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### **ORIGINAL ARTICLE**

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# **In Situ Fe and S isotope analyses in pyrite from the 3.2 Ga Mendon Formation (Barberton Greenstone Belt, South Africa): Evidence for early microbial iron reduction**

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#### **Abstract**

On the basis of phylogenetic studies and laboratory cultures, it has been proposed that the ability of microbes to metabolize iron has emerged prior to the Archaea/ Bacteria split. However, no unambiguous geochemical data supporting this claim have been put forward in rocks older than 2.7–2.5 giga years (Gyr). In the present work, we report in situ Fe and S isotope composition of pyrite from 3.28- to 3.26-Gyr-old cherts from the upper Mendon Formation, South Africa. We identified three populations of microscopic pyrites showing a wide range of Fe isotope compositions, which cluster around two  $\delta^{56}$ Fe values of −1.8‰ and +1‰. These three pyrite groups can also be distinguished based on the pyrite crystallinity and the S isotope mass-independent signatures. One pyrite group displays poorly crystallized pyrite minerals with positive  $\Delta^{33}$ S values > +3‰, while the other groups display more variable and closer to 0‰  $\Delta^{33}$ S values with recrystallized pyrite rims. It is worth to note that all the pyrite groups display positive  $\Delta^{33}$ S values in the pyrite core and similar trace element compositions.

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We therefore suggest that two of the pyrite groups have experienced late fluid circulations that have led to partial recrystallization and dilution of S isotope massindependent signature but not modification of the Fe isotope record. Considering the mineralogy and geochemistry of the pyrites and associated organic material, we conclude that this iron isotope systematic derives from microbial respiration of iron oxides during early diagenesis. Our data extend the geological record of dissimilatory iron reduction (DIR) back more than 560 million years (Myr) and confirm that microorganisms closely related to the last common ancestor had the ability to reduce Fe(III).

#### **KEYWORDS**

Archean, Barberton Greenstone Belt, dissimilatory Iron reduction, iron and Fe isotopes, Mendon cherts, SIMS

#### **1** | **INTRODUCTION**

The identification of early traces of life in the geological record is challenging due to the scarcity and complex post-depositional history of early Archean rocks. Given their micrometer size and chemical composition (dominated by organic molecules prone to oxidation), the first forms of life, Archaea and Bacteria, did not leave unambiguous fossils (Brasier, Green, Lindsay, & Steele, 2004 cf.; Schopf, Kudryavtsev, Czaja, & Tripathi, 2007). Biominerals associated with residual organic matter are among the best evidence of ancient microbial activity (Brasier et al., 2004; Schopf et al., 2007). Pyrite (FeS<sub>2</sub>) is one of them and can be found in some of the oldest sedimentary rocks on earth. Pyrite has been extensively studied for reconstructing past environmental conditions of the early earth, especially atmospheric composition. Since the canonical work of Farquhar and colleagues and the discovery of mass-independent fractionated sulfur (S-MIF) isotope signatures in Archean pyrite (Farquhar, Bao, & Thiemens, 2000), it is well established that Archean pyrites record atmospheric sulfur cycle. S-MIF signatures are expressed as  $\Delta^{33}$ S, a deviation from a mass-dependent fractionation relationship, which can be expressed as (Farquhar et al., 2000; Ono, Wing, Johnston, Farquhar, & Rumble, 2006) follows:

$$
\Delta^{33}S = \delta^{33}S - \left( \left( 1 + \delta^{34}S / 1000 \right)^{0.5152} - 1 \right) \times 1000
$$

where δ<sup>x</sup>S = ((<sup>x</sup>S/<sup>32</sup>S)<sub>sample</sub>/(<sup>x</sup>S/<sup>32</sup>S)<sub>reference</sub> − 1) × 1,000 (‰) (*x* = 33, and 34) and the reference is Canyon Diablo Troilite (V-CDT, Ding et al., 2001). Photochemical reactions of dissociation of volcanic  $SO<sub>2</sub>$  in an anoxic atmosphere can produce both elemental sulfur and sulfate aerosols with positive and negative  $\Delta^{33}$ S values, respectively. Although alternative views exist (Oduro et al., 2011), it is generally acknowledged that  $O_2$  levels below 10<sup>-5</sup> present atmospheric level (PAL) are critical for the production and preservation of S-MIF in the geological record (Farquhar et al., 2000; Harman, Pavlov, Babikov, & Kasting, 2018; Thiemens & Lin, 2019; Ueno, Ono, Rumble, & Maruyama, 2008; Zhelezinskaia, Kaufman, Farquhar, & Cliff, 2014). Both photochemical experiments and studies of sedimentary sulfides and sulfate have

then confirmed that Archean sulfate has negative  $\Delta^{33}$ S values while elementary sulfur has positive  $\Delta^{33}$ S values (Endo, Danielache, & Ueno, 2019; Farquhar et al., 2000; Farquhar, Wu, Canfield, & Oduro, 2010; Halevy, 2013; Harman et al., 2018; Muller et al., 2017; Ueno et al., 2008; Zhelezinskaia et al., 2014). Sulfur in archean sedimentary pyrite may predominantly originate either from a sulfate-derived sulfide pool (e.g., following microbial sulfate reduction) or from an elemental sulfur derived pool of polysulfides that may further react with iron sulfide precursors (Farquhar et al., 2013). Indeed diagenetic pyrite formation pathways are complex and involve various precursors either molecular (including elemental sulfur and polysulfides) or mineral (including FeS mackinawite and  $Fe<sub>3</sub>S<sub>4</sub>$  greigite; Rickard, 2012). Reconciliating S and Fe isotope compositions of sulfide with their precise diagenetic pathway is still complex as some large isotopic discrepancies between pyrite mineral and the porewater composition have been observed in modern environments (Gomes, Fike, Bergmann, Jones, & Knoll, 2018; Raven, Sessions, Fischer, & Adkins, 2016). However, it is well established that sedimentary pyrite can record past microbial metabolisms (Fike, Bradley, & Rose, 2015). While pyrite may form secondarily during late diagenesis or metamorphism following abiotic pathways, several microbial metabolisms have also been proposed to be involved in the formation of early diagenetic pyrite. These include bacterial sulfate reduction, sulfur reduction and disproportionation, and dissimilatory iron reduction (Rickard, 2012). These various metabolic processes can potentially be traced and distinguished from abiotic routes using iron and sulfur isotope compositions recorded in pyrite (Beard et al., 1999; Canfield, 2001; Johnson, Beard, & Roden, 2008; Johnston, 2011).

For instance, the oldest traces of bacterial sulfate reduction and sulfur disproportionation were evidenced from S isotope composition of pyrite in 3.47-Gyr-old barite deposits from North Pole, Australia (Philippot et al., 2007; Shen, Buick, & Canfield, 2001). Microbial dissimilatory iron reduction (DIR, a form of respiration where ferric iron is used as an electron acceptor) also plays a role in Fe sulfide formation as it promotes the reduction in ferric iron minerals into dissolved ferrous iron, which can then react with dissolved sulfide to precipitate mackinawite (FeS), a key precursor of pyrite (Rickard, 2012). DIR imparts a strong Fe isotope fractionation, enriching the product in the light isotope by ~3‰ (Crosby, Roden, Johnson, & Beard, 2007). This light Fe isotope signature was previously used as a proxy of DIR in Archean rocks as old as 2.7–2.5 Gyr (Archer & Vance, 2006; Czaja et al., 2010; Heimann et al., 2010; Johnson, Ludois, Beard, Beukes, & Heimann, 2013; Nishizawa, Maruyama, Urabe, Takahata, & Sano, 2010) and even as old as 3.8 Gyr (Craddock & Dauphas, 2011; Czaja et al., 2013). However, because an unambiguous interpretation of the 3.8 Gyr signals is hampered by the strong metamorphism experienced by early Archean rocks (Ueno, Yurimoto, Yoshioka, Komiya, & Maruyama, 2002), the Fe isotope signature recorded in 2.7 Gyr sediments can be considered as the earliest firm evidence for DIR (Czaja et al., 2013). Consistently, DIR may have evolved only after the spread of oxygen photosynthesis (Czaja et al., 2013).

In the present work, we analyzed both Fe and S isotope compositions of individual pyrite crystals in 3.28–3.26 Gyr sedimentary rocks from the well-preserved Mendon Formation (South Africa). Together with high-resolution mineralogy and trace element concentrations, the data are used to constrain the mechanisms of pyrite formation and evaluate in particular the impact of diagenesis and metamorphism on pyrite composition.

#### **2** | **SAMPLES AND METHODS**

#### **2.1** | **Geological context and samples description**

The 3.6–3.2 Ga Barberton Greenstone Belt is located in the east of the Kaapvaal Craton, South Africa. It consists of mafic, ultramafic, and felsic volcanic rocks alternating with sedimentary successions, termed the Swaziland Supergroup. Three main lithostratigraphic units compose the Swaziland Supergroup. These are the Onverwacht Group, the Fig Tree Group, and the Moodies Group, in ascending stratigraphic order (Lowe & Byerly, 2007; Viljoen & Viljoen, 1970).

The 3550- to 3260-Myr-old Onverwacht Group (Kröner, Hegner, Wendt, & Byerly, 1996) is composed of six units: Sandspruit, Theespruit, Komati, Hooggenoeg, Kromberg, and Mendon Formations (Lowe & Byerly, 1999). The Sandpruit, Theespruit, Hooggenoeg, and Kromberg Formations consist of basaltic, komatiitic, and felsic volcanic rocks and minor cherts. The Komati and Mendon Formations are composed mainly of komatiites and minor cherts. The central greenstone belt has been metamorphosed in greenschist facies conditions (i.e., ~350°C; Tice, Bostick, & Lowe, 2004). The Fig Tree and Moodies Groups consist of sandstone, shale, chert, banded iron formation, and felsic volcanic rocks, which have been dated at ca. 3,280–3,216 Myr (Byerly, Kröner, Lowe, Todt, & Walsh, 1996; Drabon, Galić, Mason, & Lowe, 2019; Hofmann, 2005; Kamo & Davis, 1994; Kröner et al., 1996; de Ronde, Wit, & Spooner, 1994).

The samples analyzed in this study come from the Barberton Drilling Project (BBDP2) drill core (Figure 1), located at 25°54′24.8″S and 31°03′23.9″E, hole azimuth 288° and dip 50° (Philippot et al., 2009). About 100 m of the uppermost Mendon Formation was recovered. The Mendon Formation was deposited between 3.34 and 3.26 Gyr and has been studied in detail (Busigny et al., 2017; Galić et al., 2016; Hofmann & Bolhar, 2007; Lowe & Byerly, 1999;

Montinaro et al., 2015; Philippot et al., 2009; Philippot, Zuilen, & Rollion-Bard, 2012; Tice et al., 2004; Trower & Lowe, 2016). The sequence consists of five volcanic cycles, termed Mv1 through Mv5, which are each capped by thin chert sequences, termed Mc1 through Mc5 (Byerly et al., 1996). In the BBDP2 drill core, the Mendon Formation is composed of volcanic rocks, mostly komatiite and silicified komatiite of the Mv5 cycle, which are overlain by ~60 m of finely laminated cherts of the Mc5 cycle (Decker, Byerly, Stiegler, Lowe, & Stefurak, 2015). The base of the Mc5 sequence was dated at 3.28 Gyr, and the top is marked by the S2 spherule layer that was deposited at 3.26 Gyr (Decker et al., 2015). Mc5 corresponds to a wide variety of chert, with mostly black chert and ferruginous gray chert, and minor amounts of black and white banded chert, banded ferruginous chert, intraclast breccia and silicified green colored ash and accretionary lapilli (Trower & Lowe, 2016). The Mendon Formation was deposited during a period of local volcanic quiescence with slow sedimentation rates under quiet water settings (Lowe & Byerly,

Twelve samples were selected based on sulfide abundance; their name indicates the location (in meter) along the drill core (Table 1). Most samples correspond to black cherts composed of microcrystalline quartz and minor amounts of organic matter and iron carbonates (Figure 2 and Table 1). The size of disseminated pyrites varies over a large range from 10 μm to 1 cm (Figure 2 and Table 1). The chert samples were affected by late fluid circulation, as illustrated by the presence of numerous quartz, carbonates, and chlorite veins linked to regional metamorphism (Busigny et al., 2017). However, primary sedimentary structures are well-preserved, with fine laminations, especially in samples 114.30 and 114.33, which have experienced only limited fluid circulation overprint (Figure 2 and Table 1). Alteration phases are present as ganterite (barium-rich mica), chlorite, and fuchsite. Minor phases of barite, iron oxide, nickel-rich iron oxide, and rutile are also present. Sulfides occur essentially as pyrite, but one sample (94.92) shows the presence of chalcopyrite and pentlandite. One sample (160.12) represents altered komatiites in which spinifex texture has been preserved, with olivine pseudomorphed by ankerite, dolomite, quartz, fuchsite, ganterite, and small grains of barite (Figure 2). Sample 160.12 displays large (mm) pyrites and smaller (>100 μm) cubic pyrites. Sample 147.38 contains one large (mm) square pyrite with visible mineral (quartz and ankerite) inclusions (Figure 2). Very small pyrites (<10 μm) are present in the chert matrix but were not studied here due to their small size compared with ion microprobe beam size.

#### **2.2** | **Methods**

1999; Trower & Lowe, 2016).

#### **2.2.1** | **SEM and TEM analyses**

Scanning electron microscopy (SEM) and energy-dispersive X-ray spectrometry (EDS) were used to characterize the microstructure and chemistry of pyrite and its inside inclusions for subsequent in situ extraction using focused ion beam milling (FIB).



**FIGURE 1** Geological map of the Barberton Greenstone Belt (from Busigny et al. 2017). The BBDP2 drill core is indicated by the red star and the samples location by yellow stars





*Note:* Qz for quartz, sid for siderite, Ank for ankerite, and C org for organic matter.

SEM observations were performed on a TESCAN VEGAII LSU at IMPMC with 15 kV accelerating voltage and a working distance of 15.4 mm according to the geometrical conditions required for EDS analyses on this microscope. SEM images were collected with both secondary (SE) and backscattered electrons (BSE) detectors. FIB ultrathin sections were extracted from different pyrite grains using an FEI Strata DB 235 at IEMN (Lille, France Schiffbauer &

Xiao, 2009; Wirth, 2009). This extraction procedure maintains textural integrity and prevents shrinkage and deformation of microscale to nanoscale pores, even in the case of highly sensitive materials (Bassim, Scott, & Giannuzzi, 2014).

Transmission electron microscopy (TEM) analyses were performed on FIB sections to characterize crystallographic orientation and textures of the pyrites. TEM observations were performed with



**FIGURE 2** Petrography images of the three groups of pyrites. Images show an optical photograph of the core (a, d, g), an optical photomicrograph of the thin section of each sample (b, e, h), and a SEM picture of the analyzed pyrite (c, f, i). The group A pyrite is exemplified by sample 104.80, the group B by sample 147.38, and the group C by sample 114.30. The core pictures show the importance of fluid circulations identified as quartz veins. We can also note the presence of thin laminations, with a little less fluid circulations in the case of sample 114.30. It is also worth to note that the pyrites occur in proximity to these fluid circulations. All the samples consist of chert with a majority of microquartz; however, samples 104.80 and 147.38 contain also more iron carbonate (ankerite) than sample 114.30. These cherts are rich in organic matter visible in the darkness of the matrix. The pyrites display different shapes but are mostly cubic. The SIMS spots are still visible on the images as well as some leftover of the gold coating. All the pyrites display inclusions of quartz and occasionally of organic matter

JEOL 2100F field emission gun (FEG) microscopes (IMPMC—Paris, France, and IEMN—Lille, France) operating at 200 kV. Scanning transmission electron microscopy (STEM) Z-contrast imaging was performed using the high-angle annular dark field (HAADF) mode. High-resolution TEM (HRTEM) images were collected in the bright field mode. Selected-area diffraction (SAED) patterns were obtained on areas of interest.

#### **2.2.2** | **Trace element composition**

The chemical composition of pyrite from 3 chert samples was investigated by Electron Probe Micro-Analyzer (EPMA) at UNIL. In large pyrite grains, EPMA traverses were performed to quantify the trace element content of pyrite using a 5 spectrometer-equipped JEOL JXA-8530F. The operational conditions were 15 kV acceleration potential **EXAMPLE SERVICE CONTROLLER IN A BUDGE OF A BU** 

difference, 40 nA beam current, and a fully focused beam (<1 µm in diameter). A set of sulfides, silicates, oxides, and native elements, including FeS<sub>2</sub> (for Fe), CuS (for S and Cu), PbS (for Pb), ZnS (for Zn),  $Mn<sub>2</sub>SiO<sub>4</sub>$ (for Mn), NiO (for Ni), and Co, was used as reference materials. All of the used reference materials were tested before the quantitative analysis. Elements and X-ray lines used for analysis were Fe (Kα), Co (Kα), Ni (Kα), Cu (Kα), Zn (Kα), S (Kα), Mn (Kα), and Pb (Mα). Detection limits were between 145 ppm by weight for Co, 160 ppm for Cu, 115 ppm for Mn, 700 ppm for Pb, 130 ppm for Mn, and 200 ppm for Zn on average. For each traverse, analysis spots were placed at 60 µm spacing.

#### **2.2.3** | **Carbon isotope analysis**

Seven rock samples from the Mendon Formation were powdered to <60 μm using a ring and puck mill. Sample powders (about 1 g) were reacted with excess HCl (6 N) at room temperature in glass beakers during one night to remove carbonates. After removing the supernatant, samples were acidified again with HCl (6 N) at 80°C and agitated during 2 hr to guarantee full sample decarbonation. After decantation, the residues were rinsed with deionized distilled water until neutral, centrifuged, and dried at 60°C overnight. TOC content and  $\delta^{13}C_{\text{org}}$  values were measured for carbonate-free residues with a Flash EA1112 elemental analyzer coupled to a Thermo Finnigan DELTA plus XP isotope ratio mass spectrometer interfaced with a ConFlo IV interface at the stable isotope laboratory of the Institut de Physique du Globe de Paris (IPGP). Our results were normalized using three internal standards with 5 different amounts, which were used to estimate the concentration of organic C (wt.%). Reproducibility on  $\delta^{13}C_{\text{ore}}$  and TOC measurements based on at least triplicate measurements of the samples is usually better than ±0.4‰ and  $\pm 0.01$  wt.%, respectively (1 $\sigma$ ).

#### **2.2.4** | **SIMS Fe and S isotopes analyses**

Iron and sulfur isotopes are expressed in delta notation ( $\delta^{56}$ Fe,  $\delta^{34}$ S, and  $\delta^{33}$ S) relative to the international standards IRMM-014 (for Fe) and V-CDT (for S) based on the following equation:

$$
\delta^i X\!=\!\left[\left(^i X/^j X_{\text{sample}}\right) / \left(^i X/^j X_{\text{standard}}\right)\!-\!1\right] \!\times\! 1000
$$

where X is Fe or S, i and j represent the heavy and light isotopes, respectively (54 and 56 for Fe and 34 or 33 and 32 for S). Sulfur mass-independent fractionation has been calculated as the deviation from the Terrestrial Fractionation Line (TFL), using the mass discrimination law (Farquhar et al., 2000), where the factor 0.5152 defines the slope of the TFL.

Iron isotope compositions were measured with a Cameca ims 1270 ion microprobe at CRPG following the procedure described in Marin-Carbonne, Rollion-Bard, and Luais (2011). Briefly, a <sup>16</sup>O<sup>−</sup> primary beam of 10–13 nA intensity was focused to a spot of about 15  $\mu$ m diameter and less than 1  $\mu$ m in depth. The mass resolution

was set to ~7,000, and <sup>54</sup>Fe<sup>+</sup> and <sup>56</sup>Fe<sup>+</sup> were measured in multi-collection mode with two off-axis Faraday cups. Typical <sup>56</sup>Fe intensity was between 5 and 6  $\times$  10<sup>7</sup> counts per second (cps). A typical analysis consisted of 2 min pre-sputtering followed by data acquisition in 30 cycles of 5 s acquisition time. The isobaric interference of <sup>54</sup>Cr on  $54$ Fe was corrected by monitoring chromium on masses 52, but chromium levels were negligible in all samples. The internal precision for  $δ<sup>56</sup>Fe$  values was typically better than ±0.1‰ (2σ), and the external reproducibility based on multiple measurements of our pyrite reference material (Balmat with  $\delta^{56}$ Fe = -0.399‰, (Whitehouse & Fedo, 2007)) was better than ±0.2‰ (2σ).

Sulfur isotope compositions were measured on the Cameca ims 1280 HR2 (CRPG) by simultaneous measurements of <sup>32</sup>S<sup>-</sup>, <sup>33</sup>S<sup>-</sup>, and  $34$ S $^{-}$  in multi-collection mode with three Faraday cups. A Cs $^{+}$  primary beam of 5 nA intensity was focused to a spot of about 15–20 µm in diameter and less than  $1 \mu m$  in depth. The mass resolution was set to 5,000 to resolve the isobaric interferences due to hydrides on sulfur isotopes. Typical  $^{32}$ S $^{-}$  intensity was between 6 and 10  $\times$  10 $^{8}$ counts per second (cps) depending on the sulfide mineral analyzed. A typical analysis consisted of 2 min of pre-sputtering followed by data acquisition in 30 cycles of 3 s each. The background of the detectors was measured during pre-sputtering and was then corrected for each analysis. Several pyrite standards Maine  $\delta^{34}$ S = -20.61‰.  $\delta^{33}S = -10.63\%$ ), Spain ( $\delta^{34}S = -1.56\%$  and  $\delta^{33}S = -0.78\%$ ), and Balmat ( $\delta^{34}$ S = 15.84‰ and  $\delta^{33}$ S = 8.12‰(Muller et al., 2017) were used to determine (a) the instrumental mass fractionation by considering the average value obtained on the three standards and (b) the reference mass discrimination line, from which  $\Delta^{33}$ S values were calculated. The internal precision achieved under these conditions was better than ±0.05‰ for  $\delta^{34}$ S and ±0.03‰ for  $\delta^{33}$ S (2 $\sigma$ ). The external precision was ±0.40‰ (2σ) for  $\delta^{34}$ S and ±0.1‰ (2σ) for  $\Delta^{33}$ S values.

#### **3** | **RESULTS**

#### **3.1** | **SEM and TEM observations**

Ultrathin sections of selected pyrites were characterized by secondary electron microscopy and transmission electron microscopy—in order to discriminate primary features from late modifications related to metamorphism and hydrothermal overprint (Figure 3). On the basis of high-resolution TEM and electron diffraction analyses, two different textures of pyrite were identified as exemplified with samples 114.30 and 147.38. Pyrites from sample 114.30 are composed of nanocrystals with various orientations embedded within a single large pyrite mineral (Figure 3a–c). Nanocrystals of pyrite support a very early diagenetic origin, consistent with the preservation of early diagenetic fine laminations. In contrast, pyrites present in sample 147.38 consist of homogeneous single crystals, typical for a post-diagenetic origin (metamorphic, hydrothermal) and indicating recrystallization features overprinting a primary texture (Figure 3d–f).



**FIGURE 3** Pyrite microphotographs (SEM and TEM) and associated diffraction patterns for samples 114.30 (a, b, c) and 147.38 (d, e, f). SEM analyses (a, d) show quartz inclusions in pyrites, whereas the matrix is composed of microquartz, altered iron carbonate (ankerite), chlorite, and ganterite (Ba-rich mica) recovered by coating. Sample 114.30 contains disoriented nm-scale pyrite particles (see powder diffraction pattern in b and HRTEM image in c) embedded within a pyrite single crystal matrix (see superimposed electron diffraction pattern in b, (<10–2> zone axis) and FFT signal of the matrix in c, displaying the same zone axis). In contrast, for sample 147.38, electron diffraction analysis (e) and HRTEM (f) indicate that pyrite is present as large single crystals (<−113> zone axis). The black bar in (a) and (b) represents the location of the FIB ultrathin section

**TABLE 2** Average trace element concentration (in ppm) of pyrite from Mendon Cherts



*Note:* Detection limits between 145 ppm for Co, 160 ppm for Cu, 115 ppm for Mn, 700 ppm for Pb, 130 ppm for Ni, and 200 ppm for Zn.

Abbreviation: b.d.l., below detection limit.



**TABLE 3** Total organic carbon (TOC, in wt.%) and  $C_{org}$  isotope composition of Mendon cherts  $(δ<sup>13</sup>C$  vs. V-PDB, in ‰)

#### **3.2** | **Trace element composition of the pyrite**

Various trace elements (Co, Ni, Cu, Zn, Pb, and Mn) have been analyzed by EPMA in 3 chert samples and are presented in Table 2. The average concentrations of trace elements are very low (<0.2 wt.%) and often below the detection limit, with for instance no detectable Co, Mn, Zn, and Pb. Each pyrite crystal shows limited internal variability, with no significant difference between rim and core (Table 2).

#### **3.3** | **Carbon isotope composition of Mendon cherts**

Bulk rock samples have organic C concentrations between 0.03 and 1.5 wt.% and  $\delta^{13}$ C values between −32.1 and −26.5‰ (Table 3 and Figure 4) in the range previously determined for Mendon formation (Hofmann & Bolhar, 2007; Walsh & Lowe, 1999). Data from Noel master's thesis (Noel, 2009) are also reported and give similar results (Figure 4). Samples with the highest organic carbon contents show the lowest  $\delta^{13}$ C values, as frequently observed in

#### **3.4** | **Iron and sulfur isotope compositions of pyrite**

In the 12 samples investigated, a total of 109 pyrites were analyzed for both S and Fe isotope compositions, with multiple analyses of the largest pyrite crystals (Table 4). Pyrites in the Mendon Formation show large isotope variability in  $\delta^{56}$ Fe,  $\delta^{34}$ S, and  $\Delta^{33}$ S, which is not related to the stratigraphic position within the core (Figure 5).  $\delta^{34}S$ and  $\Delta^{33}$ S values range from -2.66 to +6.22‰ and -0.39 to +4.25‰, respectively (Figures 5 and 6a), consistent with previous reports on the same formation (Galić et al., 2016; Montinaro et al., 2015; Roerdink, Mason, Whitehouse, & Reimer, 2013). The  $\delta^{56}$ Fe values vary from −4.02 to +2.54‰, covering almost the total terrestrial range reported so far (Dauphas, John, & Rouxel, 2017). Some samples show large  $\delta^{56}$ Fe variations up to 6‰ (Sample 160.12) with small S isotopic variability, while others have large range of  $\delta^{34}$ S values (samples 95.41 and 147.38) with smaller range of variations in  $\delta^{56}$ Fe (Figures 5 and 6). There is no correlation between  $\delta^{34}$ S and  $\delta^{56}$ Fe values (Figure 6b).

In order to better illustrate the isotopic variability of each sample, a probability density distribution was calculated for each sample. Each measurement (δ<sup>56</sup>Fe, σ) is replaced by a Gaussian curve  $f_{\delta^{56}Fe}$  such as:

$$
f_{\delta^{56}Fe,\sigma}(x) = \frac{1}{\sigma \times \sqrt{2\pi}} \times \exp\left[\frac{-\left(-x - \delta^{56}Fe\right)^2}{2\sigma^2}\right]
$$
(1)

The integral of  $f_{\delta^{56}Fe,\sigma}$  between  $x_1$  and  $x_2$  gives the probability that the true  $\delta^{56}$ Fe of the sample is between  $x_1$  and  $x_2$ . The probability density function  $F(x)$  describing  $\delta^{56}$ Fe variations in a sample is defined for *N* measurements as the sum of individual  $f_{\delta^{56}Fe,\sigma}$  divided by *N*:



**FIGURE 4** TOC and  $\delta^{13}$ C values of the different pyrite groups. Data from the same formation published by Hofmann & Bolhar, 2007 and master's thesis from Vincent Noel are also reported in white and black, respectively

$$
F(x) = \sum_{j=1}^{j=N} \frac{f_{\delta^{56}Fe_j \sigma_j}(x)}{N}
$$
 (2)

with the probability  $P(x) = \int_{0}^{+\infty} F(x) \times dx = 1$ . −∞

For example, if  $\int_{0}^{-2.5} F(x) \times dx = 0.4$ , then the probability for a nanopyrite showing δ<sup>56</sup>Fe values lying between −2.5‰ and −2.0‰ is 40%.

## **4** | **DISCUSSION**

Pyrites from the Mendon cherts display a large range of isotopic compositions both in Fe and S, which is not correlated with the size or shape of pyrites. The sample stratigraphic position does not correlate with Fe and/or S isotope compositions (Figure 5). However,  $\Delta^{33}$ S correlates with crystallinity order, with values around 0‰ for well-crystallized pyrites and values higher than 2‰ for polycrystal-hosting pyrites (Figure 7). Three different pyrite populations can be distinguished based on their  $\delta^{56}$ Fe and  $\Delta^{33}$ S values (Figure 7a). These are Group A with mean  $\delta^{56}$ Fe of -1.76 ± 0.6‰ (1 $\sigma$ ) and  $\Delta^{33}$ S of -0.05 ± 0.3‰ (1σ; samples 137.84, 104.8, and 104.83 and few pyrites from 160.12), Group B with  $\delta^{56}$ Fe of +1.05 ± 0.4‰ (1 $\sigma$ ) and  $\Delta^{33}$ S of +0.67 ± 0.5‰ (1 $\sigma$ ; samples 94.92, 95.41, 133.55, 139.80, 147.38, and 160.12), and Group C with  $\delta^{56}$ Fe averaging +1.01 ± 0.9‰ and  $\Delta^{33}$ S of +3.5 ± 0.6‰ (1 $\sigma$ ; samples 114.3 and 114.33). It is worth noting that some samples (e.g., 160.12) contain pyrite grains that belong to two different groups. The following discussion will focus on constraining the origin of the large isotopic variability in both Fe and S compositions by assessing first, the influence of fluid circulation and hydrothermal overprint, second, the different pyrite group origin and pyrite formation pathways, and lastly, the potential microbial origin of these isotope compositions.

### **4.1** | **Role of hydrothermal processes on Fe and S isotopic compositions**

All pyrites from Mendon cherts contain mineral inclusions of either quartz or carbonate and can thus be considered as a late phase related to diagenesis. Post-diagenetic and hydrothermal processes should lead to pyrite recrystallization with variable trace element enrichments such as Co, Ni, and Pb, depending on the fluid chemistry (Large, Maslennikov, Robert, Danyushevsky, & Chang, 2007). In Mendon samples, trace element concentrations of the different pyrite groups are similar (Table 2) and display no detectable enrichments in Co, Mn, Zn, and Pb above the detection limits of EPMA technique (>100 ppm, see section 2.2.2). Chemical profiles obtained across large pyrite grains show homogenous and low concentration in terms of trace element distribution patterns (Table 2). The absence of trace element enrichment argues against a late stage of pyrite overgrowth, which contrasts with SEM and TEM observations. Among the few cases where Co was detected, Co/Ni ratio was always below 0.6, supporting a sedimentary rather than a metamorphic origin for these pyrites (Gregory et al., 2015). It is worth emphasizing that Group B

**TABLE 4** S and Fe isotope composition of pyrite from the Mendon cherts







# **TABLE 4** (Continued)



#### **TABLE 4** (Continued)

L



#### **TABLE 4** (Continued)



pyrite can contain up to 1,700 ppm Ni, but that this enrichment is homogenously distributed throughout the pyrite grains. Group C pyrites show the lowest trace element concentrations, with values typically under the detection limits. Therefore even if the Mendon cherts have experienced hydrothermal fluid circulations (Busigny et al., 2017), these late events have not significantly modified the original pyrite trace element concentrations.

The effect of fluid circulation on iron isotope compositions of Mendon pyrite can also be evaluated. For an equilibration temperature of ~350°C (Tice et al., 2004), pyrite precipitated from a Fe(II)-bearing fluid should have been in, or near to, isotopic equilibrium. Assuming a hydrothermal fluid with  $\delta^{56}$ Fe values between −0.5 and 0‰ (e.g., Severmann et al., 2004; Rouxel, Sholkovitz, Charette, & Edwards, 2008; Johnson et al., 2008) and a fractionation of ~1‰ between pyrite and Fe(II) dissolved in the fluid (Polyakov et al., 2019; Syverson, Borrok, & Seyfried Jr, 2013), the  $\delta^{56}$ Fe values of pyrite precipitated from this fluid should range between +0.5 and +1‰. These values are compatible with the pyrites from Groups B and C (Figure 6), but not with those of Group A. Additionally, pyrite formed under hydrothermal conditions should be well-crystallized, which is not the case of the Group C pyrites. We therefore conclude that a secondary hydrothermal alteration process could account for the Fe isotope compositions of Group B pyrites but not for those of Groups A and C pyrites. We note that comparable Fe isotope profiles obtained in Groups B and C pyrites (Figure 8) supports a related formation process.

Similarly, the influence of fluid circulations on sulfur isotope compositions can be examined. Sulfur in late fluid circulations can have a juvenile origin characterized by a  $\delta^{34}$ S of 0‰ and no S-MIF signatures with  $\Delta^{33}$ S = 0‰ (Labidi, Cartigny, & Jackson, 2015). Sulfur isotope profiles through large pyrites (more than 10 isotopic profiles measured in different pyrite grains) from each group highlight the presence of positive  $\Delta^{33}$ S values in all the pyrite groups (Figure 8). Group C pyrite displays homogenous distributed  $\Delta^{33}$ S throughout the grains, while Group B and Group A show a zoning pattern with non-zero  $Δ^{33}S$  cores and  $Δ^{33}S \sim 0$ % rims (Figure 8). Pyrites from sample 147.38 show large isotopic variations in  $\delta^{56}$ Fe (~2‰) and  $\delta^{34}$ S values (~6‰), with high  $\delta^{56}$ Fe and low  $\delta^{34}$ S cores, and low  $\delta^{56}$ Fe and high  $\delta^{34}$ S rims. These isotopic profiles are consistent with a late stage of pyrite overgrowth associated with the infiltration of a fluid phase of different S composition as already suggested by Marin-Carbonne et al. (2014). Note, however, that some Group C pyrites (sample 114.30), displaying the same range of ~2‰  $\delta^{56}$ Fe variation from core to rim, show homogeneous  $\Delta^{33}$ S and  $\delta^{34}$ S profiles, thus arguing for different fluid sources during rim crystallization (Figure 8). The presence of S-MIF in each pyrite group indicates that the sulfur source is of atmospheric origin. The absence of a  $\Delta^{33}$ S anomaly in the rim of pyrite grains from Groups B and C indicates that a mantle-derived fluid source ( $\Delta^{33}$ S = 0‰) contributed to some of the pyrite overgrowth. This observation contrasts with the absence of core–rim variations recorded from trace element concentrations and Fe isotope compositions, which supports the conclusions



**FIGURE 5**  $\delta^{56}$ Fe,  $\delta^{34}$ S, and  $\Delta^{33}$ S isotope profiles along the BBDP2 core. There is no relationship between Fe and S isotope compositions and the sample depth. Some samples display homogeneous isotope composition, while others (e.g., 104.80 or 114.30) show large isotopic variability. Some single grains show large internal variability, up to 4‰ for  $\delta^{34}$ S. Most samples have  $\Delta^{33}$ S values close to 0‰, while only 5 samples have  $\Delta^{33}$ S values > 0‰ (94.92, 104.80, 114.30, 147.38, and 160.12)

already reached by Marin-Carbonne et al. (2014) on pyrite and Li, Konhauser, Kappler, and Hao (2013) on magnetite that there can be a decoupling between different geochemical tracers during mineral recrystallization.

Altogether, the trace element concentrations,  $\delta^{56}$ Fe and  $\Delta^{33}$ S isotope variabilities at a micrometer scale, pinpoint a sedimentary origin of these pyrites or their precursors.

#### **4.2** | **Pyrite origin and formation pathways**

Pyrite formation is a complex two-step process involving slow pyrite nucleation and fast crystal growth (Rickard, 2012). Pyrite can be formed from various precursors either iron monosulfide like mackinawite (FeS) or polysulfide like greigite (Fe<sub>3</sub>S<sub>4</sub>) (Rickard, 2012). The precursor minerals always dissolve to form aqueous FeS complexes, which then react with  $H_2S$  or polysulfides to form pyrite (Rickard, 2012).

The presence of positive  $\Delta^{33}$ S values in all pyrite groups suggests that pyrite precipitated from elemental sulfur. Elemental sulfur with positive  $\Delta^{33}$ S values was likely produced via photochemical reactions in oxygen-free Archean atmosphere (e.g., Farquhar et al., 2000) and was preferentially delivered to the sediments rather than processed in the water column, as insoluble and unreactive particles (Rickard, 2012). However, pyrite cannot form directly from particulate elemental S (Rickard & Luther, 2007). Before reacting with a dissolved pyrite precursor, an intermediate step to break  $S_8$  rings is required (Rickard, 2012). In sedimentary pore waters,  $S_8$  rings were possibly gradually opened and sulfur chains and compounds were biologically converted to  $H<sub>2</sub>S$ , for example, through disproportionation. The pyrites from Mendon Formation likely formed by late recrystallization of pyrite precursors. As discussed above, this recrystallization process modified the S isotope composition but not the Fe isotope composition. As the formation of pyrite after mackinawite only requires sulfur addition, the lack of Fe isotope composition changes suggests that mackinawite rather than greigite was the precursor phase.

How confidently pyrite  $\delta^{56}$ Fe values reflect the isotopic composition of the parent Fe(II) remains subject of debates (Beard et al.,

In addition, the marked change from positive to negative  $\delta^{56}$ Fe values across a single pyrite grain from Group B and C (Figure 8)



A recent study of Mansor and Fantle (2019) proposed that pyrite  $\delta^{56}$ Fe value can be controlled by its precipitation rate. Negative  $\delta^{56}$ Fe values would reflect the expression of kinetic isotopic effect (KIE), while positive  $\delta^{56}$ Fe values would correspond to equilibrium isotopic effect (EIE). Taken at face value, this would imply that the bimodal distribution of  $\delta^{56}$ Fe values reflects different rates of pyrite precipitation controlled either by kinetic or equilibrium fractionation. If correct, some correlation between pyrite morphology, size, and iron isotope composition would be expected. As suggested by Mansor and Fantle (2019), precipitation rate will be different for large cubic pyrite grain than for small pyrite aggregates. However, large pyrite grains show either negative (sample 104.83) or positive  $\delta^{56}$ Fe values (sample 114.30), and there is no apparent correlation between the size and Fe isotope composition of the pyrites. For example, sample 133.55 displays a large range of pyrite sizes, from 10 μm to 1 mm, while the Fe isotope composition of these pyrites is quite homogeneous (mean  $\delta^{56}$ Fe of +1.12 ± 0.28% for 24 pyrite grains with variable size).



**FIGURE 6** (a)  $\delta^{34}$ S and  $\Delta^{33}$ S compositions and (b)  $\delta^{34}$ S and  $\delta^{56}$ Fe values of the Mendon pyrites. Error bars are comprised in the dot

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would imply that the rate of pyrite precipitation changed drastically during pyrite formation. The model of Mansor and Fantle (2019) predicts that pyrite nucleation and early stages of crystal growth should be associated with high rate of precipitation (*i.e.,* kinetically dominated and low  $\delta^{56}$ Fe values), while later stages should evolve toward lower rates of precipitation (closer to equilibrium, with higher  $\delta^{56}$ Fe values). This prediction is opposite to our Fe isotope profiles in Groups B and C pyrite and is not clearly observed in Group A (Figure 8). This prediction is also not consistent with textural observations, as all pyrite cores from the different groups are polycrystalline and should therefore reflect a same precipitation rate, but group A displays negative  $\delta^{56}$ Fe values, while Group B and C pyrites display positive  $\delta^{56}$ Fe values. Additionally, if Fe isotope compositions were influenced by the rate of pyrite precipitation and kinetic effect, this should have also influenced the S isotopic compositions as suggested by modeling studies (Hegyi & Halevy, 2019; Johnston, Hemingway, Gill, & Halevy, 2019; Raiswell, Whaler, Dean, Coleman, & Briggs, 1993). Pyrite rims and cores should display different  $\delta^{34}$ S values, which has been recorded in only one sample from Group B (147.38) but not in Groups A and C pyrites and other samples from Group B (95.41, Figure 8). With the exception of one sample from Group B, there is no marked variation of  $\delta^{34}$ S values between pyrite cores and rims.

The large range of Fe and S isotope compositions recorded in Mendon pyrites can be explained by various pyrite precursors. In this case, the bimodal distribution of  $\delta^{56}$ Fe values (Figure 7) would reflect two different pyrite precursors, one with negative  $\delta^{56}$ Fe and positive  $\Delta^{33}$ S values and one with positive  $\delta^{56}$ Fe and  $\Delta^{33}$ S values. The negative  $\delta^{56}$ Fe origin of the pyrite precursor may be explained by a kinetic process (see below), while the positive  $\delta^{56}$ Fe values of the precursor would have originated from iron oxides reduction either abiotically or biologically induced by DIR (Yoshiya et al., 2015). How concomitant these pyrite precursors would have been formed is still unknown. In such a scenario, different pyrite groups would reflect different pyrite generations. The similarity in microscale mineralogy and trace element contents is difficult to reconcile with

different pyrite precursors but most likely reflects diagenetic processes. Accordingly, we suggest the Mendon pyrites were most likely formed through the transformation of the precursor mackinawite, which would be best represented by the fine-grained and polycrystalline nature (Rickard, 2012) of some pyrite cores (Group C).

#### **4.3** | **Biosignature of dissimilatory iron reduction recorded in pyrites**

Groups A and B display similar  $\Delta^{33}$ S but distinct  $\delta^{56}$ Fe values (Figures 7 and 8). The negative  $\delta^{56}$ Fe values of Group A are comparable to most Archean sedimentary pyrites (Planavsky et al., 2012), which are usually interpreted as reflecting DIR (Johnson et al., 2008), kinetic isotope fractionation associated with pyrite precipitation (Guilbaud et al., 2011; Rolison et al., 2018), and/or a distillation effect of oceanic dissolved Fe(II) due to Fe oxide precipitation (Rouxel, Bekker, & Edwards, 2005). In contrast, high  $\delta^{56}$ Fe values of Group B pyrites are more occasional for Archean pyrites (Johnson et al., 2008; Rouxel et al., 2005), but rather typical of iron oxides formed from partial oxidation of a large Fe(II) reservoir (Planavsky et al., 2012; Rouxel et al., 2005). Interestingly, the samples hosting Group B pyrites do not contain iron oxide contrary to samples from Group A (Figure 9). This suggests that Group B pyrites with positive  $\delta^{56}$ Fe values were derived from total reduction in former iron oxides, thus preserving their original Fe isotope compositions (near + 1‰). In contrast, Group A pyrites with negative  $\delta^{56}$ Fe values could have formed from dissolved Fe(II) released by partial reduction in iron oxides and, therefore, could record the composition of dissolved Fe(II). Although—as discussed above—a kinetic isotope fractionation associated with pyrite precipitation cannot be ruled out, the absence (in Group B) or presence (in Group A) of residual Fe oxides is more supportive of a prominent effect of Fe reduction. Following this scenario, the difference in Fe isotope composition between Group A and B pyrites could reflect the fractionation associated with iron oxide reduction, which can be estimated at about −2.8‰ (Figure 7). This Fe isotope fractionation is remarkably similar to the one determined experimentally



**FIGURE 7** (a) δ<sup>56</sup>Fe and Δ<sup>33</sup>S values of the Mendon pyrites display three different groups (A in blue, B in green, and C in orange/yellow). There is no correlation with the stratigraphic location (Figure 5). Error bars are comprised within the dots. The mean value of each group is reported as a square with the respective color of each pyrite group. (b) Probability density function of  $\delta^{56}$ Fe values for all pyrites showing bimodal distribution with a difference of ~2.8‰ interpreted as reflecting isotope fractionation associated with microbial Fe respiration



**FIGURE 8** SEM pictures, δ<sup>56</sup>Fe, δ<sup>34</sup>S, Δ<sup>33</sup>S isotopic profiles in individual pyrite grain from each pyrite groups. Errors bars are comprised in the dot



**FIGURE 9** SEM picture of iron oxides present in Group A pyrites. These iron oxides, potentially hematite, are embedded in the microquartz matrix and often associated with iron carbonates

for microbial DIR (Crosby et al., 2007; Johnson & Beard, 2005), but also with abiotic reduction at low temperature (e.g., Frierdich et al., 2019). However, this isotopic fractionation is not compatible with abiotic reduction related to burial and/or interaction with metamorphic fluids. A reduction associated with hydrothermal overprints such as those experienced by Mendon cherts is unlikely because reequilibration at about 350°C should have reduced, not increased the Fe isotope fractionation (Dauphas et al., 2017). Therefore, the large and consistent Fe isotope fractionation can suggest that pyrite  $\delta^{56}$ Fe values were inherited from early diagenetic microbial reduction and remained unaffected by subsequent hydrothermal alteration processes (Figure 10). Groups A and B pyrites were thus likely formed by a DIR process through partial and total Fe reduction, respectively. In contrast, the well-ordered crystallinity and  $\Delta^{33}$ S values near 0‰ of Groups A and B pyrites point to sulfide recrystallization in the presence of S-rich mantle-derived fluids (Roerdink, Mason, Whitehouse, & Brouwer, 2016). The modification of the primary S isotope signature but preservation of the original Fe isotope imprint possibly arose from interaction of a precursor sulfide (e.g., most likely mackinawite) with a S-rich and Fe-poor hydrothermal fluid. Considering the mineralogy of the rocks and the diagenetic and metamorphic processes that are known to have affected the Barberton area, production and circulation of fluids enriched in sulfides must have been common, which could account for the precipitation of pyrite with no S-MIF signatures (Agangi, Hofmann, Eickmann, & Marin-Carbonne, 2019; Agangi, Hofmann, Eickmann, Marin-Carbonne, & Reddy, 2016).

Pyrites from Group C show more variable but mostly positive  $δ<sup>56</sup>Fe$  values, averaging +1.01 ± 0.90‰ (1σ). The positive  $δ<sup>56</sup>Fe$ values together with the absence of iron oxides in the samples suggest pyrite formation after total iron oxide reduction, like for Group B pyrites. Positive  $\Delta^{33}$ S values of Group C pyrites argue for a volcanic source of sulfur equilibrated in an anoxic atmosphere (Johnston, 2011). Photochemical reactions of volcanic  $SO<sub>2</sub>$  in the Archean atmosphere are generally supposed to produce elemental sulfur (S<sup>0</sup>) and sulfate (SO $_4^2$ ) aerosols, with positive and negative  $\Delta^{33}$ S values, respectively (Farquhar et al., 2000). Accordingly, our results suggest that pyrites from Group C derived, at least partly, from Archean atmospheric elemental sulfur. Pyrite  $\delta^{34}$ S values are



**FIGURE 10** Biogeochemical model of the formation of pyrite from 3.28- to 3.26-Gyr-old cherts of the Mendon Formation. The model illustrates the environmental Fe and S compositions in the atmosphere, ocean, and sediment, as well as the various processes leading to the formation of pyrite. The three different groups of pyrites are shown in red. The DIR process took place in the sediment and was associated with sulfur disproportionation. A late metasomatic event modified S but not Fe isotope composition, implying interaction with an Fe-poor fluid

also consistent with microbial disproportionation of elemental sulfur (Figure 5, Johnston, 2011), as most disproportionating organisms produce fractionations between  $\mathsf{S}^\mathsf{0}$  and  $\mathsf{H}_2\mathsf{S}$  smaller than 8‰ (Canfield, 2001). TEM analyses of pyrites from Group C reveal the persistence of polycrystalline particles (Figure 3b,c). These observations, as well as the finely laminated texture of the cherts hosting these pyrites, suggest a sedimentary origin, with better preservation than those of Groups A and B. Moreover, carbon isotope composition of organic matter hosted by Mendon black cherts ranges between −26 and −32.1‰ (Figure 4) and is consistent with a biological origin from phototrophic organisms (Schidlowski, Hayes, & Kaplan, 1983; Zerkle, House, & Brantley, 2005). Previous studies have highlighted the presence of micro-organisms during the deposition of cherts in the Onverwacht Group (Hofmann & Bolhar, 2007; Tice & Lowe, 2006; Walsh, 1992; Walsh & Lowe, 1999) and have excluded an hydrothermal origin of the organic matter (Hofmann & Bolhar, 2007; van Zuilen, Chaussidon, Rollion-Bard, & Marty, 2007). The presence of this fossil biomass supports the possibility for DIR activity, as this metabolic pathway involves iron-based oxidation of organic matter. It is worth noting that samples hosting Group A pyrites, which have experienced partial Fe reduction, have similar organic carbon contents and  $\delta^{13}$ C values to those of Group C, which

corresponds to total Fe oxide reduction ( $C_{\text{ore}}$  ~0.82 to 1.51 wt.%,  $\delta^{13}$ C ~−31.38 to −32.30‰; Figure 4). This indicates that organic matter availability was not the main limiting factor controlling partial versus total Fe reduction. Samples hosting pyrites from Group B have lower organic carbon content ( $C_{\text{ore}}$  ~0.03 to 0.38 wt.%) and slightly less negative  $\delta^{13}$ C values ( $\delta^{13}$ C ~−26.57 to −30.18‰), showing that in this case the total reduction in iron has almost entirely consumed the organic matter (Figure 10).

#### **5** | **CONCLUSIONS**

Taken together, Fe isotope data of Mendon pyrites from Groups A, B, and C show a bimodal distribution, demonstrating a large Fe isotope fractionation process (~2.8‰, Figure 7b) that can be ascribed to DIR. Anaerobic microbial respiration can simultaneously account for (a) the measured  $\delta^{56}$ Fe systematic and variability and (b) the production of the reduced sulfur species necessary for pyrite formation. In addition, the light  $\delta^{13}$ C values recorded in associated organic matter likely reflect the residual biomass derived from phototrophic organisms and consumed through joint Fe and S metabolisms during diagenesis. In contrast, the mineral assemblages and associated isotopic signals are difficult to explain by the abiotic processes discussed above. Our data extend reliable evidence of DIR metabolism back more than ~500 Myr than previously accepted (2.7 and 2.5 Gyr, Archer & Vance, 2006; Craddock & Dauphas, 2011). This supports indirect evidence from phylogenetic analyses and microbial culture studies for DIR origin in the early Archean, and therefore a close relation to the last universal common ancestor (Lonergan et al., 1996; Lovley, 2002; Vargas, Kashefi, Blunt-Harris, & Lovley, 1998). Interestingly, some DIR bacteria are known to reduce elemental sulfur enzymatically through disproportionation reactions producing both sulfide and sulfate (Flynn, O'Loughlin, Mishra, DiChristina, & Kemner, 2014; Thamdrup, Finster, Hansen, & Bak, 1993). This metabolic process could explain the formation of sulfide even in a very low sulfate environment (Berg et al., 2019; Crowe et al., 2014), but also the occurrence of pyrite displaying S isotopic compositions inherited from elemental sulfur in the Archean geological record. The presence of DIR in the Mendon Formation predating the deposition of banded iron formations (BIF) in the overlying Fig Tree Formation is also significant, as this metabolism is considered to be essential in the formation of BIF (Konhauser et al., 2017; Li et al., 2013). Indeed, DIR could explain the low content of organic carbon, the negative carbon isotope values of iron carbonates (Craddock & Dauphas, 2011), as well as the presence of other Fe(II)-bearing minerals (Li et al., 2013) in BIF. Combined with the positive  $\Delta^{33}$ S values recorded in Mendon pyrites, this may indicate the antiquity of coupled microbial iron and sulfur metabolic pathways.

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