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Metamorphic and geochronogical study of the Triassic El Oro metamorphic complex, Ecuador: Implications for high-temperature metamorphism in a forearc zone

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ABSTRACT

In the forearc of the Andean active margin in southwest Ecuador, the El Oro metamorphic complex exhibits a well exposed tilted forearc section partially migmatized. We used Raman spectroscopy on carbonaceous matter (RSCM) thermometry and pseudosections coupled with mineralogical and textural studies to constrain the pressure-temperature (P-T) evolution of the El Oro metamorphic complex during Triassic times. Our results show that anatexis of the continental crust occurred by white-mica and biotite dehydration melting along a 10 km thick crustal domain (from 4.5 to 8 kbar) with increasing temperature from 650 to 700 °C. In the biotite dehydration melting zone, temperature was buffered at 750-820 °C in a 5 km thick layer. The estimated average thermal gradient during peak metamorphism is of 30 °C/km within the migmatitic domain can be partitioned into two apparent gradients parts. The upper part from surface to 7 km depth records a 40-45 °C/km gradient. The lower part records a quasi-adiabatic geotherm with a 10 °C/km gradient consistent with an isothermal melting zone. Migmatites U-Th-Pb geochronology yielded zircon and monazite ages of 229.3 ± 2.1 Ma and 224.5 ± 2.3 Ma, respectively. This thermal event generated S-type magmatism (the Marcabeli granitoid) and was immediately followed by underplating of the high-pressure low-temperature (HP-LT) Arenillas–Panupali unit at 225.8 ± 1.8 Ma. The association of high-temperature low-pressure (HT-LP) migmatites with HP-LT unit constitutes a new example of a paired metamorphic belt along the South American margin. We propose that in addition to crustal thinning, underplating of the Piedras gabbroic unit before 230 Ma provided the heat source necessary to foster crustal anatexis. Furthermore, its MORB signature shows that the asthenosphere was involved as the source of the heat anomaly. S-type felsic magmatism is widespread during this time and suggests that a large-scale thermal anomaly affected a large part of the South American margin during the late Triassic. We propose that crustal anatexis is related to an anomaly that arose during subduction of the Panthalassa ocean under the South American margin. Slab verticalization or slab break-off can be invoked as the origin of the upwelling of the asthenosphere.

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1. Introduction

While the generation of *HP-LT* rocks, especially blueschist facies assemblage, is characteristic of a subduction setting (e.g., Ernst, 1988; Guillot et al., 2009), *HT-LP* rocks can be produced in a wide range of geological settings. In modern belts, *HT-LP* rocks and associated crustal anatexis (S-type magmatism) are typically generated in

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collisional orogens (Hodges, 2000). In such a setting, HT conditions are reached either by crustal thickening or by post-orogenic collapse (Gardien et al., 1997). Crustal magma generation occurs on a time-scale of 10 to 30 Ma and can produce large volumes of granites (Thompson and Connolly, 1995). In a subduction context, granitoids are usually of I-type and are not associated with HT-LP metamorphism, especially in a forearc setting (Brown, 2007; Huppert and Sparks, 1988).

In the present forearc region of the Andean margin of southwest Ecuador, the El Oro metamorphic complex (Fig. 1) exhibits a continental sequence composed of metasediments (El Tigre and La Victoria units), partially molten metasediments (La Bocana unit) and intruded



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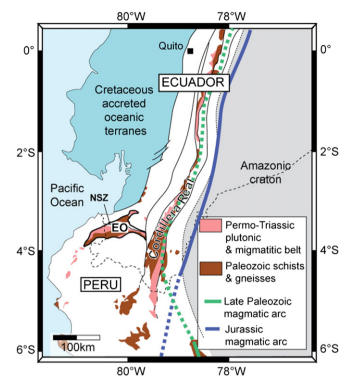


Fig. 1. Simplified geological map of Ecuador and northern Peru modified after Chew et al. (2007) showing the location of the El Oro metamorphic complex. EO, El Oro; NSZ, Naranjo Shear Zone. Continuous line: documented magmatic arc position, dashed line: inferred magmatic arc position.

by S-type granitoids (Marcabeli and El Prado plutonic rocks), juxtaposed with amphibolitic metagabbros (Piedras unit) and blueschist facies metabasalts (Arenillas–Panupalí unit) (Fig. 2). Because of the subvertical structures, this complex has been interpreted either as a shear zone active during the Triassic (Aspden et al., 1995), or as a remnant of a forearc zone tilted during a major tectonic event in the Late Triassic (Gabriele, 2002). However, little is known about the geological structures of the continental sequence and its relationships with the Piedras metagabbroic unit and the Arenillas–Panupalí blueschist unit.

Here, we focus our study on the *HT-LP* Triassic rocks of the El Oro metamorphic complex (Fig. 2). Our aims are to: (1) characterize the deformation of those units; (2) estimate the peak metamorphic P-T conditions; (3) age of migmatization; and (4) propose a model for the *HT-LP* gradient and the crustal anatexis associated with underplating of *HP-LT* rocks. Here we present the results obtained by combining field work, textural observations, mineralogical identification, geothermobarometry and geochronological studies. The analytical methods are presented in the Appendix A.

2. Geological setting and previous studies

In SE Ecuador various metamorphic terranes of both continental and oceanic affinity form the El Oro metamorphic complex (Aspden et al., 1995; Feininger, 1978). This complex is bounded to the south by the Cretaceous volcano-sedimentary sequence of the Celica–Lancones basin (Jaillard et al., 1996, 1999), to the NE by Tertiary volcanic deposits and to the NW by Cenozoic, recent sedimentary deposits (Fig. 2). This complex is characterized by its forearc position and by discordant E–W striking structures, within the NNE striking Andean orogenic belt (Fig. 1). The Late Paleozoic subduction related magmatic arc is known to occur within the Cordillera Central of Colombia (Vinasco et al., 2006), in NW Peru (Mišković, et al., 2009), and a Jurassic magmatic arc is known in Colombia and Ecuador, lying in the Subandean Zone

(Jaillard et al., 2000). Consequently, since at least Carboniferous times, the El Oro metamorphic complex has been located in a forearc position relative to Pacific subduction zone (Fig. 1).

Three tectono-metamorphic groups are distinguished for the El Oro metamorphic complex:

- (1) The Biron Complex, North of the La Palma–Guayabo Shear Zone (Fig. 2), consists of metasediments, migmatitic paragneisses, granitoids, metadiorites and amphibolites with a N-MORB-type geochemical affinity (Gabriele, 2002). Ar–Ar radiometric datings on biotite in migmatites and in metadiorites yielded cooling ages of 75.5 ± 2.3 Ma and 78.4 ± 0.5 Ma, respectively. Pb–Pb dating on monazites yielded ages of 78 ± 1 Ma and 82 ± 1 Ma, and three U–Pb zircon analyses plot on a reverse discordia with a lower intercept at 200 ± 30 Ma (Noble et al., 1997). Noble et al. (1997) interpreted the lower intercept at 200 ± 30 Ma as the age of crystallization of the granitoid, and attributed the younger monazite ages to a later episode of deformation and metamorphism.
- (2) South of the La Palma–Guayabo shear zone (Fig. 2), the Raspas complex (Feininger, 1980) consists of an ophiolitic massif, the El Toro metaperidotitic unit, and the Raspas eclogitic unit (Gabriele et al., 2003; John et al., 2010). The Lu–Hf ages of John et al. (2010) from the Raspas complex indicate that the ophiolite underwent prograde HP metamorphism at around 130 Ma. Radiometric dating yielded an age of 127–123 Ma (Ar/Ar on phengite), which was interpreted as the age of underplating of the Raspas eclogitic complex beneath the Ecuadorian margin (Feininger and Silberman, 1982; Gabriele, 2002; John et al., 2010).
- (3) The El Oro paired metamorphic belt is located between the Raspas complex and the Cretaceous Celica-Lancones Basin (Figs 1 & 2) and consists of late Paleozoic sediments (Martínez, 1970) metamorphosed during the Triassic (Aspden et al., 1995), juxtaposed with the Piedras gabbroic unit (Aspden et al., 1992b) and with the Arenillas-Panupalí blueschist unit metamorphosed in the Triassic (Gabriele, 2002). From south to north (Fig. 2), the continental sequence is composed of the El Tigre and La Victoria low-to-high-grade metapelitic units, intruded by the Marcabeli S-type granitoid at 227 ± 0.5 Ma (U–Pb on zircon, Noble et al., 1997) and of the La Bocana migmatitic unit dated at 219 ± 22 Ma (Sm/Nd whole-rock/garnet isochron; Aspden et al., 1995). In the Piedras metagabbroic unit, U–Pb ages on zircon are 221 ± 17 Ma (Noble et al., 1997) and Ar–Ar ages on amphibole yielded an age of 226 ± 1.8 Ma (Gabriele, 2002). Geochemical studies indicate a MORB-type affinity for the Piedras gabbroic unit (Aspden et al., 1995; Bosch et al., 2002; Gabriele, 2002). Both the Piedras and the Arenillas-Panupalí units were retrogressed under greenschist-facies conditions (Gabriele, 2002).

The El Tigre unit dips approximately to the north (50-60°), and is composed of turbidites (Aspden et al., 1992a). On its southern boundary the El Tigre unit is unconformably overlain by the Cretaceous sediments of the Celica-Lancones basin. In the north, the appearance of chlorite and biotite (Fig. 2) marks a gradational transition zone between the low metamorphic grade El Tigre unit and the moderate- to high-grade La Victoria unit. The La Victoria unit comprises a sequence of metapelites and metapsammites similar to that of the El Tigre unit. To the north, the La Victoria unit is typically composed of biotite \pm muscovite \pm fibrolite, albite and quartz with and alusite porphyroblasts. The Marcabeli S-type granitoid into the La Victoria unit is an elongated E–W trending laccolithic body. This pluton principally consists of medium-grained biotite \pm muscovite granodiorites, with hornblende-rich xenoliths (Aspden et al., 1995). The contact between the La Victoria unit and the La Bocana unit has been interpreted as a syn- to late-magmatic dextral shear zone (Aspden et al., 1995), which is coeval with sillimanite crystallization (Feininger, 1978). These shear zones show a sub-vertical foliation. North of Marcabeli, Feininger (1978) mapped a folded contact between the La Victoria and the La Bocana unit (Fig. 2), with a lens of the La

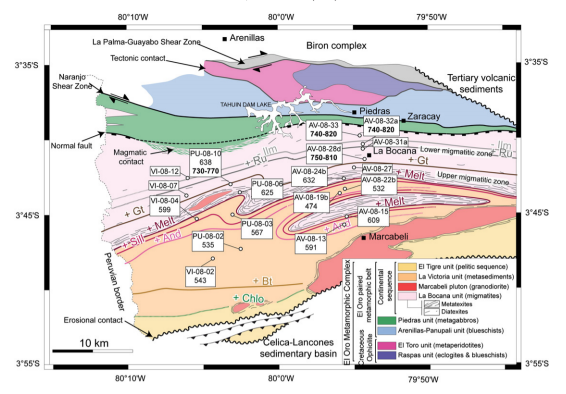


Fig. 2. Geological map of the western part of the El-Oro metamorphic complex of Ecuador modified after Feininger (1978) and Aspden et al. (1995). The map shows the different tectono-metamorphic units. The main contacts, metamorphic isograds, the location of the studied samples are also indicated, and our thermometric results are indicated in °C (regular = RSCM thermometry; Bold = pseudosection thermometry). In unmelted metasediments there is a good correlation between metamorphic isograds and estimated temperature by RSCM; in the migmatitic zone the calibration limit of the RSCM thermometer is reached (640 °C) and temperatures are underestimated in comparison with the wet solidus (~650 °C) and calculated pseudosection temperatures in garnet-bearing migmatites (~750 to 820 °C).

Bocana unit cropping out within the metasediments of the La Victoria unit. Moreover, the metamorphic isograds follow the structure, implying that folding was post-metamorphic. The La Bocana unit (Fig. 2) has been described by Aspden et al. (1995) at its type locality as predominantly composed of garnet-bearing granitoid and lesser amounts of metatexites. North of the La Bocana unit, the contact with the Piedras unit (Fig. 2) is recognized as tectonic (Gabriele, 2002). However, amphibolite xenoliths at the base of the La Bocana unit and granitic bodies within the Piedras unit suggest that the contact was originally intrusive (Aspden et al., 1995). The Piedras dips to the south (70–90°), and is 3 to 5 km thick. This unit is mainly composed of amphibolites retrogressed under greenschist-facies conditions, and shows variable textures, massive to gneissic, fine to coarse grained (Gabriele, 2002). It can be traced from the Peruvian border for about 60 km to the east (Aspden and Litherland, 1992). The contact zone between the Piedras unit and the Arenillas-Panupalí unit is defined by the Naranjo shear zone (Gabriele, 2002). The Arenillas-Panupalí unit roughly strikes E-W and presents a sub-vertical foliation. It forms a 40 km long, 0.5 to 5 km wide belt. This unit is composed of metabasalts and metasediments retrogressed under greenschist-facies, relict blueschist-facies assemblage. The peak P-T conditions of this unit are estimated by Gabriele (2002) at ~9 kbar and 300 °C from garnet + glaucophane + titanite.

The El Oro metamorphic complex was first interpreted as a continental terrane accreted to the Andean margin during Late Jurassic to Early Cretaceous times (Mourier et al., 1988). Aspden et al. (1995) interpreted the El Oro metamorphic complex as a tectonic mélange formed in a Jurassic accretionary prism, preserving in the southern part a HT metamorphosed and migmatized unit of Late Triassic age. On the basis of the presence of garnet and glaucophane, Aspden et al. (1995) considered the Arenillas–Panupalí blueschist unit as part of the Late Jurassic–Early Cretaceous Raspas ophiolitic complex (Aspden and Litherland, 1992; Aspden et al., 1992a). Additionally, Aspden et al. (1995) suggested that the emplacement of the gabbroic Piedras unit into the active shear zone provided the heat source for crustal anatexis and formation of the S-type Marcabeli granitoid in *HT-LP* conditions. More recently, based on the MORB-like geochemical affinity and radiometric ages on amphibole of 225.3 ± 1.7 Ma, Gabriele (2002) described the Arenillas–Panupalí unit as a new tectono-metamorphic blueschist unit located between the Raspas complex to the north, and the Piedras unit to the south.

Based on the geochemical MORB-type affinity and on the radiometric age of 226 ± 1.8 Ma of the Piedras unit, Gabriele (2002) interpreted this unit as having originated during a pre-Jurassic subduction period, subsequently accreted to the continental margin. However, according to Noble et al. (1994), the zircons of the Piedras unit are of magmatic origin, and hence the age of 221 ± 17 Ma is considered to be the crystallization age. Although the Piedras unit shows a MORB-type signature, it is predominantly composed of amphibolites, with relict gabbroic textures, and no pillow basalts or sediments have been observed, thus favoring a plutonic origin. The continental sequence exhibits a northward increase in metamorphic temperature conditions (Feininger, 1978). Gabriele (2002) suggested this increase in thermal conditions from low-grade metasediments (El Tigre and La Victoria units) to high-grade migmatites (La Bocana unit) was related to the thermal gradient of a crustal sequence, and the present-day geometry with sub-vertical foliation resulted from the tilting of the entire sequence during a major tectonic event in the Late Triassic (~220 Ma).

3. Field relationships and migmatite morphology

From south to north, we focused our study on two subparallel cross-sections starting from the Cretaceous contact of the Celica– Lancones basin to the eclogitic unit of Raspas. The location of the studied samples is shown in Fig. 2. We observed the same metamorphic evolution from south to north in the La Victoria unit as Feininger (1978) and Aspden et al. (1995). Kinematic indicators (Fig. 3a, b) largely exhibit dextral transtensive apparent movement. The transition from the La Victoria to the La Bocana units is marked by the appearance of *in situ* melting and numerous metatexites. The southernmost evidence of partial melting in the La Victoria is represented by leucocratic veins emplaced within the unmelted metasediments (Fig. 4e).

Along the eastern cross-section, the migmatitic succession of the La Bocana unit out crops 3 km south of La Bocana village (Fig. 2). This succession can be followed along the Río Piedras with almost

no interruption from low-melt fraction metatexites to the south, to the high-melt fraction mesocratic diatexites to the north (Figs. 2 & 4). Foliation is subvertical or strongly dipping to the south. The migmatitic succession starts with the appearance of small pockets of *in situ* melts that are only generated in fertile layers (Fig. 4, zone 2). As the fraction of melt increases in metatexitic migmatites, the sparse pockets of melts become connected (Fig. 4, zone 3). A psammite–pelite original layering, similar to the La Victoria, is still present and shows complex geometries due to contrasting rheological behavior controlled by the different amounts of melt or melt network. Leucocratic and melanocratic parts within pelitic layers are often folded together and

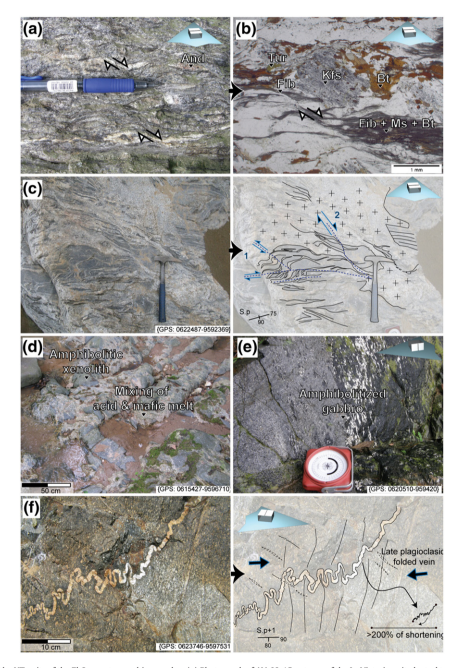


Fig. 3. Photograph of rocks of the HT units of the El Oro metamorphic complex. (a) Photograph of AV-08-15 outcrop of the La Victoria unit, dextral transtensive apparent kinematics indicators are indicated by the black arrows. (b) Photomicrograph of AV-08-15 under polarized light, K-feldspar is recrystallized at the rims by fibrolite \pm white mica \pm biotite, underlining the high temperature foliation with dextral transtensive apparent movement (black arrows); (c) metatexite of the upper migmatitic zone, melt has percolated within resistant melanocratic layers, also showing a apparent dextral transtensive movement; (d) photograph of mixing zone between the Piedras unit and the La Bocana unit, gabbroic melt and acid melt are mixed (see Fig. 5h), numerous mafic xenoliths are present; (e) photograph of the metagabbros of the Piedras unit; (f) photograph of the highly strained contact zone between the Piedras unit and the Arenillas–Panupalí unit, unfolding of the plagioclase vein yields > 200% of shortening. Insert blue arrows represent the north direction lying in a horizontal plane, the highlighted face of the cube shows the observed plane surface of the outcrop compared with the horizontal, and finally the line on the highlighted face of the cube shows the azimuth of the foliation. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

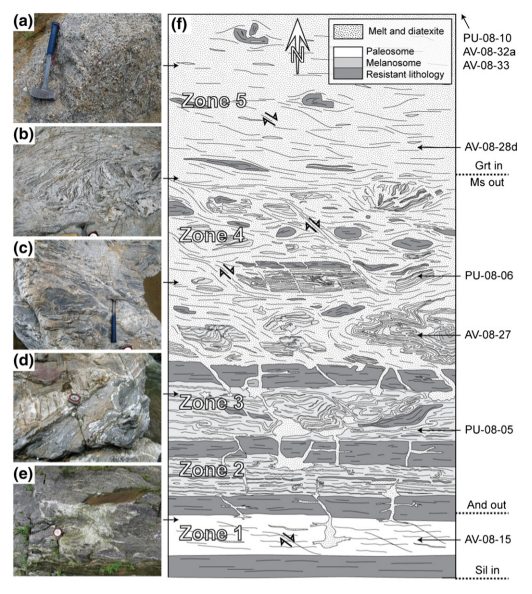


Fig. 4. Migmatitic "log" of the upper part of the La Bocana unit. (a–e), photographs of outcrops in relation to their structural level in the migmatitic log; (f) schematic log representing the upper migmatitic section for a 4 km thick; internal structures are purposely magnified and the different generations of leucosomes are not distinguished; zone 1: first appearance of melt with a very low fraction of in situ melting; zone 2: melting occurs through white-mica dehydration in the pelitic rich layers producing metatexites and melt fraction is inferior to 25%; between the zones 1 to 2, 2 to 3 and 3 to 4, psanmitic layers poor in micas, are not, or are weakly affected by in situ melting; zone 3: melt fraction increases, but the metatexitic texture is still dominant and several generations of leucosomes can be observed; they are either folded with residual layers or crosscutting the former ones; as the melt fraction increases, metatexites of zone 4 with large rafts of resistant bodies (quartz rich); zone 5: garnet appears and the volume of resistant lithologies decreases. The migmatitic acquire homogeneous diatexitic magma like rheology and no syn-migmatitic preferred orientation can be seen, at least at the macroscopic scale.

crosscut by leucocratic veins. Psammitic layers are less fertile (lower white mica content) and more competent, and show dilatant structures (Figs 3c & 4c), which allow leaching and facilitate circulation of melt throughout the crustal pile. Close to the garnet zone, melting affects the whole metasedimentary pile (Fig. 4, zone 4) forming diatexites with large amounts of rafts of resistant lithologies (quartz-rich unmelted bodies). Further north, garnet crystallizes and the volume of residual (biotite-rich) lithologies decreases significantly. At the top of this 7 km thick zone, melt is pervasive and early structures are erased. Resulting textures are mesocratic to melanocratic diatexites and have variable proportions of biotite and garnet (Fig. 4a). Within the garnet-bearing migmatitic layer, quartzite and other resistant bodies are dissolved in the melt toward the north, and are almost absent near the contact with the Piedras unit. Apparent kinematic indicators largely exhibit dextral transtension and roughly N-S dilatant structures. N-S structures of escaping melt are observed in competent layers (Fig. 4, zone 2 and 3) and show a southward direction of melt percolation (initially upward, taking into account subsequent tilting; Gabriele, 2002). Close to Piedras, the contact between the La Bocana and the Piedras units is tectonic and exhibits normal low-grade ductile faulting with top-to-the-south movement. In the western part, this contact zone exhibits mafic xenoliths (Fig. 2) and variably mixed dioritic magmas a short distance (up to 100 m) into the La Bocana unit. Resulting textures are produced by mingling and/or mixing of acidic and mafic melts (Fig. 3d). The Piedras unit as described by Aspden et al. (1995) is made of saussuritized metagabbros (now amphibolites, Fig. 3e). In addition, we found ultramafic lenses with relicts of olivine along the eastern cross-section, south of the Piedras locality. Farther north, the contact between the Piedras and Arenillas-Panupalí units (Naranjo Shear Zone) is diffused and highly deformed with dextral transpressive shear sense under low-grade metamorphic conditions (Fig. 3f). In addition, we observed a limited amount of serpentininite (up to 10%) within the Arenillas–Panupalí unit.

4. Petrography and mineral assemblages

Mineral assemblages of the continental sequence are described from the low-grade metasediments of the La Victoria unit to the south, to the high-grade molten metasediments of the La Bocana unit to the north. The location of studied samples is shown in Fig. 2 and characteristic mineral assemblages are presented in Table 1.

4.1. La Victoria metapelites

Near the boundary between the El Tigre and La Victoria units, the metapelite is composed of chlorite + biotite + quartz + plagioclase, thus defining a chlorite-biotite zone. Farther north, in the biotite zone (Fig. 2), the characteristic mineral assemblage is biotite + white mica + quartz + K-feldspar + plagioclase. In the psammitic layers, the mineral assemblage is dominated by quartz, with subordinate biotite and white mica. In the andalusite zone (Fig. 2) (AV-08-19b, Table 1), the typical mineral assemblage is biotite + white mica + and alusite + quartz + K-feldspar + plagioclase. Farther north, and alusite is progressively replaced by fibrolite (Feininger, 1978). Although cordierite is widely reported in this unit by Aspden et al. (1995), this mineral was not observed in examined samples. In the narrow and alusite-fribolite zone, the characteristic assemblage is and alusite + sillimanite + biotite + whitemica + quartz + K-feldspar + plagioclase. Weakly strained chiastolitic andalusite occurs as large porphyroblasts up to 5 cm. In rare samples andalusite is statically replaced by kyanite on its outer edge (AV-08-15). On the rim of these andalusite/kyanite porphyroblasts, kyanite is replaced by biotite and fibers of sillimanite (fibrolite) (Fig. 5a & b). The fibrolite defines the main foliation (Fig. 5b), and deflects the older fabric with apparent dextral transtensive movement.

The andalusite fribolite assemblage is transitional to the anatectic zone. Textural relationships observed in AV-08-15 sample show that the crystallization sequence was: andalusite, kyanite, and sillimanite. This suggests a clock-wise P-T path that looped around the Al_2SiO_5 triple point (Whitney, 2002).

4.2. La Bocana migmatites

On the basis of the mineralogical assemblage, we distinguished two sub-units in the La Bocana unit. Near the contact with the unmelted metasediments of the La Victoria unit, the upper migmatitic zone is characterized by metatexites without garnet (Fig. 4, zones 1 to 4). The lower migmatitic zone is characterized by garnet-bearing diatexites (Fig. 4, zones 5). On the eastern cross-section, this zone begins near the locality of La Bocana and extends northward to the Piedras unit.

In the upper migmatitic zone (Fig. 4, zone 4), the melanosome characteristic mineral assemblage is biotite + sillimanite + quartz +

K-feldspar + plagioclase \pm retrograde white-mica. Sillimanite occurs as prismatic aggregate of minerals associated with lath-like biotite and K-feldspar. Rare spinel is present as small inclusions of hercynite within large prismatic sillimanite (Fig. 5d). Cesare (1994) showed that inclusion of herycnite in prismatic sillimanite formed after breakdown of staurolite at 585–655 °C. Therefore, hercynite in the upper La Bocana migmatites likely represents relicts of prograde metamorphism at subsolidus conditions. Fibrolite occurs on the outer rim of the prismatic sillimanite often surrounded by late white-mica. Leucocratic layers exhibit larger grain size, and are dominated by quartz, plagioclase and K-feldspar. Biotite occurs as well crystallized, or interstitial, grains. Late white-mica occurs as late interstitial crystals at grain boundaries of plagioclase and K-feldspar. Vermicular intergrowth of feldspar and quartz at K-feldspar grain boundary are widely observed (Fig. 6a). Symplectic intergrowth of muscovite and quartz are also a common feature in this zone (Fig. 6a). These two last migmatitic microstructures are compatible with crystallization from melt. In this upper migmatite zone, garnet, orthopyroxene and cordierite are absent from the samples examined in this study. Biotite is present in every sample and its composition does not exhibits significant variation compared to the biotite of the La Victoria unit (Fig. 7d). Thus, natural observations are consistent with melting of the metasedimentary rocks in this zone within the stability field of the biotite at a temperature close to the solidus between 640 and 700 °C (Patiño-Douce and Harris, 1998; Vielzeuf and Holloway, 1988). Melting reactions principally involved white mica breakdown. Two major melting reactions are thought to occur in this zone, either by fluid-present, or fluid-absent white-mica dehydration melting (Patiño-Douce and Harris, 1998; Thompson and Tracy, 1979):

white mica + plagioclase + quartz + $H_2O = melt + sillimanite + biotite$ (1)

And

white mica + plagioclase + quartz = melt + sillimanite +K-feldspar + biotite (2)

In the lower migmatitic zone, the most significant microstructure associated with partial melting is the segregation of felsic material surrounding peritectic products, erasing pre-melting structures and leading to the formation of relatively homogeneous mesocratic to melanocratic migmatites (Fig. 4a). Neither preferred orientation, nor migmatitic foliation can be recognized, at least at the macroscopic scale. The characteristic mineral assemblage is biotite + garnet + sillimanite + quartz + K-feldspar + plagioclase. Garnet contains inclusions of biotite, prismatic sillimanite and plagioclase, ilmenite and/or rutile. In the north of the lower migmatitic zone, modal percent of garnet

Characteristic mineral assemblages of studied samples used in pseudosections. AV-08-15, AV-08-22b and PU-08-06 are also reported. GPS coordinates are in decimal degrees. I, inherited mineral; P, Peak thermal mineral; R, retrograde mineral. Mineral abbreviations used in figures and tables are from Kretz (1983).

	Localisation				Mineralogy									
Sample	Long	Lat	Unit	Lithology	Kfs	And	Ку	Sil	Gt	Bt	Ms	Chl	Ilm	Ru
AV-08-19b	- 79.927	- 3.755	La Victoria	Metapelite	Р	Ι	-	-	-	I + P	I + P	Р	Ι	-
AV-08-22b	-79.921	-3.714	La Victoria	Metapsammite	Р	-	-	-	-	I + P	I + P	Р	Ι	-
AV-08-13	-79.925	-3.747	La Victoria	Metapsammite	Р	Р	-	-	-	I + P	I + P	-	Ι	-
VI-08-04	-80.080	-3.755	La Victoria	Metapsammite	Р	-	-	Р	-	I + P	I + P	-	Ι	-
AV-08-15	-79.925	-3.739	La Victoria	Metapelite	Р	Ι	Ι	Р	-	I + P	I + P	-	Ι	-
PU-08-06	-80.045	-3.727	La Bocana	Sil migmatite	Р	-	-	Р	-	I + P	R	-	Ι	-
AV-08-24b	-79.927	-3.709	La Bocana	Sil migmatite	Р	-	-	Р	-	I + P + R	R	-	Ι	-
AV-08-27	-79.921	-3.702	La Bocana	Sil migmatite	Р	-	-	Р	-	I + P + R	R	-	Р	Ι
AV-08-28d	-79.904	-3.678	La Bocana	Gt migmatite	P + R	-	-	P + R	Р	P + R	R	R	Р	Ι
AV-08-32a	-79.906	-3.667	La Bocana	Gt migmatite	P + R	-	-	P + R	Р	P + R	R	R	Р	Ι
AV-08-33	-79.909	-3.664	La Bocana	Gt migmatite	P + R	-	-	P + R	Р	P + R	R	R	-	Ι
PU-08-10	-80.055	-3.716	La Bocana	Gt migmatite	P + R	-	-	P + R	Р	P + R	R	-	Ι	-

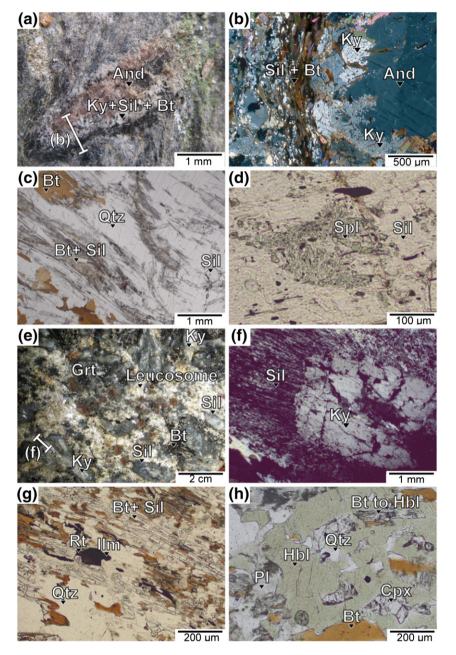


Fig. 5. Photograph of metasediments, under macroscopic view (a, e), polarized light (c, d,g, h), and polarized and analyzed light (b, f). (a) sample AV-08-15, andalusite is statically replaced by kyanite at rim and main foliation is underlined by finely crystallized fibrolite and biotite; (b) detailed microphotograph of (a), showing the microstructural relationships among the aluminosilicates; (c) biotite-sillimanite peritectic aggregates produced by dehydration melting of white-mica at the boundary of large grain of quartz; (d) spinel (hercynite) inclusions in sillimanite; (e) macroscopic view of a highly residual garnet-bearing migmatite with leucosome and associated peritectic garnet; note the associated breakdown of kyanite to sillimanite; (g) characteristic texture and mineral assemblage of garnet-bearing migmatites, rutile and ilmenite are in equilibrium; (h) typical paragenesis of the mixing zone between the Piedras gabbroic unit and the La Bocana migmatitic unit.

increases, garnet becomes skeletal and elongated, and the sillimanite content decreases. Biotite occurs, either as elongated lath-like aggregate together with sillimanite, or as interstitial grains within quartz + plagioclase + K-feldspar rich layers (Fig. 5g). While garnet is well preserved in many samples, with only limited breakdown into biotite at rims, K-feldspar belonging to residual aggregates (schlieren) appears to be largely recrystallized and/or breakdown. Fig. 6e & f show penetrative breakdown of K-feldspar in symplectic intergrowth of biotite + sillimanite + quartz + plagiocase indicating that these phases were crystallizing from the melt during cooling (Spear et al., 1999). In the lower migmatitic zone white-mica is almost absent. However, in a few samples such as AV-08-35, white mica overgrows plagioclase and K-feldspar, forming a common low-temperature alteration product

(Fig. 6b). The appearance of garnet, absence of white mica and decreasing sillimanite and biotite content toward higher metamorphic grade implies incongruent melting produced by biotite dehydration (Patiño-Douce and Jonhston, 1991; Vielzeuf and Holloway, 1988):

 $biotite + alumino-silicate + quartz = melt + garnet + K-feldspar \qquad (3)$

and

biotite + plagioclase + quartz \pm H₂O = melt + garnet \pm K-feldspar (4)

Whether reactions (3) and (4) produce K-feldspar as peritectic phase is dependent on the activity of H₂O. In the La Victoria and the

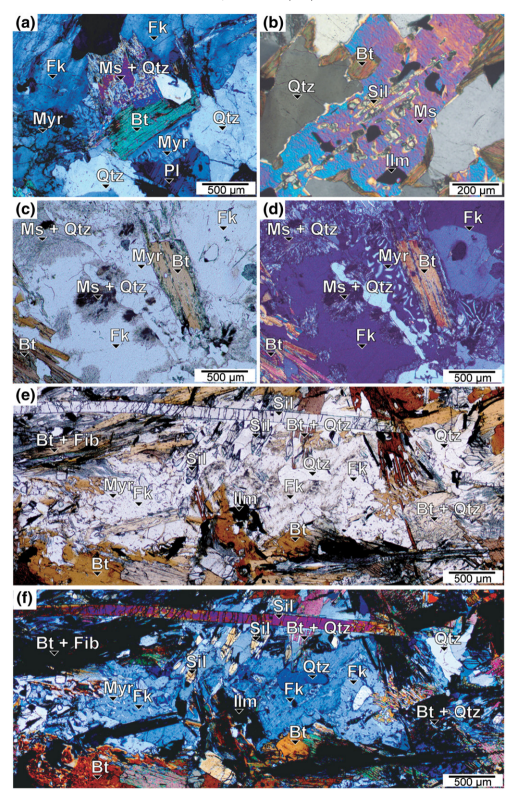
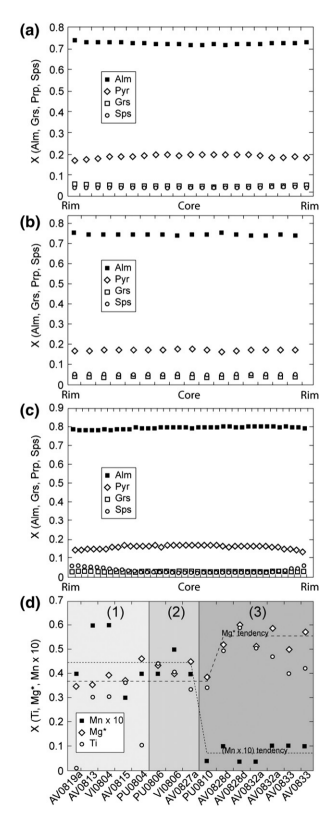


Fig. 6. Microphotograph of partially molten metasediments under polarized and analyzed light (a, b, d, f) and polarized light (c, e). (a) sample PU-08-06, interaction between residual melt and K-feldspar forming myrmekite and muscovite + quartz symplectite; (b) late crystallization of muscovite in AV-08-350 garnet-bearing sample; (c, d) AV-08-28d garnet-bearing sample, retrograde interaction between residual melt and K-feldspar forming myrmekite and likely contributed to the penetrative saussuritization of the K-feldspar in muscovite + quartz; (e, f) PU-08-10 garnet-bearing sample, retrograde interaction between residual melt and a large crystal of K-feldspar. The former K-feldspar grains are largely broken down by the formation of myrmekite, crystallization of quartz + biotite symplectite and late crystallization of crosscutting prismatic sillimanite crystals.

La Bocana units graphite is omnipresent in examined samples. Connolly and Cesare (1993) showed that C–O–H fluid produced by the equilibration of H₂O and excess graphite must maintain the atomic H/O ratio of water, 2:1. This precludes water saturated conditions during partial melting and thus favors reactions (2) and (3) of dehydration melting in the Upper and the Lower La Bocana units, respectively.

Along the western cross-section, at the transition zone between the upper and lower migmatitic zone, garnet is associated in highly residual layers. Relict kyanite is replaced by pluricentimetric sillimanite (Fig. 5e & f). For metapelitic lithologies ilmenite and rutile are the two main Ti-bearing minerals as ilmenite and rutile are stable at low and high pressure, respectively. Ilmenite and/or rutile present in the studied samples exhibits a South to North modal variability (Table 1, Fig. 2). Whereas in the la Victoria unit only ilmenite has been observed, in the



lower migmatitic zone, we observe layers containing both ilmenite and rutile in equilibrium (Fig. 5g). North of this area only rutile is observed (see Fig. 2). This mineral succession, from ilmenite in the La Victoria unit to rutile in the La Bocana unit, suggests an increase in pressure conditions toward the north. In sample AV-08-28d, ilmenite is present in the matrix with relict rutile cores, whereas rutile is not observed as inclusions in garnet. Crystallization of sillimanite from kyanite and of ilmenite from rutile shows that the continental sequence recorded heating and/or decompression, which is consistent with the clock-wise P-T path around the Al₂O₃ triple point observed in sample AV-08-15. Furthermore, the lack of rutile inclusions in garnet in sample AV-08-28d indicates that decompression started before garnet crystallization.

4.3. Piedras amphibolites

In the Piedras unit, the characteristic mineral assemblage is hornblende + plagioclase + titanite + epidote + actinolite \pm quartz \pm clinopyroxene. Clinopyroxene is rare and its rims are always replaced by hornblende (Aspden et al., 1995). Plagioclase is widely saussuritized, and epidote and actinolite represent retrogression under greenschist facies conditions. Near the locality of Piedras, ultramafic lenses (Fig. 2) with characteristic mineral assemblage of olivine + serpentine + magnetite \pm clinopyroxene characteristic mineral assemblage were identified. In the western cross-section, in the contact zone with the La Bocana migmatites, mixing of acid and basic magmas (Fig. 3d) produced a mineral assemblage composed of hornblende + clinopyroxene + biotite + quartz + plagioclase (Fig. 5h).

4.4. Arenillas-Panupalí blueschists

The Arenillas–Panupalí unit is mainly represented by metabasalts metamorphosed under blueschist facies conditions, which underwent an intense greenschist facies overprint. Pressure peak-metamorphic conditions are characterized by the garnet + glaucophane + titanite assemblage (Gabriele, 2002). The greenschist overprint is documented by crystallization of fine-grained albite, chlorite and epidote, and by glaucophane breaking down to barroisite and actinolite (Gabriele, 2002).

5. Mineral chemistry

Representative analyses of garnet are presented in Table 2. Almost all garnets are found in the lower migmatitic zone, and are interpreted as peritectic phases (no sub-solidus garnet). Garnet exhibits little or no chemical variations from core to rim (Fig. 7a–c). X_{Alm} content from sample-to-sample (Table 1) ranges from 0.70 to 0.80, X_{Prp} content from 0.13 to 0.20, X_{Grs} from 0.02 to 0.06 and X_{Sps} from 0.02 to 0.06. X_{Alm} and X_{Prp} variations between different samples reflect their differences in MgO/(MgO + FeO) bulk composition (Table 7). Representative analyses of biotite are presented in Table 3. In metasediments, the X_{Mg} in biotite ranges from 0.33 to 0.46 in metapelites, and from 0.48 to 0.62 in garnet-bearing migmatites (Fig. 7d). Titanium content per formulae unit (p.f.u.) varies from 0.10 to 0.44 for unmolten metapelites,

Fig. 7. <u>Garnet and biotite chemical composition of the El Oro metamorphic complex El</u> Oro metamorphic complex. (a, b, c) Microprobe analysis across garnet crystal of AV-08-28d (a), AV-08-32a (b) and PU-08-10 (c) samples. Most of the end-members exhibit relatively flat pattern for all samples, excepted for (b) and (c) where X_{pyr} and X_{sps} are respectively, slightly depleted and enriched at rim; (d) $X_{(Ti, Mg^*, Mn \times 10)}$ of biotite of various listed samples across the El Oro metamorphic complex. (1) La Victoria metasediments, (2) upper the La Bocana metatexites, (3) lower the La Bocana garnet-bearing migmatites. Mg^{*} = Mg/(Fe + Mg). The Mg^{*} tendency exhibits an increase from 0.36 in the zone (1) and (2) to 0.55 in the zone (3) where partial melting occurred under biotite dehydration melting. X_{Mn} (Mn p.f.u. * 10, see Table 3) drops from 0.45 in zones (1) and (2) down to <0.1 in zone (3). This coincides with the condition of partial melting of biotite. Storage of Mn in biotite in lower metamorphic grades (zone (1) and (2)) can account for the absence of garnet whereas pseudosections stabilize garnet when MnO is used. X_{Ti} (p.f.u., see Table 3) is of 0.3 to 0.4 in zone (1) and (2) and becomes >0.4 in zone (3).

Sample	PU0803	PU0810		AV0829	AV0832a	AV0828d	AV0833
Unit	La Vic.	L. B.		L. B.	L. B.	L. B.	L. B.
Position	Core	Core	Rim	Core	Core	Core	Core
SiO ₂	36	37.71	37.93	36.62	37.08	37.9	37.23
TiO ₂	0	0.08	0	0.02	0.05	0.09	0.03
Al ₂ O ₃	20.3	20.77	20.79	21.26	21.72	21.34	20.972
FeO	24.4	35.52	34.89	34.13	33.2	32.78	33.49
MnO	12.4	0.87	2.48	2.4	1.77	1.36	2.29
MgO	3	4.04	3.4	3.99	4.64	4.77	4.44
CaO	3.5	0.95	0.81	1.35	1.58	1.77	1.46
Total	99.6	99.95	100.31	99.77	100.04	100.01	99.9
Structural formulae	(12 anhydrous oxygen bo	ısis)					
Si	2.88	2.77	2.78	2.71	2.71	2.76	2.74
Ti	0	0	0	0	0	0	0
Al	1.92	1.8	1.8	1.85	1.87	1.83	1.82
Fe	1.64	2.18	2.14	2.11	2.03	1.994	2.06
Mn	0.85	0.05	0.15	0.15	0.11	0.08	0.14
Mg	0.36	0.44	0.37	0.44	0.51	0.52	0.49
Ca	0.3	0.07	0.06	0.11	0.12	0.14	0.11
Total	7.94	7.33	7.32	7.37	7.35	7.32	7.35
Endmembers (mol%)							
Ругоре	11.34	16.07	13.62	15.67	18.27	18.93	17.36
Almandine	52.16	79.25	78.4	75.17	73.31	72.96	73.45
Grossular	9.56	2.72	2.33	3.81	4.47	5.05	4.1
Spessartine	26.94	1.97	5.64	5.35	3.96	3.07	5.09
Mg/(Fe + Mg)	0.18	0.17	0.15	0.17	0.20	0.21	0.19

Representative electron microprobe analysis of garnet in the La Bocana garnet-bearing migmatitic samples. Structural formulae have been calculated on the basis of 12 atoms oxygen; calculated end members shown in the table are used to calculate the *P*–*T* conditions in pseudosections. La Vic., La Victoria, U. B., Upper La Bocana, L. B., Lower La Bocana.

to a range of 0.40 to 0.60 for migmatites (Fig. 7d). X_{Mnx10} varies from 0.30 to 0.60 for metapelites and low grade migmatites (without garnet), whereas in garnet-bearing migmatites, X_{Mnx10} varies from 0.05 to 0.10. This drop in X_{Mnx10} correlates with the apparition of garnet and with the ongoing biotite dehydration melting.

Titanium content and X_{Mg} increase in biotite is compatible with biotite progressive incongruent melting (Patiño-Douce and Harris, 1998). White mica is present in metapelites as a prograde and peak-*T* phase, whereas in the garnet-bearing migmatites, white mica occurs as a retrograde phase that results from saussuritization of feldspar or late back reaction from melt (Fig. 6c–f). In metapelites and migmatites, white mica shows no significant compositional differences and is muscovite-rich (X_{Mu} >0.80) with X_{Cel} accounting for the rest (Table 4). Silica content (p.f.u.) ranges from 3.00 to 3.05. In metapelite, plagioclase has high X_{Ab} , between 0.75 and 0.85. In high grade metapelites close to the anatectic zone, plagioclase is zoned and exhibits a slight enrichment in X_{Ab} from core to rim (0.77 to 0.83). In migmatites, plagioclase composition is more variable and X_{Ab} ranges from 0.60 to 1.00. Plagioclase commonly exhibits symplectite textures of micron-scale white mica + plagioclase with lower X_{An} .

6. Thermobarometry

Thermobarometry has been performed combining three independent approaches: the Raman Spectroscopy on Carbonaceous Matter thermometry (RSCM see Appendix A), pseudosections and Fe–Mg exchange thermometry.

6.1. RSCM thermometry and garnet-biotite thermometry

As carbonaceous material is present in all metasediments of the continental sequence, we used RSCM thermometry, which is based on the degree of organization of carbonaceous material. This method gives estimates on the maximal temperature undergone by the rocks for thermal conditions between 330 °C and 640 °C \pm 50 °C (Beyssac et al., 2002, 2004). RSCM is also applied on middle to relatively high grade metamorphic rocks up to the complete graphitization at

~640 °C (Robert et al., 2010). Table 6 shows the RSCM thermometry results. T_{max} estimated with the RSCM method, range from 475 to 640 °C, with a general increase in temperature from south to north.

PU-08-03 is the only unmolten metasediment in which small grains of garnet are present. Garnet–biotite thermometery on this sample yields temperatures of 560 ± 50 °C and 570 ± 50 °C, with the calibrations of Ferry and Spear (1978) and Perchuk and Lavrent'eva (1983), respectively. $T_{\rm max}$ with RSCM thermometry on the same sample gives 567 ± 50 °C (Table 6), which is in very good agreement with the garnet–biotite temperature. For high grade migmatitic samples, inclusions of graphite were rare and often located on the surface of the host minerals. Few analyses were carried out on these samples (3–5 spectra per sample), but the obtained spectra are of good quality ($1\sigma < 8$ °C). $T_{\rm max}$ obtained in migmatites reach the graphitization *T* and are indicative of minimum temperature. We observe no discrepancy between the peak *T* mineralogical assemblages and the RSCM thermometry.

6.2. Pseudosections

P-T and T-X pseudosections have been calculated (1) to precise metamorphic P-T conditions and more particularly conditions of partial melting, and (2) to determine the type of prograde melting reactions. Phase relations have been modeled in the Ti(Mn)NKFMASH system using Perple_X'07 software (Connolly, 2005) and using the internally consistent thermodynamic database of Holland and Powell (1998). Bulk rock composition, solution models and end-member phases considered in the modeling are listed in Tables 7 and 8, respectively. Although the TiO₂ component is a minor component, it has been included to the model in order to take into account the rutile/ilmenite transition observed along the cross-section (Fig. 2, Table 1). MnO is taken into account for garnet-bearing migmatite in order to better estimate the end-member composition of garnet. In unmelted metasediments, the only MnO-bearing phases are biotite and ilmenite as garnet which typically contains MnO has not been observed. Therefore MnO is not taken into consideration for the calculation of pseudosections of unmelted metasediments as MnO as little effect on the stability field of biotite. Water content in the P-T

Representative electron microprobe analysis of biotite in metasediments and migmatites. Structural formulae have been calculated on the basis of 11 atoms oxygen; analyzed samples are located in Fig. 2. La Vic., La Victoria, U. B., Upper La Bocana, L. B., Lower La Bocana.

Sample	AV0819a	AV0813	VI0803	VI0804	AV0815	PU0804	PU0806	VI0806	AV0827a	PU0810	AV0828d	AV0828d	AV0832a	AV0832a	AV0833	AV0833
Unit	La Vic.	La Vic.	La Vic.	La Vic.	La Vic.	La Vic.	U. B.	U. B	L. B.	L. B.	L. B.	L. B.	L. B.	L. B.	L. B.	L. B.
Na ₂ O	0.23	0.23	0.17	0.20	0.16	0.30	0.27	0.24	0.06	0.19	0.14	0.34	0.15	0.15	0.19	0.24
MgO	6.71	6.89	7.46	7.64	6.01	8.56	7.34	7.38	7.97	7.13	9.83	11.53	9.82	11.53	9.83	11.71
Al_2O_3	19.13	19.05	19.90	19.54	22.52	21.56	19.20	19.25	19.06	19.96	17.23	18.13	18.52	17.88	19.15	18.84
SiO ₂	36.46	34.98	34.70	34.75	35.10	34.80	34.60	34.95	36.42	34.06	36.77	37.33	35.89	36.90	35.19	35.53
K ₂ O	9.12	10.12	9.44	9.24	9.13	8.40	9.43	9.47	8.79	9.79	9.39	9.60	9.65	9.71	9.86	9.54
CaO	0.01	0.07	0.01	0.03	0.52	0.20	0.02	0.04	0.08	0.00	0.05	0.05	0.00	0.01	0.01	0.00
TiO ₂	0.09	2.56	2.52	2.41	2.73	0.84	3.69	3.50	2.70	2.96	4.54	5.37	4.33	4.14	3.45	3.78
MnO	0.29	0.33	0.43	0.46	0.24	0.33	0.30	0.36	0.31	0.01	0.10	0.02	0.02	0.06	0.07	0.08
FeO	22.22	21.51	20.15	20.80	18.84	18.24	19.86	20.55	18.03	20.42	17.86	14.43	16.65	14.87	17.24	15.20
Total	94.26	95.74	94.78	95.05	95.25	93.24	94.71	95.75	93.42	94.40	95.92	96.79	95.03	95.24	95.00	94.92
Structural formulae Si	(11 anhydrous 5.77	oxygen basis 5.50) 5.44	5.45	5.52	5.44	5.43	5.44	5.67	5.38	5.60	5.52	5.49	5.57	5.42	5.40
Ti Alvi	0.01	0.30	0.30	0.28	0.31	0.10	0.43	0.41	0.32	0.38	0.52	0.60	0.50	0.47	0.40	0.43
	1.34	1.03	1.12	1.06	1.57	1.42	0.98	0.97	1.17	1.10	0.69	0.68	0.83	0.75	0.90	0.78
Aliv	2.23	2.50	2.56	2.55	2.48	2.56	2.57	2.56	2.33	2.62	2.40	2.48	2.51	2.43	2.58	2.60
Al	3.57	3.53	3.68	3.61	4.05	3.98	3.55	3.53	3.50	3.72	3.09	3.16	3.34	3.18	3.48	3.38
Fe(ii)	2.94	2.83	2.64	2.73	2.41	2.39	2.60	2.67	2.35	2.70	2.28	1.78	2.13	1.88	2.22	1.93
Mn	0.04 1.58	0.04 1.61	0.06	0.06	0.03 1.37	0.04 2.00	0.04 1.72	0.05	0.04	0.00 1.68	0.01	0.00	0.00 2.24	0.01 2.59	0.01 2.26	0.01 2.65
Mg	0.07		1.74 0.05	1.78				1.71	1.85		2.23	2.54			2.26	
Na		0.07		0.06	0.05	0.09	0.08	0.07	0.02	0.06	0.04	0.10	0.04	0.04		0.07
K Ti	1.84	2.03	1.89	1.85	1.58	1.68	1.89	1.88	1.75	1.97	1.83	1.81	1.88	1.87	1.94	1.85
	0.01	0.30	0.30	0.28	0.31	0.10	0.43	0.41	0.32	0.34	0.52	0.60	0.50	0.47	0.40	0.43
Mg(Fe + Mg)	0.35	0.36	0.40	0.39	0.36	0.46	0.40	0.39	0.44	0.38	0.49	0.59	0.51	0.58	0.50	0.58
Mn*10	0.40	0.40	0.60	0.60	0.30	0.40	0.40	0.50	0.40	0.00	0.10	0.00	0.00	0.10	0.10	0.10
Ti/(Ti + Fe + Mg)	0.00	0.06	0.06	0.06	0.08	0.02	0.09	0.09	0.07	0.08	0.10	0.12	0.10	0.10	0.08	0.09

Representative electron microprobe analysis of muscovite in metasediments and migmatites. Structural formulae have been calculated on the basis of 11 atoms oxygen; analyzed samples are located in Fig. 2. La Vic., La Victoria, U. B., Upper La Bocana, L. B., Lower La Bocana.

Unit La Vic. La Vic. La Vic. La Vic. La Vic. La Vic. U. B. L. B. Na20 0.84 0.30 0.59 0.55 0.63 0.32 MgO 0.38 0.75 0.49 0.49 0.62 0.89 Al ₂ O ₃ 36.78 34.03 36.69 36.80 35.77 35.76 SiO ₂ 48.61 46.57 45.08 45.17 45.50 49.10 K2O 8.97 10.27 10.58 9.91 10.77 8.40 CaO 0.00 0.15 0.00 0.03 0.00 1.28 MnO 0.00 0.02 0.00 0.01 1.00 0.03 FeO 0.94 1.34 0.90 1.05 1.08 1.20 Zr ₂ O ₃ 0.03 0.00 0.00 0.00 0.00 0.00 Structural formulae (11 anhydrous oxygen basis) Si 3.14 3.13 3.01 3.02 3.02 3.16	Sample	AV0819a	AV0813	AV0815	VI0803	PU0806	AV08350
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	Unit	La Vic.	La Vic.	La Vic.	La Vic.	U. B.	L. B.
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	Na ₂ O	0.84	0.30	0.59	0.55	0.63	0.32
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	MgO	0.38	0.75	0.49	0.49	0.62	0.89
K20 8.97 10.27 10.58 9.91 10.77 8.40 Ca0 0.00 0.15 0.00 0.03 0.00 0.00 TiO2 0.10 0.94 0.66 0.63 0.00 1.28 MnO 0.00 0.02 0.00 0.01 1.00 0.03 FeO 0.94 1.34 0.90 1.05 1.08 1.20 Zr_2O3 0.03 0.00 0.00 0.00 0.00 0.00 0.00 Total 96.64 94.38 95.01 94.64 96.17 96.99 Structural formulae (11 anhydrous oxygen basis) 51 3.14 3.13 3.01 3.02 3.02 3.16 Ti 0.01 0.05 0.04 0.03 0.00 0.06 Aliv 1.94 1.83 1.90 1.91 1.88 1.87 Aliv 0.86 0.87 0.99 0.99 0.98 0.85 Al 2.80 <td>Al_2O_3</td> <td>36.78</td> <td>34.03</td> <td>36.69</td> <td>36.80</td> <td>36.57</td> <td>35.76</td>	Al_2O_3	36.78	34.03	36.69	36.80	36.57	35.76
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	SiO ₂	48.61	46.57	45.08	45.17	45.50	49.10
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	K20	8.97	10.27	10.58	9.91	10.77	8.40
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	CaO	0.00	0.15	0.00	0.03	0.00	0.00
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	TiO ₂	0.10	0.94	0.66	0.63	0.00	1.28
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	MnO	0.00	0.02	0.00	0.01	1.00	0.03
Total96.6494.3895.0194.6496.1796.99Structural formulae (11 anhydrous oxygen basis)Si 3.14 3.13 3.01 3.02 3.02 Ti 0.01 0.05 0.04 0.03 0.00 0.06 Alvi 1.94 1.83 1.90 1.91 1.88 1.87 Aliv 0.86 0.87 0.99 0.99 0.98 0.85 Al 2.80 2.70 2.89 2.90 2.86 2.71 Fe(ii) 0.05 0.08 0.05 0.06 0.07 Mn 0.00 0.00 0.00 0.00 0.00 Mg 0.04 0.08 0.05 0.06 0.09 Na 0.11 0.04 0.08 0.07 0.08 0.04 K 0.74 0.88 0.90 0.85 0.91 0.69 Endmembers (mol%)Pyrophilite 15.5 8.0 2.5 8.5 1.0 27.0 Céladonite 1.0 22.5 10.0 5.0 1.0 10.5 Muscovite 81.0 72.0 87.5 84.5 92.5 61.0	FeO	0.94	1.34	0.90	1.05	1.08	1.20
Structural formulae (11 anhydrous oxygen basis)Si 3.14 3.13 3.01 3.02 3.02 3.16 Ti 0.01 0.05 0.04 0.03 0.00 0.06 Alvi 1.94 1.83 1.90 1.91 1.88 1.87 Aliv 0.86 0.87 0.99 0.99 0.98 0.85 Al 2.80 2.70 2.89 2.90 2.86 2.71 Fe(ii) 0.05 0.08 0.05 0.06 0.07 Mn 0.00 0.00 0.00 0.00 0.00 Mg 0.04 0.08 0.05 0.06 0.09 Na 0.11 0.04 0.08 0.07 0.08 0.04 K 0.74 0.88 0.90 0.85 0.91 0.69 Endmembers (mol%)Pyrophilite 1.5 8.0 2.5 8.5 1.0 27.0 Céladonite 1.0 22.5 10.0 5.0 1.0 10.5 Muscovite 81.0 72.0 87.5 84.5 92.5 61.0	Zr_2O_3	0.03	0.00	0.00	0.00	0.00	0.00
Si 3.14 3.13 3.01 3.02 3.02 3.16 Ti 0.01 0.05 0.04 0.03 0.00 0.06 Alvi 1.94 1.83 1.90 1.91 1.88 1.87 Aliv 0.86 0.87 0.99 0.99 0.98 0.85 Al 2.80 2.70 2.89 2.90 2.86 2.71 Fe(ii) 0.05 0.08 0.05 0.06 0.06 0.07 Mn 0.00 0.00 0.00 0.00 0.00 0.00 0.06 Mg 0.04 0.08 0.05 0.05 0.06 0.09 Na 0.11 0.04 0.88 0.90 0.85 0.91 0.69 Endmembers (mol%) Pyrophilite 15.5 8.0 2.5 8.5 1.0 27.0 Céladonite 1.0 22.5 10.0 5.0 1.0 10.5 Muscovite 81.0	Total	96.64	94.38	95.01	94.64	96.17	96.99
Si 3.14 3.13 3.01 3.02 3.02 3.16 Ti 0.01 0.05 0.04 0.03 0.00 0.06 Alvi 1.94 1.83 1.90 1.91 1.88 1.87 Aliv 0.86 0.87 0.99 0.99 0.98 0.85 Al 2.80 2.70 2.89 2.90 2.86 2.71 Fe(ii) 0.05 0.08 0.05 0.06 0.06 0.07 Mn 0.00 0.00 0.00 0.00 0.00 0.00 0.06 Mg 0.04 0.08 0.05 0.05 0.06 0.09 Na 0.11 0.04 0.88 0.90 0.85 0.91 0.69 Endmembers (mol%) Pyrophilite 15.5 8.0 2.5 8.5 1.0 27.0 Céladonite 1.0 22.5 10.0 5.0 1.0 10.5 Muscovite 81.0	Churrent for		huduana au	(man hania)			
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$					2.02	2.02	2.16
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Mn 0.00 0.00 0.00 0.00 0.06 0.00 Mg 0.04 0.08 0.05 0.05 0.06 0.09 Na 0.11 0.04 0.08 0.07 0.08 0.04 K 0.74 0.88 0.90 0.85 0.91 0.69 Endmembers (mol%) Pyrophilite 15.5 8.0 2.5 8.5 1.0 27.0 Céladonite 1.0 22.5 10.0 5.0 1.0 10.5 Muscovite 81.0 72.0 87.5 84.5 92.5 61.0							
Mg 0.04 0.08 0.05 0.05 0.06 0.09 Na 0.11 0.04 0.08 0.07 0.08 0.04 K 0.74 0.88 0.90 0.85 0.91 0.69 Endmembers (mol%)	. ,						
Na 0.11 0.04 0.08 0.07 0.08 0.04 K 0.74 0.88 0.90 0.85 0.91 0.69 Endmembers (mol%) Pyrophilite 15.5 8.0 2.5 8.5 1.0 27.0 Céladonite 1.0 22.5 10.0 5.0 1.0 10.5 Muscovite 81.0 72.0 87.5 84.5 92.5 61.0							
K 0.74 0.88 0.90 0.85 0.91 0.69 Endmembers (mol%) <th< td=""><td>0</td><td></td><td></td><td></td><td></td><td></td><td></td></th<>	0						
Endmembers (mol%) Pyrophilite 15.5 8.0 2.5 8.5 1.0 27.0 Céladonite 1.0 22.5 10.0 5.0 1.0 10.5 Muscovite 81.0 72.0 87.5 84.5 92.5 61.0							
Pyrophilite 15.5 8.0 2.5 8.5 1.0 27.0 Céladonite 1.0 22.5 10.0 5.0 1.0 10.5 Muscovite 81.0 72.0 87.5 84.5 92.5 61.0	K	0.74	0.00	0.50	0.05	0.51	0.05
Céladonite 1.0 22.5 10.0 5.0 1.0 10.5 Muscovite 81.0 72.0 87.5 84.5 92.5 61.0	Endmembers	(mol%)					
Muscovite 81.0 72.0 87.5 84.5 92.5 61.0	Pyrophilite	15.5	8.0	2.5	8.5	1.0	27.0
	Céladonite	1.0	22.5	10.0	5.0	1.0	10.5
Phlogopite 2.5 – 2.5 0.0 2.0 5.5 1.5	Muscovite	81.0	72.0	87.5	84.5	92.5	61.0
	Phlogopite	2.5	-2.5	0.0	2.0	5.5	1.5

pseudosections is kept constant for the entire range of P–T conditions. This water content was estimated for each sample by calculating T–X_{H20} pseudosection and P–X_{H20} pseudosections. The chosen water content

Table 6

RSCM thermometry results of studied samples. Analytical uncertainties on the estimation of temperature are quoted at 1σ while the calibration error for each sample on the determination of absolute temperature is of \pm 50 °C.

Sample	No analysis	R2	Std	T °C Beyssac et al., (2002)	1σ
AV0813	9	0.11	0.03	591	4.73
AV0815	7	0.07	0.01	609	2.47
AV0819b	10	0.38	0.05	474	6.93
AV0822b	8	0.24	0.04	532	5.79
AV0824b	4	0.02	0.01	632	1.09
PU0802	9	0.24	0.05	535	7.96
PU0803	8	0.17	0.04	567	6.06
PU0806	3	0.04	0.01	625	1.65
PU0811	3	0.01	0.01	638	2.03
VI0802	6	0.22	0.04	543	6.60
VI0804	5	0.09	0.03	599	5.77

corresponds to the minimum amount of water necessary to saturate the sub-solidus assemblage just below the solidus with no free water. In these conditions C–O–H fluids in the graphitic saturated system of the metasediments of the La Victoria and the La Bocana units have not been modeled.

6.2.1. Metasediments

We selected one metapsammite (AV-08-13) and one metapelite (AV-08-15) belonging to a different structural level of the La Victoria unit in order to constrain the evolution of the P-T conditions within the metasedimentary sequence. Sample AV-08-15 and sample AV-08-13 are located in the narrow andalusite–fibrolite zone (Fig. 2).

In the metasediments of the La Victoria unit, the association of biotite + sillimanite \pm white mica defines the foliation (Fig. 5b). In sample AV-08-15, the observed mineral assemblage, X_{Mg} of biotite (0.36), X_{Ab} (0.79) shows that the $P-T_{max}$ conditions lie in the field of biotite + plagioclase + K-feldspar + muscovite + sillimanite + quartz (Fig. 8). This is consistent with RSCM thermometry of 609 ± 50 °C. Thus, associated extensional microstructures observed in sample

Representative electron microprobe analysis of plagioclase in metasediments and migmatites. Structural formulae have been calculated on the basis of 32 atoms oxygen; analyzed samples are located in Fig. 2. La Vic., La Victoria, U. B., Upper La Bocana, L. B., Lower La Bocana.

Sample	AV-08-13	AV-08-15	PU0806	AV0828d	AV0828d	PU0810	PU0810	AV0832a	AV0833
Unit	La Vic.	La Vic.	U. B.	L. B.	L. B.	L. B.	L. B.	L. B.	L. B.
Position	Core	Core	Core	Schlieren	Leucosome	Schlieren	Leucosome	Schlieren	Schlieren
Na ₂ O	9.47	9.34	9.52	6.98	7.90	9.08	10.20	8.23	7.92
MgO	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00
Al_2O_3	22.68	22.93	24.00	25.78	25.13	22.53	21.14	25.21	25.12
SiO ₂	63.36	63.62	62.71	59.39	60.27	63.90	65.72	60.42	59.87
K2O	0.12	0.23	0.14	0.51	0.22	0.22	0.33	0.22	0.24
CaO	3.40	4.14	4.04	7.38	6.71	4.51	2.64	6.64	6.74
TiO ₂	0.00	0.02	0.02	0.02	0.00	0.04	0.02	0.02	0.05
MnO	0.00	0.04	0.03	0.04	0.01	0.00	0.00	0.03	0.01
FeO	0.00	0.04	0.21	0.24	0.04	0.05	0.00	0.03	0.03
Zr_2O_3	0.00	0.02	0.00	0.00	0.00	0.00	0.00	0.00	0.00
Total	99.04	100.38	100.66	100.35	100.28	100.33	100.06	100.80	99.96
Numbers of io	ns on the basis of 3	2 0							
Si	11.38	11.42	11.26	10.66	10.82	11.47	11.80	10.85	10.75
Al	4.80	4.85	5.08	5.46	5.32	4.77	4.47	5.33	5.31
Fe(ii)	0.00	0.01	0.03	0.04	0.01	0.01	0.00	0.00	0.00
Ca	0.65	0.80	0.78	1.42	1.29	0.87	0.51	1.28	1.30
Na	3.30	3.25	3.31	2.43	2.75	3.16	3.55	2.86	2.76
К	0.03	0.05	0.03	0.12	0.05	0.05	0.08	0.05	0.05
Ba	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00
Total	20.15	20.38	20.49	20.12	20.23	20.33	20.41	20.38	20.17
Endmembers ((mol%)								
Anorthite	16.46	19.42	18.84	35.77	31.54	21.27	12.27	30.45	31.56
Albite	82.85	79.29	80.37	61.27	67.21	77.52	85.89	68.35	67.11
Orthose	0.69	1.29	0.79	2.97	1.26	1.22	1.84	1.21	1.33

Table 7
Bulk rock composition used in this study. \leq LD, below limit of detection.

Sample	AV-08-28d	AV-08-32a	AV-08-33	AV-08-13	AV-08-15	PU-08-10
SiO ₂	69.2	69.64	71.68	73.85	59.54	61.3
TiO ₂	0.78	0.8	0.66	0.52	0.90	0.94
Al ₂ O ₃	14.28	13.76	12.4	13.25	19.23	15.55
FeO	5.29	4.81	5.47	3.34	7.16	8.59
MnO	0.08	0.07	0.12	0.06	0.08	0.09
MgO	1.89	1.73	1.69	0.99	2.12	2.05
CaO	1.76	1.77	1.81	0.78	0.56	0.44
Na ₂ O	1.52	2.41	2	2.35	1.26	1.39
K ₂ 0	2.25	2.61	2.21	3.36	4.53	3.74
P ₂ O ₅	0.2	<ld< td=""><td><ld< td=""><td><ld< td=""><td>1.75</td><td>0.39</td></ld<></td></ld<></td></ld<>	<ld< td=""><td><ld< td=""><td>1.75</td><td>0.39</td></ld<></td></ld<>	<ld< td=""><td>1.75</td><td>0.39</td></ld<>	1.75	0.39
LOI	2.15	1.63	1.22	1.58	2.17	3.11
Total	99.4	99.23	99.27	100.08	99.20	97.59
MgO/(FeO + MgO)	0.26	0.26	0.24	0.23	0.23	0.19

AV-08-15 during crystallization of sillimanite and the metastability of and alusite (Fig. 3a&b) constrains the $P-T_{max}$ at 4 ± 2 kbar and $620 \pm$ 50 °C (Fig. 8).

In sample AV-08-13, the observed mineral assemblage, X_{Mg} of biotite (0.36), X_{Ab} (0.83) and RSCM thermometry (580±50 °C) show that the $P-T_{max}$ lies in the field of biotite + plagioclase + K-feldspar + muscovite + andalusite + quartz mineral assemblage (Appendix C., Fig. 1). Estimated $P-T_{max}$ are of 3 ± 2 kbar and 580 ± 50 °C.

6.2.2. Garnet-bearing migmatites

Four garnet-bearing migmatites have been selected, three samples were collected from the eastern cross-section near the La Bocana locality and represent different structural levels: AV-08-28d (Fig. 9), AV-08-32a (Fig. 10) and AV-08-33 (Appendix C. Fig. 3) (see position in Fig. 4). The fourth sample, PU-08-10 (Appendix C. Fig. 4), which is located in the same upper structural level as sample AV-08-27, belongs to the western cross-section. The bulk compositions used in the calculations are presented in Table 7. All pseudosections were made with the assumption that the bulk composition of our samples during garnet crystallization did not significantly change. However, melt loss or gain is a widely accepted feature of migmatites (e.g., Brown, 2002, 2007; Kriegsman, 2001; White and Powell, 2002) that could strongly influence the bulk composition, and consequently, the mineral assemblage of the rocks. In order to investigate the effect of melt transfer, we considered $T-X_{melt}$ pseudosections (Appendix C. Fig. 2). $T-X_{melt}$ pseudosections show the characteristic migmatitic mineral assemblage field, the positions of X_{Alm} , X_{Grs} , X_{Mg} of biotite and X_{Ab} isopleths and thus peak P-T conditions estimates at temperature above 690 °C, are adversely affected by either loss or gain of melt.

The supra-solidus domain, between the stability field of kyanite and cordierite (Figs 9 & 10), is characterized by the mineral assemblage observed in the lower La Bocaca unit: biotite + garnet + plagioclase + K-feldspar + sillimanite + quartz. Pressure-T conditions were refined by contouring the X_{Alm} and X_{Grs} of garnet, X_{Mg} of biotite and X_{Ab} of plagioclase on each P-T pseudosections.

In sample AV-08-28d, garnet is modeled in all the P-T range and muscovite is absent of sub-solidus-region. The absence of muscovite in subsolidus domain can be related to extraction of low density melt produced during associated muscovite dehydration melting. P- T_{max} equilibration of garnet (Fig. 9) is of 7 ± 2 kbar and 760 ± 50 °C $(X_{Alm} = 0.73, X_{Grs} = 0.05)$. Biotite equilibration (Fig. 9) is likely to have occurred at higher temperature ranging from 800 to 840 °C $(X_{Mg} = 0.49$ to 0.59), compatible with plagioclase composition $(X_{Ab} = 0.67)$ in schlieren. This apparent discrepancy which is also observed in AV-08-32 and AV-08-33 samples (see Fig. 10 and Figs 3 & 4 in Appendix C.) can be explained either by requilibration of garnet during crystallization of residual melt, or by the difficulty to estimate active bulk composition during partial melting due to inherited metastable phases as Ti-bearing phases.

The stability of Ti-bearing oxide provides a good constraint for pressure. Rutile-bearing assemblages that have been observed in the lowest structural domains (e.g., sample AV-08-33) are restricted to pressure conditions above 7.0 kbar (Fig. 3 in Appendix B). Whereas Ti-bearing phases have been observed in all examined samples, in modeled pseudosections Ti-bearing phases, in suprasolidus conditions, are restricted to domains of high temperature (>780-800 °C). Pseudosections model theorical mineral assemblages at each P-T conditions without taking into account metastability or heterogenous reactions during metamorphism. In order to model the metastable Ti-bearing phases in suprasolidus domain we artificially saturated each pseudosection in Ti and reported the results over the original pseudosections (Figs. 9 & 10).

Results of pseudosections show that $P-T_{max}$ conditions in garnetbearing migmatites rise from 750 ± 50 °C for PU-08-10 to 750-820 °C for AV-08-28d, AV-08-32a and AV-08-33 while pressure increase continuously from 6 to 8 kbar for AV-08-28d. AV-08-32a and AV-08-33 and is ~7 kbar for PU-08-10.

7. Geochronological results

U-Th-Pb data obtained by SHRIMP at the University of Western Australia, Perth, Australia, and LA-ICPMS methods at the University

Table 8	
Solution and end-members used for the pseudosections (see Figs. 7 & 8). See Perple_X documentation (http://www.perplex.ethz.ch) for more de	tailed informations.

Phase	Solution model label in perpleX	End members	References
Chlorite	Chl(HP)	Clinochlore-daphnite-amesite-Al free chlorite	Holland and Powell, 1998
Plagioclase	Pl(h)	High albite-anorthite	Newton et al., 1980
Garnet	Gt(HP)	Spessartine-grossular-pyrope-almandine	Holland and Powell, 1998
Cordierite	hCrd	Fe-cordierite–Mn-cordierite–cordierite	Holland and Powell, 1998
Melt	melt(HP)	H ₂ O liquid–fayalite liquid–forsterite liquid–albite liquid–anorthite liquid–K-feldspar liquid–sillimanite liquid–Silica liquid	Holland and Powell, 1998
Orthopyroxene	Opx(HP)	Enstatite-ferrosilite	Holland and Powell, 1998
White-mica	Mica(CHA)	Muscovite-celadonite-Fe-celadonite-paragonite-margarite	Holland and Powell, 1998
Staurolite	St(HP)	Fe-staurolite-Mg-staurolite-Mn-staurolite	Holland and Powell, 1998
Biotite	Bio(TCC)	Mn-biotite–Ti biotite–annite–phlogopite–eastonite–Fe3 + Biotite	Tajcmanová et al., 2009

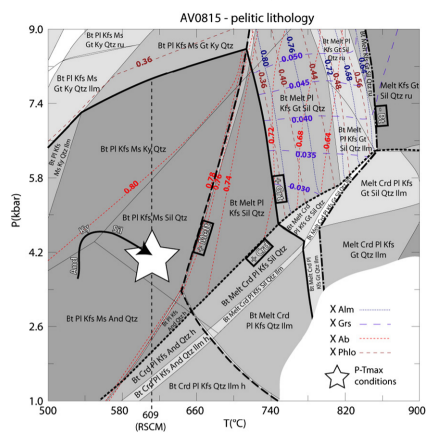


Fig. 8. *P*–*T* pseudosection of AV-08-15 metapelite in the NCKFMASHT system. The white star corresponds to $P-T_{max}$ metamorphic conditions underwent by the sample. These conditions have been estimated using the observed paragenesis and RCSM thermometry (vertical dotted line passing through the star center). Estimated conditions are supported by the compositions of biotite and plagioclase. The heavy black arrow represents the suggested P-T path before $P-T_{max}$ conditions based on aluminosilicate sequence of Fig. 5a & b. Bulk compositions used for each pseudosection are reported in Table 7. No shading = field divariant, then light to dark shading = increasing the variance (trivariant, quadrivariant, etc.). AV-08-15 sample represents the pelitic endmember lithology which in the lower the La Bocana unit underwent partial melting under biotite dehydration melting. In order to validate this hypothesis, X_{Alm} , X_{Crs} isopleths of garnet and X_{Phlo} , X_{Ab} of biotite and plagioclase are also indicated in the suprasolutis region. Reported isopleth of solid-solutions in suprasolidus region are compatible with measured compositions in samples AV-08-28d, AV-08-32 and PU-08-10 (X_{Alm} , 0.72 to 0.80; X_{Crs} , 0.025 to 0.045; X_{Phlo} , 0.5 to 0.6, X_{Ab} , 0.6 to 0.7) and also indicate that partial melting occurred under biotite dehydration melting at *P*-*T* conditions of 5 to 8 kbar and 740 to 820 °C. Same calculations were performed for AV-08-13 sample which represents the paramitic end-member lithology observed in the La Victoria unit. Results are presented in supplementary data and show similar results.

of Clermont-Ferrand, France, on zircon and monazite are presented in Tables 9 and 10, in Fig. 11 and Appendix A.

7.1. Monazites

Ten subhedral and eighteen anhedral monazite grains were analysed *in situ* in thin sections by LA-ICMPS, for the PU-08-10 and VI-08-12 garnet-bearing migmatitic samples of the La Bocana unit, respectively (Fig. 11a & b). They are compositionally homogeneous as no zoning was observed on backscattered images. Monazite 208 Pb/ 232 Th ages for sample PU-08-10 range from 228.9 ± 4.7 to 219.9 ± 4.5 and give a mean age of 223.2 ± 2.2 Ma (Fig. 11a). No inherited monazite grains were dated. Monazite 208 Pb/ 232 Th ages for VI-08-12 range from 229 ± 7 to 219 ± 5 Ma and give a mean age of 226.0 ± 1.3 Ma (Fig. 11b). Analyzed grains 1 and 6 of VI-08-12 sample exhibit a Th/U ratio of 0.6, and 208 Pb/ 232 Th ages of 235.7 ± 14.5 Ma and 236 ± 19.5 Ma (Table 9). The Th/U ratio (0.6) is consistent with xenotime rather than monazite. 208 Pb/ 232 Th dating on another xenotime grain yielded an older age (362.5 ± 12.3 Ma) with higher error and was considered in the mean age calculation.

7.2. Zircon

Fourteen zircon grains of AV-08-31 and ten zircon grains of AV-08-28d garnet-bearing migmatitic samples of the La Bocana unit were imaged and dated *in situ* by SHRIMP (Fig. 11c & d). The grains

are subhedral to rounded, and are usually composite with a core surrounded by multiple rims. Most of the crystal cores are detrital and characterized by a corroded outline. These cores exhibit a magmatic Th/U ratio (>0.1) and yield ²⁰⁶Pb/²³⁸U ages ranging from late Archean to Ordovician. Rims are characterized by metamorphic Th/U ratio (0.01 to 0.1) and are concordant. U contents are lower in the inherited core (90–120 ppm) than in the rim (230–280 ppm). ²⁰⁶Pb/²³⁸U ages on zircon rims from sample AV-08-31 range from 236 ± 3 to 224 ± 3 Ma, with a mean age of 229.3 ± 2.4 Ma (Fig. 11c). Rim ²⁰⁶Pb/²³⁸U ages of AV-08-28d range from 221 ± 4 Ma to 229 ± 3 Ma, with a mean age of 225.7 ± 6.5 Ma (Fig. 11d). No systematic ages were obtained on multiple overgrowths. However, ages obtained from the rims yielded undistinguishable Late Triassic ages within the analytical uncertainty.

8. Discussion

Petrological, thermobarometric and geochronological informations acquired in this study are combined to reconstruct the thermal structure and the possible evolution of the south-western Ecuadorian margin during Late Triassic times.

8.1. Geochronological implications

The U–Th–Pb *in situ* ages on garnet-bearing migmatites of the La Bocana unit are coherent with, but more precise than, the age of

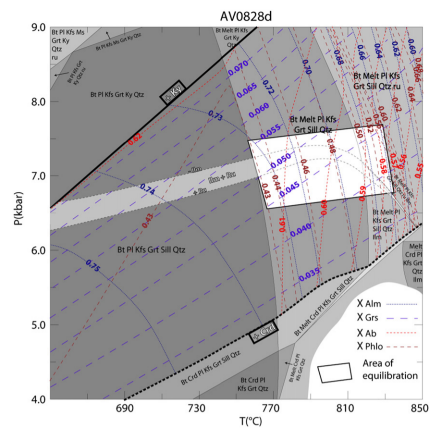


Fig. 9. *P*–*T* pseudosections of garnet-bearing migmatite AV-08-28d in the NCKFMASHT system. Isopleths of almandine (*X*_{*Alm*}),grossular (*X*_{*Grs*}), phlogopite (*X*_{*Phb*}) and albite (*X*_{*Ab*}) are plotted in the field of the observed peak *P*–*T* mineralogical assemblage; the white box represent the *P*–*T* conditions of garnet, biotite and plagioclase equilibration i.e., when measured composition matches the theoretical composition constructed by isopleths. Bulk compositions used for each pseudosection are reported in Table 7. Same shading as Fig. 8.

 219 ± 22 Ma obtained by a Sm/Nd whole-rock/garnet isochron on similar migmatites (Aspden et al., 1992a, 1992b). The Piedras unit has been dated at 221 ± 17 Ma by the U-Pb method on magmatic zircons (Noble et al., 1997), and at 226 ± 1.8 Ma by the Ar/Ar method on hornblende (Gabriele, 2002). These two ages constrain the age of emplacement and crystallization of the Piedras gabbroic unit before 226 ± 1.8 Ma. The migmatitic ages obtained in this study (235-220 Ma) are compatible with a single Triassic thermal event producing the Piedras gabbroic unit and the Marcabeli S-type granitoid unit (227.5 \pm 0.8 Ma, Noble et al., 1997). The Ar/Ar age of 225.3 \pm 1.7 Ma on amphibole (Gabriele, 2002) for the Arenillas-Panupalí blueschist unit records either the cooling of the rock below $550\pm$ 50 °C isotherm (Harrison and McDougall, 1981), or mineral crystallized below 550 ± 50 °C. The peak *P*–*T* conditions of the Arenillas–Panupalí unit have been estimated at 9 kbar and 300 °C (Gabriele, 2002), therefore the age at 225.3 ± 178 Ma dates the timing of the blueschist facies metamorphism. This age is statistically identical to the thermal event. However, the Arenillas-Panupalí blueschist unit, which was retrogressed under greenschist facies conditions (Gabriele, 2002), did not undergo any high temperature overprint. Therefore, underplating of the Arenillas-Panupalí oceanic unit postdated the thermal event. This interpretation is supported by the lack of zoning in migmatite garnets, which precludes a long cooling period. The timing of underthrusting of the Arenillas-Panupalí blueschist unit has important implications for the interpretation of the tectonic setting. The only setting to create blueschists is a subduction context (Ernst, 1988), and their exhumation mainly occurs in forearc regions (Cloos and Shreve, 1996; Guillot et al., 2009). Consequently, the age similarity for the whole Late Triassic metamorphic belt of El Oro massif implies that subduction was the only possible geodynamical setting that could produce anatexis and subsequent blueschist underplating.

8.2. P-T conditions

Estimates of P-T conditions in a metamorphic complex are of primary interest and are highly dependent on the geodynamical setting. The continental sequence of the El Oro metamorphic complex has been interpreted either as a regional shear zone undergoing partial melting (Aspden et al., 1995), or as a tilted continental sequence (Gabriele, 2002). The increase in metamorphic grade to the north from unmetamorphosed sediments to high-grade migmatites has been interpreted either as resulting from the closing from the shear zone (Aspden et al., 1995), or as due to normal deepening in the crustal sequence (Gabriele, 2002). Peak P-T data are compiled in Fig. 12; and our results show an excellent correlation between increasing temperature and pressure conditions toward the north. This is supported by the typically observed southward escape path of melt within dilatant structures, which must have been originally upward. Consequently the present geometry of the El Oro metamorphic complex is not representative of that in the Late Triassic. As proposed by Gabriele (2002) on the basis of the increasing temperature conditions, the continental sequence has been tilted to acquire its present subvertical. Because the Early Cretaceous Raspas eclogitic complex also exhibits a subvertical (Aspden et al., 1995; Gabriele, 2002), tilting must have occurred during or after exhumation of the Raspas complex, i.e., in the Early Cretaceous or later. The unconformity of the gently dipping Celica-Lancones sediments (Jaillard et al., 1999) on the vertical La Victoria unit implies that tilting occurred before Albian times. The present-day geometry has to be rotated ~90° along an east-west horizontal axis in order to reconstruct the original geometry during the thermal event. Rotation of the present day transtensive dextral movement indicates a low angle top-tothe-east or top-to-the-west normal movement during migmatization.

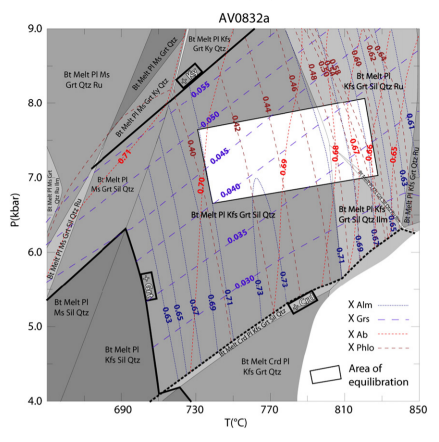


Fig. 10. P-T pseudosections of garnet-bearing migmatite AV-08-32a in the NCKFMASHT system. Isopleths of almandine (X_{Alm}), grossular (X_{Grs}), phlogopite (X_{Phlo}) and albite (X_{Ab}) are plotted in the field of the observed peak P-T mineralogical assemblage; the white box represent the P-T conditions of garnet, biotite and plagioclase equilibration; i.e., when measured composition matches the theoretical composition constructed by isopleths. Bulk compositions used for each pseudosection are reported in Table 7. Same shading as Fig. 8.

Table 9
LA-ICP-MS U–Pb analytical data for migmatitic rocks of La Bocana unit.

		Pb	Th	U			2σ error		2σ error	Age (Ma)	2σ error	
Sample	Analysis_#	(ppm)	(ppm)	(ppm)	Th/U	²⁰⁸ Pb/ ²³² Th	²⁰⁸ Pb/ ²³² Th	²⁰⁶ Pb/ ²³⁸ U	²⁰⁶ Pb/ ²³⁸ U	²⁰⁸ Pb/ ²³² Th	²⁰⁸ Pb/ ²³² Th	Position
VI-08-12												
	1	20	344	553	0.6	0.01173	0.00072	0.03728	0.00138	235.7	14.5	Core
	2	460	40053	1726	23.2	0.01077	0.00026	0.03713	0.00110	216.5	5.0	Core
	3	492	40421	1951	20.7	0.01123	0.00026	0.03852	0.00106	225.7	5.1	Core
	4	494	45777	782	58.5	0.01141	0.00026	0.03770	0.00114	229.2	5.2	Core
	5	660	54308	3344	16.2	0.01114	0.00026	0.03878	0.00108	224.0	5.0	Core
	6	32	507	917	0.6	0.01179	0.00098	0.03503	0.00152	236.9	19.5	Core
	7	599	51791	1969	26.3	0.01135	0.00024	0.03612	0.00092	228.1	4.8	Core
	8	645	53487	2881	18.6	0.01135	0.00024	0.03658	0.00092	228.1	4.8	Core
	9	783	52372	8674	6.0	0.01118	0.00024	0.03546	0.00086	224.6	4.7	Core
	10	660	47882	5640	8.5	0.01124	0.00024	0.03666	0.00090	225.9	4.7	Core
	11	56	601	987	0.6	0.01810	0.00062	0.05861	0.00158	362.5	12.3	Core
	12	545	45829	2298	19.9	0.01132	0.00024	0.03599	0.00092	227.6	4.8	Core
	13	729	62434	2600	24.0	0.01115	0.00024	0.03844	0.00114	224.1	4.8	Core
	14	69	5819	213	27.4	0.01139	0.00034	0.03698	0.00284	229.0	6.9	Core
	15	516	42647	2440	17.5	0.01134	0.00024	0.03504	0.00088	228.0	4.8	Core
	16	606	52860	2545	20.8	0.01109	0.00024	0.03491	0.00090	223.0	4.7	Core
	17	552	44673	2876	15.5	0.01104	0.00024	0.03910	0.00100	221.8	4.6	Core
PU-08-10												
	1	574	45262	3778	12.0	0.01123	0.00022	0.03560	0.00084	225.8	4.6	Core
	2	651	45939	6723	6.8	0.01095	0.00022	0.03531	0.00084	220.2	4.5	Core
	3	649	46894	6276	7.5	0.01094	0.00022	0.03581	0.00084	219.9	4.5	Core
	4	633	48186	5208	9.3	0.01104	0.00022	0.03535	0.00084	221.8	4.5	Core
	5	574	45428	3680	12.3	0.01107	0.00022	0.03602	0.00088	222.5	4.5	Core
	6	465	38568	2501	15.4	0.01096	0.00022	0.03591	0.00090	220.4	4.5	Core
	7	535	32502	6709	4.8	0.01139	0.00024	0.03546	0.00084	228.9	4.7	Core
	8	497	30691	6063	5.1	0.01124	0.00024	0.03642	0.00088	226.0	4.6	Core
	9	851	53582	1688	31.8	0.01124	0.00024	0.03697	0.00092	225.9	4.6	Core
	10	868	48776	2888	16.9	0.01100	0.00022	0.03692	0.00092	221.2	4.5	Core

Table 10
SHRIMP U–Pb analytical data for migmatitic rocks of La Bocana unit.

		Th	U			$2\sigma\text{error}$		2σ error		$2\sigma\text{error}$		2σ error	Age (Ma)	2σ error	
Sample	Analysis_#	(ppm)	(ppm)	Th/U	²⁰⁷ Pb/	²⁰⁸ Pb/	²⁰⁶ Pb/	²⁰⁶ Pb/	²⁰⁷ Pb/	²⁰⁷ Pb/	²⁰⁸ Pb/	²⁰⁸ Pb/	²⁰⁶ Pb/	²⁰⁶ Pb/	Position
					²⁰⁶ Pb	²³² Th	²³⁸ Pb	²³⁸ Pb	²³⁵ Pb	²³⁵ Pb	²⁰⁶ Pb	²⁰⁶ Pb	²³⁸ U	²³⁸ U	
AV-08-31															
	1	27	262	0.1	0.0507	5.9	0.037	1.4	0.29	10.2	0.053	6.9	233	3	rim
	2	19	297	0.1	0.1248	3.5	0.036	2.5	0.28	32.4	0.027	2.8	226	6	core
	3	19	278	0.1	0.0501	6.2	0.036	1.5	0.27	14.4	0.035	8.6	230	3	rim
	4	189	119	1.6	0.0630	5.2	0.107	1.3	0.94	7.4	0.515	1.6	653	8	core
	5	12	219	0.1	0.0479	6.6	0.036	1.1	0.29	5.5	0.047	10.3	231	2	rim
	6	4	249	0.0	0.0505	6.0	0.035	1.1	0.19	10.3	-0.022	17.0	223	2	core
	7	16	236	0.1	0.0545	6.1	0.035	1.3	0.22	15.0	0.006	8.1	224	3	rim
	8	55	90	0.6	0.0650	4.7	0.129	1.3	1.16	4.9	0.200	2.8	782	9	core
	9	23	224	0.1	0.0494	6.3	0.036	1.1	0.31	5.1	0.067	7.3	231	2	rim
	10	174	624	0.3	0.0584	2.4	0.078	1.0	0.62	2.6	0.098	1.9	482	5	rim
	11	24	507	0.0	0.0602	2.6	0.095	2.7	0.80	3.9	0.018	4.6	583	15	core
	12	435	125	3.5	0.0622	4.4	0.095	1.3	0.91	8.0	1.114	1.2	584	7	core
	13	296	146	2.0	0.0631	4.1	0.094	2.3	0.81	4.7	0.650	1.4	577	13	core
	14	26	240	0.1	0.0505	6.1	0.036	1.7	0.25	16.3	0.038	7.0	229	4	core
	15	116	180	0.6	0.1143	1.3	0.513	2.7	8.06	3.0	0.162	8.4	2671	59	core
	16	137	582	0.2	0.1472	4.7	0.358	1.0	7.27	4.8	0.071	6.2	1974	17	core
	17	27	283	0.1	0.0519	5.1	0.037	1.3	0.26	5.3	0.038	6.1	232	3	rim
	18	55	450	0.1	0.0496	4.5	0.036	1.0	0.25	4.6	0.038	5.2	228	2	rim
	19	89	172	0.5	0.0492	6.3	0.037	1.1	0.24	7.5	0.155	3.6	236	3	core
	20	28	309	0.1	0.0513	5.0	0.036	1.3	0.26	5.1	0.033	6.2	231	3	rim
AV-08-28d															
	1	26	302	0.1	0.0572	3.9	0.052	5.3	0.41	6.6	0.032	5.2	329	17	rim
	2	391	377	1.0	0.0704	2.1	0.169	3.7	1.64	4.2	0.322	4.9	1009	34	core
	3	115	286	0.4	0.0653	2.9	0.087	1.5	0.78	3.3	0.132	4.9	538	8	rim
	4	50	1745	0.0	0.0561	1.7	0.072	2.0	0.56	2.6	0.008	7.3	447	9	core
	5	62	255	0.2	0.0645	5.1	0.085	3.7	0.76	6.3	0.074	3.7	525	19	core
	6	84	460	0.2	0.0545	3.7	0.040	1.9	0.30	4.1	0.056	3.7	252	5	core
	7	18	242	0.1	0.0516	5.7	0.035	1.7	0.25	6.0	0.021	9.2	221	4	rim
	8	77	186	0.4	0.0496	5.8	0.089	2.9	0.60	6.5	0.116	2.6	547	15	core
	9	11	203	0.1	0.0546	6.0	0.035	1.5	0.26	6.2	0.019	10.3	223	3	rim
	10	26	57	0.5	0.1926	4.5	0.538	1.8	14.28	4.9	0.140	2.1	2774	41	core
	11	22	591	0.0	0.0562	2.7	0.071	2.6	0.55	3.7	0.011	4.8	443	11	rim
	12	31	74	0.4	0.0777	4.0	0.164	1.7	1.75	4.4	0.140	3.1	977	15	core
	13	9	213	0.0	0.0499	14.4	0.036	1.7	0.25	14.5	0.012	10.9	226	4	rim
	14	45	78	0.6	0.0455	5.8	0.087	2.7	0.68	6.4	0.188	3.3	536	14	core
	15	40	383	0.0	0.0496	4.6	0.036	1.5	0.08	4.8	0.033	5.2	228	3	rim
	16	21	585	0.1	0.0490	20.4	0.030	3.6	1.47	20.7	0.033	3.7	863	29	core
	10	21	238	0.4	0.0744	3.8	0.143	2.3	0.62	4.5	0.114	3.6	451	10	rim
	17	16	238	0.1	0.0021	5.4	0.072	1.5	0.02	5.6	0.073	8.1	229	3	rim
	10	10	245	0.1	0.0492	J.4	0.050	1.J	0.23	5.0	0.025	0.1	223	ر	11111

Extension during Late Triassic times is supported by our mineralogical study: relict rutile in ilmenite marks decompression before garnet crystallization and observed kyanite recrystallizing into sillimanite during migmatization. Note that extension is assumed to have occurred in north-western South America at that time (e.g., Mišković et al., 2009; Vinasco et al., 2006).

P-T estimates (Fig. 12) are direct proxies of the maximum geotherm during the early Late Triassic thermal anomaly. The pre-anatectic mineral assemblage in the La Victoria unit is recorded in the sample AV-08-15 by andalusite statically re-equilibrated into kyanite. The thermal event is marked by a strong temperature increase and active deformation in all the studied samples marked by sillimanite metamorphic foliation. It defines a hot maximum geothermal gradient (Fig. 12). This Late Triassic maximum geotherm exhibits two segments. In the upper unit, the geotherm shape is convex with high dT/dP, whereas in the lower unit, the geotherm shape is linear, almost vertical, with very low dT/dP. The migmatitic layer is of about 10 km thick. In the upper part, in the 2.5 km thick white-mica dehydration melting zone, temperature rises from sub-solidus conditions to 750 °C, thus representing a thermal gradient of 45 °C/km (down to 6 kbar) (Fig. 12). In contrast, temperature estimates in the garnet-bearing migmatites of La Bocana (7 km thick) are rather constant, ranging from 750 to 820 °C, which exhibits a thermal gradient of 10 °C/km. However, in this part pressure increases northward from 6 to 8 kbar (Fig. 12). Taking into account both melt modal percent and melt geometry we suggest that in the lower zone, maximum temperature is buffered, in part, by latent heat of biotite melting (e.g., Depine et al., 2008; Thompson and Connolly, 1995). Although the isothermal gradient could be explained by melt buffering alone, the high amount of melt and the lack of oriented structures and magmatic fabric within the garnet-bearing migmatites, suggest that convective motion occurred throughout the lower migmatitic zone.

8.3. Heat source

Migmatization of the semi-pelitic sequence of the La Bocana unit, i.e., switching from a relatively cool to a high geothermal gradient, requires heat input into the system. Three main possible mechanisms can provide additional heat to the continental crust (Bergantz, 1992; Huppert and Sparks, 1988): (1) crustal thinning and isotherm tightening, (2) crustal thickening and enhanced radiogenic production, (3) emplacement of a plutonic body close to, or within the system. The very strong convex geothermal gradient for the upper El Oro units cannot be explained by thermal conduction alone. We suggest that this strong apparent geothermal gradient results from the combined effect of crustal thinning, and heat input in the La Bocana garnet-bearing migmatites. The mixing zone between the gabbroic magma of the Piedras unit and the acid magma of the La Bocana unit implies that the Piedras gabbroic unit is an *in situ* magmatic body emplaced during Triassic times (Aspden et al., 1995; Noble et al., 1997), and is not a tectonically underplated oceanic terrane, as suggested by Gabriele (2002) and Bosch et al. (2002). Hence, we propose that crustal anatexis occurred, due to a combination of crustal thinning, heat transfer from

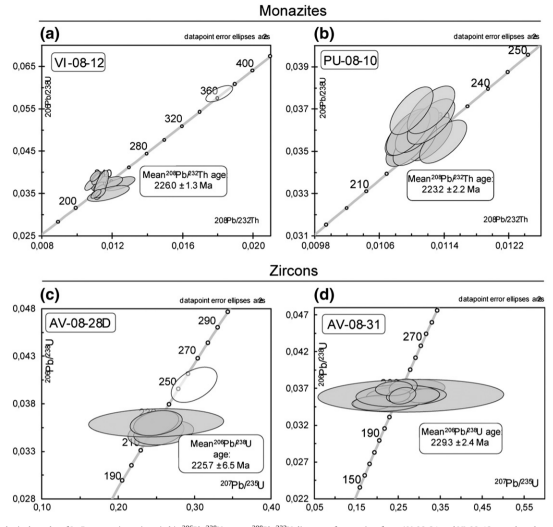


Fig. 11. Geochronological results of La Bocana migmatites. (a,b); ²⁰⁶Pb/²³⁸U versus ²⁰⁸Pb/²³²U diagram of monazites from AV-08-31 and VI-08-12 samples; the mean ²⁰⁸Pb/²³²U age is indicated; (c, d) Concordia diagram of zircons from AV-08-31 and AV-08-28d samples, the mean ²⁰⁶Pb/²³⁸U age is indicated.

the magmatic underplating of the Piedras unit, and latent heat of crystallization of the latter. Based on structural (Fig. 3a & b) and mineralogical evidence (rutile to ilmenite), the El Oro metamorphic complex underwent crustal thinning before and during migmatization. Even though crustal thinning likely played a role in the isotherms tightening, it was probably not the only heat source that caused the anatectic event.

In contrast the magmatic emplacement of the high temperature (>1000 °C) gabbroic Piedras unit before 226 ± 1.8 Ma at crustal root level (Noble et al., 1997) is likely to have provided the local heat source for migmatization (Aspden et al., 1995). This assumption is supported by the presence of a relatively thin magma mixing zone (~100 m), where gabbroic melts from the Piedras gabbroic unit and granitic melt from the La Bocana unit can be observed. If we can locally attribute the main heat source to the emplacement of the Piedras unit, the formation of the Piedras gabbro itself remains an open question. Based on the MORB origin of the Piedras gabbro (Aspden et al., 1995; Bosch et al., 2002), we propose that upwelling and subsequent decompression of the asthenosphere beneath the forearc zone were able to produce this MORB-like magma.

8.4. Geodynamical framework

Based on the similar ages of Tres Lagunas granite and Marcabeli granitoid, Noble et al. (1997) proposed that the El Oro metamorphic complex and the Cordillera Real of Ecuador (Fig. 1) were contiguous during the emplacement of S-type granites. Inherited core of zircons of sample AV-08-28d (Table 10) show two main peak at 550–525 Ma and 450–440 Ma. Those ages demonstrate that the protolith of the La Bocana migmatites is probably metasediments equivalent to the Isimanchi and Chiguinda Unit of the Cordillera Real in Ecuador, which are believed to be autochthonous to the Gondwana margin (Chew et al., 2007). Therefore these inherited ages support the idea that the El Oro metamorphic complex belonged to the Cordillera Real of Ecuador before Cretaceous tilting of the El Oro metamorphic complex and exhumation of the ophiolitic Raspas complex.

While partial melting of a significant part of a forearc continental crustal section is a rare feature in the Andes, evidence of a late Triassic tectono-thermal anomaly associated with granitic production are widespread all along the western South-American margin. From north to south, evidences of late Triassic granitoid emplacement are: the Amaga Granitic Stock in Colombia at 227.6 ± 3.2 Ma (U–Pb, Vinasco et al., 2006), Tres Lagunas peraluminous granite within the Cordillera Real of Ecuador (see Fig. 1) at 227.3 ± 2.2 Ma (U-Pb, Litherland et al., 1994; Noble et al., 1997), Sabanilla migmatite of northern Peru at 230.4 ± 3.2 Ma (Chew et al., 2008), Zongo granitoid in Bolivia at 225.1 \pm 4 Ma (U–Pb, Farrar et al., 1990), and Constitución granite in Chile at 224 ± 1 Ma (Pb–Pb, Willner, 2005a, 2005b).These authors agree that crustal thinning occurred during crustal melting and granitic intrusion. Farther East in Ecuador, ages at 245.7 ± 5.6 , 239.2 ± 2.2 , 234.2 ± 1.1 and 234.66 ± 0.95 Ma, from magmatic rims of zircons from acidic intrusive rocks the Loja Terrane and the El Oro metamorphic complex El Oro metamorphic complex, show the

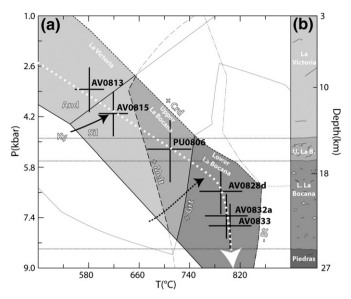


Fig. 12. *P*–*T* diagram for El Oro metamorphic complex. (a) reported peak *P*–*T*_{max} conditions of studied samples in the simplified pseudosection of AV-08-15 sample (Fig. 8). Fields in greyscale show the domain of stability of observed mineral paragenesis of the La Victoria, the Upper and Lower La Bocana units. Note that the sample AV-08-15 (Fig. 5c) indicates that the pre-anatectic thermal history is marked by the transition from andalusite to kyanite, suggesting burial of the La Victoria unit before the high-temperature event (black arrow). The white and dashed arrows present the maximum *P*–*T* condition recorded in the rock. We suggest that these last *P*–*T* conditions could represent the geothermal gradient during Triassic times. (b) Schematic structural log of the El Oro from the La Victoria unit to the South to the Piedras unit to the North.

presence of a belt of Triassic migmatites across the southern Ecuador. Comparable events involving migmatization of the crust and S-type magmatism, dated from 260 to ca. 226 Ma, are well known from Northern Colombia to Northern Peru (Cardona et al., 2010; Mišković et al., 2009; Restrepo et al., 2011) indicating a long period of S-type magmatism during late Permian to late Triassic times.

In Colombia and Ecuador, intrusions of the Amaga and Tres Lagunas granites were interpreted as related to the Tethyan rifting between North and northern South America (Aspden and Litherland, 1992). However, Cardona et al. (2010) showed that subduction-related magmatism was active until late Permian, thus North and northern South America were separated during Permian-Triassic times precluding a continental rifting model. Moreover, in a rifting model intense extension would have occurred to upwell the asthenosphere at shallow depth. In the La Victoria and the La Bocana units such intense deformation has not been observed. However, in Chile, Willner (2005b) interpreted the intrusion of the Constitución granite as indicating retreat of the subducting slab. More recently, Bustamante et al. (2011) showed that in the Central Cordillera of Colombia, amphibolitic schists related to subduction were tectonically juxtaposed with Triassic migmatites and granulites during a late Triassic subduction event. The presence of an underplated oceanic slab (Arenillas-Panupalí unit) at 225 Ma in the El Oro metamorphic complex and oceanic related underplated material in Colombia (Bustamante et al., 2011) imply that subduction was active at that time. Therefore, we interpret the thermal event affecting the Ecuadorian margin during Triassic times as the result of an anomaly arising from subduction processes.

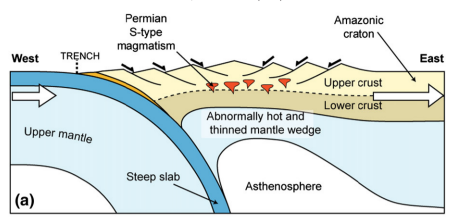
During Permo-Triassic times, at 270–250 Ma, South America underwent a major shift in drift direction from northward to eastward directed (Geunal et al., 2010; Tomezzoli, 2009; Torsvik et al., 2008). This change in plate motion resulted from the counter-clockwise rotation of Gondwana during that time and ended in Late Triassic times, when South America resumed its movement toward the north (Torsvik et al., 2008).

We propose that this major eastward migration of South America provoked a rapid trench retreat, and a generalized extensional regime associated with a steep to vertical slab (Fig. 13a). Slab verticalization, due to slab anchorage at depth, could have triggered dehydration of the slab and upwelling of the asthenosphere (Zhu et al., 2011). In this model, partial melting of the asthenosphere likely generated MORB-type identical to the Piedras gabbroic unit. Underplating of magma and melting of the crust would have started in arc and backarc position (Zhu et al., 2011) during late Permian to Triassic times and finally extended to the forearc position in late Triassic. Subsequent underplating at ~230 Ma of the Piedras gabbroic unit at crustal root level (Fig. 13b) in turn triggered migmatization of the semi-pelitic sequence of the La Bocana unit and emplacement of the Marcabeli S-type granitoid (Noble et al., 1997). At the same time (\approx 227 Ma), eastward migration of South America ceased (Torsvik et al., 2008), and we propose that slab breakoff occurred.

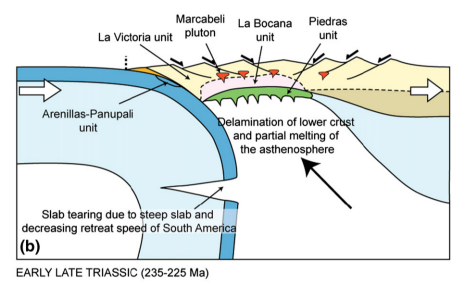
Based on U-Pb geochronology and geochemical study, Mišković et al. (2009) proposed that the Permian-Triassic magmatism in northwestern Gondwana from 260 to 220 Ma is related to a change in plate vector kinematics, resulting in thermal weakening of the subduction slab and eventually slab breakoff. They interpreted the subsequent upwelling of the asthenophere as the cause of basaltic magma emplacement at the base of the lower crust triggering a vast region of extensive crustal melting from 260 to 200 Ma along the western Gondwana. Our model is similar except that we propose that slab breakoff occurred when South America resumed its course northward after a period of ~30 Ma of progressive slab verticalization ultimately reaching the forearc regions. Slab breakoff likely provoked an isostatic rebound of the oceanic plate, thus accounting for the tectonic underplating of the Arenillas-Panupalí blueschist unit (Fig. 13c) at ~226 Ma and underplating of oceanic material in Colombia in late Triassic (Bustamante et al., 2011). The latter process would explain the rapid cooling of both the Piedras unit and the El Oro metamorphic complex (~226 Ma).

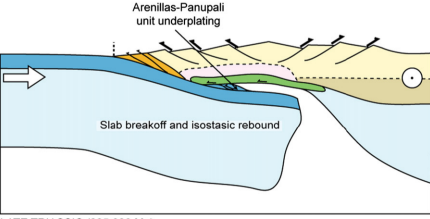
9. Conclusion

The results of our detailed thermobarometric study of the El Oro metamorphic complex, and the evolution of the Late Triassic paired metamorphic belt of southwestern Ecuador can be summarized as follows. After a period of relatively limited thinning of the continental crustal, melting of the Paleozoic psammite-pelite sequence of the La Bocana unit started at ~229 Ma. Crustal thinning and emplacement of the Piedras gabbroic unit provided the heat source that enhanced crustal anatexis. Melting of the metasediments occurred through white-mica and biotite dehydration processes. In the biotite dehydration melting zone, garnet was produced as the main peritectic phase, while modal percentage of biotite and sillimanite decreased with increasing metamorphic conditions. P-T conditions estimates showed that crustal anatexis started at about 650 °C and ~4.5 kbar, and reach maximum of 820 °C. On a 7 km thick layer the apparent geotherm was of about 40 °C/km in the upper crustal unit and 5 °C/km in the lower unit. In the lower unit we suggest that temperature was buffered by biotite dehydration melting and by convective motions of magmas. The occurrence of numerous granite intrusions during Triassic times along the South American margin indicates that western South American underwent a widespread thermal anomaly between 230 and 225 Ma, which produced large amounts of S-type granitoids. We attribute the tectono-thermal event to a slab break-off event due to eastward migration of South America during Permian-Triassic times. Slab breakoff and/or slab verticalisation provoked upwelling and decompression of the asthenosphere, which underwent partial melting and provoked the magmatic underplating of the MORB-type Piedras gabbroic unit. This tectono-thermal event was immediately followed by renewed subduction processes and the underplating of the Arenillas-Panupalí blueschist unit, which strongly contributed to the rapid cooling down of the forearc region of southwestern Ecuador by ~226 Ma.



EARLY TO MIDDLE TRIASSIC (250-235 Ma)





LATE TRIASSIC (225-220 Ma)

Fig. 13. Schematic proposed geodynamical model of the Ecuadorian margin at Triassic times.(a) middle Triassic period, south America is moving eastward and the margin is affected by S-type magmatism likely due to a steep to vertical slab; (b) early Late Triassic period, partial melting of the asthenosphere produces the emplacement of the MORB-type gabbro of the Piedras unit at crustal root level, and subsequent migmatization of the La Bocana unit; (c) shortly after slab breakoff, subduction resumes and the Arenillas–Panupalí blueschist unit is underplated.

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Appendix A. Analytical techniques

Raman study of carbonaceous material

We used Raman spectroscopy on carbonaceous matter (RSCM) in metasediments of middle to high metamorphic grade. The Raman spectra of carbonaceous matter are characterized by three bands. The band centered at ~1580 cm⁻¹ is interpreted as the G band and corresponds to the vibration mode of aromatic carbon in the graphitic structure. The band centered at ~1350 cm⁻¹ is interpreted as the D1 band and is attributed as the presence of heteroatoms (O. H. N) or defects in the structure. The third band centered at ~1620 cm⁻¹ is the D2 band and is attributed to out-plane defects (Bény-Bassez and Rouzaud, 1985). Graphitization is a process observed in metamorphic rocks and corresponds to the solid-state transformation of organic matter into carbonaceous mineral. Graphitization is thermally dependent and can be used a geothermometer (Beyssac et al., 2002). This geothermometer is based on the estimation of the area ratio R2 calculated from the three characteristic bands (D1. D2 and G).

In this study Raman spectroscopy was performed at the Ecole Normale Supérieure de Lyon, France, with a Horiba Jobin-YvonLabRam apparatus. The light source was an argon-ion laser to achieve 514.5 nm wavelengths. An OlympusTM BX30 open microscope equipped with a \times 50 long-working distance objective was coupled to a spectrometer to focus the laser output power on the sample to a 3 µm spot diameter. The Raman signal was collected in the backscattered direction. Acquisition time was 90 s distributed over three accumulation cycles. with a laser output power on the sample of 0.7 mW. The spectral resolution was 1 cm⁻¹ using 600 g/mm grating. The spectral region from 1200 to 1800 was investigated in order to characterize the vibration mode of carbonaceous matter following the method proposed by Beyssac et al. (2002). Treatments of the RSCM data were undertaken using Peakfit software.

Microprobe

Mineral chemistry has been studied by microprobe analysis on thin-sections, using a CAMECA SX 100 microprobe at the Laboratoire Magma et Volcans of the university of Clermont-Ferrand) with 15 kV acceleration potential and 15 nA beam current. Representative analyses of minerals are presented in Tables 1 and 2. Acquisition of the garnet map of the Fig. 8 was done on the same microprobe, with 15 kV acceleration potential, 100 nA beam current and 300 ms counting time.

Whole-rock chemistry

Whole-rock analyses were performed at ISTerre (University of Grenoble. France) and are presented in Table 4. Concentrations of major elements were determined by ICP-AES using a Perkin Elmer 3000 DV spectrometer using the method of Cotten et al. (1995).

Geochronology

Monazite geochronology

U–Th–Pb geochronology of monazite was conducted by laser ablation inductively coupled plasma spectrometry (LA-ICPMS) at the university of Clermont-Ferrand, on thin sections. The analyses involve the ablation of minerals with a Resonetics Resolution M-50 powered by an ultra short pulse ATL Atlex Excimer laser system operating at a wavelength of 193 nm (detailed description in Müller et al., 2009). Spot diameters of 7 µm with repetition rates of 3 Hz and 1 Hz were used. The ablated material is carried into helium, and then mixed with nitrogen and argon before injection into a plasma source of an Agilent 7500 cs ICP-MS equipped with a dual pumping system to enhance the sensitivity.

The alignment of the instrument and mass calibration was performed before every analytical session using the NIST SRM 612 reference glass. by inspecting the signal of ²³⁸U and by minimizing the ThO⁺/Th⁺ ratio (<<1%). The mean sensitivity on ²³⁸U at the instrumental conditions reported in Table 7 and using a spot size of 44 µm is about 15-20,000 cps/ppm. The analytical method for isotope dating of monazite with laser ablation ICPMS is basically similar to that developed for zircon and monazite and reported in Tiepolo (2003) and Paquette and Tiepolo (2007). The signal of the $^{204}(Pb + Hg)$. ^{206}Pb . ²⁰⁷Pb. ²⁰⁸Pb. ²³²Th and ²³⁸U masses are acquired. The occurrence of common Pb in the sample can be monitored by the evolution of the 204 (Pb + Hg) signal intensity, but no common Pb correction was applied owing to the large isobaric interference from Hg. The ²³⁵U signal is calculated from 238 U on the basis of the ratio 238 U/ 235 U = 137.88. Single analyses consisted of 30 s of background integration with the laser off followed by 1 min integration with the laser firing and a 30 s delay to wash out the previous sample (approximately 10 s for a 6 order of magnitude decrease in signal intensity) and prepare the next analysis.

Data are corrected for U–Pb and Th–Pb fractionation occurring during laser sampling and for instrumental mass discrimination (mass bias) by standard bracketing with repeated measurements of the Moacir monazite standard (Gasquet et al., 2010). At the beginning and at the end of every run, repeated analyses of the Manangoutry monazite (Paquette and Tiepolo, 2007) standard were treated as unknowns to independently control the reproducibility and accuracy of the corrections. Data reduction was carried out with the software package GLITTER® developed by the Macquarie Research Ltd. (Jackson et al., 2004; van Achterbergh et al., 2001). For each analysis, the time resolved signal of single isotopes and isotope ratios was monitored and carefully inspected to verify the presence of perturbations related to inclusions, fractures, mixing of different age domains or common Pb. Calculated ratios were exported and Concordia ages and diagrams were generated using Isoplot/Ex v. 2.49 software package of Ludwig (2003). The concentrations in U-Th-Pb were calibrated relative to the certified contents of 91500 zircon standard (Wiedenbeck, 1995) and Moacir monazite (Seydoux-Guillaume et al., 2004).

Zircon geochronology

Zircon grains were handpicked from the concentrate. Crystals were mounted in epoxy resin with several chips of the BR266 standard zircon standard (903 ppm U corresponding to an age of 559 Ma) on 25 mm diameter mounts and polished to expose zircon cross-sections. The polished mounts were carbon-coated prior to SEM examination. Zircons were imaged using a JEOL 6400 Scanning Electron Microscope at the Centre for Microscopy and Microanalysis at the University of Western Australia. Backscattered electron and cathodoluminescence (CL) images were obtained for all the mounts, and were used to characterize each grain in terms of size growth morphology and internal structure. The carbon coat was removed and the mount gold-coated for SHRIMP analysis.

Isotope analyses were performed using the SHRIMP II ion microprobe at Curtin University of Technology, Western Australia using the standard operating conditions described by Compston et al. (1992) and Smith et al. (1998). The spot size used during all the sessions was around 20 μ m. In this study, six data collection cycles (scans) were performed per analysis and count times (per scan) were 10 s for the ²⁰⁴Pb. ²⁰⁶Pb and ²⁰⁸Pb mass peaks and background and 30 s for the ²⁰⁷Pb mass peak. Unknown analyses were referenced to multiple analyses of the BR266 standard for U/Pb calibration. Standard analyses performed throughout each analysis session gave 1 σ uncertainties of 2.0% or less for Pb*/U Data reduction was carried out with the software package SQUID v1.10 and ISOPLOT v3.0 (Ludwig, 2003). Corrections for common Pb content were based on measured ²⁰⁴Pb and assuming contemporary values for common Pb derived from Stacey and Kramers (1975).

Appendix B. Microprobe composition of garnet profiles

Table 1

Electron microprobe analysis of a garnet profile of AV0828d sample.

	AV0828d	AV0828d	AV0828d	AV0828d	AV0828d	AV0828d	AV0828d	AV0828d	AV0828d	AV0828d	AV0828d	AV0828d	AV0828d	AV0828d	AV0828d	AV0828d	AV0828d	AV0828d	AV0828d	AV0828d
SiO ₂	38.34	38.34	38.20	41.15	38.41	38.39	38.71	38.64	39.62	38.58	38.58	38.49	38.51	38.70	39.66	36.91	36.54	36.46	36.36	36.43
TiO ₂	0.03	0.03	0.00	0.01	0.03	0.07	0.05	0.03	0.03	0.01	0.02	0.03	0.02	0.04	0.00	0.03	0.06	0.10	0.07	0.09
Al_2O_3	20.76	20.76	20.78	24.39	20.74	20.60	20.62	20.56	21.15	20.74	20.60	20.81	20.80	20.73	21.08	20.11	20.04	20.15	20.14	19.71
FeO	27.91	27.91	28.25	20.84	28.19	28.54	28.49	28.89	28.37	28.88	28.43	28.85	28.43	28.39	27.89	26.68	26.82	26.48	26.87	26.44
MnO	6.98	6.98	6.91	4.30	7.01	7.04	7.09	7.00	6.96	7.01	7.05	7.12	7.10	7.29	7.11	14.47	14.27	14.07	14.40	14.32
MgO	3.66	3.66	3.59	2.95	3.58	3.64	3.54	3.56	3.46	3.55	3.56	3.54	3.52	3.41	3.26	1.20	1.17	1.22	1.22	1.22
CaO	1.50	1.50	1.42	1.16	1.47	1.44	1.35	1.35	1.35	1.40	1.40	1.37	1.43	1.49	1.99	0.64	0.69	0.74	0.80	0.75
Total	99.17	99.17	99.15	94.80	99.43	99.71	99.84	100.04	100.95	100.17	99.64	100.22	99.81	100.06	100.99	100.04	99.59	99.22	99.86	98.96
Numbers of io	ns on the b	asis of 22 ()																	
Si	3.07	3.07	3.06	3.30	3.08	3.08	3.10	3.10	3.18	3.09	3.09	3.08	3.09	3.10	3.18	2.96	2.93	2.92	2.91	2.92
Ti	0.00	0.00	0.00	0.00	0.00	0.00	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	0.01
Al	1.96	1.96	1.96	2.30	1.96	1.95	1.95	1.94	2.00	1.96	1.95	1.97	1.96	1.96	1.99	1.90	1.89	1.90	1.90	1.86
Fe	1.87	1.87	1.89	1.40	1.89	1.91	1.91	1.94	1.90	1.94	1.91	1.93	1.91	1.90	1.87	1.79	1.80	1.77	1.80	1.77
Mn	0.47	0.47	0.47	0.29	0.48	0.48	0.48	0.48	0.47	0.48	0.48	0.48	0.48	0.49	0.48	0.98	0.97	0.96	0.98	0.97
Mg	0.44	0.44	0.43	0.35	0.43	0.43	0.42	0.43	0.41	0.42	0.43	0.42	0.42	0.41	0.39	0.14	0.14	0.15	0.15	0.15
Ca	0.13	0.13	0.12	0.10	0.13	0.12	0.12	0.12	0.12	0.12	0.12	0.12	0.12	0.13	0.17	0.05	0.06	0.06	0.07	0.06
Total	7.94	7.94	7.94	7.74	7.96	7.97	7.98	7.99	8.08	8.01	7.97	8.01	7.98	8.00	8.08	7.83	7.79	7.77	7.81	7.74
Endmembers ((mol%)																			
Pyrope	15.0	15.0	14.7	16.5	14.7	14.7	14.4	14.4	14.2	14.4	14.5	14.3	14.4	13.9	13.4	4.8	4.7	5.0	4.9	4.9
Almandine	64.3	64.3	65.0	65.3	64.7	64.9	65.2	65.6	65.5	65.5	65.0	65.4	65.0	64.9	64.2	60.2	60.6	60.4	60.2	60.0
Grossular	4.4	4.4	4.2	4.6	4.3	4.2	3.9	3.9	4.0	4.1	4.1	4.0	4.2	4.4	5.9	1.8	2.0	2.2	2.3	2.2
Spessartine	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.1	0.1	0.1	0.1	0.1

Table 2

Electron microprobe analysis of a garnet profile of PU0810 sample.

Sample	PU0810	PU0810	PU0810	PU0810	PU0810	PU0810	PU0810	PU0810	PU0810	PU0810	PU0810	PU0810	PU0810	PU0810	PU0810	PU0810	PU0810	PU0810	PU081
SiO ₂	37.93	37.99	37.92	37.56	38.26	38.13	37.86	37.81	38.03	37.87	37.97	37.77	37.35	37.36	37.81	37.82	37.71	37.77	38.29
TiO ₂	0.00	0.04	0.10	0.07	0.13	0.10	0.11	0.10	0.02	0.05	0.00	0.11	0.52	0.10	0.08	0.00	0.08	0.08	0.04
Al_2O_3	20.79	20.95	20.76	20.81	20.86	20.95	20.83	20.56	21.00	21.26	20.67	20.70	20.02	20.99	21.01	21.21	20.77	21.14	21.40
FeO	34.89	35.31	35.15	34.67	35.11	36.04	35.22	35.85	35.31	35.69	35.85	35.76	35.21	35.65	36.49	35.83	35.52	35.75	36.52
MnO	2.48	2.58	2.30	2.23	2.19	2.01	1.83	1.63	1.46	1.15	1.18	1.01	1.00	0.98	0.96	0.94	0.87	0.84	0.84
MgO	3.40	3.44	3.59	3.59	3.65	3.70	3.87	4.00	4.13	4.14	3.98	4.17	3.94	4.21	3.98	4.20	4.04	4.17	4.18
CaO	0.81	0.92	0.96	0.90	0.92	0.83	0.95	0.90	0.85	0.88	0.78	0.83	0.93	0.85	0.87	0.92	0.90	0.91	0.96
Total	100.31	101.22	100.78	99.82	101.13	101.75	100.68	100.86	100.80	101.04	100.42	100.35	98.96	100.15	101.20	100.92	99.90	100.67	102.24
Numbers of ior	ns on the bas	sis of 12 O																	
Si	3.04	3.05	3.04	3.01	3.07	3.06	3.03	3.03	3.05	3.03	3.04	3.03	2.99	2.99	3.03	3.03	3.02	3.03	3.07
Ti	0.00	0.00	0.01	0.00	0.01	0.01	0.01	0.01	0.00	0.00	0.00	0.01	0.03	0.01	0.01	0.00	0.01	0.00	0.00
Al	1.96	1.98	1.96	1.97	1.97	1.98	1.97	1.94	1.98	2.01	1.95	1.95	1.89	1.98	1.98	2.00	1.96	2.00	2.02
Fe	2.34	2.37	2.36	2.32	2.35	2.42	2.36	2.40	2.37	2.39	2.40	2.40	2.36	2.39	2.45	2.40	2.38	2.40	2.45
Mn	0.17	0.17	0.16	0.15	0.15	0.14	0.12	0.11	0.10	0.08	0.08	0.07	0.07	0.07	0.07	0.06	0.06	0.06	0.06
Mg	0.41	0.41	0.43	0.43	0.44	0.44	0.46	0.48	0.49	0.49	0.48	0.50	0.47	0.50	0.48	0.50	0.48	0.50	0.50
Ca	0.07	0.08	0.08	0.08	0.08	0.07	0.08	0.08	0.07	0.08	0.07	0.07	0.08	0.07	0.07	0.08	0.08	0.08	0.08
Total	7.99	8.06	8.03	7.96	8.06	8.10	8.04	8.05	8.06	8.09	8.02	8.02	7.89	8.01	8.08	8.08	7.99	8.06	8.18
Endmembers (mol%)																		
Pyrope	13.6	13.6	14.2	14.4	14.5	14.4	15.3	15.6	16.3	16.3	15.7	16.4	15.8	16.6	15.5	16.5	16.1	16.4	16.2
Almandine	78.4	78.1	77.9	77.9	78.0	78.8	77.9	78.3	78.0	78.7	79.4	79.0	79.2	78.8	79.9	78.8	79.4	79.1	79.3
Grossular	2.3	2.6	2.7	2.6	2.6	2.3	2.7	2.5	2.4	2.5	2.2	2.3	2.7	2.4	2.4	2.6	2.6	2.6	2.7
Spessartine	5.6	5.8	5.2	5.1	4.9	4.5	4.1	3.6	3.3	2.6	2.7	2.3	2.3	2.2	2.1	2.1	2.0	1.9	1.8

Table 2	(continued)
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Sample	PU0810	PU0810	PU0810	PU0810	PU0810	PU0810	PU0810	PU0810	PU0810	PU0810	PU0810	PU0810	PU0810	PU0810	PU0810	PU0810	PU0810	PU0810	PU0810
SiO ₂	38.22	38.05	37.78	37.70	37.71	37.85	37.60	38.04	37.47	38.07	38.14	38.06	37.85	37.60	37.83	38.31	37.50	37.81	37.33
TiO ₂	0.08	0.06	0.07	0.00	0.03	0.06	0.03	0.07	0.05	0.08	0.10	0.07	0.04	0.05	0.06	0.04	0.05	0.06	0.05
Al_2O_3	20.96	20.87	21.05	20.88	20.71	21.17	21.18	21.01	20.97	20.79	20.97	21.21	21.04	20.98	21.19	20.68	20.96	21.01	20.85
FeO	35.45	35.79	35.96	36.42	36.21	36.41	36.45	35.94	35.89	35.72	35.81	36.49	36.06	36.00	35.67	35.53	35.26	35.49	34.96
MnO	0.91	0.79	0.80	0.79	0.77	0.71	0.85	0.80	0.76	0.73	0.76	0.79	0.86	0.96	1.19	1.35	1.76	1.88	2.42
MgO	4.12	4.22	4.10	4.17	4.11	4.08	3.99	4.09	4.09	3.97	4.02	4.08	4.02	3.87	3.86	3.58	3.60	3.52	3.24
CaO	0.91	0.86	0.91	0.89	0.85	0.90	0.90	0.88	0.91	0.89	0.85	0.89	0.81	0.93	0.93	0.86	0.87	0.87	0.92
Total	100.65	100.62	100.67	100.84	100.39	101.17	101.00	100.83	100.15	100.25	100.67	101.59	100.68	100.39	100.74	100.35	100.01	100.64	99.77
Numbers of ior	ns on the ba	sis of 12 O																	
Si	3.06	3.05	3.03	3.02	3.02	3.03	3.01	3.05	3.00	3.05	3.06	3.05	3.03	3.01	3.03	3.07	3.01	3.03	2.99
Ti	0.01	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	0.00	0.00
Al	1.98	1.97	1.99	1.97	1.96	2.00	2.00	1.98	1.98	1.96	1.98	2.00	1.99	1.98	2.00	1.95	1.98	1.98	1.97
Fe	2.38	2.40	2.41	2.44	2.43	2.44	2.44	2.41	2.41	2.39	2.40	2.45	2.42	2.41	2.39	2.38	2.36	2.38	2.34
Mn	0.06	0.05	0.05	0.05	0.05	0.05	0.06	0.05	0.05	0.05	0.05	0.05	0.06	0.07	0.08	0.09	0.12	0.13	0.16
Mg	0.49	0.50	0.49	0.50	0.49	0.49	0.48	0.49	0.49	0.47	0.48	0.49	0.48	0.46	0.46	0.43	0.43	0.42	0.39
Ca	0.08	0.07	0.08	0.08	0.07	0.08	0.08	0.08	0.08	0.08	0.07	0.08	0.07	0.08	0.08	0.07	0.07	0.07	0.08
Total	8.06	8.05	8.05	8.06	8.02	8.09	8.07	8.06	8.01	8.01	8.05	8.12	8.05	8.02	8.05	8.00	7.98	8.02	7.94
Endmembers (mol%)																		
Pyrope	16.4	16.6	16.1	16.2	16.2	16.0	15.6	16.1	16.2	15.9	16.0	15.9	15.9	15.3	15.3	14.4	14.4	14.0	13.0
Almandine	79.0	79.2	79.5	79.6	79.7	79.9	80.0	79.6	79.5	79.9	79.8	79.8	79.9	79.9	79.3	80.1	79.1	79.3	78.8
Grossular	2.6	2.4	2.6	2.5	2.4	2.5	2.5	2.5	2.6	2.6	2.4	2.5	2.3	2.7	2.7	2.5	2.5	2.5	2.6
Spessartine	2.1	1.8	1.8	1.7	1.7	1.6	1.9	1.8	1.7	1.6	1.7	1.7	1.9	2.2	2.7	3.1	4.0	4.2	5.5

Electron microprobe analysis of a garnet profile of AV0829a sample.

Sample	AV0829a	AV0829a	AV0829a	AV0829a	AV0829a	AV0829a	AV0829a	AV0829a	AV0829a	AV0829a	AV0829a	AV0829a	AV0829a	AV0829a	AV0829a	AV0829a
SiO ₂	36.82	37.09	37.08	37.40	36.88	37.16	37.22	37.26	36.81	37.02	36.79	36.73	36.88	37.16	37.22	37.26
TiO ₂	0.02	0.01	0.00	0.04	0.03	0.03	0.01	0.07	0.07	0.05	0.06	0.00	0.03	0.03	0.01	0.07
Al_2O_3	21.11	21.17	21.30	21.33	21.16	21.24	21.12	20.91	21.04	21.05	21.11	21.18	21.16	21.24	21.12	20.91
FeO	33.76	33.30	34.05	33.60	32.83	32.96	33.57	33.43	33.55	34.00	33.97	33.69	32.83	32.96	33.57	33.43
MnO	2.19	2.42	2.26	2.24	2.19	2.17	2.23	2.32	2.29	2.16	2.28	2.25	2.19	2.17	2.23	2.32
MgO	4.27	4.21	4.38	4.32	4.21	4.27	4.30	4.41	4.37	4.42	4.18	4.29	4.21	4.27	4.30	4.41
CaO	1.31	1.33	1.26	1.37	1.48	1.33	1.28	1.34	1.31	1.32	1.26	1.38	1.48	1.33	1.28	1.34
Total	99.48	99.53	100.33	100.30	98.77	99.16	99.73	99.75	99.44	100.02	99.64	99.51	98.77	99.16	99.73	99.75
Numbers	of ions on the	basis of 22 O														
Si	2.95	2.97	2.97	3.00	2.96	2.98	2.98	2.99	2.95	2.97	2.95	2.94	2.96	2.98	2.98	2.99
Ti	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00
Al	1.99	2.00	2.01	2.01	2.00	2.01	2.00	1.98	1.99	1.99	1.99	2.00	2.00	2.01	2.00	1.98
Fe	2.26	2.23	2.28	2.25	2.20	2.21	2.25	2.24	2.25	2.28	2.28	2.26	2.20	2.21	2.25	2.24
Mn	0.15	0.16	0.15	0.15	0.15	0.15	0.15	0.16	0.16	0.15	0.15	0.15	0.15	0.15	0.15	0.16
Mg	0.51	0.50	0.52	0.52	0.50	0.51	0.51	0.53	0.52	0.53	0.50	0.51	0.50	0.51	0.51	0.53
Ca	0.11	0.11	0.11	0.12	0.13	0.11	0.11	0.12	0.11	0.11	0.11	0.12	0.13	0.11	0.11	0.12
Total	7.98	7.99	8.05	8.05	7.93	7.97	8.00	8.01	7.98	8.02	7.98	7.99	7.93	7.97	8.00	8.01
Endmemb	ers (mol%)															
Ру	16.8	16.7	17.1	17.0	16.9	17.1	17.0	17.3	17.2	17.2	16.4	16.9	16.9	17.1	17.0	17.3
Alm	74.6	74.1	74.4	74.1	73.9	74.1	74.4	73.7	74.0	74.3	74.9	74.2	73.9	74.1	74.4	73.7
Gro	3.7	3.8	3.5	3.9	4.3	3.8	3.6	3.8	3.7	3.7	3.5	3.9	4.3	3.8	3.6	3.8
Sp	4.9	5.5	5.0	5.0	5.0	4.9	5.0	5.2	5.1	4.8	5.1	5.0	5.0	4.9	5.0	5.2

Appendix C. Supplementary pseudosections

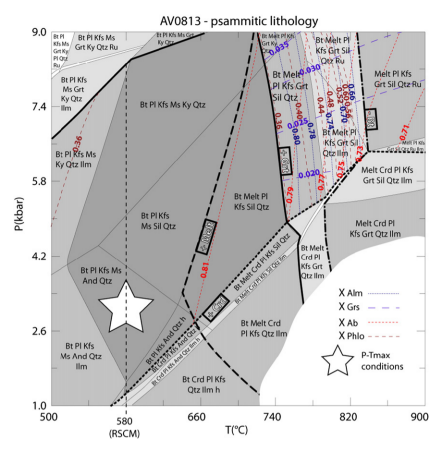


Fig. 1. *P*–*T* pseudosection of AV-08-13 metapsammite sample in the NCKFMASHT system. The white star corresponds to $P-T_{max}$ metamorphic conditions underwent by the sample. These conditions have been estimated using the observed paragenesis and RCSM thermometry. Estimated conditions are supported by the compositions of biotite and plagioclase. The heavy black arrow represents the suggested P-T path before $P-T_{max}$ conditions based on aluminosilicate sequence of Fig. 5a & b. Bulk compositions used for each pseudosection are reported in Table 5. No shading = field divariant, then light to dark shading = increasing the variance (trivariant, quadrivariant, etc.).

The sample AV-08-13 represents the psammitic endmember lithology which in the lower the La Bocana unit underwent partial melting under biotite dehydration melting. In order to validate this hypothesis, X_{Alm} , X_{Grs} isopleths of garnet and X_{Phlo} , X_{Ab} of biotite and plagioclase are also indicated in the suprasoludis region. Reported isopleth of solid-solutions in suprasolidus region are compatible with measured compositions in samples AV-08-28d, AV-08-32a, AV-08-33 and PU-08-10 (X_{Alm} , 0.72 to 0.80; X_{Grs} , 0.025 to 0.045; X_{Phlo} , 0.5 to 0.6, X_{Ab} , 0.6 to 0.7) and also indicate that partial melting occurred under biotite dehydration melting at P-T conditions of 5 to 8 kbar and 740 to 820 °C.

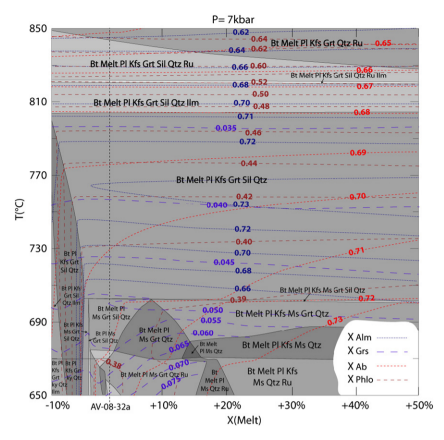


Fig. 2. *T*-*X_{Melt}* pseudosection of sample AV-08-32a in the MNNCKFMASHT system. Pressure is fixed at 7 kbar which correspond to a mean pressure in the lower La Bocana unit. Isopleths of almandine (X_{Alm}), grossular (X_{Grs}), phlogopite (X_{Phlo}) and albite (X_{Ab}) are plotted in order to estimate the influence of melt interaction on the composition of the main solid-solution of garnet, biotite and plagioclase. Variations of the bulk composition from $X_{Melt} = -10\%$ to $X_{Melt} = +50\%$ are modeled by adding or withdrawing the melt composition (in mol content) of sample AV-08-32a estimated at 7 kbar and 780 °C. No shading = field divariant, then light to dark shading = increasing the variance (trivariant, quadrivariant, etc.). This $T-X_{Melt}$ pseudosection show the influence of melt loss or gain on the mineral composition is limited in the observed field biotite + garnet + plagioclase + K-feldspar + sillimanite + quartz + melt field and supports the *P*-*T* results of the garnet-bearing migmatites.

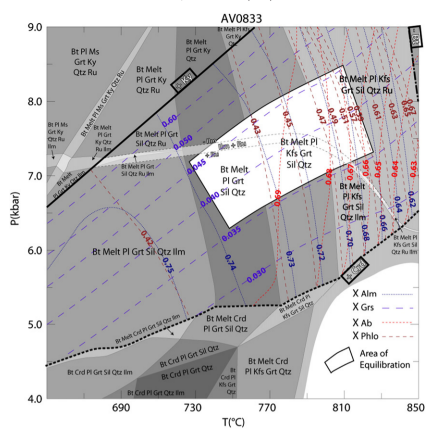


Fig. 3. *P*–*T* pseudosection of the garnet-bearing migmatitic sample AV-08-33 in the MnNCKFMASHT system. Isopleths of almandine (X_{Alm}), grossular (X_{Grs}), phlogopite (X_{Phlo}) and albite (X_{Ab}) are plotted in the field of the observed peak *P*–*T* mineralogical assemblage; the white box represents the *P*–*T* conditions of garnet, biotite and plagioclase equilibration i.e., when measured composition matches the theoretical composition constructed by isopleths. Bulk compositions used for each pseudosection are reported in Table 5. No shading = field divariant, then light to dark shading = increasing the variance (trivariant, quadrivariant, etc.).

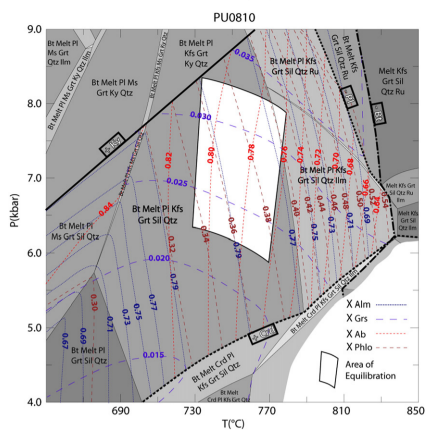


Fig. 4. *P*-*T* pseudosections of the garnet-bearing migmatitic sample PU-08-10 in the MnNCKFMASHT system. Isopleths of almandine (X_{Alm}), grossular (X_{Grs}), phlogopite (X_{Phlo}) and albite (X_{Ab}) are plotted in the field of the observed peak *P*-*T* mineralogical assemblage; the white box represents the *P*-*T* conditions of garnet, biotite and plagioclase equilibration i.e., when measured composition matches the theoretical composition constructed by isopleths. Bulk compositions used for each pseudosection are reported in Table 5. No shading = field divariant, then light to dark shading = increasing the variance (trivariant, quadrivariant, etc.).

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