Two groups of fluid inclusions in the Yunotani eclogite from the Hida-Gaien Belt: Implications for changes of fluid salinity during exhumation

Yuzuki SHINJI*, Tatsuki TSUJIMORI**, and Tatsuhiko KAWAMOTO***

*Department of Earth Science, Graduate School of Science, Tohoku University, Sendai 980–8578, Japan
**Center for Northeast Asian Studies, Tohoku University, Sendai 980–8576, Japan
***Department of Geoscience, Shizuoka University, Shizuoka 422–8529, Japan

Late Paleozoic Yunotani eclogites in the Itoigawa–Omi area of the Hida–Gaien Belt were subjected to a blueschist–facies recrystallization and deformation after the peak eclogite–facies metamorphism. We studied fluid inclusions in quartz domains of different metamorphic stages in the retrograde eclogite. Coarse-grained quartz filling pressure shadow of porphyroblastic garnets contains transgranular fluid inclusions (Group-I) with high salinity [6.0–8.0 eq mass% NaCl, with a mean value 7.2 ± 0.6 eq mass% NaCl (n = 12)]. In contrast, transgranular fluid inclusions in later stage quartz–rich band (Group–II) in parallel to a penetrative foliation developed in matrix are characterized by lower salinity [1.1–5.8 eq mass% NaCl, with a mean value 3.4 ± 1.2 eq mass% NaCl (n = 12)]. The occurrences of fluid inclusion trails suggest that each group corresponds to the fluids entrapped prior to the blueschist–facies recrystallization and the fluids entrapped during the blueschist–facies stage, respectively. The Group–II fluid inclusions formed at blueschist–facies conditions support the idea that aqueous fluids in the subduction channel have a salinity similar to seawater (~ 3.5 eq mass% NaCl). Moreover, the Group–I fluid inclusions indicate a possibility that aqueous fluids at the eclogite–facies depth in subduction channels possess a salinity higher than seawater.

Keywords: Aqueous fluid inclusion, Retrograde eclogite, Blueschist–facies recrystallization, Itoigawa-Omi area, Hida–Gaien Belt

INTRODUCTION

Aqueous fluid inclusion is a minor but ubiquitous microscopic phase in various high-pressure and ultra-high pressure [HP (–UHP)] metamorphic rocks. Since the minute aqueous fluid inclusions are trapped during and/or after metamorphic crystal growth, a great attention has been focused on metamorphic fluids in subduction zone environment (e.g., Philippot et al., 1998; Scambelluri and Philippot, 2001; Frezzotti and Ferrando, 2015). Numerous fluid inclusion studies have found common occurrence of saline fluids (1–10 eq mass% NaCl) and even brines (10–45 eq mass% NaCl) in exhumed HP-UHP rocks (e.g., Barr, 1990; Giaramita and Sorensen, 1994; Gao and Klemd, 2001; Sachan et al., 2017; Kawamoto et al., 2018; Brooks et al., 2019; Liu et al., 2019). The ubiquitous occurrence of fluid inclusions with high-salinity in subducted oceanic materials is critically important to evaluate fluid-mediated enhancement of various metamorphic reactions in subducting oceanic plates as well as fluid-mediated element cycling in solid Earth. Experimental studies have revealed that H2O–NaCl solution controls the stability limit of some metamorphic minerals and element mobility (e.g., Manning, 2004; Tropper and Manning, 2004; Newton and Manning, 2006; Kawamoto et al., 2014). Although there is a significant range of salinity of fluid inclusions entrapped in HP rocks, Kawamoto et al. (2018) suggested that aqueous fluids in the subduction channel have a salinity similar to seawater (~ 3.5 eq mass% NaCl). Brooks et al. (2019) proposed a model to form higher salinity fluids up to ~ 35 eq mass% NaCl from primary fluid inclusions with ~ 4.0 eq mass% NaCl; they suggested fluid–rock interaction...
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in a relatively closed system. On the other hand, there are many observations of different salinities among different occurrences of fluid inclusion clusters/trails in an identical specimen. For examples, primary fluid inclusions in eclogite-facies minerals have higher salinity than that of transgranular (or secondary) (Simmons and Richter, 1976; Kranz, 1983) fluid inclusions entrapped during retrograde metamorphism (e.g., Sachan et al., 2017; Liu et al., 2019). However, salinity systematics of subduction zone metamorphism is not well constrained yet. In order to better understand the systematics, we focused on transgranular fluid inclusions in two types of quartz domains formed at different metamorphic stages in a retrograde eclogite, which is an eclogite-facies rock suffered from retrograde metamorphism overprint (e.g., Zhang et al., 2003). As described below, we found two groups of fluid inclusions with different occurrences and salinities. In this paper, we present the first report of fluid compositions at different metamorphic stages in Japanese late Paleozoic eclogites with a so-called ‘Franciscan-type’ retrograde pressure-temperature (P-T) trajectory, in which the rocks retrace the almost identical path during exhumation to the prograde burial P-T path.

**GEOLOGIC SETTING**

The Hida-Gaien Belt of the Hokuriku Region is a tectonic mixture of various pre-Jurassic lithologic slices of the Oeyama, Renge, Akiyoshi, and Maizuru Belts that are widely developed in Southwest Japan. In this belt, the late Paleozoic high-pressure (HP) schists derived from the Renge Belt occur associated with jadeite-bearing antigorite serpentinite (e.g., Tsujimori, 2002; Tsujimori and Harlow, 2017). The ‘Renge Schists’ record mainly greenschist- to amphibolite-facies metamorphism and locally preserve blueschist- to eclogite-facies metamorphism. In the Omi area, the HP schists are divided into two distinct groups: an ‘Eclogitic’ (EC) unit and a ‘Non-eclogitic’ (non-EC) unit (Tsujimori, 2002; Tsujimori and Matsumoto, 2006; Yamada et al., 2019) (Fig. 1). The EC unit consists mainly of paragonite-bearing metasedimentary schists with mafic layers or blocks of epidote blueschist, garnet blueschist, and glaucophane-bearing eclogite, both of which commonly contain glaucophane and/or barroisite. Prograde-zoned porphyroblastic garnets of eclogites preserve mineral inclusions showing a prograde transition from epidote-blueschist- to eclogite-facies metamorphism (at 2.0–2.2 GPa and 550–600 °C, Tsujimori, 2002; Tsujimori and Matsumoto, 2006). The matrix of eclogite has been subjected to a blueschist-facies recrystallization and deformation after the peak eclogite-facies metamorphism. According to Tsujimori et al. (2000) and Tsujimori (2002), the matrix foliation (S1) is in parallel to an internal fabric (inclusion trail) of the peak eclogite-facies mineral assemblage in the garnet rims. However, the matrix omphacite is commonly replaced by chlorite, and initially coarse-grained glaucophane is reduced in grain-size. A pumpellyite-bearing retrograde blueschist-facies mineral assemblage is found in the garnet blueschist, an Mg-rich equivalent of the eclogite (Shinji and Tsujimori, 2019).

**SAMPLE DESCRIPTIONS**

We investigated fluid inclusions in retrograde eclogites found in the Yunotani Valley (Tsujimori et al., 2000; Tsujimori, 2002) (Fig. 1). The studied sample (YTECT02) is a well-foliated and retrograded eclogite, which characteristically contains quartz domains. The sample consists mainly of glaucophane, porphyroblastic garnet (up to ~5 mm in size), omphacite, epidote, quartz, rutile, and phengite. Titanite, chlorite, and rare albite occur as secondary minerals. A penetrative foliation (S1) is defined by a preferred orientation of neoblastic glaucophane in the matrix as well as inclusion fabrics of the mantle and rim portion.
of porphyroblastic garnets (Tsujimori, 2002). In the studied sample, two types of quartz domains occur: one is coarse–grained quartz filling pressure shadow of porphyroblastic garnets (Fig. 2a), the other is a quartz–rich band in parallel to the \( S_1 \) foliation (Fig. 2b). The presence of omphacite in some of the quartz domains at the pressure shadows suggests that the quartz domains were formed prior to the blueschist–facies recrystallization (Fig. 2a). In addition, the preservation of rutile without partial replacement by titanite suggests that the pressure shadow quartz domains were prevented from retrograde recrystallization and deformation (Fig. 2c). On the other hand, the mineral assemblage glaucophane + epidote + titanite in the quartz–rich band suggests that the crystallization (or recrystallization) of quartz in the band was postdated the peak of eclogite–facies metamorphism (Fig. 2d).

Abundant fluid inclusion trails, normally \(~ 10\)–\(~ 50 \)\( \mu \)m wide in the thin section, occur in both types of quartz domains. The fluid inclusions are commonly two-phase (liquid + vapor), liquid–rich fluid inclusions with size variation from \(~ 1 \) to \(~ 10 \)\( \mu \)m in length and angular to rounded shape (Figs. 2e and 2f).

Based on the occurrences of fluid inclusion trails, we defined Group–I and –II fluid inclusions. The Group–I inclusions are transgranular inclusions, aligned at high angles to the \( S_1 \) foliation, and do not reach the boundary between the quartz domain and matrix (Fig. 2c). In contrast, the Group–II inclusions are also transgranular inclusions aligned along sealed microfractures but mostly in parallel to the \( S_1 \) foliation. Considering these, the Group–I and –II fluid inclusions were entrapped during the formation of the pressure shadow quartz domains and the quartz–rich band, respectively.

Phase identification of fluid inclusions was performed by HORIBA XploRA PLUS Confocal Raman Microscope at the Graduate School of Environmental Studies, Tohoku University. A 532 nm solid–state Nd–YAG laser with 10 mW power was used as the light source. The Raman spectra were measured ranging from 200 to 4400 cm\(^{-1}\) using a grating of 600 g/mm. The measuring area is \(~ 2 \)\( \mu \)m; the exposure time is 50 s (5 s \( \times \) 10). The Raman shift was calibrated using a reference silicon. All of analyzed fluid inclusions show room–temperature Raman spectrum with the OH–stretching (\( \nu_{\text{OH}} \)) band at around \(~ 3200\)–\(~ 3600 \)cm\(^{-1}\), indicating the presence of liquid H\(_2\)O (Fig. 3). Any gas species were not detected.
Doubly polished wafers with a thickness of about 40 ± 15 µm were prepared from the sample YTECT02 using a Metkon Forcipol 1V grinder/polisher, a Metkon Micracut 152 precision cutter and a Maruto Prepalap MG-300 polisher at Tohoku University. Then microthermometric measurements of fluid inclusions in one of the wafers (YTECT02_ts39) were performed on a Linkam Scientific THMS600 heating/cooling stage with ×100 objective lens at Shizuoka University. Temperature was calibrated using synthetic fluid inclusion standards at −56.6 °C (melting point of CO₂), 0 °C (melting point of H₂O), and 373 °C (critical temperature of H₂O). Th (homogenization temperature) and Tm (ice melting temperature) were determined at a heating rate of 0.5 °C/min and 0.2 °C/min at around Th and Tm, respectively. These rates were controlled with a constant heating rate of 0.1 °C/min toward desired temperature. We estimated the salinity of H₂O fluid inclusions using the relationship between salinity and freezing point depression (Bodnar, 1993). Precision of the Th is within 1 °C, and that of Tm is within 0.1 °C, which results in estimated uncertainties of individual salinity measurements of 0.12–0.18 eq mass% NaCl in the range of 0 to 10 eq mass% NaCl (Bodnar, 1993).

The microthermometry results for all types of fluid inclusions are summarized in Table 1 and Figure 4a. The results are also plotted in a Th–Tm diagram (Fig. 5). The measured Group-I inclusions belong to a single trail, while those of the Group–II inclusions are located along several trails. All measured fluid inclusions show freezing between −38 to −51 °C characterized by sudden shrinkage of the vapor during cooling, and we observed first melting at ~ −20 °C in one of the Group–I inclusions (FI/Qz–#014_G1), indicating that NaCl or/and KCl present dominantly in the aqueous phase (Roedder, 1984). The salinity of the Group–I fluid inclusions ranges from 6.0 to 8.0 eq mass% NaCl, with a mean value 7.2 ± 0.6 eq mass% NaCl (1 s.d., n = 12). In contrast, the Group–II fluid inclusions are characterized by lower salinity, ranging from 1.1 to 5.8 eq mass% NaCl, with a mean value 3.4 ± 1.2 eq mass% NaCl (1 s.d., n = 12).

Table 1. Measured Tm and Th and derived calculated salinity

| Group | FI/Qz–#005_G1 | FI/Qz–#006_G1 | FI/Qz–#007_G1 | FI/Qz–#008_G1 | FI/Qz–#010_G1 | FI/Qz–#012_G1 | FI/Qz–#013_G1 | FI/Qz–#014_G1 | FI/Qz–#015_G1 | FI/Qz–#020_G1 | FI/Qz–#022_G1 | FI/Qz–#044_G1 |
|-------|---------------|---------------|---------------|---------------|---------------|---------------|---------------|---------------|---------------|---------------|---------------|---------------|
| Th (°C)| 7.2 (0.6)     | -5.1          | 7.3           | -6.6          | 7.3           | -6.6          | 7.1           | -4.4          | 6.2           | -3.9          | 6.9           | -4.3          |
| Tm (°C)| 8.0           | 4.7           | 7.4           | 4.7           | 6.8           | 6.0           | 7.3           | 7.4           | 6.8           | 6.0           | 8.0           | 7.7           |
| Derived salinity, eq mass% NaCl | 151           | 147           | 177           | 162           | 140           | 181           | 178           | 195           | 156           | 154           | 161           | 167           |
| Mean (1σ) | 7.2 (0.6)     | -4.5 (0.4)    | 164 (17)      |               |               |               |               |               |               |               |               |               |

Figure 4. (a) Histograms showing the variation of salinity of the measured fluid inclusions. Numbers indicate mean values of salinity. (b) Variations of fluid inclusion salinities in rocks similar to the studied sample (Sachan et al., 2017) and Dominican–Republican jaditeites (Kawamoto et al., 2018).
mean value of 164 ± 17 °C (1 s.d.). The $T_b$ for the Group-II fluid inclusions yield a range from 136 to 167 °C, with a mean value of 151 ± 11 °C (1 s.d.). There is no obvious correlation between $T_b$ and $T_m$, i.e. salinity (Fig. 5). A large variation of $T_m$ for the Group-I inclusions suggests that the Group-I inclusions suffered from various degrees of density/$\rho$ modification by stretching of fluid inclusion due to internal overpressure (e.g., Prezbindowski and Larson, 1987). However, the values of $T_m$ have a small variation within the Group-I inclusions, and no texture indicating fluid leakage/refilling such as fractures and/or a secondary inclusion halo around an inclusion (Sterner and Bodnar, 1989) was observed, suggesting that the Group-I inclusions preserve the original fluid composition.

It is notable that our preliminary analysis of aqueous fluid inclusions in omphacite in the same sample (YTTECT02_ts39) shows salinity lower (mean value of 2.6 ± 0.7 eq mass% NaCl, $n = 10$) than the Group-I inclusions, although the timing of the entrapment remains unknown. The detail of the fluid inclusions in the omphacite will be described in the future.

**SIGNIFICANCE**

The Yunotani eclogite records a so-called ‘Franciscan-type’ $P$-$T$ trajectory, in which the rocks retrace the almost identical path during exhumation to the prograde burial $P$-$T$ path (Tsujimori, 2002). Especially, the studied sample (YTTECT02) suffered from intensive blueschist–facades recrystallization and deformation. As we reported above, we found two groups of fluid inclusions with different types of occurrence and salinities. The occurrence of the fluid inclusion trails suggests that each group corresponds to the fluids entrapped prior to the blueschist–facades recrystallization and the fluids entrapped during the blueschist–facades stage, respectively. Considering this, the observed difference of salinities would reflect difference of fluid compositions at different depths in a paleo-subduction zone. Based on the fluid inclusion study of jadeites formed at blueschist–facades conditions, Kawamoto et al. (2018) suggest that aqueous fluids in the subduction channel have a salinity similar to or slightly higher than seawater (~3.5 eq mass% NaCl) (Fig. 4b). In our study, the Group-II fluid inclusions formed at a blueschist-facies condition support this idea. However, higher salinity in the Group-I fluid inclusions requires other explanation to have higher salinity.

There are a few reports of difference in fluid inclusion salinities among eclogite–facades condition and later retrograde conditions during exhumation in subduction setting (e.g., Sachan et al., 2017; Liu et al., 2019). Sachan et al. (2017) studied fluid inclusions in blueschist–facades retrograde eclogites in the Shergol Ophiolitic Mélange of Northwest India. They reported that eclogite–facades stage primary fluid inclusions in garnet and quartz included in garnet had higher salinity (7.8-14 eq mass% NaCl) than secondary fluid inclusions in quartz formed at blueschist-facies retrograde stage (1.7-6.4 eq mass% NaCl) (Fig. 4b). The presence of omphacite in the pressure-shadow forming quartz domains in the studied sample suggests that the Group-I fluid inclusions were captured at near peak conditions prior to blueschist-facies condition. Considering our data of the Group-I fluid inclusions together with the case of the Shergol Ophiolitic Mélangé, aqueous fluids at the eclogite–facades would be characterized by a salinity higher than seawater. For further understanding of the possible mechanism of the high salinity along the subduction channel, more systematic data focusing on fluid inclusions trapped during eclogite–facies metamorphism are required.

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