Immobilization, Trapping, and Anion Exchange of Perrhenate Ion Using Copper-based Tripodal Complexes

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Abstract: We describe a multidentate tripodal ligand in which three pendant arms carrying di(2-picolyl)amine units are linked to the ortho-positions of a tris(o-xylyl) scaffold, providing N(CH₂-o-C₆H₄-CH₂N(CH₂py)₂)₃ (L). Reaction of L with CuCl₂ in the presence of hexafluorophosphate anion afforded blue cubes of [(CuCl)₃L](PF₆)₃·5H₂O (1). Crystallographic studies of 1 revealed that the three symmetry-related arms each coordinate a {Cu^{II}Cl} unit, and two molecules of 1 are connected to one another through a Cu(µ-Cl)₂Cu bridge, extending the molecular structure to form a two-dimensional layer. These 2-D layers pack in an ABCABC... fashion with PF₆⁻ anions located in between. Reaction of 1 with a stoichiometric amount of perrhenate ion afforded blue plates of [(CuCl)₃L](PF₆)(ReO₄)₂·3H₂O (2). Compound 2 has the same lattice structure of 1, but the tricopper unit backbone now traps one ReO₄ anion through Coulombic interactions. In addition, three molecules of ${\bf 2}$ are bridged by a perrhenate ion forming a $\text{Cu}_3(\mu^3\text{-ReO}_4)$ cluster to give a different 2-D structure, displaying a rare tridentate bridging ReO₄ mode. Thus in addition to classic perrhenate trapping through weak Coulombic interactions, 2 represents an exceptional example in which the ReO₄ anion is immobilized in an extended framework through tight covalent interactions. The interlamellar PF₆⁻ anions in 1 can be exchanged with other anions including perrhenate, perchlorate, or periodate. The structural similarity between perrhenate and pertechnetate makes these materials of potential interest for pertechnetate trapping.

Introduction

Systems that trap and immobilize anions have attracted much recent attention.¹⁻⁸ Because anions regulate a variety of physiological processes, they are potential toxins. As examples, perchlorate ion can adversely affect human health by interfering with iodide uptake into the thyroid,⁹⁻¹¹ and chromate (CrO₄²⁻) is toxic, mutagenic, and a human carcinogen.^{12,13} As a result, many oxo-hydroxo anionic forms of metals and p-block elements are listed as U. S. EPA priority pollutants.¹⁴ Among anionic pollutants, pertechnetate, the tetrahedral oxoanion of technetium, is especially noteworthy. Apart from its ability to interfere with physiological processes, technetium is a nuclear fission waste product with significant radioactivity. Because of its long half-life and good environmental mobility, pertechnetate is one of the most dangerous radiation-derived contaminants and a major concern for long-term disposal of radioactive waste.

In part because of such issues, chemists are interested to develop methods and materials for trapping anions. Resins with cationic quaternary ammonium polymer chains and exchangeable counter anions are the standard in commercial anion exchangers, although they have limited thermal and chemical stability. Molecular trapping complexes, in particular ion pair receptors with recognition sites for both cations and anions, are able to bind an ion pair through cooperative interactions between co-bound ions. Ale an ion pair through the general formula [M²⁺_{1-x}M³⁺_x(OH)₂] Aⁿ⁻_{x/n}·mH₂O, where trivalent metal ions substitute for some of the divalent ones, have a net positive charge on the layered framework. These substances comprise an isostructural class of pure inorganic materials capable of exchanging interlamellar counter anions with

other anions of interest. 18-20 Metal-organic frameworks with extended cationic networks and guest anions are also potential candidates for anion exchange and trapping. 6,21-24

In the foregoing examples, the anions are typically bound through weak electrostatic and/or hydrogen bonding interactions. The binding constants of anions in these systems are small, and this problem is especially troubling for the binding of pertechnetate, which is relatively large, of low charge density, and has a small enthalpy of complexation.² As a consequence, it has proved difficult to prepare receptors for TcO₄⁻, and there is a need for new materials that can trap it with high affinity.² Recently, we reported a multidentate tripodal ligand, N(CH₂-o-C₆H₄-CH₂N(CH₂py)₂)₃ (L), which carries three di(2-picolyl)amine (dpa) units linked to the ortho-positions of a tris(o-xylyl) scaffold (Chart 1). The reaction of L with transition metal ions produced trinuclear complexes that effectively coordinate tetrahedral oxoanions, such as phosphate and arsenate, with each of the three metal ions coordinated to an oxygen atom of the central MO₄ⁿ⁻ anion.²⁵ As a result, the tridentate bridging tetraoxoanion is immobilized by the trinuclear metal complex via strong covalent bonds.

Chart 1

We therefore decided to explore the potential of the tripodal ligand L to bind pertechnetate ion, and the present article describes our findings. As a surrogate for the radioactive TcO_4^- ion, we employed the chemically and structurally similar perrhenate anion, specifically with a Cu(II) complex 1. The result is an extended, cationic 2-D structure capably of completely exchanging interlamellar PF_6^- anions with other anions, including perrhenate, perchlorate, and periodate.

Experimental Section

Synthesis of N(CH₂-o-C₆H₄-CH₂N(CH₂py)₂)₃ (L). Ligand L was synthesized following our published method,²⁵ and its identity and purity were confirmed by ¹H and ¹³C NMR spectroscopy and high-resolution electrospray ionization mass spectrometry (ESI-MS). Crystals of L suitable for single crystal X-ray diffraction were obtained from a mixture DCM/hexane (1:5) solution of L. Slow evaporation of this solution at -10 °C afforded large colorless prisms of L in 3 weeks. Detailed characterization of L is available in the Supporting Information.

Synthesis of [(CuCl)₃L](PF₆)₃·5H₂O (1). To a stirred solution of L (30.0 mg, 0.032 mmol) in 10 mL of dichloromethane at room temperature was added 14.5 mg of CuCl₂ (0.106 mmol, 3.3 equiv) dissolved in 5 mL of MeOH. The resulting clear green-blue solution was stirred overnight to give a clear blue solution. The volume of the solution at this stage was about 3 mL due to evaporation of the solvent. To this clear blue solution was added 50.0 mg of solid NH₄(PF₆) (0.30 mmol), which resulted in immediate precipitation. The precipitate was redissolved by addition of 2.0 mL of MeCN, and the

resulting clear solution was diluted with an additional 10 mL of MeOH. Slow evaporation of this solution at room temperature afforded large blue crystalline cubes of **1** in 5 days. The crystals were harvested by filtration and washed with MeOH (50.0 mg, yield 90%). FT-IR data (2% KBr pellet): 3442 (br, s), 2928 (br, m), 1633 (sh), 1611 (s), 1575 (m), 1483 (s), 1447 (s), 1389 (w), 1366 (w), 1305 (m), 1289 (m), 1251 (m), 1211 (m), 1184 (m), 1161 (m), 1128 (w), 1092 (m), 1067 (w), 1054 (m), 1031 (m), 992 (m), 961 (m), 938 (m), 842 (s), 766 (s), 739 (w), 655 (m), 558 (s), 508 (w), 483 (m), 424 (m). Anal. Calcd. for $[(CuCl)_3(C_{60}N_{10}H_{60})](PF_6)_3 \cdot 5H_2O$: C, 41.34; H, 4.05; N, 8.04. Found: C, 41.36; H, 3.97; N, 8.04. [MW = 1743 g/mol]. High resolution ESI-MS for monocation $[(CuCl)_3(C_{60}N_{10}H_{60})(PF_6)_2]^+$: calcd. 1504.1240; found: 1504.1279 (Supporting Information).

Synthesis of [(CuCl)₃**L](PF**₆)(**ReO**₄)₂·3**H**₂**O** (2). A 10.0 mg sample of **1** (0.00573 mmol) was dissolved in 2 mL of MeCN at room temperature. To the resulting clear blue solution was added 5.2 mg of NaReO₄ (0.0189 mmol, 3.3 equiv) dissolved in 5 mL of MeOH. Slow evaporation of the resulting clear solution at room temperature afforded large blue crystalline plates of **2** over 3 days. The crystals were harvested by filtration and washed with MeOH (10.0 mg, yield 91%). FT-IR data (2% KBr pellet): 3450 (br, s), 2932 (br, m), 1699 (m), 1630 (sh), 1610 (s), 1573 (m), 1482 (s), 1445 (s), 1383 (w), 1367 (m), 1302 (m), 1288 (m), 1248 (m), 1210 (m), 1183 (m), 1159 (m), 1127 (w), 1090 (m), 1067 (w), 1053 (m), 1030 (m), 993 (m), 959 (m), 910 (s), 840 (s), 768 (s), 732 (w), 654 (m), 557 (s), 507 (w), 483 (m), 422 (m). Anal. Calcd. for [(CuCl)₃(C₆₀N₁₀H₆₀)]-

(PF₆)(ReO₄)₂·3H₂O: C, 37.58; H, 3.47; N, 7.30; Cu, 9.94; Cl, 5.55. Found: C, 37.38; H, 3.47; N, 7.19; Cu, 9.83; Cl, 5.65. [MW = 1917 g/mol].

X-ray Crystallography. Data sets for complexes $[(Cu^{II}CI)_3L](PF_6)_3 \cdot 5H_2O$ (1), $[(Cu^{II}CI)_3L](PF_6)(ReO_4)_2 \cdot 3H_2O$ (2), and $N(CH_2-o-C_6H_4-CH_2N(CH_2py)_2)_3$ (L) were collected as described below. Single crystals suitable for X-ray analysis were coated with Paratone-N oil, suspended in a small fiber loop, and placed in a cold gaseous nitrogen stream on a Bruker APEX CCD X-ray diffractometer performing ϕ - and ω -scans at 100(2) K. Diffraction intensities were measured using graphite monochromated Mo $K\alpha$ radiation ($\lambda = 0.71073$ Å). Data collection, indexing, initial cell refinements, frame integration, and final cell refinements were accomplished using the program APEX2. Absorption corrections were applied using the SADABS. Scattering factors and anomalous dispersion corrections were taken from the *International Tables for X-ray Crystallography*. The structure was solved by direct methods using SHELXS²⁸ and refined against F^2 on all data by full-matrix least squares with SHELXL-97²⁹ following established refinement strategies. Details of the data quality and a summary of the residual values for the refinements are listed in Table S1.

Complex 1 crystallizes in the trigonal space group $R^{\bar{3}}$ with V = 14858(3) Å³ and Z = 6. There are two kinds of PF₆⁻ counter anions in the X-ray structure. P1 is disorder free; P2 sits on a special position and is disordered over two sites. All heavy atoms were refined anisotropically. In the crystal lattice, there are a number of badly disordered solvent molecules. As a result, the PLATON/SQUEEZE function was used to deal with this problem, and the formula of 1 was determined by a combination of X-ray

crystallography and elemental analysis. Complex 2 crystallizes in the monoclinic space group C2 with $V = 7669.0(10) \text{ Å}^3$ and Z = 4. The very small Flack parameter 0.029(8) indicates that the correct absolute structure of 2 for this crystal was obtained. The [(Cu^{II}Cl)₃L]³⁺ cation is charge-balanced by one PF₆⁻ and two ReO₄⁻ anions. A tridentate bridging ReO₄ anion binds to three Cu atoms, each from a different molecule of 2 through three Cu-O bonds, whereas the other ReO₄ anion is held in the cavity of a tricopper unit backbone by Coulombic and possible anion- π interactions. The PF₆⁻ anion was not disordered. All heavy atoms in the structure of 2 were refined anisotropically. The largest diffraction peak and hole, 3.621 and -2.129 e Å⁻³, are located close to Re1 atom (1.03 and 0.77 Å, respectively), and possibly arise from imperfect absorption corrections as often encountered in heavy-metal atom structures. In the X-ray structure of 2, no solvent molecules were located in the difference electron density maps, but three water molecules were suggested based on elemental analysis (see above). Ligand L crystallizes in the trigonal space group $R^{\overline{3}}$ with V = 32725(7) Å³ and Z = 24. There are two molecules in the asymmetric unit, one on a special position with a crystallographically required C_3 axis passing through the central N atom and the other at a general position. The PLATON/SQUEEZE function was also used to treat solvent distribution disorder in the X-ray structure of L. The identity and purity of L were further confirmed by high-resolution ESI-MS.

Anion Exchange Studies. Anion exchange of **1** with NaReO₄ was achieved by immersing crystals (10.0 mg) in a 1.0 mL of a 0.1 M NaReO₄ methanolic solution at room temperature without stirring. The exchange process was complete in 1 day as

judged by infrared spectroscopy, and blue crystals of **1**-ReO₄ were harvested by filtration and washed with MeOH (5 × 10 mL) (10.3 mg, yield 91%). FT-IR data (2% KBr pellet): 3440 (br, s), 2930 (br, m), 1632 (sh), 1610 (s), 1574 (m), 1482 (s), 1445 (s), 1386 (w), 1368 (w), 1303 (m), 1288 (m), 1251 (m), 1210 (m), 1184 (m), 1159 (m), 1127 (w), 1090 (m), 1067 (w), 1053 (m), 1030 (m), 993 (m), 962 (m), 909 (s), 863 (w), 766 (s), 733 (w), 655 (m), 512 (w), 483 (m), 424 (m). Anal. Calcd. for [(CuCl)₃(C₆₀N₁₀H₆₀)](ReO₄)₃: C, 36.60; H, 3.07; N, 7.11; Cu, 9.68; Cl, 5.40. Found: C, 36.83; H, 3.28; N, 7.07; Cu, 9.33; Cl, 5.90. [MW = 1969 g/mol].

The reversibility of this anion exchange process was studied by exposing crystals of 1-ReO₄ to 1.0 mL of a 0.1 M NaPF₆ methanolic solution. The PF₆⁻ \leftrightarrow ReO₄⁻ anion exchange processes were repeated three times without the loss of anion exchange activity. Anion exchange between 1 and NaClO₄ or NaIO₄ was also attempted under the same conditions. All materials after anion exchange were harvested by filtration, washed with methanol, dried in air, and analyzed by infrared spectroscopy.

Powder X-ray Diffraction Studies. Powder X-ray diffraction (PXRD) patterns were recorded with a PANalytical's X'Pert Pro Materials research diffractometer equipped with a theta (θ) / two theta (2θ) Bragg-Brenano geometry and using nickel-filtered Cu K α radiation ($K\alpha 1 = 1.5406$ Å, $K\alpha 2 = 1.5444$ Å, $K\alpha 2$ / $K\alpha 1 = 0.5$). Crystals of 1 were removed from the methanol solution by pipette and deposited onto a zero-background silicon plate coated with a thin layer of petroleum jelly. The excess solution was removed using a Kimwipe and the crystals were crushed carefully into the jelly. Crystals of 1-ReO₄ and 1 obtained from anion exchange were loaded and analyzed in a similar manner.

Authentic crystalline samples of NH₄Cl, NH₄(PF₆), NaPF₆, and NaReO₄ were also analyzed by PXRD to confirm that no such salts used or generated during synthesis or anion exchange remained on the surface of crystals of 1, 1-ReO₄, or 1 from anion exchange. PXRD data of these samples are reported in Supporting Information.

Results and Discussion

Synthesis and Structure of Complex 1. Recently, we reported the assembly of trimetal-phosphate clusters that are structural analogues of the trinuclear metal active sites of many enzymes involved in phosphate metabolism, using a novel multidentate tripodal ligand N(CH₂-o-C₆H₄-CH₂N(CH₂py)₂)₃ (L).²⁵ The three dpa-based arms of L can each coordinate a metal ion, and the resulting trinuclear metal site is able to capture a phosphate group, a monoprotonated (HPO₄²⁻), or even a diprotonated (H₂PO₄⁻) phosphate group, by forming three metal-oxygen bonds. These results clearly demonstrate the capability of the tris-dpa ligand to coordinate a trimetal-tetraoxoanion unit, with the tetraoxoanion acting as a tridentate bridging group. Because of our interest in trapping pertechnetate ion with extremely tight binding as a means for the regulation and removal of this dangerous and radioactive anionic pollutant, we decided to synthesize new trimetallic species, such as tricopper complexes, and then study their potential to capture pertechnetate ion. We chose copper because of its earth abundance and strong binding affinity for hard oxygen atoms.

Slow evaporation of a dichloromethane/hexane (1:5 in volume) solution of free ligand L at -10 °C afforded large colorless crystals in three weeks. The three ligand arms linked at the ortho-positions of the tris(o-xylyl) scaffold each carry a dpa unit (Figure

S5), and these arms fold to give a cavity suitable for trapping anions. Reaction of L and three equiv of $CuCl_2$ in methanol gave a clear blue solution. Addition of $NH_4(PF_6)$ caused immediate precipitation and recrystallization of the blue precipitate from MeCN/MeOH afforded large blue cubes of 1 in high yield (90%). The tricopper unit in 1 sits on a special position with a crystallographically required C_3 axis passing through the central N atom (N1, see Figure 1). The three symmetry equivalent dpa ligand arms each incorporate one N0 in through three nitrogen atoms with characteristic N1. Single bond lengths (Table 1). Each N2 ion has an additional N3 ligand at a N4 N5 charge balanced by three N5 anions, which were also located in the N5 ray structure.

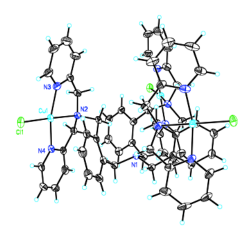


Figure 1. Thermal ellipsoid plot (50% probability) of the tricopper $[(CuCl)_3L]^{3+}$ unit of 1.

In the crystal lattice of 1, two arms of two tricopper ions are connected to one another through a $Cu(\mu\text{-}Cl)_2Cu$ quadrilateral, at a $Cu\cdots Cu$ separation of 3.432 Å (Figure 2B). The bridging of the two Cl^- ions is unsymmetrical. For each dinuclear unit, one chloride ligand is equatorial to its bound Cu^{2+} ion, but resides on the axial position of the other Cu^{2+} ion with a longer Cu(II)–Cl distance of 2.7618(9) Å (Table 1). As a result, the

 Cu^{2+} ion has distorted square pyramidal coordination geometry. These interactions extend the molecular structure to form an infinite 2-D layer. As shown in Figure 2C, the layer consists of an array of hexagons with a diameter of ~20 Å. The central N1 atoms of the bridged tricopper units sit at the vertices of these hexagons, with the two arms linked by $\{Cu(\mu-Cl)_2Cu\}^{2+}$ units forming the edges. Because the layers are constructed exclusively with tricopper $[(CuCl)_3L]^{3+}$ units, the extended structure is positively charged, forming a cationic 2-D metal-organic framework.

Table 1. Bond Distances (Å) to the Cu Ions in 1 and 2

Complex	1		2	
Atom	Cu1	Cu1	Cu2	Cu3
Cu-N	1.982(3)	1.979(8)	1.996(10)	1.995(8)
Cu-N	1.997(3)	1.991(8)	2.014(11)	1.996(9)
Cu-N	2.038(3)	2.049(8)	2.025(8)	2.068(8)
Cu-Cl	2.2536(9)	2.231(3)	2.214(3)	2.226(3)
Cu-Cl for 1 Cu-O for 2	2.7618(9)	2.399(6)	2.270(7)	2.376(7)

These cationic layers pack in an ABCABC... manner, with PF_6^- counter anions located within the interlayer spaces. Figure 3 reveals that the tricopper(II) backbones in each 2-D layer have alternating up and down orientations. Because these layers do not overlap with one another, there are no hexagonal channels in the crystal structure. Because there are only weak electrostatic interactions between PF_6^- anions and the

cationic metal-organic framework, we expected that these guests would be readily exchanged with other anions of interest (vide infra).

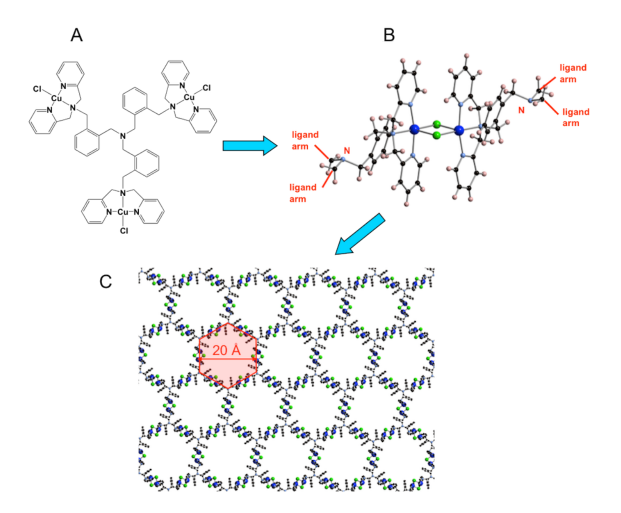


Figure 2. (A) Structure of the tricopper $[(Cu^{II}Cl)_3L]^{3+}$ unit in 1. (B) Ball-and-stick representation of the $\{Cu(\mu-Cl)_2Cu\}^{2+}$ unit that bridges two arms of two molecules in 1. The bridging of the two Cl^- ions is unsymmetrical. (C) The 2-D cationic layer extending through $\{Cu(\mu-Cl)_2Cu\}^{2+}$ interactions (Cu blue; Cl green; N light blue; C black; H pink). The hexagonal pores have diameters of ~ 20 Å.

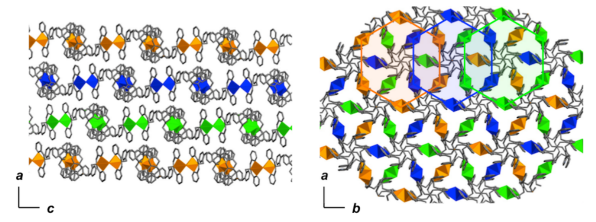


Figure 3. Combined polyhedron and stick representations showing the ABCABC... packing of cationic layers in the 3-D structure of **1** with viewed along b (left) and c (right). The coordination spheres of the Cu^{2+} ions are depicted as polyhedrons, and the ligand backbone is shown as sticks. Hydrogen atoms are omitted for clarity. The layers are indicated in orange, blue, and green to distinguish their packing positions.

Synthesis and Structural Characterization of Complex 2. After analyzing the structure of 1, we were interested to explore its potential to serve as a pertechnetate trapping reagent, anticipating that it might capture a tetrahedral oxoanion in the cavity formed by its tripodal backbone via copper-oxygen bonds, similar to our previous findings.²⁵ Because technetium is radioactive with a long half-life (~10⁶ years), we decided to use perrhenate as a model of pertechnetate for these studies. Table 2 reveals that the geometric features of perrhenate and pertechnetate, including the radii of the metal center,³⁰ metal-oxo bond lengths,³¹ and effective ionic radii of the two tetraoxoanions,^{1,32} are very similar. Because they have the same tetrahedral geometry, similar charge density, and closely related chemical properties, perrhenate is a good model of pertechnetate and should behave in a nearly identical manner in anion trapping studies.

Table 2. Structural Properties and Ionic Radii of Tetrahedral X^{VII}O₄- Anions

	Ionic Radius of X(VII) (Å)	X(VII)–O Bond Length (Å)	X ^{VII} O ₄ Ion Radius (Å)
$\mathrm{TcO_4}^-$	0.56^{a}	1.711^{b}	2.55^e
${ m ReO_4}^-$	0.53^{a}	1.719^{b}	2.60^e
$\mathrm{IO_4}^-$	0.53^{a}	1.726 ^c	2.50^e
C1O ₄ ⁻	0.27^{a}	1.440^d	2.40^e

^aThe ionic radius of X in the tetrahedral ion (X = Tc, Re, I and Cl); ³⁰ ^bTc–O and Re–O bond lengths; ³¹ ^cthe I–O bond length averaged from 21 structures in the Cambridge Crystallographic Data Centre (CCDC); ^dCl–O distance averaged from more than 50 structures in the CCDC; and ^ethe effective $X^{VII}O_4^-$ ion radius. ^{1,32}

Reaction of **1** with three equivalents of NaReO₄ in MeCN/MeOH and subsequent slow evaporation of the resulting clear blue solution gave a high yield (91%) of large blue plates of **2**. These crystals were harvested by filtration and washed with MeOH before analysis. Infrared spectra showed the presence of both ReO₄⁻ (910 cm⁻¹) and PF₆⁻ (840 and 558 cm⁻¹) anions in **2** (Figure 4),³³ an indication that perrhenate is trapped in this material. Crystallographic studies revealed that **2** crystallizes in the monoclinic space group *C2* (Table S1) with an almost identical tricopper unit as that of **1**. The three ligand arms of **2** each bind one Cu²⁺ ion through three nitrogen atoms and each copper has an additional chloride ligand. Cu(II)–N and Cu(II)–Cl bond lengths in **2** are comparable to those in **1** (Table 1). Although the three arms of **2** are not symmetry equivalent in the

crystal structure, they are chemically identical. A thermal ellipsoid plot of the tricopper $[(CuCl)_3L]^{3+}$ unit in **2** is depicted in Figure 5.

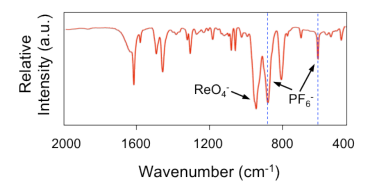


Figure 4. FTIR spectrum of complex **2**. Both ReO_4^- and PF_6^- anions are apparent.

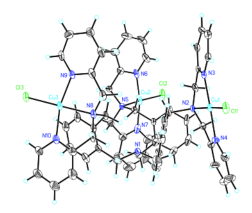


Figure 5. Thermal ellipsoid plot (50% probability) of the tricopper $[(CuCl)_3L]^{3+}$ unit in 2.

In 2, each $[(CuCl)_3L]^{3+}$ unit is charge balanced by two ReO_4^- and one PF_6^- anions, which are all located in the crystal structure without any disorder. As shown in Figure 6, one ReO_4^- anion is located in the cavity of the tricopper backbone. However,

unlike the phosphate derivatives reported previously, ²⁵ for which $H_xPO_4^{(3-x)-}$ (x = 0, 1, or 2) can bridge three metal centers through three metal-oxygen bonds, the long Cu···O(Re) distances (4.580 Å, 4.646 Å, and 4.785 Å) indicate that this ReO₄ anion (Re2) does not covalently bind to copper but is held electrostatically by the [(CuCl)₃L]³⁺ unit through Coulombic and possible anion- π interactions. The use of positively charged receptors for the complexation of anions, such as pertechnetate and perrhenate, is an important research area in supramolecular chemistry.^{2,17} Because pertechnetate and perrhenate are considered soft bases with low surface charge density, receptors having transition metal cations are attractive as trapping agents for these two anions. For example, metal complexes of cyclotriveratrylenes (CTVs) having the general formula [(ML)₃(CTV)]⁶⁺ (M = Ru(II) or Fe(II), L = arene) can bind a ReO_4^- anion in their bowl-shaped molecular cavity.34-36 In our tripodal system, the three Cu-containing arms of 2 fold to form a preorganized cavity that can encapsulate a ReO₄⁻ anion through electrostatic interactions. A space-filling model (Figure 6C) further reveals the size compatibility between the guest tetrahedral ReO₄⁻ anion and the tripodal host molecule.

Of further interest is the ability of the other ReO_4^- anion (Re1) to bridge the ligand arms of three molecules of **2**. As shown in Figure 7B, the Re1 perrhenate ion serves as a tridentate bridging unit, with three of its oxygen atoms each connected to a Cu ion from each of three different molecules. The perrhenate oxygen atom resides on the axial position of each Cu center to produce a distorted square pyramidal geometry, and the resulting Cu-O bond distances are 2.274(7), 2.371(7), and 2.399(6) Å. Because of its low surface charge density, perrhenate is in general not a good σ -donating ligand to transition metal ions. Numerous examples of free perrhenate, monodentate, bidentate, or

bridging bidentate ReO_4^- anions are known in the literature, but a structurally characterized tridentate bridging ReO_4^- unit is extremely rare.^{33,37} The $Cu_3(\mu^3-ReO_4)$ cluster in **2** is one such example of a tridentate bridging ReO_4^- anion. Apart from these interesting structural features, the tridentate binding of a perrhenate ion in the present structure significantly impacts the immobilization of the anion and as such is a model for pertechnetate trapping. As stated above, the capture of these two anions is difficult, and the need to trap them with tight binding units is an important goal in materials science.²

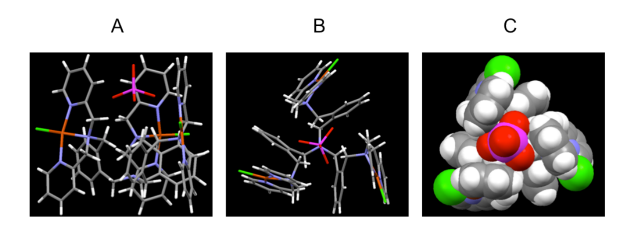


Figure 6. (A) A side view of the stick model showing a trapped ReO_4^- anion in the cavity of the tricopper unit backbone of complex **2**; (B) a top view of the trapped ReO_4^- anion in a stick model; and (C) a top view of the trapped ReO_4^- anion in a space filling model. Color code: Re, purple; O, red; Cu orange; Cl, green; N, blue; C, grey; H, white.

The intermolecular interactions described above extend the structure of **2** to form infinite 2-D layers (Figure 7C). The different solid state structures of **1** and **2** reveal the consequences of using different linkers to bridge the tricopper [(CuCl)₃L]³⁺ unit in the

formation of metal-organic frameworks. Compared to the relatively short $Cu\cdots Cu$ separation of 3.432 Å in 1, the three $Cu\cdots Cu$ vectors in the bridged $Cu_3(\mu^3\text{-ReO}_4)$ cluster of 2 has an average value of 6.274 Å. Figure 8 shows the packing of these cationic layers in the crystal lattice. Two layers having opposite orientations form a bilayer structure through weak electrostatic interactions between perrhenate anions in one layer and the ligand backbone in the other. As a result, the two kinds of perrhenate anions, now designated Re1 and Re2, are both trapped in bilayers with the Re1 anions integrated into the 2-D framework via strong covalent bonds and the Re2 anions encapsulated in the tricopper units cavities through Coulombic interactions. The bilayers are positively charged and they pack in the 3-D structure against anionic layers of PF_6 anions. The structure of 2, with alternating bilayers and PF_6 ions, is presented in Figure 8C.

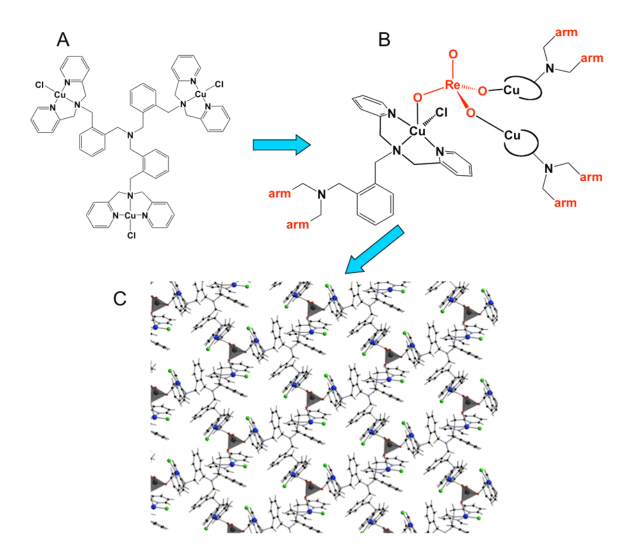


Figure 7. (A) ChemDraw structure of the tricopper $[(Cu^{II}CI)_3L]^{3+}$ unit of **2**. (B) ChemDraw representation of the $Cu_3(\mu^3-ReO_4)$ unit that bridges three arms of three molecules of **2**. The perrhenate oxo ligand sits on the axial position of each Cu ion. (C) Combined polyhedron and ball-and-stick representation showing the 2-D layer extended through $Cu_3(\mu^3-ReO_4)$ interactions $(ReO_4^-$ grey tetrahedron; O red; Cu blue; Cl green; N light blue; C black; H light pink).

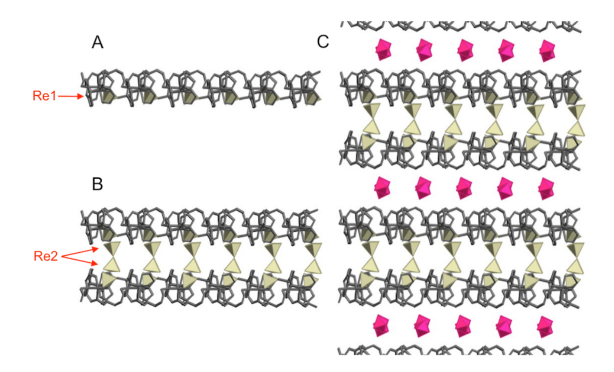


Figure 8. Combined polyhedron and stick representations of the monolayer (A), bilayer (B) and 3-D structure (C) of complex **2**. The ligand backbones are shown in sticks; ReO_4^- anions are shown in grey tetrahedrons; and PF_6^- anions are shown in purple octahedrons. Hydrogen atoms and carbon atoms that are not the main framework skeleton are omitted for clarity.

Anion Exchange Studies. Cationic metal-organic frameworks (MOFs) with positively charged extended networks and loosely bound anionic guests are of potential interest as agents for trapping and removing anionic pollutants. Recently, the anion exchange properties of a few such cationic 2-D MOFs have been described. Similar to layered double hydroxides with anion exchange and trapping features that have been extensively studied over the last two decades, the interlamellar anion guests in cationic 2-D MOFs are also exchangeable with anions, including perrhenate and perchlorate. In these examples, cationic 2-D layers pack against counter anions and solvent molecules that occupy interlayer spaces. The weak interactions between these anions and the

cationic layers facilitate anion exchange reactions for these materials. Because complex 1 also has an extended 2-D MOF structure, formed by $[(CuCl)_3L]^{3+}$ units through intermolecular $Cu(\mu-Cl)_2Cu$ linkers (Figures 2 and 3), with unbound PF_6^- counter anions located in the interlayer spaces, we were interested to investigate its potential for anion exchange reactions.

The replacement of PF₆⁻ anions in 1 by perrhenate was achieved by immersing crystals of 1 in a methanolic solution of NaReO₄ at room temperature (Figure 9). Complex 1 is not soluble in methanol. As shown in Figure S9, the methanol solution is colorless and the crystals maintain their original shape and morphology throughout the anion exchange process. The $PF_6^- \rightarrow ReO_4^-$ anion exchange was complete in one day without stirring when as-isolated crystals of 1 with an average dimension of 0.30 mm were employed. This process could be nicely followed by using infrared spectroscopy. The anion exchange process is fast, although it can take up to a week in previously reported system.⁶ Direct comparison of rate constants for the anion exchange process is difficult because such heterogeneous reactions depend on many experimental factors including crystal size, stir rate, and temperature. Figure 10 shows the FT-IR spectra of 1 and its anion-exchanged product 1-ReO₄. Complex 1 has two intense peaks at 842 and $558~\text{cm}^{-1}$ that are characteristic of octahedral PF_6^- anions. Additional strong peaks in the 1400 and 1700 cm⁻¹ region and the peak at 766 cm⁻¹ are due to the aromatic phenyl and pyridine rings of L. After anion exchange, the two strong PF₆ peaks disappear, and a new intense band at 909 cm⁻¹ appears that is characteristic of the tetrahedral ReO₄⁻ anion.33 Except for this change, all the other peaks from the cationic 2D layers are unaffected. This result confirms complete replacement of PF₆ by ReO₄ anions.

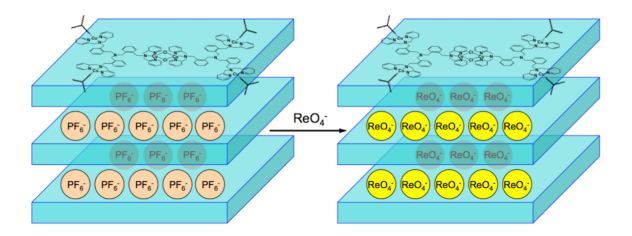


Figure 9. Schematic representation showing the interlamellar PF_6^- anions in complex 1 that are exchanged with ReO_4^- anions.

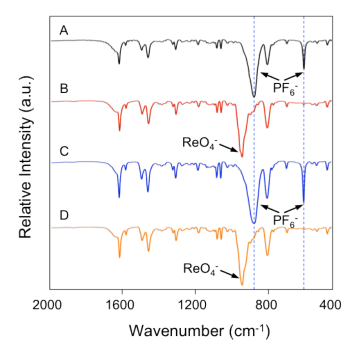


Figure 10. FTIR studies of $PF_6^- \Leftrightarrow ReO_4^-$ anion exchange. (A) initial complex 1 (black); (B) 1-ReO₄ obtained from anion exchange of A and NaReO₄ (red); (C) 1 obtained from anion exchange of B and NaPF₆ (blue); (D) 1-ReO₄ obtained from anion exchange of C and NaReO₄ (orange). The two peaks from PF_6^- anions are highlighted.

The anion exchange was further investigated by powder X-ray diffraction (PXRD) studies. Although we were unable to obtain a single crystal X-ray structure of **1**-ReO₄, the products after anion exchange are crystalline, as shown in Figure 11. Crystals of **1**-ReO₄ gave a different PXRD pattern than crystals of **1**, a finding seen in previous anion exchange studies using cationic 2-D MOFs.^{6,21} The change in the PXRD has a structural origin, for example, modification of the interlayer distances upon anion exchange. Replacement of PF₆⁻ by ReO₄⁻ is also supported by elemental analyses of samples of **1** and **1**-ReO₄ (see Experimental Section).

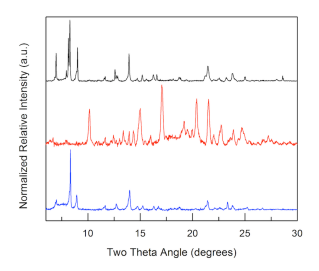


Figure 11. PXRD spectra of (A) initial complex **1** (black); (B) **1**-ReO₄ obtained from anion exchange of A and NaReO₄ (red); (C) **1** obtained from anion exchange of B and NaPF₆ (blue).

The perrhenate in **1-**ReO₄ can be back-exchanged with PF₆⁻ anions. By immersing crystals of **1-**ReO₄ in a methanolic solution of NaPF₆, we observed that all ReO₄⁻ anions were replaced by PF₆⁻ within one day at room temperature. As shown in Figure 10 (from B to C), the ReO₄⁻ peak (909 cm⁻¹) in the infrared spectrum disappears with appearance of the PF₆⁻ peaks (842 and 558 cm⁻¹). Moreover, crystals of **1** obtained from a reaction of **1-**ReO₄ with NaPF₆ behave similarly to those of **1** synthesized for PXRD (Figure 11) and anion exchange studies (Figure 10D). We repeated the PF₆⁻ \rightarrow ReO₄⁻ \rightarrow PF₆⁻ cycle three times without loss of crystal morphology and anion exchange activity. In these processes, all the other IR peaks arising from the cationic framework were unchanged.

The anion exchange properties of 1 were also examined using IO_4^- and CIO_4^- anions under the same experimental conditions. The IO_4^- anion has very similar geometric parameters as TcO_4^- (Table 2) and is another model for studying pertechnetate trapping. After exchanging PF_6^- anions of 1 with $NaIO_4$ in methanol, blue crystals of 1- IO_4 could be isolated. These crystals were washed carefully with methanol and then analyzed by infrared spectroscopy (Figure 12). A strong peak at 847 cm⁻¹ after anion exchange is attributed to IO_4^- anions. 40,41 This assignment is based on literature infrared studies of $NaIO_4$. Although the peak is close to one of the peaks of PF_6^- at 842 cm⁻¹, complete $PF_6^- \to IO_4^-$ exchange was confirmed by the disappearance of the peak at 558 cm⁻¹ arising from the PF_6^- anions. Similarly, the exchange of PF_6^- for CIO_4^- was investigated. Complex 1- CIO_4 displays a new peak at 1090 cm⁻¹ that is characteristic of CIO_4^- anions. 42,43 In both anion exchange processes, all other peaks were unaffected. These results further demonstrate the ability of 1 to exchange its interlamellar anions.

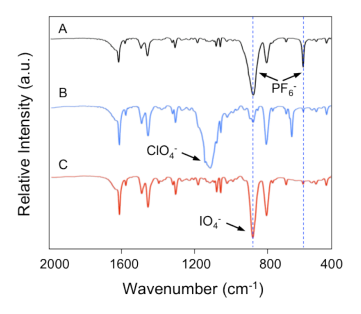


Figure 12. FTIR studies of the $PF_6^- \to XO_4^-$ (X = Cl or I) anion exchange. (A) initial complex 1 (black); (B) 1-ClO₄ obtained from anion exchange of A and NaClO₄ (blue); (C) 1-IO₄ obtained from anion exchange of A and NaIO₄ (red). The two peaks from PF_6^- anions are highlighted.

Summary and Conclusions

We report here the crystal structure of a multidentate tripodal ligand $N(CH_2-o-C_6H_4-CH_2N(CH_2py)_2)_3$ (L) and two metal-organic frameworks with trapped anions, $[(CuCl)_3L](PF_6)_3\cdot 5H_2O$ (1) and $[(CuCl)_3L](PF_6)(ReO_4)_2\cdot 3H_2O$ (2). Compound 1 has a 2-D layer structure formed by tricopper $[(Cu^{II}Cl)_3L]^{3+}$ units extended through intermolecular $Cu(\mu-Cl)_2Cu$ linkers. The PF_6^- counter anions are located within interlayer spaces and can be exchanged with other anions, including perrhenate, perchlorate, and periodate. These results suggest that 1 is an efficient anion-exchanging reagent. Compound 2 also has a layered 2-D structure but its tricopper $[(Cu^{II}Cl)_3L]^{3+}$ units are extended through intermolecular $Cu_3(\mu^3-ReO_4)$ linkages. The tridentate bridging

perrhenate unit represented in the Cu₃(μ³-ReO₄) cluster is extremely rare, probably because of the low surface charge density of the perrhenate ion and its corresponding poor σ-donor properties. The second perrhenate anion in 2 is located in a cavity formed by the tricopper unit backbone and held in place by Coulombic interactions. As a consequence, in addition to perrhenate trapping through electrostatic interactions, 2 represents an exceptional example of tight binding of perrhenate ion through the formation of three covalent Cu–O bonds and the integration of ReO₄⁻ anions into the network skeleton. The immobilization of perrhenate in extended structures, such as metal-organic frameworks, provides a promising lead for pertechnetate trapping studies, and the present investigation may therefore offer some strategic guidance for the regulation of anionic pollutants.

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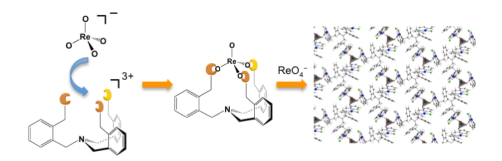
References

- 1. Gloe, K.; Stephan, H.; Grotjahn, M. Chem. Eng. Technol. 2003, 26, 1107-1117.
- Katayev, E. A.; Kolesnikov, G. V.; Sessler, J. L. Chem. Soc. Rev. 2009, 38, 1572-1586.
- 3. Kim, S. K.; Sessler, J. L. Chem. Soc. Rev. **2010**, *39*, 3784-3809.
- 4. Gale, P. A. Chem. Soc. Rev. **2010**, *39*, 3746-3771.
- 5. Ballester, P. Chem. Soc. Rev. 2010, 39, 3810-3830.

- 6. Fei, H. H.; Rogow, D. L.; Oliver, S. R. J. J. Am. Chem. Soc. **2010**, 132, 7202-7209.
- 7. Wang, S. A.; Alekseev, E. V.; Juan, D. W.; Casey, W. H.; Phillips, B. L.; Depmeier, W.; Albrecht-Schmitt, T. E. *Angew. Chem. Int. Ed.* **2010**, *49*, 1057-1060.
- 8. Yu, P.; Wang, S. A.; Alekseev, E. V.; Depmeier, W.; Hobbs, D. T.; Albrecht-Schmitt, T. E.; Phillips, B. L.; Casey, W. H. *Angew. Chem. Int. Ed.* **2010**, *49*, 5975-5977.
- 9. Wolff, J. Pharmacol. Rev. **1998**, *50*, 89-105.
- 10. Urbansky, E. T. Environ. Sci. Pollut. Res. 2002, 9, 187-192.
- 11. Greer, M. A.; Goodman, G.; Pleus, R. C.; Greer, S. E. *Environ. Health Perspect.* **2002**, *110*, 927-937.
- 12. Zayed, A. M.; Terry, N. Plant Soil **2003**, 249, 139-156.
- 13. Costa, M.; Klein, C. B. *Crit. Rev. Toxicol.* **2006**, *36*, 155-163.
- 14. Oliver, S. R. J. Chem. Soc. Rev. **2009**, *38*, 1868-1881.
- 15. Mark, R.; Findley, W. N. *Polym. Eng. Sci.* **1978**, *18*, 6-15.
- 16. Sessler, J. L.; Kim, S. K.; Gross, D. E.; Lee, C. H.; Kim, J. S.; Lynch, V. M. *J. Am. Chem. Soc.* **2008**, *130*, 13162-13166.
- 17. Thuery, P. *Inorg. Chem.* **2009**, *48*, 4497-4513.
- 18. Chibwe, K.; Jones, W. J. Chem. Soc., Chem. Commun. 1989, 926-927.
- 19. Rives, V.; Ulibarri, M. A. Coord. Chem. Rev. **1999**, 181, 61-120.
- 20. Poudret, L.; Prior, T. J.; McIntyre, L. J.; Fogg, A. M. Chem. Mater. 2008, 20, 7447-7453.
- 21. Min, K. S.; Suh, M. P. J. Am. Chem. Soc. **2000**, 122, 6834-6840.
- 22. Hamilton, B. H.; Wagler, T. A.; Espe, M. P.; Ziegler, C. J. *Inorg. Chem.* **2005**, *44*, 4891-4893.
- 23. Xu, G. C.; Ding, Y. J.; Okamura, T. A.; Huang, Y. Q.; Liu, G. X.; Sun, W. Y.; Ueyama, N. *Crystengcomm* **2008**, *10*, 1052-1062.
- 24. Michaelides, A.; Skoulika, S. Cryst. Growth Des. **2009**, *9*, 2039-2042.
- 25. Cao, R.; Müller, P.; Lippard, S. J. J. Am. Chem. Soc. **2010**, 132, 17366-17369.
- 26. Bruker AXS, I. *APEX2 v2009*, Madison, WI, **2009**.
- 27. Sheldrick, G. M. *SADABS*, 2008/1; **2008**.
- 28. Sheldrick, G. M. Acta Cryst. **1990**, A46, 467-473.
- 29. Sheldrick, G. M. *Acta Cryst.* **2008**, *A64*, 112-122.
- 30. Shannon, R. D. Acta Cryst. **1976**, A32, 751-767.
- 31. Krebs, B.; Hasse, K. D. *Acta Cryst.* **1976**, *B32*, 1334-1337.
- 32. Marcus, Y. J. Solution Chem. **1994**, 23, 831-848.
- 33. Chakravorti, M. C. Coord. Chem. Rev. **1990**, 106, 205-225.
- 34. Steed, J. W.; Junk, P. C.; Atwood, J. L.; Barnes, M. J.; Raston, C. L.; Burkhalter, R. S. *J. Am. Chem. Soc.* **1994**, *116*, 10346-10347.
- 35. Holman, K. T.; Halihan, M. M.; Jurisson, S. S.; Atwood, J. L.; Burkhalter, R. S.; Mitchell, A. R.; Steed, J. W. *J. Am. Chem. Soc.* **1996**, *118*, 9567-9576.
- 36. Gawenis, J. A.; Holman, K. T.; Atwood, J. L.; Jurisson, S. S. *Inorg. Chem.* **2002**, *41*, 6028-6031.
- 37. Inglis, R.; Jones, L. F.; Karotsis, G.; Collins, A.; Parsons, S.; Perlepes, S. P.; Wernsdorfer, W.; Brechin, E. K. *Chem. Commun.* **2008**, 5924-5926.

- 38. Zhao, W.; Fan, J.; Okamura, T. A.; Sun, W. Y.; Ueyama, N. *Microporous Mesoporous Mater.* **2005**, *78*, 265-279.
- 39. Du, M.; Zhao, X. J.; Guo, J. H.; Batten, S. R. Chem. Commun. 2005, 4836-4838.
- 40. Gymkowski, T.; Lambert, D. G.; Kimmel, H. S. *J. Inorg. Nucl. Chem.* **1972**, *34*, 1841-1846.
- 41. Levason, W. Coord. Chem. Rev. 1997, 161, 33-79.
- 42. Bünzli, J. C. G.; Mabillard, C. *Inorg. Chem.* **1986**, *25*, 2750-2754.
- 43. Krasnopoler, A.; Stuve, E. M. J. Vac. Sci. Technol., A 1995, 13, 1681-1686.

Table of Contents



The trication unit of a tricopper complex $[(CuCl)_3L](PF_6)_3\cdot 5H_2O$ (1) with $L = N(CH_2-o-C_6H_4-CH_2N(CH_2py)_2)_3$ is an efficient receptor for perrhenate. Reaction of 1 with two perrhenate ions affords $[(CuCl)_3L](PF_6)(ReO_4)_2\cdot 3H_2O$ (2), in which one ReO_4^- anion is trapped in the trication backbone though electrostatic interactions and the other bridges three molecules of 2 in a tridentate manner to form a $Cu_3(\mu^3-ReO_4)$ cluster. This interaction extends the molecular structure of 2 to a 2-D layered material.