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The Manganese Deposits of Buritirama (Pará, Brazil)

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Abstract

In the Serra do Buritirama numerous lensoid bodies of supergene manganese ore, composed of eryptomelane, lithiophorite, nsutite and busetite occur. The unaltered protore consists of manganese calc-silicate marbles with small pods of braunite. The bonates, pyroxenoids, garnets, tephroite, braunite, manganophyllite and sulfides of the protores and associated rocks were studied in detail optically, by X-ray and microprobe to evaluate the physical chemical conditions of the metamorphism in amphibolite facies that affected the region. Low variance mineral assemblages suggest temperatures of metamorphism around $500-550^{\circ}$ C, and oxygen fugacities higher than 10^{-10} bar. Fluid inclusions infer corresponding pressures of about 3 kbar, at high X_{CO_2} .

It is concluded that the ore was deposited as ^a manganiferous limestone dolomite sequence in a shallow platform basin from terrestrial weathering products.

Introduction and geological aspects

The manganese deposits of Serra de Buritirama as well as the huge iron deposits of the Serra dos Carajâs are situated in the central-east part of Para State in the region drained by the Itacaiunas river, between the 5° 30' to 6° 30' ^S latitude and 50' W Greenwich (Fig. 1).

The vast Pre-cambrian terrain is part of the Guaporé Craton. The oldest polymetamorphic gneissic and amphibolitic rocks are locally named Xingu Complex, and the overlying metasedimentary sequence belongs to the Gräo Pará Group.

In the Serra dos Carajâs region, ¹⁰⁰ km south of Buritirama, the rocks of the Gräo Para are predominantly schists, quartzites and two basic metavolcanic sequences containing the Iron Formation (Itabirites; TOLBERT et al., 1971).

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Fig. 1. Localization map and regional geology of the Serra dos Carajâs region, including the Serra de Buritirama area (modified after GALEÃO DA SILVA and others, 1974; as in GOMES and others, 1975).

Overlying the Grâo Para Group discordantly, there is ^a sequence of molassie sediments represented by conglomeratic sandstones, siltites, shales and marbles - the Uatuma Group - correlated to the Rio Fresco Formation. This sequence contains manganese deposits similar to those from Francevillien Series of Gabon (WEBER and VALARELLI, in preparation).

Many little lensoid manganese bodies averaging ¹⁸⁰ m in length are found in the area (Fig. 2). By way of exception B_4 presents three mineralized areas totaling 3.3 km in length.

The best known B_5 orebody is 1,400 m long, 35-240 m wide and 5-30 m thick. A cross section (Fig. 3) shows weathered protores (manganese calcsilicate marbles and schists) interbedded with mica schists placed beneath sugary quartzite and above sericite quartzites all in ^a same general attitude of N-60-W, 20-30° NE.

The massive, hard, concretionary, high-Mn ore $(46-47\%)$ is restricted to the first 1-3 meters, grading rapidly downwards to a finegrained mixture of

Fig. 2. Geology of the Buritirama (B-5) orebody after ANDERSON, DYER and TORRES (1974) (simplified). Legend is the same as in Fig. 3.

Fig. 3. Geological profile, 200 N of the Buritirama, B-5 orebody after ANDERSON, DYER and TORRES (1974).

manganese oxide and clay minerals that constitutes the bulk of the crude ore $(32-46\%$ Mn) until the lowest weathering level is attained at about 60-80 m.

Enveloping the lensoid concentrations of manganese ores and sometimes interlayered with them are zones of weathered schists impregnated with manganese oxides and hydroxides (Fig. 3).

The authigenic manganese minerals of the supergene ore include: cryptomelane, lithiophorite, nsutite, and buserite-like minerals or amorphous $MnO₂$. Braunite is also present as relicts in the weathered protores. The main gangue minerals are: mica, chlorite, dickite, goethite and (weathered) spessartite.

Wall rocks

The manganese protores are enclosed in a thick banded series of calc mica schists and calc-silicate marbles interbedded with biotite schists (Fig. 4) (ANDERSON et al., 1974). The most common mineral assemblages are: diopsidic pyroxene + phlogopite + calcite + dolomite, diopside + phlogopite + garnet, tremolite + phlogopite + calcite + dolomite, diopside + phlogopite + K-spar + calcite + dolomite, diopside + quartz, diopside + tremolite + calcite + dolomite + quartz, biotite + garnet + quartz + K-spar + andalusite. Sphene, apatite and tourmaline are common accessories, along veins tremolite forms after diopside and is associated with epidote.

Fig. 4. Schematic lithologie profile of the weathered zone of the B-5-9 diamond drill hole and of the fresh rocks of the B-5-10 mond drill hole. These drills are located on the Fig. ² and represented in less detail in Fig. 3. Constructed on the basis description of geologists of Cia Meridional de Mineraçao and controlled by pétrographie and mineralogical studies.

Manganese protores

The protores can he followed through a thickness of 25-30 m and are made up of manganesiferous calc-silicate marbles and manganese-rich calc-silicate schists interlayered with less manganesiferous marbles and schists. Vertical

and horizontal changes in the original sedimentary conditions make it difficult to correlate different drill data.

a) Calc-silicate marble

Poikiloblastic manganese clinopyroxene, Mn-calcite and manganese phlogopite are the main constituents of a mosaic fabric, K-feldspar becoming a major constituent in ^a few samples. Manganese amphibole may replace pyroxene in patches and along veins. Chlorite veinlets can be found cutting the rock.

b) Calc-silicate schist

Manganese phlogopite (manganophyllite), manganese clinopyroxene (johannsenite), spessartite and manganese carbonate are the main constituents. Additional K-feldspar, pyrrhotite and barite may also occur as well as chlorite formed to the expense of manganophyllite.

c) Pyroxmangite calc-silicate marble

These protores are composed of a mosaic intergrowth of manganese-rich carbonate (rhodochrosite), equant spessartite, and large xenoblastic mangite showing lamellar twinning. Secondary manganese serpentine occurs in narrow veins.

d) Alabandite-tephroite-rhodochrosite marble

Large (2—3 mm) poikiloblastic tephroite crystals are seen beside Ca-rich rhodochrosite, sphalerite, alabandite and spessartite. Accessory pyroxmangite and hausmannite are also present.

e) Braunite-manganoan calcite marble

Masses of braunite are intimately intergrown with manganoan calcite, manganophyllite, K-feldspar and manganoan clinopyroxene. Massive rhodonite and quartz occur in veins. Some secondary formation of a manganese-rich amphibole (dannemorite) after rhodonite is common. During this alteration, fine grained carbonate may be formed.

Mineralogy of the manganese protores

These minerals were normally identified by the usual X-ray and optical means. The chemical compositions of the mineral phases were determined with an ARL electron microprobe type SEMQ, equiped with ⁶ spectrometers (4 scanners and ² fixed channels for Si and Al respectively). The following technical parameters were used: ¹⁵ KV

acceleration voltage, a sample current of 50 nA for silicates and 10 to 20 nA for carbonates. A fixed beam size of 0.5 micron was applied for silicates, whereas for carbonates the beam was varied between 1 and 20 microns depending on their stability. On each grain up to ¹⁰ points were analysed and averaged. Care was taken to analyse assemblages of ² or more minerals in contact with each other. For the silicate analysis natural silicates and oxides and for the carbonate analysis natural carbonates were used as standard material. For the minor elements at least 10 000 counts were accumulated providing a standard deviation (1 σ) better than 1.0%. The data were on-line corrected for background, deadtime and instrumental drift with ^a PDP-11/05 puter. Correction procedures for X-ray absorption, X-ray fluorescence and atomic number effects were based on ^a modified MAGIC IV program fitting the CDC ⁶⁵⁰⁰ computer system at the ETH-Zürich.

Carbonates

Carbonates consist almost entirely of $MnCO₃$, $CaCO₃$ and $MgCO₃$. Small amounts (less than 1 mol $\%$) of FeCO₃ were also found. No other component could be detected. As seen from the analyses in Table ¹ and Fig. 5, the Ca/Mn ratio varies over ^a wide range, although within one sample the composition

| | | Weight % | | | | Mol $\%$ end members | | | | | |
|---------------------|-------------------|-------------------|-------------------|-------------------|-------------------|----------------------|-------------------|--------------------------|--|--|--|
| | MnCO ₃ | CaCO ₃ | MgCO ₃ | FeCO ₃ | MnCO ₃ | CaCO ₃ | MgCO ₃ | FeCO ₃ | | | |
| $B-4-14-80.8$ I | 64.2 | 26.3 | 7.8 | 0.7 | 60.7 | 28.6 | 10.1 | 0.6 | | | |
| $B-4-14-80.8$ II | 64.4 | 25.7 | 7.5 | 0.7 | 61.5 | 28.1 | 9.8 | 0.6 | | | |
| $B-4-14-80.8$ III | 71.9 | 16.9 | 8.3 | 0.7 | 69.6 | 18.7 | 11.0 | 0.6 | | | |
| $B-5-10-80.2$ I | 19.1 | 77.2 | 2.6 | 0.2 | 17.1 | 77.5 | 3.2 | 0.2 | | | |
| $B-5-10-80.2$ II | 19.8 | 73.3 | 3.1 | 0.2 | 18.3 | 79.7 | 3.9 | 0.2 | | | |
| $B-5-10-80.2$ III | 16.3 | 82.8 | 1.3 | n.d. | 14.4 | 84.1 | 1.5 | | | | |
| $B-5-10-92.0 I$ | 20.8 | 78.3 | 1.9 | n.d. | 18.3 | 79.2 | 2.3 | $\overline{}$ | | | |
| $B-5-10-92.0$ II | 20.1 | 82.1 | 1.1 | n.d. | 17.3 | 81.4 | 1.3 | $\overline{}$ | | | |
| $B-5-10-97.8$ II | 11.7 | 90.3 | 0.4 | n.d. | 10.1 | 89.4 | 0.5 | $\overline{}$ | | | |
| $B-5-14-67.1$ I | 50.4 | 38.9 | 8.3 | 0.7 | 47.1 | 41.7 | 10.6 | 0.6 | | | |
| $B-5-14-67.1$ II | 51.5 | 38.9 | 8.0 | 0.7 | 47.8 | 41.5 | 10.1 | 0.6 | | | |
| $B-5-14-69.9I$ | 45.2 | 47.5 | 5.8 | 0.7 | 41.8 | 50.3 | 7.3 | 0.6 | | | |
| $B-5-14-69.9$ II | 44.1 | 47.5 | 5.7 | 0.7 | 41.2 | 51.0 | 7.2 | 0.6 | | | |
| $B-5-14-72.0$ I | 52.4 | 37.4 | 9.2 | 0.7 | 48.3 | 39.6 | 11.6 | 0.6 | | | |
| $B-5-14-72.0$ II | 51.1 | 36.3 | 9.0 | 0.6 | 48.4 | 39.5 | 11.6 | 0.6 | | | |
| $B-5-14-72.0$ III | 52.0 | 35.7 | 9.0 | 0.6 | 49.1 | 38.7 | 11.6 | 0.6 | | | |
| $B-5-14-72.0$ IV | 52.1 | 36.5 | 9.1 | 0.6 | 48.7 | 39.2 | 11.6 | 0.6 | | | |
| $B-5-14-78.8$ I | 50.2 | 38.8 | 8.7 | 0.7 | 46.8 | 41.5 | 11.0 | 0.6 | | | |
| $B-5-14-78.8$ II | 49.1 | 40.5 | 8.6 | 0.7 | 45.5 | 43.0 | 10.8 | 0.6 | | | |
| $B-5-14-78.8$ III | 49.0 | 40.7 | 8.7 | 0.7 | 45.2 | 43.2 | 11.0 | 0.6 | | | |
| $B-5-14-78.8$ IV | 51.0 | 37.8 | 8.7 | 0.7 | 47.7 | 40.6 | 11.1 | 0.6 | | | |
| $B-5-22-64.6$ I | 39.8 | 53.3 | 6.6 | 0.6 | 36.0 | 55.3 | 8.1 | 0.5 | | | |
| $B-5-22-64.6$ II | 38.8 | 53.4 | 6.5 | 0.6 | 35.4 | 56.0 | 8.1 | 0.5 | | | |
| $B-5-22 - 64.6$ III | 38.0 | 53.8 | 6.8 | 0.6 | 35.3 | 56.0 | 8.5 | 0.5 | | | |
| $B-5-22-64.6$ IV | 39.0 | 53.0 | 6.5 | 0.6 | 35.7 | 55.6 | 8.1 | 0.6 | | | |
| $B-5-22-65.8$ I | 40.9 | 50.5 | 7.1 | 0.7 | 37.4 | 53.1 | 8.9 | 0.6 | | | |
| $B-5-22-65.8$ II | 41.2 | 51.2 | 6.7 | 0.7 | 37.5 | 53.5 | 8.3 | 0.6 | | | |
| $B-5-22-65.8$ III | 41.0 | 49.0 | 7.6 | 0.7 | 37.8 | 51.9 | 9.6 | 0.7 | | | |
| $B-5-22-65.8$ IV | 38.9 | 53.1 | 6.5 | 0.7 | 35.5 | 55.7 | 8.1 | 0.7 | | | |

Table 1. Microprobe analysis of carbonates from Mn protores of Buritirama

(First two numbers refer to borehole, third number to depth in meters and roman number to spot on thin section. $n.d. = not detected.$)

Fig. 5. (Mn) -(Ca)-(Mg+Fe) mol per cent plot of carbonates, rhodonites pyroxmangites, clinopyroxenes and tephroites of manganese protores samples from Buritirama, analyzed by microprobe. The different coexisting phases are plotted with the same graphical symbols.

varies little. The amount of Mg is positively correlated with the Mn/Ca-ratio. Comparing the results with the experimental data of GOLDSMITH and GRAF (1957) in the system $MnCO₃-CaCO₃-MgCO₃$, most analyses fall within the region of complete miscibility that even at temperatures as low as 500° ^C extends from CaCO₃ toward Ca_{0.42}Mn_{0.58}CO₃. Carbonates of one sample have the compositions: $Ca_{0.19}Mn_{0.70}Mg_{0.11}CO_3$ and $Ca_{0.29}Mn_{0.61}Mg_{0.10}CO_3$. At 500° C these would * fall within the ² phase immiscibility field, but at temperatures slightly above 550° ^C they would lie outside.

Pyroxenoids

Three types of pyroxenoids are common: pyroxmangite, rhodonite and a manganese rich clinopyroxene.

The content of $CaSiO₃$ in the pyroxmangites show little variation (3-4.5) mol $\%$), but the MgSiO₃, FeSiO₃ and MnSiO₃ contents vary strongly as can be seen from the analyses in Table ² (Fig. 5). With the present data, no correlations between the composition and the optical properties are evident.

Pyroxmangite is the more frequent pyroxenoid and coexists with the manganese richer carbonates (Fig. 5).

Rhodonite occurs in veins and also as a rock-forming mineral coexisting

with the calcium richer carbonates. The $CaSiO₃$ content in the rhodonites varies from 9-11 mol $\%$, and the MgSiO₃ + FeSiO₃ content does not exceed 15 mol $\%$ (Table 2, Fig. 5). Fig. 5 also shows that rhodonite coexists with a Mn-rich clinopyroxene. No valid correlation between the chemical composition and optical properties is evident with the present data. The manganeserich clinopyroxene occurs associated with rhodonite and calcium richer bonates of the calcite-kutnahorite series. Its $Fe + Mg$ content is rather constant, but the Mn/Ca ratios varies appreciably (Table 3). The X-ray pattern can hardly be distinguished from that of johannsenite.

| | | | | | | | Numbers of ions | | | | | |
|-------------------|------------------|------|-------------|-------|------|------|---------------------------|------|-------------|-------------------|--|--|
| | | | Weight $\%$ | | | | on the basis of 3 oxygens | | | | | |
| | SiO ₂ | MnO | CaO | MgO | FeO | Si | Mn | Ca | $_{\rm Mg}$ | Fe | | |
| Pyroxmangites | | | | | | | | | | | | |
| $B-4-14-80.8$ III | 47.7 | 44.5 | 1.09 | 5.73 | 0.95 | 1.00 | 0.79 | 0.03 | 0.18 | 0.02 | | |
| $B-5-14-67.1$ I | 49.8 | 42.6 | 1.45 | 6.31 | 1.46 | 1.01 | 0.73 | 0.03 | 0.19 | 0.02 | | |
| $B-5-14-67.1$ II | 48.2 | 42.7 | 1.47 | 6.49 | 1.34 | 1.00 | 0.75 | 0.03 | 0.20 | 0.02 | | |
| $B-5-14-69.9$ I | 48.9 | 41.8 | 2.09 | 6.23 | 1.86 | 1.00 | 0.73 | 0.05 | 0.19 | 0.03 | | |
| $B-5-14-72.0$ I | 48.5 | 42.4 | 1.43 | 6.88 | 1.34 | 1.00 | 0.74 | 0.03 | 0.21 | 0.02 | | |
| $B-5-14-72.0$ II | 48.2 | 42.4 | 1.43 | 7.13 | 1.30 | 0.99 | 0.74 | 0.03 | 0.22 | 0.03 | | |
| $B-5-14-72.0$ IV | 48.7 | 42.5 | 1.39 | 6.55 | 1.30 | 1.00 | 0.73 | 0.05 | 0.20 | 0.02 | | |
| $B-5-14-78.8$ II | 48.2 | 42.9 | 1.57 | 6.75 | 1.28 | 0.99 | 0.75 | 0.03 | 0.21 | 0.03 | | |
| $B-5-14-78.8$ III | 48.1 | 42.1 | 1.41 | 7.31 | 1.29 | 0.99 | 0.74 | 0.03 | 0.23 | 0.02 | | |
| $B-5-14-78.8$ IV | 47.8 | 42.2 | 1.38 | 7.36 | 1.26 | 0.99 | 0.74 | 0.03 | 0.23 | 0.02 | | |
| $B-5-22-64.6$ II | 52.6 | 30.2 | 1.70 | 14.98 | 1.62 | 1.01 | 0.49 | 0.03 | 0.43 | 0.03 | | |
| $B-5-22-64.6$ IV | 52.7 | 30.4 | 1.61 | 14.90 | 1.65 | 1.01 | 0.49 | 0.03 | 0.42 | 0.03 | | |
| $B-5-22-65.8$ I | 52.3 | 30.6 | 1.46 | 14.97 | 1.77 | 1.01 | 0.50 | 0.03 | 0.43 | 0.03 | | |
| $B-5-22-65.8$ II | 50.1 | 41.2 | 1.72 | 6.82 | 1.84 | 1.01 | 0.70 | 0.04 | 0.21 | 0.03 | | |
| $B-5-22-65.8$ IV | 50.2 | 41.4 | 1.94 | 6.44 | 1.94 | 1.01 | 0.71 | 0.04 | 0.19 | 0.02 | | |
| Rhodonites | | | | | | | | | | | | |
| $B-5-10-80.2$ II | 48.9 | 41.6 | 5.01 | 4.50 | 0.62 | 1.01 | 0.73 | 0.11 | 0.14 | 0.01 | | |
| $B-5-10-92.0$ I | 46.8 | 43.0 | 4.83 | 4.15 | 0.05 | 0.99 | 0.77 | 0.11 | 0.13 | 0.01 | | |
| $B-5-10-97.8$ IV | $\bf 48.5$ | 45.0 | 6.00 | 1.61 | 0.15 | 1.01 | 0.79 | 0.13 | 0.05 | 0.00 ₁ | | |

Table 2. Microprobe analysis of pyroxenoids from Mn protores of Buritirama

Table 3. Microprobe analysis of clinopyroxenes from Mn protores of Buritirama

| | | | | Weight $\%$ | | | | | | |
|------------------|---|------|------|-------------|------|---------------|-------------------|--|--|--|
| | SiO ₂ | MnO | CaO | MgO | FeO | $\rm Al_2O_3$ | Na ₂ O | | | |
| $B-5-10-80.2$ I | 53.0 | 14.9 | 15.6 | 14.9 | 1.02 | 0.23 | 0.24 | | | |
| $B-5-10-80.2$ II | 53.3 | 18.3 | 12.4 | 14.6 | 1.33 | 0.12 | n.d. | | | |
| $B-5-10-97.8$ II | 53.7 | 13.2 | 19.4 | 13.7 | 0.17 | 0.10 | n.d. | | | |
| $B-5-10-97.8$ IV | 54.1 | 11.4 | 20.4 | 14.4 | 0.12 | 0.07 | n.d. | | | |
| | Numbers of ions on the basis of 3 oxygens | | | | | | | | | |
| | Si | Mn | Ca | Mg | Fe | Al | Na | | | |
| $B-5-10-80.2$ I | 1.00 | 0.24 | 0.32 | 0.42 | 0.02 | 0.01 | 0.01 | | | |
| $B-5-10-80.2$ II | 1.01 | 0.29 | 0.25 | 0.41 | 0.02 | | | | | |
| $B-5-10-97.8$ II | 1.01 | 0.21 | 0.39 | 0.38 | | | | | | |
| $B-5-10-97.8$ IV | 1.01 | 0.18 | 0.40 | 0.40 | | | | | | |

Tephroite

Tephroite is a mineral as common in the Mn marble of Buritirama as is in other Brazilian carbonatic protores (Serra do Navio and Lafayette). In Buritirama this mineral commonly grows as oriented poikilitic crystals of up to ³ mm length.

5 measurements from 2 samples gave: $2V_x$ 62-69°.

In addition to Mn_2SiO_4 , Buritirama tephroite contains about 15 mol $\%$ of Mg_2SiO_4 (forsterite), and trifling amount of Fe_2SiO_4 (fayalite) and Ca_2SiO_4 (larnite) (Fig. 5). The forsterite contents manifests itself in a notable placement of the X-ray reflections towards lower angles as compared with those of pure Mn_2SiO_4 .

Table 4. Microprobe analysis of tephroites from Mn protores of Buritirama

| | Weight $\%$ | | | | | | Number of ions on the basis of 4 oxygens | | | | | |
|-------------------|-------------|----------|------|-------|--------------|------|---|------|--------------------------|------|------|--------------------------|
| | | SiO, MnO | CaO | MgO | $_{\rm FeO}$ | ZnO | Si | Mn | Ca | Mg | Fe | \mathbf{Zn} |
| $B-4-14-80.8$ I | 30.7 | 59.4 | 0.06 | 6.24 | 3.49 | n.d. | 0.99 | 1.61 | $\frac{1}{2}$ | 0.30 | 0.09 | |
| $B-4-14-80.8$ II | 30.9 | 59.4 | 0.08 | 6.83 | 3.42 | n.d. | 0.99 | 1.60 | $\overline{}$ | 0.31 | 0.09 | $\overline{}$ |
| $B-4-14-80.8$ III | 31.6 | 59.0 | 0.07 | 6.73 | 3.55 | n.d. | 1.00 | 1.60 | \rightarrow | 0.31 | 0.09 | and it |
| $B-5-14-78.8$ I | 31.6 | 54.2 | 0.06 | 9.98 | 4.46 | 0.16 | 0.99 | 1.42 | $\frac{1}{2}$ | 0.46 | 0.12 | $\overline{}$ |
| $B-5-14-78.8$ II | 31.0 | 54.4 | 0.07 | 9.70 | 4.72 | 0.12 | 0.98 | 1.42 | \rightarrow | 0.45 | 0.12 | |
| $B-5-14-78.8$ III | 31.3 | 54.0 | 0.07 | 9.98 | 4.63 | 0.21 | 0.98 | 1.41 | $\overline{}$ | 0.46 | 0.12 | 0.01 |
| $B-5-14-78.8$ IV | 31.0 | 53.4 | 0.06 | 10.94 | 4.58 | 0.22 | 0.97 | 1.42 | $\overline{}$ | 0.51 | 0.12 | 0.01 |

Manganese amphibole

The manganese clinoamphibole, which is generally formed in parts of the rocks where the partial pressures of $H₂O$ was high, can be named either a manganese-rich cummingtonite or a magnesium-rich dannemorite. Under the microscope this mineral is colorless, or rarely yellow. It may show polysynthetic twinning and its $2V_x$ is $\sim 85^\circ$. The genesis of this mineral seems to be related to the (hydrothermal?) alteration of pyroxenoids and pyroxenes in veins. A few microprobe data are presented in Table 5.

Table 5. Microprobe analysis of amphiboles from Mn protores of Buritirama

| | Weight $\%$ | | | | | | | | | |
|------------------|------------------|---------------|-------|---------------|--|--------------------------|---------------------------------|-------------------|--|--|
| | SiO ₂ | $\rm Al_2O_3$ | CaO | $_{\rm{MgO}}$ | MnO | FeO | TiO, | Na ₂ O | | |
| $B-5-10-92.0$ II | 56.9 | 0.06 | 11.19 | 22.0 | 5.83 | n.d. | n.d. | 0.34 | | |
| $B-5-14-69.9$ I | 56.4 | 0.06 | 0.85 | 20.1 | 18.70 | 2.41 | n.d. | 0.05 | | |
| $B-5-22-64.6$ I | 54.4 | 0.19 | 0.92 | 19.5 | 19.94 | 2.17 | n.d. | 0.08 | | |
| $B-5-22-64.6$ IV | 56.4 | 0.14 | 0.91 | 19.4 | 19.14 | 2.17 | $\mathbf{n} \cdot \mathbf{d}$. | n.d. | | |
| | | | | | Numbers of ions on the basis of $23(0, \text{OH})$ | | | | | |
| | Si | Al | Ca | Mg | Mn | Fe | Tì | Na | | |
| $B-5-10-92.0$ II | 7.99 | 0.01 | 1.68 | 4.60 | 0.69 | $\overline{}$ | | 0.09 | | |
| $B-5-14-69.9$ I | 8.03 | 0.01 | 0.13 | 4.25 | 2.25 | 0.29 | $\overline{}$ | 0.01 | | |
| $B-5-22-64.6$ I | 7.92 | 0.03 | 0.14 | 4.22 | 2.46 | 0.26 | Î. | 0.02 | | |
| $B-5-22-64.6$ IV | 7.99 | 0.02 | 0.14 | 4.14 | 2.30 | 0.26 | | | | |
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Garnets

Garnet compositions are close to the spessartite end member but usually comprise small proportions of grossularite, pyrope, almandine, and andradite (Table 6). Garnet is closely associated with carbonates either as inclusions or as ^a host to carbonatic pigments. In general coarser garnets are confined to silicate (pyroxenoid) richer parts. The cell parameter for garnets varies from 11.570 to 11.634 Â and the measured refractive indices (1.71-1.79) indicate ^a spessartitic end member with ^a tendency of some deviation towards ularite.

Manganoan phlogopite

Manganoan phlogopite (manganophyllite) is a common hydroxylbearing mineral in most assemblages. The manganese content varies little, but the amounts of iron, although low compared with magnesium, differ. The substitution of Na⁺ for K⁺ is generally negligeable (Table 7).

Ore minerals

Braunite is an abundant mineral in the braunite marble protore. It presents a massive aspect being frequently cut by veins containing: carbonates $+$ manganophyllite + manganese clinoamphibole. Bixbyite is an occasional ciate. The braunite's $SiO₂$ content is nearly constant and supports the ideal formula: $3 \text{ Mn}_2\text{O}_3 \cdot \text{MnSiO}_3$. Substitutions of Fe and Al for Mn³⁺ in the first part of this formula arc quite restricted, as are the substitution of Ca and Mg for Mn^{2+} in the second part of the formula.

Alabandite was found in one sample (B-4-14 drill) where it is associated with accessory pyrrhotite in ^a rhodochrosite-tephroite-spessartite rock. It occurs as large xenomorphic grains containing some Fe and traces of Mg.

Sphalerite occurs as an accessory. Beside Zn some Fe and Mn were also found. Its microscopic red colour makes it almost indistinguishable from pyrophanite - $MnTiO₃$ - a relatively common protore accessory. Microprobe analyses show some CaO and FeO in the latter mineral. Hausmannite is a rare accessory found in the braunite marble protore. It contains about 5 mol $\frac{9}{6}$ of $Fe₃O₄$, besides $Mn₃O₄$.

Mineral assemblages

The following manganese-bearing mineral assemblages were encountered among the analysed suits.

- 01) carbonate + pyroxmangite + tephroite
- 02) tephroite + carbonate + spessartite + pyrophanite
- 03) tephroite + carbonate + pyroxmangite + spessartite
- 04) carbonate + tephroite + spessartite + alabandite $($ + sphalerite)
- 05) carbonate + braunite + manganophyllite + manganese amphibole
- 06) manganese clinopyroxene + carbonate + manganophyllite + spessartite
- 07) manganese clinopyroxene + rhodonite + carbonate
- 08) $rho = +$ carbonate + spessartite
- 09) manganese clinopyroxene + rhodonite + carbonate + braunite
- 10) manganese clinopyroxene + manganese clinoamphibole + manganophyllite + spessartite + carbonate
- ¹¹ pyroxmangite + manganophyllite + carbonate
- 12) pyroxmangite + manganese clinoamphibole + carbonate + spessartite
- 13) pyroxmangite $+$ carbonate $+$ spessartite $+$ manganophyllite
- 14) carbonate + braunite + hausmannite $(+bixbyite)$
- ¹⁵ carbonate + hausmannite
- 16) rhodonite + carbonate + manganophyllite + K-spar + spessartite \pm baryte
- 17) quartz + rhodonite + carbonate
- 18) quartz + braunite + rhodonit

Neglecting the small amounts of Fe and assuming CO_2 , H_2O and S to be perfectly mobile components, one is left either with the ⁵ component system $Mn-Ca-Mg-Al-Si$ or, including K (in manganophyllite and K-spar), with a 6 component system. Manganophyllite free assemblages contain mostly 4 phases, which makes the system trivariant.

Fluid inclusions

In addition to chemical and mineralogical data first results from fluidinclusion research may be interesting for evaluation of P-T data.

In quartz grains coexisting with different manganese minerals several generations of fluid inclusions have been recognized which are mainly bound to small healed fractures. In general earlier generations have very high concentrations of CO_2 , whereas later ones show low values of X_{CO_2} with three phases (CO₂l, CO₂g, H₂O1). Latest stages present water solutions of high salinity, even with small solid inclusions.

The earliest inclusions, supposed to have formed during the main metamorphic event, constantly show one or two phases of $CO₂$. The observed homogenizations, always towards liquid $CO₂$, indicate hight densities. In the diagram (Fig. 6) two main populations of homogenization temperatures and corresponding densities (LANDOLT-BOERNSTEIN tables, 1960) can be recognized, indicating decompression of the fluid.

Fig. 6. Critical densities (g/cm³) of CO₂ and temperature of homogenization of 394 fluid inclusions in quartz crystals from Buritirama (13-5-10) braunite marble manganese protore. The quartz occurs in small aggregates associated with different manganese minerals.

The earlier population, with higher densities can be interpreted in two ways: a) as a true population with a main value of homogenization temperature of $8-9^{\circ}$ C and a main density of 0.87, or b) only the highest values with homogenization temperatures from $-5-0$ °C and densities of 0.95–0.93 are significant, the other values testifying later stages of decompression. Both interpretations are feasible, and at the moment no preference can be given to one or the other.

The linear extrapolation of P-V-T data for $CO₂$ (KENNEDY, 1954) from isochores $d = 0.95$ g/cm³ (maximum values) and $d = 0.87$ g/cm³ (main value) gives P-T-gradients of 10° C/78.5 bars (33[°]C/km, maximum value) or 10° C/ ⁵³ bars (54°C/km, main value). As influences of decompressibility and dilatancy of quartz can be neglected, pressures can be calculated, if temperatures are known.

Physical-chemical conditions during metamorphism

The mineral assemblages in the rocks associated with the manganeseprotores indicate a metamorphism in the lower to middle amphibolite facies. No invariant assemblages that could be used as a precise geothermometer and geobarometer were encountered. From the composition of the Ca-Mn-Carbonates ^a minimum temperature of 540° ^C can be deduced. In some rocks the isobaricly invariant assemblage diopside + tremolite + calcite + dolomite + Qz was found, indicating a high X_{CO_2} and temperatures between 500°C and 600° C depending upon the total pressure. These high partial pressures of CO_2 inferred by the mineral assemblages support the assumption that the first generation fluid inclusions with the high $CO₂$ content formed at the main metamorphic event. The mean density of $CO₂$ in most inclusions of the first generation (Fig. 6) is 0.87 g/cm². Assuming a temperature of $550 \pm 50^{\circ}$ C, a total pressure of 3 ± 0.3 Kb results.

For the estimation of the oxygen fugacity during metamorphism the assemblages manganese carbonate-braunite, carbonate-braunite-hausmannite and hausmannite-carbonate are important. In the pure system Mn-Si-C-0 (PETERS, VALARELLI and CANDIA, 1974) the second assemblage would be isobaricly invariant, that would fix log f_{Q_2} at -10 . However, the carbonate is not pure $MnCO₃$, but a solid solution of $CaCO₃$ and $MnCO₃$, meaning an activity of less than 1 for MnCO₃ in the equilibrium $6MnCO₃ + O₂ =$ $2Mn₃O₄ + 6CO₂$ resulting in an even higher f_{O2} compared with pure $MnCO₃$.

Ore genesis

The manganese ore of Buritirama has ^a complex history, of which only the latest stages can be evaluated with some confidence.

The latest event is the enrichment of manganese by supergene weathering from Mn-bearing calc-silicate marbles. Although the bulk of the carbonates has a relatively low Mn-content they weather easily, giving an ore rich in highly oxidized manganese oxides and hydroxides. A regional metamorphism in amphibolite facies has determined the mineralogy and texture of the protore. The effects of ^a later retrograde metamorphism are small. During the strong metamorphic event manganese was enriched in the silicate phases (pyroxenoides, garnets and tephroite) as they formed from the reaction of Ca-Mn-Carbonates with quartz and phyllosilicates. Contemporaneously dolomite reacted with $SiO₂$ to form diopsidic clinopyroxenes and calcite.

The strong regional metamorphism has obliterated practically all sedimentological evidences of the original deposition. The succesion of quartzites, manganiferous calc-silicate marbles and muscovite quartzites in Buritirama is comparable with the manganiferous limestone dolomite formations as defined by VARENTSOV (1964). Together with the relatively high oxidation state and high Ba-content of the Buritirama protores a succession formed in ^a shallow platform basin is indicated. The graphite-bearing Mn-rich protores of Amapa

and Lafayette associated with black bitumous pyritiferous shales formed in an enviroment transitional to deeper water. The ore material was probably derived from weathering of the gneisses and amphibolites of the underlying Xingu Complex. An additional volcanic supply cannot be excluded as there is evidence for basic volcanic activity in the Grâo Para Group in the Serra dos Carajâs region.

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