





# Evolução geodinâmica da Faixa Ribeira (SE do Brasil) baseada no estudo de inclusões fluidas e na modelação da $fO_2$

Geodynamic evolution of Ribeira Fold Belt (SE Brazil) based on fluid inclusion studies and  $fO_2$  modelling

T. Bento dos Santos<sup>1</sup>, J. Munhá<sup>1</sup>, C. Tassinari<sup>2</sup>, F. Noronha<sup>3</sup>, A. Guedes<sup>3</sup>, P. Fonseca<sup>4</sup>, C. Dias Neto<sup>2</sup>, A. Dória<sup>3</sup> (tmsantos@fc.ul.pt)

## **SUMÁRIO**

A aplicação de diversas metodologias ao estudo das inclusões fluidas e da fugacidade do oxigénio permitiu a caracterização do percurso tardio dos granulitos da Faixa Ribeira. Após o pico metamórfico, uma rápida exumação das rochas a T  $\sim$  450  $^{\circ}\text{C}$  levou à formação de inclusões fluidas de CO $_2$  pouco densas, seguidas do influxo de água no sistema a P < 1 kbar, passando o sistema de oxidado para reduzido, o que provocou a precipitação de grafite em migmatitos.

Palavras-chave: Faixa Ribeira; inclusões fluidas; fugacidade do oxigénio; evolução P-T-fluido

# SUMMARY

Several procedures applied to fluid inclusion and oxygen fugacity studies allowed to characterize the late retrograde path of Ribeira Fold Belt. After metamorphic peak cooling, at about 450  $^{\circ}\text{C}$ , a significant pressure drop occurred, leading to low-density CO<sub>2</sub> inclusion formation, followed by the influx of water at P < 1 kbar, which turned the system from highly oxidized to reduced and caused the precipitation of graphite in migmatites.

Key-words: Ribeira Belt; fluid inclusions; oxygen fugacity; P-T-Fluid evolution

### Introduction

Although fluid inclusion studies have long been a concern among metamorphic geologists to unravel the mysteries of the lower crust (Touret, 1971), work has yet to be done in order to understand the dynamics of fluid evolution in the Braziliano Cycle. Thus, this work addresses T, P,  $fO_{2}$ , age, origin and evolution of fluids in the Ribeira Belt in order to constrain the retrograding P-T-Fluid path of this granulitic belt

# Geologic setting and field observations

The studied São Fidelis – Santo António de Pádua (SFSAP) sector is located in the central-north Ribeira Belt, SE Brazil. The Ribeira Belt is

a NE-SW to NNE-SSW trending Neoproterozoic belt formed in the Braziliano Orogeny by the collision of the São Francisco and West Congo cratons. from which resulted Western Gondwana (Cordani, 1971). Ribeira Belt is a complex orogenic belt composed of several geological units, separated by deep dextral shears. The SFSAP sector is located SE to one of these mega-shears, the Além Paraíba -Santo António de Pádua shear (APPS) that vigorously deformed the area rocks imposing a NĒ-SW trending transpressive shear high-grade deformation associated with granulite facies metamorphism, producing generalized migmatization. Outcrops in the area comprise: migmatitic paragneisses (1)

<sup>&</sup>lt;sup>1</sup>Centro/Departamento de Geologia, Universidade de Lisboa, Edifício C6, 2°, Campo Grande, 1749-016 Lisboa

<sup>&</sup>lt;sup>2</sup>Instituto de Geociências, Universidade de São Paulo, Rua do Lago, 562 – Butantã, CEP: 05508-080, SP, Brasil

<sup>&</sup>lt;sup>3</sup>Centro de Geologia, Universidade do Porto, Rua do Campo Alegre, 687, 4169-007 Porto

<sup>&</sup>lt;sup>4</sup>LATTEX, Universidade de Lisboa, Edifício C6, 2°, Campo Grande, 1749-016 Lisboa

(metatexites), commonly interlayered with amphibolites and marbles; (2) diatexites; (3) massive and incipient-type charnockites; and (4) blastomilonites that resulted from late retrogression of the major rock types. Also present in the area are khondalites (graphitic gneisses) that resulted from incipient charnockitization of metatexites in areas of contact with charnockites.

## Oxygen Fugacity calculations

Oxygen fugacity was determined for 6 charnockites, 1 migmatite, 3 blastomilonites and 1 amphibolite, using the QUILF algorithm (Andersen & Lindsley, 1988). MH (Magnetite-Hematite) temperature determinations ranged from 370 to 771 °C, indicating post-metamorphic Ti-magnetite oxidation in accordance with observation of exsolution of

ilmenite from magnetite. Thus, Ti-magnetite compositions were reconstructed at the appropriate P-T range (Bento dos Santos et al., 2006) in order to estimate the metamorphic  $fO_2$ MH, conditions. OHQ (Orthopyroxene-Hematite-Quartz) and AHQ (Augite-Hematite-Quartz) (Harlov, 1992)  $fO_2$  estimates range from  $10^{-11.538}$  to  $10^{-17.799}$  bar for the calculated temperature range of 896 to 656 °C. Figure 1 shows that high-T charnockites amphibolites) have fO2 values above the QFM buffer (QFM +1), whereas migmatites and blastomilonites provide fO2 at QFM -1. Thus, the inferred fO<sub>2</sub> evolution suggests that fluids experienced metamorphic relative reduction during cooling. This is consistent with field and petrographic observations indicative of late graphite deposition in khondalites.

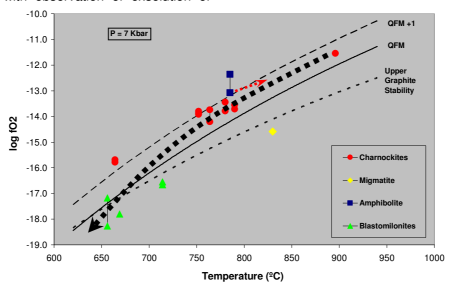


Fig.1 –  $fO_2$  of studied samples at respective metamorphic peak temperatures (Bento dos Santos et al., 2006).

## Fluid Modelling

Fluid modelling in the C-O-H system was performed in order to determine fluid compositional variations ( $H_2O$ ,  $CO_2$ , CO,  $CH_4$  and  $H_2$ ) at a given T, P and  $fO_2$ . Results show that  $CO_2$ -rich fluid inclusions should be stable during charnockite formation (at the estimated P, T,  $fO_2$  conditions), whereas aqueous fluids (with minor  $CH_4$  and  $CO_2$ ) were dominant in migmatites (and blastomilonites).

# Fluid Inclusion (FI) and Raman Studies

Fluid inclusions were analysed in 4 charnockites, 1 diatexite, 2 migmatites, 2 khondalites and 2 blastomilonites (amounting to several hundred measurements in quartz and garnet crystals). Inclusions are typically < 10  $\mu$ m in size and are referred as primary, secondary or late, according to their relative textural relations following Roeder (1984) classification. On this basis 6 fluid inclusion

groups were defined, and their evolving characteristics are summarized in Table 1. G3a is the most common FI type; G4 FI group is characteristic of khondalites (that lack G2 and G3 FI).

# **Graphite analysis**

Graphite is a common occurrence in granulite facies meta-sediments as the result of conversion of organic matter into crystalline graphite. Because mineral structure of graphite cannot be retrogressed and is, therefore, mainly dependent on temperature, it can be used to estimate crystallization temperatures (Pasteris & Wopenka, 1991). The use of analytical procedures that evaluate its mineral structure, such as Raman Spectroscopy and X-ray diffraction, and the use of appropriate formulation (Beyssac et al, 2002) supplied T estimates in the 333 to 449 °C range for graphites in the studied khondalite samples.

Table 1: Summar	v of fluid inclusion microthermometr	y and Raman Spectroscopy results.

Group	Composition	Phases	Occurrence	Flw	Tm CO <sub>2</sub>	Tm Ice	Th CO <sub>2</sub>	TH	CO <sub>2</sub>	N <sub>2</sub>	CH <sub>4</sub>	d (g/cm <sup>3</sup> )	Salinity (Wt% Eq. NaCl)
1	N <sub>2</sub> -CH <sub>4</sub>	Mono	Р	-	-	-	=	-	-	94 - 95	5 - 6	-	-
2a	CO <sub>2</sub> ; CO <sub>2</sub> -N <sub>2</sub>	Mono	Р	-	-58.1 : -59.6	-	-16.3 : 6.2 (L)	-	92 - 100	0 - 8	-	0.86 - 1.01	-
2b	CO <sub>2</sub> ; CO <sub>2</sub> -N <sub>2</sub>	Mono	P or S	-	-58.5 : -63.2	i	6.4 : 10.9 (L)	-	89 - 100	0 - 11	-	0.79 - 0.86	-
2c	CO <sub>2</sub> ; CO <sub>2</sub> -N <sub>2</sub>	Mono e Bi	P or S		-58.4 : -62.2	-	13.4 : 30.1 (L)	-	94 - 100	0 - 6	-	0.59 - 0.81	-
3a	CO <sub>2</sub> ; CO <sub>2</sub> -N <sub>2</sub>	Bi	P or S	-	-57.2 : -59.5	-	17.3 : 31.0 (C)	-	64 - 100	0 - 36	-	0.19 - 0.29	-
3b	N <sub>2</sub> -CO <sub>2</sub> ; N <sub>2</sub>	Mono	P or S	-	-	-	-	-	0 - 30	70 - 100	-	-	-
4	CO <sub>2</sub> -N <sub>2</sub> -CH <sub>4</sub> -H <sub>2</sub> O	Bi	P or S	0 - 0.1	-60.0 : -62.8	-	8.7 : 19.0 (L)	-	94 - 95	3	2 - 3	0.73 - 0.82	-
5	CO <sub>2</sub> -H <sub>2</sub> O	Bi	S - Late	0.3 - 0.7	-58.8 : -59.7	-3.7 : -5.4	9.5 : 13.1 (L)	232 : 404 (L)	100	-	-	0.56 - 0.99	6.1 - 10.5
6a	H <sub>2</sub> O	Bi	Late	0.6 - 0.95	=	-0.1 : -4.5	=	86: 367 (L)	-	-	-	0.57 - 0.93	0 - 7.2
6b	H <sub>2</sub> O	Bi	Late	0.9 - 0.95	-	-4.0 : -9.3	-	98: 174 (L)	-	-	-	0.97 - 0.99	6.5 - 13.2

## P-T-Fluid evolution

FI microthermometry indicates that the SFSAP sector rocks evolved in equilibrium with  $N_2 \pm CH_4$  and  $CO_2\text{-}N_2$  rich fluids at high metamorphic temperatures. During the retrograding path fluids became progressively enriched in water, generating  $CO_2\text{-}H_2O$  fluids and late low-salinity

H<sub>2</sub>O fluids. Representative FI were used for isochore calculations presented in Fig. 2. Observation of the P-T-Fluid evolution shows that all FI are late, (relative to the peak of metamorphism), being trapped during cooling and decompression (exhumation) of their host metamorphic rocks.

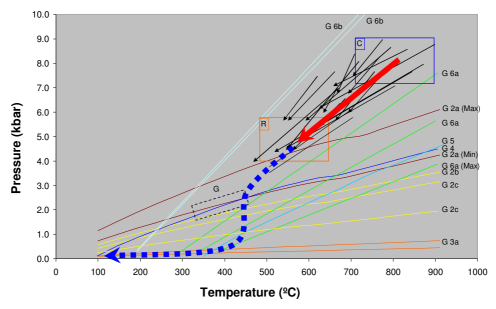


Fig.2: P-T-Fluid evolution (C and R: core and rim temperature estimates (Bento dos Santos et al., 2006); G: Graphite temperature estimates for this study).

## **Discussion**

 ${\rm CO_2}$  is the most oxidized fluid in the C-O-H system and its influx into lower crust from deep-seated sources has been advocated to explain charnockite development (Newton et al., 1980), which is consistent with the oxidized conditions estimated for SFSAP charnockites. However, Touret (1971) and Cesare et al. (2005) argued that  ${\rm Fe}^{3+}$  reduction during biotite dehydration-melting could cause graphite oxidation, producing  ${\rm CO_2}$  and globally rising  $f{\rm O_2}$  (if water is leaving the system). Thus, early  ${\rm CO_2}$ 

predominance in SFSAP fluids is interpreted as a result of relative concentration of the least mobile fluids, whereas water is preferentially removed by ascending melts. This process would also induce relative oxidation, as estimated for the studied charnockites.

The late P-T-Fluid path involved cooling and decompression until about 450 °C, followed by a significant pressure drop (probably associated with orogenic collapse). Indeed, graphite deposition in khondalites is a relatively late process that took place after significant cooling down to 450 - 330 °C. Accordingly, graphite

deposition should be coeval with late tectonic (orogenic collapse) and imbrication and consequent cooling decompression, enhancing permeability and admixture of reducing H<sub>2</sub>O-rich fluids into the system. This stage is related to the formation of (early) lowdensity CO<sub>2</sub> fluid inclusions (G3a in Fig. 2) and (late) low-salinity H<sub>2</sub>O fluids, as the rock pile progressively approached the surface. interacting with shallow aguitards/aguifers.

### **Conclusions**

Fluid evolution reflects compositional readjustments related to rapid decompression and cooling during the late stages of the Ribeira Belt exhumation path. Results indicate that high-T (> 550  $^{\circ}$ C) fluids were dominated by CO<sub>2</sub> – N<sub>2</sub> components. At 450  $^{\circ}$ C rocks were already exhumed to 3–10 km depths, producing generalized low-density CO<sub>2</sub> (+ H<sub>2</sub>O) inclusions, followed by interaction with shallower aquifer waters. fO<sub>2</sub> decreased substantially during cooling and mixture of CO<sub>2</sub> and H<sub>2</sub>O, causing late graphite deposition.

Incipient charnockitic development by "CO<sub>2</sub> influx" is possible for some khondalites, but this process does not explain the massive charnockite formation in Ribeira Belt. We suggest that CO<sub>2</sub>-rich, high-T metamorphic fluids should have resulted mainly from CO<sub>2</sub> concentration after water removal to ascending granitic melts, as originally proposed by Fyfe (1973).

## Acknowledgements

FAPESP, POCA-PETROLOG (CEGUL, UI: 263; POCTI/FEDER), GEODYN (POCTI – ISFL – 5 – 32) and a scholarship from FCT (SFRH/BD/17014/2004) financed field and analytical procedures. The authors would also like to thank the following colleagues: Teresa Palácios, Álvaro Pinto, Nuno Leal and Línia Martins.

### References

Andersen, D., Lindsley, D. H., 1988. Internally consistent solution models for Fe-Mg-Mn-Ti oxides: Fe-Ti oxides. Am. Min., 73, 714-726.

Bento dos Santos, T., Munhá, J., Tassinari, C., Dias Neto, C., Fonseca, P., 2006. Petrologia, Geoquímica e Geocronologia de Granulitos no sector São Fidélis – Santo António de Pádua, RJ, SE Brasil. Ext. Abst., VII Congresso Nacional de Geologia, 1, 241-244.

Beyssac, O., Goffé, B., Chopin, C., Rouzaud, J. N., 2002. Raman spectra of carbonaceous material in metasediments: a new geothermometer. Jour. Met. Geol., 20, 859-871.

Cesare, B., Meli, S., Nodari, L., Russo, U., 2005. Fe3+ Reduction during biotite melting in graphitic metapelites: another origin of CO2 in granulites. Contrib. Min. Pet., 149, 129-140.

Cordani, U. G., 1971. Síntese da geocronologia Pré-Cambriana da região costeira atlântica meridional da América do Sul. Ext. Abst., 25th Congresso Brasileiro de Geologia, 179-180.

Fyfe, W. S., 1973. The granulite facies, partial melting, and the Archaean crust. Phil. Trans. Royal Soc. London, A273, 457-461.

Newton, R. C., Smith, J. V., Windley, B. F., 1980. Carbonic metamorphism, granulites and crustal growth. Nature, 288, 45-50.

Pasteris, J. D., Wopenka, B., 1991. Raman spectra of graphite as indicators of degree of metamorphism. Can. Min., 29, 1-9.

Roeder, E., 1984. Fluid Inclusions. Mineralogical Society of America, Reviews in Mineralogy, 12, 644.

Touret, J., 1971. Le facies granulite en Norvège méridionale. II. Les inclusions fluides. Lithos, 4, 423-436

Touret, J., 1981. Fluid inclusions in high grade metamorphic rocks. In: Hollister, L. S., Crawford, M. L. (Eds.) Short Course in Fluid Inclusions. Mineral. Assoc. Canada, Calgary, 182-208.

Valley, J. Bohlen, S., Essene, E., Lamb, W., 1990. Metamorphism in the Adirondacks: II. The role of fluids. Jour. Pet., 31, 3, 555-596.

Valley, J., McLelland, J., Essene, E., Lamb, W., 1983. Metamorphic fluids in the deep crust: evidence from the Adirondacks. Nature, 301, 226-228.