Original paper Metamorphic reactions and textural changes in coronitic metagabbros from the Teplá Crystalline and Mariánské Lázně complexes, Bohemian Massif

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Coronitic metagabbros occur as small isolated bodies along the contact between the Mariánské Lázně and the Teplá Crystalline complexes in the northwestern part of the Teplá-Barrandian Unit in the Bohemian Massif. Metagabbros show variable metamorphic and textural characteristics with respect to their magmatic mineral assemblage and degree of metamorphism. The aim of this study has been to characterize the mineralogical, chemical, and microstructural changes related to metamorphism in selected samples. Magmatic mineral assemblages in the metagabbros represented by plagioclase, orthopyroxene, clinopyroxene, amphibole, biotite, ilmenite and occasionally involving olivine, spinel or quartz are generally well preserved. Metamorphism is mainly reflected by the formation of single or multiple coronas at the contact of plagioclase with the other primary minerals. The coronas typically consist of amphibole, garnet and orthopyroxene. The progressive breakdown of magmatic plagioclase is reflected by the formation of a mixture of An₄₀ and An_{oo} plagioclase associated with spinel, corundum and locally also kyanite. The calculated P-T conditions show an increase in metamorphic grade towards the structural footwall, i.e. from the east-southeast ($\sim 600 \pm 50$ °C; 10 ± 1.5 kbar) to the west–northwest (\sim 700±50 °C; 13.5±1.5 kbar), which corresponds to the previously reported Variscan metamorphic field gradient in this area. Since there is no gap in P-T conditions between metagabbros included in MLC and TCC and showing similar age and geochemical signature, it is concluded that both complexes were brought together before the gabbro intrusion at \sim 500 Ma. In addition, the maximum pressure of \sim 14 kbar estimated for the metamorphism of the gabbro occurring in the eclogite-bearing Mariánské Lázně Complex suggests that the eclogite-facies metamorphism might have been pre-Variscan. The chemistry of the studied metagabbros corresponds to subalkaline basalts with traceelement signatures characteristic of E-MORB, which is consistent with an interpretation that the intrusion of these rocks was related to an intracontinental rifting of the Teplá-Barrandian Unit during Late Cambrian and was not connected to any subduction processes.

Keywords: corona, gabbro, Variscan metamorphism, Mariánské Lázně Complex, Teplá Crystalline Complex, Cambro–Ordovician extension

Received: 18 December, 2015; accepted: 26 August, 2016; handling editor: P. Hasalová

1. Introduction

The Bohemian Massif represents the easternmost segment of the European Variscides (Fig. 1a), which was consolidated during Late Devonian–Carboniferous subduction–collision process following the closure of the Saxothuringian/ Rheic Ocean (e.g. Matte et al. 1990; Franke 2000; Schulmann et al. 2009). The main evolutionary stages of the Bohemian Massif summarized by Schulmann et al. (2009, 2014) include 1) subduction of the Saxothuringian oceanic plate underneath the Teplá–Barrandian Unit at 430–350 Ma, 2) crustal thickening/relamination of the orogenic root domain at 350–340 Ma, and 3) relatively fast exhumation of the orogen core at 340–330 Ma.

The Teplá–Barrandian Unit (Fig. 1a) represents the uppermost part of the orogenic structure and, at the same

time, it also provides the complete crustal section with little metamorphosed Precambrian and Lower Paleozoic rocks in the east and lower crustal rocks in the west (Peřestý 2012). The westernmost, structurally lowermost and the highest grade part of the Teplá-Barrandian Unit is represented by mainly mafic rocks of the Mariánské Lázně Complex (MLC) and the overlying Proterozoic metasediments of the Teplá Crystalline Complex (TCC). The MLC (Fig. 1b) has been interpreted as a relic of oceanic crust derived from the Saxothuringian/Rheic Ocean that was subducted to eclogite-facies conditions and exhumed during Devonian (Bowes and Aftalion 1991; Jelínek et al. 1997; Zulauf 1997; Dallmeyer and Urban 1998; Faryad 2012). In contrast, the continental crust of the TCC (Fig. 1b) records Cambro-Ordovician rift-related extension and an early Variscan orogenic



Fig. 1a – Simplified map of the Bohemian Massif (modified from Franke 2000). CBPC – Central Bohemian Plutonic Complex, CMP – Central Moldanubian Pluton. Bottom left corner – location of Bohemian Massif in the European Variscides (after Edel et al. 2003) RH – Rhenohercynian Zone, ST – Saxothuringian Zone, M – Moldanubian Zone, B – Brunia. **b** – Geological map of the contact between the Teplá Crystalline and Mariánské Lázně complexes. Simplified from the Geological map of the Czech Republic 1:500 000 (Cháb et al. 2007).

thickening followed by exhumation during Devonian (Peřestý 2012). It is interesting to note that the Cambro–Ordovician rifting was associated with numerous intrusions of gabbroids identified in the TCC but also in the MLC, thus questioning their separate Variscan history (Štědrá et al. 2002; Timmermann et. al. 2004). Moreover it is not clear whether the two complexes shared the same Devonian metamorphic evolution, possibly manifested by the continuous increase in metamorphic grade towards the structural footwall, or whether they were mutually independent with a major metamorphic gap.

In this respect, the metamorphic record in the gabbro lenses, together with their spatial arrangement across the boundary between TCC and MLC bids for a systematic study. This work is focused on coronitic metagabbros, which occur as small isolated bodies in both the mafic rocks of MLC and the felsic rocks of TCC near the contact of the two complexes (Fig. 1b). These mafic magmatic intrusives of Cambro-Ordovician age were metamorphosed during Variscan while escaping the deformation overprint localized into the surrounding country rocks. Advantageously and in contrast to their host rocks, the mechanically resistant gabbro lenses were not affected by the exhumation-related deformation overprint, preserving well the peak metamorphic assemblages. The static metamorphism in the metagabbros thus provides an excellent opportunity to describe and characterize the relationships between the primary igneous minerals and their metamorphic coronitic overprint. In addition, the sequence of metamorphic phases in coronas together with their chemistry is used to estimate P-T conditions of their formation by thermobarometric techniques.

2. Geological setting

2.1. Mariánské Lázně Complex (MLC)

The MLC is dominantly composed of amphibolites, which can be divided into several NE–SW trending belts (Kastl and Tonika 1984; Beard et al. 1995) (Fig. 1b). The central part of the MLC is occupied by the high-grade garnet- and rutile-bearing amphibolite hosting boudins of eclogite and HP granulite (Jelínek et al. 1997; Štědrá 2001; Timmermann et al. 2004; Faryad 2012; Hrouda et al. 2014). The high-grade belt is enveloped by garnet- and titanite-bearing amphibolite and serpentinite body in the northwest that is ascribed to a separate unit (Kachlík 1993).

The MLC mafic rocks are subalkaline and follow the tholeiitic trend (Beard et al. 1995; Timmermann et al. 2004). Major- and trace-element compositions of eclogites fall within the N-MORB and E-MORB fields (Timmermann et al. 2004). Peak metamorphic conditions of 16–23 kbar and 640–715 °C were estimated for the MLC eclogites (Jelínek et al. 1997; Štědrá 2001; Faryad 2012). Faryad (2012) distinguished two separate HP events recorded in the eclogites, the older one at 725 °C and 17 kbar and the younger one at 640 °C and 27–28 kbar. According to O'Brien et al. (1997) the eclogites were overprinted by short-lived granulite-facies metamorphism followed by amphibolite-facies overprint.

Based on the U–Pb zircon dating, the protolith ages of most of the MLC high-pressure rocks were Cadomian (~560–535 Ma: Timmermann et al. 2004). The Variscan metamorphic ages in MLC were constrained to 380–365 Ma by the U–Pb dating of zircon and monazite (Timmermann et al. 2004). Similar cooling ages of 397–370 Ma were obtained by the K–Ar and 40 Ar/³⁹Ar dating of hornblende and interpreted to reflect the quick exhumation of MLC (Kreuzer et al. 1989; Zulauf 1997; Dallmeyer and Urban 1998; Bowes et al. 2002).

2.2. Teplá Crystalline Complex (TCC)

The TCC is dominated by the Proterozoic metasediments with several intrusions of Cambro-Ordovician granitoid plutons (Fig. 1b) transformed into orthogneisses during the Variscan Orogeny (Dörr et al. 1998). The metasediments revealed a Variscan metamorphic field gradient with northeast-southwest trending isograds (Cháb and Žáček 1994; Žáček 1994; Cháb et al. 1997; Zulauf 2001). The metamorphic conditions in the studied area increase towards the NW starting from the staurolite isograd with estimated P-T conditions of 5.0-7.7 kbar and 530-620 °C to the kyanite isograd with 5.0-8.7 kbar and 550-645 °C (Cháb and Žáček 1994; Žáček 1994). The timing of the Variscan metamorphism and exhumation in the TCC is constrained by U-Pb monazite and Ar-Ar muscovite ages ranging between 385 and 370 Ma (Dallmeyer and Urban 1998; Timmermann et al. 2006).

Metagabbro bodies along the contact of the MLC and TCC (Fig. 1b) show well preserved magmatic textures with metamorphic corona overprint (e.g. Štědrá et al. 2002). On the basis of their compositions the metagabbros have been divided into several groups: Ti-rich, Mg-rich, alkali-rich, Rb-rich and transitional composition groups (Štědrá et al. 2002). Major- and trace-element chemistry of the metagabbros is tholeiitic (e.g. Beard et al. 1995; Štědrá et al. 2002; Timmermann et al. 2004). The geochemical character shows affinities to oceanic rocks (Beard et al. 1995) and more recent studies (Štědrá et al. 2002; Timmermann et al. 2004) show that the composition of metagabbros corresponds to E-MORB and T-MORB. Their metamorphic conditions were estimated to 8–11 kbar and 600–730 °C by Štědrá (2001), while slightly higher pressure and lower temperature of 12–13 kbar and 585–614 °C were calculated by Faryad (2012). The intrusive ages of metagabbros of 496–503 Ma were obtained by U–Pb zircon dating (Bowes and Aftalion 1991; Timmermann et al. 2004).

3. Analytical techniques

3.1. Mineral chemistry

Composition of the minerals was determined in the laboratory of electron microscopy at the Institute of Petrology and Structural Geology (Charles University in Prague) using a scanning electron microscope Tescan VEGA equipped with an energy-dispersive spectrometer X-Max 50 (Oxford Instruments). Data were acquired and processed using the software INCA. Quantitative analyses were acquired with an acceleration voltage of 15 kV and a probe current of 1.5 nA, compositional maps showing the distribution of specific elements were acquired at 15 kV and 7 nA. Chemical composition of amphiboles was recalculated and classified following Leake et al. (1997), the X_{Mg} was calculated as $Mg/(Fe^{2+}+Fe^{3+}+Mg)$. Mineral abbreviations used are according to Kretz (1983), Amp is used for amphibole.

3.2. Geothermobarometry

The P–T estimates were obtained by applying the multiequilibrium thermobarometry (average pressure–temperature calculations, Powell and Holland 1994) in the software THERMOCALC v. 3.33 (Powell and Holland 1985, 1988, recent upgrade) using the internally consistent thermodynamic data set (Holland and Powell 1998: November 2003 upgrade). In addition, the P–T conditions were estimated using the "conventional" methods, i.e. the garnet–hornblende geothermometer of Ravna (2000) and the garnet–hornblende–plagioclase geobarometer of Kohn and Spear (1990).

3.3. Whole-rock geochemistry

The chemical analyses of major and trace elements in the selected samples were carried out in order to characterize their magmatic origin. The samples were crushed with a jaw crusher and pulverized in an agate mill in the Laboratories of the Czech Geological Survey (Prague, Czech Republic). Pulverized and homogenized samples were analyzed in the Bureau Veritas Mineral Laboratories (Vancouver, Canada). The total whole rock characterization (code LF202, see http://acmelab.com/ pdfs/FeeSchedule-2016.pdf) was done using the lithium borate fusion ICP-ES (major and minor elements, code LF302) the lithium borate fusion ICP-MS (trace elements, code LF100), and Aqua Regia ICP-ES/MS (trace elements, code AQ200) methods. The geochemical data were plotted and interpreted using the Geochemical Data Toolkit (Janoušek et al. 2006).

4. Petrography and mineral chemistry

The studied samples were collected from six localities near the contact between the MLC and TCC (Fig. 1b, Tab. 1). They were divided into three groups with respect to their location: 1) metagabbros from the TCC; 2) metagabbros from the contact between the MLC and TCC; and 3) metagabbros from the MLC.

4.1. Metagabbros from the TCC

The sample **PJ5** shows high degree of metamorphic recrystallization although the primary magmatic texture is still well preserved (Fig. 2). The observed mineral phases can be assigned either to the primary (presumably magmatic) assemblage M_1 or to the secondary metamorphic assemblage M_2 , the latter resulting from recrystallization of the primary minerals and development of coronas at their contacts. The M_1 assemblage is generally coarse-grained and consists of plagioclase, amphibole, clinopyroxene, biotite, and ilmenite. Plagioclase grains up to 2 mm preserve euhedral shape

Tab. 1 Sample list with geographic coordinates, localities and their characteristics

locality	Ν	E		surrounding rocks
TCC				
PJ 5	49.94146292	13.01445046	NW of Úterý	micaschist (Ky zone)
TCC-MLC				
PJ 8	49.96539376	12.90800532	SE of Teplá	orthogneiss
PJ 7	49.97590793	12.86426051	E of Teplá	amphibolite, Ky/Sil-bearing gneiss
VP 46	50.09017808	12.98426924	near Chylice	gneiss
MLC				
PJ 6	50.02928825	12.91391029	near Otročín	garnet amphibolite with titanite
PJ 9	49.93372710	12.78617830	W of Výškovice	garnet amphibolite with titanite

and their composition ranges from andesine to labradorite (An₄₉₋₅₄; Fig. 3a) with increasing Na and decreasing Ca contents (Ab = 46 \rightarrow 54 %, An_{53 \rightarrow 46}, Or = 0–1 %) at their rims. Amphibole (pargasite–edenite, X_{Mg} = 0.51–0.58, Si = 6.18– 6.63 apfu; Fig. 3e; Tab. 2) together with clinopyroxene (augite, X_{Mg}=0.75–0.76 and Jd = 2.5–4 mol. %; Fig. 3c;



Fig. 2 Back-scattered electron images characterizing the mineral assemblage of the TCC metagabbro: \mathbf{a} – Polycrystalline aggregate consisting of clinopyroxene and amphibole in plagioclase matrix; \mathbf{b} – Detail of ilmenite grain with titanite rim and zircon grains; \mathbf{c} – Amphibole and garnet corona; \mathbf{d} – Detail of garnet corona with inclusions; \mathbf{e} – Detail of plagioclase recrystallization into bytownite and andesine mixture.

Tab. 2) usually form polycrystalline aggregates (Fig. 2a). These aggregates mostly have clinopyroxene and amphibole crystals in the central parts, and amphibole with ilmenite lamellae at their margins. Biotite grains

with $X_{Mg} = 0.57-0.58$ and Ti = 2.7-2.9 apfu (Tab. 2) locally occur in association with ilmenite, which is commonly surrounded by tiny (<1µm) zircon grains (Fig. 2b).



Fig. 3 Diagrams demonstrating composition of the main rock-forming minerals of all samples: olivine (a), plagioclase (b), pyroxenes (c) after Morimoto (1988), garnet (d) and amphibole (e-f) after Leake et al. (1997).

The M_2 mineral assemblage is characterized by growth of amphibole, garnet, plagioclase and titanite at the expense of M_1 minerals, which are enveloped by several

types of the corona sequence. The M_1 clinopyroxene occurs either alone or it is surrounded by an amphibole corona up to 200 μ m thick. M_1 amphibole is rimmed by

unit		TCC					TCC	MLC					MLC	
sample		PJ 5			P.	J 8		VP	46			ΡJ	6	9 I d
mineral	Cpx	Amp	Bt	Ю	Ρl	Cpx	Opx	Amp	Spl	Bt	Opx	Cpx	Amp	Amp
SiO_2	53.24	44.02	36.77	37.24	50.58	52.00	55.12	44.21	0.18	39.37	52.13	52.1	42.28	46.51
TiO_2	0.16	1.62	4.06	b.d.	b.d.	0.83	0.08	2.93	0.86	2.42	0.13	0.36	1.59	1.93
Cr_2O_3	0.37	0.28	b.d.	b.d.	b.d.	0.65	b.d.	0.59	26.08	b.d.	b.d.	b.d.	b.d.	b.d.
Al_2O_3	1.36	11.35	15.11	b.d.	31.50	3.05	0.43	10.88	20.88	14.95	0.97	2.39	13.12	9.61
FeO	7.66	14.32	16.73	30.00	b.d.	5.93	16.08	9.70	46.35	7.48	24.84	8.83	14.44	9.89
MnO	0.31	0.18	b.d.	0.30	b.d.	0.16	0.51	0.20	0.47	b.d.	0.69	0.26	0.16	0.09
MgO	13.42	10.94	12.96	32.30	b.d.	15.35	26.50	14.18	3.03	20.70	20.09	12.68	11.56	14.96
CaO	23.14	11.76	0.07	b.d.	14.05	20.45	1.06	11.06	b.d.	b.d.	0.53	22.07	11.12	11.38
Na_2O	0.50	1.38	0.16	b.d.	3.54	1.13	b.d.	2.99	b.d.	0.83	b.d.	0.72	2.45	1.87
K_2O	b.d.	0.83	9.70	b.d.	b.d.	b.d.	b.d.	0.92	b.d.	9.21	b.d.	0.00	0.64	0.58
V_2O_5	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	0.58	b.d.	b.d.	b.d.	b.d.	b.d.
CI	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	0.05	b.d.	b.d.	b.d.	b.d.
ZnO	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	0.93	b.d.	b.d.	b.d.	b.d.	b.d.
total	100.16	96.68	95.56	99.84	99.66	99.54	99.78	97.66	99.36	95.01	99.38	99.41	97.36	96.82
cat/O	4/6	13+Ca+Na+K/23	16/22	3/4	5/8	4/6	4/6	13+Ca+Na+K/23	3/4	16/22	4/6	4/6	13+Ca+Na+K/23	13+Ca+Na+K/23
Si	1.98	6.56	5.67	1.01	2.31	1.91	2.00	6.45	0.01	5.74	1.98	1.95	6.22	6.72
Ti	0.00	0.18	0.47	0.00	0.00	0.02	0.00	0.32	0.02	0.25	0.00	0.01	0.18	0.21
Cr	0.01	0.03	0.00	0.00	0.00	0.02	0.00	0.07	0.68	0.00	0.00	0.00	2.27	1.64
Al	0.06	1.99	2.75	0.00	1.69	0.13	0.02	1.87	0.81	2.55	0.04	0.11	0.00	0.00
$\mathrm{F}e^{2+}$	0.24	1.77	2.16	0.68	0.00	0.12	0.49	1.18	0.87	0.92	0.79	0.26	1.73	1.17
Fe^{3+}	0.00	0.02	0.00	0.00	0.00	0.06	0.00	0.01	0.41	0.00	0.00	0.02	0.07	0.04
Mn	0.01	0.02	0.00	0.01	0.00	0.00	0.02	0.02	0.01	0.00	0.02	0.01	0.02	0.01
Mg	0.74	2.43	0.00	1.30	0.00	0.84	1.43	3.08	0.00	0.02	1.14	0.71	2.53	3.22
Ca	0.92	1.88	2.98	0.00	0.69	0.81	0.04	1.73	0.15	4.58	0.02	0.89	1.75	1.76
Na	0.04	0.40	0.01	0.00	0.31	0.08	0.00	0.85	0.00	0.00	0.00	0.05	0.70	0.52
K	0.00	0.16	0.05	0.00	0.00	0.00	0.00	0.17	0.00	0.20	0.00	0.00	0.12	0.11
>	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.01	1.74	0.00	0.00	0.00	0.00
CI	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.01	0.00	0.00	0.00	0.00
Zn	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.02	0.00	0.00	0.00	0.00	0.00
X_{Mg}	0.76	0.58	0.58	0.66		0.82	0.75	0.72	0.14	0.83	0.59	0.72	0.58	0.73
End mem	nbers (mol.	(%)		An	69.0									
				$^{\mathrm{Ab}}$	31.0									

samplemineralP1 SiO_2 46.54 SiO_2 $b.d.$ Cr_2O_3 $b.d.$ $A1_2O_3$ 34.06 FeO $b.d.$ MO $b.d.$ MO $b.d.$ CaO 17.24	Pl 56.49	J 5														
mineral Pl SiO2 46.54 TiO2 b.d. TiO2 b.d. Cr2O3 b.d. Al2O3 34.06 FeO b.d. MRO b.d. MRO b.d. Cato 17.24	Pl 56.49			PJ 7				VP 46			PJ 6			PJ 9		
SiO ₂ 46.54 TiO ₂ b.d. Cr ₂ O ₃ b.d. Al ₂ O ₃ 34.06 FeO b.d. MnO b.d. MgO b.d. CaO 17.24	56.49	Amp	Amp (Pl)	Amp (Opx)	Cum	Opx	Opx	Grt (Amp)	Grt core	(Pl)	Spl	Cpx	Opx	Amp	Pl rim	ol core
TiO2 b.d. Cr2O3 b.d. Al2O3 34.06 FeO b.d. MnO b.d. MgO b.d. CaO 17.24		48.32	42.11	51.19	54.44	55.55	55.82	37.56	38.83	38.9	0.12	53.96	54.41	48.54	43.84	59.21
Cr ₂ O ₃ b.d. Al ₂ O ₃ 34.06 FeO b.d. MnO b.d. MgO b.d. CaO 17.24	b.d.	0.95	0.33	0.15	b.d.	0.13	b.d.	b.d.	b.d.	b.d.	b.d.	0.09	b.d.	0.84	b.d.	b.d.
Al ₂ O ₃ 34.06 FeO b.d. MnO b.d. MgO b.d. CaO 17.24	b.d.	0.28	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	0.42	b.d.	b.d.
FeO b.d. MnO b.d. MgO b.d. CaO 17.24	27.61	7.76	14.07	4.23	0.44	0.6	1.03	21.13	21.86	21.99	61.63	0.98	0.86	7.53	35.57	25.55
MnO b.d. MgO b.d. CaO 17.24	b.d.	12.72	16.28	14.09	22.51	16.95	14.66	25.67	23.99	23.61	22.16	5.59	17.38	8.42	b.d.	0.11
MgO b.d. CaO 17.24	b.d.	0.22	0.12	0.29	0.40	0.36	0.32	0.89	0.54	0.66	b.d.	0.09	0.23	0.13	b.d.	b.d.
CaO 17.24	b.d.	13.38	9.31	14.70	17.80	26.80	28.09	5.68	6.36	5.97	13.08	15.29	26.05	16.38	b.d.	b.d.
	9.61	12.06	11.78	10.86	0.83	0.39	0.19	7.61	8.47	9.38	0.27	23.47	0.16	12.04	19.48	7.42
Na ₂ O 1.80	6.04	0.91	0.96	0.31	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	0.45	b.d.	1.38	0.65	7.47
K ₂ O b.d.	0.14	0.43	1.26	0.19	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	0.27	b.d.	0.06
ZnO b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	0.37	b.d.	b.d.	b.d.	b.d.	b.d.
total 99.64	99.89	97.03	96.22	96.01	96.42	100.78	100.11	98.54	100.1 1	00.50	97.63	99.92	90.09	95.95	99.54	99.82
cat/O 5/8	5/8	13+Ca+Na+K/23	13+Ca+Na+K/23	13+Ca+Na+K/23	13+Ca+Na+K/2	3 4/6	4/6	8/12	8/12	8/12	3/4	4/6	4/6	13+Ca+Na+K/23	5/8	5/8
Si 2.14	2.54	7.04	6.75	7.51	7.03	2.00	2.00	2.97	3.00	2.99	0.01	1.98	1.99	7.02	2.03	2.65
Ti 0.00	0.00	0.10	0.04	0.02	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.09	0.00	0.00
Cr 0.00	0.00	0.03	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.05	0.00	0.00
Al 1.85	1.46	1.33	1.78	0.49	0.07	0.03	0.04	1.97	1.99	1.99	1.95	0.04	0.04	1.28	1.94	1.35
${\rm Fe}^{2+}$ 0.00	0.00	1.53	2.18	1.66	2.13	0.51	0.44	1.59	1.53	1.49	0.67	0.16	0.53	0.99	00.00	0.00
${ m Fe}^{3+}$ 0.00	0.00	0.03	0.00	0.10	0.64	0.00	0.00	0.10	0.02	0.03	0.03	0.01	0.00	0.03	00.00	0.00
Mn 0.00	0.00	0.03	0.02	0.04	0.04	0.01	0.01	0.06	0.04	0.04	0.00	0.00	0.01	0.02	0.00	0.00
Mg 0.00	0.00	2.91	2.23	3.22	3.43	1.44	1.50	0.67	0.73	0.68	0.00	0.84	1.42	3.53	0.00	0.00
Ca 0.85	0.46	1.88	2.02	1.71	0.11	0.02	0.01	0.64	0.70	0.77	0.28	0.92	0.01	1.86	0.97	0.36
Na 0.16	0.53	0.26	0.30	0.09	0.00	0.00	0.00	0.00	0.00	0.00	0.01	0.03	0.00	0.39	0.06	0.65
K 0.00	0.01	0.08	0.26	0.04	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.05	0.00	0.00
Zn 0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.05	0.00	0.00	0.00	0.00	0.00
X_{Mg}		0.65	0.50	0.65	0.55	0.74	0.77	0.28	0.32	0.31	0.29	0.83	0.73	0.78		
End members (m	nol. %)															
An 84.0	46.0						Prp	21.1	24.2	22.5				An	94.0	35.0
Ab 16.0	53.0						Alm	50.4	50.5	49.0				Ab	6.0	64.0
Or 0.0	1.0						Grs	20.4	23.1	25.4				Or	0.0	1.0
							Sps	1.9	1.2	1.4						

a 50-250 µm thick corona of M, amphibole, which is locally followed by a 20-200 µm thick garnet corona (Fig. 2c). M, amphibole is magnesiohornblende (Fig. 3f, Tab. 3) characterized by an increase in X_{Mg} (0.59 \rightarrow 0. 67), Si (6.80 \rightarrow 7.04 apfu) and decrease in Na (0.33 \rightarrow 0.26), K $(0.14 \rightarrow 0.08)$, Al $(1.7 \rightarrow 1.3)$ and Ti $(0.16 \rightarrow 0.10 \text{ apfu})$ in the direction from the primary amphibole and clinopyroxene towards the surrounding plagioclase. Garnet corona contains numerous plagioclase and amphibole inclusions up to 40 μ m across together with small (<1 μ m) elongated crystals of kyanite (Fig. 2d). Garnet is almandine with composition Alm₅₂₋₅₈Grs₁₈₋₂₂Prp₁₆₋₂₂Sps₂₋₆ (Fig. 3d). Garnet coronas show a slightly asymmetric compositional zoning, with respect to its central part (Alm₅₆Prp₂₂ $Grs_{18}Sps_3$; $X_{Mg} = 0.28$), characterized by a rimward decrease in $X_{M_{F}}$, Prp and Alm, and increase in Grs and Sps components. The asymmetry is revealed by a more significant chemical change towards the contact with amphibole (Alm₅₄Prp₁₈Grs₂₀Sps₄; $X_{Mg} = 0.25$) compared to the contact with plagioclase (Alm₅₃Prp₂₁Grs₂₂Sps₂; X_{Mg} = 0.28). The primary plagioclase (and esine-labradorite) is extensively replaced by a fine-grained mixture of plagioclase with two distinct compositions, calcic (bytownite; An₈₅) and sodic (andesine, An₄₅; Fig. 2e, Tab. 3), whereby the former forms elongated ribbon-like grains in the matrix dominated by the latter (Fig. 2e). Primary ilmenite occurs in association with zircon and it is locally surrounded by a thin titanite rim (Fig. 2b).

Additionally, epidote is occasionally present in the central part of the clinopyroxene and amphibole aggregates together with calcic amphibole (actinolite to hornblende, $X_{Mg} = 0.67-0.69$; Si = 7.22–7.54, Ti = 0.02–0.03, K = 0.03–0.05 and Na = 0.16–0.22 apfu) and cummingtonite ($X_{Mg} = 0.55-0.56$, Si = 7.01–7.03 apfu). Plagioclase is in places decomposed into a fine-grained mixture of albite, muscovite and epidote forming narrow zones along the grain margins or cross-cutting the primary plagioclase. Prehnite is locally present in association with titanite; biotite is partially replaced by chlorite.

4.2. Metagabbros at the contact of the TCC and MLC

Three samples with best preserved primary mineral assemblage M_1 were collected at the contact of TCC and MLC. Two of them, PJ8 and VP46, are olivine metagabbros (Fig. 4). The third sample (PJ7) contains an olivine-free M_1 mineral assemblage, which includes quartz (Fig. 5).

The M_1 assemblage of olivine metagabbro **PJ8** consists of plagioclase, olivine, and clinopyroxene with minor biotite and ilmenite, and accessoric apatite and pyrrhotite. Olivine ($X_{Mg} = 0.65-0.71$; Fig. 3b; Tab. 2) forms grains (<2 mm) with inclusions of Cr-spinel and

pyrrhotite. It is commonly crosscut by fractures filled by amphibole or minerals of the serpentine group (Fig. 4a). Plagioclase forms up to 3 mm large crystals (An₅₆₋₇₀; Tab. 2). Clinopyroxene (augite, $X_{Mg} = 0.81-0.86$ and Jd = 2-5 mol. %; Fig. 3c; Tab. 2) occurs as up to 3 mm large crystals containing inclusions of olivine, ilmenite, and occasionally Cr-spinel. Orthopyroxene (enstatite with $X_{Mg} = 0.74-0.75$; Fig. 3c; Tab. 2) is present rarely as a part of clinopyroxene-amphibole-olivine aggregates (Fig. 3a) and it contains inclusions of Cr-spinel. Amphibole (pargasite-edenite with $X_{Mg} = 0.76-0.77$, Si = 6.38-6.42 apfu and 0.30-0.36 apfu of Ti) usually occurs within clinopyroxene-orthopyroxene aggregates and is locally surrounded by, or includes tiny (<1 µm) crystals of, ilmenite (Fig. 4a). Biotite (phlogopite with $X_{M_{g}}$ = 0.79-0.81 and 1.81-2.05 apfu of Ti) forms up to 1 mm large crystals, which locally contain inclusions of ilmenite. Ilmenite is in places associated with biotite and its rims are commonly surrounded by small zircon crystals.

The M₂ mineral assemblage is characterized by the formation of coronas composed of orthopyroxene, amphibole, and spinel around the M1 minerals and by plagioclase recrystallization. Clinopyroxene is surrounded by a single layer of amphibole (pargasite, $X_{M_{\pi}} = 0.70-0.72$, Si = 5.93-6.03 apfu). Biotite and olivine grains at the contact with plagioclase are usually enveloped by double corona of orthopyroxene (<100 μ m, enstatite with X_{Mg} = 0.70–0.71) followed by a layer (<200 µm) of amphibole (pargasite with $X_{Mg} = 0.71-0.73$). Amphibole corona occasionally contains embayments of plagioclase and spinel symplectites along the contact with plagioclase (Fig. 4b). Plagioclase is locally recrystallized into a mixture of oligoclase (An_{24-29}) and anorthite $(An_{92}; Fig. 3a)$, in restricted domains along the rims of primary grains. These domains contain tiny (0.X µm) spinel grains and small tabular corundum crystals (Fig. 4b).

The second olivine metagabbro, VP46, shows similar primary mineral assemblage as the sample PJ8: olivine, plagioclase, clinopyroxene, amphibole, biotite and ilmenite; however the M₁ assemblage is less affected by the M_2 overprint. Olivine ($X_{Mg} = 0.70-0.75$) usually occurs as large crystals (up to 3 mm in size, Fig. 4c), commonly containing tiny ($<0.X \mu m$) inclusions of calcic amphibole associated with magnetite (Fig. 4c). Plagioclase forms up to 5 mm large crystals generally not affected by recrystallization (Fig. 4c), which show rather variable composition (Fig. 3d). The plagioclase domains are composed of sectors with contrasting chemistry ranging from oligoclase (An₁₈₋₂₅) to labradorite (An₅₃, Fig. 3a). Clinopyroxene (augite-diopside, $X_{Mg} = 0.77-0.80$ and Jd 3-7 mol. %) forms euhedral crystals (<1 mm) with spinel inclusions. Clinopyroxene shows compositional zoning (Fig. 6a) with distinct trends determined by the particular M, phase at the contact. The clinopyroxene cores are relatively





Fig. 5 Back-scattered electron images characterizing the mineral assemblage of the olivine-free metagabbro from the TCC–MLC boundary. \mathbf{a} – Primary magmatic texture defined by large orthopyroxene, clinopyroxene, ilmenite, and plagioclase; \mathbf{b} – Detail of plagioclase recrystallized into mixture of bytownite and andesine, including small grains of biotite; \mathbf{c} – Detail of double corona around clinopyroxene formed by quartz–calcic amphibole symplectite followed by monomineral amphibole layer; \mathbf{d} – Detail of orthopyroxene with ilmenite inclusions enveloped by triple corona of 1) cummingtonite, 2) calcic amphibole (actinolite) with quartz and 3) garnet layer.

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Fig. 4 Back-scattered electron images characterizing the mineral assemblage of the olivine-bearing metagabbros from the TCC–MLC boundary: **a** – Detail of primary olivine, clinopyroxene, amphibole (Amp₁), and orthopyroxene (Opx₁) surrounded by amphibole corona; **b** – Detail of amphibole corona with symplectite consisting of plagioclase and spinel; in the upper part of the picture recrystallization of plagioclase into anorthite and oligoclase with spinel and corundum; **c** – Primary clinopyroxene, amphibole and olivine enveloped by orthopyroxene and amphibole coronas; in bottom left corner detail of inclusions in olivine of magnetite associated with amphibole; **d** – Detail of ilmenite grain with magnetite and spinel lamellae, which is surrounded by symplectite-like texture accompanied by zircon grains; **e** – Corona sequences around olivine represented by orthopyroxene and amphibole layers alternating with spinel–plagioclase symplectite; primary biotite is surrounded by amphibole alternating with spinel–plagioclase symplectite.

Sample VP 46	
(a)	and the second
Amp ₁ Amp ₁	
Cpx	
PI PI	
Срх	
Amp ₂	
	·
<u>Opx</u> <u>0.2 mm</u>	
a share the state of the state	A State of the second s
BOARD STREET	An TE MALLER AND A REAL
Start Start Starts	
1 4 3 F 1 - 1 - 1 - 2	
A	
Sample PJ 7	
PI Amp ₂ Cpx	
Bt	
IIm Amp ₂ +Qtz	
and the second states and the	
The second	
Opx	
<u>0.2 mm</u>	A
Sample PJ 6	
(C) Den 200	
L' Brack	
· Portes	
· Aller A state	A A A A A A A A A A A A A A A A A A A
Amp ₂	
Grt Bt	
0.1 mm	
Sample PJ 9	
Grt	
Opx.	
Amp ₂ +Cpx ₂	
Opx ₂ +Pl 0.2 mm	

rich in Cr (Cr = 0.04, Na = 0.07, Al = 0.18, Ti = 0.03and Ca = 0.77 apfu; $X_{Mg} = 0.80$). The zoning towards the contact with plagioclase is characterized by a decrease in Cr, Na, Ca and $X_{_{\ensuremath{Mg}}}$ and an increase in Ti and Al (Cr = 0.01, Na = 0.05, $A\ddot{l} = 0.24$, Ti = 0.05, and Ca = 0.76 apfu; $X_{M_{rg}} = 0.77$). In contrast, towards the contact with M₁ amphibole the zoning shows a decrease in Cr, Na, Al, Ti, Ca, and X_{Mg} and an increase in Ca (Cr = 0.01, Na = 0.04, A1 = 0.06, Ti = 0.01 and Ca = 0.80 apfu; $X_{M\sigma} = 0.79$). Orthopyroxene (enstatite, $X_{Mg} = 0.70-0.78$) is observed occasionally between olivine and biotite grains and it contains inclusions of ilmenite and spinel. Anhedral crystals of amphibole (pargasite-edenite; Fig. 3e, Tab. 3) are associated with clinopyroxene (Fig. 4c) and they reveal compositional zoning (Fig. 6a) with a rimward decrease in Ti, Al and Na contents accompanied by an increase in Cr, Si and X_{Mg} (Ti = 0.58 \rightarrow 0.32, Al = 1.99 \rightarrow 1.87, Na = $0.92 \rightarrow 0.85$, $Cr = 0.05 \rightarrow 0.07$, $Si = 6.23 \rightarrow 6.51$ apfu, X_{Mg} = 0.70 \rightarrow 0.72). Biotite (phlogopite, X_{Mg} = 0.80–0.85 with highly variable Ti of 0.75-3.04 apfu; Tab. 2) forms up to 0.5 mm large crystals with inclusions of ilmenite observed between olivine grains. Ilmenite commonly occurs in association with Cr-spinel (Cr = 0.48-0.68 apfu; Tab. 2) and magnetite, which can also form lamellae inside ilmenite crystals (Fig. 4d). Ilmenite is usually rimmed by small zircon crystals (Fig. 4d).

The M₂ mineral assemblage is connected with formation of coronas including orthopyroxene, amphibole, and spinel. Olivine is surrounded by double corona consisting of up to 100 µm wide orthopyroxene domain (enstatite with X_{Mg} = 0.76–0.78; Fig. 3c; Tab. 3) followed by 10–60 µm wide corona of amphibole (pargasite, X_{Mg} = 0.74–0.79, Si = 5.63–6.01 apfu), which is occasionally alternating with an orthopyroxene–spinel symplectite (Fig. 4c, e). No corona occurs around clinopyroxene or amphibole. The contact between biotite and plagioclase is marked by a symplectitic corona (5–25 µm thick) of spinel and orthopyroxene, which is in places alternating with amphibole corona (pargasite, X_{Mg} = 0.77; Fig. 4f).

Sample **PJ7** is the only one that contains quartz as a part of the primary mineral assemblage: plagioclase, clinopyroxene, orthopyroxene, ilmenite, biotite and quartz (Fig. 5a). In addition, this sample is generally

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richer in ilmenite and plagioclase compared to the other samples. Although large crystals of the primary plagioclase preserve their original crystal shapes with twinning still recognizable, they recrystallized completely into the fine-grained mixture of sodic and calcic plagioclase. The integrated original composition corresponds to labradorite (An_{60}) . Clinopyroxene containing orthopyroxene and ilmenite lamellae (augite-diopside, $X_{M_{\sigma}} = 0.67$ and Jd = 2.3-2.7 mol. %; Fig. 3c) and orthopyroxene with lamellae of clinopyroxene and ilmenite (enstatite with X_{Mg} = 0.51-0.56) form crystals up to 3 mm across (Fig. 5a). Large orthopyroxene grains show slight compositional zoning with $X_{M_{\sigma}}$ decreasing from 0.55 to 0.52 towards their rims. Ilmenite (Fig. 5a) is usually associated with both pyroxenes and biotite and forms anhedral crystals up to 1 mm that are usually surrounded by small ($<3 \mu m$) zircon crystals.

The M₂ mineral assemblage is associated with formation of coronas composed of garnet and compositionally variable amphibole (Fig. 3e-f) and with recrystallization of the M₁ plagioclase. The primary ilmenite is surrounded by up to 100 µm wide garnet corona with inclusions of amphibole, dolomite, quartz, and kyanite. Garnet has a composition of Alm₅₅₋₅₉Grs₁₈₋₂₈Prp₁₅₋₁₇Sps₃₋₄ (Fig. 3d) and shows compositional zoning characterized by increasing Alm, Prp and Sps, and decreasing Grs contents from core to the rim. The zoning is slightly asymmetric with more significant compositional change towards the contact with ilmenite compared to the contact with plagioclase (ilmenite \rightarrow plagioclase: Alm_{58 \rightarrow 51 \rightarrow 56}Prp_{15 \rightarrow 14 \rightarrow 17}Grs_{18 \rightarrow 28} $\sum_{j=21}^{3} \operatorname{Sps}_{4 \to 3 \to 4j}; X_{Mg} = 20 \to 21 \to 22). \text{ Occasionally, a nar-}$ row corona of amphibole is present between ilmenite and the garnet domain. Clinopyroxene is enveloped by a double corona composed of a 10–100 µm thick symplectite layer consisting of amphibole and quartz that is followed by a monomineral amphibole layer up to 100 µm in width (Fig. 5c). The systematic change in amphibole composition (Fig. 6b) is demonstrated by a continual decrease of X_{Mg} and an increase in Al, Ca, K, and Na towards plagioclase in both amphibole-quartz symplectite layer (actinolite-magnesiohornblende; X_{Mg} = $0.65 \rightarrow 0.61$; Si = $7.51 \rightarrow 7.24$, Al = $0.49 \rightarrow 0.94$, Ca = $1.78 \rightarrow 1.90$, Na = 0.09 $\rightarrow 0.20$ and K = 0.04 $\rightarrow 0.09$ apfu) and the outer monomineralic amphibole zone (edenite; $X_{Mg} = 0.60 \rightarrow 0.51$; Si = 7.28 \rightarrow 6.75, Al = 0.98 \rightarrow 1.78, Ca $= 1.95 \rightarrow 2.02$, Na = 0.18 $\rightarrow 0.3$ and K = 0.11 $\rightarrow 0.25$ apfu). The garnet corona is locally developed at the contact of the M₂ amphibole with plagioclase. The most complex corona sequence is developed around the primary orthopyroxene. In the direction from orthopyroxene to plagioclase, the multiple corona consists of cummingtonite domain ($X_{Mg} = 0.55$; Si = 6.97-7.03; Tab. 3), symplectite layer of amphibole (hornblende with X_{Mg} = 0.65, Si = 7.28 apfu) and quartz, followed by garnet

Fig. 6 Compositional maps characterizing the representative minerals. **a** – Zoning of clinopyroxene, amphibole and corona sequence around olivine in the olivine-bearing sample VP 46 from the TCC–MLC boundary: BSE picture of the analysed area (left) and distributions of Cr, Al, and Ti (right); **b** – Zoning of amphibole + quartz symplectite and monomineral amphibole corona around clinopyroxene in sample PJ 7 from the TCC–MLC boundary: BSE image, Al distribution; **c** – Zoning of garnet corona between primary biotite and recrystallized plagioclase in the MLC sample PJ 6: BSE image, Mn, Fe, and Mg distributions; **d** – Corona sequence around orthopyroxene and character of plagioclase recrystallization into fine-grained matrix in the MLC sample PJ 9: BSE image, Al, Na, and Ca distributions.

layer with amphibole, quartz, and kyanite inclusions (Fig. 5d). This garnet corona shows asymmetric compositional zoning generally identical to that formed around ilmenite. The primary plagioclase is decomposed into a fine-grained mixture of andesine (An_{41-50}) and bytownite (An_{80} ; Figs 3a; 5b), the latter forming small elongated crystals embedded in the andesine matrix. Ilmenite, biotite, amphibole, and quartz occur within the recrystallized plagioclase matrix.

4.3. Metagabbros from the MLC

Metagabbros from the MLC, represented by samples PJ6 and PJ9 collected at Otročín and Výškovice, are characterized by the relatively intense recrystallization of the primary minerals; however the primary microstructure is still recognizable. In both samples, the primary mineral assemblage M_1 includes plagioclase, clinopyroxene, orthopyroxene, amphibole, biotite, and ilmenite, although the degree of recrystallization is different.

In sample PJ6, plagioclase (labradorite, An_{65-67}) is intensively recrystallized. Clinopyroxene (augite, $X_{Mg} =$ 0.71–0.82 and Jd = 5–7 mol. %; Fig. 3c; Tab. 2) occurs as large (<2 mm) crystals containing numerous lamellae of orthopyroxene and ilmenite. Orthopyroxene (enstatite, $X_{Mg} = 0.57-0.59$; Fig. 3c; Tab. 2) forms individual grains up to 0.5 mm across (Fig. 5a). Amphibole (pargasite; $X_{Mg} = 0.58-0.65$; Si = 6.04–6.40 apfu; Fig. 3e; Tab. 2) occurs as crystals up to 0.4 mm large (Fig. 7a). Biotite (with $X_{Mg} = 0.62-0.68$ and Ti = 0.24–0.55 apfu) is usually observed in association with ilmenite, the latter forming large grains (up to 1 mm) commonly surrounded by small crystals of zircon.

The M₂ mineral assemblage connected with recrystallization of the M₁ assemblage and formation of coronitic and symplectitic texture around primary minerals is represented by plagioclase, orthopyroxene, amphibole, and garnet. Similarly to the other samples, plagioclase is decomposed into a mixture of elongated crystals of bytownite–anorthite (An_{83-91}) with random orientation embedded in matrix formed by oligoclase-andesine (An₂₈₋₄₇; Fig. 3a). This microstructure occurs preferentially in the central parts of the large plagioclase crystals together with small spinel grains (Fig. 7b, Tab. 3). Clinopyroxene is usually surrounded by amphibole (pargasite, X_{Mg} = 0.54-0.58, Si = 5.90-6.08 apfu) corona, 10-100 μm in width. However at the contact with plagioclase, it is often enveloped by symplectites formed by amphibole-plagioclase or orthopyroxene-plagioclase. Similarly to clinopyroxene, the primary amphibole and orthopyroxene are enveloped by orthopyroxene-plagioclase or amphiboleplagioclase symplectites (Fig. 7a). Locally, the amphibole zone around orthopyroxene and biotite is followed by a garnet corona containing inclusions of plagioclase, amphibole, kyanite, and occasionally clinopyroxene. Garnet is dominated by the almandine component (Alm₅₀Prp₂₀₋₂₆ Grs₂₀₋₂₆Sps₁₋₂; Fig. 3d). This corona is characterized by a slight asymmetric zoning (Fig. 6c, Tab. 3) characterized by a rimward decrease in X_{Mg}, Alm and Prp components and, more significant, decrease in Prp and X_{Mg} towards the contact with amphibole. The Grs content continuously increases while Sps decreases across the corona from amphibole towards plagioclase (Alm_{50→51→49}Prp_{21→24→23} Grs_{20→23→25}Sps_{2→1→1}; X_{Mg} = 0.28→0.32→0.31). The primary mineral assemblage M₁ in sample PJ9

The primary mineral assemblage M_1 in sample PJ9 consists of clinopyroxene and orthopyroxene that are surrounded by plagioclase matrix. Plagioclase is completely recrystallized, but it still preserves its euhedral shape especially when locked in pyroxene (Fig. 7c). Clinopyroxene (augite–diopside, $X_{Mg} = 0.79-0.80$, Jd= 3–10 mol. %; Fig. 3c) forms up to 3 mm large crystals with orthopyroxene lamellae and inclusions of amphibole. The orthopyroxene (enstatite, $X_{Mg} = 0.67-0.73$) grains up to 1 mm across contain clinopyroxene lamellae and inclusions of amphibole, apatite, and rutile. Amphibole (pargasite–edenite, $X_{Mg} = 0.70-0.73$, Si = 6.44–6.72 apfu) occurs usually in association with orthopyroxene or clinopyroxene and occasionally contains inclusions of apatite and rutile. Ilmenite is mostly associated with biotite and it is usually surrounded by zircon and locally replaced by rutile (Fig. 7d).

Several types of coronas were formed around primary minerals in PJ9. Clinopyroxene is mostly surrounded by a narrow (20–60 µm thick) layer of amphibole (Fig. 7c) showing compositional zoning from magnesiohornblende at the contact with clinopyroxene to edenite next to plagioclase (Fig. 3e-f). This change is accompanied by a decrease of $X_{_{Mg}},\,Si$ and an increase of Ti, K and Na (X $_{_{Mg}}$ $= 0.78 \rightarrow 0.68$; Si = 7.02-6.17 apfu, Ti = 0.09 $\rightarrow 0.17$, Na = $0.39 \rightarrow 0.67$ and K = $0.05 \rightarrow 0.14$ apfu, Tab. 3). Orthopyroxene is enveloped by a complex corona sequence (Fig. 7e). In the direction from orthopyroxene to plagioclase, 30–200 µm thick zone of mixed amphibole (magnesiohornblende, $X_{Mg} = 0.75-0.83$; Si = 6.83-7.44 apfu; Fig. 3f) and clinopyroxene (augite-diopside, X_{Mg} = 0.82 - 0.84 and Jd = 14 - 18 mol. %; Fig. 3c) is followed by 40-200 µm thick domain of orthopyroxene-plagioclase symplectite and the sequence is completed by up

Fig. 7 Back-scattered electron images characterizing the mineral assemblage of the MCC metagabbro. **a** – Magmatic amphibole with orthopyroxene and plagioclase; **b** – Detail of primary magmatic plagioclase breakdown into andesine and bytownite with spinel; **c** – Detail of plagioclase inclusion in clinopyroxene with amphibole corona developed at the contact of both minerals; **d** – Detail of an ilmenite grain, partially replaced by rutile and accompanied by small zircon grains; **e** – Corona sequence surrounding orthopyroxene consisting of clinopyroxene and amphibole layer followed by orthopyroxene–plagioclase symplectite and garnet layer; **f** – Primary plagioclase recrystallized into fine-grained matter with spinel, corundum, and kvanite grains.



			Τ	Ŋ										MLC							
sample			[]	5					ΡJ	9							PJ 9				
nethod		geoTB			av PT			geoTB		geo	TB + av	ΡT			geo	TB				av PT	
nineral	Amp	PI	Grt	Amp	ΡI	Grt	Amp	Pl	Grt	Amp	PI	Grt	Amp	PI	Grt	Amp	Pl	Grt	Amp	PI	Grt
SiO ₂	44.80	57.08	38.32	45.09	55.68	38.42	40.80	59.50	37.56	39.51	58.07	38.61	46.51	59.39	39.48	44.80	60.05	39.01	47.97	59.78	39.19
TiO ₂	0.50	b.d.	b.d.	0.35	b.d.	b.d.	0.42	.p.d	b.d.	0.41	b.d.	b.d.	1.93	.p.d	b.d.	2.10	b.d.	b.d.	0.23	b.d.	.p.d
Cr_2O_3	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	b.d.	0.12	b.d.	b.d.							
Al ₂ O ₃	11.93	25.36	21.40	12.47	28.11	21.41	16.34	24.84	21.13	18.41	25.34	21.76	9.61	25.48	22.04	11.60	25.27	21.90	9.84	25.09	21.96
FeO	11.49	0.69	24.30	12.03	0.61	25.65	13.78	0.29	25.67	11.42	0.68	24.03	9.89	0.26	22.76	10.50	0.50	22.29	8.03	0.59	20.45
MnO	0.14	b.d.	1.28	0.11	b.d.	1.97	0.13	b.d.	0.89	b.d.	b.d.	0.69	0.09	b.d.	1.02	0.12	b.d.	0.85	b.d.	b.d.	b.d.
MgO	13.04	b.d.	6.15	13.16	b.d.	4.75	10.82	b.d.	5.68	11.44	b.d.	5.40	14.96	b.d.	8.41	13.95	b.d.	8.38	16.94	b.d.	8.86
CaO	11.59	7.99	7.61	11.55	10.07	7.38	10.94	6.89	7.61	11.73	7.54	9.71	11.38	7.12	6.51	11.36	6.84	6.94	11.69	6.81	8.21
Na,O	1.30	6.80	.p.d	1.41	5.93	b.d.	2.38	7.50	b.d.	2.46	7.23	p.d.	1.87	7.46	b.d.	2.19	7.81	b.d.	1.62	7.61	.p.d
K,0	0.66	0.23	b.d.	0.62	0.01	b.d.	0.66	0.08	b.d.	0.76	b.d.	b.d.	0.58	0.09	b.d.	0.76	0.10	b.d.	0.23	0.23	b.d.
otal	95.45	98.15	90.06	96.79	100.41	99.58	96.27	99.10	98.54	96.14	98.86	100.32	96.82	99.8	100.2	97.38	100.6	99.37	96.55	100.11	99.33
cat/O	13/23	5/8	8/12	13/23	5/8	8/12	13/23	5/8	8/12	13/23	5/8	8/12	13/23	5/8	8/12	13/23	5/8	8/12	13/23	5/8	8/12
Si	6.59	2.60	3.00	6.53	2.49	3.02	6.02	2.68	2.97	5.83	2.63	2.98	6.72	2.65	3.01	6.49	2.66	3.00	6.80	2.66	2.99
Ľ	0.06	0.00	0.00	0.04	0.00	0.00	0.05	0.00	0.00	0.05	0.00	0.00	0.21	0.00	0.00	0.23	0.00	0.00	0.02	0.00	0.00
Cr	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.01	0.00	0.00	0.00	0.00	0.00	0.00	p.d	0.00	0.00
41	2.07	1.36	1.97	2.13	1.48	1.98	2.84	1.32	1.97	3.20	1.35	1.98	1.64	1.34	1.98	1.98	1.32	1.98	1.64	1.32	1.98
⁴ e ²⁺	1.37	0.03	1.56	1.40	0.02	1.69	1.64	0.01	1.59	1.37	0.03	1.51	1.17	0.01	1.45	1.25	0.02	1.41	0.89	0.02	1.27
Fe^{3+}	0.06	0.00	0.03	0.07	0.00	0.13	0.09	0.00	0.10	0.06	0.00	0.04	0.04	0.00	0.00	0.03	0.00	0.02	0.08	0.00	0.04
Mn	0.02	0.00	0.08	0.01	0.00	00.00	0.02	0.00	0.06	0.00	0.00	0.05	0.01	0.00	0.07	0.01	0.00	0.06	0.00	0.00	0.04
Mg	2.86	0.00	0.72	2.84	0.00	0.56	2.38	0.00	0.67	2.52	0.00	0.62	3.22	00.0	0.96	3.01	0.00	0.96	3.58	0.00	1.01
Ca	1.83	0.39	0.64	1.79	0.48	0.62	1.73	0.33	0.64	1.85	0.37	0.80	1.76	0.34	0.53	1.76	0.32	0.57	1.78	0.33	0.67
Na	0.37	0.60	0.00	0.40	0.51	0.00	0.68	0.65	0.00	0.70	0.63	0.00	0.52	0.65	0.00	0.62	0.67	0.00	0.45	0.66	0.00
×	0.12	0.01	0.00	0.11	0.00	0.00	0.12	0.00	0.00	0.14	0.00	0.00	0.11	0.01	0.00	0.14	0.01	0.00	0.04	0.01	0.00
X.	0.67		0.31	0.66		0.25	0.58		0.28	0.64	0.37	0.29	0.73		0.40	0.70		0.40	0.79	0.33	0.44

to 300 µm wide garnet corona with inclusions of plagioclase, amphibole, and kyanite. Garnet in coronas has a composition of Alm₃₉₋₄₈Prp₂₉₋₃₄Grs₁₈₋₂₆Sps₁₋₂ (Fig. 3d). Garnet coronas show asymmetric zoning with increasing Prp and decreasing Grs proportions from core to rim. The Alm and Sps contents continuously decrease and $X_{\mbox{\tiny Mg}}$ increases across the corona in the direction from orthopyroxene to plagioclase $(Alm_{48 \rightarrow 41 \rightarrow 40} Prp_{31 \rightarrow 29 \rightarrow 35} Grs_{18 \rightarrow 26 \rightarrow 22} Sps_{2 \rightarrow 1 \rightarrow 1}; X_{Mg} = 0.41 \rightarrow 0.40 \rightarrow 0.46).$ A thin zone of plagioclase occurs between the orthopyroxene-plagioclase symplectite and the garnet corona and it shows compositional zoning marked by the change from oligoclase (An_{25}) to labradorite (An_{68}) in the direction from symplectite to the garnet zone (Fig. 6d). Plagioclase in the whole sample is completely recrystallized into fine-grained clusters of crystals showing strong compositional zoning (Fig. 6d) characterized by the increasing anorthite content from core (An_{35-46}) to the rim (An₇₉₋₉₄; Fig. 3a, Tab. 3). Small grains of spinel, corundum, and kyanite occur within the recrystallized plagioclase. The crystals of plagioclase show a weak shapepreferred orientation defining the fabric and suggesting that the recrystallization was associated with deformation. The other primary minerals (orthopyroxene, clinopyroxene, and ilmenite) do not show such features nor were they observed in any of the other studied samples.

5. P-T conditions

Metamorphic conditions associated with formation of the corona textures have been constrained using the metamorphic assemblages in the selected samples.

Tab. 4 Representative analyses of M, minerals used for P-T calculations

Only three samples were used for geobarometry, namely those bearing garnet-bearing coronas (PJ5 from TCC, PJ6 from MLC near the contact with TCC, and PJ9 from MLC). The reason is that they contain the plagioclase-amphibole-garnet-kyanite assemblage suitable for both multi-equilibrium and conventional geothermobarometry.

For the characterization of metamorphic P-T conditions in the studied coronitic metagabbros (and coronitic rocks in general), the primary (magmatic) mineral assemblage is considered as metastable with respect to corona formation and thus it has to be avoided in the calculations. Moreover, corona formation is typically associated with chemical potential gradients of the components present and therefore even the formation of two adjacent layers of corona sequence cannot be considered to represent a thermodynamic equilibrium. Considering this fact, the minerals used for P-T calculations were carefully selected so that they come exclusively from the garnet corona and occur in a close proximity to each other. Thus the chemical composition of garnet in the vicinity of amphibole, plagioclase and kyanite inclusions was used in the calculations (Tab. 4).

5.1. Multiequilibrium geothermobarometry

For the purpose of the average P–T calculations, activities of mineral end-members in phases considered for P–T estimates were calculated using the AX software (accessible at *http://www.esc.cam.ac.uk/research/researchgroups/research-projects/tim-hollands-software-pages/ ax*). Calculations were carried out equally for the mineral assemblage garnet–amphibole–plagioclase–kyanite in all the selected samples. First set of calculations, done with H₂O in excess, gave the following P–T conditions (Tab. 5, all errors are 2σ): sample PJ5 – 720±172 °C and 12.3±4.6 kbar; sample PJ6 – 765±100 °C and 14.5±2.8 kbar; sample PJ9 – 815±112 °C and 16.3±3.6 kbar.

However, the aH_2O values are expected to be generally low in the coronitic rocks; e.g., St-Onge and Ijewliw (1996) estimated water activities in similar coronitic metagabbros from Canada to values as low as 0.3–0.1. Indeed, the observed textures, represented by coronas and symplectites, indicate that the studied metamorphism occurred under H₂O-undersaturated conditions. In order to deal with this problem, calculations with H₂O-absent equilibria were attempted, giving the following results (Tab. 5): sample PJ5 $- 556 \pm 158$ °C and 9.1 ± 3.4 kbar; sample PJ6 - 679 ± 244 °C and 12.7 ± 5.4 kbar; sample PJ9 - 607 ± 100 °C and 11.7 ± 4.6 kbar. Thus these H2O-absent calculations led to lower mean P-T estimates, especially for samples PJ5 and PJ9, while the conditions calculated for sample PJ6 did not change significantly.

It is clear that for our studied samples the assumed *a*H₂O has large effect on the average P-T calculation results. This means that at least an approximate estimate of the aH₂O values is needed in order to obtain reasonably precise P-T results. Following the approach of Pitra et al. (2010), a series of average P-T calculations with identical mineral assemblage and varying aH₂O was performed. Examining the dependence of calculation errors and the results of the χ^2 test ("fit" value) on aH_2O_2 , one can estimate the correct values of aH₂O. The summary of the average P–T calculations with aH_2O values ranging from 0 to 1 is given in Tab. 6. The lowest uncertainties and the best "fit" values are obtained for very low aH₂O (~0.1) in samples PJ5 and PJ9. The sample PJ6 are also gives the lowest uncertainties for the $aH_2O = 0.1$, however both uncertainties and the "fit" value do not change significantly with increasing aH_2O_1 .

It is important to note that in all cases, the results of average P–T calculations passed the statistical test for the entire range of aH_2O , which means that it is not possible to estimate the aH_2O precisely. However, based on the optimal parameters (uncertainties and the "fit" value), it can be assumed that water activity in the studied rocks was generally rather low. Thus the aH_2O for the final average P–T calculations was set between 0.1 and 0.3. Using the plagioclase–amphibole–garnet– kyanite assemblage, the following range of temperatures and pressures was calculated for the selected samples ($aH_2O = 0.1 \rightarrow 0.3$, uncertainties given as 2σ ; Fig. 8): PJ5:



Fig. 8 P–T diagram showing correlation of the results obtained using conventional and multiequilibrium (average P–T calculations) geothermobarometry. The results of the average P–T calculations are plotted with 1σ uncertainties and were calculated assuming aH_2O of 0.1 (solid-line ellipse) and 0.3 (dashed ellipse).

Tab. 5 Results of multi-equilibrium thermobarometry calculations using average P-T calculations in THER-MOCALC. Conditions were estimated for the samples PJ5, PJ6 and PJ9 for H_0 -absent ($aH_0 = 0$) and H_0 present ($aH_2O = 0.1-1$) conditions

					Sam	ole P.I5				
Phase:		Amp	hibole		Plag	ioclase		Garnet		Kyanite
Endmember:	tr	fact	ts	parg	an	ab	ру	gr	alm	ky
Activity	0.240	0.000	0.004	0.018	0.710	0.570	0.017	0.017	0.150	1.000
sd(a)/a	0.156	0.996	0.700	0.482	0.050	0.055	0.498	0.497	0.198	
$aH_{2}O = 1.0$						H ₂ O abso	ent			
excluded end	member	:: spessa	rtine			excluded	l endmem	ber: spes	sartine	
Independent	set of re	actions				Independ	lent set o	f reaction	S	
1) 3tr + 8gr +	- 24ky =	= 3ts + 2	4an + 2p	у		1) 3tr + 8	8gr + 24k	y = 3ts +	24an + 2p	у
2) 5ts + 20an	= 3tr +	8gr + 2	2ky + 2I	H,O		2) 9tr + 6	6fact + 40)gr + 120	ky = 15ts	+ 120an + 10alm
3) 3fact + 8g	r + 27ky	/ = 30an	+ 5alm	+ 3H,O		3) 9ts + 6	6ab + 4py	y + 8 gr =	3tr + 6par	g + 24an
4) 9ts + 6ab -	+ 4py +	8gr = 3t	r + 6par	g + 24aı	1			-	-	

For 95% con	nfidence,	fit < 1.7	3			For 95%	confiden	ce, fit < 1	.96		
T[°C]	sd (2σ)	P [kbar]	$sd \; (2\sigma)$	cor	fit	T[°C]	$sd~(2\sigma)$	P [kbar]	$\text{sd} \ (2\sigma)$	cor	fit
720	172	12.3	4.6	0.893	1.61	556	158	9.1	3.4	0.943	1.02

					Sam	ple PJ6				
Phase:		Ampl	hibole		Plag	ioclase		Garnet		Kyanite
Endmember:	tr	fact	ts	parg	an	ab	ру	gr	alm	ky
Activity	0.091	0.000	0.001	0.056	0.590	0.650	0.029	0.036	0.100	1.000
sd(a)/a	0.293	1.119	3.900	0.316	0.050	0.050	0.432	0.404	0.258	
$aH_{2}O = 1.0$						H ₂ O abso	ent			
excluded end	member	: spessa	rtine			excluded	l endmem	ber: spes	sartine	
Independent	set of re	actions				Independ	dent set o	f reaction	S	
1) 3tr + 8gr +	- 24ky =	= 3ts + 2	4an + 2p	у		1) 3tr + 3	8gr + 24k	xy = 3ts +	24an + 2p	у
2) 3tr + 8gr +	- 27ky =	= 30an +	5py + 3	H ₂ O		2) 3fact	+ 5py = 3	str + 5alm	1	
3) 3fact + 8g	r + 27ky	/ = 30an	+ 5alm	+ 3H,0		3) 9ts +	6ab + 4py	y + 8gr =	3tr + 6parg	g + 24an
4) 9ts + 6ab -	+ 4py +	8gr = 3t	r + 6par	g + 24aı	1					

4) 9ts + 6ab + 4py + 8gr = 3tr + 6parg + 24an

For 95% con	fidence,	fit < 1.7	3			For 95%	confiden	ce, fit < 1	.96		
T[°C]	$sd\;(2\sigma)$	P [kbar]	$\text{sd}\;(2\sigma)$	cor	fit	T[°C]	$sd \; (2\sigma)$	P [kbar]	$sd\;(2\sigma)$	cor	fit
765	50	14.5	1.4	0.901	0.65	679	122	12.7	2.7	0.978	0.6
					Samp	le PJ9					
Phase:		Ampl	nibole		Plagi	oclase		Garnet		Kyaı	nite
Endmember:	tr	fact	ts	parg	an	ab	ру	gr	alm	ky	/
Activity	0.558	0.000	0.002	0.067	0.530	0.670	0.085	0.034	0.057	1.0	00
sd(a)/a	0.125	1.216	1.700	0.290	0.066	0.050	0.283	0.411	0.341	0.0	00
$aH_{2}O = 1.0$						H ₂ O abso	ent				
excluded end	member	: spessa	rtine			excluded endmember: spessartine					
Independent	set of re	actions				Independ	lent set o	f reactions	5		
1) 3tr + 8gr +	- 24ky =	3ts + 24	4an + 2p	у		1) 3tr + 3	8gr + 24k	y = 3ts +	24an + 2	ру	
2) 5ts + 20an	= 3tr +	8gr + 22	2ky + 2E	I_0		2) 3tr + 3	5alm = 31	fact + 5py			
3) 3fact + 8g	r + 27ky	- 30an	+ 5alm ·	+ 3H,O		3) 9ts +	6ab + 4py	y + 8gr = 1	3tr + 6pa	rg + 24an	

For 95% cor	fidence,	fit < 1.73	3			For 95%	confiden	ce, fit < 1	.96		
T[°C]	$sd~(2\sigma)$	P [kbar]	sd (2\sigma)	cor	fit	T[°C]	$sd\;(2\sigma)$	P [kbar]	$sd\;(2\sigma)$	cor	fit
815	56	16.3	1.8	0.873	1.42	607	100	11.7	2.3	0.972	0.96

 aH_2O – water activity; cor – correlation coefficient; fit – result of the χ^2 test end member abbreviations: tr - tremolite; fact - Fe-actinolite; ts - tschermackite; parg - pargasite an - anortite; ab - albite; py - pyrope; gr - grossular; alm - almandine; ky - kyanite

 $T = 578 \pm 78 \rightarrow 640 \pm 96 \,^{\circ}C$ $P = 9.6 \pm 2.2 \rightarrow 10.8 \pm 2.6$ k b a r, P J 6: T = $616 \pm 72 \rightarrow 681 \pm 84 \ ^{\circ}C.$ $P = 11.4 \pm 2.0 \rightarrow 12.7 \pm 2.4$ kbar and PJ9: $T = 659 \pm 58 \rightarrow 727 \pm 70 \,^{\circ}C$ $P = 12.9 \pm 2.0 \rightarrow 14.4 \pm 2.4$ kbar.

5.2. Conventional geothermobarometry

Using the methods of conventional thermobarometry (Kohn and Spear 1990; Ravna 2000; Tab. 7; Fig. 8), the results for the sample PJ5 from the TCC indicate P-T conditions of 585 °C and 9.4 kbar. Two sets of analyses from the MLC sample PJ6 yielded higher metamorphic conditions of 634 °C at 12.7 kbar and 711 °C at 12.9 kbar, respectively. Similarly, the sample PJ9 from the MLC gave 671 °C at 13.6 kbar and 681 °C at 12.7 kbar. Although the methods of conventional geothermobarometry do not allow determining uncertainties, which can be generally very large (e.g. Powell and Holland 2008), the comparison of the results with those obtained by the multiequilibrium geothermobarometry shows a generally good match of both approaches.

Considering the estimated P-T conditions, it is demonstrated that the corona formation-related metamorphism near the boundary between TCC and MLC reached the upper amphibolitefacies conditions. The lowest P-T conditions were estimated for sample PJ5 from the TCC (Fig. 8). Taking into account the spatial distribution of the selected samples, the calculated P-T conditions imply a rather continuous increase in metamorphic grade towards the structural footwall, i.e. from the east-southeast to the westnorthwest.

6. Whole-rock geochemistry

Based on the C.I.P.W. norm, all analysed samples are hypersthene-normative and, except for the sample PJ7 from the TCC-MLC boundary containing below 1 % of normative quartz, also olivine-normative. The normative plagioclase corresponds to labradorite (An_{52-57}). Sample PJ7 is a gabbro-norite, while all the other samples correspond to olivine gabbro-norite. Samples PJ8 and VP46 are

noticeably rich in normative olivine (~ 25 and 36 % respectively) suggesting that they may represent cumulates.

Since all of the studied samples have undergone some degree of metamorphic changes, which may be connected with an influx or depletion of mobile elements with respect to magmatic protolith, diagrams used here for the classification and characterization of the samples are primarily based on immobile element contents. The TAS proxy diagram Nb/Y vs. Zr/ Ti (Fig. 9a; Pearce 1996, after Floyd and Winchester 1975) shows that the studied samples correspond to subalkali basalts/basaltic andesites except for the ol-

ivine-bearing sample VP46, which falls into the alkali basalt field. Similar signatures can be deduced from the ternary plot relating the cation percentages of Al_2O_3 , FeO + Fe₂O₃ + TiO₂, and MgO (Fig 9b; Jensen 1976), where four of the studied samples correspond mostly to high-Mg tholeiitic basalts. Significant enrichment in Mg content

Tab. 6 Results of P–T calculations using multi-equilibrium thermobarometry for range of $\rm H_2O$ activities from 0 to 1

Sample	aH ₂ O	avT [°C]	sd (2σ)	avP [kbar]	sd (20)	cor	fit
PJ5	For 95% c	onfidence, fit <	1.96				
	0	556	158	9.1	3.4	0.943	1.02
	For 95% c	onfidence, fit <	1.73				
	0.1	578	78	9.6	2.2	0.867	0.75
	0.3	640	96	10.8	2.6	0.880	1.07
	0.5	672	124	11.4	3.4	0.885	1.29
	0.7	695	146	11.9	4.0	0.889	1.44
	0.9	712	164	12.2	4.4	0.892	1.56
	1.0	720	172	12.3	4.6	0.893	1.61
PJ6	For 95% c	onfidence, fit <	1.96				
	0	679	244	12.7	5.4	0.978	0.60
	For 95% c	onfidence, fit <	1.73				
	0.1	616	72	11.4	2.0	0.875	0.60
	0.3	681	84	12.7	2.4	0.888	0.43
	0.5	715	90	13.4	2.4	0.893	0.48
	0.7	739	96	13.9	2.6	0.897	0.55
	0.9	757	98	14.3	2.6	0.900	0.62
	1.0	765	100	14.5	2.8	0.901	0.65
PJ9	For 95% c	onfidence, fit <	1.96				
	0	607	200	11.7	4.6	0.972	0.96
	For 95% c	onfidence, fit <	1.73				
	0.1	659	58	12.9	2.0	0.846	0.77
	0.3	727	70	14.4	2.4	0.860	1.04
	0.5	763	86	15.1	2.8	0.866	1.19
	0.7	787	98	15.7	3.2	0.870	1.30
	0.9	807	108	16.1	3.6	0.872	1.38
	1.0	815	112	16.3	3.6	0.873	1.42

 aH_2O – water activity; cor – correlation coefficient ; fit – result of the χ^2 test

of the two olivine-bearing samples VP46 and PJ8 is then supporting the interpretation that they represent olivine cumulates.

Binary diagrams based on ratios of selected immobile elements (Nb/Yb, Nb/Th and Ti/Yb, Figs 9c-d, after Pearce 2008) were used in order to define magma types corresponding to the studied gabbro samples and subsequently to assess their likely tectonic setting. In the Nb/Yb vs. Th/Yb plot (Fig. 9c) all the studied samples have signatures close to that of the E-MORB. Since all the samples show no significant increase in the Th/Nb ratio above the 'Mantle Array' (NMORB–OIB line), it can be concluded that they

Tab. 7 P-T conditions estimated using conventional geothermobarometry

sample	T (°C)	geothermometer of Ravna (2000)	P (kbar)	geobarometer of Kohn and Spear (1990)
PJ5	585	Grt–Hbl	9.4	Grt–Hbl–Pl
PJ6	634	Grt–Hbl	12.7	Grt–Hbl–Pl
	711	Grt–Hbl	12.9	Grt–Hbl–Pl
PJ9	671	Grt–Hbl	13.6	Grt–Hbl–Pl
	681	Grt–Hbl	12.7	Grt–Hbl–Pl



Fig. 9 Classification and discriminant diagrams showing the character and tectonic setting of the studied gabbros. \mathbf{a} – The immobile element-based TAS proxy diagram (Pearce 1996, after Floyd and Winchester 1975); \mathbf{b} – Ternary plot relating the cation percentages of Al₂O₃, FeO + Fe₂O₃ + TiO₂, and MgO (Jensen 1976); \mathbf{c} – Binary diagram Nb/Yb vs. Th/Yb (Pearce 2008, 2014). The 'MORB–OIB array' is formed by average N-MORB, E-MORB and OIB compositions taken from Sun and McDonough (1989). The Th/Nb proxy is used to distinguish suprasubduction zone from midocean ridge basalts; \mathbf{d} – The Nb/Yb–TiO₂/Yb diagram to discriminate between the deep and shallow melting leading to formation of N-MORB, E-MORB, and OIB (Pearce 2008, 2014).

were generally neither affected by subduction-derived fluid/melt nor significantly contaminated by a mature crustal material. The Ti/Yb ratios of the samples (Fig. 9d) are low, typical of shallow (spinel) mantle melting and correspond to E-MORB, except the olivine cumulate VP46.

The E-MORB character of the studied gabbros is also supported by multielement plots (Fig. 10). Contents of presumably immobile trace elements normalized to N-MORB (Fig. 10a, after Pearce 2014) resemble typical E-MORB (Sun and McDonough 1989) except again for the olivine cumulate VP46 that is depleted in all of the elements compared to the other samples. Similarly in the REE spider plots normalized to chondrite composition (Fig. 10b, after Boynton 1984), the patterns correspond to the E-MORB. Positive Eu-anomaly in the sample PJ7 indicates a role of plagioclase accumulation during its formation.

7. Discussion

7.1. Geochemical characteristics of the studied metagabbro in the context of previous studies

The geochemical characteristics of the studied metagabbros from the TCC-MLC boundary show that their composition generally corresponds to tholeiitic basalts, which is in agreement with previous studies (Beard et al. 1995; Štědrá et al. 2002; Timmermann et al. 2004). Their relatively immobile trace-element signatures display a clear affinity to E-MORB with no significant contribution of subduction-derived fluid/melt or continental crust material and with composition typical of shallow mantle melting. Such signatures indicate that the formation of the gabbros was not connected to a subduction-related environment but it was rather associated to rifting of the crust. This is in agreement with interpretations that the gabbros were intruded into an already established crustal stack involving the metasedimentary TCC and eclogitebearing MLC as proposed by Štědrá et al. (2002). This intrusion may have been then related to a formation of a rift in the Teplá–Barrandian Unit that was presumably triggered by the onset of the Iapetus Ocean subduction at \sim 510 Ma as proposed by Žák et al. (2013).

7.2. Metamorphic characteristics of the studied metagabbros in the context of previous studies

The contact of the TCC and MLC represents an important lithological and tectonic boundary in the northwestern Bohemian Massif. However, the character and significance of this boundary remains unclear since the two neighboring but mechanically contrasting complexes have undergone polyphase and rather complicated history over a wide time span from ~550 to ~370 Ma. On the other hand, the studied metagabbros occurring in both complexes show formation of comparable static corona textures associated with a single metamorphic event and implying the absence of deformation overprint. Therefore the corona formation in these rocks represents an important process that allows linking and correlating at least part of the tectono-metamorphic history across the boundary between TCC and MLC.

Metamorphic conditions calculated for the sample PJ5 from the TCC indicate 578-640 °C and 9.1-10.8 kbar. This corresponds well to the P–T conditions estimated for the surrounding metapelites in kyanite-zone at 550-645 °C and 5-9 kbar (Cháb a Žáček 1994). The higher P–T conditions were calculated for the two MLC metagabbros (616-715 °C at 11.4-12.9 kbar for PJ6 and 659-727 °C at 12.7-14.4 kbar for PJ9). These estimates

are in a good agreement with the previous data on the metamorphic overprint of the MLC gabbros (600–730 °C, 8–11 kbar: Štědrá 2001; 585–614 °C, 12–13 kbar: Faryad 2012).

In conclusion, the P–T conditions calculated for the studied metagabbro samples correspond well to the peak metamorphic conditions and metamorphic field gradient recognized in the surrounding rocks, indicating an increase in P–T conditions towards the northwest (Cháb and Žáček 1994; Žáček 1994; Cháb et al. 1997; Zulauf



Fig. 10 Spider plots showing trace-element signatures in the studied samples; average E-MORB composition (Sun and McDonough 1989) is plotted for comparison. \mathbf{a} – Relatively immobile trace elements normalized to N-MORB (after Pearce 2014, normalizing values from Sun and McDonough 1989), \mathbf{b} – REE normalized by average chondrite composition (Boynton 1984).

Tab. 8 Whole-rock chemical compositions of studied metagabbros

major elements (oxides in wt. %)						
sample	PJ5	PJ6	PJ7	PJ8	PJ9	VP46
SiO ₂	49.06	47.93	51.34	45.86	47.92	43.67
TiO,	1.26	1.82	0.75	1.39	1.21	1.03
Al ₂ O ₃	15.62	16.51	17.14	11.17	17.97	7.01
Fe ₂ O ₃	9.40	11.52	7.55	14.31	8.57	16.62
Cr,0,	0.077	0.021	0.030	0.156	0.049	0.171
MnO	0.17	0.19	0.17	0.20	0.14	0.19
MgO	8.49	8.08	7.88	17.30	8.99	22.08
CaO	11.61	10.01	10.71	6.39	10.52	5.28
Na ₂ O	2.80	2.83	2.84	1.92	2.87	1.26
K,Ō	0.23	0.52	0.21	0.53	0.47	0.39
P,O,	0.15	0.19	0.07	0.20	0.15	0.12
total	98.87	99.62	98.69	99.43	98.86	97.82
trace elements (ppm)						
Ва	1	158	68	135	121	54
Co	38.1	41.9	37.0	84.4	40.8	119.6
Cs	2.3	1.2	0.7	1.2	0.8	1.1
Ga	15.7	17.9	16.4	13.4	15.3	10.5
Hf	4.5	3.7	1.0	3.8	2.9	1.7
Nb	7.8	9.2	2.4	8.5	7.3	8.0
Rb	4.8	12.4	4.5	13.0	9.8	5.2
Sr	192.2	242.5	264.6	177.8	305.5	166.2
Та	0.6	0.6	0.2	0.9	0.5	0.7
Th	0.9	1.2	0.3	1.3	1.1	0.5
U	0.2	0.4	0.1	0.7	0.5	0.1
V	243	246	265	161	177	152
Zr	159.5	139.2	39.4	139.3	100.6	62.4
Y	32.3	28.7	10.8	19.9	18.9	8.9
La	9.3	10.8	4.4	10.8	10.0	6.6
Ce	24.3	25.6	9.0	25.9	22.3	13.7
Pr	3.12	3.34	1.14	3.39	2.82	1.89
Nd	15.8	15.2	5.6	15.7	11.7	8.5
Sm	4.33	4.03	1.16	3.88	2.87	2.00
Eu	1.29	1.46	0.91	1.19	1.13	0.75
Gd	5.13	5.17	1.96	4.04	3.50	2.34
Tb	0.95	0.86	0.35	0.70	0.56	0.38
Dy	5.74	5.50	2.00	4.29	3.59	2.02
Но	1.07	1.08	0.38	0.82	0.76	0.38
Er	3.49	3.10	1.19	2.35	2.01	1.06
Tm	0.51	0.48	0.15	0.33	0.28	0.13
Yb	3.43	2.65	1.11	2.20	1.88	0.89
Lu	0.48	0.43	0.17	0.34	0.25	0.15
Мо	0.5	0.6	< 0.1	0.8	0.4	0.8
Cu	69.0	24.1	20.6	128.0	33.0	120.8
Pb	0.5	1.0	0.5	1.0	1.9	1.0
Zn	8	16	7	18	13	47
Ni	28.0	31.0	21.4	475.3	71.1	506.5
As	<0.5	1.0	0.6	2.5	< 0.5	< 0.5
Au	3.6	1.2	0.7	2.9	0.6	0.6

2001; Peřestý 2012). The peak P–T conditions obtained from the TCC metapelites have been recently ascribed to the early-Variscan horizontal east–west shortening and crustal thickening (Peřestý 2012). It is inferred that the corresponding P–T conditions obtained from the studied metagabbro were very likely associated with the same deformation phase. The subsequent exhumation, manifested by an intense tectonometamorphic overprint of surrounding rocks (Peřestý 2012), is reflected only by a weak retrogression identified in the studied TCC metagabbro PJ5.

The above-described metamorphic field gradient documented from metagabbro samples and surrounding rocks shows rather continuous increase in P–T conditions across the contact between TCC and MLC. Such a lack of obvious gap in the Variscan metamorphic conditions supports the interpretation that the thickening event recognized in the TCC rocks was also responsible for the metamorphism of the studied metagabbros. This conclusion, together with the fact that both complexes were intruded by the Late Cambrian gabbros of similar geochemical signature (496–503 Ma: Bowes and Aftalion 1991; Timmermann et al. 2004), strongly suggests that both major lithological complexes were brought together prior to the gabbro intrusion during the Late Cambrian rifting.

It is important to point out that the metagabbros within the eclogite-bearing MLC reveal maximum pressure of ~14 kbar, which is generally lower compared to estimates for the eclogites (16-28 kbar, Jelínek et al. 1997; Štědrá 2001; Faryad 2012). This led Štědrá et al. (2002) and Timmermann et al. (2004) to suggest that the eclogitefacies metamorphism in the MLC may have been pre-Variscan, presumably older than ~500 Ma as constrained by the age of the gabbro intrusions. On the other hand, this study revealed a pressure increase from ~10 kbar (sample PJ5) to ~14 kbar (sample PJ9) from the TCC to the eastern part of the MLC over a horizontal distance of ~ 15 km. It is therefore possible that the eclogitefacies conditions with pressures of ~18-20 kbar could have been achieved in the western part of the MLC in the continuity of the presumably Variscan metamorphic field gradient. This argumentation was used by Faryad (2012) to support the Variscan age of the eclogite-facies metamorphism in the MLC.

However, the presence of eclogites is not limited to the northwestern MLC and several such bodies crop out directly at the contact between MLC and TCC. These eclogite occurrences are located ~ 5 km east of the metagabbro sample PJ6 that did not reach pressures exceeding 13 kbar. Thus the occurrence of eclogites in the close proximity to metagabbros violates the otherwise continuous metamorphic field gradient and bids for a polymetamorphic character of the mafic rocks in the MLC. This stands in sharp contrast to the single metamorphic event recorded in the metagabbro. Indeed, the eclogites are overprinted by amphibolite-facies metamorphic assemblage. Based on the above arguments, it seems possible that eclogites were first intruded by gabbro and only then retrogressed under the amphibolite-facies conditions during the Variscan event. Because of the Late Cambrian intrusive age of the gabbro this in turn implies a pre-Variscan age for at least some of the MLC eclogites.

7.3. Interpretation of the metamorphic textures

The characteristic feature of the studied samples is a large variability of corona sequences developed at the contacts between primary minerals of the M₁ assemblage, namely between plagioclase and the mafic phases, olivine, orthopyroxene, clinopyroxene, amphibole, biotite, and ilmenite. While the magmatic origin of the M₁ mineral assemblage is rather convincing, the origin of the M₂ mineral assemblage needs to be discussed, since there is no general agreement on the process of corona formation. In different studies, the studied coronas were explained to result from 1) progressive reactions during magmatic crystallization (e.g. Joesten 1986), 2) subsolidus reactions during slow cooling and post-magmatic evolution (e.g. Whitney and McLelland 1973), or 3) post-magmatic metamorphism of the gabbroic rocks (e.g. Spear and Markussen 1997).

Corona sequences similar to those described in this study were already reported from several localities. Haas et al. (2002) described coronas formed between olivine and plagioclase in a gabbro from Norway and concluded that these textures were formed during a multi-stage, late-magmatic process. A comparable corona sequence around olivine (orthopyroxene \rightarrow amphibole $\rightarrow \pm$ clinopyroxene–spinel symplectite) has been also observed in a troctolitic gabbro from Argentina (Gallien et al. 2012) where it was interpreted as being formed in several stages during cooling of the gabbro intrusion at the transition to granulite.

On the other hand, similar coronas were studied in a gabbro-norite from the Baltic Shield by Larikova (2000) and their formation was interpreted as a result of a prograde metamorphism of the magmatic rocks. This author has concluded that all the layers of the observed corona sequences had grown simultaneously by the mechanism of diffusion metasomatism. Simultaneous growth of multi-layer coronas by the metasomatism between olivine/orthopyroxene and plagioclase was later simulated experimentally by Larikova and Zaraisky (2009).

The formation of coronitic textures in metagabbros from the MLC and TCC could reflect a prograde metamorphism (e.g. Štědrá et al. 2002; Timmermann et al. 2004), even though some of the phases present in the coronas, i.e. Ti-rich amphibole and orthopyroxene, were interpreted as late magmatic (e.g. Štědrá et al. 2002 and references therein). The coronas described in this work do not show a polyphase metamorphic overprint except for sample PJ5, where the growth of chlorite, albite, muscovite, and prehnite can be attributed to a later retrogression. The gabbro bodies intruded into relatively shallow depths as indicated by the occurrence of hornblende and pyroxene hornfelses in thermal aureoles around some of the gabbro bodies (Kachlík 1997; Štědrá et al. 2002). This contrasts with the relatively higher pressures estimated from coronas in the current study. At the same time, metamorphic P-T conditions of the gabbros show very good correlation with the record of Variscan metamorphism in the surrounding rocks. Based on these arguments, it can be concluded that the observed coronas, consisting of the M₂ metamorphic mineral assemblage, were formed by prograde metamorphism superimposed on the significantly older (~500 Ma) magmatic mineral assemblage M₁ during the regional metamorphic event of Variscan age.

An interesting feature observed in all studied samples is the occurrence of tiny zircon rims developed around magmatic M₁ ilmenite. Since zircon appears in such specific microstructural position, it can provide a valuable age constraints for either metamorphic or magmatic age related to M₂ or M₁ assemblage, respectively. Similar zircons have been previously reported by several authors, however their genetic interpretations did diverge. The metamorphic origin of such zircons was explained either by a direct crystallization (Sláma et al. 2007) or by exsolution of baddeleyite from magmatic ilmenite and its subsequent transformation to zircon during granulitefacies conditions (Bingen et al. 2001). The magmatic origin of the zircons was proposed by Morriset and Scoates (2008), who observed zircons around ilmenites in unmetamorphosed anorthosites from Quebec. Austrheim et al. (2008) concluded that the zircon rim around ilmenite in amphibolites and gabbros could be of magmatic origin or developed during subsequent metasomatism.

It is worth noting that the mineral assemblage enveloping ilmenite in our studied samples changes with the metamorphic grade. In the sample from the TCC, characterized by the lowest P–T conditions, zircon around ilmenite is associated with titanite (Fig. 2b). The intermediate P–T conditions recorded by the MLC and TCC–MLC boundary samples show ilmenite being rimmed only by zircon while the highest P–T conditions recorded in sample PJ9 show ilmenite partially replaced by rutile, which is surrounded by the zircon grains (Fig. 7d). Microstructural relations in the studied samples do not provide clear evidence for magmatic or metamorphic origin of the zircon rims. A simple fact that zircon occurs as rims around magmatic ilmenite in coronitic rocks may suggest that the zircon itself represents a corona formed

during metamorphism. On the other hand, the zircon grains present around ilmenite that was partially replaced by rutile in sample PJ9 may imply that zircon rim was formed before the metamorphism.

Almost all of the studied samples show an extensive recrystallization of magmatic plagioclase associated with formation of a two-plagioclase mixture. Grove et al. (1983) interpreted this microstructure with coexisting oligoclase/andesine and bytownite in terms of plagioclase immiscibility characteristic of the temperature range of ~400–575 °C, as earlier defined by Spear (1977). In the case of the studied metagabbro samples, this microstructure can be then interpreted as a result of the magmatic plagioclase breakdown during metamorphism at temperatures in which the original (magmatic) composition of ~An₅₀₋₇₀ becomes unstable.

In addition, plagioclase in the studied samples often contains numerous small inclusions or lamellae of Al-rich phases such as spinel and/or corundum. Similar phenomenon often referred to as "clouding" of plagioclase was previously described by Poldervaart and Gilkey (1953) or Goldsmith (1982). The presence of tiny second-phase crystals in plagioclase mixture was interpreted to result from a diffusion-controlled process. The samples studied in this work are characterized by growth of coronas that contain Ca-bearing phases such as amphibole and garnet at the contact of Ca-free olivine and orthopyroxene with plagioclase. Formation of such coronas requires consumption of Ca (and Al) from plagioclase. Therefore, the crystallization of spinel and corundum within plagioclase may be interpreted as an indication of the higher diffusivity of Ca compared to Al out from plagioclase accompanied by Mg and Fe diffusion into plagioclase during the corona formation.

In summary, the process of plagioclase transformation can be divided into three stages: 1) crystallization of spinel and corundum in plagioclase due to diffusion of Ca > Al from plagioclase accompanied with Mg + Fe diffusion into plagioclase, 2) breakdown of magmatic plagioclase (labradorite) to andesine–oligoclase and anorthite–bytownite, and 3) recrystallization of the magmatic plagioclase into the fine-grained clusters of zoned plagioclase grains with Ca increasing towards their rims (e.g. PJ9). While the stages 1 and 2 clearly represented a static process, the final recrystallization stage 3 in sample PJ9 was connected with late deformation–metamorphic overprint.

Most of these features were never described from the metagabbros of the TCC or MLC. On the other hand, some previous works (Zulauf 1997; Štědrá et al. 2002) have reported the presence of kyanite and zoisite grains in the calcic domains of plagioclase that were not observed in the present study. The only exception is the kyanite appearing in the plagioclase domains of the MLC sample PJ9 that yields the highest metamorphic conditions. While it is difficult to link our observations with those reported by other authors, it is possible that the textures seen in our samples represent an initial stage of plagioclase decomposition, which was followed by complete recrystallization connected by formation of zoisite and kyanite during continuing metamorphism and deformation.

8. Conclusions

Petrological and geochemical investigations of the metamorphosed gabbros occurring at the contact of the Teplá-Barrandian Unit (TBU) and the Mariánské Lázně Complex (MLC) brought some constraints on the evolution of this important tectonic boundary. The M₂ mineral assemblage associated with coronas formed around the primary magmatic M₁ minerals as a result of static metamorphic overprint of the gabbro. The closely related minerals in the garnet-bearing coronas were used to estimate metamorphic P-T conditions in several samples across the boundary of the Teplá Crystalline and Mariánské Lázně complexes (TCC and MLC, respectively). The lowest metamorphic conditions of $\sim 600 \pm 50$ °C and 10 ± 1.5 kbar were estimated from the TCC and the highest of $\sim 700 \pm 50$ °C and 13.5 ± 1.5 kbar were obtained from the MLC. The spatial arrangement of our P-T estimates suggests rather continuous increase in metamorphic grade from TCC to MLC, i.e. towards the west-northwest. The observed higher grade metamorphic field gradient across the TCC-MLC boundary corresponds well to the lower grade metamorphic field gradient recorded in the TCC where the peak metamorphic conditions have been associated with crustal thickening during Variscan event. It appears likely that the thickening was also responsible for the metamorphism of the metagabbro in the MLC thus questioning the Variscan age of the nearby eclogites. The whole-rock geochemical analyses show that metagabbros occurring in the TBU have compositions corresponding to tholeiitic basalts with trace-element signatures characteristic of E-MORB, which is consistent with interpretation that the intrusion of these rocks was related to an intracontinental rifting of the TBU during late Cambrian and was not connected to any subduction processes.

Acknowledgements. V. Peřestý and O. Lexa are thanked for their help with sampling and numerous valuable discussions in the field and beyond. We gratefully acknowledge thorough reviews by P. Pitra and V. Štědrá and editorial handling by P. Hasalová. The work was financially supported by the Czech National Grant Agency (13–16315S to P. Štípská and O. Lexa).

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