#### **REVISION 1** 1 Bicapite, KNa<sub>2</sub>Mg<sub>2</sub>(H<sub>2</sub>PV<sup>5+</sup><sub>14</sub>O<sub>42</sub>)·25H<sub>2</sub>O, a new polyoxometalate mineral with 2 a bicapped Keggin anion from the Pickett Corral mine, Montrose County, 3 Colorado, U.S.A. 4 5 ANTHONY R. KAMPF<sup>15</sup>, JOHN M. HUGHES<sup>2</sup>, BARBARA P. NASH<sup>3</sup>, AND JOE MARTY<sup>4</sup> 6 7 8 <sup>1</sup>Mineral Sciences Department, Natural History Museum of Los Angeles County, 900 9 10 Exposition Boulevard, Los Angeles, California 90007, U.S.A. <sup>2</sup>Department of Geology, University of Vermont, Burlington, VT 05405, U.S.A. 11 12 <sup>3</sup>Department of Geology and Geophysics, University of Utah, Salt Lake City, Utah 84112, U.S.A. 13 <sup>4</sup>5199 East Silver Oak Road, Salt Lake City, UT 84108, U.S.A. 14 15 ABSTRACT 16 17 Bicapite, $KNa_2Mg_2(H_2PV^{5+}_{14}O_{42})$ · 25H<sub>2</sub>O, is a new mineral species (IMA2018-048) 18 19 discovered at the Pickett Corral mine, Montrose County, Colorado. Bicapite occurs as square tablets up to about 0.2 mm on edge on montroseite-corvusite-bearing sandstone. Crystals are 20 dark red-brown, often appearing black. The streak is orange and the luster is vitreous. Bicapite is 21 22 brittle, has a Mohs hardness of $1\frac{1}{2}$ , and displays one excellent cleavage on $\{100\}$ . The measured density is 2.44(2) g·cm<sup>-3</sup>. Bicapite is uniaxial (+), $\omega = 1.785(5)$ , $\varepsilon \approx 1.81$ (white light); 23 pleochroism is red-brown; E > O, slight. The electron probe microanalysis and results of the 24 crystal structure determination provided the empirical formula (based on 67 O apfu) 25 $(K_{1,23}Na_{2,23}Mg_{1,48})_{\Sigma 4,94}[H_{2,51}P_{1,02}(V^{5+}_{13,91}Mo^{6+}_{0,07})_{\Sigma 13,98}O_{42}] \cdot 25H_2O$ . Bicapite is tetragonal, I4/m, 26 with a = 11.5446(12), c = 20.5460(14) Å, V = 2738.3(6) Å<sup>3</sup>, and Z = 2. The strongest four lines 27 in the diffraction pattern are [d in Å(I)(hkl)]: 10.14(100)(002, 101), 2.978(29)(134, 206), 28 2.809(11)(305), and 2.583(11)(420, 008). The atomic arrangement of bicapite was solved and 29

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30	refined to $R_1 = 0.0465$ for 1008 independent reflections with $I > 2\sigma I$ . The structural unit is a
31	$[H_2PV_{12}^{5+}O_{40}(V_{12}^{5+}O_{2})_2]^{7-}$ heteropolyanion composed of twelve distorted VO <sub>6</sub> octahedra
32	surrounding a central PO <sub>4</sub> tetrahedron and capped on opposite sides by two VO <sub>5</sub> square
33	pyramids; the structural unit is a modification of the $\alpha$ -isomer of the Keggin anion, $[XM_{12}O_{40}]^{n-}$ .
34	Charge balance in the structure is maintained by the $[KNa_2Mg_2(H_2O)_{25}]^{7+}$ interstitial complex.
35	The name bicapite is in recognition of this being the only known mineral with a structure based
36	on a bicapped Keggin anion. The discovery of bicapite and numerous other natural
37	polyoxometalate compounds in the Colorado Plateau uranium/vanadium deposits make that the
38	most productive region found to date for naturally occurring polyoxometalate compounds.
39	
40	Keywords: Bicapite; new mineral; crystal structure; polyoxometalate; bicapped Keggin
41	anion; Pickett Corral mine, Montrose County, Colorado, USA
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#### **INTRODUCTION AND OCCURRENCE**

The arcuate 120 km-long Uravan Mineral Belt of the Colorado Plateau has been a rich 44 source of uranium and vanadium ore since early in the twentieth century. The numerous mines in 45 the belt exploit roll-front deposits in sandstone of the Salt Wash member of the Morrison 46 47 Formation (Carter and Guiltierri 1965; Shawe 2011). The mineralization of U and V took place where solutions rich in  $U^{6+}$  and  $V^{4+}/V^{5+}$ -bearing aqueous species encountered local strongly 48 reducing conditions, typically due to the presence of organic matter. The details of the 49 geochemistry of the deposits and conditions of deposition were first examined in detail by Evans 50 51 and Garrels (1958). The uranium and vanadium deposits of the Uravan Mineral Belt have also been a rich source of new minerals; more than 30 minerals with essential vanadium have been 52 discovered there since 2008 during collecting undertaken by one of the authors (JM) and his 53 colleagues. These minerals have added extensively to the knowledge of the complexity of 54 naturally occurring vanadium compounds. 55

The new mineral bicapite was found underground in the Pickett Corral mine, Bull 56 Canyon, Montrose County, Colorado, about 13 km west of the town of Naturita. The portal to 57 the Pickett Corral mine is located at 38.195272, -108.843326; however, the partially flooded 58 59 workings of the mine were accessed through the main portal of the interconnected Blue Streak mine located at 38.199434, -108.839946. The Pickett Corral mine is in the central portion of the 60 Uravan Mineral Belt. The new mineral forms from the oxidation of montroseite-corvusite 61 62 assemblages in a moist environment. Under ambient temperatures and generally oxidizing nearsurface conditions, water reacts with pyrite to form aqueous solutions with relatively low pH; 63 this water reacts with unoxidized and oxidized phases that have been exposed by mining 64 65 operations to form bicapite and other new mineral species.

66	The name bicapite is in recognition of this being the only known mineral with a structure
67	based on a bicapped Keggin anion. The mineral was collected on March 24, 2016, by one of the
68	authors (JM), Al Wilkins and Okie Howell. It has been found very sparingly on only several
69	small specimens. It occurs on montroseite-corvusite-bearing sandstone in close association with
70	gypsum, huemulite, and thenardite. Bicapite was approved by the Commission on New Minerals,
71	Nomenclature, and Classification of the International Mineralogical Association (IMA 2018-
72	048). Two cotype specimens of bicapite are deposited in the collections of the Natural History
73	Museum of Los Angeles County, Los Angeles, California, USA, catalogue numbers 66915 and
74	66916.
75	POLYOXOMETATES
76	Polyoxometalate compounds (often abbreviated as POMs) are relatively rare in geology,
77	but of great importance in synthetic chemistry and materials science. Polyoxometalates, usually
78	anionic in nature, are clusters of three or more transition metal-centered polyhedra that are linked
79	by sharing oxygen ligands between and among the polyhedra. The linked polyhedra form a
80	three-dimensional framework. One example of a polyoxometalate compound recently found to
81	be common to numerous minerals in the Colorado Plateau deposits is the $[V_{10}O_{28}]^{6-1}$
82	isopolyanion, the decavanadate group, formed of 10 VO <sub>6</sub> octahedra linked through corner- and
83	edge-sharing. Typically, bonding within the polyoxometalate is stronger than between the
84	polyoxometalate and the surrounding crystal field.
85	Polyoxometalates are important phases in industrial chemistry, where a large effort is
86	expended in synthesizing polyoxometalates with desirable characteristics of size, charge, shape
87	and conformation. Many hundreds of papers are published annually in the chemistry literature on
88	polyoxometalates, and the number of papers has been increasing logarithmically since the early

89	1990s (Hutin et al. 2013); a similar increase in patents involving polyoxometalates is also noted
90	(Katsoulis 1998). In a detailed survey of industrial uses of polyoxometalates, Katsoulis (1998)
91	notes that 80-85% of the patents issued for polyoxometalates are in the area of catalysis, with the
92	remaining 15-20% involving the compounds in nearly a score of other applications, including
93	coatings, sorbents of gases, sensors, dyes, capacitors, cation exchangers, anti-tumor agents, and
94	bleaching of pulp paper. New uses of polyoxometalate compounds have been in medicine as
95	antiviral agents, cancer antagonists and treatment for Alzheimer's disease, and in
96	nanomagnetism with applications to semiconductors and quantum computing.
97	There are many methods of synthesis of polyoxometalates, but the most common method
98	is undertaken using acidic aqueous systems that contain the relevant polyoxometalate cations and
99	the cations needed to balance the charge of the anionic polyoxometalate at a controlled pH;
100	molybdate and tungstate polyoxometalates tend to occur at lower pH values than vanadate
101	POMs. The conditions of vanadium polyoxometalate formation in laboratory aqueous synthesis
102	are similar to the natural post-mining conditions found in the Colorado plateau vanadium-
103	uranium mines, where solutions rich in vanadium for the POM formation as well as charge-
104	balancing cations exist at a suitable pH.
105	Although polyoxometalate compounds are not common in minerals, they do exist, and a
106	growing number of POMS are being discovered in the Colorado Plateau Uravan Belt; these
107	phases are listed subsequently. In addition to those minerals with a similar Colorado Plateau
108	genesis to bicapite, POM-bearing minerals include zunyite (Baur and Ohta 1982) and murataite
109	(Ercit and Hawthorne 1995); however, in these minerals, the kegging-type polyoxometalates are

- 110 condensed into frameworks and do not contain independent polyoxometalate units. The Colorado
- 111 Plateau deposits have enlarged the number of POM-bearing minerals greatly, and that region is

112	the most productive region found to date for naturally occurring polyoxometalate compounds.
113	APPEARANCE, OPTICAL AND PHYSICAL PROPERTIES
114	Bicapite occurs as square tablets up to about 0.2 mm on edge (Figures 1 and 2); the
115	crystals are dark red-brown, often appearing black. The streak is orange and the luster is vitreous.
116	Bicapite is non-fluorescent under long- and short-wave ultraviolet light. It is brittle, has a Mohs
117	hardness of $1\frac{1}{2}$ (based upon scratch tests), and exhibits excellent cleavage on {100}; the fracture
118	is irregular, stepped. The density measured by flotation in methylene iodide – toluene is 2.44(2)
119	$g \cdot cm^{-3}$ ; the calculated density is 2.434 $g \cdot cm^{-3}$ for the empirical formula and 2.428 $g \cdot cm^{-3}$ for the
120	ideal formula. Bicapite is very slowly soluble in room-temperature $H_2O$ (~1 hour); the phase
121	decomposes in dilute HCl at room temperature and then slowly dissolves.
122	Optical observations on bicapite were made in white light. Bicapite is uniaxial (+), with $\omega$
123	= 1.785(5) and $\varepsilon \approx 1.81$ . Note that $\varepsilon$ is observed to be slightly greater than 1.80; in liquids > 1.80,
124	crystals of an unknown phase immediately grow on the surfaces of the bicapite crystals.
125	Pleochroism is red-brown; $E > O$ , slight. The Gladstone-Dale compatibility (Mandarino 2007),
126	$(1 - (K_p/K_c))$ , is -0.017), is (superior) using the empirical formula, and -0.019 (superior) using the
127	ideal formula.
128	
129	CHEMICAL COMPOSITION
130	Analyses of bicapite (4 points on 3 crystals) were performed at the University of Utah on
131	a Cameca SX-50 electron microprobe with four wavelength dispersive spectrometers, and
132	utilizing Probe for EPMA software. Analytical conditions were 15 keV accelerating voltage, 10
133	nA beam current and a beam diameter of 5 to 10 $\mu$ m. Counting times were 20 seconds on peak
134	and 20 seconds on background for each element. Raw X-ray intensities were corrected for matrix

135	effects with a $\phi \rho(z)$ algorithm (Pouchou and Pichoir 1991). A time-dependent correction was
136	applied to the data for Na to allow for an intensity decrease during analysis. All others elements
137	exhibited no time-dependence in intensity.
138	No damage from the electron beam was observed. However, as is typical of highly
139	hydrated phases with weakly held H <sub>2</sub> O, bicapite partially dehydrates under vacuum either during
140	carbon coating or in the microprobe chamber. This results in an irregular surface for analysis. In

- addition, the  $H_2O$  loss results in higher concentrations for the remaining constituents than are to
- be expected for the fully hydrated phase. Because an insufficient quantity of bicapite is available
- 143 for a direct determination of  $H_2O$ , that content has been calculated based upon the structure
- determination. Element concentrations are given in Table 1.

145 The empirical formula (based on 42 O apfu in structural unit) is

146  $(K_{1,23}Na_{2,23}Mg_{1,48})_{\Sigma 4.94}[H_{2,51}P_{1,02}(V^{5+}_{13,91}Mo^{6+}_{0.07})_{\Sigma 13.98}O_{42}] \cdot 25H_2O$ . The simplified formula is

147 KNa<sub>2</sub>Mg<sub>2</sub>(H<sub>2</sub>PV<sup>5+</sup><sub>14</sub>O<sub>42</sub>)·25H<sub>2</sub>O, which corresponds to K<sub>2</sub>O 2.35, Na<sub>2</sub>O 3.10, MgO 4.03, P<sub>2</sub>O<sub>5</sub>

148  $3.54, V_2O_563.59, H_2O23.39$ , total 100 wt%.

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## **CRYSTAL STRUCTURE EXPERIMENTAL**

151 The X-ray powder diffraction study was performed using a Rigaku R-Axis Rapid II 152 curved imaging plate microdiffractometer with monochromatized Mo $K\alpha$  radiation. A Gandolfi-153 like motion on the  $\varphi$  and  $\omega$  axes was used to randomize the samples. Observed *d* values and 154 intensities were derived by profile fitting using JADE 2010 software. Data, including unit-cell 155 parameters refined from the powder data using JADE 2010 with whole pattern fitting, are given 156 in Table 2.

157	Single-crystal X-ray studies were carried out using the same instrument and radiation
158	noted above. The Rigaku CrystalClear software package was used for processing the structure
159	data, including the application of numerical and empirical absorption corrections, the latter with
160	the multi-scan approach using ABSCOR (Higashi 2001). The structure was solved by direct
161	methods using SIR2011 (Burla et al. 2012). SHELXL-2016 (Sheldrick 2015) was used for the
162	refinement of the structure. The occupancies of the interstitial cation sites were refined, with the
163	K site refined with full joint occupancy by K and O (H <sub>2</sub> O). A residual peak 2.229(17) Å from the
164	K site was refined as an additional low-occupancy H <sub>2</sub> O site; however, based upon distances to
165	surrounding O sites, it could also accommodate additional Na. All non-hydrogen sites were
166	refined with anisotropic displacement parameters. Difference Fourier syntheses located all H
167	atom positions, except those associated with the partially occupied H <sub>2</sub> O sites and those that
168	protonate the bicapped Keggin anion. The H sites were refined with soft restraints of $0.82(3)$ Å
169	on the O–H distances and 1.30(3) Å on the H–H distances; the $U_{eq}$ of each H was set to 1.2 times
170	that of the donor O atom. Data collection and refinement details are given in Table 3, atom
171	coordinates and displacement parameters in Table 4, selected bond distances in Table 5 and
172	bond-valence sums in Table 6.
173	
174	ATOMIC ARRANGEMENT OF BICAPITE
175	The atomic arrangement of bicapite (Figures 3 and 4) consists of two distinct parts, a
176	structural unit and an interstitial complex, as suggested by Schindler and Hawthorne (2000) for
177	such hydrated minerals with a polyanion. The structural unit is a $[H_2PV^{5+}_{12}O_{40}(V^{5+}O)_2]^{7-}$
178	heteropolyanion composed of twelve distorted $VO_6$ octahedra surrounding a central $PO_4$
179	tetrahedron and capped on opposite sides by two VO <sub>5</sub> square pyramids, which share four of their

180	edges with $VO_6$ octahedra (Figure 3). Without the two capping tetrahedra (and H atoms of the
181	structural unit), the heteropolyanion is the $\alpha$ -isomer of the Keggin anion, $[XM_{12}O_{40}]^{n-}$ , which in
182	this case has the formula $[PV_{12}^{5+}O_{40}]^{15-}$ . In light of the bond valence sum (BVS) values for the
183	V sites all being slightly greater than 5 vu, all V in the mineral is likely to be in the +5 oxidation
184	state. The central PO <sub>4</sub> tetrahedron is disordered, with eight half-occupied O atoms around the P.
185	This is consistent with the P residing at the 0, 0, $\frac{1}{2}$ special position with $\frac{4}{m}$ symmetry. Attempts
186	to refine the structure in a space group that would allow P to occupy a site with -4 symmetry
187	were unsuccessful. It should be noted that this same sort of averaged orientation of the central
188	Keggin tetrahedron has been reported in other structures (cf. Nyman et al. 2007).
189	The interstitial complex includes two octahedrally coordinated cation sites, one occupied
190	by Mg and surrounded by six $H_2O$ groups and one occupied by Na and surrounded by five $H_2O$
191	groups and one O atom. The Mg(H <sub>2</sub> O) <sub>6</sub> octahedron is isolated and only links to other structural
192	components via hydrogen bonding. The NaO(H2O)5 octahedra form dimers by linking through
193	one shared H <sub>2</sub> O group. Each end of the dimer is an O atom shared with the apical O atoms of the
194	VO <sub>5</sub> square pyramids that cap different bicapped Keggin anions. Above the four square
195	"cavities" in each bicapped Keggin anion is the site jointly occupied by K and $H_2O$ . When the
196	site is occupied by K, it forms bonds to the four O atoms at the corners of the cavity, one bond to
197	an O atom in a different bicapped Keggin anion and three bonds to H <sub>2</sub> O groups belonging to the
198	NaO(H <sub>2</sub> O) <sub>5</sub> octahedra. When the K site is occupied by the O atom of an H <sub>2</sub> O group, it
199	presumably forms hydrogen bonds to surrounding O atoms. One such bond is probably to O4,
200	thereby compensating for its otherwise low BVS (1.86 vu).
201	The interstitial complex ideally has the formula [KNa <sub>2</sub> Mg <sub>2</sub> (H <sub>2</sub> O) <sub>25</sub> ] <sup>7+</sup> , which, for charge
202	balance, requires that the structural unit is doubly protonated. Although no specific H sites could

203 be identified on the periphery of the bicapped Keggin anion, among synthetic bicapped Keggin 204 anions, protonation is quite common. For example, Kato et al. (1982) described a similar bicapped Keggin anion for which they gave the formula  $H_n PV_{14}O_{42}^{(9-n)-}$ . In bicapite, the most 205 206 likely candidate O sites for partial OH character are probably O2, O3 and O4, because of their 207 somewhat low BVS values and their locations on the periphery of the structural unit, although as 208 noted above, the low BVS for O4 may be compensated by a hydrogen bond from the H<sub>2</sub>O group 209 partially occupying the K site. The low BVS for the PO<sub>4</sub> group O1 site (1.69 vu) is probably due to steric effects rather than being indicative of partial OH character. For comparison, the O atom 210 211 of the PO<sub>4</sub> group in the similar bicapped Keggin anion reported by Kato et al. (1982) is 1.74 vu 212 corresponding to a P–O bond length of 1.529 Å (identical to the P–O distance in bicapite). 213 A note on bonding within the structural unit and between the structural unit and the 214 interstitial complex is in order. The exterior oxygen atoms of the heteropolyanionic structural unit that protrude into the interstitial unit (O2, O3, O4), known as "terminal" oxygens, are all 215 bound to the vanadium atoms of the structural unit by vanadyl bonds (Table 5; Schindler et al. 216 2000), as is a fourth oxygen atom (O7) that is an also an exterior atom but not a terminal oxygen. 217 Three of these four oxygen atoms are shared with atoms of the interstitial complex (Na-O2, K-218 219 O4, K-O7). In addition, O6, although not bonded to vanadium with a vanadyl bond, is bound to 220 two vanadium atoms in addition to the interstitial, partially occupied. K site. The strongly anisodesmic nature of bonding in the polyanionic structural group is illustrated by the bond 221 valence (in valence units, vu) each of its oxygen atoms receives from the cations of the 222 223 polyanion: O1 = 1.69; O2 = 1.68; O3 = 1.69; O4 = 1.81; O5 = 1.98; O6 = 1.84; O7 = 1.88). Such 224 bonding has implications for the solution behavior of the polyanions and ion transport in the ore-225 forming fluids of the Colorado Plateau.

226 227	IMPLICATIONS
228	Over the past several years, numerous new minerals have been described from the mines
229	of the Colorado Plateau. In addition to the previously described Colorado Plateau mineral
230	sherwoodite (Evans and Konnert 1978) and numerous new $[V_{10}O_{28}]^{6-}$ decavanadate minerals
231	(Kampf et al. 2017a), there is a growing number of recently described Colorado-Plateau minerals
232	with large, complex heteropolyanions. The minerals vanarsite, packratite, morrisonite and
233	gatewayite all contain the $[As^{3+}As^{5+}_{6}V^{4+}_{2+x}V^{5+}_{10-x}O_{51}]^{(11+x)-}$ heteropolyanion, a novel
234	polyoxometalate cluster (Kampf et al. 2016). The mineral kegginite (Kampf et al. 2017b)
235	contains the $[As^{5+}V^{5+}_{12}O_{40}(VO)]^{12-}$ monocapped $\epsilon$ -isomer of the Keggin heteropolyanion, and
236	ophirite contains a heteropolytungstate tri-lacunary Keggin anion [Fe <sup>3+</sup> W <sub>9</sub> O <sub>34</sub> ) <sub>2</sub> ] <sup>11-</sup> (Kampf et al.
237	2014). Herein we describe bicapite, a mineral whose structure contains the
238	$[H_2PV^{5+}_{12}O_{40}(V^{5+}O)_2]^{7-}$ protonated bicapped $\alpha$ -isomer of the Keggin heteropolyanion. Other new
239	minerals of the region that contain large heteropolyions, both anionic and cationic, are currently
240	under study. The mines of the Colorado Plateau are a rich source of minerals with complex
241	heteropolyions that have formed in low-temperature, post-mining mineral assemblages,
242	apparently similar to the "one-pot" bench-scale synthesis of polyoxometalates in inorganic
243	chemistry (Hutin et al. 2013), and obviously forming in a self-assembling environment. The
244	discovery and study of those self-organizing nanostructures advance our knowledge and
245	understanding of mineral complexity on Earth, provide new insights into complex ions that can
246	exist in near-surface environments, and inform researchers in other fields of new avenues to
247	explore in developing phases with potential technological uses. The Colorado Plateau
248	uranium/vanadium deposits encompass the most productive region found to date for naturally
249	occurring polyoxometalate compounds.

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FIGURE 3. Bicapped Keggin anion that is the  $[H_2PV^{5+}_{12}O_{40}(V^{5+}O)_2]^{7-}$  structural unit in bicapite. The O atom sites are numbered. The **c** axis is approximately vertical.

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345						
346	TABLE 1. A	Analytica	l data (wt%) for	bicapite	2	
347						
348	Const.	Mean	Range	S.D.	Standard	Norm.
349	K <sub>2</sub> O	3.49	3.32-3.65	0.17	sanidine	2.89
350	Na <sub>2</sub> O	4.16	3.68-4.52	0.35	albite	3.44
351	MgO	3.59	3.45-3.80	0.16	diopside	2.97
352	$P_2O_5$	4.35	4.15-4.49	0.15	apatite	3.60
353	$V_2O_5$	76.25	75.72-76.69	0.54	V metal	63.04
354	MoO <sub>3</sub>	0.61	0.42-0.77	0.18	Mo metal	0.50
355	H <sub>2</sub> O*					23.57
356	Total					100.01
357						
358	*Based up	oon the c	rystal structure v	with V +	P + Mo = 1	5 and
359	O = 67 ap	ofu.				
360						
361						

362

363

# **TABLE 2**. Observed and calculated powder X-ray diffraction data (*d* in Å) for bicapite

365

Iobs	$d_{\rm obs}$		$d_{ m calc}$	<i>I</i> <sub>calc</sub>	hkl	Iobs	$d_{ m obs}$		$d_{\rm calc}$	Icalc	hkl
100	10.14	5	10.2999	16	002	11	2 5 9 2	5	2.5779	7	420
100	10.14	l	10.0602	84	101	11	2.385	l	2.5750	3	008
6	8.13		8.1519	7	110	1	2 505	5	2.5151	1	404
5	5.897		5.8994	5	103	4	2.303	l	2.4994	2	136
			5.7643	1	$2\ 0\ 0$	5	2.314		2.3136	4	145
			5.1500	1	004				2.2914	1	431
10	5 027	5	5.0301	4	202				2.2609	1	150
10	5.057	J	5.0014	6	121				2.2074	1	406
10	4.348		4.3539	8	114	2	0 1770	5	2.1858	1	503
8	4.142		4.1229	6	213	3	2.1//0	Ì	2.1769	1	228
7	2 9 1 0	5	3.8405	3	204				2.1653	1	237
/	3.810	J	3.7900	3	222	2	2.1234		2.1293	1	521
			3.7777	1	301				2.1032	1	318
2	3.439		3.4367	1	132	1	2 0005	5	2.0920	1	219
7	3.357		3.3534	7	303	4	2.0803	l	2.0702	1	514
4	3.220		3.2186	3	125	2	2 0217	5	2.0438	1	253
6	2 170	ſ	3.1641	1	116	5	2.0317	l	2.0270	1	417
0	5.170	l	3.1596	5	321				1.9665	1	309
29	2.978		2.9755	30	134				1.9214	1	600
			2.9497	1	206	5	1 2020	5	1.8888	2	602
1	1 001	5	2.8986	1	233	5	1.0939	l	1.8873	3	161
4	2.002	l	2.8821	2	$4\ 0\ 0$	2	1 9602	5	1.8691	1	338
11	2.809		2.8102	10	305	3	1.8005	l	1.8612	1	329
3	2.706		2.7173	2	330	Q	1 8100	5	1.8228	2	260
			2.6274	1	332	0	1.0199	Ì	1.8218	4	428

366 Refined unit-cell parameters: a = 11.583(13), c = 20.68(2) Å, V = 2775(7) Å<sup>3</sup>.

# 368 **TABLE 3.** Data collection and structure refinement details for bicapite

369	Diffractometer	Rigaku R-Axis Rapid II					
370	X-ray radiation / power	Mo $K\alpha$ ( $\lambda = 0.71075$ Å)/50 kV, 40 mA					
371	Temperature	293(2) K					
372	Empirical Formula	$H_{42} K_{1.49} Mg_{1.89} Na_{1.93} O_{66.98} P_1 V_{14}$					
373	Space group	4/m					
374	Unit cell dimensions	a = 11.5446(12) Å					
375		c = 20.5460(14)  Å					
376	V	$2738.3(6) \text{ Å}^3$					
377	Ζ	2					
378	Density (for above formula)	$2.434 \text{ g cm}^{-3}$					
379	Absorption coefficient	$2.594 \text{ mm}^{-1}$					
380	<i>F</i> (000)	1974.1					
381	Crystal size	$140 \times 80 \times 30 \ \mu m$					
382	θ range	3.19 to 24.98°					
383	Index ranges	$-13 \le h \le 12, -13 \le k \le 13, -24 \le l \le 24$					
384	Refls collected / unique	$10217 / 1245; R_{int} = 0.057$					
385	Reflections with $I > 2\sigma I$	1008					
386	Completeness to $\theta = 24.98^{\circ}$	99.4%					
387	Max. and min. transmission	0.926 and 0.713					
388	Refinement method	Full-matrix least-squares on $F^2$					
389	Parameters / restraints	141 / 10					
390	GoF	1.100					
391	Final <i>R</i> indices $[F_0 > 4\sigma(F)]$	$R_1 = 0.0465, wR_2 = 0.1133$					
392	<i>R</i> indices (all data)	$R_1 = 0.0578, wR_2 = 0.1214$					
393	Largest diff. peak / hole	$+0.54 / -0.60 e/A^3$					
394	* $R_{\text{int}} = \Sigma  F_o^2 - F_o^2(\text{mean})  / \Sigma [F_o^2 - F_o^2(\text{mean})] / $	$\int_{0}^{2} [GoF = S = \{\Sigma[w(F_{o}^{2} - F_{c}^{2})^{2}]/(n-p)\}^{1/2}. R_{1} = \Sigma  F_{o}  -  F_{c}  /\Sigma F_{o} . wR_{2}]$					
395	$= \{ \sum [w(F_{o}^{2} - F_{c}^{2})^{2}] / \sum [w(F_{o}^{2})^{2}] \}$	$ {}^{1/2}; w = 1/[\sigma^2(F_o^2) + (aP)^2 + bP]$ where <i>a</i> is 0.0578, <i>b</i> is 13.824 and					
396	<i>P</i> is $[2F_c^2 + Max(F_o^2, 0)]/3$ .						
397							

399

# TABLE 4. Atom coordinates and displacement parameters (Å $^2)$ for bicapite

	x/a	y/b	z/c	$U_{ m eq}$	Occupancy	
Р	0	0	0.5	0.0201(7)	1	
V1	0	0	0.31386(8)	0.0295(4)	1	
V2	0.29689(10)	-0.05685(10)	0.5	0.0306(4)	1	
V3	0.18565(7)	0.12493(7)	0.38618(4)	0.0313(3)	1	
Mg	0.5	0	0.25	0.0305(13)	0.944(15)	
Na	0.5	0.5	0.62235(19)	0.0415(17)	0.966(17)	
K/O	0.3951(3)	0.2637(3)	0.5	0.0608(17)	K: 0.372(14	4): O: 0.628(14)
OW	0.4463(12)	0.1529(14)	0.5841(8)	0.025(6)	0.184(11)	),( )
01	0.1059(5)	-0.0230(5)	0.4572(3)	0.0264(14)	0.5	
02	0	0	0.2361(3)	0.0416(18)	1	
03	0.2624(3)	0.1777(3)	0.32880(17)	0.0412(9)	1	
04	0.4277(4)	-0.0938(4)	0.5	0.0469(14)	1	
05	0.0299(3)	0.1487(3)	0.34411(19)	0.0393(9)	1	
06	0.2960(5)	0.0526(3)	0 4346(2)	0.0711(15)	1	
07	0.1688(3)	0.2447(5)	0.4368(2)	0.0792(18)	1	
OW1	0.5	0.500000	0.5	0.060(3)	1	
H1	0.52(2)	0.569(7)	0.5	0.071	1	
OW2	0.5	0	0.3506(3)	0.0447(13)	1	
H2	0.5	-0.021(6)	0.378(2)	0.054	1	
OW3	0.4516(3)	0.021(0) 0.1778(3)	0.2505(2)	0.0428(10)	1	
H3A	0.385(3)	0.186(5)	0.265(3)	0.051	1	
H3B	0.256(5) 0.456(5)	0.229(4)	0.223(2)	0.051	1	
OW4	0.5446(5)	0.223(1) 0.2936(5)	0.6219(3)	0.0746(15)	1	
H4A	0.602(5)	0.2900(2)	0.6219(3)	0.090	1	
H4B	0.550(7)	0.275(7)	0.6599(16)	0.090	1	
	$\frac{U^{11}}{U^{11}}$	$\frac{U^{22}}{U^{22}}$	$\frac{U^{33}}{U^{33}}$	$U^{23}$	$U^{13}$	$U^{12}$
Р	0.0213(10)	0.0213(10)	0.0176(16)	0	0	0
V1	0.0349(6)	0.0349(6)	0.0186(8)	ů 0	Ő	0
V2	0.0254(6)	0.0316(7)	0.0349(7)	ů 0	0	-0 0006(4)
V3	0.0329(5)	0.0386(5)	0.0223(5)	-0.0006(3)	0.0037(3)	-0.0078(3)
Mg	0.0341(16)	0.0341(16)	0.0220(0)	0	0	0
Na	0.050(2)	0.050(2)	0.024(3)	ů 0	0	0 0
K	0.047(2)	0.063(3)	0.023(3)	ů 0	0	0.0022(16)
OW	0.012(9)	0.036(11)	0.072(10)	-0.006(7)	-0.009(7)	-0.005(6)
01	0.012(3)	0.030(3)	0.027(10)	-0.003(3)	0.003(3)	0.002(3)
02	0.052(3)	0.052(3)	0.020(4)	0	0	0
03	0.039(2)	0.054(2)	0.020(1)	0.0046(17)	0,0069(16)	-0.0083(17)
04	0.027(3)	0.038(3)	0.031(2) 0.076(4)	0	0	-0.001(2)
05	0.027(3)	0.030(3) 0.041(2)	0.049(2)	-0.0127(17)	-0 0020(16)	0.001(2) 0.0022(14)
06	0.116(4)	0.033(2)	0.064(3)	0.016(2)	-0.059(3)	-0.021(2)
07	0.035(2)	0.130(4)	0.073(3)	-0.070(3)	-0.014(2)	0.025(2)
OW1	0.069(5)	0.069(5)	0.040(7)	0	0	0
OW2	0.046(3)	0.061(4)	0.028(3)	Õ	õ	0.000(3)
OW3	0.040(2)	0.001(1)	0.020(3)	0.0035(18)	0.0128(18)	0.000(0)
OW4	0.079(4)	0.059(3)	0.086(4)	0.001(3)	0.023(3)	0.015(3)
	•••••	0.007(0)	5.000(1)	5.001(5)		

448	P–O1 (×4)	1.529(6	5)	V3–O3	1.59	6(3)	Na–O2	2.338(8)	
449				V3–O7	1.74	2(4)	Na–OW4 ( $\times$ 4)	2.438(5)	
450	V1–O2	1.597(7	')	V3–O6	1.81	8(4)	Na-OW1	2.514(4)	
451	V1–O5 (×4)	1.858(3	5)	V3–O5	2.01	4(3)	<na-o></na-o>	2.434	
452	<v1–o></v1–o>	1.858		V3–O5	2.03	1(3)			
453				V301	2.38	8(6)	K04	2.833(6)	
454	V2–O4	1.569(5	5)	<v3–o></v3–o>	1.99	9	K–O7 (×2)	2.925(5)	
455	V2–O6 (×2)	1.845(4	l)				K–OW1	2.985(4)	
456	V2–O7 (×2)	1.928(4	)	Mg–OW2	(×2) 2.06	8(5)	K-O6 (×2)	3.009(5)	
457	V201	2.405(6	5)	Mg–OW3	(×4) 2.12	8(4)	K–OW4 (×2)	3.061(6)	
458	<v2–o></v2–o>	1.990		<mg–o></mg–o>	2.11	6	<ko></ko>	2.976	
459									
460	Hydrogen bo	nds							
461	D–H··· $A$		<i>D</i> –Н	$\mathrm{H}^{}A$	$D^{\cdot \cdot \cdot}A$	$\angle D$ –H– $A$			
462	OW1−H1…F	$X(H_2O)$	0.84(3)	2.16(6)	2.985(4)	166(7)			
463	OW2−H2…(	06	0.79(3)	2.22(3)	2.982(7)	162(6)			
464	OW3−H3a···	03	0.83(3)	1.93(3)	2.712(5)	157(5)			
465	OW3–H3b···	·O5	0.82(3)	1.98(3)	2.800(6)	178(6)			
466	OW4–H4a…	07	0.80(3)	2.28(3)	3.074(8)	169(8)			
467	OW4−H4b…	·OW3	0.81(3)	2.44(5)	3.131(8)	144(8)			
468	V-O vanadyl	bonds (	Schindler a	and Hawthor	rne 2000) i	in bold.			
469	2				· · · · · · · · · · · · · · · · · · ·				

# 447 **TABLE 5**. Selected bond distances (Å) and angles (°) in bicapite

### 470 **TABLE 6.** Bond-valence sums for bicapite. Values are expressed in valence units\*

471

	V	Na	Mg	Р	V1	V2	V3	H bonds		2
	K							Acc.	Don.	ዾ
01				1.27 (×4↓)		0.21	$\begin{array}{c} 0.22 \ (\times^{1/2} \downarrow \rightarrow) \\ 0.20 \ (\times^{1/2} \downarrow \rightarrow) \end{array}$			1.69
02		0.22			1.68					1.90
03							1.69	0.22		1.91
O4	0.14 (×0.37→)					1.81				1.86
05					0.86 (×4↓)		0.57 0.55	0.18		2.16
06	$\begin{array}{c} 0.09 \ (\times 2 \downarrow) \\ (\times 0.37 \rightarrow) \end{array}$					0.89 (×2↓)	0.95	0.13		2.00
07	$\begin{array}{c} 0.11 \ (\times 2 \downarrow) \\ (\times 0.37 \rightarrow) \end{array}$					0.72 (×2↓)	1.16	0.12		2.04
OW1	0.10 (×4×0.37→)	$\begin{array}{c} 0.14 \\ (\times 2 \rightarrow) \end{array}$							-0.11 -0.11	0.06
OW2			0.35 (×2↓)						-0.13 -0.13	0.09
OW3			0.31 (×4↓)					0.11	-0.22 -0.18	0.01
OW4	$\begin{array}{c} 0.08 \ (\times 2\overline{\downarrow}) \\ (\times 0.37 \rightarrow) \end{array}$	0.17 (×4↓)							-0.11	0.09
Σ	0.80	1.04	1.94	5.08	5.12	5.23	5.13			

\*Bond valence parameters are from Gagné and Hawthorne (2015). Hydrogen-bond strengths

473 based on O–O bond lengths from Ferraris and Ivaldi (1988).