The Canadian Mineralogist Vol. 57, pp. 245-253 (2019) DOI: 10.3749/canmin.1800069

IDENTIFYING PROTONATED DECAVANADATE POLYANIONS

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Abstract

The decavanadate anion, $[V_{10}O_{28}]^{6-}$, is a common constituent in vanadate minerals and synthetic materials, and both protonated, $[H_xV_{10}O_{28}]^{(6-x)-}$, and mixed-valent, $[(V^{4+}_xV^{5+}_{10-x})O_{28}]^{(6+x)-}$, varieties also occur. Here we examine recent structure refinements containing protonated decavanadates in order to develop simple criteria to help identify specific O_D atom(s) belonging to (OH) groups of the decavanadate polyanion. Non-protonated O atoms on the surface of the decavanadate polyanion (*i.e.*, [1]-, [2]-, and [3]-coordinated O atoms) have incident bond-valence sums from V atoms in the range \sim 1.7–2.0 ν u. Protonated O atoms on the exterior have incident bond-valence sums from V atoms in the range \sim 1.2–1.5 ν u and are either [3]-coordinated or [2]-coordinated by V atoms. All hydrogen bonds are directed toward a neighboring decavanadate polyanion in which the O_A (acceptor) anions are 2.61–3.08 Å from their respective O_D (donor) anions on the adjacent polyanion, and there is a linear relation between the $O_D \dots O_A$ distance and the incident bond-valence received by O_D from the bonded V atoms. For well-refined structures, the incident bond-valence sums from V atoms and the associated $O_D \dots O_A$ distances suffice to identify protonation of the decavanadate polyanion and the specific anion(s) that is (are) protonated. These criteria show that one structure reported as protonated is not protonated, and one structure reported as containing a mixed $V^{4+/5+}$ protonated decavanadate polyanion contains only V^{5+} and is not protonated, and another structure also reported as containing a mixed $V^{4+/5+}$ protonated decavanadate polyanion contains both V^{4+} and V^{5+} but is not protonated.

Keywords: decayanadate polyanion, vanadate minerals, mixed-valence $V^{4+/5+}$, protonation, bond valence.

Introduction

The $[V_{10}O_{28}]^{6-}$ decavanadate polyanion is a cluster of eight (VO₆) octahedra with [1+4+1] coordination and two (VO₆) octahedra with [2+2+2] coordination, in which the O atoms are coordinated by 1, 2, 3 or 6 V atoms (Fig. 1). It is a common constituent of secondary vanadium minerals that form from oxidation of primary vanadium minerals in ore bodies exposed to near-surface conditions. Decavanadates are also important as catalysts (*e.g.*, Csányi *et al.* 2003, Valverde *et al.* 2012) and as inhibitors of bacterial growth (*e.g.*,

Gillet *et al.* 2016, Samart *et al.* 2018). The decavanadate anion, $[V_{10}O_{28}]^{6-}$, is stable at mildly acidic conditions and relatively high Eh. At lower pH, protonated derivatives occur: $[H_xV_{10}O_{28}]^{(6-x)-}$; at more reducing conditions, mixed-valence decavanadate anions, $[(V^{4+}_xV^{5+}_{10-x})O_{28}]^{(6+x)-}$, are common. In the last 16 years, the atomic arrangements of 18 decavanadate minerals have been determined and refined (Table 1). The structures consist of negatively charged decavanadate clusters, the structural unit, linked by weakly bonded interstitial complexes consisting of monovalent and divalent (simple or

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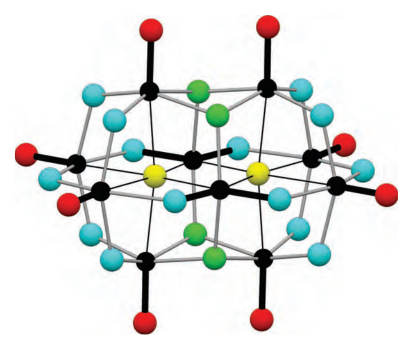


Fig. 1. The [V₁₀O₂₈]ⁿ⁻ decavanadate polyanion. V atoms = black circles, [1]-coordinated O atoms = red circles, [2]-coordinated O_C atoms = blue circles, [3]-coordinated O_B atoms = green circles, [6]-coordinated O atoms = yellow circles, V-O_{vanadyl} bonds = thick black line, V-O_{nans} bonds = thin black line, V-O_{equatorial} bonds = grey shaded line. Known O_D sites: ^[2]C- and ^[3]B-type anions (Day *et al.* 1987).

complex) cations and (H₂O) groups, and are consistent with the idea of binary representation of structure whereby a complicated structure is considered as two parts, a strongly bonded structural unit and a weakly bonded interstitial complex (Hawthorne 1983, Hawthorne & Schindler 2008, Schindler et al. 2000). In order to examine the interaction of these two constituents, we need to derive the Lewis basicity of the decayanadate polyanion, which is a function of the net charge of the polyanion, which in turn depends on (1) the amounts of V^{4+} and V^{5+} , and (2) the presence of any protonated anions in the polyanion. Cooper et al. (2019) addressed the issue of calculating the site occupancies of V^{4+} and V^{5+} in non-protonated decavanadates. The interstitial complexes of decavanadate minerals are commonly disordered and more poorly characterized relative to the more robust decavanadate polyanion. The final charge balance between the decavanadate polyanion and the interstitial complex is sometimes achieved via speculative assignment of protons to exterior anions of the decavanadate group. Here we address the issue of deriving the number of H+ ions in protonated decayanadates via a more quantitative assessment of prospective O_D (i.e., O_{DONOR}) and O_A (O_{ACCEPTOR}) anions. Issues involving OD and OA in synthetic decavanadates have been examined previously: Ferreira da Silva et al. (2003) and Correia et al. (2004) examined the effect of protonation and hydrogen bond donor-acceptor relations on the polymerization of decavanadate polyanions; and Bošnjaković-Pavlović et al. (2011) showed that the strengths of hydrogen bonds are strongly dependent on the connectivity of the anions to V, and suggest that this could be a key to understanding the physiological functions of decavanadates.

GENERAL APPROACH

Here we examine recent structure refinements containing known protonated decavanadates with the intent of developing simple criteria to help identify specific O_D and O_A atom(s) belonging to the decavanadate polyanion. Direct detection of a H atom belonging to an (OH) group from residual electrondensity in a difference-Fourier map at the final stages of structure refinement is often not possible in these structures, particularly in minerals, because of disorder in the structure, and a more general crystal-chemical approach is required.

Using bond-valence sums for decavanadate O atoms, Evans & Popev (1984) showed that long V-O

TABLE 1. RECENTLY CHARACTERIZED DECAVANADATE MINERALS

Mineral species	Ideal formula*	First reference, best structure reference				
	Decavanadates: [V ₁₀ O ₂₈	3]6-				
Ammoniolasalite	$\{(NH_4)_2Mg_2(H_2O)_{20}\}\ [V_{10}O_{28}]$	Kampf et al. 2018a				
Burroite	$\{(NH_4)_2Ca_2(H_2O)_{15}\}[V_{10}O_{28}]$	Kampf et al. 2017a				
Huemulite	${Na_4Mg(H_2O)_{24}}[V_{10}O_{28}]$	Gordillo et al. 1966, Colombo et al. 2011				
Hughesite	${Na_3Al(H_2O)_{22}} [V_{10}O_{28}]$	Rakovan et al. 2011				
Hummerite	$\{[K_2Mg_2(H_2O)_{16}\}\ [V_{10}O_{28}]$	Weeks et al. 1951, Hughes et al. 2002				
Hydropascoite**	${Ca_3(H_2O)_{24}} [V_{10}O_{28}]$	Kampf et al. 2017b				
Kokinosite	${Na_2Ca_2(H_2O)_{24}} [V_{10}O_{28}]$	Kampf <i>et al.</i> 2014a				
Lasalite	${Na_2Mg_2(H_2O)_{20}} [V_{10}O_{28}]$	Hughes et al. 2008				
Magnesiopascoite	${Ca2Mg(H2O)16} [V10O28]$	Kampf & Steele 2008				
Okieite	${Mg_3(H_2O)_{27}}[V_{10}O_{28}]$	Kampf et al. 2018b				
Pascoite	${Ca_3(H_2O)_{17}}[V_{10}O_{28}]$	Hillebrand et al. 1914, Hughes et al. 2005				
Postite	${MgAl_2(OH)_2(H_2O)_{27}} [V_{10}O_{28}]$	Kampf <i>et al.</i> 2012				
Schindlerite	$\{(NH_4)_4Na_2(H_2O)_{10}\}\ [V_{10}O_{28}]$	Kampf et al. 2013a, Kampf et al. 2016				
Wernerbaurite	$\{(NH_4)_2Ca_2(H_2O)_{16}\}\ [V_{10}O_{28}]$	Kampf et al. 2013a, Kampf et al. 2016				
Protonated Decavanadates: [H _x V ₁₀ O ₂₈] ^{(6-x)-}						
Gunterite	$\{Na_4(H_2O)_{22}\} [H_2V_{10}O_{28}]$	Kampf et al. 2011a				
Rakovanite***	{Na ₃ (H ₂ O) ₁₅ } [H ₃ V ₁₀ O ₂₈]	Kampf <i>et al</i> . 2011b				
Mixed-Valence Decavanadates: $[(V^{4+}_xV^{5+}_{10-x})O_{28}]^{(6+x)-}$						
Bluestreakite	$\{K_4Mg_2(H_2O)_{14}\}[V^{4+}_2V^{5+}_8O_{28}]$	Kampf et al. 2014b				
Nashite	${Na_3Ca_2(H_2O)_{24}}[(V^{4+}V^{5+}_9)O_{28}]$	Kampf et al. 2013b				

^{*} Expressed as the interstitial complex and the structural unit.

bonds are associated with O_D atoms. A recent analysis of bond valence in the structures of many decavanadate minerals showed that the bond-valence equation of Brown & Altermatt (1985) for V⁵⁺—O interactions is accurate in validating the formal valence of V only in decavanadates containing only V⁵⁺. Cooper *et al.* (2019) developed a bond-valence equation that predicts site-specific V⁴⁺ content; this equation may also be used to calculate accurate sums of the bond valence incident at the decavanadate O atoms for high-quality structure-refinements, as these have accurate V—O bond-lengths.

PROTONATED DECAVANADATES

Seventeen well-characterized protonated decavanadate structures that contain either inorganic or organic interstitial components were used in this work (Table 2). The structures are well ordered, recent (1987–2015), have low R values (R < 10%), and contain only V^{5+} . For many of these decavanadates, the H atoms belonging to the decavanadate (OH) group were located experimentally. The pro-

tonated decavanadate polyanions contain from one to four H atoms that form ordered (OH) groups on the exterior of the decavanadate groups at specific O sites.

Bond-valence sums at the decavanadate anions

We calculated partial bond-valence sums at the decavanadate anions resulting from the V-O interactions alone (s Σ_{OV}) using the V⁵⁺-O equation of Brown & Altermatt (1985). Non-protonated O atoms on the surface of the decavanadate polyanion (i.e., bonded to one, two, or three V atoms) have $s \Sigma_{OV}$ values from \sim 1.7–2.0 vu (Table 2), and the remaining bond-valence received by these anions to satisfy the valence-sum rule comes from interstitial constituents or from hydrogen bonds from a neighboring decavanadate (OH) group. Protonated O atoms on the exterior of the polyanion have significantly lower s $\Sigma_{\rm OV}$ values, from $\sim 1.2 - 1.5$ vu, and occur for two specific types of O atoms: (1) [3]-coordinated O atoms [denoted here as O_B] and (2) [2]-coordinated O atoms [denoted here as O_C] (nomenclature of Day et al. 1987) (Fig. 1, Table 1).

^{**} Kampf *et al.* (2017b) proposed that, in hydropascoite, a small amount of V is tetravalent and there is some protonation to balance the charge; however, the ideal formula is written with all V⁵⁺ and no protonation.

^{***} The formula for rakovanite is probably [(NH₄)₃Na₃(H₂O)₁₂] [V₁₀O₂₈].

TABLE 2. MISCELLANEOUS INFORMATION FOR 17 PROTONATED DECAVANADATES

[POV]	Interstitial component	$s \Sigma_{OV^*} (\mathit{vu})$	O _D type**	O_DO_A (Å)	Ref.
[HV ₁₀ O ₂₈]	$K_4Na(H_2O)_{10}$	1.49	С	3.03	[1]
$[HV_{10}O_{28}]$	Rb ₄ Na(H ₂ O) ₁₀	1.52	С	3.08	[2]
$[HV_{10}O_{28}]$			С	2.65	[3]
$[H_2V_{10}O_{28}]$	- , , ,		С	2.88	[4]
$[H_2V_{10}O_{28}]$	[(CH3)3CNH3]4	1.23	С	2.77	[5]
$[H_2V_{10}O_{28}]$	$Cs_4(H_2O)_4$	1.28	С	2.61	[6]
$[H_2V_{10}O_{28}]$	$[N(CH_3)_4]_4(CH_3COOH)(H_2O)_{2.8}$	1.26	С	2.75	[7]
		1.31	В	2.75	
$[H_2V_{10}O_{28}]$	$[CH_3(CH_2)_{11}N(CH_3)_3]_4(H_2O)_8$	1.22	С	2.76	[8]
[H ₂ V ₁₀ O ₂₈]	[(CH ₃) ₄ N] ₄ (H ₂ O) _{3.8}	1.29	С	2.76	[9]
	2, 2, 2, 2, 2	1.34	В	2.77	
[H ₃ V ₁₀ O ₂₈]	$(C_6H_8N)_3(H_2O)$	1.29	С	2.74	[10]
	, , , , , , , , , , , , , , , , , , , ,	1.32	В	2.80	
		1.29	С	2.75	
$[H_3V_{10}O_{28}]$	$(C_7H_{10}N)_3(H_2O)$	1.30	В	2.81	[11]
	(1 10)6(2)	1.27	С	2.74	
		1.24	С	2.73	
$[H_3V_{10}O_{28}]$	$[(C_6H_5)_4P]_3(CH_3CN)_4$	1.27	В	2.76	[12]
	2(0 0). 20(0).	1.29	С	2.76	
		1.25	С	2.78	
[H ₃ V ₁₀ O ₂₈]	$[(C_2H_5)_4N]_3(H_2O)_2$	1.24	С	2.76	[9]
	1(2 0). 10(2 /2	1.26	В	2.76	
		1.24	С	2.76	
$[H_3V_{10}O_{28}]$	[(n-C ₃ H ₇) ₄ N] ₃	1.26	С	2.79	[9]
	1(-0 //4 10	1.29	В	2.89	
		1.23	С	2.96	
$[H_3V_{10}O_{28}]$	$[(n-C_4H_9)_4N]_3$	1.26	C	2.76	[9]
	10 -4 3/4 13	1.31	В	2.81	
		1.31	C	2.78	
$[H_4V_{10}O_{28}]$	$[(n-C_3H_7)_4N]_2$	1.38	В	2.70	[9]
1 4-10-201	11 - 3: 1/4: 12	1.37	C	2.72	[-]
$[H_4V_{10}O_{28}]$	$[(n-C_4H_9)_4N]_2$	1.36	В	2.72	[9]
	1(49/412	1.36	C	2.74	[0]

References: [1] Lee & Joo (2004); [2] Yakubovich *et al.* (2015); [3] Jouffret *et al.* (2010); [4] Riou *et al.* (1998); [5] Wery *et al.* (1996); [6] Tatyanina *et al.* (1987); [7] Pecquenard *et al.* (1998); [8] Janauer *et al.* (1997); [9] Nakamura & Ozeki (2001); [10] Santiago *et al.* (1988); [11] Arrieta *et al.* (1988); [12] Day *et al.* (1987).

Hydrogen-bond-acceptor anions for decavanadate (OH) groups

For the seventeen protonated decavanadate structures examined, there are 33 individual (OH) groups associated with either $^{[3]}O_{B}$ or $^{[2]}O_{C}$ anions of the decavanadate polyanion (Table 2). All hydrogen bonds are directed toward a neighboring decavanadate polyanion in which the O_{A} anions are 2.61–3.08 Å from their respective O_{D} anions on the adjacent polyanion. Figure 2 shows the variation in $O_{D}\ldots O_{A}$ distance as a function of s Σ_{OV} values for the respective O_{D} anions. The line through the data in the lower part of Figure 2 has a negative slope that is consistent with longer $O_{D}\ldots O_{A}$ distances (and weaker

hydrogen bonds) coupled to lower incident bond-valence received by the O_D from the bonded V atoms. This relation is in good agreement with the valence-sum rule (Brown 2016). Strong V–O bonds provide $\sim 1.4~vu$ to O_D and the H atom provides $\sim 0.6~vu$ to O_D to satisfy the valence-sum rule at O_D . In turn, the valence-sum rule at the H atom requires a relatively strong hydrogen bond ($\sim 0.4~vu$) to a close O_A anion (i.e., $O_D \dots O_A \approx 2.6~\text{Å}$) of a neighboring decavanadate polyanion. Weak V–O bonds provide $\sim 1.2~vu$ to O_D and the H atom provides $\sim 0.8~vu$ to O_D to satisfy the valence-sum rule at O_D . The valence-sum rule at the H atom requires a weaker hydrogen bond ($\sim 0.2~vu$) to an O_A anion (i.e., $O_D \dots O_A \approx 2.9~\text{Å}$) of a neighboring

^{*} Bond valence from V-O_D contributions using Brown & Altermatt (1985); ** Day et al. (1987).

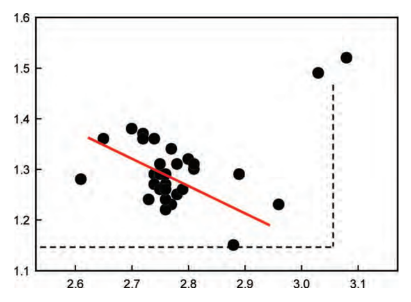


Fig. 2. O_D...O_A distances for the 33 (OH) groups as a function of the bond valence incident at O_D from the V–O bonds alone. Solid red line: hand drawn through the ordered (OH) data; dashed lines: disordered (OH) groups extrapolated to main trend.

decavanadate polyanion. Thus, accurate assessment of the incident bond-valence to an O_D atom of a potential (OH) group on a decavanadate polyanion, coupled with its distance to a prospective O_A on a neighboring decavanadate polyanion, can provide powerful confirmation of protonation without the requirement that the H atom itself be located during structure refinement.

Two data points fall off the trend in Figure 2; these belong to two isostructural decavanadates, $K_4Na(H_2O)_{10}$ [HV₁₀O₂₈] and Rb₄Na(H₂O)₁₀[HV₁₀O₂₈] (Lee & Joo 2004, Yakubovich *et al.* 2015). For this structure type, the hydrogen atom is disordered between symmetrically equivalent O_C anions on adjacent decavanadate groups (Fig. 3); whether this disorder is static or dynamic is not known. The net effect is that two local

arrangements (Fig. 3a, b) occur with equal frequency. In each local arrangement, the observed $O_D ... O_A$ distance of ~ 3.05 Å indicates that O_D will receive ~ 1.15 vu from its two bonds with V (dashed lines on Fig. 2). The remaining bond-valence (*i.e.*, 2-1.15=0.85 vu) required to satisfy the valence-sum rule will come from the H atom, and the ensuing hydrogen-bond-valence is 0.15 vu. If we then average the two different bond-valences incident from the V atoms for the two arrangements [*i.e.*, (1.15+1.85)/2=1.5 vu], the predicted long-range value of ~ 1.5 vu is in close agreement with the value calculated from the experimental (long-range-averaged) bond lengths (Table 2, Fig. 2).

Fig. 3. The two local hydrogen-bond arrangements associated with the disordered (OH) group in $K_4Na(H_2O)_{10}[HV_{10}O_{28}]$ and $Rb_4Na(H_2O)_{10}[HV_{10}O_{28}]$, resulting in an average bond-valence contribution of 1.50 vu to $O_{A/D}$ from the V atoms.

Protonated decavanadate minerals

The structures reported for rakovanite and gunterite contain protonated decayanadate polyanions with the formulae $Na_3\{H_3[V_{10}O_{28}]\}(H_2O)_{15}$ and Na_4 (H₂O)₁₆(H₂V₁₀O₂₈)(H₂O)₆, respectively (Kampf et al. 2011a, b). For rakovanite, the orange color and bond-valence sums at the V atoms (5.00-5.11 vu) indicate that all V is V5+. The decayanadate O atoms have $s \Sigma_{OV}$ sums from 1.66 to 1.98 vu and 1.73 to 1.92 vu for O_B and O_C anions specifically. The lowest s Σ_{OV} value is for a vanadyl O atom that also receives additional bond-valence from interstitial Na. Thus, all decavanadate OB and OC atoms belonging to the rakovanite polyanion contain $s \Sigma_{OV}$ sums within the range $\sim 1.7-2.0$ vu and are expected to be unprotonated by analogy with known protonated decayanadate structures (Table 2, Fig. 2). In addition, the closest anion approach between polyanions involving at least one O_B or O_C anion is 3.41 Å, which lies well outside the range of data in Figure 2. However, we note that even though all the known protonated decavanadates investigated thus far have hydrogen bonds from (OH) groups to OA anions on a neighboring polyanion (and thus closer OD...OA approaches; Fig. 2), there is no crystal-chemical rationale that excludes an interstitial constituent [e.g., the O atom of an (H₂O) group] as a possible hydrogen-bond acceptor. The strong indication of an unprotonated decavanadate polyanion in rakovanite has led us to initiate a re-evaluation of rakovanite both chemically and structurally in terms of the interstitial components. An FTIR examination of rakovanite indicates that it contains significant NH₄⁺ (Paul M. Adams, pers. commun.). As a result, a reinvestigation of rakovanite has been initiated, which is likely to lead to a revision of its formula. For gunterite, poor crystal quality and uncertainty in the refined structure does not allow conclusive evaluation of possible decavanadate protonation using the above crystal-chemical criteria; a better-quality refinement is needed, but FTIR examination of gunterite indicates that it does not contain significant NH₄⁺ (Paul M. Adams, pers. commun.).

Mixed $V^{4+/5+}$ protonated decayanadates

The occurrence of V^{4+} in a decavanadate increases the net charge of the polyanion, whereas protonation decreases the net charge of the polyanion. If attainment of electroneutrality were the only constraint on polyanion charge, one would not expect mixed V^{4+}/V^{5+} protonated decavanadates to occur. On the other hand, one expects the V^{4+}/V^{5+} ratio to be sensitive to oxygen fugacity, and hence low fO_2 and a need for a high polyanion charge to attain electroneutrality might lead to mixed V^{4+}/V^{5+} protonated decavanadates.

Protonation of a decavanadate polyanion requires an abnormally low value of $s \Sigma_{OV}$ at a potential O_D anion. This can occur *via* two mechanisms: (1) V^{5+} —O bond elongation in typical decavanadates containing only V^{5+} , or (2) incorporation of a lower-valence cation (*i.e.*, V^{4+}).

In this regard, the well-refined structure of $(H_3O)_2[M(H_2O)_6]_2[H_6V_{10}O_{28}]\cdot 4H_2O$ (M = Co,Ni)piqued our interest as the polyanion was inferred to have the composition $[H_6V^{4+}_{6}V^{5+}_{4}O_{28}]^{6-}$ (Khan *et al.* 2000). The Co and Ni variants are isostructural and we evaluated the slightly better refined (R = 5.0%) Ni structure. Application of the V⁵⁺-O bond-valence parameters to [6]V-O distances would result in an incident bond-valence sum at the V atom significantly below 5 if significant V^{4+} is ordered at the V site; a V site occupied completely by V⁴⁺ is expected to have an incident bond-valence sum of $\sim 4.35 \ vu$ (Cooper et al. 2019). Bond-valence calculations for the M = Nidecavanadate of Khan et al. (2000) using the V5+-O relation of Brown & Altermatt (1985) gives bondvalence sums from 4.99 to 5.07 vu for the V sites and from 1.70 to 1.97 vu for the O sites of the polyanion. We therefore conclude that the polyanion contains only V⁵⁺ and is unprotonated.

Hydropascoite, ideally Ca₃(V₁₀O₂₈)(H₂O)₂₄, is yellow-green and contains minor V4+ at the V4 and V5 sites (Kampf et al. 2017b). Bond-valence calculations using the V⁵⁺-O relation of Brown & Altermatt (1985) give bond-valence sums of 4.85 and 4.73 vu, respectively, for the V4 and V5 sites in hydropascoite. Cooper et al. (2019) derived an equation to predict the actual bond-valence, V_P , incident at a given V site: V_P = $1.538(V_{\rm C}) - 2.692 \ vu$, where $V_{\rm C}$ is the incident bondvalence sum calculated using the V5+O equation of Brown & Altermatt (1985). Application of this equation gives a composition of $(V^{5+}_{0.77}V^{4+}_{0.23})$ for the V4 site and $(V^{5+}_{0.58}V^{4+}_{0.42})$ for the V5 site. The resultant polyanion composition is $(V^{5+}_{9.35}V^{4+}_{0.65}O_{28})^{-6.65}$, and charge balance was inferred to occur via protonation of the decayanadate polyanion (Kampf et al. 2017b). The bond-valence sums for the O sites of the polyanion from V span 1.60–1.96 vu, with the incident bond-valences at the O_B and O_C anions spanning 1.72–1.87 vu. The atom displacement parameters for all V and O atoms of the two symmetrically distinct polyanions are reasonably uniform and do not suggest any significant local V-O bond elongation that might be coupled to a V-O bond arrangement (possibly involving V⁴⁺) that would result in a significant reduction in bond valence at the anion. Thus, all decavanadate O_B and O_C atoms belonging to the hydropascoite polyanion contain s Σ_{OV} sums in the range \sim 1.7–2.0 vu and are expected to be unprotonated by analogy with known protonated decavanadate structures (Table 1, Fig. 2). Additionally, the hydropascoite structure does not contain close anion–anion approaches between neighboring polyanions. Charge balance involving $V^{4+/5+}$ may possibly involve an unknown positively charged interstitial constituent in the vicinity of the OW24 position: the atom-displacement parameters associated with the OW24 site are extreme, and there is a minor peak in the difference-Fourier map $\sim 1.5 \text{ Å}$ from OW24. In this context, it should be noted that FTIR examination of hydropascoite indicates that it does not contain significant NH₄⁺ (Paul M. Adams, *pers. commun.*).

Conclusions

The following criteria are key indicators of decavanadate protonation:

- (1) Incident bond-valence sums ($s \Sigma_{OV}$) at the O_C or O_B decavanadate exterior anions that are in the range 1.2–1.5 vu, indicative of O_D anions;
- O<sub>D...O_A distances between neighboring decavanadate polyanions that are <3.1 Å;
 </sub>
- (3) An obvious lack of proximal interstitial constituents capable of providing substantial bond valence to the low-valence O_B or O_C anions in (1) above.

Moreover, we have found no evidence so far for mixed $V^{4+/5+}$ protonated decayanadates.

ACKNOWLEDGMENTS

We thank reviewers Fernando Cámara and Michael Schindler for their useful comments. Paul M. Adams is thanked for providing the results of his FTIR studies on rakovanite, gunterite, and hydropascoite. Financial support was provided by a Discovery Grant from the Natural Sciences and Engineering Research Council of Canada to FCH and by the John Jago Trelawney Endowment to the Mineral Sciences Department of the Natural History Museum of Los Angeles County to ARK.

REFERENCES

- ARRIETA, J.M., ARNAIZ, A., LORENTE, L., SANTIAGO, C., & GERMAIN, G. (1988) Tris(3-ethylpyridinium) decavanadate monohydrate. Acta Crystallographica C44, 1004–1008.
- Bošnjaković-Pavlović, N., Prévost, J., & Spasojević-De Biré, A. (2011) Crystallographic statistical study of decavanadate anion based-structures: Toward a prediction of noncovalent interactions. Crystal Growth and Design 11, 3778–3789.
- Brown, I.D. (2016) The Chemical Bond in Inorganic Chemistry: The Bond Valence Model, 2nd Edition. IUCr, Oxford University Press, Oxford, England, 352 pp.
- Brown, I.D. & Altermatt, D. (1985) Bond-valence parameters obtained from a systematic analysis of the inorganic

- crystal structure database. *Acta Crystallographica* **B41**, 244–247.
- COLOMBO, F., BAGGIO, R., & KAMPF, A.R. (2011) The crystal structure of the elusive huemulite. *Canadian Mineralogist* **49**, 849–864.
- Cooper, M.A., Hawthorne, F.C., Kampf, A.R., & Hughes, J.M. (2019) Determination of $V^{4+}:V^{5+}$ ratios in the $\left[V_{10}O_{28}\right]^{n-}$ decavanadate polyanion. *Canadian Mineralogist* (this issue).
- CORREIA, I., AVECILLA, F., MARCÃO, S., & PESSOA, J.C. (2004) Structural studies of decavanadate compounds with organic molecules and inorganic ions in their crystal packing. *Inorganica Chimica Acta* 357, 4476–4487.
- CSÁNYI, L.J., JÁKY, K., DOMBI, G., EVANICS, F., DEZSŐ, G., & KÓTA, Z. (2003) Onium-decavanadate ion-pair complexes as catalysts in the oxidation of hydrocarbons by O₂.

 Journal of Molecular Catalysis A: Chemical 195, 101–111.
- Day, V.W., Klemperer, W.G., & Maltbie, D.J. (1987) Where are the protons in $H_3V_{10}O_{28}^{3-}$? *Journal of the American Chemical Society* **109**, 2991–3002.
- EVANS, H.T., JR. & POPEV, M.T. (1984) Reinterpretation of five recent crystal structures of heteropoly and isopoly complexes: divanadodecamolybdophosphate, trivanadoenneamolybdophosphate, ".gamma.-dodecatungstophosphate", the dodecamolybdate-dodecamolybdomolybdate blue complex, and dihydrogen decavanadate. *Inorganic Chemistry* 23, 501–504.
- FERREIRA DA SILVA, J.L., MINAS DA PIEDADE, M.S., & DUARTE, M.T. (2003) Decavanadates: a building-block for supramolecular assemblies. *Inorganica Chimica Acta* 356, 222–242.
- GILLET, J-.M., Wu, P., WEI, Y., & SPASOJEVIĆ-DE BIRÉ, A. (2016) A combined crystallographic analysis and ab initio calculations to interpret the reactivity of functionalized hexavanadates and their inhibitor potency toward Na⁺/K⁺-ATPase. Journal of Inorganic Biochemistry 161, 27–36.
- Gordillo, C.E., Linares, E., Toubes, R.O., & Winchell, H. (1966) Huemulite, $Na_4MgV_{10}O_{28}\cdot 24H_2O$, a new hydrous sodium and magnesium vanadate from Huemul mine, Mendoza Province, Argentina. *American Mineralogist* **51**, 1–13.
- HAWTHORNE, F.C. (1983) Graphical enumeration of polyhedral clusters. Acta Crystallographica A39, 724–736.
- Hawthorne, F.C. & Schindler, M. (2008) Understanding the weakly bonded constituents in oxysalt minerals. *Zeits-chrift für Kristallographie* **223**, 41–68.
- HILLEBRAND, W.F., MERWIN, H.E., & WRIGHT, F.E. (1914) Hewettite, Metahewettite and Pascoite, Hydrous Calcium Vanadates. *Proceedings of the American Philosophical* Society 53(213), 31–54.

- Hughes, J.M., Schindler, M., Rakovan, J., & Cureton, F.E. (2002) The crystal structure of hummerite, $KMg(V_5O_{14})^{\bullet}8H_2O$: Bonding between the $[V_{10}O_{28}]^{6-}$ structural unit and the $\{K_2Mg_2(H_2O)_{16}\}6+$ interstitial complex. Canadian Mineralogist 40, 1429–1435.
- HUGHES, J.M., SCHINDLER, M., & FRANCIS, C.A. (2005) The C2/ m disordered structure of pascoite, Ca₃(V₁₀O₂₈)•17H₂O. Canadian Mineralogist 43, 1379–1386.
- Hughes, J.M., Wise, W.S., Gunter, M.E., Morton, J.P., & Rakovan, J. (2008) Lasalite, Na₂Mg₂[V₁₀O₂₈]•20H₂O, a new decavanadate mineral species from the Vanadium Queen Mine, La Sal District, Utah: Description, atomic arrangement, and relationship to the pascoite group of minerals. *Canadian Mineralogist* **46**, 1365–1372.
- JANAUER, G.G., DOBLEY, A.D., ZAVALII, P.Y., & WHITTINGHAM, M.S. (1997) Evidence for decavanadate clusters in the lamellar surfactant ion phase. *Chemistry of Materials* 9, 647–649.
- JOUFFRET, L., RIVENET, M., & ABRAHAM, F. (2010) A new polymeric chain in the dihydrogendecavanadate(V)-decavanadate(V) [NH₂(CH₂)₄]₅[V₁₀O₂₈H₂]_{0.5}[V₁₀O₂₈]_{0.5} obtained by *in situ* synthesis of the organic cation. *Inorganic Chemistry Communications* 13, 5–9.
- KAMPF, A.R. & STEELE, I.M. (2008) Magnesiopascoite, a new member of the pascoite group: Description and crystal structure. *Canadian Mineralogist* 46, 679–686.
- KAMPF, A.R., HUGHES, J.M., MARTY, J., & NASH, B. (2011a) Gunterite, Na₄(H₂O)₁₆(H₂V₁₀O₂₈)•6H₂O, a new mineral with a doubly-protonated decavanadate polyanion: Crystal structure and descriptive mineralogy. *Canadian Mineralogist* 49, 1243–1251.
- KAMPF, A.R., HUGHES, J.M., MARTY, J., GUNTER, M.E., & NASH, B. (2011b) Rakovanite, Na₃{H₃[V₁₀O₂₈]}•15H₂O, a new species of the pascoite family with a protonated decvanadate polyanion. *Canadian Mineralogist* 49, 889– 898.
- Kampf, A.R., Hughes, J.M., Marty, J., & Nash, B. (2012) Postite, Mg(H₂O)₆Al₂(OH)₂(H₂O)₈(V₁₀O₂₈)•13H₂O, a new mineral species from the La Sal mining district, Utah: Crystal structure and descriptive mineralogy. *Canadian Mineralogist* **50**, 45–53.
- Kampf, A.R., Hughes, J.M., Marty, J., & Nash, B.P. (2013a) Wernerbaurite, $\{[Ca(H_2O)_7]_2(H_2O)_2(H_3O)_2\}\{V_{10}O_{28}\}$, and schindlerite, $\{[Na_2(H_2O)_{10}](H_3O)_4\}\{V_{10}O_{28}\}$, the first hydronium-bearing decavanadate minerals. *Canadian Mineralogist* **51**, 297–312.
- KAMPF, A.R., HUGHES, J.M., MARTY, J., & BROWN, F.H. (2013b) Nashite, Na₃Ca₂([V⁵⁺₉V⁴⁺₁]O₂₈)•24H₂O, a new mineral species from the Yellow Cat Mining District, Utah and the Slick Rock Mining District, Colorado: Crystal structure and descriptive mineralogy. *Canadian Mineralogist* 51, 27–37.
- KAMPF, A.R., HUGHES, J.M., NASH, B.P., & MARTY, J. (2014a) Kokinosite, Na₂Ca₂(V₁₀O₂₈)·24H₂O, a new decavanadate

- mineral species from the St. Jude mine, Colorado: Crystal structure and descriptive mineralogy. *Canadian Mineralogist* **52**, 15–25.
- KAMPF, A.R., HUGHES, J.M., MARTY, J., NASH, B.P., CHEN, Y.-S., & STEELE, I.M. (2014b) Bluestreakite, K₄Mg₂ (V⁴⁺₂V⁵⁺₈O₂₈)·14H₂O, a new mixed-valence decavanadate mineral from the Blue Streak Mine, Montrose County, Colorado: Crystal structure and descriptive mineralogy. Canadian Mineralogist 52, 1007–1018.
- Kampf, A.R., Hughes, J.M., Nash, B.P., Marty, J., Cooper, M.A., Hawthorne, F.C., Karpenko, V.Y., Pautov, L.A., & Agakhanov, A.A. (2016) Revision of the formulas of wernerbaurite and schindlerite: Ammonium-rather than hydronium-bearing decavanadate minerals. *Canadian Mineralogist* 53, 555–558.
- KAMPF, A.R., NASH, B.P., MARTY, J., & HUGHES, J.M. (2017a) Burroite, Ca₂(NH₄)₂(V₁₀O₂₈)·15H₂O, a new decavanadate mineral from the Burro mine, San Miguel County, Colorado. *Canadian Mineralogist* 55, 473–481.
- KAMPF, A.R., NASH, B.P., MARTY, J., HUGHES, J.M., & ROSE, T.P. (2017b) Hydropascoite, Ca₃(V₁₀O₂₈)·24H₂O, a new decavanadate mineral from the Packrat mine, Mesa County, Colorado. *Canadian Mineralogist* 55, 207–217.
- Kampf, A.R., Nash, B.P., Adams, P.M., Marty, J., & Hughes, J.M. (2018a) Ammoniolasalite, $(NH_4)_2Mg_2(H_2O)_{20}$ [$V_{10}O_{28}$], a new decavanadate species from the Burro mine, Slick Rock district, Colorado. *Canadian Mineralogist* **56**, 859–869.
- KAMPF, A.R., NASH, B.P., ADAMS, P.M., MARTY, J., & HUGHES, J.M. (2018b) Okieite, IMA 2018-080. CNMNC Newsletter No. 46, December 2018. Mineralogical Magazine 82.
- Khan, I., Tabussum, M., & Zheng, C. (2000) Synthesis and characterization of decavanadates: X-ray crystal structures of $(H_3O)_2[M(H_2O)_6]_2[V_{10}O_{28}H_6]\cdot 4H_2O$ (M= Co, Ni) and Na[H₃N(CH₂)₂NH₃]_{2.5}[V₁₀O₂₈]·5H₂O. Synthesis and Reactivity in Organic and Metal-Organic Chemistry 30, 1773–1790.
- LEE, U. & Joo, H.-C. (2004) A novel monoprotonated decavanadate, K₄Na[HV₁₀O₂₈]·10H₂O. Acta Crystallographica E60, i22–i24.
- Nakamura, S. & Ozeki, T. (2001) Hydrogen-bonded aggregates of protonated decavanadate anions in their tetraalkylammonium salts. *Journal of the Chemical Society, Dalton Transactions* **2001**, 472–480.
- Pecquenard, B., Zavalii, P.Y., & Whiingham, M.S. (1998) Tetrakis(tetramethylammonium) dihydrogendecavanadate acetic acid 2.8-hydrate, [N(CH₃)₄]₄[H₂V₁₀O₂₈].CH₃ COOH.2.8H₂O. *Acta Crystallographica* **C54**, 1833–1835.
- RAKOVAN, J., SCHMIDT, G.R., GUNTER, M., NASH, B., KAMPF, A.R., MARTY, J., & WISE, W.S. (2011) Hughesite, Na₃Al(V₁₀O₂₈)•22H₂O, a new member of the pascoite family of minerals from the Sunday mine, San Miguel County, Colorado. *Canadian Mineralogist* 49, 1253–1265.

- RIOU, D., ROUBEAU, O., & FÉREY, G. (1998) Evidence for the solid state structural transformation of the network-type decavanadate (NC₇H₁₄)₄[H₂V₁₀O₂₈] into a lamellar topology (NC₇H₁₄)[V₄O₁₀]. Zeitschrift für Anorganische und Allgemeine Chemie 624, 1021–1025.
- SAMART, N., ARHOUMA, Z., KUMAR, S., MURAKAMI, H.A., CRICK, D.C., & CRANS, D.C. (2018) Decavanadate inhibits mycobacterial growth more potently than other oxovanadates. Frontiers in Chemistry, DOI: 10.3389/fchem.2018. 00519.
- SANTIAGO, C., ARNAIZ, A., LORENTE, L., & ARRIETA, J.M. (1988) Tris(3-methylpyridinium) decavanadate monohydrate. Acta Crystallographica C44, 239–242.
- SCHINDLER, M.C., HAWTHORNE, F.C., & BAUR, W.H. (2000) A crystal-chemical approach to the composition and occurrence of vanadium minerals. *Canadian Mineralogist* 38, 1443–1456.
- Tatyanina, I.V., Sergienko, V.S., Zabolotskikh, A.V., & Torchenkova, E.A. (1987) Crystal Structure of cesium decavanadate tetrahydrate, $Cs_4(H_2V_{10}O_{28})_4(H_2O)$. Koordinatsionnaya Khimiya 13, 680–688.

- Valverde, J.A., Echavarría, A., Ribeiro, M.F., Palacio, L.A., & Eon, J.-G. (2012) Decavanadate-intercalated Ni—Al hydrotalcites as precursors of mixed oxides for the oxidative dehydrogenation of propane. *Catalysis Today* 192, 136–143.
- Weeks, A.D., Cisney, E.A., & Sherwood, A.M. (1951) Hummerite and montroseite, two vanadium minerals from Montrose County, Colorado. *American Mineralogist* 36, 326–327
- WERY, A.S.J., GUTIERREZ-ZORRILA, J.M., LUQUE, A., & ROMAN, P. (1996) Influence of protonation on crystal packing and thermal behavior of tertbutylammonium decavanadates. *Polyhedron* 15, 4555–4564.
- YAKUBOVICH, O.V., STEELE, I.M., YAKOVLEVA, E.V., & DIMITROVA, O.V. (2015) One-dimensional decavanadate chains in the crystal structure of Rb₄[Na(H₂O)₆] [HV₁₀O₂₈]·4H₂O. *Acta Crystallographica* C71, 465–473.
- Received November 28, 2018. Revised manuscript accepted January 30, 2019.