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Different stages of chemical alteration on metabasaltic rocks in the subduction channel: Evidence from the Western Tianshan metamorphic belt, NW China



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ABSTRACT

To understand the geochemistry of subduction zone metamorphism, especially the large-scale mass transfer at forearc to subarc depths, we carried out a detailed study of a ~ 1.5 m size metabasaltic block with well-preserved pillow structures from the Chinese Western Tianshan high- to ultrahigh-pressure metamorphic belt. This metabasaltic block is characterized by omphacite-rich interiors gradually surrounded by abundant channelized (veins) glaucophane-rich patches toward the rims. The glaucophane-rich rims share the same peak metamorphic conditions with omphacite-rich interiors, but have experienced stronger blueschist-facies overprinting during exhumation. Representative samples from the glaucophane-rich rims and omphacite-rich interiors yield a welldefined Rb-Sr isochron age of 307 \pm 23 Ma, likely representing this overprinting event. Both glaucophane-rich rims and omphacite-rich interiors show elevated K-Rb-Cs-Ba-Pb-Sr contents relative to their protolith, reflecting a large-scale enrichment of these elements and formation of abundant phengite during subduction. Compared with the omphacite-rich interiors, the glaucophane-rich rims have gained rare earth elements (REEs, > 25%), U-Th (~75%), Pb-Sr (>100%) and some transition metals like Co and Ni (25–50%), but lost P (~75%), Na (> 25%), Li and Be (~50%); K-Rb-Cs-Ba show only 10% loss. These chemical changes would be caused by serpentinite-derived fluids during the exhumation in the subduction channel. Therefore, there are two stages of fluid action in the subduction channel. As the formation of phengite stabilizes K-Rb-Cs-Ba at the first stage, the residual fluids released from the phengite-rich metabasaltic rocks would be depleted in these elements, which are unlikely to contribute to elevated contents of these elements in arc magmas if phengite remains stable at subarc depths. In addition, the decrease of U/Pb ratios as the preferred enrichment of Pb over U in the eclogitic rocks during the first stage chemical alteration may further lead to the lower radiogenic Pb isotope component of the deeply subducted ocean crust with time, which is inconsistent with the high radiogenic Pb isotope component of high μ (=²³⁸U/²⁰⁴Pb) basalts.

1. Introduction

The subduction zone metamorphism and related geochemical processes are significant for arc magmatism and mantle heterogeneity, which have attracted much attention in recent years (e.g., Kerrick and Connolly, 2001; Kelley et al., 2005; Keppler, 1996; McCulloch and Gamble, 1991; Niu et al., 2002; Niu and O'Hara, 2003; Niu, 2009; Tatsumi, 2005; John et al., 2012; Bebout, 2014; Marschall and Schumacher, 2012; Spandler and Pirard, 2013; Zheng and Chen, 2016). These studies have found that the geochemical behaviors of chemical elements during subduction zone metamorphism are much more complex than previously inferred through subduction zone magmatism, and they are not only controlled by the presence and stability of variable minerals (e.g., El Korh et al., 2009; Hermann and Rubatto, 2009; Spandler et al., 2003; Xiao et al., 2012, 2013, 2014, 2016) but also affected by the thermal structure of subduction zones (van Keken et al., 2011; Zheng et al., 2016) and the physicochemical properties of fluids, e.g., fluids with dissolved Na-Al silicates or halogen, supercritical fluids

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(Gao et al., 2007; Gao and Klemd, 2001; Haase et al., 2015; Hermann et al., 2006; John et al., 2008, 2012; Rubatto and Hermann, 2003; Schmidt et al., 2004; Spandler and Hermann, 2006; Zack and John, 2007; Zheng et al., 2011). The complex subduction-exhumation processes (i.e., multiple subduction-exhumation cycles) in subduction channels proposed in recent studies may lead to more complicated geochemical processes and consequences (Rubatto et al., 2011; Zheng and Hermann, 2014; Li et al., 2016; Xiao et al., 2016). Hence, the information on large-scale fluid-rock interactions in subduction channels is critically needed for understanding of large-scale mass transfer from the subducting slab to the overlying mantle wedge, which is essential towards a genuine understanding of arc magmatism and mantle heterogeneity.

It has been proposed that large-scale fluid fluxes and channelized fluids can facilitate element mobility in subduction zones (e.g., Ague, 2011; Bebout, 2007; Herms et al., 2012; John et al., 2008; Spandler and Hermann, 2006; Zack and John, 2007; Li et al., 2013; Bebout and Penniston-Dorland, 2016). In order to better understand the large-scale mass transfer from the subducting slab to the overlying mantle wedge, it is best to study subduction-zone metamorphic rocks affected by largescale fluid flows, the conduit of which is manifested by large-scale veins (e.g., John et al., 2012; Spandler and Hermann, 2006; van der Straaten et al., 2008, 2012). This study focuses on a ~ 1.5 m size metabasaltic block with well-preserved pillow structures from the Chinese Western Tianshan high-pressure (HP) to ultrahigh pressure (UHP) metamorphic belt (Figs. 1 and 2). Through a combined study of mineralogy, petrology and geochemistry, we discuss the behaviors of chemical elements in response to different stages of large-scale fluid-rock interactions, which offer insights into the possible large-scale mass transfer, contributing to our understanding of subduction zone magmatism and mantle heterogeneity.

2. Field Geology and petrography

The Western Tianshan HP-UHP metamorphic belt in NW China (Fig. 1) represents a paleo-convergent plate margin associated with successive northward subduction of the South Tianshan ocean crust beneath the Tarim Block during the Carboniferous (Gao and Klemd,

2003; Gao et al., 1999; Su et al., 2010; Klemd et al., 2011; Yang et al., 2013). This HP-UHP metamorphic belt is composed of pelitic schist, marble, serpentinite, blueschist and eclogite (Gao and Klemd, 2001; Wei et al., 2009; Li et al., 2012; Lü et al., 2013; Shen et al., 2015; Klemd et al., 2011; Yang et al., 2013) with protoliths of sandstone, pelite, carbonates, peridotite and basalts respectively (Ai et al., 2006; Gao and Klemd, 2003; Xiao et al., 2012). The Sm-Nd isochron age of ~343 Ma indicates the time of subduction metamorphism (Gao and Klemd, 2003), whereas the ⁴⁰Ar/³⁹Ar and Rb–Sr ages of ~310 Ma for white mica are thought to represent the time of retrograde overprint during exhumation (Klemd et al., 2005). Based on recent findings of coesite and their distributions (e.g., Lü et al., 2008, 2009, 2013, 2014; Lü and Zhang, 2012), the Chinese Western Tianshan metamorphic belt has been classified into HP and UHP metamorphic sub-units (Fig. 1; Lü and Zhang, 2012).

Numerous studies have shown large-scale fluid-rock interactions in the Chinese Western Tianshan HP-UHP metamorphic belt (Gao and Klemd, 2001; Gao et al., 2007; John et al., 2008, 2012; van der Straaten et al., 2008, 2012; Beinlich et al., 2010; Lü et al., 2012; Li et al., 2013, 2016). Our sampling location is along the Atantayi River (Fig. 1). The studied metabasaltic block with well-preserved pillow structures (Fig. 2) is a boulder, which has recently rolled from the steep mountain slope, and has omphacite-rich domains in the interiors, gradationally surrounded by abundant channelized glaucophane-rich patches toward the rims (Figs. 2a–c and 3a). Both omphacite-rich interiors and channelized glaucophane-rich rims are characteristically dominated by omphacite, glaucophane, phengite, and epidote as well as locally accumulated garnet plus some quartz, apatite, and carbonates, but glaucophane and epidote modes increase with decreasing omphacite mode toward rims (e.g., Fig. 2d–g).

Samples for this study are taken as a drilling core from this metabasaltic block with well-preserved pillow structures, ~ 10 cm long with a diameter of 2.5 cm (Fig. 2). The drilling core is divided into 4 sections for convenience, labelled as 22A, 22B, 22C and 22D from the rim surface to the interior (Fig. 2c). These four rock samples contain similar mineral assemblages with varying modal abundances. Samples 22C and 22D represent the omphacite-rich domain dominated by omphacite and phengite (Fig. 2d and e), while 22A and 22B represent the glaucophane-



Fig. 1. Geological sketch of the Chinese Western Tianshan HP-UHP metamorphic terrane (after Lü et al., 2013; Li et al., 2013) and our sampling location for this study.



Fig. 2. Field photographs and photos for relationship of our rock samples. (a–b) show the pillow-lava structure of our studied metabasaltic rock, i.e., omphacite-rich interiors with glaucophane-rich rims, which represent the previous fluid flow passage. The white dotted curve in (a) indicates the pillow margin, and the red dashed circle in (b) indicates the drilling position for this study. Gln – glaucophane; Omp – omphacite. (c) demonstrates the division of four sections from the top surface to the bottom of the drilling core for this study. (d–g) are relevant slides for the four sections in (c). These sections clearly show the decrease of glaucophane (bluish) and epidote (yellowish green) modal abundances, but the increase of omphacite (green) abundance. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

rich rim with abundant glaucophane and epidote (Fig. 2f and g). No garnet is present in these samples.

Samples 22C and 22D are mainly composed of omphacite and phengite matrix with some epidote and glaucophane porphyroblasts (Fig. 3c–f). Carbonate is locally present together with coarse-grained omphacite, phengite, and apatite (Fig. 3d and e). Epidote and glaucophane porphyroblasts are always euhedral and contain abundant inclusions of omphacite, phengite, carbonate, rutile and titanite (Fig. 4a and b). Glaucophane porphyroblasts also contain epidote inclusions. Rutile occurs as inclusions (Fig. 4a) and is locally replaced by titanite (Fig. 4b and c). Titanite is commonly present as poikiloblasts containing omphacite, phengite and quartz inclusions (Fig. 4c).

Samples 22A and 22B mainly comprise coarse-grained epidote and glaucophane (Fig. 3b) with phengite and much decreased omphacite (Fig. 4f). Carbonate, apatite and titanite are randomly present (Fig. 4f). More epidote and glaucophane with titanite occur as poikiloblasts and contain omphacite, phengite and carbonate (Fig. 4d–f), reflecting stronger blueschist-facies overprinting. Allanite is also present in epidote porphyroblast (Fig. 4e). Rutile has been entirely replaced by titanite.

3. Analytical methods

Each rock sample (22A, 22B, 22C and 22D) from the top surface to the bottom of the drilling core from the metabasaltic block (Fig. 2c) has been further cut into three pieces, i.e., the middle chip of the section was mounted in epoxy resin and polished for mineral analysis (Fig. 2d–g), while the other two chips were ultrasonically cleaned before hand-crushed in an agate mortar into powders for bulk-rock analysis. Considering the highly heterogeneous mineral distribution in metamorphic rocks, especially the along-core modal variation, this division method can make the bulk-rock composition of sample powders to be representative and consistent with the mineral assemblages as observed in thin sections. Thin sections for the other two drilling cores from the same metabasaltic block have also been prepared for petrography study (e.g., Fig. 3).

3.1. Mineral compositions

Major elements of hydrous minerals were analyzed using a JXA-8100 electron probe micro-analyzer (EPMA) at Chang'an University, China. The analyses were performed using 15 kV accelerating voltage



Fig. 3. Photomicrographs. (a) Relationship between omphacite-rich interiors and glaucophane-rich rims. The mineral from omphacite-rich interiors can intervene glaucophane-rich rims. (b) Glaucophane-rich rim, dominated by glaucophane and epidote with phengite. (c–f) are for omphacite-rich interior, dominated by a matrix of fine-grained omphacite and phengite with some glaucophane and epidote. (e) Locally distributed coarse-grained apatite and omphacite. (f) Locally distributed carbonate, which can be present as the inclusion of glaucophane porphyroblasts. Mineral abbreviations used in this paper are: Aln – allanite; Ap – apatite; Ca – carbonate; Ep – epidote; Gln – glaucophane; Omp – omphacite; Ph – phengite; Qz – quartz; Rt – rutile; Ttn – titanite (mostly referred to Whitney and Evans, 2010).

and 10 nA probe current. Standards used for calibration are: albite for Na, quartz for Si, orthoclase for K, apatite for P and Ca, magnetite for Fe, pyrophanite for Mn and Ti, chromite for Cr and Fe, forsterite for Mg, and jadeite for Al (Xiao et al., 2013). Representative analytical results of mineral major element contents using EPMA are given in Table S.1. Mineral trace elements and major elements for anhydrous minerals were measured on polished sections by using an LA-ICP-MS (Agilent 7500a with GeoLas 2005 193 Eximer Laser sampler) at the State Key laboratory of Continental Dynamics, Northwest University, China. The repetition rate of laser ablation is 6 Hz, and the pit size is 32 µm. During each analysis, the acquisition times for the background (gas blank) and

the sample ablation are 20–30 s and 50 s respectively. United States Geological Survey (USGS) glasses BCR-2G, BHVO-2G, BIR-1G and one synthetic glass GSE-1G (Guillong et al., 2005; Jochum et al., 2005) are used as reference materials. For anhydrous minerals (i.e., omphacite, and titanite), instead of using the internal standard for calibration, the concentrations of all the elements of interest (both major and trace elements) are analyzed simultaneously and calibrated through an internal standard-independent calibration method (see Liu et al., 2008). For hydrous minerals, major elements analyzed by using EPMA are chosen as internal standards for calibration, i.e., Si for amphibole, epidote group minerals and phengite; Ca for apatite and carbonate. The



Fig. 4. SEM-EDS photographs. See caption of **Fig. 3** for mineral abbreviations. (a–c) are for omphacite-rich cores, while (d–f) are for glaucophane-rich rims. (a) and (d) display variable mineral inclusions in epidote porphyroblasts from the omphacite-rich interior and glaucophane-rich rim respectively, including omphacite, phengite, rutile, titanite and quartz. (b) A big glaucophane porphyroblast with its variable mineral inclusions including titanite and rutile, and an epidote poikiloblast with omphacite, rutile and phengite inclusions. (e) shows allanite composition core of the epidote porphyroblast. (c, f) Abundant variable mineral inclusions in titanite with rutile residuals, i.e., omphacite, phengite, and quartz. Epidote, glaucophane and titanite poikiloblasts from both omphacite-rich interiors and glaucophane-rich rims generally share similar mineral inclusions, i.e., omphacite, phengite, quartz, and rutile, which indicate that these samples have experienced similar eclogite facies metamorphism. Incl. – inclusion; porp. – porphroblast.

analytical uncertainty is within 10% and the precision determined by repeated analysis of GSE-1G is generally better than 5%. During analysis, inclusions are purposely avoided. Analytical results of mineral trace element contents are given in Table S.2.

3.2. Bulk-rock major and trace elements

Bulk-rock major and trace element contents were analyzed at Northwest University, China. The major elements were analyzed by using X-ray fluorescence (Rigaku RIX 2100 XRF) on fused glass disks with analytical precision better than 5% as determined by duplicate analyses. An additional sample powder $\sim 1 \text{ g}$ is heated in a muffle furnace at 1000 °C for loss on ignition (LOI) determination.

For the trace element analysis, 50 mg sample powders were digested in an HF + HNO₃ mix in high-pressure Teflon bombs at 190 °C for 48 h. After being dried, 3 ml 50% HNO₃ was added and sample powders was dissolved again in high-pressure Teflon bombs at 150 °C for 12 h. Rh as an internal standard was added to sample solutions, which are finally diluted to 80 g. ICP-MS (Agilent 7500a) was used for trace element analysis with analytical accuracy better than 5% for most trace elements. The analytical results of bulk-rock compositions are given in Table 1.

3.3. Sr-Nd isotope

Sample digestion for Sr-Nd isotope analysis was done at the National Cheng-Kung University, Taiwan. Sample powder of 100 mg was decomposed in a high-pressure bomb by 3 ml concentrated HF–HNO₃ mixture at 190 °C for 24 h. After being dried, 5 ml 6N HCl was added, followed by conversion to nitrate form (see Liu et al., 2015). The Sr and Nd separation was done at the Institute of Earth Sciences, Academia Sinica in Taipei through a two-column technique using Bio-Rad AG50W-X8 and Ln-B25-A (Eichron) resins respectively (Jahn et al., 2009).

Finnigan MAT-262 thermal ionization mass spectrometer (TIMS) and Finnigan Triton TIMS were used for Sr and Nd isotope analyses respectively. The Sr and Nd isotopic ratios were normalized against the value of $^{86}\text{Sr}/^{88}\text{Sr} = 0.1194$ and $^{146}\text{Nd}/^{144}\text{Nd} = 0.7219$ respectively. The 2σ values for all the analyses are less than 0.000007 for $^{87}\text{Sr}/^{86}\text{Sr}$ and less than 0.000008 for $^{143}\text{Nd}/^{144}\text{Nd}$ (Table S.3). The measured isotopic ratio for NBS987-Sr standard is $^{87}\text{Sr}/^{86}\text{Sr} = 0.710245 \pm 0.000010$ (2 σ) and that for JMC Nd standard is $^{143}\text{Nd}/^{144}\text{Nd} = 0.511817 \pm 0.000007$ (2 σ), which are consistent with the data of Shellnutt et al. (2012; 0.710248 \pm 0.00001, 2 σ) and Jahn et al. (2009; 0.511821 \pm 0.000016, 2 σ) respectively.

4. Analytical results

4.1. Mineral compositions

In chondrite normalized trace element diagram (Fig. 5), epidote from 22A (representing the glaucophane-rich rim) show greater variations of Th, U and rare earth elements (REEs) than 22D (representing the omphacite-rich domain; $10-10^4$ times vs. 10^2-10^3 times the chondritic values, Fig. 5a). Allanite as the core of an epidote crystal from 22A contains the highest REEs (e.g., 3192 ppm in analyzed point 22A-C4-1, Table S.2; Figs. 4e and 5a), while some other epidote crystals from 22A show only several ppm REEs (Table S.2). Apatite also shows consistently high REEs-Th-U-Sr (up to 10-100 times the chondritic values, Fig. 5c). Carbonate has high Sr contents (>10 times the chondritic value, Fig. 5e). All the analyzed points on phengite from both 22D and 22A generally share a similarly high contents of large ion lithophile element (LILE), including Ba-Rb-Cs (10–10³ times the chondritic values, Fig. 5b). Glaucophane shows low contents for most trace elements of interest (Fig. 5f) except Li and Be (up to > 10 ppm Li and several ppm Be; Table S.2). Glaucophane from 22D shows generally higher Be than that from 22A (several ppm vs. lower than detection limits; Table S.2).

Titanite is characterized by high Nb and Ta contents (up to 3 orders of magnitude higher than the chondritic values) and heavy rare earth elements (HREEs; ~10 times the chondritic values, Fig. 5d). Omphacite also shows low contents for most trace elements of interest (Fig. 5f), but contains the highest Li and Be among all the analyzed minerals (tens of ppm Li and several ppm Be; Table S.2).

Table 1

Bulk-rock compositions of omphacite-rich interior and glaucophane-rich rim of metabasaltic rock from the Chinese Western Tianshan HP-UHP metamorphic belt.

Sample name	22A 40% Gln + 15% Ph + 5% Omp + 40% Ep + some (Ttn + Ca)	22B 20% Gln + 15% Ph + 20% Omp + 40% Ep + some (Ttn + Ca + Ap)	22C 5% Gln + 25% Ph + 40% Omp + 25% Ep + 5% (Ttn + Ca + Ap)	22D 5% Gln + 25% Ph + 50% Omp + 15% Ep + 5% (Ttn + Ca + Ap)	Slope to data point 22A/22D	Gain/Loss relative to 22D ΔC _i /C _i °	Scaled	
							C _i ° (22D)	C _i ^A (22A)
wt.%								
SiO_2	46.27	46.01	46.30	46.46	0.996	-0.011	48.00	47.80
TiO ₂	1.51	1.56	1.56	1.52	0.993	-0.014	1.00	0.99
Al_2O_3	19.42	18.98	17.75	17.26	1.125	0.117	47.00	52.88
TFe_2O_3	7.68	7.76	6.59	6.71	1.145	0.136	46.00	52.65
MnO	0.04	0.04	0.04	0.04	1.000	-0.007	45.00	45.00
MgO	4.67	4.60	4.96	5.29	0.883	-0.124	44.00	38.84
CaO	9.78	10.49	10.49	10.35	0.945	-0.062	43.00	40.63
Na ₂ O	1.93	2.05	2.42	2.78	0.694	-0.311	42.00	29.10
R ₂ O	0.13	0.12	4.10	4.01	0.925	-0.082	38.00 41.00	10.06
	4 42	4 51	5.05	4.61	0.245	-0.048	40.00	38 35
TOTAL	99.56	99.58	99.54	99.56	0.909	0.010	10.00	00.00
ppm	8.00	9 67	12.4	15.0	0 510	0.404	12.00	6 1 2
LI Be	8.09 0.87	8.07	13.4	15.9	0.510	-0.494	12.00	0.12 6.28
Sc	30.1	31.3	30.4	29.3	1 028	0.433	10.00	10.28
v	233	237	239	237	0.979	-0.028	9.00	8.81
Cr	177	177	175	159	1.114	0.106	8.00	8.91
Co	46.0	43.6	34.6	36.6	1.255	0.245	7.00	8.78
Ni	120	117	76.5	84.2	1.424	0.414	6.00	8.54
Cu	63.6	57.8	62.4	137	0.463	-0.541	5.00	2.31
Zn	74.9	71.2	60.1	65.4	1.144	0.136	4.00	4.58
Ga	24.2	24.1	20.2	18.9	1.281	0.272	3.00	3.84
Ge	1.27	1.34	1.18	1.13	1.122	0.113	2.00	2.24
Rb	57.4	53.8	64.0	62.6	0.916	-0.090	36.00	32.99
Sr	549	563	335	262	2.095	1.080	32.00	67.05
Y 7r	19.4	20.9	15.2	13.8	1.401	0.390	19.00	20.01
Nb	13.8	14.0	104	155	0.908	-0.099	39.00	35.41
Cs	1.62	1.52	1.82	1.78	0.909	-0.098	35.00	31.81
Ba	434	400	470	461	0.941	-0.066	37.00	34.81
La	14.3	14.6	12.4	10.3	1.397	0.387	28.00	39.11
Ce	34.1	35.2	29.1	24.3	1.404	0.394	27.00	37.92
Pr	4.49	4.68	4.07	3.40	1.320	0.310	26.00	34.31
Nd	19.9	20.7	18.0	15.2	1.306	0.296	25.00	32.64
Sm	4.80	5.07	4.28	3.65	1.314	0.304	24.00	31.52
Eu	1.72	1.84	1.45	1.23	1.397	0.387	23.00	32.14
Gđ	4.83	5.24	4.26	3.64	1.327	0.318	22.00	29.20
I D Du	0.76	0.81	0.65	0.56	1.350	0.346	21.00	28.48
Ho	4.13	4.40	0.60	0.55	1.336	0.328	20.00	20.70
Er	1.86	1.97	1.46	1.38	1.347	0.337	17.00	22.90
Tm	0.22	0.23	0.17	0.17	1.331	0.321	16.00	21.29
Yb	1.23	1.26	0.95	0.89	1.392	0.382	15.00	20.88
Lu	0.16	0.16	0.13	0.12	1.365	0.355	14.00	19.11
Hf	3.52	3.74	3.64	3.47	1.015	0.007	13.00	13.19
Та	0.87	0.90	0.91	0.88	0.995	-0.012	34.00	33.82
Pb	2.72	2.74	1.60	1.24	2.185	1.169	33.00	72.10
Th	1.45	1.39	1.07	0.89	1.633	0.621	30.00	48.98
U	0.46	0.50	0.31	0.26	1.758	0.746	29.00	51.00
La/Ta	16.5	16.3	13.7	11.7				
Th/Nb	0.11	0.10	0.07	0.06				
Ta/U	1.90	1.81	2.90	3.35				
Nb/U	30.0	28.2	46.2	58.0				
Ce/Pb	12.0	12.8	18.2	19.5				
IND/18	10.0 2.17	10.0	2 /1	17.3 2.41				
Rh/Sr	0.10	2.01 0.10	0.19	0.24				
Sm/Nd	0.24	0.24	0.24	0.24				
U/Pb	0.17	0.18	0.20	0.21				

The highlighted (i.e., SiO₂-MnO-TiO₂-Ta-Zr-Hf) are reference elements for isocon ploting, and the acquired slope of isocon is 1.0073.

4.2. Bulk-rock major and trace element contents

From 22D to 22A with increasing extents of blueschist-facies overprinting, major element contents vary, reflecting controls of mineral modal abundances. Al_2O_3 content obviously increases towards glaucophane-rich rims (Table 1), consistent with the increasing modal abundance of epidote resulting from addition of Al associated with blueschist-facies overprinting. Whereas, Na₂O and P₂O₅ contents slightly decrease towards glaucophane-rich rims (Table 1; Fig. 6b), consistent with less omphacite and apatite respectively.



Fig. 5. Chondrite normalized mineral composition distributed diagrams (Sun and McDonough, 1989). (a) Epidote from both the least and the most blueschist-facies overprinted rocks with variably enriched Th-U-REEs-Pb-Sr; one allanite is highlighted. (b) Phengite from both the least and the most blueschist-facies overprinted rocks contains consistently high Ba-Rb-Cs contents. (c) Apatite with consistently high REEs-Th-U-Sr. (d) Titnaite showing high Nb-Ta, some U and REEs. (e) Carbonate with high Sr contents. (f) Glaucophane and omphacite contain insignificant trace element contents.

By using commonly accepted immobile high field strength elements (HFSEs) and HREEs (Pearce, 2008), the four rock samples in this study plot within the mid-ocean ridge basalts (MORB)-oceanic island basalts (OIB) array, close to the data point for enriched-MORB (E-MORB) and OIB (Fig. 6a). Furthermore, these rocks consistently show similar trace element patterns to that of E-MORB (Fig. 6b) except for the notably higher K-Rb-Ba-Pb-Sr contents and stronger L(light-)REE/HREE fractionation. With increasing extents of blueschist-facies overprinting from 22D to 22A, contents of Th-U-REEs-Pb-Sr and some siderophile elements (e.g., Co, Ni) variably increase (Fig. 6b; Table 1), while Li and Be contents tend to decrease (Table 1). K-Rb-Cs-Ba contents show only slight differences, and no obvious change of HFSEs or Sc-V among the four rocks (Fig. 6b; Table 1).

4.3. Sr-Nd isotopes

The four bulk samples yield a well-defined Rb-Sr isochron age of 307 ± 23 Ma (MSWD = 6.9) and an initial $^{87}\text{Sr}/^{86}\text{Sr}$ ratio of 0.70461 ± 0.00015 (Fig. 7a; using ISOPLOT 3 of Ludwig, 2003). This Rb-Sr isochron age, within error, overlaps with the age of the rehydration overprint at ~310 Ma for the Chinese Western Tianshan HP-UHP metamorphic belt in literature (Gao and Klemd, 2003; Klemd et al., 2005, 2011), and thus represents the time of blueschist-facies overprinting during exhumation. The limited range of $^{147}\text{Sm}/^{144}\text{Nd}$ ratios (0.143–0.148) does not give a useful Sm-Nd isochron age. The initial Sr-Nd isotope values of the four samples calculated at 310 Ma give ($^{87}\text{Sr}/^{86}\text{Sr})_{310\text{Ma}}$ of 0.704560 to 0.704614 and $\epsilon_{\text{Nd}}(t)$ values of +1.9 to +2.1 at t = 310 Ma (Table S.3). In Fig. 7b, all the Sr-Nd isotope data points for the four samples plot as a cluster in the MORB-OIB mantle array.

5. Discussion

5.1. Geochemical behaviors of chemical elements

The exhumed HP-UHP metamorphic rocks of oceanic subduction zones are always an integrated product, which is likely to have experienced a series of geochemical processes including seafloor alteration during protolith generation, chemical alteration during subduction and retrograde overprint during exhumation in the subduction channel (e.g., Xiao et al., 2016). As discussed above, mineral assemblages and mineral inclusions in glaucophane and epidote porphyroblasts from the glaucophane-rich rims and omphacite-rich interiors are similar (Figs. 2-4). This reflects that both glaucophane-rich rims and omphacite-rich interiors have experienced a similar eclogite-facies peak metamorphic condition. However, more epidote and glaucophane porphyroblasts replacing omphacite towards glaucophane-rich rims (Figs. 2 and 3) reflect that the glaucophane-rich rim experienced a greater extent of overprinting than the omphacite-rich interior. These petrological results indicate that our studied four rock samples have similarly experienced two main stages of geochemical alteration in the subduction channel with different extents of blueschist-facies overprinting. Therefore, it is important to identify various effects of these geochemical processes on the behaviors of chemical elements at different stages.

5.1.1. Seafloor alteration

Ba and Rb have been found to behave similarly during subduction zone metamorphism as they are preferentially sequestered into phengite (e.g., Hermann and Rubatto, 2009; Xiao et al., 2012, 2014, 2016). However, it has been reported that seafloor alteration can lead to obvious enrichment of Rb (9 times of the original) but not Ba (Kelley et al.,



Fig. 6. (a) Covariation diagram of Th/Yb and Nb/Yb for protolith fingerprint of our subduction-zone metamorphosed rock samples (after Pearce, 2008). (b) E-MORB normalized bulk-rock composition distributed diagram (Niu and O'Hara, 2003). N-MORB and OIB are also plotted for comparison (Niu and O'Hara, 2003). The rocks representing glaucophane-rich rims plot as blue circles, i.e., the bluest circles represent rock 22A, while the light blue circles represent rock 22B. The rocks representing omphacite-rich cores plot as yellowish circles, i.e., the light yellow circles represent rock 22C, while the orange circles represent rock 22D. The legends used in Figs. 7–9 are also following these. The elements plotted in the grey areas (i.e., K-Ba-Rb-Pb-Sr) are those mobilized by pervasive enrichment process covering the four rocks at the first stage geochemical alteration in the subduction channel before the blueschist-facies overprinting. (For interpretation of this article.)

2003). Hence, the rocks that have experienced seafloor alterations are expected to have low Ba/Rb ratios. In Fig. 8, the four rock samples show relatively lower Ba/Rb ratios than those of seafloor basalts while no fractionation of Ba/Rb ratios among these rock samples, probably indicating the effects of seafloor alteration on the trace element systematics of these rock samples.

5.1.2. The first stage chemical alteration during subduction

Given the E-MORB-like protolith composition as shown in Fig. 6, our rock samples show commonly elevated contents of K-Rb-Cs-Ba-Pb-Sr compared to those in E-MORB (Fig. 6b), and thus these elements

cannot be inherited from the protolith. As all these four rock samples have experienced eclogite-facies metamorphism with different extents of blueschist-facies overprinting as manifested by mineral assemblages (Figs. 2 and 3), the consistency in the elevated contents of K-Rb-Cs-Ba-Pb-Sr in these rocks relative to their E-MORB-like protolith suggests that the large-scale enrichment of these elements occurred during the eclogite-facies metamorphism when the first stage chemical alteration took place in the subduction channel before the blueschist-facies overprinting.

5.1.3. The second stage chemical alteration during exhumation

The blueschist-facies overprinting of metabasaltic pillow interiors and rims to variable extents represents the second stage chemical alteration in the subduction channel. We take omphacite-rich 22D with the least blueschist-facies overprint as representing the rock composition before this overprint. Considering the increasing extents of blueschist-facies overprinting from 22D to 22A (Figs. 2d–g and 3), the systematic elemental changes from 22D to 22A (Fig. 6b; Table 1) indicate that P-Na-Mg-K-Ca-Al-Fe, Th-U-REEs-Pb-Sr-Li-Be, some transition metals (e.g., Co and Ni) have been mobilized to varying extents, while Rb-Cs-Ba, HFSEs and Sc-V have been largely conserved with blueschistfacies overprinting at the second stage chemical alteration in the subduction channel.

To further understand the mass change with blueschist-facies overprinting during exhumation, we chose the highly overprinted glaucophane-rich 22A to compare with 22D in an isocon diagram (Fig. 9a). The slope of isocons defined by immobile SiO₂-MnO-TiO₂-Ta-Zr-Hf is 1.0073, very close to 1, and thus indicates a very small mass loss or gain (Grant, 2005). As the data points plotted above and below the isocon reflect the element gain and loss respectively, the most evident changes with blueschist-facies overprint are: (1) > 100% Pb and Sr gain; (2) \sim 75% U-Th gain; (3) > 25% REEs gain; (4) 25–50% transition metal (e.g., Ni and Co) gain; (5) \sim 25% and 75% Na and P loss respectively; (6) \sim 50% Li and Be loss. On the other hand, Sc-V-Nb and LILE (K-Rb-Cs-Ba) show insignificant changes from 22D to 22A, i.e., no more than 10% gain or loss. Given the immobility of Ta, Zr and Hf, the increase of La/Ta and Th/Nb ratios but the decrease of Ta/U and Nb/U ratios from 22D to 22A (Fig. 9b) further reflects the addition of REEs (represented by La)-Th-U with blueschist-facies overprinting.

5.2. Controls on the geochemical behaviors of chemical elements

Based on our previous studies (Xiao et al., 2014, 2016), the geochemical behaviors of trace elements during subduction zone metamorphism have been found to be controlled by a number of factors,



Fig. 7. (a) Rb-Sr isochron diagram of the four rock samples from the drilling core, yielding a well-defined isochron age of 307 ± 23 Ma (MSWD = 6.9). (b) Initial Sr-Nd isotopic compositions of the four rock samples from the drilling core are calculated back to 310 Ma, and have been plotted in the mantle array. The legends used are the same to Fig. 6. The initial Sr-Nd isotopes for metasomatic eclogites from the Chinese Western Tianshan HP-UHP metamorphic belt in the study of van der Straaten et al. (2012) are also plotted for comparison.



Fig. 8. Covariation diagrams for element contents and element ratio of bulk rocks (after van der Straaten et al., 2012). (a) Ba/Rb – K. The relatively lower Ba/Ra of our samples than seafloor basalts may reflect the effects of seafloor alteration before subduction. (b) The correlated increase of Co and Ni contents with the increasing degree of blueschist-facies overprint, indicating the large fluid fluxes are mainly externally derived from serpentinite. The legends used are the same to Fig. 6.

particularly the formation and stability of specific minerals in metamorphic rocks. Heterogeneous compositions of metamorphic minerals and bulk rocks are determined by element availability, i.e., the inherited composition and the competition between coexisting minerals (Xiao et al., 2016), which is also controlled by the presence and stability of metamorphic minerals. For an open system, variable physicochemical compositions and abundances of fluids can affect the behaviors of chemical elements (e.g., John et al., 2012; Spandler and Hermann, 2006; van der Straaten et al., 2008, 2012; Zack and John, 2007), and thus can further affect the compositions of metamorphic minerals and bulk rocks. The studied metabasaltic block has large-scale channelized (veins) glaucophane-rich rims surrounding the omphacite-rich interiors. As veins represent previous fluid flow passages, the large-scale channelized glaucophane-rich patches reflect the presence and action of



Fig. 9. (a) Isocon diagram following the method of Grant (2005), taking 22D and 22A as the least and the most blueschist-facies overprinted rocks. Elements with the individual slope within 1.000 \pm 0.020, i.e., SiO₂, MnO, TiO₂, Ta, Zr and Hf (highlighted in Table 1), are chosen as the reference elements to plot a best-fit linear array through the origin as an isocon. The relevant equation and criterion for isochon plotting are referred to Gresens (1967), Grant (2005) and van der Straaten et al. (2012). (b) Bulk-rock trace element ratio changes with the blueschist-facies overprint. Th/Nb ratio has been ten times for better clarification. The legends used are the same to Fig. 6.



large-scale fluid flux in the subduction channel (Bebout, 2007).

As K-Rb-Cs-Ba-Pb-Sr are expected to be enriched in meta-sedimentary rocks (e.g., Breeding et al., 2004; Hermann and Spandler, 2008), the common enrichments of these elements during the first stage chemical alteration in the subduction channel may have resulted from addition of these elements by fluids released from subducting sediments. Furthermore, considering the significance of phengite in hosting K-Rb-Cs-Ba (Fig. 5b; e.g., Sorensen et al., 1997; Spandler et al., 2003; El Korh et al., 2009; Zheng et al., 2011; Xiao et al., 2014, 2016), this enrichment facilitates abundant phengite formation as observed (Fig. 3), which results in the great conservation of these elements during this process as manifested by even much higher contents of these elements in the four studied rock samples than those in OIB (Fig. 6b).

With the blueschist-facies overprinting during the second stage chemical alteration in the subduction channel, the increase of REEs-Sr-Pb-Th-U with increasing extents of blueschist-facies overprinting (Fig. 9) manifests precipitation of these elements in the form of epidote from external fluids (Figs. 5a and 10; e.g., Xiao et al., 2014, 2016). The obvious decrease of Li and Be contents during this process is consistent with the loss of Li and Be associated with the replacement of omphacite by glaucophane. Because transition metals are enriched in mantle rocks relative to mantle-derived basaltic rocks and show immobility during serpentinization (e.g., ~100 ppm vs. ~30-~50 ppm for Co, ~800-~2000 ppm vs. 10 s-100 s ppm for Ni, Niu, 2004; Niu and Batiza, 1997), the correlated increase of Co and Ni contents from the omphacite-rich interior to the glaucophane-rich rim (Fig. 8b) suggests that the fluids responsible for the blueschist-facies overprinting may be derived from serpentinite (van der Straaten et al., 2008, 2012), which is the most significant fluid budget in the subducting slab and crucial for element mobility in the subduction channel (Barnes et al., 2014; Deschamps et al., 2012; Herms et al., 2012; Rüpke et al., 2002, 2004; Scambelluri et al., 2014; Shen et al., 2015; Spandler et al., 2014; Stern et al., 2006).

Whereas, K-Rb-Cs-Ba contents show rather insignificant changes (only $\sim 10\%$ loss) with blueschist-facies overprinting, although these elements were thought to be easily mobilized with fluids, especially in an open system (e.g., Breeding et al., 2004; van der Straaten et al., 2012; Xiao et al., 2014). This is different from the result of recent studies by van der Straaten et al. (2008, 2012) for metabasaltic rocks with pillow structures affected by similar blueschist-facies overprinting from the Chinese Western Tianshan HP-UHP metamorphic belt, i.e., even up to ~200% of K-Rb-Cs-Ba have been gained during blueschistfacies overprinting. The insignificant mobility of LILEs during blueschist-facies overprinting in our studied rocks may be attributed to the significant conservation of LILEs in abundant phengite, the formation of which resulted from the commonly great enrichments of LILEs during the first stage chemical alteration in the subduction channel. Meanwhile, it also reflects insignificant addition of LILEs by serpentinitederived fluids during the blueschist-facies overprinting, which thus cannot lead to the formation of more phengite. Hence, after the reaction

Fig. 10. Co-variation diagrams for trace element contents of epidote from both the least and the most blueschist-facies overprinted rocks (22D represented by the orange solid circles vs. 22A represented by the blue solid circles). Allanite core of an epidote porphyroblast has also been presented. The relationship of La with Pb (a) and Sr (b) shows higher contents of Pb and Sr in epidote from 22A than of those from 22D, resulting from the greatest addition of these elements by externally derived fluids. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

between phengite-rich metabasaltic rocks and large fluid fluxes derived from serpentinite, K-Rb-Cs-Ba can be still largely conserved and show limited mobility. However, in studies of van der Straaten et al. (2008, 2012), phengite modal abundance in the eclogitic interior is much lower (e.g., no more than ~10%, van der Straaten et al., 2008) while more phengite formed during blueschist-facies overprinting (e.g. ~20%, van der Straaten et al., 2008) due to addition of the fluids from serpentinite but with crucial addition of sediment-derived component.

5.3. Geodynamic implications

The slab-mantle interface composed of diverse lithologies and characterized by high permeability can facilitate the formation of channelized fluids as well as intense fluid-rock interactions, which may enhance element mobility (e.g. Bebout, 2007; Beinlich et al., 2010; Breeding et al., 2004; John et al., 2008; Li et al., 2013; van der Straaten et al., 2008; Zheng, 2012). The common enrichment of K-Rb-Cs-Ba in our studied rocks during the first stage chemical alteration in the subduction channel reflects the mobility and introduction of these elements, most likely derived from dehydration of the subducting sediments. This enrichment facilitates abundant phengite formation as widely observed in our samples, while the residual fluids released from these phengite-rich metabasalts may be depleted in K-Rb-Cs-Ba. Furthermore, the conservation by the already present phengite and the low contents of K-Rb-Cs-Ba in the serpentinite-derived fluids during the second stage chemical alteration in the subduction channel result in no obvious change of these elements with the infiltration of large fluid fluxes (Fig. 2a). This indicates that the reaction between phengite-rich metabasaltic rocks and large fluid fluxes mainly derived from dehydration of the serpentinites cannot make phengite breakdown/dissolved or lead K-Rb-Cs-Ba to be mobilized at forearc depths. Therefore, the elevated contents of K-Rb-Cs-Ba widespread in arc magmas (known as the arc signature, e.g., McCulloch and Gamble, 1991) may be caused by the phengite breakdown at subarc depths, where fluid mobile elements can be transferred from the subducting crust to the mantle (Zheng et al., 2016). Otherwise, it requires the involvement of supercritical fluids or hydrous melts originated from the subducting slab (e.g., Xiao et al., 2012, 2014).

On the other hand, with varying extents of element changes, element ratios (e.g., Nb/U, Ta/U, Ce/Pb, U/Pb) will change accordingly, including radioactive parent to radiogenic daughter element ratios (Fig. 9b). The common enrichment of Pb but insignificant alteration of U during the first stage chemical alteration in the subduction channel (Fig. 6b) indicates the decrease of U/Pb ratios in the highly eclogitized rocks (e.g., sample 22D), which will further result in the lower radiogenic Pb isotope component of the deep mantle with time after the continuous subduction of these materials. Hence, the high μ (HIMU; $\mu = ^{238}$ U/²⁰⁴Pb) OIB may not be caused by the long time-integrated subducting MORB (e.g., Hofmann, 1997) but by the other mechanism. For example, Castillo (2015) has proposed that subducting ancient

marine carbonates can lead to high μ component as high U but low Pb precipitated from seawater (Xiao et al., 2017).

6. Conclusions

In this study, the glaucophane-rich rims share the similar peak metamorphic conditions with omphacite-rich cores, but show stronger blueschist-facies overprinting during exhumation, the age of which has been constrained to be 307 \pm 23 Ma (MSWD = 6.9) by a well-defined Rb-Sr isochron age. Our rock samples record at least two stages of chemical alteration in the subduction channel. During the first stage chemical alteration. K-Rb-Cs-Ba-Pb-Sr are pervasively enriched in the studied rocks, producing abundant phengite during subduction. The second stage chemical alteration with blueschist-facies overprinting resulted in the enrichment of REEs (> 25%), U-Th (> 50%), Pb-Sr (> 100%) added by fluids and depletion of P (\sim 75%) – Li and Be $(\sim 50\%)$ as the result of the formation of epidote and glaucophane at the expense of apatite and omphacite. The rather insignificant loss of K-Rb-Cs-Ba (< 10%) with blueschist-facies overprinting results from the conservation of these elements by already formed abundant phengite, indicating that the reaction of large fluid fluxes with phengite-rich metabasaltic rocks has limited effects on the mobility of K-Rb-Cs-Ba during exhumation. The phengite breakdown at subarc depths is the key to the elevated contents of these elements in arc magmas; otherwise, the other mechanisms like supercritical fluids or hydrous melts from subducting sediments may be responsible for the elevated K-Rb-Cs-Ba contents in arc magmas. As the decrease of U/Pb ratios in the highly eclogitized rocks results from the common enrichment of Pb but insignificant alteration of U during the first stage chemical alteration in the subduction channel, it will further lead to the lower radiogenic Pb isotope component of the deeply subducted ocean crust with time, inconsistent with the feature of HIMU. Hence, the HIMU of OIB may be caused by the other mechanism, e.g., the subducting marine carbonates rather than the long time-integrated subducting MORB in the traditional model.

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Appendix A. Supplementary material

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.jseaes.2017.06.001.

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