

Formation of high nuclearity mixed-valent polyoxovanadates in the presence of copper amine complexes

JENCY THOMAS,^a SANJEEV SHARMA,^a S E LOFLAND,^b K V RAMANUJACHARY^b and A RAMANAN^{a,*}

^aDepartment of Chemistry, Indian Institute of Technology, Delhi, New Delhi 110 016

^bDepartment of Chemistry and Physics, Center for Materials Research and Education, Rowan University, 201 Mullica Hill Road, Glassboro, NJ 08028, USA

e-mail: aramanan@chemistry.iitd.ac.in

Abstract. Two new Müller-type clusters, a one-dimensional solid $\{\text{Cu}(\text{en})_2\}_4[\text{Cl} \subset \text{V}_{15}\text{O}_{36}] \cdot 12\text{H}_2\text{O}$ **1**, and a three-dimensional solid $\{\text{Cu}(\text{pn})_2\}_4[\text{Cl} \subset \text{V}_{18}\text{O}_{42}] \cdot 12\text{H}_2\text{O}$ **2**, have been synthesised by employing identical hydrothermal conditions except varying the nature of organic diamine. **1** crystallised in a chiral space group $P2_12_12_1$ with $a = 12.757(1)$, $b = 18.927(2)$ and $c = 28.590(3)$ Å, and $Z = 4$. **2** crystallised in a tetragonal system with space group $P4/nnc$, $a = 15.113(1)$ and $c = 18.542(3)$ Å, and $Z = 2$. Mixed-valent vanadium ions in structures **1** and **2** have been established both by magnetisation and bond-length bond-valence measurements. Chemistry of formation of high nuclearity polyoxovanadate clusters is discussed.

Keywords. Müller-type clusters; polyoxovanadate; copper amine complex; magnetism, hydrothermal conditions.

1. Introduction

Polyoxovanadate (POV) cluster based solids are attractive inorganic materials owing to their structural versatility and potential applications in the areas of catalysis, magnetism, medicine etc.^{1–3} A range of soluble nano-size vanadium oxide cluster anions formed in the aqueous solution can be stabilised as salts or composite solids by suitable choice of structure directing organic cations or metal complexes.^{4–6} An examination of the crystallographic database (ICSD and CSD) suggests that only a very few high nuclearity clusters, $[\text{V}_x\text{O}_y]^{n-}$ with $x > 15$ have been structurally characterised (figure 1). It was also observed that such clusters are invariably stabilised from solution containing substantial concentration of lower valent vanadium ions (3+ and 4+). Müller and his group assembled several cage structures $[\text{V}_{15}\text{O}_{36}]^{5-}$, $[\text{V}_{16}\text{O}_{38}]^{7-}$, $[\text{V}_{18}\text{O}_{42}]^{12-}$, $[\text{V}_{19}\text{O}_{45}]^{9-}$ and $[\text{V}_{34}\text{O}_{82}]^{10-}$ in which anions are encapsulated inside the cavity of these spherical clusters.⁷ The sizes of the anions (NO_3^- , Cl^- , Br^- , SCN^-) appear to dictate the structure and geometry of these solids. Other anions like formate and acetate invariably derivatise the vanadate group and occur on the surface of the cluster anions. A few more groups are also successful in crystallis-

ing POV clusters wherein $V > 15$.^{8–11} Hydrothermal reactions are popular soft chemistry routes to synthesise such solids. Recently, our group has been involved in rationalising critical chemistry issues involved in controlling the reaction and hence the structure of polyoxometalates under hydrothermal condition, even though such reactions are commonly referred as “black-box” in nature.¹² During our attempts to identify the phases formed in the presence of copper amine complexes, we isolated two mixed-valent microporous solids: $\{\text{Cu}(\text{en})_2\}_4[\text{Cl} \subset \text{V}_{15}\text{O}_{36}] \cdot 12\text{H}_2\text{O}$ (**1**) and a three-dimensional $\{\text{Cu}(\text{pn})_2\}_4$

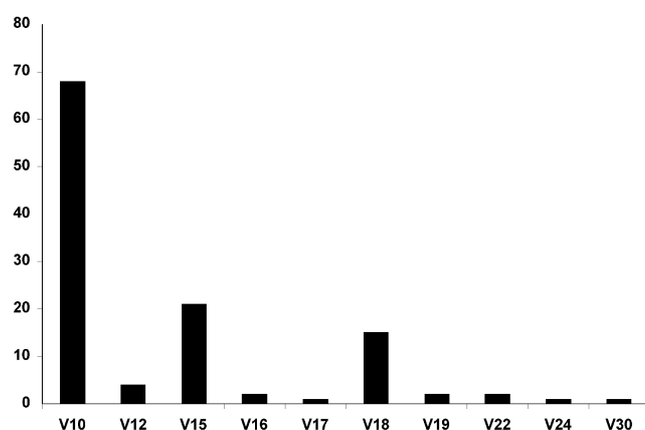


Figure 1. An overview of high nuclearity polyoxovanadates.

Dedicated to Prof J Gopalakrishnan on his 62nd birthday

*For correspondence

Table 1. Single crystal and experimental data for **1** and **2**.

Formula	[Cu(en) ₂] ₄ [Cl ⊂ V ₁₅ O ₃₆] ₄ ·12H ₂ O 1	[Cu(pn) ₂] ₄ [Cl ⊂ V ₁₈ O ₄₂] ₄ ·12H ₂ O 2
Space group	<i>P</i> 2 ₁ 2 ₁ 2 ₁	<i>P</i> 4/ <i>nnc</i>
<i>a</i> (Å)	12.757(1)	15.113(1)
<i>b</i> (Å)	18.927(2)	15.113(1)
<i>c</i> (Å)	28.590(3)	18.542(3)
<i>V</i> (Å ³)	6903.3(1)	4235.5(8)
<i>Z</i>	4	2
Molecular weight (g)	2272.43	2687.61
<i>d</i> _{calc} (g·cm ⁻³)	2.163	2.089
<i>m</i> _{MoKa} (cm ⁻¹)	3.270	2.993
Diffractometer	Bruker Smart Apex CCD	Bruker Smart Apex CCD
Radiation	MoKa	MoKa
<i>T</i> (K)	273(2)	273(2)
Crystal size (mm)	0.52 × 0.23 × 0.13	0.42 × 0.22 × 0.16
Theta range (deg)	2.27–56.60	2.27–51.00
No. of measured reflections	62300	38425
No. of unique reflection	9352	1985
No. of observed reflections (<i>I</i> > 2σ) <i>I</i>	12359	1703
No. of refined parameters	933	141
<i>R</i> ₁ (<i>I</i> > 2σ) <i>I</i>	0.0757	0.0672
<i>WR</i> ₂ (all)	0.1585	0.1534
Min/max Δ <i>r</i> , ς·Å ⁻³	−0.838/1.495	−0.631/0.765
CCDC No.	284863	284864

[Cl ⊂ V₁₈O₄₂]₄·12H₂O (**2**) employing identical reaction conditions except varying the nature of the amine. This paper reports synthesis, structure and magnetic properties of these solids.

2. Synthesis

Vanadium pentoxide (V₂O₅), cupric chloride (CuCl₂·2H₂O), ethylenediamine (*en*) and 1,2-diaminopropane (*pn*) were obtained from Aldrich. All reactions were carried out in teflon-lined stainless steel containers under autogeneous pressure. In a typical synthesis V₂O₅, CuCl₂·2H₂O, organic base and distilled water were taken together in the molar ratio 1:1:5.5:1666. The reactants were sealed into a 15 ml teflon-lined acid digestion reactor and heated at 150°C for 65 h and then cooled to room temperature. The resulting products were washed with water, acetone and allowed to dry in air.

3. Characterisation

FTIR spectra were recorded on KBr pellets using a Nicolet 5DX spectrophotometer. TG/DT analysis was carried out using a Perkin–Elmer TGA7 system on well-ground samples in flowing nitrogen atmosphere with a heating rate of 10°C/min. The magneti-

satation was measured at temperatures ranging from 5 to 300 K, in applied fields of up to 5000Oe with a quantum design physical properties measurement system. Corrections due to the diamagnetism of the sample holder have been applied to the data. Structures of **1** and **2** were determined using single crystal X-ray diffraction. Intensity data collection was carried out on a Bruker Smart Apex CCD diffractometer with a MoKa sealed tube at room temperature. Crystal structures were solved by direct methods using the SHELXTL package. Semi-empirical absorption correction was applied using the SADABS program. The vanadium atoms were first located and then the remaining atoms were deduced from subsequent difference Fourier syntheses. H atoms were located using geometrical constraints. All atoms except H were refined anisotropically. Crystal and experimental data are provided in table 1.

4. Results and discussion

TGA and DTA curves for **1** and **2** are shown in figure 2. The weight loss in the lower temperature region can be attributed to the removal of lattice water molecules. The elimination of organic occurred in a broad step; in both the cases the composition derived from TGA agreed reasonably with those ob-

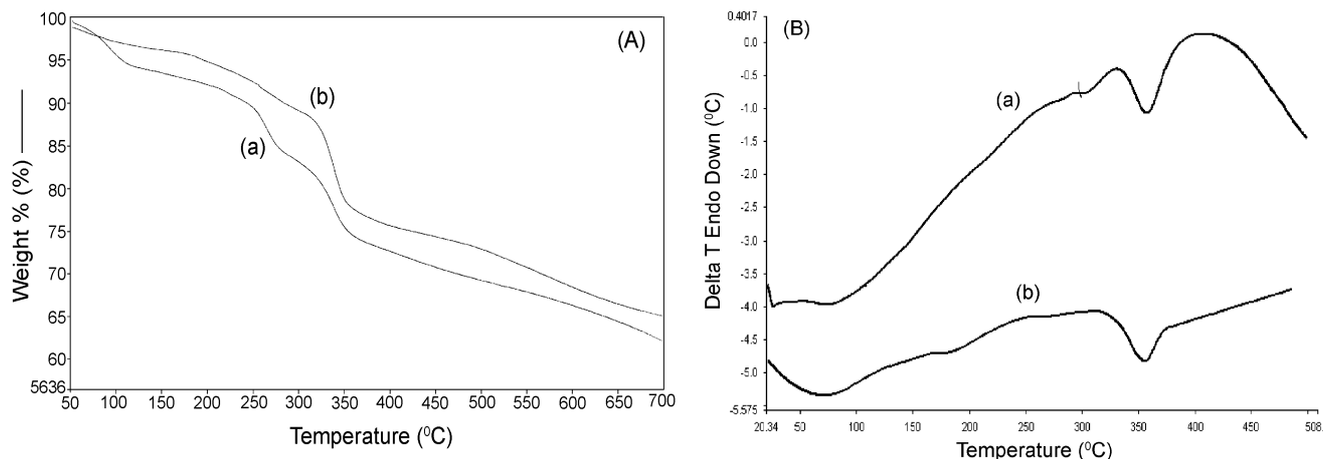


Figure 2. TGA (A) and DTA (B) curves for (a) **1** and (b) **2**.

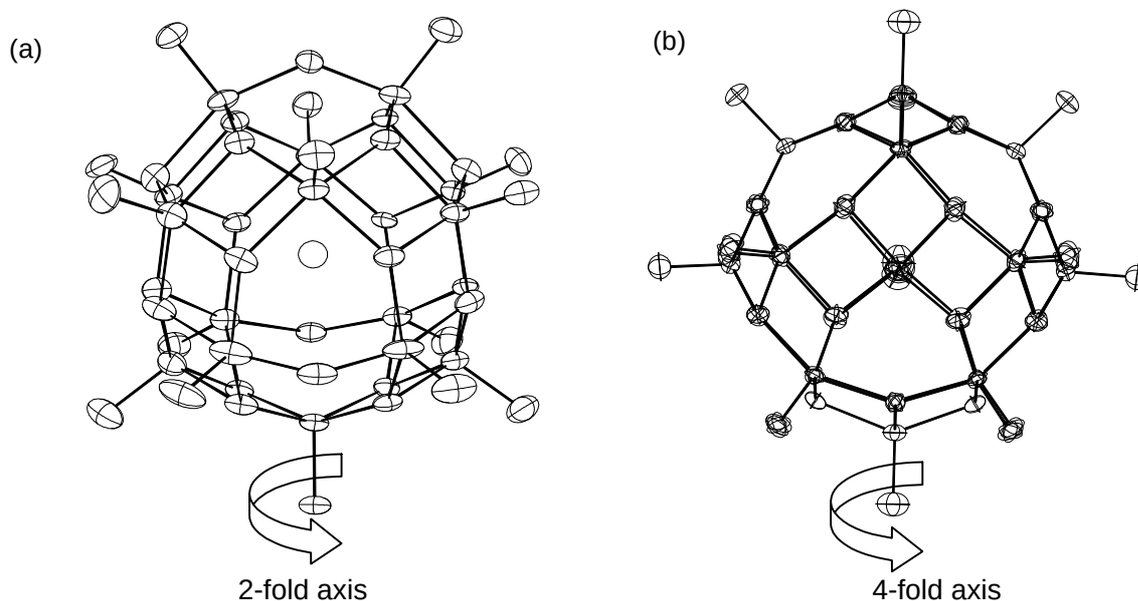


Figure 3. An ORTEP view of (a) **1** and (b) **2**.

tained from single crystal analysis. DTA of the two solids showed an endothermic peak at lower temperature due to water molecules and other at higher temperature due to the removal of organic templates in correspondence with TG analysis. FTIR spectrum of **1** showed strong bands at 965, 888, 844, 721 and 665 cm^{-1} due to V–O symmetric stretching and V–O–V anti-symmetric stretching vibrational modes.¹³ Bands in the 1626–1099 cm^{-1} region are attributed to extending vibrations of *en* groups.¹⁴ FTIR spectrum of **2** exhibited similar characteristic peaks for oxovanadate network. The strong bands at 981, 769, 732 and 660 cm^{-1} are ascribed to V–O–V symmetric

stretching and V–O–V anti-symmetric stretching vibrational modes.¹³ Bands in the range of 1632–1017 cm^{-1} are characteristic of C–C and C–N bonds of *pn* groups.¹⁴

4.1 Crystal structure of **1**

The cluster anion, $[\text{Cl} \subset \text{V}_{15}\text{O}_{36}]^{8-}$ in **1** is closely related to those found in alkali vanadium oxide/halide systems described by Müller *et al.*^{15,16} with idealized C_2 symmetry (figure 3a). The cluster $[\text{Cl} \subset \text{V}_{15}\text{O}_{36}]^{8-}$ is made of a nearly spherical oxovanadium shell built up from fifteen VO_5 distorted square pyramids

that share edges and corners through three $m\bar{2}$ - and eighteen $m\bar{3}$ -oxygen atoms, respectively. The V atoms are arranged at the surface of a sphere with a radius ~ 3.6 Å from the central Cl⁻. The fifteen apical oxygens of the VO₅ square pyramids point out of the cage (figure 3a). In the distorted {VO₅} square pyramid, V–O_b lengths are in the range 1.782(6)–2.036(6) Å, V–O_t lengths 1.605(6)–1.639(6) Å, and bond angles of O–V–O vary from 78.4(2) to 154.7(2)°. The chloride anion adopts a tricosahedral coordination, implying 23 oxygen atoms with the Cl–O distance ~ 3.4 Å, from oxygen atoms of the basal plane of the square pyramids.

Four counter cations, [Cu(en)₂]²⁺, present in the medium, coordinate with V₁₅ cluster anions on one side to form infinite chains (figure 4a) with Cu–O distances of 2.481(1) Å during self-assembly (figure 4b). The 1D chains are further linked via a weaker Cu–O bond ($\sim 2.606(1)$ Å) to form a three-dimensional network (figure 5). In addition, **1** also exhibits hydrogen-bonding interactions between the amine nitrogens and terminal and bridging oxo groups of the cluster along with the water molecules that occur as space fillers. Crystal structure analysis shows that one of the copper ethylenediamine complex rings has rather high thermal parameters. This feature arises probably from some kind of dynamic disorder because of the high volume cell and large number of atoms. Since no special problems for locating atoms were encountered, it is concluded that the structural determination is reasonably correct. Bond valence sums¹⁷ (BVS) of V₁₅ suggest delocalisation of charge over all the vanadium centres (table 2). On the basis of single crystal structure, bond valence sums and thermal analysis, we derived the composition of **1** as {Cu(en)₂}₄[Cl ⊂ V₁₅O₃₆]·12H₂O.

4.2 Crystal structure of **2**

The {V₁₈O₄₂} shell in **2** is essentially similar to Cs₁₂[(H₂O ⊂ V^{IV}₁₈O₄₂)]·9H₂O and related compounds¹⁶

Table 2. Bond valence sums for **1**

V1	4.122	V2	4.520
V3	4.655	V4	4.215
V5	4.683	V6	4.195
V7	4.648	V8	4.164
V9	4.626	V10	4.849
V11	4.722	V12	4.522
V13	4.669	V14	4.762
V15	4.694		
Average sum = 4.54			

with an idealized D_{4d} symmetry (figure 3b) except that Cl⁻ ions are trapped inside the cluster. The cluster anion [Cl ⊂ V₁₈O₄₂]⁸⁻ is composed of eighteen edge-sharing VO₅ square pyramids with an average V–V distance of ~ 3.0 Å. The encapsulated chloride

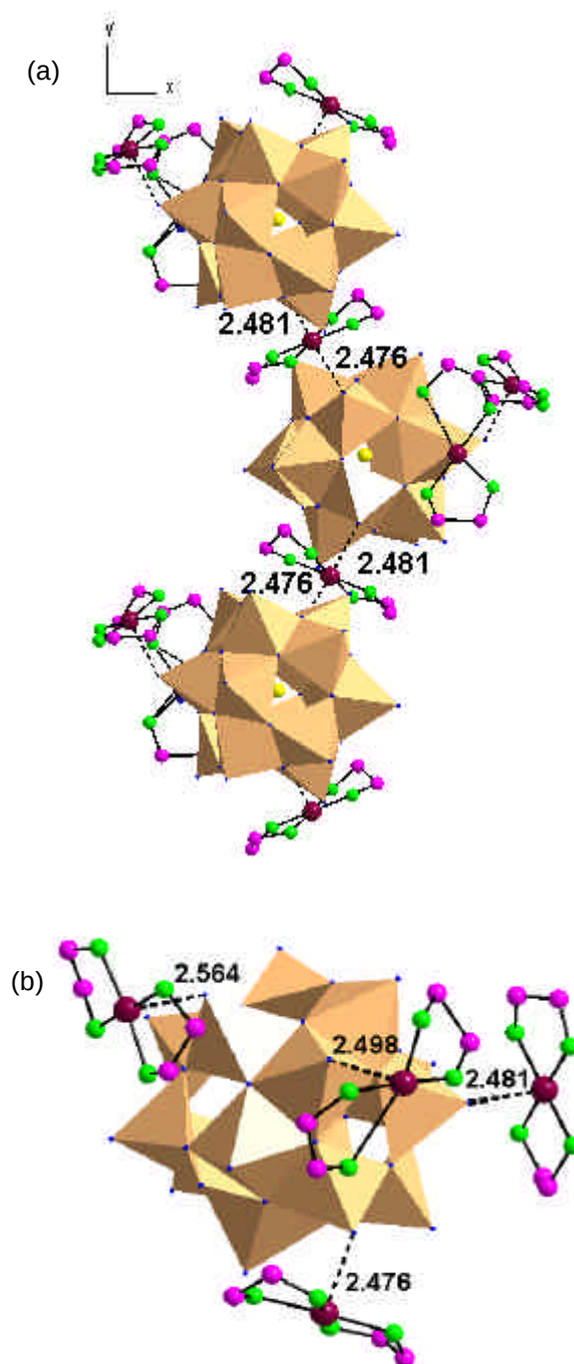


Figure 4. (a) [Cu(en)₂]²⁺ and V₁₅ forming 1D chains along z-axis in **1**. (b) Each V₁₅ cluster in 1D chain is enveloped by four [Cu(en)₂]²⁺ with short Cu–O distance (~ 2.5 Å).

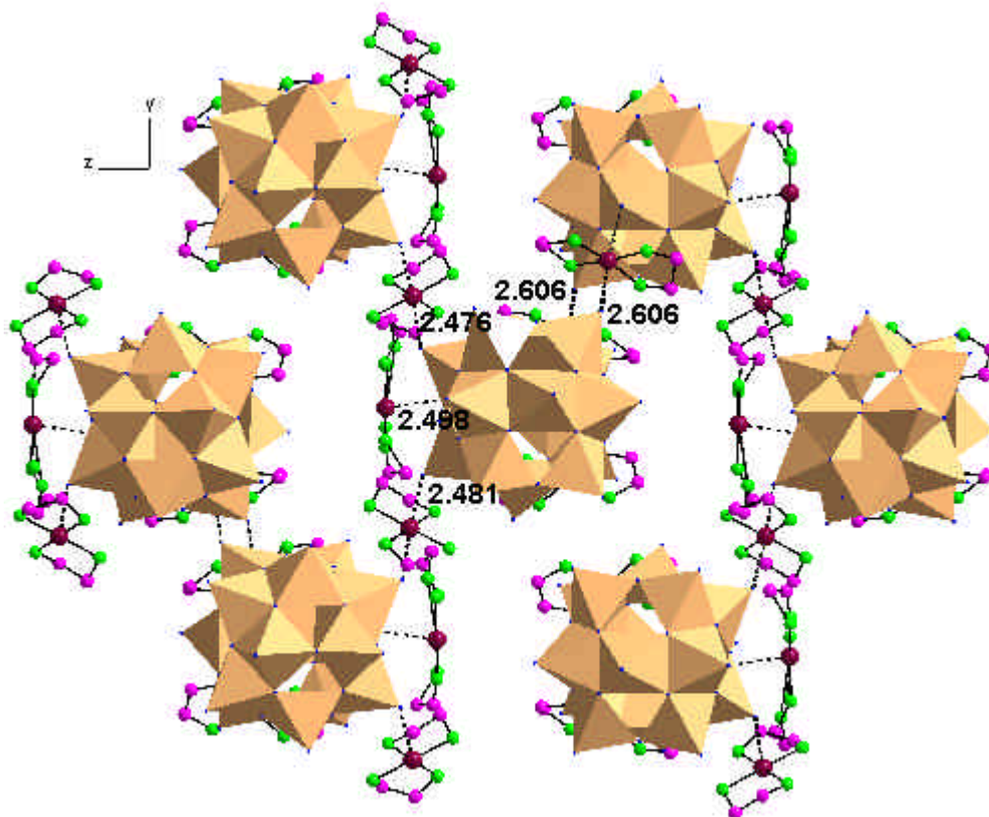


Figure 5. Interaction of 1D chains on the bc -plane through longer Cu–O distances (~ 2.6 Å). (water molecules are not shown, for clarity).

Table 3. Bond valence sums for **2**

V1	4.399
V2	4.498
V3	4.555
Average sum = 4.48	

ion in the shell has no appreciable interaction with the shell framework. The average distance of this chloride ion from the V atoms of the shell is ~ 3.78 Å. All the VO_5 units in **2** have a terminal oxo group and four m_3 -oxygen atoms. Each V_{18} cluster is enveloped by eight $[\text{Cu}(\text{pn})_2]^{2+}$ cations connected through coordinate bonding (figure 6) to form a three-dimensional network with water molecules occupying interstitial spaces (figure 7).

V–O distances and BVS calculations¹⁷ (table 3) indicate that the oxygen atoms in $[\text{Cl} \subset \text{V}_{18}\text{O}_{42}]^{8-}$ are not protonated. **2** exhibits hydrogen-bonding interactions between the amine nitrogens and terminal and bridging oxo groups of the clusters along with water molecules. The cell parameters of **2** seem to be similar to those reported recently in the literature.^{18,19} On

the basis of single crystal structure, bond valence sums and thermal analysis, we derived the composition of **2** as $\{\text{Cu}(\text{pn})_2\}_4[\text{Cl} \subset \text{V}_{18}\text{O}_{42}] \cdot 12\text{H}_2\text{O}$.

4.3 Magnetic properties

Magnetic properties of **1** and **2** were investigated between 300 and 5 K. Figures 8 and 9 show χ^{-1} versus T and χ versus T in the low temperature region. The linear portions of the plots have been fitted to the Curie–Weiss relation yielding the Weiss constants of -11 K and -6 K for the structures **1** and **2** respectively. The negative values of the Weiss constant indicate that the precursor exchange interactions are dominantly anti-ferromagnetic, although no long-range ordering was observed down to 5 K in either of the samples. The effective magnetic moments calculated for both **1** and **2** were ~ 4.4 and 5.8 BM respectively corresponding to ~ 6 and ~ 11 unpaired electrons per formula unit. However, the observed number of unpaired electrons is smaller than expected based on four copper ions and fifty percent of V^{4+} population in the vanadium clusters as estimated

from bond valence sums. It is interesting to note that in contrast to the lower $c.T$ ($\sim 1\text{--}2$ BM) values observed by Müller *et al*²⁰ for mixed-valent V_{18} clusters containing similar vanadium (IV) population, in the present samples, the d electrons seem more localized. Further studies including e.s.r. and neutron diffraction are currently underway to address this issue.

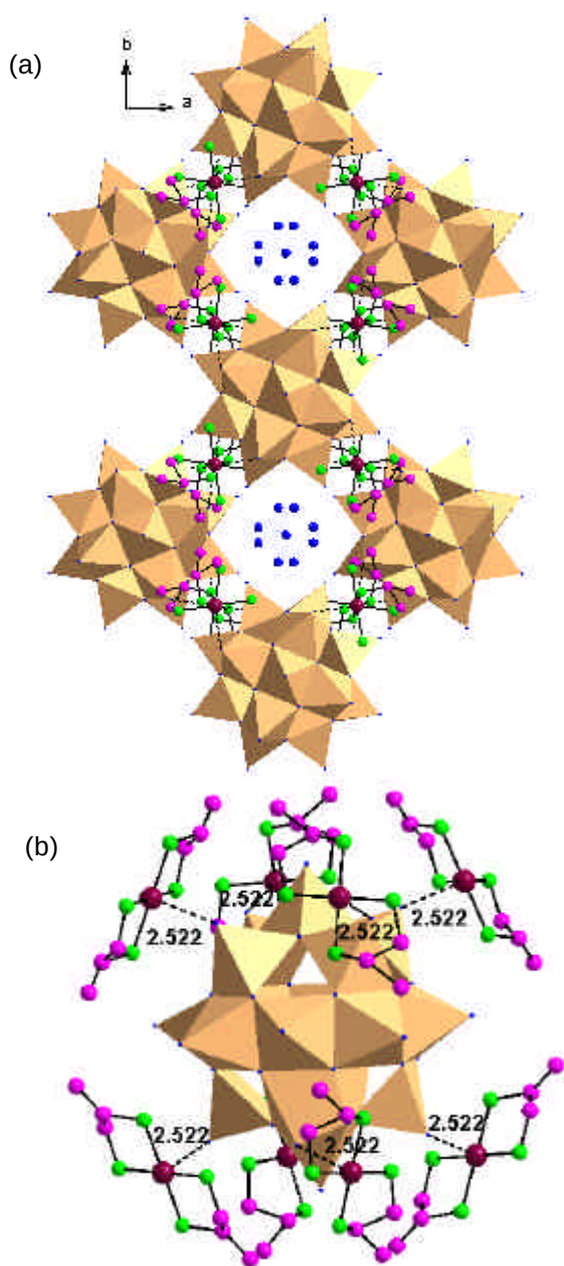


Figure 6. (a) Packing arrangement of three-dimensional network of V_{18} in **2** viewed along the z -axis. Tetragonal cavities show the presence of water molecules. (b) A supramolecular assembly of V_{18} with eight symmetrically coordinating $[Cu(pn)_2]^{2+}$ complexes.

4.4 Chemistry of formation of high nuclearity polyoxovanadates

An examination of all V_{15} and V_{18} clusters reported in the literature suggest that anions or water molecules clearly dictate the formation of this cluster especially when crystallisation is carried out from solutions containing substantial amounts of lower valent vanadium.^{16,20} There exist no example of V_{15} clusters wherein all the vanadium occurs in the +4 oxidation state. V_{18} clusters do occur with all vanadium in the +4 oxidation state but only in the presence of the alkali metals, Rb, Cs and K, as counter cations.²⁰ In fact, neither of these clusters is stabilised in the presence of sodium ions; decavanadates and Keggin clusters dominate reactions containing hydrated sodium ions. In the presence of larger counter cations such as tetra alkyl ammonium or metal complexes, their charge and size decides the overall charge on the anions and the clusters are invariably mixed-valent.^{7,21} Copper complexes present in the solution readily coordinate with these clusters and the supramolecular assemblies thus formed dictate the crystal packing of the final solid. While $[Cu(en)_2]^{2+}$ is known only in **1** (V_{15} cluster) as a counter cation, $[Cu(pn)_2]^{2+}$ occurred as a counter cation in many solids: $\{Cu(pn)_2\}_3[Cl \subset V_{15}O_{36}] \cdot 2.5H_2O$,²² $\{[Cu(pn)_2]_7\{(H_2O)_2 \subset V_{16}O_{38}\} \cdot 4H_2O$,²³ $\{Cu(pn)_2\}[Cl \subset V_{16}O_{38}] \cdot 3 \cdot 5H_2O$,²⁴ $\{Cu(pn)_2\}_4[(H_2O) \subset V_{18}O_{42}] \cdot 8H_2O$,¹⁸ $\{Cu(pn)_2\}_4[Cl \subset H_5V_{18}O_{42}] \cdot 8H_2O$,¹⁹ and $\{Cu(pn)_2\}_4[Cl \subset V_{18}O_{42}] \cdot 12H_2O$. In addition, the way different copper complexes coordinate with the anion, decides the overall symmetry of the unit cell. The high symmetry reflected in the crystal packing of **2** probably suggests that nucleation is induced by the symmetry of the supramolecular assembly.

Another interesting observation that emerges out of this work is that whenever crystallisation is carried out under slow evaporation conditions, bi-capped Keggin clusters are favoured if the solution does not contain sufficient concentration of lower-valent vanadium.^{5,13,25} Formation of V_{15} and V_{18} under our reaction conditions was due to the reduction of vanadium species by the organic amines such as *en* and *pn* under hydrothermal condition. We intentionally did not employ any vanadium metal or lower valent vanadium precursors in our reactions. On the basis of the present work and past experience,^{5,6,25–27} we infer that in many cases pH and concentration of lower-valent vanadium are the primary factors responsible for the occurrence of a particular POV cluster-based solid; in many cases the organic cations

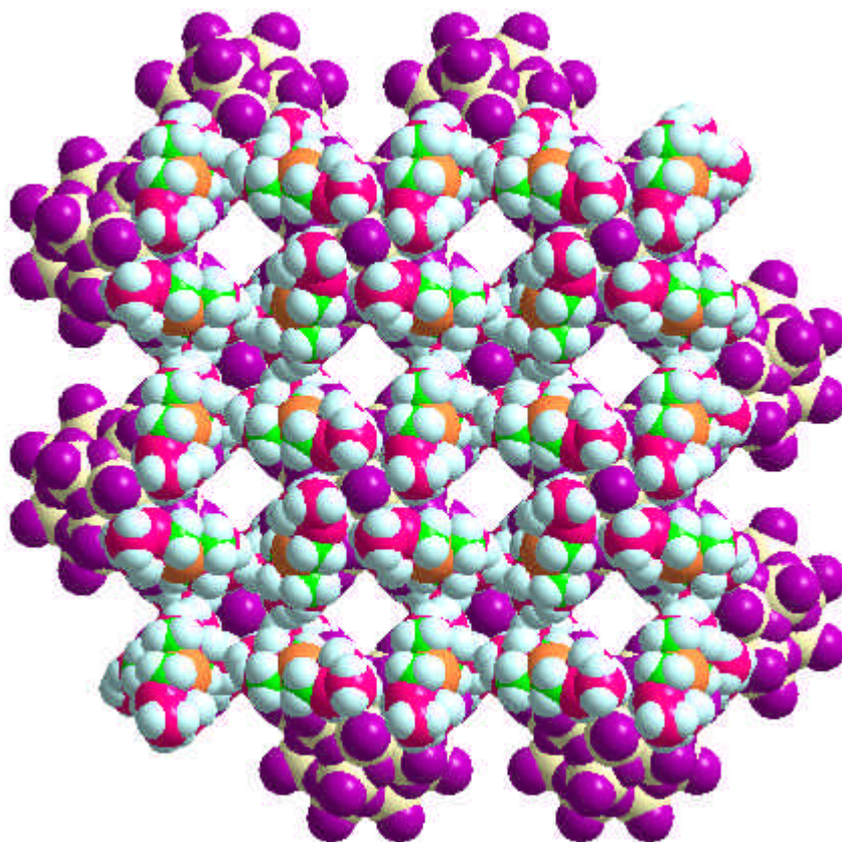


Figure 7. CPK representation of **2** (water molecules are not shown, for clarity).

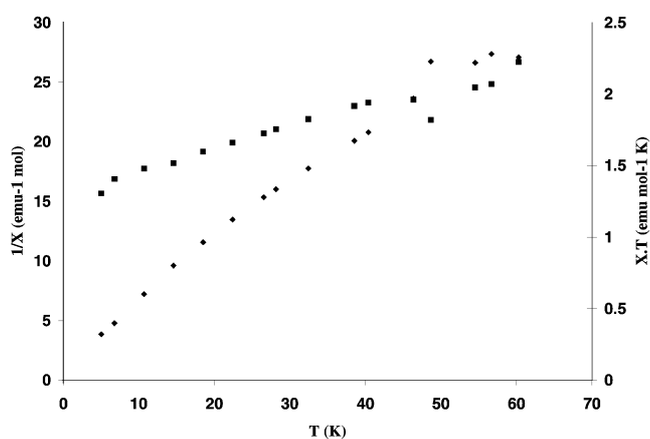


Figure 8. c^{-1} vs T (◆) and $c.T$ vs T (■) of **1** in the 5–60 K temperature range.

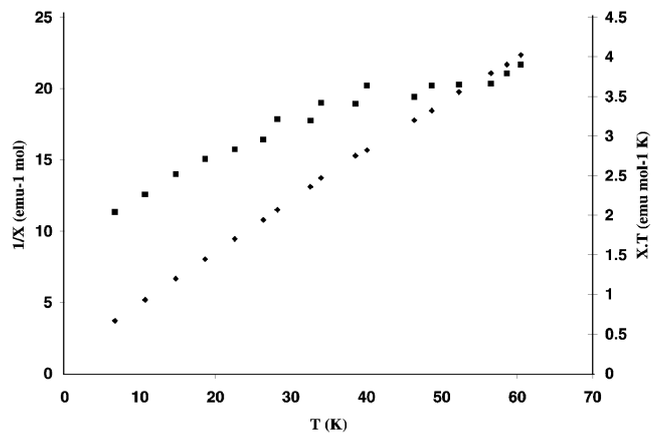


Figure 9. c^{-1} vs T (◆) and $c.T$ vs T (■) of **2** in the 5–60 K temperature range.

occur as spectator counter-cations. Hydrothermal conditions assist the reduction of vanadates by organic amines to the lower-valent state. In addition, it provides better solubility of reactants and formation of good crystals.

5. Conclusions

High nuclearity POV-based solids reported here crystallise from aqueous solution containing lower-valent vanadium species but are essentially driven

by chloride ions. The size and nature of the counter cations play a secondary role in stabilising such clusters. Hydrothermal medium is effective in solubilising reactants that are otherwise insoluble in aqueous solution and promoting reduction of vanadium species with organic amines. The coordinating ability of the counter cation with a particular POV cluster dictates the crystal packing of these hybrid solids. Hydrothermal chemistry will be more attractive for materials syntheses if a rational strategy is evolved rather than employing a cocktail of arbitrary reactants and reaction conditions.

Acknowledgments

JT acknowledges CSIR for a fellowship. AR acknowledges DST for financial support and DST-IRHPA and DST-FIST for the powder and single crystal X-ray diffraction facility to Department of Chemistry, IIT, Delhi. AR, KVR and SEL express their respect and admiration to Prof J Gopalakrishnan whose contributions have enriched our understanding of the fascinating chemistry of inorganic solids.

References

1. Coronado E and Gomez-Garcia C J 1998 *Chem. Rev.* **98** 273
2. Pope M T 1983 *Heteropoly and isopoly oxometalates* (Berlin: Springer)
3. Pope M T and Müller A 1994 In *Polyoxometalates: From platonic solids to anti-retroviral activity* (Dordrecht: Kluwer)
4. Müller A, Peters F, Pope M T and Gatteschi D 1998 *Chem. Rev.* **98** 239
5. Duraisami T, Ojha N, Ramanan A and Vittal J J 1999 *Chem. Mater.* **11** 2339
6. Duraisami T, Ramanan A and Vittal J J 2001 *Cryst. Eng.* **3** 237
7. Müller A, Rohlfing R, Krickemeyer E and Bogge H 1993 *Angew. Chem., Int. Ed. Engl.* **32** 909
8. Suber L, Bonamico M and Fares V 1997 *Inorg. Chem.* **36** 2030
9. Khan M I, Ayesh S, Doedens R J, Yuc M and O'Connor C J 2005 *Chem. Commun.* 4658
10. Hayashi Y, Fukuyama K, Takatera T and Uehara A 2000 *Chem. Lett.* 770
11. Hayashi Y, Miyakoshi N, Shinguchi T and Uehara A 2001 *Chem. Lett.* 170
12. Pavani K and Ramanan A 2005 *Eur. J. Inorg. Chem.* **15** 3080
13. Sharma S, Ramanan A and Vittal J J 2001 *Prod. Indian Acad. Sci. (Chem. Sci.)* **113** 621
14. Nakamoto K 1978 *Infrared and Raman spectra of inorganic and coordination compounds* (New York: John Wiley & Sons)
15. Müller A, Krickemeyer E, Penk M, Walberg H J and Bogge H 1987 *Angew. Chem. Int. Ed. Engl.* **26** 1045
16. Müller A, Krickemeyer E, Penk M, Rohlfing R, Armatage A and Bogge H 1991 *Angew. Chem. Int. Ed. Engl.* **30** 1674
17. Brown I D and Altermatt D 1985 *Acta Crystallogr.* **B41** 244
18. Lin B Z and Liu S X 2002 *Gaodeng Xuexiao Huaxue Xuebao* **23** 535
19. Cui X B, Zheng S T, Ding L, Ding H and Yang G Y 2003 *Jiegou Huaxue* **22** 491
20. Müller A, Sessoli R, Krickemeyer E, Bogge H, Meyer J, Gatteschi D, Pardi L, Westphal J, Hove-meier K, Rohlfing R, Doring J, Hellweg F, Beugholt C and Schmidtman M 1997 *Inorg. Chem.* **36** 5239
21. Khan M I 2000 *J. Solid State Chem.* **152** 105
22. DeBord J R D, Haushalter R C, Meyer L M, Rose D J, Zapf P and Zubieta J 1997 *J. Inorg. Chim. Acta* **256** 165
23. Lin B and Liu S 2002 *Chem. Commun.* 2126
24. Cui X, Zheng S, Sun Y and Yang G 2004 *Chem. Res. Chin. Univ.* **20** 266
25. Pavani K, Upreti S and Ramanan A 2006 *J. Chem. Sci.* (accepted)
26. Sharma S, Ramanan A and Jansen M 2004 *Solid State Ionics* **170** 93
27. Sharma S, Ramanan A, Zavalij P Y and Whittingham M S 2002 *Cryst. Eng. Comm.* **4** 601