# Parasibirskite, a new mineral from Fuka, Okayama Prefecture, Japan

# I. KUSACHI, Y. TAKECHI

Department of Earth Sciences, Faculty of Education, Okayama University, Okayama, 700-8530, Japan

C. Henmi

Department of Earth Sciences, Faculty of Science, Okayama University, Okayama, 700-8530, Japan

AND

# S. KOBAYASHI

Division of Earth Sciences, Kurashiki University of Science and the Arts, Kurashiki, 712-8505, Japan

# ABSTRACT

Parasibirskite, with the ideal formula  $Ca_2B_2O_5 \cdot H_2O$ , is a new mineral species found at Fuka, Okayama Prefecture, Japan. It is a polymorph of sibirskite, CaHBO<sub>3</sub>, and occurs as subparallel aggregates of tabular crystals up to  $40 \times 20 \times 3 \mu m$  in size. Associated minerals are takedaite, olshanskyite, sibirskite, frolovite and calcite. The mineral is white, and has a weak pearly luster. Optically, the mineral is biaxial positive,  $\alpha$  1.556(2),  $\beta$  1.593(2),  $\gamma$  1.663(2) ( $\lambda$  589 nm). The Vickers microhardness of aggregates is 121 kg mm<sup>-2</sup>. The mineral is monoclinic with space group of  $P2_1/m$ , a 6.722(4), b 5.437(2), c 3.555(2) Å,  $\beta$  93.00(5)°, V 129.8(2) Å<sup>3</sup>. The strongest lines in the X-ray powder pattern [d in Å (I)(hkl)] are 2.237(100)(300), 6.73(70)(100), 2.975(60)(011), 3.354(30)(200), 2.855(20)(210) and 1.776(20) (002). Wet chemical analysis, electron-microprobe analysis and ICP emission spectrometry give the values CaO 56.06 %, B<sub>2</sub>O<sub>3</sub> 34.10 %, H<sub>2</sub>O 9.97 % and total 100.13%. The empirical formula calculated on the basis of O = 6 is Ca<sub>1.985</sub>B<sub>1.945</sub>O<sub>4.901</sub>·1.099H<sub>2</sub>O, for Z = 1, D<sub>calc</sub> 2.54 and D<sub>meas</sub> 2.50(1) g cm<sup>-3</sup>. Parasibirskite is formed by hydrothermal alteration of takedaite.

KEYWORDS: parasibirskite, sibirskite, gehlenite-spurrite skarns, borate, Fuka, Japan.

# Introduction

DURING a mineralogical survey of the gehlenitespurrite skarns at Fuka, Okayama Prefecture, Japan, Kusachi *et al.* (1997) reported the occurrence of an unidentified hydrous borate mineral associated with sibirskite and frolovite. Subsequent study has shown the mineral to be a natural analogue of synthetic  $2CaO \cdot B_2O_3 \cdot H_2O$  reported by Schäfer (1968). The new mineral has been named parasibirskite as a polymorph of sibirskite, CaHBO<sub>3</sub>. The mineral species and its name have been approved by the Commission on New Minerals and Mineral Names, International Mineralogical Association. The type mineral of parasibirskite is deposited at the National Science Museum, Tokyo, Japan. The present paper deals with its mode of occurrence and mineralogical properties.

# Occurrence

At Fuka, Okayama Prefecture, primary skarn minerals such as gehlenite and spurrite were formed as pyrometasomatic products of limestone. In subsequent retrograde hydration reactions, most of the gehlenite crystals were altered to retrograde minerals such as vesuvianite, hydrogrossular and bicchulite (Henmi *et al.*, 1973). Retrograde changes of spurrite crystals, on the other hand, are not extensive.

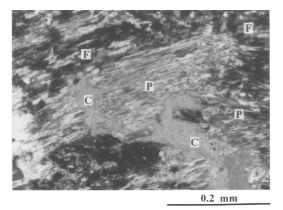


FIG. 1 Photomicrograph of parasibirskite. Crossed nicols. Abbreviations: P = parasibirskite, F = froloviteand C = calcite.

During the post-metasomatic stage, an irregular vein consisting of borate minerals penetrated along the boundary between crystalline limestone and the skarns. In the vein, an anhydrous borate, takedaite (Kusachi *et al.*, 1995*a*), was primarily formed by the reaction of boron-bearing fluids with limestone. At the circumference of the vein, hydrous borates such as nifontovite, olshanskyite (Kusachi and Henmi, 1994), frolovite (Kusachi *et al.*, 1995*b*) and sibirskite (Kusachi *et al.*, 1997) were secondarily formed by late-hydrothermal alteration of takedaite.

Parasibirskite was found in the altered part of takedaite, and occurred as subparallel aggregates of tabular crystals up to 40  $\mu$ m long, 20  $\mu$ m wide and 3  $\mu$ m thickness, in association with takedaite, olshanskyite, frolovite, sibirskite and calcite. The occurrence of parasibirskite from Fuka is shown in Fig. 1. It is likely that parasibirskite was formed by hydrothermal alteration of takedaite.

#### **Physical and optical properties**

Parasibirskite was white with a weak pearly lustre in hand specimen, and colourless in thin-section. The cleavage is perfect on (100). Optically, the mineral was biaxial positive with refractive indices

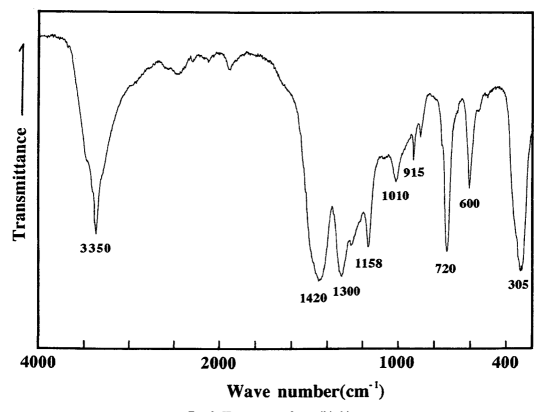


FIG. 2. IR spectrum of parasibirskite.

#### PARASIBIRSKITE, A NEW MINERAL

1				2		
Ī	d <sub>(meas)</sub>	d <sub>(calc)</sub>	h k l	d	Ι	
70	6.73	6.7128	100	6.71	70	
10	4.23	4.2250	1 1 0	4.23	10	
		3.5501	0 0 1	3.55	<5	
30	3.354	3.3564	2 0 0	3.36	30	
		3.0725	101	3.076	5	
60	2.975	2.9727	0 1 1	2.973	60	
20	2.855	2.8560	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	2.857	20	
		2.7632	$\bar{1}$ 1 1	2.760	<5	
10	2.721	2.7185	0 2 0	2.715	10	
		2.6749	1 1 1	2.676	<5	
		2.3776	2 0 1	2.378	<5	
100	2.237	2.2376	300	2.237	100	
		2.1784	2 1 1	2.181	<5	
		2.1584	021	2.158	<5	
		2.1125	2 2 0	2.112	<5	
10	2.075	2.0742	$   \begin{array}{cccc}     2 & 2 & 0 \\     \overline{1} & 2 & 1 \\   \end{array} $			
			}	2.071	10	
		2.0692	310			
10	2.036	2.0360	1 2 1	2.035	10	
		1.9393	301	1.936	<5	
5	1.842	1.8424	Ž 2 1	1.840	5	
10	1.826	1.8267	$     \begin{array}{r}       \bar{3} & 0 & 1 \\       \bar{2} & 2 & 1 \\       \bar{3} & 1 & 1     \end{array} $	1.825	10	
10	1.790	1.7897	2 2 1	1.790	10	
20	1.776	1.7752	0 0 2	1.776	20	
		1.7512	3 1 1	1.752	5	
5	1.750	1.7497	1 3 0	1.749	5 5	
a (Å) =	6.722(4)			6.7	6.719(2)	
$b(\mathbf{A}) =$	5.437(2)				5.434(3)	
c(Å) =	3.555(2)				3.555(1)	
$\beta(°) =$	93.00(5)				92.87(4)	

#### TABLE 1. X-ray powder data for parasibirskite

1. Parasibirskite from Fuka, Okayama Prefecture, Japan. The present work.

2. Synthetic Ca<sub>2</sub>B<sub>2</sub>O<sub>5</sub>·H<sub>2</sub>O (Schäfer, 1968)

 $\alpha$  1.556(2),  $\beta$  1.593(2),  $\gamma$  1.663(2), 2V 74.8° (calc.). 2V could not be measured due to the thinness of crystals. The Vickers microhardness of aggregates was 121(109–155) kg mm<sup>-2</sup> (25 g load). The measured density by heavy liquids was 2.50(1) g cm<sup>-3</sup>, the calculated density being 2.54 g cm<sup>-3</sup>. The measured density was lower than that of synthetic 2CaO·B<sub>2</sub>O<sub>3</sub>· H<sub>2</sub>O reported by Schäfer (1968), which was found to be 2.60 g cm<sup>-3</sup>.

The infrared absorption spectrum of parasibirskite was measured by the KBr method for the region 4000 to 250 cm<sup>-1</sup>, as shown in Fig. 2. The absorption band at 3350 cm<sup>-1</sup> is attributed to the OH stretching vibration. Numerous bands observed at 1420 to 305  $\text{cm}^{-1}$  are characteristic of borates.

# **Chemical composition**

Chemical analysis using an electron-microprobe analyser revealed the presence of Ca, B and O, and the absence of any other element. The concentration of Ca was obtained by an electron-microprobe analyser, the B content was determined by ICP, and the H<sub>2</sub>O content was determined from ignition loss and TGA at 900°C. Chemical analysis gave the values CaO 56.06%, B<sub>2</sub>O<sub>3</sub> 34.10%, H<sub>2</sub>O 9.97% and total 100.13%. The empirical formula calculated on the basis of O = 6is therefore  $Ca_{1.985}B_{1.945}O_{4.901} \cdot 1.099H_2O$  which gives the ideal formula  $Ca_2B_2O_5 \cdot H_2O$ . Parasibirskite easily alters to calcium borate hydrate ( $CaB_2O_4 \cdot 6H_2O$ ) and calcite in water, and is also easily soluble in dilute hydrochloric acid with no effervescence.

# X-ray studies

The X-ray powder data for parasibirskite were obtained by an X-ray diffractometer using Ni-filtered Cu- $K\alpha_1$  radiation. Single crystal studies could not be carried out because of the small crystal size. Reflections of parasibirskite from Fuka were indexed on the monoclinic cell, *a* 6.719, *b* 5.434, *c* 3.555 Å and  $\beta$  92.87°, reported for synthetic 2CaO-B<sub>2</sub>O<sub>3</sub>·H<sub>2</sub>O (Schäfer, 1968). Unit cell dimensions, refined by least squares from the present powder diffraction data for parasibirskite, were *a* 6.722(4), *b* 5.437(2), *c* 3.555(2) Å and  $\beta$  93.00(5)°. The X-ray powder data are given in Table 1, and compared with those of synthetic 2CaO-B<sub>2</sub>O<sub>3</sub>·H<sub>2</sub>O reported by Schäfer.

### **Thermal behaviour**

DTA and TG curves were obtained by heating parasibirskite from room temperature to 900°C at a rate of 10°C min<sup>-1</sup>, as shown in Fig. 3. The DTA curve has endothermic peaks at 371 and 414°C corresponding to the loss of water. An exothermic peak was not observed. On heating the mineral at 900°C, calcium borate (Ca<sub>2</sub>B<sub>2</sub>O<sub>5</sub>) appeared as a crystalline phase.

# Discussion

Schäfer(1968) synthesized single crystals of  $2CaO \cdot B_2O_3 \cdot H_2O$  from  $Ca(OH)_2$  and  $H_3BO_3$  at temperatures between 100 and 300°C under saturated water pressure. The chemical composition and X-ray powder data for parasibirskite from Fuka agree well with those for the synthetic  $2CaO \cdot B_2O_3 \cdot H_2O$ . Therefore, parasibirskite from Fuka is a natural analogue of the synthetic  $2CaO \cdot B_2O_3 \cdot H_2O$ . 2CaO  $\cdot B_2O_3 \cdot H_2O$  in an oxide form may be written as  $Ca_2B_2O_3 \cdot H_2O$  and can also be expressed as  $2(CaHBO_3)$ . The mineral

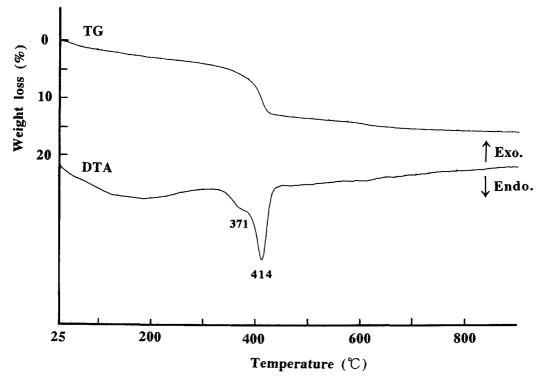


FIG. 3. DTA and TG curves of parasibirskite.

with the chemical formula of CaHBO3 is known as sibirskite. Kusachi et al. (1997) reported that chemical data of sibirskite from Fuka are consistent with the ideal formula of CaHBO<sub>3</sub>. The chemical composition of parasibirskite agrees with that of sibirskite from Fuka, and their infrared absorption spectrum and thermal behaviour are similar. X-ray powder data for parasibirskite, however, are different from those for sibirskite from Fuka. The unit cell parameters for parasibirskite are a 6.722(4), b 5.437(2), c 3.555(2) Å and  $\beta$  93.00(5)° although those for sibirskite from Fuka are a 8.643(6), b 9.523(2), c 3.567(3) Å and  $\beta$  119.23(3)° (Kusachi et al., 1997). From these data, it is considered that there is a polymorphic relationship between parasibirskite and sibirskite.

#### Acknowledgements

We would like thank Mrs M. Shimada and S. Kishi for field work conducted, and Mr L. E. Anthony of Okayama University of Science for proofreading the final manuscript. We also wish to thank the Research Instruments Center of Okayama University of Science for the use of their facilities. This work was supported in part by a Grant-in-Aid for Scientific Research from the

Ministry of Education, Science, Sports and Culture, Japan.

# References

- Henmi, C., Kusachi, I., Henmi, K., Sabine, P. A. and Young, B. R. (1973) A new mineral bicchulite, the natural analogue of gehlenite hydrate, from Fuka, Okayama Prefecture, Japan and Carneal, County Antrim, Northern Ireland. *Mineral. J.*, 7, 243-51.
- Kusachi, I. and Henmi, C. (1994) Nifontovite and olshanskyite from Fuka, Okayama Prefecture, Japan. *Mineral. Mag.*, 58, 279-84.
- Kusachi, I., Henmi, C. and Kobayashi, S. (1995*a*) Takedaite, a new mineral from Fuka, Okayama Prefecture, Japan. *Mineral. Mag.*, **59**, 549–52.
- Kusachi, I., Henmi, C. and Kobayashi, S. (1995b) Frolovite from Fuka, Okayama Prefecture, Japan. *Mineral. J.*, 17, 330-7.
- Kusachi, I., Henmi, C. and Kobayashi, S. (1997) Sibirskite from Fuka, Okayama Prefecture, Japan. *Mineral. J.*, **19**, 109–14.
- Schäfer, U. L. (1968) Synthese und röntgenographische Untersuchung der Borate 3CaO·B<sub>2</sub>O<sub>3</sub>, 2CaO·B<sub>2</sub>O<sub>3</sub> und 2CaO·B<sub>2</sub>O<sub>3</sub>·H<sub>2</sub>O. *Neues Jahrb. Mineral. Mh.*, 75-80.
- [Manuscript received 29 September 1997]