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Tracing the Scale of Fluid Flow in Subduction Zone Forearcs: Implications from Fluid-Mobile elements

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ABSTRACT

Despite the importance of fluids in subduction zone processes, the extent of mass transfer and fluid circulation from the subducting plate through the forearc region remain unclear. To estimate the fluid budget and scale of fluid circulation in subducted sediments, we assess the distribution and retention of fluid-mobile elements (FME) in metamorphically equivalent metapelites from the Kodiak complex (Alaska) and the Shimanto Belt (Japan). The temperature range of interest is 230–350 °C, i.e., encompasses the base of the seismogenic zone. We examine the Li, B, Rb, Sr, Cs, and Ba concentrations in the bulk rock, as well as in fluid inclusions and individual minerals by (LA-) ICP-MS.

The whole-rock composition in metapelites from Kodiak shows no significant loss of FME, which is in contrast to Shimanto where between 10% and 55% of FME (particularly Li, B and Cs) are leached out of rocks. The FME budget in Kodiak is consistent with a redistribution of elements between metamorphic illite and chlorite, whereas in Shimanto the loss of Li, B and Cs as temperature increases is the result of decreasing concentrations in illite and chlorite. The semi-quantitative analysis of fluid inclusions is consistent with a significant enrichment in all analyzed trace elements relative to seawater and interstitial pore fluids of seafloor sediments.

Combining the fluid compositions with the whole-rock compositions, mass balance calculations were performed for B, Cs, and Ba. In the Kodiak complex the mass balance calculations are consistent with closed-system behavior, wherein the fluid is an insignificant reservoir for FME. Conversely, in the Shimanto Belt the mass balance calculations are consistent with open-system behavior, wherein large amounts of fluid percolated through rocks and the mass water-rock ratios correspond to 0.5–2.2. We infer that such an open system behavior was promoted by a larger amount of internal strain and the proximity to a large-scale fault zone. Moreover, fluid compositions observed in this study exhibit similarities to the composition of mud volcano fluids. This similarity is consistent with extensive, focused fluid circulation originating from depths of at least 15 km and ascending to the surface through a substantial damage zone associated with an out-of-sequence thrust.

1. Introduction

Fluids play a major role in many subduction zone processes, such as the element cycling between the different reservoirs of the Earth (Bebout, 1995, 2014, 2021; Hacker et al., 2003; Peters et al., 2017; Shervais and Jean, 2012; Tatsumi and Kogiso, 2003), seismicity (Audet and Schwartz, 2013; Behr and Bürgmann, 2021), or back-arc volcanism (Hacker, 2008; Schmidt and Poli, 2014; Stern, 2011). In addition to serpentinites and altered oceanic crust (Deschamps et al., 2013; John et al., 2011; Reynard, 2013; Rupke, 2004; Scambelluri et al., 2004a, 2019; Tenthorey and Hermann, 2004), sediments are an important carrier of volatiles, such as carbon dioxide or water (Bebout et al., 2013; Bekaert et al., 2020; Kerrick and Connolly, 2001; Rea and Ruff, 1996). Despite the critical role of fluids, the extent of fluid circulation in subducted material, as well as the quantity of fluids passing through permeable horizons, remain uncertain.

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Along with burial, load and temperature increase, sediments progressively lose the water they contain, following several processes (Saffer and Tobin, 2011). First, at shallow depths (typically <5 km), the volume of pore fluid is significantly reduced due to compaction and cementation (Kastner et al., 2014; Kominz and Pekar, 2001; Saffer and Tobin, 2011). Then, starting at ~50 °C with diagenetic reactions of hydrated silica phases and quartz and proceeding down to eclogitefacies conditions, successive metamorphic reactions result in a prograde decrease in mineral-bound water (Bebout, 2007; Hacker et al., 2003; Schmidt and Poli, 2014). The majority of fluids released early in the subduction history (<5 km) circulate along the subduction interface back to the ocean (You et al., 1995; Chan and Kastner, 2000; Saffer and Tobin, 2011), whereas in deeper regions (5-15 km) expelled fluids return to the ocean along out-of-sequence thrust faults in forearcs (Bangs et al., 1996; Bangs et al., 1999; Lauer and Saffer, 2012, 2015; Moore et al., 2001; Tobin et al., 2001; Ujiie, 2003; Vrolijk et al., 1991).

Dehydration fluids have compositions that differ significantly from seawater or original pore fluids, as a result of mineral reactions, such as the smectite-to-illite transformation (see the compilation by Kastner et al., 2014; Herviou et al., 2021). In particular, they are characterized by elevated fluid-mobile elements (FME) concentrations, even at low depths (Barnes et al., 2019b; Dia et al., 1999; Dimitrov, 2002; Martin et al., 1996; Westbrook and Smith, 1983). As a result, fluid-mobile elements have been extensively used to follow fluid and fluid-mediated transfers, including distinguishing between open or closed systems (Barnes et al., 2019b; Bebout et al., 2013; Bebout et al., 2007; Bebout et al., 1999), tracking dehydration reactions and loss of volatiles (Dragovic et al., 2012; John et al., 2011; Menzies et al., 2022; Peters et al., 2017; Scambelluri et al., 2004a), or tracing the scale of mass transfer and fluid circulation (Angiboust et al., 2014; Barnes et al., 2019a; Hoover et al., 2022; Ishikawa et al., 2008; Muñoz-Montecinos et al., 2021; Shervais and Jean, 2012; Zack and John, 2007).

Presumably, the FME budget is strictly controlled by the geothermal gradient, which governs the mineral reactions that occur along the prograde metamorphic path. For instance, Bebout et al. (2013) noted that in very cold subduction zones, like those preserved in the Western Alps (the gradient of $\sim 8 \,^{\circ}\text{C/km}$), no major dehydration reactions have occurred down to \sim 80 km, thus preserving FME concentrations that are similar to those found in the potential sedimentary protolith. Conversely, metasedimentary rocks from the Catalina Schist, California, exhibit a progressive loss of FME (in particular B and Cs) along the prograde metamorphic path (Bebout et al., 2007). The B and Cs loss is attributed to the prograde devolatilization reactions, such as the progressive recrystallization of mica (Bebout et al., 2007). These studies belong to an abundant literature focused on tracking mass and fluid transfer in metamorphic series of rocks for temperature conditions above \sim 300 °C, whereas little is known for the lower conditions of the seismogenic depth interval (Oleskevich et al., 1999). The base of the seismogenic zone is of particular interest with respect to the issue of FME transfers and fluid circulations, as this range of depths satisfies both conditions of (1) rock brittle behavior, enabling a permeable crack network to form and (2) temperatures high enough for efficient fluidrock interactions.

To elucidate FME and fluid mobility in the seismogenic domain of subduction zones, this work focuses on FME concentrations and budgets (Li, B, Rb, Sr, Cs, and Ba) in metasedimentary units from two paleoaccretionary complexes, the Kodiak complex and Shimanto Belt. This study builds upon earlier research concerning mineral reactions within the same complexes (Rajič et al., 2023b), which demonstrated the absence of dehydration reactions and the preservation of the major chemical composition in the range 230 to 350 °C. In the present work, we examine the evolution of FME budgets as a function of peak-metamorphic temperatures from 230 to 350 °C, by using a combination of three datasets: FME concentrations of whole-rock, individual metamorphic phases, and vein-hosted fluid inclusions. The Kodiak accretionary complex and the Shimanto Belt were selected for the following reasons: (1) Both localities are paleo-accretionary complexes, with the majority of the exposed rocks being basally underplated (Byrne and Fisher, 1987; Ditullio and Byrne, 1990; Fisher and Byrne, 1987; Sample and Moore, 1987; Taira et al., 1982; Ujiie, 1997). (2) Both contain rock units that have experienced a range of (near-) peak metamorphic conditions that correspond to the seismogenic zone and its down-dip limit (from ~ 230 °C up to ~ 350 °C; Vrolijk et al., 1988; Rowe et al., 2009; Raimbourg et al., 2014, 2021; Palazzin et al., 2016; Rajič et al., 2023a). At each locality, syn-subduction veins with primary fluid inclusions associated with a main deformation event have been observed (Raimbourg et al., 2021; Rajič et al., 2023b). (3) Units at each locality have lithological and mineralogical similarities, as well as similarities in deformation style (Raimbourg et al., 2014; Rajič et al., 2023a). (4) The primary distinction between these two complexes is that units from the Kodiak complex are situated at a considerable distance from any largescale fault zones. Conversely, selected units in the Shimanto Belt are separated by the out-of-sequence thrust, the Nobeoka Tectonic Line (Kondo et al., 2005). The NTL was active significantly after the deposition ages of the unit in the hanging wall (Foliated Morotsuka; Fisher et al., 2019). As a result, this study provides (i) insights into fluid-rock interactions in subducted sediments at temperatures <350 °C and their impact on the chemical composition of pore fluids, as well as (ii) a framework for fluid circulation and mass transfer on the cm- to the multi-km scale.

2. Sampling areas

2.1. Kodiak accretionary complex

The Kodiak accretionary complex represents a well described paleoaccretionary complex that consists of coherent slices of metasedimentary units and tectonic mélanges (Fig. 1a), decreasing in age towards the modern trench (Byrne and Fisher, 1987; Connelly, 1978; Moore et al., 1983; Plafker et al., 1994). The Late Cretaceous Kodiak Formation consists of deep trench sediments, and is divided into three belts (Landward, Central and Seaward Belts) based on variations in bedding, cleavage orientation and peak-metamorphic temperatures (Rajič et al., 2023b; Sample and Fisher, 1986). The Kodiak Landward Belt (KLB) consists of intercalations of sandstone/siltstone and shale beds (Fig. 2a), with occasionally disrupted strata closer to the Uganik Thrust (Fisher and Byrne, 1987; Rowe et al., 2009). Veins from the Landward Belt are Mode 1 quartz veins with an orientation that is approximately perpendicular to the bedding (Fig. 2a). Temperatures derived from Raman Spectroscopy of carbonaceous material (RSCM) of the Landward Belt in Uyak Bay range from 240 to 260 °C (Rajič et al., 2023a). The Kodiak Central Belt (KCB) consists of sub-horizontal foliated shale and sandstone layers that are characterized by either (i) topto-the-trench or (ii) conjugate extensive faults and recumbent folds (Rajič et al., 2023a; Sample and Fisher, 1986; Sample and Moore, 1987). Veins used for this study are Mode 1 quartz-(calcite) veins, often in en échelon arrays (Fig. 2c; Raimbourg et al., 2021; Rajič et al., 2023b) and are characterized by crack-seal microstructures (Fisher and Brantley, 2014; Raimbourg et al., 2021). The Central Belt, examined in Seal Bay, experienced temperatures of $\sim 330~^\circ C$ and pressures of $0.30\pm 0.04~GPa$ that are inferred from RSCM and multi-phase equilibrium models (Raimbourg et al., 2021; Rajič et al., 2023b). The geothermal gradient is estimated to be between 20 and 30 °C/km (Rajič et al., 2023b). The Kodiak Seaward Belt (KSB) is characterized by intercalated sandstone and shale layers. This belt experienced strong NW-SE horizontal coaxial shortening (Rajič et al., 2023a; Sample and Moore, 1987). RSCM temperatures of the Seaward Belt range from 300 to 350 °C (Rajič et al., 2023a). Samples from the Kodiak Central and Seaward Belts experienced similar peak-metamorphic temperatures and consequently are treated together as the higher-grade units in the Kodiak complex, while the Kodiak Landward Belt constitutes the lower-grade unit. All samples from the Kodiak Formation used during this study are sampled at least 2



Fig. 1. Geological maps of (a) the Kodiak accretionary complex, Alaska, modified after Rajič et al. (2023a), and (b) the Shimanto Belt on Kyushu, Japan, modified after Raimbourg et al. (2021). The locations of individual samples used in this study are provided on each map.



Fig. 2. Field photos of vein samples used in this study from the Kodiak complex (a and c) and the Shimanto Belt (b and d). (a) Mode 1 quartz-calcite veins in coherent portion of the Kodiak Landward Belt. (b) Quartz veins dominantly located in the sandstone lenses in strongly sheared sample from the Hyuga Tectonic Mélange. (c) A set of en échelon arrays from the Kodiak Central Belt. (d) An example of en échelon arrays from the Foliated Morotsuka.

km away from recognized large-scale fault zones (Fig. 1a).

2.2. Shimanto Belt, Japan

The Shimanto Belt is interpreted as a paleo-accretionary wedge (Taira, 1988) exposed on Honshu, Kyushu and Shikoku islands of Japan, with a general SW-NE trend of units parallel to the modern Nankai Through (Fig. 1b). The Foliated Morotsuka (FM) is the higher-grade unit of the Shimanto Belt. It comprises highly flattened sandstone and basaltic boudins embedded within a metapelitic matrix (Mackenzie, 1989; Raimbourg et al., 2014; Ujiie et al., 2018). Foliation is marked by the alignment of white mica and chlorite, which formed during vertical shortening (Kameda et al., 2011; Raimbourg et al., 2009). The Foliated Morotsuka records extensional tectonics at peak metamorphic conditions in the form of conjugate extensional shear bands (Raimbourg et al., 2014). Veins examined in this study are from en échelon arrays cutting across the foliations and are folded perpendicular to the foliation (Fig. 2d), which have been interpreted to have formed at peak metamorphic conditions (Raimbourg et al., 2021). Pressure-temperature conditions of the Foliated Morotsuka are estimated to be 0.3-0.5 GPa and 300-350 °C, respectively, and are based on RSCM and the metamorphic assemblages in basaltic lenses and multi-phase equilibrium model in metapelites (Palazzin et al., 2016; Rajič et al., 2023b; Toriumi and Teruya, 1988). Based on pressure-temperature results, the geothermal gradient is estimated to be around 20 °C/km (Rajič et al., 2023b; Toriumi and Teruya, 1988). The Foliated Morotsuka represents a hanging wall of the Nobeoka Tectonic Line (NTL), an out-of-sequence thrust (Kondo et al., 2005). The Hyuga Tectonic Mélange (HTM) is the footwall of the NTL and consists of strongly sheared sandstone lenses embedded in a pelitic matrix. The dominant syn-deformation veins are Mode 1 quartz veins perpendicular to the orientation of the sandstone lenses or mineralized shear bands (Fig. 2b). Based on RSCM, the peak temperature of deformation was 245 \pm 30 °C (Palazzin et al., 2016), hence the Hyuga Tectonic Mélange is the lower-grade unit of the Shimanto Belt.

2.3. Mineralogy, textures and fluid inclusions of samples

Samples used for this study are extensively described in Rajič et al. (2023b). What follows is a summary of all relevant information. A sample list and peak-metamorphic temperatures-(pressures) are reported in Table A.1.

Whole-rock, major element compositions do not systematically change as temperature increases, consistent with the preservation of whole-rock composition and a potentially closed system. However, in restricted areas there is evidence of local mass transfer from the host rock to the veins. Main mineral phases in the host rocks in all samples are illite (21-48%), chlorite (0-24%), quartz (16-43%) and albite (5-26%). Minor smectite-illite mixed-layers are preserved only in lowergrade samples (1–5%). With increasing temperature, the major change in mineralogy is an increase in chlorite over other phyllosilicates (0-27% in low-grade samples vs 20-44% in higher-grade samples). In higher-grade samples, the smectite-to-illite transformation (later in the text referred as the illite recrystallization) is complete and is reflected by an increase in potassium in the interlayer site. Locally in higher-grade samples, quartz and albite are dissolved and reprecipitated in the veins. In lower-grade samples, from HTM and KLB, the grain size of illite and chlorite varies from 1 to 60 μ m in length in the pelitic matrix and is up to 120 µm in length when in contact with quartz ribbons (Fig. 3a-b). At higher temperatures the grain size of chlorite and illite in the KCB range from 5 to 400 μm and 1 to 80 $\mu m,$ respectively. The grain size of both phases is noticeably larger when in contact with the veins and is up to 775 µm (Fig. 3c). Similarly, samples from FM contain grains of illite and chlorite in the pelitic matrix that are characterized by significantly larger grain sizes of up to 1 mm (Fig. 3d).

Syn-subduction veins are characterized by different textures

depending on the temperature at which they formed. Mode 1 quartz tension veins in lower-grade units are characterized by euhedral crystals with intercalation of high-luminescence (CL-blue) and lowluminescence (CL-brown) zones, which correlates to the aluminum concentration (Raimbourg et al., 2021). Veins in higher-grade units contain crack-seal textures with low-luminescent quartz. The quartz in all of the veins is characterized by dense populations of fluid inclusions (FIs), with local zones containing a low density of fluid inclusions (Fig. 4a-b). In lower-grade veins, two generations of primary FIs are observed: one-phase CH₄-rich and two-phase H₂O-rich FIs (Fig. 4c). Quartz in higher-grade veins contains only two-phase H₂O-rich FIs (Fig. 4d). The salinity of primary FIs also differs (Table A.1), wherein FIs in lower-grade quartz have salinities lower than that of seawater (~ 2 wt % NaCl eq.) and higher-grade FIs have salinities closer to or higher than that of seawater (\sim 3.5 wt% NaCl eq.). The salinity of secondary FIs in all analyzed samples is <1 wt% NaCl eq.

3. Analytical and computational methods

3.1. Whole-rock trace element composition

Whole-rock trace element compositions were acquired for 16 samples from the Kodiak Formation (Landward, Central and Seaward Belt) and the Shimanto Belt (the Hyuga Tectonic Mélange and the Foliated Morotsuka). After removing any visible macro- and micro-veins, the metasedimentary samples were crushed in a mortar and washed with doubly distilled water. Subsequently, the sample powders were rinsed twice with doubly distilled water and then dried in an oven at 90 °C for 24 h. Final powder samples were analyzed at Actlabs (Activation laboratories Ltd. Ireland; analytical code ME-MS89L + B-MS89L). Samples (0.2 g) were fused with sodium peroxide at 670 °C and mixed with hydrochloric acid after cooling. Prepared solutions were analyzed by inductively coupled plasma-mass spectrometry. Boron has been analyzed separately by glassless digestion analysis to eliminate boron from labware. The detection limits of FME of interest are as follows: Li = $2 \mu g/g$; B = $8 \mu g/g$; Rb = 0.5 $\mu g/g$; Sr = $20 \mu g/g$; Cs = 0.1 $\mu g/g$; and Ba = $2 \mu g/g$. For remaining analyzed trace elements, the detection limits are given in Table A.2, as well as analyzed duplicates, standards (GIOP-111, AMIS0055, LR-LOI3, and LR-LOI5), and the sample blanks. The standards were measured as unknowns and were within 10% of accepted values (Table A.2).

3.2. Laser Ablation Inductively coupled Plasma Mass Spectrometry

3.2.1. Trace-element concentration in mineral phases

Laser Ablation Inductively Coupled Plasma Mass Spectrometry (LA-ICP-MS) was used to determine the trace element concentrations in major mineral phases (illite, chlorite, and albite in the pelitic matrix, and quartz and calcite in the veins). Sufficiently large grains (wider than 30 µm) for LA-ICP-MS analysis were carefully selected by using optical microscopy and SEM-BSE, as well as chemical mapping of thin sections (Rajič et al., 2023b). LA-ICP-MS analyses were conducted at CNRS Institut de Sciences de la Terre d'Orléans (ISTO) using a RESOlution-SE 193 nm ArF excimer laser with a S155 ablation cell coupled to an Agilent 8900 QQQ mass spectrometer. To determine trace element concentrations in illite, chlorite, albite, and vein calcite, the laser was operated at 3 Hz and a fluence of \sim 4 J/cm² on the sample surface, and a spot size of 10 µm in diameter. A SQUID was used to homogenize the ablation signal. For each phase and spot, the SiO₂ content measured by electron microprobe was used as an internal standard and NIST610 was used as an external standard. For calcite, predetermined CaO content measured by electron microprobe was used as an internal standard. NIST612 was also analyzed as unknowns for quality control. The analyzed isotopes were: ⁷Li, ¹¹B, ⁵¹V, ²⁷Al, ²⁸Si, ⁵⁶Fe, ⁶⁰Ni, ⁶³Cu, ⁶⁶Zn, ⁷⁵As, ⁸⁵Rb, ⁸⁸Sr, ⁹³Nb, ¹¹⁸Sn, ¹³³Cs, and ¹³⁷Ba. Data reduction was performed with the software GLITTERTM (Van Achterbergh et al., 2001), wherein



Fig. 3. Photomicrographs (optical microscope - a, b, d and SEM-BSE - c) images of metapelites from the Kodiak complex (a and c) and the Shimanto Belt (b and d). Low-grade metapelites characterized by intercalations of pelitic matrix and quartz ribbons from (a) the Kodiak Landward Belt (Sample KO17E) and (b) the Hyuga Tectonic Mélange (Sample HN299). Pelitic matrix contains fine-grained illite, chlorite, albite, and quartz, with only occasionally larger illite and chlorite grains. (c) A SEM-BSE (left) and a phase-map (right) of the matrix in the contact with a quartz shear vein, Kodiak Central Belt (Sample KO38). Note a grain size increase in the pelitic matrix towards the contact with the vein. (d) Metapelite with the pervasive foliation, the Foliated Morotsuka (Sample HN247). Note a significantly higher grain size than in previous samples.



Fig. 4. Quartz and entrapped fluid inclusions from syn-subduction veins. (a-b) Quartz crystals characterized by FI-rich and FI-poor zones (Samples KO17 and KO33A2). Note that the FI-poor zone in (b) still contains fluid inclusions, but in comparison with adjacent FI-rich zone in significantly lower amount. (c) An example of coexisting one-phase CH₄-rich and two-phase H₂O-rich fluid inclusions in lower-grade veins (Sample KO17). (c) Primary two-phase H₂O-rich fluid inclusions with CO₂-(CH₄) vapor bubble in higher-grade veins (Sample KO33A2).

background and ablation signals were manually selected and a linear fit was applied to all standard data. The values of quality control (NIST612) are reported in the Table A.3.

3.2.2. Bulk microanalysis of fluid inclusion populations

The size of the FIs hosted in quartz in the studied samples rarely exceeds 10 μ m and thus, the classical approach of analysis by laser ablation of individual FIs was not applicable. Instead, bulk microanalysis of FI populations, following the approach of Tuba et al. (2019), was utilized. Tuba et al. (2019) showed that this approach is in a good agreement with other methods for determining the chemical composition of FIs, such as microthermometric studies, decrepitation method or crush-leach techniques.

Doubly-polished thick sections of quartz were prepared with a thickness of ~200–250 μ m. To avoid heating and volatilization of the glue during analysis, all superglue was removed after polishing by immersing samples in acetone for up to ~24 h and then rinsing repeatedly. The margins of the thick sections were subsequently fixed to a new glass plate with superglue. The laser was operated at a frequency of 5 or 10 Hz and a fluence of 8 J/cm² on the sample surface with a spot size of 50 μ m in diameter. The isotopes that were analyzed were ⁷Li, ⁹Be, ¹¹B, ²³Na, ²⁴Mg, ²⁷Al, ²⁸Si, ³⁹K, ⁴³Ca, ⁴⁷Ti, ⁵⁵Mn, ⁵⁶Fe, ⁶⁰Ni, ⁶³Cu, ⁶⁶Zn, ⁷⁵As, ⁸⁵Rb, ⁸⁸Sr, ⁸⁹Y, ⁹⁵Mo, ¹¹⁸Sn, ¹²¹Sb, ¹³³Cs, ¹³⁷Ba, ¹⁸²W, ²⁰⁸Pb, and ²⁰⁹Bi. SiO₂ was used as the internal standard (100 wt%) and NIST610 was used as the external standard. NIST612 and BCR2G were also analyzed as unknowns for quality control.

In all veins that were examined, quartz displays distinct variations in FI abundance, which we refer to as FI-rich and FI-poor quartz zones (Fig. 4a-b). In FI-poor zones, quartz contains a significantly reduced quantity of FI (Fig. 4a-b). However, it is crucial to note that quartz is never entirely devoid of FIs (Fig. 4a-b). Before analysis, these FI-poor and FI-rich zones were qualitatively identified in each sample using

optical microscopy.

The first step involves laser ablation analyses in FI-rich and FI-poor zones in quartz, to distinguish elements concentrated within the FIs from the elements hosted in the quartz crystal lattice. Given the higher FI density in the FI-rich zones, the elements contained within FIs will exhibit systematically higher concentrations compared to the FI-poor zones. The quartz zones selected for laser ablation were specifically targeted, such that the chosen areas predominately contained primary FIs and were without mineral inclusions. While secondary FIs are frequently present, their salinities are below 1 wt% NaCl eq. (Rajič et al., 2023b), indicating that the detected signal is primarily originating from primary FIs.

For elements displaying higher concentrations in FI-rich zones compared to FI-poor zones, it is assumed that these elements primarily reside within FIs rather than within the quartz crystal lattice. The composition of the primary FIs was determined by a standard charge-balance approach (Allan et al., 2005):

$$mNa + \sum_{i} n_i \times mX_i = mCl \tag{1}$$

where mX_i represents the concentration, in molality (mol/kg) and n_i represents the charge of each cation (X_i) other than Na, respectively. Eq. (1) can be transformed into:

$$mNa = mCl \times \left(1 + \sum_{i} n_i \frac{mX_i}{mNa}\right)^{-1}$$
(2)

where $\left(\frac{mX_i}{mNa}\right)$ stands for molal ratios of all elements of interest relative to sodium. Previously reported bulk salinities of the FIs (Raimbourg et al., 2018; Rajič et al., 2023b) were used to determine the total chlorine content of the inclusions (*mCl*). The molal ratios of all elements relative

to sodium $\left(\frac{mX_i}{mNa}\right)$ were determined by LA-ICP-MS for FI-rich zones. After determining both *mCl* and all molal ratios, the absolute concentration of sodium (*mNa*) and therefore the absolute concentration of all other elements mX_i were determined.

The main problem in the quantification procedure arises if major elements (as described in $\S4.3$. – Na, K, or Ca) in the FIs are below the detection limit of the analyzed sample volume (which comprises both the signal from FIs and host quartz). Sodium and potassium are generally above detection limits, but calcium was at times below the detection limit. Therefore, the semi-quantitative results presented in this work are derived only for analyses where sodium, potassium and calcium concentrations were above detection limit.

3.3. Water-rock ratio calculations

To calculate mass water-rock (W/R) ratios present in the accretionary complexes, mass balance calculations were performed combining the FME whole-rock concentrations as well as recovered FME concentrations of fluid inclusions with theoretical porosity. The porosity of metapelitic rocks at the corresponding depths (>7 km) is expected to be below 5 vol% (Bray and Karig, 1985; Neuzil, 1994; Tsuji et al., 2006). Because the system is conservative and only mass exchange between the solid and fluid is considered, the change in the mass of an element in the solid and the fluid can be defined as

$$\frac{\left|\frac{m_i^{Ff} - m_i^{F,0}}{m_i^{Sf} - m_i^{S,0}}\right| = 1 \tag{3}$$

where $m_i^{F,f}$ is the mass of element *i* in the final fluid, $m_i^{F,0}$ is the mass of element *i* in the initial fluid, $m_i^{S,f}$ is the mass of element *i* in the final solid, and $m_i^{S,0}$ is the mass of element *i* in the initial solid, where final and initial state refer to higher and lower temperature conditions of the present study, respectively. For both the solid and the fluid, the mass of element *i* can be expressed as a concentration by

$$m_{Total}^{j} \cdot c_{i}^{j} = m_{i}^{j} \tag{4}$$

where m_{Total}^{j} is the total mass of phase *j* (fluid or solid), c_{i}^{j} is the concentration of element *i* in phase *j*, and m_{i}^{j} is the mass of element *i* in phase

j. Assuming that the total mass of the fluid and solid remains approximately constant at both final and initial stages, a valid approximation in conservative systems when considering trace elements, the concentrations of element *i* in the solid and the fluid can be expressed as

$$\frac{\left(c_i^{Ff} - c_i^{F,0}\right) m_{Total}^F}{\left(c_i^{Sf} - c_i^{S,0}\right) m_{Total}^S} = 1$$
(5)

Finally, the concentrations of element i in the solid and the fluid are then related to the mass solid/fluid ratio with the following equation:

$$\frac{\left|\left(c_{i}^{Ff}-c_{i}^{F,0}\right)\right|}{\left(c_{i}^{Sf}-c_{i}^{S,0}\right)}\right|=\frac{m_{Total}^{S}}{m_{Total}^{F}}$$
(6)

4. Results

4.1. Whole-rock trace element composition

Whole-rock trace element concentration data of FME of interest are presented in Table 1 and the concentrations are plotted as a function of peak-metamorphic temperatures in Fig. 5. The whole set of analyzed major and trace elements are reported in the Table A.2.

In the Kodiak complex, among all samples, lithium and boron concentrations range from 40 to 70 μ g/g and from 60 to 130 μ g/g, respectively (Fig. 5a-b). Concentrations of rubidium, strontium and barium range from 60 to 120 μ g/g, 110 to 280 μ g/g, and 560 to 1300 μ g/g, g, respectively (Fig. 5c, d, e). Cesium concentrations are in the range of 3 to 7 μ g/g (Fig. 5f; Table 1). In metapelites from the Shimanto Belt, lithium concentrations range from 50 to 70 μ g/g in HTM and from 30 to 40 μ g/g in FM (Fig. 5a; Table 1). Boron concentrations range from 160 to 200 μ g/g in HTM and from 60 to 100 μ g/g in FM (Fig. 5b). Concentrations of rubidium, strontium and barium are relatively consistent for the two units, ranging from 100 to 180 μ g/g, 110 to 150 μ g/g, and 430 to 700 μ g/g, respectively, for all examined samples (Fig. 5c, d, e). Cesium concentrations are 9 to 11 μ g/g in HTM and 5 to 6 μ g/g in FM (Fig. 5f).

4.2. Distribution of FME among mineral phases

The average FME concentrations and standard deviation of mineral phases in each sample are reported in Table 2 and concentrations for

Table 1

Whole-rock trace element composition of studied metapelites from the Kodiak accretionary complex and the Shimanto Belt. Raman Spectroscopy of carbonaceous material (RSCM), K₂O and MgO results are from Rajič et al. (2023b).

| | RSCM | K ₂ O | MgO | В | Li | Rb | Sr | Cs | Ва | B/ K2O | Li/ K ₂ O | Rb∕ K₂O | Sr/ K ₂ O | Cs∕ K₂O | Ba/ K ₂ O | Li/ MgO | B/ Li | Rb/ Cs |
|----------|--------|------------------|------|-----|-----|-------|-----|------|------|-----------|-------------------------|------------|-------------------------|------------|-------------------------|------------|----------|-----------|
| Sample | T (°C) | wt | wt% | µg∕ | µg∕ | µg∕g | µg∕ | µg∕ | µg/g | | | | | | | | | |
| | | % | | g | g | | g | g | | | | | | | | | | |
| Kodiak | | | | | | | | | | | | | | | | | | |
| KO17B | 256 | 1.48 | 1.94 | 64 | 39 | 59.7 | 110 | 4.2 | 555 | 43.2 | 26.4 | 40.3 | 74.3 | 2.8 | 375.0 | 20.1 | 1.6 | 14.2 |
| KO17C | 256 | 2.84 | 2.26 | 118 | 40 | 120.5 | 110 | 7.1 | 1125 | 41.6 | 14.1 | 42.4 | 38.7 | 2.5 | 396.1 | 17.7 | 3.0 | 17.0 |
| KO17D | 258 | 2.37 | 2.08 | 78 | 57 | 96.3 | 280 | 3.8 | 1015 | 32.9 | 24.1 | 40.6 | 118.1 | 1.6 | 428.3 | 27.4 | 1.4 | 25.3 |
| KO17E | 240 | 2.62 | 2.18 | 139 | 55 | 105 | 190 | 5.8 | 978 | 53.1 | 21.0 | 40.1 | 72.5 | 2.2 | 373.3 | 25.2 | 2.5 | 18.1 |
| KO33 | 333 | 1.94 | 2.83 | 115 | 52 | 93.1 | 130 | 4.2 | 1300 | 59.3 | 26.8 | 48.0 | 67.0 | 2.2 | 670.1 | 18.4 | 2.2 | 22.2 |
| KO36 | 330 | 3.29 | 3.08 | 90 | 51 | 77.1 | 260 | 3.4 | 950 | 27.4 | 15.5 | 23.4 | 79.0 | 1.0 | 288.8 | 16.6 | 1.8 | 22.7 |
| KO38 | 340 | 3.61 | 3.19 | 127 | 70 | 112.0 | 120 | 5.4 | 1195 | 35.2 | 19.4 | 31.0 | 33.2 | 1.5 | 331.0 | 21.9 | 1.8 | 20.7 |
| KO11 | 300 | 2.82 | 3.29 | 122 | 54 | 83.4 | 200 | 3.7 | 923 | 43.3 | 19.2 | 29.6 | 70.9 | 1.3 | 327.3 | 16.4 | 2.3 | 22.5 |
| KO12B | 348 | 2.63 | 2.85 | 112 | 56 | 81.1 | 180 | 3.8 | 859 | 42.6 | 21.3 | 30.8 | 68.4 | 1.4 | 326.6 | 19.7 | 2.0 | 21.3 |
| KO45C | 345 | 2.63 | 2.85 | 78 | 51 | 68.5 | 230 | 3.5 | 818 | 29.7 | 19.4 | 26.1 | 87.5 | 1.3 | 311.0 | 17.9 | 1.5 | 19.6 |
| Shimanto | | | | | | | | | | | | | | | | | | |
| HN64 | 238 | 3.29 | 1.98 | 195 | 68 | 164.5 | 110 | 10.5 | 704 | 59.3 | 20.7 | 50.0 | 33.4 | 3.2 | 214.0 | 34.3 | 2.9 | 15.7 |
| HN85 | 242 | 1.44 | 1.49 | 155 | 53 | 179.0 | 120 | 9.0 | 664 | 107.6 | 36.8 | 124.3 | 83.3 | 6.3 | 461.1 | 35.6 | 2.9 | 19.9 |
| HN299 | 228 | 2.80 | 1.80 | 167 | 54 | 167.5 | 150 | 10.1 | 655 | 59.6 | 19.3 | 59.8 | 53.6 | 3.6 | 233.9 | 30.0 | 3.1 | 16.6 |
| 18NOB23 | 341 | 3.04 | 2.44 | 97 | 44 | 106.0 | 110 | 4.9 | 430 | 31.9 | 14.5 | 34.9 | 36.2 | 1.6 | 141.5 | 18.0 | 2.2 | 21.6 |
| HN285 | 321 | 3.08 | 2.07 | 63 | 26 | 182.0 | 130 | 6.2 | 664 | 20.5 | 8.4 | 59.1 | 42.2 | 2.0 | 215.6 | 12.6 | 2.4 | 29.4 |
| HN243 | 335 | 3.00 | 1.74 | 65 | 28 | 144.5 | 110 | 5.4 | 531 | 21.7 | 9.3 | 48.2 | 36.7 | 1.8 | 177.0 | 16.1 | 2.3 | 26.8 |



Fig. 5. Whole-rock FME concentrations in metapelites from the Shimanto Belt and the Kodiak accretionary complex as a function of peak-metamorphic temperatures: (a) Li; (b) B; (c) Rb; (d) Sr; (e) Ba; and (f) Cs.

illite and chlorite are also presented in Fig. 6. All analyses are presented in Table A.4.

Lithium. Lithium is mainly hosted in chlorite and illite (Fig. 6), and to a lesser extent in quartz from the veins (Raimbourg et al., 2021). In the KLB lithium concentrations in chlorite are 240 \pm 20 $\mu g/g$ and 50 \pm 20 μ g/g in illite. In the KCB, chlorite has a lithium concentration of 250 \pm 90 μ g/g (Fig. 6a) and illite has a lithium concentration of 30 \pm 20 μ g/g (Fig. 6b). In the Kodiak complex, chlorite shows retention or even slight lithium enrichment as temperature increases, whereas illite is characterized by comparable lithium concentrations (Fig. 6b). In the Shimanto Belt, lithium concentrations in chlorite are 320 \pm 40 $\mu g/g$ and 140 \pm 20 μ g/g for samples from HTM and FM, respectively, and in illite lithium concentrations are 90 \pm 70 $\mu g/g$ and 20 \pm 20 $\mu g/g$ for HTM and FM, respectively (Fig. 6b). In the Shimanto Belt, the lithium concentration decreases in both chlorite and illite as temperature increases (Fig. 6; Table 2). The vein quartz contains Li in concentrations from $<30 \,\mu g/g$ in veins from KLB and $< 6 \,\mu g/g$ in KCB, $< 110 \,\mu g/g$ in HTM down to $< 4 \,\mu g/g$ g in FM (Raimbourg et al., 2021). Calcite from the veins contains $<1 \mu g/$ g of lithium, with the majority of analyses being below detection limit $(<0.1 \ \mu g/g; Table 2).$

Boron. Boron is dominantly carried by illite and to a lesser extent by chlorite and albite. In the Kodiak complex, illite is characterized by boron concentrations of 140 \pm 50 $\mu g/g$ and 210 \pm 40 $\mu g/g$ for KLB and KCB, respectively. Chlorite from the Kodiak complex has boron concentrations of 10 \pm 5 $\mu g/g$ and 40 \pm 30 $\mu g/g$ for KLB and KCB, respectively (Fig. 6a). In the Shimanto Belt, boron concentrations in illite are 260 \pm 60 $\mu g/g$ and 150 \pm 70 $\mu g/g$ for samples from HTM and FM, respectively (Fig. 6b). Chlorite in units from the Shimanto Belt has low boron concentrations of 40 \pm 30 $\mu g/g$ and 5 \pm 3 $\mu g/g$ in HTM and FM, respectively (Fig. 6a). Albite from the KLB has boron concentrations of $20\pm10~\mu\text{g/g},$ whereas boron concentrations in albite from HTM are 50 \pm 10 $\mu g/g$ (Table 2). The behavior of boron is similar to that of lithium in illite and chlorite. In the Kodiak complex, boron is retained with similar concentrations between lower- and higher-grade illite and chlorite, whereas in the Shimanto Belt boron concentrations decrease in both illite and chlorite (Fig. 6a-b). Calcite in veins contains boron concentrations of up to 15 μ g/g, yet the majority of analyses ranges from below detection limit (<1 μ g/g) to 5 μ g/g (Table 2).

LILE (Rb, Sr, Cs, and Ba). Major carrier minerals of large ion-

lithophile elements are illite (Fig. 6b) and albite (Table 2). Rubidium concentrations in illite from the Kodiak complex are 190 \pm 50 $\mu g/g$ and $320 \pm 30 \ \mu\text{g/g}$ in KLB and KCB, respectively (Fig. 6b). Illite from the Shimanto Belt shows slightly higher rubidium concentrations than in the Kodiak complex, with concentrations of 320 \pm 50 $\mu g/g$ and 400 \pm 40 μ g/g in illite from HTM and FM, respectively (Fig. 6b). Illite has strontium concentrations of 90 \pm 40 μ g/g in the KLB, 70 \pm 40 μ g/g in the KCB, 60 \pm 40 $\mu g/g$ in the HTM, and 30 \pm 10 $\mu g/g$ in the FM (Fig. 6b). Illite from KLB has barium concentrations of 1500 \pm 400 $\mu g/g$ and illite from the KCB has an average barium concentration of $2300 \pm 300 \,\mu\text{g/g}$ (Fig. 6b). The average barium concentration in illite from HTM is 1100 \pm 200 µg/g and illite from FM has a barium concentration of 1400 \pm 100 μ g/g (Fig. 6b). Cesium concentrations in illite are 8 \pm 3 μ g/g from KLB, $17\pm3\,\mu\text{g/g}$ from KCB, $18\pm5\,\mu\text{g/g}$ in samples from HTM, and 15 \pm 3 µg/g from FM (Fig. 6b). Albite hosts LILE as well, but due to its small grain size only analyses on the lower-grade samples were performed. In samples from both KLB and HTM, albite contains low rubidium concentrations of $20 \pm 20 \ \mu\text{g/g}$ and $40 \pm 20 \ \mu\text{g/g}$, respectively (Table 2). Measured concentrations of strontium are 240 \pm 220 µg/g in KLB and $150 \pm 80 \,\mu$ g/g in HTM, whereas barium concentrations are 200 ± 100 μ g/g in KLB and 120 \pm 90 μ g/g in HTM (Table 2). The lowest concentrations among LILE are for cesium, which are ${<}2\,\mu\text{g/g}$ in KLB and 3 ± 2 ug/g in HTM.

Calcite from the veins contains no significant rubidium or cesium concentrations (<10% of the measurements have concentrations above the detection limit; concentrations <1 μ g/g). Barium concentrations in calcite range from 1 to 45 μ g/g (average 9 μ g/g), whereas strontium concentrations are the highest in calcite among LILE, ranging from 0.16 to 1.06 wt% (Table 2).

4.3. Major and trace elements of fluid inclusion populations

A total of 393 LA-ICP-MS spot analyses were performed for FI-rich quartz domains and 208 for FI-poor quartz (Table A.5) in order to determine which elements are hosted within FIs and which ones are hosted in the quartz crystal lattice. Estimated concentrations of Be, Ni, Zn, As, Y, Mo, Sn, Sb, W, Pb, and Bi are below or very close to the detection limit without any systematic difference between FI-poor and FI-rich zones. FI-rich zones are characterized by systematically higher

Table 2

The average concentrations and standard deviation of trace elements in chlorite (chl), illite (ill), albite (ab), and calcite (cc) analyzed by LA-ICP-MS.

| | # | Phase | | Li | В | V | Ni | Cu | Zn | As | Rb | Sr | Nb | Sn | Cs | Ва |
|---------------|-----------|-------|------------|------|----------|----------|----------|------|------|------|----------|------|------|--------|---------|-----------|
| | | | | µg∕g | µg∕g | µg∕g | µg∕g | µg∕g | µg∕g | µg∕g | µg∕g | µg∕g | µg∕g | µg∕g | µg∕g | µg∕g |
| Kodiak Landw | ard Belt | | | | | | | | | | | | | | | |
| KO17A | 34 | ill | Av. | 70 | 150 | 290 | 60 | 30 | 100 | 30 | 190 | 100 | 8 | 4 | 9 | 1400 |
| | | | SD | 40 | 80 | 80 | 30 | 20 | 60 | 20 | 50 | 300 | 8 | 2 | 3 | 400 |
| | 15 | ab | Av. | 5 | 20 | 10 | 60 | 10 | 10 | - | 20 | 240 | - | - | - | 200 |
| | | | SD | 2 | 10 | 10 | 20 | 8 | 10 | - | 20 | 220 | - | - | - | 100 |
| KO17E | 26 | ill | Av. | 70 | 150 | 300 | 60 | 30 | 100 | 30 | 200 | 100 | 4 | 3 | 8 | 1600 |
| | | | SD | 40 | 30 | 60 | 50 | 20 | 90 | 20 | 30 | 90 | 4 | 1 | 2 | 300 |
| | 6 | chl | Av. | 190 | 20 | 150 | 140 | 25 | 300 | 7 | 20 | 20 | 1 | 2 | 2 | 60 |
| K017H | 15 | | SD | 50 | 10 | 50 | 40 | 8 | 100 | 4 | 10 | 9 | 1 | 1 | 1 | 20 |
| | 15 | cc | Av. | 0.15 | 1 | - | - | 0.4 | 1.8 | 0.19 | 0.04 | 6200 | - | - | - | 21 |
| | | | 5D | 0.07 | 0.4 | - | - | 0.5 | 0.6 | 0.07 | 0.02 | 900 | - | - | - | 4 |
| Kodiak Centra | l Rolt | | | | | | | | | | | | | | | |
| KO29A | 24 | ill | Av | 100 | 210 | 350 | 90 | 50 | 130 | 9 | 310 | 60 | 1 | 6 | 17 | 2110 |
| ROEM | 2. | | SD | 60 | 80 | 60 | 50 | 20 | 110 | 3 | 40 | 20 | 0 | 3 | 3 | 330 |
| | 27 | chl | Av. | 210 | 60 | 150 | 170 | 80 | 340 | 10 | 20 | 6 | 1 | 3 | 2 | 16 |
| | | | SD | 80 | 30 | 60 | 70 | 30 | 140 | 4 | 10 | 3 | 1 | 1 | 1 | 9 |
| KO33C | 24 | ill | Av. | 70 | 190 | 420 | 70 | 40 | 140 | 7 | 290 | 60 | 2 | 4 | 14 | 2270 |
| | | | SD | 60 | 30 | 50 | 60 | 60 | 120 | 1 | 40 | 40 | 5 | 2 | 3 | 290 |
| | 23 | chl | Av. | 220 | 20 | 140 | 210 | 30 | 470 | 3 | 1 | 4 | 0 | 1 | 2 | 2 |
| | | | SD | 80 | 10 | 40 | 90 | 20 | 180 | 1 | 1 | 3 | 0 | 0 | 1 | 2 |
| KO38 | 23 | ill | Av. | 80 | 190 | 490 | 90 | 20 | 140 | 10 | 280 | 80 | 2 | 3 | 15 | 2140 |
| | | | SD | 60 | 30 | 60 | 70 | 20 | 110 | 10 | 50 | 20 | 4 | 1 | 3 | 400 |
| | 19 | chl | Av. | 170 | 40 | 200 | 160 | 30 | 310 | 10 | 3 | 8 | 0 | 1 | 4 | 1 |
| | | | SD | 70 | 20 | 50 | 50 | 30 | 100 | 10 | 3 | 6 | 0 | 1 | 2 | 1 |
| KO32A | 13 | cc | Av. | 0.2 | 1.4 | - | - | 0.1 | 1.3 | 0.3 | 0.04 | 7700 | - | - | 0.06 | 3 |
| | | | SD | 0.1 | 0.7 | - | - | 0.1 | 0.7 | 0.1 | | 1190 | - | - | 0.03 | 1 |
| KO32B | 14 | cc | Av. | 0.9 | 3 | - | - | 0.3 | 2 | 0.3 | 0.1 | 8150 | - | - | 0.1 | 6 |
| | | | SD | 0.8 | 2 | - | - | 0.3 | 1.8 | 0.1 | 0.03 | 1320 | - | - | 0.1 | 2 |
| | | | | | | | | | | | | | | | | |
| Hyuga Tecton | ic Mélang | ge | | | | | | | | | | | | _ | | |
| HN299A | 46 | 111 | Av. | 100 | 250 | 220 | 50 | 49 | 87 | 40 | 310 | 110 | 6 | 5 | 16 | 1080 |
| | 47 | -1-1 | SD | 60 | 50 | 50 | 40 | 27 | 52 | 30 | 50 | 390 | 9 | 2 | 4 | 210 |
| | 47 | ciii | AV. | 230 | 40 | 20 | 80 20 | 23 | 180 | 13 | 40 | 20 | 3 | 3 1 | 4 | 20 |
| | 0 | ab | 3D Av | 90 | 40 54 | 30 | 30 | 24 | 08 | 14 | 30 40 | 150 | 4 | 1 | 2 | 20 120 |
| | 0 | aD | AV. | - | 5 | 30 10 | - | 24 | - | 5 | 20 | 130 | 4 | 2 | 3 2 | 00 |
| HN200C | 26 | -111 | Δv | - 70 | 240 | 180 | - | 320 | - | 290 | 20 | 80 | 7 | 5 | 2 18 | 1080 |
| 11112556 | 20 | 111 | SD | 40 | 50 | 40 | 490 | 470 | 100 | 160 | 50 | 80 | 50 | 3 | 5 | 230 |
| | 21 | chl | Av | 250 | 40 | 80 | 70 | 14 | 230 | 11 | 20 | 14 | 1 | 4 | 5 | 40 |
| | | | SD | 70 | 30 | 20 | 20 | 8 | 80 | 5 | 10 | 7 | 2 | 1 | 2 | 20 |
| HN65 | 17 | ill | Av. | 140 | 260 | 230 | 20 | 30 | 50 | 11 | 320 | 80 | 7 | 5 | 21 | 980 |
| | | | SD | 80 | 70 | 40 | 10 | 20 | 30 | 7 | 60 | 60 | 8 | 2 | 7 | 190 |
| | 11 | chl | Av. | 440 | 130 | 100 | 20 | 17 | 50 | 7 | 150 | 12 | 5 | 3 | 7 | 280 |
| | | | SD | 360 | 50 | 30 | 20 | 8 | 30 | 4 | 40 | 6 | 9 | 1 | 2 | 100 |
| | | | | | | | | | | | | | | | | |
| Foliated Moro | tsuka | | | | | | | | | | | | | | | |
| HN145 | 52 | ill | Av. | 60 | 180 | 220 | 40 | 140 | 120 | 40 | 380 | 40 | 10 | 6 | 17 | 1340 |
| | | | SD | 20 | 70 | 20 | 20 | 80 | 60 | 30 | 20 | 10 | 20 | 3 | 4 | 100 |
| | 4 | chl | Av. | 80 | 80 | 110 | 110 | 370 | 380 | 20 | - | - | 3 | 4 | - | - |
| | | | SD | 40 | 40 | 30 | 50 | 170 | 190 | 10 | - | - | 4 | 2 | - | - |
| HN243 | 29 | ill | Av. | 50 | 140 | 190 | 70 | 40 | 50 | 9 | 410 | 30 | 1 | 5 | 15 | 1280 |
| | | | SD | 20 | 70 | 60 | 30 | 20 | 50 | 4 | 110 | 10 | 0 | 2 | 5 | 370 |
| | 28 | chl | Av. | 120 | 70 | 110 | 70 | 50 | 290 | 20 | 8 | 3 | 1 | 3 | 3 | 5 |
| | | | SD | 50 | 20 | 30 | 30 | 30 | 130 | 10 | 5 | 2 | 1 | 1 | 2 | 3 |
| HN285 | 35 | ill | Av. | 12 | 110 | 50 | 5 | 4 | 26 | 3 | 410 | 30 | 10 | 5 | 13 | 1340 |
| | 0.2 | | SD | 2 | 10 | 20 | 2 | 2 | 4 | 1 | 20 | 10 | 13 | 1 | 2 | 110 |
| | 33 | chl | Av. | 140 | 7 | 80 | 110 | 50 | 510 | 8 | 2 | 3 | 0 | 1 | 1 | 6 |
| NOP OOP | 15 | | SD | 20 | 5 | 10 | 30 | 30 | 70 | 9 | 2 | 1 | U | U | 1 | 2 |
| NOB_23B | 15 | cc | AV. | 0.5 | 1.6 | - | - | 0.5 | 2 | 0.4 | 0.3 | 4/40 | - | - | 0.05 | 3 1 |
| | | | 5D | 0.2 | 0.8 | - | - | 0.4 | 0.5 | 0.2 | 0.2 | 1530 | - | - | 0.02 | 1 |

concentrations of B, Na, K, Ca, Mg, Fe, Rb, Sr, Cs, and Ba in comparison to FI-poor zones (Fig. A.1). An example of which elements are found in the FIs is presented in form of box plots for FI-rich zones and adjacent FI-poor zones (Fig. 7).

fluid evolution studies due to inability to detect whether they are concentrated in the quartz crystal lattice or in both quartz and FIs.

Lithium, aluminum and titanium show no systematic difference between the two zones (Fig. 7). When ablating multiple FI instead of individual FI, subtracting the host mineral from the signal becomes challenging. Thus, Li, Al, and Ti are not generally considered for the For elements present in the FIs, element ratios on a mole basis are in the range of: Ca/Na – 0.06 to 13.23; B/Na – 0.02 to 0.57; K/Na – 0.01 to 5.69; Rb/Na – 2.2×10^{-5} to 0.05; Sr/Na – 0.0005 to 0.09; Ba/Na – 1.1×10^{-8} to 5.3×10^{-6} ; and Cs/Na – 1.2×10^{-5} to 0.002 (Table 3). Sodium, calcium and potassium are major elements in FIs. Their concentrations have a large range: Na – 330 to 9700 µg/g; Ca – 360 to 10,800 µg/g; and



Fig. 6. Box plots of FME concentrations as a function of peak-metamorphic temperatures of (a) chlorite (Li and B) and (b) illite (Li, B, Rb, Sr, Cs and Ba), from the Kodiak complex and the Shimanto Belt, acquired by LA-ICP-MS. Arrows indicate the trend in chemical evolution of fluids as temperature rises, with orange representing Kodiak and gray representing Shimanto.

K – 10 to 10,600 μg/g (Table 3; Fig. 8). Potassium is the predominant cation in FIs of lower-grade veins, whereas Ca and Na have higher concentrations in FIs of higher-grade veins. Magnesium and iron are minor elements with ranges of 10 to 1200 μg/g and 20 to 1500 μg/g, respectively, with noticeably higher concentrations in higher-grade FIs (Fig. 8). The ranges of trace elements are: B – 4 to 870 μg/g; Rb – 0.4 to 40 μg/g; Sr – 10 to 380 μg/g; Cs – 0.1 to 12 μg/g; and Ba – 1 to 1800 μg/g. Trace elements show a narrower concentration range than major cations for the examined veins of each unit (Table 3; Fig. 8).

5. Discussion

5.1. Fluid-mobile elements redistribution in metasediments upon metamorphism

As mentioned in the § 1, the FME behavior is highly controlled by the geothermal gradient of the subduction zone, which in turn controls the occurrence of metamorphic reactions (e.g., Bebout et al., 2013). However, metasedimentary rocks from Kodiak and Shimanto display a similarly high geothermal gradient (20–30 °C/km) and a lack of major dehydration reactions in the range 230–350 °C (Rajič et al., 2023b). Despite experiencing similar peak pressure-temperature conditions and mineral reactions in both complexes, the FME behavior differs between two localities. Consequently, in order to elucidate the contrasting behavior of FME between Kodiak and Shimanto, alternative explanations must be considered.

5.1.1. FME budget in the solid matrix

In contrast to whole rock major element concentrations that do not

vary when temperature increases from 230 to 350 °C (Rajič et al., 2023b), some of the FME concentrations vary across this temperature range (Fig. 5). In samples from the Kodiak complex, excluding the partial loss of cesium, all other FME are retained upon increasing temperature and the concentrations in higher-grade and the lower-grade samples are similar. Conversely, in the Shimanto Belt, FME show significant and systematic loss as temperature increases (Fig. 5). The different behaviors between the Kodiak complex and the Shimanto Belt can be illustrated by devolatilization indicators that have been extensively applied to other metasedimentary units (e.g., Bebout, 1995; Bebout et al., 2013), such as Rb/Cs or B/Li. Both Rb/Cs or B/Li are scattered among samples from the Kodiak complex, whereas in the Shimanto Belt Rb/Cs ratio increases and B/Li ratio systematically decreases (Fig. 9; Table 1). In terms of mineral reactions, the two opposing trends for FME budget can be accounted for by (i) changes in FME concentrations in the major minerals in the pelitic matrix, and (ii) mineral reactions and changes in phase proportions as a function of temperature (both phenomena being connected to the extent of fluidrock interactions).

As noted by several studies, the FME budget in subducted lithologies is highly controlled by the abundance of phyllosilicates and their reactions during burial (e.g., Barnes et al., 2019a; Bebout et al., 1999, 2007, 2013; Busigny et al., 2003; Marschall et al., 2006). In the samples from this study, the main FME carriers are illite (boron, rubidium, barium, cesium \pm lithium, strontium) and chlorite (lithium \pm boron), with lower concentrations of LILE in albite (strontium \pm rubidium, barium; Table 2; Fig. 10). Upon burial and increase in P and T, the minerals present in the pelitic matrix are affected by three main mineral reactions that affect their composition and/or abundance: illite



Fig. 7. The concentrations of B, Na, K, Sr, Ba, Li, Al, and Ti in quartz obtained with the LA-ICP-MS. For each unit, colored box plots correspond to the FI-rich quartz, whereas adjacent white box plots correspond to the FI-poor quartz. Note that FI-rich quartz is enriched in B, Na, K, Sr and Ba compared to FI-poor quartz, whereas measured concentrations of Li, Ti and Al are fairly equal in both zones.

recrystallization, chlorite crystallization and the dissolution of albite (Rajič et al., 2023b).

To examine the effects of these mineralogical reactions on FME budget, we performed mass balance calculations using the modal proportions of major minerals (XRD results; Rajič et al., 2023b) and the FME concentrations of each major mineral (Fig. 10). The calculated concentrations of lithium, rubidium, cesium, and barium are in good agreement (within ~20% difference) with whole-rock concentrations. In contrast, for boron and strontium, the calculated concentrations are systematically underestimated for most of samples (from ~25% to ~50% and from ~35% to ~75%, for B and Sr, respectively, see Fig. 10). The following discussion is therefore restricted to the elements for which the budget can be consistently estimated (Li, Rb, Cs and Ba), while B and Sr are discussed in § 5.1.2.

In the Kodiak complex, lithium concentration in chlorite, the main Li carrier, remains statistically identical as temperature increases (Fig. 6). Therefore, in spite of minor modal variations due to chlorite crystallization, the whole-rock stock of Li remains constant upon T increase (Fig. 5a). In contrast, in the Shimanto Belt, metamorphic illite and chlorite show significant depletion in lithium as temperature increases (Fig. 6), which result in the whole-rock decrease in Li upon T increase (Fig. 5a).

In both localities, recrystallized illite has higher concentrations of rubidium and barium in comparison to lower-grade illite (Fig. 6). As discussed in \S 5.2, rubidium concentrations decrease in the fluid as temperature increases. Thus, newly formed illite consumes Rb from the fluid and possibly from dissolved fraction of albite. As a result, there are limited whole-rock variations in Ba and Rb concentrations upon T increase (Fig. 5c, e).

Finally, cesium shows contrasted behavior between the two geological examples. In Kodiak, similarly to Rb and Ba, cesium is retained in recrystallized illite and the whole-rock stock remains constant as temperature increases. In the Shimanto Belt example, cesium concentrations show slight decrease in illite as temperature increases, thus resulting in whole-rock decrease as well (Fig. 5f, 6).

In summary, in most cases (all elements for Kodiak, Rb and Ba for the Shimanto Belt), elements are redistributed between mineral phases upon temperature increase, without affecting the whole-rock budget. In contrast, Li and Cs are leached out of the rock in the Shimanto Belt example, as a result of a decreasing concentrations in newly formed minerals.

Consistent with the results of this study, Ramirez et al. (2021) described a similar depletion of FME in scaly fabrics (also known as pressure-solution seams) in metapelites of tectonic mélanges from the

Table 3

Average values and standard deviation of recovered concentrations of major, minor and trace elements in fluid inclusion populations and molal ratios for each studied formation. Only point analyses where Ca was successfully measured are reported.

| μg/g mole mole <t< th=""><th></th></t<> | |
|--|-----------|
| Kodiak Landward Belt Av. 110 4120 120 5550 1030 - 110 6 100 1 190 0.07 0.8 0.4 0.0004 0.008 4.09E-05 3.1 SD 30 980 60 1600 1060 - 100 3 40 0.5 80 0.01 0.4 1.5 0.0003 0.009 2.08E-05 1.4 Min 8 330 30 660 420 - 20 1 50 0.4 30 Max 200 7040 350 9090 6640 - 390 14 190 2.5 360 - < | nole |
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| Foliated Morotsuka | |
| Av. 273 3790 200 2200 6030 230 330 6.5 100 4.6 233 0.19 0.72 1.09 0.0011 0.007 0.0003 5. | 5.32E-07 |
| SD 192 2070 240 2140 2490 360 370 7.4 60 3.1 428 0.08 2.86 1.99 0.0048 0.004 0.0002 8. | 3.60E-07 |
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Fig. 8. Box plots illustrating recovered concentrations of fluid inclusions based on charge-balance approach (for B, Rb, Sr, Cs, Ba, Na, and K) plotted against peakmetamorphic temperatures. Arrows indicate the trend in chemical evolution of fluids as temperature rises, with orange representing Kodiak and gray representing Shimanto.



Fig. 9. (a) Rb/Cs, (b) B/Li and (c) Li/MgO ratios of metapelites from the Kodiak complex and the Shimanto Belt, plotted against peak-metamorphic temperatures.

Shimanto Belt and proposed that elements are remobilized by diffusion along grain boundaries into low, effective normal stress sites and incorporated into vein-forming minerals. However, vein-forming minerals (quartz and calcite) are small reservoirs for most FME, except for the limited incorporation of lithium in quartz (Raimbourg et al., 2021) or strontium in calcite (Table 2). Therefore, for most FME the whole-rock depletion implies that these elements are transferred into a fluid phase and leached out of the rock.



Fig. 10. Modal proportion of mineral phases and distribution of Li, B, Rb, Sr, Cs, and Ba among major mineral phases for selected samples from the Hyuga Tectonic Mélange (HTM), Foliated Morotsuka (FM), Kodiak Landward (KLB) and Central (KCB) Belts. The mass balance calculations of Li, B, Rb, Sr, Cs, and Ba were calculated from their concentrations in the major mineral phases (illite, chlorite, albite, and quartz) combined with XRD results (Rajič et al., 2023b) and normalized to 100% of whole-rock concentrations. Dashed lines represent 20% variations from the whole-rock concentrations.

5.1.2. Additional reservoirs for Sr and B

The mass balance calculations for strontium and boron are inconsistent with the whole-rock concentrations (Fig. 10). Strontium behaves similarly to calcium, and the presence of calcium-bearing phases, which may incorporate large amounts of strontium, can highly influence the strontium budget. In the examined samples, calcite is observed in the host rock as a minor phase. However, due to its very small grain size, FME concentrations of calcite in the pelitic matrix were not obtained and not considered in the mass balance calculations (Fig. 10). The modal fractions of calcite in the pelitic matrix range from ~ 0.5 to $\sim 1.5\%$ (Rajič et al., 2023b). Larger calcite crystals are observed in the veins (Rajič et al., 2023b) and have strontium concentrations of up to 1 wt%

(Table 2). If calcite in the pelitic matrix has similar strontium concentrations to the vein calcite (Table 2), it would correspond to 50–150 μ g/g of strontium on the whole-rock scale, which resolves the strontium underestimation in mass balance calculations (Fig. 10). Thus, the underestimation of the whole-rock strontium budget is attributed to the omission of the minor proportion of calcite in the matrix from the mass balance calculation.

Regarding the discrepancy in boron, several possibilities can be raised, such as boron being stored in organic material (Williams et al., 2001b) or variable concentrations of B in illite as a function of grain size (Clauer et al., 2018; Clauer et al., 2014). First, although kerogen may contain significant boron concentrations (Williams et al., 2001b), the amount of carbonaceous material in metapelites is estimated to be ~ 0.5 wt% (Raimbourg et al., 2017) and therefore it is a second-order reservoir. Second, Clauer et al. (2018) pointed out that illitization progressed via dissolution-reprecipitation rather than by the solid-state substitution. As a consequence, newly precipitated illite is characterized by a larger grain size and lower boron concentrations than the older, smaller



Fig. 11. (a) Ca/Na vs. K/Na molal ratios as well as (b) Rb/Na, (c) Sr/Na, (d) Cs/Na, and (e) Ba/Na molal ratios plotted against B/Na molal ratio of fluid inclusions, for each of studied formation. Arrows on the binary plots (orange – Kodiak; blue – Shimanto) represent the evolution of chemical composition as temperature rises. In (a), the pink points represent the average composition of all decrepitation mounds in individual samples of vein quartz from the Kodiak complex and the Shimanto Belt, acquired by EPMA (Rajič et al., 2023b). Gray areas represent fluid composition in metasedimentary units from the Central Alps (Gauli) determined by LA-ICP-MS on individual fluid inclusions (Rauchenstein-Martinek et al., 2016), whereas red points in (a) represent the chemical composition of fluid inclusions from the Sambagawa Belt determined by applying crush-leach technique (Yoshida et al., 2015). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

grains (Clauer et al., 2018, Clauer et al., 2014). During this study, only sufficiently large grains (exceeding 30 μ m in width) were analyzed by LA-ICP-MS (e.g. larger illite in the contact with the shear band shown in Fig. 3c); additionally, the FME concentration of smaller illite in the pelitic matrix was not measured. Therefore, the underestimation of the B content from individual mineral analyses (Fig. 10) could be explained by the grain size dependence of B concentration in illite.

In conclusion, we propose that strontium was remobilized by the fluid into dilatation sites, where it is stored in newly precipitated calcite. Regarding boron, the same general trend as Li can be observed. In the case of Kodiak, there is a slight increase in illite concentration in boron upon temperature increase. This increase, combined with the dissolution of albite (minor host of B), is consistent with the redistribution of B within the remaining illite population and constant whole-rock concentrations. In contrast, newly formed illite hosts lower proportion of B (Fig. 6) upon temperature increase in the case of Shimanto, consistent with a decrease in whole-rock concentration in B. Therefore, unlike the close-system behavior observed for Kodiak, the FME in high-temperature endmember in the Shimanto Belt was leached out by a circulating fluid.

5.2. Chemical evolution of the fluid and fluid-rock interaction

Bulk-microanalysis of FI populations revealed high variations in the major elements (sodium, potassium, and calcium) composition of the fluid (Figs. 7, 8). However, within a given geological unit, veins have similar chemical composition (Table 3; Fig. 11). Similar results were derived by using a decrepitation method for the same FIs (decrepitation of fluid inclusions occurs as a result of rapid thermal impact, where released material subsequently precipitates around the halos; Kontak, 2004): the proportion of sodium, potassium and calcium is highly variable and the majority of the analyzed evaporite mounds contained calcium as a major cation (Rajič et al., 2023b). All microprobe analyses in the decrepitation mounds show high variability; however, the average ratios in a single sample revealed similar K/Na and higher Ca/Na when compared to the results obtained by LA-ICP-MS (Fig. 11a). A discrepancy in Ca/Na ratios obtained by two methods can be explained by two reasons. First, derived Ca concentrations by LA-ICP-MS may be underestimated due to the analytical limitations as explained in § 3.2.2. Second, during the bulk microanalysis of FI population, multiple FIs were ablated, in contrast to individually analyzed evaporite mounds. This explains higher variability in EPMA point analyses when compared to LA-ICP-MS results, whereas the average composition of samples containing multiple mounds is similar to the LA-ICP-MS results (Fig. 11a).

To the best of our knowledge, other studies examining the chemical composition of FIs in subducted sediments that have experienced peaktemperatures of 230-350 °C are limited. The closest example are the higher-grade metasedimentary and metamafic units from the Sambagawa Belt, Japan, where crush-leach technique was applied to synmetamorphic quartz veins (Yoshida et al., 2015). Their results in regard to K/Na, Ca/Na, and B/Na are consistent with those reported here (Fig. 11a). Another example are fissure veins from chlorite-mica schists in the Central Alps (Gauli), which have experienced sub-greenschist to amphibolite facies metamorphism (Rauchenstein-Martinek et al., 2016) and the chemical composition of individual FIs were measured by LA-ICP-MS (Fig. 11a-e). The absolute concentrations of major elements (sodium, potassium, and calcium) and trace elements (boron, strontium, cesium, and barium) are also in agreement with our results (Rauchenstein-Martinek et al., 2016; Table 3). Similarities in the determined fluid compositions of the two previous studies (Rauchenstein-Martinek et al., 2016; Yoshida et al., 2015) and the results from this study validate the approach of LA-ICP-MS bulk microanalysis of FI populations applied in this study (Fig. 11).

The ratios on a molal basis obtained in the present work, such as K/ Na, Ca/Na, B/Na, Rb/Na, or Ba/Na are on average at least one order of magnitude higher than those of seawater and of the pore fluids of subducting sediments measured during IODP/ODP programs (Fig. 12). Only Cs/Na ratios are similar to the pore fluids (Fig. 12). This indicates that the fluids entrapped in FIs have chemical compositions that are significantly different from the initial pore fluid and that reflect the effect of diagenetic to low-grade metamorphic reactions. Upon burial of clay-rich sediments, the smectite-to-illite transformation is the critical mineral reaction that influences the evolution of the fluid, because smectite-illite contains boron, lithium, and LILE in both the interstitial site and the crystal structure. At temperatures >60 °C, alkali metals (Li, Na, K, and LILE) preferentially go into the fluid phase (Kastner et al., 2014; Seyfried et al., 1998; You and Gieskes, 2001). Most boron stored in clay minerals is released into the fluid phase at temperatures $\sim 150\ ^\circ C$ (Hüpers and Kopf, 2012; Williams et al., 2001a; You et al., 1995). Conversely, the depletion of potassium and rubidium in the fluid is expected when the transformation is completed, as illite incorporates higher concentrations of potassium and rubidium than smectite-illite (Kastner et al., 2014). Thus, major changes in the composition of pore fluids are likely to occur at temperatures <250 °C, with smaller compositional changes upon further temperature increase.

Nevertheless, upon a temperature increase from ~250 to ~350 °C, a few general geochemical trends are observed in veins from both the Kodiak complex and the Shimanto Belt: Ca/Na, B/Na, and Cs/Na ratios increase (Fig. 11a, d), whereas K/Na and Rb/Na ratios decrease as temperature increases (Fig. 11a, b). The Sr/Na ratio remains relatively constant. Only the Ba/Na ratio shows different trends between the two localities, with a slight decrease in the Kodiak complex and a slight increase in the Shimanto Belt with a temperature increase (Fig. 11e). These changes of element ratios as a function of temperature can be explained by examining the chemical evolution of the mineral phases and the mineral reactions in the pelitic matrix (Rajič et al., 2023b). For example, the K/Na ratio decreases as temperature increases because illite becomes more potassium-rich as temperature increases, and the fluid becomes enriched in sodium at higher temperatures related to the dissolution of albite in the pelitic matrix.

Furthermore, the fluid in higher-grade veins from both case studies is characterized by an enrichment in trace elements, and in both localities the increase in FME concentrations are similar (Fig. 8). In the Kodiak complex, there is no systematic loss of FME as temperature increases in the whole-rock (Fig. 5a-f). Conversely, in metapelites from Shimanto, FME show the most significant loss in whole-rock with increasing temperature (Fig. 5a-f). Thus, in both localities the same trends in the fluid were observed as temperature increased, whereas the whole-rock trends are opposite, and therefore the two case studies require different interpretations. The plausible explanations are (i) the difference in the size of fluid and rock reservoirs and (ii) the external influx of fluid in parallel to the observed metamorphic reactions responsible for release of FME from the rock.

5.2.1. Water-rock ratio

In order to explain the chemical evolution of pore fluids from 250 °C to 350 °C, the proportion of the solid and the fluid reservoirs needs to be considered. In metapelitic rocks at the depths >7 km the expected porosity is around 1–5 vol% or lower (Bray and Karig, 1985; Neuzil, 1994; Tsuji et al., 2006), meaning that the mass of pore fluid (for a fluid density of ~ 0.7 g/cm³) is 70–330 times smaller than that of the solid in given mass of a solid (Fig. 13a). Thus, if the system is closed, minor changes in the FME concentrations of the solid caused by interaction with a fluid should have significant impact on the fluid composition. This may be the case in the Kodiak complex, as no significant loss of FME is observed from the solid with increasing temperature (Fig. 5), whereas FIs record an enrichment in FME, especially for boron and cesium (Fig. 8).

In the Kodiak complex, the theoretical mass solid/fluid ratios were combined with measured concentrations of element *i* in the initial solid $(c_i^{s,0})$, initial fluid $(c_i^{F,0})$ and final fluid $(c_i^{F,f})$, allowing for concentra-



Fig. 12. Binary plots of B/Na vs. K/Na, Ca/Na, Rb/Na, Sr/Na, Cs/Na, and Ba/Na molal ratios of fluid inclusions examined in this study by LA-ICP-MS, with the compilation of possible fluid reservoirs involved into chemical evolution of subduction zone fluids in sedimentary units: The seawater composition (Bruland, 1983); The chemistry of pore fluid: IODP 372B/375 Site U1520 (Wallace et al., 2019); IODP 204 Sites 1245 and 1251 (Tréhu et al., 2003); IODP 320/321 (Pälike et al., 2009); Mud volcanoes and hot/cold springs (Barnes et al., 2019b; Dia et al., 1999); Crush-leach results from the Sambagawa Belt (Yoshida et al., 2015); Fluids released during dehydration reactions in serpentinites (Scambelluri et al., 2004a, 2015); Serpentinite mud volcano fluids from the Mariana Forearc (Hulme et al., 2010; Menzies et al., 2022; Mottl et al., 2004).



Fig. 13. The mass fluid-rock ratio calculations for the Kodiak complex and the Shimanto Belt. (a) The mass fluid-rock ratio as a function of the porosity, showing that the fluid reservoir is significantly lower than the solid. Calculations of required decrease in concentrations of (b) B and (c) Cs from the solid in the Kodiak complex, in order to create the measured enrichment of B and Cs in the pore fluid. Adjacent box plots represent the range of measured whole-rock concentrations of boron and cesium in lower- and higher-temperature samples. (d) Boron and (e) cesium concentrations decrease at whole-rock scale (primary y-axis) as a function of temperature in the Shimanto Belt, whereas the fluid shows an enrichment (secondary y-axis). (f) Mass water-rock ratios calculated for boron, cesium, and barium. Red area represents water-rock ratio to the porosity of <10 vol% in the case of closed system. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

tions in the final solid (c_i^{Sf}) to be calculated (Eq. 6). In order to cause an enrichment in boron concentrations in the fluid (from ~110 µg/g to 230–700 µg/g; Table 4), the boron concentration in the solid is required to decrease from ~100 µg/g to ~91 µg/g (Fig. 13b), which represents a < 10% decrease (assuming a porosity of 1–5 vol%). For cesium, in order to increase concentrations in the fluid from ~1 µg/g to 3–7 µg/g, the cesium concentrations in the solid is required to decrease from 5.2 µg/g to 5.1 µg/g (Fig. 13c), which is <2%. Therefore, because of the small volume of the pore fluid reservoir (compared to the solid matrix), the concentration of FME is much more sensitive to rock/fluid exchanges than the solid matrix.

Conversely, in the case of Shimanto, significantly higher amounts of FME are released from the solid (Fig. 13d, e) and therefore the fluid should be drastically enriched in all released elements. However, the enrichment is on the same order as in the fluids from the Kodiak complex (Fig. 8). Thus, another mechanism should be considered, such as the percolation of externally derived fluids, which are responsible for decrease in concentrations of FME in the solid. In order to calculate the required mass of fluid to leach the FME from the solid, the mass waterrock ratio is calculated following eq. (6). Calculated mass water-rock ratios in the Shimanto Belt range from 0.5 to 0.7, 1.2 to 1.7, and 0.7 to 2.2 for boron, cesium, and barium, respectively (Fig. 13f).

5.2.2. Closed vs. open system

Derived mass water-rock ratios in the Foliated Morotsuka (Shimanto Belt) are ~ 30 to ~ 130 times higher than expected if the system was closed and characterized by porosity lower than 5 vol% (Fig. 13f). This implies that a large flux of fluid percolated through the rocks and leached the elements out of the minerals. Two factors can be invoked to explain why fluid percolation was present at Shimanto Belt and was not

observed at Kodiak. The first is the large amount of strain the rocks experienced in the Foliated Morotsuka (Raimbourg et al., 2014), where the pervasive network of shear zones greatly enhanced rock permeability (Cox, 2002, 2010; Wibberley and Shimamoto, 2003). The second is the proximity of large-scale fault zones, which may have acted as pathways for deeper and overpressurized fluid to drive fluid-rock interactions (Faulkner et al., 2010, Faulkner et al., 2003; Herviou and Bonnet, 2023; Saffer and Tobin, 2011). The Foliated Morotsuka is located in the hanging wall of a large-scale fault zone, the Nobeoka Tectonic Line (Kondo et al., 2005), which may have promoted percolation of external fluid through these rocks. For these reasons, we propose that high water-rock ratios in the Foliated Morotsuka are the result of fluid flow from deeper levels that was focused on the structural horizon where strain is the highest. Similar observations have been reported for the Glarus Thrust in the Eastern Helvetic Alps (Abart et al., 2002; Badertscher et al., 2002; Burkhard et al., 1992; Burkhard and Kerrich, 1988; Hürzeler and Abart, 2008). Another crucial observation is that some samples (e.g., HN145) collected from the Foliated Morotsuka are located at substantial distance from the Nobeoka Tectonic Line (see Fig. 1b). This significant distance from the NTL implies a substantial damage zone influenced by the infiltration of fluids originating from the deeper regions of the subduction zone.

In the case of the Kodiak Central Belt, textural evidence around crack-seal veins is consistent with local migration of SiO_2 due to porefluid diffusion of SiO_2 from the pelitic matrix (Fisher and Brantley, 1992; Rajič et al., 2023b) and there is a lack of evidence to support longdistance transport (Fisher et al., 1995; Fisher and Brantley, 2014). In contrast to the Foliated Morotsuka, the Kodiak Central Belt is located in the central part of large turbiditic sequence far from any recognized large-scale faults (Fig. 1).

In the Kodiak complex, trace elements behaved similarly to major

elements (Rajič et al., 2023b): The system was closed at the large (typically >hundreds of meters or kilometers) scale and open at the centimeter scale, where FME were transported from the pelitic matrix towards the vein and the newly formed minerals (illite and chlorite) in contact with the vein (Fig. 14). Conversely, there is a discrepancy between the behavior of major and trace elements during prograde metamorphism in the Shimanto Belt. No systematic changes have been reported for major elements in HTM and FM, which indicates a closed system (Rajič et al., 2023b). However, fluid-mobile elements examined during this study show a systematic decrease as temperature increases, requiring high mass water-rock ratio to leach these elements out of the rocks.

Thus, we propose that the contrasting behavior between the closed system (with respect to FME) observed in the Alaskan terranes and the open system observed in the Japanese terranes is the result of their internal strain and their proximity to large-scale faults, which concentrated the fluid flow (Fig. 14).

5.3. Implications for fluid circulation in subduction zones

This study shows that boron and cesium are the most sensitive tracers of fluid-mediated processes in subduction zones, which is consistent with previous use of boron and cesium as tracers for prograde dehydration of altered oceanic crust (Angiboust et al., 2014; Bebout, 1991; Bebout et al., 1993, 2007; Lafay et al., 2013; Marschall et al., 2006; Moran et al., 1992) or serpentinites (Deschamps et al., 2010, 2011, 2013; Hattori and Guillot, 2007; Lafay et al., 2013; Li and Lee, 2006; Scambelluri et al., 2004a, 2004b). Furthermore, derived water-rock ratios suggest that high quantities of fluid circulated through the Foliated Morotsuka, where such fluids may be expelled through the subduction zone forearc along permeable horizons. Additionally, despite the absence of dehydration reactions within analyzed samples from the Shimanto Belt (Rajič et al., 2023b), there is a noticeable loss of FME. This observation is consistent with the high sensitivity of the FME not just to water loss, but also to the volume of fluids traversing through rocks. Consequently, assesing the FME evolution of both the solid and the fluid has a great potential to characterize rocks undergoing substantial fluid influx, such as metasomatic rocks formed along the subduction interface (Angiboust et al., 2014; Penniston-Dorland and Harvey, 2023).

In addition, similarities are observed between boron and cesium concentrations (or B/Na and Cs/Na ratios) in FIs acquired during this study and the chemical composition of mud volcano fluids from Trinidad (Dia et al., 1999) or hot and cold springs from New Zealand (Barnes et al., 2019b). The origin of spring fluids from New Zealand have been interpreted to be derived from dehydrated subducted sediments based on high δ^{18} O and δ D values in fluids flowing through the springs (Barnes et al., 2019b; Giggenbach et al., 1993; Reves et al., 2010). The fluids from mud volcanoes and springs (Dia et al., 1999; Barnes et al., 2019b) are characterized by a composition that is between that of the seawater and fluids from this study (Fig. 12), confirming the mixing of seawater that is percolating downwards and metamorphic fluids that are percolating upwards, as proposed for the Hikurangi accretionary prism (Reyes et al., 2010). Our study further confirms that compositionally altered pore fluids escaped from subducting plate and travel along zones with high permeability back to the seafloor.

However, the fluid composition preserved in FIs examined in this study closely resembles the potential fluid sources found in deeply subducted lithologies. In this context, the fluid that traverses the Foliated Morotsuka may originate from any of the H₂O-bearing lithologies that were subducted to significant depths, including sediments, altered oceanic crust or serpentinites. The major dehydration reactions in these lithologies and subsequent release of water occurs deeper than the depths at which the units from this study were buried (Aizawa et al., 1999; Deschamps et al., 2013; John et al., 2011; Kameda et al., 2017; Peacock, 1993; Scambelluri et al., 2004b; Zack et al., 2003). When

comparing the fluid compositions reported in this study with that of metasedimentary and metamafic units from the Sambagawa Belt (subducted to depths between 15 and 60 km; Yoshida et al., 2015), the fluid composition is relatively similar (Fig. 12). Dehydration reactions in altered oceanic crust may also contribute to the upwelling fluids. A good example of a release of such fluids from altered oceanic crust at temperatures <300 °C are serpentinite mud volcano fluids from the Mariana forearc (Hulme et al., 2010; Menzies et al., 2022; Mottl et al., 2004). However, serpentinite mud volcano fluids exhibit a composition similar to seawater (Fig. 12), implying their limited relevance in this study. Another significant source of fluids are serpentinites, with the major dehydration reactions occurring at depths between 40 and 60 km (Scambelluri et al., 2004a, 2015). Upon comparing the chemical composition of fluid from this study with those released during major dehydration reactions in serpentinites (Scambelluri et al., 2004a, 2015), only slight depletion in boron relative to our results is observed (Fig. 12). Therefore, gaining a better understanding of which subducted lithology and which specific reaction primarily contribute to the fluids percolating through the Foliated Morotsuka requires further investigation.

6. Conclusions

This study provides an insight into the behavior of fluid-mobile elements by examining whole-rock concentrations, concentrations in individual minerals, and fluid inclusions from syn-subduction veins in the Kodiak complex, Alaska and the Shimanto Belt, Japan. The Kodiak complex exhibits closed system behavior and retention of all FME at temperatures between ~230 and ~350 °C. This retention reflects the redistribution of the FME between major mineral phases in the pelitic matrix. The Shimanto Belt exhibits open system behavior and significant loss is observed for all FME at temperatures between ~230 °C and ~ 350 °C, except for barium and rubidium for which the variations are much more limited. In the Shimanto Belt strontium is transfered to calcite in the veins, whereas there is a release of lithium, boron, cesium, and potentially minor proportion of barium into the fluid.

The fluid inclusions hosted in quartz in metasedimentary rocks that experienced temperatures 230–350 °C from the two localities show identical trends of evolution with temperature, such as increase in boron and cesium. In the Kodiak complex, the enrichment in boron and cesium in the fluid from ~ 230 °C to ~ 350 °C is consistent with a very small loss of boron and cesium from the solid, below that which can be detected by whole-rock analysis. In the Shimanto Belt, the significant FME loss is observed as temperature rises, which is consistent with a decrease in FME concentrations in the solid by a percolating fluid and water-rock ratios ranging between 0.5 and 2.2. These high water-rock ratios are at least one order of magnitude larger than the in situ pore fluid. The difference between the closed (Kodiak) and open (Shimanto) system behavior is attributed to the variation in the amount of internal strain (controlling premeability) and the distance to large-scale faults (draining fluids).

Furthermore, recovered chemical composition of these fluids have FME concentrations are at least an order of magnitude higher than that of interstitial pore fluid in shallow sediments or seawater. Similarities in composition between FIs trapped in metamorphic veins and mud volcano fluids or springs in subduction zone forearcs are consistent with the contribution of fluids being expelled from sedimentary units forming the seismogenic plate interface along permeable zones through the subduction zone forearc.

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CRediT authorship contribution statement

Kristijan Rajič: Writing – original draft, Software, Methodology, Investigation, Formal analysis, Data curation, Conceptualization. Hugues Raimbourg: Writing – review & editing, Visualization,



Fig. 14. The schematic scenario of FME budget evolution at studied sites, from the location within the wedge down to thin section: (a) A sketch of an accretionary wedge redrawn after Kastner et al. (2014) with the relative locations of studied units, the Foliated Morotsuka and the Kodiak Central Belt. The Foliated Morotsuka is located in the hanging wall of the out-of-sequence thrust (Nobeoka Tectonic Line), whereas the Kodiak Central Belt was underplated in the middle of thick turbiditic sequence far from any large-scale faults. A 3D sketch of (b) the Foliated Morotsuka and (c) the Kodiak Central Belt with presented deformation structures and possible scenarios for fluid flow. (d) and (e) show details of the mineralogy of metapelites from both localities and their influence on FME budget. In the Foliated Morotsuka (d), Li, B, Cs, and \pm Ba are leached from the host rock, with only Sr stored in vein-forming calcite. Conversely, in the Kodiak Central Belt (e), Li, B, Rb, and Ba are reorganized between illite, chlorite and albite and Sr being stored in the vein calcite. Only a small proportion of Cs is potentially lost to the fluid.

Supervision, Resources, Project administration, Methodology, Funding acquisition, Conceptualization. **Austin M. Gion:** Writing – review & editing, Visualization, Software, Methodology, Data curation, Conceptualization. **Catherine Lerouge:** Supervision. **Saskia Erdmann:** Writing – review & editing, Methodology, Investigation, Formal analysis, Data curation.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data availability

Supplementary data are available through Zenodo repository at https://doi.org/10.5281/zenodo.10658679 or online at https://doi.org/10.1016/j.chemgeo.2024.122141.

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