Controlling the Reactivity of the [P₈W₄₈O₁₈₄]⁴₀⁻ Inorganic Ring and Its Assembly into POMZite Inorganic Frameworks with Silver Ions

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Abstract: The construction of pure-inorganic framework materials with well-defined design rules and building blocks is challenging. In this work, we show how a polyoxometalate cluster with an integrated pore, based on [P₂W₁₂O₄₈]¹⁴⁻ (abbreviated as [P₂W₁₂]), can be self-assembled into inorganic frameworks using silver ions, which both enable reactions on the cluster as well as link them together. The [P₂W₁₂] was found to be highly reactive with silver ions resulting in the in situ generation of fragments, forming [P₆W₂₆O₇₁] and [P₆W₂₆O₇₁] in compound (1) where these two clusters co-crystallize and are connected into a POMZite framework with 11 Ag⁺ ions as linkers located inside clusters and 10 Ag⁺ linking ions situated between clusters. Decreasing both the concentration of Ag⁺ ions, and the reaction temperature compared to the synthesis of compound (1), leads to [P₆W₂₆O₇₁] in compound 2 where the [P₆W₁₂] clusters are linked to form a new POMZite framework with 9 Ag⁺ ions per formula unit. Further tuning of the reaction conditions yields a cubic porous network compound (3) where [P₆W₁₂] clusters as cubic sides are joined by 4 Ag⁺ ions to give a cubic array and no Ag⁺ ions were found inside the clusters.

Polyoxometalate (POM)-based materials are a family of compounds known for their rich structural diversity and properties. Over the years many synthetic strategies have been developed to control the self-assembly of the POMs as effective anionic molecular inorganic building blocks. One of the most recent successes of this strategy was made possible by using the superlacunary cyclic heteropolyanion [P₈W₄₈O₁₈₄]⁴₀⁻ (abbreviated as [P₈W₄₈]) as a building block for the construction of intrinsically porous all-inorganic framework materials, named POMZites. Polyoxometalate-based framework materials, or POMZites, are an emerging class of configurable all-inorganic porous materials. Porous materials such as zeolites or metal–organic frameworks, MOFs, are ordered networks whose building units are linked with strong interactions via ionic, covalent, and coordination bonds. Silver ions are often used as a flexible linkers in coordination chemistry, for instance, silver-linked molybdenum oxide POMs are very stable, both in the solid and liquid phases. Further, silver ions play a remarkable role in chemical reactions and crystallization processes acting both as counterions and structure directors. In this work, we demonstrate that by using silver ions as linkers, two new porous polyoxotungstates (POTs) can be derived from [P₈W₄₈], which was isolated originally by Contant and Tezé in 1985. The [P₈W₄₈] cluster has a symmetry of point group D₄h and is a highly stable and versatile oligomer formed from the aggregation of four subunits of the hexavacant [P₈W₄₈O₁₈₄]⁴₀⁻. In 2005, Kortz et al. reported the first Cu-containing [P₈W₄₈] assembly, thus proving this molecule to be a large superlaccunary polyanion precursor. Since then, a series of novel structures based on [P₈W₄₈] clusters have been reported in the literature including [Cu₄₈] clusters[12] [V₄₃] aggregates,[13] [Fe₄₉] and Ln-containing cluster anions,[14] as well as organoruthenium-based composites.[15] All of these structures were based on transition metal complexes with the superlacunary [P₈W₄₈] cluster acting as a ligand. Herein, we present a unique method of extending the [P₈W₄₈]-based frameworks to form a higher nuclearity species by binding additional lacunary units, as well as building new POMZites.

The [P₈W₄₈] cluster can be viewed as a drum-like structure formed by the condensation of four [P₆W₁₂O₇₁] subunits. Around the corners of the [P₆W₁₂O₇₁] subunits on the top and bottom faces, there are eight sites where [W₄] units or other transition metal ions can be added to form new locations for further growth to extend the structures. The tungsten occupancy on these new growth points is significant, and can be differentiated from other transition metal ions like Co²⁺ or Mn²⁺.[16] Although these growth points have been noted as being occupied by both tungsten and other transition metals in previous publications,[16–19] no extended structures built upon these foundations have been reported. Our work demonstrates the formation of higher nuclearity clusters built up from the superlacunary [P₈W₄₈] cluster by extension of these growth sites (Scheme 1). The reaction of [P₈W₄₈] and a high concentration of Ag⁺ ions as starting materials led to the formation of Li₉K₅₅Ag₂[Hₚ₈W₄₈O₇₁]₅.
crystallization of the [P-W_{6}O_{35}]^{2−} and [P_{9}W_{66}O_{251}]^{2−} clusters. Both clusters have the same [P-W_{8}] base and have roughly similar sizes for the [P-W_{8}] and [P_{2}W_{12}] parts. The formation of high-nuclearity [P_{9}W_{63}O_{235}]^{2−} and [P_{10}W_{66}O_{251}]^{2−} clusters is only possible with a high concentration of Ag^{+} ions and heating/ refluxing during the synthesis process before crystallization. A careful analysis of the cluster structure found in compound (1) revealed that the central cavity of the [P-W_{8}] base is filled with Ag^{+} ions which form a [Ag_{9}] aggregate with two Cl− ions as cores, which is similar to the observation in other Ag-POM clusters.[17a] One more Ag^{+} ion was found to support and stabilize key [W_{4}] growth sites, similar to the [Ag_{9}] cluster, from within the central cavity of the cluster (Figure S2). The remaining 10 Ag^{+} ions per formula were found to link clusters which form a 3D network with a complex topology. The silver ions appear to use all the linking models of [W_{48}] that are possible with a high concentration of Ag^{+} ions.[5] The packing diagram in Figure S3 shows that the space between clusters accommodates silver ions at locations inside and between the clusters.

Compound Li_{8}K_{13}Ag_{13}[H_{12}P_{8}W_{51}O_{196}]·50H_{2}O (2) was synthesized under reaction conditions similar to (1), but at room temperature and with a lower concentration of AgNO_{3}. In the single-crystal structure determination, major silver ion positions are clearly defined with occupancies between 0.55 and 1.0. The structure can be described as [P_{8}W_{48}] rings that are virtually coplanar with ring center-to-center distances being about 22.9 and 24.2 Å in the two distinct dimensions. Each layer has a thickness of around 11.3 Å. The [P_{8}W_{48}] rings either within or between layers are linked by Ag^{+} ions (total 9 per formula, Figure S4) and the gaps and cavities are further filled by K^{+} ions.

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comparison to the above description of \((1)\). This confirms the existence of the growth points of the \([P_8W_{48}]\) in solution starting with either a minor tungstate impurity in the starting material, or the decomposition of a small amount of \([P_8W_{48}]\) during the reaction. In contrast to the \(Ag^+\) cluster found in the cavity of \([P_8W_{48}]\) rings in \((1)\), the corresponding positions in \((2)\) are instead occupied by \(K^+\) ions. The network structure of \((3)\) is best described as a “truncated cuboctahedron” (or great rhombicuboctahedron); in other words, the molecular parking of \([P_8W_{48}O_{184}]^{40^-}\) units into the network can essentially be viewed as cubic (Figure 3), a topology shared by some other notable materials.\(^{19}\) Strikingly, this topology is analogous to that of the prominent LTA (Linde Type A) zeolite framework, with a pore size of 0.4 nm.\(^{20}\) Both compound \((3)\) and POMzite-3\(^{21-25}\) [\(\text{Mn}_8(\text{H}_2\text{O})_3\text{P}_8\text{W}_{44}\text{O}_{116}]^{24-}\) have approximately spherical voids with an internal diameter of \(\approx 2.1\) nm (that is, a void volume of \(\approx 4.8\) nm\(^3\), see Figure 3) and are accessible through the integrated pores of the six surrounding \([P_8W_{48}O_{184}]^{40^-}\) anions.\(^{20}\) LTA, compound \((3)\), and POMzite-3 frameworks crystallize in the same cubic \(Pm\bar{3}m\) space group and differ only in the tiling around the central cubo-octahedral cavity.\(^{19,20}\)

Structural control between the two distinct structures, \((1)\) and \((2)\), was achieved by varying reaction temperatures and the concentration of \(Ag^+\) ions, whilst maintaining the same pH 1.53. Compound \((1)\) was obtained by heating at 80°C for 30 minutes with a much higher concentration of \(Ag^+\) ions that replaces the \(K^+\) ions in the \([P_8W_{48}]\) cavity of the starting materials and produces more \([PW_9]\) and \([P_2W_{12}]\) fragments, which fully fill the growth sites on the \([P_8W_{48}]\) base. This leads to the formation of molecular clusters \([P_8W_{40}O_{120}]\) and \([P_10W_{36}O_{180}]\) which co-crystallize in compound \((1)\). The synthesis of \((2)\) under similar conditions but with a lower concentration of \(Ag^+\) ions and without heating yields some partially filled growth points but with no extended POM structure, such as \([P_8W_{40}O_{120}]\) and \([P_10W_{36}O_{180}]\) as in \((1)\). Under such conditions \(Ag^+\) ions cannot substantially replace the \(K^+\) ions in the \([P_8W_{48}]\) cavity of the starting materials and no significant amount of \([PW_9]\) and \([P_2W_{12}]\) fragments are produced. Compound \((3)\), synthesized with the reaction conditions of \((2)\), but with a lower pH of 1.45, has no growth points filled with \([W_1]\) units. This is because lower pH is not favorable for \([P_8W_{48}]\) to disassemble to produce fragments or even \([W_1]\) units. Also lower pH is not favorable for \(Ag^+\) ions to coordinate to oxo ligands because oxo has a higher affinity to protons. Therefore, fewer \(Ag^+\) ions are included in \((3)\) and hence fewer O-Ag-O bridges are formed between the \([P_8W_{48}]\) clusters, generating a less dense cubic framework compared with compounds \((1)\) and \((2)\) (Figure S5).

In conclusion, by varying the reaction conditions—the temperature, pH and the concentration of \(Ag^+\) ions—it is possible to control the synthesis of lacunary \([PW_{9}]\) and
[P,W] fragments which in turn produce high-nuclearity tungstate clusters, [P,W] and [P,W]. Both clusters have the same [W] base and roughly similarly sized [P,W] and [P,W] additional units. These discoveries demonstrate that the [P,W] cluster is an important building block for the construction of POM materials and networks utilizing cluster paneling, via extension from the eight possible growth points. The role of Ag+ is significant by filling the [P,W] cavity to support the extended clusters, as further evidenced by the fact that Ag+ ions are needed to produce these clusters. For example, the potassium salt of the starting material’s [P,W] cluster forms readily from plenty of [P,W] fragments, but further growth cannot be achieved without adding silver ions to replace K+ ions and support the key growth sites. Future work will focus on the reaction of Ag+ ions with other lacunary POM clusters as well as exploring the properties of the framework with regard to guest uptake and reactivity of the inner pores.

**Experimental Section**

General experimental remarks: All chemicals were purchased from commercial sources and used without further purification. CCDC 1892362, 1892363 and 1892364 contain the supplementary crystallographic data for this paper. These data can be obtained free of charge from The Cambridge Crystallographic Data Centre.

**Synthesis of**

LiKAg([H,P,W]O )·H[P,W]O (1): In a 25 mL round-bottomed flask, LiNO (84 mg, 1.2 mmol) was dissolved in 12 mL of H2O, then KLi([H,P,W]O )·92H2O (102 mg, 6.9 × 10 mmol) was added and dissolved. The solution was adjusted to pH 1.53 by using HNO3 (70%), then AgNO3 (35.7 mg, 0.21 mmol) was added. The mixture was then heated at 80°C for 30 min then cooled down to room temperature. After two weeks well-behaved block colorless crystals started to form in solution. The products were isolated after several weeks. The chlorine came from the trace amount in the starting materials. Yield: 0.06 g. Characteristic IR bands (in cm−1): v(O–H), 3439 (br), 1620(m); v(O–Ag), 1384 (s); v(P–O), 1128 (s), 1086 (s); v(W–O), 1015 (w), 926 (br). Elemental analysis calcd (found) for AgClH2L2K3Li2O15P16W30: Ag 2.7 (2.9), W 54.9 (55.1), K 2.92 (3.17), Li 0.43 (0.43). TGA water loss from room temperature to 400°C, calc (found): %: 19.0 (19.5).

**Synthesis of**

LiKAg([H,P,W]O )·50H2O (2): In a 25 mL flask, LiNO (84 mg, 1.2 mmol) was dissolved in 12 mL of H2O, then KLi([H,P,W]O )·92H2O (102 mg, 6.9 × 10 mmol) was added. The solution was adjusted to pH 1.53 by using HNO3 (70%), then AgNO3 (13.7 mg, 0.08 mmol) was added. The mixture was then stirred at room temperature for 5 min. After three weeks well-behaved block, colorless crystals started to form in solution. The products were isolated after several weeks. Yield: 0.12 g. Characteristic IR bands (in cm−1): v(O–H), 3416 (br), 1626(m); v(Ag–O), 1412 (w); v(P–O), 1134 (s), 1081 (s); v(W–O), 1015 (w), 926 (br). Elemental analysis calcd (found) for AgH2L2Li2K3Ag21[P8W66O251]Cl2: Ag 2.7 (2.9), W 54.9 (55.1), K 2.92 (3.17), Li 0.43 (0.43). TGA water loss from room temperature to 400°C, calc (found): %: 19.0 (19.5).

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**Conflict of interest**

The authors declare no conflict of interest.

**Keywords**

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Controlling the Reactivity of the \([\text{P}_8\text{W}_{48}\text{O}_{184}]^{40-}\) Inorganic Ring and Its Assembly into POMZite Inorganic Frameworks with Silver Ions

Familienachwuchs: Ein Polyoxometallatcluster auf Basis des superlakunären Heteropolyanions \([\text{P}_8\text{W}_{48}\text{O}_{184}]^{40-}\) (abgekürzt als \(\text{P}_8\text{W}_{48}\)) reagiert mit Silberionen zu drei neuen, rein anorganischen, porösen Metalloxidmaterialien, welche die Familie zeolithischer Polyoxometallatgerüste erweitern. Die Silberionen ermöglichen sowohl Reaktionen an den Clustern als auch deren Verknüpfung.