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An Ir^{IV}-containing polyoxometalate†

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 $[HIr^{IV}W_6O_{24}]^{7-}, \ representing \ the \ first \ Ir-containing \ Anderson-Evans-type \ polyanion \ and \ the \ first \ structurally \ characterized \ Ir^{IV}-based \ polyoxometalate, \ has \ been \ isolated \ as \ its \ hydrated \ sodium \ salt \ and \ characterized \ by \ single-crystal \ X-ray \ diffraction, \ mass \ spectrometry, \ and \ IR, \ UV-Vis \ and \ EPR \ spectroscopy. \ Cyclic \ voltammetry \ indicates \ that \ the \ Ir^{IV} \ ions \ in \ [HIrW_6O_{24}]^{7-} \ can \ undergo \ reversible \ one-electron \ reduction \ and \ oxidation, \ resulting \ in \ Ir^{II} \ and \ Ir^V \ derivatives.$

The chemistry of polyoxometalates (POMs) incorporating noble metals represents a rapidly developing area, the interest in which is driven by various fundamental and applied aspects.¹ Members of this class demonstrate a range of promising properties, in particular, catalytic activity in water splitting^{2,3} and oxidation of various organic substrates.⁴ We stress, though, that the number of publications on POM complexes of Ru and Pd by far exceeds those dedicated to all remaining noble metals (Pt, Au, Ir, Os and Rh) taken together. In particular, the chemistry of Ir-containing POMs for a long time has been limited to various POM-supported IrI- and IrIII-organometallic complexes (e.g. $[Ir^{I}(1,5-COD)]^+$, $[Ir^{III}(Cp^*)]^{2+}$ etc.),⁵ some of which, *e.g.* $[(1,5-COD)Ir^{I}Nb_{3}P_{2}W_{15}O_{62}]^{8-}$, are pre-catalysts for cycloalkene oxygenation with O2.5b-d Recently a number of Keggin and Wells-Dawson type POMs with covalently grafted heteroleptic cyclometalated Ir^{III}-complexes exhibiting effective charge separation

between the Ir^{III}-based chromophore and the POM unit were reported.⁶ The first reliably characterized iridium-containing POM incorporating no organometallic fragments, $[(Ir^{III}Cl_4)(P_2W_{20}O_{72})]^{15-}$, was reported as late as in 2009 by Hill's group and identified as a water oxidation catalyst.³ Two years later, the synthesis, characterization and reactivity of Keggin-type complexes $[PW_{11}O_{39}Ir^{III}L]^{n-}$ (n = 4 or 5; $L = H_2O$, OH, CH₃, Cl, SCN) were reported.⁷ Yet, POMs incorporating Ir^{IV} centers, which are of interest as promoters or intermediates in oxidation reactions, remain unknown. For this oxidation state, we expect Ir-containing polyoxotungstates or molybdates to adopt the Anderson–Evans structure, in analogy to known Rh^{III},⁸ Pt^{IV},^{9,10} and even Pd^{IV} derivatives.¹¹

We report the preparation of $[HIr^{IV}W_6{}^{VI}O_{24}]^{7-}$ (**IrW**), the first polyoxometalate incorporating Ir^{IV} , and its characterization by single-crystal X-ray analysis, mass spectrometry, IR, UV-Vis and EPR spectroscopy, as well as cyclic voltammetry.

IrW assembles in condensation reactions of in situ-formed Ir^{IV} hydroxo complexes with tungstate ions in aqueous acetate media. In a first step, IrCl₃·3H₂O was dissolved in a CH₃COONa solution and the pH of the reaction mixture was adjusted to 12 with 6 M NaOH. Heating of the solution led to a gradual color change from brown-green to blue and then to purple. Previous studies on alkaline solutions of hexachloroiridate(III) have shown that the processes in such reaction systems include substitution of Cl⁻ ligands in the inner coordination sphere of the IrIII ions by OH⁻ groups and the subsequent oxidation of thus formed hydroxo-oxo complexes of Ir^{III} in air.¹² During the reaction, the color change of the Ir^{IV}-containing solutions from blue to purple is apparently caused by (not well-defined) oligomerization processes. Another hypothesis postulates transformation of the blue superoxo dimeric [(OH)₅Ir^{IV}(µ-O₂⁻)Ir^{IV}(OH)₅]³⁻ species into the purple $[(OH)_4 Ir^{IV}(\mu - O_2^{2-})(\mu - OH) Ir^{IV}(OH)_5]^{3-}$ complex, based on UV-Vis and EPR studies of 3 M KOH solutions of [IrCl₆]³⁻.^{12f} Next, the solution of the *in situ*-formed Ir^{IV} hydroxo complexes was added dropwise to a solution containing the lacunary [B-α-PW₉O₃₄]⁹⁻ polyanion under careful control of pH kept strictly between 6 and 7 by addition of diluted nitric acid.

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 $[B-\alpha-PW_9O_{34}]^{9-}$ is unstable in solution and acts as a source of free WO₄²⁻ species, which effectively assemble and condense around Ir^{IV}, resulting in IrW. This is conceptually similar to the controlled formation of Mn^{III} species by slow hydrolysis of [Mn₁₂(CH₃COO)₁₆(H₂O)₄O₁₂] in the synthesis of Mn-substituted polyoxotungstates.¹³ It should be noted that both monoprotonated and non-protonated forms of $[Ir^{IV}W^{VI}O_{24}]^{8-}$ (IrW') can be isolated as the hydrated sodium salts Na₆H[HIr^{IV}W^{VI}O₂₄]·26H₂O (Na-IrW) and Na₈[Ir^{IV}W^{VI}O₂₄]·26H₂O (Na₈-IrW'), respectively. The exact protonation state depends on the final pH of the reaction mixture.

Thus, pure Na-IrW crystals form when the final pH of the reaction mixture before crystallization is set within the range of 6.1-6.5. At higher final pH (7.0-7.5) crystals of Na₈-IrW' form as a byproduct to Na-IrW. No crystals could be isolated at pH > 7.6 where one could expect formation of pure Na_8 -IrW'. We herein focus on the monoprotonated species and provide structural data for Na₈-IrW' in the ESI.†

Our attempts to rationalize the synthetic procedure by reacting Ir^{IV} solutions directly with Na₂WO₄ at various ratios were not successful and resulted in formation of the paratungstate salt Na₁₀[H₂W₁₂O₄₂]·25H₂O as the main product, as established via single-crystal X-ray measurements and IR spectroscopy. We conjecture that an initial excess of Ir^{IV} ions coupled with the gradual release of WO_4^{2-} species into the reaction medium are key to successful preparation of IrW. Alternatively, IrW can be prepared by prolonged heating of $K_2[IrF_6]$ with Na₂WO₄ in water at neutral pH, however, this reaction is not very well reproducible and the **IrW** salts could be isolated only in a low yield ($\leq 10\%$).

The compound Na₆H[HIr^{IV}W₆^{VI}O₂₄]·26H₂O (Na-IrW) crystallizes in the triclinic space group *P*1. The IrW polyanions display an Anderson-Evans-type ring structure comprising a central Ir^{IV}O₆ octahedron surrounded by six distorted edge-sharing $W^{VI}O_6$ octahedra (Fig. 1). All metal centers in the IrW₆ core define a single plane and form a virtually regular Ir-centered W₆ hexagon (W···W: 3.223-3.345 Å; Ir···W: 3.234-3.316 Å). IrW has idealized D_{3d} symmetry and thus belongs to the A-type family of polyanions with Anderson-Evans structure which is typical for $[M'W_6O_{24}]^{n-}$ heteropolyoxotungstates. In contrast, some of the polyoxomolybdates $[M'Mo_6O_{24}]^{n-}$ may also adopt a bent conformation with C_{2v} symmetry (usually referred as a B-type Anderson-Evans structure), which was e.g. found in





Fig. 1 Structure of IrW. Color code: WO₆, blue octahedra; Ir, green; O, red. Fig. 2 Powder X-band EPR spectrum of Na-IrW at 110 K.

$[H_4PtMo_6O_{24}]^{4-}, {}^{9a,c-e} [H_2SbMo_6O_{24}]^{5-}$ (ref. 14) and heptamolybdate [Mo₇O₂₄]⁶⁻ ions.¹⁵

There are three types of oxygen atoms in IrW: μ_3 -O linking Ir^{IV} and two W^{VI} centers (Ir-O: 1.994(5)-2.022(5) Å; W-O: 2.142(5)–2.293(5) Å), μ_2 -O bridging two W^{VI} ions (W-O: 1.923(5)-1.987(5) Å) and terminal O atoms (W=O: 1.735(5)-1.747(5) Å). The average Ir-O bond length in IrW of 2.004 Å is comparable with that of 2.013 Å observed in the only structurally characterized Ir^{III}-containing all-inorganic polyoxotungstate $[(Ir^{III}Cl_4)P_2W_{20}O_{72}]^{15-,3}$ and with that in Ir^{IV} oxides, *e.g.* CaIrO₃ (2.057 Å).¹⁶ The W–O distances in **IrW** are similar to the W–O bond lengths in other $[XW_6O_{24}]^{n-}$ species (X = Pt^{IV}, Te^{IV}, Mn^{IV} , n = 8; $X = Sb^{V}$, n = 7).^{10,17} Bond valence sum calculations (Table S2, ESI[†]) suggest a disorder of a proton over two inversion center related μ_3 -O atoms (O123) in IrW. The second proton, which should be present in the structure of Na-IrW from the charge balance considerations (only six Na⁺ countercations were found both in X-ray diffraction and elemental analyses of Na-IrW), is either highly disordered over the terminal oxygens of the IrW polyanions (BVS 1.60-1.67) or, more likely, is associated with an oxygen of a co-crystallized water molecule, *i.e.* as a H₃O⁺ cation. In the crystal lattice of Na-IrW the Na⁺ countercations and the crystal water molecules assemble in the parallel infinite zig-zag {Na₆(μ_2 -H₂O)₁₀(H₂O)^t₁₄}⁶ⁿ⁺ chains. The IrW polyanions are packed between the chains (see ESI[†] for details) and form an extended net of hydrogen bonds with the H atoms of the H₂O molecules (Fig. S1 and Table S4, ESI[†]).

The oxidation state of the iridium center (Ir^{IV} , $5d^5$, S = 1/2) was further confirmed by the characteristic EPR signal around g = 2 (Fig. 2). The complexity of the powder spectrum indicates the strong contribution of quadrupole as well as hyperfine coupling $(I = 3/2 \text{ for } {}^{191}\text{Ir} \text{ [natural abundance } 37\%] \text{ and } {}^{193}\text{Ir}$ [63%]), similar to what has been observed for iridium(IV) impurities displaying a tetragonally distorted octahedral environment in MgO and CaO matrices.¹⁸ Unfortunately, low distortion of the iridium coordination sphere in IrW leads to a small split between g values, and the resulting overlap of the various components in our powder X-band spectrum precludes direct determination of the hyperfine and quadrupole coupling constants, while the large number of variables renders an exhaustive search of the parameter space impracticable.



Fig. 3 CV of 1.3 mM **Na–IrW** solution in 2 M NaOAc medium (pH 6.1) at different scan rates: 10 mV s⁻¹ (green), 25 mV s⁻¹ (pink), 50 mV s⁻¹ (blue), 100 mV s⁻¹ (red), 250 mV s⁻¹ (black) vs. Ag/AgCl.

The ESI mass spectrum obtained by ionization of Na-IrW solution in H₂O/MeOH is very complex and exhibits signals which could be assigned for the doubly negatively charged ion pairs $\{Na_4H_2[Ir^{IV}W_6^{VI}O_{24}]\}^{2-}$ and $\{Na_3H_3[Ir^{IV}W_6^{VI}O_{24}]\}^{2-}$ as well as sets of signals corresponding to various decomposition products, e.g. $[HWO_4]^-$, $[HW_2O_7]^-$, $Na_xH_{5-x}[IrW_5O_{20}]^{2-}$ etc. (see ESI[†]). While it confirms the existence of the intact polyanions in aqueous solutions the obtained data do not answer the question if the partial decomposition starts to occur already in the solution or only happens during the ionization process. The stability of the polyanions in aqueous solution as well as aqueous acetate media with different pH has been further checked by UV-vis spectroscopy (see ESI† for details). Electrochemistry studies of IrW in aqueous CH₃COONa solution at pH 6.1 (see ESI[†]) produce typical voltammograms (Fig. 3) that show two quasi-reversible redox couples with the corresponding peak potentials $E_{1/2}$ located at 0.245 V ($\Delta E = 0.190$ V at 100 mV s⁻¹) and 0.700 V ($\Delta E = 0.200$ V) vs. Ag/AgCl reference electrode (0.441 V and 0.896 V vs. SHE, respectively). These waves are attributed to the Ir^{III}/Ir^{IV} and Ir^{IV}/Ir^V redox transformations within the polyanion $[HIrW_6O_{24}]^{n-}$. The observed Ir^{III}/Ir^{IV} potential value is similar to 0.18 V found for the Ir^{III}/Ir^{IV} transition in $[PW_{11}O_{39}Ir^{III}(H_2O)]^{4-}$, and the Ir^{IV}/Ir^{V} couple is close to the potential of irreversible oxidation of the Ir centers in the same POM 0.75 V (vs. Ag/AgCl).^{7a} The significant shift to lower potentials as compared with the Ir^{III}/Ir^{IV} couple observed in the hexachloroiridates (0.669 V vs. Ag/AgCl¹⁹) is consistent with the stabilization of higher oxidation states of transition metals generally observed in the oxygen environment provided by polyoxometalates. The relative broadness of the waves could be due to uncompensated solution resistance but this can be ruled out since no change was observed upon increase of the electrolyte concentration. A deviation from reversibility is to be also excluded by the fact that the peak currents for the both processes remain proportional to the square root of the scan rate (Fig. S11, ESI⁺), indicating diffusion-controlled electrode reactions for scan rates up to 1 V s^{-1} .²⁰ The diffusion coefficient $D = 2.62 \times 10^{-6} \text{ cm}^2 \text{ s}^{-1}$ calculated for $[\text{HIr}^{\text{IV}}\text{W}_6\text{O}_{24}]^{7-}$ according to Randles–Sevcik equation²¹ compares well to the values reported for other polyoxotungstates.²² The broadness of the redox waves reflects the co-existence of multiple protonation states at pH = 6.1. This is supported by significant narrowing of the waves when the pH decreases to 4.3 (Fig. S12, ESI[†]), as well as by isolation of the non-protonated form, IrW', at pH > 6.5 (*vide supra*). Detailed studies of the protonation phenomena are underway. The redox couples associated with reduction of tungsten(vi) centers in IrW are not accessible, which is typical for W^{VI} centers in POMs possessing two terminal *cis*-oxo ligands.²³

In summary, the identified synthetic path leading to new Ir^{IV}-based polyoxotungstate with Anderson–Evans structure, $[HIr^{IV}W_6O_{24}]^{7-}$, comprises the use of *in situ*-formed Ir^{IV} oxo-hydroxo complexes and tungstate ions gradually released in solution. The **IrW** polyanion is the first example of a structurally characterized Ir^{IV}-containing POM as well as the first Ir-based POM with Anderson–Evans structure. The Ir^{IV} oxidation state in **IrW** has been directly confirmed by EPR spectroscopy and the cyclic voltammetry measurements showed the possibility of reversible oxidation and reduction of the Ir^{IV} ions in **IrW** resulting in Ir^V and Ir^{III} species. Our results suggest existence of a rich Ir^{IV}-POM and possibly even Ir^V-POM chemistry that we are planning to elucidate.

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