

## Discreditation of bobdownsite and the establishment of criteria for the identification of minerals with essential monofluorophosphate ( $\text{PO}_3\text{F}^{2-}$ )

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### ABSTRACT

Bobdownsite, IMA number 2008-037, was approved as a new mineral by the Commission on New Minerals, Nomenclature and Classification (CNMNC) as the fluorine end-member of the mineral whitlockite. The type locality of bobdownsite is in Big Fish River, Yukon, Canada, and bobdownsite was reported to be the first mineral with essential monofluorophosphate ( $\text{PO}_3\text{F}^{2-}$ ). The type specimen of bobdownsite has been reinvestigated by electron probe microanalysis (EPMA), and our data indicate that fluorine abundances are below detection in the mineral. In addition, we conducted detailed analysis of bobdownsite from the type locality by gas chromatography isotope ratio mass spectrometry, Raman spectroscopy, EPMA, and NMR spectroscopy. These data were compared with previously published data on synthetic monofluorophosphate salts. Collectively, these data indicate that bobdownsite is indistinguishable from whitlockite with a composition along the whitlockite-merrillite solid solution. Bobdownsite is therefore discredited as a valid mineral species. An additional mineral, krásnoite, has been purported to have monofluorophosphate components in its structure, but reexamination of those data indicate that  $\text{F}^-$  in krásnoite forms bonds with Al, similar to  $\text{OH}^-$  bonded to Al in perhamite. Consequently, krásnoite also lacks monofluorophosphate groups, and there are currently no valid mineral species with monofluorophosphate in their structure. We recommend that any future reports of new minerals that contain essential monofluorophosphate anions be vetted by abundance measurements of fluorine, vibrational spectroscopy (both Raman and FTIR), and where paramagnetic components are permissibly low, NMR spectroscopy. Furthermore, we emphasize the importance of using synthetic compounds containing monofluorophosphate anions as a point of comparison in the identification of minerals with essential monofluorophosphate. Structural data that yield satisfactory P-F bond lengths determined by X-ray crystallography, coupled with direct chemical analyses of fluorine in a material do not constitute sufficient evidence alone to identify a new mineral with essential monofluorophosphate.

**Keywords:** Merrillite, whitlockite, apatite, krásnoite, fluorine, SIMS standard, NMR spectroscopy, hydrogen isotopes

### INTRODUCTION

Phosphate minerals such as apatite, merrillite, and whitlockite are of importance to a wide variety of fields from Earth science to life science, material science, and planetary science (Chew and Spikings 2015; Harlov 2015; Hawthorne 1998; Hughes et al. 2006, 2008; Hughes and Rakovan 2015; Jolliff et al. 2006; McCubbin and Jones 2015; McCubbin et al. 2014; Rakovan and Pasteris 2015; Shearer et al. 2015; Webster and Piccoli 2015).

Phosphates are the major source of P on Earth and their uses range from fertilizers to detergents to insecticides. Synthetic phosphates have been used for ceramics and coatings, and have even been used for the production of fuel cells (i.e., Kendrick et al. 2007; Lin et al. 2007; Pietak et al. 2007). Phosphates have a propensity for concentrating rare earth elements (Jolliff et al. 1993; Prowatke and Klemme 2006; Shearer et al. 2011) that are used pervasively for deciphering sedimentary, igneous, and metamorphic petrogenesis. Furthermore, their ability to accommodate the radioisotopes used for dating makes them important to geochronological studies of rocks (Chew and Spikings 2015).

At present, there are 586 unique phosphate mineral spe-

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cies that have been identified (Hazen et al. 2015; IMA list of minerals updated as of January 2017). Of these, the phosphate anion is most typically represented by phosphate ( $\text{PO}_4^{3-}$ ), where phosphorus is bonded to 4 O atoms; however, there is a subset of phosphate minerals totaling 58 individual mineral species that have  $\text{OH}^-$  or  $\text{F}^-$  substituents for  $\text{O}^{2-}$  on the phosphate molecule. These substituents consist of either hydrogen phosphate anions ( $\text{PO}_3\text{OH}^{2-}$ ) or monofluorophosphate anions ( $\text{PO}_3\text{F}^{2-}$ ). Minerals with protonated phosphate groups constitute 57 of the 58 individual mineral species. The most common of these species is the mineral whitlockite (Fron del 1941; Hughes et al. 2008), although other examples include brushite, monetite, newberyite, and groatite (Cooper et al. 2009; Duff 1971; Frost et al. 2011; Terpstra 1937). Only one mineral species (bobdownsite) is reported to have an essential monofluorophosphate component ( $\text{PO}_3\text{F}^{2-}$ ) in its structure.

Bobdownsite was approved as a new mineral by the International Mineralogical Association in 2008 (IMA number 2008-037), and it was described in detail by Tait et al. (2011). The type locality of bobdownsite is in Big Fish River, Yukon, Canada. Bobdownsite is also reported to occur at several other localities on Earth (e.g., Tip Top Mine, South Dakota), and it has been reported in martian meteorites (Gnos et al. 2002; Tait et al. 2011). In addition to bobdownsite, the mineral krásnoite is reported to contain fluorophosphate groups (Mills et al. 2012), but, unlike bobdownsite, the validity of krásnoite as a species is not predicated on the assignment of fluorine to fluorophosphate groups. Although only two naturally occurring fluorophosphate compounds have been reported, there is a wide array of synthetic fluorophosphate compounds that have many versatile uses from additives in toothpaste (Ericsson 1964) to treatments for osteoporosis (Balena et al. 1998; Kleerekoper 1998; Lems et al. 2000). In the present study, we provide in-depth structural and chemical characterization of bobdownsite from the type locality in Big Fish River, Yukon, Canada, to reassess whether or not fluorine is present, and if so, how it is bonded within the structure. The results of this study will be compared and contrasted with previous studies of bobdownsite, krásnoite, as well as several well-characterized synthetic monofluorophosphate salts. The aim of our study is to establish minimum criteria for identifying and classifying minerals with an essential monofluorophosphate structural component.

## METHODS

### Samples analyzed in the present study

Bobdownsite samples from the type locality in Big Fish River, Yukon, Canada, were obtained from several sources: Samples of bobdownsite were personally collected by coauthor KTT in July 2012 at the same locality that the bobdownsite type sample was collected at Rapid Creek, Yukon, Canada. The mineral was not found in situ, but collected in the talus slope below the original find (samples identified as Bobdownsite\_Tait); bobdownsite specimens were purchased from Rod Tyson of Tysons' Fine Minerals Inc. (samples identified as bobdownsite\_A); and we also obtained the type specimen of bobdownsite from Robert Downs at the University of Arizona (sample number R050109). In addition, Robert Downs provided us with a sample of bobdownsite from the Tip Top Mine in South Dakota (sample number R070654). Finally, we purchased a sample that was described as whitlockite from Steve Covey of Amethyst Galleries' Mineral Gallery from Big Fish River, Yukon, Canada (samples identified as Yukon phosphate) that is identical in appearance to the bobdownsite from the same locality. We suspect that the material is the same as the other material from this locality, which is supported by the fact that the whitlockite specimen was purchased in 2009, prior to the publication of Tait et al.

(2011) describing bobdownsite for the first time.

In addition to using the natural samples described above, we also synthesized whitlockite ( $\text{Ca}_9\text{Mg}[\text{PO}_3\text{OH}][\text{PO}_4]_8$ ) (sample identified as MGS-008) to use as a F- and Fe-free standard to compare with the natural bobdownsite samples. The whitlockite synthesis was conducted using the methods of Adcock et al. (2013, 2014), which are optimized after Hughes et al. (2008) and Gopal et al. (1974). A solution containing 90 mL of high-purity (18.2M $\Omega$ ) water, 1.00 g of laboratory-grade hydroxylapatite (Spectrum, reagent grade) and 0.30 g magnesium nitrate hexahydrate (J.T. Baker, ACS grade) is created in a 125 mL Parr acid digestion vessel (Parr 4748) with an acid-washed polytetrafluoroethylene liner. Once the solution is mixed, it is acidified to a pH of 2.6–2.8 using concentrated phosphoric acid (Alfa Aesar, ACS grade). The vessel is then sealed and incubated in an oven at 240 °C for 7 days. At the end of 7 days, the vessel is removed from the oven and quenched in a water bath in an effort to prevent any further reaction. After cooling, the vessel is opened and the solution decanted leaving the solids. Solid material is rinsed from the vessel using ethanol, allowed to air dry for 24 h, weighed and inspected by optical microscopy for preliminary phase identification. Additional phases can form during synthesis, typically hydroxylapatite ( $\text{Ca}_5[\text{PO}_4]_3\text{OH}$ ) and monetite ( $\text{CaHPO}_4$ ), but are primarily confined to the <75  $\mu\text{m}$  fraction. Output masses are therefore brush sieved on a 200 mesh screen to remove the <75  $\mu\text{m}$  size fraction. The methods have been shown to consistently and reliably produce high-purity Mg-end-member whitlockite (Hughes et al. 2008; Adcock et al. 2013, 2017), although minor amounts of merrillite may occur within the whitlockite (Hughes et al. 2008; Adcock et al. 2017).

### Electron probe microanalysis (EPMA)

All of the natural phosphate specimens (Bobdownsite\_A, Yukon phosphate, Bobdownsite\_Tait, R050109, and R070654) were analyzed using the JEOL 8200 electron microprobe at the Institute of Meteoritics at the University of New Mexico using Probe for EPMA (PFE) software. An accelerating voltage of 15 kV, a nominal probe current of 20 nA, and a beam diameter of 5  $\mu\text{m}$  were used during each analysis. We analyzed for the elements Si, Al, Fe, Mn, Mg, Ca, Na, P, F, and Cl. F was analyzed using a light-element LDE1 detector crystal, and Cl was analyzed using a PET detector crystal. The standards used were as follows; for Ca and P, Durango apatite (Jarosewich et al. 1980) was used as the primary standard, and a natural fluorapatite from India (Ap020 from McCubbin et al. 2012) was used as a secondary check on the standardization. A synthetic  $\text{SrF}_2$  crystal from the Taylor multi element standard mount (C.M. Taylor Co.<sup>1</sup>) was used as the primary F standard, and Ap020 was used as an additional check on the F standardization. We have demonstrated previously that  $\text{SrF}_2$  is a reliable fluorine standard for the analysis of fluorine in both glasses and apatite (McCubbin et al. 2015b, 2016). A synthetic sodalite crystal from Sharp et al. (1989) was used as a primary Cl standard, and scapolite from the Smithsonian Institution (NMNH R6600-1) was used as a secondary check on the Cl standardization. Spessartine, albite, and quartz from the Taylor multi element standard mount (C.M. Taylor<sup>1</sup>) were used as primary standards for Mn, Na, and Si, respectively. Pyrope from the Taylor multi-element standard mount (C.M. Taylor<sup>1</sup>) was used for Mg and Al. Ilmenite from the Smithsonian Institution (NMNH 133868) was used as a primary standard for Fe.

The synthetic Mg-whitlockite (MGS-008) was analyzed using a JEOL JXA-8900 at the EMIL facility within the University of Nevada, Las Vegas. An accelerating voltage of 15 kV, a nominal probe current of 10 nA, and a beam diameter of 10  $\mu\text{m}$  were used during each analysis. We analyzed for the elements Si, Fe, Mn, Mg, Ca, Na, P, S, and Cl. Cl was analyzed using a PETJ detector crystal. Standards included apatite for Ca from the Smithsonian Institution (NMNH 104021) (Jarosewich 2002) and San Carlos olivine for Mg and Si. Fe and Mn were standardized using ilmenite sample NMNH 96189. Kakanui anorthoclase was used as a standard for Na, and scapolite NMNH R6600 was used for Cl. P was standardized using Micro-Analysis Consultants Ltd. Apatite from Yates Mine, Quebec (MAC EMS 80095-MINA apatite). S was standardized using Micro-Analysis Consultants Ltd. pyrite, also from Yates Mine, Quebec (MAC EMS 80095-MINA pyrite).

### Carbon reduction gas chromatography-isotope ratio mass spectrometry

Gas chromatography-isotope ratio mass spectrometry (GC-IRMS) was used to quantify water contents in Bobdownsite\_A and Yukon phosphate samples, and H isotopic values were also determined for the Yukon phosphate sample. Measurements were made in the Center for Stable Isotopes in the Department of Earth and Planetary Sciences at the University of New Mexico, using the technique and apparatus previously described in detail by Sharp et al. (2001). Briefly, the technique involves reduction of structural OH components in the phosphate samples to  $\text{H}_2$  gas

by reaction with glassy carbon at 1450 °C in a helium carrier gas. Product gases are separated in a gas chromatograph and analyzed in a mass spectrometer configured to make hydrogen isotope analyses in continuous flow mode. In the present study, the phosphate samples were wrapped in silver foil and dropped into the furnace using a commercially available autosampler (e.g., Carlo Erba AS 200-LS) mounted directly over the reduction tube. We used sample sizes of about 5–11.5 mg, which produced a sufficient volume of H<sub>2</sub> to quantitatively determine the abundances of H<sub>2</sub>O in the phosphates. The detection limits for H<sub>2</sub>O during our session were determined by running empty silver foil buckets that were used to wrap the phosphate samples, which yielded a detection limit of approximately 300 ppm H<sub>2</sub>O, which is equivalent to approximately 0.006 µL H<sub>2</sub>O that we attribute to adsorbed surface water on the silver. This base level amount of adsorbed surface water is consistent with the blank H<sub>2</sub>O contents from other investigations using this laboratory (McCubbin et al. 2012). This technique was also used to determine δD values using standard correction procedures, which produced results identical to those obtained conventionally with a precision of 4‰ (2σ) for hydrous minerals that have at least 0.1 µL H<sub>2</sub>O (Sharp et al. 2001) (equivalent to 5000 ppm H<sub>2</sub>O in the phosphate samples that were analyzed). Throughout our analysis routine, we checked the reproducibility of the column using several H<sub>2</sub>O standards including NBS 30 biotite (Sharp et al. 2001), water canyon biotite (USGS sample number 3149-11), and BUD biotite (Bindeman and Serebryakov 2011). For each phosphate sample, we analyzed four to seven separate aliquots of sample to test for homogeneity in our phosphate grains. Water contents for the phosphate samples are presented in Table 1. The 2σ uncertainty reported in Table 1 refers to the standard deviation of the mean water contents among all analyses for each sample, which was larger than the analytical uncertainty. Total time of analysis is less than two minutes for a single hydrogen and δD analysis.

## Raman spectroscopy

Raman spectra of Bobdownsite\_A and Yukon phosphate were collected using a Witec α-Scanning Near-Field Optical Microscope (SNOM) confocal Raman spectrometer. Both samples were prepared as polished mounts. The excitation source is a frequency-doubled solid-state YAG laser (532 nm) operating between 0.3 and 1 mW output power (dependent on objective), as measured at the sample using a laser power meter. Objective lenses included a ×100 LWD and a ×20 LWD with a 50 µm optical fiber acting as the confocal pin hole. Spectra were collected

on a Peltier-cooled Andor EMCCD chip, after passing through a f/4 300 mm focal length imaging spectrometer typically using a 600 lines/mm grating. The lateral resolution of the instrument is as small as 360 nm in air when using the ×100 LWD objective, with a focal plane depth of ~800 nm.

We employed this instrument to collect single spectra of the Bobdownsite\_A and Yukon phosphate samples. Single spectra mode allows the acquisition of a spectrum from a single spot on the target. Average spectra are produced using integration times of 30 s per accumulation and 10 accumulations to allow verification of weak spectral features. A cosmic ray reduction routine was used to reduce the effects of stray radiation on Raman spectra.

## Nuclear magnetic resonance (NMR) spectroscopy

NMR spectra were collected on Bobdownsite\_A, Yukon phosphate, and the synthetic Mg-whitlockite sample, MGS-008. The <sup>19</sup>F single-pulse (SP) spectra were collected on a 500 MHz (11.7 T) Varian InfinityPlus spectrometer operating at 470.179 MHz, using a Varian/Chemagnetics T3-type probe assembly configured for 3.2 mm (O.D.) rotors. The 90° pulse width was 5 µs, and 170 transients were collected at a relaxation delay of 100 s. Longer relaxation delays produced nominal increases in signal, however the NMR line shapes did not change at relaxation delays up to 600 s. Chemical shifts were measured relative to neat CFC1<sub>3</sub> set to δ<sub>F</sub> = 0 ppm. The MAS rate was 20 kHz. Additional spectra were acquired after a small amount of NaF was added to each sample, corresponding approximately to one F per formula unit (i.e., an F:P ratio of 1:7). The <sup>1</sup>H SP/MAS spectra were collected on a 400 MHz (9.4 T) Varian Inova spectrometer operating at 399.895 MHz. A probe assembly modified for low <sup>1</sup>H background and configured for 4 mm (O.D.) rotors was used, with the samples contained in ZrO<sub>2</sub> rotors having Kel-F tips, and PTFE spacers. The MAS rate was 10 kHz. The 90° pulse width was 5 µs with a pulse delay of 30 s, and 128 transients were collected. Chemical shifts were measured relative to hydroxylapatite set to δ<sub>H</sub> = 0.2 ppm. The <sup>31</sup>P NMR spectra were acquired with the 500 MHz spectrometer and a T3-type probe fitted with 4 mm spinning assembly and at a 15 kHz spinning rate. The single-pulse spectra were collected with 4 µs pulses, where the 90° pulse width was 5 µs. For the synthetic samples, spectra collected fully relaxed as a single acquisition after overnight equilibration did not differ significantly from spectra acquired with a 300 s relaxation delay. The <sup>31</sup>P chemical shifts are referenced to 85% H<sub>3</sub>PO<sub>4</sub>, set to δ<sub>P</sub> = 0 ppm.

**TABLE 1.** Average electron microprobe analyses of phosphates analyzed in present study

Oxide	Bobdownsite_A	Yukon phosphate	Bobdownsite_Tait	R050109	R070654	MGS-008
P <sub>2</sub> O <sub>5</sub>	45.52(71)	46.6(2)	46.94(39)	47.07(29)	46.03(52)	46.98(66)
SiO <sub>2</sub>	0.09(3)	0.00(0)	0.10(10)	0.00(0)	0.00(0)	0.00(0)
Al <sub>2</sub> O <sub>3</sub>	0.37(12)	0.34(14)	0.30(15)	0.01(1)	0.00(0)	n.d.
FeO	1.09(11)	1.10(19)	1.43(40)	1.33(10)	0.01(1)	0.00(0)
MnO	0.02(1)	0.01(1)	0.02(2)	0.02(1)	0.02(3)	0.00(0)
MgO	2.87(8)	2.88(13)	2.63(19)	2.77(11)	3.53(8)	3.75(10)
CaO	47.19(50)	46.5(2)	47.04(40)	47.46(21)	47.88(19)	47.65(36)
Na <sub>2</sub> O	0.51(22)	1.00(31)	0.72(29)	0.61(18)	0.19(0.04)	0.00(0)
H <sub>2</sub> O	0.71(4)	0.70(5)	n.d.	n.d.	n.d.	n.d.
SO <sub>3</sub>	0.00(0)	n.d.	n.d.	n.d.	n.d.	0.16(9)
F	0.00(0)	0.00(0)	0.00(0)	0.00(0)	0.00(0)	n.d.
Cl	0.00(0)	0.00(0)	0.01(1)	0.00(0)	0.00(0)	0.01(1)
–O=F	0	0	0	0	0	–
–O=Cl	0	0	0	0	0	0
Total	98.37	99.2	99.20	99.26	97.65	99.41
N	78	54	66	101	102	10
<b>Structural formulas based on 28 O atoms</b>						
P	6.92	7.01	7.01	7.03	6.97	6.99
Si	0.02	0.00	0.02	0.00	0.00	0.00
Al	0.08	0.07	0.06	0.00	0.00	–
Fe	0.16	0.16	0.21	0.20	0.00	0.00
Mn	0.00	0.00	0.00	0.00	0.00	0.00
Mg	0.77	0.76	0.69	0.73	0.94	0.98
Ca	9.09	8.86	8.90	8.98	9.18	8.98
Na	0.18	0.34	0.25	0.21	0.07	0.00
S	0.00	–	–	–	–	0.02
Σ Cations	17.22	17.21	17.14	17.14	17.16	16.95
F	0.00	0.00	0.00	0.00	0.00	–
Cl	0.00	0.00	0.00	0.00	0.00	0.00
Σ Anions						
OH	0.86	0.83	0.85 <sup>a</sup>	0.85 <sup>a</sup>	0.85 <sup>a</sup>	1.00 <sup>b</sup>

Notes: N = number of analyses. – = a value was not computed. n.d. = abundance was not determined. Parenthetical values represent the uncertainty in the reported abundance.

<sup>a</sup> Value is based on the average OH from Bobdownsite\_A and Yukon phosphate to aid in accurate structural formula determination.

<sup>b</sup> Value is assumed based on stoichiometry and used to calculate an appropriate structural formula.

## RESULTS

### Chemical and isotopic composition of phosphates

The compositions of all the phosphates analyzed in the present study are provided in Table 1, and the H<sub>2</sub>O abundances of samples Bobdownsite\_A and Yukon phosphate are also provided in Table 1. All of the phosphates from Big Fish River, Yukon, Canada, have similar compositions with some variations in Na, Mg, and Fe (Table 1). Sodium abundances in these samples range from 0.61 to 1.00 wt% Na<sub>2</sub>O, which corresponds to 0.21–0.34 structural formula units of Na (sfu) per 28 O atoms. Fe and Mg abundances in the samples from Big Fish River are inversely correlated and range from 1.08 to 1.43 wt% FeO and 2.63 to 2.88 wt% MgO. These abundances correspond to 0.16 to 0.21 sfu Fe and 0.69 to 0.73 sfu Mg per 28 O atoms. The sum of Fe and Mg per 28 O atoms ranges from 0.90 to 0.94 sfu. Although H<sub>2</sub>O abundances were only measured in the samples Bobdownsite\_A and Yukon phosphate, they yielded similar results of 0.71 and 0.70 wt% H<sub>2</sub>O, respectively. These abundances of H<sub>2</sub>O correspond to 0.83 to 0.86 sfu OH per 28 O atoms. Fluorine was not detected in any of the phosphates analyzed in the present study, so the chemistry of the phosphates indicates that they are whitlockite with compositions that lie along the whitlockite-merrillite solid solution.

The sample (R070654) from the Tip Top mine, South Dakota, was distinctly different from the samples from Big Fish River, Yukon, Canada, given the paucity of FeO, lower abundance of Na, and elevated abundance of Mg (Table 1). We did not detect any fluorine within sample R070654, so it is also likely a phase that lies along the whitlockite-merrillite solid solution; however direct analysis of H<sub>2</sub>O is required to definitively identify the sample as either whitlockite or merrillite. The synthetic sample MGS-008 is an end-member Mg-whitlockite and does not have detectable Fe or Na (Table 1).

In addition to elemental abundances, we measured the H-isotopic composition of the sample Yukon phosphate. Isotopic compositions are traditionally expressed in  $\delta$  notation according to the following equation:

$$\delta^M X = \left[ \frac{\frac{M}{\text{Heavy}} X_{\text{Measured}} \times \frac{M}{\text{Light}} X_{\text{Standard}}}{\frac{M}{\text{Light}} X_{\text{Measured}} \times \frac{M}{\text{Heavy}} X_{\text{Standard}}} - 1 \right] \times 1000 \quad (1)$$

where  $\frac{\frac{M}{\text{Heavy}} X_{\text{Measured}}}{\frac{M}{\text{Light}} X_{\text{Measured}}}$

is the isotopic ratio of the measured sample,  $\frac{\frac{M}{\text{Light}} X_{\text{Standard}}}{\frac{M}{\text{Heavy}} X_{\text{Standard}}}$

is the isotopic ratio of the standard, and  $\delta^M X$  is the  $\delta$  value of the isotope expressed in per mil. The standard for D/H is Vienna Standard Mean Ocean Water with D/H of  $1.5576 \times 10^{-4}$  (Coplen 1994). The H isotopic composition of the sample Yukon phosphate has a  $\delta D$  value of  $-200 \pm 12\text{‰}$ , which is similar to meteoric waters in northern Yukon, Canada, where the samples originated (e.g., Tondou et al. 2013), and it is distinct from meteoric waters in Albuquerque, New Mexico, where the samples were analyzed and stored prior to analysis (e.g., Yapp 1985).

### Raman spectroscopy

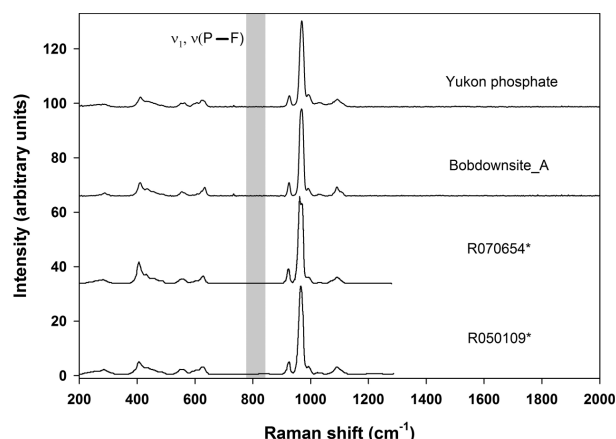
We analyzed the samples Bobdownsite\_A and Yukon phosphate by Raman spectroscopy, and Raman spectra for samples R050109 and R070654 were reported by Tait et al. (2011). Spectra from all four samples are displayed in Figure 1. All spectra exhibit the same patterns, consistent with being comprised of similar material. Peaks in the range of 900 to 1180 cm<sup>-1</sup> are associated with P-O vibrations (Socrates 2001). The most prominent peak in this region is represented by the symmetric P-O stretch ( $\nu_1$ ) in phosphate at ~970 cm<sup>-1</sup>. The broad weak peaks in the range of 1070–1120 cm<sup>-1</sup> are representative of the asymmetric P-O stretching ( $\nu_3$ ) in phosphate. The sharp medium to weak intensity peak at approximately 923–925 cm<sup>-1</sup> is in the region that has been attributed to the symmetric stretching mode ( $\nu_1$ ) of HPO<sub>4</sub><sup>2-</sup> in whitlockite (Jolliffe et al. 2006). We observe no peaks in the region of the symmetric P-F stretching mode ( $\nu_1$ ) of 776 to 840 cm<sup>-1</sup> range (highlighted in gray in Fig. 1) reported previously for the P-F symmetric stretch ( $\nu_1$ ) in PO<sub>3</sub>F<sup>2-</sup> from monofluorophosphate salts (Baran and Weil 2009; Weil et al. 2015, 2007).

### <sup>19</sup>F and <sup>1</sup>H NMR

<sup>19</sup>F is the second most sensitive naturally occurring stable nuclide for NMR spectroscopy, with a detection limit estimated to be of the order 100  $\mu\text{g F g}^{-1}$ , but no <sup>19</sup>F MAS/NMR signal was detected from the Bobdownsite\_A, Yukon phosphate, and synthetic whitlockite (MGS-008) samples. To check this result, additional spectra were acquired after addition of small amounts of NaF (4.5–6 wt%), equivalent to approximately one F per formula unit (i.e., an F:P ratio of 1:7). Spectra acquired after addition of the NaF contain a central peak at  $\delta_F = -220$  ppm and associated spinning sidebands (Fig. 2) but no other signals. This chemical shift is in good agreement with previous reports for NaF (Stebbins and Zeng 2000). Previous <sup>19</sup>F NMR studies of inorganic fluorophosphate compounds show spectra containing wide spinning sideband manifolds with distinct center band doublets for the fluorophosphate F. The <sup>19</sup>F chemical shifts for the fluorophosphate group range from -52.5 ppm for BaPOF<sub>3</sub> to -75 ppm for Na<sub>2</sub>PO<sub>3</sub>F (Weil et al. 2004, 2007; Zhang et al. 2007; Stoeger et al. 2013; Jantz et al. 2016). No signal in this chemical shift range (shaded region of Fig. 2) occurs in the spectra of the Bobdownsite\_A and Yukon phosphate samples from Big Fish River, Yukon, Canada.

The synthetic whitlockite sample yields a well-resolved, relatively narrow <sup>1</sup>H NMR peak near  $\delta_H = +10$  ppm (Fig. 3), consistent with the presence of a single hydrogen phosphate group in the crystal structure. Additional minor signals occur near +7 and +12 ppm, possibly from unidentified impurities. The <sup>1</sup>H NMR spectra of the natural Bobdownsite\_A and Yukon phosphate samples also contain a major peak centered near +10 ppm, although the peaks are significantly broader and exhibit substantially more spinning sideband intensity than the synthetic whitlockite. The broader resonances likely result from minor concentrations of paramagnetic ions in the mineral specimens (e.g., Begaudeau et al. 2012; Oldfield et al. 1983), which is consistent with the 1.1 to 1.4 wt% FeO in these samples (Table 1). An additional minor peak occurs in the synthetic whitlockite near +0.2 ppm that can be attributed to an impurity phase, possibly hydroxylapatite based on its similar chemical shift (Yesinowski and Eckert 1987).





**FIGURE 1.** Raman spectra of Yukon phosphate, Bobdownsite\_A, bobdownsite type specimen R050109, and bobdownsite from Tip top Mine, South Dakota (R070654). The sample names denoted with an asterisk indicate the data were reported in Tait et al. (2011). The region that is characteristic of P-F symmetric stretching modes in fluorophosphate salts is indicated by the shaded area (Baran and Weil 2009; Heide et al. 1985; Jantz et al. 2016; Weil et al. 2015, 2004, 2007; Zeibig et al. 1991).

### <sup>31</sup>P NMR

The <sup>31</sup>P MAS/NMR spectra of the Bobdownsite\_A and Yukon phosphate specimens contain broad, poorly resolved centerbands centered near +0.6 and 5.5 ppm FWHM (Fig. 4). In contrast, the synthetic whitlockite (MGS-008) sample yields well-resolved centerbands exhibiting sharp peaks at 2.3, 1.4, and -0.2 ppm plus shoulders at +0.7 and -0.8 ppm. Minor additional peaks occur at -3.4 and -4.3 ppm that could arise from impurity phases. Qualitatively, the centers of gravity of the Bobdownsite\_A and Yukon phosphate spectra coincide with that of the synthetic whitlockite (MGS-008). This result suggests that peak broadening by paramagnetic substituents present in the natural samples, similar to that discussed for the <sup>1</sup>H results above, prevents resolution of distinct sites. However, the spectrum of the natural Yukon phosphate sample does appear to exhibit an asymmetry that suggests the presence of at least two peaks at positions similar to those for the main peaks for the synthetic sample, namely at +2.3 and -0.2 ppm. The chemical shifts for the main peaks observed for these samples lie well within ranges reported previously for orthophosphate groups in general, and Ca- and Mg-orthophosphates in particular (Rothwell et al. 1980; Turner et al. 1986; Belton et al. 1988).

The spectrum of the synthetic whitlockite (MGS-008) is consistent with the presence of three crystallographically distinct P positions in a 3:1:3 ratio plus additional signals arising from merrillite-like local environments (Calvo and Gopal 1975). A more complete report of the NMR spectra of whitlockite and assignment of the resolved features will be presented in a separate paper.

### DISCREDITATION OF BOBDOWNSITE

Based on the similarities in chemical composition and the nearly identical Raman spectra among all phosphate samples

analyzed from Big Fish River, Yukon, Canada, we infer that all samples from this locality are comprised of the same mineral phase. Consequently, we refer to all of these samples collectively as “bobdownsite” throughout the discussion except in cases where we refer to a specific sample, in which case the sample name is used.

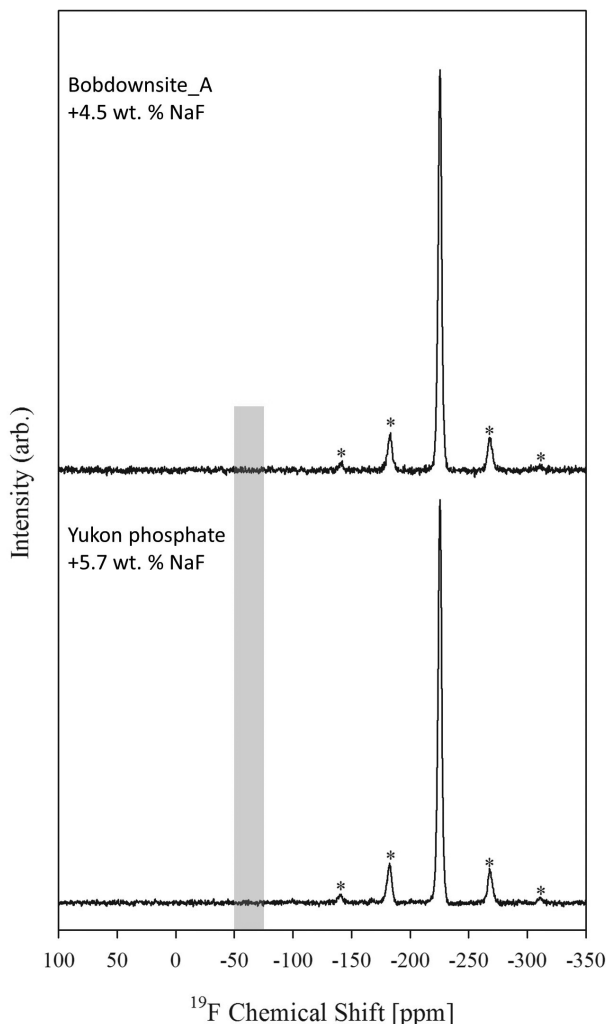
Three of the primary lines of evidence that supported the identification of bobdownsite require reassessment based on differences in results between the present study and the initial description of this mineral (i.e., Tait et al. 2011). These include the absence of OH<sup>-</sup> in the samples inferred by thermogravimetric analysis (TGA), the assignment of a Raman peak at 923 cm<sup>-1</sup> to symmetric stretching in PO<sub>3</sub>F<sup>2-</sup>, and the abundance of fluorine determined by EPMA. We will first compare data from Tait et al. (2011) and the present study and then use NMR results from the present study to resolve any remaining discrepancies.

### H abundance of “bobdownsite”

Samples of “bobdownsite” heated to 1100 °C by Tait et al. (2011) did not exhibit weight loss that could be attributed to the presence of OH in the structure. However, in the present study, “bobdownsite” was heated to 1450 °C and H was liberated from the sample, yielding approximately 0.7 ± 0.05 wt% H<sub>2</sub>O with an H isotopic composition of -200 ± 12‰. The abundance of H<sub>2</sub>O fills the equivalent of at least 83% of the OH<sup>-</sup> site in whitlockite with OH<sup>-</sup>. The presence of OH<sup>-</sup> in the “bobdownsite” structure is confirmed by our observation of <sup>1</sup>H NMR signals for a hydrogen phosphate group in samples from the Big Fish River, Yukon, Canada, locality.

### Vibrational spectroscopy of “bobdownsite”

The fluorophosphate group occurs in several inorganic salts for which crystal structures and vibrational spectra have been reported (Heide et al. 1985; Zeibig et al. 1991; Weil et al. 2004, 2007, 2015; Baran and Weil 2009; Jantz et al. 2016). Owing to a lack of significant bonding of fluorine beyond the P-F bond in most of these compounds, the P-F stretching frequency is typically observed to vary only slightly from its value for aqueous fluorophosphate ion, 795 cm<sup>-1</sup> (Siebert 1966). A small variation in the symmetric P-F stretching mode has been observed, ranging from 776 to 840 cm<sup>-1</sup> that in some cases appears to relate to the P-F bond distance (Heide et al. 1985; Zeibig et al. 1991; Baran and Weil 2009; Weil et al. 2004, 2007, 2015; Jantz et al. 2016). The wavenumber range of stretching modes attributed to P-F bonds in fluorophosphate salts falls outside that typical for the symmetric P-O stretch in phosphate and hydrogen phosphate. In contrast, Tait et al. (2011) analyzed bobdownsite by Raman spectroscopy, assigning a peak at 923 cm<sup>-1</sup> to the P-F ν<sub>1</sub> mode in PO<sub>3</sub>F<sup>2-</sup>. However, 923 cm<sup>-1</sup> falls outside the range reported for P-F symmetric stretching modes exhibited by synthetic fluorophosphate salts (Heide et al. 1985; Zeibig et al. 1991; Weil et al. 2004, 2007, 2015; Baran and Weil 2009; Jantz et al. 2016). Our analyses of bobdownsite\_A and Yukon phosphate match those of Tait et al. (2011) for samples R050109 and R070654, which supports the argument that a fluorophosphate group attributed to “bobdownsite” is misidentified (Fig. 1), despite the fact that the refined P-F bond distance from the XRD results of Tait et al. (2011) falls within the range of crystalline fluorophosphate salts [although substantially longer than previous reports for all except NaK<sub>3</sub>(PO<sub>3</sub>F)<sub>2</sub>; Durand

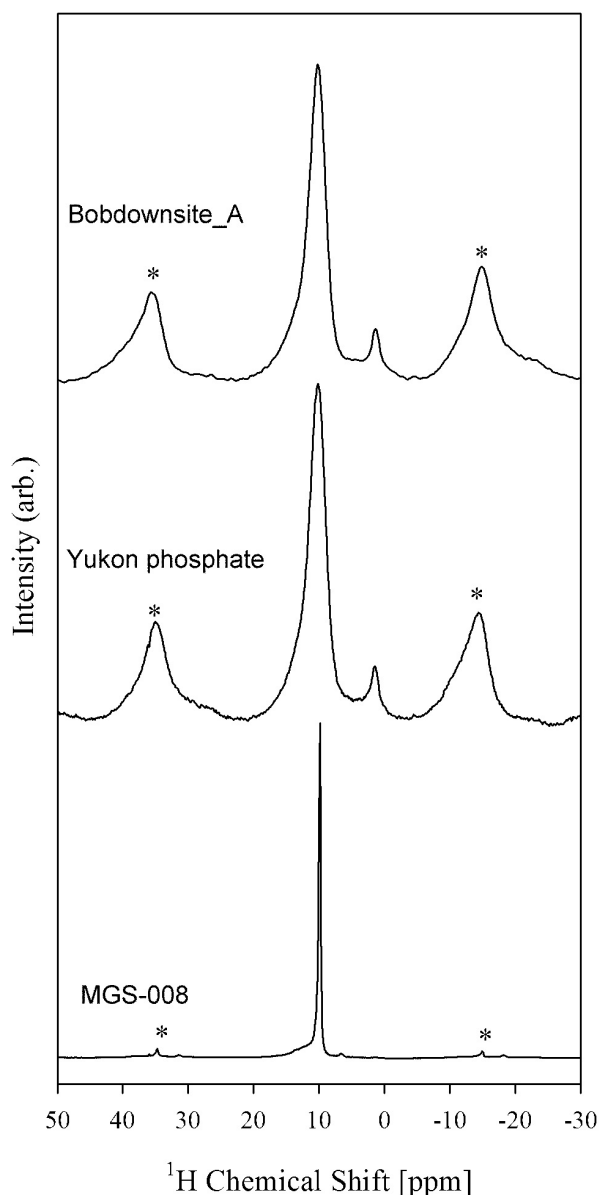


**FIGURE 2.**  $^{19}\text{F}$  MAS/NMR spectra of Bobdownsite\_A (top) and Yukon phosphate (bottom) samples with the indicated amounts of added NaF, which approximate one F atom per formula unit (7 P). Shaded region corresponds to chemical shift range for previous reports of compounds containing fluorophosphate groups (see text). Spectra acquired by direct excitation with a 100 s relaxation delay and a spinning rate of 20 kHz. Asterisks denote spinning sidebands.

et al. 1975]. Finally, the Raman spectra of the purported “bobdownsite” specimens are strikingly similar to those reported for a whitlockite specimen from the Palermo type locality (R080052) in Tait et al. (2011), which also exhibits a peak at about  $923\text{ cm}^{-1}$ . Peaks in vibrational spectra near  $923\text{ cm}^{-1}$  have been attributed in previous studies to the P-O stretch for the hydroxyl oxygen in the hydrogen phosphate group (e.g., Socrates 2001; Jolliff et al. 2006). Assignment of the  $923\text{ cm}^{-1}$  peak to the P-O(H) vibration is consistent with our observation of  $^1\text{H}$  NMR signals for a hydrogen phosphate group in samples from these localities.

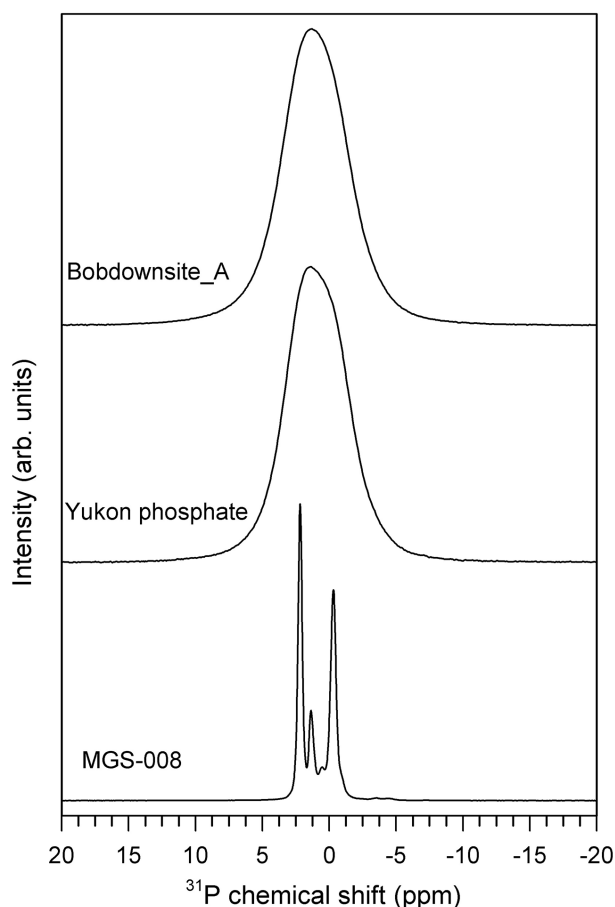
#### F abundance of “bobdownsite”

Although Tait et al. (2011) reported fluorine abundances by EPMA in samples R050109 and R070654 of 1.9 and 1.5 wt% F,



**FIGURE 3.** Comparison of the center band region of  $^1\text{H}$  MAS/NMR spectra of Bobdownsite\_A (top) and Yukon phosphate (middle) specimens and synthetic whitlockite sample MGS-008 (bottom). Spectra acquired by direct excitation with a 30 s relaxation delay at a spinning rate of 10 kHz. Spinning sidebands are denoted by asterisks and for the natural phosphate samples extend for at least four more orders owing to broadening by paramagnetic ions.

respectively, fluorine was below detection (detection limit was approximately 0.05 wt% F) in our EPMA analyses of R050109 and R070654 as well as the other phosphate samples from Big Fish River, Yukon, Canada, that we analyzed. To accurately measure fluorine by EPMA, a synthetic multilayer crystal with a large  $d$ -spacing (e.g., LDE-1, OV-60, PC0) is typically preferred over the more classically used and widely available thallium acid phosphate (TAP) crystal because the intensity of the  $\text{FK}\alpha$  peak using the multilayer crystal is approximately 14 times higher than



**FIGURE 4.** Comparison of  $^{31}\text{P}$  SP-MAS NMR spectra of Bobdownsite\_A (top) and Yukon phosphate (middle) specimens and synthetic whitlockite sample MGS-008 (bottom). Spectra were acquired with direct excitation (single-pulse) with a 300 s relaxation delay at a spinning rate of 15 kHz. Spinning sidebands fall outside the displayed spectral region. Highly resolved spectra for synthetic samples result from absence of paramagnetic substituents.

with TAP (Potts and Tindle 1989; Raudsepp 1995; Reed 2005). This enhanced intensity in the  $\text{FK}\alpha$  all third order or greater interferences on the  $\text{FK}\alpha$  line insignificant, which was one of the primary problems with analyzing F in apatite using a TAP crystal (Potts and Tindle 1989; Raudsepp 1995; Reed 2005). Furthermore, the primary difference in the EPMA routines between our study and that of Tait et al. (2011) is our use of an LDE-1 crystal to measure fluorine X-ray counts vs. use by Tait et al. (2011) of a TAP crystal (Robert Downs, personal communication); this may explain the differences between our results. We further investigated whether or not F was present in the “bobdownsite” using  $^{19}\text{F}$  MAS/NMR spectroscopy, which confirmed that F was below detection ( $\sim 100$  ppm F) in the “bobdownsite” samples, consistent with our EPMA results.

#### Whitlockite from Big Fish River, Yukon, Canada

The mineral “bobdownsite” is differentiated from whitlockite based on the presence of a monofluorophosphate group ( $\text{PO}_3\text{F}^{2-}$ ) rather than a protonated phosphate group ( $\text{PO}_3\text{OH}^{2-}$ ).

We were unable to detect fluorine in any of the phosphate specimens from Big Fish River, Yukon, Canada, including the type specimen R050904. Furthermore, measured  $\text{H}_2\text{O}$  abundances were at a level consistent with a whitlockite along the whitlockite-merrillite join. Consequently, we regretfully conclude that the mineral bobdownsite must be discredited.

#### IMPLICATIONS FOR STRUCTURAL ACCOMMODATION OF FLUORINE IN KRÁSNOITE

The mineral krásnoite [ $\text{Ca}_3\text{Al}_{7.7}\text{Si}_3\text{P}_4\text{O}_{22.9}(\text{OH})_{13.3}\text{F}_2 \cdot 8\text{H}_2\text{O}$ ] is reported to be the fluorine analog of the mineral perhamite (Mills et al. 2011). In a follow-up manuscript describing the mineral krásnoite, Mills et al. (2012) concluded that the fluorine was substituting for  $\text{OH}^-$  on a hydrogen phosphate anion ( $\text{PO}_3\text{OH}^{2-}$ ), forming a fluorophosphate group rather than substituting for one of the other  $\text{OH}^-$  sites available in the mineral. Although krásnoite is a valid species regardless of the structural accommodation mechanism of fluorine, we have examined the assignment of fluorine to a monofluorophosphate component given the paucity of such molecular groups in other minerals as exemplified by the discreditation of bobdownsite. There are two primary lines of evidence used by Mills et al. (2012) to support the occurrence of monofluorophosphate groups in krásnoite, including vibrational (Raman and IR) spectroscopic data and  $^{19}\text{F}$  MAS NMR data. We compare the data in Mills et al. (2012) with those published on synthetic monofluorophosphate salts to further elucidate the structural role of F in krásnoite.

Mills et al. (2012) reported both Raman and FTIR data for krásnoite. They assigned a Raman peak at  $920\text{ cm}^{-1}$  to the P-F symmetric stretching mode in  $\text{PO}_3\text{F}$  based on the same assignment in bobdownsite from Tait et al. (2011); however, this peak position falls outside the range of the F-P  $\nu_1$  mode in  $\text{PO}_3\text{F}$  from fluorophosphate salts (Heide et al. 1985; Zeibig et al. 1991; Weil et al. 2004, 2007, 2015; Baran and Weil 2009; Jantz et al. 2016), instead falling within the range of stretching and bending modes attributed to P-O or P-O(H) groups (Socrates 2001; Jolliff et al. 2006). Therefore, the  $920\text{ cm}^{-1}$  Raman peak is likely not indicative of a monofluorophosphate group in krásnoite. A peak at  $820\text{ cm}^{-1}$  within the FTIR spectrum of krásnoite was also assigned to a fluorophosphate group by Mills et al. (2012). The position of the P-F  $\nu_1$  modes for fluorophosphate salts ranges from  $727$  to  $824\text{ cm}^{-1}$  (Baran and Weil 2009; Weil et al. 2004, 2007, 2015; Jantz et al. 2016), so the assignment of a P-F  $\nu_1$  mode to the  $820\text{ cm}^{-1}$  in krásnoite is permissive; nevertheless, the lack of a corresponding Raman peak to corroborate this assignment does not lend confidence to the interpretation, especially given overlaps in the  $820\text{ cm}^{-1}$  region with several other peaks related to Al-O or Si-O vibrational modes (Socrates 2001). Consequently, the vibrational spectroscopic data do not provide sufficient evidence that krásnoite contains monofluorophosphate groups within its structure.

Mills et al. (2012) reported  $^{19}\text{F}$  MAS NMR results for krásnoite that were interpreted to verify the presence of F in the mineral structure. Because chemical shifts reported by Mills et al. (2012) were relative to  $\text{CaF}_2$ , we have converted their values relative to neat  $\text{CFCl}_3$ , the conventional standard for  $^{19}\text{F}$  NMR spectral comparisons (e.g., Huve et al. 1992). The chemical shift of  $^{19}\text{F}$  in  $\text{CaF}_2$  relative to neat  $\text{CFCl}_3$  is  $-110$  ppm (Wang and Grey 1998), which corresponds to a new primary peak at a chemical shift of

–130 ppm with minor peaks at chemical shifts of –110, –140, and –150 ppm. The  $^{19}\text{F}$  chemical shifts for the monofluorophosphate groups in fluorophosphate salts ranges from –52.5 ppm for  $\text{BaPOF}_3$  to –75 ppm for  $\text{Na}_2\text{PO}_3\text{F}$  (Weil et al. 2004, 2007; Zhang et al. 2007; Stoeger et al. 2013; Jantz et al. 2016), which are substantially displaced from the observed  $^{19}\text{F}$  chemical shifts for krásnoite (Mills et al. 2012). Consequently, the  $^{19}\text{F}$  MAS NMR data in Mills et al. (2012) do not support assignment of F in krásnoite to monofluorophosphate groups. Furthermore, the  $^{19}\text{F}$  NMR peak for monofluorophosphate should be represented by a doublet owing to the strong scalar coupling between  $^{19}\text{F}$  and  $^{31}\text{P}$  nuclei (e.g., Stoeger et al. 2013). In addition, Mills et al. (2012) attributed the peak at a chemical shift of –150 ppm to a minor fluorapatite impurity, whereas  $^{19}\text{F}$  NMR spectra of fluorapatite exhibit a chemical shift near –103 ppm (e.g., Braun et al. 1995; McCubbin et al. 2008; Mason et al. 2009), which is not exhibited by the  $^{19}\text{F}$  NMR spectra for krásnoite. To further elucidate the role of F in the structure of krásnoite, we have interpreted the  $^{19}\text{F}$  NMR spectra accordingly and show that the chemical shifts observed at –130, –140, and –150 ppm can be ascribed to Al-F groups (Kiczinski and Stebbins 2002; Zhang et al. 2002; Chupas et al. 2003; Kao and Chen 2003), indicating that  $\text{F}^-$  likely substitutes for  $\text{OH}^-$  bonded to Al in perhamite (i.e., Mills et al. 2006).

#### PROPOSED CRITERIA FOR IDENTIFICATION OF MINERALS WITH A MONOFLUOROPHOSPHATE ( $\text{PO}_3\text{F}^{2-}$ ) ANION

We recommend that future reports of new minerals containing essential monofluorophosphate anions be vetted by abundance measurements of fluorine, vibrational spectroscopy (both Raman and FTIR), and where paramagnetic components are permissibly low, NMR spectroscopy. Furthermore, the previous reports of bobdownsite and krásnoite highlight the importance of using synthetic compounds containing monofluorophosphate anions as a point of comparison in the identification of minerals with essential monofluorophosphate. Given that fluoride is a common substituent for hydroxyl in many phosphates and occupies crystallographic sites in phosphates that are not bonded directly to phosphorus, a concerted effort should also be made to verify that a proposed new monofluorophosphate mineral contains  $\text{PO}_3\text{F}$  molecular groups, or at minimum, P-F bonds. Although not a shortcoming of the bobdownsite and krásnoite studies (Tait et al. 2011; Mills et al. 2012), it must be emphasized that structural data that yield satisfactory P-F bond lengths determined by X-ray crystallography, coupled with direct chemical analyses of fluorine in a material, do not constitute sufficient evidence alone to identify a new mineral having essential monofluorophosphate anions.

#### IMPLICATIONS

The preponderance of data collection on whitlockite from Big Fish River, Yukon, Canada, has highlighted the potential utility of this material as a phosphate standard for H abundance and H-isotopic analysis by secondary ion mass spectrometry (SIMS). Given the heightened interest in the analysis of hydrogen abundances and hydrogen isotopes in phosphates from planetary materials (i.e., Bector et al. 2003; Greenwood et al. 2008, 2011; Boyce et al. 2010, 2012; McCubbin et al. 2010, 2012, 2014; Barnes et al. 2013, 2014; Robinson and Taylor 2014; Sarafian et al. 2013, 2014; Tartèse et al. 2013, 2014a, 2014b; Treiman et

al. 2014; Jones et al. 2014, 2016; Barrett et al. 2016; Mane et al. 2016; Robinson et al. 2016), the identification of whitlockite with consistent inter- and intra-granular  $\text{H}_2\text{O}$  abundances and an isotopically light H isotope composition is unique among available phosphate SIMS standards (cf. McCubbin et al. 2015a, and references therein). Given that many of the SIMS analyses of merrillite in planetary samples have been conducted using apatite standards (McCubbin et al. 2014; Mane et al. 2016), a whitlockite standard will provide a better matrix match for future SIMS studies of merrillite or whitlockite in Earth and planetary materials.

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## Endnote:

<sup>1</sup> Structure Probe, Inc. (1996) Taylor multi element standard documentation. C.M. Taylor Company. <https://www.2spt.com/catalog/standards/aweb/tayl-min.html>