Coupled Fe and S isotope variations in pyrite nodules from Archean shale

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1. Introduction

Variations in Fe and S isotope composition of sedimentary pyrites have placed important constraints on the chemistry and redox evolution of the Earth's ocean and atmosphere over geological time (e.g., Bekker et al., 2004; Farquhar et al., 2000; Johnson et al., 2008; Rouxel et al., 2003, 2005; Strauss, 2003). These variations record isotope fractionations during redox reactions, which in some cases might have been biologically mediated (Archer and Vance, 2006; Beard et al., 1999; Johnson et al., 2008). Sulfur isotopes have been used to document ancient microbial metabolisms, because the fractionations produced by living organisms can be large and reflect specific metabolic activity (e.g., Johnston, 2011). Indeed, in the process called Bacterial Sulfate Reduction (BSR), dissolved sulfate is used by eukaryotes, bacteria, and certain groups of Archea as an electron acceptor during organic C remineralization or H2 oxidation. In other redox reactions, hydrogen sulfide can act as an electron donor associated with O2, NO3 or CO2 reduction (Canfield, 2001). BSR preferentially metabolizes 32S relative to 34S, thereby producing fractionation of the S isotopes up to 70‰ (Canfield, 2001; Sim et al., 2011).

The discovery of mass-independent fractionation (MIF) of S isotopes in Archean sedimentary sulfides and sulfates (Farquhar et al., 2000) has deeply modified our understanding of the Precambrian sulfur cycle. The prevailing hypothesis to explain S-MIF is based on experimental studies and atmospheric models that invoke...
photochemical reactions, and suggest an absence of atmospheric oxygen before 2.4 Ga (Farquhar et al., 2000; Ono et al., 2003; Pavlov and Kasting, 2002). Farquhar et al. (2001) suggested that Archean S-MIF was created via photolysis of \( \text{SO}_2 \) and/or \( \text{SO} \) by short ultraviolet radiation (< 220 nm) that penetrated deeply into the Archean atmosphere due to the lack of \( \text{O}_2 \). \( \text{SO}_2 \) photodissociation in an oxygen-free atmosphere produces water-soluble \( \text{SO}_4^{2-} \) with negative \( \Delta^{34}\text{S} \) values and elemental sulfur aerosols, mostly \( \text{S}_8 \), with positive \( \Delta^{33}\text{S} \) values (Farquhar et al., 2000). Although alternative views are still debated (see for alternative view Oduro et al., 2011; e.g., Ohmoto et al., 2006), \( \text{O}_2 \) level below \( 10^{-5} \) Present Atmospheric Level (PAL) is considered critical for the production of mass-independent fractionation in \( \text{S} \) isotopes and its preservation in the sedimentary rock record (Thiemens, 2001). Based on photochemical experiments, it was proposed that Archean seawater sulfate had negative \( \Delta^{34}\text{S} \) values (Farquhar et al., 2000; Ono et al., 2009, 2003), whereas Archean disseminated pyrites have mostly positive \( \Delta^{33}\text{S} \) values (Farquhar and Wing, 2003). This is consistent with \( \text{S} \) isotope composition of hydrothermal barite and sulfide in base-metal barren, distal exhalite deposits (e.g., Farquhar and Wing, 2003), which derived their \( \text{S} \) from seawater sulfate in distal, hydrothermally-influenced low-energy environments.

The \( \text{Fe} \) isotope composition of sedimentary pyrite is highly sensitive to the size of dissolved \( \text{Fe}(II) \) and \( \text{S} \) reservoirs and hence can place important constrains on the redox state and chemistry of Precambrian oceans (Guilbaud et al., 2011; Rouxel et al., 2005). Iron isotopes fractionate through both redox and non-redox reactions (e.g., Johnson and Beard, 2005). Hence interpretation of iron isotope record of \( \text{Fe}(II) \)-bearing marine deposits requires an understanding of \( \text{Fe} \) sources and formation mechanisms of iron-bearing minerals, including oxides, sulfides, carbonates, and silicates, in marine sediments. Each of these minerals can have various origins, such as detrital, biochemical and hydrothermal, and thus can record different \( \text{Fe} \) isotope fractionations (Heimann et al., 2010; Johnson et al., 2008; Planavsky et al., 2009; Rouxel et al., 2005).

By coupling the \( \text{S} \) and \( \text{Fe} \) isotope systems, it is possible to gain additional insights into the processes resulting in the formation of pyrite (Archer and Vance, 2006; Fabre et al., 2011; Hofmann et al., 2009; Rouxel et al., 2008). For example, it has been proposed that coupled \( \text{Fe} \) and \( \text{S} \) isotope data can be used as a proxy for microbial \( \text{Fe}(III) \) and sulfate reduction, especially for Archean sediments (Archer and Vance, 2006). Studies of Archean rocks frequently use a bulk rock approach, although a growing number of studies focuses on individual crystals or crystal aggregates. A recent SIMS study of \( \text{S} \) isotopes of various Archean pyrites has shown large intra-grain variability in \( \Delta^{33}\text{S} \) values (Kamber and Whitehouse, 2007). Similarly, SIMS \( \text{Fe} \) isotope studies have also shown large ranges of \( \delta^{56}\text{Fe} \) values, from +0.9% to +5.2% in a single magnetite grain (Marin-Carbonne et al., 2011) and from −4.2% to +2.9% in pyrites from the 2.72 Ga Tumbiana Formation (Yoshiya et al., 2012). Such variations have been interpreted as indicating multi-stage mineral formation and/or mineral alteration processes, and highlight the importance of spatially resolved analyses to better constrain processes in the water column, and during diagenesis and, possibly, metamorphism. Pyrite nodules in Archean shales have been extensively investigated in the context of the diagenetic history of ancient sedimentary rocks and have been used as a proxy for paleoenvironmental conditions on the early Earth (Bekker et al., 2004; Kakegawa et al., 1998; Ono et al., 2009, 2003; Rouxel et al., 2005). Whether there exists any isotopic variability in \( \text{Fe} \) within individual pyrite nodules is not known.

Herein, we present bulk measurements along with \textit{in-situ} isotopic (\( \text{Fe} \) and \( \text{S} \)) and trace element analyses of pyrite nodules hosted in carbonaceous shale from the Late Archean (2.83–2.70 Ga) Bubi Greenstone Belt, Zimbabwe. We use these data to constrain the origin and growth history of these pyrite nodules and to explore the possibility of a microbially-induced fractionation.

2. Samples and methods

2.1. Samples

We investigated pyrite nodules present in Late Archean carbonaceous shales from core 690B92-02 (see supplementary material S1) drilled along the eastern margin of the Bubi Greenstone Belt (Zimbabwe) north of the Damba nickel prospect (Hofmann et al., 2014; Prendergast, 2003). The drill core intersects basalt and underlying carbonaceous shale (see supplementary material S2 for the detailed stratigraphic log), which are considered correlative with the 2.7 Ga Reliance Formation and the 2.83 to 2.70 Ga Manjeri Formation of the Beulingwe Greenstone Belt in Zimbabwe (Hofmann and Kusky, 2004; Prendergast, 2003; Stone et al., 1994). The metamorphic grade is not well-established, but rocks in the Bubi Greenstone Belt have been subjected to lower greenschist grade metamorphism at most (Dziggel et al., 1998; Saggerson and Turner, 1976).

Pyrite occurs as nodules (up to 1 cm in diameter), discontinuous laminae (up to 0.5 cm thick), rare veinlets and disseminated grains (< 20 μm in size) in the shale (Fig. 1). Pyrite formed before burial compaction at the stage of early diagenesis as indicated by ellipsoidal shape of nodules with the long axis parallel to bedding, laminations in shale that bend around the nodules and rare pytymatically folded veinlets perpendicular to bedding. Nodules and laminae consist of either massive pyrite, a fine intergrowth of pyrite with the host matrix, or concentrations of fine pyrite crystals in the host material. The margin of nodules and laminae frequently consist of relatively coarse pyrite crystals suggestive of recrystallization. Pyrite represents the only observed Fe-sulfide mineral phase, with the exception of minor chalcopirite crystals in the more coarsely-crystalline domains. Nodules frequently contain marginal pressure shadows elongated parallel to bedding that consists of quartz, phyllosilicates, and rare coarsely-crystalline pyrite. A foliation is present in the shale and oriented at an oblique angle to bedding. A summary of pyrite textures is shown in Fig. 1.

Our study focuses on a single sample of shale (at depth of 144.5 m in drill core) containing several circular to slightly ellipsoidal pyrite nodules (Fig. 2). X-ray diffraction analysis (X-PertPro, University of Johannesburg) indicates that the shale mainly consists of quartz, illite, Fe-chlorite, and minor calcite mineral phases. The shale is relatively rich in organic carbon with \( \text{C}_{\text{org}} \) content ranging from 6.4 to 9.3 wt.% (Eltra Elemental Analyzer, University of Manitoba).

2.2. Methods

A slab of shale containing two, immediately adjacent pyrite nodules (called nodule 1 and 2 in Fig. 2) was embedded in epoxy, polished with 1 μm size diamond paste and Au-coated for \textit{in-situ} isotopic measurements (see supplementary material S3).

2.2.1. Bulk sample Fe and S isotope analyses

In the following sections, iron and sulfur isotopes are expressed in delta notation (\( \delta^{56}\text{Fe} \), \( \delta^{33}\text{S} \), and \( \delta^{34}\text{S} \)) relative to the international standards IRMM 14 (for \( \text{Fe} \)) and V-CDT (for \( \text{S} \)) based on the following equation:

\[
\delta A = \frac{(2A)^{1/2}A_{\text{sample}}}{(2A)^{1/2}A_{\text{standard}}} - 1 \times 1000
\]

where \( A \) is \( \text{Fe} \) or \( \text{S} \), and 1 and 2 represent the heavy and light isotopes, respectively (54 and 56 for \( \text{Fe} \) and 34 or 33 and 32 for \( \text{S} \)).
Fig. 1. Photomicrographs (reflected light) of pyrite textures. A. Ellipsoidal pyrite nodule. B. Portion of pyrite nodule with rim and irregular central domain of recrystallized pyrite. C. Disseminated pyrite in shale matrix near pyrite nodule. D. Portion of B showing finely intergrown pyrite and matrix silicates of nodule centre, recrystallized pyrite rim and disseminated pyrite in shale matrix. E. Pyrite lamina in shale showing recrystallized rim with chalcopyrite (arrows). F. Disseminated pyrite within nodular structure. G. Portion of pyrite nodule with coarse pyrite present in pressure shadow (top right). H. Same as G showing chalcopyrite inclusions (arrows).
Fig. 2. a. Scanning electron microscopy (SEM) images of the polished surface of the investigated pyrite nodules. The white dots indicate the in-situ Fe and S isotope and trace element spot analyses. b. The Co and Ni concentration (wt.%, EPMA) profile of nodule 1 shows difference between the coarse-grained rim and the fine-grained core (see text for the definition of core and rim). c. An SEM image of the core-rim transition zone showing textural difference.

Mass-independent fractionation has been calculated as the deviation from the Terrestrial Fractionation Line (TFL), using the mass-discrimination law (Farquhar et al., 2000):

$$\Delta^{33}S = \left( \ln \left( \frac{^{33}S}{1000} + 1 \right) - 0.515 \times \ln \left( \frac{^{34}S}{1000} + 1 \right) \right),$$

where the factor 0.515 defines the slope of the TFL.

Pyrite nodules from the same sample of shale used for in-situ work were drilled, crushed and millimetric-sized pyrite particles were hand-picked. Sulfur isotope ratios of pyrite particles were determined at the Geophysical Laboratory using techniques described by Hu et al. (2003). Pyrite particles (0.5 to 1 mg) from nodules were reacted with fluorine under a 25 W CO2 infrared laser at 25–30 Torr in a vacuum chamber to produce SF6, which was then purified by dual gas chromatography. Multiple sulfur isotope ratios were measured with a Thermo Scientific MAT 253 mass spectrometer in dual inlet mode (Hofmann et al., 2009; Ono et al., 2009). The precision for $\delta^{34}S$, $\delta^{33}S$, and $\Delta^{33}S$ values was determined by the multiple analyses of CDT material and internal reference materials (Maine and Alpha Aesar pyrite) and is better than 0.34%, 0.19%, and 0.03%, respectively (2σ).

Fe isotope compositions were measured following the procedure described in Rouxel et al. (2005). Hand-picked particles from pyrite nodules were dissolved in concentrated HNO3–HCl acid mixture and Fe was purified on Bio-Rad AG1X8 anion resin. Fe isotope ratios were determined with a Thermo Scientific Neptune multicollector inductively-coupled plasma mass-spectrometer operated at IFREMER, Pole Spectrometry Ocean, Brest in France. Long-term reproducibility of $\delta^{56}Fe$ measurements was determined on duplicate analysis of reference material and is about 0.08% (2σ).

2.2.3. SIMS analyses

Iron isotope compositions were measured in-situ with a Cameca ims 1270 ion microprobe at both CRPG (Nancy, France) and UCLA (Los Angeles, USA) following the procedure described in detail in Marin-Carbonne et al. (2011). Briefly, a $^{16}O^-$ primary beam of about 10 nA intensity was focused to a spot of about 15 μm. The mass resolution was set $\sim$7,000 and $^{54}Fe^+$ and $^{56}Fe^+$ were measured in multicollection mode with two off-axis Faraday cups. The gains of these Faraday cups were determined at the beginning of the analytical session and drift was monitored by frequent analyses of standards interspersed among analyses of the unknowns. The background of each detector was measured during the pre-sputtering for 1 min, i.e. at the beginning of each analysis. Ion currents converted to count rates were typically $\sim$2.105 counts per second (cps) for $^{56}Fe$. An analysis consisted of 30 cycles with 5 s acquisition time. Chromium was monitored on the masses 52 and 53 by using electron multipliers, but chromium levels were negligible in all samples. The internal precision for $\delta^{56}Fe$ 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 and Fedo, 2007) was better than 0.2% (2σ).
Sulfur isotope compositions were measured on the Cameca ims 1280 HR2 (CRPG, Nancy, France) by simultaneous measurements of $^{35}$S, $^{34}$S, and $^{33}$S in multicollection mode with three off-axis Faraday cups. The relative gains of the Faraday cups were intercalibrated at the beginning of each analytical session. The analytical method is described in detail in Thomassot et al. (2009) and Philippot et al. (2012) and is only summarized here. A Cs$^+$ primary beam of 5 nA intensity was focused to a spot of about 15–20 μm. Typical $^{32}$S$^+$ intensity was between 6 and 10.10$^6$ counts per second (cps) depending on the sulfide mineral analyzed. Several pyrite standards (Maine, Philippot et al., 2012 and Balmat pyrite, courtesy of M. Whitehouse) were used to determine the instrumental mass fractionation, and (ii) the reference mass discrimination line, from which $^\Delta ^{32}$S values were calculated. A typical analysis consists of 2 min of presputtering followed by data acquisition in 30 cycles of 3 s each. The background of the detectors was measured during the presputtering and was then corrected for each analysis. The internal precision achieved under these conditions was better than 0.05‰ for $^\delta ^{32}$S and 0.03‰ for $^\delta ^{34}$S values (2σ). The external precision, which is the standard deviation calculated from repeated measurements on various reference materials, was 0.40‰ (2σ) for $^\delta ^{34}$S and 0.06‰ (2σ) for $^\Delta ^{33}$S values.

3. Results

Fe and S isotope compositions and EPMA trace element concentrations have been obtained across each nodule (see supplementary material S3 and S4), while LA-ICP-MS trace element concentration profiles were performed only for nodule 2 (see supplementary material S4).

3.1. $^\delta ^{56}$Fe variations

The two nodules show similar variations in Fe isotope composition (Figs. 3a, b). The variations are large for both nodules from −2.11 to +0.45‰ for nodule 1 and from −1.58 to +0.75‰ for nodule 2, corresponding to about 2/3 of the total $^\delta ^{56}$Fe range measured so far in bulk terrestrial rocks (Dauphas and Rouxel, 2006; Johnson et al., 2008). The total range for $^\delta ^{56}$Fe values from the edge to the centre is ∼2.5‰ in nodule 1 (Fig. 3a) and ∼2.3‰ in nodule 2 (Fig. 3b). The variations from the margin to the centre are consistent for both nodules. Both nodules have similar $^\delta ^{56}$Fe values for the rims with a mean value of −0.4 ± 0.4‰ for nodule 1 and −0.2 ± 0.4‰ for nodule 2. The centre of the nodules is $^{56}$Fe-depleted with respect to the margins with a mean $^\delta ^{56}$Fe value of −1.7 ± 0.3‰ for nodule 1 and −1.1 ± 0.5‰ for nodule 2. The mean $^\delta ^{56}$Fe values of the nodules calculated with SIMS data are −1.1 ± 0.7‰ for nodule 1 and −0.5 ± 0.8‰ for nodule 2 and are consistent with the bulk pyrite nodule value of −1.46 ± 0.1‰ measured on pyrite nodule fragments from the same hand-specimen.

3.2. $^\delta ^{34}$S variations

The $^\delta ^{34}$S values are highly variable across the nodules, without any systematic trend for nodule 1 (Fig. 3c). The centre of nodule 2, having $^\delta ^{34}$S values close to +1‰, is more depleted in $^{34}$S than the rim with $^\delta ^{34}$S values as high as +8‰ (Fig. 3d), although there is considerable scatter in the $^\delta ^{34}$S values of the rim. The total range of $^\delta ^{34}$S values is 5.4‰ for nodule 1 and 7.8‰ for nodule 2, with the mean values calculated with SIMS data of +2.9 ± 1.3‰ for nodule 1 and +2.2 ± 2.6‰ for nodule 2. These mean values are consistent with the bulk pyrite nodule value of +3.18 ± 0.34‰, measured on pyrite nodule fragments from the same depth.

3.3. $^\Delta ^{33}$S variations

Mass-independently fractionated sulfur (S-MIF) is present in both nodules, with $^\Delta ^{33}$S values ranging between −1.5 and +2.9‰ for nodule 1 and between −1.6‰ and +1.9‰ for nodule 2 (Fig. 3e, f). Nodule rims display mostly positive $^\Delta ^{33}$S values, while their centres show predominantly negative $^\Delta ^{33}$S values. The centre has a mean value of −0.7 ± 0.5‰ for nodule 1 (Fig. 3e) and −0.7 ± 0.6‰ for nodule 2 (Fig. 3f). The mean $^\Delta ^{33}$S values are −0.04 ± 1.4‰ (nodule 1) and −0.24 ± 1.1‰ (nodule 2), which are comparable to the bulk pyrite nodule value of 0.31 ± 0.03‰, considering the high variability of $^\Delta ^{33}$S values at a micrometer scale measured by SIMS. The negative $^\Delta ^{33}$S values of pyrite nodule cores are in the range of values measured for Archean hydrothermal barites and sulfides (Bao et al., 2007; Farquhar et al., 2000; Philippot et al., 2012; Shen et al., 2009; Ueno et al., 2008). The outer rims have positive $^\Delta ^{33}$S values up to +2.47‰, with a mean value of +0.64 ± 1.30‰ for nodule 1 and +0.6 ± 1.30‰ for nodule 2, in the range of values for disseminated pyrite in sedimentary rocks of this age (Farquhar et al., 2000; Ono et al., 2009).

4. Discussion

4.1. Nodule composed of two chemically and isotopically distinct parts: a core and a rim

Each nodule is characterized by a core (∼3000 μm in width) and a more coarsely-crystalline rim (Fig. 1). Both nodules broadly display similar Fe and S isotope trends from the core to the margin, thus potentially preserving information regarding similar processes and diagenetic histories. The rim and the core have different trace element concentrations and, more importantly, different isotopic compositions (Figs. 2 and 3). The core is characterized by predominantly negative $^\Delta ^{33}$S values, positive $^\delta ^{34}$S values, and negative $^\delta ^{56}$Fe values. The rims have predominantly positive $^\Delta ^{33}$S values, close to zero $^\delta ^{56}$Fe values and high Co and low Ni contents. The cores of both nodules show a positive correlation between $^\delta ^{34}$S and $^\Delta ^{33}$S values (Fig. 4), and an antithetic trend between $^\delta ^{56}$Fe and $^\Delta ^{33}$S values (Fig. 5).

The nodules have a bimodal distribution of Fe isotope values (Fig. 6, right), with two major peaks at −1.5‰ and 0‰. Moreover, both nodules display a $^\Delta ^{33}$S range from −1.5 to +2.9‰ (Fig. 6, left), with two modes at −1‰ and +1.5‰. Nodule 1 shows also another mode at +3‰. These Fe and S isotope distributions rule out any continuous process of pyrite nodule growth. The large ranges in isotopic composition could have been produced during either diagenesis or burial in association with fluid circulation. While metasomatic and/or metamorphic processes could fractionate both Fe and S isotopes, they cannot produce MIF of sulfur (e.g., Johnston, 2011). Moreover, textures of the pyrite nodules and hosting black shales suggest that these nodules were formed before burial compaction, and at the stage of early diagenesis. Furthermore, large range of $^\Delta ^{33}$S values, from −1.5 to +2.9‰, cannot be explained by equilibrium or kinetic dissolution–precipitation processes. Although redox processing of S may produce fractionations having a slightly different slope from that of the terrestrial fractionation line in the $^\delta ^{34}$S vs. $^\Delta ^{33}$S space (Farquhar et al., 2010; Johnston, 2011), these processes cannot explain this range in $^\Delta ^{33}$S values. Moreover, limited range of the $^\delta ^{34}$S values observed in these nodules implies small deviation from terrestrial fractionation line due to biological sulfur processing (Canfield, 2001; Sim et al., 2011). Thus, this range of $^\Delta ^{33}$S values can only be explained by different S sources that were present in the sediments, including the pyrite precursors themselves, diagenetic waters, or
Pyrite precipitation can be induced by both bacterial sulfate reduction (BSR) and dissimilatory iron reduction (DIR). Archer and Vance (2006) proposed that pyrite formed when these processes are coupled would display co-variation between δ56Fe and δ34S values, however, we do not observe such relationship (Fig. 7). The absence of co-variation between δ56Fe and δ34S values does not rule out microbial influence on pyrite formation, but suggests that

hydrothermal and metamorphic fluids. Whether those different S sources may also correspond to different Fe sources with distinct δ56Fe values requires thorough evaluation of diagenetic processes and pyrite formation pathways (Johnston, 2011). As the trend in Fe and S isotope composition is better defined for nodule 1, the following discussion will be mainly based on the results from this nodule.

Fig. 3. δ56Fe (a, b), δ34S (c, d) and Δ33S (e, f) profiles across the pyrite nodule 1 (circles) and nodule 2 (squares). Both nodules reveal large, micrometer-scale isotopic heterogeneity, but nodule 1 shows more clearly defined trends in isotopic composition from the centre to the rim. The error bars are comprised in the symbols. Note that the textural boundary between the core and the rim does not exactly match the change in isotopic composition, especially for the Δ33S values.
nodules were not simply formed by sulfate and Fe(III) reduction in tandem.

Pyrite nodules in shales are thought to be formed by the dissolution of disseminated Fe sulfides present in the shale (Berner, 1984; Coleman and Raiswell, 1981; Kakegawa et al., 1998). The chemical reactions resulting in crystallization of pyrite are complex and involve, in addition to sulfate, sulfide, and Fe(II), several intermediate species such as thiosulfate, elemental sulfur, FeS, and Fe$_2$S$_3$ among others (see Rickard, 2013 for an extensive review). However experimental studies have suggested that pyrite can be easily formed from two different precursors, mackinawite, FeS, or greigite, Fe$_3$S$_4$ (e.g., Rickard, 2013). Pyrite formation is kinetically controlled and requires iron loss from, or sulfur addition to, a pyrite precursor, which can be iron monosulfide, like FeS$_m$ (mackinawite), or iron polysulfide, like Fe$_2$S$_4$ (greigite). In the case of mackinawite as a pyrite precursor, pyrite formation requires the presence of H$_2$S:

$$\text{FeS}_m \rightarrow \text{FeS}_0 + \text{H}_2\text{S} = \text{FeS}_2 + \text{H}_2$$

If greigite is the pyrite precursor, pyrite forms via a reaction between polysulfide and soluble FeS (Rickard and Luther, 2007). In the following, we will consider that pyrite nodules were formed from pyrite precursors, monosulfides and polysulfides.

4.2. Origin of the isotopic variability in the core

The core of nodule 1 shows a large S-MIF range of 2‰ and a positive co-variation between $\Delta^{33}$S and $\delta^{34}$S values (Fig. 4). The
Fig. 6. Histogram of $\Delta^{33}S$ (left) and $\delta^{56}Fe$ (right) values for both nodules (nodule 1 in grey and nodule 2 in white). The pyrite nodules show two main $\Delta^{33}S$ peaks, one with positive $\Delta^{33}S$ value and another one with negative $\Delta^{33}S$ value and two main $\delta^{56}Fe$ peaks, one with negative $\delta^{56}Fe$ value and another one with $\delta^{56}Fe$ value close to 0%. The similar isotopic compositions suggest a shared origin of the two nodules.

Fig. 7. $\delta^{56}Fe$ and $\Delta^{33}S$ values for nodule 1 (in white) and nodule 2 (in black). Previous data obtained for the Belingwe pyrite by Archer and Vance (2006) is plotted in grey, and the DIR and BSR processes are represented by the two curves reflecting different dissolved Fe concentrations.

The co-variation on the scale of the core of the pyrite nodule reflects $S$ contribution from two different sources within the relatively small volume of hosting black shales. Although similar positive co-variation on the scale of the stratigraphically thick sedimentary sections has been linked to temporal changes in composition of atmospherically-derived sulfur-bearing aerosols (e.g., Farquhar and Wing, 2003; Kaufmann et al., 2007; Ono et al., 2009), our in situ analyses of pyrite nodules indicate mixing of sulfur derived from two end-member sources hosted within a relatively small volume of black shale sediments deposited almost instantaneously on a geological timescale. Similar $\delta^{34}S$–$\Delta^{33}S$ trends have already been observed in Archean sedimentary pyrites (Ono et al., 2003; Farquhar et al., 2013) and has been explained by mixing between two compositionally distinct sources. These two sources are probably a monosulfide pyrite precursor (FeSm) and a fluid bearing $H_2S$, according to the pyrite formation equation above. Both $\delta^{34}S$ and $\delta^{33}S$ values of the sources can be calculated from mass balance:

$$x_{FeSm} \times \delta^{3X}_{FeSm} + x_{H_2S} \times \delta^{3X}_{H_2S} = x_{pyrite} \times \delta^{3X}_{pyrite}$$

where $X$ is 3 or 4, and the $\Delta^{33}S$ value is then calculated from the $\delta^{34}S$ and $\delta^{33}S$ values and $x$ is the proportion of total $S$ in the different $S$ species, such as FeSm, $H_2S$, and pyrite. In our case, $x_{pyrite} = x_{FeSm} + x_{H_2S}$. The $\delta^{34}S$ values of FeSm and $H_2S$ reflect the isotopic composition of $H_2SO_4$ and $S_8$ aerosols, respectively, as further discussed below.

Two end-members can be defined to reproduce the $\delta^{34}S$ and $\delta^{33}S$ relationships, one with negative $\Delta^{33}S$ and unfractionated $\delta^{34}S$ values and the other one with $\delta^{34}S$ values ranging from +1.8 to +8% and $\Delta^{33}S$ values close to +0.6%. With these two end-members, almost all $S$ isotope compositions of the cores of both nodules can be reproduced (Fig. 4, right). $H_2SO_4$ aerosols are formed in the atmosphere and delivered to the ocean, and $H_2SO_4$ is preferentially processed in the water column to form $H_2S$ with negative $\Delta^{33}S$ values. $H_2S$ can then react with Fe in the water column to form small particles of iron monosulfide, such as mackinawite (Rickard and Luther, 2007), which would record the negative S-MIF of the sulfur source. During early diagenesis in sediments, mackinawite would dissolve contributing to the growth of larger pyrite nodules by Ostwald ripening effect (cf. Kakegawa et al., 1998). Elemental sulfur with positive $\Delta^{33}S$ values that was also produced via photochemical reactions in oxygen-free atmosphere was preferentially delivered to the sediments rather than utilized in the water column as $S_8$ is insoluble (Rickard, 2013) and its biological intake and processing in the Archean, oxygen-free ocean would be slower than that of dissolved sulfate. In pore waters, $S_8$ rings would be gradually open and sulfur chains and compounds would be biologically converted to $H_2S$. Various microbial organisms are able to disproportionate elemental sulfur in the sediments and promote pyrite formation (e.g., Johnston, 2011). Thus the core
of the pyrite nodule was formed by mixing of sulfur derived from the dissolution of the mackinawite and elemental sulfur present in sediment pore waters.

Crystallization of pyrite forming the nodule core and resulting from the dissolution of mackinawite can be modeled as a Rayleigh distillation process to explain the Fe isotope variations across the nodule core. In this model, the \( \delta^{56}\text{Fe} \) value of each increment of precipitated pyrite is controlled by the isotopic fractionation between dissolved FeSm and FeS2 (\( \Delta_{\text{pyrite-FeSm dissolved}} \)) and by the evolving composition of the fluid, which is controlled by the fraction (\( f \)) of FeSm remaining in solution:

\[
\delta^{56}\text{Fe}_{\text{pyrite}} = \delta^{56}\text{Fe} (\text{FeSm dissolved})^f + \Delta_{\text{pyrite-FeSm dissolved}} \times (1 + \ln f)
\]

For this calculation, we adopt an isotopic fractionation factor (\( \Delta_{\text{pyrite-FeSm dissolved}} \)) of \(-1\%\) for pyrite formation according to theoretical calculations (Polyakov et al., 2007; Polyakov and Soutanov, 2011) and recent experiments (Guilbaud et al., 2011). Rayleigh fractionation can explain the \( \delta^{56}\text{Fe} \) distribution for the core of nodule 1 by near total consumption of dissolved FeSm (Fig. 8). Considering a fluid with an initial \( \delta^{56}\text{Fe} \) value of \(-1.2\%\) and for \( f \) from 1 to 0.08, the calculated \( \delta^{56}\text{Fe}_{\text{pyrite}} \) values range from \(-2.2\) to \(+1.78\%\), which is in agreement with the measured range (from \(-2.11\) to \(+0.45\%\)). Thus, the cores of the pyrite nodules could have formed in a closed with respect to Fe pore-water system with iron supplied by the dissolution of a mackinawite in the sediments. This result also implies that the rim was also formed under similar Fe-limited pore-water conditions. Assuming that the mackinawite dissolution produced a pore water fluid with negative \( \delta^{56}\text{Fe} \) value of approximately \(-1.2\%\), reflecting the average Fe isotope composition of the mackinawite, and using the experimentally-determined Fe isotope equilibrium fractionation factor between mackinawite and Fe(II) of around \(-0.3\%\) at 25°C (Butler et al., 2005; Guilbaud et al., 2011), the mackinawite in the water column should have crystallized from a source with \( \delta^{56}\text{Fe} \approx -0.9\%\).

We infer that the core of the pyrite nodule formed during early diagenesis by the dissolution of mackinawite, which would have had negative \( \delta^{56}\text{Fe} \) values, near to 0%, \( \delta^{34}\text{S} \) values, and negative \( \Delta^{33}\text{S} \) values. Such a pyrite precursor could have formed in an aqueous solution, possibly in the water column or at the water–sediment interface. The negative \( \delta^{56}\text{Fe} \) value of FeSm reflects composition of dissolved Fe(II) depleted in heavy isotopes. In principle, such Fe isotope composition can be produced via two main pathways: (i) DIR during diagenesis releasing Fe(II) depleted in heavy isotopes (\( \delta^{56}\text{Fe} \approx -2\%\)) into pore water, which is transferred to pyrite precursor by precipitation, as in modern anoxic environments (Johnson et al., 2008, 2004; Percal-Dennett et al., 2011; Severmann et al., 2006), and (ii) oxidation to Fe(III) and precipitation of Fe-oxhydroxides during iron cycling in the water column producing a residual Fe(II) pool depleted in heavy isotopes (Rouxel et al., 2008, 2005; Tsikos et al., 2010). The presence of negative S-MIF indicates that the mackinawite was formed by sulfate reduction, likely close to the water–sediment interface, where both DIR and partial iron oxidation processes are possible and could explain the negative \( \delta^{56}\text{Fe} \) values. In the case of DIR, pore waters should have had a negative \( \delta^{56}\text{Fe} \) value of approximately \(-1\%\) to explain the isotopic composition of the pyrite precursor. Although it is known that pore waters near the seawater–sediment interface in modern marine settings can have \( \delta^{56}\text{Fe} \) values down to \(-3.4\%\) in some extreme cases (Hinokawa et al., 2009; Rouxel et al., 2008; Severmann et al., 2006), more typically they display an average value around 0% in anoxic environments (Severmann et al., 2006). The second pathway suggests that large-scale seawater redox cycling could be responsible for the origin of negative \( \delta^{56}\text{Fe} \) values of pyrite precursor (Planavsky et al., 2012; Rouxel et al., 2005). The preferential sequestration of heavy Fe isotopes into Fe-oxhydroxides would produce an isotopically negative Fe(II) pool. This pathway assumes that the pyrite precursor with negative \( \delta^{56}\text{Fe} \) values reflects the presence of an isotopically light reservoir in the water column. Evidence for this process is found in the Fe isotope composition of magnetite and hematite in iron formations, which are typically characterized by positive \( \delta^{56}\text{Fe} \) values (Johnson et al., 2003; Planavsky et al., 2012). Although it still remains largely unresolved whether precipitation of Fe oxhydroxides (cf., Rouxel et al., 2005) or iron-shuttle from continental margins to deep-water, driven by dissimilatory iron reduction (Severmann et al., 2006, 2008) was a major control over Fe isotope composition of seawater, both processes would contribute to negative Fe isotope composition of seawater. The pyrite precursor of the nodule core, mackinawite, was thus formed in the water column by sulfate reduction, as indicated by S-MIF, and from a dissolved iron pool with negative \( \delta^{56}\text{Fe} \) values.

4.3. Origin of the isotopic composition of the rim

The nodule rims display different Fe and S isotope compositions from their cores and a coarse-grained texture, clearly indicating distinct history from the cores. First and foremost, the rim has \( \delta^{56}\text{Fe} \) values close to 0% and mostly positive \( \Delta^{33}\text{S} \) values, the latter suggestive of a sulfur source derived from \( \text{S}_8 \) aerosols. Elemental sulfur is highly insoluble and less amenable to biological processes; as a result, most of it is transferred to sediments as solid S (Ono et al., 2003). Pyrite cannot form directly from elemental solid S (Rickard and Luther, 2007), which implies an intermediate step to break the \( \text{S}_8 \) rings (Rickard, 2013) and react with a dissolved pyrite precursor. Elemental solid S can react with dissolved mackinawite present in the sediment pore waters to form greigite as described in the following equation:

\[
3\text{FeSm} + S_8 = \text{FeS}_4
\]
Greigite can form directly from mackinawite in anoxic and acidic environments, in which low pH enhances the dissolution of mackinawite (Rickard and Luther, 2007). Greigite formation from mackinawite can also be promoted by sulfate-reducing bacteria (Rickard and Luther, 2007). If pyrite precursor, greigite, was formed during an early stage of diagenesis, when the mackinawite was still present in the sediment, by the reduction of the S₈ in a highly reducing fluid (e.g., due to the presence of carbonaceous matter), then the variations in Δ³³S composition of the pyrite nodule rim must be due to mixing between S from FeS₈ and S⁰ sources. With respect to Fe isotope values, pyrite nodule rims may reflect mixing between negative Fe isotope values of mackinawite and pore water fluid, derived from dissolution of reactive iron in sediments, e.g., Fe-oxyhydroxides. Thus, pyrite nodule rims could have been formed by the dissolution of the same mackinawite as the nodule cores in contact with a reducing fluid in sediments containing elemental sulfur. Different composition of cores and rims suggests that the mackinawite dissolution started earlier in the diagenesis and, subsequently, overlapped in time with the dissolution of elemental sulfur. That is why the inner part of the rim has still negative Δ³³S values, largely inherited from the mackinawite, gradually giving away to positive Δ³³S values, derived from the elemental sulfur, towards the outer part of the rim.

Complicating this picture further are trace element zonation patterns and petrographic observations. The rim has a coarse-grained texture associated with nickel depletion and cobalt enrichment, suggesting its growth during burial diagenesis or in association with post-depositional processes such as metasomatism (Large et al., 2009; Steadman et al., 2013). In this case, the fluid should have carried sulfur with positive MIF-S as observed on the rims, which implies that either the fluid had an access to an atmospheric source or was enriched in sulfur by dissolution of sulfides bearing MIF-S. The rim pyrite has different S and Fe isotope composition from that of the core of the pyrite nodule and thus, by extension, from that of the core pyrite precursor, mackinawite. The most parsimonious scenario is that the pyrite rim was formed by the dissolution of the small, disseminated greigite grains during burial. In this case, the trace element composition of the rim was acquired from the late diagenetic or metasomatic fluids different in trace element composition from the early diagenetic fluids, while the Fe and S isotope compositions of the rim reflect the pyrite precursor, i.e., greigite, and reactive iron in sediments. Disseminated pyrite could have also formed from the disseminated greigite precursor.

It would be useful to test this hypothesis by analyzing individual disseminated pyrite grains, however their small size (around 2 μm) precludes this approach.

4.4. Nodule formation model and paleoenvironmental implications

Pyrite nodules were formed from two different precursors, mackinawite and greigite, which have recorded different environmental conditions. Mackinawite formed in the water column from the sulfate pool and provides constraints about this environment, while greigite formed in the sediment via reaction of mackinawite with atmospherically-derived elemental sulfur that was present in the sediment pore-waters and records late-stage diagenetic and/or burial conditions. Mackinawite crystallized from sulfide likely produced by BSR and from dissolved iron (Fig. 9a), suggesting an Fe-rich deep water, whereas greigite was formed from dissolved mackinawite and elemental sulfur in pore waters (Fig. 9b). The pyrite nodule core was formed by the dissolution of mackinawite in a closed system with a limited-size Fe reservoir and by mixing of sulfur derived from the pyrite precursor, mackinawite, and elemental sulfur in the pore water (Fig. 9a). At some point, mackinawite was consumed and the sediment contained excess of elemental sulfur. Some of the remaining mackinawite present in sediments reacted in a solid state with elemental sulfur to form small disseminated greigite crystals (Fig. 9b). During burial at a higher temperature and in the presence of the late-stage diagenetic or metasomatic fluid, the disseminated greigite grains started to dissolve aggregating into coarse-grained pyrite nodule rims and forming small, disseminated pyrite grains in the sediment (Fig. 9c).

The δ⁵⁶Fe and δ³⁴S isotope values do not show co-variation (Fig. 7) previously inferred for the Belingwe sulfides (Archer and Vance, 2006), challenging the notion of coupled S and Fe reduction. Furthermore, while Fe isotope values become progressively more positive towards the rim and can be explained by Rayleigh distillation in the closed diagenetic system with respect to Fe, δ³⁴S values do not show any systematic trend from the core to the rim. As a result, our data is consistent with the diagenetic setting limited with respect to Fe, but unlimited with respect to S. The
rims are clearly formed after the core, but still likely with S source hosted in the same sedimentary package as evidenced by the presence of MIF in S isotopes. Considering high organic carbon loading in this setting, Fe reduction would have been quantitative and Fe isotope fractionations during dissimilatory Fe reduction should not be expressed. These observations rule out the model proposed by Archer and Vance (2006) based on coupled Fe and S reduction. Furthermore, we inferred highly fractionated by precipitation of Fe-oxyhydroxides seawater based on the average Fe isotope values of the cores of the pyrite nodules (−1.7 ± 0.3‰ for nodule 1 and −1.1 ± 0.5‰ for nodule 2), equilibrium Fe isotope fractionation during precipitation of mackinawite (−0.3 to −0.9‰; Butler et al., 2005; Guibaud et al., 2011), and possible range of δ34Fe values of the Fe sources to the Archean seawater (0.0 to −0.5‰; Rouxel et al., 2005). We concur with Guibaud et al. (2001) that pyrite can be formed by the Rayleigh distillation process based on a partial Fe(II)aq utilization during abiotic pyrite precipitation. In-situ analysis is a relevant approach to test this model. However, we did not observe the highly fractionated Fe isotope values predicted by the model. Instead, the Rayleigh distillation modeling of the fractionated Fe isotope compositions of the cores of our nodules reflects the almost total Fe-utilization in a closed system. We therefore infer that the highly fractionated, average Fe isotope composition of the cores of the pyrite nodules corresponds to the Fe isotope composition of seawater. Rouxel et al. (2005) proposed that negative Fe isotope values of bulk pyrite nodules in Archean organic matter-rich shales reflect composition of Archean seawater influenced by Fe-oxyhydroxide precipitation. Our study also shows micro-scale Fe isotope heterogeneity within pyrite nodules reflecting fractionations in the diagenetic realm.

The nodules display S-MIF with both positive and negative values, which suggest the presence of two distinct sulfur pools in the diagenetic realm; one being mackinawite derived from soluble sulfate and the other being less reactive pool of insoluble, atmospherically-derived elemental sulfur as already proposed by Farquhar et al. (2013). Importantly, these well-defined S sources were not entirely mixed on a micrometer-scale to erase their distinct S isotope signatures during S processing in sediment pore-
wides. We relate this unique preservation of atmospheric S signature during diagenetic processing to the difference in reactivity between these two pools with the insoluble sulfur compounds derived from atmospheric elemental sulfur being less reactive with respect to the seawater sulfate. Fe and S isotope variations observed in the Archean pyrite nodules reflect the contribution of two atmospherically-derived S sources and Fe pool fractionated by precipitation of iron oxohydroxides as well as by diagenetic processes involved in the growth of pyrite nodules.

5. Conclusions

In-situ coupled Fe and S isotope study of the c. 2.7 Ga pyrite nodules have revealed an extreme isotopic variability both in Fe and S at the micrometer scale (see Table 1). It has been shown for the first time that these nodules have both positive and negative \( \Delta^{33}S \) values and highly variable Fe isotope composition. They reveal a complex crystallization history with, at least, two steps of dissolution and precipitation of two different pyrite precursors involved in the Ostwald ripening process. The core of the pyrite nodules has grown by dissolution of fine-grained mackinawite precipitated in the water column, while the rims were formed during the late diagenesis by dissolution of disseminated greigite grains crystallized during early diagenesis, when remaining dissolved Fe\(_{\text{sm}}\) clusters reacted with broken S\(_8\) rings. The nodules thus record S processing in two different environments, water column and sediment pore-waters, at the time when the Earth's atmosphere was still anoxic. They also indicate the presence of two distinct and yet contemporaneous seawater sulfur pools derived from the atmosphere, soluble sulfate and insoluble elemental sulfur, which, being different in their isotopic composition, are now resolvable at a micrometer-scale within the pyrite nodules. It also implies that temporal trends in \( \Delta^{33}S \) and \( \delta^{34}S \) values earlier inferred from bulk-rock S isotope analyses showing positive co-variation between \( \Delta^{33}S \) and \( \delta^{34}S \) values (e.g., Kaufmann et al., 2007; Ono et al., 2009) represent local variations in relative contribution of these two atmospherically-derived S sources and cannot be reliably used for global correlation of hosting sedimentary successions.

This study, based on two isotopic systems, reveals that pathways for pyrite nodule formation are complex and involve several pyrite precursors, such as mackinawite and greigite, and isotopically distinct S and Fe sources. However, the isotopic and elemental records spatially preserved in these pyrite nodules do constrain the conditions for the precipitation of pyrite precursors and thus can be used as proxies for paleoenvironmental conditions in the Archaean oceans and atmosphere. The study also illustrates the importance of in-situ and coupled isotopic studies to reveal diagenetic histories and pyrite formation pathways.

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Appendix A. Supplementary material

Supplementary material related to this article can be found on-line at http://dx.doi.org/10.1016/j.epsl.2014.02.009.

References
