Discovery of datolite in a high-pressure marble from the Sanbagawa metamorphic belt: Indication of B-rich fluid activity

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Datolite [CaBSiO4(OH)] was discovered in an eclogite–facies calcite marble collected from the Eastern Iratsu body in the Sanbagawa metamorphic belt of central Shikoku. The marble was composed of calcite, diopside, and garnet that contained inclusions of omphacite. Enclosed in the marble is a pod composed mainly of quartz, with subordinate calcite, diopside, and garnet that has inclusions of datolite. The formation conditions of the datolite were estimated on the basis of mineral assemblage and the Raman elastic geobarometer to be approximately 400–650 °C and 0.8–1.3 GPa, which coincide with the conditions of the eclogite juxtaposition with the non-eclogite units in the Besshi district. Our study records the highest pressure-temperature conditions as the metamorphic datolite formation. Our findings provide evidence for the occurrence of B-rich fluid infiltration during the juxtaposition of eclogite unit with the non-eclogite unit in the Besshi district.

Keywords: Datolite, Sanbagawa metamorphic belt, Deep fluid, B-rich fluid, Borosilicate

INTRODUCTION

Datolite [CaBSiO4(OH)] is a hydrous borosilicate that is commonly formed through hydrothermal, volcanic, or low-grade metamorphic processes (Grew, 1996; Kiss et al., 2012). Although the stability limit of datolite, with respect to pressure (P) and temperature (T), is wide as <700 °C at atmospheric pressure or <5 GPa at 250 °C (Tarney et al., 1973; Goryainov et al., 2015), datolite is sensitive to the composition of coexisting fluids, such as H⁺ activity (pH) and H3BO3 activity. On the basis of the experimental data, Yang and Rosenberg (1995) calculated the stable minerals in the system of CaO–B2O3–SiO2–H2O, indicating that datolite is stable at high pH at a given αCa₂⁺/αH⁺, where low αCa₂⁺/αH⁺ or high αSiO₂ conditions favor danburite (CaB₂Si₂O₇) instead of datolite.

This study reports the occurrence of datolite in a quartz pod containing grossular, calcite, and diopside enclosed in an eclogite–facies marble. Datolite was discovered as inclusions in andradite–enriched patches in grossular for which the formation conditions were well constrained based on mineral assemblage and Raman-based geobarometry. We report the highest P–T conditions for the formation of metamorphic datolite. We also discuss the significance of datolite as a possible measure of B-rich fluid activity during the exhumation of eclogite–facies rock in the metamorphic terrain.

GEOLOGICAL BACKGROUND

The Sanbagawa metamorphic belt is a Cretaceous subduction complex where four mineral zones have been identified based on the mineral assemblages of pelitic schists. The succession with increasing peak metamorphic T is (1) chlorite zone, which is equivalent to the prehnite–pumpellyite facies (300 °C, 0.5 GPa), (2) garnet zone, (3) albite–biotite zone, and (4) oligoclase–biotite zone, which is equivalent to the epidote–amphibolite facies (610 °C, 1.0 GPa) (Higashino, 1990; Enami, 1994; doi:10.2465/jmps.200730 K. Yoshida, yoshida_ken@jamstec.go.jp Corresponding author
Aoya et al., 2013). In central Shikoku, eclogite–facies rocks, showing distinct high–pressure conditions, are widely distributed among the highest-grade parts of the albite– and oligoclase–biotite zones (Takasu, 1984; Aoya et al., 2013; Kouketsu et al., 2014a).

The Eastern Iratsu body is mainly composed of metamorphosed layered gabbro that has been subjected to the early stages of granulite–facies metamorphism and subsequent eclogite–facies metamorphism. This body is bordered by the albite–biotite zone to the east (Fig. 1; Takasu and Kohsaka, 1987). The datolite-bearing sample, EI1901, was obtained from the eastern margin area of the Eastern Iratsu body (EI), Higashi-Akaishi peridotite body (HA), Tonaru body (TN), and Seba body (SB).

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**ANALYTICAL PROCEDURE**

The chemical compositions of the constituent minerals were analyzed at Japan Agency for Marine–Earth Science and Technology (JAMSTEC, Yokosuka, Japan) using an electron microprobe (EMP) analyzer with five wavelength dispersive X-ray detectors (JXA–8500F, JEOL). Synthetic and natural standards were employed in the quantitative analysis. X-rays were obtained using an accelerating voltage of 15 kV and a beam current of 10 nA, except for datolite. For the analysis of datolite, a 5 nA beam current was used to avoid damaging the sample, and synthetic boron nitride was used as the standard for boron analysis. The representative EMP analyses of garnet and datolite are presented in Table 1. Raman microscopy was performed using a Raman spectrophotometer (RAMANtouch, Nanophoton) at JAMSTEC using a 532 nm green laser. Laser power was obtained at 2–10 mW on the sample surface.

**PETROGRAPHY AND MINERAL CHEMISTRY**

EI1901 was collected from a marble outcrop (more than 3 m in width) with many greenish mafic inclusions measuring several centimeters. Mafic rocks are recognized few meters away from the marble outcrop, although the contact between the two rock types is not visible owing to the surface soil. The studied sample is a weakly foliated marble containing several dark-colored spots of garnet (up to 5 mm) and greenish spots of diopside (<1 mm) (Fig. 2a). The matrix of the sample mainly consists of calcite, diopside, and garnet. There are two modes of occurrence of calcite: faceted fine grains, and elongated coarse and anhedral grains. The latter defines the main foliation. The garnet in the matrix is colorless under thin-section observation and subhedral, containing abundant inclusions of omphacite, epidote, quartz, phengite, aragonite, and graphite with K-feldspar and albite possibly resulting from decomposition reactions. The diopside in...
the matrix is round in shape and almost devoid of mineral inclusions. A quartz pod is elongated parallel to the main foliation and contains quartz, garnet, calcite, and diopside with trace amounts of pyrite and titanite (Fig. 2a). The garnet grains in the quartz pod are subhedral and up to 3 mm across. It is almost pure grossular (Grs) with minor amounts of andradite (Adr) and almandine (Alm) components as described below. The scanning electron microscope (SEM) showed that the garnet in the quartz pod featured isolated and fragmented Fe-poor cores with irregular outlines (Grt1) in a patchy matrix richer in Fe (Grt2) (Figs. 2b and 2c). These textures were caused by differences in the ferrie iron content: Grt1 and Grt2 comprise Grs89.99Adr1.1Alm1.6 and Grs88.92Adr1.1Alm1.6, respectively (Fig. 2e and Table 1). There is no systematic change of the compositional variation of Grt2. Grt1 contains inclusions of aragonite and minor amounts of K-feldspar, while Grt2 contains calcite and quartz.

Datolite is exclusively found as subhedral, unzoned inclusions in Grt2 (Fig. 2d). The datolite grain exhibits a weakly faceted shape that is devoid of chemical zonation (Fig. 2d). The EMP analysis showed compositions that were identical to the ideal formula of datolite, and the Raman spectra were consistent with those in the reference study (Table 1 and Fig. 3a, Lafuente et al., 2015). Datolite was neither found in garnet nor in diopside in the calcite marble hosting the quartz pod.

### FORMATION P-T CONDITIONS OF DATOLITE

The occurrence of datolite in EI1901 strongly suggests that the datolite was formed simultaneously with Grt2. The formation P-T conditions of Grt2 were estimated using a combination of quartz-Raman barometry and thermodynamic calculations. Quartz-Raman barometry is a pressure-estimation method that employs the frequency shift of quartz grains that are completely shielded in the garnet (Enami et al., 2007; Kouketsu et al., 2014b). The Raman shifts of quartz grains in Grt2 were measured, and \( \Delta \omega_{1} \) values (difference in the distance between two peaks of \( \nu_{464} \) and \( \nu_{205} \)) were calculated according to the procedure reported by Kouketsu et al. (2014b). Figure 3b shows the histogram of \( \Delta \omega_{1} \) obtained from Grt2, yielding approximately \( \Delta \omega_{1} = 4.5 \) for the most frequent value. Then, \( \Delta \omega_{1} \) values were evaluated using the P-T relationship reported by Kouketsu et al. (2014b) (Fig. 4b).

The temperature ranges were constrained using the mineral assemblage of grossular + quartz + calcite, because Grt2 contains quartz and calcite inclusions. The stability limits of grossular strongly depend on the composition of coexisting fluids involving CO$_2$. For simplicity, we considered the stable mineral assemblage in the CaO-Al$_2$O$_3$-SiO$_2$ system along with the fluid of the H$_2$O-CO$_2$ binary mixture (CASH-CO$_2$). The stability field of the assemblage grossular + quartz + calcite was bounded by the following T-sensitive reactions:

\[
\text{zoisite + calcite + quartz = grossular + H}_2\text{O + CO}_2 \quad (1)
\]

\[
\text{calcite + quartz = wollastonite + CO}_2 \quad (2).
\]

As both reactions involve CO$_2$ as a volatile component, the \( X_{\text{CO}_2} \) of the fluid strongly affects the position of
Figure 2. (a) A scan of the thinsection showing the microstructure of the studied sample (EI1901). (b) Back-scattered electron (BSE) image of the garnet in the quartz pod. (c) Fe X-ray image of the garnet in the quartz pod. The dark and bright part in the BSE image correspond to the Fe distribution. (d) BSE image of datolite found in the garnet in the quartz pod. The datolite grain shows faceted boundary and is surrounded by the Grt2. (e) Chemical composition of garnet in the quartz pod plotted in a ternary composition diagram. Abbreviations of the minerals are as follows: garnet (Grt), clinopyroxene (Cpx), quartz (Qz), calcite (Cal), aragonite (Arg), datolite (Dat), almandine (Alm), grossular (Grs), and andradite (Adr).
Datolite in a high-\(P\) marble from the Sanbagawa metamorphic belt

The occurrence of datolite instead of danburite is an indication of low \(X_{\text{CO}_2}\) conditions, where \(X_{\text{CO}_2} < 0.02\) was inferred under \(P-T\) conditions of 500 °C and 200 MPa (Burt, 1978; Grew, 1996). Considering two reactions, a polybaric \(T-X_{\text{CO}_2}\) diagram for the range of \(P = 0.8-1.2\) GPa and \(aP-T\) diagram based on the assumption of \(X_{\text{CO}_2} = 0.01\) and 0.001 were calculated using Perple_X version 6.8.5 (Connolly, 2005) (Figs. 4a, and 4b).

The \(T-X_{\text{CO}_2}\) diagram adequately constrains the upper limit of \(X_{\text{CO}_2}\) to approximately 0.01 at \(T < 550\) °C (Fig. 4a). The two reaction curves and quartz–Raman barometry (\(\Delta \omega_1 = 4.5\)) intersect at conditions of 400–650 °C and 0.8–1.3 GPa (Fig. 4b), which shows distinctly lower pressure than the estimated eclogite–facies conditions of the

**Figure 3.** (a) Raman spectra of the datolite in the studied sample and reference data obtained from the online RRUFF database (https://rruff.info: Lafuente et al., 2015). The RRUFF sample number is also shown. (b) Histogram of \(\Delta \omega_1\) values obtained from the quartz inclusions in the Grt2.

In the Besshi district, the exhumation \(P-T\) history of the rocks in the eclogite unit were considered to be as follows: (1) decompression and cooling from the eclogite facies conditions (~650 °C, 2 GPa) to the epidote–amphibolite facies (~550 °C, 1 GPa), (2) juxtaposition with the non-eclogite unit, which was also subjected to exhumation, and (3) the eclogite–non-eclogite units were heated together to the peak temperature stage (Kouketsu et al., 2014a). As the carbonate inclusion species changed from aragonite to calcite during the transition from the growth of Grt1 to Grt2, the latter is considered to have formed during decompression stage (1) or (2) listed above. The estimated \(P-T\) conditions are almost identical to that of the juxtaposition of the eclogite unit with the non-eclogite unit in the Besshi district (Fig. 4b).

The stability of the datolite in the CaO–B₂O₃–SiO₂–H₂O system is constrained by the following reactions:

\[
2\text{ datolite} + 2\text{H}^+ = \text{danburite} + \text{Ca}^{2+} + 2\text{H}_2\text{O} \\
\text{wollastonite} + \text{H}_2\text{BO}_3 = \text{datolite} + \text{H}_2\text{O} \\
\text{quartz} + \text{H}_2\text{BO}_3 + \text{Ca}^{2+} = \text{datolite} + 2\text{H}^+.
\]

These reactions qualitatively indicate that the stability of datolite favors either low \(H^+\) activity or high \(H_2\text{BO}_3\) activity. Figure 4c shows the stability relations of datolite, danburite, and wollastonite in the CaO–B₂O₃–SiO₂–H₂O system under the conditions of 500 °C and 1 GPa. Calculations were conducted using the deep earth water (DEW) model (Sverjensky et al., 2014; Huang and Sverjensky, 2019) using DEW Excel spreadsheet version 11.0.2. The thermodynamic properties of datolite were sourced from the study by Yang and Rosenberg (1995), and those of wollastonite and quartz from the study by Berman (1988). As shown in Figure 4c, an increase in the activity of \(H_2\text{BO}_3\), such as infiltration of B-rich fluid, would stabilize datolite at relatively high \(a_{\text{Ca}^{2+}}/a_{\text{H}^+}^2\) conditions. Compared to the diagram illustrating relatively lower \(T\) and \(P\) (250 °C and 100 MPa) (Yang and Rosenberg, 1995), the lower limit of log \(a_{\text{H}_2\text{BO}_3}\) was higher by approximately 1.5 (~1.7). As a high \(a_{\text{Ca}^{2+}}/a_{\text{H}^+}^2\) is also required to prevent the precipitation of danburite, if we include \(\text{CO}_2\) in the considered system and calcite precipitates from the solution, \(a_{\text{Ca}^{2+}}/a_{\text{H}^+}^2\) would decrease; and datolite is decomposed to either quartz or danburite. The datolite in our sample is enclosed in Grt2. This is an indication of the simultaneous precipitation of datolite at the early stage of the precipitation of Grt2. The timing of datolite precip-
Yoshida et al. (2011) calculated the partitioning coefficient of B between aqueous fluid and a metapelite collected from the oligoclase–biotite zone of the Besshi district under the prograde P–T path of the relevant area, indicating a considerable release of B from the subducting metapelites. Despite the origin of the fluid that affected Grt2–formation being unclear, B–enriched aqueous fluids can be derived externally from the surrounding rocks of the Eastern Iraatsu body at the timing of the juxtaposition. Sunada and Sakakibara (2004, 2006) reported tourmaline–rich layer from the adjacent area of the eclogite unit of the Besshi district. Based on the mode of occurrence and coexisting minerals, they indicated that B–rich aqueous fluid activity took place during the peak metamorphic stage of the non-eclogite unit (i.e., epidote–am-
phibolite facies conditions). In many subduction-related rocks, a common borosilicate is tourmaline as it is stable over wide $P$-$T$ range (Schreyer and Werding, 1997; Dutrow and Henry, 2011), and thus, tourmaline is used to decipher B-rich fluid activity (e.g., Kawakami et al., 2019). However, the experimental study of amphibolite and B-bearing fluids indicates that tourmaline becomes unstable under alkaline conditions ($pH > ~ 6.5$–$7.0$) regardless of the boron concentration in mafic system (Morgan and London, 1989). In contrast, datolite can be stable under alkaline (low $a_\text{H}_{2}\text{O}$) and high $a_\text{Al}_{2}\text{O}_{3}$ conditions in calcic system. Our findings regarding datolite formed under epidote-amphibolite facies conditions provide new form of the evidence of B-bearing fluid activity in calc-silicate rocks.

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