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# Thermal conditions during deformation of partially molten crust from TitaniQ geothermometry: rheological implications for the anatectic domain of the Araçuaí belt, eastern Brazil

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**Abstract.** During the Neoproterozoic orogeny, the middle crust of the Araçuaí belt underwent widespread partial melting. At the regional scale, this anatectic domain is characterized by a progressive rotation of the flow direction from south to north, suggesting a 3-D deformation of the anatectic middle crust. To better determine whether melt volumes present in the anatectic middle crust of the Araçuaí orogen were large enough to allow a combination of gravity-driven and convergence-driven deformation, we used the titaniumin-quartz (TitaniQ) geothermometer to estimate the crystallization temperatures of quartz grains in the anatectic rocks. When possible, we compared these estimates with thermobarometric estimates from traditional exchange geothermobarometers applied to neighboring migmatitic kinzigites. TitaniQ temperatures range from 750 to 900 °C, suggesting that quartz starts crystallizing at minimum temperatures of ≥ 800 °C. These results, combined with the bulk-rock chemical composition of diatexites, allows the estimation of a minimum of  $\sim 30\%$  melt and a corresponding viscosity of  $\sim 10^9 - 10^{10}$  Pa s. Such a minimum melt content and low viscosity are in agreement with interconnected melt networks observed in the field. Considering that these characteristics are homogeneous over a wide area, this supports the finding that the strength of the middle crust was severely weakened by extensive partial melting, making it prone to gravitydriven flow and lateral extrusion.

#### 1 Introduction

Geophysical profiles and geological observations in hot orogens suggest that large volumes of the middle to lower crust were partially molten during deformation (e.g., Chen et al., 1996; McNamara et al., 1996; Nelson et al., 1996; Alsdorf et al., 1998; Unsworth et al., 2005). Indeed, the presence of even a small fraction of melt may dramatically reduce the strength of rocks and affect their tectonic behavior (e.g., Rosenberg and Handy, 2005; Labrousse et al., 2011). When partial melting is widespread, it may result in the development of a thick, low-strength layer in the crust that deeply alters its rheological behavior during orogeny. In this case, numerical models (e.g., Beaumont et al., 2004) suggest that, provided the viscosity of the lower to middle crust is low enough (e.g.,  $\leq 10^{19}$  Pa's Beaumont et al., 2004;  $\sim 10^{17}$  Pa's, Grujic et al., 1996), complex deformation patterns may occur in response to a combination of tectonic and gravitational forces.

The Araçuaí belt of eastern Brazil is one of several Neoproterozoic belts that involve a large anatectic domain (Oliveira et al., 2000; Vauchez et al., 2007; Petitgirard et al., 2009; Cavalcante et al., 2013). Most previous studies of the Araçuaí belt, including those studying the anatectic domain, placed emphasis on the characterization of strain distribution and/or magmatic episodes (e.g., Pedrosa-Soares et al., 2006; Vauchez et al., 2007; Mondou et al., 2012; Cavalcante et al., 2013); only few data on the thermal conditions and partial melting during deformation are available. Preliminary pressure (*P*) and temperature (*T*) estimates in mylonites from the western part of the belt (e.g., Petitgirard et

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al., 2009) suggest high-temperature ( $\sim$  750 °C), low-pressure ( $\sim$  600 MPa) synkinematic conditions.

Recently, we have shown that the anatectic domain in the eastern Araçuaí belt displays a complex 3-D strain distribution that we tentatively interpreted as resulting from the deformation of a low-stiffness anatectic middle crust in response to a combination of tectonic forces from the South America—Africa collision and gravity-driven flow (Cavalcante et al., 2013). However, temperature estimates from the anatectic domain are lacking, and such an interpretation assumes that rheological critical conditions that allowed the development of gravity-driven flow in the anatectic middle to lower crust were reached during orogeny. It is, however, difficult to evaluate, even roughly, the synkinematic viscosity of most anatectic domains without estimating equilibrium temperatures and evaluating the melt proportion present in the anatectic rocks during deformation.

Estimates of temperatures were performed using the titanium-in-quartz (TitaniQ) geothermometer developed by Wark and Watson (2006), on migmatitic rocks of the anatectic domain. We have chosen this geothermometer because it requires analysis of only one phase, quartz, its high potential to record temperatures in a major rock-forming mineral (e.g., Negrini et al., 2014), and because anatexites often do not display adequate mineralogical assemblage required for exchange geothermometers. Furthermore, microstructural observations and microprobe analysis of minerals in anatexites, migmatitic granulite and kinzigites indicate that quartz crystallized in equilibrium with rutile, and therefore represents a natural case similar to laboratory experiments performed for the calibration of the TitaniQ geothermometer. To test the reliability of the approach, migmatitic kinzigites with suitable mineralogical assemblages that bound the anatexites eastward were analyzed with both conventional thermobarometry and TitaniQ thermometry. In addition, as the titanium concentration in quartz is pressure sensitive, we have also estimated the temperatures using the calibrations by Thomas et al. (2010) and Huang and Audétat (2012). Finally, to assess the probability of a contribution of gravity-driven flow to the overall deformation, the temperature estimates and bulk rock composition of anatectic magma were used to evaluate the degree of partial melting of the anatectic middle crust during deformation. This evaluation, combined with field and microstructural observations, allowed us to constrain the viscosity of the anatectic crust more reliably and to test whether it was lower than the viscosity threshold below which topographic loading starts contributing to tectonic forces.

### 2 Geological setting

The convergence between the South American and African continents amalgamated lithospheric blocks of different compositions and ages and formed the Ribeira–Araçuaí–West Congo orogen (e.g., Trompette, 1994; Monié et al., 2012, Fig. 1). The Araçuaí belt and its southward con-

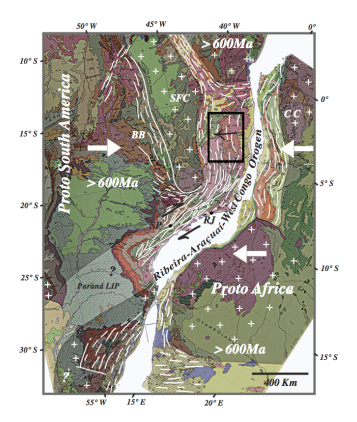


Figure 1. Schematic reconstitution of the position of South America and Africa before the opening of the South Atlantic (Vauchez et al., 2007). Shaded areas represent domains where collision occurred before  $\sim 600\,\mathrm{Ma}$  (white crosses: cratonic domains; white lines: Neoproterozoic belts). The area delimited by the rectangle represents the location of Fig. 2. Arrows show the convergence between South America and Africa. Small black arrows show the main kinematics. CC: Congo craton; BB: Brasilia belt; SP: São Paulo; RJ: Rio de Janeiro.

tinuation, the Ribeira belt, were formed by Neoproterozoic convergence between the São Francisco and Congo cratons. The Araçuaí belt is characterized by the dominant thrusting of high-temperature metamorphic allochthonous units on the São Francisco craton and subordinate orogen-parallel transcurrent motions, whereas the Ribeira belt is characterized by dominant orogen-parallel dextral transcurrent shearing combined with thrusting normal to the belt (Trompette, 1994; Egydio-Silva et al., 2002; Schmitt et al., 2004).

Schematically, the Araçuaí belt is divided into three lithological domains (Oliveira et al., 2000; Vauchez et al., 2007): the western mylonitic unit, the central plutonic unit and the eastern anatectic unit (Fig. 2a). The mylonitic domain is composed of high-temperature metasedimantary mylonites injected by a large volume of leucogranite sills. These rocks were thrust westward (top to west) onto the para-autochthonous metasediments of the São Francisco craton (Cunningham et al., 1996; Petitgirard et al., 2009). Eastward, the central plutonic domain comprises syncollisional

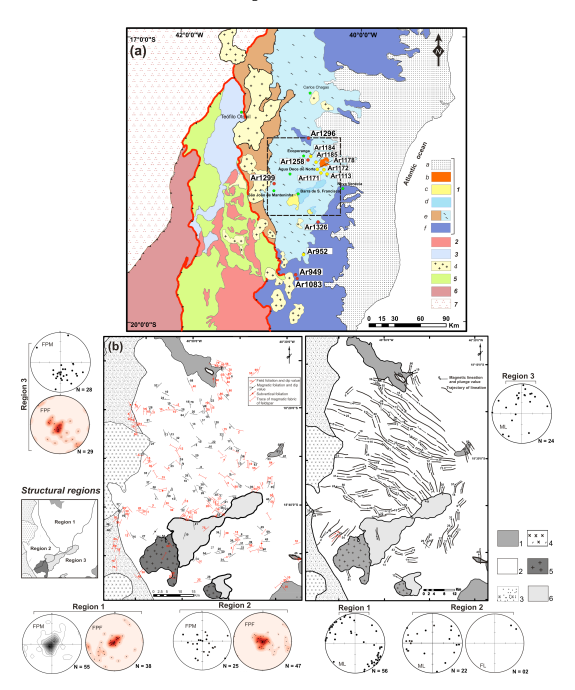


Figure 2. (a) Simplified geological map (after Oliveira et al., 2000, and Vauchez et al., 2007) showing the main domains of the Araçuaí belt and the location of the samples selected for this study (red = samples analyzed by TitaniQ and yellow = samples of geochemistry analysis): the eastern domain (1) comprises a thick (> 10 km) layer of diatexites, metatexites and anatectic granites (e) associated with migmatitic granulites (f) and intruded on by granite (c) and charnockite (d). (a) and (b) represent the Phanerozoic cover. The central domain comprises a sycollisional magmatic complex (2 = Galiléia batholith; 3 = São Vitor tonalite) intruding in HT metasediments (5). The western domain involves HT metasedimentary mylonites injected by abundant synkinematic leucocratic magma (6) thrust upon the para-autochthonous metasedimentary cover of the São Francisco craton (7). Late orogenic (~ 520 Myr, Mondou et al., 2012) porphyritic granitoids associated with charnockites (4) intrude on the stack of allochthonous units. The red line represents the boundary between lithological domains. The green pentagons represent cities. The dashed box shows the location of (b) (structural map showing the AMS and the field measurements across the anatectic domain from Cavalcante et al., 2013): left foliations; the small map on the left shows the three main structural regions as defined by their magnetic lineation pattern. The stereoplots FPM and FPF for each structural region represent the foliation poles obtained from the AMS (k3) and measured in the field, respectively. Right: lineation and lineation traces illustrating the magmatic flow. The ML (k1) and FL stereoplots for each structural region represent the lineation obtained from the AMS and measured in the field, respectively. All the stereoplots are represented in the lower hemisphere of the equal-area projection. 1, 2, 3, 4, 5 and 6 are, respectively, 1f, 1e (Carlos Chagas anatexite), 1e, 4, 1c and 1d of (a).

tonalite and granodiorite bodies deformed in the magmatic state (Mondou et al., 2012). The eastern anatectic domain is dominated by a variety of anatexites and anatectic granites that result from pervasive partial melting of metasedimentary rocks under mid-crustal P and T conditions (Vauchez et al., 2007). This anatectic domain is  $\sim 300 \, \mathrm{km}$  long and 50– 100 km wide. Migmatitic granulite forms rafts embedded in the anatexites. Migmatitic kinzigites occur east and southeast of the anatexites; they progressively grade westward to diatexites and aluminous granites, suggesting a transition from kinzigites to anatexites. Structural mapping in the anatectic unit using the anisotropy of magnetic susceptibility (AMS) highlighted a complex pattern of lineation, the trend of which rotates from WSW-ENE in the southern part of the study area to NW-SE in the northern part, and to N-S in the easternmost part (Cavalcante et al., 2013; Fig. 2b). Flow in varying directions is expected during deformation of a weakened crust that cannot support the topographic load; such deformation "is analogous to pressing on an egg sandwich: the soft filling is squeezed out and emerges at the edge of the sandwich and through any holes in the bread" (Jamieson et al., 2011). Considering the progressive transition between the various dominant orientations of lineation together with evidence supporting the finding that, independently of the flow direction, the fabric was acquired in the magmatic state (Cavalcante et al., 2013), led us to suggest that the complex deformation of the anatexites results from a single, probably protracted, tectonic event.

Thermochronological data from the mylonitic domain suggest that this region remained hot (750 to  $> 500\,^{\circ}\text{C}$ ) for tens of millions of years (580–530 Ma) due to a low cooling rate ( $\leq 5\,^{\circ}\text{C}\,\text{Myr}^{-1}$ ; Petitgirard et al., 2009). Conventional thermobarometric estimates performed by Munhá et al. (2005) in migmatitic gneisses from the southern part of the Araçuaí belt suggest that the metamorphic peak and partial melting took place at temperatures of  $820\pm30\,^{\circ}\text{C}$  and at pressures of  $650\pm50\,\text{MPa}$ .

#### 3 Samples: location and description

Thirteen samples were selected from the anatectic unit (Fig. 2a). Six samples were used for TitaniQ analyses (Ar1258, Ar1299, Ar1326, Ar1296, Ar949 and Ar1083), and seven samples for geochemistry analyses (Ar1113, Ar952, Ar1171, Ar1172, Ar1178, Ar1184 and Ar1185). All these samples are anatexites, however, with different volumes of melt and composition. Nine samples come from the diatexites (Ar1258, Ar1299, Ar1113, Ar952, Ar1171, Ar1172, Ar1178, Ar1184 and Ar1185), one from the metatexite (Ar1326), one from the migmatitic granulite of the northern domain (Ar1296), and two from the migmatitic kinzigites of the southeastern domain (Ar949 and Ar1083). At the outcrop scale, the diatexites display a clear magmatic foliation marked by the preferred alignment of biotite and feldspar

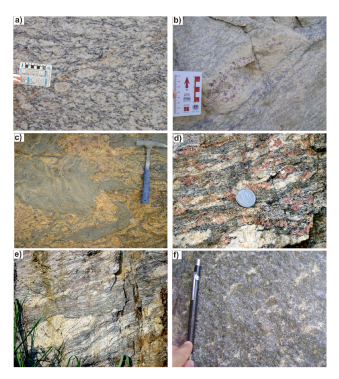


Figure 3. Pictures of outcrops representative of the studied rocks: (a) an example of diatexite with well-marked magmatic foliation characterized by the preferred alignment of biotite and feldspar; (b) diatexite with leucosome rich in garnet and feldspar; (c) diatexite displaying a scholle feature. (d) and (e): migmatitic kinzigites displaying stromatic leucosomes embedded between gneissic foliations; (f) a migmatitic granulite with small pockets of melt.

grains (Fig. 3a). In situ leucosomes are frequently observed in metatexites that preserve a gneissic banding (Fig. 4). Migmatitic features such as stromatic, nebulitic and schlieren leucosomes, rich in garnet and feldspar, are also common (Fig. 3b, c).

The migmatitic kinzigites display a gneissic banding characterized by alternating felsic and mafic layers. Stromatic and coarser-grained leucosomes, rich in feldspar, garnet and quartz, are embedded parallel to banding (Fig. 3d, e).

The migmatitic granulite is mafic in composition and does not show a clear macroscopic fabric. It does, however, contain narrow strings of leucosomes pointing to limited partial melting (Fig. 3f).

## **Petrographic observations**

Diatexites, metatexites and migmatitic kinzigites are composed of K-feldspar, quartz, plagioclase, biotite, garnet, cordierite, sillimanite, ilmenite and rutile. The migmatitic granulite also contains orthopyroxene  $\pm$  magnetite. Most minerals do not or rarely show evidence of intracrystalline deformation; this suggests a predominantly magmatic fabric free of superimposed solid-state deformation.



**Figure 4.** Pictures (left) and schematic drawing (right) of partially molten kinzigites of the Araçuaí belt. White represents the in situ leucosomes and gray represents the residuum and/or paleosome portions.

Quartz usually displays interstitial shapes (Fig. 5a), although this is less frequent in the migmatitic granulite that contains less quartz. It has irregular grain boundaries, with sizes ranging from  $\sim\!200\,\mu m$  to  $\sim\!2$  mm, and is mostly free of substructure, with the exception of an undulose extinction or faint subgrain boundaries (Fig. 5b).

Plagioclase reaches  $\sim 0.4\,\mathrm{cm}$  in size, and predominantly displays subeuhedral shapes; in many cases, it is surrounded by interstitial films of quartz along grain boundaries (Fig. 5c). In the migmatitic granulites, many crystals of plagioclase display deformation twins, attesting to the high temperature deformation during partial melting (Fig. 5d).

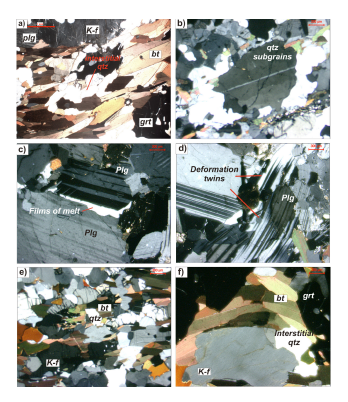
K-feldspar is  $\sim 0.3$  cm in size, and displays curved, lobate and straight grain boundaries. It often exhibits myrmekites, and is free of evidence of intracrystalline deformation.

Biotite displays a strong preferred orientation (Fig. 5a, e). It is euhedral, subeuhedral and platy in shape, with straight to curved grain boundaries. It tends to arch around large K-feldspars (Fig. 5f). In the migmatitic kinzigites, a few kinked biotites are observed.

Sillimanite is acicular and garnet is usually rounded, but locally elongate parallel to foliation, with frequent inclusions of quartz, biotite, ilmenite and rutile.

## 4 TitaniQ temperature estimates

The TitaniQ geothermometer is based on substitution of Si by Ti in quartz. Since the substitution is dependent on temperature (Wark and Watson, 2006), Ti concentration in quartz is correlated with its formation temperature. TitaniQ has already been applied to characterize thermal histories of migmatitic (Storm and Spear, 2009), mylonitic (Kohn and Northrup, 2009; Grujic et al., 2011) and volcanic rocks (e.g.,



**Figure 5.** Photomicrographs of anatexites and migmatitic granulites. (a) Large interstitial quartz and preferred orientation of biotite characterizing the magmatic foliation, (b) quartz subgrains representing the record of the intracrystalline deformation, (c) film of quartz crystallized along grain boundaries that likely represent former films of melt, (d) deformation twins in plagioclase feldspar, (e) preferred orientation of biotite grains, and (f) large K-feldspar grains associated with interstitial quartz and arched biotite.

Ehrlich et al., 2012). To develop this geothermometer, Wark and Watson (2006) synthesized quartz in the presence of rutile and aqueous fluid or hydrous silicate melt at temperatures ranging from 600 to  $1000\,^{\circ}\text{C}$  and  $1.0\,\text{GPa}$  of pressure. This geothermometer relates the temperature (T) of crystallization to the Ti content of quartz according to

$$Log(Ti_{quartz}) = 5.69 - 3765/T,$$

where  $Ti_{quartz}$  is the Ti concentration in quartz in ppm and T is the temperature in K. No adjustment was necessary for this equation because both rutile and ilmenite were present in the analyzed samples. Wark and Watson (2006) calibration however does not take into account the pressure dependence of Ti concentrations in quartz. Thomas et al. (2010), then Huang and Audétat (2012), experimentally calibrated a TitaniQ thermobarometer. Thomas et al. (2010) data are fitted

by the equation

$$\begin{split} RT \ln X_{\text{TiO}_2}^{\text{Quartz}} &= -60952 + 1520 \cdot T\left(\mathbf{K}\right) - 1741 \cdot P(\text{kbar}) \\ &+ RT \ln a_{\text{TiO}_2}, \end{split}$$

where R is the gas constant  $8.3145 \,\mathrm{J\,K^{-1}}$ , T is the temperature in Kelvin,  $X_{\mathrm{TiO}_2}^{\mathrm{Quartz}}$  is the mole fraction of  $\mathrm{TiO}_2$  in quartz, and  $a_{\mathrm{TiO}_2}$  is the activity of  $\mathrm{TiO}_2$  in the system. Those from the Huang and Audétat (2012) experiments are fitted by

$$\log \text{Ti(ppm)} = -0.27943 \times 10^4 / T$$
$$-660.53 \left( P^{0.35} / T \right) + 5.6459,$$

where *T* is given in K and *P* in kbar.

We have therefore also calculated the crystallization temperatures using these two calibrations for comparison.

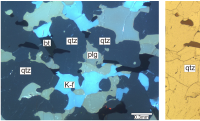
#### 5 Material and procedure

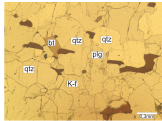
Microprobe analysis and backscattered electron images were performed and produced with a CAMECA SX-100 electron probe instrument equipped with five wavelength-dispersive X-ray spectrometers. Some of the X-ray spectrometers are equipped with a large, high-efficiency Johansson crystal. Before microprobe analysis, the selected samples were imaged by cathodoluminescence, and no evidence of zoning was observed (Fig. 6). All quartz grains display an intense homogeneous dark blue color, suggesting that multiple thermal events are unlikely. We have selected large and/or interstitial quartz grains (≥ 0.3 mm) that have similar aspects, no inclusion and no evidence of intracrystalline deformation or microfracture.

Four to five quartz grains from each sample were analyzed. Microprobe analyses were made along transverse profiles with a spacing of at least  $120\,\mu m$  (Fig. 7), and preferentially in areas free of rutile and impurities. More than 370 Ti-in-quartz analyses were performed on samples from the anatectic domain.

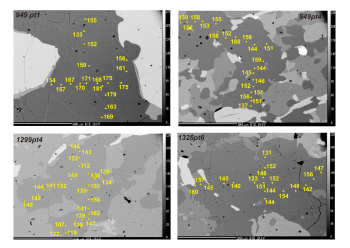
## 5.1 Analytical procedures

Accurate characterization of trace elements using electron probe micro-analysis (EPMA) requires optimal analytical conditions (e.g., Merlet and Bodinier, 1990; Bodinier et al., 1996; Kalfoun et al., 2002a, b). The choice of the accelerating voltage value is essential, because it influences the X-ray emission yield and bias. The best spatial resolution is obtained with a low accelerating voltage, whereas higher voltages are needed for transition and trace elements. These constraints require a compromise between X-ray line, peak counting statistics, background subtraction, electron beam size, beam drift, quantification uncertainty, stray radiation and sample damage. In order to find the optimum balance, recent developments in microanalysis, such as spectrum simulation and X-ray emission modeling (Fournier et al., 1999; Kalfoun et al., 2002a, b), were taken into account.





**Figure 6.** CL (right) and true-color (left) images of the migmatitic kinzigite (sample Ar949) displaying the intense homogenous dark blue of quartz grains.



**Figure 7.** Spots along transverse profiles and respective titanium concentrations (ppm) in quartz from migmatitic kinzigite (samples Ar949 and Ar1326) and from diatexite (Ar1299).

Traces of titanium in quartz are relatively easy to identify, but care must be taken against sample damage and Bremsstrahlung fluorescence (braking radiation) of Ti atoms in close proximity to rutile (Wark and Watson, 2006), ilmenite and biotite. The Bremsstrahlung effect is a form of secondary fluorescence produced by elastic interaction of the 15 kV incident electrons with the quartz lattice (e.g., Bastin et al., 1984) that can be calculated by Monte Carlo simulation. At 20 kV accelerating voltage, a minimum gap of 200 microns between these minerals is required in order to avoid erroneous estimates of Ti concentrations.

Concentrations were measured using an accelerating voltage of 20 kV, a beam current of 200 nA and a 50-micron defocused probe to reduce irradiation damage. An electron beam current was selected to compromise between a high X-ray counting rate and minimal sample damage. The Ti concentration was determined by simultaneously counting Ti K-alpha X-ray lines using three spectrometers with large pentaerythritol (LPET) crystals in a Johansson configuration and averaging the results.

Bremsstrahlung background was subtracted using a method that combines the measured and calculated

background spectra from samples for which neither spectral lines nor absorption edges were observed in the region of interest (Kalfoun et al., 2002). The peak and background acquisition time for each point for all elements and all five spectrometers was 480 s. The counting time of 480 s is the maximum time that a defocused beam of 50 microns can shine on a sample without damaging it significantly. The corresponding integration time and beam current provide a typical statistical error (peak counting) of approximately 1.5% for a 100 ppm Ti concentration, and thus a 7 ppm limit of detection. Concentrations are obtained from raw intensities by using the X-PHI quantification procedure (Merlet, 1994). Natural SiO<sub>2</sub> and TiO<sub>2</sub> are used as standards for Si and Ti, respectively.

The global uncertainty of Ti concentrations at a single point, obtained by adding the random and systematic uncertainties in quadrature, is  $\pm 6\%$  ( $2\sigma$ ) at  $\geq 100$  ppm; these estimates consider errors from peak counting (1.5%), background estimates (2%), sample damage and contamination (1.5%), and uncertainties in standard composition and quantification procedures (1%).

#### 5.2 Results

The Ti content in quartz from diatexites, metatexites, migmatitic kinzigites and granulite, and the calculated temperatures, are presented in Table 1 and Fig. 8. Temperature was calculated from each analytical spot. The average Ti content is  $\sim$  129 ppm in anatexites (diatexites and metatexite),  $\sim 142 \, \mathrm{ppm}$  in migmatitic kinzigites and  $\sim 112 \, \mathrm{ppm}$  in migmatitic granulite. The average crystallization temperature calculated for all quartz grains from the selected samples ranges from 840 to 900 °C in Huang and Audétat's (2012) calibration, from 670 to 730 °C in the Thomas et al. (2010) calibration, and from 700 to 810 °C in Wark and Watson's (2006) equation. Single quartz crystals display a rather homogenous distribution of titanium, and consequently uniform temperature values (Table 1; see values for individuals grains). Specimen Ar1296, a mafic granulite, constitutes an exception, with temperature averages for grains 1 and 3 (ar1296pt01 and ar1296pt03 in Table 1) exhibiting a variation of greater than 100 °C. This rock is derived from a mafic magmatic protolith in which the presence of a rather small amount of interstitial quartz and a large amount of plagioclase locally showing deformation twins (Fig. 5d) suggests limited partial melting during high-temperature, solid-state deformation.

For the diatexites and metatexites, calculated temperature averages obtained from the Huang and Audétat (2012), Thomas et al. (2010) and Wark and Watson (2006) calibrations are, respectively,  $848\pm20\,^{\circ}\mathrm{C}$ ,  $676\pm10\,^{\circ}\mathrm{C}$  and  $747\pm15\,^{\circ}\mathrm{C}$  for sample Ar1258,  $868\pm20\,^{\circ}\mathrm{C}$ ,  $703\pm10\,^{\circ}\mathrm{C}$  and  $780\pm16\,^{\circ}\mathrm{C}$  for sample Ar1299 from the central part of the study area, and  $881\pm20\,^{\circ}\mathrm{C}$ ,  $715\pm10\,^{\circ}\mathrm{C}$ , and  $792\pm17\,^{\circ}\mathrm{C}$  for sample Ar1326 from the southeast, and for the partially

molten kinzigites, high temperature averages are observed,  $884\pm20\,^{\circ}\text{C}$ ,  $717\pm10\,^{\circ}\text{C}$  and  $794\pm17\,^{\circ}\text{C}$  for sample Ar949 and  $877\pm20\,^{\circ}\text{C}$ ,  $710\pm10\,^{\circ}\text{C}$  and  $787\pm17\,^{\circ}\text{C}$  for sample Ar1083.

#### 6 Exchange reaction thermobarometry

Thermobarometric estimates using exchange reactions were determined from the migmatitic kinzigites (samples Ar949, Ar1083 and Ar1076) for comparison with the temperatures determined with the TitaniQ thermometry. Sample Ar1076 collected close to site Ar1083 was not analyzed for TitaniQ thermometry, but was used for exchange thermobarometry due to an adequate mineral assemblage.

Temperatures were obtained using the geothermometer developed by Ferry and Spear (1978) and Bhattacharya et al. (1988, 1991). Pressures were estimated using the garnet-plagioclase-biotite-quartz (GPBQ) geobarometer (Wu et al., 2004). We chose these geothermobarometers by considering the textural characteristics of the analyzed mineral pairs, i.e., those displaying equilibrium relations expected on the temperature interval on which the geothermobarometers were calibrated. The resulting P-T estimates are presented in Table 2.

Cores and rims of garnet, biotite, plagioclase, cordierite and orthopyroxene were analyzed (Table 3). Garnet-biotite, garnet-orthopyroxene and garnet-cordierite geothermometers and a GPBQ geobarometer applied to analyses from the cores of the crystals suggest that the minimum temperatures for peak metamorphic conditions recorded in migmatitic kinzigites range from 720 to  $840 \pm 50$  °C, and pressure from 650 to  $700 \pm 100$  MPa. These temperatures are close to those obtained with TitaniQ for the same samples. They usually are between the temperatures calculated using the Wark and Watson (2006) and Huang and Audétat (2012) calibrations: they are however higher than those computed with the Thomas et al. (2010) calibration. Significantly lower temperatures (550–780 °C) obtained in garnet, biotite and opx rims likely represent cooling during retrogressive metamorphic conditions. However, more data are required to constrain the significance of these values better.

#### 7 Discussion

Paleotemperatures recorded by the anatectic rocks of the eastern part of the Araçuaí belt have been calculated from Ti content in quartz. Individual crystals display rather homogeneous compositions (Fig. 7), suggesting that Ti diffusion after crystallization was limited. For instance, analysis from sample AR949 shows a good correlation between temperatures estimated from Ti in quartz and those estimated from garnet and biotite core compositions (Tables 1 and 2). Garnet and biotite rim compositions however have recorded cooling to significantly lower temperatures, which was not recorded

 Table 1. Titanium content and calculated TitaniQ temperatures for each analytical spot, with the average values shown for each sample.

Sample-crystal number	Lithology	Ti (ppm)		$T$ (°C), $P = 650 \mathrm{Mpa}$		
			Thomas et al. (2010)	Wark and Watson (2006)	Huang and Audétat (2012	
ar1296pt01	Migmatitic granulite	92	667	737	831	
ar1296pt01	Migmatitic granulite	98	673	745	840	
ar1296pt01	Migmatitic granulite	89	663	733	827	
ar1296pt01	Migmatitic granulite	27	557	611	692	
ar1296pt01	Migmatitic granulite	83	656	725	818	
ar1296pt02	Migmatitic granulite	112	687	761	858	
ar1296pt02	Migmatitic granulite	110	685	759	855	
ar1296pt02	Migmatitic granulite	109	684	758	854	
ar1296pt02	Migmatitic granulite	118	693	767	865	
ar1296pt02	Migmatitic granulite	136	708	785	885	
ar1296pt02	Migmatitic granulite	141	712	790	890	
ar1296pt03	Migmatitic granulite	91	665	736	830	
ar1296pt03	Migmatitic granulite	107	682	755	851	
ar1296pt03	Migmatitic granulite	156	723	804	905	
ar1296pt03	Migmatitic granulite	265	788	879	989	
ar1296pt03	Migmatitic granulite	103	678	751	846	
ar1296pt04	Migmatitic granulite	97	672	744	838	
ar1296pt04	Migmatitic granulite	91	665	736	830	
ar1296pt04	Migmatitic granulite	101	676	748	844	
ar1296pt04	Migmatitic granulite	122	696	772	869	
ar1296pt04	Migmatitic granulite	103	678	751	846	
ar1296pt04	Migmatitic granulite	88	662	732	825	
ar1296pt05	Migmatitic granulite	104	679	752 752	847	
ar1296pt05 ar1296pt05	Migmatitic granulite	112	687	761	858	
ar1296pt05 ar1296pt05	Migmatitic granulite	125	699	775	873	
ar1296pt05 ar1296pt05	Migmatitic granulite	123	698	774	873 872	
ar1296 (average)	Migmatitic granulite	112	$687 \pm 10$	$755 \pm 16$	$842 \pm 20$	
ar1299pt01	Diatexite	117	692	766	863	
ar1299pt01	Diatexite	142	713	791	891	
ar1299pt01	Diatexite	136	708	785	885	
ar1299pt01	Diatexite	123	697	773	870	
ar1299pt01	Diatexite	128	701	778	876	
ar1299pt01	Diatexite	126	700	776	874	
ar1299pt01	Diatexite	132	705	782	880	
ar1299pt01	Diatexite	153	721	801	902	
ar1299pt01	Diatexite	136	708	785	885	
ar1299pt01	Diatexite	120	694	770	867	
ar1299pt01	Diatexite	132	705	782	880	
ar1299pt01	Diatexite	130	703	780	878	
ar1299pt01	Diatexite	124	698	774	872	
ar1299pt01	Diatexite	130	703	780	878	
ar1299pt01	Diatexite	119	693	769	866	
ar1299pt01	Diatexite	132	705	782	880	
ar1299pt01	Diatexite	115	690	764	861	
ar1299pt01	Diatexite	114	689	763	860	
ar1299pt03	Diatexite	117	692	766	863	
ar1299pt03	Diatexite	125	699	775	873	
ar1299pt03 ar1299pt03	Diatexite	118	693	773 767	865	
	Diatexite	118	693	767 767	865	
ar1299nt03				787 782		
	Diatevite	127				
ar1299pt03 ar1299pt03	Diatexite	132	705 700		880 875	
	Diatexite Diatexite Diatexite	132 127 149	705 700 718	782 777 797	880 875 898	

Table 1. Continued.

Sample-crystal number	Lithology	Ti (ppm)		$T$ (°C), $P = 650 \mathrm{Mpa}$	
			Thomas et al. (2010)	Wark and Watson (2006)	Huang and Audétat (2012)
ar1299pt03	Diatexite	136	708	785	885
ar1299pt03	Diatexite	124	698	774	872
ar1299pt03	Diatexite	138	710	787	887
ar1299pt03	Diatexite	108	693	757	853
ar1299pt03	Diatexite	122	696	772	869
ar1299pt03	Diatexite	144	714	793	893
ar1299pt03	Diatexite	135	707	785	884
ar1299pt03	Diatexite	138	710	787	887
ar1299pt03	Diatexite	123	697	773	870
ar1299pt04	Diatexite	122	696	772	869
ar1299pt04	Diatexite	119	693	769	866
ar1299pt04	Diatexite	107	682	755	851
ar1299pt04	Diatexite	139	710	788	888
ar1299pt04	Diatexite	139	710	788	888
ar1299pt04	Diatexite	147	717	796	896
ar1299pt04	Diatexite	162	728	809	910
ar1299pt04	Diatexite	141	712	790	890
ar1299pt04	Diatexite	156	723	804	905
ar1299pt04	Diatexite	135	707	785	884
ar1299pt04	Diatexite	150	719	798	899
ar1299pt04	Diatexite	136	708	785	885
ar1299pt04	Diatexite	134	706	784	883
ar1299pt04	Diatexite	134	706	784	883
ar1299pt04	Diatexite	112	687	761	858
ar1299pt04	Diatexite	122	696	772	869
ar1299pt04	Diatexite	143	714	792	892
ar1299pt04	Diatexite	146	716	795	895
ar1299pt04	Diatexite	144	714	793	893
ar1299pt04	Diatexite	152	721	800	901
ar1299pt04	Diatexite	141	712	790	890
ar1299pt05	Diatexite	123	697	773	870
ar1299pt05	Diatexite	115	690	764	861
ar1299pt05	Diatexite	124	698	774	872
ar1299pt05	Diatexite	127	700	777	875
ar1299pt05	Diatexite	124	698	774	872
ar1299pt05	Diatexite	130	703	780	878
ar1299pt05	Diatexite	121	695	771	868
ar1299pt05	Diatexite	126	700	776	874
ar1299pt05	Diatexite	140	711	789	889
ar1299pt05	Diatexite	118	693	767	865
ar1299pt05	Diatexite	109	684	758	854
ar1299 (average)	Diatexite	130	$703 \pm 10$	$780 \pm 16$	$868 \pm 20$
ar1258pt01	Diatexite	82	655	724	816
ar1258pt01	Diatexite	102	677	750	845
ar1258pt01	Diatexite	79	651	720	812
ar1258pt01	Diatexite	75	646	714	805
ar1258pt01	Diatexite	82	655	724	816
ar1258pt01	Diatexite	88	662	732	825
ar1258pt01	Diatexite	85	659	728	821
ar1258pt01	Diatexite	143	714	792	892
ar1258pt01	Diatexite	92	667	737	831
ar1258pt01	Diatexite	112	687	761	858
ar1258pt01	Diatexite	93	668	739	833

Table 1. Continued.

Sample-crystal number	Lithology	Ti (ppm)		$T$ (°C), $P = 650 \mathrm{Mpa}$	
			Thomas et al. (2010)	Wark and Watson (2006)	Huang and Audétat (2012)
ar1258pt01	Diatexite	88	662	732	825
ar1258pt01	Diatexite	92	667	737	831
ar1258pt01	Diatexite	96	671	742	837
ar1258pt01	Diatexite	87	661	731	824
ar1258pt01	Diatexite	126	700	776	874
ar1258pt01	Diatexite	85	659	728	821
ar1258pt01	Diatexite	101	676	748	844
ar1258pt01	Diatexite	99	674	746	841
ar1258pt01	Diatexite	86	660	729	822
ar1258pt01	Diatexite	88	662	732	825
ar1258pt01	Diatexite	86	660	729	822
ar1258pt01	Diatexite	82	655	724	816
ar1258pt01	Diatexite	78	650	718	810
ar1258pt01	Diatexite	77	649	717	808
ar1258pt01	Diatexite	83	656	725	818
ar1258pt01	Diatexite	103	678	751	846
ar1258pt02	Diatexite	97	672	744	838
ar1258pt02	Diatexite	104	679	752	847
ar1258pt02	Diatexite	88	662	732	825
ar1258pt02	Diatexite	99	674	746	841
ar1258pt02	Diatexite	112	687	761	858
ar1258pt03	Diatexite	89	663	733	827
ar1258pt03	Diatexite	97	672	744	838
ar1258pt03	Diatexite	147	717	796	896
ar1258pt03	Diatexite	101	676	748	844
ar1258pt03	Diatexite	77	649	717	808
ar1258pt03	Diatexite	87	661	731	824
ar1258pt03	Diatexite	108	693	757	853
ar1258pt03	Diatexite	89	663	733	827
ar1258pt03	Diatexite	99	674	746	841
ar1258pt03	Diatexite	121	695	771	868
ar1258pt03	Diatexite	84	657	727	819
ar1258pt03	Diatexite	100	675	747	842
ar1258pt03	Diatexite	93	668	739	833
ar1258pt04	Diatexite	96	671	742	837
ar1258pt04	Diatexite	97	672	744	838
ar1258pt04	Diatexite	96	671	742	837
ar1258pt04	Diatexite	88	662	732	825
ar1258pt04	Diatexite	90	664	735	828
ar1258pt04	Diatexite	89	663	733	827
ar1258pt04	Diatexite	91	665	736	830
ar1258pt04	Diatexite	108	693	757	853
ar1258pt04	Diatexite	95	670	741	835
ar1258pt04	Diatexite	101	676	748	844
ar1258pt04	Diatexite	97	672	744	838
ar1258pt04	Diatexite	112	687	761	858
ar1258pt04	Diatexite	102	677	750	845
ar1258pt05	Diatexite	131	704	781	879
ar1258pt05	Diatexite	98	673	745	840
ar1258pt05	Diatexite	91	665	736	830
ar1258pt05	Diatexite	92	667	737	831
ar1258pt05	Diatexite	102	677	750	845
ar1258pt05	Diatexite	94	669	740	834
ar1258pt05	Diatexite	103	678	751	846

Table 1. Continued.

Sample-crystal number	Lithology	Ti (ppm)		$T$ (°C), $P = 650 \mathrm{Mpa}$	
			Thomas et al. (2010)	Wark and Watson (2006)	Huang and Audétat (2012)
ar1258pt05	Diatexite	111	686	760	856
ar1258pt05	Diatexite	120	694	770	867
ar1258pt05	Diatexite	181	741	824	927
ar1258pt05	Diatexite	92	667	737	831
ar1258pt05	Diatexite	93	668	739	833
ar1258pt05	Diatexite	202	754	839	944
ar1258pt05	Diatexite	100	675	747	842
ar1258pt05	Diatexite	116	691	765	862
ar1258pt05	Diatexite	99	674	746	841
ar1258pt05	Diatexite	97	672	744	838
ar1258pt05	Diatexite	137	709	786	886
ar1258pt05	Diatexite	101	676	748	844
ar1258pt05	Diatexite	98	673	745	840
ar1258pt05	Diatexite	123	697	773	870
ar1258pt05	Diatexite	106	681	754	850
ar1258pt05	Diatexite	108	693	757	853
ar1258pt05	Diatexite	95	670	741	835
ar1258pt05	Diatexite	109	684	758	854
ar1258pt05	Diatexite	129	702	779	877
ar1258pt05	Diatexite	100	675	747	842
ar1258pt05	Diatexite	85	659	728	821
ar1258 (average)	Diatexite	101	$676 \pm 10$	$747 \pm 15$	$848 \pm 20$
ar1326pt02	Metatexite	112	687	761	858
ar1326pt02	Metatexite	157	724	804	906
ar1326pt02	Metatexite	157	724	804	906
ar1326pt02	Metatexite	155	723	803	904
ar1326pt02	Metatexite	154	722	802	903
ar1326pt02	Metatexite	103	678	751	846
ar1326pt02	Metatexite	15	512	561	637
ar1326pt02	Metatexite	157	724	804	906
ar1326pt02	Metatexite	150	719	798	899
ar1326pt02	Metatexite	138	710	787	887
ar1326pt02	Metatexite	137	709	786	886
ar1326pt03	Metatexite	158	725	805	907
ar1326pt03	Metatexite	152	721	800	901
ar1326pt03	Metatexite	145	715	794	894
ar1326pt03	Metatexite	156	723	804	905
ar1326pt03	Metatexite	143	714	792	892
ar1326pt03	Metatexite	129	702	779	877
ar1326pt03	Metatexite	124	698	774	872
ar1326pt03	Metatexite	110	685	759	855
ar1326pt03	Metatexite	127	700	777	875
ar1326pt03	Metatexite	128	701	778	876
ar1326pt03	Metatexite	137	709	786	886
ar1326pt03	Metatexite	148	717	797	897
ar1326pt03	Metatexite	157	724	804	906
ar1326pt03	Metatexite	136	708	785	885
ar1326pt03	Metatexite	131	704	781	879
ar1326pt03	Metatexite	123	697	773	870
ar1326pt03	Metatexite	135	707	785	884
ar1326pt03	Metatexite	128	701	778	876
ar1326pt06	Metatexite	131	704	781	879
ar1326pt06	Metatexite	152	721	800	901

Table 1. Continued.

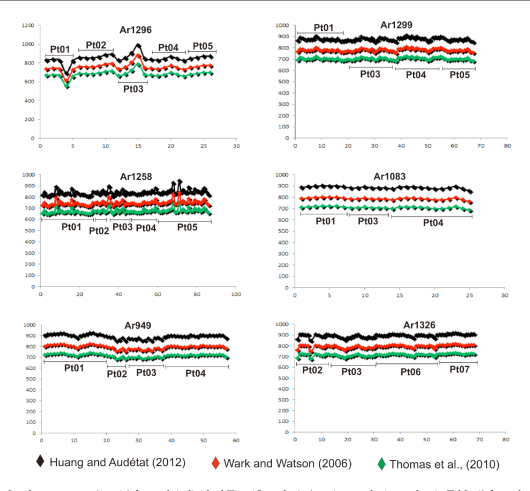
Sample-crystal number	Lithology	Ti (ppm)	$T$ (°C), $P = 650 \mathrm{Mpa}$					
			Thomas et al. (2010)	Wark and Watson (2006)	Huang and Audétat (2012)			
ar1326pt06	Metatexite	148	717	797	897			
ar1326pt06	Metatexite	144	714	793	893			
ar1326pt06	Metatexite	144	714	793	893			
ar1326pt06	Metatexite	146	716	795	895			
ar1326pt06	Metatexite	157	724	804	906			
ar1326pt06	Metatexite	162	728	809	910			
ar1326pt06	Metatexite	160	726	807	908			
ar1326pt06	Metatexite	154	722	802	903			
ar1326pt06	Metatexite	160	726	807	908			
ar1326pt06	Metatexite	157	724	804	906			
ar1326pt06	Metatexite	145	715	794	894			
ar1326pt06	Metatexite	145	715	794	894			
ar1326pt06	Metatexite	140	711	789	889			
ar1326pt06	Metatexite	123	697	773	870			
ar1326pt06	Metatexite	151	720	799	900			
ar1326pt06	Metatexite	152	721	800	901			
ar1326pt06	Metatexite	154	722	802	903			
ar1326pt06	Metatexite	148	717	797	897			
ar1326pt06	Metatexite	142	713	791	891			
ar1326pt06	Metatexite	156	723	804	905			
ar1326pt06	Metatexite	147	717	796	896			
ar1326pt06	Metatexite	124	698	774	872			
ar1326pt07	Metatexite	153	721	801	902			
ar1326pt07	Metatexite	160	726	807	908			
ar1326pt07	Metatexite	160	726	807	908			
ar1326pt07	Metatexite	154	722	802	903			
ar1326pt07	Metatexite	166	731	812	914			
ar1326pt07	Metatexite	162	728	809	910			
ar1326pt07	Metatexite	170	733	815	917			
ar1326pt07	Metatexite	171	733	816	918			
ar1326pt07	Metatexite	165	730	811	913			
ar1326pt07	Metatexite	161	727	808	909			
ar1326pt07	Metatexite	150	719	798	899			
ar1326pt07	Metatexite	159	726	806	907			
ar1326pt07	Metatexite	160	726	807	908			
ar1326pt07	Metatexite	162	728	809	910			
ar1326pt07	Metatexite	161	727	808	909			
ar1326 (average)	Metatexite	145	$715 \pm 10$	$792 \pm 17$	$881 \pm 20$			
ar1083pt01	Migmatitic kinzigites	139	710	788	888			
ar1083pt01	Migmatitic kinzigites	145	715	794	894			
ar1083pt01	Migmatitic kinzigites	150	719	798	899			
ar1083pt01	Migmatitic kinzigites	153	721	801	902			
ar1083pt01	Migmatitic kinzigites	152	721	800	901			
ar1083pt01	Migmatitic kinzigites	152	721	800	901			
ar1083pt01	Migmatitic kinzigites	145	715	794	894			
ar1083pt03	Migmatitic kinzigites	134	706	784	883			
ar1083pt03	Migmatitic kinzigites	136	708	785	885			
ar1083pt03	Migmatitic kinzigites	144	714	793	893			
ar1083pt03	Migmatitic kinzigites	135	707	785	884			
ar1083pt03	Migmatitic kinzigites	134	706	784	883			
ar1083pt03	Migmatitic kinzigites	136	707	785	885			
ar1083pt04	Migmatitic kinzigites	126	700	776	874			
ar1083pt04	Migmatitic kinzigites	142	713	791	891			

Table 1. Continued.

Sample-crystal number	Lithology	Ti (ppm)	$T$ (°C), $P = 650 \mathrm{Mpa}$					
			Thomas et al. (2010)	Wark and Watson (2006)	Huang and Audétat (2012			
ar1083pt04	Migmatitic kinzigites	144	714	793	893			
ar1083pt04	Migmatitic kinzigites	144	714	793	893			
ar1083pt04	Migmatitic kinzigites	138	710	787	887			
ar1083pt04	Migmatitic kinzigites	134	706	784	883			
ar1083pt04	Migmatitic kinzigites	127	700	777	875			
ar1083pt04	Migmatitic kinzigites	127	700	777	875			
ar1083pt04	Migmatitic kinzigites	134	706	784	883			
ar1083pt04	Migmatitic kinzigites	148	717	797	897			
ar1083pt04	Migmatitic kinzigites	127	700	777	875			
ar1083pt04	Migmatitic kinzigites	112	687	761	858			
ar1083 (average)	Migmatitic kinzigites	138	$710 \pm 10$	$787 \pm 17$	$877 \pm 20$			
ar949pt01	Migmatitic kinzigites	154	722	802	903			
ar949pt01	Migmatitic kinzigites	167	731	813	915			
ar949pt01	Migmatitic kinzigites	167	731	813	915			
ar949pt01	Migmatitic kinzigites	170	733	815	917			
ar949pt01	Migmatitic kinzigites	171	733	816	918			
ar949pt01	Migmatitic kinzigites	175	737	819	922			
ar949pt01	Migmatitic kinzigites	175	737	819	922			
ar949pt01	Migmatitic kinzigites	161	727	808	909			
ar949pt01	Migmatitic kinzigites	156	723	804	905			
ar949pt01	Migmatitic kinzigites	155	723	803	904			
ar949pt01	Migmatitic kinzigites	133	706 783		881			
ar949pt01	Migmatitic kinzigites	152	721	800	901			
ar949pt01	Migmatitic kinzigites	159	726	806	907			
ar949pt01	Migmatitic kinzigites	168	732	814	916			
ar949pt01	Migmatitic kinzigites	181	741	824	927			
ar949pt01	Migmatitic kinzigites	179	739	822	925			
		163		809	923 911			
ar949pt01	Migmatitic kinzigites		729					
ar949pt01	Migmatitic kinzigites	169	733	814	917			
ar949pt01	Migmatitic kinzigites	156	723	804	905			
ar949pt01	Migmatitic kinzigites	147	717	796	896			
ar949pt02	Migmatitic kinzigites	140	711	789	889			
ar949pt02	Migmatitic kinzigites	139	710	788	888			
ar949pt02	Migmatitic kinzigites	109	684	758	854			
ar949pt02	Migmatitic kinzigites	114	689	763	860			
ar949pt02	Migmatitic kinzigites	133	706	783	881			
ar949pt03	Migmatitic kinzigites	107	682	755	851			
ar949pt03	Migmatitic kinzigites	126	700	776	874			
ar949pt03	Migmatitic kinzigites	125	699	775	873			
ar949pt03	Migmatitic kinzigites	122	696	772	869			
ar949pt03	Migmatitic kinzigites	132	705	782	880			
ar949pt03	Migmatitic kinzigites	111	686	760	856			
ar949pt03	Migmatitic kinzigites	111	686	760	856			
ar949pt03	Migmatitic kinzigites	127	700	777	875			
ar949pt03	Migmatitic kinzigites	108	693	757	853			
ar949pt03	Migmatitic kinzigites	130	703	780	878			
ar949pt03	Migmatitic kinzigites	134	706	780 784	883			
ar949pt03	Migmatitic kinzigites	118	693	767	865			
ar949pt04	Migmatitic kinzigites	137	709	786	886			
ar949pt04	Migmatitic kinzigites	151	720	799	900			
ar949pt04	Migmatitic kinzigites	156	723	804	905			
ar949pt04	Migmatitic kinzigites	151	720	799	900			
ar949pt04	Migmatitic kinzigites	152	721	800	901			

Table 1. Continued.

Sample-crystal number	Lithology	Ti (ppm)	$T$ (°C), $P = 650 \mathrm{Mpa}$					
			Thomas et al. (2010)	Wark and Watson (2006)	Huang and Audétat (2012)			
ar949pt04	Migmatitic kinzigites	146	716	795	895			
ar949pt04	Migmatitic kinzigites	145	715	794	894			
ar949pt04	Migmatitic kinzigites	144	714	793	893			
ar949pt04	Migmatitic kinzigites	159	726	806	907			
ar949pt04	Migmatitic kinzigites	151	720	799	900			
ar949pt04	Migmatitic kinzigites	144	714	793	893			
ar949pt04	Migmatitic kinzigites	158	725	805	907			
ar949pt04	Migmatitic kinzigites	160	726	807	908			
ar949pt04	Migmatitic kinzigites	152	721	800	901			
ar949pt04	Migmatitic kinzigites	158	725	805	907			
ar949pt04	Migmatitic kinzigites	155	723	803	904			
ar949pt04	Migmatitic kinzigites	153	721	801	902			
ar949pt04	Migmatitic kinzigites	158	725	805	907			
ar949pt04	Migmatitic kinzigites	155	723	803	904			
ar949pt04	Migmatitic kinzigites	130	703	780	878			
ar949 (average)	Migmatitic kinzigites	147	$717 \pm 10$	$794 \pm 17$	$884 \pm 20$			



**Figure 8.** Calculated temperature (y axis) for each individual TitaniQ analysis (x axis = analysis number in Table 1) from the calibrations of Huang and Audétat (2012), Thomas et al. (2010) and Wark and Watson (2006).

Identification	Analysis	Thermometer	<i>T</i> (°C)	Reference	Barometer	P (Mpa)
Ar949	Core	grt-bt	802-832	Ferry and Spear (1978)	GPBQ	
	Rim	grt-bt	607-742	Ferry and Spear (1978)	GPBQ	349-447
	Core	grt-crd	797-824	Bhattacharya et al. (1988)	GPBQ	
	Rim	grt-crd	756–789	Bhattacharya et al. (1988)	GPBQ	
Ar1083	Core	grt-bt	826-842	Ferry and Spear (1978)	GPBQ	296–338
	Rim	grt-bt	633–679	Ferry and Spear (1978)	GPBQ	321–335
Ar1076c	Core	grt-bt	765–835	Ferry and Spear (1978)	GPBQ	704
Ar1076d	Core	grt-bt	721–814	Ferry and Spear (1978)	GPBQ	
	Rim	grt-bt	558-668	Ferry and Spear (1978)	GPBQ	
	Core	grt-opx	792-831	Bhattacharya et al. (1991)	GPBQ	559-612
	Rim	grt-opx	652-784	Bhattacharya et al. (1991)	GPBQ	651-652

**Table 2.** Summarized geothermobarometric results for the studied rocks.

by Ti in quartz. This suggests that post-crystallization cooling had little effect on the Ti content in quartz.

The temperatures obtained from Huang and Audétat's (2012) calibration are  $50\text{--}100\,^{\circ}\text{C}$  higher than those from the Thomas et al. (2010) and Wark and Watson calibrations (2006; Table 1). Huang and Audétat (2012) attributed the discrepancy between their calibration and Thomas et al. (2010) to the growth rate of quartz grains. Their experiments were carried out on quartz grains whose growth rates were  $4\text{--}10\,\mu\text{m}\,\text{day}^{-1}$ , and they concluded that the quartz grains of the Thomas et al. (2010) experiments, with 440  $\mu\text{m}$  to  $\sim 1\,\text{mm}$  in size, would have a growth rate 10--50 times faster ( $\sim 50\text{--}100$  to  $250\text{--}500\,\mu\text{m}\,\text{day}^{-1}$ ). According to the Huang and Audétat (2012) experiments, higher growth rates artificially increase the incorporation of titanium into quartz, and this tends to shift the resulting calibration toward lower temperatures.

We can expect grain growth in anatexites to be rather slow, especially when the cooling rate is low, and thus that temperatures computed with Huang and Audétat (2012) will be more reliable. However, considering the uncertainty that remains in the calibration of the TitaniQ geothermobarometers, in our estimate of the viscosity of the anatexites, we will use a conservative value between those obtained from the Thomas et al. (2010), Wark and Watson (2006) and Huang and Audétat (2012) calibrations. We roughly evaluate the temperature at which quartz grains started to crystalize using TitaniQ temperatures calculated from Wark and Watson (2006).

Temperatures obtained from Wark and Watson's (2006) calibration range from 700 to 810 °C, with individual average values between 750 and 800 °C (Table 1; Fig. 8). For each sample, the analyzed grains define a temperature range that likely represents the minimum temperature interval over which quartz crystallized from the magma during cooling and solidification of the anatectic domain. Data lying significantly outside this domain of temperatures and contrasting strongly with the temperature of neighboring points of

analysis may be regarded as out of equilibrium, and thus have not been taken into account further. The upper limit of this temperature interval is considered to represent the initial stage of interstitial quartz crystallization, and consequently the minimum value for the peak temperature to which the anatectic domain was submitted. Samples Ar949, Ar1083, Ar1299 and Ar1326 strongly suggest a minimum temperature of  $\sim\!800\,^{\circ}\mathrm{C}$  or slightly above. Samples Ar1258 and Ar1296 suggest a slightly lower temperature in the range 750–800 °C.

Migmatites and silicate melts containing Al-rich minerals are usually generated by partial melting of metasedimentary rocks in the crust at minimum temperatures  $\geq 700$  °C (e.g., MacRae and Nesbit, 1980; White et al., 2003). The viscosity of resulting melts has been estimated at  $\sim 10^{4.5}$  Pas (Scaillet et al., 1998) or at  $10^3$  to  $10^6$  Pas (Clemens and Petford, 1999). The viscosity of solid rocks differs from the viscosity of their melts by several orders of magnitude. Therefore, the viscosity of partially molten rocks varies widely, depending on the melt fraction, the viscosities of the molten and crystalline fractions, and on the shape and spatial distribution of solid particles (e.g., Arzi, 1978). Experiments suggest that the effect of the melt fraction on the viscosity of migmatite dominates the effects of temperature, pressure and stress, which vary less abruptly (Arzi, 1978; Rosenberg and Handy, 2005). According to Arzi (1978), a rock containing a melt fraction of  $20 \pm 10\%$  would reach the critical melt percentage (RCMP), above which, due to the connectivity of the melt fraction, its viscosity would decrease and rapidly approximate the melt's viscosity. In their compilation of experiments, Rosenberg and Handy (2005) suggest that the largest weakening resulting from partial melting occurs for melt fractions as low as 7 %.

The amount of titanium preserved in quartz is in equilibrium with the temperature prevailing when quartz crystallized from the melt. The peak temperature reached by the anatexites was therefore higher than the temperatures

**Table 3.** Summarized oxide mineral chemistry for the studied rocks, including the major element composition (wt%) of selected garnet, biotite, plagioclase, orthopyroxene and cordierite. \*-= content lower than the detection limit.

	Phase	SiO <sub>2</sub>	Al <sub>2</sub> O <sub>3</sub>	TiO <sub>2</sub>	Na <sub>2</sub> O	MgO	FeO	K <sub>2</sub> O	CaO	Cr <sub>2</sub> O <sub>3</sub>	MnO	NiO	Total
	Garnet	38.97	21.81	0.01	- 0.01	6.98	31.96	0.01	1.16	0.01	0.83	-	101.75
		38.69 38.63	21.82 21.94	0.04 0.01	0.01	7.35 7.69	31.52 30.99	_	1.17 1.22	0.01	0.75 0.77	0.01	101.35
		38.70		0.01	0.01 0.01	7.85	31.03	_	1.24		0.77		101.28 101.58
		38.78	21.91 22.04	0.02		7.83 7.84	31.05	_	1.17	0.02	0.80	- 0.01	
		38.54	21.92		-	7.68	31.26	_	1.17	0.01 0.05	0.73	0.01	101.88
		38.47	21.92	- 0.04	0.02 0.01	8.18	30.30	_	1.15	0.03	0.78	_	101.20 100.86
		38.62	21.93	0.04	- -	8.13	30.62	_	1.13	0.03	0.74	0.01	100.86
	Biotite	36.36	16.39	5.79	0.12	11.58	15.61	9.94	0.00	0.07	0.01	0.03	95.91
6		36.43	16.16	6.08	0.15	13.67	12.59	9.97	0.01	0.09	0.00	0.04	95.19
Ar949		36.93	17.43	6.13	0.37	13.33	12.39	9.43	0.03	0.10	0.03	0.03	96.18
Ā		37.09	17.18	5.11	0.21	13.91	12.47	9.88	-	0.12	_	0.02	96.00
		37.21	16.92	5.85	0.19	14.50	11.02	10.02	0.02	0.09	_	0.03	95.85
		36.45	17.32	5.29	0.34	12.85	12.40	9.32	_	0.06	0.03	0.04	94.10
		36.33	17.16	6.26	0.34	12.03	13.92	9.59	_	0.08	0.02	0.04	95.79
	Cordierite	49.33	32.99	0.01	0.11	9.32	6.49	0.01	0.01	0.01	0.05	_	98.32
		49.40	33.39	_	0.08	9.73	6.42	0.00	0.01	_	0.07	0.03	99.12
		49.05	33.03	_	0.07	9.27	6.78	0.00	_	0.01	0.07	0.02	98.32
		49.29	33.10	0.02	0.07	9.30	6.66	0.00	0.02	0.01	0.07	0.02	98.55
		49.06	33.12	_	0.10	9.24	6.83	0.01	0.01		0.08	0.01	98.45
	Plagioclase	59.77	24.92	0.01	7.78	0.01	0.02	0.26	6.74	0.01	_	_	99.52
		60.02	25.11	0.01	7.67	0.01	_	0.27	6.89	0.02	_	_	100.00
		60.15	24.93	0.03	7.71	_	0.01	0.29	6.73	_	0.01	_	99.86
	Garnet	38.60	21.81	0.03	_	7.14	32.14	-	0.93	0.02	1.08	-	101.76
		38.73	22.01	0.02	_	7.07	31.51	0.01	0.93	0.03	1.08	0.02	101.40
		38.74	21.79	0.01	_	7.21	31.66	_	0.93	0.04	1.08	0.02	101.48
		39.32	22.14	0.02	0.01	7.06	31.69	0.01	0.95	0.03	0.03	_	102.28
		38.49	21.94	_	_	7.13	31.76	_	0.91	0.04	1.09	_	101.37
~	Biotite	35.55	17.27	6.27	0.36	13.44	12.17	9.57	0.00	0.08	0.03	0.04	94.79
Ar1083		36.57	17.27	6.46	0.33	13.18	12.56	9.73	_	0.10	0.03	0.03	96.25
₹-1		36.92	17.41	6.35	0.31	12.77	12.74	9.80	_	0.07	0.03	0.01	96.41
7		36.75	16.95	5.51	0.21	11.10	15.91	9.90	0.02	0.09	0.02	0.02	96.49
		36.91	16.70	5.51	0.11	11.32	15.48	10.02	0.01	0.08	0.03	0.01	96.17
	Plagioclase	61.59	24.21	0.02	8.34	0.01	_	0.24	5.68	_	_	_	100.09
	C	62.29	23.76	0.04	8.51	_	0.02	0.31	5.23	_	0.03	0.01	100.19
		61.74	24.21	0.02	8.39	_	0.03	0.24	5.61	0.01	_	_	100.26
		61.24	24.30	0.02	8.48	_	0.09	0.16	5.87	0.02	_	_	100.19
		61.56	24.40	0.01	8.41	_	0.03	0.21	5.76	0.01	0.01	_	100.40
	Garnet	38.33	21.71	0.04	_	5.80	32.45	_	2.11	0.04	1.30	_	101.78
		38.29	21.63	0.03	_	5.29	33.24	_	1.84	_	1.36	_	101.67
Ar1076c	Biotite	36.18	16.23	4.82	0.14	10.51	18.24	9.78	0.00	0.06	0.05	0.03	96.04
r10		36.27	16.13	4.94	0.15	10.58	18.15	9.83	0.01	0.06	0.02	0.02	96.16
A		36.27	16.35	4.78	0.15	11.42	17.23	9.75	-	0.06	0.02	0.05	96.09
		36.42	16.16	4.86	0.15	10.45	18.22	9.65	_	0.05	0.03	0.02	96.02
	Plagioclase	59.97	25.40	0.02	7.60	_	0.08	0.19	7.10	0.03	0.01	_	100.40

Table 3. Continued.

Phase	$SiO_2$	$Al_2O_3$	TiO <sub>2</sub>	Na <sub>2</sub> O	MgO	FeO	K <sub>2</sub> O	CaO	$Cr_2O_3$	MnO	NiO	Total
Garnet	37.86	21.43	0.05	_	5.71	32.38	0.01	2.09	0.03	1.29	_	100.85
	37.83	21.47	0.02	0.01	4.45	34.66	0.01	1.65	0.06	1.37	_	101.53
	37.93	21.63	0.03	0.01	6.02	31.58	_	2.17	0.03	1.28	_	100.68
	38.37	21.72	0.03	_	6.27	31.58	0.01	2.11	0.03	1.23	_	101.34
	37.53	21.34	0.02	_	5.06	33.53	_	1.80	0.02	1.32	0.02	100.63
	37.35	20.99	0.03	0.03	5.08	33.86	_	1.31	0.05	1.38	_	100.08
	37.55	21.26	0.01	0.01	5.27	33.13	0.01	1.70	0.03	1.36	0.01	100.33
	39.95	22.63	_	0.01	5.36	32.10	_	1.79	0.03	1.32	_	103.19
	38.41	21.64	0.04	0.02	6.15	31.28	_	1.94	0.04	1.20	_	100.72
	38.38	21.57	0.02	0.03	6.22	31.79	_	1.95	0.01	1.25	0.01	101.24
	38.27	21.64	_	_	6.29	31.89	0.01	1.90	0.04	1.26	_	101.29
	38.17	21.71	0.02	0.01	6.29	31.95	_	1.76	0.03	1.26	_	101.20
	38.28	21.43	0.01	0.02	6.15	31.34	0.02	1.76	0.05	1.26	_	100.32
	38.38	21.70	_	_	6.18	31.90	_	1.78	0.03	1.30	_	101.27
Biotite	36.05	16.21	4.71	0.12	11.09	17.59	9.82	0.01	0.08	0.03	0.01	95.72
<del>p</del>	35.79	16.21	4.58	0.16	11.09	17.77	9.61	0.03	0.10	0.01	0.02	95.35
Ar10/6d	36.25	16.21	4.59	0.13	11.65	17.26	9.87	_	0.05	0.03	0.04	96.09
	36.24	16.20	4.56	0.13	11.55	17.39	9.85	0.05	0.06	0.03	0.02	96.09
4	36.30	16.20	4.55	0.19	13.38	14.35	9.70	0.01	0.08	0.02	0.02	94.80
	36.36	16.15	4.55	0.16	11.63	16.85	9.81	_	0.08	0.01	0.01	95.60
	36.20	16.08	4.54	0.14	11.40	17.71	9.83	0.02	0.05	0.02	0.04	96.01
	36.33	16.22	4.33	0.14	11.41	17.31	9.75	0.01	0.06	0.02	0.03	95.62
	36.42	16.22	4.32	0.10	11.33	17.01	9.79	0.01	0.06	0.03	0.03	95.33
	35.87	16.22	4.41	0.11	11.46	17.22	9.74	0.01	0.10	0.01	0.01	95.17
	36.28	16.27	4.42	0.13	11.31	17.27	9.82	0.01	0.07	_	0.03	95.58
	36.28	16.19	4.45	0.14	11.37	17.04	9.70	0.02	0.06	0.03	0.02	95.30
	35.87	16.09	4.31	0.13	11.65	17.54	9.75	0.02	0.10	0.03	0.02	95.51
Opx	49.30	2.95	0.06	0.01	15.59	30.97	_	0.15	0.02	0.49	_	99.53
_	48.99	2.91	0.09	_	15.75	30.89	0.01	0.15	_	0.49	_	99.30
	49.75	2.76	0.07	_	15.94	30.60	_	0.15	0.01	0.45	_	99.72
	49.33	3.04	0.07	_	15.67	31.17	_	0.10	0.01	0.47	0.02	99.89
Plagioclase	59.39	24.99	0.04	7.55	_	0.06	0.27	6.71	0.04	_	0.01	99.07
-	59.93	24.87	0.03	7.49	_	0.09	0.14	6.38	_	0.01	_	98.93
	59.41	24.91	0.01	7.70	_	0.03	0.31	6.70	_	0.01	_	99.06
	59.17	24.80	0.02	7.49	_	0.03	0.29	6.67	_	_	0.03	98.50

calculated using the TitaniQ calibration (750–800 °C). P-T estimates from exchange geothermobarometers suggest that the minimum temperatures and pressures for peak metamorphic conditions are recorded in samples of migmatitic kinzigites from which are derived, at least partly, the anatexites at  $\sim 800\pm 50$  °C and  $\sim 650-700\pm 100$  MPa. They are also consistent with estimates from Munhá et al. (2005;  $\sim 820$  °C, 650 MPa). A temperature of 800 °C for the anatexites seems therefore to be a conservative value for discussing their rheological characteristics further.

We have estimated the solid–liquid proportion in the neosomes of the anatexites using the rhyolite MELTS software (Gualda et al., 2012) for temperature and pressure in the ranges  $800-850\,^{\circ}\text{C}$  and  $600-700\,\text{MPa}$ ,  $H_2\text{O}$  content  $1-2\,\%$ , and bulk rock composition of seven samples of neosomes

(Table 4). The proportion of liquid in the system, inferred using these parameters, is systematically larger than 28 %, and may reach up to  $\sim 50$  %. The viscosity of the liquid phase varies in the range  $10^5$ – $10^6$  Pa s. MELTS predicts progressive crystallization of quartz (in a ratio inversely proportional to the amount of liquid in the system) with decreasing temperature. Such values are in agreement with the ubiquitous presence of films of quartz or feldspar separating inherited crystals, and also with the stromatic, schollen and locally nebulitic aspect of the anatexites (e.g., Sawyer, 2008). We may therefore consider that at conservative P–T values of 800 °C and 650 MPa, 30 % liquid present in the neosome is a reasonable value that may be used to evaluate the viscosity of the magmatic suspension. For such a proportion of solids in the magma (70 %), MELTS does not compute the viscosity

Wt%	Ar1113	Ar952	Ar1171	Ar1172	Ar1178	Ar1184	Ar1185
SiO <sub>2</sub>	72.53	72.43	71.61	72.81	71.45	69.73	71.18
$Al_2O_3$	13.78	14.59	14.31	13.89	14.29	15.11	14.59
$Fe_2O_3$	2.65	1.62	2.94	2.26	2.95	3.11	2.81
MgO	0.74	0.45	0.68	0.53	0.75	0.90	0.69
CaO	2.00	0.76	1.37	1.46	1.81	2.51	1.67
Na <sub>2</sub> O	2.42	2.57	2.46	2.55	2.52	2.85	2.73
$K_2O$	4.37	5.92	5.50	5.30	4.83	4.11	4.90
$TiO_2$	0.40	0.33	0.38	0.32	0.43	0.52	0.38
$P_2O_5$	0.19	0.28	0.12	0.19	0.21	0.21	0.24
MnO	0.05	0.01	0.04	0.03	0.05	0.05	0.04
LOI	0.8	0.9	0.5	0.6	0.6	0.8	0.7
Sum	99.89	99.86	99.92	99.93	99.89	99.86	99.90

**Table 4.** Bulk rock composition of neosomes of representative anatectic rocks.

of the system. We therefore roughly estimated this viscosity using the Roscoe (1952) equation:

$$\eta_{\rm e} = \eta (1 - 1.35\varphi)^{-2.5},$$

where  $\eta_e$  is the effective viscosity of a suspension made of a liquid of viscosity  $\eta$ , and  $\varphi$  is the solid fraction of spherical particles. The computed viscosity of the suspension (neosome) varies between  $\sim 10^8$  and  $\sim 10^{10}$  Pas for 30 % of the liquid phase, with viscosities of  $10^5$  and  $10^6$  Pas. If a rather large diversity of grain sizes is taken into account, the viscosity of the suspension decreases by about 1 order of magnitude (e.g., Kansal et al., 2002).

Restitic bodies in the anatectic domain of the Araçuaí belt, independent of their size (cm to km), are systematically immersed in neosome and anatectic granites that form a network of magma (Fig. 4) connected at the regional scale. This supports the finding that the viscosity of the partially molten crust was closer to the magma viscosity than to the solid-rock viscosity (e.g., Sawyer, 2008) and, in any case, significantly lower than the viscosity of  $10^{17}$  Pas (Grujic et al., 1996) or  $10^{19}$  Pas (Beaumont et al., 2004), below which estimates from the Himalayan orogen predict an increasing probability of large-scale horizontal flow being triggered in response to gravity forces.

The anatectic crust of the Araçuaí belt was formed during the east—west (present-day orientation) collision between the South American and African plates that lasted more than 50 Myr (Schmitt et al., 2004; Vauchez et al., 2007; Pedrosa-Soares et al., 2008; Petitgirard et al., 2009; Monié et al., 2012). Seismic receiver functions suggest that the current crustal thickness of the Araçuaí–Ribeira belt is  $\sim$  32–41 km (Assumpção et al., 2013). Considering the pressure estimates obtained in this study and by Munhá et al. (2005) for the minimum conditions for metamorphic peaks of the migmatitic kinzigites (650–700 MPa), we can infer that approximately 25 km of the continental crust thickness have been removed. It follows that the crust thickness reached at least 60 km during the continent—continent collision, suggesting high to-

pography. Such a thick crust, an abnormally high geotherm  $(> 30 \, {}^{\circ}\text{C km}^{-1})$  and a slow cooling rate  $(< 5 \, {}^{\circ}\text{C Myr}^{-1}; \text{ Pe}$ titgirard et al., 2009) resulted in widespread partial melting of the middle to lower crust and a decrease in the thermal conductivity of the middle crust, enhancing abnormally high temperatures. These conditions dropped the viscosity of the neosome to  $\sim 10^{10} \, \mathrm{Pa} \, \mathrm{s}$ , supporting the finding that the viscosity of the anatectic middle crust was low enough ( $\ll 10^{17} \, \text{Pa s}$ ) to enable a gravity-induced component of flow. Considering the cooling rate, the large volume of melt produced during orogeny, and the latent heat of crystallization release by this anatectic layer, we can hypothesize that this low-viscosity layer remained so over a very long period (tens of Myr), its viscosity increasing slowly with the decrease in temperature. This supports the proposal that the gravitydriven deformation may have combined with deformation due to east-west convergence between the South American and African proto-continents, resulting in a complex 3-D flow of the anatectic middle crust.

#### 8 Conclusions

Temperature estimates from the middle crust of the Araçuaí belt acquired using the titanium-in-quartz geothermometer in anatectic rocks are similar to those obtained from exchange geothermometry in neighboring kinzigites that have mineral assemblages suitable for such analysis. This good agreement supports that the TitaniQ geothermometer provides reliable crystallization temperatures of quartz in anatectic rocks. These estimates point to minimum temperature and pressure conditions for metamorphic peaks above 800 °C and 650–700 MPa. Solid/liquid fractions computed using conservative P and T values and the chemical compositions of several anatexites suggest that at least 30 % of the melt was present when the middle crust reached 800 °C. The proportion of melt under peak metamorphism conditions during orogeny was certainly higher than 30%, a proportion consistent with the presence of in situ leucosomes and stromatic, nebulitic and schlieren structures in the anatexites. Such a proportion of melt is large enough to promote melt interconnectivity, as mentioned by Arzi (1978) and Rosenberg and Handy (2005), for instance. Rough estimates suggest that the partially molten middle crust had a viscosity significantly lower than the 10<sup>17</sup>–10<sup>19</sup> Pas that is suggested by Grujic et al. (1996) and Beaumont et al. (2004) as the upper limit below which a rheologically weak crust is prone to being deformed through gravitational flow during orogeny. Therefore, considering the size ( $\sim 300 \, \text{km} \times 50\text{--}100 \, \text{km}$ ) of the anatectic domain of the Araçuaí belt, the large orogenic thickness of the crust, the abnormally high temperature gradient, the low cooling rate, and the transient low viscosity of the partially molten middle crust, the complex flow pattern that characterizes the anatectic domain of the Araçuaí belt (e.g., Cavalcante et al., 2013) may result from the deformation of the low-viscosity middle crustal anatectic layer through gravitydriven escape flow associated with convergence-driven short-

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