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Family of \([\text{Cu}_2]\), \([\text{Cu}_4]\) and \([\text{Cu}_5]\) Aggregates: Alteration of Reaction Conditions, Ancillary Bridges and Capping Anions

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Abstract. Phenoxido-bridged $[\text{Cu}_2]$ complex $[\text{Cu}_2(\mu-\text{H}_4\text{L}_1)(\mu-\text{OH})(\mu_{1,3-}\text{NO}_3)(\text{NO}_3)\text{(OH}_2)]\cdot\text{H}_2\text{O}$ (1), and its hierarchical $[\text{Cu}_4]$ and $[\text{Cu}_5]$ assemblies $[\text{Cu}_4(\mu-\text{H}_4\text{L}_1)_2(\mu-\text{OH})(\mu_{1,3-}\text{ClO}_4)\text{(OH}_2)_2]\text{(ClO}_4)_3\cdot2\text{H}_2\text{O}$ (2), $[\text{Cu}_5(\mu-\text{H}_4\text{L}_1)_2(\mu_3-\text{OH})(\mu_{1,3-}\text{O}_2\text{CCF}_3)_2]\text{(ClO}_4\text{COO)}_2\cdot3\text{H}_2\text{O}$ (3) were obtained from the reactions of $\text{H}_5\text{L}_1$ (2,6-bis-$(\{1,3\text{-dihydroxy-2-methylpropan-2-yl}	ext{imino})\text{methyl}\})\cdot4\text{-methylphenol}$) with three copper(II) salts. Available $\text{NO}_3^-$, $\text{ClO}_4^-$ and $\text{CF}_3\text{COO}^-$ ions have been trapped for ‘spontaneous’ anion directed ‘self-assembly’ reactions. All the synthesized complexes contain $[\text{Cu}_2(\mu-\text{H}_4\text{L}_1)(\mu-\text{OH})]^{2+}$ fragment, prone to assemble and crystallize $[\text{Cu}_4]$ and $[\text{Cu}_5]$ complexes in varying reaction conditions. They were characterized by UV–vis and IR spectroscopy, X-ray diffraction analysis and magnetic studies. A change from $\text{NO}_3^-$ to $\text{ClO}_4^-$ and $\text{CF}_3\text{COO}^-$ result different course of reaction based on $\text{Cu}_2(\mu-\text{H}_4\text{L}_1)$ fragments. Binding of $\text{NO}_3^-$ provided 1 as the isolated $[\text{Cu}_2]$ complex by trapping the reactive fragment. In 2 a perchlorate ligand, in $\mu_{1,3}$- binding mode, has been realized as a solitary support for the condensation of two $\text{Cu}_2(\mu-\text{H}_4\text{L}_1)$ fragments. The $(\{\text{Cu}_5(\mu_3-\text{OH})(\mu_{1,3-}\text{O}_2\text{CCF}_3)_2\}_{6+}$ constellation in 3 contains five $\text{Cu}^{II}$ centers with a unique $Z$-in distorted octahedral one in the central position. Binding of different anions to the copper(II) centers controls the nuclearity of the reaction products and tuning of the self-aggregate process within same ligand environment $(\mu-\text{H}_4\text{L}_1^-)$. Magnetic properties of the compounds have been studied both experimentally and using DFT calculations, revealing moderate to strong antiferromagnetic coupling in all aggregates.
Introduction

In recent years the synthesis, spectroscopic and structural characterization, and catalytic evaluation of Cu\(_2\)-compounds have received substantial attention from the viewpoint of their accessibility, structures and their relevance to active sites in biology.\(^1\) For example, the Cu\(_2\) species in tyrosinase is present in a variety of plant and animal tissues.\(^2\) Medicinal Cu\(^{II}\) chemistry deals with opportunities in relation to metal ion coordination for the design and use as therapeutic agents that are not attainable through simple organic molecules which are pharmaceutically active.\(^3,4\) In these class of copper(II) complexes the exploitation of the range of coordination numbers and geometries, formation of multinuclear aggregates, thermodynamic and kinetic control, and intrinsic properties of the complex species offer openings for their screening for reactivity patterns and properties. Synthetic endeavor for mimicking Cu enzymes is dependent on the nature and type of the donor atoms available on the ligand backbone. Schiff base ligands are one such type and their Cu\(^{II}\) complexes are known to show \textit{in vitro} anticancer potential against the HepG2 cell line.\(^5\) Also Cu\(_2\) compounds of the ligand system are known to promote the hydrolytic cleavage of double-strand plasmid DNA and inhibit the growth of GLC4 and K562 cells without affecting macrophage viability.\(^6,7\) The presence of phenol unit and adjacent imine functions within ligand frame brings two Cu\(^{II}\) (\(S = \frac{1}{2}\)) centers into close proximity and allows interaction of their electron spins. Available oxygen donors lead to the formation of a Cu\(_2\)O\(_2\) type diamond core with modulation from HO\(^-\) or O\(^{2-}\) bridges, required for further growth in aggregation. The magnetic interaction is subjective to Cu···Cu separation, the Cu–O distances, the Cu–O–Cu angles, the dihedral angle between the planes within the Cu\(_2\)O\(_2\) core, the out-of-plane displacement of substituents attached to the bridging groups, and the coordination geometry around each Cu\(^{II}\) centre.\(^1\)

Design and synthesis of coordination aggregate complexes, formed by the incorporation of a group of Cu\(_2\)-fragments, is dependent on the choice and selection of the ligand system giving ligand anion bound \{Cu\(_2\)L\} species in solution. Collapse of two such \{Cu\(_2\)L\} fragments provides structurally different [Cu\(_4\)] aggregates having cubane,\(^8\) tetrahedron,\(^9\) stepped-cubane,\(^10\) rhomboid,\(^11\) and double-cubane\(^12\) structures. Weak bridges like NO\(_3\)\(^-\) and ClO\(_4\)\(^-\) are not appropriate to show similar interactions with \{Cu\(_2\)L\} moiety. Whereas carboxylate bridges can adopt \(\mu_{1,1}, \mu_{1,3} \) or \(\mu_{1,1,3}\) modes during such process. Synergistic aggregating potential of HO\(^-\) and RCO\(_2\)\(^-\) groups have been shown to bind \{Cu\(_2\)L\} fragments for Cu\(_6\) aggregates.\(^13\)
From the functional behavior and application point of view the synthesized copper(II) complexes of phenol based multiple alcohol side arm bearing ligand system can be examined for their biomimetic catalytic oxidation efficiency and biophysical studies. The hydrogen bonding efficiency and hydrogen bonded network generation in solution from the coordinated and dangling alcohol arms of bound ligand could be useful during association of substrate molecules. In the present work, reactions of H₅L₁ (Chart 1, left) have been explored with Cu(ClO₄)₂, Cu(NO₃)₂ and Cu(O₂CCF₃) to identify the role of anions in directing the assembly process. H₃bemp, a close analog of H₅L₁, (Chart 1, 2,6-bis-[2-hydroxyethylimino)-methyl]-4-methylphenol; right) is known to give other type of [Cu₄] aggregate. Thus the primary coordinating ability of H₅L₁ and in situ generated HO⁻ ions has been judged for aggregation in presence of nitrate, perchlorate and trifluoroacetate ions (Chart S1, ESI). Interestingly three complexes of varying nuclearity has been achieved in [Cu₂(µ–H₄L₁)(µ–OH)(µ₁,₃–NO₃)(NO₃)(OH₂)]·H₂O (1), [Cu₄(µ–H₅L₁)₂(µ–OH)₂(µ₁,₃–ClO₄)(OH₂)₂](ClO₄)₂·2H₂O (2) and [Cu₅(µ–H₅L₁)₂(µ₂–OH)₂(µ₁,₃–O₂CCF₃)₂(O₂CCF₃)₂](CF₃COO)₂ (3). The reproducible synthesis, spectroscopic characterization, molecular structures determination and magnetic properties are described, discoursed and rationalized.

**Chart 1** H₅L₁ (used in this work) and related H₃bemp

**Results and Discussion**

**Synthetic Protocol**

[2,6-bis-{(1,3-dihydroxy-2-methylpropan-2-ylimino)methyl)}-4-methylphenol] (H₅L₁) was obtained in good yield (Scheme S1, ESI) from a reaction of 2,6-diformyl-4-methylphenol and 2-amino-2-methyl-1,3-propanediol in MeOH under stirring and refluxing condition. Reactions of H₅L₁ were examined with different copper(II) salts, as summarized in Scheme 1, to identify the role of anionic groups in directing the self-aggregation processes. Reaction of Cu(NO₃)₂·3H₂O and H₅L₁ in presence of CH₃CO₂Na as base in a 2:1:2 ratio in MeOH-MeCN medium at room temperature stirring condition afforded a greenish solution, from which dark-green crystals of 1 were subsequently isolated in 54% yield. Use of other
Carboxylate salts (propionate, trifluoroacetate) as base did not alter the formation and yield of 1 by incorporating these anions in the molecules. The elemental analysis and single-crystal X-ray diffraction studies confirm the composition of 1 as \[ \text{[Cu}_2(\mu-\text{H}_4\text{L1})(\mu-\text{OH})(\mu_{1,3-}\text{NO}_3)(\text{NO}_3)(\text{OH}_2)]\cdot\text{H}_2\text{O}. \]

\[
\text{H}_5\text{L1} + 2\text{Cu(NO}_3)_2 \cdot 3\text{H}_2\text{O} + 2\text{CH}_3\text{COONa} \rightarrow \text{CH}_3\text{OH} + \text{CH}_3\text{OCN} \rightarrow \text{[Cu}_2(\mu - \text{H}_4\text{L1})(\mu - \text{OH}) \quad \ldots(1)
\]

\[
(\mu_{1,3 - \text{NO}_3})(\text{NO}_3)(\text{OH}_2)] \cdot \text{H}_2\text{O} + 2\text{CH}_3\text{COOH} + 2\text{NaNO}_3 + 3\text{H}_2\text{O}
\]

Reaction of Cu(ClO\text{4})\text{2} \cdot 6\text{H}_2\text{O with H}_5\text{L1 in presence of CH}_3\text{CO}_2\text{Na in MeOH solution provided a green reaction mixture from which green block shaped crystals of 2 were isolated in 69% yield. Unique bridging coordination of ClO\text{4}^- in presence of available CH}_3\text{CO}_2^- ions result [Cu}_4\text{]} complex 2 (eq 2) through a different self-aggregation route without any abound acetate group.

\[
2\text{H}_5\text{L1} + 4\text{Cu(ClO}_4)_2 \cdot 6\text{H}_2\text{O} + 4\text{CH}_3\text{COONa} \rightarrow \text{CH}_3\text{OH} \rightarrow \text{[Cu}_4(\mu - \text{H}_4\text{L1})_2(\mu - \text{OH})_2
\]

\[
(\mu_{1,3 - \text{ClO}_4})(\text{OH}_2)_3)(\text{ClO}_4)_3 \cdot 2\text{H}_2\text{O} + 4\text{CH}_3\text{COOH} + 4\text{NaClO}_4 + 18\text{H}_2\text{O} \quad \ldots(2)
\]

Reaction of Cu(CF\text{3}COO)\text{2} \cdot 4\text{H}_2\text{O and H}_5\text{L1 in 2.5:1 molar ratio in stirring MeOH and in absence of any added base, afforded a dark solution, from which 3 (eq 3) was subsequently isolated from the reaction medium as green block shaped single crystals suitable for X-ray analysis in 81% yield.

\[
2\text{H}_5\text{L1} + 5\text{Cu(CF}_3\text{COO})_2 \cdot 4\text{H}_2\text{O} \rightarrow \text{CH}_3\text{OH} \rightarrow \text{[Cu}_5(\mu - \text{H}_4\text{L1})_2(\mu_3 - \text{OH}_2)(\mu_{1,3 - \text{O}_2\text{CCF}_3})_2(\text{O}_2\text{CCF}_3)_2)(\text{CF}_3\text{COO})_2 + 4\text{CF}_3\text{COOH} + 18\text{H}_2\text{O} \quad \ldots(3)
\]

The elemental analysis and X-ray diffraction data established the products as [Cu}_4(\mu - \text{H}_4\text{L1})_2(\mu - \text{OH})_2(\mu_{1,3 - \text{ClO}_4})(\text{OH}_2)_3)(\text{ClO}_4)_3 \cdot 2\text{H}_2\text{O and [Cu}_5(\mu - \text{H}_4\text{L1})_2(\mu_3 - \text{OH}_2)(\mu_{1,3 - \text{O}_2\text{CCF}_3})_2(\text{O}_2\text{CCF}_3)_2)(\text{CF}_3\text{COO})_2 for 2 and 3 (Scheme 1).

**Scheme 1** Synthetic route for 1–3
Evaluation by FT-IR Spectroscopy

The FT-IR spectra for 1, 2 and 3 initially identifies the presence of H₄L¹⁻ in {Cu₂(OH)L¹} fragments (Figure S1, ESI). The $\tilde{\nu}_{\text{C}-\text{N}}$ stretching frequencies were detected at 1635, 1636 and 1644 cm⁻¹ for 1, 2 and 3 respectively. The phenoxydo-bridge shows the $\tilde{\nu}_{\text{C}-\text{O}}$ stretching at 1551, 1560 and 1560 cm⁻¹ for 1, 2 and 3 respectively. Broad bands at 3400 cm⁻¹, 3338 cm⁻¹ and 3348 cm⁻¹ for 1, 2, and 3 respectively were found for alcohol arms and bound hydroxido groups. The coordination of nitrate group in 1 is verified by the presence of the $\tilde{\nu}_3(E')$ mode of vibration at 1384 cm⁻¹.¹⁷,¹⁸ Peaks at 1433 cm⁻¹ ($\tilde{\nu}_{\text{as}}$), 1329 cm⁻¹ ($\tilde{\nu}_s$) are assigned to asymmetric and symmetric stretching modes of vibration of the bridging NO³⁻ group.¹⁹,²⁰ For complex 2, the presence of two different types of perchlorate anions have been detected by identifying the characteristic $\tilde{\nu}_{\text{as}}$ and $\tilde{\nu}_s$ stretching vibration modes. For anionic ClO₄⁻ anions ($T_d$ symmetry), the $\tilde{\nu}_3(T_2)$ ($\nu_{\text{ClO}}$) band appeared at ~1088 cm⁻¹ and $\tilde{\nu}_d(T_2)$ ($\delta_{\text{DOCIO}}$) mode at 626 cm⁻¹.²⁰ In case of bridging ($C_{2V}$) ClO₄⁻ anion the $\nu_3$ mode split into three components at 1142, 1111 and 1088 cm⁻¹.¹³,²¹ In 3, two types of trifluoroacetate groups were detected by the presence of broad asymmetric stretching vibrations ($\tilde{\nu}_{\text{as(COO)}}$) at 1670 cm⁻¹ and medium intense symmetric stretching vibrations ($\tilde{\nu}_{\text{s(COO)}}$) at 1431 and 1329 cm⁻¹ respectively. The characteristic $\Delta \tilde{\nu}$ value for free CF₃CO₂⁻ group is higher (341 cm⁻¹) compared to the bridged and chelated ones ($\Delta \tilde{\nu}=239$ cm⁻¹).²²,²³ The presence of electron withdrawing fluorine atoms moves the asymmetric stretching frequency to higher region.²⁴
**Powder X-ray Diffraction Patterns**

The powder XRD patterns of the bulk materials of 1-3 were collected using a Bruker AXS X-ray diffractometer and compared with the simulations derived from the single crystal X-ray diffraction data. Figure S2 (ESI) shows that the powder patterns are in good agreement with the simulated data. The difference in intensity is due to the orientation of the powder samples during experiment. The similarity infers that the prepared powder samples are pure and have comparable composition to that of the single crystals. The molecular composition of single crystals and powder samples were same, thus corroborating the reproducibility of powder samples in multiple synthetic attempts.

**Electronic Spectral Transitions**

In MeOH solutions complexes 1, 2 and 3 register several characteristic bands within 800-200 nm range. The band maxima at 645 nm (ε = 67 L mol⁻¹ cm⁻¹), 666 nm (ε =129 L mol⁻¹ cm⁻¹), and 678 nm (ε =104 L mol⁻¹ cm⁻¹) are assigned to the low intensity d-d transitions for 1, 2 and 3, respectively. The copper(II) centers in octahedral O₆, O₅N and square pyramidal O₄N coordination geometry do register electronic spectral behavior of [Cu(en)₂(H₂O)₂]²⁺ showing corresponding band at 545 nm (ε = 64 L mol⁻¹ cm⁻¹). Free ligand H₅L₁ in MeOH solution registers n→π* transition at 338 nm (ε = 3600 L mol⁻¹ cm⁻¹) and π→π* transition at 232 nm (ε = 18200 L mol⁻¹ cm⁻¹). The intense absorptions just below 400 nm are dominated by copper(II) bound ligand-based absorptions. The characteristic phenoxido O to Cu II charge transfer transition (LMCT) is observed at 368 nm (ε = 7600 L mol⁻¹ cm⁻¹) for 1, 374 nm (ε = 10500 L mol⁻¹ cm⁻¹) for 2, and 371 nm (εₘₐₓ = 13700 L mol⁻¹ cm⁻¹) for 3. Interestingly the other ligand based strong absorption bands were seen at 259 nm with ε values of 3510, 52400 and 64500 L mol⁻¹ cm⁻¹ for 1, 2 and 3, respectively. These are due to Cu II bound intra-ligand π → π* transitions mainly centered on Cu II-bound >C=N functions (Figure S3, ESI).

**Description of Crystal Structures**

[Cu₂(µ-H₄L₁)(µ-OH)(µ₁,3-NO₃)(NO₃)(OH₂)]·H₂O (1). X-ray structural analysis shows that 1 crystallized in the monoclinic P2₁/n space group. The interatomic connectivity of the asymmetric unit of 1 is shown in Figure 1, and selected bond distances and angles are given in Table S1 (ESI). The neutral complex is formed from one coordinating Schiff base ligand, two nitrate ions and a lattice water molecule. Two octahedral Cu II centers in the dinuclear unit are bridged by the in plane phenoxido and hydroxido groups resulting in short Cu···Cu separation of 2.896 Å in two adjacent octahedral coordination geometry. Within the Cu₂O₂
diamond core the Cu–O distances range from 1.910 to 1.958 Å, where the shortest distances are established from phenoxido group. One nitrate group in µ1,3–bridging mode also binds the two copper (II) centers. The O–N–O bridging part of nitrate group connects two almost coplanar CuO₃N planes from two apical sides and register a dihedral angle of 5.68°.

Figure 1 Molecular view of I with partial atom numbering scheme. H-atoms are omitted for clarity. Color code: Cu, green, N, blue, O, red, C, black

The N₂O₅ donors of H₂L1 provided two phenoxido-bridged meridional O₂N halves to bind two Cu¹I centers. The second –CH₂OH arms on each side remain pendant and participate in hydrogen bonding interactions within the crystal lattice. For NO₅ octahedral geometry, the Cu–N and Cu–O distances range from 1.927 to 2.475 Å for Cu1 and 1.910 to 2.574 Å for Cu2 centers respectively. Cu–O bond from bridging nitrate provides the longest (2.574 Å) separation above the CuO₃N plane (for Cu1 center). The other Cu–O bond distance (for Cu2 center) from bridging nitrate is 2.432 Å, confirming the asymmetric bridging by NO₅¯ group. The water O atom is found at 2.549 Å from Cu2, whereas the O atom of the NO₅¯ group is at 2.475 Å from Cu1.

Two discrete molecular units are engaged in intermolecular hydrogen bonding interactions involving the coordinated hydroxyl group of a ligand arm with a hydroxide bridge (O4–H4···O2) and dangling hydroxide group (O3–H3···O12), producing a hydrogen bonded dimeric assembly of two Cu₂ units. The coordinated water molecule functions both as a hydrogen bond donor and acceptor and form hydrogen bonding interactions with the nitrate ion (O11–H11B···O7, O11–H11A···O9) and lattice trapped water molecule (O1W–H1WA···O11). The lattice water molecule also involved in hydrogen bonding interaction with available nitrate (O1W–H1WB···O7), bridging hydroxide (O2–H2···O1W) and dangling hydroxyl group of ligand (O12–H12···O1W). Such hydrogen-bonding interactions extend the hydrogen bonded dimeric assembly in two dimensions producing a two dimensional hydrogen bonded network (Figure 2).
\[ \text{Cu}_4(\mu-H_4L_1)\text{Cu}_4(\mu-OH)\text{Cu}_4(\mu_{1,3}\text{ClO}_4)(OH_2)_2\text{ClO}_4)_3\cdot2\text{H}_2\text{O} \] (2). The structure of the tetranuclear Cu\textsuperscript{II} complex is shown in Figure 3. Selected interatomic distances and bond angles are provided in Table S1 (ESI). It crystallizes in the triclinic \(P\bar{1}\) space group and consists of a tricationic \([\text{Cu}_4(\mu-H_4L_1)\text{Cu}_4(\mu-OH)\text{Cu}_4(\mu_{1,3}\text{ClO}_4)(OH_2)_2]\) part and three ClO\textsuperscript{4}\textsuperscript{−} anions. X-ray diffraction quality single crystals were obtained as perchlorate salts.

Anionic H\textsubscript{4}L\textsubscript{1}\textsuperscript{−} each providing a N\textsubscript{2}O\textsubscript{5} donor set is suitable to provide the Cu\textsubscript{4} complex with unique bridging mode of a perchlorate anion. Unlike complex 1, all the Cu\textsuperscript{II} centers are in square pyramidal geometry well suited for lone inter-dimeric bridging support from one ClO\textsubscript{4}\textsuperscript{−} ion. Each H\textsubscript{4}L\textsubscript{1}\textsuperscript{−} unit provides basal coordination support to two Cu\textsuperscript{II} ions which are tightly bridged by hydroxido O atom within the same plane. Herein all four Cu\textsuperscript{II} ions are in NO\textsubscript{4} square pyramidal geometry and the Cu–N and Cu–O distances range from 1.912 to 2.638 Å. Cu–O bond distances from bound water molecules are 2.354 and 2.357 Å,\textsuperscript{27} whereas Cu–O bonds from bridging perchlorate (µ\textsubscript{1,3}\text{ClO}_4\textsuperscript{−}) O atoms give longest (2.584 Å for Cu4
and 2.638 Å for Cu2) Cu–O bonds below and above the two CuO$_3$N square planes. Very weak coordination of these O atoms shows only square-pyramidal environment around each metal ion. As a result the Addison parameter $\tau$ remains in the 0.007–0.055 range (Figure 4).

The metallic core of 2 resulting from the perchlorate–mediated assembly takes the shape of a trapezium (Figure 4). Perchlorato bridged Cu$^{II}$ centers (Cu···Cu) are 7.142 Å apart compared to the phenoxido-hydroxido bridged separation of 2.921 Å. The open side registers a Cu···Cu separation of 6.747 Å. The presence of two water molecules in 2 shows hydrogen bonding network with dangling ligand alcohol arms.

For 2 the coordinated hydroxyl group of ligand arm was involved in intermolecular hydrogen-bonding interactions with dangling hydroxyl group (O4–H4···O16) and with bridging –OH group (O3–H3···O13, O14–H14···O2). Hydrogen bonding interactions of the dangling hydroxyl groups with each other (O6–H6···O17) and with lattice water molecule (O5–H5···O1W, O16–H16···O1W) are also observed. The bridging hydroxyl group and coordinated water molecule remain involved in intramolecular hydrogen-bonding interactions with bridging perchlorate (O13–H13···O10, O18–H18B···O9). Hydrogen bonding interactions are also observed between coordinated water molecule and perchlorate counter anion (O18–H18A···O21) as well as dangling –OH group and perchlorate counter anion (O17–H17···O30). Further expansion through hydrogen bonding interactions occur via lattice water molecules (O1W–H1WA···O22, O2W–H2WA···O30, O2W–H2WB···O5, O1W–H1WB···O26, O15–H15···O2W) producing a 2D sheet structure (Figure 5). These non-covalent and non-bonding interactions play an important role in the crystal lattice with entrapment of anions and lattice solvent molecules.
[Cu₅(µ-H₄L1)₂(µ₂-OH)₂(µ₁,3-O₂CCF₃)₂(O₂CCF₃)₂](CF₃COO)₂ (3). Complex 3 crystallizes in the monoclinic P2₁/c space group with Z=2 and crystallographic inversion center on Cu₃ atom. The selected metric parameters for the structure are summarized in Table S1 (ESI). The asymmetric unit of 3 contains half of the molecular structure and one trifluoroacetate counter anion. A centrosymmetric structure containing five CuⅡ centers within a [Cu₅O₄] core generates through intermetallic connections from H₄L₁⁻ and water derived HO⁻ anions. Four trifluoroacetate groups connect the ligand bound CuⅡ centers and coordinate to the central CuⅡ ion in bidentate fashion (Figure 6).

Within the Cu₂(µ-H₄L1) fragments two square pyramidal CuⅡ centers are connected by the in plane phenoxido and hydroxido groups and bringing the CuⅡ centers at a distance of 2.909 Å. The square pyramidal natures of these CuⅡ centers are known from the calculated Addison parameters (τ) of 0.01 and 0.12. The Cu–O distances within this Cu₂O₂ diamond core range.
from 1.924 to 1.995 Å. Interestingly the central Cu3 is a unique one and adopts a typical O6 octahedral coordination geometry. The environment is developed from forced bidentate coordination of two trifluoroacetato (O7, O8, O7* and O8*) groups in a basal plane. The corresponding Cu–O bond distances were longer in 1.976–2.048 Å range and characteristic with four long distances in case of copper(II) ion in octahedral geometry and susceptible to Jahn-Teller distortion. Two apical positions around this central Cu atom are occupied by the μ3–OH group providing two short Cu3–O2 bonds of 1.919 Å separations, clearly demonstrating an unusual example of a compressed coordination octahedron within a [Cu5] compound (Figure 7c).

Two such μ3–OH bridges provide lone supports to the central copper(II) for aggregation. For such bridge two types of Cu–O OH bond distances are observed, the smaller (1.919 Å) ones connect the Cu3 and longer ones (1.975 and 1.995 Å) are present within Cu2(μ-H4L1) fragments. In one of our previous work we have shown that the affinity of the copper(II) ions for long distance binding along the apical Jahn-Teller axis is responsible for the binding of six acetate ions, unlike the fluoroacetate anions in the present work, in three different coordination modes. The central copper(II) ion adopts a compressed octahedral geometry owing to the compression in apical direction, with a significant compression of the apical bonds by 0.093 Å, when compared by considering six Cu–O bonds around Cu3. In the present case the forced coordination from two alcohol side arms and availability of mononuclear copper(II) trifluoroacetate fragment in the reaction medium are responsible for the formation of the different type of [Cu5] coordination aggregate. Two other trifluoroacetate groups binding Cu1 (Cu1*) and Cu2 (Cu2*) in μ1,3 bridging mode pushes the ligand bound Cu2 fragments to entrap the central copper(II) in tetragonally compressed fashion by two μ3–OH bridges (Figure 7b). Compared to ‘paddlewheel’ structures in case of copper(II) acetates we take the advantage of perfluorinated carboxylate salt of copper(II) in the synthesis, which tend to form monomeric complexes. Within the aggregate three different types of trifluoroacetate groups were found (Figure 7a). These were μ1,3 bridging attached to Cu2(μ-H4L1) fragments, bidentate chelating for the central Cu3 atom and anionic, outside the coordination sphere. For bridging trifluoroacetate the Cu–O–C–O/O–C–O–Cu torsion angle is 17.16°/5.28. The Cu5 core of 3 is formed from trapping of central copper(II) center by two Cu2(μ-H4L1) fragments taking a shape of a modified hourglass (Figure 7d). The central copper(II) center chelated by trifluoroacetate anions remains 3.194 and 3.345 Å apart from CuII centers within Cu2(μ-H4L1) fragments. The ligand bound and phenoxido-hydroxido bridged Cu···Cu separation is shortened to 2.909 Å.
Figure 7. (a) Core view of the $\text{[Cu}_5\text{]}$ unit; (b and c) coordination environments of $\text{Cu}_2(\mu-\text{H}_2\text{L1})$ fragment and central Cu$^{\text{II}}$ centre; (d) phenoxido and hydroxido bridges with 3

Here also, the hydrogen-bonding interactions between the dangling alcohol arms of the ligand and trifluoroacetate anions play the main role in the crystal packing with entrapment of trifluoroacetate anions. Strong hydrogen-bonding interactions are seen between chelated imine alcohol arms bearing bound OH functions of the ligand with dangling alcohol arm (O3–H3A···O10) and with anionic trifluoroacetate (O4–H4···O12). Also dangling free alcohol arm showed engagement with each other (O10–H10···O9) and with trifluoroacetate (O9–H9···O12).$^{32}$ Involvement of $\mu_3$–OH group is observed with trifluoroacetate (O2–H2···O11) for a hydrogen bonded 2D sheet structure (Figure 8).

Figure 8 Intermolecular hydrogen bonding network resulted a 2D structure; inset: enlarged view of hydrogen bonding interactions involving trifluoroacetates.

**Structural Comparison of 1 with 2 and 3**

In all three compounds each Cu$^{\text{II}}$ center is bound by single imine N atom of ligand except Cu3 in 3. The average Cu–N bond lengths in all the compounds are in 1.913 to 1.939 Å range. All together a variety of Cu–O bonds were found which differ from each with respect
to the geometry and environment, e.g., square pyramidal O₄N, and octahedral O₃N and O₆.

Nine different Cu–O bonds are found from coordination of water, alcohol hydroxyl, bridging nitrate and perchlorate, bridging and chelated trifluoroacetate, bridging phenoxido and hydroxido groups. The longest ones in this set are from bridging perchlorato O atoms and next to these are from bridging nitrato O atoms. Water coordinated to the copper(II) center makes a large (2.549 Å) Cu–O separation in complex 1 compared to those in complex 2 (2.357 and 2.354 Å). The Cu–O bonds from bridging trifluoroacetate groups are longer than the chelated ones. The Cu–O bonds from bridging phenoxido group within Cu₂(µ-LH₄L₁) part in all the compounds are in 1.910–1.939 Å range. In 1 two types of coordination from nitrate anions stabilize the Cu₂(µ-H₄L₁) in a particular manner different from a single perchlorato anion for clipping two Cu₂(µ-H₄L₁) units in 2. The µ₃–OH connection in 3, connecting three Cu⁺⁺ centers, provide one short (1.919 Å) and two long (1.975 and 1.995 Å) Cu–O bonds. Interestingly the coordination of Cu⁺⁺ center to alcoholic OH arms has no effect with regard to C–O separation compared to the free and dangling arms. The trifluoroacetato-bridge clips the two Cu⁺⁺ centers in 3 and also in bidentate chelating mode to the central Cu⁺⁺ ion. In both the two cases the Cu–O separations are different, 2.201 and 2.301 Å in case of bridging mode and 1.976 and 2.048 Å for chelating mode. These two different modes of binding are responsible for varying the C–O bond distances in 3. For bridging trifluoroacetate groups the C–O distances are 1.236 and 1.246 Å, whereas for chelating mode the same are 1.138 and 1.137 Å. The Cu–O–Cu angles within the {Cu₂(µ-LH₄L₁)(OH)} fragments is in 97.73–99.59° range for phenoxido bridges vs. 94.24–97.08° for hydroxido bridges. In 3, the Cu–O(H)–Cu angle is further shortened due to the involvement of the µ₃–OH connector required for the growth of [Cu₅] aggregate. In all the complexes, due to involvement in hydrogen bonding interactions, the H atom of the bridging hydroxide group remains out of plane with respect to the Cu₂O₂ diamond core. The geometrical features with respect the Cu–O(Ph)–Cu angle (α) (97.95 for 1, 99.59 and 99.33 for 2, and 97.74 for 3), out-of-plane angle deviation (φ) (12.83 for 1, 4.77 and 5.20 for 2 and 5.57 for 3) and Cu–O–Cu–O torsion angle (τ) for the positioning of phenyl ring of bridging phenoxido group with respect to the molecular plane bearing the two Cu⁺⁺ ions can modify the nature of magnetic interactions in the solid state (vide supra) (see Figure S4 in ESI).

**Mass Spectroscopic Evidence of Fragments**

**High Resolution ESI-MS Analysis.** The existence of Cu₂(µ-LH₄L₁) based fragments in solution were established from mass spectral (ESI-MS positive) analysis of compounds 1–3
in MeOH solutions (Figures S5-S7 in ESI). Anion dependent charge neutralization of Cu$_3$(µ-H$_4$L1) unit formed in solution resulted in the [Cu$_2$], [Cu$_4$] and [Cu$_5$] aggregates as compounds 1–3. The analysis establishes the populations of key fragments in solution. All three compounds exhibited a common base peak at $m/z = 240.02$ which can be attributed to [Cu$_2$(µ–H$_4$L1)(µ–OH)]$^{2+}$ (C$_{17}$H$_{26}$Cu$_3$N$_2$O$_6$; Calcd 240.02) fragment in the solution phase. For 1, the nitrato-bridged structure found in the solid state is not survived in MeOH solution medium and no other peaks of even smaller $m/z$ value corresponding to any rational fragment is found in this analysis. The mass spectrum of 2 in MeOH medium records two peaks at $m/z = 562.97$ and 580.98 corresponding to the fragments [Cu$_2$(µ–H$_4$L1)(ClO$_4$)]$^+$ (C$_{17}$H$_{26}$ClCu$_2$N$_2$O$_6$; Calcd 562.99) and [Cu$_2$(µ–H$_4$L1)(µ–OH)(ClO$_4$)]$^+$ (C$_{17}$H$_{26}$ClCu$_2$N$_2$O$_{10}$; Calcd 580.98). Compared to the base peak of $m/z = 240.02$ for [Cu$_2$(µ–H$_4$L1)(µ–OH)]$^{2+}$ the relative intensity of these two peaks are less than 10% indicating poor abundance in solution. This is due to the fact that monodentate coordination of perchlorato anion is less stable than the bridging version as identified in 2 in the solid state (vide supra). In these two cases the monocationic charge states correspond to the species where one perchlorato anion is associated with the complex fragments. In case of complex 3 the mass spectrum displays a peak at $m/z = 575.01$ corresponding to [Cu$_2$(µ–H$_2$L1)(CF$_3$COO)]$^+$ (C$_{19}$H$_{25}$Cu$_2$F$_3$N$_2$O$_7$; Calcd 575.01). The abundance of this species was found to be close to 40% of the base peak implying the presence of this species in considerable amount in solution compared to those obtained for compound 2. Clearly this may be attributed to better coordinating potential of CF$_3$COO$^-$ as compared to both NO$_3^-$ and ClO$_4^-$.$^-$ A low intensity peak at $m/z = 596.99$ corresponding to [Cu$_2$(µ–H$_2$L1)(CF$_3$COO)]Na$^+$ (C$_{19}$H$_{25}$Cu$_2$F$_3$N$_2$NaO$_7$; Calcd 596.99) was also present in the spectrum of 3.

**MALDI-TOF Analysis.** We have used this technique to create ions from all three compounds with nominal fragmentation. In search of higher nuclearity fragments in solution which might not have survived in the ESI process the compounds were subjected to MALDI-TOF analysis using 2,5-dihydroxybenzoic acid (DHBH) as the matrix. The spectrum of all the complexes exhibit a base peak at $m/z = 339.61$ corresponding to the protonated ligand [H$_5$L1]$^+$ (C$_{17}$H$_{27}$N$_2$O$_5$; Calcd 339.19) along with a peak at $m/z = 616.65$ arising from [Cu$_2$(µ–H$_4$L1)(DHB)]$^+$ (C$_{24}$H$_{36}$Cu$_2$N$_2$O$_5$; Calcd 616.05) (Figures S9–S11 in ESI). This later species is formed from the bridging coordination of carboxy end of the anionic matrix with dinuclear fragments observed in ESI-MS analysis ([Cu$_2$(µ–H$_4$L1)(µ–OH)]$^{2+}$). For 2 a peak at $m/z = 371.42$ is observed due to the presence of [Cu$_4$(µ–H$_4$L1)$_2$(µ–OH)$_2$(µ$_{1,3}$–ClO$_4$)(OH)$_2$)$_3$$^{3+}$ (C$_{34}$H$_{58}$ClCu$_4$N$_4$O$_{19}$; Calcd 371.68) species in solution. The spectrum of 3 shows peaks at $m/z$
= 678.61, 802.96, 864.94 and 1387.36 corresponding to the species \([\text{Cu}_2(\mu-H)L_1)(\mu-\text{OH})(\text{CF}_3\text{COO})(\text{CH}_3\text{OH})_2]\text{Na}^+\) (Caled 679.05), \([\text{Cu}_3(\mu-H)L_1)(\mu-\text{OH})(\text{CF}_3\text{COO})_2(\text{CH}_3\text{OH})_2]^+\) (Caled 802.96), \([\text{Cu}_3(\mu-H)L_1)(\mu-\text{OH})(\text{CF}_3\text{COO})_2(\text{CH}_3\text{OH})_3]\text{Na}^+\) (Caled 865.01) and \([\text{Cu}_5(\mu-H)L_1)(\mu-\text{OH})(\text{CF}_3\text{COO})_2(\text{CH}_3\text{OH})_2(\text{H}_2\text{O})]\text{Na}^+\) (Caled 1387.06). This study thus clearly indicates the power of relatively soft technique responsible for low fragmentation while obtaining ions of aggregate compounds 2 and 3. Thus in the gas phase it is possible to detect the formation of tetranuclear \([\text{Cu}_4(\mu-H)L_1)(\mu-\text{OH})(\mu_1,3-\text{ClO}_4)]^{3+}\) and pentanuclear \([\text{Cu}_5(\mu-H)L_1)(\mu-\text{OH})(\text{CF}_3\text{COO})_2(\text{CH}_3\text{OH})_3(\text{H}_2\text{O})]\text{Na}^+\) species which ultimately crystallize in the solid state.

**Reasoning behind the Isolated Products**

The tuning of reaction conditions under which the \([\text{Cu}_2]\), \([\text{Cu}_4]\) and \([\text{Cu}_5]\) family formed, were analyzed in terms of anion dependence and synthetic environments. Competitive coordination of auxiliary anions to the metal ion centers influences the coordination aggregation process. Multinuclear bridging ability to connect more than two metal ions by \(\text{NO}_3^-\) ions is not frequently documented in the literature. Herein use of nitrate salt in the reaction medium trapped the product as \([\text{Cu}_2L]\) (1). Isolation of this dinuclear precursor from the reaction medium thus tempted us to examine the role of other anions for spontaneous self-aggregation reaction. Formation of \([\text{Cu}_4]\) species as complex 2 is dependent on the nature of the anion and the presence of perchlorate anion being a factor in its isolation as X-ray quality crystals. The highest possible bridging capacity of twelve metal ions by single \(\text{ClO}_4^-\) anion is known in \([\text{Cu}_{12}\text{La}_6]\) complex. Connection of two \(\text{Cu}_2L\) fragments by a single \(\text{ClO}_4^-\) anion to provide the \([\text{Cu}_4]\) complex is not common in the literature. The \(\mu_{1,3}\) mode of one non-planar perchlorate anion functioning as bridging ligand led to the isolation of 2 while other three perchlorate anions were required for charge neutralization only. In case of 3 the reaction with Cu(\(\text{CF}_3\text{CO}_2\))\(_2\)·4\(\text{H}_2\text{O}\) on the other hand resulted in the trapping of trifluoroacetate chelated copper(II) within the resulting \([\text{Cu}_3]\) complex. The trapping of \([\text{Cu}(\text{CF}_3\text{CO}_2)]\) unit by two \([\text{Cu}_2(\text{H}_4\text{L}1)(\text{CF}_3\text{CO}_2)]\) units, through \(\mu_3-\text{OH}\) bridging, is responsible for the generation of new \([\text{Cu}_3]\) structural motif in Cu\(^{II}\)-carboxylate chemistry.

**Electrochemistry**

Further characterization in solution was achieved from cyclic voltammetric measurements in MeOH for 1-3. The characteristic voltammetric responses for \{Cu\(_2(\mu-H)L_1)\}\) fragments
coming from all the compounds indicate their stability and aggregation behavior. A three-electrode setup was used consisting of a Pt working electrode, an Ag/AgCl reference electrode and a platinum auxiliary electrode.

**Figure 9** Cyclic voltammograms for (i) $\text{H}_5\text{L1}$, (ii) $\text{I}$ [inset (a) from 0.0 V to -1.2 V, (b) from 0.0 V to 1.2 V], (iii) $\text{2}$ [inset (a) from 0.0 V to -1.2 V, (b) from 0.0 V to 0.8 V] and (iv) $\text{3}$ [inset (a) from 0.0 V to -1.2 V, (b) from 0.0 V to 1.2 V] in MeOH containing 0.1 M $\text{nBu}_4\text{NClO}_4$ as supporting electrolyte at 298 K, at platinum working electrode, at a scan rate of 100 mV s$^{-1}$, using Ag/AgCl as the reference electrode.

For $\text{H}_3\text{L1}$ in MeOH an irreversible anodic peak at 0.74 V at a sweep rate of 0.1 V s$^{-1}$ was obtained for ligand centered oxidation (Figure 9-i). For $\text{I}$ two closely spaced irreversible reductions were recorded at $E_{pc} = -0.520$ and $-0.779$ V respectively, for the electron transfer processes $\text{Cu}^{II}\text{Cu}^{II} \rightarrow \text{Cu}^0\text{Cu}^0$ and $\text{Cu}^0\text{Cu}^I \rightarrow \text{Cu}^0\text{Cu}^0$ within $\{\text{Cu}_2(\mu-\text{H}_4\text{L1})\}$ fragment. During the reverse scan a small peak at 0.115 V is found due to anodic stripping, which is absent when scanned between 0.0 V and 1.2 V. The irreversible oxidative response at $E_{pa} = 0.422$ V is most probably due to $\text{Cu}^{II}\text{Cu}^{II} \rightarrow \text{Cu}^{III}\text{Cu}^{III}$ oxidation (Figure 9-ii-b inset). On scan reversal
from 1.2 V, a small cathodic peak at $E_{pc} = 0.347$ V is seen for the reduction of unstable Cu$^{III}$Cu$^{III}$ fragment to Cu$^{II}$Cu$^{II}$ species ($i_{pc}/i_{pa} \neq 1$). The response for 2 is similar to that found for 1 with irreversible cathodic reductions at $E_{pc} = -0.503$ V and $-0.665$ V respectively with anodic stripping at 0.155 V. The irreversible oxidative response for Cu$^{II}$Cu$^{II}$ → Cu$^{III}$Cu$^{III}$ occurs at $E_{pa} = 0.468$ V (Figure 9-Lii-Lb inset).

The nature of voltammogram for 3 is different from both 1 and 2. Now two well separated irreversible reductive responses were observed at $E_{pc} = -0.425$ V and $-0.822$ V corresponding to Cu$^{II}$Cu$^{II}$ → Cu$^{I}$Cu$^{I}$ and Cu$^{II}$Cu$^{I}$ → Cu$^{0}$Cu$^{0}$. Herein the anodic stripping potential for Cu$^{0}$ from the electrode surface was found at 0.275 V. Scanning in the positive direction resulted two irreversible oxidations at $E_{pa} = 0.557$ V and $E_{pa} = 0.700$ V, of which the latter was due to ligand centered oxidation ($E_{pa}$ for H$_5$L1 = 0.730 V). The former response is due to Cu$^{II}$Cu$^{II}$ → Cu$^{III}$Cu$^{III}$ oxidation and the corresponding reduction of Cu$^{II}$Cu$^{II}$ is found at $E_{pc} = 0.296$ V. The $\Delta E_p$ value found to be 261 mV ($E_{1/2} = 0.426$ V) and $i_{pc}/i_{pa}$ ratio $\approx 1$ indicate some stability of Cu$^{III}$Cu$^{III}$ species within cyclic voltammetric time scale. The similar nature of the voltammograms for 1 and 2 suggest the existence of [Cu$_2$(μ–H$_4$L1)(μ–OH)]$^{2+}$ fragments in MeOH solutions in both the two cases as also evidenced from the ESI-MS analysis (vide supra). For 3 the distinct responses were found from species like [Cu$_2$(μ–H$_2$L1)(CF$_3$COO)]H$^+$ having 40% abundance to that of the base peak in ESI-MS analysis.

**EPR Spectra**

X-band EPR spectra for powdered samples of 2 and 3 were measured at 298 K and 75 K (Figure 10). An axial EPR signal was observed for 2 with $g_{||} = 2.264$ and $g_{\perp} = 2.04$ ($g_{av} = 2.114$) (Figure 10-i). The average nuclear hyperfine constants are, $A_{||} = 180.6$ G and $A_{\perp} = 41.66$ G and no resonances below 1050 G and above 3850 G were detected. For 3 a different type of axial spectrum was obtained (Figure 10-ii). At 75 K hyperfine features of the parallel component were observed with a small $A_{||}$ value of 65.79 G. The g-tensors were found to be $g_{||} = 2.28$ and $g_{\perp} = 2.10$. The weak intensity, forbidden half field ($\Delta M_S = \pm 2$) signals were observed at 1605 G (g = 4.298) and 1605.5 G (g = 4.296) for 2 and 3 respectively, clearly indicating the presence of spin-spin interactions. Unfortunately 1 gave poor quality spectra from which reliable A and g values could not be determined.
Figure 10 X-band EPR spectra at 298 K and 75 K for microcrystalline samples of 2 (i) and 3 (ii) respectively

The X-band EPR spectrum for a polycrystalline sample of 3 in the temperature range 5 K – 115 K is shown in Figure 11 (i). At 5 K one broad symmetric transition was observed at 3312 G without any hyperfine feature. The intensity of the signal is temperature dependent and showed maxima at 10 K. Further rise in temperature led to continuous decrease in signal intensity. The rate of decrease of signal intensity is high upto 35 K then gradually decreasing at higher temperatures. The g value at the same time increases from 2.09 at 5 K to 2.12 at 115 K. This sort of change in signal intensity with rise in temperature is consistent with the presence of antiferromagnetic coupling interactions within the copper centers. The spectrum at 10 K shows the highest intensity band centered at $g = 2.09$ ($3318 \text{ G for } \nu = 9.7196 \text{ GHz}$). Simulation of the spectrum$^{36,37}$ considering a system with three electrons and three copper nuclei lead to a reasonable fit yielding the following gyromagnetic factors: $g_1 \approx g_2 = 2.04$ ($g_\perp$) and $g_3 = 2.20$ ($g_\parallel$) (Figure 11(ii)). The $g_{\text{av}}$ value of 2.09 obtained from the simulated data is in good agreement with that found experimentally. The $g_{L}$ values of axial spectrum are related by the expression$^{38}$ $G = (g_{\parallel}-2.003)/(g_{\perp}-2.003)$ and is a measure for exchange interactions between the Cu$^{II}$ centers as measured in the solid state. When $G < 4.0$, there is substantial exchange interaction in the polycrystalline sample.$^{38,39}$ In our case the $G$ value for 3 is 5.324 demonstrating the presence of weak interactions which is in agreement with the magnetic and DFT studies. The absence of fine structures at the low temperatures ( < 75 K) is consistent with the presence of intercluster exchange interactions that tend to average out the fine structures arising from the multiplets.$^{40}$ The hyperfine features in the parallel component is observed at 75 K which is not distinguishable at room temperature due to signal broadening. The $A_{\parallel}$ value of 65.79 G is 2.7 times smaller than that obtained for isolated Cu$^{II}$ site and in case of complex 2 ($A_{\parallel} \approx 180 \text{ G}$).
Figure 11 (i) X-band EPR spectra at 5 K to 115 K for microcrystalline sample of 3. (ii) Overlapping experimental and simulated EPR spectra of 3 at 10 K

Magnetic Measurements

The temperature dependence of the molar magnetic susceptibility, $\chi_M$, of 1–3 was measured under a magnetic field of 1 T for 1 and 0.1 T for 2 and 3 over the temperature range 290 – 2 K. Plots of $\chi_M T$ versus temperature, $T$, for compounds 1 and 3 is shown in Figure 12. At 290 K, the values of $\chi_M T$ are 0.78 (1) and 1.96 cm$^3$ mol$^{-1}$ K (3), which is consistent with the expected values for two Cu$^{II}$ sites in case of 1 (0.75 cm$^3$ mol$^{-1}$ K, for $g = 2.0$) and five Cu$^{II}$ centers for 3 (1.875 cm$^3$ mol$^{-1}$ K, for $g = 2.0$). For both the compounds, a decrease in $\chi_M T$ is observed upon lowering the temperature, with the data entering a plateau at low temperatures. For 1, the value of $\chi_M T$ is very close to zero below 20 K, reaching a minimum value of 0.002 cm$^3$ mol$^{-1}$ K measured at 2 K. In the case of 3, the $\chi_M T$ data also enter a plateau below 20 K, where $\chi_M T = 0.51$ cm$^3$ mol$^{-1}$ K, with a further slight decrease to $\chi_M T = 0.44$ cm$^3$ mol$^{-1}$ K at 2 K. These data indicate overall antiferromagnetic exchange interactions in both compounds.

To determine the size of the exchange, the data were fitted using the program Phi 2.0.$^{41}$ The data for 1 were fitted considering one coupling constant arising from the interaction between the two Cu$^{II}$ ions, $J_1$ (Figure S15, ESI), and also by allowing the $g$-value to vary freely.
small paramagnetic impurity of 1% (\(S = \frac{1}{2}\)) was also included in the fit. The Hamiltonian used for the fitting procedure was:

\[
\hat{H} = -2J_1(\hat{S}_{Cu1}\hat{S}_{Cu2}) + g\mu_B H \sum_{i=1}^{2} \hat{S}_i
\]

and the resulting curve is shown in Figure 12. The strength of the exchange interaction \(J_1\) was found to be \(-42.9\) cm\(^{-1}\), and the \(g\)-value yielded was 2.16. It is difficult to offer a direct comparison of these values with other compounds in the literature, as there are very few examples of two hexa-coordinate Cu\(\text{II}\) centers bridged by one phenoxido, one hydroxido, and one nitrato group. For dinuclear Cu\(\text{II}\) complexes containing two hydroxido bridges, a linear correlation was found between the coupling constant \(J\) and the size of the Cu–O–Cu angle, with larger angles leading to stronger antiferromagnetic exchange.\(^{42}\) However, this exchange interaction was also found to be dependent on the extent of displacement of the hydrogen atom of the hydroxido bridges away from the Cu\(_2\)O\(_2\) plane, with large displacements leading to much weaker antiferromagnetic exchange or even ferromagnetic exchange interactions.\(^{43,44}\) We therefore used B3LYP/6-31+G* calculations (broken symmetry approach, see theoretical methods) to evaluate the magnetic coupling constant \(J\) in 1. The theoretical value of \(J\) (\(-36.9\) cm\(^{-1}\)) is in good agreement with the experimental one (\(-42.9\) cm\(^{-1}\)) and confirms the antiferromagnetic coupling. To investigate the mechanism for the magnetic exchange coupling, the spin density distribution was analyzed. The atomic spin population values on the Cu\(\text{II}\) centers and the donor atoms of the ligands are listed in Table 1.

Figure 12 \(\chi_M T\) versus \(T\) plot for 1 and 3. The solid blue line represents a fit of the data for 1, while the solid black line is the fit of the data for 3 and the red line is a simulation based on the results of DFT calculations (see text for details)
Table 1 Atomic spin densities (e) computed for the high and low spin configuration of 1 at the B3LYP/6-31+G*

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<th>Atom Label</th>
<th>High spin</th>
<th>Low Spin</th>
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<td>Cu1</td>
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<td>Cu2</td>
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<tr>
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<td>0.11</td>
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<td>N2</td>
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<tr>
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</tr>
<tr>
<td>O3</td>
<td>0.02</td>
<td>0.02</td>
</tr>
<tr>
<td>O4</td>
<td>0.03</td>
<td>-0.03</td>
</tr>
</tbody>
</table>

For the high-spin (HS) configuration, the Mulliken spin population data shows that some spin (ca. 0.62 e) is delocalized through the ligands, and the rest (1.38 e) is supported by the CuII centers. The spin density plots corresponding to one of the ‘broken-symmetry’ wave function and the high-spin state for 1 are described in Figure 13, where α and β spin states are denoted by positive (blue) and negative (green) signs, respectively. The broken-symmetry spin population values at the magnetic centers are +0.73 on Cu(1) and −0.68 on Cu(2) and the spin delocalization is considerable (~30% of the spin is delocalized to the ligand framework). The spin on the phenoxide and hydroxide O atoms (O1/O2) is ca. 0.37 e in the HS state and −0.05 e in the broken-symmetry state of 1, consequently, the bridging oxygen atoms mediate the magnetic exchange. Plots of the magnetic orbitals are given in the ESI (Figure S13) showing the contribution of \( d_{x^2-y^2} \) atomic orbitals of the CuII centers along with the \( p_x \) and \( p_y \) orbitals of the bridging O atoms.

**Figure 13** Representation of spin density (contour 0.004 e Å\(^{-3}\)) at the high spin (a) and low spin (b) configurations of 1. Positive spin represented in blue and negative spin in green.
For 3, the fits were initially attempted using a model involving three distinct exchange interactions considering the topology of the complex as a pair of scalene triangles sharing a vertex (Cu3). However, this led to physically unreasonable exchange interactions, with the interaction between Cu1 and Cu3 or Cu2 and Cu3 found to be as strong as the interaction between Cu1 and Cu2, despite the significant differences in distance between the pairs of atoms and in their connectivity. Therefore, the data were fitted considering one exchange interaction $J_1$ between Cu1 and Cu2, (Figure S15 in ESI) and their symmetry equivalent counterparts, adopting an approach used previously for a compound containing a related ligand and the same topology. The $g$ values for the five Cu(II) ions were fixed at 2.16 based on the results for the fitting procedure for compound 1. The Hamiltonian was therefore:

$$\hat{H} = -2J_1(\hat{\mathbf{S}}_{\text{Cu1}} \cdot \hat{\mathbf{S}}_{\text{Cu2}} + \hat{\mathbf{S}}_{\text{Cu1}} \cdot \hat{\mathbf{S}}_{\text{Cu2}}) + g\mu_B H \sum_{i=1}^{5} \hat{\mathbf{S}}_i$$

yielding a value for $J_1 = -54.2$ cm$^{-1}$ (shown as the black curve in Figure 12). Simulations including weaker interactions between Cu1 and Cu2 and the central Cu3 ion did not lead to better agreement with the experimental data. DFT calculations indicated an antiferromagnetic interaction between Cu1 and Cu2 of $-61$ cm$^{-1}$, and ferromagnetic exchange between the Cu3 center and Cu1 and Cu2 centers of $+2.5$ cm$^{-1}$. The most notable aspect of all of the exchange interactions described is that they are relatively weak for coupling between the Cu$^{II}$ centers, and that they are significantly weaker than the interactions mediated by other related ligands.$^{29,10}$

As aforementioned, the B3LYP/6-31+G* calculation of the Cu1–Cu2 interaction in compound 3 reveals an antiferromagnetic coupling with $J = -61.1$ cm$^{-1}$, that is in good agreement with the experimental one ($-54.2$ cm$^{-1}$). To investigate mechanism for the magnetic exchange coupling, the spin density distribution was studied. The atomic spin population values on the Cu1 and Cu2 metal centers and the O1 and O2 bridging donor atoms of the ligands are listed in Table 2.

**Table 2** Atomic spin densities (e) computed for the High and Low spin configuration of compound 3 at the B3LYP/6-31+G*  

<table>
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<th>Atom Label</th>
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<th>Low Spin</th>
</tr>
</thead>
<tbody>
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<td>-0.68</td>
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<td>Cu2</td>
<td>0.69</td>
<td>0.71</td>
</tr>
<tr>
<td>N1</td>
<td>0.13</td>
<td>-0.14</td>
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For the high-spin (HS) configuration, the Mulliken spin population data shows that a noticeable spin (ca. 0.63 e) is delocalized through the ligands, and the rest (1.37 e) is supported by the Cu magnetic centers (see Table 2). The spin density plots of the "broken-symmetry" wave function and the high-spin state for complex 3 are described in Figure 14, where α and β spin states are represented by blue and negative (green) signs, respectively. The broken-symmetry spin population values at the magnetic centers are +0.71 on Cu(1) and −0.68 on Cu(2) and the spin delocalization is considerable (~30% of the spin is delocalized to the ligand framework). The spin on the phenoxide and hydroxide O atoms (O1/O2) are 0.16 and 0.12 e in the HS state and negligible in the broken-symmetry state of complex 3, thus confirming that the bridging oxygen atoms mediate the magnetic exchange.

Finally, we have also studied the magnetic behavior of compound 2 using DFT calculations. We have initially analyzed three distinct exchange interactions (Cu1-Cu2, Cu3-Cu4 and Cu2-Cu4). The magnetic interaction between Cu2 and Cu4 (μ1,3-ClO4 bridging, Cu⋯Cu separation > 7Å) was found to be inexistent (J = 0). In contrast, the B3LYP/6-31+G* calculation of Js between Cu1 and Cu2 and between Cu3 and Cu4 showed the existence of antiferromagnetic coupling (~99.1 cm⁻¹ and ~90.0 cm⁻¹, respectively). Thus the J values
predicted for this compound are slightly larger in absolute value than those observed for compounds 1 or 3. The atomic spin densities and spin density plots for compound 2 are given in the ESI (see Figure S14 and Tables S3 and S4). Unfortunately we were unable to obtain good magnetic data for 2, due to the presence of a paramagnetic impurity that dominated the data in the low temperature region, where a well-isolated $S = 0$ ground state can be expected based on the DFT calculations. The magnetic data for compound 2 together with a simulation based upon the results of the DFT calculations are shown in Figure S16 in ESI.

**Experimental Section**

**Reagents and Materials**

The chemicals used were obtained from the following sources: trifluoroacetic acid (99%) from SRL, India, copper(II) carbonate hydroxide, copper(II) nitrate trihydrate and triethylamine from Merck, India and 2-amino-2-methyl-1,3-propanediol from Alfa Aesar. Copper(II) perchlorate hexahydrate was freshly prepared by treating hydrated copper(II) carbonate (22.12 g, 0.1 mol) with 1:1 aqueous solution of perchloric acid. Similarly copper(II) trifluoroacetate was obtained by reacting copper(II) carbonate (11.06 g, 0.05 mol) with trifluoroacetic acid (11.40 g, 0.1 mol) in water. 2,6-diformyl-4-methylphenol was prepared following a modified literature procedure providing better yield. All the chemicals and solvents used in this work were reagent-grade materials and were used as received without further purification.

**Synthesis**

Ligand $\text{H}_2\text{L1}$ [2,6-bis-{(1,3-dihydroxy-2-methylpropan-2-ylimino)methyl}-4-methylphenol]. The Schiff base ligand $\text{H}_2\text{L1}$ was prepared by reacting 4-methyl-2,6-diformylphenol (0.820 g, 5 mmol) and 2-amino-2-methyl-1,3-propanediol (1.05 g, 10 mmol) under stirring condition for 1 h followed by refluxing for another 2 h. The solvent was removed under vacuum to yield a yellow oily mass, which was characterized by FTIR, NMR spectroscopy and used directly without further purification. FT-IR (cm$^{-1}$, KBr pellet): 3341 (br), 1638 (m), 1219 (w), 1052 (w), 772 (vs). $^1$H NMR (600 MHz, (CD$_3$)$_2$SO, δ ppm): 8.61 (2H, imine–H), 7.55 (2H, aromatic–H), 3.42 (8H, –OCH$_2$), 2.50 (3H, –CH$_3$ substituent on phenyl ring), 1.18 (6H, –CH$_3$). $^{13}$C NMR (150 MHz, (CD$_3$)$_2$SO, δ ppm): 163.58 (imine C), 124.78–160.01 (aromatic C), 66.06 (methylene C attached with O), 63.72 (tertiary C attached to imine N), 22.65 (methyl C attached with benzene ring), 18.63 (methyl C).
[Cu$_2$(μ–H$_2$L1)(μ–OH)(μ$_{1,3}$–NO$_3$)(NO$_3$(OH$_2$))·H$_2$O (1). The as-prepared solution of H$_2$L1 (25 mL), containing 1 mmol of H$_2$L1 (approx. 0.33 g), was diluted to 30 mL by MeOH. To this, a MeOH solution (15 mL) of copper(II) nitrate trihydrate (0.485 g, 2 mmol) was added slowly and the resulting green solution was stirred for ca. 15 min. Following this, a MeOH solution (10 mL) of sodium acetate (0.164 g, 2 mmol) was added during which the solution color changed to dark green and the stirring was continued for 1 h. The solution was filtered, mixed with equal volume of MeCN and kept in air at room temperature for slow evaporation. Green block shaped single crystals suitable for X-ray analysis were obtained after 8 days from this MeOH-MeCN (1:1) solvent mixture. Yield: 0.35 g (54%). Anal. calc. for C$_{17}$H$_{30}$Cu$_2$N$_4$O$_{14}$ (641.55 g mol$^{-1}$): C, 31.83; H, 4.71; N, 8.73. Found: C, 31.91; H, 4.74; N, 8.74. Selected FT-IR bands: (KBr, cm$^{-1}$, vs=very strong, br=broad, s=strong, m=medium, w=weak): 3400 (br), 1635 (vs), 1384 (vs). UV-vis spectra ($\lambda_{\text{max}}$, nm ($\epsilon$, L mol$^{-1}$ cm$^{-1}$)): (MeOH solution) 645 (67), 368 (7600).

[Cu$_4$(μ–H$_4$L1)$_2$(μ–OH)$_2$(μ$_{1,3}$–ClO$_4$)(OH$_2$)$_2$](ClO$_4$)$_3$·2H$_2$O (2). Complex 2 was prepared following a similar procedure as described above for 1 using Cu(ClO$_4$)$_2$·6H$_2$O (0.74 g, 2 mmol) instead of Cu(NO$_3$)$_2$·3H$_2$O. Green block shaped crystals suitable for X-ray analysis were obtained after 1 week. Yield: 0.494 g (69%). The elemental analysis data confirms the molecular formula [Cu$_4$(μ–H$_4$L1)$_2$(μ–OH)$_2$(μ$_{1,3}$–ClO$_4$)(OH$_2$)$_2$](ClO$_4$)$_3$·2H$_2$O. Anal. calc. for C$_{34}$H$_{60}$Cl$_4$Cu$_4$N$_4$O$_{32}$ (1432.85 g mol$^{-1}$): C, 28.50; H, 4.22; N, 3.91. Found: C, 28.08; H, 4.13; N, 3.80. Selected FT-IR bands: (KBr, cm$^{-1}$, vs=very strong, br=broad, s=strong, m=medium, w=weak): 3338 (br), 1636 (s), 1142 (vs), 1111 (vs), 1088 (vs). UV-vis spectra ($\lambda_{\text{max}}$, nm ($\epsilon$, L mol$^{-1}$ cm$^{-1}$)): (MeOH solution) 666 (129), 374 (10500).

[Cu$_6$(μ–H$_4$L1)$_2$(μ–OH)$_2$(μ$_{1,3}$–O$_2$CCF$_3$)$_2$(O$_2$CCF$_3$)$_2$](CF$_3$COO)$_2$ (3). To a stirred ‘solution of H$_5$L1’ (approx. 0.33 g, 1 mmol) in MeOH (20 mL), a MeOH solution (10 mL) of Cu(CF$_3$COO)$_2$·4H$_2$O (0.904 g, 2.5 mmol) was added. The resulting green solution was stirred for 2 h. Then the solution was filtered and kept for slow evaporation in air. Green block shaped single crystals suitable for X-ray analysis were obtained after 1 week from the MeOH reaction medium. Yield: 0.69 g (81%). Microanalytical data are consistent with the formula [Cu$_6$(μ–H$_4$L1)$_2$(μ–OH)$_2$(μ$_{1,3}$–O$_2$CCF$_3$)$_2$(O$_2$CCF$_3$)$_2$](CF$_3$COO)$_2$. Anal. calc. for C$_{46}$H$_{52}$Cu$_5$F$_{18}$N$_4$O$_{24}$ (1704.67 g mol$^{-1}$) C, 32.41; H, 3.07; N, 3.29. Found: C, 32.47; H, 3.10; N, 3.31. Selected FT-IR bands: (KBr, cm$^{-1}$, vs=very strong, br=broad, s=strong, m=medium, w=weak) 3348 (br), 1670 (vs), 1644 (s) 1431 (m), 1329 (m). UV-vis spectra ($\lambda_{\text{max}}$, nm ($\epsilon$, L mol$^{-1}$ cm$^{-1}$)): (MeOH solution) 678 (104), 371 (13700).
Caution! Complexes of metal ions containing organic ligands along with perchlorate counter anions are explosive in nature. Though we have not faced any problem while dealing with the title compounds, but it is safe to prepare small amount of sample and handle it with care.

Physical Measurements
Elemental analyses (C, H and N) of the compounds were performed with a PerkinElmer model 240C elemental analyzer. A Shimadzu UV 3100 UV–vis–NIR spectrophotometer and a PerkinElmer RX1 spectrometer were used to record the solution electronic absorption spectra and FTIR spectra respectively. The purity of the powder compounds was determined by powder X-ray diffraction (PXRD) patterns using a Bruker AXS X-ray diffractometer (40 kV, 20 mA) using Cu-Kα radiation (λ = 1.5418 Å) within 5–50° (2θ) angular range and a fixed-time counting of 4s at 25 °C. A Bruker esquire 3000 plus mass spectrometer was employed to collect the electrospray ionization (ESI) high resolution mass spectra of the three compounds. MALDI-TOF MS studies were performed with the DHBH (2,5-dihydroxybenzoic acid) matrix by using a Bruker UltrafleXtreme instrument. CHI 1120A electrochemical analyzer was used for cyclic voltammetric measurements. Electrochemical measurements were performed at 25°C in MeOH in one compartment cell with platinum as working electrode, Pt wire as counter electrode and Ag/AgCl as reference electrode. Electron paramagnetic resonance (EPR) spectra were recorded at 9.13 GHz (X-band) in continuous wave mode with a Bruker ELEXSYS 580 X-band EPR spectrometer equipped with a standard accessory for room temperature operation (298 K).

Magnetic Measurements
Direct current (dc) magnetic measurements were performed on polycrystalline samples of compounds 1 and 3 constrained in eicosane, using a Quantum Design SQUID magnetometer equipped with a 5 T magnet. The dc measurements were carried out in the temperature range 290 – 2.0 K under applied fields of 1 T (1) and 0.1 T (2 and 3). Data were corrected for the diamagnetism of the compounds and for the diamagnetic contributions of the sample holder and eicosane through measurements.

Crystal Data Collection and Refinement
Single crystal X-ray diffraction data for 1- 3 were collected on a Bruker SMART APEX-II CCD X-ray diffractometer furnished with a graphite-monochromated Mo Kα (λ = 0.71073 Å) radiation by the ω scan (width of 0.3° frame⁻¹) method at 293 K with a scan rate of 4 s per frame. SAINT and XPREP software were used for data processing and space group
determination. Direct method of SHELXS-2014\textsuperscript{47} were used to solve the structure and then refined with full-matrix least squares using the SHELXL (2014/7)\textsuperscript{48} program package included into WINGX system Version 2014.1.\textsuperscript{49} Data were corrected for Lorentz and polarization effects; an empirical absorption correction was applied using the SADABS.\textsuperscript{50} The locations of the heaviest atoms (Cu) were determined easily. The O, N, and C atoms were subsequently determined from the difference Fourier maps. These atoms are refined anisotropically. The H atoms on the bridging hydroxide were found in the difference Fourier maps and refined isotropically. The remaining H atoms were incorporated at calculated positions and refined with fixed geometry and riding thermal parameters with respect to their carrier atoms. Crystallographic diagrams were presented using DIAMOND software.\textsuperscript{51} A summary of the crystal data and relevant refinement parameters is summarized in Table 3. Short distance of 0.96 Å between H14C and H16D is due to the disorder of C16. Crystallographic data have been deposited with the Cambridge Crystallographic Data Centre as supplementary publications CCDC-1828020, 1828021, 1828022. These data can also be obtained free of cost at www.ccdc.cam.ac.uk/conts/retrieving.html (or from the Cambridge Crystallographic Data Centre).

Table 3 Crystal data and structure refinement details for 1 to 3

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<td>C$<em>{17}$H$</em>{30}$Cu$_2$N$<em>4$O$</em>{14}$</td>
<td>C$<em>{34}$H$</em>{62}$Cl$_2$Cu$_4$N$<em>4$O$</em>{32}$</td>
<td>C$<em>{40}$H$</em>{52}$Cu$<em>5$F$</em>{18}$N$<em>4$O$</em>{24}$</td>
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<td>1432.85</td>
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<td>$P$ $T$</td>
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The magnetic properties of compounds 1, 2 and 3 were calculated using the Gaussian 09 package \(^52\) by means of the density functional theory (DFT) and the broken symmetry approach. \(^53,54,55\) We have used the hybrid B3LYP \(^56\) functional combined with the 6-31+G* basis set, \(^57\) which is a good compromise between the size of the systems and the accuracy of the method. We have utilized the crystallographic coordinates and optimized the positions of the hydrogens to carry out the calculations. Moreover, the theoretical models were simplified using H atoms instead of methyl groups in the ligand.

For the calculations of the pentanuclear complex 3, we evaluated each individual coupling constant (\(J_I\) and \(J_2\)) using the methodology proposed by Alvarez group \(^58,59\) in mutinuclear systems. This methodology comprises the substitution of paramagnetic centers by diamagnetic ones to simplify the calculation of coupling constants to evaluations for magnetically coupled systems. This methodology is accurate and very convenient to estimate coupling constants individually.

Conclusions
We have demonstrated for the first time the use of same Schiff base H\(_5\)L\(_1\) for the synthesis of [Cu\(_2\)], [Cu\(_4\)] and [Cu\(_5\)] complexes. Involvement of NO\(_3^-\), ClO\(_4^-\) and F\(_3\)CCO\(_2^-\) groups has been systematically examined for their specificity and anion binding dependent aggregation. The study also recognized the potential of H\(_5\)L\(_1\) in phenoxido form to enforce the formation of the Cu\(^{II}\) aggregates of varying nuclearity and structure. From the imine side arms one of the protonated alcohol end showed coordination whereas the other adjacent one remained dangling to contribute for hydrogen bonding interactions and stabilization of the entities in crystals. Structures in three dimension and bridging potential of NO\(_3^-\), ClO\(_4^-\) and F\(_3\)CCO\(_2^-\)
between CuII centers are crucial in guiding the final outcome of different reactions. Charge density within the products was important to obtain neutral [Cu2], and cationic [Cu4] and [Cu5] compounds. Both ESI-MS (positive) and MALDI-TOF studies indicated the presence of ligand bound \{Cu2(µ-H4L1)\} fragments in solution and thus allowed fine-tuning of aggregation in presence of NO3−, ClO4− and F3CCO2− ions during solid state separation and crystallization. Solution phase characterization was also done by cyclic voltammetry analysis. Only NO3− as ancillary ligand is appropriate for the isolation of 1. Whereas ClO4− ion, as a very rare example, could bridge two \{Cu2(µ-H4L1)\} fragments in 2. We have shown that \{Cu2(µ-H4L1)(µ-OH)\}2+ units, resulting from H4L1− and water derived HO− bridges, and characterized by HRMS analysis, were accountable for the trapping of \{Cu(O2CCF3)2\} units to give 3. The magnetic exchange interactions found for the complexes were weaker than those normally found for related CuII base multinuclear complexes. The coupling is determined by the geometry in the plane of the CuII ions, with a strong influence from the proton on the bridging hydroxido ion. As our assemblies show significant hydrogen bonding interactions, it could be that by altering the nature of these supramolecular interactions that we can tune the magnetic interaction.

Electronic Supplementary information
X-ray crystallographic data in CIF format, Chart S1, Scheme S1, Figures S1–S16, Tables S1–S4. CCDC 1828020, 1828021 and 1828022 contain the supplementary crystallographic data in CIF format for complexes 1–3.

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Conflicts of interest
The authors declare no competing financial interest

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duly acknowledged. M.D. is thankful to Mr. Sudipto Khamrui for his help with EPR measurements.

References


Family of [Cu$_2$], [Cu$_4$] and [Cu$_5$] Aggregates: Alteration of Reaction Conditions, Ancillary Bridges and Capping Anions

Manisha Das,$^a$ Gavin A. Craig,$^b$ Daniel Escudero,$^c$ Mark Murrie,$^b$ Antonio Frontera,$^d$ and Debasish Ray*$^{a,a}$

Enforced coordination by NO$_3^-$, ClO$_4^-$ and CF$_3$COO$^-$ groups resulted in [Cu$_2$] (1), [Cu$_4$] (2) and [Cu$_5$] (3) complexes using H$_2$L1.