Tetrapalladium-Containing Polyoxotungstate $[Pd^{II}_{4}(\alpha-P_2W_{15}O_{56})_2]^{16-}$: a Comparative Study

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ABSTRACT The novel tetrapalladium(II)-containing polyoxometalate $[Pd^{II}_{4}(\alpha-P_2W_{15}O_{56})_2]^{16-}$ has been prepared in aqueous medium and characterized as its hydrated sodium salt Na₁₆[Pd₄(α -P₂W₁₅O₅₆)₂]·71H₂O by single-crystal XRD, elemental analysis, IR, Raman, multinuclear NMR and UV-Vis spectroscopy. The complex exists in *anti* and *syn* conformations which form in a 2:1 ratio and possesses unique structural characteristics in comparison with known {M₄(P₂W₁₅)₂} species. ³¹P and ¹⁸³W NMR spectroscopy are consistent with long-term stability of the both isomers in aqueous solutions.

INTRODUCTION

Pd^{II} complexes with polyoxotungstates (POTs) have attracted considerable attention during the last years as potential catalysts or precatalysts for various low-temperature transformations of organic substrates.^{1,2} On the other hand, the square-planar Pd^{II} coordination environment (in contrast to octahedral coordination characteristic for the first-row transition metals, the reactivity of which towards polyoxometalates (POMs) is widely investigated) opens possibilities to design polyanions with unique, so far not observed, structures and properties. In this context, a novel class of polyanions based exclusively on Pd^{II} and Au^{III} centers has been discovered in the past decade,^{2b,3} and very recently several unusual seleno- and tellurotungstates incorporating multinuclear Pd^{II}-based fragments reminiscent building blocks which constitute polyoxopalladate structures have been reported.⁴

Nevertheless the number of structurally characterized Pd-containing polyanions remains quite small in comparison with that of POMs incorporating 3*d* metals and lanthanides and the other examples include only a few Pd^{II} complexes with lacunary POTs ⁵⁻⁹ as well as 1D polymers where Pd^{II} ions link paratungstate species into infinite chains.¹⁰ In this respect it was shown that monolacunary Lindqvist-, Keggin- and Wells-Dawson-type POTs form polyanions $[Pd^{II}_{2}(W_{5}O_{18})_{2}]^{8-}$, $[Pd^{II}_{2}(\alpha-PW_{11}O_{39})_{2}]^{10-}$ and *syn/anti*- $[Pd^{II}_{2}(\alpha_{2}-P_{2}W_{17}O_{61})_{2}]^{16-}$, respectively, where two Pd^{II} ions in a square-planar environment link together the two POM ligands.⁵ When the dilacunary derivatives of Keggin-type silicotungstate are reacted with Pd^{II} acetate, they form monomeric [γ -H₂SiW₁₀O₃₆Pd₂(CH₃COO)₂]⁴⁻ species there two Pd^{II} centers grafted on the vacant site of the POT are additionally bridged by two acetates.^{1m} Interaction of this complex with dicarboxylates led to carboxylate metathesis and formation of dimeric assemblies of the constitution [{(γ -H₂SiW₁₀O₃₆Pd₂)(O₂C(CH₂)_nCO₂)}₂]⁸⁻ (*n* = 1, 3, 5).⁶ A number of sandwich-like

complexes with general formulae $[Pd_{3-y}(WO_2)_y(XW_9O_{34})_2]^{z^-}$ (E = P^V, Si^{IV}) ⁷ and $[Pd_{3-y}(WO_2)_y(XW_9O_{33})_2]^{z^-}$ (X = As^{III}, Sb^{III}, Te^{IV})⁸ are built by two trilacunary Keggin-type {XW₉} units linked either via three Pd^{II} centers (*y* = 0) or via a belt comprising one Pd^{II} and two WO₂ groups (*y* = 2) or two Pd^{II} centers and one WO₂ group (*y* = 1), depending on the exact reaction conditions. Interaction of Pd^{II} with trilacunary bismuthotungstate $[BiW_9O_{33}]^{n^-}$ resulted in the Krebs-type structure $[Pd_3(H_2O)_9Bi_2W_{22}O_{76}]^{8^-}$ where Pd^{II} ions are only weakly bound to the POT surface.⁹

At the same time almost no attention has been given to the reactivity of Pd^{II} towards lacunary derivatives of Wells-Dawson-type POMs, and the above-mentioned *syn* and *anti* isomers of $[Pd^{II}_2(\alpha_2-P_2W_{17}O_{61})_2]^{16^-}$ are the only known structurally characterized Pd^{II} complexes with POTs of this structural type. Herein we report a novel sandwich-like polyanion $[Pd_4(\alpha-P_2W_{15}O_{56})_2]^{16^-}$ (1) exhibiting unique structural features, which was obtained in the reactions of Pd^{II} ions with trilacunary Wells-Dawson type POT $[\alpha-P_2W_{15}O_{56}]^{12^-}$ and isolated as the hydrated sodium salt $Na_{16}[Pd_4(\alpha-P_2W_{15}O_{56})_2] \cdot 71H_2O$ (**Na-1**) and the tetrabutylammonium salt $[(C_4H_9)_4N]_{15}[HPd_4(\alpha-P_2W_{15}O_{56})_2]$ (**TBA-1**).

RESULTS AND DISCUSSION

Synthesis. The polyanion **1** self-assembles in the reaction of Pd^{II} nitrate and $[\alpha-P_2W_{15}O_{56}]^{12-1}$ in 0.5 M CH₃COONa medium in the pH range of 2 – 8 and the temperature range of 5 to 80 °C. Variation of the $Pd^{II} : P_2W_{15}$ ratio from 3 : 1 to 1 : 1 also does not influence the composition of the final product which is crystallized from the $Pd^{II} / \{P_2W_{15}\} / 0.5$ M CH₃COONa reaction systems, and **Na-1** has been isolated in all the cases as based on ³¹P NMR and IR spectroscopy as well as unit cell measurements. **Na-1** is well soluble in water (> 0.25 g / 1 ml) at room

temperature and is also soluble in 1 : 1 mixtures of $H_2O/(CH_3)_2CO$ and H_2O/CH_3CN mixtures, and can be repeatedly recrystallized from water and 0.5 M CH₃COONa (pH 4.2).

The tetrabutylammonium salt, **TBA-1**, soluble in common organic solvents (*e.g.* CH₃CN, $(CH_3)_2CO$, CH_2Cl_2 *etc*), was prepared by dropwise addition of an aqueous solution of **Na-1** to an aqueous solution of TBAHSO₄, followed by washing of the obtained precipitate with plenty of water and its identity and purity was confirmed using IR, ³¹P NMR and C, H, N analysis.

Crystal structure analysis. Compound **Na-1** crystallizes in the triclinic symmetry in the space group *P*-1. The polyanions **1** possess a sandwich-like structure where two phosphotungstate units $[\alpha - P_2 W_{15}O_{56}]^{12-} (= \{P_2 W_{15}\})$ are linked via a belt of four Pd^{II} centers (Fig. 1).

There is a complex disorder of $[Pd_4(\alpha-P_2W_{15}O_{56})_2]^{16^-}$ polyanions in the crystals of Na-1 implying (1) a rotation of $\{P_2W_{15}\}$ ligands by 60° which results in two symmetrically independent positions of $\{P_2W_{15}\}$ with the relative occupancies of 66 and 34%, respectively; and (2) a rotation of the Pd₄ belt by +60° and -60° resulting in three symmetrically independent positions for the Pd₄ thomb with the relative occupancies of 66% (non-rotated), 23% (rotated to +60°) and 11% (rotated to -60°). Such kind of disorder implies, on one hand, that part of the polyanions $[Pd_4(\alpha-P_2W_{15}O_{56})_2]^{16^-}$ can be turned relative to each other by either +60° or -60° during their packing in the crystals of Na-1. One the other hand, such hypothesis does not explain the non-equal distribution of the "rotated" and "non-rotated" positions for 1, which is exactly reproducible from crystal to crystal. Another explanation is the turn of only one of the $\{P_2W_{15}\}$ ligands in part of polyanions 1 relative to the remaining $\{Pd_4P_2W_{15}\}$ unit by 60° which results in *syn/anti* isomerism with the *anti-1* to *syn-1* isomers ratio of 2 : 1, respectively (66% *vs* 34). It should be noted that the rotation of $\{P_2W_{15}\}$ *vs* $\{Pd_4P_2W_{15}\}$ by either +60° or -60° would

equally lead to the *syn* derivative. The second scenario and the presence of both *anti* and *syn* isomers in this exact ratio have been confirmed by 31 P and 183 W NMR spectroscopy (*vide infra*).

The {P₂W₁₅} ligands in **1** have typical Wells-Dawson structure with one W₃O₆ "capping" group missing and consist of two central tetrahedral PO₄ templates surrounded by 15 corner- and edge-shared WO₆ octahedra (Fig. 1). The polyanions possess $C_{3\nu}$ symmetry with the C_3 axis passing through the two P^V ions. The 15 WO₆ octahedra can be formally divided into a W₃O₆ "cap" (gray in Fig.1) and an "inner" W₆O₂₇ "belt" (dark-blue) assembled around one of the two P^AO₄ units (yellow tetrahedra), and an "outer" W₆O₂₇ "belt" (light-blue) placed around the second P^BO₄ group (orange). Due to the missing second W₃O₆ "cap" the phosphotungstate {P₂W₁₅} offers a so-called vacant or lacunary site with seven nucleophilic oxygen atoms suitable for coordination to various heterometals. Six of these oxygens coordinated to six W^{VI} centers of the "outer" W₆O₂₇ "belt" form a nearly regular hexagon (O···O_{average}: 3.08 Å) which is centered by an oxygen of the PO₄ group (Fig. S1, left). The W–O and P–O bond lengths in **1** are in the usual range.

All four Pd^{II} ions in the complex with $\{P_2W_{15}\}$ exhibit square-planar coordination. Two of them, situated on the opposite side of the Pd_4 rectangle, coordinate an oxygen atom of one of the WO_6 unit (Pd–O 1.985(15) – 2.096(10) Å) and an oxygen atom of the P^BO_4 group (Pd–O 2.029(15) - 2.057(18) Å). With respect to the geometry of the vacant site of $\{P_2W_{15}\}$ the two Pd^{II} ions of this structural type bind the two opposite oxygen atoms of the O_6 hexagon and the central O atom which they share between each other (Pd···Pd 3.121(4) Å). The type of coordination of these two Pd^{II} ions is the same for the both *anti* and *syn* isomers of **1**.



Figure 1. Structure of the *anti*-1 (left) and *syn*-1 (right) polyanions. WO₆ octahedra: W_3O_6 "cap", gray; inner W_6O_{27} "belts", blue; outer W_6O_{27} "belts"), light blue. P^AO₄ and P^BO₄: yellow and orange tetrahedra, respectively; Pd: green spheres, O: red. The Pd₄ plane is emphasized as a transparent yellow rhombus. Rotation of the lower {P₂W₁₅} group by 60° transforms the isomers into each other.

Each of the other two Pd^{II} ions in the *anti* isomer of **1** coordinates two oxygens of the edgeshared W_2O_{10} unit of one POM ligand and two oxygens of the corner-shared W_2O_{11} unit of the second { P_2W_{15} } species (Pd–O: 1.978(15) – 2.215(18) Å). Thus, the *anti*-**1** polyanions are centrosymmetric and belong to C_{2h} point symmetry group (Fig. 1, left). In terms of the geometry of the vacant site of the every { P_2W_{15} } ligand these two Pd^{II} centers coordinate oxygens of two opposite sides of a centered $\{O_6\}$ hexagon which remain non-coordinated by the Pd^{II} ions of the first structural type (Fig. S1, right). The Pd···Pd distance is 5.576(3) – 5.621(3) Å for the palladium centers of this structural type and 3.182(9) – 3.192(19) Å between the palladium(II) ions of the first and second structural types.

In the *syn* isomer the Pd^{II} centers of the second structural type are not equivalent: one of them binds oxygen atoms of edge-shared W_2O_{10} unit of the every {P₂W₁₅} ligand and the second one, correspondingly, coordinates oxygens of the corner-shared W_2O_{11} groups of the each {P₂W₁₅}. Overall the *syn* derivative of **1** possesses idealized $C_{2\nu}$ symmetry (Fig. 1, right).

The 2 : 1 ratio between the *anti* and *syn* isomers of **1** does not change by reacting Pd^{II} and $\{P_2W_{15}\}$ at various pH (from 2.0 to 7.8) or different temperatures (including reaction and crystallization of **Na-1** at 5 °C as well as prolonged heating of the reaction mixture at 80 °C). It also remains unchanged by performing the reaction in other Na⁺-containing media (*e.g.* 0.5 M Na₂SO₄ and NaClO₄ aqueous solutions) as based on ³¹P NMR spectroscopy as well as the unit cell and, in some cases, complete crystal structure measurements. Importantly, ³¹P NMR spectra of the reaction mixtures of Pd^{II} and $\{P_2W_{15}\}$ in CH₃COONa solutions showed that this ratio is not driven by crystal packing of the isomers in **Na-1** and appears already before crystallization (Fig. S6). It is also interesting to note that complex **1** does not form by reacting Pd^{II} and $\{P_2W_{15}\}$ in water based on ³¹P NMR spectra of Pd^{II} / $\{P_2W_{15}\} / H_2O$ system.

The formation of the two isomers of $[Pd_4(\alpha-P_2W_{15}O_{56})_2]^{16^-}$ is in agreement with the prior observation of *syn* and *anti* derivatives in Pd^{II} complexes with monolacunary Wells-Dawson phosphotungstates { α_2 -P₂W₁₇}.^{5c} However in that case the significantly different shape of the *anti* and *syn* isomers of $[Pd_2(\alpha_2-P_2W_{17}O_{61})_2]^{16^-}$ resulted in the different solubility of their alkali metal salts and thus allowed isolations of the both derivatives in the pure state. Unfortunately the

relatively high $C_{3\nu}$ symmetry of {P₂W₁₅} ligands and the very similar stereochemistry of *anti*-1 and *syn*-1 render their separation hardly feasible at the moment. However we are trying to isolate other (Rb⁺, Cs⁺ or organic) salts of 1 with the aim of separating the isomers or to changing their relative ratio.

It is also interesting to compare the structure of 1 with that of well-known $[M_4(H_2O)_2(P_2W_{15}O_{56})_2]^{n-}$ (= { $M_4(P_2W_{15})_2$ }) complexes formed by octahedrally-coordinated transition metal centers ($M = Mn^{II}$, Fe^{III} , Co^{II} , Ni^{II} , Cu^{II} , Zn^{II} , and Cd^{II}).¹¹ Like in **1** the typical structure of a $\{M_4(P_2W_{15})_2\}$ complex comprises a tetrametal belt $M_4(H_2O)_2$ sandwiched between two $\{P_2W_{15}\}$ phosphotungstates. At the same time the higher coordination number six of the M^{m+} centers in $M_4(H_2O)_2$ (comparing with CN = 4 for Pd^{II} in 1) leads to significant structural differences. Thus every O atom associated with the P^BO_4 group at the lacunary site of $\{P_2W_{15}\}$ in $\{M_4(P_2W_{15})_2\}$ is coordinated not by two as in 1 but by three M^{m+} ions at the same time. Two of the four M^{m+} centers in $M_4(H_2O)_2$ coordinate to P^BO_4 groups of both $\{P_2W_{15}\}$ ligands (which could be compared with the two Pd^{II} of the first structural type in 1), whereas the remaining two M^{m+} ions only bind to a $P^{B}O_{4}$ of only one $\{P_{2}W_{15}\}$ ligand. This results in a parallel shift of the main axes of the $\{P_2W_{15}\}$ units in $\{M_4(P_2W_{15})_2\}$ while the C_3 axes of each $\{P_2W_{15}\}$ unit coincide and pass through the center of the Pd₄ rhombus (Fig. S2). From an alternative point of view, the three MO₆ octahedra in the M₄(H₂O)₂ belt coordinating to P^BO₄ form {M₃P₂W₁₅} units akin to β type Wells-Dawson structures, and the common $\{M_4(P_2W_{15})_2\}$ structure can be regarded as a centrosymmetrical $\beta\beta$ -isomer.¹¹ In the case of M = Co^{II} it was shown that a mixture of $\beta\beta$ - and $\alpha\beta$ - isomers (which differ from $\beta\beta$ by a 60° rotation of one of the {P₂W₁₅} units with respect to the $\{M_4P_2W_{15}\}$ part) co-exist in reaction mixtures at neutral pH values.¹¹ⁱ This is in direct analogy with *anti/syn* isomerism in the $\{Pd_4(P_2W_{15})_2\}$ complex.

In summary, the Pd^{II} -based complex **1**, $\{Pd_4(P_2W_{15})_2\}$, in comparison to conventional $\{M_4(P_2W_{15})_2\}$ -type structures exhibits structural similarities, in particular the similar rhombic arrangement of the four metal centers of the inner belt of the polyanions and the ability to form isomers which differ by 60° rotation of one of the $\{P_2W_{15}\}$ groups with respect to the $\{M_4P_2W_{15}\}$ unit. At the same time, **1** displays unique structural characteristics imparted by the square-planar coordination environment of Pd^{II} ions, implying the local C_3 axes of the two $\{P_2W_{15}\}$ groups coincide and formation of the *syn* isomers also in acidic media.

The structure of **1** is also unique in comparison with structurally characterized sandwich-like Pd^{II} complexes of the trilacunary Keggin-type species $[A-\alpha-(XO_4)W_9O_{30}]^{n-}$ and $[B-\alpha-(XO_3)W_9O_{30}]^{n-}$ (= {XW₉}), {Pd₃(XW₉)₂}.⁷⁻⁸ Compared to {XW₉} the vacant site of {P₂W₁₅} offers an additional central oxygen atom of the P^BO₄ group due to the different orientation of the XO₄ tetrahedron in the *B-α-* and *A-α-*trilacunary POT derivatives (Fig. S3). Also in {*B-α-*XW₉} this additional oxygen is absent and replaced by a lone pair on the central X^{III/IV} heteroion (X = As^{III}, Sb^{III}, Te^{IV}). This feature allows for coordination of four Pd^{II} centers in **1** while the sandwich-like complexes of Pd^{II} with {XW₉} POTs contain only up to three noble metal ions. On the other hand, Ratiu and co-workers in 1998 proposed formation of [Pd₄(*B-α*-PW₉O₃₄)₂]¹⁰⁻ polyanions based on UV-Vis, photocolorimetry and conductometry data.¹¹ {Pd₄(*B-α*-PW₉O₃₄)₂] should possess a structure similar to **1** where the {P₂W₁₅} ligands are replaced by [*B-α-*PW₉O₃₄]⁹⁻ POTs. However there still is no structural evidence for the formation of a complex with {Pd₄(*B-α*-PW₉O₂₄} stoichiometry and our attempts to isolate such derivative so far failed, leading to the well-characterized^{7a-c} [Pd₃(*A-α-*PW₉O₃₄]¹²⁻ species.

The IR spectrum of **Na-1** exhibits three absorption bands at 1090, 1065 and 1016 cm⁻¹ which could be assigned to vibrations of P–O bonds. They could be compared with the P–O vibrations

bands at 1130, 1086 and 1009 cm⁻¹ for $Na_{12}[\alpha-P_2W_{15}O_{56}]\cdot 24H_2O$ (Na-{P₂W₁₅}) (Fig. S4 and refs. 11a, 12) and at 1090 and 1012 cm⁻¹ for $K_6[\alpha-P_2W_{18}O_{62}]\cdot 14H_2O$ (Fig. S4 and refs. 12, 13). The disappearance of the band at 1130 cm⁻¹ characteristic for non-coordinated {P₂W₁₅} and appearance a new band at 1065 cm⁻¹ is in agreement with the coordination of one of the oxygen atoms of PO₄ group at the lacunary site of the {P₂W₁₅} ligands by Pd^{II} in **1**. The band characteristic for terminal W=O bonds appears at 941 cm⁻¹. The set of bands in the range of 909 to 767 cm⁻¹ could be assigned to vibrations of W–O–W and W–O–Pd bonds. The significant shift of these bands comparing to the W–O–W bands in non-coordinated {P₂W₁₅} (see Fig. S4) is also consistent with formation of a coordination complex between Pd^{II} and Na-{P₂W₁₅} in **Na-1**. A band at 563 cm⁻¹ which is absent in the spectrum of Na-{P₂W₁₅} corresponds to Pd–O valence bands.¹⁴ The IR spectrum of **TBA-1** exhibits similar features and additional bands corresponding to C–C, C–C and C–H vibrations of the TBA⁺ countercations (Fig. S5)

NMR spectroscopy. The room temperature ³¹P NMR spectrum of **1** (Fig. 2) exhibits three signals at -3.4, -3.6 and -14.6 ppm with the relative intensities of 2:1:3 which could be compared with the signals at +0.1 and -13.3 ppm for the non-coordinated {P₂W₁₅} ligands¹² and the lines at -4.3 and -14.3 ppm for {Zn₄(P₂W₁₅)₂}.^{11b} The most intense signal at -14.6 ppm stems from the P^AO₄ group of the {P₂W₁₅} ligands and it appears to be not sensitive to the *anti-/syn*-isomerization. The two downfield signals originate to the P^BO₄ close to the Pd₄ belt. Based on relative intensities of these two signals which are in a very good agreement with the solid-state X-ray data the signal at -3.4 ppm could be assigned to the *anti-1* isomer while the signal at -3.6 ppm belongs to the *syn-1* derivative. Spectra measured at 278 K (Fig. S7) and 353 K (Fig. S8) both exhibit the same three signals, although their chemicals shifts differ slightly compared to the room temperature spectrum (-2.9, -3.1 and -14.1 ppm at 278 K and -4.0, -4.2 and -15.2

ppm at 353 K). The relative intensity of the two downfield signals in these spectra indicates the same 2:1 ratio between the *anti* and *syn* isomers of **1** confirming the absence of dynamic phenomena associated with the complex dissociation and isomerization in aqueous medium at various temperatures.



Figure 2. Room temperature ³¹P NMR spectrum of **Na-1** redissolved in H_2O / D_2O (the three small signals belong to minor impurities which do not exceed 2%).

The spectrum remains unchanged for a long period of time, indicating high solution stability of the polyanions **1** in aqueous medium.

The room temperature ¹⁸³W NMR spectrum of aqueous Na-1 solution exhibits 15 signals (Fig. 3) in a good agreement with the expected pattern, based on the solid-state structure. The $\{P_2W_{15}\}$

ligands in both *anti* and *syn* isomers of **1** acquire local C_s symmetry with the vertical mirror plane intersecting the two Pd^{II} ions of the second structural type and one of the W^{VI} center of the W₃O₁₃ "cap". The mirror plane divides the W centers of each {P₂W₁₅} unit into seven pairs of symmetry-equivalent atoms and the unique polar W site, similar to the patterns observed for the {M₄(P₂W₁₅)₂} series (M = Co, Cu, Zn).^{11b} Accordingly, the observed 15-line spectrum can be attributed to the 2 : 1 mixture of the C_{2h} -symmetric *anti* and the $C_{2\nu}$ -symmetric *syn* isomers of **1**, each of which giving 8 signals in a 2 : 2 : 2 : 1 : 2 : 2 : 2 : 2 intensity ratio. Thus the more intense signals at -87 (relative intensity 4, J_{W-O-P} 1.4 Hz), -155 (4, J_{W-O-P} 0.8 Hz), -158 (2, J_{W-O-P} 1.5 Hz), -159.5 (4, J_{W-O-P} 1.8 Hz), -236 (4, J_{W-O-P} 1.9 Hz), -243 (4, J_{W-O-P} 1.75 Hz), -245.4 (4, J_{W-O-P} 1.65 Hz) belong to the *anti*-isomer of **1** while the weaker signals at -91 (2), -156 (2), -159 (1), -160.4 (2), -236.6 (2), -243.6 (2) and -244.6 (2) could be assigned to the *syn* derivative. The peak at -136 (6) is attributed to the overlapping signals from both *anti*-1 and *syn*-1 species.



Figure 3. Room temperature ¹⁸³W NMR spectrum of **Na-1** redissolved in H_2O / D_2O after exponential multiplication with a line broadening of 0.5 Hz (top); zoom into the signals (bottom)

Based on their lower relative intensity, the signals at -158 ppm (anti-1) and -159 ppm (syn-1) are attributed to the unique polar W^{VI} centers of the $\{P_2W_{15}\}$ ligands. The chemical shift values for these signals are in a good agreement with the literature data for $[\alpha-H_xP_2W_{15}Nb_3O_{62}]^{(9-x)-,15}$ $[\alpha - H_3 P_2 W_{15} O_{59} \{Al(OH_2)\}_3]^{6-16}$ and $[\alpha - P_2 W_{15} Ti_3 O_{62}]^{12-17}$ polyanions with $C_{3\nu}$ symmetry for which the signals corresponding to the W^{VI} centers of the W_3O_6 "caps" appear at -148.0, -156.6 and -148.3 ppm, respectively, as well as with observation of signal at -150.4 ppm for the unique W^{VI} centers in $\{Zn_4(P_2W_{15})_2\}$.^{11b} Unfortunately very close spacing of the signals for the both isomers complicates their direct assignment based on J_{W-O-W} values, however the attribution of the remaining signals can be based on the similarity to the ¹⁸³W NMR spectrum of the related ${Zn_4(P_2W_{15})_2}^{11b}$ complex (for which 2D INADEQUATE ${}^{183}W{}^{31}P$ } NMR measurements allowed unambiguous attribution of all peaks) and logical reasoning. Thus the three closely spaced upfield signals (at -236, -243 and -245.4 for anti-1 and -236.6, -243.6 and -244.6) most likely belong to the W^{VI} centers of the inner W_6O_{27} "belt" of $\{P_2W_{15}\}$ (arranged around the P^AO_4 group like the W₃O₆ "cap"). This proposition is in a good agreement with observation of the signals for the inner W_6O_{27} "belt" of $\{P_2W_{15}\}$ units in $\{Zn_4(P_2W_{15})_2\}$ at -238.4, -243.4, -244.7ppm. The signal at -159.5 ppm shows the same splitting due to J_{P-O-W} coupling (~1.8 Hz) as that of the unique polar W^{VI} center. This suggests that the signals at -159.5 ppm (for *anti*-1) and -160.4 ppm (for syn-1) also belong to the W centers of the W_3O_{13} "caps" connected with the P^B atom through the O atom that is common to all the three tungstens. Then the remaining three signals that are shifted to significantly higher frequencies relative the spectrum range of $\{Zn_4(P_2W_{15})_2\}$ relate to the W atoms connected to palladium. It is also logical to suggest that the

six W^{VI} centers close to the lacunary site of the {P₂W₁₅} units in **1** are less shielded compared to the other W^{VI} centers due to the square-planar coordination environment of the Pd^{II} ions and subsequently give the three most downfield signals (at -87, -136 and -155 ppm for *anti*-**1** and -91, -136 and -156 ppm for *syn*-**1**). At that the downfield shift of these signals in comparison with the corresponding signals in the ¹⁸³W NMR spectrum for {Zn₄(P₂W₁₅)₂}^{11b} is consistent with a weaker binding of the {P₂W₁₅} ligands to the Pd^{II} centers in **1** than to the Zn^{II} centers in {Zn₄(P₂W₁₅)₂}, which is in full agreement with the different coordination geometries of the Pd^{II} and Zn^{II} centers in these structures. We further hypothesize that the signal at -136 ppm may possibly correspond to the W^{VI} centers binding the oxygens coordinated by Pd^{II} centers of the first structural type as such W^{VI} centers have very similar coordination environment in the both *anti*-**1** and *syn*-**1** derivatives.

UV-Vis spectroscopy. The solutions of **Na-1** were further examined using absorption spectroscopy. The UV-Vis spectrum of **Na-1** in 0.5 M CH₃COONa aqueous solution (pH = 4.3), shown on the Fig. S7, exhibits strong absorption maximum at 236 nm (ε = 166167 M⁻¹cm⁻¹) followed by a broad shoulder at about 292 nm (ε = 81495 M⁻¹cm⁻¹) in the UV light area and a less intense absorption maximum at 477 nm (ε = 2374 M⁻¹cm⁻¹) in the visible light area. The spectrum remains unchanged for at least 1 week for both more (6.7×10⁻⁴ M) and less (4.7×10⁻⁵ M) concentrated solutions confirming the stability of polyanions **1** in 0.5 M CH₃COONa aqueous medium at pH 4.3.

The spectra of **Na-1** solutions in 1 M CH₃COOH (pH 2.0) and 1 M CH₃COONa at various pH are shown on the Fig. S10. The solutions were prepared independently taking equal amounts of **Na-1** and equal amounts of the corresponding solvent. The obtained spectra showed highest absorption at about 477 nm in the solutions with a pH range of 2 to 4 suggesting the highest

stability of **1** in these media. The spectra at pH 2.0 and 4.0 remained unchanged for at least 1 day while it was not possible to monitor the stability of **1** in the solutions with higher pH values due to lower solubility and recrystallization of **Na-1** which occurs within several hours.

CONCLUSIONS

In our investigation of the reactivity of Pd^{II} ions towards trilacunary derivative of Wells-Dawson-type phosphotungstates $[\alpha-P_2W_{15}O_{56}]^{12-}$ in 0.5 M CH₃COONa media, the tetranuclear sandwich-like complex $[Pd_4(\alpha-P_2W_{15}O_{56})_2]^{16-}$ was found to self-assemble in a wide range of pH, Pd^{II} : $\{P_2W_{15}\}$ ratios and reaction temperatures. The $[Pd_4(\alpha-P_2W_{15}O_{56})_2]^{16-}$ clusters exist as *syn* and *anti* isomers which vary in the relative orientation of the two $\{P_2W_{15}\}$ ligands and form in a 2 : 1 ratio. Due to square-planar Pd^{II} coordination the title structure differs from conventional (transition metal-substituted) $\{M_4(P_2W_{15})_2\}$ species despite the similar rhombic arrangement of the four heterometal ions in these complexes. The arrangement of Pd^{II} centers in 1 also differs markedly from Pd^{II} complexes with trilacunary Keggin-type polyoxotungstates because of the slightly different structure of the vacant sites in these POTs. ³¹P and ¹⁸³W NMR spectroscopy demonstrated a long-term solution stability of polyanions 1 in aqueous media.

EXPERIMENTAL SECTION

General methods and materials. The reagents were used as purchased without further purification. Na₁₂[α -P₂W₁₅O₅₆]·24H₂O was obtained according to the reported procedure^{11a,12} starting from K₆[α -P₂W₁₈O₆₂]·14H₂O.^{12,13} Elemental analysis results (ICP-OES) were obtained from Central Institute for Engineering, Electronics and Analytics (ZEA-3), Forschungszentrum Jülich GmbH (D-52425 Jülich, Germany). Vibrational spectra were recorded on a Bruker VERTEX 70 FT-IR spectrometer coupled with a RAM II FT-Raman module (1064 nm Nd:YAG laser) on KBr disks for the FT-IR and the solid material for the Raman measurements. UV-Vis spectra were measured using 10 mm quartz cuvettes on Analytik Jena Specord S600 spectrophotometer. ³¹P NMR spectra were recorded at room temperature in 5 mm tubes using a Bruker Avance 600-MHz spectrometer equipped with a prodigy probe, operating at 242.95 MHz for ³¹P and with a Varian Inova 400 MHz spectrometer equipped with an Auto-X-PFG-probe and with resonance frequency of 161.834 MHz. Chemical shifts are reported with respect to 85% H₃PO₄; all chemical shifts downfield of the reference are reported as positive values. ¹⁸³W NMR spectra of **Na-1** solution in H₂O / D₂O ($c \sim 1 \times 10^{-2}$ M) were recorded in 10 mm tubes on a Bruker Avance 400 MHz instrument at room temperature with a resonance frequency of 16.67 MHz. The chemical shifts are reported with respect to 1 M Na₂WO₄ aqueous solution as a reference.

Synthesis of Na₁₆[Pd₄(\alpha-P₂W₁₅O₅₆)₂]·71H₂O (Na-1). A sample of Na₁₂[\alpha-P₂W₁₅O₅₆]·24H₂O (0.450 g, 0.102 mmol) was dissolved in 9 mL of 0.5 M CH₃COONa buffer (pH 4.2) under vigorous stirring. Solid Pd(NO₃)₂·2H₂O (0.075 g, 0.281 mmol) was added to the obtained solution and the reaction mixture was stirred for another 60 min at room temperature and then filtered and left for evaporation at room temperature being divided into 2–3 vials. Brown crystals of Na-1 (rhombic plates) form within 1–2 weeks. The crystals were collected by filtration, washed with ice-cold water and dried in air. Yield: 0.27 g (55% based on {P₂W₁₅}). Elemental analysis: calculated for H₁₄₂Na₁₆O₁₈₃P₄Pd₄W₃₀ (found): Na, 3.87 (3.86); P, 1.30 (1.32); Pd, 4.48 (4.48); W, 58.03 (58.0)%. IR spectrum (KBr pellet), cm⁻¹: 3434 (s, br); 1621 (s); 1090 (s); 1065 (m); 1016 (m); 941 (s); 909 (s); 831 (s); 767 (s); 598 (m); 563 (m); 527 (m); 394 (w); 376 (w). Raman (in solid), cm⁻¹: 984 (s); 964 (s); 887 (m); 822 (w); 526 (w); 374 (w); 324 (w); 226 (w); 164 (w); 118 (w). ³¹P NMR (H₂O/D₂O): *anti***-1, \delta: –3.4, –14.6 ppm;** *syn***-1, \delta: –3.6, –14.6 ppm. ¹⁸³W NMR (H₂O/D₂O):** *anti***-1, \delta: –87 (2 W), –136 (2 W), –158 (1 W), –159.5 (2 W), –156 (2 W), –245.4 (2 W) ppm;** *syn***-1, \delta: –91 (2 W), –136 (2 W), –156 (2 W), –156 (2 W), –**

159 (1 W), -160.4 (2 W), -236.6 (2 W), -243.6 (2 W), and -244.6 (2 W) ppm. UV-Vis (0.5 M CH₃COONa buffer solution, pH 4.3): λ = 236 nm, ε = 166167 M⁻¹cm⁻¹; λ = 292 nm, ε = 81495 M⁻¹cm⁻¹; λ = 477 nm, ε = 2374 M⁻¹cm⁻¹.

Synthesis of $[(C_4H_9)_4N]_{15}[HPd_4(\alpha-P_2W_{15}O_{56})_2]$ (TBA-1). A solution of Na-1 (0.100 g, 0.010 mmol) in 3 mL of H₂O was dropwise added to an aqueous solution of $[(C_4H_9)_4N]HSO_4$ (0.080 g, 0.236 mmol, 2 mL of H₂O) under vigorous stirring. The mixture was acidified with 1 drop of 2M HNO₃. The obtained precipitate of **TBA-1** was filtered on a glass frit, washed with plenty of water, and dried in air. Elemental analysis: calculated for C₂₄₀H₅₄₁N₁₅O₁₁₂P₄Pd₄W₃₀ (found): C, 25.08 (24.04); H, 4.74 (4.77); N, 1.83 (1.90)%. IR spectrum (KBr pellet), cm⁻¹: 3468 (m, br); 2961 (s); 2934 (m); 2873 (m); 1630 (w); 1484 (m); 1383 (m); 1237 (w); 1167 (m); 1094 (s); 1064 (m); 1025 (w); 999 (w); 958 (s); 902 (s); 779 (s, br); 597 (m); 595 (m); 560 (m); 530 (m); 430 (w); 392 (m).

X-ray Crystallography. Single-crystal diffraction data for **Na-1** were collected on a SuperNova (Agilent Technologies) diffractometer with MoK α radiation ($\lambda = 0.71073$ Å) at 120 K. A crystal was mounted in a Hampton cryoloop with Paratone-N oil to prevent water loss. Absorption corrections were applied numerically based on multifaceted crystal model using CrysAlis software.¹⁸ The SHELXTL software package ¹⁹ was used to solve and refine the structure. The structure was solved by direct methods and refined by full-matrix least-squares method against $|\mathbf{F}|^2$ with anisotropic thermal parameters for all heavy POM skeleton atoms (Pd, P, W) and sodium countercations. The hydrogen atoms of the crystal waters were not located. The relative site occupancy factors for the disordered positions of tungsten, palladium and oxygen atoms due to the {Pd₄P₂W₁₅} / {P₂W₁₅} ligands rotation by 60° were refined using the PART instruction combined with EADP restrictions for the heavy atoms and then fixed at the

obtained values and refined normally. The relative site occupancy factors for the disordered solvent oxygens were first refined in an isotropic approximation with U_{iso} = 0.05 and then fixed at the obtained values and refined without the thermal parameters restrictions.

The number of crystal water molecules and sodium countercations found by XRD was smaller than that determined by elemental analysis (32 vs 71 for H₂O and 7 vs 16 for Na⁺, respectively) which could be explained by the high degree of disorder in the solid-state structure of Na-1. This is also consistent with large solvent-accessible volume remained in the structure. For the overall consistency the formula shown in the CIF file corresponds to the bulk material and has the same number of countercations and crystallization water molecules as found by elemental analysis since all the further studies are / will be performed on the isolated bulk material of Na-1.

Additional crystallographic data are summarized in Table 1. Further details on the crystal structure investigation may be obtained from Fachinformationszentrum Karlsruhe, 76344 Eggenstein-Leopoldshafen, Germany [fax (+49) 7247-808-666; e-mail crysdata@fiz-karlsruhe.de), upon quoting the depository number CSD 428389.

Empirical formula	$H_{142}Na_{16}O_{183}P_4Pd_4W_{30}$
Formula weight, g/mol	9503.96
Crystal system	Triclinic
Space group	<i>P</i> -1
<i>a</i> , Å	14.0333(3)
b, Å	14.0985(3)
<i>c</i> , Å	25.2703(5)
α	83.9425(17)°

 Table 1. Crystal data and structure refinement for Na-1

β	76.5160(19)°
γ	60.514(2)°
Volume, Å ³	4231.88(16)
Ζ	1
$D_{\text{calc}}, \text{g/cm}^3$	3.729
Absorption coefficient, mm ⁻¹	20.912
<i>F</i> (000)	4246
Crystal size, mm	$0.06 \times 0.06 \times 0.13$
Theta range for data collection	4.09° - 25.03°
Completeness to Θ_{\max}	98.5%
Index ranges	$-16 \le h \le 16,$
	$-16 \le k \le 16,$
	$-30 \le l \le 30$
Reflections collected	75309
Independent reflections	14717
R _{int}	0.0722
Observed $(I > 2\sigma(I))$	13246
Absorption correction	Empirical using spherical harmonics
T_{\min} / T_{\max}	0.0822 / 0.3732
Data / restraints / parameters	14717 / 24 / 663
Goodness-of-fit on F ²	1.109
$R_1, \mathrm{w}R_2 (I > 2\sigma(I))$	$R_1 = 0.0722,$
	$wR_2 = 0.1829$
R_1 , w R_2 (all data)	$R_1 = 0.0781,$
	$wR_2 = 0.1874$
Largest diff. peak and hole, e. $Å^{-3}$	3.876 and -3.061

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ASSOCIATED CONTENT

Supporting Information. Crystallographic data in CIF format, IR spectra and UV-Vis spectra; structure of $\{P_2W_{15}\}$ and its vacant site and comparison of the $\{Pd_4(P_2W_{15})_2\}$ structure with the structures of $\{M_4(P_2W_{15})_2\}$ and $\{Pd_3(XW_9)_2\}$ polyanions. This material is available free of charge via the Internet at http://pubs.acs.org.

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Author Contributions

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

ABBREVIATIONS

POT: polyoxotungstate; POM: polyoxometalate.

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 Pd^{II} -based sandwich complexes of lacunary $\{P_2W_{15}\}$ -type polyoxotungstates differ markedly from classical dimeric $\{M_4(P_2W_{15})_2\}$ structures incorporating octahedral first-row transition metal M^{II} centers, form *syn* and *anti* isomers in a characteristic 2:1 ratio and exhibit high stability in solution.