Petrographic and geochemical studies at giant Serra Norte iron ore deposits in the Carajás mineral province, Pará State, Brazil

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ABSTRACT: The Carajás iron ore deposits, located in the southern part of the state of Pará in Brazil, are hosted by the metavolcanosedimentary sequence of the Grão Pará Group, Itacaiúnas Supergroup. The protoliths to iron mineralization are jaspilites, under- and overlaid by basalts, both greenschist facies metamorphosed. The major Serra Norte N1, N4E, NAW, NSE and N5S iron ore deposits of the Carajás Mineral Province are distributed along, and structurally controlled by, the northern flank of the Carajás fold. High-grade iron mineralization (> 64 % Fe) is made up of hard and soft ores. The basal contact of the high-grade iron ore is defined by a hydrothermally altered basaltic rock mainly composed by chlorite and microplaty hematite. Varying degrees of hydrothermal alteration have affected jaspilites to form iron ores, from distal alteration zone, representing an early alteration stage, to intermediate and proximal alteration zones, synchronous with the main iron-ore forming event. The latter represents an advanced alteration stage (i.e., the high-grade iron ore). Jaspilites from the N4W, NSE and N5S deposits, and hard ores from N1 and N4E have a low ∑REE content, are enriched in light REE and exhibit positive europium anomalies (Eu/Eu*> 1), which is typical of Archean banded iron formations. The REE pattern defined by NSE ores is nearly flat and displays an increase in ∑REE and absence of the positive Eu anomaly. The increase in LREE was accentuated during the formation of magnetite and microplaty hematite, and the advance of martitization to form anhedral hematite, which may have favoured the relative increase of HREE in the residual fluid, resulting in an increase in HREE in advanced-stage precipitates and almost flat REE patterns associated with the advanced stage of mineralization (euhedral and tabular hematite formation). The mineralogical, geochemical and isotopic changes from jaspilites to high-grade iron ores suggest a hydrothermal origin for hard ore via interaction with an early-stage, relatively reduced magmatic fluid, which leached silica and formed magnetite, which evolved to more oxidizing conditions, with the advance of martitization, increase in the REE concentration and microplaty hematite precipitation in veins and martite borders, from interaction with modified meteoric waters.

Keywords: jaspilite, iron ore, Carajás, rare earth elements

RESUMO: ESTUDOS PETROGRÁFICOS E GEOQUÍMICOS DOS GIGANTES DEPÓSITOS DE MINÉRIO DE FERRO DA SERRA NORTE, PROVÍNCIA MINERAL DE CARAJÁS, PARÁ, BRASIL. Os depósitos de minério de ferro de Carajás, localizados no sudeste do estado do Pará no Brasil, estão hospedados na sequência metavulcanosedimentar do Grupo Grão Pará, Supergrupo Itacaiúnas. Os protólitos da mineralização de ferro são jaspilitos, soto sobrepostos a rochas basálticas, ambos meiramorizados em fácies xisto verde. Os maiores depósitos de minério de ferro da Serra Norte são N1, N4E, NAW, NSE e N5S, estão distribuídos ao longo e estruturalmente controlados pelo flanco norte da dobra Carajás. Minério de alto teor (> 64 % Fe) consiste em minérios compactos e fráveis. O contato basal dos minérios de alto teor é definido por rocha basáltica alterada hidrotermalmente, composta por clorita e hematita microlamelar. Diferentes estágios de alteração hidrotermal afetaram jaspilitos para formar minérios de ferro, da zona de alteração distal, representada pelo estágio cedo-hidrotermal, às zonas de alteração intermediária e proximal, concomitantes com o evento principal de formação de minério de ferro. A zona proximal representa o estágio de alteração avançada (i.e., minério de ferro de alto). Jaspilitos dos depósitos N4W, NSE e N5S e minérios compactos dos depósitos N1 e N4E têm baixo conteúdo ∑ETR, são enriquecidos em REE leves e exibem anomalias positivas de európio (Eu/Eu*> 1), padrão típico de formação de jaspilites bandeadas arqueanas. O padrão de ETR definido por minérios de NSE é quase horizontal e apresenta aumento em ∑ETR e ausência de anomalia positiva de Eu. O aumento acentuado em ETR ocorreu durante a formação de magnetita e hematita microlamelar. Já aumento em ETRP e padrão quase horizontal de ETR coincidem com o avanço da martitização formando hematita anidrítica, o qual pode ter favorecido o aumento relativo de ETR pesados no fluido residual, resultando em precipitados do estágio avançado de mineralização (formação de hematitas euédricas e tabulares). As mudanças mineralogicas, geoquímicas e isotópicas de jaspilitos a minérios de ferro de alto teor sugerem uma origem hidrotermal para minério compacto via interação de fluido magmático reduzido em estágio cedo-hidrotermal, lixiviando silicia e formando magnetita. Este fluido evoluiu para condições mais oxidorantes, com avanço da martitização, aumento na concentração de ETR e formação de hematita microlamelar em veios e bordas de martita, a partir de mistura com águas meteóricas modificadas.

Palavras-chave: jaspilite, minério de ferro, Carajás, elementos terras raras

1. INTRODUCTION

The Carajás Mineral Province (CMP) is located in the eastern part of the Amazon craton, east-southeastern Pará State, Brazil (Fig.1), and is considered one of most important mineral provinces of the world, with production and growing potential for Fe, Mn, Cu, Au, Ni, U, Ag, Pd, Pt, Os, Zn and W. The iron ores are hosted by the Carajás Iron Formation, which is interbedded with mafic volcanic rocks of the Grão Pará Group of the Neoarchean Itacaiúnas Supergroup (Fig. 1). The Grão Pará Group outcrops along the northern and southern flanks of a synclinal structure, known as the Carajás fold, with a major shear zone known as the Carajás shear zone along the axis of the fold. Iron ore deposits developed along the northern flank of the fold, known as the Serra Norte (Northern Range) deposits and along the southern flank known as the Serra Sul (Southern Range) deposits (Fig. 1). At present, only
some of the Serra Norte deposits are mined with extensive exploration taking place in some of the Serra Sul deposits. There are nine deposits in the Serra Norte, numbered N1 – N9 (Figs. 1 and 2), with strike lengths from hundreds of meters to tens of kilometers and some of them comprise two or more orebodies. The combined resources of the nine deposits are 17.3 billion tons at 66 wt. percent Fe (CVRD, 2007), with orebody thicknesses varying from approximately 250 to 300 meters. Four main open pits (Figs. 2A and B) have, to date, produced about 1.2 billion tons of high-grade iron ore, which made Carajás the second largest Brazilian producer of iron following the Quadrilátero Ferrífero (Iron Quadrangle) region of Minas Gerais State (Rosière et al., 2008).

Jaspilites constitute the protore in many iron ore deposits of the Carajás region (Meirelles, 1986). The term jaspilite was first applied in the Lake Superior area to name oxide-facies iron-formation in which silica is present as jasper (UNESCO, 1973). In Carajás, the term jaspilite was first used by Suszczynski (1972) to describe the banded iron formations (BIFs) in the Serra Norte deposits (Fig. 1). This type of BIF consists of bands of chert with hematite dust (i.e. jasper) intercalated with iron oxides. In this paper, and other contributions to the Carajás iron ore mineralization (e.g. Figueiredo e Silva, 2004; Figueiredo e Silva et al., 2007a; b; 2008; Hagemann et al., 2006; Lobato et al., 2005a, b; 2008, Zuccheti, 2007; Zuccheti et al., 2007), we use the term jaspilite as the BIF type described in the Serra Norte mines and surroundings. This BIF type is clearly different from BIF found in other parts of the Carajás area. A volcanogenic origin for these jaspilites is indicated by Meirelles (1986), Dardenne & Schobbehaus (2001, and references therein) and Klein & Ladeira (2002) based on the Grupo Grão Pará tectonic environment and/or the geochemical characteristics of the jaspilites. However, we stress here that we use jaspilite as a descriptive term with no genetic connotation.

The Carajás Range iron ore occurrences were discovered in August 1967 by Companhia Meridional de Mineração, the pilot area being the current N1 deposit (Beisegel et al., 1973). In 1969, 18 mineral exploration license areas were claimed by Companhia Meridional de Mineração and Companhia Vale do Rio Doce (CVRD – now named Vale) encompassing some 160 000 ha area. From 1970 to 1972, an intensive exploration program was executed (Beisegel et al., 1973). Open pit mining of the N4E deposit started in 1984, followed by the N4WC (central northwestern 4 orebody) in 1994, the N4WN (north northwestern 4 orebody) in 1996, and the NSW and NSE deposits in 1998 (Huhn et al., 2000). Presently, all iron ore deposits belong entirely to Vale, the world’s largest producer of iron ore (CVRD 2007). At the moment, the S11 orebody in the Serra Sul is the subject of an intensive exploration campaign.

Similar to other BIF iron ore districts in Brazil, such as the itabirites in the Iron Quadrangle in Minas Gerais State, jaspilites hosts soft (i.e. high porosity) and hard (i.e. low porosity) high-grade ores. Both supergene and hypogene genetic models explaining the formation of iron ores have been suggested (Morris, 1985). During the past 12 years, many research groups have studied iron ore deposits in Australia (Barley et al., 1999; Hagemann et al., 1999; Taylor et al., 2001; Thorne et al., 2004; Angerer & Hagemann, 2010), South Africa (Beukes et al., 2002; Netshiozwi, 2002; Gutzmer et al., 2006) and Brazil (Varajão et al., 2002; Cabral et al., 2003; Spier et al., 2003; Rosière and Rios, 2004; Rosière et al., 2008). Although these deposits contain hypogene iron mineralization, with silica leaching, with or without iron addition, most authors agree that supergene fluids are involved in the iron ore forming process as well. A few studies have proposed a hydrothermal origin for the Carajás hard iron ores, with hydrothermal introduction of e.g., hematite, carbonate and sulfide (Guedes, 2000; Guedes et al., 2002; Figueiredo e Silva, 2009; Figueiredo e Silva et al., 2008; Dalstra and Guedes, 2004; Lobato et al., 2005a, b). There is significant evidence, based on constraints of the paragenetic sequence and textural features, for the contemporaneous emplacement of veins and hydrothermal alteration in the jaspilite and also in the mafic wall rocks, which caused the transformation of jaspilites to high-grade ores, i.e., the mineralization process(es). This paper is only concerned with the formation of hard ores considered to be of hypogene origin and provides: (1) a reevaluation of the regional setting of the CMP, (2) descriptions of the main iron ore deposits, (3) the petrography and geochemistry of the protore and hard iron ores, (4) hydrothermal alteration zonation including vein classification and minerals that formed during the transformation of jaspilite to hard ores, and (6) an assessment of previous genetic models.

2. GEOLOGICAL SETTING OF THE CARAJÁS MINERAL PROVINCE

The Carajás Mineral Province (CMP) is divided into two Archean tectonic blocks, the southern Rio Maria granite-greenstone terraine (Huhn et al., 1988) and the northern Itacaúnas Shear Belt (Araújo et al., 1988). The CMP is characterized by: (1) the giant iron ore deposits of the Serras Norte (N1 to N9), Sul (S1 to S45) and Leste, (2) the iron oxide-copper-gold (IOCG) deposits of Salobo, Pojuca, Alemão-Igarapé Bacia, Cristalino, Sossego, Alvo 118, and Gameleira; (3) the
gold-palladium deposit at Serra Pelada, (4) the manganese deposits at Azul and Sereno; and (5) the nickel deposit at Vermelho (Fig. 1). The area covered by these deposits is characterized by metavolcano-sedimentary rocks, and Meso- to Neo-Archean basement, which consist of igneous suites and metamorphic complexes (Santos, 2003; Tassinari et al., 2000).

In the northern Itacaiúnas Shear Belt, granite-gneissic terrains, such as the tonalite-trondhjemite-granodiorite (TTG) Xingu Complex (Silva et al., 1974), form the basement underlying the volcano-sedimentary sequence. Many Archean granites intrude the basement and cover volcano-sedimentary sequences, such as the granites and diorites of the 2.74 Ga Plaquê Suite (Fig. 1) (Huhn et al., 1999), and the foliated 2736 ± 24 Ma Planalto alkali granite (Avelar et al., 1999; Huhn et al., 1999). Mafic-ultramafic intrusive rocks are represented by the 2763 ± 6 Ma Luanga suite (Machado et al., 1991).

Fig. 1. Geological map of the Itacaiúnas Shear Belt, Carajás Mineral Province (after Bizzi et al., 2001) showing major mineral resources including iron ore deposits N1, N4 and N5 (Serra Norte), Serra Sul and Serra Leste; some IOCG deposits (Gameleira, Allemão/Igarapé Bahia, Alvo 118, Sossegio, Cristalino); the Serra Pelada gold-palladium deposit; the Azul manganese deposit, and the Vermelho nickel deposit. Lithostratigraphic classification is adapted with field data from Costa (2007) and interpretation by Seoane et al., (2004), based on Landsat ETM7 RGB 321, 752 and PC1-52 images.
The Serra dos Carajás, the S-shaped mountain belt that contains the major iron ore deposits, itself is composed of metavolcano-sedimentary and metasedimentary rock units of the Itacaiúnas Supergroup (Fig. 1). In the northern part, the Itacaiúnas Supergroup includes greenschist facies metavolcano-sedimentary and metasedimentary sequences of the Igarapé Bahia (2745 ± 1 and 2747 ± 1 Ma; Galarza & Macambira, 2002), Aquiri, Grão Pará (Santos, 2003) and Rio Novo (e.g. Costa, 2007) Groups. In the south it includes the amphibolite-facies Igarapé Salobo and Igarapé Pojuca Groups (e.g., Lindenmayer, 1990). Despite their tectono-metamorphic overprint, these rocks commonly exhibit low-strain features, and original structures and textures may be preserved. Igneous or sedimentary terminology is widely used for these rocks and the prefix ‘meta’ is implicit but omitted for ease of reading in this manuscript.

The Grão Pará Group (Fig. 1) is a typical greenstone-belt succession composed of volcanic and sedimentary rocks with ages between 2.8 and 2.7 Ga (Gibbs et al., 1986; Wirth et al., 1986; Olszewsky, 1989; Macambira et al., 1996; Trendall et al., 1998) with interbeds of BIF. The lower unit of this Group, the Parauapebas Formation (Meireles et al., 1984), consists mainly of basalts and basaltic andesites, and minor basic to intermediate pyroclastic rocks. An amygdaloidal facies is found closer to the top of the unit (Meireles et al., 1984) with BIF and basic tuff intercalations. In the basal portions, rhyolite layers are found together with fluvial conglomerates and arenite. The rocks were subjected to regional greenschist metamorphism and hydrothermal alteration that caused iron enrichment (Figueiredo e Silva et al., 2008; Zucchetti, 2007). The pyroelastic or volcanoclastic rocks preserve ocean-floor hydrothermal minerals, and this suggests very low-grade metamorphism (Zucchetti & Lobato, 2004; Zucchetti, 2007). Parauapebas Formation metabasalts that are devoid of hydrothermal alteration are composed of actinolite, chlorite, epidote, quartz, and calcite, also compatible with greenschist facies metamorphism (Meirelles, 1986; Macambira et al., 1990; Teixeira & Egger, 1994; Teixeira et al., 1997).

The middle part of the Grão Pará Group is formed by the BIFs of the Carajás Formation (2751 ± 4 Ma; Krimsky et al., 2002), which host the giant iron ore deposits (Tolbert et al., 1971, 1973; Gibbs & Wirth, 1990). The Carajás Formation, defined in 1972 (DOCEGEO, 1988), contains layers and discontinuous lens-shaped bodies, or lenses, of jaspilites and high-grade iron ores, intruded by mafic sills and dikes. The mafic sills and dikes at the N4E deposit, varies from centimeters to tens of meters in width (Macambira & Silva, 1995). Jaspilites display typical intercalation of centimeter-thick, light and dark layers, or mesolayers (about 5 cm), and microlayers with iron-oxide layers intercalated with reddish to light-colored layers comprised of jasper and chert, respectively. Primary structures and textures in the jaspilites such as syn-sedimentary microfaults and spherulites (jasper or chert ringed by hematite) are still observed, despite the greenschist grade of metamorphism (Meirelles, 1986). Dolomite-bearing iron formations consist of layers of dolomite and chert in varying proportions, alternating with dark iron-oxide layers (Macambira & Schrank, 2002; Dalstra & Guedes, 2004). The dolomite-bearing iron formations are approximately 50 meters thick extending for about 400 m along strike.

In the central part of the Serra dos Carajás, the volcano-sedimentary rocks are covered by sedimentary psammo-pelitic rocks (e.g. arenites, calcarenites, siltites and conglomerates) of the Águas Claras Formation (Fig. 1), which yielded zircon U-Pb ages of 2708 ± 37 Ma (Mougeot, 1996) and 2645 ± 12 Ma (Dias et al., 1996).

Paleoproterozoic A-type, alkaline to subalkaline granites such as the Serra dos Carajás granite (Fig. 1) intrude the Itacaiúnas Supergroup and have an age of about 1880 ± 2 Ma (Machado et al., 1991). Dall’Agnoll & de Oliveira (2007) have classified these as oxidized, magnetite-series, rapakivi-type granites.

A Paleoproterozoic age for the iron mineralization has been constrained by whole-rock Sm-Nd data on hematitized mafic volcanic (basalt) wall rocks, along the contact with the high-grade iron ore (Lobato et al., 2005b). More recently, Santos et al. (2010) obtained two distinct ages using hydrothermal minerals associated with the iron oxides of mineralized samples at NSE. Anatase in hematitized mafic volcanic rock yielded 1717 ± 12 Ma, and monazite included in hydrothermal hematite in iron ore indicated 1613 ± 21 Ma. These are interpreted as representing the iron mineralization ages.

3. TECTONIC AND STRUCTURAL SETTING

Several interpretations have been presented for the tectonic evolution of the Itacaiúnas Supergroup volcano-sedimentary sequence in the Carajás region. Dardenne et al. (1988) suggest a rift-related subduction zone to account for the basin responsible for the Grão Pará Group’s evolution, which would have developed on continental crust adjacent to rift zones (Olszewsky et al., 1989). Other authors point out that the volcano-sedimentary sequence is related to intracratonic basins (Gibbs et al., 1986; Machado et al., 1991; Lindenmayer and Fyfe, 1992; Santos, 2003; Grainger et al., 2008), or that such
sequences formed in an arc environment of an evolving subduction zone (Meirelles & Dardenne, 1991; Teixeira, 1994; Lobato et al., 2005b; Rosière et al., 2006).

According to Zucchetti (2007) and Zucchetti et al. (2007), the Grão Pará Group basalts have a calc-alkaline magmatic affinity, and Nb-negative and Th-positive anomalies that are consistent with a subduction zone signature. Trace-element enrichment (e.g. high Zr/Y and Nb/Yb ratios) suggests that they formed in a continental-arc environment (Zucchetti, 2007). Basalts (Olszewsky et al., 1989), and jaspilites with negative εNd values (Lobato et al., 2005b) indicate crustal contamination of the Grão Pará Group. Thus, the trace-element geochemistry characteristics suggest a back-arc-related tectonic setting on an attenuated continental crust (Zucchetti, 2007; Zucchetti et al., 2007). These authors also suggested that the Grão Para Group is a greenstone-belt-type sequence as previously indicated by Hirata et al. (1982), Meireles et al. (1984), Araújo & Maia (1991) and Faraco et al. (1996).

The structural setting of the CMP is extensively discussed in the literature (Beiseigel et al., 1973; Silva et al., 1974; Araújo and Maia, 1991; Holdsworth & Pinheiro, 2000; Pinheiro & Holdsworth, 1997; 2000) despite a distinct lack of detailed field data. A summary of the structural framework was provided by Rosière et al. (2006) who proposed that the dominant structure is a flattened flexural fold system with axes moderately plunging ESE to WNW, intersected by several strike-slip faults that trend sub-parallel to the fold plane of the fold system. This interpretation is based on structural data taken from several iron ore deposits (N1 to N5), regional mapping and satellite image analyses. Further to the southeast (N6 to N8), other Serra Norte deposits are currently being investigated in order to better define the regional structural setting including the anticlines and synclines.

The Serra dos Carajás itself is a S-shaped syncline-anticline pair, named the Carajás fold, and is partially disrupted by the Carajás shear zone. The latter divides the entire structure into blocks that subsequently were named the Serra Norte and Serra Sul (Fig. 1). The approximately WNW-ESE-trending sinistral Carajás and Cinzento shear zones (Fig. 1) represent major structural discontinuities sub-parallel to the axial plane of several minor regional folds. The shear zones were likely developed to accommodate progressive flattening of the Carajás fold by lateral escape (Rosière et al., 2006). Another major event that influenced the structural setting of the area, particularly in the eastern part of the CMP, was the intrusion of shallow-level (Barros et al., 2001), syntectonic calc-alkaline granite intrusions (named intrusive suites in Fig. 1), such as the Estrela Complex (Barros & Barbey, 1998, 2000), into the metamorphosed volcano-sedimentary sequences of the Itacaiúnas Supergroup.

The Carajás and Cinzento shear zones (Fig. 1) played a decisive role in the iron mineralization processes by preparing the terrane (i.e. increased the porosity) during the Archean and creating pathways for hydrothermal fluids to percolate in the Proterozoic when these shear zones were reactivated, resulting in roughly north-south splays along the Serra Norte iron ore deposits.

The high-grade Serra Norte orebodies developed in zones of greatly enhanced rock permeability at the regional hinge zone of the anticlinal Carajás fold (Rosière et al., 2006), which has been subsequently disrupted and rotated by faults and related splays. Hard orebodies surrounded by soft ore are preferentially concentrated in the hinge zone of large folds such as in the NS deposit (Lobato et al., 2005a). The Serra Sul deposits are located within large (i.e., hundreds of meters to kilometer wavelength) second-order folds (Lobato et al., 2005b), but the detailed structural framework and relative timing relationships are presently not well constrained.

According to Rosière et al. (2006) and Lobato et al. (2005b), the formation of the Carajás iron orebodies postdated regional metamorphism and all Archean deformation events documented in the CMP. Regional metamorphic grade is very low in rocks of the Grão Pará Group (e.g. Beisiegel et al., 1973; Gibbs et al., 1986). Medium- to high-temperature minerals and tectonic fabrics in both ore and country rocks are found exclusively in: (i) the shear-zone domains, where high hydrothermal fluid-to-rock ratios control the mineral assemblages, or (ii) the contact metamorphic zones near granitic bodies. Most of the granites in the CMP and near the iron ore deposits are of Proterozoic age (e.g. Machado et al., 1991). High-grade iron ores also display a distinct hydrothermal texture, such as comb-textured hematite veins and breccias, which overprints the jaspilite fabric.

4. IRON ORE DEPOSITS

4.1 - Mesoscopic classification of iron ore types

The proterre at the Serra Norte iron ore deposits is defined as: (1) jaspilite, which may contain bedding-parallel quartz veins, and (2) early-stage hydrothermally altered, so-called, least-altered jaspilite. Iron ore is defined as advanced stage hydrothermally altered and mineralized jaspilite and consists of hard- and porous hard-ore. Hard ore also includes variable amounts of quartz +/- hematite vein types. Soft ores can be the direct product of
Jaspilite and hypogene ores via supergene enrichment (Tolbert, 1971), but these are not described here. The transition from the jaspilite to the hydrothermal hard ore encompasses the development of varying porosity stages that may locally characterize a high-grade soft ore type (Lobato et al., 2005a; 2008). This type of hypogene soft ore is typical of the N4E deposit.

High-grade ores can be both soft and hard, and are classified in terms of their physical hardness into hard to medium-hard, friable, soft and powdery and dusty ore (Clout & Simonson, 2005). Hard to medium-hard ores have a low porosity with an interlocking texture between martite and hematite, or martite-MpHem (microlamnate hematite) grains. Friable ores are easily disintegrated by hand, and are more porous than medium-hard ore. They commonly break into centimeter-size prisms/plates, defined by joint planes and fissile beds. Soft (0.05-1.0 mm particles) ores are those that can be dug in situ by hand or with a shovel; they are very porous, but typically not powdery. Soft orebodies are discontinuous and tabular, friable and banded, and locally contain massive hard ore lenses.

The friable ore displays primary lamination and contains 64 to 67 wt per cent Fe. It consists of friable hematitic material with gray color, metallic luster and a high porosity. High-grade hard ores may be massive, and/or compact, brecciated and banded where the layers can be compact and/or porous (Rosière & Chemale Jr., 2000). Locally, the brecciated ores exhibit fragments of dissilicified jaspilite layers, with the original fabric completely destroyed in places (Figueiredo e Silva, 2004). Compact hard ores, commonly gray blue, contain 67 wt percent Fe, have a metallic luster and low porosity. Massive hard ores are mainly observed along the contact between jaspilites and lower basaltic wall rocks as small lenses and tabular bodies (first described by Tolbert et al., 1971), and also as lenses surrounded by basaltic rock.

Basalts in the N4 and N5 deposits display hydrothermal alteration minerals, with both intense chloritization and hematitization along the contact with the orebody (Figueiredo e Silva, 2004; Zucchetti, 2007).

Iron ores and weathered basaltic rocks are covered by ferricrete (canga), which formed in the superficial portions of deposits as a supergene product. It consists of cemented goethitic/limonitic material, with colloform, vuggy textures, and has a high porosity. Where canga is located directly over iron ore, it consists of hematite blocks that are cemented by hydrated iron oxides (Lopes, 1997). The classification and description of the supergene ores are provided in Beisegel et al. (1973), Rezende & Barbosa (1972) and Ladeira & Cordeiro (1988).

Iron ore may be contaminated with: (i) manganese, particularly where in contact with lower basaltic wall rocks; (ii) aluminum and phosphorous where in contact with canga; and (iii) silica, which generally increases with depth (CVRD, 1996). The cut-off grade for iron ore is 60 wt % Fe, 2 wt % Mn, 2-2.5 wt % Si and Al, and 0.2 wt % P.

4.2 - Geological setting of the N1, N4 and N5 deposits

**N1 deposit:** At the N1 deposit (Fig. 2A), the dip of the rocks of the Carajás Formation varies from 45° to subvertical and outcrops are characterized by soft friable ore, soft lateritic hematite ore and canga cover. Hard ore is subordinate to soft ore and is found as either discordant (northeastern part of the N1 deposit) vein-controlled or discordant (southeastern part of the deposit) lenses to the soft ore. Hard ore may contain hematite veinlets.

**N4 deposit:** The N4 deposit comprises two main orebodies, namely N4E and N4W that are interpreted to be separated by a N-S oriented fault (Fig. 2B). The N4W orebody (Fig. 2B) is presently the most important in the Carajás iron district, with total reserves in excess of 0.5 billion tons, and consists mainly of soft ore and only minor hard ore. Bodies of hard ore are frequently close to NE-SW and NW-SE trending faults (Domingos, 2005). Jaspilites (Fig. 5A) have an average thickness of 220 meters (Borges, 1994) and crop out in the northern portion of the N4E deposit as well as smaller, < 40 meters thick lenses contained within the orebodies. Phosphorus, aluminum and manganese contents increase close to ferricrete cover, and along contacts with mafic volcanic wall rock and dikes (CVRD, 2004).

The N4E open pit displays an approximate J shape structure in plan-view (Fig. 2B), with a N-S extension of 5 km, width of about 500 m and an average orebody thickness of 350 m. The deposit is divided into a northern and southern domain. In the northern domain the lithological units strike mainly N-S and dip to the W. According to Ladeira & Cordeiro (1988), hard hematite orebodies and jaspilites are distributed discontinuously along a N30W trend in the northern domain of the N4E deposit, which is also the trend of the axial trace of early-stage folds (F1). The second-generation F2 folds plunge 4° toward N25-50W. Laminated soft ore predominates in the northern domain, containing many irregular and partially preserved jaspilite lenses. Lenses of hard ore are in the middle of soft ore (Fig. 4A) that is in contact with jaspilite. The jaspilites contain discordant and discordant quartz veins, as well as hematite veinlets with quartz nuclei. The soft ore contains hematite veinlets and is locally brecciated at the basal contact with basaltic rocks.
At the N4W deposit, soft ore predominates and hard ore is scarce. The soft ore consists mainly of friable and porous hematite-martite material. The open pit is approximately 7 km long, 200 to 500 m wide and 400 m deep (CVRD, 1996). The bedding of jaspilites and ores strike NW-SE with variable dips from subvertical to 40°-50° to the SW in the central part of the open pit. In the northern part of the open pit bedding strikes N20-30E, and dips 40°-50° to the NW (Pinheiro et al., 2001). Poles of these structures indicate major mine-scale folds with their axes plunging shallowly (< 30°) to the NW (Pinheiro et al., 2001). A gradational soft ore and jaspilite contact is often observed in many of these mine benches (Fig. 2).
Quartz breccias (hematite-free), as well as bedding-discordant quartz veins, are commonly observed. Quartz breccias are observed from the N4 deposit. The N5E orebody is massive hard ore overlying soft ore that are discordant to the bedding. Hydrothermal aureole, which may contain hydraulic breccias; vugs and lenses of dolomite are observed in the eastern portion of this domain at the lower contact of jaspilite with the ore sequence. The N5 deposit, where the high-grade hard hematite-type ore is abundant. Mineral abbreviations: Hem: hematite; Mt: martite; Mag magnetite. In the southern domain bedding planes strike NE-SW and dip to the NW, giving the J shape of this deposit (Pinheiro et al., 2001). Soft ore also predominates in the southern domain, enclosing lenses of dolomite-bearing and manganiferous jaspilites and, banded hard ore that contains dolomitic carbonate veins (Fig. 5B). Hydrothermally altered, hematitized basaltic rocks are found in the eastern portion of this domain at the lower contact of jaspilite with the ore sequence. Near the contact with the iron ore, the basalts are totally replaced by chloride and hematite (up to ~80 vol % of chloride and 15-20 vol % of hematite) and minor quartz, calcite, and white mica. Hard hematite orebodies (>56 wt % Fe) occur mainly near the contact with the overlying mafic metavolcanic rocks. They are typically surrounded by a hydrothermal aureole, which may contain hydraulic breccias; vugs filled with carbonate, quartz, kaolinite, MtHem, quartz-hematite veins (Fig. 5C), and fibrous chloride aggregates (Guedes 2000). Carbonate-sulfide veins are known to be present along the jaspilite-basaltic lower contact (Figueiredo e Silva et al., 2007b) in the southeastern portion of the N4E deposit.

**N5 deposit**: The N5 deposit (Figs. 2 and 3) displays geological characteristics, such as the dominance of massive hard ore overlying soft ore that are distinct from those at the N4 deposit. The N5E orebody is presently pod shaped, with jaspilites and high-grade ore, mainly hematite, surrounded by mafic rocks, which are locally hematitized where in contact with hard ore (Figs. 3 and 4B). The jaspilites and basaltic rocks have a predominantly N40W to N80W strike and dip 50°-60° to the SW (Pinheiro et al., 2001). Numerous, metallic gray hematite, stockwork veins may constitute hard ore and locally crosscut both the soft ores and jaspilites (Fig. 5E). “Christmas-tree” veinlets are also observed (Guedes et al., 2002), similar to those described by Taylor et al. (2001) and Thorne et al. (2004). Evidence of dolomitic ore has only been noted in drillcore where dolomite veinlets crosscut and/or are parallel to beds, and locally forms a carbonate-rich breccia fill (Figueiredo e Silva, 2004; Figueiredo e Silva et al., 2007a). The contacts between basaltic rock and hard ore, and jaspilite and porous hard ore, are gradational with clear lateral bedding continuity. The soft ore is enriched in silica near jaspilite lenses. Phosphorous contamination increases toward the ferricrete cover, whereas aluminum and silica increase towards the base of the orebody (CVRD, 2004). Discordant quartz veins are rarely associated with the hard ore. Zucchetti (2007) reported basaltic rocks that are completely replaced by chloride and hematite where in contact with the iron orebodies (Figs. 5F and G). Late-stage hematite veins crosscut the chlorite-hematite rocks (Fig. 5H). At the N5S deposit, jaspilites are crosscut by vuggy quartz-hematite, and hematite-only veins that are either discordant or parallel to the jaspilite beds (Fig. 5I). Soft ore predominates and silica contamination is common. The most common hard ore in the N5S deposit constitutes meter-scale banded, silica-contaminated ore. This ore differs from the typical hard ores (which are scarce in N5S) in other deposits because it contains remnants of strongly desilicified and hematitized jaspilite-like bands. Hard magnetite-rich lenses (approximately 2m thick) are unique in the Serra Norte iron ore deposits, and are found locally at the jaspilite-ore contacts. Magnetite is also found as veins that are discordant or parallel to jaspilite beds (Fig. 5J).

**5. Petrography of Jaspilite and Hard Iron Ore**

**5.1 - Jaspilite and least-altered jaspilite**

Jaspilite is characterized by alternating micro- to mesolayers (2 to 40 mm) of jasper and microcrystalline hematite (MiHem). The jasper layers (Fig. 6A) contain chert and dusty hematite, which is defined as <0.004 mm hematite particles. Locally the jasper layers contain chalcedony spherulites (0.01 to 0.1 mm in diameter). The MiHem layers (Fig. 6B) are composed of dense concentrations of very fine-grained, <0.004 mm,
hematite crystals that represent the original oxide species of the jaspilite bands.

Least-altered jaspilite contains portions of hematite-free, recrystallized chert in equilibrium with magnetite, interpreted as the early stage of hydrothermal alteration. Variably altered jaspilites may be brecciated, containing various amounts of hematite types (e.g. microplaty and anhedral), and vein-associated quartz, carbonate and sulfide minerals.

![Diagram A](image1.png)  
**Legend**
- Canga over ore
- Canga over mafic rock
- Soft ore
- Hard ore (hematite-martite)
- Carbonate-veined hard ore
- Jaspilite
- Hematitized mafic rock
- Mafic rock

![Diagram B](image2.png)  
**Legend**
- Canga over ore
- Canga over mafic rock
- Weathered mafic rock
- Soft ore
- Jaspilite
- Hard ore (hematite)
- Hematitized mafic rock
- Mafic rock

Fig. 4. A. Cross-section of the N4E orebody. B. Cross-section of the N5E orebody. Note the presence of a lens of hard ore along the contact with iron-mineralized mafic rock. Source Vale (local mine grid). For location of drill cores refer to figure 2B.

5.2 - Iron Ores

Iron ores are classified based on the oxide paragenetic sequence and detailed texture and mineralogy of oxides and gangue minerals such as quartz and carbonate (Table 1 and Fig. 7). Two main ore types are recognized: (i) hematite-martite (Figs. 7A, B and C), and (ii) hematite ores (Figs. 7D and E). Locally, the major ore types may develop certain variations, as shown in Table 1. Ores from the N4E deposit, for example, are characterized by a combination of different hematite types and martite, and may preserve layering (Fig. 7A) inherited from the jaspilitic protore; such ores may exhibit a continuous transition from a typical hard to a porous hard ore. The martite-hematite hard ore with quartz veins (Figs. 7C and H) has only minor amounts of carbonate. In contrast, carbonate and hematite-martite ores (Figs. 7B and G) contain...
Fig. 5. A. Jaspilite (Jp) in the N4E northern domain. Microbanding is defined by alternating jasper and iron oxides bands. B. High-grade massive ore with carbonate veins discordant (arrow) and along bedding in the N4E southern domain. C. Mafic rock partially hematitized with hematite veinlets, N4E deposit. D. Gradational contact between jaspilite and soft high-grade ore (SO) in N4W deposit. E. Hematite hard ore stock-work veins crosscutting soft ore and also jaspilite lens, N5E deposit. F. Hematitized basaltic rock in contact with hard orebody, NSE deposit. G. Core sample of hematitized basaltic rock where amygdales are replaced by hematite (arrows), N5E deposit. H. Core sample showing contact between hard ore and basalt replaced by chlorite and hematite, with hematite veinlets (arrow), N5E deposit. I. Core sample of banded jaspilite from the N5S deposit showing quartz-hematite vuggy veins (arrow) cutting across the bedding in the jaspilite. J. Core sample of jaspilite from N5S deposit showing magnetite veins (arrow) cross-cutting the jaspilite. Abbreviations: Jp = jaspilite; SO = soft ore; HO = hard ore; Dol = dolomite; Qtz = quartz; Hem = hematite; Mag = magnetite.
significant amounts (up to 70 vol %) of carbonate (Figs. 7B and G). The ores of the NSE deposit differ from those of other deposits because they are brecciated and comprised almost entirely of hematite, i. e., massive or porous hard ore (Figs. 7D and J), representing those hard ores where the original layers were almost obliterated (Figs. 7 I, J and K).

**Hematite-martite ore type:** This ore type is typical of the N4E and N1 deposits. Hematite-martite ore is mainly composed of microcrystalline (+/- microplaty) hematite and disperse euhedral martite crystals with or without MpHem. Most of these ores still preserve the original jaspilite texture. Euhedral martite crystals are about 100 to 300 μm in size (Fig. 6D) and overgrow the massive MiHem and/or MpHem. Locally, kenomagnetite relics are present. Microplaty (or lamellar) hematite is made up of platy crystals, commonly with diamond-shaped sections of approximately 4-8 μm, locally up to 100 to 200 μm in length. The MpHem generally occupies vein walls (Fig. 6E), grows at the expense of martite (Fig. 7F), and less commonly is found in the center of martite crystals. Anhedral hematite (AnHem) forms eye-shaped agglomerates of lobate crystals in the oxide layers and also in veins. The crystals are ~ 20 μm in size and where located in veins, they display growth lamellae (Fig. 6F). Euhedral and bladed-tabular (Ehem-THem) hematite occurs in veins, is fine- to medium-grained (200 to 300 μm) and forms internal selvages to veins (Fig. 6G). Microcrystalline hematite is recrystallized, where in contact with these Ehem-THem hematite veins.

**Hematite ore type:** These ores are typical of the N5E deposit and are essentially composed of hematite, mainly microcrystalline (<0.004 mm), besides AHem, MpHem, Ehem (+/- 0.3 mm) and THem (0.2-0.3 mm) (refer to Table 1 for explanation of abbreviations). These ores are generally brecciated, but locally preserve the original micro-to meso-bands that, if present, are defined by the intercalation of continuous and/or discontinuous layers of fine-grained (+/-0.03 mm) AHem aggregates with MiHem. The latter hematite can be recrystallized at the contact with MpHem-THem veins. Anhedral hematite mesobands are locally discontinuous and interspersed with vugs that may contain MpHem. The EHem-THem is commonly comb textured, may exhibit growth lines, and partially fill open spaces or vugs.

**Hematized wallrock:** It consists of hydrothermally altered and mineralized basalt, which are found only at the contact between the mineralized jaspilite and the under- and overlying basaltic wallrock. In the N4E deposit, they are composed of talc, kutnahorite, quartz and rare chlorite (locally in veinlets) together with MpHem (up to 0.4 mm) (Fig. 6H), with subordinate anhedral crystals of martite. The EHem-THem veinlets crosscut these rocks and exhibit growth lamellae. At the NSE deposit, the hydrothermally altered basalts are composed of chlorite (60%), with rare white mica, and MpHem (+/-35%) and martite (+/-5%). Rare kenomagnetite relics are present and locally also platy hematite with crystals up to 0.2 mm in length.

### 6. HYDROTHERMAL ALTERATION

Varying degrees of hydrothermal alteration have affected jaspilites, causing the transformation to iron ore, defined by a distinct paragenetic sequence of iron-oxide minerals with concomitant removal of silica and formation of a specific suite of veins that may contain quartz, carbonate, hematite, sulfide, monazite, and rare gold. The distal alteration zone represents an early alteration stage, whereas the intermediate and proximal alteration stages are synchronous with the main iron ore forming event. The proximal alteration stage also represents the advanced alteration stage, i.e., the high-grade ore itself. The distal alteration zone (up to ~ 80 m wide in the N4E deposit and ~ 100 m in the N4W deposit, measured vertically) in jaspilite is mainly characterized by the recrystallization of jasper and the removal of its iron, and the associated formation of magnetite (Fig. 8A). Magnetite crystals are euhedral to anhedral and form: (i) overgrowths on MiHem layers (Fig. 6C), (ii) grains in the nuclei of recrystallized chert invariably associated with crosscutting quartz veins, or (iii) grains in equilibrium with vein quartz and calcite. Magnetite is commonly replaced by hematite (martitization), leaving kenomagnetite relics (confirmed by Mössbauer analyses according to Figueiredo e Silva, 2004), which is a deficient Fe^{2+} magnetite phase (Kullerud et al., 1969).

Sulfides such as pyrite, chalcopyrite (Figs. 8C and D) may be associated with the hydrothermal alteration, but it should be noted that relatively rare, syngenetic pyrite (N4W) and calcite (N4E) crystals also occur in parallel laminae of jaspilite (Fig. 8B). The veins formed in this alteration zone lack hematite. The intermediate alteration zone is about 200 m in width (e.g. in the NSS deposit) and is characterized by martite, with or without kenomagnetite. Quartz-hematite and hematite-quartz veins are common (Fig. 8E) and contain MpHem and subordinate sulfides. The porosity increase in the altered jaspilites is significant, and affects the original jasper layers. The proximal alteration zone, represented by hard and hard porous ores, is about 20 to 50m in width. The original shape of martitic magnetite “blasts” is destroyed by the progressive martitization, forming AHe (Table 1) in abundant eye-shaped agglomerates of lobate crystals. Intense carbonate
alteration associated with the high-grade ores results in the formation of ore breccias cemented by dolomite. Examples are locally observed in the N5 deposit, and as kutnahorite cemented breccias in the N4 deposit. Hard ore with predominance of late-stage anhedral and tabular hematite lack quartz and carbonate veins. The presence of carbonate is mostly restricted to ores at depths >200 meters in the NSE deposit, and along the jaspilite-basaltic contact in the N4E deposit.

Microcrystalline hematite is stable even in the more advanced hydrothermal alteration stages. Stockwork-type veins contain tabular hematite. These zones are also characterized by discontinuous quartz ± carbonate veins with intergrown MpHem, which may be included in the hematite-martite ore type. The MpHem intergrown with quartz forms the matrix of hydrothermal breccias in the proximal alteration zone (Fig. 9F).

Hydrothermal alteration of basaltic wallrock has also been subdivided into intermediate and proximal alteration zones (Zucchetti, 2007). The hydrothermal zones are characterized by chlorite and chlorite-hematite replacement of the igneous minerals, respectively. The intermediate alteration zone has abundant chlorite, calcite, quartz, hematite, white mica, albite, titanite, magnetite, and subordinate sulfide minerals. Chlorite and MpHem from the proximal alteration zone occur along veins, or replace amygdales (Figs. 8G and H). Quartz, white mica, albite, titanite, and calcite are subordinate. The intermediate zone can locally extend up to about 30m, whereas the proximal zone is approximately 60m wide relative to non-altered basaltic rock (Fig. 5C).

Table 1. Hard iron ore types characteristics.

<table>
<thead>
<tr>
<th>Ore types</th>
<th>Hematite-martite</th>
<th>Hematite</th>
</tr>
</thead>
<tbody>
<tr>
<td>Deposits</td>
<td>N1 and N4E</td>
<td>N4E</td>
</tr>
<tr>
<td></td>
<td>Mt, MiHem, Mt</td>
<td>Mt (KenoMag), MiHem, Aivem + MtHem</td>
</tr>
<tr>
<td></td>
<td>MiHem + Mt</td>
<td>Mt (KenoMag), MiHem, Aivem + MtHem</td>
</tr>
<tr>
<td></td>
<td>MiHem, Sub-AHem, AHem-THem</td>
<td>Aivem-EHem-THem, MiHem</td>
</tr>
<tr>
<td>Mineralogy</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Veins-Veinlets</td>
<td>Qtz-Kut-MpHem</td>
<td>Qtz; MpHem, THem*</td>
</tr>
<tr>
<td>Others</td>
<td>(i)Qtz-Chalc, (ii)Qtz-Kut, (MpHem), (iii)MpHem + AHem, aside from AHem + THem*</td>
<td>Qtz; MpHem, THem*; AHem-EHem</td>
</tr>
<tr>
<td></td>
<td></td>
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</tbody>
</table>

*In MiHem-rich bands, MiHem is coarser-grained and recrystallized at the contact with these veinlets. Mineral abbreviations: MiHem = microcrystalline hematite; Mag = magnetite; KenoMag = kenomagnetite; MpHem = microplaty hematite; Mt = martite; AHem = anhedral hematite; EHem=euhedral hematite; THem = tabular hematite; Ccp = chalcopyrite; Qtz=quartz; Kut=kutnahorite; Cal=calcite; Dol=dolomite

Geonomos é publicada pelo CPMTC-Centro de Pesquisa Professor Manoel Teixeira da Costa, Instituto de Geociências, Universidade Federal de Minas Gerais
Fig. 7. Main features and characteristics of the different hard ore types. Core samples from A to E and photomicrograph from F to K. A. Hematite-martite ore type from the N4E deposit. B. Carbonate and hematite-martite (banded and brecciated) hard ore from the N4E deposit. C. Martite-hematite (brecciated and quartz veins) hard ore from the N4E deposit. D. Veined hematite (banded and/or brecciated) hard ore from the N5E deposit. E. Hematite and carbonate (brecciated) hard ore from the N5E deposit. F. Martite and MpHem crystals in hematite-martite type, N4E deposit. Note that MpHem grows along martite borders. Reflected light, under crossed polars (50X). G. Carbonate (kurmahorite) veins crosscutting MiHem and Mt bands of carbonate and hematite-martite ore, N4E deposit. Reflected light (50X). H. Mt crystals over MiHem bands in hematite-martite ore association, N4E deposit. Reflected light (50X). I. Anhedral-subhedral hematite portion characterizing brecciated hematite ore, N5E deposit. Reflected light, under crossed polars (50X). J. AHem and MpHem portions in brecciated hematite ore, N5E deposit. Reflected light (50X). K. Platy hematite (PHe) quartz vein in brecciated ore, N5E deposit. Reflected light (50X). Mineral abbreviations: MiHem = microcrystalline hematite; MpHem = microplaty hematite; Mt = martite; AHem = anhedral hematite.
6.1 - Veins and breccias

Detailed logging of about 20 diamond-drill cores, through the N4 and N5 deposits, and petrographic and textural descriptions of veins and breccias were used to establish a vein classification with respect to their location in specific hydrothermal alteration zones and associated iron mineralization and the iron oxide paragenesis (Fig. 9). It must be emphasized that the vein types from the distal alteration zone do not contain hematite, and represent an early mineralization stage. In contrast, hematite is intergrown with quartz, carbonate and sulfides (where present) in veins from the intermediate and proximal alteration zones. Two vein-breccia types characterize the distal alteration zone in jaspilite: V1a – quartz ± sulfide breccias, and V1b – carbonate ± sulfide breccia-veins. The intermediate alteration zone is represented by the following vein types: V2a – quartz ± hematite bedding-discordant veins; V2b - vug-textured quartz + hematite discordant vertical veins; V3 – hematite ± quartz veins crosscut and/or parallel to the jaspilite bedding. The proximal alteration zone is characterized by vein filling in breccia classified as V4 – carbonate-quartz breccia, and V5 – quartz ± microplaty hematite breccia, both are located in high-grade ore.

Quartz is present principally as (i) micro- to cryptocrystalline crystal aggregates of mosaic texture, probably the product of recrystallization of chalcedony and amorphous quartz (Lovering, 1972), (ii) comb-textured crystals observed in V2 and V3 veinlets and veins, and oriented perpendicular to magnetite crystal faces, and (iii) fine to very coarse (8mm), pyramidal crystals that occur along polygonal grain boundaries in V1, V2 and V5 veins.

Carbonate crystals occur in discordant V1 veins or veinlets and/or along jasplite layers, cementing ore breccias as V4 veins and locally in jasper layers. The anhedral to euhedral carbonate crystals are generally fine grained, but locally coarse grained (>10 mm). Zoned crystals display a dusty hematite pigment. They are classified as manganese-rich dolomite (i. e., kutnahorite), dolomite and calcite (Figueiredo e Silva, 2004). Calcite is only associated with V1 veins that are discordant and discordant to bedding in jaspilite.

Sulfide minerals: Pyrite is present as fine grained (0.1 to 0.2mm), subhedral and anhedral crystals or grains, and occurs in veins with associated Mphem or along the jasplite bands. In places, pyrite overgrows kenomagnetite or is intergrown (less commonly included) in martitized magnetite crystals (Fig. 8D). Chalcopyrite crystals are subhedral to euhedral, mainly in textural equilibrium with carbonate (Fig. 8C) or dispersed in anhedral quartz aggregates in V1 veins. Rare chalcopyrite crystals are included in kenomagnetite and locally in jaspilite bands. Rare covellite is associated with goethite, fine-grained hematite, chalcopyrite, and is found within quartz-rich jasplite crystals. Locally, veins of native copper and fine-grained gold are observed in jaspilite and martite aggregate, respectively.

Rare very fine monazite crystals were identified using scanning electron microscopy. They are associated with V1 veins.

7. WHOLE-ROCK GEOCHEMISTRY

Major, trace and rare-earth element (REE) analyses of 50 samples from the N1, N4E, N4W, N5E, and N5S deposits include protore jaspilite, hydrothermally altered jasplite and high-grade iron ore (Figueiredo e Silva, 2004; Lobato et al., 2007; Figueiredo e Silva et al., 2008). Analytical procedures, instruments and standards can be found in Figueiredo e Silva et al. (2008). The major-element diagram Fe₂O₃ versus SiO₂ for the N4W, N5E and N5S jaspilites exhibits a negative correlation characterized by the progressive decrease in Fe₂O₃ as SiO₂ increases (Fig. 10A). The Fe₂O₃Total contents vary between 48.7 and 63.2 wt percent (34.2 wt % to 44.2 wt % Fe), and SiO₂ contents between 35 and 50 wt percent. Two hydrothermally altered jasplite samples from the N4W deposit exhibit elevated Fe₂O₃ content of ~ 80 wt percent (c. 60 wt % Fe) similar to the ore samples. The hard ores have a high iron concentration, with N1 samples exhibiting between 68.2 and 69.7 wt percent Fe, and those from N5E between 64 and 67.5 wt percent Fe, with very low SiO₂ contents (Fig. 10A). The Al₂O₃ contents are < 0.3 wt percent in the jasplites, and < 2 wt percent in the ores; values from 2.3 to 4 wt percent were obtained for high-grade ore samples at the contact with basaltic wall rocks. Trace-element compositions in jasplites and iron ores are normalized to chondrite (Taylor & McLennan, 1985) (Fig. 10B). The elements Ba, U, Nb and Y are enriched in all types of ores when compared to jaspilites (Fig. 10B). The ores display an enrichment of U on the order of 20 to 200 times compared to jaspilite (Fig. 10B). Ores from the N5E deposit have U values up to 1000 times that of chondrite. The Ba concentration reaches ~ 100 times that of chondrite and 10 times that of jasplilte. The Nb varies up to 10 times chondrite, except in N5E jaspliltes where there is no Nb enrichment. The Y contents only exceed chondrite values in ores and hydrothermally altered basaltic rock. Jasplilte and ores from the NSE deposit display depletion of Cu, Zn, Ni (Fig. 10B) and Co, but V is enriched by ~ 20 times chondrite.
The REE data are normalized to chondrite (Nakamura, 1974). The sum of the REE in least-altered jaspilites varies in the N4E deposit from 5.6 to 23 ppm, in the NSE deposit from 4.5 to 13.8 ppm, and in the NSS deposit from 4 to 24.9 ppm. In contrast, for the ores this sum is more variable ranging from 6.5 to 14.8 ppm in the N1 deposit, except in two samples, in which ΣREE = 103 and 91 ppm, 8.1 to 44.2 ppm for the N4E deposit, and 14.6 to 97 ppm for the N5E deposit. The REE patterns of least-altered jaspilite samples from the N4W, NSE, and NSS deposits are shown in Figs. 11A, B and C. Samples from the different deposits exhibit similar fractionation patterns with light REE (LREE) enrichment (La/Sm) = 9.53 to 13.69, small to moderate positive Eu anomalies (Eu/Eu* = 1.54 to 2.34), and relatively horizontal patterns for the heavy REE (HREE) that commonly exhibit low contents (ΣHREE < 1 ppm). Samples from NSS display a wider range of REE values, mainly HREE (Fig. 11C).

Ores from the N4E deposit display two different populations (Figs. 11E and F), with higher (14.8 to 44.2 ppm) and lower (8.3 and 8.1 ppm) ΣREE. The former displays a LREE enrichment and discrete negative Eu anomaly, whereas the latter, a kutnahorite-bearing ore (Fig. 11F), presents a weakly positive Eu anomaly (1.3 and 1.5).

The characteristic pattern in ores from the N1 deposit, with ΣREE = 11.7 to 14.8 ppm (Fig. 11D), resembles that of the N4W jaspilites (Fig. 11A), except for higher Eu anomalies (2.1 to 2.9) and relative increase in HREE contents. In high-grade, hematite-martite N1 ores, oxides that formed due to the hydrothermal alteration, are microplaty hematite-martite ± tabular hematite; however original MiHem is still preserved. The slightly higher total ΣREE contents of the N1 ores, when compared to N4W jaspilites, may be related to their lack of jasper bands, and also to the larger proportion of newly formed MpHem.

Two distinct chondrite-normalized REE patterns characterize ore samples from N5E deposit. The first (Fig. 11G) shows an almost flat pattern and is characterized by veined hematite ore, locally layered with MiHem, AHem, and THem and MpHem veinlets. The second is represented by two brecciated hematite ore samples and yields the highest ΣREE contents, respectively 85.54 and 97.52 ppm (Fig. 11H). They are predominantly comprised of the advanced-stage alteration oxides, with
subhedral to aHem, and also contain tabular, lamellar and euhedral hematite veinlets. This REE pattern shows the most prominent LREE enrichment in comparison to the HREE, a slightly negative Eu anomaly (Eu/Eu* = 0.62 and 0.79), with the possible addition of iron. Michard (1989) related the increase in the concentration of REE, negative Eu anomaly and LREE enrichment to equilibration with low-pH fluids. The ubiquitous presence of newly formed MPres in these ores suggests a greater incorporation of REE, favored by the same coordination in the structure and size of the ionic radii of REE (Khan et al., 1996), from 1.032 Å (La) to 0.861 Å (Lu) (Grossi Sad & Dutra, 1987).

In comparison to least-altered jaspilites (Figs. 11A, B and C), the N1, N4E and N5E ores yield REE compositional populations progressively enriched in REE, in that order, except for a single N1 sample. In general, there is closer similarity to the HREE than to the LREE chondrite concentrations (Fig. 11D). The ∑LREEi commonly yields higher concentrations, up to 88.2 ppm in N5E ores, and elements with greater standard deviations are represented by Ce, La and Nd, in jaspilites and ores.

Very fine-grained (< 0.01 mm) monazite inclusions in some aHem crystals from N5E deposit were identified under SEM. Their influence on the especially elevated LREE enrichment cannot be discarded.

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**Fig. 10.** A. Bivariate plot of Fe2O3 and SiO2 compositions (in wt %) of least-altered jaspilite (here understood as variably altered jaspilite) and high-grade ores from the N4W, N5E and N5S deposits. Note negative correlation between Fe2O3 and SiO2 compositional values. Note that two altered jaspilites from the N4W deposit have higher iron contents (~ 80% Fe2O3) than others. B. Trace element distribution in jaspilites and iron ores. Data normalized to chondrite (Taylor and McLennan, 1985).
Fig. 11. Chondrite-normalized rare-earth elements plots for the jaspilite and iron ore of the Serra dos Carajás deposit. A. Least to variably altered jaspilites, N4W deposit. B. Least- to variably altered jaspilites, NS5 deposit. C. Least to variably altered jaspilites, NS5 deposit. D. Hematite-martite ores, N1 deposit. E. Hematite-martite ores, N4E deposit. F. Carbonate (kutnahorite) and hematite-martite ores, N4E deposit. G. Veined hematite ores (locally banded), N5E deposit. H. Brecciated hematite ores with sub- to anhedral hematite, and euhedral-tabular and lamellar hematite veins, N5E deposit. Line with boxes corresponds to average of Isua quartz-magnetite banded iron formation, West Greenland (Dymek and Klein 1988). Data normalized to chondrite (Nakamura 1974).
8. DISCUSSION

8.1 - Review of the genetic models proposed for the Carajás iron ore formation

A number of authors have proposed a hypogene genesis for the iron ore deposits at Carajás, which preceded supergene residual enrichment via leaching of silica (Tolbert et al., 1971; Rezende & Barbosa, 1972; Beisegel et al., 1973; Ladeira & Cordeiro, 1988; Macambira & Lopes, 1996a; Dardenne & Schobbenhaus, 2001; Lindemayer et al., 2001; Beukes et al., 2002; Klein & Ladeira, 2002; Guedes et al., 2002; Dalstra & Guedes, 2004; Clout & Simonson, 2005). However, no systematic work, either in the public or private domains, existed regarding the hypogene ore, with the exception of a few generalized descriptions (e.g., Dalstra & Guedes, 2004). In addition, there was only restricted information available about the jaspilites at Carajás (e.g. Lindemayer et al., 2001; Macambira, 2003).

At Carajás a volcanic origin for the so-called “primary iron ore”, which contains magnetite, titano-magnetite and ilmenite, was first suggested by Suszczynski (1972), who also mentioned a close association between the hard ore and volcanic rocks. He further suggested that magnetite developed during a magmatic stage, whereas hematite was considered to be post-magmatic. Rezende & Barbosa (1972) and Beisegel et al. (1973) also proposed a hypogene, metasomatic origin for the massive ore lenses within friable orebodies. Lenses of massive orebodies are associated with intrusive basic dikes, and are located along the hangingwall contact with these dikes, which also crosscut the soft orebodies. Based on these ideas, Melo et al. (1981) suggested that the dikes generated heat for the metasomatic process. The authors also indicated that in the Serra Leste deposits the distribution of the principal lenses of massive hematite ore is structurally controlled.

According to Tolbert et al. (1971), ore formation resulted from leaching of silica from the “iron formations” via weathering due to water percolation that generated residual enrichment of iron oxides and formation of the present orebodies.

Teixeira et al. (1997) first mentioned the occurrence of scarce dolomite lenses at the base of the “BIF”. According to the authors, these lenses are relics of former limestone beds that underwent calcite leaching during hydrothermal alteration. These authors indicated that removal of these limestone might have enhanced the “BIF” permeability via fracturing and crack propagation. This would have been followed by silica leaching resulting in iron enrichment of the “BIF”. Macambira et al. (1999) pointed out that these carbonate lenses (~ 50 meters wide) represent a “carbonate-bearing jaspilite facies” and suggested that carbonate had been leached from those rocks in order to form hard ores. Due to the local presence of finely laminated dolomite in the N4E deposit, Guedes et al. (2002) and Dalstra & Guedes (2004) defined carbonate-rich protores (45 wt % Fe), which they referred to as “dolomitic iron formation”. According to these authors, chert was displaced by dolomite along the layering, accompanied by magnetite development. Dalstra & Guedes (2004) also pointed out that a second dolomitization stage was responsible for carbonate precipitation in veins and vugs, with the local development of breccias. The intense tropical weathering leached the carbonates from “dolomitic iron formation” resulting in the development of friable, soft ore.

Nevertheless, detailed documentation of the hydrothermal alteration stages (Figueiredo e Silva, 2004; Figueiredo e Silva et al., 2008; Lobato et al., 2005a; Zucchetti, 2007) that are spatially and temporally associated with the formation of hard ore have shown that the majority of the carbonate (i.e. dolomite, ktnahorite and calcite) is in the form of discordant and discordant veins and breccias commonly in association with sulfides (e.g. chalcopyrite and pyrite), with or without subordinate magnetite and hematite. Carbonate alteration is only ubiquitous in the N4E and N5E deposits, and is lacking or less common in most other deposits. Moreover, replacement textures of chert or quartz by carbonate are only locally observed. As such, carbonate cannot be considered to be paramount to iron ore formation in the Serra Norte deposits.

8.2 - The REE fingerprint from jaspilite to iron ore

The crystals of magnetite in hydrothermally altered jaspilites are locally surrounded by recrystallized and “clean” jasper, and in equilibrium with fine (~ 0.01mm) vein quartz. In accordance with Taylor et al. (2001), the leaching of quartz in BIFs is principally seen as a function of temperature, between 150 to 250°C. This may have been the case in the deposits studied here (Rios et al., 2004; Figueiredo e Silva et al., 2008). The leaching of quartz was accompanied by an increase in martitization of magnetite (Fig. 6D). Quartz and carbonate, therefore, precipitated prior to or synchronous to the formation of magnetite in the jaspilites (Fig. 8A). The leaching of quartz may have been responsible for the general increase in total REE contents, as only magnetite and MiHem remain (Fig. 6C). With the growth of magnetite and MiHem, and the progressive martitization to form Ahem, the total REE concentration increased significantly (Figs. 11G and H). This is corroborated by the distribution coefficients $K_o$ of magnetite, which favor the fixation
of LREE (Schock, 1979; Li, 2000). This process may also have favored the relative increase in HREE in the residual hydrothermal fluid, explaining the increase in HREE and the flat REE patterns (Fig. 11) for some ore samples in comparison to least-altered jaspilites. The change in the REE patterns of the ores in relation to jaspilites, encompassing a general REE enrichment with almost flat HREE patterns for some ore samples (Fig. 11G), indicates that the fluids responsible for the early-hydrothermal stage were significantly different or evolved to those of the advanced hydrothermal stage. Indeed, it is during the advanced hydrothermal stages that the EHem and THem generations dominate (Figs. 6F and G).

Hydrothermally altered jaspilites from the N4E, N5E and N5S deposits (Figs. 11A, B and C) show that Ni and a specific group of samples from the N4E deposit (Figs. 11D to H) display positive Eu anomalies. Locally, N4E and N5E ores have a weakly negative Eu anomaly (Figs. 11E, G and H). According to Grossi & Dutra (1987), no mineral retains Eu$^{2+}$ under accentuated oxidizing conditions, since the element would be unavailable at that oxidation state. However, under reducing conditions, Eu is isolated, potentially being incorporated in the Eu$^{2+}$ state. The initial hydrothermal fluid that interacted with the Serra Norte rocks was relatively reducing (i.e. $fO_2$ in equilibrium with magnetite), probably causing Eu reduction, which may be corroborated by the formation of magnetite after original hematite. The Eu$^{2+}$ was possibly fixed in the structure of minerals, such as dolomites. For example, in the case of the N4E ores, samples with carbonate do in fact display a positive anomaly (Fig. 11F). With the evolution of the hydrothermal fluids, they become relatively more oxidizing, with the remaining Eu$^{2+}$ not incorporated in the rocks, resulting in REE patterns with a weakly negative Eu anomaly, for example in the high-grade ores (Figs. 11G and H).

Ore samples from selected Hamersley deposits show similar features, with less pronounced enrichment in LREE (Lobato et al., 2007). The overall higher REE contents of the Carajás samples compared to the Hamersley ore samples suggest that different fluid sources may have been involved in the origin of the deposits. For Carajás, a magmatic fluid source mixed with meteoric waters has been postulated (Lobato et al., 2005b; Figueiredo e Silva et al., 2007a, 2008), whereas, for example, Hagemann et al. (1999) and Thorne et al. (2004) suggest that basinal brines were involved in the origin of the Mt. Tom Price deposit.

9. CONCLUSIONS

The giant Serra Norte Carajás iron ore deposits are hosted by the metavolcano-sedimentary sequence of the Grão Pará Group, Itacaiúnas Supergroup, where the jaspilite protore and high-grade iron ores (> 64 wt % Fe) are under- and overlain by low metamorphic grade basaltic rocks. The iron ore forming age is considered to be Paleoproterozoic age (Lobato et al., 2005b, 2008; Santos et al., 2010). Detailed mineralogical, geochemical and fluid chemistry investigations on the Serra Norte Carajás iron ore deposits revealed the following:

1. Considering the hydrothermal origin of iron ore forming event at the studied Serra Norte iron ore deposits (Guedes et al., 2002; Figueiredo e Silva, 2009; Lobato et al., 2005b), the sequential order of oxides that follows the original microcrystalline hematite-MiHem is interpreted as: magnetite $\rightarrow$ martite with or without associated kenomagnetite $\rightarrow$ microplaty hematite (MpHem) $\rightarrow$ anhedral hematite (AHem) $\rightarrow$ euhedral (EHem) and/or tabular hematite (Them). Other mineralogical modifications are: (i) recrystallization and the cleansing of jasper with the formation of chert and fine quartz; (ii) progressive leaching of chert and quartz, leaving oxides and a significant volume of empty spaces as vugs; (iii) silicification and dolomitization with associated sulfides and oxides in veins, breccias and along jaspilite bands; (iv) advanced martitization with the formation of AHem, partial MiHem recrystallization to AHem and partially filling spaces with microplaty/platy hematite; (v) continued space-filling by comb-textured EHem and THem in veinlets and along bands.

2. The iron mineralization involved hydrothermal fluids probably under epizonal conditions. Comb textures defined by quartz, hematite and talc, as well as preserved primary amygdales in basaltic rocks (Zucchetti, 2007) and spherulites in jaspilites, are typical of deposits formed under epizonal conditions and preserved at shallow, low-temperature crustal conditions.

3. Two different populations of REE patterns are notable: (i) those characteristic of the jaspilites, ore samples from the N1 deposit and a group from the N4E deposit, which indicate a distinct LREE enrichment and positive Eu anomalies, similar to patterns displayed by Archean BIFs worldwide; and (ii) those of N5E ores that are almost flat, with relative enrichment in both LREE and HREE. Another group of N5E brecciated hematite ores display high total REE contents, principally LREE, which may reflect the fixation of these elements in hematite.

4. The REE enrichment in ores and the changes in the shapes of the REE patterns indicate a hydrothermal fluid evolution, where silica leaching may have resulted in a relative general REE increase during the early hydrothermal stage. The LREE increase was more accentuated during the
formation of magnetite and MpHem, and the advance of martitization to AHem. This may have favored the relative increase of HREE in the residual fluid, resulting in an increase in HREE in advanced mineralization stage precipitates and almost flat REE patterns. It is during the advanced hydrothermal stages that the generation of euhedral and tabular hematite was dominant. The increase in REE concentrations in N4E and NSE ore samples further suggests the presence of significant amounts of Fe in the mineralizing fluid.

5. The mineralogical, geochemical and isotopic changes of jaspilites and ores attest to a hydrothermal origin for the formation of the hypogene high-grade iron ores, via interaction with: (i) an early-stage, high-salinity, relatively reducing magmatic fluid (Figueiredo e Silva et al., 2008), which leached silica, formed magnetite and precipitated quartz, carbonate and sulfides; (ii) an intermediate stage fluid that evolved to more oxidizing conditions, with the advance of martitization, increase in the REE concentration and precipitation of hematite in quartz veins; (iii) a late hydrothermal fluid of both low and high salinities, associated with the development of euhedral- and/or tabular-hematite bearing veins. The main hydrothermal fluid flow was probably focused along the original jaspilite-basaltic contact (Lobato et al., 2005b) with intensive lateral fluid diffusion through the jaspilitic sequence and less intense diffusion through the basaltic rocks as these had a more isotropic fabric when compared to banded jaspilites. An inherited Archean structural framework prepared the terrane for the iron mineralizing hydrothermal fluids, as the hard orebodies formed mainly in the hinge zone (e.g., the NSE deposit) of large folds, and associated with splays from existing shear zones (Rosière et al., 2006).

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