Polyoxometalates

Classical/Non-classical Polyoxometalate Hybrids

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Abstract: Two polyanions [Se$_{12}$W$_{4}$(H$_{2}$O)$_{10}$H$_{2}$]$^{11-}$ and [Se$_{14}$Pd$_{4}$W$_{31}$O$_{90}$H$_{12}$]$^{12-}$ are the first hybrid polyoxometalates in which classical (Group 5/6 metal based) and non-classical (late transition-metal based) polyoxometalate units are joined. Requiring no supporting groups, this co-condensation of polyoxotungstate and isopolyoxopyrrolate constituents also provides a logical link between POM-Pd$^{2+}$ coordination complexes and the young subclass of polyoxopalladates. Solid-state, solution, and gas-phase studies suggest interesting specific reactivities for these hybrids and point to several potential derivatives and functionalization strategies.

The chemistry of palladium-containing polyoxometalates (POMs) has experienced impressive development over the past decade,[1,2] with progress primarily concentrated on two areas. The first is defined by conventional Pd$^{2+}$ coordination complexes of lacunary polyoxotungstates (POTs), [Pd$_{x}$($X$$_{y}$W$_{z}$O$_{p}$)$_{q}$]$^{x-}$, where Pd$^{2+}$ ions in square-planar environments coordinate oxygen atoms of vacant sites of POT ligands, resulting in a diverse range of structures incorporating one to four Pd$^{2+}$ centers.[2] Such species are convenient precursors for highly stable suspensions of POT-stabilized Pd$^{2+}$ nanoparticles, which can be obtained at mild conditions in aqueous media.[3] Some of the Pd-POT complexes were also shown to act as pre-catalysts for various organic transformations.[2m,4] In these complexes, the Pd$^{2+}$ centers typically lack a direct connection, with the only exception in [Pd$_{2}$($X$-$P$$_{2}$W$_{13}$O$_{37}$)$_{3}$]$^{18+}$, where two out of four Pd$^{2+}$ ions are bridged via O atoms of two phosphate groups.[5,6]

In the second main area, formed by so-called polyoxopalladates (POPs), the Pd$^{2+}$ centers, in contrast, act as addenda ions themselves. Here, the elementary PdO$_{4}$ building blocks are condensed via corners and edges, typically also involving external RXO$_{2}^{2-}$ heterogroups stabilizing the discrete (PdO$_{4}$)$_{n}$ entity.[6,7] About 50 of these non-classical POMs are known today, incorporating up to 84 Pd$^{2+}$ ions. One of the most stable POPs described comprises species of general composition [MPd$_{14}$O$_{59}$$(RXO$_{2}$)$_{n}$]$^{14-}$ ((MPd$_{14}$)$_{2}$), where a heterometal ion M$^{2+}$ in a cubic O$_{59}$ environment is encapsulated in the cuboid-shaped (Pd$_{14}$O$_{59}$$(RXO$_{2}$)$_{n}$) shell (RX = Se$_{4}$$^{2-}$, OAs$^{2-}$, PhAs$^{2-}$, OP$^{2-}$, PhP$^{2-}$).[6]

Recently, Cronin and co-workers also reported several polyaniions that can be considered as complexes of seleno- and telurotungstates ($X$$_{y}$W$_{z}$O$_{p}$) with selenite- or tellurate-supported multinuclear Pd$^{2+}$-based fragments.[7] In two isomeric [H$_{2}$Pd$_{10}$Se$_{8}$W$_{12}$O$_{39}$]$^{16-}$ polyanions, two {Pd$_{6}$Se$_{6}$O$_{14}$}$_{2-}$ units are coordinated to (β-se-W$_{9}$O$_{24}$)$_{2-}$ and (γ-se-W$_{9}$O$_{24}$) POT moieties. In [Pd$_{4}$Te$_{19}$Se$_{8}$W$_{12}$O$_{39}$]$^{16-}$ two identical {Pd$_{4}$Te$_{3}$O$_{12}$}$_{2-}$ groups are stabilized by six (ε-Te$_{6}$W$_{12}$O$_{39}$) lacunary POTs.[7]

Yet, up to now there was no systematic investigation on how to achieve commensurate reaction conditions that allow to co-condense, and thus cleanly interface, classical POTs and non-classical POPDs. We thus explored the possibility to prepare hybrid polyoxopalladatotungstates ($X$$_{y}$Pd$_{m}$W$_{n}$O$_{p}$)$_{q}^{x-}$, where both Pd$^{2+}$ and W$^{6+}$ centers act as addenda centers of their individual POM units, without the need for any additional external stabilizing groups. Herein we report two first examples of such hybrid palladatotungstates, [Se$_{14}$Pd$_{4}$W$_{4}$$(H$_{2}$O)$_{10}$]$^{11-}$ (1) and [Se$_{16}$Pd$_{4}$W$_{31}$O$_{90}$H$_{12}$]$^{12-}$ (2), crystallized as hydrated mixed cesium/sodium salts Cs$_{3}$Na$_{2}$[Se$_{16}$Pd$_{4}$W$_{31}$O$_{90}$H$_{12}$]·18H$_{2}$O·0.3CsOAc·0.2NaOAc (CsNa$_{1}$; OAc = acetate) and Cs$_{3}$Na$_{2}$[Se$_{16}$Pd$_{4}$W$_{31}$O$_{90}$H$_{12}$]·30H$_{2}$O (CsNa$_{2}$), respectively, and their characterization in the solid state, aqueous solutions, and gas phase.

The polyanions 1 and 2 have been prepared in reactions of [Se$_{16}$W$_{31}$O$_{90}$H$_{12}$]$^{14+}$ ([Se$_{16}$W$_{31}$O$_{90}$]$^{16+}$ with Pd$^{2+}$ nitrate in different aqueous media (Supporting Information, Scheme S1). The (Se$_{16}$W$_{31}$) precursor possesses a cyclic structure, where three (γ-se-W$_{9}$O$_{24}$) units are alternating with three trans-O(con-W$_{9}$O$_{24}$) groups. In aqueous solution it slowly decomposes, releasing (Se$_{6}$W$_{18}$O$_{42}$) fragments[8] and thus could act as a source for preparation of diverse tungstoselenite complexes.[9] The Cs$^{+}$ counterions seem to play an important role for isolation of 1 and 2 as pure crystalline materials owing to relatively low solubility of the hydrated Cs$^{+}$ salts. Alternatively, a Rb$^{+}$/Na$^{+}$ salt of 1 can be successfully prepared by replacing CsNO$_{3}$ with RbNO$_{3}$ in the synthesis of CsNa$_{1}$. With no additional counterions only the hydrated sodium salt of paratungstate-8 ([H$_{2}$W$_{12}$O$_{40}$]$^{12+}$).
could be isolated from the reaction medium for preparation of 1 as a crystalline product. The paratungstate-B salt is also sometimes present as an impurity to CsNa-1, which could be purified in this case by recrystallization from 0.25 M NaOAc aqueous solution (pH 6.7). Similar recrystallization of CsNa-2 leads to formation of a mixture of CsNa-1, CsNa-2, and other undefined products. The purity and composition of the compounds was further confirmed by elemental analysis, PXRD, TGA, and XPS (see the Supporting Information for details).

CsNa-1 crystallizes in the orthorhombic space group Pnnm. The polyanion 1 exhibits idealized C\textsubscript{2v} symmetry and comprises an [α-Se\textsubscript{2}W\textsubscript{4}O\textsubscript{13}]\textsuperscript{2-} POT moiety ([α-Se\textsubscript{2}W\textsubscript{4}]) supporting a [Pd\textsubscript{4}O\textsubscript{4}] fragment (Figure 1).

\[ \text{Figure 1. Structure of 1 (a) and the (H\textsubscript{2}O)Na-1 associate (b); comparison with the [Pd\textsubscript{4}O\textsubscript{4}(RKO\textsubscript{4})\textsubscript{3}]\textsuperscript{2-} fragment (c) in the cuboid-shaped polypseudopolyanolate } \]

\[ \text{[MPd}_{12}\text{O}_{28}(\text{RKO}_{14})_{3}]^{2-} \]

The (α-Se\textsubscript{2}W\textsubscript{4}) unit can be compared to a hypothetical tetralacunary Wells–Dawson-type (α-P\textsubscript{4}W\textsubscript{11}O\textsubscript{39}) fragment (Supporting Information, Figure S3), with two neighboring [W\textsubscript{11}O\textsubscript{39}] groups, composed of two edge-shared (WO\textsubscript{4}) octahedra, removed from the inner (W\textsubscript{i}) belts of [α-P\textsubscript{4}W\textsubscript{11}O\textsubscript{39}]; Supporting Information, Figure S3a/b), each one from one belt. The Se\textsuperscript{IV} ions in [α-Se\textsubscript{2}W\textsubscript{4}] adopt a trigonal pyramidal environment with the outwards oriented lone pair (Supporting Information, Figure S3d; Se–O 1.677(16)–1.725(15) Å). The formation of [α-Se\textsubscript{2}W\textsubscript{4}] from the [γ-Se\textsubscript{2}W\textsubscript{4}] precursor requires attachment of two additional W\textsuperscript{VI} ions to [γ-Se\textsubscript{2}W\textsubscript{4}], each of which is completing the outer (W\textsubscript{o}) cap of the POT fragment, combined with [γ-Se\textsubscript{2}W\textsubscript{4}] isomerization by rotation of both (W\textsubscript{i}) caps by 60° (Supporting Information, Figure S3). The same [α-Se\textsubscript{2}W\textsubscript{4}] building blocks have been recently observed in [Fe\textsubscript{2}Se\textsubscript{2}W\textsubscript{12}O\textsubscript{42}(OH)\textsubscript{4}]\textsuperscript{18-} polyanions.\textsuperscript{28} At the same time, the arrangement of W\textsuperscript{VI} centers in [α-Se\textsubscript{2}W\textsubscript{4}] is different from that in the actual (α-P\textsubscript{4}W\textsubscript{11}O\textsubscript{39}) moieties that, for example, form [H\textsubscript{17}Fe\textsubscript{3}P\textsubscript{4}W\textsubscript{11}O\textsubscript{39}]\textsuperscript{16-} and [(W\textsubscript{11}Mn\textsubscript{4}O\textsubscript{18})(P\textsubscript{4}W\textsubscript{11}O\textsubscript{39})\textsuperscript{38-} complexes.\textsuperscript{29} In fact, these (α-P\textsubscript{4}W\textsubscript{11}O\textsubscript{39}) building blocks are the structural isomers to the hypothetical (α-P\textsubscript{2}W\textsubscript{11}O\textsubscript{39}) units discussed above, and can be obtained from (α-P\textsubscript{4}W\textsubscript{11}O\textsubscript{39}) by removing not (W\textsubscript{11}O\textsubscript{39}) but rather corner-sharing (W\textsubscript{11}O\textsubscript{39}) units from its inner belts (Supporting Information, Figure S3c). It is also different in γ-Se\textsubscript{2}W\textsubscript{4} moieties constructing the reported [H\textsubscript{3}P\textsubscript{4}Se\textsubscript{2}W\textsubscript{12}O\textsubscript{42}]\textsuperscript{2-} (see above) and [Fe\textsubscript{2}Se\textsubscript{2}W\textsubscript{12}O\textsubscript{42}(OH)\textsubscript{4}]\textsubscript{28–29} complexes where the two (W\textsubscript{i}) caps are rotated by 60° relative to their orientation in the α isomer (Supporting Information, Figure S3e).

The four Pd\textsuperscript{II} centers in the [Pd\textsubscript{4}O\textsubscript{4}] fragment form a rectangle (Pd–Pd 3.360(2)–3.375(2) Å) and are linked by four μ-O sites (Figure 1a). All square-planar Pd\textsuperscript{II}O\textsubscript{4} (Pd–O 1.976(14)-2.010(15) Å) include two cis-positioned μ-O of the [Pd\textsubscript{4}O\textsubscript{4}] fragment as well as two Pd-O-μ atoms; two Pd\textsuperscript{II} centers bind to the (W\textsubscript{i}) caps and two to the belts of (α-Se\textsubscript{2}W\textsubscript{4}) (Figure 1a). Based on bond valence sums, the proton in 1 is disorder over the four μ-O atoms linking the Pd\textsuperscript{II} centers. These oxygens also coordinate to a (Na(OH)\textsubscript{3})\textsuperscript{+} counterion (Figure 1b; Na–O 2.42(2)–2.53(2) Å).

The direct connection between the Pd\textsuperscript{II} centers by oxo ligands as well as the complete integration of the POPd (Pd\textsubscript{4}O\textsubscript{4}) moiety in the POM framework allow to consider 1 as a genuine hybrid polypseudopolyanolatostate. Interestingly, the structure of [Pd\textsubscript{4}O\textsubscript{4}] unit in 1 compares to the [Pd\textsubscript{4}O\textsubscript{4}(RKO\textsubscript{4})\textsubscript{3}]\textsuperscript{2-} face in the cuboid-shaped (MPd\textsubscript{4}O\textsubscript{4}) POPds (Figure 1c/d), with the RKO\textsubscript{4}– groups stabilizing the [MPd\textsubscript{4}O\textsubscript{4}] core replaced by (α-Se\textsubscript{2}W\textsubscript{4}). Moreover, the Na\textsuperscript{+} attachment to [Pd\textsubscript{4}O\textsubscript{4}] in 1 is similar to the connection mode between the central M\textsuperscript{2+} ion and the [Pd\textsubscript{4}O\textsubscript{4}(RKO\textsubscript{4})\textsubscript{3}] face in [MPd\textsubscript{4}] nanocubes (Figure 1b/d). This suggests that the [Pd\textsubscript{4}O\textsubscript{4}] group in 1 possesses reactivity towards oxophilic heterometals.

The total number of metal centers in 1 allows for an analogy between 1 and Wells–Dawson-type polyanions (α-P\textsubscript{4}W\textsubscript{11}O\textsubscript{39})\textsuperscript{2+} Both POMs comprise two central heteroatoms surrounded by 18 addenda metal ions. However the [Pd\textsubscript{4}] rectangle in 1 is rotated by 45° in comparison to the [W\textsubscript{11}O\textsubscript{39}] rectangle in (α-P\textsubscript{4}W\textsubscript{11}O\textsubscript{39}) if the latter is formally decomposed into the above-mentioned hypothetical (α-P\textsubscript{4}W\textsubscript{11}O\textsubscript{39}) fragment and four W\textsubscript{VI} centers (Supporting Information, Figure S4), possibly enforced by the square-planar Pd coordination mode in 1 relative to the octahedral W\textsubscript{VI}O\textsubscript{4} groups. This analogy prompted us to probe the possibility to form lacunary derivatives of 1 at conditions similar to those for formation of (α-Se\textsubscript{2}W\textsubscript{4}) and (α-P\textsubscript{4}W\textsubscript{11}O\textsubscript{39}) from (α-Se\textsubscript{2}W\textsubscript{4}). These experiments, however, only resulted in Cs\textsubscript{4}Na\textsubscript{2}[H\textsubscript{3}P\textsubscript{4}Se\textsubscript{2}W\textsubscript{12}O\textsubscript{42}(SeO\textsubscript{4})\textsubscript{3}]·ca.20H\textsubscript{2}O·POPd\textsubscript{3+}\textsuperscript{2+},\textsuperscript{22} which suggests that decomposition of 1 proceeds first through release of Pd\textsuperscript{II} ions, followed by POT decomposition.

However the possibility of existence of unstable lacunary derivatives of (α/β/γ/δ-Se\textsubscript{2}Pd\textsubscript{4}W\textsubscript{4}) polyanions is evident from the structure of 2 obtained indirectly by reaction of (Se\textsubscript{2}W\textsubscript{4}) with Pd\textsuperscript{II} in water. The compound CsNa-2 crystallizes in the triclinic space group P1. The unit cell in CsNa-2 contains two identical polyanions 2, each of which can be imagined as a dimer of two γ-(H\textsubscript{2}O)(OH)\textsubscript{2}Pd\textsubscript{4}Se\textsubscript{2}W\textsubscript{12}O\textsubscript{42} (γ-Pd\textsubscript{4}Se\textsubscript{2}W\textsubscript{4}) units connected by two trans-O–(H\textsubscript{2}O) groups (Figure 2). In line with the
are highlighted in light purple, while ion is bound to centers. It is interesting to niti ns CsNa-2 exhibits two main signals at 1316.5 ppm and 1226.8 ppm and is significantly upfield-by (Supporting Information, Figure 2) and (1223.8 ppm)-cen- CsNa-1 77 1 T he monopro- 77 Tcations, the compounds 2 are only slightly soluble in water; howe- 77 moity (b) in the polyanion in aqueous medium in and with the observation of a singlet at 1202 ppm in the LiOAc solution (pH 6.2) exhibits as singlet at 1225.3 ppm (Figure 3), consistent with the presence of only one symmetrically non-equivalent SeV ion in the crystal structure of CsNa-1 and with the observation of a singlet at 1202 ppm in the 77 Se MAS NMR for this compound (Supporting Information, Figure S12). This indicates stability of 1 in aqueous medium in saturated solutions. The observed chemical shift is commensurate with those of ZnII (1222.5 ppm) and LuIV (1223.8 ppm)-centered cuboid (MPd4Se6) POPd6[8a] and is significantly upfield-shifted compared to an aqueous SeO2 solution (pH 6.4; 1316.3 ppm). For comparison with other tungstoselenites, the (SeW6) precursor (unstable in solution) gives a broad peak centered at 1289.1 ppm in 77 Se MAS NMR[8a].

The 77 Se MAS NMR of CsNa-2 (Supporting Information, Figure S13) shows two broad signals centered at 1255 and 1187 ppm (verified for two different spinning frequencies), in line with the symmetry of 2. Based on literature data for (SeW6) [8a] and the data obtained for CsNa-1 (see above), we tentatively assign the upfield signal to SeIV ions of the (Pd3Se14) half of the (γ-Pd3Se14) subunit (Figure 2a), and the 1255 ppm peak to the SeV ions positioned in the Pd6-free (SeW6) part of this motif. In contrast to 1, solution 77 Se NMR of 2 exhibits two main signals at 1316.5 ppm and 1226.8 ppm with 1:8:1 relative intensities (Supporting Information, Figure S14). The chemical shifts of the signals are evident of decomposition of the polyanions with the release of selenite ions (signal at 1316.5 ppm) concurrent with formation of 1 (singlet at 1226.8 ppm), in line with the formation of CsNa-1 crystals after recrystallization of CsNa-2 from aqueous acetate solutions. These solution stability observations for 1 and 2 are further supported by STEM images obtained after drop-casting of 10-4 M CsNa-1 and CsNa-2 solutions in ultra-pure water onto HOPG surface (Supporting Information, Figure S5).

The exact composition of ion pairs based on 1 and 2 that potentially exist in solutions and gas phase was probed by previous discussion, the (SeW6) structure can be understood as a (γ-Se2W3) unit, present in (Se2W6), binding a WIV to complete one of the (Wj) caps or, alternatively, as a (γ-Se2W3)(O=W(H2O)) (Supporting Information, Figure S3e), missing one WIV ion in its (Wj) cap. The two Pd2 ions in (γ-Pd2Se2W3) assume a square planar environment, each coordinating two cis-positioned oxygens of (γ-Se2W3): one from the WIV ion in the (Wj) cap and one from the (Wj) belt (Figure 2a). Furthermore, the two Pd2 ions are μ2-OH-bridged. One of the Pd2 ions additionally coordinates a terminal H2O, and its μ2-O (Pd, W) ion in the trans-position to the aqua ligand is protonated (Figure 2a/c; Supporting Information, Table S4). The second Pd2 ion is bound to trans-(O=O=W(H2O)) group through the μ2-O (Figure 2b).

Thus, the (γ-Pd2Se2W3) fragment can be considered as a lacunary derivative of a hypothetical polyanion (γ-Pd3Se2W3) polyanion, lacking two Pd2 and one WIV centers. It is interesting to note that the orientation of (Pd3O4) fragment in this (γ-Pd3Se2W3) POM, in case it exists, would be similar to that in [α-P3W11OH20]4− and not in [α-P3Pd3Se;W3OH20]. Along with the μ2-O ligand connecting it to Pd2 (see above), the WIV center of each trans-(O=O=W(H2O)) group also binds to an O atom of the neighboring (Wj) belt of the same (γ-Pd2Se2W3) monomeric unit as well as to the two O atoms of the incomplete (Wj) cap group of the second (γ-Pd2Se2W3) monomer, each of which belongs to different WIV ions (Figure 2b/c). Interestingly, one of the H2O ligands of the trans-(O=O=W(H2O)) groups is directed inward the polyanion, while the second one is pointed outward (Figure 2c). Thus, considering the protonation sites, 2 is of C1 symmetry. Otherwise, it would possess a C2 axis passing through the center of a line connecting the WIV centers of the two (O= W(H2O)) groups (Figure 2c).

Owing to the presence of large Cs+ cations, the compounds CsNa-1 and CsNa-2 are only slightly soluble in water; however, their solubility is significantly increased in 0.25–0.5 M sodium and lithium acetate solutions (pH 6–7), especially upon heating to 65–70 °C. This allowed assessment of the solution behavior of 1 and 2 by 77 Se NMR and UV/Vis spectroscopy (see the Supporting Information). Room-temperature 77 Se NMR of 1 in 0.25 M LiOAc solution (pH 6.2) exhibits a singlet at 1225.3 ppm (Figure 3), consistent with the presence of only one symmetrically non-equivalent SeV ion in the crystal structure of CsNa-1 and with the observation of a singlet at 1202 ppm in the 77 Se MAS NMR for this compound (Supporting Information, Figure S12). This indicates stability of 1 in aqueous medium in saturated solutions. The observed chemical shift is commensurate with those of ZnII (1222.5 ppm) and LuIV (1223.8 ppm)-centered cuboid (MPd4Se6) POPd6[8a] and is significantly upfield-shifted compared to an aqueous SeO2 solution (pH 6.4; 1316.3 ppm). For comparison with other tungstoselenites, the (SeW6) precursor (unstable in solution) gives a broad peak centered at 1289.1 ppm in 77 Se MAS NMR[8a].

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The exact composition of ion pairs based on 1 and 2 that potentially exist in solutions and gas phase was probed by
mass spectrometry. The negative-ion-mode ESI-MS spectrum of 1 (Figure 4) shows a set of peaks (III–VII), which can be attributed to various ion pairs \([\text{H}_2\text{Na}_{1.5}\text{Se}_2\text{Pd}_7\text{W}_{14}\text{O}_{67}^\text{III}]^\text{+}\) based on the intact polyanion 1 (Table 1), by virtue of their \(m/z\) values and analysis of the corresponding calculated and observed isotope envelopes (see Figure 4, inset; Supporting Information, Figures S17–S24). Peak II could be attributed to an ion pair based on a monovacant derivative of 1, where one of the Pd\(^{II}\) centers is missing, while peak I belongs to a dilacunary species lacking two Pd\(^{IV}\) ions with the \(\mu_2\)-bridging oxygen ion linking these metal ions together. This suggest that decomposition of 1 in gas phase (and possibly also in solution) proceeds via release of Pd\(^{IV}\) centers in a first step.

This is consistent with our observations of loss of Pd\(^{IV}\) ions and the following POT moiety decomposition during our attempts to prepare lacunary derivatives of 1, but also suggests that such species could in principle exist if adequately stabilized. The ESI-MS spectrum of 2 recorded at similar conditions (Supporting Information, Figure S25) only exhibits peaks attributed to singly charged POM decomposition products (see the Supporting Information for details), consistent with our NMR observations.

In summary, we have isolated and characterized two polyanions \([\text{Se}_2\text{Pd}_7\text{W}_{14}\text{O}_{67}^\text{III}]^\text{+}\) and \([\text{Se}_2\text{Pd}_7\text{W}_{14}\text{O}_{67}^\text{IV}]^\text{2-}\) comprising both W\(^{VI}\) and Pd\(^{IV}\) addenda sites. As such, the new hybrid species bridge the conventional POT-Pd\(^{IV}\) coordination complexes and POPds. The analysis of the structural data for CsNa-1 suggests reactivity of Pd\(^{IV}\) ions bridging Pd\(^{IV}\) ions in its (Pd\(_2\)O\(_4\)) fragment towards oxophilic metals. Hence, the (Pd\(_2\)O\(_4\)) site in 1 could serve an analogy to a vacant site of lacunary metal oxides, suggesting that decomposition of these metal ions together. This suggest that decomposition of 1 in gas phase (and possibly also in solution) proceeds via release of Pd\(^{IV}\) centers in a first step.

### Experimental Section

**Synthesis of CsNa-1:** Samples of Na\(_2\)\([\text{H}_2\text{Se}_2\text{Pd}_7\text{W}_{14}\text{O}_{67}^\text{IV}]^\text{4+}\) (0.105 g, 0.423 mmol) were dissolved in 5 mL of aqueous 0.5 M NaOAc solution (prepared by addition of solid NaOH into 0.5 M H\(_2\)O\(_4\) solution in water until pH reaches 6.7) under vigorous stirring and heating at about 50–60 °C. The obtained clear dark red reaction mixture was stirred at 50 °C for 30 min and then cooled to room temperature. After that 0.5 mL of 1 M Cs\(_2\)NO\(_3\) solution in H\(_2\)O was added to the reaction mixture under stirring leading to immediate formation of light-brown precipitate. The precipitate was collected by filtration and recrystallized from warm 0.25 M NaOAc medium (pH 6.7) resulting in an orange solution. Needle-like brown-yellow crystals of CsNa-1 form within several days. The filtrate produced additional portion of CsNa-1, although often contaminated by hydrated Cs/Na salt of paratungstate-8 (based on IR and single-crystal XRD). In this case purification is achieved by recrystallization of the obtained solid material from 0.25 M NaOAc medium (pH 6.7). The crystals of the product were collected by filtration and washed with small amount of ice cold water. Total yield: 0.177 g (33 % based on Pd).

Elemental analysis calcd (%) for C\(_{1404}\)H\(_{297}\)O\(_{516}\)Pd\(_{28}\)Se\(_{14}\)W\(_{108}\)Na\(_{10}\): C 52.6, H 5.5, O 26.9, N 3.8, Pd 4.3, Se 4.3. CSD no.: 814740. IR (KBr tablet), \(\nu\) cm\(^{-1}\): 3424 (s, br); 1625 (m); 1108 (w); 1078 (w); 934 (s); 902 (s, br); 874 (s); 840 (s); 819 (s); 774 (s); 713 (s); 676 (s); 502 (s); 451 (s). Raman (solid sample, \(\lambda_{\text{exc}} = 1064 \text{ nm}, \nu \text{ cm}^{-1}: 958\) (s); 891 (m); 872 (m); 835 (m); 787 (w); 582 (w); 417 (m); 191 (m); 161 (m, br); 130 (m); 100 (m); 75 (m). \(^{75}\)Se MAS NMR (H\(_2\)O/D\(_2\)O): 1225.3 ppm. \(^{75}\)Se MAS NMR: 1202 ppm. UV/Vis (0.25 M NaOAc buffer solution, pH 6.7): \(\lambda_{\text{max}} (\nu) = 227 (74450), 273 (shoulder, 34153), 414 nm (1484 mol\(^{-1}\) dm\(^{-1}\) cm\(^{-1}\)). CSD no.: 431484.

**Synthesis of CsNa-2:** \(\text{Na}_2\)\([\text{H}_2\text{Se}_2\text{Pd}_7\text{W}_{14}\text{O}_{67}^\text{IV}]^\text{4+}\) (0.200 g, 0.017 mmol) and Pd(NO\(_3\))\(_2\)\(\cdot\)H\(_2\)O (0.026 g, 0.105 mmol) were dissolved in 2 mL of H\(_2\)O under vigorous stirring and heating at about 50–60 °C. After the dissolution of all the reagents, the reaction mixture was stirred and further heated for 1 h and then cooled to room temperature and filtered. Three drops of 1 M aqueous Cs\(_2\)NO\(_3\) solution were added to the obtained dark red–brown filtrate. The
obtained pale brown precipitate was filtered and the evaporation of the resulting solution at room temperature led to brown crystalline material of CsNa-2 within 1–3 days. Crystals were collected by filtration, washed with ice-cold water and dried in air. Yield: 0.040 g (17% based on W).

Elemental analysis calcd (%) for CsNa-2: C, 3.31; H, 0.61; Na, 0.61; Pd, 4.49; Se, 3.33; W, 4.24. Found: C, 3.22; H, 0.61; Na, 0.61; Pd, 4.49; Se, 3.31; W, 4.25. IR (KBr pellet), ν (cm⁻¹): 3432 (s, br); 1614 (s); 954 (s); 843 (s); 768 (s); 704 (s); 662 (s, br); 491 (m); 427 (s). Raman (solid sample, ω = 1064 nm), ν (cm⁻¹): 970 (s); 914 (m); 902 (m); 885 (m); 866 (w, sh); 812 (m); 717 (w); 660 (m); 646 (m); 513 (m); 503 (w); 216 (m); 110 (m); 75 (m).

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