

Review

Polyoxidovanadates' interactions with proteins: An overview

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ABSTRACT

Polyoxidovanadates (POVs, previously named polyoxovanadates) are a subgroup of polyoxidometalates (POMs, previously named polyoxometalates) with interesting pharmacological actions that have been tested as potential antidiabetic, antibacterial, antiprotozoal, antiviral, and anticancer drugs. They contain mainly vanadium and are able to interact with proteins, affecting various biological processes. The most studied POV is the isopolyoxidovanadate decavanadate (V_{10}), which interacts with proteins and/or enzymes such as tyrosine protein phosphatases, P-type ATPases, RNA triphosphatases, myosin and actin. However, in many POVs–protein systems, the binding sites and/or the residues involved in the interaction are not identified. In the present review, the interactions of POVs, as well as linear trivanadate (V_3), both linear and cyclic tetraavanadate (V_4) and two proposed heptavanadate (V_7 ; which are better described by V_{10} molecules), with proteins are described through X-ray crystallographic studies. Interactions with POVs through theoretical and spectroscopic studies of proteins related to muscle contraction, serum, oxidative stress, and diabetes were also discussed. In sum, herein, we describe POVs' interactions with various proteins including acid phosphatase A, receptor tyrosine kinase, ectonucleoside triphosphate diphosphohydrolase (NTPDases), transient receptor potential cation channel (TRPM4), phosphoglucosyltransferases, P-type ATPases, myosin, actin, transferrin, albumin, and glucosidases, among others. The putative POVs' effects on proteins are impacted by the POV' stability and speciation. The modes of POVs' interactions include H-bond, electrostatic, H-bond + electrostatic, van der Waals, and covalent binding. The spectroscopic, X-ray and computational results, the sites and modes of binding are described in detail.

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Abbreviations

^{51}V -NMR	vanadium-51 NMR	PDB	protein data bank
ABC	ATPases, ATP binding-cassette ATPases	PG	phosphoglycerate
AcPA	acid Phosphatase A	POM	polyoxidometalate
ADP	adenosine diphosphate	POMo	polyoxidomolybdate
AMP	adenosine monophosphate	PONb	polyoxidoniobate
ATP	adenosine triphosphate	POT	polyoxidotungstate
BPG	bisphosphoglycerate	POV	polyoxido vanadate
CAT	catalase	ROS	reactive oxygen species
Ca^{2+} -ATPase	adenosine triphosphatase, calcium dependent	S_1	myosin subfragment 1
CAN	Ca^{2+} activated, non-selective channel	SOD	superoxide dismutase
Cdk	cyclin-dependent kinases	SR	sarcoplasmic reticulum
DMAPH	4-dimethylaminopyridinium	TEW	tellurium-centered Anderson polyoxidotungstate [$\text{TeW}_6\text{O}_{24}$] $^{6-}$
DMEM	Dulbecco's modified eagle medium	TRP	transient receptor potential channel
dPGM	dependent phosphoglycerate mutase	TRPM4	transient receptor potential cation channel
F-actin	filamentous polymerized actin	UDP	uridine diphosphate
G-actin	monomeric actin	UTP	uridine triphosphate
GSH	reduced glutathione	V_1	monomeric vanadate, simplest oxido vanadate
HSA	human serum albumin	V_2	dimeric vanadate, divanadate
HTf	human serum transferrin	V_3	trimeric vanadate, trivanadate
IC_{50}	half maximal inhibitory concentration	V_4	tetrameric vanadate, tetravanadate
MD	molecular docking	V_7	heptameric vanadate, heptavanadate
MHRs	N-terminal TRPM homology regions	V_{10}	decameric vanadate, decavanadate
Metf	metformin	VDAC	voltage-dependent anion-selective channel
NRF2	nuclear factor erythroid 2-related factor 2	vdW	van der Waals
NTP	nucleoside triphosphate		
NTPDase	ecto-nucleotidetriphosphate diphosphohydrolase		

1. Introduction

In the last 30 years, the number of studies of polyoxidometalates (POMs, previously named polyoxometalates) and polyoxido vanadates (POVs, previously named polyoxovanadates) associated with enzymatic inhibition [1–5] and diseases, including insulin enhancement agents [6–9] for diabetes mellitus, and inhibitors of the aggregation of amyloid β -peptides associated with Alzheimer's disease have clearly increased [10,11]. The growing interest in POMs and POVs is extending their applications to diverse areas of basic and applied sciences [12–18] with several studies about various aspects and applications of POVs published covering chemical engineering [19] catalysis [20], environmental chemistry [21], material science [18], biochemistry, biology, pharmacology, and medicine [5,13,14,22–24].

POVs' structures have widespread sizes and shapes and may include other hetero-ions such as P^{V} and As^{V} (Fig. 1). The addenda metal oxidoions $\text{M}=\text{O}$ generally contain $\text{M} = \text{W}^{\text{VI}}$, Mo^{VI} , and/or V^{V} and other transition metal ions such as Co^{II} , and/or Mn^{II} . The structures are described in excellent reviews on chemistry of these POV compounds [12,23,25,26].

In the present review, we describe recent studies on the interaction of polyoxido vanadates with proteins, depending on the diversity and nature of the particular POV, and their potential implications in biological processes as well as in biomedical

applications. In order to understand such an interaction, information on the stability and speciation under physiological conditions are necessary. In some studies, the protein interactions with other oxido vanadates, such as monomeric (V_1 , Fig. 1A), dimeric (V_2 , Fig. 1B), trimeric (V_3) and tetrameric (V_4 , Fig. 1C, D) vanadates, as well as other POVs were investigated [27]. The effects of vanadium coordination complexes are not the focus of this review, and the reader is referred to other sources for such information [28–32].

2. POVs' stability and speciation

Considering the structural complexity of POVs, their stabilities are variable and it is important to determine what species exist in solution together with the proteins. Hence, it is desired that the stability of the species is investigated under the assay conditions, as recently reviewed [5,33,34].

2.1. Stability of POVs under physiological conditions

Decavanadate, $[\text{V}_{10}\text{O}_{28}]^{6-}$ (V_{10} , Fig. 1H), is an isopolyoxido vanadate that has been described in detail and serves to showcase the relevance of POVs' speciation and stability to deduce its biological effects [34,35]. Decavanadate has an oblong, compact structure with dimensions of $8.3 \times 7.7 \times 5.4 \text{ \AA}$ [36–38]. By ^{51}V NMR

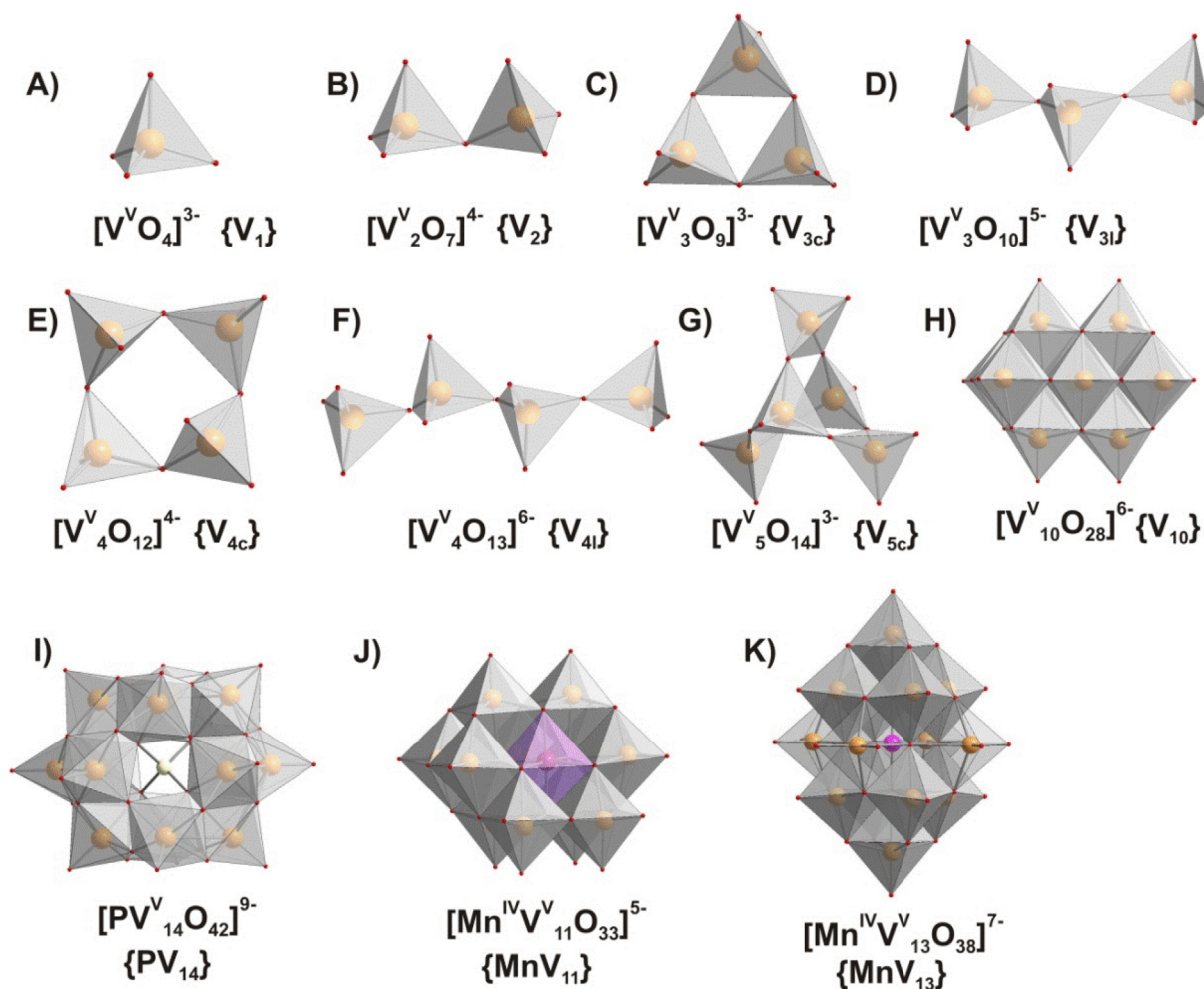


Fig. 1. Oxidovanadates and polyoxidovanadate structures with the nuclearity up to 14 addenda atoms and both tetrahedral and octahedral coordination of V. Tri- and tetra- vanadates are presented in two forms – cyclic ($\{V_{3c}\}$, C and $\{V_{4c}\}$, E) and linear ($\{V_{3l}\}$, D and $\{V_{4l}\}$, F). The linear forms of $\{V_{3l}\}$ and $\{V_{4l}\}$ have never been obtained in solid state. In the $\{MnV_{13}\}$ structure, K, four equatorial V atoms have 75 % occupancy [27] and so in total correspond to three V ions with full occupancy in the sum formula. Color code: $\{VO_x\}$, grey; O, red; P, yellow; Mn, pink. The subscript “c” stands for “cyclic”, “l” for “linear”. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

spectroscopy, three different vanadium ion environments can be distinguished in the V_{10} structure. The yellow or bright orange color of V_{10} solutions can be detected at 360 and 400 nm by UV/Vis spectroscopy [34,39] and the V_{10} stability can be followed by this technique, even for μM concentrations [39,40]. Once formed, V_{10} was shown to be persisting in solution even in the neutral and basic pH range where V_{10} is expected to be no longer thermodynamically stable [34,35]; kinetic reasons can explain this behavior. At acidic pH values, from pH 2 to 6, it is the most stable oxidovanadate. The speciation profile is shown in Fig. 2, illustrating that V_{10} has several protonation states with all the pK_a values falling in the range from pH 2 to 6.

The acidic orange decavanadate solutions can be hydrolyzed by either heating or boiling the neutral or alkaline solutions into the colorless monomeric and oligomeric vanadates [34,41,42]. ^{51}V NMR spectroscopy can be used to monitor the speciation of V_{10} in enzyme assays or in growth media and experiments can be designed to evaluate specific interactions with proteins [5,34,39,40,43,44]. For V_{10} , as well as for other POVs, particularly those that are labile such as V_2 and V_4 , respectively, it is possible by combining kinetic with spectroscopic studies to analyze their stability, recognize their specific protein interactions, and determine the specific POV species' contribution to the observed biolog-

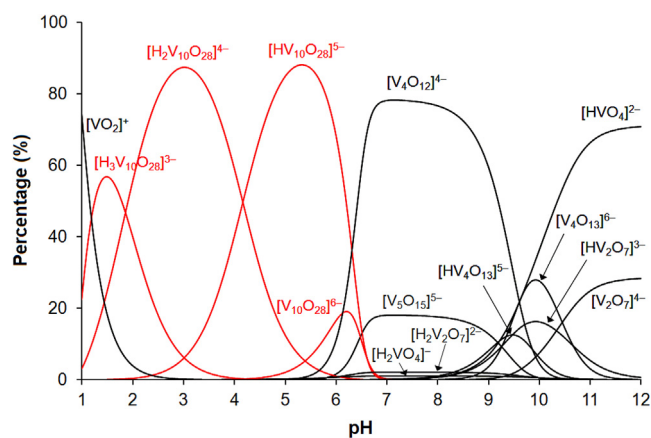


Fig. 2. Aqueous solution species distribution of V(V) as a function of pH at 0.200 M. The stability constants of the inorganic vanadates were taken from [41]. Decavanadate (V_{10}) species are shown in red. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

ical effects [34,35,45–48]. In contrast to the labile oxidovanadates, the half-life of decavanadate is much greater and in serum is 15 h

at ambient temperature [39], whereas in buffered media at 25 °C and 37 °C, the half-life decreases to 12 h and 3 h, respectively [39,49]. When V₁₀ was applied to media with growth additives for mycobacteria, the half-life reduces even further to a few hours [50]. Metal-based compounds are subject to speciation chemistry [51], which depend on buffers itself and all the other components in the solution, emphasizing the need to confirm the stability of V₁₀ under the assay conditions [34,35,39,49,52,53].

The kinetics of the decomposition of sodium and metforminium decavanadate were determined in Dulbecco's modified Eagle's medium (DMEM, CaCl₂, 0.2 g/L; KCl, 0.4 g/L; NaCl, 6.4 g/L; Na₂HPO₄, 0.109 g/L; Na₂CO₃, 3.7 g/L; glucose, 1 g/L, and 20 proteino-genic amino acids) at pH 7.4 and 25 °C [54]. In these systems, the results showed that the half-life time of [H₂VO₄]⁻ (V₁, Fig. 1A) and other smaller oxidovanadates species such as [H₂V₂O₇]²⁻ (V₂, Fig. 1B), [V₄O₁₂]⁴⁺ (V₄, Fig. 1E) and [V₅O₁₅]⁵⁻ (V₅) were 9 and 11 h for sodium and metforminium salts, consistent with earlier reports in other growth media and differences in lifetime based on counterion, temperature and, most importantly, pH [55–57].

2.2. Do proteins affect POVs' stability and/or speciation?

Although speciation studies consider all the contributions of components in solution, the effects of proteins are usually neglected [39,53]. The presence of macromolecules can significantly affect, and in some cases increase, the stability of V₁₀. In 2006, it was described that, at pH 7.4, the decavanadate's half-life increases 5.5-fold in the presence of the monomeric form of actin (G-actin) [39], while the addition of *Mycobacterium smegmatis* or *Mycobacterium tuberculosis* cells to a V₁₀ solution between pH 5.8 and 6.8 causes immediate decomposition of the decavanadate [50]. Changes in its hydrolytic stability caused by the addition of G-actin is in contrast to the stability of POVs in pure water and illustrate one more time the potential change in the POV's stability through addition of biomolecules [39,50]. Similarly, studies with V₁₀ as a phosphatase inhibitor have shown that the enzymes may facilitate hydrolysis [58]. V₁₀ is stable for weeks in pure aqueous at pH 4.0 and no change in the hydrolysis rate is observed [59].

3. POVs' interactions with proteins

In the present review we focus on POVs' interactions with proteins, which complements previous reviews on the binding of general vanadium compounds [60,61], polyoxidomolybdates- or polyoxidotungstate-protein binding [1,3,15] and updates previous reviews on V₁₀ and other POVs [5,34]. The interactions of POV (summarized in Tables 1 and 2, starting from 1989 for Table 1) with protein can be grouped into:

i) *Coordinative or covalent* interaction of an amino acid side-chain to V(V). This unusual binding type is represented by binding of β-alanine to {V₆O₂₄} [62]. The possibility that this type of bonding does exist is supported by the observation that the presence of V₁₀ caused oxidation of Cys374 in F-actin and one of the protein core cysteine residues in G-actin, presumably Cys272 [63,64] (see below in section 3.2). Oxidation of the protein supports a putative POV bond to the protein or intermolecular electron transfer.

ii) *Non-covalent* binding, more common than the *covalent* one, when the interaction occurs through secondary interactions, namely van der Waals (vdW) contacts and hydrogen bonds (H-bonds). This binding mode depends on several factors, such as the electric charge that influences the electrostatic interactions, the shape and volume size that determine the host-guest complementary, and the formation of H-bonds with the terminal oxygens [65,66]. Other variables are pH and ionic strength. The H-bonds are often directed or mediated by H₂O molecules [67]. The majority of

Table 1
POVs interactions with proteins or inhibition of enzymes, *in vitro* studies.

Protein/Enzyme/Effects	POVs (pH of stock solution)	Year	Ref.
Acid phosphatase (human prostate)	V ₁ and V ₂	1989	[48]
6-phosphogluconate dehydrogenase	V ₄	1990	[45]
Glucose-6-phosphate dehydrogenase (<i>Leuconostoc mesenteroides</i>)	V ₂ and V ₄	1990	[89]
Glycerol-3-phosphate dehydrogenase	V ₄ (and possible weak interaction by V ₂)	1991	[97]
Superoxide dismutase	V ₄	1991	[90]
Purple acid phosphatase (porcine uterine fluid)	V ₁ and oxidovanadium(IV) cation VO ₂ ⁺	1992	[98]
Fructose 1,6 bisphosphate aldolase (muscle)	V ₂ and V ₄	1992	[46]
cAMP-dependent protein kinase	V ₁₀ (7.1)	1997	[91]
DNA-binding protein	V ₁ (pH 11) and V ₁₀ (pH 7.4)	2002	[92]
Methaemoglobin reductase inhibition	V ₁ (pH 6.7) and V ₁₀ (pH 4)	2003	[55]
Myosin/actomyosin ATPase inhibition	V ₁₀ (pH 4)	2004	[49]
Muscle contraction regulation	V ₁₀ (pH 4)	2004	[99]
ATP sensitive cation channels	V ₁₀ (pH 2)	2004	[100]
TRPM4 cation channels	V ₁₀ (pH 2)	2004	[101]
G-actin polymerization inhibition	V ₁₀ (pH 4)	2006	[39]
RNA triphosphatase	V ₁ (ortho- and meta; pH not specified) and V ₁₀ (pH not specified)	2006	[102]
P2X receptor antagonist	V ₁₀ (pH 2)	2006	[103]
Back-door binding to myosin	V ₁₀ (pH 4)	2007	[104]
Porin (VDAC) modulator	V ₁₀ (pH not specified)	2007	[105]
Mitochondrial membrane depolarization; changes in the redox steady-state of cytochrome <i>b</i> (complex III) Inhibition of mitochondrial oxygen consumption	V ₁₀ (pH 4)	2007	[106]
G-Actin oxidation and oxidovanadium(IV) formation	V ₁₀ (pH 4)	2009	[63]
Na ⁺ /K ⁺ -ATPase inhibition	V ₁₀ (pH not specified)	2009	[107]
Ca ²⁺ -ATPase	V ₁₀ (pH 4)	2012	[56]
Type 1 Fc epsilon receptor	V ₁₀ (pH 7.4)	2013	[108]
ATP prevents V ₁₀ G-actin reduction	V ₁₀ (pH 4)	2017	[109]
<i>ex-vivo</i> Na ⁺ /K ⁺ -ATPase inhibition coupled with chloride secretion	PV ₁₄ (Fig. 1I)	2019	[57]
SR Ca ²⁺ -ATPase	MnV ₁₁ (Fig. 1J) MnV ₁₃ (Fig. 1K)	2019	[110]
Thaumatococcus, lysozyme, albumin, transferrin	V ₁₀ , V ₁₀ Cu (decavanadate coordinated to Cu ²⁺), V ₁₀ Co (decavanadate coordinated to Co ²⁺)	2019	[43]
V ₁₀ /Nb ₁₀ G-actin binding sites	V ₁₀ (pH 4)	2021	[64]

the possible binding modes of V₁₀ in proteins as well as a summary of the non-covalent interacting amino acids are given in Fig. 3.

Distinguishing *coordinative* from *non-covalent* bond and determining their strength is not a trivial task and computational methods are often necessary [68,69]. On the framework of *non-covalent* binding, several docking software programs offer the opportunity to examine metal-protein interactions [70–73]. In most of the cases, reproducing experimental results requires adjusting the weights of intramolecular and intermolecular H-bonds and vdW

Table 2
Protein complexes with V₁₀ and smaller oxidovanadates investigated using X-ray crystallography or cryogenic electron microscopy.

Protein	Source	PDB code	V-cluster	V-precursor added	pH of media	Image/ method	Ref.
Acid Phosphatase A (AcPA)	<i>Francisella tularensis</i>	2D1G	Binds V ₁₀ on surface (Fig. 1H) Binds V ₁ in active site	Na ₃ VO ₄	6.0	Fig. 6 / X-ray	[114]
Ecto-nucleoside triphosphate diphosphohydrolase (NTPDase)	<i>Rattus norvegicus</i>	3ZX2	V ₁₀ (Fig. 1H)	Na ₃ VO ₄	4.0	Fig. 7A / X-ray	[119]
NTPDase	<i>Legionella pneumophila</i>	4BRH	V ₁₀ (Fig. 1H)	Na ₃ VO ₄	5.0	Fig. 7B / X-ray	[120]
Tyrosine kinase	<i>Homo sapiens</i> , <i>Rattus norvegicus</i>	3GQI	V ₁₀ (Fig. 1H)	Na ₃ VO ₄	8.0	Fig. 8 / X-ray	[121]
Human transient receptor potential cation channel (TRPM4)	<i>Homo sapiens</i>	5WP6	V ₁₀ (Fig. 1H)	Na ₃ VO ₄	2.0	Fig. 9 / cryo-EM	[126]
Human cell cycle protein ckshs1	<i>Homo sapiens</i>	1DKT	V ₇ in reported model; but the actual cluster binding is V ₁₀ (Fig. 1H)	NaVO ₃	7.4	Fig. 10 / X-ray	[113]
Co-factor dependent phosphoglycerate mutase (dPGM)	<i>E. coli</i>	1E59	Linear V ₄ (Fig. 1F)	NaVO ₃	8.0	Fig. 11 / X-ray	[129]
C3 exoenzyme	<i>Clostridium botulinum</i>	1UZI	Cyclic V ₄ (Fig. 1E)	Na ₃ VO ₄	5.5	Fig. 12A / X-ray	[131]
BtuCD protein	<i>E. coli</i>	1L7V	Cyclic V ₄ (Fig. 1E)	Na ₃ VO ₄	8.0	Fig. 12B / X-ray	[132]
Phosphatase PhoE	<i>Geobacillus stearothermophilus</i>	1H2F	Linear V ₃ (Fig. 1D)	NH ₄ VO ₃	4.5 and 5.0	Fig. 13 / X-ray	[133]
Uridine phosphorylase	<i>E. coli</i>	1RXS	Unlikely isomer V ₇ , but information not sufficient to determine the cluster	Na ₃ VO ₄	7.5	No Figure shown / X-ray	[117]
Protein-tyrosine phosphatase YopH	<i>Yersinia</i>	3F9B	V ₂ (Fig. 1B)	Na ₃ VO ₄	7.5	No Figure shown / X-ray	[134]

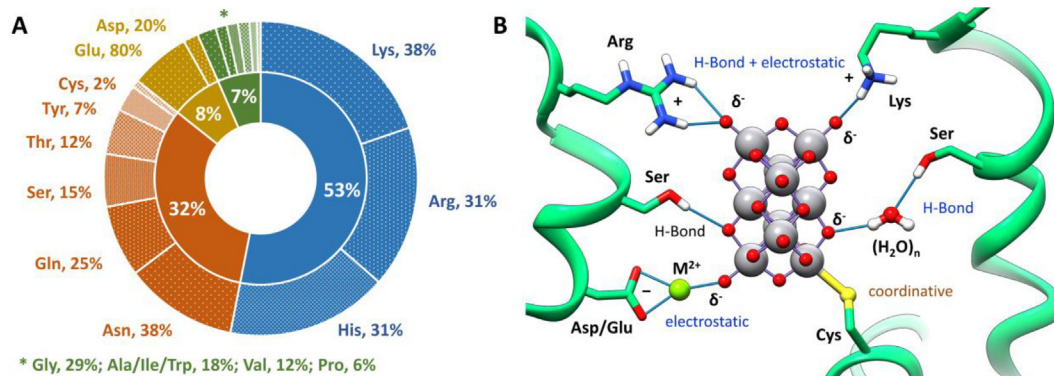


Fig. 3. A) Percentage of amino acids interacting with V₁₀ (blue basic, orange polar uncharged, yellow acidic, green unipolar) and B) possible modes of POVs' interaction with proteins. Data taken from ref. [15]. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

contacts [74]. The *coordinative* binding can be described with pure electrostatic functions [70–72] or through *covalent docking* approaches nowadays implemented in packages such as GOLD [75] or Autodock [71] and source code modifications such as CovalentDock [76] or Docktite [77]. In these cases, the metal–protein bond needs to be defined *a priori* and restrained during docking. A recently updated version of the *GoldScore* scoring function of the GOLD program was successfully validated for a large series of metal complexes demonstrating its capability to predict coordination bonds without any geometrical constraint or energy restraint [78,79].

A report in 1973 introduced V₁₀ as a microMolar inhibitor of rabbit muscle adenylate kinase [80]. These studies were followed by several papers discussing a number of other glycolytic enzymes including fructohexokinase, and phosphofructokinase, among others, all inhibited by V₁₀ [5,81–83]. These inhibition studies were

complemented early on through ⁵¹V NMR studies by Csermely *et al.* in 1985 [84], reporting that the addition of Ca²⁺-ATPase from the sarcoplasmic reticulum to a solution of both labile oxidovanadates and V₁₀ would result in ⁵¹V NMR spectra in which V₁₀ signals selectively disappeared. These results are consistent with the binding of Ca²⁺-ATPase to the V₁₀ over the simple mono-oxidovanadates (Fig. 1) [85]. The effects of V₁₀ in the structure and function of the Ca²⁺-ATPase were further explored [85,86]. Thus, it was found that only V₁₀ species, and not the V₁ species, were able to inhibit calcium accumulation coupled with ATP hydrolysis [85], as well as proton ejection [86] by the sarcoplasmic reticulum Ca²⁺-ATPase, clearly affecting energy transduction processes [85–87].

Analysis of the speciation in aqueous solution accompanied the enzyme studies [34,35,46,83–91], and in the case of 6-phosphogluconate dehydrogenase all the observed inhibition could

be attributed to the V_{4c} (Fig. 1E) species [45]. Subsequent kinetic studies confirmed inhibition with corresponding tetranuclear molybdate and ruled out inhibition with the pentanuclear molybdate [88]. Several additional studies with glycolytic enzymes including *Leuconostoc mesenteroides* glucose-6-phosphate dehydrogenase, glycerol-3-phosphate dehydrogenase, and muscle fructose 1,6-bisphosphate aldolase illustrated that V_{4c} was an inhibitor, although partial inhibition may have been contributed by a weaker inhibiting V_2 . The enzyme source can affect the selectivity as illustrated in the case of the *Leishmania* acid phosphatase where both V_{10} and simple monomeric vanadate V_1 were considered to be the inhibitors [2,58], whereas for the human prostatic phosphatase both V_1 and V_2 were inhibitors. These two reports are contrary to most other phosphatases that are inhibited potently by V_1 [93–96]. Besides the ones described above, several

studies of the interaction/effects of POVs with proteins/enzymes have been reported since 1989, using kinetic and spectroscopic methods and are summarized in Table 1. Some of the milestones for POVs' interactions with proteins and enzymes are highlighted in a chronological order in Fig. 4.

3.1. X-ray crystallographic studies of POVs with proteins

3.1.1. Overview of V-protein and POV-protein complexes

A search of the Protein Data Bank (PDB) [111,112] on the structures determined by X-ray crystallography using the terms «vanadium», «vanadate», «vanadyl» and «proteins» gives over 150 structures as shown in Fig. 5. This is about 40 structures more than reported in 2015 when an analysis of V-phosphatases complexes was reported [93–96] and a testimony to the increased interest

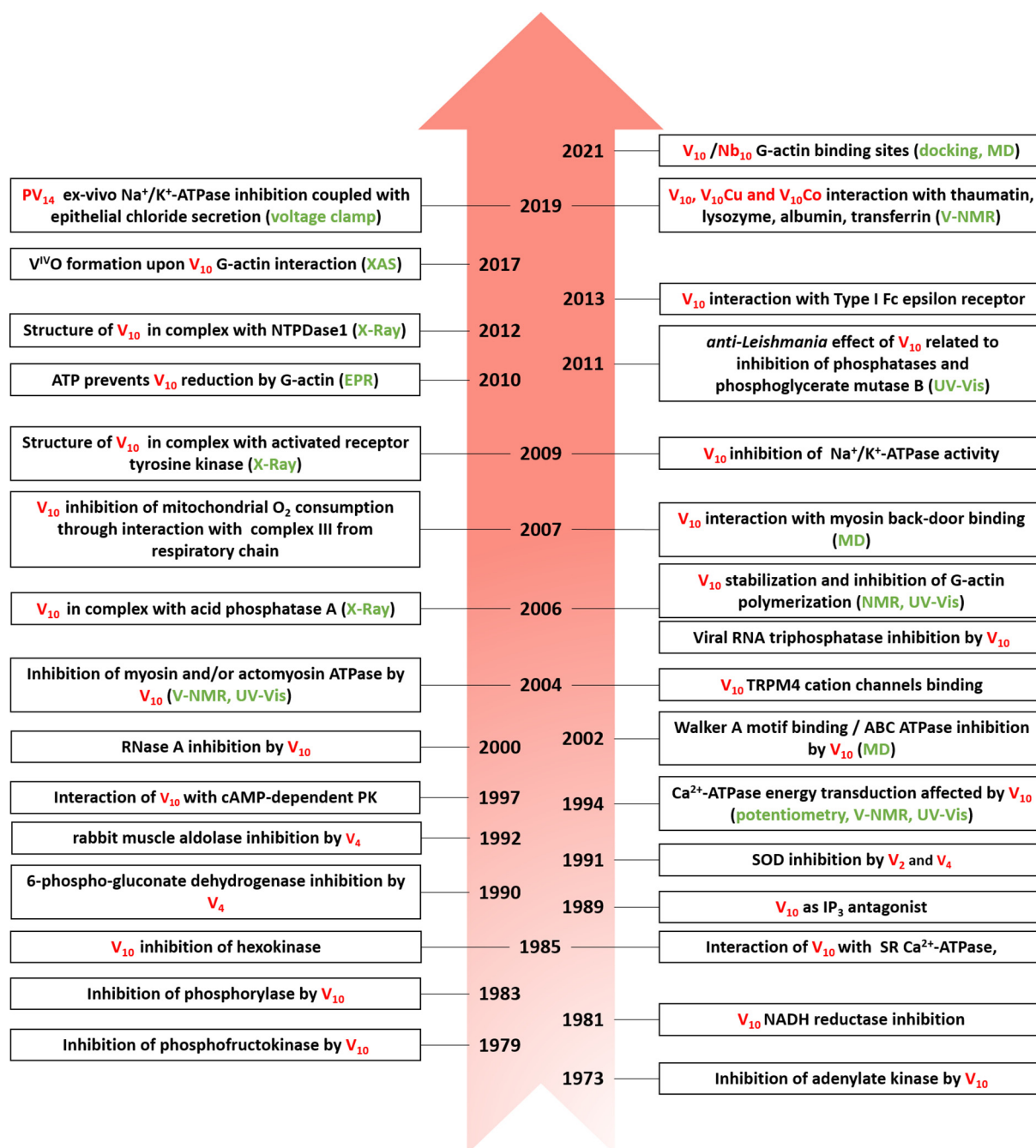


Fig. 4. Timeline for studies of POVs interactions with proteins.

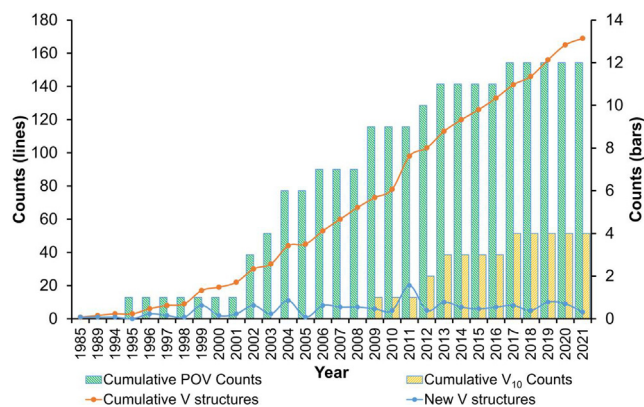


Fig. 5. A plot of POV-proteins complexes reported in the PDB database as a function of time. The right Y-axis relates to the cumulative counts of V₁₀, the left Y-axis describes counts of POVs, cumulative V-structures and new V structures per year.

of V-protein interaction. However, only four structures contain V₁₀, with a fifth structure when cryogenic electron microscopy (cryo-EM) as a technique is used (lined bars, Fig. 5) and a sixth structure where the electron density of the V₁₀ is not complete. These six structures include an acid phosphatase, a tyrosine kinase, two ecto-nucleoside triphosphate diphosphohydrolase (NTPDases), a human transient receptor potential cation channel (TRPM4) and a human cell cycle protein CksHs1 [96,111,113–116]. In addition, a few other structures exist containing other POVs (angled bars, Fig. 5), including V₂ (Fig. 1B), V_{3I} (Fig. 1D), V_{4c} (Fig. 1E), and the unusual heptaoxidovanadate (V₇) and are summarized in Table 2. The remaining vanadium-containing protein X-ray structures in the PDB display only one vanadium ion and, although most contain orthovanadate (Fig. 1A) or other protonation states collectively called V₁, some other forms do exist such as V₂ and/or V₄. We first describe the six V₁₀-protein complexes and then the protein adducts with other smaller oxidovanadates including the two species with the binding of V₇ [113,117].

The proteins bind V₁ mainly as tetrahedral and trigonal bipyramidal vanadium oxidoanions; the latter mimic the trigonal bipyramidal transition states of phosphoryl transfer reactions [93,94,96,118]. A few structures in which the binding of the vanadium to a ligand associated with a protein have been reported, along with structures showing the binding of V₁₀ or other POVs to the putative active site of the proteins, mimicking the binding of vanadate in place of its endogenous substrate. Specifically, in 2006 the first interaction of V₁₀ with the enzyme phosphatase was crystallographically characterized [114]. The enzyme was a respiratory burst-inhibiting acid phosphatase from the Centers for Disease Control and Prevention Category A bio-terrorism agent *Francisella tularensis*. The enzyme is a prototype of a super-family of acid phosphatases and phospholipases C, which was known to be inhibited by vanadate. The phosphatase in question has a serine in the active site as well as a metal ion, but, in contrast to most known phosphatases [93,94], binds V₁₀ on the surface [114] and after it V₁ in its active site.

3.1.2. POV-protein complexes: V₁₀-protein adducts

A 2006 publication describes the *Francisella tularensis* Acid phosphatase A (AcPA) as a prototype of a unique superfamily of acid phosphatases and phospholipases C [114]. The phosphatase has in the active site a serine residue interacting with the metal ion through a *coordinative* bond and hence a mechanism that deviates from other phosphatases. In the PDB there is also a structure of *Francisella tularensis* AcPA obtained by crystallization at pH 6.0 bound to V₁₀ (PDB code 2D1G) shown in Fig. 6 [114]. This form of the crystal is not described in the publication [114], but the

structure is uploaded into the PDB. AcPA is a phosphatase that binds V₁, but V₁₀ was observed to also bind to the protein when the enzyme was incubated with vanadate added to the solution in the form of sodium orthovanadate (Na₃VO₄), which converted to V₁₀ under the crystallization conditions (pH = 6) [114]. Interestingly, the crystallized protein that binds V₁₀ also contains V₁ coordinated in the active site. The larger V₁₀ anion is interacting loosely with a group of positively charged amino acids at the surface of the protein. The phosphatase interacts with the V₁₀ through several H-bonds resulting in isolation of the V₁₀-protein complex (Fig. 6B). As shown in Fig. 6B, a lysine residue is found to cap the loosely bound V₁₀.

The interaction of V₁₀ to the respiratory burst-inhibiting acid phosphatase was not expected because V₁ is clearly the form of vanadate that is known to inhibit other phosphatases [94,93]. However, the inhibition of alkaline phosphatase by V₁₀ has also been reported [58]. The inhibition of this phosphatase was different than that observed by V₁ alone, even if the behavior cannot be related to the presence of V₁₀ only. In fact, although V₁₀ was stable under the conditions of the enzyme assay, some minor hydrolysis of the V₁₀ was observed [58]. These observations are consistent with the protein causing the partial hydrolysis of V₁₀ to V₁. This latter reaction has been observed for the growth inhibition of *Mycobacterium smegmatus* and *Mycobacteria tuberculosis* where speciation studies showed in both that V₁₀ was hydrolyzed more rapidly in media containing the bacteria [44]. Furthermore, growth inhibition of these bacteria was observed by the presence of V₁₀ much more potently than by V₁ [44] or similarly by monosubstituted V₁₀ derivatives such as V₉Pt and V₉Mo [50].

A second acid phosphatase with a {VO₄N} coordination environment is also reported in the PDB (PDB code 4QIH), but has not yet been reported in the literature [115] and, hence, provides little additional insights into the binding of the phosphatase-V systems, but a POV appears to be bound to it. In addition, the human prostatic acid phosphatase was previously reported to be inhibited not only by V₁ but also by the V₂; the former at neutral and basic pH and the latter at acidic pH values (Table 1) [48]. These reports support the interpretation that X-ray data show different oxidovanadates interactions with the prostatic acid phosphatase, depending on the experimental conditions [93,94].

Nucleoside triphosphate diphosphohydrolase 1 (NTPDase1) is an ectonucleotidase that catalyze the hydrolysis of γ - and β -phosphate residues of triphospho- and diphosphonucleosides [116]. By hydrolyzing proinflammatory ATP and platelet-activating ADP to AMP, it blocks platelet aggregation and supports blood flow. NTPDase1 hydrolyzes P2 receptors (P2Y and P2X), such as ATP, ADP, UTP and UDP, with similar efficacy. Investigating the structure and dynamics of this representative member of the eukaryotic NTPDase family was done using a variant of soluble NTPDase1 (PDB codes 4BRH, 3ZX2 and 3GQI for activated receptor) lacking a putative membrane interaction loop between the two lobes of the catalytic domain, as reported by Zebisch *et al.* [119]. Notably, V₁₀ was formed under the experimental conditions from orthovanadate. The complex structure of NTPDase1 (PDB code 3ZX2) shows that V₁₀ binds electrostatically to a highly positively charged loop that is involved in binding of the nucleobase. The crystal structure shows four independent copies of the protein, each of them bound to V₁₀ interacting with the same interfacial active site cleft of the enzyme that contains several lysine groups. As shown in Fig. 7A, the lysine network in the protein interacts through H-bonds with two of the V₁₀ faces bringing the two parts of the protein together. Binding on multiple faces of the V₁₀ is also observed in the crystal structure of NTPDase1 that bound both thiamine-phosphovanadate and V₁₀ (PDB code 4BRH) [120]. V₁₀ interacts both with the lysine residues and the NTP unit (Fig. 7B), organizing two parts of the protein around the V₁₀ molecule.

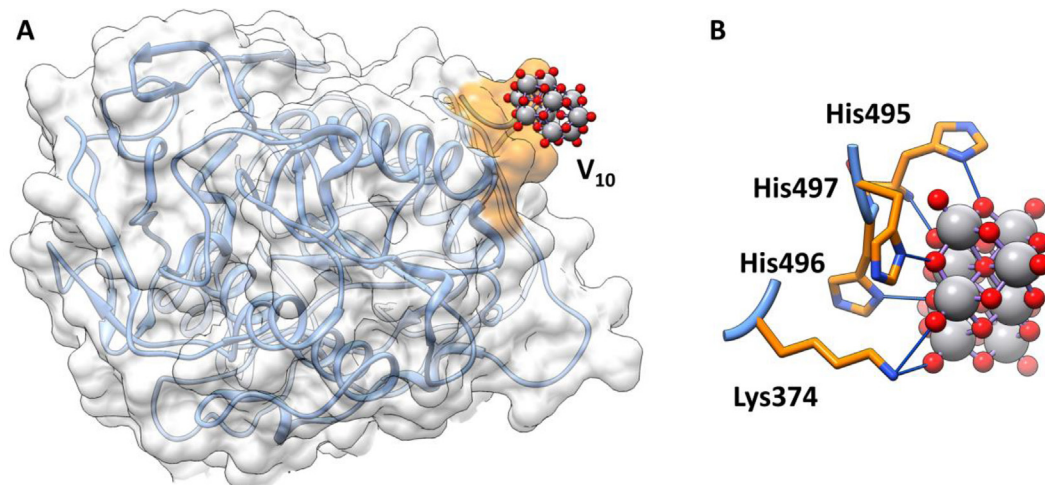


Fig. 6. Structure of *Francisella tularensis* AcPA (PDB code 2D1G) bound to V_{10} (added in form of orthovanadate and converted to V_{10}); A) full protein; B) close-up interaction of protein surface with V_{10} [114].

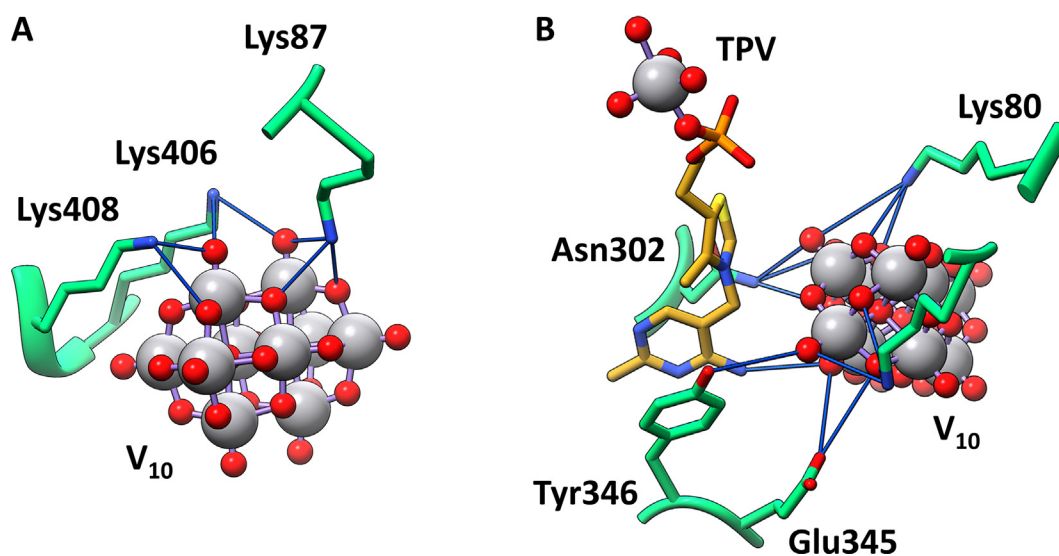


Fig. 7. Close-up view of the X-ray resolved binding sites of V_{10} in: A) apo-NTPDase1 (PDB code 3ZX2) [119]; B) NTPDase1 and thiamine-phosphovanadate (PDB code 4BRH) [120].

V_{10} also binds to an activated receptor tyrosine kinase in one known crystal structure (PDB code 3GQJ), shown in Fig. 8 [121]. Although kinases are complementary enzymes to phosphatases which interact strongly with vanadium salts, surprisingly few examples have been reported exhibiting any direct interaction of kinases with vanadate or POVs. The cAMP protein kinase that is removing phosphate groups from peptide substrates and assayed using the Kemptide substrate (Leu-Arg-Arg-Ala-Ser-Leu-Gly) was inhibited by V_{10} [91]. The mechanistic studies demonstrated that the competitive inhibition was caused by the Kemptide binding to the V_{10} anion and not V_{10} binding to the kinase. However, this report proves a rare and strong affinity for a peptide- V_{10} complex which is significantly stronger than the H-bonds observed in the V_{10} -dipeptide (Gly-Gly) complex that is crystallographically characterized [38]. Interaction of the activated receptor tyrosine kinase with V_{10} (PDB code 3GQJ) is therefore of great interest [121]. The receptor tyrosine kinase is a subclass of tyrosine kinases and not only a representative of kinases, but a protein involved in mediating cell-to-cell communication and controlling a wide range of complex biological functions, including cell growth, motility, dif-

ferentiation, and metabolism [122]. V_{10} was previously reported to initiate signaling on another tyrosine kinase receptor, the Type I Fcε receptor [108,123]. This receptor initiates intracellular signaling cascades ultimately leading to degranulation, the release of histamine, from cytoplasmic vesicles, as a key event leading to physical symptoms associated with an allergic response.

In the crystal structure of the activated receptor tyrosine kinase (PDB code 3GQJ) the V_{10} moiety is found at a turn on the protein surface and at the interface between PLC γ and FGFR1-3p protein domains [121]. The protein peptides are organized/wrapped around the V_{10} anion (Fig. 8). The authors reported the crystallization in a tris-buffered solution (pH 8.0) containing the protein, where Na_3VO_4 was added. However, the V_{10} anion was localized using idealized V_{10} coordinates placed where high electron density was present in the difference map. The buffered alkaline conditions of the crystallization would not normally favor formation of V_{10} , however in the deposited structure the authors assigned the observed density as V_{10} and optimized the geometry to fit. The V_{10} is found in close contact with a positively charged patch at the interface between FGFR1-3p and PLC γ chains (Fig. 8A) [121].

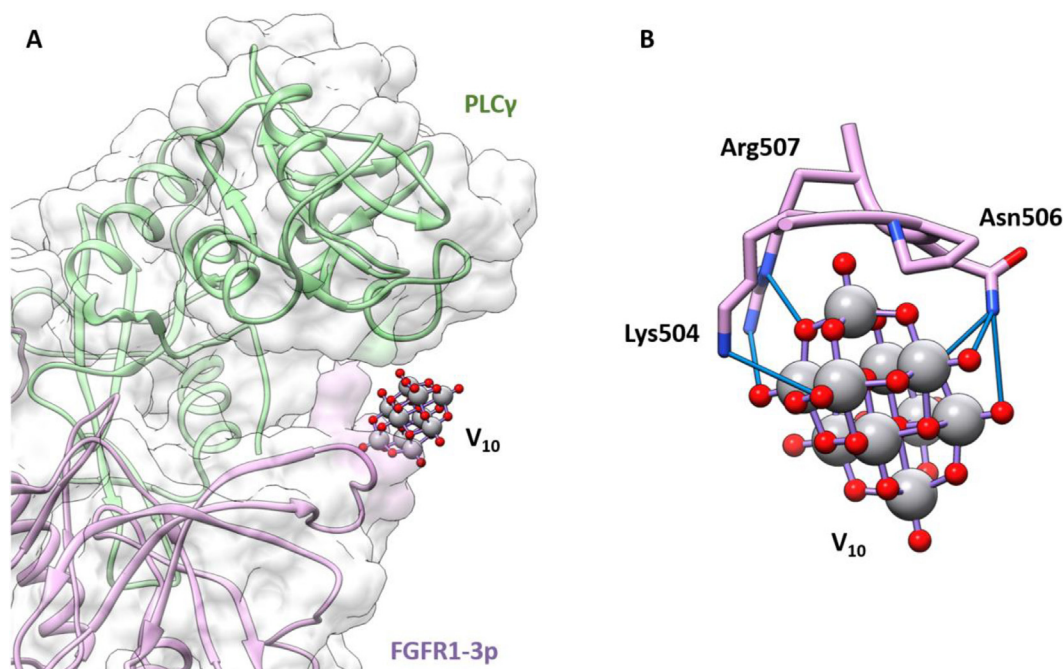


Fig. 8. Tyrosine kinase from activated receptor tyrosine kinase (PDB code 3GQI) [121]. A) interaction of V₁₀ at interface between FGFR1-3p and PLCγ chains; B) close view of the V₁₀ interaction with the FGFR1-3p chain. H-bond contacts are depicted with blue lines. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

Upon closer examination, the V₁₀ anion is H-bonding with two adjacent amino acid side chains, namely Asn506 and Arg507, the binding is further supported by Lys504 (Fig. 8B) interacting with a different V₁₀ surface part than observed with the apo-NTPDase1, NTPDase1 and thiamine-phosphovanadate shown in Fig. 7; i.e. the protein is interacting with the end of the oblong anisotropic V₁₀ anion rather than the side of the V₁₀. Given the difference in binding directionality and the use of idealized coordinates, it is possible that the cluster is actually a number of disordered smaller oxidovanadate ions that are known to form at pH 8 [41,124]. The small number of available protein complexes with V₁₀ makes it difficult to label this motif as an outlier. Such interpretation is especially difficult given the similarity in interactions between activated receptor tyrosine kinase (PDB code 3GQI) and the acid phosphatase structure (PDB code 2D1G) shown in Fig. 6. Although protein binding can stabilize V₁₀ at higher pH values [39] and some counterions have been found to significantly change fundamental properties of V₁₀ [125], we point to the unlikely formation of V₁₀ at pH 8, and the existence of an attractive interpretation that the electron density in the protein complex reported in ref. [121] can be attributed to one or more of the smaller rapidly equilibrating oxidovanadates.

Transient receptor potential channel (TRP) is a Ca²⁺-activated, non-selective cation channel that is permeable to Na⁺ and K⁺ and is found to depolarize the cell [126]. This process can be modulated by V₁₀. It is one of the eight melastatin-like transient receptor potential (TRPM) subfamily of TRP channels. TRPM family members are characteristically assembled with N-terminal TRPM homology regions (MHRs) and a C-terminal coiled-coil domain. This depolarizing modulation of cellular Ca²⁺ entry is important for cellular responses such as neuronal bursting activity, cardiac rhythm and the immune response. The cryo-EM structure was determined for the most widespread Ca²⁺-activated, non-selective (CAN) channel, human TRPM4, bound to both the agonist Ca²⁺ and the modulating V₁₀ (PDB code 5WP6). The V₁₀ was formed from a solution of 50 mM Na₃VO₄ adjusted to pH 2.0 from which 1 mM V₁₀ and 5 mM

calcium chloride was mixed with purified TRPM4 for a few hours before cryo-EM experiment. The cryo-EM structure of a human TRPM4 channel shown in Fig. 9A consists of a protein complex with Ca²⁺ and several V₁₀ [126]. The Ca²⁺ ions are necessary for formation of the crystals although the density for the Ca²⁺ was not discernible, the patch-clamp recorded data showed Ca²⁺ activation that can be blocked by either flufenamic acid or ATP⁴⁻ consistent with the Ca²⁺ being agonist and associated with the active site. The global view of TRPM4 in Fig. 9A shows eight V₁₀ moieties, four for each surface and internal characterized binding mode. Enlarged views of the two unique V₁₀'s binding modes are shown in Fig. 9B and Fig. 9C and show the high density of positively-charged residues surrounding each V₁₀. The V₁₀ at the surface sites are characterized by the interaction of four Arg, two Ser, two Lys and one Gln side chain (Fig. 9B). The internally stabilized V₁₀ is supported by three Arg, one Ser and one His side chain (Fig. 9C).

The crystal structure of the human cell cycle protein CksHs1 with kinase domain was shown to bind V₇ species, see Fig. 10 (PDB code 1DKT). However, as the authors described in the manuscript, they believed that the V-cluster in the protein was a V₁₀ cluster with three disordered vanadates [113]. This protein is involved in cell-cycle progression, which is largely regulated up or down by the activity of cyclin-dependent kinases (Cdks) [127]. The Cdks act by phosphorylation and several Cdks have cell-cycle functions in eukaryotic species. Two human gene homologs, CksHs1 and CksHs2, have been cloned and found to functionally substitute for CKS1 in *S. cerevisiae* [128]. Both phosphate and vanadate bound to CksHs1 provided information on CksHs1-ligand interactions. The vanadate binds tightly to Cks and was successfully used for crystallizing CksHs1 [113]. Two phosphate-binding sites are located near the dimer interface within a crevice formed by the dimer association (Fig. 10).

CksHs1 crystallizes as a dimer with phosphate, tungstate [113] or vanadate. However, since vanadate oligomerizes and forms V₁₀ in aqueous solution at acidic pH and V₁₀ interacts with an activated receptor tyrosine kinase, it is possible that a cluster would be asso-

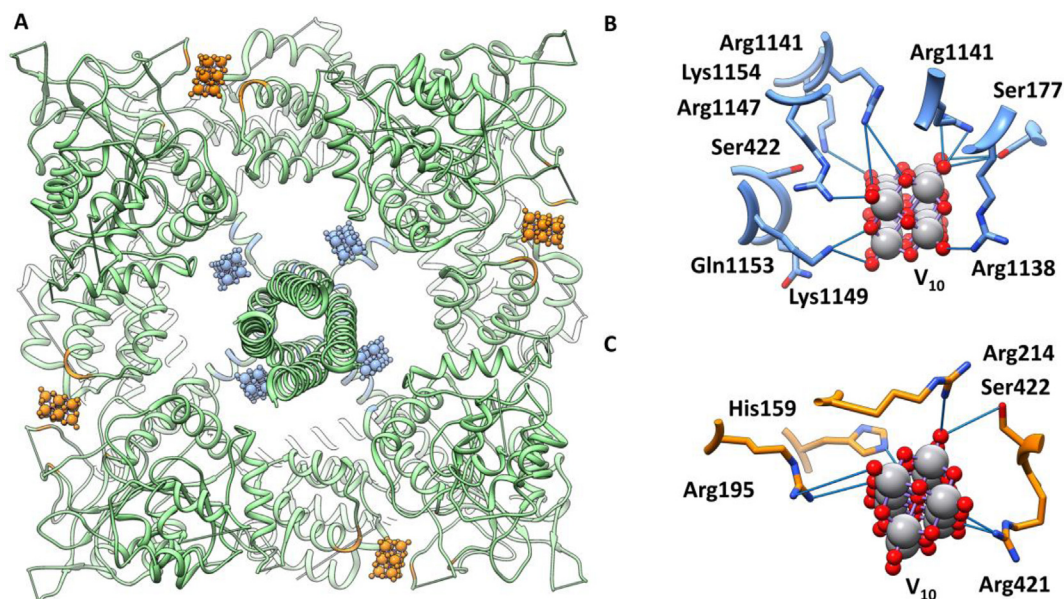


Fig. 9. Structure of the human TRPM4 (PDB code 5WP6) [126] channel shown in the complex with Ca^{2+} and V_{10} . A) global view; B) and C) close-up views of the two different binding modes of V_{10} with B) being the bound at the kink of the C-terminal helix (blue in A) and C) exposed at the interface of the N-terminal TRPM homology domain between the two adjacent subunits (orange in A). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

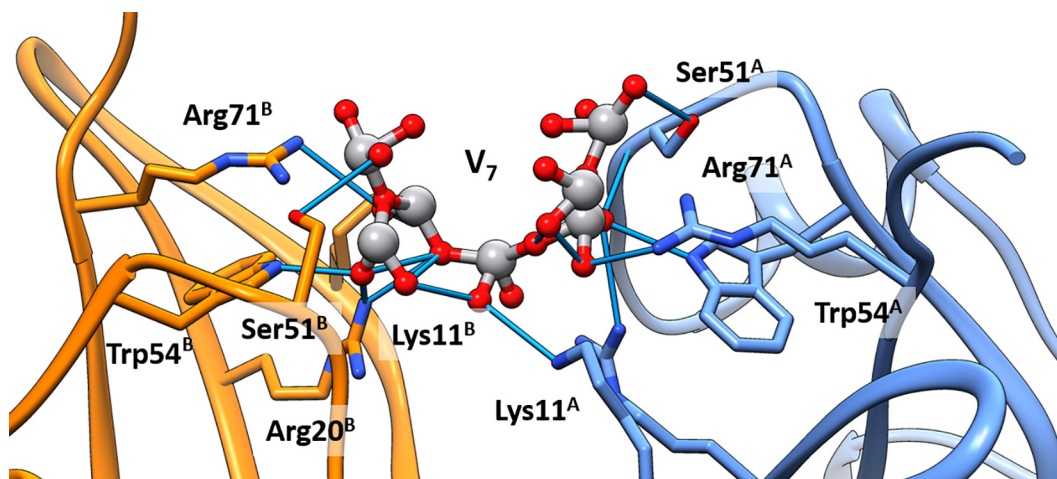


Fig. 10. Structure of human CksHs1 dimer (PDB code 1DKT) of the bound V_7 representing a partial V_{10} -cluster [113]. The authors assigned the protein complex to a Protein- V_{10} complex. Three disordered vanadate molecules would complete the putative V_{10} cluster, which on the basis of the electron density in the PDB is shown as V_7 [113]. H-bond contacts are depicted with blue lines. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

ciated with CksHs1. The authors noted a large positive region of electron density in their model, indeed larger than what is predictable for a mononuclear and even simple oligomeric species, which the authors presumed to be “metavanadate” given their crystallization conditions. As seen in the structure shown in Fig. 10, the observed electron density was modeled with seven V-atoms as a V_7 . Because V_7 is not commonly observed in aqueous solution [41], the presence of a V_7 in a protein complex would represent the report of a new oxidovanadate species. Therefore, we considered the possibility that the V_7 structure would be more accurately described as another POV, as described below.

In this highly symmetric (D_3) structure, a region of high electron density sits along the symmetry axis and this was formulated as $\{\text{V}_7\text{O}_{19}\}$ in the deposition for 1DKT in the PDB [113]. Unfortunately, distances between V and O centers and atom arrangements leading to the suggested coordination geometries are inconsistent with the formulation of a V_7 structure. One possible interpretation, sug-

gested by the authors, is that this POV is more likely to be V_{10} anion for which all the V-atoms are not fully occupied in the crystal structure. This interpretation is supported by the fact that the symmetry does not allow an asymmetric non-branching V_7 as formulated in the deposited structure and thus casting serious doubt on the interpretation of a novel V_7 species. Unlike in entry 3GQJ [121], where the authors chose to use idealized coordinates and parameters to model the V_{10} in the structure, in the 1DKT it was assumed that the actual species is V_{10} with some electron density missing. The missing electron density has likely been modeled as solvent water and thus the authors labeled the rest of the V_{10} molecule as a V_7 moiety to rationalize the diffraction data. In summary, this report does not support a new V_7 -protein complex, but an imperfect V_{10} -protein adduct missing some of the electron density needed to account for a complete V_{10} anion.

An alternative interpretation considers that the 1DKT structure, like 2D1G [114], contains more than one binding site for oxi-

dovanadates and, as such, could contain smaller species bound in addition to the postulated V_7/V_{10} . The smaller species bind in a shallow surface groove in positions occupied by phosphates in the corresponding phosphate-containing crystal structure. The V_7/V_{10} species appears to form hydrogen bonds with Lys, two Arg, and a backbone nitrogen of Ser in the modeling. Given the presence of smaller oxidovanadates in the structure, it is possible that the alternative interpretation of the V_7 is representing at least in part a mixture of smaller, disordered bound oxidovanadates with average electron density corresponding to a V_7 .

Although only six proteins were found to bind V_{10} , from the protein descriptions above some common patterns do emerge. There are fundamentally two types of binding sites: i) *surface binding*, where V_{10} binds non-specifically to the surface of the protein; and ii) *internal binding*, where V_{10} binds within a specific pocket into the three-dimensional structure of the protein. In several cases of surface binding, the V_{10} is bound to the protein along with a simple vanadate, V_1 . In each of these cases the V_{10} binding involves positive “patches” of the protein; the negatively charged POVs electrostatically interact with multiple positively charged side chains. In the larger V_{10} , there are more of these weak electrostatic interactions than in the case of V_1 , making the surface binding relatively stronger. In two of the cases, although they are still relatively close to the protein surface, the side chains of the protein's amino acids wrap around the V_{10} anion, somehow creating a specific solvent excluded pocket. In these internal binding cases, the anion is associated more specifically and solidifies the interactions between the protein and the large V_{10} . In the following section, we will describe interactions with smaller POV anions.

3.1.3. V-protein complexes: $V_2/V_3/V_4/V_7$ -protein adducts

In addition to V_{10} , there are several smaller POVs including V_2 , V_3 , V_4 (Fig. 1) and a V_7 that have been reported to form complexes with various proteins and have been investigated by X-ray diffraction methods. One of the first characterized structures of vanadate oligomers bound to proteins was the *E. coli* co-factor dependent

phosphoglycerate mutase (dPGM) and the structure with a linear V_4 (Fig. 1F) bound has been determined at a 1.3 Å resolution, PDB code 1E59 [129] (Fig. 11A).

The dPGM enzyme is a structural homolog of rat prostatic acid phosphatase from *Rattus norvegicus* and catalyzes the reaction of 3-phosphoglycerate (3-PG) to form 2-phosphoglycerate (2-PG). The mechanism goes through the phosphoryl transfer to form 2,3-bisphosphoglycerate (2,3-BPG) and a dephosphorylated dPGM intermediate. This X-ray structural determination of dPGM shows a linear V_4 bound in the active site [129] (Fig. 11B). This structure remains the only reported case of binding of the linear tetrameric POV to a protein. The studies with vanadate provided insight into the mechanism of this reaction.

In neutral solutions, a cyclic tetrameric oxidovanadate $[V_4O_{12}]^{4-}$ (Fig. 1E) is stable, while a linear oxidovanadate $[V_4O_{13}]^{6-}$ (Fig. 1F) has been reported up to pH 9.0 [130]. The authors, reporting the V_4 -dPGM structure, characterize the binding site of dPGM as basic, which is consistent with the interaction and stabilization of the linear oxidovanadate over the cyclic V_4 generally stable in aqueous solution, and observed in structures with PDB codes 1UZI and 1L7V [131,132] (Fig. 12). The structure of the V_4 -dPGM shows the two internal V-atoms (V_2 and V_3) well defined by the electron densities, whereas the two terminal V-atoms are disordered and modelled with only half occupancy. Thus, the structure may be a disordered V_2 or V_3 which are present in aqueous solution [130] and which have been observed in 1H2F, Fig. 13 [133], and 3F9B [134], where POVs bind to *Bacillus stearothermophilus* phosphatase PhoE and *Yersinia* protein-tyrosine phosphatase YopH, respectively. The highly charged binding site of dPGM forms a strong V-protein complex replacing the sulfate binding. This protein hence illustrates the versatility of the interactions of oxidovanadates with enzymes formed from solutions containing different V-species including V-Tris adducts (aqueous solutions of vanadate in Tris and other buffers will form V-buffer complexes as described previously in detail [51,135,136], however additional complexes in a crystallization medium may be important to generate the envi-

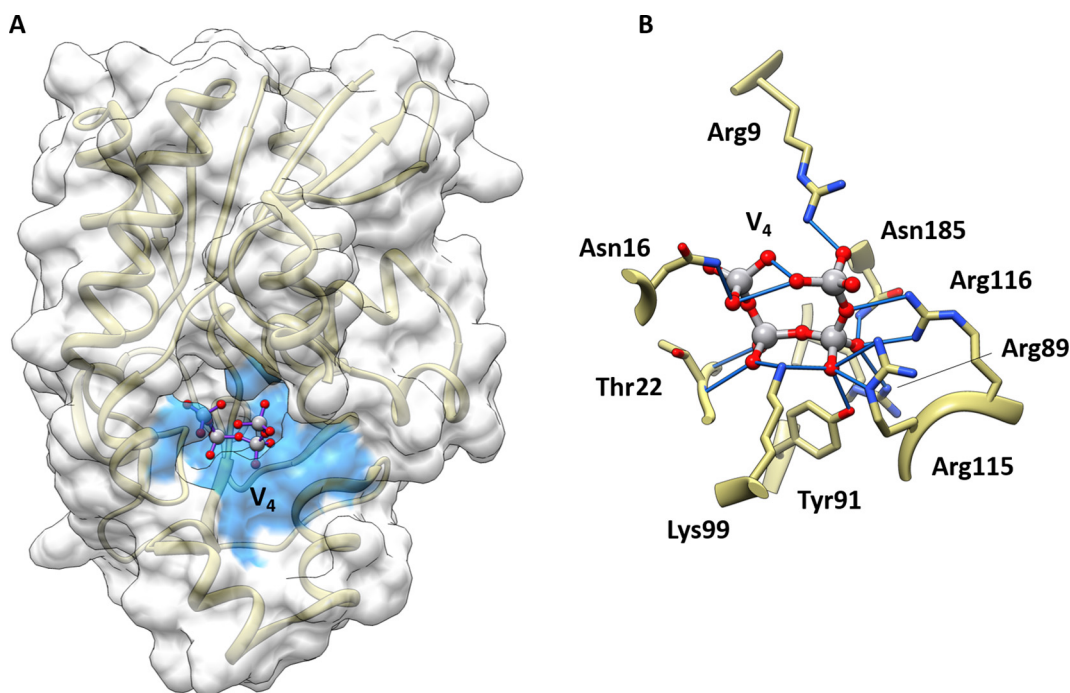


Fig. 11. The structure of linear V_4 bound to dPGM (PDB code 1E59). (A) Global view of the enzyme with V_4 in its active site; (B) close view of the H-bond network showing amino acid partners with O-atoms on the linear V_4 oxidovanadate.

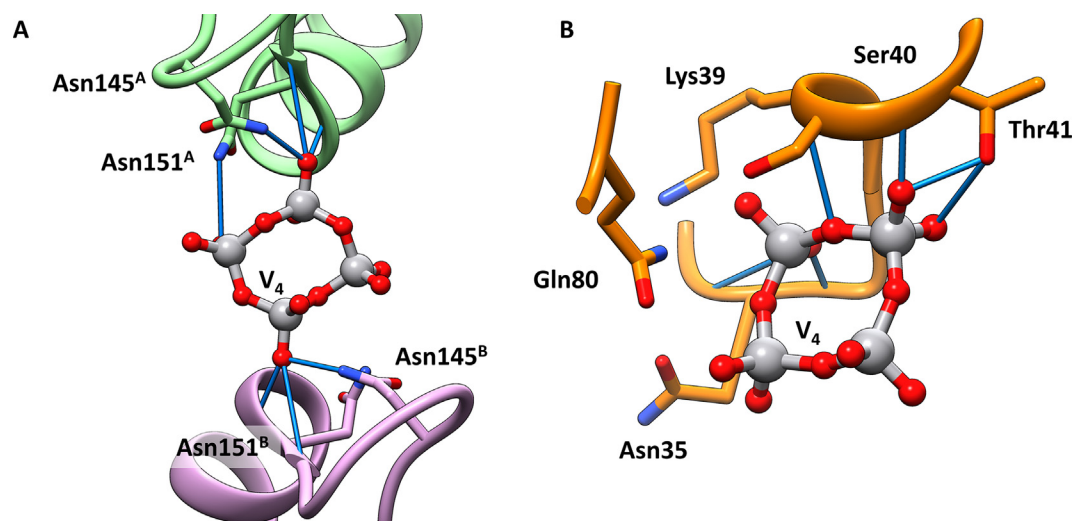


Fig. 12. Structure of the binding mode of cyclic V_4 toward: A) C3 exoenzyme for *Clostridium botulinum*, 1UZI; B) *E. coli* BtuCD protein, (PDB code 1L7V). H-bond contacts with electrostatics of V_4 the enzymes are depicted with blue lines [131,132]. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

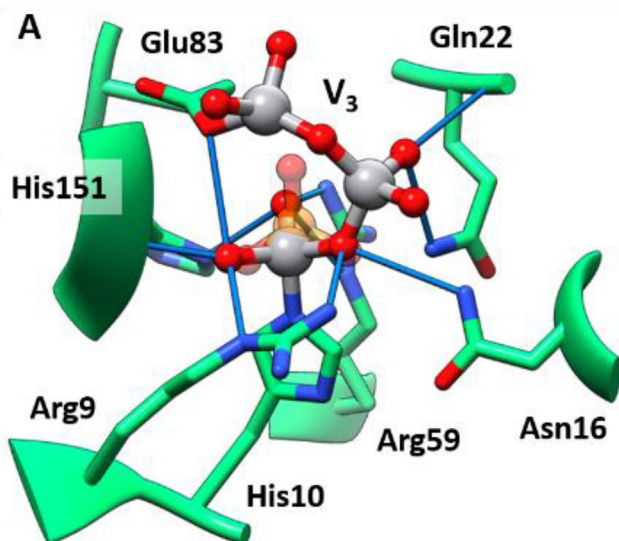


Fig. 13. Structure of the V_3 binding to the active site of phosphatase PhoE (PDB code 1H2F) along with the superposition of the native phosphate [133]. H-bond contacts are depicted with blue lines. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

ronment that allows a particular crystal to form). The linear V_4 is bound to dPGM in place of the four sulfate groups found in native dPGM structures, stabilized by an elaborate H-bond network, although the two internal V-atoms or sulfate groups are held tighter than the two terminal groups, Fig. 11B.

The two reported structures containing cyclic V_4 -protein interactions are those of the C3 exoenzyme for *Clostridium botulinum*, (C3bot1, PDB code 1UZI), Fig. 12A [131], and *E. coli* BtuCD protein (PDB code 1L7V), Fig. 12B [132]. In C3bot1, the cyclic V_4 interacts loosely with the exoenzyme displaying contacts with asparagine side chains and the protein backbone (Fig. 12A). The binding site is found between biological assemblies holding together adjacent symmetry-equivalent domains. It is interesting that vanadate (V_1) similarly serves to bring together adjacent symmetry-equivalent domains at both ends of the assembly. Vanadate here thus serves as electrostatic “glue” to hold together the adjacent

symmetry-equivalent domains. Similarly, the holding together of adjacent biological domains has been observed for other anisotropic POMs including the tellurium-centered Anderson polyoxido-tungstate (TEW, $[\text{TeW}_6\text{O}_{24}]^{6-}$), which is now used as an effective technique to promote protein crystallization [137–141].

In the *E. coli* BtuCD protein, an ABC transporter involved in B12 uptake [132], the cyclic V_4 unit has more interactions than in C3bot1 as shown in Fig. 12B. Specifically, contacts with a threonine side chain and a long interaction with a glutamine from a neighboring chain are formed, but most are with protein backbone in a helical portion of the structure.

The complex of V_3 with *Bacillus stearothermophilus* phosphatase (PhoE) [133], a member of the cofactor-dependent phosphoglycerate mutase superfamily, has been characterized by X-ray crystallography (PDB code 1H2F) [133]. The V_3 is coordinated both to catalytic residues His10 and Glu83 that are part of an elaborate H-bonding network (Fig. 13). The V_3 -PhoE complex was prepared through soaking of the PhoE-phosphate crystals prepared in 30.0% ethylene glycol, 25.0 mM AMP, and 55.0 mM sodium cacodylate buffer at pH 4.5 to which a solution of 50 mM NH_4VO_3 , 30% ethylene glycol, 20 mM sodium acetate at pH 5.0 was added [133]. Because this mixture is incubated for three days, it is likely that the solution under those conditions contains V_{10} (Fig. 2), as well as a previously characterized complex with ethylene glycol [142], and yet the crystal formed containing the less stable V_3 anion, perhaps because of the size constraints of the binding site. The image in Fig. 13 shows that V_3 replaces the phosphate, forming an extensive H-bonding network in the V_3 -PhoE complex. Hence the V_3 -PhoE adduct is significantly different than the dPGM structure containing V_4 [129], where V_4 was found to form a strong complex replacing the four sulfate groups present in the native enzyme (Fig. 11). The association of V_3 with the PhoE protein is the only reported case so far involving V_3 , with its unique availability of apoenzyme, product affinity, and phosphorylated intermediate structures. It seems likely that similar affinities will apply to other dPGM superfamily members. Indeed, modeling with adding V_3 into *E. coli* uridine phosphorylase was carried out and provided mechanistic details of this enzyme [117]. These observations are important because active, non-phosphorylated dPGM and inactive F26BPase structures as well as several other potential members of this superfamily of proteins have been identified in pathogenic organisms such as *Streptococci*, *Listeria*, *Staphylococcus aureus* and *Bacillus anthracis* [134].

A heptamer (V_7) was indicated in a deposited structure of *E. coli* uridine phosphorylase, PDB code 1RXS [117]. As mentioned above for the human cell cycle protein CksHs1 with PDB code 1DKT, a V_7 would represent a new species and was considered in detail by Arvai *et al.* [113] and now by us in preparation of this manuscript. Both Arvai *et al.* and us assigned the observed cluster to a V_{10} with missing electron density, even if Arvai *et al.* deposited the coordinates for the V_7 in PDB with code 1DKT. In the case of the structure of *E. coli* uridine phosphorylase, with PDB code 1RXS, electron density of a V -cluster assigned by Caradoc-Davies *et al.* [117] to the V_7 anion is found, even if no discussion was provided in the publication other than vanadate was used in the crystallization and for the solution of the structure. Given that the V_7 in 1DKT was just an electron density model, not an actual proposal of a structure, we favor an interpretation for the vanadium species to be something other than V_7 . However, since there are not enough details provided by Caradoc-Davies *et al.* [117], we cannot deduce whether this structure contains one or more of the smaller oxidovanadates or a V_{10} .

In addition to the V_3 , V_4 , and V_7 POV interactions with proteins, a few reports have described V_2 -protein adducts [94]. The systems range from simply pyrophosphate analogs to a diamond core complex associated with the vanadium. One very interesting case is one of the vanadate-*Yersinia* phosphatases structures (PDB code 3F9B) [134], where the dimer that is associated with the structure contains vanadium with two different coordination geometries [134]. The vanadium associated with the protein has a five-coordinate trigonal bipyramidal geometry whereas the free, terminal vanadium is four-coordinate and tetrahedral and is exemplary of the influence the protein can have on the coordination environment of oxidometalates [94].

In this section, we have discussed the binding of POVs described as a V_2 , V_3 , V_4 , and a putative V_7 . These former three structures show a greater diversity than those reported for the binding of V_{10} , whereas the reports with V_7 are likely a result of a decomposed or disordered POV or an incomplete electron density. In all these studies, complementarity between protein and POVs is high for crystals to form and in some cases so high that even less-stable POVs can give rise to tight V -protein adducts.

3.2. POVs and muscle contraction

According to Tables 1 and 2, V_{10} interacts with several proteins including those involved in the mechanism and regulation of muscle contraction – myosin, actin and Ca^{2+} -ATPase [39,49,56]. It is common for V_{10} to interact with ATP binding proteins, but it also interacts and inhibits other enzymes such as membrane channels and RNA ribonucleases (Tables 1 and 2). Sarcoplasmic reticulum (SR) calcium pump has proven to be an excellent model to study toxicology effects of POVs on P-type ATPases, such as the Na^+ , K^+ -ATPase and Ca^{2+} -ATPase. These ion pumps are involved in essential ions homeostasis, such as Na^+ , K^+ and Ca^{2+} , therefore regulating several key cellular processes. Although Table 1 represents only data since 1989, it has been known since 1982 that V_{10} interacts with SR Ca^{2+} -ATPase [84,143]. Thus, V_{10} has a higher inhibition capacity compared to V_1 alone and different abilities to induce changes in the structure and function of the enzyme, as described, for instance, for myosin, actin and calcium pump [39,49,56,99,144]. Recently, another POV, $Cs_{5,6}H_{3,4}PV_{14}O_{42} \cdot 12H_2O$ (PV_{14} , Fig. 11), was described to inhibit P-type ATPases, being particularly more potent than V_{10} ($IC_{50} = 15 \mu M$) in the inhibition of the Ca^{2+} -ATPase ($IC_{50} = 0.6 \mu M$) [57].

Regarding of actomyosin ATPase inhibition by V_{10} , the first studies were described in 1988, using flow microcalorimetry [145]. In 2004 [49], it was reported that such an inhibition implies a myosin binding site different from ATP site, and that V_{10} inhibi-

tion is non-competitive. These studies clearly demonstrated that, unlike V_1 , V_{10} species can strongly inhibit the myosin actin-stimulated ATPase activity with an IC_{50} of 6.1 μM for V_{10} , whereas no inhibitory effects were detected for V_1 up to 150 μM [49]. It was proposed, combining molecular docking simulations with kinetic studies, that V_{10} interacts with the phosphate-binding domains in the vicinity of the designated “back-door” binding site (Fig. 14) [49,104]. Thus, an intermediate myosin-MgATP- V_{10} complex is formed, which inhibits the myosin-actin interactions associated with ATP hydrolysis by actomyosin, and the blocking of the muscle contraction process most likely occurs in a pre-hydrolysis state [49,99,104]. Note that the walker A motif (corresponding to the P-loop in myosin), as well as the ABC (ATP-binding cassette) ATPases, were also described as an anion-binding domain that can interact with this POV with high affinity [92,104].

In contrast to the examples described above, for a long time there was no available information on the molecular interaction of V_{10} with actin, another major protein of the contractile system of muscle cells. The first report describing the interaction between V_{10} and actin was published in 2006, suggested that G-actin, the globular monomeric form of actin, stabilizes decavanadate species by increasing the half-life time of decomposition from 5 to 27 h [39]. V_{10} inhibits the rate of G-actin polymerization into F-actin (polymerized form of actin), with an IC_{50} of 17 μM [39], suggesting that it would affect cytoskeleton structures responsible for many processes of relevant biological significance. These studies were further explored and described protein cysteine oxidation together with vanadate reduction upon V_{10} incubation with actin [63]. Additionally, it was also shown that V_{10} binding induces an opening of the G-actin cleft. V_{10} was able to oxidize Cys374 in F-actin and one of the protein core cysteine residues, presumably Cys272 [63,64]. In fact, the interaction of V_{10} with G-actin results in a vanadate(V) reduction to oxidovanadium(IV) ion. Moreover, in the presence of ATP in the medium assay, the reduction of V^V to V^{IV} is prevented and all five G-actin cysteine residues remain in their reduced form [63,109].

In 2021, molecular dynamics and docking studies were described in order to attain an understanding at molecular level of the V_{10} -G-actin interaction, rationalizing the results obtained during the last 15 years by several spectroscopic techniques [64]. Four binding sites of V_{10} were identified, named α , β , γ , and δ , the site α being the catalytic nucleotide site located in the cleft of the enzyme at the interface of the subdomains II and IV. The site α is stabilized by H-bonds with Gln59, Arg62, Tyr69, Thr204, Arg207, and Arg211 residues. The site α is more stable than site β by more than 20 kcal·mol⁻¹ and much more for sites γ and δ [64].

A slight opening of the nucleotide binding site α is predicted after the interaction with V_{10} which varies the cleft opening (defined as the $C(\alpha)_{Lys67}-C(\alpha)_{Val339}-C(\alpha)_{Thr203}$ angle) from 28.6° in the apo form to 35.6° in the adduct with V_{10} , with a rearrangement of the H-bonds. The behavior of V_{10} was compared with the bulkier isostructural polyoxidoionobate (PONb) Nb_{10} (volume of 594.3 Å³ vs. 520.6 Å³ of V_{10}) which is more stable at the site β [64].

ATP is found at the site α , the enzymatic cleft of G-actin, and this ATP binding is more favorable by 14.1 kcal mol⁻¹ than V_{10} . MD studies indicated that the presence of ATP hinders the V_{10} binding, which is forced to reach the site β . Notably, the competition is less important for Nb_{10} because this POM shows a higher affinity for site β than for site α , suggesting that both POVs and PONbs could contemporaneously bind to G-actin, making their action synergistic, a finding that should be possible for other POMs-protein systems (Fig. 15) [64].

Finally, in the study mentioned above, it was demonstrated that the binding mode of $V^{IV}O^{2+}$ ion, formed upon the V_{10} reduction by Cys residues, is in the catalytic site α with (His161, Asp154) coor-

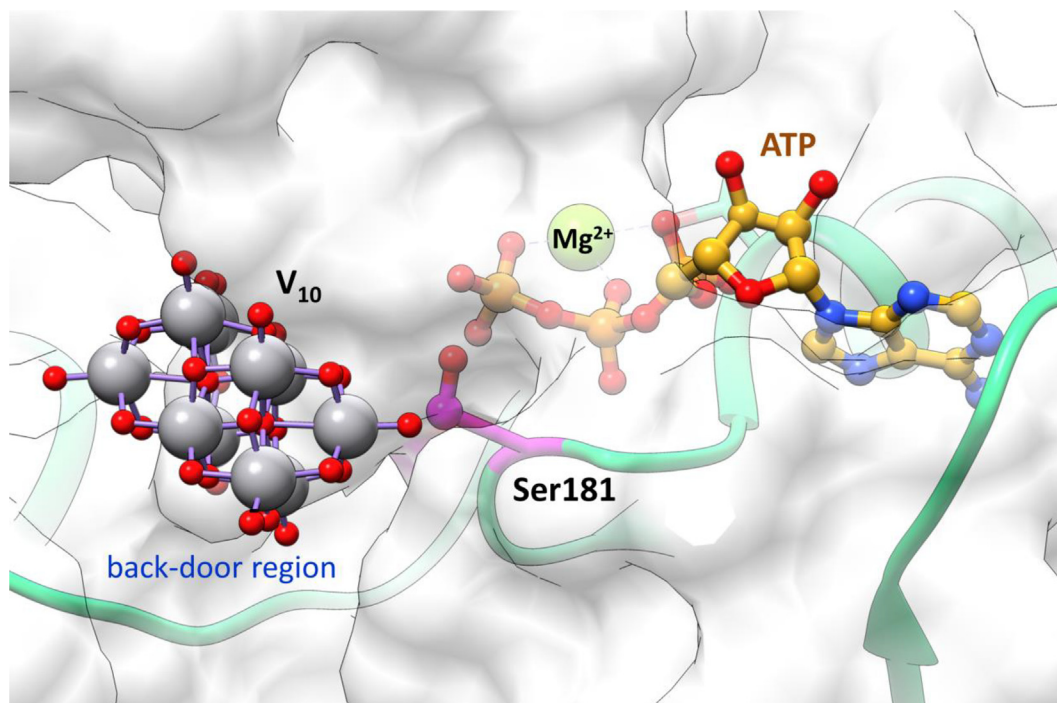


Fig. 14. Docking solution of V_{10} interacting with myosin at the phosphate-binding domain, in the vicinity of the nominated “back-door” binding site (adapted from ref. [104]). Color code: V, grey, O, red. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

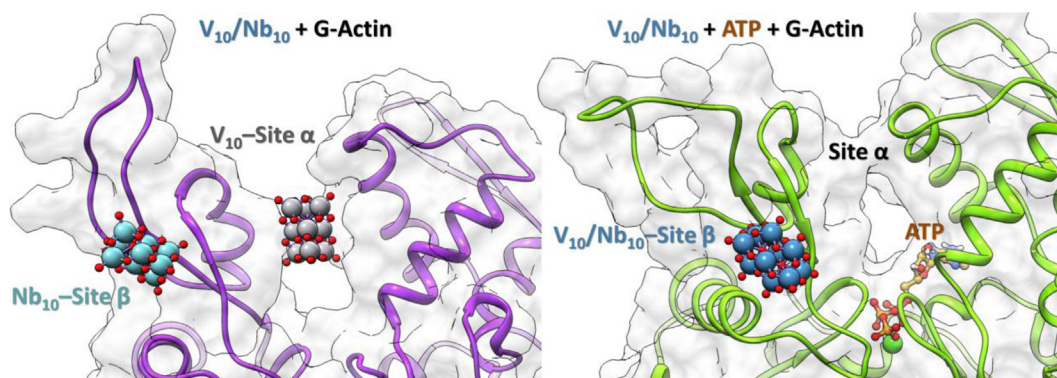


Fig. 15. V_{10}/Nb_{10} actin interactions predicted by molecular dynamics and docking studies (adapted from ref. [64]). Color code: V, grey; Nb, blue; O, red. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

dination; interestingly, this adduct overlaps significantly with the region where ATP is bound, accounting for the competition between V_{10} and $V^{IV}O^{2+}$ with ATP, observed by EPR spectroscopy [64].

3.3. POVs' interactions with serum proteins

After the administration of a POV, it may or may not reach the bloodstream intact, depending on the mode of administration [146]. The interaction of a POV with serum proteins is most likely to occur with intravenous administration or if the compound easily penetrates the cells membranes required to reach the blood, where POV would be associated with the serum fraction. In serum, a POV encounters several biomolecules and, in particular, proteins like human serum transferrin (HTf) and human serum albumin (HSA). Transferrins are a group of single-chain glycoproteins having 700 amino acids with a mass of ca. 80 kDa with a concentration in blood around 30 μM [147,148]. In human serum, HTf transports not only iron [149], but also other metal ions such as Bi, Al, Ru,

Mn, Ni, and V in its three oxidation states stable in biological systems (+III, +IV, +V) [61,150]. Considering the iron amount in the two specific sites in the N- and C-terminal, the concentration of available sites for metal binding is ca. 34 μM [151]. HSA is the most abundant blood protein with a concentration around 640 μM [148] and transports, beside essential and toxic metal ions and their compounds [152], fatty acids and endogenous and exogenous ligands [153].

It was recently described that V_{10} interacts both with HSA and HTf, presumably through non-covalent binding (see section 3.1), but surprisingly (2-hepH) $_2$ [{Co(H $_2$ O) $_5$] $_2$ V $_{10}$ O $_{28}$] $_2$ ·4H $_2$ O ($V_{10}\text{Co}$, 2-hepH = 2-hydroxyethylpyridinium) exhibits high affinity toward transferrin but does not interact with albumin [43]. It was supposed that the Co^{II} centers coordinated to {VO} groups of V_{10} occupy the binding sites of V_{10} , blocking its interaction with the albumin protein [43]. In contrast, HTf shows interaction with $V_{10}\text{Co}$, but the reasons for this different behavior should be examined in detail at an experimental and computational level. The different behavior of $V_{10}\text{Co}$ was also shown by two model pro-

teins: in proteinase K, $V_{10}Co$ interacts more strongly than thau-matin [43].

3.4. POVs and diabetes

According to the WHO, about 400 million individuals world-wide have diabetes, which can cause heart disease, stroke, kidney damage, and even death [154]. Insulin, which is produced by β -cells found in pancreatic islets, regulates the metabolism of carbohydrates. In 2021, it will be 100 years since insulin was discovered by Banting, Best and McLeod at the University of Toronto [28,155,156]. Vanadium has also long been associated with diabetes; in 1899, French scientists reported the blood glucose-lowering effect of vanadium [157]. However, it was only by 1979 that it was shown that vanadium salts exhibit insulin-enhancing effects, leading to a growing interest in vanadium compounds for the treatment of diabetes [158–160]. Since then, insulin-like effect of various vanadium compounds, mainly oxidovanadium(IV) sulfate and organic vanadates have been described [146]. One organic vanadium compound, bis(ethylmaltolato)oxidovanadium(IV), entered Phase I and II clinical trials [156]. However, vanadium coordination complexes are not the purpose of the present review, and the interested reader should address that elsewhere [158,161,162]. Therefore, the antidiabetic potential of POVs and vanadium-substituted POMs is described here.

Certain polyoxidometalates are known to have insulin-enhancing properties [163]. Also, some POVs, for instance V_{10} , have been shown to enhance glucose uptake in rat adipocytes [7]. Moreover, a hybrid decavanadate compound namely metformin (Metf) decavanadate, $(H_2Metf)_3V_{10}$, showed hypoglycemic and lipid-lowering effects in animal models of both insulin-dependent (type 1) and insulin-independent diabetes (type 2) [8] beyond those observed with metformin itself. The insulin-mimetic effect of benzylamine-decavanadate $(C_7H_{10}N)_6V_{10}$ was also studied [9]. As with $(H_2Metf)_3[V_{10}O_{28}]$, $(C_7H_{10}N)_6V_{10}$ stimulated glucose transport and normalized the lipid profile. Also recently, a second compound, $[DMAPH]_4(NH_4)_2V_{10}$ (DMAPH = 4-dimethylaminopyridinium), showed hypoglycemic and lipid-lowering effects [164].

The exact POVs mechanism remains unknown for both type 1 and 2 diabetes. However, the POVs insulin-enhancer effect might, among others, derive from V_{10} inhibition of protein tyrosinase phosphatase, an enzyme catalyzing the dephosphorylation of the insulin receptor, which turns off insulin signaling. It was also suggested that $(H_2Metf)_3V_{10}$ interferes with the mitochondrial electron transport chain, forcing lipid oxidation. In fact, V_{10} was previously described to target mitochondria strongly affecting oxygen consumption and mitochondria membrane depolarization [106], and to induce necrotic death in neonatal rat cardiomyocytes through mitochondrial membrane depolarization [165]. More recently, it was referred that a mixed POM containing vanadium, $K_{11}H[(VO)_3(SbW_9O_{33})_2]$, which exerts hypoglycemic effects in streptozotocin-induced diabetic rats, shows hepatic protective properties and stimulation of insulin synthesis [166]. These findings clearly show that V-containing POMs are not only insulin-enhancers, as suggested before for other vanadium compounds, but also stimulate the insulin synthesis [146,167].

Besides hyperglycemia and hyperlipidemia, oxidative stress contributes to the pathogenesis of diabetes. Recently, the oxidative stress in multiple tissues of alloxan-induced diabetic rats (type 1 diabetes) and the effect of $(H_2Metf)_3V_{10}$ on this stress was evaluated [168]. It was described that $(H_2Metf)_3V_{10}$ reduced oxidative stress by positively affecting the nuclear factor erythroid 2-related factor 2 (NRF2), a transcription factor responsible for cellular antioxidants. Thus, POVs mediated activation of NRF2 induces the expression of reduced glutathione (GSH), superoxide dismutase (SOD), and catalase (CAT), restoring cellular antioxidants defenses.

POVs inhibition of α -glucosidase, which converts carbohydrates into glucose, is another strategy for the treatment of diabetes. About 10 mixed polyoxidomolybdates (POMos) containing vanadium have been described to exhibited inhibitory effects on α -glucosidase [169,170,171]. One of them, $H_6[PMo_9V_3O_{40}]$, exhibits higher activity (IC_{50} , 9.6 μM) than acarbose (IC_{50} , 38.2 μM), a clinically used antidiabetic drug and α -glucosidase inhibitor [172]. For comparison, higher potency POMs α -glucosidase inhibitors include $H_6[P_2Mo_{18}O_{62}]$ (IC_{50} = 0.17 μM) and $H_3[PMo_{12}O_{40}]$ (IC_{50} = 6.1 μM)

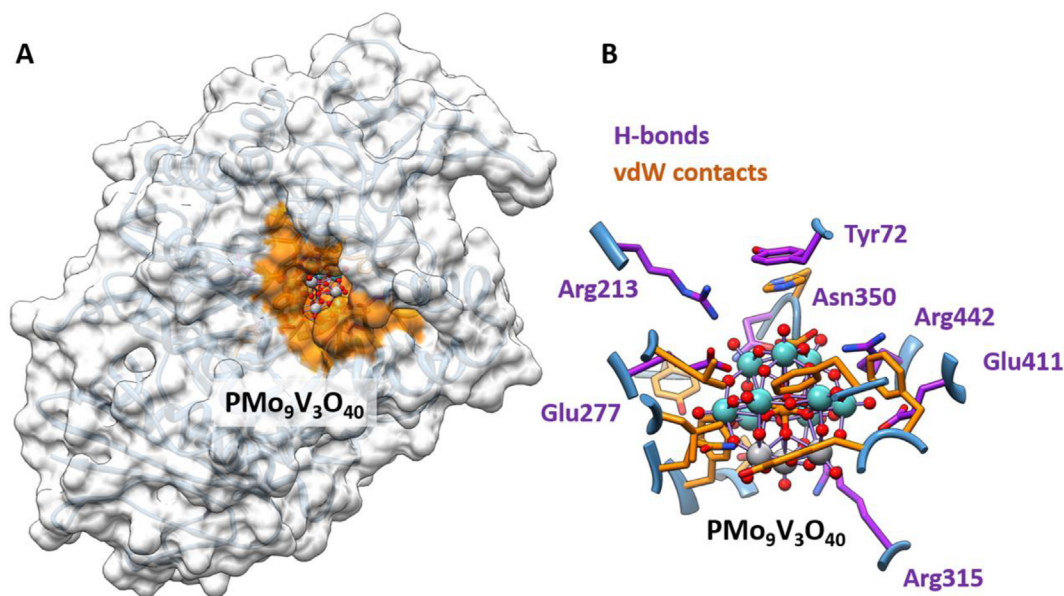


Fig. 16. Representation of the docking solution of $[PMo_9V_3O_{40}]^{6-}$ bound to α -glucosidase. A) global view of the enzyme with $[PMo_9V_3O_{40}]^{6-}$ in its active site; and (B) close view of the secondary interactions network. H-bond donors depicted in purple and vdW interactors in orange (His351, Thr306, Tyr347, Gln279, Tyr158, Asp352, Phe159, Phe178, Leu219, Val216, Phe303). Adapted from ref. [171]. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

[170,173]. For $H_6[PMo_9V_3O_{40}]$ the mixed type of inhibition was described, and the structure predicted from docking studies is shown in Fig. 16.

3.5. POVs and oxidative stress responses

As referred in the above section, the mode of action of POVs in diabetes might be due to, inter alia, direct or indirect changes in cellular oxidative responses. However, owing to the low stability of V_{10} under physiological conditions, it is expected that it dissociates into minor oxidovanadates namely V_1 , V_2 and V_4 , presumably involved in the biological effects observed. The detoxification mechanism proposed for vanadate involves bioreduction of vanadate to oxidovanadium(IV) by glutathione (GSH) [174]. Therefore, GSH is an important cellular antioxidant defense system and directly or indirectly regulates the levels of ROS [175,176]. On the other hand, in Fenton-like reactions vanadate is reduced to $V^{IV}-O^{2+}$ with production of OH^\cdot [177,178]. However, vanadium prooxidants effects are not the purpose of the review and the reader is referred elsewhere [146,167]. Since the beginning of the 21 century, the Aureliano research group has been performing POVs *in vivo* studies, mainly with V_{10} , in order to determine the contribution of V_{10} species to toxic effects observed in fish at room tem-

perature where V_{10} stability is higher [40,55,179,180]. Following V_{10} *in vivo* administration, several oxidative stress parameters were ascertained in fish: reduced GSH content, overall rate of ROS production, lipid peroxidation, and antioxidant enzyme activities [40,55,179,180]. It was concluded that V_{10} clearly induces different and, in many times, opposite effects than the ones observed for vanadate [181,182]. For instance, upon V_{10} exposure, an increase in GSH content and ROS production was observed, as well as a decrease in mitochondrial antioxidant enzymes activities such as SOD and catalase activities, whereas opposite or no effects were observed for vanadate [40,55]. Therefore, it was suggested that POVs species exposure follows different pathways than vanadate for the generation of reactive oxygen species and that they interfere differently with some of the enzymes involved in antioxidant defenses in cells [182,183].

In summary, studies describing POVs, mainly V_{10} interactions with proteins (Fig. 17) include, among others: 1) proteins associated with diabetes such as phosphatases, tyrosine kinases and glucosidases; 2) membrane channels such as TRPM4 channels, ATP sensitive channels, porin VDAC modulator and P2X receptor; 3) muscle contraction proteins such as myosin and actin; 4) serum proteins, such as albumin and transferrin; 5) P-type ATPases such as Ca^{2+} -ATPase pumps; 6) mitochondria proteins such as complex

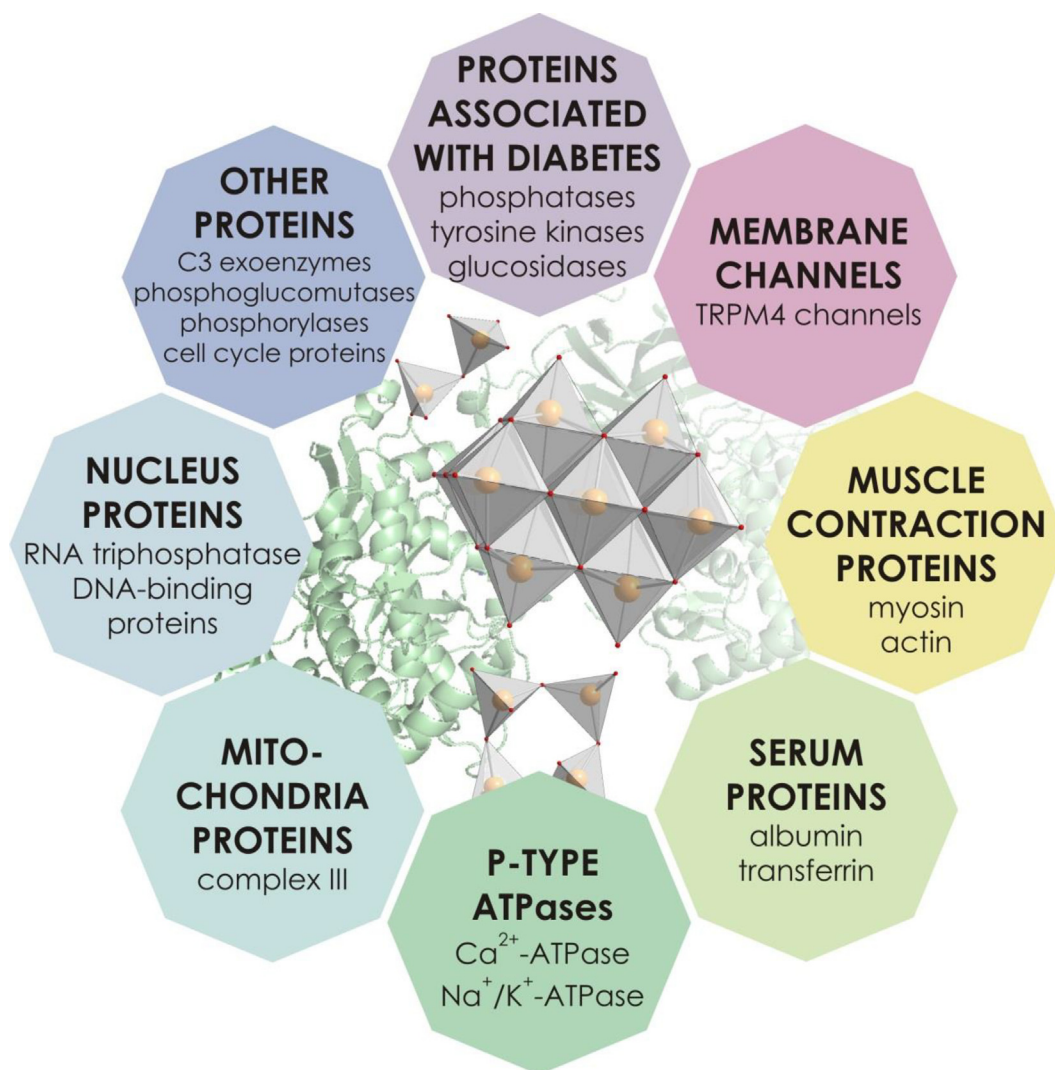


Fig. 17. Scheme of the described interactions of proteins with POVs such as V_2 , V_4 and V_{10} . POVs' interactions include muscle contraction proteins, channels, receptors and ion pumps, mitochondria proteins, nucleus proteins, serum proteins as well as those associated with diabetes, among others.

III and 7) nucleus proteins, such as RNA triphosphatase and DNA-binding protein. Smaller POV proteins interactions include, among others: 8) phosphoglucomutases; C3 exoenzymes; cell cycle proteins and phosphorylases (Fig. 17).

4. Conclusions and perspectives

In summary, we have described from a qualitative and quantitative point of view information about the interaction of several POVs, particularly V_{10} , with several key cellular proteins including phosphatases, NTPDases, TRPM4 channel, tyrosine kinases, myosin actin, transferrin, albumin and glucosidases, among others. Considering the methods available to determine speciation and mode of actions, this subgroup of POMs is likely to receive much more interest in the near future as probes for enzymes and in biomedical applications. The application of POVs, being pure, POVs-based hybrids and/or nanomaterials containing POVs in various biological systems, is a rapidly growing branch of science.

The understanding of the POVs interactions with proteins remains the key step for elucidating the biomedical mechanism of action of these compounds, and in this review, we examined both the molecular interactions through reported POV-protein structures from X-ray crystallographic reports as well as studies characterizing these interactions using computational methods. In spite of the fact that simple vanadate salts are often used for the experimental treatment, the POVs can form under the conditions of the studies. In the reported cases, POVs seem to exert their cellular effects by interacting with the numerous proteins in the respiratory chain and the cytoskeleton, with the ionic transport systems, as well as with DNA, among others, and the molecular interactions are affecting structure and function.

Besides hydrogen bonds, electrostatic and van der Waals interactions contribute to POVs binding to proteins. The X-ray structures of V_{10} -proteins show that there are two groups of POV-protein interactions, those located at the surface or at the interior of the proteins. In both cases, the binding sites are characterized by complementary interactions of a positive patch on the protein surface where the V_{10} will form several H-bonds and electrostatic salt bridges. The positively charged residues include a variety of Lys, Arg, and His and in an organized manner complementary surrounding the V_{10} anion, strengthening the surface interaction. Of particular interest is the V_{10} myosin back-door interaction and also its reduction by actin that is prevented by the native ligand MgATP. In spite the low stability of POVs at physiological pH, the half-life of V_{10} is increased in serum, suggesting an interaction with proteins that can stabilize its structure. Some proteins, such as actin, improve 6-fold the stability of V_{10} .

Examples of less common POVs forming V-protein adducts were also summarized. These systems are very interesting in that these include interactions of vanadate with kinases, signaling molecules and phosphoglucomutases and phosphoglyceratemu-tases and the formation of V_2 -protein, V_3 -protein, V_4 -protein, as well as V_1 -protein adducts was demonstrated. However, there are two reported V_7 -protein complexes, whose electronic density is often diffuse and does not lead to clear solutions. In one case the study was exhaustive and showed that the system is not likely to involve a new V_7 POV, but the results instead are better explained by V_{10} or a combination of smaller oxidovanadates.

A couple of POVs is actually binding the less stable linear V_3 and linear V_4 , which are not observed in aqueous solution at pH 7. Interestingly, in these cases the POV-proteins interactions are sufficiently stable and the result is the formation of complexes between proteins and uncommon and less stable POVs present only in small amounts from the crystallization media. The reported V-protein complexes must bind the V-cluster very tightly and be

stabilized by both primary and secondary non-covalent interactions.

POVs, especially modified V_{10} , showed promising antidiabetic effects in animal models of type 1 and 2 diabetes. However, other activities have recently been reported including antimicrobial activities. V_{10} and Pt and Mo mono-substituted V_{10} were found to inhibit growth of *Mycobacteria smegmatis* and/or *Mycobacteria tuberculosis*. Furthermore, the POVs were reported to initiate signaling and, in this case, it was not only V_{10} but also multivalent V_{14} and V_{15} POVs that were active.

In summary, this review demonstrates the quantity and quality of information now available characterizing the POVs' interaction with proteins and enzymes. Studies reported have used a range of methods to determine speciation and mode of actions. Undoubtedly, the POV subgroup of POMs is likely to receive much more interest in the near future given their ability to serve as probes for enzymes and their direct effects in biomedical applications. With so few POV-protein structures reported and the known ability of POMs to assist in crystallization, it is very likely that the number of reported POV-protein structures will increase. Moreover, we also predict that the future will show that a number of different hetero-POVs will be biologically active. Such an approach will allow for obtaining not only clusters with mixed metal ions in addition to vanadium atoms, but also multivalent POVs, i.e. containing both V^{IV} and V^V . As was illustrated in a few available studies, the structural framework from the vanadium will be maintained but substitutions will allow for more diverse redox chemistry and biological activities. The future is indeed bright for POVs.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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