New Routes in Polyoxometalate Chemistry: From Keplerates to Chemistry under Confined Conditions

– Cumulative Dissertation –

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Chapter 1

Introduction

1.1 Synopsis

The emergence of the term “nanotechnology” and its increasing use in the scientific and popular scientific literature reflects the expanding interest in the ability to gain control over the organisation of material, in order to fabricate and exploit entities with dimensions of less than 100 nm. This fascinating field opens up many new exciting possibilities for example in materials science, biomolecular transport systems, and bio-sensor technology.

A fundamental question is: how can the appropriate material be organised in the desired arrangements to provide such nano-sized entities? One strategy for achieving such entities reproducibly is the “top-down” approach. In this approach usually the desired nanostructure is designed starting from a macroscopic structure. A second strategy is a “bottom-up” approach. In this approach the nanostructures are generated from a library of building blocks, which might be a group of atoms, molecules, ions or an iterative structural moiety. It is inevitably a zone where chemists can exhibit extreme creative prowess.

Polyoxometalate chemistry, the chemistry of the inorganic metal-oxygen cluster anions, mostly based on Mo, W or V, uses the advantages of self assembly but based on covalent linking, to synthesise a variety of nano-sized entities which are based on the linking of transferable building blocks, under "one-pot" conditions. This type of chemistry has already yielded a multitude of compounds containing polyanions which display a fascinating degree of structural and functional variety comparable to that of proteins [1]. Optimal conditions for linking of fragments leading to a large variety of structures are:
1. the potential of the system to generate a versatile library of linkable units,
2. the ability to generate groups (intermediates) with high free enthalpy to drive polymerization or growth processes, e.g. based on formation of H$_2$O,
3. the possibility for easy structural changes in the building units and blocks, the ability to include hetero elements in the fragments,
4. the possibility to form larger groups which can be linked in different ways,
5. the ability to control the structure-forming processes by templates,
6. the ability to generate structural defects in reaction intermediates (e.g. leading to lacunary structures) e.g. by removing building blocks from (large) intermediates due to the presence of appropriate reactants,
7. the ability to localize and delocalize electrons in different ways in order to gain versatility,
8. the ability to control and vary the charge of building parts (e.g. by protonation, electron transfer reactions, or substitution) and to limit growth by the abundance of appropriate terminal ligands,
9. the possibility of generating fragments with energetically low-lying unoccupied molecular orbitals [2].

These conditions can be optimally fulfilled in polyoxometalate systems which possess the relevant variety of structural and electronic versatility. It is not only possible to perform a new type of chemistry with the clusters in aqueous solution but also to dissolve them in organic solvents, e.g. after encapsulating them with suitable surfactant molecules, with the option of forming monolayers, thin films, liquid crystals and hybrid materials [3]. Furthermore, it is possible to study their aggregation behavior in solution leading to the formation of novel vesicles [4].

1.2 Basic principles and highlights of the field

The basic structural principle for polyoxotungstates and molybdates is the same, since the structures are governed by the principle that each metal atom occupies an \( \{MO_x\} \) coordination
polyhedron, in which the metal atom is displaced, as a result of M-O π bonding, toward those polyhedral vertices that form the surface of the structure. However, a more detailed view of this fascinating area of chemistry shows striking differences for these compound types also with respect to the very large cluster systems [5].

1.2.1 Polyoxotungstates
The structural features of large polyoxotungstate clusters can be visualized in terms of subunits based on lacunary fragments of the Keggin anion (with the classical archetypal \( \{W_3O_{13}\} \) units) including its isomers [6].

**Keggin-isomers and trilacunary structures**
The structure of the classical Keggin anion has overall \( T_d \) symmetry and is based on a central \( XO_4 \) tetrahedron surrounded by twelve \( \{WO_6\} \) octahedra arranged in four groups of three edge-shared octahedra, \( \{W_3O_{13}\} \). These \( \{W_3O_{13}\} \) groups are linked by sharing corners to each other and to the central \( XO_4 \) tetrahedron (Figure 1.1). The Keggin ion can adopt up to five isomers (\( \alpha-\varepsilon \)) and these isomers are related to each other by a rotation of one or more edge-shared \( \{W_3O_{13}\} \) groups by \( \pi/3 \) [7, 8].

![Figure 1.1: Polyhedral representation of the α-Keggin (left) ion and of the trilacunary derivative \( \{α-A-XW_{10}\} \) (right). The metal ions form the centers of the blue polyhedra and the oxygen atoms form the apexes of the polyhedra. The central heteroatom is shown as yellow polyhedra.](image-url)
Lacunary derivatives of the Keggin type anions geometrically result from the removal of one or more WO groups. The two tri-vacant species correspond to the loss of a corner-shared group of \(\{WO_6\}\) octahedral (A-type) or an edge-shared group (B-type) (Figure 1.1) [5, 6].

**Wells-Dawson-isomers and hexalacunary structures**

The Wells-Dawson anion \([P_2W_{18}O_{62}]^{6-}\) results from the direct association of two \([A-PW_{9}O_{34}]^{9-}\) units. The structure has two types of tungsten atoms, six "polar" and twelve "equatorial" (Figure 1.2). Six isomers of this anion are theoretically possible depending upon whether the half-units are derived from \(\alpha\) or \(\beta\)-Keggin species and also whether the fragments combined in a staggered (S) or eclipsed (E) fashion [8]. Four of these isomers have been observed for \([As_2W_{18}O_{62}]^{6-}\) and three for \([P_2W_{18}O_{62}]^{6-}\) [9].

Figure 1.2: Polyhedral representation of the \(\alpha\)-Dawson (left) ion and of the hexavacant derivative \(\{\alpha-A-P_2W_{12}\}\) (right). The metal ions form the centers of the blue polyhedra and the oxygen atoms form the apexes of the polyhedra. The central heteroatom is shown as yellow polyhedra.

As in the case of the Keggin ion, lacunary derivatives of the Well-Dawson structure are also known. The most important of these are based on the most common isomer known as the \(\alpha\)-Dawson anion. Lacunary derivatives of the \(\alpha\)-Dawson anion include a metastable hexavacant species \(\{P_2W_{12}\}\) (Figure 1.2) [10]. A well known polyoxoanion which can be described as a derivative of the \(\{P_2W_{12}\}\) unit is the tetramer \([K_8\subset P_8W_{48}O_{184}]^{32-}\) or \(\{P_8W_{48}\}\) with the cyclic
structure shown in Figure 1.3. The crystal structure of the related salt reveals that the central cavity of the anion encapsulates potassium cations [11].

![Figure 1.3: Structure of the anion \(\{P_8W_{48}O_{184}\}^{40-}\), \(\{P_8W_{48}\}\) as a cyclic assembly of four lacunary \(\{P_2W_{12}\}\) groups. A ball and stick representation is shown on the left and a polyhedral representation on the right side.](image)

### 1.2.2 Polyoxomolybdates

From a unique library containing molybdenum-oxide based building blocks/fragments in aqueous solution under reducing conditions a huge variety of nanoobjects, allowing specific reactions at well-defined positions, can be generated. Examples include: the molecular big-sphere of the type \(\{Mo_{132}\}\), the molecular big-wheel of the type \(\{Mo_{154}/Mo_{176}\}\) and in addition the by far largest structurally well characterized cluster \(\{Mo_{368}\}\) with the shape of a lemon.

Specifically speaking the reasons for such a versatile behavior of polyoxomolybdates are:

1. The easy change of coordination numbers as well as easy exchange of H\(_2\)O ligands at Mo sites.
2. The moderate strength of Mo-O-Mo type bonds allowing "split and link" type processes.
3. The easy change and especially increase of electron densities without the strong tendency to form metal-metal bonds.
4. The presence of terminal Mo = O groups preventing in principle unlimited growth to extended structure [12].
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The giant spherical polyoxomolybdates

If we intend to construct a giant species similar in shape to the spherical viruses with icosahedral symmetry (containing C₅, C₃ and C₂ axes), we have to find a reaction system in which pentagonal units can first be generated, then get linked and be placed at the 12 corners of an icosahedron. Reaction mixture of molybdates under appropriate pH values and reducing conditions house a potential library ideal for such constructions (Figure 1.4). In the giant spherical clusters, the pentagonal \((Mo)Mo₅\) building blocks, each of which consists of a central pentagonal bipyramidal \(MoO₅\) unit sharing edges with five \(MoO₆\) octahedra, are placed at the 12 vertices of an icosahedron and linked by a set of 30 mono- or dinuclear spacers/linkers, such as \(Fe^{III}(H₂O)\)³⁺ [13], \(Mo^{VI}O(H₂O)\)³⁺ [14], \(V^{IV}O(H₂O)\)²⁺ [15] or \(Mo^{V}O₄(ligand)\)ⁿ⁺ (e.g. ligand = HCOO⁻, CH₃COO⁻, SO₄²⁻, H₂PO₂⁻, PO₄³⁻), respectively [16, 17]. In this context it is worthwhile to mention that the linkers span (distorted) Archimedean solids with approximately icosahedral symmetry: in the case of dinuclear metal linkers a (distorted) truncated icosahedron, \(M₂_{30}\) and in the case of mononuclear linkers the unique icosidodecahedron \(M₃₀\) (Figure 1.5).

Figure 1.4: Construction principle for a cluster with icosahedral symmetry. The \((Mo)Mo₅\) units (polyhedral model with central pentagonal bipyramid in light grey) are the basis for the formation of the \((pent₁₂(link)₈₉)\) type clusters, where the \((Mo)Mo₅\) units define the icosahedron vertices.
All such spherical clusters, which can be described by the general formula \([\{(Mo)Mo_5\}_{12}[L]_{30}\] or \([(pentagon)_{12}(linker)_{30}\], belong to the family of “Keplerate” type molecules because of their similarity to Kepler's early model of the Universe, as described in his speculative opus Mysterium Cosmographicum [18]. A Keplerate has, accordingly to our definition, one central point - whether or not occupied by an atom - and its atoms are organized in one or more spherical shells around this central point while each set of equivalent atoms forms a Platonic or a (generalized) Archimedean solid [19].

These ideas are the base of the present work.

Figure 1.5: “Sizing” the nanospheres is possible. Structure only formed by the linkers: Left: the icosidodecahedron with 12 pentagons and 20 triangles formed by mononuclear linkers. Right: the (distorted) truncated icosahedron with twelve pentagons and twenty hexagons formed by 30 dinuclear \([Mo_2]\) linkers. Colour code: central pentagonal bipyramidal \([MoO_7]\) of \([MoMo_3]\) in cyan and other octahedra in blue.

**The molecular big-wheels**

The mentioned spherical Keplerates are results of the spherical disposition of pentagonal \((Mo)Mo_3\) building blocks with a \(C_5\) symmetry whereas a circular disposition of the deformed \((Mo)Mo_3\) unit can lead to the formation of \(\{Mo_{154}\}\) and \(\{Mo_{176}\}\) type clusters, commonly known as the big-wheel and giant wheel type species, respectively [20]. Their structures can formally be represented as \([\{Mo_8\}/Mo'_{2}/\{Mo_1\}\]_n, where \(n = 14\) and 16 for \(\{Mo_{154}\}\) and \(\{Mo_{176}\}\),
respectively. (Figure 1.6) The \{Mo_8\} building block is built up by a central pentagonal \((Mo)Mo_5\) unit (containing a central \{MoO_7\} or \{MoO_6(NO)\} bipyramid sharing edges with five \{MoO_6\} octahedral) and two more weakly bonded (sharing only corners) \{MoO_6\} octahedra which can be more easily “removed”.

In addition to the \{Mo_8\} unit, the wheel type polyoxomolybdates contain as mentioned above \\{Mo’\} = \{Mo_2O_5(H_2O)\}^{2+}, formed by two corner-sharing \{MoO_6\} octahedra, together with \{Mo_1\} type units. The cluster anions are 2x14- and 2x16-fold reduced, respectively, with the related Mo (4d) electrons trapped in 14 and 16 \{Mo_5O_6\}-type “compartments”, over which they are delocalized [20]. The \{Mo_{154}\} cluster has an external diameter of 3.4 nm while that of \{Mo_{176}\} is 4.1 nm [21].

![Figure 1.6: Top (left) and side (right) polyhedral views of the hexameric \{Mo_{176}\} cluster. The \{Mo_8\}, \{Mo_2\} and \{Mo_1\} building blocks are shown below and the positions of \{Mo_1\} units are ringed on the side view on the right side.](image)
Chapter 1. Introduction

The "Blue Lemon"

The special building units abundant in a "dynamic library" can form, via a type of "split and/or link process" (for instance the mentioned \{(Mo)Mo₅\} units from larger ones which subsequently become linked) a class of molecular architectures upon slight variation of the boundary conditions. The \{Mo₃₆₈\} cluster [12, 22] is such a molecular type (Figure 1.7); comparable to the size of hemoglobin (external diameter ca. 6 nm), it contains 368 metal (1880 non-hydrogen) atoms formed by the linking of 64 \{Mo₁\}, 32\{Mo₂\}, and 40 \{(Mo)Mo₅\} type units (32 with sulfate ligands and 8 without) via a remarkable symmetry breaking process which is nicely recognizable at the cluster surface.

![Figure 1.7: \[HₓMo₃₆₈O₁₀₃₂(H₂O)₂₄₀(SO₄)₄₈\]⁴⁻ cluster anion in polyhedral representation showing the different building blocks and large areas of different local symmetry; building units \{Mo₁\} (yellow), \{Mo₂\} (red), \{(Mo)Mo₅\} (blue with blue-turquoise pentagonal MoO₇ bipyramids).](image)

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In terms of building blocks the \( \{\text{Mo}_{168}\} \) can be described as \( [H_x\{\text{Mo(Mo)5}\}'8\{\text{Mo(Mo)5}\}''32\{\text{Mo2}\}'16\{\text{Mo2}\}'''8\{\text{Mo1}\}64]^{48-} \) (x=16) and has an approximate \( D_4 \) symmetry with a central ball-shaped fragment with \( D_{8d} \) symmetry \( \{\text{Mo}_{288}\} \equiv \{\text{Mo}_{288}\text{O}_{784}(\text{H}_2\text{O})_{192}(\text{SO}_4)_{32}\} \) and two capping fragments \( \{\text{Mo}_{40}\} \equiv \{\text{Mo}_{40}\text{O}_{124}(\text{H}_2\text{O})_{24}(\text{SO}_4)_{8}\} \) with \( C_{4v} \) symmetry. The structure of the cluster anion can be considered as a hybrid between the \( \{\text{Mo}_{176}/\text{Mo}_{154}\} \) type giant molecular wheels and the \( \{\text{Mo}_{102}\} \) type clusters.

1.3 Goals of the project: defining the problem

After describing the basic principles and highlights of polyoxometalate chemistry the aims of the research developed in the course of this thesis are as bellow:

- To explore the utility of pentagonal \( \{(\text{Mo})\text{Mo}_5\}\)-type units, available in a dynamic polyoxomolybdate library as (virtual) building blocks, in the synthesis of giant spherical clusters based on the linking of the units by paramagnetic centers.
- To study metal-oxide based nucleation processes under confined conditions within the cavity of the \( \{\text{P}_8\text{W}_{48}\}\)-type cluster.

The results obtained will be described in the next sections.
Bibliography


Chapter 2
Publications
2.1. Triangular Geometrical and Magnetic Motifs Uniquely Linked on a Spherical Capsule Surface


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Contribution of A. M. Todea to the publication:
- Synthesis and characterization (electronic absorption as well as vibrational spectra, redox titrations, elemental analysis) of the new compound.

2.2. Formation of a "less stable" polyanion directed and protected by electrophilic internal surface functionalities of a capsule in growth: $[\{\text{Mo}_{6}O_{19}\}^{2-} \subset \{\text{Mo}^{\text{VI}}_{72} \text{Fe}^{\text{III}}_{30}O_{252}(ac)_{20}(H_{2}O)_{92}\}]^{4-}$

A. Müller, A. M. Todea, H. Bögge, J. van Slageren, M. Dressel, A. Stammler, M. Rusu

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Contribution of A. M. Todea to the publication:
- Synthesis and characterization (electronic absorption as well as vibrational spectra, elemental analysis) of the new compound.
2.3. Extending the \{(Mo)Mo_{5}\}_{12}M_{30} Capsule Sequence: New Cr_{30} Cluster of s = 3/2 Metal Centres with a \{Na(H_{2}O)_{12}\} Encapsulate


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2.4. Unique Properties of Mo_{72}Fe_{30} Cluster in Solution

A. Müller, A. M. Todea

Manuscript in preparation

Contribution of A. M. Todea to the publication:
- Synthesis and characterization (vibrational spectra) of the deuterated compound.
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A. Müller, M. T. Pope, A. M. Todea, H. Bögge, J. van Slageren, M. Dressel, P. Gouzerh, R. Thouvenot, B. Tsukerblat, A. Bell


Contribution of A. M. Todea to the publication:
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Contribution of A. M. Todea to the publication:
- Synthesis and characterization (electronic absorption as well as vibrational spectra, elemental analysis) of the new compound.
Triangular Geometrical and Magnetic Motifs Uniquely Linked on a Spherical Capsule Surface**

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Dedicated to Professor Francis Sécheresse on the occasion of his 60th birthday

Polygons can be placed on spherical surfaces such that periodical structures of a cyclic nature result, while these can be considered as discrete models for two-dimensional (extended) structures. If we wish to construct a chemical structure on a spherical capsule surface in the same way, we have to remember that 1) pentagons are the basic units for sphere constructions, as is well known, for example, from virus structures, 2) they exist, for example, in the form of \([\text{Mo}^{VI}]\text{Mo}^{VI}\) type units, and that 3) they occur in Keplerates of the type \([\text{Mo}^{VI}]\text{Mo}^{VI}\{\text{Linker}\}_{12}\) [1–5] (linker can be of the mononuclear \(\text{M} = \text{metal center}\) or dinuclear type \(\text{M}_2\); for the definition of Keplerates, see ref. [5b]). However, until now it was not possible to synthesize a spherical capsule surface directly by the addition of linkers to the pentagonal units that are available in a dynamic library. [1–3] It is significant that in the Keplerates the linkers describe generic Archimedean solids: in the case of dinuclear linkers \(\text{M}_2\) a distorted truncated icosahedron, \([\text{M}_2]_{10}\), and in the case of mononuclear linkers the unique icosidodecahedron (Figure 1) [6] [M30], which has—geometrically speaking—linked \(\text{M}_1\) triangles. Surprisingly the related consequences for chemistry have not been discussed until now. In the \([\text{M}_30]\) situation, there is a network of corner-shared triangles on the sphere surface, this can result unique magnetic properties as in the case of the “classical” Keplerate \([\text{Mo}^{VI}]\text{Mo}^{VI}\{\text{Fe}^{III}\}\) which otherwise have only been observed in selected one-, two-, and three-dimensional lattice spin systems. [9] Herein we report on the spherical cluster 1a where the twelve \([\text{Mo}^{VI}]\text{Mo}^{VI}\{\text{V}^{III}\}\) type units fix 30 d3 V30 linkers/centers with spin \(S = 1/2\) in the form of an icosidodecahedron, and thus 1) demonstrating for the first time that the spherical capsule/Keplerate can be directly constructed from the mononuclear linkers and the appropriate molybdate library; [8b] 2) providing the chance to obtain new information regarding the unique molecular magnetism of the \([\text{M}_30]\) type network of linkers/triangles, and 3) clarifying the quantum effects of the spin \(S = 1/2\) vanadyl linkers especially in connection with the two-dimensional \(S = 1/2\) Kagomé lattice which contains linked triangles and exhibits unique magnetic properties. [9c]

After adding vanadyl sulfate to an acidified molybdate solution, in the presence of \(K^+\) ions, compound 1 precipitates after some time in high yield. (A simpler expression for the

Figure 1. a) The M centers (small colored spheres) of the \([\text{Mo}^{VI}]\text{Mo}^{VI}\{\text{Fe}^{III}\}\) Keplerate, e.g., \(\text{M} = \text{V}^{III}\), \(\text{Fe}^{III}\) describe the icosidodecahedron shown, which is unique among the icosahedral Archimedean solids as all edges are equivalent and all dihedral angles equal. Referring to the special situation of \(\text{M} = \text{Fe}^{III}\), there are three groups (“sub-lattices”) of 10 spins (colors: red, blue, green), with all spins of a sub-lattice pointing in the same direction, while nearest-neighbor spin vectors (three are highlighted) differ in angular orientation by 120°. Also shown: b) A fragment highlighting five linked triangles around a pentagon. c) A fragment of a planar Kagomé lattice with six linked triangles around a hexagon.

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[1–5] (linker can be of the extended) structures. If we wish to construct a chemical structure on a spherical capsule surface in the same way, we have to remember that 1) pentagons are the basic units for sphere constructions, as is well known, for example, from virus structures, 2) they exist, for example, in the form of \([\text{Mo}^{VI}]\text{Mo}^{VI}\) type units, and that 3) they occur in Keplerates of the type \([\text{Mo}^{VI}]\text{Mo}^{VI}\{\text{Linker}\}_{12}^{[1–5]}\) (linker can be of the mononuclear \(\text{M} = \text{metal center}\) or dinuclear type \(\text{M}_2\); for the definition of Keplerates, see ref. [5b]). However, until now it was not possible to synthesize a spherical capsule surface directly by the addition of linkers to the pentagonal units that are available in a dynamic library. [1–3] It is significant that in the Keplerates the linkers describe generic Archimedean solids: in the case of dinuclear linkers \(\text{M}_2\) a distorted truncated icosahedron, \([\text{M}_2]_{10}\), and in the case of mononuclear linkers the unique icosidodecahedron (Figure 1) [6] [M30], which has—geometrically speaking—linked \(\text{M}_1\) triangles. Surprisingly the related consequences for chemistry have not been discussed until now. In the \([\text{M}_30]\) situation, there is a network of corner-shared triangles on the sphere surface, this can result unique magnetic properties as in the case of the “classical” Keplerate \([\text{Mo}^{VI}]\text{Mo}^{VI}\{\text{Fe}^{III}\}\) which otherwise have only been observed in selected one-, two-, and three-dimensional lattice spin systems. [9] Herein we report on the spherical cluster 1a where the twelve \([\text{Mo}^{VI}]\text{Mo}^{VI}\{\text{V}^{III}\}\) type units fix 30 d3 V30 linkers/centers with spin \(S = 1/2\) in the form of an icosidodecahedron, and thus 1) demonstrating for the first time that the spherical capsule/Keplerate can be directly constructed from the mononuclear linkers and the appropriate molybdate library; [8b] 2) providing the chance to obtain new information regarding the unique molecular magnetism of the \([\text{M}_30]\) type network of linkers/triangles, and 3) clarifying the quantum effects of the spin \(S = 1/2\) vanadyl linkers especially in connection with the two-dimensional \(S = 1/2\) Kagomé lattice which contains linked triangles and exhibits unique magnetic properties. [9c]

After adding vanadyl sulfate to an acidified molybdate solution, in the presence of \(K^+\) ions, compound 1 precipitates after some time in high yield. (A simpler expression for the...
Compound 1, which crystallizes in the monoclinic space group C2/c, was characterized by elemental analysis, thermogravimetry, spectroscopic methods (IR, Raman, UV/Vis), single-crystal X-ray structure analysis (including bond valence sum (BVS) calculations), and susceptibility measurements (including related quantum Monte Carlo calculations).

The cluster anion 1a of 1 is of the expected (Pentagon)$_{12}$-(Linker)$_{30}$ type and is a slightly compressed sphere, while the heptacoordinate Mo VI centers of the 12 pentagonal units correspondingly describe a slightly distorted icosahedron and the 30 V IV centers—acting as linkers for the pentagonal [(Mo V )Mo V ]$_{2}$ type units—describe a (slightly distorted) icosidodecahedron (Figure 2; the V IV–V IV distances in the distorted Archimedean solid vary from 6.3 to 6.6 Å). The distortion is in agreement with the fact that 20 V IV centers in the equatorial region have octahedral coordination and the two sets of five V IV centers in the polar area have square-pyramidal coordination; the distances from the 10 equatorial V IV units to the center of the cluster are a little shorter (10.3 Å) than the related distances of the other 20 V IV units (10.6 Å). Ten of the twelve [SO$_4$]$^{2-}$ ligands are coordinated by three oxygen atoms to three adjacent Mo V centers of the [(Mo V )Mo V ]$_{2}$ groups such that two [KSO$_4$]$_{5}$ rings parallel to the equator result, with the K$^+$ ions (formally) bridging the [SO$_4$]$^{2-}$ ions (Figure 3). The other two sulfate groups are disordered and act as ligands to the two polar [(Mo V )Mo V ]$_{2}$ groups. The structure of 1a comprising the twenty triangular and twelve pentagonal faces of the icosidodecahedron built up by 30 V IV centers shows an interesting relation to the much less symmetrical cluster anion 2a which has a non-complete spherical [V$_3$]$_{20}$ type net. In 2a, a strongly distorted icosidodecahedron is described by 10 Mo V and 20 V IV centers, while the equatorial [V$_3$]$_{20}$ belt-formed by 10 linked [V$_3$] triangles—is identical to the related equatorial segment of 1a.

The presence of K$^+$ and [SO$_4$]$^{2-}$ ions in the reaction medium seems to be of fundamental importance for the structure formation, as the potassium cations of the two [KSO$_4$]$_{5}$ rings of 1a attract the two negatively charged polar [(Mo V )Mo V ]$_{2}$ units thus causing the slight compression of the sphere. This distortion leads to an inclination of the adjacent [VO$_6$]$_{6}$ polyhedra and thus prevents an octahedral coordination of the 10 polar V IV centers. The sixth (H$_2$O) ligand required for octahedral coordination would be too close to the [SO$_4$]$^{2-}$ ligands of the [KSO$_4$]$_{5}$ rings.

The investigations nicely show that [(Mo V )Mo V ]$_{2}$ type units are potentially available in a dynamic polymolybdate library; remarkably, they can be “used” in the present case as

\[\text{Na}_8K_{14}(VO_2)_{10}[(Mo^{IV})Mo^{V}O_4(H_2O)_3]_{20}[(Mo^{IV})Mo^{V}O_4(H_2O)_3]_{20}[(H_2O)_{3}(SO_4)]_{12}\text{[SO}_4\text{]$_{2}$]}_{20}\]
FeIII centers by VO

{FeIII}

icosidodecahedron as is approximately the case in the 3d electrons are not "localized" at the vertices of the cluster skeleton. Therefore, the room temperature value of \( T_H \) corresponds to 30 uncoupled \( S = 5/2 \) ions. The quantum Monte Carlo (QMC) method provides accurate values of susceptibility for the Heisenberg model of the \([\text{V}_{30}]\) system for \( T > 0.5J/k_B \) and as seen in Figure 4, a very good fit to \( T_H \) is achieved for \( T > 120 \) K for the choices \( J/k_B = 245 \) K and \( g = 1.95 \). Unfortunately, reliable results cannot be obtained for lower temperatures using the QMC method owing to the "sign problem" that occurs for spin systems with antiferromagnetic exchange based on lattice geometries where the classical counterpart exhibits spin frustration.

To summarize: We have demonstrated that it is possible to "use" pentagonal units as "building units" which play, geometrically speaking, the same role as the pentagonal units in other sphere-based constructions, such as spherical viruses, fullerenes, and geodesic domes; in our context they are used as a glue for trapping magnetic centers, such that triangles are linked to form an icosidodecahedron, that is, a part of a Keplerate. As the "quantum" Keplerate \( \text{Ia} \) obtained is a new example of a frustrated magnetic system which shares a topological feature with the classical Keplerate \([\text{MoV}\text{III}_5]\text{FeV}\text{III}_{30}\) and the Kagomé-lattice antiferromagnet, its exploration is expected to provide a deeper understanding of basic aspects of magnetic frustration and the role played by the size of the intrinsic spin of the interacting magnetic ions. This study should also shed light on the parallel problem, and the focus of an intense effort, to characterize and understand the \( S = 1/2 \) Kagomé lattice, which is considered to have unique magnetic/electronic properties originating in the small value of \( S \).[17] The behavior of the magnetization of the quantum Keplerate \( \text{Ia} \) in high magnetic fields will be of key importance in studies aimed at elucidating the characteristics of its magnetic frustration.[17]

**Experimental Section**

A solution of VOSO₄·5H₂O (2.53 g, 10 mmol) in H₂O (35 mL) was added to a stirred solution of Na₂MoO₄·2H₂O (2.42 g, 10 mmol) in H₂SO₄ (0.5 m; 8 mL) in an conical flask. The resulting dark purple
mixture was stirred at room temperature for 30 min (flask closed with a rubber stopper) and then treated with KCl (0.65 g, 8.72 mmol). After additional stirring for 30 min the solution was stored in the flask which was closed with a rubber stopper. After 5 days, the purple-black rhomboic crystals of 1 were collected by filtration, washed with cold water, and finally dried in air. Yield: 1 g; elemental analysis: calcd for Na$_2$K$_2$Mo$_7$V$_3$S$_5$O$_{25}$: C, 24.8; H, 2.3; N, 0.3%. FT-Raman (solid; n$\tilde{\nu}$=1664 cm$^{-1}$); UV/Vis (in H$_2$O): $\lambda$ = 510 (vs), 689 (w), 784 (w) nm.

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[2] In this context we should refer first to the fact that molybdenum oxide building units—abundant as virtual building blocks, for instance, occur in spherical capsules/Kepler-center, see a) M. O’Keefe, B. G. Hyde, Crystal Structures, I. Patterns and Symmetry, Mineralogical Society of America, Washington, DC, 1996; b) H. S. M. Coxeter, Introduction to Geometry, Wiley, New York, 1989; c) H. S. M. Coxeter, Regular Polytides, 3rd ed., Dover, New York, 1973, in which we read (p. 18): “A quasi-regular polyhedron (like the icosidodecahedron) is defined as having regular faces, while its vertex figures, though not regular, are cyclic and equiangular (i.e., inscribable in circles and alternate-sided). It follows from this definition that the edges are all equal, say of length 2L, that the dihedral angles are all equal, and that the faces are of two kinds, each face of one kind being entirely surrounded by faces of the other kind.”


[6] The (quasi-regular) icosidodecahedron is a hybrid of the icosahedron and the dodecahedron and thus contains 20 trigonal and 12 pentagonal faces which intersect at each corner. Unlike the other related Archimedean solids, such as the [Mo$_6$O$_{19}$]-truncated icosahedron, it comprises not only equivalent vertices but also equivalent edges and equal dihedral angles (Figure 1); see a) M. O’Keefe, B. G. Hyde, Crystal Structures, I. Patterns and Symmetry, Mineralogical Society of America, Washington, DC, 1996; b) H. S. M. Coxeter, Introduction to Geometry, Wiley, New York, 1989; c) H. S. M. Coxeter, Regular Polytides, 3rd ed., Dover, New York, 1973, in which we read (p. 18): “A quasi-regular polyhedron (like the icosidodecahedron) is defined as having regular faces, while its vertex figures, though not regular, are cyclic and equiangular (i.e., inscribable in circles and alternate-sided). It follows from this definition that the edges are all equal, say of length 2L, that the dihedral angles are all equal, and that the faces are of two kinds, each face of one kind being entirely surrounded by faces of the other kind.”; d) A. Holden, Shapes, Space and Symmetry, Dover, New York, 1991.


[10] Crystal data for 1: H$_4$Na$_2$K$_2$Mo$_7$V$_3$S$_5$O$_{25}$, M$_r$=19068.10 g mol$^{-1}$, monoclinic, space group C2/c, a = 47.155(2), b = 42.536(18), c = 26.5344(12) Å, $\beta$ = 90.2510(10)$^\circ$, V = 53222.4(2) Å$^3$, Z = 4, $\rho$ = 2.380 g cm$^{-3}$, $\mu$ = 2.530 mm$^{-1}$, $F(000)$ = 36.848, crystal size = 0.30 × 0.20 × 0.02 mm$^3$. Crystals of 1 were removed from the mother liquor and immediately cooled to 183(2) K on a Bruker AXS SMART diffractometer (three circle goniometer with 1 K CCD detector, Mo$_{K\alpha}$ radiation, graphite monochromator; hemisphere data collection in $\omega$ at $0.3^\circ$ scan width in three runs with 606, 435, and 230 frames ($\phi$ = 0.88, and 180$^\circ$) at a detector distance of 5 cm). A total of 158364 reflections (1.50 < $\theta$ < 26.99$^\circ$) were collected of which 57718 reflections were unique (R(int) = 0.0612). An empirical absorption correction using equivalent reflections was performed with the program SADABS. The structure was solved with the program SHELXS-97 and refined using SHELXL-97 to $R$ = 0.0567 for 40563 reflections with $I$ > 2$\sigma(I)$. $R$ = 0.0916 for all reflections; max/min residual electron density 3.456 and –2.026 e Å$^{-3}$. (SHELXS/L, SADABS from G. M. Sheldrick, University of Göttinngen 1997; structure graphics with DIA-MOND 2.1 from K. Brandenburg, Crystal Impact GbR, 2001.)

In spite of these expected differences, there are some important features common to the two Keplerates, such as similar characteristics of the ground state and the low-lying excitation states, which arise because of their otherwise common geometrical structure and the existence of antiferromagnetic exchange between nearest-neighbor spins. As one example, the total spin quantum number of the ground state eigenvector for the “quantum” Keplerate 1 is expected to be $S = 0$, as it is for the “classical” Keplerate [Fe$_{30}$], and its direct manifestation would be in the vanishing of $T_c$ (proportional to $\langle S^2 \rangle$), in the low-temperature regime; $\chi$ is the zero-field susceptibility.

The raw $M/H$ data display a rapid rise on decreasing the temperature below 50 K that is very accurately simulated by a term proportional to $1/T$ (Curie behavior). This observation suggests the presence of additional non-interacting paramagnetic centers in the sample. In fact, fitting the low temperature data leads to a value of approximately $2S = 1/2$ centers. These centers are considered to be the non-interacting VO$_2$$^+$ (d$^1$) ions. In large cluster systems like 1, they can take the same place as the other diamagnetic cations, which are usually disordered in the large voids between the clusters and can therefore not be discovered by single-crystal structure analysis (see ref. [10]). On subtracting the contribution of these discrete paramagnetic centers, we obtain a corrected magnetization $M'$ as well as the corrected susceptibility data, $T/M''$, shown in Figure 4. The Curie behavior of the paramagnetic centers shows that they do not interact with each other and with the [V$_{15}$] cluster. The possibility that the finite $T_c$ value at low temperatures is due to a non-zero spin ground state can be precluded. Regarding the presented $T_c$ curve: As always there is uncertainty as to the appropriate choices for diamagnetic and temperature-independent paramagnetism (TIP) corrections. Additionally, because of the large voids between the clusters the VO$_2$$^+$ groups need not be present stoichiometrically in the compound; correspondingly, there is a very small error limit in the given/used number of two VO$_2$$^+$ groups which influences the correction of the raw magnetic data ((VO)$_{1.8}$ could, for example, correspond to K$_{14.4}$).

Note added in proof, April 27, 2005: Very recent measurements by H. Nojiri (Tohoku University) of $M$ versus $H$ at 0.5 K up to 27 Tesla, as well as ESR measurements at 190 GHz for several temperatures, show features which are fully consistent with our physical interpretation of a strong intrACLuster exchange constant and approximately two VO$_2$$^+$ ions per formula unit that are magnetically independent of the [V$_{15}$] cluster. Full details will be published elsewhere.
Formation of a “less stable” polyanion directed and protected by electrophilic internal surface functionalities of a capsule in growth: [{Mo$_6$O$_{19}$}]$^{2-}$ ⊂ {Mo$^{VI}$$_{72}$Fe$^{III}_{30}$O$_{252}$(ac)$_{20}$(H$_2$O)$_{92}$}]$^{4-}$

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The spherical capsule skeleton of the host–guest system [{Mo$_6$O$_{19}$}]$^{2-}$ ⊂ {Mo$^{VI}_{72}$Fe$^{III}_{30}$O$_{252}$(CH$_3$COO)$_{20}$(H$_2$O)$_{92}$}]$^{4-}$ ⊂ 1a—built up by 12 (Mo$^{VI}$)Mo$^{VI}$ type pentagonal units linked by 30 Fe$^{III}$ centers which span the unique icosahedral Archimedean solid, the icosidodecahedron—can now be constructed deliberately and with a simpler composition than before from an acidified aqueous molybdate solution containing the mentioned (virtual) pentagonal units; the encapsulated heptamolybdate—normally not formed in water—is built up in an unprecedented way concomitant with capsule growth, while being directed by the corresponding internal electrophilic surface functionalities.

It is well known that aesthetically beautiful spherical objects can, geometrically speaking, be constructed from pentagons according to the related building block principle known for instance from virology, architecture (geodesic domes) as well as from daily life. But the use of this principle in synthetic chemistry has not been deliberately exploited. Though it became known that pentagonal units of the type ((Mo$^{VI}$)Mo$^{VI}$) occur in spherical capsules/Keplerates of the general formula (Mo$^{VI}$)$_{12}$[Linker]$_{30}$ (the linker can be of a mononuclear (M) or dinuclear metal type (M$^2$)) it could only be proven in the particular case of M = VO$^{2+}$ that the spherical capsules can be directly synthesized by the addition of linkers to the dynamic library containing (virtual) pentagonal units† (see also ref. 5 regarding general remarks). The related cluster type with the skeleton (Mo$^{VI}$)Mo$_{12}$[Fe$^{III}$]$_{30}$ could until now not be synthesized in a straightforward reaction and contains inside the capsule—due to the specific preparation process—complex units of the type {Mo$_5$O$_{80}$}$^{10-}$ as ligands, which complicates the structure and, in principle, the interpretation of its (magnetic) properties.

Herein we report on the synthesis of the (Mo$^{VI}$)Mo$_{12}$[Fe$^{III}$]$_{30}$ type cluster 1a exhibiting several new features: (1) the spherical capsule/Keplerate can be directly constructed based on an appropriate molybdate library containing the above mentioned virtual pentagonal units, (2) the mentioned {Mo$_5$O$_{80}$}$^{10-}$ units are present, and most important (3) the hexamolybdate [Mo$_6$O$_{19}$]$^{2-}$ anion is encapsulated noncovalently bonded, a discovery which will have implications for future theoretical and synthetic work. Furthermore, it should be noted that the inorganic skeleton of 1a is of tremendous interest because of its unique magnetic properties.$^8$$^9$a–c

Na$_2$[Mo$_6$O$_{19}$]$^{2-}$ ⊂ {Mo$^{VI}_{72}$Fe$^{III}_{30}$O$_{252}$(CH$_3$COO)$_{20}$(H$_2$O)$_{92}$}]$^{4-}$ ca. 120 H$_2$O = Na$_2$1a · ca. 120 H$_2$O 1

If an acidified aqueous solution of sodium molybdate which contains a rather high concentration of acetic acid is treated with iron(III) chloride, yellow crystals of 1 precipitate after some days.$^†$ Compound 1, which crystallizes in the space group $\mathbf{R}$3, was characterized by elemental analysis (including ESCA), thermogravimetry (to determine the actual(!) crystal water content), spectroscopy (IR, Raman, UV-Vis), and single-crystal X-ray structure analysis (including bond valence sum (BVS) calculations)$^8$ and magnetic susceptibility measurements.

The cluster anion 1a (the anions occupy in 1 two crystallographically independent positions 1 and 3) is of the expected spherical (Mo$^{VI}$)Mo$_{12}$[Linker]$_{30}$ type mentioned above. Correspondingly, the heptacoordinated Mo centers of the 12 pentagonal (Mo$^{VI}$)Mo$^{VI}$ type units span an icosahedron, and the 30 Fe$^{III}$ linkers, the quasi-regular icosidodecahedron (Fig. 1) which is a hybrid of the icosahedron and the dodecahedron and contains correspondingly 20 trigonal and 12 pentagonal faces.$^\dagger$ (Fig. 1). The 20 acetate ligands in 1a, some of which are disordered, are located inside the sphere and are coordinated in a bidentate fashion to the metal centers, preferentially bridging Fe and Mo sites. 1a does not contain the {Mo$_5$O$_{80}$}$^{10-}$ type units in contrast to the former reported [Fe$_{30}$]$_{30}$ type compound.$^6$

The most interesting aspect of the present work is the encapsulation of the hexamolybdate [Mo$_6$O$_{19}$]$^{2-}$ in the cavity of the capsule (Fig. 2). This polyanion cannot be obtained as other polyoxomolybdates simply in water and is preferably formed in organic solvents, thereby crystallizing together with organic cations.$^{10}$ In the present system it is stabilized/protected by the comparably large number of acetates inside the cavity of 1a, which form a hydrophobic environment. It is important to note that negatively charged polynoxometalates cannot be encapsulated in (highly) negatively charged Keplerates/capsules like the {Mo$_{12}$]$_{32}$ cluster.$^3$ In the present case 1a is “approximately neutral” and the pure inorganic part/skeleton, i.e. without the acetate ligands, is even positively charged. (Note: a few deprotonated H$_2$O ligands coordinated to the Fe centers cause a small negative charge.) This allows attraction/collection of small anionic fragments with the consequence that the stepwise formation of [Mo$_6$O$_{19}$]$^{2-}$ can occur
be obtained under non-reducing conditions. Correspondingly, at low pH values and furthermore the largest isopolymolybdate to 'split-and-link' mechanism with the \{(MoVI)MoVI\}_5^{12}\{Linker\}^{30} type Keplerates, which have—because of relatively weakly bound in the \([MoVI\_{36}O_{22}(OH)_8\]^{11-} anion)—a high formation tendency, are formed from that solution in the presence of (potential) linkers like FeIII ions. The reaction is based on a 'split-and-link' mechanism with the \{(MoVI)MoVI\}_5^{12} unit being abstracted from the \{Mo_{50}\} type species.† (Related results regarding mixed metal species like \{Mo_{36}VIV\}_{36}, or \{Fe^{III}_{22}VIV\}_{12}^{12-} will be reported later.) As pentagonal \{(MoVI)MoVI\}_5^{12} building blocks of the \{Pentagon\}_{12}\{Linker\}_{30} type occur in spherical capsules/Keplerates also with the binuclear linkers \{Mo_{50}O_{24}\}^{12-} it is possible to apply also to them the synthetic principle outlined here for the mononuclear linkers.13

To characterize the magnetic properties of I, the magnetic susceptibility versus temperature as well as the magnetization as a function of field were measured (this had to be done in order to prove whether an influence of the \{Mo_{36}O_{22}F^{-}\} units exists; see above). The room temperature value \(\chi T = 120.1\ \text{cm}^3\ \text{K mol}^{-1}\) is slightly smaller than that expected for 30 noninteracting iron(III) ions (\(\chi T = 131.3\ \text{cm}^3\ \text{K mol}^{-1}\), assuming \(g = 2.00\)). On lowering the temperature, the \(\chi T\) product decreases, reaching a value of \(\chi T = 10.3\ \text{cm}^3\ \text{K mol}^{-1}\) at \(T = 1.8\ \text{K}\), which is indicative of predominant antiferromagnetic exchange interactions. A Curie–Weiss plot of \(\chi^{-1}\) versus \(T\) yields a Curie constant of \(C = 129.4 \pm 0.2\ \text{cm}^3\ \text{K mol}^{-1}\) and a Weiss temperature of \(\theta = -22.3 \pm 0.3\ \text{K}\). This Weiss temperature is virtually identical to that found for the earlier \{Mo_{36}Fe_{30}\} cluster,9b which shows that the magnetic properties of the cluster have not been affected by the changed synthetic procedure. This is confirmed by the magnetization curve which is virtually linear as a function of field. The numerical derivative shows the typical minimum at about 5.5 T, which was shown to be due to competing spin configurations.36 Also the low-temperature field-dependence of the magnetization (0–24 Tesla, 0.42 K) of I is identical to that of the previously described \{Mo_{36}Fe_{30}\}^{9a,b} (see Acknowledgments).

It has been demonstrated that it is possible to ‘use’ pentagonal ‘building units’ which play, geometrically speaking, in spherical molecules the same role as in spherical constructions in general. In the present context they can be used as a glue for trapping 30 magnetic centers \(M\) (e.g. \(M = V^{IV}O_5^-, Fe^{III}\)) to form the icosidodecahedron \{M\}_{36}. As the resulting frustrated magnetic systems share topological features with the highly celebrated Kagomé lattice14 (see Fig. 1) these types of studies are expected to provide a deeper understanding of basic aspects of geometrical/magnetic frustration and the related role of the spin sizes of the metal centers.4a,14 Correspondingly, the method will be extended by us to other magnetic centers in the future, too. The most important aspect of the work refers to the possibility to encapsulate quantum dots like polyoxometalates existing with different electron distributions in host systems. In this context it is important to note that there seems to be a general tendency for practically neutral spherical molybdenum oxide based capsules—but positively charged without the organic ligands—to have a high affinity for the (integration of) polyoxomolybdates.15 Interestingly, metal oxide cluster encapsulation is also possible in virus cavities (see: Viral Cage Directed Synthesis of Nanoclusters, chapter 10.9, p. 482 in ref. 16).

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**Fig. 1** The unique Archimedean solid: the FeIII centers (small colored spheres) of the \{(MoVI)MoVI\}_{12}\{Fe^{III}\}_{36} type Keplerate 1a (but also of the corresponding \{OV\}_{36} system) span the shown icosidodecahedron which is unique among the Archimedean solids.† (Note: in the Kagomé lattice, every vertex has the same surroundings in the sense that the polygons meeting at each vertex are in sequence: pentagon (hexagon in the Kagomé case), triangle, pentagon (hexagon), triangle; see ref. 7a, p. 13.) There are three groups (‘sublattices’) of 10 spins (red, blue, green, respectively), with all spins of a sublattice having parallel spin vectors, while nearest-neighbor spin vectors (three highlighted) differ in angular orientation by 120° (see text and ref. 9a).

**Fig. 2** Structure of 1a emphasizing the MoVI positions of the encapsulated hexamolybdate ion within the metal oxide skeleton of the capsule 1a (Mo blue, O red spheres) and the linkers in the form of FeO₆ octahedra (yellow).
Notes and references
† In this context it is worthwhile to mention that different linkers in the Kepkelets with approximately icoshedral symmetry span (distorted) Archimedean solids:13b in the case of dinuclear metal linkers a (distorted) truncated icoshedron, [M]120 and in the case of mononuclear linkers the unique icostidodecahedron [M]30 with a highly “isotropic” surface (of importance for the magnetic behavior) due to the fact that all edges and dihedral angles are equal (Fig. 1). In the [M]120 situation, a network of corner-shared triangles exists on the sphere surface (Fig. 1), which can cause in special spin situations like in the (FeIII)30 case a unique magnetic frustration.8,9 These [FeIII]30 type clusters show furthermore a novel type of assembly process in solution.

§ Elemental analysis (% calc.: Mo 40.26, C 2.58, H 2.62, Na 0.48%. Found: Mo 39.0, C 2.6, H 2.8, Na 0.5%. (The (not given) ESCA data obtained from Prof. M. Neumann (Osnabrück) correspond to the theoretical Mo to Fe ratio. There is a small error limit with respect to the charge/number of acetates.)

Characteristic IR bands: νcm⁻¹ (KBr pellet): 1622m (v(\text{H}_2\text{O})), 1451m (v(COO)), 1412w (v(COO)), 1322m (v(COO)), 1209m (v(COO)), 955m (v(Mo=O)), 854m (v(Mo=O)), 838s, 518m, 451w, 370m, 238 w-m.

Crystallographic data in CIF or other electronic format see DOI: 10.1039/b604977j. The {Mo6} type clusters exhibit the features of hexamolybdate.}

7. (a) M. O’Keeffe and B. G. Hyde, Cluster Structures, I Patterns and Symmetry, Mineralogical Society of America, Washington, D.C., 1996;
15. A. Muller, et al., unpublished work.
17. More details about this effect will be published in due course.
Extending the \([\text{Mo}_7\text{Cr}_{30}]_{12}\text{M}_{30}\) Capsule Keplerate Sequence: A \([\text{Cr}_{30}]\) Cluster of \(S = \frac{3}{2}\) Metal Centers with a \([\text{Na(H}_2\text{O})_{12}]\) Encapsulate**

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In memory of Abhilash Goplan Janardhan

In previous work it was shown that molybdenum oxide based spherical hollow clusters of the type \([\text{(Mo)Mo}_5]_{12}[\text{linker}]_{30}\)—which, because of their structural features, are also called Keplerates—can be directly synthesized by the addition of linkers, such as \(\text{V}^{\text{IV}}\text{O}^{3+}\) and \(\text{Fe}^{\text{III}}\), to a dynamic library containing (virtual) pentagonal units in solution\([1,2]\) (for general information see also refs. \([3–5]\)). In the \([\text{linker}]_{30}\) situation, a network of corner-sharing triangles exists on the sphere surface,\([1,2a]\) and this gives rise, as in the \([\text{Mo}_7\text{Fe}_{30}]\) situation, to a network of corner-sharing triangles exist on the sphere surface,\([1,2a]\) and this gives rise, as in the \([\text{Mo}_3\text{Fe}_{20}]\) case, to a geometrically frustrated magnetic system which has received a lot of attention.\([4d–f,6]\) In fact, these zero-dimensional materials share topological features with the planar Kagomé lattices which also support geometrical frustration\([7]\) and are of interest for materials science.\([7a]\) The exploration of distinct yet related spin systems, which is an important aspect of the present work, is expected to provide a deeper understanding of basic aspects of geometrical frustration and especially the role played by varying the spin quantum number of the metal centers. Herein we report the synthesis (and new perspectives for the general synthetic principles leading to these types of capsules) of the spherical, hollow \([\text{(Mo)Mo}_5]_{12}[\text{Cr}^{\text{III}}]_{30}\) type cluster \(1\) which contains an interesting encapsulated \([\text{Na(H}_2\text{O})_{12}]\) entity. This \([\text{Mo}_7\text{Cr}_{30}]\) cluster system exhibits, like the \([\text{Mo}_7\text{Fe}_{30}]\) and \([\text{Mo}_3\text{V}_{20}]\) systems, the unique Archimedean solid \([\text{Mo}_8]\) icosidodecahedron (see below and references \([1a,2a]\)).

If an acidified aqueous solution of sodium molybdate containing a rather high concentration of acetic acid is treated with chromium(III) chloride, green crystals of \(1\) precipitate after some days (the basic differences to the preparation of the \(\text{V}_8\) and \(\text{Fe}_{30}\) systems are outlined in the Experimental Section). Compound \(1\), which crystallizes in the space group \(R3\), was characterized by elemental analysis, thermogravimetry (to determine the actual water content based on the release of crystal water), spectroscopy (IR, Raman, UV/Vis), and single-crystal X-ray structure analysis.\([8a]\)

\[\left[\text{Na(H}_2\text{O})_{12}\right] \leq \left[\text{Mo}^{\text{VII}}\text{Cr}^{\text{III}}\text{O}_{25}(\text{CH}_3\text{COO})_{19}\text{(H}_2\text{O})_{19}\right]\]

ca. 120H_2O ≡ 1a - ca. 120H_2O 1

The cluster \(1\) is of the expected spherical \([\text{(Mo)Mo}_5]_{12}-[\text{linker}]_{30}\) (Linker = \(\text{Cr}^{\text{III}}\)) type (Figure 1). Correspondingly, the heptacoordinate \(\text{Mo}\) centers of the 12 pentagonal \([\text{(Mo)Mo}_5]\) units describe an icosahedron, and the 30 \(\text{Cr}^{\text{III}}\) linkers the quasiregular icosidodecahedron. The latter Archimedean solid is a hybrid of the icosahedron and the dodecahedron and contains 20 triangular and 12 pentagonal faces. It comprises not only equivalent vertices but also equivalent edges and equal dihedral angles (see also refs. \([1,2]\)). The 19 acetate ligands in \(1\), which are mainly disordered, are located inside the cavity and coordinate in a bidentate fashion to the metal centers, preferentially bridging \(\text{Cr}\) and \(\text{Mo}\) sites. In the \([\text{(Mo)Mo}_5]_{12}[\text{Cr}^{\text{III}}]_{30}\) cluster, the 30 \(\text{Cr}^{\text{III}}\) ions have their typical sixfold coordination, that is, with two bonds to the small ligands (either to two \(\text{H}_2\text{O}\) ligands or to one external \(\text{H}_2\text{O}\) and one to an internal acetate) and four bonds of the \(\text{Cr}^{\text{III}}\) to \(\text{Mo}\) units (Figure 1). This scenario can also be described as 30 \(\text{Cr}^{\text{III}}\) centers trapped in a matrix formed by the 12 pentagonal \([\text{(Mo)Mo}_5]\) units (Figure 1). This scenario can also be described as 30 \(\text{Cr}^{\text{III}}\) centers trapped in a matrix formed by the 12 pentagonal \([\text{(Mo)Mo}_5]\) units. The \(\text{Cr}^{\text{III}}\) metal ions can be replaced by others, such as \(\text{Fe}^{\text{III}}\) and \(\text{V}^{\text{IV}}\text{O}^{3+}\), in the typical way of coordination chemistry; this situation is of special interest for further syntheses and magnetic studies. The solution and solid-state electronic spectrum of \(1\) shows a band at approximately 15.6 × 10^3 cm\(^{-1}\) which arises from the presence of \(\text{Cr}^{\text{III}}\) and is assigned to the spin-allowed \(^4\text{A}_{2g} \rightarrow ^4\text{T}_{2g}\) transition.

One fascinating structural aspect is that a \([\text{Na(H}_2\text{O})_{12}]\) cluster is encapsulated in the cavity of \(1\). The O atoms of the \([\text{Na(H}_2\text{O})_{12}]\) cluster form an icosahedron, with its approximate \(C_5\) axes coinciding with the \(C_5\) axes of the metal oxide...
The structure of the spherical capsule \([(\text{Mo})\text{Mo}_{12}\text{Cr}^{3+}]_{30}\) was calculated according to a well-established method\(^{[8d]}\) by considering electrostatic energy minimization using a combination of simulated thermal annealing and Powell's algorithms (see ref. \([8e]\)). The result obtained corresponds to the minimum-energy structure when the coordinates of the \(\text{Na}^+\) ion will be published in a later paper.

Figure 2 shows the magnetic susceptibility temperature product \(\chi T\) and \(\chi^{-1}\) as functions of temperature. The \(\chi T\) value is 47.17 cm\(^3\)Kmol\(^{-1}\) at \(T = 300\) K decreasing with decreasing temperature, reaching 3.97 cm\(^3\)Kmol\(^{-1}\) at 1.8 K. The Curie–Weiss plot (Figure 3, bottom) is essentially a straight line above \(T = 100\) K, and a fit to the Curie–Weiss law yields a Curie constant of \(54.8 \pm 0.1\) cm\(^3\)Kmol\(^{-1}\) and a Weiss temperature of \(\theta = -50 \pm 0.1\) K. The magnetization as a function of field at \(T = 1.8\) K (not shown) increases rapidly with increasing field slowing down above 2\(T\), but it does not saturate. Preliminary X-band EPR measurements (not shown) show a broad line at high temperatures, which decreases on lowering the temperature and includes an evolving sharp feature. From these data the value \(g = 1.966\) was obtained.

Both the decrease of the \(\chi T\) value with decreasing temperature and the substantial and negative Weiss temperature show that the predominant exchange interactions in \(1\) are antiferromagnetic which agrees with the fact that the room-temperature \(\chi T\) value is below that for 30 uncoupled spins 3/2 with \(g = 1.966\) (54.35 cm\(^3\)Kmol\(^{-1}\)). To quantify the magnetic behavior we performed quantum Monte Carlo (QMC) calculations\(^{[9,10]}\) of the weak-field susceptibility by adopting a Heisenberg model with isotropic exchange interaction of the form \(-J \mathbf{S}_i \cdot \mathbf{S}_j\) for each pair of spins 3/2 on nearest-neighbor sites \(i\) and \(j\) of an icosidodecahedron.

The values of both \(J\) and \(g\) were optimized to fit the experimental data. The comparison between experimental and theoretical data was restricted to temperatures \(T > 30\) K, because below that temperature the QMC method is not reliable for the icosidodecahedron structure owing to spin-frustration effects.\(^{[9]}\) The best fit was achieved for \(J/k_B = -8.7 \pm 0.2\) K and \(g = 1.96 \pm 0.01\) (Figure 3, bottom). Note that also the weak upturn in \(\chi^{-1}\) with decreasing \(T\) around 50 K is reproduced well by the calculations. In spite of the long superexchange pathway (Cr\(^{3+}\)-O-Mo\(^{5+}\)-O-Cr\(^{3+}\)) the interaction is quite strong. Literature values for exchange couplings along such a pathway range from about 1 K to about 21 K.\(^{[11]}\) The strength of the interaction is determined by the overlap of the Cr d orbitals with the empty d orbitals of the Mo\(^{5+}\) ion.
The moderate $J$ value and the spin quantum number $3/2$ for the individual Cr$^{III}$ centers makes compound 1 an interesting and experimentally convenient system for future physical, theoretical, and experimental studies and especially for the comparison of its magnetic properties with those of the corresponding Fe$^{III}$[12b,13,12a] and V$^{IV}$[11] clusters (spins 5/2 and 1/2, respectively) as well as with the spin-related Kagomé systems.[7] Of special interest is the possibility of observing a narrow minimum in the differential susceptibility $dM/dH$ for $H \approx 20$ T (one-third of the saturation field) as has been seen in the Fe$^{III}$ complex.[8c] This minimum in $dM/dH$ is associated with polytopes assembled from corner-sharing triangles and the Kagomé lattice. In addition, because of its convenient spin-state transition energies (see below), 1a is excellently suited for experimental investigation (by inelastic neutron scattering) of the validity of the rotational band model.[6e] These factors and the weak exchange interaction are also responsible for the fact that quantum steps in the magnetization versus field have not been observed for accessible temperatures in contrast to other antiferromagnetically coupled systems. For $M = V^{IV}$, where the spin is small and delocalized, the system is called a “Quantum Kepler-like”.[14a] The electron delocalization creates a very strong spin coupling which has been shown by QMC calculations to be in the range of $|J|/k_B \approx -245$ K.[14a] This spin coupling causes the excited spin states to be at very high energies, which makes, for example, spin-state crossings inaccessible in conventional laboratory magnetic fields. The spin system of 1a is in between those of the other two compounds which results in correspondingly intermediate magnetic properties.

It should be possible to synthesize new related compounds with interesting and even new types of magnetic properties. These spherical cluster systems have other unique properties, as well. For example, the [Fe$_{30}$] and [Cr$_{30}$] clusters are “nanoacids” and show an unprecedented type of assembly processes leading to vesicles the size of which can be tuned by changing the pH value, a long-term goal in nanoscience. Future synthetic strategies will be determined by the general characteristics of the spherical clusters which can be compared with “coordination polymers” with spherical periodicity.[13] Coordination polymers are considered as being based on metal ions and (organic) ligand building blocks assembling to infinite one-, two-, and three-dimensional networks, the properties of which depend in a characteristic way on the dimensionality.[13] In our spherical clusters we can refer to metal–ligand interactions between the pentagonal units (the ligands) [(Mo)Mo$_5$O$_2$(H$_2$O)$_6$]$^{6-}$ (present in a dynamical library)[11a] and the metal ions. These interactions are comparable to those between lacunary polyoxometalates and metal cations.[14a]

The option of the planned syntheses of molecular spheres based on a careful selection of appropriate pentagonal ligands and metal ions with their specific coordination behavior[14a] allows for the extension of this work to other 3d metal cations which can also lead to mixed-metal as well as mixed-valence clusters with different types of magnetic properties (see for instance ref. [14b]). The preparation of mixed-metal/mixed-valence clusters would correspond to symmetry breaking for the spherical systems which can also be brought about by choosing special cavity-internal ligands attached to the metal centers.[14] A spherical product is formed preferentially if the following condition is fulfilled: ligand-type pentagonal units with appropriate bonding sites are present, as are transition-metal cations which can coordinate to the pentagonal units (that is, assemble around them) maintaining the fivfold symmetry (see Figure 1). In the clusters discussed herein, the repeating unit is [(Mo)Mo$_5$M$_{30}$] ($M = Cr$), in the larger spherical system [(Mo$^{VI}$)Mo$^{VI}$_$5$Si$_3$(Mo$^{V}$_$2$)$_{30}$] which contains binuclear linkers the repeating unit is [(Mo)Mo$_5$Mo]$_{30}$.[15b] The repeating units occur 12 times in the spherical clusters corresponding to the icosahedral symmetry.
Experimental Section

CrCl$_3·$6H$_2$O (8.6 g, 32.27 mmol) was added to a solution of Na$_2$Mo$_{72}$Cr$_{30}$C$_{38}$O$_{516}$H$_{509}$: C 2.58, H 2.89, Na 0.13; found: C 2.6, H 2.5, Na 0.2%. (A small error for the number of Na$^+$ ion.)

Magnetic susceptibility and magnetization measurements were performed on a powder sample of magnetic susceptibility were corrected for diamagnetic contributions.

In the field of quantum chemistry, magnetic susceptibility is a key parameter. The study of magnetic properties is crucial in understanding the behavior of materials at the quantum level. This is especially important in the context of quantum technologies, where precise control of magnetic interactions is necessary for the development of advanced devices. The quantum chemistry approach allows for the calculation of these properties, providing deeper insights into the material's behavior.

Crystals precipitated from the filtrate after 5 days, were collected by filtration. Green crystals were obtained.

Reflex for 45 min. The resulting green solution was cooled to room temperature.

The effectiveness of this technique for magnetic molecules has been reported to improve the efficiency of the magnetic behavior measurement. The improved technique leads to more accurate results, which are essential for the development of new magnetic materials.

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a) The present situation is formally comparable to the “poly-
association” leading, for example, to the impressive helices of J.
M. Lehn. In both cases, polytopic ligands—either preorganized
in the form of poly(bipyridine) ligand strands in the helices or
existing as pentagonal units in a dynamic library in the spherical
systems—interact with a larger number of directing metal
cations showing the expected coordination type; in the former
cases the copper (I) ions enforce the tetrahedral coordination
geometry and in the latter cases the Fe(II) ions the octahedral (see
J. M. Lehn, Supramolecular Chemistry: Concepts and Perspec-
tives, Wiley-VCH, Weinheim, 1995); b) B. Botar, P. Kögerler, A.
Müller, R. Garcia-Serres, C. L. Hill, Chem. Commun. 2005,
5621–5623; see also: A. Müller, B. Botar, H. Bögge, P. Kögerler,


226–231; see also: J. Glerup, A. Hazell, K. Michelsen, H. Weihe,

a) The \( S = 5/2 \) (Fe(II)) analogue of complex \( [\text{Fe}_6\text{Mo}_6\text{O}_{24}]^{3-} \) has some
very intriguing properties which has led to many experiment-
al and theoretical studies. Briefly, it is well described by a
simple picture of geometrical frustration whereby the \( S = 0 \)
ground state corresponds to a spin topology of three coplanar
spin sublattice vectors of length 25, each spin sublattice being
associated with ten parallel \( S = 5/2 \) ion spins, with an angular
separation of 120 degrees between the individual sublattice
vectors. In the low-temperature regime, with increasing applied
magnetic field, \( H \), the three sublattice vectors cant and these
may be pictured as a “uniformly folding umbrella”; the
projections of the sublattice vectors in the plane normal to the
external field maintain a frustration angle of 120 degrees and the
magnetization grows linearly with \( H \) and ultimately saturates at
a field \( H_{sat} \approx 17.6 \) T. Additionally, at low, nonzero temperatures
the differential susceptibility \( dM/dH \) versus \( H \) shows a character-
istic minimum at \( H_{sat}^2/3 \) which has been explained in terms
of the competition between two spin configurations (the canted
frustrated phase and a magnetically stiff configuration). Similar
behavior can be expected for compound 1, where, using our
result \( J/k_B = -8.7 \) K, we estimate \( H_{sat} \approx 60 \) T. A drawback of the
Fe(II) analogue is that owing to the weak superexchange
interaction of \( J/k_B = -1.57 \) K, the energy levels are very closely
spaced, and thus the individual spin-state crossings and tran-
sitions have not been observed in experiments. In unpublished
work by the same authors.

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which it was mentioned that in special cases parallels can exist
between infinite arrays of coordination polymers and discrete
supramolecules, as for example, Fujita’s \([\text{M}_n(\text{tpy})_n]\) (tpy corre-
sponds to a complex triazine ligand), Fujita, D. Oguro, M.
469–471. Interestingly, the pentagonal units discussed herein
also occur in network structures (see refs. 22a,b of ref. [5b]).

[14] a) The present situation is formally comparable to the “poly-
association” leading, for example, to the impressive helices of J.
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Unique Properties of Mo$_{72}$Fe$_{30}$ Cluster in Solution

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Giant molecular spheres with the general formula [(pentagon)$_{12}$(linker)$_{30}$] or [Mo(Mo)$_{3}$]$_{12}$(Linker)$_{30}$, due to their structural features also called Keplerates [1-3], with variable size can be generated by linking the fundamental pentagonal {MoMo$_{3}$} building block with various linkers, which can be of the dinuclear ({$\text{Mo}^{V}_{2}\text{O}_{4}\text{ligand}$})$_{n}^{n^{+}}$ (e.g. ligand = HCOO$^{-}$, CH$_3$COO$^{-}$, SO$_{4}^{2-}$, H$_2$PO$_2^{-}$, PO$_4^{3-}$)) or mononuclear ({$\text{Fe}^{III}(\text{H}_{2}\text{O})$})$_{3}^{3^{+}}$, {$\text{Cr}^{III}(\text{H}_{2}\text{O})$})$_{3}^{3^{+}}$, {$\text{Mo}^{V}\text{O(H}_{2}\text{O})$})$_{3}^{3^{+}}$, {$\text{V}^{IV}\text{O(H}_{2}\text{O})$})$_{2}^{2^{+}}$) type.

Here we begin to study the replacement with D$_2$O of the crystal water and of the H$_2$O ligands attached to the Fe$^{III}$ and Mo$^{VI}$ centres of the cluster with the stoichiometry: [Mo$_{72}$Fe$_{30}$O$_{252}$CH$_3$COO)$_{12}$[Mo$_2$O$_7$(H$_2$O)]$_2$ [H$_2$Mo$_2$O$_8$(H$_2$O)] (H$_2$O)$_{91}$] · ca. 150 H$_2$O (Mo$_{72}$Fe$_{30}$), which is part this class of spherical systems. The 72 Mo$^{VI}$O$_n$ polyhedra are organized into twelve pentagonal {Mo(Mo)$_{3}$} type units (comprising a central MoO$_7$ pentagonal bipyramid linked via edges to five MoO$_6$ octahedra). The Mo=O bonds are preferably directed outwards (ca. 80%), but according to a disorder also approximately 20% Mo(OH$_2$) groups are directed outwards. The twelve pentagonal units are connected by 30 Fe(OH$_2$) octahedra acting as linkers. Each of these linkers exposes a single terminal Fe(OH$_2$) outwards. Inside the near-spherical capsule there are twelve additional water molecules bound to the Fe$^{III}$ centers. The remaining 18 octahedral coordination sites on the Fe$^{III}$ linkers are bridges to molybdate (two of [Mo$_2$O$_7$(H$_2$O)] type and one of [H$_2$Mo$_2$O$_8$(H$_2$O)] type) or to acetate ligands.
The Mo$_{72}$Fe$_{30}$ clusters exist as discrete, almost neutral molecules in aqueous solution at pH < 2.9, but get deprotonated and self-associate into single-layer blackberry-type structures at higher pH while the assembly process, i.e. the size of the final species can be controlled by the pH values. The average hydrodynamic radius ($R_h$) of the aggregates decreases monotonically with increasing number of charges on the Mo$_{72}$Fe$_{30}$ macroanions (from ~45 nm at pH~3.0 to ~15 nm at pH~6.6), as studied by laser light scattering and TEM techniques.

Standard $^{17}$O-NMR investigations on 18.65mg/ml aqueous solutions of Mo$_{72}$Fe$_{30}$, limited to 2<pH<3 because of the cluster stability as it dissociates slowly at pH<2 and forms larger aggregates at pH>3, showed that the rates of exchange of waters bound to the Fe$^{III}$(OH$_2$) functional groups on the Mo$_{72}$Fe$_{30}$ cluster are $\sim10^4$ more rapid than for the simple aqua ion, Fe(H$_2$O)$_6^{3+}$, but slightly slower than the Fe$^{III}$-EDTA and its derivatives, but also that small changes in pH near the pKa of the sites have little effect on the rates of exchange of bound waters [5]. That these molecules remain intact in the aqueous phase is well established through characteristic Raman spectrum (due to their high symmetry the free clusters exhibit only a few well-defined lines).

Two strategies were employed in order to replace the H$_2$O present by D$_2$O in the Mo$_{72}$Fe$_{30}$-type cluster‡:

A) Recrystallization of Mo$_{72}$Fe$_{30}$ from heavy water

0.5 g Mo$_{72}$Fe$_{30}$ (prepared according to the method of Müller et al. [4]) were dissolved in 15 ml D$_2$O, leading to a 33.33mg/ml solution of Mo$_{72}$Fe$_{30}$. The solution was heated at $\sim$ 60°C, under argon 24h, 48h and 72h respectively. The resulting solution was cooled under 0°C and the

‡ These investigations were done for quasi elastic neutron scattering experiments.
formed precipitate was filtered off and dried under argon. The identity of \( \text{Mo}_{72}\text{Fe}_{30}-\text{D}_2\text{O} \), where most of H\(_2\text{O}\) was replaced by D\(_2\text{O}\), was established by IR.

Figures 2-4 show the IR spectra of the \( \text{Mo}_{72}\text{Fe}_{30}-\text{D}_2\text{O} \) samples precipitated from the 33.33 mg/ml heavy water solutions after heating these solutions for 24h, 48h and 72h respectively at ~60\(^\circ\)C. In this case we can notice a decrease in the intensity of the bands at 3200 cm\(^{-1}\) and 1616 cm\(^{-1}\) and the appearance of the bands at ~ 2200 cm\(^{-1}\) and ~1194 cm\(^{-1}\) due to the replacement of H\(_2\text{O}\) by D\(_2\text{O}\).

Stirring for 24h at 80\(^\circ\)C leads to the decomposition of the cluster and subsequent precipitation of an "iron-polymolybdate".


Figure 1. IR spectrum (diamond ATR) of $[\text{Mo}_{72}\text{Fe}_{30}\text{O}_{252(\text{CH}_3\text{COO})_{12}[\text{Mo}_2\text{O}_7(\text{H}_2\text{O})]_2}}$ $[\text{H}_2\text{Mo}_2\text{O}_8(\text{H}_2\text{O})](\text{H}_2\text{O})_{91}] \cdot \text{ca. 150 H}_2\text{O} \equiv \text{Mo}_{72}\text{Fe}_{30}$. 
Figure 2. IR spectrum (diamond ATR) of "Mo$_{72}$Fe$_{30}$-D$_2$O" precipitated from a heavy water solution after stirring for 24h at ~ 60°C.
Figure 3. IR spectrum (diamond ATR) of "Mo$_{72}$Fe$_{30}$-D$_2$O" precipitated from a heavy water solution after stirring for 48h at ~ 60°C.
Figure 4. IR spectrum (diamond ATR) of "Mo$_{72}$Fe$_{30}$-D$_2$O" precipitated from a heavy water solution after stirring for 72h at ~ 60°C.
B) Starting with deuterated precursors

0.3 g FeCl₃ (1.85 mmol) were added to a solution of 1.78 g MoO₃ (12.36 mmol) (from Aldrich) in a mixture of 2.87 g NaOD (40% in D₂O, 99 atom %D) (from Aldrich), 35 g D₂O (99.8 atom %D, from Acros Organics) and 16.06 g CH₃COOD 100% (98 atom %D, from Acros Organics). After acidification with 1.63 g DCl (35% in D₂O, 99 atom %D) (from Aldrich) the solution (pH≈2) was heated to 90-95°C, stirred shortly and cooled to room temperature while after 6h the yellow microcrystalline precipitate was filtered off (under argon). The yellow crystals of the pure compound precipitated from the filtrate (kept under argon) after 5 days, were filtered off and dried under argon.

The attempts to prepare the "Mo₇₂Fe₃₀-D₂O" cluster starting from deuterated precursors lead to the precipitation of compounds in which the H₂O could be replaced only to a small extent (figure 5, 6, 7 and 8). The source of H₂O could be FeCl₃. The precipitation of the iron ball with water ligands could be due to the fact this is less soluble than the heavy water enriched iron ball or to an unusual type of isotope effect.
Figure 5. IR spectrum in nujol of "Mo$_{72}$Fe$_{30}$-H$_2$O-D$_2$O" prepared starting from deuterated precursors.
Figure 6. IR spectrum (diamond ATR) of "Mo$_{72}$Fe$_{30}$-H$_2$O-D$_2$O" prepared starting from deuterated precursors.
Figure 7. IR spectrum in nujol of "Mo72Fe30-H2O" prepared starting from deuterated precursors.
Figure 8. IR spectrum (diamond ATR) of $\text{Mo}_{72}\text{Fe}_{30}\cdot\text{H}_2\text{O}$ prepared starting from deuterated precursors.
Metal-Oxide-Based Nucleation Process under Confined Conditions: Two Mixed-Valence V₆-Type Aggregates Closing the W₄₈ Wheel-Type Cluster Cavities

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Dedicated to Professor C. N. R. Rao

Understanding assembly processes based on the linking of simple XY₄-type polyhedra has been of tremendous interdisciplinary interest since the time of Pauling’s original statements referring especially to the silicates.[1a] It seems to be of importance to distinguish between self-assembly and directed-assembly, terms used for elementary processes that probably occurred on the primordial earth.[1b] In bulk solution one can mostly refer to self-assembly, but under confined conditions directed-assembly should be predominant because of the probable influence of the “environment”. In nearly all of these cases the elementary processes, especially the first related steps, are not known. The most studied system in this context is that of the polyoxometalates (POMs), in which MO₆-type species appear finally as polyhedra linked (mainly) through corners and edges.[2] Regarding directed-assembly, two reports should be mentioned: a study on a giant wheel-type species in which molecular growth is observed inside the cavity[3] and the discovery of an unprecedented stepwise molecular cascade type growth process based on nucleophilic/electrophilic interactions.[4] Herein we report a process that occurs within the highly negatively charged wheel-shaped [P,WₓO₄₈]₆₋₂⁻ species[5] in which the formation of two mixed-valence [Vₓ⁴⁺₁Vₒ³⁻(H₂O)₉₂]⁺ assemblies with linked tetrahedra and octahedra is directed by the internal surface, thereby closing the cluster cavity.[6] This type of process allows generation of a variety of magnetically interesting mixed-valence species by studying systematically the influence of the environment and internal wheel surface under variable redox conditions.

Reaction of a solution of 1 with vanadyl sulfate in aqueous medium yields crystals of 2 after some time. Compound 2, which crystallizes in the triclinic space group P1̅, was characterized by elemental analysis, thermogravimetry to determine the actual crystal water content, spectroscopic methods (IR, resonance Raman, UV/Vis, EPR, MAS-NMR), single-crystal X-ray structure analysis (including bond-valence-sum (BVS) calculations),[7] and susceptibility measurements. A view of the structure of 2 in a mixed polyhedral and ball-and-stick representation is shown in Figure 1.

K₃₋₄Li₃₋₄H₃₋₄Kₓ₋₄⊂ {P₆W₉₀O₃₄₄₄} · x H₂O (x ≈ 92) (1)[8]

Na₂₋₄K₃₋₄H₃₋₄⊂ {P₆W₉₀O₃₄₄₄} [Vₓ₋₄⁻¹⁰⁻₉⁻⁴(H₂O)₉₂] · x H₂O (x ≈ 80) (2)

The “host” anion, [K₅₋₄P₆W₉₀O₃₄₄₄]₁²⁻ (1a) with fourfold (D₄h) symmetry is derived from the linkage of four [P₆W₉₀O₃₄₄₄] lacunary fragments of the Wells–Dawson anion [P₆W₉₀O₃₄₄₄][]⁻; it encloses eight potassium cations in the positions shown in Figure 2a. The cations at the 3-, 6-, 9-, and 12-o’clock positions in the figure are disordered over two sites above and below the equatorial mirror plane; the remainder lie in that plane. Potassium cations occupy analogous positions within the new host–guest complex 2a but they are now sandwiched between the two cyclic capping groups...
[\{V^{VV},V^{IV},O_{12}(H_{2}O)_{2}\}^{4+} (3a), each with two octahedral V^{IV} and four tetrahedral V^{V} centers (Figure 2a). 3a can be considered formally as an interesting polycation exhibiting two different types of entities, namely, \{O=V(H_{2}O)\}^{2+}\[8] and V_{6}O_{12},\[8] which occur under well-defined conditions, the latter in the gas phase, the former in hydrated form in aqueous solution. The formation of 3a is based on oxidation of V^{IV} by air and does not occur in absence of oxygen. As is evident from the figure the V_{6} caps force a rhombic distortion of the square \{P_{8}W_{48}\} perimeter observed in 1a.\[9] The internal angles of the rhombus in 2a (of lines extended from the P···P vectors in each \{P_{8}W_{48}\} unit) are close to 80° and 100°. The overall anion dimensions (measured by the separation between the oxygen atom “hinges” linking the corner-shared \{WO_{6}\} octahedra at 6- and 12-o’clock positions in Figure 2a versus those at 3- and 9-o’clock positions) are 15.27 and 17.94 Å. As shown in Figure 2b the short terminal V–O bonds of the four VO^{2+} centers are directed towards the interior of the structure and point towards the centroids of groups of three encapsulated potassium cations. The orientation of the VO^{2+} groups places four weakly bound water molecules (V–OH\textsubscript{2} = 2.23 and 2.27 Å) on the surface of the anion, and this leads to the possibility of attaching (albeit weakly) other ligands at these positions. The structure is a rare example of a polyoxometalate in which M–O\textsubscript{terminal} vectors point towards the interior of the anion.\[10] Presumably space limitations (the V···V separation is 6.21 Å) preclude the enclosure of two water molecules.

Magnetic measurements were performed and Figure 3 shows the paramagnetic susceptibility–temperature product and the inverse of the susceptibility recorded on a fresh powder sample of 2 at an applied field of 1 T, both as a function of temperature. The high-temperature \(\chi T\) value of 1.34 cm\(^3\)Kmol\(^{-1}\) corresponds to the presence of 3.8 V^{IV} centers, which is in good agreement with the four [V^{IV}O\textsubscript{12}(H\textsubscript{2}O)]\(^{2+}\) units found in the crystallographic studies. From this result it can be immediately concluded that there are no additional [V^{IV}O\textsubscript{12}]\(^{2+}\) units in the lattice. The Curie–Weiss plot of the inverse of the susceptibility as a function of temperature (Figure 3) gives a Curie constant of 1.35 cm\(^3\)Kmol\(^{-1}\), which is in excellent agreement with the room-temperature value of \(\chi T\). The small Weiss temperature of \(\theta = -1.8 ± 0.2\) K shows that the ions are only very weakly interacting, which is
not surprising in view of the orientation and separation of the magnetic ("d\text{xy}" orbitals of the V\text{IV} centers. The decrease of the value of \(\chi T\) at lower temperatures is not an effect of magnetization saturation because the 0.1 T applied field susceptibility curve does not differ from that measured at an applied field of 1 T. This behavior is probably a consequence of small inter- or intramolecular exchange interactions.

The X-band EPR spectrum recorded on a powder sample exhibits a broad line with \(g = 1.95\) at room temperature, but shows a hyperfine splitting from \(I = 7/2\) V\text{IV} nuclei of the order of \(A = 200\) MHz (67 x 10\(^{-4}\) cm\(^{-1}\)) at 4.1 K (Figure 4). The magnitude of \(A\) and the pattern of hyperfine features are consistent with a triplet state arising from interactions between pairs of V\text{IV} centers rather than a single VO\text{II} center, although the “forbidden” half-field line was not observed. Raising the temperature immediately leads to line broadening and disappearance of the hyperfine structure as a result of an increase in spin-lattice relaxation. Owing to the broadness of the spectrum even at 4.1 K it is not possible to obtain an unambiguous set of \(g\) and \(A\) parameters by simulation. The simulation depicted in Figure 4 corresponds to the parameters shown in Table 1, but these are by no means optimal. Nevertheless, the hyperfine parameters are similar to those reported for the vanadyl tartrate dimers shown in Table 1 and are quite distinct from those of a typical monovanadium(V) polytungstate, also shown in the Table.

It seems plausible to assume that the “eclipsed” V\text{IV} centers (V1 and V4, Figure 2b) account for the triplet state EPR spectrum since the V1···V4A separation is 10.2 Å and would involve an unfavorable V\text{IV}(oct.)-O-V\text{IV}(tet.)-O-V\text{IV}(oct.) superexchange pathway. Superexchange between V1 and V4 involves multiple V\text{IV}(O)-W\text{VI}(O)-W\text{VI}(O)-W\text{VI}(O)-W\text{IV}(O) pathways favored by the “mixed-valence” interaction between V\text{IV} and W\text{VI} (discussed below). Similar exchange involving multiple bond pathways has been demonstrated in [K\text{IV},W\text{VI}(O),O\text{IV}]	ext{II}\text{IV}.

However, the adjacency of the two triplet states within the same anion is almost certainly responsible for the increased spin-lattice relaxation observed. A more detailed single-crystal EPR analysis is warranted for this interesting compound.

According to the Robin–Day classification\(^{[11]}\) 3a is a Class I mixed-valence fragment/cation since the two valence states are clearly distinguished by their different coordination geometries, while the electron density is predominantly localized on the octahedral site, for which the crystal-field stabilization (here applicable only in a formal sense) is larger than that for the tetrahedral site (10\(Dq_{\text{rel}}\) = \(-(4/9)10\(Dq_{\text{rel}}\)). The electronic spectra of Class I mixed-valence compounds are a combination of the spectra of the individual oxidation states with no additional absorption (intervalence) bands. The situation is a little more complicated when the complete anion 2a is considered because 3a is embedded in the polytungstate matrix. In addition to the V\text{IV}V\text{IV} Class I mixed-valence situation, there is also adjacency of V\text{IV} with W\text{VI}. As has been well established for tungstenates(IV) (e.g., [Si\text{IV}(V\text{IV},V\text{IV})\text{IV}])\text{II}, these can be treated as Class II systems with partial electron delocalization from “octahedral” V\text{IV} to “octahedral” W\text{VI}, which results in intensity-enhanced “d–d transitions” of V\text{IV} as well as V\text{IV}–W\text{VI} “intervalence” charge transfer at around 500 nm (see reference [14]). In contrast to the case of the V\text{IV}V\text{IV} situation, the enhancement of the electron-transfer rate for the V\text{IV}W\text{VI} case should also be influenced by the vibrations that change the metal–metal distances and therefore modulate the transfer integrals (see the Supporting Information). The UV/Vis spectrum of a freshly prepared solution of 2 (Figure S1 in the Supporting Information) shows just such absorption features\(^{[15]}\).

The slow decomposition of 2 in solution, together with the presence of four paramagnetic centers, precluded the observation of useful multinuclear NMR spectra. However, solid-state spectra could be recorded (see the Supporting Information).

Apart from the points that refer to general aspects of encapsulation chemistry, another interesting phenomenon was discovered. 3a can be considered to be an interesting mixed-valent polycation that is formed by a nucleation process under confined conditions, and the specific geometric environment influences the type of electron distribution and exchange interactions, which are negligible in the present case. There is no doubt that other nucleation processes can be studied in the [P\text{IV},W\text{VI}] cluster cavity under formally similar conditions such as those related to molybdenum oxide based aggregates, which were observed in larger cavities. The interesting aspect is that a variety of magnetically interesting mixed-valence species can be generated by changing the redox potentials.

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Table 1: Selected EPR g and \(A\) (10\(^{-4}\) cm\(^{-1}\)) parameters.

| Compound                  | \(g_{||}\) | \(g_{\perp}\) | \(A_{||}\) | \(A_{\perp}\) | Reference |
|---------------------------|-----------|-------------|-----------|-------------|-----------|
| 2                         | 6.21      | 1.93        | 1.93      | 75          | 30        | this work |
| VO\text{II}-tartrate dimer| 4.08      | 1.953       | 1.982     | 72.3        | 21.3      | [11]      |
| VO\text{II}-tartrate dimer| 4.35      | 1.950       | 1.984     | 73.2        | 24.6      | [11]      |
| [PV\text{IV},V\text{IV}O\text{IV}]\text{III} | 1.915     | 1.970       | 167.2     | 59.7        | [14d]     |           |
Experimental Section

VOSO₄·5H₂O (0.35 g, 1.38 mmol) was added to a stirred solution of K₃[As₂W₁₈O₆₄]·9H₂O (0.35 g, 0.024 mmol) in freshly prepared 1st Na₂CO₃, COOH buffer (30 mL, pH = 4.4). The resulting solution was heated to 50°C for 4 h and then filtered, and the filtrate was allowed to evaporate at room temperature in an open plastic bottle. Lactam flask, during which period partial aerial oxidation of vanadium occurred. After 1 week the dark purple crystals were filtered, washed with cold water, and dried in air; yield: 0.2 g (54%); elemental analysis: found: Na 1.8% K 3.8% V 3.9%.

Characteristic IR bands (KBr pellet): 1711 (m), 1528 (m), 1457 (m), 1354 (m), 1064 (m), 963 (s), 785 (s), 682 (s), 559 (s), 467 (w) cm⁻¹; characteristic Raman band (Solid/KBr dilution; 

| \( n_\text{UV} = \text{As}(W-O-V) \), 574 (w), 536 (w), 467 (w) cm⁻¹ |
| \( \text{V}^{\text{III}} \) (IVCT), 680 (sh), 890 (w) (d–d(V IV)). |

Magnetic susceptibility measurements were performed on a powder sample with a Quantum Design MPMS XL7 SQUID magnetometer. The underlying diamagnetic susceptibility was determined by susceptibility measurements on the diamagnetic [P₈W₄O₁₉]³⁻ host ion, because standard procedures with Pascal constants give values that are too low. X-band EPR spectra were recorded on a Bruker AXS SMART diffractometer (three-circle goniometer with 1 K CCD detector, Mo Kα radiation, graphite monochromator; hemisphere data collection in 0.3° scan width in three runs with 606, 435, and 230 frames (\( \epsilon = 0.82, 1.88 \) and 180°) at a detector distance of 5 cm). (SHELXS/L, SADABS from G.M. Sheldrick, University of Göttingen 1997/2001; structure graphics with DIAMOND 2.1 from K. Brandenburg, Crystal Impact GbR, 2001). Further details on the crystal structure investigations may be obtained from the Fachinformationszentrum Karlsruhe, 76344 Eggenstein-Leopoldshafen, Germany (fax: (+49) 7247-808-666; e-mail: crysdatal@fiz-karlsruhe.de), on quoting the depository number CSD-417620.

Although the nonrigidity of the [P₈W₄O₁₉]³⁻ assembly should not have been unexpected, the other reported derivatives retain the essentially “square” conformation.

Other examples include equatorial W centers in \( [\text{AsIII} \text{W}^\text{IV} \text{V}^\text{III}] \) (Y. Jeannin, J. Martin-Frède, J. Am. Chem. Soc. 1981, 103, 1664–1667) and \( [\text{PV} \text{V}^\text{IV}], \text{V}^\text{IV}] \) (C. M. Tourné, G. F. Tourné, T. J. R. Weakley, J. Chem. Soc. Dalton Trans. 1986, 2237–2242). The problem is of basic importance for larger species such as Mo₆O₁₈, Mo₃O₄Fe₄, and Mo₅S₄. In the first two species, the Mo-O bonds are preferably directed outwards (ca. 80%), but according to a disorder also approximately 20% Mo(OH)₄ groups are directed outwards. In the latter ring-type species, the situation is more complicated because of the pressure of other (different) building units. In any case, there is a preference for H₂O ligands on the outer periphery (for compound types, see, for example: A. Müller, S. Roy, J. Mater. Chem. 2005, 15, 4673–4677 and A. Müller, S. Roy, In The Chemistry of Nanomaterials: Synthesis, Properties and Applications (Eds.: C. N. R. Rao, A. Müller, A. K. Cheetham), Wiley-VCH, Weinheim, 2004, 452–475).


The extinction coefficient per V⁴⁺ center (ca. 1900 m⁻¹ cm⁻¹) in 2 is approximately three times greater than that observed in certain tungsotanades (5 P. Harskmal, M. A. Leparulo, M. T. Pope, J. Am. Chem. Soc. 1983, 105, 4286–4292). This is attributed to the larger V-O-W angles (162°–164°) in 2, which favor greater electron delocalization from V⁴⁺ to W⁴⁺.

Metal-Oxide-Based Nucleation Process under Confined Conditions: Two Mixed-Valence V₆-Type Aggregates Closing the W₄₈ Wheel-Type Cluster Cavities

Achim Müller,* Michael T. Pope,* Ana Maria Todea, Hartmut Bögge, Joris van Slageren, Martin Dressel, Pierre Gouzerh, René Thouvenot, Boris Tsukerblat and Aidan Bell

Figure S1. UV-Vis spectrum of a freshly-prepared solution of 2.
Valence localization/delocalization and Intervalance Charge Transfer Transitions


![IR spectrum of 2](image_url)
Phosphorus NMR Spectra

Room temperature $^{31}$P solid-state NMR spectra were recorded at 162 MHz in the single-pulse mode using a Bruker Avance 400 spectrometer equipped with a 4 mm MAS NMR probe. Owing to the small chemical shift anisotropy of the samples, relatively low spinning rates (= 10 kHz) were sufficient to obtain spectra free of spinning side bands. Data were acquired with 10° flip angle of 1 ms length with a repetition time of 1 s. Chemical shifts are given with respect to 85% H$_3$PO$_4$ as external standard.

The $^{31}$P MAS NMR spectrum of 1 (Figure S4) displays three partially overlapping isotropic signals at ca. -7.7, -7.3 and - 7.1 ppm (fwhm ca. 160 Hz, ca. 1 ppm). This indicates the presence of at least three different phosphorus environments compatible
with the X-ray structural analysis showing the presence of four crystallographically-independent phosphorus atoms in the crystal.\[5\] The $^{31}$P MAS NMR spectrum of 2 (Figure S5) displays a very broad isotropic feature centered at $ca.$ 0 ppm (fwhm $ca.$ 2 kHz, $ca.$ 12 ppm) which consists essentially of two overlapping signals with unequal width. According to the centrosymmetrical 2a, there should be four inequivalent phosphorus atoms. Due to the vicinity of the paramagnetic V$^{IV}$ centers the individual signals are broadened with respect to 1 and overlap each other.

Figure S4. $^{31}$P MAS-NMR spectrum of 1
Figure S5. $^{31}$P MAS-NMR spectrum of 2
Nucleation Process in the Cavity of a 48-Tungstophosphate Wheel Resulting in a 16-Metal-Centre Iron Oxide Nanocluster


Abstract: The 16-FeIII-containing 48-tungstophosphate \([\text{P}_8\text{W}_{48}\text{O}_{184}\text{Fe}_{16}(\text{OH})_{28}(\text{H}_2\text{O})_4])^{20-}\) (1) has been synthesised and characterised by IR and ESR spectroscopy, TGA, elemental analyses, electrochemistry and susceptibility measurements. Single-crystal X-ray analyses were carried out on \(\text{Li}_{12}\text{K}_{16}[\text{P}_8\text{W}_{48}\text{O}_{184}\text{Fe}_{16}(\text{OH})_{28}(\text{H}_2\text{O})_4]\cdot6\text{H}_2\text{O}\cdot2\text{KCl}\) (LiK-1), orthorhombic space group \(P2_1/a\), monoclinic space group \(C2/c\), and on the corresponding mixed sodium–potassium salt \(\text{Na}_9\text{K}_1\text{P}_8\text{W}_{48}\text{O}_{184}\text{Fe}_{16}(\text{OH})_{28}(\text{H}_2\text{O})_4]\cdot10\text{H}_2\text{O}\) (NaK-1), monoclinic space group \(C2/c\), which can be regarded as an assembly process under confined geometries. One fascinating aspect is the possibility to model the uptake and release of iron in ferritin. The electrochemical study of 1, which is stable from pH 1 through 7, offers an interesting example of a highly iron-rich cluster. The reduction wave associated with the FeIII centres could not be split in distinct steps independent of the potential scan rate from 2 to 1000 mVs⁻¹; this is in full agreement with the structure showing that all 16 iron centres are equivalent. Polyanion 1 proved to be efficient for the electrocatalytic reduction of NO₂, including nitrate. Magnetic and variable frequency EPR measurements on 1 suggest that the FeIII ions are strongly antiferromagnetically coupled and that the ground state is tentatively spin \(S=2\).
Introduction

Chemistry under confined geometries—and this in a general sense—has attractive aspects which may be related to special topics of surface,[1a] geo-[1b] and especially biosciences.[2,3] One may ask general questions, such as: what is it like for molecules/ions to “live” inside a nanosized molecular container (or generally under constrained/shielded environmental conditions) with respect to the interactions between them? In this context we can refer to two scenarios: 1) such interactions take place (nearly) independent of the cavity-interior shell functionalities (as in a nano-test-tube) or 2) they are influenced by the shell functionalities. Whereas in the first case the situation allows the spectroscopic identification/characterisation of the species under consideration more easily than under bulk conditions, in the second case one can study template-directed syntheses leading to unprecedented nanospecies. Such a process occurs in nature in different types of compartments.[3] In the case of biomineralisation we can refer to the imposition of (biological) directionality on the chemistry of growth processes (vectorial regulation).[3] In the present study we consider templated nucleation processes based on hydrate complexes of FeII (in presence of O2) and FeIII in the cavity of the cyclic 48-tungstophosphate P8W48, leading to an unprecedented 16-metal-centre iron oxide formed by linking FeO6 octahedra. This type of nucleation is based on a breaking of symmetry during the assembly process caused by the template effect of the cavity internal WO groups. The mentioned reaction of FeII in the presence of dioxygen shows an important feature: it played a key role on the early earth leading to iron-banded ores and is—regarding the confinement conditions—the basis for the formation of the iron oxide core of the metal-storage protein ferritin.[4a] In contrast with the present vectorial growth process, we should refer also to the bacterial Mo storage proteins, in which different specific pockets of the protein cavity direct in unique nucleation processes the formation of different polyoxometalates (POMs).[4b] The type of procedure/nucleation process described in this paper, which has several important interdisciplinary aspects, could in principle be extended to POMs with much larger cavities, for example, wheel-shaped polyoxomolybdates of the type Mo170.[4c]

Results and Discussion

Synthesis and structure: Although the P12W48 cluster has been known for more than 20 years,[5] only recently the first examples of metal-containing derivatives have been reported. Pope's group prepared the first lanthanide derivative, \([\text{Ln}_8(\text{H}_2\text{O})_{24}][\text{K}^+\cdot\text{P}_8\text{W}_{48}\text{O}_{184}\text{Fe}_{16}(\text{OH})_{28}]\) \([\text{Ln} = \text{La}, \text{Ce}, \text{Pr}, \text{Nd})\],[6] and Kortz and co-workers isolated the mixed cation salts LiK4[\text{P}_8\text{W}_{48}\text{O}_{184}\text{Fe}_{16}(\text{OH})_{28}]66\text{H}_2\text{O}·2\text{KCl} (\text{LIK-1}) and Na4K2[\text{P}_8\text{W}_{48}\text{O}_{184}\text{Fe}_{16}(\text{OH})_{28}]100\text{H}_2\text{O} (\text{NAK-1}),[7] (see Experimental Section). Polyanion 1 contains an unprecedented \([\text{Fe}_{16}(\text{OH})_{28}\text{(H}_2\text{O})_{4}]^{2+}\) cluster in the cavity of \(\text{P}_8\text{W}_{48}\) with 16 edge- and corner-sharing FeO6 octahedra being grafted to the inner- and corner-sharing FeO6 octahedra being grafted to the inner- and corner-sharing FeO6 octahedra being grafted to the inner- and corner-sharing FeO6 octahedra being grafted to the inner- and corner-sharing FeO6 octahedra being grafted to the inner- and corner-sharing FeO6 octahedra being grafted to the inner- and corner-sharing FeO6 octahedra being grafted to the inner- and corner-sharing FeO6 octahedra being grafted to the inner- and corner-sharing FeO6 octahedra being grafted to the inner- and corner-sharing FeO6 octahedra being grafted to the inner- and corner-sharing FeO6 octahedra being grafted to the inner- and corner-sharing FeO6 octahedra being grafted to the inner- and corner-sharing FeO6 octahedra being grafted to the inner- and corner-sharing FeO6 octahedra being grafted to the inner-

![Figure 1. Front and side view of the structure of 1 emphasising the FeO6 octahedra (brown) in polyhedral representation. Colour code: W (green), O (red), P (pink).](image)

Figure 1. Front and side view of the structure of 1 emphasising the FeO6 octahedra (brown) in polyhedral representation. Colour code: W (green), O (red), P (pink).
it is possible to use a large variety of iron salts as starting materials, ranging from mononuclear iron(II) and iron(III) complexes (the former requires addition of an oxidant) to dinuclear iron(III) carboxylates. These observations support earlier knowledge that POM syntheses in general depend very much on the boundary conditions in the reaction vessel (e.g., pH and solvent).

Polyanion 1 exhibits a highly attractive symmetrical D₆ₐ structure (see Figures 1 and 2). The large cavity (roughly 9 × 9 × 7 = 567 Å³) of the “cyclic template/host” P₈W₁₈O₆₁₀⁻ has been “decorated” with a cationic nanocluster built up by 16 FeO₆ octahedra, resulting in a smaller, central cavity (roughly 6 × 6 × 5 = 180 Å³). The related “Fe₄₈ ring” is composed of eight pairs of structurally equivalent, edge-shared FeO₆ octahedra that are connected to each other through corners. While most of the Fe-O-Fe bridges are monoprotonated, four are diprotonated (presence of H₂O ligands). This can be confirmed by looking at the related bond valence sums (BVS) of these oxygen atoms. For example, the protonated oxygen (with the corresponding BVS values) of the polyanion in the mixed lithium–potassium salt LiK₁₋₁P₈W₁₈O₆₁₀H₂O₂₋₂ is O14F (0.69), O23F (0.71), O13F (1.08), O24F (1.11), O14G (1.17), O23G (1.27), O1FE (1.31), O2FE (1.32), and O4FE (1.34), see Figure 3. The rather low, but “intermediate” (between mono- and diprotonation) BVS values of 0.69 and 0.71 for O14F and O23F, respectively, led us to believe that we are looking at a water and a hydroxo ligand disordered over these two sites. Hence, we should have a total of 28 hydroxo and 4 aqua ligands associated with 1.

These results confirm that we have indeed grafted an unprecedented, cyclic [Fe₆₇(OH)₉₆(H₂O)₄]²⁺ iron nanocluster with hydroxo and aqua ligands inside the cavity of P₈W₁₈O₆₁₀⁻ (see Figures 1 and 2). Selected bond lengths and angles of the [Fe₆₇(OH)₉₆(H₂O)₄]²⁺ unit are shown in Figure 3. The FeO₆ octahedra are only slightly distorted with Fe–O distances ranging from 1.985(12) to 2.153(12) Å.

It is of interest to compare the structure of 1 with P₈W₁₈O₆₁₀⁻ type analogues containing other transition-metal centres. For example, we notice that the grafting mode of the 16 FeIII centres in 1 is different from that of the 20 CuII centres in [Cu₂₀(H₂O)₂₀P₈W₁₈O₆₁₀]⁺⁻. In 1 each of the 16 equivalent FeIII centres is bound to P₈W₁₈O₆₁₀⁻ by a Fe-O(W) and a Fe-O(P) bond, resulting in a tight anchoring of the 16-iron hydroxo core. In the CuP₈W₁₈O₆₁₀⁻, only eight of the 20 CuII ions form two covalent Cu–O(W) bonds each to the P₈W₁₈O₆₁₀⁻ host. Hence, the eight phosphate groups of P₈W₁₈O₆₁₀⁻ are not involved in the binding to the cationic [Cu₂₀(OH)₉₆]²⁺ cluster guest. In fact, 1 is structurally most closely related to Mialane’s Cu₆₇–azide derivative [P₈W₁₈O₆₁₀Cu₂₀(N₃)₆(H₂O)₉₆]²⁺. In the latter, 16 of the 20 CuII ions are bound to the inner rim of P₈W₁₈O₆₁₀⁻ in exactly the same fashion as the FeIII centres in 1. The sites of the remaining four unique, Jahn–Teller distorted CuII ions in Mialane’s POM remain empty in 1. However, we believe that in principle these four sites could be filled in 1 as well; for example, by CuII ions. In other words, there is a good chance that a mixed-metal (e.g., 16-iron-4-copper) derivative of 1 can be prepared.
Spectra are characterised by an absorption peak located from pH 0.3 through 7. In this pH domain, its electronic ion (CV) were used to assess the stability of the title poly anions of POMs. Both techniques demonstrate that CV results underscore, at least, two important characteristics of stability studies: UV/Vis spectroscopy and cyclic voltammetry studies indicate that 1 is stable from pH 0.3 through 7 and also its precursor P₃W₄₅. Figure 4 shows the CVs of 1 and P₃W₄₅ in superposition. In the cathodic branch of the CV of 1, is observed a reduction peak located at −0.220 V versus SCE, which does not exist in the pattern of the lacunary species P₃W₄₅. This wave is assigned to the reduction of the Fe(III) centres. As a matter of fact, the Fe(III) centres are known to be more easily reduced than the W(Ⅵ) centres as seen in several iron-containing polytungstates.[14–16] For potential scan rates from 1000 down to 2 mV s⁻¹, no splitting of the single Fe-wave of 1 was observed. Controlled potential coulometry with the potential set at −0.230 V versus SCE indicates the consumption of 16 electrons per molecule. In addition, the characteristic blue colour of reduced W centres of most POMs was not observed during such reduction. These observations together confirm that this wave features the simultaneous one-electron reduction of the 16 Fe(III) centres in 1. Reduction of all the 16 structurally equivalent Fe centres in 1 in a single step suggests that they are relatively independent, a feature that is reminiscent of the reduction process of certain polymers or dendrimers. Analogous examples can be found in POM electrochemistry.[16]

A detailed electrochemical study of 1 is provided in the Supporting Information (see Figures S1 and S2). These results underscore, at least, two important characteristics of 1, which render this molecule a potential candidate for triggering electrocatalytic processes. Firstly, accumulation of several metallic centres associated with simultaneous electron

**Electrochemistry**

Stability studies: UV/Vis spectroscopy and cyclic voltammetry (CV) were used to assess the stability of the title polyanion 1 by redissolving LiK₃·P₈W₄₈·10H₂O in several aqueous media classically used as supporting electrolytes in electrochemical studies of POMs. Both techniques demonstrate that 1 is stable from pH 0.3 through 7. In this pH domain, its electronic spectra are characterised by an absorption peak located roughly at 350 nm and assigned to the Fe(III) centres in the structure and a second peak around 265 nm due to the tungstophosphate ligand framework P₆W₉₆. The locations of these peaks depend on the pH. CV experiments, the duration of which can last up to 10 h, confirm also that 1 is stable in this pH domain.

**Voltammetric studies:** UV/Vis spectroscopy and CV studies indicate that 1 is stable from pH 0.3 through 7 and also its precursor P₃W₄₅.[13] Figure 4 shows the CVs of 1 and P₃W₄₅ in superposition. In the cathodic branch of the CV of 1, is observed a reduction peak located at −0.220 V versus SCE, which does not exist in the pattern of the lacunary species P₃W₄₅. This wave is assigned to the reduction of the Fe(III) centres. As a matter of fact, the Fe(III) centres are known to be more easily reduced than the W(Ⅵ) centres as seen in several iron-containing polytungstates.[14–16] For potential scan rates from 1000 down to 2 mV s⁻¹, no splitting of the single Fe-wave of 1 was observed. Controlled potential coulometry with the potential set at −0.230 V versus SCE indicates the consumption of 16 electrons per molecule. In addition, the characteristic blue colour of reduced W centres of most POMs was not observed during such reduction. These observations together confirm that this wave features the simultaneous one-electron reduction of the 16 Fe(III) centres in 1. Reduction of all the 16 structurally equivalent Fe centres in 1 in a single step suggests that they are relatively independent, a feature that is reminiscent of the reduction process of certain polymers or dendrimers. Analogous examples can be found in POM electrochemistry.[16]

A detailed electrochemical study of 1 is provided in the Supporting Information (see Figures S1 and S2). These results underscore, at least, two important characteristics of 1, which render this molecule a potential candidate for triggering electrocatalytic processes. Firstly, accumulation of several metallic centres associated with simultaneous electron
transfer and fairly fast kinetics is a necessary condition for reactions that require multiple electron transfers carried to completion and high efficiency. Secondly, the proximity of the Fe$^{III}$ and W$^{VI}$ waves of 1 can bring about beneficial effects.

To our knowledge, 1 constitutes the first example of multi-iron-containing POMs with such a small separation between the Fe$^{III}$ and the first W$^{VI}$ waves. The benefit of this property was demonstrated in the electrocatalytic reduction of nitrite by Fe$^{III}$-monosubstituted Wells–Dawson-type tungsstates or molybdates. Further examples were encountered in the electrocatalytic reduction of dioxygen and nitrogen oxides by Cu$^{II}$-substituted tungstomolybdates.

Electrocatalytic behaviour of 1 towards NO$_x$: The electrocatalytic reduction of nitrate remains a challenge in the NO$_x$ series because very few POMs are active in such electrocatalytic reduction. The positive results observed with 1 are described in the Supporting Information. An exciting observation is described in the following.

Reversible binding of NO to 1: The interest in NO has grown considerably ever since its important role in biology, environment and industry was unveiled.

Very recently, we have demonstrated by CV that the following two POMs, [Co(H$_2$O)$_3$][H$\beta$-$\beta$-Si$_{12}$W$_{40}$O$_{136}$](OH)]$_{25}$–[16] and [[Sn(CH$_3$)$_3$(H$_2$O)]$_{25}$[Sn-(CH$_3$)$_3$]$_{25}$[Sn-(CH$_3$)$_3$]$_{25}$[A-PW$_{12}$O$_{40}$]$_{25}$–[16–18] interact reversibly with NO or related species. A similar, but much weaker interaction was detected for the plenary Wells–Dawson type tungstophosphate [PW$_{12}$O$_{40}$]$^{8-}$ and the plenary Keggin type tungstosilicate [SiW$_{12}$O$_{40}$]$^{8-}$. These observations suggest that the combination of large POM size and/or incorporation of Co$^{II}$ or diorganotin moieties favours interaction with NO, followed by efficient electrocatalytic reduction of NO$_x$.[18c]

With this prior knowledge in mind, we decided to study the interaction of 1 with NO, hoping for an associated catalytic activity of the polyanion towards the reduction of NO. For these experiments solutions of 1 at pH 1 were saturated in an alternating fashion with argon and NO, respectively. The main observations are illustrated in Figure 5: the potential domain is restricted to that of the composite wave recorded on a solution of 1 saturated with argon; it is composed of the Fe$^{III}$ reduction wave and the first W$^{VI}$ reduction wave of 1. In the presence of NO an important catalytic current is observed that sets in at a more positive potential than that of the composite wave of 1. In addition, the intensity of this catalytic wave increases strongly with time (Figure 5A). The cell was checked for leakage as explained in the Experimental Section.

An analogous behaviour is observed for PW$_{12}$ in the presence of NO (Figure S3 in the Supporting Information), albeit with a roughly seven times weaker intensity than for 1 (Figure S4). However, this example permits to highlight details of the catalytic current. Figure 5B shows in superposition this catalytic process with its current scaled down to make its peak current match that observed in the presence of argon. Three closely spaced waves can be distinguished, with the peak potential of the first one located roughly 0.1 V positive of the first wave of PW$_{12}$. In short, this positive wave should be associated with a complex between NO and PW$_{12}$. The same kind of complex is probably also present for 1, but obscured by the Fe$^{III}$-wave, thus leaving only the overall positive shift of the catalytic wave. It must be noted that such shift is larger for 1 than for PW$_{12}$. We found that NO can be eliminated by bubbling argon through the solutions and then voltammograms virtually identical to the original ones are restored for 1 and PW$_{12}$.

Together, these observations indicate a reversible interaction between NO and 1 or PW$_{12}$, followed by electrocatalytic reduction of NO.

Magnetic susceptibility and EPR studies: The molar magnetic susceptibility (χ$_{m}$) and χ$_{m}$T of LiK-1 as a function of temperature, T, are displayed in Figure 6. The observed

![Figure 5](image-url)
room-temperature $\chi_m T$ value of 21.9 emu K mol$^{-1}$, when compared with that expected (70 emu K mol$^{-1}$) for 16 non-interacting Fe$^{III}$ ($S = 5/2$, $g = 2$) ions, indicates that antiferromagnetic interactions play a dominant role in the title polyanion 1. At 1.8 K, a $\chi_m T$ value of 3.3 emu K mol$^{-1}$ suggests that the ground state is $S_T = 2$ (expected is 3 emu K mol$^{-1}$ with $g = 2$).

The highly symmetrical [Fe$_{16}$(OH)$_{28}$(H$_2$O)$_4$]$^{20+}$ magnetic cluster incorporated in 1 is composed of 16 equivalent Fe$^{III}$ centres (see Figure 2). There are three types of Fe-O-Fe bridges (see Figure 3) that require three exchange coupling constants $J_1$ (e.g., Fe1-O1Fe-Fe3(i)), $J_2$ (e.g., Fe1-O14F/O14G-Fe4) and $J_3$ (e.g., Fe2-O24F-Fe4) and the interplay between them determines the ground state of LiK-1. The presence of 16 spin states with a total spin $S_T$ ranging from 0 to 40 renders the detailed analysis of the magnetic susceptibility data of LiK-1 complicated and is beyond the scope of the present study. However, a comparison of the Fe-O bond lengths and Fe-O-Fe bond angles of 1 with the literature values for 2-hydroxo-bridged Fe$^{III}$ dimers and oligomers suggests that the magnitudes of $J_1$, $J_2$ and $J_3$ should be in the range of 20–25 cm$^{-1}$. We hypothesise that the exchange couplings might be very similar in magnitude resulting in closely spaced spin levels and therefore the ground-state $S_T = 2$ assignment can only be tentative. Although the absence of a plateau at around 3.3 emu K mol$^{-1}$ in the $\chi T$ profile supports our hypothesis, susceptibility measurements below 1.8 K are needed for confirmation.

Magnetisation ($M/N\beta$) versus field H for LiK-1 at various temperatures is plotted in Figure 7. At 1.8 K, as the field increases from 0 to 7 T, $M/N\beta$ tends to reach 4, as expected for an $S = 2$ spin system with $g = 2$. The decrease in magnetisation as the temperature increases from 1.8 to 20 K could be due to the expected $H/T$ dependence of the magnetisation.

In a further attempt to understand the magnetism of polyanion 1, electron paramagnetic resonance (EPR) spectra were collected for various frequencies (9.65–319.2 GHz) and temperatures (5–300 K) for a powder sample of LiK-1. Only one broad peak ($\Delta H_{pp} = 70 \pm 3$ mT) at $g = 2.002 \pm 0.001$ is observed at room temperature for all experimental frequencies (see Figure 8). This is reminiscent of our earlier results on the hexa-Fe$^{II}$-substituted Keggin dimer [Fe(OH)$_3$($A$-$\alpha$-GeW$_9$O$_{34}$(OH)$_3$)$_2$]$^{11-}$. Figure 9 shows some typical X-band (9.6462 GHz) and ~319 GHz spectra at a few temperatures.

At least three features are evident: 1) a strong decrease in the signal intensity with decreasing temperatures, 2) signal broadening with decreasing temperatures and 3) absence of any additional splitting of the main peak. The intensity decrease is consistent with a similar trend in the magnetic susceptibility (Figure 6), and can thus be attributed to the population of states with smaller $S_T$ values at lower temperatures. On the other hand, signal broadening could be due to dipolar broadening and/or shorter relaxation times. The lack of any fine structure at all frequencies and temperatures studied renders it meaningless to derive any conclusions about the single-ion anisotropy of the overall $S_T$ value. The low temperature $g\approx 4.3$ peak observed at X-band (indicated by * in Figure 9 top) is a Fe$^{III}$ impurity signal from the
sample. Finally, the small remnant signal observed at 319.2 GHz (see Figure 9 bottom) is attributed to a minor g~2 impurity in the system and is not considered significant to the overall focus of the present study.

**Conclusion**

We have prepared the 16-Fe$^{III}$ containing 48-tungsto-8-phosphate \([P_8W_{48}O_{184}Fe_{16}(OH)_{28}(H_2O)_4]\) (1) as the mixed cation salts Li$_4$K$_{16}$$\left[P_8W_{48}O_{184}Fe_{16}(OH)_{28}(H_2O)_{4}\right]$·66H$_2$O·2KCl (LiK-1) and Na$_4$K$_{16}$$\left[P_8W_{48}O_{184}Fe_{16}(OH)_{28}(H_2O)_{4}\right]$·100H$_2$O (NaK-1). Polyanion 1 contains 16 edge- and corner-sharing FeO$_6$ octahedra in the form of a cyclic, unprecedented \([Fe_{16}(OH)_{28}(H_2O)_{4}]^{20+}\) iron-hydroxo–aqua nanocluster, grafted on the inner surface of the crown-shaped \([H_2P_8W_{48}O_{184}]^{33–}\) \((P_8W_{48})\) precursor. The synthesis of 1 was accomplished by reaction of hydrate complexes of Fe$^{II}$ (in presence of O$_2$), Fe$^{III}$ and \([Fe_3O(CH_3COO)_3(H_2O)]^4+\) with \(P_8W_{48}\) in aqueous, acidic medium (pH$\approx$4).

Besides the unprecedented \([Fe_{16}(OH)_{28}(H_2O)_{4}]^{20+}\) iron-hydroxo–aqua nanocluster, the central (empty) cavity of the polyanion 1 has another highly interesting feature. Access of an oxidant/substrate to the "iron active site" is easily possible and therefore 1 is very attractive for catalytic applications. In fact, initial oxidation catalysis studies with air-oxygen as oxidant are highly promising.[11]

Furthermore, it is very likely that the cavity in 1 can be filled with additional metal centres, for example, those different from iron(III). We are currently engaged in the process of preparing mixed-metal derivatives of 1 (e.g., “Fe$^{III}$-M$_x$P$_8$W$^{VI}_{48}$”) with one or more of the iron centres substituted by other transition-metal ions (e.g. Mn$^{II}$, Co$^{II}$, Zn$^{II}$). Such derivatives could lead to interesting magnetic as well as catalytic properties.[24]

The electrochemistry of 1 is characterised by a single 16-electron Fe-wave featuring the simultaneous reduction of all the Fe$^{III}$ centres, in full agreement with their structural equivalence. This wave could not be split into distinct steps whatever the potential scan rate from 2 to 1000 mV s$^{-1}$. Its potential location is very close to that of the first W$^{VI}$ wave of the lacunary precursor \(P_8W_{48}\). Polyanion 1 shows efficient electrocatalytic properties regarding the reduction of NO, including nitrate. In addition, a remarkable reversible interaction between 1 and NO is observed. Such an interaction might justify investigating the biomimetic properties of this new POM.

The magnetic characterisation of LiK-1 indicates that the ground state is made up of spin \(S_f=2\), based on the data at 1.8 K. Even though we are unable to provide a quantitative estimate of the exchange interactions \(J_1, J_2\) and \(J_3\) we hypothesise that \(J_1\approx J_2\approx J_3\) and the observed \(S_f=2\) could be regarded as a tentative ground state of 1. Additional measurements in the low-temperature range (<1.8 K) are needed to ascertain this prediction.

The study of a reaction of a solution of \(P_8W_{48}\) with metal cations offers the possibility to obtain basic information about principles of directed assembly processes under geometrically confined conditions. This can also lead, as in the present case, to cationic nanoclusters not obtainable under bulk conditions (see also reference [9]). One specific reaction described here refers to the “uptake of iron” under oxidative conditions and “release” under reducing conditions (the related simple reactions can be done under different reducing conditions) thereby mimicking the process occurring in the cavity of the protein ferritin. In this context one may also think about the option to study in a nanocavity the important reaction steps occurring during the reduction of O$_2$ with Fe$^{III}$, leading to O$_2^-$ and OH$^-$[25] even under simple “single-molecule type” conditions.

**Experimental Section**

**Synthesis:** The precursor salt K$_{28}$Li$_{5}$$\left[H_7P_8W_{48}O_{184}\right]$·92H$_2$O was synthesised according to the published procedure of Contant[25] and the purity was confirmed by infrared spectroscopy. All other reagents were used as purchased without further purification.

\(\text{Li}_x\text{K}_y\text{P}_8\text{W}_{48}\text{O}_{184}\text{Fe}_{16}(\text{OH})_{28}(\text{H}_2\text{O})_4\cdot66\text{H}_2\text{O}\cdot2\text{KCl} (\text{LiK-1})\)

**Method 1 (Bremen):** A sample of K$_{28}$Li$_{5}$$\left[H_7P_8W_{48}O_{184}\right]$·92H$_2$O (0.370 g, 0.025 mmol) was dissolved in LiCH$_3$COO/CH$_3$COOH buffer (0.5 g, 20 mL) at pH 4.0. Then FeCl$_3$·6H$_2$O (0.169 g, 0.625 mmol) was added and after complete dissolution, 30% H$_2$O$_2$ (10–20 drops) was added. Then...
the solution was heated to 80°C for 1 h and filtered while hot. After cool-
ing to room temperature the filtrate was layered with around of KCl (1 mL solution). Slow evaporation in an open beaker at room tem-
perature resulted in dark yellowish crystals after about one week. Evapo-
ration was allowed to continue until the solution level had almost ap-
proached the solid product **LiK-1**, which was then collected by filtration, washed with cold water and air dried. Yield: 0.083 g (22%). IR: ν = 1191 (sh), 1064 (s), 1019 (m; all νCH(P-O)), 951 (s), 927 (s, ν(C-O)), 793 (s), 752 (s), 687 (s), 647 (sh) (νC(W-O-W)), 559 (w), 526 (w), 473 cm⁻¹ (w); elemental analysis calculated (%) for **LiK-1**: Li 0.24, K 4.73, Fe 5.35, W 58.1, P 1.60, Cl 0.28.

**Table 1.** Crystal data and structure refinement for **LiK-1**.

<table>
<thead>
<tr>
<th><strong>LiK-1</strong></th>
<th><strong>NaK-1</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>M</strong> (gm⁻³)</td>
<td>15.4737</td>
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<tr>
<td>crystal system</td>
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<tr>
<td>space group (No.)</td>
<td>Pmm (58)</td>
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<tr>
<td>a [Å]</td>
<td>36.3777(9)</td>
</tr>
<tr>
<td>b [Å]</td>
<td>13.9708(3)</td>
</tr>
<tr>
<td>c [Å]</td>
<td>26.9140(7)</td>
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<tr>
<td>V [Å³]</td>
<td>13678.64(6)</td>
</tr>
<tr>
<td>goodness-of-fit on F²</td>
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</tr>
<tr>
<td>R [F² &gt; 2σ(F²)]</td>
<td>0.059</td>
</tr>
<tr>
<td>Rg (all data)</td>
<td>0.183</td>
</tr>
</tbody>
</table>

Crystal data for **NaK-1** with dimensions 0.06 x 0.11 x 0.1 mm was removed from the mother liquor and immediately cooled to 183(2) K on a Bruker AXS SMART diffractometer (three-circle goniometer with 1 K CCD detector, MoKα radiation, graphite monochromator; hemisphere data collection in ω at 0.3° scan width in three runs with 606, 435 and 230 frames (φ = 0, 88 and 180°) at a detector distance of 5 cm). A total of 78576 reflections (1.48θ < 2θ < 0.05°) were collected of which 29084 reflections were unique (Rint = 0.1127). An empirical absorption correction using equivalent reflections was performed with the program SADABS 2.0. The structure was solved with the program SHELXS-97 and refined using SHELXL-97 to R = 0.0188 for 14766 reflections with I > 2σ(I). R = 0.1935 for all reflections; max/min residual electron density 5.573 and −2.272 eÅ⁻³. The crystallographic data collection and structure determination were performed by the Fachinformationszentrum Karlsruhe, 76344 Eggenstein-Leopoldshafen, Germany (Fax: (49)-7247-808-666; e-mail: crysdata@fiz-karlsruhe.de) on the depository number CSD-418194 (LiK-1) and CSD-418527 (NaK-1).

**U/Vis spectroscopy:** Pure water was used as solvent throughout, which was obtained by passing through a ROs 8 unit followed by a Millipore-Q Academic purification set. All reagents were of high-purity grade and were used as purchased without further purification. The UV/Vis spectra were recorded on a Perkin-Elmer Lambda 19 spectrophotometer on 1.6 x 10⁻⁶ molar solutions of **LiK-1**. Matched 1.00 cm optical path quartz cuv-
ettes were used. The following media proved useful for the present study: 0.5 mol/l HCl, pH 0.3; 1 mol/l LiOH, pH 1; 3 mol/l CH₃COOH/CH₃COONa, pH 5 to 7.

Several crystals were filtered off after 3 days; these crystals were washed with a small amount of cold water and dried in air. Yield: 0.03 g, 8%; elemental analysis calculated (%) for **NaK-1**: Na 0.18, K 4.56, Fe 5.78, W 57.1, P 1.60, Cl 0.28.

**Method 2 (Bremen):** A sample of K₂Li₃[H₃PW₁₀O₄₀Fe₄(OH)₄·6H₂O·2KCl (0.0974 g, 0.275 mmol) was dissolved in NaCH₃COO/CH₃COOH buffer (2.0 mL, pH 4.0). Then Fe(CH₃COO)₆(H₂O)₄·100H₂O (NaK-1) was added and the solution was heated to 80°C for 1 h and filtered while hot. The following steps were identical to those of Method 1. The identity of **LiK-1** (isolated in very low yield, <10%) was established by XRD and IR.

**Method 3 (Bremen):** A sample of K₂Li₃[H₃PW₁₀O₄₀Fe₄(OH)₄·6H₂O·2KCl (0.0974 g, 0.275 mmol) was dissolved in LiCH₃COO/CH₃COOH buffer (2.0 mL, pH 4.0). Then Fe(CH₃COO)₆(H₂O)₄·100H₂O (NaK-1) was added and the solution was heated to 80°C for 1 h and filtered while hot. After cooling, the solution was filtered and the yellowish crystals that were filtered off after 3 days; these crystals were washed with a small amount of cold water and dried in air. Yield: 0.09 g, 15% (based on PₓWᵧ); elemental analysis calculated (%) for **NaK-1**: Na 1.30, K 2.71; found Na 1.3, K 2.8. The identity of **NaK-1** was established by elemental analysis (in part done by Mikroanalytisches Labor Egmont Pascher, see above), IR spectroscopy and complete single-crystal X-ray structure analysis.

**Method 4 (Bielefeld):** K₂Li₃[H₃PW₁₀O₄₀Fe₄(OH)₄·6H₂O·2KCl (0.067 g, 0.046 mmol) was dissolved in NaCH₃COO/CH₃COOH buffer (1 x 30 mL, pH 4.2). After Fe(C₂O₄)₃·H₂O (0.2 g, 0.31 mmol) was added, the solution was heated to 60°C for 30 h and filtered after cooling to room temperature. Slow evaporation in an open Erlenmeyer flask at room temperature resulted in the precipitation of dark yellowish crystals that were filtered off after 3 days; these crystals were washed with a small amount of cold water and dried in air. Yield: 0.09 g, 15%; elemental analysis calculated (%) for **NaK-1**: Na 1.30, K 2.71; found Na 1.3, K 2.8. The identity of **NaK-1** was established by elemental analysis, XRD and IR spectroscopy.

**X-ray crystallography**

Crystal data for **LiK-1** with dimensions 0.06 x 0.12 x 0.33 mm² was mounted in oil on a Hampton cryoloop for indexing and intensity data collection at 173 K on a Bruker D8 APEX II CCD single-crystal diffractometer with MoKα radiation (λ = 0.71073 Å). Of the 335706 reflections collected (2θ max. = 52.8°, 99.7% complete), 14333 were unique (Rint = 0.153) and 10468 reflections were considered observed [I > 2σ(I)]. The data were processed using SAINT (from Bruker AXS) and an absorption correction was performed using the SADABS program (G. M. Sheldrick, Brucker AXS). Direct methods were used to locate the tungsten atoms (SHELXS-97), and the remaining atoms were found from successive Fourier maps (SHELXL-97). No H or Li atoms were located. The final cycles of refinement on F² over all data included the atomic coordinates, anisotropic thermal parameters (W, Fe, P, Cl and non-disordered K atoms) and isotropic thermal parameters (O and disordered K atoms), converging to R = 0.059 [I > 2σ(I)] and Rₚ = 0.183 (all data). In the final difference map the deepest hole was −3.48 eÅ⁻³ (0.94 Å⁻³ from WS) and the highest peak 4.20 eÅ⁻³ (0.73 Å⁻³ from K4). The crystallographic data are provided in Table 1.
contaminants such as NO, to be scavenged in 9M KOH. NO was bubbled through the electrolyte in the electrochemical cell for 30 min, resulting in a NO-saturated solution (1–2 mM). The electrochemical cell was checked for leaks in the following way: solutions saturated with NO were kept for several hours and used for electrocatalytic reduction of this substrate; after removing NO by bubbling pure argon, no electroactivity of NO or a related species could be detected in the potential range from +0.920 V to −0.730 V at pH 1.

The source, mounting and polishing of the glassy carbon (GC) electrodes were described previously.30 The glassy carbon samples had a diameter of 3 mm. The electrochemical set-up was an EG & G 273 A driven by a PC with the M270 software. Potentials are quoted against a saturated calomel electrode (SCE). The counter electrode was a platinum gauze of large surface area. All experiments were performed at room temperature.

Magnetic measurements: Magnetic susceptibility and magnetisation measurements were carried out on powder samples of LiK-1 using a Quantum Design MPMS SQUID magnetometer in the temperature range of 1.8–300 K and field range of 0–7 T. The data were corrected for the sample holder, TIP of Fe3+ W4+ ions, and molecular diamagnetism which was estimated from Klemm constants.31

EPR measurements: Polycrystalline powder EPR spectra of LiK-1 were recorded at frequencies ranging from 9.64 to 320 GHz at the high-field electron magnetic resonance facility at the National High Magnetic Field Laboratory in Tallahassee, FL, as described elsewhere.32 Temperature variation was carried out from room temperature to 5 K. An Oxford Instruments Teslacon superconducting magnet sweepable between 0 and 17 T was used to apply the Zeeman field. In all experiments the modulation amplitudes and microwave power were adjusted for optimal signal intensity and resolution.

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[1] a) This refers in a general sense to scenarios where “materials” grow with preferred orientations on surfaces influenced/directed by the surfaces’ geometry: V.E. Henrich, P.A. Cox, The Surface Science of Metal Oxides, Cambridge University Press, Cambridge 1994, p. 384; b) The process is quite common in the geosphere. This leads to situations where spaces with larger scale sizes (bulk condition compara- rable), but also up to smaller scales, are filled with minerals or water; the well-known geodes are objects of that type (see textbooks of mineralogy). The confinement induced changes in the water structure and dynamics, which are commonly substrate specific, play a key-role in the geosphere regarding the reactivity of mineral surfaces (see J. Wang, A.G. Kalinichev, J. Phys. Chem., B 2005, 109, 14308–14311). Important information about that topic can be obtained from POM chemistry (see: A. Oleinikova, H. Weimagertner, M. Chaplin, E. Diemann, H. Böge, A. Müller, ChemPhysChem 2007, 8, 646–649).


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Chapter 3

Summary

The thesis refers mainly to the synthesis of unique polyoxometalates having interesting magnetic properties. The compounds 1-5 which were obtained are described below.

The classical/historical polyoxometalates are diamagnetic, however they are currently receiving much attention in the field molecular magnetism due to the fact that: (i) they can act as ligands for the coordination of paramagnetic ions, like V(IV), Fe(III), Cr(III), Fe(II), Mn(II), Co(II), Ni(II) and Cu(II), leading to well-defined magnetic clusters of controlled nuclearities and topologies, (ii) they offer the opportunity to investigate both experimentally and theoretically the nature of the magnetic exchange interactions, and (iii) they stimulate the development of new theories to account for the interplay between electron delocalization and exchange interactions in high-nuclearity mixed valence clusters. In this dissertation we investigated the utility of the pentagonal \{Mo\}_5-type unit in constructing spherical systems with interesting magnetic properties.

Molybdenum-oxide based spherical, hollow clusters of the type \{(Mo)Mo\}_{12}\{linker\}_{30} – because of their structural features also called Keplerates – can for instance be directly synthesized by the addition of linkers such as V\textsuperscript{IV}O\textsuperscript{2+}, Fe\textsuperscript{III} and Cr\textsuperscript{III} to a dynamic library containing (virtual) pentagonal units in solution.

In the \{Mo\textsubscript{72}V\textsubscript{30}\}_1 cluster the 30 V\textsuperscript{IV} centers – acting as linkers for the pentagonal \{(Mo\textsuperscript{VI})Mo\textsuperscript{VI}\}_5 type units – describe a (slightly distorted) icosidodecahedron, the distortion is in agreement with the fact that 20 V\textsuperscript{IV} in the equatorial region have octahedral coordination and the two sets of five V\textsuperscript{IV} centers in the polar area have square pyramidal coordination. Ten of the twelve SO\textsubscript{4}\textsuperscript{2–} ligands are coordinated by three oxygen atoms to three adjacent Mo\textsuperscript{VI} centers of the
\{(MoVI)MoVI\}_5 \text{ groups such that two } \{KSO_4\}_5 \text{ rings parallel to the equator result with the K+ cations (formally) bridging the SO}_4^{2-} \text{ anions.}

\[
\text{Na}_8\text{K}_{14}(\text{VO})_2\{\text{Mo(Mo)}_3\text{O}_{21}(\text{H}_2\text{O})_{30}\}\{\text{Mo(Mo)}_5\text{O}_{21}(\text{H}_2\text{O})_8(\text{SO}_4)\}_2\{\text{VO(H}_2\text{O})\}_2\text{[VO]}_{10}\ \{(\text{KSO}_4\}_5\}_2 \cdot \text{ca. 150 H}_2\text{O}
\]

We know that practically neutral sherical molybdenum-oxide based capsules – but positively charged without the organic ligands – have a high affinity for integration of different guest species. In the cavity of the \{\text{Mo}_{72}\text{Fe}_{30}\}_1 \text{ capsule a hexamolybdate Lindqvist type anion } [\text{Mo}_6\text{O}_{19}]^{2-} \text{ got encapsulated, noncovalently bonded. Such hexamolybdate polyanion cannot be obtained as other polymolybdates simply in aqueous medium and it is preferably formed in organic solvents while crystallizing together with organic cations. In the present system it is stabilized/protected by the comparably large number of acetates inside the cavity forming a hydrophobic protecting environment. As the skeleton of 2a is nearly neutral and the pure inorganic part, \textit{i.e.} without the acetate ligands, is even positively charged, this allows the attraction of small anionic fragments with the consequence that the stepwise formation of the } [\text{Mo}_6\text{O}_{19}]^{2-} \text{ occurs parallel to the growth of the capsule, } \textit{i.e.} \text{ on its internal electrophilic surface.}

\[
\text{Na}_4[\text{Mo}_6\text{O}_{19}]^{2-} \subset \{\text{Mo}^{VI}_{72}\text{Fe}^{III}_{30}\text{O}_{252} (\text{CH}_3\text{COO})_{20} (\text{H}_2\text{O})_{92}\} \cdot \text{ca. 120 } \text{H}_2\text{O}
\]

In the cavity of \{\text{Mo}_{72}\text{Cr}_{30}\}_3 \text{ cluster an } \{\text{Na(H}_2\text{O)}_{12}\}_1 \text{ cluster is encapsulated where the O atoms form an icosahedron, with its approximate C}_5 \text{ axes coinciding with the C}_5 \text{ axes of the metal-oxide based skeleton. The relatively small water cluster is protected by the hydrophobic shell built up by the acetate ligands and should therefore not be formed within the cavities of the present type of capsules which contain hydrophilic interiors.}

\[
\{\text{Na(H}_2\text{O)}_{12}\}_1 \subset \{\text{Mo}^{VI}_{72}\text{Cr}^{III}_{30}\text{O}_{252}(\text{CH}_3\text{COO})_{19}(\text{H}_2\text{O})_{94}\} \cdot \text{ca. 120 } \text{H}_2\text{O}
\]
The three practically isostructural clusters/skeletons of the type \{(Mo)Mo_5\}_12\{M\}_30 with a sequence of spin values S=1/2 (V^{IV}), S=3/2 (Cr^{III}) and S=5/2 (Fe^{III}) show quite different magnetic behavior. The capsule/Keplerate with M=Fe^{III} has essentially a classical behavior with weak superexchange interaction due to the large spin 5/2 centred at 30 different corners of the \{M_{30}\} icosidodecahedron. By contrast, in the case of M=V^{IV} where the spin is small and the electrons delocalized, the system is called a “Quantum Keplerate”. The capsule with M=Cr^{III} is just in between that of the two other ones with the correspondingly intermediate magnetic properties.

The wheel-shaped \{P_8W_{48}\} polyoxotungstate provides a reaction chamber for the directed-assembly process leading to 4a, where two unprecedented mixed-valence vanadium oxide \{V^{IV}_4V^{IV}_2O_{12}(H_2O)_2\}^{4+} cavity-capping groups based on linked octahedra and tetrahedra with V^{IV} and V^{V} centers, respectively are encapsulated and to 5a, which contains – in the form of a cyclic arrangement – the unprecedented \{Fe_{16}(OH)_{28}(H_2O)_4\}^{20+} nanocluster, with 16 edge- and corner-sharing FeO_6 octahedra, grafted on the inner surface of the crown-shaped \{P_8W_{48}\} precursor.

\[\text{Na}_{12}\text{K}_3\text{H}_4\cdot\text{K}_8\subset\{\text{P}_8\text{W}_{48}\text{O}_{184}\}\{\text{V}^{IV}_4\text{V}^{IV}_2\text{O}_{12}(\text{H}_2\text{O})_2\}_2\cdot\text{ca. 80 H}_2\text{O}\] 4

\[\text{Na}_9\text{K}_{11}\{\text{P}_8\text{W}_{48}\text{O}_{184}\text{Fe}_{16}(\text{OH})_{28}(\text{OH}_2)_4\}\cdot\text{ca. 100H}_2\text{O}\] 5

In 4a the terminal V^{IV}O bonds are directed towards the interior of the composite anion and towards the corresponding bonds of the opposite capping group (V...V separation \textit{ca} 6.2Å). Potassium cations occupy analogous positions within the new host-guest complex but they are now sandwiched between the two cyclic capping groups \{V^{IV}_4V^{IV}_2O_{12}(H_2O)_2\}^{4+}. There is no doubt that other nucleation processes can be studied in the \{P_8W_{48}\} cluster cavity under formally similar conditions.
Appendix

Spectroscopic data

**Infrared and Raman spectroscopy**
Vibrational spectra were recorded with a Bruker IFS 66 / FRAU 106 spectrometer. FT-IR: KBr pellet, recorded in the range 4000-400 cm$^{-1}$; FT-Raman: KBr matrix, excitation with a Nd: YAG-laser; $\lambda e = 1064$ nm.

**UV-Vis spectroscopy**
UV-Vis spectra were recorded from degassed solutions in the range 1200-200 nm with a Shimadzu UV-160A spectrophotometer and evaluated with a program associated with the spectrometer.
IR spectrum of \( \text{Na}_8\text{K}_4\{\text{VO}\}_2\{\{\text{Mo(Mo)}_3\text{O}_{21}(\text{H}_2\text{O})_3\}_{10}\{\text{Mo(Mo)}_3\text{O}_{21}(\text{H}_2\text{O})_3(\text{SO}_4)\}_{2}\{\text{VO(H}_2\text{O)}\}_{20}\{\text{VO}\}_{10}\{\{\text{KSO}_4\}_3\}_{2}\} \cdot \text{ca. } 150\ \text{H}_2\text{O} \)

Solid state Raman spectrum of \( \text{Na}_8\text{K}_4\{\text{VO}\}_2\{\{\text{Mo(Mo)}_3\text{O}_{21}(\text{H}_2\text{O})_3\}_{10}\{\text{Mo(Mo)}_3\text{O}_{21}(\text{H}_2\text{O})_3(\text{SO}_4)\}_{2}\{\text{VO(H}_2\text{O)}\}_{20}\{\text{VO}\}_{10}\{\{\text{KSO}_4\}_3\}_{2}\} \cdot \text{ca. } 150\ \text{H}_2\text{O} \)
UV-Vis spectrum of $\text{Na}_8\text{K}_{14}(\text{VO})_2\{\text{Mo}_\text{II}(\text{Mo})_5\text{O}_{21}(\text{H}_2\text{O})_3\}_{10}\{\text{Mo}(\text{Mo})_3\text{O}_{21}(\text{H}_2\text{O})_3(\text{SO}_4)\}_2$
$\{\text{VO}(\text{H}_2\text{O})\}_{20}\{\text{VO}\}_{10}\{\text{KSO}_4\}_3\}_2 \cdot \text{ca. 150 H}_2\text{O}$

IR spectrum of $\text{Na}_4\{\text{Mo}_6\text{O}_{19}\}_2^2\subset\{\text{Mo}^{\text{VI}}_{72}\text{Fe}^{\text{III}}_{30}\text{O}_{252} (\text{CH}_3\text{COO})_{20} (\text{H}_2\text{O})_{62}\}_2 \cdot \text{ca. 120 H}_2\text{O}$
Solid state Raman spectrum of Na₄[{Mo₆O₁₉}²⁻⊂{Mo⁶⁺Fe³⁺₇₂O₂₅₂(CH₃COO)₂₀(H₂O)₉₂}]·ca. 120 H₂O

UV-Vis spectrum of Na₄[{Mo₆O₁₉}²⁻⊂{Mo⁶⁺Fe³⁺₇₂O₂₅₂(CH₃COO)₂₀(H₂O)₉₂}]·ca. 120 H₂O
IR spectrum of $[[\text{Na(H}_2\text{O})_{12}] \subset \{\text{Mo}^{VI}_{72}\text{Cr}^{III}_{30}\text{O}_{252}(\text{CH}_3\text{COO})_{19}(\text{H}_2\text{O})_{94}\}] \cdot \text{ca. 120 H}_2\text{O}$

Solid state Raman spectrum of $[[\text{Na(H}_2\text{O})_{12}] \subset \{\text{Mo}^{VI}_{72}\text{Cr}^{III}_{30}\text{O}_{252}(\text{CH}_3\text{COO})_{19}(\text{H}_2\text{O})_{94}\}] \cdot \text{ca. 120 H}_2\text{O}$
Appendix

UV-Vis spectrum of $\{Na(H_2O)_{12}\subset Mo^{VI}_{72}Cr^{III}_{30}O_{252}(CH_3COO)_{19}(H_2O)_{94}\} \cdot ca. 120 H_2O$

IR spectrum of $Na_{12}K_8H_4[K_8\subset P_8W_{48}O_{184}]\{V^{W}_4V^{IV}_{2}O_{12}(H_2O)_{2}\} \cdot ca. 80 H_2O$
Solid state Raman spectrum of $\text{Na}_{12}\text{K}_8\text{H}_4[\text{K}_8\subset\{\text{P}_8\text{W}_{48}\text{O}_{184}\}\{\text{V}^{IV}_{4}\text{V}^{IV}_{2}\text{O}_{12}\text{(H}_2\text{O})_2\}] \cdot \text{ca. 80 H}_2\text{O}$

UV-Vis spectrum of $\text{Na}_{12}\text{K}_8\text{H}_4[\text{K}_8\subset\{\text{P}_8\text{W}_{48}\text{O}_{184}\}\{\text{V}^{IV}_{4}\text{V}^{IV}_{2}\text{O}_{12}\text{(H}_2\text{O})_2\}] \cdot \text{ca. 80 H}_2\text{O}$
**IR spectrum of Na$_9$K$_{11}$[P$_8$W$_{48}$O$_{184}$Fe$_{16}$(OH)$_{28}$(OH$_2$)$_4$]·100H$_2$O**

**UV-Vis spectrum of Na$_9$K$_{11}$[P$_8$W$_{48}$O$_{184}$Fe$_{16}$(OH)$_{28}$(OH$_2$)$_4$]·100H$_2$O**
Chapter 4

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Microsoft Office (Word, Excel, PowerPoint), Origin, Corel Draw, Internet related software.

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LIST OF PUBLICATIONS


2. A. Müller, A. M. Todea, H. Bögge, J. van Slageren, M. Dressel, A. Stammler, M. Rusu, "Formation of a "less stable" polyanion directed and protected by electrophilic internal surface functionalities of a capsule in growth: [{Mo_{6}O_{19}}^{2-} in {Mo^{VI}Fe^{III}_{72}O_{252}(ac)_{20}(H_{2}O)_{92}}]^{4-}", *Chem. Commun.*, **2006**, 3066-3068.

