Heteroleptic Triple-Stranded Metallosupramolecules with Hydrophobic Inner Voids

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ABSTRACT: The systematic combination of well-defined coordination spheres and multiple types of ligands (heteroleptic) can lead to the generation of hierarchical metallosupramolecules with a high level of complexity and functionality. In particular, a specific multilevel coordination-driven assembly through the initiate generation of multinuclear clusters can form unique heteroleptic multiple-stranded supramolecular complexes. Herein, we report novel triple-stranded nickel-based supramolecules constructed from two different ditopic ligands ([1,1′:3′,1′″-terphenyl]-4,4″-dicarboxylate (TP) and 2,6-pyridinedicarboxylate (PDA)) and a nickel precursor. The solid-state structures of the as-synthesized supramolecules revealed that three PDA ligands are employed to fabricate a tetranuclear ({{Ni}}₄) cluster, and two {{Ni}}₃ clusters are assembled to form the final triple-stranded metallosupramolecules by three TP ligands. The bridging TP ligands also provide large inner voids with highly hydrophobic environments. Structural investigation of the generated complexes provided a deeper understanding of the aspects driving the formation of heteroleptic supramolecules, which is crucial for the design of multiple-strands with desired morphologies and functionalities.

INTRODUCTION

Highly programmed metallosupramolecular assemblies have attracted intensive interest in mimicking the unique topologies of various natural structures. To obtain a high level of intricacy and functionality, systematic combinations of metal ions with well-defined coordination and multiple types of ligands (heteroleptic) with proper geometric properties have been attempted, and the resulting hierarchical structures have been applied in the construction of highly ordered systems, energy conversion, catalysis, and enzyme mimics. Each ligand in a heteroleptic supramolecule can have its own structural role in driving the assembly of primary metallic clusters or combining the clusters to generate secondary assemblies within a unique structural organization. Efforts have been made to create heteroleptic metallosupramolecules. However, entropy often thwarts these efforts due to the high tendency to form uncontrolled statistical mixtures of homo- and heteroleptic supramolecules or narcissistically self-sorted mixtures. Hence, the formation of a single, integratively self-sorted species needs to be driven by significant enthalpic benefit originated from the perfect combination of two types of ligand components in the assembly to overcome entropic penalty, or the homoleptic structures are disfavored by repulsive steric or strain effects.

Among various reported discrete supramolecular species up to now, helicates and mesocates are considered to be the simplest architectures. The chirality of these two structural modes depends on the assembly of two chiral metal units. The combination of two units with the same configuration results in a chiral helicate. If two units of opposite configurations are combined, an achiral mesocate is formed. The formation of helicate or mesocate greatly depends on the nature of the metal and ligands, and sometimes on additional information introduced by template or reaction conditions. For ligands, the stereoselectivity of the self-assembly can be influenced by ligand rigidity, ligand geometry, and the number of methylene units in the alkyl spacers. Immense works have been done for the investigation of helicate and mesocate compounds, but the future will show the merit of studying this fascinating topic of metallosupramolecular chemistry.

In our reported works, we have reported a range of supramolecules in which metal ions (Co²⁺, Ni²⁺, or Mn²⁺) coordinated with 2,6-pyridinedicarboxylate (PDA) to form tetranuclear metal clusters (primary assembly), and two metal clusters are interlinked by three bridging ligands, that is, benzene-1,3-dicarboxylate (PTA) derivatives or 4,4′-carbonyldibenzoate (CDBA), to form triple-stranded helicates (TSH) with C₃ symmetry. For supramolecules constructed from...
PTA derivative, three bridging ligands with a bending angle of 120° create an empty space within the supramolecule. Depending on the substituent-functional group attached to C5 of the PTA ligand, various higher-order assembly modes were obtained such as discrete, polymeric, and cage metallosupramolecules.31–35 When moving from a one-benzene ring system (PTA) to a two-benzene ring system (CDBA), TSHs could also be obtained with C3 symmetry.36 The inner voids surrounded by three CDBA ligands are larger than those surrounded by three PTA derivative ligands. Meanwhile, the higher length and flexibility of CDBA initiate a higher potential for the generation of unprecedented higher-order assembly modes with high complexity, flexibility, and diversity. The maintenance of heteroletic structures when changing metal ions, as well as extending the ligand length and flexibility, also suggests that the symmetry and helicity of triple-stranded metallosupramolecules are energetically favorable.37

On the basis of the results about metallosupramolecules we have obtained so far, we expect that the extension in bridging ligand length, together with the variation of the substituent-functional groups in the backbone, could create metallosupramolecules with unprecedented assembly modes. The lengthening of the bridging ligand is also expected to generate voids with large size, which is meaningful in host-guest interaction. Herein, 5′-methyl-[1′,3′,1′′-terphenyl]-4,4′-dicarboxylic acid (CH3-H2TP) and 5′-hydroxy-[1′,3′,1′′-terphenyl]-4,4′-dicarboxylic acid (HO-H2TP) were prepared according to the method reported in the literature.38–40 Other chemicals were purchased from commercial sources and used without further purification. Tetrakis(triphenylphosphine)palladium(0) (Pd(PPh3)4, 95%, Fluorochem), 3,5-dibromotoluene (C7H6Br2, 98%, TCI), 3,5-dibromophenol (C6H4Br2O, 98%, TCI), 4-carboxyl-C7H5NO4, 99%, Sigma-Aldrich), N,N-dimethylformamide (DMF, 99.99%, Burdick and Jackson), hydrochloric acid (68%), nitric acid (70%, Sigma-Aldrich), nickel(II) acetate tetrahydrate (Ni(OAc)2·4H2O, 99%, Sigma-Aldrich), nickel(II) nitrate hexahydrate (Ni(NO3)2·6H2O, 99.9%, Samchun), potassium carbonate (K2CO3, 99.0%, TCI), methanol (99.5%, Sigma-Aldrich), ethanol (99.9%, Sigma-Aldrich), and acetone (99.96%, Burdick and Jackson) were used as received.

Syntheses. Synthesis of CH3-H2TP (5′-Methyl-[1′,3′,1′′-terphenyl]-4,4′-dicarboxylic acid) and HO-H2TP (5′-Hydroxy-[1′,3′,1′′-terphenyl]-4,4′-dicarboxylic acid). The synthesis of CH3-H2TP ligand was according to a method proposed in the literature with a slight modification.38–40 In a typical synthesis, 3,5-dibromotoluene (2.00 g, 8.00 mmol) and 4-carboxyphenylboronic acid (3.98 g, 24.0 mmol) were dissolved in potassium carbonate solution (16.0 g in 93 mL H2O) and 180 mL of dimethylformamide (DMF) in a 500 mL Schlenk flask. The resulting mixture was stirred vigorously and degassed for 30 min. After that, tetrakis(triphenylphosphine)-palladium(0) (1.27 g, 1.10 mmol) was added quickly to the flask, and the system was degassed for a further 10 min. The Schlenk flask was sealed and stirred at 100 °C for 3 days. The reaction mixture after cooling down to room temperature (RT) was poured into 100 mL of H2O and washed with dichloromethane (5 × 100 mL). The water phase was separated and acidified with an aqueous HCl solution (v/v = 1:1). The product was collected by centrifugation at 4500 rpm for 3 min, washed with water four times, and finally washed with methanol. The resulting white solid was dried in an oven at 90 °C for 24 h to give CH3-H2TP ligand in 55.3% yield (based on the amount of 3,5-dibromotoluene used). The 1H NMR spectrum of the CH3-H2TP ligand is shown in Figure S1.

HO-H2TP ligand was synthesized using a similar protocol, with the exception that 3,5-dibromophenol was used instead of 3,5-dibromotoluene. The product yield was 58.6% based on the amount of 3,5-dibromophenol used. Figure S2 shows the 1H NMR spectrum of the HO-H2TP ligand.

Synthesis of [Ni8(PDA)6(CH3-TP)3(H2O)6](DMF)2.1H2O.38 DMF solutions of Ni(NO3)2·6H2O (1.2 mL, 0.05 M), H2PDA (0.6 mL, 0.05 M), CH3-H2TP (0.3 mL, 0.05 M), and HNO3 (0.1 mL, 0.05 M) were mixed in a 4 mL glass vial at RT. The vial was sealed, heated to 100 °C (heating rate: 2.67 °C/min), maintained for 24 h, and then cooled to 30 °C (cooling rate: ~0.25 °C/min). After the cooling process, green hexagonal plate-shaped crystals of I, suitable for SC-XRD analysis, were successfully formed and sequentially washed with DMF (3 × 10 mL) and acetone (3 × 10 mL) and then dried in vacuo. The solid yield of I was 37.3% based on H2PDA. Anal. Calc. for C29H23N2O12Ni3 (FW = 5259.36 g/mol): C: 49.27, H: 2.84, N: 3.28. Found: C: 49.85, H: 4.28, N: 5.58.

Synthesis of [Ni8(PDA)6(HO-TP)3(DMF)6(H2O)3] (2). The synthesis of 2 was similar to that of I, except for the use of different precursors. Specifically, a mixture of Ni(OAc)2·4H2O (1.2 mL, 0.05 M), H2PDA (0.6 mL, 0.05 M), HO-H2TP (0.3 mL, 0.05 M), and HCl (0.1 mL, 1.00 M) was used. Compound 2 was obtained as green rod-shaped crystals with a solid yield of 40.5% based on H2PDA. Anal. Calc. for C31H23N2O12Ni3 (FW = 2730.41 g/mol): C: 48.78, H: 2.97, N: 4.61. Found: C: 45.33, H: 4.10, N: 7.01.

Instrumentation. The 1H nuclear magnetic resonance (1H NMR) spectra of the organic ligands were recorded on a Bruker AVANCE III-400 instrument (400 MHz) using dimethyl sulfoxide-d6 (DMSO-d6) as the NMR solvent. Thermogravimetric analysis (TGA) and differential scanning calorimetry (DSC) were performed from RT to 600 °C at a heating rate of 10 °C/min under a nitrogen atmosphere using a Q600 simultaneous DSC-DTA-TGA system from TA Instruments. Powder X-ray diffraction (PXRD) was performed using synchrotron radiation (wavelength 1.1 Å) in focused beam configuration in the 2θ range of 2°–30° at 298 K. The simulated PXRD patterns were calculated from the single-crystal X-ray diffraction (SC-XRD) data using the Mercury 3.8 program. X-ray photoelectron spectroscopy (XPS) measurement was performed on an R3000 spectrometer (VG SCIENTA, United Kingdom) with monochromated Al Kα X-ray radiation as the X-ray source for excitation. Magnetic measurements of compounds were performed using a
Quantum Design MPMS3 magnetometer for temperatures $3 \leq T \leq 300$ K with a 1000 Oe applied field.

**X-ray Crystallography.** Diffraction data obtained from the light green crystals of (1) $(0.117 \times 0.094 \times 0.084 \text{ mm}^3)$ and (2) $(0.105 \times 0.058 \times 0.027 \text{ mm}^3)$ mounted on a MiTeGen MicroMount were collected at 223 K (for 1) and 293 K (for 2) on a Rayonix MX225HS detector with a silicon (111) double-crystal monochromator (DCM) equipped with a synchrotron radiation source $(0.80000$ and $0.70000 \text{ Å for 1 and 2},$ respectively) at the 2D Supramolecular Crystallography Beamline (2D SMC), Pohang Accelerator Laboratory (PAL), Pohang, Republic of Korea. All of the calculations for the structure determination were carried out using the SHELXTL package (Ver. 2018/3).

**RESULTS AND DISCUSSION**

CH$_3$-H$_2$TP (5′-methyl-[1,1′:3′,1″′-terphenyl]-4,4″′-dicarboxylic acid) and HO-H$_2$TP (5′-hydroxy-[1,1′:3′,1″′-terphenyl]-4,4″′-dicarboxylic acid) were synthesized through a Suzuki coupling reaction according to the method reported in the literature with a slight modification.$^{38−40}$ Similar to H$_2$PTA (benzene-1,3-dicarboxylic acid) and H$_2$CDBA (4,4″′-carbonyldibenzoic acid) used in the syntheses of the previously reported triple-stranded supramolecules,$^{31−36}$ both CH$_3$-H$_2$TP and HO-H$_2$TP are ditopic ligands with a bending angle of $\sim 120°$ (Scheme 1). The fundamental difference between substituted H$_2$TP ligands and H$_2$PTA or H$_2$CDBA lies in their length and flexibility, that is, the distance between the carbon atoms in the two carboxyl groups of H$_2$TP ligand is significantly longer than those of H$_2$PTA and H$_2$CDBA (Scheme 1). In addition, H$_2$TP ligands show a higher level of flexibility when compared to H$_2$PTA or H$_2$CDBA, which can be mainly attributed to the rotations around the two single C–C bonds connected to the central phenyl ring.

Treatment of 4 equiv of Ni(NO$_3$)$_2$·6H$_2$O with 2 equiv of H$_2$PDA and 1 equiv of CH$_3$-H$_2$TP in DMF with HNO$_3$ at 100 °C led to the formation of [Ni$_8$(PDA)$_6$(CH$_3$-TP)$_3$(H$_2$O)$_6$] (1), which was isolated as a green crystalline product (Figure S3a and Scheme 2). The structure of 1 in the solid state was analyzed using single-crystal X-ray crystallography (SC-XRD)

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**Scheme 1. Chemical Structures of Ligands That Have Been Used to Synthesize Heteroleptic Triple-Stranded Supramolecules:** H$_2$PDA, H$_2$PTA Derivatives (X-H$_2$PTA, X = −H, −tBu, −OH, −Br, −I, −NH$_2$), H$_2$CDBA (Previous Works), and H$_2$TP Derivatives (X-H$_2$TP, X = −CH$_3$, −OH) (This Work)

**Scheme 2. Schematic Representation for the Formation of 1 and 2**

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**Figure 1.** X-ray crystal structure of [Ni$_8$(PDA)$_6$(CH$_3$-TP)$_3$(H$_2$O)$_6$] (1). (a) Side view of 1 with two Ni$_4$ clusters linked by three CH$_3$-TP ligands, together with a mirror plane of symmetry in 1 (green balls show cobalt, purple parts show nitrogen, and red parts show oxygen). All the coordinated and free solvent molecules, hydrogen atoms, and disorder components were omitted for clarity. (b) Presentation of two nickel clusters in form of clockwise and counterclockwise isomers.
in every experiment. The metal clusters possess chirality, either clockwise or counterclockwise, which is similar to the reported structures.\textsuperscript{31–36} Each discrete supramolecule of 1 is constructed from two metal clusters with opposite chirality, resulting in a meso-helicate (Figure 1). The existence of one mirror plane of symmetry passing through the middle point of the three CH$_3$-TP ligands initiates C$_{3v}$ symmetry of the meso-helicate. This is unique when compared to the reported triple-stranded supramolecules,\textsuperscript{31–36} which show two metal clusters of the same chirality interconnected via three PTA derivative or CDBA bridging ligands, leading to the formation of either right-handed or left-handed helicate with C$_3$ symmetry. In terms of crystal packing, the discrete molecules of 1 are arranged regularly into layers. Every two adjacent layers are staggered and separated by a gap of 3.5 Å (Figure 2a,b). Every alternating layer is eclipsed, resulting in the formation of honeycomb-shaped one-dimensional channels (Figure 2c).

The phase purity of the as-synthesized 1 was determined using powder X-ray diffraction (PXRD) (Figure S5). There are considerable similarities between the observed and the expected PXRD patterns. Thermogravimetric analysis (TGA) and differential scanning calorimetry (DSC) of 1 were shown in Figure S6. The first weight loss of 6.1% up to 168 °C is assigned to the loss of coordinated solvent within 1 and the next weight loss of 56.1% up to 600 °C corresponds to the decomposition of organic ligands.\textsuperscript{34} The endothermic peaks and weight loss observed from DSC and TGA curves at 350–420 °C indicate that the thermal decomposition of 1 occurred rapidly at this temperature range. XPS full spectrum confirmed the presence of Ni, O, C, and N in 1 (Figure S7a). The deconvolution of high-resolution O 1s XPS spectrum indicates the present C–O (530.8 eV), C–O–Ni (532.0 eV), and C=C (533.0 eV).\textsuperscript{41} While the deconvolution of high-resolution N 1s XPS spectrum suggests the existence of N (pyridine–amide) (398.7 eV) and N (amide) (399.9 eV) in 1 (Figure S7bc).\textsuperscript{42–43} The oxidation state of nickel ions was determined to be +2 by the presence of two strong peaks at 855.98 and 873.11 eV standing for Ni 2p$_{3/2}$ and Ni 2p$_{1/2}$, respectively (Figure S7d).\textsuperscript{35,36} The bond valence sum calculation also revealed that nickel ions in these clusters have a +2 oxidation state (Table S2).\textsuperscript{44,45} The measurement of temperature-dependent magnetization (emu/g) of 1 was conducted using a Quantum Design MPMS3 magnetometer for temperatures 3 K ≤ T ≤ 300 K with 1000 Oe applied fields (Figure S8a). By applying the Curie–Weiss law, the corresponding fitting (1/χ vs T) (Figure S8b) yielded a Weiss constant $\theta = -7.78$ K, suggesting the week antiferromagnetic interaction between nickel ions.\textsuperscript{46} and Curie–Weiss constant $C = 0.00383$ emu K g$^{-1}$. The measured $\chi_M T$ at 300 K was 1.20 (emu K mol$^{-1}$) (Figure S8c), and the corresponding $\mu_B$ was calculated to be 3.10 μB. This value falls within an acceptable range of experimentally observed high-spin octahedral Ni (II) ions.\textsuperscript{47}

The inner empty space of 1, as illustrated in Figure S9, is noticeable. The size of this “inner cage” is significantly larger than the similar empty spaces of the previously reported triple-stranded supramolecules prepared using PTA or CDBA ligands.\textsuperscript{31–36} The larger size of the inner space of 1 possibly allows the inclusion of external molecules through further ligand modification. In general, the hydrophobicity of a cage can be proved by the inclusion of hydrophobic molecules from the water phase.\textsuperscript{48} It was also commented that the central cavity of a cage may be hydrophobic owing to its surrounding aryl rings.\textsuperscript{49–50} Therefore, the empty space inside 1, which is

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**Figure 2.** Packing structures of 1. Space-filling representations of 1 from the views of the crystallographic (a) a-axis and (b) c-axis, showing the staggered assembly of adjacent packing layers of 1. (c) Capped sticks representation of 1 illustrating its honeycomb packing structure.

**Figure 3.** X-ray crystal structure of [Ni$_8$(PDA)$_6$(HO-TP)$_3$(DMF)$_3$H$_2$O)$_3$] (2). (a) Side view and top-down view of 2. (b) Representation of the left-handed and right-handed configurations of 2 with the respective constructing clusters.
surrounded by aryl rings, can be regarded as hydrophobic. Moreover, using SQUEEZE,\textsuperscript{51} we could confirm that solvent molecules were excluded from the void. Hence, we speculate that the inner voids are highly hydrophobic. The numerous C–H bonds in the aromatic rings are oriented toward the inner space, providing the possibility for host–guest interactions including hydrophobic and CH⋯π interactions.\textsuperscript{52,53} Thus, these inner empty spaces may exhibit selectivity toward neutral and hydrophobic guests because of the hydrophobic effect.\textsuperscript{48,54} In addition, the hydrophobic inner voids can be further applied as a reactor due to their ability to provide a suitable environment for hosting chemical reactions, as well as stabilizing reactive hydrophobic intermediates.\textsuperscript{48,55}

The effects of bridging ligand modification on the self-assembly of heteroleptic supramolecules were also investigated with HO-TP ligand. Four equiv of Ni(OAc)\textsubscript{2}·4H\textsubscript{2}O, 2 equiv of H\textsubscript{2}PDA, and 1 equiv of HO-H\textsubscript{2}TP were treated in DMF with HNO\textsubscript{3} at 100 °C, forming [Ni\textsubscript{8}(PDA)\textsubscript{6}(HO-TP)\textsubscript{3}(DMF)\textsubscript{3}(H\textsubscript{2}O)\textsubscript{3}] (2) (Scheme 2 and Figure S3b). The solid-state structure of 2 shows multinuclear clusters of either clockwise or counterclockwise isomer (Figures 3 and S10). Different from 1, two \{Ni\textsubscript{4}\} clusters are connected by three bridging ligands with unequal twisting, leading to the symmetry breaking in the structure (Figure 3a). The combination of two nickel clusters of the same chirality in a single supramolecule results in a helicate of either left-handed or right-handed configuration (Figure 3b). PXRD spectra of 2 show a good match between the experimental and the simulated patterns (Figure S11). The thermal property of 2 was shown in Figure S12. The first weight loss of 6.1% up to 168 °C is assigned to the removal of coordinated solvent, and the next weight loss of 56.1% up to 450 °C corresponds to the decomposition of organic ligands.\textsuperscript{31} The fast decomposition at the temperature range of 360–430 °C was indicated not only by the dramatic weight loss but also the thermal event observed on the DSC curve at this condition. XPS full spectra confirmed the presence of Ni, O, C, and N in 2 (Figure S13a). The deconvolution of high-resolution O 1s XPS spectrum indicates the presence of O in the form of C–O, C=O–Ni, and C=O,\textsuperscript{41} while the deconvolution of the high-resolution N 1s XPS spectrum suggests the existence of pyridine nitrogen and amide nitrogen in 2 (Figure S13b,c).\textsuperscript{42,43} The +2 oxidation state of the nickel ions was confirmed with two strong peaks at 856.69 and 873.94 eV corresponding to Ni 2p\textsubscript{3/2} and Ni 2p\textsubscript{1/2}, respectively (Figure S13d).\textsuperscript{35,36} Using the bond valence sum theory, the oxidation state of the nickel ions in the clusters was...
determined to be +2 (Table S2).\textsuperscript{44,45} The temperature-dependent magnetization (emu/g) of 2 is illustrated in Figure S14a. The Curie–Weiss fitting (1/\chi vs T) (Figure S14b) yielded a Weiss constant \( \theta = -8.06 \) K, indicating the weak antiferromagnetic interaction between nickel ions\textsuperscript{66} and Curie–Weiss constant \( C = 0.00386 \) emu K g\(^{-1}\). The value of \( \chi_M \) at 300 K was 1.31 (emu K mol\(^{-1}\)), and the corresponding \( \mu_{eff} \) was 3.23 \( \mu_B \), which falls within an acceptable range of experimentally observed high-spin octahedral Ni (II) ions.

The crystal packing of 2 revealed that the supramolecules are stacked by intermolecular interactions composed of numerous O–H···O hydrogen bonds (Figure 4). The estimated hydrogen bond lengths and bond angles of these interactions were summarized in Table S3. Notably, the O···O distances in O2O–H2O···O3 and O15–H15···O32 hydrogen bonds are 2.658 and 2.648 Å, significantly less than the sum of van der Waal radii of two oxygen atoms (3.04 Å), indicating that these hydrogen bonds are strong.\textsuperscript{66–69} Considering the C104–H10A···O4 and C24–H24···O20 hydrogen bonds, although the C···O distances are 3.327 and 3.440 Å, which are longer than the sum of van der Waals radii of carbon and oxygen (3.22 Å), the H···O distances are 2.399 and 2.586 Å, less than the Van der Waals radii of hydrogen and oxygen (2.62 Å).\textsuperscript{77} In addition, their bond angles are 165.3 and 128.6\(^\circ\), which are in the range of a common hydrogen bond (110–180\(^\circ\)).\textsuperscript{60} These data indicate that the formation of two C–H···O bonds is relatively effective. Although the H···O distances in C57–H57···O15 and C110–H11A···O33 are 2.694 and 2.703 Å, respectively, which are higher than the sum of Van der Waals radii of hydrogen and oxygen, it does not mean that no interaction exists. Instead, it suggests that these two hydrogen bonds can contribute to the stabilization energy but to a lesser extent than the four aforementioned hydrogen bonds. We assume that the stabilization energy from the various O–H···O and C–H···O hydrogen bonds compensated for the loss of entropy and the increase in enthalpy owing to the symmetry breaking of 2.

**CONCLUSION**

In conclusion, two distinctive triple-stranded metallosupramolecules with unprecedented symmetries were successfully synthesized by extending the bridging ligand from a one- to a three-benzene-ring system with a longer length, higher flexibility, and variation in functional groups. While the assembly of tetrancular clusters is similar in 1 and 2, the different modes of combining two nickel clusters into a discrete supramolecule and the dissimilarity in linker geometries lead to the fabrication of meso-helicate 1 with very high symmetry (C\(_{3h}\)) and helicate 2 with no symmetry element. The combination of nickel clusters and TP derivative ligands generates a large empty space that can play the vital roles of a reactor or host–guest interaction.

**ASSOCIATED CONTENT**

**Supporting Information**

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsomega.2c00447.

Details of crystallographic data, additional figures of metallosupramolecules, characterization data of ligands (\(^{1}H\) NMR), characterization data of supramolecules (XPS, PXRD, TGA-DSC, and magnetic property), calculation of sizes of inner voids of supramolecules (PDF), Collection of two molecules (CIF)

**Accession Codes**

CCDC 2132311 and 2132320 contain the supplementary crystallographic data which can be obtained free of charge via www.ccdc.cam.ac.uk/data_request/cif, or by emailing data_request@ccdc.cam.ac.uk, or by contacting The Cambridge Crystallographic Data Centre, 12 Union Road, Cambridge CB2 1EZ, UK; fax: + 44 1223 336033.

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**Notes**

The authors declare no competing financial interest.

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**ABBREVIATIONS**

CH\(_{3}\)-TP, S’-methyl-[1,1′:3′,1′′-terphenyl]-4,4′-dicarboxylate; HO-TP, S’-hydroxy-[1,1′:3′,1′′-terphenyl]-4,4′-dicarboxylate; H\(_2\)-TP, [1,1′:3′,1′′-terphenyl]-4,4′-dicarboxylic acid; TP, [1,1′:3′,1′′-terphenyl]-4,4′-dicarboxylate; CDBA, 4,4′-carbonyldibenzoate

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