or NaOH) for 24 h. Clearly, they maintained their structures across a wide pH range from 1 to 11 in aqueous solutions, which demonstrates their resilience in both acidic and basic environments (Fig. S29 and S30†). This stability is essential for practical applications, indicating that S-1MF can withstand harsh conditions, making it a promising candidate for applications in chiral recognition and others.

Chiral recognition properties

Chiral recognition on solid surfaces is important for related applications such as enantiomeric separation and sensing of chiral molecules and heterogeneous asymmetric catalysis. 44,45 The microflowers of S-1MF and R-1MF possess much larger surface areas than their crystalline counterparts, as well as domain-limited spaces between the petals. This is confirmed by the N₂ adsorption isotherm measurements at 77 K. As shown in Fig. S31,† at $P/P_0 \approx 0.98$, the adsorption capacities of **R-1MF** and **S-1MF** for N_2 were 32.72 cm³ g⁻¹ and 34.51 cm³ g⁻¹, respectively, which were higher than that of S-1C (16.01 cm 3 g $^{-1}$). In addition, the BET surface areas of **R-1MF** $(12.1 \text{ m}^2 \text{ g}^{-1})$ and **S-1MF** $(12.37 \text{ m}^2 \text{ g}^{-1})$ were also higher than that of S-1C (7.31 $\text{m}^2 \text{ g}^{-1}$). Notably, all three samples show rather low BET areas, which is reasonable considering that R-1 and S-1 have a densely packed layered structure. The pore size distribution determined by the BJH method⁴⁶ showed three

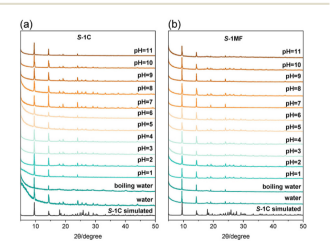


Fig. 5 PXRD patterns of the as-synthesized samples S-1C (a) and S-1MF (b) by soaking in water, boiling water and aqueous solutions in the pH range from 1 to 11 for 24 h.

peaks at 1.93, 7.98 and 14.02 nm for R-1MF and S-1MF, indicating the presence of both micro- and meso-pores in these materials. In contrast, the pore size of S-1C is concentrated around 1.29 nm, suggesting the presence of only micropores (Fig. S32†).

To investigate the chiral recognition properties of these materials, adsorption measurements were performed on activated samples of S-1MF, R-1MF, and S-1C using enantiopure R-2-butanol and S-2-butanol as probe molecules. The adsorption isotherms clearly exhibit enantioselective behaviour. At $P/P_0 \approx$ 0.98, the adsorption capacity for S-2-butanol on S-1MF (4.57 cm³ g⁻¹) is significantly higher compared to **R-1MF** $(3.06 \text{ cm}^3 \text{ g}^{-1})$ and **S-1C** $(3.45 \text{ cm}^3 \text{ g}^{-1})$ (Fig. 6a). The higher adsorption capacity of S-1MF in comparison with S-1C can be explained by the larger surface area of the microflowers than the crystalline sample. Fig. 6b and Fig. S33† demonstrate the differential adsorption of R-2-butanol and S-2-butanol on S-1MF and R-1MF. Apparently, S-1MF shows much higher adsorption for S-2-butanol (4.57 cm 3 g $^{-1}$) than for R-2-butanol $(0.86 \text{ cm}^3 \text{ g}^{-1})$, indicating a clear preference for the S-enantiomer. We note that the adsorption of S-2-butanol on **S-1MF** becomes obvious above $P/P_0 = 0.4$, suggesting a gateopening effect. Moreover, the adsorption/desorption isotherm shows a large hysteresis, with the desorption of S-2-butanol occurring below $P/P_0 = 0.25$. The result suggests that a strong interaction is present between the adsorbent S-1MF and S-2butanol.

According to the structural description, compound S-1 has a positively charged layer structure, balanced by the NO₃⁻ anion. The latter forms hydrogen bonds with the coordinated OH and water molecules as well as the uncoordinated amino groups of S-pempH⁻ (Fig. 7). As a result, the adjacent 1-phenylethylamino groups of S-pempH⁻ form chiral pockets on the surface of the layer together with the nitrate anion. The chiral guest molecule, S-2-butanol, can be inserted into the pocket and interacts with the layer through O-H...O hydrogenbonding interaction with the nitrate anion and/or C-H $\cdots\pi$ interaction with the phenyl groups in the layer. Obviously, the small chiral pocket in S-1MF favours the adsorption of S-2butanol over R-2-butanol. This may be the reason why the

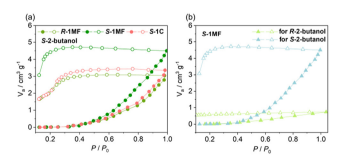


Fig. 6 (a) Comparison of the S-2-butanol adsorption (filled)/desorption (open) isotherms for the activated compounds R-1MF, S-1MF, and S-1C. (b) S- and R-2-butanol adsorption (filled) and desorption (open) isotherms for the activated compound S-1MF at 298 K.

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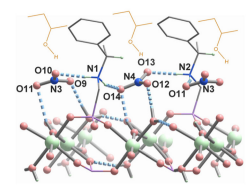


Fig. 7 The proposed adsorption mechanism of S-2-butanol (in brown color) on the layer surface of S-1MF.

adsorption capacity of S-1MF for S-2-butanol is significantly enhanced. PXRD analysis confirmed that the structures of S-1C, S-1MF, and R-1MF were preserved following the adsorption tests (Fig. S34†).

Conclusions

We report the first examples of chiral indium phosphonates, namely R-, $S-[In_2(pempH)_2(\mu_2-OH)_2(H_2O)_2](NO_3)_2$ (R-, S-1), which exhibit layered structures. The morphology of these compounds can be modulated using alcohol/water mixed solvents, leading to crystals of R-, S-1C in TEG/H2O and spiral microflowers of R-, S-1MF in IPA/H₂O. The acquisition of R-, S-1MF spiral microflowers demonstrates that chirality has been successfully transferred and amplified from the chiral ligand to the layer and finally to the morphology. Interestingly, the spiral microflowers of R-, S-1MF are stable in water. Adsorption studies using R- and S-2-butanol as probing molecules revealed that S-1MF shows enhanced adsorption capacity in comparison with its crystalline counterpart and pronounced adsorption discrimination properties. This work may encourage further exploration of chiral metal phosphonates with chiral morphologies macroscopically and interesting functions.

Author contributions

Q. T. and R. G. conducted experiments and analysed the data. Q. T. wrote the manuscript. S.-S. B. solved the crystal structure. L.-M. Z. conceived the project and revised the manuscript.

Data availability

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

Conflicts of interest

There are no conflicts to declare.

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References

- 1 X. Niu, R. Zhao, S. Yan, Z. Pang, H. Li, X. Yang and K. Wang, Small, 2023, 19, 2303059.
- 2 M. Xue, B. Li, S.-L. Qiu and B.-L. Chen, Mater. Today, 2016, 19, 503-515.
- 3 G. Sun, X. Zhang, Z. Zheng, Z.-Y. Zhang, M. Dong, J.-L. Sessler and C.-J. Li, J. Am. Chem. Soc., 2024, 146, 26233-26242.
- 4 X. Shang, I. Song, H. Ohtsu, Y.-H. Lee, T. Zhao, T. Kojima, J.-H. Jung, M. Kawano and J. H. Oh, Adv. Mater., 2017, 29, 1605828.
- 5 T. He, I. K. W. On, S. Bi, Z. Huang, J. Guo, Z. Wang and Y. Zhao, Angew. Chem., Int. Ed., 2024, 63, e202405769.
- 6 (a) S. Das, S. Xu, T. Ben and S.-L. Qiu, Angew. Chem., Int. Ed., 2018, 57, 8629; (b) J.-P. Zhao, J. Luo, Z.-H. Lin, X. Chen, G.-H. Ning, J.-Z. Liu and D. Li, Inorg. Chem. Front., 2022, 9, 4907-4912.
- 7 (a) X.-H. Niu, Y.-Q. Liu, R. Zhao, M. Yuan, H.-F. Zhao, H.-X. Li and K.-J. Wang, ACS Appl. Mater. Interfaces, 2024, 16, 17361-17370; (b) C. Li, X.-X. Yang, M.-Y. Zheng, Z.-G. Gu and J. Zhang, Adv. Funct. Mater., 2024, 34, 2401102.
- 8 (a) S. Zhang, Y. Zheng, H. An, B. Aguila, C.-X. Yang, Y. Dong, W. Xie, P. Cheng, Z. Zhang, Y. Chen and S. Ma, Angew. Chem., Int. Ed., 2018, 57, 16754; (b) A. Gheorghe, B. Strudwick, D. M. Dawson, S. E. Ashbrook, S. Woutersen, D. Dubbeldam and S. Tanase, Chem. - Eur. J., 2020, 26, 13957.
- 9 C.-X. Tan, K.-W. Yang, J.-Q. Dong, Y.-H. Liu, Y. Liu, J.-W. Jiang and Y. Cui, J. Am. Chem. Soc., 2019, 141, 17685-17695.
- 10 J. Guo, Y. Zhang, Y. Zhu, C. Long, M. Zhao, M. He, X. Zhang, J. Lv, B. Han and Z.-Y. Tang, Angew. Chem., Int. Ed., 2018, 57, 6873.
- 11 B. Shen, Y. Kim and M. Lee, Adv. Mater., 2020, 32, 1905669.
- 12 Y.-H. Liu, L.-M. Liu, X. Chen, Y. Liu, Y. Han and Y. Cui, J. Am. Chem. Soc., 2021, 143, 3509-3518.
- 13 H.-J. Jiang, L. Zhang, J. Chen and M.-H. Liu, ACS Nano, 2017, 11, 12453-12460.
- 14 Y. Jin, C. Xiao, L. Tan, Z. Chen, Z. Wen, Y. Cheng, W. Fu and P. Wang, Adv. Opt. Mater., 2023, 11, 2203068.
- 15 (a) K. Ding, J. Ai, Q. Deng, B. Huang, C. Zhou, T. Duan, Y. Duan, L. Han, J. Jiang and S. Che, Angew. Chem., Int. Ed., 2021, **60**, 19024; (b) Y.-Y. Duan, X. Liu, L. Han, S. Asahina,

Paper **Dalton Transactions**

- D.-D. Xu, Y.-Y. Cao, Y. Yao and S. Che, J. Am. Chem. Soc., 2014, 136, 7193-7196.
- 16 B. Zhao, H.-L. Yu, K. Pan, Z. Tan and J.-P. Deng, ACS Nano, 2020, 14, 3208-3218.
- 17 Metal Phosphonate Chemistry: From Synthesis to Applications, ed. A. Clearfield and K. Demadis, The Royal Society of Chemistry, 2012.
- 18 S. J. Shearan, N. Stock, F. Emmerling, J. Demel, P. A. Wright, K. D. Demadis, M. Vassaki, F. Costantino, R. Vivani, S. Sallard, I. R. Salcedo, A. Cabeza and M. Taddei, Crystals, 2019, 9, 270.
- 19 (a) S.-S. Bao, M.-F. Oin and L.-M. Zheng, Chem. Commun., 2020, 56, 12090-12108; (b) T. Zheng, W. Tan and L.-M. Zheng, Acc. Chem. Res., 2024, 57, 2973-2984.
- 20 G.-G. Weng and L.-M. Zheng, Sci. China: Chem., 2020, 63, 619-636.
- 21 X.-G. Liu, S.-S. Bao, Y.-Z. Li and L.-M. Zheng, *Inorg. Chem.*, 2008, 47, 5525-5527.
- 22 Z.-M. Zhai, T. Hou, Y. Xu, O. Teng, S.-S. Bao and L.-M. Zheng, Chem. - Eur. J., 2025, 31, e202403699.
- 23 SAINT, version 8.40A, Program for Data Extraction and Reduction, Bruker Nano., Inc., 2019.
- 24 G. M. Sheldrick, SHELXT, 2014/5, 2014; SHELXL 2018/3,
- 25 Y. Inokuchi and N. Nishi, J. Chem. Phys., 2001, 114, 7059-7065
- 26 J. Morizzi, M. Hobday and C. Rix, J. Mater. Chem., 1999, 9, 863-864.
- 27 A. F. Richards and C. M. Beavers, Dalton Trans., 2012, 41, 11305-11310.
- 28 W.-Y. Dan, X.-F. Liu, M.-L. Deng, Y. Lin, Z.-X. Chen and Y.-M. Zhou, Dalton Trans., 2015, 44, 3794-3800.
- 29 V. Chandrasekhar, J. Goura and A. Duthie, Inorg. Chem., 2013, 52, 4819-4824.
- 30 Y. Y. Enakieva, M. V. Volostnykh, S. E. Nefedov, G. A. Kirakosyan, Y. G. Gorbunova, A. Y. Tsivadze, A. G. Bessmertnykh-Lemeune, C. Stern and R. Guilard, Inorg. Chem., 2017, 56, 3055-3070.
- 31 A. Kechiche, S. A. Shehimy, L. Khrouz, C. Monnereau, Bucher, S. Parola, A. Bessmertnykh-Lemeune,

- Y. Rousselin, A. V. Cheprakov and H. Nasri, Dalton Trans., 2024, 53, 7498-7516.
- 32 J.-G. Jia, C.-C. Zhao, Y.-F. Wei, Z.-M. Zhai, S.-S. Bao, A. J. Jacobson, J. Ma and L.-M. Zheng, J. Am. Chem. Soc., 2023, 145, 23948-23962.
- 33 G.-G. Weng, X.-D. Huang, R. Hu, S.-S. Bao, O. Zou, G.-H. Wen, Y.-Q. Zhang and L.-M. Zheng, Chem. - Asian J., 2021, 16, 2648-2658.
- 34 T. Hou, C.-C. Zhao, S.-S. Bao, Z.-M. Zhai and L.-M. Zheng, Dalton Trans., 2024, 53, 4291-4298.
- 35 S.-Q. Zhang, L.-N. Li, S.-G. Zhao, Z.-H. Sun, M.-C. Hong and J.-H. Luo, J. Mater. Chem. A, 2015, 3, 15764-15768.
- 36 M. Rashid, S. Parsaei, A. Ghoorchian, K. Dashtian and D. Mowla, J. Ind. Eng. Chem., 2023, 121, 275-286.
- 37 S.-L. Zhong, Z.-G. Zou, S.-J. Lv, S.-C. Zhang, J. Geng, J.-Y. Meng, X. Liu, F.-G. Liang and J.-J. Rao, J. Alloys Compd., 2022, 903, 163845.
- 38 N. Roy, H. R. Barai, A. N. Banerjee, J. S. Kim and S. W. Joo, Chem. Eng. Sci., 2023, 277, 118834.
- 39 J. Wang, J. Yang, H. Yang, H.-Z. Huang, X.-D. Yang and L. Wei, Opt. Mater., 2023, 135, 113303.
- 40 Z.-S. Cai, Y. Shi, S.-S. Bao, Y. Shen, X.-H. Xia and L.-M. Zheng, ACS Catal., 2018, 8, 3895-3902.
- 41 Y. Huang, F. Zhou, J. Feng, H. Zhao, C. Qi, J. Ji, S. Bao and T. Zheng, Chem. Commun., 2021, 57, 1238-1241.
- 42 G.-H. Wen, X.-M. Chen, K. Xu, X. Xie, S.-S. Bao and L.-M. Zheng, Dalton Trans., 2021, 50, 17129-17139.
- 43 (a) M.-F. Qin, Q.-Q. Mu, S.-S. Bao, X. Liang, Y. Peng and L.-M. Zheng, ACS Appl. Energy Mater., 2021, 4, 4319-4326; (b) M.-F. Qin, L.-F. Zheng, S.-S. Bao, F.-H. Xue, J. Su, J. Ma and L.-M. Zheng, ACS Appl. Electron. Mater., 2023, 5, 887-
- 44 J. Sui, N. Wang, J. Wang, X. Huang, T. Wang, L. Zhou and H. Hao, Chem. Sci., 2023, 14, 11955-12003.
- 45 G. Li, Q. Yang, K. Manna, Y. Zhang, P. Merz, C. Shekhar, Y. Zhang, H. Lv, A. Markou, Y. Sun and C. Felser, Angew. Chem., Int. Ed., 2023, 62, e202303296.
- 46 M. Thommes, K. Kaneko, A. V. Neimark, J. P. Olivier, F. Rodriguez-Reinoso, J. Rouquerol and K. S. Sing, Pure Appl. Chem., 2015, 87, 1051-1069.