Si-DEFICIENT, OH-SUBSTITUTED, BORON-BEARING VESUVIANITE 
FROM THE WILUY RIVER, YAKUTIA, RUSSIA

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ABSTRACT

A low-temperature, Si-deficient variety of vesuvianite occurs in porous tetrahedral “achtarandite” pseudomorphs consisting of hibschite, along the banks of the Wiluy River, Yakutia, Russia, the type locality of grossular and wiluite. The (H₄O₄)₄⁻-for-(SiO₄)⁴⁺ hydrogarnet-type substitution is evident in the vesuvianite, a substitution that allows it to be considered an analogue of hibschite. This variety of vesuvianite belongs to a new series in the vesuvianite group, as expressed by the formula 

\[(X^{19}Y^{13}T^{0–5})(Si₂O₇)(SiO₄)₁₀–x(OH)₄W_{10}.\]

The filling of the \(X\), \(Y\), and \(T\) positions in this Si-deficient vesuvianite, where \(x\) varies from 0.67 to 2.89, is analogous to that in vesuvianite and wiluite. The Si-deficient vesuvianite is characterized by increased unit-cell parameters, \(a = 15.688(3)\) Å, \(c = 11.860(3)\) Å and by lower indices of refraction, \(n_H = 1.691(1)\), \(n_K = 1.668(1)\). In the OH-region, the FTIR and Raman spectra differ sharply from those of low-temperature vesuvianite from rodingites, but are similar to the spectra of hibschite. A line near 3620 cm⁻¹ indicates that the substitution occurs only in the isolated tetrahedra. More than 25% of these can be substituted by (H₄O₄). Contents of boron up to 2.48 apfu were detected in the Si-deficient vesuvianite. The vesuvianite formed during the hydration (serpentinization and rodingitization) of early, high-temperature skarns.

Keywords: Si-deficient vesuvianite, hydrogarnet-type substitution, boron, wiluite, infrared spectra, Raman spectra, electron-microprobe data, unit-cell parameters, Wiluy River, Russia.

SOMMAIRE

Une variété de vésuvianite déficitaire en Si, formée à basse température, fait partie d’un amas tétraédrique de hibschite formé par pseudomorphose (“achtarandite”) le long des rives de la rivière Wiluy, en Yakoutie, Russie, la localité-type du grossulaire et de la wiluite. La substitution de (H₄O₄)₄⁻ au (SiO₄)⁴⁺, comme c’est le cas dans un hydrogénat, se manifeste donc dans la

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vésuvianite, et même à ce que l’on peut considérer comme un analogue de la hibschite. Cette variété de vésuvianite fait partie d’une nouvelle série du groupe de la vésuvianite, comme l’exprime la formule \(X_{10}Y_{1}Z_{0}T_{0}(Z_{2}O_{3})(\text{ZnO})_{10}W_{10}\). Le schéma utilisé pour remplir les sites X, Y, et T dans cette vésuvianite déficitaire en Si, dans laquelle la x varie de 0.67 à 2.89, est analogue à celui qui régit la vésuvianite et la wiluite. La vésuvianite déficitaire en Si fait preuve d’une augmentation en paramètres réticulaires, à 15.688(3), \(\alpha = 11.860(3)\) \(\AA\), et d’une diminution des indices de réfraction, \(\varepsilon = 1.691(1), \omega = 1.668(1)\). Dans la région où se trouvent les bandes OH, les spectres d’absorption infrarouge (avec transformation de Fourier) et de Raman diffèrent de façon marquée de ceux de la vésuvianite de basse température provenant des rodingites, mais ressemble à ceux de la hibschite. Une bande située près de 3620 cm\(^{-1}\) indiquerait que la substitution n’implique que les tétraèdres isolés. Plus de 25\% de ceux-ci semblent remplacés par des agencements (H\(_{4}\)O\(_{4}\)). Des teneurs en bore atteignant jusqu’à 2.48 atomes par unité formaulaire sont signalées dans la vésuvianite déficitaire en Si. Cet exemple se serait formé lors d’une déshydratation (serpentinitisation et rodingitisation) d’un skarn de haute température.

**Mots-clés:** vésuvianite déficitaire en Si, substitution de type hydrogarenat, bore, wiluite, spectres infrarouges, spectres de Raman, données de microsonde électronique, paramètres réticulaires, rivière Wiluy, Russie.

**Introduction**

The common simplified formula of vesuvianite-group minerals is \(X_{10}Y_{1}Z_{0}T_{0}(Z_{2}O_{3})(\text{ZnO})_{10}W_{10}\). The X position is usually occupied by Ca, Y by Al, Mg, Fe\(^{2+}\), Fe\(^{3+}\), Mn\(^{2+}\), and Ti, T by B and Al, Z by Si, and W by O, OH, F, and Cl. The boron-dominant structural analogue of vesuvianite, wiluite, has more than 2.5 B atoms per formula unit \((\text{apfu})\) at the T position (Goat et al. 1998). The similarity of both vesuvianite and grossular in terms of structure and composition (Allen & Burnham 1992), as well as the common association of vesuvianite with hydrogarnet-type substitution in garnet-group minerals is Indicated by the formula \((\text{H}_{4}\text{O}_{4})_{4}^-\) for \((\text{SiO}_{4})_{4}^-\) hydrogarnet-type substitution in vésuvianite. However, numerous investigations of vesuvianite from various environments have failed to provide any support for a hydrogarnet-type substitution (Goat et al. 1992, Fitzgerald et al. 1992, Lager et al. 1999).

Recently, Armbruster & Gnos (2000) discovered vacancies at the tetrahedral sites in low-temperature Mn-enriched vesuvianite from South Africa, which points to the possibility of a \((\text{H}_{4}\text{O}_{4})_{4}^-\) for \((\text{SiO}_{4})_{4}^+\) substitution. Contents of Si in these samples fluctuate slightly below the ideal 18 \text{apfu}, which indicates a low degree of substitution of the Si tetrahedra.

During our investigation of the vesuvianite–wiluite series of minerals from the Wiluy occurrence, along the banks of the Wiluy River, in Yakutia, split crystals and spherulitic arrays, some as large as 200 μm, were discovered within “achtarandite”-sponge pseudomorphs of hibschite after a wadalite-like phase (Galuskin et al. 1996, Galuskin & Galuskina 2002). These were found to be Si-deficient vesuvianite displaying an apparently low degree of substitution of \((\text{SiO}_{4})_{4}^+\) by \((\text{H}_{4}\text{O}_{4})_{4}^-\) (Galuskin et al. 2002). The vesuvianite-like phase forming a tight intergrowth with hydrogarnet in the alteration products of gehlenite probably also consist of Si-deficient vesuvianite. However, such material has not been investigated sufficiently thoroughly to draw a definitive conclusion (Henmi et al. 1994).

**Methods of Investigation**

The optical properties of the vesuvianite were investigated in thin sections and immersion preparations with the aid of a polarizing microscope. The morphology of the vesuvianite was studied using electron microscopes JSM–35C [(high vacuum (HV)] and FEI/Philips XL30 with EDS (EDAX) [HV and low vacuum (LV)]. Secondary electron [SE (HV)] and back-scattered electron [BSE (HV and LV, 0.3 Torr)] detectors were used to obtain images of the vesuvianite. During work in the LV regime, non-coated samples were used.

Electron-microprobe analyses were made using a CAMECA SX–100 instrument (Warsaw), taking into account the recommendations of McGee & Anovitz (1996) concerning the monitoring of boron. Measurements of the main components were performed at 15 kV and 20 nA for 20 seconds, using natural standards. Concentrations of F and B were measured at 5 kV and 100 nA for 50–100 seconds at each point. In measuring the concentration of boron, a danburite standard with control measurements from external standards (danburite, wiluite and boron-free marialite) was used.

The unit-cell parameters were calculated using the X’Pert Plus program on the basis of a powder X-ray pattern obtained using a Philips PW3710 diffractometer under the following conditions: 40 kV, 30 mA, 0.02° step, 35 seconds measurement-time per point. Following the standard method, the infrared spectrum of vesuvianite was also investigated, in KBr pellets at room temperature (resolution 4 cm\(^{-1}\)) and at 20 K (resolution 8 cm\(^{-1}\)) using a FTIR Digilab 60V (BioRad) spectrometer.

A sample of Si-deficient vesuvianite was analyzed on a Dilor Micro Raman spectrometer with a 514.5 nm argon ion laser. With this instrument, one focuses the size of the laser beam to about 5 μm on the surface of the sample. Raman spectra of vesuvianite-group phases from rodingites from the Urales, here given for comparison, have been collected using a FTS 6000 Bio-Rad spectrometer with Raman section (with Nd:YAG Spec-
tra Physics T10 106 4c laser) with a resolution of 4 cm⁻¹ and 30,000 scans. The laser power was maintained at 200 mW on the sample.

**MORPHOLOGY AND COMPOSITION OF SI-DEFICIENT VESUVIANITE**

Light-cream spherulites and split crystals of Si-deficient vesuvianite with a characteristic nacreous luster were found inside “achtarandite” pseudomorphs (Fig. 1). These define compact intergrowths with a chlorite-group mineral and hibschite (Fig. 2). Sheaf-like crystals of vesuvianite are bounded by the {100} and {101} faces (Figs. 1a, b). The {001} pinacoid faces, as main crystallographic form, appear on subindividuals of spherulites (Fig. 1c). The spherulites consist of flat subindividuals elongate parallel to the Z axis (Figs. 1d, 2) and possess an increased porosity. Both morphological forms of the Si-deficient vesuvianite developed as a result of a splitting of the {100} prism faces by an “autodeformation mechanism” connected with sectoral heterometry (Punin 2000, Galuskin et al. 2001).

The composition of the Si-deficient vesuvianite is close to that of vesuvianite from rodingites (e.g., Groat et al. 1992, Fitzgerald et al. 1992). Si-deficient vesuvianite is characterized by a low content of Mg, Ti and Fe and a higher level of Al. This composition contrasts with that of the high-temperature wiluite associated with Si-deficient vesuvianite (Table 1, Fig. 3). There is a considerable variation in the contents of cations incorporated in the Y position, with a tendency for increasing Fe contents toward the margin of the spherulites (Fig. 3, Table 1).

Several zones, with a homogeneous structure in which aluminum is enriched, are significant. In these, the sum of the cations at the Y position is slightly more than 13 apfu (Fig. 2c, Table 1, anal. 3). This excess serves as evidence of limited replacement of Al for Si.

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**Fig. 1.** Morphology of Si-deficient vesuvianite, SEM: a Split{100}+{101} crystal on the {111}+{110} crystal of hibschite, SE. b Sheaf-like {100}+{101} crystals, SE. c Spherulites terminated by the {001}, {100}, {101} faces, BSE (LV). d Radial structure of spherulite, BSE (LV).
and of occupancy of a small part of the Al at the $T(1)$ and $T(2)$ positions (Groat et al. 1994a). In general, the sum of cations incorporated at the $Y$ position in vesuvianite is close to 13 apfu (Table 1), which indicates that the Al-for-Si substitution does not play a significant role.

The principal characteristic of the Wiluy River vesuvianite, which sets it apart from all other known samples of vesuvianite, is its low content of Si, near 15–16 apfu (Table 1, Fig. 4a). This feature demonstrates the considerable role of the $(O_4H_4)^{4–}$ for $(SiO_4)^{4–}$ substitution in Si-deficient vesuvianite from the Wiluy River, in analogy with hydrogrossular (Rinaldi & Passaglia 1989). In addition, thin late zones with increased contents of OH groups that exceed maximum possible contents of the OH groups at the $W$ positions, i.e., 9 apfu, are noted on crystals of wiluite and boron-rich vesuvianite (Fig. 4a, Galuskin et al., in prep.).

No correlation is observed between amounts of Si and OH in wiluite with low contents of OH groups (Fig. 4a), but a negative correlation between B and OH occupying incompatible positions in the wiluite structure is plainly visible (Fig. 4b). The average content of boron in Si-deficient vesuvianite is lower than that in wiluite ($B > 2.5$ apfu) and usually does not exceed 2 apfu of B although, in rare cases, the content of boron exceeds 2.48 apfu (Fig. 4b, Table 1). It is likely that boron in Si-deficient vesuvianite occupies the $T(1)$ and $T(2)$ positions in the structure, as in boron-bearing vesuvianite and wiluite (Groat et al. 1994b, 1996, 1998). There is a negative correlation between amounts of Si and OH (Fig. 4a) in Si-deficient vesuvianite, with numbers of

Fig. 2. Cross-section of split forms of Si-deficient vesuvianite. a–c Spherulite: transmitted light at parallel nicols (a) and crossed nicols with gypsum compensator (b) and BSE image (c), with sites of electron-microprobe analyses indicated (Table 1). d Sheaf-like crystals, BSE image with marked points of analyses (Table 1). Chl chlorite, Hb hibschite.
OH groups exceeding maximum possible numbers at the $W$ position (taking in account isomorphism with other anions), which corroborates the occurrence of isomorphic (O$_2$H$_2$)$^-$ for (SiO$_2$)$^2^+$ substitution, as in hydrogarnet. There is a common tendency for increasing Si contents toward the margins of aggregates of Si-deficient vesuvianite (Table 1). Similar degrees of hydration of hibschite and Si-deficient vesuvianite are evident where they occur in association (Table 1, Galuskina et al. 2001).

Minerals of the hibschite–katoite series have increased unit-cell parameters, which reflect increasing $Z$–O and Ca–O interatomic distances (Lager et al. 1989). Hydrogarnet-group minerals also have lower indices of refraction than grossular. They are characterized by the appearance of bands near 3600–3620 and 3660 cm$^{-1}$ in the region of OH-stretching vibrations, which is indicative of the replacement of part of the Si in the tetrahedra by H (Zabiński 1965, Passaglia & Rinaldi 1984, Rinaldi & Passaglia 1989, Rossman & Aines 1991, Galuskina et al. 2001).

The Si-deficient vesuvianite from the Wiluy River is differentiated from typical vesuvianite by increased unit-cell parameters: $a$ 15.688(3), $c$ 11.860(3) Å. In the vesuvianite group, only in metamict vesuvianite do both parameters increase, whereas wiluite is characterized by an increased $a$ parameter and a decreased $c$ parameter (Groat et al. 1992).

The selection of a sample for single-crystal X-ray investigation was a major problem because all crystals have a domain and fibrous structure and a high porosity. Only in one case did we obtain diffraction data for a
TABLE 1. CHEMICAL COMPOSITION (WT. %) AND DEGREE OF HYDRATION (K, %) OF SI-DEFICIENT VESUVIANITE, WILUITE AND HIBSCHITE

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1-6 - {100}+{001}+{101} spherulites of Si-deficient vesuvianite (Fig.2c), 7-11 {100}+{101} shaf-like split crystal of Si-deficient vesuvianite (Fig. 2d), 12 - {110} hibschite (fig.2d), 13- late zone of {100} growth sector of wiluite crystal.

* - total iron as Fe₂O₃, ** - water calculated by valence balance, * - in total 0.02 Cr₂O₃, ** - Fe³⁺+Cr³⁺, *** - calculated on charge 144-F and normalized on 18 Ca, n.a. - not analyzed, n.d. - not detected, K - % of vacancies in isolated Z(1,2) tetrahedra [=degree of hydration of SiO₄ tetrahedra (Rinaldi & Passaglia 1989)].
vesuvianite crystal of pinacoidal type (0.22 × 0.22 × 0.15 mm) with the crystal-chemical formula Ca19Al9.02 Mg3.22Fe2+0.58Mn2+0.16Ti3+0.02Sr13(B2O3.81Al3.09)20.9
[(SiO4)28.05(H2O)31.95Si2O7]4[Si2O7]4(OH)7.18O2.37F0.36C0.06]10 (Galuskin et al., in prep.). A structure refinement of this vesuvianite (P4/mnc, R1 = 6.36%) reveals a significant number of vacancies (~15%) in isolated Si-
tetrahedra [Z(1,2)] and an increase in Z(1)–O atomic
distances, up to 1.686 Å (Galuskin et al., in prep.). This finding confirms our inference concerning a hydro-
garnet-type substitution in Si-deficient vesuvianite.

Lower indices of refraction, ε = 1.691(1), ω = 1.668(1), are typical of Si-deficient vesuvianite (for more ferriferous varieties, see Table 1). The optic sign is variable, depending on which growth sectors of the vesuvianite faces participate in the formation of spherulites (Fig. 2b; Galuskin et al. 2001). The angle 2V could not be measured in the central part of spherulites because of the thin fibrous structure (Figs. 1d, 2a). A bi-
axial character is not evident in the thin optically
negative part at the margin of the spherulites.

FTIR spectra of the OH-region of the Si-deficient vesuvianite differ sharply from spectra typical of low-
temperature vesuvianite and display a similarity to the spectra of hibschite (Fig. 5). At room temperature, the position of absorption bands is not successfully defined on the spectra of the Si-deficient vesuvianite. Only a lower-temperature measurement at 20 K, following decomposition of the spectra using the method proposed by Handke et al. (1994), allows for the demarcation of a few absorption bands: 3618, 3534, 3442, 3351 and
at the O(11) position or of B at the T(1) position. The 3516 cm−1 band in the vesuvianite spectrum, and also a weak band near the 3534 cm−1 band in the spectrum of Si-deficient vesuvianite, are probably defined by the Fe2+–Al–OH configuration (Groat et al. 1995). The positions and shapes of bands near 3442 (Si-deficient ve-
suvianite), 3431 (hibschite) and 3423 cm−1 (vesuvianite) indicate the presence of molecules of H2O (Fig. 5).

Vesuvianite from the rodingite shows a pronounced band near 3155 cm−1 (Fig. 5, spectrum 4); it is character-
istic of low-temperature, ordered vesuvianite, with a symmetry lower than P4/mnc, and is defined by the local O(10)H(2)− ... O(10)O2− configuration (Zabiński & Paluszkiewicz 1994, Groat et al. 1995). Si-deficient vesuvianite does not possess an analogous band, but we do see very weak broad bands higher than 3200 cm−1, which may indicate a low degree of order in the vicinity of the O(10) position. Boron atoms at the T(2) position, and F and Cl at the O(10) position, will influence the character of spectra and lead to reduction of the intensity of the OH-band.

Sharp distinctions between vesuvianite from the Urals rodingites and the Si-deficient vesuvianite from Wiluy River are observed in the OH region on Raman spectra (Fig. 6). The intense band near 3622 cm−1 is analogous to the 3618 cm−1 band on the FTIR spectra (Fig. 5); it is noted in spectra of hydrogrossular and
Wiluy River are observed in the OH region on Raman spectra (Fig. 6). The intense band near 3622 cm−1 is analogous to the 3618 cm−1 band on the FTIR spectra (Fig. 5); it is noted in spectra of hydrogrossular and points to a hydrogarnet type of substitution in the isolated tetrahedra (Arredondo & Rossman 2002). This band is absent in the spectra of vesuvianite from the rodingite, but there is a broad band near 3150–3200 cm−1 that appears to be analogous to a similar band defined on FTIR spectra that is connected with the O(10)H(2)− ... O(10)O2− configuration.

The crystal-chemical formula of highly OH substi-
tuted, Si-deficient vesuvianite can be presented as X19Y13T0.25Si(SiO4)10(OH)4(OH,OH,F,Cl)10, where the occupants of the X, Y, and T positions are analogous to those in vesuvianite and wiluite, and 0.67 < x < 2.89 for Wiluy Si-deficient vesuvianite. In this series, only a phase with x > 5 could be considered as a new mineral species ("hydrovesuvianite"), according to the rule of C N M N MIMA (Nickel & Grice 1998). It is interesting that some compositions of Si-deficient ve-
suvianite are very close to the field of wiluite (Fig. 4, Groat et al. 1998), which may indicate the presence in nature of Si-deficient wiluite ("hydrowiluite").

A predominance of oxygen at the W position, for example Ca19Al9.22(Mg3.02Fe2+0.75Mn2+0.06Ti0.03)21.03
B2.06(Si2O7)4Si(SiO4)28.13(OH)7.36Al22OH2.42F0.93
C0.30Z10 (anal. 7, Table 1), is noted in Si-deficient ve-
suvianite with high contents of boron. A predominance of oxygen at the W position is characteristic of wiluite, but not of typical vesuvianite from various environments (Fitzgerald et al. 1992, Groat et al. 1992, 1998).

The formation of Si-deficient vesuvianite was a result of the hydration (serpentinization and rodingitiza-
tion) of early high-temperature layered skarns
associated with magmatic rocks of Siberian Trap Formation on the Wiluy River (Galuskina et al. 1998, 2001). Crystallization of Si-deficient vesuvianite and of hibschite progressed under non-equilibrium conditions, a fact reflected in their morphology. The conditions under which crystallization occurred are characterized by high Al/Si and Ca/Si in the mineral-forming solutions, high $P(H_2O)$ and 300 < $T$ < 350°C (Galuskina et al. 2001).

ACKNOWLEDGEMENTS

We thank Prof. J.A. Mandarino, Prof. D.Yu. Pushcharovsky and Dr. J.D. Grice for discussions connected with the problems of vesuvianite-group nomenclature. We also thank Dr. P. Dzierżanowski (University of Warsaw) for technical assistance with the electron-microprobe analyses. We also thank Dr. Padhraig Kennan and Mr. David Gordon for their comments regarding English usage, and Dr. M. Gunter for editorial work. We thank Prof. T. Armbruster, Prof. L.A. Groat, an anonymous referee, and R.F. Martin for their constructive comments on this paper. Financial support for this project was provided by the Scientific Research Fund of the University of Silesia under grant BW–22/2000 and “KBN” (Polish Committee for Research) grant 3PO4D 03622.

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Received December 2, 2002, revised manuscript accepted July 13, 2003.