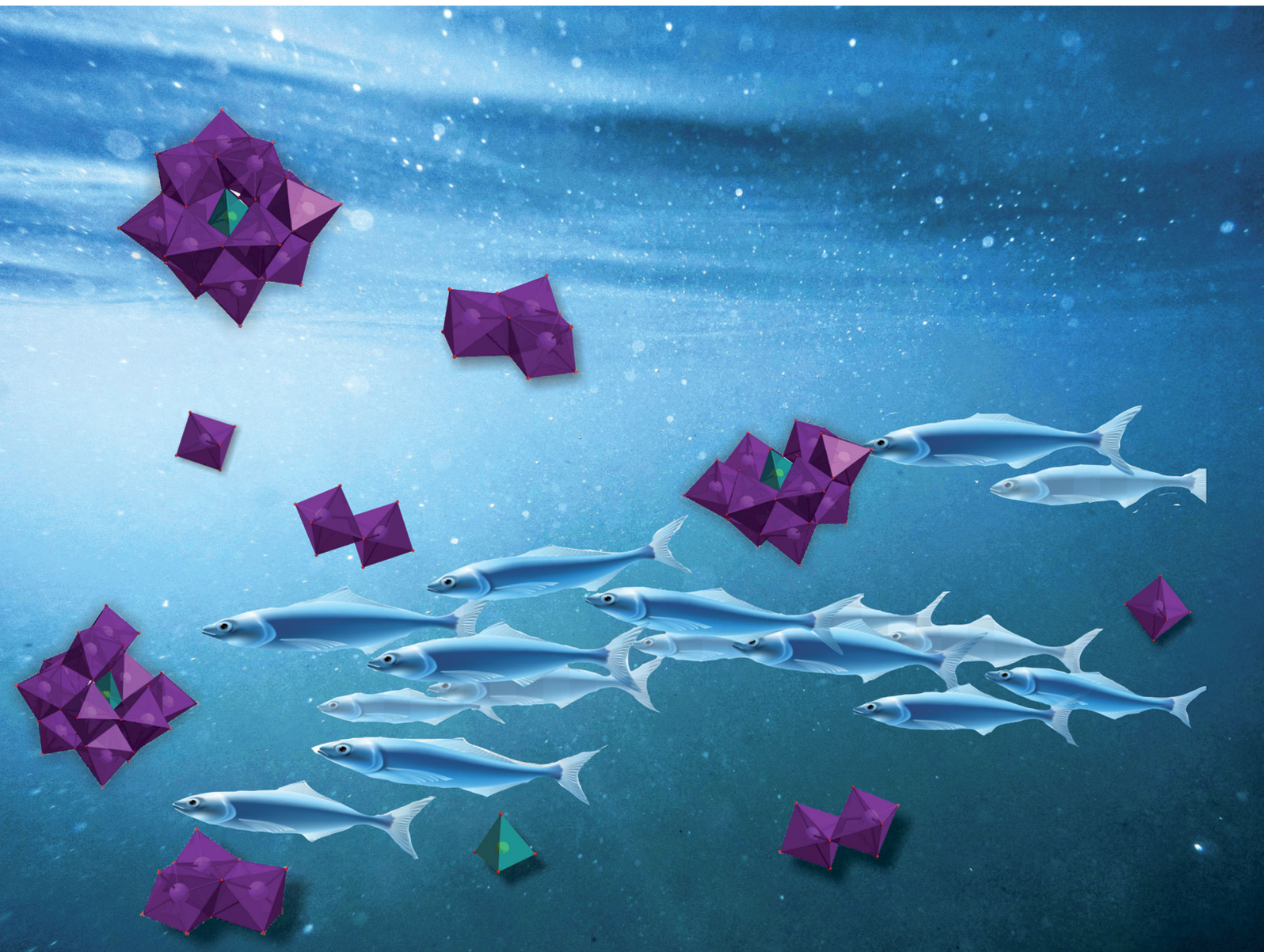


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Polyoxometalates in solution: speciation under spotlight



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Polyoxometalates in solution: speciation under spotlight†

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Polyoxometalates (POMs) are a large group of anionic polynuclear metal–oxo clusters with discrete and chemically modifiable structures. In most aqueous POM solutions, numerous, and often highly negatively charged, species of different nuclearities are formed. It is rather difficult to determine the dominant POM species or their combination, which is responsible for the specific POM activity, during a particular application. Thus, the identification of all individual speciation profiles is essential for the successful implementation of POMs in solution applications. This review article summarizes species that are present in isopoly- and heteropolyvanadates, -niobates, -molybdates and -tungstates aqueous solutions and covers their stability and transformations. The ion-distribution diagrams over a wide pH range are presented in a comprehensive manner. These diagrams are intended for the targeted use of POMs, and in a clear form shows species that are in equilibrium at the given pH value. Thus, the data accumulated in this review can serve as both a starting point and a complete reference material for determining the composition of POM solutions. Some examples are highlighted where the POM speciation studies led to a detailed understanding of their role in applications. In doing so, we aim to motivate the POM community for more speciation studies and to make the subject more comprehensible, both for synthetic POM chemists and for scientists with different backgrounds interested in applying POMs in biological, medical, electrochemical, supramolecular and nanochemistry fields, or as homogeneous catalysts and other water-soluble materials.

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Annette Rompel studied Chemistry at the Westfälische Wilhelms University of Münster where she received her doctoral degree. Besides research at the University of California, Berkeley, and the Lawrence Berkeley National Laboratory, she was a visiting scientist at the RIKEN, Institute of Physical and Chemical Research, Sendai, Japan, and the University of Southern Denmark, Odense. Since 2008, she is the Head of

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1. Introduction

1.1. Solution and solid-state structural chemistry of polyoxometalates (POMs)

Polyoxometalates (POMs) are a large group of discrete, mostly anionic polynuclear metal-oxo clusters amenable to a variety of chemical transformations.^{1–3} POMs are generally characterized in the solid state prior to dissolution, and this structural information is used as the framework on which the solution chemistry is developed. The compound isolated in crystalline form may not necessarily be the one with highest abundance in solution. For example, di- and tri-molybdates can be easily crystallized from an acidified molybdate solution at pH ~ 7,^{4,5} but they are not represented as discrete solution species, and the dominating anion at this pH is [Mo₇O₂₄]⁶⁻.⁶ In solution, POMs form species that can be protonated and undergo redox processes, which contribute to the utmost importance of the speciation characterization. For the application and/or investigation of POM complexes in aqueous solution, a thorough insight of the solution chemistry by identifying all equilibrium constants and individual speciation profiles is essential in order to understand the reaction mechanism and tune the application conditions accordingly. In rare cases where the solution chemistry is presented in literature, the solution details are often not of the main interest for the authors, and are described only in the supplementary materials. The rapidly growing number of POMs application in solution, especially their catalytic⁷ and biological ones,^{8–11} entails the need for a deeper understanding and analysis of the fundamental relationship between POM's structural behavior in the solid state and in solution. This is currently an acute drawback, often leading to an incorrect interpretation and erroneous determination of structure–activity relationships. To greatly facilitate the selection of a suitable POM cluster with classical addenda atoms (V^V, Nb^V, Ta^V, Mo^{VI} and W^{VI}) for any deliberate and purposeful use in solution, this review will serve as a guide for better understanding of the POM behavior in the liquid phase. The POM solution equilibria presented in details can and should be used to interpret the results obtained in POM solution application for fundamental vision and full understanding of the POM nature.

1.2. Speciation in chemistry: key factors affecting POMs solution behavior

The most concise definition of chemical speciation is as follows: composition, concentration, and oxidation state of each of the chemical forms of an element present in a sample.^{12,13} The term “speciation” is also used to describe the distribution of species in a particular sample, where it is synonymous with the “species distribution”.¹⁴ This notion has been accepted in such diverse fields as toxicology, clinical chemistry, geochemistry, environmental chemistry, biochemistry and inorganic chemistry. New developments in analytical instrumentation and methodology (electronical, vibrational, X-ray absorption and nuclear magnetic resonance spectroscopy,

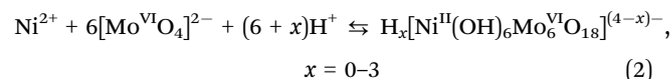
mass-spectrometry, electrochemistry) allow identifying and quantifying the species present in solution.

The key factors affecting POMs speciation and the mechanism of isopoly- and heteropolyanions formation are the added acid and metal concentrations, kind of interactions and the range of chemical conditions (ionic strength, buffer type, presence of potential heteroatoms, type of counterions, *etc.*) under which the dissolution takes place. It is not the pH alone that determines POMs speciation; rather, it is the ratio of acid (usually strong inorganic acids such as HCl, HNO₃, H₂SO₄) to monomeric oxo-ions (*e.g.* [Mo^{VI}O₄]²⁻, [W^{VI}O₄]²⁻ or [V^{VO}O₄]³⁻) concentration. This ratio determines the ‘degree of protonation’, *Z*, which is defined as the average number of protons bound to monomeric oxo-metalate in solution.¹ It is widely used in speciation studies, and we will refer to it in the cases where no pH regions are given and *Z* is used by the authors. *Z* is defined as the ratio *q/p* in the general eqn (1):



in case of M = V^V, *l* = 1, *n* = 4, *k* = 3; and M = Mo^{VI} or W^{VI}, *l* = 1, *n* = 4, *k* = 2.

The main results of speciation studies are distribution curves that represent the percentages, partial mole fractions (*α*) or equilibrium concentrations of the different chemical species present in solution under given conditions.¹⁵ Concentration distribution curves are generally presented as a function of a single variable, such as pH. For distribution curves of POMs, the ‘degree of protonation’ *Z* is commonly used as one single variable. The equilibrium concentrations of various species are calculated by solving the mathematical system of mass balance equations constructed for each component, and these mass balance equations are then solved iteratively for the concentrations of the free components.¹⁶ It should be noted that the thermodynamic equilibrium constants are based on activities that depend on temperature and pressure. Most reported stability constants for POMs are considered as stoichiometric constants, which are expressed as equilibrium concentration quotients, and thus they are valid only at a given ionic strength (*μ*, M) and in a given solvent. As an example, the concentration equilibrium constants lg *K_c* for the formation of Ni-centered Anderson-type polyoxomolybdate (POMo) eqn (2):



is calculated according to eqn (3):

$$K_c = \frac{[H_x[Ni^{II}(OH)_6Mo_6^{VI}O_{18}]^{(4-x)-}]}{[Ni^{2+}] \cdot [Mo^{VI}O_4^{2-}]^6 \cdot [H^+]^{(6+x)}}. \quad (3)$$

Potentiometric studies (Fig. 1A) allowed to calculate lg *K_c* for each anion in solution and based on these values the ion distribution diagram depending on the acidity [*α*, mole fraction = *f*(*Z*)] was built (Fig. 1B).¹⁷

To avoid questioning of inconsistent results presented by various authors, throughout this review we deliberately display distribution diagrams without considering concentration or



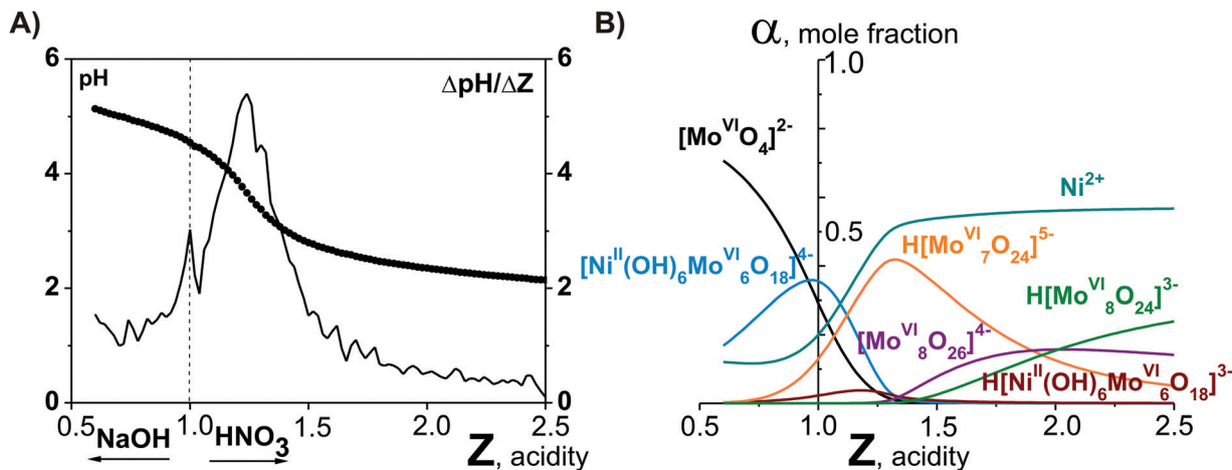


Fig. 1 pH-Potentiometry and mathematical modeling in the system $\text{Ni}^{2+} - [\text{Mo}^{\text{VI}}\text{O}_4]^{2-} - \text{H}^+ - \text{H}_2\text{O}$ acidified to mole ratio $Z = n(\text{H}^+)/n([\text{Mo}^{\text{VI}}\text{O}_4]^{2-}) = 1.00$.¹⁷ (A) Integral (points, $\text{pH} = f(Z)$) and differential (line, $\Delta\text{pH}/\Delta Z = f'(Z)$) titration curve. (B) Distribution diagrams of ions presented in solution after 60 days aging of the initial $\text{Ni}^{2+} - [\text{Mo}^{\text{VI}}\text{O}_4]^{2-} - \text{H}^+ - \text{H}_2\text{O}$ solution.

mole fraction and only focus on the pH range in which a particular POM species is stable. As an example, the maximum mole fraction at pH 5.5 for $[\text{Mo}_7^{\text{VI}}\text{O}_{24}]^{6-}$ is $\sim 75\%$ reported by Cruywagen *et al.*,⁶ and is $\sim 100\%$ by Maximovskaya *et al.*¹⁸ Only species confirmed by several methods (preferably NMR spectroscopy among them) and/or groups are included to our analysis and as a result to the diagrams. The speciation analysis for POM with V^{V} , Nb^{V} , Ta^{V} , Mo^{VI} and W^{VI} as addenda atoms and for majority of archetypes is provided; however, due to lacking a “data base” for POM anions in solution, some speciation works might have been unintentionally overlooked.

Since the majority of POMs have been formed and studied in water, this review focuses on speciation in aqueous solutions, where many applications take place, *e.g.* catalysis and biological application. Although the possible supramolecular assembly of POMs with H-binding and counterions is an important topic for POM chemistry and application, this aspect is not covered here. We first discuss the state of the art and methodology in studying polyanions in solution, which often is a complex issue. This section is followed by the detailed description of POM speciation mostly according to the solution pH and organized by their addenda atoms (V^{V} , Nb^{V} , Ta^{V} , Mo^{VI} and W^{VI}). The common trends in POM solution behavior with the aim of predicting stability trends based on the addenda atom type and structure are discussed throughout the text. Some POM applications performed without POM stability proofs are examined.

2. Methods of POM investigation in solution

The ambiguity of POMs equilibria in solution requires appropriate experimental techniques and careful interpretation of the results. Ideally, a number of different complementary and orthogonal experimental techniques are necessary to study and understand the distribution of species. However, this extensive characterization is rarely done. An exceptional example for a

detailed species analysis is an investigation of acidified orthotungstate solution using electrospray-ionization mass-spectrometry (ESI-MS), ^{183}W -NMR and Raman spectroscopy.¹⁹ The polycondensation product, heptatungstate $[\text{W}_7^{\text{VI}}\text{O}_{24}]^{6-}$, has been proven by NMR and Raman spectroscopies to be the main species in an equilibration mixture at $\text{pH} < 7$, but fails to be detected by ESI-MS due to its ESI-induced dissociation into Lindqvist $[\text{W}_6^{\text{VI}}\text{O}_{19}]^{2-}$ anion (for more details see Section 3.4.1). The reason leading to the different results obtained *via* NMR, Raman and MS is the instability of $[\text{W}_7^{\text{VI}}\text{O}_{24}]^{6-}$ upon ionization. This study also shows that ESI-MS is mainly applicable to stable polyanions, and that the method of investigation should be selected based on its strengths and weaknesses and by taking into account the specific characteristics of a particular POM.

In order to obtain a spectrum as a fingerprint and assign it to a species, the species must be either homogeneous in solution or solid. The POM solution to be examined requires observation over a varying pH, time range and, if applicable, temperature and concentration range. Currently, solution studies no longer use only potentiometry and vibration spectroscopy as their exclusive methods for characterizing metal oxide systems, but various advanced methods complement these studies, including multinuclear NMR spectroscopy, small-angle X-ray scattering (SAXS), X-ray absorption spectroscopy (XAS, consisting of Extended X-ray absorption fine structure (EXAFS) and X-ray absorption near edge structure (XANES)) investigations as well as mass-spectrometry. We briefly summarize the characterization methods in context of their application in POM systems. A detailed description of the basic principles of the method, the recording, evaluation and interpretation of the data can be found in the specialized literature. In the following, the methods for examining POMs in solutions are presented in the order of their chronological appearance.

2.1. Potentiometry

Potentiometric titration was one of the first methods that was used to investigate POM speciation.¹ In principle, the method



involves measuring the hydrogen ion concentration in polyanion solutions as a function of the acid or base added (Fig. 1A) and the total metal ion concentration.^{16,20} The values for the formation constants and stoichiometric coefficients are determined by titration, followed by the iterative calculation of the constants of one species in the presence of another and the entire system being described by calculations in an iterative process. The formation constants K_c are the basis for species distribution diagrams (Fig. 1B) showing various types of POM that exist at different pH values. However, great care should be taken to ensure that a true equilibrium has been reached for each measurement, that the activity coefficient quotients are reasonably constant (use of supporting electrolyte) and that the potentials of the liquid transitions can be controlled. At the moment, potentiometry is outdated due to the availability of other advanced methods that provide a more accurate picture of the processes in solution.

2.2. Electronic and vibrational spectroscopy

Typically, the addenda ions in POMs have d^0 electronic configuration, and as a result, only one absorption band occurs in the UV-vis between 190 and 400 nm due to the oxygen-to-metal charge transfer transition.^{21,22} The spectra of the reduced “heteropoly blue” complexes show intervalence charge-transfer transitions, e.g., $\text{Mo}^V \rightarrow \text{Mo}^{VI}$ at ~ 700 nm.²³ Electronic spectroscopy implies practically no structural information, but it is one of the easiest ways to check POM stability in solution.^{24,25} For example, at pH 1 the Keggin polyanion $[\text{P}^V\text{W}_{12}\text{O}_{40}]^{3-}$ ($\{\text{PW}_{12}\}$) (Fig. 2B) exhibits two intense absorption bands in the UV range, with maxima at about 200 and 263 nm attributed to the p_π - d_π charge-transfer transitions of the $\text{O}_t \rightarrow \text{W}^{VI}$ (O_t – terminal oxygen atom), and p_π - d_π charge-transfer transitions of the $\text{O}_{b,c} \rightarrow \text{W}^{VI}$ ($\text{O}_{b,c}$ – bridge oxygen atoms) (Fig. 2A), respectively.²⁵ The absorption band maximum at 263 nm corresponds to the electron transition $\text{O}_{b,c} \rightarrow \text{W}^{VI}$ of the intact Keggin anion and is shifted to 252.5 nm at pH 3.5

corresponding to mono-lacunary form $[\text{P}^V\text{W}_{11}\text{O}_{39}]^{7-}$ ($\{\text{PW}_{11}\}$), indicating $[\text{P}^V\text{W}_{12}\text{O}_{40}]^{3-}$ decomposition (Fig. 2).²⁵ A detailed description of the hydrolytic stability of $\{\text{PW}_{12}\}$ is given in Section 3.4.2.1.

Infrared, Raman and resonance Raman spectroscopy are broadly used in POM chemistry as diagnostic fingerprints. Similarities of spectral band positions, shapes and relative intensities for two compounds strongly indicate that both have identical structures. The characteristic spectrum region is between 1000 and 400 cm^{-1} where absorptions due to metal-oxygen stretching vibrations occur.

A good agreement between the spectra of crystalline and dissolved polyanions shows that the structure of the dissolved anion is the same as that observed in the solid state. Raman spectroscopy is more frequently used for aqueous solution studies of POMs^{26–29} and IR spectroscopy for studies in both aqueous and nonaqueous solvents.³⁰

2.3. Nuclear magnetic resonance (NMR)

Nuclear magnetic resonance spectroscopy has been carried out on POMs containing NMR-active nuclei, i.e. ^{31}P (natural abundance (NA): $\text{NA}(^{31}\text{P}) = 100\%$; nuclear spin (I): $I = 1/2$), ^{51}V (NA = 99.75%; $I = 7/2$), ^{17}O (NA = 0.04%; $I = 5/2$), ^1H (NA = 99.98%; $I = 1/2$), ^{29}Si (NA = 4.7%; $I = 1/2$), and, later, ^{95}Mo (NA = 15.87%; $I = 5/2$) and ^{183}W (NA = 14.32%, $I = 1/2$), to investigate their solution structures and dynamics since the 1970s.^{31,32} For reliable identification of a POM species in solution, it is desirable, whenever possible, to measure NMR spectra of all NMR-active core components.

2.3.1. Addenda nuclei

^{51}V NMR. So far, the largest number of measurements for POMs has been carried out at ^{51}V , a core nucleus with relatively high sensitivity, which provides spectra with line widths in the range from ~ 10 to ~ 800 Hz in diamagnetic polyanions. The chemical shifts (reference VOCl_3) in isopoly- and heteropolyvanadates fall in

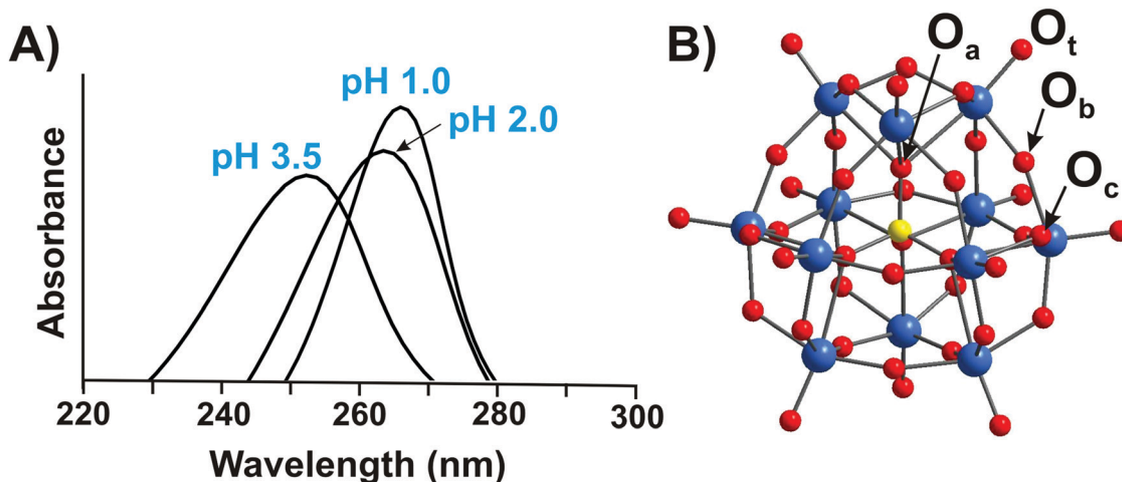


Fig. 2 (A) UV-vis spectra of aqueous solutions of $\text{H}_3[\text{P}^V\text{W}_{12}\text{O}_{40}]$ recorded at pH 1.0; 2.0 and 3.5.²⁵ The decomposition of Keggin anion can be clearly seen already at pH 3.5 by shifting of the maximum absorption from 263 nm to 252.5 nm. For more details see Section 3.4.2.1. (B) Ball-and-stick representation of an α -Keggin type anion $[\text{P}^V\text{W}_{12}\text{O}_{40}]^{3-}$ with the indicated types of oxygen atoms: μ_3 - O_a – oxygen atom connected to heteroatom P^V ; μ_2 - O_b and μ_2 - O_c – two types of bridging atoms in the structure; O_t – terminal oxygen atom. Color code: W, blue; P, yellow; O, red.



the range between -400 to -600 ppm. Even relatively small structural variations result in separable peaks due to the broad chemical shift range.^{33,34}

⁹⁵Mo NMR. Despite the existence of two NMR-active isotopes, ⁹⁵Mo (NA = 15.87%; $I = 5/2$) and ⁹⁷Mo (NA = 9.46%, $I = 5/2$), Mo NMR is less frequently used due to their low natural abundance and their low gyromagnetic ratios: $\gamma(^{95}\text{Mo}) = -1.751 \times 10^7 \text{ rad s}^{-1} \text{ T}^{-1}$ and $\gamma(^{97}\text{Mo}) = -1.788 \times 10^7 \text{ rad s}^{-1} \text{ T}^{-1}$. The ⁹⁵Mo nucleus is generally preferred over ⁹⁷Mo because of its lower quadrupolar moment. Compared with ¹⁸³W NMR, the ⁹⁵Mo NMR signals from a typical asymmetric POMo environment are strongly broadened due to the quadrupole moment, which complicates spectral measurements and their interpretation. The earlier measurements of the ⁹⁵Mo NMR spectra of isopolymolybdates (IPOMos) $[\text{Mo}_2^{\text{VI}}\text{O}_7]^{2-}$, $[\text{Mo}_7^{\text{VI}}\text{O}_{24}]^{6-}$, $[\text{Mo}_6^{\text{VI}}\text{O}_{19}]^{2-}$, and $\alpha/\beta\text{-}[\text{Mo}_8^{\text{VI}}\text{O}_{26}]^{4-}$ were mainly carried out in non-aqueous solutions after ⁹⁵Mo enrichment (96%) and, subsequently, ⁹⁵Mo NMR was applied to study aqueous Mo^{VI} solutions.¹⁸

¹⁸³W NMR. Despite its low sensitivity, the ¹⁸³W NMR is of unique importance in studying polyoxotungstates (POTs). Narrow NMR lines of ¹⁸³W with nuclear spin $I = 1/2$ allow to observe constants of the indirect spin–spin coupling, *i.e.* $^2J(\text{W-P})$, $^2J(\text{W-W})$, which provide structural information. The use of high field spectrometers significantly reduces the sensitivity limitations, although a concentrated sample solution ($\sim 1 \text{ mol L}^{-1}$) and a long acquisition time are still required. A saturated solution of sodium tungstate is recommended as a reference for the chemical shift.³² For POTs, the range of ¹⁸³W-chemical shifts lays between $+260$ and -300 ppm and even up to -670 ppm, if the POT-peroxocomplexes are taken into account.³² When looking at reduced POTs or at POTs with incorporated paramagnetic ions even larger chemical shifts from $+2500$ to -4000 ppm are observed.

⁹³Nb (NA = 100%; $I = 9/2$) and ¹⁸¹Ta (NA = 99.98%; $I = 7/2$) NMR spectroscopy is hardly used for POM investigation, due to the excessive line widths as a result of large quadrupole coupling.

2.3.2. Other nuclei (¹H, ¹⁷O, ³¹P, ¹⁹F, ¹¹B, ²⁷Al)

¹H NMR. The rapid exchange with solvent protons limits the use of high resolution proton NMR for fully inorganic POMs, while integrated resonances of non-labile protons in organic units of hybrid polyoxoanions and counteranions are routinely used for analytical purposes.¹ However, in some cases separate signals for solvent and polyanions were observed. One of the earlier successful examples was published in 1966, when Pope and Varga demonstrated the presence of two central protons in the metatungstate anion, $[\text{H}_2\text{W}_{12}^{\text{VI}}\text{O}_{40}]^{6-}$, using ¹H NMR.³⁵

¹⁷O NMR is more universal, as oxygen is an indispensable element of all POM clusters. There are two main structural types of oxygen atoms in a polyanion (Fig. 2B): terminal oxygens O=M with coordination number one and bridging ones M–O–M with coordination numbers from two to six. NMR lines for oxygen atoms of different types are well resolved and the line's form is characteristic for a structural type of oxygen atom.³¹

Although ¹⁷O is difficult to observe on account of its low natural abundance (0.04%) and its negative quadrupole moment $Q(^{17}\text{O}) = -26 \text{ mB}$, these disadvantages are compensated by its large chemical shift ranging from 1200 to -100 .^{1,31} To overcome the low natural abundance of ¹⁷O (0.04%), target POMs can be enriched with H₂¹⁷O. This allows studying the rates of oxygen-isotope exchange between solvent and POM molecules sites and has been proven useful for understanding of POMs equilibria.^{36,37}

³¹P NMR. The large number and wide variety of heteropoly compounds with phosphorus as a heteroatom led to a significant development of ³¹P NMR, which shows a high sensitivity of its chemical shift based on POMs composition. In the ³¹P NMR spectra of Keggin type H₃[P^VW^{VI}O₄₀] (Fig. 2B) and the majority of its derivatives, each form is represented only by one signal in the chemical shift range between -15 and -2.5 ppm (relative to 85% H₃PO₄), which allows to directly detect several coexisting species and to determine their concentrations.³⁸

Other nuclei. ¹¹B (NA = 80.42%; $I = 3/2$), ³⁹F (NA = 100%; $I = 1/2$), ⁴⁰Al (NA = 100%; $I = 5/2$)⁴¹ and ¹⁹⁵Pt (NA = 33.7%; $I = 1/2$)⁴² NMR spectroscopy are not as often used, but equally important for POM solution investigation.

If suitable NMR-active cations are present, NMR measurements can also probe cation–POM interactions in solution, which can affect the transformation between POM species.⁴³ Using ⁷Li,⁴⁴ ²³Na⁴⁵ and ¹³³Cs⁴⁶ NMR spectroscopy, the Nyman group successfully studied the cationic association with various POMs in solution.

2.4. Mass-spectrometry (MS)

Electrospray-ionization mass-spectrometry (ESI-MS) is suitable for the elucidation of solution phase equilibria of stable upon ionization anions, since it enables semi-quantitative detection of both cationic and anionic species in aqueous solvents with excellent detection limits. POMs are ideal candidates for mass-spectrometry studies since they exhibit complex isotopic envelopes resulting from the high number of stable isotopes as for tungsten (¹⁸²W, 26.5%; ¹⁸³W, 14.3%; ¹⁸⁴W, 30.6%; ¹⁸⁶W, 28.4%) or molybdenum (⁹²Mo, 14.8%; ⁹⁴Mo, 9.3%; ⁹⁵Mo, 15.9%; ⁹⁶Mo, 16.7%; ⁹⁷Mo, 9.6%; ⁹⁸Mo, 24.1%; ¹⁰⁰Mo, 9.6%), and are intrinsically charged.^{47,48} However, the experiments must be carefully designed in order to obtain reliable data without overinterpretation of gas phase data for the solution chemistry.⁴⁹ While ESI-MS does not yield any information beyond the mass-to-charge ratio of the analyte, it has high sensitivity and does not impose too many requirements on the system to be analyzed, and time-resolved data can be obtained on very dilute solutions. ESI-MS measurements have been used for comprehensive POM speciation studies with all kind of addenda atoms and significantly contribute to the speciation analysis given in detailed in Section 3.



2.5. Small angle X-ray scattering (SAXS)

SAXS is a well-established non-destructive method for probing the size, shape, reactivity, and interactions of dissolved species.⁵⁰ This method is very powerful, but so far, an underutilized technique to obtain speciation information on POM solutions. SAXS is fundamentally similar to X-ray crystallography, where a sample is irradiated by a collimated monochromatic X-ray beam.⁵¹ Like nanoparticles and quantum dots, many POMs exhibit high net charge, contain high electron-density elements (W, Mo and other metals) and therefore scatter X-rays strongly. Since POMs are molecular by definition, solutions in which the clusters are stable must be absolutely monodisperse, and their X-ray scattering data can be simulated very accurately by applying solid-state crystal structure data sets. To date, many POM classes have been thoroughly investigated using SAXS, including POMs of group V (Nb^{V} and Ta^{V}),^{50,51} POTs⁵² and their complexes with actinides,⁵³ large reduced POMs⁵⁴ as well as POM supramolecular assemblies.⁵⁵ SAXS can be successfully used to investigate speciation in catalytic systems, as one representative example, the Co-containing POTs speciation as a function of pH, buffer salts, and addition of a chemical oxidant during water oxidation catalysis has been carefully studied.⁵⁶

2.6. Other methods

POMs have a rich electrochemistry associated with both reduction of tungsten or molybdenum⁵⁷ and redox-reaction of heterometals (*i.e.*, incorporated cobalt, ruthenium, iridium, or nickel). These characteristic redox wave peaks can be used to identify the number of terminal oxygen atoms, metastable hydrolysis fragments, new isomers and reduced anions.^{58,59}

Extended X-ray absorption fine structure (EXAFS) and X-ray absorption near edge structure (XANES) are valuable techniques to probe both the local coordination environment and the

oxidation state of POM's atoms either in solution or in solid-state materials. Each kind of atom in the POM cluster can be accessed individually and an average spectrum for each element is observed.⁶⁰ Despite XAS (X-ray absorption spectroscopy) being a powerful technique, there are just a limited number of examples for their usage in POM structure analysis.^{61,62}

Dynamic light scattering (DLS) is aimed to determine whether particles are formed in solutions and, if present, to examine their size.⁶³ DLS has found its broadest usage in monitoring POM stability during catalytic reactions (*e.g.* water splitting systems⁶⁴).

3. POM speciation in aqueous solutions

3.1. Polyoxovanadates (POVs)

This section is divided into isopoly- and heteropolyvanadates. First, the general behavior of POVs in solution is described, and second, the characteristics of tri-, tetra-, penta- and decavanadates, as well as polyoxophosphovanadates are detailed in the order of increasing nuclearity.

3.1.1. Isopolyoxovanadates (IPOVs)

3.1.1.1. Ion-distribution diagram for IPOVs. A great variety of discrete isopolyvanadates (IPOVs) has been characterized in the solid state isolated from aqueous and non-aqueous solutions which include $[\text{V}_3\text{O}_9]^{3-}$ (Fig. 3C),⁶⁵ $[\text{HV}_4\text{O}_{12}]^{3-}$ (Fig. 3E)⁶⁶ and $[\text{V}_5\text{O}_{14}]^{3-}$ (Fig. 3G)⁶⁷ with tetrahedrally coordinated vanadium (Table 1) and $[\text{V}_{10}\text{O}_{28}]^{6-}$ $\{\text{V}_{10}\}$ ⁶⁸ (Fig. 5), $[\text{V}_{12}\text{O}_{32}]^{4-}$,⁶⁹ $[\text{V}_{13}\text{O}_{34}]^{3-}$,⁷⁰ and $[\text{V}_{15}\text{O}_{42}]^{9-}$ ⁷¹ (Fig. 6) with octahedrally coordinated V^{V} ions. The maximum species concentration and the equilibrium of IPOVs are influenced by pH, ionic strength, temperature, and vanadate concentrations. The most commonly used speciation model (Fig. 4 and Table 1) was first

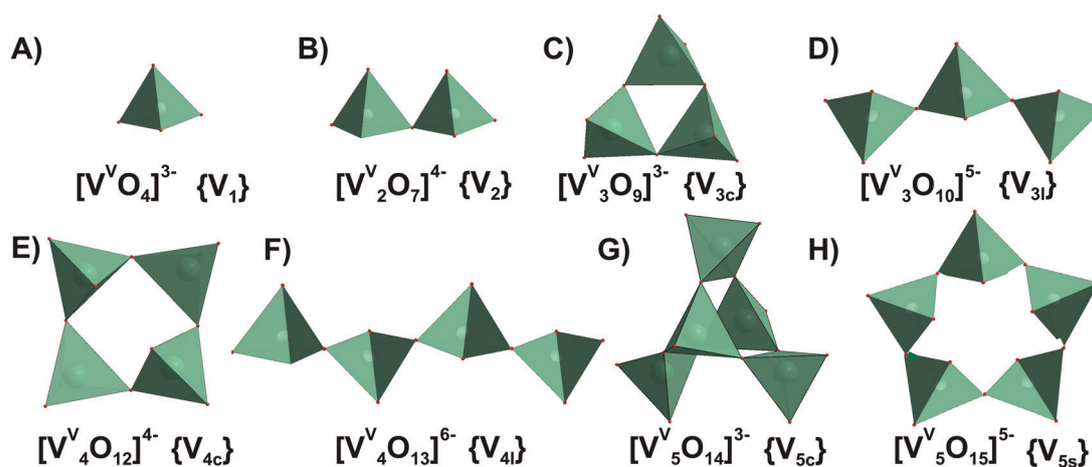


Fig. 3 IPOVs with the nuclearity up to 5 addenda atoms and tetrahedral coordination of V^{V} ((A)–(H)), which can be present in aqueous media. Tri- and tetraavanadates are presented in two forms – cyclic ($\{\text{V}_{3\text{c}}\}$, (C) and $\{\text{V}_{4\text{c}}\}$, (E)) and linear ($\{\text{V}_{3\text{l}}\}$, (D) and $\{\text{V}_{4\text{l}}\}$, (F)). The linear forms of $\{\text{V}_{3\text{l}}\}$ and $\{\text{V}_{4\text{l}}\}$ have never been obtained in solid state. Pentavanadate is depicted in two cyclic forms: $[\text{V}_5^{\text{V}}\text{O}_{14}]^{3-}$ ($\{\text{V}_{5\text{c}}\}$, (G)),⁶⁷ which was synthesized from non-aqueous solution and $[\text{V}_5^{\text{V}}\text{O}_{15}]^{5-}$ ($\{\text{V}_{5\text{s}}\}$, (H)),^{78,79} which has never been crystallized. Color code: $\{\text{VO}_4\}$, green; O, red. In abbreviations the subscript “c” stands for “cyclic”, “l” – for “linear”, “s” – for “star”.



Table 1 IPOVs stoichiometry, formation constants, chemical shifts and pK_a values

IPOV species	<i>q, p</i> ^a	pK _a	Isolated in solid state	δ(⁵¹ V), ppm ^{80,87 b}	Formation constants ^a lg K _c at 25 °C according to					
					Sigel ^{c 84}	McCann ^{d 85}	Larson ^{d 86}	Tracey ^{d 78}	Elvingson ^{e 79}	Heath ^{f 87}
[V ^V O ₄] ³⁻ (Fig. 3A)	-2, 1		Yes	-541.2	-21.31	> -23	> -22			~ -19
H[V ^V O ₄] ²⁻	-1, 1	13.4	No	-538.8	-7.91	-9.02	-8.75	-8.8	-8.17	-7.1
H ₂ [V ^V O ₄] ⁻	0, 1	7.91	No	-560.4						
[V ₂ O ₇] ⁴⁻ (Fig. 3B)	-2, 2		Yes ⁸¹	-561.0	-15.13	-19.0	-18.60		-16.19	-12.8
H[V ₂ O ₇] ³⁻	1, 2	9.74	No	-563.5	-5.39	-7.47	-7.30		-5.85	-3.9
H ₂ [V ₂ O ₇] ²⁻	0, 2	8.29	No	-572.7	2.90	2.15	2.30	2.5	2.65	3.3
H[V ₃ O ₁₀] ⁴⁻ (Fig. 3D)	-1, 3		No	~ -570						-11.4
[V ₄ O ₁₃] ⁶⁻ (Fig. 3F)	-2, 4		No	Between -566 and -585	-8.50	-16.1			-9.98	28.1
H[V ₄ O ₁₃] ⁵⁻	-1, 4	8.9	No	Between -566 and -585	0.4					
[V ₄ O ₁₂] ⁴⁻ (Fig. 3E)	0, 4		Yes ⁶⁶	-577.6	10.04	7.0	7.60	8.4	9.24	11.6
[V ₅ O ₁₅] ⁵⁻ (Fig. 3H)	0, 5		No	-586.0	12.43	7.5			11.17	14.5
[V ₁₀ O ₂₈] ⁶⁻ (Fig. 5)	4, 10		Yes ⁶⁸	-422, -496, -513	51.98				50.28	
H[V ₁₀ O ₂₈] ⁵⁻	5, 10	6.14	No	-424, -500, -516	58.12				56.90	
H ₂ [V ₁₀ O ₂₈] ⁴⁻	6, 10	3.68	Yes ⁸²	-425, -506, -524	61.80				61.07	
H ₃ [V ₁₀ O ₂₈] ³⁻	7, 10	1.57	Yes ⁸³	-427, -515, -534	63.37				62.93	
[V ^V O ₂] ⁺	2, 1		Yes	-545.0	6.97				7.00	

^a Stoichiometric coefficients and formation constants for the reaction $p[\text{H}_2\text{V}^{\text{V}}\text{O}_4]^- + q\text{H}^+ \rightleftharpoons [\text{H}_z\text{V}_p^{\text{V}}\text{O}_m]^{n-} + (q/2 + p - z/2)\text{H}_2\text{O}$. ^b δ⁵¹V relative to VOCl₃. ^c Ionic strength μ = 0.6 M (NaCl). ^d Without electrolyte addition. ^e μ = 0.15 M (NaCl). ^f μ = 2 M (NaClO₄).

constructed about thirty years ago as a result of rigorous research using ⁵¹V NMR investigations in conjunction with potentiometric titration studies.^{72,73} The proposed speciation model (Fig. 4) postulates, that in aqueous vanadium solutions of pH > 6, polymers with one to five V atoms (Fig. 3) are formed, where many of these IPOVs also undergo several individual protonation steps. In contrast, under acidic

conditions, only two major species {V₁₀} and [V^VO₂]⁺ (Fig. 4 and Table 1) have been reported.

Since ⁵¹V NMR is a powerful technique to determine POV speciation and easily accessible, the use of other techniques is rarely reported. ESI-MS analysis of ammonium metavanadate (NH₄)₃[V^VO₄] aqueous solution at pH 4.5⁷⁴ and 6⁷⁵ revealed a series of known (protonated {V₁₀} and [V^VO₄]³⁻) and

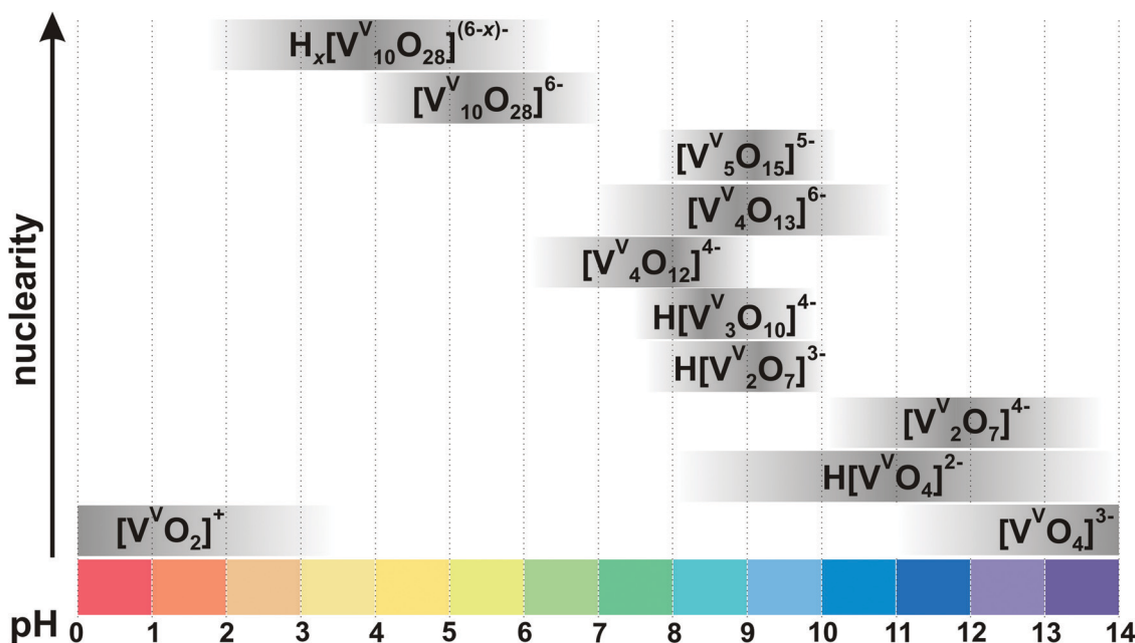


Fig. 4 Speciation of IPOVs in an aqueous solution with a total concentration of vanadium more than 0.1 mM based on ref. 34, 72, 78, 79 and 84–88. The maximum intensity of grey color in each box with a single species corresponds to its maximum concentration in the chosen pH region, e.g. the maximum concentration of [V₂O₇]⁴⁻ is at pH 12. The grey boxes along the y-axis are positioned according to increasing nuclearity, but do not show the domination over other species at a certain pH range. *x* in H_{*x*}[V₁₀O₂₈]^{6-*x*-} is 1–3. The molecular structures of the clusters are depicted in Fig. 3 and 5. The ⁵¹V chemical shifts, formation constants and pK_a values are summarized in Table 1.



“previously unknown” polyoxovanadate anions and cations, which might have been formed in gas phase due to ionization.

3.1.1.2. Tri-, tetra- and penta-vanadates. The structures of the trivanadate (Fig. 3C and D) and tetravanadate (Fig. 3E and F) in water were debated for a long time and numerous studies lead to contradictory conclusions.⁷⁶ The various structural suggestions include linear (Fig. 3D and F) or cyclic (Fig. 3C and E) trivanadate or tetravanadate ions exhibiting tetra- or penta-coordinated vanadium. Nowadays, a consensus has been reached that a linear form of tri- and tetranuclear vanadates, $[V_3^V O_{10}]^{5-}$ and $[V_4^V O_{13}]^{6-}$ (Fig. 3D and F), does exist in aqueous solution at pH < 10, which was confirmed by ^{51}V and ^{17}O NMR (Fig. 4).⁷⁷ Up to date only salts with cyclic $[V_3^V O_9]^{3-}$ (Fig. 3C), $[HV_4^V O_{12}]^{3-}$ (Fig. 3E) and $[V_5^V O_{14}]^{3-}$ (Fig. 3G) have been obtained from non-aqueous media.^{65–67} The isolation of the discrete linear tri- and tetravanadate (Fig. 3D and F) from aqueous solution has failed so far because the crystallization always goes hand in hand with the polymerization of the anion to “endless” metavanadate chains $[(V^V O_4)^{3-}]_n$.

3.1.1.3. Decavanadate $[V_{10}^V O_{28}]^{6-}$ stability. $\{V_{10}\}$ is stable for days at neutral pH;²⁹ while at pH values higher than 7 it, at least in part, turns into structurally and functionally different smaller oxovanadates such as the monomeric $[V^V O_4]^{3-}$ (Fig. 3A), dimeric $[V_2^V O_7]^{4-}$ (Fig. 3B) or tetrameric $[V_4^V O_{12}]^{4-}$ (Fig. 3E) with a half-life time depending on vanadium concentration, ionic strength and temperature.⁸⁹ In the basic pH region, $\{V_{10}\}$ will ultimately convert to the colorless metavanadates $H_x[V^V O_4]^{(3-x)-}$ ($x = 0–2$) (Fig. 4), although near neutral pH, this process may take up to three weeks depending on ionic

strength and temperature (Fig. 5). At neutral pH, $\{V_{10}\}$ can be destructed to smaller IPOVs and therewith being removed by boiling the solution.⁹⁰

^{51}V NMR and UV-vis spectroscopic investigations revealed that at pH 7 and 37 °C the half-life of $\{V_{10}\}$ decomposition is 2 h in serum-free Dulbecco's modified Eagle's medium (DMEM: CaCl₂, 0.2 g L⁻¹; KCl, 0.4 g L⁻¹; NaCl, 6.4 g L⁻¹; Na₂HPO₄, 0.109 g L⁻¹; Na₂CO₃, 3.7 g L⁻¹; glucose, 1 g L⁻¹, and 20 proteinogenic amino acids).⁹¹ At room temperature in physiological serum (pH = 7, NaCl 0.9%) the half-life of 12 h was estimated for $\{V_{10}\}$.⁹² In brain-heart infusion medium (BHI: calf brain, 12.5 g L⁻¹; beef heart, 5 g L⁻¹; peptone, 10 g L⁻¹; D-glucose, 2 g L⁻¹; NaCl, 5 g L⁻¹; Na₂HPO₄, 2.5 g L⁻¹) at a slightly higher pH of 7.2, $\{V_{10}\}$ decomposes quickly with measurable decay in as little as one hour and the dominant species after 19 hours is tetrameric $[V_4^V O_{12}]^{4-}$ (Fig. 3E).⁹³ Almost no $\{V_{10}\}$ was detected after 42 h while tetravanadate is still the dominating species after 42 h and 66 h (Fig. 5). The half-life of $\{V_{10}\}$ in aqueous buffered solution (KCl, 0.1 M; MgCl₂, 5 mM; ethylene glycol-bis(β-aminoethyl ether)-N,N,N',N'-tetraacetic acid (EGTA), 0.5 mM; imidazole, 10 mM) at pH = 7.4 with $[\{V_{10}\}] = 5$ mM at 25 °C is 43 h, and under the same conditions with a higher concentration $[\{V_{10}\}] = 50$ μM decreases to 19 h (Fig. 5).⁹⁴ In a view of the significant decomposition of $\{V_{10}\}$ in diluted solutions at room temperature, it is advisable to perform the experiments involving $\{V_{10}\}$ in more concentrated solutions and limit them to a few hours when working at room temperature. At increased 37 °C and keeping the same pH of 7.4 however in mitochondrial respiration buffer (MRB: sucrose, 0.2 M; KH₂PO₄, 5 mM; KCl, 10 mM; MgCl₂, 5 mM; Tris-HCl, 10 mM; pyruvate 5 mM; malate, 0.5 mM) 5 mM $\{V_{10}\}$

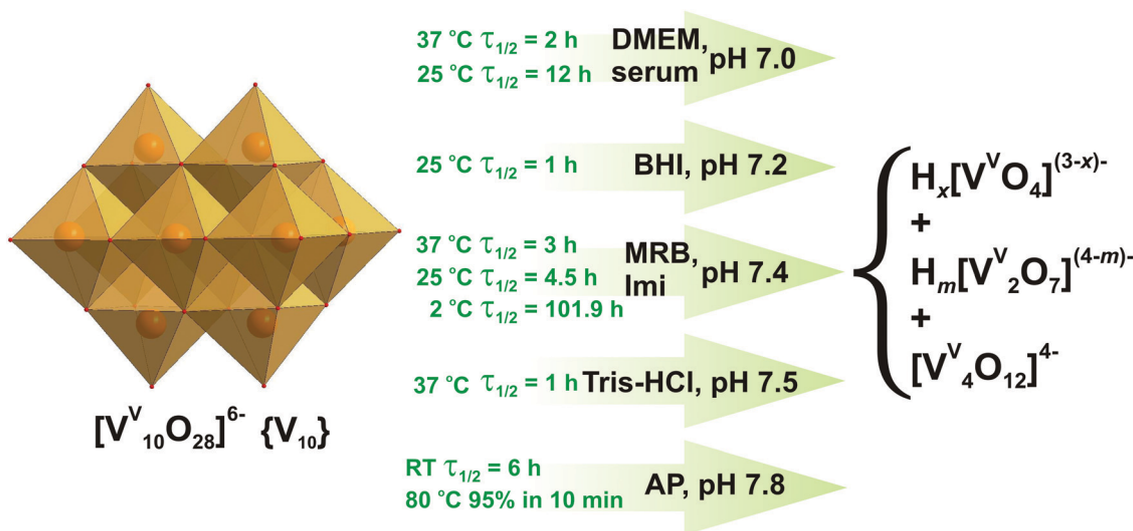


Fig. 5 Decavanadate $\{V_{10}\}$ decomposition in aqueous solutions between pH 7.0 and 7.8 based on ^{51}V NMR, UV-vis and XAS investigations.^{61,89,91–95} Half-life of $\{V_{10}\}$ decomposition is given for a total V concentration of 5 mM. DMEM – serum-free Dulbecco's modified Eagle's medium, CaCl₂, 0.2 g L⁻¹; KCl, 0.4 g L⁻¹; NaCl, 6.4 g L⁻¹; Na₂HPO₄, 0.109 g L⁻¹; Na₂CO₃, 3.7 g L⁻¹; glucose, 1 g L⁻¹, and 20 proteinogenic amino acids; MRB – mitochondrial respiration buffer, sucrose, 0.2 M; KH₂PO₄, 5 mM; KCl, 10 mM; MgCl₂, 5 mM; Tris-HCl, 10 mM; pyruvate 5 mM; malate, 0.5 mM; serum – physiological serum, NaCl, 0.9%; BHI – brain-heart infusion medium (calf brain, 12.5 g L⁻¹; beef heart, 5 g L⁻¹; peptone, 10 g L⁻¹; D-glucose, 2 g L⁻¹; NaCl, 5 g L⁻¹; Na₂HPO₄, 2.5 g L⁻¹, pH 7.2); Imi – imidazole buffer (KCl, 0.1 M; MgCl₂, 5 mM; EGTA, 0.5 mM; imidazole, 10 mM); Tris-HCl – 2 mM Tris-HCl buffer; AP – alkaline phosphatase buffer (NaCl, 5 M; Tris-HCl, 1 M; MgCl₂, 1 M). The structures of $H_x[V^V O_4]^{(3-x)-}$ ($x = 0–2$), $H_m[V_2^V O_7]^{(4-m)-}$ ($m = 0–2$) and $[V_4^V O_{12}]^{4-}$ are shown in Fig. 3A, B and E. Color code: $\{VO_6\}$, light orange; O, red.



solution shows the short half-life of just 3 h.⁹⁵ A UV-vis kinetics study of the $\{V_{10}\}$ decay in the presence of alkaline phosphatase buffer (AP: NaCl, 5 M; Tris-HCl, 1 M; MgCl₂, 1 M) at pH 7.8, carried out by following the peak disappearance at 430 nm ($p_{\pi-d_{\pi}}$ charge transfer of the $O_{\text{bridg.}} \rightarrow V^V$), indicated that $\{V_{10}\}$ degrades by approximately 15% per hour at room temperature ($t_{1/2} = 6$ h under first order kinetics) (Fig. 5).⁹³ About 95% of $\{V_{10}\}$ decomposed in 10 minutes to tetravanadate $[V_4O_{12}]^{4-}$ (Fig. 3E) in the same AP buffer (pH = 7.8) at increased 80 °C.

Under acidic conditions (pH 5.4) in Schneider's insect medium (SIM: Na₂HPO₄, 0.7 g L⁻¹; MgSO₄, 3.7 g L⁻¹; KCl, 1.6 g L⁻¹; KH₂PO₄, 0.45 g L⁻¹; NaCl, 2.1 g L⁻¹ and 20 proteinogenic amino acids), the decomposition of $\{V_{10}\}$ is substantially slower and after 2 hours, only 0.06 molar equivalent of $\{V_4\}$ (Fig. 3E) was detectable and the rest remained as $\{V_{10}\}$.⁹³

Recently EXAFS and XANES spectroscopic investigations were applied to examine the V oxidation state in $\{V_{10}\}$ solution.⁶¹ Under the physiological temperature of 37 °C $\{V_{10}\}$ began to decompose after about 1 h at pH 7.5 (2 mM Tris-HCl buffer), while at room temperature $\{V_{10}\}$ was present for the entire acquisition period (ca. 7 h). According to the XANES analysis the solution of $\{V_{10}\}$ at pH 7.5 and 37 °C contained 81.3% of $[V^VO_4]^{3-}$ (Fig. 3A) and 18.7% of the reduced oxidovanadium (V^{IV}) species ($[V^{IV}O]^{2+}$ and its complexes), evidencing $\{V_{10}\}$ reduction upon interaction with actin, and meaning that all $\{V_{10}\}$ species have been completely decomposed in solution visible also by a color change from yellow to colorless. The $\{V_{10}\}$ stability studies performed in different media and temperatures lead to sometimes contradictory results, and it is recommendable to check the $\{V_{10}\}$ integrity under the applied conditions.

3.1.1.4. The use of speciation data to understand the biological activities of decavanadate. The applicability of this review will be shown in this section using decavanadate as an example, since among all IPOVs (Fig. 3), $\{V_{10}\}$ (Fig. 5) demonstrates many important roles in fundamental biological processes.^{61,90,96-101}

Three IPOVs: $[V^VO_4]^{3-}$ (Fig. 3A), $[V_4^VO_{12}]^{4-}$ (Fig. 3E) and $\{V_{10}\}$ (Fig. 5), were tested as antibacterial agents against six strains of *Streptococcus pneumoniae* (penicillin-intermediate-resistant IID553, IID554 and penicillin-resistant BS225, BS234, BS259, BS269) without the use of any additional antibiotic.¹⁰² Prior to the evaluation of the minimum inhibitory concentrations (MIC), all IPOVs were incubated at 37 °C for 20 h in Mueller-Hinton Agar medium (agar, 17 g L⁻¹; beef infusion solids, 2.0 g L⁻¹; casein hydrolysate, 17.5 g L⁻¹; starch, 1.5 g L⁻¹; pH = 7.3), which according to works^{61,89,93,94} (Fig. 5) should lead to almost complete hydrolysis of $\{V_{10}\}$. The ion-distribution diagram for IPOVs (Fig. 4) clearly shows that $\{V_{10}\}$ is the stable and dominant species between pH 4 and 7, and the additional heating used for incubation only aggravates the $\{V_{10}\}$ hydrolysis rate. MICs of all three IPOVs, $[V^VO_4]^{3-}$, $[V_4^VO_{12}]^{4-}$ and $\{V_{10}\}$, were in the concentration range between 4 and 32 μg mL⁻¹ with slight lower numbers for $\{V_{10}\}$ solution (4–8 μg mL⁻¹). The authors, without considering the $\{V_{10}\}$ decomposition into mono-, di- and tetravanadates during the incubation, made a

conclusion about the superiority of decavanadate as an anti-bacterial agent, which in fact definitely requires a more careful study.

An example of the proper use of decavanadate is when applied as an inhibitor for bovine pancreatic ribonuclease A (RNase A).¹⁰³ Before evaluating the thermodynamic parameters for the $\{V_{10}\}$ binding to RNase A applying isothermal titration calorimetry (ITC), a stock solution of decavanadate was prepared with a pH between 3 and 5 and checked after 24 hours to verify that equilibrium was established. All experiments have been performed at 25 °C in 30 mM sodium succinate buffer (pH 6.0), which, in accordance with the IPOVs ion-distribution diagram (Fig. 4) and $\{V_{10}\}$ decomposition scheme (Fig. 5), should not lead to decavanadate decomposition. As a result, it is justified to conclude that the intact cluster interacts with RNase A.

Crans and co-authors recently conducted an even more thorough analysis of decavanadate speciation to study its effect on the growth of *Mycobacterium smegmatis* (*M. smeg*) and *Mycobacterium tuberculosis* (*M. tb*).¹⁰¹ ⁵¹V NMR spectra were recorded at pH between 5.8 and 6.8, where there should not have been any decavanadate hydrolysis according to Fig. 4. The spectra showed 100% intact decavanadate $\{V_{10}\}$ in stock solution and growth media with $\{V_{10}\}$ but without bacterial cells. The addition of growth media containing cells to $\{V_{10}\}$ stock solution immediately caused some decomposition of $\{V_{10}\}$.¹⁰¹ These results lead to the conclusion, that $\{V_{10}\}$ can interact with some cell components, leading to $\{V_{10}\}$ decomposition and emphasize the importance to investigate the influence of macromolecules on POM speciation. Ramos *et al.*⁹⁷ confirmed the influence of macromolecules on decavanadate stability by investigating the interaction of $\{V_{10}\}$ solutions with G-actin. Different macromolecules show various effects on decavanadate stability, thus, the $\{V_{10}\}$ half-life increases from 5 to 27 h in the presence of G-actin and from 5 to 18 h in the presence of sarcoplasmic reticulum vesicles at 22 °C and pH 7.5 (Tris, 2 mM; CaCl₂, 0.2 mM; KCl, 100 mM; MgCl₂, 2 mM). The addition of ATP to the medium decreases the half-life of $\{V_{10}\}$ from 27 to 10 h, while in the presence of phosphatidylcholine liposomes or myosin $\{V_{10}\}$ stability does not change. It is assumed that the decavanadate interaction with G-actin, favored by the G-actin polymerization, stabilizes the presence of intact $\{V_{10}\}$.

3.1.2. Heteropolyoxovanadates (HPOVs)

3.1.2.1. Speciation in phospho-vanadate solutions. Among the variety of heteroatoms present in vanadates only the vanadate-phosphate speciation has been investigated systematically.¹⁰⁴⁻¹⁰⁶ In the aqueous $H^- - H_2V^VO_4^- - H_2P^VO_4^-$ solution in the presence of 0.15¹⁰⁵ or 0.60¹⁰⁴ M NaCl at 25 °C three types of phosphovanadates – $[H_kP^kV^VO_7]^{(4-k)-}$ ($k = 1-4$) ($\{PV\}$), $[H_mP_2^mV^VO_{10}]^{(5-m)-}$ ($m = 1, 2$) ($\{P_2V\}$), $[H_nP^nV^VO_{14}]^{(9-n)-}$ ($n = 3-5$) ($\{PV_{14}\}$; Fig. 6A) – have been identified by pH-potentiometry and high-field ³¹P and ⁵¹V NMR spectroscopy to be present between pH 1 and 10 (Fig. 7 and Table 2).

The kinetically labile $\{PV\}$ and $\{P_2V\}$ have been reported only in solutions and never been crystallized. The predominant



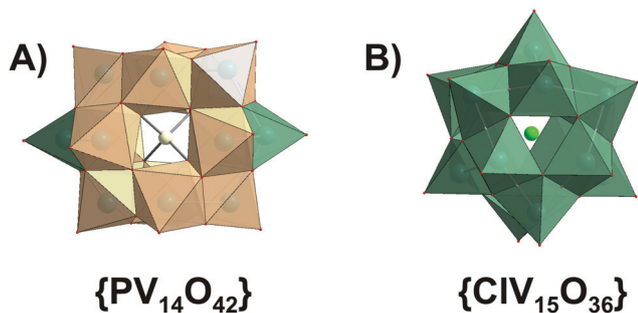


Fig. 6 Polyhedral representation of (A) $[H_n P^V V^V O_{42}]^{(9-n)-}$ ($n = 3-5$ and $[H_6 V_{12} V_2^IV O_{38} (P^V O_4)]^{5-}$ ($\{PV_{14}O_{42}\}$) and (B) $[V_7^V V_8^IV O_{36} (Cl)]^{6-}$ ($\{ClV_{15}O_{36}\}$). Color code: $\{VO_6\}$, light orange, $\{VO_4\}$ and $\{VO_5\}$, green; O, red; P, yellow; Cl, green.

equilibrium species, $\{PV_{14}\}$ (Fig. 6A), which has a trans-bicapped α -Keggin structure, has been known both in solution ($[V]/[P] = 0.5-14$, $pH = 1-6$) and solid state.¹⁰⁷ Reaching equilibrium in the system $H^+ - H_2V^VO_4^- - H_2P^VO_4^-$ is generally fast, except under acidic condition. Here, the formation of $\{PV_{14}\}$ species (Fig. 6A) is even slower and requires at least three months to establish a complete equilibrium. Naturally, high vanadate total concentrations favor the formation of $\{PV_{14}\}$ species, but also that of decavanadate $\{V_{10}\}$.

3.1.2.2. Speciation of mixed valence HPOVs and the use of speciation data to understand their biological activities. Speciation studies were carried out for mixed valence polyoxovanadates $K(NH_4)_4[H_6V_{12}V_2^IV O_{38}(P^V O_4)] \cdot 11H_2O$ (Fig. 6A) and $[(CH_3)_4N]_6[V_7^V V_8^IV O_{36}(Cl)]$ (Fig. 6B) by ^{51}V NMR and EPR spectroscopic studies in aqueous solution and in Luria-Bertani (LB) medium ($pH 7.4$; tryptone, $10 g L^{-1}$; yeast extract, $5 g L^{-1}$; NaCl,

$10 g L^{-1}$) to understand mixed valence HPOVs chemoprotective activity against the alkylating agent diethylsulphate (DES) in *Escherichia coli* DH5 α cultures.¹⁰⁸ $[(CH_3)_4N]_6[V_7^V V_8^IV O_{36}(Cl)]$ (Fig. 6B) is more stable in LB than in pure aqueous solution, and is able to react with increasing amounts of DES. According to ion-distribution diagram (Fig. 7), fully oxidized $\{PV_{14}\}$ (Fig. 6A) should be stable between $pH 3.5$ and 5.5 . The tested reduced analog $[H_6V_{12}V_2^IV O_{38}(P^V O_4)]^{5-}$ also decomposes and rearranges rapidly in LB ($pH = 7.4$), resulting in complete absence of chemoprotective activity against DES. Decomposition products, $[V^VO_4]^{3-}$ and $\{V_{10}\}$, react poorly, or even do not react, with the alkylating agent. The observation of chemoprotective activity for $[(CH_3)_4N]_6[V_7^V V_8^IV O_{36}(Cl)]$ (Fig. 6B) or its absence for $K(NH_4)_4[H_6V_{12}V_2^IV O_{38}(P^V O_4)] \cdot 11H_2O$ (Fig. 6A) against DES is clearly dependent on the chemical nature and stability of the soluble species in the culture medium after addition of the alkylating agent.

3.2. Polyoxoniobates (PONbs) and polyoxotantalates (POTas)

In contrast to POVs, POMos and POTs, polyoxoniobates (PONbs) and polyoxotantalates (POTas) can be stabilized only under basic conditions due to their high negative charge, and their chemistry was initially dominated by the Lindqvist anion $[M_6O_{19}]^{8-}$ ($M = Nb^V, Ta^V$) in solution and by its alkali salts in the solid state.¹ Owing to the narrow working pH region (> 7), the low solubility, and the low reactivity of niobate and tantalate species, the progress of PONb and POTas chemistry is far behind that of POMo or POT chemistry.¹⁰⁹ This chapter is devoted only to isopolyniobates (IPONbs) and -tantalates (IPO-Tas) due to the lack of information about heteropoly PONbs and POTas stability and speciation in solution.

3.2.1. Isopolyniobates (IPONbs) and -tantalates (IPOTas)

3.2.1.1. Ion-distribution diagram. The Lindqvist hexaniobate $[Nb_6^VO_{19}]^{8-}$ ($\{Nb_6\}$, Fig. 8A)¹¹⁰ and the related decaniobate $[Nb_{10}^VO_{28}]^{6-}$ ($\{Nb_{10}\}$, Fig. 8B)^{111,112} were for a long time the only known PONbs. Spectroscopic and potentiometric studies have shown that the Lindqvist ion is indeed the dominant species of Nb in solutions at pH higher than 7 and room temperature (Fig. 9).¹¹³ The heptaniobate $[Nb_7^VO_{22}]^{9-}$ ($\{Nb_7\}$, Fig. 8C) has been recently detected only in solution at $pH 9$ ($100 mM H_3BO_3$),^{114,115} as a part of larger structures. The formation of other Nb or Ta POM archetypes cannot be reproducibly achieved by pH change, because Nb and Ta solutions are difficult to retain except at high $pH > 12$. Therefore, forming other polyoxoniobate and polyoxotantalate clusters requires finding conditions in which the Nb or Ta precursor is soluble, without $\{Nb_6\}$ being the dominant species. Using hydrothermal synthesis allows the dissolution of Nb_2O_5 precursor without adding a huge excess of base that favors $\{Nb_6\}$ precipitation. Despite the growing number of new PONbs,¹⁰⁹ their solution behavior needs to be investigated in more details. The speciation diagram for IPONbs is rather incomplete and includes four species $\{Nb_6\}$, $\{Nb_7\}$, $\{Nb_{10}\}$ and $\{Nb_{24}\}$ (Fig. 9 and Table 3). It should be noted that aqueous polyniobate chemistry at pH lower than 5 remains to be explored.

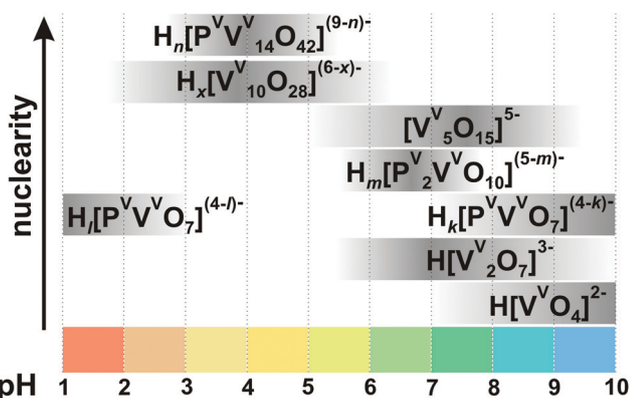


Fig. 7 Speciation in the aqueous solution of $H^+ - H_2V^VO_4^- - H_2P^VO_4^-$ with $0.15 M NaCl$ at $25 ^\circ C$ and $[V]_{tot} = 5 mM$, $[P] = 60 mM$ based on ref. 105. Protonation degree: $H_l[P^V V^V O_7]^{(4-l)-}$ ($k = 3-4$); $H_k[P^V V^V O_7]^{(4-k)-}$ ($k = 1-2$), $H_m[P_2^V V^V O_{10}]^{(5-m)-}$ ($m = 1, 2$), $H_n[P^V V^V O_{42}]^{(9-n)-}$ ($n = 3-5$), $H_x[V^V O_{28}]^{(6-x)-}$ ($x = 1-3$). The maximum intensity of grey color in each box with a single species corresponds to its maximum concentration in the chosen pH region. The grey boxes along the y -axis are positioned according to increasing nuclearity, but do not show the domination over other species at a certain pH range. The structures of the species are presented in Fig. 3, 5 and 6. The ^{51}V chemical shifts, formation constants and pK_a values are summarized in Table 2.



Table 2 Phospho-vanadates stoichiometry, formation constants, chemical shifts and pK_a values

Phospho-vanadate species	q, p, r^a	pK_a^{105}	Isolated in solid state	$\delta(^{51}\text{V})$, ppm ^{b 105}	$\lg K_c$ at 25 °C ^{a,c}
$\text{H}_3[\text{P}^{\text{V}}\text{V}_{14}\text{O}_{42}]^{6-}$ (Fig. 6A)	9, 14, 1		Yes ¹⁰⁷	-521, -572, -589	89.39 (0.19)
$\text{H}_4[\text{P}^{\text{V}}\text{V}_{14}\text{O}_{42}]^{5-}$	10, 14, 1	4.5	No	-530, -580, -598	93.93 (0.04)
$\text{H}_5[\text{P}^{\text{V}}\text{V}_{14}\text{O}_{42}]^{4-}$	11, 14, 1	2.1	No	-534, -589, -598	96.03 (0.17)
$\text{H}[\text{P}^{\text{V}}\text{V}_7\text{O}_7]^{3-}$	-1, 1, 1		No	-567.1	-5.68 (0.10)
$\text{H}_2[\text{P}^{\text{V}}\text{V}_7\text{O}_7]^{2-}$	0, 1, 1	7.19	No	-582.5	1.51 (0.08)
$\text{H}_3[\text{P}^{\text{V}}\text{V}_7\text{O}_7]^{-}$	1, 1, 1	3.82	No	-558.7	5.33 (0.07)
$\text{H}_4[\text{P}^{\text{V}}\text{V}_7\text{O}_7]$	2, 1, 1	3.04	No	-557.9	8.37 (0.07)
$\text{H}[\text{P}^{\text{V}}\text{V}_2\text{O}_{10}]^{4-}$	-1, 1, 2		No	-579.4	-3.9 (0.26)
$\text{H}_2[\text{P}_2^{\text{V}}\text{V}_2\text{O}_{10}]^{3-}$	0, 1, 2	6.3	No	-602.0	2.4 (0.12)

^a Stoichiometric coefficients and formation constants for: $p[\text{H}_2\text{V}^{\text{O}}_4]^- + q\text{H}^+ + r[\text{H}_2\text{PO}_4]^- \rightleftharpoons [\text{H}_z\text{P}_r\text{V}_p\text{O}_v]^{n-} + (q/2 + r + p - z/2)\text{H}_2\text{O}$. ^b $\delta^{51}\text{V}$ relative to VOCl_3 . ^c Ionic strength $\mu(\text{NaCl}) = 0.15$ M.

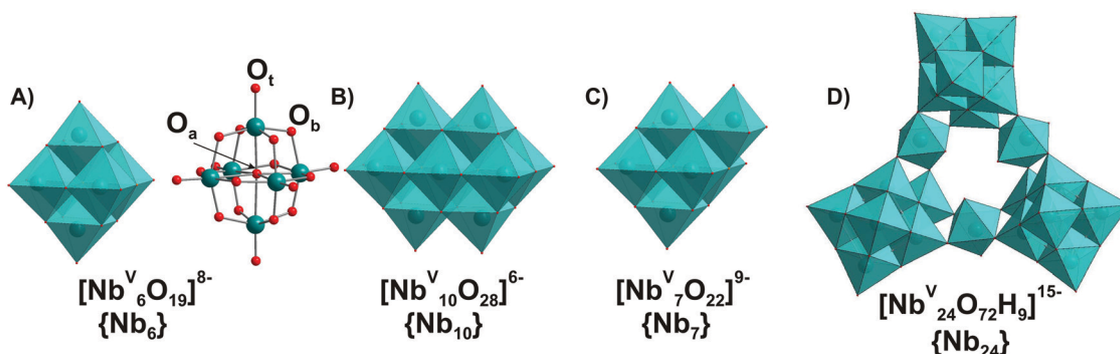


Fig. 8 IPONb species ((A)–(D)) with the nuclearity up to 24 addenda atoms, which can be present in aqueous media. The Lindqvist anion $\{\text{Nb}_6\}$ is presented in (A) as polyhedral and ball-and-stick form with the indicated types of oxygen atoms: $\mu_6\text{-O}_a$ – hexacoordinated oxygen atom; $\mu_2\text{-O}_b$ – dicoordinated bridging atom; O_t – terminal oxygen atom. Color code: $\{\text{Nb}_6\}$, blue; O, red.

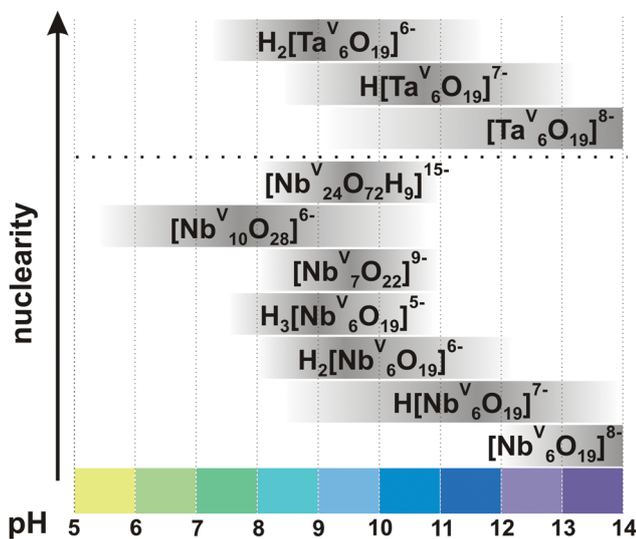


Fig. 9 Speciation of IPONbs and IPOtas in an aqueous solution based on works.^{29,113–115,118,119,126} The range of $\text{pH} < 5$ is not studied well and the precipitation of $\text{Nb}_2\text{O}_5 \cdot n\text{H}_2\text{O}$ is usually observed. The maximum intensity of grey color in each box with a single species corresponds to its maximum concentration in the chosen pH region. The grey boxes along the y-axis are positioned according to increasing nuclearity, but do not show the domination over other species at a certain pH range. The structures of IPONbs are depicted in Fig. 8 and their formation constants are given in Table 3.

3.2.1.2. *Hexaniobate and -tantalate*. The Lindqvist ion $\{\text{Nb}_6\}$ (Fig. 8A) was the first PONb studied in both the solid-state and solution. $\{\text{Nb}_6\}$ has the highest charge-density of all known POMs and readily associates with alkali metals in solutions. Nyman group studied the alkali metals ion-association of $\{\text{Nb}_6\}$ in solution *via* SAXS and showed that the completely neutralized hexaniobate, $\text{A}_6[\text{Nb}_6^{\text{V}}\text{O}_{19}]$ ($\text{A} = \text{Rb}^+, \text{Cs}^+$), is the dominant species in AOH solution, while the ‘nude’ anion $[\text{Nb}_6^{\text{V}}\text{O}_{19}]^{8-}$ dominates in TMAOH (TMA = tetramethylammonium) solutions.¹¹⁶ It is worth mentioning that the hexatantalate $[\text{Ta}_6^{\text{V}}\text{O}_{19}]^{8-}$ ($\{\text{Ta}_6\}$) exhibits lower solubility in water than the analog hexaniobate salts.¹⁰⁹ The dissolution of $\text{TBA}_6[\text{H}_2\text{Ta}_6^{\text{V}}\text{O}_{19}]$ in toluene under prolonged heating leads to the isolation of $\text{TBA}_6[\text{Ta}_{10}^{\text{V}}\text{O}_{28}] \cdot 6\text{H}_2\text{O}$, which has an isostructural anion with decavanadate $\{\text{V}_{10}\}$ (Fig. 5) and decaniobate $\{\text{Nb}_{10}\}$ (Fig. 8B).¹¹⁷

At pH higher than 12 the hexaniobate ion $\{\text{Nb}_6\}$ is deprotonated and at lower pH values (Fig. 9), it carries between one and three protons, probably located on the μ_2 -bridging O_b oxygen atoms (Fig. 8A). According to ^{17}O NMR measurements and density functional theory (DFT) calculations the protonated states of $\{\text{Ta}_6\}$ are all observed at lower pH values – mono-protonated species dominate at pH 10.5 and di-protonated at pH 9.¹¹⁸ Hexaniobate $\{\text{Nb}_6\}$ (Fig. 8A) in its various protonation states has been characterized in solution and in the solid-state (Table 3). While all hexatantalate protonation states have been



Table 3 Isopolyoniobates stoichiometry and formation constants

IPONb species	q, p^a	Isolated in solid state	Formation constants ^a $\lg K_c$ at 25 °C according to	
			Rozantsev ^{b113}	Extebarria ^{c119}
$[\text{Nb}_6\text{O}_{19}]^{8-}$ (Fig. 8A)	0, 1	Yes ¹¹⁰		
$\text{H}[\text{Nb}_6\text{O}_{19}]^{7-}$	1, 1	Yes ¹²⁰	9.44	13.63
$\text{H}_2[\text{Nb}_6\text{O}_{19}]^{6-}$	2, 1	Yes ¹²¹	15.95	23.55
$\text{H}_3[\text{Nb}_6\text{O}_{19}]^{5-}$	3, 1	Yes ¹²²	22.12	32.90
$[\text{Nb}_7\text{O}_{22}]^{9-}$ (Fig. 8B)	0.28, 1	Yes ^{123,124}		
$[\text{Nb}_{10}\text{O}_{28}]^{6-}$ (Fig. 8C)	4.4, 1	Yes ¹¹¹		
$[\text{H}_9\text{Nb}_{24}\text{O}_{72}]^{15-}$ (Fig. 8D)	4.25, 1	Yes ¹²⁵		

^a Stoichiometric coefficient and formation constants for: $p[\text{Nb}_6\text{O}_{19}]^{8-} + q\text{H}^+ \rightleftharpoons [\text{H}_q\text{Nb}_{6p}\text{O}_p]^{n-} + (q/2 + p - z/2)\text{H}_2\text{O}$. ^b Ionic strength $\mu = 0.1$ M (KCl). ^c $\mu = 3$ M (KCl).

observed in solution, only the salt in its di-protonated form was isolated in solid state.¹⁰⁹

3.2.1.3. Decaniobate transformation. Aqueous $\{\text{Nb}_{10}\}$ (Fig. 8B) is stable and soluble at neutral pH, converts to $\{\text{Nb}_6\}$ with increased pH (pH > 11) and precipitates as Nb_2O_5 by acidification (pH < 5.5) (Fig. 9).²⁹ The decaniobate ion showed no sign of protonation between pH 6–10, in contrast to the hexaniobate ion which is protonated between pH 8 and 13.¹¹⁵ The conversion of $\{\text{Nb}_{10}\}$ to $\{\text{Nb}_6\}$ was studied by ¹⁷O NMR, ESI-MS and DFT calculations and proceeds *via* a “heptaniobate” intermediate, $\{\text{Nb}_7\}$ (Fig. 8C).^{114,115} $\{\text{Nb}_7\}$ features three reactive terminal oxo-ligands leading to condensation into larger clusters such as the $\{\text{Nb}_{24}\}$ unit.¹²⁵ Recently, Nyman and co-authors showed *via* Raman spectroscopy, ESI-MS, X-ray scattering and computational

studies that $\{\text{Nb}_{10}\}$ converts to oligomers of $[\text{H}_x\text{Nb}_{24}\text{O}_{72}]^{(24-x)-}$ ($\{\text{Nb}_{24}\}$, Fig. 8D) upon adding only alkali chloride salts, even in buffered (1 M HEPES, pH = 7) neutral solutions.¹²⁷ The rate of $\{\text{Nb}_{10}\}$ to $\{\text{Nb}_{24}\}$ conversion increases in raw $\text{Cs}^+ > \text{Rb}^+ > \text{K}^+ > \text{Na}^+ > \text{Li}^+$ and cation concentration and indicates that the alkali cations open the compact $\{\text{Nb}_{10}\}$ structure and are primarily responsible for driving the reaction.¹²⁷

3.3. Polyoxomolybdates (POMs)

Since the 1960s the speciation in acidified solutions of $[\text{Mo}^{\text{VI}}\text{O}_4]^{2-}$ with different concentration and ionic strength has been a subject of comprehensive investigation by potentiometry, NMR spectroscopy, and ESI-MS and was extensively elaborated by Tytko and Glemser,¹²⁸ Cruywagen,⁶ Pettersson,¹²⁹ Maksimovskaya¹⁸ and others. However, in the last decades, this important topic receives less and less attention and for newly synthesized POMs their solution stability and transformation are rarely investigated.

3.3.1. Isopolymolybdates (IPOMs)

3.3.1.1. Ion-distribution diagram for IPOMs. The polymerization processes of $[\text{Mo}^{\text{VI}}\text{O}_4]^{2-}$ are observed in solution where the Mo^{VI} concentration is higher than 10^{-4} M. In strongly alkaline molybdate(vi) solutions the monomeric $[\text{Mo}^{\text{VI}}\text{O}_4]^{2-}$ anion is dominating and forms salts with mono- to trivalent cations, whereas in strongly acidic solutions, the predominant species are stable mono- $[\text{Mo}^{\text{VI}}\text{O}_2(\text{OH})(\text{H}_2\text{O})_3]^+$ or bi- $[\text{Mo}_2^{\text{VI}}\text{O}_5(\text{OH})(\text{H}_2\text{O})_5]^+$ oxocations, which form salts with inorganic anions and complexes with organic ligands.¹⁸ It is generally accepted that the first major product that forms upon acidification of aqueous $[\text{Mo}^{\text{VI}}\text{O}_4]^{2-}$ is heptamolybdate, $[\text{Mo}_7^{\text{VI}}\text{O}_{24}]^{6-}$ ($\{\text{Mo}_7\}$, Fig. 10A), which shows its maximum concentration at pH ~ 5 and can

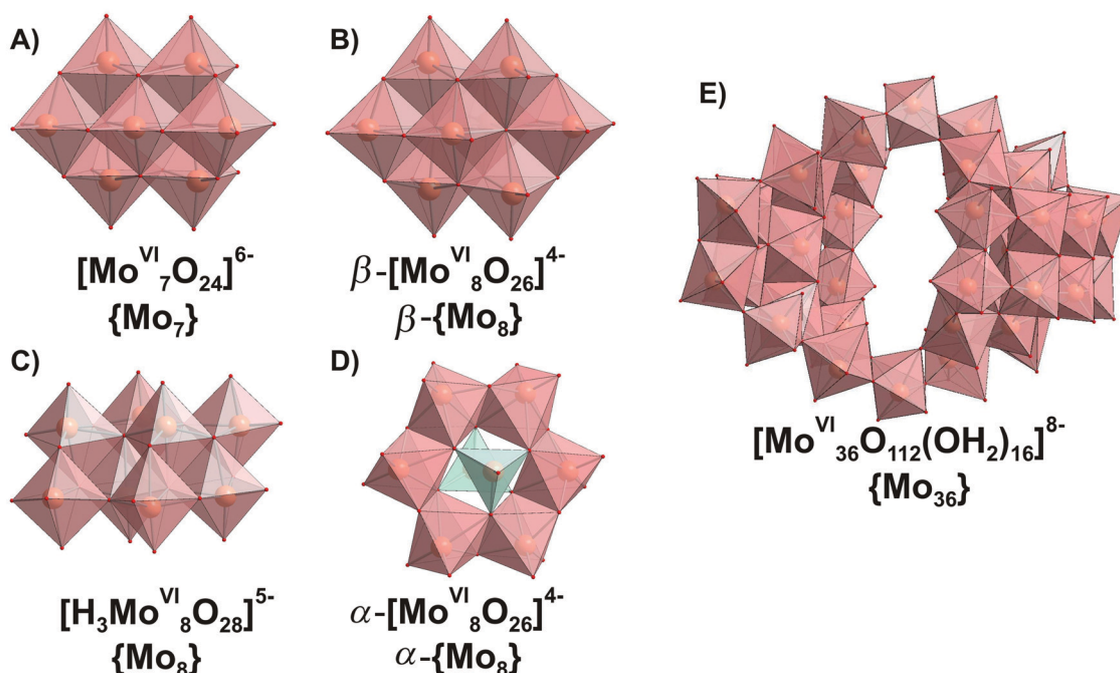


Fig. 10 IPOMs with nuclearity of up to 36 addenda atoms, which are confirmed to be present in aqueous media. Color code: $\{\text{MoO}_6\}$, pink; $\{\text{MoO}_4\}$, green; O, red.



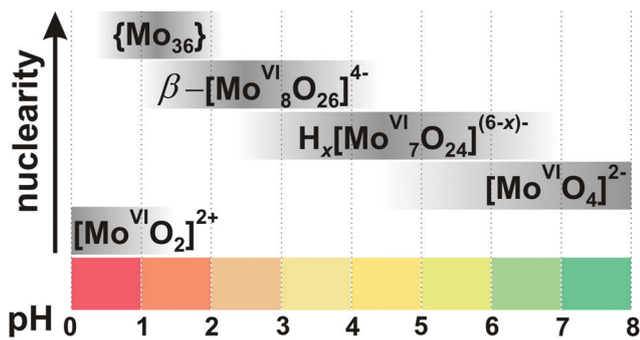


Fig. 11 Speciation of IPOMos in an aqueous solution with the concentration of Mo^{VI} 0.1–0.4 M based on works.^{6,18,128,131,132} The maximum intensity of grey color in each box with a single species corresponds to its maximum concentration in the chosen pH region. The grey boxes along the y-axis are positioned according to increasing nuclearity, but do not show the domination over other species at a certain pH range. x in $\text{H}_x[\text{Mo}^{\text{VI}}\text{O}_{24}]^{(6-x)-}$ is 0–2; $\{\text{Mo}_{36}\}$ is abbreviation for $[\text{Mo}_{36}^{\text{VI}}\text{O}_{112}(\text{OH})_{216}]^{8-}$. The structures of species are presented in Fig. 10. The ^{95}Mo and ^{17}O chemical shifts, stoichiometry, formation constants and pK_a values are summarized in Table 4.

be protonated upon further acidification (Fig. 11). The $\{\text{Mo}_7\}$ ion, its protonated forms and the octamolybdate ion $\beta\text{-}[\text{Mo}_8^{\text{VI}}\text{O}_{26}]^{4-}$ ($\beta\text{-}\{\text{Mo}_8\}$, Fig. 10B) are included in most speciation models describing potentiometric results (Table 4). In some cases, $\text{H}_3[\text{Mo}_7^{\text{VI}}\text{O}_{24}]^{3-}$ is preferred over $\beta\text{-}\{\text{Mo}_8\}$ as the only existing species, but in the majority of models both polyanions are present. There are contradictory opinions about the speciation being present in the pH region between 4.5 and 2.5.

In 1990, Howarth and co-authors¹²⁹ proposed, based on ^{95}Mo and ^{17}O NMR spectroscopic studies, that monoprotonated $\{\text{Mo}_7\}$, instead of being higher protonated, is converted to the intermediate polyanion $[\text{H}_3\text{Mo}_8^{\text{VI}}\text{O}_{28}]^{5-}$ (Fig. 10C). The structure of the latter species is believed to be the same as that of the $[\text{H}_2\text{Mo}_8^{\text{VI}}\text{O}_{28}]^{6-}$ anion, which was crystallized from aqueous solution with isopropylammonium as counteranion.¹³⁰ Later Maksimovskaya and Maksimov have shown using the same methods, ^{95}Mo and ^{17}O NMR, that only heptamolybdate and octamolybdate co-exist in the pH region from 5.5 to 2.5.¹⁸ Moreover, they showed evidence of the di-protonated anion $\{\text{Mo}_7\}$ and that chemical exchange occurs between $\text{H}_x[\text{Mo}_7^{\text{VI}}\text{O}_{24}]^{(6-x)-}$ ($x = 0\text{--}2$) and $\beta\text{-}\{\text{Mo}_8\}$ without any additional intermediate species. Notably, a recent potentiometric study confirms the presence of both octamolybdate and the up to three protonated forms of $\{\text{Mo}_7\}$ in solution ($C_{\text{Mo}} = 0.15$ M, ionic strength $\mu = 0.15$ M (NaClO_4)).¹³¹

Salts of the larger anion, $[\text{Mo}_{36}^{\text{VI}}\text{O}_{112}(\text{OH})_{216}]^{8-}$ (Fig. 10E and Table 4) can be crystallized from systems at pH lower than 2.8¹⁴¹ and its identity in solution was confirmed by ^{95}Mo NMR^{18,132} and Raman¹³³ spectroscopy. The anion is a dimer of $[\text{Mo}_{18}^{\text{VI}}\text{O}_{56}(\text{H}_2\text{O})_8]^{4-}$ with inversion center, that is why both $\{\text{Mo}_{36}\}$ and $\{\text{Mo}_{18}\}$ are often included in the ion-distribution diagram⁶ for Mo^{VI} solution in acidic region (Table 4).

Hypothetical anions such as $[\text{HMo}_{17}^{\text{VI}}\text{O}_{55}]^{7-}$, $[\text{HMo}_{13}^{\text{VI}}\text{O}_{42}]^{5-}$ and $[\text{Mo}_{10}^{\text{VI}}\text{O}_{34}]^{8-}$ have been included to theoretical models for better interpretation of potentiometric results obtained at high

molybdate concentration (>0.1 M),^{6,134} but more work is necessary to determine the existence and structure of these species in solution. At lower molybdate concentrations, the presence of the hypothetical species $[\text{H}_2\text{Mo}_6^{\text{VI}}\text{O}_{21}]^{4-}$ has been proposed.⁶ Equilibrium constants obtained by different research groups based on potentiometric and NMR spectroscopic studies under the same conditions agree quite well (Table 4). The ion-distribution diagram presented here (Fig. 11) is based on potentiometric, Raman and NMR spectroscopic investigations (Table 4).

3.3.1.2. IPOMos under physiological conditions. Two stable species, $[\text{Mo}^{\text{VI}}\text{O}_4]^{2-}$ and $\{\text{Mo}_7\}$ (Fig. 10A), have been reported to be simultaneously present at neutral pH, that makes the analysis of their equilibrium important for understanding their function in biological or catalytic application complicated. Ng *et al.* investigated isopolymolybdate solutions as a function of concentration, time and pH by Raman spectroscopy, light and X-ray scattering,¹⁴³ showing that significant amounts of both $[\text{Mo}^{\text{VI}}\text{O}_4]^{2-}$ and $\{\text{Mo}_7\}$ are present in equilibrium over a wide pH range (4.5–7) in a 0.1 M solution of Mo^{VI} . Quantitative deconvolution of the Raman spectra taken from 0.1 M solution of Mo^{VI} demonstrated that even at pH 6.6, only half of all orthomolybdate is in hepta-form and the full conversion to $\{\text{Mo}_7\}$ is completed only at pH 4.5. Ng *et al.*¹⁴³ did not observe the formation of significant quantities of $\beta\text{-}\{\text{Mo}_8\}$ (Fig. 10B), as was suggested by Tytko.¹³⁵ When studying the sorption of isopolyoxomolybdates into layered double hydroxides by *in situ* real time infrared spectroscopy, the coexistence of $[\text{Mo}^{\text{VI}}\text{O}_4]^{2-}$ and $\{\text{Mo}_7\}$ was observed only in the pH range from 5.5 to 5.7 in 0.1 M solution of Mo^{VI} , whereas at pH above 5.7, the predominant anion is the monomeric one.¹⁴⁴

Investigation of the hydrolysis of the phosphodiester bond in the DNA model substrate bis(*p*-nitrophenyl)phosphate (BNPP, 25 mM) in the presence of $\text{Na}_6[\text{Mo}_7^{\text{VI}}\text{O}_{24}]$ (25 mM) at 50 °C in the absence of buffer showed that the maximal cleavage reaction was observed at pH = 5.3.¹⁴⁵ ^{95}Mo NMR confirmed that the predominant species at this pH is $\{\text{Mo}_7\}$ (Fig. 10A), which promotes the hydrolysis. Furthermore, ^{95}Mo NMR and Mo K-edge EXAFS investigations of the different hydrolytic reaction stages showed a gradual disappearance of $\{\text{Mo}_7\}$ during the hydrolytic reaction and appearance of $[\text{P}_2\text{Mo}_5^{\text{VI}}\text{O}_{23}]^{6-}$ (Fig. 14A), which was the final compound observed at the end of hydrolytic reaction.¹⁴⁶

3.3.1.3. Hexamolybdate $[\text{Mo}_6^{\text{VI}}\text{O}_{19}]^{2-}$ and its stability in aqueous solutions. In addition to the five species shown in the ion-distribution diagram for IPOMos (Fig. 11), the Lindqvist type hexamolybdate anion $[\text{Mo}_6^{\text{VI}}\text{O}_{19}]^{2-}$ ($\{\text{Mo}_6\}$) exists in solid state and has the same structure as its niobate analog (Fig. 8A). The $\{\text{Mo}_6\}$ anion is the only colored IPOMO (yellow, $\lambda_{\text{max}} = 325$ nm, $\text{O}_b \rightarrow \text{Mo}^{\text{VI}}$ charge transfer) structurally based on six monounits MoO_t (O_t – terminal oxygen atom), rather than three *cis* MoO_2 units. $\{\text{Mo}_6\}$ does not exist in any notable concentration in pure aqueous solution and requires solvents with lower dielectric constants¹⁴⁷ or macrocycles¹³² for stabilization due



Table 4 IPOMos stoichiometry, formation constants, chemical shifts and pK_a values

IPOMo species	<i>p</i> , <i>q</i> ^a	pK _a	Isolated in solid state	δ(⁹⁵ Mo), ppm ^b	δ(¹⁷ O), ppm ^b	Raman bands, cm ⁻¹	Formation constants ^a lg K _c at 25 °C			
							Cruywagen ^{c6}	Tytko ^{d135}	Taube ^{e136}	Cruywagen ^{f137}
[Mo ^{VI} O ₄] ²⁻	0, 1		Yes			896 ¹³³				
H[Mo ^{VI} O ₄] ⁻	1, 1		No					3.48	3.40	
[Mo ^{VI} O ₂₄] ⁶⁻ (Fig. 10A)	8, 7		Yes ¹³⁸	210, 32, 15 ¹⁸	121.7, 338.6, 356.5, 395.2, 745, 754.4, 815 ¹²⁹	939 ¹³³	54.07 (0.03)	55.71	52.43	
H[Mo ^{VI} O ₂₄] ⁵⁻	9, 7	4.38	No				59.01 (0.03)	60.22	57.42	
H ₂ [Mo ^{VI} O ₂₄] ⁴⁻	10, 7	3.38 ¹³⁵	No						61.24	
H ₃ [Mo ^{VI} O ₂₄] ³⁻	11, 7	1.87 ¹³⁵	No						63.90	
[H ₃ Mo ^{VI} O ₂₈] ⁵⁻ (Fig. 10C)	11, 8		Yes ¹³⁰		261, 370, 395, 840, 850, 860 ¹²⁹		70.30 (0.03)	71.41		
α-[Mo ^{VI} O ₂₆] ⁴⁻ (Fig. 10D)	12, 8		Yes ¹³⁹			959, 918 ¹³³				
β-[Mo ^{VI} O ₂₆] ⁴⁻ (Fig. 10B)	12, 8		Yes ¹⁴⁰	100, 10 ¹⁸	53.5, 290.7, 404.3, 738.2, 861.6, 876, 878, 917.3 ¹²⁹	971, 943, 915, 904 ¹³³	74.10 (0.03)	71.52		
β-H[Mo ^{VI} O ₂₆] ³⁻	13, 8	1.83 ¹³⁵	No							
[Mo ^{VI} O ₅₆ (H ₂ O) ₈] ⁴⁻	32, 18		half unit of {Mo ₃₆ } ¹⁴¹				179.28	180.39		
[Mo ^{VI} O ₁₁₂ (OH) ₁₆] ⁸⁻ (Fig. 10E)	64, 36		Yes ¹⁴¹			983, 957, 899 ¹³³				
[Mo ^{VI} O ₂] ²⁺	4, 1		Yes ¹⁴²	-62 to -69 ¹⁸						
[Mo ^{VI} O ₂ (OH)(H ₂ O) ₃] ⁺	3, 1	0.9	No					8.09	8.81	
[Mo ^{VI} O ₅ (OH)(H ₂ O) ₅] ⁺	5, 2	~0.1-0.2	No					18.02	18.93	
[Mo ^{VI} O ₅ (H ₂ O) ₆] ⁺										19.60
[Mo ^{VI} O ₄ (OH)(H ₂ O) ₅] ⁺										18.98

^a Stoichiometric coefficients and formation constants for $p[\text{Mo}^{\text{VI}}\text{O}_4]^{2-} + q\text{H}^+ \rightleftharpoons [\text{H}_z\text{Mo}_p^{\text{VI}}\text{O}_m]^{n-} + (q - z/2)\text{H}_2\text{O}$ relative to Na_2MoO_4 . ^b $\delta^{17}\text{O}$ relative to H_2O . ^c Ionic strength $\mu = 2 \text{ M}$ (LiClO_4). ^d $\mu = 2 \text{ M}$ (NaCl). ^e $\mu = 0.3 \text{ M}$ (Na_2SO_4). ^f $\mu = 3 \text{ M}$ ($\text{Na}(\text{H})\text{ClO}_4$).

to its low charge density and rigid symmetry. UV-vis spectroscopy shows that $[\text{Mo}_6^{\text{VI}}\text{O}_{19}]^{2-}$ is formed in solution containing acetone, acetonitrile, ethanol, dimethyl formamide and 1,4-dioxane with an organic solvent content >30%.¹⁴⁷ The addition of a large γ -cyclodextrin macrocycle to the acidified aqueous solution of sodium orthomolybdate leads to the host-guest stabilization of $\{\text{Mo}_6\}$ by formation of the complex $\{\text{Mo}_6\text{O}_{19}@\gamma\text{-CD}\}^{2-}$, as a new species in the pH range between 1 and 2.5.¹³² Still the inclusion of the Lindqvist type hexamolybdate in the IPOMos distribution diagram in purely aqueous solutions is reported which should be seen critically.¹⁴⁸

3.3.1.4. Mono- and bimolybdyl cationic species in strongly acidic solutions. The successive protonation equilibria of $[\text{Mo}^{\text{VI}}\text{O}_4]^{2-}$ lead to formation of the cationic species: $[\text{Mo}^{\text{VI}}\text{O}_2(\text{OH})(\text{H}_2\text{O})_3]^+$, $[\text{Mo}_2^{\text{VI}}\text{O}_5(\text{H}_2\text{O})_6]^{2+}$, $[\text{Mo}_2^{\text{VI}}\text{O}_4(\text{OH})(\text{H}_2\text{O})_6]^{3+}$, that are well characterized by potentiometry, and the equilibrium constants have been determined for different ionic media by various authors (Table 4).^{137,149,150} The total molybdate concentration must be lower than 10^{-4} M for the mono- and binuclear species to be predominantly present in aqueous solution. Cruywagen has reported the presence of four species, $[\text{Mo}^{\text{VI}}\text{O}_3(\text{H}_2\text{O})_3]$, $[\text{Mo}^{\text{VI}}\text{O}_2(\text{OH})(\text{H}_2\text{O})_3]^+$, $[\text{Mo}_2^{\text{VI}}\text{O}_5(\text{H}_2\text{O})_6]^{2+}$, $[\text{Mo}_2^{\text{VI}}\text{O}_4(\text{OH})(\text{H}_2\text{O})_6]^{3+}$,

under strongly acidic conditions (HClO_4 with concentration 0.5–4 M and pH region from -3 to 0.5).¹³⁷ Later, applying XAS, UV-vis spectroscopy in combination with first-principles calculation methods, all four species were confirmed and the predominant species in 2 M nitric acid are $[\text{Mo}_2^{\text{VI}}\text{O}_5(\text{H}_2\text{O})_6]^{2+}$ and $[\text{Mo}_2^{\text{VI}}\text{O}_4(\text{OH})(\text{H}_2\text{O})_6]^{3+}$ (Fig. 12).¹⁴⁹

3.3.1.5. Computational investigation of IPOMos formation. The speciation diagram for IPOMos does not show species with two to six molybdenum ions (Fig. 11). It is possible that these species exist at low concentration as intermediates in the process of forming larger IPOMos. DFT calculations together with ESI-MS experiments provide^{151,152} insights into the possible formation mechanism of the Lindqvist $[\text{Mo}_6^{\text{VI}}\text{O}_{19}]^{2-}$ ion (Fig. 8A), which is preferably formed in organic media and is undergoing a transformation in aqueous solution.¹ Unstable low-nuclearity species such as $[\text{Mo}_2^{\text{VI}}\text{O}_7]^{2-}$, $[\text{Mo}_3^{\text{VI}}\text{O}_{10}]^{2-}$, $[\text{Mo}_4^{\text{VI}}\text{O}_{13}]^{2-}$, $[\text{Mo}_5^{\text{VI}}\text{O}_{16}]^{2-}$ and $[\text{Mo}_6^{\text{VI}}\text{O}_{19}]^{2-}$ were detected experimentally by ESI-MS in acetonitrile. Recently, Duarte and co-authors have shown using DFT/PBE calculations that formation of IPOMos with up to four molybdenum atoms is highly favorable in water.¹⁵³ The formation of the predominant species $[\text{Mo}_7^{\text{VI}}\text{O}_{24}]^{6-}$ and $\beta\text{-}[\text{Mo}_8^{\text{VI}}\text{O}_{26}]^{4-}$ involves a reaction between



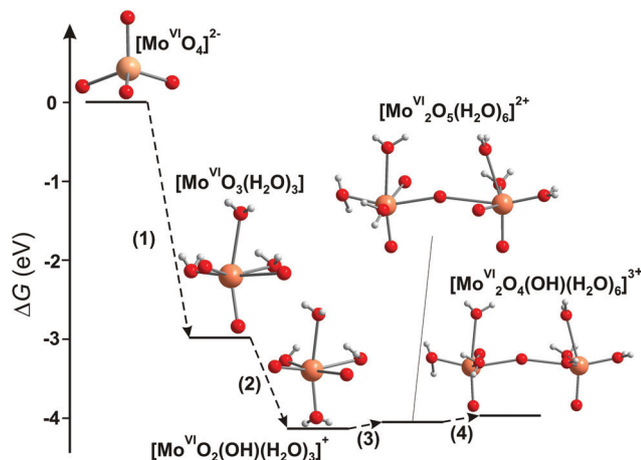


Fig. 12 Energy diagram of the free-energy change (ΔG) for the condensation reaction of Mo complexes.¹⁴⁹ Color code: Mo, orange; O, red; H, grey.

the species with six and seven Mo atoms, respectively, and the monomer $[\text{Mo}^{\text{VI}}\text{O}_4]^{2-}$, which are frequently assisted by protons through an aggregation or water condensation mechanism.¹⁵³

3.3.1.6. Comparison of IPOMos in solution and the solid state.

As with many other POMs there are some isopolymolybdates, which occur only in solution, and others only in solid state. However, all prevalent molybdate species ($[\text{Mo}^{\text{VI}}\text{O}_4]^{2-}$, $[\text{Mo}_7^{\text{VI}}\text{O}_{24}]^{6-}$ (Fig. 10A), $\beta\text{-}[\text{Mo}_8^{\text{VI}}\text{O}_{26}]^{4-}$ (Fig. 10B), $[\text{Mo}_{36}^{\text{VI}}\text{O}_{112}(\text{OH}_2)_{16}]^{8-}$ (Fig. 10E)) exist in solution and in solid state (Table 4). Low-nuclearity IPOMos, which are unstable in solution, tend to form polymeric structures with infinite two-dimensional chains that have been synthesized in solid state reactions^{154,155} or hydrothermally (Table 5).^{156,157} *In situ* Raman spectroscopy confirmed that between 170 and 190 °C and

pH 7 and 5, chain-like or discrete molecular structures of dimolybdates $[\text{Mo}_2^{\text{VI}}\text{O}_7]^{2-}$ (Fig. 13B) and trimolybdates $[\text{Mo}_3^{\text{VI}}\text{O}_{10}]^{2-}$ (Fig. 13D) are preferentially formed,¹⁵⁸ whereas heptamolybdate $\{\text{Mo}_7\}$ dominates under ambient pressure at 25 °C (Fig. 11). Long-term equilibrium in solutions of $\{\text{Mo}_7\}$ leads to its transformation and crystallization of different products depending on temperature and time (Fig. 13 and Table 5), which should undoubtedly be taken into account when using heptamolybdate. The addition of structure-directing reagents such as big cage-like organic cations (1,4-diazabicyclo[2.2.2]octane¹⁵⁹ or bis[benzidinium(1-)]¹⁵⁷) affects the speciation at a given pH value and leads to the isolation of new, sometimes polymerized, structures (Table 5).

3.3.2. Heteropolymolybdates (HPOMos). Molybdenum(vi) forms a great number of heteropolyanions, some of which have been recognized for well over 150 years and are still widely used in the quantitative and qualitative element's determination, especially for phosphorus and silicon. The universality of molybdenum in the formation of various heteropolyanionic structures with several metallic and non-metallic heteroatoms competes only with tungsten. The molybdophosphate $[\text{P}^{\text{V}}\text{Mo}_{12}^{\text{VI}}\text{O}_{40}]^{3-}$ ($\{\text{PMo}_{12}\}$) (Fig. 14C), and the analogous molybdosilicate, $[\text{Si}^{\text{IV}}\text{Mo}_{12}^{\text{VI}}\text{O}_{40}]^{4-}$ ($\{\text{SiMo}_{12}\}$) have always been the most widely studied of all heteropolymolybdates. This section is divided into parts according to the type of heteroatom and ends with sections on the stability of the two most common archetypes – Keggin (Fig. 14C) and Wells–Dawson (Fig. 14G).

3.3.2.1. Speciation in the phosphate-molybdate system. In the late 1980s Pettersson *et al.* extensively analyzed speciation in the system $[\text{Mo}^{\text{VI}}\text{O}_4]^{2-}\text{-HP}^{\text{V}}\text{O}_4^{2-}\text{-H}^+$ using potentiometry and ³¹P NMR spectroscopy in the pH range from 0.5 to 7 and the concentration range $[\text{Mo}] = 0.030\text{--}0.480\text{ M}$, $[\text{P}] = 0.005\text{--}0.040\text{ M}$, where $[\text{Mo}]$ and $[\text{P}]$ – are the total concentrations of

Table 5 IPOMos isolated from aqueous solutions at different pH values

pH or Z ^a	Species isolated from solution	Cation	Ref.
> 7	$[\text{Mo}^{\text{VI}}\text{O}_4]^{2-}$	Metal ions in ox. state 2+, 3+	160
5–7	$[\text{Mo}_7^{\text{VI}}\text{O}_{24}]^{6-}$ (Fig. 10A)	Na^+ , K^+ , Cs^+ , NR_4^+ (R = H, alkyls, alkenyls, phenyls)	161
6–7 (from $[\text{Mo}_7^{\text{VI}}\text{O}_{24}]^{6-}$ solution at 60 °C after addition of NH_4^+)	$([\text{Mo}_2^{\text{VI}}\text{O}_7]^{2-})_\infty$ (Fig. 13B)	NH_4^+	4
6–7 (aging of $[\text{Mo}_7^{\text{VI}}\text{O}_{24}]^{6-}$ solution – 12 days)	$([\text{Mo}_3^{\text{VI}}\text{O}_{10}]^{2-})_\infty$ (Fig. 13D)	Rb^+	5
6–7 (aging of $[\text{Mo}_7^{\text{VI}}\text{O}_{24}]^{6-}$ solution – weeks or HT ^b)	$([\text{Mo}_8^{\text{VI}}\text{O}_{27}]^{4-})_\infty$ (Fig. 13C)	NH_4^+ , $(\text{C}_4\text{H}_{12}\text{N}_2)^+$, $(\text{NH}_3(\text{CH}_2)_3\text{NH}_3)^{2+}$, <i>etc.</i>	162–164
6–7 (aging of $[\text{Mo}_7^{\text{VI}}\text{O}_{24}]^{6-}$ solution – under UV irradiation)	$[\text{H}_2\text{Mo}_8^{\text{VI}}\text{O}_{28}]^{6-}$ (Fig. 13A)	$(\text{C}_3\text{H}_{10}\text{N})^+$	130
6–7 (aging of $[\text{Mo}_7^{\text{VI}}\text{O}_{24}]^{6-}$ solution – several days at 80 °C)	$[\text{Mo}_{10}^{\text{VI}}\text{O}_{34}]^{8-}$ (constructed from $\{\text{Mo}_8^{\text{VI}}\text{O}_{28}\}$ unit connected at corners to two $\{\text{Mo}^{\text{VI}}\text{O}_4\}$ tetrahedra) (Fig. 13E)	NH_4^+	165
5 (HT ^b from $[\text{Mo}_7^{\text{VI}}\text{O}_{24}]^{6-}$)	$[\text{Mo}_5^{\text{VI}}\text{O}_{16}]^{2-}$ (constructed from $([\text{Mo}_4^{\text{VI}}\text{O}_{14}]^{2-})_n$ chains linked through MoO_6)	$(\text{C}_{12}\text{H}_{13}\text{N}_2)^+$	157
4 (HT ^b from $[\text{Mo}_7^{\text{VI}}\text{O}_{24}]^{6-}$)	$([\text{Mo}_3^{\text{VI}}\text{O}_{10}]^{2-})_\infty$	$(\text{C}_6\text{H}_{13}\text{N}_2)^+$	
3	$[\text{Mo}_7^{\text{VI}}\text{O}_{24}]^{6-}$ (Fig. 10A)	Zn^{2+} , protonated hexamethylenetetramine	166
2–3 (HT from $[\text{Mo}_7^{\text{VI}}\text{O}_{24}]^{6-}$)	$([\text{Mo}_8^{\text{VI}}\text{O}_{26}]^{4-})_\infty$ or $([\text{Mo}_4^{\text{VI}}\text{O}_{13}]^{2-})_\infty$ (constructed from $\gamma\text{-}[\text{Mo}_8^{\text{VI}}\text{O}_{28}]$)	$(\text{C}_6\text{H}_{13}\text{N}_2)^+$, NR_4^+ (R = H, alkyls, alkenyls, phenyls)	159
Z = 1.8–2.0	$[\text{Mo}_{36}^{\text{VI}}\text{O}_{112}(\text{OH}_2)_{16}]^{8-}$ (Fig. 10E)	K^+	141

^a Z = $n(\text{H}^+)/n(\text{MoO}_4^{2-})$. ^b HT – hydrothermal synthesis.



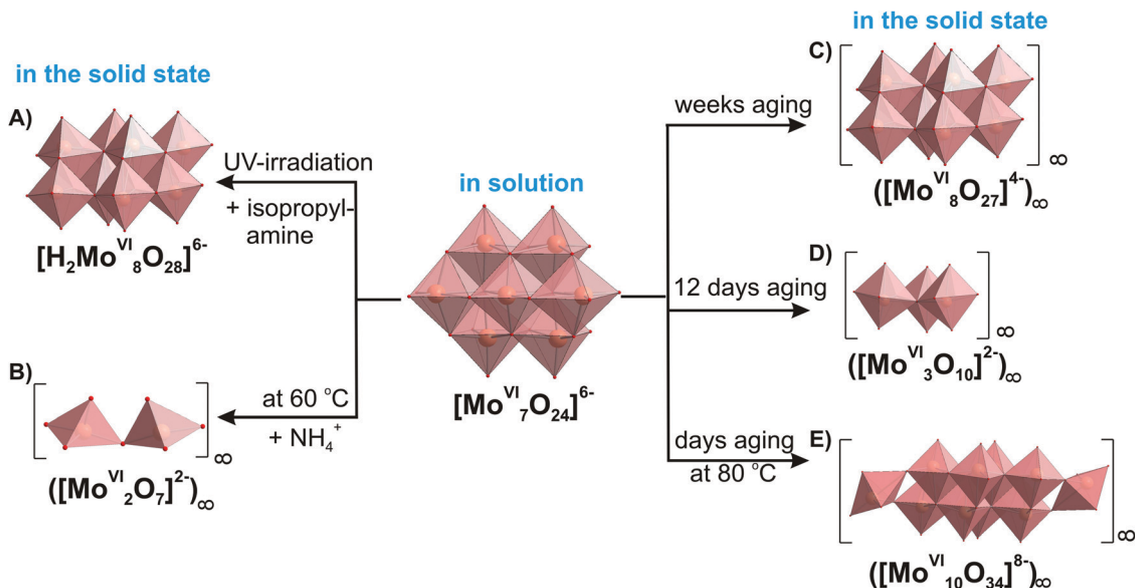


Fig. 13 Heptamolybdate (Mo_7) transformation at different synthetic conditions showing the conformity of IPOMos in solution and in the solid state based on ref. 4, 5, 130, 141, 162, 165 and 166. The time of aging for the polymeric octa- and decamolybdate has been described as "several" days or weeks in the original works^{162,165} without exact number. Color code: $\{MoO_6\}$, pink; $\{MoO_4\}$, green; O, red.

molybdenum(vi) and phosphorus(v).^{167,168} When the ratio $[Mo]/[P]$ is lower than 2.5 and the pH between 2 and 7, only colorless heteropolyanions $H_x[P_2Mo_5^VI O_{23}]^{(6-x)-}$ ($x = 0-2$, $\{P_2Mo_5\}$, Fig. 14A) are present. The $\{P_2Mo_5\}$ unit, co-called Strandberg $\{X_2Mo_5\}$ archetype, is built up from five $\{MoO_6\}$ octahedra and two $\{PO_4\}$ tetrahedra, as shown in Fig. 14A, and was isolated in solid state in all protonation states (Table 6).¹⁶⁹⁻¹⁷¹ At $[Mo]/[P] > 2.5$ and pH below 5.5, other molybdophosphate complexes with 9 $[P^VMo_9^VI O_{34}]^{9-}$, $\{PMo_9\}$, Fig. 14E and F), 11 $[P^VMo_{11}^VI O_{39}]^{7-}$,

$\{PMo_{11}\}$, Fig. 14D), 12 $[P^VMo_{12}^VI O_{40}]^{3-}$, $\{PMo_{12}\}$, Fig. 14C) and 18 $[P^VMo_{18}^VI O_{62}]^{6-}$, $\{P_2Mo_{18}\}$, Fig. 14G) Mo^{VI} atoms start to form. Besides, the $\{P_2Mo_5\}$ (Fig. 14A) and $\{PMo_{11}\}$ (Fig. 14D) series, protonated trilacunary species $\{PMo_9\}$ (Fig. 14E and F) are also formed in solution. In the acidic region with $pH < 1.8$ and at high $[Mo]/[P]$ ratios, the Keggin $\{PMo_{12}\}$ (Fig. 14C) anion is present. During aging in solution $[Mo^VI O_4]^{2-} - HP^VO_4^{2-} - H^+$ at pH lower than 3, the yellow Wells-Dawson $\{P_2Mo_{18}\}$ anion (Fig. 14G) is slowly - up to 1 month - formed. The Wells-Dawson formation is

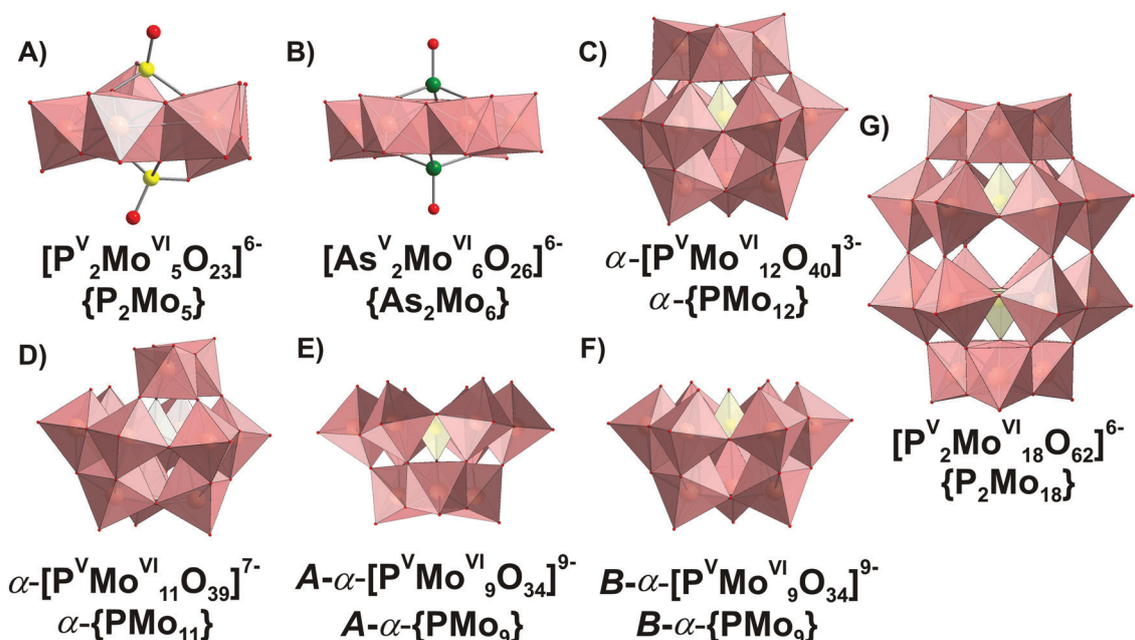


Fig. 14 Phospho- and arsenatomolybdates present in aqueous media. The data about structural isomerism of lacunary Keggin anions are given in ref. 1. Color code: $\{MoO_6\}$, pink; $\{PO_4\}$ and P, yellow; As, green; O, red.



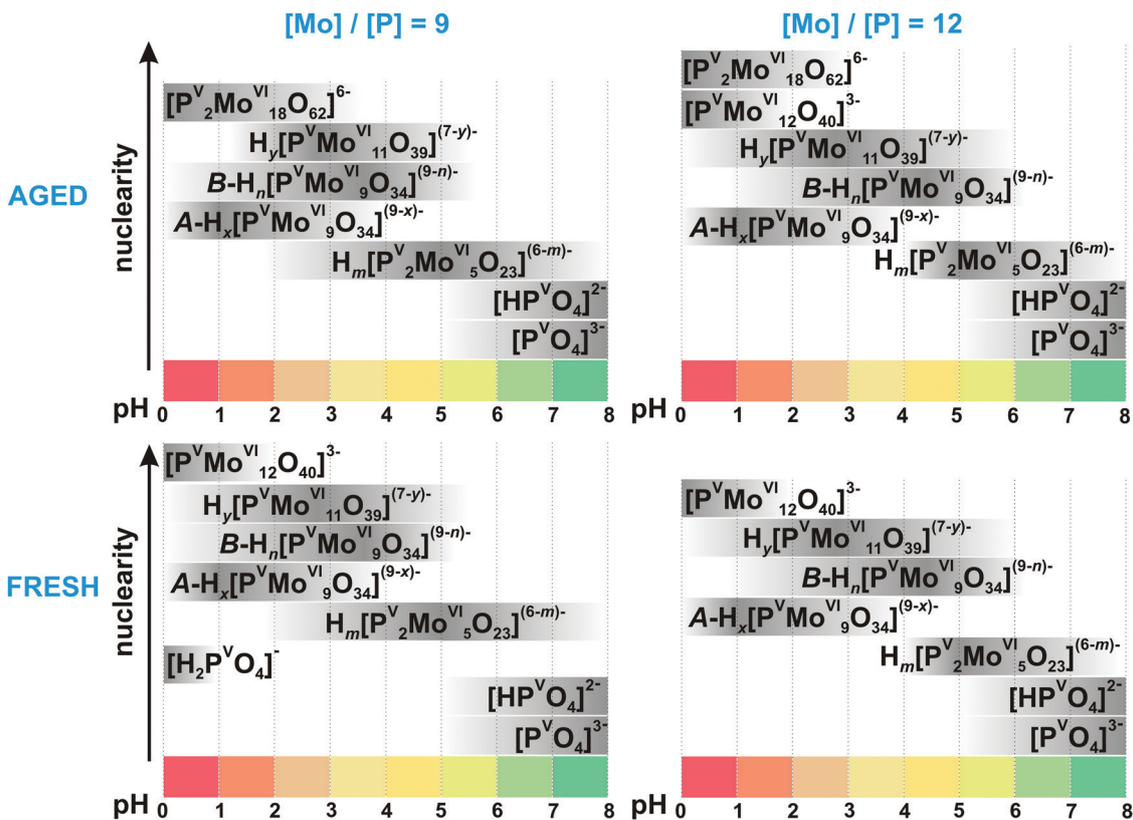


Fig. 15 Speciation of phosphomolybdates in freshly prepared and aged (up to 1 month) aqueous solution with the concentration of 0.18 M Mo^{VI} ($[\text{Mo}]/[\text{P}] = 9$) and 0.24 M ($[\text{Mo}]/[\text{P}] = 12$) based on works.^{167,168} The maximum intensity of grey color in each box with a single species corresponds to its maximum concentration in the chosen pH region. The grey boxes along the y-axis are positioned according to increasing nuclearity, but do not show the domination over other species at a certain pH range. The x value in $\text{A-H}_x[\text{P}^{\text{V}}\text{Mo}_9^{\text{VI}}\text{O}_{34}]^{(9-x)-}$ is 5–6; y in $\text{H}_y[\text{P}^{\text{V}}\text{Mo}_{11}^{\text{VI}}\text{O}_{39}]^{(7-y)-}$ is 0–2; n in $\text{B-H}_n[\text{P}^{\text{V}}\text{Mo}_9^{\text{VI}}\text{O}_{34}]^{(9-n)-}$ is 0–3; m in $\text{H}_m[\text{P}_2^{\text{V}}\text{Mo}_5^{\text{VI}}\text{O}_{23}]^{(6-m)-}$ is 0–2. The structures of species are presented in Fig. 14. The ^{31}P chemical shifts, stoichiometry, formation constants and pK_a values are summarized in Table 6.

slowed down with increasing $[\text{Mo}]$ to $[\text{P}]$ ratio. The distribution diagrams for “fresh” (after preparation) and “aged” (up to 1 month) solutions are illustrated in Fig. 15. Fig. 15 clearly demonstrates that the initially high concentration of the Keggin complex $[\text{P}^{\text{V}}\text{Mo}_{12}^{\text{VI}}\text{O}_{40}]^{3-}$ (Fig. 14C) in fresh acidic solutions with $[\text{Mo}]/[\text{P}] = 12$ diminishes considerably when the Wells-Dawson anion $[\text{P}_2^{\text{V}}\text{Mo}_{18}^{\text{VI}}\text{O}_{62}]^{6-}$ (Fig. 14G) is formed.¹⁶⁸

Mono- and trilacunary Keggin species. Raman spectroscopic studies performed by Ueda *et al.* confirmed, that the formation of Wells-Dawson type anion (Fig. 14G) takes around 30 h after solution ($[\text{P}] > 20$ mM, $[\text{Mo}] = 100$ mM, $\text{pH} = 3$) preparation and an elevated temperature of about 80 °C.¹⁷² However, unlike the results of Petterson,^{167,168} they did not detect the $\{\text{PMo}_{11}\}$ complex (Fig. 14D) in the system with $[\text{Mo}] = 100$ mM and $[\text{P}] > 7$ mM, and $\{\text{PMo}_{12}\}$ is directly converted into its trilacunary form $\{\text{A-PMo}_9\}$ (Fig. 14E). At the same time, van Veen *et al.* suggested based on ^{31}P NMR and Raman spectroscopies that $\{\text{PMo}_{11}\}$ is more prominent at higher $[\text{Mo}]/[\text{P}] = 12$ than at lower $[\text{Mo}]/[\text{P}] = 9$ ratios, while the opposite is true for the $\{\text{A-PMo}_9\}$ species.¹⁷⁸ The results of ESI-MS studies on the pH-dependence of phosphomolybdate solutions from pH between 1.7 and 10.2 confirm the presence of both mono- and trilacunary species.¹⁷³ The confusion in the literature about the existence of

mono- $\{\text{PMo}_{11}\}$ and trilacunary $\{\text{A-PMo}_9\}$ complexes is understandable, since their spectroscopic characteristics (^{31}P NMR shifts and Raman spectra (Table 6)) are indeed very similar.

3.3.2.2. Speciation in the phosphito-molybdate system. The speciation in acidified molybdate solution with trivalent phosphorus anions, such as $\text{HP}^{\text{III}}\text{O}_3^{2-}$, $(\text{C}_6\text{H}_5)_3\text{P}^{\text{III}}\text{O}_3^{2-}$ or $(\text{CH}_3)_3\text{P}^{\text{III}}\text{O}_3^{2-}$, has been determined from both potentiometric and ^{31}P NMR measurements.^{174,175} In the system with phosphite, phenyl- and methylphosphonate anions structures similar to the $\text{H}_x[\text{P}_2^{\text{V}}\text{Mo}_5^{\text{VI}}\text{O}_{23}]^{(6-x)-}$ ($x = 0-2$) anion¹⁶⁹ (Fig. 14A) dominate.¹⁷⁴ It was shown that partial oxidation of phosphite to phosphate occurs, especially in the thin-walled NMR sample tubes when exposed to fluorescent light. In $\{\text{X}_2\text{M}_5\}$ -type HPO-Mos the phosphite anion $\text{HP}^{\text{III}}\text{O}_3^{2-}$ binds weaker to the Mo^{VI} -oxoframework than phosphate or phosphonates.

3.3.2.3. Speciation in arsenato-molybdate system. Aqueous equilibria in the arsenato-molybdate system $[\text{As}^{\text{V}}\text{O}_4]^{3-}$ – $[\text{Mo}^{\text{VI}}\text{O}_4]^{2-}$ have been studied by potentiometric and spectrophotometric methods (298 K, 3.0 M NaClO_4).¹⁸² These investigations established the formation of two series of complexes: colorless anions with two As atoms and five or six Mo atoms,



Table 6 Molybdophosphates stoichiometry, formation constants, chemical shifts and pK_a values

Molybdophosphate species	Abbreviation	<i>q, p, r</i> ^a	pK _a ¹⁶⁸	Isolated in solid state	δ(³¹ P), ppm ^b	Raman bands, cm ⁻¹	Formation constants lg K _c at 25 °C ^{a,c,168}
[P ₂ Mo ₅ ^{VI} O ₂₃] ⁶⁻	{P ₂ Mo ₅ } (Fig. 14A)	8, 5, 2		Yes ¹⁶⁹	2.35 ¹⁶⁷		61.97 (2)
H[P ₂ Mo ₅ ^{VI} O ₂₃] ⁵⁻		9, 5, 2	5.10	Yes ¹⁷⁰	1.94 ¹⁶⁷		67.07 (5)
H ₂ [P ₂ Mo ₅ ^{VI} O ₂₃] ⁴⁻		10, 5, 2	3.79	Yes ¹⁷¹	1.86 ¹⁶⁷		70.86 (8)
B-α-[P ^V Mo ₉ ^{VI} O ₃₄] ⁹⁻	B-{PMo ₉ } (Fig. 14F)	14, 9, 1		No	0.47–0.08 ¹⁶⁸		98.21 (7)
B-α-H[P ^V Mo ₉ ^{VI} O ₃₄] ⁸⁻		15, 9, 1	3.83	No			102.04 (8)
B-α-H ₂ [P ^V Mo ₉ ^{VI} O ₃₄] ⁷⁻		16, 9, 1	2.85	No			104.89 (8)
B-α-H ₃ [P ^V Mo ₉ ^{VI} O ₃₄] ⁶⁻		17, 9, 1	1.53	No			106.42 (8)
A-α-[P ^V Mo ₉ ^{VI} O ₃₁ (OH)(OH ₂) ₂] ⁴⁻	A-{PMo ₉ } (Fig. 14E)	16, 9, 1		Yes ¹⁷⁶	-1.00 to -1.15 ¹⁶⁸		104.71 (8)
A-α-[P ^V Mo ₉ ^{VI} O ₃₁ (OH ₂) ₃] ³⁻		17, 9, 1	2.46	Yes ¹⁷⁶		968, 854, 716, 641 ¹⁷²	107.17 (5)
[P ₂ Mo ₁₈ ^{VI} O ₆₂] ¹⁶⁻	{P ₂ Mo ₁₈ } (Fig. 14G)	34, 18, 2		Yes ¹⁷⁷	-2.53; ¹⁶⁸ -2.4 ¹⁷⁸	971, 962, 707 ¹⁷²	217.8 (1)
[P ^V Mo ₁₁ ^{VI} O ₃₉] ⁷⁻	{PMo ₁₁ } (Fig. 14D)	17, 11, 1		Yes ¹⁷⁹ (SXR only for substituted anions ¹⁸⁰)	-0.78 to -1.20 ¹⁶⁸	—	118.68 (8)
H[P ^V Mo ₁₁ ^{VI} O ₃₉] ⁶⁻		18, 11, 1	4.42	No		—	123.10 (5)
H ₂ [P ^V Mo ₁₁ ^{VI} O ₃₉] ⁵⁻		19, 11, 1	2.95	No		974, 969, 957, 875, 763 ¹⁷²	126.05 (5)
[P ^V Mo ₁₂ ^{VI} O ₄₀] ¹³⁻	{PMo ₁₂ } (Fig. 14C)	23, 12, 1		Yes ¹⁸¹	-3.20 ¹⁶⁸	988, 965, 894, 602 ¹⁷²	139.7 (1)

^a Stoichiometric coefficients and formation constants for the equilibrium: $p[\text{Mo}^{\text{VI}}\text{O}_4]^{2-} + q\text{H}^+ + r[\text{HP}^{\text{VO}_4}]^{2-} \rightleftharpoons (\text{H}^+)_q(\text{Mo}^{\text{VI}}\text{O}_4^{2-})_p(\text{HP}^{\text{VO}_4}^{2-})_r$. ^b δ³¹P relative to H₃PO₄ (85%). ^c Ionic strength μ = 3 M (NaCl).

and yellow complexes with [As]/[Mo] ratio 1:9 ({AsMo₉}) (the same structure as {PMo₉} see Fig. 14E and Table 7). Pettersson suggests for {As₂Mo₅} the same structure as for {P₂Mo₅} (Fig. 14A),¹⁸² while the {As₂Mo₆} complex was isolated in all protonated states and characterized by SXR (Table 7 and Fig. 14B). Applying large-angle X-ray scattering (LAXS) investigations on {AsMo₉} complexes showed them to have the same basic trilacunary structure (Fig. 14E) as the [As^VMo₉^{VI}O₃₁(OH₂)₃]³⁻ anion present in Na₃[As^VMo₉^{VI}O₃₁(OH₂)₃].13H₂O.¹⁸³ Later Ueda *et al.* showed,¹⁷² that at As^V concentrations lower than 10 mM, in contrast to the phosphate system, the corresponding Mo^{VI}-As^V system did not produce the Keggin-type [As^VMo₁₂^{VI}O₄₀]¹³⁻ anion (the same structure as {PMo₁₂} see Fig. 14C) but a labile complex (tentatively formulated as [As^VMo₁₀^{VI}O₃₇H₅]⁴⁻) at ambient temperature and

higher As^V concentration. The [As^VMo₁₀^{VI}O₃₇H₅]⁴⁻ complex transforms directly into [As^VMo₆^{VI}O₂₆H₂]⁴⁻ (Fig. 14B). The Dawson-type [As^VMo₁₈^{VI}O₆₂]¹⁶⁻ complex (the same structure as {P₂Mo₁₈} see Fig. 14G), which was formed upon heating at 80 °C for 5 h, also underwent subsequent transformation into the more stable [As^VMo₆^{VI}O₂₆H₂]⁴⁻ complex (Fig. 14B) at As(v) concentrations higher than 20 mM.

3.3.2.4. Hydrolytic stability of HPOMos with Keggin structure. {PMo₁₂} and {AsMo₁₂}. According to Raman spectroscopic data, the Keggin molybdophosphate (Fig. 14C) firstly appears 1 min and 23 s after the acidified molybdate solution is mixed with the acidified phosphate solution at pH 1, which is the optimal acidity for the {PMo₁₂} formation.¹⁸⁸ The UV-vis, IR and

Table 7 Molybdoarsenates stoichiometry, formation constants and pK_a values

Molybdoarsenate species	Abbreviation	<i>q, p, r</i> ^a	pK _a	Isolated in solid state	Formation constants lg K _c at 25 °C ^{a,b,182}
[As ₂ Mo ₅ ^{VI} O ₂₃] ⁶⁻	{As ₂ Mo ₅ } (Fig. 14A)	8, 5, 2		No	60.92 (0.04)
[As ₂ Mo ₆ ^{VI} O ₂₆] ⁶⁻	{As ₂ Mo ₆ } (Fig. 14B)	10, 6, 2		Yes ¹⁸⁴	75.25 (0.04)
H[As ₂ Mo ₆ ^{VI} O ₂₆] ⁵⁻		11, 6, 2		Yes ¹⁸⁵	80.73 (0.05)
H ₂ [As ₂ Mo ₆ ^{VI} O ₂₆] ⁴⁻		12, 6, 2		Yes ¹⁸⁶	84.07 (0.07)
H ₃ [As ^V Mo ₉ ^{VI} O ₃₄] ⁶⁻	{AsMo ₉ } (Fig. 14E and F)	14, 9, 1		Yes ¹⁸³	100.11 (0.02)
H ₄ [As ^V Mo ₉ ^{VI} O ₃₄] ⁵⁻		15, 9, 1	3.70 ¹⁸²	No	103.81 (0.05)
H ₅ [As ^V Mo ₉ ^{VI} O ₃₄] ⁴⁻		16, 9, 1	2.33 ¹⁸²	No	106.14 (0.05)
H ₆ [As ^V Mo ₉ ^{VI} O ₃₄] ³⁻		17, 9, 1	1.33 ¹⁸²	No	107.47 (0.06)
[As ₂ Mo ₁₈ ^{VI} O ₆₂] ¹⁶⁻	{As ₂ Mo ₁₈ } (Fig. 14G)	34, 18, 2		Yes ¹⁸⁷	—

^a Stoichiometric coefficients and formation constants for: $p[\text{Mo}^{\text{VI}}\text{O}_4]^{2-} + q\text{H}^+ + r[\text{HAS}^{\text{VO}_4}]^{2-} \rightleftharpoons (\text{H}^+)_q(\text{Mo}^{\text{VI}}\text{O}_4^{2-})_p(\text{HAS}^{\text{VO}_4}^{2-})_r$. ^b Ionic strength μ = 3 M (NaClO₄).



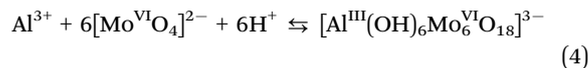
^{31}P NMR spectra of $\{\text{PMo}_{12}\}$ solution recorded at pH values between 0 and 12 manifest that already at pH 1.5 the intact Keggin anion decomposes into lacunary forms, and at pH 7, monomeric molybdate and phosphate dominate in the solution.^{189–192} $\{\text{PMo}_{12}\}$ co-exists with its monolacunary form $\{\text{PMo}_{11}\}$ between pH 1.5 and 5 (Fig. 15).¹⁹⁰

$\{\text{SiMo}_{12}\}$ and $\{\text{GeMo}_{12}\}$. Already in 1975, it was established that $\alpha\text{-H}_4[\text{Si}^{\text{IV}}\text{Mo}_{12}^{\text{VI}}\text{O}_{40}]$ ($\{\text{SiMo}_{12}\}$) acid is formed between pH 3.8 and 4.8 and the β -isomer at lower pH values between 1.0 and 1.8.^{193,194} The yield of both α - and β -forms (at pH 4.0 and 1.2, respectively) is independent of molybdate concentrations between 0.015 and 0.100 M. Later, using ESI-MS the formation of molybdosilicates was detected between pH 1 and 2.¹⁹⁵ The decomposition of $\{\text{SiMo}_{12}\}$ is 50% completed immediately after dissolution at pH 4.1.¹⁹¹ As the pH is increased, the decomposition also proceeds rapidly and at pH 5.2 the acid is completely decomposed. The $\alpha\text{-}[\text{Ge}^{\text{IV}}\text{Mo}_{12}^{\text{VI}}\text{O}_{40}]^{4-}$ ($\{\text{GeMo}_{12}\}$) Keggin complex has a stability range similar to $\{\text{SiMo}_{12}\}$ and is reported to form mono-lacunary $[\text{Ge}^{\text{IV}}\text{Mo}_{11}^{\text{VI}}\text{O}_{39}]^{8-}$ between pH 4.3 and 4.5. The formation rate of $[\text{Ge}^{\text{IV}}\text{Mo}_{11}^{\text{VI}}\text{O}_{39}]^{8-}$ is faster than that of $[\text{Si}^{\text{IV}}\text{Mo}_{11}^{\text{VI}}\text{O}_{39}]^{8-}$, while the formation rate of $\{\text{GeMo}_{12}\}$ is slower than the one of $\{\text{SiMo}_{12}\}$.¹⁹⁶ Thus, the stability of the lacunary anion relative to the Keggin anion, is higher for molybdogermanates than for molybdosilicates, which leads to a higher number of sandwich derivatives for molybdogermanates.¹⁹⁷

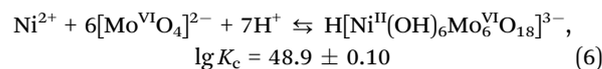
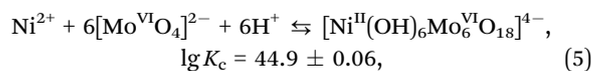
At pH above 2, the stability of $\{\text{XMo}_{12}\}$ anions can be increased by the addition of organic solvents.¹⁹⁸ Heteropolyanions with protonated amino acids as cations can be stable up to pH 6.2, which can provide valuable insights for correct application of these POMs in medical usage.¹⁹⁹

In general, complexes based on lacunary HPOMs are more unstable in aqueous solution than, for example, their tungsten analogs,¹⁹⁷ and there is still insufficient data on lacunary HPOMs stability in solution. Heteropolymolybdates follow a generally accepted trend – the substitution of addenda atoms with transition metals in Keggin clusters reduces the net charge and stabilizes therewith the anions (Table 8). Thus, monosubstituted germanomolybdates $[\text{Ge}^{\text{IV}}\text{Mo}_{11}^{\text{VI}}\text{M}(\text{H}_2\text{O})\text{O}_{39}]^{6-}$ ($\text{M} = \text{Co}^{\text{II}}, \text{Ni}^{\text{II}}$ and Mn^{II} , have the same structure as unsubstituted $\{\text{GeMo}_{12}\}$ see Fig. 14C) and stay intact in aqueous solution between pH 4 and 10.²⁰⁰

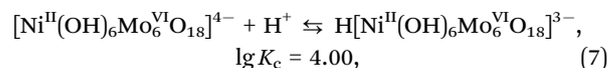
3.3.2.5. Hydrolytic stability of HPOMs with Anderson structure. HPOMs with Anderson structure can be synthesized from aqueous solution between pH 4 and 5 and are stable in near neutral aqueous media.^{1,201} There are only a few papers discussing the hydrolytic stability of the Anderson POMs with Al^{III} ,²⁰² Cr^{III} ,²⁰³ Ni^{II} ¹⁷ and Ga^{III} ²⁰⁴ as a central heteroatom (Fig. 16A and B). UV-vis studies revealed that $[\text{Cr}^{\text{III}}(\text{OH})_6\text{Mo}_6^{\text{VI}}\text{O}_{18}]^{3-}$ is stable in a fairly narrow pH range of 4.50 to 5.83 (Fig. 17).²⁰³ In solutions with an aluminum concentration of 1 mM and ratio $[\text{Mo}]/[\text{Al}]$ between 0.1 and 1, $[\text{Al}^{\text{III}}(\text{OH})_6\text{Mo}_6^{\text{VI}}\text{O}_{18}]^{3-}$ is the predominate species over IPOMs (octa- and heptamolybdate) between pH 2 and 5 (Fig. 17).²⁰² According to potentiometric and ^{27}Al NMR data, the formation constant for this species is $\lg K_c = 50.95 \pm 0.04$ in 0.6 M NaCl medium at 25 °C based on reaction (4):



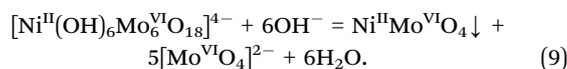
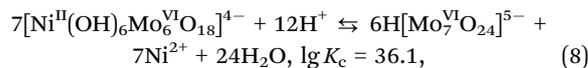
The $[\text{Ni}^{\text{II}}(\text{OH})_6\text{Mo}_6^{\text{VI}}\text{O}_{18}]^{4-}$ $\{\text{NiMo}_6\}$ complex forms in 0.1 M NaNO_3 medium at 25 °C according to reactions (5) and (6) with formation constants:



Potentiometric studies reveal that upon acidification, the $[\text{Ni}^{\text{II}}(\text{OH})_6\text{Mo}_6^{\text{VI}}\text{O}_{18}]^{4-}$ $\{\text{NiMo}_6\}$ complex is firstly protonated ($Z = n(\text{H}^+)/n([\text{MoO}_4]^{2-}) = 1-1.17$) (Fig. 1B):



and then at $Z > 1.29$ converted to $\{\text{Mo}_7\}$ (Fig. 1B) accompanied by the release of Ni^{2+} (8) and a decrease in acidity (9) leads to $\text{NiMoO}_4 \cdot x\text{H}_2\text{O}$ precipitation:¹⁷



The described decomposition pathway could be principally applied to Anderson HPOMs with different di- and trivalent ions. Noteworthy, the organic functionalization can significantly increase the stability of Anderson HPOMs. Thus, the alkoxy-functionalized Anderson gallatomolybdate (Fig. 16B) is reported by ESI-MS to be stable up to pH 9 (Fig. 17).²⁰⁴ The stability of the Anderson-like decamolybdocobaltate $[\text{H}_4\text{Co}_2^{\text{III}}\text{Mo}_{10}^{\text{VI}}\text{O}_{38}]^{6-}$ (Fig. 16C) has been verified by Raman and UV-vis spectroscopies and showed that the Co-containing HPOMs is the predominant species between pH 1 and 4.5 (Fig. 17).²¹²

3.4. Polyoxotungstates (POTs)

Polyoxotungstate species are overall more stable than their molybdate analogues due to the enthalpy factor of the condensation process when the tetrahedrally coordinated W^{VI} or Mo^{VI} ions expand their coordination spheres from four to six during POM formation reaction.^{1–3} Part of this effect can be rationalized in terms of the slightly larger force constant of the W–O bonding as compared to that of the Mo–O bonding.²¹³ Owing to the existence of a number of simultaneous equilibria, some of which are very slow to establish, and the metastability of some anions, speciation in isopolytungstates (IPOTs) solutions has not been studied as extensively as in Mo^{VI} or V^{V} systems. However, the solution stability of the phosphotungstate anion $[\text{P}^{\text{V}}\text{W}_{12}^{\text{VI}}\text{O}_{40}]^{3-}$ ($\{\text{PW}_{12}\}$, Fig. 2B) was investigated as thoroughly as no other POM species.³⁸

3.4.1. Isopolytungstates (IPOTs)

3.4.1.1. Ion-distribution diagram for IPOTs. Despite the similarity of the solid monomeric precursors for Mo^{VI} and W^{VI} ,



Table 8 Introductory information on pH stability of various sandwich-like archetypes for some HPOMos and HPOTs constructed from lacunary Keggin anions containing 3d- or 4f-metals^a

HPOMs species	Stable in the pH region	Analysis method	Fingertip characteristics	Ref.
<i>HPOMos based on lacunary forms of {PMo₁₂}</i>				
$[M(P^V Mo_{11} O_{39})_2]^{10-}$ (M = Ce ^{IV} , Th ^{IV})	1–5	³¹ P NMR	$\delta(^{31}P) = 2.55$ ppm for Ce ^{IV} and 2.52 for Th ^{IV}	205
$[M(P^V Mo_{12} O_{40})(P^V Mo_{11} O_{39})]^{6-}$ (M = Zr ^{IV} , Hf ^{IV})	2–5.3	³¹ P NMR	$\delta(^{31}P) = -3.5$ ppm for Hf ^{IV} and -3.2 for Zr ^{IV}	205
<i>HPOMos based on lacunary forms of {GeMo₁₂}</i>				
$[Ge^{IV} Mo_{11} M(H_2O)_3]^{6-}$ (M = Co ^{II} , Ni ^{II} and Mn ^{II})	4–10	UV-vis	Absorption at 207, 231 and 568 nm (only for Co ^{II})	200
$[Cu^{II}_2(\beta\text{-}Y\text{-}Ge^{IV} Mo_9^V O_{33})_2]^{12-}$	4.5–10	UV-vis	Absorption at 208 and 233 nm	206
$[Ni^{II}_4(H_2O)_2(\alpha\text{-}B\text{-}Ge^{IV} Mo_9^V O_{34})_2]^{12-}$	3–10	UV-vis	Absorption at 206 and 232 nm	206
<i>HPOTs based on lacunary forms of {PW₁₂}</i>				
$[Ni^{II}(H_2O)P^V W_{11} O_{39}]^{5-}$	pH 4 (at least for 4 h)	ESI-MS and UV-vis	$m/z = 683.5$ (for anion with net charge -4)	207
$[H_2P^V W_9^VI Ni_4^II O_{34}(OH)_3(H_2O)_6]^{2-}$	At pH 6 converted to $[Ni^{II}(H_2O)P^V W_{11} O_{39}]^{5-}$ within 7 h; at pH < 3 unstable	UV-vis	Shoulder located between 280 and 260 nm	208
$[Na_2(U^VI O_2)_2(P^V W_9^VI O_{34})_2]^{12-}$	At pH 7; at pH 6 stable for 4 days; at pH 8.1 30% decomposes after 7 days; at pH 9.4 full decomposition in 3 h	³¹ P and ¹⁸³ W NMR	$\delta(^{31}P) = -9.7$ ppm; $\delta(^{183}W) = -124.6, -141.1, -148.7, -182.2, -209.6$	209
$[Co^{II}(H_2O)_2(\alpha\text{-}P^V W_9^VI O_{34})_2]^{10-}$	6–10	ESI-MS and UV-vis	$m/z = 1202.6$ for $[Co_4^{II}(P^V W_9^VI O_{34})_2 HNa_5]^{4-}$	210
$[Co^{II}(H_2O)_6(OH)_3(HP^V O_4)_2(P^V W_9^VI O_{34})_3]^{16-}$	5.5–11	SAXS	—	56
<i>HPOTs based on lacunary forms of {SiW₁₂}</i>				
$[Ni^{II}(H_2O)Si^{IV} W_{11} O_{39}]^{6-}$	At pH 4 (at least for 4 h)	ESI-MS and UV-vis	$m/z = 684.3$ for anion with net charge -4	207
$[Co^{II}(H_2O)Si^{IV} W_{11} O_{39}]^{6-}$	6–10	ESI-MS and UV-vis	$m/z = 1406.6$ for $[(Co^{II} Si^{IV} W_{11} O_{39})K_2 H_2]^{2-}$	210
$\alpha\text{-}$ and $\beta\text{-}[Si^{IV} W_9^VI O_{37}(M(H_2O))_3]^{10-}$ (M = Co ^{II} , Ni ^{II} , Mn ^{II} , Cu ^{II})	5–10	UV-vis	No data provided	211
$\alpha\text{-}$ and $\beta\text{-}[Si^{IV} W_9^VI O_{37}(M(H_2O))_3]^{7-}$ (M = Fe ^{III} , Al ^{III} , Ga ^{III})	5–10	UV-vis	No data provided	211
<i>HPOTs based on lacunary forms of {GeW₁₂}</i>				
$[Ni^{II}(H_2O)Ge^{IV} W_{11} O_{39}]^{6-}$	pH 4 (at least for 4 h)	ESI-MS and UV-vis	$m/z = 694.8$ (for anion with net charge -4)	207
<i>HPOTs based on lacunary {As^{III}W₉}</i>				
$[Na\{2\text{-}((CH_3)_2HNCH_2)C_6H_4Sb^{III}\}As_2^{III} W_{19}^VI O_{67}(H_2O)]^{10-}$	3–7 (at least for 5 h)	UV-vis, ¹⁸³ W NMR	Absorption bands at 200 and 250–254 nm; $\delta(^{183}W) = -87.2, -95.1, -102.9, -103.3, -105.9, -138.1, -141.1, -157.8, -174.3$ ppm	24

^a The data for this table were searched in May 2020 in the scientometric databases *Scopus* and *Web of Science* with key words: “sandwich polyoxometalate” and “solution stability”.

differences do occur in the course of the aggregation reactions, while only one type of isopolyspecies, $[M_7^VI O_{24}]^{6-}$ (M = Mo^{VI} and W^{VI}), is present in both systems and all others vary. The speciation in acidified aqueous solution of $[W_7^VI O_{24}]^{6-}$ has been studied predominantly by potentiometry,^{128,214–216} Raman spectroscopy,^{26,28} ¹⁸³W, ¹⁷O and ¹H NMR spectroscopy,^{217,218} and lately by ESI-MS.^{19,219} When an aqueous solution of tungstate is acidified, condensation reactions lead to the formation

of IPOT anions, of which the following types are certainly present in solution and in the solid phase (Table 9): heptatungstate (paratungstate A) $[W_7^VI O_{24}]^{6-}$ ($\{W_7\}$, Fig. 18A); paratungstate B $[W_{12}^VI O_{40}(OH)_2]^{10-}$ ($\{W_{12}O_{42}\}$, Fig. 18B); α -metatungstate $[H_2W_{12}^VI O_{40}]^{6-}$ ($\{W_{12}O_{40}\}$, Fig. 18C) and decatungstate $[W_{10}^VI O_{32}]^{4-}$ ($\{W_{10}\}$, Fig. 18D). Studies of how the pH affects aqueous solutions of $[W_7^VI O_{24}]^{6-}$ are difficult to perform, partly due to very slow equilibria, and they are often problematic to



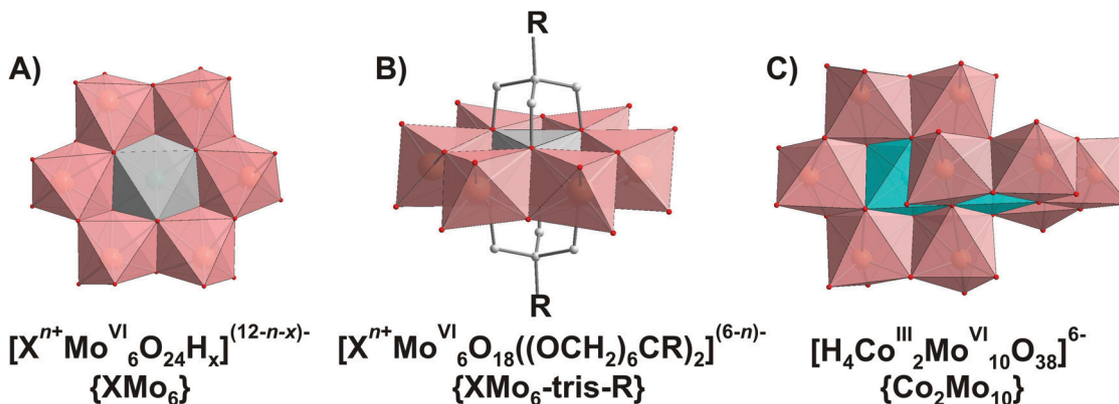


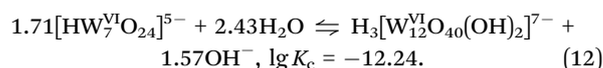
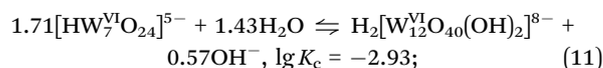
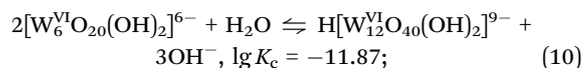
Fig. 16 HPOMos with Anderson and Anderson-like structures: (A) Anderson type anion $[X^{n+}Mo_6^VI O_{24}H_x]^{(12-n-x)-}$, where X – heteroion (e.g. Ni^{II} ,¹⁷ Al^{III} ,²⁰² or Cr^{III} ,²⁰³), $n = 2-6$, $x = 0-6$; (B) functionalized with tris-alkoxo ligands Anderson type anion $[X^{n+}Mo_6^VI O_{18}((OCH_2)_6CR)_2]^{(6-n)-}$, where X – heteroion (e.g. Ga^{III} ,²⁰⁴), $n = 2-3$, R = $-NH_2$, CH_2OH , $-CH_3$ etc.; (C) Anderson-like decamolybdocobaltate $[H_4Co_2^III Mo_{10}^VI O_{38}]^{6-}$. Color code: { MoO_6 }, pink; { XO_6 }, grey; { CoO_6 }, light blue; C, light grey; O, red.

interpret. Notably, the age of the solution and the total W^{VI} concentration (aggregation does not occur in highly diluted solutions) plays a role.

NMR studies confirm that above pH 5, the major polytungstate species are heptatungstate $\{W_7\}$ (Fig. 18A) and paratungstate B $\{W_{12}O_{42}\}$ (Fig. 18B), which have been isolated in numerous salts (Fig. 19 and Table 9). At pH lower than 5, equilibria are established very slowly, and most IPOTs anions are metastable, with the exception of the metatungstate anion $\alpha-[H_2W_{12}O_{40}]^{6-}$ (Fig. 18C). Acidification of paratungstate solutions leads to “pseudometatungstates”, which ultimately form the stable metatungstate anion (Fig. 18C). From solutions suggested to contain “pseudometatungstates”, the salt $K_6H_4[W_{11}O_{38}]\cdot 11H_2O$, with an anion of C_s symmetry (Fig. 18E), has been crystallized.²²⁰ However, the ^{183}W NMR spectrum of the similar acidified paratungstate solution shows a species with 11 resonances of equal intensity, which has tentatively been assigned to $[H_7W_{11}O_{40}]^{7-}$, a lacunary form of paratungstate-B (Fig. 18F), and also two lines attributed to decatungstate $\{W_{10}\}$ (Fig. 18D),

once known as tungstate-Y, a species that is stabilized in nonaqueous solvents.

3.4.1.2. *Aging of tungstate solution. Conversions of paratungstates.* It is a well-known experimental fact that, from aged fully equilibrated tungstate solutions close to neutral and slightly acidic pH (Fig. 19), the isolation of paratungstate B $\{W_{12}O_{42}\}$ (Fig. 18B) prevails.²¹⁸ Potentiometric studies in the $[W^{VI}O_4]^{2-} - H^+$ system with different aging times (0–20 000 min from the preparation) show that heptatungstate (Fig. 18A) and possibly unstable hexatungstate transform over time into protonated forms of paratungstate B (Fig. 18B), the existence of which was proven by crystallization,²²³ according to eqn (10)–(12).²¹⁵



Taking together the results obtained by ESI-MS, ^{183}W NMR and Raman spectroscopy, it must be concluded that the heptatungstate (Fig. 18A) anion is the main species in freshly prepared solutions under nearly neutral conditions (pH = 5.8 and 6.8).¹⁹ During aging of $Na_6[W_7^{VI}O_{24}]$ solutions, paratungstate B, $Na_{10}[W_{12}^{VI}O_{40}(OH)_2]$, crystallizes within an hour after preparation. However, ^{183}W NMR shows a very low concentration of paratungstate B $\{W_{12}O_{42}\}$ (Fig. 18B) in neutral to slightly acidic solutions, indicating that after conversion paratungstate B tends to crystallize immediately.²¹⁸ The reverse transition is also possible as upon prolonged heating of a $Na_{10}[W_{12}^{VI}O_{40}(OH)_2]$ solution, the paratungstate B (Fig. 18B) anions partly transform to $[W_7^{VI}O_{24}]^{6-}$ (Fig. 18A). To conclude, heptatungstate $\{W_7\}$ (Fig. 18A) is more stable in solution, while paratungstate B $\{W_{12}O_{42}\}$ (Fig. 18B) is more likely to crystallize from solution.

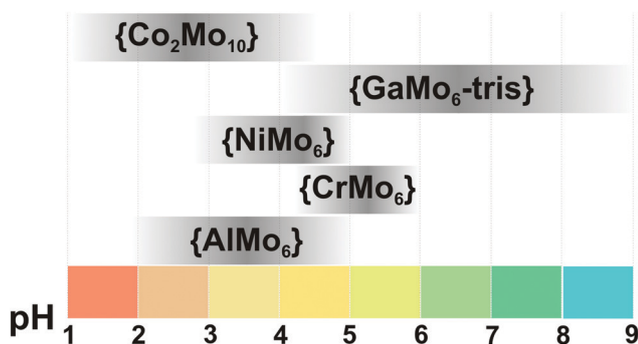


Fig. 17 Hydrolytic stability of Anderson based HPOMOs based on ref. 17, 202–204 and 212. The maximum intensity of grey color in each box with a single species corresponds to its maximum concentration in the chosen pH region. $\{AlMo_6\} - [Al^{III}(OH)_6Mo_6^VI O_{18}]^{3-}$; $\{CrMo_6\} - [Cr^{III}(OH)_6Mo_6^VI O_{18}]^{3-}$; $\{NiMo_6\} - [Ni^{II}(OH)_6Mo_6^VI O_{18}]^{4-}$; $\{GaMo_6\text{-tris}\} - [Ga^{III}Mo_6^VI O_{18}(OH)_3((OCH_2)_3CCH_2OH)]^{3-}$; $\{Co_2Mo_{10}\} - [H_4Co_2^{II}Mo_{10}^VI O_{38}]^{6-}$.



Table 9 IPOTs stoichiometry, formation constants and ^{183}W and ^{17}O chemical shifts

IPOT species	p, q^a	pK_a	Isolated in solid state	$\delta(^{183}\text{W}),^b$ ppm	$\delta(^{17}\text{O}),^b$ ppm	Formation constants ^a $\lg K_c$ at 25 °C		
						Cruywagen ^{214c}	Rozantsev ^{215d}	Rozantsev ^{216e}
$[\text{W}^{\text{VI}}\text{O}_4]^{2-}$	1, 0		Yes					
$[\text{HW}^{\text{VI}}\text{O}_4]^-$	1, 1	4.6 ²¹⁴	No					
$[\text{W}_6^{\text{VI}}\text{O}_{20}(\text{OH})]^{5-}$	6, 7		No					
$[\text{W}_6^{\text{VI}}\text{O}_{20}(\text{OH})_2]^{6-}$	6, 6		No			49.01	51.66 (1.29)	53.68 (0.28)
$[\text{W}_7^{\text{VI}}\text{O}_{24}]^{6-}$ (Fig. 18A)	7, 8		Yes ²¹⁸	268, -106, -189 ²¹⁷	71, 254, 295, 314, 595, 590, 624, 648 ²¹⁸	65.19		
$\text{H}[\text{W}_7^{\text{VI}}\text{O}_{24}]^{5-}$	7, 9	4.77 ²¹⁴	Yes ²²¹			69.96	72.31 (1.81)	76.59 (0.7)
$[\text{W}_{12}^{\text{VI}}\text{O}_{40}(\text{OH})_2]^{10-}$ (Fig. 18B)	12, 14		Yes ²²²	-109, -114, -116, -147 ²¹⁷		115.38	119.45 (2.99)	123.38 (1.27)
$\text{H}[\text{W}_{12}^{\text{VI}}\text{O}_{40}(\text{OH})_2]^{9-}$	12, 16	4.59 ²¹⁷	Yes ²²³					
$\text{H}_2[\text{W}_{12}^{\text{VI}}\text{O}_{40}(\text{OH})_2]^{8-}$	12, 16						129.01 (3.23)	
$\text{H}_3[\text{W}_{12}^{\text{VI}}\text{O}_{40}(\text{OH})_2]^{7-}$	12, 17						133.70 (3.34)	
$\alpha\text{-}[\text{H}_2\text{W}_{12}^{\text{VI}}\text{O}_{40}]^{6-}$ (Fig. 18C)	12, 18		Yes ²²⁴	-123 ²¹⁷		80, 415, 418, 723 ²¹⁷	136.98 (3.42)	149.59 (0.7)
$\beta\text{-}[\text{H}_2\text{W}_{12}^{\text{VI}}\text{O}_{40}]^{6-}$	12, 18		No	-112, -127, -136 ²¹⁷		61, 397, 442, 725 ²¹⁷		
$[\text{W}_{10}^{\text{VI}}\text{O}_{32}]^{4-}$ (Fig. 18D)	10, 16		Yes ²²⁵	-105, -118 ²¹⁷				129.63 (0.98)

^a Stoichiometric coefficients and formation constants for: $p[\text{W}^{\text{VI}}\text{O}_4]^{2-} + q\text{H}^+ \rightleftharpoons [\text{H}_q\text{W}_p^{\text{VI}}\text{O}_m]^{n-} + (q/2 - z/2)\text{H}_2\text{O}$. ^b $\delta^{183}\text{W}$ relative to Na_2WO_4 , $\delta^{17}\text{O}$ relative to H_2O . ^c Ionic strength $\mu = 1\text{ M}$ (NaCl). ^d $\mu = 0.4\text{ M}$ (KNO_3). ^e $\mu = 0.1\text{--}0.5\text{ M}$ (NaNO_3).

3.4.1.3. *Metatungstate* $[\text{H}_2\text{W}_{12}^{\text{VI}}\text{O}_{40}]^{6-}$ in solution. α -Keggin metatungstate has been reported in 4 protonation states $[\text{H}_n\text{W}_{12}^{\text{VI}}\text{O}_{40}]^{(8-n)-}$ ($n = 1\text{--}4$), in which the “non-acidic” protons occupy the central tetrahedral cavity (Fig. 18C). Launay reported the formation of a monoprotated species, $[\text{H}_1\text{W}_{12}^{\text{VI}}\text{O}_{40}]^{7-}$ during the reduction of the diprotated anion $[\text{H}_2\text{W}_{12}^{\text{VI}}\text{O}_{40}]^{6-}$.²²⁶ The NMR spectra (^{183}W , ^{17}O , ^1H) also reveal a conversion of paratungstate B, $[\text{W}_{12}^{\text{VI}}\text{O}_{40}(\text{OH})_2]^{10-}$ (Fig. 18B) to the monoprotated Keggin anion, $[\text{H}_1\text{W}_{12}^{\text{VI}}\text{O}_{40}]^{7-}$, which leads to the formation of diprotated $[\text{H}_2\text{W}_{12}^{\text{VI}}\text{O}_{40}]^{6-}$ (Fig. 18C) upon further acidification. The existence of tri-protonated $[\text{H}_3\text{W}_{12}^{\text{VI}}\text{O}_{40}]^{5-}$ or tetra-protonated $[\text{H}_4\text{W}_{12}^{\text{VI}}\text{O}_{40}]^{4-}$ has not been widely accepted, because the third or fourth proton is regarded as an external acidic proton. Thus, uncertainty still remains about the external (acidic) and internal (non-acidic) location of the protons. Under non-aqueous or mixed water-solvent conditions, $[\text{H}_3\text{W}_{12}^{\text{VI}}\text{O}_{40}]^{5-}$ and $[\text{H}_4\text{W}_{12}^{\text{VI}}\text{O}_{40}]^{4-}$ were reported, with all three or four protons assigned to the internal cyrtand tetrahedral cavity.^{227–229} Although $[\text{H}_3\text{W}_{12}^{\text{VI}}\text{O}_{40}]^{5-}$ is not formed in aqueous W^{VI} (0.05–0.50 M) solutions, the correct choice of counteraction (e.g. tetrabutylammonium) can lead to its isolation from aqueous W^{VI} (0.05 M) solutions at pH 4.²³⁰

3.4.1.4. *Comparison of IPOTs in solution and in the solid state.* Very slow equilibrium reactions in tungstate solutions result in very long crystallization times and low yields. Along with the IPOTs dominating in acidified aqueous tungstate solution, $\{\text{W}_7\}$ (Fig. 18A), $\{\text{W}_{12}\text{O}_{42}\}$ (Fig. 18B), $\{\text{W}_{12}\text{O}_{40}\}$ (Fig. 18C) and $\{\text{W}_{10}\}$ (Fig. 18D) (Table 9), the following IPOTs could be crystallized. Salts containing fragments of the heptatungstate structure, $\text{Na}_3[\text{H}_3\text{W}_6^{\text{VI}}\text{O}_{22}]^{3-}$ (Fig. 18H) and $\text{K}_7[\text{HW}_5^{\text{VI}}\text{O}_{19}]^{2-}$ (Fig. 18G), have been isolated from concentrated weakly alkaline aqueous

tungstate solutions, but there is no evidence that these anions are stable in solution. These penta- and hexatungstates contain $\{\text{WO}_6\}$ octahedra with free octahedra faces showing three terminal oxygen atoms in an unstable *fac* configuration (Fig. 18H and G). Obviously, $[\text{H}_3\text{W}_6^{\text{VI}}\text{O}_{22}]^{5-}$ and $[\text{HW}_5^{\text{VI}}\text{O}_{19}]^{7-}$ react immediately upon dissolution to become a more stable, more condensed heptatungstate ion (Fig. 18A), and therefore might exist in such low concentrations that they cannot be detected experimentally.²¹⁸

Concentrated tungstate solutions with $c([\text{W}^{\text{VI}}\text{O}_4]^{2-}) = 1.5\text{ M}$ were used to generate two high nuclearity derivatives of $[\text{H}_4\text{W}_{11}^{\text{VI}}\text{O}_{38}]^{6-}$ (Fig. 18E), $[\text{H}_4\text{W}_{22}^{\text{VI}}\text{O}_{74}]^{12-}$ (Fig. 18I) and $[\text{H}_{10}\text{W}_{34}^{\text{VI}}\text{O}_{116}]^{18-}$ (Fig. 18J) at pH 3.4 and 2.4, respectively in the presence of Na_2SO_3 .²³³ $[\text{H}_4\text{W}_{22}^{\text{VI}}\text{O}_{74}]^{12-}$ consists of two $[\text{H}_4\text{W}_{11}^{\text{VI}}\text{O}_{38}]^{6-}$ fragments, and in the case of $[\text{H}_{10}\text{W}_{34}^{\text{VI}}\text{O}_{116}]^{18-}$ two undecatungstates are connected *via* one paratungstate B cluster $\{\text{W}_{12}\text{O}_{42}\}$ (Fig. 18B). So far, the stability of high nuclearity species was only demonstrated in the gas phase with ESI-MS. The synthetic procedures for synthesizing these IPOTs require the presence of the sulfite anion, and control experiments without sulfite do not allow the isolation of the compounds in high yields. Once again, not only pH and acidity affect the speciation of POMs, but also the substance present in the medium including choice of counteraction, aging time and concentration.

3.4.1.5. *IPOTs in non-aqueous solutions.* The Lindqvist $[\text{W}_6^{\text{VI}}\text{O}_{19}]^{2-}$ (isostructural with the corresponding niobate Fig. 8A) and decatungstate anions $\{\text{W}_{10}\}$ (Fig. 18D) are well characterized in non-aqueous solutions. The $\{\text{W}_{10}\}$ complex becomes stable in the presence of many organic solvents such as acetonitrile, 1,4-dioxane, methanol, dimethylsulfoxide and *N,N*-dimethylformamide.²³⁰ All IPOTs that are stable in



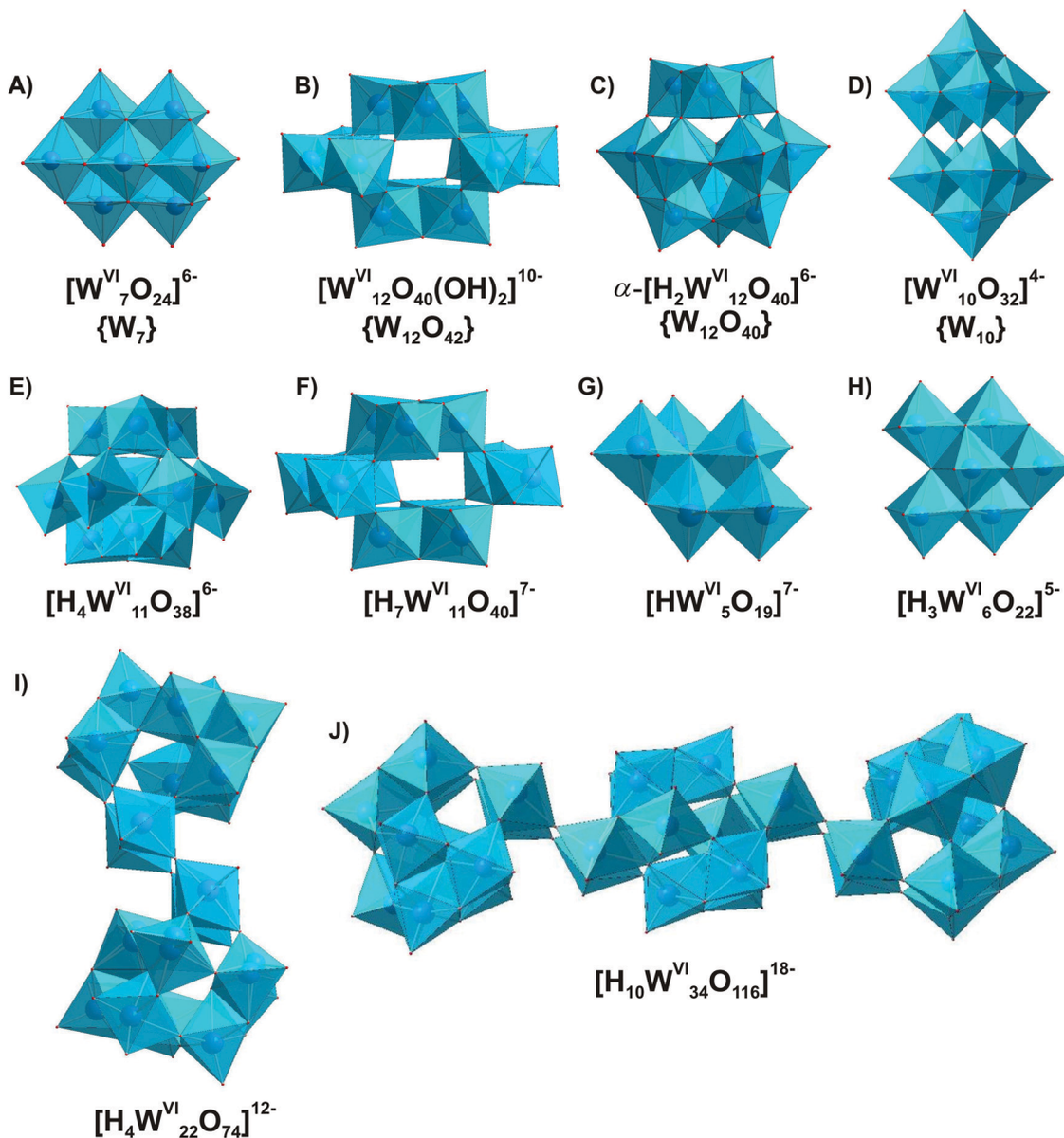


Fig. 18 I POTs with nuclearity up to 34 addenda atoms, which can be present and/or isolated from aqueous media. Color code: $\{WO_6\}$, pink; O, red.

non-aqueous media, which are α - and β - $\{W_{12}O_{40}\}$, $\{W_{10}\}$ and $[W_6^{VI}O_{19}]^{2-}$, are known to be easily reducible.⁵⁷

3.4.2. Heteropolytungstates (HPOTs). Heteropolytungstates (HPOTs) are a rapidly growing class of POM chemistry with the largest number of synthesized compounds due to the variety of stably lacunary precursors that can be easily formed from Keggin (Fig. 20) and Dawson (Fig. 24) archetypes and combined with d- or f-metals as well as with organic ligands (Table 8). The speciation and self-assembly of POMs are intimately related, therefore this section is of particular interest not only for targeted application, but also for the development of new synthetic strategies.

3.4.2.1. Hydrolytic stability of HPOTs with Keggin structure. The $[XW_{12}^{VI}O_{40}]^{n-}$ anions (Fig. 20A–C) begin to disintegrate with

decrease in acidity *via* interactions with hydroxyls. The stability towards the base hydrolysis increasing with the anion net charge as follows: $[P^VW_{12}^{VI}O_{40}]^{3-}$ $\{PW_{12}\}$ (pH < 1.5); $[Si^IVW_{12}^{VI}O_{40}]^{4-}$ $\{SiW_{12}\}$ (pH < 4.5); $[B^IIIW_{12}^{VI}O_{40}]^{5-}$ $\{BW_{12}\}$ and $[Al^IIIW_{12}^{VI}O_{40}]^{5-}$ $\{AlW_{12}\}$ (pH < 7); $[Be^IIW_{12}^{VI}O_{40}]^{6-}$ $\{BeW_{12}\}$ (pH < 9) (Fig. 22).^{1,234} Decomposition of the Keggin structure, like its formation, begins with disintegration of the $[XW_{12}^{VI}O_{40}]^{n-}$ anions by the removal of one $W=O^{4+}$ unit accompanied with the formation of a monolacunary derivative $[XW_{11}^{VI}O_{39}]^{(n-4)-}$ (Fig. 20E–H), followed by formation of di- or trilacunary anions (Fig. 20I–K).

$\{PW_{12}\}$. The hydrolytic conversions of $\{PW_{12}\}$ (Fig. 20A) have been studied for more than a century and most recently were thoroughly reviewed by Maksimovskaya and Maksimov.³⁸ The analysis of 95 previously published investigations made it



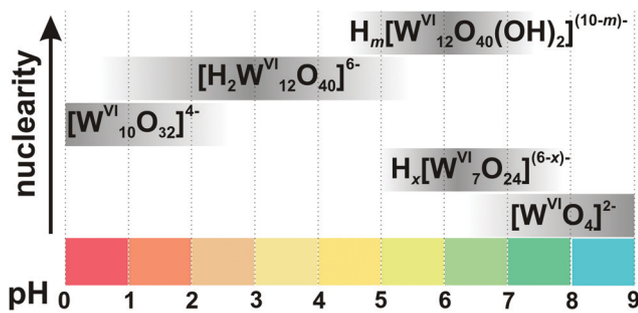


Fig. 19 Speciation of isopolytungstates in an aqueous solution with the concentration of 0.01–0.1 M W^{VI} based on works.^{132,215,217,218} The maximum intensity of grey color in each box with a single species corresponds to its maximum concentration in the chosen pH region. The grey boxes along the y-axis are positioned according to increasing nuclearity, but do not show the domination over other species at a certain pH range. x in $H_x[W^VI_7O_{24}]^{(6-x)-}$ is 0, 1; m in $H_m[W^VI_{12}O_{40}(OH)_2]^{(10-m)-}$ between 0 and 3. The structures of species are presented in Fig. 18. The ^{183}W and ^{17}O chemical shifts, stoichiometry, formation constants and pK_a values are summarized in Table 9.

possible to propose the most complete currently existing hydrolysis scheme for the “main” anion of POM chemistry (Fig. 21).³⁸

A few explanatory remarks need to be added to this scheme, while a detailed discussion is given in ref. 38. Systematical ^{31}P and ^{183}W NMR studies reveal that $\{PW_{12}\}$ exists in strongly acidic solutions with pH lower than 1.5 and converts to $[P^VW_{11}O_{39}M]^{6-}$ ($M = Li^+, Na^+, K^+$, Fig. 20D) through the intermediates $[H_nP^VW_{11}O_{39}]^{(7-n)-}$ (Fig. 20E), $n = 2-3$ and $[P^VW_{11}O_{39}(W^VI_2O_2)]^{5-}$ (Fig. 20D). In concentrated solutions ($[W^{VI}] > 0.01$ M) the latter $\{PW_{11}\}$ (Fig. 20E) anions slowly transform into several more stable species including $[P_2^VW_{21}O_{71}]^{6-}$ (Fig. 20N) and $[P_2^VW_{20}O_{70}]^{10-}$ (Fig. 20M).²³⁵ At pH higher than 7 three isomers of the trilacunary anion $A-\alpha$ -, $A-\beta$ -, $B-\alpha$ - $[P^VW_{19}O_{34}]^{9-}$ (see isostructural molybdates for α -isomers in Fig. 14E and F and β -forms Fig. 20J and K) and $[P_2^VW_{19}O_{69}]^{14-}$ –²³⁵ (Fig. 20L), an unstable derivative of $A-\alpha$ - $\{PW_9\}$, are present.

The gas-phase fragmentation of $\{PW_{12}\}$ in 50:50 acetonitrile– H_2O medium leads to pairs of product ions of general formulas $[W^VI_xO_{3x+1}]^{2-}$ and $[P^VW_{12-x}O_{39-3x}]^{(6-x)-}$ ($x = 6-9$), with the most prominent pair being $[W^VI_6O_{19}]^{2-}$ (see analog structure Fig. 8A) and $[P^VW_6O_{21}]^{(6-6)-}$.²³⁶ The formation and stability of the compact hexatungstate anion is not surprising in mixed water-organic solvent solution and was observed also during ESI-MS investigations of IPOTs solutions.¹⁹

$\{SiW_{12}\}$. The α -isomer of $[Si^IVW_{12}O_{40}]^{4-}$ (Fig. 20A) is stable in dilute solutions below pH 4.5, but the β - and γ -isomer (Fig. 20B and C) are metastable, slowly converting to the α -form (Fig. 23).¹ In their pioneering work on the hydrolysis of α - $[Si^IVW_{12}O_{40}]^{4-}$ ($\{SiW_{12}\}$), Kepert and Kyle reported that the reaction of silicotungstate with excess base proceeded via three distinct stages, with the intermediates being α - $[Si^IVW_{11}O_{39}]^{8-}$ (Fig. 20E) and $[Si^IVW_9O_{34}]^{10-}$ (see isostructural molybdates Fig. 14E and F), to finally produce $[Si^IVO_4]^{4-}$ and $[W^VI_4O_4]^{2-}$ (Fig. 23).²³⁷ ^{183}W NMR spectroscopic studies confirmed the formation of monolacunary species and its Na-substituted

analog α - $[NaSi^IVW_{11}O_{39}]^{7-}$ (Fig. 20D) in the pH range of 4–7.5 together with corresponding IPOTs $\{W_7\}$, $\{W_{12}O_{42}\}$, $\{W_{12}O_{40}\}$ (Fig. 19).^{238,239} The β - $\{SiW_{12}\}$ complex upon alkalization forms three short-lived isomers of the β - $[Si^IVW_{11}O_{39}]^{8-}$ anion (Fig. 20F–H),²³⁹ β_1 (C_s symmetry), β_2 (C_1), and β_3 (C_s), and the isomerization pathway is irreversible in solution and finally yields α - $[Si^IVW_{11}O_{39}]^{8-}$ (Fig. 20E).

Above pH 8, the trilacunary anions α - $[Si^IVW_9O_{34}]^{10-}$ (see isostructural molybdates Fig. 14E and F) and β - $[HSi^IVW_9O_{34}]^{9-}$ (Fig. 20J and K) originate from their monolacunary analogs. Fresh aqueous solutions of α - $[Si^IVW_9O_{34}]^{10-}$ (see isostructural molybdates Fig. 14E and F) between pH 9 and 10 are metastable, leading to monomeric silicate and tungstate (Fig. 23).²³⁸

γ - $[Si^IVW_{12}O_{40}]^{4-}$ (Fig. 20C) is stable only in non-aqueous or mixed water–ethanol or dioxane media. Upon dissolution of γ - $\{SiW_{12}\}$ in acidic aqueous media (pH < 4), only its α - and β -forms were detected.²⁴⁰ At pH between 4 and 7, the highly reactive²⁴¹ γ - $[HSi^IVW_{10}O_{36}]^{7-}$ (Fig. 20I) is present in freshly prepared solutions (Fig. 23). Then the β - and finally the α -isomer of $\{SiW_{11}\}$ are formed slowly by reaction of γ - $\{SiW_{10}\}$ with the IPOT species ($\{W_7\}$ or $\{W_{12}O_{42}\}$), which was shown by cyclic voltammetry²⁴⁰ and ESI-MS.²⁴² The polarogram of the γ - $\{SiW_{10}\}$ solution at pH between 4 and 6 shows one wave, which can be tentatively attributed to the intermediate species γ - $[H_xSi^IVW_{10}O_{39}]^{(8-x)-}$ in low concentration.²⁴⁰ Even though the γ -11-tungstosilicate polyanion cannot be obtained in a pure state, it has been identified as a subunit of a more complex structure $[H_2Si_2^IVW_{23}O_{78}]^{8-}$.²⁴⁰ Another labile γ lacunary complex has been determined by ESI-MS as an intermediate during the transformation from β - $\{SiW_{11}\}$ to γ - $\{SiW_{10}\}$.²⁴³ The unexpected labile precursor γ - $\{SiW_9\}$ is capable of a direct $\beta \rightarrow \gamma$ isomerization *via* a rotational transformation (Fig. 23).²⁴³

$\{AlW_{12}\}$. Weinstock *et al.* comprehensively investigated formation, isomerization, and derivatization of $[Al^{III}W_{12}O_{40}]^{5-}$ $\{AlW_{12}\}$ (Fig. 20A) using ^{27}Al and ^{183}W NMR spectroscopy, UV-vis and cyclic voltammetry.^{41,244} Contrary to $\{PW_{12}\}$ and $\{SiW_{12}\}$, β -forms of $\{AlW_{12}\}$ (Fig. 20B) and $\{AlW_{11}\}$ (Fig. 20F–H) are much more stable. Acidification of $[W^VI_4O_4]^{2-}$, followed by addition of Al^{III} at pH 7, leads to the formation of β - $[Al^{III}(Al^{III}OH_2)W_{11}O_{39}]^{6-}$ $\{Al_2W_{11}\}$, with the α derivative as a minor product with very low yield. Further acid addition to pH 0 and refluxing resulted in $H_5[Al^{III}W_{12}O_{40}]$ mostly in its β -form with the α -isomer as a minor product.²⁴⁴ Prolonged (5 days) heating of the β - $\{Al_2W_{11}\}$ solution at pH of 0–0.2 and 100 °C resulted in a mixture containing 92% α - $\{AlW_{12}\}$ (Fig. 20A) and 8% β - $\{Al_2W_{11}\}$ (Fig. 20B).⁴¹ The introductory data on pH stability of some sandwich-like HPOTs archetypes is described in (Table 8).

3.4.2.2. *The use of speciation data to understand the biological and catalytic activities of HPOTs with Keggin structure.* Apparently, in a physiological pH range as required for most biological investigations, the low charge clusters, $[P^VW_{12}O_{40}]^{3-}$ and $[Si^IVW_{12}O_{40}]^{4-}$ (Fig. 20A), should not be considered as intact species which will undergo at least partial hydrolysis. Due to their good synthetic accessibility they are often used for



biological applications without performing careful speciation studies under the applied physiological buffer conditions. A rational approach for the use of Keggin type POTs under physiological conditions has been demonstrated by our group.^{245,246} The combination of ¹²Be, ²⁷Al, ²⁹Si, ³¹P and ¹⁸³W NMR spectroscopies allowed to determine that the enzymatic inhibition behavior of the α -Keggin POTs against *Ab*PP04^{247,248} in citric buffer at pH 6.8 is charge-dependent, revealing significant inhibitory effects for POTs with lower net charge ($[P^V W_{12}^{VI} O_{40}]^{3-}$ and $[Si^{IV} W_{12}^{VI} O_{40}]^{4-}$ (Fig. 20A)) counterbalanced by their pronounced hydrolysis. A recent theoretical study by Carbo and co-authors also demonstrates that Keggin anions with lower net charge have higher affinity towards proteins.²⁴⁹

The selective peptide bond hydrolysis for human serum albumin, oxidized insulin chain B, myoglobin and cytochrome *c* by a range of Ce(IV)-substituted and Zr(IV)-substituted Lindqvist, Keggin and Wells–Dawson POMs was investigated by Parac-Vogt's group along with speciation in solution detailed in her

recent review.²⁵⁰ For instance, a combination of low-temperature ³¹P DOSY NMR spectroscopy and theoretical calculations allowed authors to conclude that in an aqueous solution with pD 6.4 the Keggin-type dimer $[\{\alpha-P^V W_{11}^{VI} Zr^{IV}(\mu-OH)(H_2O)\}_2]^{8-}$ used as enzyme mimetic for the hydrolysis of the phosphoester bonds in the DNA model substrate BNPP is present in equilibrium with its monomeric form.²⁵¹ In the proposed reaction mechanism, BNPP initially coordinates to monomeric $\{Zr^{IV} P^V W_{11}^{VI}\}$ in a monodentate fashion, which results in hydrolysis of the first phosphoester bond in BNPP and formation of nitrophenyl phosphate.

Many reactions applying POM catalysts are conducted in pure water,⁷ and so the hydrolytic stability must be taken into account. A good example is given by Neumann and co-authors investigating a sandwich Krebs type HPOT $Na_{12}[W^{VI} Zn^{II} Zn^{II}(H_2O)_2(Zn^{II} W_9^{VI} O_{34})_2]$ (two trilacunary Keggin fragments connected *via* "belt" of Zn and W ions) as a catalyst for selective alcohol oxidation with hydrogen peroxide in water–alcohol reaction media.^{252,253} Using ¹⁸³W NMR under reaction conditions (1 mmol of substrate (various alcohols),

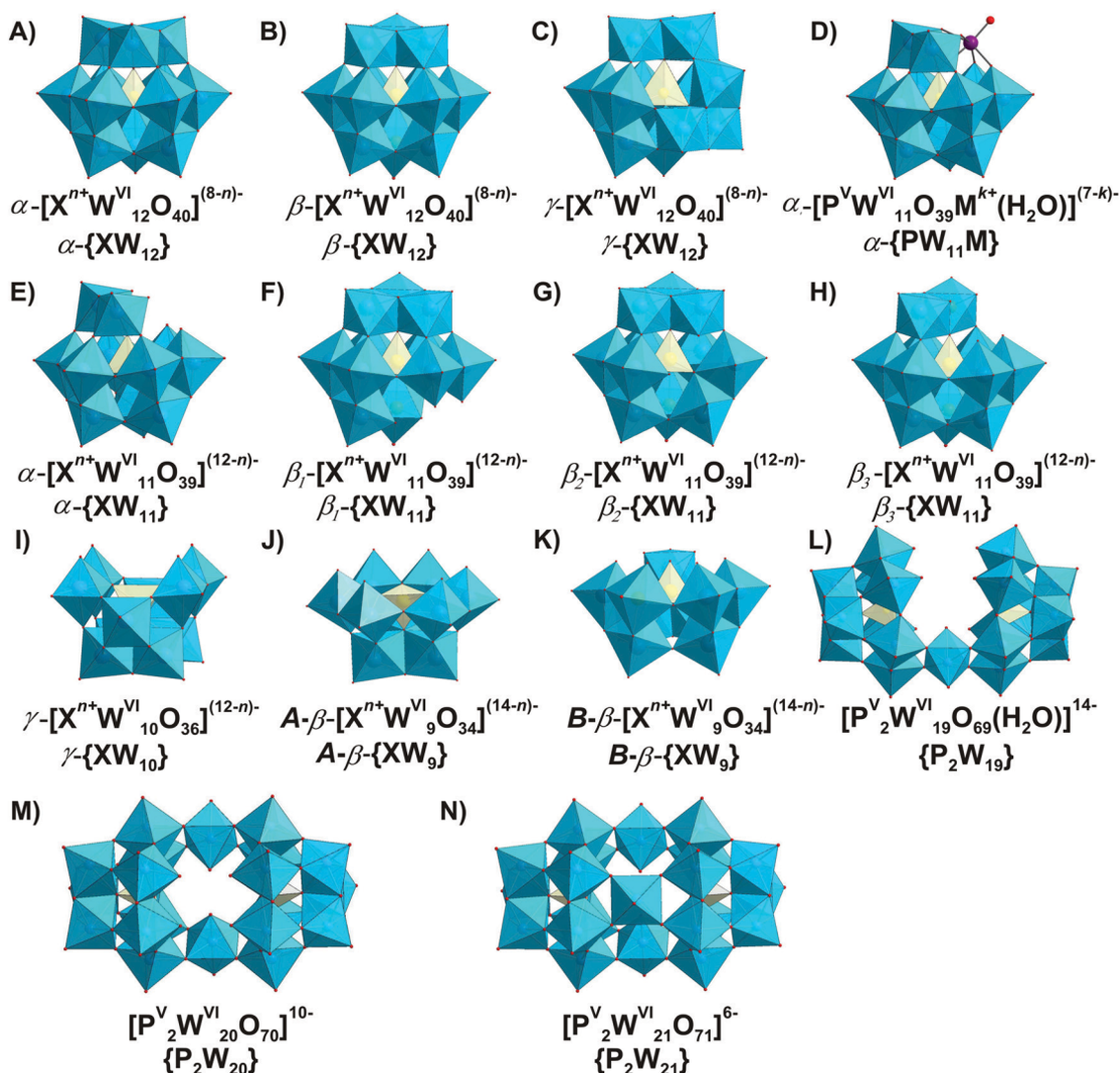


Fig. 20 HPOTs with Keggin ($[X^{n+}W_{12}^{VI}O_{40}]^{(8-n)-}$, where $X = P^V, Si^{IV}, Al^{III}$ etc.) and lacunary Keggin structures, which can be present and/or isolated from aqueous media ((A)–(N)). In (D) $[P^V W_{11}^{VI} O_{39} M^{k+} (H_2O)]^{(7-k)-}$ M^{k+} is Li^+, Na^+, K^+ , or $[W^{VI}O_2]^{2+}$. Color code: $\{WO_6\}$, blue; $\{XO_4\}$, yellow; M, purple; O, red.



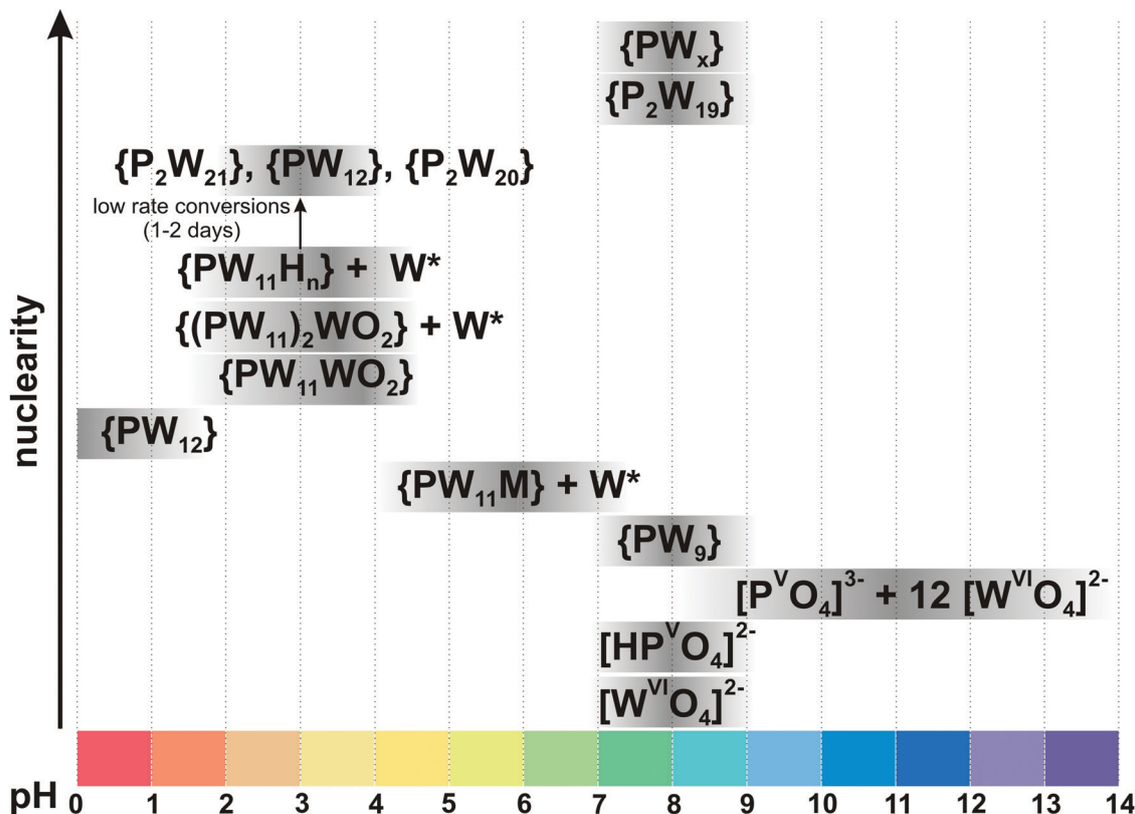


Fig. 21 Hydrolytic conversions of $\text{H}_3[\text{P}^{\text{V}}\text{W}_{12}^{\text{VI}}\text{O}_{40}]$ ($c = 0.01\text{--}0.1\text{ M}$, $[\text{W}]/[\text{P}] = 12$) alkalinized by MOH ($\text{M} = \text{Li}^+, \text{Na}^+, \text{K}^+$).³⁸ The maximum intensity of grey color in each box with a single species corresponds to its maximum concentration in the chosen pH region. The grey boxes along the y-axis are positioned according to increasing nuclearity, but do not show the domination over other species at a certain pH range. The abbreviation $\{\text{PW}_{11}\text{M}\}$ corresponds to the formula $[\text{P}^{\text{V}}\text{W}_{11}^{\text{VI}}\text{O}_{39}\text{M}]^{6-}$, where $\text{M} = \text{Li}^+, \text{K}^+, \text{Na}^+$; $\text{W}^* = [\text{H}_2\text{W}_{12}^{\text{VI}}\text{O}_{40}]^{6-}$ or $[\text{W}_{12}^{\text{VI}}\text{O}_{40}(\text{OH})_2]^{10-}$; $\{\text{PW}_{11}\text{WO}_2\} = [\text{P}^{\text{V}}\text{W}_{11}^{\text{VI}}\text{O}_{39}(\text{W}^{\text{VI}}\text{O}_2)]^{5-}$; $\{(\text{PW}_{11})_2\text{WO}_2\} = [(\text{P}^{\text{V}}\text{W}_{11}^{\text{VI}}\text{O}_{39})_2\text{W}^{\text{VI}}\text{O}_2]^{12-}$; $\{\text{H}_n\text{PW}_{11}\} = [\text{H}_n\text{P}^{\text{V}}\text{W}_{11}^{\text{VI}}\text{O}_{39}]^{(7-n)-}$, $n = 2\text{--}3$; $\{\text{P}_2\text{W}_{21}\} = [\text{P}_2^{\text{V}}\text{W}_{21}^{\text{VI}}\text{O}_{71}]^{6-}$; $\{\text{P}_2\text{W}_{20}\} = [\text{P}_2^{\text{V}}\text{W}_{20}^{\text{VI}}\text{O}_{70}]^{10-}$; $\{\text{PW}_9\} = \text{A-}\alpha\text{-, A-}\beta\text{-, B-}\alpha\text{-}[\text{P}^{\text{V}}\text{W}_9^{\text{VI}}\text{O}_{34}]^{9-}$; $\{\text{P}_2\text{W}_{19}\} = [\text{P}_2^{\text{V}}\text{W}_{19}^{\text{VI}}\text{O}_{69}(\text{H}_2\text{O})]^{14-}$; $\{\text{PW}_x\}$ – unidentified intermediate species. The structures of species are presented in Fig. 14 and 20. The ^{31}P chemical shifts and stoichiometry are summarized in ref. 38, and no formation constants have been reported for phosphotungstates.

5 mmol of H_2O_2 , 1 mL of water, $85\text{ }^\circ\text{C}$) the presence of intact anion was proved. The performed speciation analysis allowed the authors to determine the active POT species present under the chosen conditions and to suggest a catalytic mechanism.

Supporting the results of solution speciation using theoretical calculations can significantly improve understanding the reaction mechanism. Thus, the results of ^{31}P -NMR-monitoring of mixed-metal Keggin $[\gamma\text{-P}^{\text{V}}\text{V}_2\text{W}_{10}^{\text{VI}}\text{O}_{40}]^{5-}$ protonation in CH_3CN together with DFT calculations of the oxygen transfer process during the aromatic hydroxylation reaction allowed to

determine a peroxo complex $[\gamma\text{-P}^{\text{V}}\text{V}_2\text{W}_{10}^{\text{VI}}\text{O}_{38}(\text{O}_2)]^{3-}$ as the most active form of the catalyst.²⁵⁴

To illustrate a thorough speciation analysis of POM catalyst during water oxidation, six spectroscopic, scattering, and chemical experiments were used to indicate the stability of $[\{\text{Ru}_4^{\text{IV}}\text{O}_4(\text{OH})_2(\text{H}_2\text{O})_4\}(\gamma\text{-Si}^{\text{IV}}\text{W}_{10}^{\text{VI}}\text{O}_{36})_2]^{10-}$ in solution and under catalytic turnover conditions.²⁵⁵ UV-vis, cyclic voltammetry, ^{183}W and ^{29}Si NMR spectroscopy confirmed the stability of anion in aqueous solution at pH 3–4 up to 5 months, while SAXS and DLS in photocatalysis solution ($\text{Ru}(\text{bipy})_3^{2+}$, 0.5–1.0 mM; $\text{Na}_2\text{S}_2\text{O}_8$, 2.5–10 mM; $[\{\text{Ru}_4^{\text{IV}}\text{O}_4(\text{OH})_2(\text{H}_2\text{O})_4\}(\gamma\text{-Si}^{\text{IV}}\text{W}_{10}^{\text{VI}}\text{O}_{36})_2]^{10-}$, 1.25–10 μM ; sodium phosphate buffer, 20–50 mM; pH 7.2) did not show any particles indicative of RuO_2 or other possible decomposition products. Here we emphasize once again that a combination of different methods is the best strategy to get a broad picture of the POM solution behavior in aqueous media.

3.4.2.3. Hydrolytic stability of HPOTs with Wells–Dawson structure. Some aspects of the hydrolytic stability of $\{\text{X}_2\text{W}_{18}\}$ (Fig. 24) are highlighted in a general review about compounds with Wells–Dawson structure.²⁵⁶ The most common representative of Wells–Dawson HPOTs, $[\text{P}_2\text{W}_{18}^{\text{VI}}\text{O}_{62}]^{6-}$ ($\{\text{P}_2\text{W}_{18}\}$, Fig. 24A),

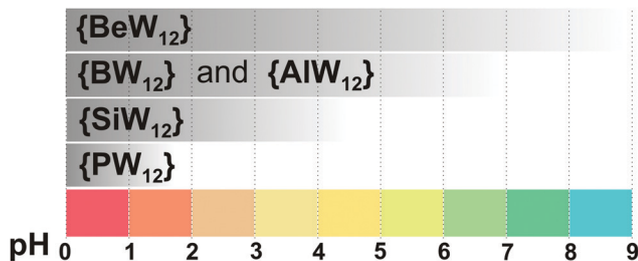


Fig. 22 pH stability regions of intact α -Keggin anions $\{\text{XW}_{12}\}$ with selected heteroatoms.



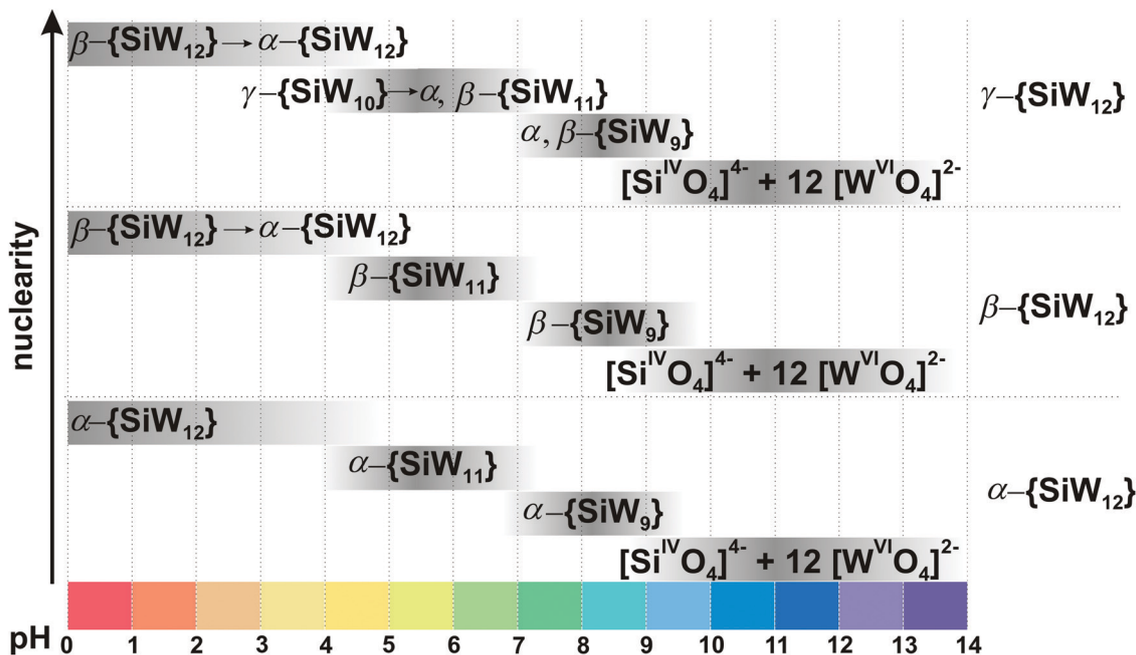


Fig. 23 Hydrolytic conversions of α -, β - and γ -[Si^{IV}W₁₂O₄₀]⁴⁻ ([SiW₁₂], [W] = 1.5 × 10⁻⁵–0.1 M, [W]/[Si] = 12) alkalinized by NaOH based on works.^{237–240} The maximum intensity of grey color in each box with a single species corresponds to its maximum concentration in the chosen pH region. The grey boxes along the y-axis are positioned according to increasing nucleicity, but do not show the domination over other species at a certain pH range. The abbreviation α -[SiW₁₁] corresponds to the mixture of α -[Si^{IV}W₁₁O₃₉Na]⁷⁻ and α -[Si^{IV}W₁₁O₃₉]⁸⁻; β -[SiW₁₁] – to the mixture of three isomers β_1 -, β_2 -, β_3 -[Si^{IV}W₁₁O₃₉]⁸⁻; α -[SiW₉] – α -[Si^{IV}W₉O₃₄]¹⁰⁻; β -[SiW₉] – β -[HSi^{IV}W₉O₃₄]⁹⁻; γ -[SiW₁₀] – γ -[HSi^{IV}W₁₀O₃₆]⁷⁻. The time-dependent conversions of β and γ forms to more stable α isomers are shown by arrows. The structures of species are presented in Fig. 14 and 20. No formation constants have been reported for silicotungstates.

shows stability at pH lower than 6, and its alkaline degradation leads to the formation of three different lacunary species: [P₂W₁₇O₆₁]¹⁰⁻ ({P₂W₁₇}, Fig. 24B), [P₂W₁₅O₅₆]¹²⁻ ({P₂W₁₅}, Fig. 24C) and H₂[P₂W₁₂O₄₈]¹²⁻ ({P₂W₁₂}, Fig. 24D).²⁵⁷ The {P₂W₁₂} species can further assemble to a very stable crown-like POT [H₇P₈W₄₈O₁₈₄]³³⁻, which is stable from pH 1 to 8. Since no detailed investigation of speciation was reported for {P₂W₁₈}, the hydrolytic conversion is presented as a scheme without exact pH ranges (Scheme 1).

Contant and Thouvenot studied the isomerism and stability of Wells–Dawson tungstates with P^V and As^V as heteroions.²⁵⁸ As for

the Keggin anions, β - and γ -isomers have structures in which one and two {W₃} groups, respectively, have been rotated by 60°. Structures in which the central {W₆} belts are staggered rather than eclipsed are denoted with an asterisk, and out of those, only the γ^* -isomer has been confirmed.¹ Spectroscopic (IR, Raman, UV-vis, ¹⁸³W and ³¹P NMR), polarographic and kinetic measurements showed that the α isomers are the only species stable in aqueous solution and all other forms isomerize to it.²⁵⁸ All isomers of {X₂W₁₈} (X = P^V and As^V) under alkaline hydrolysis firstly form monolacunary compounds. The rate of hydrolysis follows the trend for counteractions K⁺ > Na⁺ > Li⁺ and for isomers $\alpha < \beta < \gamma < \gamma^*$.

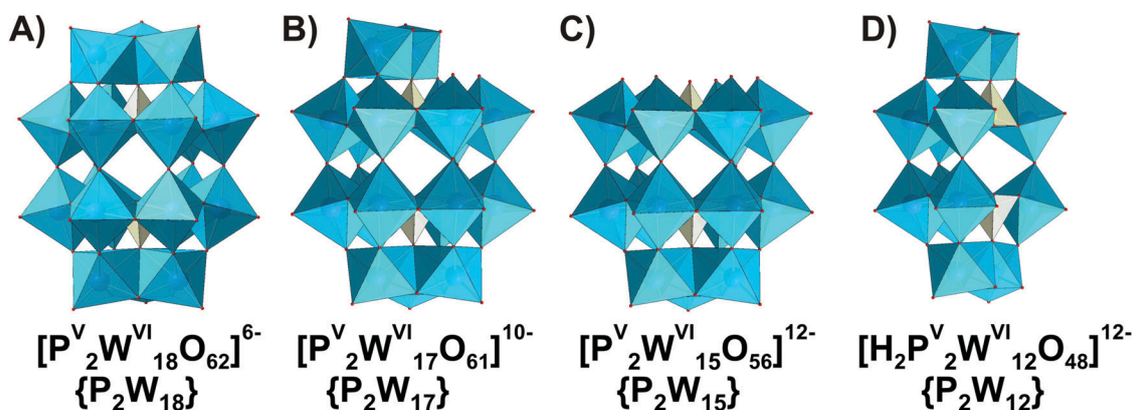
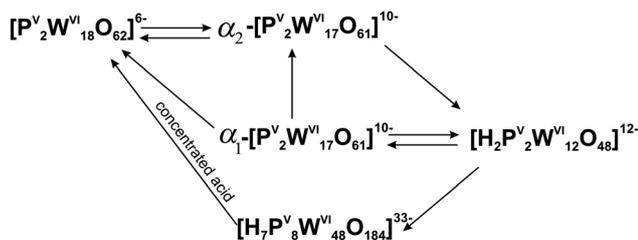


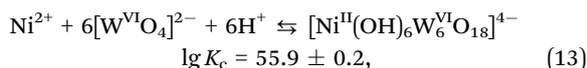
Fig. 24 Phosphotungstates with intact (A) and lacunary (B–D) α -Wells–Dawson structure. Color code: {WO₆}, blue; {PO₄}, yellow; O, red.



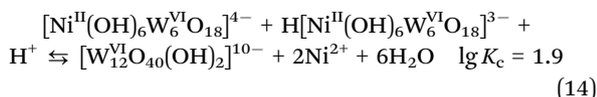
Scheme 1 Hydrolytic transformation of $\{P_2W_{18}\}$.²⁵⁷ The structures of species are shown in Fig. 24.

3.4.2.4. *The use of speciation data to understand the biological activities of HPOTs with Dawson structure.* The ability of $\{P_2W_{18}\}$ (Fig. 24A), $\{P_2W_{12}\}$ (Fig. 24D) and $\{P_2W_{15}\}$ (Fig. 24C) to affect aquaporin-3 (AQP3) activity and impair melanoma cell migration has been recently tested.²⁵⁹ The careful speciation investigation of respective POT clusters at physiological pH 7.4 (phosphate-buffered saline medium; NaCl, 137 mmol L⁻¹; KCl, 2.7 mmol L⁻¹; Na₂HPO₄, 10 mmol L⁻¹; KH₂PO₄, 1.8 mmol L⁻¹) and in water by ³¹P NMR analysis to address cluster species responsible for interaction in solution was performed. Solutions of the most promising AQP3 inhibitor, $\{P_2W_{18}\}$, contain intact Wells–Dawson anions, along with the mono-lacunary $\{P_2W_{17}\}$, while in the spectra of $\{P_2W_{12}\}$ and $\{P_2W_{15}\}$ only mono-lacunary species are present in PBS buffer. Although more investigations are needed to establish structure–activity relationships, it can be assumed that it is the intact phosphotungstate $\{P_2W_{18}\}$ that is responsible for the observed inhibitory effect on AQP3 activity and melanoma cell migration.

3.4.2.5. *Hydrolytic stability of HPOTs with Anderson structure.* Like their molybdate analogues, $\{XW_6\}$ Anderson archetype (see Fig. 16A for $\{XMo_6\}$ analog) compounds can be synthesized from slightly acidified solutions and are stable in near neutral pH range.^{1,201} However, the data on Anderson HPOTs stability is very limited and only systems with Ni^{II}, Cr^{III} and Te^{VI} have been investigated. The potentiometric studies show that $[Ni^{II}(OH)_6W_6^{VI}O_{18}]^{4-}$ ($\{NiW_6\}$) forms in 0.1 M NaNO₃ medium at 25 °C according to reaction (13) with formation constant:



After protonation of $\{NiW_6\}$ in acidified solutions it hydrolyzed to paratungstate B $\{W_{12}O_{42}\}$ (Fig. 18B) ($Z = n(H^+)/n([WO_4]^{2-}) = 1.17$) with the release of free Ni^{II} ions reaction (14):¹⁷



During the investigation of numerous successful co-crystallizations of $[Te^{VI}W_6^{VI}O_{24}]^{6-}$ ($\{TeW_6\}$) with different proteins, it was proven that $[Te^{VI}W_6^{VI}O_{24}]^{6-}$ is stable in a pH range from 4.5 to 7.5.^{260–263} In the presence of aurone synthase CgAUS²⁶⁴ $\{TeW_6\}$ was even functionalized by carboxylic fragment of protein glutamic acid on the μ_2 oxygen atoms, which proves the reactivity of Anderson POTs and demonstrates a

special kind of flexibility with respect to both geometric and functional properties.²⁶² All these examples show the importance of speciation analysis in physiological systems in the presence of biomacromolecules. The double-side trifunctionalized $[Cr^{III}((OCH_2)_3CC_2H_5)_2W_6^{VI}O_{18}]^{3-}$ is a predominant species in neutral pH between 5.7 and 7 almost without decomposition fragments, which was shown by ESI-MS.²⁶⁵

4. Conclusions

In the presented survey, the ion-distribution diagrams in a wide pH range (analog to “phase diagrams”) for POMs of V^V, Nb^V, Ta^V, Mo^{VI} and W^{VI} are given. They sum up our current understanding about POM stability and transformations in aqueous media. A brief timeline of POM speciation in aqueous media highlighting the first and the most important studies is given in Fig. S1 (ESI[†]). POM species in solution may be different to compounds that can be crystallized from the same solutions. It is advisable to use several orthogonal methods to understand and analyze the speciation. Sometimes, conclusions of the speciation differ according to the methods applied and the research method should be selected by taking into account the characteristics of a particular POM.

Throughout, this review factors that affect speciation have been shown. Among the obvious, such as pH (Section 3), ionic strength (Section 3), countercations (Section 3.2.1.3), buffer composition (Sections 3.2.1.3 and 3.4.1.4), aging time (Sections 3.3.1.6 and 3.3.2.1) and physical factors as temperature (Sections 3.2.1.3 and 3.3.2.1), attention should be paid to the presence of interfering substances as macromolecules and substrates of enzymes (Sections 3.1.1.4 and 3.4.2.5). Two intrinsic features of the POMs itself affect the stability in solution: charge density and anion symmetry (Sections 3.4.1.4 and 3.4.2.2). Stability increases with increasing charge density and symmetry of the POM anion. Strategies to stabilize POMs are: partial reduction or the replacement of addenda cation M^{VI} by a cation with lower charge.

We hope as an outcome of this review that special attention will be paid to investigation of POM behavior in solution that will facilitate the deeper understanding of their role in various applications. Reliable and accurate published data are of utmost importance for any type of POM project. For deliberate application of POMs, there is a great need for stability data obtained in non-aqueous solutions, water-organic solvent mixtures, or unusual experimental conditions (pressure, temperature, or at high concentrations), as well as in the presence of amphipathic molecules or other micelle-forming organic substances, and for complex formation processes on the surfaces of solid materials. Virtually nothing is known, about the possible role of enzymes or other physiologically occurring biological macromolecules, as well as their substrates with respect to enhancing the breakdown and/or rearrangements of POM species with some exception investigating decavanadate biomolecule interactions. We are confident that the new era of POM chemistry will be opened once the POM speciation



gets the attention it deserves and analyzing the behavior of the solution will become a common tool for understanding each POM application.

Conflicts of interest

There are no conflicts to declare.

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Notes and references

- M. T. Pope, *Heteropoly and Isopoly Oxometalates*, Springer-Verlag, New York, 1983.
- M. T. Pope and A. Müller, *Angew. Chem., Int. Ed. Engl.*, 1991, **30**, 34–48, DOI: 10.1002/anie.199100341.
- D.-L. Long, E. Burkholder and L. Cronin, *Chem. Soc. Rev.*, 2007, **36**, 105–121, DOI: 10.1039/B502666K.
- I. Knöpnadel, H. Hartl, W.-D. Hunnius and J. Fuchs, *Angew. Chem., Int. Ed. Engl.*, 1974, **13**, 823, DOI: 10.1002/anie.197408231.
- H.-U. Kreusler, A. Förster and J. Fuchs, *Z. Naturforsch., B: J. Chem. Sci.*, 1980, **35**, 242–244, DOI: 10.1515/znb-1980-0223.
- J. J. Cruywagen, A. G. Draaijer, J. B. B. Heyns and E. A. Rohwer, *Inorg. Chim. Acta*, 2002, **331**, 322–329, DOI: 10.1016/S0020-1693(02)00700-4.
- S.-S. Wang and G.-Y. Yang, *Chem. Rev.*, 2015, **11**, 4893–4962, DOI: 10.1021/cr500390v.
- A. Bijelic and A. Rompel, *Acc. Chem. Res.*, 2017, **50**, 1441–1448, DOI: 10.1021/acs.accounts.7b00109.
- A. Bijelic and A. Rompel, *ChemTexts*, 2018, **4**, 10, DOI: 10.1007/s40828-018-0064-1.
- A. Bijelic, M. Aureliano and A. Rompel, *Chem. Commun.*, 2018, **54**, 1153–1169, DOI: 10.1039/c7cc07549a.
- A. Bijelic, M. Aureliano and A. Rompel, *Angew. Chem., Int. Ed.*, 2019, **58**, 2980–2999 (*Angew. Chem.*, 2019, **131**, 3008–3029), DOI: 10.1002/anie.201803868, (DOI: 10.1002/ange.201803868).
- T. Kiss and A. Odani, *Bull. Chem. Soc. Jpn.*, 2007, **80**, 1691–1702, DOI: 10.1246/bcsj.80.1691.
- D. C. Crans, *Coord. Chem. Rev.*, 2017, **352**, 398–400, DOI: 10.1016/j.ccr.2017.11.008.
- D. M. Templeton, F. Ariese, R. Cornelis, L.-G. Danielsson, H. Muntau, H. P. van Leeuwen and R. Lobinski, *Pure Appl. Chem.*, 2000, **72**, 1453–1470, DOI: 10.1351/pac200072081453.
- T. Kiss, É. A. Enyedy and T. Jakuscha, *Coord. Chem. Rev.*, 2017, **352**, 401–423, DOI: 10.1016/j.ccr.2016.12.016.
- I. Ali and H. Y. Aboul-Enein, *Instrumental methods in metal ion speciation*, Taylor & Francis, Boca Raton, 2006, DOI: 10.1201/9781420019407.
- N. I. Gumerova, A. V. Notich, G. M. Rozantsev and S. V. Radio, *J. Solution Chem.*, 2016, **45**, 849–860, DOI: 10.1007/s10953-016-0471-0.
- R. I. Maksimovskaya and G. M. Maksimov, *Inorg. Chem.*, 2007, **46**, 3688–3695, DOI: 10.1021/ic061549n.
- L. Fan, J. Cao and C. Hu, *RSC Adv.*, 2015, **5**, 83377–83382, DOI: 10.1039/c5ra18059g.
- D. C. Crans, K. A. Woll, K. Prusinskas, M. D. Johnson and E. Norkus, *Inorg. Chem.*, 2013, **52**, 12262–12275, DOI: 10.1021/ic4007873.
- T. Yamase, *Chem. Rev.*, 1998, **98**, 307–325, DOI: 10.1021/cr9604043.
- H. So and M. T. Pope, *Inorg. Chem.*, 1972, **11**, 1441–1443, DOI: 10.1021/ic50112a061.
- G. M. Varga Jr, E. Papaconstantinou and M. T. Pope, *Inorg. Chem.*, 1970, **9**, 662–667, DOI: 10.1021/ic50085a045.
- P. Yang, Z. Lin, G. Alfaro-Espinoza, M. S. Ullrich, C. I. Rat, C. Silvestru and U. Kortz, *Inorg. Chem.*, 2016, **55**, 251–258, DOI: 10.1021/acs.inorgchem.5b02189.
- I. Holclajtner-Antunović, D. Bajuk-Bogdanović, M. Todorović, U. B. Mioć, J. Zakrzewska and S. Uskoković-Marković, *Can. J. Chem.*, 2008, **86**, 996–1004, DOI: 10.1139/V08-138.
- P. Häufe, *Fresenius' Z. Anal. Chem.*, 1982, **310**, 388–391, DOI: 10.1007/BF00483010.
- R. Q. Cabrera, S. Firth, C. S. Blackman, D.-L. Long, L. Cronin and P. F. McMillan, *J. Solid State Chem.*, 2012, **186**, 171–176, DOI: 10.1016/j.jssc.2011.12.005.
- A. F. Redkin and G. V. Bondarenko, *J. Solution Chem.*, 2010, **39**, 1549–1561, DOI: 10.1007/s10953-010-9595-9.
- M. Aureliano, C. A. Ohlin, M. O. Vieira, M. P. M. Marques, W. H. Casey and L. A. E. Batista de Carvalho, *Dalton Trans.*, 2016, **45**, 7391–7399, DOI: 10.1039/C5DT04176G.
- W. G. Klemperer and W. Shum, *J. Am. Chem. Soc.*, 1976, **98**, 8291–8293, DOI: 10.1021/ja00441a083.
- M. A. Fedotov and R. I. Maksimovskaya, *J. Struct. Chem.*, 2006, **47**, 952–978, DOI: 10.1007/s10947-006-0413-6.
- Y.-G. Chen, J. Gong and L.-Y. Qu, *Coord. Chem. Rev.*, 2004, **248**, 245–260, DOI: 10.1016/j.cct.2003.11.003.
- A. S. Tracey, G. R. Willsky and E. S. Takeuchi, *Vanadium: Chemistry, Biochemistry, Pharmacology and Practical Applications*, CRC Press, 2007, p. 200.
- D. C. Crans, P. K. Shin and K. B. Armstrong, *Adv. Chem.*, 1996, **246**, 303–328, DOI: 10.1021/ba-1995-0246.ch011.
- M. T. Pope and G. M. Varga, *J. Chem. Soc., Chem. Commun.*, 1966, 653–654, DOI: 10.1039/C19660000653.
- M. Filowitz, R. K. C. Ho, W. G. Klemperer and W. Shum, *Inorg. Chem.*, 1979, **18**, 93–103, DOI: 10.1021/ic50191a021.
- C. A. Ohlin and W. H. Casey, *Annu. Rep. NMR Spectrosc.*, 2018, **94**, 187–248, DOI: 10.1016/bs.arnmr.2018.01.001.
- R. I. Maksimovskaya and G. M. Maksimov, *Coord. Chem. Rev.*, 2019, **385**, 81–99, DOI: 10.1016/j.ccr.2019.01.014.
- R. I. Maksimovskaya and G. M. Maksimov, *Inorg. Chem.*, 2011, **50**, 4725–4731, DOI: 10.1021/ic101996f.
- M. Bugnola, R. E. Schreiber, Y. Kaufman, G. Leitus, L. J. W. Shimon and R. Neumann, *Eur. J. Inorg. Chem.*, 2019, 482–485, DOI: 10.1002/ejic.201800552.



- 41 J. J. Cowan, A. J. Bailey, R. A. Heintz, B. T. Do, K. I. Hardcastle, C. L. Hill and I. A. Weinstock, *Inorg. Chem.*, 2001, **40**, 6666–6675, DOI: 10.1021/ic0106120.
- 42 P. Klonowski, J. C. Goloboy, F. J. Uribe-Romo, F. Sun, L. Zhu, F. Gándara, C. Wills, R. J. Errington, O. M. Yaghi and W. G. Klemperer, *Inorg. Chem.*, 2014, **53**, 13239–13246, DOI: 10.1021/ic502617k.
- 43 A. Misra, K. Kozma, C. Streb and M. Nyman, *Angew. Chem., Int. Ed.*, 2020, **59**, 596–612, DOI: 10.1002/anie.201905600.
- 44 J. Xie, H. A. Neal, J. Szymanowski, P. C. Burns, T. M. Alam, M. Nyman and L. Gagliardi, *Inorg. Chem.*, 2018, **57**, 5514–5525, DOI: 10.1021/acs.inorgchem.8b00474.
- 45 M. Nyman, F. Bonhomme, T. M. Alam, J. B. Parise and G. M. B. Vaughan, *Angew. Chem., Int. Ed.*, 2004, **43**, 2787–2792, DOI: 10.1002/anie.200353410.
- 46 D. J. Sures, S. A. Serapian, K. Kozma, P. I. Molina, C. Bo and M. Nyman, *Phys. Chem. Chem. Phys.*, 2017, **19**, 8715–8725, DOI: 10.1039/C6CP08454K.
- 47 H. N. Miras, E. F. Wilson and L. Cronin, *Chem. Commun.*, 2009, 1297–1311, DOI: 10.1039/B819534J.
- 48 C. S. Truebenbach, M. Houalla and D. M. Hercules, *J. Mass Spectrom.*, 2000, **35**, 1121–1127, DOI: 10.1002/1096-9888(200009)35:9<1121::AID-JMS40>3.0.CO;2-7.
- 49 C. A. Ohlin, *Chem. – Asian J.*, 2012, **7**, 262–270, DOI: 10.1002/asia.201100763.
- 50 M. Nyman, *Coord. Chem. Rev.*, 2017, **352**, 461–472, DOI: 10.1016/j.ccr.2016.11.014.
- 51 M. Nyman and L. Fullmer, *Small angle X-ray scattering of group V polyoxometalates*, Trends in Polyoxometalates Research, Nova Science Publishers Inc., 2015, Book chapter pdf: ISBN 978-163482693-8; 978-163482656-3.
- 52 A. A. Shmakova, V. V. Volchek, V. Yanshole, N. B. Kompankov, N. P. Martin, M. Nyman, P. A. Abramov and M. N. Sokolov, *New J. Chem.*, 2019, **43**, 9943–9952, DOI: 10.1039/C9NJ01907C.
- 53 M. Dufaye, S. Duval, G. Stoclet and T. Loiseau, *Eur. J. Inorg. Chem.*, 2019, 4500–4505, DOI: 10.1002/ejic.201900732.
- 54 P. Yin, B. Wu, E. Mamontov, L. L. Daemen, Y. Cheng, T. Li, S. Seifert, K. Hong, P. V. Bonnesen, J. K. Keum and A. J. Ramirez-Cuesta, *J. Am. Chem. Soc.*, 2016, **138**, 2638–2643, DOI: 10.1021/jacs.5b11465.
- 55 P. Yin, T. Li, R. S. Forgan, C. Lydon, X. Zuo, Z. N. Zheng, B. Lee, D. Long, L. Cronin and T. Liu, *J. Am. Chem. Soc.*, 2013, **135**, 13425–13432, DOI: 10.1021/ja404777g.
- 56 S. Goberna-Ferrón, J. Soriano-López, J. R. Galán-Mascarós and M. Nyman, *Eur. J. Inorg. Chem.*, 2015, 2833–2840, DOI: 10.1002/ejic.201500404.
- 57 N. I. Gumerova and A. Rompel, *Nat. Rev. Chem.*, 2018, **2**, 0112, DOI: 10.1038/s41570-018-0112.
- 58 M. Sadakane and E. Steckhan, *Chem. Rev.*, 1998, **98**, 219–237, DOI: 10.1021/cr960403a.
- 59 T. Ueda, *ChemElectroChem*, 2018, **5**, 823–838, DOI: 10.1002/celec.201701170.
- 60 G. Bunker, *Introduction to XAFS*, Cambridge University Press, Cambridge, 2010.
- 61 M. P. M. Marques, D. Gianolio, S. Ramos, L. A. E. Batista de Carvalho and M. Aureliano, *Inorg. Chem.*, 2017, **56**, 10893–10903, DOI: 10.1021/acs.inorgchem.7b01018.
- 62 B. C. Bostick, J. Sun, J. D. Landis and J. L. Clausen, *Environ. Sci. Technol.*, 2018, **52**, 1045–1053, DOI: 10.1021/acs.est.7b05406.
- 63 S. Bhattacharjee, *J. Controlled Release*, 2016, **235**, 337–351, DOI: 10.1016/j.jconrel.2016.06.017.
- 64 J. J. Stracke and R. G. Finke, *ACS Catal.*, 2014, **4**, 909–933, DOI: 10.1021/cs4011716.
- 65 E. E. Hamilton, P. E. Fanwick and J. J. Wilker, *J. Am. Chem. Soc.*, 2002, **124**, 78–82, DOI: 10.1021/ja010820r.
- 66 J. Fuchs, S. Mahjour and J. Pickardt, *Angew. Chem., Int. Ed. Engl.*, 1976, **15**, 374–375, DOI: 10.1002/anie.197603741.
- 67 V. W. Day, W. G. Klemperer and O. M. Yaghi, *J. Am. Chem. Soc.*, 1989, **111**, 4518–4519, DOI: 10.1021/ja00194a068.
- 68 P. Y. Zavalij, T. Chirayil, M. S. Whittingham, V. K. Pecharsky and R. Jacobson, *Acta Crystallogr., Sect. C: Cryst. Struct. Commun.*, 1997, **53**, 170–171, DOI: 10.1107/S0108270196013480.
- 69 V. W. Day, W. G. Klemperer and O. M. Yaghi, *J. Am. Chem. Soc.*, 1989, **111**, 5959–5961, DOI: 10.1021/ja00197a077.
- 70 D. Hou, K. S. Hagen and C. L. Hill, *J. Am. Chem. Soc.*, 1992, **114**, 5864–5866, DOI: 10.1021/ja00040a061.
- 71 D. Hou, K. S. Hagen and C. L. Hill, *J. Chem. Soc., Chem. Commun.*, 1993, 426–428, DOI: 10.1039/C39930000426.
- 72 D. C. Crans, *Pure Appl. Chem.*, 2005, **77**, 1497–1527, DOI: 10.1351/pac200577091497.
- 73 L. Pettersson, *Mol. Eng.*, 1993, **3**, 29–42, DOI: 10.1007/bf00999622.
- 74 D. K. Walanda, R. C. Burns, G. A. Lawrance and E. I. von Nagy-Felsobuki, *Inorg. Chem. Commun.*, 1999, **2**, 487–489, DOI: 10.1016/S1387-7003(99)00129-X.
- 75 D. K. Walanda, R. C. Burns, G. A. Lawrance and E. I. von Nagy-Felsobuki, *Inorg. Chim. Acta*, 2000, **305**, 118–126, DOI: 10.1016/S0020-1693(00)00122-5.
- 76 D. C. Crans, C. D. Rithner and L. A. Theisen, *J. Am. Chem. Soc.*, 1990, **112**, 2901–2908, DOI: 10.1021/ja00164a009.
- 77 I. Andersson, L. Pettersson, J. J. Hastings and O. W. Howarth, *J. Chem. Soc., Dalton Trans.*, 1996, 3357–3361, DOI: 10.1039/DT9960003357.
- 78 A. S. Tracey, J. S. Jaswal and S. J. Angus-Dunne, *Inorg. Chem.*, 1995, **34**, 5680–5685, DOI: 10.1021/ic00126a043.
- 79 K. Elvingson, A. González Baró and L. Pettersson, *Inorg. Chem.*, 1996, **35**, 3388–3393, DOI: 10.1021/ic951195s.
- 80 D. C. Crans, B. Baruah and N. E. Levinger, *Biomed. Pharmacother.*, 2006, **60**, 174–181, DOI: 10.1016/j.biopha.2006.02.006.
- 81 R. Gopal and C. Calvo, *Acta Crystallogr., Sect. B: Struct. Crystallogr. Cryst. Chem.*, 1974, **30**, 2491–2493, DOI: 10.1107/S0567740874007400.
- 82 S. Nakamura and T. Ozeki, *J. Chem. Soc., Dalton Trans.*, 2001, 472–480, DOI: 10.1039/B008128K.
- 83 V. W. Day, W. G. Klemperer and D. J. Maltbie, *J. Am. Chem. Soc.*, 1987, **109**, 2991–3002, DOI: 10.1021/ja00244a022.
- 84 *Vanadium and its role for life; metal ions in biological systems*, ed. A. Sigel and H. Sigel, New York, Marcel Dekker, 1995, DOI: 10.1016/0307-4412(95)90238-4.



- 85 N. McCann, M. Wagner and H. Hasse, *Dalton Trans.*, 2013, **42**, 2622–2628, DOI: 10.1039/c2dt31993d.
- 86 J. W. Larson, *J. Chem. Eng. Data*, 1995, **40**, 1276–1280, DOI: 10.1021/je00022a030.
- 87 E. Heath and O. W. Howarth, *J. Chem. Soc., Dalton Trans.*, 1981, 1105–1110, DOI: 10.1039/DT9810001105.
- 88 D. C. Crans, B. J. Peters, X. Wu and C. C. McLauchlan, *Coord. Chem. Rev.*, 2017, **344**, 115–130, DOI: 10.1016/j.ccr.2017.03.016.
- 89 O. W. Howarth and M. Jarrold, *J. Chem. Soc., Dalton Trans.*, 1978, 503–506, DOI: 10.1039/DT9780000503.
- 90 M. Aureliano and D. C. Crans, *J. Inorg. Biochem.*, 2009, **103**, 536–546, DOI: 10.1016/j.jinorgbio.2008.11.010.
- 91 S. S. Soares, F. Henao, M. Aureliano and C. Gutiérrez-Merino, *Chem. Res. Toxicol.*, 2008, **21**, 607–618, DOI: 10.1021/tx700204r.
- 92 R. M. C. Gândara, S. S. Soares, H. Martins, C. Gutiérrez-Merino and M. Aureliano, *J. Inorg. Biochem.*, 2005, **99**, 1238–1244, DOI: 10.1016/j.jinorgbio.2005.02.023.
- 93 T. L. Turner, V. H. Nguyen, C. C. McLauchlan, Z. Dymon, B. M. Dorsey, J. D. Hooker and M. A. Jones, *J. Inorg. Biochem.*, 2012, **108**, 96–104, DOI: 10.1016/j.jinorgbio.2011.09.009.
- 94 P. Csermely, A. Martonosi, G. C. Levy and A. J. Ejchart, *Biochem. J.*, 1985, **230**, 807–815, DOI: 10.1042/bj2300807.
- 95 S. S. Soares, C. Gutiérrez-Merino and M. Aureliano, *J. Inorg. Biochem.*, 2007, **101**, 789–796, DOI: 10.1016/j.jinorgbio.2007.01.012.
- 96 M. Aureliano, *Dalton Trans.*, 2009, 9093–9100, DOI: 10.1039/b907581j.
- 97 S. Ramos, M. Manuel, T. Tiago, R. Duarte, J. Martins, C. Gutiérrez-Merino, J. J. G. Moura and M. Aureliano, *J. Inorg. Biochem.*, 2006, **100**, 1734–1743, DOI: 10.1016/j.jinorgbio.2006.06.007.
- 98 M. Aureliano, G. Fraqueza and C. A. Ohlin, *Dalton Trans.*, 2013, **42**, 11770–11777, DOI: 10.1039/c3dt50462j.
- 99 D. C. Crans, J. J. Smee, E. Gaidamauskas and L. Yang, *Chem. Rev.*, 2004, **104**, 849–902, DOI: 10.1021/cr020607t.
- 100 A. Al-Qatati, F. L. Fontes, B. G. Barisas, D. Zhang, D. A. Roess and D. C. Crans, *Dalton Trans.*, 2013, **42**, 11912–11920, DOI: 10.1039/C3DT50398D.
- 101 N. Samart, Z. Arhouma, S. Kumar, H. A. Murakami, D. C. Crick and D. C. Crans, *Front. Chem.*, 2018, **6**, 519, DOI: 10.3389/fchem.2018.00519.
- 102 N. Fukuda and T. Yamase, *Biol. Pharm. Bull.*, 1997, **20**, 927–930, DOI: 10.1248/bpb.20.927.
- 103 J. M. Messmore and R. T. Raines, *Arch. Biochem. Biophys.*, 2000, **381**, 25–30, DOI: 10.1006/abbi.2000.1951.
- 104 A. Selling, I. Andersson, L. Pettersson, C. M. Schramm, S. L. Downey and J. H. Grate, *Inorg. Chem.*, 1994, **33**, 3141–3150, DOI: 10.1021/ic00092a021.
- 105 I. Andersson, A. Gorzsás, C. Kerezsi, I. Tóth and L. Pettersson, *Dalton Trans.*, 2005, 3658–3666, DOI: 10.1039/B508273K.
- 106 W. Zhang, T. Zhang, G. Lv, X. Cao and H. Zhu, *Sep. Purif. Technol.*, 2019, **218**, 164–172, DOI: 10.1016/j.seppur.2019.02.025.
- 107 R. Kato, A. Kobayashi and Y. Sasaki, *Inorg. Chem.*, 1982, **21**, 240–246, DOI: 10.1021/ic00131a045.
- 108 K. Postal, D. F. Maluf, G. Valdameri, A. L. Rüdiger, D. L. Hughes, E. L. de Sá, R. R. Ribeiro, E. M. de Souza, J. F. Soares and G. G. Nunes, *RSC Adv.*, 2016, **6**, 114955–114968, DOI: 10.1039/c6ra15826a.
- 109 M. Nyman, *Dalton Trans.*, 2011, **40**, 8049–8058, DOI: 10.1039/c1dt10435g.
- 110 I. Lindqvist, *Ark. Kemi*, 1953, **5**, 247–250.
- 111 E. J. Graeber and B. Morosin, *Acta Crystallogr., Sect. B: Struct. Crystallogr. Cryst. Chem.*, 1977, **33**, 2137–2143, DOI: 10.1107/S0567740877007900.
- 112 C. A. Ohlin, E. M. Villa and W. H. Casey, *Inorg. Chim. Acta*, 2009, **362**, 1391–1392, DOI: 10.1016/j.ica.2008.06.006.
- 113 G. M. Rozantsev, O. I. Dotsenko and G. V. Taradina, *Russ. J. Coord. Chem.*, 2000, **26**, 247–253.
- 114 E. M. Villa, C. A. Ohlin, E. Balogh, T. M. Anderson, M. D. Nyman and W. H. Casey, *Angew. Chem., Int. Ed.*, 2008, **47**, 4844–4846, DOI: 10.1002/anie.200801125.
- 115 W. G. Klemperer and K. A. Marek, *Eur. J. Inorg. Chem.*, 2013, 1762–1771, DOI: 10.1002/ejic.201201231.
- 116 M. R. Antonio, M. Nyman and T. M. Anderson, *Angew. Chem., Int. Ed.*, 2009, **48**, 6136–6140, DOI: 10.1002/anie.200805323.
- 117 M. Matsumoto, Y. Ozawa, A. Yagasaki and Y. Zhe, *Inorg. Chem.*, 2013, **52**, 7825–7827, DOI: 10.1021/ic400864e.
- 118 E. Balogh, T. M. Anderson, J. R. Rustad, M. Nyman and W. H. Casey, *Inorg. Chem.*, 2007, **46**, 7032–7039, DOI: 10.1021/ic700845e.
- 119 N. Etxebarria, L. A. Fernández and J. M. Madariaga, *J. Chem. Soc., Dalton Trans.*, 1994, 3055–3059, DOI: 10.1039/DT9940003055.
- 120 A. Goiffon, E. Philippot and M. Maurin, *Rev. Chim. Miner.*, 1980, **17**, 466–476.
- 121 M. Nyman, T. M. Alam, F. Bonhomme, M. A. Rodriguez, C. S. Frazer and M. E. Welk, *J. Cluster Sci.*, 2006, **17**, 197–219, DOI: 10.1007/s10876-006-0049-x.
- 122 C. A. Ohlin, E. M. Villa, J. C. Fettinger and W. H. Casey, *Angew. Chem., Int. Ed.*, 2008, **47**, 8251–8254, DOI: 10.1002/ange.200803688.
- 123 N. P. Martin, E. Petrus, M. Segado, A. Arteaga, L. N. Zakharov, C. Bo and M. Nyman, *Chem. – Eur. J.*, 2019, **25**, 10580–10584, DOI: 10.1002/chem.201902770.
- 124 P. Müscher-Polzin, C. Näther and W. Bensch, *Z. Anorg. Allg. Chem.*, 2020, **646**, 193–198, DOI: 10.1002/zaac.201900276.
- 125 R. P. Bontchev and M. Nyman, *Angew. Chem., Int. Ed.*, 2006, **45**, 6670–6672, DOI: 10.1002/anie.20060220.
- 126 J. R. Black, M. Nyman and W. H. Casey, *J. Am. Chem. Soc.*, 2006, **128**, 14712–14720, DOI: 10.1021/ja065529w.
- 127 D. Sures, M. Segado, C. Bo and M. Nyman, *J. Am. Chem. Soc.*, 2018, **140**, 10803–10813, DOI: 10.1021/jacs.8b05015.
- 128 K.-H. Tytko and O. Glemser, *Adv. Inorg. Chem. Radiochem.*, 1976, **19**, 239–315, DOI: 10.1016/S0065-2792(08)60073-4.
- 129 O. W. Howarth, P. Kelly and L. Patterson, *J. Chem. Soc., Dalton Trans.*, 1990, 81–84, DOI: 10.1039/dt9900000081.
- 130 M. Isobe, F. Marumo, T. Yamase and T. Ikawa, *Acta Crystallogr., Sect. B: Struct. Crystallogr. Cryst. Chem.*, 1978, **34**, 2728–2731, DOI: 10.1107/S0567740878009103.



- 131 J. Torres, L. Gonzatto, G. Peinado, C. Kremer and E. Kremer, *J. Solution Chem.*, 2014, **43**, 1687–1700, DOI: 10.1007/s10953-014-0231-y.
- 132 C. Falaise, M. A. Moussawi, S. Floquet, P. A. Abramov, M. N. Sokolov, M. Haouas and E. Cadot, *J. Am. Chem. Soc.*, 2018, **140**, 11198–11201, DOI: 10.1021/jacs.8b07525.
- 133 S. Himeno, H. Niiya and T. Ueda, *Bull. Chem. Soc. Jpn.*, 1997, **70**, 631–637, DOI: 10.1246/bcsj.70.631.
- 134 P. L. Brown, M. E. Shyinq and R. N. Sylva, *J. Chem. Soc., Dalton Trans.*, 1987, 2149–2157, DOI: 10.1039/DT9870002149.
- 135 K. H. Tytko, G. Baethe and J. J. Cruywagen, *Inorg. Chem.*, 1985, **24**, 3132–3136, DOI: 10.1021/ic00214a010.
- 136 F. Taube, I. Andersson and L. Pettersson, Molybdate speciation in systems related to the bleaching of kraft pulp, in *Polyoxometalate chemistry from topology via self-assembly to applications*, ed. M. T. Pope and A. Müller, Springer, Dordrecht, 2001, pp. 161–173, DOI: 10.1007/0-306-47625-8.
- 137 J. J. Cruywagen and J. B. B. Heyns, *Polyhedron*, 2000, **19**, 907–911, DOI: 10.1016/S0277-5387(00)00331-4.
- 138 I. Lindqvist, *Acta Crystallogr.*, 1950, **3**, 159–160, DOI: 10.1107/S0365110X50000355.
- 139 J. Fuchs and H. Hartl, *Angew. Chem., Int. Ed. Engl.*, 1976, **15**, 375–376, DOI: 10.1002/anie.197603751.
- 140 J. Fuchs and I. Knöpnadel, *Z. Kristallogr.*, 1982, **158**, 165–179, DOI: 10.1524/zkri.1982.158.12.165.
- 141 I. Paulat-Böschchen, *J. Chem. Soc., Chem. Commun.*, 1979, 780–782, DOI: 10.1039/C39790000780.
- 142 H. Schulz and F. A. Schröder, *Acta Crystallogr., Sect. A: Cryst. Phys., Diffr., Theor. Gen. Crystallogr.*, 1973, **29**, 322–333, DOI: 10.1107/S0567739473000902.
- 143 K. Y. S. Ng and E. Gulari, *Polyhedron*, 1984, **3**, 1001–1011, DOI: 10.1016/S0277-5387(00)84659-8.
- 144 A. Davantès and G. Lefèvre, *J. Phys. Chem. A*, 2013, **117**, 12922–12929, DOI: 10.1021/jp408885k.
- 145 E. Cartuyvels, G. Absillis and T. N. Parac-Vogt, *Chem. Commun.*, 2008, 85–87, DOI: 10.1039/b714860g.
- 146 G. Absillis, R. Van Deun and T. N. Parac-Vogt, *Inorg. Chem.*, 2011, **50**, 11552–11560, DOI: 10.1021/ic201498u.
- 147 S. Himeno, N. Ishii, M. Hasegawa, A. Saito and T. Hori, *Inorg. Chim. Acta*, 1987, **131**, 11–13, DOI: 10.1016/S0020-1693(00)87899-8.
- 148 K. Murata and S. Ikeda, *Spectrochim. Acta, Part A*, 1983, **39**, 787–794, DOI: 10.1016/0584-8539(83)80018-X.
- 149 S. Watanabe, T. Sato, M. Nakaya, T. Yoshida and J. Onoe, *Chem. Phys. Lett.*, 2019, **723**, 76–81, DOI: 10.1016/j.cplett.2019.02.049.
- 150 K. Yokoi, N. Matsubayashi, T. Miyanaga, I. Watanabe and S. Ikeda, *Polyhedron*, 1993, **12**, 911–914, DOI: 10.1016/S0277-5387(00)81545-4.
- 151 L. Vilà-Nadal, E. F. Wilson, H. N. Miras, A. Rodríguez-Fortea, L. Cronin and J. M. Poblet, *Inorg. Chem.*, 2011, **50**, 7811–7819, DOI: 10.1021/ic200969h.
- 152 M. Marianski, J. Seo, E. Mucha, D. A. Thomas, S. Jung, R. Schlögl, G. Meijer, A. Trunschke and G. von Helden, *J. Phys. Chem. C*, 2019, **123**, 7845–7853, DOI: 10.1021/acs.jpcc.8b06985.
- 153 F. Steffler, G. Ferreira de Lima and H. Anderson Duarte, *Chem. Phys. Lett.*, 2017, **669**, 104–109, DOI: 10.1016/j.cplett.2016.12.010.
- 154 B. M. Gatehouse and P. Leverett, *J. Chem. Soc. A*, 1971, 2107–2112, DOI: 10.1039/j19710002107.
- 155 B. M. Gatehouse and P. Leverett, *J. Chem. Soc. A*, 1968, 1398–1405, DOI: 10.1039/j19680001398.
- 156 P. J. Zapf, R. C. Haushalter and J. Zubietta, *Chem. Commun.*, 1997, 321–322, DOI: 10.1039/a607022A.
- 157 S.-F. Zhang, Y.-Q. Suna and G.-Y. Yang, *Acta Crystallogr., Sect. C: Cryst. Struct. Commun.*, 2004, **60**, m299–m301, DOI: 10.1107/S010827010401090X.
- 158 J. Noack, F. Rosowski, R. Schlögl and A. Trunschke, *Z. Anorg. Allg. Chem.*, 2014, **640**, 2730–2736, DOI: 10.1002/zaac.201400439.
- 159 V. Coué, R. Dessapt, M. Bujoli-Doeuff, M. Evain and S. Jobic, *Inorg. Chem.*, 2007, **46**, 2824–2835, DOI: 10.1021/ic0621502.
- 160 H.-J. Lunk and H. Hartl, *ChemTexts*, 2017, **3**, 13, DOI: 10.1007/s40828-017-0048-6.
- 161 A. Wutkowski, B. R. Srinivasan, A. R. Naik, C. Schütt, C. Näther and W. Bensch, *Eur. J. Inorg. Chem.*, 2011, 2254–2263, DOI: 10.1002/ejic.201001154.
- 162 I. Böschchen, B. Buss and B. Krebs, *Acta Crystallogr., Sect. B: Struct. Crystallogr. Cryst. Chem.*, 1974, **30**, 48–56, DOI: 10.1107/S0567740874002184.
- 163 W. T. A. Harrison, L. L. Dussack and A. J. Jacobson, *Acta Crystallogr., Sect. C: Cryst. Struct. Commun.*, 1996, **52**, 1075–1077, DOI: 10.1107/S0108270195015642.
- 164 S. Chakrabarti and S. Natarajan, *Cryst. Growth Des.*, 2002, **2**, 333–335, DOI: 10.1021/cg0200210.
- 165 J. Fuchs, H. Hartl, W.-D. Hunnius and S. Mahjour, *Angew. Chem., Int. Ed. Engl.*, 1975, **14**, 644, DOI: 10.1002/anie.197506441.
- 166 T. Arumuganathan, A. Srinivasarao, T. Vijay Kumar and S. K. Das, *J. Chem. Sci.*, 2008, **120**, 95–103, <https://www.ias.ac.in/article/fulltext/jcsc/120/01/0095-0103>.
- 167 L. Pettersson, I. Andersson and L.-O. Öhman, *Acta Chem. Scand., Ser. A*, 1985, **A39**, 53–58, DOI: 10.3891/acta.chem.scand.39a-0053.
- 168 L. Pettersson, I. Andersson and L.-O. Öhman, *Inorg. Chem.*, 1986, **25**, 4726–4733, DOI: 10.1021/ic00246a028.
- 169 R. Strandberg, *Acta Chem. Scand.*, 1973, **27**, 1004–1018, DOI: 10.3891/acta.chem.scand.27-1004.
- 170 B. Hedman and R. Strandberg, *Acta Crystallogr., Sect. B: Struct. Crystallogr. Cryst. Chem.*, 1979, **35**, 278–284, DOI: 10.1107/S0567740879003356.
- 171 B. Hedman, *Acta Chem. Scand.*, 1973, **27**, 3335–3364, DOI: 10.3891/acta.chem.scand.27-3335.
- 172 S. Himeno, M. Hashimoto and T. Ueda, *Inorg. Chim. Acta*, 1999, **284**, 237–245, DOI: 10.1016/S0020-1693(98)00294-1.
- 173 C. E. Easterly, D. M. Hercules and M. Houalla, *Appl. Spectrosc.*, 2001, **55**, 1671–1675, DOI: 10.1366/0003702011953991.
- 174 M. Hashimoto, I. Andersson and L. Pettersson, *Dalton Trans.*, 2007, 124–132, DOI: 10.1039/b613214f.
- 175 M. Hashimoto, I. Andersson and L. Pettersson, *Dalton Trans.*, 2009, 3321–3327, DOI: 10.1039/B818530A.



- 176 R. Strandberg, *Acta Chem. Scand., Ser. A*, 1974, **28**, 217–225, DOI: 10.3891/acta.chem.scand.28a-0217.
- 177 R. Strandberg, *Acta Chem. Scand., Ser. A*, 1975, **29A**, 350–358, DOI: 10.3891/acta.chem.scand.29a-0350.
- 178 J. A. Rob van Veen, O. Sudmeijer, C. A. Emeis and H. de Wit, *J. Chem. Soc., Dalton Trans.*, 1986, 1825–1831, DOI: 10.1039/dt9860001825.
- 179 L. A. Combs-Walker and C. L. Hill, *Inorg. Chem.*, 1991, **30**, 4016–4026, DOI: 10.1021/ic00021a010.
- 180 A. J. Gaunt, I. May, M. J. Sarsfield, D. Collison, M. Helliwell and I. S. Denniss, *Dalton Trans.*, 2003, 2767–2771, DOI: 10.1039/b301995k.
- 181 R. Strandberg, *Acta Chem. Scand., Ser. A*, 1975, **29**, 359–364, DOI: 10.3891/acta.chem.scand.29a-0359.
- 182 L. Pettersson, *Acta Chem. Scand., Ser. A*, 1975, **29**, 677–689, DOI: 10.3891/acta.chem.scand.29a-0677.
- 183 G. Johansson, L. Pettersson and N. Ingri, *Acta Chem. Scand., Ser. A*, 1978, **32**, 681–688, DOI: 10.3891/acta.chem.scand.32a-0681.
- 184 X. Zhang, X. Luo, Y. Duan, Y. Huang, N. Zhang, L. Zhao and J. Wu, *J. Mol. Struct.*, 2017, **1141**, 245–251, DOI: 10.1016/j.molstruc.2017.03.110.
- 185 B. Hedman, *Acta Crystallogr., Sect. B: Struct. Crystallogr. Cryst. Chem.*, 1980, **36**, 2241–2246, DOI: 10.1107/S0567740880008503.
- 186 B.-W. Cong, Z.-H. Su, Z.-F. Zhao, B.-Y. Yu, W.-Q. Zhao, L. Xia, X.-J. Ma and B.-B. Zhou, *CrystEngComm*, 2017, **19**, 2739–2749, DOI: 10.1039/c7ce00319f.
- 187 H. Ichida and Y. Sasaki, *Acta Crystallogr., Sect. C: Cryst. Struct. Commun.*, 1983, **39**, 529–533, DOI: 10.1107/S0108270183005363.
- 188 K. Murata and S. Ikeda, *Polyhedron*, 1987, **6**, 1681–1685, DOI: 10.1016/S0277-5387(00)80771-8.
- 189 D. Bajuk-Bogdanović, I. Holclajtner-Antunović, M. Todorović, U. B. Mioć and J. Zakrzewska, *J. Serb. Chem. Soc.*, 2008, **73**, 197–209, DOI: 10.2298/JSC0802197B.
- 190 G. B. McGarvey and J. B. Moffat, *J. Mol. Catal.*, 1991, **69**, 137–155, DOI: 10.1016/0304-5102(91)80110-O.
- 191 A. Jürgensen and J. B. Moffat, *Catal. Lett.*, 1995, **34**, 237–244, DOI: 10.1007/BF00808338.
- 192 D. Bajuk-Bogdanović, S. Uskoković-Marković, R. Hercigonja, A. Popa and I. Holclajtner-Antunović, *Spectrochim. Acta, Part A*, 2016, **153**, 152–159, DOI: 10.1016/j.saa.2015.08.029.
- 193 V. W. Truesdale and C. J. Smith, *Analyst*, 1975, **100**, 203–212, DOI: 10.1039/an9750000203.
- 194 V. W. Truesdale and C. J. Smith, *Analyst*, 1975, **100**, 797–805, DOI: 10.1039/AN9750000797.
- 195 M. Takahashi, Y. Abe and M. Tanaka, *Talanta*, 2015, **131**, 301–308, DOI: 10.1016/j.talanta.2014.07.079.
- 196 H. Katano, T. Osakai, S. Himeno and A. Saito, *Electrochim. Acta*, 1995, **40**, 2935–2942, DOI: 10.1016/0013-4686(95)00225-4.
- 197 F. Li and L. Xu, *Dalton Trans.*, 2011, **40**, 4024–4034, DOI: 10.1039/c0dt00691b.
- 198 R. Massart, R. Contant, J.-M. Fruchart, J.-P. Ciabrini and M. Fournier, *Inorg. Chem.*, 1977, **16**, 2916–2921, DOI: 10.1021/ic50177a049.
- 199 S. Dianat, S. Tangestaninejad, B. Yadollahi, A.-K. Bordbar, M. Moghadam, V. Mirkhani and I. Mohammadpoor-Baltork, *J. Mol. Liq.*, 2012, **174**, 76–79, DOI: 10.1016/j.molliq.2012.07.006.
- 200 N. Wang, F. Li, Y. Wang, L. Xu, K. Cui and B. Xu, *Eur. J. Inorg. Chem.*, 2013, 1699–1705, DOI: 10.1002/ejic.201201137.
- 201 A. Blazevic and A. Rompel, *Coord. Chem. Rev.*, 2016, **307**, 42–64, DOI: 10.1016/j.ccr.2015.07.001.
- 202 L. O. Öhman, *Inorg. Chem.*, 1989, **28**, 3629–3632, DOI: 10.1021/ic00318a004.
- 203 L. Yang, Z. Zhou, P.-T. Ma, X.-F. Zhang, J.-P. Wang and J.-Y. Niu, *J. Coord. Chem.*, 2013, **66**, 1058–1067, DOI: 10.1080/00958972.2013.775429.
- 204 A. Blazevic, E. Al-Sayed, A. Roller, G. Giester and A. Rompel, *Chem. – Eur. J.*, 2015, **21**, 4762–4771, DOI: 10.1002/chem.201405644.
- 205 R. Copping, L. Jonasson, A. J. Gaunt, D. Drennan, D. Collison, M. Helliwell, R. J. Pirttijarvi, C. J. Jones, A. Huguet, D. C. Apperley, N. Kaltsoyannis and I. May, *Inorg. Chem.*, 2008, **47**, 5787–5798, DOI: 10.1021/ic800101t.
- 206 S. Li, J. Zhao, P. Ma, J. Du, J. Niu and J. Wang, *Inorg. Chem.*, 2009, **48**, 9819–9830, DOI: 10.1021/ic901330y.
- 207 K. von Allmen, R. Moré, R. Müller, J. Soriano-López, A. Linden and G. R. Patzke, *ChemPlusChem*, 2015, **80**, 1389–1398, DOI: 10.1002/cplu.201500074.
- 208 D. Jabbour, B. Keita, I.-M. Mbomekalle, L. Nadjo and U. Kortz, *Eur. J. Inorg. Chem.*, 2004, 2036–2044, DOI: 10.1002/ejic.200300830.
- 209 K.-C. Kim and M. T. Pope, *J. Am. Chem. Soc.*, 1999, **121**, 8512–8517, DOI: 10.1021/ja9909125.
- 210 D. Lieb, A. Zahl, E. F. Wilson, C. Streb, L. C. Nye, K. Meyer and I. Ivanović-Burmazović, *Inorg. Chem.*, 2011, **50**, 9053–9058, DOI: 10.1021/ic201243n.
- 211 J. Liu, F. Ortéga, P. Sethuraman, D. E. Katsoulis, C. E. Costello and M. T. Pope, *J. Chem. Soc., Dalton Trans.*, 1992, 1901–1906, DOI: 10.1039/dt9920001901.
- 212 J. Moreau, O. Delpoux, E. Devers, M. Digne and S. Lorient, *J. Phys. Chem. A*, 2012, **116**, 263–270, DOI: 10.1021/jp208534p.
- 213 L. C. W. Baker and D. C. Glick, *Chem. Rev.*, 1998, **98**, 3–50, DOI: 10.1021/cr960392l.
- 214 J. J. Cruywagen and I. F. J. van der Merwe, *J. Chem. Soc., Dalton Trans.*, 1987, 1701–1705, DOI: 10.1039/DT9870001701.
- 215 G. M. Rozantsev, S. V. Radio and N. I. Gumerova, *Pol. J. Chem.*, 2008, **82**, 2067–2080.
- 216 G. M. Rozantsev and O. I. Sazonova, *Russ. J. Coord. Chem.*, 2005, **31**, 552–558, DOI: 10.1007/s11173-005-0135-x.
- 217 J. J. Hastings and O. W. Howarth, *J. Chem. Soc., Dalton Trans.*, 1992, 209–215, DOI: 10.1039/DT9920000209.
- 218 R. I. Maksimovskaya and K. G. Burtseva, *Polyhedron*, 1985, **4**, 1559–1562, DOI: 10.1016/S0277-5387(00)87227-7.
- 219 D. K. Walanda, R. C. Burns, G. A. Lawrance and E. I. von Nagy-Felsobuki, *J. Cluster Sci.*, 2000, **11**, 5–28, DOI: 10.1023/A:1009098627616.
- 220 T. Lehmann and J. Fuchs, *Z. Naturforsch., B: J. Chem. Sci.*, 1988, **43**, 89–93, DOI: 10.1515/znB-1988-0116.



- 221 X.-F. Wang, J. Cao, K.-L. Huang, Y.-Q. Xu, Y.-N. Chi and C.-W. Hu, *Eur. J. Inorg. Chem.*, 2013, 1788–1792, DOI: 10.1002/ejic.201200810.
- 222 I. Lindqvist, *Acta Crystallogr.*, 1952, **5**, 667–670, DOI: 10.1107/S0365110X52001817.
- 223 A. L. Nolan, E. N. Wilkes, T. W. Hambley, C. C. Allen, R. C. Burns and G. A. Lawrance, *Aust. J. Chem.*, 1999, **52**, 955–964, DOI: 10.1071/CH99070.
- 224 M. Asami, H. Ichida and Y. Sasaki, *Acta Crystallogr., Sect. C: Cryst. Struct. Commun.*, 1984, **40**, 35–37, DOI: 10.1107/S0108270184002924.
- 225 J. Fuchs, H. Hartl and W. Schiller, *Angew. Chem., Int. Ed. Engl.*, 1973, **12**, 420, DOI: 10.1002/anie.197304201.
- 226 J.-P. Launay, M. Boyer and F. Chauveau, *J. Inorg. Nucl. Chem.*, 1976, **38**, 243–247, DOI: 10.1016/0022-1902(76)80402-2.
- 227 J. Fuchs and E.-P. Flindt, *Z. Naturforsch., B: Chem. Sci.*, 1979, **34**, 412–422, DOI: 10.1515/znB-1979-0311.
- 228 C. R. Sprangers, J. K. Marmon and D. C. Duncan, *Inorg. Chem.*, 2006, **45**, 9628–9630, DOI: 10.1021/ic061370c.
- 229 S. Himeno, M. Yoshihara and M. Maekawa, *Inorg. Chim. Acta*, 2000, **298**, 165–171, DOI: 10.1016/S0020-1693(99)00436-3.
- 230 S. Himeno, K. Nakajima and K. Eda, *Polyhedron*, 2010, **29**, 2595–2599, DOI: 10.1016/j.poly.2010.06.002.
- 231 H. Hartl, R. Palm and J. Fuchs, *Angew. Chem., Int. Ed. Engl.*, 1993, **32**, 1492–1494, DOI: 10.1002/anie.199314921.
- 232 J. Fuchs, R. Palm and H. Hartl, *Angew. Chem., Int. Ed. Engl.*, 1996, **35**, 2651–2653, DOI: 10.1002/anie.199626511.
- 233 H. N. Miras, J. Yan, D.-L. Long and L. Cronin, *Angew. Chem., Int. Ed.*, 2008, **47**, 8420–8423, DOI: 10.1002/anie.200802109.
- 234 A. V. Anyushin, A. I. Smolentsev, D. A. Mainichev, C. Vicent, A. L. Gushchin, M. N. Sokolov and V. P. Fedin, *Chem. Commun.*, 2014, **50**, 9083–9085, DOI: 10.1039/C4CC02456G.
- 235 C. M. Tourné and G. F. Tourné, *J. Chem. Soc., Dalton Trans.*, 1988, 2411–2420, DOI: 10.1039/dt9880002411.
- 236 M. T. Ma, T. Waters, K. Beyer, R. Palamarczuk, P. J. S. Richardt, R. A. J. O'Hair and A. G. Wedd, *Inorg. Chem.*, 2009, **48**, 598–606, DOI: 10.1021/ic8016326.
- 237 D. L. Kepert and J. H. Kyle, *J. Chem. Soc., Dalton Trans.*, 1978, 137–141, DOI: 10.1039/dt9780000137.
- 238 B. J. Smith and V. A. Patrick, *Aust. J. Chem.*, 2002, **55**, 281–286, DOI: 10.1071/ch01092.
- 239 B. J. Smith and V. A. Patrick, *Aust. J. Chem.*, 2003, **56**, 283–286, DOI: 10.1071/ch01112.
- 240 A. Tézé, J. Canny, L. Gurban, R. Thouvenot and G. Hervé, *Inorg. Chem.*, 1996, **35**, 1001–1005, DOI: 10.1021/ic951004l.
- 241 B. S. Bassil and U. Kortz, *Dalton Trans.*, 2011, **40**, 9649–9661, DOI: 10.1039/c1dt10845j.
- 242 Q. Jia, J. Cao, Y. Duan and C. Hu, *Dalton Trans.*, 2015, **44**, 553–559, DOI: 10.1039/c4dt02723j.
- 243 J. M. Cameron, L. Vilà-Nadal, R. S. Winter, F. Iijima, C. J. Murillo, A. Rodríguez-Fortea, H. Oshio, J. M. Poblet and L. Cronin, *J. Am. Chem. Soc.*, 2016, **138**, 8765–8773, DOI: 10.1021/jacs.6b02245.
- 244 I. A. Weinstock, J. J. Cowan, E. M. G. Barbuzzi, H. Zeng and C. L. Hill, *J. Am. Chem. Soc.*, 1999, **121**, 4608–4617, DOI: 10.1021/ja982908j.
- 245 J. Breibeck, N. I. Gumerova, B. B. Boesen, M. Galanski and A. Rompel, *Sci. Rep.*, 2019, **9**, 5183, DOI: 10.1038/s41598-019-41261-7.
- 246 J. Breibeck, A. Bijelic and A. Rompel, *Chem. Commun.*, 2019, **55**, 11519–11522, DOI: 10.1039/c9cc05818d.
- 247 M. Pretzler, A. Bijelic and A. Rompel, *Sci. Rep.*, 2017, **7**, 1810, DOI: 10.1038/s41598-017-01813-1.
- 248 M. Pretzler and A. Rompel, *Inorg. Chim. Acta*, 2018, **481**, 25–31, DOI: 10.1016/j.ica.2017.04.041.
- 249 A. Solé-Daura, J. M. Poblet and J. J. Carbó, *Chem. – Eur. J.*, 2020, **26**, 5799–5809, DOI: 10.1002/chem.201905533.
- 250 L. S. Van Rompuy and T. N. Parac-Vogt, *Curr. Opin. Biotechnol.*, 2019, **58**, 92–99, DOI: 10.1016/j.copbio.2018.11.013.
- 251 T. K. N. Luong, P. Shestakova, T. T. Mihaylov, G. Absillis, K. Pierloot and T. N. Parac-Vogt, *Chem. – Eur. J.*, 2015, **21**, 4428–4439, DOI: 10.1002/chem.201405810.
- 252 D. Sloboda-Rozner, P. L. Alsters and R. Neumann, *J. Am. Chem. Soc.*, 2003, **125**, 5280–5281, DOI: 10.1021/ja0344821.
- 253 D. Sloboda-Rozner, P. Witte, P. L. Alsters and R. Neumann, *Adv. Synth. Catal.*, 2004, **346**, 339–345, DOI: 10.1002/adsc.200303156.
- 254 I. Y. Skobelev, V. Yu. Evtushok, O. A. Kholdeeva, N. V. Maksimchuk, R. I. Maksimovskaya, J. M. Ricart, J. M. Poblet and J. J. Carbó, *ACS Catal.*, 2017, **7**, 8514–8523, DOI: 10.1021/acscatal.7b02694.
- 255 Y. V. Geletii, C. Besson, Y. Hou, Q. Yin, D. G. Musaev, D. Quiñonero, R. Cao, K. I. Hardcastle, A. Proust, P. Kögerler and C. L. Hill, *J. Am. Chem. Soc.*, 2009, **131**, 17360–17370, DOI: 10.1021/ja907277b.
- 256 L. E. Briand, G. T. Baronetti and H. J. Thomas, *Appl. Catal., A*, 2003, **256**, 37–50, DOI: 10.1016/S0926-860X(03)00387-9.
- 257 R. Contant and A. Tézé, *Inorg. Chem.*, 1985, **24**, 4610–4614, DOI: 10.1021/ic00220a036.
- 258 R. Contant and R. Thouvenot, *Inorg. Chim. Acta*, 1993, **212**, 41–50, DOI: 10.1016/S0020-1693(00)92306-5.
- 259 C. Pimpão, I. V. da Silva, A. F. Mósca, J. O. Pinho, M. M. Gaspar, N. I. Gumerova, A. Rompel, M. Aureliano and G. Soveral, *Int. J. Mol. Sci.*, 2020, **21**, 2467, DOI: 10.3390/ijms21072467.
- 260 S. G. Mauracher, C. Molitor, R. Al-Oweini, U. Kortz and A. Rompel, *Acta Crystallogr., Sect. F: Struct. Biol. Commun.*, 2014, **70**, 263–266, DOI: 10.1107/S2053230X14000582.
- 261 S. G. Mauracher, C. Molitor, R. Al-Oweini, U. Kortz and A. Rompel, *Acta Crystallogr., Sect. D: Biol. Crystallogr.*, 2014, **70**, 2301–2315, DOI: 10.1107/S1399004714013777.
- 262 C. Molitor, A. Bijelic and A. Rompel, *Chem. Commun.*, 2016, **52**, 12286–12289, DOI: 10.1039/C6CC07004C.
- 263 A. Bijelic, C. Molitor, S. G. Mauracher, R. Al-Oweini, U. Kortz and A. Rompel, *ChemBioChem*, 2015, **16**, 233–241, DOI: 10.1002/cbic.201402597.
- 264 C. Molitor, S. G. Mauracher and A. Rompel, *Proc. Natl. Acad. Sci. U. S. A.*, 2016, **113**, E1806–E1815, DOI: 10.1073/pnas.1523575113.
- 265 N. I. Gumerova, T. Caldera Fraile, A. Roller, G. Giester, M. Pascual-Borràs, A. C. Ohlin and A. Rompel, *Inorg. Chem.*, 2019, **58**, 106–113, DOI: 10.1021/acs.inorgchem.8b01740.

