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The Growth of Sodic Amphibole at the Greenschist- to Blueschist-facies Transition (Dent Blanche, Western Alps): Bulk-rock Chemical Control and Thermodynamic Modelling

Paola Manzotti D^{1,2}*, Michel Ballèvre³, Pavel Pitra^{3,4}, Benita Putlitz², Martin Robyr² and Othmar Müntener²

¹Department of Geological Sciences, Stockholm University, Stockholm 106 91, Sweden; ²Institute of Earth Sciences, University of Lausanne, Géopolis, Quartier Mouline, 1015 Lausanne, Switzerland; ³Université Rennes 1, CNRS, Géosciences Rennes-UMR 6118, F-35000 Rennes, France; ⁴Česká geologická služba, Prague 1, Czech Republic

*Corresponding author. Telephone: +46 (0)8 16 4745. E-mail: paola.manzotti@geo.su.se

ABSTRACT

The sodic amphibole glaucophane is generally considered as indicative of blueschist-facies metamorphism. However, sodic amphiboles display a large range in chemical compositions, owing principally to the $Fe^{2+}Mg_{-1}$ and $Fe^{3+}Al_{-1}$ substitutions. Therefore, the whole-rock composition (namely its Na₂O and FeO* content, and the Fe^{2+} - Fe^{3+} ratio), strongly controls the stability field of the sodic amphiboles at the transition from greenschist- to blueschist-facies conditions. Neglecting these variables can lead to erroneous estimates of the metamorphic conditions and consequently the tectonic framework of the rocks. This paper explores the mechanisms that control the development of sodic amphibole and sodic pyroxene within the basement of the Dent Blanche Tectonic System (Western Alps), as a result of the Alpine metamorphic history. Field, petrographic and geochemical data indicate that sodic amphibole and sodic pyroxene form in different rock types: (1) in undeformed pods of ultramafic cumulates (hornblendite), sodic amphibole (magnesioriebeckite) forms coronas around magmatic pargasite; (2) metatonalite displays patches of radiating sodic (magnesioriebeckite) and calcic (actinolite) amphiboles; (3) sodic amphibole (magnesioriebeckite-glaucophane) occurs with high-Si potassic white mica (phengitic muscovite) in fine-grained (blue) schists; (4) in mylonitized granitoids (amphibole-gneiss) metasomatized along the contact with ultramafic cumulates, sodic amphibole (magnesioriebeckite-winchite) mainly forms rosettes or sheaves, generally without a shape-preferred orientation. Only locally are the needles aligned parallel to the stretching lineation. Pale green aegirine-augite is dispersed in an albite-quartz matrix or forms layers of fine-grained fibrous aggregates. The bulk-rock chemical composition of the different lithologies indicates that sodic amphibole and sodic pyroxene developed in Na- and Fe-rich systems or in a system with high Fe^{3+} Fe*. Thermodynamic modelling performed for different rock types (taking into account the measured Fe₂O₃ contents) reveals that sodic amphibole appears at \sim 8 ± 1 kbar and 400–450 °C (i.e. at the transition between the greenschist- and blueschist-facies conditions) about 5 kbar lower than previous estimates. To test the robustness of our conclusion, we performed a review of sodic amphibole compositions from a variety of terranes and P-T conditions. This shows (1) systematic variations of composition with P-T conditions and bulk-rock chemistry, and (2) that the amphibole compositions reported from the studied area are consistent with those reported from other greenschist- to blueschist-facies transitions.

Key words: sodic amphibole; Fe³⁺; blueschist–greenschist; Dent Blanche; Alps

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INTRODUCTION

The sodic amphibole glaucophane is an iconic mineral, because of the attractive blue colour, and because it classically defines the blueschist facies. Because glaucophane may occur in a wide range of bulk-rock compositions, its use for assessing pressure-temperature (P-T) conditions (e.g. Evans & Brown, 1986; Bucher & Grapes, 2011) requires great care in multicomponent chemical systems where the number of components exceeds the number of phases. Sodic amphiboles, with the generalized chemical formula $\Box Na_2(Mg,Fe_{2\perp})$ $3(AI,Fe_{3+})Si_8O_22(OH)2$ (where \Box represents a vacancy), display a wide range of compositions, among others owing to the substitutions of divalent as well as trivalent cations. Iron can occur in both di- and trivalent states, and the transition from the greenschist to the blueschist facies is therefore very sensitive to the oxidation state of the rock (i.e. to the ratio of Fe^{2+} and Fe^{3+}), in addition to the bulk-rock Na content. This has been clearly shown on an empirical basis by Miyashiro & Banno (1958) and Brown (1974, 1977). Further theoretical development by Liou et al. (1985) and experimental data by Maruyama et al. (1986) have definitely established this point.

With the advent of thermodynamic databases, the shift of the metamorphic facies boundaries, based on the presence of a sodic amphibole in a model system for metabasite, has been calculated by Evans (1990) (Fig. 1), and later confirmed by many others calculating phase diagrams for specific bulk-rock compositions (e.g. White et al., 2000, 2002; Fitzherbert et al., 2003, 2005; Warren & Waters, 2006; Diener et al., 2007; Groppo et al., 2009a; Diener & Powell, 2010; López-Carmona et al., 2013). For example, Diener & Powell (2010) have shown that for a model mid-ocean ridge basalt (MORB) composition at 450 °C, the first appearance of glaucophane occurs at \sim 9.3 kbar in an Fe³⁺-free system, but that P decreases to \sim 5.5 kbar with increasing Fe³⁺ content for a given, constant, total amount of Fe.

Despite these caveats, sodic amphibole is considered an important indicator of high-P, low-T conditions, and is often interpreted as relic of blueschist-facies metamorphism. However, without a proper chemical characterization of the sodic amphibole, P-T evaluations may be uncertain. An excellent example is provided by the Dent Blanche Tectonic System (DBTS) in the Western Alps (Fig. 2), where reports of sodic amphibole (e.g. Diehl, 1938; Stutz, 1940; Diehl et al., 1952; Ayrton et al., 1982; Roda & Zucali, 2008; Angiboust et al., 2014) have been taken for a long time as indicative of an early blueschist-facies stage (e.g. Oberhänsli et al., 2004; Berger & Bousquet, 2008; Beltrando et al., 2010; Bousquet et al., 2012). The scarce occurrence of sodic amphibole, and its peculiar Fe³⁺-rich chemistry (Ballèvre & Kienast, 1987; Canepa et al., 1990; Roda & Zucali, 2008; Manzotti et al., 2012, 2014a), may be explained by two different hypotheses.

The first hypothesis is that the *P*–*T* path of the DBTS may have reached the blueschist facies, but the blueschist-facies parageneses have been preserved in only a limited number of rocks because of the extensive greenschist-facies overprint (path A in Fig. 1; Roda & Zucali, 2008; Angiboust *et al.*, 2014). The second hypothesis is that the rocks may have reached much lower *P*, at the transition between greenschist and blueschist facies (Ballèvre & Kienast, 1987) (consistent with path B in Fig. 1). In this hypothesis, the development of sodic amphibole may have been limited to specific bulk-rock chemistries (e.g. Na-rich, Fe-rich or high fO_2).

In this paper, we test these two hypotheses. Following an extensive mapping campaign (Manzotti & Ballèvre, 2017; Manzotti et al., 2017), we studied the rare sodic amphibole-bearing rocks as well as a much larger number of samples that do not contain sodic amphibole. This allows us to determine the chemistry of both the amphibole and the bulk-rocks in which it occurs (or not), a prerequisite for attempting to model the relevant phase diagrams. The goal of the thermodynamic modelling is primarily the qualitative understanding of the behaviour of Na-amphibole at the greenschist-blueschist-facies transition in various rock types, rather than the determination of precise P-T conditions of the DBTS. We show that the sodic amphibole develops in a variety of rock types, including metasomatic layers at the contact between ultramafic

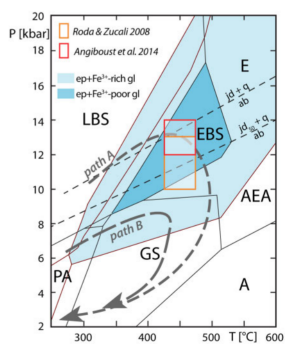


Fig. 1. Two possible *P*–*T* paths for the DBTS (Western Alps). Peak metamorphic conditions estimated in previous studies are reported (red and orange rectangles). Light and dark blue fields in the *P*–*T* diagram refer to the epidote blueschist-facies stability field calculated using two different compositions of sodic amphiboles, rich and poor in Fe^{3+} respectively (Evans, 1990). EBS, epidote-blueschist; LBS, lawsonite-blueschist; AEA, albite-epidote-amphibolite; A, amphibolite; E, eclogite; GS, greenschist; PA, pumpellyite-actinolite.

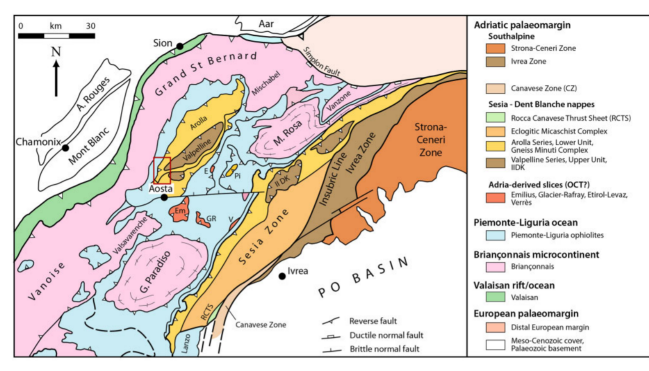


Fig. 2. Simplified structural map of the North Western Alps (modified after Manzotti *et al.*, 2014*a*, 2014*b*). The studied samples are from the frontal part of the Dent Blanche Tectonic System, north of Aosta (see red rectangle). Further details on the location of the samples may be found in Table 1 and in the Supplementary Data (Fig. S1).

cumulates and orthogneisses, at relatively low *P*; that is, at the transition from the greenschist to the blue-schist facies.

GEOLOGICAL SETTING

The Dent Blanche Tectonic System

The internal Western Alps comprise continental and oceanic units, which were assembled during the oblique convergence of Europe and the thinned Adriatic margin (Schmid et al., 2004; Handy et al., 2010). The Dent Blanche Tectonic System (DBTS) was derived from the Adriatic palaeomargin and, together with the Sesia Zone (Fig. 2), forms the Sesia-Dent Blanche nappes (e.g. Manzotti et al., 2014b, 2017). The DBTS mainly comprises polycyclic basement slices, bounded by Alpine shear zones (Manzotti et al., 2014a). The tectonometamorphic setting of the DBTS has been described in previous studies (e.g. Manzotti et al., 2014b, 2017). Therefore, only the structural and metamorphic features relevant for this study are hereafter summarized. The DBTS is made of two basement nappes, the Dent Blanche (DB) and the Mont Mary Cervino (MM) (see Manzotti et al., 2017, fig. 1), which are separated by the crustal-scale Roisan-Cignana Shear Zone (RCSZ; Manzotti, 2011; Manzotti et al., 2014a). Mesozoic metasediments deposited on top of the Palaeozoic basement are preserved in the Mont Dolin Series (Weidmann & Zaninetti, 1974; Ayrton et al., 1982) and the Roisan Zone (Manzotti et al., 2014b; Ciarapica et al., 2016).

The Valpelline Series in the DB and the Upper Unit in the MM are slivers of Permian lower continental crust and mainly consist of amphibolite- to granulite-facies metapelites, and mafic and metacarbonate rocks (Gardien *et al.*, 1994; Manzotti & Zucali, 2013; Dal Piaz *et al.*, 2015; Kunz *et al.*, 2018). The Arolla Series of the DB comprises Permian granitoids (e.g. Diehl *et al.*, 1952; Roda & Zucali, 2008; Manzotti *et al.*, 2018) and gabbros (e.g. Dal Piaz *et al.*, 1977; Monjoie *et al.*, 2007; Baletti *et al.*, 2012; Manzotti *et al.*, 2017) and their metamorphic products. The Lower Unit of the MM consists of pre-Alpine amphibolite-facies metapelites and minor granitoids, leucocratic gneisses and pegmatites (Canepa *et al.*, 1990; Dal Piaz *et al.*, 2010; Manzotti *et al.*, 2014*a*).

Alpine metamorphism in the DBTS

Alpine greenschist-facies metamorphism is dominant in the DBTS (e.g. Ayrton *et al.*, 1982). However, a few occurrences of sodic amphibole and sodic pyroxene may indicate early, blueschist-facies, metamorphism. For instance, metamorphic assemblages comprising sodic pyroxene and/or sodic amphibole have been described in the meta-igneous rocks, of acid, intermediate and basic compositions and in Mesozoic metasediments (see Supplementary Data Table S1 and references therein; Supplementary Data are available for downloading at http://www.petrology.oxfordjour nals.org).

Out of 17 studies, the bulk chemistry of the sodic amphibole-bearing rocks has been analysed in only two cases and the chemistry of sodic amphibole in five

Table	Table 1: Compilation of geographical coordinates and main petrc	oordinates and m	ain petrographical c	ographical characters of the studied samples	les	
Site	GPS coordinates	Rock type	Sample name	Preserved magmatic relice	Metamorphic minerals	
					Silicates	Non-silicates
-	45°53′13·29"N, 7°21′02·35"E	hornblendite	OL02*	am (hb-prq-ed)-Ti mt-ilm	Type 1: mrbk to wnc-ep-chl-ab-q-ttn; Type 2: ep-act-chl	ru-pv-hem
		hornblendite	OL03†	am (hb-prg-ed)-Ti mt-ilm	Type 1: mrbk to wnc-ep-chl-ab-q-ttn; Type 2: ep-act-chl	ru-py-hem
		hornblendite	OL04†	am (hb–prg–ed)–Ti mt–ilm	Type 1: mrbk to wnc-ep-chl-ab-q-ttn; Type 2: ep-act-chl	ru-py-hem
2	45°52'25.37"N, 7°20'28.71"E	orthogneiss	0L25‡	kfs–aln–zrn	q-ab-ep-mu-chl-ttn	py-hem
		orthogneiss	OL24‡	kfs–aln–zrn	q-ab-ep-mu-chl-ttn	py-hem
		orthogneiss	OL23#	kfs–aln–zrn	g-ab-ep-mu-chl-ttn	hem
		am-gneiss	OL50#	aln–zrn	q-ab-ep-mrbk-wnc-act	Ι
		am-gneiss	OL68A†	aln–zrn	q-ab-ep-ttn	mt-hem-py
		am-gneiss	OL68B*	aln–zrn	q-ab-ep-mrbk-wnc-act-aeg-aug-ttn	mt-hem
		am-gneiss	OL49C‡	aln–zrn	q-ab-ep-chl	Ι
		mu-rich layer	OL49B‡	Zrn	mu-ep	Ι
		chl-schist	OL49A‡	aln-zrn-ap	chl-ep±mu	I
	45°52′24.37"N, 7°20′28.71"E	ultramafic	OL35†		srp-Mg-chl-tr-ttn	cal-mt
		ultramafic	OL34†		srp-Mg-chl-tr-ttn	cal-mt
		ultramafic	0L33†		srp-Mg-chl-tr-ttn-tlc	cal-mt
ო	45°52'26.78"N, 7°20'25.97"E	am-gneiss	OL61‡	zrn–aln	g-ab-ep-mrbk-wnc-act	Py
			0L62†	zrn–aln	g–ab–ep–mrbk–wnc–act–ttn	I
			OL63†	zrn–aln	g-ab-ep-mrbk-wnc-act-ttn	I
4	45°52′27.29"N, 7°20′16.57"E	am-gneiss	0L21†	zrn–aln	g-ab-ep-mrbk-wnc-act	ру
	45°52′25.76"N, 7°20′24.64"E	ultramafic	OL60†		srp–Mg-chl–tr–ttn	cal-mt
പ	45°52′08·39"N, 7°19′44·77"E	am-gneiss	OL06†	aln–zrn	q–ab–ep–mrbk–wnc–act–ttn	ру
		orthogneiss	OL08#	kfs–aln–zrn	q-ab-ep-mu-chl-ttn	Ι
	45°52′13·58"N, 7°19′49·29"E	metagabbro	0L1424*		mu-zo-Mg-chl	Ι
9	45°51′43.82"N, 7°19′35.02"E	meta-tonalite	OL39†	zrn–aln–mnz–mu–q	q–ab–mrbk–wncaeg–aug–ep–ttn	ру
			OL40‡	zrn–aln–mnz–mu–q	q–ab–mrbk–wnc–aeg–aug–ep–ttn	Ι
			OL43*	zrn–aln–mnz–mu–q	q-ab-mrbk-wnc-act-aeg-aug-ep-ttn	I
			OL44†	zrn–aln–mnz–mu	q–ab–mrbk–wnc–aeg–aug–ep–ttn	ру
		meta-tonalite	OL42A†	zrn–aln–mnz–q	g-ab-mrbk-chl	:
	45°51'43.82"N, 7°19'35.02"E	chl-schist	OL42B‡	zrn-ap-mnz	Mg-chl	I
		ultramafic	OL54†		srp-Mg-chl-ttn	mt
		ultramafic	OL55†		srp-mt	mt
7	45°47′13·38"N, 7°19′01·57"E	blueschist	VP7*	zrn–aln–ap	gl-mrbk-mu-ep-chl-ttn-q-g	ру
	M. Morion (Diehl <i>et al.</i> 1952)	orthogneiss	14§		q-ab-ep-mu-chl-ttn	I
	-	-	1			

*Whole-rock chemistry + microprobe + thermodynamic modelling. †Whole-rock chemistry. ‡Whole-rock chemistry + microprobe. §Thermodynamic modelling.

cases only (Table S1), indicating mostly magnesioriebeckite and rarely glaucophane. Although some researchers described aegirine-augite in the DBTS (e.g. De Leo *et al.*, 1987; Pennacchioni & Guermani, 1993), not a single chemical analysis of sodic pyroxene has been reported.

For the Arolla orthogneisses, peak *P*–*T* conditions of 12–14 kbar and ~450 °C have been estimated on the basis of conventional thermobarometry and phase equilibria (e.g. Roda & Zucali, 2008). A pseudosection approach has been used to constrain peak and retrograde evolution of Arolla orthogneisses devoid of sodic amphibole (Angiboust *et al.*, 2014; Kirst, 2017), and it returned similar *P*–*T* conditions. For the Lower Unit of the Mont Mary, thermobarometric calculations have provided an estimate at 13 ± 2 kbar and 480 ± 50 °C (Manzotti *et al.*, 2014a).

METHODS

Out of a collection of a few hundred samples from the DBTS, we found 16 samples from seven localities in which sodic amphibole and/or sodic pyroxene have been observed (Figs 3 and 4, Table 1, Supplementary Data Fig. S1). The bulk-rock composition of all these samples has been analysed (with the exception of sample OL44, because it was too altered) and the mineral chemistry has been determined in detail in eight samples. Our study also includes 14 samples that do not contain sodic amphibole and/or pyroxene (Table 1). The bulk-rock chemistry has been obtained for the entire set of samples and the mineral chemistry has been studied in detail in six samples. Thermodynamic modelling has been applied on six samples, four of which contain sodic amphibole and/or sodic pyroxene, and two of which are devoid of these minerals. The variety of rock types and provenance of the selected samples allow us to evaluate the geochemical control (protolith bulk composition versus successive fluid-rock interactions) on the development of sodic amphibole and pyroxene.

Mineral chemistry

Samples were investigated by optical petrography in polished thin sections (Fig. 5) and back-scattered electron (BSE) imaging by scanning electron microscopy. Mineral abbreviations and symbols used in this study are listed in Supplementary Data Table S2. For practical purpose, muscovite is used for potassic white mica, whatever its Si content, because there is no formal definition of phengite and because there is no miscibility gap between muscovite and celadonite.

A JEOL JXA-8530F electron probe microanalyzer at the Institute of Earth Science of the University of Lausanne (Switzerland) was used to acquire quantitative analyses of mineral phases and X-ray compositional maps in wavelength-dispersive mode. Spot analyses were performed using an accelerating voltage of 15 keV. A current of 20 nA was used for garnet and clinopyroxene spot analysis. To avoid electron-beaminduced decomposition of minerals containing volatile elements, a current of 15 nA was applied for plagioclase, amphibole, chlorite, muscovite and epidote quantitative measurements. For these latter phases, a beam size ranging between 2 and 5 μ m was used depending on the size of the mineral. The X-ray compositional mapping was carried out at 50 ms on 0-7 μ m pixel, using an acceleration voltage of 15 keV and a current of 20 nA.

To evaluate the oxidation state of the studied samples, the oxide minerals present (magnetite and hematite) were identified optically and confirmed by Raman spectroscopy (Horiba HR Raman-FTIR spectrometer from HORIBA Scientific, an integrated Raman microprobe consisting of an Olympus BX41 confocal microscope coupled to an 800 mm focal-length spectrograph, Institute of Earth Sciences, University of Lausanne).

Because the microprobe analyses do not readily distinguish Fe^{2+} and Fe^{3+} , and because this is a fundamental aspect of our study, the procedure followed is explained in more detail below. In pyroxene, the Fe³⁺ content has been calculated by stoichiometry. Structural formulae for this mineral have been calculated on the basis of six oxygens and four cations. The recalculation procedure for amphibole has been discussed in detail by Schumacher (1997, 2007). For each analysis, a minimum, a maximum (and a mean) estimation of the Fe^{3+} can be obtained. The influence of using one (or the other) calculation scheme is displayed for some analyses in Figure 6 [i.e. the diagram proposed by Miyashiro & Banno (1958) and adopted later in the classification of sodic amphiboles, except for crossite (Leake et al., 1997)]. Specifically, we reported as large symbols the recalculation giving the maximum amount of Fe^{3+} [generally on O = 23 and sum of cations except Ca, Na, K = 13 for sodic amphiboles; see Schumacher (2007)], and the two other estimations (i.e. the minimum and the mean) are plotted in the classification diagram as a thin line joining the highest estimate to the two other ones (Fig. 6). For consistency, we plotted all amphibole analyses with a maximum amount of Fe^{3+} . This has some influence on their position in the diagram, but the uncertainty introduced by the recalculation scheme is negligible (i.e. very short lines) in most cases or moderate for a few cases. Some amphibole analyses may fall into different fields according to which calculation scheme is adopted. However, because there is no miscibility gap between the different end-members plotted in Miyashiro's diagram, the important point is not the name of the amphibole (the different fields have arbitrary limits), but the location of the analyses in the diagram, whichever of the four fields they fall in.

Representative mineral analyses are reported in the Supplementary Data (Tables S3–S10). The main chemical characteristics of sodic amphibole and sodic pyroxene are given in Table 2.

	Sample	Mineral	Si	$(Na + K)_A$	$X_{\sf Na}$	Na _B	X_{Fe}	$X_{\rm Fe3}$	n
Hornblendite	OL02	Prg	6.39–6.48	0.50-0.61	0.20-0.22		0.29-0.35	0.00-0.05	15
		Ed	6.55-7.53	0.39-0.54	0.23-0.37		0.36-0.42	_	4
		Mg-hb	6.84–7.32	0.17-0.49	0.21-0.24		0.28-0.48	0.09-0.32	5
		mrbk	7.93-8.00	0.00-0.03	0.75–0.91	1.51–1.82	0.38-0.46	0.64-0.71	10
		wnc	7.53–7.97	0.00-0.04	0.23-0.75	0.53-1.49	0.27-0.46	0.41-0.72	34
		act	7.65–7.98	0.00-0.07	0.07-0.26		0.24-0.37	0-0.70	25
Metatonalite	OL40	mrbk	7.90–8.00	0.00-0.02	0.76–0.97	1.53–1.94	0.15–0.43	0.46–0.87	61
		wnc	7.90-8.00	0.00-0.02	0.26-0.74	0.55-1.49	0.22-0.35	0.62-0.87	10
	OL42	wnc	7.86-8.00	0.06-0.16	0.18-0.52	0.53–1.14	0.20-0.27	0.00-0.63	6
	OL43	mrbk	7.94-8.00	0.00-0.03	0.75-0.97	1.51–1.93	0.26-0.44	0.58-0.73	19
		wnc	7.87–8.00	0.00-0.04	0.27-0.74	0.51–1.48	0.24-0.40	0.57-0.72	21
		act	7.89-8.00	0.00-0.01	0.04-0.25		0.24-0.35	0-0.55	20
	OL44	mrbk	7.83–7.99	0.00-0.04	0.75-0.90	1.51–1.79	0.31-0.42	0.52-0.68	36
		wnc	7.64–7.98	0.00-0.11	0.22-0.77	0.54-1.48	0.21-0.44	0.42-0.70	32
Blueschist	VP7	gl	7.87–8.00	0.00-0.07	0.94-0.99	1.87–1.98	0.39–0.50	0.22-0.50	42
		mrbk	7·11–8·00	0.00-0.08	0.87-0.97	1.73–1.94	0.27-0.51	0.45-0.68	29
Am-gneiss	OL50	gl	7.98–7.99	0.00-0.01	0.95-0.96	1.90–1.92	0.37–0.38	0.45-0.49	5
		mrbk	7.93-8.00	0.00-0.04	0.80-0.95	1.60–1.89	0.32-0.39	0.52-0.64	7
		wnc	7.75–7.96	0.01–0.05	0.28-0.69	0.55–1.39	0.25-0.32	0.62-0.72	34
		act	7.81–7.90	0.01-0.03	0.12-0.22		0.24-0.30	0.51-0.70	8
	OL61	mrbk	7.94	0.00	0.85	1.60	0.44	0.62	1
		wnc	7.84–7.99	0.00-0.06	0.21-0.65	0.51–1.30	0.20-0.37	0.45-0.72	55
		act	7.75–7.95	0.00-0.04	0.15–0.24		0.18-0.43	0.51–0.68	8
	OL68	gl	7.81–7.99	0.00	0.87-0.98	1.74–1.96	0.31-0.40	0.36-0.47	6
		mrbk	7.79–7.97	0.00-0.12	0.65-0.96	1.56–1.91	0.22-0.41	0.48-0.74	20
		wnc	7.50-7.93	0.00-0.19	0.30-0.86	0.52-1.74	0.19–0.51	0.27-0.73	23
		act	7.85–8.00	0.03–0.17	0.11–0.29		0.40-0.54		10
	Sample	Mineral	Jd	Aeg	n				
Metatonalite	OL43	aeg–aug	0.13–0.26	0.30-0.42	58				
	OL44	aeg-aug	0.04-0.34	0.23-0.44	75				
Am-gneiss	OL68	aeg-aug	0.07-0.24	0.29-0.43	30				

Table 2: Main chemical characters (major element compositions) of amphibole and pyroxene

 $X_{Na} = Na/(Na + Ca); X_{Fe} = Fe^{2+}/(Fe^{2+} + Mg); X_{Fe3} = Fe^{3+}/(Fe^{3+} + AI); n$, number of analyses.

Bulk-rock chemistry

To assess the nature and the degree of chemical modification in the rocks displaying sodic amphibole and/or sodic pyroxene in the DBTS, we analysed the major and trace element compositions of the selected set of samples (Table 4 and Tables S11-S16). We performed two or three separate bulk-rock analyses for samples that macroscopically showed different domains (e.g. OL49), to obtain the most representative chemical composition of each domain. Major elements were determined by inductively coupled plasma atomic emission spectrometry (ICP-AES; SARM-CRPG, Nancy). Bulk-rock glasses were fused at 980 °C by mixing appropriate proportions (1%) of fine-grained rock powder with di-lithium tetraborate. Then, glasses were dissolved in a mixture of HNO_3 (5%), H_2O_2 (0.5%) and glycerol (10%) prior to analysis. Details about the method used for the analyses have been given by Carignan et al. (2001). Uncertainties at 1σ are c. 5% for the ICP-AES data. In addition, separate analyses of Fe²⁺ content of wholerock samples were performed (SARM-CRPG, Nancy, France) by titration with potassium dichromate after dissolution of the sample in a HF-H₂SO₄ mixture, in the presence of H₃BO₃ and H₃PO₄.

Phase diagram calculations

P-T pseudosections were calculated with Theriak/ Domino (de Capitani & Brown, 1987; de Capitani & Petrakakis, 2010), the internally consistent thermodynamic dataset ds6.2 (Holland & Powell, 2011) and mixing models for solid solutions (converted for Theriak/Domino by D. K. Tinkham, http://dtinkham.net/ peq.html).

Phase relations were modelled in the chemical sys-Na₂O-CaO-FeO-MgO-Al₂O₃-SiO₂-H₂O-TiO₂tem $Fe_2O_3 \pm MnO \pm K_2O$ (MnNCKFMASHTO). In the first step, H₂O was considered in excess in all calculations. The phases considered in the calculations and the activity composition models are clinoamphibole and clinopyroxene (Green et al., 2016), chlorite, muscovite, garnet, biotite (White et al., 2014a, 2014b), plagioclase (Holland & Powell, 2003), epidote (Holland & Powell, 2011), ilmenite (White et al., 2000), magnetite (White et al., 2002), pumpellyite (ideal mixing on sites). Pure phases comprise quartz, titanite, and rutile. The activity-composition (a-x) relations (i.e. White et al., 2014a, 2014b) used with the internally consistent thermodynamic dataset of Holland & Powell (2011) include Fe³⁺ in several silicate phases (e.g. garnet, chlorite, biotite, staurolite, chloritoid, muscovite, paragonite, and margarite).

P–*T* pseudosections were calculated using the amount of Fe^{2+} (versus Fe^{3+}) measured in the bulk composition. As rocks can be affected by surface weathering processes including oxidation, the measured amount of Fe^{3+} is considered as the maximum value on

the range of possible Fe^{3+} contents (Diener & Powell, 2010). $P-X(Fe_2O_3)$ (expressed in wt%) pseudosections were calculated to explore the effect of variable Fe^{2+}/Fe^{3+} ratio in the bulk-rock. In these calculations, *T* was generally fixed at 450 °C, a value that, according to previous studies (e.g. Roda & Zucali, 2008; Angiboust *et al.*, 2014; Kirst, 2017) corresponds to the *T* reached by the DBTS at peak *P* conditions. In the diagrams, the abbreviation 'gl' has been used for sodic amphibole, whereas clinopyroxene is indicated with the abbreviation 'cpx'.

P-H₂O (expressed in wt%) diagrams were calculated to explore the effect of variable bulk-rock H₂O on phase equilibria and P-T estimates. These diagrams were constructed for the same bulk compositions as used for the P-T diagrams (Fig. 14d and Figs S4–S5), for a T of 450 °C and a P range of 4–12 kbar.

PETROGRAPHY AND MINERAL CHEMISTRY

In the DBTS, the dominant rock types are felsic orthogneisses of granitic to dioritic composition (the Arolla orthogneisses), paragneisses interpreted as the country-rocks of these intrusions (Manzotti et al., 2018), and a few metagabbroic complexes, displaying ultramafic cumulates, and mesocratic and leucocratic metagabbro (e.g. the Berrio Gabbro, Manzotti et al., 2017). All these rock types are devoid of sodic amphibole and sodic pyroxene. These two minerals are, however, found in minor rock types that belong to two main groups. The first one does not show obvious signs of metasomatism and comprises (1) a single occurrence of hornblendite in the Berrio Gabbro, (2) a few metatonalite pods within ultramafic cumulates from the Berrio Gabbro, and (3) rare blueschists interlayered with felsic orthogneisses (Manzotti et al., 2014a). The second group consists of amphibole-gneisses, located in reaction zones, at the boundary between the Arolla orthogneisses and the ultramafic cumulates of the Berrio Gabbro (Figs 3 and 4). These four types of rocks are described in more detail below. For comparison, we also studied in detail two rocks that contain neither sodic amphibole nor sodic pyroxene; namely, a leucocratic metagabbro and an Arolla orthogneiss.

Hornblendite lenses in the Berrio Gabbro

Decametre-long lenses of unfoliated hornblendite are found included in the mylonitic Berrio metagabbro (Supplementary Data, Fig. S1). Based on differences in their mineral assemblages, two domains may be distinguished at the microscale. The type-I domain is dominated by magmatic calcic amphiboles (plotting in the pargasite, edenite and magnesio-hornblende fields, X_{Fe} = 0·28–0·48), up to 1 mm in size and partially overgrown by coronas of blue sodic and sodic–calcic amphiboles (Mg-riebeckite, X_{Fe} = 0·38–0·46, Na_B = 1·51–1·82 p.f.u., and winchite, X_{Fe} = 0·27–0·46, Na_B = 0·53–1·49 p.f.u., respectively) (Figs 5a and 6). Epidote, chlorite, albite (<5%) and quartz (~5%) are also present and accessory 7

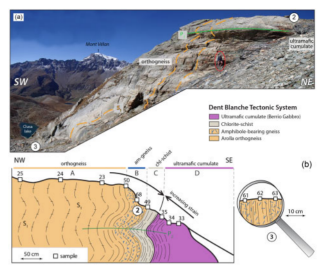


Fig. 3. (a) Contact between the Arolla orthogneiss and ultramafic cumulate from the Berrio Gabbro. The S₁ foliation in the orthogneiss and the contact between the two lithologies are affected by metre- to decametre-scale open folds. Sites 2 ($45^{\circ}52'25\cdot37^{"}N$, $7^{\circ}20'28\cdot71^{"}E$) and 3 ($45^{\circ}52'26\cdot78^{"}N$, $7^{\circ}20'28\cdot71^{"}E$) and 3 ($45^{\circ}52'26\cdot78^{"}N$, $7^{\circ}20'25\cdot97^{"}E$) are indicated in the photograph. (b) Schematic cross-section of the contact between the Arolla orthogneiss and ultramafic cumulate from the Berrio gabbro. Strain increases in both lithologies towards the contact. White squares indicate sample locations and numbers refer to the sample names.

minerals (<1%) comprise pyrite, hematite and magnetite. Fine-grained intergrowths of rutile and Ti-hematite surrounded by titanite coronas are common. Rutile and Ti-hematite are interpreted to be pseudomorphs resulting from the oxidation of original intergrowths of titanomagnetite + ilmenite, as observed in other mafic rocks (e.g. Haggert, 1991). The type-II domain is dominated by fine-grained aggregates of epidote replacing previous magmatic plagioclase and by aggregates of actinolite ($X_{\text{Fe}} = 0.24-0.35$) and chlorite. Epidote shows significant compositional zoning, with an Fe³⁺/Al³⁺ + Fe³⁺ ratio in the core of 0.09–0.20 and in the rim of 0.26– 0.32 (i.e. an almost pure pistacite) (Fig. S2).

To sum up, in sample OL02, used below for the thermodynamic modelling, the observed mineral paragenesis in the type-I domain comprises sodic and sodic-calcic amphibole (magnesioriebeckite to winchite), epidote, chlorite, quartz and titanite.

Metatonalite pods within the Berrio ultramafic rocks

Rare, medium-grained metatonalites are found as $\sim 2 \text{ m}$ long and $\sim 30 \text{ cm}$ thick layers and metre-scale pods within the massive ultramafic rocks (mainly antigorite serpentinite) of the Berrio Gabbro close to the Alpe Berrio, along and above the main path for the Col de Breuson (Fig. S1). Metatonalite pods are massive in appearance with a dense network of millimetre- to centimetre-sized patches of radiating blue and greenish amphiboles (Fig. 4a and b). Metatonalite layers locally display a weak foliation, defined by blue amphiboles and layers of fine-grained fibrous aggregates of sodic pyroxene. Large crystals of quartz show undulose extinction, irregular grain boundaries and tiny fluid inclusions aligned along fracture planes. Large albite crystals locally displaying polysynthetic twinning may contain rare corroded muscovite flakes (up to 2 mm in size, Si = 3.12-3.18 p.f.u.). Quartz, albite and muscovite, as well as accessory monazite and zircon, are

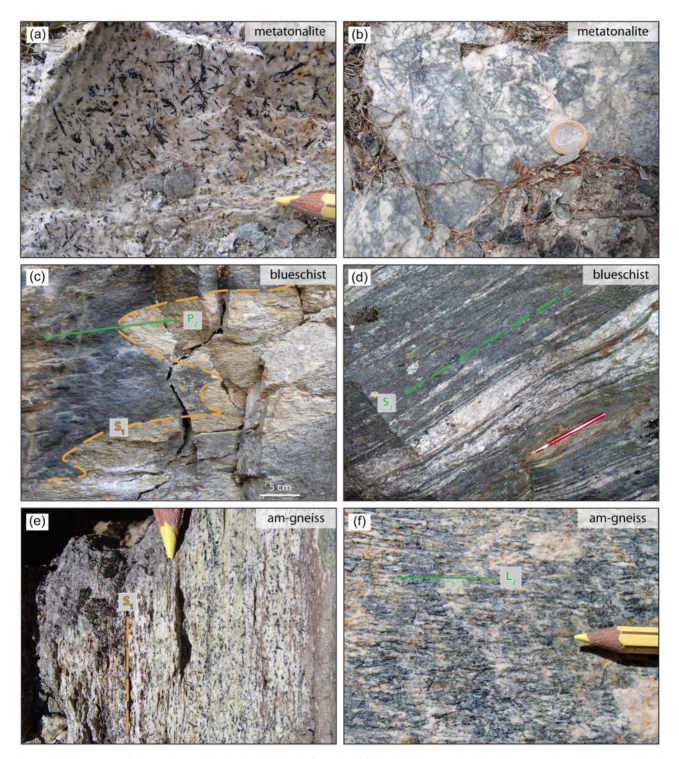


Fig. 4. Field aspects of the sodic amphibole-bearing rocks from the DBTS. (a) Metatonalite displaying needles of blue amphibole dispersed in an albite–quartz matrix (site 6, $45^{\circ}51'43\cdot82$ "N, $7^{\circ}19'35\cdot02$ "E). (b) Network of millimetre- to centimetre-sized patches of radiating blue and greenish amphiboles in a metatonalite pod (site 6, $45^{\circ}51'43\cdot82$ "N, $7^{\circ}19'35\cdot02$ "E). (c) Blueschist folded isoclinally together with leucocratic aplitic lenses (site 7, $45^{\circ}47'13\cdot38$ "N, $7^{\circ}19'01\cdot57$ "E). (d) In the blueschist, S₂ foliation is marked by epidote and quartz domains and by films of muscovite, chlorite and blue amphibole (site 7, $45^{\circ}47'13\cdot38$ "N, $7^{\circ}19'01\cdot57$ "E). (e, f) Needles of blue amphiboles in the amphibole gneiss (site 2): (e) crosscutting the mylonitic foliation S₁ ($45^{\circ}52'25\cdot37$ "N, $7^{\circ}20'28\cdot71$ "E) and (f) the crenulation lineation L₂ ($45^{\circ}52'26\cdot78$ "N, $7^{\circ}20'25\cdot97$ "E).

interpreted as relics of magmatic phases. The finegrained recrystallized matrix consists of quartz, albite, blue amphibole, locally green clinopyroxene (Fig. 5b and c) and minor epidote, titanite and pyrite (<1%). Clinopyroxene is mainly aegirine–augite (jd_{25} –aeg₃₅– quad₄₀, sample OL43; Figs 5b and 7), locally with patchy zoning (from jd_5 –aeg₂₅–quad₇₀ to jd_{30} –aeg₃₀–quad₄₀, sample OL44; Fig. 7b). Blue and green amphibole form

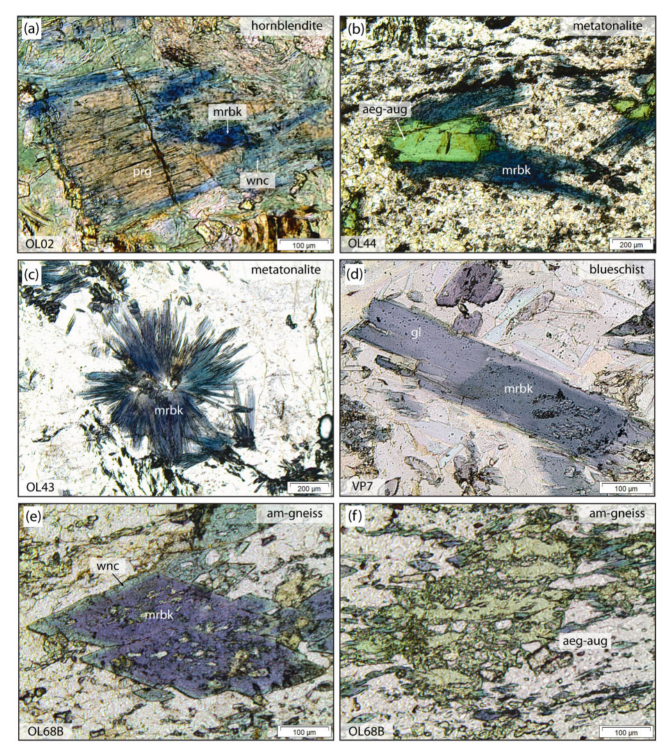


Fig. 5. Mineralogy and textures of the blue amphibole-bearing rocks from the DBTS. Plane-polarized light (PPL) photomicrographs. (a) Coronas of magnesioriebeckite and winchite around magmatic pargasite in the hornblendite (sample OL02, site 1). (b) Aegirine-augite surrounded by magnesioriebeckite in the metatonalite from the Berrio gabbro (sample OL44, site 6). (c) Needles of magnesioriebeckite in the albite matrix of a metatonalite from the Berrio gabbro (sample OL43, site 6). (d) A dark blue core of magnesioriebeckite surrounded by a pale blue glaucophane rim in blueschist (sample VP07, site 7). (e) Magnesioriebeckite with winchite rim in the amphibole-gneiss at the contact with ultramafic cumulate of the Berrio Gabbro (sample OL68, site 2). (f) Aegirine-augite crystals from the blue amphibole-gneiss at the contact with ultramafic cumulate of the Berrio Gabbro (site 2, sample OL68).

together radiating sheaves in an albite matrix. In these sheaves, blue amphibole is dominant over green amphibole. However, there is no textural argument allowing us to establish whether sodic or calcic amphiboles appear first, and thus they are considered to be stable together. Blue amphibole is slightly zoned with a darker blue core (magnesioriebeckite $X_{Fe} = 0.15-0.46$, Na_B = 1.51-1.94 p.f.u.) and a lighter blue rim of winchite ($X_{Fe} = 0.20-0.44$, Na_B = 0.51-1.49 p.f.u.) (Fig. 6). The greenish amphibole found together with magnesioriebeckite and winchite is actinolite ($X_{Fe} = 0.24-0.35$).

A 10–20 cm thick dark greenish grey band of chloriteschist marks the contact between the metatonalite and the ultramafic rocks and is considered a reaction zone between the two contrasting rock types ('the blackwall'). The transition from the metatonalite to the chloriteschist is defined by the progressive increase in the modal abundance of chlorite and the disappearance of albite (Fig. S3a). In the chlorite-schist, chlorite is zoned, with Mg content increasing from core ($X_{Mg} = 0.77-0.79$) to rim ($X_{Mg} = 0.80-0.83$). In the chlorite-schist, accessory phases include abundant zircon and rare apatite and monazite, with the latter wrapped by chlorite foliation.

To sum up, in sample OL43, used below for the thermodynamic modelling, the observed mineral paragenesis comprises quartz, albite, sodic amphibole (magnesio-riebeckite to winchite), calcic amphibole

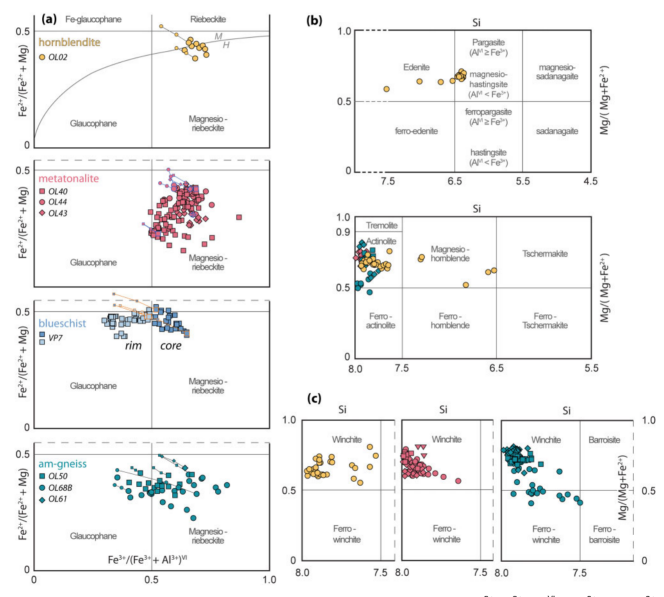


Fig. 6. Compositional diagrams for amphibole. (a) Na-amphibole compositions plotted in the $Fe^{3+}/(Fe^{3+} + AI^{VI})$ vs $Fe^{2+}/(Mg + Fe^{2+})$ diagram. The large symbols correspond to the recalculation giving the maximum amount of Fe^{3+} . The other estimations (i.e. the minimum and the mean) are plotted in the classification diagram as a fine line joining the highest estimate to the two others. The grey line marked by the letters H and M corresponds to the fO_2 values of hematite–magnetite buffer, as reported by Okay (1980). (b) Ca-amphibole compositions plotted in the Si (p.f.u.) vs Mg/(Mg + Fe²⁺) diagram [same legend as in (a)]. (c) Na–Ca-amphibole compositions plotted in the Si (p.f.u.) vs Mg/(Mg + Fe²⁺) diagram [same legend as in (a)].

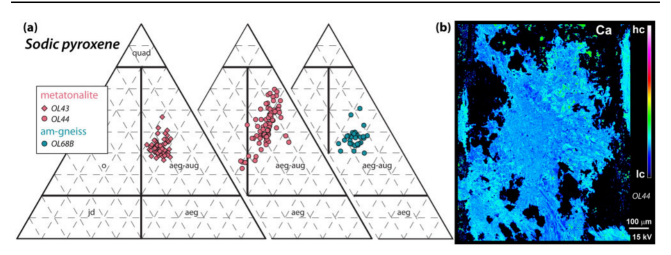


Fig. 7. Compositional diagrams for clinopyroxene. (a) Aegirine–augite compositions plotted in the jd–quad–aeg diagram. (b) Ca X-ray map of aegirine–augite from metatonalite (sample OL44).

(actinolite), clinopyroxene (aegirine-augite), epidote and titanite.

Blueschist

North of Roisan village, a rather poor outcrop of dark blue fine-grained schist occurs along the path rising from Gorrey to Berrio Nemoz. The schist is isoclinally folded together with leucocratic aplitic layers (10-20 cm thick) consisting of quartz, albite and muscovite (Fig. 4c and d). The schist comprises blue amphibole, muscovite, epidote, chlorite, guartz, titanite, rare spessartinerich garnet (alm₂₀₋₃₁py₁gr₂₁₋₂₂spss₄₇₋₅₃andr₀₋₅), apatite, zircon, and pyrite (<1%). It displays a crenulation cleavage (S_2) with relicts of an earlier foliation (S_1) marked by blue amphibole and muscovite preserved in the microlithons. The new foliation (S_2) is defined by epidote $(Fe^{3+}/Al^{3+} + Fe^{3+}$ ranges from 0.19–25 in the core to 0.28-0.32 in the rim; Fig. S2) and quartz domains, muscovite (Si = 3.36-3.42 p.f.u.) and chlorite (X_{Mg} = 0.48-0.51) films, and by blue amphibole. Muscovite contains some Fe³⁺ as analyses plot above the ideal celadonite substitution (Fig. 8a). Blue amphibole (1-3 mm long) shows a dark blue core (magnesioriebeckite, $X_{\text{Fe}} = 0.27 - 0.51$, Na_B = 1.73 - 1.94 p.f.u.) and light blue rim (glaucophane, $X_{Fe} = 0.39-0.50$, Na_B = 1.87-1.98 p.f.u.) and tiny inclusions of epidote and guartz (Figs 5d and 6).

To sum up, in sample VP7, used below for the thermodynamic modelling, the observed mineral paragenesis comprises sodic amphibole (magnesioriebeckite to glaucophane), muscovite, epidote, chlorite, quartz, titanite, and garnet.

Amphibole-gneiss at the boundary between orthogneiss and ultramafic

Sodic amphibole-bearing rocks are localized along the contact between the Arolla orthogneisses and the ultramafic rocks of the Berrio Gabbro. This contact, sharply defined at map to sample scale, is best exposed north

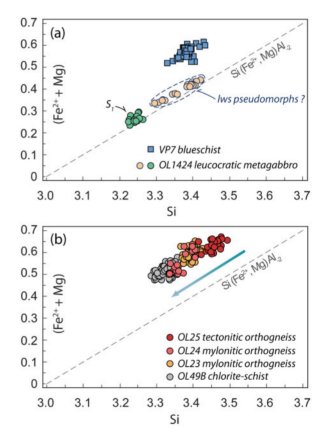


Fig. 8. Potassic white mica composition plotted in the Si vs $Fe^{2+} + Mg$ (p.f.u.) diagram from the leucocratic metagabbro and blueschist (a) and from samples along the profile of site 2 (b). The dashed line represents the ideal celadonitic substitution.

of the Clusa Lake (Fig. 3). There, three main lithological domains are distinguished, from the NW to the SE.

(1) The granitic orthogneisses display a strain gradient towards the contact with the ultramafic rocks (Fig. 3b): foliated orthogneisses with few magmatic feldspar and allanite porphyroclasts evolve into mylonites with a fine banded structure. Magmatic accessory phases include zircon, allanite, and titanite. The mylonitic foliation S_1 is defined by whitish albite-quartz bands alternating with yellowish-greenish discontinuous layers of epidote (Fe³⁺/Al³⁺ + Fe³⁺ = 0.16–0.23 in the core and 0.26–0.32 in the rim; Fig. S2), chlorite and muscovite (Si = 3.42-3.50 p.f.u., 3.34-3.41 p.f.u., 3.35-3.45 p.f.u. in OL25, OL24 and OL23 respectively). Analyses of muscovite plot above the line of ideal celadonite substitution, indicating the presence of Fe³⁺ (Fig. 8b).

(2) A reaction zone, about 0.5 m thick, is observed along the contact between the Arolla orthogneisses (A in Fig. 3b) and the Berrio ultramafic rocks (D in Fig. 3b). The reaction zone (B and C in Fig. 3b) displays two types of lithologies.

The first one is an ${\sim}10\text{--}20\,\text{cm}$ thick zone of amphibole-gneiss, yellowish-greenish in colour, sometimes showing a centimetre-scale layering. It is characterized by small needles of blue amphibole and rare pale green clinopyroxene, both displaying numerous quartz inclusions (B in Fig. 3b). Only locally are the blue amphibole needles aligned parallel to the stretching lineation L₁. In most cases, blue amphibole defines rosettes or sheaves (up to 3 mm in size), which are dispersed in a quartz-albite-rich matrix, generally without a shape-preferred orientation (Fig. 4e). Growth of these rosettes may be contemporaneous with, or later than, the crenulation lineation L_2 (Fig. 4f). Blue amphiboles are slightly zoned, with darker blue cores and lighter blue or greenish rims, whatever their structural position (Fig. 5e). Cores are magnesioriebeckite and rarely glaucophane ($X_{Fe} = 0.22-0.41$ and 0.31-0.40, respectively; Na_B = 1.56-1.91 and 1.90-1.92 p.f.u. respectively), whereas rims are winchite and rarely actinolite ($X_{Fe} =$ 0.18-0.51 and 0.40-0.54, respectively) (Fig. 6). Pale green clinopyroxene (Fig. 5f) is aegirine-augite (jd₂₅aeg₃₅-quad₄₀, sample OL68) (Fig. 7). Epidote shows significant compositional zoning, with an $Fe^{3+}/AI^{3+} + Fe^{3+}$ ratio of 0.15-0.23 in the core and of 0.25-0.32 in the rim (Fig. S2). A few titanite crystals containing tiny rutile grains are dispersed in the rock, together with idioblastic crystals of hematite, magnetite and pyrite (<1%).

Second, an \sim 10 cm thick layer of dark greenish grey fine-grained chlorite-schist (C in Fig. 3b) is found against the ultramafic rocks. At the interface between the amphibole-gneiss and the chlorite-schist, an \sim 3 mm thick layer of randomly oriented muscovite flakes (Si = 3.30-3.39 p.f.u.) is observed (Fig. S3b). Muscovite contains some Fe³⁺ because analyses plot above the line of ideal celadonite substitution (Fig. 8b). The chloriteschist is essentially made of zoned chlorite, with Mg content increasing from core ($X_{Mg} = 0.67-0.79$) to rim $(X_{Mg} = 0.82 - 0.86)$ (Fig. S3b). The foliation is defined by the shape fabric of chlorite and by aligned epidote crystals ($X_{Ps} = 0.25-0.29$; Fig. S2). The former presence of porphyroclasts is recorded by the presence of chlorite aggregates [after plagioclase(?), e.g. Müntener et al., 2010] wrapped by the foliation. In the chlorite-schist,

accessory phases include numerous zircon grains, apatite and a few allanite crystals.

(3) The Berrio ultramafic rocks (wehrlite; see Manzotti et al., 2017, for details) also display a strain gradient towards the contact with the orthogneiss. Unfoliated rocks that preserve their primary magmatic texture progressively evolve to mylonitic serpentinites (a few centimetres thick; Fig. 3b). In the undeformed domain, the rock comprises a serpentine mesh with magnetite (interpreted as pseudomorphs after olivine), and Cr-magnetite and Mg-chlorite (interpreted to replace Crspinel). Fine-grained aggregates of tremolite-calcite-Mg-chlorite-titanite are interpreted to pseudomorph clinopyroxene. The intercumulus phases are completely recrystallized into actinolite-tremolite. In the mylonitic serpentinites, a pervasive foliation, oriented parallel to the contact with the orthogneisses, is defined by an alternation of serpentine, chlorite, and magnetite layers. Talc occurs as veins crosscutting or parallel to the main foliation. The modal proportion of talc increases towards the contact.

To sum up, in sample OL68B, used below for the thermodynamic modelling, the observed mineral paragenesis comprises quartz, albite, epidote, sodic amphibole (magnesioriebeckite to winchite), clinopyroxene (aegirine–augite), and titanite.

Leucocratic metagabbro

Leucocratic metagabbro occurs as decametre-thick layers in the Berrio Gabbro and is always mylonitic. Examples can be observed in the Col Cornet-Monte Berrio area, in the western portion of the mafic complex, along the contact with the Arolla orthogneiss (Fig. S1). The inferred original assemblage plagioclase +clinopyroxene + minor spinel is replaced by Alpine metamorphic minerals. The rocks show a pervasive Alpine mylonitic foliation S_1 , marked by tiny crystals of muscovite (Si = 3.22-3.26 p.f.u.), and Mg-chlorite (X_{Mg} = 0.76-0.90) (replacing magmatic clinopyroxene), alternating with zoisite layers (replacing magmatic plagioclase). The few rectangular-shaped aggregates consisting of muscovite (Si = $3 \cdot 30 - 3 \cdot 42$ p.f.u.) and finegrained clinozoisite and Mg-chlorite may represent pseudomorphs after former lawsonite. The fit of the composition of muscovite with the theoretical line of the Si(Fe,Mg)Al_2 substitution suggests that the ${\rm Fe}^{3+}$ content is negligible (Fig. 8a). Locally, clinozoisite layers wrap around plagioclase porphyroclasts (now consisting of pure albite). Green slightly elongated patches up to 2 mm in size are scattered in the leucocratic gabbro. In thin section they consist of Cr-rich muscovite ('fuchsite'; Cr content up to 6.6 wt%) and chromian pumpellyite (Cr content up to 16.40 wt%) and are interpreted to replace Cr-rich spinel. Sodic amphibole has never been observed in the leucocratic metagabbro.

To sum up, in sample OL1424, used below for the thermodynamic modelling, the Cr-bearing phases (i.e. fuchsite and pumpellyite) are absent and the observed

synfolial mineral assemblage is epidote, chlorite, muscovite, albite, and quartz.

A summary of the relationships between deformation and growth in the studied samples (Table 3) shows that the sodic amphiboles grew at peak P-T conditions and eventually recrystallized after the mylonitic deformation in the amphibole-gneiss.

WHOLE-ROCK GEOCHEMISTRY

Hornblendite lenses in the Berrio Gabbro

Hornblendites (samples OL02, OL03 and OL04) are ultramafic rocks (SiO₂ 36·41–42·08 wt%), rich in Fe₂O₃* (13·01–21·92 wt%) and low in alkalis (0·39–2·61 wt%) (Fig. 9a). They display elevated Fe³⁺/Fe* ratios (0·29–0·37; Table 4). Calculated Mg# values fall within the range 61–67 (Fig. 9c). Their rare earth element (REE) patterns are relatively flat (1·79 < La_N/Yb_N < 2·83; 2·37 < Sm_N/Yb_N < 2·80).

Metatonalite pods within the Berrio ultramafic rocks

The metatonalites are silica-rich (SiO₂ 68·79-72·61 wt%), poor in Fe₂O₃* (1.61-2.45 wt%) and rich in Na₂O (8.76-9.71 wt%) (Fig. 9a and c). Their K₂O content is negligible (<0.1 wt%). The Fe^{3+}/Fe^* ratio varies from 0.27 to 0.40 (Table 4). They display a light REE (LREE)-enriched pattern (La_N/Yb_N up to 85.21). Three samples display negative Eu anomalies (Eu/Eu* = 0.17-0.72) and one sample shows a positive Eu anomaly (Eu/Eu* = 1.64) (Fig. 9e). The chlorite-schist at the contact with the ultramafic rocks displays a strong decrease in SiO₂ (29.23 wt%) and an important increase in Al₂O₃ (18.66 wt%), Fe₂O₃* (13.59 wt%) and MgO (25.95 wt%). The loss on ignition (LOI) strongly increases towards the reaction zone (from 0.77 wt% in the metatonalite to 11.13 wt% in the chlorite-schist) and may be attributed to the presence of H₂O structurally bound principally in chlorite. Metatonalite and chlorite-schist display similar REE patterns (Fig. 9e), suggesting that both lithologies evolved from the same protolith.

Blueschist

The blueschist displays an intermediate SiO₂ content (57.92 wt%) and a relatively high alkali content (6.41 wt%) (Fig. 9b). It is rich in Fe₂O₃* (9.09 wt%), with a relatively high Fe³⁺/Fe* ratio (0.49; Table 4). Assuming

that the protolith of the blueschist was a volcanic rock, its composition plots in the trachyandesite field in both the total alkalis–silica (TAS) (Le Maître, 2002) and Zr/Ti versus SiO₂ diagrams (Winchester & Floyd, 1977).

Reaction zone at the boundary between orthogneiss and ultramafic rock

A geochemical profile comprising 12 samples was traced across the reaction zone between the Arolla orthogneisses and the Berrio ultramafic rocks (site 2), by far the best outcrop of this reaction zone (Fig. 3a and b). The profile comprises three samples of orthogneisses, six samples of amphibole-gneisses and three samples of ultramafic rocks (Figs 3b and 10).

The Arolla orthogneisses (samples OL25, OL24 and OL23) show an acidic composition (SiO₂ = 61·75–71·19 wt%) and high alkali content (7·18–7·79 wt%). The Na₂O/K₂O ratio drastically increases from the orthogneiss to the amphibole-gneiss. The amphibole-gneisses display variable SiO₂ content (55·89–72·42 wt%). They are rich in Na₂O (5·62–10·74 wt%) but very low in K₂O (<1 wt%) (Fig. 9b and d). They show a large variation in Fe₂O₃* (0·57–6·13 wt%) content and high Fe³⁺/Fe* ratios (0·43–0·50).

Some trends in major and trace elements are evident along the profile (Fig. 10, Table S11). In the orthogneisses, MgO and Al₂O₃ contents increase towards the contact with the ultramafic rocks, whereas SiO₂ decreases (from A to C in Fig. 10b). CaO and Na₂O contents display a continuous increase (from A to B) followed by a rapid decrease close to the contact (C; Table S11). The LOI increases abruptly from 1.20 wt% in the orthogneiss (A) to 11.26 wt% in the chlorite-schist (C). Its concentration correlates with the modal abundance of chlorite (\sim 90%) in the schist and is thus attributed to the presence of structurally bound H₂O. The molar Fe^{3+} Fe* ratio decreases from 0.50 to 0.40 (A) followed by an increase up to 0.67 (B) and decrease down to 0.05 (C, chlorite-schist). Some of the large ion lithophile elements (LILE; e.g. K, Ba, Rb and Cs, Table S13) decrease in concentration towards the contact with the ultramafic rocks (A to B). This could be related to the absence of muscovite in the amphibole-gneiss. This trend is followed by a rapid and localized increase in LILE corresponding to a thin layer of muscovite (sample 49B, C; Fig. 10a) and then by an abrupt disappearance of these

Table 3: A summary of the deformation-growth relationships in the studied rocks

Lithology	Deformation	Texture	Zoning	Timing of growth
Hornblendite	Undeformed	Coronitic overgrowth on magmatic amphibole	Unzoned	Peak
Metatonalite	Undeformed	Needles and rosettes and coronas around sodic clinopyroxene	Unzoned	Prograde to peak
Blueschist	Foliation <i>S</i> ₁ + crenulation <i>S</i> ₂	Aligned along the S_1 foliation and passively reoriented along the crenulation planes S_2	Zoned (mrbk core, gl rim)	Prograde to peak
Amphibole-gneiss	Mylonitic foliation <i>S</i> 1	Parallel to the S_1 and randomly oriented cross-cutting the S_1	Zoned (mrbk at the core, wnc rim)	Peak to post

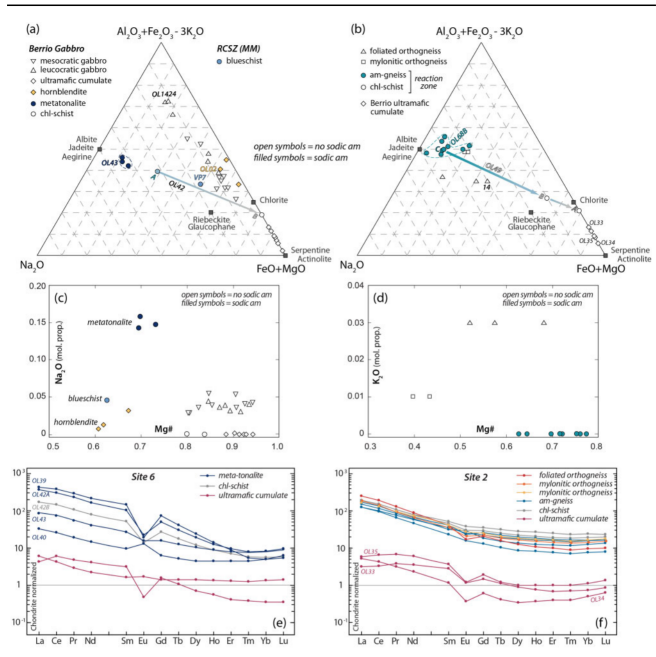


Fig. 9. Whole-rock composition of the studied samples. (a) Ternary diagram showing the bulk-rock compositions of the Berrio Gabbro rock types (sites 1 and 6) and blueschist (site 7). The analyses of leucocratic and mesocratic gabbros are from Manzotti et *al.* (2017). (b) Ternary diagram showing the bulk-rock compositions of orthogneiss, am-gneiss and ultramafic cumulate from sites 2, 3, 4, and 5. Strictly speaking, the ternary diagrams are projections from quartz, muscovite, clinozoisite/epidote and vapour. It should be noted, however, that some of these phases are not present in all lithologies (especially quartz and epidote in the chlorite-schist and in the ultramafic cumulate). (c) Mg# vs Na₂O (expressed as molecular proportions) diagram for the Berrio Gabbro lithologies and blueschist. Open symbols refer to rocks that do not contain sodic amphibole; filled symbols refer to rocks containing sodic amphibole (i.e. metatonalite, hornblendite, and blueschist). Metatonalite displays low Mg# and high Na₂O content. Hornblendite and blueschist show low Mg# and moderate Na₂O content. (d) Mg# vs K₂O (expressed in molecular proportions) diagram for the samples orthogneiss and amphibole-gneiss. Open symbols refer to rocks that do not contain sodic amphibole; filled symbols refer to rocks containing sodic amphibole-gneiss. Open symbols refer to rocks that do not contain sodic amphibole; filled symbols refer to rocks containing sodic amphibole-gneiss. Open symbols refer to rocks that do not contain sodic amphibole; filled symbols refer to rocks containing sodic amphibole. Amphibole-gneisses display very low K₂O content. (e) Trace element compositions of the samples from site 6. The similarity between the pattern of the metatonalite and chlorite-schist, indicating that both evolve from the same protolith, should be noted. (f) Trace element compositions of the samples from site 2. The similarity in the pattern of the orthog-neisses and chlorite-schist, indicating that they all evolve from the same protolith, sh

elements approaching the contact with the ultramafic rock (sample 49A, C). In the ultramafic rocks, the Fe³⁺/ Fe* ratio decreases from 0.36 to 0.20 towards the contact with the orthogneisses.

Along the profile, the orthogneiss, the amphibolegneiss and the chlorite-schist display similar LREEenriched patterns (Fig. 9f), indicating that the three lithological units were derived from the same granitic

			2	Aetamorphic rock	×			Metasomatic rock	atic rock
	Ultramafic cumulate	Mesocratic gabbro	Horn- blendite*	Meta- tonalite*	Leucocratic gabbro	Orthogneiss	Roisan blueschist*	Am-gneiss*	Chlorite- schist
Fe ₂ O ₃ * (wt%)	6.19–16.29	3·37–8·59	13.01–21.92	1.61–3.26	0.97-4.37	2.72-5.11	9·09	0.57-6.13	10.92-13.59
FeO* (wt%)	5.57-14.66	3.03-7.73	11.71–19.72	1.45–2.93	0.87–3.93	2.45-4.60	8·18	0.51-5.52	9.83–12.23
FeO* + MgO (wt%)	36.68-46.28	9.66-17.11	19·19–28·74	2.52–3.77	2·86–15·00	3.41–6.28	10.79	0.74-7.84	36.68–38.17
Mg#	88-94	66-82	61–67	69–75	85	52-68	63	71–77	80-84
Fe ^{Z+} /Fe ²⁺ + Mg	0.06-0.12	0.06-0.20	0.33-0.39	0.27-0.31	0.07-0.15	0.23-0.43	0.37	0.23-0.37	0.16-0.20
$Fe^{3+}/Fe^{3+} + Fe^{2+}$	0.15-0.40	0.18-0.66	0.29-0.37	0.27-0.40	0.18-0.60	0.27-0.61	0.49	0.35-0.67	0.03-0.05
Na/Na + Ca	0.00-0.01	0.11-0.22	0.02-0.13	0.90-0.94	0.11-0.18	0.67-0.96	0.39	0.50-0.94	0.00-0.03
*Bock types containing sorlig amphibole and/or sorlig nyroxene	d sodic amphibole	and/or sodic pyro	xana						

Table 4: Synthesis of the main chemical characters of the studied rocks

N

protolith. Considering the entire profile, the most prominent feature is the decrease of all the high field strength elements (HFSE) and LILE at the transition from the granitic orthogneiss to the ultramafic rocks.

THERMODYNAMIC MODELLING

Thermodynamic modelling was used to constrain the P-T-X (Fe₂O₃ and H₂O) conditions at which sodic amphibole ('gl') and/or sodic pyroxene ('cpx') developed in different rock types from the DBTS (i.e. hornblendite OL02, metatonalite OL43, blueschist VP7, and am-gneiss OL68B). For comparison, a leucocratic metagabbro (OL1424) and an orthogneiss (14), both devoid of sodic amphibole and sodic pyroxene, were modelled.

P-T modelling

Hornblendite OL02

In the hornblendite (OL02), blue amphibole only occurs as overgrowths on magmatic pargasite in the type-I domains (see the section 'Petrography and mineral chemistry'). To calculate phase diagrams considering only metamorphic minerals, the chemical contribution of magmatic pargasite (20%, estimated by point counting) was subtracted from the bulk chemical composition obtained by X-ray fluorescence (XRF). The corrected bulk chemistry is considered to be a better approximation of the effective bulk composition during metamorphic recrystallization and was used to calculate a P-T pseudosection (Fig. 11a) in the system MnNCFMASHTO. In this model, the transition from 'greenschist-facies' to 'blueschist-facies' assemblages occurs over a narrow P interval (from 7.5 to 8.5 kbar depending on T; Fig. 11a). In this limited P range (less than 1kbar), the Na₂O component is transferred from ab to gl. Correspondingly, the amphibole mode isopleths show that its modal abundance rapidly increases from 5 to 20 vol%. The modal abundance of sodic amphibole modelled in the field gl-ep-pa-chl-q-sph, which corresponds to the observed assemblage in thin section, is 15-20 vol%. The modelled composition of the sodic amphibole ('gl') plots in the glaucophane field, close to the boundary of Fe-glaucophane (Fig. 12a), and does not change significantly with P. The observed chemistry of sodic amphibole in the studied sample plots in the magnesio-riebeckite field: it has the same Fe^{2+} content as predicted by the numerical modelling and a slightly higher Fe³⁺ content than predicted. Paragonite is predicted to be stable above 4kbar at 350°C and 9kbar at 480°C, but has never been observed. However, its modal amount is small. For example, in the field gl-ep-pa-chl-sph-q-ab, paragonite modal amount increases from 1 to 5 vol% with increasing *P* (Fig. 11a).

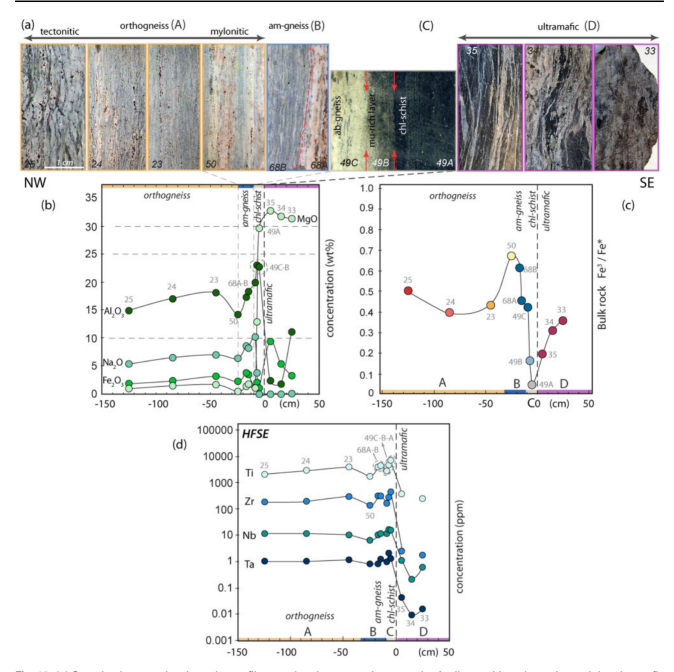


Fig. 10. (a) Sample photographs along the profile traced at the contact between the Arolla granitic orthogneiss and the ultramafic cumulates of the Berrio Gabbro (site 2, Fig. 3). (b–e) Chemical composition along the studied profile: (b, c) major elements; (d) trace elements (HFSE). For clarity, only the samples' number is reported in the figure. The colours at the bottom of the diagrams refer to the different lithologies along the section, as in Fig. 3b.

Metatonalite OL43

A *P*-*T* pseudosection (Fig. 11b) has been calculated in the MnNCKFMASHTO model system for the bulk composition obtained by XRF. Three amphiboles are present in the calculated phase diagram. A sodic amphibole is stable at rather high *P*. At low *P*, actinolite is stable at low *T*, and hornblende becomes stable at *T* higher than *c*. 450 °C. Owing to the continuous nature of the reactions, fields with two or three coexisting amphiboles (act + gl, act + hb, gl + hb, act + gl + hb) are present in the phase diagram. The transition from greenschist-facies assemblages (i.e. actinolite-bearing) to blueschist-facies assemblages (i.e. glaucophanebearing) occupies a large *P*–*T* band, up to 2 kbar wide at a given *T*. Sodic amphibole is stable down to 5 kbar at $350 \degree C$ and 7 kbar at $470 \degree C$. With increasing *P* (at a fixed T of *c*. 400 °C), the modal proportion of sodic amphibole significantly increases (from 2 to 6 vol%) once actinolite is consumed and epidote appears.

The mineral assemblage gl-act-cpx-ep-sph-q-ab, observed in thin section, is stable in the range 8–9 kbar and 430–480 °C. The chemical composition of both the

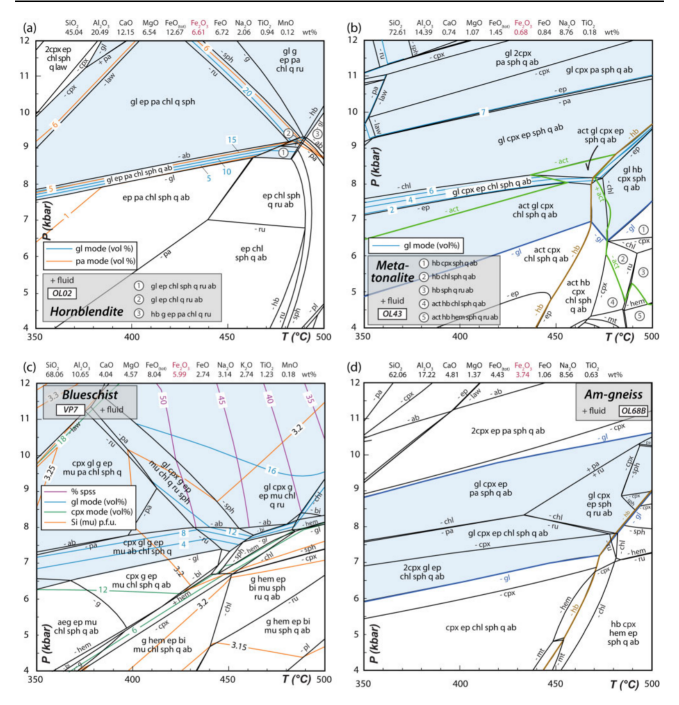


Fig. 11. *P*–*T* pseudosections for the studied samples: (a) OL02, hornblendite; (b) OL43, metatonalite; (c) blueschist VP7; (d) amgneiss OL68B. The blue field indicates the *P*–*T* space where sodic amphibole ('gl') is stable. Blue lines in (a) and (b) show the glaucophane modal amount (vol%). Muscovite compositional isopleths for Si (in p.f.u.) are reported in orange in (c). The amount of Fe₂O₃ used for the *P*–*T* calculation is highlighted in red. The grey arrows indicate two possible *P*–*T* path consistent with mineral assemblages. For clarity, some fields are not labelled; their mineral assemblages can be deduced from assemblages in adjacent fields.

sodic amphibole and the sodic pyroxene displays a decreasing $Fe^{3+}/(Fe^{3+} + AI^{3+})$ ratio with increasing *P* (Fig. 12a, c and d). The measured compositions of sodic amphibole and sodic pyroxene in this sample suggest a *P* of ~7 kbar and 7–9 kbar respectively (Figs 6 and 7).

Blueschist VP7

A *P*–*T* pseudosection (Fig. 11c) has been calculated in the MnNCKFMASHTO model system for the bulk composition obtained by XRF. Sodic amphibole ('gl') is predicted to be stable at *P* higher than \sim 6–8kbar in the *T*

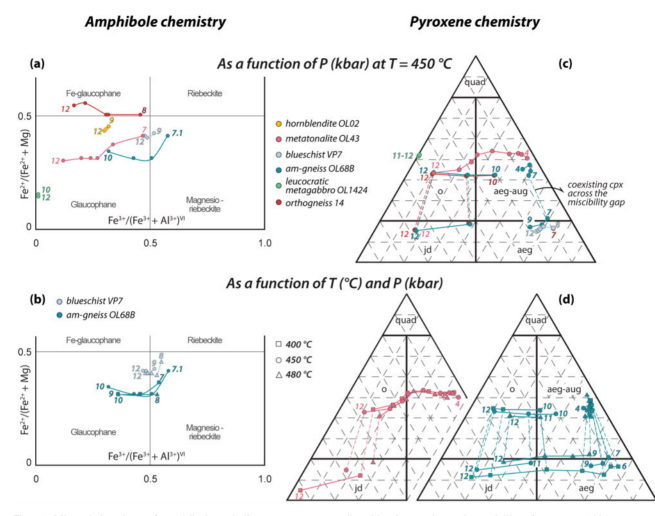


Fig. 12. Mineral chemistry of amphibole and clinopyroxene as predicted by thermodynamic modelling, from 4 to 12 kbar at 450 °C (a) and across the P-T space (b) (4–12 kbar at 400, 450 and 480 °C). Numbers are pressures in kbar. Dashed lines show coexisting minerals.

range considered (350–500 °C), always with a sodic pyroxene. The assemblage gl–cpx–g–ep–mu–chl–q–sph, which best approximates the observations, is stable in the triangular field at \sim 7.5–9 kbar and \sim 400–430 °C (Fig. 11c). Calculated isopleths suggest that 6–12 vol% pyroxene is present in this field.

The composition of the modelled sodic amphibole plots close to the boundary between the magnesioriebieckite and glaucophane (Fig. 12a and b), similar to the results of the electron microprobe (Fig. 6a). The measured compositional variations may be interpreted as recording a slight increase in *P* during amphibole growth (Fig. 6a). The composition of the modelled sodic pyroxene plots in the field of aegirine (Fig. 12c). However, sodic pyroxene has not been observed in the studied sample. The modelled garnet compositional isopleths for X_{spss} constrain the growth of garnet to about 400–450 °C (Fig. 11c).

Amphibole-gneiss OL68B

A *P*-*T* pseudosection (Fig. 11d) has been calculated in the NCFMASHTO model system for the bulk

composition obtained by XRF. The calculated phase diagram displays a large P stability field for sodic amphibole at intermediate P range and a hornblende stability field above 450 °C. Above 9-10 kbar, sodic amphibole disappears and is replaced by assemblages with two cpx, namely omphacite and jadeite (Fig. 12c and d). The sodic pyroxenes display a complex behaviour with increasing P. Considering an isothermal section at 450 °C, a sodic pyroxene with an aegirine-augite composition is stable at *P* lower than 6 kbar (Fig. 12c). At slightly higher P (6-8 kbar), two pyroxenes become stable, with a composition of aegirine-augite and aegirine. With increasing P (8-10 kbar), only one sodic pyroxene is stable, with a composition first of an aegirine then of an aegirine-augite. Above 10kbar, two pyroxenes are stable again, with an omphacite and jadeite composition. Paragonite is stable at P > 7.5-10 kbar. Chlorite disappears at P < 7.5-8 kbar at T higher than 450°C.

The observed mineral assemblage gl-cpx-ep-sphq-ab is partially reproduced in the calculated phase diagram with one additional phase (either paragonite or chlorite or rutile). In each case, their modelled modal abundance is less than ~1 vol%. The composition of the sodic pyroxene measured in the studied sample is similar to the one predicted by the numerical model at 10–11 kbar (Fig. 7). The composition of the calculated sodic amphibole displays decreasing $Fe^{3+}/(Fe^{3+} + Al^{3+})$ with increasing *P* (Fig. 12a and b). At 7–9 kbar, its composition is similar to the one measured in the studied sample (Fig. 6a).

Leucocratic metagabbro OL1424

Amongst the samples of the leucocratic metagabbro, OL1424 was chosen because it lacks Cr-bearing phases, and therefore Cr could be neglected in the numerical modelling. A P-T pseudosection (Fig. 13a) has been calculated in the NCKFMASHO model system for the bulk composition obtained by XRF. In this pseudosection, sodic amphibole (with a glaucophane composition; Fig. 12a) is predicted to be stable above 9 kbar at 370 °C and \sim 11 kbar at 500 °C, paragonite from 7 kbar at 350 °C and 9kbar at 500 °C, whereas sodic pyroxene (with an omphacite composition; Fig. 12c) appears at 8.5 kbar and 350 °C and 10.5 kbar and 500 °C. Lawsonite is stable above 7.2 kbar at 350 °C and 12 kbar at 425 °C. This collectively indicates that the glaucophane-bearing, blueschist-facies parageneses are stable at high P(>9 kbar), whereas the greenschist-facies parageneses (i.e. epmu-chl-q-ab and act-ep-mu-chl-q-ab) occupy a large Trange below \sim 9 kbar.

The observed mineral assemblage ep–mu–chl–ab–q is predicted to be stable in the range 350–400 °C and 4.5-9 kbar. The composition of syn-S₁ muscovite (Si = 3.22-3.26 p.f.u.) suggests equilibration in a *P*–*T* band between 360–390 °C at 4 kbar and 440–470 °C at ~10 kbar in the neighbouring, actinolite-bearing field. Indeed, if actinolite was also present, the assemblage would be stable in a larger field ranging from 350–400 °C at 4 kbar to 430–500 °C at 9.5 kbar (dashed lines in Fig. 13a). Calculated actinolite mode is lower than 1.2 vol% in this field, potentially explaining why this mineral was not found. Si-rich muscovite (Si > 3.30 p.f.u.), found in possible pseudomorphs after lawsonite, is predicted to be stable at low *P*–*T* conditions (*P*<8.5 kbar, *T*<370 °C).

Orthogneiss 14

A *P*–*T* pseudosection (Fig. 13b) has been calculated in the NCKFMASHTO model system for the bulk composition of an orthogneiss reported by Diehl *et al.* (1952). The rock consists of muscovite, quartz, albite, epidote, chlorite, and titanite, but systematically lacks sodic amphibole. In the pseudosection, this assemblage is predicted in the LT–LP corner, at *T* < 400 °C and *P* < 6 kbar. Sodic amphibole is predicted in a wide range of *P*–*T* conditions (at *P* higher than ~6 kbar). Its composition plots in the Fe-glaucophane field close to the limit with the glaucophane field (Fig. 12a) and displays an Fe³⁺ content decreasing with increasing *P*. The mineral assemblage gl–ep–mu–chl–sph–q–ab is modelled in the *T* interval 350–460 °C between ~6 and 8.5 kbar

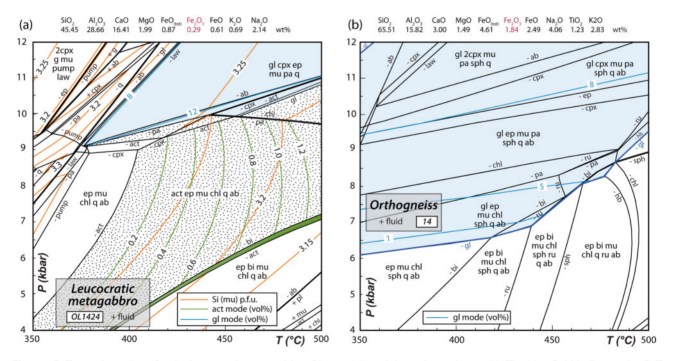


Fig. 13. *P*–*T* pseudosections for the leucocratic metagabbro OL1424 (a), and the orthogneiss 14 (b). The blue field indicates the *P*–*T* space where sodic amphibole ('gl') is stable. Blue lines show the glaucophane modal amount (vol%). The dotted field refers to the *P*–*T* space where actinolite is present. Diagram (a) is contoured (in green) for the actinolite modal amount (vol%). Muscovite compositional isopleths for Si (in p.f.u.) are reported in orange in (a). The amount of Fe_2O_3 used for the *P*–*T* calculation is indicated in red. The grey arrows indicate two possible *P*–*T* path consistent with mineral assemblages. For clarity, some fields are not labelled; their mineral assemblages can be deduced from assemblages in adjacent fields.

(Fig. 13b). In this field, sodic amphibole is modelled to be present from ~1 to ~5 vol% with increasing *P*. It is possible that the sample contained sodic amphibole and it has been replaced by chlorite–albite during decompression. Another explanation would be to accept as maximum *P*–*T* conditions at ~6.5 kbar and ~420 °C, which would explain the lack of sodic amphibole in the orthogneisses (Fig. 13b).

P-X(Fe₂O₃) modelling

 $P-X(Fe_2O_3)$ pseudosections have been calculated for the different rock types at 450 °C to explore the effect of the oxidation state of the rock (Fe³⁺–Fe²⁺ ratio) on the stability of sodic amphibole and pyroxene. A $X(Fe_2O_3)$ range of 0.05–0.9 was considered, with $X(Fe_2O_3) =$ Fe_2O_3/FeO^* [i.e. $X(Fe_2O_3) = 0$ means that all Fe is bivalent; $X(Fe_2O_3) = 0.5$ means that FeO and Fe₂O₃ are present in equal amounts]. The $P-X(Fe_2O_3)$ pseudosection (Fig. 14a) calculated at 450 °C for the hornblendite (OL02) shows that sodic amphibole is stable at relatively low Fe₂O₃ content above 7.5 kbar, whereas sodic pyroxene appears only at either very low (<0.14) or very high $X(Fe_2O_3)$ contents (0.60). Hematite and rutile are present only at high Fe₂O₃, whereas titanite is stable at low Fe₂O₃ content.

In the metatonalite (OL43), the stability field of sodic amphibole strongly varies with $X(Fe_2O_3)$ reaching the lowest P (~5.5 kbar) at a relatively low $X(Fe_2O_3)$ (~0.12; Fig. 14b). Below 5.5 kbar, the highest $X(Fe_2O_3)$ results in the appearance of hematite.

In the blueschist (VP7; Fig. 14c), hematite appears at relatively low P with increasing $X(Fe_2O_3)$ (>0.28). Sodic amphibole is stable at P > 7.5 kbar, except for very high $X(Fe_2O_3)$. From low to high $X(Fe_2O_3)$, sodic clinopyroxene is stable above \sim 9.5 kbar and \sim 6 kbar, and the composition varies from omphacite to aegirine, respectively. Above \sim 9 kbar, the two compositions coexist along a solvus in a narrow band at X(Fe₂O₃) \sim 0.33 wt%. The mineral assemblage gl-g-ep-mu-q-sphab observed in the studied sample is stable between 7.5 and 9 kbar for a $X(Fe_2O_3)$ of 0.25–0.45 (i.e. lower than the Fe^{3+}/Fe^{2+} ratio measured and used for the P-T calculation).

For the amphibole-gneiss (OL68B; Fig. 14d), the lowest *P* limit of sodic amphibole decreases from 8.5 to 7 kbar with increasing Fe₂O₃ content. Sodic amphibole disappears at $P \ge 10$ kbar for $X(Fe_2O_3) \ge 0.3$. At relatively low *P*, hematite and a sodic pyroxene (aegirine) are present at high $X(Fe_2O_3) (>0.71)$.

In the pseudosection calculated for the leucocratic metagabbro (OL1424) at 450 °C (Fig. 15a), for the measured Fe₂O₃ value, the assemblage act–ep–mu–chl–q–ab is stable between 6·5 and 9·8kbar. At these *P* conditions, an increase in the $X(Fe_2O_3)$ produces the disappearance of actinolite and an increase in the pistacite content of epidote. At *P* higher than 10kbar, sodic amphibole and then sodic pyroxene (omphacite) appear. The calculated diagram indicates that the lower *P* stability limit of sodic amphibole is not dependent on the Fe_2O_3 content of the rock.

The *P*-*X*(Fe₂O₃) pseudosection (Fig. 15b) calculated at 450 °C for the orthogneiss (14) shows that the lowest *P* limit of sodic amphibole first decreases from 9.2 to ~7 kbar with increasing *X*(Fe₂O₃) (up to ~0.61). For *X*(Fe₂O₃) higher than 0.61, the lowest *P* limit of sodic amphibole increases again up to ~8.5 kbar. Hematite is stable below 7 kbar at relatively high *X*(Fe₂O₃) (>0.41). Above 10 kbar and high *X*(Fe₂O₃), two pyroxenes (namely, aegirine and aegirine–augite) are stable.

In summary, although the stability field of sodic amphibole does depend on the oxidation state of the rock, the relation is not straightforward. In some lithologies, sodic amphibole is stabilized (at a given *T*) towards lower *P* at high Fe³⁺/Fe* ratios (e.g. Fig. 14d), whereas in others the stability domain is maximized at low Fe³⁺/Fe* ratios (e.g. Fig. 14b). In all lithologies, the chemistry of sodic amphibole, as calculated by the numerical model, displays an increasing Fe³⁺/(Fe³⁺ + Al³⁺) with increasing Fe³⁺/Fe* ratio in the bulk composition (Fig. 15c). This variation is much less pronounced in the leucocratic metagabbro; that is, the rock with the lowest Fe₂O₃* and Fe³⁺ contents, and the highest Mg#.

P-H₂O modelling

The previous calculations are based on the assumption that H₂O was in excess during the metamorphic evolution. Here, we evaluate the possibility that metamorphism occurred under H₂O-undersaturated conditions. The aim of this modelling is to test if the absence of sodic amphibole in some rock types (e.g. in the orthogneiss and leucocratic metagabbro) can be linked to a limited H₂O content in the system. The red lines in the calculated P-H₂O pseudosections (Fig. 14d and Supplementary Data Figs S4 and S5) represent the H₂O-saturation surface and divide the diagrams in a H₂O-saturated (right) and a H₂Oundersaturated (left) part. All diagrams show that H₂Oundersaturation leads to important rapid variations of the stable mineral assemblages, including crystallization of minerals commonly not expected for a given rock type. In the bulk compositions investigated, at a given T_{i} the minimum P of the stability field of sodic amphibole is generally rapidly displaced to higher P with increasing H₂O-undersaturation.

As an example, we detail here one sample (all samples are described in the Supplementary Data). A relatively large amount of H₂O (\sim 4–4.5 wt%, depending on *P*; Fig. 15d) is required to reach H₂O-saturated conditions in the hornblendite (OL02). With increasing H₂O-undersaturation, the minimum *P* of sodic amphibole stability first slightly decreases by \sim 0.5 kbar down to 8 kbar, then (for <3.4 wt% H₂O) rapidly increases up to >12 kbar (for <1.3 wt% H₂O). When H₂O-undersaturation is reached, the first changes of the mineral assemblages involve the disappearance of titanite, and the stabilization of rutile, hornblende and garnet over the entire *P* interval explored. Chloritoid appears at

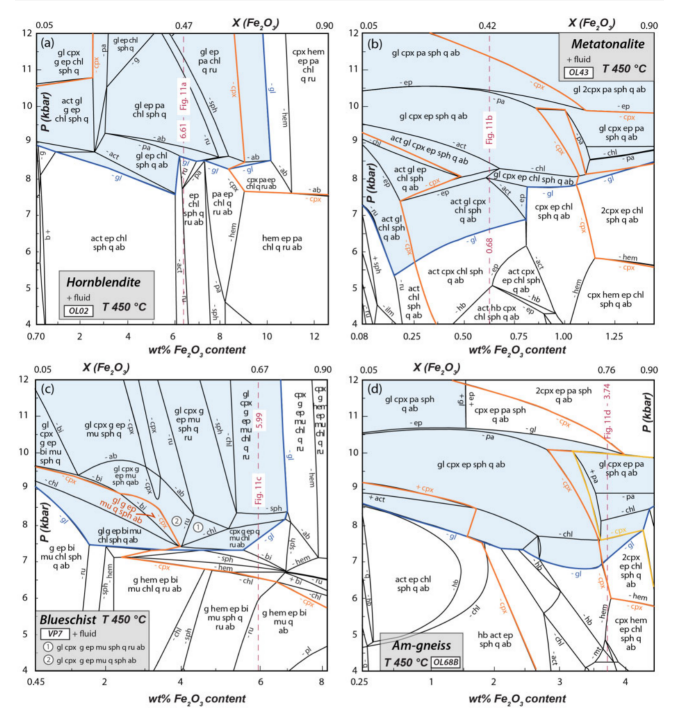


Fig. 14. $P-X(Fe_2O_3)$ pseudosections for the studied samples: (a) OL02, hornblendite; (b) OL43, metatonalite; (c) VP7, blueschist; (d) amphibole-gneiss (OL688). The blue field indicates the P-T space where sodic amphibole is stable. The red dashed line indicates the amount of Fe₂O₃ measured in the samples and used for the calculation of the P-T diagrams reported in Fig. 11. For clarity, some fields are not labelled; their mineral assemblages can be deduced from assemblages in adjacent fields.

P > 8 kbar, and staurolite for P > 5.5 kbar at H₂O ~ 1.6 – 2.6 wt%; kyanite becomes stable for H₂O contents lower than ~ 3 wt% at 12 kbar and ~ 1.8 wt% at 4 kbar. These minerals have never been observed in the studied sample. Consequently, the H₂O-saturated pseudosection yields the best match between the model and petrological observations. As a whole, the *P*-H₂O modelling (Fig. 14d and Figs S4 and S5) indicates that the assumption of excess water was probably valid for the studied rocks.

Reliability of the thermodynamic modelling

Evaluating the limits of the thermodynamic modelling is a key issue for the interpretation of the rocks. Thermodynamic data on Fe^{3+} -rich blue amphibole and

sodic pyroxene are currently under development (Green *et al.*, 2016). Because few experimental data are available and because of the uncertainty of Fe^{2+}/Fe^{3+} recalculation from microprobe analyses (as discussed above), these models may have a large degree of uncertainty. For this reason, we have carefully evaluated the match between calculated and analysed compositions of solid solution minerals and between observed and calculated parageneses in each studied sample (see above). Moreover, equilibrium conditions have not been reached in the described rocks as indicated by incompletely reacted relics (calcic amphiboles) and strong chemical zoning in blue amphibole and sodic pyroxene. This is expected during rather low-*T* metamorphism of magmatic rocks.

By essence, this work cannot evaluate the uncertainties associated with the end-member thermodynamic parameters and/or to the solid-solution models used, which are of the latest generation (Green *et al.*, 2016). For this, we should test a specific phase diagram assuming that P-T conditions are known, and compare the predictions of the phase diagram at such known P-Tconditions in terms of mineral assemblage, modal proportion of the phase in this mineral assemblage, and phase chemistry (see, e.g. Forshaw *et al.*, 2019).

However, given the large number of phase diagrams we have calculated, some general inferences may be made. First, H₂O saturation was most probably achieved in all studied samples, because of the lack of mineral phases (e.g. staurolite, kyanite, garnet, olivine, etc.) diagnostic of H₂O-undersaturated conditions, as predicted from the phase diagrams. Second, the influence of the fO_2 is fundamental but its exact relevance is difficult to assess, cumulating uncertainties in the determination of this ratio in the rock (our strategy gives the maximum amount of Fe₂O₃), and in the minerals from microprobe analyses (especially in amphibole; see above). By contrast, it is worth stressing that the composition of the pyroxene and amphibole, as calculated by the model, is in most cases similar to the one measured in our samples. When this composition is evolving with P-T conditions, the fit between calculated and observed composition is systematically in favour of the lower P range of stability for these minerals, giving some confidence in this approach.

DISCUSSION

Bulk-rock chemical control on the growth of sodic amphibole

Given the variety of rock types (from ultramafic to highly differentiated, felsic rocks) observed in the studied area, and the scarcity of rocks containing sodic amphibole and/or sodic pyroxene, the control of the bulk-rock chemistry on their occurrence should be carefully evaluated. Two groups of sodic amphibole-bearing rocks have been found. In a first group, the bulk-rock chemistry has not been significantly modified with respect to the protolith. In a second group, sodic amphibole develops in reaction zones at the contact between two contrasting lithologies; that is, in metasomatically altered rocks.

Sodic amphibole in (nearly) isochemical rocks

Sodic amphiboles (and sodic pyroxenes) may develop in a wide range of bulk-rock compositions, but are most commonly recorded in metabasites. In the studied area, metabasites constitute the bulk of the Berrio Gabbro. However, sodic amphibole is present only in the hornblendites, which differ from the other gabbros by having the lowest Mg# (61–67) (Fig. 9c). Therefore, the appearance of sodic amphiboles in this lithology may be primarily controlled by the more Fe-rich composition with respect the leucocratic and mesocratic gabbros (Mg# > 80) where no sodic amphibole has been observed (Table 4; Fig. 9c).

The metatonalites have a very different composition from the hornblendites, being much richer in Na_2O (Fig. 9c), and having a lower Fe* content. The appearance of sodic amphibole and sodic pyroxene may be therefore controlled by the high Na_2O and low Mg#.

The blueschist displays very high Fe^{*} content (Fe₂O₃^{*} = 9.09 wt%), a high molar bulk Fe³⁺/Fe^{*} ratio (0.49) and a low Mg#, similar to the hornblendite. Therefore, the occurrence of sodic amphibole in this rock seems to be strongly controlled by its high Fe^{*} content and Fe³⁺ content.

Sodic amphibole in reaction zones between granitic and ultramafic rocks

The granitic orthogneisses in the studied area are devoid of sodic amphibole and/or sodic pyroxene. Amphibole-gneiss is found at the contact between the granitic orthogneiss and serpentinized ultramafic cumulate. The major element variation (e.g. Al, Mg, Na, K) between the granitic orthogneiss and the amphibolegneiss demonstrates that the latter result from metasomatic processes at the contact between the two contrasting lithologies. With respect to the granitic orthogneiss, the amphibole-gneiss is enriched in CaO and Na₂O, and depleted in K₂O (Fig. 9d). Trace elements display a decrease in LILE, whereas HFSE remains nearly constant. The depletion of LILE is explained by a decrease in muscovite abundance along the profile from the orthogneiss towards the amphibole-gneiss. The REE patterns of the orthogneiss and the amphibolegneiss are similar. It should be noted that the protolith of the chlorite-schists is difficult to identify in the field, as nearby ultramafic rocks can be erroneously considered as their protoliths. However, the presence of magmatic zircon, apatite and minor allanite suggests a granitic protolith for the chlorite schists, which is supported by the similar REE patterns of the unaltered orthogneiss and the chlorite-schist. During the metasomatic alteration, granitoids may provide silica to the ultramafic cumulates, contributing to the formation of

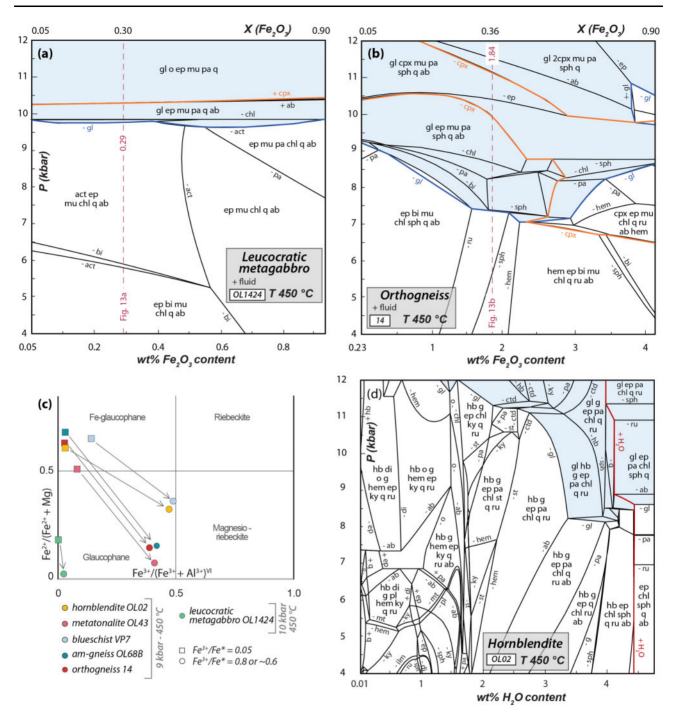


Fig. 15. $P-X(Fe_2O_3)$ pseudosection for (a) the leucocratic metagabbro (OL1424) and (b) Arolla orthogneiss (14). The blue field indicates the P-T space where sodic amphibole is stable. The red dashed line indicates the amount of Fe_2O_3 measured in the samples and used for the calculation of the P-T diagrams reported in Fig. 13. For clarity, some fields are not labelled; their mineral assemblages can be deduced from assemblages in adjacent fields. (c) Chemical composition of sodic amphibole as predicted by numerical modelling with the lowest and highest Fe_2O_3 content in the system. (d) $P-XH_2O$ pseudosections for the hornblendite (OL2). The red lines represent the H₂O-saturation surface and divide the diagram in an H₂O-saturated (right) and an H₂O-undersaturated (left) part.

talc veins in these rocks close to the contact with the granitoids (e.g. Jöns *et al.*, 2009). The preserved cumulate texture in the ultramafic rocks as well as the presence of porphyroclasts replaced by chlorite aggregates in the chlorite-schist suggest that intense deformation was very limited and strongly localized along the lithological contact.

Metamorphic history of the Dent Blanche Tectonic System

In the DBTS, the Alpine evolution is heterogeneous, with large volumes preserving the pre-Alpine structures and parageneses. For example, paragneisses and amphibolites from the country-rocks of the Permian Mont Morion granitoid record pre-intrusive partial melting at high *T* (Manzotti *et al.*, 2018), but are devoid of any sign of Alpine metamorphism. Another example is provided by the Valpelline Series, in which Permian kilometre-scale high-grade paragneisses are preserved (Nicot, 1977; Gardien *et al.*, 1994; Manzotti & Zucali, 2013; Kunz *et al.*, 2018), except for rare kyanite at biotite-cordierite boundaries and chloritoid-muscovite pseudomorphs after cordierite (Kienast & Nicot, 1971).

In pervasively deformed rocks, the dominant synkinematic parageneses are ascribed to the greenschist facies. Above we discussed in detail the few lithologies in which sodic amphibole and sodic pyroxene have been found. Two different explanations can be envisaged for the sparse occurrence of these two index minerals.

- Blueschist-facies metamorphism in the DBTS (i.e. well-developed blueschist-facies parageneses in all rocks that have been subjected to ductile deformation; Fig. 1) may have been present during an early stage of the Alpine compressional history, but has been extensively overprinted at greenschist facies during exhumation. As a consequence, the sodic amphiboles and sodic pyroxenes would represent relics of this early blueschist, only exceptionally preserved in some rocks.
- 2. According to the second hypothesis, peak *P*-*T* conditions in the studied area never reached blueschistfacies conditions, but rather the transition between pumpellyite-actinolite or greenschist facies and blueschist facies (Fig. 1). In this *P*-*T* domain, sodic amphibole and sodic pyroxene develop only in favourable chemical compositions; that is, in those especially rich in Na and Fe³⁺ and/or displaying high Fe²⁺/(Fe²⁺ + Mg) ratios (low Mg#). At higher *P*, the range of bulk compositions susceptible to develop minerals diagnostic for the blueschist facies would have expanded, and their occurrence should have been much more common than observed.

Numerical modelling of the observed parageneses is consistent with a model in which all rocks containing sodic amphibole and sodic pyroxene developed at *c*. 400–450 °C (Figs 11 and 13), in the range of 8 ± 1 kbar. In this *P*–*T* range, albite is stable in most lithologies, but Na would have been transferred from albite into sodic amphibole and/or sodic pyroxene in specific Na, Fe* and Fe³⁺-rich bulk compositions. These characters may be 'primary' (as in the hornblendites), or be acquired following metasomatic transformations (as in the amphibole-gneiss). This would explain why sodic amphibole and sodic pyroxene are scarce in the entire DBTS.

An extensive greenschist-facies overprint cannot explain the actual distribution of the sodic amphiboles and pyroxenes for the following reasons.

 In undeformed rocks, such as hornblendite in the gabbros, sodic amphibole mainly developed as a corona around the brown (magmatic) amphibole, a local domain with the highest Fe*. Away from the magmatic amphibole, in the plagioclase pseudomorphs, sodic amphibole is very rare, and actinolite is dominant. Therefore, the distribution of sodic amphibole in the undeformed rocks such as the hornblendites is not linked to a heterogeneous distribution of the greenschist-facies overprint, but to the local microchemical systems inherited from the magmatic protoliths.

- 2. In the amphibole-gneiss, field observations show that sodic amphibole is stable both during D_1 , where aligned crystals of amphibole define the stretching lineation, and during or after D_2 , where rosettes or sheaves grew across D_2 microfolds. In the orthogneiss protolith, sodic amphibole has not been observed. The continuity between the orthogneiss and the amphibole-gneiss can be ascertained thanks to the excellent and continuous outcrop conditions (Fig. 3) and both rocks must have shared the same P-T history. The lack of sodic amphibole in the orthogneiss is therefore controlled by differences in bulk-rock chemistry, already present during the early metamorphic evolution of the DBTS.
- 3. The blueschists are interbedded with felsic gneisses. Both lithologies are folded together and share the deformational history (D_1 , D_2), yet sodic amphibole is never present in the felsic gneisses, whereas it is common in the blueschists and apparently stable during both deformation events.

We thus conclude that pervasive greenschist-facies overprint cannot be the main controlling factor for the lack of sodic amphibole and pyroxene in most lithologies from the DBTS. Rather, bulk-rock chemical composition controls the growth of sodic amphibole in the most oxidized and/or the most Na- and Fe-rich rocks, pointing towards rather low $P(\sim 8 \pm 1 \text{ kbar})$ of equilibration of these assemblages.

The greenschist- to blueschist-facies transition monitored by blue amphiboles

This study demonstrates, in the specific case of the Dent Blanche Tectonic System (Western Alps), the key role of bulk-rock chemistry [high Na⁺, Fe²⁺/(Fe²⁺ + Mg) and Fe³⁺/Fe^{*}] in stabilizing sodic amphibole and/or sodic pyroxene at relatively low *P*; that is, at the transition between the greenschist and the blueschist facies. Our calculations indicate that the chemical composition of sodic amphibole displays decreasing Fe³⁺/(Fe³⁺ + Al³⁺) ratios with increasing *P* (for a given bulk-rock composition) and an increasing Fe³⁺/(Fe³⁺ + Al³⁺) with increasing Fe³⁺/Fe^{*} ratio in the bulk-rock.

To test these results further, the amphiboles from the Dent Blanche Tectonic System are compared with published analyses from other areas. Figure 16 represents a literature compilation of the chemical composition of sodic amphiboles from worldwide terrains at different metamorphic facies (i.e. *P*–*T* conditions). The following groups have been made with reference to the

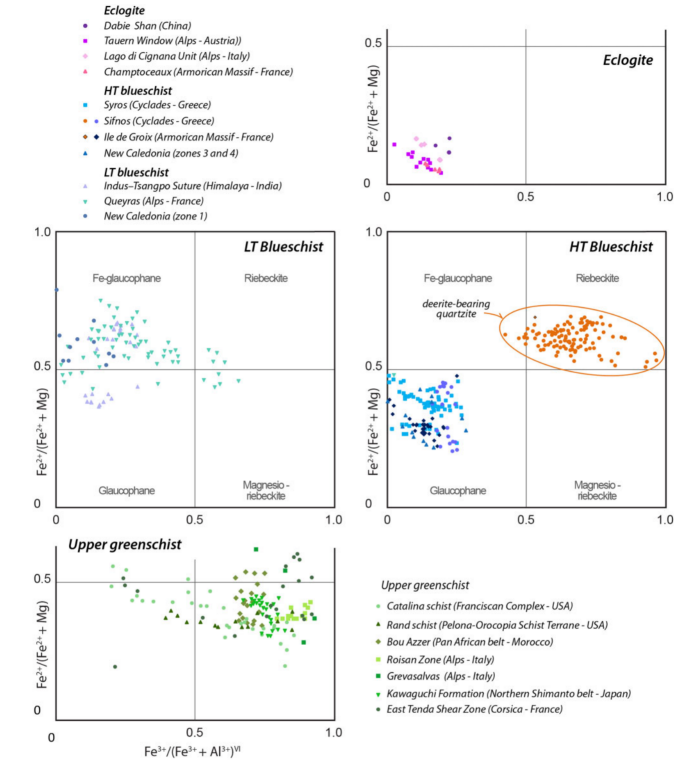


Fig. 16. Range of blueschist sodic amphiboles plotted in the Miyashiro & Banno (1958) diagram. The chemical composition of sodic amphibole has been recalculated following the procedure used in this study (see the section 'Mineral chemistry'). Upper greenschist: Catalina schist and Rand schist (Jacobson & Sorensen, 1986); Bou Azzer, Pan-African belt (Hefferan *et al.*, 2002; Bousquet *et al.*, 2008); Roisan Zone, Alps (Manzotti *et al.*, 2012); Grevasalvas, Alps (Philipp 1982); Kawaguchi Formation, Northern Shimanto belt (Aoki *et al.*, 2008); East Tenda Shear Zone, Corsica (Malasoma *et al.*, 2006; Maggi *et al.*, 2012; Rossetti *et al.*, 2015). Lower *T* blueschist: Indus–Tsangpo Suture, Himalaya (Groppo *et al.*, 2016); Queyras (Pognante & Kienast, 1989; Ballèvre & Lagabrielle, 1994); New Caledonia zone 1 (Clarke *et al.*, 1997; Fitzherbert *et al.*, 2005; Vitale Brovarone & Agard, 2013). Upper *T* blueschist: Syros (Schumacher *et al.*, 2008; Miller *et al.*, 2009; Philippon *et al.*, 2013; Skelton *et al.*, 2019); Sifnos (Evans, 1986; Groppo *et al.*, 2009); Ile de Groix, Armorican Massif (Dudek & Kienast, 1989; Ballèvre *et al.*, 2003); New Caledonia zones 3 and 4 (Clarke *et al.*, 1997; Fitzherbert *et al.*, 2013); Taetz *et al.*, 2016). Eclogite: Dabie Shan (Zhang & Liou, 1994); Tauern Window, Alps (Holland, 1979); Lago di Cignana Unit, Alps (Groppo *et al.*, 2009*b*); Champtoceaux, Armorican Massif (M. Ballèvre, unpublished data).

parageneses observed in associated metabasites and metapelites.

Upper greenschist rocks are defined by the occurrence of high-Si muscovite and chlorite and the lack of biotite in metapelites. Most mafic rocks display assemblages consisting of albite, epidote, chlorite, actinolite and titanite. However, sodic amphibole and sodic pyroxene may be present in some lithologies, such as some metabasites and felsic orthogneisses (East Tenda Shear Zone: Malasoma *et al.*, 2006; Maggi *et al.*, 2012; Rossetti *et al.*, 2015), a few Mn- and Fe-rich metasediments (Roisan Zone: Ballèvre & Kienast, 1987; Manzotti *et al.*, 2012; Grevasalvas: Philipp, 1982), but their occurrence is always scarce. Reported sodic amphibole compositions from such terranes mainly plot in the Mgriebeckite field, similar to the ones reported in this study.

The blueschist-facies occurrences have been subdivided into two groups. The low-T (LT) blueschists are distinguished by the common occurrence of lawsonite (rather than clinozoisite) in metabasites and the lack of garnet in associated metapelites (typically containing chlorite-chloritoid ± carpholite). Most sodic amphiboles in such terranes, such as the blueschists along the Indus-Tsangpo suture zone (Groppo et al., 2016), those from the External domain of the Piemonte-Liguria Ocean (i.e. Queyras in Fig. 16) in the Western Alps (metagabbros: Pognante & Kienast, 1987; marbles: Ballèvre & Lagabrielle, 1994) or those from the New Caledonia (zone 1: Clarke et al., 1997; Fitzherbert et al., 2003, 2005; Vitale Brovarone & Agard, 2013) plot in the Fe-glaucophane field, whatever the composition of the bulk-rock in which they occur.

The high-T (HT) blueschists represent the bulk of glaucophane-bearing rocks, and are defined by the occurrence of garnet-chloritoid \pm chlorite assemblages in the metapelites. Sodic amphibole is present in a large variety of bulk-rock compositions. These include the classic garnet-bearing blueschists, and glaucophaneepidote eclogites, derived from various kinds of metabasalts and metagabbros. Notable examples are found in the type locality for glaucophane Syros (Miller et al., 2009; Philippon et al., 2013; Skelton et al., 2019), the lle de Groix (Ballèvre et al., 2003), and New Caledonia (zones 3 and 4, Clarke et al., 1997; Fitzherbert et al., 2003, 2005; Vitale Brovarone & Agard, 2013; Taetz et al., 2016). In these localities, sodic amphibole is also present in some low-Al metapelites (e.g. Bosse et al., 2002), a few marbles (e.g. Syros: Schumacher et al., 2008), and the deerite-bearing rocks (Sifnos: Evans, 1986; Groppo et al., 2009a), representing the most-Ferich rocks in this spectrum. Accordingly, the chemistry of sodic amphibole changes with bulk-rock composition but they all plot in the glaucophane field (except for the deerite-bearing parageneses).

Eclogite-facies occurrences are here defined when metabasites contain garnet-omphacite-kyanite and metapelites are equilibrated in the garnet-chloritoidkyanite stability field (Tauern Window: Holland, 1979; Sesia Zone: Vuichard & Ballèvre, 1988) or at *T* above the stability field of chloritoid; that is, in the garnetkyanite stability field (Champtoceaux: Ballèvre *et al.*, 1989). Sodic-amphibole parageneses in such rocks become very rare, and sodic amphibole displays the lowest XFe^{2+} and XFe^{3+} of the entire spectrum (glaucophane is therefore close to its end-member composition). In ultrahigh-*P* (UHP) terranes, sodic amphibole close to end-member glaucophane has been reported from the Dabie Shan (Zhang & Liou, 1994) and the Lago di Cignana (Groppo *et al.*, 2009*b*). In other UHP terranes, such as the Dora Maira Massif, sodic amphibole slightly departs from the glaucophane composition (e.g. Chopin, 1986; Kienast *et al.*, 1991; Schertl *et al.*, 1991).

This discussion shows that both the Fe³⁺/(Fe³⁺ + Al³⁺) ratio and the Fe²⁺/(Fe²⁺ + Mg) of sodic amphibole decrease with increasing *P* and *T*. This general tendency is obviously counteracted by the effect of the bulk-rock composition, the best example being the deerite-bearing quartzites from lle de Groix and Sifnos, which display much higher Fe³⁺/(Fe³⁺ + Al³⁺) ratios than the other rock types from the same locality, although all were metamorphosed at high-*T* blueschist-facies conditions. The chemical composition of the sodic amphibole from the DBTS plots well in the diagram of the upper greenschist facies, again supporting the hypothesis that the bulk chemical composition controlled the growth of this mineral.

CONCLUSIONS

- In the DBTS sodic amphiboles and sodic pyroxenes have developed as a consequence of favourable whole-rock chemical composition, dominated by a combination of two or three parameters; namely, elevated Na₂O content, high Fe²⁺/Fe²⁺Mg and/or high Fe³⁺ content.
- 2. These parameters are achieved in specific rocks either because of their initial, pre-metamorphic chemistry (e.g. hornblendite, metatonalite), or because of some substantial metasomatic changes (amphibolegneiss, a reaction zone at the boundary between felsic orthogneiss and ultramafic cumulates). Sodic amphiboles and pyroxenes develop only in the generally unreactive rocks of granitic bulk compositions where these are modified by metasomatic fluid-rock interactions in an open system.
- 3. Our thermodynamic modelling shows that the rocks may have equilibrated at 8 ± 1 kbar, 400–450 °C, with excess H₂O. This study represents the first attempt to reconstruct the Alpine evolution of the DBTS by modelling the rare sodic amphibole- and sodic pyroxene-bearing rocks. Petrological models demonstrate that sodic amphibole and pyroxene formed in Fe³⁺-rich systems at relatively low P (~8±1 kbar, ~450 °C); that is, at the transition from the greens-chist- to the blueschist-facies metamorphism (path B

in Fig. 1). Therefore, these minerals record the incipient stages of HP–LT metamorphism fully developed in more internal units of the Sesia–Dent Blanche nappe stack.

4. Our conclusions are in line with the earlier studies of Miyashiro & Banno (1958) and Evans (1990). Indeed, thermodynamic modelling of specific bulk-rock compositions, using solid-solution models taking into account Fe³⁺, indicates that the *P*-*T* stability field of sodic amphiboles is largely dependent on their chemical composition. This link has been substantiated by a compilation of the chemistry of sodic amphiboles in a large range of *P*-*T* conditions, displaying how the chemistry of sodic amphibole evolves with increasing metamorphic grade.

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SUPPLEMENTARY DATA

Supplementary data are available at *Journal of Petrology* online.

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