Abstract: The Nabeba high-grade iron deposit (Republic of the Congo) is hosted by banded iron formation (BIF) in the Ivindo Basement Complex, which lies in the northwestern part of the Congo Craton. The Nabeba BIF is intercalated with chlorite-sericite-quartz schist and comprises two facies (oxide and a carbonate-oxide). In this study, whole-rock and LA-ICP-MS magnetite geochemistry of the BIF was reported. Magnetite samples from both BIF facies had fairly similar trace element compositions except for the rare earth element plus yttrium (REE + Y) distribution patterns. The high V, Ni, Cr, and Mg contents of the magnetite in the Nabeba BIF could be ascribed to the involvement of external medium-high temperature hydrothermal fluids during their deposition in relatively reduced environment. The Post-Archean Australian Shale (PAAS)-normalized REY patterns of the Nabeba BIF magnetite were characterized by LREE depletion coupled with varying La and positive Eu anomalies. Processing of the information gathered from the geochemical signatures of magnetite and the whole-rock BIF suggested that the Nabeba BIF was formed by the mixing of predominantly anoxic seawater (99.9%) with 0.1% of high-temperature (>250 °C) hydrothermal vent fluids, similar to the formation mechanism of many Archean Algoma-type BIFs reported elsewhere in the world.

Keywords: magnetite chemistry; banded iron formation (BIF); Nabeba deposit; Ivindo basement complex; Republic of the Congo

1. Introduction

Algoma-type banded iron formations (BIFs) are increasingly explored and studied due to their rich endowments of iron (e.g., the Águas Claras and Pico Mine, Brazil; [1]) and gold (e.g., the Meliadine, Canada; [2]). In addition to their economic significance, BIFs have immense scientific value in understanding the Precambrian hydrosphere, biosphere, and atmosphere [3–7]. BIFs are chemical sedimentary rocks by definition and comprise alternating chert- and iron-rich (>15% Fe) layers [8–10]. These rocks are associated with Archean to Paleoproterozoic volcano-sedimentary sequences in greenstone belts associated with back-arc or intra-arc basin setting.

The northwestern margin of the Congo Craton extends from Cameroon southward through northern Gabon to the northwestern Republic of the Congo (R.C.), where various BIF-hosted iron deposits were discovered within the Precambrian terranes and the narrow greenstone belts around them [11–15]. The BIFs in Central Africa and the Congo are much less studied than those in Cameroon. In the northwestern Congo, recent exploration by the Core Mining Ltd., Equatorial Resources, and Congo Iron SA discovered several high-grade iron deposits, including Avima, Badondo, Nabeba, and Cabosse [16] (Figure 1). Very little
is known about these BIF-hosted iron deposits, including the BIF depositional environment prior to ore enrichment [16–18]. Among these iron deposits, the Nabeba iron deposit is renowned for its strong economic potential, which attracts much attention.

The Precambrian Nabeba BIF is associated with mafic and felsic volcanic rocks (Figure 2a). Their lithological associations are similar to those of the well-studied Algoma-type Musselwhite and Meliadine BIFs. Recent studies on Algoma-type BIFs demonstrated that their depositional settings [2,20–24] involve submarine hydrothermal activity and a seawater component with varying detrital/volcaniclastic input. Despite being influenced by post-depositional processes, such as regional greenschist-/amphibolite-facies metamorphism, mineral geochemistry from Si-rich and Fe-rich layers can be used to constrain petrogenesis and depositional environment [2,25–33], especially with the recent advances in the laser ablation-inductively coupled plasma-mass spectrometry (LA-ICP-MS) analytical technology [28].

This study focuses on the BIF that hosts the Nabeba iron deposit—the deposit with the highest iron ore potential amongst the other BIFs in the Invido Basement Complex (IBC) of the Republic of Congo (Figure 1)—but its host BIF depositional environment and genesis are not well-studied. In this paper, we reported BIF petrographic features and whole-rock geochemical and LA-ICP-MS magnetite trace element compositions. Based on comparisons with well-studied BIFs around the world, we discussed the controlling factors on the magnetite trace element characteristics and the implications that all of these data have for the BIF depositional environment and petrogenesis at Nabeba.
2. Regional Geology

Due to the inaccessibility of the region to conduct regionally extensive geological works and the lack of extensive outcrops, the regional geology reported in this paper was based on the few outcrops that we encountered at the company site and environs, the company records on exposures during its geological exploration expeditions, and the scant literature available on the Nabeba area (e.g., the publication of Meloux et al. [34]). In NW Congo, a granitoid massif, which is known as the Chaillu Massif [35] and an equivalent of the IBC, contains two generations of granitoids with mainly N-S foliation. Greenstone belts (including BIF) and schist are found in several locations within the Chaillu Massif, including the Mayoko and the Zanaga Regions in southern Congo [36] and at Mt. Nabeba. To the east of Mt. Nabeba, the margins of the Chaillu Massif are overlain by low-grade metamorphic rocks of the Proterozoic Sembé-Ouesso Group (i.e., a sedimentary sequence...
of sandstone-shale and diamicite \cite{19,34,35,37} (Figure 1). Cenozoic sediments of the Congo Basin cover much of the area, including the surroundings of Mt. Nabeba.

The Precambrian Nabeba iron deposit in northwestern Congo (Figure 1) is geologically located in the Archean IBC. The IBC comprises mainly two types of rock formation, namely supracrustal and crystalline rocks (Figure 1; \cite{19,34,35,37}). The supracrustal rocks are mainly volcano-sedimentary in origin and comprise metamorphosed, EW-trending greenstone belts (commonly chlorite-sericite-quartz schists and amphibolite) and banded iron formations with mafic and/or ultramafic intrusive rocks \cite{34}. The Archean greenstone belts narrowly frame the crystalline rocks along regional NE-trending structural corridors. The crystalline rocks generally comprise, as elsewhere in the Congo Craton, cataclasized granites, orthogneiss, and migmatises. These rocks constitute the greater part of the basement (80% to 90%). However, the most widespread lithological assemblages are calc-alkaline granites, biotite, green hornblende granodiorites, and, more locally, quartz meta-diorites or metagabbros \cite{34}.

In the Souanké area, which is at the northeastern margin of the IBC, the greenstone belts and the amphibolite associated with banded iron formation (Figure 2a) are thought to be the remnants of greenstone belts, including those at Nabeba \cite{34}. The good exposures in the Souanké area reveal intrusions of Archean TTG suites and potassic granite in the amphibolite, suggesting that the Amphibolite and the BIF are at least of Precambrian age \cite{38,39}.

3. Deposit Geology

The principal rock types that represent the volcano-sedimentary units in the structurally complex Nabeba project area are BIF and chlorite-sericite-quartz schists \cite{40}. The Nabeba rocks are metamorphosed to the amphibolite facies and strongly deformed \cite{16}.

Structural geological studies revealed at least two major deformation events (D\textsubscript{1} and D\textsubscript{2}) \cite{16}. D\textsubscript{1} consists of the NE-trending S\textsubscript{1} foliation and the F\textsubscript{1} north-plunging isoclinal fold, while D\textsubscript{2} formed regional N-S-trending folds (F\textsubscript{2}) and extensive E-W-trending S\textsubscript{2} shearing \cite{16}. The Nabeba BIF was suggested to be structurally dismembered and fan-shaped, which was confined by NE-trending thrust/reverse faults. These faults converged into an E-W trending regional shear zone north of Nabeba \cite{16}.

The Nabeba BIF consists of alternating thin iron oxide-rich and cherty bands with minor iron carbonate minerals, such as siderite and ferroan-magnesite. The iron oxide minerals comprise hematite, magnetite, and martite. The BIF is crosscut by quartz veins of varying thicknesses \cite{16}.

The Nabeba BIF contains the Nabeba iron deposit: an inferred resource of 436.3 Mt at a grade of 62.3% Fe (Figure 2a; \cite{16}). Three mineralization styles are present: (1) supergene goethite-hematite ore (>62.3% Fe); (2) itabirite-type mineralization (i.e., hematite-enriched BIF; 34% Fe) (Figure 2b); (3) hypogene iron ore (45% to 55% Fe) \cite{16}.

4. Sampling and Analytical Methods

Thirty-three representative BIF samples were collected from drill cores (Figure 3) and outcrops (Figure 4a,b) for petrographic observations, from which ten samples were selected for whole-rock geochemical analysis and five for in situ LA-ICP-MS magnetite trace element analyses (Table 1).

These samples contain iron oxide bands with dominantly anhedral magnetite grains and rare hematite. Polished thin sections were prepared and examined using both transmitted and reflected light microscopy along with scanning electron microscopy (SEM) in order to select suitable geochemical analysis site with minimal intergranular matrix or mineral (e.g., quartz, siderite, monazite) inclusions.

Whole-rock major and trace (including rare earth element (REE)) analyses were performed at the ALS Laboratory (Guangzhou, China). The samples were crushed and powdered in an agate mill. Major elements were analyzed with X-ray fluorescence (XRF).
spectroscopy after the powders were fused with lithium borate into glass disks. The analytical uncertainties varied from 1% to 0.04%.

Figure 3. Drill-hole stratigraphic columns at the Nabeba BIF deposit.
Table 1. Samples collected from the Nabeba iron deposit.

| Sample Type | N° | Sample Name | Location | Coordinates East | North | Depth (m) | BIF Facies | Texture | Major Mineral Assemblage |
|-------------|----|-------------|----------|------------------|-------|-----------|------------|---------|--------------------------|
| BIF (Drill Core) |     |             |          |                  |       |           |            |         |                          |
| 1 | CG-NB 1 | NBDD012 | 387521 | 203459 | 224 | Carbonate-oxide | Fine banded, brown, grey, and black alternating bands |         | Magnetite + siderite + quartz |
| 2 | CG-NB 2 | NBDD001D | 388900 | 203208 | 86 | Oxide | Banded, alternation of white, brown, and grey |         | Magnetite + quartz |
| 3 | CG-NB 3 | NBDD001D | 388900 | 203208 | 90 | Oxide | Fine banded, alternation of white, brown, and grey |         | Magnetite + quartz + hematite |
| 4 | CG-NB 4 | NBDD012D | 387521 | 203459 | 215 | Oxide | Banded, cut by quartz veins |         | Magnetite + quartz |
| 5 | CG-NB 6 | NBDD001D | 388900 | 203208 | 96 | Oxide | Fine banded, grey, brown, and white fold bands |         | Magnetite + quartz + hematite |
| 6 | CG-NB 7 | NBDD004D | 388300 | 203588 | 156 | Oxide | Banded, brown, and white |         | Magnetite + quartz |
| 7 | CG-NB 8 | NBDD004D | 388300 | 203588 | 185 | Oxide | Banded, brown, and white, cut by quartz veins |         | Magnetite + quartz |
| 8 | CG-NB 9 | NBDD0012 | 387521 | 203459 | 220 | Oxide | Folded banded, brownish, cut by quartz veins |         | Magnetite + quartz |
| 9 | CG-NB10 | NBDD0012 | 387521 | 203459 | 208 | Oxide | Fines banded, grey to brown, crosscutting veins |         | Magnetite + quartz + siderite |
| 10 | CG-NB11 | NBDD0012 | 387521 | 203459 | 98 | Carbonate-oxide | Fines banded, gray to brown, and white alternating bands |         | Magnetite + quartz + siderite + ferroan-magnesite |
| BIF (Outcrop) |     |             |          |                  |       |           |            |         |                          |
| 11 | CG-C2 | Outcrop | 386688 | 202204 | | Oxide | Fine banded, grey, brown, and white alternating bands |         | Magnetite + quartz + hematite + martite |
| 12 | CG-C3 | Outcrop | 386221 | 202029 | | Oxide | Brown and white alternating bands |         | Magnetite + quartz + hematite + martite |
| 13 | CG-C4 | Outcrop | 386711 | 202232 | | Carbonate-oxide | Banded, brown, and grey |         | Magnetite + quartz + hematite + ferroan-dolomite |
| 14 | CG-C6 | Outcrop | 387214 | 202232 | | Carbonate-oxide | Fine banded, brown, grey, and white rhythms |         | Magnetite + quartz + hematite + ferroan-dolomite |
| 15 | CG-C12 | Outcrop | 387176 | 202145 | | Oxide | Banded, grey to brown slightly weathered, foliated, dominated by hematite bands |         | Magnetite + hematite + martite + quartz |
| 16 | CG-C14 | Outcrop | 386588 | 202279 | | Carbonate-oxide | Banded, light grey to grey, slightly weathered, presence of sugary quartz in the chert bands |         | Magnetite + quartz + hematite + ferroan-dolomite |
| 17 | CG-C21 | Outcrop | 386520 | 202354 | | Oxide | Fine banded, brown, grey, and white rhythms |         | Magnetite + quartz + hematite + martite |
| 18 | CG-C23 | Outcrop | 386547 | 202390 | | Oxide | Fine banded, grey slightly weathered, foliated, dominated by hematite-goethite bands |         | Magnetite + quartz + hematite |
| 19 | CG-C25 | Outcrop | 386720 | 202287 | | Oxide | Fine banded, grey slightly weathered, foliated, dominated by hematite-goethite bands |         | Magnetite + martite + quartz |
Figure 4. Field photos of (a) BIF outcrop at Nabeba, showing minor folding of the alternating hematite and chert layers; (b) oxide facies-BIF, showing the fine alternating layers of chert and iron oxides; (c) oxide facies-BIF, showing the fine alternating layers of chert, specularite, and magnetite; (d) slightly-folded carbonate-oxide facies-BIF, showing the alternating carbonate, magnetite, and chert layers.

Trace element compositions were analyzed with solution ICP-MS, with approximately 2% to 5% analytical uncertainties. The sample powder was mixed with distilled HF and HNO₃ acids in Teflon screw-up capsules at 140 °C and then dried and digested with HNO₃ acid at 190 °C for 48 h in an oven. The solution was dried again and digested with HNO₃ acid at 150 °C for 12 h. A solution with 800 ng/g rhodium was added into the dissolved samples as an internal standard. The precision for trace elements and REEs varies between 0.1% and 0.5%. For REE-Y, the detection limits were 0.01 ppm. Data quality was monitored by analyzing various standards between unknown samples.

LA-ICP-MS trace element analysis was performed on selected magnetite grains with an Agilent 7900 quadrupole ICP-MS (Agilent Technologies, Santa Clara, CA, USA) attached to a Photon Machines Analyte (Photon-Machines INC., Redmond, WA, USA) HE 193 nm Excimer laser ablation system equipped with a SQUID signal smoothing (Applied Spectra, INC., West Sacramento, CA, USA) device. The analysis was performed at the In-situ Mineral Geochemistry Laboratory, Ore Deposit and Exploration Centre (ODEC), Hefei University of Technology (Anhui Province, China). Sample ablation was done using He as a carrier gas, which mixed with the Ar make-up gas before entering in the ICP. The 30 µm laser spot and the 8 Hz frequency were maintained throughout the analysis. Analysis of all grains was done using laser energy at 3 J/cm². Each analysis comprised a gas blank (laser-off) background measurement of 20 s, followed by a 40 s laser-on sample signal measurement. Calibration was done using multiple reference materials (BCR-2G, SRM 612, and SRM 610), and $^{57}$Fe was used as an internal standard [41]. The ICPMSDataCal program was used for data processing, following the method described by Liu et al. [42]. This software is generally used for the integrated selection of contextual and for signal analysis, quantitative calibration, and correction of time drift. Fifty-four isotopes were measured. Forty-seven sample analysis spots were bracketed by analysis of the standard (e.g., BCR-2G, SRM610, and SSRM612). In the study, the detection limits were significantly lower than the rare earth element (REE) contents of our magnetite samples (Tables S1 and S2).

The Post-Archean Australian Shale (PAAS; [43], subscript “SN”) was used to normalize the REE+Y concentrations to minimize the influence of potential terrigenous input. The PAAS-normalized La (La_{SN}) anomalies were calculated by $(La/La^{*})_{SN} = (La/Pr)_{SN}$ after
Bau et al. [44], because these REEs were taken as reference elements due to their coherent behavior during geochemical processes and predictable fractionation [45]. The anomalous abundance of Ce\textsubscript{SN}, Eu\textsubscript{SN}, Gd\textsubscript{SN}, and Pr\textsubscript{SN} were calculated as suggested by Bau and Dulski [3]: \((\text{Ce}/\text{Ce}^*)_{\text{SN}} = \text{Ce}_{\text{SN}} / (0.5\text{La}_{\text{SN}} + 0.5\text{Pr}_{\text{SN}})\), \((\text{Eu}/\text{Eu}^*)_{\text{SN}} = \text{Eu}_{\text{SN}} / (0.67\text{Sm}_{\text{SN}} + 0.33\text{Tb}_{\text{SN}})\), \((\text{Gd}/\text{Gd}^*)_{\text{SN}} = \text{Gd}_{\text{SN}} / (0.33\text{Sm}_{\text{SN}} + 0.67\text{Tb}_{\text{SN}})\), and \((\text{Pr}/\text{Pr}^*)_{\text{SN}} = \text{Pr}_{\text{SN}} / (0.5\text{Ce}_{\text{SN}} + 0.5\text{Nd}_{\text{SN}})\).

5. Results

5.1. BIF Petrography

5.1.1. Oxide-Facies-BIF

The oxide-facies-BIF was fine- to coarse-grained and comprised alternating whitish chert- and dark-brown iron oxide-bands, which were <1 to 1.5 mm thick (Figure 4b,c). The oxide-facies-BIF consisted of quartz, magnetite, hematite, and martite, and minor fluorapatite and sulfides. Quartz occurred as isolated anhedral-euhedral fine grains (<45–400 \(\mu \text{m}\)) or formed granular aggregates 150 to >200 \(\mu \text{m}\) in size (Figure 5a,b). It locally displayed undulose extinction due to deformation. Magnetite appeared as anhedral crystals or aggregates of 25–550 \(\mu \text{m}\) in size (Figure 5a–d). Magnetite was partially replaced by hematite or as relicts in hematite grains (Figure 5d). Many magnetite grains contained quartz inclusions. Hematite occurred as massive anhedral aggregates of 25–300 \(\mu \text{m}\) in size (Figure 5c,d) and was commonly formed as pseudomorphic replacement of magnetite (Figure 5c,d). Martite occurred as granular mass with grain size of 50–300 \(\mu \text{m}\) (Figure 5c).

5.1.2. Carbonate-Oxide Facies-BIF

The Nabeba carbonate-oxide facies-BIF was dark-brown and fine- to coarse-grained (Figures 4d and 6a,b). The sample hand-specimens displayed alternating bands (<1 to 4 mm thick) of chert and ferric minerals (siderite, magnetite, and ferroan-magnesite) (Figure 6a,b). Under the microscope, the Nabeba carbonate-oxide facies-BIF contained accessory apatite,
K-feldspar, chlorite, monazite, galena, sphalerite, and sulfides (pyrite, arsenopyrite, and chalcopyrite) (Figure 6c,d).

**Figure 6.** Core sample photos (a,b) and photomicrographs of the carbonate-oxide facies-BIF (c,d): (a) folded micro-banding of alternating quartz and ferric minerals (magnetite, siderite, and ferroan-magnesite) cut by a quartz vein (sample CG-NB10); (b) alternating thick quartz-rich and ferric mineral-rich (magnetite, siderite, and ferroan-magnesite) layers (sample CG-NB10); (c) euhedral pyrite inclusions in iron oxide-rich layer (sample CG-NB1). Note the lineation of magnetite, siderite, and quartz; (d) subhedral magnetite in the carbonate layers, mostly dominated by siderite (sample CG-NB1). Abbreviations: Fe-Mgst: ferroan-magnesite; Mag: magnetite; Py: pyrite; Qtz: quartz; Sid: siderite.

Magnetite, the dominant iron-bearing ferric mineral, was euhedral-subhedral and <50–600 µm in size. Magnetite formed massive aggregates or occurred as minor inclusions in chert- or iron carbonate-rich layers (Figure 6b). Rare fine quartz and monazite inclusions were found in the massive magnetite aggregates (Figure 6c).

Siderite was often associated with ferroan-magnesite, and both minerals appeared as massive aggregates in the continuous iron carbonate-rich bands (Figure 6a,b). Siderite was often associated with magnetite or quartz (Figure 6c). Anhedral ferroan-magnesite occurred as micro-laminae in a siderite groundmass and appeared as an alteration product of siderite.

Quartz was anhedral and formed micro-laminae between magnetite-rich and carbonate-rich laminae (Figure 6b). Quartz displayed undulose extinction and was often associated with pyrite, chlorite, or ferro-microcline as inclusions in granoblastic siderite (Figure 6c).

### 5.2. Whole-Rock Geochemistry

#### 5.2.1. Major Elements

Major element analyses of the representative Nabeba BIF were performed on oxide- and carbonate-oxide-facies-BIF samples (six from outcrop and four from drill core) (Table 1). The results showed that the BIF contained mainly Fe$_2$O$_3$ and SiO$_2$, with a weak positive correlation between them ($r = 0.17$ and 0.27 for oxide-facies and carbonate-oxide-facies-BIF, respectively; Tables 2–4).
### Table 2. Major element oxide composition (wt.%) and ratios of the Nababa BIF.

| Sample Name | CG-NB3 | CG-NB6 | CG-C2 | CG-C21 | CG-C23–1 | CG-C23–2 | CG-C25 | CG-NB1 | CG-NB10 | CG-C6 |
|-------------|--------|--------|-------|--------|----------|----------|--------|--------|---------|-------|
| SiO₂        | 57.40  | 68.37  | 43.74 | 48.05  | 46.43    | 45.47    | 46.25  | 41.55  | 37.23   | 40.26 |
| TiO₂        | 0.01   | <0.01  | <0.01 | <0.01  | <0.01    | <0.01    | <0.01  | <0.01  | 0.02    | <0.01 |
| Al₂O₃       | 0.36   | 0.05   | 0.06  | 0.09   | 0.11     | 0.07     | 0.03   | 0.18   | 0.48    | 0.06  |
| TFe₂O₃      | 41.30  | 30.71  | 53.55 | 48.27  | 48.95    | 50.41    | 48.92  | 44.86  | 44.24   | 55.63 |
| Cr₂O₃       | <0.01  | <0.01  | <0.01 | <0.01  | <0.01    | <0.01    | <0.01  | <0.01  | <0.01   | <0.01 |
| MnO         | 0.01   | 0.01   | 0.01  | 0.01   | 0.11     | 0.12     | <0.01  | 0.44   | 0.30    | 0.15  |
| MgO         | 0.02   | 0.02   | 0.03  | 0.03   | 0.08     | 0.08     | 0.03   | 1.57   | 1.99    | 0.18  |
| CaO         | 0.01   | 0.01   | 0.01  | 0.02   | 0.02     | 0.02     | 0.03   | 0.11   | 0.18    | 0.01  |
| Na₂O        | <0.01  | <0.01  | <0.01 | <0.01  | <0.01    | <0.01    | <0.01  | <0.01  | <0.01   | <0.01 |
| K₂O         | 0.02   | 0.02   | 0.01  | 0.02   | 0.02     | 0.02     | 0.03   | <0.01  | <0.01   | <0.01 |
| P₂O₅        | 0.02   | 0.01   | 0.20  | 0.04   | 0.02     | 0.02     | 0.03   | 0.01   | 0.05    | 0.03  |
| SO₃         | <0.01  | <0.01  | <0.01 | 0.02   | 0.02     | 0.02     | 0.04   | 0.22   | 0.01    | 0.07  |
| LOI         | 0.14   | 0.26   | 1.72  | 2.72   | 3.52     | 3.18     | 3.94   | 11.17  | 14.95   | 3.00  |
| Total       | 99.15  | 99.18  | 97.60 | 96.53  | 95.76    | 96.22    | 95.34  | 88.94  | 84.50   | 96.39 |

### Table 3. Pearson’s correlation matrix for major element oxides of oxide-facies-BIF.

| SiO₂ | Al₂O₃ | TFe₂O₃ | MnO | MgO | CaO | K₂O | P₂O₅ | LOI |
|------|-------|--------|-----|-----|-----|-----|------|-----|
| SiO₂ | 1     | -      | -   | -   | -   | -   | -    | -   |
| Al₂O₃| -0.73630401| 1     | -   | -   | -   | -   | -    | -   |
| TFe₂O₃| 0.17460837 | -0.67069794| 1   | -   | -   | -   | -    | -   |
| MnO  | -0.69788362 | 0.54176046| -0.5597574| 1   | -   | -   | -    | -   |
| MgO  | -0.79756851 | 0.88406025| -0.7198373| 0.86809874| 1   | -   | -    | -   |
| CaO  | -0.74643832 | 0.93913917| -0.77845814| 0.7593399| 0.9745819| 1   | -    | -   |
| K₂O  | 1      | -1     | -1   | -1  | not| not| 1    | -   |
| P₂O₅| -0.27770787 | 0.52483223| -0.1171327| -0.2738272| 0.18820647| 0.33746412| not| 1   |
| LOI  | -0.76954675 | 0.90918882| -0.76268074| 0.8279835| 0.99066466| 0.99334296| 1   | 0.24325744| 1   |

### Table 4. Pearson’s correlation matrix for major element oxides of carbonate-oxide facies BIF.

| SiO₂       | Al₂O₃       | TFe₂O₃      | MnO        | MgO        | CaO         | P₂O₅       | LOI 1000 |
|------------|-------------|-------------|------------|------------|-------------|------------|--------|
| SiO₂       | 1           | -           | -          | -          | -           | -          | -      |
| Al₂O₃      | -0.83856602 | 1           | -          | -          | -           | -          | -      |
| TFe₂O₃     | 0.273337997 | -0.7532961 | 1          | -          | -           | -          | -      |
| MnO        | 0.271478474 | 0.29641895 | -0.85141724| 1          | -           | -          | -      |
| MgO        | -0.43677971 | 0.85635394 | -0.98470727| 0.74702181| 1           | -          | -      |
| CaO        | -0.60562169 | 0.94137984 | -0.93100424| 0.6012302 | 0.98035734 | 1          | -      |
| P₂O₅       | -0.9740111  | 0.69337525 | -0.0484031 | -0.48266298| 0.22167576 | 0.40964402| 1      |
| LOI 1000   | -0.5167765  | 0.89976575 | -0.96478375| 0.68346967 | 0.99585639 | 0.99423114| 0.30943453| 1      |

The oxide facies-BIF samples had SiO₂ = 43.74–68.37 wt.%, which was negatively correlated with MnO, MgO, Al₂O₃, CaO, and P₂O₅. Fe₂O₃ (30.71–55.63 wt.%) was the dominant oxide and was negatively correlated with MnO (r = −0.56), MgO (r = −0.72), and CaO (r = −0.78). Al₂O₃ (0.03–0.36 wt.%) was the second most abundant oxide and was moderately negatively correlated with Fe₂O₃ (r = −0.67) and strongly positively correlated with MnO (r = 0.54), MgO (r = 0.88), and CaO (r = 0.94). MnO content was low (0.01–0.12 wt.%) and had strong positive correlation with MgO (r = 0.87) and CaO (r = 0.75). MgO content was also low (0.02–0.08 wt.%) and had strong positive correlation with CaO (r = 0.97). These oxide facies-BIF samples had very low contents of TiO₂ (<0.01 wt.%) and K₂O (<0.02 wt.%) (Tables 2 and 3).

The carbonate-oxide facies-BIF samples had SiO₂ = 37.23–41.05 wt.% and Fe₂O₃ = 44.86–55.63 wt.%. SiO₂ had weak positive correlation with MnO (r = 0.27) and strong negative correlation with Al₂O₃ (r = −0.84), CaO (r = −0.61), and P₂O₅ (r = −0.97). Fe₂O₃ was strongly negatively correlated with MnO (r = −0.85), MgO (−0.98), and CaO (r = −0.93). The MgO content (0.18–1.99 wt.%) was strongly positively correlated with
CaO ($r = 0.98$), suggesting the incorporation of Mg in carbonate, which was the dominant phase in carbonate-oxide facies. Contents of the remaining major oxides, including TiO$_2$, Al$_2$O$_3$, MnO, CaO, K$_2$O, Na$_2$O, and P$_2$O$_5$, were below 1.00 wt.%. The Al$_2$O$_3$ content (0.06–0.48 wt.%) was strongly positively correlated with MgO ($r = 0.85$) and CaO ($r = 0.94$) (Table 4) but negatively correlated with Fe$_2$O$_3$ ($r = −0.75$). The MnO content (0.15–0.44 wt.%) was positively correlated with MgO ($r = 0.74$) and CaO ($r = 0.60$) but negatively correlated with P$_2$O$_5$ ($r = −0.48$) (Tables 2 and 4).

5.2.2. Trace Elements

Trace element concentrations of the Nabeba BIF varied widely (Table 5). The oxide facies-BIF had lower Zr (0.9–2.1 ppm), Nb (0.1–0.3 ppm), and Zn (3–21 ppm) but higher Cr (15–52 ppm) and V (1–8 ppm) contents than the carbonate-oxide facies-BIF, which had Zr (1.4–7.6 ppm), Nb (0.1–1.1 ppm), Zn (11–175 ppm), Cr (12–50 ppm), and V (5–7 ppm). Transition metals, which showed slight enrichments (relative to HFSEs), are commonly used as indicators of direct volcanogenic hydrothermal input in chemical precipitates [46]. When normalized to the average upper continental crust (UCC) [43], our BIF samples were depleted in most high field strength elements (HFSEs; e.g., Zr, Nb, Th, and REEs) and large ion lithophile elements (LILEs; K and Rb) (Figure 7).

![Figure 7. Upper continental crust (UCC)-normalized trace element patterns of the Nabeba BIF. UCC normalizing values from [47].](image-url)
Table 5. Trace element compositions (in ppm) and ratios of the Nabeba BIF.

| BIF Facies | Oxide | Carbonate-Oxide | Note: (Eu/Eu*)<sub>SN</sub>, (Sm/Yb)<sub>SN</sub>, (La/Yb)<sub>SN</sub> and (Tb/Yb)<sub>SN</sub> are normalized to chondrite [48], while (Co/Ce)*<sub>SN</sub>, (Pr/Pr*)<sub>SN</sub>, (La/Yb)<sub>SN</sub>, (Tb/Yb)<sub>SN</sub>, (Y/Y)*<sub>SN</sub>, (Sm/Yb)<sub>SN</sub>, (Eu/Eu*)<sub>SN</sub> are normalized to PAAS [49]. In the oxide facies-BIF, Nb and Al<sub>2</sub>O<sub>3</sub> showed no or weak positive correlation (r = 0.05) (Figure 8a), while for the carbonate-oxide facies-BIF, the three samples were too few for the correlative analysis and therefore were not plotted in Figure 8. In addition, Zr and Al<sub>2</sub>O<sub>3</sub> were weakly positively correlated in the oxide facies-BIF (r = 0.25) (Figure 8b). |
5.2.3. Rare Earth Elements

The Nabeba BIF had generally low total REE contents ($\Sigma$REE = 8.32–12.11 ppm), with an average of 10.06 ppm for the oxide facies-BIF and 9.81 ppm for the carbonate-oxide facies-BIF samples (Table 5). The low $\Sigma$REE content feature was also reported in many other Archean BIFs [46,50,51]. The PAAS-normalized REE+Y (REY) patterns for the studied BIF were characterized by greater depletions in LREE/HREE ((La/Yb)$_{SN}$ = 0.23 (oxide facies-BIF) and 0.28 (carbonate-oxide facies-BIF) than MREE/HREE ((Tb/Yb)$_{SN}$ = 0.84 (oxide facies-BIF) and 0.81 (carbonate-oxide facies-BIF)) (Figure 9). Strong positive Eu anomalies (Eu/Eu*)$_{SN}$ were present in both the oxide facies-BIF (1.65–2.60) and the carbonate-oxide facies-BIF (1.77–2.32) samples. The samples also had average high (La/La*)$_{SN}$ values, i.e., 1.37 for the oxide facies-BIF and 1.51 for the carbonate-oxide facies-BIF.

Determining Ce anomalies is often complicated due to the anomalous behavior of La. Bau and Dulski [3] suggested the use of Ce/Ce* vs. Pr/Pr* diagram to determine La and Ce anomalies in BIFs. The Ce/Ce* and the Pr/Pr* ratios were calculated as $\text{Ce}_{SN}/(0.5 \text{ La}_{SN} + 0.5 \text{ Pr}_{SN})$ and as $\text{Pr}_{SN}/(0.5 \text{ Ce}_{SN} + 0.5 \text{ Nd}_{SN})$, respectively. The Ce/Ce* vs. Pr/Pr* plot indicated that the majority of our samples had no negative Ce anomalies and had very small to negligible positive La and Ce anomalies (Figure 10).

| Element | Oxide facies-BIF | Carbonate-oxide facies-BIF |
|---------|-----------------|--------------------------|
| Gd      | 0.57            | 0.35                     |
| Tb      | 0.10            | 0.06                     |
| Dy      | 0.61            | 0.36                     |
| Y       | 5.0             | 3.2                      |
| Ho      | 0.14            | 0.08                     |
| Er      | 0.39            | 0.25                     |
| Tm      | 0.06            | 0.04                     |
| Yb      | 0.38            | 0.26                     |
| Lu      | 0.06            | 0.04                     |
| Hf      | <0.2            | <0.2                     |
| Ta      | 0.15            | 0.06                     |
| Pb      | 1.0             | 1.4                      |
| U       | 0.09            | <0.05                    |

Note: (Eu/Eu*)$_{CN}$, (Sm/Yb)$_{CN}$, (La/Yb)$_{CN}$, (Tb/Yb)$_{CN}$ are normalized to chondrite [48], while (Ce/Ce*)$_{SN}$, (Pr/Pr*)$_{SN}$, (La/Yb)$_{SN}$, (Tb/Yb)$_{SN}$, (Y/Y*)$_{SN}$, (Sm/Yb)$_{SN}$, (Eu/Sm)$_{SN}$, and (Eu/Eu*)$_{SN}$ are normalized to PAAS [49].
5.3. LA-ICP-MS Magnetite Chemical Compositions

5.3.1. Magnetite in the Oxide Facies-BIF

Elemental contents of magnetite from the Nabeba oxide facies-BIF varied considerably (Table S1), except for SiO$_2$ (0.07–9.40 wt.%), which was attributed to the presence of quartz near the analysis spot. Contents of other major element oxides (e.g., MgO and MnO) were very low (<0.01 wt.%), while those of most trace elements were either close to the detection limit or showed insignificant concentration change, although Ni (6.37–80.00 ppm), Co (1.06–12.70 ppm), Zn (7.34–77.80 ppm), and Y (0.17–50.80 ppm) displayed high values (Table S1). Magnetite trace element contents of the oxide facies-BIF differed widely, as shown in Mg vs. Mn (r = 0.25), Mn vs. Zn (r = 0.67), Ti vs. V (r = 0.8), Cu vs. Zn (r = 0.47), Co vs. Ni (r = 0.59), and Ti vs. Cr (r = 0.81) diagrams, indicating positive correlations. A few trace elements had narrow concentration ranges (within one order of magnitude), and they had no or weak positive correlations, i.e., Mg vs. Zn (r = 0.08) and Mg vs. Co (r = 0.08) (Figure 11b,c).

The magnetite REY contents of the Nabeba oxide facies-BIF were generally above the detection limits (Table S1). The PAAS-normalized REY patterns were characterized by LREE/HREE depletions (Nd/Yb = 0.012–0.674) and MREE/HREE depletions (Sm/Yb = 0.0282–0.9572) (Figure 12), with the exception of Mag/5–1 (sample CG-C2) that had weak MREE/HREE enrichment (Sm/Yb = 1.279). The PAAS-normalized REY patterns also showed negative to weak positive La anomalies (0.0805–0.9976) and weak (negative or positive) Y (Y/Y* = 0.4656–2.1268) and positive Eu (Eu/Eu* = 1.9963–5.4061) anomalies. Negligible to moderate positive Ce (Ce/Ce* = 0.3676–1.7234) and Gd (Gd/Gd* = 0.3034–3.745) anomalies were also found in our samples. The super-chondritic Y/Ho ratios = 18.219–45.336, with an exceptionally high ratio for Mag/4–1 (Y/Ho = 57.70) and Mag/6–1 (Y/Ho = 91.62).
Figure 11. Binary diagrams of magnetite chemical compositions from the Nabeba BIF: (a) Mn vs. Mg, (b) Co vs. Mg, (c) Zn vs. Mg, (d) Si vs. Mg, (e) Zn vs. Mn, (f) Ni vs. Co, (g) V vs. Cr, and (h) Cr vs. Ti.

Figure 12. PAAS-normalized REY patterns of selected average magnetite from the Nabeba oxide facies-BIF.
5.3.2. Magnetite in Carbonate-Oxide Facies-BIF

Apart from the high major iron oxide content, other major element oxides (e.g., MgO, MnO and TiO$_2$) had concentrations generally below 2 wt.% (Table S2).

Compared to the magnetite samples from the oxide facies BIF, REY contents of most magnetite samples from the carbonate-oxide facies BIF were below the detection limit, which might have been due to poor signals and/or alterations. This explains the very few data points used for the carbonate-oxide facies BIF. The data were not used in the bivariate plots to test for any correlations between analysed elements. Correlations based on bivariate plots for the Nabeba BIF were done using the whole rock geochemical data (Figure 8) and the magnetite LA-ICP-MS data for oxide-facies and carbonate-oxide-facies BIF (Figure 11). The magnetite LA-ICP-MS data for carbonate-oxide facies BIF were used cautiously, with emphasis on (1) the nature of the REY patterns when compared with other well-studied BIFs and the Nabeba oxide facies BIF in this study, and (2) the absolute values (i.e., amounts) of the elemental concentrations and whether they were higher or lower than the concentrations in the oxide facies BIF.’

6. Discussion

6.1. Primary Geochemical Signature of the Nabeba BIF Magnetite

The primary iron minerals (e.g., Fe-oxyhydroxide) in most Archean BIFs were modified by later diagenesis and regional metamorphism [52]. These assertions in which magnetite in BIF were formed by diagenetic and/or metamorphic processes are illustrated by the following three equations:

1. $2\text{Fe(OH)}_3 + \text{Fe}^{2+} \rightarrow \text{Fe}_3\text{O}_4 + 2\text{H}_2\text{O} + 2\text{H}^+$ [26,53];
2. $12\text{Fe(OH)}_3 + \text{CH}_2\text{O} \rightarrow 4\text{Fe}_3\text{O}_4 + \text{CO}_2 + 19\text{H}_2\text{O}$ [54];
3. $\text{Fe(OH)}_2 \cdot 2\text{Fe(OH)}_3 \rightarrow \text{Fe}_3\text{O}_4 + 4\text{H}_2\text{O}$ [32].

Recent studies [3,29,55] showed that the separation of Y and REE in the ambient seawater column represents depletions in LREEs relative to HREEs, in association with Y and La positive anomalies, which is explained by the preferential removal of these elements in organic matter, Fe oxyhydroxides, and clay minerals. Magnetite is the most abundant iron oxide mineral in both facies of the Nabeba BIF. In order to use the magnetite as a proxy for understanding the environment of deposition, it is essential to establish whether the Nabeba BIF magnetite retained the REY signature of the precursor minerals, notably Fe oxyhydroxides [1,4,56–60].

In our study, the magnetite REY patterns in the two-facies BIF samples showed LREE/HREE and MREE/HREE depletions, negative to slightly positive La anomalies, slightly negative to positive Y anomalies and positive Eu anomalies, negligible to weak positive Ce and Gd anomalies, and super-chondritic Y/Ho ratios (Table S1). The Nabeba BIF magnetite showed similar REY characteristics as those in other well-studied Archean BIF magnetites from Isua, North Caribou, Temagami [61], Badampahar [62], and Musselwhite and Meliadine BIFs [30], regardless of their diagenetic, later metamorphic, and hydrothermal histories. The similarity with magnetite data of the other well-studied BIFs indicates that the Nabeba BIF magnetite possesses the primary geochemical signatures that would be useful in determining the depositional environment and the genesis of the Nabeba BIF.

6.2. Characterization of the Nabeba BIF Depositional Environment

6.2.1. Purity of the Nabeba BIF Based on Amount of Detrital Input

The magnetite trace element contents of the Nabeba BIF were generally similar to those of other BIFs (e.g., Meliadine and Musselwhite), except for the high Y-Ni contents in the former (Figure 13a,b). Most of our samples were plotted in the extended BIF field of Gourcerol et al. [30] and overlapped with the extended field of Sun et al. [32] (Figure 14). Since BIFs are formed by chemical precipitation, any detrital input would produce high contents of Al$_2$O$_3$ and TiO$_2$ and correlations between HFSE contents and REE ratios (e.g., La/La*, Y/Ho, Pr/Yb, and Ce/Ce*) [66–68]. As we mentioned previously, the three
samples of carbonate-oxide facies BIF were not good for correlation such as in Figure 15, but we used their element ratios to discuss other aspects of the BIF characteristics. Thus, the variations and the negative correlations of low $\text{Al}_2\text{O}_3$ concentrations with $\text{SiO}_2$ and the weak $\Sigma$REE vs. $\text{Al}_2\text{O}_3$ correlations in the oxide-facies BIF samples (Tables 3 and 4; Figure 15a) suggested insignificant detrital input in the Nabeba BIF formation [5,67,69,70]. However, the strong positive $\Sigma$REE vs. $\text{P}_2\text{O}_5$ correlation in oxide-facies indicated that $\text{P}_2\text{O}_5$ controlled REY contents during the BIF precipitation (Figure 15). Slight negative $\text{Zr}$ vs. $\text{Y}/\text{Ho}$ correlation ($r = -0.11$; Figure 15c) of the oxide-facies BIF further argued against detrital (terrigenous) contamination during the BIF chemical precipitation [71].

Meanwhile, hydrothermal input into the Nabeba chemical sediments was implied from the high average $\text{Si}/\text{Al}$ (oxide facies: 772; carbonate-oxide facies: 326) and $\text{Fe}/\text{Al}$ (oxide facies: 708; carbonate-oxide facies: 474) [14,72,73]. In the $\Sigma$REE vs. ($\text{Co}+\text{Ni}+\text{Cu}$) diagram [5,74], samples of both facies were plotted close to the hydrothermal deposit field (Figure 15d), implying that the main constituents ($\text{Si}$ and $\text{Fe}$) of the Nabeba BIF originated from hydrothermal fluids. The close-superchondritic $\text{Y}/\text{Ho}$ ratio (~35) in the Nabeba BIF samples suggested that this signature was inherited from hydrothermal fluid that had not undergone significant decoupling of $\text{Y}$ from REE [70,75,76]. In Figure 13, all the Nabeba BIF magnetite data from both facies were plotted below the high-T magmatic-hydrothermal field [63] and exhibited moderate Eu anomalies (Figure 16). This suggested that the Nabeba BIF precipitated from medium-temperature fluids or a mixture of high-temperature hydrothermal vent fluids ($T \approx 250^\circ\text{C}$) with cold ambient seawater.

![Figure 13. Bulk continental crust (BCC)-normalized multi-element diagrams for the magnetite layers from the Nabeba BIF in comparison with selected BIFs (from Musselwhite, Meliadine [30], Badampahar [31], and Middleback Ranges (Magnet Iron) [33]) (all in situ analysis; [30,31,33] and high-temperature hydrothermal magnetite [63]): (a) oxide facies-BIF magnetite; (b) carbonate-oxide facies-BIF magnetite. Blue area in both diagrams denotes elemental concentration of different spots in the Nabeba BIF magnetite layers, and the grey area denotes the elemental concentration of high-temperature hydrothermal magnetite [63].](image-url)
Figure 14. (Ti + V) vs. Ni/(Cr + Mn) discrimination diagram [64] for the Nabeba BIF magnetite. Magnetite from different types of iron deposits, including Kiruna apatite-magnetite, iron-oxide-copper-gold (IOCG), porphyry Cu, magmatic Fe-Ti-V oxide, BIF, and Opemiska-type, are also shown for comparison. The extended BIF field represented by a dashed black lines results from Gourcerol et al. [30]. The general BIF field represented by a dashed blue circle is generated from this study following the magnetite concentrations plots of from this study and Middleback Ranges BIFs [33], Meliadine BIFs, Musselwhite BIFs [30], Gongchangling BIFs [32], and Sokoman iron formation [28,65].

Figure 15. Plots of (a) ΣREE vs. Al₂O₃; (b) P₂O₅ vs. ΣREE; (c) Y/Ho vs. Zr; (d) (Co+Ni+Cu) vs. ΣREE (after [5]) for the Nabeba BIF.
Figure 16. PAAS-normalized REY patterns of magnetite and whole-rock BIF (mean values) from Nabeba in relation to the average compositions of hydrothermal fluid [76] and modern seawater [71]. Magnetite geochemical data from Musselwhite, Meliadine [30], Badampahar [31], and Middleback Ranges (Iron Magnet mine) [33] BIFs are also shown for comparison.

6.2.2. Fluid Mixing and T\textsubscript{f}O\textsubscript{2} Conditions of Nabeba BIF Precipitation

The Nabeba BIF magnetite and the whole-rock samples had weak positive Ce anomalies (except for two samples of CG-NB3 and CG-NB6) (Figure 10). Their REY distribution patterns showed LREE depletions, a mild Ce incongruity, and marked positive anomalies for Eu and weakly positive to negligible Y (Figure 16). This indicated that the Nabeba BIF formation was likely influenced by reduced bottom water [77] and the chemical precipitates were derived from a mixture of anoxic seawater and medium-to-high-temperature hydrothermal fluids. Similar geochemical trends were found in the magnetite from many
well-studied Archean BIFs, such as Musselwhite and Meliadine [30], Badampahar [62], and Middle Back Ranges [33]. This implied that, despite the overprinting diagenetic, metamorphic, and hydrothermal processes, the Nabeba magnetite still retained the geochemical signature of the fluid mixing that formed the original Nabeba chemical precipitates on the paleo-seafloor.

To determine the proportion of fluid-mixing components (i.e., hydrothermal vent fluid and seawater) for the Nabeba BIF, variations of Sm/Yb and Eu/Sm ratios [78] were used (Figure 17), which yielded ~0.1% for the high-temperature hydrothermal vent fluid input [2,30]. Strong positive Eu anomalies were associated with high-temperature hydrothermal fluids, medium-to-weak Eu anomalies with low (or medium)-temperature hydrothermal fluids, and no Eu anomaly with seawater (Figure 16; [3,23,71]). Thus, the weak Eu anomalies exhibited by both facies of the Nabeba BIF (Figure 16) was consistent with the fluid mixing that dominantly comprised (~99.9%) ambient seawater with minor (~0.1%) high-temperature vent fluids.

The presence of W, As, Pb, Mo, and Sn in the precipitates from seawater was influenced by hydrothermal fluids associated with felsic rocks. In contrast, hydrothermal fluids related to mafic-ultramafic rocks would produce elevated Ti, V, Ni, Cr, and Mg contents [28,79], which was the case for the Nabeba BIF (Figures 2 and 11) [28,79]. Titanium and V contents in magnetite are sensitive indicators of redox conditions and mafic material input [80,81]. The magnetite precipitated from reduced fluids had lower Ti/V ratios than those of the magnetite precipitated from oxidized fluids [32,80–82]. Overall, the Ti/V ratios from both facies of the Nabeba BIF were considered as low. Therefore, the magnetite trace element compositions of these two facies indicated medium-high temperature and relatively reducing conditions and were likely related to mafic-ultramafic hydrothermal input during the BIF deposition on the paleo-seafloor.

However, when comparing both facies of the Nabeba BIF, magnetite from the carbonate-oxide facies-BIF had higher V content than its oxide facies-BIF counterpart (Figure 11g),

![Figure 17. Sm/Yb vs. Eu/Sm plot (modified after Alexander et al. [78]) for the Nabeba BIF and other BIFs shown in Figure 16. Average compositions of high-temperature (>350 °C) hydrothermal fluids [76], low-temperature hydrothermal fluids [83], and Pacific seawater [84] are shown for comparison.](image-url)
yielding lower magnetite Ti/V ratio for the carbonate-oxide facies (3.72–16.02 ppm, average 10.95 ppm) than that for the oxide facies (18.51–136.69 ppm, average 60.63 ppm). This suggested that the depositional environment was more reducing for the carbonate-oxide facies than for the oxide facies [28,65].

7. Conclusions
1. The Nabeba BIF comprised two facies (oxide and carbonate-oxide) and displays alternating iron-rich and quartz-rich layers. The BIF consisted of magnetite, hematite, and quartz, together with minor siderite and magnesite. Geochemical analysis suggested that the Nabeba BIF had mainly Fe and Si, with minor CO3 and trace detrital material.
2. Magnetite from both BIF facies had a wide range of trace element contents (e.g., Mn, Ti, Ni, Mg, Cr, V, and Zn), which suggested input of medium-high-temperature hydrothermal vent fluids. Redox and temperature conditions likely controlled the magnetite chemical compositions.
3. Major and trace (including REE) element compositions suggested that Fe and Si were sourced from anoxic seawater mixed with medium-high-temperature (>250 °C) hydrothermal fluids. The Nabeba BIF was likely deposited in a reducing marine environment.

Supplementary Materials: The following are available online at https://www.mdpi.com/article/10.3390/min11060579/s1, Table S1: Chemical composition of magnetite from the oxide facies BIF of the Nabeba deposit major element (wt.%) trace elements (ppm), Table S2: Chemical composition of magnetite from the carbonate facies BIF of the Nabeba deposit major element (wt.%) trace elements (ppm).

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